Chemical Process Equipment

Selection and Design

Stanley M. Walas

Butterworth-Heinemann Series in Chemical Engineering

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Stanley M. Walas

Department of Chemical and Petroleum Engineering University of Kansas

To the memory of my parents, Stanislaus and Apolonia, and to my wife, Suzy Belle

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LIST OF EXAMPLES ix

PREFACE xi

RULES OF THUMB: SUMMARY' xiif

CHAPTER 1 INTRODUCTION |

- 1.1. Process Design I
- 1.2. Equipment 1 Vendors' Questionnaires 1 Specification Forms 1
- 1.3. Categories of Engineering Practice 1
- 1.4. Sources of Information for Process Design 2 1.5. Codes, Standards, and Recommended Practices 2
- 1.6. Material and Energy Balances 3 1.7. Economic Balance 4
- 1.8. Safety Factors 6
- *1.9.* Safety of Plant and Environment 7 1.10. Steam and Power Supply 9
- 1.11. Design Basis 12
- Utilities 12
- 1.12. Laboratory and Pilot Plant Work 12 References 15

CHAPTER 2 FLOWSHEETS 19

- 2.1. Block Flowsheets 19
- 2.2. Process Flowsheets 19
- 2.3. Mechanical (P&I) Flowsheets 19
- 2.4. Utility Flowsheets 19
- 2.5. Drawing of Flowsheets 20 References 31 Appendix 2.1 Descriptions of Example Process Flowsheets 33

CHAPTER 3 PROCESS CONTROL 39

- 3.1. Feedback Control 39 Symbols 39 Cascade (Reset) Control 42
- 3.2. Individual Process Variables 4.2 Temperature 42 Pressure 42 Level of Liquid 43 Flow Rate 43 Flow of Solids 43 Flow Ratio 43 Composition 43 3.3. Equipment Control 43
- Heat Transfer Equipment 44 Distillation Equipment 47 Liquid-Liquid Extraction Towers 50 Chemical Reactors 53 Liquid Pumps 55 Solids Feeders 55 Compressors 55 References 60

CHAPTER 4 DRIVERS FOR MOVING **EQUIPMENT 61**

- 4.1. Motors 61 Induction 61 Synchronous 61 Direct Current 61
- **4.2.** Steam Turbines and Gas Expanders 62 **4.3.** Combustion Gas Turbines and Engines 65 References 68

- CHAPTER 5 TRANSFER OF SOLIDS 69
- 51 Slurry Transport 69 Pneumatic Conveying 71 5.2. Equipment 72 Operating Conditions 73 Power Consumption and Pressure Drop 74
- Mechanical Conveyors and Elevators 76 5.3 Properties of Materials Handled 76 Screw Conveyors 76 Belt Conveyors 76 Bucket Elevators and Carriers 78 Continuous Flow Conveyor Elevators 82 5.4. Solids Feeders 83
- References 88

CHAPTER 6 FLOW OF FLUIDS 91

- 6.1. Properties and Units 91
- 6.2. Energy Balance of a Flowing Fluid 92
- 6.3. Liquids 94 Fittings and Valves 95 Orifices 95 Power Requirements 98
- 6.4. Pipeline Networks 98 6.5. Optimum Pipe Diamet
- Optimum Pipe Diameter 100
- 6.6. Non-Newtonian Liquids 100 Viscosity Behavior 100
- Pipeline Design 106 6.7. Gases 109 Isentropic Flow 109 Isothermal Flow in Uniform Ducts 110 Adiabatic Flow 110 Nonideal Gases 111
- 6.8. Liquid-Gas Flow in Pipelines 111 Homogeneous Model 113 Separated Flow Models 114 Other Aspects 114
- 6.9. Granular and Packed Beds 117 Single Phase Fluids 117 Two-Phase Flow 118
- 6.10. Gas-Solid Transfer 119 Choking Velocity 119
- Pressure Drop 119 6.11. Fluidization of Beds of Particles with Gases 120 Characteristics of Fluidization 123 Sizing Equipment 123 References 127

CHAPTER 7 FLUID TRANSPORT EQUIPMENT 129

- Piping 129 7.1. Valves 129 Control Valves 129 7.2. Pump Theory 131 Basic Relations 131 Pumping Systems 133 7.3. Pump Characteristics 134 7.4. Criteria for Selection of Pumps 140 7.5. Equipment for Gas Transport 143
- Fans 143 Compressors 145 Centrifugals 145 Axial Flow Compressors 146 Reciprocating Compressors 146 Rotary Compressors 149 7.6. Theory and Calculations of Gas Compression 153 **Dimensionless Groups 153** Ideal Gases 153 Real Processes and Gases 156
 - Work on Nonideal Gases 156

Efficiency 1.59 Temperature Rise, Compression Ratio, Volumetric

Efficiency 159 7.7. Ejector and Vacuum Systems Ejector Arrangements 162 162 Air Leakage 164 Steam Consumption 165 Ejector Theory 166 Glossary for Chapter 7 166 References 167

CHAPTER 8 HEAT TRANSFER AND HEAT EXCHANGERS 169

- 8.1. Conduction of Heat 169 Thermal Conductivity 169 Hollow Cvlinder 170 Composite Walls 170 Fluid Films 170
- 8.2. Mean Temperature Difference 172 Single Pass Exchanger 172 Multipass Exchangers 173 F-Method 173 O-Method 179 Selection of Shell-and-Tube Numbers of Passes 179 Example 179
- 8.3. Heat Transfer Coefficients 179 Overall Coefficients 180 Fouling Factors 180 Individual Film Coefficients 180 Metal Wall Resistance 18.2
- Dimensionless Groups 182 8.4. Data of Heat Transfer Coefficients 182 Direct Contact of Hot and Cold Streams 185 Natural Convection 186 Forced Convection 186 Condensation 187 Boiling 187
- Extended Surfaces 188 Pressure Drop in Heat Exchangers 188 8.5.
- 8.6. Types of Heat Exchangers 188 Plate-and-Frame Exchangers 189 Spiral Heat Exchangers 194 Compact (Plate-Fin) Exchangers 194 Air Coolers 194 Double Pipes 19.5
- 8.7. Shell-and-Tube Heat Exchangers 195 Construction 195 Advantages **199** Tube Side or Shell Side 199 Design of a Heat Exchanger 199 Tentative Design 200
- 8.8. Condensers 200 Condenser Configurations 204 Desien Calculation Method 205 The Silver-Bell-Ghaly Method 206
- 8.9. Reboilers 206 Kettle Reboilers 207 Horizontal Shell Side Thermosiphons 207 Vertical Thermosiphons 207 Forced Circulation Reboilers 208 Calculation Procedures 208
- 8.10 Evaporators 208 Thermal Economy 210 Surface Requirements 211
- 8.11. Fired Heaters 211 Description of Eauinment 211 Heat Transfer 213 Design of Fired Heaters 214
- 8.12. Insulation of Equipment 219 Low Temperatures 221
 - Medium Temperatures 221

Refractories 221 8.13. Refrigeration 224 Compression Refrigeration 224 Refrigerants 226 Absorption Refrigeration 229 Cryogenics 229 References 229 9 DRYERS AND COOLING TOWERS 231 9.1. Interaction of Air and Water 231 Rate of Drying 234 Laboratory and Pilot Plant Testing 237 9.2. 9.3. Classification and General Characteristics of Dryers 237 Products 240 Costs 240 Specification Forms 240 9.4. Batch Dryers 241 Continuous Tray and Conveyor Belt Dryers 242 9.5. Rotary Cylindrical Dryers 247 9.6. Drum Dryers for Solutions and Slurries 254 Pneumatic Conveying Dryers 255 9.7 9.8. Fluidized Bed Dryers 262 9.9. 9.10. Spray Dryers 268 Atomization 276 Applications 276 Thermal Efficiency 276 Design 276 9.11. Theory of Air-Water Interaction in Packed Towers 277 Tower Height 279 9.12. Cooling Towers 280 Water Factors 285 Testing and Acceptance 285 References 285

CHAPTER 10 MIXING AND AGITATION 287

10.1. A Basic Stirred Tank Design 287 The Vessel 287 Baffles 287 Draft Tubes 287 Impeller Types 287 Impeller Size 287 Impeller Speed 288 Impeller Location 288 10.2. Kinds of Impellers 288 10.3. Characterization of Mixing Quality 290 10.4. Power Consumption and Pumping Rate 292 10.5. Suspension of Solids 295 10.6. Gas Dispersion 296 Spargers 296 Mass Transfer 297 System Design 297 Minimum Power 297 Power Consumption of Gassed Liquids 297 Superficial Liquid Velocity 297 Design Procedures 297 10.7. In-Line-Blenders and Mixers 300 10.8. Mixing of Powders and Pastes 301 References 304

CHAPTER 11 SOLID-LIQUID SEPARATION 305

- 11.1. Processes and Equipment 305 11.2 Theory of Filtration 306
- Compressible Cakes 310 11.3. Resistance to Filtration 313 Filter Medium 313 Cake Resistivity 313

CONTENTS Vii

Compressibility-Permeability (CP) Cell Measurements 314 Another Form of Pressure Dependence 315 Pretreatment of Slurries 315

- 11.4. Thickening and Clarifying 31511.5. Laboratory Testing and Scale-Up 317 Compression-Permeability Cell 317 The SCFT Concept 317 Scale-Up 318
- 11.6. Illustrations of Equipment 318
- 11.7. Applications and Performance of Equipment 320 References 334
- CHAPTER 12 DISINTEGRATION,
- AGGLOMERATION, AND SIZE SEPARATION OF PARTICULATE SOLIDS 335
- 12.1. Screening 335 Revolving Screens or Trommels 335 Capacity of Screens 335 12.2. Classification with Streams of Air or Water 337
- Air Classifiers 337 Wet Classifiers 339
- 12.3. Size Reduction 339
- 12.4. Eauiument for Size Reduction 341 Crushers 341 Roll Crushers 341
- 12.5. Particle Size Enlargement 351 Tumblers 351 Roll Compacting and Briquetting 354 Tabletting 357 **Extrusion Processes 358** Prilling 361 Fluidized and Spouted Beds Sintering and Crushing 363 362 References 370

CHAPTER 13 DISTILLATION AND GAS ABSORPTION 371

- 13.1. Vapor-Liquid Equilibria 371 Relative Volatility 374 Binary x-y Diagrams 375
- Single-Stage Flash Calculations 375 Bubblepoint Temperature and Pressure 376 13.2 Dewpoint Temperature and Pressure 377 Flash at Fixed Temnerature and Pressure 377 Flash at Fixed Enthalpy and Pressure 377 Equilibria with Ks Dependent on Composition 377
- 13.3. Evaporation or Simple Distillation 378 Multicomponent Mixtures 379
- 13.4. Binary Distillation 379 Material and Energy Balances 380 Constant Molal Overflow 380 **Basic Distillation Problem 382** Unequal Molal Heats of Vaporization 382 Material and Energy Balance Basis 382 Algebraic Method 382
- 13.5. Batch Distillation 390 Material Balances 391 13. 6. Multicomponent Separation: Generali
- Considerations 393 Sequencing of Columns 393
- Number of Free Variables 395 13.7. Estimation of Reflux and Number of Travs (Fenske-Underwood-Gilliland Method) 395 Minimum Trays 395 Distribution of Nonkeys 395 Minimum Reflux 397 Operating Reflux 397 Actual Number of Theoretical Trays 397 Feed Tray Location 397

Tray Efficiencies 397 Absorption Factor Shortcut Method of Edmister 13.8. 398 Seoarations in Packed Towers 398 13.9. Miss Transfer Coefficients 399 Distillation 401 Absorption or Stripping 401 13.10. Basis for Computer Evaluation of Multicomponent Separations 404 Specifications 405 The MESH Equations 405 The Wang-Henke Bubblepoint Method 408 The SR (Sum-Rates) Method 409 SC (Simultaneous Correction) Method 410 13.11. Special Kinds of Distillation Processes 410 Petroleum Fractionation 411 Extractive Distillation 412 Azeotropic Distillation 420 Molecular Distillation 425 13.12. Tray Towers 426 Countercurrent Trays 426 Sieve Trays 428 Valve Trays 429 Bubblecap Trays 431 13.13. Packed Towers 433 Kinds of Packings 433 Flooding and Allowable Loads 433 Liquid Distribution 439 Liauid Holdup 439 Pressure Drop 439 13.14. Efficiencies of Trays and Packings 439 Trays 439 Packed Towers 442 References 456 CHAPTER 14 EXTRACTION AND LEACHING 459 14.1. Equilibrium Relations 459 14.2. Calculation of Stage Requirements 463 Single Staee Extraction 463 Crosscurrent Extraction 464 **Immiscible Solvents 464** 14.3. Countercurrent Operation 466 Minimum Solvent/Feed Ratio 468 Extract Reflux 468 Minimum Reflux 469 Minimum Stages 469 14.4. Leaching of Solids 470 14.5. Numerical Calculation of Multicomponent Extraction 473 Initial Estimates 473 Procedure 473 14.6. Equipment for Extraction' 476 Choice of Disperse Phase 476 Mixer-Settlers' 477 Spray Towers 478 Packed Towers 478 Sieve Tray Towers 483 Pulsed Packed and Sieve Tray Towers 483 Reciprocating Tray Towers 485 Rotating Disk Contactor (RDC) 485 Other Rotary Agitated Towers 485 Other Kinds of Extractors 487 Leaching Equipment 488 References 493 CHAPTER 15 ADSORPTION AND ION

EXCHANGE 495

- 15.1. Adsorption Equilibria 495
- 15.2. Ion Exchange Equilibria 497
- 15.3. Adsorption Behavior in Packed Beds 500 Regeneration 504

Viii CONTENTS

- 15.4. Adsorption Design and Operating Practices 504
- 15.5. Ion Exchange Design and Operating Practices 506 Electrodialysis 508
- 15.6. Production Scale Chromatography 51015.7. Equipment and Processes 510
- Gas Adsorption 511 Liquid Phase Adsorption 513 Ion Exchange 517 Ion Exchange Membranes and Electrodialysis 517 Chromatographic Equipment 520 References 522

CHAPTER 16 CRYSTALLIZATION FROM SOLUTIONS AND MELTS 523

- 16.1. Solubilities and Equilibria 523 Phase Diagrams 523 Enthalpy Balances 524
- 16.2. Crvstal Size Distribution 525
- 16.3. The Process of Crystallization 528 Conditions of Precipitation 528 Supersaturation 528 Growth Rates 530
- 16.4. The Ideal Stirred Tank 533 Multiple Stirred Tanks in Series 536 Applicability of the CSTC Model 536 16.5. Kinds of Crystallizers 537 16.6. Melt Crystallization and Purification 543
- Multistage Processing 543 The Metallwerk Buchs Process 543 Purification Processes 543 References 548

CHAPTER 17 CHEMICAL REACTORS 549

- 17.1. Design Basis and Space Velocity 549 Design Basis 549 Reaction Times 549
- 17.2. Rate Equations and Operating Modes 549
- 17.2. Rate Equations and Operating Process 545
 17.3. Material and Energy Balances of Reactors 555
 17.4. Nonideal Flow Patterns 556
- Residence Time Distribution 556 Conversion in Segregated and Maximum Mixed Flows 560 Conversion in Segregated Flow and CSTR Batteries 560 **Dispersion Model 560** Laminar and Related Flow Patterns 561 17.5. Selection of Catalysts 562 Heterogeneous Catalysts 562 Kinds of Catalysts 563 Kinds of Catalvzed Organic Reactions 563 Physical Characteristics of Solid Catalysts 564 Catalyst Effectiveness 565
- 17.6. Types and Examples of Reactors 567 Stirred Tanks 567 **Tubular Flow Reactors 569** Gas-Liquid Reactions 571 Fixed Bed Reactors 572 Moving Beds 574 Kilns and Hearth Furnaces 575 Fluidized Bed Reactors 579 17.7. Heat Transfer in Reactors 582
- Stirred Tanks 586 Packed Bed Thermal Conductivity 587 Heat Transfer Coefficient at Walls, to Particles, and Overall 587 Fluidized Beds 589
- 17.8. Classes of Reaction Processes and Their Equipment 592 Homogeneous Gas Reactions 592

Homogeneous Liquid Reactions 595 Liquid-Liquid Reactions 595 Gas-Liquid Reactions 595 Noncatalytic Reactions with Solids 595 Fluidized Beds of Noncatalytic Solids 595 Circulating Gas or Solids 596 Fixed Bed Solid Catalysis 596 Fluidized Bed Catalysis 601 Gas-Liquid Reactions with Solid Catalysts 604 References 609

CHAPTER 18 PROCESS VESSELS 611

- 18.1. Drums 611
- 18.2. Fractionator Reflux Drums 612
 18.3. Liquid-Liquid Separators 612 Coalescence 613
- Other Methods 613 18.4. Gas-Liquid Separators 613
- Droplet Sizes 613 Rate of Settling 614 Empty Drums 615
 - Wire Mesh Pad Deentrainers 615

- 18.5. Cyclone Separators 616 18.6. Storage Tanks 619 18.7. Mechanical Design of Process Vessels 621 Design Pressure and Temperature 623 Shells and Heads 624 Formulas for Strength Calculations 624 References 629

CHAPTER 19 OTHER TOPICS 631

- 19.1. Membrane Processes 631 Membranes 632 Equipment Configurations 632 Applications 632 Gas Permeation 633
- 19.2. Foam Separation and Froth Flotation 635 Foam Fractionation 635 Froth Flotation 636
- 19.3. Sublimation and Freeze Drying 638 Equipment 639 Freeze Drying 639
- 19.4. Parametric Pumping 639
- 19.5. Seoarations by Thermal Diffusion 642 19.6. Electrochemical Syntheses 645 Electrochemical Reactions 646 Fuel Cells 646 Cells for Synthesis of Chemicals 648
- 19.7. Fermentation Processing 648 Processing 650 Operating Conditions 650 Reactors 654 References 660

CHAPTER 20 COSTS OF INDIVIDUAL **EQUIPMENT 663**

References 669

APPENDIX A UNITS, NOTATION, AND GENERAL DATA 671

APPENDIX B EQUIPMENT SPECIFICATION FORMS 681

APPENDIX C QUESTIONNAIRES OF EQUIPMENT SUPPLIERS 727

INDEX 747

-List of Examples -

- 1.1 Material Balance of a Chlorination Process with Recycle 5
- 1.2 Data of a Steam Generator for Making 250,000 lb/hr at 450 psia and 650°F from Water Entering at 220°F 9 Steam Plant Cycle for Generation of Power and Low
- 1.3 Pressure Process Steam 11
- Pickup of Waste Heat by Generating and Superheating 1.4 Steam in a Petroleum Refinery 11
- Recovery of Power from a Hot Gas Stream 12 1.5
- 3.1 Constants of PID Controllers from Response Curves to a Step Input 42
- Steam Requirement of a Turbine Operation 65 4.1
- 4.2 Performance of a Combustion Gas Turbine 67
- Conditions of a Coal Slurry Pipeline 70 5.1
- Size and Power Requirement of a Pneumatic Transfer 5.2 Line 77
- Sizing a Screw Conveyor 80 5.3
- 5.4 Sizing a Belt Conveyor 83
- Comparison of Redler and Zippered Belt Conveyors 88
- Density of a Nonideal Gas from Its Equation of State 91 6.1
- Unsteady Flow of an Ideal Gas through a Vessel 93 6.2
- Units of the Energy Balance 94 6.3
- 6.4 Pressure Drop in Nonisothermal Liquid Flow 97
- Comparison of Pressure Drons in a Line with Several Sets of 6.5 Fittings Resistances 101
- A Network of Pipelines in Series, Parallel, and Branches: 6.6 the Sketch, Material Balances, and Pressure Drop Equations 101
- Flow of Oil in a Branched Pipeline 101 6.7
- Economic Optimum Pine Size for Pumping Hot Oil with a 6.8 Motor or Turbine Drive 102
- 6.9 Analysis of Data Obtained in a Capillary Tube Viscometer 107
- Parameters of the Bingham Model from Measurements of 6.10 Pressure Drops in a Line 107
- Pressure Drop in Power-Law and Bingham Flow 110 6.11
- Adiabatic and Isothermal Flow of a Gas in a Pipeline 6.12 112
- 6.13 Isothermal Flow of a Nonideal Gas 113
- Pressure Drop and Void Fraction in Liquid-Gas Flow 6.14 116
- Pressure Drp in Flow of Nitrogen and Powdered 6.15 Coal 120
- Dimensions of a Fluidized Bed Vessel 125 6.16
- 7.1 Application of Dimensionless Performance Curves 132
- Operating Points of Single and Double Pumps in Parallel 7.2 and Series 133
- Check of Some Performance Curves with the Concept of 7.3 Specific Speed 136
- Gas Compression, Isentropic and True Final 7.4 Temperatures 155
- 7.5 Compression Work with Variable Heat Capacity 157
- 7.6 Polytropic and Isentropic Efficiencies 158
- Finding Work of Compression with a Thermodynamic 7.7 Chart 160
- Compression Work on a Nonideal Gas 160 7.8
- Selection of a Centrifugal Compressor 161 7.9
- Polytropic and Isentropic Temperatures 162 7.10
- 7.11 Three-Stage Compression with Intercooling and Pressure Loss between Stages 164
- Equivalent Air Rate 165 7.12
- Interstage Condensers 166 7.13
- 8.1 Conduction Through a Furnace Wall I70
- Effect of Ignoring the Radius Correction of the Overall 8.2 Heat Transfer Coefficient 171
- 8.3 A Case of a Composite Wall: Optimum Insulation Thickness for a Steam Line 171
- Performance of a Heat Exchanger with the F-Method 180 8.4

- Application of the Effectiveness and the θ Method 182 8.5
- Sizing an Exchanger with Radial Finned Tubes 193 8.6
- Pressure Drop on the Tube Side of a Vertical Thermosiphon 8.7 Reboiler 193
- Pressure Drop on the Shell Side with 25% Open Segmental 8.8 Baffles by Kern's Method 194
- Estimation of the Surface Requirements of an Air 8.9 Cooler 199
- Process Design of a Shell-and-Tube Heat Exchanger 204 8.10
- Sizing a Condenser for a Mixture by the Silver-Bell-Ghatly 8.11 Method 207
- 8.12 Comparison of Three Kinds of Reboilers for the Same Service 209
- Peak Temperatures 214 8.13
- Effect of Stock Temperature Variation 214 8.14
- Design of a Fired Heater 217 8.15
- Annlication of the Wilson-Lobo-Hottel eauation 219 8.16
- 8.17 Two-Stages Propylene Compression Refrigeration with Interstage Recycle 225
- Conditions in an Adiabatic Dryer 234 9.1
- Drying Time over Constant and Falling Rate Periods with 9.2 Constant Gas Conditions 237
- Drying with Changing Humidity of Air in a Tunnel 9.3 Dryer 238
- Effects of Moist Air Recycle and Increase of Fresh Air Rate 9.4 in Belt Conveyor Drying 239
- 9.5
- Scale-Up of a Rotary Dryer 256 Design Details of a Countercurrent Rotary Dryer 256 9.6
- Description of a Drum Drying System 260 9.1
- Sizing a Pneumatic Conveying Dryer 266 9.8
- 9.9 Sizing a Fluidized Bed Dryer 272
- 9.10 Sizing a Spray Dryer on the Basis of Pilot Plant Data 279
- Sizine of a Cooling Tower: Number of Transfer Units and 9.11 Height of Packing- 281
- 10.1 Impeller Size and Speed at a Specified Power Input 293
- Effects of the Ratios of impeller and Tank Diameters 294 10.2
- 10.3 Design of the Agitation System for Maintenance of a Slurry 299
- HP and rpm Requirements of an Aerated Agitated 10.4 Tank 301
- Constants of the Filtration Equation from Test Data 310 11.1
- 11.2 Filtration Process with a Centrifugal Charge Pump 311
- Rotary Vacuum Filter Operation 312 11.3
- Filtration and Washing of a Compressible Material 314 11.4
- 12.1 Sizing a Hydrocyclone 341
- Power Requirement for Grinding 342 12.2
- 13.1
- Correlation of Relative Volatility 375 Vanorization and Condensation of a Ternarv Mixture 378 13.2
- Temperature with the Virial add Wilson 13.3 Bubblepoint Equations 379
- Batch Distillation of Chlorinated Phenols 383 13.4
- Distillation of Substances with Widely Different Molal 13.5 Heats of Vaporization 385
- 13.6 Separation of an Azeotropic Mixture by Operation at Two Pressure Levels 387
- Separation of a Partially Miscible Mixture 388 13.7
- Enthalpy-Concentration Lines of Saturated Vapor and 13.8 Liquid of Mixtures of Methanol and Water at a Pressure of 2 aim 390
- Algebraic Method for Binary Distillation Calculation 392 13.9
- Shorcut Design of Multicomponent Fractionation 396 13.10
- Calculation of an Absorber by the Absorption Factor 13.11 Method 399
- 13.12 Numbers of Theoretical Trays and of Transfer Units with Two Values of k_L/k_G for a Distillation Process 402

X LIST OF EXAMPLES

14.1

14.2

14.4

14.5

14.6

14.7

14.8

- 13.13 Trays and Transfer Units for an Absorption Process 403 13.14 Representation of a Petroleum Fraction by an Equivalent Number of Discrete Components 413 13.15 Comparison of Diameters of Sieve, Valve, and Bubblecap Travs for the Same Service 431 13.16 Performance of a Packed Tower by Three Methods 441 13.17 Tray Efficiency for the Separation of Acetone and Benzene 451 The Equations for Tieline Data 465 Tabulated Tieline and Distribution Data for the System A = I-Hexene, B = Tetramethylene Sulfone, C = Benzene, Represented in Figure 14.1 466 Single Stage and Cross Current Extraction of Acetic Acid 14.3 from Methylisobutyl Ketone with Water 468 Extraction with an Immiscible Solvent 469 Countercurrent Extraction Represented on Triangular and Rectangular Distribution Diagrams 470 Stage Requirements for the Separation of a Type I and a Type II System 471 Countercurrent Extraction Employing Extract Reflux 472 Leaching of an Oil-Bearing Solid in a Countercurrent Battery 472
- Trial Estimates and Converged Flow Rates and 14.9 Compositions in all Stages of an Extraction Battery for a Mixture 476 Four-Component
- 14.10 Sizing of Spray, Packed, or Sieve Tray Towers 486
- Design of a Rotating Disk Contactor 488 14.11
- 15.1 Application of Ion Exchange Selectivity Data 503

- 15.2 Adsorption of n-hexane from a Natural Gas with Silica Gel 505
- Size of an Ion Exchanger for Hard Water 513 15.3
- 16.1 Design of a Crystallizing Plant 524
- Using the Phase Diagrams of Figure 16.2 528 16.2
- Heat Effect Accompanying the Cooling of a Solution of 16.3 MgSO₄ 529
- Deductions from a Differential Distribution Obtained at a 16.4 Known Residence Time 533
- Batch Crystallization with Seeded Liquor 534 16.5
- Analysis of Size Distribution Data Obtained in a 16.6 CSTC 537
- Crystallization in a Continuous Stirred Tank with Specified 16.7 Predominant Crystal Size 538
- Crystallization from a Ternary Mixture 544 16.8
- Separation of Oil and Water . 614 18.1
- Quantity of Entrainment on the Basis of Sieve Tray 18.2 Correlations 617
- Liquid Knockout Drum (Empty) 618 18.3
- Knockout Drum with Wire Mesh Deentrainer 620 18.4
- Size and Capacity of Cyclone Separators 621 18.5
- Dimensions and Weight of a Horizontal Pressure 18.6 Drum 628
- Applications of the Equation for Osmotic Pressure 633 19.1
- 19.2 Concentration of a Water/Ethanol Mixture by Reverse Osmosis 642
- Installed Cost of a Distillation Tower 663 20.1
- Purchased and Installed Cost of Some Equipment 663 20.2

Preface-

This book is intended as a guide to the selection or design of the principal kinds of chemical process equipment by engineers in school and industry. The level of treatment assumes an elementary knowledge of unit operations and transport phenomena. Access to the many design and reference books listed in Chapter 1 is desirable. For coherence, brief reviews of pertinent theory are provided. Emphasis is placed on shortcuts, rules of thumb, and data for design by analogy, often as primary design processes but also for quick evaluations of detailed work.

All answers to process design questions cannot be put into a book. Even at this late date in the development of the chemical industry, it is common to hear authorities on most kinds of equipment say that their equipment can be properly fitted to a particular task only on the basis of some direct laboratory and pilot plant work. Nevertheless, much guidance and reassurance are obtainable from general experience and specific examples of successful applications, which this book attempts to provide. Much of the information is supplied in numerous tables and figures, which often deserve careful study quite apart from the text.

The general background of process design, flowsheets, and process control is reviewed in the introductory chapters. The major kinds of operations and equipment are treated in individual chapters. Information about peripheral and less widely employed equipment in chemical plants is concentrated in Chapter 19 with references to key works of as much practical value as possible. Because decisions often must be based on economic grounds, Chapter 20, on costs of equipment, rounds out the book. Appendixes provide examples of equipment rating forms and manufacturers' questionnaires.

Chemical process equipment is of two kinds: custom designed and built, or proprietary "off the shelf." For example, the sizes and performance of custom equipment such as distillation towers, drums, and heat exchangers are derived by the process engineer on the basis of established principles and data, although some mechanical details remain in accordance with safe practice codes and individual fabrication practices.

Much proprietary equipment (such as filters, mixers, conveyors, and so on) has been developed largely without benefit of much theory and is fitted to job requirements also without benefit of much theory. From the point of view of the process engineer, such equipment is predesigned and fabricated and made available by manufacturers in limited numbers of types, sizes, and capacities. The process design of proprietary equipment, as considered in this book, establishes its required performance and is a process of selection from the manufacturers' offerings, often with their recommendations or on the basis of individual experience. Complete information is provided in manufacturers' catalogs. Several classified lists of manufacturers of chemical process equipment are readily accessible, so no listings are given here. Because more than one kind of equipment often is suitable for particular applications and may be available from several manufacturers, comparisons of equipment and typical applications are cited liberally. Some features of industrial equipment are largely arbitrary and may be standardized for convenience in particular industries or individual plants. Such aspects of equipment design are noted when feasible.

Shortcut methods of design provide solutions to problems in a short time and at small expense. They must be used when data are limited or when the greater expense of a thorough method is not justifiable. In particular cases they may be employed to obtain information such as:

- 1. an order of magnitude check of the reasonableness of a result found by another lengthier and presumably accurate computation or computer run,
- 2. a quick check to find if existing equipment possibly can be adapted to a new situation,
- 3. a comparison of alternate processes,
- 4. a basis for a rough cost estimate of a process.

Shortcut methods occupy a prominent place in such a broad survey and limited space as this book. References to sources of more accurate design procedures are cited when available.

Another approach to engineering work is with rules of thumb, which are statements of equipment performance that may obviate all need for further calculations. Typical examples, for instance, are that optimum reflux ratio is 20% greater than minimum, that a suitable cold oil velocity in a fired heater is 6ft/sec, or that the efficiency of a mixer-settler extraction stage is 70%. The trust that can be placed in a rule of thumb depends on the authority of the propounder, the risk associated with its possible inaccuracy, and the economic balance between the cost of a more accurate evaluation and suitable safety factor placed on the approximation. All experienced engineers have acquired such knowledge. When applied with discrimination, rules of thumb are a valuable asset to the process design and operating engineer, and are scattered throughout this book.

Design by analogy, which is based on knowledge of what has been found to work in similar areas, even though not necessarily optimally, is another valuable technique. Accordingly, specific applications often are described in this book, and many examples of specific equipment sizes and performance are cited.

For much of my insight into chemical process design, I am indebted to many years' association and friendship with the late Charles W. Nofsinger who was a prime practitioner by analogy, rule of thumb, and basic principles. Like Dr. Dolittle of Puddleby-onthe-Marsh, "he was a proper doctor and knew a whole lot."

RULES OF THUMB: SUMMARY

Although experienced engineers know where to find information and how to make accurate computations, they also keep a minimum body of information in mind on the ready, made up largely of shortcuts and rules of thumb. The present compilation may fit into such a minimum body of information, as a boost to the memory or extension in some instances into less often encountered areas. It is derived from the material in this book and is, in a sense, a digest of the book.

An Engineering Rule of Thumb is an outright statement regarding suitable sizes or performance of equipment that obviates all need for extended calculations. Because any brief statements are subject to varying degrees of qualification, they are most safely applied by engineers who are substantially familiar with the topics. Nevertheless, such rules should be of value for approximate design and cost estimation, and should provide even the inexperienced engineer with perspective and a foundation whereby the reasonableness of detailed and computer-aided results can be appraised quickly, particularly on short notice such as in conference.

Everyday activities also are governed to a large extent by rules of thumb. They serve us when we wish to take a course of action but are not in a position to find the best course of action. Of interest along this line is an amusing and often useful list of some 900 such digests of everyday experience that has been compiled by Parker (Rules of Thumb, Houghton Mifflin, Boston, 1983).

Much more can be stated in adequate summary fashion about some topics than about others, which accounts in part for the spottiness of the present coverage, but the spottiness also is due to ignorance and oversights on the part of the author. Accordingly, every engineer undoubtedly will supplement or modify this material in his own way.

COMPRESSORS AND VACUUM PUMPS

- 1. *Fans* are used to raise the pressure about 3% (12in. water), blowers raise to less than 40 psig, and compressors to higher pressures, although the blower range commonly is included in the compressor range.
- Vacuum pumps: reciprocating piston type decrease the pressure to 1 Torr; rotary piston down to 0.001 Torr, two-lobe rotary down to 0.0001 Torr; steam jet ejectors, one stage down to lOOTorr, three stage down to 1 Torr, five stage down to 0.05 Torr.
- 3. A three-stage ejector needs 100lb steam/lb air to maintain a pressure of 1 Torr.
- 4. In-leakage of air to evacuated equipment depends on the absolute pressure, Torr, and the volume of the equipment, V cuft, according to $w = kV^{2/3}$ lb/hr, with k = 0.2 when P is more than 90 Torr, 0.08 between 3 and 20 Torr, and 0.025 at less than 1 Torr.
- 5. Theoretical adiabatic horsepower (THP) = [(SCFM)T1/8130a]
- $[(P_2/P_1)^a 1]$, where T_1 is inlet temperature in °F+ 460 and a = (k 1)/k, $k = C_p/C_v$.
- 6. Outlet temperature $T_2 = T_1(P_2/P_1)^a$.
- 7. To compress air from 100°F, k = 1.4, compression ratio = 3, theoretical power required = 62 HP/million cuft/day, outlet temperature 306°F.
- 8. Exit temperature should not exceed 350-400°F; for diatomic gases ($C_p/C_v = 1.4$) this corresponds to a compression ratio of about 4.

- 9. Compression ratio should be about the same in each stage of a multistage unit, ratio $= (P_n/P_1)^{1/n}$, with *n* stages.
- 10. Efficiencies of reciprocating compressors: 65% at compression ratio of 1.5, 75% at 2.0, and 80–85% at 3-6.
- 11. Efficiencies of large centrifugal compressors, 6000-100,000 ACFM at suction, are 76-78%.
- 12. Rotary compressors have efficiencies of 70%, except liquid liner type which have 50%.

CONVEYORS FOR PARTICULATE SOLIDS

- Screw conveyors are suited to transport of even sticky and abrasive solids up inclines of 20° or so. They are limited to distances of 150ft or so because of shaft torque strength. A 12in. dia conveyor can handle 1000–3000 cuft/hr, at speeds ranging from 40 to 60 rpm.
- Belt conveyors are for high capacity and long distances (a mile or more, but only several hundred feet in a plant), up inclines of 30° maximum. A 24in. wide belt can carry 3000 cuft/hr at a speed of 100 ft/min, but speeds up to 600 ft/min are suited to some materials. Power consumption is relatively low.
- **3.** *Bucker elevators* are suited to vertical transport of sticky and abrasive materials. With buckets 20 x 20 in. capacity can reach 1000 cuft/hr at a speed of 100 ft/min, but speeds to 300 ft/min are used.
- **4.** *Drug-type conveyors* (Redler) are suited to short distances in any direction and are completely enclosed. Units range in size from 3 in. square to 19 in. square and may travel from 30 ft/min (fly ash) to 250 ft/min (grains). Power requirements are high.
- **5.** *Pneumatic conveyors* are for high capacity, short distance (400 ft) transport simultaneously from several sources to several destinations. Either vacuum or low pressure (6-12psig) is employed with a range of air velocities from 35 to 120 ft/sec depending on the material and pressure, air requirements from 1 to 7 cuft/cuft of solid transferred.

COOLING TOWERS

- 1. Water in contact with air under adiabatic conditions eventually cools to the wet bulb temperature.
- 2. In commercial units, 90% of saturation of the air is feasible.
- 3. Relative cooling tower size is sensitive to the difference between the exit and wet bulb temperatures:

- Tower fill is of a highly open structure so as to minimize pressure drop, which is in standard practice a maximum of 2 in. of water.
- Water circulation rate is 1-4 gpm/sqft and air rates are 1300-1800 lb/(hr)(sqft) or 300-400 ft/min.
- 6. Chimney-assisted natural draft towers are of hyperboloidal shapes because they have greater strength for a given thickness; a tower 250 ft high has concrete walls 5-6 in. thick. The enlarged cross section at the top aids in dispersion of exit humid air into the atmosphere.
- 7. Countercurrent induced draft towers are the most common in process industries. They are able to cool water within 2°F of the wet bulb.
- Evaporation losses are 1% of the circulation for every 10°F of cooling range. Windage or drift losses of mechanical draft towers

are O.1-0.3%. Blowdown of 2.5-3.0% of the circulation is necessary to prevent excessive salt buildup.

CRYSTALLIZATION FROM SOLUTION

- 1. Complete recovery of dissolved solids is obtainable by evaporation, but only to the eutectic composition by chilling. Recovery by melt crystallization also is limited by the eutectic composition.
- 2. Growth rates and ultimate sizes of crystals are controlled by limiting the extent of supersaturation at any time.
- 3. The ratio $S = C/C_{,,,}$ of prevailing concentration to saturation concentration is kept near the range of 1.02-1.05.
- In crystallization by chilling, the temperature of the solution is kept at most 1–2°F below the saturation temperature at the prevailing concentration.
- Growth rates of crystals under satisfactory conditions are in the range of 0.1-0.8 mm/hr. The growth rates are approximately the same in all directions.
- 6. Growth rates are influenced greatly by the presence of impurities and of certain specific additives that vary from case to case.

DISINTEGRATION

- 1. Percentages of material greater than 50% of the maximum size are about 50% from rolls, 15% from tumbling mills, and 5% from closed circuit ball mills.
- 2. Closed circuit grinding employs external size classification and return of oversize for regrinding. The rules of pneumatic conveying are applied to design of air classifiers. Closed circuit is most common with ball and roller mills.
- Jaw crushers take lumps of several feet in diameter down to 4 in. Stroke rates are 100-300/min. The average feed is subjected to 8-10 strokes before it becomes small enough to escape. Gyratory crushers are suited to slabby feeds and make a more rounded product.
- 4. Roll crushers are made either smooth or with teeth. A 24in. toothed roll can accept lumps 14 in. dia. Smooth rolls effect reduction ratios up to about 4. Speeds are 50-900 rpm. Capacity is about 25% of the maximum corresponding to a continuous ribbon of material passing through the rolls.
- 5. Hammer mills beat the material until it is small enough to pass through the screen at the bottom of the casing. Reduction ratios of 40 are feasible. Large units operate at 900 rpm, smaller ones up to 16,000 rpm. For fibrous materials the screen is provided with cutting edges.
- 6. Rod mills are capable of taking feed as large as 50 mm and reducing it to 300 mesh, but normally the product range is 8-65 mesh. Rods are 25–150 mm dia. Ratio of rod length to mill diameter is about 1.5. About 45% of the mill volume is occupied by rods. Rotation is at 50–65% of critical.
- 7. Ball mills are better suited than rod mills to fine grinding. The charge is of equal weights of 1.5, 2, and 3 in. balls for the finest grinding. Volume occupied by the balls is 50% of the mill volume. Rotation speed is 70–80% of critical. Ball mills have a length to diameter ratio in the range 1-1.5. Tube mills have a ratio of 4-5 and are capable of very fine grinding. Pebble mills have ceramic grinding elements, used when contamination with metal is to be avoided.
- 8. Roller mills employ cylindrical or tapered surfaces that roll along flatter surfaces and crush nipped particles. Products of 20-200 mesh are made.

DISTILLATION AND GAS ABSORPTION

- 1. Distillation usually is the most economical method of separating liquids, superior to extraction, adsorption, crystallization, or others.
- 2. For ideal mixtures, relative volatility is the ratio of vapor pressures $\alpha_{12} = P_2/P_1$.
- 3. Tower operating pressure is determined most often by the temperature of the available condensing medium, 100–120°F if cooling water; or by the maximum allowable reboiler temperature, 150 psig steam, 366°F.
- 4. Sequencing of columns for separating multicomponent mixtures: (a) perform the easiest separation first, that is, the one least demanding of trays and reflux, and leave the most difficult to the last; (b) when neither relative volatility nor feed concentration vary widely, remove the components one by one as overhead products; (c) when the adjacent ordered components in the feed vary widely in relative volatility; (d) when the concentrations in the feed vary widely but the relative volatilities do not, remove the components in the order of decreasing concentration in the feed.
- 5. Economically optimum reflux ratio is about 1.2 times the minimum reflux ratio $R_{..}$
- 6. The economically optimum number of trays is near twice the minimum value N_m .
- 7. The minimum number of trays is found with the Fenske-Underwood equation

$$N_m = \log\{[x/(1-x)]_{\text{ovhd}}/[x/(1-x)]_{\text{btms}}\}/\log \alpha.$$

8. Minimum reflux for binary or pseudobinary mixtures is given by the following when separation is esentially complete $(x_D \approx 1)$ and D/F is the ratio of overhead product and feed rates:

 $R_m D/F = 1/(\alpha - 1)$, when feed is at the bubblepoint, $(R_r + 1)D/F = \alpha/(\alpha - 1)$, when feed is at the dewpoint.

- 9. A safety factor of 10% of the number of trays calculated by the best means is advisable.
- 10. Reflux pumps are made at least 25% oversize.
- 11. For reasons of accessibility, tray spacings are made 20-24 in.
- 12. Peak efficiency of trays is at values of the vapor factor $F_s = u\sqrt{\rho_v}$ in the range 1.0-1.2 (ft/sec) $\sqrt{lb/cuft}$. This range of F_s establishes the diameter of the tower. Roughly, linear velocities are 2 ft/sec at moderate pressures and 6 ft/sec in vacuum.
- 13. The optimum value of the Kremser-Brown absorption factor A = K(V/L) is in the range 1.25-2.0.
- 14. Pressure drop per tray is of the order of 3 in. of water or 0.1 psi.
- Tray efficiencies for distillation of light hydrocarbons and aqueous solutions are 60–90%; for gas absorption and stripping, 10–20%.
- 16. Sieve trays have holes 0.25-0.50 in. dia, hole area being 10% of the active cross section.
- 17. Valve trays have holes 1.5 in. dia each provided with a liftable cap, 12-14 **caps/sqft** of active cross section. Valve trays usually are cheaper than sieve trays.
- Bubblecap trays are used only when a liquid level must be maintained at low turndown ratio; they can be designed for lower pressure drop than either sieve or valve trays.
- 19. Weir heights are 2 in., weir lengths about 75% of tray diameter, liquid rate a maximum of about 8 gpm/in. of weir; multipass arrangements are used at high liquid rates.

- 20. Packings of random and structured character are suited especially to towers under 3 ft dia and where low pressure drop is desirable. With proper initial distribution and periodic redistribution, volumetric efficiencies can be made greater than those of tray towers. Packed internals are used as replacements for achieving greater throughput or separation in existing tower shells.
- 21. For gas rates of 500 cfm, use 1 in. packing; for gas rates of 2000 cfm or more, use 2 in.
- 22. The ratio of diameters of tower and packing should be at least 15.
- 23. Because of deformability, plastic packing is limited to a 10-15 ft depth unsupported, metal to 20-25 ft.
- 24. Liquid redistributors are needed every 5-10 tower diameters with pall rings but at least every 20ft. The number of liquid streams should be 3-5/sqft in towers larger than 3 ft dia (some experts say 9-12/sqft), and more numerous in smaller towers.
- 25. Height equivalent to a theoretical plate (HETP) for vapor-liquid contacting is 1.3–1.8 ft for 1 in. pall rings, 2.5-3.0 ft for 2 in. pall rings.
- **26.** Packed towers should operate near 70% of the flooding rate given by the correlation of Sherwood, Lobo, et al.
- 27. Reflux drums usually are horizontal, with a liquid holdup of 5 min half full. A takeoff pot for a second liquid phase, such as water in hydrocarbon systems, is sized for a linear velocity of that phase of 0.5 ft/sec, minimum diameter of 16 in.
- 28. For towers about 3 ft dia, add 4ft at the top for vapor disengagement and 6 ft at the bottom for liquid level and reboiler return.
- 29. Limit the tower height to about 175 ft max because of wind load and foundation considerations, An additional criterion is that L/D be less than 30.

DRIVERS AND POWER RECOVERY EQUIPMENT

- 1. Efficiency is greater for larger machines. Motors are 85-95%; steam turbines are 42-78%; gas engines and turbines are 28-38%.
- 2. For under 100 HP, electric motors are used almost exclusively. They are made for up to 20,000 HP.
- 3. Induction motors are most popular. Synchronous motors are made for speeds as low as 150rpm and are thus suited for example for low speed reciprocating compressors, but are not made smaller than 50HP. A variety of enclosures is available, from weather-proof to explosion-proof.
- 4. Steam turbines are competitive above 100 HP. They are speed controllable. Frequently they are employed as spares in case of power failure.
- 5. Combustion engines and turbines are restricted to mobile and remote locations.
- 6. Gas expanders for power recovery may be justified at capacities of several hundred HP; otherwise any needed pressure reduction in process is effected with throttling valves.

DRYING OF SOLIDS

- Drying times range from a few seconds in spray dryers to 1 hr or less in rotary dryers and up to several hours or even several days in tunnel shelf or belt dryers.
- 2. Continuous tray and belt dryers for granular material of natural size or pelleted to 3-15 mm have drying times in the range of 10-200 min.
- Rotary cylindrical dryers operate with superficial air velocities of 5-10 ft/sec, sometimes up to 35 ft/sec when the material is coarse. Residence times are 5-90 min. Holdup of solid is 7-8%.

An 85% free cross section is taken for design purposes. In countercurrent flow, the exit gas is $10-20^{\circ}$ C above the solid; in parallel flow, the temperature of the exit solid is 100° C. Rotation speeds of about 4rpm are used, but the product of rpm and diameter in feet is typically between 15 and 25.

- 4. Drum dryers for pastes and slurries operate with contact times of 3-12 sec, produce flakes 1-3 mm thick with evaporation rates of 15-30 kg/m² hr. Diameters are 1.5-5.0 ft; the rotation rate is 2-10 rpm. The greatest evaporative capacity is of the order of 3000 lb/hr in commercial units.
- 5. Pneumatic conveying dryers normally take particles I-3 mm dia but up to 10 mm when the moisture is mostly on the surface. Air velocities are 10–30 m/sec. Single pass residence times are 0.5-3.0 sec but with normal recycling the average residence time is brought up to 60 sec. Units in use range from 0.2 m dia by 1 m high to 0.3 m dia by 38 m long. Air requirement is several SCFM/lb of dry product/hr.
- 6. Fluidized bed dryers work best on particles of a few tenths of a mm dia, but up to 4 mm dia have been processed. Gas velocities of twice the minimum fluidization velocity are a safe prescription. In continuous operation, drying times of 1-2 min are enough, but batch drying of some pharmaceutical products employs drying times of 2-3 hr.
- 7. Spray dryers: Surface moisture is removed in about 5 sec, and most drying is completed in less than 60 sec. Parallel flow of air and stock is most common. Atomizing nozzles have openings 0.012-0.15 in. and operate at pressures of 300-4000 psi. Atomizing spray wheels rotate at speeds to 20,000 rpm with peripheral speeds of 250-600 ft/sec. With nozzles, the length to diameter ratio of the dryer is 4-5; with spray wheels, the ratio is 0.5-1.0. For the final design, the experts say, pilot tests in a unit of 2 m dia should be made.

EVAPORATORS

- 1. Long tube vertical evaporators with either natural or forced circulation are most popular. Tubes are 19-63 mm dia and 12-30 ft long.
- 2. In forced circulation, linear velocities in the tubes are 15-20 ft/sec.
- 3. Elevation of boiling point by dissolved solids results in differences of $3-10^{\circ}F$ between solution and saturated vapor.
- 4. When the boiling point rise is appreciable, the economic number of effects in series with forward feed is 4-6.
- 5. When the boiling point rise is small, minimum cost is obtained with 8-10 effects in series.
- 6. In backward feed the more concentrated solution is heated with the highest temperature steam so that heating surface is lessened, but the solution must be pumped between stages.
- 7. The steam economy of an N-stage battery is approximately 0.8*N* lb evaporation/lb of outside steam.
- 8. Interstage steam pressures can be boosted with steam jet compressors of 20-30% efficiency or with mechanical compressors of 70-75% efficiency.

EXTRACTION, LIQUID-LIQUID

1. The dispersed phase should be the one that has the higher volumetric rate except in equipment subject to backmixing where it should be the one with the smaller volumetric rate. It should be the phase that wets the material of construction less well. Since the holdup of continuous phase usually is greater, that phase should be made up of the less expensive or less hazardous material.

- 2. There are no known commercial applications of reflux to extraction processes, although the theory is favorable (Treybal).
- 3. Mixer-settler arrangements are limited to at most five stages. Mixing is accomplished with rotating impellers or circulating pumps. Settlers are designed on the assumption that droplet sizes are about 150 μ m dia. In open vessels, residence times of 30-60 min or superficial velocities of 0.5–1.5 ft/min are provided in settlers. Extraction stage efficiencies commonly are taken as 80%.
- 4. Spray towers even 20–40 ft high cannot be depended on to function as more than a single stage.
- 5. Packed towers are employed when 5-10 stages suffice. Pall rings of 1-1.5 in. size are best. Dispersed phase loadings should not exceed 25 gal/(min) (sqft). HETS of 5-10 ft may be realizable. The dispersed phase must be redistributed every 5-7 ft. Packed towers are not satisfactory when the surface tension is more than 10 dyn/cm.
- 6. Sieve tray towers have holes of only 3-8 mm dia. Velocities through the holes are kept below 0.8 ft/sec to avoid formation of small drops. Redispersion of either phase at each tray can be designed for. Tray spacings are 6-24 in. Tray efficiencies are in the range of 20–30%.
- Pulsed packed and sieve tray towers may operate at frequencies of 90 cycles/min and amplitudes of 6-25 mm. In large diameter towers, HETS of about 1 m has been observed. Surface tensions as high as 30-40 dyn/cm have no adverse effect.
- 8. Reciprocating tray towers can have holes 9/16 in. dia, 50–60% open area, stroke length 0.75 in., 100-150 strokes/mitt, plate spacing normally 2 in. but in the range 1-6 in. In a 30in. dia tower, HETS is 20-25 in. and throughput is 2000 gal/(hr)(sqft). Power requirements are much less than of pulsed towers.
- 9. Rotating disk contactors or other rotary agitated towers realize HETS in the range 0.1-0.5 m. The especially efficient Kuhni with perforated disks of 40% free cross section has HETS 0.2 m and a capacity of 50 m^3/m^2 hr.

FILTRATION

- 1. Processes are classified by their rate of cake buildup in a laboratory vacuum leaf filter: rapid, 0.1-10.0 cm/sec; medium, 0.1-10.0 cm/min; slow, 0.1-10.0 cm/hr.
- 2. Continuous filtration should not be attempted if 1/8 in. cake thickness cannot be formed in less than 5 min.
- 3. Rapid filtering is accomplished with belts, top feed drums, or pusher-type centrifuges.
- 4. Medium rate filtering is accomplished with vacuum drums or disks or peeler-type centrifuges.
- 5. Slow filtering slurries are handled in pressure filters or sedimenting centrifuges.
- 6. Clarification with negligible cake buildup is accomplished with cartridges, precoat drums, or sand filters.
- 7. Laboratory tests are advisable when the filtering surface is expected to be more than a few square meters, when cake washing is critical, when cake drying may be a problem, or when precoating may be needed.
- For finely ground ores and minerals, rotary drum filtration, rates may be 1500 lb/(day)(sqft), at 20 rev/hr and 18-25 in. Hg vacuum.
- 9. Coarse solids and crystals may be filtered at rates of 6000 lb/(day)(sqft) at 20 rev/hr, 2-6 in. Hg vacuum.

FLUIDIZATION OF PARTICLES WITH GASES

1. Properties of particles that are conducive to smooth fluidization include: rounded or smooth shape, enough toughness to resist attrition, sizes in the range $50-500 \,\mu\text{m}$ dia, a spectrum of sizes with ratio of largest to smallest in the range of 10-25.

- 2. Cracking catalysts are members of a broad class characterized by diameters of $30-150 \ \mu m$, density of $1.5 \ g/mL$ or so, appreciable expansion of the bed before fluidization sets in, minimum bubbling velocity greater than minimum fluidizing velocity, and rapid disengagement of bubbles.
- 3. The other extreme of smoothly fluidizing particles is typified by coarse sand and glass beads both of which have been the subject of much laboratory investigation. Their sizes are in the range 150-500 μ m, densities 1.5-4.0 g/mL, small bed expansion, about the same magnitudes of minimum bubbling and minimum fluidizing velocities, and also have rapidly disengaging bubbles.
- 4. Cohesive particles and large particles of 1 mm or more do not **fluidize** well and usually are processed in other ways.
- 5. Rough correlations have been made of minimum fluidization velocity, minimum bubbling velocity, bed expansion, bed level fluctuation, and disengaging height. Experts recommend, however, that any real design be based on pilot plant work.
- 6. Practical operations are conducted at two or more multiples of the minimum fluidizing velocity. In reactors, the entrained material is recovered with cyclones and returned to process. In dryers, the fine particles dry most quickly so the entrained material need not be recycled.

HEAT EXCHANGERS

- 1. Take true countercurrent flow in a shell-and-tube exchanger as a basis.
- Standard tubes are 3/4 in. OD, 1 in. triangular spacing, 16 ft long; a shell 1 ft dia accommodates 100 sqft; 2 ft dia, 400 sqft, 3 ft dia, 1100 sqft.
- 3. Tube side is for corrosive, fouling, scaling, and high pressure fluids.
- 4. Shell side is for viscous and condensing fluids.
- 5. Pressure drops are 1.5 psi for boiling and 3-9 psi for other 'services.
- 6. Minimum temperature approach is 20°F with normal coolants, 10°F or less with refrigerants.
- 7. Water inlet temperature is 90°F, maximum outlet 120°F.
- 8. Heat transfer coefficients for estimating purposes, Btu/(hr)(sqft)(°F): water to liquid, 150; condensers, 150; liquid to liquid, 50; liquid to gas, 5; gas to gas, 5; reboiler, 200. Max flux in reboilers, 10,000 Btu/(hr)(sqft).
- 9. Double-pipe exchanger is competitive at duties requiring 100-200 sqft.
- 10. Compact (plate and fin) exchangers have 350 sqft/cuft, and about 4 times the heat transfer per cuft of shell-and-tube units.
- 11. Plate and frame exchangers are suited to high sanitation services, and are 25–50% cheaper in stainless construction than shell-and-tube units.
- **12.** Air coolers: Tubes are 0.75-1.00 in. OD, total finned surface 15-20 sqft/sqft bare surface, U = 80-100 Btu/(hr)(sqft bare surface)(°F), fan power input 2-5 HP/(MBtu/hr), approach 50°F or more.
- Fired heaters: radiant rate, 12,000 Btu/(hr)(sqft); convection rate, 4000; cold oil tube velocity, 6 ft/sec; approx equal transfers of heat in the two sections; thermal efficiency 70–75%; flue gas temperature 250–350°F above feed inlet; stack gas temperature 650–950°F.

INSULATION

- 1. Up to 650°F, 85% magnesia is most used.
- 2. Up to 1600-1900°F, a mixture of asbestos and diatomaceous earth is used.

- 3. Ceramic refractories at higher temperatures.
- Cyrogenic equipment (-200°F) employs insulants with fine pores in which air is trapped.
- 5. Optimum thickness varies with temperature: 0.5 in. at 200°F, 1.0 in. at 400°F, 1.25 in. at 600°F.
- 6. Under windy conditions (7.5 miles/hr), 10-20% greater thickness of insulation is justified.

MIXING AND AGITATION

- 1. Mild agitation is obtained by circulating the liquid with an impeller at superficial velocities of 0.1–0.2 ft/sec, and intense agitation at 0.7-1.0 ft/sec.
- Intensities of agitation with impellers in baffled tanks are measured by power input, HP/1000 gal, and impeller tip speeds:

Operation	HP/1000 gal	Tip speed (ft/min)
Blending	0.2-0.5	
Homogeneous reaction	0.5-1.5	7.5-10
Reaction with heat transfer	1.5-5.0	10-15
Liquid-liquid mixtures	5	15-20
Liquid-gas mixtures	5-10	15-20
Slurries	10	

- 3. Proportions of a stirred tank relative to the diameter D: liquid level = D_i turbine impeller diameter = $D/3_i$ impeller level above bottom = $D/3_i$ impeller blade width = D/15; four vertical baffles with width = D/10.
- 4. Propellers are made a maximum of 18 in., turbine impellers to 9 ft.
- 5. Gas bubbles sparged at the bottom of the vessel will result in mild agitation at a superficial gas velocity of 1 ft/min, severe agitation at 4 ft/min.
- 6. Suspension of solids with a settling velocity of 0.03 ft/sec is accomplished with either turbine or propeller impellers, but when the settling velocity is above 0.15 ft/sec intense agitation with a propeller is needed.
- **7.** Power to drive a mixture of a gas and a liquid can be 25-50% less than the power to drive the liquid alone.
- In-line blenders are adequate when a second or two contact time is sufficient, with power inputs of 0.1–0.2 HP/gal.

PARTICLE SIZE ENLARGEMENT

- The chief methods of particle size enlargement are: compression into a mold, extrusion through a die followed by cutting or breaking to size, globulation of molten material followed by solidification, agglomeration under tumbling or otherwise agitated conditions with or without binding agents.
- 2. Rotating drum granulators have length to diameter ratios of 2-3, speeds of 10–20 rpm, pitch as much as 10". Size is controlled by speed, residence time, and amount of binder; 2-5 mm dia is common.
- 3. Rotary disk granulators produce a more nearly uniform product than drum granulators. Fertilizer is made 1.5-3.5 mm; iron ore 10–25 mm dia.
- 4. Roll compacting and briquetting is done with rolls ranging from 130mm dia by 50mm wide to 910mm dia by 550mm wide. Extrudates are made l-10 mm thick and are broken down to size for any needed processing such as feed to tabletting machines or to dryers.
- 5. Tablets are made in rotary compression machines that convert powders and granules into uniform sizes. Usual maximum diameter is about 1.5 in., but special sizes up to 4in. dia are possible. Machines operate at 100 rpm or so and make up to 10,000 tablets/min.
- 6. Extruders make pellets by forcing powders, pastes, and melts

through a die followed by cutting. An 8 in. screw has a capacity of 2000 lb/hr of molten plastic and is able to extrude tubing at 150–300 ft/min and to cut it into sizes as small as washers at 8000/min. Ring pellet extrusion mills have hole diameters of 1.6-32 mm. Production rates cover a range of 30-200 lb/(hr)(HP).

- **7.** Prilling towers convert molten materials into droplets and allow them to solidify in contact with an air stream. Towers as high as 60m are used. Economically the process becomes competitive with other granulation processes when a capacity of 200-409 tons/day is reached. Ammonium nitrate prills, for example, are 1.6-3.5 mm dia in the 5-95% range.
- 8. Fluidized bed granulation is conducted in shallow beds 12-24 in. deep at air velocities of 0.1-2.5 m/s or 3-10 times the minimum fluidizing velocity, with evaporation rates of $0.005-1.0 \text{ kg/m}^2$ sec. One product has a size range 0.7-2.4 mm dia.

PIPING

- Line velocities and pressure drops, with line diameter D in inches: liquid pump discharge, (5 + D/3) ft/sec, 2.0 psi/100 ft; liquid pump suction, (1.3 + D/6) ft/sec, 0.4 psi/100 ft; steam or gas, 20D ft/sec, 0.5 psi/100 ft.
- 2. Control valves require at least 10 psi drop for good control.
- 3. Globe valves are used for gases, for control and wherever tight shutoff is required. Gate valves are for most other services.
- 4. Screwed fittings are used only on sizes 1.5 in. and smaller, flanges or welding otherwise.
- 5. Flanges and fittings are rated for 150, 300, 600, 900, 1500, or 2500 psig.
- 6. Pipe schedule number = 1000P/S, approximately, where **P** is the internal pressure psig and S is the allowable working stress (about 10,000 psi for A120 carbon steel at 500°F). Schedule 40 is most common.

PUMPS

- 1. Power for pumping liquids: HP = (gpm)(psi difference)/(1714) (fractional efficiency).
- 2. Normal pump suction head (NPSH) of a pump must be in excess of a certain number, depending on the kind of pumps and the conditions, if damage is to be avoided. NPSH = (pressure at the eye of the impeller vapor pressure)/(density). Common range is 4-20 ft.
- 3. Specific speed $N_s = (rpm)(gpm)^{0.5}/(head in ft)^{0.75}$. Pump may be damaged if certain limits of N_s are exceeded, and efficiency is best in some ranges.
- Centrifugal pumps: Single stage for 15-5000 gpm, 500ft max head; multistage for 20-11,000 gpm, 5500 ft max head. Efficiency 45% at 100 gpm, 70% at 500 gpm, 80% at 10,000 gpm.
- 5. Axial pumps for 20-100,000 gpm, 40 ft head, 65-85% efficiency.
- 6. Rotary pumps for 1-5000 gpm, 50,000 ft head, 50-80% efficiency.
- 7. Reciprocating pumps for 10–10,000 gpm, 1,000,000 ft head max. Efficiency 70% at 10 HP, 85% at 50 HP, 90% at 500 HP.

REACTORS

- 1. The rate of reaction in every instance must be established in the laboratory, and the residence time or space velocity and product distribution eventually must be found in a pilot plant.
- 2. Dimensions of catalyst particles are 0.1 mm in fluidized beds, 1 mm in slurry beds, and 2-5 mm in fixed beds.
- 3. The optimum proportions of stirred tank reactors are with liquid level equal to the tank diameter, but at high pressures slimmer proportions are economical.

- 4. Power input to a homogeneous reaction stirred tank is 0.5-1.5 HP/1000 gal, but three times this amount when heat is to be . transferred.
- 5. Ideal CSTR (continuous stirred tank reactor) behavior is approached when the mean residence time is 5-10 times the length of time needed to achieve homogeneity, which is accomplished with 500–2000 revolutions of a properly designed stirrer.
- 6. Batch reactions are conducted in stirred tanks for small daily production rates or when the reaction times are long or when some condition such as feed rate or temperature must be programmed in some way.
- Relatively slow reactions of liquids and slurries are conducted in continuous stirred tanks. A battery of four or five in series is most economical.
- 8. Tubular flow reactors ate suited to high production rates at short residence times (sec or min) and when substantial heat transfer is needed. Embedded tubes or shell-and-tube construction then are used.
- *9.* In granular catalyst packed reactors, the residence time distribution often is no better than that of a five-stage CSTR battery.
- For conversions under about 95% of equilibrium, the performance of a five-stage CSTR battery approaches plug flow.

REFRIGERATION

- 1. A ton of refrigeration is the removal of 12,000 Btu/hr of heat.
- At various temperature levels: 0-50°F, chilled brine and glycol solutions; -50-40°F, ammonia, freons, butane; -150--50°F, ethane or propane.
- Compression refrigeration with 100°F condenser requires these HP/ton at various temperature levels: 1.24 at 20°F; 1.75 at 0°F;
 3.1 at -40°F; 5.2 at -80°F.
- 4. Below -80°F, cascades of two or three refrigerants are used.
- 5. In single stage compression, the compression ratio is limited to about 4.
- 6. In multistage compression, economy is improved with interstage flashing and recycling, so-called economizer operation.
- Absorption refrigeration (ammonia to -30°F, lithium bromide to +45°F) is economical when waste steam is available at 12 psig or so.

SIZE SEPARATION OF PARTICLES

- 1. Grizzlies that are constructed of parallel bars at appropriate spacings are used to remove products larger than 5 cm dia.
- Revolving cylindrical screens rotate at 15-20 rpm and below the critical velocity; they are suitable for wet or dry screening in the range of 10-60 mm.
- 3. Flat screens are vibrated or shaken or impacted with bouncing balls. Inclined screens vibrate at 600–7000 strokes/min and are used for down to 38 μ m although capacity drops off sharply below 200 μ m. Reciprocating screens operate in the range 30-1000 strokes/min and handle sizes down to 0.25 mm at the higher speeds.
- 4. Rotary sifters operate at 500-600 rpm and are suited to a range of 12 mm to 50 μm.
- 5. Air classification is preferred for fine sizes because screens of 150 mesh and finer are fragile and slow.
- 6. Wet classifiers mostly are used to make two product size ranges, oversize and undersize, with a break commonly in the range between 28 and 200 mesh. A rake classifier operates at about 9 strokes/min when making separation at 200 mesh, and 32

strokes/min at 28 mesh. Solids content is not critical, and that of the overflow may be 2-20% or more.

7. Hydrocyclones handle up to 600 cuft/min and can remove particles in the range of 300-5 μ m from dilute suspensions. In one case, a 20in. dia unit had a capacity of 1000 gpm with a pressure drop of 5 psi and a cutoff between 50 and 150 μ m.

UTILITIES: COMMON SPECIFICATIONS

- Steam: 15-30 psig, 250-275°F; 150 psig, 366°F; 400 psig, 448°F; 600 psig, 488°F or with 100-150°F superheat.
- Cooling water: Supply at 80-90°F from cooling tower, return at 115-125°F; return seawater at 110°F, return tempered water or steam condensate above 125°F.
- 3. Cooling air supply at 85-95°F; temperature approach to process, $40^{\circ}F$.
- 4. Compressed air at 45, 150, 300, or 450 psig levels.
- 5. Instrument air at 45 psig, 0°F dewpoint.
- 6. Fuels: gas of 1000 Btu/SCF at 5-10 psig, or up to 25psig for some types of burners; liquid at 6 million Btu/barrel.
- Heat transfer fluids: petroleum oils below 600°F, Dowtherms below 750°F, fused salts below 1100°F, direct fire or electricity above 450°F.
- 8. Electricity: 1-100 Hp, 220-550 V; 200-2500 Hp, 2300-4000 V.

VESSELS (DRUMS)

- 1. Drums are relatively small vessels to provide surge capacity or separation of entrained phases.
- 2. Liquid drums usually are horizontal.
- 3. Gas/liquid separators are vertical.
- 4. Optimum length/diameter = 3, but a range of 2.5-5.0 is common.
- 5. Holdup time is 5 min half full for reflux drums, 5-10 min for a product feeding another tower.
- 6. In drums feeding a furnace, 30 min half full is allowed.
- 7. Knockout drums ahead of compressors should hold no less than 10 times the liquid volume passing through per minute.
- 8. Liquid/liquid separators are designed for settling velocity of 2-j in./min.
- 9. Gas velocity in gas/liquid separators, $V = k\sqrt{\rho_L/\rho_V 1}$ ft/sec, with k = 0.35 with mesh deentrainer, k = 0.1 without mesh deentrainer.
- 10. Entrainment removal of 99% is attained with mesh pads of 4-12 in. thicknesses; 6 in. thickness is popular.
- 11. For vertical pads, the value of the coefficient in Step 9 is reduced by a factor of 2/3.
- **12.** Good performance can be expected at velocities of 30-100% of those calculated with the given k; 75% is popular.
- 13. Disengaging spaces of 6–18 in. ahead of the pad and 12 in. above the pad are suitable.
- 14. Cyclone separators can be designed for 95% collection of 5 μ m particles, but usually only droplets greater than 50 μ m need be removed.

VESSELS (PRESSURE)

- Design temperature between -20°F and 650°F is 50°F above operating temperature; higher safety margins are used outside the given temperature range.
- The design pressure is 10% or 10-25 psi over the maximum operating pressure, whichever is greater. The maximum operating pressure, in turn, is taken as 25 psi above the normal operation.
- 3. Design pressures of vessels operating at 0-10 psig and 600-1000°F are 40 psig.

- 4. For vacuum operation, design pressures are 15 psig and full vacuum.
- 5. Minimum wall thicknesses for rigidity: 0.25 in. for 42 in. dia and 'under, 0.32 in. for 42-60 in. dia, and 0.38 in. for over 60 in. dia.
- 6. Corrosion allowance 0.35 in. for known corrosive conditions, 0.15 in. for non-corrosive streams, and 0.06 in. for steam drums and air receivers.
- 7. Allowable working stresses are one-fourth of the ultimate strength of the material.
- 8. Maximum allowable stress depends sharply on temperature.

Temp	eratur	e (°F	·)		-20-650	750	850	1000
Low	alloy	steel	SA203	(psi)	18,750	15,650	9550	2500
Туре	302	stainle	ss (ps i)	18,750	18,750	15,900	6250

VESSELS (STORAGE TANKS)

- 1. For less than 1000 gal, use vertical tanks on legs.
- 2. Between 1000 and 10,000 gal, use horizontal tanks on concrete supports.
- 3. Beyond 10,000 gal, use vertical tanks on concrete foundations.
- 4. Liquids subject to breathing losses may be stored in tanks with floating or expansion roofs for conservation.
- 5. Freeboard is 15% below 500 gal and 10% above 500 gal capacity.
- 6. Thirty days capacity often is specified for raw materials and products, but depends on connecting transportation equipment schedules.
- 7. Capacities of storage tanks are at least 1.5 times the size of connecting transportation equipment; for instance, 7500 gal tank trucks, 34,500 gal tank cars, and virtually unlimited barge and tanker capacities.

INTRODUCTION

Ithough this book is devoted to the selection and design of individual equipment, some mention should be made of integration of a number of units into a process. Each piece of equipment interacts with several others in a plant, and the range of its required

1.1. PROCESS DESIGN

Process design establishes the sequence of chemical and physical operations; operating conditions; the duties, major specifications, and materials of construction (where critical) of all process equipment (as distinguished from utilities and building auxiliaries); the general arrangement of equipment needed to ensure proper functioning of the plant; line sizes; and principal instrumentation. The process design is summarized by a process flowsheet, a material and energy balance, and a set of individual equipment specifications. Varying degrees of thoroughness of a process design may be required for different purposes. Sometimes only a preliminary design and cost estimate are needed to evaluate the advisability of further research on a new process or a proposed plant expansion or detailed design work; or a preliminary design may be needed to establish the approximate funding for a complete design and construction. A particularly valuable function of preliminary design is that it may reveal lack of certain data needed for final design. Data of costs of individual equipment are supplied in this book, but the complete economics of process design is beyond its scope.

1.2. EQUIPMENT

Two main categories of process equipment are proprietary and custom-designed. Proprietary equipment is designed by the manufacturer to meet performance specifications made by the user; these specifications may be regarded as the process design of the equipment. This category includes equipment with moving parts such as pumps, compressors, and drivers as well as cooling towers, dryers, filters, mixers, agitators, piping equipment, and valves, and even the structural aspects of heat exchangers, furnaces, and other equipment. Custom design is needed for many aspects of chemical reactors, most vessels, multistage separators such as fractionators, and other special equipment not amenable to complete standardization.

Only those characteristics of equipment are specified by process design that are significant from the process point of view. On a pump, for instance, process design will specify the operating conditions, capacity, pressure differential, NPSH, materials of construction in contact with process liquid, and a few other items, but not such details as the wall thickness of the casing or the type of stuffing box or the nozzle sizes and the foundation dimensionsalthough most of these omitted items eventually must be known before a plant is ready for construction. Standard specification forms are available for most proprietary kinds of equipment and for summarizing the details of all kinds of equipment. By providing suitable check lists, they simplify the work by ensuring that all needed data have been provided. A collection of such forms is in Appendix B.

Proprietary equipment is provided "off the shelf" in limited sizes and capacities. Special sizes that would fit particular applications more closely often are more expensive than a larger performance is dependent on the others in terms of material and energy balances and rate processes. This chapter will discuss general background material relating to complete process design, and Chapter 2 will treat briefly the basic topic of flowsheets.

standard size that incidentally may provide a worthwhile safety factor. Even largely custom-designed equipment, such as vessels, is subject to standardization such as discrete ranges of head diameters, pressure ratings of nozzles, sizes of **manways**, and kinds of trays and packings. Many codes and standards are established by government agencies, insurance companies, and organizations sponsored by engineering societies. Some standardizations within individual plants are arbitrary choices from comparable methods, made to simplify construction, maintenance, and repair: for example, restriction to instrumentation of a particular manufacturer or to a limited number of sizes of heat exchanger tubing or a particular method of installing liquid level gage glasses. All such restrictions must be home in mind by the process designer.

VENDORS' QUESTIONNAIRES

A manufacturer's or vendor's inquiry form is a questionnaire whose completion will give him the information on which to base a specific recommendation of equipment and a price. General information about the process in which the proposed equipment is expected to function, amounts and appropriate properties of the streams involved, and the required performance are basic. The nature of additional information varies from case to case; for instance, being different for filters than for pneumatic conveyors. Individual suppliers have specific inquiry forms. A representative selection is in Appendix C.

SPECIFICATION FORMS

When completed, a specification form is a record of the salient features of the equipment, the conditions under which it is to operate, and its guaranteed performance. Usually it is the basis for a firm price quotation. Some of these forms are made up by organizations such as TEMA or API, but all large engineering contractors and many large operating companies have other forms for their own needs. A selection of specification forms is in Appendix B.

1.3. CATEGORIES OF ENGINEERING PRACTICE

Although the design of a chemical process plant is initiated by chemical engineers, its complete design and construction requires the inputs of other specialists: mechanical, structural, electrical, and instrumentation engineers; vessel and piping designers; and purchasing agents who know what may be available at attractive prices. On large projects all these activities are correlated by a job engineer or project manager; on individual items of equipment or small projects, the process engineer naturally assumes this function. A key activity is the writing of specifications for soliciting bids and ultimately purchasing equipment. Specifications must be written so explicitly that the bidders are held to a uniform standard and a clear-cut choice can be made on the basis of their offerings alone.



Figure 1.1. Progress of material commitment, engineering manhours, and construction [*Matozzi*, Oil Gas. J. p. 304, (23 March 1953)].



Figure 1.2. Rate of application of engineering manhours of various categories. The area between the curves represents accumulated manhours for each speciality up to a given % completion of the project [Miller, Chem. Eng., p. 188, (July 1956)].

For a typical project, Figure 1.1 shows the distributions of engineering, material commitment, and construction efforts. Of the engineering effort, the process engineering is a small part. Figure 1.2 shows that it starts immediately and finishes early. In terms of money, the cost of engineering ranges from 5 to 15% or so of the total plant cost; the lower value for large plants that are largely patterned after earlier ones, and the higher for small plants or those based on new technology or unusual codes and specifications.

1.4. SOURCES OF INFORMATION FOR PROCESS DESIGN

A selection of books relating to process design methods and data is listed in the references at the end of this chapter. Items that are especially desirable in a personal library or readily accessible are identified. Specialized references are given throughout the **book** in connection with specific topics.

The extensive chemical literature is served by the bibliographic items cited in References, Section 1.2, Part B. The book by Rasmussen and Fredenslund (1980) is addressed to chemical **engineers** and cites some literature not included in some of the other bibliographies, as well as information about proprietary data banks. The book by Leesley (References, Section 1.1, Part B) has much information about proprietary data banks and design methods. In its current and earlier editions, the book **by** Peters and Timmerhaus has many useful bibliographies on classified topics.

For information about chemical manufacturing processes, the main encyclopedic references are Kirk-Othmer (1978–1984), McKetta and Cunningham (1976–date) and Ullmann (1972-1983) (References, Section 1.2, Part B). The last of these is in German,

but an English version was started in 1984 and three volumes per year are planned; this beautifully organized reference should be most welcome.

The most comprehensive compilation of physical property data is that of Landolt-Bornstein (1950–date) (References, Section 1.2, Part C). Although most of the material is in German, recent volumes have detailed tables of contents in English and some volumes are largely in English. Another large compilation, somewhat venerable but still valuable, is the International Critical Tables (1926-1933). Data and methods of estimating properties of hydrocarbons and their mixtures are in the API Data Book (1971–date) (References, Section 1.2, Part C). More general treatments of estimation of physical properties are listed in References, Section 1.1, Part C. There are many compilations of special data such as solubilities, vapor pressures, phase equilibria, transport and thermal properties, and so on. A few of them are listed in References, Section 1.2, Part D, and references to many others are in the References, Section 1.2, Part B.

Information about equipment sizes and configurations, and sometimes performance, of equipment is best found in manufacturers' catalogs. Items 1 and 2 of References, Section 1.1, Part D, contain some advertisements with illustrations, but perhaps their principal value is in the listings of manufacturers by the kind of equipment. Thomas Register covers all manufacturers and so is less convenient at least for an initial search. The other three items of this group of books have illustrations and descriptions of all kinds of chemical process equipment. Although these books are old, one is surprised to note how many equipment designs have survived.

1.5. CODES, STANDARDS, AND RECOMMENDED PRACTICES

A large body of rules has been developed over the years to ensure the safe and economical design, fabrication and testing of equipment, structures, and materials. Codification of these rules has been done by associations organized for just such purposes, by professional societies, trade groups, insurance underwriting companies, and government agencies. Engineering contractors and large manufacturing companies usually maintain individual sets of standards so as to maintain continuity of design and to simplify maintenance of plant. Table 1.1 is a representative table of contents of the mechanical standards of a large oil company.

Typical of the many thousands of items that are standardized in the field of engineering are limitations on the sizes and wall **thicknesses** of piping, specifications of the compositions of alloys, stipulation of the safety factors applied to strengths of construction materials, testing procedures for many kinds of materials, and so on.

Although the safe design practices recommended by professional and trade associations have no legal standing where they have not actually been incorporated in a body of law, many of them have the respect and confidence of the engineering profession as a whole and have been accepted by insurance underwriters so they are widely observed. Even when they are only voluntary, standards constitute a digest of experience that represents a minimum requirement of good practice.

Two publications by Burklin (References, Section 1.1, Part B) are devoted to standards of importance to the chemical industry. Listed are about 50 organizations and 60 topics with which they are concerned. National Bureau of Standards Publication 329 contains about 25,000 titles of U.S. standards. The NBS-SIS service maintains a reference collection of 200,000 items accessible by letter or phone. Information about foreign standards is obtainable through the American National Standards Institute (ANSI).

A listing of codes and standards bearing directly on process

TABLE 1.1. Internal Engineering Standards of a Large Petroleum Refinery'

1	Appropriations and mechanical orders (10)
2	Buildings-architectural (15)
3	Buildings-mechanical (10)
4	Capacities and weights (25)
5	Contracts (1 0)
6	Cooling towers (10)
7	Correspondence (5)
8	Designation and numbering rules for equipment and facilities (10)
9	Drainage (25)
10	Electrical (10)
11	Excavating, grading, and paving (10)
12	Fire fighting (10)
13	Furnaces and boilers (10)
14	General instructions (20)
15	Handling equipment (5)
16	Heat exchangers (10)
17	Instruments and controls (45)
18	Insulation (10)
19	Machinery (35)
20	Material procurement and disposition (20)
21	Material selection (5)
22	Miscellaneous process equipment (25)
23	Personnel protective equipment (5)
24	Piping (150)
25	Piping supports (25)
26	Plant layout (20)
27	Pressure vessels (25)
28	Protective coatings (10)
29	Roads and railroads (25)
30	Storage vessels (45)
31	Structural (35)
32	Symbols and drafting practice (15)
33	Welding (10)

'Figures in parentheses identify the numbers of distinct standards.

TABLE 1.2. Codes and Standards of Direct Bearing on Chemical Process Design (a Selection)

- A. American Institute of Chemical Engineers, 345 E. 47th St., New York, NY 10017
 - 1. Standard testing procedures; 21 have been published, for example on centrifuges, filters, mixers, firer heaters
- B. American Petroleum Institute, 2001 L St. NW, Washington, DC 20037
 - 2. Recommended practices for refinery inspections
 - 3. Guide for inspection of refinery equipment
 - 4. Manual on disposal of refinery wastes
 - 5. Recommended practice for design and construction of large, low pressure storage tanks
 - 6. Recommended practice for design and construction of pressure relieving devices
 - 7. Recommended practices for safety and fire protection
- C. American Society of Mechanical Engineers, 345 W. 47th St., New York. NY 10017
 - 8. ASME Boiler and Pressure Vessel Code. Sec. VIII, Unfired Pressure Vessels
 - 9. Code for pressure piping
 - 10; Scheme for identification of piping systems
- D. American Society for Testing Materials, 1916 Race St., Philadelphia, PA 19103
 - 11. ASTM Standards, 66 volumes in 16 sections, annual, with about 30% revision each year
- E. American National Standards Institute (ANSI), 1430 Broadway, New York, NY 10018
 - 12. Abbreviations, letter symbols, graphical symbols, drawing and drafting room practice

TABLE 1.2-(continued)

- F. Chemical Manufacturers' Association, 2501 M St. NW, Washington, DC 20037
 - 13. Manual of standard and recommended practices for containers, tank cars, pollution of air and water
 - 14. Chemical safety data sheets of individual chemicals
- G. Cooling Tower Institute, 19827 Highway 45 N, Spring, TX 77388 15. Acceptance test procedure for water cooling towers of mechanical draft industrial type
- H. Hydraulic Institute, 712 Lakewood Center N, 14600 Detroit Ave., Cleveland, OH 44107
 - 16. Standards for centrifugal, reciprocating, and rotary pumps 17. Pipe friction manual
- I. Instrument Society of America (ISA), 67 Alexander Dr., Research Triangle Park, NC 27709
 - 18. Instrumentation flow plan symbols
 - 19. Specification forms for instruments
- 20. Dynamic response testing of process control instrumentation
- J. Tubular Exchangers Manufacturers' Association, 25 N Broadway, Tarrytown, NY 10591
 - 21. TEMA standards
- K. International Standards Organization (ISO), 1430 Broadway, New York. NY 10018
 - 22. Many standards

TABLE 1.3. Codes and Standards Supplementary to Process Design (a Selection)

- A. American Concrete Institute, 22400 W. 7 Mile Rd., Detroit, MI 48219
 - 1. Reinforced concrete design handbook
 - 2. Manual of standard practice for detailing reinforced concrete structures
- B. American Institute of Steel Construction, 400 N. Michigan Ave., Chicago, IL 60611
 - 3. Manual of steel construction

 - 4. Standard practice for steel buildings and bridges
- C. American Iron and Steel Institute, 1000 16th St. NW, Washington, DC 20036
 - 5. AISI standard steel compositions
- D. American Society of Heating, Refrigerating and Air Conditioning Engineers (ASHRE), 1791 Tullie Circle NE, Atlanta, GA 30329 6. Refrigerating data book
- E. Institute of Electrical and Electronics Engineers, 345 E. 47th St., New York, NY 10017
 - 7. Many standards
- F. National Bureau of Standards, Washington, DC
 - 8. American standard building code
 - 9. National electrical code
- G. National Electrical Manufacturers Association, 2101 L St. NW, Washington, DC 20037 10. NEMA standards

design is in Table 1.2, and of supplementary codes and standards in Table 1.3.

1.6. MATERIAL AND ENERGY BALANCES

Material and energy balances arc based on a conservation law which is stated generally in the form

input + source = output + sink + accumulation.

The individual terms can be plural and can be rates as well as absolute quantities. Balances of particular entities are made around a bounded region called a system. Input and output quantities of an entity cross the boundaries. A source is an increase in the amount

of the entity that occurs without a crossing of the boundary; for example, an increase in the sensible enthalpy or in the amount of a substance as a consequence of chemical reaction. Analogously, sinks are decreases without a boundary crossing, as the disappearance of water from a fluid stream by adsorption onto a solid phase within the boundary.

Accumulations are time rates of change of the amount of the entities within the boundary. For example, in the absence of sources and sinks, an accumulation occurs when the input and output rates are different. In the steady state, the accumulation is zero.

Although the principle of balancing is simple, its application requires knowledge of the performance of all the kinds of equipment comprising the system and of the phase relations and physical properties of all mixtures that participate in the process. As a consequence of trying to cover a variety of equipment and processes, the books devoted to the subject of material and energy balances always run to several hundred pages. Throughout this book, material and energy balances are utilized in connection with the design of individual kinds of equipment and some processes. Cases involving individual pieces of equipment usually are relatively easy to balance, for example, the overall balance of a distillation column in Section 13.4.1 and of nonisothermal reactors of Tables 17.4-17.7. When a process is maintained isothermal, only a material balance is needed to describe the process, unless it is also required to know the net heat transfer for maintaining a constant temperature.

In most plant design situations of practical interest, however, the several pieces of equipment interact with each other, the output of one unit being the input to another that in turn may recycle part of its output to the inputter. Common examples are an absorber-stripper combination in which the performance of the absorber depends on the quality of the absorbent being returned from the stripper, or a catalytic cracker-catalyst regenerator system whose two parts interact closely.

Because the performance of a particular piece of equipment depends on its input, recycling of streams in a process introduces temporarily unknown, intermediate streams whose amounts, compositions, and properties must be found by calculation. For a plant with dozens or hundreds of streams the resulting mathematical problem is formidable and has led to the development of many computer algorithms for its solution, some of them making quite rough approximations, others more nearly exact. Usually the problem is solved more easily if the performance of the equipment is specified in advance and its size is found after the balances are completed. If the equipment is existing or must be limited in size, the balancing process will require simultaneous evaluation of its performance and consequently is a much more involved operation, but one which can be handled by computer when necessary.

The literature of this subject naturally is extensive. An early book (for this subject), Nagiev's Theory of Recycle Processes in Chemical Engineering (Macmillan, New York, 1964, Russian edition, 1958) treats many practical cases by reducing them to systems of linear algebraic equations that are readily solvable. The book by Westerberg et al., Process Flowsheeting (Cambridge Univ. Press, Cambridge, 1977) describes some aspects of the subject and has an extensive bibliography. Benedek in Steady State Flowsheeting of Chemical Plants (Elsevier, New York, 1980) provides a detailed description of one simulation system. Leesley in Computer-Aided Process Design (Gulf, Houston, 1982) describes the capabilities of some commercially available flowsheet simulation programs. Some of these incorporate economic balance with material and energy balances. A program MASSBAL in BASIC language is in the book of Sinnott et al., Design, Vol. 6 (Pergamon, New York, 1983); it can handle up to 20 components and 50 units when their several outputs are specified to be in fixed proportions.



Figure 1.3. Notation of flow quantities in a reactor (1) and distillation column (2). $A_{ij}^{(k)}$ designates the amount of component A in stream k proceeding from unit i to unit j. Subscripts 0 designates a source or sink beyond the boundary limits. Γ designates a total flow quantity.

A key factor in the effective formulation of material and energy balances is a proper notation for equipment and streams. Figure 1.3, representing a reactor and a separator, utilizes a simple type. When the pieces of equipment are numbered *i* and *j*, the notation $A_{ij}^{(k)}$ signifies the flow rate of substance *A* in stream *k* proceeding from unit *i* to unit *j*. The total stream is designated $\Gamma_{ij}^{(k)}$. Subscript *t* designates a total stream and subscript 0 designates sources or sinks outside the system. Example 1.1 adopts this notation for balancing a reactor-separator process in which the performances are specified in advance.

Since this book is concerned primarily with one kind of equipment at a time, all that need be done here is to call attention to the existence of the abundant literature on these topics of recycle calculations and flowsheet simulation.

1.7. ECONOMIC BALANCE

Engineering enterprises always are subject to monetary considerations, and a balance is sought between fixed and operating costs. In the simplest terms, fixed costs consist of depreciation of the investment plus interest on the working capital. Operating costs include labor, raw materials, utilities, maintenance, and overheads which consists in turn of administrative, sales and research costs. Usually as the capital cost of a process unit goes up, the operating cost goes down. For example, an increase in control instrumentation and automation at a higher cost is accompanied by a reduction in operating labor cost. Somewhere in the summation of these factors there is a minimum which should be the design point in the absence of any contrary intangibles such as building for the future or unusual local conditions.

Costs of many individual pieces of equipment are summarized in Chapter 20, but analysis of the costs of complete processes is beyond the scope of this book. References may be made, however, to several collections of economic analyses of chemical engineering interest that have been published:

1. AIChE Student Contest Problems (annual) (AIChE, New York).

EXAMPLE 1.1

Material Balance of a Chlorination Process with Recycle A plant for the chlorination has the flowsheet shown. From pilot plant work, with a chlorine/benzene charge weight ratio of 0.82, the composition of the reactor effluent is

A. C _e H _e	0.247
B. Cl ₂	0.100
C. C.H.CI	0.3174
D. C.H.Cl.	0.1559
E. HČI	0.1797

Separator no. 2 returns 80% of the unreacted chlorine to the reactor and separator no. 3 returns 90% of the benzene. Both recycle streams are pure. Fresh chlorine is charged at such a rate that the weight ratio of chlorine to benzene in the total charge remains 0.82. The amounts of other streams are-found by material balances and are shown in parentheses on the sketch per 100 lbs of fresh benzene to the system.



- Bodman, Industrial Practice of Chemical Process Engineering (MIT Press, Cambridge, MA, 1968).
- Rase, Chemical Reactor Design for Process Plants, Vol. II, Case Studies (Wiley, New York, 1977).
- 4. Washington University, St. Louis, *Case Studies in Chemical Engineering Design (22* cases to 1984).

Somewhat broader in scope are:

- 5. Wei et al., The Structure of the Chemical Processing Industries (McGraw-Hill, New York, 1979).
- Skinner et al., Manufacturing Policy in the Oil Industry (Irwin, Homewood, IL., 1970).
- 7. Skinner et al., *Manufacturing Policy in the Plastics Industry* (Irwin, Homewood, II., 1968).

Many briefer studies of individual equipment appear in some books, of which a selection is as follows:

- Happel and Jordan, *Chemical Process Economics* (Dekker, New York, 1975):
 - 1. Absorption of ethanol from a gas containing CO, (p. 403).
 - 2. A reactor-separator for simultaneous chemical reactions (p. 419).
 - 3. Distillation of a binary mixture (p. 38.5).
 - 4. A heat exchanger and cooler system (p. 370).
 - 5. Piping of water (p. 353).
 - 6. Rotary dryer (p. 414).

- Jelen et al., *Cost and Optimization Engineering* (McGraw-Hill, New York, 1983):
 - 7. Drill bit life and replacement policy (p. 223).
 - 8. Homogeneous flow reactor (p. 229).
 - 9. Batch reaction with negligible downtime (p. 236).
- Peters and Timmerhaus, *Plant Design and Economics for Chemical Engineers* (McGraw-Hill, New York, 1980):
 10. Shell and tube cooling of air with water (p. 688).
- Rudd and Watson, *Strategy of Process Engineering* (Wiley, New York, 1968):
 - 11. Optimization of a three stage refrigeration system (p. 172).
- Sherwood, A Course in Process Design (MIT Press, Cambridge, MA, 1963):
 - **12.** Gas transmission line (p. 84).
 - 13. Fresh water from sea water by evaporation (p. 138).
- Ulrich, A Guide to Chemical Engineering Process Design and Economics (Wiley, New York, 1984):

14. Multiple effect evaporator for Kraft liquor (p. 347).

 Walas, Reaction Kinetics for Chemical Engineers (McGraw-Hill, New York, 1959):
 15 Optimum number of vascele in a CSTR bettery (p. 98)

15. Optimum number of vessels in a CSTR battery (p. 98).

Since capital, labor, and energy costs have not escalated equally over the years since these studies were made, their conclusions are subject to reinterpretation, but the patterns of study that were used should be informative.

Because of the rapid escalation of energy costs in recent years,

closer appraisals of energy utilizations by complete processes are being made, from the standpoints of both the conservation laws and the second law of thermodynamics. In the latter cases attention is focused on changes in entropy and in the related availability function, $AB = AH - T_0\Delta S$, with emphasis on work as the best possible transformation of energy. In this way a second law analysis of a process will reveal where the greatest generation of entropy occurs and where possibly the most improvement can be made by appropriate changes of process or equipment. Such an analysis of a cryogenic process for air separation was made by Benedict and Gyftopolous [in Gaggioli (Ed.), *Thermodynamic Second Law Analysis*, ACS Symposium Series No. 122, American Chemical Society, Washington, DC, 1980]; they found a pressure drop at which the combination of exchanger and compressor was most economical.

A low second law efficiency is not always realistically improvable. Thus Weber and Meissner *(Thermodynamics for Chemical Engineers, John Wiley, New York, 1957)* found a 6% efficiency for the separation of ethanol and water by distillation which is not substantially improvable by redesign of the distillation process. Perhaps this suggests that more efficient methods than distillation should be sought for the separation of volatile mixtures, but none has been found at competitive cost.

Details of the thermodynamic basis of availability analysis are dealt with by Moran (*Availability Analysis*, Prentice-Hall, Englewood Cliffs, NJ, 1982). He applies the method to a cooling tower, heat pump, a cryogenic process, coal gasification, and particularly to the efficient use of fuels.

An interesting conclusion reached by Linnhoff [in Seider and Mah (Eds.), Foundations of Computer-Aided Process Design, AIChE, New York, 1981] is that "chemical processes which are properly designed for energy versus capital cost tend to operate at approximately 60% efficiency." A major aspect of his analysis is recognition of practical constraints and inevitable losses. These may include material of construction limits, plant layout, operability, the need for simplicity such as limits on the number of compressor stages or refrigeration levels, and above all the recognition that, for low grade heat, heat recovery is preferable to work recovery, the latter being justifiable only in huge installations. Unfortunately, the edge is taken off the dramatic 60% conclusion by Linnhoff's admission that efficiency cannot be easily defined for some complexes of interrelated equipment. For example, is it economical to recover 60% of the propane or 60% of the ethane from a natural gas?

1.8. SAFETY FACTORS

In all of the factors that influence the performance of equipment and plant there are elements of uncertainty and the possibility of error, including inaccuracy of physical data, basic correlations of behavior such as pipe friction or tray efficiency or gas-liquid distribution, necessary approximations of design methods and calculations, not entirely known behavior of materials of construction, uncertainty of future market demands, and changes in operating performance with time. The solvency of the project, the safety of the operators and the public, and the reputation and career of the design engineer are at stake. Accordingly, the experienced engineer will apply safety factors throughout the design of a plant. Just how much of a factor should be applied in a particular case cannot be stated in general terms because circumstances vary widely. The inadequate performance of a particular piece of equipment may be compensated for by the superior performance of associated equipment, as insufficient trays in a fractionator may be compensated for by increases in reflux and reboiling, if that equipment can take the extra load.

With regard to specific types of equipment, the safety factor practices of some 250 engineers were ascertained by a questionnaire and summarized in Table 1.4; additional figures are given by Peters and Timmerhaus (References, Section 1.1, Part B, pp. 35-37). Relatively inexpensive equipment that can conceivably serve as a bottleneck, such as pumps, always is liberally sized; perhaps as much as 50% extra for a reflux pump. In an expanding industry it is a matter of policy to deliberately oversize certain major equipment that cannot be supplemented readily or modified suitably for increased capacity; these are safety factors to account for future trends.

Safety factors should not be used to mask inadequate or careless design work. The design should be the best that can be made in the time economically justifiable, and the safety factors should be estimated from a careful consideration of all factors entering into the design and the possible future deviations from the design conditions.

Sometimes it is possible to evaluate the range of validity of measurements and correlations of physical properties, phase equilibrium behavior, mass and heat transfer efficiencies and similar factors, as well as the fluctuations in temperature, pressure, flow, etc., associated with practical control systems. Then the effects of such data on the uncertainty of sizing equipment can be estimated. For example, the mass of a distillation column that is related directly to its cost depends on at least these factors:

- 1. The vapor-liquid equilibrium data.
- 2. The method of calculating the reflux and number of trays.
- 3. The tray efficiency.
- 4. Allowable vapor rate and consequently the tower diameter at a given tray spacing and estimated operating surface tension and fluid densities.
- 5. Corrosion allowances.

Also such factors as allowable tensile strengths, weld efficiencies, and possible inaccuracies of formulas used to calculate shell and head thicknesses may be pertinent.

When a quantity is a function of several variables,

$$Y = y(x_1, x_2, \ldots),$$

its differential is

$$dy = \frac{\partial y}{\partial x_1} dx_1 + \frac{\partial y}{\partial x_2} dx_2 + \cdots.$$

Some relations of importance in chemical engineering have the form

$$\mathbf{y} = (x_1)^a (x_2)^b \cdot \cdot \cdot,$$

whose differential is rearrangable to

$$\frac{dy}{y} = a\frac{dx_1}{x_1} + b\frac{dx_2}{x_2} + \cdots$$

that is, the relative uncertainty or error in the function is related linearly to the fractional uncertainties of the independent variables. For example, take the case of a steam-heated thermosyphon reboiler on a distillation column for which the heat transfer equation is

$$q = UA\Delta T.$$

The problem is to find how the heat transfer rate can vary when the other quantities change. U is an experimental value that is known

Equipment	Design Variable	Range of Safety Factor (%)
Compressors, reciprocating	piston displacement	11-21
Conveyors, screw	diameter	8-21
Hammer mills	power input	15-21"
Filters, plate-and-frame	area	11-21″
Filters, rotary	area	14-20"
Heat exchangers, shell and tube for liquids	area	11-18
Pumps, centrifugal	impeller diameter	7-14
Separators, cyclone	diameter	7-11
Towers, packed	diameter	11-18
Towers, tray	diameter	10–16
Water cooling towers	volume	12-20

TABLE 1.4. Safety Factors in Equipment Design: Results of a Questionnaire

^a Based on pilot plant tests.

[Michelle, Beattie, and Goodgame, Chem. Eng. frog. 50,332 (1954)).

only to a certain accuracy. AT may be uncertain because of possible fluctuations in regulated steam and tower pressures. *A*, the effective area, may be uncertain because the submergence is affected by the liquid level controller at the bottom of the column. Accordingly,

,

$$\frac{dq}{q} = \frac{dU}{U} + \frac{dA}{A} + \frac{d(\Delta T)}{A}$$

that is, the fractional uncertainty of q is the sum of the fractional uncertainties of the quantities on which it is dependent. In practical cases, of course, some uncertainties may be positive and others negative, so that they may cancel out in part; but the only safe viewpoint is to take the sum of the absolute values. Some further discussion of such cases is by Sherwood and Reed, in *Applied Mathematics in Chemical Engineering* (McGraw-Hill, New York, 1939).

It is not often that proper estimates can be made of uncertainties of all the parameters that influence the performance or required size of particular equipment, but sometimes one particular parameter is dominant. All experimental data scatter to some extent, for example, heat transfer coefficients; and various correlations of particular phenomena disagree, for example, equations of state of liquids and gases. The sensitivity of equipment sizing to uncertainties in such data has been the subject of some published information, of which a review article is by Zudkevich [*Encycl. Chem. Proc. Des.* 14, 431-483 (1982)]; some of his cases are:

- 1. Sizing of isopentane/pentane and propylene/propane splitters.
- 2. Effect of volumetric properties on sizing of an ethylene compressor.
- 3. Effect of liquid density on metering of LNG.
- 4. Effect of vaporization equilibrium ratios, *K*, and enthalpies on cryogenic separations.
- 5. Effects of VLE and enthalpy data on design of plants for coal-derived liquids.

Examination of such studies may lead to the conclusion that some of the safety factors of Table 1.4 may be optimistic. But long experience in certain areas does suggest to what extent various uncertainties do cancel out, and overall uncertainties often do fall in the range of 10-20% as stated there. Still, in major cases the uncertainty analysis should be made whenever possible.

1.9. SAFETY OF PLANT AND ENVIRONMENT

The safe practices described in the previous section are primarily for assurance that the equipment have adequate performance over

anticipated ranges of operating conditions. In addition, the design of equipment and plant must minimize potential harm to personnel **and** the public in case of accidents, of which the main causes are

- a. human failure,
- **b.** failure of equipment or control instruments,
- c. failure of supply of utilities or key process streams,
- d. environmental events (wind, water, and so on).

A more nearly complete list of potential hazards is in Table 1.5, and a checklist referring particularly to chemical reactions is in Table 1.6.

Examples of common safe practices are pressure relief valves, vent systems, flare stacks, snuffing steam and fire water, escape hatches in explosive areas, dikes around tanks storing hazardous materials, turbine drives as spares for electrical motors in case of power failure, and others. Safety considerations are paramount in the layout of the plant, particularly isolation of especially hazardous operations and accessibility for corrective action when necessary.

Continual monitoring of equipment and plant is standard practice in chemical process plants. Equipment deteriorates and operating conditions may change. Repairs sometimes are made with "improvements" whose ultimate effects on the operation may not be taken into account. During start-up and shut-down, stream compositions and operating conditions are much different from those under normal operation, and their possible effect on safety must be taken into account. Sample checklists of safety questions for these periods are in Table 1.7.

Because of the importance of safety and its complexity, safety engineering is a speciality in itself. In chemical processing plants of any significant size, loss prevention reviews are held periodically by groups that always include a representative of the safety department. Other personnel, as needed by the particular situation, are from manufacturing, maintenance, technical service, and possibly research, engineering, and medical groups. The review considers any changes made since the last review in equipment, repairs, feedstocks and products, and operating conditions.

Detailed safety checklists appear in books by Fawcett and Wood (Chap. 32, Bibliography 1.1, Part E) and Wells (pp. 239-257, Bibliography 1.1, Part E). These books and the large one by Lees (Bibliography 1.1, Part E) also provide entry into the vast literature of chemical process plant safety. Lees has particularly complete bibliographies. A standard reference on the properties of dangerous materials is the book by Sax (1984) (References, Section 1.1, Part E). The handbook by Lund (1971) (References, Section 1.1, Part E) on industrial pollution control also may be consulted.

TABLE 1.5. Some Potential Hazards Energy Source Process chemicals, fuels, nuclear reactors, generators, batteries Source of ignition, radio frequency energy sources, activators, radiation sources Rotating machinery, prime movers, pulverisers, grinders, conveyors, belts, cranes Pressure containers, moving objects, falling objects Release of Material Spillage, leakage, vented material Exposure effects, toxicity, burns, bruises, biological effects Flammability, reactivity, explosiveness, corrosivity and fire-promoting properties of chemicals Wetted surfaces, reduced visibility, falls, noise, damage Dust formation, mist formation, spray Fire hazard Fire, fire spread, fireballs, radiation Explosion, secondary explosion, domino effects Noise, smoke, toxic fumes, exposure effects Collapse, falling objects, fragmentation Process state High/low/changing temperature and pressure Stress concentrations, stress reversals, vibration, noise Structural damage or failure, falling objects, collapse Electrical shock and thermal effects, inadvertent activation, power source failure Radiation, internal fire, overheated vessel Failure of equipment/utility supply/flame/instrument/component Start-up and shutdown condition Maintenance, construction and inspection condition Environmental effects Effect of plant on surroundings, drainage, pollution, transport, wind and light change, source of ignition/vibration/noise/radio interference/fire spread/explosion Effect of surroundings on plant (as above) Climate, sun, wind, rain, snow, ice, grit, contaminants, humidity, ambient conditions Acts of God, earthquake, arson, flood, typhoon, force majeure Site layout factors, groups of people, transport features, space limitations, geology, geography Processes Processes subject to explosive reaction or detonation Processes which react energetically with water or common contaminants Processes subject to spontaneous polymerisation or heating Processes which are exothermic Processes containing flammables and operated at high pressure or high temperature or both Processes containing flammables and operated under refrigeration Processes in which intrinsically unstable compounds are present Processes operating in or near the explosive range of materials Processes involving highly toxic materials Processes subject to a dust or mist explosion hazard Processes with a large inventory of stored pressure energy

Operations

The vaporisation and diffusion of flammable or toxic liquids or gases The dusting and dispersion of combustible or toxic solids

- The spraying, misting or fogging of flammable combustible materials or strong oxidising agents and their mixing
- The separation of hazardous chemicals from inerts or diluents The temperature and pressure increase of unstable liquids

(Wells, Safety in Process P/ant Design, George Godwin, London, 1980).

TABLE 1.6. Safety Checklist of Questions About Chemical Reactions

- 1. Define potentially hazardous reactions. How are they isolated? Prevented? (See Chaps. 4, 5, and 16)
- 2. Define process variables which could, or do, approach limiting conditions for hazard. What safeguards are provided against such variables?
- 3. What unwanted hazardous reactions can be developed through unlikely flow or process conditions or through contamination?
- 4. What combustible mixtures can occur within equipment?
- What precautions are taken for processes operating near or within the flammable limits? (Reference: S&PP Design Guide No. 8.) (See Chap. 19)
- 6. What are process margins of safety for all reactants and intermediates in the process?
- 7. List known reaction rate data on the normal and possible abnormal reactions
- 8. How much heat must be removed for normal, or abnormally possible, exothermic reactions? (see Chaps. 7, 17, and 18)
- How thoroughly is the chemistry of the process including desired and undesired reactions known? (See NFPA 491 M, Manual of Hazardous Chemical Reactions)
- 10. What provision is made for rapid disposal of reactants if required by emergency?
- 11. What provisions are made for handling impending runaways and for short-stopping an existing runaway?
- 12. Discuss the hazardous reactions which could develop as a result of mechanical equipment (pump, agitator, etc.) failure
- 13. Describe the hazardous process conditions that can result from gradual or sudden blockage in equipment including lines
- 14. Review provisions for blockage removal or prevention
- 15. What raw materials or process materials or process conditions can be adversely affected by extreme weather conditions? Protect against such conditions
- 16. Describe the process changes including plant operation that have been made since the previous process safety review

(Fawcett and Wood, Safety and Accident Prevention in Chemical Operations, Wiley, New York, 1982, pp. 725-726. Chapter references refer to this book.)

TABLE 1.7. Safety Checklist of Questions About Start-up and Shut-down

Start-up Mode (§4.1)

- D1 Can the start-up of plant be expedited safely? Check the following:
 (a) Abnormal concentrations, phases, temperatures, pressures, levels, flows, densities
 - (b) Abnormal quantities of raw materials, intermediates and utilities (supply, handling and availability)
 - (c) Abnormal quantities and types of effluents and emissions (§1.6.10)
 - (d) Different states of catalyst, regeneration, activation
 - (e) Instruments out of range, not in service or de-activated, incorrect readings, spurious trips
 - (f) Manual control, wrong routeing, sequencing errors, poor identification of valves and lines in occasional use, lock-outs, human error, improper start-up of equipment (particularly prime movers)
 - (g) Isolation, purging
 - (h) Removal of air, undesired process material, chemicals used for cleaning, inerts, water, oils, construction debris and ingress of same
 - (i) Recycle or disposal of off-specification process materials
 - (j) Means for ensuring construction/maintenance completed
 - (k) Any plant item failure on initial demand and during operation in this mode
 - Lighting of flames, introduction of material, limitation of heating rate

(m) Different modes of the start-up of plant: Initial start-up of plant Start-up of plant section when rest of plant down Start-up of plant section when other plant on-stream Start-up of plant after maintenance Preparation of plant for its start-up on demand

Shut-down Mode (§§4.1, 4.2)

- D2 Are the limits of operating parameters, outside which remedial action must be taken, known and measured? (CI above)
- D3 To what extent should plant be shut down for any deviation beyond the operating limits? Does this require the installation of alarm and/or trip? Should the plant be partitioned differently? How is plant restarted? (§9.6)
- D4 In an emergency, can the plant pressure and/or the inventory of process materials be reduced effectively, correctly, safely? What is the fire resistance of plant (§§9.5, 9.6)
- D5 Can the plant be shut down safely? Check the following:
- (a) See the relevant features mentioned under start-up mode (b) Fail-danger faults of protective equipment
- (c) Ingress of air, other process materials, nitrogen, steam, water, lube oil (§4.3.5)
- (d) Disposal or inactivation of residues, regeneration of catalyst, decoking, concentration of reactants, drainage, venting
- (e) Chemical, catalyst, or packing replacement, blockage removal, delivery of materials prior to start-up of plant

(f) Different modes of shutdown of plant: Normal shutdown of plant Partial shutdown of plant Placing of plant on hot standby Emergency shutdown of plant

(Wells, Safety in Process Plant Design, George Godwin, London, 1980, pp. 243-244. Paragraph references refer to this book.)

1.10. STEAM AND POWER SUPPLY

For smaller plants or for supplementary purposes, steam and power can be supplied by package plants which are shippable and ready to hook up to the process. Units with capacities in a range of sizes up to about **350,000 lb/hr** of steam are on the market, and are obtainable on a rental/purchase basis for emergency needs.

Modem steam plants are quite elaborate structures that can recover 80% or more of the heat of combustion of the fuel. The simplified sketch of Example 1.2 identifies several zones of heat transfer in the equipment. Residual heat in the flue gas is recovered as preheat of the water in an economizer and in an air preheater. The combustion chamber is lined with tubes along the floor and walls to keep the refractory cool and usually to recover more than half the heat of combustion. The tabulations of this example are of the distribution of heat transfer surfaces and the amount of heat transfer in each zone.

More realistic sketches of the cross section of a steam generator are in Figure 1.4. Part (a) of this figure illustrates the process of natural circulation of water between an upper steam drum and a lower drum provided for the accumulation and eventual **blowdown** of sediment. In some installations, pumped circulation of the water is advantageous.

Both process steam and supplemental power are recoverable from high pressure steam which is readily generated. Example 1.3 is of such a case. The high pressure steam is charged to a turbine-generator set, process steam is extracted at the desired process pressure at an intermediate point in the turbine, and the rest of the steam expands, further and is condensed.

In plants such as oil refineries that have many streams at high temperatures or high pressures, their energy can be **utilized** to generate steam or to recover power. The two cases of Example 1.4

EXAMPLE 1.2

Data of a Steam Generator for Making **250,000 lb/hr** at 450 psia and 650°F from Water Entering at **220°F**

Fuel oil of 18,500 Btu/lb is fired with 13% excess air at 80°F. Flue gas leaves at 410°F. A simplified cross section of the boiler is shown. Heat and material balances are summarized. Tube selections and arrangements for the five heat transfer zones also are summarized. The term A, is the total internal cross section of the tubes in parallel, (*Steam: Its Generation and* Use, 14.2, Babcock and Wilcox, Barberton, OH, 1972). (a) Cross section of the generator: (b) Heat balance:

Fuel input	335.5 MBtu/hr
To furnace tubes	162.0
To boiler tubes	68.5
To screen tubes	8.1
To superheater	31.3
To economizer	15.5
Total to water and steam	285.4 Mbtu/hr
In air heater	18.0 MBtu/hr

(c) Tube quantity, size, and grouping:

Screen

2 rows of $2\frac{1}{2}$ -in. OD tubes, approx 18 ft long Rows in line and spaced on 6-in. centers 23 tubes per row spaced on 6-in. centers S = 542 sqft $A_{1} = 129$ sqft



10 INTRODUCTION

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EXAMPLE 1.2—(continued)
Superheater
   12 rows of 2\frac{1}{2}-in. OD tubes (0.165-in. thick),
     17.44 ft long
  Rows in line and spaced on 3\frac{1}{4}-in. centers
  23 tubes per row spaced on 6-in. centers
  S = 3150 \text{ sqft}
  A, = 133 sqft
Boiler
  25 rows of 2\frac{1}{2}-in. OD tubes, approx 18 ft long
  Rows in line and spaced on 3\frac{1}{4}-in. centers
  35 tubes per row spaced on 4-m. centers
  s = 10,308 sqft
  A, = 85.0 sqft
Economizer
  10 rows of 2-in. OD tubes (0.148-in. thick),
     approx 10 ft long
```



- Rows in line and spaced on 3-m. centers 47 tubes per row spaced on 3-in. centers S = 2460 sqft $A_{c} = 42$ sqft Air heater 53 rows of 2-in. OD tubes (0.083-in. thick), approx 13 ft long Rows in line and spaced on $2\frac{1}{2}$ -in. centers 41 tubes per row spaced on $3\frac{1}{2}$ -in. centers S = 14,809 sqft A_{c} (total internal cross section area of 2173 tubes) = 39.3 sqft A_{c} (clear area between tubes for crossflow of air)
 - A, (clear area between tubes for crossflow of air) = 70 sqft
 - Air temperature entering air heater = 80°F



Fire 1.4. Steam boiler and furnace arrangements. [Steam, Babcock and Wilcox, Barberton, OH, 1972, pp. 3.14, 12.2 (Fig. 2), and 25.7 (Fig. 5)]. (a) Natural circulation of water in a two-drum boiler. Upper drum is for steam disengagement; the lower one for accumulation and eventual **blowdown** of sediment. (b) A two-drum boiler. Preheat tubes along the Roor and walls are connected to heaters that feed into the upper drum. (c) Cross section of a Stirling-type steam boiler with provisions for superheating, air preheating, and flue gas economizing; for maximum production of 550,000 **lb/hr** of steam at 1575 psia and 900°F.

EXAMPLE 1.3

Steam Plant Cycle for Generation of Power and Low Pressure Process Steam

The flow diagram is for the production of 5000 kW gross and 20,000 lb/hr of saturated process steam at 20 psia. The feed and hot well pumps make the net power production 4700 kW. Conditions at

key points are indicated on the enthalpy-entropy diagram. The process steam is extracted from the turbine at an intermediate point, while the rest of the stream expands to 1 in. Hg and is condensed (example is corrected from *Chemical Engineers Handbook*, 5th ed., 9.48, McGraw-Hill, New York, 1973).



EXAMPLE 1.4

Pickup of Waste Heat by Generating and Superheating Steam in a Petroleum Refinery

The two examples are generation of steam with heat from a sidestream of a fractionator in a 9000 **Bbl/day** fluid cracking plant, and superheating steam with heat from flue gases of a furnace

whose main function is to supply heat to crude topping and vacuum service in a 20,000 Bbl/day plant. (a) Recovery of heat from a sidestream of a fractionator in a 9000 Bbl/day fluid catalytic cracker by generating steam, Q = 15,950,000 Btu/hr. (b) Heat recovery by superheating steam with flue gases of a 20,000 Bbl/day crude topping and vacuum furnace.



are of steam generation in a kettle reboiler with heat from a fractionator sidestream and of steam superheating in the convection tubes of a furnace that provides heat to fractionators.

Recovery of power from the thermal energy of a high temperature stream is the subject of Example 1.5. A closed circuit of propane is the indirect means whereby the power is recovered with an expansion turbine. Recovery of power from a high pressure gas is a fairly common operation. A classic example of power recovery from a high pressure liquid is in a plant for the absorption of CO, by water at a pressure of about 4000 psig. After the absorption, the CO, is releastd and power is recovered by releasing the rich liquor through a turbine.

EXAMPLE 1.5

Recovery of Power from a Hot Gas Stream

A closed circuit of propane is employed for indirect recovery of power from the thermal energy of the hot pyrolyzate of an ethylene plant. The propane is evaporated at 500 psig, and then expanded to 100°F and 190 psig in a turbine where the power is recovered. Then the propane is condensed and pumped back to the evaporator to complete the cycle. Since expansion turbines are expensive machines even in small sizes, the process is not economical on the scale of this example, but may be on a much larger scale.



1.11. DESIGN BASIS

Before a chemical process design can be properly embarked on, a certain body of information must be agreed upon by all concerned persons, in addition to the obvious what is to be made and what it is to be made from. Distinctions may be drawn between plant expansions and wholly independent ones, so-called grassroots types. The needed data can be classified into specific design data and basic design data, for which separate check lists will be described. Specific design data include:

- 1. Required products: their compositions, amounts, purities, toxicities, temperatures, pressures, and monetary values.
- **2.** Available raw materials: their compositions, amounts, toxicities, temperatures, pressures, monetary values, and all pertinent physical properties unless they are standard and can be established from correlations. This information **about** properties applies also to products of item 1.
- **3.** Daily and seasonal variations of any data of items 1 and 2 and subsequent items of these lists.
- **4.** All available laboratory and pilot plant data on reaction and phase equilibrium behaviors, catalyst degradation, and life and corrosion of equipment.
- 5. Any available existing plant data of similar processes.
- 6. Local restrictions on means of disposal of wastes.

Basic engineering data include:

- 7. Characteristics and values of gaseous and liquid fuels that are to be used.
- **8.** Characteristics of raw makeup and cooling tower waters, temperatures, maximum allowable temperature, flow rates available, and unit costs.
- **9.** Steam and condensate: mean pressures and temperatures and their fluctuations at each level, amount available, extent of recovery of condensate, and unit costs.
- Electrical power: Voltages allowed for instruments, lighting and various driver sizes, transformer capacities, need for emergency generator, unit costs.
- 11. Compressed air: capacities and pressures of plant and instrument air, instrument air dryer.
- **12.** Plant site elevation.
- **13.** Soil bearing value, frost depth, ground water depth, piling requirements, available soil test data.

- 14. Climatic data. Winter and summer temperature extrema, cooling tower drybulb temperature, air cooler design temperature, strength and direction of prevailing winds, rain and snowfall maxima in 1 hr and in 12 hr, earthquake provision.
- **15.** Blowdown and flare: What may or may not be vented to the atmosphere or to ponds or to natural waters, nature of required liquid, and vapor relief systems.
- 16. Drainage and sewers: rainwater, oil, sanitary.
- 17. Buildings: process, pump, control instruments, special equipment.
- 18. Paving types required in different areas.
- 19. Pipe racks: elevations, grouping, coding.
- **20.** Battery limit pressures and temperatures of individual feed stocks and products.
- **21.** Codes: those governing pressure vessels, other equipment, buildings, electrical, safety, sanitation, and others.
- **22.** Miscellaneous: includes heater stacks, winterizing, insulation, steam or electrical tracing of lines, heat exchanger tubing size standardization, instrument locations.

A convenient tabular questionnaire is in Table 1.8. For anything not specified, for instance, sparing of equipment, engineering standards of the designer or constructor will be used. A proper design basis at the very beginning of a project is essential to getting a project completed and on stream expeditiously.

UTILITIES

These provide motive power and heating and cooling of process streams, and include electricity, steam, fuels, and various fluids whose changes in sensible and latent heats provide the necessary energy transfers. In every plant, the conditions of the utilities are maintained at only a few specific levels, for instance, steam at certain pressures, cooling water over certain temperature ranges, and electricity at certain voltages. At some stages of some design work, the specifications of the utilities may not have been established. Then, suitable data may be selected from the commonly **used** values itemized in Table 1.9.

1.12. LABORATORY AND PILOT PLANT WORK

The need for knowledge of basic physical properties as a factor in equipment selection or design requires no stressing. Beyond this, the state-of-the-art of design of many kinds of equipment and

I.101 Plant Location	1.107 Miscellancous Chemicals and Catalyst Supply
1.102 Plant Capacity, lb or tons/yr	In this section the operating group should outline how various miscellancous chemicals and
I. 103 Operating Factor or Yearly Operating Hours	catalysts arc to be stored and handled for consumption within the plant.
(For mos: modern chemical plants. this figure is generally 8.000 hours ${\sf per}$ year).	1.108 Atmospheric Conditions
I. 104 Provisions for Expansion	Barometric pressure range
	Temperature
1.105 Raw Material Feed (Typical of the analyses required for a liquid)	Design dry bulb temperature (°F)
Array. wt per cent min	% of summer season, this temperature is exceeded.
Impurities. wt pcr cent max	Design wet bulb temperature
Characteristic specifications	% of summer season, this temperature is exceeded.
Specific gravity	Minimum design dry bulb temperature winter condition (°F)
Distillation range °F	Level of applicable pollutants that could affect the process.
Initial boiling point 'F	Examples of these are sulfur compounds. dust and solids,
Dry end point "F	chiondes and sant water mist when the plant is at a coastal
Viscosity. centipoises	2 100 Itilities
Color APHA	2.100 Similar
Heat stability color	Characteristics of nrimary sunnly
Reaction rat (with established reagent	Voltano, nhasos curcles
Acid number	Preferred voltage for motors
Freezing point or set point °F	Ower 200 hrs
Corrosion test	Under 200 hp
Envise was	value, \$/\$ wi
such as specific density, bulk density, particle size distribution and the like are included. This physical shape information is required to assure that adequate processing and material handling operations will be provided.	load and incremental additional consumption.) 2.102 Supply Water
I.1051 Source	Cicaminicas
Supply conditions at process Max Min Normal	Corrosiveness
plant battery limits	Solids content analysis
Storage capacity (volume or day's inventory)	
Required delivery conditions at battery limits	Other Getails
Pressure	
Temperature	Pressure (at grade) Huimum Minimum
Method of transfer	Supply
1.106 Product Specifications	Return
Here again specifications would be similar 10 that of the raw material in quivalent or some- times greater detail as often trace impurities affect the marketability of the final product.	2.103 Cooling Water Well. river. sea. cooling tower, other
Storage requirements (volume or days of inventory)	Quality
Type of product storage For solid products, type of container or method of ship- ment and loading facilities should be outlined.	Value

2.104 Stam Main Normal Min High pressars, psig	Use for heat exchanger design Fouling properties Design fouling factor				2.108 Inert Gas Max Pressure, psi: Dew point, "F	Min
Line Social Construct, Fig. Image: Construct, Fig. High presents, Prig Protect CO Medium presents, Fig. Other trace improvide Medium presents, Prig Image: Construct, Fig. Temperature, Fig. Image: Construct, Fig. Construct, Fig. Image: Construct, Fig. Temperature, Fig. Image: Construct, Fig. Construct, Fig. Image: Cons	2 104 Steem	Max	Normal	Min	Per ant CO.	
Image Journer, Part (1) Protect (2) Moister, % Context transplant Moister, % Special compressor Value per thousand be Context transplant State Conceasties Special compressor Disposition Special compressor State Conceasties Special compressor Charlies and be gal Context transplant Allow per thousand be gal Context transplant Supply pressor, Pail Context transplant Total solid, pm Mu Mu Min Supply pressor, Frig Context and previous transplater moval requirements Transpersture, Fri Context and previous transplater Supply pressor, Prig </td <td>2.104 Steato</td> <td>DIEA</td> <td></td> <td></td> <td>Per cent oxygen</td> <td></td>	2.104 Steato	DIEA			Per cent oxygen	
Maisure, 7, Other trace impurities Maisure, 7, Constity setable Value per floasand is Constity setable Value per floasand is or gal Constitute, 7, Value per floasand is or gal Constitute, 7, Value per floasand is or gal Constitute, 1, Mais official content Constitute, 1, Other dealth Min Supply preasers, prise Constitute, 1, Temperature, 1, F Constitu	Temperature *F				Per cent CO	
Wate per bouand ib Quantity evalable Medium pressers, prig	Moisture. %				Other trace impurities	
Medium pressere, prig	Value Der thousand Ib				Quantity available	
Temperature, "F 2.109 Plant Åir Moisture, "A Supply Source Corp resure, pig Consistent and the construction of the constru	Medium pressure, prig				Value per thousand cu n	
Mointure, % Supply Source Value per thousand is Construction (SEL) Temperature, % Parable compressor Mointure, % Supply Source Value per thousand is Supply Source 2.105 Scan Condensts Supply Source (SSEL) Disposition Supply source (SSEL) Required pressure at lattery limits Supply pressure, pig 2.105 Scan Condensts Supply pressure, pig Disposition Supply pressure, pig Required pressure at lattery limits Supply pressure, pig Value per thousand is or gal Supply pressure, pig Disposition Supply pressure, pig Total solids, pom Supply pressure, pig Ottic datalis Ottic the advectory in the super structure, if f Total solids, pom Supply pressure Chemical sector M u Mu Min Supply pressure, TF Supply pressure of head disposal of each of these effluents Cooling water blowdown Cooling water blowdown Cooling water blowdown Cooling water blowdown Supply pressure, fig for the section of fraud effluents Cooling water blowdown Cooling water	Temperature, "F		<u> </u>		2.109 Plant Air	
Value per thousand ib	Moisture, %				Supply Source	
Low pressure, pig	Value per thousand lb				Offsite battery limits (OSBL)	
Temperature, 'F	Low pressure, psig				Portable compressor	
Moisture, %	Temperature, *F				Process air system	
Value per thousand ib Supply pressure, psig 2.105 Rectured pressure at battery limits Supply pressure, psig 2.106 Rectured pressure at battery limits Supply persersor Value per thousand ib or gal Supply pressure, psig 2.106 Rectured pressure at battery limits Supply persersor Quality Hardness, ppm It is colds, ppm Oil din and motive removal requirements Other details Oil din and motive removal requirements Other details In general, value of plant mod instrument air is usually object as the yearly over-all coarts insignificant in relation to the other utilities required. 3.101 Waste Disposal Requirements In general, value of plant mod instrument air is usually of green as the yearly over-all coarts insignificant in relation to the other utilities required. 3.101 Waste Disposal Requirements In general, value of plant mod instrument air is usually of greent. Typical items are as follows: Chemical additives M u M u Min Supply pressure Storm sever Yalue per thousand gal Chemical sever 2107 Process Water M u (If the quality of the process water is different from the make-up water or boiler feed water, visit gord of abaction of liquid effluents Yalue per thousand gal M u Yalue per thousand gal Min Supply pressure, prig Preferred material	Moisture, %				Special compressor	
2.105 Steam Condenses 2.110 Instrument Air Disposition	Value par thousand ib				Supply pressure, psig	
Disposition Required pressure at batery limits supply source (OSBL) Required pressure at batery limits Special compressor 2.106 boilsr Feed Water Supply pressure, psig Quality Hardness, ppm Hardness Supply pressure, psig Total solids, ppm Supply pressure, psig Other details Supply pressure, psig Mu Min Supply pressure, F Supply pressure, psig Yalus per thousand gal Min 2107 Process Water Mu (If the quality of the process water i different from the make-up water or boiler feed water, supply pressure, psig Storm sever Yalus per thousand gal Min Supply pressure, prig Chemical soft feed water, supply pressure, prig Quality M u Mu Min Supply pressure, prig Supply pressure, prig Temperature, i'F Supply measure, psig Temperature, i'F Supply pressure, psig Temperature, i'F Supply pressur	2.105 Steam Condensate				2.110 Instrument Air	
Required pressure at batesy limits	Disposition				supply source (OSBL)	
Value per thousand ib or gal	Required pressure at battery lim	its			Special compressor	
2.106 Boiler Feed Water Quality Hardness, ppm Total solids, ppm Total solids, ppm Other details Other details Other details Mu Mu Supply pressure Temperature, "F Value per thousand gal 2107 Process Water (If the quality of the process water is different from the make-up water or boiler feed water, separate information should be provided.) Quality Mu Supply pressure, psig Mu Temperature, "F Mu Mu Min Supply pressure, psig Mu Temperature, "F Cooling water blowdown Chemical server Storm server Storm server Storm server Storm server Storm server Storm server Facilities for treatment of gassous effecting for liquid effluents Yalue per thousand gal Storm server Storm server Storm server Storm server Facilities for treatment of gassous effluent of gassous effluent of gassous effluents Storm server Storm server Stor	Value per thousand lb or gal				Supply pressure, psig	
Quality Hardness, ppn Oii din aad mosture removal requirements silica content IO general, value of plant mod instrument air is usually not given as the yearly over-all cost is insignificant in relation to the other utilities required. Hardness Total solids, ppn Other details In general, there are three typo of wasts to be considered: liquid, solid and gaseous. The destination and disposal of each of these effluents is usually different. Typical items are as follows: Chemical additives M u M u M in Supply pressure In general, there are three typo of wasts to be considered: liquid, solid and gaseous. The destination and disposal of each of these effluents is usually different. Typical items are as follows: Cobing water blowdown Supply pressure is different from the make-up water or boiler feed water, separate information should be provided.) Quality M u Supply pressure, 'F Cooling water blowdown (If the quality of the process water is different from the make-up water or boiler feed water, spig Cooling water blowdown Supply pressure, 'F Supply pressure, 'F Storm sewer Temperature, 'F Facilities for chenical Facilities for treatment of gaseous effluents Yaius per thousand gal Facilities for treatment of gaseous effluents Facilities for treatment of gaseous effluents <	2.106 Boiler Feed Water				Dew point, "F	
Hardness, ppm IO general, value of plant mod instrument air is usually not given as the yearly over-all cost is insignificant in relation to the other utilities required. Hardness Total solids. ppm Total solids. ppm In general, value of plant mod instrument air is usually not given as the yearly over-all cost is insignificant in relation to the other utilities required. Other details In general, value of plant mod instrument air is usually not given as the yearly over-all cost is insignificant in relation to the other utilities required. Supply pressure In general, value of plant mod instrument air is usually different. Typical items are as follow: Deter details In general, value of plant mod instrument air is usually not given as the yearly over-all cost is insignificant in relation to the other utilities required. Supply pressure In general, value of plant mod instrument air is usually different. Typical items are as follow: Detained additives In general, value of plant mod instrument air is usually different. Typical items are as follow: Supply pressure In general, value of plant mod instrument air is usually different. Typical items are as follow: Cooling water blowdown Cooling water blowdown (If the quality of the provided.) M u Quality M u Min Supply pressure, psig In Min Temperature, 'F In Min	Quality				Oii din and moisture removal requirements	
Silter details 3.101 Waste Disposal Requirements Hardness Total solids. ppm Other details In general, there are three typo of waste to be considered: liquid, solid and gaseous. The destination and disposal of each of these effluents is usually different. Typical items are as follows: Chemical additives M u M u Min Supply pressure	Hardness, ppm				IO general, value of plant mod instrument air is usually not given as the ye cost is insignificant in relation to the other utilities required.	ariy over-all
Total solids. ppm	Hardness				3.101 Waste Disposal Requirements	
Other details	Total solids nom				In general, there are three typo of waste to be considered: liquid, solid and	gaseous. The
Chemical additives M u M in Destination of liquid effluents Supply pressure	Other details				destination and disposal of each of these effluents is usually different. Typical	items are as
Mu Min Cooling water blowdown Supply pressure	Chemical additives				Ionows:	
Supply pressure Temperature, 'F Value per thousand gal 2107 Process Water (If the quality of the process water is different from the make-up water or boiler feed water, separate information should be provided.) Quality Mu Supply pressure, psig Temperature, 'F Supply pressure, psig Temperature, 'F Value per thousand gal Kun Mu Min Storm sewer Storm sewer Cooling water blowdown Preferred materials of construction for Chemical sewer Cooling water blowdown Facilities for chemical Facilities for the sewer Facilities for treatment of gaseous effluents Facilities for treatment of gaseous effluents Solids disposal		м	11	Min		
Temperature, 'F Value per thousand gal 2107 Process Water (If the quality of the process water is different from the make-up water or boiler feed water, separate information should be provided.) Quality M u Mu Min Supply pressure, psig	Supply operation		_		Choning water blowdown	
Value per thousand gal 2107 Process Water (1f the quality of the process water is different from the make-up water or boiler feed water, separate information should be provided.) Quality M u Quality M u Supply pressure, psig	Temperature *F				Chemical lewer	
2107 Process Water Preferred materials of construction for (1f the quality of the process water is different from the make-up water or boiler feed water, separate information should be provided.) Mu Min Quality Mu Min Storm sever	Value per thousand sal				Storm sever	
If the quality of the process water is different from the make-up water or boiler feed water, separate information should be provided.) Cooling water blowdown Quality M u Supply pressure, psig	2107 Prome Water				Brefetted metericle of exectantics for	
(i) the provided information should be provided.) Mu Min Chemical sever Quality Mu Min Storm sever Supply pressure, psig	(If the quality of the process a	rater is different from	n the make-up water o	or boiler feed water,		
Quality M u Min Chemical struct Supply pressure, psig	separate information should be	provided.)			Chemical amor	
Supply pressure, psig	Quality	М	u	Min		
Temperature, * F	Supply pressure, psig				Becilities for chemical	
Value per thousand gal Facilities for treatment of gaseous effluents Solids disposal	Temperature, * F				treating for liquid effluents	
Solids disposal	Value per thousand gal				Facilities for treatment of gaseous effuents	
					Solids disposal	

TABLE 1.9. Typical Utility Characteristics

Pr			Steam	
	essure	(psig)	Saturation (°F)	Superheat (°F)
_	15-3	0	250-275	
	15	0	366	
	40		448	100 150
	600	J	488	100-150
_			Heat Transfer Fluids	
	۴		Fluid	d
Be	elow 6	00	petroleum oils	
Be	elow 7	50	Dowtherm and other	rs
A	bove 4	50	direct firing and elec	ctrical heating
			Refrigerants	
-	۴		Fluid	d
	40-8	0	chilled water	
	0-5	0	chilled brine and gl	ycol solutions
	-50-4	0	ammonia, freons,	butane
-	-150	-50	ethane or propane	
-	-350	-150 -300	metnane, air, nitrog	en
		-400	helium	
	Delow	-400	nenum	
Ret	urn at urn ab	110°F (s ove 125° pply at 8	Alt water) F (tempered water or Cooling Air 55-95°F	steam condensate)
	Ter Pov	nperature ver input,	approach to process, 20 HP/1000 sqft of b	40°F are surface
			Fuel	
Gas: 5-10 1000 B Liquid: at	psig, Btu/SCF 6 millio	up to 25 on Btu/b	psig for some types of arrel	burners, pipeline gas at
			Compressed Air	
		Pressure	levels of 45, 150, 300,	450 psig
			Instrument Air	

- 1.1. Process Design
- A. Books Essential to a Private Library
- 1. Ludwig, Applied Process Design for Chemical and Petroleum Plants, Gulf, Houston 1977-1983, 3 vols.
- Marks Standard Handbook for Mechanical Engineers, 9th ed., McGraw-Hill, New York, 1987.
- 3. Perry, Green, and Maloney, Perry's Chemical Engineers Handbook,

Electricity			
Driver HP Voltage			
I-100	220,440, 550		
75-250	440		
200-2500	2300, 4000		
Above 2500	4000, 13,200		

processes often demands more or less extensive pilot plant effort. This point is stressed by specialists and manufacturers of equipment who are asked to provide performance guaranties. For instance, answers to equipment suppliers' questionnaires like those of Appendix C may require the potential purchaser to have performed certain tests. Some of the more obvious areas definitely requiring test work are filtration, sedimentation, spray, or fluidized bed or any other kind of solids drying, extrusion pelleting, pneumatic and **slurry** conveying, adsorption, and others. Even in such thoroughly researched areas as vapor-liquid and liquid-liquid separations, rates, equilibria, and efficiencies may need to be tested, particularly of complex mixtures. A great deal can be found out, for instance, by a batch distillation of a complex mixture.

In some areas, suppliers make available small scale equipment that can be **used to** explore suitable ranges of operating conditions, or they may do the work themselves with benefit of their extensive experience. One engineer in the extrusion pelleting field claims that merely feeling the stuff between his fingers enables him to properly specify equipment because of his experience of 25 years with extrusion.

Suitable test procedures often are supplied with "canned" pilot plants. In general, pilot plant experimentation is a profession in itself, and the more sophistication brought to bear on it the more efficiently can the work be done. In some areas the basic relations are known so well that experimentation suffices to evaluate a few parameters in a mathematical model. This is not the book to treat the subject of experimentation, but the literature is extensive. These books may be helpful to start:

- R.E. Johnstone and M.W. Thring, Pilot Plants, Models and Scale-up Methods in Chemical Engineering, McGraw-Hill, New York, 1957.
- D.G. Jordan, *Chemical Pilot Plant Practice*, Wiley-Interscience, New York, 1955.
- 3. V. Kafarov, *Cybernetic Methods in Chemistry and Chemical* Engineering, Mir Publishers, Moscow, 1976.
- E.B. Wilson, An Introduction to Scientific Research, McGraw-Hill, New York, 1952.

McGraw-Hill, New York, 1984; earlier editions have not been obsolesced entirely.

 Sinnott, Coulson, and Richardsons, Chemical Engineering, Vof. 6, Design, Pergamon, New York, 1983.

B. Other Books

- 1. Aerstin and Street, Applied Chemical Process Design, Plenum, New York, 1978.
- Baasel, Preliminary Chemical Engineering Plant Design, Elsevier, New York, 1976.

16 INTRODUCTION

- 3. Backhurst and Harker, Process *Plant Design*, Elsevier, New York, 1973. 4. Benedek (Ed.), *Steady State Flowsheeting Of Chemical Plants*, Elsevier,
- New York, 1980. 5. Bodman, The Industrial Practice **Of** Chemical Process Engineering, MIT
- Press, Cambridge, MA, 1968.
- Branan, Process Engineers Pocket Book, Gulf, Houston, 1976, 1983, 2 vols.
- Burklin, *The Process Plant Designers Pocket Handbook of Codes and Standards*, Gulf, Houston, 1979; also, Design codes standards and recommended practices, *Encycl. Chem. Process. Des.* 14, 416-431, Dekker, New York, 1982.
- Cremer and Watkins, Chemical Engineering Practice, Butterworths, London, 1956-1965, 12 vols.
- 9. Crowe et al., *Chemical Plant Simulation*, Prentice-Hall, Englewood Cliffs, NJ, 1971.
- 10. F.L. Evans, Equipment Design Handbook for Refineries and Chemical Plants, Gulf, Houston, 1979, 2 vols.
- Franks, Modelling and Simulation in Chemical Engineering, Wiley, New York, 1972.
- 12. Institut Française du Petrole, Manual of Economic Analysis of Chemical Processes, McGraw-Hill, New York, 1981.
- Kafarov, Cybernetic Methods in Chemistry and Chemical Engineering, Mir Publishers, Moscow, 1976.
- 14. Landau (Ed.), The Chemical Plant, Reinhold, New York, 1966.
- Leesley (Ed.), Computer-Aided Process Plant Design, Gulf, Houston, 1982.
- 16. Lieberman, Process Design for Reliable Operations, Gulf, Houston, 1983.
- 17. Noel, Petroleum Refinery Manual, Reinhold, New York, 1959.
- Peters and Timmerhaus, Plant Design and Economics for Chemical Engineers, McGraw-Hill, New York, 1980.
- Rase and Barrow, Project Engineering of Process Plants, Wiley, New York, 1957.
- Resnick, Process Analysis and Design for Chemical Engineers, McGraw-Hill, New York, 1981.
- 21. Rudd and Watson, *Strategy of Process Engineering*, Wiley, New York, 1968.
- Schweitzer (Ed.), Handbook of Separation Processes for Chemical Engineers, McGraw-Hill, New York, 1979.
- Sherwood, A Course in Process Design, MIT Press, Cambridge, MA, 1963.
- 24. Ulrich, A Guide to Chemical Engineering Process Design and Economics, Wiley, New York, 1984.
- Valle-Riestra, Project Evaluation in the Chemical Process Industries, McGraw-Hill, New York, 1983.
- 26. Vilbrandt and Dryden, Chemical Engineering Plant Design, McGraw-Hill, New York, 1959.
- Wells, Process Engineering with Economic Objective, Leonard Hill, London, 1973.

C. Estimation of Properties

- AIChE Manual for Predicting Chemical Process Design Data, AIChE, New York, 1984–date.
- Bretsznajder, Prediction of Transport and Other Physical Properties of Fluids, Pergamon, New York, 1971; larger Polish edition, Warsaw, 1962.
- Lyman, Reehl, and Rosenblatt, Handbook of Chemical Property Estimation Methods: Environmental Behavior of Organic Compounds, McGraw-Hill, New York, 1982.
- Reid, Prausnitz, and Poling, *The Properties of Gases and Liquids*, McGraw-Hill, New York, 1987.
- Sterbacek, Biskup, and Tausk, Calculation of Properties Using Corresponding States Methods, Elsevier, New York, 1979.
- S.M. Walas, Phase Equilibria in Chemical Engineering, Butterworths, Stoneham, MA, 1984.

D. Equipment

- 1. Chemical Engineering Catalog, Penton/Reinhold, New York, annual.
- 2. Chemical Engineering Equipment Buyers' Guide, McGraw-Hill, New York, annual.
- 3. Kieser, Handbuch der chemisch-technischen Apparate, Spamer-Springer, Berlin, 1934-1939.

- Mead, The Encyclopedia of Chemical Process Equipment, Reinhold, New York, 1964.
- 5. Riegel, Chemical Process Machinery, Reinhold, New York, 1953.
- 6. Thomas Register of American Manufacturers, Thomas, Springfield IL, annual.

E. Safety Aspects

- 1. Fawcett and Wood (Eds.), Safety and Accident Prevention in Chemical Operations, Wiley, New York, 1982.
- 2. Lees, Loss Prevention in the Process Industries, Buttenvorths, London, 1980, 2 vols.
- 3. Lieberman, Troubleshooting Refinery Processes, PennWell, Tulsa, 1981.
- Lund, Industrial Pollution Control Handbook, McGraw-Hill, New York, 1971.
- Rosaler and Rice, Standard Handbook of Plant Engineering, McGraw-Hill, New York, 1983.
- Sax, Dangerous Properties of Industrial Materials, Van Nostrand/ Reinhold, New York, 1982.
- Wells, Safety in Process Plant Design, George Godwin, Wiley, New York, 1980.
- 1.2. Process Equipment

A. Encyclopedias

- 1. Considine, *Chemical and Process Technology Encyclopedia*, McGraw-Hill, New York, 1974.
- Kirk-Othmer Concise Encyclopedia of Chemical Technology, Wiley, New York, 1985.
- Kirk-Othmer Encyclopedia of Chemical Technology, Wiley, New York, 1978-1984, 26 vols.
- McGraw-Hill Encyclopedia of Science and Technology, 5th ed., McGraw-Hill, New York, 1982.
- McKetta and Cunningham (Eds.), Encyclopedia of Chemical Processing and Design, Dekker, New York, 1976–date.
- Ullmann, Encyclopedia of Chemical Technology, Verlag Chemie, Weinheim, FRG, German edition 1972-1983; English edition 1984-1994(?).

B. Bibliographies

- Fratzcher, Picht, and Bittrich, The acquisition, collection and tabulation of substance data on fluid systems for calculations in chemical engineering, Znt. Chem. Eng. 20(1), 19-28 (1980).
- 2. Maizell, How to Find Chemical Information, Wiley, New York, 1978.
- Mellon, Chemical Publications: Their Nature and Use, McGraw-Hill, New York, 1982.
- Rasmussen and Fredenslund, Data Banks for Chemical Engineers, Kemiigeniorgruppen, Lyngby, Denmark, 1980.

C. General Data Collections

- 1. American Petroleum Institute, Technical Data Book-Petroleum Refining, API, Washington, DC, 1971-date.
- Bolz and N. Tuve, Handbook of Tables for Applied Engineering Science, CRC Press, Washington, DC, 1972.
- 3. CRC Handbook of Chemistry and Physics, CRC Press, Washington, DC, annual.
- Gallant, Physical Properties of Hydrocarbons, Gulf, Houston, 1968, 2 vols.
- 5. International Critical Tables, McGraw-Hill, New York, 1926-1933.
- Landolt-Börnstein, Numerical Data and Functional Relationships in Science and Technology, Springer, New York, 1950-date.
- 7. Lange's Handbook of Chemistry, 13th ed., McGraw-Hill, New York, 1984.
- 8. Maxwell, Data Book on Hydrocarbons, Van Nostrand, New York, 1950.
- Melnik and Melnikov, *Technology of Inorganic Compounds*, Israel Program for Scientific Translations, Jerusalem, 1970.
- National Gas Processors Association, *Engineering Data Book*, Tulsa, 1987.
- 11. Perry's Chemical Engineers Handbook, McGraw-Hill, New York, 1984.
- 12. Physico-Chemical Properties for Chemical Engineering, Maruzen Co., Tokyo, 1977-date.

- 13. Raznjevic, Handbook of Thermodynamics Tables and Charts (SI Units), Hemisphere, New York, 1976.
- Vargaftik, Handbook of Physical Properties of Liquids and Gases, Hemisphere, New York, 1983.
- Yaws et al., Physical and Thermodynamic Properties, McGraw-Hill, New York, 1976.
- D. Special Data Collections
- 1. Gmehling et al., Vapor-Liquid Equilibrium Data Collection, DECHEMA, Frankfurt/Main, FRG, 1977-date.
- Hirata, Ohe, and Nagahama, Computer-Aided Data Book of Vapor-Liquid Equilibria, Elsevier, New York, 1976.
- Keenan et al., Steam Tables, Wiley, New York, English Units, 1969, SI Units, 1978.
- Kehiaian, Selected Data on Mixtures, International Data Series A: Thermodynamic Properties of Non-reacting Binary Systems of Organic Substances, Texas A & M Thermodynamics Research Center, College Station, TX, 1977–date.

- Kogan, Fridman, and Kafarov, Equilibria between Liquid and Vapor (in Russian), Moscow, 1966.
- Larkin, Selected Data on Mixtures, International Data Series' B, Thermodynamic Properties of Organic Aqueous Systems, Engineering Science Data Unit Ltd, London, 197%date.
- Ogorodnikov, Lesteva, and Kogan, Handbook of Azeotropic Mixtures (in Russian), Moscow, 1971; data of 21,069 systems.
- Ohe, Computer-Aided Data Book of Vapor Pressure, Data Publishing Co., Tokyo, 1976.
- 9. Sorensen and Arlt, Liquid-Liquid Equilibrium Data Collection, DECHEMA, Frankfurt/Main, FRG, 1979-1980, 3 vols.
- Starling, Fluid Thermodynamic Properties for Light Petroleum Systems, Gulf, Houston, 1973.
- 11. Stephen, Stephen and Silcock, *Solubilities of Inorganic and Organic Compounds*, Pergamon, New York, 1979, 7 vols.
- 12. Stull, Westrum, and Sinke, The Chemical Thermodynamics of Organic Compounds, Wiley, New York, 1969.
- 13. Wagman et al., The NBS Tables of Chemical Thermodynamic Properties: Selected Values for Inorganic and C₁ and C₂ Organic Substances in SI Units, American Chemical Society, Washington, DC, 1982.


plant design is made up of words, numbers, and pictures. An engineer thinks naturally in terms of the sketches and drawings which are his "pictures." Thus, to solve a material balance problem, he will start with a block to represent the equipment and then will show entering and leaving streams with their amounts and properties. Or **ask** him to describe a process and he will begin to sketch the equipment, show how iris interconnected, and what the flows and operating conditions are.

Such sketches develop into flow sheets, which are more

2.1. BLOCK FLOWSHEETS

At an early stage or to provide an overview of a complex process or plant, a drawing is made with rectangular blocks to represent individual processes or groups of operations, together with quantities and other pertinent properties of key streams between the blocks and into and from the process as a whole. Such block flowsheets are made at the beginning of a process design for orientation purposes or later as a summary of the material balance of the process. For example, the coal carbonization process of Figure 2.1 starts with 100,000 lb/hr of coal and some process air, involves six main process units, and makes the indicated quantities of ten different products. When it is of particular interest, amounts of utilities also may be shown; in this example the use of steam is indicated at one point. The block diagram of Figure 2.2 was prepared in connection with a study of the modification of an existing petroleum refinery. The three feed stocks are separated into more than 20 products. Another representative petroleum refinery block diagram, in Figure 13.20, identifies the various streams but not their amounts or conditions.

2.2. PROCESS FLOWSHEETS

Process flowsheets embody the material and energy balances between and the sizing of the major equipment of the plant. They include all vessels such as reactors, separators, and drums; special processing equipment, heat exchangers, pumps, and so on. Numerical data include flow quantities, compositions, pressures, temperatures, and so on. Inclusion of major instrumentation that is essential to process control and to complete understanding of the flowsheet without reference to other information is required particularly during the early stages of a job, since the process flowsheet is drawn first and is for some time the only diagram representing the process. As the design develops and a mechanical flowsheet gets underway, instrumentation may be taken off the process diagram to reduce the clutter. A checklist of the information that usually is included on a process flowsheet is given in Table 2.1.

Working flowsheets are necessarily elaborate and difficult to represent on the page of a book. Figure 2.3 originally was 30in. wide. In this process, ammonia is made from available hydrogen supplemented by hydrogen from the air oxidation of natural gas in a two-stage reactor F-3 and V-S. A large part of the plant is devoted to purification of the feed gases of carbon dioxide and unconverted methane before they enter the converter CV-1. Both commercial and refrigeration grade ammonia are made in this plant. Compositions of 13 key streams are summarized in the tabulation.

elaborate diagrammatic representations of the equipment, the sequence of operations, and the expected performance of a proposed p/ant or the actual performance of an already operating one. For clarity and to meet the needs of the various persons engaged in design, cost estimating, purchasing, fabrication, operation, maintenance, and management, several different kinds of flowsheets are necessary. Four of the main kinds will be described and illustrated.

Characteristics of the streams such as temperature, pressure, enthalpy, volumetric flow rates, etc., sometimes are conveniently included in the tabulation. In the interest of clarity, however, in some instances it may be preferable to have a separate sheet for a voluminous material balance and related stream information.

A process flowsheet of the dealkylation of toluene to benzene is in Figure 2.4; the material and enthalpy flows and temperature and pressures are tabulated conveniently, and basic instrumentation is represented.

2.3. MECHANICAL (P&I) FLOWSHEETS

Mechanical flowsheets also are called piping and instrument (P&I) diagrams to emphasize two of their major characteristics. They do not show operating conditions or compositions or flow quantities, but they do show all major as well as minor equipment more realistically than on the process flowsheet. Included are sizes and specification classes of all pipe lines, all valves, and all instruments. In fact, every mechanical aspect of the plant regarding the process equipment and their interconnections is represented except for supporting structures and foundations. The equipment is shown in greater detail than on the PFS, notably with regard to external piping connections, internal details, and resemblance to the actual appearance.

The mechanical flowsheet of the reaction section of a toluene dealkylation unit in Figure 2.5 shows all instrumentation, including indicators and transmitters. The clutter on the diagram is minimized by tabulating the design and operating conditions of the major equipment below the diagram.

The P&I diagram of Figure 2.6 represents a gas treating plant that consists of an amine absorber and a regenerator and their immediate auxiliaries. Internals of the towers are shown with exact locations of inlet and outlet connections. The amount of instrumentation for such a comparatively simple process may be surprising. On a completely finished diagram, every line will carry a code designation identifying the size, the kind of fluid handled, the pressure rating, and material specification. Complete information about each line-its length, size, elevation, pressure drop, fittings, etc.-is recorded in a separate line summary. On Figure 2.5, which is of an early stage of construction, only the sizes of the lines are shown. Although instrumentation symbols are fairly well standardized, they are often tabulated on the P&I diagram as in this example.

2.4. UTILITY FLOWSHEETS

These are P&I diagrams for individual utilities such as steam, steam condensate, cooling water, heat transfer media in general,



Figure 2.1. Coal carbonization block flowsheet. Quantities are in lb/hr.

compressed air, fuel, refrigerants, and inert blanketing gases, and how they are piped up to the process equipment. Connections for utility streams are shown on the mechanical flowsheet, and their conditions and flow quantities usually appear on the process flowsheet.

Since every detail of a plant design must be recorded on paper, many other kinds of drawings also are required: for example, electrical flow, piping isometrics, instrument lines, plans and elevations, and individual equipment drawings in all detail. Models and three-dimensional representations by computers also are now standard practice in many design offices.

2.5. DRAWING OF FLOWSHEETS

Flowsheets are intended to represent and explain processes. To make them easy to understand, they are constructed with a consistent set of symbols for equipment, piping, and operating conditions. At present there is no generally accepted industrywide body of drafting standards, although every large engineering office does have its internal standards. Some information appears in ANSI and British Standards publications, particularly of piping symbols. Much of this information is provided in the book by Austin (1979) along with symbols gleaned from the literature and some engineering firms. Useful compilations appear in some books on process design, for instance, those of Sinnott (1983) and Ulrich (1984). The many flowsheets that appear in periodicals such as *Chemical Engineering* or *Hydrocarbon Processing* employ fairly consistent sets of symbols that may be worth imitating.

Equipment symbols are a compromise between a schematic representation of the equipment and simplicity and ease of drawing. A selection for the more common kinds of equipment appears in Table 2.2. Less common equipment or any with especially intricate configuration often is represented simply by a circle or rectangle.

Since a symbol does not usually speak entirely for itself but also carries a name and a letter-number identification, the flowsheet can be made clear even with the roughest of equipment symbols. The

TABLE 2.1. Checklist of Data Normally Included on a Process Flowsheet

- 1. Process lines, but including only those bypasses essential to an understanding of the process
- 2. All process equipment. Spares are indicated by letter symbols or notes
- 3. Major instrumentation essential to process control and to understanding of the flowsheet
- 4. Valves essential to an understanding of the flowsheet
- 5. Design basis, including stream factor
- 6. Temperatures, pressures, flow quantities
- 7. Weight and/or mol balance, showing compositions, amounts, and other properties of the principal streams
- 6. Utilities requirements summary
- 9. Data included for particular equipment
 - a. Compressors: SCFM (60°F. 14.7 psia); APpsi; HHP; number of stages; details of stages if important
 - b. Drives: type; connected HP; utilities such as kW, Ib steam/hr, or Btu/hr
 - c. Drums and tanks: ID or OD, seam to seam length, important internals
 - d. Exchangers: Sqft, kBtu/hr, temperatures, and flow quantities in and out; shell side and tube side indicated
 - e. Furnaces: kBtu/hr, temperatures in and out, fuel
 - f. Pumps: GPM (60°F), APpsi, HHP, type, drive
 - g. Towers: Number and type of plates or height and type of packing; identification of all plates at which streams enter or leave; ID or OD; seam to seam length; skirt height
 - h. Other equipment: Sufficient data for identification of duty and size

TABLE 2.2. Flowsheet Equipment Symbols



22 FLOWSHEETS

TABLE 2.2-(continued)



letter-number designation consists of a letter or combination to designate the class of the equipment and a number to distinguish it from others of the same class, as two heat exchangers by E-112 and E-215. Table 2.4 is a typical set of letter designations.

Operating conditions such as flow rate, temperature, pressure,

enthalpy, heat transfer rate, and also stream numbers are identified with symbols called flags, of which Table 2.3 is a commonly used set. Particular units are identified on each flowsheet, as in Figure 2.3.

Letter designations and symbols for instrumentation have been

TABLE 2.2—(continued)

Convevors and Feeders

Separators



thoroughly standardized by the Instrument Society of America (ISA). An abbreviated set that may be adequate for the usual flowsketch appears on Figure 3.4. The P&I diagram of Figure 2.6 affords many examples.

For clarity and for esthetic reasons, equipment should be represented with some indication of their relative sizes. True scale is not feasible because, for example, a flowsheet may need to depict both a tower 15Oft high and a drum 2ft in diameter. Logarithmic

TABLE 2.2—(continued)



scaling sometimes gives a pleasing effect; for example, if the 150 ft tower is drawn 6in. high and the 2ft drum 0.5 in., other sizes can be read off a straight line on log-log paper.

A good draftsman will arrange his flowsheet as artistically as possible, consistent with clarity, logic, and economy of space on the drawing. A fundamental rule is that there be no large gaps. Flow is predominantly from left to right. On a process flowsheet, distillation towers, furnaces, reactors, and large vertical vessels often are arranged at one level, condenser and accumulator drums on another level, reboilers on still another level, and pumps more or less on one level but sometimes near the equipment they serve in order to minimize excessive crossing of lines. Streams enter the flowsheet from the left edge and leave at the right edge. Stream numbers are assigned to key process lines. Stream compositions and other desired properties are gathered into a table that may be on a separate sheet if it is especially elaborate. A listing of flags with the units is desirable on the flowsheet.

Rather less freedom is allowed in the construction of mechanical flowsheets. The relative elevations and sizes of equipment are preserved as much as possible, but all pumps usually are shown at the same level near the bottom of the drawing. Tabulations of instrumentation symbols or of control valve sizes or of relief valve sizes also often appear on P&I diagrams. Engineering offices have elaborate checklists of information that should be included on the flowsheet, but such information is beyond the scope here.

Appendix 2.1 provides the reader with material for the construction of flowsheets with the symbols of this chapter and possibly with some reference to Chapter 3.

Mass flow rate, lbs/hr	13,028
Molal flow rate, lbmols/hr	217
Temperature, °F	510
Pressure, psig (or indicate if psia or Torr or bar)	195 psia
Volumetric liquid flow rate, gal/min.	65.3
Volumetric liquid flow rate, bblS/day	8,500
Kilo Btu/hr, at heat transfer equipment	9,700
Enthalpy, Btu/Ib	953
Others	\frown

TABLE 2.3. Flowsheet Flags of Operating Conditions inTypicalUnits

TABLE 2.4. Letter Designations of Equipment

Equipment Letters		Equipment	Letters	
Agitator	М	Grinder	SR	
Air filter	FG	Heat exchanger	E	
Bin	l-r	Homogenizer	М	
Blender	М	Kettle	R	
Blower	JB	Kiln (rotary)	DD	
Centrifuge	FF	Materials handling	G	
Classifying equipment	S	equipment		
Colloid mill	SR	Miscellaneous"	L	
Compressor	JC	Mixer	М	
Condenser	E	Motor	PM	
Conveyor	C	Oven	В	
Cooling tower	TE	Packaging machinery	L	
Crusher	SR	Precipitator (dust or mist)	FG	
Crystallizer	K	Prime mover	PM	
Cyclone separator (gas)	FG	Pulverizer	SR	
Cyclone separator		Pump (liquid)	J	
(liquid)	F	Reboiler	E	
Decanter	FL	Reactor	R	
Disperser	М	Refrigeration system	G	
Drum	D	Rotameter	RM	
Dryer (thermal)	DE	Screen	S	
Dust collector	FG	Separator (entrainment)	FG	
Elevator	C	Shaker	М	
Electrostatic separator	FG	Spray disk	SR	
Engine	PM	Spray nozzle	SR	
Evaporator	FE	Tank	l-r	
Fan	JJ	Thickener	F	
Feeder	C	Tower	т	
Filter (liquid)	Р	Vacuum equipment	V E	
Furnace	В	Weigh scale	L	

'Note: The letter L is used for unclassified equipment when only a few items are of this type; otherwise, individual letter designations are assigned.



Fii 2.2. Block flowsheet of the revamp of a 30,000 Bbl/day refinery with supplementary light stocks (The C. W. Nofsinger Co.).



Figure 2.3. Process flowsheet of a plant making 47 tons/day of ammonia from available hydrogen and hydrogen made from natural gas (The C. W. Nofsinger Co.).



Figure 2.5. Engineering (P&I) flowsheet of the reaction section of plant for dealkylation of benzene (*Wells*, Safety in Process Design, *George Godwin*, *London*, 1980).







Figure 2.6. Engineering flowsheet of a gas treating plant. Note the tabulation of instrumentation flags at upper right (Fluor Engineers, by way of Ruse and Barrow, Project Engineering of Process Plants, Wiley, New York, 1957).

REFERENCES 31

REFERENCES

- D.G. Austin, Chemical Engineering Drawing Symbols, George Godwin, London, 1979.
- 2. Graphical Symbols for Piping System and Plant, British Standard 1553: Part 1: 1977.
- 3. Graphical Symbols for Process Flow Diagrams, **ASA** Y32.11.1961, American Society of Mechanical Engineers, New York.
- 4. E.E. Ludwig, Applied Process Design for Chemical and Petrochemical

Plants, Gulf, Houston, 1977, Vol. 1.

- H.F. Rase and M.H. Barrow, Project Engineering of Process Plants, Wiley, New York, 1957.
- R.K. Sinnott, Coulson, and Richardson, Chemical Engineering, vol. 6, Design, Pergamon, New York, 1983.
- 7. G.D. Ulrich, A Guide to Chemical Engineering Process Design and Economics, Wiley, New York, 1984.
- 8. R. Weaver, Process Piping Design, Gulf, Houston, 1973, 2 vols.

Appendix 2.1 –

Descriptions of Example Process Flowsheets

These examples ask for the construction of **flowsheets** from the given process descriptions. Necessary auxiliaries such as drums and pumps are to be included even when they are not mentioned. Essential control instrumentation also is to be provided. Chapter 3 has examples. The processes are as follows:

- 1. visbreaker operation,
- 2. cracking of gas oil,
- 3. olefin production from naptha and gas oil,
- 4. propylene oxide synthesis,
- 5. phenol by the chlorobenzene process,
- 6. manufacture of butadiene sulfone,
- 7. detergent manufacture,
- 8. natural gas absorption,
- 9. tall oil distillation,
- 10. recovery of isoprene,
- 11. vacuum distillation,
- 12. air separation.

1. VISBREAKER OPERATION

Visbreaking is a mild thermal pyrolysis of heavy petroleum fractions whose object is to reduce fuel production in a refinery and to make some gasoline.

The oil of 7.2 API and 700°F is supplied from beyond the battery limits to a surge drum F-l. From there it is pumped with J-1A&B to parallel furnaces B-1A&B from which it comes out at 890°F and 200 psig. Each of the split streams enters at the bottom of its own evaporator T-1A&B that has five trays. Overheads from the evaporators combine and enter at the bottom of a 30-tray fractionator T-2. A portion of the bottoms from the fractionator is fed to the top trays of T-IA&B; the remainder goes through exchanger E-5 and is pumped with J-2A&B back to the furnaces B-1A&B. The bottoms of the evaporators are pumped with J-4A&B through exchangers E-5, E-3A (on crude), and E-3B (on cooling water) before proceeding to storage as the fuel product.

A side stream is withdrawn at the tenth tray from the top of T-2 and proceeds to steam stripper T-3 equipped with five trays. Steam is fed below the bottom tray. The combined steam and oil vapors return to T-2 at the eighth tray. Stripper bottoms are pumped with J-6 through E-2A (on crude) and E-2B (on cooling water) and to storage as "heavy gasoline."

Overhead of the fractionator T-2 is partially condensed in E-1A (on crude) and E-1B (on cooling water). A gas product is withdrawn overhead of the reflux drum which operates at 15 psig. The "light gasoline" is pumped with J-5 to storage and as reflux.

Oil feed is 122,480 pph, gas is 3370, light gasoline is 5470, heavy gasoline is 9940, and fuel oil is 103,708 pph.

Include suitable control equipment for the main fractionator T-2.

2. CRACKING OF GAS OIL

A gas oil cracking plant consists of two cracking furnaces, a soaker, a main fractionator, and auxiliary strippers, exchangers, pumps, and drums. The main fractionator (150 psig) consists of four zones, the bottom zone being no. 1.

A light vacuum gas oil (LVGO) is charged to the top plate of zone 3, removed from the bottom tray of this zone and pumped to furnace no. 1 that operates at 1000 psig and 1000°F. A heavy vacuum gas oil (HVGO) is charged to the top plate of zone 2, removed at the bottom tray and charged to furnace no. 2 that operates at 500 psig and 925°F.

Effluents from both furnaces are combined and enter the soaker; this is a large vertical drum designed to provide additional residence time for conversion under adiabatic conditions. Effluent at **500 psig** and 915°F enters the bottom zone of the main fractionator.

Bottoms from zone 1 goes to a stripping column (5 psig). Overhead from that tower is condensed, returned partly as reflux and partly to zone 3 after being cooled in the first condenser of the stripping column. This condensing train consists of the preheater for the stream being returned to the main fractionator and an air cooler. The cracked residuum from the bottom of the stripper is cooled to 170°F in a steam generator and an air cooler in series. Live steam is introduced below the bottom tray for stripping.

All of the oil from the bottom of zone 3 (at 700° F), other than the portion that serves as feed to furnace no. 1, is withdrawn through a cooler (500°F) and pumped partly to the top tray of zone 2 and partly as spray quench to zone 1. Some of the bottoms of zone 1 likewise is pumped through a filter and an exchanger and to the same spray nozzle.

Part of the liquid from the bottom tray of zone 4 (at 590°F) is pumped to a hydrogenation unit beyond the battery limits. Some light material is returned at 400°F from the hydrogenation unit to the middle of zone 4, together with some steam.

Overhead from the top of the column (zone 4) goes to a partial condenser at 400°F. Part of the condensate is returned to the top tray as reflux; the rest of it is product naphtha and proceeds beyond the battery limits. The uncondensed gas also goes beyond the battery limits. Condensed water is **sewered**.

3. OLEFIN PRODUCTION

A gaseous product rich in ethylene and propylene is made by pyrolysis of crude oil fractions according to the following description. Construct a flowsheet for the process. Use standard symbols for equipment and operating conditions. Space the symbols and proportion them in such a way that the sketch will have a pleasing appearance.

Crude oil is pumped from storage through a steam heated exchanger and into an electric desalter. Dilute caustic is injected into the line just before the desalting drum. The aqueous phase collects at the bottom of this vessel and is drained away to the sewer. The oil leaves the desalter at 190°F, and goes through heat exchanger E-2 and into a furnace coil. From the furnace, which it leaves at 600°F, the oil proceeds to a distillation tower.

After serving to preheat the feed in exchanger E-2, the bottoms proceeds to storage; no bottoms pump is necessary because the tower operates with 65 psig at the top. A gas oil is taken off as a sidestream some distance above the feed plate, and naphtha is taken off overhead. Part of the overhead is returned as reflux to the tower, and the remainder proceeds to a cracking furnace. The gas oil also is charged to the same cracking furnace but into a separate coil. Superheated steam at 800°F is injected into both cracking coils at their inlets.

Effluents from the naphtha and gas oil cracking coils are at 1300° F and 1200° F, respectively. They are combined in the line just before discharge into a quench tower that operates at 5 psig and 235° F at the top. Water is sprayed into the top of this tower. The

bottoms is pumped to storage. The overhead is cooled in a water exchanger and proceeds to a separating drum. Condensed water and an aromatic oil separate out there. The water is **sewered** whereas the oil is sent to another part of the plant for further treating.

The uncondensed gas from the separator is compressed to **300 psig** in a reciprocating unit of three stages and then cooled to 100°F. Condensed water and more aromatic distillate separate out. Then the gas is dried in a system of two desiccant-filled vessels that are used alternately for drying and regeneration.

Subsequently the gas is **precooled** in exchanger E-6 and charged to a low temperature fractionator. This tower has a reboiler and a top refluxing system. At the top the conditions are 280psig and -75° F. Freon refrigerant at -90° F is **used** in the condenser. The bottoms is recycled to the pyrolysis coil. The uncondensed vapor leaving the reflux accumulator constitutes the product of this plant. It is used to **precool** the feed to the fractionator in E-6 and then leaves this part of the plant for further purification.

4. PROPYLENE OXIDE SYNTHESIS

Draw a process flowsheet for the manufacture of propylene oxide according to the following description.

Propylene oxide in the amount of 5000 **tons/yr** will be made by the chlorohydrin process. The basic feed material is a hydrocarbon mixture containing 90% propylene and the balance propane which does not react. This material is diluted with spent gas from the process to provide a net feed to chlorination which contains **40 mol** % propylene. Chlorine gas contains 3% each of air and carbon dioxide as contaminants.

Chlorination is accomplished in a packed tower in which the hydrocarbon steam is contacted with a saturated aqueous solution of chlorine. The chlorine solution is made in another packed tower. Because of the limited solubility of chlorine, chlorohydrin solution from the chlorinator is recirculated through the solution tower at a rate high enough to supplement the fresh water needed for the process. **Solubility** of chlorine in the chlorohydrin solution is approximately the same as in fresh water.

Concentration of the effluent from the chlorinator is 8 lb organics/100 lb of water. The organics have the composition

Propylene	chlorohydrin	75 mol	%
Propylene	dichloride	19	
Propionaldeh	6		

Operating pressure of the chlorinator is 30 psig, and the temperature is 125°F . Water and the fresh gas stream are at 80°F . Heat of reaction is 2000 Btu/lb chlorine reacted. Percentage conversion of total propylene fed to the chlorinator is 95% (including the recycled material).

Overhead from the chlorinator is scrubbed to remove excess chlorine in two vessels in succession which employ water and 5% caustic solution, respectively. The water from the first scrubber is used in the chlorine solution tower. The caustic is recirculated in order to provide adequate wetting of the packing in the caustic scrubber; fresh material is charged in at the same rate as spent material is purged. Following the second scrubber, propylene dichloride is recovered from the gas by chilling it. The spent gas is recycled to the chlorinator in the required amount, and the excess is flared.

Chlorohydrin solution is pumped from the chlorinator to the saponitier. It is mixed in the feed line with a 10% lime slurry and preheated by injection of live 25 psig steam to a temperature of 200°F. Stripping steam is injected at the bottom of the saponifier, which has six perforated trays without downcomers. Propylene

oxide and other organic materials go overhead; the bottoms contain unreacted lime, water, and some other reaction products, all of which can be dumped. Operating pressure is substantially atmospheric. Bubblepoint of the overhead is 60°F.

Separation of the oxide and the organic byproducts is accomplished by distillation in two towers. Feed from the saponifier contains oxide, aldehyde, dichloride, and water. In the first tower, oxide and aldehyde go overhead together with only small amounts of the other substances; the dichloride and water go to the bottom and also contain small amounts of contaminants. Two phases will form in the lower section of this tower; this is taken off as a partial side stream and separated into a dichloride phase which is sent to storage and a water phase which is sent to the saponifier as recycle near the top of that vessel. The bottoms are a waste product. Tower pressure is 20psig. Live steam provides heat at the bottom of this column.

Overhead from the first fractionator is condensed and charged to the second tower. There substantially pure propylene oxide is taken overhead. The bottoms is dumped. Tower pressure is 15 psig, and the overhead bubblepoint is 100°F. Reactions are

 $\begin{array}{c} Cl_2 + H_2O \rightarrow CIOH + \text{HCI} \\ C_3H_6 + Cl_2 + H_2O \rightarrow C_3H_6CIOH + \text{HCI} \\ C_3H_6 + Cl_2 & \rightarrow C_3H_6Cl_2 \\ C_3H_6CIOH & \rightarrow C_2H_5CHO + \text{HCI} \end{array} \xrightarrow{2} 2C_3H_6O + CaCl_2 + 2H_2O$

Show all necessary major equipment, pumps, compressors, refrigerant lines. Show the major instrumentation required to make this process continuous and automatic.

5. PHENOL BY THE CHLOROBENZENE PROCESS

A, portion of a plant for the manufacture of phenol from **mono**chlorbenzene and **NaOH** is in accordance with the following description.

- a. Construct a flowsheet of the process, with operating conditions and the two control instruments mentioned.
- b. Prepare a material balance showing the compositions of the process streams in the portion of the plant before the brine decanter V-103. The amount of phenol in this stream is 2000 lb/hr. Excess caustic (5%) is fed to the emulsifier.

Process description: The principal reactions in the plant are

$\begin{array}{ll} C_6H_5Cl + 2NaOH \rightarrow C_6H_5ONa + NaCl + H_2O & 2C_6H_5OH \\ C_6H_5ONa + HCI \rightarrow C_6H_5OH + NaCl & \downarrow \rightarrow (C_6H_5)_2O + H_2O \end{array}$

From storage, monochlorbenzene and 10% caustic are pumped together with diphenyl ether from decanter V-102 into emulsifier V-101 which is provided with intense agitation. The effluent from that vessel is pumped with a high pressure steam driven reciprocating pump P-103 at **4000 psig** through a feed-effluent exchanger E-101 and through the tube side of a direct fired heater R-101. Here the stream is heated to 700°F and reaction 1 occurs.

From the reactor, the effluent is cooled in E-101, cooled further to **110°F** in water cooler E-102, and then enters diphenyl ether decanter V-102. The lighter DPE phase is returned with pump P-104 to the emulsifier. The other phase is pumped with P-105 to another stirred vessel R-102 called a Springer to which 5% HCl also is pumped, with P-106; here reaction 2 occurs.

The mixture of two liquid phases is cooled in water cooler E-103 and then separated in brine decanter V-103. From that vessel the lighter phenol phase proceeds (P-108) to a basket type evaporator D-101 that is heated with steam. Overhead vapor from

the evaporator proceeds beyond the battery limit for further purification. Evaporator bottoms proceeds to waste disposal. The aqueous phase from decanter V-103 is pumped with P-109 through a feed-bottoms exchanger E-104 to the top tray of the brine tower D-102. The overhead is condensed in E-105, collected in accumulator V-104 and pumped beyond the battery limits for recovery of the phenol. Tower D-102 is provided with a steam heated **reboiler** E-106. Bottom product is a weak brine that is pumped with P-110 through the feed-bottoms exchanger and beyond the battery limits for recovery of the salt.

Two important control instruments are to be shown on the flowsheet. These are a back pressure controller in the reactor effluent line beyond exchanger E-101 and a pH controller on the feed line of the 5% HCl that is fed to springer R-102. The pH instrument maintains proper conditions in the springer.

Note: There is a tendency to byproduct diphenyl ether formation in reactor R-101. However, a recycle of 100 pph of DPE in the feed to the reactor prevents any further formation of this substance.

6. MANUFACTURE OF BUTADIENE SULFONE

A plant is to manufacture butadiene sulfone at the rate of 1250 lb/hr from liquid sulfur dioxide and butadiene to be recovered from a crude C_4 mixture as starting materials. Construct a flowsheet for the process according to the following description.

The crude C_4 mixture is charged to a 70 tray extractive distillation column T-l that employs acetonitrile as solvent. Trays are numbered from the bottom. Feed enters on tray 20, solvent enters on tray 60, and reflux is returned to the top tray. Net overhead product goes beyond the battery limits. Butadiene dissolved in acetonitrile leaves at the bottom. This stream is pumped to a 25-tray solvent recovery column T-2 which it enters on tray 20. Butadiene is recovered overhead as liquid and proceeds to the BDS reactor. Acetonitrile is the bottom product which is cooled to 100°F and returned to T-l. Both columns have the usual condensing and reboiling provisions.

Butadiene from the recovery plant, liquid sulfur dioxide from storage, and a recycle stream (also liquified) are pumped through a preheater to a high temperature reactor R-l which is of shell-and-tube construction with cooling water on the shell side. Operating conditions are 100°C and 300 psig. The combined feed contains equimolal proportions of the reactants, and 80% conversion is attained in this vessel. The effluent is cooled to 70°C, then enters a low temperature reactor R-2 (maintained at 70°C and 50psig with cooling water) where the conversion becomes 92%. The effluent is flashed at 70°C and atmospheric pressure in D-1. Vapor product is compressed, condensed and recycled to the reactor R-1. The liquid is pumped to a storage tank where 24 hr holdup at 70°C is provided to ensure chemical equilibrium between sulfur dioxide, butadiene, and butadiene sulfone. Cooling water is available at 32°C.

7. DETERGENT MANUFACTURE

The process of making synthetic detergents consists of several operations that will be described consecutively.

ALKYLATION

Toluene and olefinic stock from storage are pumped (at 80° F) separately through individual driers and filters into the alkylation reactor. The streams combine just before they enter the reactor. The reactor is batch operated 4 hr/cycle; it is equipped with a single impeller agitator and a feed hopper for solid aluminum chloride which is charged manually from small drums. The alkylation

mixture is pumped during the course of the reaction through an external heat exchanger (entering at -10° F and leaving at -15° F) which is cooled with ammonia refrigerant (at -25° F) from an absorption refrigeration system (this may be represented by a block on the FS); the exchanger is of the kettle type. HCl gas is injected into the recirculating stream just beyond the exit from the heat exchanger; it is supplied from a cylinder mounted in a weigh scale. The aluminum chloride forms an alkylation complex with the toluene. When the reaction is complete, this complex is pumped away from the reactor into a storage tank with a complex transfer pump. To a certain extent, this complex is reused; it is injected with its pump into the reactor recirculation line before the suction to the recirculation pump. There is a steam heater in the complex line, between the reactor and the complex pump.

The reaction mixture is pumped away from the reactor with an alkymer transfer pump, through a steam heater and an orifice mixer into the alkymer wash and surge tank. Dilute caustic solution is recirculated from the a.w.s. tank through the orifice mixer. Makeup of caustic is from a dilute caustic storage tank. Spent caustic is intermittently drained off to the sewer. The a.w.s. tank has an internal weir. The caustic solution settles and is removed at the left of the weir; the alkymer overflows the weir and is stored in the right-hand portion of the tank until amount sufficient for charging the still has accumulated.

DISTILLATION

Separation of the reactor product is effected in a ten-plate batch distillation column equipped with a water-cooled condenser and a Dowtherm-heated (650°F, 53psig) still. During a portion of the distillation cycle, operation is under vacuum, which is produced by a two-stage steam jet ejector equipped with barometric condensers. The Dowtherm heating system may be represented by a block. Product receiver drums are supplied individually for a slop cut, for toluene, light alkymer, heart alkymer, and a heavy alkymer distillate. Tar is drained from the still at the end of the operation through a water cooler into a bottoms receiver drum which is supplied with a steam coil. From this receiver, the tar is loaded at intervals into 50 gal drums, which are trucked away. In addition to the drums which serve to receive the distillation products during the operation of the column, storage tanks are provided for all except the slop cut which is returned to the still by means of the still feed pump; this pump transfers the mixture from the alkymer wash and surge tank into the still. The recycle toluene is not stored with the fresh toluene but has its own storage tank. The heavy alkymer distillate tank connects to the olefinic stock feed pump and is recycled to the reactor.

SULFONATION

Heart alkymer from storage and 100% sulfuric acid from the sulfuric acid system (which can be represented by a block) are pumped by the reactor feed pump through the sulfonation reactor. The feed pump is a positive displacement proportioning device with a single driver but with separate heads for the two fluids. The reactor is operated continuously; it has a single shell with three stages which are partially separated from each other with horizontal doughnut shaped plates. Each zone is agitated with its individual impeller; all three impellers are mounted on a single shaft. On leaving the reactor, the sulfonation mixture goes by gravity through a water cooler (leaving at 130°F) into a centrifuge. Spent acid from the centrifuge goes to storage (in the sulfuric acid system block); the sulfonic acids go to a small surge drum or can bypass this drum and go directly to a large surge tank which is equipped with an agitator and a steam jacket. From the surge drum, the material is sent by an extraction feed pump through a water cooler, then a "flomix," then another water cooler, then another "flomix" (leaving at 150°F), and then through a centrifuge and into the sulfonic acid surge tank. Fresh water is also fed to each of the "flomixers." Wash acid is rejected by the centrifuge and is sent to the sulfuric acid system. The "flomix" is a small vertical vessel which has two compartments and an agitator with a separate impeller for each compartment.

NEUTRALIZATION

Neutralization of the sulfonic acid and building up with sodium sulfate and tetrasodium pyrophosphate (TSPP) is accomplished in two batch reactors (5 hr cycle) operated alternately. The sodium sulfate is pumped in solution with its transfer pump from the sodium sulfate system (which can be represented by a block). The TSPP is supplied as a solid and is fed by means of a Redler conveyor which discharges into a weigh hopper running on a track above the two reactors. Each reactor is agitated with a propeller and a turbine blade in a single shaft.

Sodium hydroxide of 50% and 1% concentrations is used for neutralization. The 50% solution discharges by gravity into the reactor; the 1% solution is injected gradually into the suction side of the reactor slurry circulating pump. As the caustic is added to the reactor, the contents are recirculated through a water-cooled external heat exchanger (exit at 160°F), which is common to both reactors. When the reaction is completed in one vessel, the product is fed gradually by means of a slurry transfer pump to two double drum dryers which are steam-heated and are supplied with individual vapor hoods. The dry material is carried away from the dryers on a belt conveyor and is taken to a flaker equipped with an air classifier. The fines are returned to the trough between the dryer drums. From the classifier, the material is taken with another belt conveyor to four storage bins. These storage bins in turn discharge onto a belt feeder which discharges into drums which are weighed automatically on a live portion of a roller conveyor. The roller convevor takes the drums to storage and shipping.

Notes: All water cooled exchangers operate with water in at 75°F and out at 100°F. All pumps are centrifugal except the complex transfer, and the sulfonation reactor feed, which are both piston type; the neutralization reactor recirculation pump and the transfer pumps are gear pumps.

Show all storage tanks mentioned in the text.

8. NATURAL GAS ABSORPTION

A gas mixture has the composition by volume:

Component	N_2	СН₄	C₂H₀	C ₃ H ₈
Mot fraction	0.05	0.65	0.20	0.10

It is fed to an absorber where 75% of the propane is recovered. The total amount absorbed is 50 mol/hr. The absorber has four theoretical plates and operates at 135 psig and 100°F. All of the absorbed material is recovered in a steam stripper that has a large number of plates and operates at 25 psig and 230°F.

Water is condensed out of the stripped gas at 100°F. After compression to 50 psig, that gas is combined with a recycle stream. The mixture is diluted with an equal volume of steam and charged to a reactor where pyrolysis of the propane occurs at a temperature of 1300°F. For present purposes the reaction may be assumed to be simply $C_3H_8 \rightarrow C_2H_4 + CH$, with a specific rate k = 0.28/sec. Conversion of propane is 60%. Pressure drop in the reactor is 20 psi.

Reactor effluent is cooled to remove the steam, compressed to 285 **psig**, passed through an activated alumina drying system to remove further amounts of water, and then fed to the first fractionator. In that vessel, 95% of the unconverted propane is recovered as a bottoms product. This stream also contains 3%

ethane as an impurity. It is throttled to 50psig and recycled to the reactor. In two subsequent towers, ethylene is separated from light and heavy impurities. Those separations may be taken as complete.

Construct a flow diagram of this plant. Show such auxiliary equipment as drums, heat exchangers, pumps, and compressors. Show operating conditions and flow quantities where calculable with the given data.

9. TALL OIL DISTILLATION

Tall oil is a byproduct obtained from the manufacture of paper pulp from pine trees. It is separated by vacuum distillation (50mmHg) in the presence of steam into four primary products. In the order of decreasing volatility these are unsaponifiables (US), fatty acid (FA), rosin acids (RA), and pitch (P). Heat exchangers and reboilers are heated with **Dowtherm** condensing vapors. Some coolers operate with water and others generate steam. Live steam is charged to the inlet of every reboiler along with the process material. Trays are numbered from the bottom of each tower.

Tall oil is pumped from storage through a preheater onto tray 10 of the pitch stripper T-l. Liquid is withdrawn from tray 7 and pumped through a reboiler where partial vaporization occurs in the presence of steam. The bottom 6 trays are smaller in diameter and serve as stripping trays. Steam is fed below tray 1. Pitch is pumped from the bottom through steam generator and to storage. Overhead vapors are condensed in two units E-l and E-2. From the accumulator, condensate is pumped partly as reflux to tray 15 and partly through condenser E-l where it is preheated on its way as feed to the next tower T-2. Steam is not condenser that is connected to a steam jet ejector.

Feed enters T-2 at tray 5. There is a pump-through reboiler. Another pump withdraws material from the bottom and sends it to tower T-3. Liquid is pumped from tray 18 through a cooler and returned in part to the top tray 20 for temperature and reflux control. A portion of this pumparound is withdrawn after cooling as unsaps product. Steam leaves the top of the tower and is condensed in the barometric.

Tray 5 of T-3 is the feed position. This tower has two reboilers. One of them is a pumparound from the bottom, and the other is gravity feed from the bottom tray. Another pump withdraws material from the bottom, and then sends it through a steam generator and to storage as rosin acid product. A slop cut is withdrawn from tray 20 and pumped through a cooler to storage. Fatty acid product is pumped from tray 40 through a cooler to storage. Another stream is pumped around from tray 48 to the top tray 50 through a cooler. A portion of the cooled pumparound is sent to storage as another unsaps product. A portion of the overhead steam proceeds to the barometric condenser. The rest of it is boosted in pressure with high pressure steam in a jet compressor. The boosted steam is fed to the inlets of the two reboilers associated with T-3 and also directly into the column below the bottom tray.

The vapors leaving the primary barometric condenser proceed to a steam ejector that is followed by another barometric. Pressures at the tops of the towers are maintained at 50mmHg absolute. Pressure drop is 2 mm Hg per tray. Bottom temperatures of the three towers are 450, 500, and 540°F, respectively. Tower overhead temperatures are 200°F. Pitch and rosin go to storage at 350°F and the other products at 125°F. The steam generated in the pitch and rosin coolers is at 20 psig. Process steam is at 150 psig.

10. RECOVERY OF ISOPRENE

Draw carefully a flowsheet for the recovery of isoprene from a mixture of C_5 hydrocarbons by extractive distillation with aqueous acetonitrile according to the following description.

A hydrocarbon stream containing 60 mol % isoprene is charged at the rate of 10,000pph to the main fractionator D-l at tray 40 from the top. The solvent is acetonitrile with **10 wt** % water; it is charged at the rate of 70,000 pph on tray 11 of D-l. This column has a total of 70 trays, operates at **10 psig** and 100°F at the top and about 220°F at the bottom. It has the usual provisions for reboiling and top **reflux**.

The extract is pumped from the bottom of D-l to a stripper D-2 with 35 trays. The stripped solvent is cooled with water and returned to D-l. An isoprene-acetonitrile azeotrope goes overhead, condenses, and is partly returned as top tray reflux. The net overhead proceeds to an extract wash column D-3 with 20 trays where the solvent is recovered by countercurrent washing with water. The overhead from D-3 is the finished product isoprene. The bottoms is combined with the bottoms from the raffinate wash column D-4 (20 trays) and sent to the solvent recovery column D-5 with 15 trays.

Overhead from D-l is called the raffinate. It is washed countercurrently with water in D-4 for the recovery of the solvent, and then proceeds beyond the battery limits for further conversion to isoprene. Both wash columns operate at substantially atmospheric pressure and 100°F. The product streams are delivered to the battery limits at 100 psig.

Solvent recovery column D-5 is operated at 50 mm Hg absolute, so as to avoid the formation of an azeotrope overhead. The required overhead condensing temperature of about 55° F is provided with a propane compression refrigeration system; suction condition is 40°F and 80 psig, and discharge condition is 200 psig. Vacuum is maintained on the reflux accumulator with a two-stage steam ejector, with a surface interstage condenser and a direct water spray after-condenser. The stripped bottoms of D-5 is cooled to 100°F and returned to the wash columns. Some water makeup is necessary because of leakages and losses to process streams. The solvent recovered overhead in D-5 is returned to the main column D-l. Solvent makeup of about 20 pph is needed because of losses in the system.

Steam is adequate for all reboiling needs in this plant.

11. VACUUM DISTILLATION

This plant is for the distillation of a heavy petroleum oil. The principal equipment is a vacuum tower with 12 trays. The top tray is numbered 1. Trays 1, 2, 10, 11, and 12 are one-half the diameter of the other trays. The tower operates at 50 mm Hg.

Oil is charged with pump J-l through an exchanger E-l, through a fired heater from which it proceeds at 800°F onto tray 10 of the tower. Live steam is fed below the bottom tray.

Bottoms product is removed with pump J-3 through a steam

generator and a water cooled exchanger E-3 beyond the battery limits. A side stream is taken off tray 6, pumped with J-2 through E-1, and returned onto tray 3 of the tower. Another stream is removed from tray 2 with pump J-4 and cooled in water exchanger E-2; part of this stream is returned to tray 1, and the rest of it leaves the plant as product gas oil.

Uncondensed vapors are removed at the top of the column with a one-stage steam jet ejector equipped with a barometric condenser.

Show the principal controls required to make this plant operate automatically.

12. AIR SEPARATION

Make a flowsheet of an air purification and separation plant that operates according to the following description.

Atmospheric air at the rate of 6.1 million SCFD is compressed to 160 psig in a two-stage compressor JJ-1 that is provided with an intercooler and a knockout drum. Then it proceeds to a packed tower T-1 where it is scrubbed with recirculating caustic soda solution. Overhead from T-1 is cooled to 14°F in a refrigerated exchanger. After removal of the condensate, this stream proceeds to a dryer system that consists principally of two vessels F-1 and F-2 packed with solid desiccant.

After being **precooled** with product oxygen in exchanger E-1 and with product nitrogen in E-2, the air serves as the heating medium in reboiler E-3 of column T-2. Its pressure then is reduced to 100 psig, and it is fed to the middle of column T-2. Bottoms of T-2 is fed to the middle of column T-3. This stream contains 40% oxygen.

Columns T-3 and T-4 operate at 15 and **30 psig**, respectively. Column T-3 is located above T-4. Elevations and pressure differentials are maintained in such a way that no liquid pumps are needed in the distillation section of the plant.

Part of the overhead from T-2 (containing 96% nitrogen) is condensed in E-4 which is the reboiler for column T-3, and the remainder is condensed in E-5 which is the reboiler for T-4. Part of the condensate from E-4 is returned as reflux to T-2 and the rest of the condensates from E-4 and E-5 serve as top reflux to T-3. Overhead from T-3 contains 99.5% nitrogen. After **precooling** the feed in E-2, this nitrogen proceeds to the battery limits.

Bottoms of T-3 proceeds to the top of stripper T-4. Vapor overhead from T-4 is recycled to the middle of T-3. The bottoms product (containing 99.5% oxygen) is sent partly to liquid storage and the remainder to precooler E-1 where it is vaporized. Then it is compressed to 150psig in a two-stage compressor JJ-2 and sent to the battery limits. Compressor JJ-2 has inter- and aftercoolers and knockout drums for condensate.

PROCESS CONTROL

If processes are subject to disturbances that tend to change operating conditions, compositions, and physical properties of the streams. In order to minimize the ill effects that could result from such disturbances, chemical plants are implemented with substantial amounts of instrumentation and automatic control equipment. In critical cases and in especially large p/ants, moreover, the instrumentation is computer monitored for convenience, safety, and optimization.

for example, a typical billion **|b/yr** ethylene p/ant may have 600 control loops with control valves and 400 interacting loops with a cost of about \$6 million. (Skrokov, 1980, pp. 13, 49; see Sec. 3.1); the computer implementation of this control system Will cost another \$3 million. Figure 3.7 shows the control system of an ethylene fractionator which has 12 input signals to the computer and four outgoing reset signals to flow controllers.

In order for a process to be controllable by machine, it must represented by a mathematical mode/. Ideally, each element of a dynamic process, for example, a reflux drum or an individual trav of a fractionator, is represented by differential equations based on material and energy balances, transfer rates, stage efficiencies, phase equilibrium relations, etc., as we// as the parameters of sensing devices, control valves, and control instruments. The process as a who/e then is equivalent to a system of ordinary and partial differential equations involving certain independent and dependent variables. When the values of the independent variables are specified or measured, corresponding values of the others are found by computation, and the information is transmitted to the control instruments. For example, if the temperature, composition, and flow rate of the feed to a fractionator are perturbed, the computer Will determine the other flows and the heat balance required to maintain constant overhead purity. Economic factors also can be incorporated in process mode/s; then the computer can be made to optimize the operation continua//y.

For control purposes, somewhat simplified mathematical mode/s usually are adequate. In distillation, for instance, the Underwood-Fenske-Gil/i/and mode/ with constant relative volatilities and a simplified enthalpy balance may be preferred to a full-fledged tray-by-tray calculation every time there is a perturbation. In control situations, the demand for speed of response may not be realizable with an over/y elaborate mathematical system. Moreover, in practice not all disturbances are measurable, and the process characteristics are not known exactly. According/y feedforward control is supplemented in most instances with feedback. In a we//designed system (Shinskey, 1984, p. 186) typically 90% of the corrective action is provided by feed forward and 10% by feedback with the result that the integrated error is reduced by a factor of 10.

A major feature of many modern control systems is composition control which has become possible with the development of fast and accurate on-line analyzers. Figure 3.2 shows that 10 analyzers are used for control of ethylene composition in this p/ant within the purities shown, High speed on-line gas chromatographs have analysis times of 30- 120 sec and are capable of measuring several components simultaneously with a sensitivity in the parts/million range. Mass spectrometers are faster, more stable, and easier to maintain but are not sensitive in the ppm range. Any one instrument can be hooked up to a half-dozen or so sample ports, but, of course, at the expense of time lag for controller response. Infrared and NMR spectrometers also are feasible for on-line analysis. Less costly but also less specific analyzers are available for measuring physical properties such as refractive index and others that have been calibrated against mixture composition or product purity.

The development of a mathematical model, even a simplified one that is feasible for control purposes, takes a major effort and is we// beyond the scope of the brief treatment of process control that can be attempted here. What will be given is examples of control loops for the common kinds of equipment and operations, Primarily these are feedback arrangements, but, as mentioned earlier, feedback devices usually are necessary supplements in primarily feedforward situations.

When processes are subject on/y to slow and small perturbations, conventional feedback PID controllers usually are adequate with set points and instrument characteristics fine-tuned in the field. As an example, two modes of control of a heat exchange process are shown in Figure 3.8 where the objective is to maintain constant out/et temperature by exchanging process heat with a heat transfer medium. Part (a) has a feedback controller which goes into action when a deviation from the preset temperature occurs and attempts to restore the set point. Inevitably some oscillation of the outlet temperature will be generated that will persist for some time and may never die down if perturbations of the in/et condition occur often enough. In the operation of the feedforward control of part (b), the flow rate and temperature of the process input are continua//v signalled to a computer which then finds the flow rate of heat transfer medium required to maintain constant process outlet temperature and adjusts the flow control valve appropriate/y. Temperature oscillation amplitude and duration will be much less in this mode.

3.1. FEEDBACK CONTROL

In feedback control, after an offset of the controlled variable from a preset value has been generated, the controller acts to eliminate or reduce the offset. Usually there is produced an oscillation in the **value** of the controlled variable whose amplitude, period, damping and permanent offset depend on the nature of the system and the

mode of action of the controller. The usual controllers provide one, two, or three of these modes of corrective action:

- 1. Proportional, in which the corrective action is proportional to the error signal.
- 2. Integral, in which the corrective action at time *t* is proportional to the integral of the error up to that time.



Figure 3.1. Optimized control of an ethylene tower (Skrokov (Ed.), Mini- and Microcomputer Control in Industrial Processes, Van Nostrand/Reinhold, New York, 1980).



Figure 3.2. Plowsketch of an olefins plant and specifications of the ethylene product. AR designates a composition analyzer and controller (*after Skrokov (Ed.)*, Mini- and Microcomputer Control in Industrial Processes, *Van Nostrand/Reinhold*, *New York*, 1980).

3. Derivative, in which the corrective action is proportional to the rate at which the error is being generated.

The relation between the change in output $m - m_0$ and input e signals accordingly is represented by

$$m - m_0 = K_p \left(e + \frac{1}{K_i} \int_0^t e \, dt + K_d \, \frac{de}{dt} \right).$$

Just how these modes of action are achieved in relatively inexpensive pneumatic or electrical devices is explained in books on control instruments, for example, that of Considine (Process Instruments and Controls Handbook, Sec. 17, 1974). The low prices and considerable flexibility of PID controllers make them the dominant types in use, and have discouraged the development of possibly superior types, particularly as one-shot deals which would be the usual case in process plants. Any desired mode of action can be simulated by a computer, but at a price.

A capsule summary of the merits of the three kinds of corrective action can be made. The proportional action is rapid but has a permanent offset that increases as the action speeds up. The addition of integral action reduces or entirely eliminates the offset but has a more sluggish response. The further addition of derivative action speeds up the correction. The action of a three-mode PID controller can be made rapid and without offset. These effects are illustrated in Figure 3.3 for a process subjected to a unit step upset, in this case a change in the pressure of the control air. The ordinate is the ratio of the displacements of the response and upset from the set point.

The reason for a permanent offset with a proportional controller can be explained with an example. Suppose the temperature of a reactor is being controlled with a pneumatic system. At the set point, say the valve is 50% open and the flow rate

	Curve	Mode of Control	Prop Sensitivity, K P	Integral Time, K	Derív Time, K _d	Period of Cycle, sec	$\begin{array}{c} \text{Damping} \\ c^{-t/T} \\ T, \\ \text{sec} \end{array}$	Max Error units	Offset units
	1	Proportional derivative	16	-	0.9	32	15	0.18	0.05
	2	Prop int deriv	10	22	2.0	44	20	0.21	0
	3	Proportional	8	-	-	45	20	0.29	0.11
	4	Proportional integral	4	3	-	66	30	0.37	0
	5	Integral	-	35	-	210	100	0.69	0
0.5 0.4									
).2'	ĥ	2 3				λ			
).1			K			-	X		
0.1				-1					
0) 10	20 30	40 50	60	70	80	90 1	100 1	10 1

Figure 3.3. Response of various modes of control to step input (Eckman, Automatic Process Control, Wiley, New York, 1958).

of cooling water is fixed accordingly. Suppose the heat load is doubled suddenly because of an increase in the reactor contents. At steady state the valve will remain 50% open so that the water flow rate also will remain as before. Because of the greater rate of heat evolution, however, the temperature will rise to a higher but still steady value. On the other hand, the corrective action of an integral controller depends on displacement of the temperature from the original set point, so that this mode of control will restore the original temperature.

The constants K_p , K_i , and K_d are settings of the instrument. When the controller is hooked up to the process, the settings appropriate to a desired quality of control depend on the inertia (capacitance) and various response times of the system, and they can be determined by field tests. The method of Ziegler and Nichols used in Example 3.1 is based on step response of a damped system and provides at least approximate values of instrument settings which can be further fine-tuned in the field.

The kinds of controllers suitable for the common variables may be stated briefly:

Variable	Controller			
Flow and liquid pressure	PI			
Gas pressure	Р			
Liquid level	P or PI			
Temperature	PID			
Composition	P. PI. PID			

Derivative control is sensitive to noise that is made up of random higher frequency perturbations, such as splashing and turbulence generated by inflow in the case of liquid level control in a vessel, so that it is not satisfactory in such situations. The variety of composition controllers arises because of the variety of composition analyzers or detectors.

Many corrective actions ultimately adjust a flow rate, for instance, temperature control by adjusting the flow of a heat transfer medium or pressure by regulating the flow of an effluent stream. A control unit thus consists of a detector, for example, a thermocouple, a transmitter, the control instrument itself, and a control valve. The natures, sensitivities, response speeds, and locations of these devices, together with the inertia or capacity of the process equipment, comprise the body of what is to be taken into account when designing the control system. In the following pages will be described only general characteristics of the major kinds of control systems that are being used in process plants. Details and criteria for choice between possible alternates must be sought elsewhere. The practical aspects of this subject are treated, for example, in the References at the end of this chapter.

SYMBOLS

On working flowsheets the detectors, transmitters, and controllers are identified individually by appropriate letters and serial numbers in circles. Control valves are identified by the letters CV- followed by a serial number. When the intent is to show only in general the kind of control system, no special symbol is used for detectors, but simply a point of contact of the signal line with the equipment or process line. Transmitters are devices that convert the measured variable into air pressure for pneumatic controllers or units appropriate for electrical controllers. Temperature, for instance, may be detected with thermocouples or electrical resistance or height of a liquid column or radiant flux, etc., but the controller can accept only pneumatic or electrical signals depending on its type. When the nature of the transmitter is clear, it may be represented by an encircled cross or left out entirely. For clarity, the flowsheet can include only the most essential information. In an actual design

EXAMPLE 3.1 Constants of PID Controllers from Response Curves to a Step Input

The method of Ziegler and Nichols [*Trans ASME*, (Dec. 1941)] will be used. The example is that of Tyner and May (Process *Engineering Control*, Ronald, New York, 1967). The response to a change of 2 psi on the diaphragm of the control valve is shown. The full range of control pressure is from 3 to 15 psi, a difference of 12 psi, and the range of temperature is from 100 to 200°F, a difference of $100^{\circ}F$. Evaluate the % displacement of pressure as

Am = 100(2/12) = 16.7%.

From the curve, the slope at the inflection point is

R = 17.5/100(7.8 - 2.4) = 3.24%/min,

and the apparent time delay is the intercept on the abscissa,

L = 2.40 min.

The values of the constants for the several kinds of controllers are *Proportional:* $100/K_p = \%$ PB = $100RL/\Delta m = 100(3.24)(2.4)/$

16.7 = 46.6%.

Proportional-integral: % PB = $110RL/\Delta m = 51.2\%$

 $K_i = L/0.3 = 8 \min$

case, details of detectors and transmitters as well as all other elements of a control system are summarized on instrument specification forms. The simplified coding used in this chapter is summarized on Figure 3.4.

CASCADE (RESET) CONTROL

Some control situations require interacting controllers. On Figure 3.19(d), for instance, a composition controller regulates the **setpoint** of the temperature controller of a reactor and on Figure 3.15(g) the set point of the reflux flow rate is regulated by composition or temperature control. Composite systems made up of regions that respond with varying degrees of speed or sluggishness are advantageously equipped with cascade control. In the reactor of Figure 3.19(b), the temperature **TT-1** of the vessel contents responds only slowly to changes in flow rate of the heat transfer medium, but the temperature **TT-2** of the HTM leaving the cooling coil is comparatively sensitive to the flow rate. Accordingly, controller TC-2 is allowed to adjust the **setpoint** of the primary controller TC-1 with an overall improvement in control of the reactor temperature. The controller being reset is identified on flowsheets.

3.2. INDIVIDUAL PROCESS VARIABLES

The variables that need to be controlled in chemical processing are temperature, pressure, liquid level, **flow** rate, flow ratio, composition, and certain physical properties whose magnitudes may be influenced by some of the other variables, for instance, viscosity, vapor pressure, refractive index, etc. When the temperature and pressure are fixed, such properties are measures of composition which may be known exactly upon calibration. Examples of control Proportional-integral-derivative:

% PB =
$$83RL/\Delta m$$
 = 38.6%,
 $K_i = 2L = 4.8 \text{ min},$
 $K_d = 0.5L = 1.2 \text{ min}.$

These are approximate instrument settings, and may need to be adjusted in process. PB is proportional band.

A recent improvement of the Ziegler-Nichols method due to Yuwana and Seborg [AIChE J. 28, 434 (1982)] is calculator programmed by Jutan and Rodriguez [Chem. Eng. 91(18), 69-73 (Sep. 3, 1984)].



of individual variables are shown in the rest of this chapter with the various equipment (say pumps or compressors) and processes (say distillation or refrigeration) and on the earlier flowsketches of this and the preceding chapters, but some general statements also can be made here. Most control actions ultimately depend on regulation of a flow rate with a valve.

TEMPERATURE

Temperature is regulated by heat exchange with a heat transfer medium (HTM). The **flow** rate of the HTM may be adjusted, or the condensing pressure of steam or other vapor, or the amount of heat transfer surface exposed to condensing vapor may be regulated by flooding with condensate, which always has a much lower heat transfer coefficient than that of condensing vapor. In a reacting system of appropriate vapor pressure, a boiling temperature at some desired value can be maintained by **refluxing** at the proper controlled pressure. Although examples of temperature control appear throughout this chapter, the main emphasis is in the section on heat exchangers.

PRESSURE

Pressure is controlled by regulating the flow of effluent from the vessel. The effluent may be the process stream itself or a **non**-condensable gas that is generated by the system or supplied for blanketing purposes. The system also may be made to float on the pressure of the blanketing gas supply. Control of the rate of condensation of the effluent by allowing the heat transfer surface to flood partially is a common method of regulating pressure in fractionation systems. Throttling a main effluent vapor line usually is not done because of the expense of large control valves. Figure 3.5 shows vacuum production and control with steam jet ejectors.





LEVEL OF LIQUID

Level of liquid in a vessel often is maintained by permanent or adjustable built-in weirs for the effluent, notably on the trays of fractionators, extractors, etc., and in reactors and drums. Any desired adjustments of weir height, however, can be made only on shutdown. Control of the flow rate of effluent (sometimes of the input) is the most common other method of level control. Liquid levels often are disturbed by splashing or flow turbulence, so that rather sluggish controllers are used for this service. Conceivably, a level could be controlled by forcing effluent through an opening of fixed size with a controlled pressure, but there do not appear to be many such applications. Continual control of the weight of a vessel and its contents is another control method that is not used often. Figure 3.6 is devoted to level control.

FLOW RATE

A rate of flow is commonly measured by differential pressure across an orifice, but many other devices also are used on occasion. Simultaneous measurements of temperature and pressure allow the flow measurement to be known in mass units. Direct mass flow meters also are available. The flow measurement is transmitted to a controller which then adjusts the opening of a control valve so as to maintain the desired condition.

FLOW OF SOLIDS

Except for continuous weighing, control of the flow of solids is less precise than that of fluids. Several devices used for control of feed rates are shown schematically in Figure 3.7. They all employ variable speed drives and are individually calibrated to relate speed and flow rate. Ordinarily these devices are in effect manually set, but if the solid material is being fed to a reactor, some property of the mixture could be used for feed back control. The continuous belt weigher is capable ordinarily of $\pm 1\%$ accuracy and even $\pm 0.1\%$ when necessary. For processes such as neutralizations with lime, addition of the solid to process in slurry form is acceptable. The slurry is prepared as a batch of definite concentration and charged with a pump under flow control, often with a diaphragm pump whose stroke can be put under feedback control. For some applications it is adequate or necessary to feed weighed amounts of solids to a process on a timed basis.

FLOW RATIO

Flow ratio control is essential in processes such as fuel-air mixing, blending, and reactor feed systems. In a two-stream process, for example, each stream will have its own controller, but the signal from the primary controller will go to a ratio control device which adjusts the set point of the other controller. Figure 3.17(a) is an example. Construction of the ratioing device may be an adjustable mechanical linkage or may be entirely pneumatic or electronic. In other two-stream operations, the flow rate of the secondary stream may be controlled by some property of the combined stream, temperature in the case of fuel-air systems or composition or some physical property indicative of the proportions of the two streams.

COMPOSITION

The most common detectors of specific substances are gas chromatographs and mass spectrometers, which have been mentioned earlier in this chapter in connection with feedforward control. Also mentioned have been physical properties that have been calibrated against mixture compositions. Devices that are specific for individual substances also are sometimes available, for example pH, oxygen, and combustion products. Impregnated reactive tapes have been made as specific detectors for many substances and are useful particularly for low concentrations. Composition controllers act by adjusting some other condition of the system: for instance, the residence time in converters by adjusting the flow rate, or the temperature by adjusting the flow of HTM, or the pressure of gaseous reactants, or the circulation rate of regenerable catalysts, and so on. The taking of representative samples is an aspect of on-line analysis that slows down the responsiveness of such control. The application of continuously measuring in-line analyzers is highly desirable. Some physical properties can be measured this way, and also concentrations of hydrogen and many other ions with suitable electrodes. Composition controllers are shown for the processes of Figures 3.1 and 3.2.

3.3. EQUIPMENT CONTROL

Examples are presented of some usual control methods for the more widely occurring equipment in chemical processing plants. Other methods often are possible and may be preferable because of



Figure 3.5. Vacuum control with steam jet ejectors and with mechanical vacuum pumps. (a) Air bleed on PC. The steam and water rates are hand set. The air bleed can be made as small as desired. This can be used only if air is not harmful to the process. Air bleed also can be used with mechanical vacuum pumps. (b) Both the steam and water supplies are on automatic control. This achieves the minimum cost of utilities, but the valves and controls are relatively expensive. (c) Throttling of process gas flow. The valve is larger and more expensive even than the vapor valve of case (a). Butterfly valves are suitable. This method also is suitable with mechanical vacuum pumps. (d) No direct pressure control. Settings of manual control valves for the utilities with guidance from pressure indicator PI. Commonly used where the greatest vacuum attainable with the existing equipment is desired.



Figure 3.6. Some modes of control of liquid level. (a) Level control by regulation of the effluent flow rate. This mode is externally adjustable. (b) Level control with built in overflow weir. The weir may be adjustable, but usually only during shutdown of the equipment. (c) Overflow weir in a horizontal kettle reboiler. The weir setting usually is permanent.

greater sensitivity or lower cost. Also it should be noted that the choice of controls for particular equipment may depend on the kind of equipment it is associated with. Only a few examples are shown of feedforward control, which should always be considered when superior control is needed, the higher cost is justified, and the process simulation is known. Another relatively expensive method is composition control, which has not been emphasized here except for reactors and fractionators, but its possible utility always should be borne in mind. Only primary controllers are shown. The complete instrumentation of a plant also includes detectors and transmitters as well as indicators of various operating conditions. Such indications may be input to a computer for the record or for control, or serve as guides for manual control by operators who have not been entirely obsolesced.

HEAT TRANSFER EQUIPMENT

Four classes of this kind of equipment are considered: heat exchangers without phase change, steam heaters, condensers, and vaporizers or reboilers. These are grouped together with descriptions in Figures 3.8-3.11. Where applicable, comments are made about the utility of the particular method. In these heat



Figure 3.7. Solids feeders with variable speed drives. (a) Rotary vane (star) feeder with variable speed drive. (b) Horizontal screw feeder. (c) Belt feeder taking material from a bin with an adjustable underflow weir. (d) Rotary plate feeder: Rate of discharge is controlled by the rotation speed, height of the collar, and the position of the plow. (e) Continuously weighing feeder with variable speed belt conveyor.

Figure 3.8. Heat exchangers without phase change. PF = processfluid, HTM = heat transfer medium. (a) Feedback control of PF outlet temperature. Flow rate of HTM is adjusted as the PF outlet temperature is perturbed. The valve may be in either the input or output line. (b) Feedforward control. PF outlet setpoint T-2 and perturbations of PF input flow and temperature are fed to the monitor which adjusts the flow rate of the HTM to maintain constant PF outlet temperature T2. (c) Exchanger with bypass of process fluid with a three-way valve. The purpose of TC-2 is to conserve on that fluid or to limit its temperature. When the inherent leakage of the three-way valve is objectionable, the more expensive two two-way valves in the positions shown are operated off TC-1. (d) A two-fluid heat transfer system. The PF is heated with the HTM which is a closed circuit heated by Dowtherm or combustion gases. The Dowtherm is on flow control acting off TC-2 which is on the HTM circuit and is reset by TC-1 on the PF outlet. The HTM also is on flow control. Smoother control is achievable this way than with direct heat transfer from very high temperature Dowtherm or combustion gases. (e) Air cooler. Air flow rate is controllable with adjustable louvers or variable pitch fan or variable speed motors. The latter two methods achieve some saving of power compared with the louver design. Multispeed motors are also used for change between day and night and between winter and summer. The switching can be made automatically off the air temperature.





Figure 3.9. Steam heaters. (a) Flow of steam is controlled off the PF outlet temperature, and condensate is removed with a steam trap or under liquid level control. Subject to difficulties when condensation pressure is below atmospheric. (b) Temperature control on the condensate removal has the effect of varying the amount of flooding of the heat transfer surface and hence the rate of condensation. Because the flow of condensate through the valve is relatively slow, this mode of control is sluggish compared with (a). However, the liquid valve is cheaper than the vapor one. (c) Bypass of process fluid around the exchanger. The condensing pressure is maintained above atmospheric so that the trap can discharge freely. (d) Cascade control. The steam pressure responds quickly to upsets in steam supply conditions. The more sluggish PF temperature is used to adjust the pressure so as to maintain the proper rate of heat transfer.



Figure 3.10. Condensers. (a) Condenser on temperature control of the PF condensate. Throttling of the flow of the HTM may make it too hot. (b) Condenser on pressure control of the HTM flow. Throttling of the flow of the HTM may make it too hot. (c) Flow rate of condensate controlled by pressure of PF vapor. If the pressure rises, the condensate flow rate increases and the amount of unflooded surface increases, thereby increasing the rate of condensation and lowering the pressure to the correct value. (d) Condenser with vapor bypass to the accumulator drum. The condenser and drum become partially flooded with subcooled condensate. When the pressure falls, the vapor valve opens, and the vapor flows directly to the drum and heats up the liquid there. The resulting increase in vapor pressure forces some of the liquid back into the condenser so that the rate of condensation is decreased and the pressure consequently is restored to the preset value. With sufficient subcooling, a difference of 10-15 ft in levels of drum and condenser is sufficient for good control by this method.



Figure **3.11**. Vaporizers (reboilers). (a) Vaporizer with flow-rate of HTM controlled by temperature of the PF vapor. HTM may be liquid or vapor to start. (b) Thermosiphon reboiler. A constant rate of heat input is assured by flow control of the HTM which may be either liquid or vapor to start. (c) Cascade control of vaporizer. The flow control on the HTM supply responds rapidly to changes in the heat supply system. The more sluggish TC on the PF vapor resets the FC if need be to maintain temperature. (d) Vaporization of refrigerant and cooling of process fluid. Flow rate of the PF is the primary control. The flow rate of refrigerant vapor is controlled by the level in the drum to ensure constant condensation when the incoming PF is in vapor form.

transfer processes the object is to control the final temperature of the process fluid (PF) or the pressure of its source or to ensure a constant rate of heat input. This is accomplished primarily by regulation of the flow of the heat transfer medium (HTM). Regulation of the temperature of the HTM usually is less convenient, although it is done indirectly in steam heaters by throttling of the supply which has the effect of simultaneously changing the condensing pressure and temperature of the steam side.

DISTILLATION EQUIPMENT

As a minimum, a distillation assembly consists of a tower, reboiler, condenser, and overhead accumulator. The bottom of the tower serves as accumulator for the bottoms product. The assembly must be controlled as a whole. Almost invariably, the pressure at either the top or bottom is maintained constant; at the top at such a value that the necessary reflux can be condensed with the available coolant; at the bottom in order to keep the boiling temperature low enough to prevent product degradation or low enough for the available HTM, and definitely well below the critical pressure of the bottom composition. There still remain a relatively large number of variables so that care must be taken to avoid overspecifying the number and kinds of controls. For instance, it is not possible to control the flow rates of the feed and the top and bottom products under perturbed conditions without upsetting holdup in the system.

Two flowsketches are shown on Figures 3.1 and 3.12 of controls on an ethylene fractionator. On Figure 3.1, which is part of the complete process of Figure 3.2, a feedforward control system with a multiplicity of composition analyzers is used to ensure the high degree of purity that is needed for this product. The simpler diagram, Figure 3.12, is more nearly typical of two-product fractionators, the only uncommon variation being the use of a feed-overhead effluent heat exchanger to recover some refrigeration.

Crude oil fractionators are an example of a more elaborate system. They make several products as side streams and usually have some pumparound reflux in addition to top reflux which serve to optimize the diameter of the tower. Figure 3.13 is of such a tower operating under vacuum in order to keep the temperature below cracking conditions. The side streams, particularly those drawn off atmospheric towers, often are steam stripped in external towers hooked up to the main tower in order to remove lighter components. These strippers each have four or five trays, operate



Figure 3.12. Fractionator for separating ethylene and ethane with a refrigerated condenser. FC on feed, reflux, and steam supply. LC on bottom product and refrigerant vapor. Pressure control PC on overhead vapor product.



Figure 3.13. Crude oil vacuum tower. Pumparound reflux is provided at three lower positions as well as at the top, with the object of optimizing the diameter of the tower. Cooling of the side streams is part of the heat recovery system of the entire crude oil distillation plant. The cooling water and the steam for stripping and to the vacuum ejector are on hand control.

off level control on the main tower, and return their vapors to the main tower.

A variety of control schemes are shown separately in Figures 3.14 and 3.15 for the lower and upper sections of fractionators. To some extent, these sections are controllable independently but not entirely so because the flows of mass and heat are interrelated by the conservation laws. In many of the schemes shown, the top reflux rate and the flow of HTM to the reboiler are on flow controls. These quantities are not arbitrary, of course, but are found by calculation from material and energy balances. Moreover, neither the data nor the calculation method are entirely exact, so that some adjustments of these flow rates must be made in the field until the best possible performance is obtained from the equipment. In modern large or especially sensitive operations, the fine tuning is done by computer.

For the lower section of the fractionator, the cases of Figure 3.14

show the heat input to be regulated in these five different ways:

- 1. On flow control of the heat transfer medium (HTM),
- 2. On temperature control of the vapor leaving the reboiler or at some point in the tower,
- 3. On differential pressure between key points in the tower,
- 4. On liquid level in the bottom section,
- 5. On control of composition or some physical property of the bottom product.

Although only one of these methods can be shown clearly on a particular sketch, others often are usable in combination with the other controls that are necessary for completeness. In some cases the HTM shown is condensing vapor and in other cases it is hot oil, but the particular flowsketches are not necessarily restricted to one or the other HTM. The sketches are shown with and without pumps



Figure 3.14. The lower ends of fractionators. (a) Kettle reboiler. The heat source may be on TC of either of the two locations shown or on flow control, or on difference of pressure between key locations in the tower. Because of the built-in weir, no LC is needed. Less head room is needed than with the thermosiphon reboiler. (b) Thermosiphon reboiler. Compared with the kettle, the heat transfer coefficient is greater, the shorter residence time may prevent overheating of thermally sensitive materials, surface fouling will be less, and the smaller holdup of hot liquid is a safety precaution. (c) Forced circulation reboiler. High rate of heat transfer and a short residence time which is desirable with thermally sensitive materials are achieved. (d) Rate of supply of heat transfer medium is controlled by the difference in pressure between two key locations in the tower. (e) With the control valve in the condensate line, the rate of heat transfer is controlled by the amount of **unflooded** heat transfer surface present at any time. (f) Withdrawal on TC ensures that the product has the correct boiling point and presumably the correct composition. The LC on the steam supply ensures that the specified heat input is being maintained. (g) Cascade control: The set point of the FC on the steam supply is adjusted by the TC to ensure constant temperature in the column. (h) Steam flow rate is controlled to ensure specified composition of the PF effluent. The composition may be measured directly **or** indirectly by measurement of some physical property such as vapor pressure. (i) The three-way valve in the hot oil heating supply prevents buildup of excessive pressure in case the flow to the reboiler is throttled substantially. (j) The three-way valve of case (i) is replaced by a two-way valve and a differential pressure controller. This method is more expensive but avoids use of the possibly troublesome three-way valve.



Figure 3.14—(continued)

for withdrawal of bottom product. When the tower pressure is sufficient for transfer of the product to the following equipment, a pump is not needed.

Upper section control methods are shown on Figure 3.15. They all incorporate control of the pressure on the tower, either by throttling some vapor flow rate or by controlling a rate of condensation. In the latter case this can be done by regulating the flow or temperature of the HTM or by regulating the amount of heat transfer surface exposed to contact with condensing vapor.

Flow control of reflux is most common. It is desirable in at least these situations:

- When the temperature on a possible control tray is insensitive to the composition, which is particularly the case when high purity overhead is being made,
- 2. When the expense of composition control is not justifiable,
- 3. When noncondensables are present,
- With tall and wide columns that have large holdup and consequently large lags in interchange of heat and mass between phases,
- 5. When the process coupling of the top and bottom temperature controllers makes their individual adjustments difficult,
- 6. When the critical product is at the bottom.

In all these cases the reflux rate is simply set at a safe value, enough to nullify the effects of any possible perturbations in operation. There rarely is any harm in obtaining greater purity than actually is necessary. The cases that are not on direct control of reflux flow rate are: (g) is on cascade temperature (or composition) and flow control, (h) is on differential temperature control, and (i) is on temperature control of the HTM flow rate.



LIQUID-LIQUID EXTRACTION TOWERS

The internals of extraction towers can be packing, sieve trays, empty with spray feeds or rotating disks. The same kinds of controls are suitable in all cases, and consist basically of level and flow controls. Figure 3.16 shows some variations of such arrangements. If the solvent is lighter than the material being extracted, the two inputs indicated are of course interchanged. Both inputs are on flow control. The light phase is removed from the tower on LC or at the top or on level maintained with an internal weir. The bottom stream is removed on interfacial level control (ILC). A common type of this kind of control employs a hollow float that is weighted to have a density intermediate between those of the two phases. As indicated by Figures 3.16(a) and 3.16(d), the interface can be maintained in either the upper or lower sections of the tower. Some extractions are performed with two solvents that are fed separately to the tower, ordinarily on separate flow controls that may be, however, linked by flow ratio control. The relative elevations of feed and solvents input nozzles depend on the nature of the extraction process.

Controls other than those of flow and level also may be needed in some cases, of which examples are on Figure 3.17. The scheme of part (a) maintains the flow rate of solvent in constant ratio with the main feed stream, whatever the reasons for variation in flow rate of the latter stream. When there are fluctuations in the composition of the feed, it may be essential to adjust the flow rate of the solvent to maintain constancy of some property of one or the other of the effluent streams. Figure 3.17(b) shows reset of the solvent flow rate by the composition of the raffinate. The temperature of an extraction process ordinarily is controlled by regulating the temperatures of the feed streams. Figure 3.17(c) shows the temperature of one of the streams to be controlled by TC-2 acting on the flow rate of the HTM, with reset by the temperature of a control point in the tower acting through TC-1. When the effluents are unusually sensitive to variation of input conditions, it may be inadvisable to wait for feedback from an upset of output performance, but to institute feedforward control instead. In this kind of system, the input conditions are noted, and calculations are made and implemented by on-line computer of other changes that are needed in order to maintain satisfactory operation.

Mixer-settler assemblies for extraction purposes often are preferable to differential contact towers in order to obtain very high extraction yields or to handle large flow rates or when phase



(a)



FC # LC PF

(c)



Figure 3.15. Control modes for the upper sections of fractionators. (a) Pressure control by throttling of the overhead vapor flow. The drawbacks of this method are the cost of the large control valve and the fact that the reflux pump operates with a variable suction head. The flow of HTM is hand set. (b) Applicable when the overhead product is taken off as vapor and only the reflux portion need be condensed. Two two-way valves can replace the single three-way valve. The flow of HTM is hand set. (c) Flow rate of the HTM is regulated to keep the pressure constant. One precaution is to make sure that the HTM, for example water, does not overheat and cause scaling. The HTM flow control valve is small compared with the vapor valve of case (a). (d) Pressure control is maintained by throttling uncondensables can be handled this way. The flow of the HTM is manually set. (e) Bypass of vapor to the drum on PC: The bypassed vapor heats up the liquid there, thereby causing the pressure to rise. When the bypass is closed, the pressure falls. Sufficient heat transfer surface is provided to **subcool** the condensate. (f) Vapor bypass between the condenser and the accumulator, with the condenser near ground level for the ease of maintenance: When the pressure in the tower falls, the bypass valve opens, and the subcool liquid in the drum heats up and is forced by its vapor pressure back into the condenser. Because of the smaller surface now exposed to the vapor, the rate of condensation is decreased and consequently the tower pressure increases to the preset value. With normal subcooling, obtained with some excess surface, a difference of 10–15 ft in levels of drum and condenser is sufficient for good control. (g) Cascade control: The same system as case (a), but with addition of a TC (or composition controller) that resets the reflux flow rate. (h) Reflux rate on a differential temperature controller. Ensures constant internal reflux rate even when the performance of the condenser fluctuates. (i) Reflux is provided by a separ









(h)

(g)



Figure 3.15—(continued)



Figure 3.16. Extraction tower control. (a) Operation with heavy solvent, interface in the upper section, top liquid level on LC. (b) Same as part (a) but with overflow weir for the light phase. (c) Same as part (a) but with completely full tower and light phase out at the top. (d) Operation with interface on ILC in the lower section, removal of the light phase from the upper section by any of the methods of (a), (b), or (c).

separation is slow and much time is needed. Often, also, relatively simple equipment is adequate for small capacities and easy separations. Several designs of varying degrees of sophistication are available commercially, some of which are described by Lo, Baird, Hanson (Handbook of Solvent Extraction, Wiley, New York, 1983). The basic concept, however, is illustrated on Figure 3.18. The solvent and feed are thoroughly mixed in one chamber and overflow into another, partitioned chamber where separation into light and heavy phases occurs by gravity. Ordinarily the settling chamber is much the larger. The heavy phase is removed on interfacial level control and the light one on level control. The takeoffs also can be controlled with internal weirs or manually.

Several centrifugal contactors of proprietary nature are on the market. Their controls are invariably built in.

CHEMICAL REACTORS

The progress of a given reaction depends on the temperature, pressure, flow rates, and residence times. Usually these variables are controlled directly, but since the major feature of a chemical reaction is composition change, the analysis of composition and the resetting of the other variables by its means is an often used means of control. The possible occurrence of multiple steady states and the onset of instabilities also are factors in deciding on the nature and precision of a control system.

Because of the sensitivity of reaction rates to temperature,

control of that variable often dominates the design of a reactor so that it becomes rather a heat exchanger in which a reaction occurs almost incidentally. Accordingly, besides the examples of reactor controls of this section, those of heat exchangers in that section may be consulted profitably. Heat transfer and holding time may be provided in separate equipment, but the complete assembly is properly regarded as a reactor. An extreme example, perhaps, is the two-stage heater-reactor system of Figure 3.19(f); three or more such stages are used for endothermic catalytic reforming of naphthas, and similar arrangements exist with intercoolers for exothermic processes.

Although the bulk of chemical manufacture is done on a continuous basis, there are sectors of the industry in which batch reactors are essential, notably for fermentations and polymerizations. Such plants may employ as many as 100 batch reactors. The basic processing steps include the charging of several streams, bringing up to reaction temperature, the reaction proper, maintenance of reaction temperature, discharge of the product, and preparation for the next batch. Moreover, the quality of the product depends on the accuracy of the timing and the closeness of the control.

Small installations are operated adequately and economically by human control, but the opening and closing of many valves and the setting of conditions at precise times clearly call for computer control of multiple batch installations. Computers actually have taken over in modern synthetic rubber and other polymerization industries. Interested readers will find a description, complete with


Figure 3.17. Some other controls on extraction towers. (a) Solvent flow rate maintained in constant ratio with the feed rate. (b) Solvent flow rate reset by controlled composition of raffinate. (c) Temperature of solvent or feed reset by the temperature at a control point in the tower.

logic diagrams for normal and emergency operations, of the tasks involved in generating a computer system for a group of batch reactors in the book of Liptak (1973, pp. 536-565). Control of discontinuous processes in general is treated in the book of Skrokov (1980, pp. 128-163).

In the present discussion, emphasis will be placed on the control of continuous reactors, concentrating on the several examples of Figure 3.19 in the order of the letter designations of individual figures used there.

- (a) Stirred tanks are used either as batch or continuous flow reactors. Heat transfer may be provided with an external heat exchanger, as shown on this figure, or through internal surface or a jacket. Alternate modes of control may be used with the controls shown: (i) When the HTM is on temperature control, the pumparound will be on flow control; (ii) when the pumparound is on temperature control, the HTM will be on flow control; (iii) for continuous overflow of product, the control point for temperature may be on that line or in the vessel; (iv) for batch operation, the control point for temperature clearly must be in the vessel. Although level control is shown to be maintained with an internal weir, the product can be taken off with the pump on level control.
- (b) This shows either direct or cascade control of the temperature of a reactor with internal heat transfer surface and an internal weir. The sluggishly responding temperature of the vessel is used to reset the temperature controller of the HTM. For direct control, the TC-2 is omitted and the control point can be on the HTM outlet or the product line or in the vessel.
- (c) Quite a uniform temperature can be maintained in a reactor if the contents are boiling. The sketch shows temperature maintenance by refluxing evolved vapors. A drum is shown from which uncondensed gases are drawn off on pressure control, but the construction of the condenser may permit these gases to be drawn off directly, thus eliminating need for the drum. The HTM of the condenser is on TC which resets the PC if necessary in order to maintain the correct boiling temperature in the reactor. Other modes of pressure control are shown with the fractionator sketches of Figure 3.15 and on Figure 3.5 dealing with vacuum control.
- (d) Flow reactors without mechanical agitation are of many configurations, tanks or tubes, empty or containing fixed beds of particles or moving particles. When the thermal effects of reaction are substantial, multiple small tubes in parallel are used to provide adequate heat transfer surface. The sketch shows a single tube provided with a jacket for heat transfer. Feed to the reactor is on flow control, the effluent on pressure control, and the flow of the HTM on temperature control of the effluent with the possibility of reset by the composition of the effluent.
- (e) Heat transfer to high temperature reactions, above 300°C or so, may be accomplished by direct contact with combustion gases. The reaction tubes are in the combustion zone but safely away from contact with the flame. The control mode is essentially similar to that for case (d), except that fuel-air mixture takes the place of the HTM. The supply of fuel is on either temperature or composition control off the effluent stream, and the air is maintained in constant ratio with the fuel with the flow ratio controller FRC.
- (f) High temperature endothermic processes may need several reaction vessels with intermediate heat input. For example, the inlet temperature to each stage of a catalytic reformer is about 975°F and the temperature drop ranges from about 100°F in the first stage to about 15°F in the last one. In the two-stage assembly of this figure, the input is on FC, the outlet of the last reactor on PC, and the fuel supply to each furnace is on TC of its effluent, with the air supply on flow ratio control, as shown for example (e).
- (g) Very effective heat transfer is accomplished by mixing of streams at different temperatures. The cumene process shown here employs injection of cold reacting mixture and cold inert propane and water to prevent temperature escalation; by this scheme, the inlet and outlet temperature are made essentially the same, about 500°F. Although not shown here, the main feed is, as usual for reactors, on FC and the outlet on PC. The



Figure 3.18. Functioning and controls of a mixer-settler assembly for liquid-liquid extraction.

sidestreams are regulated with hand-set valves by experienced operators in this particular plant, but they could be put on automatic control if necessary. Other processes that employ injection of cold process gas at intermediate points are some cases of ammonia synthesis and sulfur dioxide oxidation.

(h) In catalytic cracking of petroleum fractions, an influential side reaction is the formation of carbon which deposits on the catalyst and deactivates it. Unacceptable deactivation occurs in about 10 min, so that in practice continuous reactivation of a portion of the catalyst in process must be performed. As shown on this sketch, spent catalyst is transferred from the reactor to the regenerator on level control, and returns after regeneration under TC off the reactor temperature. Level in the regenerator is maintained with an overflow standpipe. Smooth transfer of catalyst between vessels is assisted by the differential pressure control DPC, but in some plants transfer is improved by injection of steam at high velocity into the lines as shown on this sketch for the input of charge to the reactor. Feed to the system as a whole is on flow control. Process effluent from the reactor is on pressure control, and of the regenerator gases on the DPC. Fuel to regeneration air preheater is on TC off the preheat air and the combustion air is on flow ratio control as in part (e).

LIQUID PUMPS

Process pumps are three types: centrifugal, rotary positive displacement, and reciprocating. The outputs of all of them are controllable by regulation of the speed of the driver.

Controllability of centrifugal pumps depends on their **pressure**flow characteristics, of which Figure 3.20 has two examples. With the upper curve, two flow rates are possible above a head of about 65 ft so that the flow is not reliably controllable above this pressure. The pump with the lower curve is stable at all pressures within its range. Throttling of the discharge is the usual control method for smaller **centrifugals**, variable speed drives for larger ones. Suction throttling may induce flashing and vapor binding of the pump. Figures 3.21(a) and (b) are examples.

Rotary pumps deliver a nearly constant flow at a given speed, regardless of the pressure. Bypass control is the usual method, with speed control in larger sizes. Reciprocating pumps also may be controlled on bypass if a pulsation damper is provided in the circuit to smooth out pressure fluctuations; Figure 3.21(c) shows this mode.

Reciprocating positive displacement pumps may have adjustment of the length or frequency of the stroke as another control feature. These may be solenoid or pneumatic devices that can be operated off a flow controller, as shown on Figure 3.21(d).

SOLIDS FEEDERS

Several of the more common methods of controlling the rate of supply of granular, free-flowing solids are represented in Figure 3.7.

COMPRESSORS

Three main classes of gas compressors are centrifugal and axial, rotary continuous positive displacement, and reciprocating positive **56** PROCESS CONTROL



Figure 3.19. Chemical reactor control examples. (a) Temperature control of a stirred tank reactor with pumparound through an external heat exchanger, operable either in batch or continuously: Some alternate control modes are discussed in the text. Cascade control as in (b) can be implemented with external heat transfer surface. (b) Either cascade or direct control of temperature: For direct control, controller TC-2 is omitted, and the control point can be taken on the effluent line or in the vessel or on the HTM effluent line. A similar scheme is feasible with an external heat exchanger. (c) Reactor temperature control by regulation of the boiling pressure: The HTM is on TC off the reactor and resets the PC on the vent gases when necessary to maintain the correct boiling temperature. Although shown for batch operation, the method is entirely feasible for continuous flow. (d) Basic controls on a flow reactor: Feed on flow control, effluent on pressure control, and heat transfer medium flow rate on process effluent temperature or reset by its composition. (e) A fired heater as a tubular flow reactor: Feed is on FC, the product is on PC, the fuel is on TC or AC off the product, and the air is on flow ratio control. (f) A two-stage fired heater-reactor assembly: Details of the fuel-air supply control are in (e). (g) Control of the temperature of the exothermic synthesis of cumene by splitting the feed and by injection of cold propane and water into several zones. The water also serves to maintain activity of the phosphoric acid catalyst. (h) The main controls of a fluidized bed reactor-regenerator: Flow of spent catalyst is on level control, and that of regenerated catalyst is on TC off the reactor; these flows are assisted by maintenance of a differential pressure between the vessels. Details of the fuel-air control for the preheater are in (e).











(h)

Figure 3.19—(continued)



Figure 3.20. Characteristics curves of two centrifugal pumps.

displacement. The usual or feasible modes of control of pressure and flow may be tabulated:

Control Mode	Centrifugal I and Axial	Rotary PD	Reciprocating P D
Suction throttling	х		
Discharge throttling	х		
Bypass	х	х	х
Speed	х	х	х
Guide vanes	х		
Suction valves			x
Cylinder clearance			x



Figure 3.21. Control of centrifugal, rotary, and reciprocating pumps. (a) Throttling of the discharge of a centrifugal pump. (b) Control of the flow rate of any kind of pump by regulation of the speed of the driver. Although a turbine is shown, engine drive or speed control with gears, magnetic clutch, or hydraulic coupling may be feasible. (c) On the left, bypass control of rotary positive displacement pump; on the right, the reciprocating pump circuit has a pulsation dampener to smooth out pressure fluctuations. (d) Adjustment of the length or frequency of the stroke of a constant speed reciprocating pump with a servomechanism which is a feedback method whose action is control of mechanical position.



Figure 3.21—(continued)

Throttling of the suction of centrifugal and axial compressors wastes less power than throttling the discharge. Even less power is wasted by adjustment of built-in inlet guide vanes with a servomechanism which is a feedback control system in which the controlled variable is mechanical position. Speed control is a particularly effective control mode, applicable to large units that can utilize turbine or internal combustion drives; control is by throttling of the supply of motive fluids, steam or fuel.

Characteristic curves-pressure against flowrate-of centrifugal and axial compressors usually have a peak. Figure 3.22 is an example. In order to avoid surging, the flow through the com-



Figure 3.22. Characteristic curves of a centrifugal compressor at different speeds, showing surge limits.

3.3. EQUIPMENT CONTROL 59

pressor must be maintained above the magnitude at the peak in pressure. Figure 3.23(c) shows an automatic bypass for surge protection which opens when the principal flow falls to the critical minimum; recycle brings the total flow above the critical.

Smaller rotary positive displacement compressors are controlled with external bypass. Such equipment usually has a built-in relief valve that opens at a pressure short of damaging the equipment, but the external bypass still is necessary for smooth control. Large units may be equipped with turbine or gas engine drives which are speed adjustable. Variable speed gear boxes or belt drives are not satisfactory. Variable speed dc motors also are not useful as compressor drives. Magnetic clutches and hydraulic e u s e d .

Reciprocating compressors may be controlled in the same way as rotary units. The normal turndown with gasoline or diesel engines is 50% of maximum in order that torque remains within



Figure 3.23. Control of centrifugal compressors with turbine or motor drives. (a) Pressure control with turbine or motor drives. (b) Flow control with turbine or motor drives. SC is a servomechanism that adjusts the guide vanes in the suction of the compressor. (c) Surge and pressure control with either turbine or motor drive. The bypass valve opens only when the flow reaches the minimum calculated for surge protection.



Figure 3.24. Control of positive displacement compressors, rotary and reciprocating. (a) Flow control with variable speed drives. (b) Pressure control with bypass to the suction of the compressor. (c) Reciprocating compressor. SC is a servomechanism that opens some of suction valves during discharge, thus permitting **stepwise** internal bypass. The clearance unloader is controllable similarly. These built-in devices may be supplemented with external bypass to smooth out pressure fluctuations.

acceptable limits. Two other aids are available to control of reciprocating units.

- Valve unloading, a process whereby some of the suction valves remain open during discharge. Solenoid or pneumatic unloaders can be operated from the output of a control instrument. The stepwise controlled flow rate may need to be supplemented with controlled external bypass to smooth out pressure fluctuations.
- Clearance unloaders are small pockets into which the gas is forced on the compression stroke and expands into the cylinder on the return stroke, thus preventing compression of additional gas.

Figure 3.24 shows control schemes for rotary and reciprocating compressors. Vacuum pumps are compressors operating between a low suction pressure and a fixed discharge pressure, usually

atmospheric. Mechanical pumps are used for small capacities, steam jet ejectors for larger ones. Ejectors also are used as thermocompressors to boost the pressure of low pressure steam to an intermediate value. Control of suction pressure with either mechanical or jet pumps is by either air bleed [Fig. 3.5(a)] or suction line throttling [Fig. 3.5(c)]; air bleed is the more economical process. Up to five jets in series are used to produce high vacua. The steam from each stage is condensed by direct contact with water in barometric condensers or in surface condensers; condensation of steam from the final stage is not essential to performance but only to avoid atmospheric pollution. In a single stage ejector, motive steam flow cannot be reduced below critical flow in the diffuser, and water to the barometric condenser must not be throttled below 30-50% of the maximum if proper contacting is to be maintained. Control by throttling of steam and water supply, as on Figure 3.5(b), is subject to these limitations.

REFERENCES

- Chemical Engineering Magazine, Practical Process Instrumentation and Control, McGraw-Hill, New York, 1980.
- D.M. Considine, Process Instruments and Controls Handbook, McGraw-Hill, New York, 1985.
- B. Liptak, Instrumentation in the Process Industries, Chilton, New York, 1973.
- 4. F.G. Shinskey, Process Control Systems, McGraw-Hill, New York, 1979.
- 5. F.G. Shinskey, Distillation Control, McGraw-Hill, New York, 1984.
- M.R. Skrokov (Ed.), Mini- and Microcomputer Control in Industrial Processes, Van Nostrand Reinhold, New York, 1980.

DRIVERS FOR MOVING EQUIPMENT

Provided chemical processing equipment includes pumps, compressors, agitators and mixers, crushers and grinders, and conveyors. Drivers are electric motors, steam or gas turbines, and internal combustion engines. For loads under 150 HP or so electric motors are almost invariably the choice. Several criteria are applicable. For example, when a pump and a spare are provided, for flexibility one of them may be driven by motor and the other by turbine. Centrifugal and axial blowers and compressors are advantageously driven by turbines because the high operating speeds of 4000– 10,000 rpm are readily attainable whereas electric motors must operate through a speed increasing gear at extra expense. When fuel is relatively cheap or accessible, as in the field, gas turbines and internal combustion engines are preferred drivers. Turbines, internal combustion engines, and direct current motors are capable of continuous speed adjustment over a wide range. Energy efficiencies **vary** widely with the size and type of driver as shown in this table.

	Efficiency (%)						
Driver	10 kW	100 kW	1000 kW	10,000 kW			
Gas turbine andinternal conbustion engine		28	34	38			
Steam turbine		42	63	76			
Motor	8.5	92	96	97			

Since the unit energy costs are correspondingly different, the economics of the several drive modes often are more nearly comparable.

4.1. MOTORS

Although each has several subclasses, the three main classes of motors are induction, synchronous, and direct current. Higher voltages are more efficient, but only in the larger sizes is the housing ample enough to **accomodate** the extra insulation that is necessary. The voltages commonly used are

Horsepower	Voltage				
I-100	220. 440. 550				
75-250	440				
200-2500	2300, 4000				
Above 2500	4000, 13, 200				

Direct current voltages are 11.5, 230, and 600.

The torque-speed characteristic of the motor must be matched against that of the equipment, for instance, a pump. As the pump comes up to speed, the torque exerted by the driver always should remain 5% or so above that demanded by the pump.

The main characteristics of the three types of motors that bear on their process applicability are summarized following.

INDUCTION

Induction motors are the most frequent in use because of their simple and rugged construction, and simple installation and control. They are constant speed devices available as **3600** (two-pole), 1800, 1200, and 900rpm (eight-pole). Two speed models with special windings with 2: 1 speed ratios are sometimes used with agitators, centrifugal pumps and compressors and fans for air coolers and cooling towers. Capacities up to **20,000 HP** are made. With speed

increasing gears, the basic 1800 rpm model is the economical choice as drive for centrifugal compressors at high speeds.

SYNCHRONOUS

Synchronous motors are made in speeds from 1800 (two-pole) to 150 rpm (48-pole). They operate at constant speed without slip, an important characteristic in some applications. Their efficiencies are 1-2.5% higher than that of induction motors, the higher value at the lower speeds. They are the obvious choice to drive large low speed reciprocating compressors requiring speeds below 600 rpm. They are not suitable when severe fluctuations in torque are encountered. Direct current excitation must be provided, and the costs of control equipment are higher than for the induction types. Consequently, synchronous motors are not used under 50 HP or so.

DIRECT CURRENT

Direct current motors are used for continuous operation at constant load when fine speed adjustment and high starting torque are needed. A wide range of speed control is possible. They have some process applications with centrifugal and plunger pumps, conveyors, hoists, etc.

Enclosures. In chemical plants and refineries, motors may need to be resistant to the weather or to corrosive and hazardous locations. The kind of housing that must be provided in particular situations is laid out in detail in the National Electrical Code, Article 500. Some of the classes of protection recognized there are in this table of differential costs.

	% Cost above				
Туре	Drip Proof	Protection Against			
Drip proof		Dripping liquids and falling particles			
Weather protected, I and II	10-50	Rain, dirt, snow			
Totally enclosed fan cooled, TEFC, below 250 HP	25-100	Explosive and nonexplosive atmospheres			
Totally enclosed, water cooled, above 500 HP	25-100	Same as TEFC			
Explosion proof, below 3000 HP	110- 140	Flammable and volatile liquids			

TABLE 4.1. Selection of Motors for Process Equipment

	Motor Type'					
Application	A.C.	D.C.				
Agitator	la, lb, 2b	5a				
Ball mill	lc, 2b, 3a	5 b				
Blower	1a, 1b, 2b, 3a, 4	5 a				
Compressor	1a, 1b, 1c, 3a, 4	5b, 7				
Conveyor	1a, 1c, 2b, 3a	5b. 7				
Crusher	la, lc, 1d	5a, 5b				
Dough mixer	la, lb, 1c, 2b	5a, 5b				
Fan, centrifugal and propeller	la, 1b, 2c, 3a, 4	5a, 7				
Hammer mill	lc	5 a				
Hoist	1d, 2a, 3b	6				
Pulverizer	lc	5 b				
Pump, centrifugal	1a, 1b, 2b, 3a, 4	5 b				
Pump, positive displacement	Ic, 2b, 3a	5 b				
Rock crusher	3 a	5b, 6				

⁸ Code:

1. Squirrel-cage, constant speed

a. normal torque, normal starting current

- b. normal torque, low starting current
- c. high torque, low starting current
- d. high torque, high slip
- 2. Squirrel-cage, multispeed a. constant horsepower
 - b. constant torque
 - c. variable torque
- 3. Wound rotor
- a. general purpose b. crane and hoist
- Synchronous
- 5. Direct current, constant speed
- a. shunt wound
- b. compound wound
- 6. Direct current, variable speed series wound
- 7. Direct current, adjustable speed

(After Allis-Chalmers Mfg. Co., Motor and Generator Reference Book, Colorado Springs, CO).

Standard NEMA ratings for induction motors:

General purpose: $\frac{1}{2}$, $\frac{3}{4}$, 1, $1\frac{1}{2}$, 2, 3, 5, $7\frac{1}{2}$, 10. 15, 20, 25, 30, 40, 50, 60, 75, 100, 125, 150, 200, 250, 300, 350, 400, 450, 500.

Large motors: 250, 300, 350, 400, 450, 500, 600, 700,800, 900, 1000, 1250, 1500, 1750, 2000, 2250, 2500, 3000, 3500, 4000, 4500, 5000 and up to 30.000.

Clearly the cost increments beyond the basic drip-proof motor enclosures are severe, and may need to be balanced in large sizes against the cost of isolating the equipment in pressurized buildings away from the hazardous locations.

Applications. The kinds of motors that are being used successfully with particular kinds of chemical process equipment are identified in Table 4.1. As many as five kinds of AC motors are shown in some instances. The choice may be influenced by economic considerations or local experience or personal preference. In this area, the process engineer is well advised to enlist help from electrical experts. A checklist of basic data that a supplier of a motor must know is in Table 4.2. The kind of enclosure may be specified on the last line, operating conditions.

4.2. STEAM TURBINES AND GAS EXPANDERS

Turbines utilize the expansion of steam or a gas to deliver power to a rotating shaft. Salient features of such equipment are

1. high speed rotation,

- 2. adjustable speed operation,
- 3. nonsparking and consequently nonhazardous operation,

TABLE 4.2. Checklist for Selection of Motors

_

Motor Data
General
Type of motor (cage, wound-rotor, synchronous, or dc) Quantity Hp Rpm Phase Cycles. Voltage. Time rating (continuous, short-time, intermittent). Overload (if any) Overload (if any)
Induction Motors
Locked-rotor torque % Breakdown torque
Synchronous Motors
Power factor Torques: Locked-rotor. % Pull-in. % Pull-out % Excitation .volts dc Type of exciter. % If m-g exciter set, what are motor characteristics? Motor field rheostat. Motor field discharge resistor
Direct-current Motors
Shunt, stabilized shunt, compound, or series wound Speed range Non-reversing or reversing Continuous or tapered-rated
Mechanical Featured
Protection or enclosure
Mechanical Features (cont.)
Number of bearings Type of bearings Shaft extension: Flanged Standard or special length Press on half-coupling Terminal box NEMA C or D flange Round-frame or with feet Vertical: External thrust load Ibs. Type of thrust bearing. Base ring type Sole plates Accessories. Sole plates

Load Data

Typeofload
If compressor drive, give NEMA application number
Direct-connected, geared, chain, V-belt, or flat-belt drive
Wk' (inertia) for high inertia drives
Starting with full load, or unloaded If unloaded, by what means?
For variable-speed or multi-speed drives, is load variable torque, constant torque, or constant horsepower? Operating conditions

(By permission, Allis Chalmers Motor and Generator Reference Book, Bul. 51R7933, and ES. Lincoln (Ed.), Electrical Reference Book, Electrical Modernization Bureau, Colorado Springs, CO.

4. simple controls,

5. low first cost and maintenance, and

6. flexibility with regard to inlet and outlet pressures.

Single stage units are most commonly used as drivers, but above 500 HP or so multistage units become preferable. Inlet steam pressures may be any value up to the critical and with several









(b)

Figure 4.1. Efficiencies of (a) single-stage and (b) multistage turbines (Gartmann, De Laval Engineering Handbook, McGraw-Hill, New York, 1970, pp. 5.8-S. 9, Figs. 5.2 and 5.3).

hundred degrees of superheat. In larger sixes turbines may be convenient sources of low pressure exhaust steam in the plant. From multistage units, steam may be bled at several reduced pressures. When the expansion is to subatmospheric conditions, the operation is called condensing because the exhaust steam must be condensed before removal from the equipment. Although the efficiency of condensing turbines is less, there is an overall reduction of energy consumption because of the wider expansion range.

Several parameters **affect** the efficiency of steam turbines, as shown partially on Figure 4.1. Closer examination will need to take into account specific mechanical details which usually are left to the manufacturer. Geared turbines [the dashed line of Fig. 4.1(b)] have higher efficiencies, even with reduction gear losses, because they operate with especially high bucket speeds. For example, for a service of 500 HP with 300 psig steam, a geared turbine has an efficiency of 49.5% and one with a direct drive at 1800 rpm has an efficiency of 24%.

The flow rate of steam per unit of power produced is represented by

<i>m</i> = -	$-\frac{2545}{\eta(H_2-H_1)}$ lb/HP hr	
_	$\frac{3412}{\eta(H_2 - H_1)} \text{lb/kWh}$	

with the enthalpies in **Btu/lb**. The efficiency is η , off Figure 4.1, for example. The enthalpy change is that of an isentropic process. It may be calculated with the aid of the steam tables or a Mollier diagram for steam. For convenience, however, special tables have been derived which give the theoretical steam rates for typical combinations of inlet and outlet conditions. Table 4.3 is an abbreviated version.

Example 4.1 illustrates this kind of calculation and compares the result with that obtained by taking the steam to behave as an ideal gas. For **nonideal** gases with known **PVT** equations of state and low pressure heat capacities, the method of calculation is the same as for compressors which is described in that section of the book.

On a Mollier diagram like that with Example 4.1, it is clear that expansion to a low pressure may lead to partial condensation if insufficient preheat is supplied to the inlet steam. The final condition after application of the efficiency correction is the pertinent one, even though the isentropic point may be in the two-phase region. Condensation on the blades is harmful to them and must be avoided. Similarly, when carbon dioxide is expanded, possible formation of solid must be guarded against.

When gases other than steam are employed as motive fluids, the equipment is called a gas expander. The name gas turbine usually is restricted to equipment that recovers power from hot

TABLE 4.3. Theoretical Steam Rates for Typical Steam Conditions (Ib/kWh)*

							Initial p	messure,	lb∕in² gau	<i>z</i> c						
	150	250	400	600	600	850	850	900	900	1,200	1,250	1,250	1,450	+,450	1,800	2,400
	_							Initial	temp, °F							
	365.9	500	650	750	825	x 2 5	900	825	900	x 2 5	900	950	X25	950	1000	1000
_							I	nitial sup	perheat, °	1					_	
	0	94.0	201.9	261.2	336.2	297.8	372.8	291.1	366.1	256.3	326.1	376.1	232.0	357.0	377.9	337.0
Exhaust							In	itial ent	nalpy <u>, Bri</u>	u/lb						
pressure	1,195.5	1,2	2 6 1	. Х	1,334	.91,379.	61,421.4	1,410.61	,453.51,4	108.41,45	1.61,394	.71,438	+1,468.1	1,382.71,	461.21,4	80.11,460.
inf lg abs																
2.0	10.52	9.070	7,831	7.083	6.761	6,580	6.282	6555	6.256	6.451	6.133	5.944	6.408	5.900	5.668	5.633
2.5	10.88	Y. 343	x.03;	7.251	6.916	6.7.23	6.415	6.696	6.388	6.584	6.256	6.061	6.536	6.014	5.773	5.733
3.0	t1.20	9.582	8.217	7.396	7.052	6.847	6.330	6.819	6.502	6.699	6.362	6.162	6.648	6.112	5.862	5.819
+.0 11./in2 man	, II./6	9.996	8.524	1.044	1.282	7.058	6.726	7.026	6.6 ሃዛ	6.894	6.2 <u>41</u>	6.332	0.835	6.277	6.015	3.903
in/in_gage		16 57	13.01	1.05	+0.42	0 8 3 8	0 788	0.755	¥ 200	0.207	8 820	8 101	9.718	¥ 351	7 874	7 713
10	73.97	17.90	13.83	1.61	10.95	10 10	9.200	10 202	9.617	9.397 V 707	9.180	8 830	0 503	X.673	8 1 58	7 975
20	2X.61	20.44	15.33	2.68	11.90	11 10	10.43	to 987	10.327	10.490	9.801	9.415	10.240	9.227	8 642	8.421
30	33.69	22.95	16.73	3.63	12.75	11.80	11.08	11.67	10.952	11.095	10.341	9.922	10.801	9.704	9.057	x.799
40	39.39	25.52	18.08	4.51	13.54	12.46	11.66	12.304	11.52	11.646	10.831	10.380	11.309	10,134	9.427	9.136
50	46.00	2X.21	19.42	5.36	14.30	13.0;	1?.?!	12.00	12.06	12.16	11.284	10.804	11.779	10.531	9.767	9.442
60	53.90	31.07	20.76	16.IX	15.05	13.66	12.74	13.47	12.57	12.64	11.71	11.20	12.22	10.90	10.08	9.727
75	69. 1	35.77	22.81	17.40	16.16	14.50	13.51	14.28	13.30	13.34	12.32	11.77	12.85	11,43	10.53	to.12
80	75.9	37.47	23.51	17.80	16.54	14.78	13.77	14.55	13.55	13.56	12.52	11.95	t 3.05	11.60	10.67	10.25
100		45.21	26.46	19.43	IX.05	15.86	I-t.77	15.59	I-t.50	I-t.42	13.27	12.65	13.83	12.2-t	11.21	10.73
125		57.88	30.39	21.36	20.03	17.22	16.04	16.87	15.70	15.40	14.17	15.51	14.76	13.01	11.84	11.28
150		/0.3	33.40	23.83	22.14	10.17	11.33	18.18	10.91	10.4/	12.00	14.00	12.02	13.75	12.44	12.00
100		X0.X	37.37	24.19	23.03	19.17	17.65	10.57	17.41 IV 16	10.00	15.41	1-1.09	16.00	14.05	12.00	12.00
200			T1.10	20.27	24.45	20.04	20.05	20.01	10 / 5	17.70	16.84	16.05	17 30	15.23	13.62	12.27
250			6Y.I	35.40	32.x9	24.7X	23.08	23.90	22.2-t	20.57	18.68	17.81	19.11	16.73	14.78	13.69
300				42.72	40.62	2X. 5()	26.53	27.27	25.37	22.79	20.62	19.66	20.89	18.28	t5.95	14.59
400				72.2	67.0	3X.05	35.43	35.71	32.22	27.82	24.99	23.X2	24.74	21.64	IX.39	16.41
425				x-t.2	78.3	41.08	3 X . 2 6	3X.33	35.65	29.2-t	26.21	24.98	25.7X	22.55	19.03	16.87
600						ix.5	73.1	6X.11	63t	42.10	37.03	35.30	34.50	30. 16	24.06	20.29

'From Theoretical Steam Rate Table-Compatible with the 1967 ASME Steam Tables, ASME, 1969.

EXAMPLE 4.1

Steam Requirement of a Turbine Operation

Steam is fed to a turbine at 614.7 psia and 825°F and is discharged at 64.7 psia. (a) Find the theoretical steam rate, **lb/kWh**, by using the steam tables. (b) If the isentropic efficiency is 70%, find the outlet temperature. (c) Find the theoretical steam rate if the behavior is ideal, with $C_p/C_v = 1.33$.

(a) The expansion is isentropic. The initial and terminal conditions are identified in the following table and on the graph. The data are read off a large Mollier diagram (Keenan et al., Steam *Tables*, Wiley, New York, 1969).

Point	P	<i>T</i> ⁰F	Н	S
1	614. 7	825	1421.4	1.642
2	64.7	315	1183.0	1.642
3	64.7	445	1254.5	1.730

AH, $= H_2 - H_1 = -238.4$ Btu/lb

Theoretical steam rate = 3412/238.4 = 14.31 lb/kWh. This value is checked exactly with the data of Table 4.3.

(b) $H_3 - H_1 = 0.7(H_2 - H_1) = -166.9$ Btu/lb $H_3 = 1421.4 - 166.9 = 1254.5$ Btu/lb

The corresponding values of T_3 and S_3 are read off the Mollier diagram, as tabulated.

(c) The isentropic relation for ideal gases is

AH=
$$\frac{k}{k-1} RT_1[(P_2/P_1)^{(k-1)/k} - 1]$$

= $\frac{1.987(1285)}{0.25}[(64.7/614.7)^{0.25} - 1]$
= -4396 Btu/lbmol, -244 Btu/lb.



combustion gases. The name turboexpander is applied to machines whose objective is to reduce the energy content (and temperature) of the stream, as for cryogenic purposes.

Gas expanders are used to recover energy from high pressure process gas streams in a plant when the lower pressure is adequate for further processing. Power calculations are made in the same way as those for compressors. Usually several hundred horsepower must be involved for economic justification of an expander. In smaller plants, pressures are simply let down with throttling valves (Joule-Thomson) without attempt at recovery of energy.

The specification sheet of Table 4.4 has room for the process conditions and some of the many mechanical details of steam turbines.

4.3. COMBUSTION GAS TURBINES AND ENGINES

When a low cost fuel is available, internal combustion drivers surpass all others in compactness and low cost of installation and operation. For example, gas compression on a large scale has long been done with integral engine compressors. Reciprocating engines also are widely used with centrifugal compressors in low pressure applications, but speed increasing gears are needed to up the 300–600 rpm of the engines to the 3000–10,000 rpm or so of the compressor.

Process applications of combustion gas turbines are chiefly to driving pumps and compressors, particularly on gas and oil transmission lines where the low thermal efficiency is counterbalanced by the convenience and economy of having the fuel on hand. Offshore drilling rigs also employ gas turbines. Any hot process gas at elevated pressure is a candidate for work recovery in a turbine. **Offgases** of catalytic cracker regenerators, commonly at 45 psig and as high as 1250°F, are often charged to turbines for partial recovery of their energy contents. Plants for the manufacture of nitric acid by oxidation of ammonia at pressures of 100 psig or so utilize expanders on the **offgases** from the absorption towers, and the recovered energy is used to compress the process air to the reactors.

Combustion gas turbine processes are diagrammed on Figure 4.2 and in Example 4.2. In the basic process, a mixture of air and fuel (or air alone) is compressed to 5-10 atm, and then ignited and burned and finally expanded through a turbine from which power is recovered. The process follows essentially a **Brayton** cycle which is shown in Figure 4.2 in idealized forms on TS and PV diagrams. The ideal process consists of an isentropic compression, then heating at constant pressure followed by an isentropic expansion and finally cooling at the starting pressure. In practice, efficiencies of the individual steps are high:

Compressor isentropic efficiency, 85% Expander isentropic efficiency, 85-90% Combustion efficiency, 98%

GENERAL-PURPOSE STEAM TURBINE DATA SHEET CUSTOMARY UNITS

CONTRACT NO	
REV. NO.	DATE
BY	REVIEWED
SHEET	OF 2
PO NO	

Applicable to:		O Purchese	С	As - Built					
For	·			Unit					
Site No. Required									
Service Driven Equipment									
Manufacturer				Madel	Serial No				
	terester Completed	□ <u> </u>		By Marrufa	cturer				
C) OPERATING CONDIT	IONS			PERFORM	IANCE			
Operating Point	Power,	Sp	eed,	Operating Po	int/ NO.	Hand Veives	Steem Rate,		
Operating I onic	BHP	R	ΡY	Steam Condi	ition Op	en (3.4.1.4)	LDWHP H.r.		
Normal				Normal/Nor	mai				
Rated				Rated/Norm	al				
Other				Rated/Min. Inlet, Max	k. Exhaust				
				Indicate Guarantee Po	oint By *				
	STEAM CONDITIO	NS			CONSTRU	CTION			
	MAX.	NORMAL	MIN.	Turbine Typ. 0	Horiz. • V	erticel			
Inlet Press, PSIG				No. Stages	Wheel Dis., I	n.			
Inlet Temp, ^O F	()			Rotor: Built Up	Solid				
Exhaust Press (PSIG)	(In. Hg)			Bleding L 2 R o w L	<u></u>	Re Entry			
Unusual Conditions	(2.12.2.6)	<u>l</u>		Casing Split L Axia	al LI Redial	<u> </u>			
Duty 0 Continue	Sus 🔾 Standby 🤇	J Auto Start		Casing Support	Centerline	L Foot			
Evel. Steam Cost. \$/1	000 Lbs				NEMA 'P' 5250				
Payout Period, Years		/Yr		Trip Valve	ntegrai				
				nterstage <u>Seels</u>	Labyrinth	Carbon			
	e Speed, RPM			End Seals Carbon Hing, No/Box					
	ous spud. RPM								
Eirct Critical Spee	DDM			Type Madigi Bearings	0.21				
	Safe For Runaway Sne	ert (2 11 1)		Thrum Collar (2.9.8)	Replaceable		Non.		
Exh Temp. ^o F	Normal		NO LOR	Lube Oil Viscosity (2.1	(0.2) SUS	100% SI	JS @ 210°F		
Potential Max. Pov	er BHP			Lubrication O Bind	Oiled 0	Pressure			
Max, Nozzle Steem	Flow. Lbs/Hr			0 Purg	OIL Mist 0	ure Milt			
Max. Allowable Sc	BEC. RPM			O Shaft Areas Suitable	For Observing By	Non-Contactir	g Туре		
O Rotation Facing G	ov. End Occw) cw		Vibration Probes (2.6.2.2)					
O Driven Equipment	Thrust, Lbs. (2.9.3)		Up	CASING	DESIGN	INLET	EXHAUST		
(Vertical Turbine)			Down	Max. Allow. Press, PSI	G	•			
O Mount Turbine on	Baseplate Furnished by C	riven Equipm	ent Vendor	Max. Allow. Temp, ^o F					
O Water Piping Furn.	by Ovendor C	Others		Hydro Test Pressure, P	SIG				
O Oil Piping Furn. by	/ O Vendor C	Others		MATERIALS					
				High Pressure Casing					
				Exheust Casing					
	O SITE AND UTILITY			Nozzies					
Location (2.1.13)	O Indoor O Heated		eated	Blading					
	Uutdoor U Roof		nout Hoof	Wheels					
Ambient Temp. **		x.		Shert					
Unusual Conditions	O Duet O Salt At	mosphere		Under Packing	Application Me	100			
Elect Area (2.1.12)	Class C-		Div	GUV. VAIVE ITIM		eth Size			
LIUCI, MIUS (2.1.12)	Nos-beredour	, up					alay		
Control Power			Hz		li Speed Range	+%	-%		
Aux. Motors	V Ph		Hz	Speed Changer	Manual OPn	eum. O FI	ect.		
Cooling Water: Press	PSIG A	P, PSI		Mfr. Model					
Flow	, GPM 🛆	т, ^о ғ		· · · · · · · · · · · · · · · · · · ·					
Alloweble Sound Leve	I (2.1.11) di	BA OP	ft.	Turbine Weight, Lbs					

^a Also available in SI units (API Standard 611, January 1982). (Reprinted courtesy of the American Petroleum Institute.)



Figure 4.2. Combustion gas turbine arrangements and their thermodynamic diagrams. (a) Basic unit with PV and TS diagrams. (b) Unit with an air preheater and TS diagram.

EXAMPLE 4.2 Performance of a Combustion Gas Turbine

Atmospheric air at 80° F (305K) is compressed to 5 atm, combined with fuel at the rate of 1 kg/s, then expanded to 1 atm in a power



turbine. Metallurgical considerations limit the temperature to $1700^\circ F$ (1200K). The heat capacities of air and combustion products are

$$C_p = 0.95 + 0.00021T$$
 (K) kJ/kg

the heat of combustion is 42,000 kJ/kg, the furnace efficiency is 0.975, the isentropic efficiency of the compressor is 0.84, and that of the expander is 0.89. Find

a. the required air rate,

- b. the power loads of the compressor and expander, and
- c. the overall efficiency as a function of the temperature of the exhaust leaving a steam generator.

Poi nt	P	T _s	Т
1	1		305
2	5	483	517
3	5		1200
4	1	802	846
5	1		400
5	1		400

Compression:

$$k = 1.4, k/(k - 1) = 3.5,$$

$$T_{2s} = T_1 (P_2 / P_1)^{1/3.5} = 305 (5)^{1/3.5} = 483 \text{K},$$

 $T_2 = 305 + \frac{483 - 305}{0.84} = 517 \text{ K}.$

Combustion:

$$m'_{a} = \text{flow rate of air, kg/kg fuel}$$

$$0.975(42000) = \int_{305}^{1200} C_{p} dT + m'_{a} \int_{517}^{1200} C_{p} dT$$

$$= 991682 + 771985 m'_{a}$$

$$m'_{a} = 51.8$$

u Expansion:

k = 1.33, k/(k - 1) = 4.0 $T_{4s} = T_3(P_4/P_1)^{0.25} = 1200(0.2)^{0.25} = 802^{\circ}K$ $T_4 = 1200 - 0.89(1200 - 802) = 846^{\circ}K$

Power calculations:

Compressor:
$$w'_{c} = -m'_{a}\Delta H = -51.8 \frac{517}{I_{305}} C_{p} dT$$

 $= -51.8(216.98) = -11.240 \text{ kJ/s}$
Expander: $w'_{e} = -52.8 \int_{1200}^{517} C_{p} dT = 52.8(412.35) = 21,772 \text{ kJ/s}$
Steam generator: Q' = $52.8 \frac{1}{I_{T}} C_{p} dT$
 η_{t} = overall efficiency = $\frac{21772 - 11380 + Q'}{42000}$

The tabulation shows efficiency with three different values of the exhaust temperature.

Т	Q'	η _t		
846	0	0. 247		
600	14311	0. 588		
500	19937	0. 722		

68 DRIVERS FOR MOVING EQUIPMENT

Other inefficiencies are due to pressure drops of 2-5%, loss of 1-3% of the enthalpy in the expander, and 1% or so loss of the air for cooling the turbine blades. The greatest loss of energy is due to the necessarily high temperature of the exhaust gas from the turbine, so that the overall efficiency becomes of the order of 20% or so. Some improvements are effected with air preheating as on Figure 4.2(b) and with waste heat steam generators as in Example 4.2. In many instances, however, boilers on 1000°F waste gas are

economically marginal. Efficiencies are improved at higher pressure and temperature but at greater equipment cost.

Inlet temperature to the expander is controlled by the amount of excess air. The air/fuel ratio to make 1700° F is in the range of 50 lb/lb. Metallurgical considerations usually limit the temperature to this value. Special materials are available for temperatures up to 2200°F but may be too expensive for process applications.

REFERENCES

- 1. M.P. Boyce, Gas Turbine Engineering Handbook, Gulf, Houston, 1982.
- 2. F.L. Evans, Equipment Design Handbook for Refineries and Chemical Plants, Gulf, Houston, 1979, vol. 1.
- 3. H. Gartmann, De Laval Engineering Handbook, McGraw-Hill, New York, 1970.

- 4. R.T.C. Harman, Gas Turbine Engineering, Macmillan, New York, 1981.
- 5. E.E. Ludwig, Applied Process Design for Chemical and Process Plants, Gulf, Houston, 1983, vol. 3.
- Marks' Standard Handbook for Mechanical Engineers, McGraw-Hill, New York, 1987.

TRANSFER OF SOLIDS

n contrast to fluids which are transferred almost exclusively through pipelines with pumps or blowers, a greater variety of equipment is employed for moving so/ids to and from storage and between process equipment. Most commonly, solids are carried on or pushed along by some kind of conveyor. So/ids in granular form also are transported in pipelines as slurries in inert liquids or as suspensions in air or other gases.

5.1. SLURRY TRANSPORT

In short process lines slurries are readily handled by centrifugal pumps with large clearances. When there is a distribution of sizes, the fine particles effectively form a homogeneous mixture of high density in which the settling velocities of larger particles are less than in clear liquid. Turbulence in the line also helps to keep particles in suspension. It is essential, however, to avoid dead spaces in which solids could accumulate and also to make provisions for periodic cleaning of the line. A coal-oil slurry used as fuel and acid waste neutralization with lime slurry are two examples of process applications.

Many of the studies of slurry transfer have been made in connection with long distance movement of coal, limestone, ores, and others. A few dozen such installations **have** been made, in length from several miles to several hundred miles.

Coal-water slurry transport has been most thoroughly investigated and implemented. One of the earliest lines was 108 miles long, 10 in. dia, 50-60 wt % solids up to 14 mesh, at velocities of 4.5–5.25 ft/sec, with positive displacement pumps at 30-mile intervals. The longest line in the United States is 273 miles, 18in. dia and handles 4.8-6.0 million tons/yr of coal; it is described in detail by Jacques and Montfort (1977). Other slurry pipeline literature is by Wasp, Thompson, and Snoek (1971), Bain and Bonnington (1970), Ewing (1978), and Zandi (1971).

Principally, investigations have been conducted of suitable linear velocities and power requirements. Slurries of 40-50 vol % solids can be handled satisfactorily, with particle sizes less than 24-48 mesh or so (0.7-0.3 mm). At low line velocities, particles settle out and impede the flow of the slurry, and at high velocities the frictional drag likewise increases. An intermediate condition exists at which the pressure drop per unit distance is a minimum. The velocity at this condition is called a critical velocity of which one correlation is

$$u_c^2 = 34.6C_v Du_t \sqrt{g(s-1)/d}, \qquad \text{consistent units}, \qquad (5.1)$$

where

 u_c = critical flow velocity,

 u_t = terminal settling velocity of the particle, given by Figure 5.1,

 C_n = volume fraction of solids,

D = pipe diameter,

d = particle diameter,

- s = ratio of densities of solid and liquid,
- $g = acceleration of gravity, 32.2 \text{ ft/sec}^2$, or consistent units.

The numerical coefficient is due to Hayden and Stelson (1971).

Another criterion for selection of a flow rate is based on considerations of the extent of sedimentation of particles of various sizes under flow conditions. This relation is developed by Wasp, Aude, Seiter, and Thompson (1971),

$$\frac{C}{C_{\rm P}} = \exp(-2.55u_t/ku\sqrt{f}), \qquad (5.2)$$

where

- C = concentration of a particular size at a level 92% of the vertical diameter,
- C_0 = concentration at the center of the pipe, assumed to be the same as the average in the pipe,

f = Fanning friction factor for pipe flow

$$= 0.25 \frac{\Delta P}{\rho} / \frac{L}{D 2g_c}$$
(5.3)

At high Reynolds numbers, for example, Blasius' equation is

$$f = 0.0791/N_{\rm Re}^{0.25}, \qquad N_{\rm Re} \ge 10^5$$
 (5.4)

k in Eq. (5.2) is a constant whose value is given in this paper as 0.35, but the value 0.85 is shown in a computer output in a paper by Wasp, Thompson, and Snoek (1971, Fig. 9). With the latter value, Eq. (5.2) becomes

$$C/C_0 = \exp(-3.00u_t/u\sqrt{f}).$$
 (5.5)

The latter paper also states that satisfactory flow conditions prevail when $C/C_0 \ge 0.7$ for the largest particle size. On this basis, the minimum line velocity becomes

$$u = \frac{3u_t}{\sqrt{f}\ln(C_0/C)} = 8.41u_t/\sqrt{f},$$
(5.6)

where u, is the settling velocity of the largest particle present.

As Example 5.1 shows, the velocities predicted by Eqs. (5.1) and (5.6) do not agree closely. Possibly an argument in favor of Eq. (5.6) is that it is proposed by the organization that designed the successful 18 in., 273 mi Black Mesa coal slurry line.

Pressure drop in flow of aqueous suspensions sometimes has been approximated by multiplying the pressure drop of clear liquid at the same velocity by the specific gravity of the slurry. This is not borne out by experiment, however, and the multiplier has been correlated by other relations of which Eq. (5.7) is typical:

$$\Delta P_s / \Delta P_L = 1 + 69C_v \left[\frac{gD(s-1)}{u^2 \sqrt{C_D}} \right]^{1.3}.$$
(5.7)

This equation is a modification by Hayden and Stelson (1971) of a series of earlier ones. The meanings of the symbols are

- C_{v} = volume fraction occupied by the solids in the slurry,
- d = particle diameter,

D = pipe diameter,

s = ratio of specific gravities of solid and liquid.

EXAMPLE 5.1 Conditions of a Coal Slurry Pipeline Data of a pulverized coal slurry are

 $C_{v} = 0.4,$ D = 0.333 ft, f = 0.0045 (Blasius' eq. at $N_{re} = 10^{5}),$ s = 1.5.Mesh size 24 48 100

<i>d</i> (mm)	0. 707	0. 297	0. 125	0. 321
Weight fraction	0.1	0.8	0.1	1
u, (ft/sec)	0.164	0.050	0.010	0.0574

Mixture

The terminal velocities are read off Figure 5.1, and the values of the mixture are weight averages.

The following results are found with the indicated equations:

item	Eq.	24	48 100 Mixture
и _с	5.1	7.94	5. 45 3. 02
	ii ii 5. 11 5. 13	20.6 1.36	6.27 2.89 9.38 1.25 3.39 1.539 1.296

Eq. (5.1):
$$u_c^2 = 34.6(0.4)(0.333)\sqrt{32.2(0.5)} \frac{u_t}{\sqrt{d_{mm}}/304.8}$$

 $= 323 \frac{u_t}{\sqrt{d_{mm}}},$
Eq. (5.6): $u = \frac{8.41u_t}{\sqrt{0.0045}} = 125u_t,$
Eq. (5.8): $c_D = \frac{4}{3} \frac{32.2(1.5-1)}{u_t^2} \frac{d_{mm}}{304.8} = \frac{0.0704d_{mm}}{u_t^2},$
Eq. (5.11): $\frac{\Delta P_s}{\Delta P_L} = 1 + \frac{0.69}{0.4^{0.3}} \left[\frac{1}{0.0574} \sqrt{\frac{32.2(0.5)0.321}{304.8(3.39)^2}} \right]^{1.3}$
 $= 1.5391,$
Eq. (5.13): $\frac{\Delta P_s}{\Delta P_L} = 1 + 0.272(0.4) \left[\frac{0.0045(0.333)32.2(0.5)}{(0.0574)^2(3.39)} \right]^{1.3}$

With coal of sp gr = 1.5, a slurry of 40 vol % has a sp gr = 1.2. Accordingly the rule, $\Delta P_s / \Delta P_L = \text{sp gr}$, is not confirmed accurately by these results.

The drag coefficient is

$$C_D = 1.333gd(s-1)/u_t^2.$$
(5.8)

For mixtures, a number of rules has been proposed for evaluating the drag coefficient, of which a weighted average seems to be favored,

$$\sqrt{C_D} = \sum w_i \sqrt{C_{Di}},\tag{5.9}$$

where the w_i are the weight fractions of particles with diameters d_i .



Figure 5.1. Settling velocities of spheres as a function of the ratio ot densities of the two phases. Stokes law applies at diameters below approximately 0.01 cm (*based on* a chart of *Lapple* et al., Chemical Engineering Handbook, *McGraw-Hill, New York, 1984, p. 5.67*).

For particles of one size, Eqs. (5.7) and (5.8) combine to

$$\Delta P_s / \Delta P_L = 1 + 100 C_v [(u_t D/u^2) \sqrt{g(s-1)/d}]^{1.3},$$

consistent units. (5.10)

The pressure drop relation at the critical velocity given by Eq. (5.1) is found by substitution into Eq. (5.7) with the result

$$\Delta P_s / \Delta P_L = 1 + \frac{0.69}{C_{\nu}^{0.3}} [(1/u_t) \sqrt{gd(s-1)/C_D}]^{1.3}.$$
 (5.11)

With Eq. (5.10) the result is

$$\Delta P_s / \Delta P_L = 1 + 1 / C_v^{0.3}. \tag{5.12}$$

With the velocity from Eq. (5.6), Eq. (5.7) becomes

$$\Delta P_s / \Delta P_L = 1 + 0.272 C_v [fgD(s-1)/u_t^2 \sqrt{C_D}]^{1.3}$$
(5.13)



Figure 5.2. Non-Newtonian behavior of suspensions: (a) viscosity as a function of shear rate, 0.4 wt % polyacrylamide in water at room temperature; (b) shear stress as a function of shear rate for suspensions of TiO_2 at the indicated vol % in a 47.1 wt % sucrose solution whose viscosity is 0.017 Pa sec (Denn, Process Fluid Mechanics, Prentice-Hall, Englewood Cliffs, NJ, 1980).

and, for one-sized particles,

$$\Delta P_s / \Delta P_L = 1 + 0.394 C_v [(fD/u_t) \sqrt{g(s-1)/d}]^{1.3}.$$
 (5.14)

These several pressure drop relations hardly appear consistent, and the numerical results of Example 5.1 based on them are only roughly in agreement.

From statements in the literature, it appears that existing slurry lines were designed on the basis of some direct pilot plant studies.

Nonsettling slurries are formed with fine particles or plastics or fibers. Although their essentially homogeneous nature would appear to make their flow behavior simpler than that of settling slurries, they often possess non-Newtonian characteristics which complicate their flow patterns. In Newtonian flow, the shear stress is proportional to the shear strain,

stress =
$$\mu(\text{strain})$$
,

but in other cases the relation between these two quantities is more complex. Several classes of non-Newtonian behavior are recognized for suspensions. Pseudoplastic or power-law behavior is represented **by**

stress =
$$k(\text{strain})^n$$
, $n < 1$

where k is called the consistency index. Plastic or Bingham behavior is represented by

stress =
$$k_1 + \eta$$
(strain)

where η is called the plastic viscosity. Data for some suspensions are given on Figure 5.2.

The constants of such equations must be found experimentally over a range of conditions for each particular case, and related to the friction factor with which pressure drops and power requirements can be evaluated. The topic of nonsettling slurries is treated by Bain and Bonnington (1970) and Clift (1980). Friction factors of power-law systems are treated by Dodge and Metzner (1959) and of fiber suspensions by Bobkowitz and Gauvin (1967).

5.2. PNEUMATIC CONVEYING

Granular solids of free-flowing natures may be conveyed through ducts in any direction with high velocity air streams. In the normal plant, such lines may be several hundred feet long, but dusty materials such as fly ash and cement have been moved over a mile in this way. Materials that are being air-veyed include chemicals, plastic pellets, grains, and powders of all kinds. The transfer of catalysts between regenerator and reactor under fluidized conditions is a common operation. Stoess (1983) has a list of recommendations for about 150 different materials, of which Table 5.1 is a selection. Basic equipment arrangements are represented in Figure 5.3.

The performance of pneumatic conveyors is sensitive to several characteristics of the solids, of which the most pertinent ones are

- 1. bulk density, as poured and as aerated,
- 2. true density,
- 3. coefficient of sliding friction (= tangent of the angle of repose),
- 4. particle size distribution,
- 5. particle roughness and shape,
- 6. moisture content and hygroscopicity, and
- 7. characteristics such as friability, abrasiveness, flammability, etc.

Sulfur, for example, builds up an electrostatic charge and may introduce explosive risks.

In comparison with mechanical conveyors, pneumatic types must be designed with greater care. They demand more power input per unit weight transferred, but their cost may be less for complicated paths, when exposure to the atmosphere is undesirable and when operator safety is a problem. Although in the final analysis the design and operation of pneumatic conveyors demands the attention of experienced engineers, a design for orientation purposes can be made by the inexpert on the basis of general knowledge and rules of thumb that appear in the literature. An article by Solt (1980) is devoted entirely to preventive troubleshooting.

Some basic design features are the avoidance of sharp bends, a minimum of line fittings, provision for cleanout, and possibly electrical grounding. In many cases equipment suppliers may wish to do pilot plant work before making final recommendations. Figure

	Va	acuı	ım Sys	stem	(8-9) ps:	ia)				_	Low P	ressu	re	Syste	em (6	-12	psig)
	Conveying Distance					Conveying Distance												
	Wt	π	() ft	150) ft	4	250 ft	40	0 ft	Valuaitu	Protestra	10	() ft	25	50 ft	4	00 ft	- Velocity
Material	cu f	t S	at.	hp/1	' Sat.	hp/1	' Sat. hp	o∕r Sa	it hp/I	ft/sec	Factor	Sat	hp/I	'Sa	t.h	p/ľ S	a t	hp/[(ft/sec)
Alum	50	3.6	4.5	3.9	5.0	4.3	57	4.7	6.3	110	4.0	1.6	2.7	2.0	3.4	2.2	3.8	65
Alumina	60	2.4	4.0	2.8	4.7	3.4	5.7	4.0	6.4	105	5.0	1.1	2.4	1.6	3.4	1.9	3.9	6 0
Carbonate, calcium	25-30	3.1	42	3.1	5.0	3.9	5.5	4.2	6 ()	110	3.5	1.4	2.5	1.8	3.3	2.0	3.6	6 5
Cellulose acetate	22	3.2	4.7	3.5	5.1	3.8	5.7	4.1	6.0	100	3.0	1.4	28	17	34	19	36	55
Clay. air floated	30	3.3	4.5	3.5	5.0	3.9	5.5	4.2	6.0	105	4.0	15	2.0	1.8	33	19	3.6	50
Clay, water washed	40-50	3.5	5.0	3.8	5.6	4.2	6.5	4.5	7.2	115	4.5	1.6	3.0	1.9	3.9	2.1	4.4	60
Clay. spray dried	60	3.4	4.7	3.6	5.2	4.0	6.2	4.4	7.1	110	4.3	1.5	2.8	1.8	3.7	2.0	4.3	5 5
Coffee beans	4 2	1.2	2.0	1.6	3.0	2.1	3.5	2.4	4.2	75	5.0	0.6	1.2	0.9	2.1	1.1	2.5	4 5
Corn, shelled	45	19	2.5	2.1	2.9	2.4	3.6	2.8	4.3	105	5.0	0.9	1.5	1.1	2.2	1.3	2.6	55
Flour. wheat	4 0	1.5	30	1.7	3.3	20	3.7	2.5	44	90	2.5	0.7	18	0.9	22	1.1	2.7	35
Grits, corn	33	1.7	25	2.2	3.0	2.9	4.0	3.5	4.8	100	3.5	0.8	1.5	1.3	2.4	1.6	2.9	70
Lime, pebble	56	2.b	38	3.0	4.0	3.4	4.7	3.9	5.4	105	5.0	1.3	2.3	1.6	2.8	1.8	3.3	70
Lime, hydrated	30	2.1	3.3	2.4	3.9	2.8	4.7	3.4	6.0	90	4.0	0.6	1.x	0.8	2.2	0.9	2.6	4 0
Malt	28	1.8	2.5	2.0	2.8	2.3	3.4	2.8	4.2	100	5.0	0.8	15	1.1	2.0	1.3	2.5	55
Oats	25	2.3	3.0	2.6	3.5	3.0	4.4	3.4	5.2	100	5.0	1.0	1.8	1.4	2.6	1.6	3.1	55
Phosphate, trisodium	65	3.1	4.2	3.6	5.0	3.9	5.5	4.2	6.0	110	4 5	1.4	2.5	1.8	3.3	1.9	3.6	75
Polyethylene	30	1.2	2.0	1.6	3.0	2.1	3.5	2.4	4.2	80	50	0.55	1.2	0.9	2.1	1.1	2.5	70
Rubber pellets	40	2.9	4.2	3.5	5.0	4.0	6.0	4.5	7.2	110								
Salt cake	90	4.0	6.5	4.2	6.8	4.6	i.5	5.0	8.5	120	5.0	2.9	3.9	3.5	4.5	4.0	5.1	83
Soda ash, light	35	3.1	4.2	3.6	50	3.9	5.5	4.2	6.0	110	5.0	1.4	2.5	1.4	3.3	1.9	3.6	65
Soft feeds	20-40	3.0	4.2	34	4.5	3.7	5.0	4.2	5.5	110	3.8	1.3	2.5	1.7	3.1	1.9	3.7	70
Starch. pulverized	40	1.7	3.0	2.0	3.4	2.6	4.0	3.4	5 U	90	30	0.8	1.7	1.1	2.4	1.5	3.0	5 5
Sugar, granulated	50	3.0	3.7	32	4.0	ii	5.2	39	6.U	110	5.0	14	2.2	1.6	3.1	1.7	3.6	60
Wheat Wood flour	48 12-20	1.9 2.5	2.5 3 5	2.1 2 8	2.9 4.0	2.4 3.4	3.6 4.9	2.8 4.4	4.3 6.5	105 100	50	0.9	1.5	1.1	2.1	1.3	2.6	55

TABLE 5.1. Flow Rates and Power Requirements of Vacuum and Low Pressure Pneumatic Conveying Systems*

"HP/ton = (pressure factor)(hp/T)(sat.). The units of sat. are standard cuft of air/lb of solid transferred), and those of hp/T are horsepower/(tons/hr of solid transferred).

(Stoess, 1983).

5.4 shows a typical pilot plant arrangement. A preliminary design procedure is given by Raymus (1984). Many details of design and operation are given in books by Stoess (1983) and Kraus (1980) and in articles by Gerchow (1980), and Perkins and Wood (1974). Some of that information will be restated here. Pressure drop and power requirements can be figured largely on the basis of general knowledge.

EQUIPMENT

The basic equipment consists of a solids feeding device, the transfer line proper, a receiver, a solid-air separator, and either a blower at the inlet or a vacuum pump at the receiver. Four common kinds of arrangements are shown on Figure 5.3. Vacuum systems are favored for shorter distances and when conveying from several sources to one destination. Appropriate switching valves make it possible to service several sources and destinations with either a vacuum or pressure system. Normally the vacuum system is favored for single destinations and the pressure for several destinations or over long distances. Figure 5.3(b) shows a rotary valve feeder and Figure 5.3(c) a Venturi feeder which has a particularly gentle action suitable for friable materials. Figure 5.3(d) utilizes a fan to suck the solids from a source and to deliver them under positive pressure. Friable materials also may be handled effectively by the equipment of Figure 5.5 in which alternate pulses of granular material and air are transported.

Typical auxiliary equipment is shown on Figure 5.6. The most used blower in pneumatic conveying is the rotary positive displacement type; they can achieve vacua 6-8 psi below atmospheric or positive pressures up to 15 psig at efficiencies of about 65%. Axial positive displacement blowers also are used, as well as **centrifugals** for large capacities. Rotary feeders of many proprietary designs are available; Stoess (1983) and Kraus (1980) illustrate several types. Receivers may be equipped with fabric filters to prevent escape of fine particles; a **dacron** fabric suitable for up to 275° F is popular. Cyclone receivers are used primarily for entirely nondusting services or ahead of a filter. A two-stage design is shown in Figure 5.6(d). Typical dimensions are cited by Stoess (1983), for example:

line diameter (in.)	3	5	8
primary diameter (ft)	3.5	4.5	6.75
secondary diameter (ft)	2.75	3.5	5.0



Figure 5.3. Basic equipment arrangements of pneumatic conveying systems. (a) Vacuum system with several sources and one destination, multiple pickup; (b) pressure system with rotary valve feeder, one source and several destinations, multiple discharge; (c) pressure system with Venturi feed for friable materials; (d) pull-push system in which the fan both picks up the solids and delivers them [*after F. J. Gerchow*, Chem. Eng. (17 *Feb. 1975, p. 88*)].

Piping usually is standard steel, Schedule 40 for 3-7 in. IPS and Schedule 30 for 8-12 in. IPS. In order to minimize pressure loss and abrasion, bends are made long radius, usually with radii equal to 12 times the nominal pipe size, with a maximum of 8ft. Special reinforcing may be needed for abrasive conditions.

OPERATING CONDITIONS

Vacuum systems usually operate with at most a 6 psi differential; at lower pressures the carrying power suffers. With rotary air lock

feeders, positive pressure systems are limited to about 12 psig. Other feeding arrangements may be made for long distance transfer with 90-125 psig air. The dense phase pulse system of Figure 5.4 may operate at 10-30 psig.

Linear velocities, carrying capacity as **cuft** of free air per lb of solid and power input as HP/tons per hour (tph) are listed in Table 5.1 as a general guide for a number of substances. These data are for 4-, 5-, and 6-in. lines; for 8-in. lines, both Sat. and HP/tph are reduced by 15%, and for 10-in. by 25%. Roughly, air velocities in low positive pressure systems are 2000 ft/min for light materials,



Figure 5.4. Sketch of pilot plant arrangement for testing pneumatic conveying under positive pressure (*Kraus*, Pneumatic Conveying of Bulk Materials, *McGraw-Hill, New York, 1980*).

3000-4000 ft/min for medium densities such as those of grains, and 5000 ft/min and above for dense materials such as fly ash and cement; all of these velocities are of free air, at atmospheric pressure.

Another set of rules for air velocity as a function of line length



Figure 5.5. Concept of dense phase transfer of friable materials, by intermittent injection of material and air pulses, air pressures normally 10–30 psig and up to 90 psig (*Sturtevant* Engineering Co., *Boston, MA*).

and bulk density is due to Gerchow (1980) and is

line length	ft/min					
(ft)	55 lb/cuft	55-85	85-115			
200	4000	5000	6000			
500	5000	6000	7000			
1000	6000	7000	8000			

Conveying capacity expressed as vol % of solids in the stream usually is well under 5 vol %. From Table 5.1, for example, it is about 1.5% for alumina and 6.0% for polystyrene pellets, figured at atmospheric pressure; at 12psig these percentages will be roughly doubled, and at subatmospheric pressures they will be lower.

POWER CONSUMPTION AND PRESSURE DROP

The power consumption is made up of the work of compression of the air and the frictional losses due to the flows of air and solid through the line. The work of compression of air at a flow rate m'_a and $C_P/C_v = 1.4$ is given by

$$w_c = 3.5(53.3)(T + 460)m'_a[(P_2/P_1)^{0.2857} - 1] \qquad \text{(ft lbf/sec)}$$
(5.15)

with the flow rate in lb/sec.

Frictional losses are evaluated separately for the air and the solid. To each of these, contributions are made by the line itself, the elbows and other fittings, and the receiving equipment. It is conservative to assume that the linear velocities of the air and solid are the same. Since the air flow normally is at a high Reynolds number, the friction factor may be taken constant at $f_a = 0.015$. Accordingly the frictional power loss of the air is given by

$$w_1 = \Delta P_1 m'_a / \rho_a = (u^2/2g) \bigg[1 + 2n_c + 4n_f + (0.015/D) \bigg(L + \sum L_i \bigg) \bigg] m'_a$$
(ft lbf/sec). (5.16)

The unity in the bracket accounts for the entrance loss, n_c is the number of cyclones, n_f is the number of filters, L is the line length, and L_i is the equivalent length of an elbow or fitting. For long radius bends one rule is that the equivalent length is 1.6 times the actual length of the bend. Another rule is that the long bend radius is 12 times the nominal size of the pipe. Accordingly,

$$L_i = 1.6(\pi R_i/2) = 2.5R_i = 2.5D''_i$$
 ft, with D''_i in inches.
(5.17)

The value of g is 32.2 ft lb $m/(lbf sec^2)$.

The work being done on the solid at the rate of m'_s lb/sec is made up of the kinetic gain at the entrance (w_2) , the lift (w_3) through an elevation Az, friction in the line (w_4) , and friction in the elbow (ws). Accordingly,

$$w_2 = \frac{u^2}{2g}m'_s$$
 (ft lbf/sec) . (5.18)

The lift work is

$$w_3 = \Delta z \frac{g}{g_c} m'_s = \Delta z m'_s \quad \text{(ft lbf/sec)}. \tag{5.19}$$

The coefficient of sliding friction f_s of the solid equals the tangent of the angle of repose. For most substances this angle is 30-45" and

5.2. PNEUMATIC CONVEYING 75



Figure 5.6. Components of pneumatic conveying systems. (a) Rotary positive displacement blower for pressure or vacuum. (b) A rotary airlock feeder for fine materials (*Detroit Stoker* Co.). (c) A four-compartment receiver-filter (Fuller Co., Bethlehem, *PA*). (d) A two-stage cyclone receiver. (e) The Fuller-Kinyon pump for cement and other fine powders. Powder is fed into the aeration chamber with a screw and is fluidized with compressed air (Fuller Co., Bethlehem, *PA*).

76 TRANSFER OF SOLIDS

the value off, is 0.58-1.00. The sliding friction in the line is

$$w_4 = f_s Lm'_s \quad \text{(ft lbf/sec)} \quad , \tag{5.20}$$

where *L* is the line length.

Friction in the curved elbows is enhanced because of centrifugal force so that

$$w_5 = f_s \frac{u^2}{gR} \left(\frac{2\pi R}{4}\right) m'_s = 0.0488 f_s u^2 m'_s \quad \text{(ft lbf/sec)} \tag{5.21}$$

The total frictional power is

$$w_f = w_1 + w_2 + w_3 + w_4 + w_5, \tag{5.22}$$

and the total power consumption is

$$w = \frac{(w_c + w_f)}{550\eta(1.8m'_s)} \text{ [HP/(ton/hr)]}, \tag{5.23}$$

where η is the blower efficiency. Pressure drop in the line is obtained from the frictional power, the total flow rate, and the density of the mixture:

$$AP = \frac{w_f}{144(m'_a + m'_s)} \rho_m \quad \text{(psi)}. \tag{5.24}$$

The specific air rate, or saturation, is

saturation =
$$0.7854(60)D^2$$

(cuft/min of air)/(lb/min of solid)], (5.25)

where the velocity of the air is evaluated at atmospheric pressure.

Example 5.2 makes the calculations described here for power and pressure drop, and compares the result with the guidelines of Table 5.1.

5.3. MECHANICAL CONVEYORS AND ELEVATORS

Granular solids are transported mechanically by being pushed along or dragged along or carried. Movement may be horizontal or vertical or both. In the process plant distances may be under a hundred feet or several hundred feet. Distances of several miles may be covered by belts servicing construction sites or mines or power plants. Capacities range up to several hundred tons/hr. The principal kinds of mechanical conveyors are illustrated in Figures 5.7-5.13 and will be described. Many construction features of these machines are arbitrary. Thus manufacturers' catalogs are the ultimate source of information about suitability for particular services, sizes, capacities, power requirements and auxiliaries, Much of the equipment has been made in essentially the present form for about 100 years by a number of manufacturers so that a body of standard practice has developed.

PROPERTIES OF MATERIALS HANDLED

The physical properties of granular materials that bear particularly on their conveying characteristics include size distribution, true and bulk densities, and angle of repose or coefficient of sliding friction, but other less precisely measured or described properties are also of concern. A list of pertinent properties appears in Table 5.2. The elaborate classification given there is applied to about 500 materials in the FMC Corporation Catalog 100 (1983, pp. **B.27–B.35**) but is too extensive for reproduction here. For each material the table also identifies the most suitable design of screw conveyor of this company's manufacture and a factor for determining the power requirement. An abbreviated table of about 150 substances appears in the *Chemical* Engineers Handbook (1984, **p**. 7.5). Hudson (1954, **pp. 6–9)**, describes the characteristics of about 100 substances in relation to their behavior in conveyors. Table 5.3 lists bulk densities, angles of respose at rest, and allowable angles of inclination which are angles of repose when a conveyor is in motion; references to more extensive listings of such data are given in this table.

The angle of repose is a measure of the incline at which conveyors such as screws or belts can carry the material. The tangent of the angle of repose is the coefficient of sliding friction. This property is a factor in the power needed to transfer the material by pushing or dragging as in pneumatic, screw, flight, and Redler equipment.

Special provisions need to be made for materials that tend to form bridges; Figure 5.13(a) is an example of a method of breaking up bridges in a storage bin so as to ensure smooth flow out. Materials that tend to pack need to be fluffed up as they are pushed along by a screw; adjustable paddles as in Figure 5.7(d) may be sufficient.

SCREW CONVEYORS

These were invented by Archimedes and assumed essentially their present commercial form a hundred years or so ago. Although the equipment is simple in concept and relatively inexpensive, a body of experience has accumulated whereby the loading, speed, diameter, and length can be tailored to the characteristics of the materials to be handled. Table 5.4, for example, recognizes four classes of materials, ranging from light, freeflowing, and nonabrasive materials such as grains, to those that are abrasive and have poor flowability such as bauxite, cinders, and sand. Only a portion of the available data are reproduced in this table.

Lengths of screw conveyors usually are limited to less than about 150 ft; when the conveying distance is greater than this, a belt or some other kind of machine should be chosen. The limitation of length is due to structural strength of the shaft and coupling. It is expressed in terms of the maximum torque that is allowable. Formulas for torque and power of screw conveyors are given in Table 5.4 and are applied to selection of a conveyor in Example 5.3.

Several designs of screws are shown in Figure 5.7. The basic design is one in which the pitch equals the diameter. Closer spacing is needed for carrying up steep inclines, and in fact very fine pitch screws operating at the relatively high speeds of 350 rpm are used to convey vertically. The capacity of a standard pitch screws drops off sharply with the inclination, for example:

Angle (degrees)	<8	20	30	45
Percent of capacity	100	55	30	0

Allowable loadings as a percentage of the vertical cross section depend on the kind of material being processed; examples are shown in Table 5.4.

BELT CONVEYORS

These are high capacity, relatively low power units for primarily horizontal travel and small inclines. The maximum allowable inclination usually is 5-15" less than the angle of repose; it is shown as "recommended maximum inclination" in Table 5.3 for some substances, and is the effective angle of repose under moving conditions.

The majority of conveyor belts are constructed of fabric, rubber, and wire beads similarly to automobile tires, but they are made also of wire screen or even sheet metal for high temperature

EXAMPLE 5.2

Size aud Power Requirement of a Pneumatic Transfer Line

A pneumatic transfer line has 300 ft of straight pipe, two long radius elbows, and a lift of 5Oft. A two-stage cyclone is at the receiving end. Solid with a density of 125 lb/cuft is at the rate of 10 tons/hr and the free air is at 5000 ft/min. Inlet condition is 27 psia and 100°F. Investigate the relation btween line diameter and power requirement.

On a first pass, the effect of pressure loss on the density of the air will be neglected.

Mass flow rate of solid:

 $m'_s = 20,000/3600 = 5.56$ lb/sec.

Mass flow rate of air:

$$m'_a = \frac{5000}{60} \frac{\pi}{4} (0.075) D^2 = 4.91 D^2 \, \text{lb/sec.}$$

Density of air:

$$\rho_a = 0.075 \left(\frac{27}{14.7}\right) = 0.138$$
 lb/cuft.

Density of mixture:

Pm =
$$\frac{(m'_a + m'_s)}{m'_a/\rho_a + m'_s/\rho_s}$$

= $\frac{(m'_a + 5.56)}{m'_a/0.138 + 5.56/125}$

Linear velocity of air at inlet:

$$u = \frac{5000}{60} \left(\frac{14.7}{27}\right) = 45.37$$
 fps.

Assume air and solid velocities equal. Elbow radius = 120. Elbow equivalent length,

$$L_e = 1.6(\pi/2)(12D) = 30.20$$

Power for compression from 14.7 psia and 560 R to 27 psia,

$$k/(k = 1) = 3.5,$$

$$w_c = 3.5RT_1[(P_2/P_1)^{0.2857} - 1]m'_a$$

$$= 3.5(53.3)(560)[(27/14.7)^{0.2857} - 1]4.91D^2$$

$$= 973050^\circ \text{ ft } \text{ lbf/sec.}$$

Frictional contribution of air

$$w_1 = \frac{u^2}{2g} [5 + (0.015/D)(300 + 2(30.2)D]m'_a$$

 $= [(45.4)^2/64.4][5.9 + (4.5/D)](4.91D^2)$ = 157.1D²(5.9 + 4.5'0)

For the solid, take the coefficient of sliding friction to be $f_s = 1$. Power loss is made up of four contributions. Assume no slip velocity;

 $w_s = w_2 + w_3 + w_4 + w_5$ = $[u^2/2g + AZ + f_sL + 2(0.0488)f_su^2]m'_s$ = 5.56[45.4²/64.4 + 50 + 300 + 2(0.0488)45.4²] = 3242.5ft lbf/sec.

Total friction power:

 $w_f = 3242.5 + 157.1D^2(5.9 + 4.5/D).$

Pressure drop:

$$\Delta P = \frac{W_f}{144(m_a^{\prime} + m_{,}^{\prime})} \rho_m \text{ psi}$$

Fan power at $\eta = 0.5$:

$$P = \frac{w_c + w_f}{550(0.5)(10)} = \frac{w_c + w_f}{2750} \text{HP/tph},$$

seturation = $\frac{5000(\pi/4)D^2}{1100} = 11.78D^2 \text{SCEM}/(10/min)$

saturation = $\frac{5000(\pi/4)D^2}{20,000/60} = 11.78D^2 \text{ SCFM}/(\text{lb/min}).$

1	PS D	(ft)	m'a	ρ _m W _c	W _f
3	0. 2557	0.3210	2.4808	6362	3484
4	0. 3356	0.5530	1.5087	10, 959	3584
5	0.4206	0.8686	1.0142	17, 214	3704
6	0.5054	1.2542	0.7461	24, 855	3837

1 PS	ΔP (psi)	НР/ТРН	SCFM/ lb/min
3	10. 2	3. 58	0.77
4	6.1	5.29	1.33
5	4.1	7.60	2.08
6	2.9	10.44	3.00

From Table 5.1, data for pebble lime are

sat = 1.7 SCFM/(lb/min) power = 3.0 HP/TPH

and for soda ash:

sat = 1.9 SCFM/(lb/min)power = 3.4 HP/TPH.

The calculated values for a 4in. line are closest to the recommendations of the table.

services. A related design is the apron conveyor with overlapping pans of various shapes and sizes (Fig. 5.8), used primarily for short travel at elevated temperatures. With pivoted deep pans they are **also** effective elevators.

Flat belts are used chiefly for moving large objects and cartons.

For bulk materials, belts are **troughed** at angles of 20-45". Loading of a belt may be accomplished by shovelling or directly from overhead storage or by one of the methods shown on Figure 5.9. Discharge is by throwing over the end of the run or at intermediate points with plows.

TABLE 5.2. Codes for Characteristics of Granular Materials"

Major <u>Class</u>	Material Characteristics Included	Code Designation
Density	Bulk Density , Loose	Actual ibs/ft ³
	No. 200 Sieve (.0029") And Under VeryFine No 100 Sieve 1.0059") And Under No. 40 Sieve (.016") And Under	A ₂₀₀ A ₁₀₀ A ₄₀
	Fine NO 6 Sieve (.132") And Under	Bo
Size	Granular ½" And Under Granular 3" And Under	С _% D ₃
	(1)Lumpy Over 3" To Be Special X=Actual Maximum Size	D _x
	Irregular Stringy, Fibrous. Cylindrical, Slabs. etc.	F
Flowability	Very Free Flowing-Flow Funchon > 10 Free Flowing – Flow Funchon > 4 But < 10 Average Flowability – Flow Function 2 But < 4 Sluggish-Flow Funchon < 2	1 2 3 4
Abrasiveness	Mildly Abrasive — Index 1-17 Moderately Abrasive — Index 18-67 Extremely Abrasive — Index 68-416	5 6 7
Miscellaneous Properties Or Hazards	Builds Up and Hardens Generates Static Electricity Decomposes Deteriorates In Storage Flammability Becomes Plastic or Tends to Soften Very Dusty Aerates and Becomes Fluid Explosiveness Stickness-Adhesion Contaminable, Affecting Use Degradable, Affecting Use Grives Off Harmful or Toxic Gas or Fumes Highly Corrosive Middly Corrosive Mygroscopic Interlocks. Mats or Agglomerates Oils Present Packs Under Pressure Very Light and Fluffy-May Be Windswept Elevated Temperature	F G ∺ J ≚ J ≚ J X Y Q P Q R S ⊺ J ≥ ≥ X Y Z

'Example: A fine 100 mesh material with an average density of 50 lb/cuft that has average flowability and is moderately abrasive would have a code designation $50A_{100}36$; if it were dusty and mildly corrosive, it would be $50A_{100}36LT$.

(FMC Corp., Materials Handling Division, Homer City, PA, 1963).

Power is required to run the empty conveyor and to carry the load horizontally and vertically. Table 5.5 gives the equations, and they are applied in Example 5.4. Squirrel-cage ac induction motors are commonly used as drives. Two- and four-speed motors are available. Mechanical efficiencies of speed reducing couplings between motor and conveyor range from 95 to 50%. Details of idlers, belt trippers, cleaners, tension maintaining devices, structures, etc. must be consulted in manufacturers' catalogs. The selection of belt for strength and resistance to abrasion, temperature, and the weather also is a topic for specialists.

BUCKET ELEVATORS AND CARRIERS

Bucket elevators and carriers are endless chains to which are attached buckets for transporting granular materials along vertical, inclined or horizontal paths. Figure 5.10 shows two basic types: spaced buckets that are far apart and continuous which overlap. Spaced buckets self-load by digging the material out of the boot and are operated at speeds of 200–300 fpm; they are discharged centrifugally. Continuous buckets operate at lower speeds, and are used for friable materials and those that would be difficult to pick up in the boot; they are fed directly from a loading chute and are discharged by gravity. Bucket carriers are essentially forms of pan conveyors; they may be used instead of belt conveyors for shorter distances and when they can **be** made of materials that are

TABLE \$	5.3.	Bulk Densities, Angles of F	Repose,	and	Allowable
		Angles of Inclination	•		

Material	Average Weight (lb/cuft)	Angle of Repose (degrees)	Recom- mended Maximum Inclination
Alum, fine	45-50	30-45	
Alumina	50-65	2 2	10-12
Aluminum sulfate	54	32	17
Ammonium chloride	45-52	02	••
Ammonium nitrate	45		
Ammonium sulfate	45-58		
Asbestos shred	20-25		
Ashes, coal, dry, fin. max	35-40	4 0	20-25
Ashes, coal, wet, $\frac{1}{2}$ in, max	45-50	50	23-27
Ashes, fly	40-45	4 2	20-25
Asphalt, $\frac{1}{2}$ in. max	45		
Baking powder	40-55		18
Barium carbonate	7 2		
Bauxite, ground	68	35	20
Bentonite, 100 mesh max	50-60		
Bicarbonate of soda	40-50		
Borax. 1/2 in.	55-60		
Borax, fine	45-55		20-22
Boric acid, fine	55		
Calcium acetate	125		
Carbon, activated, dry, fine	8-20		
Carbon black, pelleted	20-25		
Casein	36		
Cement, Portland	84	39	20-23
Cement, Portland, aerated	60-75		
Cement clinker	75-95	30-40	18-20
Charcoal	18-25	3 5	20-25
Chips, paper mill	20-25		
Clav. calcined	80-100		
Clay, dry, fine	100-120	35	20-22
Clav, drv, lumpy	60-75	3 5	18-20
Coal, anthracite, $\frac{1}{2}$ in, max	60	35	18
Coal, bituminous, 50 mesh max	50-54	4 5	24
Coal, bituminous, 🗄 in. max	43-50	4 0	2 2
Coal, lignite	40-45	38	22
Coke breeze, ¹ / ₄ in. max	25-35	30-45	20-22
Copper sulfate	75-85	31	17
Cottonseed, dry, delinted	35	29	16
Cottonseed, dry, not delinted	18-25	35	19
Cottonseed meal	35-40	3 5	2 2
Crvolite dust	75-90		
Diatomaceous earth	11-14		
Dicalcium phosphate	40-50		
Disodium phosphate	25-31		
Earth, as excavated, dry	70-80	35	20
Earth, wet, containing clay	100-110	4 5	23
Epsom salts	40-50		
Feldspar, $\frac{1}{2}$ in, screenings	70-85	38	18
Ferrous sulfate	60-75		
Flour, wheat	35-40		
Fullers earth. drv	30-35	23	
Fullers earth, oily	60-65		
Grain, distillery, spen, dry	30		
Graphite, flake	40		
Grass seed	10-12		
Gravel, bank run	90- 100	38	20
Gravel, dry, sharp	90-100		15-17
Gravel, pebbles	90-100	30	12
Gypsum dust, aerated	60-70	4 2	23
Gypsum, $\frac{1}{2}$ in, screenings	70-80	40	21
Iron oxide pigment	25	40	2 5
Kaolin talc. 100 mesh	42-56	4 5	23
Lactose	32		
Lead arsenate	72		
	· -		

TABLE 5.3-(continued)

Lead oxides	60-I 50		
Lime, 🛓 in. max	60-65	43	23
Lime, hydrated, 🛓 in. max	40	40	21
Lime, hydrated, pulverized	32-40	4 2	2 2
Limestone, crushed	85-90	38	18
Limestone dust	80-85		20
Lithopone	45-50		
Magnesium chloride	3 3		
Magnesium sulfate	70		
Milk, dry powder	36		
Phosphate, triple super, fertilizer	50-55	4 5	30
Phosphate rock, pulverized	60	40	25
Polystyrene beads	40		
Potassium nitrate	76		
Rubber, pelletized	50-55	35	22
Salt, common, coarse	40-55		
Salt, dry, fine	70-80	25	11
Salt cake, dry, coarse	85	36	21
Salt cake, drv, pulverized	60-85		
Saltpeter	80		
Sand, bank, damp	100-130	45	20-22
Sand, bank, dry	90-l 10	3 5	16-18
Sawdust	10-13	36	22
Shale, crushed	85-90	39	22
Soap chips	15-25	30	18
Soap powder	20-25		
Soda ash briquetts	50	22	7
Soda ash, heavy	55-65	32	19
Soda ash, light	20-35	37	22
Sodium bicarbonate	41	4 2	23
Sodium nitrate	70-80	2 4	11
Starch	25-50	24	12
Sugar, granulated	50-55		
Sugar, powdered	50-60		
Trisodium phosphate, pulverized	50	40	25
Wood chips	1 0-30	-	27
Zinc oxide, heavy	30-35		
Zinc oxide, light	10-15		
· •			

Other tables of these properties appear in these publications:

1. Conveyor Equipment Manufacturers Association, *Belt* Conveyors for *Bulk* Materials, 1966. 25-33.

 Stephens-Adamson Mfg. Co. Catalog 66, 1964, pp. 634-636.
 FMC Corporation Material Handling Equipment Division Catalog 100, 1983, pp. B.27-B.35.

4. Perry's Chemical Engineers Handbook, 1984, D. 7.5.

TABLE 5.4. Sizing Data for Screw Conveyors'

(a) Diameter (rpm and cuft/hr)

	A B Maxi		Capacities, Feet_Per	Cubic Hour	Maximum Recommonded	Capacities, Cubic Feet Per Hour			
Diam. of Con-	Max. Lump Size,	Speed R.P.M.	At Maximum Recommended ed Speed	At One R.P.M.	Speed R.P.M.	At Maximu Recommented ed Speed	m At d-One R.P.M.		
Inches	Inches	Loading of l	ading of Materials in Trough Class I—45% Full			Loading of Materials in Trough Class II-30% Full			
6 9 12 14 16 18 20	11/2 21/2 3 8 1/2	165 150 140 130 120 115 105	875 1200 2700 4000 5600 7600 9975	2.27 8.0 19.3 80.8 46.6 66.1 95.0	120 100 90 85 80 75 70	180 560 1200 1790 2510 3400 4340	1.5 5.6 13.8 21.1 81.4 45.4 62.1		

'Example 5.3 utilizes these data. (Stephens-Adamson Co. Catalog, 1954, p. 66).

TABLE 5.4(a)-(continued)

		Maximum	Capacitie Feet Pe	, Cubie Hour	Maximum	Capacities, Feet Per 1	Cubic Iour
Diam. of Con-	Max. Lump Size,	Recommended Speed R.P.M.	At Maximu Recommen ed Speed	m At d- One R.P.M.	R.P.M.	At Maximum Recommend- ed Speed	At One R.P.M.
Inches	Inches	Loading of I	Materials in II X80 %	Trough Fuil	Loading of Class	Materials in T 111—15% Full	rough
6 9 12 14 16 18 20	11/2 21/2 8 31/2	60 50 50 45 45 46 40 40	90 280 665 950 1410 1850 2485	1.5 5.6 13.3 21.1 81.4 45.4 62.1	60 50 50 45 45 40 40	45 140 335 470 705 910 1240	.75 2.8 6.7 10.5 15.7 22.7 81.1

(b) Characteristics of Some Materials (A Selection From the **Original Table)**

Materials	Approx. Weight per Cubie Foot	apacity Jassifi- cation	Type of Conveyor to Use	Horse- oower Factor "F"
Alfalfa meal	17	II	A,B,C	.4
Alum. lumpy	50-60	II	G,H,J	1.5
Alum. pulverized	45-50	II	A,B,C	.8
*Alumina.	60	III	K	2.0
Aluminum, hydrate Ammonium sulphate Aspestos, shredded,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	15-20 52 20-25 35-40	II II IIX	А,В,С С,Н,Ј D	.8 1.0 2.0
Asphalt, crushed	45		A,B,C	.5
Bakelite, nowdered	8040		A,B,C	1.4
Baking powder	41		A,B,C	.6
Barley.	38		A,B,C	.4
Bauxite, crushed Beans, castor Bans, navy, dry Bentonite.	75-85 36 48 51	III I IX	A,B,C A,B,C D	1.8 .5 .4 1.0
*Bones, crushed	85-40	IIX	مممم	2.0
*Bone s, granulated or ground	50	IIX		1.7
*Bone black.	20-25	IIX		1.7
Bonechar	40	IIX		1.7
*Bone meal.	55-60	UX	D	1.7
Borax, powdered.	53	II	A,B,C	.7
Boric add powder.	80-40	II	A,B,C	.8
Bran	16	II	A,B,C	.4

(c) Factor Sin the Formula for Power P

Diameter	Type of Hanger Bearing								
of Conveyor, Inches	SEALMASTER Ball Bearing	Babbitt. Bronze or Oil-Impreg- nated Wood	Self- Lubricating Bronze	Hard Iron					
4 6 9 10 12 14 16 18 20 24	12 18 32 38 55 78 106 140 165 230	21 33 54 66 06 02 135 186 240 285 390	33 54 96 114 171 255 336 414 510 690	50 80 130 160 250 350 480 600 700 950					

(d) Limits of Horsepower and Torque

Diameter	Diameter	Maximum	Maximum
of	of	Horsepower	Torque
Conveyor,	Coupling,	at	Capacity in
Inches	Inches	100 R.P.M.	Inch Pounds
4	1	1.5	950
6,9,10	1 ½	5.0	3200
9,10,12	2	10.0	6300
12,14	2 ¼	15.0	9500
12,14,16,18	3	25.0	16000
20	8 ¼	40.0	25000



Figure 5.7. A screw conveyor assembly and some of the many kinds of screws in use. (a) Screw conveyor assembly with feed hopper and discharge chute. (b) Standard shape with pitch equal to the diameter, the paddles retard the forward movement and promote mixing. (c) Short pitch suited to transfer of material up inclines of as much as 20°. (d) Cut flight screws combine a moderate mixing action with forward movement, used for light, fine, granular or flaky materials. (e) Ribbon flights are suited to sticky, gummy **Or** viscous substances.

EXAMPLE 5.3

Sizing a Screw Conveyor

Dense soda ash with bulk density 60 lb/cuft is to be conveyed a distance of 100 ft and elevated 12 ft. The material is class II-X with a factor F = 0.7. The bearings are self-lubricated bronze and the drive is V-belt with $\eta = 0.93$. The size, speed, and power will be selected for a rate of 15 tons/hr.

Q = 15(2000)/60 = 500 cuft/hr.

According to Table 5.4(a) this capacity can be accommodated by a 12 in. conveyor operating at

$$\omega = (500/665)(50) = 37.6$$
 rpm, say 40 rpm

From Table 5.4(c) the bearing factor is

s = 171.

Accordingly,

 $\dot{P} = [171(40) + 0.7(500)(60)]100 + 0.51(12)(30,000)/10^6$ = 2.97 HP motor HP = $G\dot{P}/\eta = 1.25(2.97)/0.93 = 3.99$, torque = 63,000(2.97)/40 = 4678 in. lb.

From Table **5.4(d)** the limits for a 12 in. conveyor are 10.0 HP and 6300 in. Ib so that the selection is adequate for the required service.

A conveyor 137 ft long would have a shaft power of 4.00 HP and a torque of 6300 in. **Ibs**, which is the limit with a 2 in. coupling; a sturdier construction would be needed at greater lengths.

For comparison, data of Table 5.5 show that a 14 in. troughed belt has an allowable speed of 267fpm at allowable inclination of 19" (from Table 5.3), and the capacity is

2.67(0.6)(38.4) = 61.5 tons/hr,

far more than that of the screw conveyor.







(c)









(e)

Figure 5.8. Flight conveyors in which the material is scraped along, and apron conveyors in which the material is carried along in a closed path of interconnected pans. (a) Flight conveyor, in which the material is scraped along a trough with flights attached to a continuous chain. (b) Scraper-type of flight. (c) Roller flights. (d) Apron conveyor, in which the material is carried along in moving, overlapping pans. (e) Shallow and deep types of overlapping pans.

TABLE 5.5. Belt Conveyor Data"

(a) Capacity (tons/hr) at 100 ft/min, 100 lb/cuft, and Indicated Slope Angle

	45° Troughed Belt									
Bett Width (inch)	0 ⁰	102	20 ⁰	30 ⁰						
14 16 18 20	27.99 38.70 51.00 65.10	33.00 45.60 6000 76.20	38.40 52.50 69.00 87.90	43.80 60.00 78.60 100.0						
24 30 36 42	98.10 160.8 238.5 331.8	114.9 187.5 277.8 385.8	132.0 214.8 318.0 441.3	149.7 243.6 360.3 499.5						
48 54 60 66 72	440.1 563.1 702.0 856.5 1026.0	511.5 654.6 815.4 994.2 1190 1	584.4 747.9 931.2 1134.9	661.2 845.7 1053.0 1282.8 1535.4						

	1	Flat Belt		
Belt width (inches)	5 ⁰	10 ⁰	20 ⁰	30 ⁰
14	2.85	6.69	14.01	21.42
16	3.87	9.18	19.05	29.16
18	5.07	11.88	24.90	38.10
20	6.399	15.06	31.50	48.24
24	9.57	22.47	47.10	72.06
30	113.51	36445	76 .32	116.8
36 1	22. 86	53.73	112.6	172.3
42	31.65	74.37	155.9	238.5
48	39.84	90.15	196.2	300.0
54	53.49	112577	263.4	403.2
60	66.60	156.5	327.9	5901.99
66	81.112	190.5	3999.66	6611.11
72	96.99	2380	477.9	731.1
		1		

'Example 5.4 utilizes these data. Power = $P_{\text{horizontal}} + P_{\text{vertical}} + P_{\text{empty}}$ (HP), where $P_{\text{horizontal}} = (0.4 + L/300)(W/100)$, $P_{\text{vertical}} = 0.001.4W$, and P_{empty} obtained from part (c), with H = lift (ft), L = horizontal travel (ft), and W = tons/hr.

(a) From Conveyor Equipment Manufacturers Association, 1979; (b) from Stephens-Adamson Catalog 66, 1954; (c) from Hudson, 19541.

(b) Maximum Recommended Belt Speeds for Nondusting Service

D-14		Belt Speed in Feet per Minute										
Belt Width. Inches	Lump Stone or ore	Gravel	Lump Coal	Crushed Ore and Stone †	Slack Coal †	Sand	Wood Chips Grain					
12	250	300	2 5 0	350	350	350	400					
18	300	350	300	400	400	400	500					
24	350	400	350	450	450	450	600					
30	400	450	400	500	500	600	700					
36	450	500	450	550	550	550	800					
42	500	550	500	550	600	600	800					
48	550	600	550	550	650	600	800					
54	550	600	600	550	700	600	800					
60	550	600	650	550	700	600	300					
66	550	600	700	550	700	600	800					
72	550	600	700	550	700	600	800					

TABLE 5.5-(continued)

(c) Power to Drive Empty Conveyor



particularly suited to a process. Capacity and power data for bucket machines are given in Table 5.6. Flight and apron conveyors are illustrated in Figure 5.11.

CONTINUOUS FLOW CONVEYOR ELEVATORS

One design of a drag-type of machine is the Redler shown on Figure 5.12. They function because the friction against the flight is greater than that against the wall. Clearly they are versatile in being able to transfer material in any direction and have the often important merit of being entirely covered. Circular cross sections are available but usually they are square, from 3 to 30 in. on a side, and operate at speeds of 30-250 **ft/min**, depending on the material handled and the construction. Some data are shown in Table 5.7. Most dry granular materials such as wood chips, sugar, salt, and soda ash are handled very well in this kind of conveyor. More difficult to handle are very fine materials such as cement or those that tend to pack such as hot grains or abrasive materials such as sand or crushed stone. Power requirement is dependent on the coefficient of sliding friction. Factors for power calculations of a few substances are shown in Table 5.7.

The closed-belt (zipper) conveyor of Figure 5.13 is a carrier that is not limited by fineness or packing properties or abrasiveness. Of course, it goes in any direction. It is made in a nominal 4-in. size, with a capacity rating by the manufacturer of 0.07 cuft/ft of travel. The power requirement compares favorably with that of open belt conveyors, so that it is appreciably less than that of other types. The formula is

$$HP = 0.001[(L_1/30 + 5)u + (L_2/16 + 2L_3)T],$$
(5.26)

where

u = ft/min, T = tons/hr, $L_1 = \text{total belt length (ft)},$ $L_2 = \text{length of loaded horizontal section (ft)},$

 L_3 = length of loaded vertical section (ft).

Speeds of 200 ft/min or more are attainable. Example 5.5 shows that the power requirement is much less than that of the Redler conveyor.



Figure 5.9. Some arrangements of belt conveyors (*Stephens-Adamson* Co.) and types of idlers (*FMC Corp.*). (a) Horizontal conveyor with discharge at an intermediate point as well as at the end. (b) Inclined conveyor, satisfactory up to 20° with some materials. (c) Inclined or retarding conveyor for lowering materials gently down slopes. (d) A flat belt idler, rubber cushion type. (e) **Troughed** belt idler for high loadings; usually available in 20°, 35°, and 45" side inclinations.





Figure **5.9**—(continued)

Closing Comments. Most kinds of conveyors and elevators are obtainable from several manufacturers, each of whom builds equipment to individual standards of sturdiness, materials of construction, mechanical details, performance, and price. These differences may be decisive in individual cases. Accordingly, a selection usually must be made from a manufacturer's catalog, and ultimately with the advice of the manufacturer.

5.4. SOLIDS FEEDERS

Several types are illustrated in Figures 5.9 and 3.7. Rates are controlled by adjusting gates or rotation speeds or translation

speeds. All of these methods require free flow from a storage bin which may be inhibited by bridging or arching. The device of Figure 5.9(a) provides motion to break up such tendencies.

For the most part the devices shown provide only rough feed rate control. More precise control is achieved by continuous weighing. The equipment of Figure 3.16(l) employs measurements of belt speed and the weight impressed on one or several of the belt idlers to compute and control the weight rate of feed; precision better than 0.5% is achievable. For some batch processes, the feeder discharges into an overhead weighing hopper for accurate measurement of the charge. Similar systems are used to batch feed liquids when integrating **flow** meters are not sufficiently accurate.

EXAMPLE 5.4

Sizing a Belt Conveyor

Soda ash of bulk density 60 lb/cuft is to be transported at 400 tons/hr a horizontal distance of 1200 ft up an incline of 5". The running angle of repose of this material is 19". The conveyor will be sized with the data of Table 5.5.

Consider a 24 in. belt. From Table 5.5(a) the required speed is

u = (400/132)100 = 303 ft/min.

Since the recommended maximum speed in Table 5.5(b) is 350 fpm, this size is acceptable:

conveyor length = $1200/\cos 5^{\circ}$ = 1205 ft, rise = 1200 tan 5° = 105 ft.

With the formulas and graph (c) of Table 5.5, the power requirement becomes

Power =
$$P_{\text{horizontal}} + P_{\text{vertical}} + P_{\text{empty}}$$

= (0.4 + 1200/300)(400/100)
+ 0.001(105)(400) + 303(3.1)/100
= 69.0 HP.

Perhaps 10 to 20% more should be added to compensate for losses in the drive gear and motor.





Figure 5.10. Closed belt (zipper) for conveying in any direction (*Stephens-Adamson* Co.). (a) Arrangement of pulley, feed hopper and open and closed belt regions. (b) The tubular belt conveyor for horizontal and vertical transport; a section of the zippered closed belt is shown. (c) Showing how the zipper closes (on downward movement of the belt in this sketch) or opens (on upward movement of the belt).

TABLE 5.6. Capacities and Power Requirements of Bucket Elevator Conveyors

Size of	Capac at 1	oo ft./1	ns/h r. nin	H	p.† with	materia	al at 50	lb./cu. 1	Ì.
bucket,	Buck	et spaci	ng.in.	Per li	0-ftve lift	rtical	Per. 100-	t. hori: run	zontal
$L \times W$		Ducact spacing, in			Spa	cing of	buckets,	in.	
	18	24	36	18	24	36	18	24	36
16×15 20×15 24×20 24×20 30×20 36×20	46 58 70 104 125 159 191	35 44 52 	23 29 35 52 63 79 95	0.59 .74 .90 1.30 1.60 2.00 2.42	0.44 .56 .67 	0.30 .37 .45 .66 .80 1.00 1.21	5.32 6.32 7.34 9.20 10.92 13.70 16.30	4.24 4.97 5.74	3.04 3.54 4.04 4.85 5.74 7.08 8 40

(a) Gravity Discharge Elevators Used Primarily For **Coal**^{a,c}

(b) Capacities and Maximum Size of Lumps of Centrifugal Discharge **Elevators**^{*b,c*}

Bucket Size, Spe		Speed.	Max. lumps		Capacity, tons/hr.			
spacing, in.	by width, in.	ft./min.	All lumps	10% lumps	35-lb. material	50-lb. material	100-lb. material	
13 16 16 18 18 19	6 × 4 8 × 5 10 × 6 12 × 7 14 × 7 16 × 8	225 230 230 268 268 268 268	×** 1 × × ×	21/2 3 31/2 4 4 41/2	5 9 16 27 32 44	7 13 23 38 46 63	14 27 47 77 92 127	

(c) Centrifugal Discharge of Continuous Belt and Bucket Elevators'

Bucket	Siza	a .	Max.	Max. lumps		Capacity, tons/hr.				
spacing, in.	length by width	Speed. ft./min.	All lumps	10% lumps	35 lb. material	50 lb. material	100 lb. material			
13 16	6 X 4 8 X 5	225 258 258	3/3	2½ 3 3¼	5 11 18	7 15 26	14 30 52			
18 18	12×7 14×7 $16y_8$	298 298 298		4 4 4 1 4	30 36 53	42 52 114	85 103 152			

"Buckets 80% full.

"Buckets 75% full.

• Horsepower = 0.002 (tons/hr)(lift in feet).

(Link Belt Co.)



Figure 5.11. Drag-type enclosed conveyor-elevator (Redler Design) for transfer in any direction (*Stephens-Adamson* Mfg. Co.). (a) Head and discharge end of elevator. (b) Carrying and return runs. (c) Loading end. (d) Some shapes of flights; some are made close-fitting and edged with rubber or plastics to serve as cleanouts.









Figure 5.12. Bucket elevators and conveyors. (a) Spaced bucket elevator. (b) Bucket conveyor for vertical and horizontal travel. (c) Discharge of pivoted buckets on horizontal path. (d) Spaced buckets receive part of their load directly and part by scooping the bottom. (e) Continuous buckets are filled as they pass through the loading leg with a feed spout above the tail wheel. (f) Centrifugal discharge of spaced buckets. (g) Discharge mode of continuous buckets.

TABLE 5.7. Speed and Horsepower of Drag-Type Conveyors of Redler Design'

	Veight per		3 "	unit	s	11	" U1	nit s	Ī	19	" Uni	ts
Material	Foot, ounds	ĸ	E	F	G	Е	F	G		E	F	G
Beans, dry navy Bicarbonate of soda, dry, pulver-	54 55	100	1.5	2.9	1.4	1.2	2.3	3.1		.1	2.02	.6
ized Bran	26	0.0	3.0 4	1 8.33.	.98.1 8	2.4 3.0	5.2 5.9	$5.4 \\ 2.8$	2	2.2	4 6 5.0	1.9 2.4
Cellulose acetate dry, coarse granular Cement. dry Portland Clay. dry lumpy Clay. pulverized	10 60-90 40-100 25-80	\$8	8.0 3 6017	15.9 76 1 5	.4 4≋0 4.6 .8	$5.3 \\ 2.3 \\ 2.4 \\ 4.3 $	10.0 5.4 4.5 11.9	$3.1 \\ 4.1 \\ 3.3 \\ 4.6$		1.4 2.1 2.1 8.8	8.5 4.8 4.0 9.9	2.6 3.4 2.8 3.7
Coal. minus 4 " slack dry with I'g proportion fines	40-50	4 0	2.4	4.6	4. 4	1.9	3.5	3.1		1.7	3.1	2.6
Coal minus 14" slack very wet	45-55 50-60	20	33 2.5	.15 5.45	.4 .7	2.6 2.0	4.7 4.1	3.8 4.0	2	?3 .8	42 3.6	8.1 8.2
damp Coal, sized wet or dry Coconut, shredded	40-50 40-50 25 28	40 80 2 0	2.2 3.0	1.1 6.1	3.8 3.2	2.0 1.7 2.2 1.8	3.7 3.1 4.3 3.5	3.1 2.8 2.4 2.4		1.8 1.5	332 28 0 3	712.1
Corn flakes Flour, wheat	12 30-40	20	2.4 3.8	4.83 7.	2.3	2.6	5.2 5.0	$\frac{1.9}{2.6}$	14.4	2.1	4 4 3	11.7 2.3
Fuller's earth, dry granular Lime, burned or "quick "lump	4 2	80	32 3 .	5 0	7.3 7.0	2.3	5.0	4.9	ĺ	2.1	4.4	4.0 513.8
Lime, dry burned small lumps and dust	50	120	3.5	6.9	6.5 7 5	2.8	5.5	4.4		2.5	4.9	3.6
Lime, fine with tendency to pack Lime, hydrated Salt, dry granulated	10-25 80 75	ू0़ 100		35.5_8	7 0 5 7 6 5	5.± 7.3 1.6	22.5 3.1 2.9	4.8 4.0	1	5.0 1.5	18.1 2.8 2.6	3.8 3.2 3.6
Sand, silica coarse dry Sand very fine. dry	90-100 90-100	160 120	2.1 202	4 . 2 14.64	7.0 17.3	1.8	3.4	4.8		1.7	3.2	B.8 84.0
Sawdust, dry Soda ash, light soybean meal Starch. lump	25-35 40 2 5 %	20 20 80	4.5 2_2 1 5.4	11.4 15.86	6.5 4.6 3.8	3.3 1.7 1.6	7.8 3.5 2.9	$\frac{3.0}{4.4}$ $\frac{3.3}{2.8}$		2.8	6.6 3.1 2.1	3.6 . 8 2.4
Starch, pulverized Sugar, dirv granulated	50 40-50	160	4.4 2	8.57 5	.0 89.1	3.9 2.2	10.7 4.4 6	$\frac{4.1}{6.1}$		2.0	4 8.9 3.9 5.6	913.4 4.8 4.1
Wheat. dry fairly clean Wood chips. dry	48 15-30	44	351	7 3.35 6	.2 62.8	$1.3 \\ 2.4$	2.5 4.5	$3.6 \\ 2.2$		1.2	2.2	3.0 1.9

(a) Typical Speeds (ft/min)^b

MATERIAL HANDLED	1000 Conv.	1000 Elev.	2000 Conv.	3000 Conv.
Coal Coke	125 40	125 40	80 40	150 40
Flyash	3 0	3 0	3 0	3 0
Grain (Whole)	125	125	6 0	250
(Processed)	125	100	8 0	150
Salt	125	100	8 0	150
Wood (Chips)	100	80	80	150
(Sawdust	100	100	8 0	150

[#] HP = 0.001 (FL + GH + K)(tons/hr), where H = rise (ft), L = horizontal run (ft), F, G, and Kare factors from Table (b); factor E is not used in this formula. ^b Series 1000, 2000, and 3000 differ in the shapes and sturdiness of

the flights.

(Stephens-Adamson Mfg. Co.).

(b) Factors F, G, and Kfor Use in the Power Equation for Three Sizes of Units



Figure 5.13. Types of feeders for granular solids; also suitable are conveyors such as closed belt, Redler, and bucket types. (a) Bin discharge feeder. (b) Rotary plate feeder with adjustable collar and speed. (c) Flow controlled by an adjustable gate. (d) Rotary drum feeder, regulated by gate and speed. (e) Rotary vane feeder, can be equipped with air lock for fine powders. (f) Vane or pocket feeder. (g) Screw feeder. (h) Apron conveyor feeder. (i) Belt conveyor feeder. (j) Undercut gate feeder. (k) Reciprocating plate feeder. (1) Vibrating feeder, can transfer uphill, downhill, or on the level. (m) "Air-slide" feeder for powders that can be aerated. (n) Weighing belt feeder; unbalance of the weigh beam causes the material flow rate onto the belt to change in the direction of restoring balance.



EXAMPLE 5.5

Comparison of Redler and Zippered Belt Conveyors Soda ash of bulk density 30 **lb/cuft** is to be moved 120ft horizontally and 30 ft vertically at the rate of 350 **cuft/hr**. Compare power requirements of Redler and zippered belt conveyors for this service.

A 3-in Redler is adequate:

$$u = \frac{350}{60(\pi/4)(3/12)^2} = 118.8 \text{ fpm},$$

which is within the range of Table 5.7(a),

tons/hr = 350(30)/2000 = 5.25

Take constants from Table 5.7(b) for a Redler.

$$HP = \frac{5.25}{1000} [11.4(120) + 6.5(30) + 20] = 8.31$$

For a closed belt,

$$u = \frac{350}{0.07(60)} = 83.3$$
 fpm,

which is well under the 200 fpm that could be used,

$$L_1 = 300, \quad L_2 = 120, \quad L_3 = 30.$$

Use Eq. (5.26):

 $HP = 0.001\{(300/30 + 5)83.3 + [120/16 + 2(30)]5.25\} = 1.60.$

REFERENCES

- T.H. Allegri, Materials Handling Principles and Practice, Van Nostrand Reinhold, New York, 1984.
- A.G. Bain and S.T. Bonnington, The Hydraulic Transport Of Solids by Pipeline, Pergamon, New York, 1970.
- M.V. Bhatic and P.N. Cheremisinoff, Solid and Liquid Conveying Systems, Technomic, Lancaster, PA, 1982.
- A.J. Bobkowicz and W.G. Gauvin, The effects of turbulence in the flow characteristics of model fibre suspensions, *Chem. Eng. Sci. 22, 229-247* (1967).
- 5. R. Clift, Conveyors, hydraulic, Encycl. Chem. Process. Des. 11, 262-278 (1980).
- H. Colijn, Mechanical Conveyors for Bulk Solids, Elsevier, New York, 1985.
- Conveyor Equipment Manufacturers Association, Belt Conveyors for Bulk Materials, Van Nostrand Reinhold, New York, 1979.

- 8. D.W. Dodge and A.B. Metzner, Turbulent flow of non-newtonian systems, *AIChE J. 5*, 189 (1959).
- G.H. Ewing, Pipeline transmission, in Marks' Mechanical Engineers Handbook, McGraw-Hill, New York, 1978, pp. 11.134-11.135.
- FMC Corp. Material Handling Equipment Division, Catalog 100, Homer City, PA, 1983.
- F.J. Gerchow, Conveyors, pneumatic, in *Encycl. Chem. Process. Des.* 11, 278–319 (1980); *Chem. Eng.*, (17 Feb. 1975, 31 Mar. 1975).
- 12. H.V. Hawkins, Pneumatic conveyors, in Marks' Mechanical Engineers Handbook, McGraw-Hill, New York, 1978, pp. 10.50-10.63.
- J.W. Hayden and T.E. Stelson, Hydraulic conveyance of solids in pipes, in Zandi, Ref. 27, 1971, pp. 149-163.
- 14. W.G. Hudson, Conveyors and Related Equipment, Wiley, New York, 1954.
- E. Jacques and J.G. Montfort, Coal transportation by slurry pipeline, in Considine (Ed.), *Energy Technology Handbook*, McGraw-Hill, New York, 1977, pp. 1.178-1.187.

REFERENCES 89

- M. Kraus, Pneumatic Conveying Of Bulk Materials, McGraw-Hill, New York, 1980.
- R.A. Kulwiec (Ed.), Material Handling Handbook, Wiley, New York, 1985.
- D.E. Perkins, and J.E. Wood, Design and Select Pneumatic Conveying Systems, *Hydrocarbon Processing* 75-78 (March 1974).
- G.J. Raymus, Pneumatic conveyors, in *Perry's Chemical Engineers Handbook*, McGraw-Hill, New York, 1984, pp. 7.17–7.25.
- P.E. Solt, Conveying, pneumatic troubleshooting, *Encycl. Chem.* Process. Des. 11, 214-226 (1980).
- 21. Stephens-Adamson Mfg. Co., General Catalog 66, Aurora, IL, 1954, and updated sections.

- H.A. Stoess, *Pneumatic Conveying*, Wiley, New York, 1983.
 E.J. Wasp, T.C. Aude, R.H. Seiter, and T.L. Thompson, in Zandi, Ref. 27, 1971, pp. 199–210.
- E.J. Wasp, J.P. Kenny, and R.L. Gandhi, *Solid-Liquid Flow in Slurry Pipeline Transportation*, Trans. Tech. Publ., 1917, Gulf, Houston, 1979.
- 25. E.J. Wasp, T.L. Thompson, and P.E. Snoek, The era of slurry pipelines, *Chem. Technol.*, 552-562 (Sep. 1971).
- O.A. Williams, Pneumatic and Hydraulic Conveying of Solids, Dekker, New York, 1983.
- I. Zandi (Ed.), Advances in Solid-Liquid Flow in Pipes and Its Applications, Pergamon, New York, 1971.
The transfer of fluids through piping and equipment is accompanied by friction and may result in changes in pressure, velocity, and elevation. These effects require input of energy to maintain flow at desired rates. In this chapter, the concepts and theory of fluid mechanics bearing on these topics will be reviewed briefly and practical and empirical methods of sizing lines and auxiliary equipment will be emphasized.

6.1. PROPERTIES AND UNITS

The basis of flow relations is Newton's relation between force, mass, and acceleration, which is

 $F = (m/g_c)a. \tag{6.1}$

When F and m are in lb units, the numerical value of the coefficient is $g_c = 32.174$ lb ft/lbf sec². In some other units,

$$g_c = 1 \frac{\text{kg m/sec}^2}{\text{N}} = 1 \quad \frac{\text{g cm/sec}^2}{\text{dyn}} = 9.806 \frac{\text{kg m/sec}^2}{\text{kg_f}}$$

Since the common engineering units for both mass and force are 1 lb, it is essential to retain g_c in all force-mass relations. The interconversions may be illustrated with the example of viscosity whose basic definition is force/(velocity)(distance). Accordingly the viscosity in various units relative to that in SI units is

$$1 \text{ Ns/m}^2 = \frac{1}{9.806} \text{ kg}_f \quad \text{s/m}^2 = 10 \text{ g/(cm)(s)}$$

= 10 P = 0.0672 lb/(ft)(sec)
= $\frac{0.0672}{32.174}$ lbf sec/ft² = 0.002089 lbf sec/ft².

In data books, viscosity may be recorded either in force or mass units. The particular merit of **SI** units (kg, m, s, N) is that $g_c \approx 1$ and much confusion can be avoided by consistent use of that system. Some numbers of frequent use in fluid flow problems are

Data of densities of liquids are empirical in nature, but the effects of temperature, pressure, and composition can be estimated; suitable methods are described by Reid et al. (Properties of *Gases and Liquids*, McGraw Hill, New York, 1977), the *API Refining Data* Book (American Petroleum Institute, Washington, DC, 1983), and the *AIChE Data Prediction Manual* (1984–date). The densities of gases are represented by equations of state of which the simplest is that of ideal gases; from this the density is given by:

$$p = 1/V = MP/RT$$
, mass/volume (6.2)

where M is the molecular weight. For air, for example, with P in atm and $Tin \, {}^\circ R$,

$$\rho = \frac{29P}{0.73T}, \quad \text{lb/cuft.}$$
(6.3)

For nonideal gases a general relation is

FLOW OF FLUIDS

$$\rho = MP/zRT, \tag{6.4}$$

where the compressibility factor z is correlated empirically in terms of reduced properties T/T, and P/P_c and the acentric factor. This subject is treated for example by Reid et al. (1977, p. 26) and Walas (1985, pp. 17, 70). Many **PVT** equations of state are available. That of Redlich and Kwong may be written in the form

$$V = b + RT/(P + a/\sqrt{TV^2}), \qquad (6.5)$$

which is suitable for solution by direct iteration as used in Example 6.1.

Flow rates are expressible as linear velocities or in volumetric, mass, or weight units. Symbols for and relations between the several modes are summarized in Table 6.1.

The several variables on which fluid flow depends may be gathered into a smaller number of dimensionless groups, of which the Reynolds number and friction factor are of particular importance. They are defined and written in the common kinds of units also in Table 6.1. Other dimensionless groups occur less frequently and will be mentioned as they occur in this chapter; a long list is given in *Perry's Chemical Engineers Handbook* (McGraw-Hill, New York, 1984, p. 5.62).

EXAMPLE 6.1

Density of a **Nonideal** Gas from Its Equation of State The Redlich-Kwong equation of carbon dioxide is

$$(P+63.72(10^6)/\sqrt{T}V^2)(V-29.664) = 82.05T$$

with *P* in atm, *V* in mL/g mol and *Tin* K. The density will be found at P = 20 and T = 400. Rearrange the equation to

 $V = 29.664 + (82.05)(400)/(20 + 63.72(10^6)/\sqrt{400} V^2).$

Substitute the ideal gas volume on the right, V = 1641; then find V on the left; substitute that value on the right, and continue. The successive values of V are

 $V = 1641, 1579, 1572.1, 1571.3, 1571.2, \dots mL/g mol$

and converge at 1571.2. Therefore, the density is

 $\rho = 1/V = 111571.2$, or 0.6365 g mol/L or 28.00 g/L.

92 FLOW OF FLUIDS

TABLE 6.1. Flow Quantities, Reynolds Number, and Friction Factor

Flow	Symbol and	Typical Units		
Quantity	Equivalent	Common	SI	
Linear Volumetric Mass Weight Mass/area Weight/area	$u = uA = \pi D^{2} u/4$ $\dot{m} = \rho Q = \rho A u$ $\dot{w} = \gamma Q = \gamma A u$ $G = \rho u$ $G_{\gamma} = \gamma u$	ft/sec cuft/sec lb/sec lbf/sec lb/(sqft)(sec) lbf/(sqft)(sec)	m/sec m³/sec kg/sec N/sec kg/m* sec N/m² sec	

Reynolds Number (with A= $\pi D^2/4$)

$$\operatorname{Re} = \frac{Du\rho}{\mu} \approx \frac{Do}{v} = \frac{DG}{\mu} = \frac{4Q\rho}{\pi D\mu} = \frac{4\dot{m}}{\pi D\mu}$$
(1)

Friction Factor

$$f = \frac{\Delta P}{\rho} / \left(\frac{L}{D} \frac{u^2}{2g_c}\right) = 2g_c D \Delta P / L\rho u^2 = 1.6364 / \left[\ln\left(\frac{0.135\varepsilon}{D} + \frac{6.5}{Re}\right)\right]^2$$
(2)
(Round's equation)
$$\frac{AP}{\rho} = \frac{L}{D} \frac{u^2}{2g_c} f = \frac{8LQ^2}{g_c \pi^2 D^5} f = \frac{8L\dot{m}^2}{g_c \pi^2 \rho^2 D^5} f = \frac{LG^2}{2g_c D\rho^2} f$$
(3)

In the units

$$D = \text{ in., } \dot{m} = \text{lb/hr}$$

$$Q = \text{cuft/sec, } \mu = \text{cP}$$

$$p = \text{specific gravity}$$

$$Re = \frac{6.314\dot{m}}{D\mu} = \frac{1.418(10^6)\rho Q}{D\mu}$$

$$A P = 3.663(10^{-9})\dot{m}^2 + \mu \kappa$$
(4)

$$\frac{1}{L} = \frac{0.385(10^{-8})\dot{m}^2}{\rho D^5} f_{, atm/ft}$$
(5)

$$=\frac{0.6979\rho Q^2}{r^5} , f \text{ psi/ft}$$
(7)

Laminar Flow

$$Re < 2300$$

f = 64/Re (2a)
 $\Delta P/L = 32\mu u/D^2$

$$P/L = 32\mu u/D^{-7} = \frac{1.841(10^{-7})\mu\dot{m}}{20^{4}}, \text{ atm/ft}$$
(5a)

$$=\frac{2.707(10^{-6})\mu\dot{m}}{\rho D^{4}}, \text{ psi/ft}$$
(6a)
35.083 μQ

$$=\frac{-\frac{1}{D^4}}{D^4}, \text{ psi/ft}$$
(7a)

Gravitation Constant

 $g_c = 1 \text{ kg m/N sec}^2$

- = 1 g cm/dyn sec²
- = 9.806 kg m/kgf sec²

= 32.174 lbm ft/lbf sec²

- = 1 slug ft/lbf sec²
- = 1 lbm ft/poundal sec²

6.2. ENERGY BALANCE OF A FLOWING FLUID

The energy terms associated with the flow of a fluid are

- 1. Elevation potential $(g/g_c)z$,
- 2. Kinetic energy, $u^2/2g_c$,
- 3. Internal energy, U,
- 4. Work done in crossing the boundary, PV,
- 5. Work transfer across the boundary, W_s ,
- 6. Heat transfer across the boundary, Q.

Figure 6.1 represents the two limiting kinds of regions over which energy balances are of interest: one with uniform conditions throughout (completely mixed), or one in plug flow in which gradients are present. With single inlet and outlet streams of a uniform region, the change in internal energy within the boundary is

$$d(mU) = m \ dU + U \ dm = m \ dU + U(dm_1 - dm_2)$$

= $dQ - dW_s + [H_1 + u_1^2/2g_c + (g/g_c)z_1] \ dm_1$
- $[H_2 + u_2^2/2g_c + (g/g_c)z_2] \ dm_2.$ (6.6)

One kind of application of this equation is to the filling and emptying of vessels, of which Example 6.2 is an instance.

Under steady state conditions, d(mU) = 0 and $dm_{0} = dm_{2} = dm_{0}$, so that Eq. (6.6) becomes

$$\mathbf{AH} + \Delta u^2 / 2g_c + (g/g_c) \Delta z = (Q - W_s)/m, \tag{6.7}$$

or

$$AU + A(PV) + \Delta u^2/2g_c + (g/g_c)\Delta z = (Q \quad W_s)/m, \qquad (6.8)$$

or

$$\mathbf{AU} + \Delta(P/\rho) + \Delta u^2/2g_c + (g/g_c)\Delta z = (Q - W_s)/m. \tag{6.9}$$

For the plug Row condition of Figure 6.1(b), the balance is made in terms of the differential changes across a differential length dL of the vessel, which is

$$dH + (1/g_c)u \, du + (g/g_c) \, dz = dQ - dW_s, \tag{6.10}$$

where all terms are per unit mass.



Figure 6.1. Energy balances on fluids in completely mixed and plug flow vessels. (a) Energy balance on a bounded space with uniform conditions throughout, with differential flow quantities dm, and dm. (b) Differential energy balance on a fluid in plug flow in a tube of unit cross section.

EXAMPLE 6.2 Unsteady Flow of an Ideal Gas through a Vessel

An ideal gas at 350 K is pumped into a 1000 L vessel at the rate of 6 g mol/min and leaves it at the rate of 4 g mol/min. Initially the vessel is at 310K and 1 atm. Changes in velocity and elevation are negligible. The contents of the vessel are uniform. There is no work transfer.

Thermodynamic data:

$$U = C_v T = 5T,$$

$$H = C_p T = 7T.$$

Heat transfer:

$$dQ = h(300 - T) d\theta$$
$$= 15(300 - T) d\theta.$$

The temperature will be found as a function of time θ with both h = 15 and h = 0.

$$dn_1 = 6 d\theta,$$

$$dn_2 = 4 d\theta,$$

$$dn = dn_1 - dn_2 = 2 d\theta,$$

$$n_0 = P_0 V/RT_0 = 1000/(0.08205)(310) = 39.32 \text{ gmol}$$

$$n = n_0 + 2\theta,$$



Energy balance

$$d(nU) = n dU + U dn = nC_{u} dT + C_{u} T(2 d\theta)$$

 $= H_1 dn_1 - H_2 dn_2 + dQ - dw_s$ = $C_p (6T_1 \quad 4T) d\theta + h(300 - T) d\theta$,

This rearranges into

$$\begin{split} \int_{0}^{\theta} \frac{d\theta}{n_{0}+2\theta} &= \int_{310}^{T_{2}} \frac{dT}{(1/C_{v})[6C_{p}T_{1}+300h-(4C_{p}+2C_{v}+h)T]} \\ &= \begin{cases} \int_{310}^{T_{2}} \frac{dT}{3840-10.6T}, & h=15, \\ \int_{310}^{T_{2}} \frac{dT}{2940-7.6T}, & h=O. \end{cases} \end{split}$$

The integrals are rearranged to find T,

$$T_2 = \begin{cases} 362.26 - 52.26 \left(\frac{1}{1+0.0509\theta}\right)^{5.3}, & h = 15, \\ 386.84 - 76.84 \left(\frac{1}{1+0.0509\theta}\right)^{3.8}, & h - 0. \end{cases}$$

Some numerical values are

Τ2			I	Þ
θ	h = 15	h = 0	h = 15	h = 0
0	310	310	1	1
0.2	312.7	312.9	1.02	1.02
0.5	316.5	317.0		
1	322.1	323. 2		
5	346.5	354.4		
10	356.4	370.8	1.73	1.80
00	362.26	386.84	00	œ

The pressures are calculated from

$$P = \frac{nRT}{V} = \frac{(39.32 + 2\theta)(0.08205)T}{1000}$$

Friction is introduced into the energy balance by noting that it is a mechanical process, dW_f , whose effect is the same as that of an equivalent amount of heat transfer dQ_f . Moreover, the total effective heat transfer results in a change in entropy of the flowing liquid given by

$$T\,dS = dQ + dW_f.\tag{6.11}$$

When the thermodynamic equivalent

$$dH = V \, dP + T \, dS \tag{6.12}$$

and Eq. (6.11) are substituted into Eq. (6.10), the net result is

$$V dP + (1/g_c)u du + (g/g_s) dz = -(dW_s + dW_f),$$
 (6.13)

which *is* known as the mechanical energy balance. With the expression for friction of Eq. (6.18) cited in the next section, the mechanical energy balance becomes

$$V dP + (1/g_c)u du + (g/g_s) dz + \frac{fu^2}{2g_{sc}D} dL = -dW_s.$$
 (6.13)

For an incompressible fluid, integration may be performed term by term with the result

$$\Delta P/\rho + \Delta u^2/2g_c + (g/g_c)\Delta z = -(W_{s+}W_f).$$
(6.14)

The apparent number of variables in Eq. (6.13) is reduced by the substitution u = V/A for unit flow rate of mass, where A is the cross-sectional area, so that

$$V dP + (1/g_c A^2) V dV + (g/g_r) dz = -(dW_s + dW_f).$$
(6.15)

Integration of these energy balances for compressible fluids under several conditions is covered in Section 6.7.

The frictional work loss W_f depends on the geometry of the system and the flow conditions and is an empirical function that will be explained later. When *it* is known, Eq. (6.13) may be used to find a net work effect W_e for otherwise specified conditions.

The first three terms on the left of Eq. (6.14) may be grouped into a single stored energy terms as

$$AE = \Delta P/\rho + \Delta u^2/2g_c + (g/g_c)\Delta z, \qquad (6.16)$$

EXAMPLE 6.3 Units of the Energy Balance

In a certain process the changes in stored energy and the friction are

AE = -135 ft lbf/lb $w_f = 13 \text{ ft } lbf/lb.$

The net work will be found in several kinds of units:

$$w_s = -(AE + w_f) = 122 \text{ ft lbf/lb},$$

 $w_s = 122 \frac{\text{ft lbf}}{\text{lb}} \frac{4.448N2.204 \text{ lb}}{\text{lbf}} \frac{m}{\text{kg g g}} \frac{3.3.28 \text{ ft}}{3.3.28 \text{ ft}}$

$$= 364.62, \quad 364.6\frac{J}{kg},$$

w_s = 364.6 $\frac{N m}{kg} \frac{kgf}{9.806 N} = 37.19 \frac{m kgf}{kg}$

At sea level, numerically lbf = lb and kgf = kg. Accordingly,

$$w_{\rm s} = 122 \frac{\text{ft lbf lb} \text{kgf}}{\text{lb lbf kg}} \frac{m}{3.28 \text{ ft}} = 37.19 \frac{\text{kgf m}}{\text{kg}}$$

as before.

and the simpler form of the energy balance becomes

 $AE + W_f = -W_s. \tag{6.17}$

The units of every term in these energy balances are alternately:

ft lb_f/lb with $g_c = 32.174$ and g in ft/sec^2 (32.174 at sea level). N m/kg = J/kg with $g_c = 1$ and g in m/sec² (1.000 at sea level). kg, m/kg with $g_c = 9.806$ and g in m/sec² (9.806 at sea level).

Example 6.3 is an exercise in conversion of units of the energy balances.

The sign convention is that *work input is a negative quantity* and consequently results in an increase of the terms on the left of Eq. (6.17). Similarly, work is produced by the flowing fluid only if the stored energy AE is reduced.

6.3. LIQUIDS

Velocities in pipe lines are limited in practice because of

1. the occurrence of erosion.

economic balance between cost of piping and equipment and the cost of power loss because of friction which increases sharply with velocity.

Although erosion is not serious in some cases at velocities as high as 10-15 ft/sec, conservative practice in the absence of specific knowledge limits velocities to 5-6 ft/sec.

Economic optimum design of piping will be touched on later, but the rules of Table 6.2 of typical linear velocities and pressure drops provide a rough guide for many situations.

The correlations of friction in lines that will be presented are for new and clean pipes. Usually a factor of safety of 20–40% is advisable because pitting or deposits may develop over the years. There are no recommended fouling factors for friction as there are for heat transfer, but instances are known of pressure drops to double in water lines over a period of 10 years or so.

In lines of circular cross section, the pressure drop is represented by

$$\Delta P = f\rho \frac{L}{D} \frac{u^2}{2g_c}.$$
(6.18)

For other shapes and annular spaces, D is replaced by the hydraulic

diameter

 $D_h = 4(\text{cross section})/\text{wetted perimeter}.$

For an annular space, $D_h = D_2 - D_1$.

In laminar flow the friction is given by the theoretical Poiseuille equation

$$f = 64/N_{\rm re}, N_{\rm Re} < 2100,$$
 approximately. (6.19)

At higher Reynolds numbers, the friction factor is affected by the roughness of the surface, measured as the ratio e/D of projections on the surface to the diameter of the pipe. Values of ε are as follows; glass and plastic pipe essentially have $\varepsilon = 0$.

	ε (ft)	ε (mm)
Riveted steel	0.003-0.03	0.9-9.0
Concrete	0.001-0.01	0.3-3.0
Wood stave	0.0006-0.003	0.18-0.9
Cast iron	0.00085	0.25
Galvanized iron	0.0005	0.15
Asphalted cast iron	0.0004	0.12
Commercial steel or		
wrought iron	0.00015	0.046
Drawn tubing	0.000005	0.0015

The equation of Colebrook [J. Inst. Civ. Eng. London, 11, pp. 133-156 (1938–1939)] is based on experimental data of Nikuradze [Veer. Dtsch. Ing. Forschungsh. 356 (1932)].

$$\frac{1}{\sqrt{f}} = 1.14 - 0.869 \ln\left(\frac{\varepsilon}{D} + \frac{9.38}{N_{\rm Re}\sqrt{f}}\right), \quad N_{\rm Re} > 2100. \tag{6.20}$$

Other equations equivalent to this one but explicit in f have been devised. A literature review and comparison with more recent experimental data are made by Olujic [*Chem. Eng., 91-94,* (14 Dec. 1981)]. Two of the simpler but adequate equations are

$$f = 1.6364 \left[\ln \left(\frac{0.135\varepsilon}{D} + \frac{6.5}{N_{\text{Re}}} \right) \right]^{-2}$$
(6.21)

[Round, Can. J. Chem. Eng. 58, 122 (1980)],

$$f = \left\{-0.\ 86861 \ln \left[\frac{\varepsilon}{3.7D} - 2.1802 \ln \left(\frac{\varepsilon}{3.7D} + \frac{14.5}{N_{\text{Re}}}\right)\right]\right\}^{-2}$$
(6.22)

[Schacham, Ind. Eng. Chem. Fundam. 19(5), 228 (1980)]. These

TABLE 6.2. Typical Velocities and Pressure Drops in Pipelines

	Liquids (psi/100 ft)			
	Liquids within 50°F of Bubble Point	Light Oils and Water	Viscou Oils	
Pump suction	0.15	0.25	0.25	
Pump discharge	2.0 (or 5-7 fps)	2.0 (or 5-7 fps)	2.0 (or 3-4fps)	
Gravity flow to or from tankage, maximum Thermosynhon rehoiler	0.05	0.05	0.05	
inlet and outlet	0.2			

Pressure (psig)	o-300ft Equivalent Length	300–600 ft Equivalent Length	
–13.7 (28in. Vac)	0.06	0.03	
– 12.2 (25 in. Vac)	0.10	0.05	
—7.5 (15 in. Vac)	0.15	0.08	
0	0.25	0.13	
50	0.35	0.18	
100	0.50	0.25	
150	0.60	0.30	
200	0.70	0.35	
500	2.00	1.00	
Steam	psi/100 ft	Maximumft/min	
Under 50 psig	0.4	10,000	
Over 50 psig	1.0	7000	

Steam Condensate

To *traps*, 0.2 psi/l 00 ft. From *bucket traps*, size on the basis of 2-3 times normal flow, according to pressure drop available. *From* continuous drainers, size on basis of design flow for 2.0 psi/100 ft

Control Valves

Require a pressure drop of at least 10 psi for good control, but values as low as 5 psi may be used with some loss in control quality

Particular Equipment Lines	(ft/sec)
Reboiler, downcomer (liquid)	3-7
Reboiler, riser (liquid and vapor)	35-45
Overhead condenser	25-100
Two-phase flow	35-75
Compressor, suction	75-200
Compressor, discharge	100–250
Inlet, steam turbine	120-320
Inlet, gas turbine	150-350
Relief valve, discharge	0.5 <i>v</i> _
Relief valve, entry point at silencer	v,

^a V_c is sonic velocity.

three equations agree with each other within 1% or so. The Colebrook equation predicts values 1-3% higher than some more recent measurements of Murin (1948), cited by Olujic (Chemical *Engineering*, 91-93, Dec. 14, 1981).

For orientation purposes, the pressure drop in steel pipes may be found by the rapid method of Table 6.3, which is applicable to highly turbulent flow for which the friction factor is given by von Karman's equation

$$f = 1.3251[\ln(D/\varepsilon) + 1.3123)]^{-2}.$$
(6.23)

Under some conditions it is necessary to employ Eq. (6.18) in differential form. In terms of mass flow rate,

$$dP = \frac{8\dot{m}^2 f}{g_c \pi^2 \rho D^5} dL. \tag{6.24}$$

Example 6.4 is of a case in which the density and viscosity vary along the length of the line, and consequently the Reynolds number and the friction factor also vary.

FITTINGS AND VALVES

Friction due to fittings, valves and other disturbances of flow in pipe lines is accounted for by the concepts of either their equivalent lengths of pipe or multiples of the velocity head. Accordingly, the pressure drop equation assumes either of the forms

$$AP = f(L + \sum L_i)\rho u^2/2g_c D, \qquad (6.25)$$

$$\Delta P = [f(L/D) + \sum K_i]\rho u^2/2g_c. \qquad (6.26)$$

Values of equivalent lengths L_i and coefficients K_i are given in Tables 6.4 and 6.5. Another well-documented table of K_i is in the *Chemical Engineering Handbook* (McGraw-Hill, New York, 1984 p. 5.38).

Comparing the two kinds of parameters,

$$K_i = fL_i/D \tag{6.27}$$

so that one or the other or both of these factors depend on the friction factor and consequently on the Reynolds number and possibly ε . Such a dependence was developed by Hooper [*Chem. Eng.*, 96-100, (24 Aug. 1981)] in the equation

$$K = K_1 / N_{\rm Re} + K_2 (1 + 1/D), \tag{6.28}$$

where D is in inches and values of K_1 and K_2 are in Table 6.6. Hooper states that the results are applicable to both laminar and turbulent regions and for a wide range of pipe diameters. Example 6.5 compares the several systems of pipe fittings resistances. The K_i method usually is regarded as more accurate.

ORIFICES

In pipe lines, orifices are used primarily for measuring flow rates but sometimes as mixing devices. The volumetric flow rate through a thin plate orifice is

$$Q = C_d A_0 \left(\frac{2\Delta P/\rho}{1-\beta^4}\right)^{1/2},$$
 (6.29)

 A_{r} = cross sectional area of the orifice,

 $\beta = d/D$, ratio of the diameters of orifice and pipe.

For corner taps the coefficient is given by

$$C_d \approx 0.5959 + 0.0312\beta^{2.1} - 0.184\beta^8 + (0.0029\beta^{2.5})(10^6/\text{Re}_D)^{0.75}$$
(6.30)

(International Organization for Standards Report DIS 5167, Geneva, 1976). Similar equations are given for other kinds of orifice taps and for nozzles and Venturi meters.

	w C ₁	Nominal Pipe Size In.	Schedule Number	Value of C2	Nomina Pipe Size In.	l Schedule Number	Value of C ₂
		1/8	40 s 80 x	7.920.000 ;26.200.000	12	20 30	0.0157 0.0168 0.0175
ŧ	1500	1/2	40 S 80 X 160	93,500 186.100 4.300.000		.40 ≭ 60	0.0180 0.0195 0.0206
$W = C_1 \qquad C_1 \qquad W$		3/4	40 s 80 x 160	21,200 36,900 100,100 627,000		80 100 120 140 160	0.0231 0.0267 0.0310 0.0350 0.0423
9 .08 .07 .15 8 .06	900	1	40 5 80 X 160 xx	5,950 9,640 22,500 114.100	14	10 20 30 s 40	0.00949 0.00995 0.01046 0.01099 0.01155 0.01244
705 .2	700 - 500	1 1/2	40 S 80 X 160	904 1.656 4.630		80 100 120	0.01416 0.01657 0.01898
		2	40 S 80 X 160	236 488 899	16	160 10	0.0252 0.00463
ia_ 5 1 .025 .4 1 20 sg 1 .02 .5 − 1		21/2	40 S 80 X 160	66. 7 91.8 146.3 380. 0		20 30 s 40 x 60	0.00421 0.00504 0.00549 0.00612
	در بر مربع مربع مربع مربع مربع مربع مربع	3	40 s 80 x 160	21.4 28.7 48.3 96.6		80 100 120 140 160	0.00700 0.00804 0.00926 0.01099 0.01244
	Fhousa F+++ 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	3½	40 s 80 x	10.0 13.2	18	10 20 s	0.00247 0.00256 0.00266
· 3 → + .009 > + · - 1 → .008 ≥ 1 → .008 ≥ 1 → .008		4	40 s 80 x 120	5.11 6.75 8.94		30 × 40	0.00276 0.00287 0.00298
$\begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ $		5	40 S 80 X 120 160	18.59 1.59 2.04 2.69 3.69 3.69		60 80 100 120 140 160	0.00335 0.00376 0.00435 0.00504 0.00573 0.00669
$=$ $\frac{1}{1000}$	200 40 =	6	40 S 80 X 120 160	0. 610 0.798 1.015 1.376 1.861	20	10 20 s 30 x 40 60	0.00141 0.00150 0.00161 0.00169 0.00191
		8	20 30 40 s 60 80 x	0.133 0.135 0.146 0.163 0.185		80 100 120 140 160	0.00217 0.00251 0.00287 0.00335 0.00385
7 - 0013 1.0 - 0009 1.0 - 0009 7 - 0009 7 - 0009 1.0 - 0009			100 120 140 x x 160	0.211 0.252 0.289 0.317 0.333	24	10 20 5 30 X 40 60	0.000534 0.000565 0.000597 0.000614 0.000651 0.000741
.90008 0007 .8		10	20 30 40 s 60 x 80	0.0397 0.0421 0.0447 0.0514 0.0569		80 100 120 140 160	0.000835 0.000972 0.001119 0.001274 0.001478
			100 120 140 160	0.0661 0.0753 0.0905 0.1052	Note: in the c bers ir Strong, pipe. res	The letters s olumns of Sch idicate Stand and Double E spectively.	a, x, and xx hedule Num- dard, Extra Extra Strong

TABLE 6.3. Approximate Computation of Pressure Drop of Liquids and Gases in Highly Turbulent Flow in Steel Pipes^a

 $^{a}\Delta P_{109} = C_{1}C_{2}/\rho \text{ psi/100 ft. with } p \text{ in lb/cuft.}$ (Crane Co. Flow of Fluids through Fittings, Values and Pipes, Crane Co., New York, 1982).

EXAMPLE 6.4

Pressure Drop in Nonisothermal Liquid Flow

Oil is oumoed at the rate of 6000 lb/hr through a reactor made of commercial steel pipe 1.278 in. ID and 2000ft long. The inlet condition is 400°F and 750psia. The temperature of the outlet is 930°F and the pressure is to be found. The temperature varies with the distance, L ft, along the reactor according to the equation

 $T = 1500 - 1100 \exp(-0.0003287L)$ (°F)

The viscosity and density vary with temperature according to the equations

$$\mu = \exp\left(\frac{7445.3}{T + 459.6} - 6.1076\right), \quad cP,$$

$$\rho = 0.936 - 0.00036T, \quad g/mL.$$

Round's equation applies for the friction factor:

$$N_{\rm Re} = \frac{4\dot{m}}{\pi D\mu} = \frac{4(6000)}{\pi (1.278/12)2.42\mu} = \frac{29,641}{\mu} ,$$

$$\varepsilon/D = \frac{0.00015(12)}{1.278} = 0.00141,$$

10	! Examp1 e 6 4 ; pressure drop
	i n nonisothermalflow
20	REHD $[,P,0]!$ (D = length inc
	rement
30	DHTH 0,750,200
40	GOSUB 180
50	I1=1
60	GOSUB 150
70	Ī Ī = Ī
80	L=L+D
90	GOSUB 180
100	P=P5*D*(I1+I)
110	GOSUB 1 5 0
120	IF L>1800 THEN 140
130	GOTO70
140	END
150	DISP USING 1 6 0 ; L, T, R1/1000
	,100*F,P
160	IMAGE 0000,2X,000,0,2X,000,0
	2X,0.00,2X,000.0
170	RETURN
180	T=1500-1100*EXP(-(.0003287*L
))
130	M=EXP(7445.3/(T+459.6)-6.107
	6)
200	Ř=.93600036*T
210	R1=29641/M
220	F=1.6364/LOG(.135*.00141+6.5
	ZR1)^2
230	I= 568*F/R

240 RETURN

$$f = \frac{1.6364}{\left[\ln\left[0.135(0.00141) + 6.5/N_{\rm Re}\right]^2\right]}$$

The differential pressure is given by

$$-dP = \frac{8\dot{m}^2}{g_c \pi^2 \rho D^5} f \, dL = \frac{8(600/3600)^2}{32.2\pi^2 62.4\rho (1.278/12)^5 (144)} f \, dL$$
$$= \frac{0.568f}{\rho} dL, \text{ psi},$$
$$P = 750 - \int_0^L \frac{0.586f}{L} dL = 750 - \int_0^L I \, dL.$$

The pressure profile is found by integration with the trapezoidal rule over 200 ft increments. The computer program and the printout are shown. The outlet pressure is 700.1 psia.

For comparison, taking an average temperature of 665°F,

$$\mu = 1.670, \ \rho = 0.697$$

$$N_{\text{Re}} = 17,700, \ f = 0.00291,$$

$$P_{\text{out}} = 102.5.$$

L	Т	1086	100f	P
0 200 400 600 800 1200 1400 1600	400.0 470.0 535.5 596.9 654.4 708.2 758.5 805.7 349.9 891.3	2 . 3 4.4 7.5 11.6 16.7 22.7 29.5 37.1 45.2 53.8	4.85 3.99 3.497 3.10 2.95 2.80 2.61 2.55 2.51	750750 97436 379 67328 727.9 723.2 718.5 713.9 709.3 709.3
2000	930.0	62.7	2.47	700.1

TABLE 6.4. Equivalent Lengths of Pipe Fittings^a

	D	Ŋ	D		ģ	ā		ŋ
Pipe size, in.	Standard eil	Vedium radius ell	Long- radius ell	45-deg ell	Tee	G ate valve, open	Globe valve, open	Swing check, open
$ \begin{array}{c} 1\\ 2\\ 3\\ 4\\ 5\\ 6\\ 8\\ 10\\ 12\\ 14\\ 16\\ 18\\ 20\\ 24\\ 36\\ \end{array} $	2.7 5.5 8.1 11.0 14.0 16.0 21 26 32 36 42 46 52 63 94	2.3 4.6 6.8 9.1 12.0 14.0 18.0 22 26 31 35 40 43 53 79	$ \begin{array}{r} 1.7\\3.5\\5.1\\7.0\\8.9\\11.0\\14.0\\17.0\\20.0\\23\\27\\30\\34\\40\\60\end{array} $	1.3 2.5 3.8 5.0 6.1 7.7 10.0 13.0 15.0 17.0 19.0 21 23 28 43	$5.8 \\ 11.0 \\ 17.0 \\ 22 \\ 27 \\ 33 \\ 43 \\ 56 \\ 66 \\ 76 \\ 87 \\ 100 \\ 110 \\ 140 \\ 200 \\ $	$\begin{array}{c} 0.6 \\ 1.2 \\ 1.7 \\ 2.3 \\ 2.9 \\ 3.5 \\ 4.5 \\ 5.7 \\ 6.7 \\ 8.0 \\ 9.0 \\ 10.2 \\ 12.0 \\ 14.0 \\ 20.0 \end{array}$	27 57 85 110 140 160 220 290 340 390 430 500 560 680 1.000	6.7 13 20 27 33 40 53 67 80 93 107 120 134 160 240

^a Length of straight pipe (ft) giving equivalent resistance.

(Hicks and Edwards, Pump Application Engineering, McGraw-Hill, New York, 1971).

POWER REQUIREMENTS

A convenient formula in common engineering units for power consumption in the transfer of liquids is

$$\dot{P} = \frac{(\text{volumetric flow rate})(\text{pressure difference})}{(\text{equipment efficiency})} \\ = \frac{(\text{gals/min})(\text{lb/sq in.})}{1714(\text{fractional pump eff})(\text{fractional driver eff})} \text{ horsepower.}$$
(6.30a)

Efficiency data of drivers are in Chapter 4 and of pumps in Chapter 7. For example, with 500 gpm, a pressure difference of 7.5 psi, pump efficiency of 0.7, and driver efficiency of 0.9, the power requirement is 32.9 HP or 24.5 kw.

6.4. PIPELINE NETWORKS

A system for distribution of fluids such as cooling water in a process plant consists of many interconnecting pipes in series, parallel, or branches. For purposes of analysis, a point at which several lines meet is called a node and each is assigned a number as on the figure of Example 6.6. A flow rate from node *i* to node *j* is designated as Q_{ij} ; the same subscript notation is used for other characteristics of the line such as f, L, D, and N_{Re} .

Three principles are applicable to establishing flow rates, pressures, and dimensions throughout the network:

1. Each node *i* is characterized by a unique pressure P_i .

A material balance is preserved at each node: total flow in equals total flow out, or net flow equals zero. 3. The friction equation $P_i - P_j = (8\rho/g_c\pi^2)f_{ij}L_{ij}Q_{ij}^2/D_{ij}^5$ applies to the line connecting node *i* with j.

In the usual network problem, the terminal pressures, line lengths, and line diameters are specified and the flow rates throughout are required to be found. The solution can be generalized, however, to determine other unknown quantities equal in number to the number of independent friction equations that describe the network. The procedure is illustrated with the network of Example 6.6.

The three lines in parallel between nodes 2 and 5 have the same pressure drop $P_2 - P_5$. In series lines such as 37 and 76 the flow rate is the same and a single equation represents friction in the series:

$$P_3 - P_6 = kQ_{36}^2(f_{37}L_{37}/D_{37}^5 + f_{76}L_{76}/D_{76}^5).$$

The number of flow rates involved is the same as the number of lines in the network, which is 9, plus the number of supply and destination lines, which is 5, for a total of 14. The number of material balances equals the number of nodes plus one for the overall balance, making a total of 7.

The solution of the problem requires 14 - 7 = 7 more relations to be established. These are any set of 7 friction equations that involve the pressures at all the nodes. The material balances and pressure drop equations for this example are tabulated.

From Eqs. (4)-(10) of Example 6.6, any combination of seven quantities Q_{ij} and/or L_{ij} and/or D_{ij} can be found. Assuming that the Q_{ij} are to be found, estimates of all seven are made to start, and the corresponding Reynolds numbers and friction factors are found from Eqs. (2) and (3). Improved values of the Q_{ij} then are found



TABLE 6.5. Velocity Head Factors of Pipe Fittings^a

^a $h = Ku^2/2g_c$, ft of fluid. (Hydraulic Institute, Cleveland, OH, 1957).

TABLE 6.6. Velocity Head Factors of Pipe Fittings'

Fitting type			_ <i>K</i> 1	<i>K</i> ∞_	
		Standard $(R/D = 1)$, screwed Standard $(R/D = 1)$, flanged/welded Long-radius $(R/D = 1.5)$. all types		800 800 800	0.40 0.25 0.20
Elbows;	90°	Mitered elbows (<i>R/D</i> =1.5)	1 Weld $(90^{\circ} angle)$ 2 Weld $(45^{\circ} angles)$ 3 Weld $(30^{\circ} angles)$ 4 Weld $(22\%^{\circ} angles)$ 5 Weld $(18^{\circ} angles)$	1,000 800 800 800 800	1.15 0.35 0.30 0.27 0.25
	45°	Standard (<i>R</i> Long-radius Mitered, 1 w Mitered, 2 w	Standard ($R/D = 1$), all types Long-radius ($R/D = 1.5$), all types Mitered, 1 weld, 45° angle Mitered, 2 weld, $221/2^\circ$ angles		0.20 0.15 0.25 0.15
	1 80°	Standard $(R/D = 1)$, screwed Standard $(R/D = 1)$, flanged/welded Long radius $(R/D = 1.5)$. all types		1,000 1,000 1,000	0.60 0.35 0.30
Tees	Used as elbow	Standard, s Long-radius, Standard, fla Stub-in-type	crewed screwed anged or welded branch	500 800 800 1,000	0.70 0.40 0.80 1 .00
	Run- through l tee	Screwed Flanged or w Stub-in-type	velded branch	200 150 100	0.10 0.50 0.00
	Sate, ball, plug	Full line size, Reduced trin Reduced trin	$\beta = 1.0$ $n, \beta = 0.9$ $n, \beta = 0.8$	300 500 1,000	0.10 0.15 0.25
Valves	Globe, s Globe, a Diaphrag Butterfly	standard ngle or Y-type gm,dam type /		1,500 1,000 1,000 800	4.00 2.00 2.00 0.25
	Check	Lift Swing Tilting-disk		2,000 1,500 1,000	10.00 1.50 0.50

Note: Use R/D = 1.5 values for R/D = 5 pipe bends. 45° to 180° . Use appropriate tee values for flow through crosses.

^a Inlet, flush, K = $160/N_{\rm Re}$ + 0.5. Inlet, intruding, K = $160/N_{\rm Re}$ = 1.0. Exit, K = 1 .0. K = $K_1/N_{\rm Re} + K_2(1 + I/D)$, with D in inches. [Hooper, Chem. Eng. 96-100 (24 Aug. 1981)].

from Eqs. (4)-(10) with the aid of the Newton-Raphson method for simultaneous nonlinear equations.

Some simplification is permissible for water distribution systems in metallic pipes. Then the Hazen-Williams formula is adequate, namely

Ah =
$$\Delta P/\rho$$
 = 4.727 $L(Q/130)^{1.852}/D^{4.8704}$ (6.31)

with linear dimensions in ft and Q in cuft/sec. The iterative solution method for flowrate distribution of Hardy Cross is popular. Examples of that procedure are presented in many books on fluid mechanics, for example, those of Bober and Kenyon (Fluid Mechanics, Wiley, New York, 1980) and Streeter and Wylie (Fluid Mechanics, McGraw-Hill, New York, 1979).

With particularly simple networks, some rearrangement of equations sometimes can be made to simplify the solution. Example 6.7 is of such a case.

6.5. OPTIMUM PIPE DIAMETER

In a chemical plant the capital investment in process piping is in the range of 25-40% of the total plant investment, and the power consumption for pumping, which depends on the line size, is a substantial fraction of the total cost of utilities. Accordingly, economic optimization of pipe size is a necessary aspect of plant design. As the diameter of a line increases, its cost goes up but is accompanied by decreases in consumption of utilities and costs of pumps and drivers because of reduced friction. Somewhere there is an optimum balance between operating cost and annual capital cost.

For small capacities and short lines, near optimum line sizes may be obtained on the basis of typical velocities or pressure drops such as those of Table 6.2. When large capacities are involved and lines are long and expensive materials of construction are needed, the selection of line diameters may need to be subjected to complete economic analysis. Still another kind of factor may need to be taken into account with highly viscous materials: the possibility that heating the fluid may pay off by reducing the viscosity and consequently the power requirement.

Adequate information must be available for installed costs of piping and pumping equipment. Although suppliers quotations are desirable, published correlations may be adequate. Some data and references to other published sources are given in Chapter 20. A simplification in locating the optimum usually is permissible by ignoring the costs of pumps and drivers since they are essentially insensitive to pipe diameter near the optimum value. This fact is clear in Example 6.8 for instance and in the examples worked out by Happel and Jordan (Chemical Process Economics, Dekker, New York. 1975).

Two shortcut rules have been derived by Peters and Timmerhaus (1980; listed in Chapter 1 References) for optimum diameters of steel pipes of l-in. size or greater, for turbulent and laminar flow:

$$D = 3.9Q^{0.45}\rho^{0.13}, \text{ turbulent flow},$$
(6.32)
$$D = 3.0Q^{0.36}\mu^{0.18}, \text{ laminar flow},$$
(6.33)

$$D = 3.0Q^{0.05}\mu^{0.15}, \text{ laminar flow.}$$
(6.33)

D is in inches, Q in cuft/sec, ρ in lb/cuft, and μ in cP. The factors involved in the derivation are: power cost = 0.055/kWh, friction loss due to fittings is 35% that of the straight length, annual fixed charges are 20% of installation cost, pump efficiency is 50%, and cost of 1-in. IPS schedule 40 pipe is \$0.45/ft. Formulas that take additional factors into account also are developed in that book.

Other detailed studies of line optimization are made by Happel and Jordan (Chemical Process Economics, Dekker, New York, 1975) and by Skelland (1967). The latter works out a problem in simultaneous optimization of pipe diameter and pumping temperature in laminar flow.

Example 6.8 takes into account pump costs, alternate kinds of drivers, and alloy construction.

6.6. NON-NEWTONIAN LIQUIDS

Not all classes of fluids conform to the frictional behavior described in Section 6.3. This section will describe the commonly recognized types of liquids, from the point of view of flow behavior, and will summarize the data and techniques that are used for analyzing friction in such lines.

VISCOSITY BEHAVIOR

The distinction in question between different fluids is in their viscosity behavior, or relation between shear stress τ (force per unit area) and the rate of deformation expressed as a lateral velocity

EXAMPLE 6.5 Comparison of Pressure Drops in a Line with Several Sets of Fittings Resistances

The flow considered is in a 12-inch steel line at a Reynolds number of 6000. With $\varepsilon = 0.00015$, Round's equation gives f = 0.0353. The line composition and values of fittings resistances are:

	Table 6.4	Table 6.5		Table 6.6			
	L	K	<i>K</i> 1	K ₂	к		
Line 6 LR ells 4 tees, branched 2 gate valves, open 1 globe valve	1000 20 7 340	0.25 0.5 0.05 5.4	500 150 300 1500	0.15 0.15 0.10 4.00	0.246 0.567 0.158 4.58		
	1738	9.00			8.64		

EXAMPLE 6.6

A Network of Pipelines in Series, Parallel, and Branches: the Sketch, Material Balances, and Pressure Drop Equations Pressure drop:

$$\Delta P_{ij} = (8\rho/g_c \pi^2) f_{ij} L_{ij} Q_{ij}^2 / D_{ij}^5 = k f_{ij} L_{ij} Q_{ij}^2 / D_{ij}^5.$$
(1)



Reynolds number:

$$(N_{\rm Re})_{ij} = 4Q_{ij}\rho/\pi D_{ij}\mu.$$

Table 6.4,
$$\frac{\Delta P}{(\rho u^2/2g_c)} = \frac{f}{D}(1738) = 61.3,$$

Table 6.5, $\frac{AP}{(\rho u^2/2g_c)} = f\frac{L}{D} + \sum K_i$
 $= \frac{0.0353(1000)}{1} + 9.00 = 44.3,$
Table 6.6, $\frac{AP}{(\rho u^2/2g_c)} = 35.3 + 8.64 = 43.9.$

The value K = 0.05 for gate valve from Table 6.5 appears to be low: *Chemical Engineering Handbook*, for example, gives 0.17, more nearly in line with that from Table 6.6. The equivalent length method of Table 6.4 gives high pressure drops; although convenient, it is not widely used.

Friction factor:

$$f_{ij} = 1.6364 / [\ln(\varepsilon / D_{ij} + 6.5 / (N_{\rm Re})_{ij})]^2.$$
(3)

Pressure drops in key lines:

1

2 3 4

5 6

(2)

$$\Delta p_{12} = P_1 - P_2 - k f_{12} L_{12} Q_{12}^2 / D_{12}^5 = 0, \tag{4}$$

$$\Delta p_{23} = P_2 - P_3 - k f_{23} L_{23} Q_{23}^2 / D_{23}^2 = 0,$$
(5)

$$\Delta p_{25} = P_2 - P_5 = k f_{25}^{(1)} L_{25}^{(1)} (Q_{25}^{(1)})^2 / (D_{25}^{(1)})^5 \tag{6}$$

$$= kf_{25}^{(3)}L_{25}^{(3)}(Q_{25}^{(3)})^{2}/(D_{25}^{(3)})^{3}$$
(7)

$$= k_{1} \sum_{25} L_{25} (Q_{25})^{-} / (D_{25})^{-},$$
(8)
$$A p_{12} - P_{2} = P_{2} = k_{12} L_{12} Q_{22}^{2} / D_{25}^{5} - 0$$
(9)

$$\Delta p_{56} = P_5 - P_6 - kf_{56}L_{56}Q_{56}^2/D_{56}^5 = 0$$
(10)

Node Material Balance at Node:

$$Q_{01} - Q_{12} - Q_{14} = 0$$
(11)
$$Q_{10} - Q_{20} - Q_{21}^{(1)} - Q_{22}^{(2)} - Q_{25}^{(3)} = 0$$
(12)

$$Q_{03} + Q_{23} - Q_{36} = 0 \tag{13}$$

$$Q_{14} - Q_{40} - Q_{45} = 0 \tag{14}$$

$$\begin{array}{c} Q_{45} + Q_{25} + Q_{25} + Q_{25} + Q_{50} = Q_{56} = 0 \\ Q_{55} + Q_{55} - Q_{55} = 0 \end{array} \tag{15}$$

Overall
$$Q_{01} + Q_{03} - Q_{40} - Q_{50} - Q_{60} = 0$$
 (17)

EXAMPLE 6.7

Flow of Oil in a Branched Pipeline

The pipeline handles an oil with sp gr = 0.92 and kinematic viscosity of **5 centistokes(cS)** at a total rate of **12,000 cuft/hr**. All three pumps have the same output pressure. At point 5 the elevation is **100** ft and the pressure is 2 atm gage. Elevations at the other points are zero. Line dimensions are tabulated following. The flow rates in each of the lines and the total power requirement will be found.

Line	L (ft)	D (ft)
14	1000	0.4
24	2000	0.5
34	1500	0.3
45	4000	0.75



EXAMPLE 6.7-(continued)

$$\varepsilon = 0.00015$$
 ft,
 $h_f = \frac{8fLQ^2}{g_c \pi^{-2} D^5} = 0.0251 fLQ^2 / D^5$ ft, (3)

$$\begin{aligned} h_{f_1} &= h_{f_2} = h_{f_3}, \\ \underline{f_1 L_1 Q_1^2} &= \underline{f_2 L_2 Q_2^2} = \underline{f_3 L_3 Q_3^2} \end{aligned} \tag{4}$$

$$\frac{D_1^5}{D_1^5} = \frac{D_2^5}{D_2^5} = \frac{D_3^5}{D_3^5}$$
(5)
$$Q_2 = Q_1 \left[\frac{f_1 L_1}{C_1} \left(\frac{D_2}{D_2} \right)^5 \right]^{1/2} = 1.2352 \left(\frac{f_1}{c_1} \right)^{1/2} Q_1,$$
(6)

$$Q_{3} = Q_{1} \left[\frac{f_{1}L_{1}}{f_{3}L_{3}} \left(\frac{D_{3}}{D_{1}} \right)^{5} \right]^{1/2} = 0.3977 \left(\frac{f_{1}}{f_{3}} \right)^{1/2} Q_{1},$$
(7)

$$Q_1 \left[1 + 1.2352 \left(\frac{f_1}{f_2} \right)^{1/2} + 0.3977 \left(\frac{f_1}{f_3} \right)^{1/2} \right] = Q_4 = 3.333,$$
(8)

$$f = \frac{1.6364}{[\ln(2.03(10^{-5})/D + 6.5/N_{\rm Re}]^2}.$$

For line 45,

$$(N_{\text{Re}})_4 = 31542 \ (3.333) = 105,140,$$

 $f_4 = 0.01881,$
 $(h_f)_{45} = \frac{0.02517(0.01881)(4000)(3.333)^2}{(0.75)^5} = 88.65 \,\text{ft}$

Procedure:

- 1. As a first trial assume $f_1 = f_2 = f_3$, and find $Q_1 = 1.266$ from Eq. (8).
- 2. Find Q_2 and Q_3 from Eqs. (6) and (7).
- 3. With these values of the Q_i , find improved values of the f_i and hence improved values of Q_2 and Q_3 from Eqs. (6) and (7).
- 4. Check how closely $Q_1 + Q_2 + Q_3 3.333 = 0.$
- 5. If check is not close enough, adjust the value of Q_1 and repeat the calculations.

The two trials shown following prove to be adequate.

Q 1	a	l ₂ Q ₃	Q₄	10/3-	- Q4	f ₁
1.2660	1. 5	757 0. 4739	3. 3156	0. 00	23	0. 02069
1. 2707	1. 5	554 0. 5073	3. 3334	0.00	01	0. 02068
Summary	:					
	Li ne	N _{Re}	f	٥	h _r	
1	14	75, 152	0. 02068	1.2707	82.0	8
	24	60, 121	0.02106	1.5554	82.0	8
	34	99, 821	0. 02053	0.5073	82.0	8
	45	105, 140	0.01881	3. 3333	88.6	5

$$h_{f_{14}} = h_{f_{24}} = h_{f_{34}} = \frac{0.02517(0.02068)(1000)(1.2707)^2}{(0.4)^5} = 82.08 \text{ ft.}$$

Velocity head at discharge:

$$\frac{u_5^2}{2g_c} = \frac{1}{2g_c} \left(\frac{Q_4}{(\pi/4)D^2}\right)^2 = 0.88 \text{ ft.}$$

Total head at pumps:

$$h_p = \frac{2(2117)}{0.92(62.4)} + 100 + 0.88 + 82.08 + 88.65 = 345.36 \text{ ft.}$$

Power

$$= \gamma Q_4 h_p$$

0.92(62.4)(10/3)345.36
= 66,088 ft lb/sec
120.2 HP, 89.6 kW.

- ! Example 67 / t 1 OW in a bran 10 ched pipeline READ D1,02,03,L1,L2,L3 DATA .4,.5,.3,1000,2000,1500 20 30 INPUT Q1 48 02=1.2352*****01 03=.3977*****01 50 60 70 R1=59142*Q1 80 R2=47313*Q1 90 R3=78556*Q1 100 F1=1.6364/LOG(.135*.00015/D1 +6.5/R1)^2 110
- 110 F2=1.6364/LOG(.135*.00015/D2 +6.5/R2)^2 120 F3=1.6364/LOG(.135*.00015/D3 +6.5/R3)^2
- +6.5/R3)^2 130 Q2=1.2352*Q1*(F1/F2)^.5! im
- proved value 1 4 0 Q3= 3977*Q1*(F1/F3)^.5 ! imp roved value
- roved value 150 X=10/3-Q1-Q2-Q3! should be less than 0.0001 160 DISP X,Q1,Q2,Q3
- 160 DISPX, Q1, Q2, Q3 170 GOTO 40 ! choose another val ue of Q1 if condition of ste
- ue of W1 if condition of ste ≥ 150 is not satisfied 180 END.

EXAMPLE 6.8

Economic Optimum Pipe Size for Pumping Hot Oil with a Motor or Turbine Drive

A centrifugal pump and its spare handle 1000 gpm of an oil at 500° F. Its specific gravity is 0.81 and its viscosity is 3.0 cP. The length of the line is 600 ft and its equivalent length with valves and other fittings is 900 ft. There are 12 gate valves, two check valves, and one control valve.

Suction pressure at the pumps is atmospheric; the pump head exclusive of line friction is 120 psi. Pump efficiency is 71%. Material of construction of line and pumps is 316 SS. Operation is 8000 hr/yr.

Characteristics of the alternate pump drives are:

- a. Turbines are 3600 rpm, exhaust pressure is 0.75 bar, inlet pressure is 20 bar, turbine efficiency is 45%. Value of the high pressure steam is \$5.25/1000 lbs; that of the exhaust is \$0.75/1000 lbs.
- h. Motors have efficiency of 90%, cost of electricity is \$0.065/kWh.

Cost data are:

1. Installed cost of pipe is 7.50 \$/ft and that of valves is $600D^{0.7}$ \$ each, where *D* is the nominal pipe size in inches.

EXAMPLE **6.8**—(continued)

- Purchase costs of pumps, motors and drives are taken from Manual of Economic Analysis of Chemical Processes, Institut Francais du Petrole (McGraw-Hill, New York, 1976).
- 3. All prices are as of mid-1975. Escalation to the end of 1984 requires a factor of 1.8. However, the location of the optimum will be approximately independent of the escalation if it is assumed that equipment and utility prices escalate approximately uniformly; so the analysis is made in terms of the 1975 prices. Annual capital cost is 50% of the installed price/year.

The summary shows that a 6-in. line is optimum with motor drive, and an B-in. line with turbine drive. Both optima are insensitive to line sizes in the range of 6-10 in.

Q = 1000/(7.48)(60) = 2.2282 cfs, 227.2 m³/hr,

$$N_{\text{Re}} - \frac{4Q\rho}{\pi D\mu} = \frac{4(2.2282)(0.81)(62.4)}{\pi (0.000672)(3)D} = \frac{71,128}{D},$$

$$f = 1.6364 \left[\ln \frac{0.135(0.00015)}{D} + \frac{6.5}{71,128} \right]^{-2}$$

Pump head:

$$h_p = \frac{120(144)}{0.81(62.4)} + \frac{8fLQ^2}{g\pi^2 D^5}$$

= 341.88 + 124.98f/D⁵ ft.

Motor power:

$$P_m = \frac{Q\rho}{\eta_p \eta_m} h_p = \frac{2.2282(50.54)}{550(0.71(0.90))} h_p$$

= 0.3204h_p, HP

Turbine power:

$$P_t = \frac{2.2282(50.54)}{550(0.71)} h_p = 0.2883 h_p, \quad \text{HP}.$$

Steam

= 10.14 kg/HP (from the "manual")

$$= 10.14(0.2883)(2.204)h_p/1000 = 0.006443h_p$$
, 1000 lb/hr.

Power cost:

Steam cost:

4.5(8000)(1000 lb/hr), \$/yr.

Installed pump cost factors for alloy, temperature, etc (data in the "manual")

= 2[2.5(1.8)(1.3)(0.71)] = 8.2.

Summary:

IPS	4	6	6	10
D (ft)	0.3355	0.5054	0.6651	0.8350
100 <i>f</i>	1	.89 1	.67 1.89) 1.93
h_p (ft)	898	413	360	348
Pump efficiency	0.71	0.71	0.71	0.71
motor (kW)	214.6	98.7	86.0	83.2
Steam, 1000 lb/hr	5.97	2.66	2.32	2.25
Pump cost, 2 installed	50,000	28,000	28,000	28,000
Motor cost, 2 installed	36,000	16,000	14,000	14,000
Turbine cost, 2 installed	56,000	32,000	28,000	28,000
Pipe cost	18,000	27,000	36,000	45,000
Valve cost	23,750	31,546	38,584	45,107
Equip cost, motor drive	127,750	93,546	107,584	123,107
Equip cost, turbine drive	147,750	109,546	121,584	137,107
Power cost (\$/yr)	111,592	51,324	44,720	43,264
Steam cost (\$/yr)	208,440	95,760	83,520	80,834
Annual cost, motor drive	175,467	98,097	98,512	104,817
Annual cost, turbine drive	282,315	150,533	144,312	149,387

gradient, $\dot{\gamma} = du/dx$. The concept is represented on Figure 6.2(a): one of the planes is subjected to a shear stress and is translated parallel to a fixed plane at a constant velocity but a velocity gradient is developed between the planes. The relation between the variables may be written

$$\mathbf{f} = F/\mathbf{A} = \mu(du/dx) = \mu \dot{\gamma}, \qquad (6.34)$$

where, by definition, μ is the viscosity. In the simplest case, the viscosity is constant, and the fluid is called Newtonian. In the other cases, more complex relations between τ and $\dot{\gamma}$ involving more than one constant are needed, and dependence on time also may be present. Classifications of non-Newtonian fluids are made according to the relation between τ and $\dot{\gamma}$ by formula or shape of plot, or according to the mechanism of the resistance of the fluid to deformation.

The concept of an apparent viscosity

$$\mu_a = \tau / \dot{\gamma} \tag{6.35}$$

is useful. In the Newtonian case it is constant, but in general it can be a function of τ , $\dot{\gamma}$, and time θ .

Non-Newtonian behavior occurs in solutions or melts of

polymers and in suspensions of solids in liquids. Some $\tau - \dot{\gamma}$ plots are shown in Figure 6.2, and the main classes are described following.

1. Pseudoplastic liquids have a $\tau - \dot{\gamma}$ plot that is concave downward. The simplest mathematical representation of such relations is a power law

$$\boldsymbol{\tau} = \boldsymbol{K} \dot{\boldsymbol{\gamma}}^n, \quad \boldsymbol{n} < 1 \tag{6.36}$$

with n < 1. This equation has two constants; others with many more than two constants also have been proposed. The apparent viscosity is

$$\mu_a = \tau / \dot{\gamma} = K / \dot{\gamma}^{1-n}. \tag{6.37}$$

Since n is less than unity, the apparent viscosity decreases with the deformation rate. Examples of such materials are some polymeric solutions or melts such as rubbers, cellulose acetate and napalm; suspensions such as paints, mayonnaise, paper pulp, or detergent slurries; and dilute suspensions of inert solids. Pseudoplastic properties of wallpaper paste account for good spreading and adhesion, and those of printing inks prevent their running at low speeds yet allow them to spread easily in high speed machines.

2. Dilatant liquids have rheological behavior essentially



Shearing stress (e)

(f)

Figure 6.2. Relations between shear stress, deformation rate, and viscosity of several classes of fluids. (a) Distribution of velocities of a fluid between two layers of areas A which are moving relatively to each other at a distance x under influence of a force F. In the simplest case, $F/A = \mu(du/dx)$ with μ constant. (b) Linear plot of shear stress against deformation. (c) Logarithmic plot of shear stress against deformation rate. (d) Viscosity as a function of shear stress. (e) Time-dependent viscosity behavior of a rheopectic fluid (thixotropic behavior is shown by the dashed line). (f) Hysteresis loops of time-dependent fluids (arrows show the chronology of imposed shear stress).

opposite those of pseudoplastics insofar as viscosity behavior is concerned. The $\tau - \dot{\gamma}$ plots are concave upward and the power law applies

$$\tau = K\dot{\gamma}^n, \quad n > 1, \tag{6.38}$$

but with *n* greater than unity; other mathematical relations also have been proposed. The apparent viscosity, $\mu_a = K\dot{\gamma}^{n-1}$, increases with deformation rate. Examples of dilatant materials are pigment-vehicle suspensions such as paints and printing inks of high concentrations; starch, potassium silicate, and gum **arabic** in water; quicksand or beach sand in water. Dilatant properties of wet cement aggregates permit tamping operations in which small impulses produce more complete settling. Vinyl resin **plastisols** exhibit pseudoplastic behavior at low deformation rates and dilatant behavior at higher ones.

3. Bingham plastics require a finite amount of shear stress before deformation begins, then the deformation rate is linear. Mathematically,

$$\boldsymbol{\tau} = \boldsymbol{\tau}_0 + \boldsymbol{\mu}_B (d\boldsymbol{u}/d\boldsymbol{x}) = \boldsymbol{\tau}_0 + \boldsymbol{\mu}_B \dot{\boldsymbol{\gamma}}, \tag{6.39}$$

where μ_B is called the coefficient of plastic viscosity. Examples of materials that approximate Bingham behavior are drilling muds; suspensions of chalk, grains, and thoria; and sewage sludge. Bingham characteristics allow toothpaste to stay on the brush.

4. Generalized Bingham or yield-power law fluids are represented by the equation

$$\tau = \tau_0 + K \dot{\gamma}^n. \tag{6.40}$$

Yield-dilatant (n > 1) materials are rare but several cases of



Figure 6.3. Time-dependent rheological behavior of a rheopectic fluid, a 2000 molecular weight polyester *[after Steg and Katz, J. Appl. Polym. Sci. 9, 3177 (1965)].*

yield-pseudoplastics exist. For instance, data from the literature of a 20% clay in water suspension are represented by the numbers $\tau_0 = 7.3 \text{ dyn/cm}^2$, $K = 1.296 \text{ dyn(sec)}^n/\text{cm}^2$ and n = 0.483 (Govier and Aziz, 1972, p. 40). Solutions of OS-5.0% carboxypolymethylene also exhibit this kind of behavior, but at lower concentrations the yield stress is zero.

5. **Rheopectic fluids** have apparent viscosities that increase with time, particularly at high rates of shear as shown on Figure 6.3. Figure 6.2(f) indicates typical hysteresis effects for such materials. Some examples are suspensions of gypsum in water, bentonite sols, vanadium pentoxide sols, and the polyester of Figure 6.3.

6. Z'hixotropic fluids have a time-dependent rheological behavior in which the shear stress diminishes with time at a constant deformation rate, and exhibits hysteresis [Fig. 6.2(f)]. Among the substances that behave this way are some paints, ketchup, gelatine solutions, mayonnaise, margarine, mustard, honey, and shaving cream. Nondrip paints, for example, are thick in the can but thin on the brush. The time-effect in the case of the thixotropic crude of Figure 6.4(a) diminishes at high rates of deformation. For the same crude, Figure 6.4(b) represents the variation of pressure gradient in a pipe line with time and axial position; the gradient varies fivefold over a distance of about 2 miles after 200 min. A relatively simple relation involving five constants to represent thixotropic behavior is cited by Govier and Aziz (1972, p. 43):

$$\boldsymbol{\tau} = (\boldsymbol{\mu}_0 + c\boldsymbol{\lambda})\dot{\boldsymbol{\gamma}},\tag{6.41}$$

$$d\lambda/d\theta = a - (a + b\dot{\gamma})\lambda. \tag{6.42}$$

The constants μ_0 , *a*, *b*, and c and the structural parameter λ are obtained from rheological measurements in a straightforward manner.

7. Viscoelastic fluids have the ability of partially recovering their original states after stress is removed. Essentially all molten polymers are viscoelastic as are solutions of long chain molecules such as polyethylene oxide, polyacrylamides, sodium **carboxy**methylcellulose, and others. More homely examples are egg whites, dough, jello, and puddings, as well as bitumen and napalm. This property enables eggwhites to entrap air, molten polymers to form threads, and such fluids to climb up rotating shafts whereas purely viscous materials are depressed by the centrifugal force.

Two concepts of deformability that normally are applied only to solids, but appear to have examples of gradation between solids and liquids, are those of shear modulus *E*, which is

$$E = \text{shear stress/deformation},$$
 (6.43)

and relaxation time θ^* , which is defined in the relation between the residual stress and the time after release of an imposed shear stress, namely,

$$\tau = \tau_0 \exp(-\theta/\theta^*). \tag{6.44}$$

A range of values of the shear modulus (in kgf/cm^2) is

Gelatine	
0.5% solution	4 × 10 ⁻
10% solution (jelly)	5 x 10 ⁻²
Raw rubber	1.7×10^{2}
Lead	4.8×10^{4}
Wood (oak)	8 × 10 ⁴
Steel	8 × 10 ⁵



Figure 6.4. Shear and pipeline flow data of a thixotropic Pembina crude oil at 44.5°F. (a) Rheograms relating shear stress and rate of shear at several constant durations of shear (*Ritter and Govier*, Can. J. Chem. Eng. 48, 505 (1970)]. (b) Decay of pressure gradient of the fluid flowing from a condition of rest at 15,000 barrels/day in a 12 in. line [*Ritter and Batycky*, SPE Journal 7, 369 (1967)].

and that of relaxation time (sec) is

Water	3 x 10 ⁻⁶
Castor oil	2 × 10 ⁻³
Copal varnish	2x 10
Colophony (at 55°C)	5x 10
Gelatine, 0.5% solution	8×10^{2}
Colophony (at 12°C)	4x 10 ⁶
Ideal solids	00

Examples thus appear to exist of gradations between the properties of normally recognized true liquids (water) and true solids.

Elastic properties usually have a negligible effect on resistance to flow in straight pipes, but examples have been noted that the resistances of fittings may be as much as 10 times as great for viscoelastic liquids as for Newtonian ones.

PIPELINE DESIGN

The sizing of pipelines for non-Newtonian liquids may be based on **scaleup** of tests made under the conditions at which the proposed line is to operate, without prior determination and correlation of rheological properties. A body of theory and some correlations are available for design with four mathematical models:

$$\tau_{w} = K' (8\bar{V}/D)^{n'} \quad \begin{array}{l} \text{generalized power law,} \\ \text{(Metzner-Reed)} (AIChE J. 1,434, 1955). \\ \end{array}$$

In the last model, the parameters may be somewhat dependent on the shear stress and deformation rate, and should be determined at magnitudes of those quantities near those to be applied in the plant.

The shear stress $\hat{\tau}_w$ at the wall is independent of the model and is derived from pressure drop measurements as

$$\tau_{w} = D\Delta P/4L. \tag{6.49}$$

Friction Factor. In rheological literature the friction factor is defined as

$$f = \frac{D\Delta P}{4L\rho V^2/2g_c} \tag{6.50}$$

$$=\frac{\tau_{w}}{\rho V^2/2g_c}.$$
(6.51)

This value is one-fourth of the friction factor used in Section 6.3. For the sake of consistency with the literature, the definition of Eq. (6.50) will be used with non-Newtonian fluids in the present section.

Table 6.2 lists theoretical equations for friction factors in laminar flows. In terms of the generalized power law, Eq. (6.48),

$$f = \frac{\tau_w}{\rho V^2 / 2g_c} = \frac{K' (8V/D)^{n'}}{\rho V^2 / 2g_c}$$

= $D^{n'} V^{2-n'} \rho / g_c K' 8^{n'-1}$ (6.52)

By analogy with the Newtonian relation, f = 16/Re, the denominator of Eq. (6.52) is designated as a modified Reynolds number,

$$\operatorname{Re}_{MR} = D^{n'} V^{2-n'} \rho / g_c K' 8^{n'-1}.$$
(6.53)

The subscript MR designates Metzner-Reed, who introduced this form.

Scale *Up*. The design of pipelines and other equipment for handling non-Newtonian fluids may be based on model equations with parameters obtained on the basis of measurements with viscometers or with pipelines of substantial diameter. The shapes of plots of τ_w against $\dot{\gamma}$ or 8V/D may reveal the appropriate model. Examples 6.9 and 6.10 are such analyses.

In critical cases of substantial economic importance, it may be advisable to perform flow tests-Q against BP-in lines of moderate size and to scale up the results to plant size, without necessarily trying to fit one of the accepted models. Among the effects that may not be accounted for by such models are time

EXAMPLE 6.9

Analysis of Data Obtained in a Capillary Tube Viscometer Data were obtained on a paper pulp with specific gravity 1.3, and are given as the first four columns of the table. Shear stress τ_w and deformation rate $\dot{\gamma}$ are derived by the equations applying to this kind of viscometer (Skelland, 1967, **p**. 31; Van Wazer et al., 1963, p. 197):

$$\tau_{w} = D \Delta P/4L,$$

$$\dot{\gamma} = \frac{3n'+1}{4n'} \left(\frac{8V}{D}\right)$$

$$n' = \frac{d \ln(\tau_{w})}{d \ln(8V/D)}$$

The plot of log τ_w against log (8V/D) shows some scatter but is approximated by a straight line with equation

$$\tau_w = 1.329(8V/D)^{0.51}$$

Since

$$\dot{\gamma} = (2.53/2.08)(8V/D),$$

the relation between shear stress and deformation is given by the

EXAMPLE 6.10

Parameters of the Bingham Model from Measurements of Pressure Drops in a Line

Data of pressure drop in the flow of a 60% limestone slurry of density 1.607 g/ml were taken by Thomas [*Ind. Eng. Chem. 55*, 18-29 (1963)]. They were converted into data of wall shear stress $\tau_w = D\Delta P/4L$ against the shear rate 8V/D and are plotted on the figure for three line sizes.

The Buckingham equation for Bingham flow in the laminar region is

$$\frac{8V}{D} = \frac{\tau_w}{\mu_B} \left[1 - \frac{4}{3} \left(\frac{\tau_0}{\tau_w} \right) + \frac{1}{3} \left(\frac{\tau_0}{\tau_w} \right)^4 \right]$$
$$\approx \frac{1}{\mu_B} \left(\tau_w - \frac{4}{3} \tau_0 \right)$$

The second expression is obtained by neglecting the fourth-power term. The Bingham viscosity μ_B is the slope of the plot in the laminar region and is found from the terminal points as

$$\mu_B = (73-50)/(347-0) = 0.067 \text{ dyn sec/cm}^2$$
.

From the reduced Buckingham equation,

$$\tau_0 = 0.75 \tau_w$$
 (at $8V/D = 0$)
= 37.5.

Accordingly, the Bingham model is represented by

$$\tau_w = 37.5 + 0.067(8V/D), \, dyn/cm^2$$

with time in seconds.

Transitions from laminar to turbulent flow may be identified off

equation

$$\tau_{w} = 1.203 \dot{\gamma}^{0.51}$$





the plots:

$$D = 2.06$$
 cm, $8V/D = 465$, $V = 120$ cm/sec
4.04 215, 109
7.75 (critical not reached).

The transition points also can be estimated from Hanks' correlation [*AIChE J.* 9, 45, 306 (1963)] which involves these expressions:

$$\begin{aligned} x_c &= (\tau_0/\tau_w)_c, \\ \text{He} &= D^2 \tau_0 \rho / \mu_B^2, \\ x_c / (1 - x_c)^3 &= \text{He} / 16,800, \\ \text{Re}_{,,} &= (1 - \frac{4}{3}x_c + \frac{1}{3}x_c^4)\text{He} / 8x_c \end{aligned}$$

The critical linear velocity finally is evaluated from the critical Reynolds number of the last equation with the following results;

100 11		LOIDS			
EXAMPLES	6.10 —(a	ontinued)			
	D (cm)	10⁻⁴ H e	x _c	Re _{Bc}	V _c
	2.06 4.04 7.75	5.7 22.0 81.0	0.479 0.635 0.750	5 6 3 5 8945 14,272	114 (120) 93 (109) 77

dependence, pipe roughness, pipe fitting resistance, wall slippage, and viscoelastic behavior. Although some effort has been devoted to them, none of these particular effects has been well correlated. Viscoelasticity has been found to have little effect on friction in straight lines but does have a substantial effect on the resistance of pipe fittings. Pipe roughness often is accounted for by assuming that the relative effects of different roughness ratios ε/D are represented by the Colebrook equation (Eq. 6.20) for Newtonian fluids. Wall slippage due to trace amounts of some polymers in solution is an active field of research (Hoyt, 1972) and is not well predictable.

The scant literature on pipeline scaleup is reviewed by Heywood (1980). Some investigators have assumed a relation of the form

$$\tau_w = D\Delta P/4L = kV^a/D^b$$

100

and determined the three constants *K*, a, and *b* from measurements on several diameters of pipe. The exponent a on the velocity appears to be independent of the diameter if the roughness ratio ε/D is held constant. The exponent *b* on the diameter has been found to range from 0.2 to 0.25. How much better this kind of analysis is than assuming that a = b, as in Eq. (6.48) has not been established. If it can be assumed that the effect of differences in ε/D is small for the data of Examples 6.9 and 6.10, the measurements should plot as separate lines for each diameter, but such a distinction is not obvious on those plots in the laminar region, although it definitely is in the turbulent region of the limestone slurry data.

Observations of the performance of existing large lines, as in the case of Figure 6.4, clearly yields information of value in analyzing the effects of some changes in operating conditions or for the design of new lines for the same system.

Laminar Flow. Theoretically derived equations for volumetric flow rate and friction factor are included for several models in Table 6.7. Each model employs a specially defined Reynolds number, and the Bingham models also involve the Hedstrom number,

$$He = \tau_0 \rho D^2 / \mu_B^2. \tag{6.54}$$

These dimensionless groups also appear in empirical correlations of the turbulent flow region. Although even in the approximate Eq. (9) of Table 6.7, group He appears to affect the friction factor, empirical correlations such as Figure 6.5(b) and the data analysis of Example 6.10 indicate that the friction factor is determined by the Reynolds number alone, in every case by an equation of the form, f = 16/Re, but with Re defined differently for each model. Table 6.7 collects several relations for laminar flows of fluids.

Transitional Flow. Reynolds numbers and friction factors at which the flow changes from laminar to turbulent are indicated by the breaks in the plots of Figures 6.4(a) and (b). For Bingham models, data are shown directly on Figure 6.6. For power-law liquids an equation for the critical Reynolds number is due to Mishra and Triparthi [*Trans. IChE* 51, T141 (1973)],

$$\operatorname{Re}_{c}^{\prime} = \frac{1400(2n+1)(5n+3)}{(3n+1)^{2}}$$
(6.55)

The numbers in parentheses correspond to the break points on the figure and agree roughly with the calculated values.

The solution of this problem is based on that of Wasp et al. (1977).

The Bingham data of Figure 6.6 are represented by the equations of Hanks [AIChE J. 9, 306 (1963)],

$$(\mathbf{Re}_{B})_{c} = \frac{\mathrm{HE}}{8x_{c}} \left(1 - \frac{4}{3}x_{c} + \frac{1}{3}x_{c}^{4} \right), \tag{6.56}$$

$$\frac{x_c}{(1-x_c)^3} = \frac{\text{He}}{16,800}.$$
(6.57)

They are employed in Example 6.10.

Turbulent Flow. Correlations have been achieved for all four models, Eqs. (6.45)-(6.48). For power-law flow the correlation of Dodge and Metzner (1959) is shown in Figure 6.5(a) and is represented by the equation

$$\frac{1}{\sqrt{f}} = \frac{4.0}{(n')^{0.75}} \log_{10} [\operatorname{Re}_{n'} f^{(1-n'/2)}] - \frac{0.40}{(n')^{1.2}}.$$
(6.58)

These authors and others have demonstrated that these results can represent liquids with a variety of behavior over limited ranges by

TABLE 6.7. Laminar Flow: Volumetric Flow Rate, Friction Factor, Reynolds Number, and Hedstrom Number

Newtonian

$$f = 16/\text{Re}, \quad \text{Fie} = DV\rho/\mu \tag{1}$$

Power Law [Eq. (6.45)]

$$Q = \frac{\pi D^3}{32} \left(\frac{4n}{3n+1}\right) \left(\frac{\tau_w}{K}\right)^{1/n} \tag{2}$$

$$f = 16/\text{Re}'$$
(3)
$$\text{Re}' = \frac{\rho VD}{\rho (-4n)} \left(\frac{D}{\rho} \right)^{n-1}$$
(4)

$$\mathsf{Re} = \frac{1}{K} \left(\frac{1}{1+3n} \right)^{-1} \left(\frac{8V}{8V} \right)^{-1}$$

Bingham Plastic [Eq. (6.46)]

$$Q = \frac{\pi D^3 \tau_w}{32\mu_B} \left[1 - \frac{4}{3} \frac{\tau_o}{\tau_w} + \frac{1}{3} \left(\frac{\tau_o}{\tau_w} \right)^4 \right]$$
(5)

 $\begin{aligned} \mathsf{Re}_{B} &= D V \rho / \mu_{B} \end{aligned} \tag{6} \\ \mathsf{He} &= \tau_{0} D^{2} \rho / \mu_{B} \end{aligned} \tag{7}$

$$\frac{1}{\text{Re}_{p}} = \frac{f}{166} - \frac{\text{He}^{2}}{(6\text{Re}_{p}^{2} + \frac{\text{He}^{4}}{3f^{2}\text{Re}_{B}^{6}}} \text{ (solve for } f)$$
(8)

$$f \approx \frac{96Re_B^2}{6Re_B + He} \quad \text{[neglecting } (\tau_0/\tau_w)^4 \text{ in Eq.(5)]} \tag{9}$$

Generalized Bingham (Yield-Power Law) [Eq. (6.47)]

$$Q = \frac{\pi D^3}{32} \frac{4n}{3n+1} \left(\frac{\tau_w}{K}\right)^{1/n} \left(1 - \frac{\tau_y}{\tau_w}\right) \times \left\{1 - \frac{\tau_y/\tau_w}{2n+1} \left[1 + \frac{2n}{n+1} \left(\frac{\tau_y}{\tau_w}\right) \left(1 + n\frac{\tau_y}{\tau_w}\right)\right]\right\}$$
(10)

$$r = \frac{1}{\text{Re}'} \left(1 - \frac{1}{f\text{Re}'^2} \right) \\ \times \left\{ 1 - \frac{1}{(2n-1)} \frac{2\text{He}}{f\text{Re}'^2} \left[1 + \frac{2n}{(n+1)} \frac{2\text{He}}{f\text{Re}'^2} \left(1 + n \cdot \frac{2\text{He}}{f\text{Re}'^2} \right) \right] \right\}$$
(11)
[Re' by Ea. (4) and He by Eq. (7)]



Figure 6.5. Friction factors in laminar and turbulent flows of power-law and Bingham liquids. (a) For pseudoplastic liquids represented by $\tau_w = K'(8V/D)^{n'}$, with K' and n' constant or dependent on $\tau_w: 1/\sqrt{f} = [4.0/(n')^{0.75}] \log_{10}[\operatorname{Re}_{n'} f^{(n-2)}] - 0.40/(n')^{1.2}$, [Dodge and Metzner, AIChE J. 5, 189 (1959)]. (b) For Bingham plastics, Re, $= DV\rho/\mu_B$, He = $\tau_0 D^2 \rho/\mu_B^2$ [Planks and Dadia, AIChE J. 17,554 (1971)].

evaluating K' and n' in the range of shear stress $\tau_w = D\Delta P/4L$ that will prevail in the required situation.

Bingham flow is represented by Figure 6.5(b) in terms of Reynolds and Hedstrom numbers.

Theoretical relations for generalized Bingham flow [Eq. (6.47)] have been devised by Torrance [S. *Afr. Mech. Eng.* 13, 89 (1963)]. They are

$$\frac{1}{\sqrt{f}} = \left(\frac{2.69}{n-2} \cdot 95\right) + \frac{1.97}{n}\ln(1-x) + \frac{1.97}{n}\ln(\operatorname{Re}_T' f^{1-n/2}) + \frac{0.68}{n}(5n-8)$$
(6.59)

with the Reynolds number

$$Re, = D^n V^{2-n} \rho / 8^{n-1} K \tag{6.60}$$

and where

$$\mathbf{r} = \tau_0 / \tau_w. \tag{6.61}$$

In some ranges of operation, materials may be represented approximately equally well by several models, as in Example 6.11 where the power-law and Bingham models are applied.

6.7. GASES

The differential energy balances of Eqs. (6.10) and (6.15) with the friction term of Eq. (6.18) can be integrated for compressible fluid flow under certain restrictions. Three cases of particular importance are of isentropic or isothermal or adiabatic flows. Equations will be developed for them for ideal gases, and the procedure for **nonideal** gases also will be indicated.

ISENTROPIC FLOW

In short lines, nozzles, and orifices, friction and heat transfer may be neglected, which makes the flow essentially isentropic. Work transfer also is negligible in such equipment. The resulting theory is a basis of design of nozzles that will generate high velocity gases for



Figure 6.6. Critical Reynolds number for transition from laminar to turbulent flow of Bingham fluids. The data also are represented by Eqs. (6.56) and (6.57): (0) cement rock slurry; (A) river mud slurries; (0) clay slurry; (P) sewage sludge; (A) ThO₂ slurries; (\blacksquare) lime slurry. [Hanks and Pratt, SPE Journal, 342-346 (Dec. 1967)].

power production with turbines. With the assumptions indicated, Eq. (6.10) becomes simply

$$dH + (1/g_c)u \, du = 0, \tag{6.62}$$

which integrates into

$$H_2 - H_1 + \frac{1}{2g_c} (u_2^2 - u_1^2) \approx 0. \tag{6.63}$$

One of these velocities may be eliminated with the mass balance,

$$\dot{m} = u_2 A_2 / V_2 = u_1 A_1 / V_1 \tag{6.64}$$

so that

$$u_2^2 - u_1^2 = (\dot{m}V_2/A_2)^2 [1 - (A_2V_1/A_1V_2)^2].$$
(6.65)

For ideal gases substitutions may be made from

$$H_2 - H_1 = C_p (T_2 - T_1) \tag{6.66}$$

and

$$T_2/T_1 = (P_2/P_1)^{(k-1)/k} = (V_1/V_2)^k.$$
 (6.67)

After these substitutions are made into Eq. (6.63), the results may be solved for the mass rate of flow as

$$\dot{m}/A_2 = \frac{\left(\frac{2g_c P_1}{V_1}\right)^{1/2} \left\{\frac{k}{k-1} \left[\left(\frac{P_2}{P_1}\right)^{2/k} - \left(\frac{P_2}{P_1}\right)^{(k+1)/k}\right]\right\}^{1/2}}{\left[1 - \left(\frac{A_2}{A_1}\right)^2 \left(\frac{P_2}{P_1}\right)^{2/k}\right]^{1/2}}.$$
(6.68)

At specified mass flow rate and inlet conditions P_1 and V_1 , Eq. (6.68) predicts a relation between the area ratio A_2/A_1 and the pressure ratio P_2/P_1 when isentropic flow prevails. It turns out that, as the pressure falls, the cross section at first narrows, reaches a minimum at which the velocity becomes sonic; then the cross section increases and the velocity becomes supersonic. In a duct of constant cross section, the velocity remains sonic at and below a critical pressure ratio given by

$$\frac{P_s}{P_1} = \left(\frac{2}{k+1}\right)^{k/(k+1)}.$$
(6.69)

The sonic velocity is given by

$$u_s = \sqrt{g_c(\partial P/\partial \rho)_s} \rightarrow \sqrt{g_c k R T/M_w}, \qquad (6.70)$$

where the last result applies to ideal gases and M_w is the molecular weight.

ISOTHERMAL FLOW IN UNIFORM DUCTS

When elevation head and work transfer are neglected, the mechanical energy balance equation (6.13) with the friction term of Eq. (6.18) become

$$V dP + (1/g_c)u du + \frac{fu^2}{2g_c D} dL = 0.$$
(6.71)

Make the substitutions

$$u = G/\rho = GV \tag{6.72}$$

EXAMPLE 6.11

Pressure Drop in Power-Law and Bingham Plow

A limestone slurry of density 1.693 g/mL is pumped through a 4-in. (152 mm) line at the rate of 4 ft/sec (1.22 m/sec). The pressure drop (psi/mile) will be calculated. The slurry behavior is represented by

- **a.** The power-law with n = 0.165 and K = 34.3 dyn sec^{0.165}/cm² (3.43 Pa sec^{0.165}).
- **h.** Bingham model with $\tau_0 = 53 \text{ dyn/cm}^2 (5.3 \text{ Pa})$ and $\mu_B = 22\text{cP} (0.022 \text{ Pa sec})$.

Power law:

$$Re' = D^{n}V^{2-n}\rho/8^{n-1}K$$

= (0.152)^{0.165}(1.22)^{1.835}(1693)(8)^{0.835}/3.43
= 2957,
f = 0.0058 [Fig. 6.6(a)]

and the ideal gas relation

$$V = P_1 V_1 / P$$
 and $dV / V = -dP / P$ (6.73)

so that Eq. (6.71) becomes

$$\frac{P dP}{P_1 V_1} - \frac{G^2}{g_c} \ln\left(\frac{P_1}{P_2}\right) + \frac{fG^2}{2g_c D} dL = 0.$$
(6.74)

This is integrated term-by-term between the inlet and outlet conditions,

$$\frac{P_2^2 - P_1^2}{2P_1 V_1} + \frac{G^2}{g_c} \ln\left(\frac{P_1}{P_2}\right) + \frac{fG^2 L}{2g_c D} = 0$$
(6.75)

and may be rearranged into

$$P_{2}^{2} = P_{1}^{2} - \frac{2P_{1}V_{1}G^{2}}{g_{c}} \left[\frac{fL}{2D} + \ln\left(\frac{P_{1}}{P_{2}}\right) \right]$$
(6.76)

In terms of a density, ρ_m , at the average pressure in the line,

$$P_2 = P_1 \frac{fG^2 L}{2g_c D\rho_m}.$$
(6.77)

The average density may be found with the aid of an approximate evaluation of P_2 based on the inlet density; a second trial is never justified. Eqs. (6.76) and (6.77) and the approximation of Eq. (6.76) obtained by neglecting the logarithmic term are compared in Example 6.12. The restriction to ideal gases is removed in Section 6.7.4.

ADIABATIC FLOW

The starting point for development of the integrated adiabatic flow energy balance is Eq. (6.71) and again ideal gas behavior will be assumed. The equation of condition of a static adiabatic process, $PV^{k} = \text{const}$, is not applicable to the flow process; the appropriate

$$\frac{A}{L} = \frac{P}{2g_c D} \frac{4f\rho V^2}{2(0.152)^2} \frac{4(0.0058)(1693)(1.22)^2}{2(0.152)}$$

= 192.3 N/(m²)(m) [g_c = kgm/sec²/N],
 \rightarrow 192.3(14.7/101,250)1610 = 45.0 psi/mile

Bingham:

$$Re_{B} = \frac{DV\rho}{\mu_{B}} = \frac{0.152(1.22)(1693)}{0.022} = 14,^{270},$$

$$He = \tau_{0}D^{2}\rho/\mu_{B}^{2} = 5.3(0.152)^{2}(1693)/(0.022)^{2}$$

$$= 428,000,$$

critical Re, = 12,000 (Fig. 6.5),
 $f = 0.007$ [Fig. 6.6(b)],

$$\frac{AP}{L} = \frac{0.007}{0.0058} 45.0 = 54.3 \text{ psi/mile}.$$

one is obtained as follows. Begin with

$$dH = -d\left(\frac{u^2}{2g_c}\right) = \frac{G^2 V \, dV}{g_c} \tag{6.78}$$

$$= C_p dT = \frac{Rk}{k-1} dT = \frac{k}{k-1} d(PV),$$
 (6.79)

from which

$$d(PV) = \left(\frac{k-1}{k}\right)\frac{G^2}{g_c} V dV, \qquad (6.80)$$

and the integral is

$$PV = P_1 V_1 - \left(\frac{k-1}{k}\right) \frac{G^2}{2g_{cc}} \left(V^2 - V_1^2\right).$$
(6.81)

Also

$$V dP = d(PV) - (PV) \frac{dV}{V}$$
(6.82)

Substitutions into Eq. (6.71) result in

$$d(PV) - PV\frac{dV}{V} + \frac{G^2}{g_c}V\,dV + \frac{fG^2}{2g_cD}\,dL = 0.$$
 (6.83)

Further substitutions from Eqs. (6.80) and (6.81) and multiplying through by $2kg_c/G^2V^2$ result in

$$2\frac{dV}{V} - \left[\frac{2kg_c P_1 V_1}{G^2} + (k-1)V_1^2\right| \frac{dV}{V^3} + (k-1)\frac{dV}{V} + \frac{kf}{D}dL = 0.$$
(6.84)

Integrating from V_1 to V_2 and L = 0 to L gives

$$(k+1)\ln\frac{V_2}{V_1} + \frac{1}{2}\left[\frac{2kg_cP_1V_1}{G^2} + (k-1)V_1^2\right]\left(\frac{1}{V_2^2} - \frac{1}{V_1^2}\right) + \frac{kfL}{D} = 0$$
(6.85)

or

$$\frac{fL}{D} = \frac{1}{2k} \left[\frac{2kg_c P_1 V_1}{G^2 V_1^2} + (k-1) \right] \left[1 - \left(\frac{V_1}{V_2}\right)^2 \right] + \frac{k+1}{2k} \ln\left(\frac{V_1}{V_2}\right)^2.$$
(6.86)

In terms of the inlet Mach number,

$$M_{1} = u_{1} / \sqrt{g_{c} k R T / M_{w}} = G V_{1} / \sqrt{g_{c} k R T / M_{w}},$$
 (6.87)

the result becomes

$$\frac{fL}{D} = \frac{1}{2k} \left(k - 1 + \frac{2}{M_1^2} \right) \left[1 - \left(\frac{V_1}{V_2} \right)^2 \right] + \frac{k+1}{2k} \ln \left(\frac{V_1}{V_2} \right)^2. \quad (6.88)$$

When everything else is specified, Eqs. (6.86) or (6.88) may be solved for the exit specific volume V_2 . Then P_2 may be found from Eq. (6.81) or in the rearrangement

$$\frac{P_2 V_2}{P_1 V_1} = \frac{T_2}{T_1} = 1 + \left(\frac{k-1}{2k} M_1^2\right) \left[1 - \left(\frac{V_2}{V_1}\right)^2\right],$$
(6.89)

from which the outlet temperature likewise may be found.

6.8. LIQUID-GAS FLOW IN PIPELINES 111

Although the key equations are transcendental, they are readily solvable with hand calculators, particularly those with root-solving provisions. Several charts to ease the solutions before the age of calculators have been devised: M.B. Powley, Can. J. Chem. Eng., 241-245 (Dec. 1958); C.E. Lapple, reproduced in Perry's Chemical Engineers' Handbook, McGraw-Hill, New York, 1973, p. 5.27; O. Levenspiel, reproduced in Perry's Chemical Engineers' Handbook, McGraw-Hill, New York, 1984, p. 5.31; Hougen, Watson, and Ragatz, Thermodynamics, Wiley, New York, 1959, pp. 710-711.

In all compressible fluid pressure drop calculations it is usually justifiable to evaluate the friction factor at the inlet conditions and to assume it constant. The variation because of the effect of temperature change on the viscosity and hence on the Reynolds number, at the usual high Reynolds numbers, is rarely appreciable.

NONIDEAL GASES

Without the assumption of gas ideality, Eq. (6.71) is

$$\frac{dP}{V} + \frac{G^2}{g_c} \frac{dV}{V} + \frac{fG^2}{2g_c D} dL = 0,$$
(6.90)

In the isothermal case, any appropriate PVT equation of state may be used to eliminate either P or V from this equation and thus permit integration. Since most of the useful equations of state are pressure-explicit, it is simpler to eliminate P. Take the example of one of the simplest of the non-ideal equations, that of van der Waals

$$P = \frac{RT}{V-b} - \frac{a}{V^2},\tag{6.91}$$

of which the differential is

$$dP = \left(-\frac{RT}{(V-b)^2} + \frac{2a}{V^3}\right) dV.$$
 (6.92)

Substituting into Eq. (6.90),

$$\left(-\frac{RT}{(V-b)^2} + \frac{2a}{V^3} + \frac{G^2}{g_c}\right)\frac{dV}{V} + \frac{fG^2}{2g_cD}\,dL = 0.$$
 (6.93)

Although integration is possible in closed form, it may be more convenient to perform the integration numerically. With more accurate and necessarily more complicated equations of state, numerical integration will be mandatory. Example 6.13 employs the van der Waals equation of steam, although this is not a particularly. suitable one; the results show a substantial difference between the ideal and the **nonideal** pressure drops. At the inlet condition, the compressibility factor of steam is z = PV/RT = 0.88, a substantial deviation from ideality.

6.6. LIQUID-GAS FLOW IN PIPELINES

In flow of mixtures of the two phases in pipelines, the liquid tends to wet the wall and the gas to concentrate in the center of the channel, but various degrees of dispersion of each phase in the other may exist, depending on operating conditions, particularly the individual **flow** rates. The main patterns of flow that have been recognized are indicated on Figures 6.7(a) and (b). The ranges of conditions over which individual patterns exist are represented on maps like those of Figures 6.7(c) and (d). Since the concept of a

EXAMPLE 6.12

Adiabatic and Isothermal Flow of a Gas in a Pipeline

Steam at the rate of 7000 kg/hr with an inlet pressure of 23.2 barabs and temperature of 220°C flows in a line that is 77.7 mm dia and 30.5 m long. Viscosity is $28.5(10^{-6})N \text{ sec/m}^2$ and specific heat ratio is k = 1.31. For the pipe, $\varepsilon/D = 0.0006$. The pressure drop will be found in (a) isothermal flow; (b) adiabatic flow. Also, (c) the line diameter for sonic flow will be found.

$$V_1 = 0.0862 \text{ m}^3/\text{kg},$$

$$G = 7000/(3600)(\pi/4)(0.0777)^2 = 410.07 \text{ kg/m}^2 \text{ sec},$$

$$\text{Re}_1 = \frac{DG}{\mu} = \frac{0.0777(410.07)}{28.5(10^{-6})} = 1.12(10^6)$$

$$f = 1.6364/[\ln(0.135)(0.0006) + 6.5/1.2(10^6)]^2 = 0.0187.$$

Inlet sonic velocity:

$$u_{s1} = \sqrt{g_c kRT_1/M_w} = \sqrt{1(1.31)(8314)493.2/18.02} = 546 \text{ m/sec}$$

 $M_1 = u_1/u_{s1} = GV_1/u_{s1} = 410.07(0.0862)/546 = 0.0647.$

As a preliminary calculation, the pressure drop will be found by neglecting any changes in density:

$$\Delta P = \frac{fG^2L}{2g_c D\rho} = \frac{0.0187(410.07)^2(305)}{2(1)(0.0777)(1/0.0862)} = 5.32(10^5) \text{ N/m}^2,$$

$$\therefore P_2 = 23.2 - 5.32 = 17.88 \text{ bar.}$$

(a) Isothermal flow, Use Eq. (6.76):

$$\frac{2P_1V_1G^2}{g_c} = 2(23.2)(10^5)(0.0862)(410.07)^2 = 6.726(10^{10}),$$

$$P_2 = \left[P_1^2 - \frac{2P_1V_{1G}^2}{g_c}\left(\frac{fL}{2D} + \ln\frac{P_1}{P_2}\right)\right]^{1/2}$$

$$= 10^5 \frac{538.24 - 6.726\left(0.0187(305)/2(0.0777) + \ln\frac{23.2(10^5)}{P_2}\right)}{P_2}$$

$$= 17.13(10^5) \text{ N/m}^2,$$

and

AP = 23.2 - 17.13 = 5.07 bar.

When the logarithmic term is neglected,

 $P_2 = 17.07(10)^5 \,\mathrm{N/m^2}.$

(b) Adiabatic flow. Use Eq. (6.88):

$$\frac{fL}{D} = \frac{1}{2k} \left(k - 1 + \frac{2}{M_1^2} \right) \left[1 - \left(\frac{V_1}{V_2} \right)^2 \right] + \frac{k+1}{2k} \ln \left(\frac{V_1}{V_2} \right)^2, \quad (1)$$

$$\frac{0.0187(305)}{0.0777} = \frac{1}{2.62} \left(0.31 + \frac{2}{0.0647^2} \right) \times \left[1 - \left(\frac{V_1}{V_2} \right)^2 \right] + \frac{2.31}{2.62} \ln \left(\frac{V_1}{V_2} \right)^2, \quad (1)$$

$$73.4 = 182.47 \left[1 - \left(\frac{V_1}{V_2} \right)^2 \right] + 0.8817 \ln \left(\frac{V_1}{V_2} \right)^2, \quad (1)$$

$$\therefore \frac{V_1}{V_2} = 0.7715.$$

Equation (6.89) for the pressure:

$$\frac{P_2 V_2}{P_1 V_1} = \frac{T_2}{T_1} = \left[1 + \frac{(k-1)}{2k} M_1^2\right] \left[1 - \left(\frac{V_2}{V_1}\right)^2\right]$$

$$= 1 + \frac{0.31(0.0647)^2}{2.62} [1 - (1.2962)^2]$$

= 0.9997,
$$P_2 = 0.9997 P_1 \left(\frac{V_1}{V_2}\right) = 0.9997(23.2)(10^5)(0.7715)$$

= 17.89(10⁵) N/m²,
$$\Delta P = 23.2 - 17.89 = 5.31 \text{ bar.}$$

(c) Line diameter for sonic flow. The critical pressure ratio is

$$\frac{P_2}{P_1} = \left(\frac{2}{k+1}\right)^{k/(k-1)} = 0.5439, \text{ with } k = 1.31,$$

$$G = \frac{7000/3600}{(\pi/4)D^2} = \frac{2.4757}{D^2},$$

$$M_1 = \frac{GV_1}{U_{s_1}} = \frac{2.4757(0.0862)}{546D^2} = \frac{3.909(10^{-4})}{D^2},$$
(2)

Equation (6.89) becomes

$$0.5439(V_2/V_1) = 1 + 0.1183M_1^2[1 - (V_2/V_1)^2],$$
(3)

$$fL/D = 0.0187(305)/D = 5.7035/D$$
(4)

$$= \text{ rhs of Eq. (6.88).}$$

Procedure

- 1. Assume D. 2. Find M₁ [Eq. (2)].
- 3. Find V_2/V_1 from Eq. (6.89) [Eq. (3)]. 4. Find rhs of Eq. (6.88) [Eq. (1)].
- 5. Find D = 5.7035/[rhs of Eq. (6.88)] [Eq. (4)].
- 6. Continue until steps 1 and 5 agree.

Some trials are:

		Eq. (6.89)	Eq. (6.88)	
D	M ₁	V_{1}/V_{2}	rhs	D
0.06	0. 1086	0. 5457	44. 482	0. 1282
0.07	0.0798	0.5449	83. 344	0.06843
0.0697	0. 08046	0. 5449	81.908	0.06963

: D = 0.0697 m

- 10 ! Example 6. 12. Line dia for sonicflow
- 2 0 K=1.31 30 INPUT D ! (Trial value>
- 4 0 M=0003909/D^2 ! (Eq 2)
- 50 INPUT V! (=V1/V2)
- 60 GOSUB130
- 70 I F ABS(X1)>=.0001THEN 50
- 80 F=1/2/K*(K-1+2/M^2)*(1-V^2)+ (K+1)/2/K*LOG(V^2) ! (Eq 1) 90 D1=5 .7035/F
- 100 DISP D,D1
- 110 GOTO 3 0 (For antcher trial 1 of D if value it is not clo se enough to calculated D1)
- 120 END
- 130 Xl=-< 5439/V)+1+(K-1)/2/K*M^ 2*(1-1/V^2)
- 140 DISP XI
- 150 RETURN

particular flow pattern is subjective and all the pertinent variables apparently have not yet been correlated, boundaries between regions are fuzzy, as in (d).

It is to be expected that the kind of phase distribution will affect such phenomena as heat transfer and friction in pipelines. For the most part, however, these operations have not been correlated yet with flow patterns, and the majority of calculations of two-phase flow are made without reference to them. A partial exception is annular flow which tends to exist at high gas flow rates and has been studied in some detail from the point of view of friction and heat transfer.

The usual procedure for evaluating two-phase pressure drop is to combine pressure drops of individual phases in some way. To this end, multipliers ϕ_i are defined by

$$(\Delta P/L)_{\text{two-phase}} = \phi_i^2 (\Delta P/L)_i. \tag{6.94}$$

In the following table, subscript L refers to the liquid phase, G to the gas phase, and LO to the total flow but with properties of the liquid phase; x is the weight fraction of the vapor phase.

Subscript	Re	$\Delta P/L$	¢²
G	DGx/µ _G	$f_{c}G_{x}^{2}x^{2}/2g_{c}D\rho_{T}$	$(\Delta P/L)/(\Delta P/L)_{G}$
L	$DG(1-x)/\mu_1$	$f_{L}G^{2}(1-x)^{2}/2g_{c}D\rho_{L}$	$(\Delta P/L)/(\Delta P/L)_{L}$
LO	DG/μ_L	$f_{L0}G^2/2g_cD\rho_L$	$(\Delta P/L)/(\Delta P/L)_{L0}$

In view of the many other uncertainties of two phase flow correlations, the friction factors are adequately represented by

$$f = \begin{cases} 64/\text{Re}, \text{Re} < 2000, \text{Poiseuille equation}, \\ 0.32/\text{Re}^{0.25}, \text{Re} > 2000, \text{Blasius equation}. \end{cases}$$
 (6.95)

HOMOGENEOUS MODEL

The simplest way to compute line friction in two-phase flow is to adopt some kinds of mean properties of the mixtures and to employ the single phase friction equation. The main problem is the assignment of a two-phase viscosity. Of the number of definitions that have been proposed, that of McAdams et al. [Trans. ASME 64, 193-200 (1942)] is popular:

$$1/\mu_{\text{two-phase}} = x/\mu_G + (1-x)/\mu_L.$$
 (6.97)

The specific volumes are weight fraction additive,

$$\boldsymbol{V}_{\text{two-phase}} = \boldsymbol{x}\boldsymbol{V}_{\boldsymbol{G}} + (1 - \boldsymbol{x})\boldsymbol{V}_{\boldsymbol{L}}$$
(6.98)

so that

$$1/\rho_{\rm two-phase} = x/\rho_G + (1-x)/\rho_L,$$
 (6.99)

where χ is the weight fraction of the gas. Pressure drops by this method tend to be underestimated, but are more nearly accurate at higher pressures and higher flow rates.

With the Blasius equation (6.96), the friction factor and the pressure gradient become, with this model,

$$f = \frac{0.32}{(DG)^{0.25}} \left(\frac{x}{\mu_g} + \frac{1-x}{\mu_L}\right)^{0.25},$$
(6.100)

$$\frac{AP}{L} = \frac{fG^2}{2g_c D[x/\rho_G + (1-x)/\rho_L]}.$$
(6.101)

A particularly simple expression is obtained for the multiplier in terms of the Blasius equation:

$$\phi_{L0}^{2} = \frac{\Delta P/L}{(\Delta P/L)_{L0}} = \frac{1 - \chi + x\rho_{L}/\rho_{G}}{(1 - \chi + x\mu_{L}/\mu_{G})^{0.25}}.$$
(6.102)

Some values of $\phi_{I,0}^2$ from this equation for steam are

х	P = 0.669	bar	P = 10.3	ba
0. 01	3. 40		1. 10	
0. 10	12.16		1.95	
0. 50	80. 2		4.36	

High values of multipliers are not uncommon.

EXAMPLE 6.13

Isothermal Flow of a Nonideal Gas

The case of Example 6.12 will be solved with a van der Waals equation of steam. From the *CRC Handbook* of *Chemistry and Physics* (CRC Press, Boca Raton, FL, 1979),

a =
$$5.464 \text{ atm}(\text{m}^3/\text{kg mol})^2 = 1703.7 \text{ Pa}(\text{m}^3/\text{kg})^2$$
,
b = $0.03049 \text{ m}^3/\text{kg}$ mol = $0.001692 \text{ m}^3/\text{kg}$,
RT = $8314(493.2)/18.02 = 2.276(10^5) \text{ N m/kg}$.

Equation (6.93) becomes

$$\int_{0.0862}^{V_2} \left[\frac{-2.276(10^5)}{(V-0.00169)^2} + \frac{3407.4}{V^3} + (410.07)^2 \right] \frac{dV}{V} \\ + \frac{0.0187(410.07)^2(305)}{2(0.0777)} = 0,$$

$$\phi = \int_{0.0862}^{V_2} \left[\frac{-0.0369}{(V - 0.00169)^2} + \frac{5.52(10^{-4})}{V^3} + 0.0272 \right] \frac{dV}{V} + 1 = 0$$

The integration is performed with Simpson's rule with 20 intervals. Values of V_2 are assumed until one is found that makes $\phi = 0$. Then the pressure is found from the v dW equation:

$$P_2 = \frac{2.276(10^5)}{(V_2 - 0.00169)} \frac{1703.7}{V_2^2}$$

Two trials and, a linear interpolation are shown. The value $P_2 = 18.44$ bar compares with the ideal gas 17.13.

V ₂	¢	P2
0. 120	-0.0540	
0. 117	+0.0054	
0. 1173	0	18.44 bar



Figure 6.7. Flow patterns and correlations of flow regimes of liquid-gas mixtures in pipelines. (a) Patterns in horizontal liquid-gas flow. (b) Patterns in vertical liquid-gas flow. (c) Correlations of ranges of flow patterns according to Baker [Oil Gas J. 53(12), 185 (1954)], as replotted by Bell et al. [Chem. Eng. Prog. Symp. Ser. 66, 159 (1969)]; σ is surface tension of the liquid, and σ_w that of water. (d) Flow regimes of water/air at 25°C and 1 atm [Taitel and Dukler, AIChE J. 22, 47 (1976)]; the fuzzy boundaries are due to Mandhane et al. [Int. J. Two-Phase Flow 1, 537 (1974)].

SEPARATED FLOW MODELS

Pressure drop in two-phase flow is found in terms of pressure drops of the individual phases with empirical multipliers. The basic relation is

$$(\Delta P/L)_{\text{two-phase}} = \phi_G^2 (\Delta P/L)_G = \phi_L^2 (\Delta P/L)_L = \phi_{L0}^2 (\Delta P/L)_{L0}.$$
(6.103)

The last term is the pressure drop calculated on the assumption that the total mass flow has the properties of the liquid phase.

Some correlations of multipliers are listed in Table 6.8. Lockhart and Martinelli distinguish between the various combinations of turbulent and laminar (viscous) flows of the individual phases; in this work the transition Reynolds number is taken as 1000 instead of the usual 2000 or so because the phases are recognized to disturb each other. Item 1 of Table 6.8 is a guide to the applicability of the Lockhart-Martinelli method, which is the oldest, and two more recent methods. An indication of the attention that has been devoted to experimentation with two phase flow is the fact that Friedel (1979) based his correlation on some 25,000 data points.

Example 6.14 compares the homogeneous and Lockhart-Martinelli models for the flow of a mixture of oil and hydrogen.

OTHER ASPECTS

The pattern of annular flow tends to form at higher gas velocities; the substantial amount of work done on this topic is reviewed by

1. Recommendations			
μ, / μ _G	G (kg/m ² sec)	Correlation	
<1000	all	Friedel	
>1000	>100	Chisholm-Baroczy	
>1000	<100	Lockhart-Martinelli	

2. Lockhart-Martinelli Correlation



PAI	KAN	EI	EK	X

2	Liquid	Gas	Subscript	С
$\phi_L^* = 1 + C/X + 1/X^*$ $\phi_G^2 = 1 + CX + X^2$ $\chi^2 = (\Delta P/L)_L/(\Delta P/L)_G$	Turbulent Viscous Turbulent Viscous	t Turbuler Turbuler Viscous Viscous	nt tt nt Vt tv vv	20 12 10 5

Chisholm-Baroczy Correlation 3.

 $\phi_{L0}^{2} = 1 + (\gamma^{2} - 1)[Bx^{(2-n)/2}(1-x)^{(2-n)/2} + x^{2-n}] = (\Delta P/L)/(\Delta P/L)_{L0}$ $p_{L0} = 1 + (Y - 1)[Bx - (1 - x)]$ p = 0.25 $Y^{2} = (\Delta P/L)_{G0}/(\Delta P/L)_{L0}$ $B = 55/G^{0.5}, \quad 0 < Y < 9.5$ $= 520/YG^{0.5}, \quad 9.5 < Y < 28$ $= 15,000/Y^{2}G^{0.5}, \quad Y > 28$ x = weight fraction gas 4. Friedel Correlation
$$\begin{split} &\phi_{L0}^{2} = E + \frac{3.24FH}{Fr^{0.045}We^{0.0357}}, \quad Fr = G^{2}/g_{c}D\rho_{TP}^{2} \\ &E = (1-x)^{2} + x^{2}\frac{\rho_{L}f_{G0}}{\rho_{G}f_{L0}}, \quad We = G^{2}D/\rho_{TP}\sigma \\ &F = x^{0.78}(1-x)^{0.24}, \quad \rho_{TP} = \left(\frac{x}{\rho_{G}} + \frac{1-x}{\rho_{L}}\right)^{-1} \\ &H = \left(\frac{\rho_{L}}{\rho_{G}}\right)^{0.91} \left(\frac{\mu_{G}}{\mu_{L}}\right)^{0.19} \left(1 - \frac{\mu_{G}}{\mu_{L}}\right)^{0.7}, \quad X = \text{ weight fraction gas} \end{split}$$

1. (P.B. Whalley, cited by G.F. Hewitt, 1982). 2. [Lockhart and Martinelli, Chem. Eng. Prog. 45, 39-48 (1949); Chisholm, Int. J. Heat Mass Transfer 10, 1767-1778 (1967)]. 3. [Chisholm, Int. J. Heat Mass Transfer 16, 347-348 (1973); Baroczy, Chem. Eng. Prog. Symp. Ser. 62, 217-225 (1965)] 4. (Friedel, European Two Phase Flow Group Meeting, Ispra, Italy, Paper E2, 1979, cited by G.F. Hewitt, 1982).

EXAMPLE 6.14 Pressure Drop and Void Fraction in Liquid-Gas Flow A mixture of an oil and hydrogen at 500psia and 200°F enters a 3 in. Schedule 40 steel line. Data are:

Oil: 140,000 lb/hr, 51.85 lb/cuft, 2700 cfh, viscosity 15 cP. Hydrogen: 800 lb/hr, 0.142 lb/cuft, 5619 cfh, viscosity 2.5(10⁻⁷) lbf sec/sqft.

The pressure drop in 100 ft of line will be found, and also the voidage at the inlet condition.

$$\operatorname{Re}_{L} = \frac{4\dot{m}}{\pi Dg_{c}\mu} = \frac{4(140,000/3600)}{\pi (0.2557)(32.2)0.15},$$

$$\operatorname{Re}_{G} = \frac{4(800/3600)}{\pi (0.2557)(32.2)(2.5)(10^{-7})} = 137,500,$$

$$\frac{\varepsilon}{D} = 0.00059.$$

Round equations:

$$f = \frac{1.6434}{[\ln(0.135\varepsilon/D + 6.5/\text{Re}]^2]} = \begin{cases} 0.0272, & \text{liquid,} \\ 0.0204, & \text{gas,} \end{cases}$$
$$(\Delta P/L)_L = \frac{8f\dot{m}^2}{\pi^2 g_c \rho D^5} = \frac{8(0.0272)(38.89)^2}{\pi^2 (32.2)(51.85)(0.2557)^5}$$
$$= 18.27 \text{ psf/ft,}$$
$$(\Delta P/L)_G = \frac{8(0.0204)(0.222)^2}{\pi^2 (32.2)(0.142)(0.2557)^5} = 0.1663 \text{ psf/ft,}$$
$$X^2 = 18.27/0.1633 = 111.8.$$

Lockhart-Martinelli-Chisholm:

c = 20 for TT regime (Table 6.8),

$$\phi_L^2 = 1 + \frac{C}{X} + \frac{1}{X^2} = 2.90,$$

∴ ($\Delta P/L$) two phase = $\phi_L^2 (\Delta P/L)_L = 2.90(18.27)$
= 53.0 psf/ft, 36.8 psi/100 ft.

Check with the *homogeneous model*:

$$\begin{aligned} x &= \frac{800}{140\ 000 + 800} = 0.0057 \text{ wt fraction gas,} \\ \mu &= \left[\frac{0.0057}{2.5(10^{-7})} + \frac{0.9943}{3.13(10^{-4})}\right]^{-1} = 3.85(10^{-5})\frac{\text{lbf sec}}{\text{sqft}}, \\ \rho &= \left[\frac{0.0057}{0.142} + \frac{0.9943}{51.85}\right]^{-1} = 16.86 \text{ lb/cuft,} \end{aligned}$$

Hewitt (1982). A procedure for stratified flow is given by Cheremisinoff and Davis [*AIChE J. 25,* 1 (1979)].

Voidage of the holdup in the line is different from that given by the proportions of the incoming volumetric flows of the two phases, but is of course related to it. Lockhart and Martinelli's work indicates that the fractional gas volume is

 $\varepsilon = 1 - 1/\phi_L, \tag{6.104}$

where ϕ_L is defined in Table 6.8. This relation has been found to give high values. A correlation of Premoli et al. *[Termotecnica 25,*]

$$Re = \frac{4(39.11)}{\pi(32.2)(0.2557)3.85(10^{-5})} = 157,100$$

$$f = 0.0202,$$

$$\frac{\Delta P}{L} = \frac{8(0.0202)(39.11)^2}{\pi^2(32.2)(16.86)(0.2557)^5} = 42.2 \text{ psf/ft},$$

compared with 53.0 by the LMC method. Void fraction by Eq. (6.104):

$$\varepsilon_G = 1 - 1/\phi_L = 1 - 1/\sqrt{2.90} = 0.413,$$

compared with input flow condition of

$$\varepsilon = \frac{Q_G}{Q_G + Q_L} = \frac{5619}{5619 + 2700} = 0.675.$$

Method of Premoli [Eqs. (6.105) and (6.106)]: Surface tension $\sigma = 20$ dyn/cm, 0.00137 lbf/ft,

$$We = \frac{DG^2}{g_c \rho_L \sigma} = \frac{16\dot{m}^2}{\pi^2 g_c D^3 \rho_L \sigma}$$

= $\frac{16(38.89)^2}{\pi^2 (32.2)(0.2557)^3 (51.85)(0.00137)} = ^{64,118},$
Re = 19,196,
 $E_1 = 1.578(19196)^{-0.19} (51.85/0.142)^{0.22} = 0.8872,$
 $E_2 = 0.0273(6411.8)(19196)^{-0.51} (51.85/0.142)^{-0.08} = 7.140.$
 $y = 5619/2700 = 2.081,$
 $yE_2 = 2.081(7.140) = 14.86.$

Clearly, this term must be less than unity if Eq. (6.105a) for S is to be valid, so that equation is not applicable to this problem as it stands. If yE_2 is replaced by $y/E_2 = 0.2914$, then

$$S = 1 + 0.8872 \left(\frac{2.081}{1.2914} - 0.2914\right)^{0.5} = 2.02,$$

and the voidage is

$$\varepsilon = \frac{5619}{5619 + 2.02(2700)} = 0.51,$$

which is a plausible result. However, Eqs. (6.105) and (6.105a) are quoted correctly from the original paper; no numerical examples are given there.

17-26 (1971); cited by Hewitt, **1982]** gives the void fraction in terms of the incoming volumetric flow rates by the equation

$$\boldsymbol{\varepsilon}_{\boldsymbol{G}} = \boldsymbol{Q}_{\boldsymbol{G}} / (\boldsymbol{Q}_{\boldsymbol{G}} + \boldsymbol{S} \boldsymbol{Q}_{\boldsymbol{L}}), \tag{6.105}$$

where S is given by the series of equations

$$S = 1 + E_1[y/(1 + yE_2) - yE_2]^{1/2},$$

$$E_1 = 1.578 \text{ Re}^{-0.19} (\rho_L/\rho_G)^{0.22},$$
(6.105')

$$E_2 = 0.0273 \text{ We } \operatorname{Re}^{-0.51}(\rho_L/\rho_G)^{-0.08},$$

 $\chi = Q_G/Q_L, \quad \operatorname{Re} = DG/\mu_L, \quad \operatorname{We} = DG^2/\sigma\rho_L.$

Direct application of these equations in Example 6.14 is not successful, but if E_2 is taken as the reciprocal of the given expression, a plausible result is obtained.

6.9. GRANULAR AND PACKED BEDS

Flow through granular and packed beds occurs in reactors with solid catalysts, adsorbers, ion exchangers, filters, and mass transfer equipment. The particles may be more or less rounded or may be shaped into rings, saddles, or other structures that provide a desirable ratio of surface and void volume.

Natural porous media may be consolidated (solids with holes in them), or they may consist of unconsolidated, discrete particles. Passages through the beds may be characterized by the properties of porosity, permeability, tortuosity, and connectivity. The flow of underground water and the production of natural gas and crude oil, for example, are affected by these characteristics. The theory and properties of such structures is described, for instance, in the book of Dullien (*Porous Media, Fluid Transport and Pore Structure*, Academic, New York, 1979). A few examples of porosity and permeability are in Table 6.9. Permeability is the proportionality constant k in the flow equation $u = (k/\mu) dP/dL$.

Although consolidated porous media are of importance in chemical engineering, only unconsolidated porous media are incorporated in process equipment, so that further attention will be restricted to them.

Granular beds may consist of mixtures of particles of several sizes. In flow problems, the mean surface diameter is the appropriate mean, given in terms of the weight fraction distribution, x_i , by

$$D_p = 1/(\sum x_i/D_i). \tag{6.106}$$

When a particle is not spherical, its characteristic diameter is taken as that of a sphere with the same volume, so that

$$D_p = (6V_p/\pi)^{1/3}.$$
 (6.107)

SINGLE PHASE FLUIDS

Extensive measurements of flow in and other properties of beds of particles of various shapes, sizes and compositions are reported by

Porosity Permeability Media (%) (cm²) $1.3 \times 10^{-3} - 3.9 \times 10^{-3}$ Ber saddles 68-83 $3.8 \times 10^{-5} - 1.0 \times 10^{-5}$ Wire crimps 68-76 $4.9 \times 10^{-10} - 1.2 \times 10^{-9}$ Black slate powder 57-66 $1.3 \times 10^{-10} - 5.1 \times 10^{-10}$ Silica powder 37-49 $2.0 \times 10^{-7} - 1.8 \times 10^{-7}$ Sand (loose beds) 37-50 $2.0 \times 10^{-9} - 1.4 \times 10^{-7}$ $2.9 \times 10^{-12} - 3.0 \times 10^{-8}$ $5.0 \times 10^{-12} - 3.0 \times 10^{-8}$ Soil 43-54 Sandstone (oil sand) 8-38 $2.0 \times 10^{-11} - 4.5 \times 10^{-11}$ Limestone, dolomite 4-10 4.8 x 10⁻¹¹-2.2 x 10 Brick 12-34 $1.0 \times 10^{-9} - 2.3 \times 10^{-7}$ 9.5 x 10⁻¹⁰ - 1.2 x 10⁻¹⁰ Concrete 2-7 Leather 56-59 $3.3 \times 10^{-6} - 1.5 \times 10^{-6}$ Cork board $8.3 \times 10^{-6} - 1.2 \times 10^{-6}$ -5 Hair felt _ -7 Fiberglass 88-93 2.4 x 10 -5.1×10 -5 Cigarette filters 17-49 1.1 x 10 -10 <u>-4</u>.4 × 10⁻⁹ 2.0 x 10 Agar-agar

TABLE 6.9. Porosity and Permeability of Several Unconsolidated and Consolidated Porous Media



Leva et al. (1951). Differences in voidage are pronounced as Figure 6.8(c) shows.

A long-established correlation of the friction factor is that of Ergun (*Chem. Eng. Prog.* **48**, 89-94, 1952). The average deviation from his line is said to be $\pm 20\%$. The friction factor is

$$f_p = \frac{g_c D_p \varepsilon^3}{u^2 (1 - \varepsilon)} \left(\frac{\Delta P}{L}\right)$$
(6.108)

$$= 150/\text{Re}_p + 1.75$$
 (6.109)

with

$$\operatorname{Re}_{p} = D_{p}G/\mu(1-E).$$
 (6.110)



Figure 6.8. Friction factors and void fractions in flow of single phase fluids in granular beds. (a) Correlation of the friction factor, Re = $D_p G/(1-\varepsilon)\mu$ and $f_p = [g_c D_p \varepsilon^3/\rho u^2(1-\varepsilon)](\Delta P/L = 150/\text{Re} + 4.2/(\text{Re})^{1/6}$ [Sato et al., J. Chem. Eng. Jpn. 6, 147-152 (1973)]. (b) Void fraction in granular beds as a function of the ratio of particle and tube diameters [Leva, Weintraub, Grummer, Pollchik, and Storch, U.S. Bur. Mines Bull. 504 (1951)].

The pressure gradient accordingly is given by

$$\frac{AP}{L} = \frac{G^2(1-\varepsilon)}{\rho g_c D_p \varepsilon^3} \left[\frac{150(1-\varepsilon)\mu}{D_P G} + 1.75 \right]$$
(6.11)

For example, when $D_p = 0.005 \text{ m}$, $G = 50 \text{ kg/m}^2 \text{ sec}$, $g_c = 1 \text{ kgm/N sec}^2$, $\rho = 800 \text{ kg/m}^3$, $\mu = 0.010 \text{ N sec/m}^2$, and $\varepsilon = 0.4$, the gradient is $\Delta P/L = 0.31(10^5)$ Pa/m.

An improved correlation is that of Sato (1973) and Tallmadge (*AIChE J.* 16, 1092 (1970)] shown on Figure 6.8(a). The friction factor is

$$f_p = 150/\text{Re}_p + 4.2/\text{Re}_p^{1/6} \tag{6.112}$$

with the definitions of Eqs. (6.108) and (6.110). A comparison of Eqs. (6.109) and (6.112) is

Re _p	5	50	500	5000
, (Ergun)	31.8	4.80	2.05	1.78
, (Sato)	33. 2	5.19	1.79	1.05

In the highly turbulent range the disagreement is substantial.

TWO-PHASE FLOW

Operation of packed trickle-bed catalytic reactors is with liquid and gas flow downward together, and of packed mass transfer equipment with gas flow upward and liquid flow down.

Concurrent flow of liquid and gas can be simulated by the homogeneous model of Section 6.8.1 and Eqs. 6.109 or 6.112, but several adequate correlations of separated flows in terms of Lockhart-Martinelli parameters of pipeline flow type are available. A number of them is cited by Shah *(Gas-Liquid-Solid Reactor Design, McGraw-Hill, New York, 1979, p. 184).* The correlation of Sato (1973) is shown on Figure 6.9 and is represented by either

$$\phi = (\Delta P_{LG} / \Delta P_L)^{0.5} = 1.30 + 1.85(X)^{-0.85}, \quad 0.1 < X < 20,$$
(6.113)

or

$$\log_{10}\left(\frac{\Delta P_{LG}}{\Delta P_L + \Delta P_G}\right) = \frac{0.70}{\left[\log_{10}(X/1.2)\right]^2 + 1.00},$$
 (6.114)

where

$$X = \sqrt{(\Delta P/L)_L / (\Delta P/L)_G}$$
(6.115)

The pressure gradients for the liquid and vapor phases are calculated on the assumption of their individual flows through the bed, with the correlations of Eqs. (6.108)-(6.112).

The fraction h_L of the void space occupied by liquid also is of interest. In Sato's work this is given by

$$h_L = 0.40(a_s)^{1/3} X^{0.22}, \tag{6.116}$$

where the specific surface is

$$a_{r} = 6(1-\varepsilon)/D_{p}. \tag{6.117}$$

Additional data are included in the friction correlation of Specchia and Baldi [*Chem. Eng. Sci.* 32, 515-523 (1977)], which is represented by

$$f_{LG} = \frac{g_c D_p \varepsilon}{3\rho_G u_G^2 (1-\varepsilon)} \left(\frac{\Delta P}{L}\right),\tag{6.118}$$



Figure 6.9. Pressure drop gradient and liquid holdup in liquid-gas concurrent flow in granular beds. *[Sato, Hirose, Takahashi, and Toda, J.* Chem. Eng. Jpn. 6, 147-152 (1973)]. (a) Correlation of the two phase pressure drop gradient $\Delta P/L$, $\phi = 1.30 + 1.85X^{-0.85}$. (b) Correlation of frictional holdup h_L of liquid in the bed; a_s is the specific surface, *l/mm, d* is particle diameter, and *D* is tube diameter. $h_L = 0.4a_s^{1/3}X^{0.22}$.

$$\ln f_{LG} = 7.82 - 1.30 \ln(Z/\psi^{1.1}) - 0.0573[\ln(Z/\psi^{1.1})]^2. \quad (6.119)$$

The parameters in Eq. (6.119) are

$$Z = (Re_G)^{1.167} / (Re_L)^{0.767}, (6.120)$$

$$\psi = \frac{\sigma_w}{\sigma_L} \left[\frac{\mu_L}{\mu_w} \left(\frac{\rho_w}{\rho_I} \right)^2 \right]^{1/3}.$$
(6.121)

Liquid holdup was correlated in this work for both nonfoaming and foaming liquids.

Nonfoaming,
$$h_L = 0.\ 125(Z/\psi^{1.1})^{-0.312} (a_s D_p/\varepsilon)^{0.65}$$
, (6.122)
Foaming, $h_I = 0.06(Z/\psi^{1.1})^{-0.172} (a_s D_p/\varepsilon)^{0.65}$. (6.123)

The subscript w in Eq. (6.121) refers to water.

Countercurrent flow data in towers with shaped packings are represented by Figure 13.37. The pressure drop depends on the viscosity of the liquid and on the flow rates and densities of the liquid and gas, as well as on characteristics of the packing which are represented here by the packing factor F. Nominally, the packing factor is a function of the specific surface a, and the voidage ε , as

$$F = a_s / \varepsilon^3, \tag{6.124}$$

but calculated values are lower than the experimental values shown in the table by factors of 2-5 or so. Clearly the liquid holdup reduces the effective voidage to different extents with different packings. The voidages of the packings in the table range from 70 to **95%**, whereas voidages obtained with small spherical or cylindrical packings normally used as catalysts are less than 40% or so, which makes them impractical for countercurrent operation. However, catalysts are made in the forms of rings or saddles when very low pressure drop or countercurrent operation is desirable.

Even when they are nominally the same type and size, packings made by different manufacturers may differ substantially in their pressure drop and mass transfer behavior, so that manufacturers data should be obtained for final design.

Many data on individual packings are given by Billet (*Distillation Engineering*, Chemical Pub. Co., New York), in *Chemical Engineers Handbook* (McGraw-Hill, New York, 1984, p. 18.23) and with Figure 13.37.

The uppermost line of Figure 13.37(a) marks the onset of flooding which is the point at which sharp increase of pressure drop obtains on a plot against liquid rate. Flooding limits also are represented on Figure 13.36; in practice, it is customary to operate at a gas rate that is 70% of that given by the line, although there are many data points below this limit in this correlation.

Mesh or other open structures as vessel packing have attractive pressure drop and other characteristics, but each type has quite individual behavior so that it is best to consult their manufacturer's data.

6.10. GAS-SOLID TRANSFER

Equipment for pneumatic conveying is described in Section 5.2 along with some rules for calculating power requirements. Here the latter topic will be supplemented from a more fundamental point of view.

CHOKING VELOCITY

Although the phenomena are not clearcut, partial settling out of solids from the gas stream and other instabilities may develop below certain linear velocities of the gas called choking velocities. Normal pneumatic transport of solids accordingly is conducted above such a calculated rate by a factor of 2 or more because the best correlations are not more accurate. Above choking velocities the process is called dilute phase transport and, below, dense phase transport.

What appears to be the best correlation of choking velocities is due to Yang [AIChE J. 21, 1013-1015 (1975)], supplemented by Punwani et al. and Yang (cited by Teo and Leung, 1984, pp. 520-521). The choking velocity U_{gc} and voidage ε_c are found by simultaneous solution of the equations

$$G_s/\rho_s = (U_{gc} - U_t)(1 - \varepsilon_c) \tag{6.125}$$

or

$$\varepsilon_c = 1 \quad G_s / \rho_s (U_{gc} \quad U_t) \tag{6.126}$$

and

$$gD(\varepsilon_c^{-4.7} - 1) = 3.41(10^5)(\rho_g/\rho_s)^{2.2}(U_{gc} - U_t)^2, \qquad (6.127)$$

where G_s is the mass rate of flow of solid per unit cross section and the other terms are defined in Table 6.10. When ε_c from Eq. (6.126) is substituted into Eq. (6.127), the single unknown in that equation is readily found with a root solving routine. For the case of Example 6.15, $G_s = 29.6 \text{ kg/m}^2 \sec$, $U_t = 0.45 \text{ m/sec}$, $\rho_s = 1282 \text{ kg/m}^3$, and $\rho_g = 1.14 \text{ kg/m}^3$. Accordingly, $U_{gc} = 1.215 \text{ m/sec}$ and $\varepsilon_c = 0.9698$.

TABLE 6.10. Equations for the Calculation of Pressure Drop in Gas-Solid Transport

Solid Friction Factor $f_{\!\!s}$ According to Various Investigators	
Investigator f _s	
Stemerding (1962) 0.003	(1)
Reddy and Pei (1969) $0.046U_{p}^{-1}$	(2)
Van Swaaij, Buurman, and V van Breugel (1970) $0.080 U_p^{-1}$	(3)
Capes and Nakamura (1973) $0.048U_{p}^{-1.22}$	(4)
Konno and Saito (1969) $0.0285\sqrt{gD} U_p^{-1}$	(5)
Yang (1978). vertical $0.00315 \frac{1-\varepsilon}{\varepsilon^3} \left[\frac{(1-\varepsilon)U_t}{U_f - U_p} \right]^{-0.979}$	(6)
Yang (1976). horizontal $0.0293 \frac{1-\varepsilon}{\varepsilon^3} \left[\frac{(1-\varepsilon)U_f}{\sqrt{gD}} \right]^{-1.15}$	(7)

Free Setting Velocity

$$K = D_{p} \left[\frac{g p_{f}(\rho_{p} - \rho_{f})}{\mu_{f}^{2}} \right]^{1/3}$$

$$(8)$$

$$g D_{-}^{2}(\rho_{r} - \rho_{f})$$

$$U_{t(\text{Stokes})} = \frac{g D_p (\mu_p - \mu_f)}{18 \mu_f}, \quad K < 3.3 \qquad (9)$$

0.153 g^{0.71} D₁^{1.14} (\rho_p - \rho_s)^{0.71}

$$U_{t(\text{intermediate})} = \frac{0.153g}{\rho_{p}} \frac{D_{p}}{\rho_{f}} \frac{(p_{p} - \rho_{f})}{\mu_{f}}, \quad 3.3 < K < 43.6 \quad (10)$$

$$U_{t(Newton)} = 1.75 \left(\frac{\rho_{P} (\rho_{P} - \rho_{f})}{\rho_{f}}\right) , \qquad 43.6 < \kappa < 2360$$
(11)

	Particle Velocity	
Investigator	U_p	
Hinkle (1953) IGT (1978)	$U_{g} = U, U_{g}(1 - 0.68D_{p}^{0.92}\rho_{p}^{0.5}\rho_{f}^{-0.2}D^{-0.54})$	(12) (13)
Yang (1976)	$U_g - U_t \left[\left(1 + \frac{f_s U_p^2}{2gD} \right)^{e^{q_s/2}} \right]^{1/2}$	(14)

Voidage			
$\varepsilon = 1 - 4\dot{m}_p / \pi D^2 (\rho_p - \rho_f) U_p$	(15)		

Notation: U, is a fluid velocity, U_p is particle velocity, U_t is particle free settling velocity, \dot{m}_s is mass rate of flow of solid, D = pipe diameter, D_p is particle diameter, g = 9.806 m/sec at sea level.

(Klinzing, Gas-So/id Transport, McGraw-Hill, New York, 1981).

PRESSURE DROP

The relatively sparse data on dense phase transport is described by Klinzing (1981) and Teo and Leung (1984). Here only the more important category of dilute phase transport will be treated.

The pressure drop in simultaneous flow of gas and solid particles is made up of contributions from each of the phases. When the particles do not interact significantly, as in dilute transport, the overall pressure drop is represented by

$$AP = \rho_p (1 - \varepsilon) Lg + \rho_f \varepsilon Lg + \frac{2f_g \rho_f U_f^2 L}{D} + \frac{2f_s \rho_p (1 - \varepsilon) U_p^2 L}{D}$$
(6.128)

for vertical transport; in horizontal transport only the two frictional terms will be present. The friction factor f_g for gas flow is the normal one for pipe flow; except for a factor of 4, it is given by Eq. (6.19) for laminar flow and by the Round equation (6.21) for turbulent flow. For the solid friction factor f_s , many equations of

EXAMPLE 6.15

Pressure Drop in Flow of Nitrogen and Powdered Coal Powdered coal of 100 μ m dia and 1.28 specific gravity is transported vertically through a 1-in. smooth line at the rate of 15 g/sec. The carrying gas is nitrogen at 1 atm and 25°C at a linear velocity of 6.1 m/sec. The density of the gas is 1.14 kg/m³ and its viscosity is 1.7(10⁻⁵) N sec/m². The equations of Table 6.10 will be used for the various parameters and ultimately the pressure gradient $\Delta P/L$ will be found:

Eq. (8),
$$K = 10^{-4} \left\{ \frac{9.806(1.14)(1282 - 1.14)}{[1.7(10^{-5})]^2} \right\}^{1/3} = 3.67$$

Eq. (10), $U_r = \frac{0.153(9.806)^{0.71}(0.0001)^{1.14}(1282 - 1.14)^{0.71}}{1.14^{0.29}[1.7(10^{-5})]^{0.43}}$
= 0.37 m/sec (0.41 m/sec by Stokes' law),
Eq. (15) = 1 0.015 4.00231

Eq. (15),
$$\varepsilon = 1 - \frac{0.013}{(\pi/4)(0.0254)^2(1282 - 1.14)U_p} = 1 - \frac{0.0251}{U_p}$$
, (I)

Eq. (14),
$$U_p = 6.1 - 0.45\sqrt{1 + f_s U_p^2/2(9.806)(0.0254)}$$

Eq. (7),
$$f_s = \frac{0.00315(1-\varepsilon)\left[(1-\varepsilon)0.45\right]^{-0.979}}{\varepsilon^3}$$
 (III)

Equations (I), (II), and (III) are solved simultaneously with the results:

 $\varepsilon = 0.9959$ and $U_p = 5.608$,

For the calculation of the pressure drop,

$$f_s = 0.0031$$
 (Yang equation),
 $\operatorname{Re}_f 390 \frac{DU_f \rho_f}{\mu_f} - \frac{0.0254(6.1)(1.14)}{1.7(10^{-5})} = 10$

Therefore, Round's Eq. (6.21) applies:

$$f_f = \frac{1}{4} f_{\text{Round}} = 0.0076,$$

varying complexity have been proposed, of which some important ones are listed in Table 6.10.

These equations involve the free settling velocity U_{i} , for which separate equations also are shown in the table. At lower velocities Stokes' law applies, but corrections must be made at higher ones. The particle velocity U_p is related to other quantities by Eqs. (12)-(14) of the table, and the voidage in turn is represented by Eq. (15). In a review of about 20 correlations, Modi et al. (Proceedings, Powder and Bulk Solids Handling and Processing Conference, Powder Advisory Center, Chicago, 1978, cited by Klinzing, 1981) concluded that the correlations of Konno and Saito (1969) and of Yang (1976, 1978) gave adequate representation of pneumatic conveying of coal. They are applied in Example 6.15 and give similar results there.

6.11. FLUIDIZATION OF BEDS OF PARTICLES WITH GASES

As the flow of fluid through a bed of solid particles increases, it eventually reaches a condition at which the particles are lifted out of permanent contact with each other. The onset of that condition is Eq. (6.128),

$$\Delta P/L = 9.806[1282(1 - 0.9959) + 1.14(0.9959)] + (2/0.0254)[0.0076(1.14)(6.1)^2 + 0.0031(1282)(0.0041)(5.608)^2] = 51.54 + 11.13 + 25.38 + 40.35 = 128.4 Pa/m.$$

With Eqs. (5) and (13), no trial calculations are needed.

Therefore, the solid frictional gradient is obtained from the calculated value 40.35 in the ratio of the friction factors.

 $(\Delta P/L)_{\text{solid friction}} = 40.35(0.00242/0.0031) = 31.5 \text{ Pa/m}.$

```
10 ! Example 6.15. Pressure dro
    pin flow of nitrogen and po
    wdered coal
 20 INPUT U
 30 E=1-.0231/U ! (Eq I)
40 F=.003151*(1-E)/E^3*(.45*(1-
    E)/(6.1-U))^-.973 ! (Eq III)
 50 G=-U+6.1-.45*(1+2.007*F*U^2)
    Ā.5!
            (should =
                         0)
 60 PRINT "U="; U
 70 PRINT "G=";G
           20 ! (if G is not
 80 GOTO -
                                 suffi
    cientlr close to zero)
 30 END
u = 5.608
G = -000059348061
```

called minimum fluidization. Beyond this point the solid-fluid mass exhibits flow characteristics of ordinary fluids such as definite viscosity and flow through lines under the influence of hydrostatic head difference. The rapid movement of particles at immersed surfaces results in improved rates of heat transfer. Moreover, although heat transfer rate between particles and fluid is only moderate, 1-4 Btu/(hr)(sqft)(°F), the amount of surface is so great, 10,000–150,000 sqft/cuft, that temperature equilibration between phases is attained within a distance of a few particle diameters. Uniformity of temperature, rapid mass transfer, and rapid mixing of solids account for the great utility of fluidized beds in process applications.

As the gas flow rate increases beyond that at minimum fluidization, the bed may continue to expand and remain homogeneous for a time. At a fairly definite velocity, however, bubbles begin to form. Further increases in flow rate distribute themselves between the dense and bubble phases in some ways that are not well correlated. Extensive bubbling is undesirable when intimate contacting between phases is desired, as in drying processes or solid catalytic reactions. In order to permit bubble formation, the



Figure 6.10. Characteristics of gas-solid fluidization. (a) Schematic of the progress of pressure drop and bed height with increasing velocity, for "normal" and "abnormal" behavior. For normal systems, the rates at minimum fluidization and minimum bubbling are the same. (b) Behavior of heat transfer coefficient with gas flow rate analogous to part (a). The peak depends on the density and diameter of the particles (*Botteril*, Fluid Bed Heat Transfer, *Academic*, New York, 1975). (c) Bed expansion ratio as a function of reduced flow rate and particle size. The dashed line is recommended for narrow size range mixtures (*Leva*, 1959, *p*. 102). (d) Correlation of fluctuations in level, the ratio of the maximum level of disturbed surface to average level (*Leva*, 1959, *p*. 105). (e) Bed voidage at minimum fluidization (*Leua*, 1959). Agarwal and Storrow: (a) soft brick; (b) absorption carbon; (c) broken Raschig rings; (d) coal and glass powder; (e) Carborundum; (f) sand U.S. Bureau of Mines: (g) round sand, $\phi_s = 0.86$; (h) sharp sand, $\phi_s = 0.67$; (i) Fischer-Tropsch catalyst, $\phi_s = 0.58$; (j) anthracite coal, $\phi_s = 0.63$; (k) mixed round sand, $\phi_s = 0.86$. Van Heerden et al.: (I) coke; (m) Carborundum. (f) Coefficient C in the equation for mass flow rate at minimum fluidization (*Leva*, 1959): $G_{mf} = CD_{pg}^2 g_c \rho_F (\rho_s - \rho_F)/\mu$ and C = 0.0007 Re⁻⁶⁶³(g)-Mittimum bubbling and fluidization velocities of cracking catalysts (*Harriott and Simone, in* Cheremisinoff and Gupta, Eds., Handbook of Fluids in Motion, *Ann Arbor Science, Ann Arbor, MI*, 1983, p. 656). (h) Minimum fluidization and bubbling velocities with air as function of vessel diameter and superficial linear velocity [*Zenz and Weil*, AIChE J. 4, 472 (1958)]. (j) Good fluidization conditions (*W.V. Battcock and K.K. Pillai, "Particle size in Pressurised Combustors,"* Proc. Fifth International Conference on Fluidised Bed Combustion, Mitre Corp., Washington D.C., 1977).



particles appear to interlock to form a skin around the bubble and thus prevent free particles from raining through those spaces. Bubble sizes become large at high rates of flow and may eventually reach the diameter of the vessel, at which time slugging and severe entrainment will occur.

Onset of fluidization commonly is detected by noting a break in the plot of flow against pressure drop. For a range beyond the minimum fluidizing velocity, the pressure drop remains constant and equal to the weight of the bed but the bed level rises gradually and bubbles are generated at an increasing rate. Not in all cases, however, is the fluidization behavior entirely smooth. Figure 6.10(a) compares "normal" with a case of "abnormal" behavior. Among the reasons for abnormality are aggregation of particles because of stickiness or attractive forces between small particles and interlocking of rough surfaces. It is even possible for bubbling to occur before the onset of fluidization by formation of channels in the bed.

CHARACTERISTICS OF FLUIDIZATION

Six different regimes of fluidization are identified in Figure 6.11 and its legend. Particulate fluidization, class (b) of the figure, is desirable for most processing since it affords intimate contacting of phases. Fluidization depends primarily on the sizes and densities of the particles, but also on their roughness and the temperature, pressure, and humidity of the gas. Especially small particles are subject to electrostatic and interparticle forces.

Four main classes characterized by diameters and differences in densities of the phases are identified in Figure 6.12 and its legend. Groups A and B are most frequently encountered; the boundary between them is defined by the equation given in the legend. Group A particles are relatively small, 30-150 μm dia, with densities below 1.5 g/cc. Their bed behavior is "abnormal" in that the bed expands appreciably before bubbling sets in, and the minimum bubbling velocity always is greater than the minimum fluidization velocity. The bubbles disengage quickly. Cracking catalysts that have been studied extensively for their fluidization behavior are in this class. Group B materials have $d_p = 150-500 \,\mu\text{m}$ and are $1.5-4.0 \,\text{g/mL}$. The bed expansion is small, and minimum bubbling and fluidization velocities are nearly the same. The bubbles also disengage rapidly. Coarse sand and glass beads that have been favorite study materials fall in this group. Group C comprises small cohesive particles whose behavior is influenced by electrostatic and van der Waals forces. Their beds are difficult to fluidize and subject to channelling. Group D particles are large, 1 mm or more, such as lead shot and grains. They do not fluidize well and are usually handled in spouted beds, such as Figure 9.13(f).

Among the properties of particles most conducive to smooth fluidization are the following:

- 1. rounded and smooth shape,
- 2. in the range of 50-500 μ m diameter,
- 3. a broad spectrum of particle sizes, with ratios of largest to smallest sizes in the range of 10 to 25,
- 4. enough toughness to resist attrition.

Such tailoring of properties is feasible for many catalyst-carrier formulations, but drying processes, for instance, may be restricted by other considerations. Fluidization of difficult materials can be maintained by mechanical or ultrasonic vibration of the vessel, or pulsation of the supply of the fluid, or mechanical agitation of the contents of the vessel, or by addition of fluidization aids such as fine foreign solids.



Figure 6.11. Six regimes of fluidization identified with increasing gas superficial velocity (Grace, 198.2).

	Velocity Range	Regime	Appearance and Principal Features
(a)	0 ≤ <i>u</i> < <i>u_{mf}</i>	fixed bed	particles are quiescent; gas flows through interstices
(b)	u _{mf} ≤ u < u _{mb}	_P articulate fluid- ization	bed expands smoothly in a homoge- neous manner; top surface is well defined; some small-scale particle motion; little tendency for particles to angregate; very little fluctuation
(C)	u _{mb} ≤ U < U _{rr}	ns bubbling fluid- ization	void regions form near the distributor, grow mostly by coalescence, and rise to the surface; top surface is well defined with bubbles breaking through periodically; irregular pres- sure fluctuations of appreciable amplitude
(d)	$u_{ms} \leq u < u_k$	slugging regime	voids fill most of the column cross section; top surface rises and col- lapses with reasonably regular fre- quency; large and regular pressure fluctuations
(e)	$u_k \leq u < u_{tr}$	turbulent regime	small voids and particle clusters dart to and fro; top surface difficult to distinguish; small-amplitude pressure fluctuations only
(f)	U _{tr} ≤ U	fast fluid- ization	no upper surface to bed; particles are transported out the top and must be replaced by adding solids at or near the bottom; clusters or strands of particles move downward, mostly near the wall, while gas, containing widely dispersed particles, moves upward; at fixed solid feed rate, increasingly dilute as <i>U</i> is increased

SIZING EQUIPMENT

Various aspects of the hydrodynamics of gas-solid fluidization have been studied extensively with conclusions that afford guidance to the interpretation and extension of pilot plant data. Some of the leading results bearing on the sizing of vessels will be discussed here. Heat transfer performance is covered in Chapter 17. Example 6.16 applies to some of the cited data.



Figure 6.12. Characteristics of four kinds of groups of particles classified by **Geldart** [Powder Technol. 6, 201–205 (1972); 7, 285-292 (1973)]. The boundary between A and B is represented by the equation $d_S = 44,000\rho_F^{0.1}\mu_F^{0.9}/g(\rho_S - \rho_F)$ and that between B and D by $(\rho_S - \rho_F) d_S^2 = 10^-$ kg/m.

Feature	Group C	Group A	Group B	Group D
Distinguishing word or phrase	Cohesive	aeratable	bubble readilv	spoutable
Example	Flour	fluid cracking catalyst	sand	wheat
Particle size for $\rho_{\rm c} = 2.5 {\rm g/cm}^3$	≤20 µm	$20 < \overline{d}_{S} \le 90 \ \mu m$	$90 < \overline{d}_S \le 650 \ \mu m$	>650 µm
Channeling	Severe	little	negligible	negligible
Spouting	None	none	shallow beds only	readily
Collapse rate	-	slow	rapid	rapid
Expansion	Low because of channeling	high; initially bubble-free	m e d i u m	medium
Bubble shape	channels, no bubbles	flat base, spherical cap	rounded with small indentation	rounded
Rheological character of dense phase	high yield stress	apparent viscosity of order 1 poise	apparent viscosity of order 5 poise	apparent viscosity of order 10 poise
Solids mixing	very low	high	medium	low
Gas back mixing	very low	high	medium	low
Slugging mode	flat raining plugs	axisymmetric	mostly axi- symmetric	mostly wall slugs
Effect of $ar{d}_S$ (within group) on hydrodynamics	unknown	appreciable	minor	unknown
Effect of particle size distribution	unknown	appreciable	negligible	can cause segregation

Solids of practical interest often are mixtures of a range of particle diameters, but, for convenience, correlations are expressed in terms of a single size which is almost invariably taken as the surface average diameter given by

$$d_p = 1/\sum x_i d_i, \tag{6.129}$$

where x_i is the weight fraction of the material having a diameter d_i measured by screen analysis. Particles that deviate substantially from a spherical shape are characterized as having the diameter of a

sphere with the same volume as the particle. The sphericity is defined as the ratio

$$\phi$$
 = (surface of a sphere)/(surface of the particle with the same volume) (6.130)

and is always less than unity. Accordingly, the relation between the effective particle size d_p and that found by screen analysis is

$$d_p = \phi d_{\text{screen}}.\tag{6.131}$$

EXAMPLE 6.16

Dimensions of **a** Fluidiied Bed Vessel

A fluidized bed is to hold 10,000 kg of a mixture of particles whose true density is 1700 kg/m³. The fluidizing gas is at 0.3 m³/sec, has a viscosity of 0.017 cP or 1.7(E - 5) N sec/m² and a density of 1.2 kg/m³. The size distribution of the particles is

 d (μm)
 252
 178
 126
 89
 70
 50
 30
 10

 x (wt fraction)
 0.088
 0.178
 0.293
 0.194
 0.113
 0.078
 0.042
 0.014

 u_t (m/sec)
 3.45
 1.72
 0.86
 0.43
 0.27
 0.14
 0.049
 0.0054

The terminal velocities are found with Stokes' equation

$$u_t = \frac{g(\rho_p - \rho)}{18\mu} d_p^2 = \frac{9.81(1700 - 1.2)(E - 12)}{18[1.7(E - 5)]} [d_p \text{ (pm)}]^{\circ}.$$

(a) The average particle size is

$$d_p = 1 / \sum (x_i/d_i) = 84.5$$
 ym.

(b) With $d_p = 84.5$ and density difference of 1699 kg/m³, the material appears to be in Group A of Figure 6.12.

(c) Minimum fluidization velocity with Eq. (6.133)

$$u_{mf} = \frac{0.0093[84.5(E-6)]^{1.82}(1700-1.2)^{0.94}}{[1.7(E-5)]^{0.88}(1.2)^{0.06}}$$

= 0.0061 m/sec,

and with Eqs. (6.134) and (6.135),

$$Ar = \frac{1.2(1700 - 1, 2)(9.81)[84.5(E - 6)]^3}{[1.7(E - 5)]^2} = 4175,$$

R e, = $\sqrt{(27.2)^2 + 0.0408(41.75)} = 27.2 = 0.0313,$

$$u_{mb} = \frac{\mu}{d_p \rho} \frac{\text{Re}_{mf}}{84.5(E - 6)(1.2)} = 0.0052 \text{ m/sec.}$$

Use the larger value, $u_{mf} = 0.0061$, as the conservative one. (d) Minimum bubbling velocity, with Eq. (6.136)

 $u_{mb} = 33(84.5)(E-6)[1.2/1.7(E-5)]^{0.1} = 0.0085 \text{ m/sec},$ $\therefore u_{mb}/u_{mf} = 0.0085/0.0061 = 1.39.$

From Eq. (6.139),

 $\frac{u_{mb}}{u_{mf}} - \frac{82[1.7(E-5)]^{0.6}(1.2)^{0.06}}{9.81[84.5(E-6)]^{1.3}(1700-1.2)} = 1.35,$

which is in rough agreement.

(e) Voidage at minimum bubbling from Eq. (6.138):

$$\frac{\varepsilon_{mb}^3}{1-\varepsilon_{mb}} = 47.4 \left\{ \frac{[1.7(E-5)]^2}{9.81[84.5(E-6)]^3(1700)^2} \right\}^{0.5} = 0.1948,$$

$$\therefore \varepsilon_{mb} = 0.469.$$

It is not certain how nearly consistent this value is with those at minimum fluidization read off Figure 6.10(e). Only a limited number of characteristics of the solids are accounted for in Eq. (6.138).

(f) Operating gas velocity. The ratios of entraining and minimum fluidizing velocities for the two smallest particle sizes present are

$$0.049/0.0061 = 8.03$$
, for 30 μ m,
 $0.0054/0.0061 = 0.89$, for 10 μ m.

Entrainment of the smallest particles cannot be avoided, but an appreciable multiple of the minimum fluidizing velocity can be used for operation; say the ratio is 5, so that

$$u_f = 5u_{mf} = 5(0.0061) = 0.0305 \text{ m/sec.}$$

(g) Bed expansion ratio. From Figure 6.10(c) with $d_p = 84.5 \ \mu m$ or 0.0033 in. and $G_f / G_{mf} = 5$,

 $R = \frac{1.16}{1.22}$, by interpolation between the full lines, I 1.22, off the dashed line.

Take $\mathbf{R} = 1.22$ as more conservative. From Eq. (6.140) the ratio of voidages is

$$\varepsilon_{mb}/\varepsilon_{mf} = 5^{0.22} = 1.42.$$

From part (e), $\epsilon_{mb} = 0.469$ so that $\epsilon_{mf} = 0.469/1.42 = 0.330$. Accordingly, the ratio of bed levels is

$$L_{mb}/L_{mf} = (1 - \varepsilon_{mf})/(1 - \varepsilon_{mb}) = 0.67/0.531 = 1.262.$$

Although the value of ε_{mf} appears somewhat low, the value of *R* checks roughly the one from Figure 6.10(*c*).

(h) Fluctuations in level. From Figure 6.10(d), with $d_p = 0.0033$ in., the value of m' = 0.02, so that

$$r = \exp[0.02(5 - 1)] = 1.083.$$

(i) TDH from Figure 6.10(i). At $u_f = u_{mf} - 4(0.0061) = 0.0244$ m/sec, the abscissa is off the plot, but a rough extrapolation and interpolation indicates about 1.5 m for TDH.

(j) Dimensions of the bed and vessel. With a volumetric flow rate of $0.3 \text{ m}^3/\text{sec}$, the required diameter is

$$D = \sqrt{0.3/(0.305)(\pi/4)} = 3.54$$
 m

With a charge of 10,000 kg of solids and a voidage at minimum bubbling of 0.469, the height of the minimum bubbling bed is

$$L = \frac{10000}{1700(1-0.469)(\pi/4)D^2} = 1.13 \,\mathrm{m}.$$

This value includes the expansion factor which was calculated separately in item (g) but not the fluctuation parameter; with this correction the bed height is

$$L_b = 1.13(1.083) = 1.22$$
 m

The vessel height is made up of this number plus the TDH of 1.5 m or

vessel height =
$$1.22 + 1.5 = 2.72$$
 m.
Minimum Fluidization. The fundamental nature of this phenomenon has led to many correlations for its prediction. That of Leva (1959) applies to Reynolds numbers $\operatorname{Re}_{mf} = d_p G_{mf}/\mu < 5$, and is

$$G_{mf} = 688 D_p^{1.82} \frac{[\rho_F(\rho_S - \rho_F)]^{0.94}}{\mu^{0.88}}$$
(6.132)

in the common units G_{mf} in $lb/(hr)(sqft), D_p$ in inches, densities in lb/cuft, and viscosity in cP. In SI units it is

$$U_{mf} = \frac{0.0093 d_p^{1.82} (\rho_p - \rho_f)^{0.94}}{\mu^{0.88} \rho_f^{0.06}}$$
(6.133)

The degree of confidence that can be placed in the correlation is indicated by the plot of data on which it is based in Figure 6.10(f). An equation more recently recommended by Grace (1982) covers Reynolds numbers up to 1000:

$$\mathbf{Re}_{mf} = d_p u_{mf} \rho / \mu = \sqrt{(27.2)^2 + 0.0408(\mathbf{Ar})} = 27.2, \qquad (6.134)$$

where

Ar
$$= \rho(\rho_p - \rho)gd_p^3/\mu^2$$
. (6.135)

Here also the data show much scatter, so that pilot plant determinations of minimum fluidization rates usually are advisable.

Minimum Bubbling Conditions. Minimum bubbling velocities for Group B substances are about the same as the minimum fluidization velocities, but those of Group A substances are substantially greater. For Group A materials the correlation of Geldart and Abrahamsen *[Powder Technol* 19, 133 (1978)] for minimum bubbling velocity is

$$u_{mb} = 33d_p(\mu/\rho)^{-0.1}.$$
(6.136)

For air at STP this reduces to

$$u_{mb} = 100d_p.$$
 (6.137)

For cracking catalysts represented on Figure 6.10(g), Harriott and Simone (1983) present an equation for the ratio of the two kinds of velocities as

$$\frac{u_{mb}}{u_{mf}} = \frac{82\mu^{0.6}\rho^{0.06}}{gd_p^{1.3}(\rho_p - \rho)}.$$
(6.138)

The units of this equation are SI; the coefficient given by Cheremisinoff and Cheremisinoff (1984, p. 161) is incorrect. Figures 6.10(g) and (h) compare the two kinds of velocities over a range of particle diameters. Voidage at minimum bubbling is correlated by an equation of Cheremisinoff and Cheremisinoff (1984, p. 163):

$$\varepsilon_{mb}^{3}/(1-\varepsilon_{mb}) = 47.4(gd_{p}^{3}\rho_{p}^{2}/\mu^{2})^{-0.5}.$$
(6.139)

Bed Expansion and Fluctuation. The change of bed level with increasing gas rate is represented schematically in Figure 6.10(a). The height remains constant until the condition of minimum fluidization is reached, and the pressure drop tends to level off. Then the bed continues to expand smoothly until some of the gas begins to disengage from the homogeneous dense phase and forms bubbles. The point of onset of bubbling corresponds to a local

maximum in level which then collapses and attains a minimum. With increasing gas rate, the bed again continues to expand until entrainment develops and no distinct bed level exists. Beyond the minimum bubbling point, some fraction of the excess gas continues through the dense phase but that behavior cannot be predicted with any accuracy.

Some smoothed data of expansion ratio appear in Figure 6.10(c) as a function of particle size and ratio of flow rates at minimum bubbling and fluidization. The rather arbitrarily drawn dashed line appears to be a conservative estimate for particles in the range of $100 \,\mu$ m.

Ordinarily under practical conditions the flow rate is at most a few multiples of the minimum fluidizing velocity so the local maximum bed level at the minimum bubbling velocity is the one that determines the required vessel size. The simplest adequate equation that has been proposed for the ratio of voidages at minimum bubbling and fluidization is

$$\varepsilon_{mb}/\varepsilon_{mf} = (G_{mb}/G_{mf})^{0.22}$$
 (6.140)

$$= 2.64 \mu^{0.89} \rho^{0.54} / g^{0.22} d_p^{1.06} (\rho_p \ \rho)^{0.22}$$
(6.141)

The last equation results from substitution of Eq. (6.138) into (6.140). Then the relative bed level is found from

$$L_{mb}/L_{mf} = (1 - \varepsilon_{mf})/(1 - \varepsilon_{mb}). \tag{6.142}$$

Either ε_{mb} or ε_{mf} must be known independently before Eq. (6.141) can be applied, either by application of Eq. (6.139) for ε_{mb} or by reading off a value of ε_{mf} from Figure 6.8(c) or Figure 6.10(e). These values are not necessarily consistent.

At high gas velocities the bed level fluctuates. The ratio of maximum disturbed level to the average level is correlated in terms of G_f/G_{mf} and the particle diameter by the equation

$$r = \exp[m'(G_f - G_{mf})/G_{mf}], \tag{6.143}$$

where the coefficient m' is given in Figure 6.10(d) as a function of particle diameter.

Freeboard. Under normal operating conditions gas rates somewhat in excess of those for minimum fluidization are employed. As a result particles are thrown into the space above the bed. Many of them fall back, but beyond a certain height called the transport disengaging height (TDH), the entrainment remains essentially constant. Recovery of that entrainment must be accomplished in auxiliary equipment. The TDH is shown as a function of excess velocity and the diameter of the vessel in Figure 6.10(i). This correlation was developed for cracking catalyst particles up to 400 pm dia but tends to be somewhat conservative at the larger sizes and for other materials.

Viscosity. Dense phase solid-gas mixtures may be required to flow in transfer line catalytic crackers, between reactors and regenerators and to circulate in dryers such as Figures 9.13(e), (f). In dilute phase pneumatic transport the effective viscosity is nearly that of the fluid, but that of dense phase mixtures is very much greater. Some data are given by **Schügerl** (in Davidson and Harrison, 1971, **p**.261) and by Yates (1983). Apparent viscosities with particles of 50-550 μ m range from 700 to 1300 cP, compared with air viscosity of 0.017 cP at room temperature. Such high values of the viscosity place the flow definitely in the laminar flow range. However, information about friction in flow of fluidized mixtures through pipelines is not easy to find in the open literature. Someone must know since many successful transfer lines are in operation.

REFERENCES

General

- 1. M.M. Denn, Process Fluid Mechanics, Prentice-Hall, Englewood Cliffs, NJ, 1980.
- 2. 0. Levenspiel, Engineering Flow and Heat Exchange, Plenum, New York, 1984.
- 3. M. Modell and R.C. Reid, *Thermodynamics and Its Applications*, Prentice-Hall, Englewood Cliffs, NJ, 1983.
- 4. V.L. Streeter and E.B. Wylie, *Fluid Mechanics*, McGraw-Hill, New York, 1979.

Non-Newtonian Fluids

- G.W. Govier and K. Aziz, Flow Of Complex Mixtures in Pipes, Van Nostrand Reinhold, New York, 1972.
- N.I. Heywood, Pipeline design for non-Newtonian fluids, ht. Chem. Eng. Symp. Ser. No. 60, 33-52 (1980).
- J.W. Hoyt, The effect of additives on fluid friction, Trans. ASME J. Basic Eng., 258 (June 1972).
- 8. P.A. Longwell, Mechanics of Fluid Flow, McGraw-Hill, New York, 1966.
- R.D. Patel, Non-Newtonian *flow*, in *Handbook of Fluids in Motion*, (Cheremisinoff and Gupta, Eds.), Ann Arbor Science, Ann Arbor, MI, 1983, pp. 135-177.
- 10. A.H.P. Skelland, Non-Newtonian Flow and Heat Transfer, Wiley, New York, 1967.
- J.R. Van Wazer, J.W. Lyons, K.Y. Kim, and R.E. Colwell, Viscosity and Flow Measurement, Wiley-Interscience, New York, 1963.
- E.J. Wasp, J.P. Kenny, and R.L. Gandhi, *Solid Liquid Flow Slurry Pipeline Transportation*, Trans. Tech. Publications, Clausthal, Germany, 1977.

Two-phase Flow

13. D. Chisholm, Gas-liquid flow in pipeline systems, in Handbook of Fluids

in *Motion*, (Cheremisinoff and Gupta, Eds.) Ann Arbor Science, Ann Arbor, MI, 1983, pp. 483-513.

- 14. D. Chisholm, Two-Phase Flow in Pipelines and Heat Exchangers, George Godwin, London, 1983.
- **15.** G.W. Govier and K. Aziz, *The Flow of Complex Mixtures in Pipes*, Van Nostrand Reinhold, New York, 1972.
- G.F. Hewitt, Liquid-gas systems, in *Handbook of Multiphase Systems*, (G. Hetsroni, Ed.), Hemisphere, New York, 1982, pp. 2.1-2.94.

Gas-Solid (Pneumatic) Transport

- 17. G. Klinzing, Gas-Solid Transport, McGraw-Hill, New York, 1981.
- N.P. Cheremisinoff, and R. Gupta (Eds.), Gas-solid *flows*, in *Handbook* of *Fluids in Motion*, Ann Arbor Science, Ann Arbor, MI, 1983, pp. 623–860.
- C.S. Teo and L.S. Leung, Vertical flow of particulate solids in standpipes and risers, in Hydrodynamics of Gas-Solids Fluidization, (N.P. Cheremisinoff and P.N. Cheremisinoff, Eds.), Gulf, Houston, 1984, pp. 471-542.

Fluidization

- 20. J.S.M. Botteril, Fluid-Bed Heat Transfer, Academic, New York, 1975.
- N.P. Cheremisinoff and P.N. Cheremisinoff, Hydrodynamics of Gas-Solid Fluidization, Gulf, Houston, 1984.
- 22. J.F. Davidson and D. Harrison, Eds., *Fluidization*, Academic, New York, 1971.
- 23. J.R. Grace, Fluidization, Section 8 of G. Hetsroni, 1982.
- G. Hetsroni (Ed.), Handbook of Multiphase Systems, McGraw-Hill, New York, 1982.
- 25. M. Leva, Fluidization, McGraw-Hill, New York, 1959.
- **26.** J.C. Yates, *Fundamentals of Fluidized-Bed Chemical Processes*, Butterworths, London, 1983.

FLUID TRANSPORT EQUIPMENT

Ithough liquids particularly can be transported by operators carrying buckets, the usual mode of transport of fluids is through pipelines with pumps, blowers, compressors, or ejectors. Those categories of equipment will be considered in this chapter. A few statements will be made at the start about piping, fittings, and valves, although for the most part this is information best

gleaned from manufacturers' catalogs. Special problems such as mechanical flexibility of piping at elevated temperatures are beyond the scope here, and special problems associated with sizing of piping for thermosyphon reboilers and the suction side of pumps for handling volatile liquids are deferred to elsewhere in this book.

7.1. PIPING

Standard pipe is made in a discrete number of sizes that are designated by nominal diameters in inches, as "inches IPS (iron pipe size)." Table A5 lists some of these sizes with dimensions in inches. Depending on the size, up to 14 different wall thicknesses are made with the same outside diameter. They are identified by schedule numbers, of which the most common is Schedule 40. Approximately,

Schedule number = 1000 P/S,

where

```
P = internal pressure, psig
```

S = allowable working stress in psi.

Tubing for heat exchangers, refrigeration, and general service is made with outside diameters measured in increments of 1/16 or 1/8 in. Standard size pipe is made of various metals, ceramics, glass, and plastics.

Dimensional standards, materials of construction, and pressure ratings of piping for chemical plants and petroleum refineries are covered by ANSI Piping Code B31.3 which is published by the ASME, latest issue 1980. Many details also are given in such sources as Crocker and King, Piping Handbook (McGraw-Hill, New York, 1967), Perry's Chemical Engineers Handbook (1984), and Marks Standard Handbook for Mechanical Engineers (1987).

In sizes 2in. and less screwed fittings may be used. Larger joints commonly are welded. Connections to equipment and in lines whenever need for disassembly is anticipated utilize flanges. Steel flanges, flanged fittings, and valves are made in pressure ratings of 150, 300, 600, 900, 1500, and 2500 psig. Valves also are made in 125 and 250 psig cast iron. Pressure and temperature ratings of this equipment in various materials of construction are specified in the piping code, and are shown in *Chem. Eng. Handbook* 1984, pp. 6.75-6.78.

VALVES

Control of flow in lines and provision for isolation of equipment when needed are accomplished with valves. The basic types are relatively few, some of which are illustrated in Figure 7.1. In gate valves the flow is straight through and is regulated by raising or lowering the gate. The majority of valves in the plant are of this type. In the wide open position they cause little pressure drop. In globe valves the flow changes direction and results in appreciable friction even in the wide open position. This kind of valve is essential when tight shutoff is needed, particularly of gas flow. **Multi**- pass plug cocks, butterfly valves, slide valves, check valves, various quick-opening arrangements, etc. have limited and often indispensable applications, but will not be described here.

The spring in the relief valve of Figure 7.1(c) is adjusted to open when the pressure in the line exceeds a certain value, at which time the plug is raised and overpressure is relieved; the design shown is suitable for pressures of several hundred psig.

More than 100 manufacturers in the United States make valves that may differ substantially from each other even for the same line size and pressure rating. There are, however, independent publications that list essentially equivalent valves of the several manufacturers, for example the books of Zappe (1981) and Lyons (1975).

CONTROL VALVES

Control valves have orifices that can be adjusted to regulate the flow of fluids through them. Four features important to their use are capacity, characteristic, rangeability and recovery.

Capacity is represented by a coefficient

 $C_d = C_v / d^2,$

where d is the diameter of the valve and C_v is the orifice coefficient in equations such as the following

$$Q = C_v \sqrt{(P_1 - P_2)/\rho_w}, \text{ gal/min of liquid,}$$

$$Q = 22.7C_v \sqrt{(P_1 - P_2)P_2/\rho_a T}, \text{ SCFM of gas when } P_2/P_1 > 0.5,$$

$$Q = 11.3C_v P_1/\sqrt{\rho_a T}, \text{ SCFM of gas when } P_2/P_1 < 0.5,$$

where P_i is pressure in psi, ρ_w is specific gravity relative to water, ρ_a is specific gravity relative to air, and T is temperature "R. Values of C_d of commercial valves range from 12 for double-seated globe valves to 32 for open butterflies, and vary somewhat from manufacturer to manufacturer. Chalfin (1980) has a list.

Characteristic is the relation between the valve opening and the flow rate. Figure 7.1(h) represents the three most common forms. The shapes of plugs and ports can be designed to obtain any desired mathematical relation between the pressure on the diaphragm, the travel of the valve stem, and the rate of flow through the port. *Linear* behavior is represented mathematically by Q = kx and *equal percentage* by Q = k, $exp(k_2x)$, where x is the valve opening. *Quick-opening* is a characteristic of a bevel-seated or flat disk type of plug; over a limited range of 10–25% of the maximum stem travel is approximately linear.

Over a threefold load change, the performances of linear and equal percentage valves are almost identical. When the pressure drop across the valve is less than 25% of the system drop, the equal

percentage type is preferred. In fact, a majority of characterized valves currently are equal percentage.

Rangeability is the ratio of maximum to minimum flows over which the valve can give good control. This concept is difficult to quantify and is not used much for valve selection. A valve generally can be designed properly for a suitably wide flow range.

Recovery is a measure of the degree of pressure recovery at the valve outlet from the low pressure at the vena contracta. When flashing occurs at the vena contracta and the pressure recovery is high, the bubbles **collapse** with resulting cavitation and noise. The more streamlined the valve, the more complete the pressure recovery; thus, from this point of view streamlining seems to be an undesirable quality. A table of recovery factors of a number of valve types is given by Chalfin (1980); such data usually are provided by manufacturers.

These characteristics and other properties of 15 kinds of valves are described by Chalfin (1980).

Pressure drop. Good control requires a substantial pressure drop through the valve. For pumped systems, the drop through the valve should be at least 1/3 of the pressure drop in the system, with a minimum of 15 psi. When the expected variation in flow is small, this rule can be relaxed. In long liquid transportation lines, for instance, a fully open control valve may absorb less than 1% of the system pressure drop. In systems with centrifugal pumps, the variation of head with capacity must be taken into account when sizing the valve. Example 7.2, for instance, illustrates how the valve drop may vary with flow in such a system.

Types of valves. Most flow control valves are operated with adjustable air pressure on a diaphragm, as in Figure 7.1(d), since this arrangement is more rapid, more sensitive and cheaper than



Figure 7.1. Some kinds of manual and automatically controlled valves. (a) Gate valve, for the majority of applications. (b) Globe valve, when tight shutoff is needed. (c) Swing check valve to ensure flow in one direction only. (d) A pressure relief valve, in which the plug is raised on overpressure. (e) A control valve with a single port. (f) A double-port, reverse-acting control valve. (g) A control valve with a double port, in which the correct opening is maintained by air pressure above the diaphragm. (A) valve body; (B) removable seat; (C) discs; (D) valve-stem guide; (E) guide bushing; (F) valve bonnet; (G) supporting ring; (H) supporting arms; (J) diaphragm; (K) coupling between diaphragm and valve stem; (L) spring-retaining rod; (M) spring; (N) spring seat; (0) pressure connection. (Fischer.) (h) Relation between fractional opening and fractional flow of three modes of valve openings.

electrical motor control. Double-ported valve (d) gives better control at large flow rates; the pressures on the upper and lower plugs are balanced so that less force is needed to move the stem. The single port (e) is less expensive but gives a tighter shutoff and is generally satisfactory for noncritical service. The reverse acting valve (f) closes on air failure and is desirable for reasons of safety in some circumstances.

7.2. PUMP THEORY

Pumps are of two main classes: centrifugal and the others. These others mostly have positive displacement action in which the discharge rate is largely independent of the pressure against which they work. Centrifugal pumps have rotating elements that impart



high velocity initially and high pressure head ultimately to the liquid. Elements of their theory will be discussed here. A glossary of pump terms and terms relating primarily to centrifugal pumps are defined in the Glossary at the end of this chapter. The chief variables involved in pump theory are listed here with typical units:

D, diameter of impeller (ft or m), H, output head (ft or m), n, rotational speed (1/sec), \dot{P} , output power (HP or kW), Q, volumetric discharge rate (cfs or m³/sec), μ , viscosity (lb/ft sec or N sec/m²), ρ , density (lb/cuft or kg/m³), ε , surface roughness (ft or m).

BASIC RELATIONS

A dimensional analysis with these variables reveals that the functional relations of Eqs. (7.1) and (7.2) must exist:

$$gH/n^2D^2 = \phi_1(Q/nD^3, D^2n\rho/\mu, \varepsilon/D), \qquad (7.1)$$

$$P/\rho n^{3} D^{3} = \phi_{2}(Q/nD^{3}, D^{2} n\rho/\mu, \varepsilon/D).$$
(7.2)

The group $D^2 n \rho / \mu$ is the Reynolds number and ε / D is the roughness ratio. Three new groups also have arisen which are named

capacity coefficient,
$$C_Q = Q/nD^3$$
, (7.3)

head coefficient,
$$C_H = gH/n^2D^2$$
, (7.4)

power coefficient,
$$CP = P/\rho n^3 D^5$$
. (7.5)

The hydraulic efficiency is expressed by these coefficients as

$$\eta = gH\rho Q/P = C_H C_Q/C_P. \tag{7.6}$$

Although this equation states that the efficiency is independent of the diameter, in practice this is not quite true. An empirical relation is due to Moody [ASCE Trans. 89, 628 (1926)]:

$$\eta_2 = 1 \quad (1 - \eta_1)(D_1/D_2)^{0.25}. \tag{7.7}$$

Geometrically similar pumps are those that have all the dimensionless groups numerically the same. In such cases, two different sets of operations are related as follows:

$$Q_2/Q_1 = (n_2/n_1)(D_2/D_1)^3,$$
(7.8)

$$H_2/H_1 = (n_2 D_2/n_1 D_1)^2, (7.9)$$

$$\dot{P}_2/\dot{P}_1 = (\rho_2/\rho_1)(n_2/n_1)^3 (D_2/D_1)^5.$$
 (7.10)

The performances of geometrically similar pumps also can be represented in terms of the coefficients C_Q , C_H , C_P , and η . For instance, the data of the pump of Figure 7.2(a) are transformed into the plots of Figure 7.2(b). An application of such generalized curves is made in Example 7.1.

Another dimensionless parameter that is independent of diameter is obtained by eliminating D between C_Q and C_H with the result,

$$N_s = nQ^{0.5} / (gH)^{0.75}.$$
(7.11)

This concept is called the specific speed. It is commonly used in the

Figure 7.1—(continued)

EXAMPLE 7.1

Application of Dimensionless Performance Curves Model and prototypes are represented by the performance curves of Figure 7.2. Comparisons are to be made at the peak efficiency, assumed to be the same for each. Data off Figure 7.2(b) are:

$$\begin{split} \eta &= 0.93, \\ C_{H} &= g H/n^{2} D^{2} = 5.2, \\ C_{P} &= \dot{P}/\rho n^{3} D^{5} = 0.69, \\ C_{Q} &= Q/n D^{3} = 0.12. \end{split}$$

(a) The prototype is to develop a head of 76 m:

$$n = \left(\frac{gH}{C_H D^2}\right)^{0.5} = \left(\frac{9.81(76)}{5.2(0.371)^2}\right)^{0.5} = 32.27 \text{ rps},$$

$$Q = nD^3C_Q = 32.27(0.371)^3(0.12) = 0.198 \text{ m}^3/\text{sec},$$

$$\dot{P} = \rho n^3 D^5C_p = 1000(32.27)^3(0.371)^5(0.69)$$

$$= 0.163(10^6) \text{ W}, 163 \text{ kW}.$$

(b) The prototype is to have a diameter of 2 m and to rotate at 400 rpm:

 $Q = nD^{3}C_{Q} = (400/60)(2)^{3}(0.12) = 6.4 \text{ m}^{3}/\text{sec},$ $H = n^{2}D^{2}C_{H}/g = (400/60)^{2}(2)^{2}(5.2)/9.81 = 94.2 \text{ m},$

- $\dot{P} = \rho n^3 D^5 C_p = 1000(400/60)^3 (2)^5 (0.69)$ = 6.54(10⁶) kgm²/sec³, 6.54(10⁶) N m/sec, 6540 kW.
- (c) Moody's formula for the effect of diameter on efficiency gives

$$\eta_2 = 1 - (1 - \eta_1)(D_1/D_2)^{0.25} = 1 - 0.07(0.371/2)^{0.25}$$

= 0.954 at 2 m,

compared with 0.93 at 0.371 m.

(d) The results of (a) and (b) also are obtainable directly from Figure 7.2(a) with the aid of Eqs. (7.7), (7.8), and (7.9). Off the figure at maximum efficiency,

$$\eta = 0.93$$
, Q = 0.22, H = 97, a n d P = 218

When the new value of H is to be 76m and the diameter is to remain the same,

$$n_2 = 35.6(H_2/H_1)^{0.5} = 35.6(76/97)^{0.5} = 31.5 \text{ rps},$$

$$Q_2 = Q_1(n_2/n_1) = 0.22(H_2/H_1)^{0.5} = 0.195 \text{ m}^3/\text{sec},$$

$$\dot{P}_2 = \dot{P}_1(\rho_2/\rho_1)(n_2/n_1)^3(D_2/D_1)^5 = 218(H_2/H_1)^{1.5} = 151.2 \text{ kW}.$$

These values agree with the results of (a) within the accuracy of reading the graphs.



Figure 7.2. Performance curves in dimensional and dimensionless forms: (a) Data of a pump with a specific diameter and rotation speed. (b) Dimensionless performance curves of all pumps geometrically similar to (a). The dashed lines identify the condition of peak efficiency. (After Daugherty and Franrini, Fluid Mechanics with Engineering Applications, McGraw-Hill, New York, 1957).

EXAMPLE 7.2 Operating Points of Single and Double Pumps in Parallel and Series

The head loss in a piping system is represented by the equation

$$H_s = 50 + 6.0(Q/100)^2 + H_v$$

where H_v is the head loss in the control valve. The pump to be used has the characteristic curve of the pump of Figure 7.7(b) with an 8 in. impeller; that curve is represented closely by the equation

 $H_p = 68 - 0.5(Q/100) - 4.5(Q/100)^2$.

The following will be found (see Figure 7.17):

- (a) The values of H_v corresponding to various flow rates Q gpm.
 (b) The flow rate and head on the pumps when two pumps are connected in parallel and the valve is wide open (H_v = 0).
- (c) The same as (b) but with the pumps in series.
- (d) The required speed of the pump at 80 gpm when no control valve is used in the line.

(a) The operating point is found by equating H_s and H_p from which

$$H_v = 68 - 0.5(Q/100) - 4.5(Q/100)^2 - [50 + 6.0(Q/100)^2]$$

Some values are

Q/100	0.8	1.0	1.2	1.286
H, H,	10. 88	7.00	2. 28	0 59. 92

(b) In parallel each pump has half the total flow and the same head H_s :

$$50 + 6.0(Q/100)^2 = 68 - (0.5/2)(Q/100) - (4.5/4)(Q/100)^2$$
,
∴ Q = 157.2 gpm, $H_e = 64.83$ ft.

(c) In series each pump has the same flow and one-half the total head loss:

 $\frac{1}{2}(50 + 6.0(Q/100)^2] = 68 - 0.5(Q/100) - 4.5(Q/100)^2,$ $\therefore Q = 236.1 \text{ gpm}, H_s = 83.44 \text{ ft}.$

Series flow allows 50% greater gpm than parallel.

(d)
$$H_s = 50 + 4.8 = 54.8$$
,
 $H_p = (68 - 0.4 - 2.88)(n/1750)^2$,
 $\therefore n = 1750\sqrt{54.8/64.72} = 1610 \text{ pm}$

mixed units

$$N_s = (\text{rpm})(\text{gpm})^{0.5} / (\text{ft})^{0.75}.$$
 (7.12)

For double suction pumps, Q is one half the pump output.

The net head at the suction of the pump impeller must exceed a certain value in order to prevent formation of vapor and resulting cavitation of the metal. This minimum head is called the net positive suction head and is evaluated as

Usually each manufacturer supplies this value for his equipment. (Some data are in Figure 7.7.) A suction specific speed is defined as

$$S = (rpm)(gpm)^{0.5} / (NPSH)^{0.75}$$
. (7.14)

Standards for upper limits of specific speeds have been established, like those shown in Figure 7.6 for four kinds of pumps. When these values are exceeded, cavitation and resultant damage to the pump may occur. Characteristic curves corresponding to widely different values of N_s are shown in Figure 7.3 for several kinds of pumps handling clear water. The concept of specific speed is utilized in Example 7.3. Further data are in Figure 7.6.

Recommendations also are made by the Hydraulic Institute of suction specific speeds for multistage boiler feed pumps, with S = 7900 for single suction and S = 6660 for double suction. Thus the required NPSH can be found by rearrangement of Eq. (7.14) as

NPSH =
$$[(rpm)(gpm)^{0.5}/S]^{4/3}$$
. (7.15)

For example, at 3500 rpm, 1000 gpm, and S = 7900, the required NPSH is 34 ft.

For common fluids other than water, the required NPSH usually *is* lower than for cold water; some data are shown in Figure 7.16.

PUMPINGSYSTEMS

The relation between the flow rate and the head developed by a centrifugal pump is a result of its mechanical design. Typical curves are shown in Figure 7.7. When a pump is connected to a piping system, its head must match the head loss in the piping system at the prevailing flow rate. The plot of the flow rate against the head loss in a line is called the system curve. The head loss is given by the mechanical energy balance,

$$H_s = \frac{\Delta P}{\rho} + \frac{\Delta u^2}{2g_c} + \Delta z + \frac{fLu^2}{2gD} + H_v, \qquad (7.16)$$

where H_{ν} is the head loss of a control value in the line.

The operating point may be found as the intersection of plots of the pump and system heads as functions of the flow rate. Or an equation may be fitted to the pump characteristic and then solved simultaneously with Eq. (7.16). Figure 7.17 has such plots, and Example 7.2 employs the algebraic method.

In the normal situation, the flow rate is the specified quantity. With a particular pump curve, the head loss of the system may need to be adjusted with a control valve in the line to make the system and pump heads the same. Alternately, the speed of the pump can be adjusted to make the pump head equal to that of the system. From Eq. (7.9) the relation between speeds and pump heads at two



Figure 7.3. Performance curves of single-suction impellers corresponding to two values of the specific speed. (a) $N_s = 1550$, centrifugal pump. (b) $N_s = 10,000$, mixed and axial flow pumps.

conditions is

$$n_2 = n_1 (H_2/H_1)^{0.5}. (7.17)$$

Example 7.2 is of cases with control valve throttling and pump speed control. In large systems, the value of power saved can easily overbalance the extra cost of variable speed drives, either motor or steam turbine.

When needed, greater head or greater capacity may be obtained by operating several pumps in series or parallel. In parallel operation, each pump develops the same head (equal to the system head), and the flow is the sum of the flows that each pump delivers at the common head. In series operation, each pump has the same flow rate and the total head is the sum of the heads developed by the individual pumps at the prevailing flow rate, and equal to the system head. Example 7.1 deals with a pair of identical pumps, and corresponding system and head curves are shown in Figure 7.17.

7.3. PUMP CHARACTERISTICS

A centrifugal pump is defined in the glossary at the end of this chapter as a machine in which a rotor in a casing acts on a liquid to give it a high velocity head that is in turn converted to pressure head by the time the liquid leaves the pump. Other common nomenclature relating to the construction and performance of centrifugal and related kinds of pumps also is in that table.



Figure 7.4. Performance of several kinds of pumps. (a) Comparison of small centrifugal and turbine pumps (*Kristal and Annett, 1940).* (b) An axial flow pump operating at 880 rpm (Chem. Eng. Handbook, 1973). (c) An external gear pump like that of Figure 7.12(e) (*Viking Pump* Co.). (d) A screw-type positive displacement pump. (e) NPSH of reciprocating positive displacement pumps.



Figure 7.4—(continued)

(a) E	Efficiencies	as	% of	those	with	direc	t pist	on d	rive:		
Str Cra	oke, in ank-and-flyw	vheel	pu	mp	5	8	10	20 87	30 88	40 90	50 92
Pi: Hie	ston pump ah-pressure	pi	ımp		60 55	70 64	74 67	84 76	86 78	88 80	90 81
(b)	Efficiencies	of	cranl	cshaft-c	lriven	pum	ps of	var	ious	sizes:	•
Wa Effi	ter H ciency	P (%)	3 55	5 65	10 72	20 77	30 80	50 83	75 85	100 86	200 88

(c) % of flow above and below the mean; curve is shown for triplex double-acting:



(d) Efficiency as a function of % reduced pressure or % reduced speed:

% Full-Load	Mechanical		Mechanical
Developed Pressure	Efficiency	% Speed	Efficiency
2 0	8 2	44	93.3
4 0	88	50	92.5
60	90.5	73	92.5
80	9 2	100	92.5
100	92.5		





Figure 7.5. Data relating to the performance of piston and plunger pumps.

EXAMPLE 7.3

Check of Some Performance Curves with the Concept of Specific Speed

(a) The performance of the pump of Figure 7.7(b) with an 8in. impeller will be checked by finding its specific speed and comparing with the recommended upper limit from Figure 7.6(b). Use Eq. (7.12) for N_s

Q (gpm)	100	200	300
H (řť.)	268	255	225
N _e (calcd)	528	776	1044
N [Fig. 7.10(a)]	2050	2150	2500
NPSH	5	7	13

Clearly the performance curves are well within the recommended upper limits of specific speed.

(b) The manufacturer's recommended NPSH of the pump of Figure 7.7(c) with an 8 in. impeller will be checked against values from Eq. (7.15) with S = 7900:

Q (gpm)	100	150	200
H (ft)	490	440	300
NPSH (mfgr)	10	18	35
NPSH (Eq. (7.15))	7.4	9.7	11.8

The manufacturer's recommended NPSHs are conservative.



Figure 7.6. Upper specific-speed limits for (a) double-suction pumps (shaft through impeller eye) handling clear water at $85^{\circ}F$ at sea level, (b) single-suction pumps (shaft through impeller eye) handling clear water at $85^{\circ}F$ at sea level, (c) single-suction pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction mixed- and axial-flow pumps (overhung-impeller type) handling clear water at $85^{\circ}F$ at sea level, (d) single-suction



Figure 7.7. Characteristic curves of centrifugal pumps when operating on water at 85°F (Allis Chalmers Co.). (a) Single suction, 1750 rpm. (b) The pump of (a) operated at 3500 rpm. (c) Multistage, single suction, 3550 rpm.

The basic types of **centrifugals** are illustrated in Figure 7.9. A volute is a gradually expanding passage in which velocity is partially converted to pressure head at the outlet. The diffuser vanes of Figures 7.9(b) and 7.10(d) direct the flow smoothly to the periphery. The volute design is less expensive, more amenable to use with impellers of different sizes in the same case, and, as a consequence, by far the most popular construction. Diffuser construction is used to a limited extent in some high pressure, multistage machines. The double suction arrangement of Figure 7.9(d) has balanced axial thrust and is favored particularly for severe duty and where the lowered NPSH is an advantage. Multistage pumps, however, are exclusively single suction.

Some of the many kinds of impellers are shown in Figure 7.10. For clear liquids, some form of closed impeller [Figure 7.10(c)] is favored. They may differ in width and number and curvature of the vanes, and of course in the primary dimension, the diameter. Various extents of openness of impellers, [Figs. 7.10(a) and (b)] are desirable when there is a possibility of clogging as with slurries or pulps. The impeller of Figure 7.10(e) has both axial propeller and

centrifugal vane action; the propeller confers high rates of flow but the developed pressure is low. Figure 7.3(b) represents a typical axial pump performance.

The turbine impeller of Figure 7.10(h) rotates in a case of uniform diameter, as in Figure 7.12(j). As Figure 7.4(a) demonstrates, turbine pump performance resembles that of positive displacement types. Like them, turbines are essentially self-priming, that is, they will not vapor bind.

All rotating devices handling fluids require seals to prevent leakage. Figure 7.13 shows the two common methods that are used: stuffing boxes or mechanical seals. Stuffing boxes employ a soft packing that is compressed and may be lubricated with the pump liquid or with an independent source. In mechanical seals, smooth metal surfaces slide on each other, and are lubricated with a very small leakage rate of the pump liquid or with an independent liquid.

Performance capability of a pump is represented on diagrams like those of Figure 7.7. A single point characterization often is made by stating the performance at the peak efficiency. For example, the pump of Figure 7.7(c) with a 9 in. impeller is called a

(a) Single-suction, 1800 rpm standard pumps:



			Horsepower
Key No.	Suction and Discharge	Approximate cost	Range at 1.0 Sp Gr
1	4 x 3	\$1200	$7\frac{1}{2}-25$
2	6 x 4	1350	20-30
3	4 x 3	1200	15-25
4	5 x 4	1500	15-30
5	2x1;	750	$2-7\frac{1}{2}$
6	$2\frac{1}{2} \times 2$	1050	3-10
7	4 x 3	1200	5-15
8	5 x 4	1350	7 1 -20
9	5 x 4		-
	(1200 rpm)	1500	$3-7\frac{1}{2}$
10	$1\frac{1}{2} \times 1\frac{1}{2}$	700	1 - 2
11	2 x 2	750	1 ¹ / ₂ -3
12	$3 \times 2\frac{1}{2}$		
	(1200 rpm)	1050	1 ¹ / ₂ 3
13	4 x 3		
	(1200 rpm)	1200	2-5
14	5 x 4		
	(1200 rpm)	1350	2-5
15	$2\frac{1}{2} \times 2\frac{1}{2}$	500	$\frac{3}{4} - 1\frac{1}{2}$
16	3 x 3	600	1-2

(b) Single-suction, 3600 rpm standard pumps:



(c) Single-suction 1800 and 3600rpm refinery pumps for elevated temperatures and pressures:

 4×3

Suction and

Discharge

9

Kev

No.



1650

U.S. G. P. M

cost

Horsepower Approximate Range at 1.0 Sp Gr

1	$2x 1\frac{1}{2}$	\$3400	7 2 -30
2	3 x 2 (3600 rpm)	3700	15-50
3	4 x 3	4300	20-75
4	6 x 4	4800	40-125
5	3 x 2	4200	5-15
6	4 x3 (1800 rpm)	4500	7 ¹ / ₂ -20
7	6 x 4	5400	15-40
8	2x1;	3400	1-5
9	3 × 2 (1800 rpm)	3700	$2-7\frac{1}{2}$
10	4 x 3	4300	3-10
11	6 x 4	4800	5-15

Figure 7.8. Typical capacity-head ranges of some centrifugal pumps, their 1978 costs and power requirements. Suction and discharge are in inches (Evans, 1979, Vol. 1).



(a)







(c)



(d)



Figure 7.9. Some types of centrifugal pumps. (a) Single-stage, single suction volute pump. (b) Flow path in a volute pump. (c) Double suction for minimizing axial thrust. (d) Horizontally split casing for ease of maintenance. (e) Diffuser pump: vanes V are fixed, impellers P rotate. (f) A related type, the turbine pump.



Figure 7.10. Some types of impellers for centrifugal pumps. (a) Open impeller. (b) **Semiopen** impeller. (c) Shrouded impeller. (d) Axial flow (propeller) type. (e) Combined axial and radial flow, open type. (f) Shrouded mixed-flow impeller. (g) Shrouded impeller (P) in a case with diffuser vanes (V). (h) Turbine impeller.

175 gpm and 56Oft head pump at a peak efficiency of 57%; it requires a 15 ft suction lift, an 18 ft NPSH and 43 BHP. Operating ranges and costs of commercial pumps are given in Figure 7.8. General operating data are in Figure 7.4.

Although centrifugal pumps are the major kinds in use, a great variety of other kinds exist and have limited and sometimes unique applications. Several kinds of positive displacement types are sketched in Figure 7.12. They are essentially self-priming and have a high tolerance for entrained gases but not usually for solids unless they may be crushed. Their characteristics and applications are discussed in the next section.

7.4. CRITERIA FOR SELECTION OF PUMPS

The kind of information needed for the specification of centrifugal, reciprocating and rotary pumps is shown on forms in Appendix B. General characteristics of classes of pumps are listed in Table 7.1 and their ranges of performance in Table 7.2. Figure 7.14 shows recommended kinds of pumps in various ranges of pressure and flow rate. Suitable sizes of particular styles of a manufacturer's pumps are commonly represented on diagrams like those of Figure 7.8. Here pumps are identified partly by the sizes of suction and discharge nozzles in inches and the rpm; the key number also



Figure 7.11. Approximate efficiencies of centrifugal pumps in terms of GPM and head in feet of liquid.

identifies impeller and case size and other details which are stated in a catalog. Each combination of head and capacity will have an efficiency near the maximum of that style. Although centrifugal pumps function over a wide range of pressure and flow rates, as represented by characteristic curves like those of Figures 7.2 and 7.7, they are often characterized by their performance at the peak efficiency, as stated in the previous section. Approximate efficiencies of centrifugal pumps as functions of head and capacity are on Figure 7.11 and elsewhere here.

Centrifugal pumps have a number of good qualities:

- **1.** They are simple in construction, are inexpensive, are available in a large variety of materials, and have low maintenance cost.
- They operate at high speed so that they can be driven directly by electrical motors.
- 3. They give steady delivery, can handle slurries and take up little floor space.

Some of their drawbacks are

 Single stage pumps cannot develop high pressures except at very high speeds (10,000 rpm for instance). Multistage pumps for high



Figure 7.12. Some types of positive displacement pumps. (a) Valve action of a double acting reciprocating piston pump. (b) Discharge curve of a single acting piston pump operated by a crank; half-sine wave. (c) Discharge curve of a simplex double acting pump as in (a). (d) Discharge curve of a duplex, double acting pump. (e) An external gear pump; characteristics are in Figure 7.8(c). (f) Internal gear pump; the outer gear is driven, the inner one follows. (g) A double screw pump. (h) Peristaltic pump in which fluid is squeezed through a flexible tube by the follower. (i) Double diaphragm pump shown in discharge position (BIF unit of General Signal). (j) A turbine pump with essentially positive displacement characteristics (*data* on *Fig.* 7. 4(a)].



Figure 7.13. Types of seals for pump shafts. (a) Packed stuffing box; the sealing liquid may be from the pump discharge or from an independent source. (b) Water cooled stuffing box. (c) Internal assembly mechanical seal; the rotating and fixed surfaces are held together by the pressure of the pump liquid which also serves as lubricant; a slight leakage occurs. (d) Double mechanical seal with independent sealing liquid for handling toxic or inflammable liquids.



Figure 7.14. Range of applications of various kinds of pumps. (a) Range of applications of single and double suction pumps (*Allis-Chalmers* Co.). (b) Recommended kinds of pumps for various kinds of head and flow rate (*Fairbanks, Morse, and Co.*).

pressures are expensive, particularly in corrosion-resistant materials

- 5. Efficiencies drop off rapidly at flow rates much different from those at peak efficiency.
- 6. They are not self-priming and their performance drops off rapidly with increasing viscosity. Figure 7.15 illustrates this effect

On balance, centrifugal pumps always should be considered first in comparison with reciprocating or rotary positive displacement types, but those do have their places. Range of applications of various kinds of pumps are identified by Figure 7.14.

Pumps with reciprocating pistons or plungers are operated with steam, motor or gas engine drives, directly or through gears or belts. Their mode of action is indicated on Figure 7.12(a). They are always used with several cylinders in parallel with staggered action to smooth out fluctuations in flow and pressure. Figure 7.5(c) shows that with five cylinders in parallel the fluctuation is reduced to a maximum of 7%. External fluctuation dampers also are used. Although they are self-priming, they do deteriorate as a result of cavitation caused by release of vapors in the cylinders. Figure 7.4(e) shows the NPSH needed to repress cavitation. Application of reciprocating pumps usually is to low capacities and high pressures

of 50-1000 atm or more. Some performance data are shown in Figure 7.5.

Diaphragm pumps [Fig. 7.12(i)] also produce pulsating flow. They are applied for small flow rates, less than 100 gpm or so, often for metering service. Their utility in such applications overbalances the drawback of their intrinsic low efficiencies, of the order of 20%.

Screw pumps [Fig. 7.12(g)] are suited for example to high viscosity polymers and dirty liquids at capacities up to 2000 gpm and pressures of 200 atm at speeds up to 3000 rpm. They are compact, quiet, and efficient. Figure 7.4(d) shows typical performance data.

Gear pumps [Figs. 7.12(e) and (f)] are best suited to handling clear liquids at a maximum of about 1000 gpm at 150atm. Typical performance curves are shown in Figure 7.4(c).

Peristaltic pumps [Fig. 7.12(h)] move the liquid by squeezing a tube behind it with a rotor. Primarily they are used as metering pumps at low capacities and pressures in corrosive and sanitary services when resistant flexible tubes such as those of teflon can be used, and in laboratories.

Turbine pumps [Figs. 7.9(f), 7.12(i), and 7.4(a)] also are called regenerative or peripheral. They are primarily for small capacity and high pressure service. In some ranges they are more efficient than **centrifugals**. Because of their high suction lifts they are suited to handling volatile liquids. They are not suited to viscous liquids or abrasive slurries.

7.5. EQUIPMENT FOR GAS TRANSPORT

Gas handling equipment is used to transfer materials through pipe lines, during which just enough pressure or head is generated to overcome line friction, or to raise or lower the pressure to some required operating level in connected process equipment. The main classes of this kind of equipment are illustrated in Figures 7.18 and 7.19 and are described as follows.

- 1. *Fans* accept gases at near atmospheric pressure and raise the pressure by approximately 3% (12in. of water), usually on air for ventilating or circulating purposes.
- 2. *Blowers* is a term applied to machines that raise the pressure to an intermediate level, usually to less than 40 psig, but more than accomplished by fans.
- 3. *Compressors* are any machines that raise the pressure above the levels for which fans are used. Thus, in modern terminology they include blowers.
- Jet compressors utilize a high pressure gas to raise other gases at low pressure to some intermediate value by mixing with them.
- 5 *Vacuum* pumps produce subatmospheric pressures in process equipment. Often they are compressors operating in reverse but other devices also are employed. Operating ranges of some commercial equipment are stated in Table 7.3.
- 6. Steam jet ejectors are used primarily to evacuate equipment but also as pumps or compressors. They are discussed in Section 7.7.

Application ranges of fans and compressors are indicated on Figures 7.20 and 7.21. Some of these categories of equipment now will be discussed in some detail.

FANS

Fans are made either with axial propellers or with a variety of radial vanes. The merits of different directions of curvature of the vanes are stated in Figure 7.24 where the effect of flow rate of pressure, power, and efficiency also are illustrated. Backward curved vanes are preferable in most respects. The kinds of controls used have a marked effect on fan performance as Figure 7.23 shows. Table 7.4 shows capacity ranges and other characteristics of various kinds of

TABLE 7.1. Characteristics	s of	Various	Kinds	of	Pumps
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Pump Type	Construction Style	Construction Characteristics	Notes
Centrifugal	single-stage overhung, process type	impeller cantilevered beyond bearings	capacity varies with head
(horizontal)	two-stage overhung	two impellers cantilevered beyond bearings	used for heads above single-stage capability
	single-stage impeller between bearings	impeller between bearings; casing radially or axially split	used for high flows to 1083 ft (330 ml head
	chemical	casting patterns designed with thin sections for high-cost alloys	have low pressure and temperature ratings
	slurry	designed with large flow passages	low speed and adjustable axial clearance; has erosion control features
	canned	no stuffing box; pump and motor enclosed in a pressure shell	low head capacity limits when used in chemical services
	multistage, horizontally split casing	nozzles located in bottom half of casing	have moderate temperature-pressure ranges
	multistage, barrel type	outer casing contains inner stack of diaphragms	used for high temperature-pressure ratings
Centrifugal (vertical)	single-stage, process type	vertical orientation	used to exploit low net positive section head (NPSH) requirements
	multistage	many stages with low head per stage	low-cost installation
	inline	inline installation, similar to a valve	low-cost installation
	high speed	speeds to 380 rps, heads to 5800 ft (1770 m)	high head/low flow; moderate costs
	slump	casing immersed in sump for easy priming and installation	low cost
	multistage, deep well	long shafts	used for water well service
Axial	propeller	propeller-shaped impeller	vertical orientation
Turbine	regenerative	fluted impeller. Flow path resembles	capacity independent of head: low
		screw around periphery	flow/high head performance
Reciprocating	piston, plunger	slow speeds	driven by steam engine cylinders or motors through crankcases
	metering	consists of small units with precision flow control system	diaphragm and packed plunger types
	diaphragm	no stuffing box	used for chemical slurries; can be pneumatically or hydraulically actuated
Rotary	screw	I, 2, or 3 screw rotors	for high-viscosity, high-flow high-pressure services
	gear	intermeshing gear wheels	for high-viscosity, moderate- pressure/moderate-flow services

(Cheremisinoff, 1981).

TABLE 7.2. Typical Performances of Various Kinds of Pumps^a

Туре	Style	Capacity (gpm)	Max Head (ft)	Max P (psi)	NPSH (ft)	Мах 7 (°F)	Efficiency (%)
Centrifugal	single-stage overhung	15-5,000	492	600	6.56-19.7	851	20-80
(horizontal)	two-stage overhung	15-l ,200	1394	600	6.56-22.0	851	20-75
	single-stage impeller between						
	bearings	15-40.000	1099	980	6.56-24.9	401-851	30-90
	chemical	1000	239	200	3.94-19.7	401	20-75
	slurry	1000	394	600	4.92-24.9	851	20-80
	canned	1-20.000	4921	10,000	6.56-19.7	1004	20-70
	multistage horizontal split	20-I 1,000	5495	3000	6.56-19.7	401-500	65-90
	multistage, barrel type	20-9.000	5495	6000	6.56-19.7	851	40-75
Centrifugal	single stage	20-10.000	804	600	0.98-19.7	653	20-85
(vertical)	multistage	20-80.000	6004	700	0.98-19.7	500	25-90
	inline	20-I 2,000	705	500	6.56-19.7	500	20-80
	high speed	5-400	5807	2000	7.87-39.4	500	10-50
	sump	1 0-700	197	200	0.98-22.0		45-75
	multistage deep well	5-400	6004	2000	0.98-19.7	401	30-75
Axial	propeller	20-100.000	3 9	150	6.56	149	65-85
Turbine	regenerative	I-2000	2493	1500	6.56-8.20	248	55-85
Reciprocating	piston, plunger	1 o1 0,00 0	1.13x 10⁶	>50,000	12.1	554	65-85
	metering	O-10	1.70×10^{5}	50,000	15.1	572	2 0
	diaphragm	4-100	1.13x 10⁵	3500	12.1	500	2 0
Rotary	screw	I-2000	6.79 x 10⁴	3000	9.84	500	50-80
	gear	I-5000	11,155	500	9.84	653	50-80

 a 1 m³/min = 264 gpm, 1 m = 3.28 ft, I bar = 14.5 psi, °C = (°F - 32)/1.8.

TABLE 7.3. Operating Ranges of Some Commercial Vacuum Producing Equipment

Type of Pump	Operating Range (mm Hg)
Reciprocating piston	
I-stage	760-10
2-stage	760-1
Rotary piston oil-sealed	
I-stage	760–10 ⁻²
2-stage	$760-10^{-3}$
Centrifugal multistage (dry) liquid jet	760-200
Mercury Sprengel	$760-10^{-3}$
Water aspirator (18°C)	760-15
Two-lobe rotary blower (Roots type)	20-10-4
Turbomolecular	$10^{-1} - 10^{-10}$
Zeolite sorption (liquid nitrogen cooled)	760-10-3
Vapor jet pumps Steam ejector	
I-stage	760-100
2-stage	760-10
3-stage	760-l
4-stage	760-3x10-1
5-stage	760-5 × 10 ⁻²
Oil ejector (I-stage)	2-10-2
Diffusion-ejector	2–10 ^{–4}
Mercury diffusion with trap	1 6
1 -stage	10 ⁻ '-<10 ⁻⁶
P-stage	1-<10 ⁻ °
3-stage	10-<10 ⁻⁶
Oil diffusion	
I-stage	10 [−] '–5 × 10-s
4-stage fractionating (untrapped)	5 × 10 ⁻ '-10 ⁻⁹
Cstage fractionating (trapped)	$5 \times 10^{-1} - 10^{-12}$
Getter-ion (sputter-ion)	10_3-10_1
Sublimation (titanium)	10 [~] "-10 ⁻ '
Cryopumps (20 K)	1010
Cryosorption (15 K)	10~2-10-12

⁽Encyclopedia of Chemical Technology, Wiley-Interscience, New York, 1978-1984).

, fans. Figure 7.24 allows exploration of the effects of changes in specific speed or diameter on the efficiencies and other characteristics of fans. The mutual effects of changes in flow rate, pressure, speed, impeller diameter, and density are related by the "fan laws" of Table 7.5, which apply to all rotating propelling equipment.

COMPRESSORS

The several kinds of commercial compressors are identified in this classification:

- 1. Rotodynamic
 - a. Centrifugal (radial flow)
 - b. Axial flow
- 2. Positive displacement
 - a. Reciprocating piston
 - b. Rotary (screws, blades, lobes, etc.).

Sketches of these several types are shown in Figures 7.19 and 7.20 and their application ranges in Figures 7.20 and 7.21.

CENTRIFUGALS

The head-flow rate curve of a centrifugal compressor often has a maximum as shown on Figure 3.21, similar to the pump curve of

Figure 7.7(c). To the left the developed head increases with flow, but to the right the head decreases with increasing flow rate. At the peak the flow pulsates and the machine vibrates. This operating point is called the *surge limit* and is always identified by the manufacturer of the equipment, as shown on Figure 7.25 for those centrifugal and axial machines. Stable operation exists anywhere right of the surge limit. Another kind of flow limitation occurs when the velocity of the gas somewhere in the compressor approaches sonic velocity. The resulting shock waves restrict the flow; a slight increase in flow then causes a sharp decline in the developed pressure.

Table 7.6 shows as many as 12 stages in a single case. These machines are rated at either 10K or 12K ft/stage. The higher value corresponds to about 850 ft/sec impeller tip speed which is near the limit for structural reasons. The limitation of head/stage depends on



Figure 7.15. Effects of viscosity on performance of centrifugal pumps: (a) Hydraulic Institute correction chart for pumping liquids. (b) Typical performances of pumps when handling viscous liquids. The dashed lines on the chart on the left refer to a water pump that has a peak efficiency at 750 gpm and 100 ft head; on a liquid with viscosity 1000 SSU (220CS) the factors relative to water are efficiency 64%, capacity 95% and head 89% that of water at 120% normal capacity $(1.2Q_H)$.



Figure 7.15—(continued)

the nature of the gas and the temperature, as indicated on Figure 7.26. Maximum compression ratios of 3-4.5 per stage with a maximum of 8-12 per machine are commonly used. Discharge pressures as high as 3000-5000 psia can be developed by centrifugal compressors.

A specification form is included in Appendix B and as Table 4.4. Efficiency data are discussed in Section 7.6, Theory and Calculations of Gas Compression: Efficiency.

AXIAL FLOW COMPRESSORS

Figure 7.18(b) shows the axial flow compressor to possess a large number of blades attached to a rotating drum with stationary but adjustable blades mounted on the case. Typical operating characteristics are shown on Figure 7.22(a). These machines are suited particularly to large gas flow rates at maximum discharge pressures of 80-130 psia. Compression ratios commonly are 1.2-1.5 per stage and 5-6.5 per machine. Other details of range of applications are stated on Figure 7.20. According to Figure 7.21,



Figure 7.16. Recommended values of net positive suction head (NPSH) at various temperatures or vapor pressures: (a) NPSH of several types of pumps for handling water at various temperatures. (b) Correction of the cold water NPSH for vapor pressure. The maximum recommended correction is one-half of the cold water value. The line with arrows shows that for a liquid with 30psia vapor pressure at 100° F, the reduction in NPSH is 2.3 ft (data of Worthington International Inc.).

specific speeds of axial compressors are in the range of 1000-3000 or so.

Efficiencies are 8-10% higher than those of comparable centrifugal compressors.

RECIPROCATING COMPRESSORS

Reciprocating compressors are relatively low flow rate, high pressure machines. Pressures as high as 35,000–50,000 psi are



Figure 7.17. Operating points of centrifugal pumps under a variety of conditions. (a) Operating points with a particular pump characteristic and system curves corresponding to various amounts of flow throttling with a control valve. (b) Operating point with two identical pumps in parallel; each pump delivers one-half the flow and each has the same head. (c) Operating point with two identical pumps in series; each pump delivers one-half the head and each has the same flow.



Figure 7.18. Heavy-duty centrifugal, axial, and reciprocating compressors. (a) Section of a three-stage compressor provided with steam-sealed packing boxes (*DeLaval Steam Turbine* Co.). (b) An axial compressor (Clark Brothers Co.). (c) Double-acting, two-stage reciprocating compressor with water-cooled jacket and intercooler (*Ingersoll-Rand* Co.).











(c)

Viscosity in Seconds, Saybolt Universal (d)

4000

5000

6000

7000

Horsepower

Capacity

60 75 Pressure Ri**se ,%**

(b)

Gallons per minute

Efficiency, per cent

Brake horsepower

100% Speed

75 50

105

120

90

Figure 7.19. Some rotary positive displacement compressors (a) A two-lobe blower. (b) Performance of a two-lobe blower (*Roots-Connersville Co.*). (c) A screw pump with one power and two idle rotors (*Kristal and Annett*, 1940). (d) Performance of 3.5" screw pump handling oils at 1150 rpm against 325 psig (*Kristal and Annett*, 1940). (e) Principle of the liquid ring seal compressor (Nash Engineering Co.). (f) A sliding vane blower (Beach-Russ Co.).

15

30

45

2000

3000



Figure 7.20. Applications ranges of compressors and fans (Worthington): (a) Pressure-capacity ranges for air at 1 atm, 60°F, 0.075 lb/cuft. (b) Head-capacity ranges for all gases. Similar charts are given by Ludwig (1983, Vol. 1, p. 251) and Chemical Engineers Handbook (1984, p. 6.21).

developed with maximum compression ratios of 10/stage and any desired number of stages provided with intercoolers. Other data of application ranges are in Figure 7.20. The limitation on compression ratio sometimes is due to the limitations on discharge temperature which normally is kept below 300°F to prevent ignition of machine lubrications when oxidizing gases are being compressed, and to the fact that power requirements are proportional to the absolute temperature of the suction gas.

A two-stage double-acting compressor with water cooled cylinder jackets and intercooler is shown in Figure 7.18(c). Selected dimensional and performance data are in Table 7.7. Drives may be with steam cylinders, turbines, gas engines or electrical motors. A specification form is included in Appendix B. Efficiency data are discussed in Section 7.6, Theory and Calculations of Gas Compression: Temperature Rise, Compression Ratio, Volumetric Efficiency.

ROTARY COMPRESSORS

Four of the many varieties of these units are illustrated in Figure 7.19. Performances and comparisons of five types are given in Tables 7.8-7.9. All of these types also are commonly used as vacuum pumps when suction and discharge are interchanged.

Lobe *type units* operate at compression ratios up to 2 with efficiencies in the range of 80-95%. Typical relations between volumetric rate, power, speed, and pressure boost are shown in Figure 7.19(b).

Spiral screws usually run at 1800-3600 rpm. Their capacity ranges up to 12,000 CFM or more. Normal pressure boost is 3-20 psi, but special units can boost pressures by 60-100 psi. In vacuum service they can produce pressures as low as 2psia. Some other performance data are shown with Figure 7.19(d).

The sliding vane compressor can deliver pressures of 50 psig or



Figure 7.21. Operating ranges of single-stage pumps and compressors [*Balje*, Trans. ASME, J. Eng. Power. 84, 103 (1962)]. *Example:* atmospheric air at the rate of 100,000 SCFM is compressed to 80,000 ft lbf/ft (41.7 psig) at 12,000 rpm; calculated $N_s = 103$; in the radial flow region with about 80% efficiency, $D_s = 1.2$ -1.6, so that D = 2.9-3.9 ft.



Figure 7.22. Performances of dynamic compressors: (a) Axial compressor. (b) Centrifugal compressor. All quantities are expressed as percentages of those at the design condition which also is the condition of maximum efficiency (De Laval Engineering Handbook, *McGraw-Hill, New York, 1970*).

TABLE 7.4. Performance	Characteristics	of	Fans"
------------------------	-----------------	----	-------

	Qua (1000	ntity acfm)	Head	Opt.		Diameter (in.)				Deek
Description	Min	Max	Water	(fps)	q _{ad}	Min	Max	N _s	D _s	Eff.
Axial propeller	8	2 0	10	410	0.13	23	27	470	0.63	77
Axial propeller	2 0	90	8	360	0.12	27	72	500	0.60	80
Axial propeller	6	120	2.5	315	0.10	27	84	560	0.50	84
Radial air foil	6	100	2 2	250	0.45	18	90	190	0.85	88
Radial BC	3	35	18	260	0.63	18	90	100	1.35	78
Radial _{open} MH	2	27	18	275	0.55	18	66	97	1.45	56
Radial MH	2	27	18	250	0.55	18	66	86	1.53	71
Radial IS ,	2	27	18	250	0.55	18	66	86	1.53	66
Vane Bl flat	1	10	12	250	0.43	10	30	210	0.81	70
Vane FC	1	10	2	65	1.15	10	30	166	0.65	66

 ${}^{a}q_{ad} = 32.2H/V^{2}$, $N_{s} = NQ^{0.5}/V^{0.75}$ (specific speed), $D_{s} = DV^{0.25}/Q^{0.5}$ (specific diameter), where D = diameter (ft), H = head (ft), Q = suction flow rate (cfs), V = impeller tip speed (fps), and N = rotation speed (rpm)

TABLE 7.5. Fan Laws'

Ratio of -

cfm

ΗP

cfm

press

Fan Law

Number

b

с

а

1 а

2

(Evans. 1979).



2	a b c	ctm rpm H P	-	SIZC I/size size ²	× × ×	press" press" press	х	$1/\delta^{1/2}$ $1/\delta^{1/2}$ $1/\delta^{1/2}$
3	a b c	rpm press H P	-	1/size ³ 1/size ⁴ 1/size ⁴	× × x	cfm cfm ² cfm ³	x	1 δ δ
4	a b c	cfm press rpm	-	size ^{4/3} 1/size ^{4/3} 1/size ^{5/3}	x x x	HP ^{1/3} HP ^{2/3} HP ^{1/3}	X	1/δ ^{1/3} δ ^{1/3} 1/δ ^{1/3}
5	a b c	size rpm H P	-	cfm ^{1/2} 1/cfm ^{1/2} cfm	X X X	1/press ^{1/4} press ^{3/4} press	x	δ ^{1/4} 1/δ ^{3/4} 1
6	a b c	size press H P	~	cfm ^{1/3} cfm ^{2/3} cfm ^{5/3}	× X X	1/rpm ^{1/3} rpm ^{4/3} rpm ^{4/3}	X	1 δ δ
7	a b c	size cfm H P	-	press"' press ^{3/2} press ^{5/2}	X X X	1/rpm 1/rpm ² 1/rpm ²	x	1/δ ^{1/2} 1/δ ^{3/2} 1/δ ^{3/2}
8	a b C	size rpm press	-	1/HP ^{1/4} HP ^{3/4} HP	× × x	cfm ^{3/4} 1/cfm ^{5/4} 1/cfm	x	δ ^{1/4} 1/δ ^{3/4} 1
9	a b C	size rpm cfm		HP'" 1/HP ^{1/2} HP	x x x	1/press ^{3/4} press ^{5/4} I/press	X	δ ^{1/4} 1/δ ^{3/4} 1
10	a b	size cfm press	-	HP ^{1/5} HP ^{%,7} HP ^{2/5}	x x x	1/rpm ^{3/5} 1/rpm ^{4/3} rpm ^{4/5}	X	1/δ ^{1/5} 1/δ ^{3/5} δ ^{3/5}

Variables

Ratio

rpm

rpm²

rpm³

press"

Ratio

1

δ

δ

1/δ^{1/2}

Х

Х

х

х

х

х

Ratio

size³

size²

size

size²

Figure 7.23. Performances of fans with several kinds of controls (American Standard Co. Inc.). (a) A damper in the duct with constant-speed fan drive, (b) two-speed fan driver, (c) inlet vanes or inlet louvers with a constant-speed fan drive, (d) multiple-step variable-speed fan drive, and (e) hydraulic or electric coupling with constant-speed driver giving wide control over fan speed.

 $^{a}\delta = \rho/g_{c}$

For example, the pressure P varies as $D^2 N^2 \rho/g_c$ line 1(b), $Q^2(\rho/g_c)/D^4$ line 3(b), $\dot{P}^{2/3}(\rho/g_c)^{1/3}/D^{4/3}$ line 4(b), $Q^{2/3}N^{4/3}\rho/g_c$ line 6(b), P/Q line 8(c), and $\dot{P}^{2/5}N^{4/5}(\rho/g_c)^{3/5}$ line 10(c). (R.D. Madison, 1949).



(c)

(d)

100

Forwardly

Curved

Low

Low

Poor

Low

Poor

Poor

Small

Figure 7.24. Performances of fans with various-shaped blades (Green Fuel Economizer Co.): (a) Backward curved blades. (b) Straight radial blades. (c) Forward curved blades. (d) Comparison of characteristics of the several blade types (Sturtevant).

Frame	Normal Inlet Flow Range ^a (ft ³ /min)	Nominal Polytropic Head pee ⁻ Stage ^o p}	Nominal Polytropic Efficiency (η_ρ)	Nominal Maximum No. of Stages ^c	Speed at Nominal Polytropic Head/Stage
29M	500-8000	10,000	0. 76	10	11500
38M	6000-23.000	10,000/12,000	0. 77	9	8100
46M	20,000-35,000	10,000/12,000	0. 77	9	6400
60M	30. 000- 58. 000	10,000/12,000	0. 77	8	5000
70M	50. 000- 85. 000	10,000/12,000	0. 78	8	4100
88M	75. 000- 130. 000	10,000/12,000	0. 78	8	3300
103M	110,000160,000	10, 000	0. 78	7	2800
110M	140. 000- 190. 000	10, 000	0. 78	7	2600
25MB (H) (HH)	500-5000	12, 000	0. 76	12	11500
32MB (H) (HH)	5000-10.000	12, 000	0. 78	10	10200
38MB (H)	8000-23.000	10,000/12,000	0. 78	9	8100
46 MB	20,00035,000	10,000/12,000	0. 78	9	6400
60MB	30. 000- 58. 000	10,000/12,000	0. 78	8	5000
70 MB	50,000-85,000	10,000/12,000	0. 78	8	4100
88MB	75. 000- 130. 000	10,000/12,000	0. 78	8	3300

^a Maximum flow capacity is reduced in direct proportion to speed reduction. ${}^{b}_{C}$ Use either 10,000 or 12,000 ft for each impeller where this option is mentioned

^cAt reduced speed, impellers can be added. (Elliott Co.).



Figure 7.25. Efficiency and head coefficient q_{ad} as functions of specific speeds and specific diameters of various kinds of impellers *(Evans, 1979). Example:* An axial propeller has an efficiency of 70% at $N_s = 200$ and $D_s = 1.5$; and 85% at $N_s = 400$ and $D_s = 0.8$. See Table 7.4 for definitions of q_{ad} , N_s , and D_s .

pull a vacuum of 28in. of mercury. A two-stage unit can deliver 250psig. A generous supply of lubricant is needed for the sliding vanes. Table 7.9 shows that power requirements are favorable in comparison with other rotaries.

Liquid-her compressors produce an oil-free discharge of up to 125psig. The efficiency is relatively low, 50% or so, but high enough to make them superior to steam jet ejectors for vacuum service. The liquid absorbs the considerable heat of compression and must be circulated and cooled; a 200HP compressor requires 100 gpm of cooling water with a 10°F rise. When water vapor is objectionable in the compressed gas, other sealing liquids are used; for example, sulfuric acid for the compression of chlorine. Figure 7.19(e) shows the principle and Table 7.10 gives specifications of some commercial units.

7.6. THEORY AND CALCULATIONS OF GAS COMPRESSION

The main concern of this section is how to determine the work requirement and the effluent conditions of a compressor for which the inlet conditions and the outlet pressure are specified. Theoretical methods allow making such calculations for ideal and real gases and gas mixtures under isothermal and frictionless adiabatic (isentropic) conditions. In order that results for actual operation can be found it is **neceessary** to know the efficiency of the equipment. That depends on the construction of the machine, the mode of operation, and the nature of the gas being processed. In the last analysis such information comes from test work and its correlation by manufacturers and other authorities. Some data are cited in this section.

DIMENSIONLESS GROUPS

The theory of dimensionless groups of Section 7.2, Basic Relations, also applies to fans and compressors with rotating elements, for example, Eqs. (7.8)-(7.10) which relate flow rate, head, power, speed, density, and diameter. Equivalent information is embodied in Table 7.5. The concept of specific speed, Eqs. (7.11) and (7.12), also is pertinent. In Figures 7.21 and 7.25 it is the basis for identifying suitable operating ranges of various types of compressors.

IDEAL GASES

The ideal gas or a gas with an equation of state

$$PV = zRT \tag{7.18}$$

is a convenient basis of comparison of work requirements for real gases and sometimes yields an adequate approximation of these work requirements. Two limiting processes are isothermal and isentropic (frictionless adiabatic) flows. Changes in elevation and velocity heads are considered negligible here. With constant compressibility z the isothermal work is

w =
$$\int_{4}^{P_2} V dP = zRT \ln(P_2/P_1).$$
 (7.19)

Under isentropic conditions and with constant heat capacities, the pressure-volume relation is

$$PV^{k} = P_{1}V_{1}^{k} = \text{const.}$$
 (7.20)

where

$$k = C_p / C_v \tag{7.21}$$

TABLE 7.7. Some Sizes of One- and Two-Stage Reciprocating Compressors

(a) Horizontal, One-Stage, Belt-Driven

Diameter	Oferelas	Brake Air Prossure Up et Dated		Brake	Open	ings (in.)	
(in.)	(in.)	(cuft/min.)	rpm	(lb/sq in.)	Pressure	Inlet	Outlet
$7\frac{1}{2}$	6	106	310	80-100-125	15. 9- 17- 18	$2\frac{1}{2}$	2 ¹ / ₂
8 ¹ / ₂	9	170	300	80-100-125	25-27-29	3	3
10	10	250	285	80-100-125	36-38.5-41	$3\frac{1}{2}$	$3\frac{1}{2}$
11	12	350	270	80-100-125	51-57-60	_	4
8 ¹ / ₂	6	138	350	40-60	15-18.5		3
10	9	245	300	40-75	27-34	$3\frac{1}{2}$	31
11	10	312	285	40-75	34-43	4	4
13	12	495	270	40-75	54-70	5	5
12	9	350	300	20-45	30-42	4	4
13	10	435	285	30-45	42-52	6	6
15	12	660	270	30-50	59-74	7	7

(Worthington Corp.).

TABLE 7.7-(continued)

(b) Horizontal, One-Stage, Steam-Driven^a

Diameter, Steam Cylinder (in.)	Diameter, Air Cylinder, (in.)	Stroke (in.)	Displacement, (cuft/min)	rpm	Air Pressure, (Ib/sq in.)
7	$7\frac{1}{2}$	6	106	350	ao-loo-125
а	$8\frac{1}{2}$	9	170	300	80-100-125
9	10	10	250	285	80-100-125
10	11	12	350	270	80-100-125
7	8 ¹ / ₂	6	136	350	40-60
а	10	9	245	300	40-75 ^b
9	11	10	312	285	40-75 ^b
10	13	12	495	270	40-75 ^b
а	12	9	350	300	20–45 [°]
9	13	10	435	285	20–45 [°]
10	15	12	660	270	20–50 °

^aAll machines have piston-type steam valves. ^b110-lb steam necessary for maximum air pressure. ^c 125-lb steam necessary for maximum air pressure.

(Worthington Corp.).

(c) Horizontal, Two-Stage, Belt-Driven

Diameter			Cylinder	(in.)			Piston	
	Low	Pressure	High	Pressure	Stroke (in	n.) rpm	(cuft free air/	min)
		4		2 ¹ / ₈	4	500	28	
		6		2 ⁷ 8	6	350	6 5	
		а		3 <u>8</u>	а	300	133	
		10		4 7 8	10	275	241	

(Ingersoll-Rand Co.).

	Туре						
	Helical Screw	Spiral Axial	Straight Lobes	Sliding Vanes	Liquid Liner		
Configuration,				8	16		
features (male x female)	4 x 6	2 x 4	2 x 2	Blades	Sprockets		
Max displacement (cfm)	20,000	13,000	30,000	6,000	13,000		
Max diameter (in.)	2 5	16	la	33	4 8		
Min diameter (in.)	4	6	10	5	12		
Limiting tip speed (Mach)	0.30	0.12	0.05	0.05	0.06		
Normal tip speed (Mach)	0.24	0.09	0.04	0.04	0.05		
Max L/d, low pressure	1.62	2.50	2.50	3.00	1.1		
Normal L/d, high pressure	1 .00	1.50	1.50	2.00	1 .00		
V factor for volumetric efficiency	7	3	5	3	3		
X factor for displacement	0.0612	0.133	0.27	0.046	0.071		
Normal overall efficiency	75	70	68	7 2	50		
Normal mech. eff. at ±100 HP (%)	90	93	95	94	90		
Normal compression ratio R _c	2/3/4	3	1.7	2/3/4	5		
Normal blank-off R	6	5	5	7	9		
Displacement form-factor $A_{_{\!$	0.462	1.00	2.00	0.345	0.535		

TABLE 7.8. Summary of Rotary Compressor Performance Data

(Evans, 1979).

TABLE 7.9. Five F	Rotary Comp	ressors for a	Common	Service
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	Туре						
	Helical Screw	Spiral Axíal	Straight Lobes	Sliding Vanes	Liquid Liner		
Suction loss $ heta_i$	9.35	1.32	0.89	0.90	1.40		
Discharge loss $ heta_{oldsymbol{ ho}}$	7.35	1.04	0.70	0.70	1.10		
Intrinsic corr, B	1.185	1.023	1.016	1.016	1.025		
Adiabatic eff. η_{ad}	85.6	97.7	98.5	98.5	97.9		
Slippage <i>W_s</i> (%)	28.5	18.6	11.8	11.8	3.0		
Slip eff. $\eta_s(\%)$	71.5	83.4	88.2	88.2	97.0		
Thermal eff η_t (%)	89.2	93.7	95.8	95.5	42.5		
Volumetric eff. Evr	88.0	85.7	89.1	89.9	96.6		
Displacement (cfm)	14,700	11,650	11,220	11,120	10,370		
Rotor dia. (in.)	26.6	26.2	27.0	65.0	45.5		
Commercial size, d x <u>L</u>	25 x 25	22x33	22 x 33	46 x 92 <i>°</i>	43×48^{b}		
Speed (rpm)	3,500	1,250	593	284	378		
Motor (HP)	1,100	800	750	750	1,400		
Service factor	1.09	1.11	1.10	1.12	1.10		
Discharge temp °F	309	270	262	263	120		

Twin 32.5 x 65 or triplet 26.5 x 33 (667 rpm) are more realistic. Twin 32 x 32 (613 rpm) alternate where L = d

(Evans, 1979).

is the ratio of heat capacities at constant pressure and constant volume $% \left({{{\mathbf{r}}_{\mathbf{r}}}_{\mathbf{r}}} \right)$ and

$$C_v = R - C_p. \tag{7.22}$$

A related expression of some utility is

$$T_2/T_1 = (P_2/P_1)^{(k-1)/k}.$$
 (7.23)

Since k ordinarily is a fairly strong function of the temperature, a suitable average value must **be** used in Eq. (7.20) and related ones. Under adiabatic conditions the flow work may be written as

$$W = H_2 - H_1 = \int_{\mu}^{P_2} V \, dP. \tag{7.24}$$

Upon substitution of Eq. (7.20) into Eq. (7.24) and integration, the isentropic work becomes

$$W_{s} = H_{2} - H_{1} = P_{1}^{1/k} V_{1} \int_{P_{1}}^{P_{2}} dP / P^{1/k}$$

= $\left(\frac{k}{k-1}\right) z_{1} R T_{1} \left[\left(\frac{P_{2}}{P_{1}}\right)^{(k-1)/k} - 1 \right].$ (7.25)

TABLE 7.10. Specifications of Liquid Liner Compressors

Compressor (size)	Pressure (psi)	Capacity (cuft/min)	Motor (HP)	Speed (rpm)
	5	1020	4 0	(
K - 6	10	990	60	570
	15	870	75	1
	(²⁰	650	100	t
621	{	26	75	3500
1251	35	120	40	1750
1256		440	100	1750
621	(2 3	10	3500
1251	{ 80	110	50	1750
1256	ł	410	150	1750

(Nash Engineering Co.).

In multistage centrifugal compression it is justifiable to take the average of the inlet and outlet compressibilities so that the work becomes

$$W_s = H_2 - H_1 = \left(\frac{k}{k-1}\right) \left(\frac{z_1 + z_2}{2}\right) RT_1 \left[\left(\frac{P_2}{P_1}\right)^{(k-1)/k} - 1\right]. (7.26)$$

When friction is present, the problem is handled with *empirical* efficiency factors. The isentropic compression efficiency is defined as

$$\eta_s = \frac{\text{isentropic work or enthalpy change}}{\text{actually required work or enthalpy change}}$$
(7.27)

Accordingly,

$$W = AH = W_s/\eta_s = (\Delta H)_s/\eta_s.$$
(7.28)

When no other information is available about the process gas, it is justifiable to find the temperature rise from

$$\Delta T = (\Delta T)_s / \eta_s \tag{7.29}$$

so that

$$T_2 = T_1 (1 + (1/\eta_s) [(P_2/P_1)^{(k-1)/k} \quad 1].$$
(7.30)

A case with variable heat capacity is worked out in Example 7.5.

For mixtures, the heat capacity to use is the sum of the mol

EXAMPLE 7.4

Gas Compression, Isentropic and True Final Temperatures With k = 1.4, $P_2/P_1 = 3$ and $\eta_s = 0.71$; the final temperatures are $(T_2)_s = 1.369T_1$ and $T_2 = 1.519T_1$ with Eqs. (7.24) and (7.31).

156 FLUID TRANSPORT EQUIPMENT

fraction weighted heat capacities of the pure components,

$$C_p = \sum x_i C_{pi}. \tag{7.31}$$

REAL PROCESSES AND GASES

Compression in reciprocating and centrifugal compressors is essentially adiabatic but it is not frictionless. The pressure-volume behavior in such equipment often conforms closely to the equation

$$PV'' = P_1 V_1^n = \text{const}$$
 (7.32)

Such a process is called polytropic. The equation is analogous to the isentropic equation (7.20) but the polytropic exponent n is different from the heat capacity ratio k.

Polytropic exponents are deduced from PV measurements on the machine in question. With reciprocating machines, the PV data are recorded directly with engine indicators. With rotary machines other kinds of instruments are used. Such test measurements usually are made with air.

Work in polytropic compression of a gas with equation of state PV = zRT is entirely analogous to Eq. (7.26). The hydrodynamic work or the work absorbed by the gas during the compression is

$$W_{hd} = \int_{P_1}^{P_2} V \, dP = \left(\frac{n}{n-1}\right) z_1 R T_1 \left[\left(\frac{P_2}{P_1}\right)^{(n-1)/n} - 1 \right]. \tag{7.33}$$

Manufacturers usually characterize their compressors by their polytropic efficiencies which are defined by

$$\eta_p = \left(\frac{n}{n-1}\right) \left/ \left(\frac{k}{k-1}\right) = \frac{n(k-1)}{k(n-1)}.$$
(7.34)

The polytropic work done on the gas is the ratio of Eqs. (7.33) and (7.34) and comprises the actual mechanical work done on the gas:

$$W_{p} = W_{hd}/\eta_{p} = \left(\frac{k}{k-1}\right) z_{1} R T_{1} \left[\left(\frac{P_{2}}{P_{1}}\right)^{(n-1)/n} - 1\right]$$
(7.35)

Losses in seals and bearings of the compressor are in addition to W_p ; they may amount to 1–3% of the polytropic work, depending on the machine.

The value of the polytropic exponent is deduced from Eq. (7.34) as

$$n = \frac{k\eta_p}{1 - k(1 - \eta_p)}.$$
(7.36)

The isentropic efficiency is

$$\eta_{s} = \frac{\text{isentropic work [Eq. (7.25)]}}{\text{actual work [Eq. (7.35)]}}$$
(7.37)

$$=\frac{(P_2/P_1)^{(k-1)/k}-1}{(P_2/P_1)^{(n-1)/n}-1}$$
(7.38)

$$=\frac{(P_2/P_1)^{(k-1)/k}-1}{(P_2/P_1)^{(k-1)/k\eta_p}-1}$$
(7.39)

The last version is obtained with the aid of Eq. (7.34) and relates the isentropic and polytropic efficiencies directly. Figure 7.27(b) is a plot of Eq. (7.39). Example 7.6 is an exercise in the relations between the two kinds of efficiencies.





$$H = \frac{Ku^2}{32.2} \text{ ft / stage}$$

$$K = 0.50 - 0.65, \text{ empirical coefficient}$$

$$U = 600.900 \text{ ft / sec, impeller peripheral speed}$$

$$H = 10,000 \text{ with average values } K = 0.55 \text{ and } U = 765 \text{ fl / sec}$$

(c)

Figure 7.26. Several ways of estimating allowable polytropic head per stage of a multistage centrifugal compressor. (a) Single-stage head as a function of k, molecular weight, and temperature (Elliott Co.). (b) Single-stage head as a function of the nature of the gas (NGPSA Handbook, *Gas Processors Assn, Tulsa, OK, 1972*), obtained by dividing the total head of the compressor by number of stages. $H = Ku^2/32.2$ ft/stage, K = 0.50-0.65, empirical coefficient, u = 600-900 ft/sec, impeller peripheral speed, and H = 10,000 with average values K = 0.55 and u = 765 ft/sec. (c) An equation and parameters for estimation of head.

EXAMPLE 7.5

Compression Work with Variable Heat Capacity Hydrogen sulfide heat capacity is given by

$$C_p = 7.629 + 3.431(E - 4)T + 5.809(E - 6)T^2$$

- 2.81(E - 9)T³, cal/g mol,

with T in K. The gas is to be compressed from 100°F (310.9 K) and 14.7 psia to 64.7 psia.

Assuming the heat capacity to be independent of pressure in this low range, the isentropic condition is

AS =
$$\int_{T_1}^{T_2} (C_p/T) dT - R \ln(P_2/P_1)$$

= $\int_{310.9}^{T_2} (C_p/T) dT - 1.987 \ln(64.7/14.7) = 0$

By trial, with a root-solving program,

 $T_2 = 441.1$ K, 334.4°F (compared with 345°F from Example 7.7).

WORK ON NONIDEAL GASES

The methods discussed thus far neglect the effect of pressure on enthalpy, entropy, and heat capacity. Although efficiencies often are not known well enough to justify highly refined calculations, they may be worth doing in order to isolate the uncertainties of a design. Compressibility factors are given for example by Figure 7.29. Efficiencies must be known or estimated.

Thermodynamic Diagram Method. When a thermodynamic diagram is available for the substance or mixture in question, the flow work can be found from the enthalpy change,

The procedure is illustrated in Example 7.7 and consists of these steps:

- 1. Proceed along the line of constant entropy from the initial condition to the final pressure P_2 and enthalpy $(H_2)_s$.
- 2. Evaluate the isentropic enthalpy change (AH), = $(H_2)_s H_1$.
- 3. Find the actual enthalpy change as

$$\Delta H = (\Delta H)_s / \eta_s \tag{7.41}$$

and the final enthalpy as

$$H_2 = H_1 + (\Delta H)_s / \eta_s. \tag{7.42}$$

4. At the final condition (P_2, H_2) read off any other desired properties such as temperature, entropy or specific volume.

Thermodynamic diagrams are known for light hydrocarbons, refrigerants, natural gas mixtures, air, and a few other common substances. Unless a substance or mixture has very many applications, it is not worthwhile to construct a thermodynamic diagram for compression calculations but to use other equivalent methods.

The isentropic enthalpy change becomes

AH,
$$= \int_{310.1}^{441.1} C_p \ dT = 1098.1 \ \text{cal/g mol}$$

 $\rightarrow 1098.1(1.8)/34.08 = 58.0 \ \text{Btu/lb},$

compared with 59.0 from Example 7.7. The integration is performed with Simpson's rule on a calculator.

The actual final temperature will vary with the isentropic efficiency. It is found by trial from the equation

$$1098.1/\eta_s = \frac{T_2}{I_{1098.1}} C_p \, dT.$$

Some values are

General Method. The effects of composition of mixtures and of pressure on key properties such as enthalpy and entropy are deduced from PVT equations of state. This process is described in books on thermodynamics, for example, Reid, Prausnitz, and Sherwood (Properties of Liquids and Gases, McGraw-Hill, New York, 1977) and Walas (Phase Equilibria in Chemical Engineering, Butterworths, Stoneham, MA, 1985). Only the simplest correlations of these effects will be utilized here for illustration.

For ideal gases with heat capacities dependent on temperature, the procedure requires the isentropic final temperature to be found by trial from

AS=
$$\int_{T_1}^{T_{2s}} (C_p/T) dT - R \ln(P_2/P_1) \to 0,$$
 (7.43)

and then the isentropic enthalpy change from

$$AH = \frac{T_{2s}}{I_{T_2}} C_p \, dT. \tag{7.44}$$

The final temperature T_2 is found by trial after applying a known isentropic efficiency,

$$(\Delta H)_s / \eta_s = \int_{T_1}^{T_2} C_p \ dT. \tag{7.45}$$

The fact that heat capacities usually are represented by empirical polynomials of the third or fourth degree in temperature accounts for the necessity of solutions of equations by trial.

Example 7.5 applies this method and checks roughly the calculations of Example 7.7 with the thermodynamic diagram of this substance. The pressures are relatively low and are not expected to generate any appreciable nonideality.

This method of calculation is applied to mixtures by taking a mol fraction weighted heat capacity of the mixture,

$$C_p = \sum x_i C_{pi}.\tag{7.46}$$

When the pressure range is high or the behavior of the gas is



Figure 7.27. Efficiencies of centrifugal and reciprocating compressors. (a) Polytropic efficiencies of centrifugal compressors as a function of suction volume and compression ratio (*Clark Brothers* Co.). (b) Relation between isentropic and polytropic efficiencies, Eqs. (7.22) (7.23). (c) Isentropic efficiencies of reciprocating compressors (De Laval Handbook, *McGraw-Hill*, New York, 1970). Multiply by 0.95 for motor drive. Gas engines require 7000-8000 Btu/HP.

EXAMPLE 7.6 Polytropic and Isentropic Efficiencies

Take $\eta_p = 0.75$, k = 1.4, and $P_2/P_1 = 3$. From Eq. (7.39), n = 1.6154and $\eta_s = 0.7095$. With Figure 7.27(b), $\phi = 3^{0.2857} = 1.3687$, $\eta_s = 0.945\eta_p = 0.709$. The agreement is close. nonideal for any other reason, the isentropic condition becomes

$$AS = \int_{T_1}^{T_{2s}} (C'_p/T) \ dT \ -R \ln(P_2/P_1) + \Delta S'_1 - \Delta S'_2 \to 0.$$
(7.47)

After the final isentropic temperature T_{2s} has been found by trial, the isentropic enthalpy change is obtained from

(AH), =
$$\int_{T_1}^{T_{2r}} C'_p dT + AH; -\Delta H'_{2s}.$$
 (7.48)



In terms of a known isentropic efficiency the final temperature T_2

In terms of a known isentropic efficiency the final temperature I_2 then is found by trial from

7.6. THEORY AND CALCULATIONS OF GAS COMPRESSION 159

$$(\Delta H)_{s}/\eta_{s} = \int_{T_{1}}^{T_{2}} C'_{p} dT + \Delta H'_{1} - \Delta H'_{2}.$$
(7.49)

In these equations the heat capacity C'_p is that of the ideal gas state or that of the real gas near zero or atmospheric pressure. The residual properties AS; and $\Delta H'_1$ are evaluated at (P_1, T_1) and $\Delta S'_2$ and $\Delta H'_2$ at (P_2, T_2) . Figure 7.28 gives them as functions of reduced temperature T/T_c and reduced pressure P/P_c . More accurate methods and charts for finding residual properties from appropriate equations of state are presented in the cited books of Reid et al. (1977) and Walas (1985).

For mixtures, pseudocritical properties are used for the evaluation of the reduced properties. For use with Figure 7.28, Kay's rules are applicable, namely,

$$(P_c)_{\min} = \sum x_i P_{ci},\tag{7.50}$$

$$(T_c)_{\min} = \sum x_i T_{ci}, \tag{7.51}$$

but many equations of state employ particular combining rules.

Example 7.8 compares a solution by this method with the assumption of ideal behavior.

EFFICIENCY

The efficiencies of fluid handling equipment such as fans and compressors are empirically derived quantities. Each manufacturer will supply either an efficiency or a statement of power requirement for a specified performance. Some general rules have been devised for ranges in which efficiencies of some classes equipment usually fall. Figure 7.27 gives such estimates for reciprocating compressors. Fan efficiencies can be deduced from the power-head curves of Figure 7.24. Power consumption or efficiencies of rotary and reciprocating machines are shown in Tables 7.7, 7.8, and 7.9.

Polytropic efficiencies are obtained from measurements of power consumption of test equipment. They are essentially independent of the nature of the gas. As the data of Figure 7.27 indicate, however, they are somewhat dependent on the suction volumetric rate, particularly at low values, and on the compression ratio. Polytropic efficiencies of some large centrifugal compressors are listed in Table 7.6. These data are used in Example 7.9 in the selection of a machine for a specified duty.

The most nearly correct methods of Section 7.6.4 require knowledge of isentropic efficiencies which are obtainable from the polytropic values. For a given polytropic efficiency, which is independent of the nature of the gas, the isentropic value is obtained with Eq. (7.39) or Figure 7.27(b). Since the heat capacity is involved in this transformation, the isentropic efficiency depends on the nature of the substance and to some extent on the temperature also.

TEMPERATURE RISE, COMPRESSION RATIO, VOLUMETRIC EFFICIENCY

The isentropic temperature in terms of compression ratio is given for ideal gases by

$$(T_2)_s = T_1 (P_2 / P_1)^{(k-1)/k}.$$
(7.52)

For polytropic compression the final temperature is given directly by

$$T_2 = T_1 (P_2 / P_1)^{(n-1)/n} \tag{7.53}$$

Figure 7.28. Residual entropy and enthalpy as functions of reduced properties. (a) Residual entropy. (b) Residual enthalpy. *Drawn by* Smith and Van Ness (Introduction to Chemical Engineering Thermodynamics, McGraw-Hill, New York, 1959) from data of Lydersen et al. For illustrative purposes primarily; see text for other sources.]

EXAMPLE 7.7

Finding Work of Compression with a Thermodynamic Chart Hydrogen sulfide is to be compressed from 100°F and atmospheric pressure to 50 nsig. The isentropic efficiency is 0.70. A pressure-enthalpy chart is taken from Starling (*Fluid Thermodynamic Properties for Light Petroleum Systems*, Gulf, Houston, TX, 1973). The work and the complete thermodynamic conditions for the process will be found.

The path followed by the calculation is 1-2-3 on the sketch. The initial enthalpy is -86 Btu/lb. Proceed along the isentrop S = 1.453 to the final pressure, 64.7 psia, and enthalpy $H_2 = -27$. The isentropic enthalpy change is

 $AH_{,} = -27 - (-86) = 59 Btu/lb.$

The true enthalpy change is

AH = 59/0.70 = 84.3.

The final enthalpy is

$$H_3 = -86 + 84.3 = -1.7$$

Other conditions at points 2 and 3 are shown on the sketch. The work is

 $\dot{W} = AH = 84.3 \text{ Btu/lb}$ $\rightarrow 84.3/2.545 = 33.1 \text{ HP hr}/(1000 \text{ lb}).$

EXAMPLE 7.8

Compression Work on a Nonideal Gas

Hydrogen sulfide at 450K and 15 atm is to be compressed to 66 atm. The isentropic final temperature and the isentropic enthalpy change will be found with the aid of Figure 7.28 for the residual properties.

The critical properties are $T_c = 373.2$ K and $P_c = 88.2$ atm. The heat capacity is stated in Example 7.5:

$$T_{r1} = 4501373.2 = 1.21,$$

$$P_{r1} = 15/88.2 = 0.17,$$

$$P_{r2} = 66/88.2 = 0.75,$$

$$\therefore AS; = 0.15,$$

$$AH; = 0.2(373.2) = 75.0,$$

$$AS = \frac{T_2}{L_{450}} \frac{Cp}{T} dT - 1.987 \ln \frac{66}{15} + 0.15 - \Delta S'_2 \stackrel{?}{=} 0,$$
(1)
(2)

AH,
$$= \int_{450}^{7_2} Cp \ dT + 75.0 - AH;$$
 (2)

1. Assume a value of T_2 .

2. Evaluate T_{r_2} and AS;

- 3. Integrate Eq. (1) numerically and note the righthand side.
- 4. Continue with trial values of T_2 until AS = 0.
- 5. Find $\Delta H'_2$ and finally evaluate AH,.

Two trials are shown.

T ₂	$\Delta H_{\rm s}$
600	1487 7
626.6	1487

 $\frac{1}{100} = \frac{11.9}{5\pi 1.9}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{3}$ $\frac{1}{5\pi 1.9}$ $\frac{1}{5\pi 1.9}$ $\frac{1}{2}$ $\frac{1}$

345

440 FF

When the residual properties are neglected,

 $T_2 = 623.33$ K (compared with 626.6 real), AH, = 1569.5 (compared with 1487.7 real).

Real temperature rise:

With $\eta_s = 0.75$, the enthalpy change is 1487.7/0.75 and the enthalpy balance is rearranged to

$$\Delta = -\frac{1487.7}{0.75} + \int_{450}^{T_2} Cp^* \, dT + 75 - \Delta H_2' \stackrel{?}{=} 0$$

Trial T₂	T,	AH;	rhs
680	1.82	109	+91.7
670. 79	1.80	112	- 0. 021
670.80	1.80		+0.075

 $\therefore T_2 = 670.79$ K

For ideal gas

$$\Delta = -\frac{1569.5}{0.75} + \int_{450}^{T_2} Cp^* \, dT \xrightarrow{?} 0$$

By trial:

$$T_2 = 670.49 \text{ K}$$

Nonideality is slight in this example.

EXAMPLE 7.9 Selection of a Centrifugal Compressor

A hydrocarbon mixture with molecular weight 44.23 is raised from 41°F and 20.1 psia to 100.5 psia at the rate of 2400 lb mol/hr. Its specific heat ratio is k = 1.135 and its inlet and outlet compressibilities are estimated as $z_1 = 0.97$ and $z_2 = 0.93$. A size of compressor will be selected from Table 7.6 and its expected performance will be calculated:

2400 lb mol/hr = 1769 lb/min, 10,260 cfm

From Table 7.6, the smallest compressor for this gas rate is # 38M. Its characteristics are

$\dot{N} = 8100$ rpm at 10–12 K ft/stage $\eta_p = 0.77$



Accordingly,

$$\frac{n-1}{n} = \frac{k-1}{k\eta_p} = \frac{0.135}{1.135(0.77)} = 0.1545$$

Using Eq. (7.35) for the polytropic head,

$$H_p = \left(\frac{Z_1 + Z_2}{2}\right) \left(\frac{k}{k-1}\right) R T_1 \left[\left(\frac{P_2}{P_1}\right)^{(n-1)/n} - 1 \right]$$

= 0.95 $\left(\frac{1.135}{0.135}\right) \left(\frac{1544}{44.23}\right) (501) [5^{0.1545} - 1]$
= 39430 ft.

From Figure 7.26(a), the max head per stage is 9700, and from Figure 7.26(b) the min number of stages is about 4.5. Accordingly, use five stages with standard 10,000 ft/stage impellers. The required speed with the data of Table 7.6 is

speed =
$$8100\sqrt{39430}/10,000(5)$$
 = 7190 rpm.
Power absorbed by the gas is
 $\dot{P}_{gas} = \frac{\dot{m}H_p}{33,000\eta_p} = \frac{1769(39,430)}{33,000(0.77)} = 2745 \text{ HP}.$

Friction losses $\cong 3\%$ max;

: total power input = 2745/0.97 = 2830 HP max.

or alternately in terms of the isentropic efficiency by

$$(\Delta T)_{\text{actual}} = T_2 \quad T_1 = (\Delta T)_{\text{isentropic}} / \eta_s$$
 (7.54)

so that

$$T_2 = T_1 + (\Delta T)_s / \eta_s = T_1 \{ 1 + (1/\eta_s) [(P_2/P_1)^{(k-1)/k} - 1] \}.$$
(7.55)

The final temperature is read off directly from a thermodynamic diagram when that method is used for the compression calculation, as in Example 7.7. A temperature calculation is made in Example 7.10. Such determinations also are made by the general method for **nonideal** gases and mixtures as in Example 7.8 and for ideal gases in Example 7.4.

Compression Ratio. In order to save on equipment cost, it is desirable to use as few stages of compression as possible. As a rule, the compression ratio is limited by a practical desirability to keep outlet temperatures below 300°F or so to minimize the possibility of ignition of machine lubricants, as well as the effect that power requirement goes up as outlet temperature goes up. Typical compression ratios of reciprocating equipment are:

Large pipeline compressors	1.2 –2.0
Process compressors	1.5-4.0
Small units	up to 6.0

For minimum equipment cost, the work requirement should be the same for each stage. For ideal gases with no friction losses between stages, this implies equal compression ratios. With n stages, accordingly, the compression ratio of each stage is

$$P_{i+1}/P_i = (P_n/P_1)^{1/n}.$$
(7.56)

Example 7.11 works out a case involving a nonideal gas and interstage pressure losses.

In centrifugal compressors with all stages in the same shell, the allowable head rise per stage is stated in Table 7.6 or correlated in Figure 7.26. Example 7.9 utilizes these data.

Volumetric Efficiency. For practical reasons, the gas is not completely discharged from a cylinder at each stroke of a reciprocating machine. The clearance of a cylinder is filled with compressed gas which reexpands isentropically on the return stroke. Accordingly, the gas handling capacity of the cylinder is less than the product of the cross section by the length of the stroke. The volumetric efficiency is

$$n_v = \frac{\text{suction gas volume}}{\text{cylinder displacement}}$$

= 1 - f_c[(P_2/P_1)^{1/k} - 1], (7.57)

where

$$f_c = \frac{\text{clearance volume}}{\text{cylinder displacement volume}}$$

For a required volumetric suction rate Q (cfm), the required product of cross section **A**, (sqft), stroke length L_s (ft), and speed N (rpm) is given by

$$A_s L_s N = Q/\eta_v. \tag{7.58}$$
EXAMPLE 7.10
Polytropic and Isentropic Temperatures
Take
$$k = 1.4$$
, $(P_2/P_1) = 3$, and $\eta_p = 0.75$. From Eq. (7.34),
 $(n-1)/n = (k-1)/k\eta_p = 0.3810$

and from Eq. (7.39)

$$n_{s} = \frac{3^{0.2857} - 1}{3^{0.3810} - 1} = 0.7094$$

so that from Eq. (7.53),

$$T_2/T_1 = 3^{0.3810} = 1.5198$$
, isentropic

and from Eq. (7.54),

 $T_2/T_1 = 1 + (1/0.7094)(3^{0.2857} - 1) = 1.5197$, polytropic.

7.7. EJECTOR AND VACUUM SYSTEMS

Application ranges of the various kinds of devices for maintenance of subatmospheric pressures in process equipment are shown in Table 7.3. The use of mechanical pumps--compressors in reverse for such purposes is mentioned earlier in this chapter. Pressures also can be reduced by the action of flowing fluids. For instance, water jets at 40psig will sustain pressures of 0.5–2.0 psia. For intermediate pressure ranges, down to 0.1 Torr or so, steam jet ejectors are widely favored. They have no moving parts, are quiet, easily installed, simple, and moderately economical to operate, and readily adaptable to handling corrosive vapor mixtures. A specification form is in Appendix B.

EJECTOR ARRANGEMENTS

Several ejectors are used in parallel when the load is variable or because the process system gradually loses tightness between maintenance shutdowns-then some of the units in parallel are cut in or out as needed.

Multistage units in series are needed for low pressures. Sketches are shown in Figure 7.30 of several series arrangements. In Figure 7.30(a), the first stage drives the process vapors, and the second stage drives the mixture of those vapors with the motive steam of the first stage. The other two arrangements employ interstage condensers for the sake of steam economy in subsequent stages. In contact (barometric) condensers the steam and other



Figure 7.29. Compressibility factors, z = PV/RT, of gases. Used for the solution of Example 7.11. $P_R = P/P_c$, $T_R = TIT_c$, and $V_{r'} = P_c V/RT_c$.



Figure 7.30. Arrangements of two-stage ejectors with condensers. (a) Identification of the parts of a two-stage ejector (*Croll-Reynolds* $C_{0,L}$. (b) A two-stage ejector with interstage barometric condenser (Elliot Co.). (c) A two-stage ejector with surface condensers interstage and terminal (Elliot Co.).

164 FLUID TRANSPORT EQUIPMENT

condensables are removed with a cold water spray. The tail pipes of the condensers are sealed with a 34ft leg into a sump, or with a condensate pump operating under vacuum. Surface condensers permit recovery of valuable or contaminating condensates or steam condensate for return as boiler feed. They are more expensive than barometrics, and their design is more complex than that of other kinds of condensers because of the large amounts of noncondensables that are present.

As many as six stages are represented on Figure 7.30, combined with interstage condensers in several ways. Barometric condensers are feasible only if the temperature of the water is below its bubblepoint at the prevailing pressure in a particular stage. Common practice requires the temperature to be about 5°F below the bubblepoint. Example 7.13 examines the feasibility of installing intercondensers in that process.

AIR LEAKAGE

The size of ejector and its steam consumption depend on the rate at which gases must be removed from the process. A basic portion of such gases is the air leakage from the atmosphere into the system.

Theoretically, the leakage rate of air through small openings, if they can be regarded as orifices or short nozzles, is constant at vessel pressures below about 53% of atmospheric pressure. However, the openings appear to behave more nearly as conduits with relatively large ratios of lengths to diameters. Accordingly sonic flow is approached only at the low pressure end, and the air mass inleakage rate is determined by that linear velocity and the low density prevailing at the vessel pressure. The content of other gases in the evacuated vessel is determined by each individual process. The content of condensables can be reduced by interposing a refrigerated condenser between process and vacuum pump.

Standards have been developed by the Heat Exchange Institute for rates of air leakage into commercially tight systems. Their chart is represented by the equation

$$n = kV^{2/3} \tag{7.59}$$

where m is in lb/hr, V is the volume of the system in cuft, and the

EXAMPLE 7.11

Three-Stage Compression with Intercooling and Pressure Loss between Stages

Ethylene is to be compressed from 5 to 75atm in three stages. Temperature to the first stage is 60°F, those to the other stages are 100°F. Pressure loss between stages is 0.34 atm (5 psi). Isentropic efficiency of each stage is 0.87. Compressibilities at the inlets to the

stages are estimated from Figure 7.29 under the assumption of equal compression ratios as $z_0 = 0.98$, $z_1 = 0.93$, and $z_2 = 0.83$. The interstage pressures will be determined on the basis of equal power load in each stage. The estimated compressibilities can be corrected after the pressures have been found, but usually this is not found necessary. $\mathbf{k} = C_p/C_v = 1.228$ and $(\mathbf{k} - 1)/k = 0.1857$.



1

	1	2				
P 1	1+2	2+3				
12	27.50	28. 31				
12.5	29.85	28.94				
13.0	32.29	29.56				
12. 25						



TABLE 7.11	. Estimated Air Leakages Through Connections,
	Valves, Stuffing Boxes Etc. of Process
	Equipment ^a

Type Fitting	Estimated Average Air Leakage (lb/hr)
Screwed connections in sizes up to 2 in.	0.1
Screwed connections in sizes above 2 in.	0.2
Flanged connections in sizes up to 6 in.	0.5
Flanged connections in sizes 6 in. to 24 in. including manholes	0.8
Flanged connections in sizes 24 in. to 6 ft	1.1
Flanged connections in sizes above 6ft	2.0
Packed valves up to $\frac{1}{2}$ in. stem diameter	0.5
Packed valves above $\frac{1}{2}$ in. stem diameter	1.0
Lubricated plug valves	0.1
Petcocks	0.2
Sight glasses	1.0
Gage glasses including gage cocks	2.0
Liquid sealed stuffing box for shaft of agitators, pumps, etc. (per in. shaft diameter)	0.3
Ordinary stuffing box (per in. of diameter)	1.5
Safety valves and vacuum breakers (per in. of nominal size	1.0

⁹For conservative practice, these leakages may be taken as supplementary to those from Eq. (7.59). Other practices allow 5 lb/hr for each agitator stuffing box of standard design; special high vacuum mechanical seals with good maintenance can reduce this rate to I-2 lb/hr.

[From C.D. Jackson, Chem. Eng. Prog. 44, 347 (1948)].

coefficient is a function of the process pressure as follows:

Pressure (Tor	r) >90	20-90	3-20	I - 3	<1
k	0.194	0.146	0.0825	0.0508	0.0254

For each agitator with a standard stuffing box, 5 lb/hr of air leakage is added. Use of special vacuum mechanical seals can reduce this allowance to 1-2 lb/hr.

For a conservative design, the rate from Eq. (7.59) may be supplemented with values based on Table 7.11. Common practice is to provide oversize ejectors, capable of handling perhaps twice the standard rates of the Heat Exchange Institute.

Other Gases. The gas leakage rate correlations cited are based on air at 70°F. For other conditions, corrections are applied to evaluate an effective air rate. The factor for molecular weight M is

$$f_M = 0.375 \ln(M/2)$$
 (7.60)

and those for temperature T in °F of predominantly air or predominantly steam are

$$f_A = 1 - 0.00024(T - 70), \text{ for air,}$$
(7.61)

$$f_S = 1 - 0.00033(T - 70)$$
, for steam. (7.62)

An effective or equivalent air rate is found in Example 7.12.

EXAMPLE 7.12 Equivalent Air Rate

Suction gases are at the rate of 120 lb/hr at 300°F and have a molecular weight of 90. The temperature factor is not known as a function of molecular weight so the value for air will be used. Using Eqs. (7.61) and (7.62),

 $m = 120(0.375) \ln(90/2)[1 - 0.00024(300-70)]$ = 161.8 lb/hr equivalent air.

STEAM CONSUMPTION

The most commonly used steam is 100 psig with 10-15" superheat, the latter characteristic in order to avoid the erosive effect of liquids on the throats of the ejectors. In Figure 7.31 the steam consumptions are given as lb of motive steam per lb of equivalent air to the first stage. Corrections are shown for steam pressures other than 100 psig. When some portion of the initial suction gas is condensable, downward corrections to these rates are to be made for those ejector assemblies that have intercondensers. Such corrections and also the distribution of motive steam to the individual stages are problems best passed on to ejector manufacturers who have experience and a body of test data.



Figure 7.31. Steam requirements of ejectors at various pressure levels with appropriate numbers of stages and contact intercondensers. Steam pressure 100 psig, water temperature 85°F. Factor for 65 psig steam is 1.2 and for 200 psig steam it is 0.80 (Worthington *Corp*).

EXAMPLE 7.13

Interstage Condensers A four-stage ejector is to evacuate a system to 0.3 Torr. The

compression ratio in each stage will be

$$(P_4/P_0)^{1/4} = (760/0.3)^{1/4} = 7.09.$$

The individual stage pressures and corresponding water bubblepoint temperatures from the steam tables are



Figure 7.32. Progress of pressures, velocities, enthalpies and entropies in an ejector (*Coulson and* Richardson, Chemical Engineering, *Pergamon*, 1977, New York, Vol. 1).

GLOSSARY FOR CHAPTER 7

PUMP TERMS

Head has the dimensions [F][L]/[M]; for example, ft lbf/lb or ft; or N m/kg or m:

- **a.** pressure head = $\Delta P / \rho$;
- **b.** velocity head = $\Delta u^2/2g_c$;
- c. elevation head = $\Delta z(g/g_c)$, or commonly Az;
- d. friction head in line, $H_f = \hat{f} (L/D)u^2/2g_c$;
- e. system head H_s is made up of the preceding four items;
- **f.** pump head equals system head, $H_p = H_s$, under operating conditions;

Discharge	of	stage	0	1	2	3	4
Torr		•	0.3	2.1	15.1	107	760
°F				14	63.7	127.4	

The bubblepoint temperature in the second stage is marginal with normal cooling tower water, particularly with the practical restriction to 5°F below the bubblepoint. At the discharge of the third stage, however, either a surface or barometric condenser is quite feasible. At somewhat higher process pressure, two interstage condensers may be practical with a four-stage ejector, as indicated on Figure 7.31.

When barometric condensers are used, the effluent water temperature should be at least $5^{\circ}F$ below the bubblepoint at the prevailing pressure. A few bubblepoint temperatures at low pressures are:

Absolute (in.Hg)	0.2	0.5	1.0	2.0
Bubblepoint °F	34.6	58.8	79.0	101.1

Interstage pressures can be estimated on the assumption that compression ratios will be the same in each stage, with the suction to the first stage at the system pressure and the discharge of the last stage at atmospheric pressure. Example 7.13 examines at what stages it is feasible to employ condensers so as to minimize steam usage in subsequent stages.

EJECTOR THEORY

The progress of pressure, velocity, and energy along an ejector is illustrated in Figure 7.32. The initial expansion of the steam to point C and recompression of the mixture beyond point *E* proceed adiabatically with isentropic efficiencies of the order of 0.8. Mixing in the region from C to *E* proceeds with approximate conservation of momenta of the two streams, with an efficiency of the order of 0.65. In an example worked out by Dodge (1944, pp. 289–293), the compounding of these three efficiencies leads to a steam rate five times theoretical. Other studies of single-stage ejectors have been made by Work and Haedrich (1939) and DeFrate and Hoerl (1959), where other references to theory and data are made.

The theory is in principle amenable to the prediction of steam distribution to individual stages of a series, but no detailed procedures are readily available. Manufacturers charts such as Figure 7.31 state only the consumption of all the stages together.

- g. static suction head equals the difference in levels of suction liquid and the centerline of the pump;
- h. static suction lift is the static suction head when the suction level is below the centerline of the pump; numerically a negative number.

NPSH (net positive suction head) = (pressure head of source) + (static suction head) - (friction head of the suction line) - (vapor pressure of the flowing liquid).

Hydraulic horsepower is obtained by multiplying the weight rate of flow by the head difference across the pump and converting to horsepower. For example, HHP = (gpm)(psi)/1714 = (gpm)(sp gr)(ft)/3960. **Brake horsepower** is the driver power output needed to operate the pump. BHP = HHP/(pump efficiency).

Driver horsepower, HP = BHP/(driver efficiency) = HHP/(pump efficiency)(driver efficiency).

TERMS CONCERNING CENTRIFUGAL AND RELATED PUMPS

Axial flow is flow developed by axial thrust of a propeller blade, practically limited to heads under 50 ft or so.

Centrifugal pump consists of a rotor (impeller) in a casing in which a liquid is given a high velocity head that is largely converted to pressure head by the time the liquid reaches the outlet.

Characteristic curves are plots or equations relating the volumetric flow rate through a pump to the developed head or efficiency or power or NPSH.

Diffuser type: the impeller is surrounded by gradually expanding passages formed by stationary guide vanes [Figs. 7.2(b) and 7.3(d)].

Double suction: two incoming streams enter at the eye of the impeller on opposite sides, minimizing axial thrust and worthwhile for large, high head pumps [Fig. 7.2(b)].

Double volute: the liquid leaving the impeller is collected in two similar volutes displaced 180" with a common outlet; radial thrust is counterbalanced and shaft deflection is minimized, resulting in lower maintenance and repair, used in high speed pumps producing above 500 ft per stage.

Impeller: the rotor that accelerates the liquid.

a. Open impellers consist of vanes attached to a shaft without any

REFERENCES

Compressors

- Compressors in Encyclopedia Of Chemical Processing and Design, Dekker, New York, 1979, Vol. 10, pp. 157–409.
- F.L. Evans, Compressors and fans, in Equipment Design Handbook for Refineries and Chemical Plants, Gulf, Houston, 1979, Vol. 1, pp. 54–104.
- H. Gartmann, *DeLaval Engineering Handbook*, McGraw-Hill, New York, 1970, pp. 6.61-6.93.
- R. James, Compressor calculation procedures, in *Encyclopedia of Chemical Processing and Design*, Dekker, New York, Vol. 10, pp. 264-313.
- E.E. Ludwig, Compressors, in *Applied Process Design for Chemical and Petrochemical Plants*, Gulf, Houston, 1983, Vol. 3, pp. 251-396.
- 6. R.D. Madison, Fan Engineering, Buffalo Forge Co., Buffalo, NY, 1949.
- 7. H.F. Rase and M.H. Barrow, *Project Engineering of Process Plants*, Wiley, New York, pp. 297-347.

Ejectors

- 1. L.A. DeFrate and V. W. Haedrich, Chem. Eng. Prog. Symp. Ser. 21, 43-51 (1959).
- B.F. Dodge, *Chemical Engineering Thermodynamics*, McGraw-Hill, New York, 1944, pp. 289-293.
- F.I. Evans, Equipment Design Handbook for Refineries and Chemical Plants, Gulf, Houston, 1979, Vol. 1, pp. 105–117.
- 4. E.E. Ludwig, loc. cit., Vol. 1, pp. 206-239.
- R.E. Richenberg and J.J. Bawden, Ejectors, steam jet, in *Encyclopedia* of *Chemical Processing and Design*, Dekker, New York, Vol. 17, pp. 167–194.
- 6. L.T. Work and V.W. Haedrich, Ind. Eng. Chem. 31, 464-477 (1939).

Piping

1. ANSI Pioine Code. ASME, New York, 1980.

form of supporting sidewall and are suited to handling slurries without clogging [Fig. 7.2(a)].

- **b.** Semienclosed impellers have a complete shroud on one side [Fig. 7.3(c)]; they are essentially nonclogging, used primarily in small size pumps; clearance of the open face to the wall is typically 0.02 in. for 10 in. diameters.
- c. Closed impellers have shrouds on both sides of the vanes from the eye to the periphery, used for clear liquids [Fig. 7.3(b)].

Mechanical seals prevent leakage at the rotating shaft by sliding metal on metal lubricated by a slight flow of pump liquid or an independent liquid [Figs. 7.4(c) and (d)].

Miied flow: develops head by combined centrifugal action and propeller action in the axial direction, suited to high flow rates at moderate heads [Fig. 7.3(e)].

Multistage: several pumps in series in a single casing with the objective of developing high heads. Figure 7.6(c) is of characteristic curves.

Performance curves (see characteristic curves).

Single suction: the liquid enters on one side at the eye of the impeller; most pumps are of this lower cost style [Fig. 7.2(c)].

Split case: constructed so that the internals can be accessed without disconnecting the piping [Fig. 7.2(a)].

Stuffing box: prevent leakage at the rotating shaft with compressed soft packing that may be wetted with the pump liquid or from an independent source [Figs. 7.4(a) and (b)].

Volute type: the impeller discharges the liquid into a progressively expanding spiral [Fig. 7.2(a)].

- S. Chalfin, Control valves, Encyclopedia of Chemical Processing and Design, Dekker, New York, 1980, Vol. 11, pp. 187-213.
- F.L. Evans, Equipment Design Handbook for Refineries and Chemical Plants, Gulf, Houston, 1979, Vol. 2; piping, pp. 188-304; valves, pp. 315-332.
- J.W. Hutchinson, ISA Handbook of Control Valves, Inst. Soc. America, Research Triangle Park, NC, 1976.
- 5. R.C. King, Piping Handbook, McGraw-Hill, New York, 1967.
- J.L. Lyons, Encyclopedia of Values, Van Nostrand Reinhold, New York, 1975.
- Marks' Standard Handbook for Mechanical Engineers, McGraw-Hill, New York, 1987.
- 8. Perry's Chemical Engineers' Handbook, McGraw-Hill, New York, 1984.
- 9. R. Weaver, Process Piping Design, Gulf, Houston, 1973, 2 Vols.
- P. Wing, Control valves, in Process Instruments and Controls Handbook, (D.M. Considine, Ed.), McGraw-Hill, New York, 1974.
- 11. R.W. Zappe, Value Selection Handbook, Gulf, Houston, pp. 19.1–19.60, 1981.

Pumps

- D. Azbel and N.P. Cheremisinoff, *Fluid Mechanics and Fluid Operations*, Ann Arbor Science, Ann Arbor, MI, 1983.
- N.P. Cheremisinoff, *Fluid Flow: Pumps, Pipes and Channels*, Ann Arbor Science, Ann Arbor, MI, 1981.
- 3. F.L. Evans, loc. cit., Vol. 1, pp. 118-171.
- H. Gartmann, *DeLaval Engineering Handbook*, McGraw-Hill, New York, 1970, pp. 6.1–6.60.
- I.J. Karassik and R. Carter, Centrifugal Pump Selection Operation and Maintenance, F.W. Dodge Corp., New York, 1960.
- I.J. Karassik, W.C. Krutsch, W.H. Fraser, and Y.J.P. Messina, *Pump Handbook*, McGraw-Hill, New York, 1976.
- 7. F.A. Kristal and F.A. Annett, Pumps, McGraw-Hill, New York, 1940.
- 8. E.E. Ludwig, loc. cit., Vol. 1, pp. 104-143.
- S. Yedidiah, *Centrifugal Pump Problems*, Petroleum Publishing, Tulsa. OK. 1980.

HEAT TRANSFER AND HEAT EXCHANGERS

B asic concepts of heat transfer are reviewed in this chapter and applied primarily to heat exchangers, which are equipment for the transfer of heat between two fluids through a separating wall. Heat transfer a/so is a key process in other specialized equipment, some of which are treated in the next and other chapters. The three recognized modes of heat transfer are by conduction, convection, and radiation, and may occur simultaneously in some equipment.

8.1. CONDUCTION OF HEAT

In a solid wall such as Figure 8.1(a), the variation of temperature with time and position is represented by the one-dimensional Fourier equation

$$\frac{\partial T}{\partial \theta} = kA \frac{\partial^2 T}{\partial x^2} \tag{8.1}$$

For the most part, only the steady state condition will be of concern here, in which the case the partial integral of Eq. (8.1) becomes

$$Q = -kA\frac{dT}{dx},\tag{8.2}$$

assuming the thermal conductivity k to be independent of temperature. Furthermore, when both k and A are independent of position,

$$Q = -kA \frac{\Delta T}{\Delta x} = \frac{kA}{L} (T_0 - T_L), \qquad (8.3)$$

in the notation of Figure 8.1(a).

Equation (8.3) is the basic form into which more complex situations often are cast. For example,

$$Q = kA_{mean} \frac{\Delta T}{L}$$
(8.4)

when the area is variable and

$$Q = UA(\Delta T)_{\text{mean}} \tag{8.5}$$

in certain kinds of heat exchangers with variable temperature difference.

THERMAL CONDUCTIVITY

Thermal conductivity is a fundamental property of substances that basically is obtained experimentally although some estimation methods also are available. It varies somewhat with temperature. In many heat transfer situations an average value over the prevailing temperature range often is adequate. When the variation is linear with

$$k = k_0(1 + \alpha T), \tag{8.6}$$

the integral of Eq. (8.2) becomes

$$Q(L/A) = k_0[T_1 - T_2 + 0.5\alpha(T_1^2 - T_2^2)] = k_0(T_1 - T_2)[1 + 0.5\alpha(T_1 + T_2)],$$
(8.7)

which demonstrates that use of a value at the average temperature gives an exact result. Thermal conductivity data at several temperatures of some metals used in heat exchangers are in Table 8.1. The order of magnitude of the temperature effect on k is illustrated in Example 8.1.











Figure 8.1. Temperature profiles in one-dimensional conduction of heat. (a) Constant cross section. (b) Hollow cylinder. (c) Composite flat wall. (d) Composite hollow cylindrical wall. (e) From fluid A to fluid F through a wall and fouling resistance in the presence of eddies. (f) Through equivalent fluid films, fouling resistances, and metal wall.

TABLE	8.1.	Thermal	Conduc	tivities	of	Some	Metals
		Common	ly Used	in Hea	tΕ	xchang	gers
		[kBtu/(l	nr)(saft)(°F/ft)]			-

		Tempera	iture (°F)	
Metal or Alloy	- 100	7 0	200	1000
Steels				
Carbon		30.0	27.6	22.2
1 Cr ¹ ₄ Mo	_	19.2	19.1	18.0
410	_	13.0	14.4	_
304	_	9.4	10.0	13.7
316	8.1	9.4		13.0
Monel 400	11.6	12.6	13.8	22.0
Nickel 200	_	32.5	31.9	30.6
Inconel 600		8.6	9.1	14.3
Hastelloy C	_	7.3	5.6	10.2
Aluminum		131	133	
Titanium	11.8	11.5	10.9	12.1
Tantalum		31.8		
Copper	225	225	222	209
Yellow brass	56	69		_
Admiralty	55	64	—	_

HOLLOW CYLINDER

As it appears on Figure 8.1(b), as the heat flows from the inside to the outside the area changes constantly. Accordingly the equivalent of Eq. (8.2) becomes, for a cylinder of length N,

$$Q = -kN(2\pi r)\frac{dT}{dr}, \qquad (8.8)$$

of which the integral is

$$Q = \frac{2\pi k N (T_1 - T_2)}{\ln(r_2/r_1)}$$
(8.9)

This may be written in the standard form of Eq. (8.4) by taking

$$A_{,,,} = 2\pi L N r_{\rm im} \tag{8.10}$$

and

$$L = r_2 - r_1, (8.11)$$

where

$$r_{\rm lm} = (r_2 \quad r_1)/\ln(r_2/r_1)$$
 (8.12)

EXAMPLE 8.1

Conduction through a Furnace Wall

A furnace wall made of fire clay has an inside temperature of 1500°F and an outside one of 300°F. The equation of the thermal conductivity is k = 0.48[1+5.15(E-4)T]Btu/(hr)(sqft)(°F/ft). Accordingly,

$$Q(L/A) = 0.48(1500 - 300)[1 + 5.15(E - 4)(900)] = 0.703.$$

If the conductivity at 300°F had been used, Q(L/A) = 0.554.

is the logarithmic mean radius of the hollow cylinder. This concept is not particularly useful here, but logarithmic means also occur in other more important heat transfer situations.

COMPOSITE WALLS

The flow rate of heat is the same through each wall of Figure 8.1(c). In terms of the overall temperature difference,

$$Q = UA(T_1 - T_4), \tag{8.13}$$

where U is the overall heat transfer coefficient and is given by

$$\frac{1}{U} = \frac{1}{k_a/L_a} + \frac{1}{k_b/L_b} + \frac{1}{k_c/L_c}.$$
(8.14)

The reciprocals in Eq. (8.14) may be interpreted as resistances to heat transfer, and so it appears that thermal resistances in series are additive.

For the composite hollow cylinder of Figure 8.1(d), with length N,

$$Q = \frac{2\pi N(T_1 - T_4)}{\ln(r_2/r_1)/k_a + \ln(r_3/r_2)/k_b + \ln(r_4/r_3)/k_c},$$
(8.15)

With an overall coefficient U_i based on the inside area, for example,

$$Q = 2\pi N r_i U_i (T_1 - T_4) = \frac{2\pi N (T_1 - T_4)}{1/U_i r_{i_1}}.$$
(8.16)

On comparison of Eqs. (8.15) and (8.16), an expression for the inside overall coefficient appears to be

$$\frac{1}{U_i} = r_i \left[\frac{\ln(r_2/r_i)}{k_a} + \frac{\ln(r_3/r_2)}{k_b} + \frac{\ln(r_4/r_3)}{k_c} \right]$$
(8.17)

In terms of the logarithmic mean radii of the individual cylinders,

$$\frac{1}{U_i} = r_i \left[\frac{1}{k_a r_{ma}/(r_2 - r_1) + k_b r_{mb}/(r_3 - r_2)} + \frac{1}{k_c r_{mc}/(r_4 - r_3)} \right],$$

which is similar to Eq. (8.14) for flat walls, but includes a ratio of radii as a correction for each cylinder.

FLUID FILMS

Heat transfer between a fluid and a solid wall can be represented by conduction equations. It is assumed that the difference in temperature between fluid and wall is due entirely to a stagnant film of liquid adhering to the wall and in which the temperature profile is linear. Figure 8.1(e) is a somewhat realistic representation of a temperature profile in the transfer of heat from one fluid to another through a wall and fouling scale, whereas the more nearly ideal Figure 8.1(f) concentrates the temperature drops in stagnant fluid and fouling films.

Since the film thicknesses are not definite quantities, they are best combined with the conductivities into single coefficients

$$h = k/L \tag{8.18}$$

so that the rate of heat transfer through the film becomes

$$Q = hA \land T. \tag{8.19}$$

Through the five resistances of Figure 8.1(f), the overall heat

EXAMPLE 8.2 Effect of Ignoring the Radius Correction of the Overall Heat Transfer Coefficient

The two film coefficients are 100 each, the two fouling coefficients are 2000 each, the tube outside diameter is 0.1 ft, wall thickness is 0.01 ft, and thermal conductivity of the metal is 30:

$$\begin{aligned} r_i/r_o &= 0.04/0.05 = 0.8, \\ r_m &= (0.05 - 0.04)/\ln 1.25 = 0.0448, \\ r_m/r_o &= 0.8963, \\ U_o &= [1/100(0.8) + 1/2000(0.8) + 1/(30/0.01)(0.8963) \\ &+ 1/100 + 1/2000].^{\prime} = 41.6721. \end{aligned}$$

transfer coefficient is given by

$$\frac{1}{U} = \frac{1}{h_1} + \frac{1}{h_2} + \frac{1}{k_3/L_3} + \frac{1}{h_4} + \frac{1}{h_5}, \qquad (8.20)$$

where L_3 is the thickness of the metal.

If the wall is that of hollow cylinder with radii r_i and r_o , the overall heat transfer coefficient based on the outside surface is

$$\frac{1}{U_o} = \frac{1}{h_1(r_i/r_o)} + \frac{1}{h_2(r_i/r_0)} + \frac{1}{k_3/L_3(r_m/r_0)} + \frac{1}{h_4} + \frac{1}{h_5} , \qquad (8.21)$$

Basing on the inside area,

$$U_i = [1/100 + 1/2000 + [(30/0.01)(0.0448/0.04)]^{-1} + 0.8/100 + 0.8/2000]^{-1} = 52.0898.$$

Ignoring the corrections,

$$U = (2/100 + 2/2000 + 1/30/0.01)^{-1} = 46.8750.$$

The last value is very nearly the average of the other two.

where r_m is the mean radius of the cylinder, given by Eq. (8.12).

Since wall thicknesses of heat exchangers are relatively small and the accuracy of heat transfer coefficients may not be great, the ratio of radii in Eq. (8.21) often is ignored, so that the equation for the overall coefficient becomes simply

$$\frac{1}{U} = \frac{1}{h_1} + \frac{1}{h_2} + \frac{1}{k_3/L_3} + \frac{1}{h_4} + \frac{1}{h_5}.$$
(8.22)

The results of the typical case of Example 8.2, however, indicate that the correction may be significant. A case with two films and two solid cylindrical walls is examined in Example 8.3.

EXAMPLE 8.3

A Case of a Composite Wall: Optimum Insulation Thickness for a Steam Line

A 3 in. IPS Sched 40 steel line carries steam at 500°F. Ambient air is at 70°F. Steam side coefficient is 1000 and air side is 3 Btu/(hr)(sqft)(°F). Conductivity of the metal is 30 and that of insulation is 0.05 Btu/(hr)(sqft)(°F/ft). Value of the steam is \$5.00/MBtu. cost of the insulation is \$1.5/(yr)(cuft). Operation is 8760 hr/yr. The optimum diameter d of insulation thickness will be found.

 $d_o = 0.2917$ ft, $d_{1} = 0.2557$ ft, $\ln(d_o/d_i) = 0.1317.$

Insulation:

$$\ln(d_o/d_i) = \ln(d/0.2917). \tag{1}$$

Heat transfer coefficient based on inside area:

$$U_{i} = d_{i} \left[\frac{1}{1000d_{i}} + \frac{0.1317}{30} + \frac{\ln(d/0.2917)}{0.05} + \frac{1}{3d} \right]^{-1},$$
(2)
$$Q/A_{i} = U_{i}\Delta T = 430U_{i}.$$

Steam cost:

$$C_1 = 5(10^{-6})(8760)Q/A_i$$

= 0.0438Q/A_i, \$ (yr)(sqft inside). (3)



60 U=1/U1/D2 70 Q=430*U

10

40 50

- 80
- C1=.0438*Q C2=1.5*(D^2-01^2)/D2^2 90
- 100 C=C1+C2 ! (rea'd to be minim um)
- 110 PRINT USING 120; D.U.C1,C2,
- IMAGE .ODD/X/ .DOD/X/DD.DD/X/ 120 00.00,%,00.0000

130 GOTO 40



Heat transfer coefficients are empirical data and derived correlations. They are in the form of overall coefficients U for frequently occurring operations, or as individual film coefficients and fouling factors.

8.2. MEAN TEMPERATURE DIFFERENCE

In a heat exchanger, heat is transferred between hot and cold fluids through a solid wall. The fluids may be process streams or independent sources of heat such as the fluids of Table 8.2 or sources of refrigeration. Figure 8.2 shows such a process with inlet and outlet streams, but with the internal flow pattern unidentified because it varies from case to case. At any cross section, the differential rate of heat transfer is

$$dQ = U(T - T') dA = -mcdT = m'c' dT'.$$
(8.23)

The overall heat transfer rate is represented formally by

$$Q = UA(\Delta T)_m. \tag{8.24}$$

The mean temperature difference (AT), depends on the terminal temperatures, the thermal properties of the two fluids and on the flow pattern through the exchanger.

Total cost:

$$C = C_1 + C_2 \rightarrow \text{minimum.}$$
(5)

Substitute Eqs. (2)-(4) into Eq. (5). The outside diameter is the key unknown.

The cost curve is fairly flat, with a minimum at d = 0.50 ft, corresponding to 1.25 in. thickness of insulation. Some trials are shown with the computer program. A more detailed analysis of insulation optima is made by Happel and Jordan [*Chem. Process Econ.*, 380 (1975)], although their prices are dated. Section 8.12 also discusses insulation.



Figure 8.2. Terminal temperatures and temperature differences of a heat exchanger, with unidentified internal flow pattern.

SINGLE PASS EXCHANGER

The simplest flow patterns are single pass of each fluid, in either the same or opposite directions. Temperature profiles of the main kinds of thermal behavior are indicated on Figure 8.3(a). When the unbroken lines [cases (a)-(e)] are substantially straight, the mean temperature is expressed in terms of the terminal differences by

$$(\Delta T)_m = (\Delta T)_{\log mean} = \frac{(\Delta T)_2 - (\Delta T)_1}{\ln[(\Delta T)_2/(\Delta T)_1]}.$$
(8.25)

This is called the logarithmic mean temperature difference. The temperature profiles are straight when the heat capacities are

TABLE 8.2. Properties of Heat Transfer Media

Medium	Trade Name	Phase	°F	atm, gage	Remarks		
Electricity	_		100-4500	-	_		
Water	-	vapor	200-l 100	O-300			
Water		liquid	300-400	6-15	—		
Flue gas		gas	100-2000	o-7	-		
Diphenyl-diphenyl oxide eutectic	Dowtherm A	liquid or vapor	450-750	o-9	nontoxic, carbonizes at high temp		
Di + triaryl cpds	Dowtherm G	liquid	20-700	o-3	sensitive to oxygen		
Ethylene glycol, inhibited	Dow SR-1	liquid	-40250	0	acceptable in food industry		
Dimethyl silicones	Dow Syltherm 800	liquid	-40-750	0	low toxicity		
Mixed silanes	Hydrotherm	liquid	-50-675	0	react with oxygen and moisture		
Aromatic mineral oil	Mobiltherm, Mobil	liquid	100-600	0	not used with copper based materials		
Chlorinated biphenyls	Therminol, Monsanto	liquid	50-600	0	toxic decomposition products		
Molten nitrites and nitrates of K and Na	Hi-Tee, DuPont	liquid	300-1100	0	resistant alloys needed above 850°F		
Sodium-potassium eutectic		liguid	100-1400	0	stainless steel needed above 1000°F		
Mercury		vapor	600-1000	0-12	low pressure vapor, toxic, and expensive		



Figure 8.3. Temperature profiles in heat exchangers. (a) In parallel or countercurrent flow, with one or two phases. (b) One shell pass, two tube passes. (c) Two shell passes, four tube passes.

substantially independent of temperature over the range of the process, or when a phase change occurs at constant temperature.

When the profiles consist of linear sections, as in cases (f) and (g), the exchanger can be treated as a three-section assembly, each characterized by its own log mean temperature difference, for which intermediate temperatures may be found by direct calculation or by trial. Heat transfer for a case such as (h) with continuously curved profile must be evaluated by integration of Eq. (8.23).

MULTIPASS EXCHANGERS

For reasons of compactness of equipment, the paths of both fluids may require several reversals of direction. Two of the simpler cases of Figure 8.3 are (b) one pass on the shell side and two passes on the tube side and (c) two passes on the shell side and four on the tube side. On a baffled shell side, as on Figure 8.4(c), the dominant flow is in the axial direction, so this pattern still is regarded as single pass on the shell side. In the cross flow pattern of Figure 8.5(c), each stream flows without lateral mixing, for instance in equipment like Figure 8.6(h). In Figure 8.6(i) considerable lateral mixing would occur on the gas side. Lateral mixing could occur on both sides of the plate exchanger of Figure 8.6(h) if the fins were absent.

Mean temperature differences in such flow patterns are obtained by solving the differential equation. Analytical solutions have been found for the simpler cases, and numerical ones for many important complex patterns, whose results sometimes are available in generalized graphical form.

f-METHOD

When all of the terminal temperatures are known, the mean temperature difference is found directly from

$$(\Delta T)_m = F(\Delta T)_{\log mean}, \tag{8.26}$$

where the correction factor F depends on the flow pattern and is



Figure 8.4. Example of tubular heat exchangers (see also Fig. 8.14). (a) Double-pipe exchanger. (b) Scraped inner surface of a double-pipe exchanger. (c) Shell-and-tube exchanger with fixed tube sheets. (d) Kettle-type reboiler. (e) Horizontal shell side thermosiphon reboiler. (f) Vertical tube side thermosiphon reboiler. (g) Internal reboiler in a tower. (h) Air cooler with induced draft fan above the tube hank. (i) Air cooler with forced draft fan below the tube bank.



(a)

Figure 8.5. Correction factor F, effectiveness and number of transfer units in multipass and cross flow heat exchangers (Bowman *et al.*, Trans ASME 283, 1940; Kays and London, 1984):

$$P = \frac{T_i - T_o}{T_i - T_i'}, \quad R = \frac{T_i' - T_o'}{T_i - T_o}$$

T on the tubeside, *T* on the shellside. i = input, $\theta = \text{output}$. (a) One pass on shellside, any multiple of two passes on tubeside. (b) Two passes on shell side, any multiple of four on tubeside. (c) Cross flow, both streams unmixed laterally. (d) Cross flow, one stream mixed laterally. (e) Cross flow, both streams mixed laterally. (f) Effectiveness and number of transfer units in parallel and countercurrent flows. (g) Three shell passes, multiples of six on tubeside. (h) Four shell passes, multiples of eight on tubeside. (i) Five shell passes, multiples of ten on tubeside. (j) Six shell passes, multiples of 12 on tubeside.

$176\,$ Heat transfer and heat exchangers



(d)







Figure 8.5—(continued)



(f)











expressed in terms of these functions of the terminal temperatures:

$$P = \frac{T_o - T_i}{T_i' - T_i} = \frac{\text{actual heat transfer}}{\text{maximum possible heat transfer}},$$
(8.27)

$$R = \frac{T_i - T_o}{T_o' - T_i'} = \frac{mc}{m'c'}.$$
(8.28)

Some analytical expressions for *Fare* shown in Table 8.3, and more graphical solutions in Figure 8.5.







This method is especially easy to apply when the terminal temperatures are all known, because then F and $(\Delta T)_{\log mean}$ are immediately determinable for a particular flow pattern. Then in the heat transfer equation

$$Q = UAF(\Delta T)_{\rm lm} \tag{8.29}$$

any one of the quantities Q, U, or A may be found in terms of the others. A solution by trial is needed when one of the terminal temperatures is unknown, as shown in Example 8.4. The next



Figure 8.6. Examples of extended surfaces on one or both sides. (a) Radial fins. (b) Serrated radial fins. (c) Studded surface. (d) Joint between tubesheet and low fin tube with three times bare surface. (e) External axial fins. (f) Internal axial fins. (g) Finned surface with internal spiral to promote turbulence. (h) Plate fins on both sides. (i) Tubes and plate fins.

 TABLE 8.3. Formulas for Mean Temperature Difference and Effectiveness in Heat Exchangers



1. Parallel or countercurrent flow,

$$(\Delta T)_m = (\Delta T)_{logmean} = (AT, -\Delta T_2)/ln(\Delta T_1/\Delta T_2).$$

2. In general,

 $(\Delta T)_m = F(\Delta T)_{\text{logmean}}$

o r

(AT), = $\theta(T_i - T'_i)$,

where F and θ depend on the actual flow paths on the shell and tube sides and are expressed in terms of these quantities:

$$\begin{split} C &= C_{\min}/C_{\max'} \\ P &= (T_o - T_i)/(T_i' - T_i) = \text{actual heat transfer} \\ & (\text{maximum possible heat transfer}), \\ R &= (T_i - T_o)/(T_o' - 6') = m'c'/mc. \end{split}$$

3. Number of transfer units, N or NTU, is

 $N = UA/C_{\min} \approx P/\theta$,

where \textit{C}_{min} is the smaller of the two values mc or m'c' of the products of mass rate of flow times the heat capacity.

- 4. In parallel flow, $P = N\theta = \{I = \exp[-N(1 + C)]\}/(1 + C)$.
- 5. In countercurrent flow, $P = N\theta = \{1 \exp[-N(1 C)]\}/\{1 C \exp[-N(1 C)]\}$.

6. One shell pass and any multiple of two tube passes,

$$F = \frac{\sqrt{R^2 + 1}}{R - 1} \cdot \ln\left(\frac{1 - P}{1 - PR}\right) / \ln\left[\frac{2 - P(R + 1 - \sqrt{R^2 + 1})}{2 - P(R + 1 + \sqrt{R^2 + 1})}\right], \quad R \neq 1,$$

$$F = \frac{P}{1 - P} \cdot \sqrt{2} / \ln\left[\frac{2 - P(2 - \sqrt{2})}{2 - P(2 + \sqrt{2})}\right], \quad R = 1,$$

$$P = 2\left\{1 + C + (1 + C^2)^{1/2} \frac{1 + \exp[-N(1 + C^2)^{1/2}]}{1 - \exp[-N(1 + C^2)^{1/2}]}\right\}^{-1}.$$

7. Two shell passes and any multiple of four tube passes,

$$F = \left[\frac{\sqrt{R^2 + 1}}{2(R - 1)} \ln \frac{1 - PR}{1 - PR}\right] / \ln \left[\frac{2/P - 1 - R + (2/P)\sqrt{(1 - P)(1 - PR)} + \sqrt{R^2 + 1}}{2/P - 1 - R + (2/P)\sqrt{(1 - P)(1 - PR)} - \sqrt{R^2 + 1}}\right]$$

8. Cross flow,

- (a) Both streams laterally unmixed, $P=1 = \exp\{[\exp(-NCn)-1]/Cn\}$, where $n = N^{-0.22}$.
- (b) Both streams mixed, $P = \{1/[1 \exp(-N)] + C/[1 \exp(-NC)] 1/N\}^{-1}$.

(c) C_{max} mixed, C_{min} unmixed, $P = (1/C)\{1 - \exp[-C(1 - e^{-N})]\}$.

(d) C_{\min} mixed, C_{\max} unmixed, $P=1 - \exp\{-(1/C)[1 - \exp(-NC)]\}$

 For more complicated patterns only numerical solutions have been made. Graphs of these appear in sources such as Heat Exchanger Design Handbook (HEDH, 1983) and Kays and London (1984). method to be described, however, may be more convenient in such a case.

B-METHOD

One measure of the size of heat transfer equipment is the number of transfer units N defined by

$$N = UA/C_{\min},\tag{8.30}$$

where C_{\min} is the smaller of the two products of mass flow rate and heat capacity, mc or m'c'. N is so named because of a loose analogy with the corresponding measure of the size of mass transfer equipment.

A useful combination of P and N is their ratio

$$\theta = \frac{P}{N} = \frac{C_{\min}(T_o - T_i)}{UA(T_i' - T_i)} = \frac{Q}{UA(T_i' - T_i)} = \frac{(\Delta T)_m}{(T_i' - T_i)},$$
(8.31)

where $(T_o - T_i)$ is the temperature change of the stream with the smaller value of mc. Thus θ is a factor for obtaining the mean temperature difference by the formula:

$$(AT)_{i,j} = \theta(T'_i - T_j) \tag{8.32}$$

when the two inlet temperatures are known.

The term P often is called the exchanger effectiveness. Equations and graphs are in Table 8.3 and Figure 8.4. Many graphs for θ , like those of Figure 8.7, may be found in the *Heat* Exchanger *Design Handbook* (HEDH, 1983). When sufficient other data are known about a heat exchange process, an unknown outlet temperature can be found by this method directly without requiring trial calculations as with the F-method. Example 8.5 solves such a problem.

SELECTION OF SHELL-AND-TUBE NUMBERS OF PASSES

A low value of F means, of course, a large surface requirement for a given heat load. Performance is improved in such cases by using several shells in series, or by increasing the numbers of passes in the same shell. Thus, two l-2 exchangers in series are equivalent to one large 2-4 exchanger, with two passes on the shell side and four passes on the tube side. Usually the single shell arrangement is **more** economical, even with the more complex internals. For economy, F usually should be greater than 0.7.

EXAMPLE

A shell side fluid is required to go from 200 to $140^{\circ}F$ and the tube side from 80 to $158^{\circ}F$. The charts of Figure 8.5 will be used:

$$P = (200 - 140)/(200 - 80) = 0.5,$$

$$R = (158 - 80)/(200 - 140) \approx 1.30.$$

For a 1-2 exchanger, $F = 0.485$:
2-4 0.92
4-8 0.98

The 1-2 exchanger is not acceptable, but the 2-4 is acceptable. If the tube side outlet were at 160 instead of 158, F would be zero for the 1-2 exchanger but substantially unchanged for the others.

8.3. HEAT TRANSFER COEFFICIENTS

Data are available as overall coefficients, individual film coefficients, fouling factors, and correlations of film coefficients in terms of

EXAMPLE 8.4

Performance of a Heat Exchanger with the F-Method Operation of an exchanger is represented by the sketch and the equation

$$Q/UA = 50 = F(\Delta T)_{\rm lm}$$



The outlet temperature of the hot fluid is unknown and designated by T. These quantities are formulated as follows:

$$P = \frac{200 - T}{200 - 80}$$

$$R = \frac{200 - T}{120 - 80'}$$

$$(\Delta T)_{\rm im} = \frac{T - 80 - (200 - 120)}{\ln[(T - 80)/(200 - 120)]}$$

F is represented by the equation of Item 6 of Table 8.3, or by Figure 8.4(a). Values of *T* are tried until one is found that satisfies $G \equiv 50 - F(\Delta T)_{1m} \approx 0$. The printout shows that

T = 145.197.

The sensitivity of the calculation is shown in the following

physical properties and operating conditions. The reliabilities of these classes of data increase in the order of this listing, but also the ease of use of the data diminishes in the same sequence.

OVERALL COEFFICIENTS

The range of overall heat transfer coefficients is approximately 10-200 Btu/(hr)(sqft)(°F). Several compilations of data are available, notably in the *Chemical Engineers Handbook* (McGraw-Hill, New York, 1984, pp. 10.41-10.46) and in Ludwig (1983, pp. 70-73). Table 8.4 qualifies each listing to some extent, with respect to the kind of heat transfer, the kind of equipment, kind of process stream, and temperature range. Even so, the range of values of *U* usually is two- to three-fold, and consequently only a rough measure of equipment size can be obtained in many cases with such data. Ranges of the coefficients in various kinds of equipment are compared in Table 8.5.

FOULING FACTORS

Heat transfer may be degraded in time by corrosion, deposits of reaction products, organic growths, etc. These effects are accounted for quantitatively by fouling resistances, $1/h_f$. They are listed separately in Tables 8.4 and 8.6, but the listed values of coefficients include these resistances. For instance, with a clean surface the first listed value of U in Table 8.4 would correspond to a clean value of U = 1/(1/12-0.04) = 23.1. How long a clean value could be

tabulation:

10

20

30

4⊰ 50

ĥЙ

```
T P R (\Delta T)_{lm} F G

145.0 0.458 1.375 72.24 0.679 0.94

145.197 0.457 1.370 72.35 0.691 0.00061

145.5 0.454 1.363 72.51 0.708 -1.34

! Example 8.4. The F-method

SHORT P, R, F, T1

INPUT T

P=(200-T)/120

R=(200-T)/40

T1=(T-160)/L0G((T-80)/80)
```

70	E=(R^2+1)^.5
80	E=E/(R-1)*100((1-P)/(1-P*R))
90	F=F/L0G((2-P*(R+1-E))/(2-P*(
95	G=50-F*T1
100	PRINT "T=";T
110	PRINT "G=";G
120	PRINT "P=";P
120	PDINT "P=";P
130 140 150 160 170	PRINT "F=";F PRINT "T1=";T1 GOTO 3 0 END
T=	
P= (190240286
R= 1	145669
F= .	69109
T1=	72 346

maintained in a particular plant is not certain. Sometimes fouling develops slowly; in other cases it develops quickly as a result of process upset and may level off. A high coefficient often is desirable, but sometimes is harmful in that excessive subcooling may occur or film boiling may develop. The most complete list of fouling factors with some degree of general acceptance is in the TEMA (1978) standards. The applicability of these data to any particular situation, however, is questionable and the values probably not better than $\pm 50\%$. Moreover, the magnitudes and uncertainties of arbitrary fouling factors may take the edge off the importance of precise calculations of heat transfer coefficients. A brief discussion of fouling is by Walker (1982). A symposium on this important topic is edited by Somerscales and Knudsen (1981).

INDIVIDUAL FILM COEFFICIENTS

Combining individual film coefficients into an overall coefficient of heat transfer allows taking into account a greater variety and range of conditions, and should provide a better estimate. Such individual coefficients are listed in Tables 8.6 and 8.7. The first of these is a very cautious compilation with a value range of 1.5- to 2-fold. Values of the fouling factors are included in the coefficient listings of both tables but are not identified in Table 8.7. For clean service, for example, involving sensible heat transfer from a medium organic to heating a heavy organic,

$$U = 10,000/(57 - 16 + 50 - 34) = 175$$



Figure 8.7. θ correction charts for mean temperature difference: (a) One shell pass and any multiple of two tube passes. (b) Two shell passes and any multiple of four tube passes. [(HEDH, 1983); after Mueller in Rohsenow and Hartnett, Handbook of Heat Transfer, Section 18, McGraw-Hill, New York, 1973. Other cases also are covered in these references.]

EXAMPLE 8.5

Application of the Effectiveness and the θ Method Operating data of an exchanger are shown on the sketch. These data include

$$UA = 2000,$$

n'c' = 1000, mc = 800,
 $C = C_{min}/C_{max} = 0.8.$



The equation for effectiveness *P* is given by item 6 of Table 8.3 or it can be read off Figure 8.4(a). Both *P* and θ also can be read off Figure 8.4(a) at known N and $\mathbf{R} = C_2/C_1 = 0.8$. The number of

compared with a normal value of

$$U = 10,000/(57 + 50) = 93,$$

where the averages of the listed numbers in Table 8.6 are taken in each case.

METAL WALL RESISTANCE

With the usual materials of construction of heat transfer surfaces, the magnitudes of their thermal resistances may be comparable with the other prevailing resistances. For example, heat exchanger tubing of 1/16 in. wall thickness has these values of $1/h_w = L/k$ for several common materials:

Carbon steel	$1/h_w = 1.76 \times 10^{-4}$
Stainless steel	5.54 x 10 ⁻⁴
Aluminum	0.40 x 10 ⁻⁴
Glass	79.0×10^{-4}

which are in the range of the given film and fouling resistances, and should not be neglected in evaluating the overall coefficient. For example, with the data of this list a coefficient of 93 with carbon steel tubing is reduced to 88.9 when stainless steel tubing is substituted.

DIMENSIONLESS GROUPS

The effects of the many variables that bear on the magnitudes of individual heat transfer coefficients are represented most logically and compactly in terms of dimensionless groups. The ones most pertinent to heat transfer are listed in Table 8.8. Some groups have ready physical interpretations that may assist in selecting the ones appropriate to particular heat transfer processes. Such interpretations are discussed for example by Gröber et al. (1961, pp. 193-198). A few are given here.

The Reynolds number, $Du\rho/\mu = \rho u^2/(\mu u/D)$, is a measure of the ratio of inertial to viscous forces.

transfer units is

N = UA/C_{min} = 2000/800 = 2.5,
C = C_{min}/C_{max} = 0.8,
D =
$$\sqrt{1 + C^2}$$
 = 1.2806,
P = $\frac{2}{1 + C + D[1 + \exp(-ND)]/1 - \exp(-ND)}$ = 0.6271,
 θ = P/N = 0.2508,
AT, = θ (200 = 80) = 30.1,
Q = UA(ΔT)_m = 2000(30.1) = 60,200,
= 800(200 - T₂) = 1000(T'_2 = 80),
∴ T₂ = 124.75,
T'_2 = 140.2.

T₂ also may be found from the definition of **P**:

$$P = \frac{\text{actual AT}}{\text{max possible AT}} = \frac{200 - T_2}{200 - 80} = 0.6271,$$

$$\therefore T_2 = 124.78.$$

With this method, unknown terminal temperatures are found without trial calculations.

The Nusselt number, hL/k = h/(k/L), is the ratio of effective heat transfer to that which would take place by conduction through a film of thickness L.

The Peclet number, DGC/k = GC/(k/D) and its modification, the Graetz number wC/kL, are ratios of sensible heat change of the flowing fluid to the rate of heat conduction through a film of thickness D or L

The Prandtl number, $C\mu/k = (\mu/\rho)/(k/\rho C)$, compares the rate of momentum transfer through friction to the thermal diffusivity or the transport of heat by conduction.

The Grashof number is interpreted as the ratio of the product of the buoyancy and inertial forces to the square of the viscous forces.

The Stanton number is a ratio of the temperature change of a fluid to the temperature drop between fluid and wall. Also, St = (Nu)/(Re)(Pr).

An analogy exists between the transfers of heat and mass in moving fluids, such that correlations of heat transfer involving the Prandtl number are valid for mass transfer when the Prandtl number $C\mu/k$ is replaced by the Schmidt number $\mu/\rho k_d$. This is of particular value in correlating heat transfer from small particles to fluids where particle temperatures are hard to measure but measurement of mass transfer may be feasible, for example, in vaporization of naphthalene.

8.4. DATA OF HEAT TRANSFER COEFFICIENTS

Specific correlations of individual film coefficients necessarily are restricted in scope. Among the distinctions that are made are those of geometry, whether inside or outside of tubes for instance, or the shapes of the heat transfer surfaces; free or forced convection; laminar or turbulent flow; liquids, gases, liquid metals, non-Newtonian fluids; pure substances or mixtures; completely or partially condensable; air, water, refrigerants, or other specific substances; fluidized or fixed particles; combined convection and radiation; and others. In spite of such qualifications, it should be

			Velocities (ft/sec)		0	Tomo	Estimated Fouling		
In Tubes	Outside Tubes	Type Equipment	Tube	Shell	Coefficient	Range (°F)	Tube	Shell	Overall
A. Heating-cooling									
Butadiene mix. (Super-heating)	steam	H	25-35	-	12	400-100	_		0.04
Solvent	solvent	Н	_	1.0-l .a	35-40	110-30	—		0.0065
Solvent	propylene (vaporization)	K	I - 2		30-40	40-O		—	0.006
C ₄ unsaturates	propylene (vaporization)	K	20-40		13-18	100-35	-	_	0.005
Solvent	chilled water	Н			35-75	115-40	0.003	0.001	
Oil	oil	H			60-85	150-100	0.0015	0.0015	
Ethylene-vapor	condensate and vapor	K		_	90-125	600-200	0.002	0.001	
Ethylene vapor	chilled water	Н			50-80	270-100	0.001	0.001	
Condensate	propylene (refrigerant)	K-U	—	_	60-135	60-30	0.001	0.001	
Chilled water	transformer oil	Н	—		40-75	75-50	0.001	0.001	
Calcium brine-25%	chlorinated C,	Н	1-2	0.5-I .o	40-60	-20-+10	0.002	0.005	
Ethylene liquid	ethylene vapor	K-U	-		IO-20	-170-(-100)			0.002
Propane vapor	propane liquid	Н		—	6-15	-25-100		—	0.002
Lights and chlor, HC	steam	U			12-30	-30-260	0.001	0.001	
Unsat. light HC, CO, CO,, H ₂	steam	Н	_		IO-2	400-l 00			0.3
Ethonolamine	steam	Н	_		15-25	400-40	0.001	0.001	
Steam	air mixture	U	_		10-20	-30-220	0.0005	0.0015	~
Steam	styrene and tars	U (in tank)	_		50-60	190-230	0.001	0.002	-
Chilled water	freon-12	H	4-7		100-130	90-25	0.001	0.001	
Water ^b	lean copper solvent	Н	4-5	_	100-120	180-90			0.004
Water	treated water	Н	3-5	I-2	100-125	90-l 10			0.005
Water	C ₂ -chlor. HC, lights	Н	2-3		6-10	360-100	0.002	0.001	
Water	hydrogen chloride	Н		_	7-15	230-90	0.002	0.001	
Water	heavy C2-chlor.	Н			45-30	300-90	0.001	0.001	
Water	perchlorethylene	Н		_	55-35	150-90	0.001	0.001	
Water	air and water vapor	Н		-	20-35	370-90	0.0015	0.0015	
Water	engine jacket water	Н	_	_	230-160	175-90	0.0015	0.001	
Water	absorption oil	Н			80-115	130-90	0.0015	0.001	
Water	air-chlorine	U	4-7	_	8-18	250-90	_	_	0.005
Water	treated water	Н	5-7		170-225	200-90	0.001	0.001	-
P. Condensing									
B. Condensing	nronylono rofrig	v	v		E0 C0	60.25			0.005
V ₄ unsat lights	propylene refrig.	n V	v		50-00	45-2		_	0.005
nu ulisal. ligilis Putadiana	propylene reirig.	n V	v	_	50-60	40-0	_	-	0.0055
bulaulelle Uvdrogon oblasida	propylene retrig.	N. U.	v		11 0 60	20-35	0.012	0.001	0.004
nyarogen chioriae	propylene retrig.	м			11 U-60	0-15	0.012	0.001	

TABLE 8.4. Overall Heat Transfer Coefficients in Some Petrochemical Applications, U Btu/(hr)(sqft)(°F)^a

TABLE **8.4**—(continued)

			Velo (ft)	cities (sec)	Overall	Tomp	Esti	mated Fou	uling
In Tubes	Outside Tubes	Type Equipment	Tube	Shell	Coefficient	Range (°F)	Tube	Shell	Overall
Lights and chloro-ethanes	propylene refrig.	КU	-	_	15-25	130-(-20)	0.002	0.001	
Ethylene	propylene refrig.	ΚU	-		60-90	120-(-10)	0.001	0.001	_
Unsat. chloro HC	water	Н	7-a		90-120	145-90	0.002	0.001	
Unsat. chloro HC	water	Н	3-8	_	180-140	11 O-90	0.001	0.001	-
Unsat. chloro HC	water	Н	6		15-25	130–(–20)	0.002	0.001	-
Chloro-HC	water	ΚU		_	20-30	IIO-(-10)	0.001	0.001	-
Solvent and non cond.	water	Н	-	_	25-15	260-90	0.0015	0.004	-
Water	propylene vapor	Н	2-3	_	130-150	200-90	-		0.003
Water	propylene	Н			60-100	130-90	0.0015	0.001	_
Water	steam	H		-	225-110	300-90	0.002	0.0001	-
Water	steam	H	-		190-235	230-130	0.0015	0.001	_
Treated water	steam (exhaust)	H			20-30	220-I 30	0.0001	0.0001	—
Oil	steam	H	-	-	70-I 10	375-130	0.003	0.001	-
Water	propylene cooling and cond	н			25-50	30-45 (C) 🚶	0 0015	0.001	_
Water	propyrene cooring and correct	11			(110-150	15-20 (Co) J	0.0010	0.001	
Chilled water	air-chloring (part and cond)	П			∫ 8-15	8-15 (C) \	0 0015	0 0 0 5	_
onnica water	an enterne (part and cond.)	v			20-30	10-15 (Co) ∫	0.0010	0.000	
Water	light HC, cool and cond.	H	_	-	35-90	270-90	0.0015	0.003	—
Water	ammonia	н		-	140-165	120-90	0.001	0.001	-
Water	ammonia	U	—	-	280-300	11 0-90	0.001	0.001	-
Air-water vapor	freon	КU	_	-	{10-50 }	60-10			0.01
C. Debelling					(10-20)				
C. Rebbiling	stoom	u	7-6	_	120-150	190-160		_	0.005
C unsat	steam	n H	7-0	_	95-1 15	95-1 50	_	_	0.005
C_4 unsat:	steam	" VT		_	35-25	300-350	0.001	0 001	0.0003
Chloro unsat HC	steam	vt	_		100-140	230-130	0.001	0.001	_
Chloro, ethane	steam	VT			90-1 35	300-350	0.001	0.001	
Chloro, ethane	steam	U	—		50-70	30-1 90	0.002	0.001	_
Solvent (heavy)	steam	H	—	_	70-115	375-300	0.004	0.0005	
Mono-di-ethanolamines	steam	vт	—		210-155	450-350	0.002	0.001	_
Organics, acid, water	steam	VT	_	_	60-100	450-300	0.003	0.0005	-
Amines and water	steam	VT	_	_	120-140	360-250	0.002	0.0015	_
Steam	naphtha frac.	Annulus	-	_	15-20	270-220	0.0035	0.0005	
	-	Long. F.N.							
Propylene	C_{2}, C_{2}^{-}	κυ			120-140	150-40	0.001	0.001	_
Propylene-butadiene	butadiene, unsat.	H	-	25-35	15-18	400-100	-	-	0.02

^aFouling resistances are included in the listed values of U. ^bUnless specified, all water is untreated, brackish, bay or sea. Notes: H = horizontal, fixed or floating tuba sheet, U = U-tuba horizontal bundle, K = kettle type, V = vertical, R = reboiler, T = thermosiphon, v = variable, HC = hydrocarbon, (C) = cooling range At, (Co) = condensing range At. (Ludwig, 1983).

Equipment	Process	U
Shell-and-tube exchanger [Fig. 8.4(c)]	gas (1 atm)–gas (1atm) gas (250 atm)–gas (250 atm)	l - 6 25-50
	liquid-gas (1 atm)	2-12
	liquid-gas (250 atm)	35-70
	liquid-liquid	25-200
	liquid-condensing vapor	50-200
Double-pipe exchanger [Fig. 8,4(a)]	gas (1 atm)–gas (1 atm)	2-6
	gas (250 atm)–gas (250 atm)	25-90
	liquid-gas (250 atm)	35-I 00
	liquid-liquid	50-250
Irrigated tube bank	water-gas (1 atm)	3-10
·	water-gas (250 atm)	25-60
	water-liquid	50-160
	water-condensing vapor	50-200
Plate exchanger [Fig. 8.8(a)]	water-gas (1 atm)	3-10
	water-liquid	60-200
Spiral exchanger [Fig. 8.8(c)]	liquid-liquid	120-440
	liquid-condensing steam	160-600
Compact [Fig. 8.6(h)]	gas (1 atm)–gas (1 atm)	2-6
	gas (1atm)–liquid	3-10
Stirred tank, jacketed	liquid-condensing steam	90-260
	boiling liquid-condensing steam	120-300
	water-liquid	25-60
Stirred tank, coil inside	liquid-condensing steam	120-440
	water-liquid	90-210

TABLE 8.5. Ranges of Overall Heat Transfer Coefficients in Various Types of Exchangers [U Btu/(hr)(sqft)(°F)]

^a 1 Btu/(hr)(sqft)(°F) = 5.6745 W/m² K. Data from (HEDH, 1983).

borne in mind that very few proposed correlations are more accurate than $\pm 20\%$ or so.

Along with rate of heat transfer, the economics of practical exchanger design requires that pumping costs for overcoming friction be taken into account.

DIRECT CONTACT OF HOT AND COLD STREAMS

Transfer of heat by direct contact is accomplished in spray towers, in towers with a multiplicity of segmented baffles or plates (called shower decks), and in a variety of packed towers. In some processes heat and mass transfer occur simultaneously between phases; for example, in water cooling towers, in gas quenching with water, and in spray or rotary dryers. Quenching of pyrolysis gases in transfer lines or towers and contacting on some trays in fractionators may involve primarily heat transfer. One or the other, heat or mass transfer, may be the dominant process in particular cases.

Data of direct contact heat transfer are not abundant. The literature has been reviewed by Fair (1972) from whom specific data will be cited.

One rational measure of a heat exchange process is the number of transfer units. In terms of gas temperatures this is defined by

$$N_{g} = \frac{T_{g,\text{in}} - T_{g,\text{out}}}{(T_{g} - T_{L})_{\text{mean}}}$$
(8.33)

The logarithmic mean temperature difference usually is applicable. For example, if the gas goes from 1200 to 150°F and the liquid countercurrently from 120 to 400°F, the mean temperature

difference is 234.5 and $N_g = 4.48$. The height of a contact zone then is obtained as the product of the number of transfer units and the height H_{α} of a transfer unit. Several correlations have been made of the latter quantity, for example, by Cornell, Knapp, and Fair (1960) and modified in the Chemical Engineers Handbook (1973, pp. 18.33, 18.37). A table by McAdams (1954, p. 361) shows that in spray towers the range of H_{ρ} may be 2.5-10 ft and in various kinds of packed towers, 0.4-4 ft or so.

Heat transfer coefficients also have been measured on a volumetric or cross section basis. In heavy hydrocarbon fractionators, Neeld and O'Bara (1970) found overall coefficients of 1360-3480 Btu/(hr)(°F)(sqft of tower cross section). Much higher values have been found in less viscous systems.

Data on small packed columns were correlated by Fair (1972) in the form

$$Ua = CG^{m}L^{n}, \quad Btu/(hr)(cuft)(^{\circ}F), \qquad (8.34)$$

where the constants depend on the kind of packing and the natures of the fluids. For example, with air-oil, 1 in. Raschig rings, in an 8 in. column

$$Ua = 0.083G^{0.94}L^{0.25}.$$
 (8.35)

When G and L are both 5000 lb/(hr)(sqft), for instance, this formula gives Ua = 2093 Btu/(hr)(cuft)(°F).

In spray towers, one correlation by Fair (1972) is

$$h_g a = 0.043 G^{0.8} L^{0.4} / Z^{0.5} \text{ Btu}/(\text{hr})(\text{cuft})(^\circ\text{F}).$$
 (8.36)

TABLE 8.6. Typical Ranges of Individual Film and Fouling Coefficients [h Btu/(hr)(sqft)(°	F)]
---	-----

Fluid and Process	Conditions	P (atm)	(AT),,,,, (°F)	10 ⁴ h	10 ⁴ h _r
Sensible					
Water	liquid			7.6-11.4	6-14
Ammonia	liquid			7.1-9.5	O-6
Light organics	liquid			28-38	6-11
Medium organics	liquid			38-76	9-23
Heavy organics	liquid heating			23-76	11-57
Heavy organics	liquid cooling			142-378	11-57
Very heavy organics	liquid heating			189-568	23-170
Very heavy organics	liquid cooling			378-946	23-170
Gas		I - 2		450-700	O-6
Gas		10		140-230	O-6
Gas		100		57-113	O-6
Condensing transfer					
Steam ammonia	all condensable	0.1		4.7-7.1	O-6
Steam ammonia	1% noncondensable	0.1		9. 5-14. 2	O-6
Steam ammonia	4% noncondensable	0.1		19-28	O-6
Steam ammonia	all condensable	1		3.8-5.7	O-6
Steam ammonia	all condensable	10		2.3-3.8	O-6
Light organics	pure	0.1		28-38	O-6
Light organics	4% noncondensable	0.1		57-76	O-6
Light organics	pure	10		8-19	O-6
Medium organics	narrow range	1		14-38	6-30
Heavy organics	narrow range			28-95	11-28
Light condensable mixes	narrow range	1		23-57	0-11
Medium condensable mixes	narrow range	1		38-95	6-23
Heavy condensable mixes	medium range	1		95-190	11-45
Vaporizing transfer					
Water		<5	4 5	5. 7- 19	6-12
Water		<100	36	3.8-14	6-12
Ammonia		<30	36	11-19	6-12
Light organics	pure	20	36	14-57	6-12
Light organics	narrow range	20	27	19-76	6-17
Medium organics	pure	20	36	16-57	6-17
Medium organics	narrow range	20	27	23-95	6-17
Heavy organics	pure	20	36	23-95	11-28
Heavy organics	narrow range	20	27	38-142	11-45
Very heavy organics	narrow range	2 0	27	57-189	11-57

Light organics have viscosity <1 cP, typically similar to octane and lighter hydrocarbons. Medium organics have viscosities in the range 1-5 cP, like kerosene, hot gas oil, light crudes, etc. Heavy organics have viscosities in the range 5-100 cP, cold gas oil, lube oils, heavy and reduced crudes, etc. Very heavy organics have viscosities above 100 cP, asphalts, molten polymers, greases, etc. Gases are all noncondensables except hydrogen and helium which have higher coefficients. Conversion factor: 1 Btu/(hr)(sqft)(°F) = 5.6745 W/m² K. (After HEDH, 1983, 3.1.4-4).

In a tower with height Z = 30ft and with both G and L at 5000 lb/(hr)(cuft), for example, this formula gives $h_g a = 21.5$.

In liquid-liquid contacting towers, data cited by Fair (1972) range from 100–12,000 Btu/(hr)(cuft)(°F) and heights of transfer units in the range of 5 ft or so. In pipeline contactors, transfer rates of 6000–60,000 Btu/(hr)(cuft)(°F) have been found, in some cases as high as 200,000.

In some kinds of equipment, data only on mass transfer rates may be known. From these, on the basis of the Chilton-Colburn analogy, corresponding values of heat transfer rates can be estimated.

NATURAL CONVECTION

Coefficients of heat transfer by natural convection from bodies of various shapes, chiefly plates and cylinders, are correlated in terms of Grashof, Prandtl, and Nusselt numbers. Table 8.9 covers the most usual situations, of which heat losses to ambient air are the most common process. Simplified equations are shown for air. Transfer of heat by radiation is appreciable even at modest temperatures; such data are presented in combination with convective coefficients in item 16 of this table.

FORCED CONVECTION

Since the rate of heat transfer is enhanced by rapid movement of fluid past the surface, heat transfer processes are conducted under such conditions whenever possible. A selection from the many available correlations of forced convective heat transfer involving single phase fluids, including flow inside and outside bare and extended surfaces, is presented in Table 8.10. Heat transfer resulting in phase change, as in condensation and vaporization, also is covered in this table. Some special problems that arise in interpreting phase change behavior will be mentioned following.

		Kind of Heat	Transfer
Fluid	Sensible	Boiling	Condensing
Aromatic liquids			
Benzene, toluene, ethylbenzene,			
styrene	0.007	0.011	0.007
Dowtherm	0.007		—
Inorganic solutions			
CaCl ₂ Brine (25%)	0.004		_
Heavy acids	0.013	—	
NaCl Brine (20%)	0.0035	_	
Misc. dilute solutions	0.005	_	
Light hydrocarbon liquids			
C ₃ , C ₄ , C ₅	0.004	0.007	0.004
Chlorinated hydrocarbons	0.004	0.009	0.007
Miscellaneous organic liquids			
Acetone	0.007		_
Amine solutions			
Saturated diethanolamine and MONO-			
ethanolamine (CO, and H_2S)	0.007	_	-
Lean amine solutions	0.005	<u></u>	_
Oils			
Crude oil	0.015	_	-
Diesel oil	0.011	_	-
Fuel oil (bunker C)	0.018		_
Gas oil			
Light	0.0125		0.015
Heavy (typical of cat. cracker feed)	0.014		0.018
Gasoline (400" EP)	0.008	0.010	0.008
Heating oil (domestic 30° API)	0.010	-	_
Hydroformate	0.006	_	
Kerosine	0.009	_	0.013
Lube oil stock	0.018	_	
Naphthas			
Absorption	0.008	0.010	0.006
Liaht virain	0.007	0.010	0.007
Light catalytic	0.006	0.010	0.007
Heavy	0.008	0.011	0.0085
Polymer (C _e 's)	0.008	0.010	0.008
Reduced crude	0.018		_
Slurry oil (fluid cat. cracker)	0.015		_
Steam (no noncondensables)			0.001
Water			
Boiler water	0.003	_	<u>-</u> -
Cooling tower (untreated)	0.007	_	
Condensate (flashed)	0.002	_	
River and well	0.007		_
Sea water (clean and below 125°F)	0.004	_	
Gases in turbulent flow			
Air, CO, CO,, and N_2	0.045		
Hydrocarbons (light through naphthas)	0.035		

TABLE 8.7. Individual Film Resistances (I/h) Including Fouling Effects, with **h** in **Btu/(hr)(sqft)(°F)**

(Fair and Rase, Pet Refiner 33 (7), 121, 1854; Rase and Barrow, Project Engineering of Process Plants; 224, Wiley, 1957.)

CONDENSATION

Depending largely on the nature of the surface, condensate may form either a continuous film or droplets. Since a fluid film is a partial insulator, **dropwise** condensation results in higher rates of condensation. Promoters are substances that make surfaces nonwetting, and may be effective as additives in trace amounts to the vapor. Special shapes of condensing surfaces also are effective in developing **dropwise** condensation. None of these effects has been generally correlated, but many examples are cited in HEDH and elsewhere. Condensation rates of mixtures are influenced by both heat and mass transfer rates; techniques for making such calculations have been developed and are a favorite problem for implementation on computers. Condensation rates of mixtures that form immiscible liquids also are reported on in HEDH. Generally, mixtures have lower heat transfer coefficients in condensation than do pure substances.

BOILING

This process can be nuclear or film type. In nuclear boiling, bubbles detach themselves quickly from the heat transfer surface. In film boiling the rate of heat transfer is retarded by an adherent vapor film through which heat supply must be by conduction. Either mode

 TABLE 8.8. Dimensionless Groups and Units of Quantities

 Pertaining to Heat Transfer

Symbol	Number	Group
Bi	Biot	hL/k
Fo	Fourier	kθ/ρCL ²
Gz	Graetz	wC/kL
Gr	Grashof	$D^3 \rho^2 g \beta \Delta T / \mu^2$
NU	Nusselt	hD/k
Pe	Peclet	DGC/k = (Re)(Pr)
Pr	Prandtl	$C\mu/k$
Re	Reynolds	DG/μ, Duρ/μ
S C	Schmidt	$\mu/\rho k_d$
St	Stanton	hC/G = (Nu)/(Re)(Pr)
Notation	Name	and Typical Units
С	heat capacity [Btu/(lb)	(°F), cal/(g)(°C)]
D	diameter (ft, m)	
g	acceleration of gravity	$[ft/(hr)^2, m/sec^2]$
Ğ	mass velocity [lb/(hr)($(t)^2, kg/sec)(m)^2$
h	heat transfer coefficien W/(m) ² (sec)]	t [Btu/(hr)(sqft)(°F),
k	thermal conductivity [cal/(sec)(cm ²)(C/cm)	Btu/(hr)(sqft)(°F/ft),)]
k _d	diffusivity (volumetric)	[ft ² /hr, cm ² /sec]
L	length (ft, cm)	
T, AT	temperature, temperature	difference (°For °R, ℃ or K)
u	linear velocity (ft/hr, c	m/sec)
U	overall heat coefficient	(same as units of <i>h</i>)
W	mass rate of flow (lb/l	hr, g/sec)
β	Thermal expansion coe	fficient (1/°F, 1/°C)
θ	time (hr. Sec)	
μ	viscosity [lb/(ft)(hr), g	/(cm)(sec)]
Р	density [lb/(ft) ³ , g/(cm) ³]

can exist in any particular case. Transition between modes corresponds to a maximum heat flux and the associated critical temperature difference. A table of such data by McAdams (Heat *Transmission*, McGraw-Hill, New York, 1954, p. 386) shows the critical temperature differences to range from 42-90°F and the maximum fluxes from 42-126 KBtu/(hr)(sqft) for organic substances and up to 410 KBtu/(hr)(sqft) for water; the nature of the surface and any promoters are identified. Equations (40) and (41) of Table 8.10 are for critical heat fluxes in kettle and thermosyphon reboilers. Beyond the maximum rate, film boiling develops and the rate of heat transfer drops off very sharply.

Evaluation of the boiling heat transfer coefficient in vertical tubes, as in thermosyphon reboilers, is based on a group of equations, (42)-(48), of Table 8.10. A suitable procedure is listed following these equations in that table.

EXTENDED SURFACES

When a film coefficient is low as in the cases of low pressure gases and viscous liquids, heat transfer can be improved economically by employing extended surfaces. Figure 8.6 illustrates a variety of extended surfaces. Since the temperature of a fin necessarily averages less than that of the bare surface, the effectiveness likewise is less than that of bare surface. For many designs, the extended surface may be taken to be 60% as effective as bare surface, but this factor depends on the heat transfer coefficient and thermal conductivity of the fin as well as its geometry. Equations and corresponding charts have been developed for the common geometries and are shown, for example, in HEDH (1983, Sec. 2.5.3) and elsewhere. One chart is given with Example 8.6. The efficiency η of the extended surface is defined as the ratio of a realized heat transfer to the heat transfer that would be obtained if the fin were at the bare tube temperature throughout. The total heat transfer is the sum of the heat transfers through the bare and the extended surfaces:

$$Q = Q_b + Q_e = U_b A_b (1 + \eta A_e / A_b) (T_b - T_{\text{fluid}}).$$

A, is the tube surface that is not occupied by fins. Example 8.6 performs an analysis of this kind of problem.

8.5. PRESSURE DROP IN HEAT EXCHANGERS

Although the rate of heat transfer to or from fluids is improved by increase of linear velocity, such improvements are limited by the economic balance between value of equipment saving and cost of pumping. A practical rule is that pressure drop in vacuum condensers be limited to OS-1.0 psi (25-50 Torr) or less, depending on the required upstream process pressure. In liquid service, pressure drops of 5-10 psi are employed as a minimum, and up to 15% or so of the upstream pressure.

Calculation of tube-side pressure drop is straightforward, even of vapor-liquid mixtures when their proportions can be estimated. Example 8.7 employs the methods of Chapter 6 for pressure drop in a thermosiphon reboiler.

The shell side with a number of segmental baffles presents more of a problem. It may be treated as a series of ideal tube banks connected by window zones, but also accompanied by some bypassing of the tube bundles and leakage through the baffles. A hand calculation based on this mechanism (ascribed to K.J. Bell) is illustrated by Ganapathy (1982, pp. 292–302), but the calculation usually is made with proprietary computer programs, that of HTRI for instance.

A simpler method due to Kern (1950, pp. 147-152) nominally considers only the drop across the tube banks, but actually takes account of the added pressure drop through baffle windows by employing a higher than normal friction factor to evaluate pressure drop across the tube banks. Example 8.8 employs this procedure. According to Taborek (HEDH, 1983, 3.3.2), the Kern predictions usually are high, and therefore considered safe, by a factor as high as 2, except in laminar flow where the results are uncertain. In the case worked out by Ganapathy (1982, pp. 292–302), however, the Bell and Kern results are essentially the same.

8.8. TYPES OF HEAT EXCHANGERS

Heat exchangers are equipment primarily for transferring heat between hot and cold streams. They have separate passages for the two streams and operate continuously. They also are called recuperators to distinguish them from regenerators, in which hot and cold streams pass alternately through the same passages and exchange heat with the mass of the equipment, which is intentionally made with large heat capacity. Recuperators are used mostly in cryogenic services, and at the other extreme of temperature, as high temperature air preheaters. They will not be discussed here; a detailed treatment of their theory is by Hausen (1983).

Being the most widely used kind of process equipment is a claim that is made easily for heat exchangers. A classified directory of manufacturers of heat exchangers by Walker (1982) has several hundred items, including about 200 manufacturers of shell-and-tube equipment. The most versatile and widely used exchangers are the shell-and-tube types, but various plate and other types are valuable and economically competitive or superior in some applications. These other types will be discussed briefly, but most of the space following will be devoted to the shell-and-tube types, primarily

(16)

TABLE 8.9. Equations for Heat Transfer Coefficients of Natural Convection

Vertical plates and cylinders, length	
$X_{L} = (Gr)(Pr) = \left(\frac{L^{2}\rho_{f}^{2}g\beta_{f}\Delta t}{\mu_{f}^{2}}\right) \left(\frac{c_{p}\mu}{\kappa}\right)_{f}$	(1)
$hL/k = 0.13X_L^{1/3}$, turbulent, $10^9 < X_1 < 10^{12}$	(2)
$h = 0.19(\Delta t)^{1/3}$, for air, At in "F, h in Btu/(hr)(sqft)(°F)	(3)
$hL/k = 0.59X_L^{1/4}$, laminar, $10^4 < X_i < 10^9$	(4)
$h = 0.29(\Delta t/L)^{1/4}$, for air, L in ft	(5)

Single horizontal cylinder, diameter D_0

$$X_{D} = \frac{D_{0}^{3} \rho_{s}^{2} g \beta_{s} \Delta t}{\mu_{s}^{2}} \left(\frac{c_{p} \mu}{k}\right)$$

$$h D_{0} / k \approx 0.53 X_{D}^{1/4}, \ 10^{3} < X_{D} < 10^{9}$$

$$h = 0.18 (\Delta t)^{1/3}, \ \text{for air, } \ 10^{9} < X_{D} < 10^{12}$$

$$h = 0.27 (\Delta t / D_{0})^{1/4}, \ 10^{4} < X_{D} < 10^{9}$$
(6)
(7)
(8)
(9)

Horizontal plates, rectangular, l the smaller dimension

$$X_{L} = \frac{L^{3} \rho_{f}^{2} g \beta_{f} \Delta t}{\mu_{f}^{2}} \left(\frac{c_{p} \mu}{k}\right)_{f}$$
(10)

Heated plates facing up or cooled facing down

$hL/k = 0.14X_{L}^{1/3}, 2(10^7) < X_L < 3(10^{10}), \text{ turbulent}$	(11)
$h = 0.22(\Delta t)^{1/3}$, for air	(11)
$hL/k = 0.54X_L^{1/4}, 10^\circ < X_L < 2(10'), \text{laminar}$	(12)
$h = 0.27 (\Delta t/L)^{1/4}$	(13)

Heated plates facing down, or cooled facing up

$hL/k = 0.27X_L^{1/4}, 3(10^5) < X_L < 3(10^{10}),$ laminar	(14)
/⊨ 0.12(∆ <i>t/L</i>) ^{1/4} , for air	(15)

Combined convection and radiation coefficients, $h_c + h_u$ for horizontal steel or insulated pipes in a room at 80°F

Nominal	(At),, Temperature Difference (°F) from Surface to Room														
(in.)	50	100	150	200	250	300	400	500	600	700	600	900	1000	1100	1200
12	2.12	2.48	2.76	3.10	3.41	3.75	4.47	5.30	6.21	7.25	8.40	9.73	11.20	12.81	14.65
Ť	2.03	2.38	2.65	2.98	3.29	3.62	4.33	5.16	6.07	7.11	8.25	9.57	11.04	12.65	14.48
2	1.93	2.27	2.52	2.85	3.14	3.47	4.18	4.99	5.89	6.92	8.07	9.38	10.85	12.46	14.28
4	1.84	2.16	2.41	2.72	3.01	3.33	4.02	4.83	5.72	6.75	7.89	9.21	10.66	12.27	14.09
8	1.76	2.06	2.29	2.60	2.89	3.20	3.88	4.68	5.57	6.60	7.73	9.05	10.50	12.10	13.93
12	1.71	2.01	2.24	2.54	2.82	3.13	3.83	4.61	5.50	6.52	7.65	8.96	10.42	12.03	13.84
2 4	1.64	1.93	2.15	2.45	2.72	3.03	3.70	4.48	5.37	6.39	7.52	8.83	10.28	11.90	13.70

(McAdams, Heat Transmission, McGraw-Hill, New York, 1954).

because of their importance, but also because they are most completely documented in the literature. Thus they can be designed with a degree of confidence to fit into a process. The other types are largely proprietary and for the most part must be process designed by their manufacturers.

PLATE-AND-FRAME EXCHANGERS

Plate-and-frame exchangers are assemblies of pressed corrugated plates on a frame, as shown on Figure 8.8(a). Gaskets in grooves around the periphery contain the fluids and direct the flows into and out of the spaces between the plates. Hot and cold flows are on opposite sides of the plates. Figure 8.8(b) shows a few of the many combinations of parallel and countercurrent flows that can be maintained. Close spacing and the presence of the corrugations result in high coefficients on both sides-several times those of shell-and-tube equipment-and fouling factors are low, of the order of $1-5 \times 10^{-5}$ Btu/(hr)(sqft)(°F). The accessibility of the heat exchange surface for cleaning makes them particularly suitable for fouling services and where a high degree of sanitation is required, as in food and pharmaceutical processing. Operating pressures and temperatures are limited by the natures of the available gasketing materials, with usual maxima of 300 psig and 400°F.

Since plate-and-frame exchangers are made by comparatively few concerns, most process design information about them is proprietary but may be made available to serious enquirers. Friction factors and heat transfer coefficients vary with the plate spacing and the kinds of corrugations; a few data are cited in HEDH (1983, 3.7.4-3.7.5). Pumping costs per unit of heat transfer are said to be lower than for shell-and-tube equipment. In stainless steel

TABLE 8.10. Recommended Individual Heat Transfer Coefficient Correlations'

A. Single Phase Streams

a. Laminar Flow, Re $\,<\,$ 2300 Inside tubes

Nu, =
$$\sqrt[3]{3.66^3 + 1.61^3 \text{Pe}(d/L)}$$
, 0.1 < Pe(d/L) < 10⁴ (1)

Between parallel plates of length L and separation distance s

Nu, =
$$3.78 + \frac{0.0156[\operatorname{Pe}(s/L)]^{1.14}}{1 + 0.058[\operatorname{Pe}(s/L)]^{0.64}\operatorname{Pr}^{0.177}}$$
 0.1 < $\operatorname{Pe}(s/L) < 10^3$ (2)

In concentric annuli with d_i inside, d_o outside, and hydraulic diameter $d_h = d_o - d_i$. I, heat transfer at inside wall; ||, at outside wall; III, at both walls at equal temperatures'

$$Nu_{T} = Nu_{\infty} + f\left(\frac{d_{i}}{d_{o}}\right) \frac{0.19[Pe(d_{h}/L)]^{0.8}}{1 + 0.117[Pe(d_{h}/L)]^{0.467}}$$
(3)

Case I:
$$Nu_{i\infty} = 3.66 + 1.2 \left(\frac{d_i}{d_o}\right)^{-0.5}$$
 (4)

Case II:
$$Nu_{ox} = 3.66 + 1.2 \left(\frac{d_i}{d_o}\right)^{0.5}$$
 (5)

Case III:
$$Nu_b = 3.66 + \left[4 - \frac{0.102}{(d_i/d_o) + 0.2}\right] \left(\frac{d_i}{d_o}\right)^{0.04}$$
 (6)

Case I:
$$f\left(\frac{d_i}{d_o}\right) = 1 + 0.14 \left(\frac{d_i}{d_o}\right)^{-0.5}$$
 (7)

Case II:
$$f\left(\frac{d_i}{d_o}\right) = 1 + 0.14 \left(\frac{d_i}{d_o}\right)^{1/3}$$
 (8)

Case III:
$$f\left(\frac{d_i}{d_o}\right) = 1 + 0.14 \left(\frac{d_i}{d_o}\right)^{0.1}$$
 (9)

b. Turbulent Flow, Re > 2300

Inside tubes

Nu = 0.0214(Re^{0.8} - 100) Pr^{0.4}
$$\left[1 + \left(\frac{d}{L} \right)^{2/3} \right]$$
, 0.5 < Pr < 1.5 (10)

Nu = 0.012(Re^{0.87} 280) Pr^{0.4}
$$\left[1 + \left(\frac{d}{L}\right)^{2/3}\right]$$
, 1.5 < Pr < 500 (11)

Concentric annuli: Use d_h for both Re and Nu. Nu_{tube} from Eqs. (10) or (11)

Case I:
$$\frac{\mathrm{Nu}_{i}}{\mathrm{Nu}_{\mathrm{tube}}} = 0.86 \left(\frac{d_{i}}{d_{o}}\right)^{-0.16}$$
Case II:
$$\frac{\mathrm{Nu}_{o}}{\mathrm{Nu}_{\mathrm{tube}}} = 1 - 0.14 \left(\frac{d_{i}}{d_{o}}\right)^{0.6}$$
(13)

Case III:
$$\frac{Nu_b}{Nu_{tube}} = \frac{0.86(d_i/d_o)^{0.84} + [1 - 0.14(d_i/d_o)^{0.6}]}{1 + d_i/d_o}$$
(14)

Across one row of long tubes: d = diameter, s = center-to-center distance, a = s/d, $\psi = 1 - \pi/4a$, $L = \pi d/2$

$$\begin{array}{ll} {\rm Re}_{\psi,L} = wL/\psi\nu & (15) \\ {\rm Nu}_{o,row} = 0.3 + \sqrt{{\rm Nu}_{L,iam}^2 + {\rm Nu}_{L,turb}^2} & (161) \\ {\rm Nu}_{L,lam} = 0.664\sqrt{{\rm Re}_{\psi,L}} {\rm Pr}^{1/3} & (17) \\ {\rm Nu}_{L,turb} = 0.037 \ {\rm Re}_{\psi,L}^{0.8} {\rm Pr}/[1 + 2.443 \ {\rm Re}_{\psi,L}^{-0.1} ({\rm Pr}^{2/3} - 1)] & (18) \\ {\rm Nu}_{L,row} = \alpha L/\lambda & (19) \end{array}$$

[&]quot;Special notation used in this table: α = heat transfer coefficient (W/m² K) (instead of h), η = viscosity (instead of μ), and α = thermal conductivity (instead of k). (Based on HEDH, 1983).

TABLE 8.10—(continued)

Across a bank of n tubes deep:



ψ=1−π/4a ifb≧	≥1	(20)
$\psi=1-\pi/4ab$ if	b<1	(21)

$$Nu_{o,bank} = \alpha L/\lambda = f_A Nu_{o,row}/K, \quad n \ge 10$$

$$Nu_{o,bank} = [1 + (n - 1)f_A] Nu_{o,row}/Kn, \quad n < 10$$
(22)
(23)

[Nu_{o,row} from Eq. (16)]

$f_{A,\text{in-line}} = 1 + (0.7/\psi^{1.5})[(b/a 0.3)/(b/a + 0.7)^2]$	(24)
$f_{A,\text{stag}} = 1 + 2/3b$	(25)
$K = (\Pr/\Pr_w)^{0.25}$, for liquid heating	(26)
$K = (P_r / P_r)^{10.11}$, for liquid cooling	(27)
$K = (T/T_w)^{0.12}$, for gases	(28)

Subscript w designates wall condition

Banks of radial high-fin tubes: $\varepsilon = (bare tube surface)/(total surface of finned tube)$

In line:

$Nu = 0.30 \text{Re}^{0.625} \varepsilon^{-0.375} \text{Pr}^{0.333}$	5<ε<12,	5000 < Re < 10 ⁵	(29)
--	---------	-----------------------------	------

Staggered: $a = s_1/d$, $b = s_2/d$, s = spacing of fins

$$Nu = 0.19(a/b)^{0.2}(s/d)^{0.18}(h/d)^{-0.14} \operatorname{Re}^{0.65} \operatorname{Pr}^{0.33}, \quad 100 < \operatorname{Re} < 20,000$$
(30)

Banks of radial low-fin tubes: D = diameter of finned tube, s = distance between fins, h = height of fin; following correlation for D = 22.2 mm, s = 1.25 mm, and h = 1.4 mm

Nu = 0.0729 Re ^{0.74} Pr ^{0.36} , 5000 < Re < 35,000	(31)
Nu = 0.137 Re ^{0.68} Pr ^{0.36} , 35,000 < Re < 235,000	(32)
$Nu = 0.0511 \text{ Re}^{0.76} \text{ Pr}^{0.36}$, 235,000 < Re < 10 ⁶	(33)

B. Condensation of Pure Vapors

On vertical tubes and other surfaces; $\dot{\Gamma}$ = condensation rate per unit of periphery

$$\frac{\tilde{\alpha}}{\lambda_l} \left[\frac{\eta_l^2}{\rho_l (\rho_l - \rho_a) g_n} \right]^{1/3} = 1.47 \left(\frac{4\dot{\Gamma}}{\eta_l} \right)^{-1/3}$$
(34)

On a single horizontal tube: Γ = condensation rate per unit length of tube

$$\frac{\alpha}{\lambda_l} \left[\frac{\eta_l^2}{\rho_l (\rho_l - \rho_g) g_n} \right]^{1/3} = 1.51 \left(\frac{4\dot{\Gamma}}{\eta_l} \right)^{-1/3}$$
(35)
(continued)

TABLE 8.10—(continued)

On a bank of N horizontal tubes: Γ = condensation rate per unit length from the bottom tube

$$\frac{\alpha}{\lambda_{l}} \left[\frac{\eta_{l}^{2}}{\rho_{l}(\rho_{l} - \rho_{g})g_{n}} \right]^{1/3} = 1.51 \left(\frac{4\dot{\Gamma}}{\eta_{l}} \right)^{-1/3} N^{-1/6}$$
(36)

C. Boiling

Single	immersed	tube: ġ l	heat flux	(W/m²),	$p_c = critical$	pressure,	bars. p _r	$= p/p_c$
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$$\alpha = 0.1000 \dot{q}^{0.7} p^{0.69} [1.8 p_r^{0.17} + 4 p_r^{1.2} + 10 p_r^{10}], \quad W/m^2 \text{ K}$$
(37)

Kettle and horizontal thermosiphon reboilers

$$\alpha = 0.27 \exp(-0.027BR)\dot{q}_{c}^{0.7} p_{c}^{0.69} p_{r}^{0.17} + \alpha_{nc}$$
(38)

BR = difference between dew and bubblepoints (°K); if more than 85, use 85

$$\alpha_{nc} = \frac{250 \text{ W/m}^2 \text{ K, for hydrocarbons}}{11000 \text{ W/m K, for water}}$$
(39)

Critical heat flux in kettle and horizontal thermosiphon reboilers

$$q_{\max} = 80,700 p_c p_c^{0.35} (1 - p_r)^{0.9} \psi_{b'} \quad W/m^2$$
(40)

 ψ_b = (external peripheral surface of tube bundle)/ (total tube area); if >0.45, use 0.45

Boiling in vertical tubes: thermosiphon reboilers

Critical heat flux: p_c critical pressure, bars; D_i tube ID, m; L tube length, m

$$\dot{q} = 393,000(D_i^2/L)^{0.35} p_c^{0.61} p_r^{0.25} (1 p_r), W/m^2$$
(41)

Heat transfer coefficient with Eqs. (42)-(48) and following procedure

$$\alpha_{tp} = \alpha_{nb} + \alpha_c \tag{42}$$

$$(\dot{m}(1-x)D)^{0.8}/nC_{\rm e})^{0.4}\lambda_c$$

$$\alpha_c = 0.023 \left(\frac{1}{\eta_i}\right) \left(\frac{1}{\lambda}\right)_i \frac{1}{D} F$$
(43)

$$\alpha_{nb} = 0.00122 \left(\frac{\lambda_l^{0.75} C_{pl}^{0.35} \rho_l^{0.32}}{\sigma^{0.5} \eta_l^{0.29} \Delta h_v^{0.24} \rho_g^{0.24}} \right) \Delta T_{\text{sat}}^{0.24} \Delta p_{\text{sat}}^{0.75} S$$
(44)

$$F = 1 \quad \text{for } 1/\Lambda_{tt} \le 0.1 \tag{43}$$

$$F = 2.35(1/X) + 0.213^{10.736} \quad \text{for } 1/X > 0.1 \tag{46}$$

$$F = 2.55(1/\lambda_{tt} + 0.213) \qquad (40)$$

$$S = 1/(1 + 2.53) \qquad 10^{-6} \text{Re}^{1.17} \qquad (47)$$

$$\begin{aligned} & \mathcal{S} = \{ (1 - x)/x \}^{0.9} (\rho_g/\rho_l)^{0.5} (\eta_l/\eta_g)^{0.1} \end{aligned} \tag{48}$$

Procedure for finding the heat transfer coefficient and required temperature difference when the heat flux \dot{q} , mass rate of flow \dot{m} and fraction vapor χ are specified

1. Find X_{tt}, Eq. (48)

- 2. Evaluate F from Eqs. (45), (46)
- 3. Calculate α_c , Eq. (43)
- 4. Calculate $\operatorname{Re}_{tp} = mF^{1.25}(1 \times x)D/\eta_1$
- 5. Evaluate S from Eq. (47)
- 6. Calculate α_{nb} for a range of values of AT,,,
- 7. Calculate α_{tp} from Eq. (42) for this range of AT, , , values
- 6. On a plot of calculated $\dot{q} = \alpha_{tp} \Delta T_{sat}$ against $\alpha_{tp'}$ find the values of α_{tp} and ΔT_{sat} corresponding to the specified \dot{q}

EXAMPLE 8.6

Sizing an Exchanger with Radial Finned Tubes

A liquid is heated from 150 to 190°F with a gas that goes from 250 to 200°F. The duty is 1.25 MBtu/hr. The inside film coefficient is 200, the bare tube outside coefficient is $h_b = 20$ Btu/(hr)(sqft)(°F). The tubes are 1 in. OD, the fins are $\frac{5}{8}$ in. high, 0.038 in. thick, and number 72/ft. The total tube length will be found with fins of steel, brass, or aluminum:

LMTD = $(60 - 50)/\ln(60/50) = 54.8$, $U_{\rm b} = (1/20 + 1/200)^{-1} = 18.18$.

Fin surface:

 $A_{2} = 72(2)(\pi/4)[(2.25^{2}-1)/144] = 3.191 \text{ sqft/ft}.$

Uncovered tube surface:

 $A_{s} = (\pi/12)[1 - 72(0.038/12)] = 0.2021 \text{ sqft/ft},$ $A_{e}/A_{b} = 3.191/0.2021 = 15.79,$ $y_{b} = \text{half-fin thickness} = 0.038/2(12) = 0.00158 \text{ ft}.$

Abscissa of the chart:

 $\begin{aligned} \mathbf{x} &= (r_e - r_b) \sqrt{h_b / y_b k} = [(2.25 \quad 1)/24] \sqrt{20/0.00158k} \\ &= 5.86 / \sqrt{k}, \\ r_e / r_b &= 2.25, \\ A_b &= Q / U_b \Delta T (1 + \eta A_e / A_b) \\ &= 1.25 (10^6) / 18.18 (54.8) (1 + 15.7917) \text{ sq ft.} \end{aligned}$

Find η from the chart. Tube length, $L = A_b/0.2021$ ft.

	k	x	η	А,,	L
Steel	26	1.149	0.59	121.6	602
Brass	60	0.756	0.76	96.5	477
Al	120	0.535	0.86	86.1	426



EXAMPLE 8.7

Pressure Drop on the Tube Side of a Vertical Thermosiphon Reboiler

Liquid with the properties of water at 5 atm and $307^{\circ}F$ is reboiled at a feed rate of 2800 lb/(hr)(tube) with 30wt % vaporization. The tubes are 0.1 ft ID and 12 ft long. The pressure drop will be figured at an average vaporization of 15%. The Lockhart-Martinelli, method will be used, following Example 6.14, and the formulas of Tables 6.1 and 6.8:

$$\begin{array}{c|c} Liquid & Vapor\\ \hline \textbf{m} (lb/hr) & 2380 & 420\\ \mu (lb/fthr) & 0.45 & 0.036\\ \rho (lb/cuft) & 57.0 & 0.172\\ Re & 67340 & 148544\\ f & 0.0220 & 0.0203\\ \Delta P/L (psi/ft) & 0.00295 & 0.0281\\ \hline X^2 = 0.00295/0.0281 = 0.1051,\\ C = 20,\\ \phi_L^2 = 1 + 20/X + 1/X^2 = 72.21,\\ (\Delta P/L) \text{ two phase} = 72.21(0.00295) = 0.2130,\\ AP = 0.2130(12) = 2.56 \text{ psi}, 5.90 \text{ ft water.} \end{array}$$



Average density in reboiler tubes is

$$\rho_{m} = \frac{2800}{2380/57 + 420/0.172} = 1.13 \text{ lb/cuft}$$

Required height of liquid in tower above bottom of tube sheet

$$\rho_L h = 2.56(144) + 1.13(12),$$

h = 382.2/57 = 6.7 ft.

EXAMPLE 8.8 Pressure Drop on the Shell Side with 25% Open Segmental Baffles, by Kern's Method (1950, p. 147) Nomenclature and formulas:

hydraulic diameter $D_h = \begin{cases} 1.1028P_t^2/D_t - D_t, & \text{triangular pitch,} \\ 1.2732P_t^2/D_t - D_t, & \text{square pitch,} \end{cases}$



 $D_{s} = \text{distance between baffles,}$ N = number of baffles, $A_{r} = \text{flow area} = D_{s}BC/P_{t},$ $G_{s} = \dot{m}/A_{s}, \quad \text{lb}/(\text{hr})(\text{sqft}),$

$$Re = D_h G_s / \mu,$$

$$f = 0.0121 Re^{-0.19}, \quad 300 < Re < 10^6, \quad 25\% \text{ segmental baffles}$$

$$\Delta P = \frac{f G_s^2 D_s (N+1)}{2g \rho D_h} = \frac{f G_s^2 D_s (N+1)}{5.22(10^{10}) s D_h}, \quad \text{psi},$$

$$s = \text{specific gravity.}$$

Numerical example:

tit = 43,800 lb/hr, s = 0.73 sp gr, $\mu = 0.097$ lb/ft hr, $D_t = 1$ in., $P_t = 1.25$ in., triangular pitch, C = 1.25 - 1.00 = 0.25 in., $D_s = 21.25$ in., 1.77 ft., $D_h = 0.723$ in., 0.0603 ft., B = 5 in., N = 38 baffles, $A_r = 21.25(0.25)(5)/1.25(144) = 0.1476$ sqft, $G_s = 43,800/0.1476 = 296,810$ lb/(hr)(sqft), Re = 0.0603(296,810)/0.97 = 18,450, $f = 0.0121(18,450)^{-0.19} = 0.00187,$ $\Delta P = \frac{0.00187(296,810)^2(1.77)(39)}{5.22(10^{10})(0.73)(0.0603)} = 4.95$ psi.

construction, the plate-and-frame construction cost is 50-70% that of shell-and-tube, according to Marriott (*Chem. Eng.*, April 5, 1971).

A process design of a plate-and-frame exchanger is worked out by Ganapathy (1982, p. 368).

SPIRAL HEAT EXCHANGERS

As appears on Figure 8.8(c), the hot fluid enters at the center of the spiral element and flows to the periphery; flow of the cold fluid is countercurrent, entering at the periphery and leaving at the center. Heat transfer coefficients are high on both sides, and there is no correction to the log mean temperature difference because of the true countercurrent action. These factors may lead to surface requirements 20% or so less than those of shell-and-tube exchangers. Spiral types generally may be superior with highly viscous fluids at moderate pressures. Design procedures for spiral plate and the related spiral tube exchangers are presented by Minton (1970). Walker (1982) lists 24 manufacturers of this kind of equipment.

COMPACT (PLATE-FIN) EXCHANGERS

Units like Figure 8.6(h), with similar kinds of passages for the hot and cold fluids, are used primarily for gas service. Typically they have surfaces of the order of $1200 \text{ m}^2/\text{m}^3$ (353 sqft/cuft), corrugation height 3.8-11.8 mm, corrugation thickness 0.2-0.6 mm, and fin density 230-700 fins/m. The large extended surface permits about four times the heat transfer rate per unit volume that can be achieved with shell-and-tube construction. Units have been designed for pressures up to 80 atm or so. The close spacings militate against fouling service. Commercially, compact exchangers are used in cryogenic services, and also for heat recovery at high temperatures in connection with gas turbines. For mobile units, as

in motor vehicles, the designs of Figures 8.6(h) and (i) have the great merits of compactness. and light weight. Any kind of arrangement of cross and countercurrent flows is feasible, and three or more different streams can be accommodated in the same equipment. Pressure drop, heat transfer relations, and other aspects of design are well documented, particularly by Kays and London (1984) and in HEDH (1983, Sec. 3.9).

AIR COOLERS

In such equipment the process fluid flows through finned tubes and cooling air is blown across them with fans. Figures 8.4(g) and (h) show the two possible arrangements. The economics of application of air coolers favors services that allow $25-40^{\circ}F$ temperature difference between ambient air and process outlet. In the range above 10 MBtu/(hr), air coolers can be economically competitive with water coolers when water of adequate quality is available in sufficient amount.

Tubes are 0.75-1.00 in. OD, with 7-11 fins/in. and 0.5-0.625 in. high, with a total surface 15-20 times bare surface of the tube. Fans are 4-12 ft/dia, develop pressures of 0.5-1.5 in. water, and require power inputs of 2-5 HP/MBtu/hr or about 7.5 HP/ 100 sqft of exchanger cross section. Spacings of fans along the length of the equipment do not exceed 1.8 times the width of the cooler. Face velocities are about IO ft/sec at a depth of three rows and 8 ft/sec at a depth of six rows.

Standard air coolers come in widths of 8, 10, 12, 16, or 20 ft, lengths of 4–40 ft, and stacks of 3-6 rows of tubes. Example 8.8 employs typical spacings.

Three modes of control of air flow are shown in Figure 3.3(e). Precautions may need to be taken against subcooling to the freezing point in winter.



Figure 8.8. Plate and spiral compact exchangers. (a) Plate heat exchanger with corrugated plates, gaskets, frame, and corner portals to control flow paths. (b) Flow patterns in plate exchangers, (i) parallel-counter flows; (ii) countercurrent flows; (iii) parallel flows throughout. (c) Spiral exchanger, vertical, and horizontal cross sections.

(c)

Forced draft arrangement, from below the tubes, Figure 8.4(h), develops high turbulence and consequently high heat transfer coefficients. Escape velocities, however, are low, 3 m/sec or so, and as a result poor distribution, backmixing and sensitivity to cross currents can occur. With induced draft from above the tubes, Figure 8.4(g), escape velocities may be of the order of 10 m/sec and better flow distribution results. This kind of installation is more expensive, the pressure drops are higher, and the equipment is bathed in hot air which can be deteriorating. The less solid mounting also can result in noisier operation.

Correlations for friction factors and heat transfer coefficients are cited in HEDH. Some overall coefficients based on external bare tube surfaces are in Tables 8.11 and 8.12. For single passes in cross flow, temperature correction factors are represented by Figure **8.5(c)** for example; charts for multipass flow on the tube side are given in HEDH and by Kays and London (1984), for example. Preliminary estimates of air cooler surface requirements can be made with the aid of Figures 8.9 and 8.10, which are applied in Example 8.9.

DOUBLE-PIPES

This kind of exchanger consists of a central pipe supported within a larger one by packing glands [Fig. 8.4(a)]. The straight length is limited to a maximum of about 20 ft; otherwise the center pipe will sag and cause poor distribution in the **annulus**. It is customary to operate with the high pressure, high temperature, high density, and corrosive fluid in the inner pipe and the less demanding one in the **annulus**. The inner surface can be provided with scrapers [Fig. 8.4(b)] as in **dewaxing** of oils or crystallization from solutions. External longitudinal fins in the annular space can be used to improve heat transfer with gases or viscous fluids. When greater heat transfer surfaces are needed, several double-pipes can be stacked in any combination of series or parallel.

Double-pipe exchangers have largely lost out to shell-and-tube units in recent years, although Walker (1982) lists 70 manufacturers of them. They may be worth considering in these situations:

- 1. When the shell-side coefficient is less than half that of the tube side; the annular side coefficient can be made comparable to the tube side.
- 2. Temperature crosses that require multishell shell-and-tube units can be avoided by the inherent true countercurrent flow in double pipes.
- 3. High pressures can be accommodated more economically in the **annulus** than they can in a larger diameter shell.
- At duties requiring only 100-200 sqft of surface the double-pipe may be more economical, even in comparison with off-the-shelf units.

The process design of double-pipe exchangers is practically the simplest heat exchanger problem. Pressure drop calculation is straightforward. Heat transfer coefficients in annular spaces have been investigated and equations are cited in Table 8.10. A chapter is devoted to this equipment by Kern (1950).

8.7. SHELL-AND-TUBE HEAT EXCHANGERS

Such exchangers are made up of a number of tubes in parallel and series through which one fluid travels and enclosed in a shell through which the other fluid is conducted.

CONSTRUCTION

The shell side is provided with a number of baffles to promote high velocities and largely more efficient cross flow on the outsides of the

TABLE 8.11. Overall Heat Transfer Coefficients in Air Coolers [U Btu/(hr)(°F)(sqft of outside bare tube surface)]

Liquid Coolers					Condensers		
	Mate	rial	Heat-Transfer Coefficient, [Btu/(hr) (ft²)(°F)]	Material	Heat-Transfer Coefficient [Btu/(hr)(ft²)(°F)]	Material	Heat-Transfer Coefficient [Btu/(hr)(ft²)(°F)]
Oils, 20"	' API		IO-16	Heavy oils, 8–14° API		Steam	140-150
200°F	avg.	temp	10–16	300°F avg. temp	6-10	Steam	
300°F	avg.	temp	13-22	400°F avg. temp	10-16	10% noncondensibles	100-110
400°F	avg.	temp	30-40	Diesel oil	45-55	20% noncondensibles	95-100
				Kerosene	55-60	40% noncondensibles	70-75
Oils, 30"	' API			Heavy naphtha	60-65	Pure light hydrocarbons	80-85
150°F	avg.	temp	12-23	Light naphtha	65-70	Mixed light hydrocarbons	65-75
200°F	avg.	temp	25-35	Gasoline	70-75	Gasoline	60-75
300°F	avg.	temp	45-55	Light hydrocarbons	75-80	Gasoline-steam mixtures	70-75
400°F	avg.	temp	50-60	Alcohols and most		Medium hydrocarbons	45-50
				organic solvents	70-75	Medium hydrocarbons	
0ils, 40	° API					w/steam	55-60
150°F	avg.	temp	25-35	Ammonia	100-120	Pure organic solvents	75-80
200°F	avg.	temp	50-60	Brine, 75% water	90-110	Ammonia	100-110
300°F	avg.	temp	55-65	Water	120-140		
400°F	avg.	temp	60-70	50% ethylene glycol			
	-			and water	100-120		

Vapor Coolers

		Heat-Transfe	r Coefficient (Btu ,	/(hr)(ft ²)(°F)]	
Material	10 psig	50 psig	100 psig	300 psig	500 psig
Light hydrocarbons	15-20	30-35	45-50	65-70	70-75
Medium hydrocarbons and organic solvents	15-20	35-40	45-50	65-70	70-75
Light inorganic vapors	IO-15	15-20	30-35	45-50	50-55
Air	8-10	15-20	25-30	40-45	45-50
Ammonia	IO-15	15-20	30-35	45-50	50-55
Steam	10-15	15-20	25-30	45-50	55-60
Hydrogen 400%	00.00	45 50	05 70	05.05	05 400
75% vol	20-30 17-28	45-50 40-45	60-65	85-95 80-85	95-100 85-90
50% vol	15-25	35-40	55-60	75-80	85-90
25% vol	12-23	30-35	45-50	65-70	80-85

[Brown, Chem. Eng. (27 Mar. 1978)].

TABLE 8.12. Overall Heat Transfer Coefficients in Condensers, $Btu/(hr)(sqft)(°F)^{a}$

	Liquid	Coolants	
Vapor		Coolant	Btu/(hr)(sqft)(°F)
Alcohol		water	100-200
Dowtherm		tall oil	60-80
Dowtherm Hydrocarbons		Dowtherm	80-120
high boiling under vacu	um	water	18-50
low boiling		water	80-200
intermediate		oil	25-40
kerosene		water	30-65
kerosene		oil	20-30
naphtha		water	50-75
naphtha		oil	20-40
Organic solvents		water	100-200
Steam		water	400-1000
Steam-organic azeotrope		water	40-80
Vegetable oils		water	20-50

A	ir Coolers
Vapor	Btu/(hr) (bare sqft)(℉)
Ammonia	100-120
Freons	60-80
Hydrocarbons, light	80-100
Naphtha, heavy	60-70
Naphtha, light	70-80
Steam	130-140

"Air cooler data are based on 50mm tubes with aluminum fins 16-18 mm high spaced 2.5-3 mm apart; coefficients based on bare tube surface. Excerpted from HEDH, 1983.





Figure 8.9. Required surfaces of air coolers with three rows of tubes. (a) CJ = 140. (b) U = 120. (c) U = 100. (d) U = 80. (e) U = 60. [Lerner, Hyd. Proc., 93–100 (Fed. 1972)].

150 200 300 400 500


Figure 8.10. Required surfaces of air coolers with six rows of tubes. (a) $U = 100 \text{ Btu}/(\text{hr})(\text{sqft})(^{\circ}\text{F})$. (b) U = 80. (c) U = 60. (d) U = 40. (e) U = 20. (f) U = 10. [Lemmer, Hyd. Proc., 93-100 (Feb. 1972)].

EXAMPLE 8.9

Estimation of the Surface Requirements of an Air Cooler An oil is to be cooled from 300 to 150° F with ambient air at 90° F, with a total duty of 20 MBtu/hr. The tubes have 5/8 in. fins on 1 in. OD and 2-5/16 in. triangular spacing. The tube surface is given by

A = 1.33NWL, sqft of bare tube surface, N = number of rows of tubes, from 3 to 6, W = width of tube bank, ft, L = length of tubes, ft.

According to the data of Table 8.12, the overall coefficient may be taken as U = 60 Btu/(hr)(°F)(sqft of bare tube surface). Exchangers with 3 rows and with 6 rows will be examined.

Approach =
$$150 - 90 = 60^{\circ}F$$
,
Cooling range = $300 - 150 = 150^{\circ}F$,
From Figure 8.9(f), 3 rows,

A = 160 sqft/MBtu/hr) $\rightarrow 160(20) = 3200 \text{ sqft}$

= 1.33(3)WL.

tubes. Figure 8.4(c) shows a typical construction and flow paths. The versatility and widespread use of this equipment has given rise to the development of industrywide standards of which the most widely observed are the TEMA standards. Classifications of equipment and terminology of these standards are summarized on Figure 8.11.

Baffle pitch, or distance between baffles, normally is 0.2-1.0 times the inside diameter of the shell. Both the heat transfer coefficient and the pressure drop depend on the baffle pitch, so that its selection is part of the optimization of the heat exchanger. The window of segmental baffles commonly is about 25%, but it also is a parameter in the thermal-hydraulic design of the equipment.

In order to simplify external piping, exchangers mostly are built with even numbers of tube passes. Figure 8.12(c) shows some possible arrangements, where the full lines represent partitions in one head of the exchanger and the dashed lines partitions in the opposite head. Partitioning reduces the number of tubes that can be accommodated in a shell of a given size. Table 8.12 is of such data. Square tube pitch in comparison with triangular pitch accommodates fewer tubes but is preferable when the shell side must be cleaned by brushing.

Two shell passes are obtained with a longitudinal baffle, type F in Figures 8.11(a) or 8.3(c). More than two shell passes normally are not provided in a single shell, but a 4-8 arrangement is thermally equivalent to two 2-4 shells in series, and higher combinations are obtained with more shells in series.

ADVANTAGES

A wide range of design alternates and operating conditions is obtainable with shell-and-tube exchangers, in particular:

- Single phases, condensation or boiling can be accommodated in either the tubes or the shell, in vertical or horizontal positions.
- Pressure range and pressure drop are virtually unlimited, and can be adjusted independently for the two fluids.

When W = 16 ft, L = 50 ft.

Two fans will make the ratio of section length to width, 25/16 = 1.56 which is less than the max allowable of 1.8. At 7.5 HP/100 sqft,

Power =
$$\frac{16(50)}{100}$$
 7.5 = 60 HP.

From Figure 8.10(c), 6 rows,

A = 185 sqft/(MBtu/hr)→ 185(20) = 3700 sqft. = 1.33(6)WL.

When W = 16 ft, L = 29 ft.

Since L/W = 1.81, one fan is marginal and two should be used:

Power = [16(29)/100]7.5 = 34.8 HP.

The 6-row construction has more tube surface but takes less power and less space.

- Thermal stresses can be accommodated inexpensively.
- A great variety of materials of construction can be used and may be different for the shell and tubes.
- Extended surfaces for improved heat transfer can be used on either side.
- A great range of thermal capacities is obtainable.
- The equipment is readily dismantled for cleaning or repair.

TUBE SIDE OR SHELL SIDE

Several considerations may influence which fluid goes on the tube side or the shell side.

The tube side is preferable for the fluid that has the higher pressure, or the higher temperature or is more corrosive. The tube side is less likely to leak expensive or hazardous fluids and is more easily cleaned. Both pressure drop and laminar heat transfer can be predicted more accurately for the tube side. Accordingly, when these factors are critical, the tube side should be selected for that fluid.

Turbulent flow is obtained at lower Reynolds numbers on the shell side, so that the fluid with the lower mass **flow** preferably goes on that side. High Reynolds numbers are obtained by multipassing the **tube** side, but at a price.

DESIGN OF A HEAT EXCHANGER

A substantial number of parameters is involved in the design of a shell-and-tube heat exchanger for specified thermal and hydraulic conditions and desired economics, including: tube diameter, thickness, length, number of passes, pitch, square or triangular; size of shell, number of shell baffles, baffle type, baffle windows, baffle spacing, and so on. For even a modest sized design program, Bell (in HEDH, 1983, 3.1.3) estimates that 40 separate logical designs may need to be made which lead to $2^{40} = 1.10 \times 10^{12}$ different paths through the logic. Since such a number is entirely too large for normal computer processing, the problem must be simplified with





1.	SHELL	8.	FLOATING HEAD FLANGE	16.	TRANSVERSE BAFFLES OR
2.	SHELL COVER	9.	CHANNEL PARTITION		SUPPORT PLATES
3.	SHELL CHANNEL	10.	STATIONARY TUBESHEET	16.	IMPINGEMENT BAFFLE
€.	SHELL COVER END FLANGE	11.	CHANNEL	17.	VENT CONNECTION
6.	SHELL NOZZLE	12.	CHANNEL COVER	18.	DRAIN CONNECTION
6,	FLOATING TUBESHEET	13.	CHANNEL NOZZLE	19.	TEST CONNECTION
7.	FLOATING HEAD	14.	TIE RODS AND SPACERS	20.	SUPPORT SADDLES
				21.	LIFTING RING



Figure 8.11, Tubular Exchanger Manufacturers Association classification and terminology for heat exchangers. (a) TEMA terminology for shells and heads of heat exchangers. (b) Terminology for parts of a TEMA type AES heat exchanger. The three letters A, E, and S come from part (a).

some arbitrary decisions based on as much current practice as possible.

A logic diagram of a heat exchanger design procedure appears in Figure 8.13. The key elements are:

- 1. Selection of a tentative set of design parameters, Box 3 of Figure 8.13(a).
- 2. Rating of the tentative design, Figure 8.13(b), which means evaluating the performance with the best correlations and calculation methods that are feasible.
- 3. Modification of some design parameters, Figure 8.13(c), then rerating the design to meet thermal and hydraulic specifications and economic requirements.

A procedure for a tentative selection of exchanger will be described following. With the exercise of some judgement, it is feasible to perform simpler exchanger ratings by hand, but the present state of the art utilizes computer rating, with in-house programs, or those of HTRI or HTFS, or those of commercial services. More than 50 detailed numerical by hand rating examples are in the book of Kern (1950) and several comprehensive ones in the book of Ganapathy (1982).

TENTATIVE DESIGN

The **stepwise** procedure includes statements of some rules based on common practice.

- 1. Specify the flow rates, terminal temperatures and physical properties.
- 2. Calculate the LMTD and the temperature correction factor *F* from Table 8.3 or Figure 8.5.
- 3. Choose the simplest combination of shell and tube passes or number of shells in series that will have a value of F above 0.8 or so. The basic shell is 1-2, one shell pass and two tube passes.
- 4. Make an estimate of the overall heat transfer coefficient from Tables 8.4-8.7.
- 5. Choose a tube length, normally 8, 12, 16, or 20 ft. The 8 ft long exchanger costs about 1.4 times as much as the 20 ft one per unit of surface.
- 6. Standard exchanger tube diameters are 0.75 or 1 in. OD, with pitches shown in Table 8.13.
- 7. Find a shell diameter from Table 8.13 corresponding to the selections of tube diameter, length, pitch, and number of passes made thus far for the required surface. As a guide, many heat exchangers have length to shell diameter ratios between 6 and 8.
- 8. Select the kinds and number of baffles on the shell side.

The tentative exchanger design now is ready for detailed evaluation with the best feasible heat transfer and pressure drop data. The results of such a rating will suggest what changes may be needed to satisfy the thermal, hydraulic, and economic requirements for the equipment. Example 8.10 goes through the main part of such a design.

8.8. CONDENSERS

Condensation may be performed inside or outside tubes, in horizontal or vertical positions. In addition to the statements made in the previous section about the merits of tube side or shell side: When freezing can occur, shell side is preferable because it is less likely to clog. When condensing mixtures whose lighter components are soluble in the condensate, tube side should be adopted since drainage is less complete and allows condensation (and dissolution) to occur at higher temperatures. Venting of noncondensables is more positive from tube side.



Figure **8.12.** Arrangements of cross baffles and tube-side passes. (a) Types of cross baffles. (b) Rod baffles for minimizing tube vibrations; each tube is supported by four rods. (c) Tube-side multipass arrangements.



Figure 8.13. A procedure for the design of a heat exchanger, comprising a tentative selection of design parameters, rating of the performance, modification of this design if necessary, and re-rating to meet specifications (see *also Bell, in* Heat Exchanger Design Handbook, *Section 3.1.3, Hemisphere Publishing Company, 1983*).

Τ

I-Tube Heat Exchangers"										
Heat	Excha	anger '	Tube S	Shect I	Layout	Count	Table	c		
25	231/4	211/4	191/4	171/4	151/4	131/4	12	10	8	I.D. of Shell (In.)
550	191	201	207	917	102	125	105	69	33	3/ on 15/ A

37	35	33	31	29	27	25	231/4	211/4	191/4	171/4	151/4	131/4	12	10	8	I.D. of Shell	(In.)	
1269 1127 965 699 595	$1143 \\1007 \\865 \\633 \\545$	1019 889 765 551 477	$\begin{array}{r} 881 \\ 765 \\ 665 \\ 481 \\ 413 \end{array}$	783 667 587 427 359	663 577 495 361 303	$553 \\ 493 \\ 419 \\ 307 \\ 255$	481 423 355 247 215	391 343 287 205 179	307 277 235 163 139	247 217 183 133 111	193 157 139 103 83	135 117 101 73 65	105 91 85 57 45	69 57 53 33 33	33 33 33 15 17	34 on 154 A 34 on 1 A 34 on 1 D 1 on 14 A 1 on 14 A 1 on 14 A	Fixe Tubd	*
1242 1088 946 688 584	1088 972 840 608 523	964 858 746 530 460	846 746 644 462 402	734 646 560 410 348	626 556 486 346 298	528 468 408 292 248	452 398 346 244 218	370 32G 280 204 172	300 264 222 162 136	228 208 172 126 106	166 154 126 92 76	124 110 94 62 56	$94 \\ 90 \\ 78 \\ 52 \\ 40$	5 % 56 48 32 26	$32 \\ 2 \\ 26 \\ 16 \\ 12$	³ ⁄ ₄ " on ¹⁵ ∕ ₁₆ "∆ ³ ⁄ ₄ " on 1 A ³ ⁄ ₄ " on 1 □ 1' on 11⁄ ₄ " A 1" on 1%' □	Fixed Tubes	-
$1126 \\ 1000 \\ 884 \\ 610 \\ 526$	1008 882 778 532 464	882 112 688 466 406	$768 \\ 674 \\ 586 \\ 396 \\ 356 \\ 356 \\ \end{array}$	648 5GG 506 340 304	558 484 436 284 256	$\begin{array}{r} 460\\ 406\\ 362\\ 234\\ 214\end{array}$	398 336 304 192 180	304 270 242 154 134	234 212 188 120 100	180 158 142 84 76	1 34 108 100 58 58	94 72 72 42 38	$64 \\ 60 \\ 52 \\ 26 \\ 22$	34 26 30 8 12	8 8 1 X X 1 X X 1	³ ⁄4" on 1 ⁵ ⁄16"∆ ³ ⁄4" on 1"∆ ³ ⁄4" on 1"□ ¹ on 1%' A 1" on 1%' □	o Tubes ² "	
$1172 \\ 1024 \\ 880 \\ 638 \\ 534$	1024 913 118 560 476	$904 \\ 802 \\ 688 \\ 486 \\ 414$	788 692 590 422 3G0	680 596 510 368 310	576 508 440 308 2G0	484 424 366 258 214	412 3GO 308 212 188	332 292 242 17G 142	266 2 3 2 192 138 1 1 0	196 180 142 104 84	154 134 12G 78 74	108 96 88 GO 48	84 72 72 44 40	$ \begin{array}{r} 4 & 8 \\ 4 & 4 \\ 4 & 8 \\ 2 & 4 \\ 2 & 4 \end{array} $	$ \begin{array}{c} $	$\begin{array}{c} {}^{34}_{4} \text{ on } {}^{15}_{16} \text{ A} \\ {}^{34}_{4} \text{ on } {}^{1}_{4} \Delta \\ {}^{34}_{4} \text{ on } {}^{1}_{4} \Box \\ {}^{34}_{4} \text{ on } {}^{1}_{4} \Box \\ {}^{34}_{1} \text{ on } {}^{14}_{4} \Box \end{array}$	Fixes	á ľ
1092 968 \$52 584 500	976 852 748 508 440	S52 744 660 444 384	$740 \\ 648 \\ 560 \\ 376 \\ 336 \\$	$\begin{array}{c} 622 \\ 542 \\ 482 \\ 522 \\ 286 \end{array}$	$534 \\ 462 \\ 414 \\ 266 \\ 238$	$438 \\ 386 \\ 342 \\ 218 \\ 198$	378 318 286 178 166	$286 \\ 254 \\ 226 \\ 142 \\ 122$	218 198 174 110 90	166 146 130 74 66	122 98 90 50 50	84 64 G4 36 32	5G 52 44 20 16	28 2 0 24 XX XX	XX XX XX XX XX	34° on 154° Δ 34° on 1° A 34° on 1° \Box 1° on 114° Δ 1° on 114° \Box	U _{Tubcs} ²)@
1106 964 818 586 484	$964 \\ 852 \\ 224 \\ 514 \\ 430 \\ -$	844 744 634 442 3GS	732 640 536 382 318	(3)L 54S 460 338 268	532 464 394 274 226	$\begin{array}{r} 440\\ 388\\ 324\\ 226\\ 184 \end{array}$	372 323 2GG 182 154	294 25s 212 150 116	2 3 0 202 158 112 88	174 15G 116 82 66	116 104 78 56 44 -	8 66 5 34 X	0 XX 4 XX X	XX X XX XX XX XX XX X	(X X) X X X X) X X X X X X	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Fixed Tubes	• • • • •
1058 940 820 562 478	944 826 718 488 420	826 720 632 426 362	716 626 634 356 316	596 518 458 304 268	510 440 392 252 224	416 366 322 206 182	35s 300 268 168 152	$272 \\ 238 \\ 210 \\ 130 \\ 110$	206 184 160 100 80	156 134 118 68 60	110 88 80 42 42	74 5 (5 6 30 XX	XX X X X XX XX	XX*'' X X XX XX XX ⊠	XX '3 S X XX XX ⊠ 1'	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	U Tubes²	
1040 902 760 542 438	10 598 GG2 466 388	790 694 576 400 334	$\begin{array}{r} 682 \\ 588 \\ 490 \\ 342 \\ 2S0 \end{array}$	576 496 414 298 230	484 422 352 240 192	398 344 286 190 150	332 28G 228 154 128	258 224 174 120 94	198 170 132 90 74	140 124 94 66 XX	94 82 XX XX XX XX	X X X X XX XX XX	XX XX XX X X XX	X X X X XX X X X X	$\begin{array}{c} X & X \\ \chi \\ X \\ X \\ X \\ 1 \end{array}$	34° on 15_{16}° A XX 34° on 1° Δ 34° on 1° Δ XX XX 1° on $11_{4}^{\circ}^{\circ}$ on $11_{4}^{\circ}^{\circ}$	Fixe ^{cs} Tub	ш р
1032 908 792 540 456	916 796 692 464 396	796 692 608 404 344	$688 \\ 600 \\ 512 \\ 340 \\ 300$	578 498 438 290 254	490 422 374 238 206	398 350 306 190 170	342 286 254 154 142	254 226 194 118 98	190 170 146 90 70	$142 \\ 122 \\ 106 \\ 58 \\ 50$	102 82 70 38 34	68 52 48 2 X X	XX X X X X 4 X X	XX X X X X X X	XX XX XX XX XX XX	34 ° on 15/16 A 34 ° on 1 * ∆ 34 ° on 1 * □ 34 ° on 1 * □ XX XX 1 ° on 1 ` on 1 4 ° □	⊂ U Tu`≁s²	
37 I	35 I	33	31	29	27	25	231/4	211/4	191/4	171/4	151/4	131/4 1	2 1		8	I.D. of Shell (in.)		
1 11	lowance	made fr	nr Tio	Rode														

- Stratit

2 R.O.B. = $2\frac{1}{2} \times$ Tube Dia. Actual Number of "U" Tubes is one-half the above figures. *A 3/4 in. tube has 0.1963 sqft/ft, a 1 in. OD has 0.2618 sqft/ft. Allowance made for tie rods. * R.O.B. = $2\frac{1}{2} \times$ tube dia. Actual number of "U" tubes is one-half the above figures.

EXAMPLE 8.10

Process Design of a Shell-and-Tube Heat Exchanger An oil at the rate of 490,000 lb/hr is to be heated from 100 to 170°F

with 145,000 **Ib/hr** of kerosene initially at 390°F. Physical properties are





Kerosene outlet:

T = 390 - (490,000/145,000)(0.49/0.61)(170 - 100)= 200°F, LMTD = (220 - 100)/ln 2.2 = 152.2, P = (170 - 100)/(390 - 100) = 0.241, R = (390 - 200)/(170 - 100) = 2.71.

From Figure 8.5(a), F = 0.88, so a 1-2 exchanger is satisfactory:

AT = 152.2(0.88) = 133.9.

From Table 8.6, with average values for medium and heavy organics,

 $U = 10^4/(57 + 16 + 50 + 34) = 63.7$, $Q = 490,000(0.49)(170 - 100) = 1.681(10^7)$ Btu 1 hr, $A = Q/U\Delta T = 1.681(10^7)/63.7(133.9) = 1970$ sqft, 1970/0.2618 = 7524.8 ft of 1 in. OD tubing.

Use $1\frac{1}{4}$ in. pitch, two tube pass. From Table 8.13,

	Required	D_{shell} (number	of tubes)	
L (ft)	No. Tubes	Triangular	Square	
8 12 16 20	940 627 470 376	35 (608) 31 (462) 29 (410)	37 (584) 33 (460) 31 (402)	

CONDENSER CONFIGURATIONS

The several possible condenser configurations will be described. They are shown on Figure 8.14.

Condensation Inside Tubes: Vertical Downflow. Tube diameters normally are 19-25 mm, and up to 50 mm to minimize critical pressure drops. The tubes remain wetted with condensate which assists in retaining light soluble components of the vapor. Venting of noncondensables is positive. At low operating pressures, larger tubes may be required to minimize pressure drop; this may have the effect of substantially increasing the required heat transfer surface. A disadvantage exists with this configuration when the coolant is fouling since the shell side is more difficult to clean. Use 16 ft tubes on $1\frac{1}{4}$ in. square pitch, two pass, 33 in. shell

$$L/D = 16/(33/12) = 5.82,$$

which is near standard practice. The 20 ft length also is acceptable but will not be taken.

The pressure drops on the tube and shell sides are to be calculated.

Tube side: 0.875 *in. ID*, 230 tubes, *32ft* long: Take one velocity head per inlet or outlet, for a total of 4, in addition to friction in the tubes. The oil is the larger flow so it will be placed in the tubes.

 $\dot{m} = 490,000/230 = 2130.4 \text{ lb/(hr)(tube)}.$

Use formulas from Table 6.1

Re = 6.314(2130.4)/0.875(3.5) = 4392, **f** = $1.6364/[\ln(5(10^{-7})/0.875 + 6.5/4392)]^2 = 0.0385$, $\Delta P_f = 5.385(10^{-8})(2130)^2(32)(0.0385)/0.85(0.875)^5$ = 0.691 psi.

Expansion and contraction:

 $\Delta P_e = 4\rho(u^2/2q_e) = 4(53.04)(3.26)^2/(64.4)(144) = 0.243 \text{ psi},$ $\therefore \Delta P_{\text{tube}} = 0.691 + 0.243 = 0.934 \text{ psi}.$

Shellside. Follow Example 8.8:

 $\begin{aligned} D_h &= 1.2732(1.25/12)^2/(1/12) - 1/12 = 0.0824 \text{ ft}, \\ B &= 1.25 \text{ ft between baffles}, \\ E &= 0.25/12 \text{ ft between tubes}, \\ D_s &= 33/12 = 2.75 \text{ ft shell diameter}, \\ A_{,} &= 2.75(1.25)(0.25/12)/(1.25/12) = 0.6875 \text{ sqft}, \\ G_s &= 145,000/0.6875 = 210,909 \text{ lb/(hr)(sqft)}, \\ \text{Re} &= 0.0824(210,909)/0.4(2.42) = 17,952, \\ \mathbf{f} &= 0.0121(17,952)^{-0.19} = 0.00188, \\ \Delta P_{\text{shell}} - 0.00188(210,909)^2(2.75)(13)/5.22(10^{10})(0.82)(0.0824) \\ &= 0.85 \text{ psi.} \end{aligned}$

The pressure drops on each side are acceptable. Now it remains to check the heat transfer with the equations of Table 8.10 and the fouling factors of Table 8.6.

Condensation Inside Tubes: Vertical Upflow. This mode is used primarily for refluxing purposes when return of a hot condensate is required. Such units usually function as partial condensers, with the lighter components passing on through. Reflux condensers usually are no more than 6-10 ft long with tube diameters of 25 mm or more. A possible disadvantage is the likelihood of flooding with condensate at the lower ends of the tubes.

Condensation Outside Vertical Tubes. This arrangement requires careful distribution of coolant to each tube, and requires a sump and a pump for return to a cooling tower or other source of coolant. Advantages are the high coolant side heat transfer



Figure 8.14. Some arrangements of shell-and-tube condensers. (a) Condensate inside tubes, vertical upflow. (b) Inside tubes, vertical downflow. (c) Outside tubes, vertical downflow. (d) Condensate outside horizontal tubes. (HEDH, 1983, 3.4.3).

coefficient and the ease of cleaning. The free draining of condensate is a disadvantage with wide range mixtures.

Condensation Inside Horizontal Tubes. This mode is employed chiefly in air coolers where it is the only feasible mode. As condensation proceeds, liquid tends to build up in the tubes, then slugging and oscillating flow can occur.

Condensation Outside Horizontal Tubes. Figure 8.14(d) shows a condenser with two tube passes and a shell side provided with vertically cut baffles that promote side to side flow of vapor. The tubes may be controlled partially flooded to ensure desired subcooling of the condensate or for control of upstream pressure by regulating the rate of condensation. Low-fin tubes often are advantageous, except when the surface tension of the condensates

exceeds about 40 dyn/cm in which event the fins fill up with stagnant liquid. The free draining characteristic of the outsides of the tubes is a disadvantage with wide condensing range mixtures, as mentioned. Other disadvantages are those generally associated with shell side fluids, namely at high pressures or high temperatures or corrosiveness. To counteract such factors, there is ease of cleaning if the coolant is corrosive or fouling. Many cooling waters are scale forming; thus they are preferably placed on the tube side. On balance, the advantages often outweigh the disadvantages and this type of condenser is the most widely used.

DESIGN CALCULATION METHOD

Data for condensation are described in Section 8.4 and given in Tables 8.4-8.7, and a few additional overall coefficients are in Table



Figure 8.15. Model for partial condensation in the presence of uncondensed material: $U(T_i - T_L) = h_g(T_g - T_i) + \lambda k_g(p_g - p_i)$. [A.P. Colburn and O.A. Hougen, Ind. Eng. Chem. 26, 1178-1182 (1934)].

8.12. The calculation of condensation of pure vapors is straightforward. That of mixtures occurs over a range of temperatures and involves mass transfer resistance through a gas film as well as heat transfer resistance by liquid and fouling films. A model due to Colburn and Hougen (1934) is represented by Figure 8.15. The overall rate of heat transfer is regarded as the sum of the sensible heat transfer through a gas film and the heat of condensation of the material transferred by diffusion from the gas phase to the interface. The equation of this heat balance is, in terms of the notation of Figure 8.15,

$$U(T_{i} - T_{L}) = h_{g}(T_{g} - T_{i}) + \lambda k_{g}(p_{g} - p_{i}).$$
(8.37)

The temperature T_L of the coolant is related to the heat transfer Q by

$$dQ = \dot{m}_L C_L \, dT_L$$

or the integrated form

$$T_L = T_{L0} + \Delta Q / \dot{m}_L C_L. \tag{8.38}$$

A procedure will be described for taking the vapor from its initial **dewpoint** T_{g0} to its final **dewpoint** corresponding to the required amount of condensation. Gas temperatures are specified at intermediate points and the heat balance is applied over one interval at a time.

- 1. Prepare the condensing curve, a plot of the vapor temperature T_g against the amount of heat removed Q, by a series of isothermal flashes and enthalpy balances.
- 2. Starting at the inlet temperature T_{g0} , specify a temperature T_g a few degrees less, and note the heat transfer AQ corresponding to this temperature difference from the condensing curve.
- 3. Find the temperature T_L of the coolant with Eq. (8.38).
- 4. Assume an interfacial temperature T_i , then find the corresponding vapor pressure p_i and latent heat λ .
- 5. From available correlations, find values of the coefficients h_g , k_g , and U which are temperature- and composition-dependent, although they sometimes may be taken as constant over some ranges.
- 6. Check if these values satisfy the heat balance of Eq. (8.37). If not, repeat the process with other estimates of T_i until one is found that does satisfy the heat balance.

- 7. Continue with other specifications of the vapor temperature T_g , one interval at a time, until the required outlet temperature is reached.
- 8. The heat transfer area will be found by numerical integration of

$$A = \int_0^Q \frac{dQ}{U(T_i - T_L)} \,. \tag{8.39}$$

Examples of numerical applications of this method are in the original paper of Colburn and Hougen (1934), in the book of Kern (1950, p. 346) and in the book of Ludwig (1983, Vol. 3, p. 116).

The Silver-Bell-Ghaly Method

This method takes advantage of the rough proportionality between heat and mass transfer coefficients according to the Chilton-Colburn analogy, and employs only heat transfer coefficients for the process of condensation from a mixture. The sensible heat Q_{sv} of the vapor is transferred through the gas film

$$dQ_{sv} = h_g(T_g - T_i) \, dA. \tag{8.40}$$

In terms of an overall heat transfer coefficient U that does not include the gas film, the total heat transfer Q_T that is made up of the latent heat and the sensible heats of both vapor and liquid is represented by

$$dQ_T = U(T_i - T_L) \, dA. \tag{8.41}$$

When the unknown interfacial temperature T_i is eliminated and the ratio Z of sensible and total heat transfers

$$Z = dQ_{sv} / dQ_T \tag{8.42}$$

is introduced, the result is

$$dQ_T = \frac{U(T_g - T_L)}{1 + ZU/h_g} dA,$$
 (8.43)

which is solved for the heat transfer area as

$$A = \int_0^{Q_T} \frac{1 + ZU/h_g}{U(T_g - T_L)} dQ_T.$$
 (8.44)

Since the heat ratio Z, the temperatures and the heat transfer coefficients vary with the amount of heat transfer Q_T up to a position in the condenser, integration must be done numerically. The coolant temperature is evaluated from Eq. (8.38). Bell and Ghaly (1973) examine cases with multiple tube passes.

The basis of the method was stated by Silver (1947). A numerical solution of a condenser for mixed hydrocarbons was carried out by Webb and McNaught (in Chisholm, 1980, p. 98); comparison of the Silver-Bell-Ghaly result with a Colburn-Hougen calculation showed close agreement in this case. Bell and Ghaly (1973) claim only that their method predicts values from 0 to 100% over the correct values, always conservative. A solution with constant heat transfer coefficients is made in Example 8.11: A recent review of the subject has been presented by McNaught (in Taborek et al., 1983, p. 35).

8.9. REBOILERS

Reboilers are heat exchangers that are used primarily to provide **boilup** for distillation and similar towers. All types perform partial vaporization of a stream flowing under natural or forced circulation

EXAMPLE 8.11 Sizing a Condenser for a Mixture by the Silver-Bell-Ghatly Method

A mixture with initial **dewpoint 139.9°C** and final bubblepoint **48.4°C** is to be condensed with coolant at a constant temperature of 27°C. The gas film heat transfer coefficient is $40 \text{ W/m}^2 \text{ K}$ and the overall coefficient is 450. Results of the calculation of the condensing curve are

```
T(°C) 139.9 121.6 103.3 85.0 66.7 48.4 
Q(W) 0 2154 3403 4325 5153 5995
```

In the following tabulation, over each temperature interval are

shown the average gas temperature, the value of Z, and the value of the integrand of Eq. (8.44). The integrand is plotted following.

Interv	al	1	2	3	4	5
$(T_a)_m$		130. 75	112.45	94.15	75.85	57.4
Z	_	0.1708	0. 1613	0. 1303	0.0814	0. 0261
lntegrandx	(10 ⁵)	6.26	7.32	8.31	8.71	9.41

The heat transfer surface is the area under the stepped curve, which is $a = 0.454 \text{ m}^2$. A solution that takes into account the substantial variation of the heat transfer coefficients along the condenser gives the result $\mathbf{A} = 0.385 \text{ m}^2$ (Webb and McNaught, in Chisholm, 1980, p. 98).



conditions. Sketches of a kettle and two types of thermosiphon reboilers are in Figure 8.4. Internal reboilers, with a tube bundle built into the tower bottom, also have some application. Flow through a vertical unit like that of Figure 8.4(f) may be forced with a pump in order to improve heat transfer of viscous or fouling materials, or when the vaporization is too low to provide enough static head difference, or when the tower skirt height is too low. A summary guide to the several types of reboilers is in Table 8.14.

KETTLE REBOILERS

Kettle reboilers consist of a bundle of tubes in an oversize shell. Submergence of the tubes is assured by an overflow weir, typically **5–15 cm** higher than the topmost tubes. An open tube bundle is preferred, with pitch to diameter ratios in the range of 1.5-2. Temperature in the kettle is substantially uniform. Residence time is high so that kettles are not favored for thermally sensitive materials. The large shell diameters make kettles uneconomic for high pressure operation. Deentraining mesh pads often are incorporated. Tube bundles installed directly in the tower bottom are inexpensive but the amount of surface that can be installed is limited.

HORIZONTAL SHELL SIDE THERMOSIPHONS

The fraction vaporized in thermosiphon reboilers usually can be made less than in kettles, and the holdup is much less. Less static head difference is needed as driving force for recirculation in comparison with vertical units. Circulation rate can be controlled by throttling the inlet line. Because of the forced flow, there is a temperature gradient, from the inlet bubblepoint to the exit bubblepoint, whereas in a kettle the boiling temperature is more nearly uniform, at the exit bubblepoint. Consequently, for the same percentage vaporization, the mean temperature difference between shell and tube sides will be greater for thermosiphons than for kettles. Or for the same mean temperature difference, the percentage vaporization can be made less. Large surface requirements favor horizontal over vertical thermosiphons. Horizontal tube bundles are easier to maintain. The usual arguments for tube side versus shell side also are applicable.

VERTICAL THERMOSIPHONS

Circulation is promoted by the difference in static heads of supply liquid and the column of partially vaporized material. The exit

		Reboiler	Туре	
Process Conditions	Kettle or Internal	Horizontal Shell-Side Thermosiphon	Vertical Tube-Side Thermosiphon	Forced Flow
Operating pressure				
Moderate	E	G	В	E
Near critical	B-E	R	Rd	Е
Deep vacuum	В	R	Rd	Е
Design AT				
Moderate	E	G	В	E
Large	В	R	G - R d	E
Small (mixture)	F	F	Rd	Р
Very small (pure component)	В	F	Р	P
Fouling				
Clean	G	G	G	E
Moderate	Rd	G	В	Е
Heavy	Р	Rd	В	G
Very heavy	P	Р	Rd	В
Mixture boiling range				
Pure component	G	G	G	E
Narrow	G	G	В	E
Wide	F	G	В	E
Very wide, with viscous liquid	F-P	G-Rd	р	В

TABLE 8.14. A Guide to the Selection of Reboilers

^a Category abbreviations: B, best; G, good operation; F, fair operation, but better choice is possible; Rd. risky unless carefully designed, but could be best choice in some cases; R, risky because of insufficient data; P, poor operation; E, operable but unnecessarily expensive. (HEDH, 1983, 3.6.1).

weight fraction vaporized should be in the range of 0.1-0.35 for hydrocarbons and 0.02-0.10 for aqueous solutions. Circulation may be controlled with a valve in the supply line. The top **tube** sheet often is placed at the level of the liquid in the tower. The flow area of the outlet piping commonly is made the same as that of all the tubes. Tube diameters of 19-25 mm diameter are used, lengths up to 12ft or so, but some 20ft tubes are used. Greater tube lengths make for less ground space but necessitate taller tower skirts.

Maximum heat fluxes are lower than in kettle reboilers. Because of boiling point elevations imposed by static head, vertical thermosiphons are not suitable for low temperature difference services.

Shell side vertical thermosiphons sometimes are applied when the heating medium cannot be placed on the shell side.

FORCED CIRCULATION REBOILERS

Forced circulation reboilers may be either horizontal or vertical. Since the feed liquid is at its bubblepoint, adequate NPSH must be assured for the pump if it is a centrifugal type. Linear velocities in the tubes of 15–20 ft/sec usually are adequate. The main disadvantages are the costs of pump and power, and possibly severe maintenance. This mode of operation is a last resort with viscous or fouling materials, or when the fraction vaporized must be kept low.

CALCULATION PROCEDURES

Equations for boiling heat transfer coefficients and maximum heat fluxes are Eqs. (37) through (48) of Table 8.10. Estimating values are in Tables 8.4-8.7. Roughly, boiling coefficients for organics are 300 $Btu/(hr)(sqft)(^{\circ}F)$, or 1700 W/m^2 K; and for aqueous solutions, 1000 $Btu/(hr)(sqft)(^{\circ}F)$, or 5700 W/m^2 K. Similarly, maximum fluxes are of the order of 20,000 Btu/(hr)(sqft), or 63,000 W/m^* , for organics; and 35,000 Btu/(hr)(sqft) or 110,000 W/m^* , for aqueous systems.

The design procedure must start with a specific geometry and heat transfer surface and a specific percentage vaporization. Then the heat transfer coefficient is found, and finally the required area is calculated. When the agreement between the assumed and calculated surfaces is not close enough, the procedure is repeated with another assumed design. The calculations are long and tedious and nowadays are done by computer.

Example 8.12 summarizes the results of such calculations made on the basis of data in *Heat Exchanger Design Handbook* (1983). Procedures for the design of kettle, thermosiphon and forced circulation reboilers also are outlined by Polley (in Chisholm, 1980, Chap. 3).

8.10. EVAPORATORS

Evaporators employ heat to concentrate solutions or to recover dissolved solids by precipitating them from saturated solutions. **They** are reboilers with special provisions for separating liquid and vapor phases and for removal of solids when they are precipitated or crystallized out. Simple kettle-type reboilers [Fig. 8.4(d)] may be adequate in some applications, especially if enough freeboard is provided. Some of the many specialized types of evaporators that are in use are represented on Figure 8.16. The tubes may be horizontal or vertical, long or short; the liquid may be outside or inside the tubes, circulation may be natural or forced with pumps or propellers.

Natural circulation evaporators [Figs. 18.16(a)-(e)] are the most popular. The forced circulation type of Figure 18.16(f) is most versatile, for viscous and fouling services especially, but also the most expensive to buy and maintain. In the long tube vertical design, Figure 8.16(d), because of vaporization the liquid is in annular or film flow for a substantial portion of the tube length, and accordingly is called a rising film evaporator. In falling film

EXAMPLE 8.12

Comparison of Three Kinds of **Reboilers** for the Same Service The service is reboiling a medium boiling range hydrocarbon mixture at 10 atm with a duty of 14,600 kW. The designs are calculated in HEDH (1983, 3.65) and are summarized here.

In each case a specific geometry and surface are assumed; then the heat transfer coefficients are evaluated, and the area is checked. When agreement between assumed and calculated areas is not close, another design is assumed and checked.

Of the three sets of calculations summarized here, only that for the kettle need not be repeated. Both the others should be repeated since the assumed designs are too conservative to be economical.

Quantity	Kettle	Horizontal TS	S Vertical TS
Rated area (m ²)	930	930	480
Tube length (m)	6.1	6.1	4.9
Tube OD (mm)	19	19	<u></u>
Tube (D (mm)	-	_	21.2
Vaporization (%)	30	2 5	2 5
$U(W/m^2K)$	874	674	928
$(\Delta T)_{-}$	2 5	44.8	44.8
Calculated area (m ²)	866	483	350
Calculated \dot{q} (W/m ²)	16,859	30,227	41,174
\dot{q}_{max} (W/m ²)			67,760



evaporators, liquid is distributed to the tops of the individual tubes and flows down as a film. The hydrostatic head is eliminated, the pressure drop is little more than the friction of the vapor flow, and heat transfer is excellent. Since the contact time is short and separation of liquid and vapor is virtually complete, falling film evaporation is suitable for thermally sensitive materials.

Long tube vertical evaporators, with either natural or forced circulation are the most widely used. Tubes range from 19 to 63 mm

diameter, and 12–30 ft in length. The calandria of Figure 8.16(b) has tubes 3-5 ft long, and the central downtake has an area about equal to the cross section of the tubes. Sometimes circulation in calandrias is forced with built in propellors. In some types of evaporators, the solids are recirculated until they reach a desired size. In Figure 8.16(f), fresh feed is mixed with the circulating slurry. In Figure 8.16(g) only the clear liquid is recirculated, and small more nearly uniform crystals are formed.



Figure 8.16. Some types of evaporators. (a) Horizontal tube. (b) Calandria type. (c) Thermocompressor evaporator. (d) Long tube vertical. (e) Falling film. (f) Forced circulation evaporator-crystallizer. (g) Three types of "Oslo/Krystal" circulating liquid evaporator-crystallizers.





THERMAL ECONOMY

Thermal economy is a major consideration in the design and operation of evaporators. This is improved by operating several vessels in series at successively lower pressures and utilizing vapors from upstream units to reboil the contents of downstream units. Figure 8.17 shows such arrangements. Thermal economy is expressed as a ratio of the amount of water evaporated in the complete unit to the amount of external steam that is supplied. For a single effect, the thermal economy is about 0.8, for two effects it is 1.6, for three effects it is 2.4, and so on. Minimum cost usually is obtained with eight or more effects. When high pressure steam is available, the pressure of the vapor can be boosted with a steam jet compressor [Fig. 8.16(c)] to a usable value; in this way savings of one-half to two-thirds in the amount of external steam can be

achieved. Jet compressor thermal efficiencies are 20-30%. A possible drawback is the contamination of condensate with entrainment from the evaporator. When electricity is affordable, the pressure of the vapor can be boosted mechanically, in compressors with efficiencies of 70-75%.

Because of the elevation of boiling point by dissolved solids, the difference in temperatures of saturated vapor and boiling solution may be 3-10°F which reduces the driving force available for heat transfer. In backward feed [Fig. 8.17(b)] the more concentrated solution is heated with steam at higher pressure which makes for lesser heating surface requirements. Forward feed under the influence of pressure differences in the several vessels requires more surface but avoids the complications of operating pumps under severe conditions.

Several comprehensive examples of heat balances and surface



Figure 8.17. Forward and backward of liquid flow with respect to steam flow in triple-effect evaporators. (a) Forward flow of liquid by action of pressure differences in the vessels. (b) Backward-pumped flow of liquid through the vessels.

requirements of multiple effect evaporation are worked out by Kern (1950).

SURFACE REQUIREMENTS

The data of Tables 8.4-8.7 and particularly 8.10 for boiling liquids are applicable to evaporators when due regard is given the more severe fouling that can occur. For example, cases have been cited in which fouling presents fully half the resistance to heat transfer in evaporators. Some heat transfer data specifically for evaporators are in Figure 8.18. Forced circulation and falling film evaporators have the higher coefficients, and the popular long tube vertical, somewhat poorer performance.

With such data, an estimate can be made of a possible evaporator configuration for a required duty, that is, the diameter, length, and number of tubes can be specified. Then heat transfer correlations can be applied for this geometry and the surface recalculated. Comparison of the estimated and calculated surfaces will establish if another geometry must be estimated and checked. This procedure is described in Example 8.12.

8.11. FIRED HEATERS

a babababa

High process temperatures are obtained by direct transfer of heat from the products of combustion of fuels. Maximum Aame temperatures of hydrocarbons burned with stoichiometric air are



Figure 8.18. Overall heat transfer coefficients in some types of evaporations. (a) Water and sugar juice evaporators; (b) Sea water evaporators. [F.C. Standiford, Chem. Eng., 157-176 (9 Dec. 1963)].

about 3500°F. Specific data are cited by Hougen, Watson, and Ragatz (*Chemical Process Principles*, Vol. I, Wiley, New York, 1954, p. 409) and in *Marks Mechanicat Engineers Handbook, (1978,* p. 4.57). With excess air to ensure complete combustion the temperatures are lower, but still adequate for the attainment of process temperatures above 2000°F when necessary. Lower temperatures are obtained with heat transfer media such as those of Table 8.2 which are in turn serviced in direct-fired heaters.

DESCRIPTION OF EQUIPMENT

In fired heaters and furnaces, heat is released by combustion of fuels into an open space and transferred to fluids inside tubes which are ranged along the walls and roof of the combustion chamber. The heat is transferred by direct radiation and convection and also by reflection from refractory walls lining the chamber.

Three zones are identified in a typical heater such as that of Figure 8.19(a). In the *radiant* zone, heat transfer is predominantly (about 90%) by radiation. The *convection* zone is "out of sight" of the burners; although some transfer occurs by radiation because the temperature still is high enough, most of the transfer here is by convection. The application of extended surfaces permits attainment of heat fluxes per unit of bare surface comparable to those in the radiant zone. *Shield section* is the name given to the first two rows or so leading into the convection. On balance these tubes receive approximately the same heat flux as the radiant

Stack Hood 000 000 Convection 000 section 000 000 000 0 Shield 0 0 tubes 0 0 0 0 Oil outlet 0 Radiant 0 Radiant tubes 0 0 section Oil inlet 'ur ne 0 0 Flue and 0 0 to stack Firebox 0 Recirculating 0 0 flue gas fan 0 0 Recirculating flue gas (a) (b) REFRACTORY STACK STACK 0 ONVECTION TUBES CONVECTION RADIAN TUBES TUBES 00000000 REFRACTORY 0000 0000 0000 O ADIANT TUBES 000000000 0000000 00 č BURNERS 00000 Bυ NERS 000 ò ñ (c) (d)

Figure 8.19. Some types of process fired heaters (See also Fig. 17.16 for a radiation panel heater). (a) Radiant, shield, and convection sections of a box-type heater. (b) Heater with a split convection section for preheating before and soaking after the radiant section (*Lobo and Evans*, 1939). (c) Vertical radiant tubes in a cylindrical shell. (d) Two radiant chambers with a common convection section.

tubes because the higher convection transfer counteracts the lesser radiation due to lack of refractory wall backing. Accordingly, shield tubes are never finned.

The usual temperature of flue gas entering the shield section is $1300-1650^{\circ}F$ and should be $200-300^{\circ}F$ above the process temperature at this point. The proportions of heat transferred in the radiant and convection zones can be regulated by recirculation of hot flue gases into the radiant zone, as sketched on Figure 8.19(b). Such an operation is desirable in the thermal cracking of hydrocarbons, for instance, to maintain a proper temperature profile; a negative gradient may cause condensation of polymeric products that make coke on the tubes. Multiple chambers as in Figure 8.19(d) also provide some flexibility. In many operations, about 75% of the heat is absorbed in the radiant zone of a fired heater.

Horizontal tube supports are made of refractory steel to withstand the high temperatures. Hangers for vertical tubes make for a less expensive construction per unit of tube surface. Furnaces are lined with shaped light weight refractory brick 5-8 in. thick. A 1 in. layer of insulating brick is placed between the lining and the metal shell.

Differences of opinion exist among designers with respect to housing shapes and tube arrangements. Nelson (Petroleum Refinery *Engineering*, McGraw-Hill, New York, 1958, p. 587), for example, describes a dozen types. The most common are cylindrical shells with vertical tubes and cabin or box types with horizontal tubes. Figures 8.19 and 17.16 are of typical constructions. Convection zones are most commonly at the top. Process fluid goes first through the convection section and usually leaves the radiant tubes at the top, particularly when vaporization occurs in them. In the more complex flow pattern of Figure 8.19(b), some of the convection tubes are used for preheat and the remainder to maintain the process fluid at a suitable reaction temperature that was attained in the radiant tubes. Some of the convection zone also may be used for steam generation or superheating or for other heat recovery services in the plant.

Capacities of 10-200 MBtu/hr can be accommodated in heaters with single radiant chambers, and three to four chambers with a common convection section are feasible. Stoichiometric combustion air requirements of typical fuels are tabulated:

	LUV	Combustion Air			
Fuel	(Btu/lb)	lb/lb	ib/1000 Btu		
Methane	21,500	17.2	0.800		
Propane	19,920	15.2	0.763		
Light fuel oil	17,680	14.0	0.792		
Heavy fuel oil	17,420	13.8	0.792		
Anthracite	12,500	4.5	0.360		

Burners may be located in the floor or on the ends of the heaters. Liquid fuels are atomized with steam or air or mechanically. A particularly effective heater design is equipped with radiant panel (surface combustion) burners, illustrated in Figure 17.16(a), (b). The incandescent walls are located 2-3 ft from the tubes. The furnace side of the panel may reach 2200°F whereas the outer side remains at 120°F because of continual cooling by the air-gas mixture. Radiant panel burners require only 2-5% excess

air compared with 10–20% for conventional burners. Heaters equipped with radiant panels cost more but provide better control of temperatures of reactions such as pyrolysis of hydrocarbons to ethylene for instance.

Distances between tube banks are of the order of 20 ft or so. A rough guide to box size is about 4 cuft/sqft of radiant transfer surface, but the ultimate criterion is sufficient space to avoid impingement of flames on the tubes. Some additional notes on dimensions are stated with the design procedure of Table 8.18.

Tubes are mounted approximately one tube diameter from the refractory walls. Usual center-to-center spacing is twice the outside tube diameter. Wider spacings may be employed to lower the ratio of peak flux at the front of the tube to the average flux. For single rows of tubes, some values of these ratios are

Center-to-center/diameter	1	1.5	2	2.5	3
Max flux/avg flux	3.1	2.2	1.8	1.5	1.2

Less is gained by extending the ratio beyond 2.0. Excessive fluxes may damage the metal or result in skin temperatures that are harmful to the process fluid.

A second row of tubes on triangular spacing contributes only about 25% of the heat transfer of the front row. Accordingly, new furnaces employ only the more economical one-row construction. Second rows sometimes are justifiable on revamp of existing equipment to marginally greater duty.

HEAT TRANSFER

Performance of a heater is characterized by the average heat flux in the radiant zone and the overall thermal efficiency. Heat fluxes of representative processes are listed in Table 8.15. Higher fluxes make for a less expensive heater but can generate high skin temperatures inside and out. Thermal sensitivity of the process fluid, the strength of the metal and its resistance to corrosion at elevated temperatures are factors to be taken into account in limiting the peak flux. Because of the refractory nature of water, however, allowable fluxes in steam boilers may reach 130,000 Btu/(hr)(sqft), in comparison with a maximum of about 20,000 in hydrocarbon service. Example 8.13 is a study of the effect of tube spacing on inside film peak temperatures.

A certain amount of excess air is needed to ensure complete combustion. Typical minimum excess requirements are 10% for gaseous fuels and 15–20% for liquids. Radiant panel burners may get by with 2-5% excess air.

Éfficiency is the ratio of total heat absorbed in radiant,

TABLE 8.15. Typical Radiant Fluxes and Process Temperatures

Service	Average Radiant Rate (Btu/hr/ft²) (Based on OD)	Temperature (ºF)
Atmospheric crude heaters	10.000-14.000	400-700
Reboilers	10,000-12,000	400-550
Circulating oil heaters	8000-I 1,000	600
Catalytic reformer change and reheat	7500-1 2,000	800-1000
Delayed coking heater	10,000-11,000	925
Visbreaker heaters-heating section	9000-l 0,000	700-950
Soaking section	6000-7000	950
Lube vacuum heaters	7500-8500	850
Hydrotreater and hydrocracker charge heaters	10,000	700-850
Catalytic-cracker feed heaters	10,000-11,000	900-1050
Steam superheaters	9000-1 3,000	700-1 500
Natural gasoline plant heaters Ethylene and propylene synthesis	1 0,000- 12,000 10,000–15,000	 1300-1650

EXAMPLE 8.13 Peak Temperatures

An average flux rate is 12,000 Btu/(hr)(sqft) and the inside film coefficient is 200 Btu/(hr)(sqft)(°F). At the position where the average process temperature is 850°F, the peak inside film temperature is given by $T = 850 + 12,000^{\circ}R/200$. At the several

convection, and heat recovery sections of the heater to the heat released by combustion. The released heat is based on the lower heating value of the fuel and ambient temperature. With standard burners, efficiencies may be in the range 60-80%; with radiant panels, 80-82%. Within broad limits, any specified efficiency can be attained by controlling excess air and the extent of recovery of waste heat.

An economical apportionment of heat absorption between the radiant and convection zones is about 75% in the radiant zone. This can be controlled in part by recirculation of flue gases into the radiant chamber, as shown in Figure 8.19(b).

Because of practical limitations on numbers and possible locations of burners and because of variations in process temperatures, the distribution of radiant flux in a combustion chamber is not uniform. In many cases, the effect of such nonuniformity is not important, but for sensitive and chemically reacting systems it may need to be taken into account. A method of estimating quickly a flux distribution in a heater of known configuration is illustrated by Nelson (1958, p. 610). A desired pattern can be achieved best in a long narrow heater with a multiplicity of burners, as on Figure 17.16 for instance, or with a multiplicity of chambers. A procedure for design of a plug flow heater is outlined in the Heat Exchanger Design Handbook (1983, 3.11.5). For most practical purposes, however, it is adequate to assume that the gas temperature and the heat flux are constant throughout the radiant chamber. Since the heat transfer is predominantly radiative and varies with the fourth power of the absolute temperature, the effect of even substantial variation in stock temperature on flux distribution is not significant. Example 8.14 studies this problem.

DESIGN OF FIRED HEATERS

The design and rating of a fired heater is a moderately complex operation. Here only the completely mixed model will be treated. For this reason and because of other generalizations, the method to be described affords only an approximation of equipment size and performance. Just what the accuracy is, it is hard to say. Even the relatively elaborate method of Lobo and Evans (1939) is able to predict actual performance only within a maximum deviation of 16%.

EXAMPLE 8.14

Effect of Stock Temperature Variation

A combustion chamber is at 2260°R, a stock enters at 1060°R and leaves at 1360°R. Accordingly, the heat fluxes at the inlet and outlet are approximately in the ratio $(2.26^4 - 1.06^4)/2.26^4 - 1.36^4) = 1.095$. The small effect of even greater variation in flux on a mild cracking operation is illustrated in Figure 8.22.

tube	spacings	the	peak	temperat	ures	are					
	Center-to-	cente	r/diame	eter	1		1.5	2	2.5	3	
	Peak (°F)				1036	5 98	32	958	948	9.22	

For heavy liquid hydrocarbons the upper limit of 950°F often is adopted.

Pertinent equations and other relations are summarized in Table 8.16, and a detailed **stepwise** procedure is listed in Table 8.17. A specific case is worked out in detail in Example 8.15. Basically, a heater configuration and size and some aspects of the performance are assumed in advance. Then calculations are made of the heat transfer that can be realized in such equipment. Adjustments to the design are made as needed and the process calculations repeated. Details are given in the introduction to Example 8.16. Figures 8.20, 8.21, and 8.22 pertain to this example. Some of the approximations used here were developed by Wimpress (1963); his graphs were converted to equation form for convenience. Background and more accurate methods are treated notably by Lobo and Evans (1939) and more briefly by Kern (1950) and Ganapathy (1982). Charts of gas emissivity more elaborate than Figure 8.23 appear in these references.

An early relation between the heat absorption Q in a radiant zone of a heater, the heat release Q_f , the effective surface A, and the air/fuel ratio R lb/lb is due to Wilson, Lobo, and Hottel [Znd. Eng. Chem. 24, 486, (1932)]:

$$Q_f/Q_{=}1 + (R/4200)\sqrt{Q_f/A_{cp}}$$
 (8.45)

Although it is a great simplification, this equation has some utility in appraising directional effects of changes in the variables. Example 8.16 considers changes in performance with changes in excess air.

Heat transfer in the radiant zone of a fired heater occurs largely by radiation from the flue gas (90% or so) but also significantly by convection. The combined effect is represented by

$$Q/A = h_r (T_g^4 - T_s^4) + h_c (T_g - T_s), \qquad (8.46)$$

where T_g and T_s are absolute temperatures of the gas and the receiving surface. The radiative properties of a gas depend on its chemical nature, its concentration, and the temperature. In the thermal range, radiation of flue gas is significant only from the triatomic molecules H_2O , CO,, and SO,, although the amount of the last is small and usually neglected. With fuels having the composition C_xH_{2x} , the ratio of partial pressures is $p_{H_2O}/p_{CO_2} = 1$. In Figure 8.23, the emissivity of such a gas is represented as a function of temperature and the product *PL* of the partial pressures of water and carbon dioxide and the path of travel defined by the mean beam length. Item 8 of Table 8.16 is a curve fit of such data.

When other pertinent factors are included and an approximation is introduced for the relatively minor convection term, the heat transfer equation may be written

$$Q/\alpha A_{cp}F = 1730[(T_g/1000)^4 - (T_s/1000)^4] + 7(T_g - T_s). \quad (8.47)$$

Here the absorptivity depends on the spacing of the tubes and is given by item 5 of Table 8.16. The cold plane area *A*, is the product of the number of tubes by their lengths and by the center-to-center spacing. The combination αA_{cp} is equal to the area of an ideal black plane that has the same absorptivity as the tube

TABLE 8.16. Equations and Other Relations for Fired Heater Design

1. Radiant zone heat transfer

$$\frac{Q_R}{\alpha A_R F} = 1730 \left[\left(\frac{T_g + 460}{1000} \right)^4 - \left(\frac{T_t + 460}{100} \right)^4 \right] + 7(T_g - T_t)^2$$

2. Radiant zone heat balance

$$\frac{Q_{R}}{\alpha A_{R}F} = \frac{Q_{n}}{\alpha A_{R}F} \left(1 + \frac{Q_{g}}{Q_{n}} + \frac{Q_{f}}{Q_{n}} - \frac{Q_{L}}{Q_{n}} - \frac{Q_{g}}{Q_{n}}\right)$$

 $Q_{\rm R}$ is the enthalpy absorbed in the radiant zone, $Q_{\rm s}$ is the enthalpy of the entering air, $Q_{\rm f}$ that of the entering fuel, $Q_{\rm L}$ is the enthalpy loss to the surroundings, $Q_{\rm g}$ is the enthalpy of the gas leaving the radiant zone; $Q_{\rm s}$ and $Q_{\rm f}$ are neglected if there is no preheat, and $Q_{\rm L}/Q_{\rm p}$ is about 0.02-0.03; $Q_{\rm p}$ is the total enthalpy released in the furnace 3. Enthalpy $Q_{\rm s'}$ of the stack gas, given by the overall heat balance

 $Q_s/Q_n = 1 + (1/Q_n)(Q_a + Q_f - Q_L - Q_R - Q_{convection})$

4. Enthalpy ${\it Q}_{a\prime}$ of the flue gas as a function of temperature, °F

 $Q_g/Q_n = [a+b(T/1000-0.1)](T/1000-0.1)$ z = fraction excess air a = 0.22048 ~ 0.35027z + 0.92344z² b = 0.016086 + 0.293932 - 0.48139z²

5. Absorptivity, α_r of the tube surface with a single row of tubes

 $\alpha = 1 - (0.0277 + 0.0927(x - 1)](x - 1)$ x = (center-to-center spacing)/(outside tube diameter)

6. Partial pressure of CO, + H₂O

 $P = 0.288 - 0.229x + 0.090x^2$ x =fraction excess air

7. Mean beam lengths L of radiant chambers

Dimensional Ratio ª Rectangular Furnaces	Mean Length <i>L</i> (ft)
1. I-I-I to I-I-3 I-2-1 to I-2-4	2/3 ∛furnace volume, (ft ³)
2. I-I-4 to 1-1-∞	1.0 x smallest dimension
3. I-2-5 to I-2-8	1.3 x smallest dimension
4. I-3-3 to 1-∞-∞	1.8 x smallest dimension
Cylindrical Furnaces	
5 d×d	2/3 diameter
$\vec{h} \propto \vec{h} \propto \vec{h} $ of $\vec{h} \propto \vec{h}$	1 x diameter
⁸ Length, width, height in an	y order.

8. Emissivity ϕ of the gas (see also Fig. 8.20).

 $\phi \approx a + b(PL) + c(PL)^{2}$ PL = product of the partial pressure (6) and the mean beam length (7) $z = (T_{g} + 460)/1000$ $a = 0.47916 - 0.198472 + 0.022569z^{2}$ $b \approx 0.047029 + 0.0699z - 0.01528z^{2}$ $c = 0.000803 \quad 0.00726 + 0.001597z^{2}$

9. Exchange factor F

 $F = a + b\phi + c\phi^{2}$ $\phi = \text{gas emissivity, (8)}$ $z = A_{w} / \alpha A_{R}$ $a = 0.00064 + 0.0591z + 0.00101z^{2}$ $b = 1.0256 + 0.4908z - 0.058z^{2}$ $c = -0.144 - 0.552z + 0.040z^{2}$

TABLE 8.16—(continued)

10. Overall heat transfer coefficient U_c in the convection zone

 $U_c = (a + bG + cG^2)(4.5/d)^{0.25}$ G= flue gas flow rate, Ib/(sec)(sqft open cross section) d = tube outside diameter, (in.) Z = T_f/1000, average outside film temperature a = 2.461 - 0.759z + 1.625z² b = 0.7655 + 21.3732 - 9.6625z² c = 9.7938 - 30.809z + 14.333z²

11. Flue gas mass rate G_i

 $\frac{10^6 G_f}{4} = \begin{bmatrix} 640 + 8.0x, & \text{with fuel oil} \\ 822 + 7.78x, & \text{with fuel gas} \end{bmatrix}$ Ib/MBtu heat release x = fraction excess air

TABLE 8.17. Procedure for the Rating of a Fired Heater, Utilizing the Equations of Table 8.18

- 1. Choose a tube diameter corresponding to a cold oil velocity of 5-6 ft/sec
- 2. Find the ratio of center-to-center spacing to the outside tube diameter. Usually this is determined by the dimensions of available return bends, either short or long radius
- 3. Specify the desired thermal efficiency. This number may need modification after the corresponding numbers of tubes have been found
- 4. Specify the excess combustion air
- Calculate the total heat absorbed, given the enthalpies of the inlet and outlet process streams and the heat of reaction
- 6. Calculate the corresponding heat release, (heat absorbed)/efficiency
- 7. Assume that 75% of the heat absorption occurs in the radiant zone. This may need to be modified later if the design is not entirely satisfactory
- Specify the average radiant heat flux, which may be in the range of 8000– 20,000 Btu/(hr)(sqft). This value may need modification after the calculation of Step 28 has been made
- 9. Find the needed tube surface area from the heat absorbed and the radiant flux. When a process-side calculation has been made, the required number of tubes will be known and will not be recalculated as stated here
- 10. Take a distance of about 20ft between tube banks. A rough guide to furnace dimensions is a requirement of about 4 cuft/sqft of radiant transfer surface, but the ultimate criterion is sufficient space to avoid flame impingement
- 11. Choose a tube length between 30 and 60ft or so, so as to make the box dimensions roughly comparable. The exposed length of the tube, and the inside length of the furnace shell, is 1.5 ft shorter than the actual length
- 12. Select the number of shield tubes between the radiant and convection zones so that the mass velocity of the flue gas will be about 0.3-0.4 |b/(sec)(sqft free cross section). Usually this will be also the number of convection tubes per row
- 13. The convection tubes usually are finned
- 14. The cold plane area is

 A_{cp} = (exposed tube length)(center-to-center spacing) (number of tubes exclusive of the shield tubes)

16. The refractory area A_w is the inside surface of the shell minus the cold plane area A_{cp} of Step 14

 $A_{w} = 2[W(H+L) + H \times L)] - A_{cp}$

where W, H, and L are the inside dimensions of the shell

- 16. The absorptivity α is obtained from Eq. (5) when only single rows of tubes are used. For the shield tubes, α = 1
- 17. The sum of the products of the areas and the absorptivities in the radiant zone is

 $\alpha A_{R} = A_{\text{shield}} + \alpha A_{co}$

18. For the box-shaped shell, the mean beam length L is approximated by

 $L = \frac{2}{3}$ (furnace volume)^{1/3}

TABLE 8.17-(continued)

- 19. The partial pressure P of CO, + H_2O is given in terms of the excess air by Eq. (6)
- 20. The product PL is found with the results of Steps 18 and 19
- 21. The mean tube wall temperature T_t in the radiant zone is given in terms of the inlet and outlet process stream temperatures by

$$T_{1} = 100 + 0.5(T_{1} + T_{2})$$

22. The temperature T_g of the gas leaving the radiant zone is found by combining the equations of the radiant zone heat transfer [Eq. (1)] and the radiant zone heat balance [Eq. (2)]. With the approximation usually satisfactory, the equality is

$$\frac{Q_n}{\alpha A_R F} \left(1 - 0.02 - \frac{Q_g}{Q_n}\right) = 1730 \left[\left(\frac{T_g + 460}{1000}\right)^4 - \left(\frac{T_t + 460}{1000}\right)^4 \right] + 7(T_g - T_t)$$

The solution of this equation *involves* other functions of $T_{g'}$ namely, the emissivity ϕ by Eq. (8), the exchange factor F by Eq. (9) and the exit enthalpy ratio Q_n/Q_n by Eq. (4)

- 23. The four relations cited in Step 22 are solved simultaneously by trial to find the temperature of the gas. Usually it is in the range 1500–1800°F. The Newton-Raphson method is used in the program of Table 8.18. Alternately, the result can be obtained by interpolation of a series of hand calculations
- 24. After T_{σ} has been found, calculate the heat absorbed $Q_{\rm R}$ by Eq. (1)
- 25. Find the heat flux

 $Q/A = Q_R/A_{\text{radiant}}$

and compare with value specified in Step 8. If there is too much disagreement, repeat the calculations with an adjusted radiant surface area

- 26. By heat balance over the convection zone, find the inlet and outlet temperatures of the process stream
- 27. The enthalpy of the flue gas is given as a function of temperature by Eq. (4). The temperature of the inlet to the convection zone was found in Step 23. The enthalpy of the stack gas is given by the heat balance [Eq. (3)], where all the terms on the right-hand side are known. O_s/O_n is given as a function of the stack temperature T_s by Eq. (4). That temperature is found from this equation by trial
- 28. The average temperature of the gas film in the convection zone is given in terms of the inlet and outlet temperatures of the process stream and the flue gas approximately by

$$T_{f} = 0.5 \left[T_{L1} + T_{L0} + \frac{(T_{g1} - T_{L1}) - (T_{s} - T_{L0})}{\ln[(T_{g1} - T_{L1})/(T_{s} - T_{L0})]} \right]$$

The flow is countercurrent

- 29. Choose the spacing of the convection tubes so that the mass velocity is G = 0.3-0.4 lb/(sec)(sqft free cross section). Usually this spacing is the same as that of the shield tubes, but the value of G will not be the same if the tubes are finned
- 30. The overall heat transfer coefficient is found with Eq. (10)
- 31. The convection tube surface area is found by

 $A_c = Q_c / U_c$ (LMTD)

and the total length of bare of finned tubes, as desired, by dividing A_c by the effective area per foot

32. Procedures for finding the pressure drop on the flue gas side, the draft requirements and other aspects of stack design are presented briefly by Wimpress.

[Based partly on the graphs of Wimpress, Hydrocarbon Process. 42(10), 115-126 (1963)].

EXAMPLE 8.15 Design of a Fiied Heater

The fuel side of a heater used for mild pyrolysis of a fuel oil will be analyzed. The flowsketch of the process is shown in Figure 8.20, and the tube arrangement finally decided upon is in Figure 8.21. Only the temperatures and enthalpies of the process fluid are pertinent to this aspect of the design, but the effect of variation of heat flux along the length of the tubes on the process temperature and conversion is shown in Figure 8.22. In this case, the substantial differences in heat flux have only a minor effect on the process performance.

Basic specifications on the process are the total heat release (102.86 MBtu/hr), overall thermal efficiency (75%), excess air (25%), the fraction of the heat release that is absorbed in the radiant section (75%), and the heat flux (10,000 Btu/(hr)(sqft).

In the present example, the estimated split of 75% and a

EXAMPLE **8.15**—(continued)

radiant rate of 10,000 lead to an initial specification of 87 tubes, but 90 were taken. The final results are quite close to the estimates, being 77.1% to the radiant zone and 9900 Btu/(hr)(sqft) with 90 tubes. If the radiant rate comes out much different from the desired value, the number of tubes is changed accordingly.

Because of the changing temperature of the process stream, the heat flux also deviates from the average value. This variation is estimated roughly from the variation of the quantity

 $\beta = 1730(T_g^4 - T_L^4) + 7.0(T_g - T_L),$

where the gas temperature T_g , in the radiant zone is constant and T_L is the temperature of the process stream, both in "R. In comparison with the average flux, the effect is a slightly increased preheat rate and a reduced flux in the reaction zone. The inside skin temperature also can be estimated on the reasonable assumptions of heat transfer film coefficients of more than 100 before cracking starts and more than 200 at the outlet. For the conditions of this example, with Q/A = 9900 and $T_g = 2011^{\circ}$ R, these results are obtained:

<i>T_</i> (°F)	β/β ₇₂₄	h	T _{skin} (°F)		
547	1.093	>100	<655		
724	1	>100	<823		
900	1.878	>200	<943		

The equation numbers cited following are from Table 8.16. The step numbers used following are the same as those in Table 8.17:

- 1. Flow rate = 195,394/3600(0.9455)(62.4) = 0.9200 cfs,
 - velocity = 5.08 fps in 6-5/8 in. OD Schedule 80 pipe.
- 2. Short radius return bends have 12 in. center-to-center.
- 3. $\eta = 0.75$.
- 4. Fraction excess air = 0.25.
- 5. From the API data book and a heat of cracking of 332 Btu/(lb gas + gasoline):

$$H_{900} = 0.9(590) + 0.08(770) + 0.02(855) = 609.6 \text{ Btu/lb},$$

 $Q_{\text{total}} = 195,394(609.6 - 248) + 19,539(332) = 77.14(E6).$

6. Heat released:

$$Q_n = 77.14/0.75 = 102.86(E6)$$
 Btu/lb.

7. Radiant heat absorption:

$$Q_R = 0.75(77.14)(E6) = 57.86(E6)$$

- 8. (Q/A) rad = 10,000 Btu/(hr)(sqft), average.
- 9. Radiant surface:

- Tube length = 5786/1.7344 = 3336 ft; 40 foot tubes have an exposed length of 38.5 ft; N = 3336/38.5 = 86.6, say 92 radiant tubes.
- **12.** From Eq. (11) the flue gas rate is
 - $G_f = 102.85(1020) = 104,907 \text{ lb/hr}.$

With four shield tubes, equilateral spacing and 3 in. distance to walls,

$$G = \frac{104,907(12)}{3600(38.5)(27.98)} = 0.325$$
 lb/sec sqft.

- 13. The 90 radiant tubes are arranged as shown on Figure 8.22:4 shields, 14 at the ceiling, and 36 on each wall. Dimensions of the shell are shown.
- 14. $A_{r} = (38.5)(1)(90 \quad 4) = 3311$ sqft. **15.** Inside surface of the shell is

$$A_{,} = 2[20(37 + 38.5) + 37(38.5)] = 5869 \text{ sqft}$$

Refractory surface,

$$A_{\rm c} = 5869 - 3311 = 2558$$
 sqft.

16. (Center-to-center)/OD = 12/6.625 = 1.81,

- $\alpha = 0.917$, single rows of tubes [Eq. (5)].
- 17. Effective absorptivity:

 $\alpha A_R = 4(38.5)(1) + 0.917(3311) = 3190$ sqft, $A_w/\alpha A_r = 255813190 = 0.8018.$

18. Mean beam length:

$$L = (2/3)(20 \times 37 \times 38.5)^{1/3} = 20.36.$$

19. From Eq. (6), with 25% excess air,

P = 0.23.

- 20. PL = 0.23(20.36) = 4.68 atm ft.
- 21. Mean tube wall temp: The stream entering the radiant section has absorbed 25% of the total heat.

$$H_1 = 248 + 0.25(77.14)(E6)/195,394 = 346.7,$$

 $T_1 = 565^{\circ}F,$
 $T_r = 100 + (565 + 900)/2 = 832.5.$

22-24. Input data are summarized as:

$$PL = 4.68,$$

$$D_1 = 0.8018,$$

$$D_2 = 0.25,$$

$$T_1 = 832.5,$$

$$Q_1 = Q_n / \alpha A_R = 102.86(E6)/3190 = 32,245.$$

From program "FRN-1",

$$T_{g} = 1553.7,$$

$$F = 0.6496 [Eq. (9)],$$

$$Q_{R} = \alpha A_{R} F \left\{ 1730 \left[\left(\frac{T_{g} + 460}{1000} \right)^{4} - \left(\frac{T_{t} + 460}{1000} \right)^{4} \right] + 7(T_{g} - T_{t}) \right\}$$

$$= 3190(0.6496)(28,679) = 59.43(E6).$$

Compared with estimated 57.86(E6) at 75% heat absorption

EXAMPLE 8.15—(continued)

in the radiant section. Repeat the calculation with an estimate of 60(E6)

$$H_1 = 248 + (77.14 - 60)(E6)/195,394 = 335.7,$$

$$T_1 = 542,$$

$$T_t = 100 + 0.5(542 + 900) = 821,$$

$$T_g = 1550.5,$$

$$F = 0.6498,$$

$$Q_R = 3190(0.6498)(28,727) = 59.55(E6).$$

Interpolating,

Q _{assumed}	<i>T</i> ₁	T _t	T _g (2 _{calcd}	Q/A
57.88	565	832.5	1553.7	59.43	
60.00	542	821	1550.5	59.55	
nterpolation	547		1551.2	59.50	9900]

26-27.

$$Q_{conv} = (77.14 - 59.50)(E6)$$

= 17.64(E6),

Fraction lost in stack gas

$$Q_s/Q_n = 1 - 0.02 - 0.75 = 0.23.$$

bank, and is called the equivalent cold plane area. Evaluation of the exchange factor F is explained in item 9 of Table 8.16. It depends on the emissivity of the gas and the ratio of refractory area A, to the equivalent cold plane area αA_{cp} . In turn, $A_{,,,} = A - A_{,,}$ where A is the area of the inside walls, roof, and floor that are covered by refractory.

In the convection zone of the heater, some heat also is transferred by direct radiation and reflection. The several contributions to overall heat transfer specifically in the convection zone of fired heaters were correlated by Monrad [*Ind. Eng. Chem.* 24,505 (1932)]. The combined effects are approximated by item 10 of Table 8.16, which is adequate for estimating purposes. The relation depends on the temperature of the gas film which is taken to be the sum of the average process temperature and one-half of the log mean temperature difference between process and flue gas over the entire tube bank. The temperature of the gas entering the convection zone From (Eq. (4),

 $T_{s} = 920^{\circ} \text{F}.$

28-31.

LMTD = 735.6

mean gas film temp is

$$T_f = 0.5(400 + 547 + 735.6) = 841.3.$$

Since G = 0.325 lb/(sec)(sqft),

$$V_c = 5.6 \text{ Btu/(hr)(sqft)(°F)} [(Eq. (10)],$$

$$A_{conv} = \frac{17.64(E6)}{735.6(5.6)} = 4282 \text{ sqft},$$

$$\frac{4282}{1.7344(38.5)} = 64.1 \text{ bare tubes}$$

or 16 rows of 4 tubes each. Spacing the same as of the shield tubes.

Beyond the first two rows, extended surfaces can be installed.

Total rows =
$$2 + \frac{14}{2} = 9$$
.

is found with the trial calculation described in Steps 22-23 of Table 8.17 and may utilize the computer program of Table 8.18.

8.12. INSULATION OF EQUIPMENT

Equipment at high or low temperatures is insulated to conserve energy, to keep process conditions from fluctuating with ambient conditions, and to protect personnel who have occasion to approach the equipment. A measure of protection of the equipment metal against atmospheric corrosion also may be a benefit. Application of insulation is a skilled trade. Its cost runs to 8-9% of purchased equipment cost.

In figuring heat transfer between equipment and surroundings, it is adequate to take account of the resistances of only the insulation and the outside film. Coefficients of natural convection are in Table 8.9 and properties of insulating materials at several

EXAMPLE 8.16 Application of the Wilson-Lobo-Hottel Equation

In the case of Example 8.15, 25% excess air was employed, corresponding to 19.0 lb/air/lb fuel, the heat release was $Q_f = 102.86(10^6)$ Btu/hr, and $\alpha A_{cp} = 3036$. The effect will be found of changing the excess air to 16% (16.72 lb air/lb fuel) on the amount of fuel to be fired while maintaining the same heat absorption.

Ratioing Eq. (8.45) to yield the ratio of the releases at the two conditions.

$$\frac{Q_{f2}}{102.86(106) = 1 + (19.0/4200)\sqrt{Q_{f2}/3036}}$$

$$=\frac{1+0.0722\sqrt{Q_{f2}(10^{-6})}}{1.8327}$$

$$\therefore Q_{f2} = 95.82(10^{\circ}) \text{ Btu/hr},$$

which is the heat release with 10% excess air.

With 25% excess air, $Q/Q_f = 1/1.8327 = 0.5456$, With 10% excess air, $Q/Q_f = 0.5456(102.86/95.82) = 0.5857$,

which shows that approximately 7% more of the released heat is absorbed when the excess air is cut from 25% down to 10%.



Figure 8.20. Flowsketch of process of Example 8.16.



Figure 8.21. Tube and box configuration of the fired heater of Example 8.16.



Figure 8.22. Effects of three modes of heat flux distribution on temperature and conversion in pyrolysis of a fuel oil: (1) two levels, 12,500 and 7500; (2) linear variation between the same limits; (3) constant at 10,000 Btu/(hr)(sqft). Obtained by method of Example 8.16.

temperature levels are in Tables 8.19-8.21. Outdoors under windy conditions, heat losses are somewhat greater than indoors at natural convections. Tabulations of economic thicknesses in *Chemical Engineers Handbook* (McGraw-Hill, New York, 1984, 11.55-11.58) suggest that 10–20% greater thickness of insulation is justified at wind velocity of 7.5 miles/hr.

The optimum thickness of insulation can be established by economic analysis when all of the cost data are available, but in practice a rather limited range of thicknesses is employed. Table 8.22 of piping insulation practice in one instance is an example.

The procedure for optimum selection of insulation thicknesses is exemplified by Happel and Jordan [*Chem. Process Economics*, 380 (1975)]. They take into account the costs of insulation and fuel, payout time, and some minor factors. Although their costs of fuel are off by a factor of 10 or more, their conclusions have some validity if it is recognized that material costs likewise have gone up by roughly the same factor. They conclude that with energy cost of \$2.5/million Btu (adjusted by a factor of 10), a payout time of 2 years, for pipe sizes of 2-8 in., the optimum thicknesses in



Figure 8.23. Total emissivity of carbon dioxide and water with $P_{H_2O}/P_{CO_2} = 1$ and a total pressure of 1 atm [*Hadvig*, J. Inst. Fuel 43, 129 (1970)].

TABLE 8.18. Program for Finding the Radiant Gas Temperature by Steps 22 and 23 of Table 8.17

```
10 ! Example 8.16. Design of a
    fired heater.Radiant eas tem
    p by step 22. Program "FRN-1
    ",tape 2
 20 ! P≃PL, product of Partial P
    ressures of CO2+H20 and mean
     beam length
 30
    ( O1=Aw∠αAr
 40
    ł
      D2=fraction excess air
 5й
    1
      T1=tube surface temperatur
 60
    I
      Q1=Qn∕αθr
 70
    1
      Q2=Q3/Qn, Eq.8
 ЗŴ
    1
      F1=emissivity, Ea.6
 90
      F≃exchange factor, Eq.
                               7
    1
    ! J = RHS-LHS of step 22
100
    SHORT T
110
    READ P,01,02,T1,Q1
120
    DATA 4.1,.9605,.25,672,42828
130
    INPUT'T
140
    GOSUB 270
150
160
    J1=J
    T=1.0001#T
170
180-
    GOSUB 270
190
    J2=J
200
    DISP T
210
    H=.0001*T*J1/(J2-J1)
220
    T=T/1.0001-H
230
    IF ABS(H/T)<= 0001 THEN 250
240
    GOTO 150
250
    PRINT "RADIANT GAS TEMP=";T
260
    END
279
    Z1=(T+460)/1000
    A1=.47916-.19847*Z1+.022569*
280
    Z1^2
290
    B1=.047029+.0699*Z1-.01528*Z
    1^2
300 C1=-.000803-.00726*Z1+.00159
    7*21^2
    F1=A1+B1*P+C1*P^2
310
320
    Z2=01
330
    A2=:00064+:0591*Z2+:00101*Z2
    \sim \bigcirc
    B2=1.0256+.4908*Z2-.058*Z2^2
340
    C2=-.144-.552*Z2+.04*Z2^2
350
360
    F=82+82*F1+C2*F1^2
370
    Z3=02
    A3=,22048-,35057*23+,92344*2
380
    3^2
390
    83=.016086+.29393*23-.48139*
    Z3^2
400 Q2=(A3+B3*(T/1000-.1))*(T/10
    00 - 1 >
410 J=-(Q1/F*(.98-Q2))+1730*(((T
    +460)/1000)^4-((T1+460)/1000
    >^4>+7*(T-T1)
420 RETURN
```

insulation depend on the process temperature according to:

and the state of the

Τ (°F)		200	400	600	
Thickness	(in.)	0.5	1.0	1.25	

The data of Table 8.22 are roughly in agreement with these

calculations. Optimum thicknesses of pipe insulation also are tabulated in *Chemical Engineers Handbook (1984,* 11.56); they cover both indoor and outdoor conditions, temperature ranges of 150-1200°F and energy costs of 1-8 dollars/million Btu.

For very large tanks storing volatile liquids and subject to pressure buildup and breathing losses, it is advisable to find economic thickness of insulation by economic analysis. The influence of solar radiation should be taken into account; a brief treatment of this topic is in the book of Threlkeld *(Thermal Environmental Engineering, Prentice-Hall, Englewood Cliffs, NJ, 1970).* In at least one application, rigid urethane foam sprayed onto storage tanks in 2in. thickness and covered with a 4mil thickness of neoprene rubber for weather proofing was economically attractive.

Although resistance to heat transfer goes up as the thickness of pipe insulation is increased, the external surface also increases; a thickness may be reached at which the heat transfer becomes a minimum and then becomes larger. In accordance with this kind of behavior, heat pickup by insulated refrigerated lines of small diameters can be greater than that of bare lines. In another instance, electrical transmission lines often are lagged to increase the rate of heat loss. An example worked out by Kreith (*Principles of Heat Transfer*, Intext, New York, 1973, p. 44) reveals that an insulated 0.5 in. OD cable has a 45% greater heat loss than a bare one.

LOW TEMPERATURES

Insulants suited to cryogenic equipment are characterized by multiple small spaces or pores that occlude more or less stagnant air of comparatively low thermal conductivity. Table 8.19 lists the most common of these materials. In application, vapor barriers are provided in the insulating structure to prevent inward diffusion of atmospheric moisture and freezing on the cold surface with resulting increase in thermal conductivity and deterioration of the insulation. Sealing compounds of an asphalt base are applied to the surface of the insulation which then is covered with a weatherproof jacket or cement coating. For truly cryogenic operations such as air liquefaction and rectification in which temperatures as low as -300°F are encountered, all of the equipment is enclosed in a box, and then the interstices are filled with ground cork.

MEDIUM TEMPERATURES

Up to about 600°F, 85% magnesia has been the most popular material. It is a mixture of magnesia and asbestos fibers so constructed that about 90% of the total volume is dead air space. Equivalents are available for situations where asbestos is undesirable. Such insulants are applied to the equipment in the form of slabs or blankets which are held in place with supports and clips spotwelded to the equipment. They are covered with cement to seal gaps and finished off with a canvas cover that is treated for resistance to the weather. A galvanized metal outer cover may be preferred because of its resistance to mechanical damage of the insulation.

A mixture of diatomaceous earth and an asbestos binder is suitable for temperatures up to the range of 1600-1900°F. Johns-Manville "Superex" is one brand. Since this material is more expensive than 85% magnesia, a composite may be used to save money: sufficient thickness of the high temperature resistant material to bring its external surface to below 600°F, finished off with 85% magnesia in appropriate thickness. Table 8.22(c) is one standard specification of this type.

REFRACTORIES

Equipment made of metal and subject to high temperatures or abrasive or corrosive conditions often is lined with ceramic material.

Material	Bulk/ Density, (Ib/cuft)	Temp (°F)	h	Material	Bulk Densitv, (Ib∕cuft)	Temp (° F)	h
Corkboard	6.9	100	0.022	Rubber board,			
		-100	0.018	expanded,			
		-300	0.010	"Rubatex"	4.9	100	0.018
Fibreglas with						-100	0.015
asphalt coating						-300	0.004
(board)	11.0	100	0.023	Silica aerogel,			
		-100	0.014	powder	5.3	100	0.013
		-300	0.007	"Santocel"		0	0.012
Glass blocks, expanded,						-100	0.010
"Foamglas"	10.6	100	0.036	Vegetable fiber-			
-		-100	0.033	board, asphalt	14.4	100	0.028
		-300	0.018	coating		-100	0.021
Mineral wool				-		-300	0.013
board,							
"Rockcork"	14.3	100	0.024	Foams:	2.9	-100	0.015
		-100	0.017	Polystyrene'	5.0	-100	0.019
		-300	0.008	Polvurethane			

TABLE 8.19. Thermal Conductivities of Insulating Materials for Low Temperatures [k Btu/(hr)(sqft)(°F/ft)]

Test space pressure, 1.0 atm; k = 0.0047 at 10^{-3} mm Hg. Test space pressure, 1.0 atm; k = 0.007 at 10^{-3} mm Hg.

(Marks Mechanical Engineers Handbook, 1978, p. 4.64).

When the pressure is moderate and no condensation is likely, brick construction is satisfactory. Some of the materials suited to this purpose are listed in Table 8.21. Bricks are available to withstand 3000°F. Composites of insulating brick next to the wall and stronger brick inside are practical. Continuous coats of insulants are formed by plastering the walls with a several inch thickness of concretes of various compositions. "Gunite" for instance is a mixture of 1 part cement and 3 parts sand that is sprayed onto walls and even irregular surfaces. Castable refractories of lower density and greater insulating powers also are common. With both brickwork and castables, an inner shell of thin metal may be provided to guard against leakage through cracks that can develop in the refractory

lining. For instance, a catalytic reformer 4ft OD designed for 650 psig and 1100°F has a shell 1.5 in. thick, a light weight castable lining 4-5/8 in. thick and an inner shell of metal 1/8 in. thick. A catalytic cracker 10 ft dia designed for 75 psig and 1100°F has a 3 in. monolithic concrete liner and 3in. of blanket insulation on the outside. Ammonia synthesis reactors that operate at 250atm and 1000°F are insulated on the inside to keep the wall below about 700°F, the temperature at which steels begin to decline in strength, and also to prevent access of hydrogen to the shell since that causes embrittlement. An air gap of about 0.75 in. between the outer shell and the insulating liner contributes significantly to the overall insulating quality.

TABLE 8.20. Thermal Conductivities	of Insulating	Materials for High	h Temperatures	[k Btu/	(hr)(sqft)°F/	'ft)]
------------------------------------	---------------	--------------------	----------------	---------	---------------	-------

Material	Bulk Density, lb/cuft	Max Temp (°F)	100°F	300°F	500°F	1000°F	1500°F	2000°F
Asbestosaper,laminated	2 2	400	0.038	0.042				
Asbestopaperçorrugated	16	300	0.031	0.042				
Diatomaceous earth, silica, powder	18.7	1500	0.037	0.045	0.053	0.074		
Diatomaceous earth, asbestos and bonding material	18	1600	0.045	0.049	0.053	0.065		
Fiberglas block, PF612	2.5	500	0.023	0.039				
Fiberglas block, PF614	4.25	500	0.021	0.033				
Fiberglas block, PF617	9	500	0.020	0.033				
Fiberglas, metal mesh blanket, #900		1000	0.020	0.030	0.040			
Glass blocker,age values	14-24	1600	-	0.046	0.053	0.074		
Hydrous calcium silicate, "Kaylo"	11	1200	0.032	0.038	0.045			
85% magnesia	12	600	0.029	0.035				
Micro-quartz fiber, blanket	3	3000	0.021	0.028	0.042	0.075	0.108	0.142
Potassium titanate, fibers	71.5		_	0.022	0.024	0.030		
Rock wool, loose	8-12		0.027	0.038	0.049	0.078		
Zirconia grain	113	3000	_	-	0.108	0.129	0.163	0.217

/Marks, Mechanical Engineers Handbook, 1978, p. 4.65).

(a) Chemical Composition of Typical Refractories

												Resistanc	e to	
No.	Refractory Type	SiO ₂	Al ₂ O ₃	Fe ₂ O ₃	TiO ₂	CaO	MgO	Cr ₂ O ₃	SiC	Alkalies	Siliceous Steel-Slag	High-lime Steel-Slag	Fused Mill-Scale	Coal-Ash Slag
f	Alumina (fused)	8-10	85-90	1-1.5	1.5-2.2		-		-	0.8-1 .3 ^a	E	G	F	G
2	Chrome	6	2 3	15 ⁶		_	17	38	-	_	G	E	E	G
3	Chrome (unburned)	5	18	12 ^b	_	-	32	30		<u></u>	G	E	E	G
4	Fire clay (high-heat duty)	50-57	36-42	1.5-2.5	1.5-2.5	-	—	—	-	1–3.5 [°]	F	P	P	F
5	Fire clay (super-duty)	52	43	1	2	-	-			2 ^c	F	Р	F	F
6	Forsterite	34.6	0.9	7.0	-	1.3	55.4							
7	High-alumina	22-26	68-72	1-1.5	3.5		—	-	-	1–1.5°	G	F	F	F
8	Kaolin	52	45.4	0.6	1.7	0.1	0.2			_	F	Р	G″	F
9	Magnesite	3	2	6	-	3	86	-	-	-	Р	E	E	E
10	Magnesite (unburned)	5	7.5	8.5		2	64	10		_	Р	E	E	E
11	Magnesite (fused)	-			—		-	-	-	_	F	E	E	E
12	Refractory porcelain	25-70	25-60	-		—		-	-	I - 5	G	F	F	F
13	Silica	96	1	1		2				_	E	Р	F	Р
14	Silicon carbide (clay bonded)	7-9	2-4	0.3-l	1		_	_	85-90	ć	E	G	F	E
15	Silimanite (mullite)	35	62	0.5	1.5	—		-		0.5	G	F	F	F
16	Insulating fire-brick (2600°F)	57.7	36.8	2.4	1.5	0.6	0.5				Р	Р	G	Р

(b) Physical Properties of Typical Refractories^g

		Fusion Point		Deformation under	Deformation under		wt. of
F	Refractory No.	۴	Pyrometric Cone	Load (% at °F and Ib/in.)	Spalling Resistance'	after 5 hr (% °F)	9 in Brick (Ib)
	1	3390+	39+	1 at 2730 and 50	G	+0.5 (2910)	9–10.6
	2	3580+	41+	shears 2740 and 28	P	-0.5-1.0 (3000)	11.0
	3	3580+	41+	shears 2955 and 28	F	-0.5-1.0 (3000)	11.3
	4	3060-3170	31-33	2.5-10 at 2460 and 25	G	±0–1.5 (2550)	7.5
	5	3170-3200	33-34	2-4 at 2640 and 25	E	±0–1.5 (2910)	8.5
	6	3430	4 0	10 at 2950	F		9.0
	7	3290	36	I-4 at 2640 and 25	E	-2-4 (2910)	7.5
	8	3200	34	0.5 at 2640 and 25	E	-0.7-1.0 (2910)	7.7
	9	3580+	41+	shears 2765 and 28	Р	-1-2 (3000)	10.0
	10	3580+	41+	shear 2940 and 28	F	-0.5-1.5 (3000)	10.7
	11	3580+	41+		F	_	10.5
	12	2640-3000	16 + 30		G		
	13	3060-3090	31-32	shears 2900 and 25	P	+0.5-0.8 (2640)	6.5
	14	3390	39	O-I at 2730 and 50	E	+2 (2910)	8-9.3
	15	331 O-3340	37-38	O-O.5 at 2640 and 25	E	-0-0.8 (2910)	8.5
	16	2980-3000	29-30	0.3 at 2200 and 10	G	-0.2 (2600)	2.25

^a Divide by 12 to obtain the units k Btu/(hr)(sqft)(°F/ft). As FeO. Includes lime and magnesia. Excellent if left above 1200°F.

[°]/[°]Oxidizing atmosphere. [°]/[°] E = Excellent. G = Good. F = Fair. P = Poor. [°][Some data from Trostel, *Chem. Met. Eng. (Nov.* 1938)]. [°]Marks, *Mechanical Engineers Handbook,* McGraw-Hill, New York, 1978, pp. 6.172–6.173.

TABLE 8.22. **Specifications** of Thicknesses of Pipe Insulation for Moderate and High **Temp**eratures, in Single or Double Strength as May **B**e Needed

(a) Insulation of 85% Magnesia or Equivalent up to 600°F

Pipe Size (in.)	Standard Thick (in.)	Double Standard Thick (in.)
1-1/2 or less	7/8	l - 15116
2	1-1/32	2-5/32
2-1/2	1–1/32	2–5/32
3	1–1/32	25/32
4	1-1/8	2–1/4
5	1–1/8	2–5/16
6	1–1/8	2–5/16
8	1–1/4	2–1/2
10	1–1/4	2–1/2
12-33	1-1/2	3

(b)	Molded	Diatomaceous	Earth	Base	Insulation,	to	1900°F,
Sin	gle or Do	ouble Thickness	as Ne	eded			

Pipe Size (in.)	Thickne	ess (in.)
1-1/2	2	2
2	1-1/4	2–1/8
2–1/2	1-5/16	l - 13116
3	1-9/16	2–1/16
4	1-9/16	2-1/16
5	1-1/2	2
6	1-1/2	2-1/16
8	1-1/2	2
10	1-9/16	2–1/8
12	1-9/16	2–1/8
14-33	11/2	2

(c) Combination Insulation, Inner Layer of Diatomaceous Earth Base, and Outer of 85% Magnesia or Equivalent, for High Level Insulation to **1900°F**

Inner	Layer	Outer	Layer
Pipe Size (in.)	Thickness (in.)	Nominal Pipe Size (in.)	Thickness (in.)
1-1/2 or less	2	no outer layer	
2	1-1/4	4-1/2	1-1/2
2–1/2	1-5/16	5	1-1/2
3	1–9/16	6	1-1/2
4	1-9/16	7	1–1/2
5	1–1/2	8	2
6	1–1/2	9	2
8	1-1/2	11	2
10	1-9/16	14	2
12	1-9/16	16	2
14-33	1–1/2	17-36	2

Data of an engineering contractor.

8.13. REFRIGERATION

Process temperatures below those attainable with cooling water or air are attained through refrigerants whose low temperatures are obtained by several means:

- Vapor compression refrigeration in which a vapor is compressed, then condensed with water or air, and expanded to a low pressure and correspondingly low temperature through a valve or an engine with power takeoff.
- Absorption refrigeration in which condensation is effected by absorption of vapor in a liquid at high pressure, then cooling and

expanding to a low pressure at which the solution becomes cold and flashed.

3. Steam jet action in which water is chilled by evaporation in a chamber maintained at low pressure by means of a steam jet ejector. A temperature is 55°F or so is commonly attained, but down to 40°F may be feasible. Brines also can be chilled by evaporation to below 32°F.

The unit of refrigeration is the ton which is approximately the removal of the heat of fusion of a ton of ice in one day, or 288,000 **Btu/day**, 12,000 **Btu/hr**, 200 Btu/min. The reciprocal of the efficiency, called the coefficient of performance (COP) is the term employed to characterize the performances of refrigerating processes:

COP

A commonly used unit of COP is (tons of refrigeration)/ (horsepower input). Some of the refrigerants suited to particular temperature ranges are listed in Tables 1.10, 8.23, and 8.24.

COMPRESSION REFRIGERATION

A basic circuit of vapor compression refrigeration is in Figure 8.24(a). After compression, vapor is condensed with water cooling and then expanded to a low temperature through a valve in which the process is essentially at constant enthalpy. In large scale installations or when the objective is liquefaction of the "permanent" gases, expansion to lower temperatures is achieved in turboexpanders from which power is recovered; such expansions are approximately isentropic. The process with expansion through a valve is represented on a pressure-enthalpy diagram in Figure 8.24(b).

A process employing a circulating brine is illustrated in Figure 8.24(c); it is employed when cooling is required at several points distant from the refrigeration unit because of the lower cost of circulation of the brine, and when leakage between refrigerant and process fluids is harmful.

For an overall compression ratio much in excess of four or so, multistage compression is more economic. Figure 8.24(d) shows two stages with intercooling to improve the capacity and efficiency of the process.

Many variations of the simple circuits are employed in the interest of better performance. The case of Example 8.17 has two stages of compression but also two stages of expansion, a scheme due originally to Windhausen (in 1901). The flashed vapor of the intermediate stage is recycled to the high pressure compressor. The numerical example shows that an improved COP is attained with the modified circuit. In the circuit with a centrifugal compressor of Figure 8.25, the functions of several intermediate expansion valves and flash drums are combined in a single vessel with appropriate internals called an economizer. This refrigeration unit is used with a fractionating unit for recovering ethane and ethylene from a mixture with lighter substances.

Low temperatures with the possibility of still using water for final condensation are attained with cascade systems employing coupled circuits with different refrigerants. Refrigerants with higher vapor pressures effect condensation of those with lower vapor pressures. Figure 8.26 employs ethylene and propylene in a cascade for servicing the condenser of a demethanizer which must be cooled to -145°F. A similar process is represented on a flowsketch in the **book** of Ludwig (1983, Vol. 1, p. 249). A three element cascade with methane, ethylene and propylene refrigerants is calculated by



Figure 8.24. Simpler circuits of compression refrigeration (see also Example 8.17). (a) Basic circuit consisting of a compressor, condenser, expansion valve and evaporator (load). (b) Conditions of the basic circuit as they appear on a pressure-enthalpy diagram; the primed points are on the vapor-liquid boundary curve. (c) Circuit with circulation of refrigerated brine to process loads. (d) Circuit with two-stage compression and intercooling.

EXAMPLE 8.17

Two-Stage Propylene Compression Refrigeration with Interstage Recycle

A propylene refrigeration cycle operates with pressures of 2.56, 64, and 16psia. Upon expansion to 64 psia, the flashed vapor is

recycled to the suction of the high pressure stage while the liquid is expanded to 16psia to provide the needed refrigeration at -9° F. The ratios of refrigeration to power input will be compared without and with interstage recycle.

Basis: 1 lb of propylene to the high pressure stage. Conditions



226 HEAT TRANSFER AND HEAT EXCHANGERS

EXAMPLE **8.17**—(continued)

are shown on the pressure-enthalpy and flow diagrams. Isentropic compression and isenthalpic expansion are taken. Without recycle,

refrigeration = 452-347 = 105 **Btu/lb**, work = 512-452 = 60 **Btu/lb**, COP = 105/60 = 1.75.

With recycle,



Figure 8.25. A refrigeration system for the overhead condenser of a fractionator for recovering ethane and ethylene. Freon-12 is the refrigerant. The economizer combines the functions of several expansion valves and flash drums for intermediate recycle of flashed vapors.

interstage vapor = (347-305)/(468-305) = 0.2577 lb/lb, refrigeration = (452-305)0.7423 = 109.1 Btu/lb, work = (495-468)0.2577 + (512-452)0.7423 = 51.5 Btu/lb, COP = 109.1/51.5 = 2.12,

which points out the improvement in coefficient of performance by the interstage recycle.

Bogart (1981, pp. 44-47); it attains -240°F with a maximum pressure of 527 psia.

REFRIGERANTS

Several refrigerants commonly used above -80°F or so are compared in Table 8.23. Ethylene and butane also are in use, particularly in refineries where they are recoverable from the process streams. Properties of the freons (also known by the trade name genetrons) are listed in Table 8.24. Freon 12 is listed in both tables so some comparisons of all of these refrigerants is possible. The refrigerants of Table 8.23 have similar performance. When ammonia or some hydrocarbons are made in the plant, their election as refrigerants is logical. Usually it is preferred to operate at suction pressures above atmospheric to avoid inleakage of air. The nonflammability and nontoxicity of the freons is an attractive quality. Relatively dense vapors such as Ref-12, -22, and -500 are preferred with reciprocating compressors which then may have smaller cylinders. For most equipment sizes, Ref-12 or -114 can be adopted for greater capacity with the same equipment. Ref-22 and -500 are used with specially built centrifugals to obtain highest capacities.

Ammonia absorption refrigeration is particularly applicable when low level heat is available for operation of the stripper reboiler and power costs are high. Steam jet refrigeration is the large scale system of choice when chilled water is cold enough, that is above 40°F or so.



Figure 8.26. A cascade refrigeration system employing ethylene and propylene for condensing the overhead of a demethanizer at -145°F. The diagram is somewhat simplified.

TABLE 8.23. Comparative Data of Refrigerant9

Evaporator Temp (°F)		- 8 0	- 6 0	-40	-20	0	2 0	40	6 0
Evaporator	ammonia		5.55	10.4	18.3	30.4	49.0	73.0	107.5
pressure	propylene	7.20	12.5	20.7	32.1	49.0	70.0	96.0	131
(psia)	propane	5.55	9.78	16.2	25.5	38.1	56.0	80.0	110
	freon 12	2.88	5.36	9.3	15.3	23.8	35.7	51.7	72.4
	Condensed Ar	Liquid T nmonia 19	emperature 7; Propyler	95°F; Cond ne 212; Pro	denser Pres pane 177;	sure in p 12 123	sia:		
lb refrigerant/	ammonia		0.454	0.446	0.438	0.432	0.426	0.422	0.418
min/ton	propylene	2.07	1.96	1.87	1.79	1.72	1.66	1.60	1.54
refrigeration	propane	2.18	2.04	1.93	1.83	1.74	1.67	1.59	1.53
0	freon 12	5.18	4.89	4.65	4.42	4.22	4.05	3.88	3.74
CME of	ammonia		20.4	11 1	6 4 5	3 96	2 5 2	1 69	1 14
refrigerant/	propylene	27 1	15.7	9.18	5 85	3 84	2.52	1.00	1.14
min/ton	propylene	37 4	20.0	12.0	7 29	4 77	2.00	2 1 3	1.50
refrigeration	freon 12	59.9	31.7	18.0	10.8	6.79	4.44	3.00	2.09
Proko	ommonio		4.24	2 2 2	2.44	1 70	1.06	0 9 2 5	0.482
brsopowor/	annonia	5 00	2.06	3.23	2.41	1.70	1.20	0.035	0.405
top refrigeration	propano	1 0 0	2 97	3.10	2.35	1.74	1.20	0.030	0.485
ton reingeration	freen 12	4.90	1 33	3.03	2.32	1.75	1.24	0.000	0.458
	Condensed Ammo	Liquid Te onia 303; F	mperature Propylene 3	125°F; Con 14; Propar	denser Pres ne 260; Fre	ssure in p on 12 184	osia:		
lb refrigerant	ammonia		0.492	0.483	0.474	0.466	0.460	0.454	0.450
mir/ton	propylene	2.67	2.50	2.35	2.22	2.11	2.01	1.93	1.86
refrigeration	propane	2.86	2.63	2.44	2.29	2.16	2.04	1.94	1.84
Ū.	freon 12	6.42	5.98	5.61	5.28	5.00	4.75	4.53	4.33
CFM of	ammonia		22.0	12.0	6.97	4.26	2.72	1.82	1.23
refrigerant/	propylene	35 2	20.0	11.5	7 32	4 72	3 08	2 18	1.56
ton/	propane	50.0	25.8	15.4	9.16	5.94	3.79	2.63	1.80
refrigeration	freon 12	74.0	38.8	21.7	12.9	8.05	5.21	3.50	2.42
Brake	ammonia		5 68	4 38	3 33	2 54	1.90	1.38	0 952
horsepower/	nronvlene	7 4 9	5.96	4 71	3.66	2.04	2 03	1.55	1 10
ton refrigeration	propyrene	7 47	5 85	4 60	3 59	2.73	2.00	1.50	1.03
	NUDUIU	1.71	0.00	T. U U	0.00	2.01	4.01	1.00	1.00

"The horsepowers are based on centrifugal compressor efficiencies without economizers.

a subodies



Figure 8.27. An ammonia absorption refrigeration process for a load of 50 tons at 30°F. The conditions were established by Hougen, Watson, and Ragatz (Thermodynamics, *Wiley, New York, 1959, pp. 83.5-842).*

Refrigerant Number (ARI Designation)	11	1 2	2 2	113	114	500	502
Chemical name	trichloromono- fluoromethane	dichlorodi- fluoromethane	monochlorodi- fluoromethane	trichlorotri- fluoroethane	dichlorotetra- fluoroethane	azeotrope of dichlorodi- fluoromethane and difluoroethane	azeotrope of monochlorodi- fluoromethane and monochloropenta- fluoroethane
Chemical formula	CCl₃F	CCl ₂ F ₂	CHCIF,		C ₂ Cl ₂ F ₄	73.8% CCI₂F₂ 26.2% CH₃CHF₂	48.8% CHCLF, 51.2% CCIF ₂ –CF ₃
Molecular wl Gas constant // ((ft lb/lb R)) Boiling point at 1 atm (°F)	137.38 11.25 74.7	120.93 12.78 -21.62	86.48 17.87 -41.4	187.39 8.25 117.6	170.93 9.04 38.4	99.29 15.57 -28.0	111.64 13.87 -50.1
Freezing point at 1 atm (°F) Critical temperature (°F) Critical pressure (psia)	-168 388.0 635.0	-252 233.6 597.0	-256 204.8 716.0	-31 417.4 495.0	-137 294.3 474.0	-254 221.1 631.0	a 194.1 618.7
Specific heat of liquid, 86°F Specific heat of vapor, C_p 60°F at 1 atm Specific heat at vapor, C_v 60°F at 1 atm	0.220 a a	0.235 0.146 0.130	0.335 0.149 0.127	0.218 a a	0.238 0.156 0.145	0.300 0.171 0.151	0.305 0.164 0.161
Ratio $C_p/C_v = K$ (86°F at 1 atm)	1.11	1.14	1.18	1.12	1.09	1.13	1.023
Saturation pressure (psia) at -50°F 0°F 40°F 105°F	0.52 2.55 7.03 25.7	7.12 23.85 51.87 141.25	11.74 38.79 83.72 227.65	a 0.84 2.86 11.58	1.35 5.96 15.22 50.29	8.395 27.96 60.94 167.85	14.74 45.94 94.90 244.40
Net refrigerating effect (Btu/Ib) 40-105°F (no subcooling)	67.56	49.13	66.44	54.54	43.46	59.82	43.72
Cycle efficiency (% Carnot cycle) 40–105°F	90.5	83.2	81.8	87.5	94.9	82.0	76.1
Liquid circulated 40–105°F {(lb/min/ton)]	2.96	4.07	3.02	3.66	4.62	3.35	4.58
Theoretical displacement 40–105°F (cuft/min/ton)]	16.1	3.14	1.98	39.5	9.16	2.69	2.04
Theoretical horsepower per ton 40–105°F	0.676	0.736	0.75	0.70	0.722	0.747	0.806
Coefficient at performance 40-105°F (4.71/HP per ton)	6.95	6.39	6.29	8.74	6.52	6.31	5.86
Cost compared with R 11	1.00	1.57	2.77	2.15	2.97	2.00	5.54

^a Data not available or not applicable. (Carrier Air Conditioning Co.).

TABLE 8.24. Comparative Data of Freon Refrigerants

ABSORPTION REFRIGERATION

The most widely used is ammonia absorption in water. A flowsketch of the process is in Figure 8.27. Liquid ammonia at a high pressure is obtained overhead in a stripper, and then is expanded through a valve and becomes the low temperature vapor-liquid mixture that functions as the refrigerant. The low pressure vapor is absorbed in weak liquor from the bottom of the stripper. Energy input to the refrigeration system is primarily that of the steam to the stripper reboiler and a minor amount of power to the pump and the cooling water circulation.

This kind of system has a useful range down to the atmospheric boiling point of ammonia, -28° F or -33° C, or even lower. Two or three stage units are proposed for down to -94° F. Sizing of equipment is treated by Bogart (1981).

REFERENCES

Constant and the second second

- K.J. Bell and M.A. Ghaly, An approximate generalized design method for multicomponent partial condensers, *Chem. Eng. Prog. Symp. Ser.* 131, 72-79 (1973).
- V. Cavaseno et al. (Eds.), Process *Heat Exchange*, McGraw-Hill, New York, 1979.
- D. Chisholm (Ed.), Developments in Heat Exchange Technology I, Applied Science, London, 1980.
- J.R. Fair, Process heat transfer by direct fluid-phase contact, *Chem. Eng. Prog. Symp.* Ser. 118, 1-11 (1972); *Chem. Eng.*, (12 June 1972).
- V. Ganapathy, Applied Heat Transfer, PennWell Books, Tulsa, OK, 1982.
- H. Gröber, S. Erk, and U. Grigull, Fundamentals of Heat Transfer, McGraw-Hill, New York, 1961.
- H. Hausen, Heat Transfer in Counterflow, Parallel Flow and Cross Flow, McGraw-Hill, New York, 1983.
- HEDH, Heat Exchanger Design Handbook (E.U. Schlinder et al., Eds.), Hemisphere, New York, 1983-date, 5 vols.
- 9. M. Jakob, Heat Transfer, Wiley, New York, 1957, Vol. 2.
- S. Kakac, A.E. Bergles, and F. Mayinger (Eds.), *Heat Exchangers: Thermal-Hydraulic Fundamentals and Design*, Hemisphere, New York, 1981.
- W.M. Kays and A.L. London, *Compact Heat Exchangers*, McGraw-Hill, New York, 1984.
- 12. D.Q. Kern, Process Heat Transfer, McGraw-Hill, New York, 1950.
- S.K. Kutateladze and V.M. Borishanskii, Concise Encyclopedia of Heat Transfer, Pergamon, New York, 1966.
- E.E. Ludwig, Applied Process Design for Chemical and Petrochemical Plants, Gulf, Houston, 1983, Vol. 3, pp. 1-200.
- P.E. Minton, Designing spiral plate and spiral tube exchangers, *Chem. Eng.*, (4 May 1970); (18 May 1970).
- R.K. Neeld and J.T. O'Bara, Jet trays in heat transfer service, *Chem. Eng. Prog.* 66(7), 53 1970.
- P.A. Schweitzer (Ed.), Handbook of Separation Techniques for Chemical Engineers, McGraw-Hill, New York, 1979, Sec. 2.3, Evaporators, Sec. 24, Crystallizers.
- L. Silver, Gas cooling with aqueous condensation, Tram. Inst. Chem. Eng. 25, 30-42 (1947).
- 19. E.F.C. Somerscales and J.G. Knudsen (Eds.), Fouling of Heat Transfer Equipment, Hemisphere, New York, 1981.
- 20. J. Taborek, G.F. Hewitt, and N. Afgan (Eds.), *Heat Exchangers Theory and Practice*, Hemisphere, New York, 1983.
- 21. TEMA Standards, Tubular Exchanger Manufacturers Association, Tarrytown, NY, 1978.
- 22. G. Walker, Industrial Heat Exchangers, Hemisphere, New York, 1982.*

Another kind of absorption refrigerant system employs aqueous lithium bromide as absorbent and circulating water as the refrigerant. It is used widely for air conditioning systems, in units of 600-700 tons producing water at 45°F.

CRYOGENICS

This term is applied to the production and utilization of temperatures in the range of liquid air, -200°F and lower. A great deal of information is available on this subject of special interest, for instance in Chemical Engineers *Handbook (1984, 12.47-12.58)* and in the book of Arkhanov et al. (1981).

Fired Heaters (see also Ganapathy, HEDH, and Kern above)

- F.A. Holland, R.M. Moores, F.A. Watson, and J.K. Wilkinson, *Heat Transfer*, Heinemann, London, 1970.
- H.C. Hottel, in *McAdams Heat Transmission*, McGraw-Hill, New York, 1954.
- W.E. Lobo and J.E. Evans, Heat transfer in the radiant section of petroleum heaters, *Trans. AlChE* 35, 743 (1939).
- C.C. Monrad, Heat transmission in the convection section of pipe stills, Ind. Eng. Chem. 24, 505 (1932).
- D.W. Wilson, W.E. Lobo, and H.C. Hottel, Heat transmission in the radiant section of tube stills, *Ind. Eng. Chem. 24, 486* (1932).
- R.N. Wimpress, Rating fired heaters, *Hydrocarbon Process.* 42(10), 115-126 (1963); Generalized method predicts fired-heater performance, *Chem. Eng.*, 95-102 (22 May 1978).

Selected American Petroleum Institute Standards (API, Washington, D.C.)

- Std. 660, Shell-and-Tube Heat Exchangers for General Refinery Services, 1982.
- **30.** Std. 661, Air-Cooled Heat Exchangers for General Refinery Services, 1978.
- 31. Std. 665, API Fired Heater Data Sheet, 1966, 1973.

Insulation

- 32. Marks Mechanical Engineers Handbook, McGraw-Hill, New York, 1978, pp. 6.169-6.177.
- H.F. Rase and M.H. Barrow, *Project Engineering* of *Process Plants*, Wiley, New York, 1957, Chap. 19.
- 34. G.B. Wilkes, Heat Insulation, Wiley, New York, 1950.

Refrigeration

- A. Arkhanov, I. Marfenina, Ye. Mikulin, Theory and Design of Cryogenic Systems, Mir Publishers, Moscow, 1981.
- ASHRE, *Thermophysical Properties of Refrigerants*, American Society of Heating, Refrigeration and Air-Conditioning Engineers, Atlanta, GA, 1976.
- M. Bogart, Ammonia Absorption Refrigeration in Industrial Processes, Gulf, Houston, 1981.
- Carrier System Design Manual, Carrier Air Conditioning Co., Syracuse, NY, 1964, Part 4, Refrigerants, brines and oils.
- F.L. Evans, Equipment Design Handbook for Refineries and Chemical Plants, Gulf, Houston, 1979, Vol. 1, pp. 172-196.
- T.M. Flynn and K.D. Timmerhaus, Cryogenic processes, in *Chemical Engineers Handbook*, 1984, pp. 12.46-12.58.
- W.B. Gosney, *Principles of Refrigeration*, Cambridge University Press, Cambridge, 1982.
- 42. E.E. Ludwig, Applied Process Design for Chemical and Petroleum Plants, Gulf, Houston, 1983, Vol. 1, pp. 201-250.
- Y.R. Mehra, Refrigerating properties of ethylene, ethane, propylene and propane, *Chem. Eng.*, 97 (18 Dec. 1978); 131 (15 Jan. 1979); 95 (12 Feb. 1979); 165 (26 Mar. 1979).

^{*} The book by Walker (Appendix D, 1982) has a guide to the literature of heat transfer in book form and describes the proprietary services **HTFS** (Heat Transfer and Fluid Services) and HTRI (Heat Transfer Research Inc.).

DRYERS AND COOLING TOWERS

The processes of the drying of solids and the evaporative cooling of process water with air have a common foundation in that both deal with interaction of water and air and involve simultaneous heat and mass transfer. Water cooling is accomplished primarily in packed towers and also in spray ponds or in vacuum spray chambers, the latter for exceptionally low temperatures. Although such equipment is comparatively simple in concept it is usually large and expensive, so that efficiencies and other aspects are considered proprietary by the small number of manufacturers in this field.

In contrast, a great variety of equipment is used for the drying of so/ids. Thomas Register lists about 35 pages of U.S. manufacturers of drying equipment, classified with respect to type or the nature of the material being dried. In a major respect, dryers are so/ids handling and transporting equipment, notable examples being perforated belt conveyors and pneumatic conveyors through which hot air is blown. Solids being dried cover a range of sizes from micron-sized particles to large slabs and may have varied and distinctive drying behaviors. As in some other long-established industries, drying practices of necessity have outpaced drying theory. In the present state of the art, it is not possible to design a dryer by theory without experience, but a reasonably satisfactory design is possible from experience plus a little theory.

Performances of dryers with simple flow patterns can be described with the aid of laboratory drying rate data. In other cases, theoretical principles and correlations of rate data are of value largely for appraisal of the effects of changes in some operating conditions when a basic operation is known. The essential required information is the residence time in the particular kind of dryer under consideration. Along with application of possible available rules for vessel proportions and internals to assure adequate contacting of solids and air, heat and material balances then complete a process design of a dryer.

In order to aid in the design of dryers by analogy, examples of dimensions and performances of the most common types of dryers are cited in this chapter. Theory and correlation of heat and mass transfer are treated in detail elsewhere in this book, but their use in the description of drying behavior will be indicated here.

9.1. INTERACTION OF AIR AND WATER

Besides the obvious processes of humidification and dehumidification of air for control of environment, interaction of air and water is a major aspect of the drying of wet solids and the cooling of water for process needs. Heat and mass transfer then occur simultaneously. For equilibrium under adiabatic conditions, the energy balance is

$$k_g \lambda (p_s - p) = h(T - T_w), \qquad (9.1)$$

where p_s is the vapor pressure at the wet bulb temperature T_w . The moisture ratio, H lb water/lb dry air, is related to the partial pressure of the water in the air by

$$H = \frac{18}{29} \frac{p}{P - p} \approx \frac{18}{29} \frac{p}{P}, \qquad (9.2)$$

the approximation being valid for relatively small partial pressures. Accordingly, the equation of the adiabatic saturation line may be written

$$H_s - H = (h/\lambda k)(T - T_w)$$
(9.3)

$$= (C/\lambda)(T-T_s). \tag{9.4}$$

For water, numerically $C \simeq h/k$, so that the wet bulb and adiabatic saturation temperatures are identical. For other vapors this conclusion is not correct.

For practical purposes, the properties of humid air are recorded on **psychrometric** (or humidity) charts such as those of Figures 9.1 and 9.2, but tabulated data and equations also are available for greater accuracy. A computer version is available (Wiley Professional Software, Wiley, New York). The terminal properties of a particular adiabatic humification of air are located on the same saturation line, one of those sloping upwards to the left on the charts. For example, all of these points are on the same saturation line: (T, H) = (250, 0.008), (170, 0.026) and (100, 0.043); the saturation enthalpy is 72 Btu/lb dry, but the individual enthalpies are less by the amounts 2.5, 1.2, and 0, respectively.

Properties such as moisture content, specific volume, and enthalpy are referred to unit mass of dry air. The units employed on Figure 9.1 are lb, cuft, °F, and Btu; those on Figure 9.2 are SI. The data are for standard atmospheric pressure. How to correct them for minor deviations from standard pressure is explained for example in *Chemical Engineers' Handbook* (McGraw-Hill, New York, 1984, 12.10). An example of reading the charts is with the legend of Figure 9.1. Definitions of common humidity terms and their units are given following.

1. Humidity is the ratio of mass of water to the mass of dry air,

$$H = W_w / W_a. \tag{9.5}$$

2. Relative humidity or relative saturation is the ratio of the prevailing humidity to the saturation humidity at the same temperature, or the ratio of the partial pressure to the vapor pressure expressed as a percentage,

$$\% RH = 100 H/H_s = 100 p/p_s. \tag{9.6}$$

3. The relative absolute humidity is

$$(H/H_s)_{\text{absolute}} = \left(\frac{p}{P-p}\right) / \left(\frac{p_s}{P-p_s}\right)$$
(9.7)

4. Vapor pressure of water is given as a function of temperature by

$$p_s = \exp(11.9176 - 7173.9/(T + 389.5)), \text{ atm}, \text{ "F.} (9.8)$$



Figure 9.1. Psychrometric chart in English units (*Currier Corp.* Syracuse, NY). Example: For air at 200°F with H = 0.03 lb/lb: $T_s = 106.5$ °F, $V_h = 17.4$ cuft/lb dry, $100H/H_s = 5.9\%$, $h = h_s + D = 84 - 1.7 = 82.3$ Btu/lb dry.



Figure 9.2. Psychrometric chart for a wide temperature range, 32-600°F (Proctor and Schwartz, Inc., Horsham, PA).
Conditions in an Adiabatic Dryer The air to a dryer has a temperature of 250°F and a wet bulb temperature of 101.5°F and leaves the process at 110°F. Water is



5. The humid volume is the volume of 1 lb of dry air plus the volume of its associated water vapor,

$$V_h = 0.73(1/29 + h/18)(T + 459.6)/P,$$

cuft/(lb dry air). (9.9)

6. Humid specific heat is

$$C_h = C_a + C_w H = 0.24 + 0.45H$$
, Btu/(F)(lbdryair). (9.10)

- 7. The wet bulb temperature T_w is attained by measurement under standardized conditions. For water, T_w is numerically nearly the same as the adiabatic saturation temperature T_s .
- **8.** The adiabatic saturation temperature T_s is the temperature attained if the gas were saturated by an adiabatic process.
- 9. With heat capacity given by item 6, the enthalpy of humid air is

$$h = 0.24T + (0.45T + 1100)H.$$
(9.11)

On the psychrometric chart of Figure 9.1, values of the saturation enthalpy h_s and a correction factor *D* are plotted. In these terms the enthalpy is

$$h = h_s + D. \tag{9.12}$$

In Figure 9.2, the enthalpy may be found by interpolation between the lines for saturated and dry air.

In some periods of drying certain kinds of solids, water is brought to the surface quickly so that the drying process is essentially evaporation of water from the free surface. In the absence of intentional heat exchange with the surrounding or substantial heat losses, the condition of the air will vary along the adiabatic saturation line. Such a process is analyzed in Example 9.1.

For economic reasons, equilibrium conditions cannot be approached closely. In a cooling tower, for instance, the effluent air is not quite saturated, and the water temperature is not quite at the wet bulb temperature. Percent saturation in the vicinity of 90% often is feasible. Approach is the difference between the temperatures of the water and the wet bulb. It is a significant determinant of cooling tower size as these selected data indicate:

Approach (°F)	5	10	15	20	25
Relative tower volume	2.4	1.6	1.0	0.7	0. 55

Other criteria for dryers and cooling towers will be cited later.

evaporated off the surface of the solid at the rate of 1500 lb/hr. Linear velocity of the gas is limited to a maximum of 15 ft/sec. The diameter of the vessel will be found.

Terminal conditions of the air are read off the adiabatic saturation line and appear on the sketch:

Dry air
$$=0.043 \frac{1500}{0.010} = 45,455$$
 lb/hr
 $\rightarrow \frac{45,455(18.2)}{3600} = 229.8$ cfs
 $D = \sqrt{229.8/15(\pi/4)} = 4.4$ ft.

9.2. RATE OF DRYING

In a typical drying experiment, the moisture content and possibly the temperature of the material are measured as functions of the time. The inlet and outlet rates and compositions of the gas also are noted. From such data, the variation of the rate of drying with either the moisture content or the time is obtained by mathematical differentiation. Figure 9.3(d) is an example. The advantage of expressing drying data in the form of rates is that their dependence on thermal and mass transfer driving forces is more simply correlated. Thus, the general drying equation may be written

$$\frac{1}{A}\frac{dW}{d\theta} = h(T_g - T) = k_p(P - P_g) = k_H(H - H_g), \qquad (9.13)$$

where subscript g refers to the gas phase and H is the moisture content, (kg/kg dry material), corresponding to a partial or vapor pressure P. Since many correlations of heat and mass transfer coefficients are known, the effects of many changes in operating conditions on drying rates may be ascertainable. Figures 9.3(g) and (h) are experimental evidence of the effect of humidity of the air and (i) of the effect of air velocity on drying rates.

Other factors, however, often complicate drying behavior. Although in some ranges of moisture contents the drying process may be simply evaporation off a surface, the surface may not dry uniformly and consequently the effective amount of surface may change as time goes on. Also, resistance to diffusion and capillary flow of moisture may develop for which phenomena no adequate correlations are known. Furthermore, shrinkage may occur on drying, particularly near the surface, which hinders further movement of moisture outwards. In other instances, agglomerates of particles may disintegrate on partial drying.

Some examples of drying data appear in Figure 9.3. Commonly recognized zones of drying behavior are represented in Figure 9.3(a). Equilibrium moisture contents assumed by various materials in contact with air of particular humidities is represented by (b). The shapes of drying rate curves vary widely with operating conditions and the physical state of the solid; (b) and others are some examples. No correlations have been developed or appear possible whereby such data can be predicted. In higher ranges of moisture content of some materials, the process of drying is essentially evaporation of moisture off the surface, and its rate remains constant until the surface moisture is depleted as long as the condition of the air remains the same. During this period, the rate is independent of the nature of the solid. The temperature of



Figure 9.3. (a) Classic drying curve of moisture content against time; a heat-up period in which no drying occurs also is usually present (*Proctor and Schwartz, Inc.; Schweitzer, p.* 4.144). (b) Equilibrium moisture content as a function of relative humidity; many other data are tabulated in Chemical Engineers Handbook (*McGraw-Hill, New York, 1984, 20.12*). (*These data are from National Academy of Science, copyright 1926.*) (c) Rate of drying as a function of % saturation at low (subscript 1) and high (subscript 2) drying rates: (A) glass spheres, 60 μ m, bed 51 mm deep; (B) silica flour, 23.5 μ m, 51 mm deep; (C) silica flour, 7.5 μ m, 51 mm bed; (D) silica flour, 2.5 μ , 65 mm deep (data of Newitt et al., Trans. Inst. Chem. Eng. 27, 1 (1949). (d) Moisture content, time and drying rates in the drying of a tray of sand with superheated steam; surface 2.35 sqft, weight 27.125 lb. The scatter in the rate data is due to the rough numerical differentiation (*Wenzel, PhD. thesis, University of Michigan, 1949*). (e) Temperature and drying rate in the drying of sand in a tray by blowing air across it. Dry bulb 76.1°C, wet bulb 36.0°C (*Ceaglske and Hougen, Trans. AIChE 33, 283 (1937).* (f) Drying rates of slabs of paper pulp of several thicknesses *lafter McCready and McCabe, Trans. AIChE 29, 131 (1933)].* (g) Drying of asbestos pulp with air of various humidities [*McCready and McCabe, Trans. AIChE 27, 118 (1932)].* (i) Effect of air velocity on drying of clay slabs. The data are represented by $R = 2.0u^{0.74}(H_w - H)$. The dashed line is for evaporation in a wetted wall tower (*Walker, Lewis, McAdams, and Gilliland, Principles of Chemical Engineering, McGraw-Hill, New York, 1937*).













Drying Tie over Constant and Falling Rate Periods with Constant Gas Conditions

The data of Figure 9.3(d) were obtained on a sample that contained 27.125 lb dry sand and had an exposed drying surface of 2.35 sqft. Take the case of a sample that initially contained 0.168 lb moisture/lb dry material and is to be dried to W = 0.005 lb/lb. In these units, the constant rate shown on the graph is transformed to

$$-\frac{1}{2.35} \frac{dW}{de} = \frac{0.38}{21.125} \quad (lb/lb)/(hr)(sqft),$$

which applies down to the critical moisture content $W_c = 0.04$ lb/lb. The rate behavior over the whole moisture range is

$$-\frac{dW}{d\theta} = \begin{cases} 0.03292, & 0.04 < W < 0.168, \\ 0.823W, & W < 0.04. \end{cases}$$

Accordingly, the drying time is

$$\theta = \frac{W - W_c}{0.03292} + \frac{1}{0.823} \ln\left(\frac{W_c}{W}\right)$$
$$= \frac{0.168 - 0.04}{0.03292} + \frac{1}{0.823} \ln\left(\frac{0.04}{0.005}\right)$$
$$= 6.42 \text{ hr.}$$

This checks the reading off the plot of the original data on Figure 9.3(d).

the evaporate assumes the wet bulb temperature of the air. Constant rate zones are shown in (d) and (e), and (e) reports that temperatures are truly constant in such a zone.

The moisture content at which the drying rate begins to decline is called critical. Some of the variables on which the transition point depends are indicated in Figures 9.3(c) and (g). The shape of the falling rate curve sometimes may be approximated by a straight line, with equation

$$-\frac{dW}{d\theta} = k(W - W_e), \qquad (9.14)$$

where W_e is the equilibrium moisture content. When W_e is zero as it often is of nonporous granular materials, the straight line goes through the origin. (d) and (h) illustrate this kind of behavior. The drying time is found by integration of the rate plots or equations. The process is illustrated in Example 9.2 for straight line behavior. Other cases require numerical integration. Each of the examples of Figure 9.3 corresponds to a particular substantially constant gas condition. This is true of shallow bed drying without recirculation of humid gas, but in other kinds of drying equipment the variation of the rate with time and position in the equipment, as well as with the moisture content, must be taken into account.

An approximation that may be justifiable is that the critical moisture content is roughly independent of the drying conditions and that the falling rate curve is linear. Then the rate equations may be written

$$-\frac{1}{A}\frac{dW}{d\theta} = \begin{cases} k(H_s - H_g), & W_c < W < W_0, \\ \frac{k(H_s - H_g)(W - W_e)}{W_c - W_e}, & W_e < W < W_c. \end{cases}$$
(9.15)

Examples 9.3 and 9.4 apply these relations to a countercurrent dryer in which the humidity driving force and the equilibrium moisture content vary throughout the equipment.

LABORATORY AND PILOT PLANT TESTING

The techniques of measuring drying of stationary products, as on trays, are relatively straightforward. Details may be found in the references made with the data of Figure 9.3. Mass transfer resistances were eliminated by **Wenzel** through use of superheated steam as the drying medium.

In some practical kinds of dryers, the flow patterns of gas and solid are so complex that the kind of rate equation discussed in this section cannot be applied readily. The sizing of such equipment is essentially a scale-up of pilot plant tests in similar equipment. Some manufacturers make such test equipment available. The tests may establish the residence time and the terminal conditions of the gas and solid. Dusting behavior and possible need for recycling of gas or of dried material are among the other factors that may be noted.

Such pilot plant data are cited for the rotary dryer of Example 9.6. For the pneumatic conveying dryer of Example 9.8, the tests establish heat and mass transfer coefficients which can be used to calculate residence time under full scale operation.

Scale-up factors as small as 2 may be required in critical cases, but factors of 5 or more often are practicable, particularly when the tests are analyzed by experienced persons. The minimum dimensions of a test rotary dryer are 1 ft dia by 6 ft long. A common criterion is that the product of diameter and rpm be in the range 25-35. A laboratory pneumatic conveying dryer is described by Nonhebel and Moss (1971). The veseel is 8 cm dia by about 1.5 m long. Feed rate suggested is 100 g/min and the air velocity about 1 m/sec. They suggest that 6-12 passes of the solid through this equipment may be needed to obtain the requisite dryness because of limitations in its length.

The smallest pilot spray dryer supplied by **Bowen** Engineering Co. is 30 in. dia by 2.5-6.0 ft high. Atomization is with 15 SCFM of air at 100 psig. Air rate is 250 actual cfm at 150-1000°F. Evaporation rates of 15-80 lb/hr are attained, and particles of product range from 5 to 40 μ m.

A pilot continuous multitray dryer is available from the Wyssmont Co. It is 4 ft dia by 5 ft high with 9 trays and can handle 25-200 **lb/hr** of feed.

Batch **fluidized** bed dryers are made in quite small sizes, of the order of 100 lb/hr of feed as the data of Table 9.14(a) show, and are suitable for pilot plant work.

9.3. CLASSIFICATION AND GENERAL CHARACTERISTICS OF DRYERS

Removal of water from solids is most often accomplished by contacting them with air of low humidity and elevated temperature. Less common, although locally important, drying processes apply heat radiatively or dielectrically; in these operations as in freeze drying, the role of any gas supply is that of entrainer of the humidity.

The nature, size, and shape of the solids, the scale of the operation, the method of transporting the stock and contacting it with gas, the heating mode, etc. are some of the many factors that

Drying with Changing Humidity of Air in a Tunnel Dryer A granular material deposited on trays or a belt is moved through a tunnel dryer countercurrently to air that is maintained at 170°F with steam-heated tubes. The stock enters at 1400 lb dry/hr with W = 1.16 lb/lb and leaves with 0.1 lb/lb. The air enters at 5% relative humidity ($H_g = 0.0125$ lb/lb) and leaves at 60% relative humidity at 170°F ($H_g = 0.203$ lb/lb). The air rate found by moisture balance is 7790 lb dry/hr:



Drying tests reported by Walker, Lewis, McAdams, and Gilliland, *Principles of Chemical Engineering,* McGraw-Hill, New York, (1937, p. 671) may be represented by the rate equation

$$-100 \frac{dW}{d\theta} = \begin{cases} 0.28 \text{ (lb/lb)/hr}, & 0.58 < W < 1.16, \\ 0.28(W - W_e)/(0.58 - W_e), & W_e < W < 0.58. \end{cases}$$
(1)

The air was at 95°F and 7% relative humidity, corresponding to a humidity driving force of $H_s - H_g = 0.0082$. Equilibrium moisture content as a function of the fraction relative humidity (RH), and assumed independent of temperature, is represented by

$$W_{\rho} = 0.0036 + 0.1539(\text{RH}) - 0.097(\text{RH})^2.$$
 (2)

The critical moisture content is assumed indpendent of the drying rate. Accordingly, under the proposed operating conditions, the rate of drying will be

$$-100 \frac{dW}{d\theta} = \begin{cases} \frac{0.28(H_s - H_g)}{0.0082}, & 0.58 < W < 1.16, \\ \frac{0.28(H_s - H_g)(W - W_e)}{0.0082(0.58 - 0.014)}, & W_e < W < 0.58. \end{cases}$$
(3)

With moisture content of the stock as a parameter, the humidity of the air is calculated by moisture balance from

$$H_g = 0.0125 + (1400/7790)(W = 0.1).$$
⁽⁴⁾

The corresponding relative humidities and wet bulb temperatures and corresponding humidities H_s are read off a **psychrometric** chart. The equilibrium moisture is found from the relative humidity by Eq. (2). The various corrections to the rate are applied in Eq. (3). The results are tabulated, and the time is found by integration of the rate data over the range $0.1 \le W \le 1.16$.

W	Hg	H,	RH	W,	Rate	1 / Rate
1.16	0. 203	0. 210			0. 239	4. 184
1.00	0.174	0.182			0.273	3.663
0.9	0.156	0.165			0.303	3. 257
0.8	0.138	0.148			0.341	2.933
0.7	0.120	0.130			0.341	2.933
0.58	0.099	0.110	0.335	0.044	0.356	2.809
0.50	0.094	0.096	0.29	0.040	0.333	3.003
0.4	0.066	0.080	0.24	0.035	0.308	3. 247
0.3	0.049	0.061	0.18	0.028	0.213	4.695
0.2	0.030	0.045	0.119	0.021	0.162	6.173
0.1	0.0125	0.0315	0.050	0.011	0.102	9.804

The drying time is

$$\theta = \int_{1.16}^{0.10} \frac{dw}{\text{rate}} = 4.21 \text{ hr}, \text{ by trapezoidal rule.}$$

The length of tunnel needed depends on the space needed to ensure proper circulation of air through the granular bed. If the bed moves through the dryer at 10 ft/hr, the length of the dryer must be at least 42 ft.



have led to the development of a considerable variety of equipment. The most elaborate classification of dryers is that of **Kröll** (1978) which assigns one of 10 letters for the kind of solid and one of seven numbers for the kind of operation. As modified by Keey (1972), it comprises 39 main classes and a total of 70 with subclasses. Less comprehensive but perhaps more practical classifications are shown in Table 9.1. They take into account the method of operation, the physical form of the stock, special features, scale of production, and drying time.

In a later section, the characteristics and performances of the most widely used equipment will be described in some detail. Many types are shown in Figure 9.4. Here some comparisons are made. Evaporation rates and thermal efficiencies are compared in Table 9.2, while similar and other data appear in Table 9.3. The wide spreads of these numbers reflect the diversity of individual designs of the same general kind of equipment, differences in moisture contents, and differences in drying properties of various materials. Fluidized bed dryers, for example, are operated as batch or continuous, for pharmaceuticals or asphalt, at rates of hundreds or many thousands of pounds per hour.

An important characteristic of a dryer is the residence time distribution of solids in it. Dryers in which the particles do not move relatively to each other provide uniform time distribution. In spray, pneumatic conveying, fluidized bed, and other equipment in which the particles tumble about, a substantial variation in residence time develops. Accordingly, some particles may **overdry** and some remain wet. Figure 9.5 shows some data. Spray and pneumatic conveyors have wide time distributions; rotary and **fluidized** bed units have narrower but far from uniform ones. Differences in particle size also lead to nonuniform drying. In pneumatic conveying dryers particularly, it is common practice to recycle a portion of the product continuously to ensure adequate overall drying. In other cases recycling may be performed to improve the handling characteristics when the feed material is very wet.

EXAMPLE 9.4 Effects of Moist Air

Effects of Moist Air Recycle and Increase of Fresh Air Rate in Belt Conveyor Drying

The conditions of Example 9.3 are taken except that recycle of moist air is employed and the equilibrium moisture content is assumed constant at $W_e = 0.014$. The material balance in terms of the recycle ratio R appears on the sketch:



A = air, W = water, S = dry solid

Humidity of the air at any point is obtained from the water balance

$$H_g = \frac{1581.4R + 97.4 + 1400(W - 0.1)}{7790(R + 1)} \tag{1}$$

The vapor pressure is

$$p_s = \exp[11.9176 - 7173.9/(T_s + 389.5)]$$
 atm. (2)

The saturation humidity is

$$H_s = (18/29)p_s/(1-p_s). \tag{3}$$

The heat capacity is

$$C = 0.24 + 0.45H_g.$$
 (4)

With constant air temperature of 170°F, the equation of the adiabatic saturation line is

$$170 - T_s = \frac{\lambda}{C} (H_s - H_g) \simeq \frac{900}{C} (H_s - H_g).$$
⁽⁵⁾

The drying rate equations above and below the critical moisture content of 0.58 are

R	×	0

W	T _s	1 / Rate
1.16 1.00 .90 .70 .50 .40 .20 .10	150.21 145.92 142.86 139.45 135.62 131.24 126.19 120.25 113.08 104.15 92.45	3.9627 3.4018 3.1043 2.8365 2.5918 2.3680 2.5187 2.8795 3.5079 4.8223 9.2092

$$-100\frac{dW}{d\theta}$$

$$=\begin{cases} 34.15(R+1)^{0.8}(H_s-H_g), & 0.58 < W < 1.16, (6) \\ 60.33(R+1)^{0.8}(H_s-H_g)(W-0.014), & W < 0.58. \end{cases}$$
(7)

When fresh air supply is simply increased by a factor $\mathbf{R} + 1$ and no recycle is employed, Eq. (1) is replaced by

$$H_g = \frac{97.4(R+1) + 1400(w-0.1)}{7790(R+1)}$$
(8)

The solution procedure is:

- 1. Specify the recycle ratio R (Ibs recycle/lb fresh air, dry air basis).
- 2. Take a number of discrete values of W between 1.16 and 0.1. For each of these fmd the saturation temperature T_s and the drying rates by the following steps.
- 3. Assume a value of T_s .
- 4. Find H_g , P_s , H_s , and C from Eqs. (1)-(4).
- 5. Find the value of T_s from Eq. (5) and compare with the assumed value. Apply the Newton-Raphson method with numerical derivatives to ultimately find the correct value of T_s and the corresponding value of H_r .
- 6. Find the rate of drying from Eqs. (6), (7).
- 7. Find the drying time by integration of the reciprocal rate as in Example 9.3, with the trapezoidal rule.

The printout shows saturation temperatures and reciprocal rates for R = 0, 1, and 5 with recycle; and for R = 1 with only the fresh air rate increased, using Eq. (8). The residence times for the four cases are

$$R = 0$$
, moist air, $\theta = 3.667$ hrs
= 1, moist air, = 2.841
= 5, moist air, = 1.442
= 1, fresh air, = 1.699.

Although recycling of moist air does reduce the drying time because of the increased linear velocity, an equivalent amount of fresh air is much more effective because of its lower humidity. The points in favor of moist air recycle, however, are saving in fuel when the fresh air is much colder than 170°F and possible avoidance of case hardening or other undesirable phenomena resulting from contact with very dry air.

R = 1, fresh air

w	T _s	1 / Rate
1.16 1.00 .90 .70 .50 .50 .40 .30 .20	132.62 128.81 126.19 123.35 120.25 116.85 116.12 113.08 108.89 104.15 98.74 92.45	1.3978 1.2989 1.2395 1.1815 1.1248 1.0693 1.0582 1.1839 1.4112 1.7979 2.6014 5.2893

EXAMPLE 9.4—(continued)

R	=	1.	moist	air
п	-	ч,	moist	an

w	Τ _s	1 / Rate		
1.16 1.00 .90 .80 .70 .50 .50 .40 .30 .20 .10	150 21 148 15 146 77 145.33 143 81 142 21 141 88 140 52 138 72 136 82 134.79 132 62	2.2760 2.1043 2.0088 1.9181 1.8323 1.7509 1.7351 1.9534 2.3526 3.0385 4.4741 9.3083		

- 10 ! Example 9.4. Belt conveyor dryin4
- 20 R=1! change for other cases
- 30 INPUT W
- 40 H1=(1581.4*(R+1)+97.4+1400*(W-.1))/7790/(R+1) 45 ! H1=(97.4*(R+1)+1400*(W-.1)
- //7790/(R+1)! Replace l i n e
 40 with this when no recycle
 is used.
 50 C=.24+.45*H1
 60 T=120! Trial sat temp
 70 GDSUB 200
- 30 Y1=Y
- 90 T=1.0001*T
- 100 GDSUB 200
- 110 Y2=Y
- 120 K=.0001*T*Y1/(Y2-Y1)
- 1 3 0 T=T/1.0001-K ! Newton-Raphso n

PRODUCTS

More than one kind of dryer may be applicable to a particular product, or the shape and size may be altered to facilitate handling in a preferred kind of machine. Thus, application of **through**circulation drying on tray or belt conveyors may require prior extrusion, pelleting, or briquetting. Equipment manufacturers know the capabilities of their equipment, but they are not always reliable guides to comparison with competitive kinds since they tend to favor what they know best. Industry practices occasionally change over a period of time. For example, at one time rotary kilns were used to dry and prepare fertilizer granules of a desired size range by accretion from concentrated solutions onto the mass of drying particles. Now this operation is performed almost exclusively in fluidized bed units because of economy and controlability of dust problems.

Typical examples of products that have been handled successfully in particular kinds of dryers are listed in Table 9.4. The performance data of later tables list other examples.

 $\mathbf{R} = 5$, moist air

W	T _s	1 / Rate
1 16 1 00 90 80 70 60 58 50 40 30 20	150.21 149.54 149.11 148.68 148.24 147.79 147.70 147.33 146.87 146.40 145.92 145.43	.9451 .9208 .9060 .8916 .8776 .8630 .8604 .9918 1.2302 1.6364 2.4824 5.3205

140 IF ABS(K/T) <= .00001 THEN 160 150 GOTO 7 0 160 PRINT USING 170 ; W, T, 1/R1 ! SubstR 2 for R1 when W<.58 170 IMAGE D.DD,2X,DDD.DD,2X,D.DD nn 175 GOTO 3 0 150 END ! SR for sat temp P=EXP(11.9176-7173.9/(T+389. 200 210 5)) H=.621*P/(1-P) ! sat humidit 220 230 Y=170-T-900*(H-H1)/C 240 R1=34.15*(R+1)^.8*(H-H1) 250 R2=60.33*(R+1)^.8*(H-H1)*(W-014> 260 RETURN 270 END

COSTS

Differences in thermal economies are stated in the comparisons of Table 9.2 and other tables. Some equipment cost data are in Chapter 20. When the capacity is large enough, continuous dryers are less expensive than batch units. Those operating at atmospheric pressure cost about 1/3 as much as those at vacuum. Once-through air dryers are one-half as expensive as recirculating gas equipment. Dielectric and freeze driers are the most expensive and are justifiable only for sensitive and specialty products. In the range of 1-50 Mtons/yr, rotary, fluidized bed and pneumatic conveying dryers cost about the same, although there are few instances where they are equally applicable.

SPECIFICATION FORMS

A listing of key information relating to dryer selection and design is in Table 9.5. Questionnaires of manufacturers of several kinds of dryers are in Appendix C.

9.4. BATCH DRYERS

Materials that require more than a few minutes drying time or are in small quantity are treated on a batch basis. If it is granular, the material is loaded on trays to a depth of 1-2in. with spaces of approximately 3 in. between them. Perforated metal bottoms allow drying from both sides with improved heat transfer. Hot air is blown across or through the trays. Cross velocities of 1000 ft/min are feasible if dusting is not a problem. Since the rate of evaporation increases roughly with the 0.8 power of the linear velocity, high velocities are desirable and are usually achieved by internal recirculation with fans. In order to maintain humidity at operable levels, venting and fresh air makeup are provided at rates of **5–50%** of the internal circulation rate. Rates of evaporation of 0.05-0.4 lb/(hr)(sqft tray area) and steam requirements of 1.5-2.3 lbs/lb evaporation are realized.

Drying under vacuum is commonly practiced for sensitive materials. Figure 9.6 shows cross and through circulation tray





(b)

Classification of dryers based on physical $form \mbox{ of feed}$

wet feed

<u>Evar</u> Liquid	porate Evap or bac Pumpable slurry or suspension	orate ck-mix Soft paste or sludge	Press Preformed paste	Gri Hard paste or matrix	nd Gri Of pro Free-flowing granular or crystalline solid	nd eform Fibrous solid	Sheet
Liquid	Slurry	Paste	Reform	Hard	Granular	Fibrous	Sheet
Agitated batch Drum Spray	Agitated batch Vacuum band Drum Spray	Vacuum tray Agitated batch Convection tray Fluid bed Vacuum band Drum Spray Pneumatic Convection band Cont. tray	Vacuum tray Convection tray Batch through- circulation Fluid bed Pneumatic Convection band Cont. tray Cont. through- circulation	Vacuum tray Convection tray Indirect rotary Cont. tray	Vacuum tray Agitated batch Convection tray Batch through- circulation Fluid bed Indirect rotary Pneumatic Direct rotary cont. tray Cont. through- circulation	Vacuum tray Convection tray Batch through- circulation Fluid bed Indirect rotary Pneumatic Direct rotary Convection band Cont tray Cont. through- circulation	Vacuum tray Convection tray Drum Cont. tray

TABLE 9.1-(continued)





Classification of dryers by suitability for special features



arrangements. The typical operating data of Table 9.6 cover a wide range of drying times, from a fraction of an hour to many hours. Charging, unloading, and cleaning are labor-intensive and time-consuming, as much as 5-6 hr for a 200-tray dryer, with trays about 5 sqft and 1-1.5 in. deep, a size that is readily handled manually. They are used primarily for small productions of valuable and thermally sensitive materials. Performance data are in Tables 9.6(b) and (c).

Through circulation dryers employ perforated or open screen bottom tray construction and have baffles that force the air through the bed. Superficial velocities of 150 ft/min are usual, with pressure drops of 1 in. or so of water. If it is not naturally granular, the material may be preformed by extrusion, pelleting, or briquetting so that it can be dried in this way. Drying rates are greater than in cross flow. Rates of 0.2-2 lb/(hr)(sqft tray area) and thermal efficiencies of 50% are realized. Table 9.7(d) has performance data.

Several types of devices that are used primarily for mixing of granular materials have been adapted to batch drying. Examples appear in Figure 9.8. They are suited to materials that do not stick to the walls and do not agglomerate during drying. They may be jacketed or provided with heating surfaces in the form of tubes or platecoils, and are readily arranged for operation under vacuum when handling sensitive materials. The double-cone tumbler has

been long established. Some operating data are shown in Table 9.7. It and V-shaped dryers have a gentle action that is kind to fragile materials, and are discharged more easily than stationary cylinders or agitated pans. The fill proportion is 50-70%. When heated with 2 atm steam and operating at 10 Torr or so, the evaporation rate is 0.8-1.0 lb/(hr)(sqft of heating surface).

Fixed cylinders with rotating ribbons or paddles for agitation and pans with vertical agitators are used to a limited extent in batch operation. Pans are used primarily for materials that become sticky during drying. Table 9.7 and Figure 9.7 are concerned with this kind of equipment.

A detailed example of capital and operating costs of a jacketed vacuum dryer for a paste on which they have laboratory drying data is worked out by Nonhebel and Moss (1971, p. 110).

Fluidized bed dryers are used in the batch mode on a small scale. Table 9.14(a) has some such performance data.

9.5. CONTINUOUS TRAY AND CONVEYOR BELT DRYERS

Trays of wet material loaded on trucks may be moved slowly through a drying tunnel: When a truck is dry, it is removed at one end of the tunnel, and a fresh one is introduced at the other end. Figure 9.8(c) represents such equipment. Fresh air inlets and humid

Equipment	Figure 9.4	(lb/hr)/sqft	(lb/hr)/cuft	Efficiency" (%)
Belt conveyor	e			46-58
Shelf				
Flow through	а	0.02-2.5		18-41
Flow past	а	0.02-3.1		18-41
Rotary				
Roto-louvre		7.2-15.4		23-66
Parallel current direct fired		6.1-16.4		65
Parallel current warm air	f	6.1-16.4		50
Countercurrent direct fired		6.1-16.4		60
Countercurrent warm air	f	6.1-16.4		4 5
Steam tube	h	6.1-16.4		8 5
Indirect fired	g	6.1-16.4		2 5
Tunnel				36-42
Pneumatic				
0.5 mm dia granules	0		6.2	26-63
1.0 mm			1.2	26-63
5mm			0.25	26-63
Spray	m		0.1-3	21-50
Fluidized bed	n		50-l 60	20-55
Drum	1	1.4-5.1		36-73
Spiral agitated				
High moisture	1	1-3.1		36-63
Low moisture	1	0.1-0.5		36-63
Splash paddle	k		5.6	65-70
Scraped multitray	d	0.8-1.6		

TABLE 9.2. Evaporation Rates and Thermal Efficiencies of Dryers

"Efficiency is the ratio of the heat of evaporation to the heat input to the dryer.

TABLE 9.3. Comparative Performances of Basic Dryer Types

	Basic Dryer Tvoe					
	Tray	Conveyor	Rotary	Spray	Flash	Fluid Bed
Product	filter cake	clay	sand	TiO₂	spent grain	coal
Drying time (min)	1320	9.5	12	<1.0	<1.0	2.0
Inlet gas temperature (°F)	300	420	1650	490	1200	1000
Initial moisture (% dry basis)	233	2 5	6	100	150	16
Final moisture (% drv basi̯s)	1	5.3	0.045	0	14	7.5
Product loading (Ib dry/ft ²)	3.25	16.60	N.A.	N.A.	N.A.	21 in. deep
Gas velocity (ft/min)	500	295	700	50	2000	1000
Product dispersion in gas	slab	packed bed	gravity flow	spray	dispersed	fluid bed
Characteristic product shape	thin slab	extrusion	granules	spherical drops	grains	¹ / ₂ -in. particles
Capacity []b evap./(h)(dryer area))	0.34	20.63	1.35	0.27'	10 "	285
Energy consumed (Btu/lb evap.)	3000	1700	2500	1300	1900	2000
Fan [hp/(lb evap./h)]	0.042	0.0049	0.0071	0.019	0.017	0.105

^alb evap./(h)(dryer, volume).

(Wentz and Thygeson, 1979: tray column from Perry, *Chemical Engineers' Handbook*, 4th ed., p. 20-7; conveyor and spray columns from Proctor and Schwartz, Division of SCM; rotary, flash, and fluid bed columns from Williams-Gardner, 1971, pp. 75, 149, 168, 193).

air outlets are spaced along the length of the tunnel to suit the rate of evaporation over the drying curve. This mode of operation is suited particularly to long drying times, from 20 to 96 hr for the materials of Table 9.6(e).

In the rotating tray assembly of Figure 9.8(a), material enters at the top and is scraped onto successive lower trays after complete revolutions. A leveler on each tray, shown in Figure 9.8(b), ensures uniform drying. Although the air flow is largely across the surface of the bed, the turnover of the material as it progresses downward makes the operation more nearly through-circulation. A cooling zone is readily incorporated in the equipment. The contacting process is complex enough that laboratory tray drying tests are of little value. A pilot plant size unit is cited in Section 9.2. Some industrial data on rotary tray drying are in Table 9.9, and some other substances that have been handled successfully in this equipment are listed in Table 9.4.

An alternate design has fixed jacketed trays for indirect heating. Scrapers attached to the central shaft drop the material from tray to tray. Like the rotating tray equipment, this equipment is limited to free flowing materials, but has the advantage of being essentially dust free.

Equipment developed essentially for movement of granular solids has been adapted to drying. Screw conveyors, for instance, have been used but are rarely competitive with belt conveyors,



(k)





(b)

Fan 🚺

Vacuum

chamber



Feed

Exhaust

Rotating

shell

Steam tube (hot gas)

rotary dryer

Feed

Heat

Exhaust

Dry products

Steam

Steam supply

and condensate removal

(f)

Steam

/ tubes

Product

(g)

Heaters

To vacuum

source

Trays of

material

Heated

shelves

(a)













Figure 9.4. Types of dryers cited in Tables 9.1 and 9.2. (a) Tray or compartment. (b) Vacuum tray. (c) Vertical agitated batch vacuum drier. (d) Continuous agitated tray vertical turbo. (e) Continuous through circulation. (f) Direct rotary. (g) Indirect rotary. (h) Agitated batch rotary (atmos or vacuum). (i) Horizontal agitated batch vacuum drier. (j) Tumble batch dryer. (k) Splash dryer. (I) Single drum. (m) Spray. (n) Fluidized bed dryer. (o) Pneumatic conveying *(mostly after Nonhebel and Moss, 1971)*.



Figure 9.5. Residence time distribution in particle dryers. (a) Four types of dryers (*McCormick*, 1979). (b) Residence time distribution of air in a detergent spray tower; example shows that 27% (difference between the ordinates) has a residence time between 24 and 32sec [Place et al., Trans. Inst. Chem. Eng. 37,268 (1959)]. (c) Fluidized bed drying of two materials (*Vanacek* et al., Fluidized Bed Drying, 1966).

particularly for materials that tend to degrade when they are moved. From the point of view of drying, belt conveyors are of two types: with solid belts and air flow across the top of the bed, called convection drying, or with perforated belts and through circulation of the air. The screw conveyor of Figure 9.8(f) has indirect heating.

Solid belts are used for pastes and fine powders. Through

TABLE 9.4. Examples of Products Dried in Specific Kinds of Equipment

- Spray dryers: rubber chemicals, sulfonates, inorganic phosphates, ceramics, kaolin, coffee, detergents, pharmaceuticals, pigments, inks, lignosulfonate wood waste, melamine and urea formaldehyde resins, polyvinyl chloride, microspheres, skim milk, eggs, starch, yeast, silica gel, urea, salts
- 2. Drum dryers: potatoes, cereals, buttermilk, skim milk, dextrins, yeasts, instant oat meal, polyacylemides, sodium benzoate, propionates, acetates, phosphates, chelates, aluminum oxide, m-disulfuric acid, barium sulfate, calcium acetate-arsenate-carbonate-hydrate-phosphate, caustic, ferrous sulfate, glue, lead arsenate. sodium benzene sulfonate. and sodium chloride
- 3. Vacuum drum dryers: syrups, malted milk, skim milk, coffee, malt extract, end glue
- 4. Vacuum rotary dryers: plastics, organic polymers, nylon chips, chemicals of all kinds, plastic fillers, plasticizers, organic thickeners, cellulose acetate, starch, and sulfur flakes
- 4. Belt conveyor dryers: yeast, charcoal briquettes, synthetic rubber, catalysts, soap, glue, silica gel, titanium dioxide, urea formaldehyde, clays, white lead, chrome yellow, and metallic stearates
- Pneumatic conveyor dryers: yeast filter cake, starch, whey, sewage sludge, gypsum, fruit pulp, copper sulfate, clay, chrome green, synthetic casein, and potassium sulfate
- Rotary multitray dryer: pulverized coal, pectin, penicillin, zinc sulfide, waste slude, pyrophoric zinc powder, zinc oxide pellets, calcium carbonate, boric acid, fragile cereal products, calcium chloride flakes, caffein, inorganic fluorides, crystals melting near 100°F, prilled pitch, electronic grade phosphors, and solvent-wet organic solids
- Fluidized bed dryer: lactose base granules, pharmaceutical crystals, weed killer, coal, sand, limestone, iron ore, polyvinyl chloride, asphalt, clay granules, granular desiccant, abrasive grit, and salt
- Freeze dryers: meat, seafood, vegetables, fruits, coffee, concentrated beverages, pharmaceuticals, veterinary medicines,
- and blood plasma
 Dielectric drying: baked goods, breakfast cereals, furniture timber blanks, veneers, plyboard, plasterboard, water-based foam plastic slabs, and some textile products
- Infrared drying: sheets of textiles, paper and films, surface finishes of paints and enamels, and surface drying of bulky nonporous articles.

circulation belts are applied to granules more than about 3 mm in narrowest dimension. When the feed is not in suitable granular form, it is converted in a preformer to a size range usually of 3-15 mm. Belts are made of chain mail mesh or metal with 2 mm perforations or slots of this width.

Several arrangements of belt dryers are shown *in* Figures 9.8(c)-(e). In the wet zone, air flow usually is upward, whereas in the drier and cooling zones it is downward in order to minimize dusting. The depth of material on the belt is 1-8 in. Superficial air velocities of 5 ft/sec usually are allowable. The multizone arrangement of Figure 9.8(e) takes advantage of the fact that the material becomes lighter and stronger and hence can be loaded more deeply as it dries. Each zone also can be controlled separately for air flow and temperature. The performance data of Table 9.9 cover a range of drying times from 11 to 200 min, and thermal efficiencies are about 50%.

Laboratory drying rate data of materials on trays are best obtained with constant air conditions. Along a belt conveyor or in a tray-truck tunnel, the moisture contents of air and stock change with position. Example 9.3 shows how constant condition drying tests can be adapted to belt conveyor operation. The effects of recycling moist air and of increasing the air velocity beyond that studied in the laboratory tests are studied in Example 9.4. Recycling does reduce drying time because of the increased air velocity, but it

246 DRYERS AND COOLING TOWERS

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TABLE 9.5. Specification Form for a Dryer"

1.	Operation	m o d e	batch/continuous
		operating cycle	- h
2.	Feed	(a) material to be dried	
		(b) feed rate	—— kg/h
		(c) nature of feed	solution/slurry/sludge/granular/
			fibrous/sheet/bulky
		(d) physical properties of solids:	
		initial moisture content	– kg/kg
		hygroscopic-moisture content	—— kg/kg
		heat capacity	─── kJ/kg°C
		bulk density, wet	—— kg/m°
		particle size	- m m
		(e) moisture to be removed:	
		chemical composition	
		boiling point at 1 bar	 °C
		heat of vaporization	MJ/kg
		heat capacity	— kJ/kg°C
		(f) feed material is	scaling/corrosive/toxic/abrasive/
		• •	explosive
		(q) source of feed	
3.	Product	(a) final moisture content	—— kg/kg
		(b) equilibrium-moisture content at 60% r.h.	— kg/kg
		(c) bulky density	- kg/m ³
		(d) physical characteristics	granular/flaky/fibrous/powdery/
4.	Design restraints	(a) maximum temperature when wet	— °C
		when dry	<u> </u>
		(b) manner of degradation	
		(c) material-handling problems.	
		when wet	<u> </u>
		when dry	
		(d) will flue-gases contaminate product?	
		(A) snare limitations	
5.	Utilities	(a) steam available at	bar pressure (10 ⁶ N/m ²)
•••		maximum quantity	— ka/h
		costing	
		(h) other fuel	+/g
		at at	ka/h
		with heating value	— M.J/ka
		costing	
		(c) electric power	- V
		frequency	h z
		nhases	
		costing	
6.	Present method of drving	ooonig	4,
7.	Rate-of-drving data under co	onstant external conditions:	
			<u> </u>
	<i>or</i> data from existing plant		
	• • •		
			
8.	Recommended materials of	construction	
		(a) parts in contact with wet material	
		(b) parts in contact with vapors	

'Questionnaires of several manufacturers are in Appendix C. (Keev, 1972, p. 325).

is not as effective in this regard as the same increase in the amount of fresh air. Recycling is practiced, however, to reduce heat consumption when the fresh air is cold and to minimize possible undesirable effects from over-rapid drying with low humidity air. Parallel current operation also avoids overrapid drying near the end. For parallel flow, the moisture balance of Example 9.4 becomes

$$H_g = \frac{97.4(R+1) + 1400(1.16 - W)}{7790(R+1)}$$
(9.16)

and replaces line 30 of the computer program.





Figure 9.6. Tray dryer arrangements, batch and continuous. Performance data are in Table 9.5. (a) Air flow across the surfaces of the trays. (b) Air circulation forced through the beds on the trays (Proctor *and Schwartz Inc.*). (c) Continuous drying of trays mounted on trucks that move through the tunnel; air Row may be in parallel or countercurrent (*P.W. Kilpatrick, E. Lowe, and W.B. Van Arsdel*, Advances in Food Research, *Academic, New York, 1955, Vol. VI, p. 342*).

The kind of data desirable in the design of through-circulation drying are presented for a particular case by Nonhebel and Moss (1971, p. 147). They report on effects of extrusion diameters of the original paste, the bed depth, air linear velocity, and air inlet humidity, and apply these data to a design problem.

9.6. ROTARY CYLINDRICAL DRYERS

Rotating cylindrical dryers are suited for free-flowing granular materials that require drying times of the order of 1 hr or less. Materials that tend to agglomerate because of wetness may be preconditioned by mixing with recycled dry product.

Such equipment consists of a cylindrical shell into which the wet material is charged at one end and dry material leaves at the other end. Figure 9.9 shows some examples. Drying is accomplished by contact with hot gases in parallel or countercurrent flow or with heat transfer through heated tubes or double shells. Designs are available in which the tubes rotate with the shell or are fixed in space.

Diameters typically are 4-10 ft and lengths are 4-15 diameters. The product of **rpm** and diameter is typically between 25 and 35. Superficial gas velocities are 5-10 ft/sec; but lower values may be needed for fine products, and rates up to 35 ft/sec may be allowable for coarse materials. To promote longitudinal travel of the solid, the shell is mounted on a slope of 1 in 40 or 20.

In a countercurrent dryer the exit temperature of the solid approaches that of the inlet gas. In a parallel current dryer, the exit gas is $10-20^{\circ}C$ above that of the solid. For design purposes the temperature of the exit solid in parallel flow may be taken as $100^{\circ}C$.

Flights attached to the shell lift up the material and shower it as a curtain through which the gas flows. Cross sections of some dryers are shown in Figure 9.10. The shape of flights is a compromise between effectiveness and ease of cleaning. The number is between 2 and 4 times the diameter of the shell in feet, and their depth is between $\frac{1}{12}$ and $\frac{1}{8}$ of the diameter. Holdup in the dryer depends on details of design and operation, but 7-8% is a usual figure. Cross-sectional holdup is larger at the wet end than at the dry end. An 85% free cross section commonly is adopted for design purposes; the rest is taken up by flights and settled and cascading solids.

Residence time depends on the nature of the material and mechanical features of the dryer. The performance data of Table 9.10 show a range of **7–90 min**. A formula cited by **Williams**–Gardner (1971, p. 133) for the geometrical residence time is

$$\boldsymbol{\theta} = \boldsymbol{k} \boldsymbol{L} / \boldsymbol{n} \boldsymbol{D} \boldsymbol{S}, \tag{9.17}$$

where L is the length, D is the diameter, n is rpm, and S is the slope (in./ft). The coefficient k varies from 3 to 12 for various countercurrent single shell dryers. The formula may be of some value in predicting roughly the effects of changes in the quantities included in it.

The only safe way of designing a rotary dryer is based on pilot plant tests or by comparison with known performance of similar operations. Example 9.5 utilizes pilot plant data for upscaling a dryer. The design of Example 9.6 also is based on residence time and terminal conditions of solid and air established in a pilot plant.

When heating by direct contact with hot gases is not feasible because of contamination or excessive dusting, dryers with jacketed shells or other kinds of heat transfer surfaces are employed. Only enough air to entrain away the moisture is employed. The temperature of the solid approaches the boiling temperature of the water in the constant rate period. Figure 9.10 shows designs in which the heating tubes are fixed in space or are attached to the rotating shell. Table 9.10 gives some performance data.

Combined indirect and indirect dryers pass the hot gases first through a jacket or tubes, and then wholly or in part through the open dryer. Efficiencies of such units are higher than of direct units, being in the range 60-80%. Table 9.10(d) shows performance data. Since the surfaces are hot, this equipment is not suitable for

TABLE 9.6. Performance Data of Batch Tray and Tray-Truck Dryers

(a) Cross-Flow Operation

	Coated Tablets	PTFE	Aspirin Base Granules	Stearates	Chalk	Filter Cake	Filter Cake	Filter Cake
Capacity, wet charge (Ib)	120	80	56	20,000	1800	3000	2800	4300
Number of trays	4 0	2 0	2 0	320	72	80	80	80
Tray area (ft ²)	140	70	70	4800	1130	280	280	280
Depth of loading (in.)	0.5	1.0	0.5	2.0	2.0	1.0	1.0	1.5
Initial moisture (% w/w basis)	2 5	25-30	15	71	46	70	70	80
Final moisture (% w/w basis)	nil	0.4	0.5	0.5	2.0	1.0	1.0	0.25
Maximum air temperature (°F)	113	284	122	200	180	300	200	200
Loading (Ib/ft ²)	0.9	1.2	0.4	0.9	0.91	3.25	3.04	11.7
Drying time (hr)	12	5.5	14	24	4.5	2 2	4 5	12
Overall drying rate (Ib/hr)	2.6	5.3	0.84	62.5	185	96.6	43.2	S O
Evaporative rate (lb/hr/ft ²)	0.0186	0.05	0.008	0.013	0.327	0.341	0.184	0.317
Total installed HP	1	1	1	4 5	6	4	2	2

(Williams-Gardner, 1971, p. 75, Table 12: first three columns courtesy Calmic Engineering Co.; last five columns courtesy A.P.V.-Mitchell (Dryers) Ltd.)

(b) Vacuum Dryers with Steam Heated Shelves

	Soluble Aspirin	Paint Pigment	Ferrous Glutinate	Ferrous Succinate	Lithium Hydroxide	Tungsten Alloy	Stabilized Diazamin
Capacity, wet_product (Ib/h)	4 4	30.5	41.6	52.5	36.8	12.8	4.6
Tray area (ft ²)	108	108	108	108	54	215	172
Depth of loading (in.)		2	0.5	1	1	0.5	0.75
Initial moisture (% w/w basis)	72.4	49.3	2 5	37.4	59	1.6	22.2
Final moisture (% w/w basis)	1.25	0.75	0.5	18.8	0.9	nil	0.5
Max temp (°F)	104	158	203	203	122	239	95
Loading [lb charge (wet) ft ²]	6.1	102	2.3	1.94	3.08	7.16	1.22
Drying time (hr)	15	36	6	4	4.5	12	4.8
Overall drying rate (lb moisture evaporated/ft ² /hr)	0.293	0.14	0.11	0.11	0.034	0.013	0.0058
Total installed HP	6	6	6	6	3	2	5
Vacuum (in. Hg)	29.5	28	27	27	27	29	22-23

(Williams-Gardner, 1971, p. 88, Table 15: courtesy Calmic Engineering Co.)

(c) Vacuum Dryers with Steam-Heated Shelves

Material	Sulfur Black	Calcium Carbonate	Calcium Phosphate	
Loading (kg dry material/m')	2 5	17	3 3	
Steam press(weared gauge)	410	410	205	
Vacuum (mm Hg)	685-710	685-710	685-7 10	
Initial moisture content (%, wet basis)	50	50.3	30.6	
Final moisture content (%, wet basis)	1	1.15	4.3	
Drying time (hr)	8	7	6	
Evaooration rates (kg/sec m ²)	8.9×10^{-4}	7.9×10^{-4}	6.6×10^{-4}	

(Chemical Engineers' Handbook, McGraw-Hill, New York, 1984, p. 20.23, Table 20.8).

(d) Through Circulation Dryers

Kind of Material	Granular Polymer	Vegetable	Vegetable Seeds	
Capacity (kg product/hr)	122	42.5	27.7	
Number of trays	16	24	24	
Tray spacing (cm)	4 3	4 3	4 3	
Tray size (cm)	91.4x 104	91.4x 104	85 × 98	
Depth of loading (cm)	7.0	6	4	
Physical form of product	crumbs	0.6-cm diced	washed	
		cubes	seeds	
Initial moisture content (%, dry basis)	11.1	669.0	100.0	
Final moisture content (%, dry basis)	0.1	5.0	9.9	
Air temperature (°C)	88	77 dry-bulb	36	
Air velocity, superficial (m/sec)	1.0	0.6-1 .0	1.0	
Tray loading (kg product/m ²)	16.1	5.2	6.7	
Drying time (hr)	2.0	8.5	5.5	
Overall drying rate (kg water evaporated/hr m ²)	0.89	11.86	1.14	
Steam consumption (kg/kg water evaporated)	4.0	2.42	6.8	
Installedpower (kW)	7.5	19	19	

TABLE 9.6-(continued)

(e) Tray and Tray-Truck Dryers

Material	Color	Chrome Yellow	Toluidine Red	Half-Finished Titone	Color
Type of dryer	Z-truck	16-tray dryer	16-tray	3-truck	Z-truck
Capacity (kg product/hr)	11.2	16.1	1.9	56.7	4.8
Number of trays	8 0	16	16	180	120
Tray spacing (cm)	10	10	10	7.5	9
Tray size (cm)	60 imes 75 imes 4	65 x 100 x 2.2	65 × 100 × 2	60x70x3.8	60 x 70 x 2.5
Depth of loading (cm)	2.5-5	3	3.5	3	
Initial moisture (%, bone-dry basis)	207	46	220	223	116
Final moisture (%, bone-dry basis)	4.5	0.25	0.1	2 5	0.5
Air temperature (°C)	85-74	100	50	95	99
Loading (kg product/m')	10.0	33.7	7.8	14.9	9.28
Drying time (hr)	33	21	41	2 0	96
Air velocity (m/sec)	1.0	2.3	2.3	3.0	2.5
Drying (kg water evaporated/hr m ²)	0.59	6 5	0.41	1.17	0.11
Steam consumption (kg/kg water evaporated)	2.5	3.0	_	2.75	
Total installed power (kW)	1.5	0.75	0.75	2.25	1.5

(Proctor and Schwartz Co.).

TABLE 9.7. Performance of Agitated Batch Dryers (See Fig. 9.7)

(a) Double-Cone Tumbler

	Tungsten Carbide	Polyester Resin	Penicillin	Hydroquinone	Prussian Blue Pigment
Volatile ingredient	naphtha	water	acetone	water	water
Physical nature of charge	heavy slurry	pellets	powder	powder	filtercake
Dryer dia (ft)	2	2	2	2	2
Dryer capacity (ft ³)	2.5	2.5	2.5	2.5	2.5
Method of heating	hot water	steam	hot water	hot water	steam
Heating medium temperature (°F)	180	240	140	150	225
Vacuum (mm Hg abs)	40-84	12-18	40	50-100	40-110
Initial volatile content (% w/w basis)	18.0	0.34	27.9	5.0	83
Final volatile content (% w/w basis)	nil	0.01	nil	0.25	4.8
Weight of charge (Ib)	640	130	55	61	142.5
Bulk density of charge (Ib/ft ³)	256	51.5	21.5	26.5	58.5
Dtvina time (min)	155	215	90	50	480

(Courtesy Patterson Division, Banner Industries Inc.; Williams-Gardner, 1971).

(b) Paddle, Ribbon, and Pan"

	Time of	Size of (m)	Dryer n)	Driving	Wet	Filling	Initial Moisture Content	Absolute Pressure	Jacket	Drying	Mean Overall Coeff.
Material	Dryer	Length	Dia	(HP)	(kg)	Ratio (%, Wet Basis) (mb) (°C) (hr)	(₩/m̃ ℃)				
Organic paste	HCRP	5500	1200		4000	0.36	30	200	8 0	15	3 5
Different fine	HCRP	3800	1350	15	2260	0.2	68	265	125	6	4 5
aromatic organic }	HCRP	3800	1350	15	4660	0.4	75	265	125	8	6 0
compound crystals	HCRP	5500	1200		2100	0.2	6	200	125	4	2 5
Anthracene (water											
and pyridine)	HCRP	8900	1800	35	37000	0.72	76	665-1000	170	16	75
Dyestuff paste	HCSB	2750	1200	10	2000	0.3	70	265	105	14	30
Different organic 🔪	PVP		1800	15	1080	0.4	41	1000	125	3 2	3 5
pastes	PVP		2450	25	800	0.4	35	665	125	7 <u>1</u>	2 5
Different dyestuff	PVP		1800	15	1035	0.4	61	1000	125	11	135
pastes	PVP		2450	20-30	2400	0.7	64	470	125	12	115

 $^{\it 0}$ HCRP = paddle agitator; HCSP = ribbon agitator; PVP = pan with vertical paddles. (Nonhebel and Moss, 1971).

250 dryers and cooling towers

TABLE 9.7—(continued)

(c) Pan Dryer

	Sodium Thiosulphate	Potassium Zeolite	Arsenic Pentoxide
Dryer diameter	6 ft 0 in.	2 ft 3 in.	8ftOin.
Dryer depth	2ftOin.	1 ft 0 in.	2ftOin.
Capacity (Ib product)	12cwt	14 lb	2 ¹ / ₂ ton/day
Initial moisture (% w/w basis)	37	4 0	3 5
Final moisture (% w/w basis)	0	1	2-3
Method of heating	steam	steam	steam
Atmospheric (a) or vacuum (b)	(b) 26 Hg	(a) 60 lb/in. ² /gauge	(b)
Drying temperature: material (°F)			
Drying temperature: shelf (°F)		153c	
Bulk density product (lb/ft ³)			
Drying time (hr/batch)	5	3	8
Material of construction	SS	MS	SS

[Courtesy A.P.V.-Mitchell (Dryers) Ltd., Williams-Gardner, 1971].





Figure 9.7. Tumbling and agitated heated dryers for atmospheric and vacuum batch operation. (a) Double cone tumbler, performance data in Table 9.6(a) (*Pennsalt Chem. Co.*). (b) V-shaped tumbler. (c) Ribbon agitated cylinder; performance data in Table 9.6(b). (A) jacketed shell; (B) heads; (C) charging connections; (D) discharge doors; (E) agitator shaft; (F) stuffing box; (G) shaft bearings; (H) agitator blades; (J) vapor outlets; (K) steam inlets; (L) condensate outlets; (M) discharge siphon for shaft condensate (*Buflovak Equip. Div., Blaw Knox* Co.) (d) Paddle agitated cylinder. Performance data in Table 9.6(b). (e) Horizontal pan with agitator blades. Data are Table 9.6(b).





Figure 9.7-(continued)



Figure 9.8. Rotary tray, through-circulation belt conveyor, and heated screw conveyor dryers. (a) Rotary tray dryer (*Wyssmont Co.*). (b) Action of a rotating tray and wiper assembly (*Wyssmont* Co.). (c) A single conveyor belt with air **upflow** in wet zone and **downflow in** dry (*Proctor and Schwartz Inc.*). (d) A two-stage straight-through belt conveyor dryer. (e) A three-belt conveyor dryer; as the material becomes dryer, the loading becomes deeper and the belt longer (*Proctor and Schwartz Inc.*). (f) Screw conveyor dryer with heated hollow screw (*Bepex* Corp.).





(f)

Figure 9.8—(continued)

TABLE 9.8. Performance of Rotary Tray and Pan Dryers

(a) Multitray Dryers at Atmospheric Pressure

	China Clay	Bread Crumbs	Cu-Ni Concentrate	Catalyst Pellets	Kaolin	Calcium Chloride	Urea	Vitamin Powder
Dryer height	_	_	_	23ft	23 ft	47ft	47ft	12 ft
Dryer diameter	~	_		19 ft	19 ft	31 ft	15ft	9 ft
Tray area (ft ²)	7000	2000 (drying) 1000 (cooling)	2900	-	-	-	_	-
Capacity (lb/product/hr)	31,000	1680	19,000	4200	10,000	24,000	5000	200
Initial moisture (% w/w basis)	30	36	2 2	4 5	3 5	2 5	2 0	2 0
Final moisture (% w/w ba	sis) 10	5	5	18	5	1	0.2	5
Product temperature (°F) 160	100	200	_	_	<u> </u>	_	
Residence time (min)	40	40 (drying) 20 (cooling)	2 5	-	-	-		-
Evaporation rate (Ib/ft ² /hr)	9.100	804	4060	2050	4600	11.000	100	37
Method of heating	external	steam	external	external	external	internal	external	external
Ũ	oil		oil	gas	oil	gas	steam	steam
Heat consumption (Btu/lb moisture	9			340		320		
evaporated)	1750		2200	1750	1850	1800	3500	2700
Installed HP	80	2 5	6 0	2 3	4 7	6 5	75	2 ¹ / ₂

Williams-Gardner, 1971). (First three columns courtesy Buell Ltd.; last five columns courtesy The Wyssmont Co., Inc.).

(b) Multiple Vacuum Pan Dryer

	Sodium Hydrosulphite	Maneb	Melamine	Activated Carbon	
Dryer diameter (pans) (m)	2	2	2	2	
Number of pans	5	17	11	17	
Area (approx)(m ²)	12.4	42.8	27.6	42.8	
Dry product (Ib/hr)	1100	660	1870	440	
Initial moisture (% w/w)	4	2 3	11	6 2	
Final moisture (% w/w)	0.1	0.5	0.03	3	
Heating	hot water	steam	steam	steam	
		1.3 atm	2.5 atm	2.5 atm	
Pan temperature (°C)	98	105	125	125	
Evaporation rate (Ib/ft ² /hr)	0.325	0.325	0.79	0.78	
Drying time (min)	15	170	12	30	

(Data of Krauss-Maffei-Imperial GmbH).

TABLE 9.9. Performance of Through-Circulation Belt Conveyor Dryers [See Figs. 9.8(c)-(e)]

(a) Data of A.P.V.-Mitchell (Dryers) Ltd.

	Fertilizers	Bentonite	Pigment	Nickel Hydroxide	Metallic Stearate
Effective dryer length	42 ft 6 in.	60ftOin.	24 ft 0 in.	24ftOin.	41 ft 3 in.
Effective band width	8 ft 6 in.	8 ft 6 in.	4ftOin.	4 ft 0 in.	6ftOin.
Capacity (Ib product/hr)	2290	8512	100	125	125
Method of feeding Feedstock preforming	oscillator	oscillator	extruder	extruder	extruder
Initial moisture (% w/w basis)	45.0	30	58.9	75	75
Final moisture (% w/w basis)	2.0	10.0	0.2	0.5	0.2
Drying time (min)	16	14	60	70	60
Drying rate (lb evaporated/ft ² /hr)	7.0	6.5	2.0	7.5	1.5
Air temperature range (°F)	-				
Superficial air velocity (ft/min)	200	200	180	180	125
Heat consumption (Btu/lb evaporated)	-				
Method of heating	direct oil	direct oil	steam	steam	steam
Fan installed HP	35	50	14	14	28

(Williams-Gardner, 1971).

(b) Data of Krauss-Maffei-Imperial GmbH

	Aluminium Hydrate	Polyacrylic Nitrile	Sulfur	Calcium Carbonate	Titanium Dioxide
Effective dryer length	32 ft 9 in.	43 ft 0 in.	28ftOin.	50ftOin.	108ftOin.
Effective band width	6 ft 6 in.	6 ft 6 in.	6 ft 6 in.	6 ft 3 in.	9 ft 6 in.
Capacity (lb product/hr)	615	2070	660	1800	6000
Method of feeding Feedstock preforming	<i>grooved</i> drum	extruder	extruder	extruder	extruder
Initial moisture (% w/w basis)	38.0	55.0	45.0	60.0	50.0
Final moisture (% w/w basis)	0.2	1.0	1.0	0.5	0.5
Drying time (min)	26	52	110	40	45
Drying rate (Ib evaporated/hr/ft	2.88	3.37	3.57	5.73	6.0
Air temperature range (°F)	233	186/130	194/230	320	314/392
Superficial air velocity (ft/min)	140	100/216	140	160	150
Heat consumption (Ib steam/Ib		•			
evaporated)	1.7-1.8	1.8-1.9	1.8-1.9	1.7-1.8	1.8-1.9
Method of heating	50 lb/in. ²	25 lb/in. ²	90 lb/in.2	160 lb/in ²	260 lb/in. ²
	steam	steam	steam	steam	steam
Fan installed hp (approx.)	25	65	20	35	80

(Williams-Gardner, 1971)

(c) Data of Proctor and Schwartz Inc.

Kind of Material	Inorganic Pigment	Cornstarch	Fiber Staple	Charcoal Briquettes	Gelatin	Inorganic Chemical
Capacity (kg dry product/hr)	712	4536	1724	5443	295	862
· · · · · · ·			Stage A Stage B			
Approximate dryer area (m ²)	22.11	66.42	57.04 35.12	52.02	104.05	30.19
Depth of loading (cm)	3	4		16	5	4
Air temperature (°C)	120	115-140	130-100 100	135-120	32-52	121-82
Loading (kg product/m ²)	18.8	27.3	3.5 3.3	182.0	9.1	33
Type of conveyor (mm)	1.59 by 6.35	1.19 by 4.76	2.57.diameter	8.5	4.23 X 4.23 mesh	1.59 × 6.35 slot
	slots	slots	holes, perforated	× 8.5 mesh	screen	
			plate	screen		
Preforming method or feed	rolling extruder	filtered and scored	fiber feed	pressed	extrusion	rolling extruder
Type and size of preformed	6.35-diameter	scored filter	cut fiber	64 X 51 X 25	2-diameter	6.35-diameter
particle (mm)	extrusions	cake			extrusions	extrusions
Initial moisture content (% bone-dry basis)	120	85.2	110	37.3	300	111.2
Final moisture content (% bone- dry basis)	0.5	13.6	9	5.3	11.1	1.0
Drying time (min)	35	24	11	105	192	70
Drying rate [kg water evaporated/(hr m ²)]	38.39	42.97	17.09	22.95	9.91	31.25
Air velocity (superficial)(m/sec)	1.27	1.12	0.66	1.12	1.27	1.27
Heat source per kg water	gas	steam	steam	waste heat	steam	gas
evaporated [steam kg/kg gas (m ³ /kg)]	0.11	2.0	1.73		2.83	0.13
Installed power (kW)	29.8	119.3	194.0	82.06	179.0	41.03

(Perrys Chemical Engineers Handbook, McGraw-Hill, New York, 1984).



Figure 9.9. Rotary dryer assemblies. (a) Parts of the shell of a direct fired rotary dryer (*C.E. Raymond Bartlett* Snow Co.). (b) Assembly of a rotary dryer with pneumatic recycle of fines (*Standard Steel Corp.*). (c) Steam tube dryer with mechanical conveyor for partial recycle of product for conditioning of the feed.

thermally sensitive materials and, of course, may generate dust if the gas rate through the open dryer is high.

In the Roto-Louvre design of Figure 9.10(b) the gas enters at the wall, flows first through the bed of particles and subsequently through the shower of particles. Performance data are in Tables 9.10(b) and (c).

A formula for the power required to rotate the shell is given by Wentz and Thygeson (1979):

$$P = 0.45W_{t}v_{r} + 0.12BDNf, \qquad (9.18)$$

where *P* is in watts, W_i is the weight (kg) of the rotating parts, v, is the peripheral speed of the carrying rollers (**m/sec**), *B* is the holdup of solids (kg), *D* is diameter of the shell (m), N is rpm, and *f* is the number of flights along the periphery of the shell. Information about weights may be obtained from manufacturers catalogs or may be estimated by the usual methods for sizing vessels. Fan and driver horsepower are stated for the examples of Tables 9.10(a)-(c). The data of Table 9.10(a) are represented roughly by

$$P = 5 + 0.11DL, (9.19)$$

where P is in HP and the diameter D and length L are in feet.

9.7. DRUM DRYERS FOR SOLUTIONS AND SLURRIES

Solutions, slurries and pastes may be spread as thin films and dried on steam heated rotating drums. Some of the usual arrangements are shown on Figure 9.11. Twin drums commonly rotate in opposite directions inward to nip the feed, but when lumps are present that could damage the drums, rotations are in the same direction. Top feed with an axial travelling distributor is most common. Dip feed is shown in Figure 9.11(d) where an agitator also is provided to keep solids in suspension. When undesirable boiling of the slurry in the



Figure 9.10. Cross sections of rotary dryers. (a) Action of the flights in cascading the drying material. The knockers are for dislodging material that tends to cling to the walls. (b) Cross section of chamber of rotolouvre dryer showing product depths and air flows at feed and discharge ends. The air enters at the wall and flows through the bed as well as through the cloud of showered particles (*Link-Belt* Co.). (c) Showering action in a dryer with fixed steam tubes and rotating shell. (d) Section and steam manifold at the end of a dryer in which the steam tubes rotate with the dryer.

pan could occur, splash feed as in Figure 9.11(c) is employed. Example 9.7 describes some aspects of an actual installation.

For mechanical reasons the largest drum made is 5 ft dia by 12 ft with 188 sqft of curved surface. A 2×2 ft drum also is listed in manufacturers' catalogs. Performance data are in Tables 9.11 and 9.12.

The material comes off as flakes 1-3 mm or less thick. They are broken up to standard size of about $\frac{1}{4}$ in. square. That process makes fines that are recycled to the dryer feed. Drying times fall in the range of 3-12 sec. Many laboratory investigations have been made of drying rates and heat transfer coefficients, but it appears that the only satisfactory basis for sizing plant equipment is pilot plant data obtained with a drum of a foot or more in diameter. Usually plant performance is superior to that of pilot plant units because of steadier long time operation.

Rotation speeds of the examples in Table 9.12 show a range of 1-24rpm. Thin liquids allow a high speed, thick pastes a low one. In Table 9.13(c) the evaporation rates group in the range 15-30 kg/m² hr, but a few of the data are far out of this range. The few data in Table 9.13(a) show that efficiencies are comparatively high, 1.3 lb steam/lb water evaporated.

A safe estimate of power requirement for double drum dryers is approx 0.67 HP/(rpm)(100 sqft of surface). Maintenance can be as high as 10%/yr of the installed cost. Knives last from 1 to 6 months depending on abrasiveness of the slurry. Competitors for drum dryers are solid belt conveyors that can can handle greater thicknesses of pasty materials, and primarily spray dryers that have largely taken over the field.

9.8. PNEUMATIC CONVEYING DRYERS

Free-flowing powders and granules may be dried while being conveyed in a high velocity air stream. The necessary equipment is variously called pneumatic conveying dryer, pneumatic dryer, air lift dryer, or flash dryer. The basic system consists of an air heater, solids feeding device, vertical or inclined drying leg, cyclone or other collector and an exhaust fan. Figure 9.12 shows some of the many commercial equipment. Provision for recycling some of the product generally is included. Some of the materials being handled successfully in pneumatic dryers are listed in Table 9.5.

Readily handled particles are in the size range l-3 mm. When the moisture is mostly on the surface, particles up to 10 mm have been processed. Large particles are brought down to size in dispersion devices such as knife, hammer or roller mills.

Typical performance data are summarized in Table 9.13. In practice air velocities are 10-30 m/sec. The minimum upward velocity should be 2.5-3 m/sec greater than the free fall velocity of the largest particles. Particles in the range of 1-2 mm correspond to an air velocity of 25 m/sec. Since agglomerates may exist under drying conditions, the safest design is that based on pilot plant tests or prior experience.

EXAMPLE 9.5 Scale-Up of a Rotary Dryer Tests on a laboratory unit come up with the stated conditions for drying a pelleted material at the rate of 1000 lb dry/hr:



The residence time is 20min. The speed is 3-4rpm. On the average, 7.5% of the cross section is occupied by solid. Because of dusting problems, the linear velocity of the air is limited to

12 ft/sec. The diameter and length will be found. Since the inlet and outlet conditions are specified and the moisture transfer is known, the heat balance can be made. The heat capacity of the solid is 0.24:

moisture evap =
$$1000(0.6-0.05) = 550$$
 lb/hr
air rate = $550/(0.0428 - 0.013) = 18.456$ lb/hr

Off a **psychrometric** chart, the sp vol of the air is 15.9 **cuft/(lb** dry). The diameter is

$$D = \left(\frac{18,456(15.9)}{3600(12)(1-0.075)\pi/4}\right)^{1/2} = 3.06 \text{ ft, say 3.0 ft}$$

The length is

$$L = \frac{30(20/60)}{0.075\pi D^2/4} = 18.9 \text{ ft}$$

EXAMPLE 9.6

Design Details of a Countercurrent Rotary Dryer Pilot plants indicate that a residence time of 3 hr is needed to accomplish a drying with the conditions indicated on the sketch. For reasons of entrainment, the air rate is limited to 750 lbs dry/(hr)(sqft cross section). Properties of the solid are 50 lb/ cuft and 0.22 Btu/(lb)(°F). Symbols on the sketch are A = dry air, S = dry solid, W = water:



In terms of the dry air rate, ${\bf A}~{\bf lb/hr},$ the average moist heat capacity is

$$C = 0.24 + 0.45[0.008 + \frac{1}{2}(333/A)] = 0.2436 + 74.93/A.$$

In the dryer, the enthalpy change of the moist air equals the sum of the enthalpy changes of the moisture and of the solid. Add 7% for heat losses. With steam table data,

$$\begin{array}{l} (0.2436 + 74.93/A)A(290 - 136) = 1.07[333(1120.3) + 1(228) \\ + 1000(0.22)(260 - 60) - 334(28)] \\ = 1.07(407,936) = 43,649], \\ \vdots \ \mathbf{A} = 11.633 \ \mathrm{lb/hr} \end{array}$$

The exit humidity is

H = 0.008 + 333/11,633 = 0.0366 lb/lb,

which corresponds to an exit dewpoint of 96°F, an acceptable value.

With the allowable air rate of 750 lb/hr sqft, the diameter of the dryer is

$$D = \sqrt{11,633/750\pi/4} = 4.44 \text{ ft}, \text{ say } 4.5 \text{ ft}.$$

Say the solid occupies 8% of the cross section. With a solids density of 50 lb/cuft, the dryer volume,

$$v = 3(1000/50)/0.08 = 750$$
 cuft,

and the length is

$$L = 750/(4.5)^2 \pi/4 = 47.2$$
 ft.

The standard number of flights is 2-4 times the diameter, or

number = (2-4)4.5 = 9-18, say 12.

The product of rpm and diameter is 25-35

$$rpm = (25-35)/4.5 = 5.5-7.8$$
, say 6.7.

The stm heater duty is

 $Q_s = 11,633(0.2436)(290 - 60) = 651,733$ Btu/hr, 150 psig stm, stm = 651,733/857 = 760.5 lb/hr.

Evaporation efficiency is

 $\eta = 333/760.5 = 0.438$ lb water/lb stm.

The efficiency of the dryer itself is

 $\eta_d = 407,936/651,733 = 0.626$ Btu/Btu.

(a) Direct Heated Dryers

	Sugar Beet Pulp ^a	Calcium Carbonate"	Blast Furnace Slag ^a	Lead Concentrate	Sand ^b	Zinc Concentrate	Ammonium Sulphate	Fine Salt"	Crystals ^d	$\mathbf{Chemicals}^d$
Air flow	parallel	parallel	parallel	parallel	parallel	parallel	counter	counter	counter	indirect counter
Dryer length	9 ft 2 in.	6 ft 3 in.	7 ft 2 in.	4 ft 6 in.	4 ft 6 in.	7 ft 6 in.	9 ft 0 in.	5 ft 0 in.	10 ft 0 in.	4 ft 6 in.
Dryer length	46ftOin.	34ftOin.	40 ft 0 in.	35ftOin.	32 ft 6 in.	60ftOin.	40 ft 0 in.	40ftOin.	60 ft 0 in.	27 ft 0 in.
Method of heating	oil	oil	oil	oil	gas	oil	gas	steam	steam	Louisville
										steam tube
Method of feed	screw	belt	belt	screw	chute	screw	conveyor	feeder	screw	screw
Initial moisture (% w/w)	a 2	13.5	33	14	5.65	18	2.5	5.0	7.0	1.5
Final moisture (火 w/w)	10	0.5	nil	а	0.043	а	0.2	0.1	a.99	0.1
Evaporation (Ib/hr)	34,000	6000	11,600	1393	701	8060	1120	400	1150	63
Capacity (Ib evaporated/f? dryer volume)	11	6	7	2.5	1.35	2.3	0.5	0.52	0.245	-
Efficiency (Btu supplied/water evaporated)	1420	1940	1710	2100	2550	1850	1920	2100	1650	-
Inlet air temperature (°F)	1560	1560	1560	1300	1650	1500	400	280	302	_
Outlet air temperature (°F)	230	220	248	200	222	200	180	170	144	
Residence time (av. min)	2 0	2 5	30	2 0	12	2 0	15	40	70	30/40
Fan HP	70	4 0	50	20	5	75	2 5	a	_	10
Motive HP	15	2 0	2 5	10	10	55	60	15	60	_
Fan capacity (std. air ft ³ /min)	45000	a500	ı a.000	2750	2100	12,000	18,500	6500	-	

257

^{*}Courtesy Buell Ltd. ^b Courtesy Head Wrightson (Stockton) Ltd. 'Courtesy Edgar Allen Aerex Ltd. ^d Courtesy Constantin Engineers Ltd.-Louisville Dryers; Williams-Gardner, 1971.

(b) Roto-Louvre Dryers

	Bone Meal	Sugar'	Sulfate of Ammonia"	Bread Crumbs	Bentonite
Dryer diameter	7 ft 6 in.	7 ft 6 in.	7 ft 6 in.	4 ft 6 in.	8 ft 10 in.
Dryer length	12 ft 0 in.	25ftOin.	25 ft 0 in.	20 ft 0 in.	30ftOin.
Initial moisture (% w/w basis)	17.0	1.5	1.0	37	4 5
Final moisture (% w/w basis)	7.0	0.03	0.2	2.5	11
Method of feed	screw	screw	chute	chute	chute
Evaporation rate (Ib/hr)	1660	500	400	920	7100
Efficiency (Btu supplied/lb evaporation)	74.3	4 0	-	55	62.5
Method of heating	steam	steam	steam	gas	oil
Inlet air temperature (°F)	203	194	246	572	642
Outlet air temperature (°F)	122	104	149	158	176
Residence time, min	9.3	12.5	9.0	25.7	37.3
Fan HP (absorbed)	49.3	52.2	55	13.7	54.3
Motive HP (absorbed)	а	12.5	15	2.3	20.0
Fan capacity (ft ³ /min)					
Inlet	9560	18,000	16,000	5380	20,000
Outlet	14,000	22,300	21,000	5100	25,000

TABLE 9.10—(continued)

(c) Roto-Louvre Dryers

Material Dried	Ammonium Sulfate	Foundry Sand	Metallurgical Coke	
Dryer diameter	2 ft 7 in.	6 ft 4 in.	10 ft 3 in.	
Dryer length	10 ft	24ft	30 ft	
Moisture in feed (% wet basis)	2.0	6.0	18.0	
Moisture in product (% wet basis)	0.1	0.5	0.5	
Production rate (Ib/hr)	2500	32,000	38,000	
Evaporation rate (Ib/hr)	50	2130	8110	
Type of fuel	steam	gas	oil	
Fuel consumption	255 lb/hr	4630 ft ³ /hr	115 gal/hr	
Calorific value of fuel	837 Btu/lb	1000 Btu/ft ³	150,000 Btu/gal	
Efficiency (Btu supplied per lb evaporation)	4370	2170	2135	
Total power reauired (HP)	4	41	78	

(FMC Corp.; Chemical Engineers' Handbook, 1984, p. 20.20).

(d) Indirect-Direct Double Shell Dryers

	Indire	ct-Direct Double	Shell	
	Coal	Anhydrite	Coke	
Dryer diameter Dryer length	7 ft 6 in. 46 ft 0 in.	5ft10in. 35 ft 0 in.	5 ft 10 in. 35ftOin.	
Initial moisture content (% w/w basis)	2 2	6.0	15	
Final moisture (% w/w basis)	6	1.0	1.0	
Evaporation rate (Ib/hr)	5800	2300	1600	
Evaporation-volume ratio (lb/ft ³ /hr)	3.5	3.15	2.2	
Heat source	coal	oil	oil	
Efficiency (Btu supplied/lb water evaporated)	1250	1250	1340	
Inlet air temperature (°F)	1200	1350	1350	
Outlet air temperature (°F)	160	160	200	

(Courtesy Edgar Allen Aerex Ltd.; Williams-Gardner, 1871).

(e) Steam Tube Dryers

	Class 1	Class 2	Class 3
Class of materials	high moisture organic, distillers' grains, brewers' grains, citrus pulp	pigment filter cakes, blanc fixe, barium carbonate, precipitated chalk	finely divided inorganic solids, water- ground mica, water-ground silica, flo- tation concentrates
Description of class	wet feed is granular and damp but not sticky or muddy and dries to granular meal	wet feed is pasty, muddy, or sloppy, product is mostly hard pellets	wet feed is crumbly and friable, product is powder with very few lumps
Normal moisture content of wet feed (% dry basis)	233	100	54
Normal moisture content of product (% dry basis)	11	0.15	0.5
Normal temperature of wet feed (K)	31 0-320	280-290	280-290
Normal temperature of product (K)	350-355	380-410	365-375
Evaporation per product (kg)	2	1	0.53
Heat load per Ib product (kJ)	2250	1190	625
Steam pressure normally used (kPa gauge)	860	860	860
Heating surface required per kg product (m ²)	0.34	0.4	0.072
Steam consumption per kg product (kg)	3.33	1.72	0.85

(Chemical Engineers' Handbook, **1984).**



Figure 9.11. Drum dryers for solutions and thin slurries (*Buflovak Equip. Div., Blaw* Knox Co.). (a) Single drum dryer with dip feed and spreader. (b) Double drum dryer with splash feed. (c) Double drum dryer with top feed, vapor hood, knives and conveyor. (d) Double drum dryer with pendulum feed, enclosed for vacuum operation.

Single pass residence times are $0.5-3 \sec$, but most commercial operations employ some recycling of the product so that average residence times are brought up to 60sec. Recycling also serves to condition the feed if it is very wet. The spread of residence times in pneumatic dryers, as indicated by Figure 9.5(a), is broad, so feed that has a particularly wide size distribution may not dry uniformly. Recycling, however, assists uniformity, or several dryers in series or preclassification of particle sizes may be employed.

Since the contact time is short, heat-sensitive materials with good drying characteristics are particularly suited to this kind of dryer, but sticky materials obviously are not. Moreover, since attrition may be severe, fragile granules cannot be handled safely. Other kinds of dryers should be considered for materials that have substantial falling rate drying periods.

Pilot plant work is essential as a basis for full scale design. It may be directed to finding suitable velocities, temperatures and drying times, or it may employ more basic approaches. The data provided for Example 9.8, for instance, are of particle size distribution, partial pressure of water in the solution, and heat and mass transfer coefficients. These data are sufficient for the

EXAMPLE 9.7

Description of a Drum Drying System

A detergent drying plant handles 86,722 lb/day of a slurry containing 52% solids and makes 45,923 lb/day of product containing 2% water. The dryers are two sets of steam-heated double drums, each 3.5 ft dia by 10 ft, with a total surface of 440 sqft. Each drum is driven with a 10 HP motor with a variable

speed transmission. Each trolley top spreader has a 0.5 HP motor. Each side conveyor has a 1 HP motor and discharges to a common belt conveyor that in turn discharges to a bucket elevator that feeds a **flaker** where the product is reduced to flakes less than 0.25 in. square. Fines are removed in an air grader and recycled to the dryer feed tank.

TABLE 9.11. Performance Data of Drum Dryers

(a) Drum Dryers

	Yeast Cream	Stone Slop	Starch Solutions	Glaze	Zirconium Silicate	Brewers Yeast	Clay Slip
Feed solids (% by weight)	16	40	36	64	7 0	2 5	75
Product moisture (% w/w basis)	5.7	0.2	5	0.2	0.2	5	9
Capacity (lb prod./hr)	168	420	300-400	225	1120	146	4000
Dryer type (a) single, (b) twin, (c) double Drum	(a)	(a)	(a)	(a)	(a)	(a)	(a)
diameter length	4ftOin. 10 ft 0 in.	2 ft 6 in. 5ftOin.	48 in. 120 in.	18 in. 36 in.	36 in . 72 in.	28 in. 60 in.	48 in. 120 in.
Type of feed method	top roller	dip	top roller	side	dip	center nip	side
Steam pressure (Ib/in ² gauge)	80	60	80	_	80	40	4 0
Atmospheric or vacuum	atmos.	atmos.	atmos.	atmos.	atmos.	atmos.	atmos.
Steam consumption (Ib/Ib evaporated)	_	_	1.3	1.3		_	1.35
Average effective area (%)		_	86				65
Evaporation/ft ² /hr	6.5	4	5	9	8.4	6	8.4

(Courtesy A.P.V. Mitchell Dryers, Ltd.; Williams-Gardner, 1971).

(b) Drum Dryers in the Size Range 0.4 x 0.4-0.8 x 2.25 \mathbf{m}^{a}

Type of Dryer and Feed Size by Letter, A, B, or C	Drum Speed (rev/min)	Steam Press (bar.9)	Type of Material	Physical Form of Feed	Solids in Feed (%)	H₂O in Product (%)	output of Dried Product (g/sec m ²)	Evaporation Rate of Water (g/sec m ²)
			inorganic salts					
Single (dip)	4.4	3.5 ^b	alk. carbs	—	50	8-12	5.5	4.9
Single (splash)	1	3.0	Mg(OH) ₂	thick slurry	35	0.5	1.9	1.5
Twin (splash) A	3	3.0	Fe(OH) ₃	thin slurry	2 2	3.0	4.3	1.3
Double	3-8	5.0	Na Acetate	solution	2 0	0.4-10	2.0-7.0	8-24
Double and twin	7-9	2-3	Na₂SO₄	solution	24	0.15-5.5	4.7-6.1	11-12
Double and twin	5-9	4-6	Na₂HPO₄	solution	44	0.8-0.9	8.2-l 1.1	9-14
Twin (dip) A	5	5.5	organic salts	solution	27	2.8	1.9	5.2
Twin A	3	5.5	organic salts	solution	33	13.0	1.4	2.6
Twin B	2 "	3.5	organic salts	solution	2 0	1.0	1.0	3.8
Twin C	5	5.5	organic salts	solution	39	0.4	3.9	6.1
Twin C	5 ¹ / ₂	5.5	organic salts	solution	4 2	1.0	2.1	4.6
Twin C	6	5.5	organic salts	solution	35	5.0	4.1	7.2
Twin (splash) A	3-5	5.0	organic salts	thin slurry	2 0	1.7-3.1	1.0-I .9	3.7-7.3
Double A	5 <u>1</u>	6.0	organic salts	solution	11	_	1.1	9
Double B	6 <u>1</u>	5-6	organic salts	solution	4 0	3	3.4	4.9
Twin (dip) A	5	5.5		thin slurry	30	1.2	2.4	5.5
Twin A	5	5.0	organic	viscous soln.	28	10.5	1.9	4.2
Double	2	3.0	compounds	viscous soln.	_	6.0	0.7	—
Double	4 <u>1</u>	3.5		thin slurry	2 5	1.0	0.4-1.9	3.5-5.0
Twin (dip)	5	5.0	organic	(a) solution	2 5	0.5	0.3	0.8
Twin	10	5.5	compounds of	(b) thick slurry	30	2.5	2.0	4.6
Twin	10	5.5	low surface	(c) thick slurry	35	_	3.1	
			tension					
Double	11	5.5	similar	(b) thin paste	46		6.4	7.3
			letters					
Double	1 2	5.5	for same	(c) thick paste	58	_	6.0	4.3
Double	11	5.5	compound	(a) solution	2 0	0.5	0.24	1.0

^{(D}Dryer dia and width (m): (A) 0.457 x 0.457; (B) 0.71 x 1.52; (C) 0.91 x 2.54. ^b Plus external hot air flow. ^(Stainless steel drum.)

(Nonhebel and Moss, 1971).

TABLE 9.12. Performance of Drum Dryers

(a) Single, Double drum and Vacuum Drums

		Mathad	Moisture (% W e	Content, et Basis)	Steam	Drum	Feed	Capacity	Vacuum
	Material	of Feed	Feed	Product	(Ib/sq in.)	(rpm)	(°F)	(hr)(sq ft)]	(in. Hg)
Double-drun	n dryer								
Sodium	sulfonate	trough	53.6	6.4	63	8 ¹ / ₂	164	7.75	
Sodium	sulfate	trough	76.0	0.06	56	7	150	3.08	
Sodium	phosphate	trough	57.0	0.9	90	9	180	8.23	
Sodium	acetate	trough	39.5	0.44	70	3	205	1.51	
Sodium	acetate	trough	40.5	10.03	67	8	200	5.16	
Sodium	acetate	trough	63.5	9.53	67	8	170	3.26	
Single-drum	dryer								
Chromium	n sulfate	spray film	48.5	5.47	50	5	_	3.69	
Chromium	n sulfate	dip	48.0	8.06	50	4		1.30	
Chromium	n sulfate	pan	59.5	5.26	24	23	158	1.53	
Chromium	n sulfate	splash	59.5	4.93	55	1 3	150	2.31	
Chromium	n sulfate	splash	59.5	5.35	53	4 <u>3</u>	154	3.76	
Chromium	n sulfate	dip	59.5	4.57	53	5 <u>3</u>	153	3.36	
Vegetable	glue	pan	60-70	10-12	20-30	6-7	_	1-1.6	
Calcium	arsenate	slurry	75-77	0.5-1.0	45-50	3-4	_	2-3	
Calcium	carbonate	slurry	70	0.5	4 5	2-3	_	1.5-3	
Twin-drum	dryer								
Sodium	sulfate	dip	76	0.85	55	7	110	3.54	
Sodium	sulfate	top	69	0.14	6 0	9 ¹ / ₂	162	4.27	
Sodium	sulfate	top	69	5.47	3 2	9 ¹ /2	116	3.56	
Sodium	sulfate	splash	71	0.10	60	6	130	4.30	
Sodium	sulfate	splash	71.5	0.17	60	12	140	5.35	
Sodium	sulfate	splash	71.5	0.09	60	10	145	5.33	
Sodium	phosphate	splash	52.5	0.59	58	5 ¹ / ₂	208	8.69	
Sodium	phosphate	dip	55	0.77	60	$5\frac{1}{2}$	200	6.05	
Sodium	sulfonate	top	53.5	8-10	63	8 <u>1</u>	172	10.43	
Vacuum s	ingle-drum dryer								
Extract		pan	59	7.75	3 5	8	_	4.76	27.9
Extract		pan	59	2.76	35	6	—	1.92	27.9
Extract		pan	59	2.09	36	4	—	1.01	atmos.
Extract		pan	56.5	1.95	3 5	$7\frac{1}{2}$		3.19	22.7
Extract		pan	56.5	1.16	50	2 ¹ /2	—	0.75	atmos.
Skim mi	lk	pan	65	2-3	10-12	4-5	_	2.5-3.2	
Malted n	nilk	pan	60	2	30-35	4-5	—	2.6	
Coffee		pan	65	2-3	5-10	11 1	—	1.6-2.1	
Malt extr	act	spray film	65	3-4	3-5	0.5-1.0	-	1.3-I .6	
Tanning	extract	pan	50-55	8-10	30-35	8–10		5.3-6.4	
Vegetable	glue	pan	60-70	10-12	15-30	5-7	-	2-4	

(Perrys Chemical Engineers Handbook, McGraw-Hill, 1950 edition)

calculation of residence time when assumptions are made about terminal temperatures.

9.9. FLUIDIZED BED DRYERS

Free flowing granular materials that require relatively short drying times are particularly suited to fluidized bed drying. When longer drying times are necessary, multistaging, recirculation or batch operation of fluidized beds still may have advantages over other modes.

A fluidized bed is made up of a mass of particles buoyed up out of permanent contact with each other by a Rowing fluid. Turbulent activity in such a bed promotes high rates of heat and mass transfer and uniformity of temperature and composition throughout. The basic system includes a solids feeding device, the fluidizing chamber with a perforated distributing plate for the gas, an overflow **duct** for removal of the dry product, a cyclone and other equipment for collecting fines, and a heater and blower for the gaseous drying medium.

Much ingenuity has been applied to the design of fluidized bed drying. Many different arrangements of equipment are illustrated and described in the comprehensive book of Kröll (1978) for instance. Figure 9.13(a) depicts the basic kind of unit and the other items are a few of the many variants. Tables 9.14 and 9.15 are selected performance data.

Shallow beds are easier to maintain in stable fluidization and of course exert a smaller load on the air blower. Pressure drop in the air distributor is approximately 1 psi and that through the bed equals the weight of the bed per unit cross section. Some pressure drop data are shown in Table 9.14. The cross section is determined by the gas velocity needed for fluidization as will be described. It is usual to allow 3-6 ft of clear height between the top of the bed and the air exhaust duct. Fines that are entrained are collected in a cyclone and blended with the main stream since they are very dry

TABLE 9.12—(continued)

(b) Single and Double Drum with Various Feed Arrangements

		Moisture Content	Vapor	Potation	Linit Deaduat	Daving
Kind of Dıy er, Kind of Stock	ln (%)	out (%)	Absolute (bar)	Speed (1/min)	Capacity (kg/m² hr)	Rate (kg/m ² hr)
Single drum, dip feed						
Alkali carbonate	50	8 bis 12	. 3.5	4.4	2 0	17.8
Double drum, dip feed						
Organic salt solution	73	2.8	5.5	5	8.8	18.8
Organic compound, dilute slurry	70	1.2	5.5	5	8.6	19.6
Organic compound, solution	75	0.5	5.0	5	1.1	1.9
Single drum with spreading rolls						
Skim milk concentrate	50	4	3.8	24	15.8	14.2
Whey concentrate	4 5	4.3	5.0	16	10 bis 11.8	7.4 bis 8.8
Cuprous oxide	58	0.5	5.2	10	11.0	14.3
Single drum, splash feed						
Magnesium hydroxide, dense slurry	65	0.5	3.0	1	6.8	5.4
Double drum, splash feed						
lron hydroxide, dilute slurry	78	3.0	3.0	3	15.4	4.7
Organic salt, dilute slurry	80	1.7 bis 3.1	5.0	3 bis 5	3.6 bis 6.8	13.3 bis 26.2
Sodium acetate	50	4.0	6.0	5	10.0	9.3
Sodium sulfate	70	2.3	7.8	5	18.0	40.4
Double drum, top feed						
Beer yeast	80	8.0	6.0	5	10.0	38.2
Skim milk, fresh	91.2	4.0	6.4	12	6.2	61.5
Organic salt solution	89		6.0	5.5	4	32.3
Organic salt solution	60	3	5 bis 6	6.5	12.2	17.7
Organic compound, dilute slurry Double drum with spreading rolls	75	1	3.5	4.5	1.4 bis 6.8	12.6 bis 18
Potato pulp	76.2	11.4	8	5	22.5	61.1

(Kröll, 1978, p. 348).

due to their small size. Normally entrainment is 5-10% but can be higher if the size distribution is very wide. It is not regarded as feasible to permit high entrainment and recycle back to the drying chamber, although this is common practice in the operation of catalytic cracking equipment.

Mixing in shallow beds is essentially complete; Figure 9.5(c) shows some test data in confirmation. The corresponding wide

distribution of residence times can result in nonuniform drying, an effect that is accentuated by the presence of a wide distribution of particle sizes. Multiple beds in series assure more nearly constant residence time for all particles and consequently more nearly uniform drying. The data of Table 9.14(b) are for multiple zone dryers. Figures 9.13(c) and (d) have additional zones for cooling the product before it leaves the equipment. Another way of assuring

TABLE SITULT CITOTINATION DATA OF THE ANALY CONTENTING DIVERSION CONCENTES IN THE SITE	onveying Dryers (Sketches in Fig. 9.12)
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(a) Raymond Flash Dryer

	Fine Mineral	Spent Grain	Organic Chemical	Chicken Droppings	Fine Coal Filter Cake
Method of feed	pump	belt	screw	pump	screw
Material size, mesh	- 100	—	-30	_	-30
Product rate (Ib/hr)	27,000	9000	900	2300	2000
Initial moisture content (% w/w basis)	2 5	60	37	70	30
Final moisture content (% w/w basis)	nil	12	3	12	8.5
Air inlet temperature (°F)	1200	1200	450	1300	1200
Air outlet temperature (°F)	200/300	200/300	200/300	200/300	200/300
Method of heating	direct	direct	direct	direct	direct
	oil	oil	oil	oil	oil
Heat consumption (Btu/Ib water evaporated)	1.6 × 10 ³	1.9 x 10 ³	3.1 x 10 ³	1.9 x 10 ³	1.4x 10 ³
Air recirculation	n o	n o	n o	n o	no
Material recirculation	yes	yes	n o	yes	no
Material of construction	MS	MS/SS	MS	MS	MS
Fan capacity (std. ft ³ /min)	18,000	22,000	4300	8500	1500
Installed fan HP	110	180	30	50	10
Product exit temperature (°F)	200	-	200	—	135

(Courtesy International Combustion Products Ltd.; Williams-Gardner, 1971)

TABLE 9.13—(continued)

(b) Buttner-Rosin Pneumatic Dryer

	Metallic Stearate	Starch	Adipi Acid	Fiber	Coal Filter Cake
Method of feed	sling	sling	screw	distributor	distributor
Material size	fine	fine	-30 mesh	$-\frac{1}{4}$ in.	-30 mesh
Product rate (Ib/hr)	280	13,236	10,000	2610	67,200
Initial moisture (% w/w basis)	4 0	34	10	62.4	3 2
Final moisture (% w/w basis)	0.5	13	0.2	10	6
Air inlet temperature (°F)	284	302	320	752	1292
Air outlet temperature (°F)	130	122	149	230	212
Method of heating	steam	steam	steam	oil	PF
Heat consumption (Btu/Ib/water evaporated)	2170	1825	2400	1720	1590
Air recirculation	n o	n o	no	n o	yes
Material recirculation	yes	n o	yes	yes	yes
Fan capacity (std. ft ³ /min)	1440	26,500	9500	12,500	27,000
Installed fan HP	15	220	65	60	250
Product exit temperature (°F)	104	95	120	140	158

(Courtesy Rosin Engineering Ltd.; Williams-Gardner, 1971).

(c) Pennsalt-Berks Ring Dryer

	Metal ^a Stearates	Spent" Grains	Sewage' Sludge	Starches	Polystyrene Beads
Method of Feed	belt feeder rotary valve	back mixer rotary valve	vibratory feeder rotary valve	cascading rotary valve screen	vibratory feeder rotary valve
Product rate (Ib/hr)	240	1120	4300	5000	1000
Initial moisture (% w/w basis)	55	80	4 5	3 5	2.0
Final moisture (% w/w basis)	1	5	12	10	0.2
Air inlet temperature (°F)	250	500	600	300	175
Air outlet temperature (°F)	150	170	170	130	115
Method of heating	steam	gas	oil	steam	steam
Heat consumption (Btu/lb water evaporated)	2900	1800	1750	2000	5000
Air recirculation	no	no	no	n o	no
Material recirculation	ves	ves	ves	no	no
Material of construction	ss	MSG	MS	MSG	SS
Fan capacity (std ft ³ /min)	3750	16,500	8250	15,000	900
Installed fan HP	2 0	75	6 0	60	7.5

^a Ring dryer application. (Courtesy Pennsalt Ltd.; Williams-Gardner, 1971).

(d) Various Pneumatic Dryers

		Tube T Dia He (cm) (Tube	Gas Rate (m ³ /hr) (NTP)	Gas Temp(°C) Solid		Solid Temp (°C) Moisture (%)			Air/Solid Ratio		Water		
Material Location	Height (m)		In		Out	- Rate (kg/hr)	In	Out	In	out	(m` (NTP)	"/kg) (kg/kg)	Evaporated (kg/hr)	
Ammonium sulphate	Japan	18	1	1100	215	76	950	38.5	63	2.75	0.28	1.2	1.5	23.5
Sewage sludge filter cake Coal 6 mm	U.S.A. U.S.A.		-	1200 50,000	700 371	121 80	2270 51,000	15 15	71 57	8 0 9	10 3	5.3 1.0	7.2 1.3	1590 4350
Hexamethylene tetramine	Germany	3 0	38″	3600	93	50	2500	_	48	6-10	0.08-0.15	1.4	1.9	18.1

"23 m vertical, 15 m horizontal.

(Nonhebel and Moss, 1971).



Figure 9.12. Examples of pneumatic conveying dryers; corresponding performance data are in Table 9.13. (a) Raymond flash dryer, with a hammer mill for disintegrating the feed and with partial recycle of product (Raymond *Division, Combustion Engineering)*. (b) Buttner-Rosin pneumatic dryer with separate recycle and disintegration of large particles (*Rosin Engineering Ltd.*). (c) Berks ring dryer; the material circulates through the ring-shaped path, product is withdrawn through the cyclone and bag filter (*Pennsalt Chemical Co.*).

complete drying is a recirculation scheme like that of Figure 9.13(e). In batch operation the time can be made as long as necessary.

Stable fluidization requires a distribution of particle sizes, preferably in the range of a few hundred microns. Normally a size of 4mm or so is considered an upper limit, but the coal dryers of Tables 9.15(a) and (b) accommodate sizes up to 0.5 in. Large and uniformly sized particles, such as grains, are dried successfully in spouted beds [Fig. 9.13(f)]: Here a high velocity gas stream entrains the solid upward at the axis and releases it at the top for **flow** back through the **annulus**. Some operations do without the mechanical draft tube shown but employ a naturally formed central channel.

One way of drying solutions or pastes under fluidizing conditions is that of Figure 9.13(g). Here the tluidized mass is of auxiliary spheres, commonly of plastic such as polypropylene, into which the solution is sprayed. The feed material deposits uniformly on the spheres, dries there, and then is knocked off automatically as it leaves the drier and leaves the auxiliary spheres behind. When a mass of dry particles can be provided to start a fluidized bed drying process, solutions or pastes can be dried after deposition on the seed material as on the auxiliary spheres. Such a process is employed, for instance, for growing fertilizer granules of desired larger sizes, and has largely replaced rotary dryers for this purpose.

A few performance data of batch fluid dryers are in Table

Sizing a Pneumatic Conveying Dryer

A granular solid has a moisture content of 0.035 kg/kg dry material which is to be reduced to 0.001 kg/kg. The charge is at the rate of 9.72 kg/sec, is at 60°C and may not be heated above 90°C. Inlet air is at 450°C and has a moisture content of 0.013 kg/kg dry air.



Specific gravity of the solid is 1.77 and its heat capacity is 0.39 cal/g "C. The settling velocity of the largest particle present, 2.5 mmdia, is 10 m/sec. Heat capacity of the air is taken as 0.25 cal/g °C and the latent heat at 60°C as 563 cal/g. Experimental data for this system are reported by Nonhebel and Moss (1971, pp. 240ff) and are represented by the expressions:

Heat transfer coefficient:

ha = 0.47 cal/(kg solid)(C)

Vapor pressure:

 $P = \exp(13.7419 - 5237.0/T), atm, K$

Mass transfer coefficient:

 $k_{p}a = \exp(-3.1811 - 1.7388 \ln w - 0.2553(\ln w)^2)$

where **w** is the moisture content of the solid (kg/kg) in the units kg water/(kg solid)(atm)(sec).

In view of the strong dependence of the mass transfer coefficient on moisture content and the **35-fold** range of that property, the required residence time and other conditions will be found by analyzing the performance over small decrements of the moisture content.

An air rate is selected on the assumption that the exit of the solid is at 85° C and that of the air is 120° C. These temperatures need not be realized exactly, as long as the moisture content of the exit air is below saturation and corresponds to a partial pressure less than the vapor pressure of the liquid on the solid. The amount of heat transferred equals the sum of the sensible heat of the wet solid and the latent heat of the lost moisture. The enthalpy balance is based on water evaporating at 60° C:

$$\bar{m}_s[(0.39 + 0.001)(85 - 60) + (0.035 - 0.001)(85 - 60 + 563)] = \bar{m}_a[(0.25 + 0.480(0.001))(450 - 120) + 0.48(0.034)(120 - 60)],$$

$$\bar{m}_{a} = \frac{29.77 \bar{m}_{s}}{63.04} = \frac{29.77 (9.72)}{63.04} = \begin{cases} 3.46 \text{ kg/sec}, \\ 7.08 \text{ m}^{3}/\text{sec} \text{ at } 450^{\circ}\text{C}, \\ 3.85 \text{ m}^{3}/\text{sec} \text{ at } 120^{\circ}\text{C}. \end{cases}$$

At a tower diameter of 0.6 m,

$$U = \frac{Q}{0.36\pi/4} = \begin{cases} 25.0 \text{ m/sec at } 450^{\circ}\text{C}, \\ 13.6 \text{ m/sec at } 120^{\circ}\text{C}. \end{cases}$$

These velocities are great enough to carry the largest particles with settling velocity of 10 m/sec.

Equations are developed over intervals in which $W_1 \rightarrow W_2$, $T_1 \rightarrow T_2$, and $T'_1 \rightarrow T'_2$. The procedure will be:

1. Start with known W_1 , T_1 , and T'_1 .

- 2. Specify a moisture content W_2 .
- 3. Assume a value T_2 of the solid temperature.
- 4. Calculate T'_2 from the heat balance.
- 5. Check the correctness of T_2 by noting if the times for heat and mass transfers in the interval are equal.

$$\theta_{h} = \frac{Q}{ha(\Delta T)_{\text{lm}}} = \frac{Q}{0.47(\Delta T)_{\text{lm}}}$$
$$\theta_{m} = \frac{w_{1} - w_{2}}{k_{g}a(\Delta P)_{\text{lm}}}$$

Heat balance:

$$\begin{split} & \bar{m}_s [0.391(T_2 - T_1) + (W_1 - W_2)(T_2 - T_1 + 563)] \\ &= \bar{m}_a \{ [0.25 + 0.48(0.001](T_1' - T_2) \\ &+ 0.48(W_1 - W_2)(T_2' - 60)). \end{split}$$

Substitute $\bar{m}_s/\bar{m}_a = 9.72/3.46 = 2.81$ and solve for T'_2 .

$$T'_{2} = -0.25048T'_{1} + 28.8(W_{1} \quad W_{2}) + 2.81 \\ \times [0.39(T_{2} - T_{1}) + (W_{1} - W_{2})(T_{2} - T_{1} + 563)] \\ \hline 0.48(W_{1} - W_{2}) - 0.25048$$
(1)

$$g_1 = 0.013 + \frac{\bar{m}_s}{\bar{m}_a}(W_1 - 0.013) = 0.013 + 2.81(W_1 - 0.013).$$
 (2)

$$P_1 = \frac{g_1}{18/29 + g_1} = \frac{g_1}{0.6207 + g_1}$$
(partial pressure in air). (3)
$$g_1 = 0.012 + 2.81(W = 0.012)$$
(4)

$$g_2 = 0.013 + 2.61(w_2 - 0.013). \tag{4}$$

$$P_2 = \frac{\delta^2}{0.6207 + g_2}.$$
 (5)

$$Pa_1 = \exp[13.7419 - 5237.9/(T_1 + 273.2)],$$
 vapor pressure. (6)
 $Pa_2 = \exp[13.7419 - 5237.9/(T_2 + 273.2)].$ (7)

$$(\Delta P)_{\rm im} = \frac{(Pa_1 - P_1) - (Pa_2 - P_2)}{\ln[(Pa_1 - P_1)/(Pa_2 - P_2)]} \tag{8}$$

$$(\Delta T)_{\rm lm} = \frac{T_1' - T_1 - (T_2' - T_2)}{\ln[(T_1' - T_1)/(T_2' - T_2)]} \tag{9}$$

$$AQ = 0.391(T_2 - T_1) + (W_1 - W_2)(T_2 - T_1 + 563),$$

per kg of solid. (10)

$$\bar{W} = 0.5(W_1 + W_2). \tag{11}$$

$$k_g a = \exp[-3.1811 - 1.7388 \ln \bar{W} - 0.2553 (\ln \bar{W})^2].$$
(12)

$$\theta_h = \Delta Q / ha(\Delta T)_{\rm lm} = \Delta Q / 0.43 (\Delta T)_{\rm lm}, \text{ heating time.}$$
(13)
$$\theta_m = (W_1 - W_2) / k_e a(\Delta P)_{\rm lm}, \text{ mass transfer time.}$$
(14)

$$\mathbf{Z} = \boldsymbol{\theta}_{h} - \boldsymbol{\theta}_{m} \rightarrow 0$$
 when the correct value of T_{2}

(15)

After the correct value of T_2 has been found for a particular interval, make $W_2 \rightarrow W_1$, $T_2 \rightarrow T_1$, and $T'_2 \rightarrow T'_1$. Specify a

EXAMPLE 9.8—(continued)

decremented value of W_2 , assume a value of T_2 , and proceed. The solution is tabulated.

w	Τ	T′	0 (sec)
0. 035	60	450	0
0.0325	73.04	378. 2	0.0402
0.03	75.66	352.2	0.0581
0.025	77.41	315.3	0.0872
0.02	77.23	286.7	0.1133
0.015	76.28	261.3	0.1396
0. 01	75.15	236.4	0.1687
0.005	74.67	208.4	0.2067
0.003	75.55	192.4	0.2317
0.001	79.00	165.0	0.2841

When going directly from 0.035 to 0.001,

$$T_2 = 80.28,$$

 $T'_2 = 144.04,$
 $\theta = 0.3279$ sec.

The calculation could be repeated with a smaller air rate in order to reduce its exit temperature to nearer 120°C, thus improving thermal efficiency.

In the vessel with diameter = 0.6 m, the air velocities are

 $u_a = \begin{cases} 25.0 \text{ m/sec} & \text{at } 450^{\circ}\text{C} \text{ inlet} \\ 5.15 \text{ m/sec} & \text{at } 165^{\circ}\text{C} \text{ outlet} \end{cases}$

20.1 m/sec average.

The vessel height that will provide the needed residence time is $H = \tilde{u}_a \theta = 20.1(0.2841) = 5.70$ m.

Very fine particles with zero slip velocity will have the same holdup time as the air. The coarsest with settling velocity of 10 m/sec will have a net forward velocity of

 $\bar{u}_{s} = 20.1 - 10 = 10.1 \text{ m/sec},$

which corresponds to a holdup time of

 $\theta = 5.7/10.1 = 0.56$ sec,

which is desirable since they dry more slowly. After the assumption of T_2 , other quantities are evaluated in the order shown in this program.

```
10 ! Example 9.3. Pneumat 1C con
  vering dryer
20 ! Finding the exit solids te
  meT2bytrial, then all dep
  sndent quantities
```

9.14(a). This process is faster and much less labor-intensive than tray drying and has largely replaced tray drying in the pharmaceutical industry which deals with small production rates. Drying rates of 2-10 lb/(hr)(cuft) are reported in this table, with drying times of a fraction of an hour to several hours. In the continuous operations of Table 9.15, the residence times are at most a few minutes.

Thermal efficiency of fluidized bed dryers is superior to that of many other types, generally less than twice the latent heat of the water evaporated being required as heat input. Power requirements are a major cost factor. The easily dried materials of Table 9.15(a) show evaporation rates of 58-103 lb/(hr)(HP installed) but the more difficult materials of Table 9.15(d) show only 5-18 lb/(hr) (HP

- 30 INPUT W1, W2, T1, A1! A1 is th e inlet air tempTl' 40 INPUT T2 ! Trial value 50 A2=(2.81*(.391*(T2-T1)+(W1-W
- 2)*(T2-T1+563))-.25048*A1+28 8*(W1-W2))/(.48*(W1-W2)-.25 048)
- 60 G1=.013+2.81*(W1-.013)
- 70 P1=G1/(.6207+G1)
- 80 G2=.013+2.81*(W2-.013)
- 90 P2=G2/(.6207+G2)
- 100 Q1=EXP(13.7419-5237.9/(T1+27 3.2))! vapor pressure
- 110 Q2=EXP(13.7419-5237.9/(T2+27 3.2>>
- 120 P3=(Q1-P1-Q2+P2)/LOG((Q1-P1) /(Q2-P2)) ! (AP)1m
- 130 T3=(A1-T1-A2+T2)/LOG((A1-T1)) /(A2-T2)) ! (&T)1m
- 140 Q=.391*(T2-T1)+(W1-W2)*(T2-T 1+563)
- H1=Q/.47/T3! heatingtime 150
- 160 W=.5*(W1+W2) 170 K=EXP(-3.1811-1.7388*LOG(W)-.2533*LOG(W)^2)
- 180 H2=(W1-W2)/K/P3 ! vaporizati on time
- Z=H1-H2 ! time difference sh 190 ould be zero
- DISP Z 200
- 210
- DISP A2,H1 GOTO 40 ! if Z is not near e 220 nough to zero; otherwise the correct value of T2 has bee n found
- 230 END

Data	for	the	first	interval
		-		
W1 =	.03	5 195		
W2=	- 03	20		
11=	(). - 45	194 Ga		
(1 -	- 40	10		
T2=	73.	04		
Ť2+*	= 37	8.16	69691	11
Time	= 4	.023	28366	0795E-2

installed). The relatively large power requirements of fluidized bed dryers are counterbalanced by their greater mechanical simplicity and lower floor space requirements.

Air rates in Table 9.15 range from 13 to 793 SCFM/sqft, which is hardly a guide to the selection of an air rate for a particular case. A gas velocity twice the minimum fluidization velocity may be taken as a safe prescription. None of the published correlations of minimum fluidizing velocity is of high accuracy. The equation of Leva (Fluidization, McGraw-Hill, New York, 1959) appears to be as good as any of the later ones. It is

$$G_{mf} = 688D_p^{1.83}[\rho_g(\rho_s - \rho_g)]^{0.94}/\mu^{0.88}, \qquad (9.20)$$

where G_{mf} is in lb/(hr)(sqft), ρ_g and ρ_s are densities of the gas and solid (lb/cuft), D_p is the particle diameter (in.), and μ is the gas viscosity (cP). In view of the wide scatter of the data on which this correlation is based, shown on Figure 6.14(f), it appears advisable to find the fluidization velocity experimentally for the case in hand.

Although it is embarrassing again to admit the fact, unfortunately all aspects of fluidized bed drying must be established with pilot plant tests. The wide ranges of performance parameters in Tables 9.14 and 9.15 certainly emphasize this conclusion. A limited exploration of air rates and equipment size can be made on the basis of a drying rate equation and fluidization correlations from the literature. This is done in Example 9.9. A rough approximation of a drying rate equation can be based on through circulation drying of the granular material on a tray, with gas flow downward.

9.10. SPRAY DRYERS

Suitable feeds to a spray dryer are solutions or **pumpable** pastes and slurries. Such a material is atomized in a nozzle or spray wheel, contacted with heated air or flue gas and conveyed out of the equipment with a pneumatic or mechanical type of conveyor. Collection of fines with a cyclone separator or filter is a major aspect of spray dryer operation. Typical equipment arrangements and flow patterns are shown in Figure 9.14.

The action of a high speed spray wheel is represented by Figure 9.14(e); the throw is lateral so that a large diameter vessel is required with this form of atomization, as shown in Figure 9.14(a). The flow from nozzles is largely downward so that the dryer is slimmer and taller. Parallel flow of air and spray downward is the most common



Figure 9.13. Fluidized bed dryers. (a) Basic equipment arrangement (McCabe and Smith, Unit Operations in Chemical Engineering, McGraw-Hill, New York, 1984). (b) Multiple bed dryer with dualflow distributors; performance data are in Table 9.14(b) (Romankov, in Davidson and Harrison, Fluidisation, Academic, New York, 1971). (c) A two-bed dryer with the lower one used as cooler: (a, b, c) rotary valves; (d) drying bed; (e) cooling bed; (f, g) air distributors; (h, i) air blowers; (k) air filter; (1) air heater; (m) overflow pipe; (n) product collector (Kroll, 1978). (d) Horizontal multizone dryer: (a) feeder; (b) air distributor; (c) fluidized bed; (d) partitions; (e) dust guard; (f) solids exit; (g) drying zone; (h) cooling zone; (i, k) blowers; (1, m) air plenums; (n) air duct; (o) dust collector; (p) exhaust fan (Kroll, 1978). (e) Circulating fluidized bed used for removal of combined water from aluminum hydroxide: (a) feed; (b) fluidized bed; (c) solids exit; (d) fuel oil inlet; (e) primary air inlet; (f) secondary air inlet; (g) gas exit (Kroll, 1978). (f) Spouted bed with draft tube for drying coarse, uniform-sized granular materials such as grains [Yang and Keairns, AIChE Symp. Ser. 176, 218 (1978), Fig. 1]. (g) Fluidized bed dryer for sludges and pastes. The fluidized solids are fine spheres of materials such as polypropylene. The wet material is sprayed in, deposits on the spheres and dries thre. At the outlet the spheres strike a plate where the dried material is knocked off and leaves the dryer as flakes. The auxiliary spheres remain in the equipment: (a) feed; (b) distributor; (c) spheres loaded with wet material; (d) returning spheres; (e) striking plate; (f) hot air inlet; (g) air and solids exit (Kroll, 1978).



Figure 9.13—(continued)

arrangement, but the left-hand figure of Figure 9.14(d) is in counterflow. Figure 9.14(c) has tangential input of cooling air. In some operations, the heated air is introduced tangentially; then the process is called mixed flow. Most of the entries in Table 9.16(a) are parallel flow; but the heavy duty detergent is in counterflow, and titanium dioxide is either parallel or mixed flow. **Counterflow** is thermally more efficient, results in less expansion of the product

particles, but may be harmful to thermally sensitive products because they are exposed to high air temperatures as they leave the dryer. The flat bottomed dryer of Figure 9.14(c) contacts the exiting solids with cooling air and is thus adapted to thermally sensitive materials.

Two main characteristics of spray drying are the short drying time and the porosity and small, rounded particles of product. Short
TABLE 9.14. Performance Data of Fluidized Bed Dryers: Batch and Multistage Equipment

(a) Batch Dryers

	Ammonium Bromide	Lactose Base Granules	Pharmaceutical Crystals	Liver Residue	Weed Killer
Holding capacity (Ib wet product)	100	104	160	280	250
Bulk density, dry (Ib/ft ³)	75	30	2 0	30	35
Initial moisture (% w/w basis)	6	10	6 5	50	20-25
Final moisture (% w/w basis)	1	2	0.4	5.0	1.0
Final drying temperature (°F)	212	158	248	140	140
Drying time (min)	2 0	90	120	75	210
Fan capacity (ft ³ /min at 11 in. w.g.)	750	1500	3000	4000	3000
Fan HP	5	10	2 0	2 5	2 0
Evaporation rate (Ib H ₂ O/hr)	1 5	5.7	5 2	100	17

(Courtesy Calmic Engineering Co. Ltd.; Williams-Gardner, 1971).

(b) Multistage Dryers with Dual-flow Distributors [Equipment Sketch in Fig. 9.13(b)]

Function	Heater	Cooler	Drier	Cooler
Material	Wheat Grains	Wheat Grains	Slag	Quartz Sand
Particle size (diameter)(mm)	5 x 3	5 x 3	0.95	1.4
Material feed rate (metric tons/hr)	1.5	1.5	7.0	4.0
Column diameter (m)	0.90	0.83	1.60	1.70
Perforated trays (shelves):				
Hole diameter (mm)	2 0	20	20; 10	2 0
Proportion of active section	0.4	0.4	0.4; 0.4	0.4
Number of trays	10	6	1; 2	2 0
Distance between trays (mm)	2 0	2 0	25; 40	15
Total pressure drop on fluidized bed (kgf/m ²)	113	64	70*	40
Hydraulic resistance of material on one tray (kgf/m ²)	7.8	9.2	20; 10	1.8
Inlet gas temperature (°C)	265	38	300	2 0
Gas inlet velocity (m/sec)	8.02	3.22	4.60	0.74
Material inlet temperature (°C)	68	175	2 0	350
Material discharge temperature (°C)	175	54	170	2 2
Initial humidity (% on wet material)	2 5	_	8	-
Final humidity (% on wet material)	2.8	_	0.5	-
Blower conditions				
Pressure (kgf/m²)	450	250	420	250
Throughput (m ³ /min)	180	130	360	100
	(80°C)	(50°C)	(70°C)	(35°C)
Power consumption (HP)	50	2 0	7 5	7.5

⁸ With grids and two distributor plates.

(Romankov, in Davidson and Harrison, Fluidisation, Academic, New York, 1971).

drying time is a particular advantage with heat sensitive materials. Porosity and small size are desirable when the material subsequently is to be dissolved (as foods or detergents) or dispersed (as pigments, inks, etc.). Table 9.17 has some data on size distributions, bulk density, and power requirements of the several types of atomizers.

The mean residence time of the gas in a spray dryer is the ratio of vessel volume to the volumetric flow rate. These statements are made in the literature regarding residence times for spray drying:

Source	Time (sec)
/feat Exchanger Design Handbook (1983)	5-60
McCormick (1979)	2 0
Masters (1976)	20-40 (parallel flow)
Nonhebel and Moss (1971)	<60
Peck (1983)	5-30
Wentz and Thygeson (1979)	<60
Williams-Gardner (1971)	∫ 4-10 (<15 ft dia)
) 10–20 (>15 ft dia)

Residence times of air and particles are far from uniform; Figure 9.5(a) and (b) is a sample of such data.

Because of slip and turbulence, the average residence times of particles are substantially greater than the mean time of the air, definitely so in the case of countercurrent or mixed flow. Surface moisture is removed rapidly, in less than 5 sec as a rule, but falling rate drying takes much longer. Nevertheless, the usual drying operation is completed in 5-30 sec. The residence time distribution of particles is dependent on the mixing behavior and on the size distribution. The coarsest particles fall most rapidly and take longest for complete drying. If the material is heat-sensitive, very tall towers in parallel flow must be employed; otherwise, countercurrent or mixed flows with high air temperatures may suffice. In some cases it may be feasible to follow up incomplete spray drying with a pneumatic dryer.

Drying must be essentially completed in the straight sided zones of Figures 9.14(a) and (b). The conical section is for gathering and efficient discharge of the dried product. The lateral throw of spray wheels requires a vessel of large diameter to avoid

TABLE 9.15. Performance Data of Continuous Fluidized Bed Dryers

(a) Data of Fluosatatic Ltd.

	Coal	Sand	Silica Sand	Limestone	Iron Ore
Material size, mesh	¹ / ₂ -0	-25-0	-18-0	$\frac{3}{16} - 0$	- 3 -0
Method of feed	twin	bucket	conv.	co"".	conv.
	screw	elev.			
Product rate (Ib product/hr)	448,000	22,400	112,000	67,000	896,000
Initial moisture (% w/w basis)	11	6	6	15	3
Final moisture (% w/w basis)	5.5	0.1	0.1	0.1	0.75
Residence time (min)	1	1.25	1.5	1.25	0.5
Dryer diameter (ft)	10	3.0	7.25	5.5	8.5
Fluid bed height fin.)	18	12	12	12	18
Air inlet temperature (°F)	1000	1200	1200	1200	1200
Air outlet temperature (°F)	170	212	212	212	212
Air quantity (ft ³ /min std.)	40,000	2000	9000	13,000	45,000
Material exit temperature (°F)	140	220	220	220	220
Evaporation (Ib/hr)	24,640	1430	6720	11,880	20,400
Method of heating	coal	gas	oil	oil	oil
Heat consumption (Btu/Ib water evaporated)	1830	1620	1730	1220	2300
Fan installed HP	240	2 0	8 0	115	350

(Williams-Gardner, 1971).

(b) Data of Head Wrightston Stockton Ltd.

	Coal	Siliciou Grit	Glass Sand	Sand	Asphalt
Method of feed	screw feeder	chute	chute	c h u t e	chute
Material size	- 1/2 in.	-&in.	-36 mesh	-&in.	-&in.
Product rate (b product/hr)	190,000	17,920	15,680	33,600	22,400
Initial moisture (% w/w basis)	14	5	7	5	5
Final moisture (% w/w basis)	7	0	0	0	0.5
Residence time (min)	2	11	3	3	10
Dryer diameter	7 ft 3 in.	3 ft Ö in.	4 ft 6 in.	6 ft 6 in.	8 ft 0 in.
Fluid bed height (in.)	21	12	12	12	2 4
Air inlet temperature (°F)	1000	1400	1400	1400	470
Air outlet temperature (°F)	135	230	230	230	220
Air quantity (ft ³ /min std)	20,000	2000	2000	3500	7000
Material exit temp (°F)	140	230	230	230	220
Evaporated rate (Ib/hr)	11,200	896	1097	1680	1120
Method of heating	coke-	gas oil	t o w n	gas oil	gas oil
-	oven	-	gas	•	
	gas				
Heat consumption (Btu/Ib water evaporated)	2000	2250	2000	2200	1800
Fan installed HP	210	32 ¹ / ₂	18	3 0	90

(Williams-Gardner, 1971).

(c) Data of Pennsalt Ltd.

	Abrasive Grit	Clay Granules	Sand	Granular Desiccant	Household Salt
Product rate (Ib/hr)	2200	1000	14,000	150	13,500
Initial moisture (% w/w basis)	9	2 2	6	2 5	4
Final moisture (% w/w basis)	dry	3	dry	7	0.03
Air inlet temperature (°F)	580	160	325	300	390
Air outlet temperature (°F)	210	120	140	205	230
Method of heating	gas	steam	gas	gas	steam
Heat consumption (Btu/Ib water evaporated)	2700	3800	2700	3600	5100
Bulk density (Ib/ft ³)	120	60	90	30	60
Average drying time (min)	2.5	30	3	2 4	4
Fan capacity (ft ³ /min std.)	2.5	1.35	1.05	0.84	1.05
Installed fan HP	10	4 5	2 5	5	5 0

(Williams-Gardner, 1971).

272 DRYERS AND COOLING TOWERS

TABLE 9.15—(continued)

(d) Data of Rosin Engineering Ltd.

	Sodium Perborate	Weed Killer	PVC	Coal	Sand
Method of feed Material size	screw 30-200	vibrator 5-I mm	screw 60-1 20	vibrator 3 mesh–	vibrator 30-120
Product rate (Ib product/hr)	mesh 11,400	flake 5100	mesh 10,075	zero 440,000	mesh 112,000
Initial moisture (% w/w basis)	3.5	14	2.0	8	8
Final moisture (% w/w basis)	0.0	0.2	0.2	1	0.2
Residence time (min)	1.5	11	30	0.3	0.45
Drier bed size (ft x ft)	22.5 x 5.5	18 x 4.5	23 x 6	16 x 6.6	12.5 x 3.2
Fluid bed height (in.)	4	3	18	5	6
Air inlet temperature (°F)	176	212	167	932	1202
Air outlet temperature (°F)	104	150	122	180	221
Air quantity (ft ³ /min std)	6600	14,200	5400	67.330	8000
Material exit temperature (°F)	104	205	122	180	212
Evaporation (lb/hr)	400	720	183	33,440	9750
Method of heating	steam	steam	steam	coke-oven	oil
Heat consumption (Btu/Ib water ovenerated)	2100	2060	4640	gas	2200
Fan installed HP	3 3	40	4640 34	600	70

(Williams-Gardner, 1971).

EXAMPLE 9.9

Sizing a Fluidiied Bed Dryer

A wet solid at 100°F contains W = 0.3 lb water/lb dry and is to be dried to W = 0.01. Its feed rate is 100 lb/hr dry. The air is at 350°F and has $H_{g0} = 0.015$ lb water/lb dry. The rate of drying is represented by the equation

$$-\frac{dW}{d\theta} = 60(H_s - H_g), \quad (lb/lb)/min.$$

The solid has a heat capacity 0.35 $Btu/(lb)({}^{\circ}F)$, density 150 lb/cuft, and average particle size $0.2 \,\mu m$ (0.00787 in.). The air has a viscosity of 0.023 cP and a density of 0.048 lb/cuft. The fluidized bed may be taken as a uniform mixture. A suitable air rate and dimensions of the bed will be found



Symbols used in the computer program are in parentheses.

Minimum fluidizing rate by Leva's formula:

$$G_{mf} = \frac{688D_p^{1.83}[0.048(150 - 0.048)]^{0.94}}{\mu^{0.88}}$$

$=\frac{688(0.00787)^{1.83}[0.048(150 \quad 0.048)]^{0.94}}{(0.023)^{0.88}}$

= 17.17 lb/(hr)(sqft).

Let $G_f = 2G_{m_f} = 34.34 \text{ lb/(hr)(sqft)}$. Expanded bed ratio

 $(L/L_0) = (G_f/G_{m_f})^{0.22} = 2^{0.22} = 1.16.$

Take voidage at minimum fluidization as

$$\boldsymbol{\varepsilon}_{m_f} = 0.40,$$

 $\boldsymbol{\cdot}, \boldsymbol{\varepsilon}_{\boldsymbol{\epsilon}} = 0.464$

Drying time:

$$\theta = \frac{W_0 - W \quad 0.3 \quad 0.01}{60(H_s - H_g) = 60(H_s - H_g)}$$
(1)

Since complete mixing is assumed, H_s and H_g are exit conditions of the fluidized bed.

Humidity balance:

$$\dot{A}(H_g - H_{g_0}) = S(W_0 - W),$$

 $H_g = 0.015 + 0.29\dot{S}/\dot{A}.$ (2)

Average heat capacity:

$$C_g = \frac{1}{2}(C_{g_0} + C_g) = 0.24 + 0.45[(0.015 + H_g)/2]$$

= 0.2434 + 0.225H_g. (3)

Heat balance:

$$AC_g(T_{g_0} - T_g) = S[(C_s + W)(T_s - T_{s_0}) + \lambda(W_0 - W)],$$

(A/S)C_g(350 - T_g) = 0.36(T_s - 100) + 900(0.29). (4)

TΞ

EXAMPLE **9.9**—(continued)

Adiabatic saturation line:

$$T_{g} - T_{s} = \frac{\lambda}{C_{g}} (H_{s} - H_{g}) = \frac{900}{C_{g}} (H_{s} - H_{g}).$$
(5)

Vapor pressure:

$$P_s = \exp[11.9176 - 7173.9/(T_s + 389.5)].$$
(6)

Saturation humidity:

$$H_{s} \frac{P_{s} 18}{291 - P_{s}}.$$
 (7)

Eliminate T_3 between Eqs. (4) and (5):

$$T_{s} = 350 - \frac{0.36(T_{4} - 100) + 261}{RC_{g}}$$
$$= T_{4} + \frac{900(H_{4} - H_{3})}{C_{g}}, \ [T_{3} \equiv T_{g}, \ T_{4} \equiv T_{s}].$$
(8)

Procedure: For a specified value of $R = \dot{A}/\dot{S}$, solve Eqs. (6), (7), and (8) simultaneously.

T _g	T _s	Hg	H,	θ (min)
145. 14	119.84	0.0730	0.0803	0.862
178.11	119.74	0.0633	0.0800	0.289
220.09	119.60	0.0513	0.0797	0.170
245.72	119.52	0.0440	0.0795	0.136
262.98	119.47	0.0392	0.0794	0.120
	T _g 145. 14 178. 11 220. 09 245. 72 262. 98	T _g T _s 145. 14 119. 84 178. 11 119. 74 220. 09 119. 60 245. 72 119. 52 262. 98 119. 47	T _g T _s H _g 145. 14 119. 84 0. 0730 178. 11 119. 74 0. 0633 220. 09 119. 60 0. 0513 245. 72 119. 52 0. 0440 262. 98 119. 47 0. 0392	T _g T _s H _g H _s 145.14 119.84 0.0730 0.0803 178.11 119.74 0.0633 0.0800 220.09 119.60 0.0513 0.0797 245.72 119.52 0.0440 0.0795 262.98 119.47 0.0392 0.0794

Take

R = 10 lb air/lb solid. $\dot{A} = 10(100) = 1000 \text{ lb/hr},$ $\theta = 0.136$ min.

Cross section:

 $\dot{A}/G_f = 1000/34.34 = 29.12$ sqft, 6.09 ft dia.

Avg density:

 $\frac{1}{2}(1/20.96 + 1/19.03) = 0.0501$ lb/cuft.

Linear velocity:

$$u = \frac{G_f}{\rho \varepsilon(60)} = \frac{34.34}{0.0501(0.464)(60)} = 24.62$$
 fpm.

Bed depth:

 $L = u\theta = 24.62(0.136) = 3.35$ ft.

Note: In a completely mixed fluidized bed, the drying time is determined by the final moisture contents of the air and solid. When drying is entirely in the falling rate period with rate equation

$$-\frac{dW}{d\theta}=\frac{k(H_s-H_g)}{W_c}W, \quad W\leq W_c,$$

the drying time will be

$$\theta = \frac{W_c}{k(H_s - H_g)W}$$

1

15

263.0 288.4

where H_s , H_g , and W are final conditions. When the final W is small, 0.01 in the present numerical example, the single stage drying time will be prohibitive. In such cases, multistaging, batch drying, or some other kind of drying equipment must be resorted to.

```
10 ! Example 9.9. Fluidized bed
     dryer
                  R!=R/S, ratio of rat
    INPUT
 20
 es of flow of air and solid
30 H3=.015+.29/R ! =H9
 40 C1=.2434+.225*H3
    INPUT T4!Trialvalue o
                                  f
 50
 60
    GOSUB 2 0 9
    Y1=Y
 78
    T4=1.0001*T4
 80
 90
    GOSUB 200
100
    Y2=Ŷ
    K=.0001*Y1/(Y2-Y1)
110
120
    T4=T4/1 000 1 -K
130
    DISPT4
    IFABS(K/T4) <= .00001 THEN 16
140
    GOTO 60
150
    DISPUSING 170; R, T3, T4, H3,
160
    H4.T5
170 IMAGE DD,X,DDD.D,X,DDD.D,X,.
    DDDD XX.0000XX.DDD
180
    END
200
    ! SR for T4
210 P=EXP(11.9176-7173.9/(T4+389
     55)
220
    H4=18*P/29/(1-P) ! ≃ Hs
230
    T3=T4+900*(H4-H3)/C1!
                                     Τ9
249 Y=-T3+350-(.36*(T4-100)+261)
    /R/C1
T5=.29/(H4-H3)/60!
                                  time
258
                             =
    RETURN
268
270
    END
             Т
                     Η
                            Hs
                                 Time
     Τ.
  R
              s
    145.1
           119.84
  5
                     0730
                           0803
                                   662
                                  .289
.170
    178.1
                     0633
0513
                \frac{74}{61}
  58
    228.
              ģ
                           .0795
           119.53
                     0440
                                  .136
  Й
2
    245.7
```

119.47

119.42

0794

0792

.120

.108

0392

0343



Exhaustair

Drying tower

Cvclone

Product

AIR 120

б

60

210

270

330

PRODUCT

181

241

30

`350 AIR

30

LO

30

(d)

🗲 FEED

Sweeper air

product to collector



Figure 9.14. Spray dryer arrangements and behavior. (a) Spray dryer equipped with spray wheel; straight section LD = 0.5-1.0 (Proctor and Schwartz Inc.). (b) Spray dryer equipped with spray nozzle; straight section LD = 4-5 (Nonhebel and Moss, 1971). (c) Spray dryer for very heat sensitive products; flat bottom, side air ports and air sweeper to cool leaving particles. (d) Distribution of air temperatures in parallel and countercurrent flows (Masters, 1976, p. 18, Fig. 1.5). (e) Droplet-forming action of a spray wheel (Stork-Bowen Engineering Co.).

TABLE 9.16. Performance Data of Spray Dryers

(a) Data of Kröll (1978)

	Moi: Con	sture itent	0		Ai Tempe	r rature
Kind of Stock	In (%)	out (%)	Spray Device	Pattern	In (° C)	out (°C)
Skim milk, $d = 60 \ \mu m$	48-55	4	wheel or	parallel	250	95-100
	50-60	4	nozzle			
			170-200 bar	parallel	250	95-100
Whole milk	50-60	2.5	wheel or	-		
			nozzle			
			100—140 bar	parallel	170-200	
Eggs, whole	74-76	2-4	wheel or			
			nozzle	parallel	140-200	50-80
Eggs, yolks	50-55	2-4	wheel or			
			nozzle	parallel	140-200	50-80
Eggs, whites	87-90	7-9	wheel or			
			nozzle	parallel	140-200	50-80
Coffee, instant, 300 μ m	75-85	3-3.5	nozzle	parallel	270	110
Tea, instant	60	2	nozzle,			
			27 bar	parallel	1 SO-250	
Tomatoes	65-75	3-3.5	wheel	parallel	140-150	
Food yeast	76-78	8	wheel	parallel	300-350	100
Tannin	50-55	4	wheel	parallel	250	S O
PVC emulsion, 90% $>$ 80 μ m	40-70	0.01-0.1	wheel or			
<60 μ m			nozzle or			
			pneumatic	parallel	165-300	
Melamine-urethane-formaldehyde resins	30-50	0	wheel			
			140-160 m/sec	parallel	200-275	65-75
Heavy duty detergents	35-50	8-13	nozzle,			
			30-60 bar	counter	350-400	90-110
Kaolin	35-40	1	wheel	parallel	600	120

(b) Performance of a Dryer **18 ft** Dia by **18 ft** High with a Spray Wheel and a Fan Capacity of 11,000 cfm at the Outlet"

	Air Te	mp (° F)	0/ 10/-/	Francisco
Material	In	out	in Feed	Rate (Ib/hr)
Blood, animal	330	160	6 5	780
Yeast	440	140	86	1080
Zinc sulfate	620	230	5 5	1320
Lignin	400	195	63	910
Aluminum hydroxide	600	130	93	2560
Silica gel	600	170	95	2225
Magnesium carbonate	600	120	9 2	2400
Tanning extract	330	150	4 6	680
Coffee extract A	300	180	70	500
Coffee extract B	500	240	47	735
Magnesium chloride	810	305	53	1140
				(to dihydrate)
Detergent A	450	250	50	660
Detergent B	460	240	63	820
Detergent C	450	250	4 0	340
Manganese sulfate	600	290	50	720
Aluminum sulfate	290	170	70	230
Urea resin A	500	180	60	505
Urea resin <i>B</i>	450	190	70	250
Sodium sulfide	440	150	50	270
Pigment	470	140	73	1750

[#]The fan on this dryer handles about 11,000 cuft/min at outlet conditions. The outlet-air temperature includes cold air in-leakage, and the true temperature drop caused by evaporation must therefore be estimated from a heat balance.

(Bowen Engineering Inc.).

TABLE 9.17. Particle Diameters, Densities, and Energy Requirements

(a) Atomizer Performance

Туре	Size Range (pm)	Power input (kWh/1000 L)
Single fluid nozzle	8-800	0.3-0.5
Pneumatic nozzle	3-250	
Spray wheel	2-550	0.8-1.0
Rotatina CUD	25-950	

(b) Dry Product Size Range

Product	μm
Skim milk	20-250
Coffee	50-600
Eggs	5 - 5 0 0
Egg white	1 - 4 0
Color pigments	1-50
Detergents	20-2000
Ceramics	15-500

(c) Bulk Density of Sprayed Product as Affected by Air Inlet Temperature and Solids Content of Feed"



^a The full lines are against temperature, the dashed ones against concentration: (a) sodium silicate; (b) coffee extract, 22%; (c) water dispersible dye, 19.5%; (d) gelatin.

[Data of Duffie and Marshall, Chem. Eng. Prog. 49, 417 480 (1953)].

accumulation of wet material on the walls; length to diameter ratios of 0.5-1.0 are used in such cases. The downward throw of nozzles permits small diameters but greater depths for a given residence time; L/D ratios of 4-5 or more are used.

ATOMIZATION

Proper atomization of feed is the key to successful spray drying. The three devices of commercial value are pressure nozzles, pneumatic nozzles, and rotating wheels of various designs. Usual pressures employed in nozzles range from 300 to 4000 psi, and

orifice diameters are 0.012-0.15 in. An acceptably narrow range of droplet sizes can be made for a feed of particular physical properties by adjustment of pressure and diameter. Multiple nozzles are used for atomization in large diameter towers. Because of the expense of motive air or steam, pneumatic nozzles are used mostly in small installations such as pilot plants, but they are most suitable for dispersion of stringy materials such as polymers and fibers. The droplet size increases as the motive pressure is lessened, the range of 60-100 psi being usual. The action of a rotating wheel is indicated in Figure 9.14(e). Many different shapes of orifices and vanes are used for feeds of various viscosities, erosiveness, and clogging tendencies. Operating conditions are up to 60,000 lb/hr per atomizer, speeds up to 20,000 rpm, and peripheral speeds of 250400 ft/sec.

The main variables in the operation of atomizers are feed pressure, orifice diameter, flow rate and motive pressure for nozzles and geometry and rotation speed of wheels. Enough is known about these factors to enable prediction of size distribution and throw of droplets in specific equipment. Effects of some atomizer characteristics and other operating variables on spray dryer performance are summarized in Table 9.18. A detailed survey of theory, design and performance of atomizers is made by Masters (1976), but the conclusion is that experience and pilot plant work still are essential guides to selection of atomizers. A clear choice between nozzles and spray wheels is rarely possible and may be arbitrary. Milk dryers in the United States, for example, are equipped with nozzles, but those in Europe usually with spray wheels. Pneumatic nozzles may be favored for polymeric solutions, although data for PVC emulsions in Table 9.16(a) show that spray wheels and pressure nozzles also are used. Both pressure nozzles and spray wheels are shown to be in use for several of the applications of Table 9.16(a).

APPLICATIONS

For direct drying of liquids, slurries, and pastes, drum dryers are the only competition for spray dryers, although fluidized bed dryers sometimes can be adapted to the purpose. Spray dryers are capable of large evaporation rates, 12,000–15,000 lb/hr or so, whereas a 300sqft drum dryer for instance may have a capacity of only 3000 lb/hr. The spherelike sprayed particles often are preferable to drum dryer flakes. Dust control is intrinsic to spray dryer construction but will be an extra for drum dryers. The completely enclosed operation of spray dryers also is an advantage when toxic or noxious materials are handled.

THERMAL EFFICIENCY

Exit air usually is maintained far from saturated with moisture and at a high temperature in order to prevent recondensation of moisture in parallel current operation, with a consequent lowering of thermal efficiency. With steam heating of air the overall efficiency is about 40%. Direct fired dryers may have efficiencies of 80–85% with inlet temperatures of 500–550°C and outlet of 65-70°C. Steam consumption of spray dryers may be 1.2-1.8 lb steam/lb evaporated, but the small unit of Table 9.19(b) is naturally less efficient. A 10% heat loss through the walls of the dryer often is taken for design purposes. Pressure drop in a dryer is 15-50in. of water, depending on duct sizes and the kind of separation equipment used.

DESIGN

The design of spray dryers is based on experience and pilot plant determinations of residence time, air conditions, and air flow rate. Example 9.10 utilizes such data for the sizing of a commercial scale spray dryer.

Variable Increased	Factors Increased	Factors Decreased
Chamber inlet temperature	Feed <i>rate</i> and thus: product rate, particle size (<i>b</i>), product moisture content, chamber wall build-up (<i>a</i>)	bulk density (<i>b</i>)
Chamber outlet temperature	product thermal degradation (<i>a</i>)	feed rate and thus: product rate particle size (b) product moisture content chamber wall build-up
Gas volume rate	feed rate and thus: product rate, particle size (b), product moisture content, chamber wall build-up (a)	residence time
Feed concentration	product rate, bulk density (<i>b</i>), particle size (<i>b</i>)	
Atomizer speed	p	
Atomizer disc diameter		
For stable lattices	bulk density	particle size and thus: product moisture content chamber wall build-up
For unstable lattices	coagulation (<i>a</i>) and thus: particle size, product moisture content, chamber wall build-up	
Atomizer vane depth Atomizer vane number	bulk density (<i>b</i>)	particle size (b) and thus: product moisture content, chamber wall build-up
Atomizer vane radial length		For unstable lattices particle size chamber wall build-up
Feed surface tension Chamber inlet gas humidity	bulk density (b) product moisture content, chamber wall build-up (a)	particle size (<i>b</i>)

TABLE 9.18. Effects of Variables on Operation of Spray Dryers

^a This factor will only occur if a critical value of the variable is exceeded. ^b Not for suspensions. (Nonhebel and Moss, 1971).

The smallest pilot unit supplied by Bowen Engineering has a diameter of 30 in. and straight side of 29 in., employs parallel flow, up to 25 ACFM, 150–1000°F, particle sizes $30-40 \,\mu$ m average, either pneumatic nozzle or spray wheel. The performance of this unit is given in Table 9.19. The magnitude of the "product number" is arrived at by pilot plant work and experience; it increases with increased difficulty of drying or thermal sensitivity or both. Although much useful information can be obtained on this small scale, Williams-Gardner (1971) states that data on at least a 7 ft dia dryer be obtained for final design of large capacity units.

9.11. THEORY OF AIR-WATER INTERACTION IN PACKED TOWERS

The key properties of mixtures of air and water vapor are described in Section 9.1. Here the interactions of air and water in packed towers under steady flow conditions will be analyzed. The primary objectives of such operations may be to humidify or dehumidify the air as needed for particular drying processes or other processes, or to cool process water used for heat transfer elsewhere in the plant. Humidification-dehumidification usually is accomplished in spray towers, whereas cooling towers almost invariably are filled with some type of packing of open structure to improve contacting **but** with minimum pressure drop of air. Analysis of the interaction of air and water involves the making of material and enthalpy balances. These are made over a differential section of the tower shown on Figure 9.15(a) and are subsequently integrated to establish the size of equipment for a given performance. In terms of empirical heat, k_h , and mass, k_m , transfer coefficients, these balances are

$$Gdh = LC_L dT = L dT \tag{9.21}$$

$$= k_m(h_s - h) dz \tag{9.22}$$

$$=k_h(T-T_s)\,dz.\tag{9.23}$$

In Eq. (9.21) the heat capacity of water has been taken as unity. The approximations that are involved in making an enthalpy difference a driving force are discussed for example by Foust et al. (1980). Rearrangement and integration leads to the results

$$NTU = \frac{k_m Z}{L} \tag{9.24}$$

$$= \int_{T_1}^{T_2} \frac{dT}{h_s - h}$$
(9.25)

$$\frac{\mathbf{G}}{\mathbf{L}} \frac{\mathbf{h}_2}{\mathbf{h}_1} \frac{dh}{\mathbf{h}_s - \mathbf{h}}.$$
(9.26)

 TABLE 9.19. Product Numbers and Performance of a 39 x

 29 in. Pilot Plant Spray Dryer

(a)	Product	Numbers	of	Selected	Materials
-----	---------	---------	----	----------	-----------

Material	Product number
l. COLOURS Reactive dyes	5- 6
Pigments	5-11
Dispersed dyes	16-26
2. FOODSTUFFS	
Carbohydrates	14-20
Milk	17
Proteins	16-28
3. PHARMACEUTICALS	
Blood insoluble/soluble	11-22
Hydroxide gels	6-10
Riboflavin	15
Tannin	16-20
4. resins	
Acrylics	1011
Formaldehvde resin	18-28
Polystyrene	12-15
5 055 41100	
J. CERAMICS	
Alumina	1115
Ceramic colours	10

(Bowen Engineering Inc.).

(b) **Performance** of the Pilot Unit as a Function of Product **Number**^a



'Example: For a material with product number = 10 and air inlet temperature of 500°F, the evaporation rate is 53 lb/hr, input Btu/lb evaporated = 1930, and the air outlet temperature is 180°F. (Bowen Engineering).

Both forms of the integral are employed in the literature to define the number of transfer units. The relation between them is

$$k_m Z/G = (L/G)(\text{NTU}). \tag{9.27}$$

The height of a transfer unit is

$$HTU = Z/(NTU) = L/k, = (L/G)(G/k_m).$$
(9.28)

The quantity G/k_m sometimes is called the height of a transfer unit expressed in terms of enthalpy driving force, as in Figure 9.16, for example:

$$G/k_m = (G/L)(HTU). \tag{9.29}$$

Integration of Eq. (9.21) provides the enthalpy balance around one end of the tower,

$$L(T - T_1) + G(h - h_1).$$
(9.30)

Combining Eqs. (9.22) and (9.23) relates the saturation enthalpy and temperature,

$$h_s = h + (k_m/k_h)(T - T_s). \tag{9.31}$$

In Figure 9.15(c), Eq. (9.31) is represented by the line sloping upwards to the left. The few data that apparently exist suggest that the coefficient ratio is a comparatively large number. In the absence of information to the contrary, the ratio commonly is taken infinite, which leads to the conclusion that the liquid film resistance is negligible and that the interface is at the bulk temperature of the water. For a given value of T, therefore, the value of h_s in Eq. (9.25) is found from the equilibrium relation (h_s, T_s) of water and the corresponding value of h from the balance Eq. (9.30). When the coefficient ratio is finite, a more involved approach is needed to find the integrand which will be described.

The equilibrium relation between T_s and h_s is represented on the **psychrometric** charts Figures 9.1 and 9.2, but an analytical representation also is convenient. From Section 9.1,

$$h_s = 0.24T_s + (18/29)(0.45T_s + 1100)[p_s/(1-p_s)], \qquad (9.32)$$

where the vapor pressure is represented by

$$p_s = \exp[11.9176 - 7173.9/(T_s + 389.5)]. \tag{9.33}$$

Over the limited ranges of temperature that normally prevail in cooling towers a quadratic fit to the data,

$$h_s = a + bT_s + cT_s^2$$

may be adequate. Then an analytical integration becomes possible for the case of infinite k_m/k_h . This is done by Foust et al. (1980) for example.

The Cooling Tower Institute (1967) standardized their work in terms of a Chebyshev numerical integration of Eq. (9.25). In this method, integrands are evaluated at four temperatures in the interval, namely,

$$T_{2} + 0.1(T_{2} - T_{1}), \quad \text{corresponding integrand } I_{1},$$

$$T_{2} + 0.4(T_{2} - T_{1}), \quad \text{corresponding integrand } I_{2},$$

$$T_{1} - 0.4(T_{2} - T_{1}), \quad \text{corresponding integrand } I_{3},$$

$$T_{1} - 0.1(T_{2} - T_{1}), \quad \text{corresponding integrand } I_{4}.$$
(9.34)

Then the integral is

$$\int_{T_1}^{T_2} \frac{dT}{h_s - h} = 0.25(T_2 - T_1)(I_1 + I_2 + I_3 + I_4).$$
(9.35)

EXAMPLE 9.10

Siziig a Spray Dryer on the Basis of Pilot Plant Data

Feed to a spray dryer contains 20% solids and is to be dried to 5% moisture at the rate of 500 lb/hr of product. Pilot plant data show that a residence time of 6 sec is needed with inlet air of 230°F, H = 0.008 lb/lb, and exit at 100°F. Ambient air is at 70°F and is heated with steam. Enthalpy loss to the surroundings is 10% of the heat load on the steam heater. The vessel is to have a 60° cone. Air rate and vessel dimensions will be found.

Enthalpy, humidity, and temperatures of the air are read off the psychrometric chart and recorded on the sketch.



Enthalpy loss of air is $0.1(69.8 - 28.0) \approx 4.2$ Btu/lb. Exit enthalpy of air is h = 69.8 - 4.2 = 65.6.

At 100°F and this enthalpy, other properties are read off the psychrometric chart as

$$H = 0.0375 \text{ lb/lb},$$

 $V = 14.9 \text{ cuft/lb}.$

Air rate is

$$A = \frac{1900 - 25}{0.0375 - 0.008} = 63,559 \text{ lb/hr}$$

$$\rightarrow \frac{63,559}{3600} \left(\frac{17.6 + 14.9}{2}\right) = 287 \text{ cfs.}$$

With a residence time of 6 sec, the dryer volume is

$$V_d = 287(6) = 1721.4$$
 cuft.

Make the straight side four times the diameter and the cone 60":

1721.4 =
$$4D(\pi D^2/4) + \frac{0.866\pi D^3}{12} = 3.3683D^3$$
,
∴ $D = 8.0$ ft.

When $k_m/k_h \rightarrow \infty$, evaluation of the integrands is straightforward. When the coefficient ratio is finite and known, this procedure may be followed:

- **1.** For each of the four values of T, find h from Eq. (9.30).
- 2. Eliminate h_s between Eqs. (9.31) and (9.32) with the result

$$\begin{aligned} h + (k_m/k_h)(T - T_s) \\ &= 0.24T_s + (18/29)(0.45T_s + 1100)[p_s/(1 - p_s)]. \end{aligned}$$
(9.36)

- **3.** Substitution of Eq. (9.33) into (9.36) will result in an equation that has T_s as the only unknown. This is solved for with the Newton-Raphson method.
- **4.** Substitution of this value of T_s back into Eq. (9.31) will evaluate h_{s} .
- 5. The integrand $1/(h_s h)$ now may be evaluated at each temperature and the integration performed with Eq. (9.35).

Example 9.11 employs this method for finding the number of transfer units as a function of liquid to gas ratio, both with finite and infinite values of k_m/k_h . The computer programs for the solution of this example are short but highly desirable. Graphical methods have been widely used and are described for example by Foust et al. (1980).

TOWER HEIGHT

The information that is ultimately needed about a cooling tower design is the height of packing for a prescribed performance. This equals the product of the number of transfer units by the height of each one,

$$\mathbf{Z} = (\mathrm{NTU})(\mathrm{HTU}). \tag{9.37}$$

Some HTU data for cooling tower packing have been published, for example, those summarized on Figure 9.16. Other data appear in the additional literature cited for this chapter. Several kinds of tower fill made of redwood slats are illustrated in Figure 9.17. The numbers N of such decks corresponding to particular NTUs and (L/G)s are given by the equation

$$N = \frac{[(NTU) - 0.07](L/G)^{b}}{(9.38)}$$

Values of *a* and *b* are given for each type of fill with Figure 9.17. These data are stated to be for 120°F inlet water. Although the authors state that corrections should be estimated for other temperatures, they do not indicate how this is to be done. For example, with deck type C, NTU = 2 and L/G = 1.2: N = $(2 - 0.07)(1.2)^{0.60}/0.092 = 23.4$ decks, or a total of 31.2 ft since the deck spacing is 16 in. The data of Figure 9.16 are used in Example 9.11.



Figure 9.15. Relations in a packed continuous flow air-water contactor. (a) Sketch of the tower with differential zone over which the enthalpy and material balances are made. (b) Showing equilibrium and operating lines from which the integrand $1/(h_s - h)$ can be found as a function of liquid temperature T. (c) Showing interfacial conditions as determined by the coefficient ratio k_m/k_h ; when this value is large, interfacial and saturation temperatures are identical.



Figure 9.16. Data of heights of transfer units of packings characterized by the specific surface $a_d(\operatorname{sqft}/\operatorname{cuft})$. The ordinate is $G/k_m = Z/\int dh/(h_s h)$, which is related to the form of NTU used in this chapter by

HTU =
$$Z/NTU = Z/\int \frac{dT}{h_s - h} = L/k_m = (G/k_m)(L/G)$$

The equation of the London line is equivalent to

 $HTU = 5.51(L/G)^{0.59}$

(Sherwood et al., 1975).

9.12. COOLING TOWERS

Cooling of water in process plants is accomplished most economically on a large scale by contacting it with air in packed towers. For reasons of economy, the tower fill is of a highly open structure. Efficient ring and structured packings of the sort used for distillation and other mass transfer processes are too expensive and exert too high a power load on the fans. Standard cooling tower practice allows a maximum of 2in. of water pressure drop of the air. Water loadings range 500-2000 lb/(hr)(sqft) or 1-4 gpm/sqft. Gas loadings range 1300–1800 lb/(hr)(sqft) or between 300 and 400 ft/min. The liquid to gas ratio L/G normally is in the range 0.75-1.50 and the number of transfer units or the tower characteristic, HTU = $k_m Z/L$, vary from 0.5 to 2.50.

The most common fill is of wooden slats of rectangular or triangular cross section arranged as in Figure 9.17. Corrugated sheets of asbestos-concrete have some application and also PVC construction unless the temperatures are above 160°F.

Fan power consumption is the major operating cost and can be counterbalanced in part by greater investment in natural draft construction. In the majority of process applications, fan-operated towers are preferred. Very large installations such as those in power plants employ chimney assisted natural draft installations. A limited use of atmospheric towers is made in areas where power costs are especially high.

The main types of cooling towers are represented on Figure 9.18. Their chief characteristics and some pros and cons will be discussed in order.

EXAMPLE 9.11 Sizing of a Cooling Tower: Number of Transfer Units and Height of Packing

Water is to be cooled from 110 to 75°F by contact with air that enters countercurrently at 90°F with a **dewpoint** of 60°F. The data of London et al. (1940) of Figure 9.16 for height of transfer unit are applicable. Calculations will be made for two values of the coefficient ratio k_m/k_h , namely, 25 and ∞ Btu/(°F) (lb dry air), of Eq. (9.31). The effect of the ratio of liquid to gas rates, *L/G*, will be explored.



The maximum allowable L/G corresponds to equilibrium between exit air and entering water at 110. The saturation enthalpy at 110°F is 92, so that Eq. (9.30) becomes

$$\left(\frac{L}{G}\right)_{\max} = \frac{92 - 27}{110 - 75} = 1.857.$$

The several trials will be made at L/G = (0.6, 1.0, 1.4, 1.7).

The applicable equations with numerical substitutions are listed here and incorporated in the computer program for solution of this problem [Eqs. (9.30)-(9.33)]:

$$h = 27 + (L/G)(T - 75),$$

$$h_s = h + 25(T - 75),$$

$$h_s = 0.24T + (18/29)(0.45T + 1100)P_s/(1 - P_s),$$

$$P_s = \exp[11.9176 - 7173.9/(T_s + 389.5)].$$

When $k_m/k_h \rightarrow \infty$, T_s in Eq. (9.33) is replaced by T.

The four temperatures at which the integrands are evaluated for the Chebyshev integration are found with Eq. (9.34) and tabulated in the calculation summary following.

Equations (9.30) and (9.31) are solved simultaneously for h and h_s with the aid of the Newton-Raphson method as used in the computer program; the integrands are evaluated and the integration are completed with Eq. (9.35).

The number of transfer units is sensitive to the value of L/G, but the effect of k_m/k_h is more modest, at least over the high range used; data for this ratio do not appear to be prominently recorded. Figure 9.16 shows a wide range of heights of transfer units for the different kinds of packings, here characterized by the surface a_d (sqft/cuft) and substantial variation with L/G. The last line of the calculation summary shows variation of the tower height with L/G

Data of London et al. (1940) of Figure 9.16:

$$(G/L)(HTU) = 5.51(G/L)^{0.4}$$

or

$$HTU = 5.51(L/G)^{0.59}$$

Tower height:

Z = (HTU)(NTU).

For several values of *L/G*:

Evaluation of interfacial temp and the NTU for L/G = 1 with $k_m/k_h \approx 25$:

т	h	T _s	$1/(h_s - h)$
78.5	30.5	78.099	0.0864
89	41	88. 517	0.0709
96	48	95.400	0.0575
106.5	58.5	105. 581	0. 0385

0. 2533

$$\therefore$$
 NTU = (110 - 75(0.2533)/4 = 2.217.

For other values of L/G:

		1/(h _s - h)				
т	h	L/G=0.0	6 1	1.4	1.7	
78.5	30.5	0. 0751	0.0864	0.0943	0. 1043	
89	41	0.0518	0.0709	0.1167	0. 2200	
96	48	0.0398	0.0575	0.1089	0.3120	
106.5	58 . 5	0. 0265	0.0385	0.0724	0. 1987	
		-	-	-	-	
		0. 1933	0. 2533	0.3923	0.8350	
NTU	→	1.691	2.217	3. 433	7.306	

With $k_m/k_h \rightarrow \infty$:

			1/(h _s — h)		
т	h	L/G=0.6	1	1.4	1.7	
78.5	30. 5	0.0725	0.0807	0.90	0. 1006	
89	41	0.0494	0.0683	0.1107	0.2070	
96	48	0. 0376	0.0549	0.1020	0. 2854	
106.5	106.5	0. 0248	0.0361	0.0663	0.1778	
		-	-	-	-	
		0. 1844	0.2400	0.3700	0.7708	
NTU -	→	1.613	2.100	3. 238	8.745	
Z-	→	6.58	11. 57	21.76	50.86	

10 / Example 9.11 with Km/Kh = 25

20 READ T1,L 30 DATA 106.5,1.7

40 H2=27+L*(T1-75)

50 INPUT T ! = Ts

- 60 GOSUB 180
- 70 Y1=Y
- 80 T=1.0001*T 90 GOSUB 180

```
EXAMPLE 9.11—(continued)
110 K=.0001*Y1/(Y2-Y1)
120 T=T/1.0001-K
    DISP T
130
140
   IF ABS(K/T) <= 00001 THEN 160
150
    GOTO 60
160
    DISP T,H,1/(H-H2)
170
    END
      SR FOR T, H
180
    P=EXP(11.9176-7173.9/(T+389
190
    5 > >
200
    M=18*P/29/(1-P)
    H=.24*T1+(.45*T1+1100)*M
210
220
    H1=H2-25*(T-T1)
230
    Y = H1 - H ! = 0 ?
```



Figure 9.17. Kinds of fill made of redwood slats for cooling towers, and factors for determining the required number of decks with inlet water at 120° F (*Cheremisinoff and Cheremisinoff*, 1981).

- 240 RETURN 250 END Example 9.11 with infinite 10 Km/Kh 20 L=1.4 INPUT TI 30 H2=27+L*(T1-25) 40 P=EXP(11 9176-7173.9/(T1+389 50 500M=18*P/29/(1-P) ! = Hs 69. H=.24*T1+(.45*T1+1100)*M 70 80 DISP 1/(H-H2) 90 GOTO 30 100 END
- a. Atmospheric towers are effective when prevailing wind velocities are 5 miles/hr or more. For access to the wind they are narrow but long, lengths of 2000 ft having been constructed. Water drift losses are relatively large. The savings because of elimination of tall chimney or fan power is counterbalanced by increased size because of less efficient cross flow and variations in wind velocity.
- **b.** Chimney assisted natural draft towers also eliminate fans. Most of the structure is the chimney, the fill occupying only 10–12% of the tower height at the bottom. The temperature and humidity of the air increase as the air flows upward so that its buoyancy increases and results in rapid movement through the chimney. Smaller units are made as circular cylinders since these can be built rapidly. The hyperboloidal shape has greater strength for a given wall thickness. In towers as large as 25Oft dia and 450ft high, wall thicknesses of 5–8 in. of reinforced concrete are adequate. The enlarged cross section at the top converts some kinetic energy into pressure energy which assists in dispelling the exit humid air into the atmosphere.

The ratio of base diameter to height is 0.75-0.85, the ratio of throat and base diameters is 0.55-0.65, and the ratio of vertical depth of air opening to base diameter is 0.1-0.12. Air velocity through the tower is 3-6 ft/sec, water flow rates range from 600 to 1800 lb/(hr)(sqft). Two towers each 375 ft high are able to service a 500 MW power plant. Natural draft towers are uneconomical below heights of 70 ft. The upper limit is imposed principally by environmental visual considerations; towers 500 ft high are in existence. A cost comparison is made with item d.

- c. Hyperbolic fan assisted towers can have as much as three times the capacity of the same size natural draft towers. The fans provide greater control than the natural draft systems; for example, they may be turned on only at peak loads. Rules of thumb cited by Cheremisinoff and Cheremisinoff (1981) for relative sizing is that fan assisted hyperbolic towers may have diameters 2/3 and heights 1/2 those of purely natural draft designs.
- d. Countercurrent-induced draft construction is the most widely used type in process industries. Mechanical draft is capable of a greater degree of control than natural draft and such towers are able in some cases to cool the water within 2°F of the wet bulb temperature of the air. The elevated fan location introduces some structural and noise problems. The flow of air is quite uniform across the cross section and its discharge is positive and at high velocity so that there is little backflow of humid air into the tower. A cost comparison (dated 1978) with hyperbolic towers is made by Singham (1983, Sec. 3.12.4.1). The case is for a water rate of 6.1 m³/sec, cooling range of 8.5°C, approach of 10°C, and wet bulb of 17°C. The cost of the natural draft tower





Figure 9.18. Main types of cooling towers. (a) Atmospheric, dependent on wind velocity. (b) Hyperbolic stack natural draft. (c) Hyperbolic assisted with forced draft fans. (d) Counterflow-induced draft. (e) Crossflow-induced draft. (f) Forced draft. (g) Induced draft with surface precooler for very hot water; also called wet/dry tower. [(b)-(e) f_{jrom} Cheremisinoff and Cheremisinoff, 1981).



Figure 9.19. Typical cooling tower performance curves (Cheremisinoff and Cheremisinoff, 1981).

TABLE 9.20. Selected Data Required with Bids of Cooling Towers

A. Cooling Tower 1. Number of cells 2. Cell dimensions, ft, in. 3. Tower length, ft, in. 4. Tower width, ft. in. 5. Tower height, ft, in. 6. Casing, material and dimensions	3. Type and manufacturer
7. Structure, material and dimensions 8. Fill decks, material and dimensions 9. Partitions and baffles, materials and dimensions 10. Drift eliminators, material and dimensions 11. Fan stacks, material and dimensions 12. Fan deck, material and dimensions	H. Distribution System 1. Number and size of inlet flanges 2. Height of water inlet above curb, ft. in. 3. Header material 4. Lateral material 5. Nozzle, or downspout material
13. Louvers, material and dimensions 14. Board feet of fill 15. Board feet total tower 16. Height of fan stacks, ft, in. 17. Post extension below curb, ft. in. 18. Total shipping weight, lb 19. Total operating weight, lb 19. Fans 10. Number of units 10. Total operating weight, lb 10. Total operating weight, lb 10. Total operating weight, l	J. Design Performance 1. Pumping head from top of basin curb, ft 2. Spray loss, max % 3. Evaporation loss, max % 4. Fill wetted surface, ft ² 5. Total wetted surface, ft ² 6. Effective splash surface, ft ³ 7. Effective cooling volume, ft ³ (from eliminators to water level) 8. Air volume per fan, cfm 9. Static pressure, inches of water 10. Output horsepower/motor/(turbine) 11. Tower loading, gpm/ft ²
7. rpm 8. Tip speed, fpm 9. Mechanical efficiency, % 10. Static efficiency, % 11. Weight, Ib C. Motors 1. Number of units	K. Drawings and Performance Curves 1. Tower outline elevation 2. Foundation outline 3. Fill rack details 4. Drift eliminator details 5. Tower sheeting arrangement 6. A series of guaranteed performance curves within limits of CTI

was 1.2 M pounds and that of the mechanical draft was 0.75 M pounds, but the fan power was 775 kW. The opinion was expressed that mechanical draft towers are more economical at water rates below 1.25 m^3 /sec (19,800 gpm).

- e. Crossflow induced draft offer less resistance to air flow and can operate at higher velocities, which means that less power and smaller cell sizes are needed than for counterflows. The shorter travel path of the air makes them less efficient thermally. The cross flow towers are made wider and less high, consequently with some saving in water pumping cost.
- f. Forced draft towers locate the fans near ground level which requires simpler support structures and possibly lower noise levels. A large space must be provided at the bottom as air inlet. Air distribution is **poor** because it must make a 90" turn. The humid air is discharged at low velocity from the top of the tower and tends to return to the tower, but at the same time the drift loss of water is less. The pressure drop is on the discharge side of the fan which is less power-demanding than that on the intake side of induced draft towers.
- g. Wet-dry towers employ heat transfer surface as well as direct contact between water and air. Air coolers **by** themselves are used widely for removal of sensible heat from cooling water on a comparatively small scale when cooling tower capacity is limited. Since dry towers cost about twice as much as wet ones, combinations of wet and dry sometimes are applied, particularly when the water temperatures are high, of the order of 160°F, so that evaporation losses are prohibitive and the plumes are environmentally undesirable. The warm water flows first through tubes across which air is passed and then enters a conventional

REFERENCES

Drying

- W.L. Badger and J.T. Banchero, Introduction to Chemical Engineering, McGraw-Hill, New York, 1955.
- 2. C.W. Hall, Dictionary of Drying Dekker, New York, 1979.
- 3. R.B. Keey, *Drying Principles and Practice*, Pergamon, New York, 1972. 4. R.B. Keey, *Introduction to Industrial Drying Operations*, Pergamon,
- New York, 1978.
 5. K. Kröll, Trockner und Trocknungsverfahren, Springer-Verlag, Berlin, 1978.
- P.Y. McCormick, Drying, in Encyclopedia of Chemical Technology, Wiley, New York, 1979, Vol. 8, pp. 75-113.
- 7. K. Masters, Spray Drying George Godwin, London, 1976.
- A.S. Mujumdar (Ed.), Advances in Drying. Hemisphere, New York, 1980-1984, 3 vols.
- 9. G. Nonhebel and A.A.H. Moss, Drying of Solids in the Chemical Industry, Butterworths, London, 1971.
- R.E. Peck, Drying solids, in Encyclopedia of Chemical Processing and Design, Dekker, New York, 1983, Vol. 17, pp. 1-29.
- 11. E.U. Schlinder, Dryers, in *Heat Exchanger Design Handbook*, Hemisphere, New York, 1983, Sec. 3.13.
- 12. G.A. Schurr, Solids drying, in *Chemical Engineers Handbook*, McGraw-Hill, New York, 1984, pp. 20.4–20.8.
- T.H. Wentz and J.R. Thygeson, Drying of wet solids, in Handbook of Separation Techniques for Chemical Engineers, (Schweitzer, Ed.), McGraw-Hill, New York, 1979.

packed section where it is cooled further by direct contact with air. Separate dampers for air to the dry and wet sections can throw greater load on the wet section in summer months.

WATER FACTORS

Evaporation losses are about 1% of the circulation for every 10°F of cooling range. Windage or drift losses are 0.3-1.0% for natural draft towers and 0.1-0.3% for mechanical draft. Usually the salt content of the circulating water is limited to 3-7 times that of the makeup. Blowdown of 2.5-3% of the circulation accordingly is needed to maintain the limiting salt concentration.

TESTING AND ACCEPTANCE

At the time of completion of an installation, the water and air conditions and the loads may not be exactly the same as those of the design specification. Acceptance tests performed then must be analyzed to determine if the performance is equivalent to that under the design specifications. Such tests usually are performed in accordance with recommendations of the Cooling Tower Institute.

The supplier generally provides a set of performance curves covering a modest range of variation from the design condition, of which Figure 9.19 is a sample. Some of the data commonly required with bids of cooling tower equipment are listed in Table 9.20, which is excerpted from a lo-page example of a cooling tower requisition by Cheremisinoff and Cheremisinoff (1981).

14. A. Williams-Gardner, Industrial Drying, Leonard Hill, Glasgow, 1971.

Cooling Towers

- 1. N.P. Cheremisinoff and P.N. Cheremisinoff, Cooling Towers: Selection, Design and Practice, Ann Arbor Science, Ann Arbor, MI, 1981.
- 2. Cooling Tower Institute, Performance Curves, CTI, Spring, TX, 1967.
- 3. A.S. Foust et al., Principles of Unit Operations, Wiley, New York, 1980.
- 4. D.Q. Kern, Process Heat Transfer, McGraw-Hill, New York, 1950.
- T.K. Sherwood, R.L. Pigford, and CR. Wilke, *Mass Transfer*, McGraw-Hill, New York, 1975.
- J.R. Singham, Cooling towers, in *Heat Exchanger Design Handbook*, Hemisphere, New York, 1983, Sec. 3.12.

Data on Performance of Cooling Tower Packing

- 1. Hayashi, Hirai, and Okubo, Heat Transfer Jpn. Res. 2(2) 1-6 (1973).
- 2. Kelly and Swenson, *Chem Eng. Prog.* 52, 263 (1956), cited in Figure 9.16.
- 3. Lichtenstein, Trans. ASME 66, 779 (1943), cited in Figure 9.16.
- 4. London, Mason, and Boelter, Trans. ASME 62, 41 (1940). cited in Figure 9.16.
- Lowe and Christie, Proceedings, International Heat Transfer Conference, Boulder, CO, 1961, Part V, pp. 933-950.
- 6. Simpson and Sherwood, Refrig. Eng. 52, 535 (1946), cited in Figure 9.16.
- 7. Tezuka, Heat Transfer Jpn. Res. 2(3), 40-52 (1973).

gitation is a means whereby mixing of phases can be accomplished and by which mass and heat transfer can be enhanced between phases or with external surfaces. In its most general sense, the process of mixing is concerned with all combinations of phases of which the most frequently occurring ones are

1. gases with gases.

- 2. gases into liquids: dispersion.
- 3. gases with granular solids: fluidization, pneumatic conveying, drying,
- 4. liquids into gases: spraying and atomization.
- 5. liquids with liquids: dissolution, emulsification, dispersion.
- 6. liquids with granular solids: suspension.
- 7. pastes with each other and with solids.
- 8. solids with solids: mixing of powders.

Interaction of gases, liquids, and solids also may take place, as in hydrogenation of liquids in the presence of a *slurried solid catalyst where the gas must be dispersed as bubbles* and the solid particles must be kept in suspension.

Three of the processes involving liquids, numbers 2, 5,

and 6, employ the same kind of equipment; namely, tanks in which the |iquid is circulated and subjected to a certain amount of shear. This kind of equipment has been studied most extensive/y. Although some unusual cases of liquid mixing may require pilot p/ant testing, genera/ rules have been developed with which mixing equipment can be designed somewhat satisfactorily. This topic will be emphasized in this chapter.

The other mixing operations of the list require individual kinds of equipment whose design in some cases is less quantified and is based largely on experience and pilot plant work. Typical equipment for such purposes will be illustrated later in this chapter. Phase mixing equipment which accomplishes primarily mass transfer between phases, such as distillation and extraction towers, a/so are covered elsewhere. Stirred reactors are discussed in Chapter 77.

Circulation and shear of the liquid in a vessel can be accomplished with external pumps and appropriate location of suction and discharge nozzles, but a satisfactory combination of vertical and lateral flows is obtained more economically by internal impellers, baffles, and draft tubes. Some genera/ statements about dimensions, proportions, and internals of a liquid mixing vessel can be made.

10.1. A BASIC STIRRED TANK DESIGN

The dimensions of the liquid content of a vessel and the dimensions and arrangement of impellers, baffles and other internals are factors that influence the amount of energy required for achieving a needed amount of agitation or quality of mixing. The internal arrangements depend on the objectives of the operation: whether it is to maintain homogeneity of a reacting mixture or to keep a solid suspended or a gas dispersed or to enhance heat or mass transfer. A basic range of design factors, however, can be defined to cover the majority of cases, for example as in Figure 10.1.

THE VESSEL

A dished bottom requires less power than a flat one. When a single impeller is to be used, a liquid level equal to the diameter is optimum, with the impeller located at the center for an all-liquid system. Economic and manufacturing considerations, however, often dictate higher ratios of depth to diameter.

BAFFLES

Except at very high Reynolds numbers, baffles are needed to prevent vortexing and rotation of the liquid mass as a whole. A baffle width one-twelfth the tank diameter, $w = D_i/12$; a length extending from one half the impeller diameter, d/2, from the tangent line at the bottom to the liquid level, but sometimes terminated just above the level of the eye of the uppermost impeller. When solids are present or when a heat transfer jacket is used, the baffle width. Four radial baffles at equal spacing are standard; six are only slightly more effective, and three appreciably less so. When the mixer shaft is located off center (one-fourth to

one-half the tank radius), the resulting flow pattern has less swirl, and baffles may not be needed, particularly at low viscosities.

DRAFT TUBES

A draft tube is a cylindrical housing around and slightly larger in diameter than the impeller. Its height may be little more than the diameter of the impeller or it may extend the full depth of the liquid, depending on the flow pattern that is required. Usually draft tubes are used with axial impellers to direct suction and discharge streams. An impeller-draft tube system behaves as an axial flow pump of somewhat low efficiency. Its top to bottom circulation behavior is of particular value in deep tanks for suspension of solids and for dispersion of gases. About a dozen applications are illustrated by Sterbacek and Tausk (1965, pp. 283ff) and a chapter is devoted to their use by Oldshue (1983, 469ff).

IMPELLER TYPES

A basic classification is into those that circulate the liquid axially and those that achieve primarily radial circulation. Some of the many shapes that are being used will be described shortly.

IMPELLER SIZE

This depends on the kind of impeller and operating conditions described by the Reynolds, Froude, and Power numbers as well as individual characteristics whose effects have been correlated. For the popular turbine impeller, the ratio of diameters of impeller and vessel falls in the range, d/D, = 0.3-0.6, the lower values at high rpm, in gas dispersion, for example.



Figure 10.1. A basic stirred tank design, not to scale, showing a lower radial impeller and an upper axial impeller housed in a draft tube. Four equally spaced baffles are standard. H = height of liquid level, $D_t =$ tank diameter, d = impeller diameter. For radial impellers, $0.3 \le d/D_t \le 0.6$.

IMPELLER SPEED

With commercially available motors and speed reducers, standard speeds are 37, 45, 56, 68, 84, 100, 125, 155, 190, and 320rpm. Power requirements usually are not great enough to justify the use of **continously** adjustable steam turbine drives. Two-speed **drives** may be required when starting torques are high, as with a settled slurry.

IMPELLER LOCATION

Expert opinions differ somewhat on this factor. As a first approximation, the impeller can be placed at 1/6 the liquid level off the bottom. In some cases there is provision for changing the position of the impeller on the shaft. For off-bottom suspension of solids, an impeller location of 1/3 the impeller diameter off the bottom may be satisfactory. Criteria developed by Dickey (1984) are based on the viscosity of the liquid and the ratio of the liquid depth to the tank diameter, h/D. Whether one or two impellers are needed and their distances above the bottom of the tank are identified in this table:

Viscosity	Maximum	Number of	Impeller	Clearance	
[cP (Pa sec)]	h/D _t	Impellers	Lower	Upper	
<25,000 (<25)	1.4	1	h/3		
<25,000 (<25)	2.1	2	D,/3	(2/3)h	
>25,000 (>25)	0.8	1	h/3		
>25,000 (>25)	1.6	2	D _t /3	(2/3)h	

Another rule is that a second impeller is needed when the liquid must travel more than 4 ft before deflection.

Side entering propellors are placed 18-24 in. above a flat tank floor with the shaft horizontal and at a 10" horizontal angle with the centerline of the tank; such mixers are used only for viscosities below 500 cP or so.

In dispersing gases, the gas should be fed directly below the impeller or at the periphery of the impeller. Such arrangements also are desirable for mixing liquids.

10.2. KINDS OF IMPELLERS

A rotating impeller in a **fluid** imparts flow and shear to it, the shear resulting from the flow of one portion of the fluid past another. Limiting cases of **flow** are in the axial or radial directions so that impellers are classified conveniently according to which of these flows is dominant. By reason of reflections from vessel surfaces and obstruction by baffles and other internals, however, flow patterns in most cases are mixed. When a close approach to axial flow is particularly desirable, as for suspension of the solids of a slurry, the impeller may be housed in a draft tube; and when radial flow is needed, a shrouded turbine consisting of a rotor and a **stator** may be employed.

Because the performance of a particular shape of impeller usually cannot be predicted quantitatively, impeller design is largely an exercise of judgment so a considerable variety has been put forth by various manufacturers. A few common types are illustrated on Figure 10.2 and are described as follows:

- a. The three-bladed mixing propeller is modelled on the marine propeller but has a pitch selected for maximum turbulence. They are used at relatively high speeds (up to 1800 rpm) with low viscosity fluids, up to about 4000 cP. Many versions are available: with cutout or perforated blades for shredding and breaking up lumps, with sawtooth edges as on Figure 10.2(g) for cutting and tearing action, and with other than three blades. The stabilizing ring shown in the illustration sometimes is included to minimize shaft flutter and vibration particularly at low liquid levels.
- **b.** The turbine with flat vertical blades extending to the shaft is suited to the vast majority of mixing duties up to 100,000 **cP** or so at high pumping capacity. The simple geometry of this design and of the turbines of Figures 10.2(c) and (d) has inspired extensive testing so that prediction of their performance is on a more rational basis than that of any other kind of impeller.
- c. The horizontal plate to which the impeller blades of this turbine are attached has a stabilizing effect. Backward curved blades may be used for the same reason as for type e.
- **d.** Turbine with blades are inclined **45°** (usually). Constructions with two to eight blades are used, six being most common. Combined axial and radial flow are achieved. Especially effective for heat exchange with vessel walls or internal coils.
- e. Curved blade turbines effectively disperse fibrous materials without fouling. The swept back blades have a lower starting torque than straight ones, which is important when starting up settled slurries.
- f. Shrouded turbines consisting of a rotor and a **stator** ensure a high degree of radial flow and shearing action, and are well adapted to emulsification and dispersion.
- g. Flat plate impellers with sawtooth edges are suited to emulsification and dispersion. Since the shearing action is localized, baffles are not required. Propellers and turbines also are sometimes provided with sawtooth edges to improve shear.
- **h.** Cage beaters impart a cutting and beating action. Usually they are mounted on the same shaft with a standard propeller. More violent action may be obtained with **spined** blades.







(c)







(e)











(j)





(h)



(k)



(i)



290 MIXING AND AGITATION

- i. Anchor paddles fit the contour of the container, prevent sticking of pasty materials, and promote good heat transfer with the wall.
- **J.** Gate paddles are used in wide, shallow tanks and for materials of high viscosity when low shear is adequate. Shaft speeds are low. Some designs include hinged scrapers to clean the sides and bottom of the tank.
- k. Hollow shaft and hollow impeller assemblies are operated at high tip speeds for recirculating gases. The gas enters the shaft above the liquid level and is expelled centrifugally at the impeller. Circulation rates are relatively low, but satisfactory for some hydrogenations for instance.
- This arrangement of a shrouded screw impeller and heat exchange coil for viscous liquids is perhaps representative of the many designs that serve special applications in chemical processing.

10.3. CHARACTERIZATION OF MIXING QUALITY

Agitation and mixing may be performed with several objectives:

1. Blending of miscible liquids.

- 2. Dispersion of immiscible liquids.
- 3. Dispersion of gases in liquids.
- 4. Suspension of solid particles in a slurry.
- 5. Enhancement of heat exchange between the fluid and the boundary of a container.
- 6. Enhancement of mass transfer between dispersed phases.

When the ultimate objective of these operations is the carrying out of a chemical reaction, the achieved specific rate is a suitable measure of the quality of the mixing. Similarly the achieved heat transfer or mass transfer coefficients are measures of their respective operations. These aspects of the subject are covered in other appropriate sections of this book. Here other criteria will be considered.

The uniformity of a multiphase mixture can be measured by sampling of several regions in the agitated mixture. The time to bring composition or some property within a specified range (say within 95 or 99% of uniformity) or spread in values-which is the blend time-may be taken as a measure of mixing performance. Various kinds of tracer techniques may be employed, for example:

- 1. A dye is introduced and the time for attainment of uniform color is noted.
- 2. A concentrated salt solution is added as tracer and the measured electrical conductivity tells when the composition is uniform.
- The color change of an indicator when neutralization is complete when injection of an acid or base tracer is employed.
- 4. The residence time distribution is measured by monitoring the outlet concentration of an inert tracer that can be analyzed for accuracy. The shape of response curve is compared with that of a thoroughly (ideally) mixed tank.

The last of these methods has been applied particularly to chemical reaction vessels. It is covered in detail in Chapter 17. In most cases, however, the **RTDs** have not been correlated with impeller characteristics or other mixing parameters. Largely this also is true of most mixing investigations, but Figure 10.3 is an uncommon example of correlation of blend time in terms of Reynolds number for the popular pitched blade turbine impeller. As expected, the blend time levels off beyond a certain mixing intensity, in this case beyond Reynolds numbers of 30,000 or so. The acid-base indicator technique was used. Other details of the test work and the scatter of the data are not revealed in the published information. Another practical solution of the problem is typified by Table 10.1 which relates blend time to power input to



Figure 10.3. Dimensionless blend time as a function of Reynolds number for pitched turbine impellers with six blades whose W/D = 1/5.66 [Dickey and Fenic, Chem. Eng. 145, (5 Jan. 1976)].

vessels of different sizes and liquids of various viscosities. A review of the literature on blend times with turbine impellers has been made by Brennan and Lehrer [*Trans. Inst. Chem. Eng.* 54, 139-152 (1975)], who also did some work in the range $10^4 < N_{\rm Re} < 10^5$ but did not achieve a particularly useable correlation.

An impeller in a tank functions as a pump that delivers a certain volumetric rate at each rotational speed and corresponding power input. The power input is influenced also by the geometry of the equipment and the properties of the fluid. The flow pattern and the degree of turbulence are key aspects of the quality of mixing. Basic impeller actions are either axial or radial, but, as Figure 10.4 shows, radial action results in some axial movement by reason of deflection from the vessel walls and baffles. Baffles contribute to turbulence by preventing swirl of the contents as a whole and elimination of vortexes; offset location of the impeller has similar effects but on a reduced scale.

Power input and other factors are interrelated in terms of certain dimensionless groups. The most pertinent ones are, in common units:

$N_{\rm Re}=10.75Nd^2S/\mu,$	Reynolds number,	(10.1)
$N_P = 1.523(10^{13})P/N^3 d^5 S,$	Power number,	(10.2)
$N_Q = 1.037(10^5)Q/Nd^3$,	Flow number,	(10.3)
$t_b N$,	Dimensionless blend time,	(10.4)

TABLE 10.1. Blending Data for Four-Bladed 45° Turbines"

V-→		1000 ga	1		5000 ga	1	1	0, 00 0 g	al	2	0.000 ga	ıl
₩ ↓ () →	6	12	30	6	12	30	6	12	30	6	12	30
100			\odot	3	2	1	3	2	1	71	5	2
250		\bigcirc	\bigcirc	3	2	l	5	3	1 1	10	71	3
500	(3 †)		(†)	3	2	1	71	5	2	15	10	5
1000	11	(21)		5	3	112,	15	7 <u>‡</u>	3	20	15	71

"Motor horsepowers for various batch volumes, viscosities in $\ensuremath{\mathsf{CP}}$, blend times in minutes.

• Denotes single four-bladed, 45" axial-flow impeller (unshaded selections).

† Denotes portable geardrive mixer with single 1.5-pitch propeller ("shaded" selections).

(Oldshue, 1983, p. 91).



Figure 10.4. Agitator flow patterns. (a) Axial or radial impellers without baffles produce vortexes. (b) Offcenter location reduces the vortex. (c) Axial impeller with baffles. (d) Radial impeller with baffles.

(10.5)

	()
d = impeller diameter (in.),	
D = vessel diameter (in.),	
N = rpm of impeller shaft,	
P = horsepower input,	
Q = volumetric pumping rate (cuft/sec),	
S = specific gravity,	

Froude number

$$t_b =$$
 blend time (min) ,

$$\mu$$
 = viscosity (cP).

 $N_{\rm T} = 7.454(10^{-4})N^2 d$

The Froude number is pertinent when gravitational effects are significant, as in vortex formation; in baffled tanks its influence is hardly detectable. The power, flow, and blend time numbers change with Reynolds numbers in the low range, but tend to level off above $N_{\rm Re} = 10,000$ or so at values characteristic of the kind of impeller. Sometimes impellers are characterized by their limiting N_p , as an $N_p = 1.37$ of a turbine, for instance. The dependencies on Reynolds number are shown on Figures 10.5 and 10.6 for power, in Figure 10.3 for flow and in Figure 10.7 for blend time.

Rough rules for mixing quality can be based on correlations of power input and pumping rate when the agitation system is otherwise properly designed with a suitable impeller (predominantly either axial or radial depending on the process) in a correct location, with appropriate baffling and the correct shape of vessel. The power input per unit volume or the superficial linear velocity can be used as measures of mixing intensity. For continuous flow reactors, for instance, a rule of thumb is that the contents of the vessel should be turned over in 5-10% of the residence time. Specifications of superficial linear velocities for different kinds of operations are stated later in this chapter. For baffled turbine agitation of reactors, power inputs and impeller tip speeds such as



Figure 10.5. Power number, $N_p = Pg_c/N^3 D^5 \rho$, against Reynolds number, $N_{\text{Re}} = ND^2 \rho/\mu$, for several kinds of impellers: (a) helical shape (*Oldshue*, 1983); (b) anchor shape (*Oldshue*, 1983); (c) several shapes: (1) propeller, pitch equalling diameter, without baffles; (2) propeller, s = d, four baffles; (3) propeller, s = 2d, without baffles; (4) propeller, s = 2d, four baffles; (5) turbine impeller, six straight blades, without baffles; (6) turbine impeller, six blades, four baffles; (7) turbine impeller, six curved blades, four baffles; (8) arrowhead turbine, four baffles; (9) turbine impeller, inclined curved blades, four baffles; (10) two-blade paddle, four baffles; (11) turbine impeller, six blades, four baffles; (12) turbine impeller with stator ring; (13) paddle without baffles (data of Miller and Mann); (14) paddle without baffles (data of White and Summerford). All baffles are of width 0.1D [after Rushton, Costich, and Everett, Chem. Eng. Prog. **46**(9), 467 (1950)].





the following may serve as rough guides:

Operation	HP/1000 gal	Tip Speed (ft/sec)
Blending	0.2-0.5	
Homogeneous reaction	0.5-1 .5	7.5-10
Reaction with heat transfer	1.5-5.0	10-15
Liquid-liquid mixtures	5	15-20
Liquid-gas mixtures	5-10	15-20
Slurries	10	

The low figure shown for blending is for operations such as



incorporation of TEL into gasoline where several hours may be allowed for the operation.

Example 10.1 deals with the design and performance of an agitation system to which the power input is specified. Some degree of consistency is found between the several rules that have been cited.

10.4. POWER CONSUMPTION AND PUMPING RATE

These basic characteristics of agitation systems are of paramount importance and have been investigated extensively. The literature is



Figure 10.6. Power number against Reynolds number of some turbine impellers [Bates, Fondy, and Corpstein, Ind. Eng. Chem. Process. Des. Dev. 2(4) 311 (1963)].



Figure 10.7. Flow number as a function of impeller Reynolds number for a pitched blade turbine with $N_p = 1.37$. D/T is the ratio of impeller and tank diameters. [Dickey, 1984, 12, 7; Chem. Eng., 102-110 (26Apr. 1976)].

reviewed, for example, by Oldshue (1983, pp. 155–191), Uhl and Gray (1966, Vol. 1), and Nagata (1975). Among the effects studied are those of type and dimensions and locations of impellers, numbers and sizes of baffles, and dimensions of the vessel. A few of the data are summarized on Figures 10.5–10.7. Often it is convenient to characterize impeller performance by single numbers; suitable ones are the limiting values of the power and flow numbers at high Reynolds numbers, above 10,000–30,000 or so, for example:

EXAMPLE 10.1

Impeller Size and Speed at a Specified Power Input

For a vessel containing 5000 gal of liquid with specific gravity = 0.9 and viscosity of 100 cP, find size and speed of a pitched turbine impeller to deliver 2 HP/1000 gal. Check also the superficial linear velocity and the blend time.

The dimensions of the liquid content are 9.5 ft high by 9.5 ft dia. Take

$$d = 0.40 = 0.4(9.5)(12) = 45.6 \text{ in., say 46 in., impeller,}$$

$$P = 2V = 2(5) = 10 \text{ HP,}$$

$$N_{\text{Re}} = \frac{10.75\text{SN}d^2}{\mu} = \frac{10.75(0.9)(46)^2\text{N}}{1000} = 20.47\text{N}$$

$$N_P = \frac{1.523(10^{13})P}{N^3D^5S} = \frac{1523(10^{13})(10)}{0.9(46)^5N^3} = \frac{821,600}{N^3}$$

Solve for N by trial with the aid of curve 6 of Figure 10.6.

Trial **N N_{Re} N_p N (Eq. (2)]** 56 1146 1.3 85.8 a4 1720 1.3 85.8

Туре	No. baffles	Np	Na
Propeller	0	0.3	
Propeller	3-8	0.33-0.37	0.40-0.55
Turbine, vertical blade	0	0.93-I .08	0.33-0.34
Turbine, vertical blade	4	3-5	0.70-0.85
Pitched turbine, 45"	0	0.7	0.3
Pitched turbine, 45"	4	1.30-1.40	0.60-0.87
Anchor	0	0.28	

A correlation of pumping rate of pitched turbines is shown as Figure 10.7.

Power input per unit volume as a measure of mixing intensity or quality was cited in Section 10.3 and in Chapter 17. From the correlations cited in this section, it is clear that power input and Reynolds number together determine also the pumping rate of a given design of impeller. This fact has been made the basis of a method of agitator system design by the staff of Chemineer. The superficial linear velocity-the volumetric pumping rate per pnit cross section of the tank-is adopted as a measure of quality of mixing. Table 10.2 relates the velocity to performance of three main categories of mixing: mixing of liquids, suspension of solids in slurries, and dispersion of gases. A specification of a superficial velocity will enable selection of appropriate impeller size, rotation speed, and power input with the aid of charts such as Figures 10.6 and 10.7. Examples 10.1 and 10.2 are along these lines.

The combination of HP and rpm that corresponds to a particular superficial velocity depends on the size of the tank, the size of the impeller, and certain characteristics of the system. Tables 10.3, 10.4, and 10.5 are abbreviated combinations of horsepower and rpm that are suitable at particular pumping rates for the three main categories of mixing. More complete data may be found in the literature cited with the tables.

1. For mixing of liquids, data are shown for a viscocity of 5000 cP, but data also have been developed for 25,000 cP, which allow for

Take N = 84 rpm. According to Figure 10.7 at d/D = 0.4,

 $N_Q = 0.61,$ $Q = N_Q N d^3 = 0.61(84/60)(46/12)^3 = 48.1 \text{ cfs},$ $u_s = 48.1/[(\pi/4)(9.5)^2] = 0.68 \text{ fps}.$

This value corresponds to moderate to high mixing intensity according to Table 10.2.

From Figure 10.3, at $N_{\text{Re}} = 1720$, blend time is given by

$$t_b N(d/D)^{2.3} = 17.0$$

or

$$t_b = \frac{17}{84(0.4)^{2.3}} - 1.67 \text{ min}$$

According to Table 10.1, the blend time is less than $6 \min$, which agrees qualitatively.

TABLE 10.2. Agitation Results Corresponding to Specific Superficial Velocities

ft/sec	Description	ft/sec	Description
Liquid Syst	ems		c. suspend all solids with the design settling velocity
0.1-0.2	low degree of agitation; a velocity of 0.2 ft/sec will a. blend miscible liquids to uniformity when specific gravity differences are less than 0.1 b. blend miscible liquids to uniformity if the ratio of viscosities is less than 100		completely off the bottom of the vessel d. provide slurry uniformity to at least one-third of the liquid level e. be suitable for slurry drawoff at low exit nozzle locations
	c. establish liquid movement throughout the vessel	0.6-0.8	when uniform solids distribution must be approached; a velocity of 0.6 ft/sec will
0.3-0.6	characteristic of most agitation used in chemical processing: a velocity of 0.6 ft/sec will		f. provide uniform distribution to within 95% of liquid level
	e. blend miscible liquids to uniformity if the specific gravity differences are less than 0.6		g. be suitable for slurry drawoff up to 80% of liquid level
	f. blend miscible liquids to uniformity if the ratio of viscosities is less than 10.000	0.9-1 .0	when the maximum feasible uniformity is needed. A velocity of 0.9 ft/sec will
	g. suspend trace solids (less than 2%) with settling rates of 2-4 ft/min		h. provide slurry uniformity to 98% of the liquid level i. be suitable for slurry drawoff by means of overflow
0.7-1.0	n. produce surface rippling at low viscosities high degree of agitation: a velocity of 1.0. ft/sec will	Gas Dispers	ion
0.7-1.0	 i. blend miscible liquids to uniformity if the specific gravity differences are less than 1.0 j. blend miscible liquids to uniformity if the ratio of viscosities is less than 100,000 k. suspend trace solids (less than 2%) with settling rates of 4-6 ft/min 	0.1-0.2	used when degree of dispersion is not critical to the process; a velocity of 0.2 ft/sec will a. provide nonflooded impeller conditions for coarse dispersion b. be typical of situations that are not mass transfer limited
	I. produce surging surface at low viscosities	0.3-0.5	used where moderate degree of dispersion is needed; a velocity of 0.5 ft/sec will
Solids Susp	ension		c. drive fine bubbles completely to the wall of the
0.1-0.2	minimal solids suspension; a velocity of 0.1 ft/sec will a. produce motion of all solids with the design settling velocity		vessel d. provide recirculation of dispersed bubbles back into the impeller
	b. move fillets of solids on the tank bottom and suspend them intermittently	0.6-1.0	used where rapid mass transfer is needed; a velocity of 1.0 ft/sec will
0.3-0.5	characteristic of most applications of solids suspension and dissolution; a velocity of 0.3 ft/sec will		e. maximize interfacial area and recirculation of dispersed bubbles through the impeller

[Chemineer, Co. Staff, Chem. Eng., 102-110 (26 April 1976); 144-150 (24 May 1976); 141-148 (19 July 1976)].

EXAMPLE 10.2

Effects of the Ratios of Impeller and Tank Diameters Power and rpm requirements will be investigated and compared with the data of Table 10.3. The superficial velocity is 0.6 ft/sec, V = 5000 gals, Sp Gr = 1.0. Viscosities of 100 cP and 5000 cP will be considered.

With h/D = 1, D = h = 9.47 ft,

pumping rate Q = $0.6(\pi/4)(9.47)^2$ = 42.23 cfs,	
$N_Q = 1.037(10^5)Q/Nd^3 = 4.3793/Nd^3$	(1)
$\tilde{N_{\text{Re}}} = 10.7Nd^2S/\mu = 0.00214Nd^2, \mu = 5000,$	(2)
- 11 13 150 /4 500 /4 013	()

$$P = N_p N^3 d^3 S / 1.523(10^{10}), \tag{3}$$

N_n from Figure 10.6.

For several choices of d/D, solve Eqs. (1) and (2) simultaneously with Figure 10.7. With $\mu = 5000$ cP;

d/D	d	N	<i>N</i> a [Eq. (1)]	N _{Re} [Eq. (2)]	N _a (Fig. 10.7)	N _p	<i>P</i> (HP)
0.25	28.4	300	0.637	518	0.64	1.4	45.9
0.33	37.5	145	0.573	436	0.57	1.45	21.5
0.50	56.8	52	0.460	359	0.45	1.5	8.2

With $\mu = 100$ cP, turbulence is fully developed.

					Na		
d/D	d	N	Na	N _{Re}	(Fig. 10.7)	N _p	Р
0.25	28.4	228	0.839	18,990	0.84	1.3	18.7
0.33	37.5	112	0.742	16,850	0.74	1.3	8.9
0.50	56.8	40	0.597	13,800	0.60	1.3	3.2

Table 10.3 gives these combinations of HP/rpm as suitable: 25/125, 20/100, 10/56, 7.5/37. The combination 10/56 checks roughly the last entry at 5000 cP. Table 10.3 also has data for viscosities of 25,000 cP, thus allowing for interpolation and possibly extrapolation.

			Volume	(gal)		
		5000 cP			25, 000 cP	
ft/sec	1000	2000	5000	1000	2000	5000
0.1	2/280	2/190	2/100	2/125	2/84	7.5/125
	1/190	1/100		1.5/84	1.5/56	5/100
						5/84
	0/000	0/405	= /+ ==		E /40E	3/56
0.2	2/190	2/125	5/125	3/84	5/125	10/84
	1/100	2/84	3/84	2/84	3/84	/.5/68
		1.5/84	3/68	1.5/50	3/68	5/45
0.0	0/105	2/04	2/45	E /10E	2/40	3/3/
0.3	2/120	3/04	/.5/125	5/125	15/155	15/00
	1.5/04	1.5/50	5/100	2/04	7.5/00 E/AE	10/00
			2/64	3/00	2/45	10/40
0.4	2/8/	5/125	3/50	75/9/	10/8/	20/100
0.4	15/56	3/68	7 5169	5/56	75/45	25/84
	1.5/50	3/56	5/A5	3,30	7.5745	20/68
		2/45	3/37			10/37
0.5	5/125	7.5/125	15/100	15/155	25/125	75/190
	3/84	5/84	10/68	10/100	20/100	60/155
	•,•	-,	7.5/45	10/84	15/84	40/100
			,	7.5/68	10/56	15/45
0.6	5/100	15/155	25/125	20/155	25/100	40/84
	3/68	10/100	20/100	15/125	15/68	30/68
	3/56	7.5/84	10/56		15/56	25/56
	2/45	3/37	7.5/37		10/45	20/37
0.7	7.5/125	10/84	15/68	25/155	40/155	75/125
	5/84	7.5/68	15/56	15/84	30/100	50/84
		5/45	10/45		25/84	30/45
			10/37		20/68	
0.8	10/125	10/68	30/100	30/155	50/155	75/100
	7.5/100	7.5/56	25/84	25/125	40/125	60/84
			20/68	20/100		50/68
			15/45		(40/56
0.9	15/155	15/84	60/155	40/155	75/190	75/84
	10/100	10/56	40/100	30/125	60/155	60/68
	7.5/84	/.5/45		25/100	40/100	50/56
	10/04	20/155	50/000	40 405	30/68	105/405
1.0	10/84	30/155	50/100	40/125	50/125	125/125
	7.2/22	25/125	40/84	30/100	50/100	75/00
		20/100	30/08		20/84 40/84	/0/00 60/56
		15/00	20/00		40/04	00/90

TABLE	10.3.	Mixing of Liquids; Power and Impeller Speed (hp/rpm) for
		Two Viscosities, as a Function of the Liquid Superficial
		Velocity; Pitched Blade Turbine Impeller

[Hicks, Morton, and Fenic, Chem. Eng., 102-110 (26 April 1976)].

interpolation and possibly extrapolation. The impeller is a pitched-blade turbine.

- 2. For suspension of solids, the tables pertain to particles with settling velocities of 10 ft/min, but data are available for 25 ft/min. The impeller is a pitched-blade turbine.
- 3. For gas dispersion the performance depends on the gas rate. Data are shown for a superficial inlet gas rate of 0.07 ft/sec, but data are available up to 0.2 ft/sec. Four baffles are specified and the impeller is a vertical blade turbine.

Distant Street of

Example 10.2 compares data of Table 10.4 with calculations based on Figures 10.6 and 10.7 for all-liquid mixing. Power and rpm requirements at a given superficial liquid velocity are seen to be very sensitive to impeller diameter. When alternate combinations of **HP/rpm** are shown in the table for a particular performance, the design of the agitator shaft may be a discriminant between them. The shaft must allow for the torque and bending moment caused **by** the hydraulic forces acting on the impeller and shaft. Also, the

impeller and shaft must not rotate near their resonant frequency. Such mechanical details are analyzed by Ramsey and Zoller [*Chem. Eng.*, 101-108 (30 Aug. 1976)].

10.5. SUSPENSION OF SOLIDS

Besides the dimensions of the vessel, the impeller, and baffles, certain physical data are needed for complete description of a slurry mixing problem, primarily:

- 1. Specific gravities of the solid and liquid.
- 2. Solids content of the slurry (wt %).
- 3. Settling velocity of the particles (ft/min).

The last of these may be obtained from correlations when the mesh size or particle size distribution is known, or preferably experimentally. Taking into account these factors in their effect on suspension quality is at present a highly empirical process. Tables

			Vol ume	(gal)		
		10 ft/min		_	25 ft/min	
ft/sec	1000	2000	5000	1000	2000	5000
0.1	1/190	2/190	5/125	2/190	2/125	5/125
		1/100	3/84	1/190	2/84	3/84
		·	3/68	1/100	1.5/84	3/68
			2/45		1.5/56	2/45
0.2	1/100	2/125	7.5/125	2/125	3/84	15/155
	·	1.5/84	5/100			10/100
			5/84			7.5/68
			3/56			5/45
0.3	2/190	2/84	3/37	1.5/84	5/125	10/84
		1.5/56			3/68	
					2/45	
0.4	2/155	5/155	7.5/84	2/84	7.5/155	7.5/45
	1.5/100		5/56	1.5/56	5/100	
					3/56	
0.5	1.5/84	3/84	15/155	2/68	7.5/125	15/84
	2/125		10/100	2/56	5/84	10/56
			7.5/68			7.5/37
			5/45			
0.6	2/100	5/125	10/84	3/84	5/56	25/125
	1.5/68	3/68				20/100
		3/56				15/68
		2/45			_	10/45
0.7	2/84	7.5/155	15/84	7.5/155	15/155	30/100
	1.5/56	7.5/125	10/56	5/125	10/100	25/84
		5/84	7.5/45	5/100	7.5/84	20/68
			7.5/37	3/68	7.5/68	15/56
0.8	3/84	7.5/84	25/125	7.5/125	1 0/84	60/155
		5/56	20/100	5/84		40/100
			15/68			30/68
	/		10/45		45 10 4	25/56
0.9	7.5/155	15/155	40/155	10/125	15/84	/5/190
	5/125	10/100	30/100	7.5/100		60/125
	5/100	7.5/68	25/84			50/100
	3/68	20 /1 00	20/68	45 /455	20/155	40/84
1.0	7.5/125	20/100	50/100	15/155	30/155	/5/125
	5/84	15/84	40/84	10/100	25/125	75/100
		10/84	30/68		20/100	60/84
			25/56			50/84

TABLE 10.4. Suspension of Solids; Power and Impeller Speed (hp/rpm) for Two Settling Velocities, as a Function of the Superficial Velocity of the Liquid; Pitched Blade Turbine Impeller

[Gates, Morton, and Fondy, Chem. Eng., 144-150 (24 May 1976)].

10.2-10.5 are one such process; the one developed by Oldshue (1983) will be examined shortly.

Suspension of solids is maintained by upward movement of the liquid. In principle, use of a draft tube and an axial flow impeller will accomplish this flow pattern most readily. It turns out, however, that such arrangements are suitable only for low solids contents and moderate power levels. In order to be effective, the cross section of the draft tube must be appreciably smaller than that of the vessel, so that the solids concentration in the draft tube may become impractically high. The usually practical arrangement for solids suspension employs a pitched blade turbine which gives both axial and radial flow.

For a given tank size, the ultimate design objective is the relation between power input and impeller size at a specified uniformity. The factors governing such information are the slurry volume, the slurry level, and the required uniformity. The method of Oldshue has corrections for these factors, as F_1 , F_2 , and F_3 . When multiplied together, they make up the factor F_4 which is the ordinate of Figure 10.8(d) and which determines what combinations

of horsepower and ratio of impeller and vessel diameters will do the required task. Example 10.3 employs this method, and makes a comparison with the Chemineer method of Tables 10.2 and 10.3.

10.6. GAS DISPERSION

Gases are dispersed in liquids usually to facilitate mass transfer between the phases or mass transfer to be followed by chemical reaction. In some situations gases are dispersed adequately with spargers or porous distributors, but the main concern here is with the more intense effects achievable with impeller driven agitators.

SPARGERS

Mixing of liquids and suspension of solids may be accomplished by bubbling with an inert gas introduced uniformly at the bottom of the tank. For mild agitation a superficial gas velocity of 1 ft/min is used, and for severe, one of about 4 ft/min.

TABLE 10.5. Dispersion of Gases; Power and Impeller Sp	eed
(hr/rpn) for Two Gas Inlet Superficial Velocit	ies,
as a Function of the Liquid Superficial Velocit	y;
Vertical Blade Turbine Impeller	

			Volume	(gal)		
		0.07 ft/sec			0. 20 ft/sec	;
ft/sec	1500	3000	5000	1500	3000	5000
0.1	2/56	5/84	7.5/68	3/56	7.5/68	1 0/45
0.0	0.45	7 5 /4 05	15/155	0/45	10/100	
0.2	2/45	7.5/125	10/84	3/45	15/155	15/68
			7.3743	5/100	10/04	20/100
03	3/84	7.5/68	10/45	5/100	7 6/45	25/125
0.0	3/68	5/45	10/56		1.5/45	
	3/56	7.5/84	10,00			
	-,	5/56				
0.4	5/125	10/84	15/68	5/84	10/45	30/1 55
	5/84	1 0/100	20/1 00	7. 51155	10/56	20/68
	5/100	1 0/45	15/84	5/56	-,	15/45
	5/45	10/56	20/68	•		15/56
0.5	7.5/125	15/1 <i>55</i>	25/1 <i>2</i> 5	7.5/125	15/68	25/84
	7.5/155	15/68	25/84	7.5/68	15/84	25/100
	7.5/68	15/84	25/100	7.5/84	15/45	25/56
	7.5/84	15/45	25/56		15/56	
0.6	10/84	20/1 <i>00</i>	30/1 <i>55</i>	1 0/84	20/1 <i>00</i>	30/100
	10/100	20/68	30/100	10/100	20/68	30/125
		20/45	30/1 <i>2</i> 5			30/68
			30/68			30/45
0.7	10/56	25/1 <i>25</i>	40/1 55	10/56	25/125	40/155
		25/84	40/84		25/84	40/84
		25/1 00	40/1 00		25/100	40/100
		25/56	40/56		25/56	40/56
0.8	15/1 55	30/1 55	50/1 00	15/155	30/155	50/100
	15/84	30/100	50/68	15/84	30/100	50/68
		30/125	50/84		30/1 25	50/84
0.0	15 100	20/00	50/45	45 100	00/00	50/56
0.9	15/68	30/68	60/1 25	15/68	30/68	60/125
			60/155			60/155
			60/84			60/84
1.0	20/100	40/155	00/50	25/1 25	40/1 55	00/00
1.0	20/68	40/84	75/100	20/120 25/84	40/155	75/190
	20/00		75/125	20/04	40/04	75/100
			10/120			70/120

[Hicks and Gates, Chem. Eng, 141-148 (19 July 1976)].

MASS TRANSFER

The starting point of agitator design is properly a mass transfer coefficient known empirically or from some correlation in terms of parameters such as impeller size and rotation, power input, and gas flow rate. Few such correlations are in the open literature, but some have come from two of the industries that employ aerated stirred tanks on a large scale, namely liquid waste treating and fermentation processes. A favored method of studying the absorption of oxygen is to measure the rate of oxidation of aqueous sodium sulfite solutions. Figure 10.9 summarizes one such investigation of the effects of power input and gas rate on the mass transfer coefficients. A correlation for fermentation air is given by Dickey (1984, **12-17**):

$$k_L a = \text{rate/(concentration driving force)}$$

= 0.064(P_g/V)^{0.7}u_g^{0.2}, 1/sec, (10.6)

with P_g/V in HP/1000 gal and superficial gas velocity u_g in ft/sec. A general correlation of mass transfer coefficient that does not have

power input as a factor is given by Treybal (*Mass Transfer Operations*, McGraw-Hill, New York, 1980, 156); presumably this is applicable only below the minimum power input here represented by Figure 10.11.

When mass transfer coefficients are not determinable, agitator design may be based on superficial liquid velocities with the criteria of Table 10.2.

SYSTEM DESIGN

The impeller commonly used for gas dispersion is a radial turbine with six vertical blades. For a liquid height to diameter ratio $h/D \le 1$, a single impeller is adequate; *in* the *range* $1 \le h/D \le 1.8$ two are needed, and more than two are rarely used. The lower and upper impellers are located at distances of 1/6 and 2/3 of the liquid level above the bottom. Baffling is essential, commonly with four baffles of width 1/12 that of the tank diameter, offset from the wall at 1/6 the width of the baffle and extending from the tangent line of the wall to the liquid level. The best position for inlet of the gas is below and at the center of the lower impeller; an open pipe is commonly used, but a sparger often helps. Since ungassed power is significantly larger than gassed, a two-speed motor is desirable to prevent overloading, the lower speed to cut in automatically when the gas supply is interrupted and rotation still is needed.

MINIMUM POWER

Below a critical power input the gas bubbles are not affected laterally but move upward with their natural buoyancy. This condition is called gas flooding of the impeller. At higher power inputs the gas is dispersed radially, bubbles impinge on the walls and are broken up, consequently with improvement of mass transfer. A correlation of the critical power input is shown as Figure 10.10.

POWER CONSUMPTION OF GASSED LIQUIDS

At least partly because of its lower density and viscosity, the power to drive a mixture of gas and liquid is less than that to drive a liquid. Figure 10.11(a) is a correlation of this effect, and other data at low values of the flow number Q/Nd^3 are on Figure 10.11(b). The latter data for Newtonian fluids are correlated by the equation

$$P_{\sigma}/P = 0.497(Q/Nd^3)^{-0.38}(N^2d^3\rho_L/\sigma)^{-0.18},$$
(10.7)

where the last group of terms is the Weber number, ρ_L is the density of the liquid, and σ is its surface tension.

SUPERFICIAL LIQUID VELOCITY

When mass transfer data are not known or are not strictly pertinent, a quality of mixing may be selected by an exercise of judgment in terms of the superficial liquid velocity on the basis of the rules of Table 10.2. For gas dispersion, this quantity is related to the power input, **HP/1000** gal, the superficial gas velocity and the ratio d/D in Figure 10.12.

DESIGN PROCEDURES

On the basis of the information gathered here, three methods are possible for the design of agitated gas dispersion. In all cases the size of the tank, *the* ratio *of impeller and* tank diameters and the gas feed rate are specified. The data are for radial turbine impellers with six vertical blades.



Figure 10.8. Suspension of solids. Power and ratio of diameters of impeller and tank, with four-bladed 45" impeller, width/diameter = 0.2. [method of *Oldshue (1983)*]. (a) The factor on power consumption for slurry volume, F_1 . (b) The factor on power requirement for single and dual impellers at various h/D ratios, F_2 . (c) The effect of settling velocity on power consumption, F_3 . (d) Suspension factor for various horsepowers: $F_4 = F_1F_2F_3$.

EXAMPLE 10.3

Design of the Agitation System for Maintenance of a Slurry These conditions are taken:

$$V = 5000$$
 gal.

h/D = 1, settling velocity = 10 ft/min, solids content = 10 wt %

Reading from Figure 10.8,

$$F_{1} = 4,$$

$$F_{2} = 1.1,$$

$$F_{3} = \begin{cases} 3.0, & \text{off bottom}, \\ 10.0, & \text{uniform}, \end{cases}$$

$$F_{4} = F_{1}F_{2}F_{3} = \begin{cases} 13.2, & \text{off bottom}, \\ 44, & \text{uniform}. \end{cases}$$

The relation between the ratio of impeller and vessel diameters,

1. Start with a known required mass transfer coefficient. From a correlation such as Figure 10.9 or Eq. (10.6) the gassed power per unit volume will become known, and the total gassed power to the tank will be P_g . The ratio of gassed power to ungassed power is represented by Figure 10.11(a) and the equations given there; at this stage the rotation speed N is not yet known. This value is found by trial by simultaneous solution with Figure 10.6 which relates the Reynolds and power numbers; the power here is the ungassed power. The value of N that results in the precalculated P_g will be the correct one. Curve 2 of Figure 10.6 is the one applicable to gas dispersion with the data of this section. **2.** Start with a choice of superficial liquid velocity u_L made in accordance with the criteria of Table 10.2. With the aid of the known gas velocity u_s and d/D, find P_g/V from Figure 10.12. Then proceed to find N by trial with Figures 10.11(a) and 10.6 as in method 1.



Figure 10.9. Typical data of mass transfer coefficients at various power levels and superficial gas rates for oxidation of sodium sulfite in aqueous solution. d/D = 0.25-0.40 (Oldshue, 1983).

d/D and HP is read off Figure 10.8(d).

	H	I P
d/D	Off btm	Uniform
0.2	20	65
0.4	7.5	25
0.6	4	12

Comparing with readings from Tables 10.2 and 10.3,

Superficial liq. velocity	HP/rpm
0.3 (off btm)	1 0/45, 1 0/56
0.6 (uniform)	30/155, 30/125, 30/100, 30/68

These results correspond roughly to those of the Oldshue method at d/D = 0.4. The impeller sizes can be determined with Figures 10.6 and 10.7.

3. As soon as a superficial liquid velocity has been selected, a suitable combination of **HP/rpm** can be taken from Table 10.5.

These procedures are applied in Example 10.4.

As general rules, levels of 5-12 HP/1000 gal are typical of aerobic fermentation vessels. and 1-3 HP/1000 gal of aerobic waste treatment; concentrations and oxygen requirements of the microorganisms are different in the two kinds of processes.



Figure 10.10. Minimum power requirement to overcome flooding as a function of superficial gas velocity and ratio of impeller and tank diameters, *d/D.* [*Hicks and Gates,* Chem. Eng., 141-148 (19 July 1976)].



Figure 10.11. Power consumption. (a) Ratio of power consumptions of aerated and unaerated liquids. Q is the volumetric rate of the gas: (0) glycol; (\mathbf{x}) ethanol; ($\mathbf{\nabla}$) water. [After Calderbank, Trans. Inst. Chem. Eng. 36, 443 (1958)]. (b) Ratio of power consumptions of aerated and unaerated liquids at low values of Q/Nd^3 . Six-bladed disk turbine: (\Box) water; ($\mathbf{\Theta}$) methanol (10%); (A) ethylene glycol (8%); (A) glycerol (40%); P_g gassed power input; P = ungassed power input; Q = gas flow rate; N = agitator speed; d = agitator-impeller diameter. [Luong and Volesky, AIChE J. 25, 893 (1979)].

(b)

10.7. IN-LINE BLENDERS AND MIXERS

When long residence time is not needed for chemical reaction or other purposes, small highly powered tank mixers may be suitable, with energy inputs measured in HP/gal rather than HP/1000 gal. They bring together several streams continuously for a short contact time (at most a second or two) and may be used whenever the effluent remains naturally blended for a sufficiently long time, that is, when a true solution is formed or a stable emulsion-like mixture. When it is essential that the mixing be immediate each stream will



Figure 10.12. Relation between power input, P/V HP/1000 gal, superficial liquid velocity u_L ft/sec, ratio of impeller and tank diameters, d/D, and superficial gas velocity u_s ft/sec. [Hicks and Gates, Chem. Eng., 141-148 (19 July 1976)].

have its own feed nozzle, as in Figure 10.13(b), but usually the streams may be combined externally near the blender and then given the works, as in Figure 10.13(a).

One manufacturer gives these power ratings:

Tank size	(gal)	1	5	10	30
Motor HP	-	0.5	1	2	3

Another ties in the line and motor sizes:

Line size, (in.) I-4 6-8 10-12 Motor HP 0.5 1 2

But above viscosities of 10 cP a body one size larger than the line size is recommended.

Other devices utilize the energy of the flowing fluid to do the mixing. They are inserts to the pipeline that force continual changes of direction and mixing. Loading a section of piping with tower packing is an example but special assemblies of greater convenience have been developed, some of which are shown in Figure 10.14. In each case manufacturer's literature recommends the sizes and pressure drops needed for particular services.

The Kenics mixer, Figure 10.14(a), for example, consists of a succession of helical elements twisted alternately in opposite directions. In laminar flow for instance, the flow is split in two at each element so that after n elements the number of striations becomes 2". The effect of this geometrical progression is illustrated in Figure 10.14(b) and points out how effective the mixing becomes after only a few elements. The Reynolds number in a corresponding empty pipe is the major discriminant for the size of mixer, one manufacturer's recommendations being

N _{Re}	Number of Elements
Less than 10	2 4
10-2000	12-18
More than 2000	6

Besides liquid blending applications, static mixers have been **used** for mixing gases, **pH** control, dispersion of gases into liquids, and dispersion of dyes and solids in viscous liquids. They have the advantages of small size, ease of operation, and relatively low cost. The strong mixing effect enhances the rate of heat transfer from viscous streams. Complete heat exchangers are built with such

EXAMPLE 10.4

HP and 'pm Requirements of an Aerated Agitated Tank A tank contains 5000 gal of liquid with sp gr = 1.0 and viscosity 100 cP that is aerated and agitated. The ratio of impeller to tank diameters is d/D = 0.4. Two sets of conditions are to be examined.

- **a.** The air rate is 972 SCFM or 872ACFM at an average submergence of 4 ft. The corresponding superficial gas velocity is 0.206 ft/sec or 0.063 m/sec. A mass transfer coefficient $k_L a = 0.2$ /sec is required; Dickey's equation (10.6) applies. Find the power and rpm needed.
- **b.** The air rate is 296 ACFM, 0.07 ft/sec, 0.0213 m/sec. The required intensity of mixing corresponds to a liquid superficial velocity of 0.5 ft/sec. Find the power, rotation speed, and mass transfer coefficients for sulfite oxidation and for fermentation.

a. d = 0.4(9.47) = 3.79 ft, 45.46 in., $k_L a = 0.064(P_g/V)^{0.7} u_g^{0.2} = 0.2$, $P_g/V = [0.2/0.064(0.206)^{0.2}]^{1/0.7} = 8.00$ HP, $P_g = 5(8.0) = 40.0$ HP/5000 gal, $Q/Nd^3 = 872/(379)^3N = 16.02/N$, $N_{\text{Re}} = 10.75Nd^2S/\mu = 10.75(45.46)^2N/100 = 222N$.

Equation (10.2),

$$N_n = 1.523(10^{13})P/N^3d^5S = 78,442P/N^3.$$

Curve 2 of Figure (10.6) applies. P_g/P from Figure 10.10(a). Solve by trial.

N	Q/Nd°	P _g /P	N _{Re}	N _p	Р	Pg
100	0.160	0.324	22, 200	4	51	16.5
150	0. 107	0.422	33, 300	4	172	72.6
127	0. 1261	0.3866	28, 194	4	104.5	40. 4- 40. 0

The last entry of P_g checks the required value 40.0. Find the corresponding superficial liquid velocity with Figure 10.12:

$$X = (P/V)(d/D)^{1.85} = 8.04(0.4)^{1.85} = 1.48,$$

at $u_G = 0.206$ ft/sec, Y = 2.0,

 $\therefore u_L = 2/10(0.4)^{1.2} = 0.60$ ft/sec.

mixing inserts in the tubes and are then claimed to have 3-5 times normal capability in some cases.

10.8. MIXING OF POWDERS AND PASTES

Industries such as foods, cosmetics, pharmaceuticals, plastics, rubbers, and also some others have to do with mixing of high viscosity liquids or pastes, of powders together and of powders with pastes. Much of this kind of work is in batch mode. The processes are so diverse and the criteria for uniformity of the final product are so imprecise that the nonspecialist can do little in the way of equipment design, or in checking on the recommendations of equipment manufacturers. Direct experience is the main guide to selection of the best kind of equipment, predicting how well and quickly it will perform, and what power consumption will be. For From Table 10.2, a liquid velocity of 0.6-0.7 ft/sec will give moderate to high dispersion. Table 10.5 gives possible HP/rpm combination of 30/125, somewhat less than the value found here.

b. With liquid circulation velocity specified,

$$u_L = 0.5 \text{ ft/sec.}$$

Use Figure 10.12:

$$Y = iou_L (d/D)^{1.2} = 10(0.5)(0.4)^{1.2} = 1.67,$$

X= 0.8,
 $P_R/V = 0.8/(0.4)^{1.85} = 4.36$ HP/1000 gal

(this does exceed the minimum of 1.6 from Figure 10.11),

$$P_{g} = 5(4.36) = 21.8,$$

$$\frac{Q}{Nd^{3}} = 296/(3.79)^{3}N = 5.437/N,$$

$$N_{Re} = 222N \quad (part a),$$

$$N_{=}\frac{78,442P}{N} \quad (part a).$$

Solve by trial, using Figure 10.10(a) and curve 2 of Figure 10.6.

Ν	Q/Nd ³	Pg/P	N _{Re}	NĻ,	P	Pg
100	0.0544	0.5194	22, 200	4	51	26.5
94	0.0576	0.5130		4	42.35	21.7 ≅ 2.8

The closest reading from Table 10.5 is HP/rpm = 25/100 which is a good check.

For sulfite oxidation, at $u_g = 0.07$ ft/sec,

 $P_g/V = 4.36$ HP/1000 gal, from Figure 10.9, $k_e a = 0.07$ lb mol/(cuft)/(hr)(atm).

For fermentation, Eq. 10.6 gives

$$k_L a = 0.064(4.36)^{0.7}(0.07)^{0.2}$$

= 0.105 lb mol/(cuft)(sec)
lb mol/cuft

projects somewhat out of direct experience and where design by analogy may not suffice, testing in pilot plant equipment is a service provided by many equipment suppliers.

A few examples of mixers and blenders for powders and pastes are illustrated in Figure 10.15. For descriptions of available equipment-their construction, capacity, performance, power consumption, etc.--the primary sources are catalogs of manufacturers and contact with their offices. Classified lists of manufacturers, and some of their catalog information, appear in the *Chemical Engineering Catalog* (Reinhold, New York, annually) and in the *Chemical Engineering Equipment Buyers Guide* (McGraw-Hill, New York, annually). Brief descriptions of some types of equipment are in *Perry's Chemical Engineers Handbook* (McGraw-Hill, New York, 1984 and earlier editions). Well-classified descriptions, with figures, of paste mixers are in Ullmann (1972,



Figure 10.13. Motor-driven in-line blenders: (a) Double impeller made by Nettco Corp.; (b) three-inlet model made by Cleveland Mixer Co.



Figure 10.14. Some kinds of in-line mixers and blenders. (a) Mixing and blending with a recirculating pump. (b) Injector mixer with a helical baffle. (c) Several perforated plates (orifices) supported on a rod. (d) Several perforated plates flanged in. (e) **Hellical** mixing elements with alternating directions (Kenics Corp.). (f) Showing progressive striations of the flow channels with Kenics mixing elements.





(a)



(c)



(e)



(d)



(f)





(h)

Figure 10.15. Some mixers and blenders for powders and pastes. (a) Ribbon blender for powders. (b) Flow pattern in a double cone blender rotating on a horizontal axis. (c) Twin shell (Vee-type); agglomerate breaking and liquid injection are shown on the broken line. (d) Twin rotor; available with jacket and hollow screws for heat transfer. (e) Batch muller. (f) Twin mullers operated continuously. (g) Double-arm mixer and kneader (*Baker-Perkins Inc.*). (h) Some types of blades for the double-arm kneader (*Baker-Perkins Inc.*).

Vol. 2, pp. 282-300) and a similar one for powder mixers (*loc. cit.*, pp. 301-311). Since this equipment industry has been quite stable,

older books are still useful, notably those of Riegel (1953), Mead (1964), and particularly Kieser (1934-1939).

REFERENCES

- R.S. Brodkey (Ed.), Turbulence in Miring Operations, Academic, New York, 1975.
- Chemineer Co. Staff, Liquid Agitation, Reprint of 12 articles from Chemical Engineering, 8 Dec. 1975-6 Dec. 1976.
- D.S. Dickey, In Handbook of Chemical Engineering Calculations, (N.P. Chopey and T.G. Hicks Eds.), McGraw-Hill, New York, 1984.
- S. Harnby, M.F. Edwards, and A.W. Nienow, *Mixing in the Process Industries*, Butterworths, Stoneham, MA, 1985.
- A.J. Kieser, Handbuch der chemisch-technischen Apparate, Springer-Verlag, Berlin, 1934-1939.
- W.J. Mead, Encyclopedia of Chemical Process Equipment, Reinhold, New York, 1964.

- S. Nagata, Mixing Principles and Applications, Wiley, New York, 1975.
- J.Y. Oldshue, *Fluid Mixing Technology*, McGraw-Hill, New York, 1983.
- 9. E.R. Riegel, Chemical Process Machinery, Reinhold, New York, 1953.
- Z. Sterbacek and P. Tausk, *Miring in the Chemical Industry*, Pergamon, New York, 1965.
- J.J. Ulbrecht and G.K. Patterson, Mixing of Liquids by Mechanical Agitation, Gordon & Breach, New York, 1985.
- 12. V. Uhl and J.B. Gray (Eds.), Mixing Theory and Practice, Academic, New York, 1966, 1967, 2 vols.
- Ullmann's Encyclopedia of Chemical Technology, Verlag Chemie, Weinheim, Germany, 1972, Vol. 2, pp. 249-311.

SOLID-LIQUID SEPARATION

olid–liquid separation is concerned with mechanical processes for the separation of liquids and finely divided insoluble solids.

11.1. PROCESSES AND EQUIPMENT

Much equipment for the separation of liquids and finely divided solids was invented independently in a number of industries and is of diverse character. These developments have occurred without benefit of any but the most general theoretical considerations. Even at present, the selection of equipment for specific solid-liquid separation applications is largely a process of scale-up based on direct experimentation with the process material.

The nature and sizing of equipment depends on the economic values and proportions of the phases as well as certain physical properties that influence relative movements of liquids and particles. Pressure often is the main operating variable so its effect on physical properties should be known. Table 11.1 is a broad classification of mechanical processes of solid-liquid separation. Clarification is the removal of small contents of worthless solids from a valuable liquid. Filtration is applied to the recovery of valuable solids from slurries. Expression is the removal of relatively small contents of liquids from compressible sludges **by** mechanical means.

Whenever feasible, solids are settled out by gravity or with the aid of centrifugation. In dense media separation, an essentially homogeneous liquid phase is made by mixing in finely divided solids (less than 100 mesh) of high density; specific gravity of 2.5 can be attained with magnetite and 3.3 with ferrosilicon. Valuable ores and coal are floated away from gangue by such means. In flotation, surface active agents induce valuable solids to adhere to gas bubbles which are skimmed off. Magnetic separation also is practiced when feasible. Thickeners are vessels that provide sufficient residence time for settling to take place. Classifiers incorporate a mild raking action to prevent the entrapment of fine particles by the coarser ones that are to be settled out. Classification also is accomplished in hydrocyclones with moderate centrifugal action.

TABLE 11 1.	Chief Mechai	nical Means of
	Solid-Liquid	Separation

1.	Settling
	a by gravity
	in thickeners
	II. IN CLASSIFIERS
	D. by centritugal force
	c. by air flotation
	d. by dense media flotation
	e. by magnetic properties
2.	Filtration
	a. on screens, by gravity
	b. on filters
	i by vacuum
	ii. by pressure
	iii. by centrifugation
3.	Expression
	a. with batch presses
	b. with continuous presses
	, sorew presses
	ii. rolls
	iii. discs

Freely draining solids may be filtered by gravity with horizontal screens, but often filtration requires a substantial pressure difference across a filtering surface. An indication of the kind of equipment that may be suitable can be obtained by observations of sedimentation behavior or of rates of filtration in laboratory vacuum equipment. Figure 11.1 illustrates typical progress of sedimentation. Such tests are particularly used to evaluate possible flocculating processes or agents. Table 11.2 is a classification of equipment based on laboratory tests; test rates of cake formation range from several **cm/sec** to fractions of a **cm/hr**.

Characteristics of the performance of the main types of commercial SLS equipment are summarized in Table 11.3. The completeness of the removal of liquid from the solid and of solid from the liquid may be important factors. In some kinds of equipment residual liquid can be removed by blowing air or other gas through the cake. When the liquid contains dissolved substances that are undesirable in the filter cake, the slurry may be followed by



Figure 11.1. Sedimentation behavior of a slurry, showing loose and compacted zones (Osborne, 1981).
TABLE 11.2. Equipment Selection on the Basis of Rate of Cake Buildup

Process	Туре	Rate of Cake Buildup	Suitable Equipment
Rapid filtering		0.1-10 cm/sec	gravity pans; horizontal belt or top feed drum; continuous pusher type centrifuge
Medium filtering		O.I-IOcm/min	vacuum drum or disk or pan or belt; peeler type centrifuge
Slow filtering		0.1-10 cm/hr	pressure filters; disc and tubular centrifuges; sedimenting centrifuges
Clarification		negligible cake	cartridges; precoat drums; filter aid systems; sand deep bed filters

(Tiller and Crump, 1977; Flood, Parker, and Rennie, 1966).

pure water to displace the residual filtrate. Qualitative cost comparisons also are shown in this table. Similar comparisons of filtering and sedimentation types of centrifuges are in Table 11.19.

Final selection of filtering equipment is inadvisable without some testing in the laboratory and pilot plant. A few details of such work are mentioned later in this chapter. Figure 11.2 is an outline of a procedure for the selection of filter types on the basis of appropriate test work. Vendors need a certain amount of information before they can specify and price equipment; typical inquiry forms are in Appendix C. Briefly, the desirable information includes the following.

- Flowsketch of the process of which the filtration is a part, with the expected qualities and quantities of the filtrate and cake.
- 2. Properties of the feed: amounts, size distribution, densities and chemical analyses.
- 3. Laboratory observations of sedimentation and leaf filtering rates.
- 4. Pretreatment options that may be used.
- 5. Washing and blowing requirements.
- 6. Materials of construction.

A major aspect of an SLS process may be conditioning of the slurry to improve its filterability. Table 11.4 summarizes common pretreatment techniques, and Table 11.5 lists a number of flocculants and their applications. Some discussion of pretreatment is in Section 11.3.

11.2. THEORY OF FILTRATION

Filterability of slurries depends so markedly on small and unidentified differences in conditions of formation and aging that no correlations of this behavior have been made. In fact, the situation is so discouraging that some practitioners have dismissed existing filtration theory as virtually worthless for representing filtration behavior. Qualitatively, however, simple filtration theory is directionally valid for modest scale-up and it may provide a structure on which more complete theory and data can be assembled in the future.

As filtration proceeds, a porous cake of solid particles is built up on a porous medium, usually a supported cloth. Because of the fineness of the pores the flow of liquid is laminar so it is represented by the equation

$$Q = \frac{dV}{dt} = \frac{A\Delta P}{\mu R}.$$
(11.1)

The resistance R is made up of those of the filter cloth R_f and that of the cake R_c which may be assumed proportional to the weight of the cake. Accordingly,

$$Q = \frac{dV}{dt} = \frac{AAP}{\mu(R_f + R_c) = \mu(R_f + \alpha cV/A)}$$
(11.2)
 α = specific resistance of the cake (m/kg),
c = wt of solids/volume of liquid (kg/m³),
 μ = viscosity (N sec/m²)
 P = pressure difference (N/m²)
 A = filtering surface (m²)
 V = volume of filtrate (m³)
 Q = rate of filtrate accumulation (m³/sec).

 R_f and α are constants of the equipment and slurry and must be evaluated from experimental data. The simplest data to analyze are those obtained from constant pressure or constant rate tests for which the equations will be developed. At constant pressure Eq. (11.2) is integrated as

$$\frac{A\Delta P}{\mu}t = R_f V + \frac{\alpha c}{2A}V^2 \tag{11.3}$$

and is recast into linear form as

$$\frac{t}{V/A} = \frac{\mu}{\Delta P} R_f + \frac{\mu \alpha c}{2\Delta P} \frac{V}{A}.$$
(11.4)

The constants R_f and α are derivable from the intercept and slope of the plot of t/V against V. Example 11.1 does this. If the constant pressure period sets in when $t = t_0$ and $V = V_0$, Eq. (11.4) becomes

$$\frac{t - t_0}{V - V_0} = \frac{\mu}{A\Delta P} R_f + \frac{\mu\alpha c}{2A^2 \Delta P} (V + V_0).$$
(11.5)

A plot of the left hand side against $V + V_0$ should be linear. At constant rate of filtration, Eq. (11.2) can be written

$$Q = \frac{V}{t} = \frac{AAP}{\mu(R_f + \alpha c V/A)}$$
(11.6)

and rearranged into the linear form

$$\frac{\Delta P}{Q} = \frac{\Delta P}{V/t} = \frac{\mu}{A} R_f + \frac{\mu \alpha c}{A^2} V.$$
(11.7)

The constants again are found from the intercept and slope of the linear plot of $\Delta P/Q$ against *V*.

After the constants have been determined, Eq. (11.7) can be employed to predict filtration performance under a variety of constant rate conditions. For instance, the slurry may be charged with a centrifugal pump with a known characteristic curve of output pressure against flow rate. Such curves often may be represented by parabolic relations, as in Example 11.2, where the data are fitted by an equation of the form

$$P = a - Q(b + cQ).$$
 (11.8)

The time required for a specified amount of filtrate is found by integration of

$$t = \frac{V}{I_0} \frac{dV}{Q}.$$
 (11.9)

	Product Parameters		Fe	Feed Conditions Favoring Use		Equip	Equipment Characteristics			Direct Costs		
	Solids in Liquid Product	Liquid in Solid Product	Wash* Possibilities	Solids Concentration	Solids Density	Particle Size	Power	Space	Holdup	Initial	Operating	Maintenance
Filtration			đ									
Vacuum drum filter	F	G	E	high to med.	—	medium	high	medium	medium	high	high	medium
Disc filters	F	G	P to F	m e d i u m		fine	high	medium	medium	med. to high	high	m e d i u m
Horizontal filter	F	G	G to E ^d	high to med	-	coarse	high	medium	medium	medium	high	medium
Precoat filter	E	P**	P to F**	very low	-	slimy	high to med	medium	medium	high	very high	m e d i u m
Leaf (Kelly) filter Sedimentation	G to E ^d	F	F to G	l o w	-	fine, slimy	med. to low	medium	medium	m e d i u m	very high	m e d i u m
Thiskses	C to F	Р	n		4	modium	low	Von	Von	mad to	Law	vory low
Inickener	GTOE	'	r	mearum	aense	meurum	10 w	high	high	low	10 W	very low
Clarifier	G	Р	very P	low	med. dense	fine	very low	very high	very high	med.to low	low	very low
Classifier	Р	Ρ	P to F	m e d i u m	dense	coarse	low	high	high	med.to	low	l o w
Centrifugation												
Disc	F to G	P	Р	low to med.	medium	fine	high	low	low	high	high	high
Solid bowl	Р	F	Pto F	med. to high	medium	med.to fine	high	low	low	med.to high	high	high
gasket	P to F	E	Ed	med. to high		coarse	high	low	low	medium	high	high
Liquid cyclones												
Large	Р	Pto F	P	low to med.	high	medium	med.to low	low	low	very low	medium	high
Small multiple	Pto F	P	very P	l o w	med.to high	fine	med.to	low	low	low	medium	m e d i u m
Screens	Р	P to F	P	med. to high		coarse to med.	low	very	very	very	medium	med. to high
Ultrafiltration	E	P to F	Р	l o w	-	very fine	med. to high	high	high	high	high	very high

^a P = Poor. F = Fair. G = Good. E = Excellent. • Decantation wash always possible. ^d Displacement wash feasible. **Solids product contaminated by precoat material. (Purchas, 1981).



Figure 11.2. Experimental routine for aiding the selection of solid-liquid separation equipment (Davies, 1965).

Action On	Technique	Effects
1. Liquid	 heating dilution with solvent degassing and stripping 	reduction of viscosity, thereby speeding filtration and settling rates and reducing cake moisture content prevents gas bubbles forming within the
2. Solid particles	1. coagulation by chemical additives	medium or cake and impeding filtration destabilizes colloidal suspensions, allow- ing particles to agglomerate into microflocs
	2. flocculation by natural or forced convection	microflocs are brought into contact with each other to permit further agglomera- tion into large flocs
	3. aging	size of individual particles increases, e.g., by crystal growth
3. Concentration of solids	 increase by appropriate first-stage device such as settling tank, cyclone flotation cell or filter/thickener 	rate of filtration increased, especially if initial concentration 12%
	2. classify to eliminate fines, using sedimentation	rate of filtration increased and cake moisture content reduced
	 add filter powder (e.g., diatomite) or other solids to act as 'body aid' 	rate of filtration increased by more porous cake and possibly by high total solid concentration
4. Solid/liquid interaction	1. heat treatment, e.g., Porteus process involving pressure	physical methods which condition sludge
	2. freeze/thaw 3. ultrasonics	and induce coagulation and/or flocculation
	5. addition of wetting agents	reduces the interfacial surface tension, improves the draining characteristics of the cake, and decreases the residual moisture content

TABLE 11.4. Action and Effects of Slurry Pretreatments

(Purchas, 1981).

TABLE 11.5. Natures and Applications of Typical Flocculants

Trade Name	Composition	Typ eor Mech anism	Typical Application	Normal Range of pH Effectiveness	Normal Effective Concentration	Approx Price per lb	Manufacturer
Alum	$Al_2(SO_4)_3$ ·XH ₂ O	electrolytic and coagulation	water treatment	5-10	15 ppm	2¢	inorganic chemical manufacturers
Ferric sulfate	Fe ₃ (SO ₄)XH ₂ O	electrolytic coagulation	water treatment and chemical processing	any	5-100 ppm	2¢	inorganic chemical manufacturers
Sodium CMC	sodium carboxy- methylcellulose	coagulation and bridging	mineral processing	3-9	0.03-0.5 lb/ton	50¢	Hercules, DuPont
Kelgin W	algins	coagulation and bridging	water treatment	4-11	up to 5 ppm	\$1.50	Kelco Co.
Separan	acrylamide polymer	bridging	chemical processing	2-10	0.2-10 ppm	\$1.00-\$2.00	Dow Chemical Co.
Fibrefloc	animal glue	electrolytic	waste treatment	I - 9	5-30 ppm	18¢	Armour and Co.
Corn starch	corn starch	bridging	mineral processing	2-10	10 lb/ton	7¢	_
Polynox	polyethylene oxide	bridging	chemical processing	2-10	I-50 ppm	\$2.00	Union Carbide
Silica sol	activated silica sol	electrolytic coagulation	waste treatment	4-6	l-20 ppm	1,5¢as sodium silicate	inorganic chemical manufacturers
Sodium aluminate	sodium aluminate	coagulation	water treatment	3-12	2-10 ppm	10¢	National Aluminate
Guar gum	guar gum	bridging	mineral processing	2-12	0.02-0.3 lb/ton	35¢	General Mills
Sulfuric acid	H₂SO₄	electrolytic	waste treatment	1-5	highly variable	1¢	inorganic chemical manufacturers

^a 1966 prices, for comparison only. (Purchas, 1981). EXAMPLE 11.1 Constants of the Filtration Equation from Test Data Filtration tests were performed on a CaCO₃ slurry with these properties:

C = 135 kg solid/m³ liquid, $\mu = 0.001$ N sec/m².

The area of the filter leaf was 500 cm^2 . Data were taken of the volume of the filtrate (L) against time (sec) at pressures of 0.5 and 0.8 bar. The results will be analyzed for the filtration parameters:

		0	. 5 bar	0	. 8 bar
(L)	V/A	t	t/(V/A)	t	t/(V/A)
0.5	0.01	6.8	680	4.8	480
1	0.02	19.0	950	12.6	630
1.5	0.03	36.4	1213	22.8	760
2	0.04	53.4	1335	35.6	890
2.5	0.05	76.0	1520	50.5	1010
3	0.06	102.0	1700	69.0	1150
3.5	0.07	131.2	1874	88.2	1260
4	0.08	163.0	2038	112.0	1400
4.5	0.09			-	_
5	0.10			165. 0	1650

The units of V/A are m^3/m^2 . Equation (11.2) is

$$\frac{d(V/A)}{dt} = \frac{AP}{\mu(R_f + \alpha CV/A)}$$

whose integral may be written

$$\frac{R_f}{\Delta P/\mu} + \frac{\alpha C}{2(\Delta P/\mu)} \frac{v}{A} = \frac{t}{V/A}$$

Intercepts and slopes are read off the linear plots. At 0.5 bar,

$$\Delta P/\mu = 0.5(10^5)/0.001 = 0.5(10^8),$$

$$R_f = 600\Delta P/\mu = 3.0(10^{10}) \text{ m}^{-1},$$

Basic filtration Eq. (11.2) is solved for the amount of filtrate,

$$V = \frac{A}{\mu c \alpha} \left(\frac{A \Delta P}{Q} - \mu R_f \right). \tag{11.10}$$

Equations (11.8) and (11.10) are solved simultaneously for AP and Q at specified values of V and the results tabulated so:



Integration is accomplished numerically with the Simpson or trapezoidal rules. This method is applied in Example 11.2.

When the filtrate contains dissolved substances that should not remain in the filter cake, the occluded filtrate is blown out; then the cake is washed by pumping water through it. Theoretically, an amount of wash equal to the volume of the pores should be sufficient, even without blowing with air. In practice, however, only $\alpha = [18,000(2)/C]\Delta P/\mu = 36,000(0.5)(10^8)/135$ = 1.333(10¹⁰) m/kg.

At 0.8 bar,

$$\Delta P/\mu = 0.8(10^8),$$

 $R_f = 375(0.8)(10^8) = 3(10^{10}) \text{ m}^{-1},$
 $\alpha = 12,750(2)(0.8)(10^8)/135 = 1.511(10^{10}) \text{ m/kg}$

Fit the data with Almy-Lewis equation, Eq. (11.24),

$$\alpha = kp'',$$

$$n = \frac{\ln(\alpha_1/\alpha_2)}{\ln(P_1/P_2)} = \frac{\ln(1.511/1.333)}{\ln(0.8/0.5)} = 0.2664$$

$$k = 1.511(10^{10})/0.8^{0.2664} = 1.604(10^{10}),$$

$$\alpha = 1.604(10^{10})P^{0.2664}, \quad \text{m/kg}, P \text{ in bar.}$$



30–85% of the retained filtrate has been found removed by one-displacement wash. Figure 11.3(b) is the result of one such test. A detailed review of the washing problem has been made by **Wakeman** (1981, pp. 408-451).

The equations of this section are applied in Example 11.3 to the sizing of a continuous rotary vacuum filter that employs a washing operation.

COMPRESSIBLE CAKES

or

Resistivity of filter cakes depends on the conditions of formation of which the pressure is the major one that has been investigated at length. The background of this **topic** is discussed in Section 11.3, but here the pressure dependence will be incorporated in the filtration equations. Either of two forms of pressure usually is taken,

$$\boldsymbol{\alpha} = \boldsymbol{\alpha}_0 \boldsymbol{P^n} \tag{11.11}$$

$$\alpha = \alpha_0 (1 + kP)^n. \tag{11.12}$$

EXAMPLE 11.2

Filtration Process with a Centrifugal Charge Pump

A filter press with a surface of 50m²handles a slurry with these properties:

$$\mu = 0.001 \text{ N sec/m}^2,$$

C = 10 kg/m³,
 $\alpha = 1.1(10^{11}) \text{ m/kg},$
R_f = 6.5(10¹⁰) m-i.

The feed pump is a centrifugal with a characteristic curve represented by the equation

$$AP = 2 - Q(0.00163Q - 0.02889), \quad bar \tag{1}$$

with Q in m^3 hr. Find (a) the time required to obtain $50 m^3$ of filtrate; (b) the volume, flow rate, and pressure profiles. Equation (11.2) of the text solved for V becomes

$$V = \frac{A}{\alpha\mu C} \left(\frac{A\Delta P}{Q} - \mu R_f \right) = \frac{50}{1.1(10^9)} \left[\frac{50(10^5)\Delta P}{Q/3600} - 6.5(10^7) \right]$$
$$= 818.1 \left(\frac{\Delta P}{Q} - 0.0036 \right)$$
(2)

Equations (1) and (2) are solved simultaneously to obtain the tabulated data. The time is found by integration with the

The first of these does not extrapolate properly to resistivity at low pressures, but often it is as adequate as the more complex one over practical ranges of pressure.

Since the drag pressure acting on the particles of the cake varies from zero at the face to the full hydraulic pressure at the filter cloth, the resistivity as a function of pressure likewise varies along the cake. A mean value is defined by

$$\frac{1}{\bar{\alpha}} \simeq \left(\frac{1}{\alpha}\right)_{\text{mean}} = \frac{1}{\Delta P_c} \int_0^{\Delta P_c} \frac{dP}{\alpha} \frac{1}{\Delta P} \int_0^{\Delta P} \frac{dP}{\alpha}, \qquad (11.13)$$

where ΔP_c is the pressure drop through the cake alone. In view of the roughness of the usual correlations, it is adequate to use the overall pressure drop as the upper limit instead of the drop through the cake alone.

With Eq. (11.12) the mean value becomes

$$\bar{\alpha} = \frac{\alpha_0 k (1 - n) \Delta P}{(1 + k \Delta P)^{1 - n} - 1}.$$
(11.14)

The constants α_0 , k, and n are determined most simply in compression-permeability cells as explained in Section 11.4, but those found from filtration data may be more appropriate because the mode of formation of a cake also affects its resistivity. Equations (11.14) and (11.2) together become

$$\frac{d(V/A)}{dt} = \frac{\Delta P}{\mu} \left[R_f + \frac{\alpha_0 c k (1-n) \Delta P}{(1+k \Delta P)^{n-1} - 1 A} \right]^{-1}, \quad (11.15)$$



which integrates at constant pressure into

$$\frac{2t}{V/A} = \frac{2\mu}{\Delta P} R_f + \frac{\alpha_0 c k \mu (1-n)}{(1+k\Delta P)^{1-n} - 1} (V/A)$$
(11.16)

The four unknown parameters are α_0 , *k*, *n*, and R_f . The left-hand side should vary linearly with *V/A*. Data obtained with at least three different pressures are needed for evaluation of the parameters, but the solution is not direct because the first three parameters are involved nonlinearly in the coefficient of *V/A*. The analysis of constant rate data likewise is not simple.

The mean resistivity at a particular pressure difference can be evaluated from a constant pressure run. From three such runs-AP,, ΔP_2 , and ΔP_3 —three values of the mean resistivity- $\bar{\alpha}_1, \bar{\alpha}_2$, and $\bar{\alpha}_3$ —can be determined with Eq. (11.2) and used to find the three constants of the expression for an overall mean value,

$$\bar{\alpha} = \alpha_0 (1 + k\Delta P)^n, \tag{11.17}$$

which is not the same as Eq. (11.12) but often is as satisfactory a representation of resistivity under practical filtration conditions. Substituting Eq. (11.17) into Eq. (11.2), the result is

$$\frac{d(V/A)}{dt} = \frac{AP}{\mu[R_f + \alpha_0 c(1 + k\Delta P)^n(V/A)]}$$
(11.18)

Integration at constant pressure gives the result

$$\frac{\alpha_0 c\mu (1+k\Delta P)^n}{2\Delta P} \frac{V}{A} + \mu R_f / \Delta P = \frac{t}{V/A}.$$
(11.19)



Figure 11.3. Laboratory test data with a vacuum leaf filter. (a) Rates of formation of dry cake and filtrate. (b) Washing efficiency. (c) Air flow rate vs. drying time. (d) Correlation of moisture content with the air rate, pressure difference AP, cake amount W lb/sqft, drying time θ_d min and viscosity of liquid (*Dahlstrom and Silverblatt*, 1977).

EXAMPLE 11.3 Rotary Vacuum Filter Operation A TiO₂ slurry has the properties

> c = 200 kg solid/m² liquid, ρ_s = 4270 kg/m³, μ = 0.001/3600 N hr/m², α = 1.6(*E*12) m/kg (item 4 of Fig. 11.2), ε = 0.6.

Cloth resistance is $R_f = 1(E10) \text{ m}^{-1}$. Normal peripheral speed is about 1 m/min. Filtering surface is 1/3 of the drum surface and washing surface is 1/6 of the drum surface. The amount of wash equals the pore space of the cake. The cake thickness is to be limited to 1 cm. At suitable operating pressures, find the drum

speed in rph and the drum diameter:

cake thickness = 0.01 m =
$$\frac{c}{\rho_s(1-\varepsilon)} \frac{V_f}{A}$$

= $\frac{200}{4270(0.4)} \frac{V_f}{A}$,
 $\frac{V_f}{A} = \frac{0.01(4270)(0.4)}{200} = 0.0854 \text{ m}^3/\text{m}^2$
wash liquid = pore volume
= $0.01(0.6) = 0.006 \text{ m}^2 \text{m}^2$. (1)

With the pressure difference in bar,

$$\frac{d(V/A)}{dt} = \frac{10^{5} \Delta P_{b}}{(0.001/3600)[10^{10} + 160(10^{10})V/A]}$$
$$= \frac{36 \Delta P_{h}}{1 + 160V/A}.$$
(2)

The integral at constant pressure is

$$80(V_f/A)^2 + V_f/A = 36\Delta P_b t_f$$
(3)

With $V_f/A = 0.0854$,

$$\begin{aligned} \Delta P_b t_f &= 0.01858, \\ t_f &= 0.01858 / \Delta P_b = 1/3 \dot{n}_f \\ \dot{n}_f &= 17.94 \Delta P_b, \end{aligned} \tag{4}$$

where \dot{n}_{f} is the rph speed needed to make the 1 cm thick cake. From Eq. (2) the washing rate is

$$r_{w} = \frac{36\Delta P_{b}}{1^{+} 160(0.0854)} = 2.455\Delta P_{b}.$$
 (6)

Washing time:

$$t_{w} = \frac{0.006}{2.455\Delta P_{b}} = \frac{0.00244}{\Delta P_{b}} \ge \frac{1}{\dot{n}_{w}},\tag{7}$$

Eq. (11.19) could be written in terms of $\tilde{\alpha}$ from Eq. (11.17) and would then have the same form as Eq. (11.2), but with only R_f as a parameter to be found from a single run at constant pressure. In Example 11.1, the mean resistivity is found from the simpler equation

$$\bar{\alpha} = \alpha_0 (\Delta P)^n. \tag{11.20}$$

Analysis of the filtration of a compressible material is treated in Example 11.4.

11.3. RESISTANCE TO FILTRATION

The filtration equation

$$\frac{Q}{A} = \frac{\Delta P}{\mu(R_f + \alpha c V/A)}$$
(11.2)

considers the overall resistance to flow of filtrate to be made up of contributions from the filter medium R_f , and from the cake with specific resistance α .

FILTER MEDIUM

In practice, a measured R_f includes the effects of all factors that are independent of the amount of the cake; in a plate-and-frame press, for instance, piping and entrance and exit losses will be included, although most of the resistance usually is due to the medium itself. Aging and the resulting increase in resistance is a recognized behavior, particularly of media made of fibers. Particles are gradually occluded in the media so thoroughly that periodic cleaning cannot restore the original condition. The degree of penetration of the medium depends on the porosity, the pore sizes, particles sizes, and velocity. Normally R_f is found to depend on the operating pressure; on plots like those of Example 11.1, the two intercepts may correspond to different values of R_f at the two pressures.

Data for some filter media are shown in Table 11.6. Although these porosities and permeabilities are of unused materials, the relative values may be useful for comparing behaviors under filtration conditions. Permeability K_p normally is the property

$$\dot{n}_w \le 68.3 \Delta P_b \tag{8}$$

Comparing (5) and (8), it appears that an rph to meet the filtering requirements is 68.3/17.94 = 3.8 times that for washing and is the controlling speed.

With a peripheral speed of 60 m/hr

$$60 = \pi Dn, D = 60/\pi n = 19.1/\dot{n}.$$
(9)

The parameters at several pressures are

ΔP_b (bar)	0.2	0.4	0.6	0.8
ń (rph)	3. 59	7.18	10.76	14.35
D (m)	5.3	2.66	1.78	1.33

If the peripheral speed were made 1.22 m/min, a drum 1.0 m dia would meet the requirements with AP = 0.8 bar. Another controllable feature is the extent of immersion which can be made greater or less than 1/3. Sketches of a rotary vacuum filter are in Figure 11.12.

reported rather than the resistivity that has been discussed here. It is defined by the equation

$$Q/A = K_{p} \Delta P/\mu L, \qquad (11.21)$$

where L is the thickness. The relation to the resistivity is

$$\boldsymbol{R}_{\boldsymbol{p}} = \boldsymbol{L}/\boldsymbol{K}_{\boldsymbol{p}}.\tag{11.22}$$

Thus the filtration resistivity of the medium includes its thickness. Typical measured values of R_f are of the order of 10^{10} m⁻¹; for comparison, the fine filter sheet of Table 1.6, assuming it to be 1 mm thick, has $L/K_r = 0.001/0.15(10^{-12}) = 0.7(10^{10})$ m⁻¹.

CAKE RESISTIVITY

A fundamental relation for the flow resistance of a bed of particles is due to Kozeny (*Ber. Wien. Akad.* 135a, 1927, 271-278):

$$\alpha = Ks_0^2 (1 - \varepsilon) / \varepsilon^3, \tag{11.23}$$

K = approximately 5 at low porosities,

- s_0 = specific surface of the particles,
- ρ_s = density of the particles,
- $\boldsymbol{\varepsilon}$ = porosity, volume voids/volume of cake.

Because the structure of a cake is highly dependent on operating conditions and its history, the Kozeny equation is only of qualitative value to filtration theory by giving directional effects.

At increasing pressures, the particles or aggregates may be distorted and brought closer together. The rate of flow also may affect the structure of a cake: at low rates a loose structure is formed, at higher ones fine particles are dragged into the previously formed bed. The drag pressure at a point in a cake is the difference between the pressure at the filter medium and the pressure loss due to friction up to that point. As the drag pressure at a distance from the filter cloth increases, even at constant filtering pressure, the porosity and resistance adjust themselves continuously. Figure 11.4(a) shows such effects of slurry concentration and filtering rates

EXAMPLE 11.4 Filtration and Washing of a Compressible Material A kaolin slurry has the properties

c = 200 kg solid/m³ filtrate, μ = 0.001 N sec/m², 2.78(*E* - 7) N hr/m², ρ_s = 200 kg/m³, α = 87(*E*10)(1 + *P*/3.45)^{0.7} m/kg with *P* in bar, ε = 1 0.460(1 + *P*/3.45)^{0.12}.

The equations for α and ε are taken from Table 11.8.

Filtration will proceed at a constant rate for 15 min, the pressure will rise to 8 bar and filtration will continue at this pressure until the end of the operation. Filter cloth resistance is $R_f = 1(10^{10}) \text{ m}^{-1}$. The down time per batch is 1 hr.

a. Find the maximum daily production of filtrate.

b. The filtrate will be blown and then washed with a volume of water equal to the pore space of the cake. Find the maximum daily production of filtrate under these conditions.

Part (a)

Basis 1 m² of filtering surface. At P = 8 bar, or 8(10⁵) Pa

 $\begin{aligned} \alpha &= 87(10^{10})(1+8/3.45)^{0.7} = 2.015(10^{12}) \text{ m/kg}, \\ \varepsilon &= 1 - 0.46(1+8/3.45)^{0.12} = 0.47, \\ \mu c \alpha &= (0.001/3600)(200)(2.015)(10^{12}) = 1.12(10^8) \text{ N hr/m}^4. \end{aligned}$

The filtration equation (11.2) is

 $\frac{dV}{dt} = \frac{AAP}{\mu(R_f + \alpha CV/A) = (0.001/3600)[10^{10} + 2.015(10^{12})(200)V]}$ $= \frac{AP}{2780 + 1.12(10^8)V}.$

The rate when t = 0.25h and AP = 8(10⁵) Pa,

$$Q = \frac{8(10^3)}{2780 + 1.12(10^8)Qt} = \frac{8(10^3)}{2780 + 0.28(10^8)Q} = 0.1691 \text{ m}^3/\text{m}^2 \text{ hr.}$$

The amount of filtrate at this time is

 $V_0 = Qt = 0.1691(0.25) = 0.0423 \text{ m}^3.$

on the parameters of the correlating equation

 $\alpha = \alpha_0 (\Delta P)^n.$

The integral of the rate equation at constant P is

$$2780(V_f - 0.0423) + 0.56(10^8)(V_f^2 - (0.00423)^2] = 8(10^5)(t_f - 0.25).$$

Filtering period is

$$t_f = 0.25 + 0.0035(V_f - 0.0423) + 70.0(V_f^2 - 0.0018).$$

Daily production rate,

$$R_{d} = (\text{no of batches/day})(\text{filtrate/batch})$$

$$= \frac{24V_{f}}{t_{d} + t_{f}} = \frac{24V_{f}}{1 + t_{f}}, \text{ m}^{3}/(\text{m}^{2})(\text{day})$$

$$= \frac{24V_{f}}{1.25 + 0.0035(V_{f} - 0.0423) + 70(V_{f}^{2} - 0.0018)}$$

The tabulation shows that R_d is a max when $V_f = 0.127$.

	ţ,		R _d
		1.3507	
		1.3526	
1	. 2533	1.3527	(max)
		1.3526	
		1.3525	
		1.3522	

Part(b)

Amount of wash liquid =
$$\frac{cV_f\varepsilon}{\rho_s(1-\varepsilon)} = \frac{200(0.47)}{2500(0.53)} = 0.0709V_f$$

wash rate = filtering rate at the conclusion of the filtration

$$= \frac{\Delta p}{\mu(R_f + \alpha cV_f)} = \frac{8(10^5)}{2780 + 1.12(10^8)V_f}, \quad \text{m}^3/\text{hr},$$

$$t_w = \text{wash time} = \frac{0.709V_f[2780 + 1.12(10^8)V_f]}{8(10^5)}$$

$$= V_f(0.000246 + 9.926V_f),$$

$$R_d = \frac{24V_f}{1 + t_f + t_w}$$

$$= \frac{24V_f}{[1 + 0.0035(V - 0.0423) + 7010(V_f^2 - 0.0018) + V_f(0.000246 + 9.926V_f)]}.$$

The optimum operation is found by trial:

$$V_f = 0.105,$$

 $t_f = 1.0805,$
 $t_w = 0.1095,$
 $R_d = 1.1507$ (max), daily production rate.

COMPRESSIBILITY-PERMEABILITY (CP) CELL MEASUREMENTS

The measurements were obtained with a small filter press. Clearly, the resistivity measured at a particular rate is hardly applicable to predicting performance at another rate or at constant pressure.

(11.24)

The probable success of correlation of cake resistivity in terms of all the factors that have been mentioned has not been great enough to have induced any serious attempts of this nature, but the effect of pressure has been explored. Although the α 's can be deduced from

Porosity (%)	
Wedge wire screen	5-10
Perforated sheet	20
Wire mesh:	
Twill weave	15-25
Square	30-35
Porous plastics, metals, ceramics	30-50
Crude kieselguhr	50-60
Porous ceramic, special	70
Membranes, plastic foam	80
Asbestos/cellulose sheets	80
Refined filter aids (diatomaceous earth ex-	80-90
panded perlite)	
Paper	60-95
Scott plastic foam	97
Permeability, $10^{12} K_p$ (m ²) (compare Eq. (11.22))	
Filter aids	
Fine	0.05-0.5
Medium	1-2
Coarse	4-5
Cellulose fibre pulp	1.86
Cellulose fibre + 5% asbestos	0.34
Filter sheets	
Polishing	0.017
Fine	0.15
Clarifying	1.13
Sintered metal	
3 μm pore size	0.20
8 μm pore size	1.0
28 μm pore size	7.5
75 μm pore size	70

TABLE 11.6. Porosities and Permeabilities of Some Filter Media

(Purchas, 1981).

filtration experiments, as done in Example 11.1, a simpler method is to measure them in a CP cell as described briefly later in this chapter. Equation (11.24) for the effect of pressure was proposed by **Almy** and Lewis (1912). For the materials of Figure 1.2(b), for instance, it seems to be applicable over at least moderate stretches of pressure. Incidentally, these resistances are not represented well by the Kozeny porosity function $(1 - \varepsilon)/\varepsilon^3$; for substance 6, the ratio of resistivities at 100 and 1 psia is 22 and the ratio of the porosity functions is 2.6. The data of Table 11.7 also show a substantial effect of pressure on resistivity.

Since the drag pressure varies along the cake as a result of friction, porosity and resistivity also will vary with position. Figure 11.5 shows such data at three different overall pressures. The axial profile of the normalized pressure, $P_{\text{local}}/P_{\text{face}}$, appears to be a unique function of fractional distance along the cake, independent of the filtering pressure. The resistivity will vary along the cake just as the porosity does. As the cake builds up, moreover, the drag pressure, porosity, and resistivity at a particular distance from the filter medium also will vary. Consequently, since the resistivity does not necessarily change linearly with position, any mean value also is likely to vary as the cake builds up. Thus, in the filtration equation even a mean value of α has to be expressed as a function of P and V. The proper mathematical representation of a filtration process is by means of an integro-differential equation with a moving boundary (the face of the cake). Such an analysis was made by Wakeman (1978) and a similar one by Tiller, Crump, and Ville (1979). At present, unfortunately, such a mathematical approach to filtration problems is more of academic than practical value. One of the factors that is not taken into account is the effect of flow rate on

the formation and stability of loose cake structures; such behavior normally is not reproducible.

ANOTHER FORM OF PRESSURE DEPENDENCE

Equation (11.24) cannot be entirely valid because it predicts zero resistivity at zero pressure, whereas cakes do have structures and significant resistivities even at minimal operating pressures. Modified Eq. (11.12) is extrapolatable, and is rewritten here as

$$\boldsymbol{\alpha} = \boldsymbol{\alpha}_0 (1 + \boldsymbol{k} \boldsymbol{P}) \tag{11.25}$$

with a similar one for porosity

$$\boldsymbol{\varepsilon} = 1 - (1 - \boldsymbol{\varepsilon}_0)(1 + \boldsymbol{k}\boldsymbol{P})^n. \tag{11.26}$$

Some data fitted to these equations by Tiller et al. (1979) are in Table 11.8; here the constant k is the same for both α and ε , although this is not necessarily generally the case. Unfortunately, these data show that the parameters are not independent of the pressure range. Apparently the correlation problem has not been solved. Perhaps it can be concluded that insofar as the existing filtration theory is applicable to real filtering behavior, the approximation of Almy and Lewis may be adequate over the moderate ranges or pressures that are used commonly, somewhere between 0.5 and 5 atm.

PRETREATMENT OF SLURRIES

Since the sizes of particles and agglomerates of the slurry are a main determinant of a rate of filtration, any methods of influencing these sizes are of great practical value. For example, Figures 1.2(b) and (c) show $CaCO_3$ and TiO_2 each to be precipitated at two different values of pH with resultant great differences in resistivity and porosity. At 10 psia, for instance, the resistivities of the two $CaCO_3$'s are in the ratio of 5, with corresponding differences in rate of filtration. Pretreatment of a slurry to enhance coagulation and particle growth is an important aspect of filter process design. Another method of long standing for improving filtration behavior is the formation of an open cake structure by addition of relatively large and rigid particles of a filter aid. The common methods of pretreatment are listed in Table 11.4, and some chemical flocculants that are of practical value are described in Table 11.5. These effects cannot be predicted safely and must be measured.

11.4. THICKENING AND CLARIFYING

When dilute slurries are encountered on a large scale, it is more economical to concentrate them before filtering. This is accomplished by sedimentation or thickening in tanks for an appropriate period. Typical designs of thickeners are sketched in Figure 11.6. The slurry is introduced at the top center, clear liquid **overflows** the top edge, whereas the solids settle out and are worked gradually towards the center with slowly rotating rakes towards the discharge port at the bottom center. The concentrated slurry then is suitable for tiltration or other further processing. Clarifiers are similar devices, primarily for recovering clear liquids from dilute suspensions. Some characteristics of sedimentation equipment are given in Table 11.3 and typical applications are listed in Table 11.9 and 14.7. Sedimentation rates often are assisted by addition of flocculating agents, some of which are listed in Table 11.5. Specifically, pilot plant testing is advisable when

- 1. The expecting filtering area is expected to be substantial, measured in tens of \mathbf{m}^2 .
- 2. Cake washing is critical.



Figure 11.4. Data of compressibilities and porosities of filter cakes. (a) Parameters of the correlation $\alpha = \alpha_0 (\Delta P)^n$ for resistivity of CaSiO₃ filter cakes at two rates and two concentrations (*Rushton and Katsoulas*, 1984). (b) Resistivity as a function of pressure measured in a compressibility-permeability (CP) cell [Grace, Chem. Eng. Prog. 49, 303, 367, 427 (1953)]. (c) P orosity as a function of pressure for the same six materials (Grace, Zoc. cit.).

3. Cake drying is critical.

4. Cake removal may be a problem.

5. Precoating may be needed.

11.5. LABORATORY TESTING AND SCALE-UP

Laboratory filtration investigations are of three main kinds:

- 1. observation of sedimentation rates;
- 2. with small vacuum or pressure leaf filters;
- 3. with pilot plant equipment of the types expected to be suitable for the plant.

Sedimentation tests are of value particularly for rapid evaluation of the effects of aging, **flocculants**, vibration, and any other variables that conceivably could affect a rate of filtration. The results may suggest what kinds of equipment to exclude from further consideration and what kind is likely to be worth investigating. For instance, if sedimentation is very rapid, vertical leaves are excluded, and top feed drums or horizontal belts are indicated; or it may be indicated that the slurry should be **preconcentrated** in a thickener before going to filtration. If the settling is very slow, the use of filter aids may be required, etc. Figure 11.1 illustrates typical sedimentation behavior. Figure 11.2 summarizes an experimental routine.

Vacuum and pressure laboratory filtration assemblies are shown in Figure 11.7. Mild agitation with air sometimes may be preferable to the mechanical stirrer shown, but it is important that any agglomerates of particles be kept merely in suspension and not broken up. The test record sheet of Figure 11.8 shows the kind of data that normally are of interest. Besides measurements of filtrate and cake amounts as functions of time and pressure, it is desirable

TABLE 11.7	. Specific	Resistances	of	Some	Filter
	Cakes				

Material	Filtration Pressure psi	Resistance SI Units, m/kg
High grade kieselguhr	_	1.64 x 10 ⁹
Ordinary kieselguhr	2 5	1.15x IO"
	100	1.31 x 10¹¹
Carboraffin charcoal	1.4	3.14 x 10 ¹⁰
	10	5.84 x 10 ¹⁰
Calcium carbonate	2 5	2.21 x 10 ¹¹
(precipitated)	100	2.68×10^{11}
Ferric oxide (pigment)	2 5	8.04 x 10"
	100	14.12 x 10"
Mica clay	2 5	4.81 x 10"
	100	8.63 $ imes$ 10"
Colloidal clay	2 5	5.10 x 10 ¹²
	100	$6.47 imes 10^{12}$
Magnesium hydroxide	2 5	3.24 x 10 ¹²
(gelatinous)	100	6.97 x 10 ¹²
Aluminium hydroxide	2 5	2.16 x 10 ¹³
(gelatinous)	100	4.02×10^{13}
Ferric hydroxide	2 5	1.47 x 10 ¹³
(gelatinous)	100	4.51 x 10 ¹³
Thixotropic mud	80	$6.77 imes 10^{14}$
Theoretical figures for		
rigid spheres:		
d= 10 μm		6.37 × 10 ⁹
$d = 1 \mu m$		6.37 × 10"
<i>d</i> = 0.1 pm		6.37 × 1 0 ¹³

(Carman, 1938).

to test washing rates and efficiencies and rates of moisture removal with air blowing. Typical data of these kinds are shown in Figure 11.3. Detailed laboratory procedures are explained by Bosley (1977) and Dahlstrom and Silverblatt (1977). Test and scale-up procedures for all kinds of SLS equipment are treated in the book edited by Purchas (1977).

Before any SLS equipment of substantial size is finally selected, it is essential to use the results of pilot plant tests for guidance. Although many vendors are in a position to do such work, pilot equipment should be used at the plant site where the slurry is made. Because slurries often are unstable, tests on shipments of slurry to the vendors pilot plant may give misleading results. It may be possible to condition a test slurry to have a maximum possible resistivity, but a plant design based on such data will have an unknown safety factor and may prove uneconomical.

COMPRESSION-PERMEABILITY CELL

Such equipment consists of a hollow cylinder fitted with a permeable bottom and a permeable piston under controlled pressure. Slurry is charged to the slurry, cake is formed with gentle suction, and the piston is lowered to the cake level. The rate of flow of filtrate at low head through the compressed cake is measured at a series of pressures on the piston. From the results the resistivity of the cake becomes known as a function of pressure. The data of Figures 11.4(b) and (c) were obtained this way; those of Figure 11.4(a) by filtration tests.

There is much evidence, however, that the resistivity behavior of a cake under filtration conditions may be different from that measured in a CP cell. The literature is reviewed by **Wakeman** (1978). CP cell data are easily obtained and may be of value in a qualitative sense as an indication of the sensitivity of resistivity to pressure, but apparently are not of acceptable engineering accuracy for the design of filtration equipment. The deduction of resistivities from filtration tests is illustrated in Example 11.1.

THE SCFT CONCEPT

No serious attempt has yet been made to standardize filtration tests and to categorize filtration behavior in generally accepted terms. A possibly useful measure of filterability, however, has been proposed by Purchas (1977; 1981). The time in minutes required to form a cake 1 cm thick when the cell is operated with a differential of 500 **Torr** (0.67 bar) is called the Standard Cake Formation Time (SCFT), t_F . The pressure of 500Torr is selected because it is obtained easily with common laboratory equipment. The procedure suggested is to make a series of tests at several cake thicknesses and to obtain the SCFT by interpolation, rather than to interrupt a single test to make observations of cake thickness. A direct relation exists, of course, between the SCFT and resistivity α ; some examples are

Material	a (m/kg)	SCFT t _F (min)
Filter aid	1.64(E9)	0.26
CaCO ₃	2.21 (EI 1)	34.6
Colloidal clay	5.10(E12)	798

Full scale filtration equipment requirements can be estimated quickly in terms of t_F . For instance, when the resistance of the filter medium is neglected, the constant pressure Eq. (11.3) may be written as

$$\Delta Pt = \frac{\alpha c}{2} \left(\frac{V}{A}\right)^2 = \frac{\alpha c}{2} \left[\frac{(1-\varepsilon)L}{c}\right]^2, \qquad (11.27)$$

where L is the thickness of the cake in meters. Upon rationing in



Figure 11.5. Axial distribution of pressure and porosity of an ignition-plug clay measured in a CP cell. (a) Normalized pressure distribution as a function of normalized distance [(- - -) experimental filtration data; theoretical curves: (x) AP = 98 kN m⁻²; (\bigoplus) AP = 294 kN m⁻²; (A) ΔP = 883 kN m⁻²]. (b) Porosity distributions at three pressures. The curves are by Wakeman (1978).

the SCFT data for 0.01 m,

$$\frac{\Delta Pt}{0.67t_F} = (100L)^2, \tag{11.28}$$

with AP in bar. From this relation the filtering time can be found at a specified pressure and cake thickness and when t_F is known.

SCALE-UP

Sizing of full scale equipment on the basis of small scale tests requires a consideration of possible ranges of at least the following variables:

- 1. filterability as measured by cake and medium resistivity;
- 2. feed rate and concentration;
- operating conditions, particularly pressure and high initial rates;
 behavior of the filter cloth with time.

Safety factors for scale up from laboratory leaf tests are difficult to generalize. On the basis of pilot plant work, adjustments of ll-21% are made to plate-and-frame filter areas or rates, and 14-20% to continuous rotary filters, according to Table 1.4.

The performance of solid-liquid separation equipment is difficult to predict by the engineer without some specific experience in this area. Unfortunately, it must be again recommended that the advice of experienced vendors should be sought, as well as that of expert consultants.

11.6. ILLUSTRATIONS OF EQUIPMENT

Equipment for solid-liquid separation is available commercially from many sources. About 150 names and addresses of suppliers in the United States and abroad are listed by Purchas (1981). Classifications of vendors with respect to the kind of equipment are given, for instance, in *Chemical Engineering Catalog* (Reinhold, New York, annual) and in *Chemical Engineering Equipment Buyers Guide* (McGraw-Hill, New York, annual).

The variety of solid-liquid separation equipment is so great that only a brief selection can be presented here. The most extensive modern picture gallery is in the book of Purchas (1981). The older encyclopedia of Kieser (Spamer-Springer, Berlin, 1937, Vol. 2) has 250 illustrations in 130 pages of descriptions; the pictures do not appear to have aged particularly. Illustrations in manufacturers catalogs are definitive and often reveal the functioning as well as aspect of the equipment. The selected figures of this chapter are primarily line drawings that best reveal the functioning modes of the equipment.

Figure 11.9 shows two models of sand filters whose purpose is to remove small contents of solids from large quantities of liquids. The solids deposit both on the surface of the bed and throughout the bed. They are removed intermittently by shutting off the main

TABLE 11.8. Parameters of Equations for Resistivity a and Porosity *£* of Some Filter Cakes

 $\alpha = \alpha_0 \left(1 + \frac{p_s}{p_s}\right)^n$

$(1 - \varepsilon) = (1 - \varepsilon_0) \left(1 + \frac{\rho_a}{\rho_a}\right)^{\beta^*}$								
Material	Pressure range, kPa	ρ _a , kPa	¤₀, m kg-' [™] x 10 ^{−10}	n	$(1 - \varepsilon_0)$	β*		
CaCO ₃ (ref. 7)	3-480	1	11	0. 15	0. 209	0.06		
CaCO ₃ (ref. 8)	7- 550	7	5.1	0. 2	0. 225	0. 06		
	550- 7000	790	8.1	0. 9	0. 263	0. 22		
Darco-B (ref. 8)	7- 275	1.7	1.1	0.4	0. 129	0. 08		
	275- 7000	520	4.7	1.8	0. 180	0. 18		
Kaolin-Al ₂ SO ₄ (ref. 8)	7- 415	7	43	0. 3	0. 417	0. 04		
	415- 7000	345	87	0. 7	0. 460	0. 12		
Solka-Floc (ref. 8)	7- 275	2. 75	0. 00058	1.0	0. 132	0. 16		
	275- 7000	260	0. 13	2.0	0. 237	0. 26		
Tal c-C (ref. 8)	7- 1400	5.5	4.7	0.55	0. 155	0. 16		
	1400- 7000	1400	35	1.8	0. 339	0. 25		
TiO ₂ (ref. 8)	7-7000	7	18	0.35	0. 214	0.1		
Tungsten (ref. 8)	7- 480	7	0. 39	0. 15	0. 182	0. 05		
	480- 7000	520	0. 38	0. 9	0. 207	0. 22		
Hong Kong	1- 15	1	42	0. 35	0. 275	0. 09		
pink kaolin (ref. 9)	15- 1000	12	70	0. 55	0. 335	0. 1		
Gairome clay (ref. 10)	4-1000	3.4	370	0.55	0.309	0.09		

(Tiller et al, 1979)

flow and backwashing with liquid. The concentrated sludge then must be disposed of in some way. Beds of charcoal are employed similarly for clarification of some organic liquids; they combine adsorption and mechanical separation.

Clarification of a large variety of liquids is accomplished with cartridge filters which come in a large variety of designs. Usually the cartridges are small, but liquid rates in excess of **5000 gpm** have been designed for. The filtering surface may be a fine metal screen or an assembly of closely spaced disks whose edge face functions as the filtering surface, or woven or matted fibers. The operation is intermittent, with either flushing back of the accumulated solids or replacement of the filtering elements in the body of the cartridge, or in some instances the solids are scraped off the filtering surface with a built-in mechanism and then flushed out in concentrated form. The variety of cartridge filters are described in detail in books by Warring (1981), Purchas (1981), and Cheremisinoff and Azbel (1983). Table 11.10 is a selected list of some of their applications and the minimum sizes of particles that are removed.

Figure 11.6 is of two types of sedimentation equipment, and Figure 12.2(e) of another. They are used for clarifying a valuable liquid or for preparing a concentrated slurry for subsequent filtration. They depend on gravitational sedimentation. Removal is assisted by rake action, or by the conical sides of the vessel of Figure 11.6(b).

Figure 11.10 is of the main kinds of filters that can be operated at superatmospheric pressures which may be necessary with otherwise slow filtering slurries. Commercial sizes are listed in Table 11.11. They all operate on intermittent cycles of cake formation, washing, dewatering with air blowing and cake removal. The plate-and-frame design of Figure 11.10(a) is the most widely recognized type. In it, cake removal is effected after separating the plates. The horizontal plate design of Figure 11.10(b) is popular in smaller sizes under, 2ft dia or so; the plates are lifted out of the casing for cake removal. The other units all have fixed spacings between the leaves. From them the cakes may be blown back with air or flushed back or scraped off manually. The Vallez unit of Figure 11.10(f) ordinarily does not require the case to be opened for cleaning.

Figure 11.11 is of continuous horizontal filtering equipment that operate primarily with vacuum, although they could be housed in pressure-tight casings for operation at superatmospheric pressure or with volatile liquids. Both the belt and the rotary units are well suited to rapidly settling and free draining slurries. In comparison with rotary drum vacuum filters, the horizontal equipment of Figure 11.11(c) has the merit of more readily accessible piping, a real advantage from a servicing point of view.

Figure 11.12 represents the main kinds of rotary drum filters. Commercial sizes are listed in Table 11.14. The flowsketch of Figure 11.12(a) identifies the main auxiliaries required for this kind of filtration process. Feed to the drum may be dip-type as in Figure 11.12(b), but top feed designs also are widely used. The unit with internal filtering surface of Figure 11.12(c) is suited particularly to rapidly settling solids and has been adapted to pressure operation.

Cake removal usually is with a scraper into a screw or belt conveyor, but Figure 11.12(d) depicts the use of a drum with a filtering belt that is subject to a continual cleaning process. Some filters have a multi parallel string discharge assembly whose path follows that of the belt shown.

The double drum filter of Figure 11.12(e) has obvious merit particularly when top feeding is desirable but it is not used widely nowadays. Disk filters of the type of Figure 11.12(f) are the most widely used rotary type when washing of the cake is not necessary.

Figure 11.13 is of a variety of devices that utilize centrifugal force to aid in the separation of solid and liquid mixtures. Figure



Figure 11.6. Thickeners for preconcentration of feed to filters or for disposal of solid wastes [see also the rake classifier of Fig. 12.2(e)]. (a) A thickener for concentrating slurries on a large scale. The rakes rotate slowly and move settled solids towards the discharge port at the center. Performance data are in Table 11.11 (Brown, Unit Operations, *Wiley, New* York, 1950). (b) Deep cone thickener developed for the National Coal Board (UK). In a unit about 10 ft dia the impellers rotate at about 2 rpm and a flow rate of 70 m³/sec with a solids content of 6 wt %, concentrates to 25-35 wt % (*Svarovsky*, 1981).

11.13(a) performs cake removal at reduced rotating speed, whereas the design of Figure 11.13(d) accomplishes this operation without slowing down. The clarifying centrifuge of Figure 11.13(e) is employed for small contents of solids and is cleaned after shutdown. The units of Figures 11.13(b) and (c) operate continuously, the former with discharge of cake by a continuous helical screw, the latter by a reciprocating pusher mechanism that operates at 30-70 strokes/min and is thus substantially continuous.

Hydrocyclooes generate their own, mild centrifugal forces. Since the acceleration drops off rapidly with diameter, hydrocy-

TABLE 11 .9. Performances of Sedimentation Equipment (a) **Thickeners**⁴

	%	solids	Unit area,
	Feed	Underflow	sq. ft. /ton. day
Alumina, Bayer process:			
Red-mud primary settlers	3-4	10–25	20-30
Red-mud washers	6-8	15-20	10-15
Red-mud final thickener	6-8	20-35	10-15
Trihydrate seed thickener	2-8	30-50	12-30
Cement, West process	16-20	60-70	15-25
Cement kiln dust	9-10	45-55	3-18
Coral	12-18	45-55	15-25
Cyanide slimes	16-33	40-55	5-13
Lime mud:			
Acetylene generator	12-15	30-40	15-33
Lime-soda process	9-11	35-45	15-25
Paper industry	8-10	32-45	14-18
Magnesium hydroxide from brine	8-10	25-50	60-100
Metallurgical (flotation or gravity concentration):			
Copper concentrates	14-50	40-75	2-20
Copper tailings	10-30	45-65	4-10
Lead concentrates	20-25	60-80	7-18
Zinc concentrates	10-20	50-60	3-7
Nickel:			
Leached residue	20	60	8
Sulfide concentrate	3-5	65	2 5
Potash slimes	I - 5	6-25	40-l 25
Uranium:			
Acid leached ore	1 0-30	25-65	2-10
Alkaline leached ore	2 0	60	10
Uranium precipitate	1 - 7	10-25	50-125

(b) Clarifiers

Application	Overflow rate, gal./min.,sq.ft.	Detention time, hr.
Primary sewage treatment (settleable-solids removal)	0.4	2
clarifiers-activated sludge and trickling filters) Water clarification (following 30-	0.55-0.7	1.5-2
min. flocculation)	0.4-0.55	3
rate—upflow units) Industrial wastes	1.5 Must be tested for	2 each application

"See also Table 14.7.

(Perry's Chemical Engineers Handbook, McGraw-Hill, New York, 1963, pp. 19.49, 19.52).

clones are made only a few inches in diameter. For larger capacities, many units are used in parallel. The flow pattern is shown schematically in Figure 11.13(f). The shapes suited to different applications are indicated in Figure 11.13(g). 10 Figure 11.13(h), the centrifugal action in a hydrocyclooe is assisted by a high speed impeller. This assistance, for example, allows handling of 6% paper pulp slurries in comparison with only 1% in unassisted units. Hydrocyclones are perhaps used much more widely for dust separation than for slurries.

11.7. APPLICATIONS AND PERFORMANCE OF EQUIPMENT

Data of commercially available sizes of filtration equipment, their typical applications, and specific performances are available only to a limited extent in the general literature, but more completely in



Figure 11.7. Two types of laboratory filter arrangements. (a) Vacuum test filter arrangement; standard sizes are 0.1, 0.05, or 0.025 sqft (*Dahlstrom and Silverblatt*, 1977). (b) Laboratory pressure filter with a vertical filtering surface and a mechanical agitator; mild air agitation may be preferred (*Bosley*, 1977).

manufacturers' literature. Representative data are collected in this section and summarized in tabular form. One of the reasons why more performance data have not been published is the difficulty of describing each system concisely in adequate detail. Nevertheless, the limited listings here should afford some perspective of the nature and magnitude of some actual and possibly potential applications.

Performance often is improved by appropriate pretreatment of the slurry with flocculants or other means. An operating practice that is finding increasing acceptance is the delaying of cake deposition by some mechanical means such as scraping, brushing, severe agitation, or vibration. In these ways most of the filtrate is

11 7. APPLICATIONS AND PERFORMANCE OF EQUIPMENT 321

expelled before the bulk of the cake is deposited. Moreover, when the cake is finally deposited from a thickened slurry, it does so with an open structure that allows rapid filtration. A similar factor is operative in belt or top feed drum filters in which the coarse particles drop out first and thus form the desirable open structure. A review of such methods of enhancement of filtration rates is by Svarovsky (1981).

The relative suitability of the common kinds of solid-liquid separation equipment is summarized in Table 11.3. Filtration is the most frequently used operation, but sedimentation as a method of pretreatment and centrifugation for difficulty filterable materials has many applications. Table 11.15 gives more detail about the kinds of filters appropriate to particular services.

Representative commercial sizes of some types of pressure filters for operation in batch modes are reported in Table 11.11. Some of these data are quite old, and not all of the equipment is currently popular; thus manufacturers should be consulted for the latest information. Commercially available size ranges of continuous belt, rotary drum, rotary disk, and horizontal rotary filters are listed in Table 11.12. For the most part these devices operate with vacua of 500 Torr or less.

Sedimentation equipment is employed on a large scale for mineral and ore processing. These and other applications are listed in Table 11.9(a). The clarification operations of Table 11.9(b) are of water cleaning and sewage treatment. The sludges that are formed often are concentrated further by filtration. Such applications are listed in Table 11.16 along with other common applications of plate-and-frame filter presses. Sludge filter cakes are compressible and have high resistivity so that the elevated pressures at which presses can be operated are necessary for them. Among the kinds of data given here are modes of conditioning the slurries, slurry concentrations, cake characteristics, and cycle times.

Clarification of a great variety of industrial liquids is accomplished on smaller scales than in tank clarifiers by application of cartridge filters; some of these applications are listed in Table 11.10.

Cycle times, air rates, and minimum cake thicknesses in operation of rotary drum filters are stated in Table 11.13. A few special applications of horizontal belt filters are given in Table 11.14, but in recent times this kind of equipment is taking over many of the traditional functions of rotary drum filters. Belt filters are favored particularly for freely filtering slurries with wide range of particle sizes.

The applications listed in Table 11.17 and 11.18 are a few of those of rotary drum, rotary disk, and tipping or tilting pan filters. The last type employs a number of vacuum pans on a rotating circular track; after the cake is formed, the pans are blown back with air and then tipped to discharge the cake. The data of these tables include particle size range, moisture content of the cake, filtering rate, solids handling rate, vacuum pump load and degree of vacuum. Clearly a wide range of some of these variables occurs in practice.

Characteristics of centrifugal filters and sedimentation centrifuges are in Table 11.19. The filtering types are made to handle from less than 5 tons/hr to more than 100 tons/hr of solids, with g-levels ranging from 30 to 3000. For sedimentation types, the g-levels listed range up to 18,000, but high values can be used only with small diameter equipment because of metal strength limitations. Capacity of sedimentation types is measured in terms of liquid rates, the maximum listed here being 100,000 L/hr. An outstanding feature of centrifugal separators is the small sizes of particles that can be handled satisfactorily; the values in the table cover the range 1–400 μ m. Short retention time is a feature of centrifuge operation that may be of interest when unstable materials need to be processed.

322 solid-liquid separation

C _{om} Addr	pany •ss						FILT		ION Mat'i	LEA as Soli	Reco ds: _ Analy	EST Fived	DATA : Date _	SHEET	- _	VACU	UM ANI) PR	ESSUF	RE	Test N Dote 1 By	o	I			
Filte Used	r Type 1 Shim:	No _	Leaf	Size1	(es_		F	: _{†.} 2	1	Liqu Pre	uid: Anoig coot	rsis Form	ning Liq	, , , , , , , , , , , , , , , , ,	1			1	1	1	Location	n Tem	p		•	°F∕°C
		v م	in F	Feed	Vai Pi	c vum ressu	- in. re - P	Hg. \$1.	L			E MI	IN.	Air F	'low 1	Filtrate			₩0	rsh T		Cake Weights				
Run No.	Filter Media and/or Precoat Typ	Feed Temp., ^P F	As Prepared	Back Calculated	Fora	Ma sh	Dry	After Cake Crocks	Forn	Dewater	Wash	Dry	To Crack or Gas Breakthrough Afte Form Wash	(1)	Temp., °F °C	ML,	Clarity	Precoat Penetration	ML.	Temp., °F. °C	Cake / Precoo Thickness, In	Dia. of Shave Area, in.	Tare GMS.	Wet & Tare GMS,	Dry & Tare GMS.	Dish No.
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Figure 11.8. A filtration leaf test data sheet (Dahlstrom and Silverblatt, 1977).



Figure 11.9. Deep bed sand filters for removal of small contents of solids from large quantities of liquids. Accumulations from the top and within the bed are removed by intermittent backwashing. Charcoal may be used instead of sand for clarifying organic liquids. (a) Gravity operation. (b) Pressure operation.

TABLE	11.10.	Application	tion	of	Cartrid	ge	Filters	in	Industry	and
		Typical	Part	ticle	Size	Ra	nges l	Rem	oved	

Industry and Liquid	Typical Filtration Range
Chenrical Industry	
Alum	60 mesh-60 μm
Brine	100-400 mesh
Ethyl Alcohol	5–10 µm
Ferric Chloride	30-250 mesh
Herbicides/Pesticides	100-700 mcsh
Hydrochloric Acid	100 mesh to $5-10 \ \mu m$
Mineral Oil	400 mesh
Nitric Acid	40 mesh to 5-10 μm
Phosphoric Acid	100 mesh to $5-10\mu\text{m}$
Sodium Hydrosidc	$1-3$ to $5-10\mu m$
Sodium Hypochiorite	$1-3$ to $5-10 \mu m$
Sodium Sultate	5-10 μm
Sulfuric Acid	250 mesh to I-3 μ m
Synthetic Oils	25-30 pm
Petroleum Industry	
Atmospheric Reduced Crude	25-75 μm
Completion Fluids	200 mesh to I-3 µm
DEA	250 mcsh to 5-10 µm
Dcasphaltcd Oil	200 mcsh
Decant Oil	60 mesh
Dicsel Fuel	100 mesh
Gas Oil	25-75 µm
Gasoline	I-3 μm
Hydrocarbon Wax	25-30 μm
Isobutane	250 mesh
MEA	200 mesh to 5-10 mm
Naphtha	25-30 pm
Produced Water for Injection	I-3 to IS-20 pm
Residual Oil	25-50 µm
Seawater	5-10 µm
Steam Injection	5-10 µm
Vacuum Gas Oil	25-75 μm
All Industries	
Adhcsivcs	30-150 mcsh
Bouer Feed Water	5-10 μm
Caustic Soda	250 mesh
Chiller Water	200 mcsh
City Water	500 mcsh to I-3 μ m
Clay Slip (ceramic and china)	20-700 mesh
Coal-Based Syntuci	60 mcsh
Condensate	200 mcsh to $5-10 \mu m$
Coolant Water	500 mcsh
Cooling Tower Watch	150-250 mcsh
Deionized Water	100-250 mesh
Ethylene Glycol	100 mcsh to 1-3 pm
l loor Polish	250 mesh
Clyccrinc	5-10 µm
Inks	40-150 mesh
Liquid Detergent	40 mcsn
Machine Oli	150 mesh
Pelletizer water	250 mesh
Phonolic Resin Binder	60 mesh
Photographic Chemicals	25-30 μm
Pump Scal Water	200 mesh to $5-10 \mu\text{m}$
Quench Water	250 mcsh
Resins	30-150 mesh
Scrubber Water	40-100 mesh
wax	20-200 mesh
Wcllwatcr	60 mcsh to 1-3 µm

TABLE 11 .11. Sizes of Commercial Discontinuous Pressure Filters

(a) Approximate Area and Cake Capacity for Various Sizes of Plate and Frame Filters

Size of filter	Effective Filtr per Cham	ation area ber (m)	Cake-Holding Capacity per Chamber per 25 mm of Chamber Thickness I					
plate (mm)	Cast Iron	Wood	Cast Iron	Wood				
250	0.096	0.054	1.2	0.6				
360	0.2	0.123	2.5	1.43				
470	0.35	0.21	4.4	2.5				
630	0.66	0.45	6.3	5.4				
600	1.1	0.765	13.7	9.3				
1000	1.74	1.2	21.62	14.6				
1200	2.5	1.76	31.4	21.36				
1450	3.7	2.46	46.24	30.2				

(b) Sizes of Kelly Filters (in.)

	30 × 49	40 × 108	48 × 120	60 × 108
Number of frames Spacing between frames (in.)	4 50	8 25;:	10 4	12
Filter area (sqft)			450	65;:

(c) Standard Sweetland Filter

No.	ID⁵ (in.)	Length of Shell (in.)	No. Leaves 2 in. Space	No. No. Leaves Leaves 2 in. 4 in. Space Space		Filter Area 4 in. Spacing (sqft)	Total Weight ^c of Filter (lbs)
1	10	$20\frac{1}{2}$	9	5	8	4 ¹ / ₂	550
2	16	$36\frac{1}{2}$	16	9	46	23	2150
5	25	61		15			
7	25	82	3041	20	252 185	123 92	7300 9350
10	31	109	54	27	523	262	16500
12	37	145	7 2	36	1004	502	29600

(d) Vallez Filter (Largest Size Only, 20 ft Long, 7 ft high, 7 ft wide)^d

Spacing) f Leaves (in.)	No. of Leaves	O.D. of Leaf (in.)	Filter Area (sqft)	Cake Capacity (cuft)
3	52	52	1232	6 5
4	39 31	52 52	924	7 2
ő	23	52	734 646	7992

(e) Characteristics of Typical Vertical-Tank Pressure Leaf Filters'

Tank Diam (in.)	Filter Area (sqft)	No of Leaves	Leaf Spacing (in.)	Max. Cake Capacity (cuft)	Tank Volume (gal)	Approx. Overall Height (ft)	Approx. Shipping Weight (Ib)
18	19	5	3	1.8	38	5.5	625
18	24	5	3	2.3	45	6.0	650
18	27	7	2	1.7	38	5.5	650
18	35	7	2	2.2	45	6.0	675
30	80	9	3	7.2	128	6.5	1125
30	95	9	3	8.7	132	7.0	1200
30	110	12	2	6.6	128	6.5	1180
30	125	12	2	8.0	132	7.0	1275
48	320	16	3	30.0	435	6.8	2900
48	370	16	3	35.0	500	9.3	3050
48	440	21	ž	28.0	435	8.8	3125
48	510	21	2	32.0	500	9.3	3325

⁸F. H. Schule, Ltd. ^b Diameter of leaf 1 in. less.

'Filled with water.

^d There are smaller sizes with leaves the outside diameters of which are $44\frac{1}{2}$, 36, 30, and 22 in.; for the 30 in. leaves, four lengths of shell are available.

^eT.Shriver & Co.. Inc.

(Courtesy of Ronningen-Petter Division, Dover Corporation, Portage, MI; Cheremisinoff and Azbel, 1983).



(a)







(c)



Figure 11.10. Pressure filters for primarily discontinuous operation. (a) Classic plate-and-frame filter press and details; the plates are separated for manual removal of the cake (*T. Shriver* Co.). (b) Horizontal plate filter; for cleaning, the head is removed and the plates are lifted out of the vessel (Sparkler Mfg. Co.). (c) Pressure leaf filter; the leaf assembly is removed from the shell and the cake is scraped off without separating the leaves (*Ametek Inc.*). (d) The Kelly filter has longitudinal leaves mounted on a carriage; for cleaning, the assembly is slid out of the shell (Oliver *United Filters*). (e) The Sweetland filter has circular leaves and a split casing; the lower half of the casing is dropped to allow access for removal of the cake (Oliver *United Filters*). (f) The Vallez filter has circular leaves rotating at about 1 rpm to promote cake uniformity when the solids have a wide size range; removal of blown-back or washed back cake is accomplished with a built-in screw conveyor without requiring the shell to be opened (*Gosh-Birmingham Co.*).





Figure 11.10.-(continued)





Figure 11.11. Continuous horizontal vacuum filters especially suited to free settling and draining solids. (a) Principle of the conveyor belt filter; units may operate up to 0.5 m/sec with a cycle time up to 10 min and produce cake thicknesses up to 15 cm. (b) Showing the construction of a grooved rubber belt support for the filter cloth of the belt filter (*Purchas*, 1981). (c) Rotating horizontal vacuum filter; the unit has readily accessible piping and is amenable to thorough washing of free draining solids (*Dorr-Oliver Inc.*).



(a)



Figure **11.12.** Continuous rotary drum filters. (a) Flowsketch of continuous vacuum filtration with a rotary drum filter. The solids are taken away with a screw or belt conveyor (*McCabe and Smith*, Unit Operations of Chemical Engineering, *McGraw-Hill, New York*, 19.56). (b) Cross section of a dip-type rotary drum filter showing the sequence of cake formation, washing, dewatering and cake removal; units also are made with top feed (Oliver United Filters). (c) Cross section of a rotary drum filter with internal filtering surface, suited particularly to free settling slurries (*Oliver United Filters*). (d) Rotary filter with a filtering belt that is discharged and cleaned away from the drum; in the similarly functioning string discharge filters, the filtering cloth remains on the drum but the string assembly follows the path shown here for the belt. (e) Double drum filter, particularly suited to rapidly settling slurries, and may be adapted to cake washing which is not shown in this unit (*System Gerlach, Nordhausen, E.* Germany). (f) Vacuum disk filter, the main kind in use when cake washing is not required (*Dorr–Oliver Inc.*).





(e)



TABLE 11.12. Sizes of Commercial Continuous Vacuum Filters

(a) Horizontal Belt Filters'

Series	Ft ² Range	No. Vac . Pans
2600	1 0-45	1
4600	45-200	1
6900	150-700	1
9600	130-500	2
13,600	600-1200	2

(Eimco).

(b) Rotary Drum, Disk, and Horizontal Filters

					Rota	ry Dr	um C	omp	onent	Filte	rs ^b		
					F	ilter	Surfa	ice A	rea (s	qft)			
Drum Diam (ft)	,						Len	gth	(ft)				
		4	6	6	5 1	01	2	14	16	16	20	22	24
6	7	′6	113	151	18	9 22	6						
8				200	25	0 30	0 3	50	400				
10					31	0 37	2 4	34	496	558	620		
1 2						45	65	32	608	684	760	836	912
				C)isk (Comp	onen	t Filt	ers ^d				
	Disk	dia	m (f	t) "		6		7	6	9		10	11
Number	of	disks											
Min.						2		3	4	5		6	7
Max.						8		9	10	11		12	13
Filtering	area	per	disl	k (sc	(ft)	47	6	7	90	117	14	7	180
					Но	rizonta	al F	ilters					
Dia (f	t)*	6	6	10	13	15	16	17	7 16	19	20	22	24
Area (so	(ft)												
Nom		28	50	78	133	177	201	227	254	283	314	380	452
Eff		25	45	65	120	165	191	217	244	273	304	372	444

⁶Filtrate IO-1600 lb/(hr)(sqft). ⁶Adaptable to knife, wire, string, belt, or roll discharge. ⁶All-plastic construction filters also available in 3 and 4 ft drum dia, providing filter areas of 9 to 100 sqft. All disks are composed of 10 sectors. Disk spacing is 16 in.

The American filter, a similar disk filter, also available in 4ft diameter, with 20 sqft disk. 'Also available in 3, 4, and 11.5 ft diameter.

(Dorr-Oliver Inc.).

Figure 11.12—(continued)



Figure 11.13. Filtering centrifuges. (a) Top suspended batch centrifugal filter; the cake is scraped off the screen intermittently at lowered rotation speeds of 50 rpm or so, cake thicknesses of 2-6 in., cycle time per load 2-3 min (*McCabe* and Smith, Unit Operations of Chemical Engineering, *McGraw-Hill, New* York, 1956). (b) A solid bowl centrifugal filter with continuous helical screw discharge of the cake (Bird Machine Co.). (c) Pusher type of centrifuge in which the cake is discharged with a reciprocating pusher mechanism that operates while the machine is at full speed (Baker-Perkins Co.). (d) Horizontal centrifugal with automatic controls for shutting off the feed, washing the cake and scraping it off, all without slowing down the rotation (*Baker-Perkin* Co.). (e) Supercentrifuge for removing small contents of solids from liquids; dimensions 3-6 in. by 5 ft, speed 1000 rps, acceleration 50,000 g, 50-500 gal/hr, cleaned after shutdown. (f) Pattern of flow in a hydrocyclone. (g) The shape of hydrocyclone adapted to the kind of service. (h) Centrifugal action of a cyclone assisted by a high speed impeller (*Voight Gmbh*).



Figure 11.13—(continued)

TABLE 11.13.	Typical	Applications	of	Industrial	Filters
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	Characteristics		Equipment Type'					Vacuum	Pressure
Material		kg/(m ²)(hr)	Α	В	С	D	Е	(Torr)	(atm)
Flotation concentrates	minerals, 10.3 m	300-1000	-	I	х	1	х	450-600	~
Sedimentation concentrates	>0.3 mm	6000-42.000		-	х	—	х	50 - 150	
Crystals and granules	0.05–0.3 mm	600-2000		-	х	_	х	100-300	
Beverages, juices	worthless solids, use filter aids	150-5000	x	x	ڪني		-		2.5-3.5
Pigments	smeary, sticky, 0.06 mm	120-300		-	x	x	-	500-680	~
		batch mode	х	x			_		2.5-4
Limestone, oxide minerals	fine, high density	200-1000		-	х		_	450-600	
		batch mode	х	х	_	х	-		2.5-4
Cane sugar mud	fibrous, viscous		х	х	х	_	<u> </u>		
Mineral oils	high viscosity,								
	1–20% bleaching clays	100-1000		х			_		4
Liquid fuels	low viscosity,								
	bleaching clays	800-2500		х	-	—	—	~	<4
Varnishes, lacquers	cloudy, viscous,								
	solid adsorbents	15-18	х	-	—		—		1
	worthless solids,								
Fats, oils, waxes	50–70°C	500-800	х	х		-	_		-
Sewage sludge	colloidal, slimy	15-150	_	_	х	—		550-600	
Pulp and paper	fibrous, free filtering fine limestone.	150-500	-	—	X		-	150-500	-
Cement	shale, clay, etc	300-1000	-	-	X	-	-	450-630	-

^a Equipment type: (A) filter press; (B) leaf pressure filters, such as Kelly, Sweetland, etc.; (C) continuous vacuum filter; (D) batch rotary filter; (E) continuous rotary filter.

TABLE 11.14. Design and Operating Factors for Continuous Vacuum Filters

(a) Typical Factors for Cycle Design

	Submergence			% (
Filter type	Apparent	Effective Maximum	under ^b Active Vac or Pressure	Max^c for Washing	Max ford Dewatering Only	Required for Cake Discharge
Drum						
Standard scraper	35	30	80	29	50-60	20
Roll discharge	35	30	80	29	50-60	20
Belt	35	30	75	29	45-50	25
Coil or string	35	30	75	29	45-50	25
Precoat	35, 55, 85	35, 55, 85	93	30	10	5
Horizontal belt	as reg'd	as reg'd	lengthen	as reg'd	as read	0
		•	as reg'd	•		
Horizontal table	as req'd	as req'd	80	as req'd	as reg'd	20
Tilting pan	as req'd	as req'd	75	as req'd	as req'd	25
Disc	35	28	75	none	45-50	25

'Total available for effective subm., cake washing, drying, etc. 'Value for bottom feed filters assume no trunnion stuffing boxes, except for precoat. Consult manufacturers for availability of higher submergences.

Maximum washing on a drum filter starts at horizontal centerline on rising side and extends to 15 past top dead center. ^dDewatering means drainage of liquor from cake formed during submergence.

(b) Typical Air Flow Rates

Type of Filter	Air Flow at 500 Torr Vacuum [m ³ /(h)(m ²)]
Rotary drum	50-80
Precoat drum	100-150
Nutsche	30-60
Horizontal belt or pan	100-150

(c) Minimum Cake Thickness for Effective Discharge

	Minimum Design Thickness				
Filter Type	(in.)	(mm)			
Drum					
Belt	1/8-3/16	3-5			
Roll discharge	1/32	1			
Std scrapter	1/4	6			
Coil	1/8–3/16	3-5			
String discharge	1/4	6			
Precoat	0–1/8 max	O-3 max			
Horizontal belt	1/8-3/16	3-5			
Horizontal table	3/4	2 0			
Filting pan	3/4-1	20-25			
Disc	3/8-1/2	10-13			

[(a, b) Purchas, 1981; (c) Purchas, 1977].

	Filter area,	Slu	urry feed acteristics	. 4	Wash ratio (wt/wt based	Solubles recovery	Final cake moisture
Application	m²	% solids	рН	t/hr	on dry solids)	%	%
Dewatering metallic concentrates Brine	8	4 0	_	2 0	00 0 0	 ,	7
precipitate	25	12		,	0		5.0
Calcine leach	60	45	10	79	1	SU 00 7	14
Uranium leach pulp	120	50	1-2	300	0.4	99.3	1 4
Cyanide leach gold pulp	120	50	10-11	80	0.6	99.6	20

TABLE 11.15. Typical Performance Data for Horizontal Belt Filters

(Delfilt Ltd.; Purchas, 1981).

Type of Material	Nature and level of conditioning	Filtration cycle time (Hr)	Solids feed Wt/Wt (%)	Cake Wt/Wt (%)	Cakø thickness (mm)	Remarks
Fine waste slurry	Polyelectrolytes 0.05-0.3 lb/ton	0.52	15-35	75-82	25-40	More than 80% below
Frothed tailings	Polyelectrolytes 0.05–0.3 lb/ton	1-2.5	15-35	73-80	25-40	240BS mesh
Primary sewage sludge	5–25% lime with 5-I 5% copperas. 5-25 lime and 3–6% ferric chloride	% 3-7 1.5-2	4 - 7 3 - 6	4055 35-50	25-32 25-32	
Digested sewage sludge	or 1-2% ACH(Al ₂ O ₃)	2 - 3				
Heat treated sludge		I - 2	12-15	50-70	32	
Mixed sewage sludge including surplus	Up to 3% aluminium chlorohydrate (Ala Oa basic)	3 - 6	UP to 4	30-45	32	Proportion of surplus
activated	or 30% lime with 30% copperas or 3-8% F ECI 3	2 - 4	up to 4	3040	25	by weight
Paper Mill Humus sludge	1% ACH	8	0.5-1 .5	3045	25	
Paper Mill pool effluent sludge	10% lime, 10% copperas of 1% FECI 3	1 - 3	1-1.5	40-55	25	
Pickling end plating sludge	Up to 10% lime if required	1.5-3	2 - 3	30-45	25-32	
Potable water trestment sludge	In some instances no conditioning is required 0.2-1 1% polyelectrolyte (Fre- Quently it is possible to decent large quantities of clarified water after conditioning end before filtration).	3-8	0.5-3	25-35	19-25	
Brine sludge		1.5-3	10-25	60-70	20-25	
Hydroxide sludge	1 mg/l polyelectrolyte or 10% lime	1.5-3	0.5-I . 5	35-45	25-32	
Lead hydroxide sludge		0.5	45	80	32	

TABLE 11.16. Examples of Filter Press Performance for Dewatering of Wastes in Municipal, Potable Water and Industrial Effluents

(Edwards and Jones Ltd.).

Application	Type of vacuum	Solids content	Solids handling	Mois ure	Air flov	Air flow	
	filter frequency used^b	of feed, wt/wt	rate. kg dry solids h ⁻¹ m ⁻² filter surface'	content of cake, wt/wt	m ³ h ⁻¹ m ⁻² filter surface ^d	Vacuum. mmHg	
Chemicals							
Alumina hydrate	Top feed drum	40	450-750	15	90	125	
Barium nitrate	Top feed drum	80	1250	5	450	250	
			50			500	
Betráwbronase⊌lpenatesoda	Drum	5 0	1750	32	546	300	
Calcium carbonate	Drum	50	125	2 2	36	500	
Calcium carbonate (precipitated)	Drum	30	150	40	36	550	
Calcium sulphate	Tipping pan	35	600	30	90	450	
Caustic lime mud	Drum	30	750	50	108	375	
Sodium hypochlorite	Belt discharge drum	12	150	30	54	500	
Titanium dioxide	Drum	30	125	4 0	36	500	
Zinc Stearate	Drum	5	25	65	54	500	
Minerals							
Frothed coal (coarse)	Top feed drum	30	400	18	7 2	300	
Frothed coal (fine)	Drum or disc	35	200	2 2	54	375	
Frothed coal tailings	Drum	4 0		30	36	550	
Copper concentrates	Drum	50	300	10	36	525	
Lead concentrates	Drum	70	1000	12	54	550	
Zinc concentrates	Drum	70	750	10	54	500	
Flue dust (blast furnace)	Drum	4 0	150	2 0	90	500	
Fluorspar	Drum	50	1000	12		375	

TABLE 11.17. Operating Data of Some Vacuum Filter Applications

Notes:

^a The information given should only be used as a general guide, for slight differences in the nature. size range and concentration of solids, and in the nature and temperature of liquor in which they are suspended, can significantly affect the performance of any filter. ^b It should **not** be assumed that the type of filter stated is the only suitable unit for each application. Other types may be suitable, and the ultimate selection will

normally be a compromise based on consideration of many factors regarding the process and the design features of the filter. ^c The handling rate (in kg $h^{-1}m^{-2}$) generally refers to dry solids except where specifically referred to as filtrate. ^d The air volumes stated are measured at the operating vacuum (i.e. they refer to attenuated air).

(Osborne, 1981).

TABLE 11.18. Typical Performance Data of Rotary Vacuum Filters

	Annewingto	Food colida	Filtration rate (0)	Vacuum F	Pump (9)
Material	particle size	conc. wt %	kg/(m ²)(hr)	m³/(m²)(min)	mm Hg
Disc fitter Flotation coal Copper concentrates Magnetic concentrates Coal refuse Magnesium hydroxide	33-43%-200 mesh 90%-200 mesh 80-95%-325 mesh 35-50%-250 mesh 15 microns av. size	22-26 60-70 55-65 35-40 Io-15	300600 250450 10002000 100125 4060	1.5 0.5 2.5-3 0 0.6 0.6	500 500 600–650 500 500
Drum filter (1) Sugar cane mud CaCO ₃ mud recausticising (2) Corn starch Sewane sludge	Limed for flocculation 15-18 microns, av. size	7-18 by vol. 35-40 32-42	25-75 500–600 110–150	0.2 1.8–2 0.9-I	500 250– 38 0 560
Primary Primary digested (3) Leached uranium ore	Flocculated Flocculated 50–60%–200 mesh Flocculated	5-8 4 –7 50–60	15-30 1 O-20 150–220	05 05 05-	500 509 500
Kraft pulp (4) Kaolin clay Belt drum filter	Long fibre 98–75%–2 micron	1-1 25-35	220-300 30-75	Barometric leg 05	600
(5) Sugar cane mud Sewage sludge Primary Primary digested Corn gluten Corn starch	Seperan flocculated Flocculated Flocculated Self flocculating 15-1 8 microns, av. size	7-18 by vol. 5-8 4-7 1620 oz/U.S . gal 32-42	90250 30-50 1535 15-30 1 80-250	0.2 0.5 0.5 06 0.9-I	500 500 500 500 500 500
 (3) Gold cyanide leached off (3) Spent vegetable carbon Dextrose processing Steel mill dust (3) Sodium hypochlorite 	65%–200 mesh 98%–325 mesh 20–40%–2 microns Fine	50–60 100-1 30 gm/litres 40–50 12	300–600 30–50 1 70–300 150	0.5 1.5 0.6-1 . 2 0.9	500 500 500 500
Top feed drum Iron ore concentrates (6) Sodium Chloride	2–4%–200 mesh 8 mesh top size 5-I 0%–100 mesh	35 25-35	6300–7300 1000-1 500	15 30	150 150
Bone char (6) Ammonium sulphate	1%-70 mesh 515%35 mesh	a20 35-40% by vol.	1200-1700 1000-1700	4 0 45-60	9 0 75
(7) Gypsum from digested phosphate rock	40-50 micron av.	35-40	600900	1.2-1.5	500
(8) Leached cobalt residue (8) Alumina-silica gel catalyst (7) Pentaerythritol	-200 mesh	45–50 12 30–40	250 270 75–100	3 0.9 3.6	360 500 500

Notes: (1) Filtrate very dirty-must be recirculated back to clarifier-cake washed.

(2) String discharge filter.(3) Cake washed.

(4) Roller discharge drum filter.

(5) Filtrate very clean-goes directly to evaporation-cake washed.

(6) lop feed filter drier. (7) Two or three stages of counter-current washing.

(8) Three stages of counter-current washing.

(9) Based on tottal fifitter a percea.

(Data of Envirotech Corp.).

TABLE 11.19. Data of Centrifugal Filters and Sedimentation Centrifuges (Purchas, 1977)

(a) Operating Ranges of Main Types of Centrifugal Filters

Type of Centrifuge	Continuous	Automatically Discharged at Full Speed	Automatically Discharged at Reduced Speed	g-Factor Range (F_c)	Minimum Solid Concentration in Feed [% by Volume (C,)	Possibility of Washing	Minimum Particle Size, mm	Minimum Filtrability Coefficient (k)(m/sec)	Maximum Retention I i m e (Sec)
Oscillating	х			30-120	40	no	0.3	5 x 10 ⁻⁴	6
Tumbler	Х			50-300	40	no	0.2	2 x 10 ⁻⁴	6
Worm Screen	Х			500-3000	20	poor	0.06	1 × 10 ⁻⁵	15
Pusher	х			300-2000	30	good	0.08	5 x 10 ⁻⁵	60
Peeler		Х	Х	300-1 600	5	verv good	0.01	2 × 10 ⁻⁷	as wanted
Pendulum			Х	200-1200	5	very good	0.005	1 × 10 ⁻⁷	as wanted

(Hultsch and Wilkesmann; Purchas, 1977).

TABLE **11.19**—(continued)

(b) Criteria for Selection of Sedimentation Centrifuges

Parameter	Tubular Bowl	Skimmer Pipe	Disc	Scroll
Solids concentration. vol./vol.	<1%	up to about 40%	up to about 20%	any as long as it remains pumpable
Particle size range processable for density difference under 1 g/cc and liquor viscosity 1 CP	¹ / ₂ –50μm	10 µm–6 mm	I-400 µm	5 μ m-6 mm
Settling time of 1 litre under 1 g	Few hours to infinity	¹ / ₂ hr to davs	several hours	¹ / ₂ −1hr
Settling time of 50 cc at 2000 g	5-15 min	Î-5 min	5-10 min	Î-5 min
Approximation maximum throughput for largest machine	5000 litre/hr	15,000 litre/hr	100,000 litre/hr	70,000 litre/hr
Approximate nominal throughput for largest machine	1250 litre/hr	12,000 litre/hr	40,000 litre/hr	30,000 litre/hr
Nature of bottle spun solids	Can be any consistency	Must be fluid to pasty	Must not be too cohesive	Preferably compact and cohesive
Batch or continuous	Batch	Semi	Semi or continuous	Continuous
Floc applicable	Possibly but not usual	Yes	Νο	Yes
$oldsymbol{g}$ levels used	Up to 18,000. 80,000 Laboratory model	Up to 1600	4500-1 2,000	500-4000
Maximum sigma value x 10 ⁷ cm ²	5	4	10	1 4

(F.A. Records).

REFERENCES

- C. Almy and W.K. Lewis, Factors determining the capacity of a filter press, *Ind. Eng. Chem. 4, 528* (1912).
- N.P. Cheremisinoff and D. Azbel, *Liquid Filtration*, Ann Arbor Science, Ann Arbor, MI, 1981.
- 3. R. Bosley, Pressure vessel filters, in Purchas, Ref. 14, 1977, pp. 367-401.
- D.A. Dahlstrom and C.E. Silverblatt, Continuous filters, in Purchas, Ref. 14, 1977, pp. 445-492.
- E. Davies, Filtration equipment for solid-liquid separation, Trans. Inst. Chem. Eng. 43(8), 256-259 (1965).
- J.E. Flood, H.E. Parker, and F.W. Rennie, Solid-liquid separation, *Chem. Eng.* 163-181 (30 June 1966).
- M.P. Freeman and J.A. Fitzpatrick (Eds.), Theory, practice and process principles for physical separations, Proceedings of the Engineering Foundation Conference, Pacific Grove California, Oct.-Nov. 1977, Engineering Foundation or AIChE, 1981.
- C. Gelman, H. Green, and T.H. Meltzer, Microporous membrane filtration, in Azbel and Cheremisinoff, Ref. 3, 1981, pp. 343-376.
- C. Gelman and R.E. Williams, Ultrafiltration, in Cheremisinoff and Azbel, Ref. 3, 1981, pp. 323-342.
- J. Gregory (Ed.), Solid-Liquid Separation, Ellis Horwood, Chichester, England, 1984.
- K.J. Ives, Deep bed filtration, in Svarovsky, Ref. 17, 1981, pp. 284-301.
- D.G. Osborne, Gravity thickening, in Svarovsky, Ref. 17, 1981, pp. 120–161.

- D.G. Osborne, Vacuum filtration, in Svarovsky, Ref. 17, 1981, pp. 321-357.
- D.B. Purchas, (Ed.), Solid-Liquid Separation Equipment Scale-Up, Uplands Press, London, 1977.
- D.B. Purchas, Solid-Liquid Separation Technology, Uplands Press, London, 1981.
- A. Rushton and C. Katsoulas, Practical and theoretical aspects of constant pressure and constant rate filtration, in Gregory, Ref. 10, 1984, pp. 261-272.
- L. Svarovsky (Ed.), Solid-Liquid Separation, Butterworths, London, 1981.
- F.M. Tiller (Ed.), Theory and Practice Of Solid-Liquid Separation, University of Houston, Houston, 1978.
- 19. F.M. Tiller and J.R. Crump, Solid-liquid separation: an overview, *Chem. Eng. Prog.*, **73**(10), 65-75 (1977).
- 20. F.M. Tiller, J.R. Crump, and C. Ville, Filtration theory in its historical perspective; a revised approach with surprises, Second World Filtration Congress, The Filtration Society, London, 1979.
- R.J. Wakeman, A numerical integration of the differential equations describing the formation of and flow in compressible filter cakes. *Trans. Inst. Chem. Eng. 56, 258-265* (1978).
- R.J. Wakeman, Filter cake washing, in Svarovsky, Ref. 17, 1981, pp. 408–451.
- 23. R.H. Warring, Filters and Filtration Handbook, Gulf, Houston, 1981.
- Solids Separation Processes, International Symposium, Dublin, April 1980, EFCE Publication Series No. 9, Institution of Chemical Engineers, Symposium Series No. 59, Rugby, England, 1980.

DISINTEGRATION, AGGLOMERATION, AND SIZE SEPARATION OF PARTICULATE SOLIDS

rom the standpoint of chemical processing, size reduction of so/ids is most often performed to make them more reactive chemically or to permit recovery of valuable constituents. Common examples of comminution are of ores for separation of valuable minerals from gangue, of limestone and shale for the manufacture of cement, of coal for combustion and hydrogenation to liquid fuels, of cane and beets for recovery of sugar, of grains for recovery of oils and flour, of wood for the manufacture of paper, of some flora for recovery of natural drugs, and so on.

Since the process of disintegration ordinarily is not high/y selective with respect to size, the product usually requires separation into size ranges that are most suitable to their subsequent processing. Very small sizes are necessary for some applications, but in other cases intermediate sizes are preferred. Thus the byproduct fines from the crushing of coal are briquetted with pitch binder into 3–4–in. cubes when there is a demand for coal in lump form. Agglomeration in general is practiced when larger sizes are required for ease of

handling, or to reduce dust nuisances, or to densify the product for convenient storage or shipping, or to prepare products in final form as tablets, granules, or prills.

Comminution and size separation are characterized by the variety of equipment devised for them. Examples of the main types can be described here with a few case studies. For real, it is essential to consult manufacturers' catalogs for details of construction, sizes, capacities, space, and power requirements. They are properly the textbooks for these operations, since there are few generalizations in this area for prediction of characteristics of equipment. A list of about 90 U.S. and Canadian manufacturers of size separation equipment is given in the Encyclopedia of Chemical Technology [21, 137 (1983)], together with identification of nine equipment types. The Chemical Engineering Equipment Buyers Guide (McGraw-Hi//, New York) and Chemical Engineering Catalog (Penton/Reinhold, New York) a/so provide listings of manufacturers according to kind of eauipment.

12.1. SCREENING

Separation of mixtures of particulate solids according to size may be accomplished with a series of screens with openings of standard sizes. Table 12.1 compares several such sets of standards. Sizes smaller than the $38 \,\mu m$ in these tables are determined by elutriation, microscopic examination, pressure drop measurements, and other indirect means. The distribution of sizes of a given mixture often is of importance. Some ways of recording such data are illustrated in Figure 16.4 and discussed in Section 16.2.

The distribution of sizes of a product varies with the kind of disintegration equipment. Typical distribution curves in normalized form are presented in Figure 12.1, where the size is given as a percentage of the maximum size normally made in that equipment. The more concave the curves, the greater the proportion of fine material. According to these correlations, for example, the percentages of material greater than 50% of the maximum size are 50% from rolls, 15% from tumbling mills, and only 5% from closed circuit conical ball mills. Generalization of these curves may have led to some loss of accuracy since the RRS plots of the data shown in Figure 12.1(c) deviate much more than normally from linearity.

In order to handle large lumps, separators are made of sturdy parallel bars called grizzlies. Punched plates are used for intermediate sizes and woven screens for the smallest sizes. Screening is best performed dry, unless the feed is the product of wet grinding or is overly dusty and an equipment cover is not feasible. Wetting sometimes is used to prevent particles from sticking together. Types of screens and other classifiers to cover a range of sizes are shown in Figure 12.2. Usually some kind of movement of the stock or equipment is employed to facilitate the separations.

REVOLVING SCREENS OR TROMMELS

One type is shown in Figure 12.2(a). They are perforated cylinders rotating at 15–20 rpm, below the critical velocity. The different-

sized perforations may be in series as shown or they may be on concentric surfaces. They are suitable for wet or dry separation in the range of 60-10 mm. Vertically mounted centrifugal screens run at 60-80 rpm and are suitable for the range of 12-0.4 mm.

Examples of performance are: (1) a screen 3 ft dia by 8 ft long with 5-mesh screen at 2 rpm and an inclination of 2" has a capacity of 600 cuft/hr of sand; (2) a screen 9 ft dia by 8 ft long at 10 rpm and an inclination of 7" can handle 4000 cuft/hr of coke.

Flat screens are vibrated or shaken to force circulation of the bed of particles and to prevent binding of the openings by oversize particles. Usually several sizes are arranged vertically as in Figures 12.2(b) and (c), but sometimes they are placed in line as in the cylindrical screen of Figure 12.2(a). Inclined screens vibrate at 600–7000 strokes/min. They are applicable down to 38 μ m or so, but even down to 200 mesh at greatly reduced capacity. Horizontal screens have a vibration component in the horizintal direction to convey the material along; they operate in the range of 300-3000 strokes/min.

Shaking or reciprocating screens are inclined slightly. Speeds are in the range of 30–1000 strokes/min; the lower speeds are used for coal and nonmetallic minerals down to 12 mm, and higher speeds may size down to 0.25 mm. The bouncing rubber balls of Figure 12.2(c) prevent permanent blinding of the perforations.

Rotary sifters are of either gyratory or reciprocating types. They operate at 500-600 rpm and are used for sizes of 12 mm- 50 μ m, but have low capacity for fine sizes.

CAPACITY OF SCREENS

For coarse screening, the required area per unit of hourly rate may be taken off Figure 12.3. More elaborate calculation procedures that take into account smaller sizes and design features of the equipment appear in the following references:

Mathews, Chem. Eng. 76 (10 July 1972) and presented in Chemical

TABLE 12.1. Comparisor	n Table of United States,	Tyler, Canadian, British,	, French, and German	Standard Sieve Series
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U.S.A. (1)		TYLER (2)	TYLER (2) CANADIAN (3)		BRITISH (4)		FREN	CH (5)	GERMAN (6)
"Standard	Alternate	Mesh Designation	Standard	Alternate	Nominal Aperture	Nominal Meeh No .	Ope. M.M.	No.	Opg.
125 mm	5" 4.24"		125 mm 106 mm	5″ 4.24″			1	Ī	
100 mm	4"		100 mm	4"					
7s mm	3"		75 mm	3"					
63 mm	2%		63 mm	215			-		
50 mm	2"		50 mm	2"					
46 mm 37.6 mm			45 mm 37.6 mm						
31.5 mm	1%*		31.5 mm	1%					
26.5 mm 26.0 mm	1.06*	1.05"	26.5 mm 25.0 mm	1.06" 1 "					26.0 mm
22.4 mm	<i>X</i> .	.883"	22.4 mm	1					20.0 mm
15.0 mm	- 74	.146	19.0 1111	<u>/1</u>					18.0 mm
16.0 mm	530"	.624″	16.0 mm	520"					16.0 mm
12.5 mm	K	.020	12.6 mm						12.6 mm
11.2 mm	16"	.441**	11.2 mm	1/10-			-		10.0 mm
9.5 mm	8.	.371″	9.5 mm	5.					10.0 mini
6.7 mm	.265	3	6.7 mm	.265"					8.0 mm
6.3 mm	<u>ד</u>	0.0/	6.3 mm	א "					6.3 mm
5.0 mm	NO. 373	3%	5.6 mm	NO.			5.000	38	5.0 mm
4.75 mm 4.00 mm	4	4 5	4.75 mm 4.00 mm	4			4.000	37	4.0 mm
3.35 mm	6	6	3.35 mm	Ğ	3.35 mm	5		- 0.0	0.10
2.80 mm	7	7	2 80 mm	7	2.60 mm	j 6	3.150	36	3.16 mm
2.36 mm	.8	8	2.36 mm	8	2.40 mm	1	2.600	35 34	2.5 mm
2.00 mm 1.70 mm	10	10	1.70 mm	12	4. 0 8 mm	10	ĩ.600	33	1.6 mm
1.40 mm	14	12	1.40 mm	14	1.40 mm	12	1 250	32	1.25 mm
1.18 mm	16	14	1.18 mm	16	1.20 mm	14	1.200	01	1.0
1.00 mm 850 µm	18 20	16 20	1.00 mm 850 متب	18 20	mm 1.00 mm 850 مبر	16	1.000	31	1.0 mm
		9.4	710 mm	 0 E	710	0.0	.800	30	800 µm
710 mm	25	24		20		~ ~ ~	.630	29	س ر 630
600 μΩ 500 μΩ	30 35	28 32	500 متس متسر 500	30 35	500 متسر متسر 500	25 30	.500	28	<u>مم</u> ر 500
425 µm	40	35	425 µm	40	می 420	36	400	27	400 um
355 µm	4 5	60	335 µm	46	356 נומע	44	.400	£1 0.0	
300 mm	50	65 60 48	21 µnum	60	anu 300 متلي	62	.315	26	מז 16 (11
250 µm	60			60	250 µm	60	.250	25	250 µm
212 µm	70			70	210 מת ע	72	.200	24	m 200
ממע 180	80	80	mس 180	80	180 µm	85	160	23	160 um
150 µm	100	100		100	150 متثبر	- m	.100		
125 µm 106 µm	120	115	125 µm 106 µm	120	125 מחש 105 מחש	120	.125	22	125 µm
	170	100		170	90 µm	170	.100	21	100 µm 90 µm
συμπα	170	170	<u></u>	170		170	.080	20	80
76 μπ	200	200	75 μπ	200	75 µDA	200			mu 71
63 µm	230	250	63 µm	230	63 µm	240	.063	19	63 µm
53 um	270	270	53 um	270	5 <u>3 µm</u>	300			σομΒ
48	210 00 7	205		0.07	45	250	.050	18	50 µm 45 µm
40 µm	325	325	45 µm	325	45 μΩ	350	.040	17	40 µm
38 µ703	400	400	38 µm.	400			1		

U.S.A. Sieve Series-ASTM Specification E-I 1-70
 Tyler Standard Screen Scale Sieve Series.
 Canadian Standard Sieve Series 8-GP-1d.

(4) British Standards Institution, London BS-410-62.
 (5) French Standard Specifications, AFNOR X-I I-501.
 (6) German Standard Specification D1N 4188.

* These sieves correspond to those recommended by ISO (International Standards Organization) as an International Standard and this designation should be used when reporting sieve analysis intended for international publication.



Figure 12.1. Normalized cumulative size distribution curves of comminuted products. (a) From various kinds of crushing equipment. (b) From rod and ball mills. (c) RRS plots of two curves (Z'aggart, 1951).

Engineers' Handbook, McGraw-Hill, New York, 1984, p. 21.17.

- Kelly and Spottiswood, *Introduction to Mineral Processing*, 1982, p. 193.
- V.K. Karra, Development of a model for predicting the screening performance of a vibrating screen, *CIM Bull.* 72, 167-171 (Apr. 1979).
- The last of these procedures is in the form of equations suitable for use on a computer.

12.2. CLASSIFICATION WITH STREAMS OF AIR OR WATER

Entrainment of particles with streams of air or water is particularly suitable for removal of small particles from mixtures. Complete distribution curves can be development by employing several stages operating at suitable conditions in series.

AIR CLASSIFIERS

Although screens of 150 mesh and finer are made, they are fragile and slow, so that it is often preferable to employ air elutriation to



(c)



Figure 12.2. Equipment for classifying particulate solids by size from more than 0.5 in. to less than 1.50 mesh. (a) Rotating cylinder (trommel) for sizing particles greater than 0.5 in., 2-10 rpm, $10-20^{\circ}$ inclination. (b) Heavy duty vibrating screen, 1200-1800 vib/min (Tyler-Niagara, Combustion Engineering Inc.). (c) Three-product reciprocating flat screen, 500-600 rpm, with bouncing rubber balls to unbind the openings, dry products to 100 mesh (Rotex Inc.). (d) Air classifier for products less than 150 mesh. Feed enters at A, falls on the rotating plate B, fines are picked up by air suction fans C, transferred to zone D where they separate out and fall to the discharge, and air recirculates back to fans C (Sturtevant Mill Co.). (e) Dorr drag rake wet classifier. (f) Hydrocyclone.





remove fine particles. The equipment of Figure 12.2(d) employs a rotating plate that throws the particles into the air space from which the finer particles are removed and subsequently recovered.

WET CLASSIFIERS

These are used to make two product size ranges, oversize and undersize, with some overlap. The break commonly is between 28 and 200 mesh. A considerable variety of equipment of this nature is available, and some 15 kinds are described by Kelly and **Spot**tiswood (1982, pp. 200-201). Two of the most important kinds, the drag rake classifier and the hydrocyclone, will be described here.

The classifier of Figure 12.2(e) employs two set of rakes that alternately raise, lower, and move the settled solids up the incline to the discharge. Movement of the rakes is sufficient to keep the finer particles in suspension and discharge them at the lower end. More construction detail of the Dorr classifier may be found in older books, for example, the 1950 edition of the *Chemical Engineers Handbook* (McGraw-Hill, New York). The stroke rate may be 9/min when making separation at 200 mesh and up to 32/min for 28 mesh rapid settling sands. Widths range from 1 to 20 ft, lengths to 40 ft, capacity of 5-850 tons slurry/hr, loads from 0.5 to 150HP. The solids content of the feed is not critical, and that of the overflow may be 2-20% or more.

Hydrocyclones, also called hydroclones, employ self-generated mild centrifugal forces to separate the particles into groups of predominantly small and predominantly large ones. Because of bypassing, the split of sizes is not sharp. The characteristic diameter of the product is taken as d_{50} , the diameter than which 50 wt % of the material is greater or less. The key elements of a hydrocyclone are identified on Figure 12.2(f). A typical commercial unit made by Krebs Engineers has an inlet area about 7% of the cross-sectional area between the vessel wall and the vortex finder, a vortex finder with diameter 3.5-40% that of the vessel, and an apex diameter not less than 25% that of the vortex finder. For such a unit, the

equation for the cut point is

$$d_{50} = \frac{13.2D^{0.675} \exp(-0.301 + 0.0945V - 0.00356V^2 + 0.0000684V^3)}{(\Delta P)^{0.3}(S-1)^{0.5}}$$
(12.1)

and the slurry flow rate is

$$Q = 0.7(\Delta P)^{0.5} D^2 \tag{12.2}$$

in the units $d_{50} \mu m$, vessel diameter *D* in inches, V = vol % of solids in the feed, AP is the pressure drop in psi, S = specific gravity, and Q is the flow rate in gpm (Mular and Jull, in Mular and Bhappu 1978, p. 397). Performance characteristics of one line of commercial hydrocyclones are shown in Figure 12.3(b). Comparison of the chart and equations is made in Example 12.1.

Hydrocyclones are small and inexpensive separators for handling feeds up to about 600 cuft/min and removing particles in the range of 300-5 μ m from dilute suspensions. Large diameters (up to about 24in.) have greater volumetric capacity but also a greater cutpoint on particle diameter. Series and parallel arrangements may be made for any desired compromise between these quantities. In comparison with drag rake classifiers, hydrocylones are smaller, cost about the same to operate but have lower costs for capital and installation. They are preferred in closed circuit grinding.

12.3. SIZE REDUCTION

Crushing is applied to large lumps of feed stock and grinding to smaller lumps, often the products of crushing, but the size distinction is not overly sharp. The process of size reduction results in a range of product sizes whose proper description is with the complete cumulative size distribution, but for convenience a characteristic diameter corresponding to 80% pass in the cumulative distribution curve is commonly quoted.

Some devices employ impact (hammers) and others employ crushing by nipping (rolls or jaws). Within limits, kinetic energy and dimensions of crushing elements can be selected to give a desired reduction ratio. Because of the deformability of solid materials, however, a theoretical limit does exist to the size of particles that can be crushed. These limits are 1 μ m for quartz and 3–5 μ m for limestone. The products of crushing these sizes, of course, can be very much smaller, so that really there is no practical lower limit to grinding.

In practical operations, only about 1% of the input energy to the mill appears as new surface energy of the product. Nevertheless, empirical relations for power consumption based on the extent of size reduction have been developed. One such relation is

$$W = 10W_i(1/\sqrt{d} - 1/\sqrt{d_i}), \text{ kWh/ton},$$
 (12.3)

where d and d_i are the final and initial diameters (μ m) corresponding to 80% cumulative passing. The work index W_i is related to the crushing strength of the material; typical values appear in Table 12.2. Example 12.2 compares a result from this formula with direct data from a manufacturer's catalog.

Characteristics of the main common types of size reduction equipment are listed in Table 12.3, including size of feed, size of product, capacity, power consumption, and average reduction ratio. Coarse comminuters perform with reduction ratios less than 10, fine ones with ratios of 100 or more. From very large to very fine may require several operations in series, as in the flowsketch of Figure 12.4(b), where three stages of crushing and two of classification are shown.



Figure 12.3. Performances of screens and hydrocyclones. (a) Capacities of screens for various products (*Denver Equipment* Co.). (b) Capacity, separation range and pressure drop of hydrocyclones (*Krebs* Engineers). Example: A 380 mm dia vessel has a separation range of 50-110 μ m, and can handle between 200 and 450 gpm at a pressure drop of 7.5 psi.

Toughness, hardness, and temperature sensitivity are some of the properties that influence choice of equipment and operating conditions. Fibrous materials require cutting rather than crushing action. Temperature-sensitive materials such as plastics and rubber need to be cooled with ambient or refrigerated air. Cryogenic processing that involves immersion of the material in liquid nitrogen is employed even for such prosaic materials as scrap automobiles

and rubber tires; the low temperatures enhance brittleness and result in lowered power consumptions.

The kinds of equipment used for certain materials are identified in Table 12.4. Usually several kinds are more or less equally suited. Then the choice may be arbitrary and based on experience or on marginal considerations. Table 12.5 presents a broader range of materials that are being ground in four of the principal kinds of fine

EXAMPLE 12.1 Sizing a Hydrocyclone

A hydrocyclone assembly is required to handle 10,000 gpm of slurries of a solid with specific gravity 2.9 with a cutoff point of $d_{50} = 100 \ \mu\text{m}$. The allowable pressure is $\Delta P = 5$ psi. Several slurry concentrations V will be examined. Substituting into Eq. (12.1) with z the function of V in parentheses,

$$100 = \frac{13.2D^{0.675}e^z}{5^{0.3}1.9^{0.5}}$$

whence

$$D = (16.92/e^z)^{1.4815}$$

The corresponding capacity of one hydrocyclone is

$$Q = 0.7(5)^{0.5} D^2$$

grinders. Performances of attrition, cutter and jet mills with some materials are given with Table 12.7. Additional operating data arranged by material are referred to in Table 12.10.

Closed circuit grinding employs a means whereby only material smaller than a specified size appears in the product. A less precise mode of operation employs an air stream through the equipment at such a rate that only the appropriately fine material *is* withdrawn and the rest remains until it is crushed to size. Ball mills sometimes are operated in this fashion, and also the ring-roller mill of Figure 12.4(a). For closer size control, all of the crushed material is withdrawn as it is formed and classified externally into product and recycle. The other examples of Figure 12.4 illustrate several such schemes.

Wet grinding with water is practiced when dusting is a problem, or when subsequent processing is to he done wet, as of ores that are later subjected to separation by flotation or sink-float processes. Removal of a slurry from a ball mill is easier than of dry material; there are cases where this advantage is controlling. Because of the lubricating effect of the water, power consumption of wet milling is less per ton, but this advantage may be outweighed by corrosion of the equipment.

12.4. EQUIPMENT FOR SIZE REDUCTION

Some of the many available kinds of size reduction equipment can be described here. Manufacturers' catalogs have the most complete descriptions of the equipment and almost always provide typical or expected performance data. Useful compilations of such information are by Taggart (1945) and the *Chemical Engineers Handbook* (1984, Section 8, as well as older editions).

CRUSHERS

Lumps as large as several feet in diameter are crushed in gyratory or jaw crushers. Figure 12.5(a) shows a type of crusher that is made in widths from about 5 to 70 in. and with gaps from 4 to 60 in. Stroke rates vary from 300 to 100/min. The average feed is subjected to 8-10 strokes before it becomes small enough to escape. The jaw crusher is suited to hard, abrasive, and also sticky feeds; it makes minimum fines but the product may be slabby because of the long, narrow exit. Gyratory crushers are more suited to slabby feeds and make a more rounded product.

The results are tabulated following at several values of *V*:

v	ez	in.	mm	Q	No. Units in Parallel
5	1.0953	57.7	1466	5214	2
10	1.4282	39.0	989	2375	4
20	2.0385	23.0	584	828	12
30	3. 2440	11.6	293	209	48

From Figure 12.3(b), with 5 psi a 660 mm unit will handle 1000 gpm and have a cutoff between 50 and $150 \,\mu$ m. This corresponds to the calculated data with *V* about 19 vol %. For a more detailed study of hydrocyclone sizing, the article of Mular and Jull (in Mular and Bhappu, 1980, pp. 376-403) may be consulted. The pressure drop can be adjusted to compensate for changes in slurry concentration.

ROLL CRUSHERS

Toothed rolls such as those of Table 12.8(b) can handle relatively large lumps, for example, 14 in. maximum with 24 in. rolls according to the table. To smooth rolls, the feed size is limited by the angle of nip which depends on the surface conditions but often is approximately 16" or arccos0.961. Accordingly the relation between the diameters of the roll d_r and feed d_f and the gap d_0 between the rolls is given by

$$d_{f} = (0.961d_{f} - d_{0})/0.039. \tag{12.4}$$

For example, with $d_f = 1$ in. and $d_0 = 0.25$ in., the roll diameter is figured as 18 in. Table 12.8(b) lists 16 in. as the smallest size suitable for this service, which appears to be somewhat marginal in comparison with the calculated result. According to the formula, 1 in. lumps could be nipped by 16 in. rolls with a spacing of 0.34 in. It is not possible to state who is smarter, the formula or the manufacturer.

Figure 12.5(b) shows a smooth roll assembly. Usually only one of the rolls is driven and one is spring mounted to prevent damage by uncrushable material in the feed. Reduction ratios shown in Table 12.8(c) range only between 2:1 and 4:1. The proportion of fines is comparatively small. Sets of rolls in series with decreasing settings are used to achieve overall high reduction ratios. The rolls of a pair can he driven at the same or different speeds, within a range of 50-900rpm. The capacity generally is about 25% of the maximum corresponding to a continuous ribbon of material passing between the rolls. A sample listing of materials that are ground in roll mills is in Table 12.5(a). In the arrangement of Figure 12.4(c), the upper pair of rolls is the primary crusher whereas the lower pair works on recycle of the oversize.

Hammer mills employ rotating elements that beat the material until it is small enough to fall through the screen at the bottom of the casing. Product size is determined by the speed of the hammers and the size of the screen openings. Table 12.9(a) shows the former effect. The units of this table operate at speeds up to 900 rpm and make size reductions of 40: 1 or so. The smaller units of Table 12.9(b) operate at speeds to 16,000rpm and make very fine powders. Because of the heating effect, they often are operated with a stream of ambient or refrigerated air for cooling. Under these conditions even heat softening materials such as natural resins or chicle can be ground satisfactorily. Hammer mills are the
TABLE 12.2. Typical Values of the Work Index W, kWh/ton,of Eq. (12.3)

Material	Work Index Wi	Material	Work Index
All materials tested	12 01	Kvanito	40.07
Andesite	22 13		10.07
Rarita	6 24	Lead-zinc ore	11.40
Basalt	20.41	Limestone	11.61
Bauxite	9.45	Limestone for cement	10 18
Cement clinker	13.49	Manganese ore	12.46
Cement raw material	10.57	Magnesite, dead burned	16.80
Chrome ore	9.60	Mica	134.50
Clav	7.10	Molybdenum	12.97
Clav, calcined	1.43	Nickel ore	11.88
Coal	11.37	Oil shale	18.10
Coke	20.70	Phosphate fertilizer	13.03
Coke, fluid petroleum	38.60	Phosphate rock	10.13
Coke, petroleum	73.80	Potash ore	8.88
Copper ore	13.13	Potash salt	8.23
Coral	10.16	Pumice	11.93
Diorite	19.40	Pyrite ore	8.90
Dolomite	11.31	Pyrrhotite ore	9.57
Emery	58.18	Quartzite	12.18
Feldspar	11.67	Quartz	12.77
Ferro-chrome	8.87	Rutile ore	12.12
Ferro-manganese	7.77	Sandstone	11.53
Ferro-silicon	12.83	Shale	16.40
Flint	26.16	Silica	13.53
Fluorspar	9.76	Silica sand	16.46
Gabbro	18.45	Silicon carbide	26.17
Galena	10.19	Silver ore	17.30
Garnet	12.37	Sinter	8.77
Glass	3.08	Slag	15.76
Gneiss	20.13	Slag, iron blast furnace	12.16
Gold ore	14.83	Slate	13.83
Granite	14.39	Sodium silicate	13.00
Graphite	45.03	Spodumene ore	13.70
Gravel	25.17	Syenite	14.90
Gypsum rock	8.16	Tile	15.53
limenite	13.11	Tin ore	10.81
Iron ore	15.44	litanium ore	11.88
Hematite Hematite Organization	12.68	Ігар госк	21.10
nematite-Specular	15.40	uranium ore	17.93
Uulitic	11.33	ZINC OFE	12.42
Limanite	8.45 10.21		
magnetite	10.21		
raconite	14.8/		

[F.C. Bond, Bri. Chem. Eng. 6, 378-385, 643-548 (1961)].

EXAMPLE 12.2

Power Requirement for Grinding

Cement clinker is to be reduced from an initial $d_{80} = 1500 \ \mu \text{m}$ to a final d_{80} of 75 μm . From Table 12.2 the work index is $W_i = 13.49$. Substituting into Eq. (12.3)

 $W = 10(13.49)(1/\sqrt{75} - 1/\sqrt{1500}) = 12.1 \text{ kW/(ton/hr)}.$

According to Table 12.7(b), a 3 ft x 24 in. ball mill requires 10 HP for a rate of 0.5 tons/hr, or 14.9 kW/(ton/hr), a rough check of the result from the equation.

principal equipment for cryogenic processing when products of 50-100 mesh are adequate. Scrap automobiles and rubber tires are chilled with liquid nitrogen and are thereby made highly brittle to facilitate grinding. Nitrogen consumption runs about 0.25 kg/kg steel and up to 0.65 kg/kg rubber [Biddulph, *Chem. Eng.*, (11 Feb. 1980)].

This equipment is particularly suited to crushing of soft, friable materials to cube-shaped products with small proportions of fines. For fibrous materials, the screen is provided with cutting edges. Some data are in Table 12.7(c). A list of materials that are handled in hammer mills is in Table 12.5(a), and other products are referred to in Table 12.10.

Tumbling mills consist of vessels rotating about the horizontal and charged with a mass of relatively small elements that tumble and crush the process material as they fall. Their function may be to mix as well as grind, in batch or continuous operation, in open or closed circuit. Figure 12.4(d) shows a closed circuit arrangement with a ball mill. The crushing elements most commonly are steel balls of several sizes, or ceramic pebbles, or rods the length of the shell, or a range of sizes of the process material that is thus made to grind itself. In processing of minerals, tumbling mills often operate wet with slurries of about 80% solids, just thin enough to permit flow in and out of the equipment.

The mode in which the material grinds itself is called *auto*genous grinding. Such operation can achieve size reduction from 25 cm to 0.1 mm in one step. Autogeneous mills operate at 80–85% of the critical speed, which is the speed at which the grinding media are thrown to the wall and cling to it. They are desirable for mineral treatment since they release the mineral content without overgrinding which could complicate a subsequent flotation process, for instance. Materials for which the process is used are friable and grainy, such as silica rock, asbestos, basic slag, bauxite, cement clinker, dolomite, ferrosilicon, limestone, specular hematite, and taconite. In comparison with ball milling, steel consumption is largely eliminated **but** energy costs are greater by between 25 and 100% because of lower impacting forces with low density materials.

Rod mills [Fig. 12.5(f)] are capable of taking feed as large as 50 mm and reducing it to 300 mesh, but ordinarily the cutoff point is larger. The performance data of Table 12.6(e) shows a product range from 8 to 65 mesh. Rods in use range from 25–150 mm dia; smaller ones tend to bend and break. The ratio of rod length to vessel diameter is kept in the range of 1.4-1.6. Ratios below 1.25 tend to result in tangling. Maximum usable rod length is about 6 m; above this they tend to bend. About 45% of the bulk volume of the mill is occupied by rods. Rotation is at 50–65% of critical speed. Rod consumption normally is in the range of 0.1-1.0 kg steel/ton of ore for wet grinding, and about 10–20% less for dry grinding. Because the coarse feed tends to spread the rods at the feed end, grinding takes place preferentially on the large particles and results in a product of relatively narrow size range. Accordingly, rod mills are nearly always run in open circuit.

Ball mills serve as a final stage of comminution. Balls have a greater ratio of surface area to weight than rods so they are better suited to fine grinding. The length to diameter ratio ranges from less than 1 to about 1.5. Rotation speed is greater than that of rod mills, being 70–80% of critical. Mills that are subjected to vibration can operate above the critical speed. The bulk volume of balls is about 50% of the mill volume.

The Denver ball mills for which operating data are shown in Table 12.6(a) normally are charged with equal weights of 2-, 3-, and 4-in. balls; or for finer grinding, with equal weights of 1.5-, 2-, and 3-in. balls. Figure 12.5(d) is of the widely used conical shape of mill in which a range of sizes of balls group themselves axially during operation. The balls range from 5 in. down, the large ones for crushing the large lumps and the small ones acting on the small

Equipment	Size of Feed (mm)	Size of Product (mm)	Reductio Ratio	Capacity (tons/hr)	Power Consumption (kW)
Gyratory crushers	200-2000	25-250	8	100-500	100-700
Jaw crushers	100-1000	25-100	8	10-1000	5-200
Cone crushers	50-300	5-50	8	10-1000	20-250
impact breakers	50-300	I-10	40	10-1000	100-2000
Rod mills	5-20	0.5-2	10	20-500	100-4000
Ballmills	I-10	0.01-0.1	100	10-300	50-5000
Hammer mills	5-30	0.01-0.1	400	0.1-5	I-100
let mills	I-10	0.003-0.05	300	0.1-2	2-100

TABLE 12.3. Operating Ranges for Commonly Used Size Reduction Equipment

lumps. The performance data of Table 12.6(b) are for wet grinding; dry grinding capacities are 10-20% less. Segregation of balls by size also is achieved in cylindrical shapes with spiral twists in the liner profile (Trelleborg AB).

Tube mill is the term applied to a mill of uniform diameter with greater ratio of length to diameter, in the range of 4-5. Because of the greater length, and correspondingly greater residence time, a finer product can be obtained. Figure 12.5(e) shows a tube mill with three compartments.

Pebble mills are single compartment tube mills with ceramic balls as the grinding medium. They are used when contamination with iron must be avoided, as for grinding and mixing of light colored pigments, food products, and pharmaceuticals. Since the grinding rate is roughly proportional to the weight of the balls, the grinding rate with pebbles is only about 1/3 that with steel balls of the same volume. This is clear from the data of Table 12.6(b). The batch operating data of Table 12.6(c) are for grinding sand of

TABLE	12.4.	Size	Reduction	Equipment	Common	y Used	in
		the C	Chemical Pr	ocess Indu	stries		

Material	Equipment
Asbestos and mica	roll crushers, hammer, and jet mills
Cement	gyratory, jaw and roll crushers, roller, and ball mills
Clays	pan crushers, ring-rollers, and bead mills
Coal	roll crushers, pulverizers, ball, ring-roller, and bowl mills
Coke	rod, ball, and ring-roller mills
Colors and pigments	hammer, jet, and ring-roller mills
Cosmetics and pharmaceuticals	dispersion and colloid mills
Cotton and leather	rotary cutters
Flour and feed meal	roller, attrition, hammer, and pin mills
Graphite	ball, tub, ring-roller, and jet mills
Hard rubber	roller mills
Lime and Shells	hammer and ring-roller mills
Metallic minerals	gyratory and jaw crushers, tumbling mills
Paper and plastics	cutters and slitters
Phosphates	ball and ring-roller mills
Polymers	pulverizers, attrition mills
Pressed cakes	hammer and attrition mills
Refractories	gyratory and jaw crushers, pan and ball mills
Salts	cage and hammer mills
Soaps	hammer, multicage, and screen mills
Starch	hammer and pin mills
Stone and aggregate	gyratory, jaw, and roll crushers
Sulfur	ring-roller mills
Talc and soapstones	roll crushers, ring-roller, pebble, and jet mills

density 100 **lb/cuft** with a charge of 50 **vol** % of pebbles or 33 **vol** % of steel balls. The obvious advantage of batch grinding is that any degree of fineness can be obtained by allowing sufficient time.

Roller mills [Figs. 12.4(a) and 12.6(f)]. Such equipment employs cylindrical or tapered surfaces that roll along flatter surfaces and crush nipped particles. In the ring-roller mill of Figure 12.4(a), spring-loaded rolls are forced against a rotating ring and crush the material that is thrown between them with ploughs. In another design the ring is stationary, and the assembly of rolls is rotated and maintained in contact with the ring by centrifugal force. The unit shown is equipped with built-in air classification: as fines are formed, they are removed with an air stream. For closer control of size distribution, ring-roller mills often are operated with external air classification similar to the process of Figure 12.4(d). The performance data of Table 12.6(c) are for products ranging from 20 to 200 mesh, with appropriate control of air rates. Ring roller mills are used widely for grinding of materials from coal to hard rock and cement clinker; some applications are cited in Tables 12.5(b) and 12.10.

Some special equipment is shown in Figure 12.6.

- a. *Gyratory crusher* was described earlier. The sketch shows the cone-shaped element rotating in a tapered passage. Gyratories are made to handle even larger lumps than jaw crushers. Very large lumps are broken first by explosives or pneumatic hammers before feeding to crushers.
- **b.** *The squirrel-cage disintegrator* illustrated has four concentric cages with horizontal impact bars. The cages rotate in alternately opposite directions, strike the feed and disintegrate or tear it apart. Coal, other friable materials, and fibrous materials are handled in this equipment.
- *c. Disc-type attrition mills* have surfaces that rotate past each other with close clearance and high speed, usually in opposite directions but sometimes in the same direction at different speeds. Clearances are adjustable with spring loading in increments of 0.001 in. Maximum feed sizes are 10–25 mm, diameters 12-48 in., and speeds 1200-7000 rpm. Table 12.5(b) is a list of materials that have been ground in disc mills. Some data are in Table 12.7(b).
- d. *Colloid mills* are **used** to grind and disperse solids in liquids and to prepare emulsions. Adjustable clearances are between 0.001 and 0.050 in., and peripheral speeds to 10,000 **ft/min**. They are used, for example, to make lubricating greases by dispersion of calcium stearate in hydrocarbon oils.
- e. *Buhrstone mills* are an ancient example of an attrition mill. Nowadays they are used mostly for fine grinding of paints, inks, and pharmaceuticals.
- f. Roller mills, also called spindle mills, act by crushing nipped materials between a rotating cylindrical or tapered surface and a

TABLE 12.5. Materials that Have Been Ground in Particular Kinds of Mills

(a) Crusher Rolls ammonium nitrate asbestos barley malt bauxite beet pulp bone casein catalyst beads cereals charcoal cheese chemicals coal cocoa cakes coconut shells coffee cork corn corn cobs corn meal cottonseed cracker meal crackings crimping grains dog food cakes DDT dolomite lime dried biscuits dried apple pulp (b) Disc Attrition Mills alloy powders alum aluminum chips apples, dried asbestos bark borax brake lining scrap brass chips caustic soda cereals chalk charcoal chemical salts chips cloves cocoa coconut shells copper powders copra cork corn cottonseed and hulls drugs dye stuffs egg shells feathers fertilizers fish meal glue (c) Roller Mills alum

barytes

feedstuffs flaxseed floor tile flour fuller's earth glue grains gun powder insulating materials iron oxide lumpy chemicals and flour magnesium oxide malt malted milk meat scraps mustard seed oil bearing seeds pelletized feeds pepper pharamaceuticals plastics reclaim rubber resin salt soy beans spices sponge iron starch uranium concentrates oum arabic hops leather metal powder mica nuts and shells oil cake paris green peanuts and hulls pepper phosphates plaster potash potatoes numice rice and hulls roots rosin rubber sawdust salt suds soy beans spices starch shavings tankage tobacco stems wood pulp hematite insecticide roots

bentonite clay cement clinker chalk cocoa D D T dolomite feldspar graphite gypsum

(d) Hammer Mills aluminum tristearate

animal glue antioxidants asbestos asphalt aspirin bagasse barley bentonite, dried bone char brewer's yeast carbonate calcium calcium phosphate carbon black cellulose acetate cinnamon clav coal cocoa cake cocoa-sugar mixtures coconut shells corn meal cottonseed cake diatomaceous earth dvestuffs etching powder ginger (e) Fluid Jet Mills aluminum aluminum oxide antibiotics asbestos barytes benzene hexachloride carbon Carborundum coal cocoa cryolite DDT dieldrin fatty acids feldspar ferrochrome frite fuller's earth graphite iron oxide lead oxide mica

kaolin lime limestone mica phosphate rock resins soy bean cake sulfur talc titanium dioxide

graphite guar gum gum acacia gypsum irish moss lactose lead, red licorice root lime, hydrated mica milo grain oats, rolled ovster shells pentaerythritol perlite pigments plastic molding compounds potato flour pyrethrum saccharin sage soya flour sugar talc tobacco stems vermiculite

molybdenum disulfide nephelene syenite phenolics PVC pyrethrum resins rotenone calte shellacs silica gel silicon silicon carbide sugar sulfa drugs sulfur talc titanium dioxide toluidine red vanilla beans vitamins waxes yeast

(After Mead, "Encyclopedia of Chemical Process Equipment," Reinhold, N.Y., 1994.)



Figure 12.4. Closed-circuit grinding processes, in which coarse products are captured and recirculated until they are brought down to size. (a) Ring-roller mill (Raymond) with built-in air classification; crushing action is by rotating vertical rolls acting on a revolving bowl ring. (b) Flowsketch of closed-circuit grinding with three stages of grinding and two of classifying (McCabe and Smith, Unit Operations, McGraw-Hill, New York, 1976). (c) A two-pair high roller mill (Schutz–O'Neil Co.); recycle is reground in the lower rolls; Table 12.5(c) lists materials ground by this equipment. (d) A Hardinge conical ball mill in a closed circuit with an air classifier and dust collectors (Hurdinge Co.).









(b)



(c)





Figure 12.5. Jaw, roll, impact, and tumbling equipment for size reduction. (a) Blake-type jaw crusher operates at 200-300 strokes/min (*Allis-Chalmers* Co.). (b) Smooth roll crusher, for which operating data are in Table 12.8(b). (c) Swing hammer mill; operating data in Table 12.7(a). (d) View of a conical ball mill, showing distributions of balls and material and crushing ranges; data in Table 12.6(b) (*Hardinge* Co.). (e) Tube mill with three compartments, length to diameter ratio 3-5. (f) Rod mill in a cylindrical tumbler, L/D = 1.2-1.6; data in Table 12.6(d).

Denver Ball Mill Size Dia. X Lgt. (ft)	-Capacity 2-in to 35 mesh	y (tons per 1-in to 48 mesh	24 hrs) M Jý-in to 65 mesh	edium-hard Jg-in to 100 mesh	l Quartz !4-in to 200 mesh	Rpm Mill	Horse To Run	power Of Motor
3×2	15	11	9	6		33	71/2	10
3×3	2 0	16	14	9		33	10	15
3×4	2 5	21	19	12	7	33	12	15
3 x 6	3 5	31	29	18	9	33	171/2	20
3 x 9	5 0	4 G	44	27	13	33	2 4	25
4 X 3	4 2	34	30	22	12	28	17	20
4 x 5	63	5 5	50	31	16	28	28	30
4 x 10	116	108	103	62	26	28	49	50
5 x 3	ii	63	5 5	40	22	26	3 4	4 0
5 x 6	130	116	110	67	33	26	57	60
5 x 12	250	236	224	136	54	2 G	103	125

(a) Capacities of Some Straight-Sided Ball Mills on Quartz to Various Meshes

TABLE 12.6. Performance of Ball, Pebble, and Rod Mills in Continuous and Batch Modes

(Denver Equipment Co.).

(b) Hardinge Conical Ball Mills in Continuous Wet Grinding: Dry Grinding Rates Are 10-20% Less

3' ∑X @24"ົ	Weight ЗДАЙО	of 12 n4109	Weight of Balls MaximuMO(lim)	Rpm8	Motor (maxL (hp)	Capaci 1½-in to 10 mèsh	ty (tons p 12-in to 100 mcsh 12	er 24 hrs) 34-in to 98 % - 325 mesh 4
5' 🗙 22"	10,200	8,000	8,309	30.4	4 0	140	49	19
6' × 36"	17,100	11,709	17,500	27.7	75	282	97	38
8' 🗙 48"	29,000	23 000	43,500	23.8	200	820	274	108
10' imes 66''	50,600	35:000	83,500	21.2	450	1,900	632	249

(Hardinge Co.).

(c) Hardinge Conical Pebble Mills in Continuous Wet Grinding; Dry Grinding Rates \mbox{Are} 10-20% Less

	Weight	of	Weight of Balls	5	Motor	Capac	ty (tons p	er 24 hrs)
Size	Mill	Lining	Maximum (lbs)	Rom•	(max. hp)	10 mesh	100° mesh	-325 mesh
3' 🗙 24''	3,000	1,300	700	40.4	5	15	5.5	2.1
5' X 22''	9,600	4,000	2,300	31.2	15	54	19	7.5
6′.X3G"	16,500	6,500	4,800	28.2	30	117	4 2	17
8' X 48"	19,400	12,300	12,700	24.1	76	32 6	117	4 5
10' X FG"	35,900	16,800	25,500	21.4	150	675	242	95

(Hardinge Co.).

(d) Pebble and Balls Mills for Batch Grinding of Sand of 100 lb/cuft; Pebble Charge 50 vol %, Steel Ball Charge 33 vol %

				Pebble	M ills-	- Bal	Mills-
M N 81	iii 6A	I.D. of Steel Cylinder 24 X 24"	Capac Grindi Porcelain 104	ity. Dry ng (ibs) Buhrstone 9C	Approx. rpm 40	Grinding (lbs)	Rpm 3 6
8 1 6	ζĊ Α	2 4 X 36" 3 0 X 36"	$\begin{array}{c}164\\273\end{array}$	145 245	86	280 440	36 32
5	2	$36 \times 42"$	440	440	32	740	2 9
* ? 2	2 A	4 2 X 48 60 X 60"	1916	1890	29 19	1152 2936	27 17
1 1	A C	72 X 72" 72 X 120"	$\begin{array}{c} 3456 \\ 5900 \end{array}$	$\begin{array}{c} 3410 \\ 5850 \end{array}$	16 16	$\begin{array}{c} 5076 \\ 8460 \end{array}$	14 14

(e) Performance of Marcy Rod Mills

	Red		Mill		Capaci	ty, tom	/24 hr.	
Size, ft.	tons	100 Inp. 20	speed, r.p.m.	No. 8 sieve	No. 20 sieve	No. 35 sieve	No. 48 sieva	No. 65 sicve
2 × 4 3 × 6 4 × 8 5 × 10 6 × 12 7 × 15 8 × 12 9 × 12	0.9 3.6 7.6 14.5 24.1 42.1 43.4 54 7	4 6 18-22 44 48 85-95 135-150 225-250 230-250 310-340	38 30 25 21 1714 15 13.2 12.5	28 105 240 525 855 1600 1675 2240	15 80 180 390 640 1200 1250 1680	12 65 145 315 510 965 1000 1350	10 50 120 260 425 600 830 1115	7 40 90 195 320 600 625 835

(Paul Abbe Co.).

(Mine and Smelter Division, Kennedy Van Saun Co).

TABLE 12.7. Some Other Kinds of Disintegrators

(a) Ring Roller Mills to Make Down to 100 Mesh

Barytes, 8 to 10 tons/hr to 40 mesh Coal, 5 to 6 tons/hr to 40 mesh Coke (96 hour) 3+ to 4 tons/hr to 20	Gannister, 10 to 12 tons/hr to 14 mesh Iron borings, 8 to 10 tons/hr to 20 mesh Limestone. 8 to 12 tons/hr to 20 mesh
mesh	Limestone, 3 to 4 tons/hr to 85%-200 mesh
Fire clay, 8 to 11 tons/hr	Manganese, 2 to 4 tons/hr to 80 mesh
Florida pebble, 7 t. to 85%-60 mesh	Marble, 3 to 4 tons/hr to 95%—100 mesh
Florida pebble, 3 t. to 95%—100 mesh	Oyster shells, 4 to 5 tons/hr to 60 mesh
The No. 2 mill has 50 per cent larger capacitie	s .
The No. 0 capacity is approximately 35 per ce	ent of the figures for No. $1\frac{i}{2}$ in this table.

Size of feed: 1" to $1\frac{1}{2}$ ".

	DIMENSIONS AND	SPEEDS FOR STURTE	VANT RING ROLL	MILLS	
	Ring	Rolls	Ring Speed		
Size	Diam. x Face	Diam. x Face	(rpm)	Horsepower	
NO. 0	24' x 7"	14 "× 7"	125	8to 15	
No. 1 ¹ /2	45 "× 8"	16" ×1012″	64 45	to 50	
No. 2	44" × 14"	18 " × 14"	70	75	

(Sturtevant Mill Co.).

(b) Attrition Mills for Tough Organic Materials

Material	Size-reduction details	Unit*	Capacity Ib./hr.	Hp.
Alkali cellulose	Shredding for xanthation	В	4,860	5
Asbestos	Fluffing and shredding	С	1,500	50
Bagasse	Shredding	В	1,826	5
Bronze chips	fin. to No. 100 sieve size	Α	50	10
Carnauba wax	No. 4 sieve to 65% < No. 60 sieve	D	1,800	20
Cast-iron borings	$\frac{1}{4}$ in. to No. 100 sieve	Α	100	10
Cast-iron turnings	$\frac{1}{4}$ in. to No. 100 sieve	E	500	50
Cocoanut shells	2 X 2 X $\frac{1}{4}$ in. to 5/100 sieve	В	1,560	17
	5/100 sieve to 43% < No. 200 sieve	D	337	20
Cork	2/20 † sieve to 20/1 20 < No. 200 sieve	D	145	1 5
Corn cobs	1 in. to No. 10 sieve	F	1,500	150
Cotton seed oil and solvent	Oil release from 10/200 sieve product	В	2,400	30
Mica	4 x 4 x $\frac{1}{4}$ in. to 3/60 sieve	В	2,800	6
	8/60 to 75% < 60/200 sieve	D	510	7.5
Oil-seed cakes (hydraulic)	$1-\frac{1}{2}$ in. to No. 16 sieve	F	15,000	100
Oil-seed residue (screw press)	1 in. to No. 16 sieve size	F	25,000	100
Oil-seed residue (solvent)	$\frac{1}{4}$ in. to No. 18 sieve	F	35,000	100
Rags	Shredding for paper stock	В	1,440	11
Ramie	Shredding	В	820	10
Sodium sulfate	35/200 sieve to 80/325 sieve	В	11,880	10
Sulfite pulp sheet	Fluffing for acetylation, etc.	С	1,500	50
Wood flour	10/50 sieve to 35% < 100 sieve	D	130	15
Wood rosin	4 in. max. to 45% < 100 sieve	В	7,200	15

* A-8 in. single-runner mill

B-24 in. single-runner mill

C-36 in. single-runner mill D-20 in. double-runner mill E-24 in. double-runner mill

F-36 in. double-runner mill t 2/20, or smaller than No. 2 and larger than No. 20 sieve size.

(Sprout-Waldron Co.).

TABLE 12.7—(continued)

(c) Rotary Cutters for Fibrous Materials

Material	Screen opening	Feed rate, Ib./hr.	Нр.	Air Remarks on product
Amosite asbestos	1 <u>1</u> ″	1000	11	Yes Finer fiber bundles average length 2"
Cellophane bags	$\frac{11}{32}''$	200	10	Yes Finer than $\frac{5}{16}$ "
Cork	<u>3</u> "	525	16	Yes 90% 4/24" sieve
Chemical cotton	60 mesh	120	15	Yes Flock; 35% under No. 100 sieve
Leather scrap	4	600	20	Yes Precutting before shredding
Fiberglass	$\frac{3}{16}$ "	300	18	Yes 1" (approx.) lengths
Waste paper	<u>5</u> ″ 16	338	13	Yes Through No. 4 sieve and finer
Sheet pulp	40 mesh	150	15	Yes Flock; 85%. 40/100 sieve
Tenite scrap	<u>5</u> "	340	12	No Granulated for reuse
Vinylite scrap	$\frac{7}{32}$ "	300	15	Yes 35%. 6/10 sieve; granular
¹ ₃ ″ Geon sheet	<u>5</u> " 16	540	11	No 99%. 4/20 sieve; for mold- ing granules
Cotton rags	3,, 4	500	11	Yes No linting
Buna scrap	10 mesh	264	12	Yes Granular
Neoprene scrap	30 mesh	90	14	Yes 20°F. temperature rise
Soft-wood chips	<u>1</u> ″	960	12	Yes 90%. 10/50 sieve
Hard-wood chips	$\frac{1}{16}''$	290	11	Yes 83%. 20/100 sieve

* 9per cent 4/24 sieve, *i.e.*, 90 per cent is through No. 4 and on No. 24 sieve. (Sprout-Waldron Co.).

TABLE 12.8. Performance of Jaw and Roll Crushers

(a) Capacities and Data on Blake Type Jaw Crushers (Selected $|{\rm tems})^{\rm a}$

Size of Jaw Opening (in.)	(1 - 1	Ca ton t Open 12	pacit 0 20 Side 2	y (toi cuft = Şei -2 ¹ / ₂	ns/h Capa tting	r) acity) (in.) 4		Jaw Motion (in.)	Horse Power Req .	rpm
10x7	8	1 ⁹	18				₿]	Ş	7 ¹ / ₂	300
20x10		15 24	2 0 3 2	24 40	31 49		A∫ B	58	15	275
30x18			4 8	38 60	45 74	61 102	А В }	<u>11</u> 16	4 0	250

'A-straight jaw plates; B-nonchoking jaw plates.

*(Data supplied by Allis-Chalmers Mfg. Co., Milwaukee, WI).

((b)	Double	Toothed-Roll	Crushers	on	Coal
		DOUNIO		01001010		

Roll Size (in.)		Maximum Size	Dell	Capacity (TPH)	Minimum
Dia	Face	Lump (in.)	(rpm)	Reducing to $1\frac{1}{4}$ to 2	(HP)
18	18	4	150	39-67	8
18	2 0	4	150	46-75	8-10
18	24	4	150	52-88	10-12
24	18	14	125	46-74	12-18
24	2 0	14	125	54-82	15-20
24	24	14	125	62-98	15-20

(Stephens-Adamson Co.)

(c) Relation of Capacity, Size of Feed, Roll Setting, and Speed of Rolls for Sturtevant Balanced Crushing Rolls; Screening in **Closed Circuit** (Average Rock, Which Can Be Nipped at Speeds **Named**)

Size of Roll Dia x Face (in.)	Feed Cube (in.)	Roll Setting (in.)	Speed (rpm)	Capacitγ (tons/hr)
16x10	1.25	0.61	200	26.6
	1	0.25	212	11.6
	0.75	0.2	225	9.8
	0.50	0.125	245	6.67
	0.25	0.065	272	3.86
24x15	2	1	115	56.4
	1.5	0.54	130	34.4
	1	0.25	140	17.15
	0.75	0.2	150	14.7
	0.5	0.125	163	10
36x20	3	1.5	59	87
	2.5	1	6 2	61
	2	0.5	70	34.2
	1.5	0.37	78	29.2
	1	0.25	8 5	20.9

(Sturtevant Mill Co.).

TABLE 12.9. Performance of impact Disintegrators

(a) Hammer Mills

	tons/ hr				
	Limestone, $\frac{1}{8}$ in. Slots	Limestone, $\frac{1}{4}$ in. Slots	Burnt lime, $\frac{1}{4}$ in. Slots		
0 Swing-sledge	2-4	4-7	7-9		
Swing-sledge	6-10	12-15	18-20		
2 Swing-sledge	12-15	20-30	60-70		
00 Hinged-hammer pulverizer	-2	2-4	4-6		

Approximate	Screen	Analysis	of	Product,	Reducing	3	in.	Limestone
-------------	--------	----------	----	----------	----------	---	-----	-----------

•			Passing thro	ugh Mesh Stat	ed
Grate Spacing (in.)	. <mark>1</mark> ₄ in.	10 mesh	50 mesh	100 mesh
<u>1</u> 4		99.8%	85%	50%	40%
1 8			99	70	6 0
1		hat the grat		1	

 $\frac{1}{8}$ in. slots means that the grating space was $\frac{1}{8}$ in

Dimensions and Speeds							
			Ins	ide	Food	Bullow	Annex
	Length	Width	Diameter	Width	Opening	Speed (rpm)	HP
0 Swing-sledge	4ft 3in.	4ft lin.	24 in.	10 in	13x 11 in.	1 zoo-1 500	12
Swing-sledge	5ft 1in.	5ft 8in.	30 in.	2 0 in.	17 🗙 20 in.	1000-1300	4 0
2 Swing-sledge	6ft	7ft	3 6 in.	30 in.	20 x 30 in.	1000-1200	75
00 Hinged-hammer pulverizer	2ft 5in.	3 ft	1 6 in.	11 in.	12 x 12 in.	1 ZOO-3600	5-20
O-24 in. Hinged hammer pulverizer	3ft 7in.	5ft 8in .	24 in.	24 in.	12 ¹ / ₂ x 24in.	1000-1200	15-20

(Data supplied by Sturtevant Mill Co., Boston, MA).

(b) High Speed "Mikro-Pulverizer"

Material	Mesh Fineness	No.1 (5 HP)	No. 2 (15 HP)	No. 3 (40 HP)	No. 4 (75 HP)
Aluminum Hydrate	99.8% through 200 98% through 325 99% through 300 70% through 200 99.99% lhrough 325 94% through 325 94% through 325	600	1,800	4.800	9,000
Ball Clay		600	1.800	4.800	9,000
Calcium Arsenate		1.250	3.750	10.000	18.750
Bituminous Coal		500	1,500	4.000	7.500
Carbon Black		450	1,350	3.600	6.750
Cellulose Acetate (Pulp)		200	600	1.600	3.000
Chrome Yellow	99.9% through 200 Smooth Slurry Good Blend 88% through 100 95% through 325 99.9% through 325	1,250	3.750	10.000	18,750
Dry Color Slurry		800	2.400	6.400	12.000
Face Powder Mixture		600	1.800	4.800	9.000
Gypsum, Raw		1,650	5,000	13.200	24.750
Iron Blue		750	2.250	6.000	11.250
Kaolin		750	2,250	6.000	11.250
Malted Milk	99% through 20 90% through 16 96% Ihrough 20 94% through 100 99% through 100	625	1,875	5.000	9.400
Molding Compound		750	2,250	6.000	11,250
Soap Powder		1,500	4,500	12.000	22.500
Soybean Flake		300	900	2.400	4.500
Sugar		600	1,800	4.800	9.000
Tile Clay Body	100% through 16	1.650	5.000	13.200	24,750
Titanium Dioxide	99.8% through 325	600	1.800	4.800	9.000
White Lead	99.99% through 325	1,000	3.000	8.000	15.000
Zinc Oxide	99.9% lhrough 325	600	1.800	4.600	9.000

Top Rotor Speeds— Approximate Idle Loads						
Unit	Speed HP					
No. 1	9,600 RPM	11/2				
No. 2	6,900 RPM	4				
No. 3	No. 3 4,600 RPM 12					
No. 4	3.450 RPM	18				

TABLE 12.10. Mill Performance Data for Grinding of Specific Products

Material	Equipment	Handbook Table No.
Anthracite	ball mill CC	46
Barite	wet Hardinge ball mill	35
Cement clinker	three-compartment wet tube mill	42
Fertilizers	hammer mill	41
Fuller's earth	roller	48
Grain	attrition	32
Gypsum rock	ring-roller	4 5
Iron oxide	ring-roller	47
Limestone	ring-roller	34
Limestone	wet Hardinge ball mill	35
Metal stearates	hammer mill	50
Oyster shells	hammer mill	38
Phosphates	ball mill	39-40
Quicklime	ball mill CC	44
Rubber	roller mill	51
Seed cake	hammer mill	33
Siliceous refractories	pebble mill	36
Slate	three-compartment wet tube mill	43
Sodium carbonate	roller	4 8
Sulfur	ring-roller	4 9

Note: CC is closed circuit grinding; the ring-roller mill has built-in air classification.

(From Chemical Engineers' Handbook, McGraw-Hill, New York, 1984, pp. 8.48-8.60).

flatter surface. When the rotating surface is cylindrical and the flat surface is horizontal, the equipment is called a dry pan mill. The equipment shown throws the crushed material outwards where it is picked up and removed with an air stream. Table 12.5(c) is a list of materials that are being ground in roller mills. The ring-roller mill of Figure 12.4(a) is in this class.

g. *Fluid jet pulverizers* have opposed high speed gas jets that cause collision and disintegration of the particles. A size classifier and fan return larger sizes to the jet stream. The "Majac" jet mill of Figure 12.6(g) is a related kind of device; it has a horizontal section in which high speed gas jets act on the particles. These mills are used primarily for specialty fine grinding of high-value materials. Performance data of **Micronizers** are in Table 12.9(c); those of the Majac pulverizer are expected to be similar.

12.5. PARTICLE SIZE ENLARGEMENT

For many purposes, lumps of materials of intermediate sizes are the most desirable forms, neither too small nor too large. For instance, beds of overly small granules of catalysts exhibit too great resistance to flow of reacting fluids, and too small particles in suspensions settle out or filter too slowly. Other situations that benefit from size enlargement of particles are listed in Table 12.11.

Because of adhesive forces, particles tend to stick together, particularly small particles that have a large ratio of surface to mass. If a mass is vibrated or shaken lightly, for instance, smaller particles penetrate the interstices between larger ones with increase of contact area and adhesion of the mass. Substances differ naturally in their tendency to agglutinate; as examples, the following groups of materials are listed in the order of increasing tendency to

agglutination:

- 1. Superphosphate, sulfates, and NPK fertilizers.
- 2. Carbamide and diammophosphate.
- 3. Ammonium phosphate, potassium chloride, potassium bicarbonate, and salt.

Adhesion of any mass of particles can be developed by sufficiently high pressure, but lower pressures suffice upon addition of liquid or syrupy binders. Table 12.12 is a list of some commercial agglomerations and the binders that they employ, and Table 12.13 shows how much moisture is needed.

The main types of processes used industrially for particle size enlargement are five in number, defined as follows:

- Compaction is achieved either by compression or extrusion. Compression is done either into a mold to give a final shape or into a sheet or block that is later broken up to proper sizes. Extrudates are formed under pressure in dies of a variety of cross sections; as they leave the die they are broken up or cut to size.
- Agglomeration is accomplished under tumbling or otherwise agitated conditions, with or without binding agents. Size is controlled by adjusting the residence time and by gradual addition of feed and binder, slurry or solution.
- 3. *Globulation* is the formation of droplets of solution, slurry, or melt followed by solidification by prilling, spray drying, or fluidized bed operation. Control of particle size is best achieved in fluidized beds.
- 4. *Heat bonding* is of two types: *nodulization* in which material is tumbled while heated to give hard rounded granules and *sintering* in which the product is an integrated mass that is subsequently broken to size.
- 5. *Flocculation*, coagulation and growth of particles in dilute slurries, to assist in subsequent sedimentation and filtration.

A particular industry may employ more than one of these techniques, for instance the manufacture of solid catalysts. Spherical catalysts are made in rotating pan granulators (Fig. 12.7). If the **rheological** properties are suitable, the material can be extruded (Fig. 12.8), then cut into short cylinders, and subsequently tumbled (Fig. 12.9) into rounded shapes. Smaller spherical beads, for instance, of catalysts for moving bed processes, are made by precipitation or coagulation in an immiscible fluid. Pellets or rings are made on tabletting machines (Fig. 12.10). Although the process is more expensive than extrusion, the product is more nearly uniform. Both extrusion and tabletting result in diffusion resistant skins that, however, usually are eliminated on drying or calcination of the catalyst. Ammonia synthesis catalyst is made by sintering (Fig. 12.11) or fusion of **the** several ingredients, then crushed and used as irregular lumps of size ranges 1.5-3, 6-10, and 12-21 mm.

In the following, the main equipment for particle size enlargement will be illustrated and discussed.

TUMBLERS

The particles of a granular mass will cohere when they are tumbled and sprayed lightly with a liquid binder which often is water or a concentrated solution of the material being agglomerated. The growth may be due to agglomeration of small particles or to layering of material evaporated from the sprayed solution. Rotary kilns of the kind used for drying or chemical reaction (cement or lime burning, for instance) are adapted to size enlarging service. Usually the tumbling action is less intense, only enough to expose the material to sprays. The sprays are fine and are applied to the surface of the bed of particles. The tumbling action then distributes the liquid uniformly through the mass.

The disk granulator of Figure 12.7 is a shallow pan, inclined 45–70° to the horizontal and rotating at speeds of 10–30 rpm. The ratio of pan diameter to collar height is 3-5. The variety of materials to which this equipment is applied is indicated by the listing of part (e) with this figure and in Table 12.14. As the rotation proceeds, fresh solids and spray are charged continually. The finer particles settle to the bottom, the largest remain at the top and then overflow the collar and constitute the product. Because of the size stratification, the product of disk granulation is more uniform in size than of drum granulators which discharge a mixed product. Some

performance data in addition to those shown with Figure 12.7 are:

Material	Diameter (mm)	kg/(min)(m ²)
Iron ore	10-25	11.4
Cement flour		18
Fertilizer	1.6-3.3	14.3

The data of Figure 12.7(c) for cement kiln feed are $42-44 \text{ kg/(min)(m^2)}$.

Pans also are made with height more nearly equal to the diameter. In one such device the material is continually lifted onto



Figure 12.6. Examples of mostly less common devices for size reduction. (a) Schematic of a gyratory crusher for very large lumps. (b) Squirrel-cage disintegrator with four cages. (c) Disc-type attrition mill, rotating at 1200-7000 rpm, clearances adjustable by increments of 0.001 in. (d) Schematic of colloid mill, clearance adjustable between 0.001 and 0.050 in., peripheral speeds to 10,000 ft/min. (e) Buhrstone attrition mill, used for making flour and grinding paints, printers inks and pharmaceuticals. (f) Roller or spindle mill; the crushed material is thrown outwards and removed with an air stream. (g) Majac fluid energy mill making a -200 mesh product; opposed air jets cause high speed collisions and disintegration of the material.



Figure 12.6-(continued)

TABLE	12.11.	Such A	s of Size	enlargement	and Example	es of
						-

	Benefit	Examples of Application
1.	Production of useful structu- ral forms and shapes	pressing of intricate shapes in powder metallurgy; Mahufac - ture of spheres by planetary rolling
2.	Preparation of definite quan- tity units	metering, dispensing, and ad- ministering of drugs in phar- maceutical tablets
3.	Reduced dusting losses	briquetting of waste fines
4.	Creation of uniform, non - segregating blends of fine materials	sintering of fines in the steel industry
5.	Better product appearance	manufacture of fuel briquets
6.	Prevention of caking and lump formation	granulation of fertilizers
7.	Improvement of flow properties	granulation of ceramic clay for pressing operations
6.	Greater bulk density to im- prove storage and shipping of particulates	pelleting of carbon black
9.	Reduction of handling ha- zards with irritating and ob- noxious materials	flaking of caustic
10.	Control of solubility	production of instant food products
11.	Control of porosity and surface-to-volume ratio	pelleting of catalyst supports
12.	Increased heat transfer rates	agglomeration of ores and glass batch for furnace feed
13.	Removal of particles from liquids	pellet flocculation of clays in water using polymeric bridging agents
14.	Fractionation of particle mix- tures in liquids	selective oil agglomeration of coal particles from dirt in water
15.	Lower pressure drop in packed beds	reactors with granular catalysts



an internal sizing screen from which the oversize is taken off as product (Sherrington and Oliver, 1981, p. 69).

Rotating Drum Granulators The equipment of Figure 12.9 is largely free of internals to promote mixing, but provides just sufficient turnover to effect redistribution of the spray throughout the mass. With heavy sprays and little tumbling action, excess and nonuniform agglomeration will occur. Granules 4-6mm dia commonly are made by layering from the sprayed solutions. Fertilizer granules made this way are larger, more dense, and harder than those made by prilling. The trend in the industry has been for prilling towers to be replaced by drum granulators and for those in turn to be replaced by fluidized bed granulators in which dusting problems are most controllable.

A pitch of as much as 10" is used to assist material transport over the length of the drum. Length to diameter ratios of 2-3 are used and speeds of 10-20 rpm. Recommended speeds are about 50% of the critical speed for the dry material; then adequate cascading occurs and the range of particle size distribution is narrowed. Figure 12.9(b) shows the results of such tests on small scale granulation of a fertilizer.

Another application of tumblers is to the manufacture of mixed fertilizers, in which solid ammonium nitrate, liquid ammonia, liquid phosphoric acid, and liquid sulfuric acid are charged separately and reacted. The incidental agglomeration is excessive, however, and the process must be followed by appropriate crushing and size classification.

Various designs of powder blenders can be equipped with sprays and used as granulators. Figure 12.12 is of a trough equipped with two sets of paddles that rotate in opposite directions and throw the particles to the center where they are wetted. Since the mixing is predominantly lateral rather than axial, a measure of plug flow exists and results in a narrower distribution of sizes than from an empty drum, and may approach that from inclined disk granulators. The shape is not quite as rounded, but can be improved if desired by tumbling subsequently in a dry drum. Dimensions and performance of commercial units are shown with Figure 12.12.

TABLE 12.12. List of Agglomerated Products and Their Binders

Material	Binder	Agglomeration	Equipment
Activated Charcoal	Lignosulfonate	Turbulator*	
Alumina	Water	Turbulator*/Disc	
Animal Feed	Molasses	Ring Extruder	
Boric Acid	Water	Disc Pelletizer	
Carbon Black & Iron Powder	Alcohol-Carbowax	Turbulator*	
Carbon, Synthetic Graphite	Sodium Silicate	Turbulator*/Disc	
Cement, Raw Mix	Water	Disc Pelletizer	
Cement Kiln Dust	Water	Turbulator*/Disc	
Charcoal	Starch Gel	Briquetter	
Chrome Carbide	Alcohol	Disc Pelletizer	
Clay, Attapulgite	Water	Turbulator [•] /Disc	
Clay, Bentonite	Water	Turbulator [•]	
Coal, Anthracite	Pitch	Briquetter	
Coal. Bituminous	Lignosulfonate	Disc Pelletizer	
Coal Dust	Waler	Turbulator•	
Coke, Petroleum	Pitch	Briquetter	
Continuous Casting Flux	Waler	Turbulator [•] /Disc	
Copper Smelter Dust	Sodium Silicate	Turbulator [•]	
Copper Sulphite Concentrate	Sodium Silicate	Disc Pelletizer	
Detergent Dust	Water	Disc Pelletizer	
Dolomite Kiln Dust	Water	Turbulator*/Disc	
Dye Pigment	Lignosulfonate	Turbulator*/Disc	
Electric Furnace Dust	Water	Turbulator*/Disc	
Fertilizer	Ammonia	Drum	
Flourspar	Sodium Silicate	Disc Pelletiter	
Flourspar	Lime-Molasses	Briquetter	
Flyash (boiler)	Water	Turbulator*/Disc	
Flyash (high carbon)	Lignosulfonate	Briquetter	
Glass Batch	Caustic Soda	Disc Pelletizer	
Glass Batch	Water	Briquetter	
Herbicide	Lignosulfonale-Water	Turbulator*/Disc	
Herbicide	Clay-Carbowax	Briquetter	
Iron Ore	Bentonite-water	Drum	
Lignite	Gilsonile-Water	Turbulator•/Disc	
Limestone	Clay-Waler	Turbulator•	
Manganese Ore	Lime-Molasses	Briquetter	
Manganese Oxide	Sulfuric Acid	Turbulator*/Disc	
Phosphate Rock	Phosphoric Acid	Turbulator*/Disc	
Plastic Powder	Alcohol	Disc Peiletizer	
Potash Fines	Water	Disc Pelletiter	
Sodium Borate	Sulfuric Acid	Turbulator*/Disc	
Sulfur Powder	Clay	Compactor	
Tungsten Carbide	Alcohol	Disc Pelletizer	
Zeolite	Clay-Water	Turbulator*/Disc	

(Koerner and MacDougal, 1983).

Sticky, very fine, and highly aerated materials can be granulated in drums with high speed impellers with pegs or pins instead of paddles. In Figure 12.13, the material enters at one end, is immediately wetted, and emerges as pellets at the other end. Residence times are under a minute. The data with this figure show that the bulk density of carbon black is increased by a factor of 11, although with about 50% binder in the product.

As mentioned, other powder blending devices can be adapted to granulation, but unless most of the equipment is on hand, it is best to adopt a proven design with which some manufacturer has experience. If the stakes are high enough, the cost of a development program with other equipment may be justifiable.

ROLL COMPACTING AND BRIQUETTING

Agglomeration of finely divided materials is accomplished at high rates and low costs by compression between rolls. The form of product may be continuous sheets that subsequently are broken up to desired size or it may be lumps or briquets of finished form and size. A few shapes are shown in Figure 12.14, but a great variety of simple geometrical shapes can be made with readily available rolls. Briquets are a low cost product, rough in shape, and not of highly uniform weight. When smooth appearance and weight uniformity are demanded, tabletting is the process to be used. The great variety of materials that have been compacted with rolls is indicated

Raw material	Approximate size analyses of raw material, less than indicated mesh	Moisture content of balled product (% H ₂ O)
Precipitated calcium carbonate	200	29.5-32.1
Hydrated lime	325	25.7-26.6
Pulverized coal	48	20.8-22.1
Calcined ammonium metavananiate	200	20.9-21.8
Lead-zinc concentrate	20	6.9-7.2
Iron pyrite calcine	100	12.2-I 2.8
Specular hematite concentrate	150	9.4-9.9
Taconite concentrate	150	9.2-10.1
Magnetic concentrate	325	9.8-10.2
Direct shipping open pit ores	10	10.3-10.9
Underground iron ore	0.25 in.	10.4-10.7
Basic oxygen converter fume	1 µm	9.2-9.6
Row cement meal	150	13.0-13.9
Utilities-fly ash	150	24.9-25.8
Fly ash-sewage sludge composite	150	25.7-27.1
Fly ash-clay slurry composite	150	22.4-24.9
Coal-limestone composite	100	21.3-22.8
Coal-iron ore composite	4 8	12.8-13.9
lron ore-limestone composite	100	9.7-10.9
Coal-iron ore-limestone composite	14	13.3-14.8

TABLE 12.13. Moisture Requirements for Successful Granulation in Tumbling Machines

Courtesy McDowell Wellman Company.



Dish size (m)	Motor (kW)	Capacity (kg s⁻¹)	Material	Remarks
0.36	0.18	0.013	Tungsten carbide	16 x 60 mesh micropellets
0.36	0.18	0.0044	Alumina	
0.99	0.55	0.13	Phosphate rock	85% 4 x 30 mesh product
0.99	0.55	0.076	Bituminous coal filter cake	Feed to pan dryer
0.99	0.55	0.076	Beryl ore mix	Feed to sinter belt
0.99	0.55	0.15	Copper precipitate	
1.37	2.2	0.28	Frit enamel mix	Feed to furnace
2.59	11	8.5	Zinc concentrate sinter mix	Micropelletized sinter machine feed
2.59	11	0.85	Chromate	For electric ore furnace
2.59	7.3	0.93	Bituminous coal fines	For coking furnace
3.05	15	1.7-2.3	Raw shale fines	For expanding in rotary kiln
3.05	18	2.8	Bituminous coal filter cake	
3.66	22	3.4	Zinc sulphide ore	For fluid bed roasting of 4 x 30 mesh pellets
4.27	37	11	Nitrogen fertilizer material	Feed: hot melt and recycle
5.49	44	11	Magnetite ore	Feed to travelling grate – indurating section

(c)

Figure 12.7. Rotating dish granulator applications and performance (Sherrington and Oliver, 1981). (a) Edge and face view of a dish granulator, diameters to 25 ft, Froude no. $n^2D/g_c = 0.5-0.8$. (b) Stratification of particle sizes during rotation. (c) Typical applications of dish granulation (Dravo Corp.). (d) Capacity and power (Dravo Corp.). (e) Performance on cement kiln feed.

	70 lb./cu. ft	125 lb.	/cu. ft .	
	Pelleti	zing	Pellet	izing
Disk size, ft.	Approx. capacity, tons/hr.	HOW- power	Approx. capacity, tons/hr.	Horse- power
18	30	40	40	50
15	18	25	25	30
12	10	12	15	16
9	5	6	10	$7\frac{1}{2}$
6	3	3	5	5
3 %	1/2	1	1	1

Diameter (m)	3.6	4				
Depth (cm)	91	91				
Speed (rpm)	17.5	14.0				
Drive (kW)						
Installed	30	37				
Used	26	25-30				
Feed rate (kg s ⁻¹)	7.1	8-5-10-1				
Moisture (%)	12.5-13.5	12.5-13.5				
Granule porosity (%)	26	26				
Granule compressive	strength (kg) $2 \cdot 7 - 6 \cdot 7$	2.7-6.7				
Powder feed position	Bottor	n centre				
Water feed positions						
Main	Jets above powder feed					
Secondary	Fine sprays in t	op section of pan				





(e)



(a)

Number of tooling stations Output (tablets/min) Max. tablet diameter (in.) Fill depth (in.): Standard Optional Max. operating pressure (tons) Pressure release adjustment (tons) Upper punch entrance (in.)		-2,700	33 1,200-3,300 $1\frac{1}{16}$ $0-\frac{11}{16}$ 10 0-10 37 16-16	45 1,600-4,500 a $0-\frac{12}{16}$ 16-18 10 O-10 3-7 16-13
		(b)		
Series Number of stations	37 37	45 45	55	61
Max. opera phass ure (tons) Max. depth of fii (in.)	10 13	6.5 11	6.5 11	6.5
Max. tablet diameter (in.) Output (tablets/min)	1 888-3,552	\$ 2,050-8,200	7 2,500–10,000	7 16 2,775-1 1,100

(c)

Figure 12.8. Operation and specifications of rotary tabletting machines (*Carstensen*, 1984). (a) Action of the punches of a rotary tabletting machine. (b) Specifications of a Sharples Model 328 (*Stokes-Pennwalt* Co.). (c) Specifications of a Manesty Rotapress Mk 11 (*Manesty* Machines *Ltd. and Thomas Engineering Inc.*).



Figure 12.9. Rolling drum granulator sketch and performance. (a) Sketch of a rolling drum granulator (*Sherrington and Oliver*, 1981). (b) Effect of rotational speed on size distribution: (1) at 20% of critical speed; (2) at 50%. (c) Performance data on commercial units (*Capes and Fouha*, 1984).

by the listing of Tables 12.15 and 12.16. A survey of equipment currently marketed worldwide is made by Pietsch (1976); an excerpt is in Table 12.16.

Compacting of specific materials can be facilitated with certain kinds of additives. Binders are additives that confer strength to the agglomerates, and lubricants reduce friction during the operation. Some additives may function both ways. A few of the hundreds of binders that have been tried or proposed are listed in Table 12.12. Lubricants include the liquids water, glycerine, and lubricating oils; and typical solids are waxes, stearic acid, metallic stearates, starch, and talc.

Successful compacting has been accomplished at temperatures as high as 1000° C. Extrudates are 1-10 mm thick. The information of commercially available equipment of Table 12.17 is representative. Rolls range in size from 130 mm dia by 50 mm wide to 910mmdia by 550mm wide. Capacities are lo-6000 kg/h, and energy requirements, 2-16 kWh/ton. Compacting of mixed fertilizers and similar materials is accomplished by pressures of 30-1200 atm, of plastics and resins by 1200–2500 atm, and of metal powders above 5000 atm. Feed supply may be in the vertical or horizontal direction, by gravity or forced feed. Horizontal feeding is less bothered by entrapped air.

Figures 12.10(a) and (b) show product in sheet form which is subsequently broken down to size. Pellets of large size also may be made for subsequent crushing. For instance, pellets 5 cmdia by 1 cm thick are made for the pharmaceutical trade for breaking up to serve as coarse granular feed to tabletting.

TABLETTING

Rotary compression machines convert powders and granules into hard tablets of quite uniform weight, notably of pharmaceuticals, but also of some solid catalyst formulations. The process is illustrated in Figure 12.8(a). A powder is loaded into a die where it is retained by a lower punch; then it is compressed with an upper punch, and the tablet is ejected by raising both punches.

Most tablets are small, the largest shown in Figure 12.8 is 1-3/16 in. dia and the greatest depth of till is 1-3/8 in., but other machines make tablets 4 in. dia and exert forces of 100 tons. The degree of weight uniformity normally aimed at is indicated by the specifications of the U.S. **Pharmacopeia**. This states that, of a sample of 20 tablets, only two may differ from the mean by the percentage stated following and only one may deviate by twice the percentage stated:

Weight of Tablet (mg)	% Deviation
Equal to or less than 13	15
13-130	10
130-324	7.5
More than 324	5

Greater weight uniformity is achieved with coarse powders or granules as feed. Too large a proportion of fines may cause the tablets to come apart upon ejection. Satisfactory feed can be prepared by first making large tablets in another machine and then



Figure 12.10. Equipment for compacting, briquetting, and pelleting. (a) Flowsketch of a process for compacting fine powders, then granulating the mass **(AU&Chalmers** Co.). (b) Integrated equipment for roll compacting and granulating **(Fitzpatrick Co.). (c)** A type of briquetting rolls. (d) A gear pelleter. (e) A double roll extruder.

breaking them into coarse granules, or by batch-fluidized bed granulation. For pharmaceuticals the range of allowable additives to facilitate tabletting is limited. Magnesium stearate is a common lubricant to the extent of 0-2%, and corn starch is a common binder in the range of 0-5%. Disintegrants and fillers also are used. Preparation of such mixes is accomplished in powder blenders and fixed by granulation.

The Stokes machines of Figure 12.8 operate at 35-100 rpm and the large Manesty at 45-180 rpm. Maximum forces for these sizes of

tablets are 10 tons, but up to 100 tons may be needed for tablets 2.5-4 in. dia. The largest machines shown can be driven with about 50 HP.

EXTRUSION PROCESSES

Powders, pastes and melts are pelleted by extrusion through a die followed by cutting. Binders and lubricants may need to be incorporated in the feed, but the process usually is not feasible for



TABLE	12.14.	Industries	that	Employ	Disk	Granulators	and	Some	of	the
		Products	They	Process	5					

Industry	Typical Application					
Steel	Electric Furnace Baghouse Dust, BOF Dust, OH Dust, Coke Fines, Raw Materials, Iron Ore Pellet - izing					
Foundry	Baghouse Dust, Mold Sand Fines					
Ferroalloy	Silicon, Ferrosilicon, Ferromanganese, Ferro - chrome					
Copper	Concentrates, Smelter Dust, Precipitates					
Lead/Zinc	Concentrates, Sinter Mix, Flue Dust, Drosses					
Other Metals	Tungsten, Molybdenum, Antimony, Brass, Tin, Berrylium, Precious Metals. Aluminum, Silicon, Nickel					
Glass	Glass Rawmix, Furnace Dust, Glass Powder					
Ceramics	Alumina, Catalyst, Molecular Sieves, Substrates, Insulator Body, Tilemix , Press Feed, Proppants, Frits, Colors					
Refractories	Bauxite, Alumina, Kiln Dust, Blends					
Cement/Lime	Raw Meal, Kiln Dust					
Chemicals	Soda Ash, Sodium Sulfate, Detergents, Cleaners, Zinc Oxide, Pigments, Dyes, Pharmaceutical Compounds, Industrial Carbons. Carbon Black					
Ag-Chemicals	Fertilizers, Pesticides, Herbicides, Insecticides. Soil Conditioners, Aglime , Dolomite, Trace Min- erals, N-P-K raw Materials					
Foods	Instant Drink Mix, Powdered Process Foods, Sugar, Sweetners, Confectionary Mix					
coal	Coal Fines					
Power	Coal Fines, Fly Ash, FGD Sludge, Boiler Ash, Wood Ash					
Nonmetallic Minerals	Clay, Talc, Kaolin, Fluorspar, Feldspar, Diatoma- ceous Earth, Fullers Earth, Perlite					
Pulp, Paper, Wood	Paper Dust. Wood Fines, Sander Dust, Boiler Ash					
Solid Waste	Incinerator Ash, Refuse Fines, Mixed Refuse, Dried Sludge					

(Koerner and MacDougal, 1983).



Blunger dimensions	
Length (m)	4.5
Height (m)	1.4
Screw diameter (m) Pitch (m)	0·8 1.7
Shaft speed (rpm)	55
Capacity (kg s ⁻¹) Installed power (kW)	23-25 104

(a)

MODEL	MATERIAL BULK Density lb/ft³	APPROXIMATE Capacity (Tons/HR)	SIZE (WIDTH X LENGTH) (FT)	SPEED (RPM)	DRIVE (HP)
А	2 5	8	2 x 8	56	15
	50	15	2 x 8	56	2 0
	75	2 2	2 x 8	56	2 5
	100	30	2 x 8	56	30
	2 5	30	4 x 8	56	30
	50	60	4 x 8	56	50
	75	90	4 x 8	56	75
	100	120	4 x 8	56	100
	25	3 0	4x 12	56	50
	50	60	4x 12	56	100
	75	90	4x 12	56	150
	100	120	4x 12	56	200
	125	180	4x 12	56	300

(b)

Figure **12.12.** Paddle blending granulator and typical performance. (a) Sketch of a double paddle trough granulator (*Sherrington and Oliver, 1984*). (b) Performance in granulation of fertilizers (Feeco International).

abrasive materials. Economically feasible power requirements correspond to the range of **100–200 lb/HP** hr. The main types of machines are illustrated in Figures 12.10(e) and 12.15.

Screw *extruders* usually are built with a single screw as shown, hut may have as many as four screws and the die may have multiple holes of various cross sections. An 8 in. dia screw can have a capacity of 2000 lb/hr of molten plastics. Tubing can be extruded at 150–300 ft/min. To make pellets, the extrudate goes to cutting machines in which parts as small as washers can be made at rates as high as 8000/min. The extrusion of plastics is described at length by Schwartz and Goodman (*Plastics Materials and Processes*, Van Nostrand Reinhold, New York, 1982); such equipment is applicable to other situations.

Ring pellet mills consist of a power driven rotating ring with radial holes, friction driven rolls to force the material through the holes and knives to cut the extrudate to desired lengths. The feed is charged with screw feeders into the spaces between the rolls and the feed distributor flights. Hole diameters range from 1.6 to 32mm. The force of compaction is due to flow friction through the die. Differing flow and compression characteristics are accommodated by varying the thickness of the ring. A production rate of 200 **lb/HP** hr has been quoted for a normal material through 0.25 in.



(a)

Carbon black feed		
Rate, Mg/day	26.3'	
Bulk density, kg/m³	51.3	
Pellets produced		
Wet basis		
Production rate, kg/h	2108.3	
Bulk density, kg/m ³	562.3	
Denslfication ratio	11.0	
Drv basis		
Production rate, kg/h	1096.3	
Bulk density, kg/m ³	394.1	
Binder		
Specific gravity	1.05	
Injection rate, kg/h	1011.5	
Use ratio, weight of binder to		
weight of wet pellets	0.92	
Power Consumption ^b		
Rate, kW	18.5	
Per Mg of wet pellets, kWh	15.0 ^c	
Production quality		
Rotap test (5 min), %	1.4 (avg. of 45 samp	les)
Crushing strength, g	25 (avg. of 73 samp	oles)

^aAverage from S-day test. plus subsequent production. ^bAmmeter readings. ^cCold shell.

Figure 12.13. Pin mixers which operate at high speed for granulation of fine and aerated powders. (a) **Pinmixer** for the granulation of wetted fine powders. (b) Performance of a pinmixer, dimensions 0.67 m dia by 2.54 m, for pelleting a furnace oil carbon black (Capes, 1980).

(b)

holes. The life of a die is measured in hours. Units of 300 HP are made. Some applications are cited in Table 12.17.

The large tonnage application is to the preparation of animal feeds but many smaller scale applications also are being made. A survey of this literature is made by Sherrington and Oliver (1981).

PRILLING

In the process a molten material is disintegrated into droplets which are allowed to fall and to solidify in contact with an air stream. The equipment is similar to that for spray drying, but the mechanism is simpler in that no evaporation occurs and as one consequence the product is less porous and stronger. A sketch of a prilling process is in Figure 12.16. At one time the chief application was to fertilizers, but a list of many other **prillable** materials is given in Table 12.18.

Dimensional and some operating data for prilling of urea and ammonium nitrate also are in Table 12.18. Towers as high as 60 m have been installed. Because of the expense of towers, prilling is not competitive with other granulation processes until capacities of 200-400 tons/day are reached.



(a)



(b)



Figure 12.14. Common shapes and sizes of pellets made by some agglomeration techniques. (a) Sizes and shapes of briquets made on roll-type machines. (b) Catalyst pellets made primarily by extrusion and cutting *(Imperial Chemical Industries)*. (c) Some of the shapes made with tabletting machines.

Materials suitable for prilling are those that melt without decomposition, preferably have a low heat of solidification and a high enough melting point to permit the use of ambient air for cooling. Because of high viscosities, spray wheels are preferred to spray nozzles. The wheels often are equipped with scrapers to prevent clogging. Since several wheels are needed for capacity, they are often arranged in line and the cross section of the tower is made rectangular.

The density of the prills is reduced substantially when much evaporation occurs: with 0.2-0.5% water in the feed, ammonium nitrate prills have a specific gravity of 0.95, but with 3-5% water it falls to 0.75. Prilled granules usually are less dense than those made by layering growth in drum or fluidized bed granulators. The latter processes also can make larger prills economically. To make large prills, a tall tower is needed to ensure solidification before the bottom is reached. The size distribution depends very much on the character of the atomization but can be made moderately uniform. Some commercial data of cumulative % less than size are:

% Less than size	0	5	50	95	100
Dia (mm)	1.2	1.6	2.4	3.5	4.8

Cooling of the prills can be accomplished more economically in either rotary drums of fluidized beds than in additional tower height. Fluidized bed coolers are cheaper and better because more easily dust controllable, and also because they can be incorporated in the lower section of the tower. After cooling, the product is screened, and the fines are returned to the melter and recycled.

FLUIDIZED AND SPOUTED BEDS

In fluidized bed granulation, liquid or solution is sprayed onto or into the bed, and growth occurs by agglomeration as a result of binding of small particles by the liquid or by layering as a result of evaporation of solution on the surfaces of the particles. The granules grown by layering are smoother and harder. Some attrition also occurs and tends to widen the size distribution range of the product. Larger agglomerates are obtained when the ratio of droplet/granule diameters decreases. Increase in the rate of the fluidizing gas and in the temperature of the bed decreases penetration and wetting of the bed and hence leads to smaller granule sizes. A narrower and more concentrated spray wets a smaller proportion of the particles and thus leads to larger size product. The bed is often made conical so that the larger particles are lifted off the bottom and recirculated more thoroughly.

Initial particle size distribution often is in the range of 50-250 μ m. The product of Table 12.19(a) is 0.7-2.4 mm dia.

A wide range of operating conditions is used commercially. Performance data are in Table 12.19. Gas velocities cover a range of 3-20 times the minimum fluidizing velocity or 0.1-2.5 m/sec. Bed expansion ratios are up to 3 or so. As in fluidized bed drying, bed depths are low, usually between 12 and 24 in. Evaporation rates are in the range 0.005-1.0 kg/(sec)(m²).

Batch fluidized bed granulation is practiced for small production rates or when the residence time must be long. Figure 12.17(a) and Table 12.19(a) are of an arrangement to make granules as feed to pharmaceutical tabletting. A feature of this equipment is the elaborate filter for preventing escape of fine particles and assuring their eventual growth. A continuous process for recovery of pellets of sodium sulfate from incineration of paper mill wastes is the subject of Figure 12.17(b) and Table 12.19(b).

TABLE 12.15. Alphabetical List of Some of the Materials that Have Been Successfully Compacted by Roll Presses

Acrylic resins, activated carbon. adipic acid. alfalfa, alga powder, alumina, aluminium, ammonium
chloride, animal feed, anthracite. asbestos
Barium chloride, barium sulfate. battery masses, bauxite. bentonite, bitumen, bone meal, borax, brass
turnings
Cadmium oxide, calcined dolomite, calcium chloride, calcium oxide, carbomethylccllulose (CMC), carbonates, catalysts. cellulose acetate, ceramics, charcoal, clay, coal, cocoa powder, coffee powder, coke, copper, corn starch
Detergents, dextrine, dimethylterephthalate (DMT), dolomite. ductile metals, dusts, dyes
Earthy ores, eggshells, elastomers. emulsifiers, epoxy resins
Feldspar, ferroalloys, ferrosilicium. fertilizers, flue dusts. fluorspar, fly ash, foodstuffs, fruit powders,
riur wastes, rungiciues Gineuro alass making mixturas, alass nowdar, arain wasta, aranhita, aray iron chins and turnings
Herb teas, herbicides, hons, hydrated lime
Lee inorganic salts iron oxide iron nowder insecticides
Kaolin, kieselur, kieserite
Lead, lead oxide, feather wastes. LD-dust , lignite. lime, limestone, lithium carbonate. lithium fluoride.
lithium hydroxide
Magnesia, magnesium carbonates, magnetite. maleic anhydrate, manganese dioxide, metal powders,
molding compounds, molybdenum, monocalciumphosphate (MCP)
Naphthalene, nickel pwders, nickel ores. niobium oxide
Ores, organic chlorides, organic silicates. oil shale, oyster shells
Pancreas powder, penicillin, pharmaceuticals, phosphate ores. plastics. polyvinylchloride (PVC), potash. potassium compounds, protein pigments. pyrites, pyrocatechol
Raisin seeds. reduced ores, refractory materials. rice starch, rock salt
Salts. sawdust, scrap metals, shales, silicatcs, soda ash. sodium chloride, sodium compounds, sodium cyanide, sponge iron, steel turnings. stone wool, sugar, sulfur
Teas, tin, titanium sponge, turnings
Urea, urea formaldehyde
Vanadium, vermiculite, vitamins
Waxes. welding powder, wood dust, wood shavings
Yeast (dry)
Zinc oxide, zirconium sand

(Pietsch, 1976)

The multicompartment equipment of Figure 12.17(c) permits improved control of process conditions and may assure a narrower size distribution because of the approach to plug **flow**.

Some of the fluidized bed dryers of Figure 9.13 could be equipped with sprays and adapted to granulation. The dryer performance data of Tables 9.14 and 9.15 may afford some concept of the sizes and capacities of suitable granulators, particularly when the sprays are somewhat dilute and evaporation is a substantial aspect of the process.

Spouted beds are applicable when granule sizes larger than those that can be fluidized smoothly. Above 1 mm or so, large bubbles begin to form in the bed and contacting deteriorates. Two arrangements of spraying into a spouted bed are shown in Figure 12.17(d). Particles grow primarily by deposition from the evaporated liquid that wets them as they flow up the spout and down the **annulus**. As the performance data of Table 12.19(c) indicate, particles up to 5 mm dia can be made and even quite dilute solutions can be processed. The diameter of the spout can be deduced from the given gas rates and the entraining velocities of the particles being made. Figure 9.13(f) is a more complete sketch of a spouted bed arrangement. Example 9.9 is devoted to sizing a fluidized bed dryer, but many aspects of that design are applicable to a granulation process.

SINTERING AND CRUSHING

This process originally was developed to salvage iron ore fines that could not be charged directly to the blast furnace. Although other applications appear to be feasible, the original application to iron ore fines seems to be still the only one, possibly because the scale of the operation is so great. Figure 12.11 is a sketch of the process. A mixture of ore fines, some recycle, 14-25% of calcite or dolomite as fluxes, and 2.5–5% solid fuel is placed on a conveyor to a depth of 12in. or so. It is conveyed into an ignition furnace, burned and fused together, and then cooled and crushed to size. Fines under 6 mm are recycled. Sinter feed to blast furnaces is about 40-50 mm in the major dimension. The equipment is very large. One with a conveyor 5 m wide and 120m long has a capacity of 27,000 tons/day.

Nodulizing is another process of size enlargement by fusion. This employs a rotary kiln like those used for cement manufacture. The product is uniform, about 0.5 in. dia, and more dense than sinter.

Sintering of powdered metals such as aluminum, beryllium, tungsten, and zinc as well as ceramics under pressure is widely practiced as a shaping process, but that is different from the sintering process described here.

Model	Roll	dia./mm	Max. roll width/mm	Position of nor rolls/feeder	Max. force/ metric tons	Overload system	Approx. capacity/ kg -1	Feeder	type	Press drivel kW	Feeder drive / kW	Roll shapes	Max. feed temp./ °C
L 200/50	200		50 ho	oriz./vert.	~ 10	None	10-100	screw		3/4	0.5	smooth/corrugated/	80
K 26/100 K 27/200	200 300		100 he 200	prizontal/vertica	-20 -40	**	100-200 200-500	••		22	3 7·5		80 80
K 27/300 cs 25	300 230		300 65	55 55	- 8 0 25	hydraulic	500-l 000 loo-300	**		30 7	7.5 3	**	80 120
cs 50 MS 75	406 500		119 230	**	50 75	••	300-l ooo 1 000–10 000	**		22 75	5 7.5	••	150 120
MS 150 MS200	500 710		280 460	I.	150 200	••	3000-15000 up to 50 000			300	11	9 9	1 000
MS 300 MS 350	710 910		550 250	99 18	300 350	••	up to 60 000 up to 40 000	screw(s) screw		400 250	15		1 000 1 000

(b)

B 100 B 150	130	50	vertical/horizontal	10	hydraulic	20	saew		smooth/corrugated/ pocketed	ambient
B 220	200	75	11	20		200	**		**	**
B 300 8400 B 500 D 100 D 150 D 200	300 380 460 610 130 200	75 100 150 200	11 11 11 11 11 11	30 60 125 250 20 40 70	59 88 89 	1 500 3 000 5 000 15(50 200 3 000	97 99 99 99 99 99	as required	97 19 19 11 11 11	99 99 99 99 99
DH 400 DH 500 DH 600	330 520 710 920		horizon&/vertical	140 270 500	99 99 91	20000 50,000	99 99		11	800 800 800
=	520		11	550		20,000	n		*1	000

[(a) Hutt Gmbh; (b) K.R. Komarek, Inc. From Pietsch, 1976].

Material	Reason To Pellet	LB/HP/HR (KG /KW/HR)	Pellet Size (Inches Diameter) (Millimeter Dia.)
Asbestos Shortr	Density, Reduce Dust In at 201b/ft3 (320kg/m3) Out at 651b/ft3 (1041 kg/m3)	45 (27)	3/8″ (9.5mm)
Acrylamide-Dry wet	Handling	80 (49) 170 (103)	114" (6.4mm)
Bagasse	Denslfy, Reduce Dust In at 51b/ft3 (80kg/m3) Out at 301b/ft3 (480kg/m3)	80 (49)	3/8* (9.5mm)
Bauxite	Handling	300 (182)	1/2" (12.8mm)
Brewers Grain (Spent)	Densify, Handling In at 13ib/ft3 (208kg/m3) Out et 36ib/ft3 (577kg/m3)	150 (91)	1/4" (6.4mm)
clay ease Material	Handling, Densify, Calcine	100-300 (61-182)	1/8" to 314" (3.2mm to 19mm)
Cryolite Filter Cake	Handling	loo (61)	3/8" (9.5mm)
Domolite	Handling	200 (122)	114" to 3/16″ (6.4mm to 4.8mm)
Herbicide	Handling, Control, Solubility	150 (91)	12/64" (4.8mm)
Insecticide	Defined Form, Reduce Dust	120 (73)	1/8″ (32mm)
Iron Oxide	Calcining, Reduce Dust	50-100 (30- 61)	lb" to 1/4* (3.2mm to 6.4mm)
Lignite	Eliminate Fines	100 (61)	1/8" to 114' (3.2mm to 6.4mm)
Nylon Film Scrap	Denslfy	60 (36)	1/8" (3.2mm)
Paper scrap	Densify	83 (50)	1/2" (12.7mm)
Phenollc Molding Compound	Reduce Dust, Handling	60 (36)	1/8″ (3.2mm)
Polyethelyene Fllm	Denslfy from 51b/ft3 (80kg/m3) to 20 lb/ft3 (320kg/m3)	30 (18)	1/8" to 3/16" (3.2mm to 4.8mm)
Polystyrene Foam	Denslfy from 41b/ft3 (64kg/m3) to 241b/ft3 (384kg/m3)	164 (100)	1/8" (3.2mm)
Polypropylene Film	Denslfy	40 (24)	lb" (3.2mm)
Rubber Accelerator	Reduce Dust, Handling	192 (117)	12/64″ (4.8mm)
Starch	Handling	75 (46)	12/64" (4.8mm)
sswdust	Bum	60 (36)	1/4" (6.4mm)
Salt	Handling, Reduce Dust	70 (43)	1/8″ (3.2mm)

(Sprout Waldron Co.).

.



Figure 12.15. Two types of extrusion pelleting equipment. (a) Screw-type extruder for molten plastics: The die is turned 90" in the illustration from its normal position for viewing purposes. The extruded material is cooled and chopped subsequently as needed (U.S. Industrial Chemical Co.). (b) Ring extruders: material is charged with screw conveyors to the spaces between the inner rolls and the outer perforated ring, the ring rotates, material is forced through the dies and cut off with knives.



Figure 12.16. A **prilling** tower for ammonium nitrate, product size range 0.4-2.0 mm. The dryer is not needed if the moisture content of the melt is less than about 0.5%.

TABLE 12.18. List of Typical Prillable Materials and Performances of Some Prilling Operations

(a) List of Typical Prillable Materials

Adhesives	Pentachlorophenol
Adipic Acid	Petroleum wax
Alpha naphthol	Phenolic resins-Novalak resin
Ammonium nitrate and additives	Pine rosin
Asphalt	Polvethvkne ruins
Bisphenol-A	Polystyrene resins
Bitumen	Polypropylene-maleic anhydride
Carbon pitch	Potassium nitrate
Caustic soda	Resins
Cetyl alcohol	Sodium glycols
Coal-derived waxes	Sodium nitrate
coal tar pitch	Sodium nitrite
Dichloro-benzidine	Sodium sulphate
Fatty acids	Stearic acid
Fatty alcohols	Stearyl alcohol
Epoxy resins	Substituted aliphatics
Hydrocarbon resins	substituted amides
High-melting inorganic salts	Sulphur
Ink formulations	Urea and additim
Lauric acid	Urea-sulphur mix
Myristic acid	Wax-resin blends
Myristyl alcohol	
Paraffins	

(b) Data for the Prilling of Urea and Ammonium Nitrate

Tower size			
Prill tube height, ft		130	
Rectangular cross section, ft		11 by 21.4	
Cooling air			
rate, lb/h		360.000	
inlet temperature		ambient	
temperature rise, °F		15	
Melt			
Турс	Urea		Ammonium Nitrate
rate, lb/h	35.200 (190 lb H ₂ O)		43.720 (90 lb H ₂ O)
inlet temperature, • F	275		365
Prills			
outlet temperature, °F	120		225
size. mm	approximately 1 to 3		

(HPD Inc.).

TABLE 12.19. Performance of Fluidized Bed and Spouted Bed Granulators

(a) Batch Fluidized Bed Granulator to Make Feed to Pharmaceutical Tablets; the Sketch Is in Figure 12.17(a)

	APPROXI	MATE RANGE
Batch load, dry basis, lb	20 to	400 ^a
Volume of container for static bed, ft ³	2 to	15
Fluidizing air fan, hp	5 to	25
Air (Steam) heating capacity, Btu/h	70.000 to	600,000
Drying sir temperature, °C	40 to	80
Granulating liquid spray*	Two	fluid nozzle
Air volume	to	2 SCFM
Liquid volume	500 to	1500 cm ³ /min
Batch processing time, min	30 to	50
Average granule size	24 to	8 mesh

^e Batch capacity exceeds 1500 lb in the largest modem units. *Typical granulating liquids are gelatin or sodium cuboxymethyl cellulose solutions.

(Capes and Fouda, 1984).

(b) Performance of Fluidized Bed Granulation of Two Waste Products; Sketch Is in Figure 12.17(b) for Paper Mill Waste

TYPE OF SLUDGE	INCINERATOR	BED SIZE TEMPERATURE	CAPACITY	GRANULAR PRODUCT COMPOSITION
Oil refinery waste sludge (85–95% water)	40 ft high; 20 ft ID at base increasing to 28 ft at top	1330°F	31 x 10 ³ lb/hr of sludge	Start-up material was silica sand; replaced by nodules of various ash components such as CaSO ₄ , Na, Ca, Mg silicates, Al ₂ O ₃ after
Paper mill waste liquor^a (40% solids)	20 ft ID at top	1350° F	31 x 10 ³ lb/hr	operation of incinerator. Sulfur added to produce 90–95% Na₂SO4 and some Na₂CO3

(Capes and Fouda, 1984).

(c) Applications of Spouted Bed Granulations

	Feed	Feed solu tion		Product		nperature		
Material	Moisture content (%)	Temperature (°C)	Size (mm)	Moisture (%)	Inlet (°C)	Outlet (°C)	Gas flow rate (m ³ s ⁻¹)	Capacity (kg h⁻¹)
Complex fertilizer Potassium chloride	27 68	15	3-3·5 4-5	2.4	170 200	70 60	13.9 13.9	4000 1000
Sulphurium nitrate Inorganic pigments,	4	135	2·2- 4	0.2	15	5 5	1-11×10 ⁻²	954 0)
Organic dyes acid blue black Ammonium sulphate Sodium chloride	45 60 77	70	3-5 1-3 ~2 ~4.5	6.5	280 226 190 120	1 54 83 70	~1.3 x 10 ⁻³ ~1.8 x IO-'	ı -2.7 ~1.2

(Sherrington and Oliver, 1981).



Figure 12.17. Fluidized bed and spouted **bed** granulators. (a) A batch fluidized bed granulator used in the pharmaceutical industry; performance data in Table 12.19(a). (b) Part of a fluidized bed incineration process for paper mill waste recovering sodium sulfate pellets; performance data in Table 12.19(b). (c) A three-stage fluidized bed granulator for more complete control of process conditions and more nearly uniform size distribution. (d) Two modes of injection of spray to spouted beds, into the body on the left and at the top on the right; performance data in Table 12.19(c).

REFERENCES

Size Reduction and Separation

- W.L. Badger and J.T. Banchero, *Introduction 10 Chemical Engineering*, McGraw-Hill, New York, 1955.
- W.M. Goldberger, Solid-solid systems, in *Chemical Engineers'* Handbook, McGraw-Hill, New York, 1984.
- E.G. Kelly and D.J. Spottiswood, *Introduction to Mineral Processing*, Wiley, New York, 1982.
- 4. W.J. Mead (Ed.), Balling devices, Briquet machines, Grinding, Mills colloid, Mills roller, Screening, in *Encyclopedia of Chemical Process Equipment*, Reinhold, New York, 1964.
- A.L. Mular and R.B. Bhappu (Eds.), Mineral Processing Plant Design, AIMME, New York, 1980.
- 6. E.J. Pryor, Mineral Processing, Elsevier, New York, 1965.
- 7. E.R. Riegel, Chemical Process Machinery, Reinhold, New York, 1953.
- G.C. Sresty, Crushing and grinding equipment, in *Chemical Engineers'* Handbook, McGraw-Hill, New York, 1984, pp. 8.9-8.59.
- 9. A.F. Taggart (Ed.), Handbook of Mineral Dressing, Wiley, New York, 1945.
- 10. A.F. Taggart, Elements of Ore Dressing, Wiley, New York, 1951.
- 11. B.A. Wills, Mineral Processing Technology, Pergamon, New York, 1985.

Size Enlargement

- 12. C.E. Capes, Particle Sire Enlargement, Elsevier, New York, 1980.
- 13. C.E. Capes, Size englargement, in *Chemical Engineers Handbook*, McGraw-Hill, New York, 1984, pp. 8.60–8.72.
- 14. C.E. Capes and A.E. Fouda, Agitation methods, in Ref. 17, pp. 286-294.
- **15.** A.E. Capes and A.E. Fouda, **Prilling** and other spray methods, in Ref. 17, pp. **294–307**.
- 16. J.T. Carstensen, Tabletting and pelletization in the pharmaceutical industry, in Ref. 17, pp. 252-268.
- M.E. Fayed and L. Otten, (Eds.), Handbook of Powder Science and Technology, Van Nostrand Reinhold, New York, 1984.
- R.M. Koerner and J.A. MacDougal (Eds.), Briquetting and Agglomeration, Institute for Briquetting and Agglomeration, Erie, PA, 1983.
- 19. R.A. Limons, Sintering iron ore, in Ref. 17, pp. 307-331.
- 20. W. Pietsch, Roll Pressing, Heyden, London, 1976.
- 21. P.J. Sherrington and R. Oliver, Granulation, Heyden, London, 1981.
- 22. N.E. Stanley-Wood (Ed.), *Enlargement and Compaction of Particulate Solids*, Butterworths, London, 1983.

DISTILLATION AND GAS ABSORPTION

he feasibility of separation of mixtures by distillation, absorption, or stripping depends on the fact that the compositions of vapor and liquid phases are different from each other at equilibrium. The vapor or gas phase is said to be richer in the more volatile or lighter or less soluble components of the mixture. Distillation employs heat to generate vapors and cooling to effect partial or total condensation as needed. Gas absorption employs a liquid of which the major components are essentially nonvolatile and which exerts a differential solvent effect on the components of the gas. In a complete plant, gas absorption is followed by a stripping operation for regeneration and recycle of the absorbent and for recovering the preferentially absorbed substances. In reboiled absorbers, partial stripping of the lighter components is performed in the lower part of the equipment. In distillation, absorption, or rectification and stripping are performed in the same equipment. Figures 13.1 and 13.2 show the basic types of equipment.

These distinctions between the two operations are partly traditional. The equipment is similar, and the mathematical treatment, which consists of material and energy balances and phase equilibrium relations, also is the same for both. The fact, however, that the bulk of the liquid phase in absorption-stripping plants is nonvolatile permits some simplifications in design and operation.

Equipment types are of two kinds, tray-type or packed, stagewise or continuous. The trays function as individual stages and produce stepwise changes in concentration. In packed towers concentration changes occur gradually. Until recently packed towers were used only in small equipment and where their construction was an advantage under corrosive conditions or when low pressure drop was mandatory. The picture now has changed somewhat and both types often are competitive over a wide range of sizes.

13.1. VAPOR-LIQUID EQUILIBRIA

This topic is concerned with the relations between vapor and liquid compositions over a range of temperature and pressure. Functionally, the dependence of the mol fraction y_i of component *i* in the vapor phase depends on other variables as

$$y_i = f(T, P, x_1, x_2, \dots, x_n).$$
 (13.1)



Figure 13.1. Distillation column assembly.

The dependence on composition alone often is approximated by

$$\mathbf{y}_i = \mathbf{K}_i \mathbf{x}_i, \tag{13.2}$$

where K_i , the vaporization equilibrium ratio (VER) is assumed to depend primarily on the temperature and pressure and the nature of the substance, and only secondarily on the composition. Equation (13.2) can be viewed as suggested by Raoult's law,

$$y_i = (P_i^{\text{sat}}/P)x_i \tag{13.3}$$

with

$$(K_i)_{\text{ideal}} = P_i^{\text{sat}}/P, \qquad (13.4)$$





where P_i^{sat} is the vapor pressure of component *i* and *P* is the system pressure. Several correlations have been developed for VERs, chiefly for hydrocarbon systems, for example, the one in Figure 13.3. The effect of composition is expressed in terms of a convergence pressure, which is explained for instance in the *API* Data Book (1969-date). The correction is small for system pressures under 10 atm or so and is neglected in this book.

A more nearly complete expression of K_i is derived upon

noting that at equilibrium partial fugacities of each component are the same in each phase, that is

$$\hat{f}_i^v = \hat{f}_i^L \tag{13.5}$$

or in terms of fugacity and activity coefficients,

$$y_i \hat{\phi}_i^v P = x_i \gamma_i \phi_i^{\text{sat}} P_i^{\text{sat}}.$$
(13.6)



Figure 13.3. Vaporization equilibrium ratios (Hadden and Grayson, 1961; Courtesy Mobil Oil Corp., New York).

The relation between mol fractions becomes

$$y_i = \frac{\gamma_i \phi_i^{\text{sat}} P_i^{\text{sat}}}{\hat{\phi}_i^v P} x_i, \tag{13.7}$$

which makes the vaporization equilibrium ratio

$$K_i = \frac{\gamma_i \phi_i^{\text{sat}} P_i^{\text{sat}}}{\hat{\phi}_i^v P}.$$
(13.8)

Additionally usually small corrections for pressure, called Poynting factors, also belong in Eq. (13.6) and following but are omitted here. The new terms are

- γ_i = activity coefficient in the liquid phase,
- ϕ_i^{sat} = fugacity coefficient of the pure component at its vapor pressure,
- $\hat{\phi}_i^v$ = partial fugacity coefficient in the vapor phase.

Equations for fugacity coefficients are derived from equations of state. Table 13.1 has them for the popular Soave equation of state. At pressures below 5-6 atm, the ratio of fugacity coefficients in Eq. (13.8) often is near unity. Then the VER may be written

$$K_i = \gamma_i P_i^{\text{sat}} / P \tag{13.9}$$

and is independent of the nature of the vapor phase.

Values of the activity coefficients are deduced from experimental data of vapor-liquid equilibria and correlated or extended by any one of several available equations. Values also may be calculated approximately from structural group contributions by methods called UNIFAC and ASOG. For more than two components, the correlating equations favored nowadays are the Wilson, the NRTL, and UNIQUAC, and for some applications a solubility parameter method. The first and last of these are given in Table 13.2. Calculations from measured equilibrium compositions are made with the rearranged equation

$$\gamma_i = \frac{\phi_i^v P}{\phi_i^{\text{sat}} P_i^{\text{sat}}} \frac{y_i}{x_i}$$
(13.10)

$$= \frac{P}{P_i^{\text{sat}}} \frac{y_i}{x_i}.$$
 (13.11)

The last approximation usually may be made at **pressures below** 5-6 atm. Then the activity coefficient is determined by the vapor pressure, the system pressure, and the measured equilibrium compositions.

Since the fugacity and activity coefficients are mathematically complex functions of the compositions, finding corresponding compositions of the two phases at equilibrium when the equations are known requires solutions by trial. Suitable procedures for making flash calculations are presented in the next section, and in greater detail in some books on thermodynamics, for instance, the *one* by Walas (1985). In making such calculations, it is usual to start by assuming ideal behavior, that is,

$$\hat{\phi}_i^v / \phi_i^{\text{sat}} = \gamma_i = 1. \tag{13.12}$$

After the ideal equilibrium compositions have been found, they are used to find improved values of the fugacity and activity coefficients. The process is continued to convergence. TABLE 13.1. The Soave Equation of State and Fugacity Coefficients

Equation of State

$$P = \frac{RT}{V-b} - \frac{a\alpha}{V(V+b)}$$
$$z^{3} - z^{2} + (A-B-B^{2})z - AB = 0$$

Parameters

a = $0.42747R^2T_c^2/P_c$, b = $0.08664RT_c/P_c$ α = $[1 + (0.48508 + 1.551710 - 0.15613\omega^2)(1 - <math>T_r^{0.5})]^2$ α = 1.202 exp($-0.30288T_r$) and for hydrogen (Graboski and Daubert, 1979) $A = a\alpha P/R^2T^2 = 0.42747\alpha P_r/T_r^2$ $B = bP/RT = 0.08664P_r/T_r$

Mixtures

$$a\alpha = \sum \sum y_i y_j (a\alpha)_{ij}$$
$$b = \sum y_i b_i$$
$$A = \sum \sum y_i y_j A_{ij}$$

$$B = \sum_{i} y_i B_i$$

Cross parameters

 $\begin{aligned} (a\alpha)_{ij} &= (1 - k_{ij})\sqrt{(a\alpha)_i}(a\alpha)_j \\ k_{ij} \text{ in table} \\ k_{ii} &= 0 \text{ for hydrocarbon pairs and hydrogen} \end{aligned}$

Correlations in Terms of Absolute Differences between Solubility Parameters of the Hydrocarbon, $\delta_{\rm HC}$ and of the Inorganic Gas

Gas	K _{ij}
H₂S CO₂ N₂	$\begin{array}{l} 0.0178 + 0.0244 \hspace{0.1cm} \left \delta_{HC} \hspace{0.1cm} 8.80 \right \\ 0.1294 - 0.0292 \hspace{0.1cm} \left \delta_{HC} \hspace{0.1cm} - 7.12 \right - 0.0222 \hspace{0.1cm} \left \delta_{HC} \hspace{0.1cm} - 7.12 \right ^2 \\ - 0.0836 + 0.1055 \hspace{0.1cm} \left \delta_{HC} \hspace{0.1cm} - 4.44 \hspace{0.1cm} 0.0100 \hspace{0.1cm} \left \delta_{HC} \hspace{0.1cm} - 4.44 \right ^2 \end{array}$

Fugacity Coefficient of a Pure Substance

$$\ln \phi = z - 1 - \ln \left[z \left(1 - \frac{b}{V} \right) \right] - \frac{a\alpha}{bRT} \ln \left(1 + \frac{b}{V} \right)$$
$$= z - 1 - \ln(z - B) - \frac{A}{B} \ln \left(1 + \frac{B}{z} \right)$$

Fugacity Coefficients in Mixtures

$$\ln \phi_{i} = \frac{b_{i}}{b}(z-1) - \ln\left[z\left(1-\frac{b}{V}\right)\right]$$
$$+ \frac{a\alpha}{bRT}\left[\frac{b_{i}}{b} - \frac{2}{a\alpha}\sum_{j}y_{j}(a\alpha)_{ij}\right]\ln\left(1+\frac{b}{V}\right)$$
$$= \frac{B_{i}}{B}(z-1) - \ln(z-B) + \frac{A}{B}\left[\frac{B_{i}}{B} - \frac{2}{a\alpha}\sum_{j}y_{j}(a\alpha)_{ij}\right]\ln\left(1+\frac{B}{z}\right)$$

(Walas, 1985).

TABLE 13.2. Activi	ty Coefficients from	Solubility	Parameters and	from the	Wilson E	quation
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Binary Mixtures					
Name	Parameters	In γ_1 and In γ_2			
Scatchard-Hildebrand	δ_1, δ_2	$\frac{V_1}{RT}(1-\phi_1)^2(\delta_1-\delta_2)^2$			
		$\frac{V_2}{RT}\phi_1^2(\delta_1-\delta_2)^2$			
	$\phi_1 = V_1 x_1 / (V_1 x_1 + V_2 x_2)$				
Wilson	$\lambda_{12}, \lambda_{21}$	$-\ln(x_1 + \Lambda_{12}x_2) + x_2 \left(\frac{\Lambda_{12}}{x_1 + \Lambda_{12}x_2} - \frac{\Lambda_{21}}{\Lambda_{21}x_1 + x_2}\right)$			
		$-\ln(x_2 + \Lambda_{21}x_1) - x_1 \left(\frac{\Lambda_{12}}{x_1 + \Lambda_{12}x_2} - \frac{\Lambda_{21}}{\Lambda_{21}x_1 + x_2}\right)$			
	$\Lambda_{12} = \frac{V_2^L}{V_1^L} \exp\left(-\frac{\lambda_{12}}{RT}\right)$	$\Lambda_{21} = \frac{V_1^L}{V_2^L} \exp\left(-\frac{\lambda_{21}}{RT}\right)$			
	V_i^L molar volume of	pure liquid component i.			
	Ternar	y Mixtures			
	$\ln \gamma_1 = 1 - \ln(x_1 \Lambda_{i1} + x_2 \Lambda_{i2})$	$+x_{3}\Lambda_{i3}$ - $\frac{x_{1}\Lambda_{1i}}{x_{1}+x_{2}\Lambda_{12}+x_{3}\Lambda_{13}}$			
	$=\frac{x_2\Lambda_{2i}}{x_1\Lambda_{2i}+x_2\Lambda_{2i}}$	$\frac{X_3\Lambda_{3i}}{X_2\Lambda_{3i}}$			
	$\Lambda_{1}\Lambda_{21} + \Lambda_{2} + \Lambda_{3}\Lambda_{3}$ $\Lambda_{ii} = 1$	$x_1 x_{31} + x_2 x_{32} + x_3$			
	Multicompo	onent Mixtures			
Eauation	Parameters	ln γ,			
Scatchard-Hildebrand	δ_i	$\frac{V_i}{RT} \left[\delta_i - \sum_j \frac{x_j V_j \delta_j}{\sum\limits_k x_k V_k} \right]^2$			
Wilson	$\Lambda_{ij} = \frac{V_j^L}{V_i^L} \exp\left(-\frac{\lambda_{ij}}{RT}\right)$	$-\ln\left(\sum_{j=1}^{m} x_j \Lambda_{ij}\right) + 1 - \sum_{k=1}^{m} \frac{x_k \Lambda_{ki}}{\sum\limits_{j=1}^{m} x_j \Lambda_{kj}}$			
	$\Lambda_{ij} = \Lambda_{jj} = 1$				

RELATIVE VOLATILITY

The compositions of vapor and liquid phases of two components at equilibrium sometimes can be related by a constant relative volatility which is defined as

$$\alpha_{12} = \frac{y_1}{x_1} / \frac{y_2}{x_2} = \left(\frac{y_1}{1 - y_1}\right) / \left(\frac{x_1}{1 - x_1}\right)$$
(13.13)

Then

$$\frac{y_1}{1-y_1} = \alpha_{12} \frac{x_1}{1-x_1} \,. \tag{13.14}$$

In terms of vaporization equilibrium ratios,

$$\alpha_{12} = K_1 / K_2 = \gamma_1 P_1^{\text{sat}} / \gamma_2 P_2^{\text{sat}}, \qquad (13.15)$$

and when Raoult's law applies the relative volatility is the ideal

value,

$$\alpha_{\text{ideal}} = P_1^{\text{sat}} / P_2^{\text{sat}}. \tag{13.16}$$

Usually the relative volatility is not truly constant but is found to depend on the composition, for example,

$$\alpha_{12} = k_1 + k_2 x_1. \tag{13.17}$$

Other relations that have been proposed are

$$\frac{y_1}{1-y_1} = k_1 + k_2 \left(\frac{x_1}{1-x_1}\right) \tag{13.18}$$

and

$$\frac{y_1}{1-y_1} = k_1 \left(\frac{x_1}{1-x_1}\right)^{k_2}.$$
(13.19)

A variety of such relations is discussed by Hala (Vapor-Liquid

EXAMPLE 13.1

Correlation of Relative Volatility

Data for the system ethanol + butanol at 1 atm are taken from the collection of Kogan et al. (1966, #1038). The values of x/(100 - x), y/(100 - y), and α are calculated and plotted. The plot on linear coordinates shows that relative volatility does not plot linearly with x, but from the linear log-log plot it appears that

$\frac{Y}{100-y} =$	4.364 (1	$\left(\frac{x}{00-x}\right)^{1}$	045 or	$\alpha = 4.364 \left(\frac{10}{10}\right)$	$\frac{x}{0-x}\Big)^{0.045}$
	X	Y	x	Y	
	0	0	39.9	74.95	
	3.45	12.5	53.65	84. 3	
	6.85	22.85	61.6	88. 3	
	10.55	32.7	70.3	91.69	
	14.5	41.6	79.95	95.08	
	18.3	49.6	S0. 8	97.98	
	28.4	63.45	6 100.0	100. 0	
	X	а	x/100-x	y/100 – y	
	3.5	4.00	0.04	0.14	
	6.9	4.03	0.07	0.30	
	10.6	4.12	0.12	0.49	
	14.5	4.20	0.17	0.71	
	18.8	4.25	0.23	0.98	
	26.4	4.38	0.40	1.74	
	39. 9	4.51	0.66	2.99	
	53. 7	4.64	1.16	5.37	
	61.6	4.70	1.60	7.55	
	70.3	4.66	2.37	11.03	
	80.0	4.85	3.99	19. 33	
	S0. 8	4.91	9.87	48.50	

Equilibria, Pergamon, London, 1967). Other expressions can be deduced from Eq. (13.15) and some of the equations for activity coefficients, for instance, the Scatchard-Hildebrand of Table 13.2. Then

$$\alpha_{12} = \frac{y_1}{x_1} / \frac{y_2}{x_2} = \frac{P_1^{\text{sat}}}{P_2^{\text{sat}}} \exp\left\{\frac{(\delta_1 - \delta_2)^2}{RT} \left[V_1(1 - \phi_1)^2 - V_2\phi_1^2\right]\right\},$$
(13.20)

where

$$\phi_1 = \frac{V_1 x_1}{V_1 x_1 + V_2 x_2} \tag{13.21}$$

is the volume fraction of component 1 in the mixture.

Beyond a certain complexity these analytical relations between vapor and liquid compositions lose their utility. The simplest one, Eq. (13.14), is of value in the analysis of multistage separating equipment. When the relative volatility varies modestly from stage to stage, a geometric mean often is an adequate value to use. Applications are made later. Example 13.1 examines two ways of interpreting dependence of relative volatility on composition.

BINARY x-y DIAGRAMS

Equilibria between the components of a binary mixture are expressed as a functional relation between the mol fractions of the usually more volatile component in the vapor and liquid phases,

$$\mathbf{Y} = \boldsymbol{f}(\boldsymbol{x}). \tag{13.22}$$

The definition of relative volatility, Eq. (13.14), is rearranged into this form:

$$y = \frac{\alpha x}{1 + (\alpha - 1)x}$$
(13.23)

Representative x-y diagrams appear in Figure 13.4. Generally they are plots of direct experimental data, but they can be calculated from fundamental data of vapor pressure and activity coefficients. The basis is the bubblepoint condition:

$$y_1 + y_2 = \frac{\gamma_1 P_1^{\text{sat}}}{P} x_1 + \frac{\gamma_2 P_2^{\text{sat}}}{P} (1 - x_1) = 1.$$
(13.24)

In order to relate y_1 and x_1 , the bubblepoint temperatures are found over a series of values of x_1 . Since the activity coefficients depend on the composition of the liquid and both activity coefficients and vapor pressures depend on the temperature, the calculation requires a respectable effort. Moreover, some vapor-liquid measurements must have been made for evaluation of a correlation of activity coefficients. The method does permit calculation of equilibria at several pressures since activity coefficients are substantially independent of pressure. A useful application is to determine the effect of pressure on azeotropic composition (Walas, 1985, p. 227).

13.2. SINGLE-STAGE FLASH CALCULATIONS

The problems of interest are finding the conditions for onset of vaporization, the bubblepoint; for the onset of condensation, the



Figure 13.4. Some vapor-liquid composition diagrams at essentially atmospheric pressure. This is one of four such diagrams in the original reference (*Kirschbaum*, Destillier und Rektifiziertechnik, *Springer, Berlin, 1969*). Compositions are in weight fractions of the first-named.

dewpoint; and the compositions and the relative amounts of vapor and liquid phases at equilibrium under specified conditions of temperature and pressure or enthalpy and pressure. The first cases examined will take the K_i to be independent of composition. These problems usually must be solved by iteration, for which the Newton-Raphson method is suitable. The dependence of K on temperature may be represented adequately by

$$K_i = \exp[A_i - B_i / (T + C_i)].$$
(13.25)

An approximate relation for the third constant is

$$C_i = 18 - 0.19T_{b_i}, \tag{13.26}$$

where T_{b_i} is the normal boiling point in ^oK. The dependence of K on pressure may be written simply as

$$\boldsymbol{K}_{i} = \boldsymbol{a}_{i} \boldsymbol{P}^{\boldsymbol{b}_{i}}.$$

Linear expressions for the enthalpies of the two phases are

$$\boldsymbol{h}_i = \boldsymbol{a}_i + \boldsymbol{b}_i \boldsymbol{T}, \tag{13.28}$$

$$H_i = c_i + d_i T, \tag{13.29}$$

assuming negligible heats of mixing. The coefficients are evaluated by readings off Figure 13.3, for example, and tabulations of pure component enthalpies. First derivatives are needed for application of the Newton-Raphson method:

$$\partial K_i / \partial T = B_i K_i / (T + C_i)^2, \qquad (13.30)$$

$$\partial K_i / \partial P = b_i K_i / P. \tag{13.31}$$

BUBBLEPOINT TEMPERATURE AND PRESSURE

The temperature at which a liquid of known composition first begins to boil is found from the equation

$$f(T) = \sum K_i x_i - 1 = 0, \qquad (13.32)$$

where the K_i are known functions of the temperature. In terms of Eq. (13.25) the Newton-Raphson algorithm is

$$T = T - \frac{-1 + \sum K_i x_i}{\sum [B_i K_i x_i / (T + C_i)^2]}.$$
 (13.33)

Similarly, when Eq. (13.27) represents the effect of pressure, the bubblepoint pressure is found with the N-R algorithm:

$$f(P) = \sum K_i x_i - 1 = 0, \qquad (13.34)$$

$$P = P - \frac{-1 + \sum a_i P^{b_i} x_i}{\sum a_i b_i P_i^{b_i - 1} x_i}.$$
(13.35)

DEWPOINT TEMPERATURE AND PRESSURE

The temperature or pressure at which a vapor of known composition first begins to condense is given by solution of the appropriate equation,

$$f(T) = \sum y_i / K_i - 1 = 0, \qquad (13.36)$$

$$f(P) = \sum y_i / K_i - 1 = 0.$$
(13.37)

In terms of Eqs. (13.25) and (13.27) the N-R algorithms are

$$T = T + \frac{-1 + \sum y_i/K_i}{\sum [(y_i/K_i^2) \partial K_i/\partial T]} = T + \frac{-1 + \sum y_i/K_i}{\sum [B_i y_i/K_i(T + C_i)^2]},$$
(13.38)
$$P = P + \frac{-1 + \sum y_i/K_i}{\sum [(y_i/K_i^2) \partial K_i/\partial P]} = P + \frac{(-1 + \sum y_i/K_i)P}{\sum (b_i y_i/K_i)}.$$
(13.39)

FLASH AT FIXED TEMPERATURE AND PRESSURE

At temperatures and pressures between those of the bubblepoint and dewpoint, a mixture of two phases *exists* whose amounts and compositions depend on the conditions that are imposed on the system. The most common sets of such conditions are fixed T and P, or fixed H and P, or fixed S and P. Fixed T and P will be considered first.

For each component the material balances and equilibria are:

$$Fz_i = Lx, + Vy_i, \tag{13.40}$$

$$\mathbf{y}_i = \mathbf{K}_i \mathbf{x}_i. \tag{13.41}$$

On combining these equations and introducing $\beta = V/F$, the fraction vaporized, the **flash** condition becomes

$$f(\beta) = -1 + \sum x_i \approx -1 + \sum \frac{z_i}{1 + \beta(K_i - 1)} = 0, \quad (13.42)$$

and the corresponding N-R algorithm is

$$\beta = \beta + \frac{-1 + \sum [z_i/(1 + \beta(K_i - 1))]}{\sum \{(K_i - 1)z_i/[1 + \beta(K_i - 1)]^2\}}$$
(13.43)

After β has been found by successive approximation, the phase compositions are obtained with

$$x_i = \frac{z_i}{1 + \beta(K_i - 1)},$$
 (13.44)

$$\mathbf{y}_i = \mathbf{K}_i \mathbf{x}_i. \tag{13.45}$$

A starting value of $\beta = 1$ always leads to a converged solution by this method.

FLASH AT FIXED ENTHALPY AND PRESSURE

The problem will be formulated for a specified final pressure and enthalpy, and under the assumption that the enthalpies are additive (that is, with zero enthalpy of mixing) and are known functions of temperature at the given pressure. The enthalpy balance is

$$H_{F} = (1 - \beta) \sum x_{i} H_{iL} + \beta \sum y_{i} H_{iV}$$
(13.46)

$$= (1 - \beta) \sum \frac{z_i H_{iL}}{1 + \beta(K_i - 1)} + \beta \sum \frac{K_i z_i H_{iV}}{1 + \beta(K_i - 1)}.$$
 (13.47)

This equation and the flash Eq. (13.42) constitute a set:

$$f(\beta, T) = -1 + \sum \frac{z_i}{1 + \beta(K_i - 1)} = 0, \qquad (13.48)$$

$$g(\beta, T) = H_F - (1 - \beta) \sum \frac{z_i H_{iL}}{1 + \beta(K_i - 1)}$$

$$-\beta \sum \frac{K_i z_i H_{iV}}{1 + \beta(K_i - 1)} = 0, \qquad (13.49)$$

from which the phase split β and temperature can be found when the enthalpies and the vaporization equilibrium ratios are known functions of temperature. The N-R method applied to Eqs. (13.48) and (13.49) finds corrections to initial estimates of β and T by solving the linear equations

$$h\frac{\partial f}{\partial \beta} + k\frac{\partial f}{\partial T} + f = 0, \qquad (13.50)$$

$$h\frac{\partial g}{\partial \beta} + k\frac{\partial g}{\partial T} + g = 0, \qquad (13.51)$$

where all terms are evaluated at the assumed values (β_0, T_0) of the two unknowns. The corrected values, suitable for the next trial if that is necessary, are

$$\beta = \beta_0 + h, \qquad (13.52)$$

$$T = T_0 + K.$$
 (13.53)

Example 13.2 applies these equations for dewpoint, bubblepoint, and flashes.

EQUILIBRIA WITH KS DEPENDENT ON COMPOSITION

The procedure will be described only for the case of bubblepoint temperature for which the calculation sequence is represented on Figure 13.5. Equations (13.8) and (13.32) are combined as

$$f(T) = \sum \frac{\gamma_i \phi_i^{\text{sat}} P_i^{\text{sat}}}{\hat{\phi}_i P} x_i - 1 = 0.$$
(13.54)

The liquid composition is known for a bubblepoint determination, but the temperature is not at the start, so that starting estimates must be made for both activity and fugacity coefficients. In the flow diagram, the starting values are proposed to be unity for all the variables. After a trial value of the temperature is chosen, subsequent calculations on the diagram can be made directly. The correct value of *T* has been chosen when $\sum y_i = 1$.

Since the equations for fugacity and activity coefficients are complex, solution of this kind of problem is feasible only by computer. Reference is made in Example 13.3 to such programs. There also are given the results of such a calculation which reveals the magnitude of deviations from ideality of a common organic system at moderate pressure.
EXAMPLE 13.2

Vaporization and Condensation of a Ternary Mixture For a mixture of ethane, n-butane, and n-pentane, the bubblepoint and dewpoint temperatures at 100 psia, a flash at 100°F and 100 psia, and an adiabatic flash at 100 psia of a mixture initially liquid at 100°F will be determined. The overall composition z_i , the coefficients *A*, *B*, and C of Eq. (13.22) and the coefficients a, *b*, *c*, and *d* of Eqs. (13.28) and (13.29) are tabulated:



The bubblepoint temperature algorithm is

$$T = T - \frac{-1 + \sum K_i x_i}{\sum [B_i K_i x_i / (T + C_i)^2]},$$
(13.33)

and the **dewpoint** temperature algorithm is

$$T = T + \frac{-1 + \sum y_i / K_i}{\sum [B_i y_i / K_i (T + C_i)^2]}$$
(13.38)

Results of successive iterations are

Bubblepoint	Dewpoint
1000. 0000	700.0000
695.1614	597.8363
560. 1387	625.9790
506. 5023	635. 3072
496. 1742	636.0697
495. 7968	636.0743
495. 7963	636.0743

The algorithm for the fraction vapor at specified T and P is

$$\beta = \frac{V}{F} = \beta + \frac{-1 + \sum z_i / (1 + \beta / (K_i - 1))}{\sum (K_i - 1) z_i / (1 + \beta (K_i - 1))^2},$$
(13.43)

and the equations for the vapor and liquid compositions are

$$\begin{aligned} x_i &= z_i / (1 + \beta(K_i - 1)), \end{aligned} \tag{13.44} \\ y_i &= K_i x_i. \end{aligned}$$

Results for successive iterations for β and the final phase compositions are

β	4	x _i	Y _i
1.0000 C,	0.3	0. 1339	0. 7231
0. 8257 nĈ	0.3	0.3458	0. 1833
0.5964 nC ₅	0.4	0. 5203	0.0936
0. 3986			
0. 3038			
0. 2830			
0.2819			

Adiabatic flash calculation: Liquid and vapor enthalpies off charts in the API data book are fitted with linear equations

$$h = a + bT \quad (^{\circ}\mathrm{F}), \tag{13.28}$$

$$H = c + dT$$
 ("F). (13.29)

The inlet material to the flash drum is liquid at 100°F, with $H_0 = 8,575.8$ Btu/lb mol. The flash Eq. (13.43) applies to this part of the example. The enthalpy balance is

$$H_{0} = 8575.8$$

= (1 - β) $\sum M_{i}x_{i}h_{i} + \beta \sum M_{i}y_{i}H_{i}$ (13.46)

$$= (1 - \beta) \sum \frac{M_i z_i h_i}{1 + \beta(K_i - 1)} + \beta \sum \frac{K_i M_i z_i H_i}{1 + \beta(K_i - 1)}.$$
 (13.47)

The procedure consists of the steps.,

1. Assume T.

- 2. Find the K_i , h_i , and H_i .
- 3. Find β from the flash equation (13.43).
- Evaluate the enthalpy of the mixture and compare with H₀, Eq. (13.47).

The results of several trials are shown:

Τ(°R) β Η

 530.00
 0.1601
 8475.70

 532.00
 0.1681
 8585.46

 531.82
 0.1674
 8575.58~ 8575.8, check.

The final VERs and the liquid and vapor compositions are:

	K	х	y
C ₂	4.2897	0.1935	0.8299
nC₄	0.3534	0.3364	0. 1189
nC5	0.1089	0.4701	0.0512

The numerical results were obtained with short computer programs which are given in Walas (1985, p. 317).

13.3. EVAPORATION OR SIMPLE DISTILLATION

As a mixture of substances is evaporated, the residue becomes relatively depleted in the more volatile constituents. A relation for binary mixtures due to **Rayleigh** is developed as follows: The differential material balance for a change dL in the amount of liquid remaining is

Upon rearrangement and integration, the result is

$$\ln\left(\frac{L}{L_0}\right) = \int_{x_0}^x \frac{dx}{x-y} \,. \tag{13.56}$$

In terms of a constant relative volatility

$$y = \frac{\alpha x}{1 + (\alpha - 1)x},\tag{13.57}$$

$$-y \, dL = d(LX) = L \, dx + X \, dL. \tag{13.55}$$



Figure 13.5. Calculation diagram for bubblepoint temperature (Walas, Phase Equilibria in Chemical Engineering, Butterworths, Stoneham, MA, 1985).

the integral becomes

$$\ln \frac{L}{L_0} = \frac{1}{x - 1} \ln \frac{x(1 - x_0)}{x_0(1 - x)} + \ln \frac{1 - x_0}{1 - x}.$$
(13.58)

MULTICOMPONENT MIXTURES

Simple distillation is not the same as flashing because the vapor is removed out of contact with the liquid as soon as it forms, but the process can be simulated by a succession of small flashes of residual liquid, say 1% of the original amount each time. After n intervals,

EXAMPLE 13.3 Bubblepoint Temperature with the Virial and Wilson Equations

A mixture of acetone (1) + butanone (2) + ethylacetate (3) with the composition $x_1 = x_2 = 0.3$ and $x_3 = 0.4$ is at 20 atm. Data for the system such as vapor pressures, critical properties, and Wilson coefficients are given with a computer program in Walas (198.5, **p**. 325). The bubblepoint temperature was found to be 468.7 K. Here only the properties at this temperature will be quoted to show deviations from ideality of a common system. The ideal and real K_i differ substantially.

the amount of residual liquid F is

$$F = L_0(1 - 0.01n) \tag{13.59}$$

and

$$\beta = \frac{V}{F} = \frac{0.01L_0}{(1 - 0.01n)L_0} = \frac{0.01}{1 - 0.01n}.$$
(13.60)

Then the flash equation (13.42) becomes a function of temperature,

$$f(T_n) = -1 + \sum \frac{z_i}{1 + 0.01(K_i - 1)/(1 - 0.01n)} = 0.$$
(13.61)

Here z_i is the composition at the end of interval n and K_i also may be taken at the temperature after interval n. The composition is found by material balance as

$$Lz_i = L_0(1 - 0.01n)z_i = L_0 \left[z_{i0} - 0.01 \sum_{k=1}^n y_{ik} \right],$$
(13.62)

where each composition y_{ik} of the flashed vapor is found from Eqs. (13.44) and (13.45)

$$y_i = K_i x_i = \frac{K_i z_i}{1 + 0.01(K_i - 1)/(1 - 0.01n)}$$
(13.63)

and is obtained during the process of evaluating the temperature with Eq. (13.61). The VERs must be known as functions of temperature, say with Eq. (13.25).

13.4. BINARY DISTILLATION

Key concepts of the calculation of distillation are well illustrated by analysis of the distillation of binary mixtures. Moreover, many real systems are essentially binary or can be treated as binaries made up of two pseudo components, for which it is possible to calculate upper and lower limits to the equipment size for a desired separation.

The calculational base consists of equilibrium relations and material and energy balances. Equilibrium data for many binary systems are available as tabulations of x vs. y at constant temperature or pressure or in graphical form as on Figure 13.4. Often they can be extended to other pressures or temperatures or expressed in mathematical form as explained in Section 13.1. Sources of equilibrium data are listed in the references. Graphical calculation of distillation problems often is the most convenient

Component	ϕ^{sat}	φ `	φ	at∕∳ ^v	γ
1	0.84363	0.843	53 1.0	0111	1.00320
2	0. 79219	0. 790	71 1.	00186	1.35567
3	0. 79152	0. 783	56 1.0	00785	1.04995
Compor	ient	K _{ideal}	K _{real}	1	Y
1	1	. 25576	1. 25591	0.3	779
2	0	. 72452	0.98405	0.2	951
3	0	. 77266	0.81762	0. 32	270

method, but numerical procedures may be needed for highest accuracy.

MATERIAL AND ENERGY BALANCES

In terms of the nomenclature of Figure 13.6, the balances between stage η and the top of the column are

$$V_{n+1}y_{n+1} = L_n x_n + D x_n, (13.64)$$

$$V_{n+1}H_{n+1} = L_nh_n + Dh_D + Q_c \tag{13.65}$$

$$= L_n h_n + DQ', \qquad (13.66)$$
where

$$Q' = h_D + Q_c / D \tag{13.67}$$

is the enthalpy removed at the top of the column per unit of overhead product. These balances may be solved for the liquid/vapor ratio as

$$\frac{L_n}{V_{n+1}} - \frac{y_{n+1} - x_D}{x_n - x_D} - \frac{Q' - H_{n+1}}{Q' - h_n}$$
(13.68)

and rearranged as a combined material and energy balance as



Figure 13.6. Model of a fractionating tower.

Similarly the balance between plate m below the feed and the bottom of the column can be put in the form

$$y_{m} = \frac{Q'' - H_{m}}{Q'' - h_{m+1}} x_{m+1} + \frac{h_{m+1} - H_{m}}{h_{m+1} - Q''} x_{B},$$
(13.70)

where

$$\mathbf{Q}'' = \mathbf{h}_B - \mathbf{Q}_b / B \tag{13.71}$$

is the enthalpy removed at the bottom of the column per unit of bottoms product.

For the problem to be tractable, the enthalpies of the two phases must be known as functions of the respective phase compositions. When heats of mixing and heat capacity effects are small, the enthalpies of mixtures may be compounded of those of the pure components; thus

$$H = yH_{a+} (1-y)H_{b}, (13.72)$$

$$h = xh_a + (1 - x)h_b, (13.73)$$

where H_a and H_b are vapor enthalpies of the pure components at their dewpoints and h_a and h_b are corresponding liquid enthalpies at their bubblepoints.

Overall balances are

$$F = D + B$$
, (13.74)

$$Fz_{,} = Dx_{,} + Bx_{,},$$
 (13.75)

$$FH_F = Dh, + Bh,. \tag{13.76}$$

In the usual distillation problem, the operating pressure, the feed composition and thermal condition, and the desired product compositions are specified. Then the relations between the reflux rates and the number of trays above and below the feed can be found by solution of the material and energy balance equations together with a vapor-liquid equilibrium relation, which may be written in the general form

$$f(x_n, y_n) = 0. (13.77)$$

The procedure starts with the specified terminal compositions and applies the material and energy balances such as Eqs. (13.64) and (13.65) and equilibrium relations alternately stage by stage. When the compositions from the top and from the bottom agree closely, the correct numbers of stages have been found. Such procedures will be illustrated first with a graphical method based on constant molal overflow.

CONSTANT MOLAL OVERFLOW

When the molal heats of vaporization of the two components are equal and the *tower* is essentially isothermal throughout, the molal flow rates L_n and V_n remain constant above the feed tray, and L_m and V_m likewise below the feed. The material balances in the two sections are

$$\mathbf{y}_{n+1} = \frac{L_n}{\mathbf{V}_{n+1}} \mathbf{x}_n + \frac{D}{\mathbf{V}_{n+1}} \mathbf{x}_D, \tag{13.64}$$

$$y_m = \frac{L_{m+1}}{V_m} x_{m+1} - \frac{B}{V_m} x_B.$$
(13.78)

The flow rates above and below the feed stage are related by the liquid-vapor proportions of the feed stream, or more generally by the thermal condition of the feed, q, which is the ratio of the heat

required to convert the feed to saturated vapor and the heat of vaporization, that is,

$$q = (H_F^{\text{sat}} - H_F) / (\Delta H)_{\text{vap}}$$
(13.79)

For instance, for subcooled feed q > 1, for saturated liquid q = 1, and for saturated vapor q = 0. Upon introducing also the reflux ratio

$$\boldsymbol{R} = \boldsymbol{L}_n / \boldsymbol{D}, \tag{13.80}$$

the relations between the flow rates become

$$L_m = L_n + qF = RD + qF,$$
 (13.81)
 $V_m = L_m - B = RD + aF - B.$ (13.82)

Accordingly, the material balances may be written

$$y = \frac{R}{R+1} x_n + \frac{1}{R+1} x_D,$$
 (13.83)

$$y_{m} = \frac{RD + qF}{RD + qF - B} x_{m+1} - \frac{B}{RD + qF - B} x_{B}.$$
 (13.84)

The coordinates of the point of intersection of the material balance lines, Eqs. (13.83) and (13.84), are located on a "q-line" whose equation is

$$y = \frac{q}{q-1}x + \frac{1}{q-1}x_F.$$
 (13.85)

Figure 13.7(b) shows these relations.











Figure 13.7. Features of McCabe-Thiele diagrams for constant molal overflow. (a) Operating line equations and construction and minimum reflux construction. (b) Orientations of q-lines, with slope = q/(q-1), for various thermal conditions of the feed. (c) Minimum trays, total reflux. (d) Operating trays and reflux. (e) Minimum reflux determined by point of contact nearest x_D .

BASIC DISTILLATION PROBLEM

The basic problem of separation by distillation is to find the numbers of stages below and above the feed stage when the quantities x_F , x_D , x_B , F, D, B, and R are known together with the phase equilibrium relations. This means that all the terms in Eqs. (13.83) and (13.84) are to be known except the running x's and y's. The problem is solved by starting with the known compositions, x_D and x_B , at each end and working one stage at a time towards the feed stage until close agreement is reached between the pairs (x, y_n) and (x_m, y_m) . The procedure is readily implemented on a programmable calculator; a suitable program for the enriching section is included in the solution of Example 13.4. A graphical solution is convenient and rapid when the number of stages is not excessive, which depends on the scale of the graph attempted.

Figure 13.7 illustrates various aspects of the graphical method. A minimum number of trays is needed at total reflux, that is, with no product takeoff. Minimum reflux corresponds to a separation requiring an infinite number of stages, which is the case when the equilibrium curve and the operating lines touch somewhere. Often this can occur on the q-line, but another possibility is shown on Figure 13.7(e). The upper operating line passes through point (x_D, x_D) and $x_D/(R + 1)$ on the left ordinate. The lower operating line passes through the intersection of the upper with the q-line and point (x_B, x_B) . The feed tray is the one that crosses the intersection of the operating lines on the q-line. The construction is shown with Example 13.5. Constructions for cases with two feeds and with two products above the feed plate are shown in Figure 13.8.

Optimum Reflux Ratio. The reflux ratio affects the cost of the tower, both in the number of trays and the diameter, as well as the cost of operation which consists of costs of heat and cooling supply and power for the reflux pump. Accordingly, the proper basis for choice of an optimum reflux ratio is an economic balance. The sizing and economic factors are considered in a later section, but reference may be made now to the results of such balances summarized in Table 13.3. The general conclusion may be drawn that the optimum reflux ratio is about 1.2 times the minimum, and also that the number of trays is about 2.0 times the minimum. Although these conclusions are based on studies of systems with nearly ideal vapor-liquid equilibria near atmospheric pressure, they often are applied more generally, sometimes as a starting basis for more detailed analysis of reflux and tray requirements.

Azeotropic and Partially Miscible Systems. Azeotropic mixtures are those whose vapor and liquid equilibrium compositions are identical. Their x-y lines cross or touch the diagonal. Partially miscible substances form a vapor phase of constant composition over the entire range of two-phase liquid compositions; usually the horizontal portion of the x-y plot intersects the diagonal, but those of a few mixtures do not, notably those of mixtures of methylethylketone and phenol with water. Separation of azeotropic mixtures, as illustrated by Example 13.6 for ethanol-water mixtures. Partially miscible constant boiling mixtures usually can be separated with two towers and a condensate phase separator, as done in Example 13.7 for n-butanol and water.

UNEQUAL MOLAL HEATS OF VAPORIZATION

Molal heats of vaporization often differ substantially, as the few data of Table 13.4 suggest. When sensible heat effects are small, however, the condition of constant molal overflow still can be preserved by adjusting the molecular weight of one of the components, thus making it a pseudocomponent with the same

molal heat of vaporization as the other substance. The x-y diagram and all of the compositions also must be converted to the adjusted molecular weight. Example 13.5 compares tray requirements on the basis of true and adjusted molecular weights for the separation of ethanol and acetic acid whose molal heats of vaporization are in the ratio 1.63. In this case, the assumption of constant molal overflow with the true molecular weight overestimates the tray requirements. A more satisfactory, but also more laborious, solution of the problem takes the enthalpy balance into account, as in the next section.

MATERIAL AND ENERGY BALANCE BASIS

The enthalpies of mixtures depend on their compositions as well as the temperature. Enthalpy-concentration diagrams of binary mixtures, have been prepared in general form for a few important systems. The most comprehensive collection is in Landolt-Bornstein **[IV4b**, 188, (1972)] and a few diagrams are in *Chemical Engineers Handbook* (1984), for instance, of ammonia and water, of ethanol and water, of oxygen and nitrogen, and some others. Such diagrams are named after Merkel.

For purposes of distillation calculations, a rough diagram of saturated vapor and liquid enthalpy concentration lines can be drawn on the basis of pure component enthalpies. Even with such a rough diagram, the accuracy of distillation calculation can be much superior to those neglecting enthalpy balances entirely. Example 13.8 deals with preparing such a Merkel diagram.

A schematic Merkel diagram and its application to distillation calculations is shown in Figure 13.9. Equilibrium compositions of vapor and liquid can be indicated on these diagrams by tielines, but are more conveniently used with associated x-y diagrams as shown with this figure. Lines passing through point P with coordinates (x_D, Q') are represented by Eq. (13.69) and those through point Q with coordinates (x_B, Q'') by Eq. (13.70). Accordingly, any line through P to the right of PQ intersects the vapor and liquid enthalpy lines in corresponding (x_n, y_{n+1}) and similarly the intersections of random lines through Q determine corresponding (x_{m+1}, y_m) . When these coordinates are transferred to the x-y diagram, they determine usually curved operating lines. Figure 13.9(b) illustrates the stepping off process for finding the number of stages. Points P, F, and Q are collinear.

The construction for the minimum number of trays is independent of the heat balance. The minimum reflux corresponds to a minimum condenser load Q and hence to a minimum value of $Q' = h_D + Q_c/D$. It can be found by trial location of point *P* until an operating curve is found that touches the equilibrium curve.

ALGEBRAIC METHOD

Binary systems of course can be handled by the computer programs devised for multicomponent mixtures that are mentioned later. Constant molal overflow cases are handled by binary computer programs such as the one used in Example 13.4 for the enriching section which employ repeated alternate application of material balance and equilibrium stage-by-stage. Methods also are available that employ closed form equations that can give desired results quickly for the special case of constant or suitable average relative volatility.

Minimum Trays. This is found with the Fenske-Underwood equation,

$$N_{\min} \frac{\ln[x_D(1 - x_B)/x_B(1 - x_D)]}{\ln \alpha}$$
(13.86)

EXAMPLE 13.4

Batch Distillation of Chlorinated Phenols

A mixture of chlorinated phenols can be represented as an equivalent binary with 90% 2,4-dichlorphenol (DCP) and the balance 2,4,6-trichlorphenol with a relative volatility of 3.268. Product purity is required to be 97.5% of the lighter material, and the residue must be below 20% of 2,4-DCP. It is proposed to use a batch distillation with 10 theoretical stages. Vaporization rate will be maintained constant.

- **a.** For operation at constant overhead composition, the variations of reflux ratio and distillate yield with time will be found.
- **b.** The constant reflux ratio will be found to meet the overhead and bottoms specifications.

a. At constant overhead composition, $y_D = 0.975$: The composition of the residue, x_{10} , is found at a series of reflux ratios between the minimum and the value that gives a residue composition of 0.2.

```
10 ! Example 13. 9. Distillation
    a t c
A=3.268
           constantYd
 20
    OPTION BASE 1
 30
    DIM X(10), Y(12)
 40
    Y(1)=.975
 50
60 INPUTR
 70
   FOR N=1 TO 10
80 X(N)=1/(A/Y(N)-A+1)
90 Y(N+1)=1/(R+1)*(R*X(N)+Y(1))
100 NEXT N
110 Z = (Y(1) - .9) / (Y(1) - X(10)) ! =
    L/Lo
120 I=(R+1)/(Y(1)-X(10))^2! Int
    earand of Eq. 4
```

- 130 PRINT USING 140 ; R,X(10),Z, I 140 IMPEE D DDDD 24 DDDD 24 D D
- 140 IMAGE D.DDDD,2X,.DDDD,2X,D.D DDD,2X,DDD.DDDDD 150 GOTO 60
- 160 END

With q = 1 and $x_n = 0.9$,

$$y_n = \frac{\alpha x_n}{1 + (\alpha - 1)x_n} = \frac{3.268(0.9)}{1 + 2.268(0.9)} \quad 0.9671,$$

$$R_m / (R_m + 1) = \frac{0.975 - 0.9671}{0.975 - 0.9} = 0.1051$$

$$\therefore R_m = 0.1174.$$

The btms compositions at a particular value of \mathbf{R} are found by successive applications of the equations

$$x_n = \frac{y_n}{\alpha - (\alpha - 1)y_n},\tag{1}$$

$$y_{n+1} = \frac{R}{R+1} \chi_n + \frac{1}{R+1} y_D \tag{2}$$

Start with $y_1 = y_D = 0.975$. The calculations are performed with the given computer program and the results are tabulated. The values of L/L, are found by material balance:

$$L/L_{2} = (0.975 - 0.900)/(0.975 - x_{L})$$
 (3)

The values of V/L, are found with Eq. (13.111).

$$\frac{V}{L_0} = (y_D - x_{L_0}) \int_{x_{L_0}}^{x_L} \frac{R+1}{(y_D - x_L)^2} dx_L$$

= (0.975 - 0.900) $\int_{0.9}^{x_L} \frac{R+1}{(0.975 - x_L)^2} dx_L.$ (4)

From the tabulation, the cumulative vaporization is

$$V/L_0 = 1.2566.$$

The average reflux ratio is

$$\bar{R} = \frac{V - D}{D} = \frac{V}{D} - 1 = \frac{V}{L_0 - L} - 1 = \frac{V/L_0}{1 - L/L_0} - 1$$
$$= \frac{I \cdot 2566}{1 - 0.0968} - 1 = 0.3913.$$

R	х _т	L/L	Integrand	V/L	t/ t
1174	0000	1 0001	198 69073	0 0 0 0 0	0 0 0 0
.11(4	2000	0000	1/5 17000	1116	0.000 001
.1300	0710	.0202	199 74019	1140	.021
.2000	8/h	.7383	122.74013	282 U 4995	. 224
2500	.8571	.6362	89.94213	4335	. 340
.3000	.8341	. 5321	65,43739	5675	,452
.3500	.8069	4461	4 7 75229	6830	
.4000	7760	3768	35,33950	. 7793	. 620
4500	7422	3222	26.76596	8580	683
5000	7069	2797	20 86428	9210	733
6000	6357	2210	13 99672	1 0170	่ ผู้ผู้ว่า
7000	EC04	10/0	10.02002	1.0100	őse
. (9999		- 1273	10 33322	1.0741	.000
.8000	.5111	.1617	8.36392	1.1150	.867
. 9000	.4613	1460	7.20138	1 i 440	
1.0000	4191	1349	6.47313	1.1657	. 928
1.2000	.3529	1206	5.68356	1 1959	. 952
1.4000	3040	. 1118	5.32979	1 2160	. 968
1 6000	2667	1059	5 18287	1 2708	.979
1 8000	5775	1017	5.14847	1 2421	<u></u> 988
9 0000	2141		5 15139	1 9511	300
2.0000 9.1400	2141	.0700	5.13136		, 990
2.1400	. 2002	8760	3.43097	1.2300	1.000

EXAMPLE **13.4**—(continued)

This is less than the constant reflux, $\mathbf{R} = 0.647$, to be found in part b.

At constant vaporization rate, the time is proportional to the cumulative vapor amount:

$$\frac{t}{\bar{t}} = \frac{V}{V_{\text{final}}} = \frac{V/L_0}{1.2566^7}$$
(5)

Also

$$D/L_0 = 1 - L/L_0.$$
 (6)

From these relations and the tabulated data, Q/L, and R are plotted against reduced time t/\bar{t} .

b.At constant reflux: A reflux ratio is found by trial to give an average overhead composition $\bar{y}_D = 0.975$ and a residue composition $x_L = 0.2$. The average overhead composition is found with material balance

$$\bar{y}_D = [x_{L0} - (L/L_0)x_L]/(1 - L/L_0).$$
⁽⁷⁾

The value of L/L, is calculated as a function of y_D from



At a trial value of R, values of x_{10} are found for a series of assumed y_D 's until x_{10} equals or is less than 0.20. The given computer program is based on Eqs. (1) and (2). The results of two trials and interpolation to the desired bottoms composition, $x_L = 0.200$, are

YD

XL

 $1/(y_D - x_L) = L/L_0$

ΫD

Reflux ratio R = 0.6

0.99805	0.9000	10. 2035		
0.99800	0.8981	10.0150	0.9810	
0. 99750	0.8800	8.5127	0.8295	
0.99700	0.8638	7.5096	0.7286	
0.99650	0.8493	6. 7917	0.6568	
0.99600	0.8361	6. 2521	0.6026	
0.99550	0.8240	5.8314	0.5602	
0.99500	0.8130	5.4939	0. 5263	
0.99400	0.7934	4. 9855	0.4750	
0.99300	0.7765	4.6199	0. 4379	
0.99200	0.7618	4. 3436	0.4100	
0.99100	0.7467	4. 1270	0. 3879	
0. 99000	0.7370	3.9522	0.3700	
0.98500	0.6920	3. 4135	0. 3135	
0. 98000	0.6604	3. 1285	0. 2827	
0.97500	0.6357	2.9471	0. 2623	
0.97000	0.6152	2.8187	0. 2472	
0.96500	0.5976	2.7217	0. 2354	
0.96000	0. 5819	2.8450	0. 2257	
0.95500	0.5678	2.5824	0. 2176	
0.95000	0.5548	2. 5301	0. 2104	
0. 90000	0. 4587	2.2662	0. 1671	
0. 85000	0.3923	2. 1848	0. 1441	
0. 80000	0.3402	2. 1751	0. 1286	
0. 75000	0. 2972	2. 2086	0. 1171	
0. 70000	0.2606	2.2756	0. 1079	0.9773
0.65000	0. 2286	2.3730	0. 1001	0.9746
0.60000	0. 2003	2. 5019	0. 0933	0.9720
0. 60000	0. 2003	2. 5019	0. 0933	0. 9720
0. 60000	0. 2003	2. 5019	0. 0933	0. 9720
0. 60000	0. 2003 Refi	2.5019 ux ratio R =	0. 0933 0.7	0. 9720
0. 60000	0. 2003 Refi	2. 5019	0. 0933 0.7	0. 9720
0. 99895	0. 2003 Refi 0. 9000	2. 5019 ux ratio R = 10. 1061	0. 0933 0.7	0. 9720
0. 60000 0. 99895 0. 99890	0. 2003 Refl 0. 9000 0. 8963	2. 5019 ux ratio R = 10. 1061 9. 7466	0. 0933 0.7 0. 9639	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885	0. 2003 Refi 0. 9000 0. 8963 0. 8927 0. 8927	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 0. 1201	0. 0933 0.7 0. 9639 0. 9312	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 9. 1241 0. 5055	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 9015	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99880 0. 99870	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8824	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 9. 1241 8. 5985 8. 5985	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8488	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99870 0. 99860 0. 99860	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8892 0. 8892 0. 8824 0. 8758	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 9. 1241 8. 5985 8. 1433 7. 4010	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.8032 0.8032	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99870 0. 99860 0. 99840	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8633	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 9. 1241 8. 5985 8. 1433 7. 4019 0. 5020	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 7288	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99880 0. 99860 0. 99840 0. 99820	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8518	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 9. 1241 8. 5985 8. 1433 7. 4019 6. 8306 0. 2004	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 9654	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99870 0. 99840 0. 99820 0. 99820 0. 99820	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8518 0. 8510	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.0875	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 6254 0. 4857	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99870 0. 99840 0. 99820 0. 99800 0. 99800 0. 99800 0. 99800	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965	2. 5019 10. 1061 9. 7466 9. 4206 9. 1241 8. 5985 8. 1433 7. 4019 6. 8306 6. 3694 4. 9875 4. 9807	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 6254 0. 4857 0. 4150	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99880 0. 99840 0. 99840 0. 99820 0. 99800 0. 99600	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 7631	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 1241 8. 5985 8. 1433 7. 4019 6. 8306 6. 3694 4. 9875 4. 2937 0. 2020	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 6254 0. 4857 0. 4160	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99860 0. 99840 0. 99820 0. 99820 0. 99800 0. 99700 0. 99600 0. 99500	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8518 0. 8518 0. 7965 0. 7631 0. 7370 0. 7370	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 9. 1241 8. 5985 8. 1433 7. 4019 6. 8306 6. 3694 4. 9875 4. 2937 3. 8760 2. 5059	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 6254 0. 4857 0. 4160 0. 3739 0. 245c	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99860 0. 99840 0. 99800 0. 99800 0. 99700 0. 99500 0. 99500 0. 99500 0. 99500	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 76631 0. 7370 0. 7159 0. 6020	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.9875 4.2937 3.8760 3.5958 2.2022	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.8032 0.7288 0.6716 0.6254 0.4857 0.4160 0.3739 0.3456	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99870 0. 99860 0. 99820 0. 99800 0. 99800 0. 99700 0. 99600 0. 99500 0. 99400 0. 99300	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 7631 0. 7370 0. 7159 0. 6983 0. 6983	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.9875 4.2937 3.8760 3.5958 3.3933 2.9415	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 6254 0. 4857 0. 44857 0. 44857 0. 44857 0. 3456 0. 3249 0. 904	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99870 0. 99840 0. 99840 0. 99800 0. 99800 0. 99600 0. 99500 0. 99500 0. 99300 0. 99400 0. 994	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 7631 0. 7631 0. 7759 0. 6983 0. 6835 0. 6835	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.9875 4.2937 3.8760 3.5958 3.3933 3.2415 2.6092	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.8032 0.7288 0.6716 0.6254 0.4857 0.4160 0.3739 0.3456 0.3249 0.3094	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99840 0. 99840 0. 99840 0. 99800 0. 99500 0. 99500 0. 99500 0. 99500 0. 99300 0. 99200 0. 99200 0. 99200 0. 99200	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 7631 0. 7370 0. 7370 0. 7370 0. 6983 0. 6983 0. 6985 0. 6076 0. 6076	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 1241 8. 5985 8. 1433 7. 4019 6. 8306 6. 3694 4. 9875 4. 2937 3. 8760 3. 5958 3. 3933 3. 2415 2. 6082 2. 2049	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.6032 0.7288 0.6716 0.6254 0.4857 0.4160 0.3739 0.3456 0.3249 0.3094 0.2969	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99870 0. 99860 0. 99840 0. 99820 0. 99800 0. 99700 0. 99600 0. 99500 0. 99300 0. 99300 0. 99200 0. 99100 0. 99200	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8822 0. 8824 0. 8758 0. 8633 0. 8518 0. 7631 0. 7370 0. 7159 0. 6835 0. 6076 0. 6594 0. 5095	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.9875 4.2937 3.8760 3.5958 3.3933 3.2415 2.6082 3.0248 2.5674	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.8032 0.7288 0.6716 0.6254 0.4857 0.4160 0.3739 0.3456 0.3249 0.3094 0.2969 0.2869 0.2869	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99860 0. 99840 0. 99840 0. 99840 0. 99800 0. 99700 0. 99600 0. 99500 0. 99400 0. 99300 0. 99200 0. 99100 0. 99000 0. 90000 0. 900000 0. 900000 0. 9000000 0. 9000000 0. 900000 0. 900000	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 76631 0. 7370 0. 7159 0. 6983 0. 6835 0. 6076 0. 6594 0. 5905 0. 5551	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.9875 4.2937 3.8760 3.5958 3.3933 3.2415 2.6082 3.0248 2.5674 2.3929	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 6254 0. 4857 0. 4160 0. 3739 0. 3456 0. 3249 0. 3094 0. 2969 0. 2366 0. 2151	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99870 0. 99840 0. 99840 0. 99800 0. 99700 0. 99600 0. 99500 0. 99400 0. 99300 0. 99400 0. 99300 0. 99400 0. 99300 0. 99400 0. 994	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 7631 0. 7370 0. 7159 0. 6983 0. 6835 0. 6076 0. 6594 0. 5505 0. 5521 0. 5521 0. 5524	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.9875 4.2937 3.8760 3.5958 3.3933 3.2415 2.6082 3.0248 2.5674 2.3929 2.2046	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.6032 0.7288 0.6716 0.6254 0.4857 0.4160 0.3739 0.3456 0.3249 0.3094 0.3094 0.2869 0.2869 0.2869 0.2366 0.2151 0.9015	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99870 0. 99840 0. 99840 0. 99800 0. 99800 0. 99600 0. 99500 0. 99500 0. 99400 0. 99300 0. 99200 0. 99100 0. 99200 0. 992	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 7631 0. 7370 0. 7159 0. 6983 0. 6835 0. 6076 0. 6554 0. 5521 0. 5521 0. 5242 0. 5012	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 4206 9. 1241 8. 5985 8. 1433 7. 4019 6. 8306 6. 3694 4. 9875 4. 2937 3. 8760 3. 5958 3. 3933 3. 2415 2. 6082 3. 0248 2. 5674 2. 3929 2. 2946 2. 2946 2. 2946	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.8032 0.7288 0.6716 0.6254 0.4857 0.4160 0.3739 0.3456 0.3249 0.3094 0.2969 0.28669 0.2366 0.2151 0.2015 0.9112	0. 9720
0. 60000 0. 99895 0. 99890 0. 99885 0. 99880 0. 99840 0. 99840 0. 99840 0. 99820 0. 99800 0. 99500 0. 99500 0. 99500 0. 99300 0. 99400 0. 99300 0. 99400 0. 994	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 7370 0. 7370 0. 7370 0. 7370 0. 7370 0. 7370 0. 6383 0. 6835 0. 6076 0. 6594 0. 5521 0. 5242 0. 5013 0. 4916	2. 5019 ux ratio R = 10. 1061 9. 7466 9. 1241 8. 5985 8. 1433 7. 4019 6. 8306 6. 8306 6. 8306 4. 9875 4. 2937 3. 8760 3. 5958 3. 3933 3. 2415 2. 6082 3. 0248 2. 5674 2. 3929 2. 2946 2. 2287 2. 1815	0.0933 0.7 0.9639 0.9312 0.9015 0.8488 0.8032 0.7288 0.6716 0.6254 0.4857 0.4160 0.3739 0.3456 0.3249 0.3094 0.2969 0.2366 0.2151 0.2015 0.1913 0.193	0. 9720
0. 60000 0. 99895 0. 99890 0. 99880 0. 99880 0. 99880 0. 99840 0. 99840 0. 99840 0. 99800 0. 99700 0. 99500 0. 99500 0. 99100 0. 99200 0. 99200 0. 99200 0. 99200 0. 99200 0. 99200 0. 99200 0. 99100 0. 99000 0. 90000 0. 900000 0. 900000 0. 90000 0. 90000 0. 90000 0. 90000 0. 9	0. 2003 Refl 0. 9000 0. 8963 0. 8927 0. 8892 0. 8824 0. 8758 0. 8633 0. 8518 0. 8410 0. 7965 0. 7661 0. 7370 0. 7159 0. 6983 0. 6835 0. 6076 0. 6594 0. 5905 0. 5521 0. 5521 0. 5524 0. 5013 0. 4816 0. 4820 0. 4816 0. 4816 0. 4816 0. 4820 0. 4816 0. 4820 0. 4816 0. 4816 0. 4820 0. 4816 0. 4816 0. 4820 0. 4816 0. 4820 0. 4816 0. 4820 0. 4816 0. 4820 0. 4816 0. 4820 0. 4816 0. 4820 0. 4816 0. 481	2.5019 ux ratio R = 10.1061 9.7466 9.4206 9.1241 8.5985 8.1433 7.4019 6.8306 6.3694 4.9875 4.2937 3.8760 3.5958 3.3933 3.2415 2.6082 3.0248 2.5674 2.3929 2.2946 2.2287 2.1815 2.1455	0. 0933 0.7 0. 9639 0. 9312 0. 9015 0. 8488 0. 8032 0. 7288 0. 6716 0. 6254 0. 4857 0. 4160 0. 3739 0. 3456 0. 3249 0. 3094 0. 2366 0. 2366 0. 2366 0. 2366 0. 2151 0. 2015 0. 1913 0. 1832 0. 1762	0. 9720

2. 1182

2.0982

2.0803

2.0454

2.0610

2.1101

2.1877

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2 4254

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0.92000

0.91000

0.90000

0.85000

0.80000

0.75000

0.70000

0.65000

0.60000

0.55000

0.50000

0.4479

0.4334

0.4193

0.3611

0.3148

0.2761

0.2429

0.2137

0.1877

0.1643

0.1431

0.1704

0.1652

0.1605

0.1423

0 1294

0.1194

0.1112

0.1041

0 0979

0.0923

0.0872

0.9773

0.9748

0.9723

EXAMPLE 13.5.

Distillation of Substances with Widely Different Molal Heats of Vaporization

The **molal** heats of vaporization of ethanol and acetic acid are 9225 and 5663 cal/g mol. A mixture with ethanol content of $x_F = 0.50$ is to be separated into products with $x_B = 0.05$ and $x_D = 0.95$. Pressure is 1 atm, feed is liquid at the boiling point, and the reflux ratio is to be 1.3 times the minimum. The calculation of tray requirements is to be made with the true molecular weight, 60.05, of acetic acid and with adjustment to make the apparent molal heat of vaporization the same as that of ethanol, which becomes

60.05(9225/5663) = 98.14.

The adjusted mol fractions, x' and y', are related to the true ones by

$$x' = \frac{x}{x + 0.6119(1 - x)}, \quad y' = \frac{y}{y + 0.6119(1 - y)},$$

The experimental and converted data are tabulated following and plotted on McCabe-Thiele diagrams. The corresponding compositions involved in this distillation are:

In terms of the true molecular weight, minimum reflux is given by

$$x_D/(R_{min} + 1) = 0.58,$$

whence

 $R_m = 0.6379,$ R = 1.3(0.6379) = 0.8293, $x_D/(R + 1) = 0.5193,$ $x'_D/(R + 1) = 0.5296.$

Taking straight operating lines in each case, the numbers of trays

are

N = 11.0 with true molecular weight of acetic acid, N' = 9.8 with adjusted molecular weight.

In this case it appears that assuming straight operating lines, even though the **molal** heats of vaporization are markedly different, results in overestimation of the number of trays needed for the separation.



a. Construction with true molecular weight, N = 11.



b. Construction with adjusted molecular weight, N = 9.8.



Figure 13.8. Operating and q-line construction with several feeds and top products. (a) One feed and one overhead product. (b) Two feeds and one overhead product. (c) One feed and two products from above the feed point.

TABLE 13.3. Economic Optimum Reflux Ratio for	Typical Petroleum Fraction Distillation near 1 atm
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	Factor for optimum reflux $f = (R_{opt}/R_m) - 1$ $R_{opt} = (1 + f)R_m$						Factor	for optimu N _{opt} /N _m	n trays		
	N _m = 10 R _m			$N_m = 20$ R_m		N _m = 50 R _m		N _m = 10 R _m	N _m = 20 R _m	N _m = 50 R _m	
	1	3	10	1	3	10	1	10	1 to 10	1 to 10	1 to 10
Base case . Payout time	0.20	0.12	0.10	0.24	0.17	0.16	0.31	0.21	2.4	2.3	2.1
Payout time 5 yr	0.13	0.09	0.07	0.17	0.13	0.10	0.22	0.15	2.2	2.5	2.2
Steam cost \$0.30/M lb Steam cost	0.22	0.13	0.11	0.27	0.16	0.14	0.35	0.22	2.3	2.1	2.0
\$0.75/M Ib $G_a = 50$	0.18	0.11	0.09	0.21	0.13	0.11	0.29	0.19	2.5	2.3	2.1
Ib mole/(hr)(sqft)	0.06	0.04	0.03	0.08	0.06	0.05	0.13	0.08	3.1	2.8	2.4

⁸ The "base case" is for payout time of 2 yr, steam cost of \$0.50/1000 lb, vapor flow rate $G_a = 15$ lb mol/(hr)(sqft). Although the capital and utility costs are prior to 1975 and are individually far out of date, the relative costs are roughly the same so the conclusions of this analysis are not far out of line. Conclusion: For systems with nearly ideal VLE, R is approx. 1.2 R_{min} and N is approx. 2.0 N_{min} .

(Happel and Jordan, Chemical Process Economics, Dekker, New York, 1975).

Minimum Reflux. Underwood's method employs two relations. First an auxiliary parameter θ is found in the range $1 < \theta < \alpha$ by solving

$$\frac{\alpha x_F}{\alpha - \theta} + \frac{1 - x_F}{1 - \theta} = 1 - q \tag{13.87}$$

$$(1 - q)\theta^{2} + [(a - 1)x_{F} + q(\alpha + 1) - \alpha]\theta - \alpha q = 0,$$
(13.88)

or in two important special cases:

or

when
$$q = 0$$
, $\theta = \alpha - (\alpha - 1)x_F$, (13.89)

when
$$q = 1$$
, $\theta = \frac{\alpha}{(\alpha - 1)x_F + 1}$ (13.90)

Then R_m is found by substitution into

$$R_m = -1 + \frac{\alpha x_D}{\alpha - \theta} + \frac{1 - x_D}{1 - \theta e}$$
(13.91)

Formulas for the numbers of trays in the enriching and stripping sections at operating reflux also are due to Underwood (Trans. *Inst. Chem.* Eng. **10**, 112-152, 1932). For above the feed, these groups of terms are defined:

$$K_{n} = L_{n}/V_{n} = R/(R+1),$$
 (13.92)

$$\phi_1 = K_1(\alpha - 1) / (K_1 \alpha - 1). \tag{13.93}$$

Then the relation between the compositions of the liquid on tray 1

EXAMPLE 13.6

Separation of an Azeotropic Mixture by Operation at Two Pressure Levels

At atmospheric pressure, ethanol and water form an azetrope with composition x = 0.846, whereas at 95 Torr the composition is about x = 0.94. As the diagram shows, even at the lower pressure the equilibrium curve hugs the x = y line. Accordingly, a possibly feasible separation scheme may require three columns, two operating at 760 Torr and the middle one at 9.5 Torr, as shown on the sketch. The basis for the material balance **used** is that 99% of the ethanol fed to any column is recovered, and that the ethanol-rich products from the columns have x = 0.8, 0.9, and 0.995, resp.

Although these specifications lead to only moderate tray and reflux requirements, in practice distillation with only two towers and the assistance of an azeotropic separating agent such as benzene is found more economical. Calculation of such a process is made by Robinson and Gilliland (1950, **p.** 313).





388 DISTILLATION AND GAS ABSORPTION

EXAMPLE 13.6(continued)



EXAMPLE 13.7

Separation of a Partially Miscible Mixture

Water and *n*-butanol in the concentration range of about 50-98.1 mol % water form two liquid phases that boil at 92.7°C at one atm. On cooling to 40° C, the hetero-azeotrope separates into phases containing 53 and 98 mol % water.

A mixture containing 12 mol % water is to be separated by distillation into products with 99.5 and 0.5 mol % butanol. The accompanying flowsketch of a suitable process utilizes two columns with condensing-subcooling to 40°C. The 53% saturated solution is refluxed to the first column, and the 98% is fed to the second column. The overhead of the second column contains a small amount of butanol that is recycled to the condenser for recovery. The recycle material balance is shown with the sketch.

The three sets of vapor-liquid equilibrium data appearing on the x-y diagram show some disagreement, so that great accuracy cannot be expected from determination of tray requirements, particularly at the low water concentrations. The upper operating line in the first column is determined by the overall material balance so it passes through point (0.995, 0.995), but the initial point on the operating line is at $\chi = 0.53$, which is the composition of the reflux. The construction is shown for 50% vaporized feed. That result and those for other feed conditions are summarized:

q	R _m	<i>R_m</i> = 1.3 <i>R_m</i>	N
1	2.02	2.62	12
0.5	5.72	7.44	8
0	9.70	12.61	6



Water	12	0.44	18.4139	0.7662	19.1801	6.8539	12.3262	11.56
Dutanal			6.1379	0.1916	6.3295	6.0779	0.2516	0.06
Butanoi	100	88.98	24.5518	0.9578	25.5096	12.9318	12.5778	11.62
% Wate	r 12	2 0.5	75	80	75.19	53	98	99.5

6

EXAMPLE 13.7—(continued)

In the second column, two theoretical trays are provided and are able to make a 99.6 mol % water waste, slightly better than the 99.5 specified. The required L/V is calculated from compositions read off the diagram:

L/V = (0.966 - 0.790)/(0.996 - 0.981) = 13.67.

If live steam were used instead of indirect heat, the bottoms concentration would be higher in water. This distillation is studied by Billet (1979, p. 216). Stream compositions are given below the flowsketch.

Equilibrium stage requirements for the separation of water and n-butanol.

and that on tray **n** is

$$(K_1\alpha)^{n-1} = \frac{1/(1-x_1) - \phi_1}{1/(1-x_n) - \phi_1}.$$
(13.94)

Since the overhead composition x_D is the one that is specified rather than that of the liquid on the top tray, x_1 , the latter is eliminated from Eq. (13.94). The relative volatility definition is applied

$$\frac{\alpha x_1}{1-x} = \frac{x_D}{1-x}, \qquad (13.95)$$

from which

$$\frac{1}{1-x_{o}} = \frac{x_{D} + \alpha(1-x_{D})}{\alpha(1-x_{D})}.$$
(13.96)

With this substitution, Eq. (13.94) becomes

$$(K_1\alpha)^{n-1} = \frac{[x_D + \alpha(1 - x_D)]/\alpha(1 - x_D) - \phi_1}{1/(1 - x_n) - \phi_1}.$$
 (13.97)

The number of trays above the feed plus the feed tray is obtained after substituting the feed composition x_F for x_n . Below the feed.

$$K_2 = V_m / L_m = (RD + qF - B) / (RD + qF),$$
(13.98)

$$\phi_2 = (\alpha - 1) / (K_2 \alpha - 1). \tag{13.99}$$

The relation between the compositions at the bottom and at tray m is

$$(K_2 \alpha)^m = \frac{1/x_B - \phi_2}{1/x_m - \phi_2}.$$
 (13.100)

The number of trays below the feed plus the feed tray is found after replacing x_m by x_F . The number of trays in the whole column then is

$$N = m + n - 1. \tag{13.101}$$

Example 13.9 applies these formulas.

TABLE 13.4. **Molal** Heats of Vaporization at Their Normal Boiling Points of Some Organic Compounds That May Need To Be Separated from Water

			Molecul	ar Weight
Compound	NBP (° C)	cai/g mol	True	Adjusted'
Water	100	9717	18.02	18.02
Acetic acid	118.3	5663	60.05	103.04
Acetone	56.5	6952	58.08	81.18
Ethylene alvcol	197	11860	62.07	50.85
Phenol	181.4	9730	94.11	94.0
n-Propanol	97.8	9982	60.09	58.49
Ethanol	78.4	9255	46.07	48.37

^e The adjustment of molecular weight is to make the molal heat of vaporization the same as that of water.

13.5. BATCH DISTILLATION

A batch distillation plant consists of a still or reboiler, a column with several trays, and provisions for reflux and for product collection. Figure 13.10(c) is a typical equipment arrangement with controls. The process is applied most often to the separation of mixtures of several components at production rates that are too small for a continuous plant of several columns equipped with individual reboilers, condensers, pumps, and control equipment.

EXAMPLE 13.8

Enthalpy-Concentration Lines of Saturated Vapor and Liquid of Mixtures of Methanol and Water at a Pressure of 2 atm A basis of 0°C is taken. Enthalpy data for methanol are in *Chemical* Engineers' Handbook (McGraw-Hill, New York, 1984, p. 3.204) and for water in Keenan et al. (Steam Tables: SI Units, Wiley, New York, 1978).



The number of continuous columns required is one less than the number of components or fractions to be separated. Operating conditions of a typical batch distillation making five cuts on an 8-hr cycle are in Figure 13.11.

Operation of a batch distillation is an unsteady state process whose mathematical formulation is in terms of differential equations since the compositions in the still and of the holdups on individual trays change with time. This problem and methods of solution are treated at length in the literature, for instance, by Holland and Liapis (*Computer Methods for Solving Dynamic Separation* Problems, 1983, pp. 177-213). In the present section, a simplified analysis will be made of batch distillation of binary mixtures in columns with negligible holdup on the trays. Two principal modes of operating batch distillation columns may be employed:

- With constant overhead composition. The reflux ratio is adjusted continuously and the process is discontinued when the concentration in the still falls to a desired value.
- With constant reflux. A reflux ratio is chosen that will eventually produce an overhead of desired average composition and a still residue also of desired composition.

Both modes usually are conducted with constant vaporization rate at an optimum value for the particular type of column construction. Figure 13.10 represents these modes on McCabe– Thiele diagrams. Small scale distillations often are controlled

> Water: $T = 120.6^{\circ}$ C $H_v = 11,652$ cal/g mol, $h_L = 2180$ cal/g mol, AH, = 9472 cal/g mol,

Experimental x-y data are available at 1 and 3 atm (Hirata, 1976, **#517**, **#519**). Values at 2 atm can be interpolated by eye. The lines show some overlap. Straight lines are drawn connecting enthalpies of pure vapors and enthalpies of pure liquids. Shown is the tie line for x = 0.5, y = 0.77.





Figure 13.9. Combined McCabe-Thiele and Merkel enthalpy-concentration diagrams for binary distillation with heat balances. (a) Showing key lines and location of representative points on the operating lines. (b) Completed construction showing determination of the number of trays by stepping off between the equilibrium and operating lines.

manually, but an automatic control scheme is shown in Figure 13.10(c). Constant overhead composition can he assured by control of temperature or directly of composition at the top of the column. Constant reflux is assured by flow control on that stream. Sometimes there is an advantage in operating at several different reflux rates at different times during the process, particularly with multicomponent mixtures as on Figure 13.11.

MATERIAL BALANCES

Assuming negligible holdup on the trays, the differential balance between the amount of overhead, dD, and the amount L remaining in the still is

$$y_D \ dD = -y_D \ dL = -d(Lx_L) = -L \ dx_L - x_L \ dL, \qquad (13.102)$$

which is integrated as

$$\ln(L/L_0) = \int_{x_{L_0}}^{x_L} \frac{1}{y_D - x_L} \, dx_L. \tag{13.103}$$

The differences $y_D - x_L$ depend on the number of trays in the

column, the reflux ratio, and the vapor-liquid equilibrium relationship. For constant molal overflow these relations may be taken as

$$y_{n+1} = \frac{R}{R+1} x_n + \frac{1}{R+1} y_D, \qquad (13.104)$$

$$y_n = f(x_n).$$
 (13.105)

When the overhead composition is constant, Eq. 13.103 is integrable directly, but the same result is obtained by material balance,

$$\frac{L}{L_0} = \frac{y_D - x_{L_0}}{y_D - x_L}.$$
(13.106)

With variable overhead composition, the average value is represented by the same overall balance,

$$\bar{y}_D = \frac{x_{L_0} - (L/L_0) x_L}{1 - (L/L_0)},$$
(13.107)

EXAMPLE 13.9

Algebraic Method for Binary Distillation Calculation

An equimolal binary mixture which is half vaporized is to be separated with an overhead product of 99% purity and 95% recovery. The relative volatility is 1.3. The **reflux** is to be selected and the number of trays above and below the feed are to be found with the equations of Section 13.4.6.

The material balance is

Component	F	D	X _D	B	Х _в
1	50	49. 50	0.99	0.50	0.0100
2	50	0.48	0.01	49. 52	0.9900
Total	-	49 98		50.02	

Minimum no. of trays,

1

$$N_m = \frac{\ln(0.99/0.01)(0.99/0.01)}{\ln 1.3} = 35.03.$$

For minimum reflux, by Eqs. (13.88) and (13.91),

$$\begin{split} 0.5\theta^2 + & [0.3(0.5) + 0.5(2.3) - 1.3]\theta - 1.3(0.5) = 0, \\ \theta^2 &= 1.3, \\ \theta &= 1.1402, \end{split}$$

but it is also necessary to know what reflux will result in the desired overhead and residue compositions.

For constant overhead composition at continuously varied reflux ratios, the total vaporization is found as follows. The differential balance is

$$dD = dV - dL = (1 - dL/dV) dV$$
(13.108)

The derivative dL/dV is the slope of the operating line so that

$$-\frac{dL}{dV} = 1 - \frac{R}{R+1} = \frac{1}{R+1}.$$
 (13.109)



$$R_{m} = -1 + \frac{1.3(0.99)}{1.3} + \frac{0.01}{1-0.1402} = 6.9813,$$

$$R = 1.2R_{m} = 8.3775,$$

$$K_{*} = \frac{R}{R+1} = 0.8934,$$

$$\phi_{1} = \frac{0.8934(1.3-1)}{0.8934(1.3)-1} = 1.6608,$$

$$\frac{1}{1-x_{1}} = \frac{0.99+1.3(0.01)}{1.3(0.01)} = 77.1538,$$

$$(K_{1}\alpha)^{n-1} = (1.1614)^{n-1} = \frac{77.1538-1.6608}{1/(1-0.5)-1.6608} = 222.56,$$

$$\therefore n = 37.12,$$

$$K_{2} = \frac{8.3775(49.98)+0.5(100)-50.02}{468.708} = 0.8933$$

$$\phi_{2} = \frac{1.3-1}{0.8933(1.3)-1} = 1.8600,$$

$$[0.8933(1.3)]^{m} = \frac{1/0.01-1.8600}{1/0.5-1.8600} = 701.00,$$

$$\therefore m = 43.82,$$

$$\therefore N = m + n - 1 = 37.12 + 43.82 - 1 = 79.94 \text{ trays.}$$

Substitution from Eqs. (13.103), (13.106), and (13.109) into Eq. (13.108) converts this into

$$dV = L_0(x_{L_0} - \bar{y}_D) \frac{R+1}{(x_L - \bar{y}_D)^2} dx_L, \qquad (13.110)$$

from which the total amount of vapor generated up to the time the residue composition becomes \mathbf{x}_{L} is

$$V = L_0(x_{L_0} - \bar{y}_D) \int_{x_{L_0}}^{x_L} \frac{R+1}{(x_L - \bar{y}_D)^2} dx_L.$$
(13.111)



Figure **13,10**. Batch distillation: McCabe-Thiele constructions and control modes. (a) Construction for constant overhead composition with continuously adjusted reflux rate. (b) Construction at constant reflux at a series of overhead compositions with an objective of specified average overhead composition. (c) Instrumentation for constant vaporization rate and constant overhead composition. For constant reflux rate, the temperature or composition controller is replaced by a flow controller.



Figure 13.10—(continued)



Figure 13.11. Operation of a batch distillation with five cuts.

At constant vaporization rate the time is proportional to the amount of vapor generated, or

$$t/\bar{t} = V/V_{\text{total}}.$$
(13.112)

Hence the *reflux* ratio, the amount of distillate, and the bottoms composition can be related to the fractional distillation time. This is done in Example 13.4, which studies batch distillations at constant overhead composition and also finds the suitable constant reflux ratio that enables meeting required overhead and residue specifications. Although the variable **reflux** operation is slightly more difficult to control, this example shows that it is substantially more efficient thermally-the average reflux ratio is much lower-than the other type of operation.

Equation (13.97) can be used to find the still composition-x, in that equation-at a particular reflux ratio in a column-reboiler combination with n stages. Example 13.4 employs instead a computer program with Equations (13.104) and (13.105). That procedure is more general in that a constant relative volatility need not be assumed, although that is done in this particular example.

13.6 MULTICOMPONENT SEPARATION: GENERAL CONSIDERATIONS

A tower comprised of rectifying (above the feed) and stripping (below the feed) sections is capable of making a more or less sharp separation between two products or pure components of the mixture, that is, between the light and heavy key components. The *light* key is the most volatile component whose concentration is to be controlled in the bottom product and the *heavy* key is the least volatile component whose concentration is to be controlled in the overhead product. Components of intermediate volatilities whose distribution between top and bottom products is not critical are called *distributed* keys. When more than two sharply separated products are needed, say n top and bottom products, the number of columns required will be n - 1.

In some cases it is desirable to withdraw sidestreams of intermediate compositions from a particular column. For instance, in petroleum fractionation, such streams may be mixtures of suitable boiling ranges or which can be made of suitable boiling range by stripping in small auxiliary columns. Other cases where intermediate streams may be withdrawn are those with minor but critical impurities that develop peak concentrations at these locations in the column because of inversion of volatility as a result of concentration gradient. Thus, pentyne-1 in the presence of n-pentane in an isoprene-rich C_5 cracked mixture exhibits this kind of behavior and can be drawn off as a relative concentrate at an intermediate point. In the rectification of fermentation alcohol, whose column profile is shown in Figure 13.12(a), undesirable esters and higher alcohols concentrate at certain positions because their solubilities are markedly different in high and low concentrations of ethanol in water, and are consequently withdrawn at these points.

Most distillations, however, do not develop substantial concentration peaks at intermediate positions. Figure 13.12(b) is of normal behavior.

SEQUENCING OF COLUMNS

The number n of top and bottom products from a battery of n - 1 columns can be made in several different ways. In a direct method, the most volatile components are removed one-by-one as overheads in successive columns with the heaviest product as the bottoms of the last column. The number of possible ways of separating components goes up sharply with the number of products, from two arrangements with three products to more than 100 with seven products. Table 13.5 identifies the five possible arrangements for



Figure 13.12. Concentration profiles in two kinds of distillations. (a) Purifying column for fermentation alcohol; small streams with high concentrations of impurities are withdrawn as sidestreams (*Robinson and Gilliland*, Elements of Fractional Distillation, *McGraw-Hill*, *New* York, 1939 *edition*). (b) Typical concentration profiles in separation of light hydrocarbon mixtures when no substantial inversions of relative volatilities occur (*Van Winkle*, Distillation, *McGraw-Hill*, *New York*, 1967).

separating four components with three columns. Such arrangements may differ markedly in their overall thermal and capital cost demands, so in large installations particularly a careful economic balance may be needed to find the best system.

TABLE 13.5. The Five Possible Sequences for the Separation of Four Components ABCD by Three Columns

Column 1		Column 2		Column 3	
Ovhd	Btms	O v h d	Btms	O v h d	Btms
Α	BCD	В	CD	С	D
Α	BCD	BC	D	В	C
AB	CD	Α	В	С	D
ABC	D	Α	ВC	В	C
ABC	D	AB	С	Α	В

The literature of optimum sequencing of columns is referenced by King (1980, pp. 711-720) and Henley and Seader (1981, pp. 527-555). For preliminary selection of near optimal sequences, several rules can be stated as guides, although some conflicts may arise between recommendations based on the individual rules. Any recommended cases then may need economic evaluations.

- 1. Perform the easiest separation first, that is, the one least demanding of trays and reflux, and leave the most difficult to the last.
- 2. When neither relative volatility nor concentration in the feed varies widely, remove the components one-by-one as overhead products.
- 3. When the adjacent ordered components in the process feed vary widely in relative volatility, sequence the splits in the order of decreasing relative volatility.
- 4. When the concentrations in the feed vary widely but the relative

volatilities do not, sequence the splits to remove components in the order of decreasing concentration in the feed.

NUMBER OF FREE VARIABLES

The performance of a given column or the equipment requirements for a given separation are established **by** solution of certain mathematical relations. These relations comprise, at every tray, heat and material balances, vapor-liquid equilibrium relations, and mol fraction constraints. In a later section, these equations will be stated in detail. For now, it can be said that for a separation of C components in a column of *n* trays, there still remain a number, C + 6, of variables besides those involved in the cited equations. These must be fixed in order to define the separation problem completely. Several different combinations of these C + 6 variables may be feasible, but the ones commonly fixed in column operation are the following:

ltem	Name	Number of Variables
1	feed rate	t
2	feed composition	C - I
3	feed enthalpy	1
4	ratio of overhead and feed rates	1
5	reflux enthalpy	1
6	reflux ratio, L/D or L/V	1
7	number of trays	1
8	column pressure	1
		<i>C</i> + 6

A common alternate specification is of the overhead and bottoms compositions expressed through distribution of the keys (two variables) as a replacement of items 4 and 7.

13.7. ESTIMATION OF REFLUX AND NUMBER OF TRAYS (FENSKE-UNDERWOOD-GILLILAND METHOD)

The first step in the design of distillation equipment is specification of the required distribution of light and heavy key components. Then the specific operating conditions and equipment size are established, ultimately on the basis of an economic balance or simply by exercise of judgment derived from experience. The design parameters that need to be determined include intermediate ones such as limiting reflux and trays that are needed for establishing a working design. These design parameters are the following:

- 1. Minimum number of theoretical trays,
- 2. Distribution of **nonkeys** between the overhead and bottoms products,
- 3. Minimum reflux,
- 4. Operating reflux,
- 5. Number of theoretical trays,
- 6. Location of the feed tray,
- 7. Tray efficiencies.

In packed towers, the variation of conditions from top to bottom is continuous and not interrupted as at trays. Nevertheless, it is convenient to speak of packing heights equivalent to a theoretical tray (HETU), so that tray tower theory can be applied to the design of packed towers.

All of the values of this list can be established at least approximately by rapid shortcut methods. In some instances such values may be useful as final ones, but ordinarily they are for exploratory purposes or as a starting basis for a computer design. Computer design of fractionation is an iterative process which depends for rapid convergence on good starting estimates of the principal quantities. The background of shortcut methods is well treated in the books of King (1980) and Henley and Seader (1981). Here attention will be directed to application of the techniques. These shortcut methods assume constant molal overflow in the rectifying and stripping zones and constant relative volatilities, which may be taken at the conditions of the feed tray or as a geometric mean of the values at the top and bottom of the column. Since the top conditions are not known completely in advance, evaluation of a mean relative volatility is an iterative process that can be started with the value at the feed tray or at the feed condition. Particular modes of variation of α sometimes are assumed. The method of Winn assumes that the vaporization equilibrium ratios vary as

$$K_{1k} = \beta K_{hk}^{\delta} \tag{13.113}$$

$$\alpha = K_{1k} / K_{hk} = \beta K_{hk}^{\delta - 1}$$
(13.114)

The constants β and δ for the conditions of the tower are deduced from log-log plots of *K*'s, which usually are available for hydrocarbons and natural gas constituents but can be evaluated from

$$K = \gamma P^{\text{sat}} / P, \tag{13.115}$$

with activity coefficient y of unity if no better information is known.

MINIMUM TRAYS

or

This is found from the relative volatility and the distribution of the keys between the overhead and bottoms by the Underwood-Fenske equation

$$N_m = \frac{\ln[(x_D/x_B)_{1k}/(x_D/x_B)_{hk}]}{\ln(\alpha_{1k}/\alpha_{hk})} = \frac{\ln[(d/b)_{1k}/(d/b)_{hk}]}{\ln(\alpha_{1k}/\alpha_{hk})}.$$
 (13.116)

In terms of the variation of VERs according to Eq. (13.113),

$$N_{m} = \frac{\ln[(d/b)_{1k}/(d/b)_{hk}^{\delta}]}{\ln\beta}$$
(13.117)

DISTRIBUTION OF NONKEYS

A convenient approximation is that the distributions of **nonkeys** require the minimum number of trays as given by Eq. (13.116). Designating the **nonkey** by subscript *nk*, that equation becomes

$$\ln(d/b)_{nk} = \ln(d/b)_{1k} + N_m \ln(\alpha_{nk}/\alpha_{1k})$$
(13.118)

$$(d/b)_{nk} = (d/b)_{1k} (\alpha_{nk}/\alpha_{1k})^{N_m}.$$
(13.119)

The distribution of **nonkeys** actually depends somewhat on the reflux ratio. For instance, in the case of Example 13.10, the distributions at minimum trays (total reflux) and minimum reflux are substantially different. Often it turns out, however, that the distributions predicted by Eq. (13.119) are close to those at finite reflux whenever R is near $1.2R_m$, which is often near the economic value for the reflux ratio. Further discussion of this topic is by Hengstebeck (Distillation, 1961) and Stupin and Lockhart (1968) whose work is summarized by King (1980, p. 434). Knowledge of the complete distribution is needed for estimation of top and bottom temperatures and for determination of the minimum reflux by the method to be cited.

EXAMPLE 13.10

Shortcut Design of Multicomponent Fractionation

A mixture of the given composition and relative volatilities has a thermal condition q = 0.8 and a pressure of 10 atm. It is to be fractionated so that 98% of component C and 1% of component E will appear in the overhead. The tray and reflux requirements are to be found. In the following table, the quantities in brackets are calculated in the course of the solution. f_i, d_i , and b_i are the mols of component *i* per mol of total feed.

	a	f	d	b
Α	3.1	0.03	[0.0300]	[1.5(<i>E</i> 5)]
В	2.6	0.07	IO.06981	[0.0002]
C lk	2.2	0.15	0.147	0.0030
D	1.3	0.33	[0.0481]"	[0.2819] "
D hk	1.0	0.30	0.003	0.297
F	0.8	0.12	[0.0000]	[0.1200]

^aThe corrected distribution of component D will be found along with the minimum reflux.

The minimum number of trays is

$$N_m = \frac{\ln\left[\frac{0.147}{0.003} / \frac{0.003}{0.297}\right]}{\ln 2.2} = 10\ 76$$

The distribution of component A is found as

$$\begin{pmatrix} \frac{d}{b} \\ i \end{pmatrix}_{i} = \begin{pmatrix} \frac{f-b}{b} \\ i \\ 0.0037 \end{pmatrix}_{i} = \begin{pmatrix} \frac{d}{b} \\ \frac{d}{b} \\ 10.76 \\ i \\ 1962 \end{pmatrix} = 1962$$

$$b_{i} = \frac{f_{i}}{1+(d/b)_{i}} - \frac{0.03}{1+1962} = 1.5(E-5)_{i}$$

$$d_{i} = f_{2} - b_{i} = 0.03 - 1.5(E-5) = 0.300.$$

Distributions of the other components are found in the same way. Since component D is distributed, two values of θ are found from Eq. (13.120):

$$\frac{3.1(0.03)}{3.1-\theta} + \frac{2.6(0.7)}{2.6-\theta} + \frac{2.2(0.15)}{2.2-\hat{e}} + \frac{1.3(0.33)}{1.3-\theta} + \frac{1(0.3)}{1-\theta} + \frac{0.8(0.12)}{0.8-\theta} = 1 - 0.8$$

$$\therefore \theta_1 = 1.8817, \quad \theta_2 = 1.12403.$$

The overhead content d_{D} of component D and the minimum reflux are found from the two equations

$$(\mathbf{R}, + 1)D = (\mathbf{R}, +1)(0.2498 + d,)$$

= $\frac{3.1(0.03)}{3.1 - \theta_1} - \frac{2.6(0.07)}{2.6 - \theta_1} + \frac{2.2(0.147)}{2.2 - \theta_1}$

$$\frac{1.3d_{\rm D}\ 0.003}{1.3 - \theta_1\ 1 - \theta_1}$$

= $\frac{3.1(0.03)}{3.1 - \theta_2} + \frac{2.6(0.007)}{2.6 - \theta_2} + \frac{2.2(0.147)}{2.2 - \theta_2}$
+ $\frac{1.3d_{\rm D}0.003}{1.3 - \theta_2\ i - e_1}$,

Upon substituting $\theta_1 = 1.8817$, $\theta_2 = 1.12403$,

$$d_{D} = 0.09311,$$

 $D = 0.2498 + 0.09311 = 0.3429,$
(R, + 1)D = 1.1342,
 $R_{m} = 2.3077.$

Let R = 1.2 $R_m = 1.2(2.3077) = 2.7692$. Apply Eq. (13.124):

$$X = \frac{R - R_m}{R + 1} - \frac{0.2(2.3077)}{3.7692} = 0.1225,$$

Y= 0.5313
$$N = \frac{N_m + Y}{1 - Y} = \frac{10.76 + 0.5313}{1 - 0.5313} = 24.1.$$

Feed plate location:

$$\frac{N_{\text{above}}}{N_{\text{below}}} = \frac{\ln\left(\frac{0.147}{0.15} / \frac{0.003}{0.300}\right)}{\ln\left(\frac{0.15}{0.003} / \frac{0.3}{0.297}\right)} = 1.175$$

Since $N_{above} + N_{below} = 24.1$,

feed tray =
$$\frac{24.1}{1 + 1/1.175}$$
 = 13 from the top.

For comparison, apply Eqs. (13.129) and (13.130):

$$\frac{N_r^*}{24 - N_r^*} = \left[\frac{0.6572}{0.3428} \left(\frac{0.30}{0.15}\right) \left(\frac{0.003/0.6572}{0.003/0.3428}\right)^2\right]^{0.206}$$

= 1.0088,
 $N_r^* = 12.05$,
 $N_r = 12.05 - 0.5 \log 24 = 10.46$ from the top.

Presumably 10.46 from the top is more accurate than 13.0, but it also may be in error because of the approximate fashion in which the distributions of **nonkeys** were found.

Note that the predicted distributions of component D do not agree closely.

			d	b
From	minimum	trays	0.0481	0.2819
From	minimum	reflux	0.09303	0.2370

MINIMUM REFLUX

The method of Underwood employs auxiliary parameters $\boldsymbol{\theta}$ derived from the equation

$$\sum_{i=1}^{C} \frac{\alpha_i x_{Fi}}{\alpha_i - \theta} = 1 - q, \qquad (13.120)$$

where q is the thermal condition of the feed and the summation extends over all the components in the feed. The only roots required are those in numerical value between the relative volatilities of the light and heavy keys. For instance, if there is one distributed component, subscript $_{dk}$, the required roots θ_1 and θ_2 are in the ranges

$$\begin{aligned} &\alpha_{1k} > \theta_1 > \alpha_{dk}, \\ &\alpha_{dk} > \theta_2 > \alpha_{hk}. \end{aligned}$$

Then the minimum reflux and the distribution of the intermediate component are found from the two equations that result from substitution of the two values of θ into Underwood's second equation

$$R_m + 1 = \frac{1}{D} \sum \frac{\alpha_i d_i}{\alpha_i - \theta}.$$
(13.121)

The number of values of θ and the number of Eqs. (13.121) is equal to 1 plus the number of components with relative volatilities between those of the light and heavy keys. When there is no distributed component, Eq. (13.121) may be used in terms of mol fractions and only a single form is needed for finding the minimum reflux,

$$R_m + 1 = \sum \frac{\alpha_i x_{iD}}{\alpha_i - \theta}.$$
(13.122)

Occasionally the minimum reflux calculated by this method comes out a negative number. That, of course, is a signal that some other method should be tried, or it may mean that the separation between feed and overhead can be accomplished in less than one equilibrium stage.

OPERATING REFLUX

As discussed briefly in Section 13.4, the operating reflux is an amount in excess of the minimum that ultimately should be established by an economic balance between operating and capital costs for the operation. In many cases, however, as stated there the assumptions $R = 1.2R_m$ often is close to the optimum and is used without further study unless the installation is quite a large one.

ACTUAL NUMBER OF THEORETICAL TRAYS

An early observation by Underwood (*Trans. Inst. Chem. Eng.* 10, pp. 112-152, 1932) of the plate-reflux relation was

$$(R - R_m)(N - N_m) = \text{const},$$
 (13.123)

but no general value for the constant was possible. Several correlations of calculated data between these same variables have since been made. A graphical correlation made by Gilliland (*Ind. Eng. Chem. 32*, 1101, 1940) has found wide acceptance because of its fair accuracy and simplicity of use. Of the several representations of the plot by equations, that of Molokanov et al. [*Int. Chem. Eng.*]

12, 209-212 (1972)] is accurate and easy to use:

$$Y = \frac{N - N_{\min}}{N+1} = 1 - \exp\left[\left(\frac{1+54.4X}{11+117.2X}\right)\left(\frac{X-1}{X^{0.5}}\right)\right],$$
 (13.124)

where

$$X = \frac{R - R_{\min}}{R + 1},$$
 (13.125)

from which the number of theoretical trays is

$$N = \frac{N_m + Y}{1 - \bar{y}},$$
(13.126)

The Gilliland correlation appears to be conservative for feeds with **low** values of q (the thermal condition of the feed), and can be in error when there is a large difference in tray requirements above and below the feed. The principal value of the correlation appears to be for preliminary exploration of design variables which can be refined by computer calculations. Although it is often used for final design, that should be done with caution. Other possibly superior but more difficult to use correlations have been proposed and are described in standard textbooks; for example, Hines and Maddox (1985).

FEED TRAY LOCATION

Particularly when the number of trays is small, the location of the feed tray has a marked effect on the separation in the column. An estimate of the optimum location can be made with the Underwood-Fenske equation (13.116), by applying it twice, between the overhead and the feed and between the feed and the bottoms. The ratio of the numbers of rectifying N, and stripping N_s trays is

$$\frac{N_r}{N_s} \approx \frac{\ln[(d/f)_{1k}/(d/f)_{hk}]}{\ln[(f/b)_{1k}/(f/b)_{hk}]}$$
(13.127)

$$= \frac{\ln[(x_d/x_f)_{1k}/(x_d/x_f)_{hk}]}{\ln[(x_f/x_b)_{1k}/(x_f/x_b)_{hk}]}$$
(13.128)

An improved relation that. however, requires more information is due to Akashah, Erbar, and Maddox [*Chem. Eng. Commun. 3,461* (1979)]. It is

$$N_{r} = N_{r}^{*} - 0.5 \log(N_{t}), \qquad (13.129)$$

where N, is the total number of trays in the column and N_r^* is given by the empirical Kirkbride (Petrol. *Refiner 23* (9), 321, 1944) equation,

$$\frac{N_r^*}{N_r - N_r^*} = \left[\frac{B}{D} \left(\frac{x_{1k}}{x_{hk}}\right)_f \left(\frac{x_{B1k}}{x_{Dhk}}\right)^2\right]^{0.206}.$$
(13.130)

TRAY EFFICIENCIES

The calculations made thus far are of theoretical trays, that is, trays on which vapor-liquid equilibrium is attained for all components. Actual tray efficiencies vary widely with the kind of system, the flow rates, and the tray construction. The range can be from less than 10% to more than 100% and constitutes perhaps the greatest uncertainty in the design of distillation equipment. For hydrocarbon fractionation a commonly used efficiency is about 60%. Section 13.14 discusses this topic more fully.

13.8. ABSORPTION FACTOR SHORTCUT METHOD OF EDMISTER

This method finds the product distribution ratio b/d for each component in a column with known numbers of trays above and below the feed and with a known reflux ratio. The flowsketch and nomenclature appear on Figure 13.13.

An absorption factor for each component i on each tray j is defined as

$$A_{i} = L_{j}/V_{j}K_{ij},$$
 (13.131)

but usually it is understood to apply to a specific component so the subscript i is dropped and the absorption factors on tray j become

$$A_j = L_j / V_j K_j. \tag{13.132}$$

Similarly a stripping factor for each component is defined as

$$S_j = K_j V_j / L_j.$$
 (13.133)

The ratio of bottom and overhead flow rates for each component is

$$\frac{b - \phi_1 + (L_d/DK_d)\phi_2 \quad (1 - q)F}{\psi_1 + (V_b/B)\psi_2 - 1},$$
(13.134)

with which the individual flow rates of each component are found

$$b_i = \frac{f_i}{1 + (b/d)_i},$$
(13.135)

$$\boldsymbol{d}_i = \boldsymbol{f}_i - \boldsymbol{b}_i. \tag{13.136}$$



Figure 13.13. Sketch and nomenclature for the absorption factor method.

The function ϕ and ψ are defined as

A

$$\phi_1 = \frac{A_e^{n+1} - 1}{A_e - 1},\tag{13.137}$$

$$\phi_2 = (A_1 A_n)^{n/2}, \tag{13.138}$$

$$\psi_1 = \frac{S_e^{m+1} - 1}{2}, \qquad (13.139)$$

$$\psi_2 = (S_1 S_m)^{m/2}.$$
(13.140)

The effective absorption and stripping factors in each zone are approximately

$$\mathbf{A}_{r} = -0.5 + \sqrt{A_{n}(A_{1} + 1)} + 0.25, \tag{13.141}$$

$$S_e = -0.5 + \sqrt{S_m(S_1 + 1) + 0.25}.$$
(13.142)

A certain number of initial estimates must be made when applying Edmister's method which are improved by iteration.

- **1.** Initial estimates must be made of the top and bottom temperatures so that the *A*, and S_1 can be estimated. These estimates will be adjusted by bubblepoint calculations after *b* and *d* have been found by the first iteration.
- The temperature at the feed zone may be found by taking a linear temperature gradient.
- **3.** Estimates must be made of V/L at the top and bottom and the feed zone. In distillation problems, assumption of constant molal overflow in each zone probably is within the accuracy of the method. In stripping or absorption columns, first iteration evaluations of the amounts of stripping or absorption will provide improved estimates of V/L at the key points in the columns.

A distillation problem is worked out by this method by **Edmister** [Pet. *Eng.*, 128-142 (Sept. 1948)]. The method is developed there.

For independent absorbers and strippers, the Kremser-Brown formulas apply. The fraction absorbed is

$$\phi_a = \frac{A_e^{n+1} - A_e}{A_e^{n+1} - 1},\tag{13.143}$$

and the fraction stripped is

$$\phi_s = \frac{S_e^{m+1} - S_e}{S_e^{m+1} - 1}.$$
(13.144)

An absorber is calculated by this method in Example 13.11.

13.9. SEPARATIONS IN PACKED TOWERS

Continuous changes in compositions of phases flowing in contact with each other are characteristic of packed towers, spray or wetted wall columns, and some novel equipment such as the **HIGEE** contactor (Fig. 13.14). The theory of mass transfer between phases and separation of mixtures under such conditions is based on a two-film theory. The concept is illustrated in Figure 13.15(a).

In its simplest form, the rate of mass transfer per unit area across these films is

$$N/A = k_G(y - y^*) = k_L(x^* - x).$$
(13.145)

Two special cases are commonly recognized.

EXAMPLE 13.11

Calculation of an Absorber by the Absorption Factor Method A mixture of a given composition is to have 60% of its n-butane removed by scrubbing with an oil in a 4-tray tower operating essentially isothermally at a pressure of 4 atm. The oil feed rate per 100 **mol** of feed gas will be found. The data are

	4	к	¢
C1	0. 253	54	
C ₂	0.179	14	
C3	0. 222	3.5	
nC ₄	0.240	0.5	0.600
nC ₅	0.105	0.2	
	1.000		

The Kremser-Brown formula (Eq. (13.143)) for the fraction absorbed is applied to nC_4 :

$$\phi = (A_e^5 - A_e)/(A_e^5 \quad 1) = 0.6,$$

 $\therefore A_e = 0.644, \text{ by trial.}$

Estimate that 27 mol of gas is absorbed. Let L_d represent the lean oil rate: For nC_4

$$A_1 = \frac{L_d}{KV_1} = \frac{L_d}{0.5(73)}, \quad A_n = \frac{L_d + 27}{0.5(100)}.$$

Substitute into Eq. (13.141),

$$A_e = -0.5 + \left[\frac{(L_d + 27)}{50} \left(\frac{L_d}{36.5} + 1\right) + 0.25\right]^{1/2} = 0.644,$$

$$\therefore L_d = 12.46, \text{ by trial.}$$

For the other components,

$$\begin{split} A_e &= -0.5 + \left[\frac{12.46 + 27}{100K} \left(\frac{12.46}{73K} + 1 \right) + 0.25 \right]^{1/2}, \\ \phi &= \frac{A_e^5 - A_e}{A_e^5 - 1}, \\ b &= 100 z_f \phi. \end{split}$$

The results are tabulated and show that the calculated value, 27.12, is close to the assumed, 27.00.

	Z _f	К	A,	φ	b
C1	0. 253	54	0.00728	0.00728	0. 18
C2	0.179	14	0. 02776	0.02776	0.50
C3	0. 222	3.5	0.1068	0.1068	2.37
nC₄	0.240	0.5	0.644	0.600	14.40
nC ₅	0. 105	0.2	1.4766	0. 9208	9.67
	1.000				27. 12

- Equimolal counterdiffusion between the phases, as in distillation with McCabe-Thiele approximations.
- Diffusion through a stagnant film, as in absorption or stripping processes involving the transfer of a single component between liquid and vapor phases. Since there is a concentration gradient

of the diffusing substance in the films, a correction is applied to the mass transfer coefficient. It is shown in books on mass transfer that the effective coefficient of a stagnant film is

$$(k_G)_{\text{effective}} = k_G / (y - y^*)_{\text{log mean}}, \qquad (13.146)$$

where

$$(y - \gamma^*)_{\log \text{mean}} = \frac{(1 - y) - (1 - y^*)}{\ln[(1 - y)/(1 - y^*)]} - \frac{(y^* - y)}{\ln[(1 - y)/(1 - y^*)]},$$
(13.147)

MASS TRANSFER COEFFICIENTS

Numerous investigations have been conducted of mass transfer coefficients in vessels with a variety of kinds of packings. Many of the more acceptable results are cited in recent books on mass transfer, for instance, those of Sherwood et al. (Mass Transfer, McGraw-Hill, New York, 1975), Cussler (*Diffusion*, Cambridge, 1984), and Hines and Maddox (1985). A convenient correlation of mass transfer coefficients in granular beds covering both liquid and vapor films is that of Dwivedi and Upadhyay [*Ind. Eng. Chem.* Process *Des. Deu.* 16, 157 (1977)], namely,

$$\varepsilon j_{d} = \frac{0.765}{\text{Re}^{0.82}} + \frac{0.365}{\text{Re}^{0.386}}$$
(13.148)

$$j_{d} = (\text{Sh})/(\text{Re})(\text{Sc})^{2/3}$$
(Chilton-Colburn factor), (13.149)
Sh = kd/\mathcal{D}
(Sherwood number), (13.150)
SC = $\mu/\rho \mathcal{D}$
(Schmidt number), (13.151)
Re = $du\rho/\mu = 4w/\pi d^{2}\mu$
(Reynolds number), (13.151)



Figure 13.14. A centrifugal packed fractionator, trade name HIGEE, Imperial Chemical Industries. Units have been operated with 500 times gravitational acceleration, with 3-18 theoretical stages, up to 36in. dia, employing perforated metal packing. For distillation, one unit is needed for rectification and one for stripping. Units have been used primarily for gas stripping and on offshore platforms because of compactness [Ramshaw, Chem. Eng., 13-14 (Feb. 1983)].



Figure 13.15. Mechanism, nomenclature, and constructions for absorption, stripping and distillation in packed towers. (a) Two-film mechanism with equilibrium at the interface. (b) Sketch and nomenclature for countercurrent absorption or stripping in a packed tower. (c) Equilibrium and material balance lines in absorption, showing how interfacial concentrations are found. (d) Equilibrium and material balance lines in stripping, showing how interfacial concentrations are found. (e) Equilibrium and material balance lines in distillation, showing how interfacial concentrations are found.

d = particle diameter,

- \mathcal{D} = diffusivity of the substance being transferred,
- k = mass transfer coefficient,
- \boldsymbol{u} = linear velocity of the fluid,
- w = mass rate of flow of the fluid,
- $\boldsymbol{\varepsilon}$ = fractional **voidage** between particles,
- p = density of the fluid,

$$\mu$$
 = viscosity of the fluid. (13.152)

Most of the properties change somewhat from one end to the other of industrial columns for effecting separations, so that the mass transfer coefficients likewise vary. Perhaps the property that has the most effect is the mass rate of flow which appears in the Reynolds number. Certainly it changes when there is a substantial transfer of material between the two phases in absorption or stripping; and even under conditions of constant molal overflow in distillation processes, the mass rate of flow changes because of differences of the molecular weights of the substances being separated. As a practical expedient, however, mass transfer coefficients are evaluated at mean conditions in a column.

DISTILLATION

Only the important case of constant molal **overflow** will be considered. The material balance around the lower end of the column of Figure 13.15(b) is

$$Gy + L_1 x_1 = G_1 y_1 + L x, (13.153)$$

which becomes at constant molal overflow

$$y = \frac{L}{G}x + \left(y_1 - \frac{L}{G}x_1\right). \tag{13.154}$$

The rate balance on an element of height dz of a column of unit cross section is

$$-dN = d(Gy) = G \, dy = k_G a(y \quad y^*) \, dz \tag{13.155}$$

$$= d(Lx) = L dx = k_L a(x^* - x) dz, \qquad (13.156)$$

where a is the interfacial surface per unit volume of the packed bed. These equations relate the interfacial concentrations (x^*, y^*) to those in the bulks of the liquid and gas phases (x, y); thus

$$\frac{y^* - y}{x^* - x} = -\frac{k_L}{k_G}$$
(13.157)

The bulk concentrations (x, y) are related by the material balance Eq. (13.144), and the equilibrium concentrations (x^*, y^*) from experimental data in graphical, tabular, or equation form,

$$y^* = f(x^*) \tag{13.158}$$

for instance, at constant relative volatility,

$$y^* = \frac{\alpha x^*}{1 + (\alpha - 1)x^*}.$$
 (13.159)

Corresponding points (y, y^*) in a column where the ratio k_L/k_G is known are found as follows: At a particular composition x, the

value of y is known from Eq. (13.154). Then corresponding values (x^*, y^*) are related linearly by Eq. (13.157). Substitution into Eq. (13.158) then will establish the value of y^* corresponding to the selected y. Figures 13.14(c), (d), (e) display graphical procedures for this operation.

By rearrangements of Eqs. (13.155) and (13.156) the height of the column is given by

$$Z = \frac{G}{k_G a} \int_{y_1}^{y_2} \frac{dy}{y^* - y}$$
(13.160)

$$=\frac{L}{k_{L}a}\int_{x_{1}}^{x_{2}}\frac{dx}{x-x^{*}}$$
(13.161)

The integrals in these equations are measures of the difficulty of the separation. Under some conditions they are roughly equal to the number of theoretical trays for the same change in concentration (y_1, y_2) or (x_1, x_2) . Accordingly, they are called numbers of transfer units.

$$NTU_G = \int_{y_1}^{y_2} \frac{dy}{y^* - y},$$
 (13.162)

$$NTU_L = \int_{x_1}^{x_2} \frac{dx}{x - x^*}.$$
 (13.163)

Consequently, it is natural to call the coefficients of the integrals the height of a transfer unit,

$$HTU, = G/k_{R}a, \tag{13.164}$$

$$HTU, = L/k_L a. \tag{13.165}$$

These terms sometimes are used interchangeably with height equivalent to a theoretical stage (HETS), but they are nearly the same only when the ratio k_L/k_G is a large number in the case of HTU_G . Example 13.12 studies this difference.

The concepts NTU and HTU are defined only for binary distillations and the transfer of a single substance in absorption or stripping. Since most processes of industrial interest involve multicomponents, the HETS of packed towers is the more useful concept, and may be evaluated readily from test data and tray calculations.

ABSORPTION OR STRIPPING

Neither mass nor molal flow rates are constant in these operations. In cases where essentially only one component is being transferred between phases, it is sometimes convenient to recognize the flow rates G' and L' of solute-free phases. They are related to the total flow rates by

$$G' = G(1 - y) = G_1(1 - y_1), \qquad (13.166)$$

$$L' = L(1 - x) = L_1(1 - x_1).$$
(13.16/)

The material balance around the lower end of the column of Figure 13.15(b),

$$Gy + L_1 x_1 = G_1 y_1 + Lx (13.168)$$

can be written

$$\frac{y}{1-y} = \frac{L'}{G'} \left(\frac{x}{1-x}\right) + \left(\frac{y_1}{1-y_1} - \frac{L'}{G'} \frac{x_1}{1-x_1}\right)$$
(13.169)

or in the linear form

$$Y = \frac{L'}{G'} X + \left(Y_1 - \frac{L'}{G'} X_1\right)$$
(13.170)

with the substitutions

$$X = \frac{x}{1-x},\tag{13.171}$$

EXAMPLE 13.12

Numbers of Theoretical Trays and of Transfer Units with Two Values of k_L/k_G for a Distillation Process

An equimolal mixture at its boiling point is to be separated into 95 and 5% contents of the lighter component in the top and bottoms products. The relative volatility is $\alpha = 2$, the minimum reflux is 1.714, and the operating reflux is 50% greater. The two values of k_L/k_G to be examined are -1 and ∞ .

The relation between interfacial and bulk concentrations is that of Eq. (13.157), $(y^* y)/(x^* - x) = -k_L/k_G$. At a series of values of x, corresponding values of y * and y may be read off with the graphical constructions shown on Figures (b) and (c) of this example. The values for slope = -1 are tabulated, but those for slope = ∞ are calculated from the equations of the equilibrium and operating lines and are not recorded. The integrands of Eq. (13.160) also are tabulated for both cases, and the numbers of transfer units are obtained by integration with the trapezoidal rule:

$$NTU = \int_{y_1}^{y_2} \frac{dy}{y^* - y}$$

- a. The number of theoretical trays stepped off on the McCabe-Thiele diagram is 16.2.
- **b.** With $k_L/k_G = 1$, the number of transfer units is 30.7.
- c. With $k_L/k_G = \infty$, the number of transfer units is 15.4.



(a) McCabe-Thiele construction showing that 16.2 trays are needed to contain 95 and 5% of the lighter substance in the products from a 50% boiling liquid feed.

$$Y = \frac{y}{1 - y}.\tag{13.172}$$

The equilibrium curve also can be transformed into these coordinates. These transformations are useful for graphical determinations of numbers of theoretical trays rather than for determination of numbers of transfer units. Example 13.13 employs both sets of units.



(b) Construction with $k_L/k_G = 1$, showing takeoff of vapor concentrations in the bulk, y, and at the interface, y*. Number of transfer found by integration = 15.4.



(c) Construction with $k_L/k_G = \infty$. Number of transfer units found by integration = 30.6.

EXAMPLE 13.12—(continued)

Within the_{accdfac}the trapezoidal rule integration and of the graphical determination of the number of trays, the numbers 16.2 and 15.4 are substantially the same. The infinite value of the ratio of mass transfer coefficients k_L/k_G means that all of the

resistance to mass transfer is in the film:

x	Y 🔰	1/(y*	y) 1/(y*- y
0. 05	0.05	0.068	22.105	55.56
0. 10	0.114	0.149	14. 745	28.57
0. 15	0.178	0.209	12.067	32.26
0.2	0.242	0. 279	10. 949	27.03
0. 25	0.306	0.345	10.638	25.64
0.3	0.370	0.411	10. 924	24.39
0. 35	0. 434	0.474	11.832	25.00

EXAMPLE 13.13

Trays and Transfer Units for an Absorption Process

The solute content of a gas with $y_1 = 0.40$ is to be reduced to $y_2 = 0.05$. The entering solvent is solute-free, $x_1 = 0$, and is to leave with $x_2 = 0.19$. The equilibrium relationship is represented by the equation

 $y^* = x^*(1 + 5x^*),$

and the ratio of mass transfer coefficients is $k_L/k_G = 1$.

In terms of solute-free coordinates, the equation of the material balance line is

Y = 2.6176X + 0.0526,

calculated with the given terminal concentrations. In terms of mol fractions the material balance line is curved, with equation

$$y = \frac{2.6176x/(1-x) + 0.0526}{2.6176x/(1-x) + 1.0526}$$

x	Ŷ	y †	$1/(y_{\infty}^* - y)$	$1/(y_1^* - y)$
0.4	0.498	0.536	13. 619	26.31
0.45	0.526	0.593	17.039	32.26
0.5	0.626	0.648	24.590	45.45
0.55	0.662	0.687	20.974	40.00
0.6	0.698	0.728	19. 231	33. 33
0.65	0.734	0.763	18.560	34.48
0.7	0.770	0.798	18.681	35. 71
0.75	0.806	0.832	19. 533	38.46
0.8	0.842	0.870	21.327	35. 71
0.85	0.878	0.902	24.439	41.67
0.9	0.914	0.933	29.969	52.63
0.95	0.950	0.965	41,053	66.67

The equation of the equilibrium curve in solute-free coordinates is

$$\frac{Y}{1+Y} = \frac{X}{1+X} \left(1 + \frac{5X}{1+X} \right).$$

Constructions for the numbers of trays in both sets of coordinates are made. They agree within the accuracy of graphical constructions on this scale, N = 4.7 with (x, y) and N = 4.5 with (X, Y).

For the transfer unit determination with the given ratio of mass transfer coefficients, corresponding values of (y, y^*) are found by intersections of the material balance and equilibrium lines with lines whose slopes are $-k_L/k_G = -1$ as indicated on Figure (a) and in detail with Example 13.12. These values are tabulated together with the corresponding integrands. The number of transfer units is found by trapezoidal rule integration of

NTU =
$$\int_{0.05}^{0.40} \frac{dy}{(1-y)\ln[(1-Y^*)/(1-y)]}$$

= 6.52.



The two values of N should be the same, but there is a small disagreement because of construction inaccuracies on this scale: (a) construction with mol fraction coordinates, N = 4.7; (b) construction with solute-free coordinates, N = 4.5.

EXAMPLE	13.13-(co)	ntinued)			х	v	v*	Integrand
	x	y	y *	Integrand				-
					0.10	0. 2556	0. 180	13.888
	0	0.05	0.009	24.913	0. 11	0.2733	0.202	14.703
	0.01	0.0733	0.020	19.296	0.12	0. 2906	0.224	15.709
	0.02	0.0959	0.036	17.242	0. 13	0.3074	0.246	16.998
	0.03	0.1178	0.052	15.757	0. 14	0.3237	0.268	18.683
	0.04	0.1392	0.069	14.818	0. 15	0.3397	0.290	20.869
	0.05	0.1599	0.086	14. 119	0. 16	0.3553	0.312	23.862
	0.06	0. 1801	0.102	13.405	0. 17	0.3706	0.335	28.877
	0.07	0. 1998	0.122	13.469	0. 18	0.3854	0.358	37.304
	0.08	0.2189	0. 141	13.467	0. 19	0.4000	0.381	53.462
	0.09	0. 2375	0.160	13. 548				

The rate balance on an element of height dz of a column of unit cross section, as in Figure 13.15(b), is

$$-dN = d(Gy) = (k_G)_{\text{eff}} a(y - y^*) dz$$
(13.173)
= $d(Lx) = (k_I)_{\text{eff}} a(x^* - x) dz.$ (13.174)

$$= a(Lx) = (\kappa_L)_{\text{eff}} a(x - x) az.$$
(1)

Expanding the differential of Eq. (13.163),

$$d(Gy) = d\left(\frac{G'}{1-y}\right) = \frac{G'}{(1-y)^2} \, dy = \frac{G}{1-y} \, dy.$$
(13.175)

Introducing Eqs. (13.146) and (13.175) into Eq. (13.173) and integrating, the height becomes

$$Z = \left(\frac{G}{k_G a}\right)_{\text{mean}} \int_{y_1}^{y_2} \frac{(y - y^*)_{1\text{m}}}{(1 - y)(y - y^*)} \, dy.$$
(13.176)

On replacing the log mean term by Eq. (13.147), the result becomes

$$Z = \left(\frac{G}{k_g a}\right)_m \int_{y_1}^{y_2} \frac{1}{(1-y)\ln[(1-y)/(1-y^*)]} \, dy.$$
(13.177)

The variable flow rate G is used here instead of the constant G' because the mass transfer coefficient k_G depends more directly on G. As used in Eqs. (13.176) and (13.177), a mean value of the coefficient is preferred in practice in preference to accounting for its variation within the integral.

The integrals are defined as numbers of transfer units for absorption or stripping,

NTU,
$$= \int_{y_1}^{y_2} \frac{1}{(1-y) \ln[(1-y)/(1-y^*)]} dy,$$
 (13.178)

$$NTU_L = \int_{x_1}^{x_2} \frac{1}{(1-x) \ln[(1-x)/(1-x^*)]} dx, \qquad (13.179)$$

and the heights of transfer units are

$$HTU_G = (G/k_G a)_{\text{mean}}, \tag{13.180}$$

$$\mathrm{HTU}_{L} = (L/k_{L}a)_{\mathrm{mean}}.$$
 (13.181)

HTUs vary with the kind of packing, the flow rates, the distribution of flow across the cross section, and sometimes with the packing height and column diameter. They are necessarily experimental data. Some of these data are discussed at the end of this chapter.

The way in which interfacial concentrations y^* are related to the bulk concentrations y required for evaluation of the integrand of Eq. (13.176) is explained on Figure 13.14(c), (d), and in Example 13.13, which finds trays and transfer units for an absorption problem.

13.10. BASIS FOR COMPUTER EVALUATION OF MULTICOMPONENT SEPARATIONS

Until the advent of computers, multicomponent distillation problems were solved manually by making tray-by-tray calculations of heat and material balances and vapor-liquid equilibria. Even a partially complete solution of such a problem required a week or more of steady work with a mechanical desk calculator. The alternatives were approximate methods such as those mentioned in Sections 13.7 and 13.8 and pseudobinary analysis. Approximate methods still are used to provide feed data to iterative computer procedures or to provide results for exploratory studies.

The two principal tray-by-tray procedures that were performed manually are the Lewis and Matheson and Thiele and Geddes. The former started with estimates of the terminal compositions and worked plate-by-plate towards the feed tray until a match in compositions was obtained. Invariably adjustments of the amounts of the components that appeared in trace or small amounts in the end compositions had to be made until they appeared in the significant amounts of the feed zone. The method of Thiele and Geddes fixed the number of travs above and below the feed, the reflux ratio, and temperature and liquid flow rates at each tray. If the calculated terminal compositions are not satisfactory, further trials with revised conditions are performed. The twisting of temperature and flow profiles is the feature that requires most judgement. The Thiele-Geddes method in some modification or other is the basis of most current computer methods. These two forerunners of current methods of calculating multicomponent phase separations are discussed briefly with calculation flowsketches by Hines and Maddox (1985).

Computer programs for multistage operations embodying heat and material balances and sophisticated phase equilibrium relations are best left to professionals. Most such work is done by service organizations that specialize in chemical engineering process calculations or by specialists in engineering organizations. A few valuable programs appear in the open literature:

- A Wang-Henke program appears in J. Christensen (Ed.) (Stagewise Computations-Computer Programs for Chemical Engineering Education, Sterling Swift Publishing, Manchaia, TX, 1972).
- A Naphthali-Sandholm program appears in Fredenslund, Gmehling, and Rasmussen (Vapor-Liquid Equilibria Using UNIFAC, Elsevier, New York, 1977).
- A Newton-Raphson SC (simultaneous correction) program of Newman is reproduced by King (Separation Processes, McGraw-Hill, New York, Appendix E).

Abundant descriptions of the theoretical basis and procedures for computer methods appear in recent literature and are summarized in books by Holland (1981), King (1980, Chap. 10), and Henley and Seader (1981, Chap. 15). The present chapter will be devoted to the basic equations, the kinds of process specifications that can be made and met, and convergence criteria applicable to iterative calculations of problems of distillation, absorption, and stripping. To a certain extent, the same methods are applicable to liquid-liquid extraction and other phase separation processes.

SPECIFICATIONS

The variables most commonly fixed in operations of distillation columns are listed in Section 13.6. Detailed calculation processes of column performance may require other intermediate or tentative specifications whose nature depends on the particular computer algorithm used. These specifications are identified with the descriptions of the three chief methods of this section.

THE MESH EQUATIONS

The letters of this acronym refer to Material balances, Equilibria between vapor and liquid, Summations of mol fractions to unity, and Heat or enthalpy balances. The quantities and notation pertaining to a single equilibrium stage and to an assembly of them are represented on Figure 13.16. In the simplest case a distillation stage exchanges two inlet and two outlet streams with adjacent stages. In addition, some stages will have in or out material or heat flows. Computer programs can be written in general form to include these factors on each stage to accommodate multiple feeds, side streams, and intermediate condensing or boiling. Enthalpy transfers sometimes are effected with hollow trays through which a heat transfer medium is circulated, or commonly by pumping a sidestream through an external heat exchanger and returning it to the column. The latter practice is particularly common for



Figure 13.16. Flow patterns and nomenclature of a single equilibrium stage and a cascade of them (after Henley and Seader, 1981). (a) A single equilibrium stage. (b) An assembly of N stages.



Figure 13.17. Algorithm of the BP (bubblepoint) method for distillation separations [Wang and Henke, Hydrocarbon Processing **45**(8), 155–166 (1963); Henley and Seader, 1981].

petroleum fractionation as an aid in controlling the wide range of vapor rates that accompany the difference of SIC-600°F between top and bottom of a crude oil fractionator. Side reflux of this kind requires more trays than all top reflux, but an overall benefit in equipment cost results because of diameter reduction.

For every component, C in number, on every stage, N in number, there are material, equilibrium, and energy balances, and the requirement that the **mol** fractions of liquid and vapor phases on each tray sum to unity. The four sets of these equations are:

1. *M* equations-Material balance for each component (C equations for each stage):

$$M_{ij} = L_{j-1}x_{i,j-1} + V_{j+1}y_{i,j+1} + F_j z_{ij} - (L, + U_j)x_{ij} - (V_j + W_i)y_{ij} = 0.$$
(13.182)

2. *E* equations-phase Equilibrium relation for each component (C equations for each stage):

$$E_{i,j} = y_{ij} - K_{ij} x_{ij} = 0, (13.183)$$

where K_{ij} is the phase equilibrium ratio.

3. S equations--mole fraction Summations (one for each stage):

$$(S_y)_j = \sum_{i=1}^C y_{ij} - 1.0 = 0,$$
 (13.184)

$$(S_x)_j = \sum_{i=1}^C x_{ij} - 1.0 = 0.$$
 (13.185)

Start

4. H equation—energy balance (one for each stage):

$$H_{j} = L_{j-1}H_{L_{j-1}} + V_{j+1}H_{V_{j+1}} + F_{j}H_{F_{j}} - (L_{j} + U_{j})H_{L_{j}} - (V_{j} + W_{j})H_{V_{i}} - Q_{j} = 0, \qquad (13.186)$$

where kinetic and potential energy changes are ignored.

In order to simplify these equations, the liquid rate at each stage is eliminated with the substitutions

$$L_{j} = V_{j+1} + \sum_{m=1}^{j} (F_{m} - U_{m} - W_{m}) - V_{1}, \qquad (13.187)$$

and the vapor compositions by the equilibrium relations

$$y_{ij} = K_{ij} x_{ij}.$$
 (13.188)

Three other variables occurring in the MESH equations are functions of more fundamental variables, namely,

$$K_{ij} = K(T_j, P_j, x_{ij}, y_{ij}),$$
(13.189)

$$H_{LJ} = H_L(T_j, P_j, x_j),$$
 (13.190)

$$H_{Vj} = H_V(T_j, P_j, y_j).$$
(13.191)

The reboiler load is determined by the overall energy balance,

$$Q_N = \sum_{j=1}^{N} (F_j H_{F_j} - U_j H_{L_j} - W_j H_{V_j}) - \sum_{j=1}^{N-1} Q_j - V_1 H_{V_1} - L_N H_{L_N}.$$
(13.192)

When all of the following variables are specified,

N,
$$F_i$$
, z_{ij} , T_j , P_j , U_j , W_j , and Q_j (except Q_1 and Q_N),
for $i = 1$ to C and $j = 1$ to N,

the MESH equations reduce in number to N(2C + 3) in the same number of variables, and are hence in principle solvable. The equations are nonlinear, however, and require solution by some iterative technique, invariably involving linearization at some stage in the calculation process.

Almost all computer programs employed currently adopt the Thiele-Geddes basis; that is, they evaluate the performance of a column with a specified feed, bottoms/overhead ratio, reflux ratio, and numbers of trays above and below the feed. Specific desired product distributions must be found by interpolation between an appropriate range of exploratory runs. The speed and even the possibility of convergence of an iterative process depends on the values of starting estimates of the variables to be established eventually. Accordingly, the best possible starting estimates should be made by methods such as those of Sections 13.7 and 13.8, or *on* the basis of experience.

After values of the variables T_i and V_j , called tear variables, are specified, Eqs. (182)ff become a linear set in the x_{ij} variables. Initial estimates of the vapor flows are made by assuming constant molal overflow modified by taking account of external inputs and outputs, and those of the temperatures by assuming a linear gradient between estimated top and bottom temperatures. Initially, also, the K_{ij} are taken as ideal values, independent of composition, and for later iterations the compositions derived from the preceding one may be used to evaluate corrected values of K_{ij} . With appropriate substitutions,

$$A_{j}x_{i,j-1} + B_{j}x_{ij} + C_{j}x_{i,j+1} = D_{j}, \qquad (13.193)$$

where

$$A_{j} = V_{j} + \sum_{m=1}^{j-1} (F_{m} - W_{m} - U_{m}) - V_{1}, \ 2 \le j \le N,$$
(13.194)

$$B_{j} = -\left[V_{j+1} + \sum_{m=1}^{j} (F_{m} - W_{m} - U_{m}) - V_{1} + U_{j} + (V_{j} + W_{j})K_{i,j}\right]$$

$$1 \le j \le N$$
(13.195)

$$C_{j} = V_{j+1}K_{i,j+1}, \ 1 \le j \le N - 1,$$
(13.196)

$$D_{i} = -F_{i}Z_{ii}, \ 1 \le j \le N,$$
(13.197)

the modified MESH equations can be written as a tridiagonal matrix, thus



The tridiagonal matrix is readily solved by computer by a method due to Thomas which is explained by Wang and Henke [Hydrocarbon Proc. 45(8), 155-163 (1966)] and by Henley and Seader (1981). A FORTRAN program is given by Gerald and Wheatley [Appl. Numer. Anal., 146, (1984)] and King (1980, Appendix E). A program in BASIC language is by Pachner (Handbook of Numerical Analysis Applications, McGraw-Hill, New York, 1984, Program P103).

After solution of the matrix for the liquid phase mol fractions x_{ij} , the next step is to make improved estimates of T_j and V_j for the next iteration. Three different procedures have been commonly employed for proceeding to succeeding trials, differing in simplicity or particular merit for certain kinds of problems.

- 1. BP (bubblepoint) methods. Temperatures are corrected iteratively by determinations of bubblepoints. The method is satisfactory for mixtures with relatively narrow ranges of volatilities. The parent program of this type is that of Wang and Henke (1966) which is flowsketched on Figure 13.17 and described in the next section. The availability of a FORTRAN program was cited earlier in this section.
- 2. SR (sum-rates) method. The new liquid flow rates are taken



Figure 13.18. Algorithm for the SR (sum rates) method for absorbers and strippers [*Birningham and* Otto, Hydrocarbon Processing 46(10), 163-170 (1967); Henley and Seader, 1981].

proportional to the nonnormalized sums of mol fractions, the vapor rates by subsequent material balances, and the new temperatures by enthalpy balances. A flowsketch of the calculation process is in Figure 13.18, and a brief description also is given subsequently. This method is particularly suited to separations involving substances with widely differing volatilities, as in absorbers and strippers, where the bubblepoint method breaks down.

3. SC (simultaneous correction) method. The MESH equations are reduced to a set of N(2C + 1) nonlinear equations in the mass flow rates of liquid components l_{ij} and vapor components v_{ij} and the temperatures T_j . The enthalpies and equilibrium constants K_{ij} are determined by the primary variables l_{ij} , v_{ij} , and T_j . The nonlinear equations are solved by the Newton-Raphson method. A convergence criterion is made up of deviations from material, equilibrium, and enthalpy balances simultaneously, and corrections for the next iterations are made automatically. The method is applicable to distillation, absorption and stripping in single and multiple columns. The calculation flowsketch is in Figure 13.19. A brief description of the method also will be given. The availability of computer programs in the open literature was cited earlier in this section.

THE WANG-HENKE BUBBLEPOINT METHOD

The procedure is outlined in Figure 13.17. The input data are listed above Box 1 and include all external material and enthalpy flows except condenser and reboiler loads, the number of trays, the reflux rate, and the reboiler load. The process is iterative, starting with estimates of temperature and vapor flow rates on each tray and making successive improvements in these values until a convergence criterion on temperatures is satisfied.

- Box 1. Initial estimates of the temperature are made by taking linear variation between estimated overhead **dewpoint** and bottoms bubblepoint. The vapor rates are estimated on the basis of constant rnolal overflow with due regard to input or output sidestreams.
- Box 2. The system represented by the matrix Eq. (13.198) consists of linear equations that are solved for the liquid mol fractions x_{ij} .
- Box 3. In general the mol fractions will not sum to unity, so that they are normalized as

$$(x_{ij})_{\text{normalized}} = x_{ij} / \sum_{i=1}^{C} x_{ij}.$$
 (13.199)



Figure **13.19.** True boiling point (TBP) curve of a crude oil, with superimposed TBP curves of five fractions into which it is separated by a typical fractionating system like that of Figure 15.20. The separations are not sharp, with as much as 50° F difference between the end point of a light product and the initial of the next heavier one. It is common to speak of the gap between the 95 and 5% points rather than the end points.

Box 4. New values of the stage temperatures T_j are calculated as bubblepoints with the normalized x_{ij} . Initially the effect of vapor compositions y_{ij} on K_{ij} is ignored and the vapor compositions are found with

$$\mathbf{v}_{ij} = \mathbf{K}_{ij} \mathbf{x}_{ij} \tag{13.200}$$

Subsequently, the values of y_{ij} from the previous iteration can be used in the evaluation of K_{ij} .

- Box 5. The enthalpies H_{Vj} and H_{Lj} can be evaluated with Eqs. (13.190) and (13.191) since T_j , P_j , x_{ij} , and y_{ij} have been estimated. The condenser load Q_1 is figured with Eq. (13.186) and the reboiler load Q_N with Eq. (13.192).
- Box 6. The new vapor rates V_j are found with the heat balances, Eqs. (13.201)-(13.205), and the new liquid rates with Eq. (13.187):

$$\alpha_j V_j + \beta_j V_{j+1} = \gamma_j, \qquad (13.201)$$

where

$$\boldsymbol{\alpha}_{j} = \boldsymbol{H}_{\boldsymbol{L}_{j-1}} \quad \boldsymbol{H}_{\boldsymbol{V}_{j}}, \tag{13.202}$$

$$\beta_{j} = H_{V_{j+1}} - H_{L_{j}}, \tag{13.203}$$

$$\gamma_{j} = \left[\sum_{m=1}^{N} (F_{m} - W_{m} - U_{m}) - V_{I}\right] (H_{L_{j}} - H_{L_{j-1}}) + F_{j}(H_{L_{j}} - H_{F_{j}}) + W_{j}(H_{V_{j}} - H_{L_{j}}) + Q_{j}, \qquad (13.204)$$

$$V_{j} = \frac{Y_{j-1} - \alpha_{j-1}V_{j-1}}{\beta_{j-1}}$$
(13.205)

Box 7. The convergence criterion imposes a tolerance on the differences between successive iterations of the temperatures

$$\tau = \sum_{j=1}^{N} (T_j^{(k)} - T_j^{(k-1)})^2 \le 0.01N.$$
(13.206)

Box 8. If the criterion is not satisfied, the values of T_i found in Box 4 and the vapor rates V_i of Box 6 are the new starting values to be input to Box 2.

THE SR (SUM-RATES) METHOD

In this method, temperatures for succeeding iterations are found by enthalpy balances rather than by bubblepoint determinations, after new values of the liquid and vapor flow rates have been estimated from solution of the equations for the liquid mol fractions. This procedure is suited to absorption and stripping problems for which the BP method breaks down because of the wide range of relative volatilities involved. The algorithm appears in Figure 13.18. Input data are the same as for the BP method.

- Box 1. Initial temperatures and vapor flow rates are estimated in the same way as in the BP method.
- Box 2. The mol fractions are found by solution of the tridiagonal matrix as in the BP method.
- Box 3. At this point the x_{ij} are not normalized but their sum is applied to estimate new liquid flow rates from the relation

$$L_{j}^{(k+1)} = L_{j}^{(k)} \sum_{i=1}^{C} x_{ij}.$$
(13.207)

The corresponding vapor rates are obtained by the material balance, which is a rearrangement of Eq. (13.187),

$$V_j = L_{j-1} - L_N + \sum_{m=j}^{N} (F_m - W_m - U_m).$$
(13.208)

Box 4. Then the x_{ij} are normalized by

$$(\mathbf{x}_{ij})_{\text{normalized}} = \mathbf{x}_{ij} / \sum_{i=1}^{C} \mathbf{x}_{ij}; \qquad (13.209)$$

the values of y_{ij} are obtained by

$$\mathbf{y}_{ij} = \mathbf{K}_{ij} \mathbf{x}_{ij} \tag{13.210}$$

and also normalized,

$$y_{ij} = y_{ij} / \sum_{i=1}^{C} y_{ij}.$$
 (13.211)

When the K_{ij} depend on the vapor phase compositions, values of y_{ii} from the previous iteration are used.

Box 5. New temperatures are calculated from the enthalpy balances Eq. (13.186). The temperature is implicit in these equations because of its involvement in the enthalpies and the K_{ij} . Accordingly, the temperature must be found by the Newton-Raphson method for simultaneous nonlinear equations.

Box 6. The convergence criterion is

$$\tau = \sum \left(T_j^{(k)} - T_r^{(k-1)} \right)^2 \le 0.01 N?$$
(13.212)

Box 7. If the convergence criterion is not satisfied, the values of V_i

from Box 3 and the temperatures from Box 5 are input to Box 2.

SC (SIMULTANEOUS CORRECTION) METHOD

A brief description of this procedure is abstracted from the fuller treatment of Henley and Seader (1981). The MESH equations (13.182)–(13.186) in terms of mol fractions are transformed into equations with molal flow rates of individual components in the liquid phase l_{ij} and vapor phase v_{ij} as the primary variables. The relations between the transformed variables are in this list:

$$L_{j} = \sum_{i=1}^{C} l_{ij}, \quad V_{j} = \sum_{i=1}^{C} v_{ij}, \quad x_{i,j} = \frac{l_{i,i}}{L_{i}}, \quad y_{ij} = \frac{v_{ij}}{V_{i}}, \\ f_{ij} = F_{j} z_{ij}, \quad s_{j} = U_{j}/L_{j}, \quad S_{j} = W_{j}/V_{j}. \quad (13.213)$$

The balance equations become three groups totalling N(2C + 1) in number:

Material balance:

$$M_{i,j} = l_{i,j}(1+s_j) + v_{ij}(1+S_j) - l_{ij-1} - v_{ij+1} - f_{ij} = 0.$$
(13.214)

Phase equilibria:

$$E_{i,j} = K_{ij} l_{ij} \frac{\sum_{\kappa=1}^{C} v_{\kappa j}}{\sum_{\kappa=1}^{C} l_{\kappa j}} - v_{ij} = 0.$$
(13.215)

Energy balance:

$$H_{j} = H_{L_{j}}(1+s_{j}) \sum_{i=1}^{C} l_{ij} + H_{V_{j}}(1+S_{j}) \sum_{i=1}^{C} v_{ij} - H_{L_{j-1}} \sum_{i=1}^{C} l_{ij-1} - H_{V_{j+1}} \sum_{i=1}^{C} v_{ij+1} - H_{F_{j}} \sum_{i=1}^{C} f_{ij} - Q_{j} = 0.$$
(13.216)

When N and all f_{ij} , P_F , P_j , s_j , S., and Q_i are specified, there remain N(2C + 1) unknowns, the same as the number of MEH equations (13.214)–(13.216). They are nonlinear equations in the primary variables l_{ij} , v_{ij} , and T_j for i = 1 to C and j = 1 to N. The T_j are involved implicitly in equations for the enthalpies and equilibrium constants.

The convergence criterion adopted is

$$\tau_{3} = \sum_{j=1}^{N} \left\{ (H_{j})^{2} + \sum_{i=1}^{C} \left[(M_{ij})^{2} + (E_{ij})^{2} \right] \le \varepsilon_{3} \\ = N(2C+1) \left(\sum_{j=1}^{N} F_{j}^{2} \right) 10^{-10}.$$
(13.217)

It will ensure that the converged variables will be accurate to generally at least four significant figures.

The algorithm of the procedure is in Figure 13.20.

- Box 1. Initial estimates of the stage temperatures are taken from linear variations between estimated overhead **dewpoint** and bottoms bubblepoint temperatures. Those of the vapor rates are based on the assumption of constant molal overflow with due regard to sidestreams, and those of the liquid rates are made consistent with the material flow balances.
- Box 2. With the initializations of Box 1, the matrix of the MEH equations is tridiagonal like Eq. (13.198) and may be solved for the l_{ii} and v_{ii} by the Thomas algorithm.

Box 3. Evaluate the discrepancy function made up of deviations from zero of the mass M, equilibrium E, and enthalpy H functions of Eqs. (13.214)–(13.216):

$$\tau_3 = \sum_{j=1}^{N} \left\{ (H_j)^2 + \sum_{i=1}^{C} \left[(M_{ij})^2 + (E_{ij})^2 \right] \right\}$$
(13.218)

Box 4. The discrepancy function τ_3 is compared with the tolerance ε_3

$$\varepsilon_3 = N(2C+1) \left(\sum_{j=1}^N F_j^2\right) 10^{-10}.$$
 (13.219)

If $\tau_3 \le \varepsilon_3$, the process has converged and final data are evaluated in Boxes 5 and 6. If $\tau_3 > \varepsilon_3$, proceed to the next iteration by way of Box 7.

Box 5. The total flow rates are found by summing up the component flow rates

$$L_j = \sum_{i=1}^{C} l_{ij} \tag{13.220}$$

and

$$V_j = \sum_{i=1}^{C} v_{ij}.$$
 (13.221)

- Box 6. Evaluate condenser and reboiler loads by heat balances if they have not been specified.
- Box 7. When $\tau_3 > \varepsilon_3$, corrections to the l_{ij} , v_{ij} , and T_j are calculated from the nonlinear MEH equations by the **Newton**-Raphson method. In these equations the enthalpies and equilibrium constants usually are nonlinear functions of the temperatures.
- Box 8. Employ a process for evaluating the optimum fraction of a calculated correction of each variable to be applied to the next trial. That is,

$$(\Delta a)_{\text{optimum}} = t(\Delta a)_{\text{calculated}}, \quad 0 < t \le 1.$$
(13.222)

The selection process is described by Henley and Seader. The optimally corrected values of l_{ij} , v_{ij} , and T_j are input to Box 4 for the next iteration.

13.11. SPECIAL KINDS OF DISTILLATION PROCESSES

Conditions sometimes exist that may make separations by distillation difficult or impractical or may require special techniques. Natural products such as petroleum or products derived from vegetable or animal matter are mixtures of very many chemically unidentified substances. Thermal instability sometimes is a problem. In other cases, vapor-liquid phase equilibria are unfavorable. It is true that distillations have been practiced successfully in some natural product industries, notably petroleum, long before a scientific basis was established, but the designs based on empirical rules are being improved by modern calculation techniques. Even unfavorable vapor-liquid equilibria sometimes can be ameliorated by changes of operating conditions or by chemical additives. Still, it must be recognized that there may be superior separation techniques in some cases, for instance, crystallization, liquid-liquid extraction, supercritical extraction, foam fractionation, dialysis, reverse osmosis, membrane separation, and others. The special distillations exemplified in this section are petroleum, azeotropic, extractive, and molecular distillations.



Figure 13.20. Algorithm of the SC (simultaneous correction) method for all multistage separations of fluid mixtures [Naphthali and Sandholm, AIChE J. 17, 148 (1971); Henley and Seader, 1981].

PETROLEUM FRACTIONATION

Crude oils are mixtures of many substances, mostly unidentified chemically, that cover a boiling range of less than 0°F to more than **1000°F**. Lower molecular weight substances are identifiable and may be recovered as pure substances, but the usual products of petroleum fractionation are mixtures with relatively narrow boiling ranges that have found consumer acceptance as final products or are suitable for further processing in the plant. On the typical refinery flow diagram of Figure 13.21, several of the processes represented as blocks either involve or are followed by distillation.

Important properties of petroleum and its fractions are measured by standardized procedures according to the API or ASTM. A particularly distinctive property is the true boiling point (TBP) curve as a function of the volume percent distilled under standardized conditions. Figure 13.19 is the TBP curve of a whole crude on which are superimposed curves of products that can be taken off sidestreams from a main distillation column, as in Figure 19.21. As samples of the distillate are collected, their densities and other properties of interest also are measured. The figure with Example 13.14 is of such measurements.

A representative petroleum fractionation process is summarized on Figure 13.22. Steam stripping of the sidestreams removes light ends and narrows the 95-5% temperature gap discussed in Example 13.14. The only source of heat supply to the column is at the feed point. A sufficient portion of the feed must be vaporized to be equivalent to the sum of all the products removed from the column above the feed point. Usually an additional amount of 2-5%, called overflash, may be needed to cover heat losses and reflux requirements. Because of the large temperature gradient and the high temperatures, the vapor volumes are large and also change greatly as the temperature falls along the column and sidestreams are withdrawn. Optimization of the size and cost of the fractionator usually requires removal of heat and provision of reflux at intermediate points rather than exclusively at the top as in most distillations, despite the need for additional trays to maintain efficient fractionation. The vapor rates at sidestream drawoffs usually are critical ones so they are checked by heat balances. Empirical rules have been developed for reflux ratios at drawoffs that ensure quality of these products.

The older, empirical practices for the design and operation of petroleum fractionators are stated in books such as that of Nelson



Figure 13.21. Petroleum refinery block diagram. Several of the processes identified by blocks include distillation or are followed by distillation (Gary and Handwerk, Petroleum Refining, Dekker, New York, 1975).

(Petroleum *Refinery Engineering*, McGraw-Hill, New York, 1958). Some such rules are collected in Table 13.6. A recent coverage of this subject is by Watkins (*Petroleum Refinery Distillation*, 1979), and an estimation procedure for distillations of naphthas without sidestreams is described by Broughton and Uitti [*Encycl. Chem. Process. Des.* 16, 186-198 (1982)]. An engineer versed in these techniques can prepare a near optimum design in a few days. For the most part, nowadays, only rough estimates of tray numbers and heat balances need be made as starting estimates for eventual computer design of the process.

The basis of the fractionation design is the true boiling point curve. This is replaced by a stepped curve made up of fractions boiling over ranges of 10-25°F. The lighter components up to pentanes or hexanes are treated as such, but the other components are pseudocomponents characterized by their average boiling points, specific gravities, molecular weights, and other properties necessary to calculation of the distillation behavior. For full range crude oil fractionation, as many as 50 pseudocomponents may be required to represent the real TBP curve. In the case of naphtha fractionators without sidestreams, 20 pseudocomponents may be sufficient. Calculated compositions of products in terms of pseudocomponents can be reconstituted into smooth TBP curves to ensure that conventional specifications such as initial and final boiling points are met. The operation of converting a mixture characterized by TBP and specific gravity curves into a mixture of a discrete number of components with compositions expressed in mol fractions is performed in Example 13.14.

EXTRACTIVE DISTILLATION

In such a process an additive or solvent of low volatility is introduced in the separation of mixtures of low relative volatilities or for concentrating a mixture beyond the azeotropic point. From an extractive distillation tower, the overhead is a finished product and the bottoms is an extract which is separated down the line into a product and the additive for recycle. The key property of the additive is that it enhance the relative volatilities of the substances to be separated. From a practical point of view, the additive should be stable, of low cost, require moderate reboiler temperatures particularly for mixtures subject to polymerization or thermal degradation, effective in low to moderate concentrations, and easily recoverable from the extract. Some common additives have boiling points 50–100°C higher than those of the products.

Selection of an Additive. Ultimately the choice of an extractive distillation solvent will require a certain amount of experimental work, but some screening process should be employed to limit its scope. Examination of solvents that are being used or have been studied for successful commercial operations is a starting point. Some rules involving similarities or differences in polarities or hydrogen bonding have been proposed. The less soluble of a pair of substances usually will have the enhanced volatility. Accordingly, a comparison of solubility parameters may be a guide: A good additive should have a solubility parameter appreciably different from one of the components and closer to that of the other. Such an

EXAMPLE 13.14

Representation of a Petroleum Fraction by an Equivalent

Number of Discrete Components true boiling point and specific gravity variation with the volume The percent distilled are found by standard ASTM procedures. In the present case, the smooth TBP curve is replaced by a stepped curve of eleven pseudo components characterized by their 50% boiling points and specific gravities. Their molecular weights are obtained with the general correlation of Figure (c); then the mol fractions are calculated. Vaporization equilibrium ratios and relative volatilities can be read off charts such as Figure 13.3, which are available for higher boiling ranges than this one. Then any required distillation can be calculated by any suitable standard method.



(a) Experimental true boiling point and specific gravity curves, and the equivalent stepped curve. (b) Mol fraction composition of the 11 pseudo components with equivalent vaporization behavior. (c) Standard correlations of properties of petroleum fractions.


Figure 13.22. Material and energy flows in distillation of 20,000 BPSD (263,000 lb/hr) of 28.4" API crude oil into five products. The main tower is 11 ft dia by 94 ft TT, and the stripper is 3 ft dia by 54 ft TT.

explanation may be correct for the enhancement of the volatility of isooctane (7.55) relative to that of toluene (8.91) in the presence of phenol (12.1) or aniline (11.5), both of which are commercially feasible additives. The data of Figure 13.23(a) do show that the volatility of isooctane is enhanced by the presence of phenol. The numbers in parentheses are the solubility parameters. In the case of acetone (9.8), chloroform (9.3), and methyl-isobutylketone (8.3),

the data of Figure 13.23(b) show that chloroform has the enhanced volatility, although its solubility parameter is closer to that of the solvent. A possible interpretation of the data is that association of the ketones as a consequence of their hydrogen bonding capabilities reduces the volatility of the acetone. Explanations of the effects of dissolved solids, as in Figures 13.23(c) and (d), are more obscure, although a substantial number of other cases also is known.

(a) Draw Tray Temperature T_{dt} as a Function of the Bubblepoint T_{bp} of the Stream: $T_{dt} = \begin{cases} \exp(0.0040T_{bp} + 4.404), & 200 \le T_{bp} \le 325^{\circ}F \\ \exp(0.002952T_{bp} + 4.744). & 325 \le T_{bp} \le 600^{\circ}F \end{cases}$





(c) Gap and Overlap between Sidestream Products in Terms of Reflux and Plates



Numbers on the streams are °F differences between the 50% points of the streams. Dashed lines are with stripping steam, full ones without [Packie, Trans. AIChE 37, 51 (1941)].

(d) Number of Trays between Drawoffs

Separation	Number of Trays			
Light naphtha to heavy naphtha	6 to 8			
Heavy naphtha to light distillate	6 to 8			
Light distillate to heavy distillate	4 to 6			
Heavy distillate to atmospheric gas oil	4to 6			
Flash zone to first draw tray	3 to 4			
Steam and reboiled stripping sections	4			

(e) Normal Stripping Steam Usage

Product	lb Steam/gal
Naphtha	0.2-0.5
Kerosene or diesel fuel	0.2-0.6
Gas oil	0.1-0.5
Neutral oils	0.4-0.9
Topped crude oil	0.4-1 .2
Residual cylinder stock	1.0 up

(f) Superficial Linear Velocities in Towers

Operation	Pressure (psia or mm)	Trav Spacing (in.)	Superficial Tower Velocity (ft/sec)
Topping	17 lb	2 2	2.6-3.3
Cracking	40 lb	2 2	1.5-2.2
Pressure dist. rerun	20 lb	2 2	2.8-3.7"
Solution rerun	25 lb	2 2	2.8-3.5
Pressed dist. rerun	25 lb	2 2	2.8-3.9"
Pressed dist. rerun	60 mm	24	6.0-9.0
Vacuum	30 mm	30	9.0-1 2.0
Vaccum	90 mm	24	5.0-8.0
Stabilizer	1601b	18	2.2-2.8
Nat. gaso, absorber	50 lb	14	1.0-1.3
Nat. gaso. absorber	400 lb	18	0.5-0.8

'Greatly dependent on quantity of steam.

(g) Pressure Drop 0.1-0.2 psi/tray

(h) Overflash into Tower Feed Zone is 2-5%



Figure 13.23. Examples of vapor-liquid equilibria in presence of solvents. (a) Mixture of i-octane and toluene in the presence of phenol. (b) Mixtures of chloroform and acetone in the presence of methylisobutylketone. The mole fraction of solvent is indicated. (c) Mixture of ethanol and water: (a) without additive; (b) with 10 g CaCl₂ in 100 mL of mix. (d) Mixture of acetone and methanol: (a) in 2.3M CaCl₂; (b) salt-free. (e) Effect of solvent concentration on the activity coefficients and relative volatility of an equimolal mixture of acetone and water (*Carlson and Stewart, in Weissbergers Technique of Organic Chemistry IV, Distillation,* 1965). (f) Relative volatilities in the presence of acetonitrile. Compositions of hydrocarbons in liquid phase on solvent-free basis: (1) 0.76 isopentane + 0.24 isoprene; (2) $0.24 iC_5 + 0.76 IP$; (3) $0.5 iC_5 + 0.5$ 2-methylbutene-2; (4) 0.25-0.76 2MB2 + 0.75-0.24 IP [Ogorodnikov et al., Zh. Prikl. Kh. 34, 1096–1102 (1961)].

Measurements of binary vapor-liquid equilibria can be expressed in terms of activity coefficients, and then correlated by the Wilson or other suitable equation. Data on all possible pairs of components can be combined to represent the vapor-liquid behavior of the complete mixture. For exploratory purposes, several rapid experimental techniques are applicable. For example, differential ebulliometry can obtain data for several systems in one laboratory day, from which infinite dilution activity coefficients can be calculated and then used to evaluate the parameters of correlating equations. Chromatography also is a well-developed rapid technique for vapor-liquid equilibrium measurement of extractive distillation systems. The low-boiling solvent is deposited on an inert carrier to serve as the adsorbent. The mathematics is known from which the relative volatility of a pair of substances can be calculated from the effluent trace of the elutriated stream. Some of the literature of these two techniques is cited by Walas (1985, pp. 216-217).

Some Available Data. A brief list of extractive distillation processes of actual or potential commercial value is in Table 13.7; the column of remarks explains why this mode of separation is adopted. The leading applications are to the separation of close-boiling aromatic, naphthenic, and aliphatic hydrocarbons and of olefins from diolefins such as butadiene and isoprene. Miscellaneous separations include propane from propylene with acrylonitrile as solvent (DuPont, U.S. Pat. 2,980,727) and ethanol from propanol with water as solvent [Fig. 13.24(b)].

Earlier explorations for appropriate solvents may have been conducted by the Edisonian technique of trying whatever was on the laboratory shelves. An extensive list of mixtures and the extractive distillation solvents that have been studied is in the book of Kogan (*Azeotropic and Extractive Distillation*, Leningrad, 1971, pp. 340-430, in Russian).

Some of the many solvents that have been examined for certain hydrocarbon separations are listed in Table 13.8; part (c) for n-butane and butene-2 separations includes data showing that addition of some water to the solvent enhances the selectivity. The diolefins butadiene and isoprene are available commercially as byproducts of cracking operations and are mixed with other close-boiling saturated, olefinic and acetylenic hydrocarbons, often as many as 10–20 different ones. The most widely **used** extractive

solvents are n-methylpyrrolidone (NMP), dimethylformamide (DMF), furfural and acetonitrile (ACN), usually with 10-20% water to improve selectivity, although at the expense of reduced solvent power and the consequent need for a greater proportion of solvent. A few of the many available data for these important separations appear in Figure 13.23(f) and Table 13.9. They show the effects of hydrocarbon proportions, the content of solvent, and the concentration of water in the solvent. Sufficient data are available for the major pairs of commercial mixtures to permit evaluation of parameters of the Wilson or other equations for activity coefficients in multicomponent mixtures, and thus to place the design of the equipment on a rational basis. Another distinction between possible additives is their solvent power. Table 13.10, for example, shows that diolefins are much more soluble in DMF than in ACN, and thus DMF circulation need be less.

Calculation Methods. An often satisfactory approximation is to take the mixture in the presence of the solvent to be a pseudobinary of the keys on a solvent-free basis, and to employ the McCabe-Thiele or other binary distillation method to find tray and reflux demands. Since the relative volatility varies with concentration of the solvent, different equilibrium curves are used for above and below the feed based on average loads in those zones. Figure 13.25 is of such a construction.

When data of activity coefficients of all pairs of components are known, including those with the solvent, any of the standard calculation procedures for multicomponent distillation, which include ternaries, may be used. Composition profiles found by tray-by-tray calculations in two cases appear in Figure 13.24.

To the number of trays found by approximate methods, a few trays are added above the solvent feed point in order to wash back any volatilized solvent. Nonvolatility is a desirable property, **but** most otherwise suitable solvents do have appreciable volatilities.

Extractive Distillation Recovery of Isoprene. A typical flowsketch and material balance of distillation and solvent recovery towers for extracting isoprene from a mixture of cracked products with aqueous acetonitrile appears in Figure 13.26. A description of the flowsheet of a complete plant is given in Example 2.10. In spite of the fact that several trays for washing by reflux are provided, some volatilization of solvent still occurs so that the complete plant

Additive	Mixture To Be Separated	Δ <i>T</i> (°C) <i>*</i>	Remarks
Aniline	n-heptane-methylcyclohexane	2.7	ideal mixture (α = 1.07)
	benzene-cycfohexane	0.7	azeotrope
	n-heptane-toluene	12.8	
			non-ideal mixtures;
Phenol	n-heptane-toluene	12.8	asymptotic approach of
	iso-octanetoluene	11.4	equilibrium curve to
	methylcyclohexane-toluene	9.5	diagonal
Ethyleneglycol monobutylether	methylethylketone-water	20.4	azetrope
Diethylether	ethanol-water	21.6	azeotrope
Higher ketones and alcohols	acetone-methanol	8.5	azeotrope
Higher esters and alcohols	ethylacetate-ethanol	1.3	azeotrope
Higher ketones and chloro compounds	acetone-chloroform	5.0	azetrope

 TABLE 13.7. Examples of Extractive Distillation Processes for the Separation of Ideal, Nonideal, and Azeotropic Systems

^a ΔT is the difference in atmospheric boiling points, °C.

TABLE 13.8. Relative Volatilities of Three Binary Systems and Their Enhancement in the Presence of Several Solvents

Solvent	Mole per cent in liquid phase	т. °С. (av.)	Av. rel. volatility, ¤ß	Improve- ment factor. $\alpha_{\rm B}/\alpha$	Ref. No.
Aniline.	92	139	1.52	1.42	1
	78	121	1.40	1.31	1
	70	110	1.27	1.19	2
	58	113	1.26	1.18	1
Furfural	79		1.35	1.26	1
Phenol	81		1.31	1.24	1
Nitrobenzene	82	10110.	1.31	1.24	1
Dichlorodiethyl ether.	81		1.28	1.20	1
Aminocyclohesane	76		1.16	1.08	1
Pyridine	70	_	1.4	1.31	2
Ethanol	70		1.3	1.21	2
n-Butanol	70	_	1.3	1.21	2
tert-Butanol.	70		1.25	1.17	2
Acetic acid	70		1.27	1.19	2
None	-	-	1.07	1.00	2

(a) n-Heptane/Methylcyclohexane with Relative Volatility of 1.07

Griswold, Andres, Van Berg, and Kasch, Znd. Eng. Chem., 38, 66 (1946).

¹ Fenske, Carlson, and Quiggle, Ind. Eng. Chem., 39, 1322 (1947).

(b) Cyclohexane/Benzene with Relative Volatility of 1.02

Solvent	Mole per cent in charge	t, °C.	Relative volatility, ¤ ₈	Improve- ment factor. α ₈ /α
Acetic acid	69.0	84	1.75	1.78
Methanol.	67.3	53	1.58	1.61
Ethanol	67.3	65	1.36	1.38
n-Propanol	70.5	79	1.26	1.28
Isopropanol	67.9	70	1.22	1.24
Dioxane	67.4	86	1.75	1.78
Chlorex (dichlorodiethyl ether).	67.5	105	2.31	2.36
Methyl Cellosolve	66.7	85	1.84	1.88
Cellosolve	67.5	95	1.58	1.61
Carbitol	66.8	87	1.99	2.03
Acetone.	66.3	55	2.03	2.07
Methyl ethyl ketone	65.1	72	1.78	1.81
Diacetone	67.3	89	1.82	1.85
Pyridine	66.9	93	1.83	1.86
Aniline.	66.8	93	2.11	2.16
Nitromethane	67.8	74	3.00	3.06
Nitrobenzene.	68.2	102	2.25	2.30
Acetonitrile	67.3	65	2.85	2.92
Furfural	67.1	79	3.10	3.16
Phenol.	66.8	92	2.01	2.05

Updike, Langdon, and Keyes, Trans. Am. Inst. Chem. Engrs., 41, 717 (1945).

TABLE 13.8—(continued)

(c) Butane/P-Butene with Relative Volatility of about	1.08° The	asterisks	denote	that
data are included for both dry and wet solvents				

Solvent	Vol/vol HC	Temperature, °F	$= \gamma c_{\bullet} / \gamma c_{\bullet}$
Hydroxyethylacetate	4	133	1.54
Methylsalicylate	2	156	1.46
Dimethylphthalate	2	142	1.41
Ethyl oxalate	2	154	1.38
Carbitolacetate	2	160	1.35
Diethyl carbonate	2	172	1.28
Amylacetate	2	180	1.21
* Acetonitrile	2	137	1.49
Butyronitrile	2	161	1.42
Acrylonitrile	2	156	1.23
Acetonyl acetone	2	141	1.43
Cyclohexanone	2	171	1.32
Acetophorone	2	145	1.31
Methylhexyl ketone	2	166	1.27
Methylamyl ketone	2	173	1.23
Methylisobutyl ketone	2	171	1.23
Methyldiisobutyl ketone	2	177	1.18
Nitromethane	1.8	134	1.00
Nitroethane	2	146	1.46
I-Chloro-I-nitropropane	2	155	1.46
Nitrobenzene	2	150	1.41
o-Nitrotoluene	2	155	1.38
o-Nitroanisole	2	130	1.30
n-Formylmorpholine	4.6	133	1.60
Morpholine	2	160	1.41
Pyridine	2	176	1.35
Quinoline	2	148	1.33
Picoline	2	188	1.29
Benzyl alcohol	3	144	1.48
* Phenol	2	138	1.47
Diacetone alcohol	2	146	1.32
Butyl alcohol	2	152	1.21
2-Ethyl butyl alcohol	2	161	1.20
o-Hexanol	2	159	1.18
tert-Butyl alcohol	2	154	1.16
Benzaldenyde	2	145	1.42
7 Furiurai	3	138	1.40
S,4-Diethioxybelizaidenyde	2	125	1.11
A nilino	2	103	1.09
a Chloroanilina	3	150	1.05
Methylaniline	2	132	1.44
o-Toluidine	2	148	1 38
Dimethyl aniline	2	169	1 37
n-Tributyl amine	2	176	1.09
Cellosolye	2	152	1.40
Dichloroethylether	2	152	1.39
Anisole	2	175	1.28
Butyl Cellosolve	2	163	1.24
Diethyl Cellosolve	2	179	1.23
Diethyl carbitol	2	173	1.23
n-Butylether	2	197	1.10
-			
Solvents with water			
* Furfural, 96 wt %	3.7	128	1.78
* Aniline, 96.5 wt %	4.4	132	1.77
Methylacetoacetate (90 vol %)	3.6	134	1.67
* Phenol (90 vol %)	2.5	133	1.66
Acetonylacetone (95 vol %)	3	128	1.58
Acetonitrile (90 vol %)	4	133	1.58
Benzyl alcohol (95 vol %)	2.5	133	1.51
o-Chlorophenol (90 vol %)	5.0	180	1.50

[Data from Hess, Narragon, and Coghlin, Chem. Eng. Prog. Symp. Ser. 2, 72-96 (195211.

TABLE 13.9. Relative Volatilities of C_4 and C_5 Hydrocarbons in Various Solvents (a) Volatilities of Butenes Relative to Butadiene at 40°C

Mole						Acet	onitrile		
Hydro-		DMF with 12%	l₂0		Dry			13% H₂O	
Solution	1-Sutene	trans-2-Butene	cis-2-Butene	1-Butene	trans-2-Butene	cis-2-Butene	1 - Butene	trans-2-Butene	cis-2-Butene
0.05	2.10	1.66	1.6		_	_	1.86	1.47	1.41
0.10	1.90	1.62	1.43	1.81	1.44	1.37	1.73	1.38	1.31
0.15	2.15	1.65	1.40	1.83	1.50	1.35	1.73	1.43	1.20

(Galata. Kofman, and Matveeva, Chem. Chem. Tech. (in Russian) 2, 242-255 (1962)].

(b) Volatilities at Infinite Dilution and 20°C Relative to Butadiene

			Solvent		
Compound	None	Furfural	DMF	NMP	ACN
n-Butane	0.88	3.0	3.04	3.84	3.41
lsobutylene	1.03	2.03	2.00	2.45	2.20
Butene-1	1.01	1.97	1.95	2.44	2.16
t-Butene-2	0.86	1.42	1.54	2.02	1.70
c-Butene-2	-	1.29	1.40	1.76	1.56

(Evans and Sarno, Shell Development Co.).

(c) Enhancement of Relative Volatilities of ${\rm C_4}$ and ${\rm C_5}$ Hydrocarbons

Class and Nama	No.	a/a, in Pre	sence of	Solvent
of Hydrocarbons	α _o	Furfural	DMF	ACN
Alkanes/alkenes				
n-Butane/I-butene	0.83	1.53	1.83	1.70
i-Butene/1-butene	1.14	1.52	1.83	1.70
i-Pentane/2-methyl-1-butene	1.08	1.66	1.85	-
n-Pentane/2-methyl-1-butene	0.84	1.61	1.80	-
Alkanes/dienes				
1-Butene/butadiene	1.04	1.57	1.91	1.69
Trans-2-butene/butadiene	0.83	1.44	1.85	1.71
cis-2-butene/butadiene	0.76	1.41	1.84	1.71
2-Methyl-2-butene/isoprene	0.87	1.45	1.84	_
Isoprene/3-methyl-1-butene	0.88	-	1.37	-

(Galata et al., loc cit.)

 TABLE 13.10. Solubilities (wt %) of Classes of C₄ and C₅

 Hydrocarbons in Various Solvents

DMF with		ACN	with	Furfural with		
Compounds	8% Water	12% Water	8% Water	12% Water	4% Water	
Saturated	7	5	14	9	7	
Olefins	18	13	3 2	2 2	18	
Diolefins	58	41	3 2	2 2	2 4	
Acetylenes	69	50	-	-	_	

[Galata, Kofman, and Matveeva, Chem. Chem. Tech. (in Russian) 2, 242-255 (1962)).

also has water wash columns on both hydrocarbon product streams. A further complication is that acetonitrile and water form an azeotrope containing about 69 mol % solvent. Excess water enters the process in the form of a solution to control polymerization of the unsaturates in the hotter parts of the towers and reboilers.

Two feasible methods for removal of as much water as desired from the azeotrope are depicted on Figure 13.27. The dual pressure process takes advantage of the fact that the azeotropic composition is shifted by change of pressure; operations at 100 and 760Torr result in the desired concentration of the mixture. In the other method, trichlorethylene serves as an entrainer for the water. A ternary azeotrope is formed that separates into two phases upon condensation. The aqueous layer is rejected, and the solvent layer is recycled to the tower. For economic reasons, some processing beyond that shown will be necessary since the aqueous layer contains some acetonitrile that is worth recovering or may be regarded as a pollutant.

AZEOTROPIC DISTILLATION

The objective of azeotropic distillation is the separation or concentration beyond the azeotropic point of mixtures with the aid of an entrainer to carry some of the components overhead in a column. An azeotrope is a constant boiling mixture with vapor and liquid phases of the same composition. A related class of systems is that of partially miscible liquids that also boil at constant temperature. The two phases exert their individual vapor pressures so that the boiling temperature and vapor composition remain constant over the full range of immiscibility, but the compositions of vapor and overall liquid phases are only accidentally the same. In most cases of immiscible liquids, the horizontal portion of an x-y diagram crosses the x = y line, for instance, the system of n-butanol and water of Example 13.5. The system of methylethylketone and water is one of the few known exceptions for which the immiscible boiling range does not cross the x = y line of Figure 13.28(b). Artificial systems can be constructed with this behavior. Thus Figures 13.28(c) and (d) are of diagrams synthesized with two different sets of parameters of the Margules equation for the activity coefficients; one of the x-y lines crosses the diagonal and the other does not. Figure 13.28(a) for acetone and water is representative of the most common kind of homogeneous azeotropic behavior.

The overhead stream of the distillation column may be a low-boiling binary azeotrope of one of the keys with the entrainer or more often a ternary azeotrope containing both keys. The latter kind of operation is feasible only if condensation results in two liquid phases, one of which contains the bulk of one of the key components and the other contains virtually all of the entrainer which can be returned to the column. Figure 13.29(a) is of such a flow scheme. When the separation resulting from the phase split is not complete, some further processing may make the operation technically as well as economically feasible.

Data of Azeotropes. The choice of azeotropic entrainer for a desired separation is much more restricted than that of solvents for extractive distillation, although many azeotropic data are known. The most extensive compilation is that of Ogorodnikov, Lesteva, and Kogan (Handbook of Azeotropic Mixtures (in Russian), 1971). It contains data of 21,069 systems, of which 1274 are ternary, 60 multicomponent, and the rest binary. Another compilation (Handbook of Chemistry and Physics, 60th ed., CRC Press, Boca Raton, FL, 1979) has data of 685 binary and 119 ternary azeotropes. Shorter lists with grouping according to the major substances also are available in Lange's Handbook of Chemistry

(12th ed., McGraw-Hill, New York, 1979). Data of some ternary systems are in Table 13.11.

Commercial Examples. The small but often undesirable contents of water dissolved in hydrocarbons may be removed by distillation. In drying benzene, for instance, the water is removed overhead in the azeotrope, and the residual benzene becomes dry enough for processing such as chlorination for which the presence of water is harmful. The benzene phase from the condenser is refluxed to the tower. Water can be removed from heavy liquids by addition of some light hydrocarbon which then is cooked out of the liquid as an azeotrope containing the water content of the original heavy liquid. Such a scheme also is applicable to the breaking of aqueous emulsions in crude oils from tar sands. After the water is removed



Figure 13.24. Composition profiles and flowsketches of two extractive distillation processes. (a) Separation of methylcyclohexane and toluene with phenol as solvent (*data calculated by Smith*, 1963). (b) Separation of aqueous ethanol and isopropanol, recovering 98% of the ethanol containing 0.2 mol % isopropanol, employing water as the solvent. Flow rates are in mols/hr (*data calculated by Robinson and Gilliland*, 1950).



x (solvent free)

Figure 13.25. Illustrating McCabe-Thiele construction of **pseudobi**nary extractive distillation with smaller relative volatility below the feed plate.

azeotropically, solids originally dissolved or entrained in the aqueous phase settle out readily from the dry hydrocarbon phase. Even in the evaporation of water from caustic, the addition of kerosene facilitates the removal of water by reducing the temperature to which the **pot** must he heated.



Ordinary rectification for the dehydration of acetic acid requires many trays if the losses of acid overhead are to be restricted, so that azeotropic processes are used exclusively. Among the entrainers that have been found effective are ethylene dichloride, n-propyl acetate, and n-butyl acetate. Water contents of these azeotropes are 8, 14, and 28.7 wt %, respectively. Accordingly, the n-butyl acetate is the most thermally efficient of these agents. The n-propyl acetate has been used in large installations, in the first stage as solvent for extraction of acetic acid and then as azeotropic entrainer to remove the accompanying



Figure 13.26. Flowsketch for the recovery of isoprene from a mixture of C_5s with aqueous acetonitrile. Flow quantities in lb/hr, pressures in psia, and temperatures in "F. Conditions are approximate. (Data of The C. W. Nofsinger Co.)

Figure 13.27. Separation of the azeotropic mixture of acetonitrile and water which contains approximately 69 mol % or 79.3 wt % of acetonitrile. (*Pratt*, Countercurrent Separation Processes, *Elsevier*, New York, 1967, pp. 194, 497). (a) A dual pressure process with the first column at 100 Torr and the second at 760 Torr. (b) Process employing trichlorethylene as entrainer which carries over the water in a ternary azeotrope that in turn separates into two phases upon condensation.



Figure 13.28. Vapor-liquid equilibria of some azeotropic and partially miscible liquids. (a) Effect of pressure on vapor-liquid equilibria of a typical homogeneous azeotropic mixture, acetone and water. (b) Uncommon behavior of the partially miscible system of methylethylketone and water whose two-phase boundary does not extend byond the y = x line. (c) x-y diagram of a partially miscible system represented by the Margules equation with the given parameters and vapor pressures $P_1^0 = 3$, $P_2^0 = 1$ atm; the broken line is not physically significant but is represented by the equation. (d) The same as (c) but with different values of the parameters; here the two-phase boundary extends beyond the y = x line.

water. Extractive distillation with a high boiling solvent that is immiscible with water upon condensation is technically feasible for acetic acid drying but is a more expensive process.

Ethanol forms an azeotrope containing 5 wt % water. In older installations, dissolved salts were employed to break the azeotrope. Typical data are in Figure 13.28(c). Several substances form ternary azeotropes with ethanol and water, including benzene, gasoline, and trichlorethylene. The first is not satisfactory because of slight decomposition under distillation conditions. A flowsketch of a

process employing benzene is in Figure 13.29(a). In a modernization of the benzene process (Raphael **Katzen** Associates, Cincinnati, OH), a high purity ethanol is made by controlling the distillation so that the lower 10 trays or so are free of benzene. Another entrainer, diethyl ether, has the desirable property of forming an azeotrope with water but not with ethanol. The water content of the azeotrope is small so that the operation is conducted at 8 atm to shift the composition to a higher value of 3% water. In small installations, drying with molecular sieves is a competitive



(b)

Figure W.29. Composition profiles and flowsketches of two azeotropic distillation processes (adapted by King, 1980). (a) Separation of ethanol and water with benzene as entrainer. Data of the composition profiles in the first column were calculated by Robinson and Gilliland, (1950); the flowsketch is after Zdonik and Woodfield (*in* Chemical Engineers Handbook, *McGraw-Hill, New York, 19.50, p. 652)*. (b) Separation of n-heptane and toluene with methylethylketone entrainer which is introduced in this case at two points in the column (*data calculated by Smith, 1963*).

process. Separations with membranes of both vapor and liquid phases, supercritical extraction with carbon dioxide and many other techniques have been proposed for removal of water from ethanol.

Formic acid can be dehydrated with propyl formate as entrainer. Small contents of formic acid and water in acetic acid can be entrained away with chloroform which forms binary azeotropes with water and formic acid hut no other azeotropes in this system.

Some hydrocarbon separations can be effected azeotropically. Figure 13.29(b) shows an operation with methylethylketone which

entrains n-heptane away from toluene. Hexane in turn is an effective entrainer for the purification of methylethylketone by distilling the latter away from certain oxide impurities that arise during the synthesis process.

Design. When the vapor-liquid equilibria are known, in the form of UNIQUAC parameters for instance, the calculation of azeotropic distillation may be accomplished with any of the standard multicomponent distillation procedures. The Naphthali-

Sandholm algorithm (Fig. 13.20) and the θ -method of Holland (1981) are satisfactory. Another tray-by-tray algorithm is illustrated for azeotropic distillation by Black, Golding, and Ditsler [*Adv. Chem.* Ser. 115, 64 (1972)]. A procedure coupling the tower, decanter, and stripper of Figure 13.29(a) is due to Prokopakis and Seider [*AIChE J. 29, 49* (1983)]. Two sets of composition profiles obtained by tray-by-tray calculations appear in Figures 13.29(a) and (b).

MOLECULAR DISTILLATION

This process is an evaporation that is conducted at such low pressures that the distance between the hot and condensing surfaces is less than the mean free path of the molecules. Each unit is a single stage, but several units in series are commonly employed. Molecular distillation is applied to thermally sensitive high molecular weight materials in the range of 250-1200 molecular

TABLE 13.11. Selected Ternary Aezotropic Systems at Atmospheric Pressure

(a) Systems with Water and Alcohols

	B.P . 76	0 mm.	% By weight		
	Other com- ponent	Azeo- trope	Water	Alcoho	Other com- ponent
A-Ethyl Alcohol (B.P. 78.3°)					
Ethyl acetate (6)	77.1	70.3	7.6	9.0	83.2
Diethyl formal (7)	67.6	73.2	12.1	18.4	69.5
Diethyl acetal (6)	103.6	77.6	11.4	27.6	61.0
Cyclohexane	69.6	62.1	7	17	76
Benzene (4)	80.2	64.9	7.4	18.5	74.1
Chloroform Carbon totrachlarida	61.2	55.5	3.5	4.0	92.5
Carbon tetrachioride	70.8	01.0	4.3	9.7	86.0
Ethylono chlorido	72.3	66 7	5 5	17	00 79
Ethylene Chloride	03.7	00.7	5	17	10
B-n-Propyl Alcohol (B.P. 97.2')					
n-Propyl formate (6)	80.9	70.8	13	5	8 2
<i>n</i> -Propyl acetate (6)	101.6	82.2	21.0	19.5	59.5
Di-n-propyl formal (7)	137.4	86.4	8.0	44.8	47.2
Di-n-propyl acetal (6)	147.7	87.6	27.4	51.6	21.0
Di-n-propyl ether (7)	91.0	74.8	11.7	20.2	68.1
Cyclonexane Berrone (4)	80.8	69.5	0.0	10.0	81.5
Benzene (4) Carbon totrachlarida	80.2	66.4	5	11	64
Diethyl ketone	102.2	81.2	20	20	60
biolity kolono					
C-Isopropyl Alcohol (B.P. 82.5")					
Cyclohexane	80.8	64.3	7.5	18.5	74.0
Benzene (4)	80.2	66.5	7.5	18.7	73.8
D-n-Butyl Alcohol (B.P. 117.6")					
n-Butyl formate (6)	106.6	83.6	21.3	10.0	68.7
n-Butyl acetate (6)	126.2	89.4	37.3	27.4	35.3
Di-n-butyl ether (7)	141.9	91	29.3	42.9	27.7
E-Isobutyl Alcohol (B.P. 108.0°)					
Isobutyl formate (6)	94.4	80.2	17.3	6.7	76.0
Isobutyl acetate	117.2	86.8	30.4	23.1	46.5
R-tert-Rutul Alcohol (B.P. 82.6')					
Benzene (4)	80.2	67.3	8.1	21.4	70.5
Carbon tetrachloride (9)	76.8	64.7	3.1	11.9	85.0
$Q = 4 \text{mm} \left\{ (1 + 1 + 1) \right\} = 0$					
G-n-Amyl Alcohol (B.P. 137.8°)		04.4		24.2	
n-Amyl formate (6)	131.0	91.4	37.6	21.2	41.2
n-Amyl acetate (0)	148.8	94.8	56.2	33.3	10.5
H-Isoamyl Alcohol (B.P. 131.4')					
Isoamyl formate (6)	124.2	89.8	32.4	19.6	48.0
isoamyl acetate (6)	142.0	93.6	44.8	31.2	24.0
I-Allyl Alcohol (B.P. 97. 0°)					
n - Hexane	69.0	59.7	5	5	90
Cyclohexane	80.8	66.2	8	11	81
Benzene	80.2	68.2	8.6	9.2	82.2
Carbon tetrachiorida	/6.8	65.2	5	11	64

(Lange, Handbook of Chemistry, McGraw-Hill, New York, 1979).

TABLE 13.11—(continued)

(b) Other Systems

Component A Mole $\% A \equiv 100 - B$ (C)	Components B and C	Mole % B and C	Temp.,
Water	Carbon tetrachloride Ethanol	57.6 23.0	61.6 2 phase
	Trichloroethylene Ethanol	38.4 41.2	67.23 2 phase
	Trichloroethylene Allyl alcohol	9::	71.4 2 phase
	Trichloroethylene Propyl alcohol (n)	51.1 16.6	71.55 2 phase
	Ethanol Ethyl acetate	12.4 60. I	70.3
	Ethanol Benzene	22.8 53.9	64.66
	Allyl alcohol Benzene	9.5 62.2	66.3
	Propyl alcohol (n) Benzene	8.9 62.8	66.48
Carbon disulfide.	Methanol Ethyl bromide	24.1 35.4	33.92
Methyl format	Ethyl bromide Isopentane	23.0 31.0	16.93
	Ethyl ether Pentane (n)	7.2 46.2	20.4
Ropyl lactate (n)	Phenetol Menthene	35.2 34.1	163.0

weights, such as oils, fats, waxes, essential oils and scents, vitamins and hormone concentrates, and to the deodorization of high molecular weight materials.

Operating pressures are in the range of 1 m Torr. For example, the mean free paths of normal triglycerides of 800 molecular weight are

P (m Torr)	Path (mm)
8	7
3	2 5
1	50

The theoretical Langmuir equation for the rate of evaporation is

$$w' = 2100P^0 \sqrt{M/T} \text{ kg/m}^2 \text{ hr}$$
 (13.223)

with the vapor pressure P^0 in Torr, the temperature in K, and with M as the molecular weight. Industrial apparatus may have 80-90% of these rates because of inefficiencies. Some numerical values at 120°C are:

Compound	м	P ^o (Torr)	w' (kg/m² hr)
Stearic acid	284	35.0	1.87
Cholesterin	387	0.5	2.02
Tristearin	891	0.0001	2.74

From Langmuir's equation it is clear that it is possible to separate substances of the same vapor pressure but different molecular weights by molecular distillation.

Apparatus and Operating Conditions. The main kinds of commercial units are illustrated in Figure 13.30. In the falling film type, the material flows by gravity as a thin film on a vertical heated cylinder, evaporates there, and is condensed on a concentric cooled surface. Diameters range from 2 to 50 cm, heights 2 to 10 m, and feed rates from 1 to 60 L/hr. In order to prevent channelling, the surface of the evaporator is made rough or other means are

employed. The cross section of a wiped film commercial still is shown in Figure 13.30(b). Contact times in commercial apparatus may be as low as 0.001 sec.

In the centrifugal still, the material that is charged to the bottom creeps up the heated, rotating conical surface as a thin film, is evaporated, and then condensed and discharged. The film thickness is 0.05-0.1 mm. Rotors are up to about 1.5 m dia and turn at 400-500 rpm. Evaporating areas are up to 4.5 m^2 per unit, feed rates range from 200 to 700 L/hr, and distillates range from 2 to 400 L/hr, depending on the service. From three to seven stills in series are used for multiple redistillation of some products. Two stills 1.5 m dia can process a tank car of oil in 24 hr. A typical pumping train for a large still may comprise a three-stage steam ejector, two oil boosters and a diffusion pump, of capacity 1000–5000L/sec, next to the still. Equivalent mechanical pumps may be employed instead of the ejectors, depending on the economic requirements.

The evaporator of Figure 13.30(d) is for service intermediate between those of ordinary film evaporators and molecular stills, with greater clearances and higher operating pressures than in the latter equipment. The rotating action permits handling much more viscous materials than possible in film evaporators.

13.12. TRAY TOWERS

Contacting of vapor and liquid phases on trays is either in countercurrent flow or with cross flow of liquid against vapor flow upward. The spacing of trays is determined partly by the necessity of limiting carryover of entrainment from one tray to another, and is thus related to the vapor velocity and the diameter of the vessel. For reasons of accessibility of trays to periodic servicing, however, their spacing commonly is 20-24in. Then workmen can go up or down the tower through removable sections of the trays and have enough room to work in. For the same reason, tray diameters are restricted to a minimum of 30in. When a smaller size is adequate, cartridge trays that can be lifted out of the vessel as a group, as in Figure 13.31(b), or packed towers are adopted. A data sheet for recording key data of a tray tower is in Table 13.12. The tedious calculations of many mechanical details of tray construction usually are relegated to computers.

COUNTERCURRENT TRAYS

The three main kinds of trays with countercurrent flow of liquid and gas are:

- Dualflow, with round holes in the 1/8–1/2 in. range, extensively tested by Fractionation Research Inc. (FRI).
- Turbogrid, with slots 1/4-1/2 in. wide, developed by Shell Development Co.
- Ripple trays, made of perforated corrugated sheets, with vapor flow predominantly through the peaks and liquid through the valleys, developed by Stone and Webster.

Although some of the vapor and liquid flows through the openings are, continuous, the bulk of the flows pass alternately with a surging action. The absence of downcomers means a greater bubbling area and consequently a greater vapor handling capacity, and also allows a close spacing to be used, as little as 9 in. in some applications. The action in such cases approaches those of towers filled with structured packings. Their turndown ratio is low, that is, the liquid drains completely off the tray at lowered vapor rates. Consequently, countercurrent trays have never found widespread use.



Figure 13.30. Molecular distillation and related kinds of equipment. (a) Principle of the operation of the falling film still (Chemical Engineers Handbook, *McGraw-Hill, New* York, 1973). (b) Thin-layer evaporator with rigid wiper blades (*Luwa* Co., *Switzerland*). (c) The Liprotherm rotating thin film evaporator, for performance intermediate to those of film evaporators and molecular stills (*Sibtec Co., Stockholm*). (d) Centrifugal molecular still [*Hickman*, Ind. Eng. Chem. 39, 686 (1947)].



Figure 13.31. Assembled sieve tray towers. (a) Flowsketch of a sieve tray tower (*Treybal*, 1980). (b) Cartridge type sieve tray tower in small diameters (*Pfaudler* Co.).

On crossflow trays, the path of liquid is horizontal from downcomer to weir and in contact with vapor dispersed through openings in the tray floor. Such flows are illustrated in Figures 13.31 and 13.32. Depending on the rate and on the diameter, the liquid flow may be single, double, or four-pass. A common rule for dividing up the flow path is a restriction of the liquid rate to a maximum of about 8 gpm/in. of weir length. Usually towers 5 ft dia and less are made single pass. Since efficiency falls off as the flow path is shortened, a maximum of two passes sometimes is specified, in which cash flow rates may approach 20 gpm/in. of weir.

The main kinds of cross flow trays with downcomers in use are sieve, valve, and bubblecap.

SIEVE TRAYS

A liquid level is maintained with an overflow weir while the vapor comes up through the perforated floor at sufficient velocity to keep most of the liquid from weeping through. Hole sizes may range from 1/8 to 1 in., but are mostly 1/4-1/2 in. Hole area as a percentage of the active cross section is 5-15%, commonly 10%. The precise choice of these measurements is based on considerations of pressure drop, entrainment, weeping, and mass transfer efficiency. The range of conditions over which tray operation is satisfactory and the kinds of malfunctions that can occur are indicated roughly in Figure 13.33(a) and the behavior is shown schematically on Figure 13.32(e).

The required tower diameter depends primarily on the vapor rate and density and the tray spacing, with a possibly overriding restriction of accommodating sufficient weir length to keep the gpm/in. of weir below about 8. Figure 13.33(b) is a correlation for the flooding velocity. Allowable velocity usually is taken as 80% of the flooding value. Corrections are indicated with the figure for the fractional hole area other than 10% and for surface tension other than 20 dyn/cm. Moreover, the correction for the kind of operation given with Figure 13.34 for valve trays is applicable to sieve trays.

Weir heights of 2 in. are fairly standard and weir lengths about 75% of the tray diameter. For normal conditions downcomers are



(International Critical Tables, McGraw-Hill, New York, 1929).

sized so that the depth of liquid in them is less than 50% and the residence time more than 3 sec. For foaming and foam-stable systems, the residence time may be two to three times this value. The topic of tray efficiency is covered in detail in Section 13.6, but here it can be stated that they are 80–90% in the vicinity of $F = u_v \sqrt{\rho_v} = 1.0$ (ft/sec)(lb/cuft)^{1/2} for mixtures similar to water with alcohols and to C_{C-C_a} hydrocarbons.

A detailed design of a tray includes specification of these items:

- 1. Hole dia, area, pitch and pattern.
- 2. Blanking of holes for less than eventual load.
- 3. Downcomer type, size, clearance, and weir height.
- 4. Tray thickness and material.
- 5. Pressure drop.
- 6. Turndown ratio before weeping begins.
- 7. Liquid gradient.

Correlations for checking all of these specifications are known. An example is worked out by Fair (in Smith, 1963, Chap. 15). The basis is holes 3/16 in. dia, fractional open area of 0.10, weir height of 2 in. and tray spacing 24 in.

The correlation of Figure 13.33 has no provision for multipass

liquid flow. Corrections could be made by analogy with the valve tray correlation, as suggested at the close of Example 13.15.

VALVE TRAYS

The openings in valve trays are covered with liftable caps that adjust themselves to vapor flow. Illustrations of two kinds of valves are in Figure 13.32(b). The caps rest about 0.1 in. above the floor and rise to a maximum clearance of 0.32 in. The commonest hole diameter is 1.5 in. but sizes to 6 in. are available. Spacing of the standard diameter is 3-6 in. With 3 in. spacing, the number of valves is 12-14/sqft of free area. Some of the tray cross section is taken up by the downcomer, by supports, and by some of the central manway structure.

In spite of their apparent complexity of construction in comparison with sieve trays, they usually are less expensive than sieve trays because of their larger holes and thicker plates which need less support. They are more subject to fouling and defer to sieves for such services.

Tray diameters may be approximated with Figure 13.34 which is for "normal" systems, 24 in. tray spacing, and 80% of flooding. For other tray spacings, corrections may be approximated with the

430 DISTILLATION AND GAS ABSORPTION



Figure 13.32. Internals and mode of action of trays in tray towers. (a) Some kinds of bubblecaps (*Glitsch*). (b) Two kinds of valves for trays. (c) Vapor directing slot on a Linde sieve tray [Jones and Jones, Chem. Eng. Prog. 71, 66 (1975)]. (d) Vapor flow through a bubblecap. (e) Sieve tray phenomena and pressure relations; h_h is the head in the downcomer, h_i is the equivalent head of clear liquid on the tray, h_f is the visible height of froth on the tray, and h_i is the pressure drop across the tray (*Bolles*, in *Smith*, 1963). (f) Assembly of and action of vapor and liquid on a bubblecap tray.



Figure 13.33. Operating ranges of malfunctions and flooding velocity correlation of sieve trays. (a) Performance of a typical sieve tray, showing ranges of weeping, dumping, entrainment, and flooding. (b) Correlation of flooding at various tray spacings. For normal operation, take 80% of the flooding rate as a design condition. To correct for surface tension (dyn/cm), multiply the ordinate by ($\sigma/20$)^{0.2}. To correct for other than 10% hole area, multiply the ordinate by 0.9 for 8% and by 0.8 for 6% [after Fair and Matthews, Pet. Refin. **37**(4), 153 (1958)].

sieve tray correlation of Figure 13.33(c). Factors for correcting the allowable volumetric rate for various degrees of foaming are given with the figure.

Formulas and procedures for calculation of detailed tray specifications are presented, for example, by Glitsch Inc. (Bulletin 4900, Ballast Tray Design Manual, Dallas, TX, 1974), and illustrated with .a completely solved numerical problem.

BUBBLECAP TRAYS

Bubblecap assemblies serve to disperse the vapor on the tray and to maintain a minimum level of liquid. A few of the many kinds that have been used are in Figure 13.32, together with illustrations of their mode of action and assembly on a tray. The most used kinds are 4 or 6in. dia round caps. Because of their greater cost and

EXAMPLE 13.15

 $W_{\rm H} = 271,500 \ \text{lb/hr}$ of vapor,

Comparison of Diameters of Sieve, Valve, and Bubblecap Trays for the Same Service

A C_3 splitter has 24 in. tray spacing and will operate at 80% of flooding. These data are applicable:

 $\begin{array}{l} Q_v = 27.52 \ \text{cfs} \ \text{of vapor}, \\ W_L = 259,100 \ \text{lb/hr} \ \text{of liquid}, \\ Q_L = 1100 \ \text{gpm}, \\ \rho_V = 2.75 \ \text{lb/cuft}, \\ \rho_L = 29.3 \ \text{lb/cuft}. \\ \hline \text{Sieve tray: Use Figure 13.33(b):} \\ abscissa = (259,100/271,500)\sqrt{2.75/29.3} = 0.2924, \\ \text{ordinate, C} = 0.24, \\ u_G = C\sqrt{(\rho_L - \rho_V)/\rho_V} = 0.24\sqrt{29.3/2.75 - 1} = 0.746 \ \text{fps} \end{array}$

Allowable velocity at 80% of flooding,

 $u_G = 0.8(0.746) = 0.597$ fps, ∴ $D = \sqrt{Q_v/(\pi/4)u_G} = \sqrt{27.52/(\pi/4)(0.597)} = 7.67$ ft.

Valve tray: Use Figure 13.34:

(cfs) $\sqrt{\rho_V/(\rho_L - \rho_V)}$ = 27.52 $\sqrt{2.75/(29.3 - 2.75)}$ = 8.86, ∴ $D = \begin{cases} 9.4 \text{ ft, one pass,} \\ 7.6 \text{ ft, two passes.} \end{cases}$

Bubblecap tray: Use Eq. (13.224):

K = 4.2 for 24 in. tray spacing, ∴ D = 0.0956[271,500/4.2 $\sqrt{29.3(2.75)}$]^{1/2} = 8.11 ft.

The correlations for sieve and bubblecap trays have no provision for multipass flow of liquid. Their basic data may have been obtained **on** smaller towers with liquid flow equivalent to two-pass arrangement in towers 8 ft dia. The sieve tray correlation should be adapted to multipass flow by comparison with results obtained by the valve tray correlation in specific cases.



Service	Factor		
Nonfoaming, regular systems	1.00		
Fluorine systems, e.g., BF,, Freon	0.90		
Moderate foaming, e.g., oil absorbers, amine and			
glycol regenerators	0.85		
Heavy foaming, e.g., amine and glycol absorbers	0.73		
Severe foaming, e.g., MEK units	0.60		
Foam-stable systems, e.g., caustic regenerators	0.30-0.60		

Figure 13.34. Chart for finding the diameters of valve trays. Basis of 24in. tray spacing and 80% of flood for nonfoaming services. Use Figure 13.32(b) for approximate adjustment to other tray spacings, and divide the $V_{\text{load}} = V \sqrt{\frac{\rho_V}{\rho_L - \rho_V}}$ by the given "system factor" for other services (*Glitsch Inc., Bulletin 4900, Dallas, TX, 1974*).

problems with hydraulic gradient, bubblecap trays are rarely installed nowadays, having lost out since about 1950 to the other two kinds. Since they do have a positive liquid seal and will not run dry, they are used sometimes in low liquid flow rate situations such as crude vacuum towers, but even there they have lost out largely to structured tower packings which have much lower pressure drop.

The allowable vapor velocity and the corresponding tray diameter are represented by the work of Souders and Brown, which is cited in standard textbooks, for example Treybal (1980). Its equivalent is the "Jersey Critical" formula,

$$D = 0.0956 (W_v / K \sqrt{\rho_L \rho_v})^{1/2}, \text{ ft}$$
(13.224)

with the factor K dependent on the tray spacing as follows:

Tray spa	cing (in.)	18	24	30	30+
κ.	• • •	3.4	4.2	4.7	5.0

Here

 W_{ν} = vapor flow rate (lb/hr), ρ_{ν} = vapor density (lb/cuft), ρ_L = liquid density (lb/cuft).

Example 13.15 compares diameters of sieve, valve, and

bubblecap trays calculated with the relations cited in this section. All of these relations presumably are based on limiting the amount of entrainment to a level that does not affect efficiency appreciably. Accordingly, the differences in diameters found in that example are due less, perhaps, to differences in performances of the different kinds of trays than to the particular data on which the correlations are based.

A factor that is of concern with bubblecap trays is the development of a liquid gradient from inlet to outlet which results in corresponding variation in vapor flow across the cross section and usually to degradation of the efficiency. With other kinds of trays this effect rarely is serious. Data and procedures for analysis of this behavior are summarized by Bolles (in Smith, 1963, Chap. 14). There also are formulas and a numerical example of the design of all features of bubblecap trays. Although, as mentioned, new installations of such trays are infrequent, many older ones still are in operation and may need to be studied for changed conditions.

13.13. PACKED TOWERS

In comparison with tray towers, packed towers are suited to small diameters (24in. or less), whenever low pressure is desirable, whenever low holdup is necessary, and whenever plastic or ceramic construction is required. Applications unfavorable to packings are large diameter towers, especially those with low liquid and high vapor rates, because of problems with liquid distribution, and whenever high turndown is required. In large towers, random packing may cost more than twice as much as sieve or valve trays.

Depth of packing without intermediate supports is limited by its deformability; metal construction is limited to depths of 20-25 ft, and plastic to 10–15 ft. Intermediate supports and liquid redistributors are supplied for deeper beds and at sidestream withdrawal or feed points. Liquid redistributors usually are needed every $2\frac{1}{2}$ -3 tower diameters for Raschig rings and every 5-10 diameters for pall rings, but at least every 20 ft.

The various kinds of internals of packed towers are represented in Figure 13.35 whose individual parts may be described **one**by-one:

- (a) is an example column showing the inlet and outlet connections and some of the kinds of internals in place.
- (b) is a combination packing support and redistributor that can also serve as a sump for withdrawal of liquid from the tower.
- (c) is a trough-type distributor that is suitable for liquid rates in excess of 2 gpm/sqft in towers two feet and more in diameter. They can be made in ceramics or plastics.
- (d) is an example of a perforated pipe distributor which is available in a variety of shapes, and is the most efficient type over a wide range of liquid rates; in large towers and where distribution is especially critical, they are fitted with nozzles instead of perforations.
- (e) is a redistribution device, the rosette, that provides adequate redistribution in small diameter towers; it diverts the liquid away from the wall towards which it tends to go.
- (f) is a holddown plate to keep low density packings in place and to prevent fragile packings such as those made of carbon, for instance, from disintegrating because of mechanical disturbances at the top of the bed.

KINDS OF PACKINGS

The broad classes of packings for vapor-liquid contacting are either random or structured. The former are small, hollow structures with large surface per unit volume that are loaded at random into the vessel. Structured packings may be layers of large rings or grids, but are most commonly made of expanded metal or woven wire screen that are stacked in layers or as spiral windings.

The first of the widely used random packings were Raschig rings which are hollow cylinders of ceramics, plastics, or metal. They were an economical replacement for the crushed rock often used then. Because of their simplicity and their early introduction, Raschig rings have been investigated thoroughly and many data of their performance have been obtained which are still, useful, for example, in defining the lower limits of mass transfer efficiency that can be realized with improved packings.

Several kinds of rings are shown in Figure 13.36. They are being made in a variety of internal structure and surface modifications. Pall rings in metal and plastics are perhaps the most widely used packings. One brand, "Hy-Pak," has corrugated walls and more intrusions than the standard designs shown in the figure. Cascade minirings, with height less than the diameter, appear to have improved efficiency in comparison with some other pall rings. Saddles are more efficient because of greater surface and improved hydrodynamics. In plastic construction, Figure 13.36(h), they are made with a variety of holes and protrusions to enlarge the specific surface. When ceramic construction is necessary, saddles are the preferred packings. A survey of efficiencies of packed beds is in Table 13.13.

Whenever possible, the ratio of tower and packing diameters should exceed 15. As a rough guide, 1 in. packing is used for gas rates of about 500 cfm and 2 in. for gas rates of 2000 cfm or more.

Structured packings are employed particularly in vacuum service where pressure drops must be kept low. Because of their open structure and large specific surfaces, their mass transfer efficiency is high when proper distribution of liquid over the cross section can be maintained. Table 13.14 is a comparison of various features of five commercial makes of structured packings. The **HIGEE** centrifugal fractionator of Figure 13.14 employs structured packing in the form of perforated metal.

Ultimately, the choice of packing is based on pressure drop and mass transfer efficiency. Since packings of individual manufacturers differ in detail, the manufacturers pressure drop data should be used. A few such data are in Figures 13.37 and 13.38. Mass transfer efficiency is discussed in the next section.

FLOODING AND ALLOWABLE LOADS

The main operating limitation of the operation of a packed bed is the onset of flooding. Then the interstices tend to fill with liquid, the gas becomes unable to flow smoothly, and the pressure drop begins to rise sharply. The classic correlation of the flood point is due to Sherwood and Lobo et al. It is shown in Figure 13.39. Clearly, there is much scatter and many more recent kinds of packings are not covered. Nevertheless, it is fairly standard practice to design for a flow rate of 70–80% of that given by the correlation. In case the liquid is a foaming type, the factor is 40% of the flooding rate, or some means of eliminating the foam is found.

The correlation of Eckert (Fig. 13.37) combines a pressure drop relation and safe flow rates insofar as staying away from the flooding point is concerned. A flooding line corresponds to pressure drops in excess of 2 in. water/ft. In use, a pressure drop is selected, and the correlation is applied to find the corresponding mass velocity G from which the tower diameter then is calculated. Another correlation recommended by a manufacturer of packings appears in Figure 13.40. Example 13.16 compares these correlations for a specific case; they do not compare any more closely than could be expected from the scatter of flooding data.







(c)



(d)





Figure 13.35. Packed column and internals. (a) Example packed column with a variety of internals [*Chen*, Chem. Eng. 40, (5 *Mar. 1984*)]. (b) Packing support and redistributor assembly. (c) Trough-type liquid distributor. (d) Perforated pipe distributor. (e) Rosette redistributor for small towers. (f) Hold-down plate, particularly for low density packing.











(a)

(b)

(c)

(d)

(e)





(g)



(h)

(i)



(j)

(f)

(k)

(m)



(I)



(n)





(o)



(p)

Figure 13.36. Some kinds of tower packings: (a) Raschig ring; (b) partition or Lessing ring; (c) double spiral ring; (d) metal pall ring; (e) plastic pall ring; (f) ceramic Berl saddle (*Maurice A. Knight* Co.); (g) ceramic intalox saddle (Norton Co.); (h) plastic intalox saddle (Norton Co.); (i) metal intalox saddle (*Norton* Co.); (j) Tellerette (*Chem-Pro* Co.); (k) plastic tripak (Polymer Piping and Metals Co.); (m) wood grid; (n) section through expanded metal packing; (o) sections of expanded metal packings placed alternatively at right angles (*Denholme* Co.); (p) GEM structured packing (*Glitsch Inc.*).

TABLE	13.13.	Survey	of	Efficiencies	of	Packed	Beds
-------	--------	--------	----	--------------	----	--------	------

		Packing		Bed			System
	Dia.	·3	Size,	Depth	HETP,	HTU,	press.,
System	m	Туре	m	m	m	m	kPa
Hydrocarbona							
Absorber	091	Pall rings	005	70	0 85		5.964
L 0 top fractionator	091	Pall rings	005	52	0 76		1.083
L 0 bottom fractionator	1 22	Pall rings	0 05	52	0 85		1.083
Desthanizer top	046	Pall rings	0038	61	088		2,069
	0.59	Pall 1005	0038	5 5 4 88	0.98		2.069
Depropanizer bottom	0 59	Pall rings	0 038	7 32	0 73	-	1,862
Debutanizer top	0 50	Pall IIngs	0 038	366	0 73		621
Debutanizer bottom	050	Pall rings	0038	5 49	061		621
Pentane-isopentane	046	Pall rings	0025	2 74	0.46	-	101
Light/heavy naphtha	0 38	Pall rings	0025	305	062	054	13
	038	Intalox	0025	305	0 76	0.61	13
les satessitslusss	0.38	Raschig rings	0025	305	0 71	0 52	13
iso-octane/toluene	0.38		0025	305	043	045	13
Gas plant absorber	122	Pall rings	0025	70	0.88	0 31	6.206
2.2.4-trimethyl-pentane/							
methylcyclo-hexane	091	Stedman		76	0 88		101
	335	Stedman		2 1	0 13	-	101
4							
Hydrocarbons/water							
Acetone	0 36	Intalox Doll report	0025	396	046		101
Mothanol (house)	046	raii iliys	0025	8 38	037	076	101
Methanol	041	Pall (IDOS	0038	4 2 7	-	0.52	101
liotianoi	0 30	Intalox	0025	823	046	_	101
Isopropanol	0 53	Plastic pall	0038	4 88	-	084	101
-	0 33	Intalox .	0025	6 40		0 76	101
	046	Intalox	0025	335	048		101
Ethylene glycol	1 07	Pall rings	0038	4 88	U 91		31
Propylene glycol	025	Intalox	0013	1 83	0.61	0 86	101
Formula and	0.51	Intalox Poll (10/06	0038	549	0 76	_	101
Acetone (absorption)	061	Pail 11195	0.036	1067	0.0	0.4.6	101
Benzolchloride/	001	II ILLIOX	0030	5 45		040	
benzene/steam	061	Pall rings	0025	5 18		107	101
Tall Oil/steam	3 66	Intalox	0 05	1046	0 76		101
Methylisobutyl							
ketone/steam	107	Intalox	0038	853	1 22	-	101
Acetone	0 38	Intalox	005	290	053	047	101
	0 38	Pall rings	0038	290	046	055	101
	0 38	Pail Intys	0025	290	044	0 34	101
	038	Berl	0025	290	0 52	0.34	101
	0.38	Cer Raschin	0025	290	105	0 36	101
	038	Raschig rings	0025	2 90	0 52	036	101
	0 38	Pall rings	0016	290	040	032	101
Methanol	038	Pall rings	0025	290	066	067	101
Polar hydrocarbon8							
Methyl furan/							
methyl tetra-hydrofuran	061	Intalox	0038	14 63	053		101
Benzoic acid/							
toluene	061	Intalox	0038	6 40	046		101
Methone (5, 5 dimethyl							
1, 3 cyclonexaneolone).							
batch	0 56	Pall rings	0038	9 75	0 49		101
Monochloro							
acetic anhydride	025	Intalox	0025	2 74		0.86	8.0
Tar acid distillation							
(batch)	046	Pall rings	0038	9 14	0 49	-	13
Cresols (batch)	046	Pall rings	0038	914	085		13
Benzene/							
monochloro-benzene	038	Intalox	0038	2 90	1 80	0 52	101
Methylethylketone/	0 38	Intalox	0025	2 90	1 13	0 /6	101
toluene	038	Pall rings	0025	2 00	0.35	0 29	101
	038	Raschig fings	0025	2 90	0 30	0 31	101
	0 38	Intalox	0025	2 90	0 23	027	101
	038	Berl	0025	290	0 31	031	101
	038	Cer Raschig	0025	2 90	046	0 30	101
	0 38	Pall rings	0025	290	040	U 28	101
	0 38	Intelox	0025	290	0 35	030	101
	0.38	Rod	0025	2 50	0 29	0 20	101
	0.38	Cer Raschla	0025	290	031	027	101
Phenol/ortho-creosol	046	Pall rings	0038	9 14	0 49	-	13
Fatty acid	0 76	Pall rings	0038	12 19	085		4 94
Benzene/mono-		-					
chloro-benzene	183	Intalox	0038	9 75	1 07		13
DMPG/							
DMPC CIESOIS/	0.42		0.000	0.44	0.42	_	0.07
	0.46	rai iniya	0038	9 14	U 49		267
CH-CI-/							
CHCI,/							
CCI	048	Intalox	0025	20 73	046		101
Methylene/							
light ends	046	Intalox	0025	1341	0 46	-	101
Methylenel	0.01	Inteles	0000	10 70	0.4.0		101
Chloroform/	0 64	Intalox	0038	13 72	040	_	101
product	056	Intalox	0038	27 43	046		101
F	200		5000				.51

[Eckert, Chem. Eng. Prog. 59(5) 76 (1963)].

TABLE 13.14. Comparison of Structured Tower Packings

,	Goodloe packing	Hyperfil packing	Koch-Sulzer packing	Neo-Kloss packing	Leva film trays
General information:					
Туре	Knitted multifilament	Knitted multifilament	Corrugated woven-wire fabric	Rolled screen with spacers	Multiple unsealed downcomer trays on
Approximate number of units 12 in and larger sold through 1975	610	9 0	500	43	120
Largest diameter sold to date Materials in which available	5 ft,8 in	5 ft	11 ft	6 ft	14 ft , 6 in
Process and system considentions:					
Minimum head pressure, torr Liquid considerations:			0.5	0. 5'	5
Minimum rates, gal/(min)(ft ²)	0.016	0.1'	0.08	0.1'	0.05
Maximum rate, gal/(min)(ft ²)' Maximum viscosity, cP	>4.9 200	4. 7'	>8	3. 8'	$^{5}_{>100}$
Holdup, fraction of total volume, typical	0.07-0. 12	0.1	0.04	0.03	
Sensitivity to uneven initial liquid distribution	Moderate	Moderate	Fairly low	High ^d	Moderate
Vapor F factor, based on internal cross- sectional area of shell typical	0. 5- 1. 5	1.0	1.8	2. 5-3. 0	
Maximum	1.7	1.4	3.3	4.0	2.0
Minimum	low	0.14		0.16	0. 25
HETS, in:					
Range	$3^{1}/_{2} - 8^{1}/_{2}$	31/2-9	4-10	4-18	12-24
Typical	5	5		8	18
Pressure drop, mmHg per foot of packed height Pressure drop per theoretical stage			See Fig. 3	See Fig. 2	See Fig;. 4
Fouling considerations:	v		Madamaalish		
Sensitive to particulate solids?	Yes	Y e s	Moderately	N 0	N O
Sensitive to fouling by tarry substances?	Yes	Yes	Moderately "	Yes	N O
Machanical considerations:	res	res	Moderately	res	IN U
Is the device furnished as a package including	Optional	Optional	Optional	Y e s	Optional
Can it be installed through shell manholes?	Voc'	Vac	Ves	NO	N O
Can it be installed in an existing shell with only minor modifications?	Yes'	Y e s	Y e s	110	Yes ^k
Test facilities:					
Are pilot test facilities available?	Y e s	Y e s	Y e s	Y e s	Y e s

"Any metal capable of being drawn into wire.

^bAny metal which can be fabricated into the required shapes.

"These liquid rates are generally those claimed by the manufacturers. Very low liquid loadings [below 0.2 gal/(min)(ft*)] always require special attention to the design of the liquid-distribution system.

Neo-Kloss packing requires highly precise initial liquid 'distribution because liquid cannot spread from one layer of screen to the next. Care is needed in the design of the distribution system (provided by vendor), in its installation, and in prevention of fouling.

No general curves are available for estimating from F factor. See vendor bulletins for calculational methods.

Vendor bulletins indicate that pressure drop per theoretical stage will be about 0.5 mmHg or less.

"For Koch-Sulzer packing, Neo-Kloss packing, and Leva film trays, a preliminary estimate can be made by dividing the pressure drop "Relatively good irrigation properties minimize the **potential** for dry spots which promote fouling.

'Vendor considers those values to be extremes normally used but not absolute limits.

Techniques have been developed to permit installation through a manhole. However, it is preferred and usually less costly to provide full shell flanges on either new or existing columns.

'Full shell opening required.



⁽P.G. Nygren, in Schweitzer, 1979).



			Nominal Packing Size, In									
Type of Packing	Mat'l.	1/4	*	1/2	%	3/4	1	1¼	1 1/2	2	3	3½
Super Intalox	Ceramic	_	-	-	-	-	60	-	-	30	-	-
Super Intalox	Plastic	-	-		_	-	33	-	-	21	16	-
Intalox saddles	Ceramic	725	330	200	-	145	98	-	52	40	22	-
Hy-Pak rings	Metal	-	-			-	42	-	-	18	15	-
Pall rings	Plastic	_	-	_	97	-	52	_	40	25	-	16
Pall rings	Metal	-	-	-	70	_	48		28	20	-	16
Berl saddles	Ceramic	900×	_	240×	-	170 ^h	110 ^h		65 ^ħ	45×	-	-
Raschig rings	Ceramic	1,600 ^{5,×}	1,000 ^{b,×}	580°	380°	255°	155ª	125°.×	95°	65 ¹	375.×	-
Raschig rings 32-in wall	Metal	700×	390×	300×	170	155	115×	-	_	-	_	_
Raschig rings	Metal	_	_	410	290	220	137	110×	83	57	32×	_
Tellerettes	Plastic	-		-	-	-	40	-	-	20	-	-
Maspak	Plastic	-		1	1	-	1	1	-	32	20	-
Lessing exp.	Metal	-	1	-	-		-	- 3	0	-	ł	ł
Cross partition	Ceramic	-	, I		L .	-	-		 _	-	70	1
 ⁶/32 Wall ⁶/3, Wall ⁶/3, Wall ⁶/3, Wall ⁶ Packing factors obtained ⁶/2. Wall ⁶/3, Wall												

(c)

Figure 13.37. Corrrelation of flow rates, typical pressure drop behavior, and packing factors of random packed beds. [*Eckert*, *Foote, and Walter*, Chem. Eng. Prog. **62**(1), 59 (1966); *Eckert*, Chem. Eng. (14 *Apr. 1975*)]. (a) Correlation of flow rate and pressure drop in packed towers. (b) Typical pressure drop data: 2 in. porcelain intalox saddles, with F = 40, in a bed 30 in. dia by 10 ft high. (c) Packing factors, F, of wet random packings.

LIQUID DISTRIBUTION

Liquid introduced at a single point at the top of a packed bed migrates towards the walls. Relatively high liquid rates, as in distillation operations where the molal flow rates of both phases are roughly comparable, tend to retard this migration. When liquid rates are low, the maldistribution is more serious. In any event good distribution must be provided initially. A common rule is that the number of liquid streams should be 3-5/sqft in towers larger than 3ft dia, and several times this number in smaller towers. Some statements about redistribution are made at the beginning of this section.

LIQUID HOLDUP

The amount of liquid holdup in the packing is of interest when the liquid is unstable or when a desirable reaction is to be carried out in the vessel. A correlation for Raschig rings, Berl saddles, and intalox saddles is due to Leva (Tower Packings and Packed Tower Design, U.S. Stoneware Co., Akron, OH, (1953):

$$L_{\rm w} = 0.0004 (L/D_{\rm p})^{0.6}$$
, cuft liquid/cuft bed (13.225)

with L in lb liquid/(hr)(sqft) and D_p is packing size (in.). For instance, when L = 10,000 and $D_p = 2$, then $L_w = 0.066$ cuft/cuft.

PRESSURE DROP

Although several attempts have been made to correlate data of pressure drop in packed beds in accordance with the general theory of granular beds, no useful generalization has been achieved. In any event, all manufacturers make available such data measured for their packings, usually only for the air-water system. Samples of such data are in Figures 13.37, 13.38, and 13.40.

13.14. EFFICIENCIES OF TRAYS AND PACKINGS

The numbers of theoretical or equilibrium stages needed for a given vapor-liquid separation process can be evaluated quite precisely when the equilibrium data are known, but in practice equilibrium is not attained completely on trays, and the height of packing equivalent to a theoretical stage is a highly variable quantity. In a few instances, such as in large diameter towers (10 ft or so), a significant concentration gradient exists along the path of liquid flow, so that the amount of mass transfer may correspond to more than that calculated from the average terminal compositions. Mass transfer performance of packed beds is most conveniently expressed in terms of HETS (height equivalent to a theoretical stage), particularly when dealing with multicomponent mixtures to which the concept of HTU (height of a transfer unit) is difficult to apply. In addition to the geometrical configuration of the tray or packing, the main factors that affect their efficiencies are flow rates, viscosities, relative volatilities, surface tension, dispersion, submergence, and others that are combined in dimensionless groups such as Reynolds and Schmidt.

TRAYS

In spite of all the effort that has been expended on this topic, the prediction of mass transfer efficiency still is not on a satisfactory basis. The relatively elaborate method of the AIChE Bubble-Tray Manual (AIChE, New York, 1958) is based on the two-film theory but has not had a distinguished career. A number of simpler correlations have been proposed and have some value as general guidance. That literature has been surveyed recently by Vital, Grossel, and Olsen [Hyd. Proc., 55-56 (Oct. 1984); 147-153 (Nov. 1984); 75-78 (Dec. 1984)].

Efficiency of mass transfer is expressed as the ratio of the actual change in **mol** fraction to the change that could occur if equilibrium were attained

$$E = \Delta y / (\Delta y)_{\text{equilibrium}}.$$
 (13.226)

Because of concentration gradients along the tray, primarily in the liquid phase, the overall efficiency is different from a point efficiency. Since the hydraulics of the tray usually cannot be known accurately, point and overall efficiencies are difficult to relate. In Table 13.15, for instance, three kinds of efficiencies are shown:

- E_{0G} is an overall efficiency based on average changes in the vapor phase mol fraction.
- E_{mv} is the Murphree efficiency, in which $(\Delta y)_{equilib}$ is used as in equilibrium with the liquid leaving the tray.
- E_0 is the ratio of theoretical trays needed for a given separation to the actual number required, and is called the overall efficiency.

Since the efficiency may vary with the position on an individual tray and on the position of the tray in the tower, the three kinds are not the same. When more than one value is shown in Table 13.15 or other literature, the smallest value should be taken as the overall efficiency when that number is needed.

The values of Tables 13.15 and 13.16 probably are not the optima in all cases. The graphs of Figure 13.41 indicate that efficiencies depend markedly on the vapor flow factor, $F = u\sqrt{\rho}$, and there often is a peak in the efficiency curve. Figure 13.42 shows the effect of liquid flow rate across the tray and through the downcomer, measured as a percentage of the flow required to fill the downcomer of this particular tray.

Some of the available methods for estimating tray efficiencies will be described. A useful summary of the AIChE bubble-cap tray method is in the book of King (1980, pp. 621-626). Some of the literature that has found fault with this method is cited by Vital et al. (1984).

The method of O'Connell is popular because of its simplicity and the fact that predicted values are conservative (low). It expresses the efficiency in terms of the product of viscosity and relative volatility, $\mu\alpha$, for fractionators and the equivalent term HP/μ for absorbers and strippers. The data on which it is based are shown in Figure 13.43. For convenience of use with computer programs, for instance, for the Underwood-Fenske-Gilliland method which is all in terms in equations not graphs, the data have been replotted and fitted with equations by Negahban (University of Kansas, 1985). For fractionators,

$$E = 53.977 - 22.527(\log x) + 3.0700(\log x)^2 - 11.000(\log x)^3,$$
 (13.227)

where $\mathbf{x} = \boldsymbol{\mu}\boldsymbol{\alpha}, \boldsymbol{\mu}$ is viscosity (cP) and $\boldsymbol{\alpha}$ is relative volatility. For absorbers and strippers,

$$E = 39.425 + 20.034(\log x) + 1.3480(\log x)^2 - 0.3528(\log x)^3,$$
(13.228)

where $\mathbf{x} = \mathbf{HP}/\mu$, *H* is Henry's law constant [lbmol/(cuft)(atm)], *P* is in atm, and μ is in **cP**.

The equation of McFarland, Sigmund, and Van Winkle [Hyd. Proc., 111-114 (Jul. 1972)] is based primarily on data obtained in pilot plant and laboratory units. It shows a weak dependence on several dimensionless groups. About 800 data points were correlated. The absolute average deviation was 10.6%, and 90% of the calculated values were within 24% of the experimental ones.





(c)

0.06

 $C^2 = V_S^2 D_V / (D_L - D_V)$

0.10

0.12

0.14

0.16

0.06

Figure 13.38. Capacity and pressure drop in beds of pall ("ballast") rings (*Glitsch, Inc.*). (a) Capacity chart for pall rings. V_s = vapor velocity (ft/sec). Example with 1 in. rings,

fractional loading = C/C, $= \begin{cases} 10.161/0.188 = 0.856 \text{ at constant } V/L \\ 0.161/0.188 = 0.856 \text{ at constant } V/L \end{cases}$ 0.161/0.200 = 0.805 at constant gpm.

(b) Pressure drop at 85% of flooding. (c) Pressure drops with 1 and 3.5 in. metal and plastic rings at a range of flow rates.

440

(c)



Figure 13.39. The Sherwood-Lobo correlation of flooding limit in random packed beds. μ_G is the superficial linear velocity of the gas, μ_L/μ_w is the ratio of viscosities of the liquid and water, S_B is the specific surface of the packing (sqft/cuft), ρ_G and ρ_L are densities of gas and liquid, and ε is the fraction voids; the ratio S_B/ε^3 is the factor F of the table with Figure 13.37 [Sherwood, Shipley and Holloway, Ind. Eng. Chem. 30, 765 (1938); Lobo, Friend, and Hashmall, Trans. AIChE 41, 693 (1945)].

EXAMPLE 13.16

Performance of a Packed Tower by Three Methods A packed tower with 3 in. metal pall rings will be analyzed for the system of Example 13.15. The packing factor is F = 15 sqft/cuft. (a) Use the correlation of Figure 13.37:

abscissa =
$$(L/G)\sqrt{\rho_V/(\rho_L - \rho_V)}$$

= $(259,100/271,500)\sqrt{2.75/(29.3 - 2.75)} = 0.3071.$

The ordinate y is read off the figure for several values of $\Delta P/L$; then the flow rate G' and the cross sectional areas are calculated from

$$G' = \sqrt{g_c \rho_V(\rho_L - \rho_V)/F} = 12.52\sqrt{y}, \quad \text{lb/(sec)(sqft)},$$

A = 271,500/3600G' = 75.417/G', sqft,

 ΔP/L
 y
 G'
 A
 D

 0.25
 0.019
 1.726
 43.69
 7.45

 0.50
 0.035
 2.34
 32.23
 6.41

 1.00
 0.046
 2.74
 27.52
 5.92

(b) Apply the method of Figure 13.38 for a tower 7.5 ft dia, 44.18 sqft.

abscissa = 1100/A = 24.9 gpm/sqft, ordinate C = $(27.52/A)\sqrt{2.75/(29.3 - 2.75)}$ = 8.857/44.18 = 0.2004.

The intersection of the line through the origin and the operating point (24.9, 0.2004) with the 3 in. ring line (interpolated) is at

$$C_{F} = 0.27.$$

Therefore,

% flooding = 100(0.2004/0.27) = 74%.

Check the pressure drop by this method.

 $C^2 = (0.2004)^2 = 0.040.$

Interpolating on Figure 13.38(c) to 3 in. pall rings, at 1100/44.18 = 24.9 gpm/sqft,

 $\Delta P/L = 0.35$ in. water/ft,

EXAMPLE **13.16**—(continued)

which is a rough check of the value $\Delta P/L = 0.25$ by method (a). Method (b), however, predicts that flooding would occur when A = 32.23, whereas method (a) says this size is acceptable if the **pressure drop** of 0.50 can be tolerated.

(c) Check the flooding by the Sherwood-Lobo correlation (Figure 13.39):

$$(L/G)\sqrt{\rho_G/\rho_L} = (259,100/271,500)\sqrt{2.75/29.3} = 0.2924$$

The equation is

$$E_{\rm MV} = 7.0 (D_g)^{0.14} ({\rm Sc})^{0.25} ({\rm Re})^{0.08}, \qquad (13.229)$$

where

$$\begin{split} E_{MV} &= \text{percent efficiency,} \\ D_g &= \sigma_L/\mu_L U_V, \\ \mathbf{Sc} &= \mu_L/\rho_L D_{LK}, \\ \mathbf{Re} &= h_W U_V \rho_V / [(\mu_L)(\mathbf{FA})], \\ \sigma_L &= \text{surface tension (lb/hr^2),} \\ \mu_L &= \text{liquid viscosity (lb/hr hr),} \\ U_V &= \text{superficial vapor velocity (ft/hr),} \\ D_{LK} &= \text{diffusivity of light key component (ft^2/hr),} \\ h_W &= \text{height of weir (ft),} \\ \rho_L &= \text{liquid density (lb/ft^3),} \\ \rho_V &= \text{vapor density (lb/ft^3),} \\ \mathbf{FA} &= \text{fractional free area available for vapor flow.} \end{split}$$

The equation of Bakowski [Br. Chem. Eng. 14, 945 (1969); 8, 384, 472 (1963)] is

$$E_{\rm oc} = \frac{1}{1 + 3.7(10^4) KM/h' \rho_l T}$$
(13.230)

where

The equation of Chu, Donovan, Bosewell, and Furmeister [J. Appl. Chem. 1, 529 (1951)] is

$$\log_{10} E = 1.67 + 0.30 \log_{10}(L/V) - 0.25 \log_{10}(\mu_L \alpha) + 0.30h_L,$$
(13.231)

where

L, V = the liquid and vapor flow rates (kmol/sec),

$$\mu_L$$
 = the viscosity of the liquid feed (mN sec/m²),
 α = the relative volatility of the key components,
 h_L = the effective submergence (m), taken as the distance
from the top of the slot to the weir lip plus half the
slot height.

Four of these relations are applied in Example 13.17. The McFarland and Bakowski methods bracket experimental values,

y = 0.53 =
$$[(Q/\pi D^2)^2/16g_c](\rho_G/\rho_L)(u_L/u_w)^{0.1}$$

= (27.52/ πD^2)²2.75/16(32.2)(29.3),
 $D = \begin{cases} 5.64 \text{ ft, at flooding,} \\ 6.31 \text{ ft, at 80\% of flooding.} \end{cases}$

These values are more nearly consistent with the data of Figure 13.37.

whereas the other methods give low values. This comparison, of course, probably is not generally valid. Even experimental values are not exact, unless they are found for exactly the desired operating conditions, the same tray design, and for the same key components. Nevertheless, the collected experimental data and the several correlations that have been cited supply a background on which judgement can be applied to specific problems.

PACKED TOWERS

The most useful measure of the separating power of packed towers is the HETP, the height equivalent to a theoretical plate or stage. It is evaluated simply as the ratio of packed height used for a certain degree of separation to the theoretical number of stages. Its relation to the fundamental quantity, HTU, or the height of a transfer unit, is

$$\text{HETP} = \text{HTU} \ \frac{\ln(mV/L)}{mV/L - 1}, \tag{13.232}$$

where m is the slope of the equilibrium curve. In distillation, the equilibrium and operating lines diverge below the feed point and converge above it. As a result the value of mV/L averages approximately unity for distillation so that HETP and HTU become essentially equal. Usually this is not true in absorption-stripping processes.

Data also are reported as mass transfer coefficients. For the gas phase, the relation to the HTU is

$$(HTU)_{,} = G/k_{G}aP_{,}$$
 (13.233)

where G is the molal flow rate of the gas, say in the units lbmol/(hr)(sqft), P is the total pressure and k_Ga has the units lbmol/(hr)(cuft)(unit of pressure). The liquid phase relation is

$$(HTU), = Lk_L a\rho_L, \tag{13.234}$$

where *L* is the molal flow rate of the liquid [lbmol/(hr)(sqft)], $k_L a$ has the units lbmol/(hr)(cuft)(unit of concentration difference), and ρ_L is the liquid density. The individual HTUs are combined into overall expressions by

$$(\text{HTU})_{0G} = (\text{HTU}), + (m'V/L)H_L,$$
 (13.235)

$$(\text{HTU})_{0L} = (\text{HTU}), + (L/m'V)H_G.$$
 (13.236)

The positions of the slopes m' and m'' of the equilibrium curve are identified in Figure 13.44(a).

Selected data of HETP, $k_G a$, HTU and pressure drop are in Figures 13.45 and 13.46.

Mass Transfer Coefficients. A relation covering liquid and vapor phase mass transfer coefficients is cited in Section 13.9.





Figure 13.40. Comparison of pressure drops through several kinds of packed beds. (a) 2 in. Raschig and pall rings [Ecker! et al., Chem. Eng. Prog. 54(1), 70 (1958)]. (b) 1 in. Tellerettes [Teller and Ford, Ind. Eng. Chem. 50, 1201 (1958)]. (c) 2in. plastic Tripack and other 2 in. packings (Polymer Piping and Metals Co.).

TABLE 13.15. Survey of Tray Efficiencies

	Cal			Woir				
	dia	Droce	Tomp	hoight	Trav	efficiency	96	
System	uia., m	1033.,	•¥	m m	F.	F	F	Remarks
System	111	rгa	n		►0G	-mv	- 0	remains
Bubble-cap								
Ethanol/Water	0.46					98		
						70	95	
	0.11					64.6		
	0.46	101			80-90	90-130		with splash baffle
						95		
	-				61			± 1 1%
	0.46			0.61		85		± 5%
	0.15					99.8		Carey (1934)
	0.196	101				8.0		
Methylcyclohexanel	0 11					64 6		
toluene	0.11					01.0		
Air/water	1 5 0		200				83	
Carbon dioxide/water	1.52		290			0 0	125	no mixing
Carbon dioxide/water	0.076					00	120	
A							100	50% mixing
Acetic acid/water	0.46	101				65		
• • • • •	0.59	56				54		± 5%
Deuterium/hydrogen	0.027		20			50	44	± 3%
Oxygen/nitrogen							76	± 9%
Acetone/water	5.49				91			
	_				83			± 2%
Ethylene dichloride/	_			0.032		95		
tofuene								
Sugar/water	1 52					8.0		
CHCLICCL	1.02				0.0	00		↓ 5%
	0.005	101	2.0.2		90	77		<u>1</u> 3 %
Ammonia/water	0.305	101	283			11		
Methanol/water	1.0					90		
	—					70		
Acetone/benzene					79			+1%
Methanol/Isopropanol/	0 4 5				69	6.8		1,1,0
water					70	00		
	_				6.8			
Acetone/methanol/water	0.45				60			
A second methanol water	0.45				0.0			
	-				60			
Canalina atabilizara	4 44	4 000			80	4.0.0		D (1000)
	1.44	1,620			15	100		Brown (1936)
benzene/loluene	0.2					60		Lewis (1930)
	0.2					70		Carey (1934)
	0.15					58		Carey (1934)
Aniline/water	0.2					58		Carey (1934)
Naphtha/water	2.74					65		Lewis (1928)
Isopropanol/water	0.45					78		
Methanol/isopropanol	0.45					64		
Acetone/methanol	0.45					61		
Benzene/toluene/xylene	0.2					75		Lewis (1930)
Naphtha/pinene/aniline	0.2					90		Lewis (1930)
Siovo								(`````)
	0 070	101			45 5			
Ethanol/water	0.0/6	101			40.5	0 5		
	0.127					85		
	0.196	101				90		
						71.4		CA-1 00455C (1970)
	Lab						120	Brown (1936)
Methylethylketone/Water	0.08				41			
Acetone/water	0.05	101			25.5			
	0.11				43.5			± 11.5%
	0.15		373			80		
Benzene/water	0.05	101			9.6			
Toluene/water	0.05	101			7.1			
n-Heptane/	0 04	101			1.1	77.6		+ 2.07%
methylcyclohexane	0.04							T 2:01 /0
n-Hentane/ovolohovano	4.0	105		0.05			0 5	
перанелсускопехане	1.2	105		0.00			00	
	2.44	165		0.05			75	
T . 1	1.2	164		0.05		90		60% flood
ioiuene/	0.15	101					54.6	± 15%
methylcyclohexane	-	27					55.5	± 5%

TABLE 13.15—(continued)

	Col.			Weir				
Sustam	dia.,	Press.,	Temp.,	height,	Tray	efficiency,	%	Demente
System Sieve cent	111	KPa	- N	m	EoG	⊂ mv	c,	Remarks
Sieve, cont.								
Methylcyclohexane/	0.05	101				0 0	91	
Propane/butane						0 0	100	+ 5%
Carbon dioxide/water	0.08					8 0	125	no mixing
						.	100	50% mixing
n-Octane/toluene	0.15	111	298	0 0 2 5		89.7	3.8	
Air/water/ammonia	0.08		298	0.08	85.7	98.4		±0.1
	-			0.03	70	<u>.</u>		
Oxygen/water/ammonia	0 15					65 75	50	+7%
Ammonia/water	0.3	101	283			89		2 · /·
Methylisobutylketone/	0.08		298	0.08	41.5	64		± 1.7
water Ethylopo dichlorido/	0.05	404					75	
tofuene	0.05	101					15	
Methylethylketone/toluene	0.15			0.05			88	
Air/ethanol				0.1	80			± 20%
Air/propanoi Mothanol/CCI	0 11			0.03	11			± 5%
Methanol/water	0.11				20.7			+ 13%
	0.11				79.2			T 10/0
	1.0					93		0.4.4.00.4FFO (4070)
AcetonalCCL	0 11				5.0	90		CA-1 UU455C (1970)
Isopropanol/water	0.11				50		72.9	CA-100455C (1970)
Benzene/toluene	0.127				76 5	75		J 7 10%
	10.7	101	353	0.08	10.5	80.5		1 pass
	-					80.5		2 pass
				0.14		85.2		4 pass
Benzene/methanol	0.18	690		0.05	86.4	94.2	85	
Ethylbonzonolstvrono	2.4	103					70	
Emylocitzeneistyrene	0.5	13		0.038		75		
Helium/					90			±5%
methylisobutylketone				0.038	0.0			
Nitrogen/cyclohexanol	_			0.051	80 70			+20%
Acetic acid/water	0.46	101				75		
Benzene/propanol	0.46					58.6		96.6% flood
toluono	0.038				78			
tordene	_				93			
n-Heptane/toluene	-			0.02		4 5		
n-Hentanelbenzene				0 0 2		62 55		
in heptaneisenzene				0.02		68		
CC&/benzene	0.032						71	no vapor pulsing
Isobutaneln-butane		2 069				110	73	vapor pulsing
Ethanol/water/furfural	_	2,000				80		CA-165714Y (1980)
n-Hexane/ethanol/	0.1	101	333		70	•••		+0.9
methylcyclopentane	-				70.3			- 2.5
n-Hexane/ethanol/	0 1	101	333		71 55			- 4.4 + 14.3
methylcyclopentane/	J. I	101	000		60			-1.5
benzene	_				6			-1.9
Benzene/n-propanol	0.46	101	366	0.08		54 57		± 5% , 60% flood
Toluene/n-propanol	0.46	101	366	0.08		61	65	± 5%, 80% flood
Deservation	0.03	101	366			57		± 8%, 60% flood
Beer/water	Comm.						120	Brown (1936)

TABLE 13.15—(continued)

System	cot. dia., m	Press., kPa	Тетр., °К	Weir height, m	Tray E₀ g	v efficiency, E mv	% E,	Remarks
Sieve, cont.								
Air/water	-	101	297	0.05		62		
triethylunenlvcol								
APV-West								
Methanol/water	1.0					81		
Kascade								
Ethanol/water	0.2					54.1		
•• • • • • • • • •	0.2						70	
Methylcyclohexane/toluene	0.2					44.6	7 0	
Oxvgen/water	0 2					84	12	
Tunnel	0.2							
Furfural/isobutane & butylene	4.0	593	318			25		± 4%, ΔP = 12.5
Furfural/n-butane & butylene	4.0	593	318			2 5		± 7%, ΔP = 12.5
Turbogrid								
Ammonia/water	0.3	101	283			75		
Etnanol/water	-	0.24				85	85	⊥ 11 0 ⁄0
Methanol/water		101				87	00	I 11 70 + 2%
	0.1	101	298			95		+ 3.9%
Methanol/isopropanol/	Small 0.15					86 66.4		-
water Methanol/isopropanol	Small					65		
V-Grid								
Air/water/ammonia						70	60	
Combination valve-sieve								
Benzene/propanol	0.46				32		0.0	
Etnylenebenzene/styrene	_	13					00	
Wyatt Perfavalve	0.46					767		74 7% flood
Propanolibenzene	0.46					55.5		44.4% flood
Valve								
Benzene/toluene/xvlene	2.43					69		+ 3%
Ethanol/water	0.032						70	T = 12
	0.06					56		CA-1 07562W (1975)
n-Propanol/benzene	0.46					73		± 5%, 60% flood
n-Propanol/toluene Ethylbenzene/styrene	0.46	13				85		± 5%, 80% 11000
Benzene/C aromatics	2/3	15			11	00	8.8	top
	2.45				66		00	bottom
Round valve	0 196	101				9.2		
	0.130	101				01		
E-Type valve	0.196	101				68		
Nutter valve								
C ₆ /C ₇	_	165					96	float valve
iC₄/nC₄		1,131					121	float valve
Propanollbenzene	0.5					63.8		29.6% flood
Propanol/toluene	0.5		303			75.3	0.0	20% flood
	1.2						90	20 /0 1100u
Koch Valve	1 0						03 0	50% flood
Benzene/toluene/xvlene	2.43	92	346	0.08		48	88	
Glitsch	-							
n-Butanelisobutane	1.3	1,138		0.05			122	V-I Ballast
Cyclohexaneln-heptane	1.3	165		0.05		99.5		V-I Ballast
Mathemal/water	1.2						97	21% flood, Valve
wethanol/water	0.1	101		0 05		8.8	пıgn	+ 2%, Value (downcomerc)

TABLE 13.15—(continued)

coi. Weir								
System	dia., m	Press., kPa	Temp., °K	height, m	Tray E _o a	efficiency, [[] mv	% E。	Remarks
Ripple								
Methanol/water	1.0					73		
Ammonia/water	0.3	101				8 2		
Light gasoline	2.0						48	CA-88739C (1968)
Unifiex								
Methanol/water	1.0					87		
Baffle								
Toiuenelmethylcyclohexane		101				87		
Angie		404						
Methanol/water		101					70	CA-1 809/g (1972)
Crossflow plate								
Benzene/toluene						80		
Ethanol/water	_					70		
Jet								
Air/water/		101	297	0.05		6 5		
Methylethylketone/toluene	0.15			0.03			93	

[References given in the original: Vital, Grossel, and Olsen, Hyd.

Proc., 147-153 (Nov. 1984)].

TABLE 13.16. Efficiency Data of Some Operations with Bubblecap, Ripple, and Turbogrid Trays

		Column	Tray	n	Static	Effic	cy, %]
Disperser	System	diameter, ft	in	psia	submergence. in	Emo*	Eact	Remarks	Ref
Bubble cap	Ethanol-water Methanol-water Ethyl benzene-styrene cyc Johexane-n-heptane	1.31 1.31 2.5 3.2 2.6 4.0	10.6 163 14 15.7 197 24	147 14 7 14.7 14.7 1.9 14 7 24 50	/ 18 1 78 1.2 10 02 0.25 4.25	83-87 84-97 80-85 90-95 55-68 65-90 65-90 65-90			
	Cyclohexane-n-heptane Benzene-toluene Toluene-isooctane	4.0 1.5 5.0	2 4 15.7 2 4	5 2 4 14.7 14.7	06 1.5 0.4	6 5-85 75-100 70-60	60-80	Tunnel caps	6 7 8
Ripple sieve	Methanol-water Ethanol-water Methanol-water Ethyl benzene-styrene Benzene-toluene Methyl alcohol-n-propyl alcohol-	3.2 2.5 3.2 2.6 1.5	151 14 15.7 197 15.7	14.7 14.7 14 7 19 14.7	1.0 1.57 0.75 30	70-90 75-85 90-100 70 60-80		10.8% open 10.4% 4.8% open 12.3% open 18% open	3 2 3 9 7
	sec-butyl alcohol Mired xylenes + Cg-C10 paraffins and naphthenes Cyclohexane-n-heptane Isobutane-n-butane	6.0 1 3 0 4.0 4.0 4.0 4.0 4.0 4.0	18 21 24 24 24 24 24 24 24 24 24	18 25 5 24 5 165 165 300	1.33 1.25 2.0 2.0 2.0 2.0 2.0	60-70 80 70-60 110 120 110	64 86	14% open 14% open 6% open 14% open 8% open 6% open	10 5 13 13 13 12 13 12 12
Turbogrid valve	Methanol-water Ethanol-water Ethyl benzene-styrene Cyclohezane-n-heptane n-Butane-isobutene Benzene-toluene	4.0 3.2 2.5 2.6 4.0 4.0 1.5	24 15.7 14 19.7 24 24 24 15 7	400 14 7 14.7 19 20 165 14.7	20 1.0 0.75 3.0 3.0 3.0 3.0	100 7080 7585 7585 7580	5096 104-121	6% open 14.7% open Rect. valves Rect. valves	12 3 2 4 11 11 7

 Benzene-totuene
 Lot
 Aut

 References
 1. Kirschbaum, 2. Ver. Disch. Ing. Beth. Verfahrenstech., (5). 131 (1938); (3), 69 (1940).
 Xirschbaum, 2. Ver. Disch. Ing. Beth. Verfahrenstech., (5). 131 (1938); (3), 69 (1940).

 2. Kirschbaum, Distillier-Rektifizieriechnik, 4th. ed., Springer-Verlag, Berlin and Heidelberg. 1969.
 X. Katanek and Standart, Sep. Sci. 9, 439 (1967).

 3. Katanek and Standart, Sep. Sci. 9, 439 (1967).
 Billet, Chem. Ing. Tech... 38, 825 (1966); 40.377 (1968).

 5. AIChE Research Committee. Tray Efficiency in Distillation Columns, final report. University of Delaware, Newark, 1958.

 6. Raichle and Billet, Chem. Ing. Tech... 35, 801 (1963).

 7. Zuiderweg, Verburg, and Hinds, Ind. Eng. Chem., 49, 2051 (1957).

 9. Billet, Froc. Intn. Symp., Brighton, England, 1960.

 8. Manning. Marple, and Hinds, Ind. Eng. Chem., 49, 2051 (1957).

 9. Billet, Church. Green, Lee, and Rasmussen, Ind. Eng. Chem., 44, 2238 (1952).

 11. Fractionation Research, Inc.. "Report of Tests of Nutter Type B Flat Valve Tray." July 2. 1964, from Nutter Engineering Co., T 'ulsa.

 12. Sakata and Yanagi. Inst. Chem. Eng. Supp. Ser., no. 56, 32/21 (1979).

 13. Yanagi and Sakata, Ind. Eng. Chem. Process Des. Dev. 21.712 (1982).

 Ysee Eq. (18-20).

 Ysee Eq. (18-20).

 Ysee Ze, (18-20).

 Ysee Ze, (18-20).

 NOTE: To convert feet to meters NOTE: To convert feet to meters. multiply by 0.3048; to convert inchest to centimeters. multiply by 2.54, and to convert pounds-force per square inch to kilopascals, multiply by 6.895.

(Chemical Engineers' Handbook, McGraw-Hill, New York, 1984).

100

HETP Correlations. Most of the data available for correlation laboratory data and not indicative of large scale behavior except are perhaps on a comparative basis. Some guidelines for full scale tray behavior are stated by Frank [Chem. Eng. 111, (14 Mar. 1977)] in this table:

Type of Packing/Application	HETP (m)			
25 mm dia packing	0.46			
38 mm dia packing	0.66			
50 mm dia packing	0.9			
Absorption duty	1.5-I <u>.</u> 8			
Small diameter columns (<0.6 m dia.)	column diameter			
Vacuum columns	values as above +0.1 m			



A correlation for Raschig rings and Berl saddles by Murch [Ind. Eng. Chem. 45, 2616 (1953)] covers columns up to 30in. dia and 10 ft high. His relations are

HETP =
$$C_1 G'^{C_2} d_c^{C_3} Z^{1/3} \alpha \mu_L / \rho_L$$
, (13.237)

where

G' = mass velocity of vapor $(kg/m^2 sec)$ of tower area,

 d_c = column diameter (m),

Ζ = packed height (m),

- = relative volatility, α
- $\mu_L = \text{liquid viscosity} \quad (\text{N sec/m}^2),$ $\rho_L = \text{liquid density} \quad (\text{kg/m}^3).$



Figure 13.41. Efficiencies of some fractionations with several types of trays as a function of vapor factor $F = u\sqrt{\rho}$ or linear velocity. (a) Data of methanol/water in a column 3.2 ft dia [data of Kastanek, Huml, and Braun, Inst. Chem. Eng. Symp. Ser. 32(5), 100 (1969)]. (b) System 5.100, (1969)].



Figure 13.41—(continued)

Values of the constants C_i appear in this tabulation:

Type of Packing	Size (mm)	<i>С</i> , (× 10 ⁻⁵)	C ₂	C3
Rings	6			1.24
-	9	0.77	-0.37	1.24
	12.5	7.43	-0.24	1.24
	25	1.26	-0.10	1.24
	50	1.80	0	1.24
Saddl es	12.5	0.75	-0.45	1.11
	25	0.80	-0.14	1.11

A correlation for 25 and 70mm Raschig rings by Ellis [*Birmingham Univ. Chem.* Eng. **5**(1), 21 (1953)] with HETP (m) is

$$\text{HETP} = 18d_r + 12m[(G'/L' - 1)], \qquad (13.238)$$



Figure 13.42. Efficiency of Glitsch V-l valve trays on isobutane/butane and cyclohexane/n-heptane as a function of vapor density and percent of flood, measured by Fractionation Research Inc. (*Glitsch Inc., Bulletin 160, Dallas, TX, 1958*).



- 5: Sieve plate, $F_{c} = 0.042$, $h_{w} = 40$ mm, d = 4.0 mm
- 6: APV-West plate, $F_{c} = 0.1072$, $h_{w} = 45$ mm
- 7: Glitsch valve plate, Al, $F_c = 0.14$, $h_w = 50$ mm
- 8: Turbogrid, set IV, $F_{c} = 0.147, d = 4.5 \text{ mm}$

(f)

where

 d_r is the diameter of the rings (m), m is the average slope of equilibrium curve,

G' is the vapor mass flow rate,

L' is the liquor mass flow rate.

HTU data have been correlated by Cornell et al. (1960) and updated by Bolles and Fair *[Inst. Chem. Eng. Symp. Ser.* **56**(2), 3.3/35 (1979)]. Pall rings, Raschig rings, and saddles are covered in the original article, but only the pall ring results are quoted here. Separate relations for the liquid and vapor phases are represented by Eqs. (13.239) and (13.240) and Figure 13.44.

$$H_L = \phi(\mathrm{Sc}_L)^{0.5} (C) (Z/10)^{0.15}$$
(13.239)

where

 H_L = height of liquid phase transfer unit (ft), ϕ = parameter from figure, SC, = liquid phase Schmidt number, $\mu_L/\rho_I D_L$,

C = flooding correction factor from figure,

Z = height of column packing (ft);

$$H_{V} = \frac{\psi(\mathrm{Sc}_{V})^{0.5}}{(L_{m}f_{1}f_{2}f_{3})^{0.6}} \left(\frac{d_{c}}{12}\right)^{1.24} \left(\frac{Z}{10}\right)^{1/3},$$
(13.240)

where

$$\begin{split} H_V &= \text{height of gas film transfer unit} \quad (\text{ft}), \\ \text{Sc,} &= \text{vapor-phase Schmidt number,} \\ d_c &= \text{column diameter} \quad (\text{in.}), \\ Z &= \text{packing height} \quad (\text{ft}), \\ \psi &= \text{parameter from figure,} \\ L_m &= \text{liquid flow rate} \quad (\text{lb/hr}) \text{ ft}^2), \\ f_1 &= (\mu_I / 1.005)^{0.16}, \ \mu_L (\text{cP}), \\ f_2 &= (1/\rho_L)^{1.25}, \ \rho_L (\text{g/cm}^3), \\ f_3 &= (72.8/\sigma)^{0.8}, \ \sigma (\text{dyn/cm}). \end{split}$$


Figure 13.43. Efficiencies of fractionators and absorber-strippers. The original curves of O'Connell [Trans. AIChE 42, 741 1946)] have been replotted and fitted with equations, as shown on the figures, by S. Negahban (University of Kansas, 1985). (a) Fractionators (the viscosity μ is in cP). (b) Absorbers and strippers; H = Henry's law constant in lb mol/(cuft)(atm), P is in atm, and μ is in cP.

Tray Efficiency for the Separation of Acetone and Benzene (a) Method of McFarland et al.: The operating data are taken from their article, as follows:

Acetone mole fraction, $x_1 = 0.637$, Benzene mole fraction, $x_2 = 0.363$, Temperature T ("F) = 166, Superficial vapor mass velocity G (lb/hr sqft) = 3820, Vapor velocity u_{ν} (ft/hr) = 24,096, Weir height, h_{w} (ft) = 0.2082, Fraction free area FA = 0.063.

The pertinent physical properties of the mixture are

$$\begin{split} \mu_L &= 0.609 \;\; \text{lb/ft hr}, \; 0.252 \;\; \text{cP}, \\ \sigma_L &= 5.417(10^5) \;\; \text{lb/sqft hr}, \; 18.96 \;\; \text{dyn/cm}, \\ D_{\text{light key}} &= 2.32(10^{-4}) \; \text{sqft/hr}. \end{split}$$

The dimensionless groups appearing in the correlation are

$$\begin{split} N_{Dg} &= \sigma_L / \mu_L u_V = 37, \\ N_{Sc} &= \mu_L / \rho_L D_{LK} = 55, \\ N_{Re} &= h_w G / \mu_L (\text{FA}) = 2.07(10^4). \end{split}$$

The tray efficiency is found with Eq. (13.229):

$$E = 7.0 (N_{Dg})^{0.14} (N_{Sc})^{0.25} (N_{Re})^{0.08} = 71\%.$$

(b) Method of O'Connell: The relative volatility is 3.24 at x = 0.05 and 1.63 at x = 0.95, or a geometrical mean value of

 $\alpha = 2.30. \text{ Accordingly},$ $\mu \alpha = 0.252(2.3) = 0.58,$ and, from Figure 13.43 or Eq. (13.227), E = 56%.(c) Method of Bakowski, Eq. (13.230): K = y/x = 1.20 at x = 0.637, M = 58, h' = 50 mm, $p, = 820 \text{ kg/m}^3,$ T = 348 K, $E = \frac{100}{1 + 37,000(1.20)(58)/50(820)(348)} = 84.7\%.$

(d) Method of Chu et al., Eq. (13.231):

$$L/V = \begin{cases} 0.8, & \text{above the feed,} \\ 1.2, & \text{below the feed,} \end{cases}$$

log $E = 1.67 - 0.25 \log(0.58) + 0.3(0.05) + 0.3 \log(L/V)$
 $= 1.744 + 0.3 \log(L/V),$
 $E = \begin{cases} 51.9\%, & \text{above the feed,} \\ 58.6\%, & \text{below the feed.} \end{cases}$

(e) Experimental data: Table 13.15 shows E = 79% for acetone/ benzene in bubblecap tower and E = 85% for methanol/benzene with sieve trays. Figure 13.41 shows that efficiencies above 80% are readily attainable near $u\sqrt{\rho} = 1.0$.



Figure 13.44. Factors in Eqs. (13.239) and (13.240) for **HTUs** of liquid and vapor films; and slopes m' and m'' of the combining Eqs. (13.235) and (13.236): [Bolles and Fair, Inst. Chem. Eng. Symp. Ser. **56**(2), 3.3/3.5, (1979)]. (a) Definitions of slopes m' and m'' in Eqs. (13.235) and (13.236) for combining liquid and gas film **HTUs**; $\beta = 1$ for equimolal counter diffusion; $\beta = (x_B)_{mean}$ for diffusion through a stagnant film. (b) Factor ϕ of the liquid phase Eq. (13.239). (c) Factor C of the liquid phase, Eq. (13.239). (d) Factor ψ of the gas phase, Eq. (13.240), for metal pall rings.



Figure 13.45. Number of stages per meter (reciprocal of HETP), pressure loss per meter and pressure loss per theoretical stage in a 500 mm dia column filled with metal pall rings. Other charts in the original show the effects of packing height and column diameter, as well as similar data for Raschig rings (Billet, 1979). (a) Methanol/ethanol at 760Torr and total reflux in a column 500 mm dia. (b) Ethylbenzene/styrene at 100 Torr and total reflux in a column 500 mm dia.



Figure 13.46. Data of HETP, HTU, and Ka for several systems and kinds of tower packings. (a) K_{ga} for absorption of ammonia in NaOH with various packings (Polymer Piping and Materials Co.). (b) HETP of several packings as functions of the gas rate (I. Eastham, cited by Coulson and Richardson, 1978, Vol. 2, p. 515). (c) HETP for ethylbenzene/styrene at 100 Torr; curve 1 for 2in. metal pall rings [Billet, Chem. Eng. Prog. 63(9), 55 (1967)]; curve 2 for 1 in. metal pall rings (Billet, loc. cit.); curve 3 for Sulzer packing (Koch Engineering Co.). (d) K_{ga} for absorption of CO, in NaOH with various packings; CMR are cascade mini rings which are pall rings with heights about one-half the diameters (Muss Transfer International). (e) HTU for absorption of ammonia from air with water (Wen, S.M. Thesis, University of West Virginia, 1953). (g) Typical HETP for distillation with 2 in. metal pall ("ballast") rings (redrawn from Bulletin 217, Glitsch Inc.).



(c)



NB The differential performance between No3 Metal CMR and the **other rings** in this test is increased by a further 10/15% in K₂ CO, systems. This is confirmed by extensive feed-back data from operating plants up to 13 ft (4 m) diameter.

🗕 No 3 META	L CMR
No2 Hy-F	Pak®
2in META	L PALL RING
COLUMN DIA	= 30 ins
PACKED HEIGHT	= 10 ft
GAS RATE	$= 900 lbs/ft^2 hr$
GAS CONC	= 1% co,
LIQUID CONC	= 4% NaOH
LIQUID TEMP	= 75°F
CARBONATE CONC	= 25%

Figure 13.46(continued)





REFERENCES

General

- R. Billet, *Distillation* Engineering, Chemical Publishing Co., New York, 1979.
- J.M. Coulson and J.F. Richardson, *Chemical* Engineering, Pergamon, New York, 1978, Vol. 2.
- J.R. Fair, Liquid-gas systems, in *Chemical Engineers Handbook*, McGraw-Hill, New York, 1984, Section 18.
- 4. R.J. Hengstebeck, Distillation, Reinhold, New York, 1961.
- E.J. Henley and J.D. Seader, *Equilibrium-Stage Processes in Chemical* Engineering, Wiley, New York, 1981.
- A.L. Hines and R.N. Maddox, Mass Transfer Fundamentals and Applications, Prentice-Hall, Englewood Cliffs, NJ, 1985.
- 7. C.D. Holland, Fundamentals of Multicomponent Distillation, McGraw-Lin ?.,^....^L_1001

- C.S. Robinson and E.R. Gilliland, Elements of Fractional Distillation, McGraw-Hill, New York, 1950.
- J.D. Seader, Distillation, in *Chemical Engineers Handbook*, McGraw-Hill, New York, 1984, Section 13.
- B.D. Smith, Design of Equilibrium Stage Processes, McGraw-Hill, New York, 1963.
- 13. R.E. Treybal, Mass Transfer Operations, McGraw-Hill, New York, 1980.
- S.M. Walas, Phase Equilibria in Chemical Engineering. Butterworths, Stoneham, MA, 1985.

Special Topics

- D.B. Broughton and K.D. Uitti, Distillation estimates for naphtha cuts, in *Encyclopedia of Chemical Processing and Design*, Dekker, New York, 1982, Vol. 16, pp. 186198.
- 9 IS Eckert Design of nacked columns in Reference & 1070

REFERENCES 457

- P.G. Nygren, High efficiency low pressure drop packings, in Ref. 8, 1979, pp. 1.241-1.253.
- D.F. Othmer, Azeotropic and extractive distillation, *Encycl. Chem.* Tech. 3, 352-377 (1978).
- G. Prokopakis, Azeotropic and extractive distillation, *Encycl. Chem. Technol. Suppl.*, 145-158 (1984).
- P.A. Schweitzer, Editor, Handbook of Separation Techniques for Chemical Engineers, McGraw-Hill, New York, 1979.
- W.J. Stupin and F.J. Lockhart, Distillation, thermally coupled, in Encyclopedia of Chemical Processing and Design, Dekker, New York, 1982, Vol. 16, pp. 279-299.
- T.J. Vital, S.S. Grossel, and P.I. Olsen, Estimating tray efficiency, *Hyd. Proc.*, 55-56 (Oct. 1984); 147-153 (Nov. 1984); 75-78 (Dec. 1984).

- Vapor-Liquid Equilibrium Data Collections
- API Technical Data Book-Petroleum Refining, American Petroleum Institute, Washington, D.C., 1983-date, Chaps. 8 and 9.
- J. Gmehling, U. Onken et al., Vapor-Liquid Equilibrium Data Collection, DECHEMA, Frankfurt/Main, Germany, 1979–date.
- 3. M. Hirata, et al., Computer Aided Data Book of Vapor Liquid Equilibria, Elsevier, New York, 1976.
- 4. V.B. Kogan, et al., *Equilibria between Vapor and Liquid* (in Russian), **Izdatelstvo** Nauka, Moscow, 1966.
- 5. Landolt-Boernstein Zahlenwerte und Funktionen, II2a, 1960; IV4b, 1972. New Series Group IV, Vol. 3, 1975, Springer, Berlin.
- 6. NGPSA Engineering Data Book, Natural Gas Processors Suppliers Associations, Tulsa, OK, 1972, Chap. 18, and later editions.

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EXTRACTION is a process whereby a mixture of several substances in the liquid phase is at least partially separated upon addition of a liquid solvent in which the original substances have different solubilities. When some of the original substances are so/ids, the process is called leaching. In a sense, the role of solvent in extraction is analogous to the role of enthalpy in distillation. The solvent-rich phase is called the extract, and the solvent-poor phase is called the raffinate. A high degree of separation may be achieved with several extraction stages in series, particularly in countercurrent flow.

Processes of separation by extraction, distillation, crystallization, or adsorption sometimes are equally possible. Differences in Solubility, and hence of separability by extraction, are associated with differences in chemical structure, whereas differences in vapor pressure are the basis of separation by distillation. Extraction often is effective at near-ambient temperatures, a valuable feature in the separation of thermally unstable natural mixtures or pharmaceutical substances such as penicillin.

The simplest separation by extraction involves two substances and a solvent. Equilibria in such cases are represented convenient/y on triangular diagrams, either equilateral or right-angled, as for example on Figures 14.1 and 14.2. Equivalent representations on rectangular coordinates a/so are shown. Equilibria between any number of substances are representable in terms of activity coefficient correlations such as the UNIQUAC or NRTL. In theory, these correlations involve only parameters that are derivable from measurements on binary mixtures, but in practice the resulting accuracy may be poor and some multicomponent equilibrium measurements a/so should be used to find the parameters. Finding the parameters of these equations is a complex enough operation to require the use of a computer. An extensive compilation of equilibrium diagrams and UNIQUAC and NRTL parameters is that of Sorensen and Arlt (1979- 1980). Extensive bibliographies have been compiled by Wisniak and Tamir (1980- 1981).

The highest degree of separation with a minimum of

solvent is attained with a series of countercurrent stages. Such an assembly of mixing and separating equipment is represented in Figure 74.3(a), and more schematically in Figure 14.3(b). In the laboratory, the performance of a continuous countercurrent extractor can be simulated with a series of batch operations in *Separatory* funnels, as in Figure 14.3(c). As the number of operations increases horizontally, the terminal concentrations E_1 and R_3 approach asymptotically those obtained in continuous equipment. Various kinds of more sophisticated continuous equipment also are wide/y used in laboratories; some are described by Lo et al. (1983, pp. 497-506). Laboratory work is of particular importance for complex mixtures whose equilibrium relations are not known and for which stage requirements cannot be calculated.

In mixer-separators the contact times can be made long enough for any desired approach to equilibrium, but 80–90% efficiencies are economically justifiable. If five stages are required to duplicate the performance of four equilibrium stages, the stage efficiency is 80%. Since mixer-separator assemblies take much floor space, they usually are employed in batteries of at most four or five units. A large variety of more compact equipment is being used. The simplest in concept are various kinds of tower arrangements. The relations between their dimensions, the operating conditions, and the equivalent number of stages are the key information.

Calculations of the relations between the input and output amounts and compositions and the number of **extraction stages** are **based** on material balances and equilibrium relations. Know/edge of efficiencies and capacities of the equipment then is applied to find its actual size and configuration. Since extraction processes usually are performed under adiabatic and isothermal conditions, in this respect the design problem is simpler than for thermal separations where **enthalpy** balances a/so are involved. On the other hand, the design is complicated by the fact that extraction is feasible on/y of **nonideal** liquid mixtures. **Consequently**, the activity coefficient behaviors of two liquid phases must be taken into account or direct equilibrium data must be **available**.

14.1. EQUILIBRIUM RELATIONS

On a ternary equilibrium diagram like that of Figure 14.1, the limits of mutual solubilities are marked by the binodal curve and the compositions of phases in equilibrium by tielines. The region within the dome is two-phase and that outside is one-phase. The most common systems are those with one pair (Type I, Fig. 14.1) and two pairs (Type II, Fig. 14.4) of partially miscible substances. For instance, of the approximately 1000 sets of data collected and analyzed by Sorensen and Arlt (1979), 75% are Type I and 20% are Type II. The remaining small percentage of systems exhibit a considerable variety of behaviors, a few of which appear in Figure 14.4. As some of these examples show, the effect of temperature on phase behavior of liquids often is very pronounced.

Both equilateral and right triangular diagrams have the property that the compositions of mixtures of all proportions of two mixtures appear on the straight line connecting the original mixtures. Moreover, the relative amounts of the original mixtures corresponding to an overall composition may be found from ratios of line segments. Thus, on the figure of Example 14.2, the amounts of extract and raffinate corresponding to an overall composition \mathbf{M} are in the ratio $E_1/R_N = MR_N/E_1M$.

Experimental data on only 26 quaternary systems were found by Sorensen and Arlt (1979), and none of more complex systems, although a few scattered measurements do appear in the literature. Graphical representation of quaternary systems is possible but awkward, so that their behavior usually is analyzed with equations. To a limited degree of accuracy, the phase behavior of complex mixtures can be predicted from measurements on binary mixtures, and considerably better when some ternary measurements also are available. The data are correlated as activity coefficients by means of the UNIQUAC or NRTL equations. The basic principle of application is that at equilibrium the activity of each component is the same in both phases. In terms of activity coefficients this



Figure 14.1. Equilibria in a ternary system, type 1, with one pair of partially miscible liquids; A = 1-hexene, B = tetramethylene sulfone, C = benzene, at 50°C (R.M. De *Fre, thesis, Gent, 1976*). (a) Equilateral triangular plot; point *P* is at 20% A, 10% B, and 70% C. (b) Right triangular plot with **tielines** and **tieline** locus, the amount of A can be read off along the perpendicular to the hypotenuse or by difference. (c) Rectangular coordinate plot with **tieline** correlation below, also called Janecke and solvent-free coordinates.

condition is for component *i*,

$$\gamma_i x_i = \gamma_i^* x_i^*, \tag{14.1}$$

where * designates the second phase. This may be rearranged into a relation of distributions of compositions between the phases,

$$\boldsymbol{x}_i^* = (\boldsymbol{\gamma}_i / \boldsymbol{\gamma}_i^*) \boldsymbol{x}_i = \boldsymbol{K}_i \boldsymbol{x}_i, \tag{14.2}$$

where K_i is the distribution coefficient. The activity coefficients are functions of the composition of the mixture and the temperature. Applications to the calculation of stage requirements for extraction are described later.

Extraction behavior of highly complex mixtures usually can be known only from experiment. The simplest equipment for that purpose is the separatory funnel, but complex operations can be simulated with proper procedures, for instance, as in Figure 14.3(c). Elaborate automatic laboratory equipment is in use. One of them employs a 10,000–25,000 rpm mixer with a residence time of 0.3–5.0 sec, followed by a highly efficient centrifuge and two chromatographs for analysis of the two phases (Lo et al., 1983, pp. 507).

Compositions of petroleum mixtures sometimes are represented adequately in terms of some physical property. Three examples appear in Figure 14.5. Straight line combining of mixtures still is valid on such diagrams.

Basically, compositions of phases in equilibrium are indicated with tielines. For convenience of interpolation and to reduce the clutter, however, various kinds of **tieline** loci may be constructed, usually as loci of intersections of projections from the two ends of the tielines. In Figure 14.1 the projections are parallel to the base and to the hypotenuse, whereas in Figures 14.2 and 14.6 they are horizontal and vertical.

Several **tieline** correlations in equation form have been proposed, of which three may be presented. They are expressed in weight fractions identified with these subscripts:

CA solute C in diluent phase A CS solute C in solvent phase S SS solvent S in solvent phase S AA diluent A in diluent phase A AS diluent A in solvent phase S SA solvent S in diluent phase A.







Figure 14.2. Equilibria in a ternary system, type II, with two pairs of partially miscible liquids; A = hexane, B = aniline, C = methylcyclopentane, at 34.5°C [Darwent and Winkler, J. Phys. Chem. 47, 442 (1943)]. (a) Equilateral triangular plot. (b) Right triangular plot with tielines and tieline locus. (c) Rectangular coordinate plot with tieline correlation below, also called Janecke and solvent-free coordinates.



Figure 14.3. Representation of countercurrent extraction batteries. (a) A battery of mixers and settlers (or separators). (b) Schematic of a three-stage countercurrent battery. (c) Simulation of the performance of a three-stage continuous countercurrent extraction battery with a series of batch extractions in separatory funnels which are designated by circles on the sketch. The numbers in the circles are those of the stages. Constant amounts of feed F and solvent S are mixed at the indicated points. As the number of operations is increased horizontally, the terminal compositions E_1 and R_3 approach asymptotically the values obtained in continuous countercurrent extraction (*Treybal*, 1963, p. 360).



Figure 14.4. Less common examples of ternary equilibria and some temperature effects. (a) The system 2,2,4-trimethylpentane + nitroethane + perfluorobutylamine at 25°C; the Roman numerals designate the number of phases in that region [Vreeland and Dunlap, J. Phys. Chem. 61, 329 (1957)]. (b) Same as (a) but at 51.3°C. (c) Glycol + dodecanol+ nitroethane at 24°C; 12 different regions exist at 14°C [Francis, J. Phys. Chem. 60, 20 (1956)]. (d) Docosane + furfural + diphenylhexane at several temperatures [Varteressian and Fenske, Ind. Eng. Chem. 29, 270 (1937)]. (e) Fornic acid + benzene + tribromomethane at 70°C; the pair formic acid/benzene is partially miscible with 15 and 90% of the former at equilibrium at 25°C, 43 and 80% at 70°C, but completely miscible at some higher temperature. (f) Methylcyclohexane + water + -picoline at 20°C, exhibiting positive and negative tieline slopes; the horizontal tieline is called solutropic (Landolt-Börnstein II2b).



Figure 14.4—(continued)

Ishida, Bull. Chem. Soc. Jpn. 33, 693 (1960):

$$X_{\rm CS}X_{\rm SA}/X_{\rm CA}X_{\rm SS} = K(X_{\rm AS}X_{\rm SA}/X_{\rm AA}X_{\rm SS})^n.$$
(14.3)

Othmer and Tobias, 2nd. Eng. Chem. 34, 693 (1942):

$$(1 - X_{SS})/X_{SS} = K[(1 - X_{AA})/X_{AA}]^n.$$
(14.4)

Hand, J. Phys. Chem. 34, 1961 (1930):

$$X_{\rm CS}/X_{\rm SS} = K(X_{\rm CA}/X_{\rm AA})^n.$$
(14.5)

These equations should plot linearly on log-log coordinates; they are tested in Example 14.1.

A system of plotting both binodal and tieline data in terms of certain ratios of concentrations was devised by Janecke and is illustrated in Figure 14.1(c). It is analogous to the enthalpy-concentration or Merkel diagram that is useful in solving distillation problems. Straight line combining of mixture compositions is valid in this mode. Calculations for the transformation of data are made most conveniently from tabulated tieline data. Those for Figure 14.1 are made in Example 14.2. The x-y construction shown in





Figure 14.2 is the basis for a McCabe-Thiele construction for finding the number of extraction stages, as applied in Figure 14.7.

14.2. CALCULATION OF STAGE REQUIREMENTS

Although the most useful extraction process is with countercurrent flow in a multistage battery, other modes have some application. Calculations may be performed analytically or graphically. On flowsketches like those of Example 14.1 and elsewhere, a single box represents an extraction stage that may be made up of an individual mixer and separator. The performance of differential **contactors** such as packed or spray towers is commonly described as the height equivalent to a theoretical stage (HETS) in ft or m.

SINGLE STAGE EXTRACTION

The material balance is

feed + solvent = extract + raffinate,

$$F + S = E + R$$
. (14.6)

This nomenclature is shown with Example 14.3. On the triangular diagram, the proportions of feed and solvent locate the mix point



(c)

Figure 14.5. Representation of solvent extraction behavior in terms of certain properties rather than direct compositions [*Dunstan* et al., Sci. Pet., 1825-1855 (1938)]. (a) Behavior of a naphthenic distillate of VGC = 0.874 with nitrobenzene at 10°C. The viscosity-gravity constant is low for paraffins and high for naphthenes. (b) Behavior of a kerosene with 95% ethanol at 17°C. The aniline point is low for aromatics and naphthenes and high for paraffins. (c) Behavior of a dewaxed crude oil with liquid propane at 70°F, with composition expressed in terms of specific gravity.

M. The extract E and raffinate R are located on opposite ends of the tieline that goes through M.

IMMISCIBLE SOLVENTS

The distribution of a solute between two mutually immiscible solvents can be represented by the simple equation,

$$\mathbf{Y} = \mathbf{K}' \mathbf{X},\tag{14.7}$$

where

X = mass of solute/mass of diluent, Y = mass of solute/mass of solvent.

CROSSCURRENT EXTRACTION

In this process the feed and subsequently the raffinate are treated in successive stages with fresh solvent. The sketch is with Example 14.3. With a fixed overall amount of solvent the most efficient process is with equal solvent flow to each stage. The solution of Example 14.3 shows that crosscurrent two stage operation is superior to one stage with the same total amount of solvent.



Figure **14.6.** Construction of points on the distribution and operating curves: Line *ab* is a **tieline**. The dashed line is the **tieline** locus. Point e is on the equilibrium distribution curve, obtained as the intersection of paths *be* and *ade*. Line Pfg is a random line from the difference point *P* and intersecting the binodal curve in *f* and g. Point j is on the operating curve, obtained as the intersection of paths g and *fhj*.

When K' is not truly constant, some kind of mean value may be applicable, for instance, a geometric mean, or the performance of the extraction battery may be calculated stage by stage with a different value of K' for each. The material balance around the first stage where the raffinate leaves and the feed enters and an intermediate stage k (as in Fig. 14.8, for instance) is

$$EY_{F} + RX_{k-1} = EY_{k} + RX_{k}.$$
 (14.8)

In terms of the extraction ratio,

$$A = K(E/R), \tag{14.9}$$

EXAMPLE 14.1

The Equations for Tieline Data

The **tieline** data of the system of Example 14.1 are plotted according to the groups of variables in the equations of Ishida, Hand, and Othmer and **Tobias** with these results:

Ishida: $y = 1.00x^{0.67}$ [Eq. (14.3)],	
Hand: $y = 0.078x^{1.11}$ [Eq. (14.5)],	
Othmer and Tobias: $y = 0.88x^{0.90}$ [Eq.	(14.4)]



The last correlation is inferior for this particular example as the plots show.

X _{AA}	XCA	X _{SA}	XAS	x _{cs}	x _{ss}
98.945	0.0	1.055	5.615	0.0	94. 385
92.197	6.471	1.332	5.811	3.875	90. 313
63. 572	14.612	1.816	6.354	9.756	83. 889
75.356	22. 277	2.367	7.131	15.365	77.504
68.283	26.376	3. 341	8.376	20.686	70. 939
60.771	34. 345	4.884	9. 545	26.248	64.207
54.034	39. 239	6.727	11.375	31.230	57.394
47.748	42.649	9.403	13. 505	35.020	51.475
39. 225	45. 594	15. 181	18. 134	39.073	42.793

X _{CS} X _{SA} X _{CA} X _{SS}	1 X _{AA} 1	1 1 x_{ss}	x _{ca} /x _{aa}	x _{cs} /x _{ss}
0	0.0107	0.0595	0	0
0.0088	0.0846	0. 1073	0.070	0.043
0.0129	0. 1966	0. 1928	0.178	0.116
0. 0211	0. 3270	0. 2903	0.296	0.198
0.0343	0.4645	0.4097	0.416	0. 292
0.0581	0.6455	0.5575	0.565	0.409
0.0933	0.8507	0.7423	0.726	0.544
0.1493	1.0943	0.9427	0.897	0.680
0. 3040	1. 5494	1.3368	1.162	0. 913
	x _{CS} x _{SA} x _{CA} x _{SS} 0 0.0088 0.0129 0.0211 0.0343 0.0581 0.0933 0.1493 0.3040	X _{CS} X _{SA} X _{AA} 0 0.0107 0.0088 0.0846 0.0129 0.1966 0.0211 0.3270 0.0343 0.4645 0.0581 0.6455 0.0933 0.8507 0.1493 1.0943 0.3040 1.5494	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

EXAMPLE 14.2 Tabulated Tieline and Distribution Data for the System A = 1-Hexene, B = Tetramethylene Sulfone, C = Benzene, Represented in Figure 14.1

Experimental tienne data in mor 7	experimental	tieline da	ta in mol	8
-----------------------------------	--------------	------------	-----------	---

Left Phase			Right Phase			
Α	C	В	Α	с	В	
98.945	0.0	1.055	5.615	0.0	94. 385	
92. 197	6.471	1.332	5.811	3.875	90. 313	
83. 572	14.612	1.816	6.354	9.758	83. 888	
75. 356	22. 277	2.367	7.131	15.365	77.504	
68. 283	28.376	3. 341	8.376	20.686	70. 938	
60. 771	34. 345	4.884	9.545	26.248	64.207	
54.034	39. 239	6.727	11.375	31.230	57.394	
47. 748	42.849	9.403	13. 505	35.020	51.475	
39. 225	45. 594	15. 181	18.134	39.073	42.793	

the material balance becomes

А

R_N

$$(A/K)Y_F + X_{k-1} = AX_k + X_n. (14.10)$$

When these balances are made stage-by-stage and intermediate compositions are eliminated, assuming constant A throughout, the result relates the terminal compositions and the number of stages. The expression for the fraction extracted is

$$\phi = \frac{X_F - X_n}{X_F - Y_S/K} = \frac{A^{n+1} - A}{A^{n+1} - 1}.$$
(14.11)

This is of the same form as the Kremser-Brown equation for gas **absorption** and stripping and the Turner equation for leaching. The

Calculated ratios for the Jänecke coordinate plot of Figure 14.1:

Left	Phase	Right	Phase
B A + C	C A+C	B A + C	c A+C
0. 0108	0	16. 809	0
0.0135	0.0656	9.932	0.4000
0.0185	0.1488	5. 190	0.6041
0.0248	0. 2329	3. 445	0.6830
0.0346	0. 2936	2.441	0.7118
0.0513	0.3625	1.794	0. 7333
0.0721	0. 4207	1.347	0. 7330
0. 1038	0.4730	1.061	0.7217
0.1790	0. 5375	0.748	0.6830

The x-y plot like that of Figure 14.6 may be made with the **tieline** data of columns **5** and 2 expressed as fractions or by projection from the triangular diagram as shown.

solution for the number of stages is

$$n = -1 + \frac{\ln[(A - \phi)/(1 - \phi)]}{\ln A}$$
(14.12)

When A is the only unknown, it may be found by trial solution of these equations, or the Kremser-Brown stripping chart may be used. Example 14.4 applies these results.

14.3. COUNTERCURRENT OPERATION

In countercurrent operation of several stages in series, feed enters the first stage and final extract leaves it, and fresh solvent enters the last stage and final raffinate leaves it. Several representations of



(a)

Figure 14.7. Locations of operating points P and Q for feasible, total, and minimum extract reflux on triangular diagrams, and stage requirements determined on rectangular distribution diagrams. (a) Stages required with feasible extract reflux. (b) Operation at total reflw and minimum number of stages. (c) Operation at minimum reflux and infinite stages.





Figure 14.7—(continued)

such processes are in Figure 14.3. A flowsketch of the process together with nomenclature is shown with Example 14.5. The overall material balance is

$$F + S = E_1 + R_N = M \tag{14.13}$$

or

$$F - E_1 = R_N - S = P. (14.14)$$

The intersection of extended lines *FE*, and $R_N S$ locates the operating point *P*. The material balance from stage 1 through *k* is

$$F + E_{k+1} = E_1 + R_k \tag{14.15}$$

or

$$F - E_1 = R_k - E_{k+1} = P. (14.16)$$

Accordingly, the raffinate from a particular stage and the extract from a succeeding one are on a line through the operating point *P*. Raffinate R_k and extract E_k streams from the same stage are located at opposite ends of the same **tieline**.



(b)

(c)

The operation of finding the number of stages consists of a number of steps:

^xCR_N ... x_C in raffinate ×CR_o

- 1. Either the solvent feed ratio or the compositions E_1 and R_N serve to locate the mix point *M*.
- 2. The operating point *P* is located as the intersection of lines *FE*, and $R_N S$.
- 3. When starting with E_1 , the raffinate R_1 is located at the other end of the tieline.
- 4. The line PR, is drawn to intersect the binodal curve in E,.

The process is continued with the succeeding values R_2 , E_3 , R_3 , E_4 ,... until the final raffinate composition is reached.

When number of stages and only one of the terminal compositions are fixed, the other terminal composition is selected by trial until the **stepwise** calculation finds the prescribed number of stages. Example 14.6 applies this kind of calculation to find the stage requirements for systems with Types I and II equilibria.

Evaluation of the numbers of stages also can be made on rectangular distribution diagrams, with a McCabe-Thiele kind of construction. Example 14.5 does this. The Janecke coordinate plots like those of Figures 14.1 and 14.2 also are convenient when many stages are needed, since then the triangular construction may

Single Stage and Cross Current Extraction of Acetic Acid from Methylisobutyl Ketone with Water

The original mixture contains 35% acetic acid and 65% MIBK. It is charged at 100 kg/hr and extracted with water.

a. In a single stage extractor water is mixed in at 100 kg/hr. On the triangular diagram, mix point M is midway between F and S. Extract and raffinate compositions are on the **tieline** through M. Results read off the diagram and calculated with material





balance are

E	R
0.185	0.16
0.035	0.751
0.78	0.089
120	80
	E 0.185 0.035 0.78 120

b. The flowsketch of the crosscurrent process is shown. Feed to the first stage and water to both stages are at 100 kg/hr. The extract and raffinate compositions are on the **tielines** passing through mix points M_1 and M_2 . Point M is for one stage with the same total amount of solvent. Two stage results are:

	4	R ₁	4	R ₂
Acetic acid	0.185	0.160	0.077	0.058
MIBK	0.035	0.751	0.018	0.902
Water	0.780	0.089	0.905	0.040
kg/hr	120	80	113.4	66.6





become crowded and difficult to execute accurately unless a very large scale is adopted. The Janecke method was developed by Maloney and Schubert [Trans. *AIChE* 36, 741 (1940)]. Several detailed examples of this kind of calculation are worked by Treybal (1963), Oliver (*Diffusional Separation Processes*, Wiley, New York, 1966), and Laddha and Degaleesan (1978).

MINIMUM SOLVENT/FEED RATIO

Both maximum and minimum limits exist of the solvent/feed ratio. The maximum is the value that locates the mix point M on the binodal curve near the solvent vertex, such as point M_{max} on Figure 14.7(b). When an operating line coincides with a tieline, the number of stages will be infinite and will correspond to the minimum solvent/feed ratio. The pinch point is determined by the intersection of some tieline with line $R_N S$. Depending on whether the slopes of the tielines are negative or positive, the intersection that is closest or farthest from the solvent vertex locates the operating point for minimum solvent. Figure 14.9 shows the two

cases. Frequently, the ticline through the feed point determines the minimum solvent quantity, but not for the two cases shown.

EXTRACT REFLUX

Normally, the concentration of solute in the final extract is limited to the value in equilibrium with the feed, but a countercurrent stream that is richer than the feed is available for enrichment of the extract. This is essentially solvent-free extract as reflux. A flowsketch and nomenclature of such a process are given with Example 14.7. Now there are two operating points, one for above the feed and one for below. These points are located by the following procedure:

- 1. The mix point is located by establishing the solvent/feed ratio.
- 2. Point Q is at the intersection of lines $R_N M$ and $E_1 S_E$, where S_E refers to the solvent that is removed from the final extract, and may or may not be of the same composition as the fresh solvent S. Depending on the shape of the curve, point Q may be inside



Figure 14.8. Model for liquid-liquid extraction. Subscript *i* refers to a component: i = 1, 2, ..., c. In the commonest case, F_1 is the only feed stream and F_N is the solvent, or F_k may be a reflux stream. Withdrawal streams U_k can be provided at any stage; they are not incorporated in the material balances written here.

the binodal curve as in Example 14.7, or outside as in Figure 14.7.

3. Point *P* is at the intersection of lines $R_N M$ and $E_1 S_E$, where S_E refers to the solvent removed from the extract and may or may not be the same composition as the fresh solvent *S*.

Determination of the stages uses Q as the operating point until the raffinate composition R_k falls below line FQ. Then the operation is continued with operating point P until R_N is reached.

MINIMUM REFLUX

For a given extract composition E_1 , a pinch point develops when an operating line through either P or Q coincides with a tieline. Frequently, the tieline that passes through the feed point F determines the reflux ratio, but not on Figure 14.7(c). The tieline that intersects line FS_E nearest point S_e locates the operating point Q_m for minimum reflux. In Figure 14.7(c), intersection with tieline *Fcde* is further away from point S_E than that with tieline abQ_m , which is the one that locates the operating point for minimum reflux in this case.

MINIMUM STAGES

As the solvent/feed ratio is increased, the mix point *M* approaches the solvent point *S*, and poles *P* and *Q* likewise do so. At total reflux all of the points *P*, *Q*, *S*, *S*_{*E*}, and *M* coincide; this is shown in Figure 14.7(b).

Examples of triangular and McCabe-Thiele constructions for feasible, total, and minimum reflux are shown in Figure 14.7.

EXAMPLE 14.4

Extraction with an Immiscible Solvent

A feed containing 30wt % of propionic acid and 70wt % trichlorethylene is to be extracted with water. Equilibrium distribution of the acid between water (Y) and TCE (X) is represented by Y = K'X, with K' = 0.38. Section 14.3 is used.

a. The ratio *E/R* of water to TCE needed to recover 95% of the acid in four countercurrent stages will be found:

$$\begin{split} X_F &= 30/70, \\ X_n &= 1.5170, \\ Y_S &= 0, \\ \phi &= (30 - 1.5)/(30 - 0) = 0.95 = (A^5 - A/(A^5 - 1)). \end{split}$$

By trial,

$$A = 1.734,$$

 $E/R = A/K' = 1.734/0.38 = 4.563.$

b. The number of stages needed to recover 95% of the acid with E/R = 3.5 is found with Eq. 14.12.

$$A = K'E/R = 0.38(3.5) = 1.330, \quad \phi = 0.95,$$

$$n = -1 + \frac{\ln[(A - \phi)/(1 - \phi)]}{\ln A} = -1$$

$$+ \ln[(1.330 - 0.95)/(1 - 0.95)]/\ln(1.330) = 6.11$$



Countercurrent Extraction Represented on Triangular and Rectangular Distribution Diagrams

The specified feed F and the desired extract E_1 and raffinate R_N compositions are shown. The solvent/feed ratio is in the ratio of the line segments MS/MF, where the location of point M is shown as the intersection of lines E_1R_N and FS.

Phase equilibrium is represented by the **ticline** locus. The equilibrium distribution curve is constructed as the locus of intersections of horizontal lines drawn from the right-hand end of a

tieline with horizontals from the left-hand end of the tielines and reflected from the 45" line.

The operating curve is drawn similarly with horizontal projections from pairs of random points of intersection of the binodal curve by lines drawn through the difference point P. Construction of these curves also is explained with Figure 14.6.

The rectangular construction shows that slightly less than eight stages are needed and the triangular that slightly more than eight are needed. A larger scale and greater care in construction could bring these results closer together.



Naturally, the latter constructions are analogous to those for distillation since their forms of equilibrium and material balances are the same. References to the literature where similar calculations are performed with Janecke coordinates were given earlier in this section.

Use of **reflux** is most effective with Type II systems since then essentially pure products on a solvent-free basis can be **made**. In contrast to distillation, however, extraction with reflux rarely is beneficial, and few if any practical examples are known. A related kind of process employs a second solvent to wash the extract countercurrently. The requirements for this solvent are that it be only slighly soluble in the extract and easily removable from the extract and raffinate. The sulfolane process is of this type; it is described, for example, by Treybal (1980) and in more detail by Lo et al. (1983, pp. 541-545).

14.4. LEACHING OF SOLIDS

Leaching is the removal of solutes from admixture with a solid by contracting it with a solvent. The solution phase sometimes is called the overflow, but here it will be called extract. The term underflow or raffinate is applied to the solid phase plus its entrained or occluded solution.

Equilibrium relations in leaching usually are simpler than in liquid-liquid equilibria, or perhaps only appear so because few measurements have been published. The solution phase normally contains no entrained solids so its composition appears on the hypotenuse of a triangular diagram like that of Example 14.8. Data for the raffinate phase may be measured as the holdup of solution by the solid, K lb solution/lb dry (oil-free) solid, as a function of the concentration of the solution, y lb oil/lb solution. The correspond-

Stage Requirements for the Separation of a Type I and a Type II System

a. The system with A = heptane, B = tetramethylene sulfone, and C= toluene at 50°C [Triparthi, Ram, and Bhimeshwara, J. Chem. Eng. Data 20, 261 (1975)]: The feed contains 40% C, the extract 70% C on a TMS-free basis or 60% overall, and raffinate 5% C. The construction shows that slightly more than two equilibrium stages are needed for this separation. The compositions of the streams are read off the diagram:

	Feed	Extract	Raffinat
Heptane	60	27	2
TMS	0	13	93
Toluene	40	60	5

The material balance on heptane is

$$40 = 0.6E + 0.05(100 - E)$$

whence E = 63.6 lb/100 lb feed, and the TMS/feed ratio is

$$0.13(63.6) + 0.93(36.4) = 42$$
 lb/100 lb feed.

b. The type II system with A= octane, B = nitroethane, and C = 2,2,4-trimethylpentane at 25°C [Hwa, Techo, and Ziegler, J. Chem. Eng. Data 8, 409 (1963)]: The feed contains 40% TMP, the extract 60% TMP, and the raffinate 5% TMP. Again, slightly more than two stages are adequate.



ing weight fraction of oil in the raffinate or underflow is

$$x = K_V / (K + 1). \tag{14.17}$$

Since the raffinate is a mixture of the solution and dry solid, the equilibrium value in the raffinate is on the line connecting the origin

with the corresponding solution composition y, at the value of xgiven by Eq. (14.17). Such a raffinate line is constructed in Example 14.8.

Material balance in countercurrent leaching still is represented by Eqs. (14.14) and (14.16). Compositions R_k and E_{k+1} are on a line through the operating point P, which is at the intersection of



Figure 14.9. Minimum solvent amount and maximum extract concentration. Determined by location of the intersection of extended tielines with extended line $R_N S$. (a) When the tielines slope down to the left, the furthest intersection is the correct one. (b) When the tielines slope down to the right, the nearest intersection is the correct one. At maximum solvent amount, the mix point M_m is on the binodal curve.

Countercurrent Extraction Employing Extract Reflux The feed F, extract E_1 , and raffinate R_N are located on the triangular diagram. The ratio of solvent/feed is specified by the location of the point M on line SF.

Other nomenclature is identified on the flowsketch. The solvent-free reflux point R_0 is located on the extension of line SE,. Operating point Q is located at the intersection of lines SR_0 and $R_N M$. Lines through Q intersect the binodal curve in compositions of raffinate and reflux related by material balance: for instance, R_n and E_{n+1} . When the line QF is crossed, further constructions are

made with operating point P, which is the intersection of lines FQand SR_N.

In this example, only one stage is needed above the feed F and five to six stages below the feed. The ratio of solvent to feed is

$$S/F = FM/MS = 0.196$$

and the external reflux ratio is



EXAMPLE 14.8

Leaching of an Oil-Bearing Solid in a Countercurrent Battery Oil is to be leached from granulated halibut livers with pure ether as solvent. Content of oil in the feed is 0.32 lb/lb dry (oil-free) solids and 95% is to be recovered. The economic upper limit to extract concentration is 70% oil. Ravenscroft [Ind. Eng. Chem. 28, 851 (1934)] measured the relation between the concentration of oil in the solution, y_1 and the entrainment or occlusion of solution by the solid phase, K lb solution/lb dry solid, which is represented by the equation

 $K = 0.19 + 0.126y + 0.810y^2.$

The oil content in the entrained solution then is given by

$$\mathbf{x} = \mathbf{K}/(\mathbf{K} + 1)\mathbf{y}$$
, wt fraction,

and some calculated values are

Y	0	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.6
x	0	0.0174	0.0397	0.0694	0.1080	0.1565	0.2147	0. 2821	0.3578

Points on the raffinate line of the triangular diagram are located on lines connecting values of y on the hypotenuse (solids-free) with the origin, at the values of \mathbf{x} and corresponding y from the preceding tabulation.

Feed composition is $x_F = 0.32/1.32 = 0.2424$. Oil content of extract is $y_1 = 0.7$. Oil content of solvent is $y_s = 0$. Amount of oil in the raffinate is 0.32(0.05) = 0.016 lb/lb dry, and the corresponding entrainment ratio is

$$K_N = 0.016/y_N = 0.19 + 0.126y_N + 0.81y_N^2$$

EXAMPLE 14.8—(continued)

Solving by trial,

 $y_N = 0.0781,$ $K_N = 0.2049,$ $x_N = 0.0133$ (final raffinate composition).

The operating point P is at the intersection of lines FE, and SR,. The triangular diagram construction shows that six stages are needed.

The equilibrium line of the rectangular diagram is constructed with the preceding tabulation. Points on the material balance line are located as intersections of random lines through P with these results:

Y	0	0.1	0.2	0.3	0.4	0.5	0.6	0.7
x	0.013	0.043	0.079	0.120	0. 171	0. 229	0. 295	0.366

The McCabe-Thiele construction also shows that six stages are needed.

Point P is at the intersection of lines E_1F and SR, Equilibrium compositions are related on lines through the origin, point A. Material balance compositions are related on lines through the operating point P.



lines *FE*, and *SR*_N. Similarly, equilibrium compositions R_k and E_k are on a line through the origin. Example 14.8 evaluates stage requirements with both triangular diagram and McCabe-Thiele constructions. The mode of construction of the McCabe-Thiele diagram is described there.

These calculations are of equilibrium stages. The assumption is made that the oil retained by the solids appears only as entrained solution of the same composition as the bulk of the liquid phase. In some cases the solute may be adsorbed or retained within the interstices of the solid as solution of different concentrations. Such deviations from the kind of equilibrium assumed will result in stage efficiencies less than 100% and must be found experimentally.

14.5. NUMERICAL CALCULATION OF MULTICOMPONENT EXTRACTION

Extraction calculations involving more than three components cannot be done graphically but must be done by numerical solution of equations representing the phase equilibria and material balances over all the stages. Since extraction processes usually are adiabatic and nearly isothermal, enthalpy balances need not be made. The solution of the resulting set of equations and of the prior determination of the parameters of activity coefficient correlations requires computer implementation. Once such programs have been developed, they also may be advantageous for ternary extractions, particularly when the number of stages is large or several cases must be worked out. Ternary graphical calculations also could be done on a computer screen with a little effort and some available software.

The notation to be used in making material balances is shown on Figure 14.8. For generality, a feed stream F_k is shown at every stage, and a withdrawal stream U_k also could be shown but is not incorporated in the balances written here. The first of the double subscripts identifies the component *i* and the second the stage number *k*; a single subscript refers to a stage.

For each component, the condition of equilibrium is that its activity is the same in every phase in contact. In terms of activity coefficients and concentrations, this condition on stage k is written:

$$\gamma_{ik}^{E} y_{ik} = \gamma_{ik}^{R} x_{ik} \tag{14.18}$$

or

$$y_{ik} = K_{ik} x_{ik}, \tag{14.19}$$

where

$$K_{ik} = \gamma^R_{ik} / \gamma^E_{ik} \tag{14.20}$$

is the distribution ratio. The activity coefficients are functions of the temperature and the composition of their respective phases:

$$Y_{ik}^{E} = f(T_k, y_{1k}, y_{2k}, \dots, y_{ck}),$$
(14.21)

$$\gamma_{ik}^{\kappa} = f(T_k, x_{1k}, x_{2k}, \dots, x_{ck}).$$
 (14.22)

The most useful relations of this type are the NRTL and UNIQUAC which are shown in Table 14.1.

Around the kth stage, the material balance is

$$R_{k-1}x_{i,k+1} + E_{k+1}y_{i,k+1} + F_k z_{ik} - R_k x_{ik} - E_k y_{ik} = 0.$$
(14.23)

When combined with Eq. (14.19), the material balance becomes

$$R_{k-1}x_{i,k-1} - (R_k + E_k K_{ik})x_{ik} + E_{k+1}K_{i,k+1}x_{i,k+1} = -F_k z_{ik}.$$
(14.24)

In the top stage, k = 1 and $R_0 = 0$ so that

$$-(R_1 + V_1K_{i1})x_{i1} + E_2K_{i2}x_{i2} = -F_1z_{i1}.$$
(14.25)

In the bottom stage, k = N and $E_{N+1} = 0$ so that

$$R_{N-1}x_{i,N-1} - (R, + E_N K_{iN})x_{iN} = -F_N z_{iN}.$$
(14.26)

The overall balance from stage 1 through stage k is

$$R_{k} = E_{k+1} - E_{1} + \sum_{1}^{k} F_{k}, \qquad (14.27)$$

which is used to find raffinate flows when values of the extract flows have been estimated.

For all stages for a component *i*, Eqs. (14.24)-(14.26) constitute a tridiagonal matrix which is written

$$\begin{bmatrix} B_{1} & C_{1} & & & \\ A_{2} & B_{2} & C_{2} & & \\ & A_{j} & B_{j} & C_{j} & & \\ & & A_{N-1} & B_{N-1} & C_{N-1} \\ & & & & A_{N} & B_{N} \end{bmatrix} \begin{bmatrix} x_{i1} \\ x_{i2} \\ x_{ij} \\ x_{iN-1} \\ x_{iN} \end{bmatrix} = \begin{bmatrix} D_{1} \\ D_{2} \\ D_{j} \\ D_{N-1} \\ D_{N} \end{bmatrix}$$
(14.28)

When all of the coefficients are known, this can be solved for the concentrations of component i in every stage. A straightforward method for solving a tridiagonal matrix is known as the Thomas algorithm to which references are made in Sec. 13.10, "Basis for Computer Evaluation of Multicomponent Separations: Specifications."

INITIAL ESTIMATES

Solution of the equations is a process in which the coefficients of Eq. (14.28) are iteratively improved. To start, estimates must be made of the flow rates of all components in every stage. One procedure is to assume complete removal of a "light" key into the extract and of the "heavy" key into the raffinate, and to keep the solvent in the extract phase throughout the system. The distribution of the keys in the intermediate stages is assumed to vary linearly, and they must be made consistent with the overall balance, Eq. (14.27), for each component. With these estimated flowrates, the values of x_{ik} and y_{ik} are evaluated and may be used to find the activity coefficients and distribution ratios, K_{ik} . This procedure is used in Example 14.9.

PROCEDURE

The iterative calculation procedure is outlined in Figure 14.10. The method is an adaptation to extraction by Tsuboka and Katayama (1976) of the distillation calculation procedure of Wang and Henke [*Hydrocarb. Proc.* **45**(8), 1.55-163 (1967)]. It is also presented by Henley and Seader (1981, pp. 586-594).

- 1. The initial values of the flowrates and compositions x_{ik} and y_{ik} are estimated as explained earlier.
- The values of activity coefficients and distribution ratios are evaluated.
- 3. The coefficients in the tridiagonal matrix are evaluated from Eqs. (14.24)-(14.26). The matrix is solved once for each component.
- 4. The computed values of iteration (r + 1) are compared with those of the preceding iteration as

$$\tau_1 = \sum_{i=1}^{C} \sum_{k=1}^{N} |x_{ik}^{(r+1)} - x_{ik}^{(r)}| \le \varepsilon_1 = 0.01 NC.$$
(14.29)

The magnitude, 0.01NC, of the convergence criterion is arbitrary.

5. For succeeding evaluations of activity coefficients, the values of the mol fractions are normalized as

$$(x_{ik})_{\text{normalized}} = x_{ik} \Big/ \sum_{i=1}^{C} x_{ik},$$

$$(y_{ik})_{\text{normalized}} = y_{ik} \Big/ \sum_{i=1}^{C} y_{ik}.$$
(14.30)

6. When the values of x_{ik} have converged, a new set of y_{ik} is calculated with

$$\mathbf{y}_{ik} = \mathbf{K}_{ik} \mathbf{x}_{ik}.\tag{14.19}$$

7. A new set of extract flow rates is calculated from

$$E_{k}^{(s+1)} = E_{k}^{(s)} \sum_{i=1}^{C} y_{ik}, \qquad (14.31)$$

where *s* is the outer loop index number.

TABLE 14.1. NRTL and UNIQUAC Correlations for Activity Coefficients of **Three**-Component **Mixtures**^a

 $\begin{aligned} \overline{NRTL} \\ \ln \gamma_i &= \frac{\tau_{1i}G_{1i}x_1 + \tau_{2i}G_{2i}x_2 + \tau_{3i}G_{3i}x_3}{G_{1i}x_1 + G_{2i}x_2 + G_{3i}x_3} \\ &+ \frac{x_1G_{i1}}{x_1 + G_{12}x_2 + G_{13}x_3} \left[\tau_{i1} - \frac{x_2\tau_{21}G_{21} + x_3\tau_{31}G_{31}}{x_1 + x_2G_{21} + x_3G_{31}} \right] \\ &+ \frac{x_2G_{i2}}{G_{12}x_1 + x_2 + G_{32}x_3} \left[\tau_{i2} - \frac{x_1\tau_{12}G_{12} + x_3\tau_{32}G_{32}}{x_1G_{12} + x_2 + x_3G_{32}} \right] \\ &+ \frac{x_3G_{i3}}{G_{13}x_1 + G_{23}x_2 + x_3} \left[\tau_{i3} - \frac{x_1\tau_{13}G_{13} + x_2\tau_{23}G_{23}}{G_{13}x_1 + G_{23}x_2 + x_3} \right] \\ &\tau_{ii} = 0 \\ G_{ii} = 1 \end{aligned}$

UNIQUAC

$$\ln \gamma_{i} = \ln \frac{\phi_{i}}{x_{i}} + 5q_{i} \ln \frac{\theta_{i}}{\phi_{i}} + l_{i} - \frac{\phi_{i}}{x_{i}} (x_{1}l_{1} + x_{2}l_{2} + x_{3}l_{3}) + q_{i}[1 - \ln(\theta_{1}\tau_{1i} + \theta_{2}\tau_{2i} + \theta_{3}\tau_{3i})]$$

$$\frac{\theta_{1}\tau_{i1}}{\theta_{1} + \theta_{2}\tau_{21} + \theta_{3}\tau_{31}} - \frac{\theta_{2}\tau_{i2}}{\theta_{1}\tau_{12} + \theta_{2} + \theta_{3}\tau_{32}} - \frac{\theta_{3}\tau_{i3}}{\theta_{1}\tau_{13} + \theta_{2}\tau_{23} + \theta_{3}}$$

$$\tau_{ii} = 1$$

$$\phi_i = \frac{r_i x_i}{r_1 x_1 + r_2 x_2 + r_3 x_3}$$

$$\theta_i = \frac{q_i x_i}{q_1 x_1 + 42 x_2 + 43 x_3}$$

$$l_i = 5(r_i - q_i) - r_i + 1$$

⁸ NRTL equation: There is a pair of parameters g_{jk} and g_{kj} for each pair of substances in the mixture; for three substances, there are three pairs. The other terms of the equations are related to the basic ones by

 $\begin{aligned} \tau_{jk} &= g_{jk} / RT, \\ G_{jk} &= \exp(-\alpha_{jk} \tau_{jk}). \end{aligned}$

For liquid-liquid systems usually, $\alpha_{ik} = 0.4$.

UNIQUAC equation: There is a pair of parameters u_{jk} and u_{kj} for each pair of substances in the mixture:

 $\tau_{ik} = \exp(-u_{ik}/RT).$

The terms with single subscripts are properties of the pure materials which are usually known or can be estimated.

The equations are extended readily to more components. (See, for example, Walas, Phase Equilibria in Chemical Engineering, **Butterworths**, 1985).

8. The criterion for convergence is

$$\tau_2 = \sum_{k=1}^{N} (I - E_k^{(s)} / E_k^{(s+1)})^2 \le \varepsilon_2 = 0.01N.$$
(14.32)

The magnitude, **0.01***N*, of the convergence criterion is arbitrary.

- 9. If convergence has not been attained, new values of R_k are calculated from Eq. (14.27).
- 10. Distribution ratios K_{ik} are based on normalized values of x_{ik} and y_{ik} .
- 11. The iteration process continues through the inner and outer loops.

EXAMPLE 14.9

Trial Estimates and Converged Flow Bates and Compositions in All Stages of an Extraction Battery for a Four-Component Mixture

Benzene is to be recovered from a mixture with hexane using aqueous dimethylformamide as solvent in a five-stage extraction battery. Trial estimates of flow rates for starting a numerical solution are made by first assuming that all of the benzene and all of the solvent ultimately appear in the extract and all of the hexane appears in the raffinate. Then **flow** rates throughout the battery are assumed to vary linearly with stage number. Table 1 shows these estimated flowrates and Table 2 shows the corresponding mol fractions. Tables 3 and 4 shows the converged solution made by Henley and Seader (1981, p. 592); they do not give any details of the solution but the algorithm of Figure 14.10 was followed.

TABLE 1. Estimated mol/hr

Extract				Raffinate						
Stage	Total	Н	B	D	W	Total	Η	B	D	W
0	_									
1	1100	0	100	750	250	400	300	100	0	0
2	1080	0	80	750	250	380	300	80	0	0
3	1060	0	60	750	250	360	300	60	0	0
4	1040	0	40	750	250	340	300	40	0	0
5	1020	0	20	750	250	320	300	20	0	0
N + 1	1000	0	0	750	250	300	300	0	0	0

Solutions of four cases of three- and four-component systems are presented by Tsuboka and Katayama (1976); the number of outer loop iterations ranged from 7 to 41. The four component case worked out by Henley and Seader (1981) is summarized in Example 14.9; they solved two cases with different water contents of the solvent, dimethylformamide.

14.6. EQUIPMENT FOR EXTRACTION

Equipment for extraction and leaching must be capable of providing intimate contact between two phases so as to effect transfer of solute between them and also of ultimately effecting a complete separation of the phases. For so general an operation, naturally a substantial variety of equipment has been devised. A very general classification of equipment, their main characteristics and industrial applications is in Table 14.2. A detailed table of comparisons and ratings of 20 kinds of equipment on 14 characteristics has been prepared by Pratt and Hanson (in Lo et al., 1983, **p.** 476). Some comparisons of required sizes and costs are in Table 14.3.

Selected examples of the main categories of extractors are represented in Figures 14.11-14.15. Their capacities and performance will be described in general terms insofar as possible, but sizing of liquid-liquid extraction equipment always requires some pilot plant data or acquaintance with analogous cases. Little detailed information about such analogous situations appears in the open literature. Engineers familiar with particular kinds of equipment, such as their manufacturers, usually can predict performance with a minimum amount of pilot plant data.

Stone	_		Yıj	x _{ij}			
Stage j	н	В	D	W	н	В	DW
1	0.0	0. 0909	0. 6818	0. 2273	0. 7895	0. 2105	0.0 0.0
2	0.0	0.0741	0.6944	0.2315	0.8333	0.1667	0.0 0.0
3	0.0	0.0566	0.7076	0. 2359	0.8824	0.1176	0.0 0.0
4	0.0	0. 0385	0.7211	0.2404	0.9375	0.0625	0.0 0.0
5	0.0	0. 0196	0. 7353	0. 2451	1.0000	0.0	0.0 0.0

TABLE 3. Converged Mol Fractions

Stago		y,	I		× _{ij}			
j	н	В	D	W	н	В	D	W
1	0. 0263	0. 0866	0. 6626	0. 2245	0. 7586	0. 1628	0. 0777	0.0009
2	0. 0238	0.0545	0.6952	0. 2265	0.8326	0.1035	0.0633	0.0006
3	0. 0213	0.0309	0.7131	0. 2347	0.8858	0.0606	0.0532	0.0004
4	0.0198	0.0157	0.7246	0. 2399	0. 9211	0.0315	0.0471	0.0003
5	0.0190	0.0062	0.7316	0.2432	0.9438	0.0125	0.0434	0.0003

TABLE 4. Converged mol/hr

	Extract	Raffinate
Hexane	29.3	270. 7
Benzene	96.4	3.6
DMF	737.5	12.5
Water	249.0	0.1
Total	1113. 1	286.9

Literature data is almost entirely for small equipment whose capacity and efficiency cannot be scaled up to commercial sizes, although it is of qualitative value. Extraction processes are sensitive because they operate with small density differences that are sensitive to temperature and the amount of solute transfer. They also are affected by interfacial tensions, the large changes in phase flow rates that commonly occur, and even by the direction of mass transfer. For comparison, none of these factors is of major significance in vapor-liquid contacting.

CHOICE OF DISPERSE PHASE

Customarily the phase with the highest volumetric rate is dispersed since a larger interfacial area results in this way with a given droplet size. In equipment that is subject to backmixing, such as spray and packed towers but not sieve tray towers, the disperse phase is made the one with the smaller volumetric rate. When a substantial difference in resistances of extract and raffinate films to mass transfer exists, the high phase resistance should be compensated for with increased surface by dispersion. From this point of view, Laddha and Degaleesan (1978, pp. 194) point out that water should be the dispersed phase in the system water + diethylamine+ toluene. The dispersed phase should be the one that wets the material of construction less well. Since the holdup of continuous phase usually is greater, the phase that is less hazardous or less expensive should be continuous. It is best usually to disperse a highly viscous phase.



Figure 14.10. Algorithm for computing flows and compositions in an extraction battery of a specified number of stages (after *Henley and* Seader, 1981).

MIXER-SETTLERS

The original and in concept the simplest way of accomplishing extractions is to mix the two phases thoroughly in one vessel and then to allow the phases to separate in another vessel. A series of such operations performed with series or countercurrent flows of the phases can accomplish any desired degree of separation. Mixer-settlers have several advantages and disadvantages, for instance:

Pros. The stages are independent, can be added to or removed as needed, are easy to start up and shut down, are not bothered by suspended solids, and can be sized for high (normally 80%) efficiencies.

Cons. Emulsions can be formed by severe mixing which are hard to break up, pumping of one or both phases between tanks may be required, independent agitation equipment and large floor space needs are expensive, and high holdup of valuable or hazardous solvents exists particularly in the settlers.

Some examples of more or less compact arrangements of mixers and settlers are in Figures 14.11 and 14.14(c). Mixing equipment is described in Chapter 10 where rules for sizing, blending, mixing intensity, and power requirements are covered, for instance Figure 10.3 for blend times in stirred tanks. Mixing with impellers in tanks is most common, but also is accomplished with pumps, jet mixers [Fig. 14.11(b)], line mixers and static mixers.

TABLE 14.2. Features and Industrial Applications of Liquid-Liquid Extractors

Types of extractor	General features	Fields of industrial application
Unagitated columns	Low capital cost Low operating and maintenance cost Simplicity in construction Handles corrosive material	Petrochemical Chemical
Mixer-settlers	High-stage efficiency Handles wide solvent ratios High capacity Good flexibility Reliable scale-up Handles liquids with high viscosity	Petrochemical Nuclear Fertilizer Metallurgical
Pulsed columns	Low HETS No internal moving parts Many stages possible	Nuclear Petrochemical Metallurgical
Rotary-agitation columns	Reasonable capacity Reasonable HETS Many stages possible Reasonable construction cost Low operating and maintenance cost	Petrochemical Metallurgical Pharmaceutical Fertilizer
Reciprocating- plate columns	High throughput Low HETS Great versatility and flexibility Simplicity in construction Handles liquids containing suspended solids Handles mixtures with emulsifying tendencies	Pharmaceutical Petrochemical Metallurgical Chemical
Centrifugal extractors	Short contacting time for unstable material Limited space required Handles easily emulsified material Handles systems with little liquid density difference	Pharmaceutical Nuclear Petrochemical

(Reprinted by permission from T. C. Lo, Recent Developments in Commercial Extractors, Engineering Foundation Conference on Mixing Research, Rindge, N.H., 1975).

Capacities of line mixers such as those of Figure 10.13 and of static mixers such as those of Figure 10.14 are stated in manufacturers catalogs. A procedure for estimating mixing efficiencies from basic correlations is illustrated by Laddha and Degaleesan (1978, p. 424).

Separation of the mixed phases is accomplished by gravity settling or less commonly by centrifugation. It can be enhanced by inducing coalescence with packing or electrically, or by shortening the distance of fall to a coalesced phase. Figures 14.11(d), 18.2, and 18.3 are some examples. Chapter 18 deals with some aspects of the separation of liquid phases.

A common basis for the design of settlers is an assumed droplet size of $150 \,\mu$ m, which is the basis of the standard API design method for oil-water separators. Stokes law is applied to find the settling time. In open vessels, residence times of 30-60min or superficial velocities of 0.5–1.5 ft/min commonly are provided. Longitudinal baffles can cut the residence time to 5–10 min. Coalescence with packing or wire mesh or electrically cut these

478 EXTRACTION AND LEACHING

		Extraction	Equ	ipment			
TABLE	14.3.	Comparisons	Of	Performance	and	Costs	Of

(a)	Some	Comparisons	and	Other	Performance	Data
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		Total Flow Capacity (Imp. gal/hrft²)
System	Equipment	Pilot Plant Plant
co-ni-d2ehpa H ₂ SO ₄	Mixco agitated Karr reciprocating	(4 in.) 300 (60 in.) 170 (3 in.) 900
Zr-Hf-TBP	Mixco agitated	(2 m.) 900 (30 in.) 184
HNO,	sieve plate pulse (steel)	(2 in.) 500
	sieve plate pulse (Teflon) RDC	(2 in.) 1345 (10 in.) 1345 (30 in.) 135
Hf-Zr-MIBK SCN	spray column	(4 in.) 2450
Rare earths– D2EHPA H₂SO₄	Podbielniak centrifuge	(4 feed dia) 30,000 gal/hr
U-amine-solvent- in-pulp H ₂ SO ₄	sieve plate pulse	(2 in.) 600 (10 in.) 900
Cu-Lix 64N H₂SO₄	mixer settlers	60-l 20
Cu-Ni-amine HCI	mixer settlers	60-120

(b) Cost Comparison, 1970 Prices, for Extraction of 150 gpm of Aqueous Feed Containing 5 g/L of **Cu** with 100 gpm Solvent, Recovering 99% of the Copper

	Equipment		Required			
Contactor	No.	Dia. Len No. (ft) (f		Equip. cost \$ × 1000	Total cost \$ × 1000	
Mixer settler	2	-	_	60	151.2	
Mixco	3	5	16	100	246.7	
Pulse	1	5	60	160	261.5	
Kenics	3	2	28	230	336.1	
Podbielniak		3-D36	_	300	378.0	
Graesser	1 5	5	3.0	88	308.0	

'Mixers have 150 gal capacity, settlers are 150 sqft by 4 ft deep with 9 in. solvent layer.

(G.M. Ritcey and A.W. Ashbrook, Solvent Extraction, 1979, Vol. II).

times substantially. A chart for determining separation of droplets of water with a plate pack of 3/4 in. spacing is reproduced by Hooper and Jacobs (in Schweitzer, 1979, 1.343-1.358). Numerical examples of settler design also are given in that work. For especially difficult separations or for space saving, centrifuges are applied. Liquid hydrocyclones individually have low efficiencies, but a number in series can attain 80–85% efficiency overall. Electrical coalescence is used commonly for separation of brine from crude oil; the subject is treated by Waterman (*Chem. Eng. Prog.* 61(10), 51 1965).

A control system for a mixer-settler is represented by Figure 3.19.

SPRAY TOWERS

These are empty vessels with provisions for introducing the liquids as dispersed or continuous phases and for removing them. Figure 14.12(a) shows both phases dispersed, which may be demanded when substantial changes in volumetric or physical properties result from solute transfer. Capacities of spray towers are high because of their openness, and they are not bothered by suspended solids. Backmixing is severe in towers of more than a few inches in diameter. Without operating experience to the contrary, even towers 20-40 ft high cannot be depended upon to function as more than single stages. The cross section is determined by the flooding velocity; that of the continuous phase is correlated by the equation

$$V_{CF} = \frac{4000\Delta\rho^{0.28}}{[0.483\mu_C^{0.075}\rho_C^{0.5} + d_P^{0.056}\rho_D^{0.5}(V_D/V_C)^{0.5}]^2} \quad \text{ft/hr}, \quad (14.33)$$

where a factor of 0.4 suggested by Treybal has been incorporated for safe design. The units are ft lb hr; the viscosity μ_C lb/ft hr = 2.42 cP. For large capacities, several parallel towers of at most 2 ft dia should be used. Commercially, spray towers are suitable for liquid-liquid processes in which rapid, irreversible chemical reactions occur, as in neutralization of waste acids. The substantial literature of flooding, holdup, mass transfer and axial mixing in small spray towers is reviewed by Laddha and Degaleesan (1978, pp. 221-255) and more briefly by Cavers (in Lo et al., 1983, pp. *320-328*).

PACKED TOWERS

Since mass transfer in packed or spray towers occurs differentially rather than stagewise, their performance should be expressed in terms of the number of transfer units (NTU) rather than the number of theoretical stages (NTS). For dilute systems, the number of transfer units is given in terms of the terminal concentrations and the equilibrium relation by

$$NTU = \int_{x_1}^{x_2} \frac{dx}{x - x_{equilib}}$$
(14.34)

In order to permit sizing a tower, data must be available of the height of a transfer unit (HTU). This term often is used interchangeably with the height equivalent to a theoretical stage (HETS), but strictly they are equal only for dilute solutions when the ratio of the extract and raffinate flow rates, E/R, equals the distribution coefficient, $K = x_E/x_R$ (Treybal, 1963, p. 350). Extractor performance also is expressible in terms of mass transfer coefficients, for instance, $K_E a$, which is related to the number and height of transfer units by

$$\frac{K_E a \Delta C}{E/S} = \frac{\text{NTU}}{Z} = \frac{1}{\text{HTU}},$$
(14.35)

where *E/S* is the extract flow rate per unit cross section and AC is mean concentration difference of the solute. Correlations of this quantity based on data from towers of 1-2 in. dia have been made, for example, by Laddha and Degaleesan (1978). They may be of qualitative value in predicting performance of commercial equipment when combined with some direct pilot plant information. In commercial size towers, HETS of 2-5 ft may be realized. Mass transfer drops off sharply with axial distance, so that the dispersed phase is redistributed every 5-7 ft. A sketch of a redistributor is in Figure 14.12(e). Extractors with three or more beds are not uncommon.

Packed towers may be employed when 5-10 stages suffice. They are not satisfactory at interfacial tensions above 10 dyn/cm. Even at this condition, sieve trays have greater efficiency, and at much higher interfacial tensions some form of agitated tower is required.

Metal and ceramic packings tend to remain wetted with the



Figure 14.11. Some types and arrangements of mixers and settlers. (a) Kemira mixer-settler (*Mattila*, Proc. Solvent Extraction Conference, *ZSEC* 74, *Inst. Chem. Eng.*, London, 1974); (b) Injection mixer and settler (*Ziolkowski*, 1961). (c) Gravity settler; "rag" is foreign material that collects at the interface. (d) Provisions for improving rate of settling: (top) with packing or wire mesh; (bottom) with a nest of plates. (e) Compact arrangement of pump mixers and settlers [*Coplan et al.*, Chem. Eng. Prog. 50, 403 (1954)]. (f) Vertical arrangement of a battery of settlers and external mixers (*Lurgi Gesellschaften*).



Figure 14.12. Tower extractors without agitation. (a) Spray tower with both phases dispersed. (b) Two-section packed tower with light phase dispersed. (c) Sieve tray tower with light phase dispersed. (d) Sieve tray construction for light phase dispersed (left) and heavy phase dispersed (right). (e) Redistributor for packed tower with light phase dispersed (*Treybal*, 1963).



Figure 14.13. Towers with reciprocating trays or with pulsing action. (a) Assembly of a 36 in. Karr reciprocating tray column (*Chem.* Pro. Co.). (b) Sieve trays used in reciprocating trays columns; (left) large opening trays for the Karr column; (middle) countermotion trays with cutouts; (right) countermotion trays with downpipes for heavy phase. (c) Rotary valve pulsator, consisting of a variable speed pump and a rotary valve that alternately links the column with pairs of suction and discharge vessels. (d) Sieve tray tower with a pneumatic pulser [Proc. Int. Solv. Extr. Conf. 2, 1571 (1974)]. (e) A pulser with a cam-operated bellows.

liquid that first wets them, so that the tower should be charged first with the continuous phase. Thermoplastics tend to be preferentially oil wetted, but they can be wetted by aqueous phase if immersed in it for several days.

Intalox saddles and pall rings of 1–1.5-in. size are the most commonly used packings. Smaller sizes tend to be less effective since their voids are of the same order of magnitude as drop

sizes. The flooding correlation of Figure 14.16 is recommended by Eckert (1984); a safe design is about 70% of the value obtained with this correlation. Dispersed phase loadings should not exceed 25 gal/(min)(sqft). Dispersion is best accomplished with perforated plates in which hole sizes are 3/16-1/4 in. Velocities through the holes should not exceed 0.8 ft/sec, but if short riser tubes are employed the velocities can be as high as 1.5 ft/sec.



Figure 14.14. Tower extractors with rotary agitators. (a) RDC (rotating disk contactor) extraction tower (Escher B.V., Holland). (b) Oldshue-Rushton extractor with turbine impellers and stator rings (*Mixing Equip.* Co.). (c) ARD (asymmetric rotating disk) extractor: (1) rotating disk rotor; (2) mixing zone; (3) settling zone (*Luwa A.G.*). (d) Kuhni extractor, employing turbine impellers and perforated partitions (*Kühni Ltd.*). (e) EC (enhanced coalescing) extractor [*Fischer et al.*, Chem. Ing. Tech., 228 (Mar. 1983)]. (f) Model of Scheibel extractor employing baffled mixing stages and wire mesh separating zones (E.G. Scheibel Inc.). (g) Model of Scheibel extractor employing shrouded turbine impellers and flat stators, suited for larger diameter columns (E.G. Scheibel Inc.).





SIEVE TRAY TOWERS

In sieve or perforated tray towers, the continuous phase runs across each tray and proceeds to the next one through a downcomer or riser. The dispersed phase is trapped as a coalesced layer at each tray and redispersed. The designs for light phase or heavy phase dispersion are shown in Figure 14.12(d). Either phase may be the dispersed one, but usually it is the raffinate. Both the reduced axial mixing because of the presence of the trays and the repeated dispersion tend to improve the efficiency over the other kinds of unagitated towers.

Hole diameters are much smaller than for vapor-liquid contacting, being 3-8 mm, usually on triangular spacing of 2-3 dia, and occupy from 15 to 25% of the available tray area. The area at a downcomer or riser is not perforated, nor is the area at the support ring which may be an inch or two wide. Velocities through the holes are kept below about 0.8 ft/sec to avoid formation of very small droplets. The head available for flow of the continuous pulse is the tray spacing. It is estimated as 4.5 velocity heads and thus is given by the equation

$$h = 4.5 V_d^2 \rho_C / 2g_c \tag{14.36}$$

where V_d is the linear velocity in the downcomer. It is usual to fill the downcomer with packing to coalesce entrainment; then the downcomer cross section must be made correspondingly larger.

Diameter of the Tower. The cross section of the tower must be made large enough to accommodate the downcomer and the perforated zone. Diameters of 12 ft or more are common.

Tray spacing is from 6-24 in., the larger dimension to facilitate servicing the trays in place when necessary. Both the downcomer cross section and the depth of coalesced layer are factors related to the spacing, and so is the efficiency. The depth of coalesced layer at each tray must be sufficient to force the liquid through the holes. In the range of 1 ft/sec through the holes, surface tension does not affect the flow significantly, so that the head-velocity relationship is the common one through orifices, namely,

$$V = 0.67\sqrt{2g_c}h\Delta\rho/\rho_D. \tag{14.37}$$

A correction also can be applied for the ratio of perforated and total tray areas. For the case of Example 14.10, the depth of coalesced layer is 1.6 in. according to this equation.

Tray Efficiency. A rough correlation for tray efficiency is due to Treybal (1963); as modified by Krishnamurty and Rao [*Ind. Eng. Chem. Process. Des. Dev.* 7, 166 (1968)] it has the form

$$E = (0.35Z_T^{0.5}/\sigma d_0^{0.35})(V_D/V_C)^{0.42}, \qquad (14.38)$$

where the interfacial tension σ is in dyn/cm and the tray spacing Z_T and hole diameter d_0 are in ft. Efficiencies and capacities of several kinds of extractors are summarized in Figure 14.17.

Application of the rules given here for sizing extraction towers without mechanical agitation is made in Example 14.10. The results probably are valid within only about 25%. The need for some pilot plant information of the particular system is essential.

PULSED PACKED AND SIEVE TRAY TOWERS

A rapid reciprocating motion imparted to the liquid in a tower results in improved mass transfer. This action can be accomplished without parts and bearings in contact with the process liquids and consequently has found favor for handling hazardous and corrosive liquids as in nuclear energy applications. Most of the applications still are in that industry, but several other installations are listed by Lo et al. (1983, pp. 345, 366). Packed columns up to 3mdia and 10 m high with throughputs in excess of 200 m^3/hr are in use.

Both packed and perforated plate towers are in use. The most



Figure 14.15. A horizontal rotating extractor and two kinds of centrifugal extractors. (a) The RTL (formerly Graesser raining bucket) horizontal rotating extractor; both phases are dispersed at some portion of the rotation (*RTL S. A., London*). (b) Operating principle of the Podbielniak centrifugal extractor; it is made up of several concentric perforated cylinders (Baker-Perkins Co.). (c) The Luwesta centrifugal extractor (schematic diagram) (*Luwa* Corp.).

commonly used packing is 1 in. Raschig rings. A "standard" geometry for the plates is 3mmdia holes on triangular spacing to give 23% open area, plate thickness of 2 mm, and plate spacing of 50mm. Reissinger and **Schröter** (1978) favor 2mm holes and **100 mm** plate spacing. The action of the plates is to disperse the heavy phase on the upstroke and the light phase on the down stroke.

Pulsing is uniform across the cross section, and accordingly the height needed to achieve a required extraction is substantially independent of the diameter as long as hydrodynamic similarity is preserved. Although correlations for flooding, holdup, and HTU are not well generalized, a major correlating factor is the product of frequency f and amplitude A_p ; in practical applications fA_p is in the range of 20-60 mm/sec.

One large user has standardized on a frequency of 90 cycles/min and amplitudes of vibration of 6-25 mm. Three kinds of pulsing modes are shown in Figures 14.13(c)-(e). The rotary valve pulsator consists of two reservoirs each on the suction and discharge of a variable speed **centifugal** pump and hooked to a rotating valve. Pneumatic and reciprocating pump **pulsers** also are popular.

Extraction efficiency can be preserved over a wide range of throughputs by adjusting the product fA_p . A comparison of several correlations of HTU made by Logsdail and Slater (in Lo et al., 1983, pp. 364) shows a four- to five-fold range, but a rough conservative rule can be deduced from these data, namely

$$HTU = 3.7/(fA_p)^{1/3}, \quad 20 \le fA_0 \le 60 \text{ mm/sec}, \quad (14.39)$$

which gives an HTU of 1 m at $fP_p = 50$ mm/sec. In small diameter extractors, data for HETS of 0.2–0.5 m or less have been found, as appear in Figure 14.17.

Flooding, holdup, and mass transfer rates are highly interdependent and are not simply related. Reissinger and Schröter (1978) state that tray towers in comparison with other types have good efficiencies at 60 m^3/m^2 hr at frequencies of 60–90/min and amplitudes of 10 mm. Packed towers have about 2/3 the capacities of tray towers. Also in comparison with unagitated towers, which are limited to interfacial tensions below 10 dyn/cm, pulsed towers are not limited by interfacial tension up to 30–40 dyn/cm. Some



Figure 14.16. Flooding velocities in liquid-liquid packed towers [J.S. Eckert, Encycl. Chem. Process. Des. 21, 149-165 (1984)]. V = ft/hr (superficial velocity); C = continuous phase; D = disperse phase; a = sqft area of packing/cuft; A = difference; $\varepsilon =$ void fraction in packing; $\mu =$ viscosity centipoise continuous phase; $\rho = \text{lb/cuft}; \sigma = (\text{dynes/cm})$ interfacial surface tension; F = packing factor.

further comparisons are made in Tables 14.3 and 14.4 and Figure 14.17.

RECIPROCATING TRAY TOWERS

Desirable motion can be imparted to the liquids by reciprocating motion of the plates rather than by pulsing the entire liquid mass. This mode employs much less power and provides equally good extraction efficiency. A 30in. dia tower 20 ft high is sufficiently agitated with a **1.5 HP** motor. Some arrangements of **such** extractors are shown in Figure 14.13.

The holes of reciprocating plates are much larger than those of pulsed ones. Typical specifications of such extractors are: Holes are 9/16 in. dia, open area is 50-60%, stroke length 0.5-1.0 in., 100-150 strokes/min at 0.75 in. stroke length, plate spacing normally 2 in. but may vary from 1-6 in. when the physical properties vary significantly in different parts of the tower. In towers about 30in. dia, HETS is 20-25 in. and throughputs are up to $40 \text{ m}^3/\text{m}^2$ hr (2000 gal/hr sqft). Scaleup formulas for HETS and reciprocating speed, fA_p , are stated by the manufacturer, Chem Pro Corp.:

$$(\text{HETS})_2/(\text{HETS})_1 = (D_2/D_1)^{0.36}, \qquad (14.40) (fA_p)_2/(fA_p)_1 = (D_1/D_2)^{0.14}. \qquad (14.41)$$

The performance of a reciprocating tower is compared with several other small extractors in Figure 14.17.

An extractor with countermotion of alternate plates is known as the VPE (vibrating plate extractor). Figure 14.13(b) shows the arrangement. This model also is constructed with segmented plates or with downcomers for passage of the continuous phase. At least during some portion of the cycle, the tight phase coalesces and is trapped below the tray, just as in static tray extractors. The capacity of these units is greater than of those with full trays and the efficiency remains high. Some data (Lo et al., 1984, p. 386) indicate that some commercial extractions are completed satisfactorily in towers 4-8 m high at rates of 35-100 m^3/m^2 hr.

ROTATING DISK CONTACTOR (RDC)

The concept of arranging a battery of mixer-settlers in a vertical line in a single shell has been implemented in a variety of ways. In the RDC (Rotary Disk Contactor) extractor, the impellers are flat disks, the mixing zones are separated by partial diametral baffles called stators, but distinct settling zones are not provided. Figure 14.14(a) is a sketch. Because of its geometrical simplicity and its effectiveness, the RDC is one of the most widely employed of agitated extractors. The situations in which it may not be suitable are when only a few stages are needed, in which case mixer-settlers will be satisfactory and cheaper; or when their large holdup and long residence times may be harmful to unstable substances; or for systems with low interfacial tensions and low density differences because then stable emulsions may be formed by the intense agitation.

According to the comparisons of small units in Figure 14.17, the RDC is intermediate in stage efficiency and throughput. The value of HETS = 0.3 m from this figure compares roughly with the HTU = 0.4 or 0.75, depending on which phase is dispersed, of the pilot plant data of Example 14.11.

The design procedure used by Kosters, of Shell Oil Co., who developed this equipment, requires pilot plant measurements on the particular system of HTU and slip velocity as functions of power input. The procedure for scaleup is summarized in Table 14.5, and results of a typical design worked out by Kosters (in Lo et al., 1983, pp. 391-405) are summarized in Example 14.11. Scaleup by this method is said to be reliable in going from 64mm dia to 4-4.5 m dia. The data of Figure 14.18 are used in this study.

OTHER ROTARY AGITATED TOWERS

One of the first agitated tower extractors was developed by Scheibel (*AIChE. J.* 44, 681, 1948). The original design, like Figure 14.14(f), employed settling zones packed with wire mesh, but these were found unnecessary in most cases and now flat partitions between mixing zones are used. The Mixco [Fig. 14.14(b)] and Scheibel-York [Fig. 14.14(g)] units differ primarily in the turbine impellers, the Mixco being open and the other shrouded. In spite of the similarity of their equipment, the manufacturers have possibly different ranges of experience. Since extractor selection is not on an entirely rational basis, a particular body of experience may be critical for fine tuning.

Enhanced coalescing between stages is provided in the designs of Figure 14.14(e). The **Kühni** extractor of Figure 14.14(d) employs shrouded turbine impellers and perforated plate partitions between compartments and extending over the entire cross section. The ARD (asymmetric rotating disk) extractor has lateral spaces for settling between agitation zones.

Some performance data are cited for the Kiihni by Ritcey and Ashbrook (1979, p. 102):

%	Free	Cross	Section	m ³ /m ² hr	HETS (m)
		10		10	0.08
		40		50	0. 20

Although not all equipment is compared, Figure 14.17 shows the Kiihni to have a high efficiency but somewhat lower capacity than the RDC and other units.

Most of these types of equipment have at least several hundred installations. The sizing of full scale equipment still requires pilot planting of particular systems. The **scaleup** procedures require geometrical and hydrodynamic similarities between the pilot and full scale plants. Hydrodynamic similarity implies equalities of
EXAMPLE 14.10

Sizing of Spray, Packed, or Sieve Tray Towers Five theoretical stages are needed for liquid-liquid extraction of a system with these properties:

 $\begin{aligned} Q_D &= 600 \text{ cuft/hr,} \\ Q_c &= 500 \text{ cuft/hr,} \\ \rho_D &= 50 \text{ lb/cuft,} \\ \rho_c &= 60 \text{ lb/cuft,} \\ \mu_D &= 0.5 \text{ cP,} \\ \mu_c &= 1.0 \text{ cP, } 2.42 \text{ lb/ft hr,} \\ \sigma &= 10 \text{ dyn/cm, interfacial tension,} \\ d_0 &= 0.0208 \text{ ft } (0.25''), \text{ hole size,} \\ d_p &= 0.02 \text{ ft (droplet diameter).} \end{aligned}$

Spray tower: The flooding velocity is found with Eq. (14.33):

$$V_D = \frac{4000(10)^{0.28}}{[0.483(2.42)^{0.075}(60)^{0.5} + (0.02)^{0.056}(50)^{0.5}(1.2)^{0.5}]^2}$$

= 13.0 ft/hr,
A, = 600173.0 = 8.22 sqft,
D = 3.24 ft.

To accommodate five stages, a total height of 100 ft or so would be needed. Two towers each 3.5 ft dia by 50 ft high would be suitable.

Packed tower: Flooding velocity is obtained with Figure 14.17. For 1 in. metal pall rings,

 $\begin{aligned} F\varepsilon^2 &= a_p/\varepsilon = 6310.94 = 67.02, \\ (\mu_c/\Delta p)(\sigma/\rho_c)^{0.2}(a_p/\varepsilon)^{1.5} = (1/10)(10/60)^{0.2}(67.02)^{1.5} = 38.34, \\ \therefore 200 &= V_c [1 + (V_D/V_c)^{0.5}]^2 \rho_c/a_p \mu_c \\ &= V_c [1 + 1.2^{0.5}]^2 60/63(1), \\ V_c &= 47.83 \text{ ft/hr, at the flooding point.} \end{aligned}$

Take 70% of flooding:

 $A_{r} = 500/0.7(47.83) = 14.93$ sqft D = 4.36 ft.

Take a conservative HETS = 5 ft. Then the tower will be 4.5 ft dia with 25 ft of packing and two redistributors, a total of about 35 ft.

Sieve tray tower: Take 1.5 ft tray spacing, 0.25 in. holes on 0.75 in. triangular spacing. The downcomer area is found with Eq. (14.36):

Ah = 1.5 =
$$\frac{4.5V_D^2\rho_c}{2g_c\Delta\rho} = \frac{4.5(60)}{2(418)(10^8)(10)}V_D^2$$

 $V_D = 6815$ ft/hr, A. = 600/6815 = 0.088 sqft, $D_d = 4.02$ in., downcomer diameter.

Take hole velocity = 0.8 ft/sec, 2880 ft/hr:

total hole area = $\frac{600}{2880}$ = 0.2083 sqft. $\frac{\text{tray area}}{\text{hole area}} = \frac{0.866d_s^2}{\frac{1}{2}(\pi/4)d_0^2} = 2.21(\frac{ds}{d})^2$ = 2.21(3)² = 19.89, tray area = 19.89(0.2083) = 4.14 sqft.

Add area of two 4in. pipes, 4.5 in. OD = 0.11 sqft.

area = 4.14 + 0.11 = 4.25 sqft, dia = 21.9 in.

Add 2 in. for support rings, making the diameter 30 in. Tray efficiency from Eq. (14.38):

$$E = \frac{0.35(Z_t)^{0.5}}{d_0^{0.35}} \left(\frac{V_D}{V_C}\right)^{0.42} = \frac{0.35(1.5)^{0.5}}{10.(0.0208)^{0.35}(1.2)^{0.42}} = 0.18$$

number of trays = 5/0.18 = 27.8,

tower height = 1.5(28) + 6 = 48 ft, including 3 ft at each end.



Summary:

	Height	Diamete
Spray	100	3.2
Packed	35	4.5
Sieve tray	48	2.5

droplet diameters, fractional holdups, and linear superficial velocities. Also preserved are the specific radial discharge rates, defined by Q/DH = (volumetric flow rate)/(vessel dia) (compartment height).

A detailed design of an ARD extractor based on pilot plant work is presented by Misek and Marek (in Lo et al., 1983, pp. 407-417). The design and operating parameters of the ARD extractor are related to the vessel diameter D (mm); thus: Free cross section = 25%. Disk diameter = 0.490. Chamber height = $1.3D^{0.67}$. Agitator rpm = $15,000/D^{0.78}$.

A manufacturer's bulletin on a 150 mm dia ARD extractor gives HETS = 0.4 m and capacity 15 m^3/m^2 hr.



Figure 14.17. Efficiency and capacity range of small diameter extractors, 50-150 mm dia. Acetone extracted from water with toluene as the disperse phase, $V_d/V_c = 1.5$. Code: AC = agitated cell; PPC= pulsed packed column; PST= pulsed sieve tray; RDC = rotating disk contactor; PC = packed column; MS = mixersettler; ST= sieve trav [*Stichlmair*. Chem. Ine. Tech. **52**(3). 253–255 (1980)].

Less specific information about the other kinds of extractors mentioned here is presented by Lo et al. (1983, pp. 419-448) but no integrated examples. The information perhaps could be run down in the abundant literature cited there, or best from the manufacturers.

OTHER KINDS OF EXTRACTORS

Some novel types and variations of basic types of extractors have been developed, most of which have not found wide acceptance, for instance pulsed rotary towers. The literature of a few of them is listed by Baird (in Lo et al., 1983, pp. 453-457). Here the extractors illustrated in Figure 14.15 will be described.

Graesser Raining Bucket Contactor. The Graesser "raining bucket" contactor consists of a horizontal rotating shell with a shaft that carries a number of diametral partitions extending to the wall. Between the partitions are buckets that carry the liquid and cascade it through each phase. No attempt is made to effect dispersion beyond simply emptying the buckets. The light and heavy phases are alternately both dispersed. They are introduced and withdrawn at opposite ends. The speed of rotation is 1-40 rpm, depending on the diameter, and is gauged to effect proper mass transfer and yet avoid emulsification. An approximately linear relation exists between mass transfer rate and diameter up to about 2m. Production contactors usually are designed to provide about 0.3 theoretical stages per compartment. Phase flow ratios of 6:1 are accommodated readily and ratios of up to 20:1 also have been designed for. Solids can be leached in this equipment; for instance, bitumen is dissolved away from tar sands with kerosene.

A commercial unit 5 ft dia by 18 ft long has 26×7 -in, wide compartments each with 16×8 -in. buckets and provides six theoretical stages. A unit 12 in. dia by 3 ft long has a capacity of 30 gal/hr at 8 rpm. A unit 6 ft dia has a capacity of 6000 gal/hr at 1.4 rpm.

Centrifugal Contactors. These devices have large capacities per unit, short residence times, and small holdup. They can handle systems that emulsify easily or have small density differences or large interfacial tensions or need large ratios of solvent to feed. Some types are employed as separators of mixtures made in other equipment, others as both mixers and settlers, and some as differential contactors.

The Podbielmak contactor is a differential type. It is constructed of several perforated concentric cylinders and is shown schematically in Figure 14.15(b). Input and removal of the phases at each section are accomplished through radial tubes. The flow is countercurrent with alternate mixing and separating occurring respectively at the perforations and between the bands. The position of the interface is controlled by the back pressure applied on the light phase outlet.

Residence time can be as short as 10 sec. One 750 gpm unit is said to have a total liquid holdup of 200gal. From 3-10 stages per unit have been reported, although Table 14.6 shows a range of 1.8-7.7. A 65 in. dia casing can accommodate throughputs up to 25,000 gal/hr. An economic comparison of a Podbielniak with other

TABLE 14.4. Maximum Loads and Diameters of Extractors

Column Type	Maximum Load (m³/(m²)(h))	Maximum Column Diameter (m)	Maximum Throughput (m³/h)
Graesser contactor	<10	1.0	380
Scheibel	<20	1.0	16
Asymmetric rotating-disk	≈25	3.2-5.0	200
Lurgi tower	≈30	8.0	1500
Pulsed packed	≈40	2.8	250
Rotating-disk contactor	a40	4.0	500
Kiihni	≈50	3.0	350
Pulsed sieve-tray extractor	≈60	3.0	420
Karr	80-100	1.0	<80

These data apply at a high interfacial tension (30-40 dyn/cm), a viscosity similar to water, an inlet ratio of the phases of 1:1 parts by volume. and a density difference of approximately 0.6 g/cm³.

(Reissinger and Schröter, 1978).

Example **14.11**

Design of a Rotating Diik **Contactor** A hydrocarbon mixture containing 10% aromatics and at the rate of 55.5 m^3/hr is to be treated with a solvent at the rate of 173.6 m^3/hr . Ten stages are needed for the extraction. Pilot plant data are available for the HTU and the slip velocity; they are shown on the graphs for solvent either continuous or dispersed. The procedure of Table 14.5 was applied by Kosters (in Lo et al., 1983, pp. 391-405) with the following results:

	Solvent Continuous	Solvent Dispersed
Vessel dia (m)	2.1	1.7
Stator dia (m)	1.47	1.19
Rotor dia (m)	1.26	1.02
нтบ (m)	0.41	0.75
(HTU) _{eff} (m)	0.663	1.107
Number of compartments	40	81
Compartment height(m)	0.20	0.17
Total height (m)	10.4	15.7
Rotor speed (rpm)	15-60	15-70
Power (theoretical kW)	4.6	2.0



extractors is made in Table 14.3(b). Although its basic cost is high, it requires few auxiliaries so that the overall cost of an extraction plant is not drastically out of line in every instance. Nevertheless, this equipment is used primarily when short residence time and other characteristic features are indispensable.

Other kinds of **centrifugals** also are used widely. Some are described by **Hafez** (in Lo et al., 1983, pp. 459-474) and performance data are presented in Table 14.6. Characteristics of **centrifugals** that are used primarily for removal of solids from slurries are summarized in Table 11.18.

LEACHING EQUIPMENT

In leaching processes, finely divided solids are contacted with solvents to remove soluble constituents. Usually some kind of multistage and countercurrent operation is desirable. The most bothersome aspect is handling of the wet solids.

In the leaching battery of Figure 14.19(a), the solids are transported between vessels with slurry pumps and are mixed in line

with countercurrent solution from the next stage. For the process to be effective, the solids must settle freely. The tanks have sloped bottoms and slowly moving rakes that scrape the solids towards the center discharge. Units employed for treating ores, for example, are very large, 100-200 ft dia. A few performance data of settlers are in Table 14.7.

Solids being extracted remain fixed in the cells of the battery of Figure 14.20(b). Fresh solvent is charged to the cell that is most nearly exhausted and next to be taken off stream, then solution proceeds through the other cells in series and leaves as finished extract from the cell that has been charged most recently. For sugar beet extraction a battery normally consists of 10-14 cells. Cells have volumes ranging from 4-12 m³ and height to diameter ratios as high as 1.5. Since leaching is faster at elevated temperatures, the solutions are heated between cells. Leaching time is 60-100 min. The amount of solution made is 110 kg/100 kg beets and contains 13-16% sugar. Various kinds of barks and seeds also are extracted in this kind of equipment. Further details of the equipment

- 1. Stator opening diameter, S = 0.7D, where *D* is vessel diameter 2. Rotor diameter, R = 0.6D
- 3. Height to diameter ratio of a compartment:

D (m)0.5-1.01.0-I.51.5-2.52.5H/D0.150.120.10.08-0.1

- 4. Power input, Figure 14.20(a)
- 5. Fractional holdup at flooding, $h_{\rm fr}$, from Figure 14.20(b)
- 6. Slip velocity V_s preferably is obtained experimentally, but is given approximately by Figure 14.20(c)
- 7. Superficial velocity of the continuous phase at flooding,

 $V_{Cf} = \frac{V_{s} \exp(-h_{f})}{V_{D}/V_{C}h_{f} + 1/(1 - h_{r})^{2}}$

where V_{C} and V_{D} are the superficial velocities of the continuous and dispersed phases

8. Holdup hat an operating velocity V_c , say 70–80% of flooding,

 $V_{C} = \frac{V_{s} \exp(-h)}{V_{D}/V_{C}h + 1/(1-h)^{T}}$

solve by trial for h when other quantities are specified

(W.C.G. Kosters, Shell Oil Co.).

9. Effective height of a transfer unit,

(HTU)_{eff =} (HTU)_{pilot plant} + HDU

in terms of a value obtained in a pilot plant and a calculated height of a diffusion unit (HDU)

10. Height of a diffusion unit, HDU = $H(1/\text{Pe}_{C} + 1/\text{Pe}_{D})$ 11. Factors E_{C} and E_{D} for evaluating the Peclet numbers,

$$E_{c} = 0.5V_{c}H + 0.012RNH(S/D)^{2}$$

$$E_{c} = E_{c}[4.2(10^{5})(V_{c}/h)^{3.3}/D^{2}];$$

when the correction in brackets is less than unity, make ${\it E}_{\it D_{\it E}}$ = ${\it E_{\it C}}$ 12. The Peclet numbers are

$$\frac{1}{Pe_{C}} = E_{C}(1 - h)/HV_{C}$$
$$\frac{1}{Pe_{D}} = E_{D}h/HV_{D}$$

13. Final expression for height of a diffusion unit is

HDU =
$$E_c(1-h)/V_c + E_dh/V_D$$



Figure 14.18. Holdup at flooding, power input, and slip velocity in an RDC (*Kosters, in Lo, Baird, and Hanson, 1983*). (a) Fractional holdup at flooding, h_{f} , as a function of flow ratio of the phases. (b) Power input to one rotor as a function of rotation speed N and radius *R. (c)* Slip velocity versus power input group for density difference of 0.15 g/mL, at the indicated surface tensions (dyn/cm).

TABLE	14.6.	Performanceof	Centri fugal	Extractors	

Extractor	Model	Volume, m³	Capacity, m³/hr	rpm	Motor Mounting	Motor Power, k W	/ Diameter, m
Podbielniak	E48	0.925	113.5	1,600	Side	24	1.2
Quadronic	Hiatchi 4848	0.9	72	1,500	Side	5 5	1.2
α -Laval	ABE 216	0.07	21	6, 000	Тор	30	
UPV			6	1,400	Bottom	14	
Luwesta	EG 10006		5	4, 500	Bottom		
Robatel SGN	LX6 70NL	0.072	3. 5	1,600	Top, side		1.3
Robatel BXP	BXP 800	0. 220	50	1,000	Тор	15	0.8
Westfalia	TA 15007	0. 028	30	3, 500	Тор	63	0. 7
SRL/ANL		0.003	0.05	3,500	Top		0.1
MEAB	SMCS-10	0.00012	0.3	22, 000	Bottom		

SPECIFICATIONS"

^{*a*}Operating pressures are in the range 300-1 750 kPa; operating temperatures cover a very wide range; operating flow ratios cover the range $\frac{10}{1}$ • $\frac{10}{10}$ easily.

PERFORMANCE

			Operati	S	Number of	
Extractor	System	rpm	$R = Q_h/Q_l$	Q_t , m ³ /hr	Flooding, %	Stages
Podbielniak						
B-10	Kerosene-NBA ^a -water	3000	0.5	5.1	73	6-6.5
D-18	Kerosene-NBA-water	2000	0.5	11.1	58	5-5.5
A-l	Oil-aromatics-phenol ^b	5000	3.5	0.01-0.02	33-66	5-7.7
9000	Broth-penicillin B -pentacetate	2900	4.4	7.5		1.8
		2900	3.4	1.5		2.04
		2900	2.4	7.5		2.21
9500	Some system	2900	3.5	7.5		2.04
		2700	3. 5	7.5		2.19
		2500	3.5	7.5		2.30
		2300	3.5	7.5		2.36
	C&aromatics-furfural	2000	4.0	12.0	90	3 - 6
A-1	IAA ^c -boric acid-water	5000	l-o.3	0.01-0.03	44-95	3. 5-7.7
		3000	1.0	0.01	44	2.3
		4075	1.0	0.01	44	2.8
		4600	1.0	0.01	44	2.96
UPV	Oil-aromatics-phenol ^b	1400	0.8-1.2	6	75	2-5.8
Robatel SGN						
LX-168N	Uranyl nitrate-30% TBP	1500	1 - 0.2	2.14.5		7
LX-324	Some system	3100	1.6	24-63		3. 4-3. 9
SRL single stage	Uranyl nitrate- Ultrasene	1790	0. 5- 1. 5	6. 4-12	33-96	0. 92-0. 99
ANL single stage	Uranyl nitrate- TBP/dodacane	3500	0.3-4	0.8-1.6	50	0. 97- l

"Normal butyl amine. ^bContaining 1.74% water. ^cIsoamyl alcohol. "Number of theoretical and actual stages.

(M. Hafez, in Lo et al., 1983, pp. 459-474).



(a)



Figure 14.19. Continuous leaching equipment. (a) A battery of thickeners of the type shown, for example, in Figure 13.9(a), used in countercurrent leaching. The slurry is pumped between stages counter to the liquid flow: (A) mixing line for slurry and solution; (B) scraper arms; (C) = slurry pumps. (b) A bucket elevator with perforated buckets used for continuous extraction, named the **Bollmann** or **Hansa-Muchle** system [Goss, J. Am. Oil Chem. Soc. 23, 348 (1946)]. (c) A countercurrent leaching system in which the solid transport is with screw conveyors; a similar system is named Hildebrandt. (d) The Bonotto multi-tray tower extractor. The trays rotate while the solid is scraped and discharged from tray to tray. The solid transport is similar to that of the rotary tray dryer of Figure 9.8(a) [Goss, J. Am. Oil Chem. Soc. 23, 348 (1946)]. (e) Rotocel extractor, which consists of about 18 wedge-shaped cells in a rotating shell. Fresh solvent is charged to the last cell and the drained solutions are pumped countercurrently to each cell in series (*Blaw-Knox* Co.).



Figure 14.1%(continued)

arrangement are given by Badger and McCabe (Elements of *Chemical Engineering*, McGraw-Hill, New York, 1936).

Continuous transport of the solids against the solution is employed in several kinds of equipment, including screw, perforated belt, and bucket conveyors. One operation carries a bed of seeds 3-4 ft thick on a perforated belt that moves only a few feet per minute. Fresh solvent is applied 1/5 to 1/3 of the distance from the discharge, percolates downward, is collected in pans, and is redistributed by pumps countercurrently to the travel of the material.

The vertical bucket elevator extractor of Figure 14.19(b) stands 40-60 ft high and can handle as much as 50 tons/hr with 1-2 HP.

The buckets have perforated bottoms. As they start to descend, they are filled with fresh flaked material and sprayed with dilute intermediate extract. The solution percolates downward from bucket to bucket. As the travel turns upward, the buckets are subjected to countercurrent extraction with solution from fresh solvent that is charged about 1/3 the distance from the top. There is sufficient travel time for drainage before discharge of the spent flakes.

Countercurrent action is obtained in the Bonotto extractor of Figure 14.19(d). It has a number of trays arranged in vertical line and provided with scrapers to discharge solids through staggered openings in the trays. The principle of this mode of solid transport

		Rate of Solids Solid; is Size Feed in Under						olid; in Under-	ı
N	o.	(ft)	Slurry	Mesh	(tons/day)	Feed	flow(%)	Remarks
4	6	Х	5	Paint pigment	300	39	5.7%	33	Solubles washed out
3	16	Х	8	Iron oxide	300	162	10	33	C.C.D. washing
1	20	Х	8	Zinc, copper, lead	99.5%	400	20	40	
				ore	-200				
2	25	Х	10	Calcium carbonate	200	450	10	38	Feed is 14" Bé caustic
						each			liquor
1	40	Х	10	Flotation tailings	65%	800	20	55	To recover the water
					-200				
1	40	Х	12	Flotation mill con-		1050	25	56	
				centrates					

TABLE 14.7. Performance of Settling Tanks

(Hardinge Co.).



Figure 14.20. Single tank and battery of tanks as equipment for batch leaching. (a) A single tank extractor of the type used for recovering the oil from seeds. (b) Principle of the leaching battery. Cells are charged with solid and solvent is pumped through heaters and cells in series. In the figure, cell 1 has been exhausted and is being taken off stream and cell 3 has just been charged. (Badger and McCabe, Elements of Chemical Engineering, McGraw-Hill, New York, 1936).

is similar to that of Figure 9.8(b). Solvent is charged at the bottom of the tower and leaves at the top, and the spent solid is removed with a screw conveyor.

Few performance data of leaching equipment have found their

REFERENCES

- P.J. Bailes, C. Hanson, M.A. Hughes, and M.W.T. Pratt, Extraction, liquid-liquid, Encycl. *Chem. Process. Des.* 21, 19-125 (1984).
- A.E. Dunstan et al., Eds., in *Science* of *Petroleum*, Solvent extraction methods of refining, 1817-1929, Oxford University Press, Oxford, 1938, Section 28.
- 3. J.S. Eckert, Extraction, liquid-liquid, packed tower design, Encycl. *Chem.* Process. Des. 21, 149-166 (1984).
- C. Hanson Ed., Recent Advances in Liquid-Liquid Extraction, Pergamon, New York, 1971.
- E.J. Henley and J.D. Seader, Equilibrium-Stage Separation Operations in Chemical Engineering, Wiley, New York, 1981.
- A.E. Karr, Design scale up and application of the reciprocating plate extraction column, *Sep. Sci. Technol.* 15, 877-905 (1980).
- G.S. Laddha and T.E. Degaleesan, Transport Phenomena in Liquid Extraction, Tata McGraw-Hill, New York, 1978.
- T.C. Lo, M.H.I. Baird, and C. Hanson, Eds., Handbook of Solvent Extraction, Wiley, New York, 1983.
- K.H. Reissinger and J. Schröter, Selection criteria for liquid-liquid extractors, *Chem. Eng.*, 109–118 (6 Nov. 1978); also Encycl. *Chem. Process. Des.* 21, 125-149 (1984).
- G.M. Ritcey and A.W. Ashbrook, Solvent Extraction with Applications to Process Metallurgy, Elsevier, New York, 1979, Parts I, II.

way into the open literature, but since these processes have long been exploited, a large body of information must be in the files of manufacturers and users of such equipment.

- L.A. Robbins, Liquid-liquid extraction, in Ref. 13, pp. 1.256-1.282; in *Chemical Engineers' Handbook*, McGraw-Hill, New York, 1984, pp. 15.1–15.20, 21.56-21.83.
- H. Sawistowski and W. Smith, Mass Transfer Process Calculations, Wiley, New York, 1963.
- P.A. Schweitzer, Ed., Handbook of Separation Techniques for Chemical Engineers, McGraw-Hill, New York, 1979.
- J.M. Sorensen and W. Arlt, Liquid-Liquid Equilibrium Data Collection, DECHEMA, Frankfurt/Main, Germany, 1979–1980.
- 15. R.E. Treybal, Liquid Extraction, McGraw-Hill, New York, 1951, 1963.
- 16. R.E. Treybal, Mass Transfer Operations, McGraw-Hill, New York, 1980.
- 17. T. Tsuboka and T. Katayama, Design algorithm for liquid-liquid separation processes, J. Chem. Eng. Jpn. 9, 40-45 (1976).
- S.M. Walas, *Phase Equilibria in Chemical Engineering*, Butterworths, Stoneham, Mass., 1985.
- J. Wisniak and A. Tamir, Liquid-Liquid Equilibrium and Extraction Bibliography, Elsevier, New York, 1980.
- J. Wisniak and A. Tamir, *Phase Diagrams: A Literature Source Book*, Elsevier, New York, 1981.
- Z. Ziolkowski, Liquid Extraction in the Chemical Industry (in Polish), PWT, Warsaw, 1961.

eparation of the components of a fluid can be effected by contacting them with a solid that has a preferential attraction for some of them. Such processes are quantitatively significant when the specific surfaces of the solids are measured in hundreds of m^2/q . Suitable materials are masses of numerous fine pores that were generated by expulsion of volatile substances. The most important adsorbents are activated carbon, prepared by partial volatilization or combustion of a carbonaceous body, and activated alumina, silica gel, and molecular sieves which are all formed by expulsion of water vapor from a solid. The starting material for silica gel is a coagulated silicic acid and that for molecular sieves is hydrated aluminum silicate crvstals that end up as porous crvstal structures. Porous g/asses made by leaching with alkai have some application in chromatography. Physical properties of common adsorbents are listed in Tab/es 75.7 and 15.2. Representative manufacturing processes are represented on Figure 15.7.

The amount of adsorption is limited by the available surface and pore volume, and depends a/so on the chemical natures of the fluid and solid. The rate of adsorption a/so depends on the amount of exposed surface but, in addition, on the rate of diffusion to the external surface and through the pores of the so/id for accessing the internal surface which comprises the bulk of the surface. Diffusion rates depend on temperature and differences in concentration or partial pressures. The smaller the particle size, the greater is the utilization of the internal surface, but a/so the greater the pressure drop for flow of bulk fluid through a mass of the particles.

15.1. ADSORPTION EQUILIBRIA

The amount of adsorbate that can be held depends on the concentration or partial pressure and temperature, on the chemical nature of the **fluid**, and on the nature, specific surface, method of preparation, and regeneration history of the solid. For single adsorbable components of gases, the relations between amount adsorbed and the partial pressure have been classified into the six types shown in Figure 15.2. Many common systems conform to Type I, for example, some of the curves of Figure 15.3. Adsorption data are not highly reproducible because small contents of impurities and the history of the adsorbent have strong influences on their behavior.

One of the simplest equations relating amount of adsorption and pressure with some range of applicability is that of Freundlich,

$$w = aP^n \tag{15.1}$$

and its generalization for the effect of temperature

 $\omega = aP^n \exp(-b/T). \tag{15.2}$

The exponent n usually is less than unity. Both gas and liquid adsorption data are fitted by the Freundlich isotherm. Many liquid data are fitted thus in a compilation of Landolt-Börnstein (II/3, Numerical Data and Functional Relationships in Science and Technology, Springer, New York, 19.56, pp. 525–528), but their gas

In ion exchange equipment, cations or anions from the fluid deposit in the so/id and displace equivalent amounts of other ions from the solid. Suitable so/ids are not necessarily porous; the ions are able to diffuse through the solid material. A typical exchange is that of H^+ or OH^- ions from the solid for some undesirable ions in the solution, such as Ca⁺⁺ or SO_4^{--} . Eventually all of the ions in the so/id are replaced, but the activity is restored by contacting the exhausted so/id with a high concentration of the desired ion, for example, a strong acid to rep/ace lost hydrogen ions.

For economic reasons, saturated adsorbents and exhausted ion exchangers must be regenerated. Most commonly, saturation and regeneration are performed alternately and intermittently, but equipment can be devised in which these processes are accomplished continuously by countercurrent movement of the solid and fluid streams. On/y a few such operations have proved economically feasible. The UOP and Toray processes for liquid adsorption are not true continuous processes but are effectively such.

Desorption is accomplished by elevating the temperature, or reducing the pressure, or by washing with a suitable reagent. The desorbed material may be recovered as valuable product in concentrated form or as a waste in easily disposable form. Adsorbent carbons used for water treating often must be regenerated by ignition in a furnace. Relatively small amounts of adsorbents that are difficult to regenerate are simply discarded.

data are presented in graphical form only (LB IV 4/b, 1972, pp. 121-187). The effect of temperature also is correlated by a theory of Polanyi, whereby all data of a particular system fall on the same curve; Figure 15.4 is an example. For isothermal data, a combination of the Freundlich and Langmuir equations was developed by Yon and Turnock (*Chem. Eng. Prog. Symposium Series* 117, 67, 1971):

$$w = kP^n/(1 + kP^n).$$
 (15.3)

Individuals of multicomponent mixtures compete for the limited space on the adsorbent. Equilibrium curves of binary mixtures, when plotted as x vs. y diagrams, resemble those of vapor-liquid mixtures, either for gases (Fig. 15.5) or liquids (Fig. 15.6). The shapes of adsorption **curves** of binary mixtures, Figure 15.7, are varied; the total adsorptions of the components of the pairs of Figure 15.7 would be more nearly constant over the whole range of compositions in terms of liquid volume fractions rather than the mol fractions shown.

Higher molecular weight members of homologous series adsorb preferentially on some adsorbents. The desorption data of Figure 15.8 attest to this, the hydrogen coming off first and the pentane last. In practical cases it is not always feasible to allow sufficient time for complete removal of heavy constituents so that the capacity of regenerated adsorbent becomes less than that of fresh, as Figure 15.9 indicates. Repeated regeneration causes gradual deterioration

TABLE 15.1. Physical Properties of Adsorbents

	Particle Form'	Mesh Size	Effective Diameter D _p , ft.	Bulk Density <i>p_b</i> , Lb/cu.ft.	External Void Fraction F_a	External Surface a _v , sq.ft .	Specific Heat C _x , Btu/lb °F	Reactivation Temperature °F	Examples
Activated Carbon	P P G	4x6 6×8 8x 10 4x 10 6×16	0.0128 0.0092 0.0064 0.0110 0.0062	30 30 30 30 30 30	0.34 0.34 0.34 0.40 0.40	310 446 645 460 720	0.25 " 0.25	200-1000 11 11 11 11 11 11	Columbia L '' '' '' '' Pittsburgh BPL ''
Silica Gel	G G G	4x 10 3x8 6×16 4×8	0.0105 0.0127 0.0062 0.0130	28 45 45	0.44 0.35 0.35 0.35	450 230 720 300	0.22	11 11 250-450 11 11 300-450	Witco 256 Davison 03
Activated Alumina.	3 6 6	4×8 8×14 14×28 (1/40)	0.0130 0.0058 0.0027	50 52 52 54	8:25 0:25 0.25	380 380 480 970	0.22	350-600 11 11 12 10 1000	Alcoa Type F
Molecular Sieves	S G P S	'(1/8") ¹ (1/16") (1/8") 4×8	0.0208 0.0104 0.0027 0.0060 0.0104 0.0109	34 30 45 45	0:30 0:25 0:34 0.34 0.37	400 970 650 400 347	0.22	300-600	Davison, Linda
	S	8x12	0.0067	45	0.37	565	11	11 EL 12 EL	**

• P = pellets; G = granules; S = spheroids

(Fair, 1969).

TABLE 15.2. Data of Molecular Sieves

(a) Structures and Applications

				Effective Channel	
	Cationic	Formula of Typical		Diameter	
Framework	Form	Unit Cell	Window	(Å)	Application
	Na	$Na_{12}[(AlO_2)_{12}(SiO_2)_{12}]$	8-ring	3.8	Desiccant. CO2 removal
			(obstructed)		from natural gas
	Ca	$Ca_5Na_2[(AlO_2)_{12}(SiO_2)_{12}]$	8-ring	4.4	Linear paraffin
Α			(free)		separation. Air separation
	К	$K_{12}[(AlO_2)_{12}(SiO_2)_{12}]$	8-ring (obstructed)	2.9	Drying of cracked gas containing C_2H_4 , etc.
	(Na	Na86[(AlO2)86(SiO2)106]	12-ring	8.4	Pressure swing H ₂ purification
Х	Ca	$Ca_{40}Na_6[(AlO_2)_{86}(SiO_2)_{106}]$	12-ring	8.0	Removal of mercaptans from
	Sr. Ba"	$Sr_{21}Ba_{22}[(AlO_2)_{86}(SiO_2)_{106}]$	12-ring	8.0	Xylene separation
V	∫ Na	Na56[(AlO2)56(SiO2)136]	12-ring	8.0	Xylene separation
Ť	lκ	K ₅₆ [(AlO ₂) ₅₆ (SiO ₂) ₁₃₆]	12-ring	8.0	Xylene separation
Mordenite	{ Ag н	$Ag_8[(AlO_2)_8(SiO_2)_{40}]$ $H_8[(AlO_2)_8(SiO_2)_{40}]$	12-ring	7.0	I and Kr removal from nuclear off-gases ⁽³²⁻³⁴⁾
Silicalite	_	(SiO ₂) ₉₆	IO-ring	6.0	Removal of organics from water
ZSM-5	Na	$Na_3[(AlO_2)_3(SiO_2)_{93}]$	IO-ring	6.0	Xylene separation ⁽³¹⁾

'Also K-BaX.

(Ruthven, 1984).

Basic type	Nominal pore diameter, angstroms	Available form	Bulk density, lb/ft³	Heat of adsorption (max), Btu/lb HzO	Equilibrium H₂O capacity, % wt	Molecules adsorbed	Molecules excluded	Applications
3A	3	Powder Ho-in pellets %-in pellets	30 44 44	1800	2 3 20 2 0	Molecules with an effective diameter < 3 Å, including H ₂ O and NH,	Molecules with an effective diameter > 3 Å, e.g., ethane	The preferred molecular sieve adsorbent for the commerical dehydration of unsaturated hydrocarbon streams such as cracked gas, propylene, butadiene, and acetylene. It is also used for drying polar liquids such as methanol and ethanol.
4 A	4	Powder Hein pellets U-in pellets 8 X 12 beads 4 × 8 beads 14 × 30 mesh 4	30 45 45 45 45 45 45 4	1800	28.5 22 22 22 22 22 22 22	Molecules with an effective diameter < 4 A, including ethanol, H ₉ S, CO ₂ , SO ₂ , C ₂ H ₄ , C ₂ H ₄ , and C ₉ H ₆	Molecules with an effective diameter > 4 A, e.g., propane	The preferred molecular sieve adsorbent for static dehydration in a closed gas or liquid system. It is used as a static desiccant in household refrigeration systems; in packaging of drugs, electronic components and perishable chemicals; and as a water scavenger in paint and plastic systems. Also used commercially in drying saturated hydrocarbon streams.
5A	5	Powder ¼e-in pellets ¼-in pellets	30 43 43	1800	28 21.5 21.5	$\begin{array}{llllllllllllllllllllllllllllllllllll$	Molecules with an effective diameter > 5 A, e.g., iso compounds and all 4-carbon rings	Separates normal paraffins from branched-chain and cyclic hydrocarbons through a selective adsorption process.
10x	8	Powder ¼e-in pellets ¼e-in pellets	30 36 36	1800	36 28 2.8	Iso paraffins and olefins, C ₄ H ₆ , molecules with an effective diameter < 8 Å	Di-n-b&amine and larger	Aromatic hydrocarbon separation .
13x	10	Powder he-in pellets h-in pellets 8 × 12 beads 4 × 8 beads 14 × 30 mesh	3 0 3 8 3 8 4 2 4 2 3 8	1800	36 28.5 28.5 28.5 28.5 28.5 28.5	Molecules with an effective diameter < 10 Å	Molecules with an effective diameter > 10 A, e.g., (C4F9)sN	Used commercially for general gas drying, air plant feed purification (simultaneous removal of H_2O and CO_2), and liquid hydrocarbon and natural gas sweetening (H_2S and mercaptan removal).

or

TABLE **15.2**—(*continued*) (b) Typical Properties of Union Carbide Type X Molecular Sieves

(Kovach, 1978).

of adsorbent; Figure 15.10 reports this for a molecular sieve operation.

Representation and generalization of adsorption equilibria of binary and higher mixtures by equation is desirable, but less progress has been made for such systems than for vapor-liquid or liquid-liquid equilibria. The Yon and Turnock equations (1971) applied to components 1 and 2 of binary mixtures are

$$w_1/w_{1,sat} = k_1 P_1^{n_1} \theta,$$
 (15.4)

$$w_2/w_{2,sat} = k_2 P_2^{n_2} \theta,$$
 (15.5)

$$\theta = 1/(1 + k_1 P_1^{n_1} + k_2 P_2^{n_2}). \tag{15.6}$$

They have been found useful as an empirical correlation method for adsorption on molecular sieves [Maurer, *Am. Chem.* Soc. Symp. Ser. 135, 73 (1980)]. Other attempts at prediction or correlation of multicomponent adsorption data are reviewed by Ruthven (1984). In general, however, multicomponent equilibria are not well correlatable in general form so that design of equipment is best based on direct laboratory data with the exact mixture and the exact adsorbent at anticipated pressure and temperature.

Adsorption processes are sensitive to temperature, as the data of Figures 15.3, 15.5, and 15.11 show. Thus practical adsorption processes are complicated by the substantial heats of adsorption

that necessarily develop. These are of the same order of magnitude as heats of condensation. Some data are in Figure 15.4

15.2. ION EXCHANGE EQUILIBRIA

Ion exchange is a chemical process that can be represented by a stoichiometric equation, for example, when ion A in solution replaces ion B in the solid phase,

A (solution) + B (solid)
$$\rightleftharpoons \mathbf{A}$$
 (solid) + B (solution) (15.7)

$$\mathbf{A} + \mathbf{\bar{B}} / \mathbf{\bar{A}} + \mathbf{B}, \tag{15.8}$$

where the overstrike designates a component in the solid phase. The equilibrium constant is called the selectivity, designated by K_{AB} ,

$$K_{\mathbf{AB}} = C_{\bar{\mathbf{A}}} C_{\mathbf{B}} / C_{\mathbf{A}} C_{\bar{\mathbf{B}}}$$
(15.9)

$$= \mathbf{x}_{\bar{\mathbf{A}}} \mathbf{x}_{\mathbf{B}} / \mathbf{x}_{\mathbf{A}} \mathbf{x}_{\bar{\mathbf{B}}}$$
(15.10)

$$= \left| \frac{x_{\bar{\mathbf{A}}}}{1 - x_{\bar{\mathbf{A}}}} \right| / \left| \frac{x_{\mathbf{A}}}{1 - x_{\mathbf{A}}} \right|. \tag{15.11}$$

The last equation relates the mol fractions of the ion originally in the solution at equilibrium in the liquid (x_A) and solid $(x_{\bar{A}})$ phases.



Figure 15.1. Processes for making adsorbents. (a) Flowsketch of a process for making molecular sieve adsorbents. (b) Process for reactivation of bone char. (c) Silica gel by the BASF process. The gel is formed and solidifies in air from sodium silicate and sulfuric acid, then is washed free of sodium sulfate with water (*Ullmann*, Encyclopedia of Chemical Technology, *Verlag Chemie, Weinheim, Germany*).



Figure 15.2. Types of adsorption isotherms: (I) monomolecular layer; (II and III) multimolecular layers; (IV and V) multimolecular layers and condensation in pores; (VI) phase transition of a monomolecular layer on the surface (after *Brunauer*, Physical Adsorption, *Princeton Univ. Press*, 1945).

The residual mol fraction in the liquid phase corresponding to a given mol fraction or degree of saturation in the solid phase is

$$\boldsymbol{x}_{\mathbf{A}} = \frac{1}{1 + K_{\mathbf{A}\mathbf{B}}(1 - \boldsymbol{x}_{\bar{\mathbf{A}}})/\boldsymbol{x}_{\bar{\mathbf{A}}}}.$$
 (15.12)

Approximate values of the selectivity of various ions are shown in Table 15.3; for a particular pair, K_{AB} is the ratio of tabulated values for each.

When the exchanged ion D is divalent, the reaction is

$$\mathbf{D} + 2\mathbf{B} \rightleftharpoons \mathbf{D} + 2\mathbf{B},\tag{15.13}$$



Figure 15.3. Effects of temperature, pressure, and kind of adsorbent on the amount of ethane adsorbed: (1) activated carbon at 25° C; (2) type 4A molecular sieve (MS) at 0° C; (3) type 5A MS at 25° C; (4) type 4A MS at 25° C; (5) type 4A MS at 75° C; (6) silica gel at 25° C; (7) type 4A MS at 150° C. (*Data from Union Carbide Corp.*)



Figure 15.4. Polanyi characteristic curve for effect of temperature on adsorption of n-butane on silica gel [*Al-Sahhat et al.*, Ind. Eng. Chem. Process. Des. Dev. 20, 658 (1981)].

and the equilibrium constant or selectivity is given by

$$(C/C)K_{DB} = x_{B}^{2}x_{\bar{D}}/x_{\bar{B}}^{2}x_{D}$$
(15.14)
= $(x_{\bar{D}}/x_{D})[(1 - x_{D})/(1 - x_{\bar{D}})]^{2}$, (15.15)

where C and \bar{C} are the total concentrations of the two kinds of ions in the solution and in the solid, respectively.



Figure 15.5. Adsorption of binary mixtures: (1) ethane + ethylene. Type 4A MS 25°C, 250 Torr; (2) erhane + ethylene. Type 4A MS, 25°C, 730Torr; (3) ethane + ethylene. Type 4A MS, 75°C, 730Torr; (4) carbon dioxide + hydrogen sulfide. Type 5A MS, 27°C, 760Torr; (5) n-pentane + n-hexane, type 5A MS, 100°C, 760Torr; (6) ethane + ethylene, silica gel, 25°C, 760Torr; (7) ethane + ethylene, Columbia G carbon, 25°C, 760Torr; (8) acetylene + ethylene. Type 4A MS, 31°C, 740 Torr. (*Data from Union Carbide Corp.*)



Figure 15.6. Binary liquid adsorption equilibria on X-Y diagrams: (1) toluene + iso-octane on silica gel (Eagle and Scott, 1950); (2) toluene+ iso-octane on charcoal (Eagle and Scott, 1950); (3) ethylene dichloride + benzene on boehmite (Kipling); (4) ethylene dichloride + benzene on charcoal (Kipling). (*Kipling in* Proceedings of the Second International Congress of Surface Activity. (1957), Vol. III, p. 462.)



Figure 15.7. Adsorption of liquid mixtures on charcoal. Chloroform + acetone and benzene + ethanol. The ordinate gives the amount of each individual substance that is adsorbed, the abscissa the mol fraction of chloroform (mixed with acetone) or the mol fraction of benzene (mixed with ethanol). (Data *gathered by Kipling.* Adsorption from Solutions of Non-Electrolytes, 1965).



Figure 15.8. Variation of isosteric heat of adsorption with coverage showing the difference in trends between polar and nonpolar sorbates. $nC_4H_{10} - 5A$ (data of Schirmer et al.); CF_4 -NaX, SF,-NaX (data of Barrer and Reucroft); CO_2 -NaX, (data of Huang and Zwiebel), NH_3 -5A (data of Schirmer et al.); H_2O -LiX, NaX, and CsX, (data of Avgul et al.). [*Ruthven*, Sep Purification Methods 5(2), 189 (1976)].

Example 15.1 is concerned with such an exchange and regeneration process.

15.3. ADSORPTION BEHAVIOR IN PACKED BEDS

Adsorption is performed most commonly in fixed vertical beds of porous granular adsorbents. Flow of adsorbing fluid usually is down through the bed, that of regenerant usually is upward. Moving and fluidized beds have only a limited application in the field.

If the time is sufficient, the adsorbent nearer the inlet of the fluid becomes saturated at the prevailing inlet fluid concentration but a concentration gradient develops beyond the saturation zone. Figure 5.12 depicts this behavior. The region of falling concentration is called the mass transfer zone (MTZ). The gradient is called the adsorption wave front and is usually S-shaped. When its leading edge reaches the exit, breakthrough is said to have been attained. Practically, the breakthrough is not regarded as necessarily at zero concentration but at some low value such as 1% or 5% of the inlet that is acceptable in the effluent. A hypothetical position, to the left of which in Figure 15.12(b) the average adsorbate content equals the saturation value, is called the stoichiometric front. The distance between this position and the exit of the bed is called the length of unused bed (LUB). The exhaustion time is attained when the effluent concentration becomes the same as that of the inlet, or some practical high percentage of it, such as 95 or 99%.

The shape of the adsorption front, the width of the MTZ, and the profile of the effluent concentration depend on the nature of the adsorption isotherm and the rate of mass transfer. Practical bed depths may be expressed as multiples of MTZ, values of 5-10 multiples being economically feasible. Systems that have linear adsorption isotherms develop constant **MTZs** whereas **MTZs** of convex ones (such as Type I of Figure 15.1) become narrower, and those of concave systems become wider as they progress through



Figure 15.9. Concentrations in adsorption beds as a function of position and of effluent as a function of time. (a) Progress of a stable mass transfer front through an adsorption bed and of the effluent concentration (*Lukchis*, 1973). (b) The mass transfer zone (MTZ), the length of unused bed (LUB), stoichiometric front, and profile of effluent concentration after breakthrough.

the bed. The last types are called unfavorable isotherms; separations in such cases usually are accomplished more economically by some other kind of process. The narrower the MTZ, the greater the degree of utilization of the bed.

The rate of mass transfer from fluid to solid in a bed of porous granular adsorbent is made up of several factors in series:

- 1. Diffusion to the external surface.
- 2. Deposition on the surface.
- 3. Diffusion in the pores.
- 4. Diffusion along the surface.

Various combination of shapes of isotherms and mass transfer factors have been taken into account by solutions of the problem in the literature. One of the simpler cases was adopted by Hougen and Marshall (1947, see Figure 15.13), who took a linear isotherm and diffusion to the external surface as controlling the rate. They developed the solution in analytical form, of which several approximations that are easier to use are mentioned for instance by Vermeulen et al. (1984, p. 16.28). A graphical form of the solution appears in Figure 15.13. This shows the effluent concentration ratio,

 C/C_0 , in terms of a time parameter τ , at a number of values of a parameter Z', which involves the bed length Z. In Example 15.2, this chart is used to find the concentration profile of the effluent, the break and exhaustion times, and the % utilization of the adsorbent bed. In this case, the model affords a fair comparison with experimental data.

Many investigations have been conducted of the mass transfer coefficient at the external surfaces of particles and of other diffusional mechanisms. Some of the correlations are discussed in Chapters 13 and 17. A model developed by Rosen [*Ind. Eng. Chem.* 46, 1590 (1954)] takes into account both external film and pore diffusional resistance to mass transfer together with a linear isotherm. A numerical example is worked out by Hines and Maddox (1985, p. 485).

In the model developed by Thomas [J. Am. Chem. Soc. 66, 1664 (1944)], the controlling mechanism is the surface kinetics represented by the Langmuir isotherm. Extensions of this work by Vermeulen et al. (1984) incorporate external surface and pore diffusional resistances.

No comprehensive comparisons of the several models with each other and with experimental data appear to have been published.



Figure 15.10. Incomplete regeneration of adsorbent bed by a thermal-swing cycle.



Figure 15.11. Effect of temperature on molecular sieve type 5A, silica gel and activated alumina at water vapor pressure, 13.3 kPa (100 mm Hg). A, molecular sieve type 5A; B, silica-type adsorbent; C, alumina-type adsorbent. (*Chi and Cummings, 1978*).

TABLE 15.3. Gas Phase Adsorption Cycles, Steam Requirements, and Operating Parameters

(a) Typical Cycle Times in Hours for Adsorber Operation

	High Pre Gas Dr	essure 'yer	Organic S Recover	Solvent y Unit
	Α	В	Α	В
Dnstream	. 24	24	2.00	1.00
Depressure/purge	. 2	1		
Hot gas	. 10	13		• • • • •
Steam			0.75	0.67
Hot gas			0.33	
Cold gas	. 5	8	0.42	0.33
Pressure/standby	. 7	2	0.50	
			-	-
	24	24	2.00	1.00

(Fair, 1969).

(b) Steam/Solvent Ratios and Amount of Adsorbate for a Coconut-Shell Carbon 6-12 Mesh, 1200 m^2/g



(c) Typical Operating Parameters for Gas Phase Adsorption

	Range	Design
Superficial gas velocity	20 to 50 cm/s (40 to 100 ft/min)	40 cm/s (80 ft/min)
Adsorbent bed depth	3 to 10 MTZ	5 MTZ
Adsorption time	0.5 to 8 h	4 h
Temperature	-200 to 50°C	
Inlet concentration		
Adsorption base	100 to 5000 vppm	
LEL base	40%	
Adsorbent particle size	0.5 to 10 mm	4 to 8 mm
Working charge	5 to 20% wt	10%
Steam solvent ratio	2:1 to 8:1	4:I
Adsorbent void volume	38 to 50%	45%
Steam regeneration temperature	105 to 1 10°C	
Inert gas regenerant termperature	100 to 300°C	
Regeneration time	¹ / ₂ adsorption	time
Number of adsorbers	1 to 6	2 to 3

(Kovach, 1979).

EXAMPLE 15.1

Application of Ion Exchange Selectivity Data The $SO_4^=$ ion of an aqueous solution containing C = 0.018 eq/L is to be replaced with Cl- ion from a resin with $\ddot{C} = 1.2$ eq/L. The reaction is

 $SO_4^{\neq}(solution) + 2Cl^{-}(resin) \rightleftharpoons SO_4^{\neq}(resin) + 2Cl^{-}(solution),$ $D + 2\bar{B} \rightleftharpoons \bar{D} + 2B$.

From Table 15.3, the selectivity ratio $K_{DB} = 0.15/1.0 = 0.15$, and

 $K_{\rm DB}\bar{C}/C = 0.15(1.2)/0.018 = 10.$

Then Eq. (15.15) becomes

$$x_{\bar{\mathbf{D}}}/(1-x_{\bar{\mathbf{D}}})^2 = 10x_{\mathbf{D}}/(1-x_D)^2$$
.

For several values of mol fraction $x_{\rm D}$ of SO_4^- in solution, the corresponding mol fractions $x_{\tilde{D}}$ in the resin are calculated and tabulated:

	X _{SOT}	
In	Solution	In Resin
	1	1
	0.1	0.418
	0.05	0.284
	0.01	0.0853

For regeneration of the resin, a 12% solution of NaCl will be used;

its ion concentration is 2.23 eq/L. Other values for the system remain at $\tilde{C} = 1.2$ eq/L and $K_{DB} = 0.15$. Accordingly,

$$K_{\rm DB}\bar{C}/C = 0.15(1.2)/2.23 = 0.0807$$

and Eq. (15.15) becomes

$$x_{\tilde{D}}/(1-x_{\tilde{D}})^2 = 0.0807x_{D}/(1-x_{D})^2$$
.

The values of $x_{SO_{\overline{A}}}$ in the liquid phase will be calculated for several values in the resin. Those results will be used to find the minimum amount of regenerant solution needed for each degree of regeneration

SOT	i regenerant/
In Solution	L resin
0.455	1.06
0.319	1.60
0.102	5.22
	4 so ₹ In Solution 0.455 0.319 0.102

Sample calculation for the last entry of the table: The equivalents of SO_4^- transferred from the resin to the solution are

$$0.99(1.2) = 1.188 \text{ eq/L}.$$

The minimum amount of solution needed for this regeneration is

$$\frac{1.188}{0.102(2.23)}$$
 = 5.22 L solution/Liter.



Figure 15.12. Multicomponent mixtures, adsorption, and desorption. (a) Concentrations of the components of a ternary mixture in continuous adsorption, as in a moving bed unit (Kovach, 1979). (b) Composition of a desorbed stream consisting of several components as a function of time.



Figure 15.13. Dependence of the concentration ratio, C/C_0 of the effluent from an adsorber on parameters of bed length and time; for the case of a linear isotherm, zero initial adsorbate content and constant inlet composition C_0

 $Z' = (K_f a / \epsilon u_i) Z$, bed length parameter, $\tau = (K_f a / K_D \rho_b)(t - Z/u_i)$, time parameter, $K_D = q/C$, coefficient of linear adsorption isotherm, u_i = interstitial velocity in the bed, ϵ = voidage of the bed, Z = length of the bed, $K_f a$ = mass transfer coefficient, $(L^3 \text{ fluid})/(L^3 \text{ bed})$ (time).

(Hougen and Watson, Chemical Process Principles, Wiley, New York, 1947, p. 1086; Hougen and Marshall, Chem. Eng. Prog. 43, 197 (1947); Vermeulen et al., Chemical Engineers' Handbook, McGraw-Hill, New York, 1984, p. 16.29.)

Moreover, they are all based on isothermal behavior and approximations of adsorption isotherms and have not been applied to multicomponent mixtures. The greatest value of these calculation methods may lie in the prediction of effects of changes in basic data such as flow rates and slopes of adsorption isotherms after experimental data have been measured of breakthroughs and effluent concentration profiles. In a multicomponent system, each substance has a different breakthrough which is affected by the presence of the other substances. Experimental curves such as those of Figure 15.14 must be the basis for sizing an adsorber.

Since taking samples of adsorbent from various positions in the bed for analysis is difficult, it is usual to deduce the shape of the adsorption front and the width of the MTZ from the effluent concentration profile which may be monitored with a continuous analyzer-recorder or by sampling. The overall width of the MTZ, for instance, is given in terms of the exhaustion and breakthrough times and the superficial velocity as

width = $u_s(t_e - t_h)/\varepsilon$.

REGENERATION

Adsorbents are restored to essentially their original condition for reuse by desorption. Many hundreds of cycles usually are feasible, but eventually some degradation occurs, as in Figure 15.15 for instance, and the adsorbent must be discarded.

The most common method of regeneration is by purging the bed with a hot gas. Operating temperatures are characteristic of the adsorbent; suitable values at atmospheric pressure are shown in Table 15.1. The exit temperature of the gas usually is about 50°F higher than that of the end of the bed. Typical cycle times for adsorption and regeneration and steam/adsorbent ratios are given in Table 15.4. Effluent composition traces of a multicomponent system are in Figure 15.9. Complete removal of adsorbate is not always economically feasible, as suggested by Table 15.4(b). The effect of incomplete removal on capacity is shown schematically by Figure 15.10. Sufficient heat must be supplied to warm up the adsorbent and the vessel, to provide heat of desorption and enthalpy absorption of the adsorbate, and to provide for heat losses to the surroundings. Table 15.4(c) suggests that regeneration times be about one-half the adsorption times. For large vessels, it may be worthwhile to make the unsteady heating calculation by the general methods applicable to regenerators, as presented, for instance, by Hausen (Heat Transfer in Counterflow, Parallel Flow and Crossflow, McGraw-Hill, New York, 1983).

Purging of the adsorbate with an inert gas at much reduced pressure is feasible in high pressure adsorption plants. The adsorption of Example 15.2, for instance, is conducted at 55 atm, so that regeneration could be accomplished at a pressure of only a few atmospheres without heating. If the adsorbate is valuable, some provision must be made for recovering it from the desorbing gas.

Ignition of adsorbents in external furnaces is practiced to remove some high molecular weight materials that are difficult to volatilize, This is done, for example, for reactivation of carbon from water treating for trace removal of impurities such as phenol. Caustic solution can convert the phenol into soluble sodium phenate in readily disposable concentrated form as an alternate process for regeneration.

Displacement of the adsorbate with another substance that is in turn displaced in process is practiced, for instance, in liquid phase recovery of paraxylene from other C_8 aromatics. In the Sorbex process, suitable desorbents are toluene and paradiethylbenzene. This process is described later.

15.4. ADSORPTION DESIGN AND OPERATING PRACTICES

When continuous operation is necessary, at least two adsorbers are employed, one on adsorption and the other alternately on regeneration and cooling. In cases where breakthrough is especially harmful, three vessels are used, one being regenerated, the other two **onstream** with the more recently regenerated vessel downstream, as in Figure 15.16.

Beds usually are vertical; adsorbers 45 ft high and 8-10 ft dia are in use. When pressure drop must be minimized, as in the recovery of solvents from atmospheric air, horizontal vessels with shallow beds are in common use. Process gas flow most often is downward and regenerant gas flow is upward to take advantage of **counterflow** effects. **Upflow** rates are at most about one-half the fluidizing velocity of the particles. Vertical and horizontal types are represented on Figure 15.17.

A major feature of adsorber design is the support for the granular adsorbent, preferably one with a low pressure drop. The combination of Figure 15.18(a) of grid, screens, and support beams is inexpensive to fabricate and maintain, has a low heat capacity and a low pressure drop. The construction of Figure 15.18(b) is suited to adsorbers that must be dumped frequently. Supports of layers of ceramic balls or gravel or anthracite, resting on the bottom of the vessel, are suited to large vessels and when corrosion-resistant construction is required. Typical arrangements are shown in Figures 17.26 and 17.27. The successive layers increase in diameter by factors of 2-4 up to 1 in. or so. **Holddown** balls also may be

EXAMPLE 15.2

Adsorption of *n*-Hexane from a Natural Gas with Silica Gel Hexane is to be recovered from a natural gas with silica gel. Molecular weight of the gas is 17.85, the pressure is 55.4 atm, temperature is 94°F, and the content of n-hexane is 0.853 mol % or 0.0982 lb/cuft. The bed is 43 in. deep and the superficial velocity is 11.4 ft/min. Other data are shown with the sketch:



Z = 3.58 ft, bed depth, $u_s = 11.4$ ft/min, superficial velocity, $D_p = 0.01$ ft, particle diameter, a = 284 sqft/cuft, packing external surface, $\rho_b = 52$ lb/cuft, bed density, $\varepsilon = 0.35$ bed voidage.

From these and physical property data, the Schmidt and Reynolds numbers are calculated as

$$Sc = 1.87$$
, $Re = 644$.

The equation of Dwivedi and Upadhyay, Eq. (13.148), is applicable:

$$J_d = \frac{k_g}{u_s} \operatorname{Sc}^{2/3} = \frac{1}{\varepsilon} \left(\frac{0.765}{\operatorname{Re}^{0.82}} - \frac{0.365}{\operatorname{Re}^{0.386}} \right),$$

$$\therefore k_g = \frac{11.4}{0.35(1.87)^{2/3}} (0.0038 + 0.0301) = 0.7268 \text{ ft/min},$$

$$k_g a = 0.7268(284) = 206.4 \text{ cuft gas/(cuft solid)(min)}.$$

Saturation content of adsorbate is 0.17 lb/lb solid. Accordingly, the coefficient of the linear adsorption isotherm is

$$k_d = \frac{0.17}{0.0982} = 1.731 \frac{\text{lb hexane/lb solid}}{\text{lb hexane/cuft gas}}$$

Use the Hougen-Marshall chart (Fig. 15.13):

$$Z' = \frac{k_g a Z}{u_s} - \frac{206.4(3.58)}{11.4} = 64.82$$

$$t \frac{\underline{z} - k_d \rho_b}{k_g a} = \frac{1.731(52)}{u_s / \varepsilon} \frac{3.58}{11.4/0.35}$$

= 0.436\tau + 0.11 min.

Values of *t* are read off Figure 15.13 and converted into values oft:

ር/ቤ	r	t (min)
0.01	40	17.56
0.05	45	19.74
0.1	50	21.92
0.2	63	23.23
0.4	60	26.28
0.6	65	28.46
0.8	73	31.95
0.9	79	34.57
0.95	82	35.87
0.99	92	40.24

The total amount adsorbed to the breakpoint, at $C/C_0 = 0.01$, per sqft of bed cross section is

0.0982(11.4)(17.56) = 19.66 lb/sqft cross section.

The saturation amount for the whole bed is

3.58(0.17)(52) = 31.65 lb/sqft cross section.

Accordingly,

utilization of bed = (19.66/31.65)(100%) = 62.1%.

The calculated concentration profile is compared in the figure with experimental data, Run 117, of McLeod and Campbell, *Soc. Pet. Eng. J.*, 166 (June 1966):





Figure 15.14. Breakthrough curves in the adsorption of a mixture of hydrocarbons with composition n-butane 0.4 mol %, n-pentane 25.9, n-hexane 23.9 iso and cyclic hydrocarbons 49.8 mol % (Lee, in Recent Advances in Separation Science, *CRC Press, Boca Raton, FL, 1972, Vol. II, pp. 75–110*).

provided at the top to prevent disturbance of the top layer of adsorbent by incoming high velocity gas or entrainment by upflowing gases. When regeneration is by heating, a drawback of the ball support arrangement is their substantial heat capacity, which slows up the heating rate and subsequent cooling to process temperature.

Representative values and ranges of operating parameters are summarized in Table 15.3. Cycle times for some adsorptions are adjusted to work shift length, usually multiples of 8 hr, with valve adjustments made by hand. When cycle times are short, as for solvent recovery, automatic opening and closing of valves is necessary.



Figure 15.15. Capacity decline with service of a molecular sieve (*plant data, Davison* Sieve 562). Flow, 8150 kgmol; pressure, 3600 **kPa** (36 atm); temperature, 15°C; water content, 96 kg/hr; minimum cycle time, 24 hr. (*Chi and Cummings, 1978*).

Steam rates for regeneration of a particular adsorbent carbon are shown in Table 15.3(b). Steam/solvent ratios as high as 8 sometimes are necessary.

Data for liquid phase adsorption are typified by water treating for removal of small but harmful amounts of impurities. Some conditions are stated by Bemardin [*Chem. Eng., (18* Oct. 1976)]. Water flow rates are 5–10 gpm/sqft. When suspended solids are present, the accumulation on the top of the bed is backwashed at 15-20 gpm/sqft for 10–20 min/day. The adsorbent usually is not regenerated in place but is removed and treated in a furnace. Accordingly, a continuous operation is desirable, and one is simulated by periodic removal of spent adsorbent from the bottom of the vessel with a design like that of Figure 15.18(b) and replenishing of fresh adsorbent at the top. The pulses of spent and fresh carbon are 2-10% of the total bed. Height to diameter ratio in such units is about 3.

A carbon adsorber for handling 100,000 gal/day of water consists of two vessels in series, each 10 ft dia by 11 ft sidewall and containing 20,0001b of activated carbon. Total organic carbon is reduced from 650 mg/L to 25 mg/L, and phenol from 130 mg/L to less than 0.1 mg/L.

The capacity of regeneration furnaces is selected so that they operate 80-90% of the time. In multiple-hearth furnaces the loading is 70-80 lb/(sqft)(day). In countercurrent direct fired rotary kilns, a 6% volumetric loading is used with 45 min at activation temperature.

Details of the design and performance of other liquid phase adsorptions such as the Sorbex processes are proprietary.

15.5. ION EXCHANGE DESIGN AND OPERATING PRACTICES

Ion exchange processes function by replacing undesirable ions of a liquid with ions such as H^+ or OH- from a solid material in which the ions are sufficiently mobile, usually some synthetic resin. Eventually the resin becomes exhausted and may be regenerated by contact with a small amount of solution with a high content of the desired ion. Resins can be tailored to have selective affinities for

TABLE 15.4. Properties of Ion-Exchange Materials

(a) Physical Properties

		Bulk wot	Moisture	Swelling	Maximum		Exchange capacity	
Material	Shape • of particl es	density (drained), kg/L	(drained). % by weight	due to exchange, %	operating temperature, t	Operating pH range	Dry. equivalent/kg	Wet. equivalent/L
Cation eachangers: strongly acidic Polystyrene sulfonate Homogeneous (gel) resin 4% -lin ked 6% cross-linked 12% cross-linked 16% cross-linked 20% cross-linked Macroporous structure 10~12% cross-linked	5	0.75-0.85 0.76-0.86 0.77-0.67 0.78-0.88 0.79-0.69 0.80-0.90	64-70 56-65 48-60 44-46 42-46 40-45 50-55	IO-12 8-10 6-8 5 4 3 4-6	120-150	0-14	5.0-5.5 4.6-5.2 4.4-4.9 4.2-4.6 3.9-4.2 4.5-5.0	1.2-1.6 1.3-1.8 1.4-1.9 1.5-2.0 1.7-2.1 1.8-2.0 1.5-1.9
Sulfonated phenolic resin Sulfonated coal	G	0.74-0.85	50-60	7	50-90	0-14	2.9-2.5	0.7-0.9
Cation exchangers: weakly acidic Acrylic (pK 5) or methacrylic (pK 6) Homogeneous (gel) resin Macroporous Phenolic resin Polystyrene phosphonate Polystyrene aminodiacetate Polystyrene amidoaime Polystyrene thiol	s s G, s s s s s s	0.70-0.75 067-0.74 0.70-0.80 0.74 0.75 - 0 7 5 -0.75	45-50 50-55 ~50 50-70 66-75 58 45-50	20-80 10-100 L0-25 <40 <100 10	120 120 45–65 120 75 50 60	4-14 0-14 3-14 3-14 1-11 1-13	8.3-10 -8.0 2.5 6.6 2.9 26 - 5	3 3-4.0 2.5-3.5 1.0-1.4 3.0 0.7 0.8-0.9 2.0
Cellulose Phosphonate Methylene carboxylate Greensand (Fe silicate) Zeohie (Al silicate) Zirconium tungstate	F F, P, G G G G	1.3 0.85-0.95 1.15-1.25	1-5 40-45 - 5	0 0 0	60 60 >150	6-8 6-8 2-10	-7.0 -07 0.14 1.4 1.2	0 18 0.75 I O
Anion exchangers: strongly basic Polystyrene-based Trimethyl benzyl ammonium (type								
I) Homogeneous, 8% CL Macro porous, 11% CL Dimethyl hydroxyethyl ammonium (twee II)	S S	0.70 0.67	46-50 57-60	-20 15-20	60-80 60-80	0-14 0-14	3.4-3.8 3.4	1.3-1.5 1.0
Homogeneous, 8% CL Macroporous, 10% CL	S S	0.71 067	-42 -55	15-20 12-15	40- 80 40- 80	0-14 0-14	3.8-4 0 3.6	1.2 1.1
Acrylic-based Homogeneous (gel) Macroporous Cellulose-based	S S	072 067	-70 -60	- 1 5 - 1 2	40-80 40-80	0-14 0-14	-5.0 3.0-3.3	1.0-1.2 0.8-0.9
Ethyl trimethyl ammonium Triethyl hydroxypropyl ammonium	F				100 100	4-10 4-10	062 0.57	
Anion exchangers: intermediately basic (pK 11) Polystyrene-based Epoxy-polyamine	S S	0.75 0.72	- 5 0 - 8 4	15-2s 8-10	65 75	0-10 o-7	4.6 6.5	1.8 1.7
Anion exchangers: weakly basic (pK 9) Aminopolystyrene Homogeneous (gel) Mącroporous	s s	0.67 061	- 4 5 55-60	8-12 ~25	100 100	0-7 0-9	5.5 4.9	1.8 1.2
Acrylic-based amine Homogeneous (gel) Macroporous Cellulose-based	S S	072 0.72	-63 ~68	8-10 12-15	80 60	0-7 0-9	6.5 5.0	1.7 1.1
Aminoethyl Diethyl aminoethyl	P P						1.0 -0.9	

Shapes: C. cylindrical pellets; C. granules: P. powder; S. spheres. tWhentwo temperatures are shown, the first applies to H form for cation. or OH farm for anion. exchanger, the second to salt ion NOTE: To convert kilograms per liter to pounds per cubic t, multiply by 6.238 × 10¹; °F = % °C + 32.

(Chemical Engineers' Handbook, McGraw-Hill, New York, 1994; a larger table complete with trade names is in the 5th edition, 1973).

(b) Selectivity Scale for Cations on 8% Crosslinked Resin

Li+	1.0	Zn ²⁺	3.5	(c) Approximate S	electivity Scale fo	r Anions on Strong-Ba	se Resins
H ⁺ Na ⁺ NH: K ⁺ Bb ⁺	1.3 2.0 2.6 2.9 3.2	Co ²⁺ Cu ²⁺ Cd ²⁺ Be ²⁺ Mn ²⁺	3.7 3.8 3.9 4.0 4.1	I⁻ NO₃ Br⁻ HSO:	8 4 3 1.6	HCO; CH₃COO⁻ F⁻ OH-(Type_I)	0.4 0.2 0.1 0.05-0.07
Cs ⁺ Ag ⁺ UO ²⁺ Mg ²⁺	3.3 8.5 2.5 3.3	Ni ²⁺ Ca ²⁺ Sr ²⁺ Pb ²⁺ Ba ²⁺	3.9 5.2 6.5 9.9 11.5	NO; CN ⁻ Cl ⁻ BrO ₃ OH-(Type II)	1.3 1.3 1.0 1.0 0.65	so:- CO3 ⁻ HPO3 ⁻	0.15 0.03 0.01

(Bonner and Smith, J. Phys. Chem. 61, 1957, p. 326).

(Bonner and Smith, J. Phys. Chem. 61, 1957, p. 326).



Figure 15.16. A three-vessel drying system for a cracked light hydrocarbon stream. Valve operation usually is on automatic timer control. Recycled process gas serves as regenerant.

particular kinds of ions, for instance, mercury, boron, ferrous iron, or copper in the presence of iron. Physical properties of some commercial ion exchange resins are listed in Table 15.3 together with their ion exchange capacities. The most commonly used sizes are -20 + 50 mesh (0.8–0.3 mm) and -40 + 80 mesh (0.4–0.18 mm).

Rates of ion exchange processes are affected by diffusional resistances of ions into and out of the solid particles as well as' resistance to external surface diffusion. The particles are not really solid since their volume expands by 50% or more by imbibition of water. For monovalent exchanges in strongly ionized resins, **half** times with intraparticle diffusion controlling are measured in seconds or minutes. For film diffusion, half times range from a few minutes with 0.1N solutions up to several hours with 0.001N solutions. Film diffusion rates also vary inversely with particle diameter. A rough rule is that film diffusion is the controlling mechanism when concentrations are below 0.1–1.0N, which is the situation in many commercial instances. Then the design methods can be same as for conventional adsorbers.

Ion exchange materials have equilibrium exchange capacities of about 5 meq/g or 2.27 g eq/lb. The percentage of equilibrium exchange that can be achieved practically depends on contact time, the concentration of the solution, and the selectivity or equilibrium constant of the particular system. The latter factor is discussed in Section 15.2 with a numerical example.

Commercial columns range up to 6 m dia and bed heights from 1 to 6 m, most commonly 1-3 m. Freeboard of **50–100%** is provided to accommodate bed expansion when regenerant **flow** is upward. The liquid must be distributed and withdrawn uniformly over the cross section. Perforated spiders like those of Figure 15.19 are suitable. The usual support for the bed of resin is a bed of gravel or layers of ceramic balls of graded sizes as in Figure 17.27. Balls sometimes are placed on top of the bed to aid in distribution or to prevent disturbance of the top level. Since the specific volume of the material can change 50% or more as a result of water absorption and ion-ion exchange, the distributor must be located well above the initial charge level of fresh resin.

Liquid flow rates may range from 1 to 12 gpm/sqft, commonly 6-8 gpm/sqft. When the concentration of the exchange ion is less than 50 meq/L, flow rates are in the range of 15-80 bed volumes (BV)/hr. For demineralizing water with low mineral content, rates as high as 400 BV/hr are used. Regenerant flow rates are kept low, in the range of 0.5-5.0 BV/hr, in order to allow attainment of equilibrium with minimum amounts of solution.

The ranges of possible operating conditions that have been stated are very broad, and averages cannot be depended upon. If the proposed process is similar to known commercial technology, a new design can be made with confidence. Otherwise laboratory work must be performed. Experts claim that tests on columns 2.5 cm dia and 1 m bed depth can be scaled up safely to commercial diameters. The laboratory work preferably is done with the same bed depth as in the commercial unit, but since the active exchange zone occupies only a small part of a normal column height, the exchange capacity will be roughly proportional to the bed height, and tests with columns 1 m high can be dependably scaled up. The laboratory work will establish process flow rates, regenerant quantities and flow rates, rinsing operations, and even deterioration of performance with repeated cycles.

Operating cycles for liquid contacting processes such as ion exchange are somewhat more complex than those for gas adsorption. They consist of these steps:

- 1. Process stream Row for a proper period.
- 2. A rinse for recovering possibly valuable occluded process solution.
- 3. A backwash to remove accumulated foreign solids from the top of the bed and possibly to reclassify the particle size distribution.
- 4. The flow of regenerant for a proper period.
- 5. Rinse to remove occluded regenerant.

As complex a cyclic process as this may demand cycle times of more than a few hours. Very high ion concentrations or high volumetric rates may require batteries of vessels and automatic switching of the several streams, or continuously operating equipment. Several continuous ion exchange plants are being operated successfully. The equipment of Figure 15.20 employs pulsed transfer of solid between exchange and regenerant zones as often as every 4 min to every 20 or 30 min. Attrition of the resin may require replacement of as much as 30% of the resin each year in water conditioning applications.

Fluidized bed units such as the multistage unit of Figure 15.20 suffer from some loss of efficiency because the intense mixing eliminates axial concentration gradients. They do have the merit, however, of not being bothered by the presence of foreign solid particles.

The economic break between fixed bed and continuous operation has been estimated as ion concentrations of 0.5N, or flow rates above 300 gpm, or when three or more parallel beds are required to maintain continuous operation. The original application of continuous ion exchange was to treatment of radioactive wastes, but some installations of ordinary water treating have been made.

Resin requirements for two extremes of ion concentration are analyzed in Example 15.3. The high concentration stream clearly is a candidate for continuous ion exchange.

ELECTRODIALYSIS

In this process, dissolved electrolytes are removed by application of electromotive force across a battery of semipermeable membranes constructed from cation and anion exchange resins. The cation membrane passes only cations and the anion membrane only anions. The two kinds of membranes are stacked alternately and separated about 1 mm by sheets of plastic mesh that are still provided with flow passages. When the membranes and spacers are compressed together, holes in the corners form appropriate conduits for inflow and outflow. Membranes are 0.15-0.6 mm thick. A commercial stack may contain several hundred compartments or pairs of membranes in parallel. A schematic of a stack assembly is



Figure 15.17. Two designs of fixed bed gas adsorbers. (a) Vertical bed with balls on top for hold-down and distribution of feed (Johnson, Chem. Eng. 79, 87 (27 Nov. 1972)]. (b) Horizontal fixed bed for low pressure drop operation [*Treybal*, Mass Transfer Operations, *McGraw-Hill, New York, 1980; Logan, U.S. Pat. 5180,712* (1939)].



(b)

Figure **15.18.** Two types of supports for adsorbent beds [Johnston, Chem. Eng., (27 Nov. **1972**)]. See also Figures 17.23 and 17.24. (a) Common type of flat screen support. (b) Conical-type of support suited to frequent removal of adsorbent.

in Figure 15.21. Properties of commercially produced membranes are in Table 15.5 and performance data are in Table 15.6.

Membranes may be manufactured **by** mixing powdered ion exchange resin with a solution of binder polymer and pouring the heated mixture under pressure onto a plastic mesh or cloth. The concentration of the ion exchanger is normally 50–70%. They are chiefly copolymers of styrene and divinylbenzene, sulfonated with sulfuric acid for introduction of the cation exchange group.

Standard cell sizes are up to 30 by 45in. In an individual stack the compartments are in parallel, but several stacks in series are employed to achieve a high degree of ion exchange. The ion exchange membrane is not depleted and does not need regeneration. The mechanism is that an entering cation under the influence of an emf replaces an H^+ ion from the resin and H^+ from solution on the opposite face of membrane replaces the migrating cation.

Table 15.6 shows that pressures drops may be as high as 900 psi. Flow rates in a single stage are about 1 gal/(hr)(sqft of available membrane surface). The process is distinguished by very low power requirements: the desalination of sea water, for instance, consumes 11-12 kWh/1000 gal. One stage effects a reduction of about 50% in salt content, so several stages in series are used for high performance. A flow sketch of a three-stage electrodialysis plant is in Figure 15.21(c).

Like many other specialities, electrodialysis plants are purchased as complete packages from a few available suppliers. Membrane replacement is about 10% per year. Even with prefiltering the feed, cleaning of membranes may be required at intervals of a few months. The comparative economics of electrodialysis for desalting brackish waters is discussed by Belfort (1984): for lower salinities, electrodialysis and reverse osmosis are competitive, but for higher ones electrodialysis is inferior. Electrodialysis has a number of important unique applications, for removal of high contents of minerals from foods and pharmaceuticals, for recovery of radioactive and other substances from dilute solutions, in electro-oxidation reduction processes and others.

15.6. PRODUCTION SCALE CHROMATOGRAPHY

When a mixture of two substances is charged to a chromatographic column, one of them may be held more strongly than the other. Elution with an inert fluid will remove the more lightly held substance first, then the other. Separations even between very similar substances can be very sharp. Figure 15.22(a) is an example of a chromatogram. Only fluid-solid chromatography is an adsorptive process, but gas-liquid and liquid-liquid are used more frequently since liquids with suitable absorption properties are easier to find than solid adsorbents. The active sorbent is a high-boiling solvent deposited on a finely divided inert solid carrier. The process is one of absorption, but the behavior is much like that of adsorption. The principal application is to chemical analysis. Relative retention times on various sorbents are key data which are extensively tabulated, for instance in Meites (*Handbook of Analytical Chemistry*, McGraw-Hill, New York, 1963).

Chromatographic separations are necessarily intermittent with alternate injections and elutions, although a measure of continuity can be achieved with an assembly of several units, or with suitably sized surge tanks. A process flowsketch appears in Figure 15.22(b). Information on production scale chromatography is provided by Conder (1973). Only separations difficult to achieve by other means are economical with chromatography.

Individual drums are provided for each product fraction. A detector monitors the separation and provides signals for controlling the injection and collection sequence. The operation of partial condensers for the dilute eluted streams presents challenges because of aerosol formations. When a valuable carrier such as nitrogen is used, it must be cleaned up and recycled.

A 1968 estimate of the cost breakdown for a plant with a column 4 ft dia by 15 ft high and a throughput of 200-920 tons/yr has been converted to a percentage basis in Table 15.7 because of its age. The costs are said to not vary greatly with throughput or the nature of the separation, although this analysis has been made specifically for the separation of α - and β -pinenes. The temperature was 165°C and the solvent was Carbowax 20M. The design was based on data in a 4in. dia column which had a capacity of 200-1500 mL/hr.

Some of the materials for which chromatographic separation should be considered are essential oils, terpenoids, steroids, alkaloids, pharmaceuticals, metal chelates, isotopes, and close-boiling isomers. For easy separations, vacuum distillation, liquid-liquid extraction, and fractional crystallization are less expensive.

15.7. EQUIPMENT AND PROCESSES

Adsorbents are made in pellet form by extrusion or pressing or in granular form by crushing and classification of larger masses or in spherical or globular form by precipitation in an inert gaseous or liquid medium. Typical processes for some adsorbent preparations are represented in Figure 15.1. The BASF process of Figure 15.1(c)



Figure 15.19. The Toray simulated continuous adsorption process. (a) Showing the main valving for a seven-chamber adsorption system [*Otani et al., U.S. Put. 3, 761, 533, (25 Sep. 1973)*]. (b) Flowsketch for recovery of paraxylene by continuous adsorption [*Otani et al., Chem. Economy Eng. Rev. 3(6), 56-59 (1971)*].

for silica gel employs a special mixing nozzle in which water glass is mixed intermittently with a stream of sulfuric acid to form an unstable sol that is sprayed directly into air where the globules solidify immediately. Mixing time in the nozzle is 0.1-1.0 sec. The sodium sulfate is washed out in a tower to which the particles fall. The process of initial activation or reactivation is shown for bone char. In general a heating process, sometimes combined with oxidation, is required to drive gases out of the solid mass and thus to make it porous.

GAS ADSORPTION

The usual equipment for gas adsorption is a number of vessels containing **fixed** beds of the adsorbent, at least two vessels for achieving overall continuous operation, Figure 15.17 shows suitable vertical and horizontal vessels. The vertical ones are less likely to form channels and usually are favored. Bed depths as high as 45 ft are in use. Horizontal vessels are preferred when pressure drops

must be kept low, as in recovery of solvents from air in printing or paint establishments. Modes of support of granular beds are shown in Figures 15.18, 15.24, and 17.27.

A three-bed adsorption unit is illustrated in Figure 15.16. It is used to dry the feed to a distillation column with a top temperature of -70° F; thus a water **dewpoint** of -90° F is required. One of the vessels always is on regeneration and cooling down, and the other two in series on adsorption, with the more recently reactivated one downstream. A bleed off the process stream is diverted to use as regenerant. After the gas leaves the vessel being regenerated, the water is condensed out by cooling and the gas returns to the process downstream of a control valve that maintains a 10 psi differential.

Normally adsorption is conducted at or as near ambient temperature as possible and regeneration is at 350-500°F. A new process developed by Union Carbide Corp, the AHR process (adsorption heat recovery), allows the temperature to rise to 200-400°F as a result of the heat of adsorption and effects the desorption at comparable temperatures with a stream of inert gas 1





(c)





(d)

Figure 15.20. Fixed bed ion exchange vessels and arrangements. (a) Typical design of a water softener, showing bed support, distributor, and effluent collector. (b) Vessel with radial-type distributors and collectors *(Illinois Water Treatment Co.)*. (c) A double-dish underdrain system (*Permutit* Co.). (d) Some arrangements of vessels for cation and anion exchange.

EXAMPLE 15.3

Size of an Ion Exchanger for Hard Water

A hard water contains 120ppm of $C_{a}CO_{3}$, 90% of which is to be removed with a hydrogen exchange resin of capacity 5 meq/g. By the method of Example 15.1 it is ascertained that under these conditions 98% of H⁺ ion of the resin will be replaced by the Ca^{++} at equilibrium. The minimum amount of resin will correspond to the equilibrium value. That amount will be calculated for treating 100 gpm of water on a 24 hr cycle. The mol wt of $C_{a}CO_{3} = 100.06$.

resin capacity = 0.98(0.005)(100.06)= 0.490 lb CaCO₃/lb resin, CaCO₃ removed = $0.9(8.34)(100)(1440)(120)(10^{-6})$ = 129.7 lb/24 hr, resin needed = 129.7/0.49= 264.7 lb, or 4.71 cuft of resin with sp gr = 0.9.

such as nitrogen or carbon dioxide which is recycled after the water is condensed out. The process is applied to removal of as much as 20% of water from ethanol with cycle times of the order of 1 hr, instead of the common 24 hr, even for much smaller contents of water (Anon, *Chem. Eng.*, April 15, 1985, **p**. 17).

Continuous **fluidized** bed equipment has been utilized for gas adsorption, but usually attrition losses of comparatively expensive adsorbents have been prohibitive and the loss of **efficiency** because of **axial** mixing has been a serious handicap. Drying equipment such as those of Figure 9.13 presumably can be operated in reverse to recover valuable substances' from a vapor phase, and the forward mode applied for regeneration in associated equipment. Other possibly suitable fluidized bed configurations are those of the reactors of Figures 17.32(a), (c), and (d).

Moving bed gas adsorbers also have been proposed and used, patterned after moving bed gas oil crackers. In the Hypersorber of For comparison, the amount of resin needed to remove the Na^+ from a 3.5% solution of NaCl at the rate of 100 gpm in 24 hr will be found:

resin capacity = 5 meq/g = 0.005 lb mol/lb, Na⁺ removed = 0.035(8.34)(100)(1440)/58.5 = 718.5 lb mol/day.

Accordingly,

resin = 718.5/0.005 = 142,700 lb,

pointing out that a fixed bed unit on such a long cycle may not be practical for such a high concentration of ion to be exchanged.

Figure 15.23, flows of gas and solids are countercurrent in a single vessel. After saturation, the solid is stripped with steam and removed at the bottom of the tower, and gas is lifted to cooling and adsorption zones. The control mechanism for solids flow and typical performance for ethylene recovery from cracked gases also are shown with the figure. Partly because of attrition losses and the advent of competitive processes for ethylene recovery, the Hypersorber was abandoned after a few years. The simpler Nofsinger moving bed adsorber of Figure 15.24 also has not proved commercially attractive.

LIQUID PHASE ADSORPTION

A major application of liquid phase adsorption is to the removal of relatively small amounts of impurities or color bodies in water treating, sugar refining, and other processes. Both batch and



Figure 15.21. Electrodialysis equipment and processes. (a) View of the components of an electrodialysis stack (*Lacey*, 1978). (b) Flow pattern through an electrodialyzer for removal of NaCl from water (*Ionics Inc.*). (c) Electroreduction with the use of an ion exchange diaphragm. (d) Flowsketch of a three-stage electrodialysis for treatment of brackish water (Rogers, in Belfort, 1984).



Figure 15.21—(continued)

TABLE 15.5. Properties of Membranes for Electrodialysis

Manufacturer	Name of membranes	Membrane	Thickness (mm)	Capacity (meq/gm)	Electrical resistance ($\Omega \text{ cm}^2$ in 0.1 N NaCl)	Reinforcement
Ionac Chemical Co.	Ionac	MC-3 142	0.15	1.06	9.1	Yes
New Jersey		MC-3470	0.35	I.05	10.5	Yes
		MA-3 148	0.17	0.93	10.1	Yes
		MA-3475	0.40	I.13	23	Yes
		IM-12	0.13		4	Yes
American Machine and	A.M.F.	C-60	0.30	I.5	6	No
Foundry Connecticut		A-60	0.30	1.6	5	No
Ionics Inc.	Nepton	CR61 AZL 183	0.60	2.7	9	Yes
Massachusetts	<u>^</u>	AR III BZL 183	0.60	1.8	14	Yes
Asahi Glass Co. Ltd.	Selemion	CMV	0.15	I.4	6.1	Yes
Tokyo, Japan		AMV	0.14		4.0	Yes
Tokuyama Soda Ltd.	Neosepta	CL 25 T	0.16	I .8-2.0	3.5	Yes
Tokyo, Japan		AV 4 T	0.15	1.5-2.0	4.0	Yes
Asahi Chemical Industry	A.C.I. or	DK 1	0.23	2.6	6.5	Yes
Co. Ltd. Tokyo, Japan	Acipex	DA 1	0.21	1.5	4.5	Yes
Ben-Gurion University of	Neginst	NEGINST-HD	0.35	0.8	12	Yes
the Negev, Research &	0	NEGINST-HD	0.35	0.8	10	Yes
Development Authority		NEGINST-HC	0.2	1.6	6	No
Beersheva, Israel		NEGINST-HC	0.2	1.7	8	No

(Belfort, 1984).

TABLE 15.6. Performance of Electrodialysis Equipment on Treatment of 3000 ppm Brackish Water

	Single stack, MY II. four stages	Single stack, MK 111. three stages	Single stack, MY III, one stage	Three stacks in series. MY III
Typical hydraulic flow rate U.S. gal/24-h day U.S. gal/min Pressure drop at typical flow, lb/in* Number of membranes Size of membranes, in X in Total area of membranes, ft ² % total area available for transfer Approximate weight, lb Approximate overall height, including legs Demineralization per pass (25°C, high-Clwater, typical flow), %	16.700 11.6 47 540 18 X 20 1,350 62 1,300 4'6" 88.5	55,600 38.6 44 900 18 X 40 4,500 64 2,800 6'10" 88.3	166,700 116 14 900 18 X 40 4,500 64 2,800 6'8" 52	166,700 116 2.7: 18 X 40 13.506 64 8,400 6'8" 90.0
Current required for 3000-ppm feed, A	Stages 1 and 2: 19 Stages 3 and 4: 8	Stages 1 and 2: 36 Stage 3: 12	46	Stage 1: 46 Stage 2: 24 Stage 3: 12
Voltage required for 3000-ppm feedt	Stages l and 2: 180 Stages 3 and 4: 150	Stages 1 and 2: 350 Stage 3: 150	640	stage 1: 640 Stage 2: 509 Stage 3: 426
Direct-current kW/stack for 3000-ppm feedt	4.6	14.1	29	Stage 1: 29 Stage 2: 12 stage 3: 5
Direct-current kWh/1000 gal product for 3000-ppm feed	7.4	6.8	4.7	5. age 5. 5 7.4

"lonics, Incorporated, Watertown, Mass. 1979. These units use the EDR process, in which polarity and fluid flow are periodically reversed. In general, addition of acid and antiprecipitant to the feed is not necessary in this process.

For typical brackish water containing a high proportion of sodium chloride. Approximately 10% of flow wasted during reversal.

(Spiegler, 1984).

continuous equipment are illustrated in Figure 15.25. The batch process consists of **slurrying the** liquid with powdered adsorbent and then separating the two phases by filtration. The saturated adsorbent--carbon from water treating or fullers earth from oil treating-is regenerated by ignition as in the block diagram of Figure 15.1(b), or sometimes by treatment with suitable reactive solvents such as sodium hydroxide for adsorbed phenol from water. In the semicontinuous process of Figure 15.25(b), pulses of adsorbent are withdrawn periodically from the bottom and fresh

material is charged in at the top. The pulses are 2-10% of the volume of the bed. Some data of adsorbent treating of water were given in Section 15.4. Attrition losses in moving beds for liquid treating are less than for gas treating. In the similar process of ion exchange of Figure 15.25(a), ion exchange losses of 30% per year are mentioned.

The successful simulation of continuous moving bed adsorption process developed by UOP (Universal Oil Products) is illustrated in Figure 15.26. For the process being simulated, part (a) of the figure



Figure 15.22. A process for recovering uranium from mine waters. The absorption column is 2.16 m dia, water flow rate is 28.5 m/hr, resin transfer off the top of the absorption column is 87 L every 3 hr, inlet concentration 3-6 mg U/L, outlet 0.002-0.009 mg U/L (*Himsley and Bennett, in Naden and Streat, 1984, pp. 144-168; U.S. Pat.* 4,018,677).

TABLE 15.7. Economic Data for a Chromatographic Process with Throughput of 400-920 tons/yr, with Column 4ft die by 15ft high*

Equipment Cost	Percent	Annual Operating Cost	Percent
Feed preparation and injection	9.4	Maintenance and taxes	19.9
Column	13.1	Operating labor	13.7
Detection and control	2.6	Utilities and supplies	5.7
Fraction collection and heat exchange	16.7	Packing replacement	46.8
Carrier recycle	11.2	Depreciation (10year)	19.9
Process piping and building Engineering and construction	18:5 28:5		100
	100		

'Data are given only on percentage bases because of their age. (Abcor Inc., 1968).





Compositions, vol % Component Feed gas Bottoms Overhead Purge gas Hydrogen. 39.8 31.6 61.8 . 1.7 1.4 2.5 Nitrogen monoxide 0.9 0.8 1.3 0.1 0.1 0.2 0.1 Methane..... 51.3 66.1 33.7 dioxide. 0.2 2.9 0.1 Acetylene. 0.2 3.6 5.8 92.7 0.4 Ethylene. Ethane Tr. 0.7 100.0 100.0 100.0 100.0

(c)

Figure **15.23.** Hypersorber continuous moving bed gas phase adsorption system *(See Mantell, Adsorption, McGraw-Hill, New York, 1951).* (a) Schematic pattern of flows of gas and solid adsorbent *(Hengstebeck, Petroleum Processing, McGraw-Hill, New York, 1959).* (b) Solids flow rate control mechanism. (c) Typical separation performance.



Figure 15.24. Nofsinger continuous moving bed adsorber [Spangler and Price, U.S. Pat. 3, 442, 066 (6 May 1969)].

shows flows of adsorbent and fluids and the composition profiles along the tower. The simulated process employs 12 fixed beds in a single vessel, in which input and output streams are individually controlled. The points of entry and withdrawal of the four external streams-feed, extract, raffinate, adsorbent-are controlled with a single special rotary valve. Periodically each stream is switched to the adjacent bed so that the four liquid access positions are always maintained the same distance apart. Satisfactory operation is assured by uniform feeds and withdrawals and flushing of lines between their uses for regeneration and other purposes. The internal constructions of the tower, such as the mechanism of feed and withdrawal at individual beds, are not revealed in the literature. As of 1984, some 60 large capacity installations for various hydrocarbon isomer separations with molecular sieves have been made. The largest column mentioned is 22 ft dia. The distribution across the cross section has been worked out so that scale-up from 3 in. to commercial size is reliable. The process is described briefly in articles by Broughton (1978; 1984) and in several patents listed in the first of these articles.

A variant of this process developed by Toray industries is

illustrated in Figure 15.19. The beds are individual rather than in a single shell and utilize individual on-off valves. One application, to the separation of aromatics, is mentioned in the literature [Otani, *Chem. Eng.* **80**(9), 106 (17 Sep. 1973)].

ION EXCHANGE

Because of the large volumes of dilute electrolytes that sometimes need to be treated, continuous processing with ion exchange materials is more common than liquid phase adsorption, although fixed bed processes still are predominant. Typical arrangements of fixed beds appear in Figure 15.20. Any particular ion exchange resin is capable of exchanging only cations or anions. The two kinds of resins may be mixed and incorporated in the same vessel or they may be used separately in their own vessels. Cation exchange resins may be strongly or weakly acid, and anion exchange resins, strongly or weakly basic. The choice of an ion exchange system depends on the composition of the feed, the product quality required, the scale of the operation, and the economics of the process. Three of the many possible arrangements of vessels are sketched in Figure 15.20(d). Series combinations of vessels are employed when leakage is highly undesirable. The inlet to the last stage is monitored and the information is taken as a guide to transfer of the first vessel in line to regeneration.

All of the continuous processes of Figures 15.27 and 15.22 employ intermittent transfer of spent resin out of the primary vessel to regeneration facilities. Although all of the operations of exchange, rinsing, and regeneration can be performed in elegantly designed equipment such as Figure 15.27(a), greater flexibility is inherent in a multivessel plant such as Figure 15.22. This is of an operating plant for which some size and operating data are given here, and more appear in the reference article.

Performances of four fluidized bed ion exchange plants are described by Cloete (in Naden and Streat, 1984, pp. 661-667). One of the exchange columns is 4.85 m dia, has 12 stages each 1 m high, with perforated trays having holes 12mm dia with a capacity of $640 \text{ m}^3/\text{hr}$ of uranium mine waters.

Ion exchange resins are used widely as heterogeneous catalysts of processes that require acid or base catalysis, for example, hydration of propylene to isopropanol, reaction of isobutylene with acetonitrile, and many others. The same kind of equipment is suitable as for ion exchange, but usually regeneration is not necessary, although some degradation of the resin naturally occurs over a period of time.

ION EXCHANGE MEMBRANES AND ELECTRODIALYSIS

Only one basic design of electrodialysis equipment for demineralization appears to be in use. This is an assembly of alternate cation and anion ion exchange sheets separated by spacers in groups of several hundred clamped together between electrodes. The assembly physically resembles a plate-and-frame filter press. Figures 15.21(a) and (b) show such assemblies, and some dimensional data were stated in Section 15.5, Electrodialysis.

The concentrates of salt solutions made by electrodialysis of seawater are suited as feed to the evaporators of salt manufacturing plants with considerable savings in overall energy requirements. Other applications also are based on the concentrating effects of electrodialysis, for instance, tenfold increases of concentrations of depleted streams from nickel and copper plating plants are made routinely.

In applications such as the electroreduction of Figure 15.21(c) and electrolysis of brine to caustic and chlorine, single membranes serve as diaphragms between electrodes, permitting passage of ions but retarding **diffusion** and mixing of feed and product solutions.



Figure 15.25. Liquid phase adsorption processes for water treated with activated carbon and petroleum treated with clay adsorbents. (a) A two-stage slurry tank and filter process. (b) Continuous pulsed bed operation, individual pulses 2-10% of bed volume as needed.



Figure 15.26. Continuous and UOP simulated continuous moving bed liquid adsorption processes [*Broughton*, Sep. Sci. Technol. 19, 723-736 1984–1985)]. (a) Continuous moving bed liquid adsorption process flows and composition profiles. (b) UOP Sorbex simulated moving bed adsorption process.



(b)

Figure 15.26—(continued)



Figure 15.27. Continuous ion exchange equipment. (a) The Higgins moving bed unit; the consolidated resin bed is recirculated upwards with a hydraulic pulse. [U.S. Pat. 3,580,842, (2.5 May 1942)]. (b) Operating sequence in the Higgins contactor. The double-acting piston simultaneously sucks liquid from the top of the column and delivers it to the bottom: (a) solution pumping for several minutes; (b) resin movement for 3-5 sec; (c) solution pumping for several minutes (*Coulson and Richardson*, Chemical Engineering, Pergamon, New York, 1979, Vol. 3, p. 520). (c) In the Asahi system, resin is transferred between the adsorber and the regenerator at 10-60 min intervals (Asahi Chem. Ind., U.S. Pat. 3, 152, 072). (d) The Himsley multistage slurry adsorber; the resin is pumped as a slurry, one stage at a time into and from the regenerator [Himsley, Can. Pat. 980,467, (23 Dec. 1975)].





Figure 15.27—(continued)

CHROMATOGRAPHIC EQUIPMENT

The components of a commercial process employing **chromato**graphic separation are represented on the flowsketch of Figure 15.28. The process is intermittent with very short cycles. The cost breakdown of a plant for the separation of α - and β -pinenes is given in Table 15.7, which is based on pilot plant work in a 4 in. dia column. That company is no longer in that business; thus the test

data are not available, and the operating conditions are not known. Other data for the same separation, however, are presented in Table 15.8. The largest column considered there, 0.4 m dia, has a production rate of 4300 kg/(m²)(day), with cycle times of about 1 min. Valentin (1981) refers to literature where columns of 1 and 2 m dia are described and mentions that diameters up to 5 m are feasible and may have throughputs of 10,000 metric tons/yr.



Figure 15.28. Chromatographic separations. (a) Typical chromatogram produced by gas-liquid chromatography. (b) Flowsketch of a production scale chromatographic unit [Ryan, *Timmins, and O'Donnell*, Chem. Eng. Prog. 64, 53 (Aug. 1968)].



Figure 15.28—(continued)

TABLE 1	5.8. Examples	of Chromatographic	Separations	in a Colur	nn 125 mm dia	1
(a) Five	Separations					

Case	Mixture	Charge (mol %)	Product (mol %)	Temperature (°C)/ Pressure (Torr)	Remarks
1	Pentane	99.3	99.995	225/760	extreme purification
	Isopentane	0.7	50 ppm		
2	Benzyl alcohol	99.5	99.6	150/50	selective removal of
	Benzyl aldehyde	0.1			small impurities
	Others	0.4			
3	α-Pinene	70.0	99/2	160/760	simultaneous purification
	β-Pinene	30.0	1/98		of two materials
4	Essence of cloves			220/760	recovery of a thermally
	Eugenol	75	99.6		sensitive natural product
5	Bromo-3-thiophene	91.6	99.0	150/760	a difficult separation
	Bromo-2-thiophene	4.6	1.0		of position isomers
	Others	3.6			·

(Bonmati and Guiochon, Parfums, Cosmetiques, Aromes, 37-59, Sept/Oct 1976).

(b) Details of the Separation of a-Pinene and $\beta\text{-Pinene}$

Operating Conditions	Other Data		ELF–SRTI Proces		
Column					
diameter (mm)	100	305	125	400	
Length (m)	2.7	unknown	1.5	1.5	
Baffles	yes	yes	n o	n o	
Programmer	peak	deflection	time-ba	ased	
Carrier gas	Нe	Нe	Нe	H₂	
Velocity (cm/sec)	9.2	_	9.5	9	
Temperature (°C)	160	160	160	160	
Cycle (sec)	60	-	60	65	
Purity	96	.5-96.6	99.1-	97.6	
Productivity (kg/day)	19	160	40	540	
REFERENCES

Adsorption

- R.A. Anderson, Adsorption (general), Encycl. Chem. Process. Des. 2, 174-213 (1977).
- G.B. Broughton, Adsorptive separation (liquids), *Encycl. Chem. Technol.* 1, 563-581 (1978).
- D.B. Broughton, Production scale adsorption of liquid mixtures by simulated moving bed technology, Sep. Sci. Technol. 19, 723-736 (1984-1985).
- 4. C.H. Chi and W.P. Cummings, Adsorptive separation (gases), *Encycl.* Chem. Technol. 1, 544-563 (1978).
- 5. J.R. Fair, Sorption processes, Chem. Eng., 90-110 (14 Jul. 1969).
- A.L. Hines and R.N. Maddox, Mass Transfer Fundamentals and Applications, Prentice-Hall, Englewood Cliffs, NJ, 1985.
- J.L. Kovach, Gas adsorption, in *Handbook of Separation Processes for* Chemical *Engineers*, Schweitzer, (Ed.), McGraw-Hill, New York, 1979, pp. 3.3-3.47.
- M.N.Y. Lee, Novel separations with molecular sieve adsorption, in *Recent Developments in Separation Science* II, CRC Press, Boca Raton, FL, 1972, pp. 75–110.
- C.M. Lukchis, Adsorption systems, *Chem. Eng.*, (11 June 1973); (9 July 1973); (6 Aug. 1973).
- 10. D.M. Ruthven, Principles of Adsorption and Adsorption Processes, Wiley, New York, 1984.
- T. Vermeulen et al., Adsorption design, Encycl. Chem. Process. Des. 2, 162-174 (1977); Adsorption separation, Encycl. Chem. Technol. 1, 531-544 (1978); Adsorption and ion exchange, in Chemical Engineer's Handbook, McGraw-Hill, New York, 1984, Sec. 16.

ion Exchange

 R.E. Anderson, Ion exchange separation, in Handbook of Separation Techniques for Chemical Engineers, (Schweitzer, Ed.), pp. 1.359-1.414.

- D. Naden and M. Streat, Eds., *Ion Exchange Technology*, Horwood, Chichester, England, 1984.
- 3. Vermeulen et al., Ref. A.ll.
- R.M. Wheaton and E.J. Lefevre, Ion exchange, Encycl. Chem. Technol. 13, 678-705 (1981).

Electrodialysis and Ion Exchange Membranes

- 1. G. Belfort, Ed., Synthetic Membrane Processes, Academic, New York, 1984.
- 2. D.S. Flett, Ed., Ion Exchange Membranes, Horwood, Chichester, England, 1983.
- R.E. Lacey, Ion exchange separations, in Schweitzer, Ed., Handbook of Separation Techniques for Chemical Engineers, McGraw-Hill, New York, 1979, pp. 1.449–1.465.
- K.S. Spiegler, Electrodialysis, in *Chemical Engineers' Handbook*, McGraw-Hill, New York, 1984, pp. 17.37-17.45.
- 5. H. Strathmann, Electrodialysis and its application in the chemical process industry, *Sep. Purification Methods* **14**(1), 41-66 (1985).

Chromatography

- R. Bonmati and G. Guiochon, Parfums Cosmetiques Aromes, 37-59 (Sep./Oct. 1976).
- J.R. Conder, Production Scale Gas Chromatography, in New Developments in Gas Chromatography, H. Purnell, (Ed.), Wiley, New York, 1973.
- A.E. Rodrigues and D. Tondeur, Eds., *Percolation Processes: Theory* and Applications, Sijthoff and Noordhoff, Alphen aan den Rijn, Netherlands, 1981.
- 4. P. Valentin, in Rodrigues and Tondeur, loc cit., pp. 141-196.

issolved or molten substances are recoverable in solid form by precipitation upon cooling or upon removal of the solvent or by addition of precipitating agents. For convenience a distinction is made between two kinds of processes:

- In solution crystallization, the crystals are separated away from a so/vent, often water. In the case of inorganic solids particularly, the operating temperature is far below their melting points.
- In me/t crystallization, two or more substances of comparable melting points are separated by some degree of cooling. The degree of completeness of such separations depends on the phase equilibrium relations. When the crystals must be refined to remove occluded substances, the recovered material may leave the process in molten form. Subsequent/y, it may be solidified as flakes or sprayed granules.

The design of crystallizers is based on know/edge of phase equilibria, *solubilities*, rates and amounts of nuclei generation, and rates of crystal growth. Each system is

unique in most of these respects and not often predictable. The kind of information needed for design of a continuous crystallizer is indicated by the data supplied for Example 76.1.

Although theoretical advances are being made, the current state of the art of crystallization requires pilot p/ant evaluation of parameters of equations and of such operating variables as

- 1. crystal size distribution,
- 2. effects of impurities and additives,
- 3. residence time,
- 4. circulation rate,
- 5. mixing efficiency,
- 6. allowable degrees of supersaturation or subcooling,
- 7. heat transfer characteristics,

and others peculiar to the particular kind of equipment under consideration for the full scale p/ant.

This chapter will discuss the main concepts associated with crystallization practice, and will describe the main types of equipment used nowadays, together with some indications of their performance and applicability.

16.1. SOLUBILITIES AND EQUILIBRIA

The variation of the solubilities of most substances with temperature is fairly regular, and usually increases with temperature. When water is the solvent, breaks may occur in solubility curves because of formation of hydrates. Figure 16.1(a) shows such breaks, and they can be also discerned in Figures 16.2(b) and (c). Unbroken lines usually are well enough represented by second degree polynomials in temperature, but the Clapeyron-type equation with only two constants, $\ln x = A + B/T$, is of good accuracy, as appears for some cases on Figure 16.1(b).

A convenient unit of solubility is the mass of solute per unit mass of solvent, or commonly g solute/100 g solvent. Interconversions with molal units and mol fractions are made readily when densities of the solutions are known.

Under quiescent conditions a concentration substantially in excess of normal solubility or a temperature lower than the normal saturation temperature can be maintained. The maximum supersaturation appears to be a fairly reproducible quantity, but is reduced or even eliminated by stirring or by the introduction of dust or seed crystals. Some data are shown in Figure 16.1(c) and in Table 16.1. They are expressed as $AC = C - C_{sat}$ or as AC/C,..., or as $AT = T - T_{sat}$. According to the data of Table 16.1(d), subcooling correlates roughly with the heat of solution. The increments AC and AT can be quite substantial quantities.

The several regions of varying stability are represented by Figure 16.1(d). At concentrations above or temperatures below those represented by the supersaturation line, nuclei form and crystals grow spontaneously, although the rates of these processes do depend on the depth of penetration of the unstable region. Little control can be exercised on behavior in this region. In the metastable region, growth of crystals will occur even under quiescent conditions when dust or seeds are introduced and nuclei can be generated by agitation. Behavior in the metastable region is

largely controllable so that it is the practical operating region for production of crystals of significant sizes.

Practically feasible extents of supersaturation or subcooling are fairly small and depend on the substance and the temperature. Some data appear in Table 16.2. Since the recommended values are one-half the maxima listed, they rarely are more than $2^{\circ}C$ or so. This means that very high circulation rates through heat exchangers are needed. Thus, in the urea process of Example 16.1, the temperature rise is $2^{\circ}F$, and the volumetric circulation rate is about 150 times the fresh feed rate.

PHASE DIAGRAMS

Equilibria between liquid and solid phases over wide ranges of temperature are represented compactly on phase diagrams. The effect of moderate pressure on condensed phases is negligible. Aqueous systems often are complicated by the formation of hydrates, and other substances also may form intermolecular compounds. Of the substances of Figure 16.2, KC1 does not form a hydrate, but **NaCl** and **MgSO**₄ do. Mixtures always have lower melting points than those of the pure components. The lowest temperature and the corresponding composition at which a liquid phase can be present identify the eutectic ("easy melting"), for example, point C on Figure 16.2(a) and point *B* on Figure 16.2(b). Binary and ternary eutectics also are identified on the ternary diagram [Fig. 16.2(f)].

The effects of evaporation or chilling on the amounts and compositions of the liquid and solid phases can be followed on the diagrams. Example 16.2 does this. Mixtures that form eutectics cannot be separated completely by chilling. The amount and nature of a separated solid phase depends on the temperature and the overall composition. Examples 16.2(c) and (d) make such calculations. Mixtures that are completely miscible in both liquid and solid phases, such as Figure 16.2(d), can be separated

EXAMPLE 16.1

Design of a Crystallizing Plant

A plant is to make 10,000 lb/hr of urea crystals from a solution that contains 75% dissolved salt. The material balance and operating conditions are shown on the sketch. Key crystallization data are given by Bennett (1981, **p**. 452) as

1. The residence time is 3.4 hr.

2. The temperature rise across the heater is 2°F.

Other information deduced from pilot plant work is:

- 3. The feed contains 75% solids, but 1200 lb/hr of wash water from the centrifuge also is returned to the crystallizer.
- 4. The liquor contains 66.8% dissolved urea and has a specific gravity of 1.17 at the operating temperature of 130°F.
- The slurry contains 28 lb crystals/cuft and has a specific gravity of 1.354.
- 6. A purge stream amounting to 7% of the feed liquor is withdrawn as shown on the sketch.
- The pressure is 60 Torr, at which the saturation temperature of steam is 106°F. The superheat of 24°F is neglected in figuring the vapor density and velocity.
- 8. Depth of liquid in the vessel should not exceed 10 ft and the vapor velocity should not exceed that given by the equation, $u = 0.06\sqrt{\rho_L/\rho_g} 1$. A wire mesh deentrainer is not feasible because of encrustation.
- 9. Heat capacity of the solid is 0.62 Btu/(lb)(°F) and the heat of crystallization is 104 Btu/lb.
- 10. For sizing the vacuum ejector, air leakage is estimated at 25 lb/hr and carbon dioxide is 34 lb/hr.
- 11. The coefficient of heat transfer in the exchanger is 200 Btu/(hr)(sqft)(°F).

Calculations:

vapor rate = 4471(296)/3600 = 367.6 cfs, slurry holdup = 10,000(3.4)/28 = 1214.3 cuft, $u_{max} = 0.06\sqrt{84.5/0.0034} - 1 = 9.46$ fps, $D_{min} = \sqrt{367.6/9.46(\pi/4)} = 7.03$ ft.

The corresponding liquid depth is

$$h = 1214.3/(\pi/4)D^2 = 31.0$$
 ft,

which is too great a value.

Try D = 12.5 ft:

dished head capacity = 152 cuft (Figure 18.5), straight side = $(1214.3 - 152)/(12.5)^2(\pi/4) = 8.66$ ft, say 9.0 ft.

Together with the depth of liquid in the dished head, the total depth will be close to the 10 ft specified as the maximum. From Figure 18.5, a free board of 5.5 ft is adequate in the absence of a deentraining pad. Accordingly, the vessel will have a diameter of 12.5 ft, a straight side of 14.5 ft, and dished heads designed for full vacuum. The sketch is to scale.

Sufficient data are given for finding the heat balance and the liquor circulation rate, and for sizing the auxiliaries such as lines, pump, heat exchanger and vacuum system, but those calculations will not be made.



essentially completely in multistage equipment, although such processes are not often feasible. The possible extent of separation of multicomponent mixtures can be interpreted with a phase diagram like those of Figure 16.2(f) and Example 16.3. Phase diagrams are fairly plentiful, but published ones usually seem to be of the system they were interested in and not of the one you are interested in. Fortunately, nowadays phase diagrams can be developed at moderate cost and expenditure of time with differential scanning calorimeters.

Estimates of phase diagrams can be made on the assumption of ideal behavior or with activity coefficient data based on binary measurements that are more easily obtained. In such cases, clearly, it should be known that intermolecular compounds do not form.

The freezing behaviors of ideal mixtures over the entire range of temperatures can be calculated readily. The method is explained for example by Walas (*Phase Equilibria in Chemical Engineering*, Butterworths, Stoneham, MA, 1985, Example 8.9).

ENTHALPY BALANCES

Although the thermal demands of crystallization processes are small compared with those of possibly competitive separation processes such as distillation or adsorption; nevertheless, they must be known. For some important systems, enthalpy-composition diagrams have been prepared, like those of Figure 16.3, for instance. Calculations also may be performed with the more widely



Figure 16.1. Solubility relations. (a) Linear plot of solubilities against temperature (*Mullin*, 1972). (b) Solubility against temperature plotted according to the equation $z = \exp(A + B/T)$ (*Mullin*, 1972). (c) Normal and supersolubilities of two salts (*data collected by Khamskii*, 1969). (d) Identification of regions on solubility plots. In the unstable region, nucleation and growth are spontaneous. In the metastable region growth can occur on externally introduced particles. Along *u*-*d* to the left or along *u*-*d'* upwards, nucleation and growth can start at c or c', but a substantial nuclei growth rate will not be achieved until *d* or *d'* are reached.

available data of heat capacities and heats of solution. The latter are most often recorded for infinite dilution, so that their utilization will result in a conservative heat balance. For the case of Example 16.3, calculations with the enthalpy-concentration diagram and with heat of solution and heat capacity data are not far apart.

16.2. CRYSTAL SIZE DISTRIBUTION

Crystal size distribution (CSD) is measured with a series of standard screens. The openings of the various mesh sizes according to the U.S. Standard are listed in Example 6.6, and according to the British Standard in Figure 16.4. Table 12.1 is a complete listing. The size of a crystal is taken to be the average of the screen openings of successive sizes that just pass and just retain the crystal.

The cumulative wt % either greater or less than a specified screen opening is recorded. The amount of a size less than a

particular screen opening and greater than the next smaller size is called the differential amount. Typical size distribution data on Figure 16.4 are plotted in two cumulative modes, greater than or less than, and as differential polygons or histograms. For some purposes the polygon may be smoothed and often is shown that way. Some theoretical cumulative and differential distribution curves of similar nature are shown in Figure 16.5; the abscissas are proportional to the crystal length.

Cumulative data often are represented closely by the Rosin-Rammler-Sperling (RRS) equation

$$y = 100 \exp[-(d/d_m)^n],$$
 (16.1)

where *d* is the diameter, d_m is a mean diameter corresponding to y = 100/e = 36.8% and *n* is called the uniformity factor. The greater *n*, the more nearly uniform the distribution. The log-log plot of this



Figure 16.2. Some phase diagrams. (a) The water end of the system potassium chloride and water. (b) The water end of the system sodium chloride and water. (c) The water end of the system magnesium sulfate and water; the heptahydrate goes to the mono at 150°C, and to anhydrous at 200°C. (d) β -methylnaphthalene and β -chloronaphthalene form solid solutions. (e) Mixtures of formamide and pyridine form a simple eutectic. (f) These mixtures form binary eutectics at the indicated temperatures and a ternary eutectic at mol fractions 0.392 dibenzyl, 0.338 diphenyl, and 0.27 naphthalene.

TABLE 16.1.	Data of	Supersatura	tion and	Subcooling	of Solutions
(a) Maximum	Supersa	aturation of	Solutions	at 20°C, β	$= \Delta C/C_{\rm h}$

Solute	Fovbin and Krasnova's Jata [144] Gorbachev and Shlykov's data	isher's data 152]*
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.095 0.39 0.05€ 0.102 0.029 - 0.36 1.08 0.10 - 0.64 - 0.93 0.37 0.41 - 0.93 0.37 0.41 - 0.93 0.37 0.41 - 0.43 - 0.54 - - 2.71 1.60	

*Fisher's results were ok alined at 25°C.

(c) Maximum Supercooling of Salt Solutions at Various Temperatures

Salute	Heat of solu- tion λ, cal/mole	to−t, °C	λ(t ₀ - t), cal-mole ⁻¹ , deg ⁻¹
KC1 KBr KI KBrO3 KBrO3 KBrO3 KBrO3 KC103 KC103 KC104 KC105 KC104 KC105 KC105 NaNO3 NaC103 NaC1 NAC1 NH4C1 NH4C1 NBC104 NBC105 NBC104 NBC105 NBC104 NBC105 NBC104 NBC105 NBC104 NBC105 NBC104	4046 5080 5110 9760 9950 8800 12100 6100 5600 12200 3880 2370 3300 2370 3300 6360 9400	$\begin{array}{c} 19.6\\ 16.3\\ 15.5\\ 8.8\\ 13.5\\ 6.6\\ 13.0\\ 13.0\\ 13.0\\ 13.0\\ 12.0\\ 51.0\\ 20.0\\ 24.0\\ 10.3\\ 25.0\\ 36.7\\ 25.0\\ 36.7\\ 20.0\\ 9.0\\ \end{array}$	78897 80804 79205 84788 91490 65670 114400 76230 79300 65399 67200 62220 77600 56880 65016 82500 80925 72000 76320 84660

equation should be linear. On Figure 16.4(c) the scatter about the straight line is small, but several of the plots of commercial data of Figure 16.6 deviate somewhat from linearity at the larger diameters.

Two other single numbers are used to characterize size distributions. The median aperture, MA or d_{50} , is the screen opening through which 50% of the material passes. The coefficient

(b)	Temperature	D	epeno	dence	of	the	Maximum
Su	persaturation	of	Salt	Soluti	ions	5	

Solute	۰, °C	C ₀ , moles/liter	C. moles/liter	$a = C - C_0$	β
KNO,	0	1.25	2.03	0.78	0.62
	10	1.96	2.78	0.81	0.41
	20	2.76	3.75	0.99	0.36
	30	3.83	4.84	I.01	0.26
	40	4.97	6.00	1.03	0.20
кс1	0	3.33	3.88	0.55	0.16
	10	3.72	4.12	0.40	0.11
	20	4.03	4.42	0.39	0.095
	30	4.29	4.45	0.16	0.037
	40	4.45	4.58	0.13	0.029
ксіо _з {	10 20 30 40	0.40 0.58 0.90 1.11	0.65 0.82 1.05 1.32	0.25 0.24 0.25 0.21	$\begin{array}{c} 0.62 \\ 0.41 \\ 0.32 \\ 0.19 \end{array}$
K ₂ CrO ₄	10	2.68	2.96	0.28	0.11
	20	2.74	3.00	0.26	0.093
	30	2.82	3.03	0.21	0.073
	40	2.98	3.07	0.19	0.065

(d) Dependence of the Maximum Supercooling of Solutions on Heat of Solution

Solute	ło, °C	1, °C	0 =t₀t, °C	Solute	to, °C	t, °C	8 <i>=t</i> 0−t, °C
KNO3	20 30 40 50 60 70	-1.0 8.9 18.9 28.8 38.8 48.9	21.0 21.1 21.1 21.2 21.2 21.2 21.1	KBr	10.5 20 30 40 50 60	-1.8 8.0 17.8 28:0 37.9 47.7	12.3 12.0 12.2 12.0 12.1 12.3
ксі {	50 60 70 80 90	6.7 16.6 26.7 36.7 46.6	43.3 43.4 43.3 43.3 43.4	I K ₂ SO ₄ (80 90 100	67.7 3.0 13.0	12.3 87.0 87.0

(Khamskii, 1969).

of variation is defined by the equation

$$CV = 100(d_{16} - d_{84})/2d_{50}.$$
 (16.2)

The origin of this concept is that the fraction of the total area under a normal distribution curve between the 16 and 84% points is twice the standard deviation. The smaller CV, the more nearly uniform the crystal sizes. Products of DTB crystallizers, for instance, often have CVs of 30–50%. The number is useful as a measure of consistency of operation of a crystallizer. Some details are given by Mullin (1972, pp. 349, 389).

TABLE 16.2. Maximum Allowable Supercooling AT (°C) and Corresponding Supersaturation AC (g/100 g water) at 25°C^a

Substance	A T	A C
NH, alum	3.0	1.0
NH₄CI	0.7	0.3
NH₄̃NO ₃	0.6	3.0
$(NH_4)_2SO_4$	1.8	0.5
NH4H2PO4	2.5	2.3
CuSO ₄ .5H ₂ O	1.4	1.0
FeSO₄.7H ₂ O	0.5	0.6
K alum	4.0	1.0
KBr	1.1	0.6
KCI	1.1	0.3
KI	0.6	0.4
KH₂PO₄	9.0	4.6
KNO,	0.4	0.6
KNO,	0.8	0.8
K ₂ SO ₄	6.0	1.3
MgSO ₄ .7H ₂ O	1.0	1.3
NiSO₄.7H₂O	4.0	4.4
NaBr.2H ₂ O	0.9	0.9
Na ₂ CO ₃ .10H ₂ O	0.6	2.8
Na ₂ CrO ₄ .10H ₂ O	1.6	0
NaCl	4.0	0.2
Na ₂ B ₄ O ₇ .10H ₂ O	4.0	0.9
Nal	1.0	1.7
NaHPO ₄ .12H ₂ O	0.4	1.5
NaNO ₃	0.9	0.7
NaNO2	0.9	0.6
Na ₂ SO ₄ .10H ₂ O	0.3	0.7
Na ₂ S ₂ O ₃ .5H ₂ O	1.0	2.2
Urea	2.0	

^aWorking values usually are not more than one-half the maxima. (After Mullin, 1972).

16.3. THE PROCESS OF CRYSTALLIZATION

The questions of interest are how to precipitate the crystals and how to make them grow to suitable sizes and size distributions. Required sizes and size distributions are established by the need for subsequent recovery in pure form and ease of handling, and by traditional commercial practices or consumer preferences.

CONDITIONS OF PRECIPITATION

The most common methods of precipitating a solid from a solution are by evaporation of the solvent or by changing to a temperature at which the solubility is lower. Usually solubility is decreased by lowering the temperature. Some examples are in Figure 16.1. The limit of removal is determined by the eutectic composition. According to the data of Figure 16.2, for instance, a 24.6% solution of KC1 will solidify completely at -11°C and a 3.5% solution of **MgSO**₄ will do so at 4°C; these values represent the limits to which salt is recoverable by chilling. Complete recovery, however, is accomplished by evaporation.

A precipitate may be formed as a result of chemical reaction between separately soluble gases or liquids. Commercial examples are productions of sodium sulfate, ammonium sulfate, and ammonium phosphate.

Precipitation also can be induced by additives, a process generally called salting out because salts with ions common to those whose precipitation is desired are often used for this purpose. For instance, ammonium chloride is recovered from spent Solvay liquors by addition of sodium chloride and the solubility of $BaCl_2$ can be reduced from 32% to 0.1% by addition of 32% of $CaCl_2$. Other kinds of **precipitants** also are used, for instance, alcohol to precipitate aluminum sulfate from aqueous solutions.

Foreign substances even in minute amounts may have other kinds of effects on crystallization: They may inhibit or accelerate growth rate or change the shape of crystals, say from rounded to needlelike, or otherwise. One of the problems sometimes encountered with translating laboratory experience to full scale operation is that the synthetic liquors used in the laboratory may not contain the actually occurring impurities, and thus give quite different performance. Substances that modify crystal formation are very important industrially and many such materials have been the subject of patents.

SUPERSATURATION

A saturated solution is one that is in equilibrium with the solid phase and will remain unchanged indefinitely at a particular

EXAMPLE 16.2

Using the Phase Diagrams of Figure 16.2

- a. Evaporation of a solution of MgSO₄ at 30°C: As water is removed, the composition moves along the horizontal. When the salt concentration reaches about 6%, precipitation of heptahydrate begins and is completed at about 13%. Between 13 and 14% salt, the precipitate is a mixture of solid hepta and solid hexa hydrates. Beyond 14%, the mixture consists of mixtures of solid hepta and mono hydrates in proportions determined by the amount of water present overall.
- b. Chilling of a 6% solution of MgSO₄: precipitation of heptahydrate begins at about 35°C. At about 2°C, the mixture consists of solid dodecahydrate and unsaturated solution. Below -4°C complete solidification exists; the product is a mixture of pure dodecahydrate and an intimate eutectic mixture of ice and dodeca crystals.
- c. Recovery of pyridine: As appears on the diagram of Figure

16.2(e), the eutectic contains 33% formamide and 67% pyridine. When the mixture contains 80% pyridine, the maximum possible recovery of pure pyridine is

$$P = (0.8 - 0.67)/(1 - 0.67) = 0.39$$
, or 39%.

d. Recovery of formamide: When the mixture of Figure 16.2(e) contains 80% formamide, the maximum recovery of the pure material is

$$F = (0.8 - 0.33)/(1 - 0.33) = 0.70$$
, or 70%.

- e. At 50°C, the liquid phase of Figure 16.2(d) contains 35% and the solid phase 74% of β -chloronaphthalene.
- **f.** The progress of crystallization of a ternary mixture such as that of Figure 16.2(f) is described in Example 16.8.

EXAMPLE 16.3

Heat Effect Accompanying the Cooling of a Solution of $MgSO_4$ A 30% solution of $MgSO_4$ is cooled from 150F to 50F. Data of the initial and final conditions are taken off the equilibrium diagram, Fig. 16.3(b). At the lower temperature, 27% of the mixture crystallizes out as the heptahydrate.

	Onininal	Final (at 50°F)			
	at 150°F	Total	Liquid	Solid	
Water (Ib)	38.6	38.6	38.6		
MgSO ₄ ·7H ₂ O (lb)	61.4	61.4	34.4	27.0	
Total (Ib)	100	100	73.0	27.0	
H (Btu/lb)	-3	-82	-53	-161	
Accordingly the change	in enthalpy	is			

AH = -82 - (-3) = -79 Btu/lb.

This value will be compared with a calculation using data of heat capacities and heat of solution. From *Perry's Chemical Engineers' Handbook* (1984), the heat solution of the heptahydrate is -39.2 Btu/lb and its heat capacity is $0.36 \text{ Btu/(lb)}(^{\circ}\text{F})$. The enthalpy change of the cooling and crystallization process is

$$AH = [0.386 + 0.614(0.36)](50 - 150) + 0.27(-39.2)$$

= -71.3 Btu/lb,

which is a poor check of the value found with the aid of the equilibrium diagram. Possible sources of error of the second method include the use of heat of solution at infinite dilution instead of the prevailing concentration and the assumption that the heat capacities are additive.



Figure 16.3. Enthalpy-composition diagrams of some salt solutions. Several other diagrams are in the compilation of Landolt-Bernstein, IV 4b, 1972, pp. 188-224. (a) sodium sulfate/water; (b) magnesium sulfate/water (after Chemical Engineers' Handbook, 1963 edition, McGraw-Hill, New York); (c) sodium carbonate/water.



Figure 16.3—(continued)

temperature and composition of other constituents. Greater than normal concentrations also can be maintained in what is called a supersaturated condition which is metastable. Metastability is sensitive to mechanical disturbances such as agitation, ultrasonics, and friction and the introduction of solid particles. Under those

B.S. mesh number	Sieve uperture. µm	Fractional weight per cent retained	Cumulative weight per cent oversize	Cumulative weight per cent undersize
7	2360	1.2	1.2	98.8
10	1700	2.9	4.1	95.9
14	1180	18.8	22.9	77.1
18	850	28.8	51.7	48·3
25	600	22.0	73.7	26.3
36	425	11-1	84.8	15.2
52	300	60	90.8	9.2
12	212	3.9	94.7	5.3
100	150	1.8	96.5	3.5
150	106	1.3	97.8	2.2
> 150		2.2		

(a)

conditions, solids will separate out until normal saturation is obtained. When great care is taken, the metastable state is reproducible. A thermodynamic interpretation of metastability can be made in terms of the Gibbs energy of mixtures. In Figure 16.5(a), the solid line u-b is of unsaturated solution and the straight line b-e is of mixtures of all proportions of pure solid and saturated solution represented by point b. Points c and d are at the points of inflection of the plot and represent the limits of metastability. Thus line b-c represents the range of concentrations between the saturated and supersaturated values.

Several measures of supersaturation are being used in terms of the saturation concentration C_0 ; thus

$$\begin{split} \alpha &= \Delta C_s = C - C_0, & \text{the difference in concentrations,} \\ \beta &= \Delta C_s / C, & \text{the relative difference,} \\ \gamma &= C / C_0 = \beta + 1, & \text{the concentration ratio,} \end{split}$$

with similar definitions for subcooling or superheating. The data of Figure 16.1(c) and Table 16.1 show that excess concentration and metastable cooling can be quite substantial amounts.

GROWTH RATES

Crystallization can occur only from supersaturated solutions. Growth occurs first by formation of nuclei and then by their gradual growth. At concentrations above supersaturation, as at point d' on Figure 16.1(d), nucleation is conceived to be spontaneous and rapid. In the metastable region, nucleation is caused by mechanical shock or friction and secondary nucleation can result from the breakup of already formed crystals. It has been observed that the rate depends on the extent of supersaturation; thus

$$\frac{dC_N}{d\theta} = k(C - C_0)^m. \tag{16.3}$$

Values of the exponent m have been found to range from 2 to 9, but have not been correlated to be of quantitative value for prediction.



Figure 16.4. Several ways of recording the same data of crystal size distribution (CSD) (Mullin, 1972). (a) The data. (b) Cumulative wt % retained or passed, against sieve aperture. (c) Log-log plot according to the RRS equation P = exp[(-d/d),"]; off this plot, $d_{50\%} = 850$, $d_m = 1000$, n = 1.8. (d) Differential polygon. (e) Differential histogram.



Nucleation rates are measured by counting the numbers of crystals formed over periods of time.

The growth rates of crystals depend on their instantaneous surface and the linear velocity of solution past the surface as well as the extent of supersaturation, and are thus represented by the equation

$$\frac{dW_c}{d\theta} = kuA(C-C_0)^n. \tag{16.3'}$$

Values of the exponent have been found of the order of 1.5, but



Figure 16.5. Supersaturation behavior. (a) Schematic plot of the Gibbs energy of a solid solute and solvent mixture at a fixed temperature. The true equilibrium compositions are given by points b and e, the limits of metastability by the inflection points C and d. For a salt-water system, point d virtually coincides with the 100% salt point e, with water contents of the order of 10^{-6} mol fraction with common salts. (b) Effects of supersaturation and temperature on the linear growth rate of sucrose crystals [data of Smythe (1967) analyzed by Ohara and Reid, 1973].



Figure 16.6. Crystal size distributions of several materials in several kinds of crystallizers (Bamforth, 1965).

Cod	e Crystallizer	Substance	d _m	n
а	Escher-Wyss	NaCl	0.7	4.7
b	Giavanola	adipic acid	0.4	8.1
с	Matusevich	NaNO ₃	0.37	4.0
d	Kestner	Na₂SO₄	0.92	4.7
е	Oslo-Krystal	(NH ₄) ₂ SO ₄	3.2	2.1
f	Oslo-Krystal	$(NH_{4})_{2}SO_{4}$	2.35	6.0
g	Sergeev	$(NH_4)_2 SO_4$	-	1.5(?)
ĥ	DTB	(NH₄)₂SO₄	1.6	5.7
i	Standard saturator	(NH ₄) ₂ SO ₄	0.62	2.6

The parameters are those of the RRS equation, Eq. 16.1.

again no correlation of direct use to the design of crystallizers has been achieved. The sucrose growth data of Figure 16.5(b) are not quite log-log linear as predicted by this equation.

In laboratory and commercial crystallizations, wide size distributions usually are the rule, because nuclei continue to form throughout the process, either spontaneously or by breakage of already formed crystals. Large crystals of more or less uniform size are desirable. This condition is favored by operating at relatively low extents of supersaturation at which the nucleation rate is low but the crystals already started can continue to grow. The optimum extent of supersaturation is strictly a matter for direct experimentation in each case. As a rough guide, the data for allowable subcooling and corresponding supersaturation of Table 16.2 may serve. Since the recommended values are one-half the maxima shown, it appears that most crystallizations under

commercial conditions should operate with less than about 2° C subcooling or the corresponding supersaturation. The urea plant design of Example 16.4 is based on 2° F heating.

Growth rates of crystals also must be measured in the laboratory or pilot plant, although the suitable condition may be expressed simply as a residence time. Table 16.3 gives a few growth rate data at several temperatures and several extents of super-saturation for each substance. In most instances the recommended supersaturation measured as the ratio of operating to saturation concentrations is less than 1.1. It may be noted that at a typical rate of increase of diameter of 10^{-7} m/sec, the units used in this table, the time required for an increase of 1 mm is 2.8 hr.

Batch crystallizers often are seeded with small crystals of a known range of sizes. The resulting crystal size distribution for a given overall weight gain can be estimated by an approximate

EXAMPLE 16.4 Deductions from a Differential Distribution Obtained at a Known Residence **Time**

The peak of the differential distribution obtained with a residence time of i = 2 hr corresponds to $L_{pr} = 1.2$ mm. Assuming ideal mixing, $L_{pr}/G\bar{t} = 1.2/2G = 3$, and G = 0.2 mm/h. With this knowledge of G, crystal size distributions could be found at other residence times.

relation known as the McCabe Delta-L Law, which states that each original crystal grows by the same amount AL. The relation between the relative masses of the original and final size distributions is given in terms of the incremental AL by

$$R = \frac{\sum w_i (L_{0i} + \Delta L)^3}{\sum w_i L_{0i}^3}$$
(16.4)

When R is specified, AL is found by trial, and then the size distribution is evaluated. Example 16.5 does this.

Some common substances for which crystallization data are reported in the literature and in patents are listed in Table 16.4.

16.4. THE IDEAL STIRRED TANK

All continuous crystallizers are operated with some degree of mixing, supplied by internal agitators or by pumparound. The important limiting case is that of ideal mixing in which conditions are uniform throughout the vessel and the composition of the effluent is the same as that of the vessel content. In crystallization literature, this model carries the awkward name MSMPR (mixed suspension mixed product removal). By analogy with the terminology of chemical reactors it could be called CSTC (continuous stirred tank crystallizer). Several such tanks in series would be called a CSTC battery. A large number of tanks in series would approach plug flow, but the crystal size distribution still would not be uniform if nucleation continued along the length of the crystallizer.

The process to be analyzed is represented by Figure 16.4. What will be found are equations for the cumulative and differential size distributions in terms of residence time and growth rate. The principal notation is summarized here.

Q = volumetric feed rate,

 V_c = volume of holdup in the tank

- n = number of crystals per unit volume
- L =length of the crystal
- G = linear growth rate of the crystal
- t = time

 $\bar{t} = V_c/Q$, mean residence time

- $x = L/G\overline{t}$, reduced time
- ϕ_m = cumulative mass distribution
- n^0 = zero side nuclei concentration, also called zero size population density
- $B^0 =$ nucleation rate
- a, = volume shape factor = volume of crystal/(length)³ = n/6 for spheres, = 1 for cubes.

The case being considered is that in which the feed contains no

 TABLE 16.3. Mean Overall Growth Rates of Crystals (m/sec) at Each Face^a

Crystallising substance	°C	S	v (m/s)
$(\mathrm{NH}_4)_2\mathrm{SO}_4 \cdot \mathrm{Al}_2(\mathrm{SO}_4)_3 24\mathrm{H}_2\mathrm{O}$	15	1.03	1.1 × 10 ⁻⁸ *
	30	1.03	1.3 x 10 ⁻⁰
	40	1.08	1.0×10^{-7}
NH ₄ NO ₃	40	1.05	8.5 x 10 ⁻⁷
(NH ₄),SO ₄	30	1.05	2.5 x 10 ^{−7} •
	60	1.05	4.0×10^{-7}
	90	1.01	3.0×10^{-8}
NH ₄ H ₂ PO ₄	20	I-06	6.5 × 10 ⁻⁸
	30	1.02	3.0×10^{-8}
	30 40	1.02	1.1×10^{-8}
	40	1.02	/0 10-8+
MgSO ₄ ·7H ₂ O	20	1.02	4.5 x 10 °*
	30	1.02	1.5×10^{-7}
NISO (NH) SO (H O	25	1.02	5 2 x 10 ⁻⁹
$1030_4 \cdot (101_4)_2 \cdot 30_4 \cdot 01_2 \cdot 00_2 \cdot $	25	1.05	3.2×10^{-8}
	25	i ∙20	4.0×10^{-8}
K_3SO_4 Al ₂ $(SO_4)_3 \cdot 24H_3O_4$	15	1.04	1.4×10^{-8}
	30	1.04	2.8 x 10 ⁻⁸ *
	30	1.09	1·4 x 10 ⁻⁷ *
	40	1.03	5·6 x 10 ⁻⁸ *
KCI	20	1.02	2.0×10^{-7}
	40	1.01	6.0×10^{-7}
KNO ₃	20	1.05	4.5×10^{-8}
K 50	40	1.00	1.5 × 10
R2504	20	1.18	2.8 X 10
	30	1.10	4.2 x 10 ⁻⁸ *
	50	1.06	7.0 x 10 ⁻⁸ *
	50	1.12	$3.2 \times 10^{-7*}$
KH ₂ PO ₄	30	1.07	3.0×10^{-8}
	30	1.21	2.9×10^{-7}
	40	I+06 1+18	5.0×10^{-9}
NaCl	40	1.002	4.0 × 10
	50 50	1.002	2'5 x 10 ° 6:5 x 10 ⁻⁸
	70	1.003	9.0×10^{-8}
	70	1.003	1.5 x 10 ⁻⁷
$Na_2S_2O_3 \cdot 5H_2O$	30	1.02	1·1 x 10 ⁻⁷
	30	1.08	5.0×10^{-7}
Citric acid monohydrate	25	1.05	3·0 x 10 ^{−8}
	30	1.01	1.0×10^{-8}
	30	1.05	4·0 x 10 -8
ucrose	30	1.13	1·1 x 10 ⁻⁸ *
	30	1.27	$2.1 \times 10^{-8*}$
	70 70	1.15	9.3 x 10 ° 1.5 x 10 ⁻⁷
	70	1.12	1.5 X 10

^a The supersaturation is expressed by S = C/C,,, with Cthe amount dissolved and C_0 the normal solubility (kg crystals/kg water). The mean growth velocity is that at one face of the crystal; the length increase is $G = 2\bar{v}$ (m/sec). Data are for crystals in the size range 0.5-1.0 mm in the presence of other crystals. The asterisk denotes that the growth rate probably is size-dependent.

(Mullin, 1972).

EXAMPLE 16.5 Batch Crystallization with Seeded Liquor Seed crystals with this size distribution are charged to a batch crystallizer: L ₀ , length (mm) 0.251 0.178 0.127 0.089 0.064 w (wt fraction) 0.09 0.26 0.45 0.16 0.04	140 PRINT "INCREMENT L=";L 150 PRINT "SUMMATION=";S 160 PRINT "WEIGHT RATIO=";S/.003 93455 170 END
 On the basis of the McCabe AL law. these results will be found: a. The length increment that will result in a 20-fold increase in mass of the crystals. b. The mass growth corresponding to the maximum crystal length of 1.0 mm. When <i>L</i> is the increment in crystal length, the mass ratio is R = ∑ w_i(L_{0i} + L)³ - ∑ w_i(L_{0i} + L)³ = 20	W L L +L .090 .251 .2510 .260 .178 .1780 .450 .127 .1270 .160 .089 .0890 .040 .064 .0640 INCREMENT L 0 SUMMATION= .00393458 126 WE I GHT RATIO = 1 .00000032024
a. By trial, the value of $L = 0.2804$ mm. b. When $L = 1 - 0.251 = 0.749$, $R = 181.79$. The size distributions and the computer program are tabulated. 10 ! Example 16.4. Batch crysta 11:zation with seeded liquor 20 OPTION BASE 1 25 DIM L0(5),W(5),S(5) 30 MAT READ L0,W 40 DATA 221, 178, 127, 089, 06	.090 .251 .5314 .260 .178 4584 .450 .127 4074 .160 .089 3694 .040 .064 .3444 INCREMENTL = .2804 SUMMATION= 7 167685 11336E-2 WEIGHT RATIO= 19 9962514763
409,.26,.45,.16,.04 5.0 INPUT L 60 S=0 70 FOR I=1 TO 5 80 S(I)=(L0(I)+L)^3*W(I) 100 S=S(I)+S 110 PRINT USING 120 , W(I),L0(I) ,L0(I)+L 120 IMAGE .DDD,2X,.DDD,2X,D.DDDD 130 NEXT I 135 PRINT	090 .251 1.0000 .260 178 .9270 .450 127 .8760 .160 .089 8380 .040 .064 8130 INCREMENT L= .749 SUMMATION= .71526668218 WEIGHT RATIO= 181 789843434

nuclei but they are generated in the tank. The balance on the number of crystals is

rate of generation = rate of efflux

or

$$V_c \frac{dn}{dt} = Qn. \tag{16.5}$$

Upon substituting for the linear growth rate

 $G = dL/dt \tag{16.6}$

and rearranging,

$$\frac{dn}{n} = \frac{Q}{V_c} dt = \frac{Q}{V_c G} dL = \frac{dL}{tG} = dx$$
(16.7)

where

$$\bar{t} = V_c/Q \tag{16.8}$$

is the mean residence time and

$$x = L/Gt = t/i \tag{16.9}$$

is the dimensionless time. Integration of the equation

$$\int_0^n \frac{dn}{n} = \frac{1}{Gi} \int_0^L dL \tag{16.10}$$

is

$$n = n^0 \exp(-L/G\overline{t}) = n^0 \exp(-x), \qquad (16.11)$$

TABLE 16.4. Some Common Substances for which Crystallization Data Are Reported in the Literature and in **Patents**^a

Compound	Remark or aspect referred 10	Compound	Remark or aspect referred to
Ag-halides Ag₂Cr O₄ Al F₃	growth kinetics	SrSO₄ ZnSO₄ anthracene	
Al $_2O_3$ -corundum Al NH $_4$ (SO $_4$) $_2$ Al K (SO $_4$) $_2$ Al (OH) $_3$ H $_3BO_3$ Na $_2B_4O_7$ BaSO $_4$	influence of supersaturation oleic acid conducive nucleation growth	adipic acid sugars citric acid phenols xylenes naphthalene paraffin	methods and parameters of
BaCO₃ BaTiO₄	habit	urea	crystallisation NH ₄ Cl, MgCO ₃ glyoxal, cyanuric acid
$CaSO_4$ $CaCO_3$ $CaCl_2$ $Ca(NO_3)_2$	citiates, SO ² , elevated temp metaphosphate conducive	Na-acetate NaK-tartarate pentaerythrite	surface-active agents
$K_2Cr_2O_7$ $CuSO_4$ $CuCl_2$ $FeSO_4$	excess H_2SO_4 detrimental	pepsine terephthalic acid	
H₂O NH₄J	nucleation growth nucleation	'(The references, som Appendix A).	e 400 in number, are given by Nyvlt, 1971
K-halides KH ₂ PO ₄ KNO ₃	Pb^{2+}, Zn^{2+} conducive	where	
K_2SO_4 K_2CrO_4 $MgSO_4$ $MgCl_2$	$t = 45^{\circ}$ C. borax conducive	$n^0 = \lim_{L \to 0} \frac{dn}{dL}$	(16.12)
MnCl ₂ LiF LiCl		is the concentration of c also is called the zero The nucleation rate	crystals of zero length which are the nuclei; it size population density.
Li₂SO₄ NaCl	Pb, Fe, Al, Zn conducive; caking inhibited by ferrocyanides ; urea leads	$B^0 = \lim_{L \to 0} \frac{dn}{dt} = \lim_{L \to 0}$	$\left(\frac{dL}{dt}\frac{dn}{dL}\right) \tag{16.13}$
Na₂CO₃ NaHCO₃ Na₂SO₄	Na_2SO_4 conducive wetting agents conducive	$= Gn^{0}.$	(16.14)
Na ₂ S ₂ O ₃ NaClO ₃			
NaCN NH₄NO₃	paraffin, urea, dyes methods of crystallising effect of additives: conducive	n= 0	
(NH₄)₂ SO₄	urea, Fe^2 +, Mg^2 +, $tannin$, $pH5$; Al^3 + and Fe^2 + lead to needle formation removal of admixtures crystal growth methods of crystallising		N N N N N N N N N N N N N N N N N N N
(NH ₄) ₂ S ₂ O ₅ NH ₄ HCO ₃ NH ₄ Cl	coarse grained , stabilisation Zn⁺⁺ , Pb⁺⁺ , NH₄⁺ , wood extract		
H_3PO_4 $NH_4H_2PO_4$ $(NH_4)_2HPO_4$ NiSO	Fe^{3+} and NH_{4}^{+} conducive		
$Pb(NO_3)_2$ $PbCO_3$		Figure 16.7. Materia crystallizers (CSTC). (battery with overall res	a) balancing of continuous stirred tank (a) The single stage CSTC. (b) Multistage idence time $\tilde{t} = (l/Q) \sum_{i=1}^{k} V_{ci}$.

536 CRYSTALLIZATION FROM SOLUTIONS AND MELTS

The number of crystals per unit volume is

$$n_{c} = \int_{0}^{\infty} n \, dL = \int_{0}^{\infty} n^{0} \exp(-L/G\bar{t}) \, dL = n^{0}G\bar{t}.$$
 (16.15)

The total mass of crystals per unit volume is

$$m_{c} = \int_{0}^{\infty} mn \, dL = \int_{0}^{\infty} a_{v} \rho_{c} L^{3} n^{0} \exp(-L/G\bar{t}) \, dL$$

= $6a_{v} \rho_{c} n^{0} (G\bar{t})^{4}$, (16.16)

where a_v is the volumetric shape factor and ρ_c the crystal density. Accordingly, the number of crystals per unit mass is

$$n_c/m_c = 1/6a_v \rho_c (G\bar{t})^3.$$
 (16.17)

The mass of crystals per unit volume with length less than L or with dimensionless residence time less than \boldsymbol{x} is

$$m_L = \frac{L}{I_0} mn \ dL = a_v \rho_c (G\bar{t})^4 n_{I_0}^0 x^3 e^{-x} \ dx.$$
(16.18)

The value of the integral is

$$\int_0^x x^3 e^{-x} dx = 6[1 - e^{-x}(1 + x + x^2/2 + x^3/6)].$$
(16.19)

This expression has a maximum value at $\chi = 3$ and the corresponding length L_{pr} is called the predominant length

$$L_{\rm pr} = 3Gi.$$
 (16.20)

The cumulative mass distribution is

$$\phi_m = m_L/m_c = 1 \quad e^{-x}(1 + x + x^2/2 + x^3/6),$$
 (16.21)

and the differential mass distribution is

$$d\phi_m/dx = x^3 e^{-x}/6,$$
 (16.22)

which has a maximum value of 0.224 at $\boldsymbol{x} = 3$.

The nucleation rate must generate one nucleus for every crystal present in the product. In terms of M', the total mass rate of production of crystals,

$$B^{0} = \frac{M'}{n_{c}/m_{c}} = \frac{M'}{6a_{v}\rho_{c}(G\bar{t})^{3}} = \frac{1.5M'}{a_{v}\rho_{c}L_{\rm pr}^{3}}.$$
 (16.23)

The principal quantities related by these equations are ϕ_m , $d\phi_m/dx$, L, L_{pr} , t, n^0 , and B^0 . Fixing a certain number of these will fix the remaining one. Size distribution data from a CSTC are analyzed in Example 16.6. In Example 16.7, the values of the predominant length L_{pr} and the linear growth rate G are fixed. From these values, the residence time and the cumulative and differential mass distributions are found. The effect of some variation in residence time also is found. The values of n^0 and B^0 were found, but they are ends in themselves. Another kind of condition is analyzed in Example 16.4.

MULTIPLE STIRRED TANKS IN SERIES

Operation in several tanks in series will provide narrower size distributions. Equations were developed by Nyvlt (1971) for two main cases. With generation of nuclei in the first stage only, the

cumulative and differential distributions for k stages are

$$\phi_m = 1 - e^{-kx} \sum_{j=0}^{k+2} \frac{n^j x^j}{j!}, \qquad (16.24)$$

$$\frac{d\phi_m}{dx} = \frac{k(kx)^{k+2}}{(k+2)!} e^{-kx}.$$
(16.25)

The multistage distributions are plotted in Figure 16.8 for several values of the number of stages. Maxima of the differential distributions occur at

$$x_{\max} = 1 + 2/k,$$
 (16.26)

and the values of those maxima are represented by

$$\left(\frac{d\phi_m}{dx}\right)_{\max} = \frac{k^{k+3}(1+2/k)^{k+2}}{(k+2)!} \exp[-(k+2)].$$
 (16.27)

Some numerical values are:

k l 2 3 4 5 10

$$X_{max}$$
 3 2 1.67 1.5 1.4 l.2
 $(d\phi_m/dx)_{max}$ 0.224 0.391 0.526 0.643 0.745 1.144

Nyvlt (1971) also develops equations for multistage crystallizers in which nuclei form at the same rate in all stages. For two such stages, the cumulative distribution is represented by

$$\phi_m = 1 - 0.5e^{-x}[1 + x + x^2/2 + x^3/6] - 0.5e^{-2x}[1 + 2x + 2x^2 + (4/3)x^3 + (2/3)x^4].$$
(16.28)

A comparison of two-stage crystallizers with nucleation in the first stage only and with nucleation in both stages appears in Figure 16.9. The uniformity of crystal size is not as good with nucleation proceeding in every stage; the difference is especially pronounced at larger numbers of stages, which are not shown here but are by Nyvlt (1971).

As in the operation of chemical reactors, multistaging requires shorter residence time for the same performance. For the same L/G ratio, the relative crystallization times of k stages and one stage to reach the peaks are given by Eq. (16.26) as

$$t_k/t_1 = (1 + k/2)/3,$$
 (16.29)

which is numerically 0.4 for five stages. Not only is the time shortened, but the size distribution is narrowed. What remains is how to maintain substantial nucleation in only the first stage. This could be done by seeding the first stage and then operating at such low supersaturation that spontaneous nucleation is effectively retarded throughout the battery. Temperature control also may be feasible.

APPLICABILITY OF THE CSTC MODEL

Complete mixing, of course, is not practically realizable and in any event may have a drawback in that intense agitation **will** cause much secondary nucleation. Some rules for design of agitation of solid suspensions are discussed in Chapter 10, notably in Table 10.2; internal velocities as high as 1.0 **ft/sec** may be desirable.

Equations can be formulated for many complex patterns, combinations of mixed and plug flow, with decanting of supernatant liquor that contains the smaller crystals and so on. A modification to the CSTC model by Jancic and Garside (1976) recognizes that linear crystal growth rate may be size-dependent; in one instance

EXAMPLE 16.6

Analysis of Size Distribution Data Obtained in a CSTC Differential distribution data obtained from a continuous stirred tank crystallizer are tabulated.

W	L	∑w/L³
0.02	0.340	0. 5089
0.05	0.430	1. 1377
0.06	0.490	1.6477
0.08	0.580	2.0577
0. 10	0.700	2.3493
0.13	0.820	2. 5851
0.13	1.010	2.7112
0.13	1.160	2.7945
0.10	1.400	2.8310
0.09	1.650	2.8510
0.04	1.980	2.8562
0.03	2.370	2.8584

The last column is of the summation $\sum_{0}^{L} w_i/L_i^3$ at corresponding values of crystal length L. The volumetric shape factor is $a_v = 0.866$, the density is 1.5 g/mL, and the mean residence time was 2.0 hr. The linear growth rate G and the nucleation rate B^0 will be found.

The number of crystals per unit mass smaller than size L is

$$N = \frac{1}{a_{v}\rho} \sum_{0}^{L} \frac{w_{i}}{L_{i}^{3}}.$$
 (1)

It is also related to the CSTC material balance by

$$dN/dL = n = n^0 \exp(-L/Gt).$$
⁽²⁾

Integration of Eq. (2) is

$$N = \int_{\theta}^{L} n^{0} \exp(-L/G\overline{t}) dL = G\overline{t}n^{0}[1 - \exp(-L/G\overline{t})].$$
(3)

Combining Eqs. (1) and (3),

$$\sum w_i/L_i^3 = a_v \rho \bar{Gin^0} [1 - \exp(-L/\bar{Gi})].$$
⁽⁴⁾

The two unknowns G and n^0 may be found by nonlinear regression with the 12 available data for L_i . However, two representative

they find that

$$G = G^0 (1 + L/G^0 \bar{t})^{0.65}.$$

Other studies have tried to relate sizes of draft tubes, locations and sizes of baffles, circulation rate, and so on to crystallization behavior. So far the conclusions are not general enough to do a designer much good. A possibly useful concept, the separation index (SI), is mentioned by **Mullin** (1976, **p**. 293):

SI = (kg of 1 mm equivalent crystals)/m³ hr.

For inorganic salts in water at near ambient temperature, a value of **SI** in the range of $100-150 \text{ kg/m}^3/\text{hr}$ may be expected. An illustration of the utilization of pilot plant data and plant experience in the design of a urea crystallizer is in Example 16.1.

In general, the design policy to be followed is to utilize as much

values of L_i are taken here, and the unknowns are solved for by simultaneous solution of two equations. When

$$L = 0.58, \quad \sum = 2.0577,$$

 $L = 1.40, \quad \sum = 2.8310.$

Substituting into Eq. (4) and ratioing,

$$\frac{2.8310}{2.0577} = \frac{1 - \exp(-1.4/Gt)}{1 - \exp(-0.58/Gt)};$$

by trial,

$$G\bar{t} = 0.5082$$

 $G = 0.508212 = 0.2541.$

With L = 1.4 in Eq. (4),

$$2.8310 = 0.866(1.5)(0.5082)n^{0}[1 - \exp(-1.4/0.5082)].$$

from which

$$n^0 = 4.58$$
 nuclei/mm⁴
= 4.58(10)¹² nuclei/m⁴.

Accordingly,

$$B^0 = Gn^0 = 0.2541(10)^{-3}(4.58)(10)^{12} = 1.16(10)^9$$
 nuclei/m³ hr.

The cumulative mass size distribution is represented by

$$\phi_m = 1 - e^{-x}(1 + x + x^2/2 + x^3/6)$$

with

$$x = L/Gt = L/0.5082.$$

This distribution should be equivalent to the original one, but may not check closely because the two points selected may not have been entirely representative. Moreover, although the data were purportedly obtained in a CSTC, the mixing may not have been close to ideal.

laboratory and pilot plant information as possible, to work it into whatever theoretical pattern is applicable, and to finish off with a comfortable safety factor. There may be people who know how; they should be consulted.

16.6. KINDS OF CRYSTALLIZERS

The main kinds of crystallizers are represented in Figure 16.10. They will be commented on in order. Purification of products of melt crystallization is treated separately.

Batch crystallizers are used primarily for production of fine chemicals and pharmaceuticals at the rate of 1-100 tons/week. The one exception is the sugar industry that still employs batch vacuum crystallization on a very large scale. In that industry, the syrup is concentrated in triple- or quadruple-effect evaporators, and crystallization is completed in batch vacuum pans that may or may not be equipped with stirrers [Fig. 16.11(g)].

EXAMPLE 16.7

Crystallization in **a** Continuous Stirred Tank with Specified Predominant Crystal Size

Crystals of citric acid monohydrate are to made in a CSTC at 30°C with predominant size $L_{pr} = 0.833$ mm (20 mesh). The density is 1.54 g/mL, the shape factor a, = 1 and the solubility is 39.0 wt %. A supersaturation ratio C/C, = 1.05 is to be used.

Take the growth rate, $G = 2\bar{v}$, to be one-half of the value given in Table 16.3:

 $G = dL/d\theta = 4(10^{-8}) \text{ m/sec}, 0.144 \text{ mm/hr}.$

The predominant size is related to other quantities by

 $L_{\rm pr} = 0.833 = 3G\bar{t},$

from which

$$t = 0.833/(3)(0.144) = 1.93$$
 hr.

For a mass production rate of 15 kg/hr of crystals, C = 15, the nucleation rate is

$$B^{0} = \frac{1.5C}{a_{\nu}\rho_{c}L_{\rm pr}^{3}} - \frac{1.5(15)}{1(1.5)[0.833(E-3)]^{3}}$$

= 2.595(10)¹⁰ nuclei/m³ hr.

The zero size concentration of nuclei is

$$n^{0} = B^{0}/G = 2.595(10)^{10}/4(10)^{-8} = 6.49(10)^{17}$$
 nuclei/m⁴.

Accordingly, the equation of the population density is

$$n = n^0 \exp(-L/G\bar{t}) = \exp(41.01 - 360L).$$

The cumulative mass distribution is

$$\phi_m = 1 - e^{-x}(1 + x + x^2/2 + x^3/6),$$

where

$$x = L/G\bar{t} = 3.60L$$
, with L in mm.

Natural circulation evaporators like those shown on Figure 8.16 may be equipped for continuous salt removal and thus adapted to crystallization service. For large production rates, however, forced circulation types such as the DTB crystallizer of Figure 16.10(g), with some control of crystal size, are the most often used. The lower limit for economic continuous operation is 1-4 tons/day of crystals, and the upper limit in a single vessel is 100-300 tons/day, but units in parallel can be used for unlimited capacity.

Many special types of equipment have been developed for particular industries, possibly extreme examples being the simple open ponds for solar evaporation of brines and recovery of salt, and the specialized vacuum pans of the sugar industry that operate with syrup on the **tubeside** of calandrias and elaborate! internals to eliminate entrainment. Some modifications of basic types **of** crystallizers often carry the inventor's or manufacturer's name. For their identification, the book of Bamforth (1965) may be consulted.

The basic equipment descriptions following carry the letter designations of Figure 16.10.

The differential distributions are differences between values of ϕ_m at successive values of crystal length *L*. The tabulation shows cumulative and differential distributions at the key i=1.93 hr, and also at 1.5 and 3.0 hr. The differential distributions are plotted and show the shift to larger sizes as residence time is increased, but the heights of the peaks are little affected.



(a) Jacketed pipe scraped crystallizers. These are made with inner pipe 6-12 in. dia and 20-40 ft long, often arranged in tiers of three or more connected in series. Scraper blades rotate at 15-30 rpm. Temperatures of -75 to $+100^{\circ}$ F have been used and viscosities in excess of 10,000 cP present no problems. Although the action is plug flow with tendency to uniform crystal size, the larger particles settle to the bottom and grow at the expense of the smaller ones that remain suspended, with the result that a wide range of sizes is made. Capacity is limited by rates of heat transfer; coefficients of 10-25 Btu/(hr)(sqft)(°F) usually are attainable. Higher coefficients are obtainable in Votators (Cherry Burrell Co.) that have more intense scraping action. Pilot units of 4in. by 4ft and larger are made.

(b) Swenson-Walker type. In comparison with jacketed pipes, they have the advantage of being more accessible for cleaning. The standard unit is 24 in. wide, 26 in. high, and 10 ft long. Four units in line may be driven off one shaft. Capacity is limited by heat transfer rates which may be in the range of 10-25 Btu/(hr)(sqft)(°F), with an



Figure 16.8. Theoretical crystal size distributions from an ideal stirred tank and from a series of tanks with generation of nuclei only in the first tank. Equations of the curves and for the peak values are in the text. (a) Cumulative distributions. (b) Differential distributions.

effective area of 3 sqft/ft of length. According to data in *Chemical* Engineers' *Handbook* (3rd ed., McGraw-Hill, New York, 1950, p. 1071), a 40 ft unit is able to produce 15 tons/day of trisodium phosphate, and a 50ft unit can make 8 tons/day of Glaubers salt. The remarks about crystal size distribution made under item (a) apply here also.

(c) Batch stirred and cooled types. Without agitation, crystallization time can be 2-4 days; an example is given in *Chemical Engineer's Handbook* (1950, **p**. 1062). With agitation, times of 2-8 hr are sometimes cited. The limitation is due to attainable rates of heat transfer. Without encrustation of surfaces by crystals, coefficients of 50-200 **Btu/(hr)(sqft)(°F)** are realizable, but temperature differences are maintained as low as $5-10^{\circ}$ F in order to keep supersaturation at a level that prevents overnucleation. Stirring breaks corners off crystals and results in secondary nucleation so that crystal size is smaller than in unagitated tanks. Larger crystal sizes are obtained by the standard practice of seeding with an appropriate range of fine crystals. Calculation of the performance of such an operation is made in



Figure 16.9. Cumulative size distribution in continuous stirred tanks. (a) one tank; (b) two tanks in series, nucleation in both; (c) two tanks in series, nucleation in only the first.

Example 16.5. Teflon heat transfer tubes that are thin enough to flex under the influence of circulating liquid cause a continual descaling that maintains good heat transfer consistently, 20-65 Btu/(hr)(sqft)(°F). Circulating types such as Figures (d) and(e) often are operated in batch mode, the former under vacuum if needed. High labor costs keep application of batch crystallizers to small or specialty production.

(d) Circulating evaporators. Some units are built with internal coils or calandrias and are simply conventional evaporators with provisions for continual removal of crystals. Forced circulation and external heat exchangers provide better temperature control. High velocities in the tubes keep the surfaces scoured. Temperature rise is limited to **3–10°F** per pass in order to control supersaturation and nucleation. Operation under vacuum often is practiced. When the boiling point elevation is not excessive, the off vapors may be recompressed and used again for heating purposes. Multiple effect units in series for thermal economy may be used for crystallizing evaporators as they are for conventional evaporation. Pilot units of 2 ft dia are made, and commercial units up to 40 ft dia or so.

(e) Circulating cooling crystallizers. Such operations are feasible when the solubility falls sharply with decreasing temperature. Coolers usually are applied to smaller production rates than the evaporative types. Cooling is $1-2^{\circ}F$ per pass and temperature differences across the tubes are $5-15^{\circ}F$.

The special designs of Figure 16.11 mostly feature some control of crystal size. They are discussed in order.

(a) Draft tube baffle (DTB) crystallizer. The growing crystals are circulated from the bottom to the boiling surface with a slow moving propeller. Fine crystals are withdrawn from an annular space, redissolved by heating to destroy unwanted nuclei and returned with the feed liquor. The temperature rise caused by mixing of heated feed and circulating slurry is 1-2°F. The fluidized bed of large crystals at the bottom occupies 25–50% of the vessel volume. Holdup time is kept sufficient for crystal growth to the desired size. Products such as KCl, (NH₄)₂SO₄, and (NH₄)H₂PO₄ can be made in this equipment in the range of 6-20 mesh. Reaction and crystallization can be accomplished simultaneously in DTB units. The reactants can be charged into the recirculation line or into the draft tube. Examples are the production of ammonium sulfate from ammonia and sulfuric acid and the neutralization of waste acids with lime. The heat of reaction is removed by evaporation of water.

(b) Direct contact refrigeration. Such equipment is operated as



Figure 16.10. Basic types of batch and continuous crystallizers. (a) Jacketed scraped pipe and assembly of six units (*Riegel*, Chemical Process Machinery, Reinhold, N.Y., 1953). (b) Swenson-Walker jacketed scraped trough (Swenson *Evaporator* Co., *Riegel*, 1953). (c) Batch stirred tank with internal cooling coil (Badger, and *McCabe*, Elements of Chemical Engineering, *McGraw-Hill*, *New York*, *1936*). (d) Crystallization by evaporation, with circulation through an external heater (*Schweitzer*, loc. cit., p. 2.170). (e) Crystallization by chilling, with circulation through an external cooler. (*P.A. Schweitzer, Ed.*, Handbook of Separation Techniques for Chemical Engineers, *McGraw-Hill*, *New York*, *1979*, p. 2.166).









Figure 16.11. Examples of special kinds of crystallizers. (a) Swenson draft tube baffle (DTB) crystallizer; crystals are brought to the surface where growth is most rapid, the baffle permits separation of unwanted fine crystals, resulting in control of size. (b) Direct chilling by contact with immiscible refrigerant, attains very low temperatures and avoids encrustation of heat transfer surfaces. Freons and propane are in common use. (c) Oslo "Krystal" evaporative classifying crystallizer. Circulation is off the top, the fine crystals are destroyed by heating, large crystals grow in the body of the vessel. (d) Twinned crystallizer. When one chamber is maintained slightly supersaturated and the other slightly subsaturated, coarse crystals can be made. (*Nyult*, 1971). (e) APV-Kestner long tube salting evaporator; large crystals (0.5 mm or so) settle out. (f) Escher-Wyss or Tsukushima DP (double propeller) crystallizer. The double propeller maintains upward flow in the draft tube and downward flow in the **annulus**, resulting in highly stable suspensions. (g) A vacuum pan for crystallization of sugar (Honolulu *Iron* Works).



low as -75°F. Essentially immiscible refrigerant is mixed with the liquor and cools it by evaporation. The effluent refrigerant is recovered, recompressed, and recycled. Direct contacting eliminates the need for temperature difference across a heat transfer tube which can be economically more than 5-15°F, and also avoids scaling problems since the liquor must be on the outside of the tubes when refrigerant is used. Examples are crystallization of caustic with freon or propane and of p-xylene with propane refrigerant.

(c) Oslo "Krystal" evaporative classifying crystallizer. The supernatant liquid containing the fines is circulated through the external heater where some of the fines are redissolved because of the temperature rise. The settled large crystals are withdrawn at the bottom. The recirculation rate is much greater than the fresh feed rate. In one operation of $MgSO_4$.7H₂O crystallization, fresh feed saturated at 120°C is charged at 2000 kg/hr to the vessel maintained at 40°C and is mixed with a recirculated rate of 50,000 kg/hr to produce a mixture that is temporarily at 43°C, which then evaporates and cools. Vessel sizes as large as 15 ft dia and 20 ft high are mentioned in the literature. The same principle is employed with cooling type crystallization operations.

(d) Twinned crystallizer. Feed is to the right chamber. The rates of recirculation and forward feed are regulated by the position of the center baffle. Improved degree of uniformity of crystal size is achieved by operating one zone above saturation temperature and the other below. Fine particles are dissolved and the larger ones grow at their expense. Even with both zones at the same temperature, the series operation of two units in series gives more nearly uniform crystal size distribution than can be made in a single stirred tank. It is not stated if any such crystallizers are operated outside Nyvlt's native land, Czechoslovakia, that also produces very fine tennis players (Lendl, Mandlikova, Navratilova, Smid, and Sukova)

(e) APV-Kestner long tube vertical evaporative crystallizers are used to make small crystals, generally less than 0.5 mm, of a variety of substances such as NaCl, Na₂SO₄, citric acid, and others; fine crystals recirculate through the pump and heater.

(f) Escher-Wyss (Tsukushima) double propeller maintains flow through the draft tube and then annulua and maintains highly stable suspension characteristics.

(g) Sugar vacuum pan. This is an example of the highly specialized designs developed in some long-established industries. **Preconcentration** is effected in multiple effect evaporators; then crystallization is accomplished in the pans.

16.6. MELT CRYSTALLIZATION AND PURIFICATION

Some mixtures of organic substances may be separated advantageously by cooling and partial crystallization. The extent of such recovery is limited by the occurrence of eutectic behavior. Examples 16.2 and 16.8 consider such limitations. Sometimes these limitations can be circumvented by additions of other substances that change the phase equilibria or may form easily separated compounds with one of the constituents that are subsequently decomposed for recovery of its constituents.

Thus the addition of n-pentane to mixtures of p-xylene and m-xylene permits complete separation of the xylenes which form a binary eutectic with 11.8% **para**. Without the n-pentane, much **para** is lost in the eutectic, and none of the **meta** is recoverable in pure form. A detailed description of this process is given by Dale (1981), who calls it extractive crystallization. Other separation processes depend on the formation of high melting molecular compounds or clathrates with one of the constituents of the mixture. One example is carbon tetrachloride that forms a compound with p-xylene and alters the equilibrium so that its separation from m-xylene is

facilitated. Hydrocarbons form high-melting hydrates with water; application of propane hydrate formation for the desalination of water has been considered. Urea forms crystalline complexes with straight chain paraffins such as the waxy ingredients of lubricating oils. After separation, the complex may be decomposed at **75–80°C** for recovery of its constituents. This process also is described by Dale (1981). Similarly thiourea forms crystalline complexes with isoparaffins and some cyclic compounds.

Production rates of melt crystallization of organic materials usually are low enough to warrant the -use of scraped surface crystallizers like that of Figure 16.10(a). A major difficulty in the production of crystals is the occlusion of residual liquor on them which cuts the overall purity of the product, especially so because of low temperatures near the eutectic and the consequent high viscosities. Completeness of removal of occluded liquor by centrifugation or filtration often is limited because of the fragility and fineness of the organic crystals.

MULTISTAGE PROCESSING

In order to obtain higher purity, the first product can be remelted and recrystallized, usually at much higher temperatures than the eutectic so that occlusion will be less, and of course at higher concentration. In the plant of Figure 16.12, for instance, occlusion from the first stage is 22% with a content of 8% p-xylene and an overall purity of 80%; from the second stage, occlusion is 9% with a PX content of 42% but the overall purity is 95% PX; one more crystallization could bring the overall purity above 98% or so.

Because the handling of solids is difficult, particularly that of soft organic crystals, several crystallization processes have been developed in which solids do not appear outside the crystallizing equipment, and the product leaves the equipment in molten form. For organic substances, crystalline form and size usually are not of great importance as for products of crystallization from aqueous solutions. If needed, the molten products can be converted into flakes or sprayed powder, or in extreme cases they can be recrystallized out of a solvent.

THE METALLWERK BUCHS PROCESS

The Metallwerk **Buchs** (MWB) process is an example of a batch crystallization that makes a molten product and can be adapted to multistaging when high purities are needed. Only liquids are transferred between stages; no filters or centrifuges are needed. As appears on Figure 16.13, the basic equipment is a vertical thin film shell-and-tube heat exchanger. In the first phase, liquor is recirculated through the tubes as a film and crystals gradually freeze out on the cooled surface. After an appropriate thickness of solid has accumulated, the recirculation is stopped. Then the solid is melted and taken off as product or transferred to a second stage for recrystallization to higher purity.

PURIFICATION PROCESSES

As an alternative to multistage batch crystallization processes with their attendant problems of material handling and losses, several types of continuous column crystallizers have been developed, in which the product crystals are washed with their own melts in countercurrent **flow**. Those illustrated in Figures 16.14-16.17 will be described. Capacities of column purifiers as high as **500 gal/(hr)** (sqft) have been reported but they can be less than one-tenth as much. Lengths of laboratory size purifiers usually are less than three feet.

Schildknecht Column [Fig. 16.14(a)]. This employs a rotating spiral or screw to move the solids in the direction against the flow of

EXAMPLE 16.8

Crystallization from a Ternary Miiture

The case is that of mixtures of the three isomeric nitrotoluenes for which the equilibrium diagram is shown. Point P on the diagram has the composition 0.885 para, 0.085 meta, and 0.030 ortho. The temperature at which crystals begin to form must be found experimentally or it may be calculated quite closely from the heats and temperatures of fusion by a method described for instance by Walas (Example 8.9, Phase Equilibria in Chemical Engineering, Butterworths, Stoneham, MA, 1985). It cannot be found with the data shown on the diagram. In the present case, incipient freezing is at 46°C, with para coming out at point P on the diagram. As cooling

continues, more and more pure para crystals form. The path is along straight line PS which corresponds to constant proportions of the other two isomers since they remain in the liquid phase. At point S, -13° C, which is on the eutectic trough of meta and ortho, the meta also begins to precipitate. Para and meta continue to precipitate along the trough until the ternary eutectic *E* is reached at -40°C when complete solidification occurs. The cooling path is shown on the phase diagram. The recovery of pure para at equilibrium at various temperatures and the composition of the liquid phase are tabulated. (Coulson and Warner, A Problem in Chemical Engineering Design: The Manufacture of Mononitrotoluene, Inst. of Chem. Eng., Rugby, England, 1949).





the fluid. The conveyor is of open construction so that the liquid can flow through it but the openings are small enough to carry the solids. Throughputs of 50 L/hr have been obtained in a 50 mm dia column. Because of the close dimensional tolerances that are needed, however, columns larger than 200mm dia have not been successful. Figure 16.14(a) shows a section for the formation of the crystals, but columns often are used only as purifiers with feed of crystals from some external source.

30

20

10

0

- 10

66.7

75.0

796

82.3

84.8

25.0

20.4

17.7

15.2

bottom. Crystals are charged from an external crystallizer and forced downwards with a reciprocating piston or with pulses from a pump. The washing liquid reflux flows from the melting zone where it is formed upward through the crystal bed and out through the wall filter. Pulse displacement is 0.3-0.6 cm/sqcm of column cross section, with a frequency of 200-250/min. For many applications reflux ratios of 0.05-0.60 are suitable. Evaluation of the proper combination of reflux and length of purifier must be made empirically.

54.0

43.7

35.3

27.1

34.0

41.6

48.0

53.9

12.0

14-7

16.7

17.0

Philips Crystallization Process [Fig. 1414(b)]. The purifying equipment consists of a vessel with a wall filter and a heater at the

From a feed containing 65% p-xylene, a column 1000 sqcm in cross section can make 99% PX at the rate of 550 kg/hr, and 99.8%



Figure 16.12. Humble two-stage process for recovery of p-xylene by crystallization. Yield is 82.5% of theoretical. ML = mother liquor, PX =p-xylene (*Haines, Powers and Bennett, 1955*).



Figure 16.13. MWB (Metallwerk Buchs) batch recirculating crystallizer, with freezing on and melting off insides of thin film heat exchanger tubes; adaptable to multistage processing without external solids handling (*Miitzenberg and Saxer*, 1971).

PX at 100 kg/hr; this process has been made obsolete, however, by continuous adsorption with molecular sieves. Similarly, a feed of 83 mol % of 2-methyl-5-vinyl pyridine has been purified to 95% at the rate of 550 g/hr cm² and 99.7% at 155 g/hr cm². At one time, columns of more than 60 cm dia were in operation.

Brodie Crystallizer-Purifier [Fig. 1614(c)]. This equipment combines a horizontal scraped surface crystallizer with a vertical purifying section. The capacity and performance of the purifier

depends strongly on the sizes of the crystals that enter that zone. In order to ensure adequate crystal size, residence times in the crystallizing zone as long as 24 hr may be needed. No data of residence times are stated in the original article. Some operating data on the **recovery** of **para-dichlorbenzene** from a mixture containing 75% of this material are reported for a purifier that is 1.14 sqft cross section as follows, as well as data for some other materials.

Aeflux ratio	2	0.5	0.25
Feed rate (gal/hr)	29	60	90
Residue rate (gal/hr)		20	30
Product rate (gal/hr)	20	40	60
PDCB in residue (%)		25	25
Product purity (%)	99.997	99.99	99.5

TNO Bouncing Ball Purifier (Fig. 16.15). The basis for this design is the observation that small crystals melt more readily and have a greater solubility than large ones. The purifier is a column with a number of sieve trays attached to a central shaft that oscillates up and down. As the slurry flows through the tower, bouncing balls on each tray impact the crystals and break up some of them. The resulting small crystals melt and enrich the liquid phase, thus providing an upward refluxing action on the large crystals that continue downward to the melting zone at the bottom. Reflux is returned from the melting zone and product is taken off.

Specifications of a pilot plant column are:

diameter, 80 mm, hole size, 0.6 x 0.6 mm, number of balls/tray, 30, diameter of balls, 12 mm, amplitude of vibration, 0.3 mm, frequency, **50/sec**, number of trays, 13, tray spacing, 100 mm.

For the separation of benzene and thiophene that form a solid solution, a tray efficiency of more than 40% could be realized. Flow rates of 100-1000 kg/m^2 hr have been tested. The residence time of crystals was about 30 min per stage. Eutectic systems also have been handled satisfactorily. A column 500 mm dia and 3 m long with 19 trays has been built; it is expected to have a capacity of 300 tons/yr.

Kureha Double-Screw Purifier (*Fig. 1616*). This unit employs a double screw with intermeshing blades that express the liquid from the crystal mass as it is conveyed upward. The melt is formed at the top, washes the rising crystals countercurrently, and leaves as residue at the bottom. A commercial unit has an effective height of 2.6 m and a cross section of 0.31 m^2 . When recovering 99.97% p-dichlorbenzene from an 87% feed, the capacity is 7000metric tons/yr. The feed stock comes from a tank crystallizer and filter. Data on other eutectic systems are shown, and also on separation of 99.87% naphthalene is obtained in this equipment.

Brennan-Koppers Purifier (*Fig. 16.17*). This equipment employs top melting like the Kureha and wall filters like the Philips. Upward movement of the crystals is caused by drag of the flowing fluid. The crystal bed is held compact with a rotating top plate or piston that is called a harvester. It has a corrugated surface that scrapes off the top of the top of the bed and openings that permit the crystals to enter the melting zone at any desired rate. The melt flows downward through the openings in the harvester, washes the



(c)

Figure 16.14. Three types of crystal purifiers with different ways of transporting the crystals. (a) Spiral or screw conveyor type, laboratory scale, but successful up to 200 mm dia [Schildknecht, Z. Anal. Chem. 181, 254 (1961)]. (b) Philips purifier with reciprocating piston or pulse pump drive [McKay, Dale, and Weedman, Ind. Eng. Chem. 52, 197 (1960)]. (c) Combined crystallizer and purifier, gravity flow of the crystals; purifier details on the right (Brodie, 1971).

16.6. MELT CRYSTALLIZATION AND PURIFICATION 547



Figure 16. 15. TNO Crystal Purifier (Arkenbout et al., 1976, 1978).



- 2. Screw Conveyer
- 3. Melter
- 4. Bottom Filter Plate
- 5. Feed Charger
- Output of Product 6.
- 7. Outlet of Bottom Liquid (b)

upwardly moving crystals, and leaves through the sidewall filter as residue. The movement of crystals is quite positive and not as dependent on particle size as in some other kinds of purifiers. Data are given in the patent (U.S. Pat. 4,309,878) about purification of 2,6-ditertiary butyl para cresol; the harvester was operated at 40-60 rpm and filtration rates of 100 lb/(hr)(sqft) were obtained. Other information supplied directly by E.D. Brennan are that a 24in. dia unit stands 9ft high without the mixer and that the following performances have been achieved:

	Diameter	Purity	(wt %)	Prod. Rate	
	(in.)	Feed	Product	(lb/hr/ft²)	
A. Pilot plant tests					
Acetic acid	3	83	99.85	100	
p-Dichlorbenzene	6	70	99.8	380	
Naphthalene (high					
sulfur)	6	68	98	220	
Di-t-butyl-p-cresol	3	85	99.1	210	
<i>,</i> ,	6	8 5	99.1	230	
B. Commercial operation					
Di-t-butyl-p-cresol	24	90	99.5	340	
All feeds were prepared	in Armstron	g scrap	ed surface	crystallizers.	

000 6 2 А Ζ (c)

Figure 16.16. Kureha continuous crystal purifier (KCP column) (Yamada, Shimizu, and Saitoh, in Jancic and DeJong, 1982, pp. 265-270). (a) Flowsketch. (b) Dumbbell-shaped cross section at AA. (c) Details of column and screw conveyor.



Figure 16.17. Brennan-Koppers crystal purifier (Brennan, 1982).

REFERENCES

Crystallization from Solutions

- 1. A.W. Bamforth, Industrial Crystallization, Leonard Hill, London, 1965.
- R.C. Bennett, Crystallization design, Encycl. Chem. Process. Des. 13, 421-455 (1981).
- R.C. Bennett, Crystallization from solution, in *Chemical Engineers'* Handbook, McGraw-Hill, New York, 1984, pp. 19.24–19.40.
- E.D. DeJong and S.I. Jancic, Industrial Crystallization 1978, North-Holland, Amsterdam, 1979.
- Industrial Crystallization, Symposium of Inst. Chem. Eng., Inst. Chem. Eng, London, 1969.
- S.I. Jancic and E.J. DeJong, Eds., *Industrial Crystallization 1981*, North-Holland, Amsterdam, 1982.
- 7. S.I. Jancic and J. Garside, in Ref. 10, 1976, p. 363.
- E.V. Khamskii, Crystallization from Solution, Consultants Bureau, New York, 1969.
- 9. J.W. Mullin, Crystallization, Butterworths, London, 1972.
- J.W. Mullin, Ed., Symposium on Industrial Crystallization, Plenum, New York, 1976.
- J.W. Mullin, Crystallization, Encycl. Chem. Technol. 7, 1978, pp. 243-285.
- J.W. Mullin, Bulk crystallization, in Crystal Growth (Pamplin, Ed.), Pergamon, New York, 1980, pp. 521–565.
- J. Nyvlt, Crystallization as a unit operation in chemical engineering, in Ref. 5, 1969, pp. 1-23.
- J. Nyvlt, Industrial Crystallization from Solutions, Butterworths, London, 1971.
- 15. J. Nyvlt, Industrial Crystallization: The Present State of the Art, Verlag Chemie, Weinheim, 1978.
- M. Ohara and R.C. Reid, *Modelling Crystal Growth Rates from Solution*, Prentice-Hall, Englewood Cliffs, NJ, 1973.
- A.D. Randolph and M.A. Larson, *Theory of Particulate Processes*, Academic, New York, 1971.
- 18. G. Singh, Crystallization from Solution, in Handbook of Separation

Techniques for Chemical Engineers (Schweitzer, Ed.), McGraw-Hill, New York, 1979, pp. 2.151-2.182.

Melt Crystallization

- R. Albertins, W.C. Gates, and J.E. Powers, Column crystallization, in Ref. 32, 1967, pp. 343-367.
- G.J. Arkenbout, Progress in continuous fractional crystallization, Sep. Purification Methods 7(1), 99-134 (1978).
- G.J. Arkenbout, A. vanKujik, and W.M. Smit, Progress in continuous fractional crystallization, in Ref. 10, 1976, pp. 431–435.
- E.D. Brennan (Koppers Co.), Process and Apparatus for Separating and Purifying a Crystalline Material, U.S. Pat. 4,309,878 (12 Jan. 1982).
- J.A. Brodie, A continuous multistage melt purification process, Mech. Chem. Eng. Trans., Inst. Eng. Australia, 37-44 (May 1971).
- G.H. Dale, Crystallization: extractive and adductive, Encycl. Chem. Process. Des. 13, 456–506 (1981).
- 25. R.A. Findlay, Adductive crystallization, in New Chemical Engineering Separation Techniques (Schoen, Ed.), Wiley-Interscience, New York, 1958.
- 26. R.A. Findlay and J.A. Weedman, Separation and purification by crystallization, in Advances in Petroleum Chemistry and Refining, Wiley-Interscience, New York, 1958, Vol. 1, pp. 118-209.
- H.W. Haines, J.M. Powers, and R.B. Bennett, Separation of xylenes, Ind. Eng. Chem. 47, 1096 (1955).
- J.D. Henry and C.C. Moyers, Crystallization from the melt, in *Chemical Engineers' Handbook*, McGraw-Hill, New York, 1984, pp. 17.2–17.12.
- D.L. McKay, Phillips fractional solidification process, in Ref. 32, 1967, pp. 427-439.
- A.B. Mützenberg and K. Saxer, The MWB crystallizer, Dechema Monographien 66, 313-320 (1971).
- J. Yamada, C. Shimizu, and S. Saitoh, Purification of organic chemicals by the Kureha Continuous Crystal Purifier, in Ref. 6, 1982, pp. 265-270.
- M. Zief and W.R. Wilcox, Fractional Soldification, Dekker, New York, 1967, Vol. 1.

17.1. DESIGN BASIS AND SPACE VELOCITY

This chapter summarizes the main principles of chemical kinetics and catalysis; also it classifies and describes some of the variety of equipment that is suitable as chemical reactors. Because of the diversity of the behavior of chemical reactions, few rules are generally applicable to the design of equipment for such purposes. Reactors may be stirred tanks, empty or packed tubes or vessels, shell-and-tube devices or highly specialized configurations, in any of which heat transfer may be provided. These factors are balanced in individual cases to achieve economic optima. The general rules of other chapters for design of pressure vessels, heat exchangers, agitators, and so on naturally apply to reactors.

DESIGN BASIS

Although the intent of this chapter is not detailed design, it is in order to state what is included in a proper design basis, for example at least these items:

- 1. Stoichiometry of the participating reactions.
- 2. Thermal and other physical properties.
- 3. Heats of reaction and equilibrium data.
- 4. Rate of reaction, preferably in equation form, relating it to composition, temperature, pressure, impurities, catalysts and so on. Alternately tabular or graphical data relating compositions to time and the other variables listed.
- 5. Activity of the catalyst as a function of onstream time.
- 6. Mode of catalyst reactivation or replacement.
- 7. Stability and controllability of the process.
- 8. Special considerations of heat and mass transfer.
- 9. Corrosion and safety hazards.

REACTION TIMES

In practical cases reaction times vary from fractions of a second to many hours. The compilation of Table 17.1 of some commercial practices may be a basis for choosing by analogy an order of magnitude of reactor sizes for other processes.

For ease of evaluation and comparison, an apparent residence time often is used instead of the true one; it is defined as the ratio of the reactor volume to the inlet volumetric flow rate,

$$\bar{t}_{\rm app} = V_r / V_0'.$$

On the other hand, the true residence time must be found by integration,

$$\bar{t} = \int dV_r/V' = \int dn'/rV'.$$

Since the rate of reaction r and the volumetric flow rate V' at each position depend on T, P, and local molal flow rate n' of the key component of the reacting mixture, finding the true residence time is an involved process requiring many data. The easily evaluated apparent residence time usually is taken as adequate for rating sizes of reactors and for making comparisons.

A related concept is that of space velocity which is the ratio of a flow rate at $STP(60^{\circ}F, 1 \text{ atm usually})$ to the size of the reactor.

The most common versions of space velocities in typical units are:

- GHSV (gas hourly space velocity) = (volumes of feed as gas at **STP/hr)/(volume** of the reactor or its content of catalyst) = (SCFH gas feed)/cuft.
- LHSV (liquid hourly space velocity) = (Volume of liquid feed at 60°F/hr)/volume of reactor or catalyst) = (SCFH liquid feed)/cuft.
- WHSV (weight hourly space velocity) = (lb of feed/hr)/(lb of catalyst). Other combinations of units of the flow rate and reactor size often are used in practice, for instance.
- **BPSD/lb** = (barrels of liquid feed at 60° F per stream day)/(lb catalyst), but it is advisable to write out such units in each case to avoid confusion with the standard meanings of the given acronyms. Since the apparent residence time is defined in terms of the actual inlet conditions rather than at standard T and P, it is not the reciprocal of GHSV or LHSV, although the units are the same.

17.2. BATE EQUATIONS AND OPERATING MODES

The equations of this section are summarized and extended in Table 17.2. The term "rate of reaction" used here is the rate of decomposition per unit volume,

$$r_a = -\frac{1}{V}\frac{dn_a}{dt}$$
, mol/(unit time)(unit volume). (17.1)

A rate of formation will have the opposite sign. When the volume is constant, the rate is the derivative of the concentration

$$r_a = -\frac{dC_o}{dt}$$
, at constant volume. (17.2)

In homogeneous environments the rate is expressed by the law of mass action in terms of powers of the concentrations of the reacting substances

$$\mathbf{r}_a = -\frac{1}{V} \frac{d\mathbf{n}_a}{dt} = k C_a^{\alpha} C_b^{\beta} \cdots$$
(17.3)

When the reaction mechanism truly follows the stoichiometric equation

$$\mathbf{v}_a \mathbf{A} + \mathbf{v}_b \mathbf{B} + \dots \rightarrow \text{products},$$
 (17.41)

the exponents are the stoichiometric coefficients; thus,

$$\boldsymbol{r}_{\boldsymbol{a}} = \boldsymbol{k} (\boldsymbol{C}_{\boldsymbol{a}})^{\boldsymbol{v}_{\boldsymbol{a}}} (\boldsymbol{C}_{\boldsymbol{b}})^{\boldsymbol{v}_{\boldsymbol{b}}} \dots, \qquad (17.5)$$

but α , β , . . . often are purely empirical values-integral or nonintegral, sometimes even negative.

The coefficient k is called the specific rate. It is taken to be independent of the concentrations of the reactants but does depend primarily on temperature and the nature and concentration of

TABLE 17.1. Residence Times and/or Space Velocities in industrial Chemical Reactors

	Product		React	or	Cond	litions	Residence time or space	Source
	(raw materials)	Туре	phas	e Catalyst	<i>T,</i> °C	P, atm	velocity	and page
1.	Acetaldehyde (ethylene, air)	FΒ	L	Cu and Pd chlorides	50-100	а	6-40 min	[2] 1, [7] 3
2.	Acetic anhydride (acetic acid)	то	L	Triethyl phosphate	700-800	0.3	0.25-5 S	[2]
3.	Acetone (i-propanol)	МT	LG	Ni	300	1	2.5 h	[1] 1314
4.	Acrolein (formaldehyde, acetaldehyde)	FL	G	MnO, silica gel	280-320	1	0.6 s	[<i>1</i>] / 384, [7] 33
5.	Acrylonitrile (air, propylene, ammonia)	FL	G	Bi phosphomolybdate	400	1	4.3 s	[3] 684, [2] 47
6.	Adipic acid (nitration of cyclohexanol)	то	L	co naphthenate	125-160	4-20	2 h	[2] 51, [7] 49
7.	Adiponitrile (adipic acid)	FΒ	G	H₃BO₃ H₃PO₄	370-410	1	3.5-5 s 350-500 GHSV	[<i>1</i>] 2 152, [<i>7</i>] 52
8.	Alkylate (<i>i</i> -C ₄ , butenes)	CST	L	H ₂ SO ₄	5-10	2-3	5-40 min	[4] 223
9.	Alkylate (<i>i</i> -C ₄ , butenes)	CST	L	HF	25-38	8-11	5-25 min	[4] 223
10.	Allyl chloride (propylene. Cl_2)	то	G	N.A.	500	3	0.3-1.5 S	[<i>1</i>] 2 416, [7] 67
11.	Ammonia (H_2, N_2)	FB	G	Fe	450	150	28 s 7,800 GHSV	[<i>6</i>] 61
12.	Ammonia (H_2 , N_2)	FΒ	G	Fe	450	225	33 s 10,000 GHSV	[6] 61
13.	Ammonia oxidation	Flame	G	Pt gauze	900	8	0.0026 \$	[6] 115
14.	Aniline (nitrobenzene, H ₂)	В	L	$FeCl_2$ in H_2O	95-100	1	8 h	[1] 3 289
15.	Aniline (nitrobenzene, H ₂)	FΒ	G	Cu on silica	250-300	1	0.5-100 S	[7] 82
16.	Aspirin (salicylic acid, acetic anhydride)	В	L	None	90	1	> 1 h	[7] 89
17.	Benzene (toluene)	TU	G	None	740	38	48 s 815 GHSV	[6] 36, [9] 109
18.	Benzene (toluene)	TU	G	None	650	3 5	128 S	[<i>1</i>] 4 183, [7] 98
19.	Benzoic acid (toluene, air)	SCST	LG	None	125-175	9-13	0.2-2 h	[7] 101
20.	Butadiene (butane)	FΒ	G	Cr_2O_3 , Al_2O_3	750	1	0.1-I S	[7] 118
21.	Butadiene (I-butene)	FΒ	G	None	600	0.25	0.001 s 34,000 GHSV	[3] 572
22.	Butadiene sulfone (butadiene, SO ₂)	CST	L	<i>t</i> -butyl catechol	34	12	0.2 LHSV	[1] 5 192
23.	i-Butane (n-butane)	FΒ	L	AICI ₃ on bauxite	40 -120	18-36	0.5-1 LHSV	[4] 239, [7] 683
24.	i-Butane (n-butane)	FΒ	L	Ni	370-500	20-50	I-6 WHSV	[4] 239
25.	Butanols (propylene hydroformylation)	FΒ	L	PH ₃ -modified Co carbonyls	150-200	1,000	100 g/L-h	[1] 5 373
26.	Butanols (propylene hydroformylation)	FΒ	L	Fe penta- carbonyl	110	10	1 h	[7] 125
27. 28.	Calcium stearate Caprolactam	B CST	L I L	None Polyphos-	180 80-110	5 1	l-2 h 0.25-2 h	[/] 135 [<i>1</i>] 6 73,
29.	(cyclohexane oxime) Carbon disulfide (methane, sulfur)	Furn.	G	phoric acid None	500-700	1	1.0 S	[/] 139 [<i>1</i>] 6 322, [7] 144
30.	Carbon monoxide oxidation (shift)	TU	G	Cu-Zn or Fe ₂ O ₃	390-220	26	4.5 s 7,000 GHSV	[6] 44
30'.	Port. cement	Kiln	S		1400-1700	t	10 h	[11]

TABLE 17.1—(continued)

	Product		Reacto	or	Cond	litions	Residence time or space	Source
	(raw materials)	Туре	phase	e Catalyst	<i>T,</i> ℃	P, atm	velocity	and page
31.	Chloral (CI,, acetaldehyde)	CST	LG	None	20-90	1	140 h	[7] 158
32.	Chlorobenzenes (benzene, Cl_2)	SCST	LG	Fe	40	1	24 h	[1] 8 122
33.	Coking, delayed theater)	TU	LG	None	490-500	15-4	250 s	[1] 108
34.	Coking, delayed (drum, 100 ft max.)	В	LG	None	500-440	4	0.3-0.5 ft/s vapor	[1] 108
35.	Cracking, fluid- catalytic	FL	G	SiO_2 , Al_2O_3	470- 540	2-3	0.5-3 WHSV	[4] 162
36.	Cracking, hydro- (gas oils)	FΒ	LG	Ni, SiO ₂ , Al ₂ O ₃	350-420	100-150	I-2 LHSV	[11]
37.	Cracking (visbreaking residual oils)	TU	LG	None	470-495	IO-30	450 s 8 LHSV	[11]
38.	Cumene (benzene, propylene)	FΒ	G	H₃PO₄	260	35	23 LHSV	[11]
39.	Cumene hydroperoxide (cumene, air)	CST	L	Metal porphyrins	95-120	2-15	l-3 h	[7] 191
40.	$\begin{array}{llllllllllllllllllllllllllllllllllll$	FΒ	G	Nion Al_2O_3	150-250	25-55	0.75-2 LHSV	[7] 201
41.	Cyclohexanol (cyclohexane, air)	SCST	LG	None	185-200	48	2-10 min	[7] 203
42.	Cyclohexanone (cyclohexanol)	CST	L	N.A.	107		0.75 h	[8] (1963)
43.	Cyclohexanone (cyclohexanol)	ΜТ	G	Cu on pumice	250-350		4-12 s	[8] (1963)
44.	Cyclopentadiene Idicyclopentadiene)	TU	G	None	220-300	I - 2	0.1-0.5 LHSV	[7] 212
45.	DDT (chloral, chlorobenzene)	В	L	Oleum	O-15		8 h	[7] 233
46.	Dextrose (starch)	CST	L	H₂SO₄	165		20 min	[<i>8</i>] (1951)
47.	Dextrose (starch)	CST	L۰	Enzyme	60		100 min	[7] 217
48.	Dibutylphthalate (phthalic anhydride, butanol)	В	L	H₂SO₄	150-200		l-3 h	[7] 227
49.	Diethylketone (ethylene, CO)	то	L	Co oleate	150-300	200-500	0.1-10 h	[7] 243
50.	Dimethylsulfide (methanol, CS_2)	FB	G	Al ₂ O ₃	375-535	5	150 GHSV	[7] 266
51.	Diphenyl (benzene)	ΜT	G	None	730	2	0.6 s 3.3 LHSV	[7] 275, [<i>8</i>] (1938)
52.	Dodecylbenzene (benzene, propylene tetramer)	CST	L	AICI3	15-20		I-30 min	[7] 283
53.	Ethanol (ethylene, H ₂ O)	FΒ	G	H₃PO₄	300	82	1,800 GHSV	[2] 356, [7] 297
54.	Ethyl acetate (ethanol, acetic acid)	TU, CST	L	H₂SO₄	100	t	0.5-0.8 LHSV	[<i>10</i>] 45, 52, 58
55.	Ethyl chloride (ethylene, HCl)	то	G	ZnCl ₂	150-250	6-20	2 s	[7] 305
56.	Ethylene (ethane)	TU	G	None	860		1.03 s 1,880 GHSV	l31 411, [6] 13
57.	Ethylene (naphtha)	ΤU	G	None	550-750	2-7	0.5-3 s	[7] 254
58.	Ethylene, propylene chlorohydrins (Cl ₂ , H ₂ O)	CST	LG	None	30-40	3-10	0.5-5 min	[7] 310 , 580

(continued)

TABLE **17.1**—(continued)

							Residence	
	Product		React	or	Cor	ditions	time or space	Source
	(raw materials)	Type	phase	e Catalyst	<i>T.</i> °C	P, atm	velocity	and page
59.	Ethylene glycol	то	LG	1% H₂SO₄	50-70	1	30 min	[2] 398
60.	Ethylene glycol (ethylene oxide, H_2O)	то	LG	None	195	13	1 h	[2] 398
61.	Ethylene oxide (ethylene, air)	FL	G	Ag	270-290	1	1 s	[2] 409 , [7] 322
62.	Ethyl ether (ethanol)	FΒ	G	WO₃	120- 375	2-100	30 min	[7] <i>326</i>
63.	Fatty alcohols (coconut oil)	В	L	Na, solvent	142	1	2 h	[<i>8</i>] (1953)
64.	Formaldehyde (methanol, air)	FΒ	G	Ag gauze	450-600	1	0.01 S	[2] 423
65.	Glycerol (allyl alcohol, H ₂ O ₂)	CST	L	H₂WO₄	40-60	1	3 h	[7] 34 7
66.	Hydrogen (methane, steam)	МΤ	G	Ni	790	13	<i>5.4 s</i> <i>3,000</i> GHSV	[6] 133
67.	Hydrodesulfurization of naphtha	то	LG	Co-Mo	315- 500	20- 70	1. 5-8 LHSV 125 WHSV	[4] 285, [6] 179, [9] 201
68 .	Hydrogenation of cottonseed oil	SCST	LG	Ni	130	5	6 h	[6] 161
69.	Isoprene (<i>i</i> -butene,	FB	G	HCI, silica nel	250-350	1	1 h	[7] 389
70.	Maleic anhydride (butenes, air)	FL	G	V ₂ O ₅	300- 450	2-10	0.1-5 s	[7] 406
71.	Melamine (urea)	В	L	None	340- 400	40-150	5-60 min	[7] 410
72.	Methanol (CO, H ₂)	FB	G	ZnO, Cr_2O_3	350-400	340	5, 000 GHSV	[7] 421
73.	Methanol (CO, H_2)	FB	G	ZnO,Cr ₂ O ₃	350-400	254	28, 000 GHSV	[3] 562
74.	o-Methyl benzoic acid (xylene, air)	CST	L	None	160	14	0.32 h 3.1 LHSV	[3] 732
75.	Methyl chloride (methanol, Cl,)	FB	G	Al ₂ O ₃ gel	340- 350	1	275 GHSV	[2] 533
76.	Methyl ethyl ketone (2-butanol)	FB	G	ZnO	425-475	2-4	0. 5- 10 min	[7] 437
77.	Methyl ethyl ketone (2-butanol)	FB	G	Brass spheres'	450	5	<i>2.1 s</i> 13 LHSV	[10] 284
78.	(benzene HNO ₂)	CST	L	H ₂ SO ₄	45-95	1	3-40 min	[7] 468
79.	Nitromethane	то	G	None	450- 700	5-40	0.07-0.35 s	[7] 474
80.	Nylon-6 (caprolactam)	TU	L	N a	260	1	<i>12</i> h	[7] 480
81.	Phenol (cumene hydroperoxide)	CST	L	SO,	45-65	2-3	<i>15</i> min	[7] <i>520</i>
82.	Phenol (chloro- benzene steam)	FB	G	Cu. Ca phosphate	430-450	1-2	2 WHSV	[7] 522
83.	Phosgene (CO, Cl_2)	MT	G	Activated	50	5-10	16 s 900 GHSV	[11]
84.	Phthalic anhydride (o-xylene, air)	ΜТ	G	V_2O_5	350	1	1.5 s	[3] 482, 539, [7] 530
85.	Phthalic anhydride (naphthalene, air)	FL	G	V_2O_5	350	1	5 s	[7] 329 [9] 136, [10] 335
86.	Polycarbonate resin (bisphenol-A, phospene)	В	L	Benzyltri- ethylammonium chloride	30-40	1	0. 25-4 h	[7] 452
87.	Polyethylene	TU	L	Organic	180-200	1. 000- 1. 700	0. 5 - 50 min	[7] 547
88.	Polyethylene	TU	L	Cr_2O_3 , Al_2O_3 , SiO ₂	70-200	20- 50	0.1-1,000 s	[7] 549

TABLE 17.1—(continued)

	Desident		Deserve	_	Cond	litiano	Residence time or	Source
	(raw materials)	Туре	phase	Catalyst	<i>T,</i> °C	P, atm	velocity	and page
89.	Polypropylene	то	L	R₂AICI, TiCl₄	15-65	IO- 20	15-100 min	[7] <i>559</i>
90.	Polyvinyl chloride	В	L	Organic peroxides	60	10	5. 3- 10 h	[6] 139
91.	<i>i</i> -Propanol (propylene, H ₂ O)	то	L	H_2SO_4	70-110	2-14	0. 5-4 h	[7] 393
92.	Propionitrile (propylene, NH ₃)	TU	G	Co0	350- 425	70-200	0.3-2 LHSV	[7] 578
93.	Reforming of naphtha $(H_2/hydrocarbon = 6)$	FΒ	G	Pt	490	30-35	<i>3</i> LHSV <i>8,000</i> GHSV	[6] 99
94.	Starch (corn, H ₂ O)	В	L	SO₂	25-60	1	18-72 h	[7] 607
95.	Styrene (ethylbenzene)	ΜТ	G	Metal oxides	600- 650	1	<i>0.2 s</i> <i>7,500</i> GHSV	[5] 424
96.	Sulfur dioxide oxidation	FΒ	G	$V_{2}O_{5}$	475	1	<i>2.4</i> S 700 GHSV	161 86
97.	<i>t</i> -Butyl methacrylate (methacrylic acid, <i>i</i> -butene)	CST	L	H₂SO₄	25	3	0.3 LHSV	' [I] 5 328
98.	Thiophene (butane, S)	TU	G	None	600- 700	1	0.01-l s	[7] 652
99.	Toluene diisocyanate (toluene diamine, phosgene)	В	LG	None	200-210	1	7 h	[7] 657
100.	Toluene diamine (dinitrotoluene, H_2)	В	LG	Pd	80	6	10 h	[7] 656
101.	Tricresyl phosphate (cresyl, POCl ₃)	то	L	MgCl ₂	150-300	1	0.5-2.5 h	[2] 850, [7] 673
102.	Vinyl chloride (ethylene, Cl ₂)	FL	G	None	450- 550	2-10	0.5-5 s	[7] 699

Abbreviations

Reactors: batch (B), continuous stirred tank (CST), fixed bed of catalyst (FB), fluidized bed of catalyst (FL), furnace (Furn.), multitubular (MT), semicontinuous stirred tank (SCST), tower (TO), tubular (TU). Phases: liquid (L), gas (G), both (LG).

Space velocities (hourly): gas (GHSV), liquid (LHSV), weight (WHSV). Not available (N.A.)

REFERENCES

- J. J. McKetta, ed., "Encyclopedia of Chemical Processing and Design," Marcel Dekker, New York, 1976 to date (referenced by volume).
- W.L. Faith, D.B. Keyes, and R.L. Clark, "Industrial Chemicals," revised by F.A. Lowenstein and M.K. Moran, John Wiley & Sons, New York, 1975.
- 3. G.F. Froment and K.B. Bischoff, "Chemical Reactor Analysis and Design," John Wiley & Sons, New York, 1979.
- R.J. Hengstebeck, "Petroleum Processing," McGraw-Hill, New York, 1959.
- V.G. Jenson and G.V. Jeffreys, "Mathematical Methods in Chemical Engineering," 2nd ed., Academic Press, New York, 1977.

- H.F. Rase, "Chemical Reactor Design for Process Plants: Vol. 2, Case Studies," John Wiley & Sons, New York, 1977.
- M. Sittig, "Organic Chemical Process Encyclopedia," Noyes, Park Ridge, N.J., 1969 (patent literature exclusively).
- Student Contest Problems, published annually by AIChE, New York (referenced by year).
- 9. M.O. Tarhan, "Catalytic Reactor Design," McGraw-Hill, New York, 1983.
- K.R. Westerterp, W.P.M. van Swaaij, and A.A.C.M. Beenackers, "Chemical Reactor Design and Operation," John Wiley & Sons, New York, 1984.
- 11. Personal communication (Walas, 1985).

catalysts. Temperature dependence usually is represented by

$$k = k_{\infty} \exp(-E/RT) = \exp(a' - b'/T), \qquad (17.6)$$

where E is the energy of activation.

Specific rates of reactions of practical interest cannot be found by theoretical methods of calculation nor from correlations in terms of the properties of the reactants. They must be found empirically in every case together with the complete dependence of the rate of reaction on concentrations, temperature, and other pertinent factors. The analysis of experimental data will be ignored here since the emphasis is placed on the use of known rate equations.

Integration of the rate equation is performed to relate the composition to the reaction time and the size of the equipment. From a rate equation such as

$$-\frac{dC_a}{dt} = kC_a^{\alpha}C_b^{\beta}C_c^{\gamma},\tag{17.7}$$

1. The reference reaction is

 $v_s A + v_b B + \cdots \rightarrow v_r R + v_s S + \cdots$ $\Delta v = v_r + v_s + \cdots = (v_a + v_b + \cdot)$

2. Stoichiometric balance for any component i,

$$\begin{split} n_i &= n_{i0} \pm (v_i / v_s) (n_{s0} - n_s) \\ & \left\{ \begin{array}{l} + \text{ for product (right-hand side, RHS)} \\ - \text{ for reactant (left-hand side, LHS)} \end{array} \right. \\ C_i &= C_{i0} \pm (v_i / v_s) (C_{s0} - C_s), \text{ at constant Tand V only} \\ n_t &= n_{t0} + (\Delta v / v_s) (n_{s0} - n_s) \end{split} \end{split}$$

3. Law of mass action

$$r_{a} = -\frac{1}{V_{r}} \frac{dn_{a}}{dt} = kC_{a}^{v_{a}}C_{b}^{v_{b}} \cdots$$

= $kC_{a}^{v_{a}}[C_{b0} \quad (v_{b}/v_{a})(C_{a0} \quad C_{a})]^{v_{b}}$
 $r_{a} = kC_{a}^{\alpha}[C_{b0} - (v_{b}/v_{a})(C_{a0} - C_{a})]^{\beta}$

where it is not necessarily true that $\alpha = v_a, \beta = v_b, .$ 4. At constant volume, $C_a = n_a/V_r$

$$kt = \int_{C_{a}}^{C_{b0}} \frac{1}{C_{a}^{\alpha} [C_{b0} - (v_{b}/v_{a})(C_{a0} - C_{a})]^{\beta} \dots} dC_{a}$$

$$kt = \int_{n_{a}}^{n_{b0}} \frac{V_{a}^{-1+\alpha+\beta}}{n_{a}^{\alpha} [n_{b0} + (v_{b}/v_{a})(n_{a0} - n_{a})]^{\beta} \dots} -dn_{a}$$

Completed integrals for some values of α and β are in Table 17.3 5. Ideal gases at constant pressure:

$$V_r = \frac{n_r RT}{P} = \frac{R}{P} \frac{T}{P} \left[n_{t0} + \frac{\Delta v}{v_a} (n_{a0} - n_a) \right]$$

$$r_a = kC_a^{\alpha}$$

$$kt = \left(\frac{RT}{P}\right)^{\alpha - 1} \int_{n_a}^{n_{a0}} \frac{[n_{t0} + (\Delta v/v_a)(n_{a0} - n_a)]^{\alpha - 1}}{n_a^{\alpha}} dn_a$$

6. Temperature effect on the specific rate:

$$\mathbf{k} = k_{\infty} \exp(-E/RT) = \exp(a' \quad b'/T)$$

E = energy of activation

- 7. Simultaneous reactions: The overall rate is the algebraic sum of the rates of the individual reactions. For example, take the three reactions:
 - 1. $A + B \xrightarrow{k_1} C + D$. 2. $C + D \xrightarrow{k_2} A + B$. 3. $A + C \xrightarrow{k_3} E$.

The rates are related by:

$$\begin{split} r_{a} &= r_{a1} + r_{a2} + r_{a3} = k_{1}C_{a}C_{b} = k_{2}C_{c}C_{d} + k_{3}C_{a}C_{c} \\ r_{b} &= -r_{d} = k_{1}C_{a}C_{b} = k_{2}C_{c}C_{d} \\ r_{c} &= -k_{1}C_{a}C_{b} + k_{2}C_{c}C_{d} + k_{3}C_{a}C_{c} \\ r_{\theta} &= -k\&C, \end{split}$$

The number of independent rate equations is the same as the number of independent stoichiometric relations. In the present example, reactions 1 and 2 are a reversible reaction and are not independent. Accordingly, C_c and C_{dr} , for example, can be

eliminated from the equations for r_a and r_b which then become an integrable system. Usually only systems of linear differential equations with constant coefficients are solvable analytically. Many such cases are treated by Rodiguin and Rodiguina (1964) Consecutive Chemical Reactions, Van Nostrand, N.Y.

8. Mass transfer resistance:

$$C_{ai} = \text{interfacial concentration of reactant A}$$

$$r_{o} = -\frac{dC_{o}}{dt} = k_{d}(C_{o} - C_{ai}) = kC_{ai}^{\alpha} = k\left(C_{a} - \frac{r_{a}}{k_{d}}\right)^{\alpha}$$

$$kt = \int_{C_{a}}^{C_{o0}} \frac{1}{(C_{o} - r_{a}/k_{d})^{\alpha}} dC_{o}$$

The relation between r_a and C_a must be established (numerically if need be) from the second line before the integration can be completed

- 9. Solid-catalyzed reactions, some Langmuir-Hinshelwood mechanisms for the reference reaction A + $B \rightarrow R$ + S.
 - 1. Adsorption rate of A controlling

$$\begin{aligned} r_{B} &= -\frac{1}{V}\frac{dn_{a}}{d} = \frac{kP_{a}g_{VV}}{kP_{a}} \\ \theta_{V} &= 1 / \left[1 + \frac{K_{a}P_{c}P_{a}}{K_{b}} + K_{b}P_{b} + K_{r}P_{r} + K_{s}P_{s} + K_{l}P_{l} \right] \\ K_{a} &= P_{r}P_{s}/P_{a}P_{b} \text{ (equilibrium constant)} \end{aligned}$$

I is an adsorbed substance that is chemically inert 2. Surface reaction rate controlling:

$$r = kP_{a}P_{b}\theta_{v}^{2}$$

$$\theta_{v} = 1 / (1 + \sum K_{j}P_{j}),$$

summation over all substances absorbed

 Reaction A,+ B→R +S, with A, dissociated upon adsorption and with surface reaction rate controlling:

$$\begin{aligned} r_{a} &= kP_{a}P_{b}\theta_{v}^{3} \\ \theta_{v} &= 1/(1 + \sqrt{K_{a}P_{a}} + K_{b}P_{b} + \cdots) \end{aligned}$$

4. At constant P and T the P_i are eliminated in favor of n_i and the total pressure by

$$P_{a} = \frac{n_{a}}{n_{t}} P$$

$$P_{i} = \frac{n_{i}}{n_{t}} P = \frac{n_{i0} \pm (v_{i}/v_{a})(n_{a0} - n_{a})}{n_{t0} + (\Delta v/v_{a})(n_{a0} - n_{a})} P$$

$$\begin{cases} + \text{ for products, RHS} \\ -\text{for reactants, LHS} \end{cases}$$

$$V = \frac{n_{t}RT}{P}$$

$$kt = \int_{n_{a}}^{n_{a0}} \frac{1}{VP_{a}P_{b}\theta_{v}^{2}}, \text{ for a case (2) batch reaction}$$



TABLE 17.2—(continued)

10. A continuous stirred tank reactor battery (CSTR) Material balances:

$$n_{a0}' = n_a' + r_{a1}V_{r1}$$

 $n'_{a,i-1} = n'_{ai} + r_{ai} V_{ri}$, for the jth stage

For a first order reaction, with $r_a = kC_{ar}$

$$\frac{C_{\theta j}}{C_{\theta 0}} = \frac{1}{(1 + k_1 \tilde{t}_1)(1 + k_2 \tilde{t}_2) \cdots (1 + k_j \tilde{t}_j)} = \frac{1}{(1 + k_1 \tilde{t}_j)^{j'}}$$

for jtanks in series with the same temperatures and residence times $\bar{t}_i = V_{ri}/V'_{i}$, where V is the volumetric flow rate

11, Plug flow reactor (PFR);

$$r_{\theta} = -\frac{dn'_{\theta}}{dV_r} = k_{\theta} \mathcal{C}_{D}^{\alpha} \mathcal{C}_{L}^{\beta}.$$
$$= k(n'_{\theta}/V')^{\alpha} (n'_{D}/V')^{\beta}...$$

the concentrations C_b and C_c first must be eliminated with the aid of the stoichiometric equation of the process. Item 4 of Table 17.2 is an example. When several reactions occur simultaneously, the overall rate of a particular participant is the algebraic sum of its rates in individual reactions. Item 7 of Table 17.2 is an example. The number of differential equations representing the reacting system is the same as the number of independent stoichiometric equations; appropriate concentrations are eliminated with stoichiometry to develop an integrable set of equations. Integrals of **compnon** isothermal, constant volume rate equations are summarized in Table 17.3, and a simple case of a process at constant pressure is item 5 of Table 17.3.

An overall conversion rate may depend on rates of mass transfer between phases as well as chemical rates. In the simplest case, mass transfer and chemical transformation occur in series; advantage is taken of the equality of these two rates at steady state conditions to eliminate interfacial concentrations from the rate equations and thus to permit integration. Item 8 of Table 17.2 is an example.

Rates of fluid phase reactions catalyzed by solids also can be represented at least approximately by powers of the concentrations. A more fundamental approach, however, takes into account mechanisms of adsorption and of reaction on the catalyst surface. A few examples of resulting equations are in item 9 of Table 17.2.

Practical solid-catalyzed rate processes also may be influenced by rates of diffusion to the external and internal surfaces. In the latter case the rate equation is modified by inclusion of a catalyst effectiveness to become

$$\mathbf{r}_{a} = k\eta f(C_{a}). \tag{17.8}$$

The effectiveness is a measure of the utilization of the internal surface of the catalyst. It depends on the dimensions of the catalyst particle and its pores, on the diffusivity, specific rate, and heat of reaction. With a given kind of catalyst, the only control is particle size to which the effectiveness is proportional; a compromise must be **made** between effectiveness and pressure drop. In simple cases η can be related mathematically to its parameters, but in such important practical cases as ammonia synthesis its dependence on parameters is complex and strictly empirical. Section 17.5 deals with this topic.

Reaction processes may be conducted under nonflow or steady



12. Material and energy balances for batch, CSTR and PFR are in Tables 17.4, 17.5, and 17.6

13. Notation

A, \mathbf{B} , \mathbf{R} , \mathbf{S} are participants in the reaction; the letters also are used to represent concentrations

 $C_i \approx n_i / V_r$ or n_i' / V' , concentration

 $n_i \approx$ mols of component *i* in the reactor

n' = molal flow rate of component i

 $V_{r} =$ volume of reactor

V' = volumetric flow rate

 $v_i =$ stoichiometric coefficient

 r_i = rate of reaction of substance *i* [mol/(unit time)(unit volume)]

 α,β = empirical exponents in a rate equation

flow conditions. One mode of the latter is tubular flow or, in the limiting case, plug flow, in which all molecules have substantially the same residence time. The rate equation for a plug flow reactor (PFR) is

$$r_a = -\frac{dn'_a}{dV_r} = kC^{\alpha}_a C^{\beta}_b \dots = k \left(\frac{n'_a}{V'}\right)^{\alpha} \left(\frac{n'_b}{V'}\right)^{\beta} \dots, \qquad (17.9)$$

where V_r is the reactor volume and the primes (') designate flow rates. Flow reactions of gases take place at substantially constant pressure so that V' will depend on the extent of conversion if there is a change in the number of mols. Item 11 of Table 17.2 is an example of the rate equation for such conditions.

The other mode of flow reaction employs one or more stirred tanks in series, which is called a continuous stirred tank (CSTR) battery. The rate of reaction in a single tank is

$$r_{a} = \frac{n_{a0}' - n_{a}'}{V_{r}} \simeq \frac{C_{a0} - C_{a}}{V_{r}/V'} = \frac{C_{a0} - C_{a}}{\bar{t}} = k C_{a}^{\alpha} C_{b}^{\beta} \cdots .$$
(17.10)

The relation in terms of concentrations is valid if the volumetric rates into and out of the tank are substantially the same. Stirring is assumed sufficient to maintain uniform composition and temperature in the tank; then the effluent conditions are the same as those of the tank. Relations for several tanks in series are in item 10 of Table 17.2.

17.3. MATERIAL AND ENERGY BALANCES OF REACTORS

All chemical reactions are accompanied by some heat effects so that the temperature will tend to change, a serious result in view of the sensitivity of most reaction rates to temperature. Factors of equipment size, controllability, and possibly unfavorable product distribution of complex reactions often necessitate provision of means of heat transfer to keep the temperature within bounds. In practical operation of **nonflow** or tubular Row reactors, truly isothermal conditions are not feasible even if they were desirable. Individual continuous stirred tanks, however, do maintain substantially uniform temperatures at steady state when the mixing is intense enough; the level is determined by the heat of reaction as well as the rate of heat transfer provided.

In many instances the heat transfer aspect of a reactor is

TABLE 17.3. Some Isothermal Rate Equations and Their Integrals

1. $A \rightarrow$ products:

$$-\frac{dA}{dt} = kA$$

$$\frac{A}{A_0} = \begin{cases} \exp[-k(t-t_0)], & \alpha = 1\\ \left[\frac{1}{1+kA_0^{\alpha-1}(t-t_0)}\right]^{1/(\alpha-1)}, & \alpha \neq 1 \end{cases}$$

2. $A + B \rightarrow products$:

$$-\frac{dA}{dt} = kAB = kA(A + B_0 - A_0)$$

k(t t_0) = $\frac{1}{B_0 - A_0} \ln \frac{A_0(A + B_0 - A_0)}{AB_0}$

3. Reversible reaction A $\xrightarrow{k_1}_{k_2}$ B:

$$-\frac{dA}{dt} = k_1 A - k_2 (A_0 + B_0 - A) = (k_1 + k_2) A - k_2 (A_0 + B_0)$$
$$(k_{1+} k_2)(t - t_0) = \ln \frac{k_1 A_0 - k_2 B_0}{(k_1 + k_2) A - k_2 (A_0 + B_0)^{-1}}$$

4. Reversible reaction, second order, $A + B = \frac{k_1}{k_2} R + S$

$$\begin{aligned} \frac{\partial A}{\partial t} &= k_1 A B - k_2 R S = k_1 A (A + B_0 - A_0) \\ &- k_2 (A_0 + R_0 - A) (A_0 + S_0 - A) \\ &= \alpha A^2 + \beta A - \gamma \\ \alpha &= k_1 - k_2 \\ \beta &= k_1 (B_0 - A_0) + k_2 (2A_0 + R_0 + S_0) \\ \gamma &= k_2 (A_0 + R_0) (A_0 + S_0) \\ q &= \sqrt{\beta^2 + 4\alpha\gamma} \\ k(t - t_0) &= \begin{cases} \frac{2\alpha A_0 + \beta}{2\alpha A + \beta}, & q = 0 \\ \frac{1}{q} \ln \left[\left(\frac{2\alpha A_0 + \beta - q}{2\alpha A + \beta + q} \right) \left(\frac{2\alpha A + \beta + q}{2\alpha A + \beta - q} \right) \right], & q \neq 0 \end{cases}$$

5. The reaction $v_a A + v_b B \rightarrow v_r R + v_s S$ between idea/ gases at constant

T and P

$$\frac{dn_{a} \quad Kn_{a}}{dt \quad V^{\alpha-1}}$$

$$V = n_{t} \frac{RT}{P} = \left[n_{t0} + \frac{\Delta v}{v_{a}} (n_{e0} - n_{e}) \right] \frac{RT}{P}$$

$$k(t - t_{0}) = \begin{cases} \int_{n_{a}}^{n_{e0}} \frac{V^{\alpha-1}}{n_{a}^{\alpha}} dn_{e}, \text{ in general} \\ \frac{RT}{P} \left[n_{b0} + \frac{\Delta v}{v_{a}} \left(\frac{1}{n_{a}} - \frac{1}{n_{e0}} \right) \\ - \frac{\Delta v}{v_{a}} \ln \left(\frac{n_{e0}}{n_{a}} \right) \right], \text{ when } \alpha = 2$$

6. Equations readily solvable by Laplace transforms. For example:

$$A \xleftarrow{n}{k_3} B \xrightarrow{k_2} C$$

Rate equations are

$$-\frac{dA}{dt} = k_1 A - k_2 B$$
$$-\frac{dB}{dt} = -k_1 A + (k_2 + k_3) B$$
$$-\frac{dC}{dt} = -k_2 B$$

Laplace transformations are made and rearranged to

$$(s + k_1)\bar{A} + k_3\bar{B} = A_0 -k_1\bar{A} + (s + k_2 + k_3)\bar{B} = B_0 -k_2\bar{B} + s\bar{C} = C_0$$

These linear equations are solved for the transforms as

$$\begin{split} D &= s^2 + (k_1 + k_2 + k_3)s + k_1k_2 \\ \mathrm{A} &= [A_0s + (k_2 + k_3)A_0 + K_3B_0]/D \\ \bar{B} &= [B_0s + k_1(A_0 + B_0)]/D \\ \bar{C} &= (k_2\bar{B} + C_0)/s \end{split}$$

Inversion of the transforms can be made to find the concentrations *A*, *B*, and C as functions of the time *t*. Many such examples are solved by Rodiguin and Rodiguina (*Consecutive Chemical Reactions*, Van Nostrand, New York, 1964).

paramount. Many different modes have been and are being employed, a few of which are illustrated in Section 6. The design of such equipment is based on material and energy balances that incorporate rates and heats of reaction together with heat transfer coefficients. Solution of these balances relates the time, composition, temperature, and rate of heat transfer. Such balances are presented in Tables 17.4-17.7 for four processes:

- 1. Nonflow reactors.
- 2. Plug flow reactors.
- 3. Continuous stirred tanks.
- 4. Flow reactor packed with solid catalyst.

The data needed are the rate equation, energy of activation, heat of reaction, densities, heat capacities, thermal conductivity, diffusivity, heat transfer coefficients, and usually the stoichiometry of the process. Simplified numerical examples are given for some of these cases. Item 4 requires the solution of a system of partial differential equations that cannot be made understandable in concise form, but some suggestions as to the procedure are made.

17.4. NONIDEAL FLOW PATTERNS

The CSTR with complete mixing and the PFR with no axial mixing are limiting behaviors that can be only approached in practice. Residence time distributions in real reactors can be found with tracer tests.

RESIDENCE TIME DISTRIBUTION

In the most useful form the test consists of a momentary injection of a known amount of inert tracer at the inlet of the operating vessel and monitoring of its concentration at the outlet. The data are used most conveniently in reduced form, as $E = C/\bar{C}_0$ in terms of $t_r = t/\bar{t}$, where

C = concentration of tracer at the outlet,

 \bar{C}_0 = initial average concentration of tracer in the vessel,

 $i = V_r / V'$ = average residence time.

The plotted data usually are somewhat skewed bell-shapes. Some

TABLE 17.4. Material and Energy Balances of a Nonflow Reaction

Rate equations:

$$r_{\sigma} = -\frac{1}{V_{r}} \frac{dn_{\sigma}}{d\theta} = kC_{\sigma}^{\alpha} = k \left(\frac{n_{\sigma}}{V_{r}}\right)^{\alpha}$$
(1)
$$k = \exp(a' - b'/T)$$
(2)

Heat of reaction:

$$\Delta H_r = \Delta H_{r298} + \int_{298}^{T} AC, \quad dT$$
(3)

Rate of heat transfer:

$$Q' = UA(T_s - T) \tag{4}$$

(the simplest case is when UA and T_s are constant) Enthalpy balance:

$$\frac{dT}{dn_{g}} = \frac{1}{\rho V_{r} \tilde{C}_{p}} \left[\Delta H_{r} + \frac{UA(T_{g} - T)}{V_{r} k(n_{g}/V_{r})^{\alpha}} \right]$$

$$\frac{dT}{dC_{g}} = \frac{1}{\rho \tilde{C}_{p}} \left[\Delta H_{r} + \frac{UA(T_{g} - T)}{V_{r} k C_{g}^{\alpha}} \right]$$

$$T = T_{0} \quad \text{when} \quad C_{g} = C_{g0}$$

$$\tilde{C}_{p} = \frac{1}{\rho V_{r}} \sum n_{i} C_{pi}$$

$$(5)$$

Solve Eq. (6) to find $T = f(C_a)$; combine Eqs. (1) and (2) and integrate as

$$\theta = \int_{C_{\theta}}^{C_{\theta}} \frac{1}{C_{\theta}^{\alpha} \exp[a' - b'/f(C_{\theta})]} dC_{\theta}$$
(9)

Temperature and time as a function of composition are shown for two values of UA/V_r for a particular case represented by

$$\frac{dT}{dC_{g} \cdot 5} \frac{1}{5} \left[-5000 - 5T + \frac{UA(300 - T)}{V_{r}kC_{g}^{2}} \right]$$

$$k = \exp(16 - 5000/T)$$

$$T_{0} = 350$$

$$C_{g0} = 1$$



TABLE 17.5. Material and Energy Balance of a CSTR

The sketch identifies the nomenclature Mean residence time:

$$i = V_r / V' \tag{1}$$

Temperature dependence:

$$= \exp(a' - b'/T) \tag{2}$$

Rate equation:

k

$$r_{a} = kC_{a}^{\alpha} = kC_{a0}^{\alpha}(1-x)^{\alpha}, x = (C_{a0} - C_{a})/C_{a0}$$
(3)

Material balance:

$$\begin{aligned} C_{s0} &= C_s + k\bar{t}C_s \end{aligned} \tag{4} \\ x &= k\bar{t}C_{s0}^{\alpha-1}(1-x)^{\alpha} \end{aligned} \tag{5}$$



Enthalpy balance:

$$\sum_{i} n_{i0}^{\prime} H_{i} - \sum_{i} n_{i0}^{\prime} H_{i0} = 0^{\prime} - \Delta H_{r} (n_{a0}^{\prime} - n_{a}^{\prime})$$
(6)

$$H_i = \int_{298} C_{\rho i} \, dT \tag{7}$$

$$\Delta H_r = \Delta H_{r298} + \int_{298}^{r} \mathbb{AC}, \quad dT$$
(8)

For the reaction $aA + bB \rightarrow rR + sS$,

$$\Delta C_{p} = rC_{pr} + sC_{ps} \quad aC_{ps} = bC_{pb} \tag{9}$$

When the heat capacities are equal and constant, the heat balance is

$$\tilde{C}_{\rho}\rho V'(T-T_0) = Q' - \Delta H_{r^{298}}V'(C_{a0} - C_a)$$
⁽¹⁰⁾

Example:

K = exp(16-5500/7)								
$C_{a0} = 5 \text{ gm}$	nol/L								
V' = 2000	L/hr								
$\Delta H_r = -5 \text{ kcal/g mol}$									
$\rho C_p = 0.9 \text{ km}$	cal/(L)(K)								
$\alpha = 2$									
$T_0 = 350$									
	x =	= 0.90	x =	0.95					
_ T	Τ_	Q′_	Т	<u>Q'</u>					
1	419.5	80	471.3	171					
2	398.5	42	444.9	123					
2 3	398. 5 387. 1	42 22	444. 9 430. 8	123 98					
2 3 4	398.5 387.1 379.4	42 22 8	444. 9 430. 8 421. 3	123 98 81					
2 3 4 5	398.5 387.1 379.4 373.7	42 22 8 2	444. 9 430. 8 421. 3 414. 2	123 98 81 68					
2 3 4 5 7;	398. 5 387. 1 379. 4 373. 7 369. 1	42 22 8 2 - 11	444. 9 430. 8 421. 3 414. 2 408. 6	123 98 81 68 58					
2 3 4 5 7; 8	398.5 387.1 379.4 373.7 369.1 365.3 362.1 -2	42 22 8 2 - 11 3 - 17	444. 9 430. 8 421. 3 414. 2 408. 6 404. 0	123 98 81 68 58 50					
2 3 4 5 7; 8	398.5 387.1 379.4 373.7 369.1 365.3 362.1 -2	42 22 8 2 - 11 13 - 17	444. 9 430. 8 421. 3 414. 2 408. 6 404. 0 400. 0	123 98 81 68 58 50 43					
2 3 4 5 7; 8 9	398. 5 387. 1 379. 4 373. 7 369. 1 365. 3 362. 1 - 2 359. 3	42 22 8 - 2 - 11 3 - 17 - 28	444. 9 430. 8 421. 3 414. 2 408. 6 404. 0 400. 0 396. 6	123 98 81 68 58 50 43 36					

(continued)
TABLE 17.5—(continued)

Eqs. (2) and (5) combine to

$$T = \frac{5500}{16 - \ln[x/5t(1-x)^2]}$$

and Eq. (10) becomes

 $Q' = 2[0.9(T - 350) \quad 25x], Mcal/hr$

The temperature and the rate of heat input ${\cal Q}'$ are tabulated as functions of the residence time for conversions of 90 and 95%

TABLE 17.6. Material and Energy Balances of a Plug Flow Reactor (PFR)

The balances are made over a differential volume dV_r of the reactor Rate equation:

$$dV_r = \frac{-dn'_a}{r_a}$$
(1)
= $-\frac{1}{k} \left(\frac{V'}{n'_a}\right)^{\alpha} dn'_a$ (2)

$$= -\exp(-a' + b'/T) \left(\frac{n'_{t}RT}{Pn'_{a}}\right)^{\alpha} dn'_{a}$$
(3)

Enthalpy balance:

AH, = AH,... +
$$\int_{298}^{T}$$
 AC, dT (4)

$$dQ = U(T_s - T) dA_p = \frac{4U}{D}(T_s - T) dV_r$$
$$= -\frac{4U(T_s - T)}{Dr_s} dn'_a$$
(5)

$$dQ + AH, dn'_{a} = \sum n_{i} dH_{i} = \sum n_{i}C_{pi} dT$$
(6)

$$\frac{dT}{dn'_{a}} = \frac{AH, -4U(T_{s} - T)/Dr_{\rho}}{\sum n_{i}c_{\rho i}} = f(T, T_{s}, n'_{a})$$
(7)



At constant T_{sr} Eq. (7) may be integrated numerically to yield the temperature as a function of the number of mols

 $T = \phi(n_a)$

actual data are shown in Figure 17.1 together with lines for ideal CSTR and PFR. Such shapes often are represented approximately by the Erlang statistical distribution which also is the result for an n-stage stirred tank battery,

$$E(t_r) = \frac{C}{\bar{C}_0} = \frac{n^n t_r^{n-1}}{(n-1)!} \exp(-nt_r),$$
(17.11)

where n is the characterizing parameter; when n is not integral, (n - 1)! is replaced by the gamma function F(n). C_0 is the initial average concentration. The variance,

$$\sigma^{2} = \frac{\sigma}{I_{0}} E(t_{r} - 1)^{2} dt_{r} = 1/n$$
(17.12)

of this distribution is a convenient single parameter characterization of the spread of residence times. This quantity also is related to the **Peclet** number, $Pe = uL/D_e$, by

$$\sigma^{2} = 2/\text{Pe} - [1 - \exp(-\text{Pe})]/\text{Pe}^{2}, \qquad (17.13)$$

where

u = linear velocity in the axial direction,

L = distance in the axial direction,

 D_e = axial eddy diffusivity or dispersion coefficient.

Then	the	reactor	volume	is	found	by	integration
,	V. =	п' _{а0}			1	.,	dn'

$$r' = \lim_{n_{a}} \exp[a' - b'/\phi(n_{a}')] [Pn_{a}'/n_{t}' R\phi(n_{a}')]^{\alpha} - dn_{a}'$$
(9)

(10)

Adiabatic process:

(8)



The balance around one end of the reactor is

$$\sum n_{i0}H_{i0} - \sum H_{r0}(n'_{a0} - n'_{a}) = \sum n_{i}H_{i} = \sum n_{i}\int C_{pi} dT$$
(11)

With reference temperature at T_{0} , enthalpies $H_{i0} = 0$

$$\Delta H_{r0} = \Delta H_{r298} + \int_{298}^{\tau_0} \Delta C_{\rho} \, dT \tag{12}$$

Substituting Eq. (12) into Eq. (10)

$$\left[-\Delta H_{r298} + \int_{298}^{\tau_0} \Delta C_p \, dT\right] (n'_{a0} - n'_{a}) = \sum n_i \int_{\tau_0}^{\tau} C_{pi} \, dT \tag{13}$$

Adiabatic process with AC, = 0 and with constant heat capacities

$$T = T_0 - \frac{\Delta H_{r296}(n'_{a0} - n'_{a})}{\sum n_i C_{pi}}$$
(14)

This expression is substituted instead of Eq. (8) to find the volume with Eq. (9)

TABLE 17.7. Material and Energy Balances of a Packed Bed Reactor

Diffusivity and thermal conductivity are taken appreciable only in the radial direction

Material balance equation:

$$\frac{\partial x}{\partial z} - \frac{D}{u} \left(\frac{\partial^2 x}{\partial r^2} + \frac{1}{r} \frac{\partial x}{\partial r} \right) - \frac{\rho}{u_0 C_0} r_c = 0 \tag{1}$$

Energy balance equation:

$$\frac{\partial T}{\partial z} - \frac{k}{GC_{\rho}} \left(\frac{\partial^2 T}{\partial r^2} + \frac{1}{r} \frac{\partial T}{\partial r} \right) + \frac{\Delta H_r \rho}{GC_{\rho}} r_c = 0$$
⁽²⁾

At the inlet:

$$x(0, r) = x_0$$

 $T(0, r) = T_0$
(3)
 $r R = (4)$

G

u_

At the center:

At the wall:

$$r = R, \quad \frac{\partial x}{\partial r} = 0 \tag{6}$$

$$\frac{\partial T}{\partial r} = \frac{U}{k} (T' - T) \tag{7}$$

When the temperature T' of the heat transfer medium is not constant, another enthalpy balance must be formulated to relate T' with the process temperature T.

A numerical solution of these equations may be obtained in terms of finite difference equivalents, taking m radial increments and n axial ones. With the following equivalents for the derivatives, the solution may be carried out by direct iteration:

$$r = m(\Delta r)$$

$$z = n(\Delta z)$$
(8)

$$\frac{\partial T}{\partial z} = \frac{T_{m,n+1} - T_{m,n}}{Az}$$
(9)

$$\frac{\partial T}{\partial r} \overline{\int_{r}^{m+1,n} \frac{-T_{m,n}}{Ar}}$$
(10)

$$\frac{\partial^2 T}{\partial r^2} = \frac{T_{m+1,n} - 2T_{m,n} + T_{m-1,n}}{(\Delta r)^2}$$
(11)

Expressions for the x-derivatives are of the same form:

 r_c = rate of reaction, a function of g and TG= mass flow rate, mass/(time)(superficial cross section) U = linear velocity D = diffusivity

k= thermal conductivity

At large values of Pe, the ratio Pe/n approaches 2. The superficial Peclet number in packed beds,

$$Pe = u_0 d_p / D_e$$

is very roughly correlated (Wen and Fan, Models for Flow Systems



Figure 17.1. Residence time distributions of some commercial and pilot fixed bed reactors. The variance, the equivalent number of CSTR stages, and the Peclet number are given for each.

and Chemical Reactors, Dekker, NY, 1975) in terms of the dimensionless groups Re = $u_0 d_p \rho / \mu$ and Sc = $\mu / \rho D_m$, where

 d_p = particle diameter, D_m = molecular diffusivity, ε = fraction voids in the bed.

The correlations are

$$\varepsilon Pe = 0.20 + 0.011 Re^{0.48}$$
, for liquids, standard deviation 46%, (17.14)

$$\frac{1}{Pe} = \frac{0.3}{Re Sc} + \frac{0.5}{1 + 3.8/Re Sc}, \text{ for gases.}$$
(17.15)

There are no direct correlations of the variance (or the corresponding parameter n) in terms of the geometry and operating conditions of a vessel. For this reason the RTD is not yet a design tool, but it does have value as a diagnostic tool for the performance of existing equipment on which tracer tests can be made. **RTDs** obtained from tracer tests or perhaps estimated from dispersion coefficient data or correlations sometimes are applicable to the prediction of the limits between which a chemical conversion can take place in the vessel.

CONVERSION IN SEGREGATED AND MAXIMUM MIXED FLOWS

In some important cases, limiting models for chemical conversion are the segregated flow model represented by the equation

$$\bar{C}/C_0 = 1 - \mathbf{x} = \prod_{I_0}^{\infty} (C/C_0)_{\text{batch}} E(t_r) dt_r = \int_{t_0}^{\infty} (C/C_0)_{\text{batch}} E(t) dt$$
(17.16)

and the maximum-mixedness model represented by Zwietering's equation. For a rate equation $r_c = kC^{\alpha}$ this equation is

$$\frac{dx}{dt_r} - kC_0^{\alpha - 1} x^{\alpha} + \frac{E(t_r)}{1 - \int_0^{t_r} E(t_r) dt_r} x = 0, \qquad (17.17)$$

with the boundary condition

$$dx/dt_r = 0 \quad \text{when } t_r \to \infty, \tag{17.18}$$

which is used to find the starting value x_{∞} from

$$kC_{\infty}^{m-1}x_{\infty} - \frac{E(t_r)}{1 - \int_0^{\infty} E(t_r) dt.} x_{\infty} = 0.$$
 (17.19)

Numerical integration of the equation is sufficiently accurate by starting at $(\mathbf{x}_{\infty}, \mathbf{t}_r \simeq 4)$ and proceeding to $\mathbf{t}_r = 0$ at which time the value of x is the conversion in the reactor with residence time distribution $E(\mathbf{t}_r)$.

With a given RTD the two models may correspond to upper and lower limits of conversion or reactor sizes for simple rate equations; thus

	Conversion	Limit
Reaction Orde	r Segrated	Max-Mix
More than 1	upper	lower
Less than 1	lower	upper
Complex	?	?

Relative sizes of reactors based on the two models are given in Figure 17.2 for second- and half-order reactions at several conversions. For first order reactions the ratio is unity. At small values of the parameter n and high conversions, the spread in reactor sizes is very large. In many packed bed operations, however, with proper initial distribution and redistribution the value of the parameter n is of the order of 20 or so, and the corresponding spread in reactor sizes is modest near conversions of about 90%. In such cases the larger predicted vessel size can be selected without undue economic hardship.

The data also can be rearranged to show the conversion limits for a reactor of a given size.

When the rate equation is complex, the values predicted by the two models are not necessarily limiting. Complexities can arise from multiple reactions, variation of density or pressure or temperature, incomplete mixing of feed streams, **minimax** rate behavior as in autocatalytic processes, and possibly other behaviors. Sensitivity of the reaction to the mixing pattern can be established in such cases, but the nature of the conversion limits will not be ascertained. Some other, possibly more realistic models will have to be devised to represent the reaction behavior. The literature has many examples of models but not really any correlations (Naumann and **Buffham**, 1983; Wen and Fan; Westerterp et al., 1984).



Figure 17.2. Relative volumes of maximum-mixed and segregated flow reactors with the same **RTDs** identified by $n = 1/\sigma^2$, as a function of conversion for second- and half-order reactions. For first-order reactions the ratio is unity throughout.

CONVERSION IN SEGREGATED FLOW AND CSTR BATTERIES

The mixing pattern in an n-stage CSTR battery is intermediate between segregated and maximum mixed flow and is characterized by residence time distribution with variance $\sigma^2 = 1/n$. Conversion in the CSTR battery is found by solving *n* successive equations

$$\frac{C_{j-1}}{C_0} = \frac{C_j}{C_0} + \frac{kt}{n} C_0^{\alpha-1} \left(\frac{C_j}{C_0}\right)^{\alpha} \quad \text{for } j = 1-n$$
(1720)

for $C_n/C_0 = 1 - x$. The ratio of required volumes of CSTR batteries and segregated flow reactors is represented by Figure 17.3 for several values of *n* over a range of conversions for a second order reaction. Comparison with the maximum mixed/segregated flow relation of Figure 17.2 shows a distinct difference between the two sets of ratios.

DISPERSION MODEL

Although it also is subject to the limitations of a single characterizing parameter which is not well correlated, the **Peclet** number, the dispersion model predicts conversions or residence times unambiguously. For a reaction with rate equation $r_c = kC^{\alpha}$, this model is represented by the differential equation

$$\frac{1}{\text{Pe}}\frac{d^2x}{dz^2} - \frac{dx}{dz} + k\bar{t}C_0^{\alpha-1}(1-x)^{\alpha} = 0$$
(17.21)



Fire 17.3. Ratio of volumes of an n-stage CSTR battery and a pegregated flow reactor characterized by a residence time distribution with variance $\sigma^2 = 1/n$. Second-order reaction.

with the boundary conditions

at
$$z = 0$$
, $\left(1 - x + \frac{1}{\text{Pe}} \frac{dx}{dz}\right)_0 = 1$, (17.22)
at $z = 1$, $\frac{dx}{dt} = 0$, (17.23)

where

x = 1 - C/C, fractional conversion,

z = axial distance/length of reactor.

An analytical solution can be found only for a first-order reaction. The two-point boundary condition requires a special numerical procedure. Plots of solutions for first and second order reactions are shown in Figures 17.4 and 17.5.

LAMINAR AND RELATED FLOW PATTERNS

A tubular reactor model that may apply to viscous fluids such as polymers has a radial distribution of linear velocities represented by

$$u = (1 + 2/m)\bar{u}(1 - \beta^m), \qquad (17.24)$$

where $\beta = r/R$. When m = 2, the pattern is Poiseuille or laminar flow, and, when m is infinite, it is plug flow. The residence time along a streamline is

$$t = \bar{t}/(1+2/m)(1-\beta^m). \tag{17.25}$$

The average conversion over all the stream lines is

$$\frac{Cl}{C_0} = \frac{1}{\pi R^2} \int \left(\frac{c}{c}\right)_{\text{streamline}} d(\pi r^2) = 2 \int_0^1 \left(\frac{C}{C_0}\right)_{\text{streamline}} \beta \, d\beta.$$
(17.26)

For first-order reaction, for example

$$\frac{\bar{C}}{C_0} = 2 \int_0^1 \exp\left[\frac{-k\bar{t}}{(1+2/m)(1-\beta^m)}\right] \beta \, d\beta$$
(17.27)

and for second-order

$$\frac{C}{C_0} = 2 \int_0^1 \frac{1}{1 + kC_0 t/(1 + 2/m)(1 - \beta^m)} \beta \, d\beta.$$
(17.28)

These integrals must be evaluated numerically. Variation in residence time will contribute, for example, to the spread in molecular weight distribution of polymerizations.



Figure 17.4. Dispersion model. Conversion of first-order reaction as function of the **Peclet** number.



Figure 17.5. Second-order reaction with dispersion identified by the Peclet number, $Pe = uL/D_r$.

17.5. SELECTION OF CATALYSTS

A catalyst is a substance that increases a rate of reaction by participating chemically in intermediate stages of reaction and is liberated near the end in a chemically unchanged form. Over a period of time, however, permanent changes in the **catalyst**—deactivation-may occur. Inhibitors are substances that retard rates of reaction. Many catalysts have specific actions in that they influence only one reaction or group of definite reactions. An outstanding example is the living cell in which there are several hundred different catalysts, called enzymes, each one favoring a specific chemical process.

The mechanism of a catalyzed reaction-the sequence of reactions leading from the initial reactants to the final products-is changed from that of the uncatalyzed process and results in a lower overall energy of activation, thus permitting a reduction in the temperature at which the process can proceed favorably. The equilibrium condition is not changed since both forward and reverse rates are accelerated equally. For example, a good hydrogenation catalyst also is a suitable dehydrogenation accelerator; the most favorable temperature will be different for each process, of course.

A convenient classification is into homogeneous and heterogeneous catalysts. The former types often are metal complexes that are soluble in the reaction medium, but acids and bases likewise have a long known history of catalytic action. The specific action of a particular metal complex can be altered by varying the ligands or coordination number of the complex or the oxidation state of the central metal atom. Advantages of homogeneous catalysts are their specificity and low temperature and pressure requirements. Their main drawbacks are difficulty of recovery from the process fluid, often rapid degradation, and relatively high cost. Classic examples of homogeneous catalysis are the inversion of sugar with mineral acids, olefin alkylation with hydrofluoric acid, and the use of ammonia in the Solvay process and of nitrogen oxides in the Chamber process. A modern development is the synthesis of acetic acid from methanol and CO in the presence of homogeneous rhodium complexes.

The problem of separating the catalyst at the end of the operation can be eased in some cases by attaching the catalyst to a solid support, for instance, liquid phosphoric acid in the pores of a solid carrier for the vapor phase synthesis of cumene and the fairly wide application of enzymes that are attached (immobilized) by various means to solid polymers. Some metal ligands also are being combined with solid polymers.

HETEROGENEOUS CATALYSTS

By far the greatest tonnages of synthetic chemicals are manufactured in fluid phases with solid catalysts. Such materials are cheap, are easily separated from the reaction medium, and are adaptable to either flow or **nonflow** reactors. Their drawbacks are a lack of specificity and often high temperature and pressure requirements. The principal components of most heterogeneous catalysts are three in number:

- 1. A catalytically active substance or mixture.
- 2. A carrier of more or less large specific surface on which the catalyst proper is deposited as a thin film, either for economy when the catalyst is expensive or when the catalyst itself cannot be prepared with a suitable specific surface.
- Promoters, usually present in relatively small amount, which enhance the activity or retard degradation.

Some composite catalysts are designed to promote several reactions of a sequence leading to the final products. A basic catalyst often can be selected with general principles, but subsequent fine tuning of a commercially attractive design must be done in a pilot plant or sometimes on a plant scale.

Analogy to what is known to be effective in chemically similar problems usually provides a start for catalyst design, although a scientific basis for selection is being developed. This involves a study in detail of the main possible intermediate reactions that could occur and of the proton and electron receptivity of the catalyst and possible promoters, as well as reactant bond lengths and crystals lattice dimensions. Several designs are made from this fundamental point of view in the book of Trimm (1980). A thorough coverage of catalytic reactions and catalysts arranged according to the periodic table is underway in a series edited by Roiter (1968–date). Industrial catalyst practice is summarized by Thomas (1970) who names manufacturers of specific catalysts. Specific processes and general aspects of catalysis are covered in three books edited by Leach (1983-1985): In a chapter by Wagner, there are lists of 40 catalysts with the kinds of processes for which they are effective and of 49 catalytic processes with the homogeneous or heterogeneous catalysts that have been used with them. Many industrial processes are described by Satterfield (1980). Cracking, reforming, partial oxidation, hydrodesulfurization, and catalysis by transition-metal complexes are treated in detail by Gates et al. (1979) and the catalytic conversion of hydrocarbons by Pines (1981). The mechanisms and other aspects of organic catalysis are described in one of the volumes of the series edited by Bamford and Tipper (1978). A vast literature exists for enzyme processes; that technology is well reviewed in two articles in *Ullmann's Encyclopedia* (Biotechnologie, Enzyme) as well as by Bailey and Ollis (1986). In the present text, Table 17.1 identifies the catalyst used in most of the 100 processes listed.

Intermediate processes of catalyzed organic reactions may involve neutral free radicals $R^{\, }$, positive ions R^{+} , or negative ions R^{-} as short-lived reactants. A classification of catalysts and processes from the point of view of elementary reactions between reagents and catalysts is logically desirable but has not yet been worked out. However, there is a wealth of practice more or less completely documented, some proprietary but available at a price. The ensuing discussions are classified into kinds of catalysts and into kinds of processes.

KINDS OF CATALYSTS

To a certain extent, it is known what kinds of reactions are speeded up by certain classes of catalysts, but individual members of the same class may differ greatly in activity, selectivity, resistance to degradation, and cost. Even small differences in these properties can mean large sums of money on the commercial scale. Solid catalysts, the most usual kind, are not particularly specific or selective, so that there is a considerable crossing of lines in classifications between kinds of catalysts and kinds of reactions they favor. Nevertheless, leading relations can be brought out.

Strong acids are able to donate protons to a reactant and to take them back. Into this class fall the common acids, aluminum halides, and boron trifluoride. Also acid in nature are silica, alumina, aluminosilicates, metal sulfates and phosphates, and sulfonated ion exchange resins. The nature of the active sites on these kinds of solids still is not completely understood. The majority of reactions listed subsequently are catalytically influenced to some extent by acidic substances. Zeolites are dehydrated aluminosilicates with small pores of narrow size distribution, to which is due their highly selective catalytic action since only molecules small enough to enter the pores can react. In cracking operations they are diluted to 10–15% in silica-alumina to restrain their great activity; the composite catalyst still is very active but makes less carbon, makes lower amounts of C_3-C_4 products, and has a longer life. Their greater activity has led to the supplanting of fluidized bed crackers by riser-tube reactors. When zeolites are incorporated in reforming catalysts, they crack isoparaffins into straight chains that enter the pores and convert into higher octane substances.

Base catalysis is most effective with alkali metals dispersed on solid supports or in the homogeneous form as aldoxides, amides, and so on. Small amounts of promoters may be added to form organoalkali compounds that really have the catalytic power. Basic ion exchange resins also are useful. Some base-catalyzed processes are isomerization and oligomerization of olefins, reaction of olefins with aromatics, and hydrogenation of polynuclear aromatics.

Metal oxides, sulfides, and hydrides form a transition between acid-base and metal catalysts. They catalyze hydrogenationdehydrogenation as well as many of the reactions catalyzed by acids such as cracking and isomerization. Their oxidation activity is related to the possibility of two valence states which allow oxygen to be released and reabsorbed alternately. Common examples are oxides of cobalt, iron, zinc, and chromium; and hydrides of precious metals which can release hydrogen readily. Sulfide catalysts are more resistant than metallic catalysts to formation of coke deposits and to poisoning by sulfur compounds; their main application is to hydrodesulfurization.

Metals and alloys. The principal industrial metallic catalysts are found in periodic group VIII which are transition elements with almost completed 3d, 4d, and 5d electron orbits. According to one theory, electrons from adsorbed molecules can fill the vacancies in the incomplete shells and thus make a chemical bond. What happens subsequently will depend on the operating conditions. Platinum, palladium, and nickel, for example, form both hydrides and oxides; they are effective in hydrogenation (vegetable oils, for instance) and oxidation (ammonia or sulfur dioxide, for instance). Alloys do not always have catalytic properties intermediate between those of the pure metals since the surface condition may be different from the bulk and the activity is a property of the surface. Addition of small amounts of rhenium to Pt/Al_2O_3 results in a smaller decline of activity with higher temperature and slower deactivation rate. The mechanism of catalysis by alloys is in many instances still controversial.

Transition-metal organometallic catalysts in solution are effective for hydrogenation at much lower temperatures than metals such as platinum. They are used for the reactions of carbon monoxide with olefins (hydroformylation) and for some **oligomeri**zations. The problem of separating the catalyst from solution sometimes is avoided by anchoring or immobilizing the catalyst on a polymer support containing pendant phosphine groups and in other ways.

KINDS OF CATALYZED ORGANIC REACTIONS

A fundamental classification of organic reactions is possible on the basis of the kinds of bonds that are formed and destroyed and the natures of eliminations, substitutions, and additions of groups. Here a more pragmatic list of 20 commercially important individual kinds or classes of reactions will be discussed.

- 1. Alkylations, for example, of olefins with aromatics or isoparaffins, are catalyzed by sulfuric acid, hydrofluoric acid, BF,, and AlCl₃.
- Condensations of aldehydes and ketones are catalyzed homogeneously by acids and bases, but solid bases are preferred, such as anion exchange resins and alkali or alkaline earth hydroxides or phosphates.
- 3. Cracking, a rupturing of carbon-carbon bonds, for example, of gas oils to gasoline, is favored by silica-alumina, zeolites, and acid types generally.
- 4. Dehydration and dehydrogenation combined utilizes dehydration agents combined with mild dehydrogenation agents. Included in this class of catalysts are phosphoric acid, silicamagnesia, silica-alumina, alumina derived from aluminum chloride, and various metal oxides.
- 5. Esterification and etherification may be accomplished by catalysis with mineral acids of BF,; the reaction of isobutylene with methanol to make MTBE is catalyzed by a sulfonated ion exchange resin.
- 6. Fischer-Tropsch oligomerization of CO + hydrogen to make hydrocarbons and oxygenated compounds. Iron promoted by potassium is favored, but the original catalyst was cobalt which formed a carbonyl in process.
- 7. Halogenation and dehalogenation are catalyzed by substances that exist in more than one valence state and are able to accept and donate halogens freely. Silver and copper halides are used for gas-phase reactions, and ferric chloride commonly for liquid phase. Hydrochlorination (the absorption of HCl) is promoted

by **BiCl₃** or **SbCl₃** and hydrofluorination by sodium fluoride or **chromia** catalysts that fluoride under reaction conditions. Mercuric chloride promotes addition of HCI to acetylene to make vinyl chloride.

- **8.** Hydration and dehydration employ catalysts that have a strong affinity for water. Alumina is the principal catalyst, but also used are aluminosilicates, metal salts, and phosphoric acid or its metal salts on carriers and cation exchange resins.
- 9. Hydrocracking is catalyzed by substances that promote cracking and hydrogenation together. Nickel and tungsten sulfides on acid supports and zeolites loaded with palladium are used commercially.
- 10. Hydrodealkylation, for example, of toluene to benzene, is promoted by chromia-alumina with a low sodium content.
- 11. Hydrodesulfurization uses sulfided cobalt/molybdena/alumina, or alternately with nickel and tungsten substituted for Co and Mo. .
- **12.** Hydroformylation, or the **OXO** process, is the reaction of olefins with CO and hydrogen to make aldehydes. The catalyst base is cobalt naphthenate which transforms to cobalt hydrocarbonyl in place. A rhodium complex that is more stable and functions at a lower temperature also is used.
- 13. Hydrogenation and dehydrogenation employ catalysts that form unstable surface hydrides. Transition-group and bordering metals such as Ni, Fe, Co, and Pt are suitable, as well as transition group oxides or sulfides. This class of reactions includes the important examples of ammonia and methanol syntheses, the Fischer-Tropsch and **OXO** and synthol processes and the production of alcohols, aldehydes, ketones, **amines**, and edible oils.
- 14. Hydrolysis of esters is speeded up by both acids and bases. Soluble. alkylaryl sulfonic acids or sulfonated ion exchange resins are satisfactory.
- 15. Isomerization is promoted by either acids or bases. Higher alkylbenzenes are isomerized in the presence of AlCl₃/HCl or BF₃/HF; olefins with most mineral acids, acid salts, and silica alumina; saturated hydrocarbons with AlCl₃ or AlBr₃ promoted by 0.1% of olefins.
- 16. Metathesis is the rupture and reformation of carbon-arbon bonds, for example of propylene into ethylene plus butene. Catalysts are oxides, carbonyls or sulfides of Mo, W, or rhenium.
- 17. Oxidation catalysts are either metals that chemisorb oxygen readily such as platinum or silver, or transition metal oxides that are able to give and take oxygen by reason of their having several possible oxidation states. Ethylene oxide is formed with silver, ammonia is oxidized with platinum, and silver or copper in the form of metal screens catalyze the oxidation of methanol to formaldehyde.
- Polymerization of olefins such as styrene is promoted by acid or base catalysts or sodium; polyethylene is made with homogeneous peroxides.
- 19. Reforming is the conversion primarily of naphthenes and alkanes to aromatics, but other chemical reactions also occur under commercial conditions. Platinum or platinum/rhenium are the hydrogenation-dehydrogenation component of the catalyst and alumina is the acid component responsible for skeletal rearrangements.
- 20. Steam reforming is the reaction of steam with hydrocarbons to make town gas or hydrogen. For town gas a representative catalyst composition contains 13 wt % Ni, 12.1% U, and 0.3 wt % K; it is particularly resistant to poisoning by sulfur. To make hydrogen, the catalyst contains oxides of Ni, Ca, Si, Al, Mg, and K. Specific formulations are given by Satterfield (1980).

PHYSICAL CHARACTERISTICS OF SOLID CATALYSTS

Although a few very active solid catalysts are used as fine wire mesh or other finely divided form, catalysts are mostly porous bodies whose total surface is measured in m^2/g . These and other data of some commercial catalysts are shown in Table 17.8. The physical characteristics of major importance are as follows.

- **1.** Particle *size*. In gas fluidized beds the particle diameters average less than 0.1 mm; smaller sizes impose too severe loading on entrainment recovery equipment. In slurry beds the particles can be about 1 mm dia. In fixed beds the range is 2-5 mm dia. The competing factors are that the pressure drop increases with diminishing diameter and the accessibility of the internal surface decreases with increasing diameter. With poorly thermally conducting materials, severe temperature gradients or peaks arise with large particles that may lead to poor control of the reaction and the development of undesirable side reactions like carbonization.
- **2.** Specific surface. Solid spheres of 0.1 mm dia have a specific surface of $0.06 \text{ m}^2/\text{mL}$ and an activated alumina one of about $600 \text{ m}^2/\text{mL}$. Other considerations aside, a large surface is desirable because the rate of reaction is proportional to the amount of accessible surface. Large specific surfaces are associated with pores of small diameters and are substantially all internal surface.
- **3.** *Pore diameters and their distribution.* Small pores limit accessibility of internal surface because of increased resistance to diffusion of reactants inwards. Diffusion of products outward also is slowed down and may result in degradation of those products. When the catalyst is expensive, the inaccessible internal surface is a liability. A more or less uniform pore diameter is desirable, but the distribution usually is statistical and only molecular sieves have nearly uniform pores. Those catalyst granules that are extrudates of compacted masses of smaller particles have bimodal pore size distribution, between the particles and within them. Clearly a compromise between large specific surface and its accessibility as measured by pore diameter is required in some situations.
- **4.** *Effective diffusivity*. Resistance to diffusion in a catalyst pore is due to collisions with other molecules and with the walls of the pore. The corresponding diffusivities are called bulk diffusivity and Knudsen diffusivity D_{K} . Many data and correlations of the former type exist; the latter is calculable from the following formula (Satterfield, 1970, p. 42):

$$D_K = \frac{19,400\theta^2}{S_g \rho_p} \left(\frac{T}{M}\right)^{1/2},$$

where

 θ = fraction porosity, S_g = specific surface per unit mass, ρ_p = density, T = temperature (K) , M = molecular weight.

This equation applies to uniform cylindrical pores whose length equals the thickness of the catalyst through which the diffusion takes place. The actual diffusivity in common porous catalysts usually is intermediate between bulk and Knudsen. Moreover, it depends on the pore size distribution and on the true length of

TABLE 17.8. Physical Properties of Some Commercial Catalysts and Carriers'

Desig- nation	Nominal Size	Surface Area (m²/g)	Total Void Fraction	D ^b _{eff} x 10' (cm ² /sec)	Average Tortu- osity Factor τ, Parallel-Path Pore Model	r _e = 2 V _e /S _e (Å)	τ_n Based on Average Pore -Radius
T-126 T-1258 T-826 T-314 T-310	3/16 x 1/8 in.	197 302 232 142 154	0.384 0.478 0.389 0.488 0.410	29.3 33.1 37.7 20.0 16.6	$\begin{array}{c} 3.7 \pm 0.2 \\ 3.8 \pm 0.2 \\ 3.9 \pm 0.1 \\ 7.1 \bullet 0.9 \\ 3.8 + 0.1 \end{array}$	29 23.6 21.4 41.5 34.3	0.45 0.41 0.26 1.2 0.67
G-39 G-35 T-606 G-58	3/16 x 3/16 in.	190 6.4	0.354 0.354 0.115 0.389	17.5 18.2 27.7 87.0	$\begin{array}{c} 4.8 \ \pm \ 0.3 \\ 4.9 \ \pm \ 0.1 \\ 2.9 \ \pm \ 0.2 \\ 2.8 \ \pm \ 0.3 \end{array}$	22.4 543.	0.53
T-126 T-606 G-41 G-52	1/4 x 1/4 in.	165	$\begin{array}{c} 0.527 \\ 0.092 \\ 0.447 \\ 0.436 \end{array}$	38.8 0.71 21.9 27.4	$\begin{array}{c} 3.6 \ \pm \ 0.3 \\ 79 \ \pm \ 28 \\ 4.4 \ \pm \ 0.1 \\ 3.9 \ \pm \ 0.2 \end{array}$	49.0	0.79
G-56 BASF	1/2 x 1/2 in. 5 × 5 mm	42 87.3	0.304 0.500	8.1 11.8	11.1 ± 1.1 7.3 + 0.7	84. 41.	3.74 2.05
Harshaw Haldor Topsoe	1/4 x 1/4 in. 1/4 x 1/4 in.	44 143	0.489 0.433	13.3 15.8'	7.2 ± 0.1 2.8	91. 25.8	3.95 0.83

Catalyst	• Description					
T-126,	Activated y-alumina					
T-1258	Activated y-alumina					
T-826	3% CoO, 10% MoO ₃ , and 3% NiO on alumina					
T-314	About 8-10% Ni and Cr in the form of oxides on an activated alumina					
T-310	About IO-12% nickel as the oxide on an activated alumina					
T-606	Specially compounded refractory oxide support					
G-39	A cobalt-molybdenum catalyst, used for simultaneous					
	hydrodesulfurization of sulfur compounds and hydrogenation of olefins					
G-35	A cobalt-molybdenum catalyst supported on high - purity alumina, used for hydrodesulfurization of organic sulfur compounds					
G-41	A chromia-alumina catalyst, used for hydrodealkyla - tion and dehydrogenation reactions					
G-58	Palladium-on-alumina catalyst, for selective hydro- genation of acetylene in ethylene					
G-52	Approximately 33 wt % nickel on a refractory oxide support, prereduced. Used for oxygen removal from hydrogen and inert gas streams					
G-56	A nickel-base catalyst used for steam reforming of hydrocarbons					
BASF	A methanol synthesis catalyst, prereduced					
Harshaw	A methanol synthesis catalyst, prereduced					
Haldor Topsoe	A methanol synthesis catalyst, prereduced					

^a The measured effective diffusivities are those of hydrogen in nitrogen at room temperature and pressure except that of Haldor Topsoe which is of helium in nitrogen.

[Satterfield and Cadle, Ind. Eng. Chem. Process Design and Development, 7, 256 (1968)].

path. Two tortuosity factors are defined:

- τ_p = ratio of measured diffusivity to that calculated with the known pore size distribution and bulk diffusivity and the thickness of the catalyst mass.
- τ_m = ratio of measured diffusivity to that calculated from the Knudsen formula with a mean pore diameter.

The data of Table 17.8 exhibit a fairly narrow range of τ_p , an average of about 4, but there seems to be no pattern to τ_m , which is not surprising since the diffusions actually are intermediate between bulk and Knudsen in these cases. In order to be able to calculate the effective diffusivity, it is necessary to know the pore size distribution, the specific surface, the porosity, and bulk diffusivity in the reaction mixture under reaction conditions. Such a calculation is primarily of theoretical interest. Practically it is more useful to simply measure the diffusivity directly, or even better to measure the really pertinent property of catalyst effectiveness as defined next.

CATALYST EFFECTIVENESS

Catalyst effectiveness is a measure of the extent of utilization of internal surface; it is the ratio of a rate of reaction actually achieved with the catalyst particle to the rate that would prevail if all of the internal surface were exposed to the reactant concentration at the external surface of the particle. The rate equation accordingly is modified to

(17.30)
(17.3)

where η is the catalyst effectiveness and C_s is the concentration of the reactant at the external surface. For isothermal reactions, η

566 CHEMICAL REACTORS

always is less than unity, but very large values can develop for exothermic reactions in poorly conducting catalysts.

A great deal of attention has been devoted to this topic because of the interesting and often solvable mathematical problems that it presents. Results of such calculations for isothermal zero-, first-, and second-order reactions in uniform cylindrical pores are summarized in Figure 17.6. The abscissa is a modified Thiele modulus whose basic definition is

$$\boldsymbol{\phi} = \boldsymbol{R}/\boldsymbol{k}_{v} \boldsymbol{C}_{s}^{n-1}/\boldsymbol{D}_{\text{eff}},\tag{17.31}$$

where R is a linear dimension (the radius of a sphere, for example), k_v the specific rate on a volumetric basis, C_s the surface concentration, n the order of the reaction, and D_{eff} the effective diffusivity. For nonisothermal reactions, those with variable volume and with rate equations of the Langmuir-Hinshelwood or other complex types, additional parameters are involved. Although such calculations can be made, they still require measurements of effective diffusivity as well as a number of unverifiable assumptions. Accordingly in practical cases it is preferable to make direct measurements of catalyst effectiveness and to correlate them with operating parameters. The effectiveness is deduced by comparing conversion with the reference particle size with those with successively small particle sizes until the effect disappears. Two examples are presented to illustrate the variables that are taken into account and the magnitudes of the effects.

For synthesis of ammonia the effectiveness has been measured by Dyson and Simon [*Ind. Eng. Chem. Fundam. 7,605610* (1968)] and correlated by the equation

$$\eta = b_0 + b_1 T + b_2 x + b_3 T^2 + b_4 x^2 + b_5 T^3 + b_6 x^3, \qquad (17.32)$$

where T is in K, x is fractional conversion of nitrogen, and the b_i depend on pressure as given in this table:



Figure 17.6. Generalized chart of catalyst effectiveness for reactions of order n in particles with external surface A_n and volume V_p . The upper curve applies exactly to zero-order reaction in spheres, and the lower one closely for first- and second-order reactions in spheres.

The reference mixture has $H_2/N_2 = 3$ and contains 12.7% inert; other ratios had slightly different effectiveness. The particle diameters are 6–10 mm. Some calculations from this equation at 225 atm are:

Т	X	η
700	0. 25	0. 81
700	0.10	0.57
650	0.25	0.91

For oxidation of sulfur dioxide, measurements of effectiveness were made **by Kadlec**, Pour, and Regner [*Coll. Czech. Chem. Commun.* 33, 2388, 2526 (1968)] whose data are shown following. They are at atmospheric pressure. The initial content of SO, and the conversion have little effect on the result. Both increase in size of granule and temperature lower the effectiveness, although the effect of temperature is somewhat erratic.

Experimentally Determined Effectiveness Factors

			Conversion						
°C	% so,	0.4	0.5	0.6	0.7	0.8	0. 9		
Irregul	ar grain	shape,	fractio	on 5-6 m	m				
460	7	0.84	0.84	0.82	0.83	0.82	0.81		
480	7	0.60	0.62	0.62	0.62	0.60	0.60		
500	7	_	0.54	0.51	0.50	0.50	0.52		
520	7		0.35	0.35	0.35	0.38	0.38		
Cyl i ndr	rical gran	ules of	6 mm	di ameter	and	12 mm l	ength		
460	7	0.57	0.57	0.59	0.60	0.60	0.60		
	10	0.58	0.62	0.63	0.63	0.62	0.62		
480	7	0.53	0.54	0.56	0.57	0.56	0.57		
	10	0.44	0.45	0.45	0.46	0.45	0.47		
500	7	0.25	0.25	0.27	0.28	0.27	0.31		
	10	0.26	0.27	0.30	0.30	0.31	0.30		
520	7		0. 21	0.21	0. 22	0.22	0.23		
	10	—	0. 20	0.21	0. 21	0. 22	0.24		

The rate equations of both these processes are quite complex, and there is little likelihood that the effectiveness could be deduced mathematically from fundamental data as functions of temperature, pressure, conversion, and composition, which is the kind of information needed for practical purposes. Perhaps the only estimate that can be made safely is that, in the particle size range below 1 mm or so, the effectiveness probably is unity. The penetration of small pores by liquids is slight so that the catalysts used in liquid slurry systems are of the low specific surface type or even nonporous.

17.6. TYPES AND EXAMPLES OF REACTORS

Almost every kind of holding or contacting equipment has been used as a chemical reactor at some time, from mixing nozzles and centrifugal pumps to the most elaborate towers and tube assemblies. This section is devoted to the general characteristics of the main kinds of reactors, and also provides a gallery of selected examples of working reactors.

The most obvious distinctions are between **nonflow** (batch) and continuous operating modes and between the kinds of phases that are being contacted. A classification of appropriate kinds of reactors on the basis of these two sets of distinctions is in Figure 17.7.

When heterogeneous mixtures are involved, the conversion rate often is limited by the rate of interphase mass transfer, so that a large interfacial surface is desirable. Thus, solid reactants or catalysts are finely divided, and fluid contacting is forced with mechanical agitation or in packed or tray towers or in centrifugal pumps. The rapid transfer of reactants past heat transfer surfaces by agitation or pumping enhances also heat transfer and reduces harmful temperature gradients.

Batch processing is used primarily when the reaction time is long or the required daily production is small. The same batch equipment often is used to make a variety of products at different times. Otherwise, it is *not* possible to generalize as to the economical transition point from batch to continuous operation. One or more batch reactors together with appropriate surge tanks may be used to simulate continuous operation on a daily or longer basis.

STIRRED TANKS

Stirred tanks are the most common type of batch reactor. Typical proportions are shown on Figures 17.8 and 10.1, and modes of level control on Figure 3.6. Stirring is used to mix the ingredients initially, to maintain homogeneity during reaction, and to enhance heat transfer at a jacket wall or internal surfaces. The reactor of Figure 17.9(b) employs a pumparound for mixing of the tank contents and for heat transfer in an outside exchanger. Pumparound or recycle in general may be used to adapt other kinds of vessels to service as batch mode reactors; for example, any of the packed vessels of Figure 17.10(a)-(e). A pumparound tubular flow reactor is employed for the polymerization of ethylene on Figure 17.11(c); as the polymer is formed, it is bled off at a much lower rate than that of the recirculation, so that in a sense the action of this equipment approaches batch operation.

Some special industrial stirred reactors are illustrated in Figure 17.10: (b) is suitable for pasty materials, (c) for viscous materials, and the high recirculation rate of (d) is suited to intimate contacting of immiscible liquids such as hydrocarbons with aqueous solutions.

Many applications of stirred tank reactors are to continuous processing, either with single tanks or multiple arrangements as in Figures 17.9(c)-(d). Knowledge of the extent to which a stirred tank does approach complete mixing is essential to being able to predict its performance as a reactor. The other limiting case is that of plug flow, in which all nonreacting molecules have the same residence time. Deviations from the limiting cases of complete mixing (in a CISTR) and no axial mixing (in a PFR) are evaluated with residence time distributions (RTDs) based on analyses of tracer tests.

At present, RTD behavior has not been well correlated with operating or design factors, but the technique is of diagnostic value with existing equipment. CISTR (continuous ideal stirred tank reactor) behavior is approached when the mean residence time is 5-10 times the mixing time, which is in turn the length of time needed to achieve homogeneity of a mixture of several inputs. Often this is achieved by 50-200 revolutions of a properly designed stirrer. Although mixing times have been the subject of many studies in the literature (Westerterp et al., 1984, p. 254), no useful generalizations have been deduced. The mixing time depends on the geometry and the speed and power of the agitator. A propeller above and a turbine below on the same shaft, baffles attached to the wall of the tank, and possibly a draft tube around the shaft for effective recirculation of the contents constitute a basic design. However, no completely rational design of mixing equipment is possible at this time, so that in critical cases experts should be consulted. Chapter 10 also deals with this topic.

Power input per unit volume and impeller tip speeds are often used measures of the intensity of stirring, assuming correct proportions of the vessel and proper baffling. Appropriate ranges



Figure 17.7. Classification of reactors according to the mode of operation and the kinds of phases involved.



for some reaction conditions are as tabulated:

Operation	kW/m ³ /	Tip speed (m/sec)
Blending	0.05-0. 1	
Homogeneous reaction	0.1-0.3	2.5-3.3
Reaction with heat transfer	0.3-1 .0	3.5-5.0
Gas-liquid, liquid-liquid	1-2	5-6
Slurries	2-5	
[#] 1 kW/m ³ = 5.08 HP/1000	gal	

Heat transfer coefficients in stirred tank operations are discussed in Section 17.7.

For a given load and conversion, the total volume of a CSTR (continuous stirred tank reactor) battery decreases with the number of stages, sharply at first and then more slowly. When the reaction is first order, for example, r = kC, the ratio of total reactor volume V_r of n stages to the volumetric feed rate V'_0 is represented by

$$kV_r/V_0' = n[(C_0/C)^{1/n} - 1].$$
(17.33)

At conversions of 95 and 99%, some values from this equation are

n	1	2	3	4	5	10
kV,/V₀ at 95%	19	6.9	5.1	4.5	5.1	3.5
V/V at 99%	99	18.0	10.9	9.7	7.6	5.9

Since the cost of additional controls, agitators, and pumps can counterbalance the savings in volume, four or five tanks in a battery normally prove to be an optimum number, but a larger number of stages may be economical with a single shell design like Figure

Figure 17.8. Typical proportions of a stirred tank reactor with radial and axial impellers, four baffles, and a sparger feed inlet.



Figure 17.9. Stirred tank reactors, batch and continuous. (a) With agitator and internal heat transfer surface, batch or continuous. (b) With pumparound mixing and external heat transfer surface, batch or continuous. (c) Three-stage continuous stirred tank reactor battery. (d) Three-stage continuous stirred tank battery in a single shell.

17.9(d), particularly when the stages are much less efficient than ideal ones.

For some purposes it is adequate to assume that a battery of five or so **CSTRs** is a close enough approximation to a plug flow reactor. The tubular flow reactor is smaller and cheaper than any comparable tank battery, even a single shell arrangement. For a first order reaction the ratio of volumes of an n-stage CSTR and a PFR is represented by

$$(V_r)_{\rm CSTR} / (V_r)_{\rm PFR} = n[(C_0/C)^{1/n} - 1] / \ln(C_0/C).$$
(17.34)

For example, when n = 5 and conversion is 99%, the ratio is 1.64. For second-order and other-order reactions a numerical solution for the ratio is needed, one of which is represented by Figure 17.12. For a second order reaction the ratio is 1.51 at 99% conversion with five stages.

A further difference between CSTR batteries and PFRs is that of product distributions with complex reactions. In the simple case, $A+B \rightarrow C$ for example, a higher yield of intermediate product B is obtained in a PFR than in a single CSTR. It is not possible to generalize the results completely, so that the algebra of each individual reacting system must be worked out to find the best mode.

TUBULAR FLOW REACTORS

The ideal behavior of tubular flow reactors (TFR) is plug flow, in which all nonreacting molecules have equal residence times. Any backmixing that occurs is incidental, the result of natural turbulence or that induced by obstructions to **flow** by catalyst granules or tower packing or necessary internals of the vessels. The action of such obstructions can be two-edged, however, in that some local backmixing may occur, but on the whole a good approach to plug flow is developed because large scale turbulence is inhibited. Any required initial blending of reactants is accomplished in mixing nozzles or by in-line mixers such as those of Figures 10.13-10.14. As a result of chemical reaction, gradients of concentration and temperature are developed in the axial direction of **TFRs**.

TFRs may be of pipe diameters ranging from 1 to 15 cm or so, or they may be vessels of diameters measured in meters. Figure 17.13 is of a variety of vessel configurations. Single tube reactors more than **1000 m** long are used, in which case they are trombone-shaped as on Figures 17.14(f) and 17.15(c). The selection of diameter is a result of compromise between construction cost, pumping cost, and required heat transfer. In some cases it may be necessary to avoid the laminar **flow** region, which is below Reynolds numbers of 2300-4000 or so, if the **reaction** is complex and a spread of residence times is harmful.

When many tubes in parallel are needed, a shell-and-tube construction like that of heat exchangers is employed; the vessel then may be regarded as a heat exchanger in which a reaction occurs incidentally. Heat transfer to single tubes is accomplished with jackets in Figure 17.14(f) and in a fired heater in Figure 17.15(c). Some of the many designs of fired heaters that are suitable for pyrolysis and other high temperature reactions are illustrated on



Figure 17.10. Several modes of mixing in commercial tank reactors. (a) Steam-jacketed autoclave, 120 gal, 200 psig, 300°F (courtesy *Blaw-Knox* Co.). (b) Horizontal autoclave, 650 gal, 100 psig (courtesy *Blaw-Knox* Co.). (c) Ball-mill sulfonator [*Groggins*. Courtesy *McGraw-Hill*, New York]. (d) Horizontal heat-exchange reactor (courtesy Stratford Engineering Corp. patents issued and pending).

Figure 17.16. In the process for making phenol, monochlorbenzene, and aqueous caustic are reacted at 320° C and 200 atm in multipass tubes of 10 cm dia or so in a fired heater.

In general, the construction of **TFRs** is dictated by the need for accommodation of granular catalysts as well as for heat transfer. Some of the many possible arrangements are illustrated on Figure 17.13 and elsewhere in this section.

Some unusual flow reactors are shown in Figure 17.14. The residence times in the units for high temperature pyrolysis to make acetylene and ethylene and for the oxidation of ammonia are measured in fractions of a second; acetic anhydride is made by mixing reactants quickly in a centrifugal pump; NO is formed at very high temperature in an electric furnace; and ethylene is polymerized at high or low pressures in the two units shown.



Figure 17.11. Types of contactors for reacting gases with liquids; many of these also are suitable for reacting immiscible liquids. Tanks: (a) with a gas entraining impeller; (b) with baffled impellers; (c) with a draft tube; (d) with gas input through a rotating hollow shaft. (e) Venturi mixer for rapid reactions. (f) Self-priming turbine pump as a mixer-reactor. (g) Multispray chamber. Towers: (h) parallel flow falling film; (i) spray tower with gas as continuous phase; (j) parallel flow packed tower; (k) counter flow tray tower. (1) A doublepipe heat exchanger used as a tubular reactor.



Gas

Liaui d

(k)



Figure 17.12. Ratio of volumes of an n-stage CSTR battery and a plug flow eactor as a function of residual concentration ratio C/C, with a rate equation $r = kC^2$.



GAS-LIQUID REACTIONS

Except with highly volatile liquids, reactions between gases and liquids occur in the liquid phase, following a transfer of gaseous participants through gas and liquid films. The rate of mass transfer always is a major or limiting factor in the overall transformation process. Naturally the equipment for such reactions is similar to that for the absorption of chemically inert gases, namely towers and stirred tanks. Figure 17.11 illustrates schematically types of gas-liquid reactors. Figure 17.17 shows specific examples of such reactors: In the synthesis of butynediol, acetylene at high pressure is bubbled into aqueous formaldehyde at several positions along a tower in (a). The heat of absorption of nitrogen oxides in water to make nitric acid is removed in two ways in the equipment of (b) and (e). Fats are hydrogenated in a continuous multistage stirred reactor in (c) and under batch conditions in a coil-cooled stirred tank in (d). A thin film reactor is used for the sulfonation of dodecylbenzene with SO, in (f). Hydrogen is recirculated with a hollow-shaft agitator to convert nitrocaprolactam in (g). A shell-and-tube design is used for the reaction of ammonia and adipic acid in (h).

Reactions between gases and liquids may involve solids also, either as reactants or as catalysts. Table 17.9 lists a number of examples. The lime/limestone slurry process is the predominant one for removal of SO, from power plant flue gases. In this case it is known that the rate of the reaction is controlled by the rate of mass transfer through the gas film.

Some gases present in waste gases are recovered by scrubbing with absorbent chemicals that form loose compounds; the absorbent then may be recovered for reuse by elevating the temperature or lowering the pressure in a regenerator. Such loose compounds may exert appreciable back pressure in the absorber, which must be taken into account when that equipment is to be sized.

572 CHEMICAL REACTORS

In all cases, a limiting reactor size may be found on the basis of mass transfer coefficients and zero back pressure, but a size determined this way may be too large in some cases to be economically acceptable. Design procedures for mass transfer equipment are in other chapters of this book. Data for the design of gas-liquid reactors or chemical absorbers may be found in books such as those by Astarita, Savage, and Bisio (Gas *Treating* with *Chemical* Solvents, Wiley, New York, 1983) and Kohl and Riesenfeld (Gas *Purification*, Gulf, Houston, TX, 1979).

FIXED BED REACTORS

The **fixed** beds of concern here are made up of catalyst particles in the range of 2-5 mm dia. Vessels that contain inert solids with the sole purpose of improving mass transfer between phases and developing plug flow behavior are not in this category. Other uses of inert packings are for purposes of heat transfer, as in pebble heaters and induction heated granular beds-these also are covered elsewhere.

The catalyst in a reactor may be loaded in several ways, as:

- 1. a single large bed,
- 2. several horizontal beds,
- 3. several packed tubes in a single shell,

- 4. a single bed with imbedded tubes,
- 5. beds in separate shells.

Some of the possibilities are illustrated in Figures 17.13 and 17.18. Variations from a single large bed are primarily because of a need for control of temperature by appropriate heat transfer, but also for redistribution of the flow or for control of pressure drop. There are few fixed bed units that do not have some provision for heat transfer. Only when the heat of reaction is small is it possible to regulate the inlet temperature so as to make adiabatic operation feasible; butane dehydrogenation, for example, is done this way.

Because of their long industrial histories and worldwide practice, the sulfuric acid and ammonia industries have been particularly inventive with regard to reactors. A few designs for SO, oxidation are illustrated in Figure 17.19. Their dominant differences are in modes of temperature control to take advantage of high rates of reaction at high temperature and favorable equilibrium conversion at lower temperatures. Figure 17.19(g) shows the temperature profile achieved in that equipment, and Figure 17.20 presents patterns of temperature control in the production of SO,, ammonia, and methanol.

A selection of ammonia reactors is illustrated in Figures 17.21 and 17.22. These vessels incorporate particularly elaborate means for temperature regulation. The basic flow pattern is indicated in



Figure 17.13. Multibed catalytic reactors: (a) adiabatic; (b) interbed coldshot injection; (c) shell and tube; (d) built-in interbed heat exchanger; (e) external interbed exchanger; (f) autothermal shell, outside influent-effluent heat exchanger; (g) multishell adiabatic reactor with interstage fired heaters; (h) platinum-catalyst, fixed bed reformer for 5000 bpsd charge rate; reactors 1 and 2 are 5.5 ft dia by 9.5 ft high and reactor 3 is 6.5×12.0 ft.

Figure 17.21(a), and some temperature profiles in Figures 17.22(d) and **17.23(e)**. For modern high capacity performance in single units, reactors with short travel paths through the catalyst and pressures below 200atm are favored. Comparative performance data over a range of conditions appear in Figure 17.22.

Thermal effects also are major factors in the design of reactors

for making synthetic fuels. The units of Figure 17.24 for synthesis of methanol and gasoline are typical fixed bed types.

Catalytic reformers upgrade low octane naphthas into gasoline in the presence of hydrogen to retard deposition of carbon on the catalyst. Temperatures to 500°C and pressures to 35 atm are necessary. Representative reactors are shown in Figure 17.25.



Figure 17.14. Some unusual reactor configurations. (a) Flame reactor for making ethylene and acetylene from liquid hydrocarbons [*Patton et* al., Pet Refin **37**(*11*) *180*, (*1958*)]. (b) Shallow bed reactor for oxidation of ammonia, using Pt-Rh gauze [*Gillespie and Kenson*, Chemtech, 625 (Oct. *1971*)]. (c) Schoenherr furnace for fixation of atmospheric nitrogen. (d) Production of acetic acid anhydride from acetic acid and gaseous ketene in a mixing pump. (e) Phillips reactor for low pressure polymerization of ethylene (closed loop tubular reactor). (f) Polymerization of ethylene at high pressure.



Figure 17.14—(continued)

Feedstocks to such units usually must be desulfurized; a reactor like that of Figure 17.26 hydrogenates sulfur compounds to hydrogen sulfide, which is readily removed.

Fluid flow through fixed bed reactors usually is downward. Instead of screens for supporting catalyst in the vessel, a support of graduated sizes of inert material is used, as illustrated in Figure 17.27. Screens become blinded by the small particles of catalyst. A similar arrangement is used at the top to prevent disturbance of the catalyst level by the-high velocity fluids.

MOVING BEDS

In such vessels granular or lumpy material moves vertically downward as a mass. The solid may be a reactant or a catalyst or a heat carrier. The reactor of Figure 17.28(a) was used for the fixation of nitrogen in air at about 4000°F. The heat-carrying pebbles are heated by direct contact with combustion gases, dropped into a reaction zone supplied with reacting air, and then recycled with elevators to the reheating zone. The treated air must be quenched



Figure 17.15. A fired heater as a high temperature reactor. (a) Arrangement of tubes and burners: (1) radiant tubes; (2) radiant panel burners; (3) stack; (4) convection chamber tubes (*Sukhanov*, Petroleum Processing, *Mir*, Moscow, 1982). (b) Radiant (surface-combustion) panel burner: (1) housing; (2) ceramic perforated prism; (3) tube; (4) injector; (5) fuel gas nozzle; (6) air throttle (*Sukhanov*, Petroleum Processing, *Mir*, *Moscow*, 1982). (c) Fired tubular cracking furnace for the preparation of ethylene from naphtha.



Figure 17.16. Basic types of tubular furnaces [Nelson, Petroleum Refinery Engineering, McGraw-Hill, 1958. Courtesy McGraw-Hill, New York].

at the rate of about 25,000°F/sec to retain a concentration of about 1% nitrogen oxides. In another such operation, two units are used in parallel, one being heated while the other is reacting.

The pebble heater, Figure 17.28(b), is used in the same manner; its application to the pyrolysis of oils to make ethylene also did not prove competitive and has been abandoned.

Units like that of Figure 17.28(c) were employed at one time in the catalytic cracking of gas oils. The catalyst is transferred between regenerating and reacting zones with bucket elevators or air lifts. Some data for this equipment are given with the figure.

Two examples in which the solid itself is reactive are the shale oil retorts of Figure 17.29. Crushed oil shale is charged at the top,

air and gaseous fuel at the bottom. When the shale moving downward reaches a temperature of 900°F, the kerogen decomposes into oil vapor, gas, and carbonaceous residue. There are many designs of pilot plant retorts, but the only commercial units at present are in the USSR and China.

KILNS AND HEARTH FURNACES

These units are primarily for high temperature services, the kilns up to 2500° F and the furnaces up to 4000° F. Usual construction is steel-lined with ceramics, sometimes up to several feet in thickness.

Vertical kilns are used for materials that do not fuse or soften,

576 CHEMICAL REACTORS

as for the burning of limestone or dolomite. Many such operations are batch: the fresh solid is loaded into the kiln, heated with combustion products until reaction is complete, and then dumped. The lime kiln of Figure 17.30(c), however, operates continuously as a moving bed reactor. These vessels range in size from 8 to 15 ft dia and are 50-80 ft high. For calcination of lime the peak temperatures are about 2200°F, although decomposition proceeds freely at 1850°F. Fuel supply may be coke mixed with the limestone if the finished lime can tolerate the additional ash, or gaseous or liquid fuels. Space velocity is 0.8-1.5 lb CaO/(hr)(cuft of kiln), or 45-100 lb CaO/(hr)(sqft of kiln cross section), depending on the

size and modernity of the kiln, the method of firing, and the lump size which is in the range of 4-10 in.

Rotary kilns have many applications as reactors: between finely divided solids (cement), between liquids and solids (salt cake from salt and sulfuric acid), between gases and solids, and for the decomposition of solids (SO, and lime from $CaSO_4$). The kiln is a long narrow cylinder with a length-to-diameter ratio of 10-20. General purpose kilns are 100-125 ft long, but cement kilns as large as 12ft dia by 425 ft long are operated. An inclination to the horizontal of 2-5 deg is sufficient to move the solid along. Speed of rotation is 0.25-2 rpm. Lumps up to 1 in. dia or fine powders are



Figure 17.17. Examples of reactors for specific liquid-gas processes. (a) Trickle reactor for synthesis of butinediol 1.5 m dia by 18 m high. (b) Nitrogen oxide absorption in packed columns. (c) Continuous hydrogenation of fats. (d) Stirred tank reactor for batch hydrogenation of fats. (e) Nitrogen oxide absorption in a plate column. (f) A thin film reactor for making dodecylbenzene sulfonate with SO,. (g) Stirred tank reactor for the hydrogenation of caprolactam. (b) Tubular reactor for making adiponitrile from adipic acid in the presence of phosphoric acid.





Figure 17.17—(continued)

TABLE 17.9. Examples of Fluidized Bed Processes

A. Catalytic Processes

- 1. Oil cracking and reforming
- 2. Recovery of high concentrations of benzene from gas oils
- 3. Olefin production from crude oil
- 4. Chlorine by oxidation of HCI
- 5. Acetylene from methane
- 6. Preparation of unsaturated aldehydes
- 7. Reduction of nitro compounds to amines
- 6. Oxidation of SO, to SO,
- 9. Phthalic anhydride from naphthalene or o-xylene
- 10. Maleic acid anhydride from benzene
- 11. Formaldehyde from methanol
- 12. Chlorination of methane and ethylene
- 13. Fischer-Tropsch synthesis of gasoline
- 14. Hydrogenation of ethylene
- 15. Oxidation of ammonia
- 16. Ethylene oxide from ethylene
- 17. Butadiene from ethanol
- 18. Dehydrogenation of isopropanol
- 19. Isomerization of n-butane
- 20. Post-chlorination of PVC
- 21. Decomposition of ozone
- 22. Preparation of chlorinated hydrocarbons
- 23. Preparation of melamine resins
- 24. Isoprene synthesis
- 25. Reduction of vinyl acetate
- 26. Preparation of acrylonitrile

B. Noncatalytic Processes

- 1. Gasification of coal
- 2. Fluid bed coking
- 3. Pyrolytic cracking of methane
- 4. Preparation of activated carbon
- 5. Ethylene by cracking of petroleum fractions
- 6. Combustion of coal

- 7. Burning of oil shale
- 8. Combustion of municipal and industrial wastes
- 9. Burning of black liquor (paper industry)
- 10. Roasting of sulfides of iron, copper, and zinc
- 11. Combustion of sulfur in a sand bed
- 12. Decomposition of waste sulfuric acid and sulfates
- 13. Cracking of chlorides such as FeCl₂, NiCl₃, and AlCl₃
- 14. Volatilization of rhenium
- 15. Burning of limestone and dolomite
- 16. Cement burning
- 17. Reduction of iron ores and metallic oxides
- 18. Chlorination of ores of aluminum, titanium, nickel, cobalt, and tin
- 19. Chlorination of roasted pyrites and iron ores
- 20. Chlorination of lime
- 21. Calcination of aluminum hydroxide to alumina
- 22. Preparation of aluminum sulfate from bauxite
- 23. Preparation of fluorides aluminum trifluoride, uranium tetra- and hexafluorides
- 24. Preparation of pure tungsten from the fluoride
- 25. Calcination of phosphates
- 26. Preparation of phosphorus oxychloride
- 27. Preparation of carbon disulfide
- 28. Preparation of hydrazine
- 29. Preparation of nitric acid
- 30. Preparation of nitrates of ammonia and sodium
- 31. Preparation of sodium carbonate
- 32. Preparation of hydrogen cyanide
- 33. Hydrochlorination of uranium fuel elements
- 34. Preparation of uranium trioxide from the nitrate
- 35. Recovery of uranium from nuclear fuels
- 36. Removal of fluorine from offgases of aluminum electrolysis
- 37. Heating of heat transfer media such as sand
- 38. Cooling of granular masses such as fertilizers
- 39. Drying of finely divided materials such as flotation ores and raw phosphates
- 40. Coating of fuel elements by pyrolytic cracking of chlormethylsilanes



Figure 17.18. Heat transfer in fixed-bed reactors: (a) adequate preheat; (b) internal heat exchanger; (c) annular cooling spaces; (d) packed tubes; (e) packed shell; (f) tube and thimble; (g). external heat exchanger; (h) multiple shell, with external heat transfer (*Walas*, 1959).

usual. Heating mostly is with combustion gases, but some low temperature heating may be accomplished through heated jackets. Figures 17.30(a) and (b) show the temperature profiles of gas and stock in a cement kiln and space velocities of a number of kiln processes.

Multiple-hearth furnaces are suited to continuous handling of solids that exhibit a limited amount of fusion or sintering. In the kind shown on Figure 17.30(e), the scrapers rotate, in other kinds the plates rotate, and in still others the scrapers oscillate and discharge the plates at each stroke. Material is charged at the top, moves along as rotation proceeds, and drops onto successively lower plates while combustion gases or gaseous reactants flow upward. This equipment is used to roast ores, burn calcium sulfate or bauxite, and reactivate the absorbent clays of the petroleum industry. A reactor with nine trays, 16 ft dia and 35 ft high can roast about 1,250 lb/hr of iron pyrite, at a residence time of about 4-5 hr.

Hearth furnaces consist of one or more flat or concave pans, either moving or stationary, usually equipped with scraper-stirrers. Although such equipment is used mostly for ore treating and metallurgical purposes, a few inorganic chemical processes utilize them, for example, Leblanc soda ash, sodium sulfide from salt cake and coal, sodium sulfate and hydrogen chloride from salt and sulfuric acid, and sodium silicate from sand and soda ash. A kind of salt-cake furnace is shown in Figure 17.30(d). Salt and sulfuric acid are charged continuously to the center of the pan, and the rotating scrapers gradually work the reacting mass towards the periphery where the sodium sulfate is discharged over the edge. The products leave at about 1000°F. Pans of 11-18 ft dia can handle 6-10 tons/24 hr of salt. For comparison, a Laury horizontal rotating cylindrical kiln, 5 \times 22 ft, has a capacity of 1 ton/hr of salt cake.

Very high temperature operations such as those of production of glass or metals utilize single-hearth furnaces, often with heat



Figure 17.19. Reactors for the oxidation of sulfur dioxide: (a) Feed-product heat exchange. (b) External heat exchanger and internal tube and thimble. (c) **Multibed** reactor, cooling with charge gas in a spiral jacket. (d) Tube and thimble for feed against product and for heat transfer medium. (e) BASF-Knietsch, with autothermal packed tubes and external exchanger. (f) Sper reactor with internal heat transfer surface. (g) Zieren-Chemiebau reactor assembly and the temperature profile (*Winnacker-Weingartner*, Chemische Technologie, Carl Hanser Verlag, Munich, 1950-1954).



Figure 17.19—(continued)

regenerators for fuel economy. The Siemens-Martin furnace of Figure 17.30(d) with a hearth 13ft wide and 40ft long has a production rate of 10 tons/hr of steel with a residence time of 10 hr. The hearth volume is about 5000 cuft and the total regenerator volume is about 25,000 cuft,

FLUIDIZED BED REACTORS

This term is restricted here to equipment in which finely divided solids in suspension interact with gases. Solids fluidized by liquids are called slurries. Three phase fluidized mixtures occur in some coal liquefaction and petroleum treating processes. In dense phase gas-solid fluidization, a fairly definite bed level is maintained; in dilute phase systems the solid is entrained continuously through the reaction zone and is separated out in a subsequent zone.

The most extensive application of fluidization has been to catalytic cracking of petroleum fractions. Because the catalyst degrades in a few minutes, it is circulated continuously between reaction and regeneration zones. Figure 17.31(a) is a version of such equipment. The steam stripper is for the removal of occluded oil

before the catalyst is to be burned. The main control instrumentation of a side-by-side system is shown in Figure 3.6(h).

Fluid catalytic vessels are very large. Dimensions and performance of a medium capacity unit (about 50,000 BPSD, 60 kg/sec) are shown with the figure. Other data for a reactor to handle 15,000 BPSD are a diameter of 25 ft and a height of 50 ft. Catalyst holdup and other data of such a reactor are given by Kraft, Ulrich, and O'Connor (in Othmer (Ed.), *Fluidization*, Reinhold, New York, 1956) as follows:

Item	Quantity		
Unit charge, nominal	15,000 BPSD		
Catalyst inventory, total	250 tons		
Catalyst inventory, regenerator bed	100 tons		
Superficial velocity, regenerator	2.5 fps		
Bed density, regenerator	28.0 lb/cuft		
Flue gas plus solids density, cyclone inlet	0.5 lb/cuft		
Catalyst circulation rate, unit	24.0 tons/min		
Catalyst circulation rate, to cyclones	7.0 tons/min		
Catalyst loss rate, design expectation	2.0 tons/day		

Figure 17.31(b) is of a unit in which most of the cracking occurs



Figure 17.20. Control of temperature in multibed reactors so as to utilize the high rates of reaction at high temperatures and the more favorable equilibrium conversion at lower temperatures. (a) Adiabatic and isothermal reaction lines on the equilibrium diagram for ammonia synthesis. (b) Oxidation of SO, in a four-bed reactor at essentially atmospheric pressure. (c) Methanol synthesis in a four bed reactor by the ICI process at 50 atm; not to scale; 35% methanol at 250°C, 8.2% at 300°C, equilibrium concentrations.

in a transfer line, an operation that became feasible with the development of highly active zeolite catalysts. The reaction is completed in the upper zone, but the main function of that zone is to separate product and spent catalyst. In contrast to the dense-phase bed of a large reactor, in which mixing can approach ideality, the dilute phase transfer line is more nearly in plug flow. Accordingly, a much smaller reaction zone suffices; moreover, superior product distribution and greater gasoline yield result. Similar reactor configurations are shown in Figures 17.31(c) and (d) of other petroleum processes.

The mechanism of interaction between catalyst and gas in a large **fluidized** bed is complex and is not well correlated with design factors. In the bed itself, large bubbles of a foot or more in diameter form and are irrigated with a rain of catalyst particles. This process occurs in parallel with a well-mixed fluidized bed. Above the bed level and before the entrained catalyst is recovered in cyclones, the reaction continues in dilute phase plug flow. Since even the physical behavior of fluidized beds is not well understood, the design of such reactors is done largely on the basis of fairly large pilot plants and by analogy with earlier experience in this area.



(c)

Figure 17.21. Some recent designs of ammonia synthesis converters. (a) Principle of the autothermal ammonia synthesis reactor. Flow is downwards along the wall to keep it cool, up through tubes imbedded in the catalyst, down through the catalyst, through the effluent-influent exchanger and out. (b) Radial flow converter with capacities to 1800 tons/day (*Haldor Topsoe Co., Hellerup*, *Denmark*). (c) Horizontal three-bed converter and detail of the catalyst cartridge. Without the exchanger the dimensions are 8 X 85 ft, pressure 170 atm, capacity to 2000 tons/day (Pullman *Kellogg*). (d) Vessel sketch, typical temperature profile and typical data of the ICI quench-type converter. The process gas follows a path like that of part (a) of this figure. Quench is supplied at two points (*Imperial Chemical Industries*).



Typical Data for	ICI Quench	Converters	of Various	Sizes
Capacity (short tpd)	660	990	1100	1650
Pressure (psig)	4700	3200	4250	3220
Inlet gas composition (%) Ammonia	4.0	3.0	3.2	T 4
Inerts	15.0	12.0	15.0	12.0
Inlet g flow (MM scfh)	10.6	18.0	18.5	24.5
Catalyst volume (ft')	740	1170	1100	2400
Pressure vessel Internal diameter (in.) Length (in.)	80 437	96 493	95 472	109
Weight (short ton) Cartridge shell Heat exchanger Pressure vessel (less cover)	14.2 15.5 130	34.2 30.0 128	22.8 25.4 182	56.4 23.8 240

Typical Data for an ICI Quench Converter of 1300 Short

Tons/Day Capacity				
Pressure (psig)	2200	3200	4000	4700
Inlet gaow (MM scfh)"	25.8	21.2	19.8	19.0
Catalyst volume (ft')	2600	1730	1320	1030
Pressure vessel	170	103	0.6	00
Length (in.)	663	102 606	528	488
Weight (short ton)				
Cartridge shell	68.5	40.8	29.2	23.6
Heat exchanger	37.1	25.4	20.7	17.9
Pressure vessel (less cover)	186	184	187	189
Converter pressure drop (psi)	140	104	87	91

"Composition: 2% NH,, 12% inerts (CH₄+A), 21.5% N₂, 64.5% H₂ by vol.

Figure 17.21—(continued)

The earliest fluidized process was the noncatalytic Winkler process for gasification of coal in 1921. Other noncatalytic processes, and some catalytic ones, are listed in Table 17.9. A few noncatalytic reactors are shown in Figure 17.32. Cracking of naphthas to ethylene with circulating hot sand as the heat carrier is shown in part (a); at the operating temperature of 720-850°C, much carbon deposits on the sand but is not at all harmful as it would be on the surfaces of tubular cracking units. In the dilute phase process of calcination of alumina, part (b), the circulating solid is the product itself; combustion products from sprays of oil and auxiliary air furnish the motive power. The calcining unit for lime of part (c) is an example of a successful multistage reactor; residence time in the calcining zone is 2 hr, in the cooling zone 0.5 hr, and in each of the preheating zones 1 hr. Multibed units for petroleum operations have not been feasible, but some units have been built with a degree of baffling that simulates staging in a rough fashion. The catalyst of the phthalic anhydride reactor of part (d) does not need to be regenerated so the fluidized bed remains in place; since the reaction is highly sensitive to temperature, the oxidation is kept under control with much imbedded heat transfer surface and by cold injections. A modern coal gasifier appears in part (e); a thirtyfold circulation of spent char is employed along with the fresh feed to counteract the agglomeration tendency of many coals. The H-Coal reactor of part (f) operates with a three-phase mixture. The catalyst does not circulate but bubbles in place. Activity is maintained by bleeding off and replenishing 1-2% of the catalyst

holdup per day. Operating conditions are 450°C and 3000 psig. Both coal and heavy petroleum residua are handled successfully. The unit is known as an "ebullating bed."

The literature of fluidization phenomena and technology is extensive. A good although dated bibliography is in *Ullmann's Encyclopedia* (1973, Vol. 3, pp. 458–460). The book by Cheremisinoff and Cheremisinoff (1984) has more than 500 abstracts of articles on fluidization hydrodynamics, mixing and heat transfer, but little on reactor technology. Other literature on fluidization is cited in the References of Chapter 6.

17.7. HEAT TRANSFER IN REACTORS

Maintenance of proper temperature is a major aspect of reactor operation. The illustrations of several reactors in this chapter depict a number of provisions for heat transfer. The magnitude of required heat transfer is determined by heat and material balances as described in Section 17.3. The data needed are thermal **conductivities** and coefficients of heat transfer. Some of the factors influencing these quantities are associated in the usual groups for heat transfer; namely, the Nusselt, Stanton, Prandtl, and Reynolds dimensionless groups. Other characteristics of particular kinds of reactors also are brought into correlations. A selection of practical results from the abundant literature will be assembled here. Some modes of heat transfer to stirred and fixed bed reactors are represented in Figures 17.33 and 17.18, and temperature profiles in



Figure 17.22. Representative ammonia converters operating at various pressures and effluent concentrations (*Vancini*, 1971). (a) Original Uhde design operating at 12.5 atm; typical dimensions, $1.4 \times 7 \text{ m}$. (b) Haber-Bosch-Mittasch converter operating at 300atm; typical dimensions, $1.1 \times 12.8 \text{ m}$. (c) Claude converter operating at 1000 atm; typical dimensions $1.2 \times 7 \text{ m}$. (d) Fauser-Montecatini (old style) converter operating at 300 atm with external heat exchange, showing axial profiles of temperature and ammonia concentration.

	Comparison	of Perfo	ormance	
Process	Pressure (bar)	Effluent ammonia (%)	TPD/m ³	Catalyst life (yr)
Uhde	125	7-a	10	>2
Haber-Bosch	300	13-15	25	2
Claude	1000	22-24	120	0.25
Fauser	300	12-17	25	2



Figure 17.22—(continued)



Figure 17.23. Representative temperature profiles in reaction systems (see also Figs. 17.20, 17.21(d), 17.22(d), 17.30(c), 17.34, and 17.35). (a) A jacketed tubular reactor. (b) Burner and reactor for high temperature pyrolysis of hydrocarbons (*Ullmann*, 1973, *Vol.* 3, *p.* 355); (c) A catalytic reactor system in which the feed is preheated to starting temperature and product is properly adjusted; exo- and endothermic profiles. (d) Reactor with built-in heat exchange between feed and product and with external temperature adjustment; exo- and endothermic profiles.









Figure 17.24. Types of reactors for synthetic fuels [Meyers (Ed.), Handbook of Synfuels Technology, *McGraw-Hill, New York, 1984*]. (a) ICI methanol reactor, showing internal distributors. C, D and E are cold shot nozzles, F = catalyst dropout, L = thermocouple, and 0 = catalyst input. (b) ICI methanol reactor with internal heat exchange and cold shots. (c) Fixed bed reactor for gasoline from coal synthesis gas; dimensions 10 x 42 ft, 2000 2-in. dia tubes packed with promoted iron catalyst, production rate 5 tons/day per reactor. (d) Synthol fluidized bed continuous reactor system for gasoline from coal synthesis gas.





J-pipe connection for inlet of starting material; 2-zonal thermocouple; 3-pipe connection for product discharge; 4-external thermocouple; 5-pipe connection for discharge of product during ejecting of system (in catalyst regeneration); 6-pipe connection for catalyst discharge; 7-light-weight fireclay; g-reactor housing; 9-gunite lining; 10-porcelain balls

(a)

l-multizonal thermocouple; 2-reactor housing; 3-tining; 4-surface thermocouple; 5-porcelain balls; 6-pipe connection for discharge of product during electing of system (in catalyst regeneration); 7-pipe connection for catalyst discharge; 8-catalyst. Lines: 1-gas-product mixture; 11-reaction products

(b)

Figure 17.25. Catalytic reforming reactors of axial and radial flow types. The latter is favored because of lower pressure drop (*Sukhanov*, Petroleum Processing, *Mir*, Moscow, 1982). (a) Axial flow pattern. (b) Radial flow pattern.

several industrial reactors appear in Figures 17.20-17.23, 17.30, 17.34. and 17.35.

STIRRED TANKS

Values of overall coefficients of heat transfer are collected in Tables 17.10–17.12. Two sets of formulas for tank-side film coefficients are in Tables 17.13 and 17.14. They relate the Nusselt number to the Reynolds and Prandtl numbers and several other factors. In the equation for jacketed tanks, for example,

$$h_{0}(\text{jacket}) \frac{T}{k} = 0.85 \left(\frac{D^{2} N \rho}{\mu}\right)^{0.66} \left(\frac{C_{p} \mu}{k}\right)^{0.33} \left(\frac{\mu}{\mu_{s}}\right)^{0.14} \\ \times \left(\frac{Z}{T}\right)^{-0.56} \left(\frac{D}{T}\right)^{0.13}$$
(17.35)

the rpm, the tank and impeller diameters, and the liquid depth as well as a viscosity ratio are involved. Table 17.14 identifies the kind of impeller that was used in the investigation, but in general test results have shown that approximately the same heat transfer coefficient is obtained with flat-blade turbines, pitched-blade turbines, or propellers. Axial flow turbines produce the most circulation for a given power input and heat transfer is related directly to the flow, so that this kind of impeller usually is favored. From Eq. (17.35), the coefficient is proportional to the 0.66 power of the rpm, $N^{0.66}$, and from Chapter 10, the power input at high Reynolds numbers varies as the cube root of N. Accordingly it appears that the coefficient is proportional to the 0.22 exponent of the power input to the stirred tank,

$$h \propto P^{0.22}$$



1-bousing: q-support ring; 3-inlet pipe connection; 4-discharge pipe connection; 5-spere; 6-catalyst discharge pipe connection; 7-porcelain balls; e-catalyst space; 9-reactor lining (mesh-reinforced gunite): 12-distribution tray (for uniform distribution of catalyst during charging); 11-union for zonal thermocoupie; 12-union for thermocoupie; 13-lightweight fireday brick; 14baltie

Figure 17.26. Reactor for hydrofining diesel oils, with ceramic lining (*Sukhanov*, Petroleum Processing, *Mir*, Moscow, 1982).

and consequently that the coefficient of heat transfer is little affected by large increases of power input.

Since most of the literature in this area is relatively old, practitioners apparently believe that what has been found out is adequate or is kept confidential. Table 17.14 has the recommended formulas.

PACKED BED THERMAL CONDUCTIVITY

The presence of particles makes the effective conductivity of a gas greater than the molecular conductivity by a factor of 10 or more. The nature of the solid has little effect at Reynolds numbers above 100 or so; although the effect is noticeable at the lower values of Re, it has not been completely studied. Besides the Reynolds, Prandtl, and Peclet numbers, the effective diffusivity depends on the molecular conductivity, porosity, particle size, and flow conditions. Plots in terms of Re, Pr, and Pe (without showing actual data points) are made by Beek (1962, Fig. 3), but the simpler plots obtained by a number of investigators in terms of the Reynolds number alone appear on Figure 17.36(a). As Table 17.15 shows, most of the data were obtained with air whose Pr = 0.72 and



Figure 17.27. Catalyst packed adiabatic reactor, showing application of ceramic balls of graduated sizes for support at the bottom and hold-down at the top (Ruse, Chemical Reactor Design for Process Plants, *Wiley, New York, 1977*).

 $k_f = 0.026 \text{ kcal/(m)(hr)(°C)}$ at about 100°C. Accordingly, the data could be generalized to present the ratio of effective and molecular conductivities as

$$k_e/k_f = 38.5k_e. (17.36)$$

Equations of the highest and lowest lines on this figure then may be written

$$k_e/k_f = 8.08 + 0.1027 \text{Re}$$
 (Kwong and Smith), (17.37)
 $k_e/k_f = 13.85 + 0.0623 \text{Re}$ (Quinton and Storrow). (17.38)

At higher temperatures, above 300°C or so, radiation must contribute to the effective conductivity, but there are so many other uncertainties that the radiation effect has not been studied at length.

HEAT TRANSFER AT WALLS, TO PARTICLES, AND OVERALL

The correlations cited in Tables 17.16 and 17.18 are of the Nusselt number in terms of the Reynolds and Prandtl numbers, or of the Reynolds alone. They are applicable only above specified Reynolds numbers, about 40 in most cases; clearly they do not predict correctly the coefficient of natural convection, at Re = 0.

Wall coefficients are obtainable from particle-fluid data by a rule of Beek (1962),

$$h_w = 0.8h_p$$
 (17.39)

This is how Eq. (8) of Table 17.18 is deduced; Eq. (9) represents



Figure 17.28. Reactors with moving beds of catalyst or solids for heat supply. (a) Pebble reactor for direct oxidation of atmospheric nitrogen; two units in parallel, one being heated with combustion gases and the other used as the reactor [*Ermenc*, Chem. Eng. Prog. 52, 149 (1956)]. (b) Pebble heater which has been used for making ethylene from heavier hydrocarbons (*Batchelder and Ingols*, *U.S. Bureau of Mines Report* Invest. No. 4781, 1951). (c) Moving bed catalytic cracker and regenerator; for 20,000 bpsd the reactor is 16ft dia, catalyst circulation rate 2–7 lbs/lb oil, attrition rate of catalyst 0.1-0.5 lb/ton circulated, pressure drop across air lift line is about 2 psi (*L. Berg in Othmer (Ed.)*, Fluidization, *Reinhold, New York*, 1956).



Figure 17.29. Moving bed reactors for cracking and recovery of shale oil. (a) Kiviter retort, USSR 200-300 tons/day [J. W. Smith, in Meyers (Ed.), Handbook of Synfuels Technology, McGraw-Hill, New York, 1984]. (b) Paraho retort for shale oil recovery (Paraho Oil Shale Demonstration, Grand Junction, CO).

the same data but is simply a curve fit of Figure 17.36(c) at an average value Pr = 0.65.

Data of heat transfer between particle and fluid usually are not measured directly because of the experimental difficulties, but are deduced from measurements of mass transfer coefficients assuming the Colburn analogy to apply,

$$(Sherwood)(Schmidt)^{2/3} = (Nusselt)(Prandtl)^{2/3}$$

= function of Reynolds. (17.40)

Thus, in Figure 17.36(c), if the Nusselt number is replaced by the Sherwood and the Prandtl by the Schmidt, the relation will be equally valid for mass transfer.

The ratio, L/D, of length to diameter of a packed tube or vessel has been found to affect the coefficient of heat transfer. This is a dispersion phenomenon in which the **Peclet** number, uL/D_{disp} , is involved, where D_{disp} is the dispersion coefficient. Some 5000 data points were examined by **Schlünder** (1978) from this point of view; although the effect of L/D is quite pronounced, no clear pattern was deduced. Industrial reactors have L/D above 50 or so; Eqs. (6) and (7) of Table 17.18 are asymptotic values of the heat transfer coefficient for such situations. They are plotted in Figure 17.36(b).

Most investigators have been content with correlations of the Nusselt with the Reynolds and Prandtl numbers, or with the Reynolds number alone. The range of numerical values of the Prandtl number of gases is small, and most of the investigations have been conducted with air whose Pr = 0.72 at 100°C. The effect of Pr is small on Figure 17.36(c), and is ignored on Figure 17.36(b) and in some of the equations of Tables 17.17 and 17.18.

The equations of Table 17.18 are the ones recommended for coefficients of heat transfer between wall and fluid in packed vessels.

For design of equipment like those of Figure 17.28, coefficients of heat transfer between particle and fluid should be known. Direct measurements with this objective have been made with metallic packings heated by electrical induction or current. Some correlations are given in Table 17.17. Glaser and Thodos [*AIChE J.* 4, 63 (1958)] correlated such data with the equation

$$(h_p/C_p G)(C_p \mu/k)^{2/3} = \frac{0.535}{(\text{Re}')^{0.3} - 1.6}, \quad 100 < \text{Re}' < 9200,$$
(17.41)

where

Re' = $\phi \sqrt{a_p} G/\mu (1 - E)$, a_p = surface of a single particle, ϕ = sphericity of the particle.

The formulas of Table 17.18 also could be used by taking $h_p = 1.25h_w$. In moving bed catalyst regenerators, heat fluxes of the order of 25,000 Btu/(hr)(cuft) have been estimated to occur between fluid and particle. Fluid and particle temperatures consequently differ very little.

FLUIDIZED BEDS

A distinctive feature of fluidized beds is a high rate of heat transfer between the fluid and immersed surfaces. Some numerical values are shown on Figure 17.37. For comparison, air in turbulent flow in pipelines has a coefficient of about 25 Btu/(hr)(sqft)(°F). (a) is of calculations from several correlations of data for the conditions identified in Table 17.19; (b) shows the effect of diameters of quartz particles; and (c) pertains to 0.38 mm particles of several substances.



Process	Lb/(hr)(cuft of kiln)
Cement, dry process	1.1-2.6
Cement, wet process, 44% water	0.7-1 .7
Limestone calcination	1.1-2.1
Dolomite calcination	1.0-I .4
Alumina preparation	1.2-I . 7
Barium sulfide preparation	0.9-2.0
Ignition of inorganic pigments	0.4-4.6
Iron pyrite roasting	0.7-0.9

(b)



(c)





Figure 17.30. Kilns and hearth furnaces (*Walas*, 19.59). (a) Temperature profiles in a rotary cement kiln. (b) Space velocities in rotary kilns. (c) Continuous lime kiln for production of approximately 55 tons/24 hr. (d) Stirred salt cake furnace operating at 1000°F, 11–18 ft dia, 6-10 tons salt/24 hr. (e) Multiple-hearth reactor; one with 9 trays, 16 ft dia and 35 ft high roasts 12.50 lb/hr iron pyrite. (f) Siements-Martin furnace and heat regenerators; a hearth 13 ft wide and 40 ft long makes 10 tons/hr of steel with a residence time of 10 hr.



Figure 17.31. Fluidized bed reactor processes for the conversion of petroleum fractions. (a) Exxon Model IV fluid catalytic cracking (FCC) unit sketch and operating parameters. (*Hetsroni*, Handbook of Multiphase Systems, *McGraw-Hill*, *New York*, 1982). (b) A modern FCC unit utilizing active zeolite catalysts; the reaction occurs primarily in the riser which can be as high as 45 m. (c) Fluidized bed hydroformer in which straight chain molecules are converted into branched ones in the presence of hydrogen at a pressure of 1500 atm. The process has been largely superseded by fixed bed units employing precious metal catalysts (*Hetsroni*, loc. cit.). (d) A fluidized bed coking process; units have been built with capacities of 400–12,000 tons/day.





Figure 17.31—(continued)

Temperature in a fluidized bed is uniform unless particle circulation is impeded. Gas to particle heat flow is so rapid that it is a minor consideration. Heat transfer at points of contact of particles is negligible and radiative transfer also is small below 600°C. The mechanisms of heat transfer and thermal conductivity have been widely studied; the results and literature are reviewed, for example, by Zabrodsky (1966) and by Grace (1982, pp. 8.65-8.83).

Heat transfer behavior is of importance at the walls of the vessel where it determines magnitudes of heat losses to the surroundings and at internal surfaces used for regulation of the operating temperature. The old correlations for heat transfer coefficients of Wender and Cooper (1958) (shown on Fig. 17.38) and those of Vreedenburg (1960) (shown in Table 17.17) still are regarded as perhaps the best. (See Table 17.20.) A fair amount of scatter of the data obtained by various investigators is evident in Figure 17.38. Vreedenburg utilized additional data in his correlating, and, consequently, his figures show even more scatter. On Figure 17.37(a) also there is much disagreement; but if lines 8 and 9 by the same investigators and part of line 3 are ignored, the agreement becomes fair.

Some opinion is that the correlations for vertical tubes should be taken as standard. Coefficients at the wall appear to be about 10% less than at vertical tubes on the axis of the vessel and those for horizontal tubes perhaps 5-6% less (Korotjanskaja et al., 1984, p. 315).

As appears on Figure 17.37, a peak rate of heat transfer is attained. It has been correlated by Zabrodsky et al. (1976) for

particles smaller than 1 mm by the equation

$$h_{\max}d_s/k_g = 0.88 \text{Ar}^{0.213}, \quad 100 < \text{Ar} < 1.4 \text{ x} 10^5, \quad (17.42)$$

Ar = $g\rho_g(\rho_s - \rho_g)d_s^3/\mu_g^2$. (17.43)

17.8. CLASSES OF REACTION PROCESSES AND THEIR EQUIPMENT

In this section, industrial reaction processes are classified primarily with respect to the kinds of phases participating, and instances are given of the kind of equipment that has been found suitable. As always, there is much variation in practice because of local or historical or personal circumstances which suggests that a certain latitude in new plant design is possible.

HOMOGENEOUS GAS REACTIONS

Ethylene is made by pyrolysis of hydrocarbon vapors in tubes of 50–100 mm dia and several hundred meters long with a reaction time of several seconds; heat is supplied by mixing with superheated steam and by direct contact of the tube with combustion gases.

Reactors to make polyethylene are 34-50 mm dia with 10-20 m long turns totalling 400-900 m in length. The tube is jacketed and heated or cooled at different positions with pressurized water.

A flow reactor is used for the production of synthesis gas, CO + H_2 , by direct oxidation of methane and other hydrocarbons in the presence of steam. Preheated streams are mixed and react in a



Figure 17.32. Other fluidized bed reaction systems. (a) Cracking of naphtha to ethylene with circulating hot sand at 720–850°C (Lurgi). (b) Circulating fluidized bed process for production of alumina by calcination (*Lurgi*). (c) Multibed reactor for calcination of limestone (*Dorr-Oliver*). (d) Synthesis of phthalic anhydride; cooling surface is in the bed (*Badger-Sherwin-Williams*). (e) Coal gasifier with two beds to counteract agglomeration, with spent char recirculating at 20-30 times the fresh feed rate (Westinghouse). (f) Ebbulating bed reactor of the H-Coal and H-Oil process for converting these materials at high temperature and pressure into gas and lighter oils (Meyer, Handbook of Synfuels Technology, *McGraw-Hill, 1984*).


Figure 17.32—(continued)



Figure 17.33. Heat transfer to stirred-tank reactors: (a) jacket; (b) internal coils; (c) internal tubes; (d) external heat exchanger; (e) external reflux condenser; (f) fired heater (*Walas*, 1959).



Figure 17.34. Temperature and conversion profiles in a watercooled shell-and-tube phosgene reactor, 2-in. tubes loaded with carbon catalyst, equimolal CO and Cl,.



Figure 17.35. Temperature and conversion profiles of mild thermal cracking of a heavy oil in a tubular furnace with a back pressure of 250 psig and at several heat fluxes [Btu/hr(sqft)].

flow nozzle. Burning and quenching are performed in different zones of a ceramic-lined tower.

HOMOGENEOUS LIQUID REACTIONS

Almost innumerable instances of such reactions are practiced. Single-batch stirred tanks, CSTR batteries, and tubular flow reactors are all used. Many examples are given in Table 17.1. As already pointed out, the size of equipment for a given purpose depends on its type. A comparison has been made of the production of ethyl acetate from a mixture initially with 23% acid and 46% ethanol; these sizes were found for 35% conversion of the acid (Westerterp, 1984, pp. 41-58):

	Re	actor	<i>V, / V</i> 'o [m ³ /(kg/day)]
Batch	(1/3	downtime)	1.04
PFR			0.70
CSTR			1.22
3-stage	CS	TR	0.85

Some of the homogeneous liquid systems of Table 17.1 are numbers 2, 16, 22, 28, 42, 53, 54, and 96, some in batch, mostly continuous.

LIQUID-LIQUID REACTIONS

Such reactions can take place predominantly in either the continuous or disperse phase or in both phases or mainly at the interface. Mutual solubilities, distribution coefficients, and the amount of interfacial surface are factors that determine the overall rate of conversion. Stirred tanks with power inputs of 5-10 HP/1000 gal or extraction-type equipment of various kinds are used to enhance mass transfer. Horizontal TFRs usually are impractical unless sufficiently stable emulsions can be formed, but mixing baffles at intervals are helpful if there are strong reasons for using such equipment. Multistage stirred chambers in a single shell are used for example in butene-isobutane alkylation with sulfuric acid catalyst. Other liquid-liquid processes listed in Table 17.1 are numbers 8, 27, 45, 78, and 90.

GAS-LIQUID REACTIONS

Intimate contacting between chemically reacting gas and liquid phases is achieved in a variety of equipment, some examples of which follow

- a. Tanks equipped with turbine agitators with or without internal gas recirculation. An example is air oxidation of cyclohexane to cyclohexanol and cyclohexanone.
- **b.** Bubble towers with parallel flow of the phases, gas dispersed, with or without trays or packing. In such equipment isobutene from a mixture of C_4 hydrocarbons forms tertiary butanol in contact with aqueous sulfuric acid.
- c. Countercurrent flow of the two phases in tray or packed towers, as in ordinary absorption processes. Absorption of nitrogen oxides in water to make nitric acid is a prime example.
- **d.** Tubular or multitubular reactors are usable when the volumetric rate of the gas is so much greater than that of the liquid that substantial mixing of phases exists. Adipic acid **nitrile** is made from gaseous ammonia and liquid adipic acid with a volumetric ratio of 1500; the residence time of the gas phase is about 1 sec, and that of the liquid 180-300 sec.
- e. Liquid ejector for entraining the gas. This is used to remove dilute acid or other impurities from waste air by scrubbing with aqueous solutions; the liquid is recirculated so that a gas/liquid volumetric ratio of 100-200 is maintained.
- f. Pumps of centrifugal or turbine types are effective mixing devices and can constitute a reactor when the needed residence time is short. Such a device is used, for instance, to make acetic anhydride from acetic acid and ketene [Spes, *Chem. Zng. Tech.* 38, 963 (1966)].
- g. Thin film reactors are desirable when the liquid viscosity is high, the reaction is highly exothermic and short reaction times are adequate. Such a process is the sulfonation of dodecylbenzene with dilute SO₃ [Ujhidy et al., *Chem. Tech.* **18**, 625 (1966)].
- b. Packed tower reactors in parallel flow are operated either top-to-bottom or bottom-to-top. Distribution, holdup, and pressure drop behavior can be predicted from mass transfer correlations. Downflow towers have lower pressure drop, but upflow of liquid assures greater liquid holdup and longer contact time which often are advantages.

NONCATALYTIC REACTIONS WITH SOLIDS

The chief examples are smelting for the recovery of metals from ores, cement manufacture, and lime burning. The converters, roasters, and kilns for these purposes are huge special devices, not usually adaptable to other chemical applications. Shale oil is recovered from crushed rock in a vertical kiln on a batch or continuous basis-moving bed in the latter case-sometimes in a hydrogen-rich atmosphere for simultaneous denitrification and desulfurization. The capacity of ore roasters is of the order of 300-700 tons/(day)(m³ of reactor volume). Rotary kilns for cement have capacities of 0.4-1.1 tons/(day)(m³); for other purposes the range is 0.1-2.

FLUIDIZED BEDS OF NONCATALYTIC SOLIDS

Fluidized bed operations sometimes are alternates to those with fixed beds. Some of the successful processes are fluid bed combustion of coal, cracking of petroleum oils, ethylene production from gas oils in the presence of fluidized sand as a heat carrier, fluidized bed coking, water-gas production from coal (the original fluidized bed operation), recovery of shale oil from rock, reduction of iron ore with hydrogen at 30 atm pressure, lime burning, HCN from coke + ammonia + propane in a fluidized electric furnace, and many others. Many of these processes have distinct equipment configurations and space velocities that cannot be generalized, except insofar as general relations apply to fluidized bed stability,

Fluid Inside Jacket	Fluid In Vessel	Wall Material	Agitation	U
Steam	water	enameled cast iron	O-400 rpm	96-l 20
Steam	milk	enameled C.I.	none	200
Steam	milk	enameled C.I.	stirring	300
Steam	milk boiling	enameled C.I.	none	500
Steam	milk	enameled C.I.	200 rpm	86
Steam	fruit slurry	enameled C.I.	none	33-90
Steam	fruit slurry	enameled C.I.	stirring	154
Steam	water	C.I. and loose lead lining	agitated	4-9
Steam	water	C.I. and loose lead lining	n o n e	3
Steam	boiling SO ₂	steel	none	6 0
Steam	boiling water	steel	none	187
Hot water	warm water	enameled C.I.	none	70
Cold water	cold water	enameled C.I.	none	4 3
Ice water	cold water	stoneware	agitated	7
Ice water	cold water	stoneware	none	5
Brine, low	nitration		35-58 rpm	32-60
velocity	slurry			
Water	sodium alcoholate	"Frederking"	agitated,	80
e .	solution	(cast-incoil)	baffled	
Steam	evaporating water	copper	-	381
Steam	evaporating	enamelware	-	36.7
e /	water			140
Steam	water	copper	none	148
Steam	water	copper	simple	244
Steam	hoiling water	copper	none	250
Steam	paraffin wax	copper	none	27 4
Steam	naraffin wax	cast iron	scraper	107
Water	paraffin wax	copper	none	24.4
Water	paraffin wax	cast iron	scraper	72.3
Steam	solution	cast iron	double	175-210
			scrapers	
Steam	slurry	cast iron	double	160-175
			scrapers	
Steam	paste	cast iron	double	125-150
			scrapers	
Steam	lumpy mass	cast iron	double	75-96
			scrapers	
Steam	powder (5%	cast iron	double	41-51
	moisture)		scrapers	

 TABLE 17.10. Overall Heat Transfer Coefficients in Agitated Tanks

 [U Btu/(hr)(sqft)(°F)]

(LIGHTNIN Technology Seminar, Mixing Equipment Co., 1982).

particle size distribution, heat transfer, multistaging, and possibly other factors.

process the solid remains in the reactor and need not circulate through a heating zone.

CIRCULATING GAS OR SOLIDS

High temperatures are generated by direct or indirect contact with combustion gases. A circulating bed of granular solids heated in this way has been used for the fixation of nitrogen from air in the range of 2300°C. Pebble heaters originally were developed as pyrolysis reactors to make ethylene, but are no longer used for this purpose. Pebbles are 5–10 mm dia, temperatures of 1700°C are readily attained, heat fluxes are in the vicinity of 15,000 Btu/(hr)(°F)(cuft of pebbles) and contact times are fractions of a second. These characteristics should be borne in mind for new processes, although there are no current examples. Induction heating of fluidized particles has been used to transfer heat to a reacting fluid; in this

FIXED BED SOLID CATALYSIS

This kind of process is used when the catalyst maintains its activity sufficiently long, for several months or a year or two as in the cases of some catalytic reforming or ammonia synthesis processes. A few processes have operated on cycles of reaction and regeneration of less than an hour or a few hours. Cycle timers on automatic valves make such operations completely automatic. A minimum of three vessels usually is needed: One on-stream, one being regenerated, and the last being purged and prepared for the next cycle. Adsorption processes are conducted this way. The original Houdry cracking process employed 10 min on-stream. One catalytic reforming process employs seven or so reactors with one of them

				Overall	U*
Jacket f	luid	Fluid in vessel	Wall material	Btu/(h·ft ² ·°F)	$J/(m^2 \cdot s \cdot K)$
Steam		Water	stainless steel	150–300	850-1700
Steam		Aqueous solution	Stainless steel	80–200	450-1140
Steam		Organics	Stainless steel	50–150	285- 850
Steam		Light oil	Stainless steel	60–160	340- 910
Steam		Heavy oil	stainless steel	10- 50	57- 285
Brine		Water	stainless steel	40–180	230 1625
Brine		Aqueous solution	Stainless steel	35–150	200 850
Brine		Organics	stainless steel	30–120	170 680
Brine		Light oil	Stainless steel	35–130	200 740
Brine		Heavy oil	Stainless steel	Io- 30	57 170
Heat-transfer	oil	Water	Stainless steel	50–200	285-1140
Heat-transfer	oil	Aqueous solution	Stainless steel	40–170	230- 965
Heat-transfer	oil	Organics	Stainless steel	30–120	170- 680
Heat-transfer	oil	Light oil	Stainless steel	35-130	200- 740
Heat-transfer	oil	Heavy oil	Stainless steel	10–40	57- 230
Steam		Water	class-lined CS	70-100	400- 570
Steam		Aqueous solution	Glass-lined CS	50- 85	285- 480
Steam		Organics	Glass-lined CS	30- 70	170- 400
Steam		Light oil	Glass-lined CS	40- 75	230- 425
Steam		Heavy oil	Glass-lined CS	10- 40	57- 230
Brine		Water	Glass-lined CS	30- 80	170- 450
Brine		Aqueous solution	Glass-lined CS	25- 70	140- 400
Brine		Organics	Glass-lined CS	20-60	115- 340
Brine		Light oil	Class-lined CS	25- 65	140- 370
Brine		Heavy oil	Glass-lined CS	10- 30	57- 170
Heat-transfer	oil	Water	Class-lined CS	30- 80	170- 450
Heat-transfer	oil	Aqueous solution	Glass-lined CS	25- 70	140- 400
Heat-transfer	oil	Organics	Glass-lined CS	25- 65	140- 370
Heat-transfer	oil	Light oil	Glass-lined CS	20- 70	115- 400
Heat-transfer	oil	Heavy oil	Glass-lined CS	lo- 35	57- 200

'Values listed are for moderate nonproximity agitation. CS = carbon steel.

(Perry's Chemical Engineers Handbook, McGraw-Hill, New York, 1984).

down every week. Regeneration usually is done in place, but eventually the catalyst must be removed and replaced. Platinum and other precious metals are recovered from the catalyst carriers in the factory.

A granular catalyst sometimes serves simultaneously as tower packing for reaction and separation of the participants by

distillation, particularly when the process is reversible and removal of the product is necessary for complete conversion to take place. This is the case of the reaction of methanol and isobutene to make methyl tertiary-butyl ether (MTBE) in the presence of granular acid ion exchange resin catalyst. MTBE is drawn off the bottom of the tower and excess methanol off the top. Such a process is applicable

TABLE 17.12. Overall Heat Transfer Coefficients with Immersed Coils	[L	expressed in	Btu/	(h.	ft ² '	°F)]
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The of coil	Coil spacing, in.	Fluid in coil	Fluid in vessel	Temp. Irange, °F.	Ut without cement	U with heat- ransfer cement
$\frac{3}{8}$ in. o.d. copper tubing attached	2	5 to 50 lb./sq, in. gage	Water under light agitation	158-210	1-5	42-46
with bands at W-in. spacing	31/8	steam	0.00	158-210	1-5	50-53
	61/4			158-210	1-5	60-64
	12 ¹ / ₂ or greater			158-210	1-5	69-72
³ / ₈ in. o.d. copper tubing attached	2	50 lb./sq. in. gage steam	No. 6 fuel oil under light	158-258	1-5	20-30
with bands at 24-in. spacing	31/8		agitation	158-258	1-5	25-38
	6¼		-	158-240	1-5	30-40
	$12\frac{1}{2}$ or greater			158-238	1-5	3546
Panel coils		50 lb./sq. in. gage steam	Boiling water	212	29	48-54
		Water	Water	158-212	8-30	19-48
		Water	No. 6 fuel oil	228-278	6-15	24-56
			Water	130-150	7	15
			No. 6 fuel oil	130-150	4	9-19
*Data courtesy of Thermon Manu	acturing Co.		•			

*Data courtesy of Thermon Manufacturing Co. †External surface of tubing or side of panel coil facing tank.

TABLE 17.13. Summ	ary of Heat-Transfer	Coefficients on	the Agitated Side
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	General Equation : $\frac{h(L)}{\lambda_{f}} = \alpha \left(\frac{\rho N D_{I}^{2}}{\mu}\right)^{m} \left(\frac{c_{p} \mu}{\lambda_{f}}\right)^{b} \left(\frac{\mu_{b}}{\mu_{w}}\right)^{c} \text{(other terms)}$									
Agitator Type	Transfer Surface	Approx. Reynolds Number Rang	e	L a	m	b	с	Other Terms	Additional Comments	Ref.
Turbine 6-blade, flat (baffled)	jacket	10–10 ⁵	D	0.13	0.65	0.33	0.24	-	Use for standard configuration. See p . 357	1,25
	coil	400-1.5 x 10 ⁶	d _e ,	0.17	0.67	0.37	See Note	$\left(\frac{D_{\rm l}}{D}\right)^{0.1} \left(\frac{d_{\rm ct}}{D}\right)^{0.5}$	for details See Note 2. Applies for standard configuration with $D_c/D = 0.7$ and	29
	vertical baffle- type	103-2 x 10 ⁶	d _{ct}	0.09	0.65	0.33	0.4	$\left(\frac{D_{\rm I}}{D}\right)^{0.33} \left(\frac{2}{n_{\rm bv}}\right)^{0.2}$	$S_{c}/d_{ct} = 2 - 4; Z_{c}/D = 0.15$	32
6-blade, retreating blade (curved blade)	jacket	10 ³ -10 ⁶	D	0.68	0.67	0.33	0.14		Revised. See Note 3	27,30
no baffles	coil	10 ³ -10 ⁶	d _{ct}	1.40	0.62	0.33	0.14		Revised	27.30
6-blade , 45" pitched	jacket	20-200	D	0.44″	0.67	0.33	0.24		Baffles have no effect in Reynolds number range studied in 12-in diameter vessel	33
3-blade retreating	jacket	2 x 10 ⁴ -2 x 10 ⁶	D	0.37^b 0	.67 ().33().14		For glass-lined vessels with finger-type baffle	27
Propeller	jacket	2 x 10 ³	D	0.54	0.67	0.25	0.14	_	Limited data, but a large. 5 ft diameter tank used, marine-type impeller used at 45" pitch and located at the midpoint of tank. No baffles used	34
Paddle	jacket	600-5 × 10 ⁵	D	0.112	0.75	0.44	0.25	$\left(\frac{D}{D_{\rm I}}\right)^{0.40} \left(\frac{w_{\rm I}}{D_{\rm I}}\right)^{0.13}$		26.35
	coil	3 x 10 ² -	dcı	0.87	0.62	0.33	0.14			37
Anchor	jacket	2.6 x 10° 10-300 <i>30040,000</i>	D D	1.0 0.36	0.5 <i>0.67</i>	0.33 <i>0.33</i>	0.18 0.18	-		34,36 34,36

Notes.

1. $\ln c = -0.202 \text{ In } \mu - 0.357$, with μ in cp.

2. For unbaffled case with coils use 0.65 of h calculated for **baffled** case (29).

3. With baffles and $N_{Re} < 400$ use value calculated. In fully developed turbulent region baffles increase calculated h by approximately 37%(1)

New nomenclature: d_{et} is outside tube diameter of coil, D_e is coil diameter, n_{bv} is number of vertical baffle-type coils, S_e is coil spacing, w_1 is impeller blade width, and Z_e is height of coil from tank bottom.

^a For impelkr 4½-in from bottom, 0.535 for impelkr 1 1-in from bottom

• For steel impeller, 0.33 for glassed-steel impeller

(Rase, 1977, Vol. 1).

TABLE 17.14. Equations for Heat Transfer Coefficients inside Stirred Tanks^a

1. To jackets, with paddles, axial flow, and flat blade turbines^{1,6,7}

$$h_{0}(\text{jacket}) \frac{T}{k} = 0.85 \left(\frac{D^{2} N \rho}{\mu}\right)^{0.66} \left(\frac{C_{\rho \mu}}{k}\right)^{0.33} \\ \times \left(\frac{\mu}{\mu_{s}}\right)^{0.14} \left(\frac{Z}{T}\right)^{-0.56} \left(\frac{D}{T}\right)^{0.13}$$

2. To helical coils^{3,5}

$$h_{0}(\text{coil}) \frac{D}{k} = 0.17 \left(\frac{D^{2} N \rho}{\mu}\right)^{0.67} \left(\frac{C_{p} \mu}{k}\right)^{0.3'} \\ \times \left(\frac{D}{T}\right)^{0.1} \left(\frac{d}{T}\right)^{0.5} \left(\frac{\mu}{\mu_{s}}\right)^{m} \\ \text{m} = 0.714/\mu^{0.21}, \mu \text{ in } cP$$

3. To vertical tubes²

$$\begin{split} h_{0}(\text{tubes}) \frac{D}{K} &= 0.09 \Big(\frac{D^{2} N \rho}{\mu} \Big)^{0.65} \Big(\frac{C_{\rho} \mu}{k} \Big)^{0.3} \\ &\times \Big(\frac{D}{7} \Big)^{0.33} \Big(\frac{2}{n_{b}} \Big)^{0.2} \Big(\frac{\mu}{\mu_{s}} \Big)^{0.14} \end{split}$$

4. To plate coils⁴

$$\begin{split} h_0(\text{plate coil}) & \frac{L}{K} = 0.1788 \Big(\frac{ND^2\rho}{\mu}\Big)^{0.448} \Big(\frac{C_{D}\mu}{k}\Big)^{0.33} \Big(\frac{\mu}{\mu_f}\Big)^{0.50} & \text{for } N_{\text{Re}} < 1.4 \times 10^3 \\ h_0(\text{plate coil}) & \frac{L}{K} = 0.0317 \Big(\frac{ND^2\rho}{\mu}\Big)^{0.658} \Big(\frac{C_{D}\mu}{k}\Big)^{0.33} \Big(\frac{\mu}{\mu_f}\Big)^{0.50} & \text{for } N_{\text{Re}} > 4 \times 10^3 \end{split}$$

^{*b*} Nomenclature: *d* = tube diameter, *D* = impeller diameter, *L* = plate coil height, *N* = impeller rotational speed, n_b = number of baffles or of vertical tubes acting as baffles, T = tank diameter, *Z* = liquid height.

REFERENCES

- 1. G. Brooks and G.-J. Su, Chem. Eng. Prog. 54, (Oct. 1959).
- I.R. Dunlap and J.H. Rushton, Chem. Eng. Prog. Symp. Ser. 49(5), 137 (1953).
- 3. J.Y. Oldshue and A.T. Gretton, Chem. Eng Prog. 50(12) 615 (1954).
- 4. D.K. Petree and W.M. Small, AIChE Symp. Series. 74(174), (1978).
- A.H.P. Skelland, W.K. Blake, J.W. Dabrowski, J.A. Ulrich, and T.F. Mach, AIChE. J. 11(9), (1965).
- 6. F. Strek, Int. Chem. Eng. 5, 533 (1963).
- 7. V.W. Uhl and J.B. Gray, *Mixing Theory and Practice*, Academic, New York, 1966, Vol. 1.

(Recommended by Oldshue, 1983).

Figure 17.36. Effective thermal conductivity and wall heat transfer coefficient of packed beds. Re' = $d_p G/\mu$, $d_p = 6V_p/A_p$, $\varepsilon =$ porosity. (a) Effective thermal conductivity in terms of particle Reynolds number. Most of the investigations were with air of approx. $k'_f = 0.026$, so that in general $k'_e/k'_f = 38.5k'_e$ [Froment, Adv. Chem. Ser. 109, (1970)]. (b) Heat transfer coefficient at the wall. Recommendations for L/d_p above 50 by Doraiswamy and Sharma are line H for cylinders, line J for spheres. (c) Correlation of Gnielinski (cited by Schlünder, 1978) of coefficient of heat transfer between particle and fluid. The wall coefficient may be taken as $h_w = 0.8h_p$.



		Heating.		Particles		
Authors	Method of Measurement	or Cooling o Gas	Gas	Material	Shape	Diameter D_p , mm
Bakhurov and Bores- kov (1947)	Radial temper- ature and con- centration pro- files	С	Air	Glass, porce- lain, metals, etc.	Spheres, rings, cylinders, granules	3-19
Brötz (1951)	,,	Н	N ₂ , CO ₂ , H ₂	Glass, catalyst	Spheres, granules	2-10
Bunnell, Irvin, Olson, and Smith (1949)	Radial temper ature profiles	- C	Air	Alumina	Cylinders	3
Campbell and Hunt - ington (1952)	,,	H, C	Air, natural gas (82 % CH ₄)	Glass, alumina, aluminum	Spheres, cylinders	5-25
Coberly and Marshall (1951)	"	н Н	Air	Celite	Cylinders	3-12
⊾ ^r aeda (1952)	37	С	Air	Catalyst	Cylinders	3-10
Quinton and Storrow (1956)	,,	Н	Air	Glass	Spheres	4.4
Aerov and Umnik (1951b,c)	Packed bed heat exchanger. Single radial temperature	C	Air, CO ₂ , H ₂	Glass, cata- lyst, porce- lain, sand	Spheres, tablets, rings	0.4-10
Hougen and Piret (1951) Molino and Hougen (1952)	Packed bed heat exchanger	С Н	Air	Celite	Spheres, cylinders	2-12
Kling (1938b)	,,	Н	Air	Steel, glass	Spheres	3-4
Verschoor and Schuit (1960)	,,	Н	Air, H ₂	Lead, glass, etc.	Spheres, cylinders, granules	3-10
Bernard and Wilhelm (1950)	Mass diffusion		(Water)	Glass, lead, alumina, etc.	Spheres, cylinders, granulos	18

TABLE 17.15. Data for the Effective Thermal Conductivity, *K*, (kcal/mh°C), and the Tube Wall Film Coefficient, *h*_w (kcal/m²h°C), in Packed Beds*

when the reaction can be conducted satisfactorily at boiling temperatures; these can be adjusted by pressure.

A variety of provisions for temperature control of fixed beds is described in Section 17.6 and following.

- a. Single beds are used when the thermal effects are small. Jacketed walls usually are inadequate for heat transfer to beds, but embedded heat transfer **tubes sometimes are used**.
- **b.** Multitubular units with catalyst in tubes and heat transfer medium on the shell side are popular. A reactor for making phosgene from carbon monoxide and chloride has 2-in, dia tubes

8 ft long filled with activated carbon catalyst and cooling water on the shell side.

- c. **Multibed** units with built-in interstage heat transfer surface. Economical when the amount of surface is not large. In comparison with type (d), this design may have more difficult maintenance, less flexibility and higher cost because of the shortness of the tubes that may have to be used. The Sper-Rashka converter for SO, oxidation has three beds and three large internal exchangers in a single shell (Ullmann, 3rd ed., 1964, Vol. 15, p. 456).
- d. Multibed units with external heat exchangers. Several variations

ĺ	Tube		N	Range of		
	Diameter, <i>Dt</i> , mm	Length L, m	$(D_p \text{ and } D_t \text{ in } \mathbf{m})$	${}^{N}\mathrm{Re'}$ or other variable	Remarks	
	00 square	. 1	$N_{\rm Pe}' = 2.4$ to 6.3	a ~ 1000 kg/m ² h		
	100	0.2-0.4	Graphs in paper	Gas velocity : 2500-7200 m/h	Special apparatus at low velocities	
	50	0.05-0.2	$\frac{K_r}{k_g} = 5.0 + 0.061 \left(\frac{D_p G}{\mu}\right)$	30-100		
	50-100		$\frac{K_r}{k_g} = 10.0 + 0.267 \left(\frac{G}{\mu a_v}\right); h_w = 2.07 \left(\frac{G}{\mu a_v}\right)^{0.47}$	20-500		
	127	up to 1	$K_r = 0.27 + 0.00146 \frac{\sqrt[]{a_p G}}{\mu}$; $h_w = 8.51 \ G^{0.33}$	G (kg/m²h): 850-6000		
	25-100	0.1-0.4	$\left(5.5 \pm 0.05 \left(\frac{D_p G}{\mu} \right) \right)$	O-30	See also Hatta	
			$\frac{K_r}{k} = \begin{cases} 1.72 \left(\frac{D_p G}{\mu}\right)^{0.41} \end{cases}$	30-100	a, b, 1949), Maeda (1950), and Maeda	
			$\left(\begin{array}{c} \lambda_g \\ 0.209 \left(\frac{D_p G}{\mu}\right)^{0.87} \end{array}\right)$	1001000	and Kawazoe (1953)	
	41	0.75	$K_r = 0.36 + 0.00162 \left(rac{D_p G}{\mu} ight)$; h_w = 0.04 a	30-1100 500 <i><</i> G <i><</i> 17000		
	6 5	0.3	$\frac{K_r}{k_g} = 10.5 + 0.076 N_{\rm Pr} \left(\frac{4G}{\mu a_v}\right)$	1 O-3500	For rings see paper	
		ł	$\frac{h_w D_p}{k_g} = 0.155 N_{\rm Pr}^{\frac{1}{6}} \left(\frac{4G}{\mu a_v}\right)^{0.75}$	150-4000		
	35-95	0.16-0.32	$\frac{K_r}{k_g} = 1.23 \left(\frac{\sqrt{a_p}G}{\mu}\right)^{0.43}$ No correction for h_w	100-3000	The first paper gives a slightly different formula for cooling	
	60	0.3	See graph in paper. No correction for h_w	400-3500		
	30-50	0.2-0.3	$\frac{K_r}{k_g} = \frac{K_{r, 0}}{k_g} + 0.10(a_v D_t)^{0.50} \left(\frac{G}{\mu a_v}\right)^{0.69}$	10-1000	No correction for h_w	
	50		Graph in paper of N_{Pe} ' vs. N_{Re} '	5-2400	Diffusion of meth- ylene blue in water	

TABLE 17.15—(continued)

"a, = surface of a particle, a, = surface/volume in the bed. (Kjaer, Measurement and Calculation of Temperature and Conversion in Fixed-Bed Catalytic Converters, Haldor Topsoe, Copenhagen, 1958).

of this design with steam generation or feed gas as means of cooling are used for the catalytic oxidation of SO,.

- e. **Multibed** unit with interstage injection of temperature controlled process fluid or inert fluid for temperature control of the process. In the synthesis of cumene from propylene and benzene in the presence of supported phosphoric acid catalyst, interstage injection of cold process gas and water is used for temperature control and maintenance of catalyst activity (Figure 13.18(g)).
- f. Autothermal multitubular unit with heat interchange between feed on the shell side and reacting gas in the packed tubes and between feed and reacted gas in an external or built-in

exchanger. Many complex variations of this design have been or are being used for ammonia synthesis.

g. **Multibed** units in individual shells with interstage heat transfer. From three to seven stages are adopted by different processes for the catalytic reforming of naphthas to gasoline.

FLUIDIZED BED CATALYSIS

Such processes may be conducted to take advantage of the substantial degree of uniformity of temperature and composition and high rates of heat transfer to embedded surfaces. Orthophthalic

	Mathad af	Heating	=		Particles	
Authors	Method of Measurement	or Cooling of Gas		Material	Shape	$Diameter D_p, mm$
Campbell and Hunt ington (1952)	Packed bed hea exchanger	н, с	Air, natural gas (82 % CH ₄)	<pre>#lass, alumina aluminum</pre>	Spheres, cylinders	5-25
Chu and Storrow (1952	"	Н	Air	Glass, steel, lead, Socony - Vacuum cata- lyst beads	Spheres	1-6
Colburn (1931)	,,	Н	Air	Porcelain, zinc, etc.	Spheres, granules	5-25
Kling (193813)	,,	Н	Air	Steel, glass	Spheres	3-4
Leva (1947)	"	Н	Air, CO,	Glass, clay,	Spheres	3-13
Leva and Grummer (1948)	99	Н	Air	Glass, clay, metals, etc.	Spheres, cylinders, granules, etc.	2-25
Leva, Weintraub, Grummer, and Clark (1948)	,,	C -	Air, CO,	Glass, porce- lain	Spheres	3-13
Leva (1950)	,,	Н	Air	Glass, clay, porcelain, metal	Spheres, rings cylinders	4-18
Maeda (1952)	"	С	Air	Catalyst	Cylinders	3-10
Maeda and Kawazoe (1953)	"	С	Air		Granules, rings, saddles	3-25
Verschoor and Schuit (1950)		Н	Air, H ₂	Lead, glass, etc.	Spheres, cylinders, granules	3–10
Tasker (1946)	Phthalic anhy - dride synthesis	C	Air	Catalyst on quartz (?)	Granules	1.7-2.0

TABLE 17.16. Data for the Overall Heat Transfer Coefficient, u (kcal/m²h^oC), in Packed Beds

anhydride is made by oxidation of naphthalene in a fluidized bed of V_2O_5 deposited on silica gel with a size range of 0.1-0.3 mm with a contact time of 10–20 sec at 350–380°C. Heat of reaction is removed by generation of steam in embedded coils. No continuous regeneration of catalyst is needed. Acrylonitrile and ethylene dichloride also are made under conditions without the need for catalyst regeneration.

From the standpoint of daily capacity, the greatest application

of fluidized bed catalysis is to the cracking of petroleum fractions into the gasoline range. In this process the catalyst deactivates in a few minutes, so that advantage is taken of the mobility of **fluid**ized catalyst to transport it continuously between reaction and regeneration zones in order to maintain its activity; some catalyst also must be bled off continuously to maintain permanent poisons such as heavy metal deposits at an acceptable level.

Several configurations of reactor and regenerator have been in

TABLE 17.16-(continued)

Tu	be	Formula	Range of	
Diameter D_t, mm	Length L. m	$(D_p \text{ and } D_t \text{ in } \mathbf{m})$	N_{Re}' o∎other variable	Remarks
50-150		$\frac{U}{GC_p} = 0.76 \ e^{-0.0225 \ a_v D_t} \left(\frac{G}{\mu a_v}\right)^{-0.42}$	30-1000	U refers to tube axis temperature
25	0.3-1.2	$\frac{UD_t}{k_g} = 0.134 \left(\frac{D_p}{D_t}\right)^{-1.13} \left(\frac{L}{D_t}\right)^{-0.90} \left(\frac{D_p G}{\mu}\right)^{1.17}$	$\frac{D_t G}{\mu} < 1600$	
		$\frac{UD_t}{k_g} = 15 \left(\frac{D_p}{D_t}\right)^{-0.90} \left(\frac{L}{D_t}\right)^{-1.82} \left(\frac{D_pG}{\mu}\right)^n$	1600 <	
		$n = 0.55 \left(\frac{L}{D_t}\right)^{0.165}$	$\left[\frac{D_t G}{\mu} < 3500\right]$	
35-80	0.51.2	$U = f\left(rac{D_p}{D_t} ight) G^{0.83}$	Range of G (kg/m^{\$}h) : 4500–45000	Function f given in paper . Maxi- mum 0.045 for $\frac{D_p}{D_l} = 0.15$
	0.3	Graph given in paper	400-3500	
15-52	0.3-0.9	$\mu_p / D Q 0.90$	50-3500	
21-52	0.3-0.9	$\frac{\partial D_t}{k_g} = 0.813 e^{-6} \overline{D_t} \left(\frac{D_p \sigma}{\mu} \right)$	100-4500	Correction factor used for metallic packings
21-52	0.3-0.9	$\frac{UD_t}{k_g} = 3.50 \ e^{-4.6 \frac{D_p}{D_t}} \left(\frac{D_p G}{\mu}\right)^{0.7}$	150-3000	
lb-52	0.3–0.9	$\frac{UD_p}{k_g} = 0.125 \left(\frac{D_p G}{\mu}\right)^{0.75}$	500–12000	Correlation valid for high values of $\frac{D_p}{D_t}$
25-100	0.1-0.4	$\frac{UD_{t}}{k_{g}} = 4.9 e^{-2.2 \frac{D_{p}}{D_{t}}} \left(\frac{D_{p}G}{\mu}\right)^{0.60}$	100-600	
52-154		See original paper	30-900	Formula varies with shape of ma- terial
30-50	0.2–0.3	$\begin{aligned} \frac{UD_{t}}{k_{g}} &= 5.783 \frac{K_{r, 0}}{k_{g}} + 0.085 \left(\frac{D_{p}}{D_{t}}\right)^{-0.50} \left(\frac{D_{p}G}{\mu}\right)^{0.69} \\ &+ 0.066 \left(\frac{D_{t}}{L}\right) \left(\frac{D_{p}}{D_{t}}\right)^{-1} \left(\frac{D_{p}G}{\mu}\right) \end{aligned}$	40-4000	K_{F,0} is thermal conductivity of bed with stagnant gas
38	0.4-0.7	$U^{-1} = 0.00123 + 0.54 G^{-0.83}$	Range of G (kg/m [®] h): 3000-12000	U refers to tube axis temperature and is corrected for radiation

use, two of which are illustrated in Figures 17.31(a) and (b). Part (a) shows the original arrangement with separate vessels side by side for the two operations. The steam stripper is for removal of occluded oil from the catalyst before it is burned. In other designs the two vessels are in vertical line, often in a single shell with a partition. Part (b) is the most recent design of transfer line cracking

which employs highly active zeolite catalysts that are effective at short contact times. The upper vessel is primarily a catalyst disengaging zone. A substantial gradient develops in the transfer line and results in an improvement in product distribution compared with that from mixed reactors such as part (a).

Hundreds of fluidized bed crackers are in operation. The

		Heating		-	Particles	
Authors	Method of Measurement	Or Cooling of Gas	Gas	Material	Shape	$\substack{ \text{Diameter,} \\ D_p, \text{ mm} }$
Furnas (1930a, b, c, 1932)	Unsteady heat transfer	H, C	Air, flue gas	Iron ore, lime- stone, coke, etc.	Granules	4-70
Löf and Hawley (1948)	- 91	С	Air	Granitic gravel	Granules	8-34
Saunders and Ford (1940)	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	C	Air	Steel, lead, giass	Spheres	1.6-6.4
Tsukhanova and Sha- patina (1943) Chukhanov and Sha- patina (1946)	. ,,	C	Air	Steel, cha- motte, copper	Spheres, cylinders, granules	2-7
Dayton et al. (1962)	Cyclic varia- tions		Air	Glass	Spheres	3-6
Glaser (1966)	,, ,,		Air	Stoneware	Raschig rings	5–17
Gamson, Thodos, and Hougen (1943)	Drying		Air	Porous celite	Spheres, cylinders	2-19
Wilke and Hougen (1946)	- 99		Air	Porous celite	Cylinders	2-19
Taecker and Hougen (1949)	33		Air	Porous clay- kieselguhr	Raschig rings, Berl saddles	6–50
Eichhom and White (1962)	Dielectrical heating		Air	Plastic	Spheres	0.1-0.7
Satterfield and Res- nick (1964)	Decomposition of H₂O₂		Vapors of $I_{1_{2}0}$ and $\mathbf{H}_{2}0_{2}$	Catalyst	Spheres	6

TABLE 17.17. Heat Transfer Coefficient between Particle and Gas

vessels are large, as much as 10 m or so in diameter and perhaps twice as high. Such high linear velocities of vapors are maintained that the entire catalyst content of the vessels circulates through the cyclone collectors in an hour or so. Electrical precipitators after the cyclone collectors have been found unnecessary.

Two other fluidized bed petroleum reactors are illustrated as Figures 17.31(c) and (d) and several nonpetroleum applications in Figure 17.32.

GAS-LIQUID REACTIONS WITH SOLID CATALYSTS

The number of commercial processes of this type is substantial. A list of 74 is given by Shah (1979). A briefer list arranged according to the kind of reactor is in Table 17.21. Depending on the

circumstances, however, it should be noted that some reactions are conducted industrially in more than one kind of reactor.

Leading characteristics of five main kinds of reactors are described following. Stirred tanks, fixed beds, slurries, and three-phase fluidized beds are used. Catalyst particle sizes are a compromise between pressure drop, ease of separation from the fluids, and ease of fluidization. For particles above about 0.04 mm dia, diffusion of liquid into the pores and, consequently, accessibility of the internal surface of the catalyst have a minor effect on the overall conversion rate, so that catalysts with small specific surfaces, of the order of $1 \text{ m}^2/g$, are adequate with liquid systems. Except in trickle beds the gas phase is the discontinuous one. Except in some operations of bubble towers, the catalyst remains in the vessel, although minor amounts of catalyst entrainment may occur.

Tube		ıbe		Range of		
	Díameter, D_t , mm	Length L. m	$(D_p \text{ in } m)$	NRe' or other variable	Remarks	
	150–230	0.5-1	$h_v = A \frac{G^{0.7}}{D_p^{(0.9)}} T^{0.3} 10^{1.68e - 3.56e^2}$	Range of G (kg/m ² h): 2300-9200	A given in paper. Corrections for temperature and voids not very re- liable	
	ca. 300 square	0.9	$h_v = 1.82 \left(rac{G}{D_p} ight)^{0.7}$	G: 300-I 600		
	50200	0.09-0.34	Graphs in paper	a: 2670-5340	Correlated by Löf and Hawley (1948) as: h,, = 0.152;	
		0.01-0.25	Graphs and $\frac{hD_p}{k_g} = 0.24 \left(\frac{D_p G}{\mu}\right)^{0.83}$	Gas velocity : 0.7-Z m/sec.		
		- 0.05	Graphs in paper	100-1000		
i.	350		$\frac{hD_p}{k_g} = A \left(\frac{D_p G}{\mu}\right)^{0.61}$	130-2000	A varies from 0.590t00.713. See also Glaser (1938)	
			$j_h = rac{h}{GC_p} N_{ m Pr}^{-\frac{3}{2}} = 1.064 \left(rac{D_p G}{\mu} ight)^{-0.41}$	350-4000		
			$j_h = rac{h}{GC_p} N_{\mathrm{Pr}}^{-\frac{2}{3}} = 1.96 \left(rac{D_p G}{\mu} ight)^{-0.51}$	50-350		
			Rings : $j_h = 1.148 \left(\frac{\sqrt{a_p G}}{\mu}\right)^{-0.41}$	100-20000		
_	_		Saddles: $j_h = 0.920 \left(\frac{\sqrt{a_p G}}{\mu}\right)^{-0.34}$	70-3000		
	38 square		Granns in paper	1-18		
	47-75	0.024	$j_h = \overline{0.992 \left(\frac{D_p G}{\mu}\right)^{-0.34}}$	15-160		

TABLE 17.17—(continued)

 ${}^{a}a_{D}$ = Surfac3 of a particle, a, = surface/volume in the bad. (Kjaer, Measurement and Calculation of Temperature and Conversion in Fixed-Bed Catalytic Converters, Haldor Topsoe, Copenhagen, 1958).

1. Stirred tanks with suspended catalyst are used both in batch and continuously. Hydrogenation of fats or oils with Raney nickel or of caprolactam usually are in batch. Continuous processes include **some** hydrogenations of fats, some fermentation processes with cellular enzymes and air and the hydrogenation of nitrogen monoxide to hydroxylamine. The gas is distributed with spargers or introduced at the eye of a high-speed impeller in a draft tube. Internal recirculation of the gas also is practiced. The power input depends on the settling tendency of the particles and the required intimacy of gas-liquid mixing. It is greater than in the absence of solids; for example, the solid catalyzed hydrogenation of nitrogen monoxide employs a power input of about 10 kW/m³ (51 HP/ 1000 gal) compared with 5-10 HP/1000 gal for ordinary liquid-liquid mixing.

2. In ebullated (liquid fluidized) beds the particles are much larger (0.2-1 mm) than in gas fluidization (O-O.1 mm). Little

expansion of the bed occurs beyond that at minimum fluidization, so that the bed density is essentially the same as that of the fixed bed. Because substantial internal circulation of the liquid is needed to maintain fluidization, the fluids throughout the reactor are substantially uniform. In the hydrodesulfurization and hydrocracking of petroleum fractions and residua at 100 atm and 400°C, a temperature variation of only 2°C or so obtains in the reactor.

3. Slurry reactors (bubble towers) are fluidized with continuous flow of gas. The particles are smaller (less than 0.1 mm) than in the liquid fluidized systems (0.2-1 mm). In some operations the liquid and solid phases are stationary, but in others they circulate through the vessel. Such equipment has been used in Fischer-Tropsch plants and for hydrogenation of fatty esters to alcohols, **furfural** to **furfuryl** alcohol, and of glucose to sorbitol. Hydrogenation of benzene to cyclohexane is done at 50 bar and 220–225°C with Raney nickel of 0.01-0.1 mmdia. The relations between gas velocities, solids

TABLE 17.18. Formulas for the Heat Transfer	Coefficient at the Walls of Packed Vessels
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	Name	Geometry	Formula
1.	Beek (1962)	spheres	Nu = 0.203 Re ^{1/3} Pr ^{1/3} + 0.220 Re ^{0.8} Pr ^{0.4} , Re < 40
2.	Beek (1962)	cylinders	Nu = 2.58 Re ^{1/3} Pr ^{1/3} + 0.094 Re ^{0.8} Pr ^{0.4} , Re $<$ 40
3	. Yagi-Wakao (<i>Chem. Eng.</i> <i>Eng.</i> Sci. 5, 79, 1959)	spheres	$Nu = 0.186 Re^{0.8}$
4.	Hanratty (<i>Chem.</i> Eng. Sci. 3, 209, 1954)	cylinders	$Nu = 0.95 Re^{0.5}$
5.	Hawthorn (AIChE J		Nu = 0.28 $\text{Re}^{0.77} \text{Pr}^{0.4}$
	14, 69 1968)		
6.	Doraiswamy and	spheres	Nu = 0.17 Re ^{0.79} , $L/d_t > 50$, $20 < \text{Re} < 7600$, $0.05 < d_p/d_t < 0.30$
	Sharma (1984)	-	
7.	Doraiswamy and Sharma (1984)	cylinders	Nu = 0.16 Re ^{0.93} , $L/4 > 50$, $20 < \text{Re} < 800$, $0.03 < d_p/d_t < 0.02$
8.	Gnielinski-Martin,		Nu/(2.5–1.5 ε) = 0.8[2 + $F(\text{Re}/\varepsilon)^{1/2}(\text{Pr})^{1/3}]$
	Schlünder (1978)		()
			$F = 0.664 \left\{ 1 + \left[\frac{0.0577(\text{Re}/\varepsilon)^{0.3} \text{Pr}^{0.67}}{1 + 2.44(\text{Pr}^{2/3} - 1)(\text{Re}/\varepsilon)^{-0.1}} \right]^{Z} \right\}^{1/2}$
9	. Gnielinski–Martin–		Nu na
	Schlünder (1978)		$\ln \frac{1}{2.5 - 1.5\varepsilon} \approx 0.750 + 0.1061 \ln(\text{Re}/\varepsilon) + 0.0281[\ln(\text{Re}/\varepsilon)]^2$

*Definitions: Nu = $h_w d_p / k_r$, Pr = $(C_p \mu / k)_r$, h_w = wall coefficient, d_p = particle diameter = $6V_p / A_p$, k, = fluid molecular conductivity, ε = porosity, Re = $d_p G / \mu$, G = superficial mass velocity per unit cross section.



Figure 17.37. Some measured and predicted values of heat transfer coefficients in fluidized beds. 1 Btu/hr(sqft)(°F) = 4.88 kcal/(hr)(m²)(°C) = 5.678 W/(m))(°C). (a) C o mp arison of correlations for heat transfer from silica sand with particle size 0.15 mm dia fluidized in air. Conditions are identified in Table 17.19 (*Leva*, 1959). (b) Wall heat transfer coefficients as function of the superficial fluid velocity, data of Varygin and Martyushin. Particle sizes in microns: (1) ferrosilicon, d = 82.5; (2) hematite, d = 173; (3) Carborundum, d = 137; (4) quartz sand, d = 140; (5) quartz sand, d = 198; (6) quartz sand, d = 216; (7) quartz sand, d = 428; (8) quartz sand, d = 51.5; (9) quartz sand, d = 650; (10) quartz sand, d = 1110; (11) glass spheres, d = 1160. (*Zabrodsky* et al., 1976, Fig. 10.17). (c) Effect of air velocity and particle physical properties on heat transfer between a fluidized bed and a submerged coil. Mean particle diameter 0.38 mm: (I) BAV catalyst; (II) iron-chromium catalyst; (III) silica gel; (IV) quartz; (V) marble (*Zabrodsky* et al., 1976, Fig. 10.20).





concentrations, bubble sizes, and rates of heat transfer are extensively documented in the literature.

4. In trickle bed reactors the gas and liquid both flow downward through a fixed bed of catalyst. The gas phase is continuous, and the liquid also is continuous as a film on the particles. Provided that the initial distribution is good, liquid distribution remains substantially uniform at rates of $10-30 \text{ m}^3/$

 m^2 hr superficially, but **channelling** and hot spots may develop at lower rates. Redistributors sometimes are used. The many correlations that have been developed for packed bed mass transfer are applicable to trickle bed operation. Commercial reactors are 1-4 m dia and 10–30 m long. Hydrocracking and hydrodesulfurization of petroleum and hydration of olefins are commonly practiced in trickle beds at superficial liquid velocities of 3-90 m/hr.



Figure 17.38. Heat transfer coefficient in fluidized beds [Wender and Cooper, AIChE J. 4, 1.5 (1958)]. (a) Heat transfer at immersed vertical tubes. All groups are dimensionless except $k_g/C_g\rho_g$, which is sqft/hr. The constant C_R is given in terms of the fractional distance from the center of the vessel by $C_R = 1 + 3.175(r/R) - 3.188(r/R)^2$. (b) Heat transfer at the wall of a vessel. L_H is bed depth, D_T is vessel diameter.



Figure 17.38—(continued)

TABLE 17.19. Experimental Investigations of Heat Transfer in Fluidized Be

Refer- ence	Solids	Voidage range	Absolute density, lb per cu ft	Particle- sise range, ft	Type of PPtex;tus operation	Vessel dism., in.	Height of heat- transfer area, in.	Bed height in.	Fluids	Flow trange. lb / (br) (sq ft .)	Temp, °F
1	Sands, graphite, soft brick	Dense	- 83-166	- 6-14 mesh to 36-72	Steam-jacketed	1.5	14.5	 	Air	150-1,200	. <u> </u>
2	Iron powder. sands. glass beads catalyst	18.8-75	119-434	mesh 0.000198- 0.00288	Central electric	1.25 in. 5.5	4	10	Air	1 a-605	23.5-65.0
8 4	Sand, aluminum, calcium carbonate Glass beads	54-95 Dilute	169-167	0.000277- 0.000822 0.00023-	Wall electric heat Electric heat	4.0 1.959	30 12	30	Air Air	96-935 95-3,78Ю	300-450
6	Sand, aluminum. graphite, copper	phase Dense phase	24.6-27.2	0.0036 0.00079- 0.0126	from outside Central cooling	2.31	Immersed cooling		Air	40-100	87-145
9 13	catalyst Aerocat, coke, iron powder Carborundum iron	52-69 Dense	121-466	0.000363 0.000560 0.000262-	Wall steam heating Wall water cools	2.06 and 3.07	coil 23 and 26.5	2-13	Air	50-300	200-220
17	oxide, coke, lead fly ash, alloy Glass beads	phase Dense	154	0.00213	ing Internal heating	3.0	4	10	Air, Chi, Coi, town gas, H ₁ and N ₂ mixtures Air, CO ₁ , Freon-12	44-779	10-30°C
26. 29	Band. iron catalyst,	phase 35-75	80- 500	0.0011 0.000129-	by electric wire Wall steam heat	2.0 and	25 and 26	12-25	He, H ₂ , H ₂ and N ₁ mixtures Air, CO ₂ , He. N ₂	.47-1,095	258413
30		Dense phase	A	0.00149 0.000432- 0.00386	Wall cooling (air)	4.0 4.0	24		Air	50-1,100	
31	coal	1.7-86.2	43.6-180,	0.000250- 0.0142	Wall electric heating	4.0	³ sections. 2, 5, and 2 in.	lo-30	Air	79-4.350	
30 37	Glass beads, micro- spheres Glass beads	Dense phase Dilute	138-153 151-177	0.00022- 0.00027 0.000133	Small electric heater probe Internal and ex-	2.875 and	A	18-20	He, air. CH₄ , argon _i Air	lo-150 2,700	506
38 42	Silicon carbide. AlrO, silica gel	Dense phase Dense	70-243	0.00149 0.000287~ 0.000817	cooling	1.00 2.0	22	101224 6	Air. He, CO2	5.4-200	120-414
43.44	Sand, iron ore	phase	165-330	0.000179-	ing	4.75	each 5 in. high	47 68	Air	65300	Approx
50	Carborundum sand, aluminum powder, lead powder, glass	phase Dense phase	160-700	0.00197 0.00020- 0.010	Internal heating by small cylin - drical element	22.2 3.94			,Air. CO3, H2	00-000	100-400
	Dergs										

"Another list of 29 sources is given by Zabrodsky (1966). (Leva, 1959).

TABLE 17.20. Heat Transfer Coefficients in Fluidized Beds"

1. At vertical tubes [Vreedenburg, Chem. Eng. Sci. 11, 274 (1960)]:

$$[h(D - d_t)/k_a](d_t/D)^{1/3}(k_a/C_s\mu_a)^{1/2} = C[u(D - d_t)\rho_s/\mu_a]^{1/2}$$

Conditions:

$$\begin{split} \rho_s d_s u / \mu_g &< 2050, \\ \left\{ \begin{array}{l} \rho_s u (D - d_t) / \mu_g &< 2.4 \times 10^5, \ C = 2.7 \times 10^{-16}, \ n = 3.4 \\ \rho_s u (D - d_t) / \mu_g &> 2.4 \times 10^5, \ C = 2.2, \ n = 0.44 \\ \left[h (D - d_t) / k_g \right] (d_t d_s k_g / [(D - d_t) C_s \mu_g]^{1/3} &= C [u (D - d_t) g^{0.5} d_s^{1.5}]^n \end{split} \end{split} \end{split}$$

Conditions:

 $\rho_{s}d_{s}u/\mu_{q} > 2550,$ $u(D-d_t)g^{0.5}d_s^{1.5} < 1070, \quad C = 1.05 \times 10^{-4}, \quad n = 2.0$ $I u(D - d_t)g^{0.5}d_s^{1.5} > 1070, \quad C = 240,$ n = 0.8

For off-center locations, the factor C is multiplied by C_R which is

given in terms of the fractional distance from the center by

$$C_R = 1 + 3.175(r/R) - 3.188(r/R)^2$$

- 2. At vertical tubes, see the correlation of Wender and Cooper on Figure 17.17(a)
- 3. At horizontal tubes (Vreedenburg, loc. cit.; Andeen and Glicksman, ASME Paper 76-HT-67, 1976):

$$(hd_t/k_g)(k_g/C_s\mu_g)^{0.3} = 0.66[\rho_s d_t u(1-\varepsilon)/\mu_g \varepsilon]^{0.44}, \\ \rho_s d_s u/\mu_g < 2500 \\ (hd_t/k_g)(k_g/C_s\mu_g)^{0.3} = 900(1-\varepsilon)(d_t u\mu_g/d_s^3\rho_s g)^{0.326}, \\ \rho_s d_s u/\mu_g > 2550$$

4. At vessel walls, see Figure 17.17(b) for the correlation of Wender and Cooper.

^a Notation: Subscript s for solid, subscript g for gas, d_r = tube diameter, D = vessel diameter, g = acceleration of gravity.

5. Upflow fixed beds. The liquid phase is continuous and the gas phase dispersed. This mode of operation has the advantages of better mixing, higher rates of mass and heat transfer, better distribution of liquid flow across the cross section, and better scouring of deactivating deposits from the surface of the catalyst. The disadvantages relative to trickle beds are higher pressure drop,

the possibility of occurrence of flooding, and the need for mechanical restraint to prevent tluidization and entrainment of the catalyst. The most prominent example of upflow operation is the SYNTHOIL coal liquefaction process, but this mode of operation is competitive in other cases with the trickle bed, depending on the balance of advantages and disadvantages in particular situations.

TABLE 17.21. Examples of Industrial Gas-Liquid-Solid Reaction Processes

A. Fixed-bed reactors 1. Trickle beds (downflow)	b. Hydrogenation of acetone and nitrocaprolactam c. Aerated fermentation with cellular enzymes
a. Catalytic hydrodesulfurization, hydrocracking and hydrogenation b. Butynediol from acetylene and aqueous formaldehyde	d. Reaction between methanol and hydrogen chloride with $ZnCl_2$ catalyst
c. Sorbitoi from glycerol	2. Slurry towers
d. Oxidation of SO, in the presence of activated carbon	a. Fischer-Tropsch process
e. Hydrogenation of aniline to cyclohexylaniline	b. Hydrogenation of methyl styrene and carboxy acids
2. Upflow (bubble) reactors	c. Oxidation and hydration of olefins
a. Coal liquefaction by SYNTHOIL process	d. Polymerization of ethylene
b. Fischer-Tropsch process	e. Calcium hydrophosphite from white phosphorous and lime slurry
c. Selective hydrogenation of phenylacetylene and styrene	f. Lime/limestone process for removal of SO ₂ from flue gases
	3. Fluidized bed of catalyst
B. Suspended solid reactors	a. Calcium acid sulfite from $CaCO_3 + SO_1 + H_2O_2$
1. Stirred tanks	b. Coal liquefaction
a. Catalytic hydrogenation of fats and oils	c. Hydrocracking and hydrodesulfurization

a. Catalytic hydrogenation of fats and oils

REFERENCES

General

- 4. H.F. Rase, Chemical Reactor Design Of Process Plants, Wiley, New York, 1977, 2 vols.
- 5. Y.T. Shah, Gas-Liquid-Solid Reactor Design, McGraw-Hill, New York, 1979.
- 6. M.O. Tarhan, Catalytic Reactor Design, McGraw-Hill, New York, 1983.
- 7. S.M. Walas, Reaction Kinetics for Chemical Engineers, McGraw-Hill, New York, 1959.
- 8. S.M. Walas, Chemical reactor data, Chem. Eng., 79-83 (14 Oct. 1985).
- 9. K.R. Westerterp, W.P.M. Van Swaaij, and A.A.C.M. Beenackers, Chemical Reactor Design and Operation, Wiley, New York, 1984.
- 1. J. Beek, Design of packed catalytic reactors, Adv. Chem. Eng. 3, 203-271 (1962).
- 2. L.K. Doraiswamy and M.M. Sharma, Heterogeneous Reactions: Analysis, Examples and Reactor Design, Wiley, New York, 1984, 2 vols.
- 3. E.B. Nauman and B.A. Buffham, Mixing in Continuous Flow Systems, Wiley, New York, 1983.

610 CHEMICAL REACTORS

Types of Reactors

 Ullmann, Reaktionsapparate, in *Encyclopedia of Chemical Technology*, Verlag Chemie, Weinheim, Germany, 1973, Vol. 3, pp. 320–518.

Catalysts and Chemical Processes

- J.E. Bailey and D.F. Ollis, *Biochemical Engineering Fundamentals*, McGraw-Hill, New York, 1986.
- C.H. Bamford and C.F.H. Tipper (Eds.), Complex Catalytic Processes, Comprehensive Chemical Kinetics Vol. 20, Elsevier, New York, 1978.
- B.C. Gates, J.R. Katzer, and G.C.A. Schuit, *Chemistry of Catalytic Processes*, McGraw-Hill, New York, 1979.
- B.E. Leach (Ed.), *Applied* Industrial Catalysis, Academic, New York, 1983-1985, 3 vols.
- 5. H. Pines, Chemistry of Catalytic Conversions of Hydrocarbons, Academic, New York, 1981.
- V.A. Roiter (Ed.), Handbook of Catalytic Properties of Substances (in Russian), Academy of Sciences, Ukrainian SSR, Kiev, USSR, 1968–date, 4 vols. to date.
- C.N. Satterfield, Mass Transfer in Heterogeneous Catalysis, MIT Press, Cambridge, MA, 1970.
- C.N. Satterfield, *Heterogeneous Catalysis in Practice*, McGraw-Hill, New York, 1980.
- S. Strelzoff, Technology and Manufacture of Ammonia, Wiley, New York, 1981.
- C.L. Thomas, Catalytic Processes and Proven Catalysts, Academic, New York, 1970.
- 11. D.L. Trimm, Design of Industrial Catalysts, Elsevier, New York, 1980.
- 12. Ullmann's Encyclopedia of Chemical Technology: Biotechnologie, Vol. 8,

pp. 497-526, 1972; Enzyme, **Vol.** 10, pp. 471-561, 1975, Verlag Chemie Weinheim, Germany,

13. C.A. Vancini, Synthesis of Ammonia, Macmillan, New York, 1971.

Heat Transfer in Reactors

- N.P. Cheremisinoff and P.N. Cheremisinoff, Hydrodynamics of Gas-Solids Fluidization, Gulf, Houston, 1984, abstract section.
- M.B. Glaser and G. Thodos, Heat and momentum transfer in flow of gases through packed beds, AIChE J. 4, 63-74 (1958).
- 3. J.R. Grace, Fluidized bed heat transfer, in *Handbook of Multiphase Systems* (Hetsroni, Ed.), Hemisphere, New York, 1982.
- L.A. Korotjanskaja et al. cited by Doraiswamy and Sharma, *loc. cit.*, 1984, p. 323.
- 5. M. Leva, Fluidization, McGraw-Hill, New York, 1959.
- 6. J.Y. Oldshue, Fluid Mixing Technology, McGraw-Hill, New York, 1983.
- 7. H.F. Rase, Chemical Reactor Design for Process Plants, Wiley, New York, 1977, 2 vols.
- E.U. Schliinder, Transport phenomena in packed bed reactors, in *Chemical Reactor Engineering Reviews-Houston*, ACS Symposium 72, American Chemical Society, Washington, DC, 1978.
- H.A. Vreedenberg, Heat transfer between a fluidized bed and a horizontal tube, *Chem. Eng. Sci.* 9, 52-60 (1958); Vertical tubes, *Chem. Eng. Sci.* 11, 274-285 (1960).
- L. Wender and G.T. Cooper, Heat transfer between fluidized beds and bounding surfaces-correlation of data, AIChE J. 4, 15-23 (1958).
- S.S. Zabrodsky, Hydrodynamics and Heat Transfer in Fluidized Beds, MIT Press, Cambridge, MA, 1966.
- 12. S.S. Zabrodsky, N.V. Antonishin, and A.L. Parnas, On fluidized bed to surface heat transfer, Can. J. Chem. Eng. 54, 52-58 (1976).

PROCESS VESSELS

essels in chemical processing service are of two kinds: those substantially without internals and those with internals. The main functions of the first kinds. called drums or tanks. are intermediate storage or surge of a process stream for a limited or extended period or to provide a phase separation by settling. Their sizes may be established by definite process calculations or by general rules based on experience. The second category comprises the she//s of equipment such as heat exchangers, reactors, mixers, fractionators, and other equipment whose housing can be designed and constructed largely independently of whatever internals are necessary. Their major dimensions are established by process requirements described in other chapters, but considerations of adequate strength of vessels at operating pressures and temperatures will be treated in this chapter.

The distinction between drums and tanks is that of size and is **not** sharp. Usually they are cylindrical vessels with f/at or curved ends, depending on the pressure, and either horizontal or vertical. In a continuous p/ant, drums have a holdup of a few minutes. They are located between major equipmenr or supply feed or accumulate product. Surge drums between equipment provide a measure of stability in that fluctuations are not transmitted free/y along a chain, including those fluctuations that are characteristic of control instruments of normal sensitivity. For example, reflux drums provide surge between a condenser and its tower and downstream equipment; a drum ahead of a compressor will ensure freedom from liquid entrainment and one ahead of a fired heater will protect the tubes from running dry; a drum following a reciprocating compressor will smooth out pressure surges, etc. Tanks are larger vessels, of several hours holdup usually. For instance, the feed tank to a batch distillation may ho/d a day's supply, and rundown tanks between equipment may provide several hours holdup as protection of the main storage from possible off-specification product and opportunity for local repair and servicing without disrupting the entire process.

Storage tanks are regarded as outside the process battery limits, on tank farms. Their sizes are measured in units of the capacities of connecting transportation equipment: 34,500 gal tank cars, 8000 gal tank trucks, etc., usually at least 1.5 times these sizes. Time variations in the supply of raw materials and the demand for the products influence the sizes and numbers of storage tanks.

Liquid storage tanks are provided with a certain amount of vapor space or freeboard, commonly 15% below 500 gal and 70% above 500 gal. Common erection practices for liquid storage tanks are:

- a. For less than 7000 gal, use vertical tanks mounted on legs.
- b. Between 1000 and 10,000 gal, use horizontal tanks mounted on concrete foundation.
- c. Beyond 10,000 gal, use vertical tanks mounted on concrete foundations.

Liquids with high vapor pressures, liquefied gases, and gases at high pressure are stored in elongated horizontal vessels, less often in spherical ones. Gases are stored at substantially atmospheric pressure in gas holders with floating roofs that are sealed with liquid in a double wall. Liquefied gases are maintained at subatmospheric temperatures with external refrigeration or autorefrigeration whereby evolved vapors are compressed, condensed, cooled, and returned to storage.

Liquids stored at near atmospheric pressure are subject to breathing losses: As the tank cools during the night air is drawn in, then vaporization occurs to saturation, and the vapor mixture is expelled as the tank warms up during the day. Volatile liquids such as gasoline consequent/y suffer a material loss and also a change in composition because of the selective loss of lighter constituents.

In order to minimize such effects, several provisions are made, for example:

- A floating roof is a pad which floats on the surface of the stored liquid with a diameter of about a foot less than that of the tank. The annular space between the float and the she// may be sealed by one of several available methods.
- 2. An expansion roof allows thermal expansion of the vapor space. It rides with the changing vapor and is sealed with liquid in a double wall.
- **3.** A bag of vapor resistant fabric is allowed to expand into a housing of much smaller diameter than that of the storage tank. This is a lower cost construction than either of the other two.

Weather resistant so/ids such as coal or sulfur or ores are stored in uncovered pi/es from which they are retrieved with power shovels and conveyors. Other solids are stored in silos. For short-time storage for process use, so/ids are stored in bins that are usually of rectangular cross section with cone bottoms and hooked up to process with conveyors. A// aspects of the design of such equipment are covered in books by Reisnerand Rothe (1971) and Stepanoff (1969).

18.1. DRUMS

Liquid drums usually are placed horizontal and gas-liquid separators vertical, although reflux drums with gas as an overhead product commonly are horizontal. The length to diameter ratio is in the range 2.5-5.0, the smaller diameters at higher pressures and for liquid-liquid settling. A rough dependence on pressure is



The volume of a drum is related to the flow rate through it, but it depends also on the kinds of controls and on how harmful would be the consequences of downstream equipment running dry. Conventionally, the volume often is expressed in terms of the

612 **PROCESS VESSELS**

number of minutes of flow on a half-full basis. For many services, 5-10 min half-full is adequate but two notable exceptions are:

- 1. Fired heater feed surge drum for which the size is 10-30 min half-full.
- 2. Compressor feed liquid knockout drum which is made large enough to hold 10-20 min of liquid flow, with a minimum volume of 10 min worth of gas flow rate.

Other major services require more detailed consideration, as follows.

18.2. FRACTIONATOR REFLUX DRUMS

Commonly their orientation is horizontal. When a small amount of a second liquid phase (for example, water in an immiscible organic) is present, it is collected in and drawn off a pot at the bottom of the drum. The diameter of the pot is sized on a linear velocity of 0.5 **ft/sec**, is a minimum of 16in dia in drums of 4-8 ft dia, and 24 in. in larger sizes. The minimum vapor space above the high level is 20% of the drum diameter or 10 in (Sigales, 1975).

A method of sizing reflux drums proposed by Watkins (1967) is based on several factors itemized in Table 18.1. A factor F_3 is applied to the net overhead product going downstream, then instrument factors F_1 and labor factors F_2 which are added together and applied to the weighted overhead stream, and finally a factor F_4 is applied, which depends on the kind and location of level indicators. When *L* is the reflux flow rate and *D* the overhead net product rate, both in gpm, the volume of the drum (gal) is given by

$$V_d = 2F_d(F_1 + F_2)(L + F_3D)$$
 gal, full. (18.1)

For example, with L = 400 gpm and D = 200 gpm, at average conditions $F_1 = 1, F_2 = 1.5, F_3 = 3, F_4 = 1.5$, and

$$V_d = 2(1.5)(1+1.5)(400+3(200)) = 7500$$
 gal, full

TABLE 18.1 Factors for Sizing Reflux Accumulators a. Factors F_1 and F_2 on the Reflux Flow Rate

	Instrument	Factor F ₁	Lab	or Facto	r F₂
Operation	w/ Alarm	w/o Alarm	Good	Fair	Poor
FRC LRC	12	112	1 1	1.5 1.5	2 2
TRC	1 <u>1</u>	2	1	1.5	2

b. Factor $\textit{\textbf{F}}_{\!\!3}$ on the Net Overhead Product Flow to External Equipment

Operating Characteristics	F ₃
Under good control	2.0
Under fair control	3.0
Under poor control	4.0
Feed to or from storage	1.25

c. Factor F₄ for Level Control

	F ₄
Board-mounted level recorder	1.0
Level indicator on board	1.5
Gage glass at equipment only	2.0

or, 6.25 min half-full. With the best of everything, $F_1 = 0.5$, $F_2 = 1$, $F_3 = 2$, $F_4 = 1$, and

$$V_d = 2(0.5 + 1)(400 + 2(200)) = 2400 \text{ gal}, \text{ full}$$

or 2.0 min half-full. The sizes figured this way are overruled when the destination of the net product is to a fired heater or a compressor; then the numbers cited in Section 18.1 are applicable.

Although this method seems to take in a number of pertinent factors, it is not rigorous and may not dissuade some practitioners from continuing to size these drums on the basis of 5 min half-full.

18.3. LIQUID-LIQUID SEPARATORS

Vessels for the separation of two immiscible liquids usually are made horizontal and operate full, although some low rate operations are handled conveniently in vertical vessels with an **overflow** weir for the lighter phase. The latter mode also is used for particularly large flows at near atmospheric pressures, as in the mixer-settler equipment of Figure 3.19. With the usual L/D ratio of three or more, the travel distance of droplets to the separated phase is appreciably shorter in horizontal vessels.

Since the rise or fall of liquid droplets is interfered with by lateral flow of the liquid, the diameter of the drum should be made large enough to minimize this adverse effect. A rule based on the Reynolds number of the phase through which the movement of the liquid drops occurs is proposed by Hooper and Jacobs (1979). The Reynolds number is $D_h u \rho / \mu$, where D_h is the hydraulic diameter and u is the linear velocity of the continuous phase. The rules are:

N _{Re}	Effect
Less than 5000 5000–20.000	little problem some hindrance
20.000-50.000	major problem may exist
Above 50,000	expect poor separation

The jet effect of an inlet nozzle also may interfere with the phase separation. Ideally the liquid should be introduced uniformly over the cross section, but a baffle at the inlet nozzle may reduce **such** a disturbance adequately. More elaborate feed diffusers sometimes may be worthwhile. Figure 18.1 shows a perforated baffle.

Fall or rise of droplets of one liquid in another is represented closely by Stokes law,

$$u = g_c(\rho_2 - \rho_1)d^2/18\mu.$$
(18.2)

In common units,

$$u = 9.97(10^6)(\rho_2 - \rho_1)d^2/\mu, \quad \text{ft/min}, \tag{18.3}$$

where the ρ_i are specific gravities, *d* is the droplet diameter (ft), and μ is the viscosity of the continuous phase (cP).

The key property is the droplet diameter, of which many studies have been made under a variety of conditions. In agitated vessels, experience shows that the minimum droplet diameters are in the range of 500-5000 μ m. In turbulent pipeline flow, Middleman (1974) found that very few droplets were smaller than 500 μ m. Accordingly, for separator design a conservative value is 150 μ m, which also has been taken as a standard in the API Manual on Disposal of Refinery Wastes (1969). With this diameter,

$$u = 2.415(\rho_2 - \rho_1)/\mu, \quad \text{ft/min.}$$
 (18.4)

Which phase is the dispersed one can be identified with the



Figure 18.1. Drums for distillation tower reflux and for reciprocating compressor surge. (a) A reflux drum with a pot for accumulation and removal of a heavy phase. The main liquid is removed on level control through a vortex breaker. When the pot is large enough, it can accommodate an interface control for automatic drainage; otherwise the drain valve is hand set and monitored by an operator. (b) Arrangement of a surge drum for eliminating the high frequency response of a reciprocating compressor. Details are given by Ludwig (Applied Process Design for Chemical and Petrochemical Plants, Gulf *Houston, 1983*, Vol. 3).

factor

$$\psi = \frac{Q_L}{Q_H} \left(\frac{\rho_L u_H}{\rho_H u_L} \right) \mathbf{r}^{0.3} \tag{18.5}$$

Docult

with the statements of this table (Selker and Schleicher, 1965):

Ψ	Result
co.3	light phase always dispersed
0.3-0.5	light phase probably dispersed
0.5-2.0	phase inversion probable, design for worst case
2.0-3.3	heavy phase probably dispersed
>3.3	heavy phase always dispersed

These relations are utilized in Example 18.1 and the resulting design is represented on Figure 18.2.

COALESCENCE

The rate of separation of liquid phases can be enhanced by shortening the path through which the droplets need rise or fall or

by increasing their diameters. Both effects are achieved by forcing the flow between parallel flat or crimped plates or through tower packing or through a mass of packed fibers. The materials should be wetted by the disperse phase and preferably rough. Fine droplets will impinge on the surfaces and will grow by accretion of other droplets. The separator in such cases will consist of a coalescing section and an open section where the now enlarged droplets can separate freely. Figure 18.3 is of a separator equipped with a coalescer that is especially suited to the removal of relatively small quantities of dispersed liquid. Cartridge-type coalescers are described by Redmon (1963). Packed separators have been studied by Davies, Jeffrys, and Azfal (1972) and the subject is reviewed by Laddha and Degaleesan (1983). Coalescence also can be induced electrically, a process that is used widely for the precipitation of brine from crude oils. Proprietary equipment is available for this purpose. The subject is discussed by Waterman (1965) and in detail by Fronczak (1983).

OTHER METHODS

Very fine dispersions can be separated effectively with disk-type centrifuges. Commercial units have capacities of 5-500 gpm and are capable of removing water from hydrocarbons down to the ppm range. A mild centrifugal action is achieved in hydrocyclones. They have been studied for liquid-liquid separation by Sheng, Welker, and Sliepcevich (1974), but their effectiveness was found only modest. The use of hydrocyclones primarily for the recovery of solid particles from liquids is described in the book of Bradley (1965). A symposium on coalescence has papers by Belk (1965), Jordan (1965), Landis (1965), and Waterman (1965).

18.4. GAS-LIQUID SEPARATORS

Droplets of liquid are removed from a gas phase by three chief methods:

- 1. Settling out under the influence of gravity.
- 2. Settling out under centrifugal action.
- 3. Impingement and coalescence on solid surfaces followed by settling.

Available methods for the design of liquid separators are arbitrary in some respects but can be made safe economically. Figure 18.4 illustrates some of these methods.

DROPLET SIZES

The period of time needed for settling out depends on the size distribution of droplets and the required completeness of removal. Under most conditions the droplet diameter is an elusive quantity. A few observations are mentioned by York (1983). Garner et al. (1954) found 95% of evaporator entrainment to be smaller than 18-25 μ m. From spray nozzles the droplets are 90 wt % greater than 20 μ m. Spray disks made droplet diameters in the range 100–1000 ym. Sprays resulting from splashing and pickup by vapors off condensed liquid films are as large as 5000 μ m. Some mists are very fine, however; those in sulfuric acid plants are mostly less than 10 μ m, and in some equipment 50 wt % are less than 1 μ m (Duros and Kennedy, 1978). On the whole, sprays in process equipment usually are greater than 20 pm, mostly greater than 10 μ m.

The amount of entrainment has been studied mostly in distillation equipment. Figure 18.5 summarizes some of these data, and they are applied in Example 18.2. Equation 18.11 incorporates entrainment data indirectly.

A common belief is that 95% of entrainment can be removed in economically sized gravity separators, in excess of 99% with wire EXAMPLE 18.1 Separation of Oil and Water

Find the dimensions of a drum for the separation of oil and water at these conditions:

Oil at 180 cfh, sp $\mathbf{gr} = 0.90$, viscosity = 10 cP. water at 640 cfh, sp $\mathbf{gr} = 1.00$, viscosity = 0.7 cP.

Take a droplet size to be 150 μ m (0.0005 ft) and that the holdup in the tank is in the same proportions as in the feed. The geometry of the cross section:



$$A_{1} = \frac{180}{120} \frac{\pi}{4} D^{2} = \frac{D_{2}}{8} (\theta - \sin \theta),$$

$$\therefore \theta = 2.192 \text{ rad},$$

$$A_{1} = 0.7805 \frac{\pi}{4} D^{2} = 0.6130D^{2},$$

$$L = D \sin(\theta/2) = 0.88940,$$

$$S_{2} = D \left(\pi - \frac{\theta}{2}\right) = 2.0456D.$$

Hydraulic diameter of heavy liquid

2

$$D_h = \frac{4A_2}{L+S_2} = \frac{4(0.6130)D}{0.8894 + 2.0456} = 0.8354D.$$

The dispersion discriminant is

$$\psi = \frac{Q_L}{Q_H} \left(\frac{\rho_L}{\rho_H} \frac{\mu_H}{\mu_L} \right)^{0.3} = \frac{180}{640} \left(\frac{0.9(0.7)}{10} \right)^{0.3} = 0.123$$

< 0.30.

Therefore, oil is the dispersed phase:

$$N_{\rm Re} = \frac{D_h \mu \rho}{\mu} = \frac{D_h \rho}{\mu} \frac{Q}{4\pi D^2} = \frac{(62.4)640}{42(0.7)\pi} \frac{(0.8354)}{D}$$

= $\frac{25,076}{D}$.

Velocity of rise:

$$u_r = \frac{2.492(1.00 - 0.90)}{0.7} = 0.356 \text{ ft/min.}$$

Time of rise:

$$t = \frac{0.7286D}{0.356} = 2.0466D \text{ min.}$$

Forward velocity:

$$u_{H} = \frac{QH}{A_2} - \frac{640}{60(0.6130D^2)} = \frac{17.40}{D^2} \text{ ft/min.}$$

Flow distance:

i

$$L_f = tu_H = 2.0466D\left(\frac{17.40}{D^2}\right) = \frac{35.60}{D}$$
 ft.

The tangent to tangent length of the drum will be approximately 24 in. greater than L_f to accommodate inlet and outlet nozzles and baffles.

The Reynolds number identifies the quality of the separation, $N_{\rm Re} < 5000$ being good.

Some trials are

D (ft)	N _{Re}	t	щ	<i>L</i> , (ft)
5	5015	10. 23	0.696	7.12
3.5	7165	7.16	1.420	10.17
3	8358	6.14	1.933	Il.87

A vessel 5×9 ft would give excellent separation; 3×14 ft might be acceptable. A sketch of the proposed drum is in Figure 18.1.

mesh pads and other solid surfaces on which impingement and coalescence are forced, and approaching 100% in scrubbers and high speed centrifuges.

RATE OF SETTLING

The terminal or maximum settling velocity of a small droplet or particle in a gas is governed by one of Newton's equations.

$$u = f \sqrt{g_c D(\rho/\rho_g - 1)}. \tag{18.6}$$

In laminar flow the friction factor becomes a simple function of the Reynolds number,

$$f = \frac{18}{(Du\rho_g/\mu_g)}.$$
 (18.7)

When this substitution is made, the falling velocity becomes

$$u = g_c(\rho - \rho_g) D^2 / 18\mu, \qquad (18.2')$$

which is Stokes' equation. In view of the uncertainties with which droplet sizes are known in practical situations, Stokes equation usually is regarded as sufficiently descriptive of settling behavior. For example, it predicts that 100 μ m droplets of water fall at the rate of 1 .0 ft/sec in atmospheric air.

Another approximation of Newton's equation is written

$$\boldsymbol{\mu} = \boldsymbol{K} \sqrt{\rho / \rho_g - 1}, \tag{18.8}$$

where the coefficient **K** depends on the system. For the $100 \,\mu m$ droplets of water in air just cited, the coefficient becomes



Figure 18.2. A design of an oil-water separator for the conditions of Example 18.1, showing particularly the diffuser at the inlet nozzle and baffles at the outlets. *(Ifooper and Jacobs, 1979).*



Figure 18.3. Drums with coalescers for assisting in the separation of small amounts of entrained liquid. (a) A liquid-liquid separating drum equipped with a **coalescer** for the removal of small amounts of dispersed phase. In water-hydrocarbon systems, the pot may be designed for 0.5 ft/sec (Facet Enterprises, Industrial Division). (b) An oil-water separator with corrugated plate coalescers (General Electric Co.).

K = 0.035, and for other sizes it varies as the square of the diameter.

EMPTY DRUMS

The cross section of a vertical settling drum is found from the vapor rate and the allowable linear velocity with the equation

$$u = 0.14\sqrt{\rho/\rho_g - 1}, \text{ ft/sec},$$
 (18.9)

in which the coefficient of Eq. (18.8) has been evaluated for $200 \,\mu\text{m}$. The vertical dimension is more arbitrarily established. The liquid holdup is determined as in Section 18.2 and Table 18.1. For the vapor space, Watkins (1967) proposes the rules illustrated in Figure 18.6. When the calculated length to diameter ratio comes out less than 3, the length is increased arbitrarily to make the ratio 3; when the ratio comes out more than 5, a horizontal drum is preferably employed. Rules for horizontal drums also are shown on Figure 18.6. The vapor space is made a minimum of 20% of the drum volume which corresponds to a minimum height of the vapor space of 25% of the diameter, but with the further restriction that this never is made less than 12 in. When a relatively large amount of liquid must be held up in the drum, it may be advisable to increase the fraction of the cross section open to the vapor.

The diameter again is figured from the volumetric rate of the vapor and the linear velocity from Eq. (18.9). Since the upward drag of the vapor is largely absent in a horizontal drum, however, the coefficient K often is raised by a factor of 1.25. Example 18.3 deals with the design of both kinds of drums.

WIRE MESH PAD DEENTRAINERS

Pads of fine wire mesh induce coalescence of impinging droplets into larger ones, which then separate freely from the gas phase. Tower packings function similarly but are less effective and more difficult to install. The pads are made of metal wires or plastic strands or fiber glass. These data apply to stainless steel construction:

Efficiency			<u>к</u>			
(%)	lbs/ cuft	sqft/cuft	Pressure	Vacuum		
99.0	5-7	65	0.40			
99.5	9	8 5	0.35	0 00 0 07		
99.9	12	115	0.35	0.20-0.27		
99.9	13-14	120	0.25			
	Efficiency (%) 99.0 99.5 99.9 99.9	Efficiency (%) Ibs/cuft 99.0 5-7 99.5 9 99.9 12 99.9 13-14	Efficiency Ibs/cuft sqft/cuft 99.0 5-7 65 99.5 9 85 99.9 12 115 99.9 13-14 120	Efficiency Ibs/ cuft sqft/cuft Pressure 99.0 5-7 65 0.40 99.5 9 85 0.35 99.9 12 115 0.35 99.9 13-14 120 0.25		

A pad thickness of 4in. is minimum, 6in. is popular, and up to 12 in. may be required for fine mists.

The values of K in the preceding table are with a standard disengaging height of 10 in. The effect of other heights h is given by the equation

$$K = 0.021 + 0.0325h, \quad 3 \le h \le 12, \tag{18.10}$$

with a maximum value of 0.40. This relation is for standard efficiency pads. Lower values can be expected in aqueous systems where the surface tension has been reduced by surfactants.

When the pad is installed in a vertical or inclined position, the values of K should be taken 2/3 of the horizontal ones.

At high liquid rates droplets tend to be reentrained and the pad may become flooded. Some data obtained by Poppele (1958) are cited by York (1983, p. 194). A graphical correlation credited to the **Fluor** Co. is represented by **Branan** (1983, p. 67) by the equation

$$K = -0.0073 + \frac{0.263}{x^{1.294} + 0.573}, \quad 0.04 \le x \le 6.0, \tag{18.11}$$



Figure 18.4. Typical installations of mesh pads in equipment (Metal Textile Corp, Bulletin ME-7; from Ludwig, 1977, Vol. I, p. 159).

where \boldsymbol{x} is a function of the weight flow rates and densities of the phases

$$\mathbf{x} = (\mathbf{W}_L/\mathbf{W}_V)\sqrt{\rho_V/\rho_L} \tag{18.12}$$

Good performance can be expected at velocities of 30-100% of those calculated with the given Ks. Flooding velocities are at 120–140% of the design rates. At low velocities the droplets drift through the mesh without coalescing. A popular design velocity is about 75% of the allowable. Some actual data of the harmful effect of low velocities were obtained by Carpenter and Othmer (1955); they found, for example, that 99% of 6 μ m droplets were removed at 6.8 ft/sec, but 99% of 8 μ m at the lower velocity of 3.5 ft/sec.

Pressure drop in pads usually is small and negligible except at flooding; the topic is discussed by York (1983).

In existing drums or when the drum size is determined primarily by the required amount of liquid holdup, the pad dimensions must conform to the superficial velocities given by the design equation. This may necessitate making the pad smaller than the available cross section of the drum. Figure 18.7 shows typical installations. On the other hand, when the pad size is calculated to be greater than the available cross section and there develops a possibility of reentrainment of large droplets from the exit surface of the pad, a downstream settling drum or a high space above the mesh can be provided.

Good design practice is a disengaging space of 6-18 in., the more the better, ahead of the pad and 12in. above the pad. Other details are shown on Figure 18.8. A design is provided in Example 18.4.

18.5. CYCLONE SEPARATORS

In addition to those already discussed, a variety of proprietary and home-made devices can remove entrainment more or less effectively. Some of them are represented on Figure 18.8.

- a. A simple change of direction and impingement on the walls of the drum.
- **b.** Impingement on a baffle.
- c. Tangential inlet at high velocity and change of direction.
- d. Multiple baffles, without or with coarse spray irrigation.
- e. A pipeline deentrainer.

The capacity and effectiveness of proprietary devices such as items c to e cannot be estimated from general knowledge, but manufacturers usually claim that they can be sized to remove 99% of 8 μ m droplets or particles. "Separators, entrainment" is an entry in the index of the *Chemical Engineering Catalog* which is a guide to manufacturers who may be consulted about the performance of their equipment.

In cyclone separators the gas enters tangentially at a high velocity, rotates several times, acd leaves through a central pipe. Such equipment has been studied widely, particularly for the removal of dusts and catalyst fines in **fluidized** bed systems. The literature is reviewed by Rietema and Verver (1961), Maas (1979), Zenz (1982), and Semrau (1984).

Typical cyclone dimension ratios are indicated on Figure 18.9. For liquid knockout the bottom head often is made dished as on Figure 18.10 which also shows standard dimensions. Inlet velocities



Figure 18.5. Entrainment from sieve trays in the units mols liquid entrained/mol of liquid downflow; LM_I is the weight rate of flow of liquid and VM_V is the weight rate of flow of vapor. The flooding correlation is Figure 13.32(b). [Fair *and* Matthews, Pet. Refiner **37**(4), 153 (1958)].

should be in the range 100-150 ft/sec, the higher the better, but may be limited by the occurrence of reentrainment and unacceptable pressure drop. The pressure drop is estimated in terms of velocity heads, a value of four being commonly taken. Accordingly,

$$\mathbf{AP} = 4\rho V^2 / 2g = 4.313\rho (\text{ft/sec}/100)^2, \text{ psi.}$$
(18.13)

For atmospheric air, for instance, this becomes

$$AP = 0.323 (ft/sec/100)^2$$
, psi. (18.14)

EXAMPLE 18.2

Quantity of Entrainment on the Basis of Sieve Tray Correlations

The conditions of Example 13.15 will be used. This is the case of a standard sieve tray with 24in. spacing and to operate at 80% of Aooding. The entrainment correlation is Figure 18.4 for which the value of the abscissa was found to be

$$w_1/w_G \sqrt{\rho_G/\rho_L} = 0.2924.$$

At 80% of **flooding** the ordinate of Figure 18.4 is



Figure 18.6. Knockout drums. Key dimensions of vertical and horizontal types.

For the design of Figure 18.10, the size of the inlet is selected at a specified inlet velocity and required volumetric rate; the other dimensions then are fixed as given for this standard.

Very high velocities tend to skim the liquid film off the vessel wall and off the liquid at the bottom. The liquid also tends to creep up the wall and down the exit pipe where it is picked up by the exit gas. The skirt shown on Figure 18.9 is designed to prevent reentrainment of the creeping liquid, and the horizontal plate in Figure 18.10 prevents vortexing of the accumulated liquid and pickup off its surface.

Efficiencies of 95% for collection of 5 μ m droplets can be

 $\psi = 0.008$ mol entrained liquid/mol liquid downflow.

Since $w_L/w_G = 259,100/271,500 = 0.954$ mol liquid/mol vapor (assuming the same molecular weights), the entrainment expressed with reference to the vapor flow is

 $\psi = 0.008(0.954) = 0.0076 \text{ mol liquid/mol vapor flow.}$

The linear velocity of the vapor was found in Example 13.15 to be 0.597 ft/sec for this condition.

EXAMPLE 18.3 Liquid Knockout Drum (Empty)

Gas at the rate of 3000 cfm and liquid at 25 cfm enter a drum in which entrainment is to be removed. Holdup of liquid in the drum is 10 min. The properties are those of air and water at atmospheric conditions. Find the size of the drum needed to remove droplets greater than 200 μ m dia.

Vertical drum, with Eq. (18.9):

 $u = 0.14\sqrt{62.4/0.075 - 1} = 4.04$ ft/sec. $D = \sqrt{3000/60(\pi/4)(4.04)} = 3.97$ ft, say 4.0 ft.

From Figure 18.5, the vapor space is a minimum of 5.5 ft. The liquid depth is

$$L_{\rm liq} = \frac{250}{(\pi/4)D^2} = 19.9 \, {\rm ft} = 10 \, {\rm min} \, {\rm holdup},$$

L = 19.9 + 5.5 = 25.4 ft, LID = 25.4/4 = 6.35.

achieved by proper design of cyclone separators. For applications such as knockout drums on the suction of compressors, however, it is sufficient to remove only droplets greater than 40-50 μ m.

Capacity and efficiency depend on the inlet velocity and the dimensions of the vessel. Correlated studies have been made chiefly for the design of Figure 18.9 with a rectangular inlet whose width is D/4 (one-fourth of the vessel diameter) and whose height is 2-3 times the width. A key concept is a critical particle diameter which is the one that is removed to the extent of 50%. The corresponding % removal of other droplet sizes is correlated by Figure 18.11. The

With D = 4.5 ft, L = 15.7 + 5.5 = 21.2, and L/D = 4.71. Horizontal drum: The allowable velocity is 25% greater:

u = 1.25(4.04) = 5.05 ft/sec.

Try several fractional vapor cross sections ϕ :

 $D = \sqrt{50/5}.05(\pi/4)\phi = \sqrt{12.61/\phi}$ $L = 250/(1-\phi)(\pi/4)D^2 = 25.24\phi/(1-\phi),$ h = depth of liquid.

φ	h/D	D	L	L/D
0.2	0.75		8.31 (6.2)	0.78
0.3	0.66	6.48 (6.5)	10.82 (10.8)	1.66
0.4	0.58	5.61 (5.5)	16.83 (17.5)	3. 18
0.5	0.50	5.02 (5.0)	25.24 (25.5)	5.10

Accordingly, a horizontal vessel between 5.0 and 5.5 ft dia with a liquid depth between 58 and 50% of the diameter falls in the usual economic range.

equation for the critical particle diameter is

$$(D_{p})_{\rm crit} = \frac{9\mu D}{4\pi N_{t} V(\rho - \rho_{g})} \bigg]^{0.5},$$
(18.15)

where D is the diameter of the vessel and V is the inlet linear velocity. The quantity N, is the number of turns made by the gas in the vessel. A graphical correlation given by Zenz (1982) can be



Figure 18.7. Key dimensions of knockout drums equipped with mesh pads. (a) Vertical knockout drum. (b) Horizontal knockout drum.



Figure 18.8. Principles of entrainment separation and some commercial types of equipment. (a) Basic principles of entrainment separating equipment: (i) change of direction; (ii) impingement on a baffle; (iii) tangential inlet resulting in centrifugal force. (b) Wire or fiber mesh pad, typical installations as in Figure 18.7. (c) A separator combining impingement and centrifugal force (V.D. Anderson Co.). (d) Equipment with impingement and change of direction (*Wright-Austin* Co.). (e) Multiple zig-zag baffle arrangement (*Peerless Mfg.* Co.).

represented by the equation

$$N_t = \begin{bmatrix} 0.1079 & 0.00077V + 1.924(10^{-6})V^2 \end{bmatrix} V$$
(18.16)

with V in ft/sec. With a height of opening equal to 2.5 times the width, the volumetric rate is

$$Q = AV = 2.5D^2 V / 16. \tag{18.17}$$

These relations are used in Example 18.5 to find the size of a separator corresponding to a specified critical particle diameter, and to the reverse problem of finding the extent of removal of particles when the diameter of the vessel and the velocity are specified.

To obtain a high efficiency, the vessel diameter must he small, but in order to accommodate a required volumetric rate, many units in parallel may be needed. These units, called multicyclones, may be incorporated in a single shell at a cost that may be justifiable in view of greater efficiency and lower pressure drop.

18.8. STORAGE TANKS

Cylindrical tanks for the storage of inflammable liquids above or under ground at near atmospheric pressure are subject to standards of Underwriter Laboratories or of the API. Underwriters covers some smaller sizes. Both sets of standards are restricted to steel construction for essentially noncorrosive service. Various manufacturers supply Underwriter or API tanks as a matter of course.

Standard tanks are made in discrete sizes with some latitude in combinations of diameter and length. For example, Table 18.2 shows the several heights of 30ft diameter tanks among API standard sizes. The major specification is that of metal wall thickness. In smaller sizes the thickness is determined by requirements of rigidity rather than strength. Some general statements about metal thickness of tanks may be given.

Horizontal tanks. Above ground they are limited to 35,000 gal. Normally they are supported on steel structures or concrete saddles at elevations of 6 to 10 ft. The minimum thickness of shell and heads is 3/16 in. in diameters of 48-72 in. and 1/4 in. in diameters of 73-132 in.

EXAMPLE 18.4

Knockout Drum with Wire Mesh Deentrainer

For the flow conditions of Example 18.2, design a drum with a standard efficiency stainless steel wire mesh pad. For this condition, k = 0.35, so that

$$u = 0.35\sqrt{62.4/0.075} - 1 = 10.09 \text{ ft/sec},$$

$$D = \sqrt{50/(\pi/4)u} = 2.51 \text{ ft}.$$

With 2 in. support rings the pad will have a diameter of 34 in. The size of the drum is set largely by the required liquid holdup of 250 cuft. On the basis of Figure 18.7, the height of vessel above the liquid level is 4 ft. As in Example 18.2, take the diameter to be 4.5 ft. Then

$$\begin{split} L_{\rm liq} &= 25 [10/(\pi/4)(4.5)]^2 = 15.7 ~{\rm ft}, \\ L &= 15.7 ~+ 4.0 = 19.7 ~{\rm ft}, \end{split}$$
L/D = 19.7/4.5 = 4.38

This ratio is acceptable. As a check, use Eqs. (18.11) and (18.12):

$$\begin{aligned} x &= (W_L/W_v)\sqrt{P_v/P_L} = V_L/V_v\sqrt{P_L/P_v} \\ &= 25/3000\sqrt{62.4/0.075} = 0.24, \\ k &= -0.0073 + 0.263/[(0.24)^{1.294} + 0.5731] \\ &= 0.353, \end{aligned}$$

which is close to the assumed value, k = 0.35.





Figure 18.9. Typical dimension ratios of a cyclone separator.



10 24 54 36 48 14 28 68 45 60 6 16 30 76 50 66 7 18 36 80 54 72 8 20 42 94 63 84 9 10 24 48 108 72 96

 3
 6
 38
 22
 30

 4
 8
 38
 22
 30

 6
 12
 38
 22
 30

8 18 42 26 36

Figure 18.10. Dimensions of standard liquid knockout drums with tangential inlets.



Figure 18.11. Percent removal of particles in a cyclone as a function of their diameters relative to the critical diameter given by Eqs. (18.15) and (18.16) (*Zenz*, 1982).

Vertical tanks. Those supported above ground are made with dished or conical bottoms. Flat bottomed tanks rest on firm foundations of oiled sand or concrete. Supported flat bottoms usually are 1/4 in. thick. Roof plates are 3/16 in. thick. Special roof

constructions that minimize vaporization losses were mentioned earlier in this chapter; they are illustrated by Mead (1964), by Riegel (1953), and in manufacturers catalogs. The curved sides are made of several courses of plate with thicknesses graduated to meet requirements of strength. The data of the selected API tanks of Table 18.2 include this information. Figure 18.12 illustrates the facilities that normally are provided for a large storage tank.

In order to minimize hazards, storage tanks for inflammable or toxic materials may be buried. Then they are provided with an overburden of 1.3 times the weight of water that the tank could hold in order to prevent floating after heavy rainfalls.

Cylinders with curved heads are used for pressure storage at 5-230 psig. In the range of 5-10 psig, spheroids and other constructions made up with curved surfaces, as in Figure 18.12(c) are being used in quite large sizes, often with refrigeration to maintain sufficiently low pressures. More illustrations of such equipment appear in manufacturers' catalogs and in the books of Mead (1964) and Riegel (1953).

Mention of vessels for the storage of gases was made at the beginning of this chapter, and Figure 18.12(d) shows the principles of some suitable designs. Design for storage of granular solids includes provisions for handling and withdrawal, as in the case of Figure 18.13.

18.7. MECHANICAL DESIGN OF PROCESS VESSELS

Process design of vessels establishes the pressure and temperature ratings, the length and diameter of the shell, the sizes and locations of nozzles and other openings, all internals, and possibly the material of construction and corrosion allowances. This information must be supplemented with many mechanical details before fabrication can proceed, notably wall thicknesses.

Large storage tanks are supported on a concrete pad on the ground. Other vessels are supported off the ground by various means, as in Figure 18.14.

For safety reasons, the design and construction of pressure vessels are subject to legal and insurance standards. The ASME

EXAMPLE 18.5

Size and Capacity of Cyclone Separators

Air at 1000 cuft/sec and density of 0.075 lb/cuft contains particles with density 75 lb/cuft. 50% of the 10 μ m diameter particles are to be recovered. Find the sizes and numbers of cyclones needed with inlet velocities in the range of 50-150 ft/sec. The inlet is rectangular with width D/4 and height 2.5D/4, where D is the diameter of the vessel.

Equation (18.15) becomes

$$\frac{D}{N_{t}V} = \frac{4\pi(\rho - \rho_{g})D_{p}^{2}}{9\mu}$$
$$= \frac{4\pi(75 - 0.075)}{9(1.285)(10^{-5})} \left(\frac{10}{304,800}\right)^{2} = 0.00876$$

where N, is given by Eq. (18.16). The number of vessels in parallel is

$$n = \frac{\mathbf{Q}}{AV} = \frac{100}{(2.5/16)D^2V} = \frac{6400}{D^2V}.$$

The results at several velocities are summarized.

V (cfs)	N _t	D (ft)	n
50	3.71	1.62	48.8
100	5.01	4.39	3. 32
144	5.32	6.71	1.0

From Figure 18.11, the percentage recoveries of other-sized particles are:

$D_p / (D_p)_{crit}$	% Recovered
0.3	9
0.5	22
0.6	30
1	50
2	70
6	S 0
9	98.5

When the smallest of these cyclones, 1.62 ft dia, is operated at 150 cuft/sec,

$$N_{t} = 5.35,$$

$$(D_{p})_{crit} = \left[\frac{9(1.285)(10^{-5})(1.62)}{4\pi(5.35)(150)(75 - 0.075)}\right]$$

$$= 1.574(10^{-5}) \text{ ft}, 4.80 \ \mu\text{m}.$$

TABLE 18.2. Storage Tanks, Underwriter or API Standard, Selected Sizes

Canacity		Weight		
Gallons	Diameter	Length	Thickness	in pounds
280	42"	4'-0"	$\frac{3}{16}''$	540
550	48″	6'-0"	$\frac{3}{16}''$	800
1000	48″	10'8"	3/16	1260
1000	64″	6'-0"	$\frac{3}{16}$ "	1160
1500	64″	9'-0"	$\frac{3}{16}$ "	1550
2000	64″	12'-0"	$\frac{3}{16}$ "	1950
3000	64″	18′-0″	<u>3</u> "	2730
4000	64″	24'-0"	<u>3</u> "	3510

a. Small Horizontal Underwriter Label

b. Horizontal or Vertical with Underwriter Label

Nominal		Dimensions			
Capacity Gallons	Diameter	Approx. Length	Thickness	Weight	No. of Supports
5, 000	6'-0"	23'-9"	<u>1</u> "	5, 440	3
5,000	7'-0"	17'-6″	<u>1</u> ″	5, 130	2
6,000	8'-0"	16'-1"	<u>1</u> "	5, 920	2
6,000	8'-0"	16'-1"	<u> </u>	6, 720	2
8,000	8'-0"	21'-4"	$\frac{1}{4}$ "	7, 280	2
8,000	8'-0"	21'-4'	4 <u>5</u> ″	8, 330	2
10, 000	8'0"	26' - 7"	16 <u>1</u> "	8, 860	3
10, 000	8′-0″	26' - 7"	<u>5</u> "	10, 510	3
10, 000	10'0"	17'-2"	$\frac{1}{4}$ "	8, 030	2
10, 000	10′—0″	17'-2"	<u>5</u> "	9, 130	2
10, 000	10'-6"	15'-8"	$\frac{1}{4}$ "	8, 160	2
10, 000	10'-6"	15′-8″	<u>5</u> "	9, 020	2
15, 000	8'-0"	39'-11"	<u>1</u> "	13, 210	4
15, 000	8'-0"	39'-11"	4 <u>5</u> "	14, 620	4
20, 000	1 0'-0 "	34'-1"	10 1/1	14, 130	3
20, 000	10'-0"	34'-1"	<u>5</u> "	16, 330	3
25, 000	10'-6"	38'-9"	$\frac{16}{1}$	17, 040	4
25, 000	10'-6"	38′-9″	<u>5</u> " 16	19, 010	4

c. Large Vertical, API Standard

Dimensi	ions	Ca	pacity	_			Shell Pla	ites (But	t Welded)			
Diameter	Height	42 gal per bbl	U.S. Gal	Bottom Plates	Ring 1	Ring 2	Ring 3	Ring 4	Ring 5	Ring 6	Ring 7	Top Angle	R o o f Plates
21' 0" 24' 0" 30' 0" 30' 0" 30' 0" 30' 0" 30' 0" 30' 0" 40' 0" 50' 0" 60' 0"	18'0 ³ / ₄ " 24'0" 24'0" 29'11 ¹ / ₄ " 35'10 ¹ / ₄ " 37'10 ³ / ₄ " 41'9 ³ / ₄ " 33'10 ³ / ₄ " 47'9" 39'10" 40'1"	1, 114 1, 933 3, 024 3, 769 4, 510 4, 766 5, 264 7, 586 16, 700 20, 054 27, 472	46, 788 81, 186 127, 008 158, 300 189, 420 200, 161 221, 088 318, 612 701, 400 842, 268 1 153, 824	1" 1" 1" 1" 1" 1" 1" 1" 1" 1" 1" 1" 1" 1	$ \frac{3}{16}, \frac{3}{16}, \frac{3}{16}, \frac{3}{16}, \frac{3}{16}, \frac{3}{16}, \frac{3}{16}, \frac{3}{16}, \frac{1}{16}, \frac{1}{4}, \frac{3}{16}, \frac{3}$	$\begin{array}{c} 3\\ 3\\ 16\\ 3\\ 16\\ 3\\ 16\\ 3\\ 16\\ 3\\ 16\\ 3\\ 16\\ 3\\ 16\\ 1\\ 1\\ 4\\ 0.29\\ 0.27\\ 0.32^{\circ} \end{array}$	$\frac{3}{16}, \frac{3}{16}, \frac{3}$	3 16 " 16 " 16 3 16 3 16 3 16 3 16 3 16	3 16 3 16 3 16 3 16 3 16 3 16 3 16 3 16	3" 16" <u>3</u> 16" <u>3</u> 16" <u>1</u>	<u>3</u> ″	3 " × 3" × $\frac{1}{4}$ " 3 " × 3" × 3" × $\frac{1}{4}$ " 3 " × 3" × 3" × 3" × 3" 4 " 4 " 5 " × 3" × 3" × 3" × 3" 5 " × 3" × 3" × 3" × 3" 5 " × 3" × 3" × 3" × 3" × 3" × 3" × 3" ×	316 316 316 316 316 316 316 316 316 316
100' 0" 150' 0"	40' 0" 48' 0"	55, 960 151, 076	2. 350. 320 6. 345. 192	4 <u>1</u> " <u>1</u> " 4	0. 57" 1. 03"	0. 45" 0. 85"	0. 33" 0. 68"	4 1 <u>4</u> 0. 50"	4 1/ 4 0. 33"	1 <u>"</u>		3" × ³ " × ≍″ ³ " × ³ " × ≍″ ³ " × ³ " × ≍″8 ³	16 3" 16" 3"



Figure 18.12. Examples of equipment for storage of liquids and gases in large quantities. (a) A large tank and its appurtenances, but with no provision for conservation of breathing losses (Graver *Tank* and Mfg. Co.). (b) Schematic of a covered floating roof tank in which the floating roof rides on the surface of the liquid. They also are made without the fixed roof [*R. Martin*, Petro/Chem. Eng., 23, (Aug. 1965)]. (c) Cutaway of a 40,000 Bbl spheroid for operation at 10 psig (*Chicago Bridge and Iron* Co.). (d) Design principles of tanks for storage of gases or liquids subject to breathing losses at atmospheric pressure: water seal, dry seal with flexible curtain, and variable vapor space controlled by a flexible curtain.

Codes apply to vessels greater than 6in. dia operating above 15 psig. Section VIII Division 1 applies to pressures below 3000 psig and is the one most often applicable to process work. Above **3000 psig** some further restrictions are imposed. Division 2 is not pressure limited but has other severe restrictions. Some of the many details covered by Division 1 are indicated by the references to parts of the code on Figure 18.15.

DESIGN PRESSURE AND TEMPERATURE

In order to allow for possible surges in operation, it is customary to raise the maximum operating pressure by 10% or 10-25 psi, whichever is greater. The maximum operating pressure in turn may

be taken as 25 psi greater than the normal. The design pressure of vessels operating at O-10 psig and $600-1000^{\circ}F$ is 40 psig. Vacuum systems are designed for 15 psig and full vacuum. Between -20 and 650°F, 50°F is added to the operating temperature, but higher margins of safety may be advisable in critical situations. When subzero temperatures have an adverse effect on the materials of construction, the working temperature is reduced appropriately for safety.

Allowable tensile stresses are one-fourth the ultimate tensile strength of the material of construction. Values at different temperatures are given in Table 18.4 for some steels of which shells and heads are made. Welded joint efficiencies vary from 100% for double-welded butt joints that are fully radiographed to 60% for



Figure 18.W. Equipment for handling, storing and withdrawing of granular solids in a glass manufacturing plant (*Stephens-Adamson Mfg. Co.*).

single-welded butt joints without backing strips and without radiographing. The Code has details.

SHELLS AND HEADS

Although spherical vessels have a limited process application, the majority of pressure vessels are made with cylindrical shells. The heads may be flat if they are suitably buttressed, but preferably they are some curved shape. The more common types of heads are illustrated on Figure 18.16. Formulas for wall thicknesses are in Table 18.3. Other data relating to heads and shells are collected in Table 18.5. Included are the full volume V_0 and surface S as well as the volume fraction V/V_0 corresponding to a fractional depth H/D in a horizontal vessel. Figure 18.17 graphs this last relationship. For ellipsoidal and dished heads the formulas for V/V_0 are not exact but are within 2% over the whole range.

FORMULAS FOR STRENGTH CALCULATIONS

The **ASME** Code provides formulas that relate the wall thickness to the diameter, pressure, allowable stress, and weld efficiency. Since they are theoretically sound only for relatively thin shells, some restrictions are placed on their application. Table 18.3 lists these

Figure 18.14. Methods of supporting vessels. (a) Saddle supports for horizontal vessels, usually of concrete. (b) Bracket or lug supports resting on legs, for either vertical or horizontal vessels. (c) Bracket or lug supports resting on steel structures, for either vertical or horizontal vessels. (d) Straight skirt support for towers and other tall vessels; the bearing plate is bolted to the foundation. (e) Flared skirt for towers and other tall vessels, used when the required number of bolts is such that the bolt spacing becomes less than the desirable 2 ft.





Figure 18.15. References to items covered in the ASME Code for Unfired Pressure Vessels, Section VIII Division 1 (Chuse and Eber, 1984).

TABLE 18.3. Formulas for Design of Vessels under Internal Pressure'

ltem	Thickness t(in.)	Pressure P(psi)	Stress S(psi)	Notes
Cylindrical shell	P R S E – 0.6P	$\frac{SEt}{R+0.6t}$	<u><i>P(R ≠</i> 0.6t)</u> t	$t \le 0.250, P \le 0.385SE$
Flat flanged head (a)	$D\sqrt{0.3P/S}$	$t^2S/0.3D^2$	$0.3D^2P/t^2$	
Torispherical head (b)	0.885 <i>PL</i> SE-0.1P	<u>SEt</u> 0.885L + 0.12	<u>P(0.885L + 0.1 t)</u> t	$r/L = 0.06, L \le D + 2t$
Torispherical head (b)	PLM 2SE - 0.2 P	2SEt LM + 0.2t	<u>P(LM + 0.2t)</u> 2t	$M = \frac{3 + (L/r)^{1/2}}{4}$
Ellipsoidal head (c)	<u>P D</u> 2 S E 0.2P	2SEt <i>D</i> + 0.2 <i>t</i>	$\frac{P(D + 0.2t)}{2t}$	h/D = 4
Ellipsoidal head (cl	<u>PDK</u> 2SE – 0.2P	$\frac{2SEt}{DK + 0.2t}$	<u>P(DK + 0.2t)</u> 2Et	$K = [2 + (D/2h)^2]/6, 2 \le D/h \le 6$
Hemispherical head (d) or shell	P R 2 S E - 0.2P	$\frac{2SEt}{R + 0.2t}$	<u>P(R + 0.2t)</u> 2t	t≤0.178 <i>D, P</i> ≤0.685 <i>SE</i>
Toriconical head (e)	$\frac{P D}{2 (SE - 0.6P) \cos \alpha}$	$\frac{2SEt \ cos \ \alpha}{D+1.2t \cos \alpha}$	$\frac{P(D+1.2t \cos \alpha)}{2t \cos \alpha}$	$lpha \leq 30^{\circ}$

Nomenclature: l = diameter (in.), l = joint efficiency (0.6–1.0), l = crown radius (in.), P = pressure (**psig**), h = inside depth of ellipsoidal head (in.), r = knuckle radius (in.), R = radius (in.), S = allowable stress (psi), t = shell or head thickness (in.).

Note: Letters in parentheses in the first column refer to Figure 18.16.

TABLE 18.4. Maximum Allowable Tensile Stresses (psi) of Plate Steels

(a) Carbon and Low Alloy Steels

A.S.M.E.		Nominal	Spec. min.	<u> </u>	For temperatures not exceeding °F.						
No.	Grade	composition	tensile strength	-20to 650	700	800	900	1000	1100	1200	
Carbon Steel											
SA515	55	C-Si	55, 000	13, 700	13, 200	10, 200	6, 500	2, 500			
SA515	70	C-Si	70, 000	17, 500	16, 600	12,000	6, 500	2, 500			
SA516	55	C-Si	55, 000	13, 700	13, 200	10, 200	6, 500	2, 500			
SA516	70	C-Si	70, 000	17, 500	16, 600	12,000	6, 500	2, 500			
SA265	Α		45,000	11, 200	11,000	9,000	6, 500				
SA285	В	· _	50, 000	12, 500	12, 100	9,600	6, 500				
SA285	С	1.1	55, 000	13, 700	13, 200	10, 200	6, 500				
Low-Alloy Steel											
SA202	Α	Cr-Mn-Si	75,000	18, 700	17, 700	12,600	6, 500	2,500			
SA202	В	Cr-Mn-Si	85,000	21, 200	19,800	12,800	6, 500	2, 500			
SA387	D*	2 <mark>1</mark> Cr-I M	60, 000	15, 000	15, 000	15, 000	13, 100	2, 800	4, 200	1, 600	

(b) High Alloy Steels

A.S.M.E.		Neminal	Specified minimum	For temperatures not exceedi				ng °F.						
No.	Grade	composition	strength	- 20 to 100	200	400	700	900	1000	1100	1200	1300	1400	1500
SA- 240	304	18 Cr-8 Ni	75, 000	1'8,700	15, 600	12, 900	11, 000	10, 100	9, 700	8, 800	6, 000	3, 700	2, 300	1, 400
SA- 240	304L†	18 Cr-8 Ni	70, 000	15, 600	13,300	10,000	9,300							
SA- 240	310s	25 Cr-20 Ni	75, 000	18, 700	16, 900	14, 900	12, 700	11,600	9,800	5,000	2,500	700	300	200
SA- 240	316	16 Cr-12 Ni-2 M o	75,000	18, 700	16, 100	13, 300	11, 300	10, 800	10, 600	10, 300	7,400	4, 100	2, 200	1, 700
SA- 240	410	13 Cr	65, 000	16, 200	15, 400	14, 400	13, 100	10, 400	6, 400	2, 900	1, 000			

t

(ASME Publications).

relations for cylindrical and spherical shells and for all but the last of the heads of Figure 18.16. For unusual shapes there are no simple methods of design; experience and testing may need to be resorted to if such shapes are required.

The formulas are expressed in terms of inside dimensions.



Figure 18.16. Fractional volumes of horizontal cylinders and curved heads at corresponding fractional depths, H/D.

Although they are rarely needed, formulas in terms of outside dimensions, say D_o , may be derived from the given ones by substitution of $D_o - 2t$ for D. For the 2: 1 ellipsoidal head, for instance,

$$=\frac{PD}{2SE-0.2P}=\frac{P(D_o-2t)}{2SE-0.2P}=\frac{PD_o}{2SE+1.8P}.$$
(18.18)

Example 18.6 investigates the dimensions and weight of a vessel to meet specifications. It is brought out that pressure vessels with large L/D ratios are lighter and presumably cheaper. A drawback may be the greater ground space needed by the slimmer and longer construction.

In addition to the shell and heads, contributions to the weight of a vessel include nozzles, **manways**, any needed internals, and supporting structures such as lugs for horizontal vessels and skirts for vertical ones. Nozzles and **manways** are standardized for discrete pressure ratings; their dimensions and weights are listed in manufacturers' catalogs. Accounting for these items may contribute **10–20%** to the calculated weight of the vessel.

Mechanical design specification sheets (Appendix B) summarize the information that a fabricator needs in addition to the general specifications of the vessel codes. Not all of the data on the specification summary are necessarily in the province of the process engineer; it may depend on the stage of the design and on who else in the organization is available to do the work.

TABLE 18.5. Heads and Horizontal Cylinders: Formulas for Partially Filled Volumes and Other Data





Figure **18.17**. Types of heads for cylindrical pressure vessels. (a) Flat flanged: KR= knuckle radius, SF= straight flange. (b) Torispherical (dished). (c) Ellipsoidal. (d) Spherical. (e) Conical, without knuckle. (f) Conical, with knuckle. (g) Nonstandard, one of many possible types in use.



EXAMPLE 18.6

Dimensions and Weight of a Horizontal Pressure Drum A drum is to operate at 500°F and 350 psig and to hold 5000 gal at a depth H/D = 0.8. Dished heads are to be used. The material is SA285A. Examine the proportions L/D = 3 and 5. Formulas are in Table 18.5:

 $V_{\text{tank}} = 5000/7.48 = 668.4$ cuft.

Two heads, capacity with H/D = 0.8,

$$V_h = V_0(V/V_0) = 2[0.0778D^3(2)(H/D)^2(1.5 - H/D)]$$

= 0. 1394 D^3 .

Shell capacity with H/D = 0.8,

$$\theta = 2 \operatorname{arccos}(1 - 1.6) = 4.4286 \text{ rad},$$

$$V_s = V_0(V/V_0) = (\pi/4)D^2L(1/2\pi)(\theta - \sin \theta)$$

$$= 0.6736D^2L$$

$$V_{\text{liquid}} = 668.4 = 0.1394D^3 + 0.6736D^2L$$

with L/D = 3,

$$D = \left(\frac{668.4}{2.1601}\right)^{1/3} = 6.76 \text{ ft, say } 6.5 \text{ ft,}$$
$$L = \frac{668.4 - 6736D^2}{0.6736D^2} = 22.1 \text{ ft, say } 22.0.$$

Allowable stress S = 11,200 psi.

Say joint efficiency is E = 0.9:

$$t_{\text{shell}} = \frac{PR}{SE - 0.6P} = \frac{350(39)}{0.9(11,200) - 0.6(350)} = 1.38 \text{ in.}$$

Dished head with L = D and r/L = 0.06:

$$t_h = \frac{0.885(350)(78)}{0.9(11,200) - 0.1(350)} = 2.41 \,\mathrm{in}.$$

Surfaces:

shell, $S = \pi DL = 449.3$ sqft, heads, $S = 2(0.842)D^2 = 71.2$ sqft, Weight = [449.3(1.4) + 71.2(2.4)]491/12 = 32,730 Ibs.

The results for L/D = 3 and 5 are summarized.

ltem	L/D=3	L/D=5
D (ft)	6.5	5. 5
L (ft)	22.0	32.0
t _{shell} (in.)	1.38 (1.4)	0.957 (1.0)
t _{head} (in.)	2.41 (2.4)	1.67 (1.7)
Weight (lb)	32, 730	26, 170

The completed vessel will include the weights of nozzles, a manway and reinforcing around the openings, which may total another 10–20%. The weights of these auxiliaries are stated in manufacturers' catalogs.

REFERENCES

- API Manual on Refinery Wastes, Am. Pet. Inst., Washington, D.C., 1969.
- D. Azbel and N.P. Cheremisinoff, Chemical and Process Equipment Design: Vessel Design and Selection, Butterworths, London, 1982.
- T.E. Belk, Effect of physical chemical parameters on coalescence, *Chem. Eng. Prog.* 61(10), 72-76 (1965).
- 4. D. Bradley, The Hydrocyclone, Pergamon, New York, 1965.
- 5. C. Branan, *The Process Engineers Pocket Handbook*, Gulf, Houston, 1976, 1983, Vol. 1, pp. 101-110, Vol. 2, p. 67.
- 6. C.L. Carpenter and D.F. Othmer, Entrainment removal by a wire mesh separator, *AIChE J.* 1, 549-557 (1955).
- R. Chuse, and S.M. Eber, Pressure Vessels: The ASME Code Simplified, McGraw-Hill, New York, 1984.
- G.A. Davies, G.V. Jeffrys, and M. Azfal, A new packing for coalescence and separation of dispersions, Br. Chem. Eng. 17, 709–714 (1972).
- D.R. Duros and E.D. Kennedy, Acid mist removal, *Chem. Eng. Prog.* 74(9), 70–77 (1978).
- F.L. Evans, Equipment Design Handbook for Refineries and Chemical Plants, Gulf, Houston, 1980, Vol. 2, pp. 153-165.
- R.V. Fronczak, Electrical desalting, in *Encyclopedia of Chemical Processing and Design*, Dekker, New York, 1983, Vol. 17, pp. 223-251.
- F.H. Garner, S.R.M. Ellis, and J.A. Lacey, Size distribution and entrainment of droplets, *Trans. Inst. Chem. Eng.* 32, 222-235 (1954).
- W.B. Hooper and L.J. Jacobs, Decantation, in Handbook of Separation Methods for Chemical Engineers (P. A. Schweitzer, Ed.), McGraw-Hi& New York, 1979, pp. 1.343-1.358.
- M.H. Jawad and J.R. Farr, Structural Analysis and Design of Process Equipment, Wiley, New York, 1984.
- G.V. Jordan, Coalescence with porous materials, *Chem. Eng. Prog.* 61(10), 64-71 (1965).
- 16. G.S. Laddha and T.E. Degaleesan, in *Handbook of Solvent Extraction* (Lo, Baird, and Hanson, Eds.) Wiley, New York, 1983, p. 125.
- D.M. Landis, Centrifugal coalescers, Chem. Eng. Prog. 61(10), 58-63 (1965).
- E.E. Ludwig, Applied Process Design for Chemical and Petrochemical Plants, Gulf, Houston, 1977, Vol. 1, pp. 144–180.
- J.H. Maas, Cyclone separators, in *Handbook of Separation Methods for* Chemical Engineers (P. A. Schweitzer, Ed.), McGraw-Hill, New York, 1979, pp. 6.10–6.17.

- W.J. Mead, Hoppers and bins and Tanks, in *Encyclopedia of Chemical Process Equipment*, Reinhold, New York, 1964, pp. 546-559 and 941–957.
- S. Middleman, Drop size distribution produced by turbulent pipe flow of immiscible liquids through a static mixer, *Ind. Eng. Chem.*, *Process. Des. Dev.* 13(1), 78-83 (1974).
- H.L. O'Brien, Petroleum Tankage and Transmission, Graver Tank and Mfg. Co., East Chicago, IN, 1951.
- O.C. Redmon, Cartridge type coalescers, Chem. Eng. Prog. 59(9), 87-89 (1963).
- 24. W. Reisner and M.E. Rothe, Bins and Bunkers for Handling Bulk Materials, Trans Tech Publication, Clausthal, Germany, 1971.
- E.R. Riegel, Tanks, in *Chemical Process Machinery*, Reinhold, New York, 1953, pp. 112-131.
- K. Rietema and C.G. Verver, Cyclones in Industry, Elsevier, New York, 1961.
- S.S. Safarian and E.C. Harris, *Design and Construction of Silos and Bunkers*, Van Nostrand Reinhold, New York, 1985.
- A.H. Selker and C.A. Schleicher, Factors affecting which phase will disperse when immiscible liquids are stirred together, *Can. J. Chem. Eng.* 43, 298–301 (1965).
- K.T. Semrau, Gas solid separation-cyclones, in *Chemical Engineers* Handbook, McGraw-Hill, New York, 1984, pp. 20.77-20.121.
- H.P. Sheng, J.R. Welker, and C.M. Sliepcevich, Liquid-liquid separations in a conventional hydroclone, *Can. J. Chem. Eng.* 52, 487-491 (1974).
- B. Sigales, How to design reflux drums, *Chem. Eng.* 157-160 (3 Mar. 1975); How to design settling drums, *Chem. Eng.*, 141-143 (23 June 1975); More on how to design reflux drums, *Chem. Eng.*, 87-90 (29 Sep. 1975).
- 32. A.J. Stepanoff, *Gravity Flow of Bulk Solids and Transport of Solids in Suspension*, Wiley, New York, 1969.
- L.C. Waterman, Electrical coalescers, Chem. Eng. Prog. 61(10), 51–57 (1965).
- 34. R.N. Watkins, Sizing separators and accumulators, Hydrocarbon Proc. 46(11), 253-256 (1967).
- O.H. York, Entrainment separation, in *Encyclopedia of Chemical Processing and Design*, Dekker, New York, 1983, Vol. 19, pp. 168-206.
- 36. F.A. Zenz, Cyclones, in Encyclopedia of Chemical Processing and Design, Dekker, New York, 1982, Vol. 14, pp. 82-97.

everal kinds of processes and equipment that do not fit readily into the categories of earlier chapters have achieved limited but nevertheless valuable applications in chemical processing. Many of them are offered by only a few manufacturers and the performance and economic data are highly proprietary. Here the objective is primarily to describe the principles involved, to point out the

19.1. MEMBRANE PROCESSES

Membranes, usually of organic polymers, can be constructed for separation of liquids from dissolved or suspended substances of a wide range of sizes smaller than those normally processed by the kind of filtration equipment described in Chapter 11. The full range of sizes of molecules and particles is illustrated in Figure 19.1. For small dissolved molecules, a phenomenon known as osmosis is the basis for a means of separation. Osmosis becomes manifest when two solutions at the same temperature and pressure but of different concentrations are separated by a semipermeable membrane, namely one that allows passage of the solvent but not the solute. Figure 19.2 illustrates this process. One theory of the action of a semipermeable membrane is that the solvent dissolves in the membrane at the face of higher concentration or higher partial pressure and is released at the other face where the concentration is lower.

The natural tendency is for the solvent to flow in the direction that will equalize the concentrations. It turns out, however, that if a certain pressure, called the osmotic pressure, is imposed on the more concentrated solution, flow the solvent can be forced in the main applications, and to refer to sources of more information. Specific manufacturers can be identified by reference to the usual directories of chemical process equipment (References, Chapter 1).

In addition, a number of other topics that also have been ignored in earlier chapters but are of interest to chemical processing will be covered in similarly brief fashion.

direction from the more concentrated to the more dilute solution. For the case of pure solvent on the low pressure side of the membrane, the osmotic relation is

$$\ln \gamma_{w} x_{w} = -\frac{1}{RT} \int_{0}^{P_{\text{osm}}} \bar{V}_{w} dP = -\frac{\bar{V}_{w}}{RT} P_{\text{osm}}, \qquad (19.1)$$

where γ is the activity coefficient, x is the **mol** fraction, and \overline{V} is the partial **molal** volume; subscript w identifies the solvent. For ideal solutions, the activity coefficient is unity. Since nonideality is of common occurrence, this equation may be used to find activity coefficients from measurements of osmotic pressure. Example 19.1 illustrates this process, and points out incidentally how rapidly the osmotic pressure falls off with increasing molecular weight of the solute.

Size ranges for membrane processing by reverse osmosis, ultrafiltration and microfiltration are shown in Figure 19.1. Reverse osmosis is effective in removing solvents away from dissolved molecules. Because of limitations in crushing strengths of membranes, pressures are limited to maxima of about 1000 psi

	T in the second s							
		Scanning	Electron Microsco	pe 🛛 Optical	Microscope	Visible T	o Naked Eye	
	lonic Bance	Motocui	ar Bases	Macro Molecular	Micro Particle 8	l Macr	o Particle Bange	
Micrometern	ionic nange		ar nange	Range		lange macr	o Particle hange	
(Log Scale)	0.0	01 0.	01 0	1 1	.0 1	b 11	00 10	00
Angstrom Units (Log Scale)	1 1 2 3 5 8 1 1 1 1 1 1		0 ************************************	0, -0, -0, -0, -0, -0, -0, -0, -0, -0, -)4 1, 2 3 5 8 1 1 1 1 1 1			2, 2,
Approx. Molecular Wt. (Saccharide Type - No Scale)	100 20	0 1000 10,000 20	.000 100,000	500,000				
	Aqueous	Salts	Carbon Black	Paint Pign	nent	L Human I	Hair	
Relative		Pyrogen			, Yeast	Cells	Beach Sand	
Size of	Metal Ion .		Virus		Bacteria	· · · ·	Mist	
Common			Taba	and Smake	Cool	Duct		
Materials			1004		Coar			
				Lung (Damaging Blood			
	s	ugars C	olloidal Silica /Partic	ies	Cells	Pollens		
	Atomic Radii	L	Albumin Protein		Milled	Flour		
PROCESS	REVERSE OSMOS (Hyperfiltration)	SIS	M	CROFILTRAT	ION			
SEPARATION		ULTRA	FILTRATION		PARTICLE FILTRATION			

Figure 19.1. Range of molecular weights and particle or droplet sizes of common materials, how they are measured, and the methods employed for their removal from fluids (Osmonics Inc.).



Figure 19.2. Diagram of osmotic behavior and the effect of solute concentration and molecular weight on osmotic pressure. (a) Osmotic-pressure behavior of solutions; ΔP_{osm} is the excess pressure on the solution required to stop flow of solvent through the semipermeable membrane. (b) Effects of solute concentration and molecular weight on osmotic pressure.

(68 atm). Flow rates of 2-200 gal/(sqft)(day) or 0.001-0.1 kg water/m² sec are attained in various units. Ultrafiltration operates at 1-10 atm differential and is effective for the molecular weight range of 1000-200,000 which includes many proteins, viruses, and bacteria. Ultra and micro filtrations somewhat overlap. Pressures for microfiltration are about 1 atm differential. Since these processes are relatively expensive, their applications are limited largely to analytical purposes and in water treatment for pharmaceutical manufacturing. Some specific applications are listed in Table 19.1.

MEMBRANES

The first commercially successful membrane was the anisotropic or asymmetric structure invented by Loeb and Sourirajan (1960; cited by Sourirajan, *Reverse* Osmosis, Academic, N.Y., 1970). It is made

of cellulose acetate and consists of a dense layer 0.2-0.5 μ m thick deposited on a porous structure 50–100 μ m thick with pores 0.1-1.0 μ m dia. The thin film has the desired solute retention property while offering little resistance to flow and the porous substructure offers little resistance to flow but provides support for the skin. The characteristics of available membranes for reverse osmosis and ultrafiltration are listed in Tables 19.2-19.4.

Hollow fiber membranes are primarily homogeneous. In use, their lower permeability is compensated for by large surface per unit volume of vessel. Fibers are 25-250 μ m outside dia, wall thickness 5-50 μ m. The cross section of a vessel for reverse osmosis may have 20-35 millionfibers/sqft and a surface of 5500-9000 sqft/cuft of vessel. Recently developed hollow fibers for gas permeation processes have anisotropic structures.

EQUIPMENT CONFIGURATIONS

Four principal kinds of membrane assemblies are in use:

- a. Tubular, in which the membrane is deposited either on the inside or the outside of porous tubes, most commonly inside for reverse osmosis and outside for ultrafiltration. Figure 19.3(a) shows a single-tube construction but units with 7 or 19 tubes in a single shell are made as standard items. Table 19.5 lists some available sizes. "Dynamic membranes" may be deposited on porous stainless steel tubes from a feed solution containing 50-100 mg/L of membrane-forming solution which consists of polyacrylic acid and hydrous zirconium oxide. Such a membrane can be deposited in 1 hr and replaced as quickly. Fluxes are very high; IOOgal/(sqft)(day) is shown in Table 19.6(a). Some applications are described by Turbak (1981, Vol. II, pp. 434-453).
- **b.** Plate-and-frame construction is shown in Figures 19.3(b) and (c). It is used more commonly for ultrafiltration. A related kind of equipment is the electrodialysis plate-and-frame equipment of Figure 15.21.
- c. Spiral wound assemblies are illustrated in Figure 19.4. They consist of a long envelope of membrane sealed on the edges and enclosing a porous material which serves as a channel for the flow of the permeate. The spacer for the feed solution flow channel is a **meshlike** material through which the solution is forced under pressure. The modules listed in Table 19.5 are 2-8in. dia, up to 3 ft long, and provide about 250sqft of membrane **surface/cuft** of vessel. Dimensions are shown on Figure 19.4(c). According to Table 19.6, reverse osmosis rates of 2500 gal/(sqft)(day) are attained.
- **d.** Hollow fiber assemblies function as one-ended shell-and-tube devices. At one end the fibers are embedded in an epoxy tubesheet and at the other end they are sealed. Overall flows of feed solution and permeate thus are in counterflow. Flow of permeate is into the tubes which takes advantage of the great crushing strengths of the small diameter fibers, and constitutes a "fail-safe" operation since collapse of fibers results in closure whereas bursting would result in leakage. The most serious drawback is some difficulty of cleaning. A widely used equipment of this type is illustrated in Figure 19.5.

APPLICATIONS

The greatest use of membranes is for reverse osmosis desalination of seawater and purification of brackish waters. Spiral wound and hollow fiber equipment primarily are applied to this service. Table 19.6 has some operating data, but the literature is very extensive and reference should be made there for details of performance and economics.

EXAMPLE 19.1

Applications of the Equation for Osmotic Pressure

a. The osmotic pressure of a sucrose solution is 148.5 atm at 20°C. The concentration is 1.43 kg sucrose/kg water, corresponding to a mol fraction 0.0700 of sucrose. The partial molal volume of water is approx 0.018 L/g mol. Accordingly, the activity coefficient of the water is

$$\gamma = \frac{1}{1. - 0.07} \exp\left[-\frac{0.018(148.5)}{0.082(293.2)}\right] = 0.9622,$$

The difference from unity appears to be small but is nevertheless significant. At this concentration, if the activity coefficient were unity, the osmotic pressure would be

$$P = -\frac{0.082(293.2)}{0.018}\ln(1 - 0.07) = 97 \text{ atm.}$$

which is considerably in error.

Because of the low energy requirements of separations by reverse osmosis, much attention has been devoted to other separations of aqueous solutions, at least on a laboratory scale, for instance, of ethanol/water. Membranes have been found that are moderately effective, but the main obstacle to the process is the very high pressures needed to remove water from high concentrations of ethanol against pure water on the low pressure side. A practical method of circumventing this problem is to replace the water on the low pressure side by a solution of sufficiently high concentration to allow the application of only moderate pressure. The case examined in Example 19.2 utilizes a solution of ethylene glycol on the low pressure side of such concentration that concentration of ethanol above the azeotropic composition can be achieved with a pressure of only 1000 psig. The glycol is easily separated from water by distillation.

TABLE 19.1. Examples of Applications of Ultrafiltration

(a) Applications Involving Retained Colloidal Particles

Material	Application			
Pigments and diS - persed dyes	Concn/purification of organic pigment slurries; separation of solvents, etc. from pigment/resin in electropaints; concn of pigments in printing effluents			
Oil-in-water emul- sion globules	Concn of waste oils from metal working/textile scouring; concn of lanolin/dirt from wool scouring			
Polymer lattices and dispersions	Concn of emulsion polymers from reactors and washings			
Metals/nonmetals/ oxides/salts	Concn of silver from photographic wastes; concn of activated carbon slurries; concn of inorganic sludges			
Dirt, soils, and clays	Retention of particulates and colloids in turbid water supplies; concn of fines in kaolin processing			
Microorganisms	Retention of microbiological solids in activated sludge processing; <i>concn</i>			
Plant/animal Cellu- lar materials	of viral/bacterial cell cultures; separation of fermentation products from broth; retention of cell debris in fruit juices, etc.; retention of cellular matter in brewery/distillery wastes			

b. The effect of molecular weight on ideal osmotic pressures of a variety of solutions containing 0.1 kg solute/kg water is demonstrated in this tabulation:

Mol Weight	Mol fraction x	ldeal Osmotic P (atm)		
58.5 (NaCl)	0.0374	50.9		
100	0.0177	23.9		
342 (sucrose)	0.00524	7.0		
1000	0.0018	2.4		
10,000	1.8(<i>E</i> = 4)	0.24		
100,000 (virus and protein)	1.8(E 5)	0.024		

Figure 19.1 identifies sizes of common molecules and particles. Clearly, osmotic pressures are essentially negligible for molecular weights above 10,000 or so.

GAS PERMEATION

Differences in rates of permeation of membranes by various gases are utilized for the separation of mixtures, for instance, of hydrogen from ammonia plant gas, of carbon dioxide from natural gas, and of helium from natural gas. The successful "Prism" process of Monsanto Co. employs hollow fibers of a porous polysulfone base coated with possibly a thin film of silicone rubber. The fibers are about 800 μ m outside and 400 μ m inside dia. They are housed in vessels 4–8 in. dia and 10–20 ft long and may contain 10,000– 100,000 fibers per bundle. A schematic of such a unit is in Figure 19.6(a). Pressures up to 150atm are allowable. A unit 4in. dia by 10 ft long was able to upgrade 290,000 SCFD of ammonia plant purge gas, making a product with 90% hydrogen and a waste of 20% hydrogen from a feed with 37% hydrogen.

(b) Applications Involving Soluble Macromolecules

Material	Application
Proteins and polypeptides	Concn/purification of enzymes; concn/purification of casein and whey proteins; concn/purification of gluten/rein; concn/purification of gelatin; concn/purification of animal blood; reten- tion of haze precursors in clear beverages; retention of antigens in antibiotics solu- tions; concn/purification of vegetable pro- tein extracts; concn/purification of egg albumen; concn/purification of fish protein extracts; retention of proteins in sugar
Polysaccharides and oligo- saccharides	Concn of starch effluents; concn of pectin extracts
Polyphenolics Synthetic water-soluble polymers	Concn/purification of lignosulphonates Concn of PVA/CMC desize wastes

TABLE 19.2. Data of Commercial Equipment for Reverse **Osmosis and Ultrafiltration**

(a) Equipment of Amicon Corp.

Diaflo®	Nominal mol wt cutoff	Apparent pore diam, Å	water flux, gal/ft²/day at 55 lb/in²
UM 05	500	2 1	10
UM2	1,000	30	20
UM 10	10.000	38	6 0
PM 10	10.000		550
PM 30	30,000	47	500
XM 50	50.000	66	250
XM 100A	100,000	110	650
XM 300	300,000	480	1300

(b) Equipment of Nucleopore Corp.

Specified		Nominal	Nominal	Typical flow rates at 10 lb/in* (gage), 4 <i>P</i> , 70°F		
pore size, μm	Pore-size range, µm	density, pores/cm ²	thickness, µm	Water, gal/(min)(ft²)	N2, ft³/(min)(ft²)	
8.0 5.0	6.9-8.0 4 3-5 0	1 × 10 ⁵ 4 × 10 ⁵	8.0 8.6	$144.0 \\ 148.0$	$138.0 \\ 148.0$	
3.0	2.5-3.0	$2 \times 10^{\circ}$ 2 × 10'	11.0 11.5	121.0	128.0	
0.8	0.64-0.80	3 × 10'	11.6	48.3	76.0	
0.4	0.32-0.40	1 x 10 ⁸	11.6	17.0	33.0	
0.1	0.16-0.20	3×10^{6} 3×10^{8}	5.3	1.9	5.3	
0.08	0.040-0.050 0.024-0.030	6×10^{9} 6×10^{9}	$5.4 \\ 5.4 \\ 5.4$	1.12 0.006	1.3 0.19	

(c) Equipment of Koch Membrane Systems (Formerly Abcor)

Membrane Type (1)	Nominal M. W . cutoff (2)	Max. Temp (°C) (3)	Range pH (4)	Maximum Pressure PSI (Kg/cm ²)	Configuration (5)
MSD-324*	1,500	90	1-13	150 (10.5)	S
HFK-132	3,500	90	1-13	150 (10.5)	S
HFK-131	5,000	90	1-13	150 (10.5)	S
HFD-300	8,000	80	2-12	150 (10.5)	Т
HFM-100	10,000	90	1-13	150 (10.5)	S,T
HFA-251*	15,000	50	2-8	150 (10.5)	T
HFM-180	18,000	90	1-13	150 (10.5)	S,T
HFM-163	18,000	60	2-12	150 (10.5)	S,T
HFP-276	35,000	90	1-13	150 (10.5)	S,T
MSD-181	200,000	90	1-13	150 (10.5)	S.T
MSD-400*	100.000	90 1	I-13	150 (10.6)	S,T
- M S D - 4	4 0 5 2 3 0,000	90	1-13	150 (10.5)	S,T
MMP-406'	0.2 Microns	9 0	1-13	150 (1	0.5) S.T ;
MMP-404*	0.4 Microns	90	1-13	150 (10.5)	S,T
MMP-516*	2 Microns	90	1-13	150 (10.5)	S,T
MMP-407'	2-3 Microns	90	1-13	50 (3.5)	S,T
MMPSOO	1-2 Microns	90 İ	1-13	50 (3.5)	S,T
MMP-602	2-3 Microns	90	1-13	50 (3.5)	S

 Membranes beginning with "H" designation are stock items.
(* = hydrophyllic)
The nominal molecular weight (M. W.) cutoff is provided as a guide to the relative pore size for these membranes. Since many factors influence the actual MW cutoff, tests must be run to confirm retention for any specific confirm. application. (3) At pH -6. (4) At 25°C. (5) F = Flat Sheet

S = Spiral T = Tubular. Because of the long, narrow configuration, the equipment appears to function in countercurrent mode. Other data of experiments with gas permeators as continuous columns appear in Figures 19.6(b) and (c); the original paper has data on other binary and some complex mixtures.

Permeability of a membrane is determined partly by gas diffusivity, but adsorption phenomena can exist at higher pressures. Separation factors of two substances are approximately in the ratios of their penneabilities, $\alpha_{AB} = P_{oA}/P_{oB}$. Some data of permeabilities and separation factors are in Table 19.7, together with a list of membranes that have been used commercially for particular separations. Similar but not entirely consistent data are tabulated in the *Chemical Engineers Handbook* (McGraw-Hill, New York, 1984, pp. 17.16, 17.18).

19.2. FOAM SEPARATION AND FROTH FLOTATION

Foams are dispersions of gas in a relatively small amount of liquid. When they are still on the surface of the liquid from which they were formed, they also are called froths. Bubbles range in size from about 50 μ m to several mm. The data of Table 19.8 show densities of water/air foams to range from 0.8 to 24 g/L. Some dissolved or finely divided substances may concentrate on the bubble surfaces. Beer froth, for instance, has been found to contain 73% protein and 10% water. Surface active substances attach themselves to dissolved materials and accumulate in the bubbles whose formation they facilitate and stabilize. Foam separation is most effective for removal of small contents of dissolved impurities. In the treatment of waste waters for instance, impurities may be reduced from a

content measured in parts per million to one measured in parts per billion. High contents of suspended solids or liquids are removed selectively from suspension by a process of froth flotation.

FOAM FRACTIONATION

Some dissolved substances are attracted to surfactants and thus are concentrated and removed with a foam. Such operations are performed in batch or continuous stirred tanks, or in continuous towers as in the **flowsketch** of Figure 19.7. Compressed air may be supplied through a sparger or ambient air may be drawn into a high speed rotating gas disperser. Improved separation is achieved by staged operation, so that a packed tower is desirable. Moreover, packing assists in the formation of a stable foam since that is difficult to do in an empty tower of several feet in diameter. Larger contents of surfactant usually are needed in large towers than in laboratory units. In pilot plant work associated with the laboratory data of Table 19.8, a tower 2 ft square by 8 ft high was able to treat 120 gal/hr of feed. The laboratory unit was 1 in. dia, so that the gas rate of 154 cm³/min of Table 19.8 corresponds to a superficial gas velocity of 1.1 ft/min.

Most of the work on foam fractionation reported in the literature is exploratory and on a laboratory scale. A selected list of **about** 150 topics has been prepared with literature references by Okamoto and Chou (1979). They are grouped into separation of metallic ions, anions, colloids, dyes and organic acids, proteins, and others.

Stable foams that leave the fractionator are condensed for further processing or for **refluxing**. Condensation may be effected

(a) Cellulose Acet	ate Membranes				
Membrane	Manufacturer	Volumetric flow rate (1/m ² d)	NaCI retention (%)	Pressure (bar)	NaCl concentration (%)
CA*					
(Tubing & Flat) CA′	KALLE	500 - 2500	98.6-60	40	0.5
(Flat) CA [•]	DDS***	350 – 220	99-78	42	0.0s
(Tubing)	PATERSON Candy Int.		97-so	40	0,5
CA•					
(In-situ cast"; pipe) CA•	ABCOR	450 - 900	98-90	4 2	0.5
(Wound module)	univ. Oil Prod. (GULF)	650	98	70	3,5
(Hol low fiber)	Monsanto	130	94	18	3.5
{Hollow fiber) CTA**	DOWCHEM.COMP.	50	99.5	70	3.5
Ultra-thin wound <i>module</i>	univ. Oil Prod. (gulf)	550	99.3	70	3.5

TABLE 19.3. Properties of Membranes for Reverse Osmosis

(A. Walch, Proceedings, Membrane Conference, Lund, Sweden, 1976; Ullmann's Encyclopedia of Chemical Technology, Verlag Chemie, Weinheim, 1978, Vol. 16, pp. 515-535).

* CA: Cellulose-2,5-acetate, • * CTA: Cellulosetriacetate, • ** De Danske Sukkerfabriken

636 OTHER TOPICS

TABLE 19.3—(continued)

(b) Other Kinds of Membranes

			Volumetric f bw	Molecular	
Membrane	Туре	Mfgr	(ml/min • cm² • bar)	range	Conditions
Polyelectrolyte (composite)	um 05 -um 10	AMICON	0,005 - 0,08	500 – 10000	max. 4 bar; 50°C; pl·l 4 – 10
Polysulfone (asymmetric hollow fibers)	PM 10 -PM 30	Λ ΜΙCON	0,4 - 13	10 000 - 30 000	max. 4 brr; 115°C; pH 0-12
Hod acrylics (asymmetric hollow fibers)	XM 50 -XM 300	AMICON	0.6 - 2.0	50000-300000	max. 2 bnr; 70°C; pH o-12
Polyelectrolyte (composite)	PSAC. PSDM	MILLIPORE	0,18-0,15	1000-40000	max. 7 bnr; 50°C; pH 4-10
Cellulose triacetate(60µm) PEM	GELMAN	0,004	50000	max. 80°C:pH 1-10
Regenerated cellulose (symmetric hollow fibers)	BF 50	DOW CHEM. COM	P. 🛥	5000	max. 70°C; pH 1 – 12
Cellulose acetate	BF 80	DOW CHEM. COM	P. 🖛	30000	max. 50°C: pH 2-8
(symmetric hollow fibers)					·
Regenerated cellulose (100 um)	115	Sartorius	0.001 -0,1	20000-160000	max. 80°C; pH 1-12
Cellulose acetate (100 pm'	117	SARTORIUS	0,001 -0.1	20000 – 160000	max. 80°C; pH 2-8
Cellulose nitrate(I00 µm)	121	SARTORIUS	0.005 - 0.02	10 000 - 50 000	max. 80°C; $pH 1 = 10$
Zr02/Carbon	Ucarscp	UNION CARBIDE	0,02	30 000 etc.	max. 10 bar;
(asymmetric tube bund 1 e) Polyamide, polyimide (asym, composite, hollow).	вм 10- вм 500	BERGHOF	0,004 - 0,7	1000-50000	$100^{\circ}C; pH = 14$ max. $100^{\circ}C;$ pH 2-10
Co-polyacryloni tri le (sym-		RHONE-POULENC	0.017	70000	max. 2 bar; pH 2-12
Cellulose acetate (asym)	800 - 500	DDS***)	0,01 - 0,04	6000-60000	max. 20 – 100 bar; 30 – 50 °C; pH 3-7
Polysulfone (asym)	GR 5, 6. 8	DDS***)		10000-20000	max. 80°C; pH O-14
Polyacrylonitrile (asym)	ғрв – GPA	Dorr-Oliver	0,1 -0.35	1200-100000	max, 2-4 bar; 70-80°C; pH1-13
Cellulose acetate (asym)	T 2/A - T 5/A	PCI®		1000 - 20 000	max. IO-25 bar; 30-50°C; pH 3-7
Cellulose derivative	T6/B	PCI ⁰⁾		120000	max. 10 bar; 60°C; pH 2 – 11
Cellulose acetate (asym)	HFA/100 	ABCOR	0,005 - 0.3	15000-50000	max. 14-100 bar; 30 – 50°C; pH 3 - 7
Polyamide (asym)	•	ABCOR		50 000	
Cellulose acetate (asym)	UF6 -UF 100	Kalle	0.01 -0.1	6000 – 100000	3 - 10 bar; $30 - 50^{\circ}$ C; pH 3-7
Polyamide (asym)	PA 40 - PA 100	KALLE	0,06 0,1	40 000 100000	3-6 bar; max. 70°C; pH I − 11

• * BM: Berghof-Membran ** DDS: DE DANSKE SUKKERFABRIKEN

PCI: PATERSON CANDY INTERNATIONAL

by a blast of steam, by contact with a hot surface, by chemical antifoaming agents, sonically or ultrasonically, or by contact with a high speed rotating disk as appears in the flowsketch, Figure 19.7.

FROTH FLOTATION

Finely divided solids or immiscible liquids can be made to adhere to gas bubbles and then can be removed from the main liquid. Affinity of a solid for an air bubble can be enhanced with surfactants which adhere to the surface of the solid and make it nonwetting. The main application of froth flotation is to the separation of valuable minerals from gangue. Ores of Cu, Zn, Mo, Pb, and Ni are among those commercially preconcentrated in this way. Reagent requirements of each ore are unique and are established by test. A large amount of experience exists, however, and information is supplied freely by reagent manufacturers. Some recipes are given with descriptions of flotation processes in books on mineral dressing, for example, that of Wills (1985).

Promoters or collectors give the mineral the water-repellent coating that will adhere to an air bubble. Frothers enhance the formation and stability of the air bubbles. Other additives are used to control the pH, to prevent unwanted substances from floating, or to control formation of slimes that may interfere with selectivity.

Air is most commonly dispersed with mechanical agitation. Figure 19.8 illustrates a popular kind of flotation cell in which the gas is dispersed and the pulp is circulated with impellers. Such vessels have capacities of 300-400 cuft. Usually several are connected in series as in Figure 19.8(b). The froth is removed from

	Membrane	Mígr	Volumetric flow rate (1/m ² d)	NaCl retention (%)	Press (bar)	NaCl concen (%)	Stability
Polyamide	Aromatic polyamide	DUPONT	50	9 5	28	0,15	max. 35°C
	(asym hollow fiber 89) Copolyamide asym hollow fiber Bl0	DUPONT	3 0	98.5	56	3.0	pH = 11 max. 30 C pH 5-9
	Polyamide hyd razide asym wound module	DUPONT	500	99	70	3.5	
	Polypi perazinamide asym Iibers	Montecatini	600	98	80	1	chlor-
	Aromatic polyamide ultrathin wound modules	UNIV. OIL PROD.	1700	98.9	70	3.5	-
Polyurea	Ethyleneimine, toluylene" diisocyanate NS 100 ultrathin wound module	NORTH STAR Res. Inst. (Univ. Oil prod.)	500	99.6	70	3.5	pH 2-12
Polyfuran	Furfuryl alcohol. H ₂ SO4,;NS 100 ultra thin	North Star Res. INST .	1000	99.6	70	3.5	pH 2-12
Poiyether	Polyphenylene öxide sulfone (5 pm on polypropylene tube	GENERAL ELECTRI	c 1 500	84	77	0.1	max. 54°C
	Polysulfone	BUANT DATUENCE	`	0 0	60	35	may 60 C
Polyhetero-	Polybenzimidazole	CELANESE	800	95	41	0.5	max.oo o
aromatics	asym tube Polybenzimidazole asym hollow fiber	CELANESE	50	99.4	70	3.5	
	Polyimide, methoxyl 10 p m	BATTELLE (BRD) 2	2 0	99.1	100	3.5	
Fluoropolymer	Naflon	DUPONT	3	85	100	3.5	
	(Suir, 250µm) Permion (pyridine, 25 pm)	RAI- Res. Cow.	1	98.8	loo	3.5	
Inorganic	Glass hollow fiber	STASFORD RES. INS	ат. IS	83	102	0.5	
membranes	Glass hollow fiber Graphite, oxidized	WESTINGHOUSE (UNION CARBIDE COMP.)	50	98 80	41	0.5	
Dynamically	Zr0 ₂ /poly acry lic acid	OAK RIDGE NAT. LAB.	5000	90	70	0.3	
membranes	Polyacrylic acid	UNIV. OIL PROD. (GULF)	2000	8 0	102	0.3	
Vinyl polymers	Polyvinylpyrrolidone cross linked 75 pm	Univ, Oil prod. (GULF)	0,5	99	34	0.8	
	Polyvinyl alcohol	PRINCETON UNIVERSITY	3	93	4 2	0.6	
	Polyvinyl carbonate	AEROJET- GEN. CORP.	3	94	102	3.5	
	Vinyl copolymer 10 μm	BATTELLE (BRD)	70	96	100	0.5	

TABLE 19.4. Properties of Membranes for Ultrafiltration

(A. Walch, Proceedings, Membrane Conference, Lund, Sweden, 1976; Ullmann's Encyclopedia of Chemical Technology, Verlag Chemie, Weinheim, 1978, Vol. 16, pp. 515-535).



Figure 19.3. Tubular and plate-and-frame membrane modules for reverse osmosis and ultrafiltration. (a) Construction and flow pattern of a single 1 in. dia tube with membrane coating on the inside; in Table 19.4, the "Ultracor" model has seven tubes in a shell and the "**Supercor**" has 19 *[Koch Membrane Systems (Abcor)]*. (b) Assembly of a plate-and-frame ultrafiltration module (Danish Sugar Co.). (c) Flow in a plate-and-frame ultrafiltration module.

each cell as it is formed, but the pulp goes through the battery in series. The froth is not highly stable and condenses readily without special provisions as it overflows. Since some entrainment of gangue occurs, usually it is desirable to reprocess the first froth. The flowsketch of Figure 19.9 illustrates such reprocessing. The solids to the first stage are ground here to -65 mesh, which normally is fine enough to release the mineral, and to -200 mesh in the final stage.

Total residence time in a bank of cells may range from 4 to 14min. A table of approximate capacities of several makes of flotation cells for a pulp with 33% solids of specific gravity = 3 is given in the Chemical *Engineers' Handbook* (McGraw-Hill, New York, 1984, p. 21.49); on an average, an 8-cell bank with 4-min holdup has a capacity of about 1.5 tons solid/(hr) (cuft of cell) and a power requirement of about 0.6 HP/(cuft of cell).

The chief nonmineral application of froth flotation is to the removal or oil or grease or fibrous materials from waste waters of refineries or food processing plants. Oil droplets, for instance, attach themselves to air bubbles which rise to the surface and are skimmed off. Coagulant aids and frothers often are desirable. In one kind of system, the water is saturated with air under pressure and then is pumped into a chamber maintained under a partial vacuum. Bubbles form uniformly throughout the mass and carry out the impurities. The unit illustrated in Figure 19.10 operates at 9 in. mercury vacuum and removes both skimmed and settled sludges. Because of the flocculation effect it is able to process waste water at an enhanced rate of about 5000 gal/(sqft)(day) instead of the usual rate of 800-1000.

In another application, particles of plastics in waste stream are chopped to diameters of 5 mm or less, passed through flotation cells containing proprietary surfactants, and removed as an air froth.

19.3. SUBLIMATION AND FREEZE DRYING

Sublimation is the transformation of a solid directly into vapor and desublimation is the reverse process of condensing the vapor as a

Module	Length	Membrane Area/Module
Tubular UF 1"	5 ft.	1.1 sq . ft.
(2.5 cm) dia.	(1.5 m)	(.10 ma)
Tubular UF 1″	10 ft.	2.2 sq. ft.
(2.5 cm) dia.	(3 m)	(.20 m²)
ULTRA-COR [™] uF ™um	10 ft.	7.4 sq. ft.
.5″ (1.27 cm) dia.	(3 m)	(.68 m²)
SUPER-COR™	10 ft.	24 ft.2
UF Tubes	(3 m)	(2.2 m²)
Tubular RO ½//	12 ft.	48 sq. ft.
(1.27 cm) dia.	(36 m)	(4.4 m²)
Spiral UF 2"	1.2 ft.	2.5 sq. ft.
(5 cm) dia.	(.36 m)	(.23 m²)
Spiral UF 4″	3 ft.	35-60 sq. ft.
(10 cm) dia.	(.9 m)	(3.2-5.5 m²)
Spiral UF 8"	3ft.	150-250 sq. ft.
(20 cm) dia.	(.9m)	(13.923 m ²)
Spiral RO 4"	3 ft.	60 sq. ft.
(10 cm) dia.	(.9 m)	(5.5 ma)

TABLE 19.5. Specifications of Spiral and Tubular Equipment for Reverse Osmosis and Ultrafiltration'

^eThe "Ultracor" model has 7 and the "Supercor" has 19 tubes/shell.

(Koch Membrane Systems, formerly Abcor).

solid. The term pseudosublimation is applied to the recovery of solid condensate from the vaporization of a liquid.

The goal of a commercial sublimation is the separation of a valuable material from nonvolatile ones at temperatures low enough to avoid thermal degradation. The preservation of cell structure (and taste) is a deciding factor in the choice of freeze drying, a special instance of sublimation, foods, pharmaceuticals, and medical products.

Only a few solids have vapor pressures near atmospheric at safe temperatures, among them CO,, UF_6 , $ZrCl_4$, and about 30 organics. Ammonium chloride sublimes at 1 atm and 350°C with decomposition into NH, and HCl, but these recombine into pure NH_4Cl upon cooling. Iodine has a triple point 113.5°C and 90.5 Torr; it can he sublimed out of aqueous salt solutions at atmospheric pressure because of the entraining effect of vaporized water.

Sublimation pressures down to 0.001 bar are considered feasible. At lower pressures and in some instances at higher ones, entrainer gas is used, usually air or nitrogen or steam. By such means, for instance, salicyclic acid is purified by sublimation at 150°C with an entrainer of air with sufficient CO, to prevent decarboxylation of the acid. At the operating temperature, the vapor pressure is only 0.0144 bar. Operating conditions corresponding to equilibrium in the sublimer appear in Figure 19.11. Equilibrium may be approached in equipment where contact between phases is intimate, as in fluidized beds, **but** in tray types percent saturation may be as low as 10%.

Among substances that are sublimed under vacuum are anthranilic acid, hydroxyanthraquinone, naphthalene, and β -naphthol. Pyrogallol and d-camphor distill from the liquid state but condense as solids. Several metals are purified by sublimation, for instance, magnesium at 600°C and 0.01-0.15 Torr.

The common carrier gases are air or nitrogen or steam. Condensate from a carrier usually is finely divided, **snowlike** in character, which is sometimes undesirable. Substances which are sublimed in the presence of a carrier gas include anthracene, anthraquinone, benzoic acid, phthalic anhydride, and the formerly mentioned salicylic acid.

A partial list of substances amenable to sublimation is in Table 19.9.

EQUIPMENT

The process of sublimation is analogous to the drying of solids so much the same kind of equipment is usable, including tray dryers (Fig. 9.6), rotary tray dryers (Fig. 9.8), drum dryers [Fig. 9.11(b)], pneumatic conveying dryers (Fig. 9.12), and fluidized beds (Fig. 9.13). The last of these requires the subliming material to be deposited on an inert carrier which is the fluidized material proper.

Condensers usually are large air-cooled chambers whose walls are kept clear with brushes or scrapers or even swinging weights. Scraped or brushed surface crystallizers such as Figure 16.10(a) should have some application as condensers. When a large rate of entrainer gas is employed, a subsequent collecting chamber will be needed. One of the hazards of entrainer sublimation with air is the possibility of explosions even of substances that are considered safe in their normal states.

FREEZE DRYING

Preservation of cell structure, food taste, and avoidance of thermal degradation are reasons for the removal of moisture from such materials by sublimation. The process is preceded by quick freezing which forms small crystals and thus minimum damage to cell walls, and is likely to destroy bacteria. Some of the materials that are being freeze dried commercially are listed in Table 19.9(b).

The most advanced technique of quick freezing is by pouring the material onto a freezing belt. Before drying, the material is granulated or sliced to improve heat and diffusional mass transfer. These operations are conducted in cold rooms at about -46° C.

Sublimation temperatures are in the range of -10 to -40°C and corresponding vapor pressures of water are 2.6-0.13 mbar. Cabinet tray dryers are the most commonly used type. The trays are lifted out of contact with hot surfaces so the heat transfer is entirely by radiation. Loading of 2.5 lb/sqft is usual for foodstuffs. Drying capacity of shelf-type freeze dryers is 0.1-1.0 kg/(hr)(m² exposed surface). Another estimate is 0.5-1.6 lb/(hr)(sqft). The ice surface has been found to recede at the rate of 1 mm/hr. Freeze drying also is carried out to a limited extent in vacuum pans, vibrating conveyors, and fluidized beds. Condensers operate as low as -70°C.

Typical lengths of cycles for food stuffs are 5–10 hr, for bacterial pellets 2-20 hr, and for biological fluids 20-50 hr. A production unit with capacity of 500L may have 75 kW for refrigeration and 50 kW for heating. Conditions for the preparation of freeze dried coffee are preparation of an extract with 20-25% solids, freezing at $-25--43^{\circ}$ C, sublimation at approx. 200 Torr to a final final moisture content of 1-3%, total batch processing time of 6-8 hrs.

19.4. PARAMETRIC PUMPING

A class of operations has been devised in which the process fluid is pumped through a particular kind of packed bed in one direction for a while, then in the reverse direction. Each flow direction is at a different level of an operating condition such as temperature, pressure, or **pH** to which the transfer process is sensitive. Such a periodic and synchronized variation of the flow direction and some operating parameter was given the name of parametric pumping by Wilhelm (1966). A difference in concentrations of an **adsorbable**desorbable component, for instance, may develop at the two ends of the equipment as the number of cycles progresses.

TABLE	19.6.	Performanc	e Data	of Reverse	Osmosis	Membrane	Modules
(a) Data	ofB	Belfort (1964).				

Module design	Packing density (ft²/ft³)	Water flux at 600 psi (gal/ft ² day)	Salt rejection	Water output per unit volume (gal/ft ³ day)	Flow channel size (in.)	Ease of cleaning
Tubular Brine flow inside tube	30-50	10	Good	300-500	0.5-1.0	Very good
Brine flow out -side tube"	140	10	Good	1400	0.0-0. 125"	Good
Spiral wrap ^b Fiber	250	10	Good	2500	0.1	Fair
Brine flow inside fiber ^c	1000	5	Fair	5000	0.254	Fair
Brine Row out -side fiber	5000-2500	1-3	Fair	5000-7500	0.002	Poor
Flat plate^d Dynamic membrane'	35 5 0	10 100	Good Poor	350 5000	0.01-0.02 -0.25	Good Good

(b) Data of Crits [Ind. Water Eng., 20-23 (Dec. 1976-Jan. 1977)]

	Tri-acetate hollow fibers	Polyamide hollow fibers	Cellulose acetate spiral-wound
Module sizes and Row, gal/day at 400 lb/in ²	5 x 48 in, 4000 gall day; 10 x 48 in, 20,000 gal/day	4 x 48 in, 4200 gal/day (1); 8 x 48 in, 14,- 000 gal/day (1)	4 in x 21 ft (6), 4200 gal/day; 8 in x 21 ft (6) 24,- 000 gal/day
Recommended operating pressure, lb/in² Flux, permeate rate, gal/	400	400	400
day/ft ²	1.5	2	15-18
Seals, pressure	2	2	12
Recommended max operating temp, °F	86	95	85
(guaranteed %			
rejection)	90	90	90
pH range	4-7.5	4-11 0.1 > pH 8.0	4-6.5
Chlorine tolerance	0.5-1.0	0.25 > pH 8.0	0.5-1.0
(relative—FI*)	FI-<4	FI<3	FI<15
Recommended influent			
quality	FI-<3	FI<3	FI<3
Permeate back pressure	75	75	0
Biological attack	15	13	0
resistance	Resistant	Most resistant	Least resistant
Flushing cleaning	Not effective	Not effective	Effective
Module casing	Epoxy-coated steel	FRP	Epoxy-coated steel and FRP
Field membrane			
replacement	Yes	No (future yes)	Yes

* FI = fouling index.' (1) Initial flow. (6) Six modules per 4200 gal/day.

A schematic of a batch parametric pumped adsorption process is sketched in Figure 19.12(a), whereas Figure 19.12(b) shows the synchronized temperature levels and flow directions. At the start, the interstices of the bed and the lower reservoir are filled with liquid of the initial composition and with the same amount in both. The upper reservoir is empty. The bed is kept cold while the liquid is displaced from the interstices into the upper reservoir by liquid pumped from the lower reservoir. Then the temperature of the bed is raised and liquid is pumped down through the bed. Adsorption occurs from cold liquid and desorption to the hot liquid. For the system of Figure 19.12(c), the separation factor is defined as the ratio of concentrations of the aromatic component in the upper and lower reservoirs; very substantial values were obtained in this case. Data of partial desalination of a solution with an ion exchange resin are in Figure 19.12(d), but here the maximum separation ratio is only about 10.

An intermittent-simulated continuous-operation is described with Figure 19.13. Feed input and withdrawals of products are



Figure 19.4. The spiral wound membrane module for reverse osmosis. (a) Cutaway view of a spiral wound membrane permeator, consisting of two membranes sealed at the edges and enclosing a porous structure that serves as a passage for the permeate flow, and with mesh spacers outside each membrane for passage of feed solution, then wound into a spiral. A spiral 4 in. dia by 3 ft long has about 60 sqft of membrane surface. (b) Detail, showing particularly the sealing of the permeate flow channel. (c) Thickness of membranes and depths of channels for flows of permeate and feed solutions.



Figure 19.5. The "Permasep" hollow fiber module for reverse osmosis. (a) Cutaway of a DuPont "Permasep" hollow fiber membrane module for reverse osmosis; a unit 1 ft dia and 7 ft active length contains 15–30 million fibers with a surface area of 50,000–80,000 sqft; fibers are 25-250 μ m outside dia with wall thickness of 5-50 μ m (DuPont Co.). (b) The countercurrent flow pattern of a "Permasep" module.

EXAMPLE 19.2 Concentration of a Water/Ethanol Mixture by Reverse Osmosis

The pressure required to drive water out of mixtures of various concentrations of alcohol against pure water at 30°C is calculated from the osmotic equation

$$1 - x_{\rm alc} = \exp\left[-\frac{0.018P}{0.082(303.2)}\right]$$

with the results:



accomplished with periodic openings and closings of valves without shutting down the equipment at any time. Other modes of operation also can be devised.

Theoretical studies also have applied this cycling principle to liquid-liquid extraction processes with immobilized solvents, and to reversible chemical reactions. Quite comprehensive reviews of the literature of cycling zone separations have been made by Sweed (1972), Wankat (1974), Wankat et al., and (1976).

Although parametric pumping appeared on the academic scene in 1966, no commercial installations appear to have been made, at least no widely publicized ones. Periodic heating and cooling of beds of granules or even of periodic compression and decompression of the process gas appear to be serious economic obstacles.

19.5. SEPARATIONS BY THERMAL DIFFUSION

Separation of mixtures based on differences in thermal diffusivity at present are feasible only for analytical purposes or for production on a very small scale of substances not otherwise recovered easily. Nevertheless, the topic is of some interest to the process engineer as a technique of last resort. It appears that the pressures needed to make higher than azeotropic composition are beyond the strength of available membranes. A pressure of **1000 psi** (68 atm) is feasible. With this pressure the concentrations of solute on the two sides of the membrane are related by

$$\frac{1-x_1}{1-x_2} = \exp\left[-\frac{0.018(68)}{0.082(303.2)}\right] = 0.9520$$

whence

 $x_2 = 1 - (1 - x_1)/0.9520$

As long as the mol fraction of solute on the low pressure side is kept above the value given by this equation, water can be driven from the side with mol fraction x_1 across the membrane. The solute on the low pressure side should be one that is easily separated from water and any alcohol that may bleed through. Ethylene glycol is such a material which also has the advantage of a relatively low molecular weight, 62. The required minimum concentrations of glycol corresponding to various alcohol concentrations on the high pressure side (68 atm) are tabulated.

Wt % Alcohol	Mol Fraction Alcohol	Moi Fraction Glycol	Wt % Glycol
10	0.0417	≥0	≥0
50	0. 261	≥0.2447	≥52.74
90	0. 779	20. 7679	291.93
96	0. 9038	20. 8989	≥96.84

The flowsketch shows a feed stream consisting of 100 kg/hr alcohol and 900 kg/hr of water, and making a stream with 96% alcohol. If pure glycol is charged countercurrently at the rate of 106.9 kg/hr, the % glycol at the point in the column where the alcohol content is 50% will be 52.74%, which is high enough to ensure that water can be driven out by a pressure of 68 atm. Beyond this point also, the content of glycol will be high enough to ensure transfer of water out of the alcohol solution. The aqueous glycol will be distilled and recycled. A small increase in its amount will permit some water to be present in the recycle stream.

In a vessel with a temperature gradient between a hot and cold surface, a corresponding concentration gradient of a fluid likewise can develop. The substance with the smaller molecular volume usually concentrates in the high temperature region, but other factors including that of molecular shape also affect the relative migrations of components of mixtures. Thus, the **sequence** of separation of hydrocarbons from hot to cold regions **generally** is: light normal paraffins, heavy normal paraffins, naphthenes and monocyclic aromatics, and bicyclic aromatics. Isotopes with small differences in molecular weights were the first substances separated by thermal diffusion, but isomers which have identical molecular weights also are being separated.

The basic construction of a horizontal thermal diffusion cell is sketched in Figure 19.14(a). When gases are to be separated, the distance between the plates can be several mm; for liquids it is a fraction of a mm. The separation effects of thermal diffusion and convection currents are superimposed in the equipment of Figure 19.14(b), which is called a thermogravitational or Clusius-Dickel column after the inventors in 1938. A commercially available column used for analytical purposes is in Figure 19.14(c). Several such columns in series are needed for a high degree of separation.





Figure 19.6. Gas permeation equipment and performance. (a) Cutaway of a Monsanto "Prism" hollow fiber module for gas separation by permeation. (b) Flowsketch of a continuous column membrane gas separator. (c) Composition profiles of a mixture of CO, and O, in a column 5 m long operated at total reflux [Thorman and Hwang in (*Turbak, Ed.*), Synthetic Membranes II, *American Chemical Society, Washington DC, 1981, pp. 259–279*].

644 OTHER TOPICS

TABLE 19.7. Data of Membranes for Gas Permeation Separation

(a) Permeabilities of Helium, Nitrogen, and Methane in $Sev{\circ}ral$ Membranes at $20^\circ\!C$

	$P_{\rm s} \cdot 10^7 [\rm cm^2/s \ bar]$				
Membrane	He	N ₂ .	CH4		
Silicon rubber	17.25	11.25	44.25		
Polycarbonate	5.03	0.35	0.27		
Teflon FEP	4.65	0.19	0.11		
Natural rubber	2.70	0.79	-		
Polystyrene	2.63	0.17	0.17		
Ethyl cellulose	2.33	0.21	0.83		
Polyvinyl chloride					
(plasticized)	1.05	-	0.15		
Polyethylene	0.75	0.14	_		
Polyvinylfluoride	0.14	0.0014	0.00048		

(b) Separation Factors $\alpha_{AB} = P_{0A}/P_{0B}$ for Three Mixtures

Membrane	He/CH ₄	He/O,	H ₂ /CH ₄
Polyacrylonitrile	60000	_	10000
Polyethylene	264	35.5	162
Polytetrafluoroethylene Regenerated cellulose	166 400	45 48	68.5
Polyamide 66	214	39	Time .
Polystyrene	14.6	5.5	21.2
Ethylcellulose	4 8	3.2	6.6

(c) Examples of Commercial Separations and the Kinds of Membranes Used

Separation process	Membrane
0, from air	Ethylcellulose, silicon rubber
He from natural gas	Teflon FEP, asymmetric cellulose acetate
H ₂ from refinery gas	Polyimide, polyethylene- terephthalate, polyamide 6
CO, from air	Silicon rubber
NH, from synthesis gas	Polyethylene terephthalate
H₂S from natural and refinery gas	Silicon rubber, polyvinylidene fluoride
H ₂ purification	Pd/Ag alloys

[Membranen, in Ullmann's Encyclopedia of Chemical Technology, Verlag Chemie, Weinheim, 1978, Vol. 16, p. 515. Many more data are collected by Hwang, Choi, and Kammermeyer, Separation Sci. 9(6), 461-478 (1974)].

Clusius and **Dickel** used a column 36 m long to make 99+% pure isotopes of chlorine in HCI. The cascade of Figure 19.15 has a total length of 14 m; most of the annular diameter is 25.4 mm, and the annular widths range from 0.18 to 0.3 mm. The cascade is used to recover the heavy isotope of sulfur in carbon disulfide; a production rate of a 90% concentrate of the heavy isotope of 0.3 g/day was achieved.

Separation of the hydrocarbon isomers of Table 19.10(a) was accomplished in 48 hr in the column of Figure 19.14(c) with 50°C hot wall and 20°C cold wall. The concentration gradient that develops in such a column is shown in Figure 19.14(d). The equilibrium terminal compositions depend on the overall composition, as indicated in Figure 19.14(e). Other kinds of behaviors also occur. Thus mixtures of benzene and cyclohexane are not sepa-

1'ABLE 19.8. Data of Foam Separation Experiments Made in a 1 in. Dia Column on a Waste Water Containing Radioactive Components and Utilizing Several Diferent Surfactants

		Flow ra	ites (cm³/min)		Feem	Average	
Surfactant	Surf. conc. (gm/liter)	Gas, V	foam, Q	Foam cond., <i>F</i>	density, ρ _f (gm/liter)	diameter, 6 (cm)	
Aerosol AY	6.5	154	176	0.197	1.12	0.06	
Alipal CO-436	0.375	154	186	0.950	5.10	0.05	
Alipal LO-529 Deriphat 170C	0.4 0.5	154 154	174 60	0.415 4.92	2.40 74	0.06 0.025	
lgepon CN-42 Tergitol 7	0.12 2.0	154 154	72 202	1.6 0.763	24 3.77	0.038 0.05	
Ultrawet SK	0.08	154	173	0.137	0.79	0.10	

(Davis and Haas, in Adsorptive Bubble Separation Techniques (Lemlich, Ed.), Academic, New York, 1972, pp. 279-297).

rated, nor can mixtures of benzene and octadecane when the latter is in excess.

Examples of separations of isotopes are in Table 19.10(b). The concentration of U-235 listed there was accomplished in a cascade of 2100 columns, each with an effective height of 14.6 m, inner tube 5 cm dia, gap 0.25 mm, hot surface 87-143°C, and cold surface 63°C, just above the condensation temperature at the operating pressure of 6.7 MPa. Although the process was a technical success,



File 19.7. Sketch of a foam fractionating column. Surfactants or other foaming agents may be introduced with the feed or separately at a lower feed point. Packing may be employed to minimize axial mixing.



Figure 19.8. The interaction of air and pulp in a froth flotation cell and a series arrangement of such cells: (a) Sectional schematic of flotation cell. Upper portion of rotor draws air down the standpipe for thorough mixing with pulp. Lower portion of rotor draws pulp upward through rotor. Disperser breaks air into minute bubbles. Larger flotation units include false bottom to aid pulp *flow.* (*WEMCO Division, Envirotech* Corp.). (b) A bank of three flotation cells. The floating concentrate is withdrawn continuously from each stage but the remaining pulp flows in series through the cells.

it was abandoned in favor of separation by gaseous diffusion which had only 0.7% of the energy consumption.

For separation of hydrocarbons, thermal requirements are estimated to range from 70,000 to **350,000 Btu/lb**, compared with heats of vaporization of 1.50 Btu/lb.

Although thermal diffusion equipment is simple in construction and operation, the thermal requirements are so high that this method of separation is useful only for laboratory investigations or for recovery of isotopes on a small scale, which is being done currently.

19.6. ELECTROCHEMICAL SYNTHESES

Electrolysis plays a role in the manufacture of some key inorganic chemicals on an industrial scale, but rather a minor one in the manufacture of organic chemicals. Chlorine, alkalis, metals, hydrogen, oxygen, and strong oxidizing agents such as $KMnO_4$, F_2 , and Cu_2O are made this way. Electroorganic processes of commercial or potentially commercial scale are listed in Table 19.11, which implies that much research is being done in pilot plants and may pay off in the near future. In the United States, the four large tonnage applications are to the manufacture of adiponitrile,



Figure 19.9. Flotation section of a flowsheet for concentration of 350 tons/day of a copper ore (*data of Pima Mining Co., Tucson, AZ*).



Figure 19.10. **Vacuator** of the "constant-level" type. The cylindrical tank with a dome-shaped cover is under a constant vacuum of about 9 in. of mercury. Sewage enters a central draft tube from which it is distributed by means of a flared-top section. Floating solids, buoyed up by fine air bubbles, are skimmed from the liquid surface and carried to a trough. Settled solids are removed from the bottom with a scraper mechanism. (Courtesy of Engineering News-Record).



Figure 19.11. Sublimation of salicyclic acid at 1 bar. Vapor pressures are 14.4 mbar at 150°C and 0.023 mbar at 40°C. The air rate shown corresponds to equilibrium in the sublimer, but in some kinds of vessels percent saturation may be as low as 10%. The conditions are those of Mullin [Crystallisation, *Butterworths*, London, 288 (1972)].

the Nalco process for lead tetraethyi, which is being phased out from the gasoline industry, the Miles process for dialdehyde starch, which is on standby until the demand picks up, and the 3M electrofluorination process for a variety of products.

Pros and cons of electrochemical processes are not always clear cut. In a few cases, they have lower energy requirements than conventional chemical methods **but** not usually according to the survey of Table 19.12. The Monsanto process for adiponitrile by electrochemical reduction of acetonitrile is an outstanding example; moreover, comparison of the performances of the original and improved cells [sketched on Figs. 19.16(e) and (f)] suggests the often great leeway in cell design. Small scale electrode processes frequently are handicapped because of the expense of developing

TABLE 19.9. Materials That May Be Purified by Sublimation or Are Being Freeze-Dried

(a) Substances Amenable to Purification by Sublimation^a

Aluminium chloride	Naphthalene
Anthracene	β-Naphthol
Anthranilic acid	Phthailic anhydride
Anthraquinone	α-Phthalimide
Benzanthrone	Pyrogallol
Benzoic acid	Salicylic acid
Calcium	Sulphur
Camphor	Terephthalic acid
Chromium chloride	Titanium tetrachloride
Ferric chloride	Thymol
lodine	Uranium hexafluoride
Magnesium	Zirconium tetrachloride

'Some others are mentioned in the text.

(b) Products which Are Being Freeze-Dried Commercially.

Foodstuffs	Pharmaceuticals	Animal Tissues and Extracts
Coffee extract	Antibiotics	Arteries
Fish and seafood	Bacterial cultures	Blood
Fruits	Serums	Bones
Fruit juices	Virus solutions	Hormones
Meat		Skin
Milk		Tumors
Tea extract		
Vegetables		

efficient components of cells such as electrodes, diaphragms, membranes, and electrolytes which usually can be justified only for large scale operation.

In comparison with chemical oxidations and reductions, however, electrode reactions are nonpolluting and nonhazardous because of low pressure and usually low temperature. Although electricity usually is more expensive than thermal energy, it is clean and easy to use. Electrolytic processes will become more attractive when inexpensive sources of electricity become developed.

ELECTROCHEMICAL REACTIONS

An equilibrium electrical potential is associated with a Gibbs energy of formation by the equation

$$E^{0} = -\Delta G^{0}/23.06n$$

where *n* is the number of gram equivalents involved in the stoichiometric equation of the reaction, ΔG^0 is in kcal/g mol, and E^0 is the potential developed by the reaction in volts. Thus, for the reaction $H_2O \rightleftharpoons H_2 + \frac{1}{2}O_2$ at 25°C,

 $E^0 = 54.63/(2)(23.06) = 1.18 \text{ V}$

and for $HCl \rightleftharpoons \frac{1}{2}H_2 + \frac{1}{2}Cl_2$ at 25°C,

 $E^0 = 22.78/23.06 = 0.99$ V.

Practically, reactions are not conducted at equilibrium so that amounts greater than equilibrium potentials are needed to drive a reaction. Major contributions to inefficiency are friction in the electrolyte and other elements of a cell and particularly the overvoltages at the electrodes. The latter are due to adsorption or buildup of electrolysis products such as hydrogen at the electrode surfaces. Figure 19.17(a) shows magnitudes of hydrogen **overvol**tages at several metals and several currents. The several contributions to voltage drops in a cell are identified in Figures 19.17(b) and (c), whereas Figure 19.17(d) indicates schematically the potential gradient in a cell comprised of five pairs of electrodes in series.

Electrochemical cells are used to supply electrical energy to chemical reactions, or for the reverse process of generating electrical energy from chemical reactions. The first of these applications is of current economic importance, and the other has significant promise for the near future.

FUEL CELLS

A few chemical reactions can be conducted and controlled readily in cells for the production of significant amounts of electrical energy at high efficiency, notably the oxidations of hydrogen or carbon monoxide. Some data of such processes are in Figure 19.18. The basic processes that occur in hydrogen/air cells are in Figure 19.18(a). Equilibrium voltage of such a cell is in excess of 1.0 V at moderate temperatures, but under practical conditions this drops off rapidly and efficiency may become less than 40%, as Figure 19.18(b) shows. Theoretical cell potentials for several reactions of fuel cell interest are in Figure 19.18(c); in theory at least, the oxidations of hydrogen and carbon monoxide are competitive. High temperatures may be adopted to speed up the electrode processes, but they have adverse effects on the equilibria of these particular reactions. Figure 19.18(d) shows the characteristics of major electrochemical fuel systems that have been emphasized thus far. Most of the development effort has been for use in artificial satellites where cost has not been a primary consideration, but



Figure 19.12. Batch parametric processing of solid-liquid interactions such as adsorption or ion exchange. The bottom reservoir and the bed interstices are filled with the initial concentration before pumping is started. (a) Arrangement of adsorbent bed and upper and lower reservoirs for batch separation. (b) Synchronization of temperature levels and directions of flow (positive upward). (c) Experimental separation of a toluene and n-heptane liquid mixture with silica gel adsorbent using a batch parametric pump. (Reprinted from Wilhelm, 1968, with permission of the American Chemical Society). (d) Effect of cycle time τ on reservoir concentrations of a closed system for an NaCl-H₂O solution with an ion retardation resin adsorbent. The column is initially at equilibrium with 0.05M NaCl at 25°C and $\alpha = 0.8$. The system operates at 5° and 55°C. [Sweed and Gregory, AIChE J. 17, 171 (1971)].



Figure 19.13. Parametric cycle operating intermittently in five periods; valves that are open each time are identified with their flow rates V_{i} , and the low and high temperature levels are identified with an asterisk in the proper column.

spinoff to industrial applications has some potential for the near future.

CELLS FOR SYNTHESIS OF CHEMICALS

Cells in which desired chemical reactions can be conducted and controlled are assemblages of pairs of anodes and cathodes between which the necessary potential difference is impressed. The regions near the electrodes may be separated by porous diaphragms to minimize convective mixing of the products formed at the individual electrodes. In recent years, semipermeable or ion-exchange membranes have been employed as diaphragms. In Figure 19.16(a), the membrane allows only Na^+ ions to pass so that the caustic that is made in the cell is essentially free of NaCl. In the mercury cell of Figure 19.16(b), no partition is necessary because the released Na dissolves in the mercury; the amalgam is reacted with water in an electrically neutral zone of the cell to make salt-free caustic. Because of pollution by escaped mercury, such cells have been largely phased out for production of salt-free caustic.

The same process sometimes can be performed efficiently in cells either with or without diaphragms. Figures 19.16(e) and (f) are for making adiponitrile by reduction of acetonitrile. In the newer design, Figure 19.16(f), the flow rate of the electrolyte is high

enough to sweep out the generated oxygen quickly enough to prevent reverse oxidation of the product.

Either parallel, called monopolar, or series, called bipolar, electrical connections can be made to the pairs of electrodes in a complete ceil. The monopolar types have individual connections to each electrode and thus require only individual pair potential to be applied to the cell assembly. The bipolar mode has electrical connections only to the terminal electrodes. One design such as Figure 19.16(f) has 48 pairs of electrodes in series and requires 600V. The equipment of Figure 19.19(a) also has bipolar connections. The voltage profile in such equipment is indicated schematically in Figure 19.17(d). Bipolar equipment is favored **because** of its compactness and, of course, the simplicity of the electrical connections. No adverse comments appear to be made about the high voltages needed.

Although the basic cell design shown schematically in Figures 19.16(a) and 19.19(d) is effective for many applications when dimensions and materials of construction are properly chosen, many special designs have been developed and used, of which only a few can be described here. For the cracking of heavy hydrocarbons to olefins and acetylenes, for instance, the main electrodes may be immersed in a slurry of finely divided coke; the current discharges from particle to particle generate the unsaturates. Only 100-200 V appears to be sufficient.

The most widely used brine electrolytic cells are the Hooker and Diamond Shamrock which are both monopolar, but bipolar designs like that of Figure 19.19(a) also are popular. That figure does not indicate the presence of a diaphragm but one must be used.

Rotating electrodes characterize the BASF cell of Figure 19.19(b), which is used for making adiponitrile. The cell described in the literature has 100 pairs of electrodes 40 cm dia spaced 0.2 mm apart. The rapid flow rate eliminates the need for diaphragms by sweeping out the oxygen as it is formed.

Lead alkyls are made by the action of Grignard reagents on lead anodes in the equipment of Figure 19.19(c). Lead pellets serve as the anode and are replenished as they are consumed. Several tubes 5 cm dia are housed in a single shell for temperature control and as required for capacity.

The simplest kind of cell construction, shown in Figure 19.19(d), suffices for the production of hydrogen by electrolysis of water and for the recovery of chlorine from waste HCI. The term filter-press cell is applied to this kind of equipment because of the layered construction. These two **electrolyses** are economically feasible under some conditions. Some details are given by Hine (1985).

It has been mentioned already that only a few inorganic and organic electrochemical processes have made it to commercial scale, but the potential may be there and should not be ignored. Recent surveys of the field and of the literature have been made by Hine (1985), Pletcher (1982), and Roberts et al. (1982).

19.7. FERMENTATION PROCESSING

Industrial fermentation is any process involving microorganisms that results in useful products. Among the useful microorganisms are molds, yeasts, algae, and bacteria. They are distinguished from plants and animals by being made of cells of only one kind. Although some kinds are grown as food, yeast or algae, for instance, the main interest here is in chemical manufacture with their assistance. This they accomplish by creating enzymes which catalyze specific reactions. In many respects biochemical processing is like ordinary chemical processing. The recovery and purification of biochemical products, however, often is a more demanding task and offers opportunities for the exercise of ingenuity and the



Figure 19.14. Construction and performance of thermal diffusion columns. (a) Basic construction of a thermal diffusion cell. (b) Action in a thermogravitational column. (c) A commercial column with 10 takeoff points at 6 in. intervals; the mean dia of the **annulus** is 16 mm, width 0.3 mm, volume 22.5 mL (Jones *and Brown*, 1960). (d) Concentration gradients in the separation of cis and trans isomers of 1,2-dimethylcyclohexane (Jones *and Brown*, 1960). (e) Terminal compositions as a function of charge composition of mixtures of cetane and cumene; time 48 hr, 50°C hot wall, 29°C cold wall (Jones *and Brown*, 1960).

application of techniques that are exotic from the point of view of conventional processing. A distinction also is drawn between processes that involve whole cells and those that utilize their metabolic products, enzymes, as catalysts for further processing. A brief glossary of biochemical terms is in Table 19.13.

Major characteristics of microbial processes are:

- a. The reaction medium is aqueous.
- **b.** The products are made in low concentration, rarely more than 5-10% for chemicals and much less for enzyme recovery.
- c. Reaction temperatures with microorganisms or isolated enzymes are low, usually in the range of $10-60^{\circ}C$, but the optimum spread in individual cases may be 5°C or less.
- **d.** With only a few exceptions, such as potable ethanol or glucose isomerate, the scale of commercial processes is modest, and for enzymes it is measured only in kilograms per day.
- e. Batch processing is used preponderantly, but so many conditions must he regulated carefully that computer control is common.

Because of the small scale of enzyme production, laboratory



Figure 19.15. The liquid thermal diffusion system for the recovery of heavy sulfur isotope in carbon disulfide. The conditions prevailing at the time after 90% ³⁴S is reached. Each rectangle in the cascades represents a column, each height being proportional to the length of the column. The two cascades have a combined height of 14 m, annular dia 25.4 mm, and annular width 0.18-0.3 mm. Production rate of 90% concentrate of ³⁴S was 0.3 g/day [W.M. Rutherford, Ind. Eng. Chem. Proc. Des. Dev. 17, 17-81 (1978)].

types of separation and purification operations are often feasible, including: dialysis to remove salts and some low molecular weight substances, ion exchange to remove heavy metals, ultrafiltration with pore sizes under 0.5 μ m and pressures of 1-10 atm to remove substances with molecular weights in the range of 15,009-1 million, reverse osmosis to remove water and to concentrate low molecular weight products, and gel permeation chromatography to fractionate a range of high molecular weight substances. Conventional processes of filtration and centrifugation, of drying by freezing or vacuum or spraying, and colloid milling also are used for processing enzymes.

PROCESSING

The three main kinds of fermentation processes are:

- **a.** Growth of microorganisms such as **bacteria**, fungi, yeasts and others as end products.
- **b.** Recovery of enzymes from cell metabolism, either intracellularly or as secretions, mostly the latter.
- c. Production of relatively low molecular weight substances by enzyme catalysis, either with isolated enzymes or with the whole cell.

Some industrial products are listed in Table 19.14. Chemical and fermentation syntheses sometimes are competitive, for instance, of ethanol, acetone, and butanol.

Enzymes are proteins with molecular weights in the range of 15,000–1,000,000 or so. In 1968, for instance, about 1300 were known, but only a few are of industrial significance. They are named after the kinds of reactions that they promote rather than to identify the structure which often is still unknown. Some kinds of enzymes are:

Amylase, which converts polysaccharides (starch or cellulose) to sugars.

Cellulase, which digests cellulose.

Glucose oxidase, which converts glucose to dextrose and levulose. Isomerase, which converts glucose to fructose.

Lipase, which splits fats to glycerine and fatty acids.

Protease, which breaks down proteins into simpler structures.

Biochemical manufacturing processes consist of the familiar steps of feed preparation, reaction, separation, and purification. The classic mode handles the microorganisms in slurry form in a stirred reactor. Enzyme-catalyzed processes also are performed primarily in stirred tanks, but when the enzymes can be suitably immobilized, that is, attached to solid structures, other kinds of reactor configurations may be preferred. Microbes also are grown in pans or rotating drums under moistened conditions, processes known as solid culture processing. Figure 19.20(a) shows the three modes of microbe culture. Processes that demand extensive handling of moist solids are practiced only on a small scale or when stirred tank action is harmful to cell structures. The process of Figure 19.20(b) consists largely of feed preparation steps.

OPERATING CONDITIONS

The optimum ranges of conditions for microbe growth or enzyme activity are quite narrow and must be controlled closely.

Concentration. A major characteristic of microbial growth and enzymatic conversion processes is low concentrations. The rates of these processes are inhibited by even moderate concentrations of most low molecular weight organic substances, even 1 g/L often being harmful. Nutrients also must be limited, for instance, the following in g/L:

Ammonia	5
Phosphates	10
Nitrates	5
Ethanol	100
Glucose	100

In the fermentation for ethanol, the concentration limit normally is about 8 wt % ethanol, but newer processes have been claimed to function at 10% or so. The search is on for microorganisms, or for creating them, that tolerate high concentrations of reaction products and higher temperatures.

Temperature. Most microbe metabolisms and enzymatic processes function well only in the range of $10-60^{\circ}$ C, but in particular cases the active spread of temperatures is only $5-10^{\circ}$ C. A classification of microorganisms that is sometimes made is with respect to peak activities near 15°C or near 35°C or near 55°C. The maximum heat effects of metabolic processes can be estimated from heats of formation when the principal chemical participants are known, for instance:

glucose-t ethanol, heat of reaction 0.10 kcal/g glucose, glucose+ CO, + H₂O, heat of reaction 3.74 kcal/g glucose.

	Vol	Mol		F comy	Final position, ol. %	Senara-
Components	%	wt.	Density	Тор	Bottom	tion, %
n-Heptane Triptane	<u>5</u> 8	100	0.6837 0.6900	95 95	10 90 j	75.4.
Isoöctane n-Octane	30	114	8:6819	58 42	40 60) 114 ·
1-Methylnaphthalene 2-Methylnaphthalene trans-1 2 Dimethylcyclo-	50 50	142 142	$1.0163 \\ 0.9905$	44.5 55.5	57.5 42.5	} 13.1
hexane cis-1,2-Dimethylcyclo-	40	112	0.7756	100	0 }	100
hexane	60	112	0.7963	0	100	
o-Xylene*	50	106	0.8799	9 8	1000	92
o-Xylene*	50	106	0.8739	10 0	819	80
m-Xylene	50	106	0.8669	50	50	0

TABLE 19.10. Examples of Separations by Thermal Diffusion (a) Hydrocarbon isomers

(A. L. Jones and G. R. Brown, in Advances in Petroleum Chemistry and Refining (McKetta and Kobe, Eds.), Wiley, New York, 1960, Vol. $\|l|$, pp. 43-76).

^ao-Xylene contains paraffinic impurity (see Fig. 9 in McKetta and Kobe).

(b) Isotopes

Working fluid	Isotope separated	mol % product	Phase	Single column (S) or cascade (C)	Investigator	Year
HCI	³⁵ Cl	99.6	Gas	S	Clusius and Dickel	1939
	³⁷ Cl	99.4				
Kr	⁸⁴ Kr	98.2	Gas	S	Clusius and Dickel	1941
	⁸⁶ Kr	99.5				
02	¹⁷ 0	0.5	Gas	С	Clusius and Dickel	1944
	¹⁸ O	99.5				
UF ₆	235U	0.86	Liquid	С	Manhattan Dist.	1945
N ₂	¹⁵ N	99.8	Gas	S	Clusius & Dickel	1950
Xe	¹³⁴ Xe	1	Gas	С	Clusius et al.	1956
	¹³⁶ Xe	99				
He	"Не	10	Gas	С	Bowring and Davies	1958
А	³⁶ A	99.8	Gas	С	ORNL	1961
	"А	23.2				
Ne	²⁰ Ne	99.99	Gas	С	ORNL	1961
	²² Ne	99.99	Cub			
Kr	⁷⁸ Kr	10	Gas	С	ORNL	1961
	⁸⁶ Kr	961		-		- /
He§	³ He	99	Gas	С	Mound Lab.[‡]	1962
Ne	²¹ Ne	33.9	Gas	С	ORNL	1963
CH4	¹³ C	90	Gas	С	Mound Lab.	1963
Xe	¹²⁴ Xe	4.4	Gas	С	ORNL	1964

† Oak Ridge National Laboratory, U.S. AEC, Oak Ridge, Tennessee.

‡ Mound Laboratory, US. AEC, Miamisburg, Ohio.

§ Feed not of normal abundance, contained 1 percent ³He from nuclear reaction. (Benedict et al., 1981).

TABLE 19.11. Electroorganic Synthesis Processes Now Applied Commercially or Past the Pilot Plant Stage

Product ^a	Raw Material^a	Company (country)	Scale	Type of Process
Commercialized				
Adiponitrile	Acrylonitrile	Monsanto (US) Monsanto (UK) Asahi (Japan)	10 ⁸ kg/yr 10 ⁸ kg/yr 2 X 107 kg/yr	Reductive coupling
p-Aminophenol	Nitrobenzene	(Japan) Holliday (UK)	Not available Not available	Reductive rearrangement
Anthraquinone	Anthracene	Holliday (UK)	Not available	Indirect oxidation
2,5-Dimethoxydihydrofuran	Furan	(Japan) BASF (West Germany)	Not available Not available	Oxidative addition
Fluorinated Organics	Hydrocarbons, aliphatic carboxylic acids, sulfonic acids, amines , etc.	Dia Ninpon (Japan) 3M (US)^b	Not available Not available	Anodic substitution
Cluconic Acid	Glucose	(India)	3×10^5 kg/yr	Oxidation of functional group
Glyoxylic Acid	Oxalic acid	(Japan)	Not available	Reduction of functional group
Hexahydrocarbazole	Tetrahydrocarbazole	BASF (West Germany)	Not available	Reduction
Piperidine	Pyridine	Robinson Bros. (UK)	1.2 X 10 ⁵ kg/yr	Reduction
Succinic Acid	Maleic acid	(India)	6 x 10 ⁴ kg/yr	Reduction
Hexadecanedioic Acid	Monomethylazelate	Soda Aromatic Co. (Japan)	Not available	Crum Brown-Walker ^C
Tetraethyl Lead	Ethylmagnesium halide	Nalco (US)	Not available	Anodic
Propylene Oxide	Propylene	BASF (West Germany) others in UK and West Germany	Past pilot-plant Past pilot-plant	Paired synthesis
4,4'-bis-Pyridinium Salts	Pyridinium salts	(Japan)	Past pilot-plant	Paired synthesis
Salicylaldehyde	Salicylic acid	(India)	Past pilot-plant	Reduction of functional group
Sebacid Acid Diesters	Adipic acid half esters	BASF (West Germany) (Japan) (USSR)	Past pilot-plant Past pilot-plant Commercial?	Crum Brown-Walker ^C
Benzaldehyde	Toluene	(India)	Past pilot-plant	Indirect oxidation [Mn(III)]
Dihydrophthalic Acid	Phthalic acid	BASF (West Germany)	Commercial?	Reduction
Hydroquinone or Quinone	Benzene	Several	Past pilot-plant	Paired synthesis or anodic oxidation + chemical reduction
Maltol	Furfuryl alcohol	Otsuka (Japan)	Past pilot-plant	Oxidation
Pinacol	Acetone	(Japan) BASF (West Germany)	Past pilot-plant	Reductive coupling

^a Formulas are given in Appendix A.

^bAdded by author.

^c Oxida tive coupling.

[M. M. Baizer, J. Appl, Electrochem. 10,285 (1980)].

Some of the energy is used to form the cell structure. Reactions catalyzed by enzymes may be either **endo**- or exothermic depending on the particular stoichiometry. Because of the diluteness of the solutions normally handled, temperature control is achieved readily. Stirred fermenters are provided with cooling jackets. Internal cooling oils are undesirable because of the difficulty of cleaning them. Fixed beds of immobilized enzymes do not lend themselves readily to jacket cooling, but in many instances the heat effect is so low that the temperature travel can be maintained within the required limits by adjustment of the feed temperature. Multitubular reactors with cooling medium on the shell side may be practical with enzymes immobilized on granules.

Sterilization. This is necessary to prevent the growth of foreign microorganisms. Air is sterilized adequately by the heat of

compression. Filters at the inlet remove oil and any microbes that may be present, and filters at the air outlet prevent backflow of foreign microorganisms. The inoculum is prepared under sterile conditions in the laboratory. The substrate is sterilized in an external vessel by holding it at 120°C or so for 1 hr or so.

Aeration. Since metabolism of microorganisms is an oxidative process, the substrate should be kept as nearly saturated as possible. At usual fermenter operating conditions the solubility of oxygen is **about** 0.03 mmol/L. When the content falls to 0.01 mmol/L, the growth rate falls to about one-half the maximum. Compressed air is introduced through spargers. Dispersion with high-speed agitators rarely is feasible because of possible mechanical destruction of cells. In some sensitive systems, all of the necessary agitation may be provided with an adequate air flow.

		kcal/kg		
Chemical		Electrochemical ^a	Chemical	
Adiponitrile	b	43,177 (10,520)*	65,808	C
Aniline Nitrobenzene route Phenol route	b b	36,172	13,919 16,736	с
Sorbitol Terephthalic Acid		9,649 17,382	958 700	
Phenol Methyl Ethyl Ketone	b	35,592 6,187	12,251 6,690	с
Melamine Hydroguinone	b b	30,159 52,739	3,233 15,472 30,814	c
Dichloroethane HCI route Cl2 route	b b	17,773	6,131 14,819	с

TABLE 19.12. Comparative Energy Requirements of Electrochemical and Chemical Processes

^eElectrochemical energy adjusted for generating plant efficiency. ^b Improved Monsanto process.

'Energy charged is for hydrocarbon raw materials (different compounds); other compounds begin with the same raw materials.

^d Chemical route energy given by Rudd et al.; others estimated by Beck et al.

T. Beck et al., A Survey of Organic Electrolytic Processes, ANL/OEPM 79-5, Electrochemical Technology Corp., 1979.

Agitation. The purpose of agitation is to keep the microorganisms in suspension, to maintain uniformity to eliminate concentration gradients and hot spots, and to improve heat transfer to the cooling jacket. Rules for the design of agitation systems are covered in Chapter 10. In vessels of a 1000 gal or more, a power input of about 10 HP/1000 gal and impeller tip speeds of 15–20 ft/sec are adequate, but the standard fermenter described in Table 19.15 is supplied with about four times this power.

pH. Biochemical processes are highly sensitive to hydrogen-ion

concentration. Most enzymes function best in the range of pH from 5 to 7, but some extremes are pepsin at pH of 1.5 and araginase at pH of 10. For classes of microorganisms, these ranges are common:

Complex	cells	6.5-7.5
Bacteria		4-6
Molds		3-7
Yeasts		3-6

Control of pH is accomplished by additions of dilute acid or alkali.



Figure 19.16. Basic designs of electrolytic cells. (a) Basic type of two-compartment cell used when mixing of **anolyte** and catholyte is to be minimized; the partition may be a porous diaphragm or an ion exchange membrane that allows only selected ions to pass. (b) Mercury cell for brine electrolysis. The released Na dissolves in the Hg and is withdrawn to another zone where it forms salt-free **NaOH** with water. (c) Monopolar electrical connections; each cell is connected separately to the power supply so they are in parallel at low voltage. (d) Bipolar electrical connections; 50 or more cells may be series and may require supply at several hundred volts. (e) Bipolar-connected cells for the Monsanto adiponitrile process. Spacings between electrodes and membrane are 0.8-3.2 mm. (f) New type of cell for the Monsanto adiponitrile process, without partitions; the stack consists of SO-200 steel plates with 0.0-0.2 mm coating of Cd. Electrolyte velocity of 1-2 m/sec sweeps out generated 0,.





Figure 19.16—(continued)

Zon **Concentration.** Heavy metals, particularly calcium, inhibit enzyme activity. The only feasible method of removing them is with ion exchange resins.

Foam Control. Fermentations tend to froth because **metabolites** have surfactant properties. Prevention commonly is by addition of **antifoam** agents such as oils, heavy alcohols, fatty acids, or silicones. High-speed rotating impellers destroy **bubbles by direct** impact and by throwing them against the wall of the vessel.

REACTORS

Stirred tanks are the chief kind of reactors for handling microorganisms or dissolved isolated enzymes, either as batch units or as **continuous stirred tank batteries.** When the enzymes are immobilized, a variety of reactor configurations is possible and continuous operation is easily implemented. The immobilization may be on granules or on sheets, and has the further advantage of making the enzymes reusable since recovery of dissolved enzymes rarely is feasible.

	Current Density (amp cm ⁻²)		nsity "2 ₎			
Metal	0	0.01	0.10		Amalgam cell	Diaphragm cell
Platinized Platinum	0.005	0.035	0.055		at 100 A/dm ⁻	at 25 A/dm ⁻
fron	6.68	0.56	0.82	Decomposition voltage	3.16 V	2.17 v
Smooth Platinum	0.09		0.39	Anode overvoltage. DSA	0.20 v	0.03 v
Silver	0.15	0.76	0.90	Cathode overvoltage	0.05 v	0.30 V
Nickel	0.21	0.65	0.89	Solution IR, including bubble effects	0.44 v	0.35 v
copper	0.23	0.58	0.82	Diaphragm		0.60 V
Lead	0.64	1.09	1.20	Metal hardware	0.05 v	0.20 v
Zinc	0.10	0.75	1.06			
Mercury	0.78	1.10	1.18	(Sum) Terminal voltage	3.91 v	3.65 V
	(;	a)		(b)	(continued

Figure 19.17. Overvoltage and distribution of voltage drops in cells (*Him, 1985*). (a) Overvoltage of hydrogen on some metals. (b) Voltage distribution in two kinds of cells for electrolysis of brine. (c) Variation of voltage distribution with current density in the electrolysis of **HCl**. (d) Schematic of voltage profile in a bipolar cell with five pairs of electrodes.



Figure 19.18. Data of electrochemical fuel cells. (a) Processes in a fuel cell based on the reaction between hydrogen and oxygen. (b) Voltage-current characteristic of a hydrogen-air fuel cell operating at 125° C with phosphoric acid electrolyte [*Adlhart*, in Energy Technology Handbook (*Considine*, Ed.), 1977, *p*. 4.61). (c) Theoretical voltages of fuel cell reactions over a range of temperatures. (d) Major electrochemical systems for fuel cells (*Adlhart*, in *Considine*, loc. cit., 1977, *p*. 4.62).

	. .	Operating	Electrode catalyst	Reactants			
Electrolyte	Current transport	temperature,		Fuel	Oxidant	State of development	
Aqueous potassium hydroxide (KOH)	он-	20-90	Nickel, silver, platinum metals	Hydrogen, hydrazine	Oxygen, scrubbed air, HeOg	Multikilowatt systems developed by several manufacturers.	
Aqueous sulfuric acid (H ₂ SO4)	H*	80-40	Tungsten carbide, platinum metals, carbon	Impure hydrogen*	Air	Long life demonstrated in laboratory cells.	
Concentrated phosphoric rfd (H ₃ PO ₄)	H.	70–175	Platinum metals	Impure hyd rogen *	Air	Multikilowatt system developed and larger systems in development. Long life has been demonstrated.	
Fused alkali carbonate	œ ,	600800	Nickel, silver	Impure hydrogen*	Air	Several months' lib' for small cells demonstrated.	
Stabilized zirconium oxide	0	700-1000	Bese metal oxides	impure hydrogen*	Air	10,000 hours' life demonstrated in single cells. Multikilowatt systems In design.	
			(d)				

Figure **19.18**—(continued)

Many aspects of the design of biochemical reactors are like those of ordinary chemical reactors. The information needed for design are the kinetic data and the dependence of enzyme activity on time and temperature. Many such data are available in the literature, but usually a plant design is based on laboratory data obtained with small fermenters. Standard sizes of such units range from 50 to 1000 L capacity. A sketch of a plant size fermenter and some of its auxiliaries is in Figure 19.21. Although not shown here, a bottom drive mechanical agitator usually is provided. The standard specification, Table 19.15, of one make of commercial fermenter includes a listing of the many openings that are required, as well as other general information.



Figure 19.19. Some special designs of electrolytic cells. (a) Glanor bipolar diaphragm-type cell assembly for chlor-alkali production (*PPG* Industries). (b) BASF capillary gap cell has 100 pairs of graphite plates with gaps of 0.2 mm used for adiponitrile synthesis; anodes are electroplated with lead dioxide *[Beck and Guthke, Chem. Ing. Tech.* 41, 943 (1969)]. (c) Principle of the shell-and-tube reactor for electrolytic oxidation of Grignard reagents to lead alkyls. Lead shot serves as consumable anode which is replenished continuously. Individual tubes are 5 cm dia by 75 cm long [Danly, Encycl. Chem. Technol. 8, 702 (1979)]. (d) Simple cells of the type used for electrolysis of HCl and water; voltage breakdown is shown in Figure 19.16(c).



Figure 19.19—(continued)

TABLE 19.13. A Biochemical Glossary

Microorganisms (microbes) are living cells, single or in multiples of the same kind, including bacteria, yeasts, fungi, molds, algae and protozoa. Their metabolic products may be of simple or complex structure

Fermentation is a metabolic process whereby microorganisms grow in the presence of nutrients and oxygen, sometimes in the absence of oxygen. The terms used are aerobic (in the presence of oxygen) and anaerobic (in the absence of oxygen)

Substrate consists of the nutrients on which a microorganism subsists or the chemicals upon which an enzyme acts

 $\it Enzymes$ are made by living cells, and are proteins with molecular weights ranging from about 15,000 to 1,000,000. They are able to catalyze specific reactions

Enzymes, immobilized, are attached to a solid support by adsorption or chemical binding or mechanical entrapment in the pores of a gel structure, yet retain most of their catalytic powers

-ase is a suffix identifying that the substance is an enzyme. The main part of the name describes the nature of the chemical reaction that can be catalyzed, as in **Cellulase**, an enzyme that catalyzes the decomposition of cellulose

TABLE 19.14. Industrial Products of Microbial and Related Processes

A.	Significant or marginal products Acetic acid
	Amino acids
	Butyric acid
	Citric acid
	Ethanol
	Fructose from glucose
	Glucose from starch
	Gluconic acid
	Methane
	Nucleotides (glutamic acid, guanyl acid, xanthyllic acid)
В.	Products under development or absolesced from microbial synthesis Acids: fumaric, lactic, malic, oxalic and some others Acetone Butanol Butanol
	Butaneuloi

Glycerine

Lipids

Polyalcohols and other substances

- C. Enzymes (extensive lists with properties and industrial suppliers are in the book by Godfrey and Reichelt, 1983).
- D. Antibiotics (Lists with major characteristics, sources and manufacturing methods are in, for example, the book of Bailey and Ollis, *Biochemical Engineering Fundamentals*, McGraw-Hill, New York, 1985).





Figure 19.20. Plowsketches of two processes employing fermentation. (a) Process for enzyme production, showing the use of growing trays, growing drums and stirred tank. Purification steps are the same for all three modes of culture growth. (b) Production of methane-rich gas by anaerobic digestion of finely divided waste solids in a 10–20% slurry. Residence time in the digester is *five* days [D.M. Considine (Ed.), Energy Technology Handbook, McGraw-Hill, New York, 1977].



- *1 Temperature controller and recorder la Resistance thermometer
- lb Control valve
- *2 Fermenter level
- *3 pH recorder and controller 3a pH electrode system 3b Control valve
- *4 Yeast concentration recorder
- *5 Recorder controller
- 6 Phosphate feed rotameter
- 7 Water feed rotameter
- 8 Trace element feed rotameter

- s 9 Foam controller
 - 9a Foam detector
 - 9b Control valve
- 10/11 Dosage control unit
- 10a Molasses feed
- 1 la Nitrogen feed
- 12 Rotor jet
- 12a Power unit
- *13 Air controller recorder
- 13a Venturi
- 13b Power operated air control valve
- * Indicates panel-mounted instruments.

Figure 19.21. Sketch of a fermenter with its auxiliary equipment. In most cases supplemental agitation by mechanical stirrers is common (A.J. C. Olsen, Chem. Ind., 416 (1960)].

660 OTHER TOPICS

TABLE 19.15. Standard Specifications of a Fermenter

- 1. Surfaces in contact with culture are 316 SS, all others 304 SS; free of crevices, mechanically ground and polished to approx 220 grit 2. Approx proportions: height/diameter = 2, impeller/vessel
- diameter = 0.35, baffle width/vessel diameter = 0.1
 - 3. Maximum working volume = 75-80%. minimum = 25%
 - 4. Ports and penetrations are 20 in number, namely
 - A. Steam-sterilizable inoculation/addition port
 - B. Combination viewing window/filling port on headplate
 - C. Light entrance window and lamp on headplate
 - D. Air inlet line
 - E. Air exhaust line
 - F. Well for temperature control sensor and temperature recorder sensor
 - G. Well for thermometer
 - H. Water inlet line to jacket of vessel
 - I. Water outlet line from jacket of vessel
 - J. Rupture disc on headplate and pressure relief valve on jacket
 - K. Diaphragm-type pressure gauge
 - L. Steam-sterilizable sample port
 - M. Steam-sterilizable bottom drain port, discharge valve is flush-bottom

(New Brunswick Scientific Co.).

REFERENCES

Membrane Processes

- 1. G. Belfort, Materials Science Of Synthetic Membranes: Fundamentals and Water Applications, Academic, New York, 1984.
- 2. T.D. Brock, Membrane Filtration: A User's Guide and Reference Manual, Science Tech, 1983.
- 3. S.T. Hwang, C.K. Choi, and K. Kammermeyer, Gaseous transfer coefficients in membranes, Separation Sci. 3, 461-478 (1974).
- 4. N.N. Li and W.S.W. Ho, Membrane processes, in Chemical Engineers' Handbook, McGraw-Hill, New York, 1984, pp. 17.14-17.34.
- 5. D.R. Lloyd (Ed.), Materials Science of Synthetic Membranes, ACS Symposium Series 269, American Chemical Society, Washington, DC, 1985
- 6. M.C. Porter, Membrane filtration, in Handbook of Separation Techniques for Chemical Engineers (P. A. Schweitzer, Ed.), McGraw-Hill, New York, 1979, pp. 2.3-2.103.
- 7. S. Sourirajan, Reverse Osmosis, Academic, New York, 1970.
- 8. A.F. Turbak (Ed.), Synthetic Membranes: Vol. I, Desalination; Vol. 2, Hyper and Ultrafiltration Uses, ACS Symposium Series 153 and 154, American Chemical Society, Washington, DC, 1981.

Foam Separation and Flotation

- 1. R. Lemlich, Adsorptive Bubble Separation Techniques, Academic, New York, 1972.
- 2. Y. Okamoto and E.J. Chou, Foam separation processes, in Handbook of Separation Techniques for Chemical Engineers (P. A. Schweitzer, Ed.), McGraw-Hill, New York, 1979, pp. 2.183-2.197.
- 3. P. Somasundaran, Foam separation methods, a review, In Separation and Purification Methods (Perry and van Oss, Eds.), 1972, Vol. 1, pp. 117-199
- 4. T.C. Sorensen, Flotation, in Chemical Engineers' Handbook, McGraw-Hill, New York, 1984, pp. 21.46-21.52.
- 5. B.A. Wills, Mineral Processing Technology, Pergamon, New York, 1985.

Sublimation and Freeze Drying

1. W. Corder, Sublimation, in Chemical Engineers' Handbook, McGraw-Hill, New York, 1984, pp. 17.12-17.14.

- N. Side-entry port for pH electrode
 - 0. Top-entering or side-entering (size-dependent) port for installation of the dissolved oxygen electrode

 - P. Top-entering port for foam sensor Q. Side-entering ports for acid, base, and antifoam addition (valved and piped as required)
- R. Spare penetrations on headplate for insertion of additional sensors i.e., 1 1/8 in. NPT, 1 3/8 in. NPT, 1 3/4 in. NPT

5. Foam breaking: Injection port provided for chemical breaking; mechanical breaker optional, consists of a double disk rotated at high speed with its own drive

6. Agitation system has three six-bladed turbine impellers adjustable along the shaft, maximum tip speed of 1200 ft/min, standard drive of 40 HP for a 5000-L vessel, bottom drive standard, top drive optional

7. Controls and monitors: liquid level, pH, dissolved oxygen, reduction-oxidation (Redox) potential, air rate, temperature, optional automatic sterilization cycle control, rupture disk on vessel, relief valve on iacket

- 2. N. Ganiaris, Freeze drying, in Chemical and Process Technology Encyclopedia (Considine, Ed.), McGraw-Hill, New York, 1974, pp. 523-527.
- 3. C.A. Holden and H.S. Bryant, Sublimation, Separation Sci. 4(1), 1 (1969)
- 4. C.J. Major, Freeze Drying, Chemical Engineers' Handbook, McGraw-Hill, New York, 1973 edition, pp. 17.26-17.28.
- 5. G. Matz, Sublimation, in Ullmann's Encyclopedia of Chemical Technology, Verlag Chemie, Weinheim, 1972, Vol. 2, pp. 664-671.
- 6. J.W. Mullin, Sublimation, in Crystallization, Butterworths, London, 1972, pp. 284-290.

Parametric Pumping and Cycling Zone Separation

- 1. N.H. Sweed, in Recent Developments in Separation Science (N. N. Li, Ed.), Vol. I, pp. 59-74.
- 2. P.C. Wankat, Cycling separation processes, Separation Sci. 9(2), 85-116 (1974).
- 3. P.C. Wankat, J.C.D. Ore, and W.C. Nelson, Cycling zone separations, in Separation and Purification Methods, CRC Press, Boca Raton, FL, 1976, Vol. 4, pp. 215-266.
- 4. R.H. Wilhelm et al., Ind. Eng. Chem. Fundam. 5, 141-144 (1966); 7, 337-349 (1968).

Thermal Dijusion

- 1. M. Benedict, T.H. Pigford, and H.W. Levi, Nuclear Chemical Engineering, McGraw-Hill, New York, 1981.
- 2. A.L. Jones and G.B. Brown, Liquid thermal diffusion, in Advances in Petroleum Chemistry and Refining (McKetta and Kobe, Eds.), Wiley, New York, 1960, Vol. III, pp. 43-76.
- 3. G. Vasaru et al., The Thermal Diffusion Column, VEB Deutscher Verlag der Wissenschaften, Berlin, 1969.

Electrochemical Syntheses

1. F. Hine, Electrode Processes and Electrochemical Engineering, Plenum, New York, 1985.

REFERENCES 661

- 2. D. Pletcher, Industrial Electrochemistry, Chapman and Hall, London, 1982.
- 3. R. Roberts, R.P. Ouellete, and P.N. Cheremisinoff, *Industrial Applications* **Of** *Electroorganic Synthesis*, Ann Arbor Science, Ann Arbor, MI, 1982.

Fermentation Processing

1. B. Atkinson, Biochemical Reactors, Pion Ltd., London, 1974.

- B. Atkinson and F. Mativuna, Biochemical Engineering and Biotechnology Handbook, Macmillan, Surrey, England, 1983.
 J.F. Bailey and D.F. Ollis, Biochemical Engineering Fundamentals,
- J.E. Bailey and D.F. Ullis, Biochemical Engineering Fundamenta McGraw-Hill, New York, 1986.
- 4. T. Godfrey and J. Reichelt, *Industrial Enzymology*, Macmillan, Surrey, England, 1983.
- 5. P.F. Stanbury and A. Whitaker, *Principles of Fermentation Technology*, Pergamon, New York, 1984.

COSTS OF INDIVIDUAL EQUIPMENT

The choice of appropriate equipment often is influenced by considerations of price. A lower efficiency or a shorter life may be compensated for by a lower price. Funds may be low at the time of purchase and expected to be more abundant later, or the economic life of the process is expected to be limited. Alternate kinds of equipment for the same service may need to be considered: water-cooled exchangers vs. air coolers, concrete cooling towers vs. redwood, **filters** vs. centrifuges, pneumatic conveyors vs. screw or bucket elevators, and so on.

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In this chapter, the prices of classes of the most frequently used equipment are collected in the form of correlating equations. The prices are given in terms of appropriate key characteristics of the equipment, such as sqft, gpm, lb/hr, etc. Factors for materials of construction and performance characteristics other than the basic ones also are provided. Although graphs are easily read and can bring out clearly desirable comparisons between related types of equipment, algebraic representation has been adopted here. Equations are capable of consistent reading, particularly in comparison with interpolation on logarithmic scales, and are amenable to incorporation in computer programs.

Unless otherwise indicated, the unit price is \$1000, \$K. Except where indicated, notably for fired heaters, refrigeration systems, and cooling towers (which are installed prices), the prices are purchase prices, FOB, with delivery charges extra. In the United States delivery charges are of the order of 5% of the purchase price, but, of course, dependent on the unit value, as cost per lb or per

EXAMPLE 20.1 Installed Cost of a Distillation Tower Shell and trays are made of AISI 304 stainless steel. Dimensional data are:

$$\begin{split} D &= 4 \text{ ft,} \\ L &= 120 \text{ ft,} \\ N &= 58 \text{ sieve trays,} \\ \text{wall thickness } t_p &= 0.50 \text{ in. for pressure,} \\ t_b &= 0.75 \text{ in. at the bottom,} \\ \text{flanged and dished heads weigh } 325 \text{ lb each,} \\ \text{weight } W &= (n/4)(16)(120(0.5/12)(501) + 2(325) = 32,129 \text{ lb} \\ C_b &= \exp[7.123 + 0.1478(10.38) + 0.02488(10.38)^2 \\ &+ 0.0158(120/4)\ln(0.75/0.50)] \\ &= 101,726 \\ f_1 &= 1.7, \\ f_2 &= 1.189 + 0.0577(4) = 1.420, \end{split}$$

 $f_{3} = 0.85,$ $f_{4} = 1,$ $C_{p1} = 204.9(4)^{0.6332}(120)^{0.8016} = 22,879,$ purchase price C = 1.7(101,726) + 58(1.42)(0.85)(753.4) + 22,879 = \$248,646

From Table 20.1, the installation factor is 2.1 so that the installed price is

 $C_{\text{installed}} = 2.1(248,646) = $522,156$

A tower packed with 2 in. pall rings instead of trays:

packing volume $V_p = (\pi/4)(4)^2(120) = 1508$ cuft, $C_{\text{installed}} = 2.1[1.7(101,726) + 1508(23.0) + 22,879)]$ = \$484,044

EXAMPLE 20.2

Purchased and Installed Prices of Some Equipment a. A box type fired heater with CrMo tubes for pyrolysis at 15OOpsig with a duty of 40 million Btu/hr. From Item No. 10 (Table 20.1), the installed price is

> $C_{\text{installed}} = 33.8(1.0 + 0.10 + 0.15)(40)^{0.86}$ = 1008.32 K\$, \$1,008,320.

b. A 225 HP-reciprocating compressor with motor drive and belt drive coupling. Items Nos. 2 and 13 (Table 20.1). The installation factor is 1.3.

compressor C = $5960(225)^{0.61}$ = 162,210, motor, 1800 rpm, TEFC, C = 1.2 × exp[4.5347 + 0.57065(5.42) + 0.04069(5.42)²] = \$8113, belt drive coupling, C = 1.2 $\exp[3.689 + 0.8917(5.42)]$ = \$6008, total installed cost, $C_{total} = 1.3(162,210 + 8113 + 6008)$ = \$229,230.

c. A two-stage steam ejector with one surface condenser to handle **200 lb/hr** of air at **25 Torr**, in carbon steel construction. From Table 20.3 the installation factor is 1.7.

X=200/25=8, $f_1 = 1.6, f_2 = 1.8, f_3 = 1.0$ purchase C = 11(1.6)(1.8(1.0)(8))^{0.41} = 74.31 K\$, \$74,310, installed C = 1.7C_p = \$126,330. **cuft**. Multipliers have been developed whereby the installed cost of **various** kinds of equipment may be found. Such multipliers range from 1.2 to 3.0, but details are shown in Table 20.3.

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Data are taken from a number of published sources and are updated to the beginning of 1985 with the cost indexes of *Chemical Engineering Magazine*, a selection of which is in Table 20.2. The main sources and the dates of their prices are Hall et al. (1981), Institut Francais du Petrole (1975), and Evans et al. (1979). References also are made to price data of some equipment not covered here. Many data as of mid-1982 have been collected by Ulrich (1984). Perry's *Chemical Engineers Handbook* (1984) has many data scattered throughout; the page numbers having such data are listed in the reference (Green, 1984).

Material of construction is a major factor in the price of equipment so that multipliers for prices relative to carbon steel or other standard materials are given for many of the items covered here. Usually only the parts in contact with process substances need be of special construction, so that, in general, the multipliers are not always as great as they are for vessels that are made entirely of special materials. Thus, when the tube side of an exchanger is special and the shell is carbon steel, the multiplier will vary with the amount of tube surface, as shown in that section.

As with most collections of data, the price data correlated here exhibit a certain amount of scatter. This is due in part to the incomplete characterizations in terms of which the correlations are made, but also to variations among manufacturers, qualities of construction, design differences, market situations, and other factors. Accordingly, the accuracy of the correlations cannot be claimed to be better than $\pm 25\%$ or so.

TABLE 20.1. Index of Equipment

•	
1. Agitators	Falling film
2. Compressors, turbines, fans	10. Fired heaters
Centrifugal compressors	Box types
Reciprocating compressors	Cylindrical types
Screw compressors	11. Heat exchangers
Turbines	Shell-and-tube
Pressure discharge	Double pipe
Vacuum discharge	Air coolers
Fans	12. Mechanical separators
3. Conveyors	Centrifuges
Troughed belt	Cyclone separators
Flat belt	Heavy duty
Screw, steel	Standard duty
Screw, stainless	Multiclone
Bucket elevator	Disk separators
Pneumatic	Filters
4. Cooling towers	Rotary vacuum belt discharge
Concrete	Rotary vacuum scraper discharge
Wooden	Rotary vacuum disk
5. Crushers and grinders	Horizontal vacuum belt
Cone crusher	Pressure leaf
Gvratory crusher	Plate-and-frame
Jaw crusher	Vibrating screens
Hammer mill	13. Motors and couplings
Ball mill	Motors
Pulverizer	Belt drive coupling
6. Crystallizers	Chain drive coupling
External forced circulation	Variable speed drive coupling
Internal draft tube	14. Pumps
Batch vacuum	Centrifugal
7 Distillation and absorption towers	Vertical mixed flow
Distillation trav towers	Vertical axial flow
Absorption tray towers	Gear numns
Packed towars	Reciprocating numps
	15 Petrigeration
0. Diyels Potary combustion are bested	16 Steam electors and vacuum number
Rotary, compusiton gas neated	To. Steam ejectors and vacuum pumps
Rolary, not all neated	
Rotary, steam tube neated	vacuum pumps 17 Vessels
Cabinet aryers	I/. VESSEIS
Spray dryers	Horizontai pressure vesseis
Multiple hearth furnace	Vertical pressure vessels
9. Evaporators	Storage tanks, shop tabricated
Forced circulation	Storage tanks, field erected
Long tube	

TABLE 20.2. Purchase Prices of Process Equipment (Basic: CE Plant Cost Index = 325, Middle 1985)

1. Agitators [Meyers and Kime, Chem. Eng., 109-112 (27 Sep. 1976)] $C = \exp[a + b \ln HP + c(\ln HP)^{2}]$ \$, 1 < HP < 400 Single Impeller Dual Impeller Speed 1 2 2 3 1 3 Carbon 8.57 8.43 8.31 8.80 8.50 6.43 а 0.1195 -0.0880 -0.1368 -0.1981 steel b 0.1603 0.0257 0.0819 c 0.1123 0.1015 0.0659 0.0878 0.1239 Type 316 6.62 8.55 6.52 9.25 8.82 6.72 -0.1225 0.2474 0.0308 -0.1802 0.2801 0.1235 b с 0.0654 0.0643 0.1158 0.0542 0.0818 0.1075 Speeds 1: 30, 37, and 45 rpm 2: 56, 68, 84, and 100 rpm 3: 125, 155,190, and 230 rpm 2. Compressors, turbines, and fans (K\$) Centrifugal compressors, without drivers (IFP, 1981): $C = 6.49(HP)^{0.62} K$ \$, 200 < HP < 30,000Reciprocating compressors without drivers (IFP): $C = 5.96(\text{HP})^{0.61} \text{ K}$ 100 < HP < 20.000 Screw compressors with drivers (IFP): $C = 1.49(HP)^{0.71}$ K\$. 10 < HP < 800 Turbines (IFP); $C = 0.31(HP)^{0.81}$ K\$. Pressure discharge, 20 < HP < 5000 $C = 0.69(HP)^{0.81}$ K\$. discharge. 200 < HP < 8000vacuum Fans with motors (Ulrich) $C = f_m f_n \exp[a + b \ln Q + c(\ln Q)^2]$ installed cost, K\$, Q in KSCFM b a а С 0.4692 0.1203 0.0931 Radial blades 2-500 Backward 0.0400 0.0786 2-900 curved 0.1821 Propeller -0.4456 0.2211 0.0820 2-300 Propeller, with guide vanes -1.0181 0.3332 0.0647 2-500 materials factor, f_m Carbon steel 2.2 Fiberglass 4.0 Stainless steel 5.5 Nickel alloy 11.0 Pressure Factors, Fp

	Centrifugal		Axial	
Pressure (kPa[gage])	Radial	Backward curved	Prop.	Vane
1	1.0	1.0	1.0	1.00
2	1.15	1.15	—	1.15
4	1.30	1.30		1.30
6	1.45	1.45	-	—
16	1.60	-		-

3. Conveyors (IFP) KS Troughed belt: C= $1.40L^{0.66}$, 10 < L < 1300ftFlat belt: $C = 0.90L^{0.66}$, 10 < L < 1300 ft Screw (steel): $C = 0.40L^{0.78}$, 7 < L < 100 ft Screw (stainless steel): $C = 0.70L^{0.78}$, 7 < L < 100 ft Bucket elevator: $C = 4.22L^{0.63}$, 10 < L < 100 ft Pneumatic conveyor (Chemical Engineers' Handbook, McGraw-Hill, New York, 1984). 600 ft length $C = \exp[3.5612 - 0.0048 \ln W + 0.0913(\ln W)^{2}], \quad 10 < W < 100 \text{ klb/hr}$ 4. Cooling towers, installed KS Concrete (IFP) C = $135fQ^{0.61}$, 1 < Q < 60 K gal/min: At (°C) 15 10 12 1.5 2.0

f 1.0 1.5 2.0 Redwood, without basin (Hall): C = $33.9Q^{0.85}$, 1.5 < Q < 20 K gal/min

5. Crushers and grinders (IFP) K\$

Cone crusher: $C = 1.55 W^{1.05}$, 20 < W < 300 tons/hrGyratory crusher: $C = 8.0W^{0.60}$, 25 < W < 200 tons/hrJaw crusher: $C = 6.3W^{0.57}$, 10 < W < 200 tons/hrHammer mill: $C = 2.44W^{0.78}$, 2 < W < 200 tons/hrBall mill: $C = 50.0W^{0.69}$, 1 < W < 30 tons/hrPulverizer: $C = 22.6W^{0.39}$, 1 < W < 5 tons/hr

6. Crystallizers (IFP, Chemical Engineers' Handbook, p. 19.40) External forced circulation:

> $C = f \exp[4.868 + 0.3092 \ln W + 0.0548(ln W)^2],$ 10 < W < 100 klb/hr of crystals

Internal draft tube: C= $178 fW^{0.58}$, 15 < W < 100 klb/hr of crystals Batch vacuum: C = $8.16 fV^{0.47}$, 50 < V < 1000 cuft of vessel

Туре		Material	f
Forced	circulation	Mild steel Stainless type 304	1.0 2.5
Vacuum	batch	Mild steel Rubber-lined Stainless type 304	1.0 1.3 2.0

7. Distillation and absorption towers, tray and packed (Evans et **al.,** 1984) prices in \$

Tray towers:

$$C_{t} = f_{1}C_{b} + Nf_{2}f_{3}f_{4}C_{t} + C_{p1}$$

Distillation:

$$C_b = \exp[7.123 + 0.1478(\ln W) + 0.02488(\ln W)^2 + 0.01580(L/D) \ln(T_b/T_p)],$$

9020 < W < 2,470,000 lbs of shell exclusive of nozzles and skirt
$$C_t = 375.8 \exp(0.1739D), \ 2 < D < 16 \text{ ft tray diameter}$$

N = number of trays

$C_{\rho 1} = 204.9D^{-1}L^{-1}L^{-1}$	2 < D < 24, 57 < L < 170	ft (platforms and	d ladders)

Material	f ₁	f ₂	
Stainless steel, 304 Stainless steel, 316	1.7 2.1	1.189 + 0.05770 1.401 + 0.07240	
Carpenter 20CB-3 Nickel-200	3.2 5.4	1.525 + 0.07880	
Monel-400 Inconel-600 Incolov-825	3.6 3.9 3.7	2.306 + 0.1120 <i>D</i>	
Titanium	7.7		(continued)
TABLE 20.2—(continued)

Tray Types	f3
Valve	1.00
Grid	0.80
Bubble cap	1.59
Sieve (with downcomer)	0.85

 $f_{A} = 2.25/(1.0414)^{N}$, when the number of trays N is less than 20

 T_{D} is the thickness of the shell at the bottom, T_{D} is thickness required for the operating pressure, D is the diameter of the shell and tray, L is tangent-to-tangent length of the shell

Absorption:

 $C_b = \exp(6.629 + 0.1826(\ln W) + 0.02297(\ln W)^2],$

4250 < W < 980,000 lb shell $C_{0,1} = 246.4D^{0.7396}L^{0.7068}, \quad 3 < D < 21,$

27 < L < 40 ft (platforms and ladders),

 f_1, f_2, f_3 , and f_4 as for distillation

Packed towers:

 $C = f_1 C_b + V_p C_p + C_{p1}$

 V_{p} is volume of packing, C_{p} is cost of packing \$/cuft

Packing Type	C_p (\$/cuft)
Ceramic Raschig rings,, 1 in.	19.6
Metal Raschig rings, 1 in.	32.3
Intalox saddles, 1 in.	19.6
Ceramic Raschig rings, 2 in.	13.6
Metal Raschig rings, 2 in.	23.0
Metal Pall rings, 1 in.	32.3
Intalox saddles, 2 in.	13.6
Metal Pall rings, 2 in.	23.0

8. Dryers (IFP)

Rotary combustion gas heated: $C = (1 + f_g + f_m) \exp[4.9504 - 0.5827(\ln A) + 0.0925(\ln A)^2]$, 200 < A < 30,000 sqft lateral surface

Rotary hot air heated: $C = 2.38(1 + f_g + f_m)A^{0.63}$, 200 <A < 4000 sqft lateral surface

Rotary steam tube: C= $1.83FA_t^{0.60}$, 500 $< A_t < 18,000$ sqft tube surface, F = 1 for carbon steel, F = 1.75 for 304 stainless

Cabinet dryer: C= $1.15 f_o A^{0.77}$, 10 < A < 50 sqft tray surface

Pressure	f _p
Atmospheric pressure	1.0
Vacuum	2.0
Material	f _m
Mild staal	1.0
willu Steal	

Dr	ying Gas	fg
Hot air		0.00
Combustion gas (direct contact)	0.12
Combustion gas	(indirect contact)	0.35
N	laterials	f _m
Mild steel		0.00
Mild steel Lined with stainles	ss 304–20%	0.00 0.25

Spray dryers:

 $C = F \exp(0.8403 + 0.8526(\ln x) - 0.0229(\ln x)^2)$

30 < x < 3000 lb/hr evaporation

Material	F
Carbon steel	0.33
304, 321	1.00
316'	1.13
Monel	3.0
Inconel	3.67

Multiple hearth furnaces (Hall et al., 1984)

 $C = \exp(a + 0.88N)$, 4 < N < 14 number of hearths

Diameter (ft)	6.0	10.0	14.25	16.75	18.75	22.25	26.75
Sqft/hearth,	12	36	89	119	172	244	342
approx a	5.071	5.295	5 521	5 719	5.853	6.014	6 0 9 4

9. Evaporators (IFP; also Chemical Engineers Handbook, p. 11.42)

Forced circulation: $C = f_m \exp[5.9785 - 0.6056(\ln A) + 0.08514(\ln A)^2]$, 150 < A < 8000 sqft heat transfer surface Long tube: $C = 0.36 f_m A^{0.86}$, 300 < A < 20,000 sqft

Falling film (316 internals, carbon steel shell)

C = exp[3.2362 – 0.0126(ln A) + 0.0244(ln A)²], 150 < A < 4000 sqft

Forced-Circulation Evaporators

Construction Material: Shell/Tube	f _m
Steel/copper	1.00
Monel/cupronickel	1.35
Nickel/nickel	1.80

Long-Tube	Evanorators
Long-Tube	

Construction Material: Shell/Tube	f _m
Steel/copper	1.0
Steel/steel	0.6
Steal/aluminum	0.7
Nickel/nickel	3.3

10. Fired heaters, installed (Hall) KS

Box type: $C = k(1 + f_d + f_p)Q^{0.86}$, 20 < Q < 200 M Btu/hr

k
25.5 33.8 45.0
f _d
0 0.10
0.35
f _p
0 0.10 0.15 0.25 0.40 0.60

Cylindrical type: $C = k(1 + f_d + f_p)Q^{0.82}$, 2 < Q < 30 M Btu/hr

Tube Material	k	
Carbon steel CrMo steel	27.3 40.2	
Stainless	42.0	

(continued)

Design Type		fa
Cylindrical Dowtherm		00.33
Design Pressure	(psi)	fp
up to 500 1,000 1,500		00.15 0.20

11. Heat exchangers

Shell-and-tube (Evans): C= $f_d f_m f_p C_{b'}$ price in \$

$C_{L} = C_{A} P[0, 02] = 0.30003(11) A + 0.0001 (11) A(1), 130 < A < 12,000 S$	31 (ln A) ²], 150 <a <="" th="" ′<=""><th>31 (In</th><th>0.0681</th><th>A) +</th><th>0.30863(In</th><th>exp[8.821 =</th><th>Сь =</th>	31 (In	0.0681	A) +	0.30863(In	exp[8.821 =	Сь =
---	---	--------	--------	------	------------	-------------	------

Туре		fa	
Fixed-head Kettle reboiler	exp[-1.1 1.35	156 + 0.09	06(In A)]
U-tube	exp[-0.9	816 + 0.083	30(in <i>A)]</i>
Pressure Range (p	sig)	f,	
100-300 300-600 600-900	0 1 1	0.7771 + 0.04 1.0305 + 0.0 1.1400 + 0.1	1981 (In A) 7140(In A) 2088(In A)
	_	$f_m = g_1 +$	g ₂ (in A)
Material		g 1	9 2
Stainless steel 3 Stainless steel 3 Stainless steel 3 Nickel 200 Monel 400 Inconel 600 Inconel 600 Inconel 825 Titanium Hastelloy	16 04 47	0.8603 0.8193 0.6116 1.5092 1.2989 1.2040 1.1854 1.5420 0.1549	0.23296 0.15984 0.22186 0.60859 0.43377 0.50764 0.49706 0.42913 0.51774

Double pipe (IFP): C = $900f_m f_n A^{0.18}$, 2 < A < 60 sqft, price in \$

Material: Shell/Tube	f _m
cs/cs cs/304L stainless	1.0 1.9
cs/316 stainless	2.2
Pressure (bar)	f _p
≤4	1.00
4–6	1.10
6-7	1.25

Air coolers (Hall): C = $24.6A^{0.40}$, 0.05 <A < 200 K sqft, price in K\$

12. Mechanical separators Centrifuges: solid bowl, screen bowl or pusher types

C=a+bW, K\$

	Inorganic	Process	Organic	Process
Material	а	b	а	b
Carbon steel	4 2	1.63		_
316	65	3.50	98	5.06
Monel	70	5.50	114	7.14
Nickel	84.4	6.56	143	9.43
Hastelloy	-	—	300	10.0

10 < W < 90 5 < W < 40 tons/hr

Disk separators, 316 stainless (IFP):

$$C = 8.0Q^{0.52}$$
, $15 < Q < 150$ gpm, K \$

Cyclone separators (IFP): K\$

 Filters (Hall), prices in \$/sqft: rotary vacuum belt discharge: $C = exp[11.20 \ 1.2252(ln A) + 0.0587(ln A)^2]$, 10 < A < 800 sqft rotary vacuum drum scraper discharge: $C = exp[11.27 - 1.3408(ln A) + 0.0709(ln A)^2]$ \$/sqft, 10 < A < 1500 sqft rotary vacuum disk: $C = exp[10.50 - 1.008(ln A) + 0.0344(ln A)^2]$ \$/sqft, 100 < A < 4000 sqft horizontal vacuum belt: $C = 28300/A^{0.5}$ \$/sqft, 10 < A < 1200 sqft pressure leaf: $C = 695/A^{0.29}$ \$/sqft, 30 < A < 2500 sqft plate-and-frame: (Chemical Engineers' Handbook): $C = 460/A^{0.45}$ \$/sqft, 10 < A < 1000 sqft vibrating screen (IFP): $C = 3.1A^{0.59}$ K\$, 0.5 < A < 35 sqft 13. Motors and couplings, prices in \$

Motors and couplings, prices in **S** Motors: $C = 1.2 \exp[a_1 + a_2(\ln HP) + a_3(\ln HP)^2]$ Belt drive coupling: $C = 1.2 \exp[3.689 + 0.8917(\ln HP)]$ Chain drive coupling: $C = 1.2 \exp[5.329 + 0.5048(\ln HP)]$ Variable speed drive coupling:

C = 12,000/(1.562 + 7.877/HP), HP < 75

Coefficients

Туре	a ₁	a ₂	a ₃	HP limit
Open, drip-proof				
3600 rpm	4.8314	0.09666	0.10960	1-7.5
	4.1514	0.53470	0.05252	7.5-250
	4.2432	1.03251	-0.03595	250-700
1800 rpm	4.7075	-0.01511	0.22888	1-7.5
	4.5212	0.47242	0.04820	7.5-250
	7.4044	-0.06464	0.05448	250-600
1200 rpm	4.9298	0.30118	0.12630	1-7.5
	5.0999	0.35861	0.06052	7.5-250
	4.6163	0.88531	-0.02188	250-500
Totally enclosed,	fan-cooled			
3600 rpm	5.1058	0.03316	0.15374	1-7.5
	3.8544	0.83311	0.02399	7.5-250
	5.3182	1.08470	-0.05695	250-400
1800 rpm	4.9687	-0.00930	0.22616	7.5-250
	4.5347	0.57065	0.04609	250-400
1200 rpm	5.1532	0.28931	0.14357	1-7.5
	5.3858	0.31004	0.07406	7.5-350
Explosion-proof				
3600 rpm	5.3934	-0.00333	0.15475	1-7.5
	4.4442	0.60820	0.05202	7.5-200
1800 rpm	5.2851	0.00048	0.19949	1-7.5
	4.8178	0.51086	0.05293	7.5-250
1200 rpm	5.4166	0.31216	0.10573	1-7.5
	5.5655	0.31284	0.07212	7.5-200

14. Pumps

Centrifugal (Evans) prices in \$: $C = F_M F_T C_b$, base cast-iron, 3550 rpm vsc

$C_{\rm p} = 1.55 \exp[8.833$	- 0.6019(In Q√H)			
2	+ 0.0519(ln $Q\sqrt{H})^2$],	Q in apm.	H in	ft head

Material	Cost	Factor	F _M
Cast steel		1.35	
304 or 316 fittings		1.15	
Stainless steel, 304 or 316		2.00	
Cast Gould's alloy no. 20		3.502.00	
Nickel			
Monel			
ISO B		3,304,95	
ISO c		4.60	
Titanium		9.70	
Hastellov C		2.95	
Ductile iron		115 190	
Bronze			
	11-01	$\sqrt{10^2}$	(
$F_{T} = \exp[D_{1} + D_{2}(\ln U \vee H) + D_{3})$	un Qu	v m)]	(continueu)

TABLE 20.2—(continued)

		Ť	уре		<i>b</i> 1	4	4
One-s One-s One-s Two-s Multis	tage, tage, tage, tage, tage, stage,	1750 3550 1750 3550 3550	rpm, rpm, rpm, rpm, rpm,	VSC HSC HSC HSC HSC	5.1029 0.0632 2.0290 13.7321 9.6849	-1.2217 0.2744 -0.2371 -2.6304 -1.6164	0.0771 -0.0253 0.0102 0.1542 0.0834
	T	ype			Flow Range (gpm)	Head Rang (ft)	ge HP (max)
One-stage, One-stage, One-stage, One-stage, Two-stage	3550 1750 3550 1750 3550	rpm, rpm, rpm, rpm, rpm,	VSC VSC HSC HSC HSC		50-900 50-3500 100-1500 250-5000 50-1100	50-400 50-200 100-450 50-500 300-1100	75 200 150 250
Two-stage,	3550	rpm,	HSC		100-1500	650-3200	1450

Vertical mixed flow (IFP): $C = 0.036(gpm)^{0.82} K$ \$, 500 < gpm c 130,000 Vertical axial flow (IFP): $C = 0.020(gpm)^{0.78} K$ \$, 1000 < gpm < 130,000 Gear pumps (IFP): $C = exp[-0.0881+ 0.1986(ln Q) + 0.0291(ln Q)^2] K$ \$, 10 < Q < 900 gpm

Reciprocating (Pikulik and Diaz, 1979), without motor,

Cast iron: $C = 40.0Q^{0.81}$ K\$, 15 < Q < 400 gpm Others: $C = 410FQ^{0.52}$ K\$, 1 < Q < 400 gpm

F = 1.00
1.40
1.66
2.20

15. Refrigeration (IFP): C = 146FQ^{0.65} K\$, 0.5 < CI < 400 M Btu/hr, installed prices

Temperature Level (°C)	F
0	1.00
-10	1.55
-20	2.10
-30	2.65
-40	3.20
-50	4.00

16. Steam ejectors and vacuum pumps (Pikulik and Diar, 1979):

Ejectors: C= $11.0f_1f_2f_3X^{0.41}$ K\$, 0.1 < X < 100

X= (lb air/hr)/(suction pressure in Torr)

T	/pe	f ₁	No. Stages	f ₂	Material	ß
No condense 1 surface co 1 barometric 2 surface co 2 barometric	r ondenser condenser ondensers condensers	1.0 1.6 1.7 2.3 1.9	1 2 3 4 5	1.0 1.8 s 2.1 h 2.6 4.0	carbon steel stainless steel astelloy	1.0 2.0 3.0

Vacuum pumps: $C = 8.15X^{1.03}$ K\$, 0.3 < X < 15 (lbs air/hr)/(suction Torr).

17. Vessels (Evans) prices in **\$** Horizontal pressure vessels: $C = F_M C_b + C_a$

 $C_b = \exp[8.571 - 0.2330(\ln W) + 0.04333(\ln W)^2],$ 800 < W < 914,000 lb shell weight $C_s = 1 \ 370D^{0.2029}, \ 3 < 0 < 12$ ft diameter (platforms and ladders)

Vertical vessels: $C = F_M C_b + C_a$

 $C_b = \exp[9.100 - 0.2889(\ln W) + 0.04576(\ln W)^2],$

5000 < W < 226,000 lb

$$C_{\rm a} = 246D^{0.7396}L^{0.7068}, \quad 6 < D < 10,$$

12 < L < 20 ft tangent-to-tangent

FM

Material	Cost	Factor
Stainless steel, 304 Stainless steel, 316 Carpenter 20CB-3 Nickel-200 Monel-400 Inconel-600 Incoloy-825 Titanium		1.7 2.1 3.2 5.4 3.6 3.9 3.7 7.7

- Storage tanks, shop fabricated: $C = F_M \exp[2.631 + 1.3673(\ln V) 0.06309(\ln V)^2]$, 1300 < V < 21,000 gal
- Storage tanks, field erected: C = F_M exp[11.662 − 0.6104(ln V) + 0.04536(ln V)²], 21,000~ V < 11,000,000 gal

Material of Construction	Cost	Factor	FM
Stainless steel 316 Stainless steel 304 Stainless steel 347 Nickel Monel Inconel Zirconium Titanium		2.7 2.4 3.0 3.5 3.3 3.8 11.0 11.0	
brick-and-polyester-lined steel Rubber- or lead-lined steel Polyester, fiberglass-reinforced Aluminum Copper Concrete		2.75 1.9 0.32 2.7 2.3 0.55	

TABLE 20.3. Multipliers for Installed Costs of Process Equipment^a

Equipment	Multiplier	Equipment	Multiplier
Agitators, carbon steel	1.3	Chimneys and stacks	1.2
stainless steel	1.2	Columns, distillation, carbon steel	3.0
Air heaters, all types	1.5	distillation, stainless steel	2.1
Beaters	1.4	Compressors, motor driven	1.3
Blenders	1.3	steam on gas driven	1.5
Blowers	1.4	Conveyors and elevators	1.4
Boilers	1.5	Cooling tower, concrete	1.2
Centrifuges, carbon steel	1.3	Crushers, classifiers and mills	1.3
stainless steel	1.2	Crystallizers	1.9
			(continued)

TABLE 20.3—(continued)

Equipment	Multiplier	Equipment	Multiplier	
Cyclones	1.4	Pumps, centrifugal, carbon steel	2.8	
Dryers, spray and air	1.6	centrifugal, stainless steel	2.0	
other	1.4	centrifugal, Hastelloy trim	1.4	
Ejectors	1.7	centrifugal, nickel trim	1.7	
Evaporators, calandria	1.5	centrifugal, Monel trim	1.7	
thin film, carbon steel	2.5	centrifugal, titanium trim	1.4	
thin film, stainless steel	1.9	all others, stainless steel	1.4	
Extruders, compounding	1.5	all others, carbon steel	1.6	
Fans	1.4	Reactor kettles, carbon steel	1.9	
Filters, all types	1.4	kettles, glass lined	2.1	
Furnaces, direct fired	1.3	kettles, carbon steel	1.9	
Gas holders	1.3	Reactors, multitubular, stainless steel	1.6	
Granulators for plastic	1.5	multitubular, copper	1.a	
Heat exchangers, air cooled, carbon steel	2.5	multitubular, carbon steel	2.2	
coil in shell, stainless steel	1.7	Refrigeration plant	1.5	
glass	2.2	Steam drums	2.0	
graphite	2.0	Sum of equipment costs, stainless steel	1.8	
plate, stainless steel	1.5	Sum of equipment costs, carbon steel	2.0	
plate, carbon steel	1.7	Tanks, process, stainless steel	1.8	
shell and tube, stainless/stainless steel	1.9	Tanks, process, conner	1.9	
shell and tube, carbon/stainless steel	2.1	process. aluminum	2.0	
Heat exchangers, shell and tube, carbon steel/aluminum	2.2	storage, stainless steel	1.5	
shell and tube, carbon steel/copper	2.0	storage, aluminum	1.7	
shell and tube, carbon steel /Monel	1.8	storage, carbon steel	2.3	
shell and tube, Monel/Monel	1.6	field erected, stainless steel	1.2	
shell and tube, carbon steel/Hastelloy	1.4	field erected, carbon steel	1.4	
Instruments, all types	2.5	Turbines	1.5	
Miscellaneous, carbon steel	2.0	Vessels pressure stainless steel	1.7	
stainless steel	1.5	pressure, carbon steel	2.8	

^a [J. Gran, Chem. Eng., (6 Apr. 1981)].

Installed Cost = (purchase price)(multiplier).

Year	1970	1975	1980	Oct . 1985
CE Plant Cost Index	125.7	182.4	261.2	325.8
Equipment costs	123.8	194.7		347.5
Fabricated equipment	122.7	192.2		335.5
Process machinery	122.9	84.7		333.3
Piping, valves, and fittings	132.0	217.0		385.3
Process instruments and controls	132.1	181.4		323.9
Pumps and compressors	125.6	208.3		421.1
Electrical equipment	99.8	142.1		251.9

TABLE 20.4. Chemical Engineering Magazine Cost Indexes

REFERENCES

- 1. Chemical Engineering Magazine, Modern Cost Engineering, McGraw-Hill, New York, 1979.
- 2. Chemical Engineering Magazine, Modern Cost Engineering II, McGraw-Hill, New York, 1984.
- 3. L.B. Evans, A. Mulet, A.B. Corripio, and K.S. Chretien, Costs of pressure vessels, storage tanks, centrifugal pumps, motors, distillation and absorption towers, in Ref. 2, pp. 140-146, 177-183.
- J. Gran, Improved factor method gives better preliminary cost estimates, in Ref. 2, pp. 76–90.
- D.W. Green and J.O. Maloney (Eds.), *Perry's Chemical Engineers' Handbook*, McGraw-Hill, New York, 1984, cost data on pp. 6.7, 6.22, 6.112, 6.113, 6.121, 7.19, 11.19, 11.20, 11.21, 11.29, 11.42, 17.27, 17.33, 18.45, 18.46, 18.47, 19.13, 19.40, 19.45, 19.65, 19.89, 19.101, 19.102,

 $20.37, \ 20.38, \ 21.22, \ 21.45, \ 22.134, \ 22.135, \ 25.69, \ 25.73-25.75.$

- R.S. Hall, J. Matley, and K.J. McNaughton, Current costs of process equipment, in Ref. 2, pp. 102–137.
- Institut Francaise du Petrole (IFP), Manual of Economic Analysis of Chemical Processes, Technip 1976, McGraw-Hill, New York, 1981.
- 8. B.G. Liptak, Costs of process instruments, in Ref. 1, pp. 343-375.
- A. Pikulik and H.E. Diaz, Costs of process equipment and other items, in Ref. 1, pp. 302–317.
- G.P. Purohit, costs of shell-and-tube heat exchangers, *Chem. Eng., (22* Aug. 1983, 4 Mar. 1985, 18 Mar. 1985).
- 11. G.D. Ulrich, A Guide to Chemical Engineering Process Design and Economics, Wiley, New York, 1984.
- W.M. Vatavuk and R.B. Neveril, Costs of baghouses, electrostatic precipitators, venturi scrubbers, fanc carbon adsorbers, flares and incinerators, in Ref. 2, pp. 184-207.

UNITS, NOTATION, AND GENERAL DATA

- 1. Units and conversions 671
- 2. Notation 672
- 3. Properties of steam and water 673
- 4. Properties of air and steam at atmospheric pressure 674

TABLE AI. Units and Conversions

```
Prefixes for Unit Multiples and
Submultiples:
10<sup>-18</sup>
            atto a IO'
                                      deca d a
10^{-15} femto f
10^{-12} pico p
                           10<sup>2</sup>
                                      hecto h
pico p
10<sup>--9</sup> nano n
10<sup>--6</sup> mic
                           10<sup>3</sup>
10<sup>6</sup>
                                      kilo
                                                 k
                                      mega M
          micro μ 10<sup>9</sup>
                                      giga
                                                 G
 10<sup>-3</sup> milli
                    m 10<sup>12</sup>
                                                  Т
                                      tera
10<sup>-2</sup>
          centi c
 10
           deci
                    d
```

Length:

1 ft = 0.3048 m = 30.48 cm = 304.8 mm

Volume:

1 cuft = 0.0283 cum = 7.481 U.S. gal 1 cum = 35.34 cuft = 1000 L

Standard gas volume:

22.414 L/g mol at 0°C and 1 atm 359.05 cuft/lb mol at 32°F and 1 atm

Gas constant R:

Temperature	Mole	R
"Rankine	lb	4.969 x 10 ⁴
"Rankine	lb	1544
"Rankine	lb	0.7302
"Rankine	lb	10.73
"Rankine	lb	1.987
"Rankine	lb	7.805×10^{-4}
"Rankine	lb	5.819 x 10 ⁻⁴
Kelvin	g	8.314
Kelvin	kg	8.314 x 10 ³
Kelvin	kg	8.478 × 10 ²
Kelvin	g	82.0562
Kelvin	g	1.987
	Temperature "Rankine "Rankine "Rankine "Rankine "Rankine "Rankine Kelvin Kelvin Kelvin Kelvin Kelvin Kelvin	TemperatureMole"RankineIb"RankineIb"RankineIb"RankineIb"RankineIb"RankineIb"RankineIbKelvingKelvinkgKelvingKelvingKelvingKelvingKelvingKelvingKelvingKelving

Gravitational constant:

 $g_c = 1 \text{ kg mass/N sec}^2$

- $= 1 \text{ g cm/dyn sec}^2$
- = 9.806 kg mass/kg force \sec^2

= 32.174 lb mass/lb force sec^2

Mass:

1 lb = 0.4536 kg1 kg = 2.2046 lb

- 5. Properties of steel pipe 675
- 6. Standard gauges of sheets, plates, and wires 676
- 7. Weights and angles of slide of various materials

677

8. Petroleum products, typical compositions **680**

Density:

1 **lb/cuft** = 16.018 kg/cum 1 **gm/cucm** = 62.43 **lb/cuft** "API = 141.5/(specific gravity) - 131.5 specific gravity = 141.5/(°API + 131.5)

Force:

1 lb force = 0.4536 kg force = 4.448 Newtons

Pressure:

```
1 atm = 760 Torr = 760 mm Hg = 101,325 N/sqm
= 1.01325 bar = 10,330 kg/sqm = 14.696 lbf/sq in
= 2,116.2 lbf/sqft
1 bar = 100,000 N/sqm
1 Pa = 1 N/sqm
```

Energy, work, and heat:

Btu = 252.16 cal = 1055.06 J = 0.2930 W hrs = 10.41 L atm
HP hr = 0.7457 kWh = 778 ft lbf = 2545 cal
1 cal = 4.1868 J
J = 1 N m = 1 W sec = 0.2388 cal = 0.000948 Btu

Power:

1 ft lbf/sec = 0.0018182 HP = 1.356 W = 0.0012856 Btu/sec = 0.3238 cal/sec 1 W = 1 J/sec = 1 N m/sec

Temperature:

K (Kelvin) = °C (centigrade) + 273.16 = [°F (Fahrenheit)
+ 459.6]/1.8 = °R (Rankine)/1.8
°R =
$$1.8 \text{ K} = °F + 459.6$$

°C = (°F - 32)/1.8

Temperature difference:

 $1^{\circ}C = 1^{\circ}K = 1.8^{\circ}R = 1.8^{\circ}F$

Heat capacity and entropy:

$$1 \text{ cal/(g)(°K)} = 4.1868 \text{ J/(g)(°K)} = \text{Btu/(lb)(°R)}$$

Specific energy:

TABLE AI-(continued)

Volumetric flow:

1 cuft/sec = 0.028316 cum/sec = 28.316 L/sec

Heat flux:

 $1 \text{ Btu/(hr)(ft^2)} = 3.1546 \text{ W/m}^2$ $= 2.172 \text{ kcal/(hr)(m^2)}$

Heat transfer coefficient:

1 $Btu/(hr)(ft^2)(F) = 5.6783 W/m^2K$

Surface tension:

 $1 \text{ dyn/cm} = 1 \text{ erg/cm}^2$ $= 0.001 \text{ N sec/m}^2$

Viscosity, dynamic:

1 cP = 0.001 N sec/m² = 0.001 Pa sec = 0.000672 lb_m/ft sec = 2.42 lb_m/ft hr = 0.0752 lb_f hr/ft²

Viscosity, kinematic:

1 centistoke = $0.00360 \text{ m}^2/\text{hr}$ = $0.0388 \text{ ft}^2/\text{hr}$

TABLE A2. Notation*

 $C_{\rm p}$ = heat capacity at constant pressure C_{i} = heat capacity at constant volume g_c = gravitational constant (numerical values in Table AI) h = individual heat transfer coefficient H = enthalpyk = thermal conductivity $k = C_n / C_v$ K = y/x = vaporization equilibrium ratio, VER m. = mass fraction of component *i* of a mixture = molecular weight P = pressureQ = volumetric flow rate Q = heat transfer rate R = gas constant (numerical values in Table AI) S = entropyT = temperature, usually °R or °K U = linear velocity U = overall heat transfer coefficient V = volume x_i = mol fraction of component i in the liquid phase y_i = mol fraction of component i in the vapor phase z_i = mol fraction of component *i* in a mixture z = PV/RT, compressibility μ = viscositv ρ = density σ = surface tension

'Most symbols are defined near where they are used in equations. Unless defined otherwise locally, certain notations have the meanings in this list.

Temp., °F	Absolute pressure, lb/sq in.	Latent heat of vaporation, Btu/lb	Specific volume of steam, cu ft/lb	Density of quid water Ib/cu ft	7 iscosity of quid water, entipoises	'hermal con- uctivity o f Iguid water. (Btn)(ft)/ °F)(ft ²)(hr)
82 35 40 45 50	0.0885 0.1000 0.1217 0.1475 0.1781	1075.8 1074.1 1071.3 1068.4 1065.6	3306 2947 2444 2036.4 1703.2	62.42 62.42 62.42 62:42 62:42 62.39	$\begin{array}{c} 1.786 \\ 1.689 \\ 1.543 \\ 1.417 \\ 1.306 \end{array}$	$\begin{array}{c} 0.320 \\ 0.322 \\ 0.326 \\ 0.329 \\ 0.333 \end{array}$
55	0.2141	1062.7	1430.7	62.39	1.208	0. 336
60	0.2563	1059.9	1206.7	62.35	1.121	0. 340
65	0.3056	1057:1	1021.4	62.30	1.044	0. 343
70	0.3631	1054.3	867.9	62.28	0.975	0. 346
75	0: 4298	1051.5	740.0	62.23	0.913	0. 349
80	0.5069	1043.6	633.1	62.19	0.857	0.352
85	0.5959	1045. a	543.5	62.14	0.807	0.355
90	0.6982	1042.9	468.0	62.12	0.761	0.358
95	0.8153	1040.1	404.3	62.03	0.719	0.360
100	0.9492	1037.2	350.4	62.00	0.681	0.362
105	1.1016	1034.3	304.5	61.92	0.646	0. 364
110	1.275	1031.6	265.4	61.85	0.614	0. 367
115	1.471	1028.7	231.9	61.80	0.585	0. 369
120	1.692	1025.8	203.27	61.73	0.557	0. 371
125	1.942	1022.9	178.61	61.66	0.532	0.373
130	2.222	1020.0	157.34	$\begin{array}{c} 61.55\\ 61.46\\ 61.39\\ 61.28\\ 61.21 \end{array}$	0.509	0. 375
135	2.537	1017.0	138.95		0.487	0. 376
140	2.889	1014.1	123.01		0.467	0. 378
145	3.281	1011.2	109.15		0.448	0. 379
150	3.718	1008.2	97.07		0: 430	0. 381
155 160 165 170 175	$\begin{array}{c} 4.203 \\ 4.741 \\ 5.335 \\ 5.992 \\ 6.715 \end{array}$	1005.2 1002.3 999.3 996.3 993.3	$\begin{array}{c} 86.52 \\ 77.29 \\ 69.19 \\ 62.06 \\ 55.78 \end{array}$	61.10 61.01 60.90 60.79 60.68	0.414 0.398 0.384 0.370 0.357	0. 382 0. 384 0. 385 0. 386 0. 387
180	7.510	990.2	50, 23	60.58	0.345	0. 388
185	8.383	987.2	45, 31	60.47	0.334	0. 389
190	9.339	984.1	40, 96	60.36	0.333	0. 390
195	10.385	981.0	37, 09	60.25	0.312	0.391
200	11.526	977.9	33, 64	60.13	0.303	0.392
205 210 212 215 220	$\begin{array}{c} 12.777\\ 14.123\\ 14.696\\ 15.595\\ 17.188\end{array}$	974.8 971.6 970.3 968.4 965.2	30.57 27.82 26.80 25.37 23.15	60.02 59.88 59.75 59.70 59.64	0.293 8 % 0: 277 0.270	0. 392 0. 393 0. 393 0. 393 0. 393 0. 394
225	18.93	962.0	21.17	59.48	0.262	0.394
230	20.78	958.8	19.382	59.39	0.255	0.395
235	22.80	955: 5	17.779	59.24	0.248	0.395
240	24.97	952.2	16.323	59.10	0: 242	0.396
2 45	27.31	948.9	15.012	58.93	0.236	0.396
250	$\begin{array}{c} 29.82 \\ 35.43 \\ 41.86 \\ 49.20 \\ 57.55 \end{array}$	945.5	13.821	58.83	229	0.396
260		938.7	11.763	58.52	8: 218	0.396
270		931.8	10.061	58.24	0.208	0.396
280		924.7	8.645	57.94	0.199	0:396
290		917.5	7.461	57.64	0.191	0.396
300	67.01	910.1	6.466	57.31	0.183	0. 396
310	77.68	902.6	5.626	56.98		0. 396
320	89.66	894.9	4.914	56.55		0. 395
330	103.06	887.0	4.307	56:31		0. 393
340	118.01	879.0	3.788	55.96		0. 392
350	134.62	870.7	3.342	55.59		0.390
360	153.04	862.2	2.957	55.22		0.388
370	173.37	853.5	2.625	54.85		0.387
380	195.77	844: 6	2.335	54.46		0.385
390	220.37	835.4	2.0836	54.05		0.383
400	247.31	826.0	1.8633	63.65		0. 382

TABLE A3. Properties of Steam and Water

Source: Condensed from **Keenan** and Keyes, Thermodynamic Properties of Steam, Wiley, New York, 1936).

 TABLE
 A4.
 Properties of Air and Steam at Atmospheric Pressure

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Air

-					
0	0.086	0 239	1 1 1 0	0.130	0.0133
32	0.081	0.240	1.165	0.145	0.0140
100	0.071	0.240	1.285	0.180	0.0154
200	0.060	0.241	1.440	0.239	0.0174
300	0.052	0.243	1.610	0.306	0.0193
400	0.046	0.245	1.750	0.378	0.0212
500	0.0412	0.247	1.890	0.455	0.023 l
600	0.0373	0.250	2.000	0.540	0.0250
700	0.0341	0.253	2.14	0.625	0.0268
800	0.0314	0.256	2.25	0.717	0.0286
900 (0.029 1	0.259	2.36	0.815	0.0303
1000	0.0271	0.262	2.47	0.917	0.0319
1.500	0.0202	0.276	3.00	1.47	0.0400
2000	0.0161	0.286	3.45	2.14	0.0471
2500	0.0133	0.292	3.69	2.80	0.051
3000	0.0114	0.297	3.86	3.39	0.054
		!	L .		

Steam

0.0372	0.451	0.870	0.234	0.0145
0.0328	0.456	1.000	0.303	0.0171
0.0288	0.462	1.130	0.395	0.0200
0.0258	0.470	1.265	0.490	0.0228
0.0233	0.477	1.420	0.610	0.0257
0.0213	0.485	1.555	0.725	0.0288
0.0196	0.494	1.700	0.855	0.032 1
0.0181	0.50	1.810	0.987	0.0355
0.0169	0.51	1.920	1.13	0.0388
0.0149	0.53	2.14	1.44	0.0457
0.0133	0.55	2.36	1.78	0.053
0.0120	0.56	2.58	2.14	0.061
0.0109	0.58	2.81	2.58	0.068
0.0100	0.60	3.03	3.03	0.076
0.0083	0.64	3.58	4.30	0.096
0.0071	0.67	4.00	5.75	0.114
<u> </u>	L _	L _	L _	
	$\begin{array}{c} 0.0372\\ 0.0328\\ 0.0288\\ 0.0258\\ 0.0233\\ 0.0213\\ 0.0196\\ 0.0181\\ 0.0169\\ 0.0149\\ 0.0133\\ 0.0120\\ 0.0109\\ 0.0100\\ 0.0083\\ 0.0071\\ \end{array}$	$\begin{array}{c ccccc} 0.0372 & 0.451 \\ 0.0328 & 0.456 \\ 0.0288 & 0.462 \\ 0.0258 & 0.470 \\ 0.0253 & 0.477 \\ 0.0213 & 0.485 \\ 0.0196 & 0.494 \\ 0.0181 & 0.50 \\ 0.0169 & 0.51 \\ 0.0149 & 0.53 \\ 0.0133 & 0.55 \\ 0.0120 & 0.56 \\ 0.0109 & 0.58 \\ 0.0109 & 0.58 \\ 0.0003 & 0.64 \\ 0.0071 & 0.67 \\ \end{array}$	$\begin{array}{c cccccc} 0.0372 & 0.451 & 0.870 \\ 0.0328 & 0.456 & 1.000 \\ 0.0288 & 0.462 & 1.130 \\ 0.0258 & 0.470 & 1.265 \\ 0.0233 & 0.477 & 1.420 \\ 0.0213 & 0.485 & 1.555 \\ 0.0196 & 0.494 & 1.700 \\ 0.0169 & 0.51 & 1.920 \\ 0.0149 & 0.53 & 2.14 \\ 0.0133 & 0.55 & 2.36 \\ 0.0120 & 0.56 & 2.58 \\ 0.0109 & 0.58 & 2.81 \\ 0.0083 & 0.64 & 3.58 \\ 0.0071 & 0.67 & 4.00 \\ \end{array}$	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$

Nom- inal pipe		Sched-		·low area	Surfac lin ft,	ce per ft²	Weight
size, in.	OD, in.	ule No.	ID, in.	n. ² pipe,	Outside	Inside	per lin it, lb steel
1/8	0.405	40†	0.269	0.058	0.106	0.070	0.25
14	0.540	801 40	$0.215 \\ 0.364$	0.036 0.104	0.106 0.141	$0.056 \\ 0.095$	0.32 0.43
		80	0.302	0.072	0.141	0.079	0.54
3/8	0.675	4 0	0.493	0.192	0.177	0.129	0.57
		80	0.423	0.141	0.177	0.111	0.74
1/2	0.840	40	0.622	0.304	0.220	0.163	0.85
		80	0.546	0.235	0.220	0.143	1.09
3/4	1.05	4 0	0.824	0.534	0.275	0.216	1.13
		80	0.742	0.432	0.275	0.194	1.48
1	1.32	4 0	1.049	0.864	0.344	0.274	1.68
		8 0	0.957	0.718	0.344	0.250	2.17
11/4	1.66	4 0	1.380	1.50	0.435	0.362	2.28
		8 0	1.278	1.28	0.435	0.335	3.00
11/2	1.90	4 0	1.610	2.04	0.498	0.422	2.72
		8 0	1.500	1.76	0.498	0.393	3.64
2	2.38	4 0	2.067	3.35	0.622	0.542	3.66
		80	1.939	2.95	0.622	0.508	5.03
$2\frac{1}{2}$	2.88	4 0	2.469	4.79	0.753	0.647	5.80
		80	2.323	4.23	0.753	0.609	7.67
3	3.50	4 0	3.068	7.38	0.917	0.804	7.58
		80	2.900	6.61	0.917	0.760	10.3
4	4.50	4 0	4.026	12.7	1.178	1.055	10.8
		8 0	3.826	11.5	1.178	1.002	15.0
6	6.625	4 0	6.065	28.9	1.734	1.590	19.0
		8 0	5.761	26.1	1.734	1.510	28.6
8	8.625	4 0	7.981	50.0	2.258	2.090	28.6
		8 0	7.625	45.7	2.258	2.000	43.4
10	10.75	40	10.02	78.8	2.814	2.62	40.5
		6 0	9.75	74.6	2.814	2.55	54.8
12	12 75	3.0	12 09	115	3.338	3.17	43.8
16	16.0	30	15.25	183	4.189	4.00	62.6
20	20.0	2 0	19.25	291	5.236	5.05	78.6
24	24.0	20	23.25	425	6.283	6.09	94.7
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TABLE A5. Properties of Steel Pipe

† Schedule 40 designates former "standard" pipe.‡ Schedule 80 designates former "extra-strong" pipe.

Sheet mills roll steel sheets to U. S. gauge unless otherwise ordered. Plate mills usually roll heavy plates, ³/₄ and heavier, and light plate No. 8 to No. 12, to Birmingham gauge. In figuring weights of steel plates add to above the allowance for overweight, adopted by Association American Steel Manufacturers. All steel sheets in our stock are rolled to the U. S. Standard Gauge. Brass is rolled to thickness by Brown & Sharpe's American Gauge. Copper is rolled to thickness by Stubs' or Birmingham Gauge.



THICKNESS AND WEIGHT OF SHEETS AND PLATES								WIRE
No. of	U.S. S Ado G Ma	tandaro pted by overnm arch 1,	d Gauge 7 U. S. ent 1937	Birmingham or Stubs' Gauge		American or Brown & Sharpe's Gauge		Washburn & Moen Gauge
Gauge	Thic Inc	kness hes	Weigh‡ Lbs. per Sq. Ft.	Thick ness, Inche	Weight Lbs. per Sq. Ft.	hick- ness nches	Weigh Lbs. Der Sq. Ft.	l'hick- ness, nches
000 00 0 1 2 3	3/8 11/4 21/4 19/4 19/4 19/4		15.00 13.75 12.50 11.25 10.625 10.00	.425 .380 .340 .300 .284 .259	17.28 15.45 13.82 12.20 11.55 10.53	.410 .365 .325 .289 .258 .229	16.71 14.88 13.26 11.80 10.51 9.36	.363 .331 .307 .283 .263 .244
456789	15/27 E / E / E / E / E	.224 .209 .194 .179 .164 .149	9.375 8.15 8.125 7.50 6.875 6.25	.238 .220 .203 .180 .165 .148	9.68 8.95 8.25 7.32 6.71 6.02	.204 .182 .162 .144 .128 .114	8.34 7.42 6.61 5.89 5.24 4.67	.225 .207 .192 .177 .162 .148
10 11 12 13 14 15	1/8 1/10 \$/52 5/64	.134 .120 .105 .09 .075 .067	6.625 5.00 4.375 8.75 3.125 2.813	.134 .120 .109 .095 .083 .072	5.46 4.88 4.43 3.86 8.37 2.93	.102 .091 .08 .072 .064 .057	4.16 3.70 3.30 2.94 2.62 2.33	.135 .120 .105 .092 .080 .072
16 17 18 19 20 21	1/15 3/64 	.060 .054 .048 .042 .036 .033	2.50 2.25 2:00 1.75 1.60 1.375	.065 .058 .049 .042 .035 .032	2.64 2.36 1.99 1.71 1.42 1.30	.05 .045 .036 .032 .028	2.07 1.85 1.64 1.46 1.31 1.16	.063 .054 .047 .041 .035 .032
22 23 24 25 26 27	· · · · · · · · · · · · · · · · · · ·	.030 .027 .024 .021 .018 .016	1.25 1.125 1.00 .875 .750 .687	.028 .025 .022 .020 .018 .016	1.14 1.02 .895 .813 .732 .651	.025 .023 .020 .018 .016 .014	1.03 .922 .82 .73 .649 .579	.027 .026 .023 .020 .018 .017
28 29 30	 	.015 .014 .012	.625 .563 .500	.014 .013 .012	.569	.01 2 .011 .01	.514 .461 .408	.016 .015 .014

TABLE A7. Weights and Angles of Slide of Various Materials

*Weights of Materials—The following list gives weights in pounds per cubic foot. Unless otherwise noted, weights are for material in loose, least compacted form. In figuring Horse Powers, weights should be increased in proportion to their compressibility.

†Angles of Slide—The angles given are the **minimum** at which the various materials will slide on **a** steel plate, **under best** condition, for determination of friction. The minimum angle will **increase** as size of particles decrease and with **higher** moisture content. For definite recommendations refer to S-A Engineers. The inclination of **chuter must be steeper than** minimum angle of slide and S-A Engineers should be consulted for minimum chute slopes.

Friction Factors—The moving-friction factor for any material listed, sliding on steel plate, equals the natural tangent of the "angle of slide" given for that material. See table of natural functions of angles-listed in data section of book. For example, the friction factor of cement equals .809 (the natural tangent of 39°, which is the angle of slide for cement).

Specific Gravity-The specific gravity of a material is its weight (in a solid block) compared with that of water at 62" F. Example: As water weighs 62.4 pounds per cubic foot and sulphur weighs 125 pounds, the specific gravity of sulphur is twice that of water or 2.0.

Green Timber-Usually weighs from one-fifth to nearly one-half more than dry. Ordinary building timbers, tolerably seasoned, weigh about one-sixth more.

▲ Solid Cube of material-weights of broken or crushed material decrease, for example, see figures given for coal and for limestone.

• * Figures listed are for best conditions (dry, sized and without dust)-The minimum, angle will **increase** as size of particles decrease and with higher moisture content. For other conditions refer to S-A Engineers for definite **recommendations**.

DESCRIPTION	Average Wt. per cu. ft. pounds*	Minimum Angle of Slide †	DESCRIPTION	Average Wt.per cu.it. pounds*	Minimum Angle of Slide †
Air (Atmospheric at 60°F., Under pressure of one atmosphere, 14.7 lbs. per square in., weighs 1/815 asmuch as water) Alabaster (marble) A Alabaster (Real, a compact white plaster of Paris) ▲ Alabaster (Ceal, a compact white plaster of Paris) ▲ Alabaster (Real, a compact white plaster of Paris) ▲ Alabaster (Real, a compact white plaster of Paris) ▲ Alabaster (Real, a compact white plaster of Paris) ▲ Alumina, Sized or briquette. Alumina, Fine, Granulated Aluminum, Sulphate, Ground. Aluminum, Sulphate, Cranular, … Ammonium, Sulphate, Damp Granulated. Argols, Roasted, Pulverized. Ash, Black, Ground " Fly, Powdered " Damp Ashes of Bituminous Coal. " Damp Asphaltum Baassu, Nuts, ground Baasesu, Wet Sawdust Baaske, Wet Sawdust Baaske, Wet Sawdust Baast (Similar to Marble) A. Bathstone, Oolite A. Bauxite, Calcined (Granular without dust). " Ground dried. " Mine run Bentonite, pulverized. Bismuth ▲ Boreax, Dehydrated, powdered " Ground Aried .	$\begin{array}{c} .0765\\ 168\\ 144\\ 15\\ 9\\ .5\\ .5\\ .5\\ .5\\ .5\\ .62\\ .13\\ .5\\ .4\\ .40\\ .63\\ 105\\ .45\\ .35\\ .45\\ .35\\ .45\\ .45\\ .35\\ .45\\ .45\\ .45\\ .45\\ .45\\ .47\\ .144\\ .184\\ .131\\ .70\\ .68\\ .85\\ .50\\ .612. 4\\ .50\\ .612. 4\\ .50\\ .16-26\\ .140\\ .120\\ .75\\ .16-26\\ .140\\ .120\\ .37\ to\ 144\\ .100\\ .27\\\\ .50\\ \end{array}$	31°** 31°** 32° 35°** 32° 40°+5°** 40°+5°** 40° 50° 40°*** 40° 50° 40°*** 40° 50° 40°*** 40° 50° 40°** 40° 50° 40° 50° 50° 40° 50° 50° 50° 50° 50° 50° 50° 5	Cellulose Acetate, granular. Cement, Portland (per Bbl. net 376 lbs.) (per bag 94 lbs.) "Mortar, Portland, 1:2 ½. Chalk, A	$\begin{array}{c} 10 \\ 90-100 \\ 135 \\ 137 \\ 18 \\ 60 \\ 26.4 \\ 15.30 \\ 40 \\ 75 \\ 40-45 \\ 57 \\ 40-45 \\ 57 \\ 40-100 \\ 45 \\ 60 \\ -100 \\ 25-80 \\ 50-95 \\ 100 \\ 97 \\ 52 \\ to 57 \\ 56 \\ to 60 \\ \\ 40-50 \\ 40-50 \\ 30 \\ \\ 23 \\ to 32 \\ 33-36 \\ 112 \\ 155 \\ 190 \\ 75 \\ 6 \\ 52 \\ 80-120 \\ 46 \\ 181 \\ 40 \\ 55-65 \\ 55-65 \\ 72 \\ to 80 \\ 72 \\ to 80 \\ \end{array}$	33° 39° 27° 35° 22° 45° 25° 35°** 35°** 35°** 35° 45° 35° 45° 35° 35° 22° 22° 22° 22° 22° 22° 40° 40° 33° 27″ 40° 31°** 30° 32° 25° 41°
Casein, granular.	38-43	30°	" Fullers, Raw	42	35°

▲ • † ** See notes preceding table.

(continued on foi wing page)

678 UNITS, NOTATION, AND GENERAL DATA

TABLE A7—(continued)

DESCRIPTION	Average Wt. per cu. fi. pounds+	Minimum Angle of Slide†	DESCRIPTION	Average Wt. per cu. ft. pounds+	Minimum Angle of Slide†
Eggs Feldspar, Pulverized, (Crushed)	48 50–60 100	40° 32°	Gypsum . "In Irregular Lumpe	142 82 56	45° 30° 40°
Ferric Sulphate A Ferrous Sulphate (Copperas) A Flint ▲	194 119 162		Hornblende, Solid A. Humus Ice ▲	$187 \\ 30-40 \\ 57.4$	
Fuller's Earth (See Earth) Gelatin, Granulated Glass, Common and Plate A. Batch, Average Mix.	38 161 100	38" 	Ilmenite. granulated "Fine Ground Iron Oxide Pigment	148 120 25	31° 40° 40°
Glue (Pellet) Gneiss, Solid A. " in Loose Piles.	45 168 90–100	25°	Ivory▲ Kalsomine (powder). Kaolin, Green Crushed "Pulverized	114 32 64 22	42° 35° 45°
Grains, Seeds and Cereals Barlev (48 lb. per bushel) Beans, Cocoa "Navy	38 37 54	25°	Kieselguhr. crushed. Lead, #70 Red. "Sificate, Granulated. "Sulphate, Basic Pulverized	15 230 230 184	40° 40° 30° 45°**
Beans, Soy—Cake. ""Meal	45 40 27	32° 27° 40°	Leather▲ Lime, Briquette "Burned, Pebble (sized).	59 60 53	26° 30°
Beans, Soy—Crushed ""Whole ""Split	$34 \\ 45-50 \\ 44$	35° 22° 25°	Lime, Burned, Pulverized "Fine (Spent Dry Carbide) Mason "Burned or Quick, crushed	27 45 17 50	43° 40° 40° 40°
Bran Brewers Grits Buckwheat (46 lbs., per bu.).	33 34.5	30° 24° 25°	" Hydrated Limestone, solid A. " Pulverized	10-25 166 85	42°
Clover Seed (60 lbs., per bu.). Cocoa Nibs Cocoanut Meal Shredded	48 32 32 25	28° 26° 38° 27°	 Mixed Sizes Coarse, Sized Liquids 	105 98	35° 25°
Coffee Beans, Green "Steel Cut Corn, Field (on cob)	42 28 45	25° 23° 20°	Alcobol, Benzine Milk	49.3 53.1 64.3	
" Shelled (56 lbs. per bu.). Corn Flakes	45 12 25 25	22° 25° 36°	Naptna Oils, Vegetable Oi ls, Mineral Petroleum	58.7 57.1 54.8	
Cornneal (50 lbs. per bu.) " Muffin Mixture Cotton Seed. " _{Meal}	40 28 25 33	35° 45° 29° 35°	Turpentine Water Pure Rain Distilled @ 32° F. Bar. 30 in """""" " " " 212° F. "" " "	62.41 f32.35 59.7	
Farina. Feed Gluten " Mol bss es.	44 31 25	29° 34° 40°	Water, Sea. Wood, Spirit. Lithapone, Granulated.	64.08 49.9 70 73	40°
Flax Seed Ground	28 45 35 to 40 26	35° 21° 31° 40°	Magnesium Carbonate Powdered. Masonry of Granite or Limestone, Well Dressed " Brickwork (See Brickwork).	9 165	36°
Grain, Brewers Spent (Sloppy Wet) Grass, Blue, Seed (14 lbs. per bu.)	84 11.5		Metals Aluminum ▲ Babbitt A Brass (7.8 Copper to 8.4 Zinc) ▲	$165 \\ 500-650 \\ 504$	
Hay in Bales Hemp Seed (48 lbs. per bu.) Hominy	20 36 45	210	 # Rolled A. Bronze (8 Copper to 1 Tin) (Gunmetal) A. Cadmium ▲. 	534 552 539	·····
Linseed Meal Linseed Rolled. Malt, Dry " Spent Dry	27 25 32 10	34° 34° 21° 28°	Chromium ▲ Copper Cast ▲ Copper Rolled ▲	432 543 555	
Malt Sugar (Ground) Hygroscopic. "(Unground) ''"" Oats, 32 Lbs. Per Bushel "Rolleti	35 30 26 18	31° 31° 21° 28°	Iron, Cast Aura Magnetite "Iron Ore, Hematite, Magnetite ", Taconite	446 135 115 710	·····
Pablum . "Waste Rice	9 14 50	32° 38° 20°	Magnesium ▲ Magnesium Sulfate (Epsom Salts) Crystals Manganese A	109 55 475	25°
Rye (56 Lbs. Per Bu.). Timothy Seed. (45 Lbs. Per Bu.). Wheat (60 Lbs. Per Bu.). Timothy Seed. Germany German	45 36 48 32 32	23 28° 23° 30° 36°	Mercury at 32° F. ▲ Nickel ▲	849 537 655 489.6	
Granite Solid A. Granite, Gneiss A. Gravel. Green Stone, Trap, Solid A. " " Quarried in Loose Piles	159 175 120 187 107	30°	Tin. Cast A. Zinc or Ypelter A. Mica, Solid A. "Ground Milk, Powdered	439 437.5 183 13.5 40	36° 45°

A $\,^{\bullet}\,^{\dagger}\,^{\bullet\bullet}$ See notes preceding table

TABLE A7-(continued)

	i	-	1		
DESCRIPTION	Average Wt.per cu.ft. pounds*	Minimum Angle of Slide †	DESCRIPTION	Average Wt. per cu. ft. pounds+	Minimum Angle of Slide †
Molasses, Powdered Molybdenumitc Ore, Powdered Mortar, Hardened	21 107 103	45° 40°	Starch, (Powdered). " (Lump & Pelleted) " Tablet, Granular Crystals	25–45 30 40	45° 28° 24°
Mud Wet Fluid Nails and Spikes, 106 <u>I.bs</u> , Per Keg Nitrate of Soda, Pellet Type Granular	108 	····· 24°	Straw in Bales Stucco (Tubed and Untubed), Powdered Sulphur, Pulverized	25 50 50 76	36° to 38° 45°
Nuchar, Granular Paste, Dried, Flaky	22 10	30° 36°	Sugar, Brown	40-50	45°
Phosphate, Powdered	60 60 51 60	40° 30° 45° 26°	" Powdered " Granulated. " Tailings	45 50 57	45° 35° 38°
 Tri-Sodium Pulverized. Rock A Florida #20 Mesh, Air Cleaned. Mono Calcium, Powdered. 	50 200 93 61	40° 27° 40°	Talc, Solid (Soapstone) A " Micaceous, Granulated. Tankage, Ground	169 62 49	36° 32°
Phthalic Anhydride, Flaky Pigment (For Rubber Tires) Powdered Pitch ▲ "Flocka	42 52 70 42	24° 45°	Tar▲ Tartaric Acid (Cream of Tartar), Granular Tin, Oxide, Ground	75 60 100	35° 35°
Plaster of Paris, Powdered Potassium, Chloride ▲ "Sulphate ▲	50 124 167	40°	Titanium Dioxide (paint pigment), powdered Tobacco Stems, Chopped, Coarse Trap, Rock ▲	25 16 187 107 80	45° 23° 40°
Powder, Face Powder, Pudding Pumice, Pulverized	36 40-45 40	45° 36-40° 45°	Vegetables and Fruits Apples (56# per Bushel).	45	
Pyrethrum flowers, coarse ground " spentflowers Pyro, Powdered	20 32 56	30° 40° 33°	" Dried (22# per Bushel) Beets, Shredded Sugar Onions (60# per Bushel)	17 7.0 48	···31°····
Quartz ▲ Kesin, Synthetic (from plant), Crushed Resin and Wood Flour, Powdered	165 40 19	30° 40°	Peas (64 Lbs. Per Bushel)	51 48 33	 20°
Rock, Phosphate Pulv "Solid ▲ "Florida Phosphate #20 Mesh Air Cleaned	60 200 93	40° 	"Meal, Ground "Expeller Cake Ground. "Expeller Cake Chopped	40 32 29	39° 30° 20°
Rosin (From Crude Turpentine) ▲ Rubber, Scrap (Ground) Rutile, Red Oxide of Lead, Fme Ground " Powdered	67 23 132 107	35° 32° 40°	Walnut Shells, Ground (320 Mesh)	21 60	55°
Salt, Granulated " Rock Crushed, sized Sand, Mine run Sand Crasse sized	81 75 90-120 90-100	31° 25° 35°** 30°	Cedar Cherry (Perfectly Dry) Chestnut (Dry)	22 44 30	·····
" Very fine " Core" " Voids Full of Water.	90-1 10 65 10 to 130	32° 39° 45°**	Chips (Dry) Cypress Elm (Perfectly Dry)	15-32 32 35	22°
Sandstone, Solid ▲ " Quarried & Piled Serpentine (Talc) ▲	147 82 169	· · · · · · · · · · · · · · · · · · ·	Fir, Eastern Hemlock (Perfectly Dry) Hickory (Perfectly Dry).	25 29 48	
Shales, Slate ▲ Silex	172 70 80	44° 45°	Lignumvitae, Dry Locust, Dry Mahogany	41 to 83 46 35–53	
Slag▲ " Furnace Granulated " Birmingham	60 to 180 122 82	25° 25°	Maple, Dry Oak, Live, Perfectly Dry. " Red, Perfectly Dry	33-40 59 41 to 45	
Slate, Solid A. "Fine Ground "Granules, Flaky. "Flour	175 82 87 45	35° 28° 45°	Pine, White Perfectly Dry "Yellpw Perfectly Dry Short leaf " Long leaf	27 38 44	
Snow (Fresh Fallen)	5 to 12 15 to 50		Poplar, Dry Red Wood, California Dry	29 26 to 30	
Soap Chips Soapstone, Talc ▲ "Fine Ground	5-15 169 60	28–32° 40°	Sawdust, Dry. "Ground	20	45°
Soda Ash, Light " Dense " Briquette " 4 and Silica Sand (1 to 1.85)	25-35 66 50 80	37° 32° 22°	Springles Per 1000, Short 900 Lba. Long 1400 Lbe Spruce, Dry, California. Sycamore (Perfectly Dry).	25 37	
" Bicarbonate	39-55 72	42° 25–50°	Walnut, Black (Perfectly Dry) Wool Mineral. Jinc calcines, Powdered "Ore recasted Granular	42 10 85 110	30° 54° 38°
Antimonate Crushed Carbonate, Powdered (Soda Ash) Nitrate, Granular Phosphates	49 68	31 ^{8**}	" Oxide" " "Leaded, Ground" " Sulphate, Powdered	20 25-40 72	46° 50° 44°
" Sulfite, Powdered " Sulfate (fine and lumps)	96 88	40° 31°	Jonolite, Fine, Granular	7	30°

▲ • † ** See notes preceding table.

Data of Stephens-Adamson Co., Catalog 66, Aurora, IL, 1954. See also Table 5.3.

TABLE A8. Petroleum Products, Typical Compositions

		Sun	nmary	of I	Produ	ıct 7	[ypes	s Pro	duced	l Fro	m Pe	etrole	um								
Numher of Carbon Atoms Boiling Point of Normal Paraffin at 760 mm	C1 -161 -259	C2 -8{ -12'	C3 -42 -44	C4 -0.5 +31	Cs +36 97	C ₆ 69 156	°7 98 209	C8 126 258	с s 151 303	C10 174 345	C11 196 384	C12 216 421	C13 235 456	C14 253 486	с15 270 519	C 1 6 2 8 7 5 4 8	C17 302 575	C18 316 601	C19 329 625	C20 343 649	> C21
Liquefied Petroleum Gas Precipitation Naphtha VM&P Naphtha Mineral Spirits Reformate Gasoline Kerosene, Diesel Fuel Aviation Turbine Fuel.					-	+		+	+ + + + + + + + + + + + + + + + + + + +		-										
Gas Oil, Fuel Oil. Transformer Oil. Lubricating Oil. Asphalt, Pitch. W a x																		•			

Source: Humble Oil Co.

Appendix B-

EQUIPMENT SPECIFICATION FORMS

Specification forms make provision for key data of the equipment and its position in the process, including design characteristics and required performance. A major purpose is a basis for price quotations by suppliers of the equipment.

Although a few forms have been prepared by industry-wide organizations such as TEMA, Hydraulic Institute, and API, most have been prepared for their own use by engineering contractors and design divisions of chemical process organizations. Most private firms have their own libraries, but the merits of some forms have been recognized and copied widely so that much uniformity exists.

The selection of forms in this Appendix is made available through the courtesy of individual companies identified on each form, although one major contributor did not wish to be identified. They have been collected over a period of time and consequently may not be the latest forms in use. For the most part, only the most often used kind of equipment is represented, except for the few items in category K.

A related kind of information is the suppliers inquiry forms of Appendix C.

Following is a listing of the equipment specification forms, included in this Appendix. The number appearing in italic is the page number on which the form appears.

- A. Agitators and mixers
 - 1. Agitator data sheet 682
 - 2. Agitator specification sheet (2 pp.) 682
 - 3. Tank mixer, top entering 683
 - 4. Tank mixer, side entering 684
 - 5. Syphon pum(pmixing tee) 684
- B. Compressors (see also items 44 and 45)
 - 6. Centrifugal compresspecification 685
 - '7. Reciprocating compressor specification sheet 685
 - 8. Fan or blower data sheet 686
 - 9. Vacuum pumpata sheet 686
- 10. Steam ejector data sheet (2 pp.) (see also item 22) 687 C. Drivers
 - 11. Electric motor data sheet 688
 - 12. High voltage induction motors (less than 1500 HP)
 - **13.** Gas engine data 689
 - 14. Generabuspessen turbine specification 689
 - 15. Gear drive data sheet 690
- D. Heat transfer equipment
 - 16. Exchanger specification sheet (TEMA) 690
 - 17. Air cooled exchanger specification sheet 691
 - 18. Fin fan heat exchangers 691
 - 19. G-fin tank heater specifications 692
 - 20. Compressor cooler data sheet 692
 - 21. Plate and frame exchanger specification sheet 693
 - 22. Steam surface condenser and ejector data sheet (4 PP.) 693
 - 23. Fired heater specification sheet 695
 - 24. Wiped film evaporator data sheet (4 pp.) 696
 - 25. Rotary vacuum dryer data sheet (2 pp.) 698

- 26. Boiler data sheet 699
- 27. Package boiler data sheet 699
- 28. Induced draft cooling tower 700
- E. Instruments
 - 29. Instrument specification 700
 - 30. Differential pressure type flow instrument 701
 - 31. Temperature and pressure instruments 701
 - 32. Level instruments specification sheet 702
 - 33. Gauge glasses and cocks specification sheet 702
 - 34. Control valve specification sheet 703
 - 35. Valve operator specification sheet 703
 - 36. Relief valves 704
- F. Pumps
 - 37. Centrifugal pumpata sheet 704
 - 38. Centrifugal pump 705
 - 39. Reciprocating pump specification sheet 705
 - 40. Rotary pumpata sheet 706
 - 41. Rotary pump data sheet (2 pp.) 706
 - 42. Motor driven proportioning and reciprocating pump data sheet 707
- G. Refrigeration
 - 43. Packaged refrigeration system 708
 - 44. Reciprocating compressor 708
 - 45. Positive displacement rotary compressor 709
 - 46. Flowsketch 709
 - 47. Instrumentation and controls 710
- H. Solids handling equipment
 - 48. Vibrating feeder data sheet 710
 - 49. Screw conveyor data sheet (2 pp.) 711
 - 50. Bucket elevator data sheet 712
- I. Solid fluid separating equipment
 - 51. Pressure leaf filter 712
 - 52. Intake air filter 713
 - 53. Line filter specification (2 pp.) 713
 - 54. Centrifuge data sheet (2 pp.) 714
 - 55. Vibrating screen data sheet (2 pp.) 715
- J. Vessels

- 56. Column specifications 716
- 57. Bubble tower specification 717
- 58. Vertical vessel data sheet 717
- 59. Horizontal vessel data sheet 718
- 60. Glass lined vessel (2 pp.) 718
- 61. Fractionator tray design data sheet 719
- 62. Tray specification 720
- 63. Tower internals data sheet 720
- K. Other equipment
 - 64. Chlorination system 721
 - 65. Coalescers 7 2 1
 - 66. Deaerator (spray-tray) (2 pp.) 722
 - 67. Demineralizer 723
 - 68. Cation or anion exchanger sketch 723
 - 69. Air dryers (heatless) 724
 - 70. Mechanical scale data sheet 724
 - 71. Rotary valve data sheet 725

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	P
	AGITATOR
	SPECIFICATION SHEET
KS	Page 1 of 2.
REMAR	Project
	1 Service of Unit Pos. No 2 Number Required Location
	3 TypeMountingManufacturer
Y.R.Y	
ž	6 Class of Operation 7 Type of Agitation Period of Agitation
DATE	c Cycle. Batch: Smallest gql, Normal gql,
UPRV.	9 Continuous: Rate of flow GPM
¥D S	10 Liquid Liquid Solids or Gas
ر	I Material
•	12 Quantity 13 Viscosity °F °F °F °F
REV	14 Specific Gravity @ °F @°F @°F
	1 6 Solids Characteristics: Size Description
_	17 Mixture Characteristics: Sp.G?F Viscosity cp (@?F
	19 Operating Conditions: Normal Temp°F Press, psig
	Max. Temp °F Press psig 20 Vessel Volume: Gals; Diam: <u>: Str.</u> Shell:
	21 MECHANICAL DESIGN DATA
i	(Attached)
	22 Vessel Ref. Druwing:; No; Size; Speed nrpm; Spacing
	24 Shaft Length-in. Diameterin.
	26 Seal Type
	27 Seal Mfr.
	28 Lubricant Type Lubricant
	30 Stuffing Box Type
	3 1 StabilizerBaffles: No.:Width:
	32 Mounting Nozzle: Rating: Facing:
	33 Size of Nozzle for Impeller Installation:
	DATE BLDG. NO.
	DRAWING NO.
	þ.



	TANK MIXER TOP ENTERING (TYPICAL)
	SERV I CE
	SOLIDS: TYPVCONCENTRATION.
	SP. GR. Q OP. TEMP DMAX. VISCOSITY O OP. TEMPCMAX.
	DES GN PARAMETERSI PRESSTEMPDESIGN LEVEL
	TANK- ITEM NO CAPACITY SIZE
	TANK MIXERJIEM NO DESIGN
0	MANUF
12	REDUCT. GEAR EFFIC S E R V I C E FACTOR - C A L . P R I . P U M P I N <u>G C A P</u> . DES GN LOADS* TORQUESTAT I C MOMENT
3	DYNAMIC MOMENT DOWNWARD WE I CHT
4	IMPETIERS IN NO TYPE NATIONST
õ	ACTUAL H. P. ABSORBED & IMPELLER PITCH - R E M O V E I M P H R U
7	SHAFT: DIAMETER LENGTHMAT' L. OF CONST
β	OUTPUT SHAFT BEARINGETYPEAFBNAL-TO LIFE-LUB. TYPE
ő	MOUNTING FLANGE TYPE I Z E MAT'L OF CONST.
1	SEAL TYPE:MODEL NOMODEL NO
2	MAT'L. OF CONSTIROTATIN <u>G</u> FACESINTL'S. METALLICFLEX.
3	BALANCED. YESINO: TANK SHUTOFF. YESINO. FLUSH FLU ID
5	GEAR HOUS I NGI V-BELT/GEAR. MAT' L. OF CONSTA G M A G E A R C L A S S -
6	TYPE OF SUPPORT GEAR BOX LUB. TYPE
7	totor: Item NoMtrModel NoUty =
8	Enclosure I n s u l a r i o n : Temp. Rise OC above OC Ry & SE
0	Space Heater: VoltsPhaseHertz
1	Current Eff. P.F. Full Brg. Allow-Thrust Space
2	No KU Load Potor TA A A PPM No (1) B/KG IB/KG Watts
4	
5	WEIGHT: SHIPPING
6	PAINT SPECIFICATION
8	NUTESI
9	
0	
1	VENDOR TO SHOW LOC. AND TOP VIEW SHOW LOC. OF
23	HIG.FLG: DIMENSIONS MIG.FLG: DIM. OF ROTATION
4	
5	
6	
8	
9	
0	
	5 KELLOGG MANAGING OFFICE ENGINEERING OFFICE
	DATA SHEET/SKETCH
	CLIENT
	LOCATION008 NO
1	no. DATE ENCINEER CHK/REVIEW APRROVE FOR N.N.O. PAGE D.E.

[TANK MIXER SIDE	ENTERING (TYPICAL)
 2 3	SERVICEI	INDOOR/OUTDOOR ENGLISH/METRIC UNITS
4	DPERAT INC PARAMETER9 PRESS.	TEMP
5	SP. GR. Q OP. TEMP ØMAX	VISCOSITY & OP. TEMPQMAX
6	DES GN PARAMETER9 PRESS.	TEMPDES I GN LEVEL
6	DESIGN	BAFFIES
9	TANK MIXER-ITEM NO.	QTY
10	MANUF.	MODEL NO.
	REDUCT. GEAR EFFIC SERVICE	FACTORCAL. PR PUMP NG CAP
13	DESTIGN LUADSI FORQUE	DOWNWARD WE LIGHT
4	FIRST CRITICAL SHAFT SPEED	
15	PROPELLERI SIZE TYPE	
16	ACTUAL H. P. ABSORBED @ PROPELLER	PITCH REMOVE PROP. THRU
17	SHAFTIDIAMETER LENGTH	MAT' L. OF CONST.
19	OTHER BEAR INCS. TYPE	
20	MOUNTINC FLANGE: TYPE	S I ZEMAT' L. OF CONST.
21	ANGLE -	UNTEDIR. OF ROTATFROM MOTOR END
22	SEAL TYPE:MANUF.	MODEL NO.
23	MAT L. OF CONSTRUTATING FACEST	I NIL' \$, METALI C FLEX
25	SEAL DES IGN PRESS.	TFMP.
26	GEAR HOUSING V-BELT/GEAR MAT'L.	OF CONSTAGMA GEAR CLASS-
27	TYPE OF SUPPORT	GEAR BOX LUB. TYPE
28	Area Class Gr Div /Non-Ha	zardous: Volts'- Phase Hertz
30	Enclosure Insulation ; T	emp. Rise ^Q above ^Q By . Q .S.F.
31.	Space Heater: VoltsPh a s	se Hertz
32	Current Eff. P.F. Full	Al low-Infrust Space
33 34	No KW load Rotor 4 4 4 4 8 RPN	No (1) LB/KG LB/KG watts
35	ixer	
36	WEIGHT SHIPPING	INSTALLED
37	PAINT SPECIFICATION	
38 39	NOTES:	
40		
41	VE	NDOR TO SHOW LOCATIONS
42		ID ANGLE.
43		MENSION
45		(+ + -
46		
47	G GEAR 4.	
40 /0		
50	<u> </u>	
i	6	MANAGING OFFICE ENGINEERING OFFICE
	2 4	
		DATA SHEET/SKETCH
		ITEM
	0	ITEM NO SPEC. NO
1	NU. DATE ENGINEER CHK/REVIEW APRROVE	REQ' N. NO. PAGE OF

		PROCESS DESIGN SECTION CORPORATE ENGINEERING		SYPHON PUMP DATA S	p (mixing tee) Heet
		EQUIP. POSITION NO.			
		SYPHON PUMP NO.			
	.≚I	FLOWSHEET NO.			
	1	PROCESS DATA			
		INLET FLUID #1			
		DESCRIPTION			
		SPECIFIC GRAVITY			
		VISCOSITY; CP			
	.H. el	PRESSURE; PSIG			
		TEMPERATURE, F			
	DATE	FLOW			
		INLET FLUID #2			
	Sura	DESCRIPTION			
	HKD	SPECIFIC GRAVITY			
	10	VISCOSITY. CP			
	BY	PRESSURE; PSIG			
\Diamond	چ <				
<u> </u>					
		TEMPERATORE.			
ģ	Ϋ́,				
ы Т	ŝ	MATERIALS OF CONSTIN			
PRO	DES	SIZE NO			
	ž	PIPE CONN'S. INCHES			
	ED 1	PRESS INLET			
	HECI	SUCT. & DELIVERY			
	°				
	ΒY	-			
	MM	NOTES			
	DRJ				
	a CL				
ATE	LTN N				
<u> </u>	<u>. : </u>	JOB NO	PLANT LOCATION		
		PROJECT NO	REF DWG NOSEE	ABOVE LIST	ISSUE
				(DATE BLDG. NO.
		POSITION NO SEE ABOVE LIST SERIES	<u> </u>	STEP NO	$\underline{\frown}$
				DRAWING N	
					N
E 14	4089	POSITION NO SEE ABOVE LIST SERIES		STEP NO.	

GENERAL ENGINEERING DIVISION FORM 66-55 3H 7 34 G.S. SHET NO. THE FLUOR CORPORATION LTD. DATE VENOOR NUST FURNISH ALL PERTINENT DATA FOR THIS BEFECIFICATION SHEET ERFORMED RETURNING ALL PERTINENT DATA FOR THIS

REV.

CHK'D.

ITEM NO.	SERVICE						JOB N	0.	
NO. REO'D.	DRIVE			MA	NUFACTURER				
GAS HANDLED			-		SIZE & TYPE				STAGES
CORROSION FACTORS			· · · · · · · · · · · · · · · · · · ·	_	DESCRIPTION				
OPERATING CONDITION	1:	2:	3:	_		COMPR	ESSOR MAT	ERIALS	
CAPACITY					CASE				
CFM @ INLET CONDS					DIAPHRAGMS				
WT. FLOW, LBS/HR.			-	_	GUIDE VANES				
(M ² SCFD) (SCFM)*					INTER STAGE LAB	TRINTHS			
TEMP., INLET. °F					IMPELLER HUBS	COVERS			
DISCH, *F				_	IMPELLER VANES				
PRESS., INLET, PSIA				_	SHAFT				
DISCH, PSIA	_			_	SHAFT SLEEVES				
DIFFER, PSI				-	BALANCE DRUM C	RDISC			
COMP. RATIO				_		CONST	RUCTION D	ETAILS	
MOLECULAR WT.			l	_	CASE: SWP		PSIG, MAX. H	YDTEST	PSIG
5P. GR. (AIR=1)				_	SPLIT: (HO	RIZONTAL:	VERTICAL: E	ARREL)	
REL. HUMIDITY, %			<u> </u>	_	SUPPORT:	(FOOT: PEL	ESTAL: BRAG	CKET)	
"K" VALUE, CP/CV					IMPLELLER: TYPE	: (OPEN: S	EMI . ENCLOS	ED: ENCLOSE	0)
COMPRESSIBILITY. "Z"					CON	STR: (CAST	FORGED: R	VETED: WEL	DED)
ELEV. ABOVE SEA LEVEL		_		-	DIAM	ETER	", VANE	THICKNESS	
BHP (INCL. GEAR LOSS)				-	SHAFT: DIAM. AT	IMP	", AT BE	1G\$	
RРM				_	SPAN; C-	C BRGB	", IMP C	VERHANG	
IMP. TIP VEL. FPS		_			CRITICAL	SPEED			RPM
WATER RATE, #/HP/HR					BRGS: LOCATION	(INTERNAL	: EXTERNAL)		
	1				RADIAL; T	YPE	: PROJ	AREA	59 "
MAX. CAP., CFM @ INL.		: MAX. BHP			THRUST; 1	YPE	; EFF.	AREA	\$Q "
MIN. CFM. (SURGE POINT)					SEALS:				
CONTROL: (SPEED: SUCTION	N VALVE: INLET VA	NES)		_					
SOURCE: (MANUAL: FC:	PC: TC)				COUPLING			CP	LG GRD; YES, NO
STEAM: PRESS.	PSIG: TEMP.	۲; I	хн.	_	BASE PLATE:				
POWER:VC	DLTS: PH.	CYC., COOLING	WATER	۰,	-	,			
MEGR SHALL SUPPLY THE F	OLLOWING	DRAWING	NO		NOZZLES	BIZE	RATING	FACING	LOCATION
PERFORMANCE CURVE				-	INLET				
SECTIONAL DRAWING					DISCHARGE		-		
OUTLINE DRAWING					DRAINS				
ROTATION FACING CPLG:	CW: CCW	·			TURB. INLET				
LUBRICATION	SYSTEM	COMPRESSO	RWEIGHTS	_	TURB. EXHAUST				!
COMBINED WITH DRIVER; Y	ES, NO	WT., TOP HALF		LBS					
TWIN OIL COOLERS		WT., ROTOR		LBS	TESTS: MECHANI	CAL RUN IN	; YES, NO		
TWIN OIL FILTERS		WT., COMPR. BARE		LBS	WITNESS	D PERFORM	ANCE; YES,	NO	
MAIN OIL PUMP		WT., BASEPLATE		LBS	OVERSPE	ED IMPELLE	RS TO		RPM
AUX. OIL PUMP		WT., ACCESSORIES		L89	OVERSPEI	ED ROTOR T	0		RPM
OIL PRESS. SHUT DOWN		SHIPG WT. LESS DRIV	/E	LBS	HYDROST	ATIC TEST P	RÉSS.		PSIG
OIL TEMP. ALARM		FLOOR SPACE			SHOP INSPECTIO	N: YES, NO			
						RE	DUCTION G	EAR	
DRIVER: MAKE					MFGR:				
ТҮРЕ			1		TYPE:				
RATED HP			+		ARRANGEMENT	(COUPLED)	(INTEGRAL)		
RPM					LOW SPEED SHA	FT,			RPM
SHIPPING WEIGHT	—				RATING				нр
CHIPPING BOINT					SERVICE FACTOR				
SHIPHENT MONTHE	_		<u> </u>		LUBRICATION:				
REMARKS									

THE C.W. NOFSINGERCOMPANY Kansas City. Missouri

-								EFT NO	
		RECI P	ROCAT	I NG	COM	PRESSOR	SI		
		SPE	CI FI (CATION	N SE	IEET			
сш	STOWER						C1		
10	CATION						G A	161 JOB NO	
SE	RVICE			Ν	IO. REO	UIRED	BY		
N.F	R MOI	ELNO		s	ERIAL	NO.	RE	VISED	
F									
	UPERATING CUNUT	110#2					CONSTRUCTIO	IN DETAILS	
1	VAPOR GAS	MOL.WT (I	NCL H ₂ 0)	FRAME	RATING:	HP	RPM	
2	ELEVATION, FT.	BAROMETER	R. PSIA		LUBE:	TYPE	F	ESERVOIR CAP	.NEN
3	INDOOR - OUTDOOR	HEATED:	YES - N	10	FLO	W, GPM		EMP. IN/OUT,	°F
4	CORR/EROS PROPERTIES				WATER	GPM		EMP. IN/OUT,	°F
5	0	NORM	R	TED	WEIGH	TS: COMP	R	ORIVER	
6	MMSCFD/SCFN (14.7 PSIA-60 °F)				FLY	WHEEL		SHIPPING	
12	WT.FLOW, LB/HR		_		TESTS	SHOP IN	SPECT	PERFORMA	NCE
8	SUCTION, PSIA AT F		_		RAD	ROSTATIC,	PSIG	WITNESS	
1	DISCHARGE PRESS, PSIA	-	_		ACCES	S: SUCT.V	A. UNLOADING,	MANUAL-AUTO	1
10	CP/CV, K (SUCT)				CLE	ARANCE PO	CKETS, FIXED	VARIABLE	
ш	COMPRESSIBILITY, Z (SUCT)				OUTLI	NE DWG.NO		API 618:	YES-NO
		PE	RFORMA	NCE (E/	ACH ST.	AGE)			
12	STAGE - CYLINDERS/STAGE		IST		2ND	3	RD	4TH	5TH
13	DISCHARGE PSIA PF								
14	RPM								
15	BORE & STROKE								
16	CYL.DISPLACEMENT, ACFM								
17	CLEARANCE, %								
18	COMPRESSION RATIO								
19	VOLUMETRIC EFF.								
20	CAPACITY, MNCFD								
21	BHP/STAGE								
22	VALVE: TYPE								
23	LIFT (IN/OUT), WILS								
24	GAS VELOCITY, FPM								
25	TEST PRESS (WORKING P x 1.5), PS	16							
26	ROD LOAD: TENSION, PSI								
27	COMPRESSION, PSI								
28	INTAKE NOZZLE: SIZE-RATING-FACI	NG							-
29	DISCH. NOZZLE: SIZE-RATING-FACI	NG							1
	MATERIALS								
30	CYLINDER/LINER								
31	PISTON/RINGS		1						
32	RODS								
33	VALVES/SPRINGS								1
34	ROD PACKING MOTOD DELVED						TIDE INC	IVER	
2		n ev			-		TORDINE DA		
31				NED	01		100		-
37	ENCLOSURE RPN			KP		R P N	WAT	ER RATE LR/HS	- HR
, 3	8 HP SF			INLET	STN P	RESS, PSIG	/ NORN	MA NA	x
Ť	9 FRAME INSULAT	LION		INLET	STM. TE	NP, ^O F	N O R N	MA	X
4	D VOLTS/PH/HZ TEMP R	ISE ^o f		EXHAUS	ST <u>s</u> tm.	PRESS, PSII	NORN	MA	x
41	BEARINGS LUBE			BEARIN	IGS		LUB	1	
42	DRIVE TRAIN: BY MF	R .		NOZZLE	S	\$ ZE	RATING	FACING	LOCATION
43	MODEL RA	T10		INLET					
44	TORSIONAL STUDY: YES-NO BY			EXHAUS	at 🗌				1
45	SEPARATE SPEC IF: DIESEL, GAS			AP1-61	11: YE	S - NO			

VENDOR TO SUPPLY INFORMATION MARKED

*--STANDARD' CONDITIONS ARE 60"F & 14.7 PSIA

	PROCESS DESIGN SECTION FAN OR BLOWER CORPORATE ENGINEERING DATA SHEET
	OPERATING CONDITIONS
s s	
ž	ar. dn. folin = 1.0 Polynol = olynol
M N	STD. CFM: DISCH. PRESS: PSIA
R	FLOW TEMP: OF DIFFERENTIAL PRESS:PSIA
	MECHANICAL DATA
1	
	NOZZLES SIZE ASA RATING FACING POSITION METHOD OF SEALING:
Ť.	SUCTION LUBRICATION
Å.	DISCH. COOLING
`	COUPLING MFG:
5	
à	
è	MATERIALS OF CONSTRUCTION
3	CASE: WEAR RINGS:
¥	IMPELLER: COUPLING:
- Ĵ	SHAFT: GASKETS:
2	SHAFT SLEEVE:
~ 5	PERFORMANCE
< > [#	
	NO. OF STRAGES MOTELT VERSION FACING COUPLING END:
E de	MOTOR DRIVE DATA
80	ITEM NOMTD, BY VOLTS/PHASE/CYCLE/
۵	H P - R P I F R A M E BEARINGS:LUBE:
	MFG FULL LOAD AMPS:
ŝ	TYPE:INSULL,R, AMPS:
CKED	ENCLOSURE: CLASS I GROUP D
CHEC	MISCELLANEOUS
<u> </u>	OUTLINE DWG. NO ACTUAL IMPELLER DIA
>	SECTION DWG. NO TEST CURVE NO
ž	FAN OR BLOWER SERIAL NO WEIGHT POUNDS NET
MAR	REMARKS
ā	
jα	
P 12	
2	
	PROJECT NO BEE DWG NO
A	DATE BLDG. NO.
38	POSITION NO SERIES STEP NO.
2	
3	DRAWING NO.
	r (* 1

∣ ⊧						
	OPERATMG	CONDITIONS				
s.	GAS HANDLED:	SEAL FLUID:				
A A K	MOLECULAR WGT:					
Ē	DISCHARGE PRESS: "Hg VACUUM	DENSITY.				
æ		VISCOSITY:CPS				
	DISCHARGE TEMP:	VAPOR PRESS: PSIA				
	CAPACITY:LB./HR.	HEAT EXCHANGER COOLING WATER:				
	CAPACITY: SCFM	INLET TEMP:P				
	"K" VALUE:	INLET PRESS PSIC				
2	RELATIVE HUMIDITY:%	ALLOWABLE PRESS. DROP PSI				
A P P.	ALTITUDE FT.					
w	MECHAN	CAL DATA				
DAT	MFG	RECEIVER				
2	ŞIZE 🔥 TYPE:	CAPACITY:GALS.				
ŝ	NOZZLES SIZE ASA RATING FACING POSITION					
4¥0	DISCH.	CODED: ON NON CODED: O				
ô	HEAT EXCH.	HEAT EXCHANGER:				
à	COOLING	SURFACE AREA SQ. FT				
>	SEAL LIQUID	DUTY:BTU/HI				
E I		TUBE DESIGN: PSIG-TEMP PSIG-TEMP.				
	TYPE:	MAX. CASING PRESS:PSIG.				
F		CONSTRUCTION				
┓	PUMP:	HEAT EXCHANGER:				
	CASING:IMPELLER:	SHELL:				
	SHAFT:	TUBES:				
_						
⊧	PERFOR	MANCE				
	RATED CAP <u>ACITY, CFM @ "Hg.</u> VAC.	SEAL FLUID FLOW RATE: GPN				
	NO. OF STAGES.	COOLING WATER FLOW RATEGPW ROTATION FACING PUMP END				
	RATED BHP					
h	MOTOR DRIVE DATA	ACCESSORIES				
ľ	тем <u> NO.</u> мтр. ВҮ					
	VOLTS/PHASE/CYCLE/					
	HPR PFRAME - BEARINGS:	SOLENOID VALVE:				
	TYPE INSUL: LR AMPS:	SEPARATOR:				
	ENCLOSURE. CLASS I GROUP					
		N SC.4LE				
	PROJECT NO REF. DWG. NO.					
	TITLE	DATE BLDG. NO.				
ja j	POSITION NO. SERIES	STEP NO.				
- 1						

	ľ	PROCESS DESIGN SE	CTION		STEAM EJECTOR							
		CORPORATE ENGIN	EERING	±,		DATAS	HEEI S	HEET 1 of 2				
		Service			Manufacturer							
	5	No.of Unite		M	odel No			·				
	-ă-	PROCESS CONDITIONS										
	a a a	Non-Condensables (b/m										
		Condensables Vapore (Air etc.)		1b./hr. M.)	N							
		Steam		ib.fhr.								
		Suction Conditions at Ejector Abs	Press.	ň	imHg. Temp	F.						
		Discharge Conditions at Ejector:		p	sig Temp	°F.	10-	4				
	- []-	Expected Air Leakege		CUIT. N h/hr	IAX. EVACUATIN	g Time:		ites				
	Č.A	Duty		hrs /day		days	week					
	- YF	Steam Conditions Normal : Press										
	E S	Consumption at Rated Load		b./hrt Presspsig TempPF.								
	à	Water Pressure Normal: psig		Min:	PEIG Te	mp						
	1	Consumption of hated L o a d :	MEQUA									
	- 8-	EJECIOR	1st Stage	2 nd Steam	3 rd Stege	4 th Stage	5 th Stege	6 th Stage				
	Ĭ	Size			1							
-	Ť	Suction Conn.Size										
	à	Discharge Conn. Size										
$\overline{\wedge}$	2	Steam Conn. Size						-				
\geq	~	Test Pressure, paig						1 1				
		-MATERIALS: -						1				
		Suction Chamber										
-		Steam Chest										
DNG.	P84	Steam Nozzie										
7	51							1				
ŭ.	30	Location: (Precondenser.Intercon	denser etc.)									
		Size	'									
	e e	Condenser Water Rate gpm										
	EX.	Outlet WaterTemp. F.										
	E	Iniat Water Pressure Regid; paig	1									
		Vapor Outlet Nozzie Size										
	ΒY	Water Inlet Nozzie Size			1							
	MM	Water Outlet Nozzle Size										
	R O	Design Pressure, psig										
	l	Test Pressure, psig		L								
	НО	Shell		h								
μ	TIAT	Battles										
۲d	ÎN Î	Spray Nozzle										
		JOP NO	PLANT	LOCATION .	1	ł .	\frown	SCALE				
		PROJECT NC	REF	WG NO		[ISSUE					
) <u></u>				(BLDG, no.				
		POSITION NO. SER	IES	STEP	NO	<u>- </u>						
						DRAWING NO).	\sim				
								ΝZ				
								1 V_				

	PROCESS DESIGN SECTION CORPORATE ENGINEERING	STEAM EJE DATA SHEE	STEAM EJECTOR DATA SHEET SHEET 2 of 2										
	MECHANIC	AL DESIGN CONT	D.										
	SURFACE CONDENSER	· · · · · · · · · · · · · · · · · · ·											
RXS	Location Precondenser, Intercondenser, etc.												
NA N	Size.	· · · · ·											
1	Tubes Ne vOD z BWG vie n g t bi	x x x	X X X	* * *									
	Water Flow Rate:gpm												
	Water Temp. In/Out °F	1	1	/									
	Water Sid. AP: psig												
	Tubeside Nozzle Sizes; In./Out	/	/	1									
2	Shellside Nozzle Sizes; In/Out	1	/	/									
H.d.	Design Press;psig Shell/Tubes		/	/									
	Test Press; psig Shell/Tubes	/	/	/									
E,	MATEBRIALS												
a	Tub Shart	<u>├</u>											
ž	Channel	<u> </u>											
S.	Shell	<u> </u>											
ž	Sheli Baffles												
- 0	Impingement PI.14												
۱	REMARKS												
$\overline{}$	Unit to have complete stability at zero load and to be designed for 25% overcapacity												
2	Teil Pip furnished by												
"	Interconnecting piping by												
<u> </u>	JOB NOPLANT PROJECTNOREF. [TITIF	LOCATION		SCALE									
	POSITION NOS E	r i E 8 - 8 i E P	DRAWING NO.										
			1	N									

162 B 20

				IE LOS ANGELES 58. CALIFORNIA SHI	ZETO
			ELECTR	NIC MOTOR DATA SHEET	Ľ,
		1	CUSTOMER	EQUIP. NO.	
		2	PLANT LOCATION	FILE NO.	
	11	3	SERVICE		
		4	MANUFACTURER	TYPE FRAME NO	
		8	INDUCTION MOTOR	SYNCHRONOUS MOTOR	
		•	HORSEPOWER	HORSEPOWER	
		7	RPM VOLTE	RPM	
			PUACTO	VOLIS	
		10	CYCLEB	CYCLES	
		11	FULL LOAD AMPERES	REQUIRED TORQUE	
		12	LOCKED ROTOR AMPERES	LOAD WR ²	
		13	STARTING TORQUE - PERCENT	FULL LOAD AMPERES	
		14	BREAKDOWN TORQUE - PERCENT	LOCKED ROTOR AMPERES	
		15	FULL LOAD RPM	STARTING TOROUS . PERCENT	
		16	EFFICIENCY - PERCENT	PULL-IN TOROUE - PERCENT	
		17	FULL LOAD	PULL-OUT TORQUE - PERCENT	
		18	* LOAD	POWER FACTOR RATING	
		19		FIELD AMPS @ FULL LOAD	
		²⁰		EFFICIENCY - PERCENT	
		3.0	* 104D	FULL LOAD	
		23	Va LOAD	Vi LOAD	
	-	24	ELEV. ABOVE SEA LEVEL FT.	POWER FACTOR	
21U>	٥z	25	SERVICE FACTOR	FULL LOAD	
		26	TEMPERATURE RIGE	% LOAD	
		27	BOTATION FACING END OPP. COUR	Va LOAD	
		28	CLASSIFICATION	ELEV. ABOVE SEA LEVEL - FT.	
	ΤV	29	GENERAL PURPOSE	SERVICE FACTOR	
-++	F	30	SPLASH PROOF	ROTATION FACING END OPP. COUP.	
	à	31		ENCLOSURE	
		32	TEFC EXPLOSION PROOF	RADIAL BEARINGS	
		33			
	L	34			
44440>		38	SHAFT - SOUD HOLLOW	WEIGHT	
			SHAFT DIAMETER	OUTLINE DRAWING	
		38	LOAD THRUST - LBS	SERIAL NUMBER	
DATE	_	39	MAX. ALLOWABLE THRUST	EXCITER - TYPE	
		40	TYPE INSULATION	DRIVE	
BY		41	RADIAL BEARINGS - BALL OR SLEEVE	(IF MOTOR, DESCRIBE AT LEFT)	
	_	42	THRUST BEARINGS - BALL OR SLEEVE	RPM	
CHECK	.0	43	LUBRICATION - OIL OR GREASE	VOLTS. D.C.	
ENGINE	-	44	DRIVE CONNECTION	AMPS @ FULL LOAD	
2E		45	BASE	SHUNT OR COMPOUND WOUND	
PROJ. EN	IG.	46	WEIGHT	SERVICE FACTOR	
		47	OUTLINE DRAWING NUMBER		
	_	48	SERIAL NUMBER	ENCLOSURES	
		49		TYPE INSULATION	
		50			
		51	COOLING WATER @ "F & GPM	BASE	
		52	FORCED FEED LUBRICATION	WEIGHT	-
		23	PUMP BUILT-IN OR MOTOR DRIVEN		
		54			

		HIGH VOLTAGE INDUCTION MOTORS (LESS THAN 1500 HP/1119 KW)
ETOF	1	O Motor Design Data O English OMetric Units O Shop Inspection and Test
Ê	2	Mfr. Model No Required Witnes. J
· ·	3	Serial No (tyShop_Inspection OO
1	4	Driven Equip. Testing Per NEMA(MG-1) O
	5	Altitude Amb Tomp May QC Min 90 Section List Company
	5 7	Aititude Auto, temp. rdax special fests Listed below:
	8	Area Class Gr Div /Non-bazardous
	9	Basic Data:
	10	Volts Phase Hertz Couplings Supplied By:
	11	Nameplate H.P./KW S.F. Mfr. Model No.
	12	RPM OMotor Mfr. ODriven Equip.Mfr. OPurch.
	13	Insulation: Class Iype to Mount Motor Half.
	14	Temp. Rise ^O C above ^O C By Painting: Mfr. Std. Spec. No.
	15	NEMA DesignType: OHor. OVert. Shipment: ODomestic OExcort
	16	Starting: OExport Boxing Required
	17	DFull Voltage OREGUCED Voltage XOOutdoor Storage for Over Three (3) Months
	18	Doaded Unioaded Data
	19	Dividuale up % Prame No.: Full Load RPM
	20	Direct connected Ofear European Expression 274172
	21	Outher Offer Offer Offer Offer
	23	Enclosure: Locked Rotor Power Factor
	2.4	OOpen Drin Proof Oweather Protected
	2 5	Diotally Enclosed Fan Cooled lock_Rtr. W/Stnd Time (Hot Start) Sec.
	26	O0ther Torques: Full Load
	27	Driven Equipment Inertia Locked Rotor
	28	Bearing Type Lub. Pull-Up
	2 9	No. Starts: /hr,Motor @ Amb.Temp0 Breakdown
	30	No. Starts:/hr,Motor @ Rated Temp. C Acceleration Time (Motor & Load @ 100%/80%
	31	Rotation Facing Coupling End: Rated Voltage)S e c/Sec.
	32	Vibration: ONEMA Std. O Rotor WK2
	33	Noise ONEMA Std. 0 Allow. WK ² @ Motor Shart
	34	Space Heaters: Ores ONO Min. Allowable Acceleration Volt.
	30	Volts rhase heriz livit End Float to
	30	Machine Sheath Temp.
	20	OBacenlatas Osolenlates Max Stator Winding Temm · Of Alarm
	39	OMfr. Std. Fans ONon-Sparking Fans
	40	Winding Temp. Detector: No./Phase Thrust Brg.(ap.(Vert.): Up Down
	4.1	Type Dimensions:
	4 2	Selector Switch & Ind. By: OPurch. OMIT. Length Width Height
	4 3	Winding Temp. Detector & Space Htr. Leads: Weight: .
	44	OSeparate Conduit Box OSame Conduit Box Net Rotor Shipping
	4 5	Specifications: No <u>te</u> :
	4 6	
	47	
	48	
	49	
	50	Note: O Indicates information to be completed by Purchaser; D 3y Manufacturer
		6 MANAGING OFFICE ENGINEERING OFFICE
		5
		2 4 EQUIPMENT DATA SHEET
		CLIENT
	[
	L	1 ITEM
	F	0 ITEM NO. SPEC. NO.
	1	INO DATE ENGINEER CHKVREVIEW APPROVE REO'N. NO. PAGE OF

KELLOGG The M. W. Kellogg Comp	DANY JOB NO ITEN NO
8	REQ. NO.
	DATE:REV. NO
CAS ENGINE DATA	
	ACCESSORIES
RATED HPRATED RPM	AIR INLET FILTER DRY []OIL BATH []MFR. STD.
	MANUFACTURERMODEL
IJSEPARATE	EXHAUSI SILENCER MODEL
IO. POWER CYLINDERS	
FUEL RATE BTU / BHP HR. # RATED LOAD	EXHAUST MANIFOLD WATERCOOLED INSULATED
	TACHOMETER
POWER CYLINDERS WET DRY	ELECT. ENCL. MECH. VIBR. REED
	TACHOMETER RANGE
TURBOCHARGED	FLYWHEEL TURNING BAR AND STAND
LOW FIRE HAZARD IGNATION	MANUAL AIR JACK AIRPSIC
GOVERNOR	UBE OIL FILTER FULL FLOW SLIPSTREAM
RESET BY: PNEUMATIC SIGNAL	
EL ECTRONIC SIGNAL	UEL GAS SURGE DRUM BY: PURCHASER FNG MFR
MANUAL	RECOMMENDED VOLUME FT 3
PEED RANGE: RPM MAX, MIN.	TARTING AIR COMPR. BY: PURCHASER ENG MER
IGNAL RANGE: RPM MAX. MIN.	DRIVER: MOT DR GAS GASOLINE
DN SIGNAL FAILURE VALVE TO: OPEN CLOSE	NO. REQUIRED CAPACITY CFM
SOVERNOR MFR. TYPE:	MANUFACTURER YODEL
REGULATION: NEMA CLASS	AUTOMATIC START , STOP CONTROL
	STARTING AIR RECEIVER BY: PURCHASER BONG MFR.
WEIGHTS AND DIMENSIONS	NO. RECEIVER HORIZ. VERT.
	RECEIVER CAPACITY FT PRESSURE PSIC
NET WEIGHT:	NO. CONSECUTIVE STARTSSTARTS / HR.
MAX, ERECTION WEIGHT MAX, MAINT, WIL	CO, FI: /START
HEIGHT	
ADDITIONAL DISTANCE TO REMOVE POWER RODS	
AUVILLARY 64	
AUXILIART STS	
ENGINE MANUFACTURER SHALL FURNISH POWER CYLINDE	R COULING WATER PIPING FROM A SINGLE INLET FLANGE TO
A SINGLE DISCHARGE FLANGE.	
CIDCUL ATING WATER DUMP DOLVEN BY ENGINE SHAFT	
CINCOLATING WATER FOMF DRIVEN OF ENGINE SHAFT.	
REMARKS:	
REHARKS:	5 6 7 8 9 10 11 12
REMARKS:	5 6 7 8 9 10 11 12 5 6 7 8 9 10 11 12
REMARKS:	5 6 7 8 9 10 11 12

GENERAL PURPOSE STEAM TURBINE SPECIFICATION

_	U Potential Max. Horsepo	wer at Normal Steam Cond	litions							
3	U Outline Dwg. No.	Sectional Dwg. I	NO.							
4	Turbine Mounting	U Ver	tical			Horizontal				
5	Support		terine			Foot				
6	Case Split		dial		C	Axiai				
7	No. of Stages	🗆 wh	eel Diam.							
8	Single Stage	<u> </u>	tow 🗌 3 F	ow		Reentry				
9	Interstage Gland Seal	U Ca	rbon		O Labyrinth					
10	End Gland Seal	U Ca	rbon			Labyrinth				
11	□ No. of Rings/Box				-					
12	Bearing Type	U Ra	Idiai		L	Thrust				
13	C rpm Max. Cont.	Trip								
14	u rpm Max. Allowable				1st Criti	cai	~	_		
15	Case Max. Allow. Works	ng Press. inlet			psig Exh.	aust		<u>م</u>		
1€	Max, Intêt Temp.	"F Max.Exhaus	it Temp,		*					
17	Min. Allowable Exhaust Pre	issure			osig					
18	C R.V. Set Press. Exh.				05+9	Capacity				
19	Hydrotest Press.	iniet			psig	Exhaust				
20	Lube System O R	ing Oil Ο Ρι	rchasers			O Other				
21	U Corrosion Resistant Dra	in Lines								
22	Oil Required		MA TERLE			gpm				
23			MATERIAL	.						
24	High Press. Casing			Exhaust Casi	ng					
25			C Blades							
26				Wheels						
27	Under Packing			-						
28	Trip Valve Body		<u> </u>	Gov. Valve 1	nm					
29		c	CONNECTIONS							
30		SIZE	ANSI	RATING	FA	POSITION				
31	Inlet		_		-		_			
32	Exhaust									
33	Drain		_							
34	Cooling Water									
			oments on no.	zies on separa	te sheet.					
35	C Manufacturer to give al	lowable piping forces & mo								
35 36	Manuracturer to give a	lowable piping forces & mo	TESTS	-		-				
35 36 37	Manufacturer to give a	RE	TESTS				REQ'E	wi		
35 36 37 38	Hydrostatic	Re	TESTS	Aux. Eq	uipment	-	REG.C	wi (
35 36 37 38 39	Hydrostatic No Load Run	RE	TESTS 40'D. WITN 80 0 80 0	Aux. Eq Disassen	nuipment nbly After Test			(
35 36 37 38 39 40	Hydrostatic No Load Run Performance	owable piping forces & m	TESTS 20'D. WITN 38 0 38 0 0 0	Aux. Ec Disassen Test Dat	nuipment nbly After Test			() (
33 36 37 38 39 40 41	Hydrostatic Hydrostatic No Load Run Performance Gov. Response to Contr. S	ignal	TESTS (C) WITN (C) (C) (C) (C) (C) (C) (C) (C) (C) (C)	Aux. Ec Disassen Test Dat Inspecti	nuipment nbly After Test ta Sheets onO Su	rface	REQ'E U O O Parts	(
35 36 37 38 39 40 41 42	Hydrostatic No Load Run Perfyrmance Gov. Response to Contr. S	iawabie piping torces & ma RE	TESTS Q'D. WITN X O X O O O O O O O O O O O O O O	Aux. Ec Disassen Test Dat Inspecti	nbly After Test a Sheets on 0 Su	rface	REQ'C U O O Parts	() (
35 36 37 38 39 40 41 42 43	Manufacturer to give an Hydrostatic No Load Run Performance Gov. Researce to Contr. S	iawabie Diging Torces & ma	TESTS (Q'D. WITN) (Q) (Q) (Q) (Q) (Q) (Q) (Q) (Q	Aux. Ec Disassen Test Dat Inspecti	nbiy After Test a Sheets on 0 Su	rface	REQ'E U O O Parts	() ()		
35 36 37 38 39 40 41 42 43 44	Manufacturer to give a Hydrostatic No Load Run Performance Gov. Response to Contr. S Turbine	ignal	TESTS (Q'D. WITN) (Q) (Q) (Q) (Q) (Q) (Q) (Q) (Q	Aux. Ec Disassen Test Dat Inspecti TS Mounted	uipment nbly After Test ia Sheets on O Su	rface Ib.	REQ'E U O Parts	(
35 36 37 38 39 40 41 42 43 44 45	Manufacturer to give a Hydrostatic No Load Run Performance Gov. Response to Contr. S Turbine Turbine	awabie Diging Torces & m RE ignal Ib Cogler AP	TESTS 20'D. WITN. 30 0 00 0 WEIGH Ib PLICABLE SI	Aux. Ec Disassen Test Dat Inspecti TS Mounted ECLFICATIO	iuipment hbly After Test ia Sheets on Su NS	rface ID.	REQ'E U O Parts			
35 36 37 38 39 40 41 42 43 44 45 46	Manufacturer to give a Mydrostatic Mydrostatic No Load Run Performance Gov. Response to Contr. S Turbine API 611 Gen, Purpose 1	Ignal ID Cogler AP Steam Turbines	TESTS 20'D. WITN. 30 0 00 0 WEIGH IB 0 PLICABLE SI	Aux. Ec Disassen Test Dat Inspecti TS Mounted ECIFICATIO	Iulipment hobly After Test ia Sheets on Su NS R-1 Painting	rface Ib.	REQ'E	() ()		
35 36 37 38 39 40 41 42 43 44 45 46 47	Manufacturer to give a Hydrostatic No Load Run Performance Gov. Response to Contr. S Turpine O API 611 Gen. Purpose O	Ignal ID Cogler APP Steam Turbines GN-1 Noise	TESTS 2010. WITN 2000	Aux. Ec Disassen Test Dat Inspecti TS Mounted FCIFICATION	Iuipment hbly After Test ia Sheets on O Su NS R-1 Painting	rface Ib.	REG'E	() ()		
33 36 37 38 39 40 41 42 43 44 45 46 47 48	Manufacturer to give a Mydrostatic No Load Run Performance Gov. Response to Contr. 5 Turpine O API 611 Gen. Purpose 1 O	Ignel ID Cooler Construction Cooler C	TESTS 59'D. WITN. 80 0 00 0 WEIGH 15 2 PLICABLE SF	Aux. Ec Disassen Test Dai Inspecti TS Mounted ECLFICATIO	Nument holy After Test a Sneets on Su NS R-1 Painting	iface Ib.	PEQ'E U O Parts	wir ((
33 36 37 38 39 40 41 42 43 44 45 46 45 46 47 48 49	Manufacturer to give a Hydrostatic No Load Run Performance Gov. Response to Contr. S Turbine API 611 Gen. Purpose C REMARKS:	In the second se	TESTS CONTINUE	Aux. Ec Disassen Test Dai Inspecti TS Mounteg FECIFICATIO	IUIDMENT holy After Test la Sheets on O Su NS R-1 Painting	rface ib.	PEG'E	wi ((
33 36 37 38 39 40 41 41 42 43 44 45 46 47 48 49 50	Manufacturer to give a Mydrostatic No Load Run Performance Gov. Response to Contr. S Turbine O API 611 Gen. Purpose O REMARKS:	RE ignal ib Cooler AP Steam Turbines GN-1 Noise	TESTS GOD. WITN SOD SOD O O O O O O O O O O O O O	Aux. Ec Disassen Test Dai Inspecti TS Mounteg ECIFICATION	IUIDMENT hoby After Test is Sheets on O Su NS R-1 Painting	rface ID.	REGIE	() ()		
33 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51	Manufacturer to give a Mydrostatic No Load Run Performance Gov. Response to Contr. S Turbine O API 611 Gen. Purpose 1 O REMARKS:	ID Cogler Covering the District of a monomous of the Cogler ID Cogler API Steam Turbines - GN-1 Noise	TESTS Q.D. WITN SOU SOU WEIGH ID C PLICABLE SI	Aux. Ec Disassen Test Dai Inspecti TS Mounted ECIFICATION	Iulpment hbly After Test ia Sneets on O Su NS R-1 Painting	rface Ib.	REQ'E U O Parts	wir ((
33 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 51 2 52	Manufacturer to give a Hydrostatic No Load Run Performance Gov. Response to Contr. S Turbine API 611 Gen. Purpose REMARKS:	In Cogler Diping forces & m, Rf	TESTS GOD. WITN 8 O 0 O 0 O 0 O WEIGH ID D PLICABLE SI	Aux. Ec Disassen Test Dat Inspection TS Mounted ECIFICATION	iuipment nbiy After Test ia Sneets on O Su NS R-1 Painting	rface Ib.	REQ'E	wi ¹		
33 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 52 53	Manufacturer to give a Hydrostatic No Load Run Performance Gov. Researce to Contr. 5 Turbine O API 611 Gen. Purpose 1 REMARKS:	ID Cooler Coo	TESTS EQ'D. WITN SO O O O WEIGH Ib C PLICABLE SI	Aux. Ec Disassen Test Dat Insoectiv TS Mounted ECCIFICATIO CO	unioment noty After Test a Sneets on O Su NS R-1 Painting	rface ID.	REQ'E			
33 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 52 53 54	Manufacturer to give a Mydrostatic No Load Run Performance Gov. Response to Contr. S Turbine O API 611 Gen. Purpose 1 O REMARKS:	Ignal	TESTS 120'D. WITN 80 O 00 O 00 O WEIGH 15 F PLICABLE SI	Aux. Ec Disassen Test Da Inspecti TS Mounted Mounted CECIFICATION	uidment holy After Test ta Sheets on O Su NS R-1 Painting	rface ib.	REQ'E			
33 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 52 53 54 55	Manufacturer to give a Mydrostatic No Load Run Performance Gov. Response to Contr. S Turbine API 611 Gen. Purpose REMARKS:	Ignal ID Cogier AP Staam Turbines GN-1 Norse	TESTS GOD. WITN SOLO COD COD WEIGH IB 5 PLICABLE SI	Aux, Ec Disassen Test Dat Inspecti TS Mounted ECIFICATIO	INIDMENT	rface Ib.	REQ'C O O Parts			

0 RM NO 2868 4 56 MP&S		SOUTHWES 4800 SANTA	TERN ENGINEERING COMPANY	SHEETOF.
		SWECO	GEAR DRIVE DATA SHEET	R E V
		CUSTOMER	EQUIP. NO.	
	z	PLANT LOCATION	FILE NO.	
	3	SERVICE		
	4	MANUFACTURER	TYPE SIZE	
	5	OPERATION	BEARINGS	
	6	CONTINUOUS OR INTERMITTENT	H.S. SHAFT - OUTBOARD	
	7	GEAR RATING - BHP	H.S. SHAFT - INBOARD	
	8	THERMAL RATING - HP	INTERMEDIATE SHAFT - OUTBOARD	
	9	SERVICE RATING - HP	INTERMEDIATE SHAFT - INBOARD	
	10	MECHANICAL HP RATING	L.S. SHAFT - OUTBOARD	
	11	AGMA SERVICE FACTOR	L.S. SHAFT - INBOARD	
	12	AGMA CLASS GEARS		
	13	GEAR RATIO	COUPLINGS	
	14	EFFICIENCY @ BHP RATING	H.S. SHAFT COUPLING - TYPE	
	15	NO. OF REDUCTIONS OR INCREASES	SIZE	
	16	HIGH SPEED SHAFT RPM	MFG.	
	17	LOW SPEED SHAFT RPM	L.S. SHAFT COUPLING - TYPE	
	18	ACTUAL DRIVEN MACHINE RPM	SIZE	
	19	ROTATION H.S. SHAFT FACING DRIVER	MFG.	
	20	ROTATION L.S. SHAFT FACING DRIVER	COUPLING GUARDS	
	21			
	22	CONSTRUCTION & MATERIAL	TESTING	
	23	CASE TYPE	WITNESS PERFORMANCE TEST	
-	24	CASE MATERIAL	INSPECTION	
₩>- #-0Z	25	H. S. PINION OR WORM		
	26	H.S. GEAR	MISCELLANEOUS	
	27	H.S. SHAFT	WEIGHT	
TTT	28	INTERMEDIATE PINION OR WORM	OUTLINE DWG. NO.	
	29	INTERMEDIATE GEAR	CROSE EECTION DWG. NO.	
°	30	INTERMEDIATE SHAFT	SERIAL NO.	
	31	I S PINION OF WORM		
	32	L. S. CEAR	PRIME MOVER	
	33	L.S. SHAFT	TYPE - MOTOR, TURGINE, OTHER	
	34	H.S. SHAFT SINGLE OR DB'L. EXT.	RATED HP	
a a a o > w o	35	LS SHAFT SINGLE OR DB'L EXT.	DRIVER RPM /FULL LOAD RPM	/
	36	H.S. SHAFT B.D. AT CLPS.	ROTATION FACING END OFF. CLPG.	
	37	L.G. GHAFT O.D. AT GEPS	MANUFACTURER	
	38	CASE GASKETS	TIPE /allE	/
DATE	39	CASE BOLTS	WEIGHT	
	40	BEDFLATE	NOM. SHAFT O.D. AT CLPG.	
BY	41		DRAWING NO.	
	42			
HECKED	43	LUBRIGATION	BRIVEN MACHINERY	
	44	TYPE LUBRICATION - GEARS	MACHINE DRIVEN	
NGINEER	45	TTPE LUBRICATION - BEARINGS	BHP REQUIRED	
	46	OIL PUMP TYPE	REQUIRED DRIVEN RPM	
NUJ. ENG.	47	OIL PUMP DRIVE	ROTATION FACING SHAFT END	
	48	VISIBLE LUBRICATOR SIGHT GLASS	MANUFACTURER	
	49	OIL FILTER OR STRAINER	TYPE /5IZE	/
	50	OIL COOLER REQUIRED	WEIGHT	
	51	TYE OIL COOLER	NOM. SHAFT O.D. AT CLPG.	
	52	COOLING WATER @ "F & GPM	DRAWING NO.	
	53	SIZE COOLING WATER CONN.		
	-	REMARKS:		
	20		*	1

MANUFACTURERS ASSOCIATION STANDARDSOFTUBULAREXCHANGER

EXCHANGER SPECIFICATION SHEET

CUSTOMER		JOB NO. REFERENCE NO
ADDRESS		INQUIRY NO
PLANT LOCATION		DATE
SERVICE OF UNIT		ITEM NO.
SIZE	TYPE	CONNECTED IN
GROSS SURFACE PER UNIT	SHELLS PER UNIT	SURFACE PER SHELL
	PERFORMANCE OF ONE UNIT	
	SHELL SIDE	TUBE SIDE
FLUID CIRCULATED		
TOTAL FLUID ENTERING		
VAPOR		
LIQUID		
STEAM.	I	
NON-CONDENSABLES		
FLUID VAPORIZED OR CONDENSED		
STEAM CONDENSED		
GRAVITY-LIQUID		· · · · · · · · · · · · · · · · · · ·
VISCOSITY-LIQUID		
MOLECULAR WEIGHT-VAPORS		
SPECIFIC HEAT-LIQUIDS	BTU/LB/°F	BTU/LB/*F
LATENT HEAT-VAPORS	BTU/LB/°F	BTU/LB/*F
TEMPERATURE IN	• F	· · · · · · · · · · · · · · · · · · ·
TEMPERATURE OUT	· F	• F
OPERATING PRESSURE	PSI	PSI
NUMBER OF PASSES PER SHELL		
VELOCITY	FT./SEC.	FT./SEC.
PRESSURE DROP	PSI	PSI
FOULING RESISTANCE		
HEAT EXCHANGED-BTU/HR.	M.T.D. (Co	xrrected)
TRANSFER RATE-SERVICE	EFF. SURFACE PER UNIT	- · · · · · · · · · · · · · · · · · · ·
	CONSTRUCTION	
DESIGN PRESSURE	PSI	PSI
TEST PRESSURE	PSI	PSI

36 DESIGN TEMPERATURE ۰F F 37 TUBES O.D. NO. BWG. LENGTH PITCH 38 SHELL I.D. O.D. SHELL COVER 39 FLOATING HEAD COVER 40 CHANNEL CHANNEL COVER TUBE SHEETS-STATIONARY FLOATING 42 BAFFLES_CROSS TYPE 43 BAFFLE-LONG TYPE TUBE SUPPORTS GASKETS 45 CONNECTIONS-SHELL-IN OUT RATING PSI CHANNEL-IN OUT RATING P81 48 CORROSION ALLOWANCE-SHELL SIDE TUBE SIDE 49 CODE REQUIREMENTS TEMA CLASS 50 WEIGHTS-EACH SHELL BUNDLE FULL OF WATER NOTE: INDICATE AFTER EACH PART WHETHER STRESS RELIEVED (S. R.) AND WHETHER RADIOGRAPHED (X-R) 51 52 REMARKS:

46

DD FORM 2000 11-	63				
AIF	COOLED	LOCATION	IOB NO	FILE	NO.
ΕX	CHANGER	DEDADED DV			
Nothern S.D.E.C.				DATE	
	SHEET	APPROVED BY		DATE	
Company	011221	AINDICATES REV	ISION	LIVE TO LEASE	
ERVICE OF UNIT				NO. OF B	AYS
IZE	ТҮРЕ				
URFACE ITEMEXT	ERNAL	8	ARE TUBE		SQ F
EAT EXCHANGEDB	TU/HR		FFECTIVE MTD		•
RANSFER RATEEXt	SURF.	В	ARE TUBE SURF,	<u>8</u>	TU/HR SQ FT*
		PERFORMAN	CE DATA		
		TUBE S	IDE		
LUID CIRCULATED				TEMPERATURE IN	°.
TOTAL FLUID ENTERI	NG		LBS/HR	TEMPERATURE OUT	°I
VAPOR				INLET PRESSURE	PSIC
LIQUID				GRAVITY-LIQUID	
STEAM				VISCOSITY CPSEF	
NON-CONDENSABLE	<u></u>			VISCOSITY CPS@*F	
VAPOR CONDENSED				MOLECULAR WEIGHT	
STEAM CONDENSES	2			SPECIFIC HEAT	BTU/LB°
DENSITY: VAPOR-LIG	UID MIXTURE		LBS/CU FT	LATENT HEAT	BTU/LI
CONDUCTIVITY		BT	U FT/HR SQ FT°F	ALLOW, PRESS, DROP	PS
FOULING RESISTANCE	E 1.5.		HR SQ FT *F/BTU	DESIGN PRESS, DROP	PS
		AIR S	IDE		
AIR OTY	LBS/HR SCEM	TEMP IN	°F	ALTITUDE	
AIR GTY/FAN	ACFM	TEMP OUT	°F		
		CONSTRU	CTION		
DESIGN PRESSURE	P51	TEST PRESSURE	PSI	DESIGN TEMP	°F
SECTI	ON	HEAD	ER	TUBE	
SIZE	ROWS	ТҮРЕ		MATERIAL	SEAMLESS
NO./BAY		MATERIAL			WELDED
ARKANGLMENT		NU. PASSES		OD IN BWG	AVG MIN WALI
SECTIONS IN PARA	ALLEL FF	MATERIAL		NO./SECTION	
	L3			ENGIH	
BAYS IN PARALLEI	_	GASKET MATERIAL			NAUV
IES		CORRUSION ALLOWAN	ICE IN.	FIN	
MISC		SIZE INCET NOZZI P	IN.	MATERIAL	
C TRUCTURE		SIZE OUTLET NOZZLI	IN.		
STRUCTURE				NU./IN.	
				TTPE	
		MECHANICAL	FOLIPMENT		
FAN		DRIV	FR	SPEED RED	UCER
TYPE		MER		MER	
NO /BAY		TYPE		TYPE	
TIP SPEED	ET/MIN	NO. / BAY		NO /BAY	
DIAM ET R	PM	HP/DRIVER	DDM	MODEL	
HP/FAN	TOU ADJUST	ENCLOSURE	ISP W	AGMA HP RATING	
NO BLADES				RATIO	,
RIADE MATI	LIG01 9. VA8				/
NOTES					
<u>, , , , , , , , , , , , , , , , , , , </u>					

DIMENSIONS: LENGTH WIDTH REPRODUCTION OF STANDARD NO. 53.0 1-3

WIDTH

HEIGHT

SPECIFICATIONS FOR FIN FAN HEAT EXCHANGERS FLUOR • GRISCOM RUSSELL

Address Date Plant	Custo	mer			Proposal No.	
Fine Fin Fan Model Service INSIDE TUBES Lass of Fluid Altitude Specific Gravity ● 60 F Data Dansity ● An. Conditions #/Cu.Ft. Molecular workgint Data Henry Conductivity Centipoises at Fars: Model Viscosity Centipoises at Diameter Quantity—Liquid Centipoises at Diameter Vapor East Model #/Hr. N. C. Ges #/Hr. Honeyower per Fan Fild Vap. or Condud. #/Hr. Make Operating Temperature In "F. : Out "F. Drive Shaft: Operating Temperature #/Cl" Make Test Pessure Design Temperature #/Cl" Number Number Design Temperature Staft Pessure #/Cl" Number Design Temperature #/Cl" Number	Addre	55			Date	
Fin Fen Model Service Service Class of Fund Specific Gravity @ 60 F Specific Gravity @ 60 F Specific Gravity @ 60 F Data Frequency Barting @ An. Conditions #//Cu.K. Steam Conductivity Conductivity Conductivity Conductivity Conductivity Context in Frequency Contropises at Vapor of Gas Countity-Liquid Contropises at Vapor of Gas Contropises at Vapor of Gas Contropises at Vapor of Cas Contropises at Vapor of Cas Cas Vapor of Cas Cas Pressure Pressure Cas Design Pressure In Steam Condemed Design Pressur	Plant					
Service INSIDE TUBES AIR Class of Fluid INSIDE TUBES AIR Specific Gravity @ 60 F Intrude Fee Specific Gravity @ 60 F Intrude Fee Specific Gravity @ 60 F Intrude Intrude Specific Gravity @ 60 F Intrude Intrude Specific Gravity @ 60 F Intrude Intrude Dansity @ An. Conditions #//Cu.Ft. Static Pressure Conductivity Fans: Model MICHANICAL RQUIPMENT Conductivity Centipolies at F. Viscolity Centipolies at F. Vapor Ges Cu.P.124 Mg. 60/F Vapor of Ges Cu.P.124 Mg. 60/F Pressure Vapor of Ges Cu.P.124 Mg. 60/F Press Steam #/Hr. Make Steam #/Hr. Make Steam Condensed #/Hr. Make Operating Pressure #/I/1" Motors: Type Design Temperature In "F. Out Design Temperature In T. Nunber	Fin Fa	n Model				
Service INSIDE TUBES AIR DATA Class of Fluid Altitude Fee Specific Gravity @ 60 F Instant Ar Temp. Air Specific Fluid @ An Temp. B.T.U./# Outlet Air Temp. Density @ An. Conditions #/Cu.Ft. Static Pressure Molecular weight B.T.U./# Outlet Air Temp. Conductivity Entiple and Conductivity Fars: Model Viscosity Centipoises at F. Viscosity Centipoises at F. Viscosity Centipoises at F. Vapor or Ges Cu.EtC4 int @ 14/1 & Aus. & 60/5 P.P.II Vapor #/Hr. Honseldes Steam #/Hr. Honseldes Fluid Vap. or Conduct #/Hr. Honseldes Operating Temperature In *F. Operating Temperature In *F. Pressure Drop #/CI* Motors: Type Design Temperature *F. H.P. Inst Pressure #/CI* Surface per Section Sorfa Surface Surface per Section Sorfa Surface Moteral Members Indeel H.P. Fluid Members Steam In. #/CI* Materades						
Service INSIDE TUBES AIR: DATA Class of Fund Altitude Fee Specific Gravity @ 60 F Not Famp. Air Latent Heal @ An Temp. B.T.U/# Outlet Air Temp. Fr Density @ An. Conditions #/Cu.Ft. Static Pressure */HC Moticular Weight Density Fams: Model */HC Conductivity Fams: Model */HC */HC Viscosity Centipoises at */F. Number Vapor or Ges Cu.FL/24 Hr @ 14,7 Abs. & 60*F. No. or Blades Vapor or Ges Cu.FL/24 Hr @ 14,7 Abs. & 60*F. No. or Blades Vapor or Ges Cu.FL/24 Hr @ 14,7 Abs. & 60*F. No. or Blades Vapor or Ges Cu.FL/24 Hr @ 14,7 Abs. & 60*F. No. or Blades Vapor or Ges Cu.FL/24 Hr @ 14,7 Abs. & 60*F. No. or Blades Vapor or Ges Cu.FL/24 Hr @ 14,7 Abs. & 60*F. No. or Blades Vapor or Ges # //Hr. Hersonward # //Hr. Hersonward # //Hr. N.C. Ces # //Hr. Make Static Pressure Field Vep. or Conded. # //Hr. Make Static Pressure Design Temperature In *F. Out Motors: Type Design Temperature *F. H.P. <td< td=""><td></td><td></td><td>_</td><td></td><td></td><td></td></td<>			_			
INSIDE TUBES AIR DATA Class of Fund Alitude Fee Specific Gravity @ 60 F Instant Air Temp Fe Specific Hest @ Am, Temp. Air BT, U/# Latert Heat @ Am, Temp. B, IU/# Static Pressure Density @ Am. Conditions #/Cu. Ft. Static Pressure Onductivity Fans: Model Mumber Conductivity Centipolies at F. Vapor of Gas Cu. Ft./24 Hr. @ 14,7.3. Abs. 6:00'F # P. M. Vapor of Gas Cu. Ft./24 Hr. @ 14,7.3. Abs. 6:00'F # P. M. Vapor of Gas Cu. Ft./24 Hr. @ 14,7.3. Abs. 6:00'F # P. M. Vapor of Gas Cu. Ft./24 Hr. @ 14,7.3. Abs. 6:00'F # P. M. Operating Temperature In * F. Operating Temperature Operating Temperature In * F. Outors: Type Design Temperature # //I' Motors: Type Design Temperature # //I' Mumber Cust Surface per Section Sq. Ft. H.P. Surface per Section Sq. Ft. MAKe Task Members In * F. H.P. Tube Plugs AND CONSTRUCTION Make Make Tube Plugs Gas Engines: Model Make	Servic	e				
Class of Floid Altitude Fee Specific Gravity @ 60 F Inder Air Temp. Fee Density @ Ava. Conditions B.T.U./# Outler Air Temp. # Density @ Ava. Conditions #/Cu. Ft. Static Pressure *H/Ft. Molecular weight MECHANICAL Equipment ** ** Conductivity Centipolies at *. Number ** Viscosity Centipolies at *. Number ** ** Vapor or Ges Curl FL/24 Hr. @ 14,7 & Ata & 60/F # # # # ** ** ** Vapor or Ges Curl FL/24 Hr. @ 14,7 & Ata & 60/F # # # # ** * <t< td=""><td></td><td></td><td>INSIDE TUBES</td><td></td><td>AIR DATA</td><td>Ą</td></t<>			INSIDE TUBES		AIR DATA	Ą
Specific Gravity @ 60 F Inst Air Temp. Air #/Hr. Specific Heat @ Are Temp. Air Air #/Hr. Lahert Heat @ Are Temp. B. T. U/# Air #/Hr. Density @ Are. Conditions #/Cu. Ft. Static Presure */Hr. Molecular weight Centipoises at F. Number Conductivity Centipoises at F. Number Viscosity Centipoises at F. Number Quantity-Liquid Centipoises at F. Number Vapor or Ges Cu. FL/24 Hr. @ H/1 & Als. & 6/F # Arr. N. C. Ges # //Hr. Horsepower per Fan Steam # //Hr. Kats. & 6/F # are at Operating Temperature In * F. Outre Static: Operatic: Operating Temperature In * F. Outre Static: Operatic: Operatic	Class	of Fluid			Altitude	Feet
Specific Heat @ An, Temp. Air #/Hr. Latent Heat @ An, Conditions #/Cu, Ft. Static Pressure *H/Er. Molecular weight #/Cu, Ft. Static Pressure *H/E Molecular weight Fans: Model #ECHANICAL EQUIPMENT *H/E Conductivity Centipoises at * Diameter *H/E Quantity—Liquid Centipoises at * Diameter * Quantity—Liquid Cu_E1/24 Hr. @ H(7 # Abs. 5 & 0/F #P ur * * Vapor #/Hr. No. C Blades * * * * Vapor #/Hr. No. C Blades *	Specif	ic Gravity 🤁 60 F			Inlet Air Temp.	'F.
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Density @ An. Conditions #/Cu.Ft. Static Pressure "H_G Molecular weight	Latent	Heat 🝘 An. Temp.		B. T. U./#	Outlet Air Temp.	E.
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Price:	F.O.E.	Ship	ping	Poi	nt,	not	t	ere	cted	•	•		\$					
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											TO	TAL	PRICE	ş				
Estimate	d Erec	tion	Labor											\$				
Shipping	; Weij	ht																lbs.

LBS

SHIP. Wt

FORM NO. 724

THE GRISCOM-RUSSELL CO. G-FIN TANK HEATER SPECIFICATIONS

1	Customer			Item No.	
2	Address			Ing. No.	
3	Plant			Date	
4 1	Unit Required				
8					
6	Service			Type Internal Ext	ernal
7		SHELLS		TUBES	
8 (Class of Fluid				
9	Specific Gravity @ 60°F.				
10	Specific Heat @ Ave. Temp.				
ШĮ	Latent Heat @ Ave. Temp.		B.T.U. /#		8.T.U. /#
12	Viscosity	S.S.U. at	"F.		
13	Viscosity	Centipoises at	"F.	Centipoises at	<u>۴.</u>
14	Viscosity	Centipoises at	°F.	Centipoises at	°F.
15	Quantity	GPM	∦/Hr.		Į/Hr.
16	Operating Temperature	In 'F. : Out	"F.	In "F. : Out	° F .
17	Operating Pressure I		#/□		//¤'
18	No. Passes per Shell	1		2	
19	Design Pressure		∦/□ ″		1/0*
20	Test Pressure		#/□*		∦ /□*
21	Design Temperature (Max. Metal Temp),		"F.		"F.
22	Heat Exchanged	BTU /Hr.	Surface Per Us	nit	Sq. Ft.
23 ं	Corrected M.T.D. Design Rate		Surface Per Sh	nell	Sq. Ft.
24		MATERIALS AND CON	ISTRUCTION		
25	SHELL: Steel	CHANNEL:	Steel	Baffles: Steel Stand	lard Spacing
26	INLET CONN. SIZE	INLET CONN. SIZE:		Tie Rods and Spacers :	Steel
27	Drilling: 150#	Drilling:	150# P.T	Gaskets: Durabla	
28	OUTLET CONN. SIZE	OUTLET CONN. SIZE		Saddles: Steel (Type H	ISU-CSU only)
291	Drilling: 150#	Drilling:	150# P.T.	Structural Codes	
30	ank Manhole Flange Size : 0.D	Tubes-34" IPS Seamless Ste	el	Inspection by:	
31	Bolt Circle No. Dia.	Fins-Steel. Nom. Length:		OVERALL LENGTH	
32	Tube Sheet—Steel with Cone Seat Tuba Join	nts and Look_Nuts	(Balance)	OUTSIDE SHELL DIAMET	ER
33	Support Plats on Open Shell Unit Steel1"	Thick		Reference Print	-
34	GUARANTEE: This unit will give full capac	ity withtank liquid level 2 Ft. or	more above un	t.	

SUGGESTIONS:

1. steam trace all oil lines

2. Use steam trap large enough to handle indicated steam condensate

3. Vent steam side of unit.

4. Provide drains for steam side to take care of shut down during cold weather.

5. This is an instantaneous heater; steam required only when pumping oil.

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			DATA	SHEE	T	REQ NO					
	/					• 8663	۹ <u> </u>	OF			
				INTER	COOLERS AND	AFTERCOOLE	RS				
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BETWEEN STAGE	NUMBER										
OLER FURNISH	ED: Y										
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COOLEMNODEL	OW TYPE										
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	TUBES OF	R INNER PIP	۴								
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	TEMPERA	TURE OUT	•F								
	PRESSUR	E DROP PS									
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					MOISTURE SI	PARATORS					
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	TYPE IN										
	SHELL M	ATERIAL									
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	TO BE FU	IRI ISHED B		ESSOR M	ANUFACTURE		PURCHA	SER			
					AIR REC	EIVER					
FURNISHED BY: C	COMPRESS	OR MANU	FACTURER		PURCHASER						
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OPERATING	PRESS	URE	PSIG											
NUMBER OF	PASSÉS		FT 1050									_		
PRESSURF				1	<u> </u>		+					_		
FOULING RE	SISTANC	E (MIN	1)				1					+		
HEAT EXCH	ANGED	BTU/HR	1				1	MTD	(COR	RECTED].	OF . THE	ORETIC	AL	
TRANSFER	RATE·	SERVICE			•	CLEA	N							
					CONS	STRUCTIO	N							
DESIGN PR	ESSURE		P S I						I]
TEST PRES	SURE		PSI										-	
UC 816 N	CHPERAI	UKE	~F											
MATERIALS	OF CO	NSTRUCT	ION	RESS	J R E P L A T E				ONN	ECTINGP	LATE			
ENDSP	LATES	TING		THER	MALPLATE:	S NG			FRA	SIDE CA				
PROTECTION	SHRO	UD		PAIN					0100	SIDE GA	SKET			
00000505														
CONNECTIO	INS HO	I SIDE			1 N									SERIES SERIES
FABRICATI	ON SPEC	aidt												
INSPECTION	N& TEST								TEST	PRESSURE	PSIG			
WEIGHT L	BS EMP	ΓY		FULLO	FWATER				OPER	ATING				
REMARKS									-					
nements														

ENG EQUIP 705-1 (2-76)



STEAM SURFACE CONDENSER DATA SHEET

STOMER	EST. NO	
CATION	JOG NO.	
	PAGE NO	•
RVICE OF UNIT	ITEM NO.	
	DATE	
Condenser		
Manufacturer		
a. Name:		
b. Location:		
Condenser surface (total effective) :		4 ft
Number of water passes:		
Design basis operating conditions and performance		
a Nature and source of circulating water:		
b. Duty (net heat rejected to circulating water) :		
		Btu per hr
c. Circulating water quantity:		gpm
d. Circulating water inlet temperature:		F
e. Cleanliness factor:		%
f. Average circulating water velocity in tubes:		fp
g. Absolute pressure (to be measured at point specified in h Deacration — maximum free oxygen in condensate	the Heat Exchange Inst	itute Standards) : in. H _i
(42) (14) (7) — (Indicate) :		ppl
i. Circulating water friction loss through clean tubes and	waterboxes:	f
i. Circulating water friction loss through clean tubes and J. Supplementary (expected) performance on curve sheet	waterboxes:	1
 i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes 	waterboxes: s:	f
 i. Circulating water friction loss through clean tubes and J. Supplementary (expected) performance on curve sheet Tubes a. Main Condensing Sections: 	waterboxes: 8:	1
 (ii) (i) (i) (i) (i) (i) (ii) (iii) (ii	waterboxes:	ſ
 i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: 1) No Size: in. 	waterboxes:	f
i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: 1 1) N o . : S i z e : i n . I2 Length, effective:	waterboxes:	f
i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: I l) N o . : S i z e : i n . I2 Length, effective: Ov (3) M a t e r i a 1:	waterboxes:	f
 (1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	BW(
 (1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	BW(
 (1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	BW(
 (1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	BW(
 (1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	f BW(BW(
 (1) (1) (0) (1) (1) (1) (1) (1) (1) (1) (1) (1) (1	waterboxes:	f BW(BW(
(1) (waterboxes:	BW(
i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: i 1) N o . : S i z e : - i n . 12 Length, effective:	waterboxes:	BW(
i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: i. 1) N o . : S i z e : - i n . 12 Length, effective:	waterboxes:	f t
i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: I 1) N o . : S i z e : i n . 12 Length, effective:	waterboxes:	f BW(f f BW(f
i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: I 1) N o . : S i z e : - i n . 12 Length, effective: Ov (3) M a t e r i a 1: Ov (3) M a t e r i a 1: Ov (3) M a t e r i a 1: Ov (3) M a t e r i a 1: Ov (3) Material: Ov (2) Length, effective: Ov (3) Material: Ov (1) No.: Size: Ov (3) Material: Ov (1) No.: Size: Ov (3) Material: Ov (3) Material: Ov (3) Material: Ov (4) Method of connection Inlet end: (5) Method by:	waterboxes:	f
i. Circulating water friction loss through clean tubes and j. Suppkmentary (expected) performance on curve sheet Tubes a. Main Condensing Sections: I I) N o . : S i z e : i n . 12 Length, effective:	waterboxes:	f BW(f BW(
(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	BW(BW(BW(BW(BW(
(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	BW(BW(BW(BW(
(1) (1) (r) (r) (r) (r) (r) (r) (r) (r) (r) (r	waterboxes:	BW(
(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	BW(
(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	waterboxes:	f

ENG EQUIP 705-2 (2-76)	
STEAM SURFACE CONDENSER DATA SHEET (con't.) EST. NO	
CUSTOMER JOB NO	STEAM SURFACE CONDENSER DATA SNEET (con't.)
LOCATION PAGE NO	EST. NO.
SERVICE OF UNIT ITEM NU, DATE	LOCATION PAGE NO.
	SERVICE OF UNIT ITEM NO
7. Shell	DATE
a. Material :	B. Stum Jot Air Ejector(s)
b. Thickness:Number of sections:	1. Manufacturer
c. Method of connecting sections:	a. Name:
d. Provision for tube expansion:	b. Location:
e. Number and size of venting connections:	2. Total number of Ejector Units:
	3. Sixeandtype:No. of elements:No. of stages:
	mounted on:
a Hotwell	
a. Type:gais	4. Operating Conditions and Performance
b. Material :	a Design canacity - 100 ner cent -
c. Thickness:Number of sections:	Suction pressure:
	Suction presenter
9. Tube sheets	
a. Number and size: Thickness:	Dry air leakage:SLIM corresponding to :- per
h Material -	Non-condensible gases other than air leakage:()lbe per hr
D. Matchai .	()lbe per h
	()lbs per h
10. Tube support plates	Associated saturated water vapor :
a. Number ;T h i c k n e s s :	Total gas-vapor mixture:
b. Material :	Design capacity each element:per cent
	Number of elements operating for 100% capacity
11. Steam inlat node and anharet connection	Currie (a) of comparison of consistence of the second seco
II. Steam met net and exhaust connection	b. Steam conditions
a. viacual:	Maximum initial pressure raig Total temp
b. Thickness:Number of sections:	Navinum initial pressureperg Fotal temp
c. Connection to shell (flanged) or To turbine (nanged) (r (welded);	Minimum operating pressure (at housing);poing lotal (cmp;
	c. Steam 'Consumption (100 per cent design capacity) :log per h
d. Height top of waterbox hange to turbine exhaust including expansion joint (if any)	5. Condenser Data
	a. Surface sq ft:Inter-condenserAtter-condenser
e. Provision for vertical expansion (condenser hung) (spring supports) (expansion joint).	b. Surface:per cent d&i capacit
f. Spring supports (if used). Type:	c. Tube size:in. OD;BWG;Length (effective-overall)
g. Expansion joint (if used), Type:	d. Method of fastening tubes:
Material:	e. Cooling Water (condensate) (raw water)
Height: Ends — (flanged) or	Minimum flow required at:per cent design capacity:gpu
(welded)	Maximum design flow:
	Friction loss at maximum design flow:
12. Approximate weights, condenser with tubes	f Watayhayaa Maximum daaiga maasuutay
a. Dry:lbs In service:lbs	1. waterboxes Maximum dealgn presoure:
b. Steam space only filled with water: <u>lbs</u>	6. Materials
	a. Suction chambers:
	b. Diffusers
13. Preuminary drawing(s)	c. Steam nozzles:
14. Descriptive bulletin(s) (or specifications) +	d. Steam chests (nozzle heads) :
	e. Condenser shells:

hr

ENG EQUIP 705-4 (2-76)	KELLOGG				SOUTHWEST	ERN ENGINEERI	NG COMPANY	SHEETOF
				SWECO	FIRED I	HEATER SPECIFICATION	N SHEET	RE
STEAM SURFA	ACE CONDENSER DATA SH	EET (con'+.)				· · · · · · · · · · · · · · · · · · ·		
		EST. NO					EQUIP NO	
CUSTOMER		JOG NØ.	— пп	3 SERVICE	ATION			
		PAGE NO.	— 1111	4 MANUFACT	URER	TYPE	SIZE	
SERVICE OF UNIT		DATE	i///	6		THERMAL DESIGN		
6 Water showless.				7 FLUID CIRCULA 8 TOTAL FLUID E	NTED			
i. water chambers:				9		INLET	OUTL	ET
g. Tube Silleets:			— 1111	10 TOTAL VAPOR	#/HR			
h. Tubes:				12 GRAVITY LIQUI	D*API			
7. Fittings and Accessories included				13 DENSITY VAPO	R #/CU.FT.			
a. Air leakage meter:				15 VISCOBITY VAP	0R CP			
b. Other:			[]]]	16 SPECIFIC HEAT	BTU/#			
				17 LATENT HEAT	BTU/#			
			Ibe	19 OPERATING PR	ESSURE PSIG			
8. Approximate weight, each eje	ector unit:		0s	20 MOLECULAR W	EIGHT OF VAPOR			
9. Preliminary drawing(s) :				22				
10. Descriptive bulletin (s) (or specific	ations) +		⊢╄╋╋	23 ALLOWABLE PR	ESSURE DROP PSI	CALCULAT	ED PRESSURE DROP PSI	
			≈ =>-∞-0	24 TYPE OF FLOW		SERIES	PARALLEL	-
C. Hogging Ejectors				20 HEAT ABBORPT	ION BTU/HR.	RADIANT	CONVECT	ION
1. Manufacturer				27 MEAT ABSORPT	ION BTU/HR.	TOTAL		
a. Namr:								
h Location :				30 RADIANI TRAN	5. RATE BTU/SQ. FT./HR.	AVERAGE	MAXIMUM	
2 Steem) (Air) Weter) Operated	Number per condenser:	Size:		32 EFFICIENCY	% - LHV/HHV	AVERAGE	MAXIMUM	
2. Stealif (All) water) Operated. I				33 PLUE GAS TEM	P. (APPROX.) F			
3. Operating Conditions and Perform	nance		<4440>W	35 LHY OF FUEL	BTU/			
a. Capacity:lb p	per hr dry air at	in. Hg abs Suction Pr	ressure	36		· · · · · · · · · · · · · · · · · · ·		
b. Steam consumption:	lb per hr at	psig	F	37 PRESSURE AVAI	LABLE AT BURNER - PSIG	FOR GAS		MAXIMUM
c. Maximum design steam condition	18:-	psig	F DATE	39	· · · · · · · · · · · · · · · · · · ·	FOR STEAM	NORMAL	MAXIMUM
d. Air consumption:		SCFM atps	i g BY	4				
e. Water consumption:		gpm_at	DSIgCHECKED	42		MECHANICAL DESIGN	t	· · · · · · · · · · · · · · · · · · ·
4 Anneximate weight each:		51	lbs	43 NO. OF TUBES	RAD		CONVECTION	
4. Approximate weight each.			ENGINEER	45 RADIANT TUBE	S INCHES O.	D. TUBEWALL I	NCHES MATERIAL	
5 Preliminary drawing(s) :			PROJ. ENG.	46 CONVECTION TU	JBES INCHES O	D. TUBEWALL I	NCHES MATERIAL	
6. Fittings and accessories included: -				47 RETURN BENDS			ERIAL	
7. Descriptive bulletin § I (or specifi	ications) :			49 DESIGN	RADIANT PRESSURE	PSIG TEM	PERATURE 'F	
				50 51 CODE	CONVECTION PRESSURE	PSIG TEM	PERATURE 'F	
D. Remarks :				52 BURNERS - NO	түре	MFG.	PILOTS RI	EQ D.
				53 HEATER TYPE		HEATER WEIGHT		
				55	NSTRUCTION			
				56 REMARKS:				

	PROCESS DESIC CORPORATE EI	GN SECTION NGINEERING	WIPED FILM DATA SHEE	EVAPORATOR T	SHEET OF 4		PROCESS DESIGN SECTION CORPORATE ENGINEERING	i	WIPED FILM EVAPOR DATA SHEET	SHEET 2 OF 4
EMARKS	SCOPE:					EMARKS	FLUID DESCRIPTION	OPERATING CON	JACKET	CONDENSER
Ē						æ	TEMPERATURE	FEED°C DISTILLATE°C CONCENT°C	INLETOCO	
7E APP'R'V	DENTIFICATION	PERIORM.	ANCE REQUIREMENTS	STILLATE CU	NCENTRATE	ATE APP'R'V	VAPORIZATION TEMP. DESIGN TEMP. MAX. ALLOW. TEMP. OPERATING PRESS. DESIGN PRESS. DESIGN PRESS.	°C °C °C PSIG PSIG	°c. °c °c PSIG PSIG	●C ●C ●C ●SIG ●SIG
CHKD CHKD	NORMAL OPERA DESIGN COMPOSITION:						MAX. FLOW RATE MAX. FLOW RATE PRESSURE DROP MAX. ALLOW. PRESS DROP MAX. AIR LEAKAGE RATE			
	PHYSICAL PROPERT MOLECULAR WE SPECIFICGRAVITY	TIES					BEATT I E S : PRE-HEAT VAPORZATION CONDENSING SURFACE REQUIRED VAPORIZATION CONDENSATION			
DES SUPRV.	VISCOSITY: CP					D BY DES SUPRV	MAIN VESSEL JACKET SHAFT WIPER BLADES DEADINGS	MATERIAL MEC F S SEA ''0''	S H. SEAL OTATING FACE TATIONARY FACE I RING	
WN BY CHECKE	SPECIFIC HEAT LATENT HEAT. THERMAL COND VAPOR PRESS	г. ВТU/# ⁰ F втU/# ВТU/# ⁰ F FT. М ^{MH} G				AWN BY CHECKE	INTERNAL COND. TUBES	MAI	GASKETS	
	MELTING POINT	■ °c				INITIATOR	ILEM NOMID BY. # ?RPM MFR TYPE ENCLOSURE. CLASS 1 GROUP 1	FRAME BEA FUL FUL INSUL LR. D-DIV.	INTRASEIL	UBE
0846*	ICBN 0 AOJECT NO ITLE OSITION NO.	PLANT LC REF DWG SERIES	NO NO STEP NO.		BLDG NO.	E230B46 SHEET 2	0BN 0 ROJECT NO ITLE 40SITION NO+-	PLANT LOCATION REF DWG. NO. SERIES	STEP NO,	ISSUE DATE BLDG NO.



PROCESS DESIGN CORPORATE ENGI	SECTION	ROTARY VACUU DATA SHEET	IM DRYER PAGE 1 OF 2
	OPERATING		
MAT'L TO BE DRIED:		OPERATING PRESS.	PSIA PSIG PSIG
BULK DENSITY: AVG	LB./CU. FT.	HEAT TRANSFER RATE:	BTU/HR.
MOISTURE: INITIAL MOISTURE CONT FINAL MOISTURE CONTE	ENT:WT. %	ELECTRICITY: VOLTS/PHAS WATER: TEMP. MIN. PRESS,	9F MAX9F
AMT. OF MAT'L PER BAT	CH LBS. I'D CU. FT.	STEAM: PRESS, TEMP	PSIG 9F
C MAX, TEMP: WET DRYING RATE: EST, DRYING TIME/BATC	0F DHY 0F LB./HR. H HRS.	TEMP	OF
MOISTURE RECOVERY:			
	MECHAN	ICAL DATA	
MFR. SIZE&TYPE TOP OPENING: SIZE T.	Y P E	VACUUM SEAL: TYPE PACKING TYPE SIZE	NO. RINGS
BOT. OUTLET: SIZE -	түре	MECH. SEAL: MFR TYPE	
DRIVE: MFR		COUPLING: MFR TYPE	
	MAT	ERIALS	
SHELL: INTERNAL FINISH #: JACKET: VACUUM TUBE FILTER: PACKING:		MECH. SEAL: ROTATING F. STATIONARY BOT, OUTLET VALVE! VACUUM PIPING: JACKET PIPING: GASKETS:	ACE:
	MOTOR I	DRIVE DATA	
ITEM NO./MTD. BY HP/RPM/FRAME	DRYER /	VAC. PUMP / / /	/ /
MFR. TYPE/INSUL.	1	1	1
VOLTS/PHASE/CYCLE BEARINGS/LUBE.	1 1	/	/ /
FULL LOAD AMPS/SF L.R. AMPS	/	1	
DB NO ROJECT NO	PLANT LOCATI REF DWG. NO.	ON	SCALE
POSITION NO	SERIES	STEP NO	BLDG. NO
		DRAV	

	CORPORATE ENGINEERING	DATA SHEET PAGE 2 OF
	M	ISCELLANEOUS DATA
REMARK	LIST OF ACCESSORIES TO BE FURNISHED BY VE COMPLETE VACUUM SYSTEM INCL, PUMP, CO COMPLETE HEATING SYSTEM INCL PUMP, EXC	:NDOR' NDENSER, RECEIVER AND PIPING CHANGER AND PIPING
	HEATING CONTROL SYSTEM AUTOMATIC JOGGING AND POSITIONING SYSTE	EM FOR DRYER
	DINSTRUMENT SPECIFICATIONS FOR PURCHASE	BY H.LR.
V.R.V		
APE	UNMOUNTED EXPANDABLE SLEEVES	
DATE		
é		
a X		REMARKS
5 ≻		
2		
1.		
JPR V.		
DES SI		
28		
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>		
8 NA		
HQ		
тоя		
4111A		
	JOB N O PLAN	
-		DWG NO
<u></u>	PROJECT NO REF	ISSUE BLDG. NO
19835	PROJECT NO REF TITLE - POSITION NO, SERIE	ISSUE BLDG. NO

KG-1135 (6-81)

BOILER D	ATA SHEET
BOILER SHALL BE FURNISHED IN ACCORDANC	WITH THE M. W. KELLOGG COMPANY SPEC.
1. OPERATING CONDITIONS	11. EXPECTED PERFORMANCE
RATED CAPACITYPPH CONTINUOUS PER UNIT	
STEAM PRESSUREPSIG AT S.H. OUTLET	STEAM ACTUAL M. LB/HR
STEAM TEMPF AT S.H. OUTLET	BLOWDOWN %
ONFIRED OPERATION	FUEL
FEED WATER TEMPF. BLOWDOWN%	LOAD DURATION
FUEL	FURNACE LIBERATION
	M.B.T.U./CU. FT./HR.
A.P.I GRAVITY	FURNACE HEAT RATE
	M.B.T.U./SQ. FT./HR.
% BY	EXCESS AIR LEAVING BLR. %
ASH	
HYDROGEN	FLUE GAS ENT. A.H.
CARBON	
SULPHUH	STEAM AT SH OUTLET - PSIG
	SUPER HEATED STEAM
	FLUE GAS LEAVING BLB
	S WATER ENT, ECON.
	WATER ENT. BLR.
	AIR ENT. A.H.
	AIR LEAVING A.H.
	FURNACE
	BOILER AND S.H.
H.H. V. BTU/	40 FLUES
L.H.V. BTU/	OZ NET DRAFT LOSS
PSIG @ BURNER	ACROSS INST. CONNECT
TEMP. @ BURNER F.	BURNERS WINDBOX
TYPE OF INSTALLATION:	
CASING:	NET RESISTANCE
	20
AMBIENT TEMP:F	gzi
ALTITUDE:FT. ABOVE SEA LEVEL	
DESIGN WIND LOADING	
EARTH QUAKE LOADING:ZONE.	LB/HB IN H2Q F
INSTRUMENT AIR:PSIG	
E	
	· · · · · ·
START-UP	
	MANAGING OFFICE ENGINEERING OFFICE
	KELLOGG
	EQUIPMENT DATA SHEET
<u><u><u> </u></u></u>	
š 2	
	M
	NO SPEC NO.
NO DATE ENGINEER CHK/REVIEW APPROVE I RE	2'N, N O PAGE OF

KG-1124 (7-81)

	PACKAGE BC	DILER	2 C	DATA SHEET
	III EQUIPMENT		Ň	IN ACCORDANCE WITH KELLOGG SPEC.
	TYPE		AC	HEIGHT ET: DIAMETER
	DESIGN PRESSURE, PSIG		5	ARRANGEMENT.
	FURNACE TYPE		5.2	· · · · · · · · · · · · · · · · · · ·
	FURNACE VOLUME, CU. FT.		25	FURNISHED BY
			E R	
	HEATING SURFACE		PLA PLA	IN ACCORDANCE WITH KELLOGG SPEC
~	CONVENTION SURFACE: CIRCUMFERNTIAL BASIS	S)	-	
Ľ,	FURNACE, SQ. FT.		۳.	BOILER
ē	BOILER, SQ. FT.		S04	SUPERHEATER
	TOTAL, SQ. FT.			AIR HEATER
	C RUNS		ž s	SETTING (OVERALL)
	NO. REQUIRED		ME S	UNIT (OVERALL)
	UPPER DIMENSION			
	LOWER DIMENSION		ŝ	STEAM
	VERT. HT. (C.L. TO C.L.)		NN	WATER FEED
÷.	тург		S⊢	BLOW DOWN
ATE	TEMP. CONTROL			
오픈	SURFACE AREA, SQ. FT.		H	SHIPPINGLBS
٥œ			VEI	OPERATINGLBS
S S S S S S S S S S S S S S S S S S S			>	MAX.ERECTIONLBS
ΞE				IV GUARANTEE CONDITIONS
٩e	TYPE			
₹5	SURFACE AREA, SQ. FT.			FUEL
s		Antre		CAPACITY, P.P.H
ER.	REQ. MAKE RANGE	RANGE		(@ S.H. OUTLET)
ŝ	GAS	· ·	1	THERMAL EFFIC. %
ā				MAX. NET DRAFT LOSS
_				MAX, NET AIR RESISTANCE
No No				BLOW-DOWN
F S			1	FEED WATER TEMP. DEG. F.
B N				MAX ALLOWABLE BOILER
õõ				CONCENTRATION P.P.M.
	FURNISHED BY:			
			1	
			1	
+				
		- 1	KELLO	
2		1		EQUIPMENT DATA SHEET
2 3		CLIEN	NT _	
5 7		LOC	ATION	JOB NO
r	, 	ITEN	M	
N	D DATE ENGINEER CHK/REVIEW APPROVE	E REO'	NO NN 1	

I NDUCED DRA	AFT COOLING TOWER
SERV CEI MANUFACTURERI	ENGLISH/METRIC UNIT
SITE ELEVATION:	PREVAILING WIND DIRECTION
DIMENSION: LENGTH	JWER DESIGN WIDTH HE GHT
HEIGHT BASIN CURS TO FAN DECK_	FAN STACK
H. W. INLET ABOVE BAS IN CURB	
ACCESS TO TOP OF TOWER	ASTN CURB/IPERIMETER-INTERNAL
AIR TRAVEL THROUGH FILL	AVG. FILL HEIGHT
FUL SLAT SPACING HORIZONTAL	AIR VELOCITY THROUGH FILL
LOUVER ANGLE GROSS LOUVERE	D AREA FAN DECK LIVE LOAD
MATERIALS	OF CONSTRUCTION
FAN STACK:PARTITIO	INSSHEATH I NC
THEMS RECEIVING TRESERVATIVE TREA	
FAN BLADES	SFILL SUPPORTS
RING LOINT CONN	
FAN HUE	DR I VE SHAFT
	EQU I PMENT
HUB DIAMETER	MODEL NO
SPEED REDUCER MANUFACTURER	MODEL NO.
DRIVE SHAFT: MANUFACTURER	MODEL NO
RAIED H. P	
Elect. Area: Class: Class Gr.	Div. /Non-Hazaedowsus; Vogetss Ph. Hz
Enclosure Insulation Temp.	RiseOC aboveOC BvO_S.F. Temp Code
Item HP Full Locked 4 3 4 3 10	ad Frame Lub. Weight Up/Down Heater's F
No. KW Load Rotor 4 4 4 4 RP	M No. (1) LB/KG LB/KG Watts
├── │ 	
FANSTACK	VELOCITY RECOVERY YES/NO. HEIGHT
DISCHARGE AREA	DISCHARGE VELOCITY
PRESSURE DROP	
	STOP;
	FLOAT CONTROL
LABOR BY: UNLOAD I NG	G & HAULING BY-SUPERVISION BY-
_ EST. MANHOURS REQ. FOR ERECT I ON	FREIGHT
MW KELLOGG SPECIELCATION: 1141-E2 SI	I SCELLANEOUS NOW LOAD SPECIFICATION
WIND & EARTHQUAKE SPECIFICATION	PAINT SPECIFICATION
NOTES.	
6	MANAGING OFFICE ENGINEERING OFFICE
n_5	KELLOGG
	DATA SHEET
	LIENT
	ITEM INDUCED DRAFT COOL ING TOWER
	ITEM HOSPEC. NO
NUL DATE ENGINEER CHR/REVIEW APRROVE	REO'N. NOL PAGE OF

		Smonth J		CORDING CORDO	DATION	Item Page		
		BIONE C	K WEBSTER EN	EDECIEICATIO	N	Prelin	inery	
		INS	SIRUMENI 3	SPECIFICATIO	N	Final		
Client					J. (0. No.	1.1. Marca - 1	
Apparatus					Pre	jest No.	~~~~	
Service					Da	te	By	
Based on	INSTRUM	IENT CASE		BU	LBS OR T	HERMOCOUPLES	3	
Lonetton	1			Location				
Туре	-			Bulb Type				
Case Finish	_			Thermocouple Mati				
Connections	Back - Bo	ttom		Socket Material				
Mounting	Flush — Pr	rojecting		Socket Length				
Chart or Dial	No	Size	in. diameter	Ext Neck Length				
Range	Flow	Stat	tic Press.	Standard Pipe Thread		Above	A	
Control Boint	Temp.	Diff	. rress.	Tubing or Land		Below	R	
Chart Volume	Measured of	n • F	Ib. Ga Basis	Provide on Long	RES	PONSES		
Clock	Electric -	Spring Wound, 2	Day Hour	Throttling Band				
Current		Volta DC	Cycles	Reset	Automatic Manual			
Accessories	Ink. 100 Ct	harts etc to be incl	uded	Rate Response	Yes — No			
				Pneumatic Set		VALVE IS		
	INST	RUMENT		CON	RULLED	VALVE - Item	NO.	
Location	-			Bervice				
Floet Body	Care Mai	61	Trim	Valve	Size	in. Drilli	D#	
Float Flange Conn.	8444	Drđ	ling	Plug Type Value	Ralanced F	rd Parabolic V-Port or Equal		
Float	8/24	in. Met	1	Туре	Disphrage	Bpring Loaded		
Stuffing Box	With Orme	e Seal		Body	Mati	Tri	m	
Orider Flangts	Matl	Det	lling	Stem	Sitting -	Rotary		
Orldee Plate	Matl			Stuffing Ros	With Gree	en Benj		
Instr Body Niff Range	- •	in. Water Co	above liquid level	Operating Medium	Open	9	Ib ga Correction	
instr Pody Located			It islow orther plate liquid level	Valve Action	Open with	Failure of		
Mercury Included			it from pride place	Valve Positioner	Yes No -	ith Rypan		
Sealing Medium				Redisting Fine	Ves - No			
	1		CONDITIONS AN	REQUIREMENTS	I			
In			Control Valva	In	1		Control Valve	
Fluid				Nor Quantity @	7		Change and	
Sp Gr @ 60" F Water =	1			Max Quantity @	1		L	
So Gr @ Nor T & P Water	= 1			Min Quantity @	P			
Viscosity		ep @ •F	ep@ •7	Nor Pressure (Up Stream)	lb ga	1	
Corrosion Present	Yes -	- No	Yes — No	Max Pressure (Up Stream)	1b ga	. 1	
		in.	in.	Down Stream Pressure		lb ga	1	
Line Size	1	•7	1 °F	r resture Drop		16		
Line Size Normal Temperature		**	•=	Amount of Superheat		• •		
Line Size Normal Temperature Max Temperature Supply Metal Tag. according	urely wired, ma	•rked with Item N	•r	Amount of Superheat		•7		
Line Size Normal Temperature Max Temperature Supply Metal Tag, seco REMARKS	prely wired, ma	*r	•p	Amount of Superheat				
Line Size Normal Temperature Max Temperature Supply Metal Tag. sec- BEMARKS	orely wired, ma	•r	• p	Amount of Superheat		•ŗ		
Line Size Normal Temperature Max Temperature Supply Metal Tag, sect HEMARKS	urely wired, ma	•r	• p	Amount of Superheat		• 9		
Line Size Normal Temperature Max Temperature Supply Metal Tag, sect HEMARKS	urely wired, ma	•r	•r	Amount of Superheat		• •		
Lice Size Normal Temperature Mart Temperature Supply Metal Tag, sec HEMARKS	urely wired, ma	•r	• p	Amount of Superheat		• • •		
Line Size Norma Temperature Max Temperature Sundy Metal Tag, seen HEMARKA	orely wired, ma	•r	• p	Amount of Buperheat		•,		
Line Size Normal Temperature Max Temperature Sunniv Metal Tar, see UEMARKS	urely wired, ma	rked with Item N	epober	Amount of Superbeat		•,		
Lino Size Normal Temperature Normal Temperature Supply Metal Tag, see UEFMARKS	urely wired, ma	•p	spuber	Amount of Superheat		• •		
Lizo Size Nornal Temperature Max Temperature Supply Metal Tas, see UEWARKS	prely wired, ma	• y	ynber	Amount of Superbest		•,		
Lizo Size Normal Temperature Max Temperature Sunniv Metal Tag, acc HFMARKS	urely wired, ma	• p	prober	Amount of Buperheat		•,		
Line Size Normal Temperature Max Temperature Sunniv Metal Tax, see UEMARKA For Typical Installati For Typical Installati	orely wired, ma	rked with Item N	ppber	Amount of Buperheat		•,		
Line Size Normal Temperature Nat Temperature Suganly Metal Tag, see UEWARKS For Typical Installati General Specifications	orely wired, ma	• r rked with Item N	pruber	Amount of Buperheat		·,		
Lion Size Normal Temperature Nat Temperature Sunniv Metal Tas, see UEFMARKS For Typical Installati General Specifications Copy to	ureiv wired, ma	rked with Item N	puber inter in	Amount of Buperheat Amount of Buperheat		· p	Upte	

DIFFERENTIAL PRESSURE TYPE FLOW INSTRUMENT

		SERVICE DATA	PRESE	NT	NEW
		ITEM NO.			
		LOCATI ON			
	÷	SERVI CE			
2	8				
<u>-</u>	2	LINE SIZE			<u>t</u>
	Date	ORIFICE SIZE (Bever)			
		MANOMETER (Range & migi)	(hid		
		CONNECTIONS (Flange & Series)			
		ELOWING FLID			
		SP. GR. AT 60F			
		FLOWING TEMP.			
		FLOWING PRESSURE			
		SEAL MATERIAL			
		SEAL TEMP.			
		SEAL GRAVITY AT 60F			
		MAX. FLOW			
		CHART NO			
			(
			I NSTRUMENT		
		PRIMARY METER		REMARKS	
		TYPE = <u>Recording</u> , Indicating	; Blind, Transmitting		
		MFR. MODEL			
		MANOMETER = 1500 psi = Carbo	n Steel		
RE	2	RECEI VER			
	1	TYPE • Conventional, Miniature, None			
		MFR. MODEL			
	.p'd.	HOUNTING Control Board, Lo	cal		
		CHART TYPE • Circular 24 Hour, Strip			
~		CHART DRIVE <u>spring</u> , Pneuma	atic		
e l	4				
		CONTROLLER			
		TYPE • Conventional, Force Balance, Pneumatic		4	
		MFR. MODEL	ald	-	
		MANUAL CONTROL BYDASS - Yoo	No	ł	
		MANUAL CUNIRUL BIPASS <u>IEA, NO</u>		t	
	ę	CONTROL FRANKS • Prop. Band. Reset. Rate		†	
	Da.	INCREASE IN FION OUTPUT TO	1		
		REQUISITION OR REQUEST -		1	
		REF. SPEC. SHEET-		1	
		REF. DRAWING -		1	
		INST. TAG DATA		1	
				<u> </u>	
		By	HUMBLE OIL & REFIN	ING CO.	Appr'n.
		App'd Bi	AYTOWN ENGINEERING	DIVISION	SubZone
Re	-	Date	INSTRUMENT SPECIFI	CATION	NO
		· · · · · · · · · · · · · · · · · · ·			

165 BROAD NEW YORK. U.	CORPORATION TEMPPRESS. WAY INSTRUMENTS S. A.												
CUSTOMER'S NAME	REQUISITION NO.												
CONTRACT No.	DATE												
11 EM-	PAGE No.												
SERVICE-	CHANGE No.												
INSTRU <u>I</u> H	ENT												
No. Req'd menu Type m													
Location Case No.	Reg'd Bulb Mat'l												
Mounting	lb. Elev. Instrum												
RangeScaleTy	pe Socket Mat'1												
Chart E	ngth Below Threads												
Thermal System HIM F EX	P. T. Size												
Control Point.	bing Mat'l												
Control. HI I CONTROL I I CONTROL I I CONTROL I CONTROL	bing Length												
ранция и на волоти на													
Ciock Volts Cycle Phase													
Connections													
Accessories:,type air reducing valve,													
air filtercharts, ink set and air gauges													
Special													
CONTROLLED	VALVE												
Type Op Location Value	erating Medium												
Body Material Ends	ve to With Failure of Ai Cooled Bonnet Required												
Body Material Blac Ends Air- Faced and Drilled King Air-	Vec toWith Failure of Ai Cooled Bonnet Required re Positioner Required												
Body Material Ends Air- Faced and Drilled For the second seco	Vec toWith Failure of Ai Cooled Bonnet Required re Positioner Required ra Reducing Valve and Filter Required												
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Rody Material Ends Air- Faced and Drilled Faced and Drilled State of the state o	Ve toWith Failure of Ai Cooled Bonnet Required re Positioner Required ra Reducing Valve and Filter Required cial												
Rody Material Ends Air- Faced and Drilled Faced and Drilled Faced and Drilled Trim Stuffing Bor Sealed, Isolating Valve-, Stuffing Bor Sealed, Isolating Valve-, N	Ve toWith Failure of Ai Cooled Bonnet Required ve Positioner Required ra Reducing Valve and Filter Required raine												
Rot. Redu Size For the second	Ve toWith Failure of Ai Cooled Bonnet Required ve Positioner Required ra Reducing Valve and Filter Required cial D REQUIREMENTS VALVE												
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No. Redu Size Formula Air Body Material Ends Valve Faced and Drilled Formula Size Trim Size Size Valve Plug Sealed, Isolating Valve-, Size Stuffing Bor Sealed, Isolating Valve-, No SERVICE CONDITIONS No Sp.Gr. @ 60° F. Air 1 No	We toWith Failure of Ai Cooled Bonnet Required Positioner Required Ta Reducing Valve and Filter Required Ta Reducing Valve and Filter Required Ta Reducing Valve Ta Re												
No. nequ Size Proprint No. nequ Air Body Material Ends International Size Air Faced and Drilled Sealed, Isolating Valve-, Ext Stuffing Bor Sealed, Isolating Valve-, International Size Stuffing Bor Sealed, Isolating Valve-, International Size Sp.Gr. @ 60° F. Material Material No Sp.Gr. @ 60° F. Waterial No No	We toWith Failure of Ai Cooled Bonnet Required re Positioner Required ra Reducing Valve and Filter Required ratial D REQUIREMENTS D REQUIREMENTS T. Quan. @9 F ax. Quan. @9 F T. Press. (Upstream) b. Ga												
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Not. Redu Size For parameters Air- Faced and Drilled Ends Katerial Katerial Finid Sealed, Isolating Valve-, Sealed, Isolating Valve-, Sealed, Isolating Valve-, Fluid Sp.Gr. @ 60° F. Air =1 Material No Sp.Gr. @ 60° F. Air =1 Material No Na Sp.Gr. @ 60° F. Air =1 Material Material Na Normal Temp. ° F. Material Material Na	We toWith Failure of Ai Cooled Bonnet Required re Positioner Required ra Reducing Valve and Filter Required rial D REQUIREMENTS D REQUIREMENTS T. Quan. @OF												
No. Require Size Size Formula Size Air- Body Material Ends Air- Faced and Drilled Ends Ends Trim Ends Ends Valve Plug Sealed, Isolating Valve-, Sealed, Isolating Valve-, Suffing Bor Sealed, Isolating Valve-, Material Sp.Gr. @ 60° F. Air Material Material Sp.Gr. @ 60° F. Air Material Material Normal Temp., ° F. Material Material Normal Temp., ° F. Material Material	We toWith Failure of Air Cooled Bonnet Required re Positioner Required ra Reducing Valve and Filter Required ra Reducing Valve and Filter Required D REQUIREMENTS D												
No. Require size Size Formula size Air Body Material Ends Air Faced and Drilled Formula size Size Valve Plug Sealed, Isolating Valve-, System Stuffing Bor Sealed, Isolating Valve-, M Sp.Gr. @ 60° F. Wit = -1 M M Sp.Gr. @ 60° F. Wit = -1 M M Sp.Gr. @ Nor. Con. (T&P) M M Viscosity @ ° F. M M Ine Size F M Max Temp., ° F M M REMARKS: Normal Temp., ° F M	We toWith Failure of Ai Cooled Bonnet Required rea Reducing Valve and Filter Required rea Reducing Valve and Filter Required rea Reducing Valve and Filter Required D REQUIREMENTS												
No. Redu Size Formula in the second	Ve toWith Failure of Ai Cooled Bonnet Required re Positioner Required ra Reducing Valve and Filter Required rial D REQUIREMENTS D REQUIREMENTS D REQUIREMENTS VALVE Ax. Quan. @ F Ax. Quan. @ F S. Ga , S. Ga , ssure Drop												
Rody Material Size Formula Air- Faced and Drilled Ends Image: Size Kar Trim Size Sealed, Isolating Valve-, Material Image: Size Not Normal Temp., o F, Material Image: Size Normal Temp., o F, Material Image: Size Image: Size<	We to												
No. requ Size I before the second se	We to												
Not. Red Q Size Propringing of Notes in the Air- Body Material Ends Intermediation of Notes in the Air- Frind Sealed, Isolating Valve-, Sealed, Isolating Valve-, Stuffing Bor Sealed, Isolating Valve-, Intermediation of Notes in the Air- Stuffing Bor Sealed, Isolating Valve-, Intermediation of Notes in the Air- Sp.Gr. @ 60° F. Air = 1 Sp.Gr. @ Nor. Con. (TK Water=1) Intermediation of Notes in the Air- Sp.Gr. @ Nor. Con. (TK Water=1) Intermediation of F. Intermediation of Provide in the Air- Normal Temp. ° F. Intermediation of F. Intermediation of Provide in the Air- REMARKS: Normal Temp. ° F. Intermediation of Provide in the Air- REMARKS: Normal Temp. ° F. Intermediation of Provide in the Air- Remarks: Normal Temp. ° F. Intermediation of Provide in the Air- Remarks: Normal Temp. ° F. Intermediation of Provide in the Air-	Ve toWith Failure of Ai Cooled Bonnet Required re Positioner Required ra Reducing Valve and Filter Required ra Reducing Valve and Filter Required ra Reducing Valve and Filter Required												
FORM NO. 2788				EDD FO	RM 2792 1-6	4							
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	LOCATION	JOB NO.	FILE NO.	—— I []	GAU	IGE GLASSE			IOE	3 NO.	F	ILE NO.	
INSTRUMENTS	PREPARED BY		DATE		A	ND COCKS	PREPARE	D BY			C	ATE	
Nether SPECIFICATION	CHECKED BY		DATE		Northern S.P.	ECIFICATI	O N CHECKED	BY			C.	ATE	
Gen SHEET	APPROVED BY		DATE		Ges Company	SHEET	APPROVE	D BY				DATE	
	A INDICATES REVISI	ION						ATES RE	VISION				
GENERAL					5	SAUGE GLAS	SES			_G A	U <u>GE COC</u>	<u>(S</u>	
I TYPE		-		1 SU	PPLY U		AUGES & COCK	5 15	SUPPLY			DNLY	
3746 110				2 TYF		ES REQUIRED TRANSPAREN	TOTUBULAROREELE	× 17	ONNECTION	S-NPT	VESSEL	CALICE	DRAIN
AVESSEL OR EQUIPMENT NO.				3 CC		ı/2''	34 11	_ "ÌÌ [MALE	FE MALE	FE MALE
BODY						ТОР ВОТТОМ	SIDE BAC	к.			[]1/2"	1/2 "	1/2"
5MATERIAL					са м	WELDING PAD	OTHER					3/4.*	3/4
7 BTM CONN LOCATION				5M IN	MUM RATING	SIEEL	PSIG	_°F					
BCONN-SIZE										BODY	(TRIM	
9CONN SCREWED OR FLANGED								18	AAT'L				٩.
IOCASE MOUNTING									MINIMU			IG 🖷	f
12ROTATABLE HEAD		-++		1				20 0	омят. [PLAIN	CLOSING	QUICK	CLOSING
13					-	ACCESSORIE	s	1	Ľ	HANDV	VHEEL	LEVER	HANDLE
FLOAT OR DISPLACER				6 ILLU	MINATORS:	- <u>)</u>		21	CESSEL	PLAIN U		SOLID	SHANK
					TING-COOLIN	G INTERNAL		22 0	AUGE	PLAIN U	INION	PLAIN	OTHER
				9	CHAMBERS	EXTERNAL _			onn . 🗋	SPHERI	CAL UNION	отн	ER
L 7 TOROUE TUBE MATER AL				10 NOM	FROSTING TYP	PE :		23	BONNET	SCREW	ED		BOLTED
18AIR FIN		_		11 CAL 12 SUE	BRATED SCAL PORT PLATES	-E		24	CREW	SEAT			NO
19 TRANSMITTER				13 GUA	RD RODS:			26	ALL CHEC	кs			NO
20 TYPE			I	14 OTH	IER			27	ACKING	MFR S	TD		OTHER
2.000TPUT				NO	FS ·			28	AFR MODEL	NO			
22RECEIVERS ON SHEET NO									OTES :				
23 TYPE													
24 PROPORTIONAL . RESET													
250 <u>UTPUT</u>	<u> </u>												
260N LEVEL INCREASE OUTPUT		+ +											
ACCESSORIES						NO OF	VISIBLE	MER M	OD ODED	0050		ACCER	1
281FILTER AND REGULATOR				REV.	QUAN. TAG	S NO. SECT.	LENGTH CPLG	5 NO.	PRESS	TEMP.	SERVICE	SORIES	NOTES
29GAGE GLASS CONNECTIONS													
3.1 PURGE CONNECTION								-					
32ELECTRIC SWITCH													
331													
35UPPER LIQUID													
36 LOWER LIQUID								-				+	-
37SP GR UPPER LOWER	╉───┼───┼───									I			
38PRESS MAX, NORM	+ $+$ $ +$ $-$			-+ }				-		—			
SELEMP MAX, INORM		-+	+ -				} _		No.				
									_			<u> </u>	
REMARKS:													
				——				-	_				
							t		_			+	.
							1 1	1					1
				REPROE	UCTION OF	STANDARD NO.	53.02-2						

REPRODUCTION OF STANDARD NO. 53 02-1

CONTROL	VALVE	SPECIFICATION	SHEET	

SPECIFICATION No...

DATE: DATE:								CONT. NO. OF					
MA	DE BY:			CHECKE	D BY:			W.N. NO.			PA	GE	
DA	TE						1						
BY							I	1		L			
RE	VISION	1	2	3	4	5	6	7	8	9		10	11
	our d'Albad.				+						I		
	MARKS:										1		
0	ON ACTUAT	ING FLUIE) FAILU	RE. VALVE	TO:						L		
F		CLOSE V	VITH IN	CREASE OF	-								
5	VALVE TO:	OPEN W	ITH INC	REASE OF									
7	ACTUATING	FLUID											
υŀ		FINS	REQUI	RED							1		
ōŀ		PILOT	RANG	θE							1		
ž į	ACCESSORIES	5: VALVI	E POS	ITIONER							1		
븠ᅣ	TYPE OF S	STUFFING	BOX										
αŀ	TYPE OF P	LUG									î –		
зŀ		SEAT									1		
₽₽		PLUG			<u> </u>						í –		
Ξf	MATERIALS:	BODY									1		
ζt	LUBRICATOR	REQUI	RED								1		
_۴	FLANGE FA	CING									-	`	
ľ	BODY CONN	ECTIONS	& RATI	NG							i –		
Ľ,	REDUCED A	REA POR	TS					1			-		
Ē	NOMINAL	IZE. INCH	IES										
+			10.	<u> </u>									
	CPS (a °F												
h	VIS. OF LIC	,	CP	5 @ °F	· · · · · · · · · · · · · · · · · · ·			-			• • •		
- f			T NOR	M. OP. TEN	ир.			1					
a f	SP. GR. OF	LIQ. REL.	TO WA	TER AT 60	°ŕ			1					
있는	M.W. OF V.	APOR			-								
υt				TOTA	L			-					
ωŀ				LIQUI	D								
ŝ	MAX. FLOW	AT NORM	. OP. TI	EMP VAPO	R								
Î				TOTAL									
ΰĒ				LIQUI	D								
Z l	NORM. FLOW	AT NOR	V. OP.	TEMP., VAP	OR								
				MAXIMU	JM								
-	PRESSURE	DROP, PS	51.	NORM	A L								
Ξf				MAXIMU	м								
Ξt	NLET Op.	PRESS.,	PSIG,	NORM	A L								
υf				MAXIMU	JM								
1	INLET OD	TEMP °F		NORM	A.I.								
ŕ		RROSIVE	DUE TO	- -									
_			LVE		1			1					-
INS	TRUMENT	OR CONT	ROL S	ERVICE				_					
TAG	6 NUMBER												
ITE	M NUMBER	R											
в/м	NUMBER												

FORM 2834 5-64			PAGE OF
VALVE ITEM NO.		JOB NO,	W. 0. NO.
OPERATOR PREPARED	BY		DATE
Northern SPECIFICATION CHECKED	BY		DATE
Company SHEET APPROVED	BY	EVISION	DATE
	UNIED R		
	INF	D_R MATION	
OPERATOR	6	MANUFACTURE :	ALVE
ELECTRIC PNEUMATIC HYDRAUL	IC 7	FIGURE NO.	SIZE LOT
PNEUMATIC HYDRAULIC	6	DESIGN PRESSURE	PSI PSI
TIME FULL OPEN MIN MAX	10	VALVE STEM POSITIO	N:
FULL CLOSE MIN MAX		HORIZONTAL	VERTICAL
MINIMUM TOROUE OUTPUT W/FULL	1	GEAR RATIO	
POWER GAS <u>PR</u> ESSURE	12 PERATO	TOROUE REQUIREMENT	
ELECTRIC		PNEUMATIC	OR HYDRAULIC
POWER SUPPLY:	17	POWER GAS:	
VOLTAGE <u>P H A</u> S EFREQ	— [_]	L AIR	GAS
CLASS I GP D GENERAL PURPOSE	18 19	DRIVER MOTOR:	PSI
OTHER	. /	TURBINE	VANE PISTON
BREATHER AND DRAIN			—
DUTY RATING:		VANE MOTOR W/G	EARING
	DU S		
¢	ONTRO		
PILOT VALVE	25	EMERGENCY SHUT	DOWN SYSTEM
	26	ESO OPERATION WHEN	PILOT:
	27	SYSTEM PRESSURF -	DEFERCISIONIZED
VALVE ACTUATION:	2.8	VALVE ON SHUTDOWN	
PI LOT VALVE		TO OPEN	TO CLOSE
L ACTUATES OF CONTRACTUATES	29	ESD TO OVER-RIDE	REMOTE OPERATION
VOLTAGE		MANUAL	OPERATION
PILOT PRESSURE FOR VALVE ACTUATION	3 0	OVER-RIDE:	
INCREASES DECREASES		MANUAL	AUTOMATIC
CONTROL SYSTEM:		PUS	HEULTON
	32	MOUNTED:	
	IC COI		
	35	HEATERS:	SPACE HEATERS
		h h	WITH THERMOSTAT
	ESSOR	155	•
	<u>тр.</u> Т		TYPE OF ELECT. SWITCH
REMOTE SLIDE WIRE TRANSMITTER		NORMALLY OPEN I	N EA. DIR. OF TRAVEL
REMOTE POSITIVE INDICATOR W/PWR. SUP.		NORMALLY CLC	ISED IN EA. DIR. OF TRAVI
INTEGRALLY MOUNTED REVERSING STARTER	43	AOJ. TORQUE SW:	
MANUAL HMO VALVE:	4.4	ENCLOSURE:	GENERAL PURPOSE
LIMIT SWITCHES: ELECTRIC PREUMA	ATIC 4S	MANUAL HAND PUI	MP
ADJUSTABLE POSITION	46	GAS HYDRAULIC T	ANKS
GEAR DRIVE	47	ALL NECESSARY AL	APTERS TO FIT UNIT. TO VA
MICO	48	DUS	0002
MISC	LLLAND	000	
ALL WIDING REQUELT TO EVELOCION SECON	20	ALL TURING REQUI	SHT TO BULKHEAD FITTIME

ARTHUR G. MCKEE & COMPANY FORM 439

	ARTHUR G. M RELIEF ITEM No. TAG NO. VALVE SERVICE	CKEE & COMPANY	
	MANUFACTURER & TYPE No. NUMBER VALVES REQ'D. NORMAL SYSTEM PRESS., PSIG. NORMAL SYSTEM TEMP., °F. GOVERNING UPSET CONDITION ACCIUMULATION, DESCENT		
	FLOWING FLUID FLOWING FLUID FLOW QUANTITY, GPM, SCFM, #/HR. FLOW SP. GR. OR MW. FLOW SP. GR. OR MW. FLOW VISCOSITY SET PRESSURE, PSIG. ACCUM. INLET PRESSURE, PSI. ABS. BACK OPESS DUC / DEPESS DUC / DEPESS		
MADE E DATE	ROTO ORIFICE AREA, SQ. IN. ROMINAL SIZE, INS. ORIFICE AREA, SQ. IN. VALVE TOTAL ACTUAL AREA, SQ. IN. Z DODY CONN. & RATING-INLET		
	OUTLET BODY & BONNET TRIM SPRING CONTRACTOR STYLE TOP TRIM STYLE TOP		
MADE BY. Date APP. BY.	REMARKS:		
¢ J	WN No. PAGE	OF CONT. No.	B/M No.

ED-325-1 PRINTED IN USA.

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1	6 A P								10.8	NO						
+	PLANT								PUYP	NO						
、 †	LOCATION								NO	UNITS						
- Si ti -	SERVICE	SERVICE						YOTOR DRIVE								
The second se	VENDOR-SIZE NOD	EL			TURAINE DRIVE											
÷.	TYPE				SEDIAL NO											
Ĕ.	1116		_	0	DATI	NA	C O N	DITIO	IS							
- 1 - 1 - 1	£1.01 0			-	MARMAN	SPY 0	PT		NPSH	AVA	п			FT FLUID		
	PUNP TEMP *F				DESIGN	PH O	PT		NPSI	RE	Q'D .			WATER		
1 I I I I I I I I I I I I I I I I I I I	SPCR ODP.T				DISCHARG	E PRE	SS PSIC		NPS	H RE	Q'D.			FLUID		
2.2	VIP. PR 🔘 PT-PSIÅ				SUCTION	PRESS-	PSIG									
2 2	VISC OD PT-SSU				Ар		P S I	FT								
12					C	ES	IGN									
3 B	CASE SPLIT				RADIAL B	RG TYPE	2		SHA	F T -	HAX	DIAN	' A1	CPLC		
55	SUPPORT				THRUST B	RC TYPE	1		WEA	R 8	ic CI	LEAR	ANCE			
11	IMPELLER TYPE				BRG LUBI	3			STUR	FING	B O X	I D	' I	EPTH		
ι.	CORROSSION ALLOW				VISIBLE	LUBRI CA	TOR		BAS	E PL	A T E					
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a a	COOLING DATEN	11.00	P 5.	16 1	NUZZLES	POSITIO	NI SIZE	K A T TNG	TEC	TC				TT THE COLD		
E E	C.W JACKETS	KE Q'D	6 P	Ĭ	SUCT				TES	15			NEGD	VIINESSED		
y at	KAU BKG			-	DISCH				5107	183 05747	PEGI	01				
Ê i	INKUSI BAS			_	DBATN				0.00	CAD NA	10					
ž	PEDECEAL		-	-	SEAL				PERI	UNHA	NUE	-				
¥ Å	PEDESTAL				JEAL			c 1	nr.) N						
8 F				—	N.	A.I.E_	K I A L	<u>ا</u>	THRO	AT 0		10				
2	INPELLER			-+	Nº & SI	ZE RING			CASI	NG G	ASKET	1				
	CASE WRG RING					NL.			GLA	10						
έł.	INP WRG RING				ROTATIN	G FACE			BAS	PL	TE					
21	SHAFT			1	STATIO	AR" FAC	E		COUP	LING						
Ţŝ	SHAFT SLEEVE				SEAL	AING										
1	LANTERN RING				AUX. S											
₩					PE	RFC	RMA	NCE								
F .	NO. STAGES				ROTATION FACING PUNP CPLG					WEIGHTS :						
22	NAX. INPELLER - INC	HES			SHUTOFF	HD W/D	ESIGN INP	PSI	PUN	P &	CPL	6.		#		
	BID INPELLER - INC	HES			NAX.W.P.		PSI6 (10)	•F	BAS	E				#		
Ĕ ġ	YIN IMPELLER - INC	HES		_	HYDRO T	EST PRE	SSURE	PSIG	TOT	AL				#		
54	EVE AREA - SO MG	NE S		_	FURNISHED	81	VENDOR	K.H.PAKSUNS	SPAC	E NE	QUIRE	HEN IS) → W III	DRIVER		
	NESION EEE 'A				P U	SF			001	PALI	- LCR	161N Th		INCHES		
To be	DESIGN LIFE A			_	COL	DIINC			011	INC	0.00	4		INVILU		
8 ē				-	000	AUARD			CROS	S- SEC	TION	#				
n a	MAX B HP MAY IMP					0.R			PERI	ORYAN	CE	CURVE	#			
deal .	MAK. 011, MAK (4)			-	TURE	INF			G F	p #			-			
th be				-	104	DR	VER		<u> </u>							
p s	NOTOR HP		RPN		TURBINE		<u>нр —</u>	RPN								
22	VOLTS PHA	SE	CYCLE	8	STEAM		PSIC *	F SUPER HT								
1	ENCL.				EXHAUST		PSIG	/R#/HP - HR								
÷ 2	FRAME NA. N	IT	ŧ	۴	TOTAL	STEAN *	/HR	¥1 #	1							
1.3	SPEC Nº				SPEC NL											
	THE RALPH M P	ARSONS														
	COMPANY												Γ			
	LOS ANGELES			N 1	DATE		REVISIONS		BY	C H'X	APP'D	APP'D.	Γ			

FORM 495 500 3-56

				CENTRIFUG	AL PU	MP						
SERVICE				DRIVE	OTY		SEF	RIAL		ITEM		
			OPER									
			SPARE									
GENERAL SPEC					HFR							
PUMP CLASS SIZE AND TYPE												
				PROCE	SS-DAT	A						
LIQUID			GPM Н	от			SUBMERGENCE	AVAIL	FT.			
PUMP-TEMP F			DISCH	PRESS PS	IG			SUBMERGENCE	REODE	T.		
SP GR @ PT			SUCT	PRESS PSI	G			TOTAL SUCT	LIFT F	r		
VAP PR @ PT PSIA			DIFF	HEAD PSI				NPSH AVAIL	FŤ			
VIS @ PT SSU			DIFF	HEAD FT				NPSH REQD	T WATE	R		
CORR DUE TO			MAX S	UCT - PRESS	PSIG			NPSH REQD	TOIL	_		
SOLIDS			IF CR	ITICAL, M	IN GPM							
				REQUI	RÉMENT	5						
AXIS			MINC	ASE CORR-	ALLOW			CPLG GUARD				
SPLIT			SUPPO	RT				BASEPLATE				
IMPELLER			BRGL	UBE				DRIVERS BY				
COOLING WATER F	a	PSIG	FLUSH		REQD	FLUID	GPM	DRIVERS MT	D BY			
C-W JACKETS	REQD	GPM	CAGE	RING				TESTS		REQD	WITNESS	
RADIAL BEARING			WEAR	RING				SHOP INSPE	CTION			
THRUST BEARING			GLAND					HYDROSTATI	с			
PACKING BOX		1	THROA	T BUSH				RUNNING				
PEDESTAL		1	SEAL					NPSH				
MATERIALS												
CASE			PACKI	NG BOX				DIFFUSERS				
INNER CASE			THROA	T-BUSH OR	-RING			BAL DRUM				
			PACKI	NG				BAL SLEEVE				
CASE WRG-RING			CAGE	RING				BASEPLATE	BASEPLATE			
IMP WRG-RING			GLAND	AND BUSH				CASE STUDS				
SHAFT			месна	NICAL SEA	L		GLAND STUDS					
SHAFT SLEEVE			ROT	AT ING FAC	E		CASE GASKET					
SPACER SLEEVE			SIA	TIONARY F	ACE			DISCH COLUMN				
CASE BUSH			0-F	ING				SURFACE HEAD				
			AUX	GLAND				SUCT BELL				
				DE	SIGN							
NO STAGES			ROTA	TION FACE	NG CPL	G		CPLG				
MAX IMPELLER			SEAL					CPLG SIZE				
BID IMPELLER			PACK	ING SIZE				RADIAL BRO	1			
			NO O	PACKING	RINGS			THRUST BRO	3			
EYE-AREA SO IN			SHUT	OFF HD,	IM	P. PSI		COLUMN BRO				
SPEED RPM			MAX	NP	PS	IG @	F	DESIGN THE	RUST, LE	3		
DESIGN EFF			HYDR	0 TEST-PR	ESS PS	G		MAX THRUST	, LB			
DESIGN BHP			NOZZ	LE OR LENT	SIZE	RATI	NG	THRUST DIR	RECTION			
MAX BHP. BID IMPELLE	R		SUCT					WT W/O DR	VER. LE	3		
MAX BHP. MAX IMPELLE	R		DISC	н				OUTLINE				
MOTOR HP VENT CROSS SECTION												
TURBINE BHP DRAIN PERFORMANCE												
REMARKS SUPPLIER FI	LLS IN	ALL	DATA MA	RKED X								
						•						

RECI PROCATI NG PUMP ITEM NO SPECI FI CATI ON SHEET CUN 106 NO CUST JOB NO 041E 041E 87 ERVICE NO REQUIRED REVISED IFR NODEL NO SERIAL NO 0PERATING CONDITIONS PERFORMANCE 1 LIQUID GPM AT PT: NORM 41 CURVE NO.					
OCATION BY SERVICE NO REQUIRED REVISED FR NODEL NO. SERIAL NO OPERATING CONDITIONS PERFORMANCE 1 LIQUID GPM AT PT: NORM					
NO RECURED REVISEO FR MODEL NO SERIAL NO					
FR NODEL NO. Serial NO PERFORMANCE OPERATING CONDITIONS PERFORMANCE 1 LIQUID GPW AT PT: NORM 41 CURVE NO.					
OPERATING CONDITIONS PERFORMANCE 1 LIQUID GPM AT PT: NORM 41 CURVE NO.					
1 LIQUID GPM AT PT: NORM 41 CURVE NO.					
2 PT. ^o f RATED 42 DUTLINE DWG. NO.					
3 SP GR AT PT, DISCH PRESS, PSIG 43 PUMP SPEED, RPM					
4 VAP PRESS AT PT SUCT PRESS, PSIG 44 PISTON: STROKES/WIN FI	PS				
5 VIS AT PT: SSU - CP DIFF PRESS, PSI 45 EFF AT DESIGN					
6 CORR/EROS PROPERTIES DIFF HEAD, FT 46 BHP: DES MAX					
7 NPSHA, FT 47 SHUT-OFF HEAD, FT					
CONSTRUCTION 48 RELIEF VA: INT - EXT					
8 ARRANGEMENT: HORIZ - VERT					
9 DIRECT - SINGLE - DOUBLE ACTING 50 CLG WATEN, GMM					
10 SIMPLEX - DUPLEX - TRIPLEX POW.FRAME: PISTON - PLUNGER 50 DEDESTAL					
11 CASE DES PRESS, PSIG SWP, PSIG 50 DESCE DESIG TOP					
12 TAPPED CONNECTIONS: VENT - DRAIN - GAUGE 33 FRESS, FSIG 1F					
13 NOZZLES SIZE RATING FACE LOCATION 55 LARGETED YES NO					
14 SUCTION 56 STM PRESS PSIG T °F					
15 DI SCHARGE 57					
16 SHOP TESTS					
17 PUMP SIZE, IN: BORE STROKE COLOUR PUMP SIZE					
18 CYLINDER LINER 38 SHUF INSPECTION 50 DERGOMMANFE					
19 VALVES(LIU) ITPE NUMBER AREA, SULIN. OF TENTOMANUE					
20 SUCTION 00 NTURQUARTIC, TOTO/ F					
21 DISCRARGE 62					
22 DEARTAGE AND AL (1000) 23 LINEF 01 - ROFASE 01/EE: YES - NO 63 WEIGHTS:					
24 CPLS WER NORFL GUARD YES - NO 64 PUMP DRIVER					
25 PACKING BASE: MFR.STD OTHER 65 REDUCER SHPG					
LIQUID CYLINDER MATERIALS DRIVER CYLINDER MATERIALS					
26 CYLINDER VALVES CYLINDER VALVES					
27 CYL LINER PISTON ROD PISTON GASKETS					
28 LANTERN RINGS GASKETS PISTON RINGS					
29 PISTON - PLUNGER PISTON ROD					
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY	MFR TYPE				
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY 31 MFR TYPE MFR TYPE TYPE 30 ENCLOSURE RPM HP RPM MATER RATE, LB/NP-H	R				
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY 31 MFR TYPE MFR TYPE 32 ENCLOSURE RPM HP RPM WATER RATE, LB/HP-H 33 HP SF INLET STM PRESS, PSIG: NORM MAX	R				
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY 31 MFR TYPE MFR TYPE 32 ENCLOSURE RPM HP RPM WATER RATE,LB/HP-H 33 HP SF INLET STM PRESS, PSIG: NORM MAX 34 FRAME INSULATION INLET STM TEMP. OF: NORM MAX	IR				
30 Supplied by Mounted by Supplied by Mounted by 31 MFR TYPE MFR TYPE 31 MFR TYPE MFR TYPE 32 ENCLOSURE RPM HP RATER RATE,LB/NP-H 33 HP SF INLET STM PRESS,PSIG: NORM MAX 34 FRAME INSULATION INLET STM TEMP. "FISE," NORM MAX 35 VOLTS/PM /NZ TEMP RISE," FF EXALUST STM PRESS,PSIG.NORM OTHER	IR				
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY 31 NFR TYPE NFR TYPE 32 ENCLOSURE RPM NFR TYPE 33 NP SF INLET STM PRESS, PSIG: NORM NAX 34 FRAME INSULATION INLET STM TEMP. ^O F: NORM NAX 35 VOLTS/PN /NZ TEMP RISE, ^O F EXHAUST STM PRESS, PSIG: NORM OTHER 36 BEARINGS LUBE DEARINGS LUBE DEARINGS LUBE	IR				
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY 31 MFR TYPE MFR TYPE 32 ENCLOSURE PPM HP TYPE 33 HP SF INLET STM PRESS, PSIG: NORM MAX 34 FRAME INSULATION INLET STM TEMP, PF: NORM MAX 35 VDLS/PM/NZ TEMP RISE, PF EXAUST STM PRESS, PSIG: NORM OTHER 36 BEARINGS LUBE BEARINGS LUBE % PEED REBUCER: INTEGRAL SEPARATE' INCEZES SIZE RATING FACE L0	IR CATION				
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY 31 MFR TYPE MFR TYPE 32 ENCLOSURE RPM HP RPM WATER RATE_LB/MP-H 33 HP SF INLET STM PRESS, PSIG: NORM MAX 34 FRAME INSULATION INLET STM TEMP. °F: NORM MAX 35 VQLTS/PH/MZ TEMP RISE, °F EXHAUST STM PRESS, PSIG: NORM OTHER 36 BEARINGS LUBE BEARINGS LUBE DE 94 PEED REDUCER: INTEGRAL SEPARATE' NMAZZES SIZE RATING FACE LO 94 MFR MODEL EXHAUST LUBE DE DE	CATION				
30 SUPPLIED BY MOUNTED BY SUPPLIED BY MOUNTED BY 31 NFR TYPE NFR TYPE 31 NFR TYPE NFR TYPE 32 ENCLOSURE RPM NP RATER RATE, LB/NP-H. 33 HP SF INLET STM PRESS, PSIG: NORM MAX 34 FRAME INSULATION INLET STM TEMP. OFF: NORM NAX 35 VOLTS/PM /NZ TEMP RISE, OFF EXHAUST STM PRESS, PSIG: NORM OTHER 36 BEARINGS LUBE BEARINGS LUBE 36 BEARINGS LUBE BEARINGS LUBE 39 NFR MODEL EXHAUST STM INGER LD 39 NFR MODEL EXHAUST STM INGER ART DR/YER SPEC VEE 39 NFR MODEL EXHAUST STM INGER ART DR/YER SPEC VEE INGER ART DR/YER SPEC VEE 39 NFR MODEL EXHAUST STM INGER ART DR/YER SPEC VEE INGER ART DR/YER SPEC VEE	CATION S • NO				

VENDOR TO SUPPLY INFORMATION MARKED

ENG EQUIP 46 (5 77)	
FOR ROTARY PUMP	DATA SHEET
	SHEETHO
	SHEELAU
	PAGE
	N/C
OPERATING CO	ONDITIONS
PRODUCT HANDLED	DISCHARGE PRESSUREPSI(
CORROSIVE DUE TO	SUCTION PRESSURE PSIG
PUMPING TEMPERATURE DEG F VAPOR PRESSPSIA	DIFFERENTIAL PRESSUREPSI
GRAVITY AT CODEG. F. SP. GR	DESIGN HEAD
SP GP AT SUC COND SP GR AT DIS COND	NPSH* AVAIL FT N P S H REQUIRED FT
VI COSITY AT DEG F	NPSHINCLUDES
CARACITY - G B D AT 60 DEG E NORMAL	CTEAN ANTANA
	STEAM PRESSURE
	6 SATUHOR TOTAL TEMP
	EXHAUST STEAM PRESSURE
DESIGN	I MOTOR CHARACTERISTICSVOLTS PHASE CYCLE
SPECIFI	CATIONS
MANUFACTURER	HVD HP -NORMAL DESIGN
	PUNPEFF % NORMAL DESIGN
SIZE	BRAKE Hp -NORMAL DESIGN
RATED CAPACITY G P M atRPI	BH P AT RATED CAPACITY
RUTATION-COUNTER-CLOCKWISE FACING COUPLING END	MOTOR H P TURBINE H P
SUC FLANGE SIZE	DRIVER (S) TO BE FURNISHED BY
DISFLANGE SIZE	PUMP BASE
CASE DESIGN PRESS	DRIVER BASE
RELIEF VALVETO SE FURN BY	COUPLING
MAX PEL VALVE SET TO PREVENT MOTOR OVERLOAD PSIG	
N A T E R I A L S	A) DETAILS
CASE MIN THICK	LUBRICATION
END COVERS	PACKING BOX-TYPE
FLANGES	SIZEI.D
SHAFT	PACKING ARRANGEMENT
SHAFTSLEEVES	
R 0 T 0 R \$	SPARE PACKING
DRIVE GEARS	PACKING GLAND-TYPE
RADIAL SEARING	GLAND STUDS
THRUST SEARING	LANTERN RING
GENERAL	NFORMATION
COOLING WATER DEG F GPM	
PACKING BOXES GPM	INSPECTION-HVDRO TEST
COOLING OIL LANTERN RINGSDEG F G PM	PERFORM TEST
ELUSHING ON TO THROAT BUSHING	
COOLING WATER PIPING SYSTEM BY	DIMEN PRINT NO SERIAN O
IN ANI OU PIPING ST	
	CONNECTION BOLTS TO BE CHRNICHED BY VELLOCS
GLAND UL PERME BT FLUSHING OIL PIPING BY	FOUNDATION BOLTS-TO BE FURNISHED BY KELLOGG

CHEMICAL PROCESS PLANTS DEPARTMENT ROTARY PUMP CORPORATE ENGINEERING DATA SHEET Sheet 1 of 2 OPERATING CONDITIONS LUID PUMPED CAPACITY NORWAL GPM LUID CHARACTERISTICS CAPACITY DESIGN ____ G P N TEMP MAX ____OF MIN____OF NORMAL___ ___OF TOH DESIGN___ FT. FLUID VISCOSITY AT MIN. P T SUCTION PRESS DISCH, PRESS. SPECIFIC GRAVITY AT MINP T NPSH AVAILABLE FTFLUID VAPOR PRESS AT MAX P T HYDRAULIC HP TYPE OF OPERATION CONTINUOUS DINTERMITTENT HRS./DAY -DUTDOORS LOCATION. MATERIALS BODY MECH SEAL: HEADS ROTATING FACE W ROTORS STATIONARY FACE 🗕 SHAFT . METAL PARTS PACKING GLAND RELIEF VALVE LANTERN RING CASING GASKET CONSTRUCTION DATA PUMP TYPE VUMF TYPE MFR. MECH. SEAL TYPE MFR _____ CODE ž NOTES SEAL FLUID RELIEF VALVE BUILT-IN SEPARATE MOTOR DRIVE DATA TEST CERT. DATA SHOP TESTS NOT WINESSED WINESSED ITEM NO. _____ ___ MTD. 0Y __ H.P. _____ RPM ___ _ FRAME _ Running Pert. MFR. ___ CPLG. GUARD NPSH ENG ТҮРЕ ____ - INSUL -DES SUFRA ENC'L CLASS I-GROUP DIVISION 1 HYDROSTATIC 20 PSIG OTHER VOLTS/PHASE/k BEARINGS KEDBY FULL LOAD ANPS _ SF _ LR AMPS R MANUFACTURER'S DATA MER. PUMP BEARINGS MODEL NO.______ RPM _____ NOZZLES SIZE ASA RATING FACING POSITION SUCTION PUMP LUBRICATION PRESS IN SEALING CHAMBER PSIG WN BY INCHES SHAFT DIA. IN SEALING CHAMBER 🛛 WATER COOLING AVAIL AT DISCH. MAX. ALLOW. W.P. _____ PSIG PSIG BEARINGS TEM GPM B C NPSH REQ'D. (WATER) FT. GPM DESIGN EFF. _____ DESIGN BHP __ PEDESTAL GPM SHUT OFF HEAD FT TOTALWATER REQ'D. GPM ROTATION FACING CP'LG END PACKING LUBRICATION CP'LG MFR SEAL LUBRICATION Ę ТҮРЕ CALE JOB ND PLANT LOCATION PROJECT NO - REF DWG NO -DATE LDG NO TITLE POSITION NO SERIES STEP NO DRAWING E 142810* (REV. 4/83)

NO. 0 1 2 3 4 6 7 8 10 ä 9 11 12 CHECK APPRVD.

706

	CHEMICAL PROCESS PLANTS DEPARTMENT Corporate engineering	ROTARY PUMP DATA SHEET Sheet 2 of 2
	MANUFACTURER'S DATA CONT'D.	
REMARKS	VEIGHTS: PUMP EASE VEAL FLUSH SEAL QUENCH & DRAIN SPEED REDUCER MFR TYPE BASEPLATE	
	SPECIAL REQUIREMENTS & NOTES	
	VENDOR IS TO QUOTE MOST ENERGY.EFFICIENT PUMP-ORIVE COMBINATION AS A UNIT(s) OFFERED. GROUNDING LUG TO BE PROVIDED IN MOTOR CONDUIT BOX. GROUNDING LUG TO BE PROVIDED IN NOTOR CONDUIT STANDARD PROCUREMEN EIG: 1509, PROCESS ELECTRICAL EQUIPMENT - WEDIUM VOLTAGE MOTORS, ALTE EIG: 1519, PROCESS ELECTRICAL EQUIPMENT - WEDIUM VOLTAGE MOTORS, ALTE EIG: 1609, PROCESS ELECTRICAL EQUIPMENT - VARIABLE FREQUENCY AC MO' MOTOR SHALL BE PROVIDED IN ACCORDANCE WITH DATA GIVEN ON STANDARD ELECTRICAL ALTERNATING CURRENT MOTORS.	NLTERNATE IF DIFFERENT THAN NT SPECIFICATION: INATING CURRENT. TERNATING CURRENT. TORS DATA SHEET FORM E242P1
ROJ. ENG. JES SUPRV		
DATE MITI TION DHAWNBY CHECKEDBY	105	
<u> </u>	JUEno project ko. REF DWG NO.	ISSUE BLDG NO
	POSITION NO SERIES STEP NO	
		\sim

Rotary **Pump**—(continued)

KELLOGG MOTOR DRIVEN PRO			3 & F FFT	RECIPI	ROCAT	ſING			
···			_	JOB NO					
0 C A T I O N			_	ITEM NO					
			-	PAGE		OF			
SERVICE CONDITIONS			_						
LUIDPUMPED	D	ISCHARGE PI	RESSURE .					PSIG	
ORROSIVE DUE TO		INLET PRESSURE PSIG							
NLET TEMPERATURE									
SPECIFIC GRAVITY AT INLET TEMP			RIF AT TOP		P FOUNDA1	[ION		FT	
CAPACITY AT INLET TEMP DESIGN	N	PSH REQUIRE	DAT TOP O	F PUMP FO	UNDATION			FT	
		SPEC	- D.A.T	E 0.					
WECHANICAL DATA (POMPS TO BE FORMISHED IN ACCORDANCE WITH	INELLUG	ALVE WHAT							
NANUFACTURER	<u> </u>	KAKE KWNO	RMAL		DESI	5 N			
NO CYLINDERS SPW	— h	SCH_SIZE NO	RATING						
LIQUID END DESIGN PRESSURE :	PSIG ST	EAM IN SIZE							
STALLING PRESSURE	PSIG I	XHAUST SIZE							
PROPORTIONING PUMP DATA		ECIPROCATING	G PUNP D	ATA					
CAPACIN CYI A MAX MIN		TEAM PRESS	URE						
		% SATURATION	OR TOTAL	TENP .					
TYPE CAPACITY ADJ.	F	- XHAUST STEA	N PRESSU	RE					
RANGE OF ADJ	Т	OTAL STEAM	LB/HR NG	DRMAL		DES	5 I G N		
BORE CYL. ACYL. B	· · ·								
STROKE CYL. A CYL. B	s	STEAM END PACKING							
MATERIALS									
CYLINDER									
PLUNGER	- 1								
VALVE	—+:								
VALVE SEAT		PISTON RING							
VALVE SPRINGS									
MECHANICAL DETAILS	_		TYPE		NO OF	BUIKES			
COMMON BASE FOR PUMP AND MOTOR BY		PACKING BUX	SIZE		NOOF,	DD.		LENGTH	
COUPLING WITH GUARD BY									
		100% 200% SPARE PACKING (BY MFGR.)							
REVOLUTION COUNTER		□ LANTERN RING IN CENTER WITH CONN. PLUGGED							
INSULATION: STM END BYUC END BY	-								
		COOLING WATER							
MOTOR DATA (MOTOR TO BE FURNISHED IN ACCORDANCE WITH KE	LLOGO	SPEC.	DATED						
NANUEACTURER		FRAME	_						
KILOWATTSRPM	-	RATED LOAD	CURRENT						
VOLTS PHASE CYCLE		LOCKEDROTOR	CURREN	T					
ENCLOSURE		EFFICIENCY 4	4 _	3/	4	1/2 -			
SERVICE FACTOR	1	POWER FACTOR	4/4 _	3/	4	1/2 _			
TEMPERATURE RISE	C	SPEED REDUCI	ER NPE		KAI	10			
GENERAL									
APPROX BASE SIZE		TOTAL WEIGH	T 187				KG	<u></u>	
EQUIPMENT TO BESUITABLE FOR OUTDOOR INSTALLATION				SERIAL					
PNPSH NET POSITIVE SUCTION HEAD TO TOP OF PUMP FOUNDATION				a c K I A L					
NANUFACTURER SHALL SUBMIT THIS SHEET FULLY COMPLETED WIT	'HHIS I	QUOTATION							
REMARKS:				r					
NO. 0 1 2 3 4	5	6	7	8	9	10	11	12	
Ó DATE					I	L	· · · · ·		
₿Y	+						L		
CHECK							l	ł	
APPRVD	1		L	L	I	L	L	1	

PACKAGED REFRIGERATION SYSTEM	PACKAGED REFRI GERATI ON SYSTEM
Refrigerant:	Oper. Parameters LOW STAGE HIGH STAGE Units Capacity %
Design: ASME Code Section VIII, Division 1, Edition and all Mandatory Addenders.	Inlet Flow
Stamped: Yes/No, TEMA Edition 6	Inlet Press.
Item No. Service Refrig. Temp. (In/Out) Refrig. Press.(In/Out) Duty	Outlet Temp
Evaporator	RHP
0 0il Cooler 0	Pockets/Valve Open
n / / / / / / / / / / / / / / / / / / /	1 Mfr. Model No Qty
2 Temp. In/Out Total Flowrate	2 Cylinder Data: Low Stg. High Stg. Low Stg. High Stg. Low Stg. High Stg.
4 Maximum Capacity 0 RPM Uty	4 Cylinder Type Piston Speed Norm/Max.
15 Condenser (TEMA): Item No. Type Size Qty. 1	5 Cyl. Liner Yes/No Rod Diameter
6 Design Press. Shell Tubes Mat'l.: <u>Shell</u> Tubes 17 7 Cooling Medium: Flow Inlet Outlet &Press. 17	7 Bore/Stroke (In.)
18 Receiver: Item No. Type S i z e Mat'l. Qty. 14	8 Piston Displ Cyl. Press/Temp, Max
20 Knock-Out Drum: Item No. Type Size Mat 7, Oty. 20	0 No. In/Out // Hydro. Test Pressure
1 Design Press./Temp. / Oper. Press./Temp. / 22	1 Mat'1 of Const: LOW Stg. High Stg. Low Stg. High T
22 Sub-cooler: Yes/No Item No Iype Size Mat I Qty 23	3 Cylinder Liners Valve Springs
24 Oil Separator: Item No. Type Size Mat ⁻¹ . Oty.	4 Pistons
3 Design Press./Temp. // Uper.Press./Temp. // 26 26 Oil Heater: Qty. Volts Phase Hertz Watts 26	6 Piston Rods
27 Dil Cooler (TEMA):Item No. Type Size Qty. 2	7 Cooling Water: Comp. Cyl. Comp. Cyl.
9 Cooling Medium Flow Maximum △ Press. 2	9 Temp. In/Quut: / Max. Press.
30 Oil Filters: Item NoType Micron SizeQty 30	U Packing: Cooling Water; Yes/No. Vented to
32 Flow Rate: Normal Maximum △ Press Mat 1 33	2 Start/Stop; Yes/No, 2/3/5 Step Control. Suction Valve Lifters: Manual/Automatic
33 Oil Pump: Mfr Model No Type Qty 33	3 Air/Power Failure, Compressor shall Load/Unload. Mathematical Advances Gear Low Speed CPLG Belt Drive Low Speed CPLG Belt Drive
35 Purger: MfrQtyQty	5 Mfr.
36 Size Mat'I. 31 37 Meter: Compressor (Less than 200 HP) (11 Pump 3	b Model No. Size
Mctor. compressor (ress chain 200 m) orr rump 38 Mfr. /Mod. No. /Qty.	8 Ratio/Efficiency / No. Required
39 Volts/Phase/Hertz / / / 3 40 Enclosure/Insulation 4	Separate Moisture Separators with Traps: By Compressor With
Area Class. Class Groug Division /Non-Hazardous.	1 Driven Oil Pump/Hand Operated Pump for Starting.
42 Temp. Rise Cobert above Coy S.r.; Temp. Code 43	43 Electric Heater w/Thermostat: SettingNo. HeatersWatts
44 Current Effy. P.F. Full Brg. Allow-Thrust Space 4	4 Cylinder Lubrication: Lubricator to be driven by Comp. Shaft/Electric Motor
10 Item <u>HP</u> Full Locked <u>4</u> <u>β</u> <u>4</u> <u>3</u> Lbad Frame Lub. Weight Mtg. υρ/μοwn Htr. 5.r. <u>4</u> 46 No. KW Load Rotor <u>4</u> <u>4</u> <u>4</u> <u>4</u> <u>4</u> <u>4</u> <u>8</u> <u>4</u> <u>8</u> <u>8</u> <u>8</u> <u>8</u> <u>4</u> <u>4</u> <u>8</u>	46 Electric Heater w/Thermostat: Setting No. Heaters Watts
47	17 Remarks:
40 Remarks: 4	49
50 5	5 O
6 MANAGING OFFICE ENGINEERING OFFICE	5
EQUIPMENT DATA SHEET	EQUI PMENT DATA SHEET
CLIENT	
O ITEM NO. SPEC. NO.	NO DATE ENGINEER CHK/REVIEW APPROVE REO'N NO PAGE 0 F

PACKAGED REFRIGERATION SYSTEM (CONT'D)	PACKAGED REFRIGERATION SYSTEM (CONT'D)
PACKAGED REFRIGERATION SYSTEM (CONT'D) 1 POSITIVE DISPLACEMENT RUTARY COMPRESSOR 2 MTT.: Model No. 3 Speed: Max. Allow. RPM Critical RPM Ist 2nd 4 Casing: Casing split: Thickness Corr. Allow. 5 Max. Working Press. Max. Rated Press. Temp::Max. Work Min Oper. 6 Test Pressure Hydro. No. of Lobes: Material 7 Type Fabrication: Type No. of Lobes: Material 8 Type Fabrication: No. of Lobes: Material Min Oper. 9 Rotor Length to Dia. Ratio (L/D) No tor Clearance Max. Deflection Shaft: Tapered/Cylindrical 9 Rotor Length to Dia. Ratio (L/D) No tor Clearance Max. Deflection Shaft: 11 Sleeves at Shaft Seals Shaft Seal Type Seal Sys. Type 12 Inter 0il Leakage Guarantee (Vol./Day/Seal) Type Buffer Gas Tapered/Cylindrical 13 Buffer Gas Flow (Per Seal) #/Mininum Press. Press. 14 Normal #/Mininum Press. Maxer	PACKAGED REFRIGERATION SYSTEM (CONT'D) The vendor shall furnish as a minimum the equipment and components shown in the sketch and as called for on the data sheets.
Crub. nalves Keyed(1) or C or C or Hydr. CPLC 25 Spacers Required	Controls shown outside Skid Limit are by the Purchaser. Rotary Compressor Rotary Compre
46 46 46 47 48 49 50 50 1 1 2 1 1 1 1 1 2 1 1 1	Symo Size Rating Facing Weights LBS/KG a Dimensions Length Width Height Erection b Skid No. 1 Maintenance Operating c Skid No. 1 Shipping b Skid No. 1 Maintenance c Skid No. 1 Bit c Shipping Shipping Shipping c I Image: Shipping EQUIPMENT c I Image: Shipping Image: Shipping c I Image: Shipping Image: Shipping c Image: Shipping Image: Shipping Image: Shipping d Image: Shipping Image: Shipping Image: Shipping d Image: Shipping Image: Shipping Image: Shipping d Image: Shiping Imag

	PACKAGED REFRIGERATION SYSTEM (CONT'D)				
1 2	INSTRUMENTATION AND CONTROLS Panel: Freestanding/Other Covered/Uncovered				
3 4	Electrical Instruments: NEMA Enclosures Panel: Vibration Isolators/Strip Heaters/Purge/Other				
6 7	MINIMUM DISPLAY ON LOCAL PANEL MINIMUM NO. ALARMS AND SHUTDOWN ALARM SHUTDOWN				
8 9 10 11 1 2 1 3 1 4 1 5 1 6 1 7 1 8 1 9 2 0 2 1	Cube Off Temperature Gauge Fubbe Off Low Pressure Compressor Suction Pressure Gauge Compressor Low Suction Press. Compressor Discharge Pressure Gauge Compressor High Discharge Temperature Gauge Compressor Discharge Temperature Gauge System High Discharge Temp. K. 0. Drum High Level * K. 0. Drum High Level * Sub-Cooler High Hevel * High Water Temperature * Receiver High Level *				
2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	 Remarks: 1) Shutdown switches shall incorporate time delay relays, where required, to permit starting-up the system. 2) Packaged refrigeration system shall operate completely automatic after start-up. 3) Manual/automatic selector switch shall be provided, by the vendor, for manual start/stop of each refrigeration unit. 4) Condenser cooling water valve shall "fail open" and provide no less than minimum rated flow. 5) Liquid sub-cooler level control valve shall "fail open" on instrument air failure. 6) All control valves, furnished by the vendor, shall have blocks and by-passes. 7) A pair of "N.0." contacts shall be provided, by vendor, for purchasers shutdown alarm to be activated only when the system malfunctions. 8) A horn shall be mounted on the unit and wired to the shutdown circuit. 9) Copper, brass, bronze, aluminum and aluminum alloys shall not be used where exposed to ammonia vapors, ammonia solutions or the general plant atmosphere, without the specific approval of M.W. Kellogg Engineering. 10) Vendor shall include but not be limited to the following: all components, piping, valves, instruments, alarms, shutdowns, control, control panels, special tools and equipment necessary to operate the unit at its rated capacity. 11) All relief valves shall be routed to the Skid limit for purchaser connections. 				
45 46 47					
48 49 50					
00					
	2 CLIENT VCATION JOB NO.				
	0 11EW K0 SPEC. NO. 0 DATE ENGINEER CHK/REVIEW APPROVE REC'N,NO PAGE OF				

PROCESS DESIGN SECTION VIBRATING FEEDER CORPORATE ENGINEERING DATA SHEET FEED MATERIAL CHARACTERISTICS FEED MATERIAL. WEIGHT 🗙 NOISTURE. oF SULK DENSITY: LBS./CU.FT. TEMPERATURE. PARTICLE SIZE: AVERAGE: A B R A S I V E N E S S : MAXIMUM: TENDENCY TO PACK: PRODUCT DESCRIPTION: DESCRIPTION: OPERATING CONDITIONS о и т в о о к 🛛 LOCATION: INDOOR FEED RATE LBS/HR, NORMAL MAX. ROOM TEMP: _ MAX. REQ'D, WIDTH OF FEED INLET TO OUTLET LENGTH: DUSTY 🛛 ATNOSPHERIC CONDITIONS: REQ'DDEPTHOFFEED CORROSIVE EXPLOSIVE NODE OF OPERATION OTHER ____ DATE DESIGN DATA 4 TROUGH TYPE. COVER CLAMPS: LENGTH/WIDTH/DEPTH FREQUENCY CPM: Å, THICKNESS: _ - SPEED OF MAT'L, CONVEYED-FPM NATURAL FREQUENCY: REO'D: _ COVER: REQ'D, SPRINGS: TYPE MOUNTING: FLOOR-SUSPENSION AMPLIFICATION INCLINE _____ DECLINE __ ISOLATION _ REV CONTROL . HANGERS- TYPE: BASE _ _NO. MATERIALS OF CONSTRUCTION TROUGH_ SPRINGS - AMPLIFICATION _ DES SUPRV COVER _ ISOLATION _ BASE _ DRIVE COVER _ GASKETS MOTOR DRIVE DATA CHECKED BY _LUBE _ ITEM NO. MTD. BY BEARINGS FULL LOAD AMPS SF MFG. _ L.R. AMPS FRAME TYPE_ RPM HP___ воттом 🗆 OVERHEAD MOUNTING: VOLTS/PHASE/CYCLE DRAWN BY MISCELLANEOUS TOTAL WEIGHT VENDOR DWG, NO. PAINTING FEEDER SERIAL NO. GALVANIZING NITIATOR FINISH REQ'D. SCALE PLANT LOCATION REF DWG NO FROJECT NO ISSUE TITLE DATE BLDG NO E227B43 SERIES OSITIONNO. STEP NO RAWINGNO

Packaged Refrigeration System-(continued)

	FEED MATERIA	L CHARACTERISTICS
ARKS	FEEDWATERIAL BULK DENSITY: IRS / CIL FT	
ĒM	PARTICLE SIZE	DUSTINESS
۳.	MAXIMUM SIZE	CHENICAL ACTION
	ANGLE OF REPOSE PRODUCT DESCRIPTION	¥F¥F¥F
 2	OPERATIN	G CONDITIONS
H d	CAPACITY: LBS./HR. NORMAL	LOADING CLASSIFICATION
<	MAX	LOCATION: INDOOR OUTDOOR
DATE	CONSTRUCTION	
VHAD		
ġ-	MAT'L.	TROUGHEND
Ē	THICKNESS	TYPEHANGERS
<u>ک</u>	COVER: TYPE	SADDLES
	MATTL. Thickness	BEARINGS: TYPE + HARD IRON
ώ	FLIGHT: PITCH	
œ		BABBILED
<u>«</u>	WAT'L.	BRONZE
¥	NAT'L	ANTI-FRICTION
	NAT'L. THICKNESS FLATVALVEPLATEGATE RACK ĜIPINION GATE → FLAT VALVE PLATE	ANTI-FRICTION
4	HATTL FLATVALVEPLATEGATE RACK & PINION GATE CURVED VALVE PLATE	BRONZE
8	HAT'L. THICKNESS FLATVALVEPLATEGATE RACK & PINION GATE	ANTI-FRICTION
<u>د</u>	HATYL	MOTOR DATA
8	HATUL	MOTOR DATA
8	HATTL	MOTOR DATA ITEW NOMMDTOR DATA ITEW NOERANEE
A A	HATTL	BRONZE ANTI-FRICTION JACKETEDTROUGH MOTOR DATA ITEN NO. MTD. BY ITEN NO. FRAME ITEN NO. VOLTS/PHASE/CYCLE
	HAT'L. THICKRESS FLATVALVEPLATEGATE RACK & PINION GATE • FLAT VALVE PLATE CURVED VALVE PLATE CURVED VALVE PLATE DRIVE DATA CHAIN-ASANO. SPROKETS • DRIVER NO. TEETH V-BELT NO. & SIZE SHEAVES DRIVER-PITCH DIA DRIVER-PITCH DIA	BRONZE
	HAT'L. THICKNESS FLATVALVEPLATEGATE RACK & PINION GATE • FLAT VALVE PLATE CURVED VALVE PLATE DRIVE DATA CHAIN-ASANO. SPROKETS • DRIVER NO. TEETH V-BELT NO. & SIZE SHEAVES DRIVER-PITCH DIA REDUCER = MFGR.	BRONZE
	HAT'L	MOTOR DATA JACKETEDTROUGH MOTOR DATA ITEW NO. MTD. BY BY WFG. FRAME BY VOLTS/PHASE/CYCLE / JACKETEDTROUGH P ERAINES LUBE FULLIONANPS S F
	HAT'L	MOTOR DATA JACKETEDTROUGH MOTOR DATA ITER NO. MTD. BY BY #FG. FRATE FRATE #FG. FRATE BY
4	HAT'L. THICKNESS FLATVALVEPLATEGATE RACK & PINION GATE - FLAT VALVE PLATE CURVED VALVE PLATE DRIVE DATA CHAIN-ASA NO SPROKETS - DRIVER NO. TEETH V-BELT NO. & SIZE SHEAVES DRIVER-PITCH DIA DRIVEN-PITCH DIA REDUCER = MFGR NO BELNO SERVICE FACTOR INOU ELNO SERVICE FACTOR OUTPUT RPM	BRONZE
4	HAT'L	BRONZE ANTI-FRICTION JACKETEDTROUGH MOTOR DATA ITEW NO. MTD. BY WFG. TTPE RAWE NTJ./PLASE/CYCLE VOLTS/PHASE/CYCLE FULLIOADAWPS S FULLIOADAWPS S RAWGE
4	HAT'L	BRONZE ANTI-FRICTION JACKETEDTROUGH ANTI-FRICTION JACKETEDTROUGH MOTOR DATA ITEN NO. MTD. BY NTD. BY VOLTS/PHASE/CYCLE / BEARINGS LUBE FULLIOADANDS F RANGE RANGE
4	HATTL	BRONZE ANTI-FRICTION JACKETEDTROUGH MOTOR DATA ITEN NO. MTD. BY WFG. FRANE NTD. BY WTD. BY WFG. FRANE NTD. BY Prove the second sec
	HAT'L. THICKRESS FLATVALUEPLATEGATE RACK & PINION GATE • FLAT VALVE PLATE CURVED VALVE PLATE DRIVE DATA CHAIN-ASANO. CHAIN-ASANO. CHAIN-ASANO. CHAIN-ASANO. CHAIN-ASANO. CHAIN-ASANO. DRIVER NO. TEETH V-BELT NO. & SIZE SHEAVES DRIVER-NOTCH DIA. DRIVEN-PITCH DIA. DRIVEN-PITCH DIA. REDUCER • MEGR. WORELNO SERVICE FACTOR INVIT RPM OUTPUT RPM OUTPUT RPM COUPLING MEG'R. TYPE	BRONZE ANTI-FRICTION JACKETEDTROUGH ANTI-FRICTION JACKETEDTROUGH MOTOR DATA ITEN NO. MTD. BY WFG. FRANE NTD. BY NTD. BY WFG. RPN VOLTS/PHASE/CYCLE / BEARINGS LUBE FULLIOADANPS S RANGE SCALE
	HAT'L	MOTOR DATA JACKETEDTROUGH MOTOR DATA ITEN NO. MTD. BY MTD. BY WTP FRATE HTG. TTPE FRATE HTG. SEARINGS LUBE FRATE FULLLOADANDS S F RANGE SCALE SCALE
	HAT'L	BRONZE ANTI-FRICTION JACKETEDTROUGH ANTI-FRICTION JACKETEDTROUGH MOTOR DATA ITEW NO. MTD. BY #F6. FRAME TTPE FRAME #F0. FRAME YOLTS/PHASE/CYCLE / F81110010AMPS LUBE F91110010AMPS F LRAMPS F RAMOE SCALE
	HAT'L	BRONZE ANTI-FRICTION JACKETEDTROUGH ANTI-FRICTION MOTOR DATA MTD. BY #F6. FRAME TTPE FRAME #70. VOLTS/PHASE/CYCLE VOLTS/PHASE/CYCLE / BERATINGS LUBE FULLIONDAMPS F LRAMPS F LRAMPS F STEP NO SCALE

MISCE LANEOUS	
99 PAINTING:	
VENOOR DWG. NO CONVEYOR SERIAL NO FINSH OF INTERNAL PARTS:	
REMARKS	
08 NO PLANT LOCATION 10/15/1 NO. REF DWG. NO. ISSUE	E
	N O .

	PROCESS DESIGN SECTION CORPORATE ENGINEERING	BUCKET ELEVATOR DATA SHEET			
	OPERATING	CONDITIONS			
MARKS	MATERIAL CONVEYED:	WOISTURE CONTENT:WGT.% CAPACITY: TONSIHR. NORMAL MAX MODE OF OFERATION:			
RE I	BULK DENSITY.	BUCKET SPEED: FPM			
1	SCREEN ANALYSIS: %				
ĺ					
2	TYPE: CENTRIFUGAL CONTINUOUS	BELTCARRIER FOR BUCKETS:			
APP	dASING CROSS SECTION DIMENSION:	WT. OF <u>DUCK:</u> COVER THI <u>CKNESS</u>			
1	CASING GAUGE	TYPE OF BELT SPLICE:			
Ā	CASING HD. SECT GAUGE OISCH. OPENING: BOOT -TYPE:	TAIL PULLEY DIA: FACE WIDTH:			
- 2	G A U G E :	HEAD SHAFT DIA: TAIL SHAFT DIA:			
5	LOADING OPENINGS	TYPE OF BEARINGS:			
Ę	CHAIN CARRIER FOR BUCKETS	HEAD SHAFT SPEED:RPM			
<u> </u>	HEADSHAFTSPROCKET P.D :	LOCATIONS:			
2	TAIL SHAFT P.D:	INLET CHUTE ANGLE:			
-	SIZE OF BUCKETS:	DISCHCHUTEANGLE:			
1	BUCKET BOLTS-TYPE:	INLET OPENING W/LOADING LEG.			
	NO. PER BUCKET:				
	MATERIALS OF CONSTRUCTION				
, H	CASING:	CHAIN:			
S,	HEAD SECTION:	HEAD 🌡 TAIL SPROCKETS:			
UES	B00T:				
>	HEAD & TAIL SHAFTS:	BUCKET FILLER PIECES.			
D,	HEAD 🌡 TAIL SHAFT BEARINGS:	GASKETS:			
CKP	BELT & COVERS:	JOINTS.			
5	HEAD & TAIL PULLEYS.	WORS & MANWAYS:			
ž	MOTOR	DRIVE DATA			
7	ITEN NOMTD BY	L R A M P S			
		REDUCTION UNIT			
	TYPEINSUL.				
	ENCLOSURE CLASSIGROUP D	NFG			
	VOLTS/PHASE/CYCLE/ TYPE DRI	IVE CHAIN LIBELT L			
	FULLLOAD AMPS S.F	NFG.			
)BO PLANT LOCATI	ION SCALE			
	REF DWG NO				
B41		DATE BLDG, NO.			
225	SSITION NO. SERIES	STEPNO			
щ		BRAWING NO.			

		PROCESS DESIGN SECTION CORPORATE ENGINEERING	PRESSURE LEAF FILTER DATA SHEET	
		OPERATING	CONDITIONS	
	IARKS	FLUID: CHARACTERISTICS:	WORKING PRESSPSIG Allowable ApmaxPSIG	
	BEN	SPECIFIC GRAVITY:	VASH WEDIA:	
		TEWPERATURE:OF VISCOSITY:CP SOLIDS CONTENT:WT.%	TYPE OPERATION:	
		ŚLOW RATE::		
	.H.dd	MECHANIC	CAL DATA	
	DATE /	NFR. SIZE & TYPE	TANK: VERTICAL HORIZONTAL D	
	SUPRV.	NO. OF LEAVESSPACING Bubble Ring	CARRIAGE DRIVE	
	СНКО	LAKE CAPACITY	LEAF SPACERS VIBRATOR	
	ВY	SLUICER DRIVE JACKETFORPSIG	CONNECTIONS-TYPE	
$\overline{\diamond}$	N N	MATE	RIALS	
		TANK:	O-RING LEAF GASKETS	
4		SURFACEMEMBER	JACKETEXTERIOREXTERIOR	
ES SUF		ACCES	SSORIES	
CHECKED BY D		PRE-COAT OR SLURRY TANK: VERTICAL HORIZONTAL DIAM. <u>y o L U</u> m e MATERIAL FILISH	PRECONTOR FILTER FEED PUMP: CAPACITY: ATERIAL: WATERIAL: SIZERTYPE:	
DRAWN BY		LINING COVER TYPE LEVEL INDICATOR: YES D NO D AGITATOR & P N	VOLTS/PHASE/CYCLE	
INITIATOR		NOTE: ALL MOTORS TO BE CLASS +GROUP-O EXP. PROOF	II F R	
		DBNO PLANT LOCATI ROJECT.NO REF DWG. NO. IYLE		
	21837	OSITION NO. SERIES	STEPNO DATE DLUG. NU.	
	E2		DRAWING NO.	

T	— T	PROFESS RESI ON CROTTON	
INTAKE AIR FILTER T		PROCESS DESIGN SECTION	LINE FILTER
		CURPURATE ENGINEERING	SPECIFICATION SHEET 1 OF
Indoor/Outdoor; English/Metric Units			
s Site: Alt. Amb. Temp.: Max. Min. : Wind Velocity: Max			
Unusual Conditions: Dust/Fumes/Sand/Snow/Other	KS ST	1. FUNCTION.	
5 Air Flow: Normal Rated @ Temp. Press.	E MA		
6 AFI Dust Spot Efficiency (Atmospheric Dust):%.	× ×		
7 Unit Shall Remove of all Particles Microns & Larger.			
8 Rated Data: Stage No. 1 Stage No. 2			
9 Filter Type			
10 model no.			
2 Pressure Drop (Clean Filter)			
13 Max. Press. Drop (Dirty Filter)	2		
14 No. of Elements	APT		
15 Size of each Element	w	2. SERVICE CONDITIONS	
16 Filtration Area	DAT		
1/ Fittation material		FLUID FLUWING:	
19 Dust Holding Capacity	a la	FLOW RATE: NORMAL	GPM
20 Weight of Element	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	мах	GPM
21 Element Retention Method in Frame	CHK	178 17K.	
It Louvers Filter Inlet: Size Lype Material 9t Inlet: Size Winner Countrol Metanial	<u> </u>	PRESSURE.	PSIG
24 Filter Housino: Material WIFE <u>Gauge</u> Material 24 Filter Housino: Material Internal/External Paint	á	TEMPERATURE:	•F
25 Access Doors: Qty Blow in Doors: Qty Filter Outlet: Side/Bottom/Back Conn.			
26 Motor: Mfr Model No Oty.	✓ ^z	FLUID VISCOSITY:	GP
27 Area ClassGrDiv/Non-Hazardous; VoltsPhase Hertz		FLUID SPECIFIC GRAVITY:	
28 EnclosureInsulation; lerb. RiseCaboye by eS.F.		PARTICLE SLZE ANALYSIS	
29 Service We Eurient Erry, P.F. Full Brg. Allow, Infust Space		TARTCEL SIZE ANALISIS	
31 R/KG Watch 4 4 4 4 4 RPM No	ų v		
	SC E		
3: Diff. Press. Switch: Range Set Press Qty.	RO SES	SOLUDS LOADING.	
34 Diff. Press. Ind. Type		SOLIDS LOADING.	
35 Specifications: Paint WindM.W.K	λa a	FILTER PRESSURE DROP:	PSIG
77 Notes: 1) The structure, consisting of the plenum chamber & filter, shall be of such	8	FILTER RATING:	
38 design that it can be supported by means of four (4) steel columns; one (1) at each	1EC		
39 corner. 2) Plenum chamber shall be of sufficient strength & bracing to prevent wall,	<u> </u>		
40 floor or ceiling from pulsating or oscillating. 3) For ammonia/urea plants or other		3. TYPE AND DESCRIPTION:	
11 installations with annionia in the atmosonere, no copper, atum num or their diloys 1 A2 shall be used for parts evolved to the air A) Seals & filter material shall be	β		
43) resistant to vapors present in the inlet air (ammonia, hydrocarbons, etc.). 5) Unit	YW1		
44 shall be completely assembled and shipped as one unit. 6) Instruments & controls to	RO		
45 be completely piped and wired. 7) Outlet connection screen (0.25 in.) (6 mm) wire			
40 mesn, with back up bars to withstand Seven (7) psi/0.5 kg/cm2 △ Press. across the	<u>e</u>	——————————————————————————————————————	
48	2 E		
49	DA IN		
50		08 NO.	PLANT LOCATION
MANAGINGOFFICE ENGINEERING OFFICE		ROJECTNO.	REF. DWG, HO
5		ITLE	DATE BLDG. no.
9 4 EQUIPMENT DATA SHEET		POSITION HO.	SERIESS T E P N O .
			DRAWING NO.
осатіонJOB NOJOB NO			, KA
0 ITEN NO SPEC NO.	E 121 85*		
NO DATE ENGINEER CHK/REVIEW APPROVE REC'N NO PAGE OF			

713

	PROCESS DESIGN SECTION LINE FILTER CORPORATE ENGINEERING SPECIFICATION SHEET 2 C	DF	PROCESS DESIGN SECTION CORPORATE ENGINEERING	CENTRIFUGE DATA SHEET SHEET 1 OF 2
REMARKS	4. <u>MECHANICAL</u> MATERIALS OF CONSTRUCTION HOUSING:	REMARKS	OPERATING FEED MATERIAL CHARACTERISTICS DENSITY OF SLURRY: LB./CU. FT. PERCENT SOLIDS. VISCOSITY OF SLURRY: CP FEED RATE. NORM. MAX.	CONDITIONS CAKE CAPACITY: CU. FT. CAKE DENSITY: LB/CU. FT. CAKE MOISTURE % SOLIDS PARTICLE SIZE: MAT'L
- A		<u>بر</u> ۲	SG. SOLIDS SG. L I <u>Q U I D</u> MECHANI	REQ'D, CAPACITY:LB/HH CAL DATA
CHKD SUPRY. DATE APP	GASKE 15:	CHKD BUNN DATE APP	MFRBASKET DIAMBASKET DIAMBASKET DEPTHBASKET RING OPENING DIAMCARE THICKNESSCARACTLY UNDER TOO DINGCU ET	CONNECTIONS SIZE TYPE FEED PIPE SPRAY PIPE ALGART GLASS SIGHT GLASS VERT
REV BY	DESI GN_PRESSURE: PSIG DESI GN_TEMPERATURE:		BASKET RPM DRIVETYPE. ELECTRIC HYDRAULIC COVERTYPE. FULL FUME TIGHTNON-FUME TIGHT MATE	
CHECKED BY DES SUPRV.			CASE	FEED PIPE SPRAY PIPE DRAIN NOZZLE VENTNOZZLE VENTNOZZLE PURGE NOZZLE UNLOADING PLOW SXMMING TUBE FILTER CLOTH GASKETS GASKETS
ATOR DRAWN BY	A,S.ME code. required when filter is larger than 6" IN diameter or holds more than 5 cu. ft. in volume, or if design pressure is greater than 15 p.s.i.g.		ACCES AUTOMATIC CENTRIFUGING CYCLE CONTROLS (SEE REMARKS COLUMN FOR PROPOSED CYCLE) CONTROL PANEL	SSORIES
	JOB NO. PLANT LOCATION SCALE PROJECT NO. REF. DWG. NO. ISSUE TITLE DATE BLDG. NO. POSITION NO. SERIES STEP NO. DRAWING NO.	E222938	08 NO PLANT LOCATI ROJECT NO REF DWG. ND ITLE '0\$ITI0N NO SERIES	STEP NO DRAWING NO.

Line Filter Specification-(continued)

714



		PROCESS DESIGN SECTION CORPORATE ENGINEERING	VIBRATING SCREEN DATA SHEET SHEET 1 OF 2				
		FEED MATERIAL	CHARACTERISTICS				
	REMARKS	FEED MATERIALLBS./CU, FT. BULK DENSITYLBS./CU, FT. ANGLE OF REPORT WEIGHT % HOISTURE PRODUCT DBSCRIPTION	ABRASIVENESS				
	2	WATERIALTOSESCREENEDOUT PART. SIZE ON IST SCREEN,% RETAINED PART. SIZE ON ZND SCREEN/% RETAINED PART. SIZE ON ZND SCREEN/% RETAINED					
	APP'R	OPERATING CONDITIONS					
	SUPAV. DATE	FEEDRATE	SUCTION IN DISCH. LINE PRESS. IN DISCH. LINE LOCATION INDOOR OUTDOOR O NODE OF OPERATION:				
	СНКD	DESIG	N DATA				
$\overline{\bigcirc}$	REV BY	NO. OF SCREENS	DISCH. SIZE TABLE FEED DISTRIBUTOR REC'D , SCREEN ANGLE: DESIGN				
D BY DES SUPAV.		ARDERFERNESS IST SCREEN DISCN. SIZE 2ND SCREEN DISCN. SIZE SCREEN AREA FREE/GROSS IST SCREEN SQ. FT. SCREEN AREA FREE/GROSS 2ND SCREEN SQ. FT. SCREEN AREA FREE/GROSS 3ND SCREEN SQ. FT. SCREEN BACKING REQ'D./TYPE IST SCREEN SCREEN BACKING REQ'D./TYPE IST SCREEN SCREEN BACKING REQ'D./TYPE IST SCREEN SCREEN SCREEN BACKING REQ'D./TYPE IST SCREEN	OVER REC'D				
ECKE		MATERIALS OF CONSTRUCTION					
ИПЛАТОН ОНАМИВУ СН		SCREEN DECKS	HOUSING				
		MISC:					
	3842	JOB 0 PLANT LOCATI PROJECTNO REF DWG NO TITLE POSITIONNO. SEPTRE	STEP NO.				
	E22(





716

S&WFORM B-1347	S & W FORM B-1347						
0 W II 1 0 101 # 1011	STONE & WEBSTER ENGINEERING CORPORATION						
	BUBBLE TOWER					Rev. 1	
			-			Rev. 2	
1 Client		Location			J. O. No.		
2 Apparatus					Project No.		
°	Date By						
Based on					Rating Page	Dated	
		0 D II-/-	DE	SIGN CONDI			Ver No.
•		Deer Pr ID/q	in ga Dor T	lemp	• r <u>H</u> •₽ Fie	ld Hyd Tost	Yes No
		Wind Proi Area	h/an ft Self S	upporting	Yes No Oth	er Tests	
•		Max. Horizontal Defle	etion at Top 1	Fray	10		
10	Earthquake 0.2 Operating Weight applied at e.g. Yes so						
11			МЕ	CHANICAL	DATA		
12		Code	Stress Reliev	e Yes No	Shell & Head	s ASTM Spec	
18			Radiograph	Yes No	Type of Hea	ds	x , , , , , ,
14		THICKNESS:	Calculated_	Corrosion	Total Thick	Liner Mat	Liner Thick
15		Shell	110	111	10		in
18			10 in	in	in		in
18		Top Head	in	in	in		in
19		Intermediate Head	in	in	in		in
80			in	in	in		in
\$ 1		Bottom Head	in	in	in		in
39		Cone Section	in	in	in		in
18		Skirt	in	in	in		in
36 az		Insulation Stiffoner Bings	Number Rec	21 Bire	[111] in X	in	
36		NOZZLES:	Number	Туре	Series	Size	Mark No.
\$7		Feed				in	
28		O H Vapor				in	
29		Beflux In				in	
30		Reflux Out				in in	
**		Bottoms Design				in	
33		Behailer Vapor				in	
34		Reboiler Liquid				in	
\$5						in	
\$6						in	
\$7		Safety Valve				in	
38		Manholes				in	
89		Thermocouples				in	
40			· · · · · · · · · · · · · · · · · · ·			in in	
47		Gage Glass				in	
45		Pr Gage				in	
44		LevelControl		İ	1 1	in	
45						in	
46		Vent				in /	
47		steam out TRAVS.			CAPS	ш	
48		Trov No Diamate	er Spacing	Type Ma	teriala Size	Туре	Materials
50 1 1 1 1			, 7 ,	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	ł	Ţ	1
51 Estimated Weight	lb						
11				1		1	
58		Baffles, etc.					Sold
54 Insulation Supports	Yes No: No	umber Req'd	Size	in X	in: S	nop l Vd Show	Field
Platform clips: Yes	No: Number Req'd	Shop Field	Ladder Clips:	I CE NO:	. amoer neg	u Snop	ricia
57 For Funther Details	Ireq For All Covers Wei	gning Over 15 LD					
58 Remarks:	INCICI IO.						
59 Copy To		Date	Thecked	Date	Approve	d D	ate



THE FLUOR CORPORATION LTD.

TYPE I

VESSEL DAVIT

wr.

DRG.NO.

м =

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PSI 🚯

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K HEADS

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PSI

PSI

PSI

LBS.

LBS.

LBS.

LBS.

LBS.

FT. LBS.

FT. LBS.



	GLASS GLASS			GLASS LIN	D VESSEL
		CORPORA		DATA SHEET	Sheet 1 of 2
		<	Reactor]; Tank]; Other		
	2	RAL	Vertical 🛄 ; Horizontal 🛄 ; Jacketed 🛄 ; Unjecked 🛄		
	NAMA IN	ENE	Rated capacity, gais; Flanged head []; One piece construction [].		
	Ĩ	ves ves	Standard Head Openings []; Non standard head required. [_] Specification Based on Pfaudler Series or equal.		
		 			
			Manhole 🔄 ar Handhole 💭 with 4" sightglass (``); Agitator Opening (``).		
		HE S	Size "for 4" light glass Size "for		
7		N L	Size ** for Size ** for		
			Size		
·	۲	вотто	M HEAD Opening		
	ă	<u>تا ۲</u>	Shell Openings, Standard 🗍 : Special		
	MPN.	ACK	Jacket Head Openings, Standard []; Special		
	1ª		Orientation of Process openings - By yender []: HLR approval of Venders dwo. re	etd ()	
	<u>–†</u>	10	Other	_	
	à	T N	Orientation of Jacket Openings; By Vendor []; HLR approval of Vendors dwg. req ⁴	d. []	
$\overline{\mathbf{C}}$	2	l ä	Other	-	
\sim	1		Working Conditions: Water []; Steam]; Brine]; Hat Oil];		
		KET	Other		
		۲ ۲	Working temperature: °F to °F		
LNG.	PRV.		Weshing Procession: Attorphonis []; prig: Vacuum []]		
. TO	3		Leg Couplings: _ Size ''; Number		
	Ē	ORT	Vendor to supply: - Adjusting Nuts and Pedistols []; Legs [])		
	6	L SUF	Distance from face of bottor		
	HECK	SSEI	Position: — On ę of manhale 🛄 ; 45º off ę Manhale 🛄 ;' below face of flu	an ge;	
	<u> </u>		" abave face of battom outlet.		
	Ł	┣───	Vessel weight if special Ibs.		
	R.	۰ م	'' x'' glass-lined flush battom valve 🛄		
	280	DRIE -	Reducing flanges:' x' ; [] '' x' x	[] " • _	"·
		1-55	Others:		
w	to to	ACC			
UVU	i i				
		JOB NO.			SCALE
		"Im u	ад <u>ана (ст. Сид</u> ио.		BLDG. NO.
		POSITION			
		· ·	DRAW	VING NO.	
			-		K X



		PROCESS DESIGN SECTION CORPORATE ENGINEERING	FRACTIONATOR TRAY DESIGN		
		REBOILER TYPE (KETTLE OR THERMOSYPHON) TRAYSNUMBEREDFROMTOPOR BOTTOM	SERVICE		
	ARKS				
	REM	TOTAL TRAYS IN SECTION			
		TRAY SPACING (INCHES) (SEE NOTE			
		NO. OF LIQUID PASSES (SEE NOTE)			
	Ĩ	MAX. ΔP () /TRAY TRAY EFFICIENCY (SEE NOTE 1)			
		ENTRAINMENT (SEE NOTE)			
	+	INTERNAL CONDITIONS			
	P.R.	VAPORTOTRAY MAX.			
	AP	RATE (#/HR.) MIN.			
	3	PRESSURE (PSLA)			
	DAT	TEMPERATURE (°F)			
	-				
	SUPR	LIQUID FROM TRAY MAX.			
-	e	RAIE (#/HK) MIN. DENSITY (* ETI			
	5	TEMPERATURE (°F)			
	~	VISCOSITY, CP			
_		SURFACE TENSION (DYNES/CM)			
\sim	5	FOAMING TENDENCY			
\leq	1 °	MECHANICA	. DATA		
		TOWERMANHOLE <u>I.D. (INC</u> HES) (NOTE 2)	TYPE OF TRAY INSTALLATION:		
		DECK	BOLTED		
-		C A P	HATCHWAY		
	PRV.	HOLDOWN			
	SUI	NUTS & BOLTS			
	DES	DECK THICKNESS (GAGE). (NOTE 1)			
-		SUPPORT RING WIDTH & THK. (INCHES) (NOTE		
	β	DOWNCOMER BOLT BAR THK. (INCHES) (NOTE)			
	9	CORROSION ALLOWANCE			
	HEC	TRAYS (INCHES)			
	U.	TRAYS INSTALLED FROM TOP OF BOTTOM			
	78 7	NOTES: I. VENDOR SHALL FILL IN ALL BLANKS AND RE	TURN WITH QUOTATION.		
	W.Y.	2. SMALLEST I.D. THROUGH WHICH TRAY PARTS	MUST PASS.		
	ő	3. WEIR ADJUSTABILITY OF ± 1/2" REQUIRED.			
		4. VENDORS GUARANTEE: THE VENDOR SHALL BE EXPECTED TO GUARANTEE HIS DESIGN:			
	B	DOWNCOMER AREA, HE SHOULD SUBMIT A DE	SIGN WHICH HE CAN GUARANTEE FOR THE		
	TIAT	SPECIFIED SERVICE.			
	IN.				
		JOB NO. PLANT LO	CATION		
		PROJECT NOREF DWG.			
		ITLE	DATE BLDG. NO.		
		POSITION NO SERIES	STEP NO.		
		OLAILO			
			L L		
144	B12*		N.		

Glass Lined **Vessel**—(*continued*)

CATALYTIC CONSTRUCTION COMPANY

•	+50/R00×05/	12-16-20	

	I I.D. VESSEL			
	2 ITEM HO.			
	3 DESIGN TENP. °F			
	41.D. VESSEL MANHOLE	-		
	5 NO. & TYPE TRAYS			
	G TRAY MATERIAL			
I I E	7 TRAY CORROSION ALLOW. (Total)		 	
	P D'CH'R AREA-TOP Sa Inches		 	
-	IO OUTLET WEIR PERIPHERY		 	
	IL QUITLET WEIR HEIGHT			
	2 OUTLET WEIR ADJUSTMENT			
	3 OUTLET WEIRNOTCH			
	WINLET WEIR HEIGHT			
	IS TRAY MANWAY ¹¹	-		
	6 TRAY SPACING	-		
v ≠	7 D'CH'R CLEARANCE			
·	ISSUMP FORAMOFFREQUIREMENTS			
Ī	9 NO. CAPS PER TRAY			
	20 CAP & RISER MATERIAL			
	21 CAP & RISER ASSY, TYPF			
2	22 SIZE OF CAPS			
	73 CAP SPACING			
2	24 CAP SKIRT CLEARANCE			
	25 C A P SLOTSPEC.			
2	26 TOP OF SLOT ABOVE TRAY			
2	2 7 TYPEO F SLOTINCAP			
	28 I.D. & NEIGHT OF RISERS			
2	9 SIZE OF TRAY SUPPORT BAR			
	30 TRAY SUPPORT . MATERIAL			

	PROCESS DESIGN SECTION CORPORATE ENGINEERING	TOWER INTERNALS DATA SHEET
BPRV REAARS	DISTRIBUTOR TOWER INSIDE DIA MATERIAL CORROSION ALLOWANCE LIQUID FLOW-NORMAL LIQUID DENSITY DESIGN TEMP°F or°C MIN. DIA. OF ACCESS SPECIAL REQUESTS	SUPPORT PLATE TOWER INSIDE DIA. MATERIAL CORROSIN ALLOWANCE DESIGN LOAD (LIQUID) LB/SQ. FT/HI DESIGN LOAD (VAPOR) LB/SQ. FT/HI DESIGN TEMP. PACKING SIZE & TYPE PACKING SIZE & TYPE PACKING DEPTH SPECIAL REQUESTS
REV BY CHKO SUMMY D	HOLD-DOWN PLATE TOWER INSIDE DIA. MATERIAL CORROSION ALLOWANCE DESIGN TEMP. °E Or °C MN. DIA. OF ACCESS PACKING SIZE & TYPE SPECIAL REQUESTS	BED LIMITER TOWER INSIDE DIA. MATERIAL CORROSION ALLOWANCE DESIGN TEMP. DINN. DIA. OF ACCESS PACKING SIZE &TYPE: SPECIAL REOUESTS
PROLENG.	COLLECTOR PLATE TOWER INSIDE DIA	REDISTRIBUTOR TOWER INSIDE DIA. MATERIAL CORROSION ALLOWANCE LIQUID FLOW-NORMAL MAX. MAX. MAX. MAX. MAX. MAX. MAX. MIN. LIQUID DENSITY SUPPORT LEDGE WIDE x THM DESIGN TEMP. °F or MN. DIA. OF ACCESS SPECIAL REQUESTS
		I
DATE		
DATE	JOB NOPLANT LOCAT PROJECT NOREF DWG NO. TITLE	
DATE	JOB NO PLANT LOCA' PROJECT NO.	E S S T E P N O

Service: Indoor/Outdoor: English/Metric Unit Type of Water to be Chlorinated: Act/Stance Act/Stance Service: Indoor/Outdoor: English/Metric Chlorine: Liquid/Gas: Desage: Ret/Scnedule Service: Indoor/Outdoor: English/Metric Very or finite in pressure Chlorine in pressure Chlorine in pressure Service: Service: Indoor/Outdoor: English/Metric Win: Material Material Material Material Material Material Wr. Material Material Model No. Temp Year Mol. %: Vapor Capacity: Mol. Wt. Mol. % Nmr. Material Model No. Chlor./Mater Out Sp. Gr. 0 Oper. Press. Vapor Capacity: Mol. Wt. Mol. %: Mol. %: Mir. Mir. Material Mol. No. Sp. Gr. 0 Oper. Press. Solids %: Pert. Size @ Oper. Nol. %: Mol. %: Vapor Capacity: Mol. Mt. Mol. Mt. Mol. %: Wapor Capacity: Mol. %: Mol. %: Sp. Gr. 0 Oper. Press. Witer In Chlor./Material Mol. %:
Type of there to be chlorinated: Induity dubled, injection 2017. Type of there to be chlorinated: Model No. Reg. Chlorine in Pressure Currinator: Jeen No. QUY Capacity: Chlorine in No. QUY. Wir. Material Wir. Material Wir. Material Mir. Max./Shock Flow: Chlorine in No. QUY. QUY. Waterial Mir. Max./Shock Flow: Chlorine in No. QUY. Wir. Material Mir. Max./Shock Flow:
Chlorine: Liquid/Gas: Dosage: Rate/Scnedule
Reg. Chlorine In. Pressure Chlorinator/Ejector Distance Edition and all Mandatory Addendasy. Mir. Chlorine In. Pressure Corr All Low, Stamped N Mir. Material Material Model No. Corr Material Eigector: Item No. Qty. Waterial Mol. %. Mech. Design Press. Temp. All Low, Stamped N Bres. Value: Aid Sector Model No. Temp. Mol. %. Mol. %. Bres. Corr: Item No. Qty. Material Mol. %. Mol. %. Bres. Corr: Mire Model No. Temp. Mol. %. Mol. %. Bres. Corr: Item No. Qty. Material Mol. %. Mol. %. Mir. Model No. Material Mol. %. Mol. %. Mol. %. Mir. Model No. Material Mol. %. Mol. %. Mol. %. Ventilation System Type: Nodel No. Material Mol. %. Mol. %. Mir. Model No. Range Samp Material and Mandatory Addendasy. Mir. Mor. Material and Mandatory Adde
Ch_lorinator: Item No. Qty. Corr. Corr. All row. Standed Stander St
MB.r./Max./Shock Flow
Bater name brack Qty. -Water in Press./Temp. // Andre in Temp. //
MFr. Model No. Vapor Capacity: Mol. Wt. Mol. Wt. Mol. Wt. Min./Max./Shock Flow. Qty. Material Ilquid Capacity: Mol. Wt. Mol. Wt. Mol. Wt. Mol. Wt. Min./Max./Shock Flow.
Min. /kax. /Shock Flow / Material Model Material Mir.
FJ ector: Item No. Uty. Material Vendors dualariateed Removal %. Vendors dualariateed Removal %. Mfr.
main main
Chlorine Residual Analyzer: Item No. Qty. Mat'T. Inhibitors: Type Concentration PH of Liq. Mr. Motel No. Water Test Kit: Mfr. Div. Volts PH HZ Power K Water Ion System, Type: Mat'I. of Const. Inhibitors: Type Concentration Pack/Other Water Test Kit: Mfr. Model No. Range Mat'I. of Const. Meterials of Construction: Heads set Support (Int.) Specifications: Painting Model No. Range Meterials of Construction: Heads Set Support (Int.) RecularED CL2 Model No. Range Meterials of Construction: Beauforments: Bvp Purchaser/Vendors; Diff. Press. Gauge, Level Gauge, Press. Gauge, Relief Valve, Temp. Indicator with Thennowell. Specifications: FOR CHLORINE HKd, SAMPLE-1 Notes: I) Elastomer 0-rings are not permitted in hydrocarbon service. 2) Slip-on inch thick herringbone knitted stainless stele (Als: Demister pads shall consist of four (4) demist pads separated from each other by glass wool mats (three required) and held toge by end support grids to form a removable cartridge. Demister pads shall be four permitted in hydrocarbon service. 2) Slip-on inch thick herringbone knitted stainless stele (Als: The service conditions. All H EVAPORATOR HKd, SAMPLE-1 Notes: I) Elastomer 0-rings are not permitted in hydrocarbon service. 2) Slip-on inch thick herringbone knitted stainless stele (Als: The serv
Mrr. Model No. PH HZ Power K W Chlorinator House: Item No. Otts. PH HZ Power K W Ventilation System, Type:
Electrical: Area Class <u>Gr.</u> Div. Volts_PH_HZ_rower with the construction: Packets Div. Volts_PH_HZ_rower with the construction: Madei a Electricals of Construction: Madei a Support (Int.) Accessories: Control
With Tractor Weil accessories: Construction Media Support (Int.) Water Test Kit: Mfr. Model No. Range Nozzies Media Support (Int.) Specifications: Painting Model No. Range Media Support (Int.) The Vendor shall furnish as a minimum the equipment and components shown on this sheet. RegultATED Cl_2 For RegultATED Cl_2 Paint Paint Accessories: For For CHLORINE For CHLORINE For CHLORINE For ChloRINE For CHLORINE For Nozzies Seconditations: Wind The vendor shall furnish as a minimum the equipment and components shown on this sheet. The Vendor shall For CHLORINE For CHLORINE For CHLORINE For Reside further of a consist of four (4) demist paiss sparated from each other by glass wool mats (three required) and held toge by end support grids to form a removable cartridge. Demister pais shall be four Motes: For For <td< td=""></td<>
Accessories: Control Instruments: Notes Control Model No. Range Control Notes Notes Control Notes Control Notes Control Notes Control Notes Control Notes Notes Control Notes Control Notes
Water Test Kit: Mfr. Model No. Range The Vendor shall furnish as a minimum the equipment and components shown on this sheet. Painting REGULATED CL2 CHLORINE CHLORINE Korr CHLORINE CHLORINE Korr CHLORINE CHLORINE Korr CHLORINE CHLORINE Korr CHLOR
water rest N1: MTr. model No. kange Specifications: Painting The Vendor shall furnish as a minimum the equipment and components shown on this sheet. Painting REGULATED CL2 CL2 REGULATED CL2 CHLORINE RESULATED CL2 CHLORINE FOR CHLORINE FOR CHLORINE FOR CHLORINE CHLORINE Instruments: ANALYZER Hot SAMPLE- Int Int CHLORINE Int
The Ventration of the Ventrati
REGULATED CL2
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a) H EVAPORATOR H (b) CHLORINATOR H (b) CHLORINATOR H (b) CHLORINATOR H (b) CHLORINE Pack and the construction of the constru
The two dot in the second of
b LIQUID c IANALYZER IANALYZER IANALYZER inch thick herrinobone knitted stainless steel (AISI Type 316) wire mesh. Glass mats shall consist of a four (4) inch thickness, compressed to two (2) inch of 15 per cu. ft., standard glass wool batting combined with an inert binder suitable the service conditions.
CHLORINE CHL
T A A A A A A A A A A A A A A A A A A A
The vendor shall furnish as a minimum the equipment and components shownonthis
$ \begin{array}{c} \bullet \\ \bullet \\ \bullet \\ \bullet \\ \bullet \\ \bullet \\ \bullet \\ \bullet \\ \bullet \\ \bullet $
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$\langle f \rangle$ in Purch. Piping 39 $\langle f \rangle$
Sym clisize Rating Facing $\rightarrow 007$ WATER IN $\rightarrow 000$ WATER IN $\rightarrow 000$
b I I 43 b Pair: 2 LEVEL f t
44
e Dimensions Length Width Heiaht Weiaht LBS/KG 45
f Analyzer Anal (Inst) 47 Sym Dize Kating racing
y g 1 I Chlorinator (Inst) 48 b Weight LBS/KG
L Cleur Cran Linst 49 C Dimensions Length Width Height Shipping
Deperating
EQUIPMENT DATA SHEET
0 3 CLIENT EQUIPMENT DATA SHEET
0 11EM
NO. DATE ENGINEER CHURAEVIEW APPROVE REC'N. NO. PAGE OF DATE ENGINEER CHURAEVIEW APPROVE REC'N. NO. SPEC. NO.

DEAERATOR (SPRAY-TRAY)	DEAERATOR (SPRAY-TRAY)
MANUFACTURERMOCELOTYENGLISWMETRIC UNITS WATER STORAGE CAPACITYHCRIZONTAL/VERTICAL DESIGN PRESSURE: ASME BPV CODE SECT. VIIDIV. I PRESS AND FULL VACUUM	The vendor shall furnish as a minimum the equipment and components shown on the sketch and as called for on the data sheets.
OPERATING DATA 11 RATED WATER INLET OTY/PRESS/TEMP	Flow Balance 1 + 2 + 3 = 4 + 5 $R \cdot y + R = 1$ $R \cdot y + R = 1$ $R \cdot y + R = 1$
MAX. QTY/PRESS/TEMP: / // // 3 MI N. QTY/PRESS/TEMP: / // 4 STEAM 15 MAX. QTY/PRESS/TEMP: / 6 MIN. QTY/PRESS/TEMP: / 7	$e + \frac{1}{2} + $
8 UEAERATOR UN IT: 19 DEAERATOR AND STORAGE MATERIAL: ASME 20 HEATER AND STORAGE/ CORROS I ON ALLOWANCE 21 HEATER INTERNAL MATERIALS: 22 TRAY MATER I AL: 23 SPRAY VALVE MATERIAL: 24 MANWAYS S I ZE HEATER/ STORAGE:	BRIDLES TYPICAL-DC (FOUR PLACES) COVERPLOW LEVEL STOLAGE STOLA
25 VORTEX BREAKERS: 26 Ti WATER AND I PH. CONDENSATE NOZZLE: SI ZE 27 21 2PH. Condensate Nozzle: Size 28 3L LOW PRESSURE STEAM NOZZLE: SI ZE 29 DEAERATCR WATER Outlet Nozzle: Size 20 PUMP BALANCE LI NE NOZZLE: SI ZE/OTY, 31 DEAERATCR DRAIN NOZZLE: SIZE	C II C I Sample S Sample S Sample S SAMPLE
PUMP K ICKBACK/Sparger Nozzle: Size 33 ACCESSOR I ES: 34 REL I EF VALVWAC. BRKR: S I ZE/SET. PRESS. 35 [4] VENT VALVE:SIZE 36 INLET CONTROL: 37 LEVEL CONTROL MANUF. /MODEL NO.	Overflow'
18 OVERFLOW CONTROL: BY: VENDOR/KELLOGG 39 CONTROL VALVE: S ZE/MFR/MCD, NO. /_/ 40 CONTROL VALVE MATER ALS: / 41 LEVEL CONTROL MANUF. /MODEL NO. / 42 .INSTRUMENT ALS:	
14 LEVEL S# I TCHES/ LEVEL GAUGE MFR/MOD. NO. 2000000000000000000000000000000000000	$\begin{array}{c c c c c c c c c c c c c c c c c c c $
6 MANAGING OFFICE ENGINEERING OFFICE 5 ERLIGGE MANAGING OFFICE 3 CLIENT	Vendor
2 2 COCATION	21 3 CLIENT 3 2

	PROPOSAL DATA	SHEET		
1	NO. & TYPE VESSELS	1		
2	SIZE: DIAM. X STR. SIDE			·····
3	DESIGN PRESSURE/CONSTRUCTION	•		•
4	LINING TYPE & THICKNESS			
5	INTERNALS: TYPE AND MATERIAL			
6	UPPER DISTRIBUTOR			
7	REGENERANT DISTRIBUTOR			
8	INTERFACIAL COLLECTOR			
9	UNDERDRAIN			
0	MAIN PIPING . SIZE/MATERIAL/LINING			
1	CONTROL VALVES: TYPE			
2	MATERIAL BODY/LINING			
3	RESIN VOLUME PER VESSEL			
4	CATION/ANION			
5	MANUFACTURER & I.D. NUMBER			
5	RATING KGR/FT3			
4	BED DEPTH/FREE BOARD			
8	ACID DOSAGE: LBS/F13 & LBS/REGEN.			
3	LAUSTIC DUSAGE LBS/FT3 & LBS/REGEN.			
	GALLONS PER REGEN. : GROSS/NET		· · · · · · · · · · · · · · · · · · ·	
님	FLUW: GPM/SU.FT. AT MAX. FLUW			
4	PRESSURE DRUP AT MAX. FLUW			
54	PIDING PER SPECIFICATION			
5	INSTR DER SPECIFICATION			
5	INSTR. FER SPECIFICATION			
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30	EXCEPTIONS TO SPECIFICATION			
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		KELLOGG	MANAGING OFFICE	ENGINEERING OFFICE
	y			
			DATA SHEET	
		CLIENT		
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		ITEM DEMIN		
ji.		ITEM NO	SPEC	C. NO
1	INO WITE ENGINEER URK. REVIEW APPROVE	RED'N. NO.		PAGEOF



CATION OR ANION UNIT TO BE PREASSEMBLED ON A STRUCTURAL STEEL SKID. WITH EQUIPMENT TUBED AND/OR WIRED TO THE SOLENOID VALVE BOX. SUBJECT TO SHIPPING CLEARANCES, MORE THAN ONE VESSEL CAN BE MOUNTED ON A COMMON SKID.

CONDUCTIVITY INSTRUMENT AND CELL USUALLY NOT FURNISHED FOR CATION UNITS. (REFER TO DATA SHEETS).

INTERCONNECTING HEADERS BY VENDOR. FOR FIELD INSTALLATION BY M. W. KELLOGG. WHERE t TEMS ARE MARKED (OPTIONAL). SEE DATA SHEETS.

PANEL MOUNTED INSTRUMENTS SHOWN **OUTSIDE** THE SKID LIMITS SHALL BE PROVIDED BY THE VENDOR.

6						MANAGING OFFICE	ENGINEERIN	G OFFICE
5								
4						EQUIPMENT SKE	ГСН	
3					CLIENT			
2					LOCATION	JOB NO		
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	Ĩ	PROCESS DESIGN SECTION, CORPORATE ENGINEERING	MECHANICAL SCALE DATA SHEET			
		SERVICE	10C4110N			
	-	NO. BEOUIRED	MANUFACTURER			
	Ś	DUTY	MODEL NO.			
	č	TYPE				
	S.					
	ž		PROCESS CONDITIONS			
	CHARACTERISTICS OF MATERIAL					
	_	MATERIAL				
	-	STATE OF MATERIAL				
		TEMP OF				
	.>	DULY DENSITY IN /ET 3				
	÷.	AMOUNT/DATCH				
	×	AMOUNT/BATCH				
	μ	TOTAL VOLUME OF MATERIAL				
		AVERAGE DENSITY				
	Ē	TOTAL WEIGHT OF MATERIA!				
	Ę.	ACCURACY	SENSITIVITY			
	2 1	ATMOSPHERE				
			MECHANICAL DESIGN DATA			
	Í	DESIGN CAPACITY NET WEIGHT				
\sim	í.	WEIGHT OF CONTAINER EMPTY				
\sim	Ē	SIZE OF CONTAINER				
		LOCATION OF DIAL WITH RESPEC	TTO SCALE (HEIGHT)			
			(menori)			
		LOCATION OF SUPPORTING LEV	EL WITH RESPECT TO SCALE (HEIGHT)			
		TOTAL SCALE CAPACITY				
	MAXIMUM MOVEMENT WHEN FULLY LOADED					
	DIAL GRADUATIONS (MINIMUM)					
	ADJUSTABLE MARKERS REQUIRED					
		TARE BEAM	CAPACITY BEAM			
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Π	PROCESS DESIGN SECTION ROTARY VALVE CORPORATE ENGINEERING DATA SHEET
	OPERATING DATA
REMARKS	TREDUCERIPTI:
× <	MECHANICAL DATA
DATE APP	MFGR:
ŕ	MATERIALS OF CONSTRUCTION
BY CHKD	CASING
	P A C K I W G C A W D
	MOTOR DRIVE DATA
KED BY DES SUPRV.	ITEM NO: MTD, BY: HP: RPM: FRAME: FRAME: MFG'R: INSUL TYPE: INSUL ENCLOSURE: CLASS I GROUP D / VOLTS/PHASE/CYCLE / BEARINGS: LUBE: FULL LOAD AMPS: S.F.
H	SPEEDREDUCER MFG'R
DHAWN	MISCELLANEOUS TOTAL WEIGHT:
2	JOE NO PLANT LOCATION PROJECT NO REF DWG.NO
224840	TITLE DISCUSSION NO SERIES STEP NO.
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Appendix C

QUESTIONNAIRES OF EQUIPMENT SUPPLIERS

Equipment is supplied on the basis of information about the needs provided by the ultimate purchaser and user. Although the more information, within limits, the better, suppliers do require a certain minimum amount before they can make recommendations of equipment and price. Some kinds of equipment are sufficiently standardized to allow their specification by standard forms like those of Appendix B. The questionnaires of individual Suppliers sampled in Appendix C are mostly of specialized equipment that requires custom designing to a greater extent. This is a random selection and other suppliers of each type are available to provide service. Following is a listing of questionnaires contained in this Appendix C. The number appearing in italic is the page number on which the form appears.

A. Comminution

- 1. Crushing, grinding, etc. (Sturtevant) 728
- 2. Size reduction (Williams) 728
- B. Conveying
 - 3. Pneumatic conveying (Ducon) 729
 - 4. Tubular, bucket, oscillating, etc. (Hapman) 729
 - 5. Conveying elevator (Meyer) 730
- C. Drying
 - 6. Spray drying (Bowen) (2 pp.) 730
 - 7. Evaporators (Goslin-Birmingham) 731
 - 8. Dryers (Krauss-Maffei) 732

9. Drying with dessicants (Selas) 732 10. Rotary equipment (Vulcan) 733 D. Mechanical separations 11. Mist elimination (Glitsch) 733 12. Air classifiers (Raymond) 734 13. Cvclones (Krebs) 734 14. Plate filters (Sperry) (2 pp.) 735 15. Continuous centrifugal filter (Bird) (2 pp.) 736 16. Centrifuges (Robatel) (2 pp.) 737 E. Mixing and blending 17. Mixing and blending (Abbe) 738 18. Mixing and blending (Bépex) 738 19. Membrane separators (Monsanto) 739 F. Separations 20. Distillation (Glitsch) 739 21. Solvent extraction (Robatel) (2 pp.) 740 22. Crystallization (Swenson) 741 G. Thermal processing 23. Incineration (Vulcan) 741 24. Thermal processing (Bépex) 742 H. Other processes 25. Agglomeration (Feeco) 742 26. Briquetting (Bépex) (3 pp.) 743 I. Safety data 27. Material safety data sheet (U.S. Dept. of Labor) (2 PP.) 744



103 Clayton **Street** Boston, MA 02122 617 **825**-6500 Cable EMERYSTONE Telex 94 0677



To allow us to better assist you in solving dry processing problems, please use this form when requesting information on Sturtevant equipment.

1. Material to be Processed: ____

2. Processing Method(s) Required (Number in order needed):

a. Crushing	f. Mixing
b. Grinding	g. Screening
c. Air Separatmg	h Conveying
d. Micronizing	I. Elevating
e. Blending	i Other (specify)
3. Characteristics of Material being Processe	ed:
a. Specific Gravity	d. Moisture Content%
b Hardness	e. Handling Hazards
c Feed Size	f. Temperature Limits°F.or°C.
4. Production Requirements:	
a Product Size Required	c. Contamination Tolerance
b. Capacity Required	
5. If Air Separating is required:	
a. Amount of product entrained IN feed%	d. Is heating or cooling required in separation
b. Grinding media pre- ceding separation	e. Product size tolerance, if any
c. Preferred load if	

6. Any comments which would help our engineers evaluate your requirements:



Thank you for your interest in Williams size reduction equipment and/or systems. As a first step toward giving you a highly personalized answer to your need, we would appreciate your supplying the following information.

Thank you for your help. We will be back to you soon.

Form No 943

Name	_Title	
Company	Address	
City	State	Zip Code



SYSTEMS DATA



Hapman Conveyor Co.

6002 E. Kilgore Road Kalamazoo, Michigan 49003 (616) 343-1675 Telex. 224468

data sheet

(Fill out completely)

CUSTOMER:	CATE · REPRESENTATIVE:		DATE
		CUSTOMER	ZIP
		ATTN :	TITLEPHONE ()
Name Particle size	PRODUCT CHARACTERISTICS	LOCATION OF INSTALLATION	() OSCILLATING CONVEYOR () OTHER
	VEN ASSUMED GIVEN GIVEN	ASSUMED DESCRIPTION OF MATERIAL	SERVICE REQUIREMENTS
Bulk density lbs/cu. ft Angle of repo se	Hygroscopic Corrosive Multi-backrospanubrr	1. Material	Required Capacity (in cubic feet/hour or min.)
Panicle shape Surface texture Change of state temp ^o F Hzzardus (explos., toxic, ● tc)	Temp of product °F Specific beat Is breakage objectionable? If so, what % permissible?	Particle Size (Mesh or Fraction of Inch) Max. Min. Proportions	Operation Continuous Intermittent Operating Period. Hours
Tendency to bridge Moisture content % free		3. Actual Weight per Cubic Foot	Frequency
Capacity lbs/min (instantaneous/ay Horizontal Distance , feet Vertical lift , feet 45° Elbows 90° Elbows Ambient temp at blower inlet °F	SYSTEM REQUIREMENTS 1 2 3	4. Fercentage and Type of Equilibrium Present	MOTOR CHARACTERISTICS Voltage AC/DC Phase:Cycles () Totally Enclosed - Fan Cooled () Explosion Proof. Class G r o u p
Material of construction in contact with product	DESCRIPTION (Use back for simple system sketch)	Sluggish MildVe	CONSTRUCTION SPECIFICATIONS ery 1. Casing - Mild Steel
		Explosive Light and Fluffy	2. Chain Sealed Pins der Mild St1. #304 S.S. Open Pin Construction
		Describe any possible contamination problems	3. Flight Material — Factory Spec Other
	<u>G E N E R A L</u>	Pressure zones	Miscellaneous
Starters: furnished; 0 Available currentv / Available watersgpm	Notor enclosure Motor enclosure Phase H Z Open drip proof Psig OF Totally enclosed	IMPORTANT: FURNISH SKETCH ON BACK OF TH SHEET OR ON SEPARATE SHEET. Show plan, e elevation and/or side elevation properiv indicated as su	HIS SALES REPRESENTATIVE: nnd u.c.h
Available airscfmscf	psig Explosion proof	with all pertinent dimensions . Show your idea of p ferred conveyor path together with possible alternat Also indicate all clearances and obstructions to all	ore- ore- tes. Note: When sending jn Sample, attach label showir low Kind of Material, Company Name, Representative, ar

from your sketches

840 FIRST AVENUE . KING OF PRUSSIA, PA. 19406 . PHONE (215) 337-3770 . TELEX 84 . 6325

us to select the most practical circuit should thrs differ Date. Three to four cubic feet of material are required for complete testing.

OVER



DESIGNERS AND MANUFACTURERS MACHINE COMPANY OF CONVEYING AND PROCESSING EQUIPMENT

Gentlemen

Please quote us on a SIMPLEX CONVEYING ELEVATOR to meet the specifications and approximate dimensions as indicated below and on the reverse side of this sheet.

PRODUCT TO BE HANDLED	will be treated in strict confidence. We appreciate that some data may not be available or relevant. Please
APPROXIMATE WEIGHT OF PRODUCT PER CUBIC FOOT	BOWEN ENGINEERING INC., P.O. Box 898, Somerville, New Jersey 08876, U.S.A.
DESIRED MAXIMUM CAPACITY, pounds per hour	Telephone: (201) 725 3232 Telex: 4 (5408) (11)
IS PRODUCT CORROSIVE' DUSTY, FINE POWDER' GRANULES?	COMPANY Ref./Project No:
STICKY' A B R A S I V E ' - PIECES, AVERAGE SIZE OF PIECES	Company Name: Personnel:
TYPE OF BUCKETS PREFERRED Fabricated Stainless Steel Mild Steel Plastic	
Cast Aluminum Alloy Other	تبت
is dust-tight/weather-tight enclosure REQ'D?	
END DISCHARGE ONLY-WITH SIDE DISCHARGES NUMBER OF SIDE DISCHARGES	MATERIAL/PROCESS Name/Composition:
(Indicate on drawing (Fig. 3) on reverse side if discharges are to be Left Hand Of Right Hand when viewed from intake end of SIMPLEX Also indicate approximate location of discharges).	Chemical formulae: Process: dry/cool/heat/agglomerate/react/absorb/congeal/calcine/
SIDE DISCHARGES TO BE MANUALLY CONTROLLED Of SOLENOID CONTROLLED for remote	Type of dryer envisaged ; spray
operation	Heat source: direct/indirect: gas/LPG/oil/steam/electricity/HT gil,
TYPE OF FEEDER/HOPPER PREFERRED STANDARD INTAKE HOPPER only BUILT IN FEEDER,	Has material bean dried before: yes/no. If so, how Temperatures
less hopper (for feeding from bin)	CAPACITY DATA Feed rate lb/h at Feed CONC.% solids Evap, rate lb/h
SIMPLEX. (for manual feeding) with ECCENTRIC DRIVE with ELECTRO MAGNETIC	For minimum evaporation rate :
DRIVE Feed Trough and Hopper to be Stainless Steel Mild S t <u>eel</u>	For Maximum evaporation rate:
or Galvanized	Product rate: Ib/h : mm uni, normal, m a x unin n me design
	Feed rate : ID/h : [T][T] Inn anno y max mananana _
Ir nead 1001115 restricted, state Celling neight	FEED DATA
Per	Type: solution/suspension/emulsion
	Feed rheology (liquid): Newtonian/pseudoplastic/dilatant/Bingham plastic/thixotropic/rheopectic/viscoelastic
Company	Feed rheology (solid): free flowing/friable/cohesive/sticky/paste/
Address	Does rheology change markedly with feed COncentration? yes/no. If so, add notes overleaf
	Moisture: water/other (see solvent data overleaf). Impurities in feed.
Diverse Ma	If feed is SUSPENSION, what concentration of solubles in liquid.
Date	Does feed form stable foam? yes/no. Material of construction ID CONtact with feed Feed temperature:

QUESTIONNAIRE DRYING SYSTEMS

₽40₽\$

The information below is required to enable us to prepare a design and quotation. All information given

PRODUCT DATA

Moisture content:
Is product heat sensitive: yes/no. If yes, what maximum temperature for dry solids
Material of constructton IN contact with dry solid:
Required particle \$126; min micron, av micron, max micron, max
Required bulk density range: to lb/ft ³ Required final discharge temperature:
Melting point:
Other special properties:
Toxic: yes/no. Flammable: yes/no. Explosive: yes/no. Abrasive: yes/no. Hygroscopic: yes/no.
Thermoplastic: yes/no. If yes. at what temperature: *F

SOLVENT DATA

Name: ·····	Toxic:	yes/no.	Corrosive: yes/no.	
Impurities:				
Flammable: ves/no. If yes, explosive limits:		Flast	n point	
Boiling point: F Latent heat of vaporisa	ation:			BTU/Ib.
Specific heat liquid BTU/Ib. °F	Specific	heat vapour:.		BTU/lb. 'F.
Other data:				

ANALYTICAL METHODS

Moisture	content:	
Other:		

ENVIRONMENT/SAFETY

ENVIRONMENT/SAFETY			
Permissable emmission in exhaust air:	grn/ft ³	Effluent water Imits	
Maximum noise level: In plant	dBAat 3 ft.	Community	dBA at ft.
If area classified hazardous, give division, group, etc.			
ls dust flammable: yes/no. Ignition temperature, o	loud	layer	°Fat inch thick
Is dust explosive: yes/no. Dust class Ma	ax.explosive pressure	Max.	rate press. [ise

LOCAL CONDITIONS

Plant	location:	indoors/ou	tdoors.	Existing	building:	yes/no.	lf yes. s	ubmit drav	vings.		
Attitude		ft. a	above sea	level.	Max. wind	velocity			Earthquake:	Richter	
Ambient	temper	ature: Mir	nimum ,,,		°F	Average		F	Maximum .		F
Range	ambient co	onditions fo	or design	evaporat	tion rate:	Winter	°Fat	%RH,	Summer	°Fat	%RH max.
Special	conditions	: high du	st loadin	g/freezing	fog/				•••••		

UTILITIES Steam pressure at plant: Compressed air:	psig. psig.	Towns water:
Electricity: power V,		ph Hz; Control V AC/DC.
Fuel gas: Name Gross	calorific	: value BTU/scf. Pressure psig./inch wg.
Fuel oil: Grade	viscosi	Sity Gross calorific value BTU/lb.
Cooling water: Supply [•] F	Return	⁶ F _{max.} Available pressure differential

R	EM	AR	KS

GOSLIN-BIRMINGHAM, INC. EVAPORATOR INFORMATION SHEET

plant address or JOBSIT	Е		
FEED SOLUTION	PPH 9	6 Dissolved Sol	ids
Temperature	Specific Gravity	Specific	Heat
Analysis			
% Suspended Solids	Viscosity	pH	:
Temperature Limit	Thermal	Conductivity	
PRODUCT	% Dissolved	% Suspen	ded Solids
Viscosity	at Temperature	Specific	Gravity
Scaling?	Foamy?	Specific	Heat
Thermal Conductivity			
OTHER DATA	Availa	ble Steam	
Low Pressure	Psig	Temp.	cost
High Pressure	Psig	ſemp.	cost
Cooling Water	High Temp.	Low Temp.	
Power AC		D C	Cost/KW
Vo	lts Hert		Phase
Material of Constructio	n		
Liquid			
Vapors			

Spray Drying-(continued)

COMPANY	KMC PROJECT NO.
CONTACT PERSON	KMC CONTACT PERSON
PRODUCT: formula. trade name	4. DRY PRODUCT:
DRYER CAPACITY: 2.1 Wet product feed	4.1 Requested form of dry product 4.2 Admissible dry product temp°F 4.3 Dry product to be cooled- yes no to°F 4.4 Is there some isothermalabsorption known of the dry product? (Please enclose) yes no 4.5 Electrostatic charging yes no 4.6 Risk of oxidation yes no 4.7 Is the product hygroscopic? yes no
3. <u>WET PRODUCT FEED</u> 3.1 Originates from: filter centrifuge other	4.8 Meltmg point°F 4.9 Conveyability?
3.2 Feed. continuous batch other 3.3 Volume/batchgal. 3.4 Batch cycleMIN 3.5 Consistency: liquid sticky past y friable thixotropic	 Do you dry the product IN your plant now? What dryer type? With what results?
3.6 What conveyors worked satisfactorily?	6. If MOISTURE IS a solvent, IS recovery required?
3.8 Wet product temperature°F	7. Which materials of construction should be used or must not be used? For contact w/wet product For contact w/fresh gas For contact w/fresh gas
3.9 Composition of product % m0isture (vapor pressure % curves?) % 3.10 Moisture determination: %	 SPECIAL REFERENCES, REQUESTS, SUGGESTIONS PLEASE ENCLOSE SAFETY DATA SHEETS RELATING TO HAZARDOUS PROPERTIES (TOYIC IGNITABLE INFL AMMABLE ETC.)
constant weight min, under vacuum l' Hg other	9. AMBIENT AIR CONDITIONS 9.1 Temperature°F 9.2 Humidity% or Lbs. H ₂ 0/lb. Dry Air
3.13 True product bulk density way of measuring the bulk density?	10. <u>UTILITIES</u> 10.1 Electrical characteristics V,PH,Cycles
3.15 Spec, heat wet product BTU/lb 3.16 Spec, heat solids BTU/lb	10.2 Heating Media: @ PSIG



Date_

Selas	Corj	porat	ion	of	America	
Dreshe	r,	Pa.				

Attn: General Industry Division

Gentlemen:

I am interested in a Selas Dehydrator to meet the following specifications. Please send proposal, with the understanding that I am under no obligation:

GAS AS SUPPLIED TO DEHYDRATOR FOR DRYING

Type of gas:		
Volume at standard atmospheric conditions .dm	or at working	pressure:
Gas temperature entering Dehydrator:.	°F	
Working pressure:	•	
Moisture content: saturated at entering temperature 🗌 or what	% of saturation	
Specific gravity:		
Specificheat:		
GAS LEAVING DEHYDRATOR		
Final moisture content: dewpoint		. °F
o r grainsper cubic foot at		
Final temperature		°F
DENYDRATOR OPERATING SCHEDULE		
Continuous , or, if intermittent, hours on, hours	off.	
TYPE OF OPERATION		
Manual Semi-automatic	Fully Automatic	
TYPE OF FIFTDICAL CONSTDUCTION	<u> </u>	
	Explosion proof	
	Other	
	Uniti	
	Phase	Cycle
Power D NA D SA Volts	Phase	Čvcle
Steam Pressure		
Cooling water temperature: Summer °F Winter °F		
Cooling water pressure:psig minimum		
Preferred heat source for desiccant readivation.		
MECHANICAL CONSTRUCTION		
	·	
REMARKS:		
FIRM NAME		
Subl inity DT:		
11NE:		



3.18 ph value ___

3.19 Max. admissible wet product temp. 3.20 Dryer atmosphere: air, inert, vacuum

_°F

M 318

.....

ROTARY EQUIPMENT INQUIRY

Date

TO:	VULCA	AN IRON	WOR	ks, inc	C.	
	Miners	National	Bank	Bldg.,	Rm.	1050
	Wilkes	-Barre, I	Penns	ylvania	187	01

We are interested in equipment for the following process: drying . . . , cooling . , calcining (other)

Our interest is related to our: economic evaluation preliminary design final analysis & purchasing budget estimate

Process data available includes:

type of material material wt./cu.ft.; wet , dry . material specific heat . material particle size (screen analysis desirable) capacity-feed rate . discharge rate or moisture content, initial % and final %. material temperature, Initial and final maximum temperature of material without injury . dutv cvcle hours per dav and davs per week. final product temperature desired .

type of fuel to be used: 0il@ ... btu∕gal, psig and ssu. Gas@ btu/cu. ft., and psig. Other

Burner is to be automatically or manually controlled. Material can , cannot be processed in contact with gases of combustion. The . sticky, a b r a s i v e , corrosive nature of the material requires special consideration.

Motors:	т	Е	F	С		Open		F	hase	9		C	ycle		Volts
Installation	is to	be	inde	oor,								c	utdoor		
Elevation of	fins	stalla	tion	, .		feet		i	abov	е			sea		level.
Dust col	lecti	on	eq	uipmen	t to	be	inclu	ded					(yes	or	no).
Exhaust fa	in to	be	inc	luded		(У	е	s	C)	r	n	ο).
Your re	spor	ıse	01	n or	befor	re ((Date)							. is	desired
Please rep	oly to	o: (Com	pany)											

(Address) .				• •		 	 			
(Attention)	 	•	• .			 	 	 		
(Telephone)	 				 	 		 	• • •	

GLITSCH, INC.

PLANT & HOME OFFICE LOCATION: 4900 SINGLETON BLVD. , DALLAS, TEXAS @ 214/631-3841 , TELEX 73-329

MIST ELIMINATION AND LIQUID DROPLET COALESCING EQUIPMENT APPLICATION DATA

General Information: Operating: Pressure-Temperature	MIST ELIMINATORS Is the mist eliminator to go into an existing vessel? Yes D No D
Vapor (Gas) Phase: Composition	If yes, please provide a detailed sketch or blueprint of the vessel to be used Do you wish to have Glitsch quote on a suitable vessel?
Rate	Yes No LIQUID DROPLET COALESCERS Note: The Coalererr are generally sold only as a complete unit with vessel.
Liquid Phase: Composition wt. % or mol. % (state which)	SPACE LIMITATION Are there any physical limitations due to vessel size? Yes No If yes, please describe
Rate O operating temp.	SUITABLE MATERIALS OF CONSTRUCTION (See list below.) Shell :
Droplet size range Source of droplets (condensation. spray, etc.)	Mist eliminator and/or COalescer Support grids 304 Stainless Inconel Copper
Does liquid present a fouling problem when collected? Yes D No D	316 Stainless Carpenter 20 Carbon Steel 317 Stainless Incoloy Bethanized Steel 430 Stainless Hastelloy C Kynar ®
If yes, what type Please describe any problems encountered	434A Stainless Titanium Polypropylene Monel Tantalum Teflon ® Nickel Aluminum Other (specify)

Does the vapor or liquid phase have any insoluble solids in

it? Yes No If yes, describe their nature.

Does the liquid phase have any dissolved solids in it? Yes cl No 🔲 If yes, describe their nature.-

If the potential application involves a number of units, please state the anticipated annual requirement.

●A corrosion allowance is acceptable for the shell and support grids. Other wetted parts should have the ability to withstand corrosion.

Raymond" Jet-Strea	m" Classi	f le rs			Kre	DS ana DE	hlam
C-E Raymond's experience in the the <i>Jet-Stream</i> " classifier in pro- can be put to work for you to he no-cost evaluation of your applica fold and mail. If you need assist	classification of p viding consistently elp provide the kind tion, simply remov ance, call us at [3 (please	wowdered materials a sharp cut-points a d of quality product e this sheet, answe t12] 236-4044. print)	and the success of at full production capac you demand. For a er the questions below,	cities		cription of the cyclone	
Your name	C	ompany					
Address	_City	State	Zip				
Telephone ()	Best ti	me to call:		am pm	(Please com	plete flow diagram be	DW.)
Material to be classified:		Bulk density (Ibs/cu	ft)		Feed slurry	characteristics	
Specific gravity	Feed	temperature (°F)			SOLIDS	lbs/hour	Specific G
Moisture in feed (% by weight)		Feed capacity (I	lbs/ħr)			lbs/hour	Specific G
Elevation above sea level (ft)			·		SLURRY	ibs/hour	Specific G
		— •			Liquid Viscosity	A CPS	H Temp. (*F
EIN EIN					Custom Drass		
	Cumulative percent of	quirements	icron size rhown		System Pressu	re, psi PM)	
Micron Size Feed to Cla	Cumulative percent of assifier	quirements product smaller than m Fine Product	licron size rhown corn. Product		System Pressu Ficewrate (USG	re, psi PM)	atch
Micron Size Feed to Cla	Cumulative percent of	QUITEMENTS product <u>smaller than</u> m Fine Product	corn. Product		System Pressu Flowrate (USG Other feed info	ve, psi PM) mnabon (corrosive, abrasive	. etc.)
Micron Sizer Feed to Cla	IENESS KE	QUIPEMENTS roduct <u>smaller than m</u> Fine Product	Sileron size rhown corn. Product		System Press Flowrate (USG Other feed thit	re, psi PM) rmation (corrosive, abrasive	. etc.)
Micron Size Feed to Cla	Cumulative percent of	QUIPEMENTS roduct smaller then m Fine Product	Corn size rhown corn. Product		System Press. Flowratin (USG Other feed fill Overflow al * Dearied med	re, psi Pla) ministon (corrosive, abrasive 	etc.) fine fraction)
Micron Stare Feed to Cla	Cumulative percent of assifier	QUIPEMENTS product <u>smaller than m</u> Fine Product	Silicon size rhown corn. Product		System Pressu Flowren (1860 Other feed the Overflow si * Desired mea - Desired mea - Desired weap	re, pai Pla) imation (corroave, abrasve unry characteristics of separation M percent solids	fine fraction)
Micron Size Feed to Cla	Cumulative percent of assifter	QUIPEMENTS aroduct <u>smaller than m</u> Fine Product	Corn. Product		System Press. Flowratin (USG Other feed fills) Overflow al * Desired mest - Desired mest Inclustions	re, pai Pla) rmation (corrosive, abrasive urrry characteristics i nof separation m percent solids	fine fraction)
Micron Size Feed to Cla Feed to Cla Method of analysis (<i>Ro-Tap</i> ® siev	e shaker, SediGra	QUIPEMENTS product <u>smaller than m</u> Fine Product ph® particle size a	nicron size rhown corn. Product		System Pressu Finanza (1950 Other feed the Desired med Desired weig Underflow a Underflow a	re, pel PM) PM) urry characteristics of separation ht percent solids percent solids	fine fraction)
Micron Size Feed to Cla Feed to Cla Method of analysis (<i>Ro-Tap</i> ® siev	e shaker, SediGra	Quirements product <u>smaller than m</u> Fine Product ph® particle size a	analyzer, or other equipr	ment)*	System Press. Flowrath (USC) Other feed the Overflow at * Deared metil Jonderflow a Deared weigh Deared weigh Special reg	re, pai Pui) Pui) urry characteristics i nol separation M percent solids percent solids ulrements	fine fraction)
Micron Size Feed to Cla Feed to Cla Method of analysis (Ro-Tap® siev Electric power available: Phase	e shaker, SediGra	QUIPEMENTS product <u>smaller than m</u> Fine Product ph® particle size a _Hz	Alecton size rhown corn. Product	ment)*	System Pressu Fixerrain (1850 Other fead this *Desared mean - Desared wean Underflow a Desared wean Special req	ma, pai Pia) Pia) mission (corroave, abrasve mission (corroave, abrasve urry characteristics (of separation M percent solids siturry characteristics percent solids ulrements	fine fraction) (coarse fraction)
Micron Size Feed to Cla Feed to Cla Method of analysis (Ro-Tap® siev Electric power available: Phase Type of final dust collection equ	e shaker, SediGra	Quirements product smaller than m Fine Product ph® particle size a _Hz	analyzer, or other equipr	ment)*	System Press. Flowren (180) Other feed into Overflow al "Deared mea Juderflow a Deared weigh Special req	re, pai Pla) ministion (corroavel, abrasive urry characteristics (of separation m percent solids siturry characteristics percent solids ulrements	etc.) fine fraction) (coarse fraction)
Micron Size Feed to Cla Feed to Cla Method of analysis (Ro-Tap® siev Electric power available: Phase Type of final dust collection equ Special materials of construction	e shaker, SediGra	Quirements product smaller than m Fine Product ph® particle size a _Hz	analyzer, or other equipr	ment)*	System Pressu Flowran (JBC) Other fead tift Overflow si * Desired mean - Desired weigh Special req	re, pel Pla) Imitation (corrosive, abrasve umy characteristics of separation Impercent solids siturny characteristics percent solids ulrements	(coarse fraction)
Micron Size Feed to Cla Feed to Cla Method of analysis (Ro-Tap® siev Electric power available: Phase Type of final dust collection equ Special materials of construction	e shaker, SediGra	Quirements product <u>smaller than m</u> Fine Product pph® particle size a _Hz	Alecon size rhown corn. Product	ment)*	System Pressu Flowren (1860 Other feed the Desired mean Desired weigh Special req	re, pai Pla) Pla) Imation (corrosive, abreave urry characteristics (of separation M percent solids sturry characteristics percent solids ulrements	etc.) fine fraction) (coarse fraction)
Micron Size Feed to Cla Feed to Cla Method of analysis (Ro-Tap [®] siev Electric power available: Phase Type of final dust collection equ Special materials of construction	e shaker, SediGra	Quirements product smaller than m Fine Product ph® particle size a _Hz	Analyzer, or other equipr	ment)*	System Press. Flowren (180) Other feed filt Overflow al * Deared weigh Juderflow s Deared weigh Special req * *********************************	re, pai Pui) imation (corroave, abrasve urry characteristics i rol separation if percent solids sturry characteristics percent solids ulrements uton-Particle size that equi EERS, 1205 CHARSLER OR	etc.) fine fraction) (coarse fraction 4a 1-3% plus by weigh VE, MENLO PARK (
Micron Size Feed to Cla Feed to Cla Method of analysis (Ro-Tap® siev Electric power available: Phase Type of final dust collection equ Special materials of construction	e shaker, SediGra	Quirements product smaller than m Fine Product ph® particle size a _Hz	Analyzer, or other equips	ment)*	System Pressu Flowratin (USC) Other feed this Descred weight Descred weight Special req "Mesh of separation KREEDS ENGINI	re, pel Pu) Pu) Imation (corroave, abrasve imation (corroave, abrasve urry characteristics of separation M percent solids percent solids percent solids ulrements tion—Particle size that equa tion—Particle size that equa	etc.) fine fraction) (coarse fraction) (coarse fraction) (coarse fraction)
Micron Size Feed to Cla Feed t	e shaker, SediGra	Quirements product smaller than m Fine Product pph® particle size a _Hz	Analyzer, or other equipr	ment)*	System Pressu Flowmen (1860 Other feed into Overflow al * Desired read Jnderflow 3 Desired ready Special req * Mesh of aspara KREBS ENGINI	re, put Put) imation (corrosive, abreave urry characteristics (of separation M percent solids sturry characteristics percent solids ulrements ulrements tion—Particle size that equal	elc.) fine fraction) (coarse fraction) (coarse fraction) /s 1-3% plus by weight /s US PARK, C

company City Zip Yourname elephone Date

Specific Gravity Specific Gravity Specific Gravity Temp. (*F,*C)

Mesh/Micron/mm	Ind. Wt.%	Cum. Wt.%
	•	
1]
		1

s (fine fraction)

* Desired mesh of separation • Desired weight percent solids	Flow diagram	OVERFLOW TO: n Thickener (Settler)
Underflow slurry characteristics (coarse fraction) Desired weight percent solids	FEED FROM:	Centrifuge Filter Evaporator Screen Other
Special requirements	Crystalizer Cryst	UNDERFLOW TO: Centrifuge Filter Classifier Screen Other

equals 1-3% plus by weight of overflow solids.

DRIVE, MENLO PARK, CA. 94025 E TELEPHONE: 415-325-0751 E TELEX: 34-8403

FILTER QUESTIONNAIRE	Filtration Temperature	°F Max. Permissible.	°F.
	Min. Permissible	°F.	
d. r. sperry 🎦 Co.	Size of Batch	gals.	
112 N. Grant Street	Closed or Open Discharge Desired		
North Aurora, Illinois 60542	Filtration Pressure	psig	
Date	Filter Cycle Desired	hrs.	
Company	ls ProductCake	Filtrate	Both
Address	Cake Washing Necessary (Yes. No)	Name of Wash Liquid	
Telephone No.	Temperature of Wash Liquid	°F.	
Name of Individual Reporting	Desirable Dryness of Cake: % Solrds		
Title	Physrcal Condition of Cake		
Type of Operation (Chemical, Tannery. Brewery. etc.	(pastry, sticky,	dry, cracks)	
Location	Thickness of Cakein.		
	Quality of Filtrate Desire or Allowed	Brilliant	<u> </u>
	Turbid	Decolorized	
(Please answer as many questions as possible)	If Turbid • Mex. Allowable particle size of solids	3	microns
Description of Matenal to be Filtered	If Filtrate is the Product		
Liquid	Filteraid to be used Y e s	No	
Solrds	If Yes: Precoat	Quantity	
Specific Gravity of Slurry®	Body Feed	%	
Emulsion? Solution?	Both		
Viscosity pH	Type, Brand, Grade		%
(Units)	Activated Carbon		9/
Avg. Particle Size of Solids	or Bleaching Earth used Yes or	No	70
Concentration of Solrds(ppm, lbs., 76, etc.)	Flocculation AgentType_	Brand	%
Physical Nature of Solrds (Sticky, granular, flocculent. etc.)	Recommended Materials of Construction		
Specific Gravity of Solrdsgms/cc			
Chemical Composition of Solids	Materials of Constructron not Permitted		
Nature of Filtrate-	Filter Media: Cloth	Paper	
(Water, alcohol, varnish, mixture, etc.)	Туре	Grade	
Specific Gravity of Filtrate gms/cc @ F	Filter Area	<u>ft</u> , ²	
Vapor Pressure of FiltratemmHg	Filter Cake Volume	ft	,
Dissolved Salts		Laboratory Filter Tests	
Solubility: @ 32°F Units	Filter Type		
@ 68°FUnits	Area of Eilter		
@ °F Units			
Conditions end Recommendations	Care volume or Filler		
General Comments			
	RESULTS		

PRODUCT	DATA	Max. impurities in solids Max. SS in clarified ML	Diop
Company	Project Status : Cl Pilot Plant	Material of const.	_
AddressState	Semi-works C. In production	Wash separation from ML	Equ
Plant address	Customer Contacts	Wash Requirements	Size
CityState Project References Bid data	R&D Engineering	Temp°F	Pre
Completion date	Purchasing	Composition	To
PROCESS	DATA	Ratio	By:
Present problems		Disposition of wash	Disp
I I I I I I I I I I I I I I I I I I I		Å	of
Feed 6 Susp. solids (ss)	Size Distribution Sieve% On Cumul. %	FLOW	SHE
Feed 5 Susp. solids (ss) p. grViscosity(cp) at(°F)	Size Distribution Sieve%OnCumul. % +M +M	FLOW	SHE
Feed 6 Susp. solids (ss) p. grViscosity(cp) at(°F) Process Femp(°F) Pressure-(PSIG) Rate(GPM)(lbs/hr ss) Valuable Component(s)	Size Distribution Sieve % On Cumul. % + M	FLOW	SHE
Feed 6 Susp. solids (ss) p. grViscosity(cp) at(°F) Process Femp(°F) Pressure-(PSIG) Rate(GPM)(lbs/hr ss) Valuable Component(s) BOD or COD (for waste)	Size Distribution	FLOW	SHE
Feed 6 Susp. solids (ss) p. grViscosity(cp) at(°F) Process Femp(°F) Pressure-(PSIG) Rate(GPM)(lbs/hr ss) Valuable Component(s) BOD or COD (for waste) Fype (origin) of waste	Size Distribution Sieve % On Cumul. % + M	FLOW	SHE
Feed % Susp. solids (ss) p. grViscosity(cp) at(°F) Process Temp(°F) Press ure - (PSIG) Rate(GPM) (lbs/hr ss) Valuable Component(s) BOD or COD (for waste) Type (origin) of waste Solids	Size Distribution Sieve% On Cumul. % +M +M Sp gr,Shape Ash content Grease content Permissable Flocculants Bulk density Mother Liquor Composition	FLOW	SHE
Feed % Susp. solids (ss) Sp. grViscosity(cp) at(°F) Process Temp(°F) Pressure-(PSIG) Rate(GPM) (lbs/hr ss) Valuable Component(s) BOD or COD (for waste) Type (origin) of waste Solids	Size Distribution Sieve% OnCumul. % +M +M Sp gr,Shape Ash content Grease content Permissable Flocculants Bulk density Mother Liquor Composition Dissolved: Solids-% Impurities%	FLOW	SHI

Bird Machine Company, Inc., South Walpole, Massachusetts 02071 . (617) 668-0400

Bird Machine Company, Inc., South Walpole, Massachusetts 02071 • (617) 668-0400

Disposition of cake product

Disposition of Mother Liquor (ML)_

Equipment preference

Precautions necessary

Disposal: of testing residue

Size of sample available_____

To be witnessed:

of test samples

Testing Information

□ Large scale □ Preliminary □ Field

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	1122-1	cucion	INCOLUCIO	1011
1.	Company			
2.	Address			
3	Submitted by Telephone	Date		
	· · · · · · · · · · · · · · · · · · ·			
PROCES	ESS REQUIREMENTS			
4.	Name of Product			
5.	. Production rate desiredlbs/hr ., bone dry h	oasis, or	other	
б.	 Purpose (circle those which apply): recover solids, c classify solids, recover liquor, clarify liquor, separa other 	lewater s ate 2 l:	olids, wash iquids,	ı solids
MATERI	RIAL TO BE CENTRIFUGED			
7.	. Name of solids in the feed Amount	of solid	ls	å by wt.
	Amount	of solid	ls	% by vol
8.	. Feed. Temp OF min, OF max, OF normal. Pres	sure		psig.
	Viscosity Cps at OF . Bulk density	lbs/cu % min	1 ft. pH	
٩	re feed produced continuously or by batch For	hrs/2	4 hrs.	
10	Solide characteristics (circle those which apply)		•	
10.	evologive amorphous fragile min	eral	granular	
	inflammable colloidal free draining nat	ural	hygrosc	opic
	toxic crystalline slow draining syr abrasive fibrous gelatinous org	thetic anic	slimy	
	Other	,41110		
11.	. Solid particle size distribution. Apparatus used: T	/ler, ot!	her <u>T</u> est	ed wet,
	retained on_ mesh _% by wt. retained on_ me u u u u u u	sh _*	bywt. W	
			11 11	
		-	n	
12.	. Solids specific gravity. Real App	arent		
	Liquor in feed Name & by yol Vic (Cos) SD (эн	

Liquor characteristics (circle those which apply): explosive, inflammable, toxic, foamy, heat sensitive, dermal sensitive, corrosive, other

	15.	Feed Pate gpm min, gpm normal.
	16.	Solids discharge requirements lbs/hr., bone dry basis, Residual moisture by wt% min,% max,% normal, how determinedBulk densitylbs/cu ft. Allowable crystal breakage%
	17.	Liquor discharge requirements gpm. Allowable solids% by wt.
	18.	¹⁵ washing of solids required Name of wash liquor Allowable wash tempto OF Amount Ibs wash liquor/1b solids.
	19.	Must wash liquor be kept separate from mother liquor
	20.	Are additives permitted type and amount
	21.	For liquid-liquid separation & of liquid (name) is allowable in liquid (name) Density & solids in either phase
	22.	Type of operation (circle one): Batch manual, batch automatic, continuous.
	23.	Operating requirements (circle those which apply): vapor tight, venting, inert atmosphere, pressurized. Other
D.	CENTR	IFUGE MACHINE SPECIFICATIONS & MISC.
	24.	Materials of construction for parts in contact with process material
	25.	Motor enclosure (circle one): drip proof, TER, explosion proof, other
	26.	Available power: 1 phase,voltshertz 3 phase, voltshertz
	27.	What handling precautions are necessary
	28.	Is product now being separatedMachine type Diameter & length of bowl or basketX " Cake Thickness Bowl Capacity per cycle in lbsOr Feed Rate in lbs/hr or GPM BPM Feed,; RPM Spin RPM Discharge; Feed Time;Spin Time; Wash Time;; Spin Time Discharge Time, Automatic, Semi-Automatic, Manual (circle one) What problems exist (circle those which apply): erosion, corrosion, capacity, efficiency, blinding, crystal breakage, Cake moisture, other

CENTRIFUGE APPLICATION

INFORMATION PAGE 2

D.

ROBATEL INC.

737

Centrifuge
NPAUL O. ABBÉ 🕯 🕻

QUESTIONNAIREFORMIXINGORBLENDINGEQUIPMENT

In order to help our Engineering staff recommend the correct type and <u>size</u> mixer to <u>satisy</u> your individual needs, it is essential that you supply us with as much information as possible. The answers to the questions listed here will greatly assist us <u>in making</u> a prompt, <u>intelligent</u> reply to your inquiry. All mformatron received will be held in strict confidence.

Comp	any	Department	Name of Material:	
Address .		Location of Plant	Shipped Prepaid from	
Inqu	ry initiated by	Date	Sample of Finis	
	Name (if possible) material to be mixed or give names of similar	well-know materials also bulk density	Return Tested Mater	
-			Address:	
	s this material currently being mixed by you?	If so, what type and size mixer is being used and	Other Disposition I	
	what HP 15 required?			
	Will the material be mixed in dry, liquid or paste form?			
	Should this operation be batch or continuous?			
	What size batch do you wish to mix at one time , or ho hours?	w much productron is tequired per hour, or per day of	Statement of Mixing	
	What is the approximate weight per cu. ft. or gallon of the co	ombined ingredients?		
7.	Describe the consistency or viscosity of the material			
8.	Is the material abrasive?			
9.	Will it be necessary to heat or cool the material during the m	ixing process?		
	If so, what temperature is required?			
10.	If steam heat is to be used, what pressure will be used in the	Jacket?		
11.	s the mixing to be done under pressure or vacuum? If so, to	what extent?		
			Blending or Disper	
12.	Is special dust tight construction required?		Capacity Per Hour	
13.	What clearance is desired under the discharge?		Caution If Explosiv	
14.	Should parts coming in contact with your material be made	of special materials such as stainless steel, bronze, aluminum,	Characteristics Unu	
	Monel, etc? If so, what m	aterial is preferred?	Chilling Permissible	
			Normal Moisture (
15.	Should motor be standard open, totally enclosed or explosio	n proof?	Drying Permissible	
	Give voltage, phase, and cycles of your electric current		Abrasive As Contai	
16.	Outlme any specific problems which may be involved in mix	ing this material and let us have any other mformatron which	Bulk Density:	
	you feel would be helpful to us in providing the best recomm	nendations	Method of Packing	

Name:	Title:		
Company:	Division:		
Street			
City:state:	Zip;	「elephone: ()	
Name of Material:	Amount:	Value:	
Shipped Prepaid from:	Via:	Date:	
(Name of Company,	e Vie	Data	
Between Tested Meterial to:		Date.	
(Name Of Company)	Ann:		
Address:		Amount:	
Other Disposition Instructions:			
TECHNI			
Statement of Mixing Application (Include Proportions of Ingred	lients):		
Blending or Dispersion Required:			
Capacity Per Hour Wanted -Min. and Max .:			
Caution If Explosive, Inflammable, Poisonous, Noxious:			
Characteristics Unusual as Hygroscopic, Volatile, Corrosive:			
Temperature Limitations: Softens: Me	elts:	Decomposes:	
Chilling Permissible:	Heating Permissible:		
Normal Moisture Content:	%. Oil, Fat, Etc., Cont	ent:	%
Drying Permissible to:%, A	ddition of Water Permissib	le to:	%
Abrasive As Containing Silica, Quartz, etc.:			
Bulk Density: lbs./cu. ft.; Product:		; Ingredients:	
Method of Packing and Preserving Perishable Materials:			
Solubility, and Fluids Recommended for Cleaning Equipment:.			
Power Available: Volts: Phase:		Cycles:	

MIXING TEST INQUIRY DATA

PAULO.ABBEINC.

148 CENTER AVENUE

LITTLE FALLS, NEW JERSEY

Information Request Form

For Internal use: P.P._ date_

MAIL TO ONE OF THE ADDRESSES SHOWN BELOW

I would like a preliminary estimate of the cost and performance of Prism® separators for the application described below

Note: If more than one stream is to be considered, please machine copy this form and complete the information requested for each stream

Feed stream data

a. source		D			
b. Temperature		C Press	sure		
	Maximum_				
d. Flowrate	Expected				
	Minimum_				
	component	Vol. %		Component	Vol. %
			-		
-			-		
e. Feed gas composition			-		
			-		
Desired Results:					
	Recovery		Purity	Pr	essure
permeate					
non-normeate					

4. Use/Special Requirements for the non permeate gas

NAME:			
TITLE:			
COMPANY:			
ADDRESS			
CITY	STATE.	ZIP	
COUNTRY:			
FELEPHONE ()			
TELEX:			

Monsanto Company Separations Business Group 800 No. Lindbergh Blvd. 5t. Louis, Missouri 63167 USA Telephone: (314) 694-8000 Telex: 44-7282	Monsanto Europe S.A. Separations Business Group 270-272 Av. de Tervaren 1150 Brussels, Belgium Telephone: 2/762-11-12 Telex: 62927	Monsanto Japan Limited 520 Kokusai Building 1-1, Marunouchi 3-chome Chiyoda-ku Tokyo 100, Japan Telephone: (03) 287-1251 Telem: 10214	Monsanto Far East Limited Separations Business Group 1304 Great Eagle Centre 23 Harbour Road Hong Kong Telephone: 5:740738 Telep: HX 73440	Monsanto Australia Limited Separations Business Group East Tower Princes Gate 151 Flinders Street Melbourne, Australia Telephone: (03) 658-6666 Teley: Ad 30268
		Telex: J22614	Telex: HX 73440	Telex: AA 30288



FRITZ W. GLITSCH & SONS, INC.

P. 0. Box 6227 . Dallas, Texas 75222



DISTILLATION **Process Design Data Sheet**

Item No. or Service				
Tower diameter ID	-			
Tray spacing inches				
Total trave in costion				
May A P mm Hg				
Conditions at Tray No				
conditions at may no				
Vapor to tray °F				
Pressure.				
Compressibility				
*Density lh/cu, ft				
'Rate, lb./hr.				
Cll. ft./sec., (afs)				
$cfs \sqrt{D_v/(D_L-D_v)}$				
Liquid from tray, °F				
Surface tension .				
Viscosity, cp				I
*Density, h/cu ft I			I	
Bate lb./hr.				
CPM hot liquid				
Foaming tendency	None	- Moderate	High	Severe

*These values are required in this form for direct computer input.

NOTES:

- 1. This form may be used for several sections of trays in one tower, for several towers, or for various loading cases. Use additional sheets if necessary.
- 2. Is maximum capacity at constant vapor-liquid ratio desired?
- 3. Minimum rate as % of design r a t e : %
- 4. Allowable downcomer velocity (if specified) : ft/sec
- 5. Number of flow paths or passes: ______FWG Choice: ______ Bottom tray downcomer: Total draw_______Other _____
- 6. Trays numbered: top to bottom _____; bottom to top ____;
- 7. Enclose tray and tower drawings for existing columns.
- 8. Manhole size, I.D., _____ inches.
- Manways removable: top _____, bottom _____, top & bottom _____,
 Corrosion allowance: c.s. _____; other _____; other _____; no _____;
- not required 12 Packing material if required
- 13. Tray material and thickness
- 14. Valve material
- 15. Ultimate user 16. Plant location
- 17. Other _____

Form No. PE-8

ROBATEL INC. SOLVENT EXTRACTION QUESTIONNAIRE Page 2

1. NATURE OF THE OPERATION

- 2. PHASE TO EXTRACT
 - Nature of diluent
 - . Nature of solution to extract
 - . Concentration in solution
 - . Density . Output
- 3. SOLVENT
 - . Nature of solvent
 - . Density
 - . Output

4. REQUIRED CHARACTERISTICS OF EXTRACT IMPOVERISHED IN SOLUTION

- . Concentration in solution
- . Approximate density

5. REQUIRED CHARACTERISTICS OF EXTRACT ENRICHED IN SOLUTION

. Concentration in solution Approximate density

6. NUMBER OF THEORETIC STAGES CORRESPONDING TO THE REOUIRED EXTRACTION AMOUNT

(if possible, enclose adiagram of balance or the maximum information concerning the miscibility of the solvent and of the diluent).

7. OPTIMUM TEMPERATURE OF EXTRACTION

8. SOLIDS IN SUSPENSION

Does one of the 2 phases comprise any solid in suspension before extraction? Which one?

Are these particles soluble in the other phase?

Can these particles be eliminated before introduction into the extractor?

- , by filtration
- , by centrifugal decantation
- If particles can be eliminated by centrifugal decantation state:
- . time of decantation
- . Decanter bowl diameter
- . Speed of rotation

With the two initial phases having no particles in suspension, do any precipitates appear during the process of extraction7 In this case, after centrifugation are these precipitates situated:

- . within one of the phases
- , on the surface of the light phase
- . at the bottom of the heavy phase
- . in the zone of separation between the 2 phases
- 9. SPECIAL CHARACTERISTICS

Does an emulsion appear when the two phases are vigorously stirred?

In case of emulsion:

- . If the phases separate themselves by static decantation state:
- time required
- . If the phases separate themselves by centrifugal decantation state:
- time required
- decanter bowl diameter
- speed of rotation

After decantation is the surface of separation between the 2 phases very distinct?

Does one of the 2 phases stay cloudy7

Which one?

10. DIVERSE INFORMATION

Materials of construction Electricity supply available Explosion and flameproofing requirements Type of supporting base or structure

SWENSON[®]CRYSTALLIZER CHECKLIST

CHECKLIST OF DATA REQUESTED ON CRYSTALLIZER INQUIRIES

Fill in or attach as-much information as is available so the most economic design can be prepared

1 Desident being south line d	Production		
Liquid Properties East	Kate	Liquor	Purge
Bete	mother	Elquoi	i uige
Composition			
Density			
Viscosity			
к (thermal conductivity)			
Cp (specific heat)			
Boiling point elevation			
3. Evaporation Rate			
For Organic vapors only give latent heat of evaporation/spec	cific volume/vapor	viscosity	
4. Solubility of solute IN solvent at vanous temperatures (solubilit	y curve) or phase	diagram if se	veral solutes are present
5. Product Heat	Heat	of Reaction if	Specific
Crustallization Gravity	Produc	es Crystalizer	Heat
, , , , , , , , , , , , , , , , , , ,		<i>,</i>	
6 Crystal shape if known			
7. Product SI2P required			
8. Suitable materials of construction for operating temperature	range		
BodiesTubesPumps		Vacuum	Equipmen
9. Type of seals required in adjitators or pumps and their materials	of construction		
10. Gasket material required if known			
11. Utilities Available Steam	Cooling		Current
Pressure Wa	ter Temperature	e	Characteristics
12. Describe flowsheet required or used and whether one or several	bodies are requi	ired (or the eco	phomic factors which are
to be used in determining the number of stages or effects The	se factors Include	steam cost, wate	er cost, electric@ cost,
amortization period, etc.)			1
1.3. Any other data which the customer feelswould be of help such product is produced and type of equipment used, present CNVS	h as heat transfer tal SIZC; and effe	data,present CO	nditions underwhich the additives and/or temper

14. If the solution is organic and test work is desired. give the flash point of the solution ______

ature on crystal size, shape, recovery, or purity

VULCAN IRON WORKS, INC.

1050 UNITED PENN BANK BUILDING WILKES-BARRE. PENNSYLVANIA 18701 (717) 822-2161 TELEX: 831-831

Questionnaire

INQUIRY DATA

A. PROCESS REQUIREMENTS (PLEASE FURNISH ALL RELATED INFORMATION THAT IS READILY AVAILABLE)

B. INCINERATION (THERMAL OXIDATION) WITH OR WITHOUT HEAT RECOVERY. (PLEASE FURNISH ALL INFORMATION THAT IS READILY AVAILABLE) TYPE OF PLANT AND PROCESS

WASTES TYPE OF HEAT RECOVERY THAT CAN BE UTILIZED SOLIDS QUANTITY LBS /HB SOLIDS SOLIDS WASTE TYPE STEAM GENERATION: HEAT VALUE BTU/LB SOLIDS SOLIDS ASH CONTENT % SOLIDS MOISTURE CONTENT % SOLIDS _____QUANTITY LBS /HR SLUDGES ______TYPE OF SLUDGE SLUDGES ASH CONTENT % SLUDGES ______WATER CONTENT % SLUDGES - COMBUSTIBLES CONTENT % SLUDGES -HEAT CONTENT OF COMBUSTIBLES CONTENT % HEAT CONTENT OF COMBUSTIBLES SLUDGES BTU/HR LIQUIDS QUANTITY GAL/HR ____WASTE LIQUID TYPE LIQUIDS HEAT VALUE BTU/GAL LIQUIDS LIQUIDS - VISCOSITY LIQUIDS ______SPECIFIC GRAVITY ASPHALT HEATING FUMES QUANTITY SCFM FUMES TEMP FTEMP RISE EUMES EFFLUENT TYPE FUMES - E F F L U E N T Q U A N T I T Y INDICATE ADDITIONAL INFORMATION

_____LBS./HR REQUIRED PSI OPERATING PRESSURE F. OPERATING TEMP THERMAL LIQUID HEAT LOAD BTU/HR _____HEATING STATIONS F TEMP AT STATIONS FUME PRE-HEATER (GAS TO GAS HEAT EXCHANGER) HEAT RECOVERY EFFICIENCY _____F GAS INLET TEMP HOT WATER HEATING GAL/MIN TEMP RISE

GAL /MIN

AND COMMENTS ON REVERSE SIDE



MAIL TO: VULCAN IRON WORKS, INC. 1050 UNITED PENN BANK BLDG. . WILKES-BARRE. PA. 18701 U.S.A.

TELEX: 831-831

	CENTRAL LABO	RATORY				3-1*
D E P EX corporation	TEST INQUIRY	DATA CESSING	Project No Date	FEECO INTERNATIONAL, INC 3913 ALGOMA RD. GREEN BAY. WISCONSIN 54301		AGGLOMERATIO
Company.				PHONE 414-468-1000		FEEGU
Address: Contact:				TELEX 263456		INTERNATIONAL
	MATERIAL SHIPMENT	AND RETURN		Company		
Test Material:	Lbs Of Dry Material,	L	bs Wet Feed Is Being Shipped Prepaid	Address		AGGLOMERATION
From	Via	DateCor	Date	Commercial Name of Material to be pelleti	zed	
Test Material Return Instru				Chemical Formula	Specific Heat	DATA
				CAPACITY	. (h	
				Agglomerator will operate here por dev	s./hr Feed Rate or Discharge rate	SHEET
	PURPOSE OF	TEST		s feed uniform in quantity moistu	-days per week weeks per year	•
				MATERIAL	temperature	
				Moisture Content % before	pelletizing% product	
	MATERIAL	DATA		LBS /CU FT		
Dry Material Name	Fi	inal Moisture (Wet Basis)	Particle Size Average	Smallest Larges	t Hygroscopic
Liquid Name		Chemically Free	% Combined	Furnish screen Mesh		
Spec Gravity, Solids:	Fe	eed Consistency		analysis Passing if possible %		
Loose Bulk Density, Lb/Ft ³ : -	WetDry M	eltingF, Softe	ening: ^O F, Sticking: ^O F	Material is sticky Abrasive	Corrosive Plastic Bri	ttle Dusty Heterdous
Angle Of Repose	WetDry A	brasive	_Hygroscopic	Special additive (binder) required?		
Spec. Heat:S	SolidsLiquid S	oluble (If Soluble In	Liquid Attach Solubility Curve)	Describe the process before and after pellet	izing	
Heat, FusionHydration	n:Crystalliz Pa	article Size Feed	Prod Regid Prod Allowed	POWER 110 wells and an and	40 50	
Latent Heat, Liquid	Boiling Point %			220 volt phase cycle	AC DC	
Initial Mojeture (Wet Basis)		=011		440 voit phase cycle	AC DC	
Chemically Free	% Combined %	5 - On		AGGLOMERATION EXPERIENCE		
				We have been Aggiomeration		Operating conditions
Dention I Cine	ANALTTICAL PR	UCEDURES		Capacity	iomiono steel Clather C	
Partical Size				Aggiomerator to be made of carbon steel [] si		
volatiles.	PROCESS	DATA		DELIVERY		
Production Pate	FROCESS I	Hrs/Vear :	Lbs/Hr	Drawings required Propos	al required	
	Temperature In.		0F	Information furnished by	In derivery date	
Inert Gas Required		Contact With (Combustion Products OK ?	Return to		
Quality Criteria						
From What Equipment Is Feed	d Delivered ?					
Ultimate Product Use		<u>Construction</u> Materi	als			
Present Process And Equipment	nt:					
Equipment Cleaning Method.	L				and the second	
	UTILITIE	S				
Electricity	Steam AtPsig	°F, Dowtherm	AtPsigOF	3913 ALGOMA ROAD	effenning antional	GREEN BAY, WISCONSIN 54301
Hot Oil At	PsigOF, Molten Salt At	Psig_	YF			
Fuel Gas AtBtu/	/Cu Ft, Fuel Oil At	Btu/Lbs, Cooling Wat	ter AtPsigYF		a state of the sta	
	ENVIRONM	IENT				
Plant A1Ft Abo	ove Sea Level,	YF Design Fo	r% Relative Humidity			



10225 Higgins Road Rosemont Illinois 60018 (312) 825-8010 Tel ex 25-3279

APPLICATION DATA

BRIQUETTING/COMPACTING - GRANULATING MACHINERY & SYSTEMS

Please provide as much information as is presently available, so that we may determine your application requirements. This information, which we will consider confidential, is essential to determine equipment that would be necessary for laboratory tests, budget estimates, and occasionally, to allow us to supply a firm quotation.

- 1 We are interested in:
 - () Briquetting for a definite shape and size
 - () Compaction/granulation for a granular material

Agglomerating systems in general

2 - Our immediate need is for:

Costs for laboratory evaluation

Budget estimate for economic evaluation

- () A detailed proposal and quotation, if obtainable from this information
- 3 Material to be processed

damage

Bulk density (lbs./cu. ft.)		
Specific gravity		
Screen analysis _ % + _mesh _	양	+ _mesh
* + mesh _	90	+ _mesh
Moisture% Free% Combined		
() Water () Other		
Temperature required for removal		
Maximum temperature the material can withsta	ind	without

Estimated feed temperature to the machine

Estimated maximum temperature of the feed material from the previous processing step that could be considered for machine feed

Physical nature of the material:

K-G

Rietz

	() Sticky	() Abrasive) Corrosive
--	---	----------	--------------	-------------

- () Hygroscopic () Thixotropic () Toxic
- () Aerated () Irritant

If corrosive or abrasive, state some typical materials of construction you consider acceptable with this application.

If	toxi	ic	or	irritating,	give	safety	precautions	and	the	LD50
val	ue,	if	kn	own						

- 4 The material (can () cannot be processed in contact with the gases of combustion
 - () A special atmosphere is required for processing

Specify

() The material contains combustible matter which could be considered as a source of heat if the material is heated to processing temperature by direct flame.

Combustible matter

Percentage by weight

Estimate BTU/lb.

- 5 What operation(s) will be performed immediately preceeding the agglomeration step
- Estimated production rate required () tons () pounds per hour. Operation will be () continuous () intermittent

Estimated hrs./day _____days/week _____days/yr.

- 7 Final product desired:
 - () Briquets or pellets ____ x ___ x
 - () Granular material mesh + mesh

() Other _____ Moisture (solvent) ____ that may remain %

- 8 Most materials are briquetted or compacted by pressure alone. However, in some cases, a binder must be utilized to supplement the mechanical forces required for agglomeration. The binder has to be inert and non-contaminating to the material. Please state whether this approach would be permissible with your material.
 - () Yes () No Why?
- 9 Electrical power available:

Volts	Cy	cles	I	Ηz.

ALL	motors	over	н.г.	are	τo	be	IOT		VOLT	service.
								-		

All motors are to be () TEFC () Explosion proof,

Class _____ () Open () Industrial standard

- () Other
- 10 How is the agglomeration equipment to be used.
 - () Completely new product proposed for production
 - () Installation in a current process to improve or change the material's properties
 - () Expansion of a current process using agglomeration equipment
 - () Improvement of a current agglomeration process
- 11 Why is agglomeration being considered at this time?
 - () Improve product properties
 - () Reduce dusting
 - () Improve efficiency in a current process
 - () To produce a specific size and shape
 - () Meet a competitive condition
 - () Reduce process costs

Briquetting-(continued)

U.S Occupatior	5. DE nal Sa	PARTME	I Health Administration	VIB NO. 44	ved -R1387
MATERIA	L S	SAFE	TY DATA SHEET		
Required under USDL Shipbuilding, a	. Safe nd Si	ty and He 1ipbreaking	alth Regulations for Ship Repairing, (29 CFR 1915, 1916, 1917)		
		0507			
ANUFACTURER'S NAME		SECT	EMERGENCY TELEPHO	NE NO.	
DDRESS (Number. Street, City, State. and ZIP Co	ode)				
HEMICAL NAME AND SYNONYMS			TRADE NAME AND SYNONYMS		
HEMICAL FAMILY			FORMULA		
SECTION	111 -	HAZAR	DOUS INGREDIENTS		
PAINTS, PRESERVATIVES, & SOLVENTS	%	TLV (Units)	ALLOYS AND METALLIC COATINGS	%	TLV (Units)
IGMENTS	1		BASE METAL		
ATALYST	1		ALLOYS		
VEHICLE	+		METALLIC COATINGS		
OLVENTS	+		FILLER METAL PLUS COATING OR CORE FLUX		
ADDITIVES	\top		OTHERS		
THERS					
HAZARDOUS MIXTURE	SOF	OTHER LI	LUIDS, SOLIDS, OR GASES	%	TLV (Units)
· · · · · · · · · · · · · · · · · · ·					
00					
SEC		N 111 D			
BOILING POINT ("F.)	+-		PERCENT. VOLATILE		
(APOR PRESSURE (MM H9.)	+		BY VOLUME (%)		
APOR DENSITY (AIR=1)	_		_(=1)		
OLUBILITY IN WATER	_				
PPEARANCE AND ODOR					
SECTION IV	FIR	E AND	EXPLOSION HAZARD DATA		
FLASH POINT (Method USEd)			FLAMMABLE LIMITS		Uei
XTINGUISHING MEDIA			-		
PECIAL FIRE FIGHTING PROCEDURES					
JNUSUAL FIRE AN,, EXPLOSION HAZARDS					

		SE	CTION V . H	EALTH HAZARD DATA	
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	FFECTS OF OV	EREXPOSURE			
SECTION VI · REACTIVITY DATA TABILITY UNSTABLE CONDITIONS TO AVOID STABLE Image: Conditions TO AVOID WATABOUS DECOMPOSITION PRODUCTS WATABOUS Image: Conditions TO AVOID WATABOUS DECOMPOSITION PRODUCTS WATABOUS Image: Conditions TO AVOID WILL NOT OCCUR Image: Conditions TO AVOID SECTION VIL NOT OCCUR SECTION INFORMATION SECTION SECTION INFORMATION RESPIRATORY PROTECTION ISpecial PROTECTION INFORMATION SECTION SECTION RESPIRATORY PROTECTION ISpecial PROTECTION INFORMATION SECTION SECTION RESPIRATORY PROTECTION ISpecial PROTEC	MERGENC AND	FIRST AID PROCEDU	IRES		
SECTION VI · REACTIVITY DATA TABILITY UNSTABLE STABLE CONDITIONS TO AVOID WCARDOUS DECOMPOSITION WARDOUS DECOMPOSITION WARDOUS DECOMPOSITION WARDOUS DECOMPOSITION WARDOUS DECOMPOSITION WARDOUS MAY WALL NOT OCCUR CONDITIONS WILL NOT OCCUR MAY WILL NOT OCCUR MAY WILL NOT OCCUR MAY WILL NOT OCCUR MAY SECTION VIL SPILL-OR LEAK PROCEDURES TEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED ASTE DISPOSAL METHOD EXECTION VIII SPECIAL PROTECTION INFORMATION RESETTION MCALE EXHAUST MECHANICAL (General) MOTECTIVE GLOVES EYE PROTECTION INFER PROTECTIVE EQUIPMENT SECTION IX · SPECIAL PRECAUTIONS MERE PROTECTIVE EQUIPMENT SECTION IX · SPECIAL PRECAUTIONS THER PROTECTIVE EQUIPMENT SECTION IX · SPECIAL PRECAUTIONS THER PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING THER PRECAUTIONS SECTION IX · SPECIAL PRECAUTIONS <td></td> <td></td> <td></td> <td></td> <td></td>					
SECTION VI · REACTIVITY DATA TABILITY UNSTABLE CONDITIONS TO AVOID NCOMPATABILITY (Microits to avoid) WAY OCCUR WAY ARDOUS MAY OCCUR CONDITIONS TO AVOID WILL NOT OCCUR WILL NOT OCCUR MAY OCCUR WILL NOT OCCUR SECTION VII · SPILL-OR LEAK PROCEDURES TEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED RASTE DISPOSAL METHOD SECTION VIII · SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) ENTITATION LOCAL EXHAUST MECHANICAL (General) OTHER ROTECTIVE GLOUPSES EYE PROTECTION ITHER PROTECTIVE CLOUPS EYE PROTECTION RECAUTIONS TO BE TAKEN IN HANDLING AND STORING STHER PRECAUTIONS SPECIAL PRECAUTIONS					
			25051011 1/		
	TABULTY		SECTION VI		
	ABILITY	UNSTABLE	CONDITI	ONS TO AVOID	
ALARDOUS DECOMPOSITION PRODUCTS AAAPDOUS DECOMPOSITION AAY OCCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUN AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUR AAY OCUN AAY OCUN AAY OCUN AAY OCUN AAY OCUN AAY OCUN AAY OCUN AAY OCUN AAY OCUN AAY OU AAY OCUN AAY OU		STABLE (Materials to avoid)			
WAARDOULS OLYMERIZATION MAY OCCUR CONDITIONS TO AVOID WILL NOT OCCUR Interview Interview SECTION VIL - SPILL-OR LEAK PROCEDURES SECTION VIL - SPILLEOR LEAK PROCEDURES SECTION VIL - SPILLEOR SPILLED SECTION VIL - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION Specify type ENTILATION LOCAL EXHAUST MECHANICAL (General) OTHER SECTION IX - SPECIAL PROTECTION RECOULTIONS TO BE TAKEN IN HANDLING AND STORING THER PRECAUTIONS COLSPAN= TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS COLSPAN= TAKEN IN HANDLING AND STORING COLSPAN= TAKEN IN HANDLING AND STORING	AZARDOUS DEC		CTS		
AZAROUS VII NOT OCCUR VII NOT OCCUR VII NOT OCCUR SECTION VII - SPILL-OR LEAK PROCEDURES TEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED VASTE DISPOSAL METHOD SECTION VII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) RENTILATION LOCAL EXHAUST SPECIAL OTHER ROTECTIVE GLOVES FYE PROTECTION SECTION IX - SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING PROTECTIVE				CONDITIONS TO AVOID	
WILL NOT OCCUR SECTION VII - SPILL-OR LEAK PROCEDURES STEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED VASTE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) RESPIRATORY PROTECTION (Specify type) RESPIRATORY PROTECTIVE (General) DOTACTIVE GLOVES EYE PROTECTION SECTION IX - SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS	HAZARDOUS POLYMERIZATION	MAY OCCUR		CONDITIONS TO AVOID	
SECTION VII - SPILL-OR LEAK PROCEDURES ITEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED VASIE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) VENTILATION LOCAL EXHAUST MECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION DTHER PROTECTIVE EQUIPMENT EYE PROTECTION SECTION IX - SPECIAL PRECAUTIONS SPECAUTIONS TO BE TAKEN IN HANDLING AND STORING		WILL NOT	CCUR		
SECTION VII - SPILL-OR LEAK PROCEDURES TEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED TASTE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION ESPIRATORY PROTECTION (Specify type) ENTILATION LOCAL EXHAUST MECHANICAL (General) OTHER ROTECTIVE GLOVES EYE PROTECTION SECTION IX · SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING THER PRECAUTIONS THER PRECAUTIONS THER PRECAUTIONS					
TEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED VASTE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) RENTILATION LOCAL EXHAUST DECIAL DECHANICAL (General) OTHER RETECTIVE GLOVES EYE PROTECTION SECTION IX - SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING FRO 054.110		SECT			
VASTE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) PROTECTION (Code EXHAUST SPECIAL OTHER PROTECTIVE GLOVES EYE PROTECTION THER PROTECTIVE EQUIPMENT SECTION IX - SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING PTHER PRECAUTIONS THER PRECAUTIONS PTHER PRECAUTIO	TEPS TO BE TAP	SECTI Ken IN Case Materi	AL IS RELEASED O	R SPILLED	
VASTE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) PROTECTION LOCAL EXHAUST SPECIAL DECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION SECTION IX - SPECIAL PRECAUTIONS PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING PROTECTIVE FYO 054.110				-	
ASTE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) ZENTILATION LOCAL EXHAUST MECHANICAL (General) COTHER PROTECTIVE GLOVES EYE PROTECTION SECTION IX - SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING COTHER FRO 054.110 COMPARING OMPARING COMPARING COMPARING COMPARING COMPARING COMPARING COMPARING COMPARING COMPARIN					
ASSTE DISPOSAL METHOD SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) RENTILATION LOCAL EXHAUST MECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION SECTION IX - SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING PTHER PRECAUTIONS COLUMN					
SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) rentilation LOCAL EXHAUST MECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION THER PROTECTIVE EQUIPMENT SECTION IX - SPECIAL PRECAUTIONS PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS	VASTE DISPOSAL	METHOD			
SECTION VIII - SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) rentilation LOCAL EXHAUST MECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION THER PROTECTIVE EQUIPMENT SECTION IX - SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS					
SECTION VIII SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) ENTILATION LOCAL EXHAUST MECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION THER PROTECTIVE EQUIPMENT SECTION IX · SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING STHER PRECAUTIONS					
SECTION VIII . SPECIAL PROTECTION INFORMATION RESPIRATORY PROTECTION (Specify type) rentilation LOCAL EXHAUST MECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION SECTION IX · SPECIAL PRECAUTIONS PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING					
		SECTION	VIII . SPECIA	L PROTECTION INFORMATION	1
EVENTILATION EDUCAT EARNOST MECHANICAL (General) OTHER PROTECTIVE GLOVES EYE PROTECTION SECTION IX · SPECIAL PRECAUTIONS RECAUTIONS TO BE TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS FRECAUTIONS RECAUTIONS FREC	RESPIRATORY PRO	STECTION (Specify i)	'pe)		
	ENTILATION	EOCAL EXHAUST		SPECIAL	
PROTECTIVE GLOVES EYE PROTECTION EYE PROTECTION SECTION IX · SPECIAL PRECAUTIONS PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING THER PRECAUTIONS FIG. 054.110 COUNTY OF C		MECHANICAL (Ge	heral)	OTHER	
SECTION IX · SPECIAL PRECAUTIONS PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS	PROTECTIVE GLO	DVES		EYE PROTECTION	
SECTION IX - SPECIAL PRECAUTIONS PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS	THER PROTECTI	VE EQUIPMENT			
SECTION IX - SPECIAL PRECAUTIONS PRECAUTIONS TO BE TAKEN IN HANDLING AND STORING OTHER PRECAUTIONS EPO 054 110 PO 054 110 EPO 054 110 EPO 054 EPO					
CTHER PRECAUTIONS		S	ECTION IX · S	SPECIAL PRECAUTIONS	
CP0 054 110	PRECAUTIONS TO	BE TAKEN IN HAND	LING AND STORIN	G	
CP0 054110					
CP0 054110	THER PRECAUT	IONS			
GF0 054.110					
			GP	0 054.110	Ec

Material Safety Data Sheet-(continued)

Index-

Absorption, 371-357 column assembly, 371 Acetic acid/MIBK/water extraction, 468 Acetic acid purificatión, 547 Acetic anhydride reactor, 573 Acetone/methanol equilibria, 416 Acetone/water equilibria, 423 Acetonitrile azeotropic drying, 422 extractive distillation solvent, 416,419, 420,422 Activity coefficient, 372 solubility parameter, 374 Wilson equation, 374 Adiabatic saturation temperature, 234 Adiponitrile reactor, 576, 577 Adsorbents data, 496 manufacture, 510,511 manufacture flowsketch, 498 Adsorption breakthrough curves, 506 design example, 505 desorption profiles, 502,503 effluent profile, 504 gases, 511, 513 liquid phase, 506, 513,515 mechanism, 501 MTZ (mass transfer zone). 500. 501 multicomponent data, 503 operating cycles, 502 502 operating parameters, operating practices, 504 packed beds, 500-504 regeneration, 502, 504 regeneration steam, 502 Adsorption equilibria, 495,497 binary mixtures, 499,500 heat of adsorption, 500 isotherms, 499 liquids, 500 temperature effect, 499 Adsorption equipment, 510, 511,513 AHR process (Union Carbide), 511 drying system flowsketch, 508,511 fixed beds, 509 fluidized beds, 513 gases, 511,513 hypersorber moving bed, 513,516 liquid phase process, 513, 518 moving beds, 513 Nofsinger moving bed, 513,517 pulsed bed process, 518 sizing example, 505 supports for beds, 510 Toray process, 511,517 UOP simulated moving bed, 515, 517-519 Agglomeration, 351 binders, 354. 355 351 products, 354 Agitation. See also Mixers flow patterns, 291 HP and rpm, 292 performance, 294 power consumption, 292

Air classifiers, 338 Air coolers, 194-199 heat transfer, 196-198 heat transfer example, 199 sketches, 175 Air leakage, vacuum systems, 164, 165 other gases, 165 Air properties, 674 Air separation process, 37 Air-water interaction. 231-234 packed towers, 277-280 Alumina by calcination, reactor, 593 Ammonia absorption refrigeration, 226, 229 flowsketch, 277 Ammonia oxidation reactor, 573 Ammonia synthesis flowsheet, 27 reactors, 572, 581-584 performance data, 582,583 temperature profiles, 582, 584 Angle of slide, data, 677-679 ASME pressure vessel code, 625 Autogenous grinding, 342 Axial flow comressors 146.150.153 application range, 149,156 characteristics, 150 figure, 147 Axial flow pumps, 134, 136, 140 application range, 150 Azeotrope separation, 387,388,420-426 Azeotropic distillation, 420-426 acetonitrile/water separation, 422 commercial examples, 421-424 design method, 424 n-heptane/toluene/MEK process, vapor-liquid 424 424 process, vapor-liquid equilibrium data, 421, 423, 425,426 Ball mills, 342 closed circuit operation, 345 Denver, 342 Hardinee. 346. 347. 351 pebble type, 343,347 Batch distillation, 390 chlorinated phenols, column profiles, 383.384 constant overhead composition, 390 constant reflux ratio, 390 instrumentation diagram, 393 material balances, 391 McCable-Thiele diagram, 392 operating profiles, 393 Batch reactors, 557 Belt conveyors, 76 arrangements, 82,83 sizing calculations, 83 sizing data, 81, 82 Belt filters performance data, 328 sketch, 325 Benzene dealkylation process, 28,29 Berks ring dryer, 265 performance, 264 Binary distillation, 379

algebraic method, 382,387, 389 algebraic method example, 392 azeotropic mixtures, 387,388 batch. 383.390-393 constant molal overflow, 380 material and energy balances, 380 McCabe-Thiele diagram, 381 model sketch, 380 multiple feeds and products, 386 packed towers. 398-404 partially miscible liquids, 388, 389 q-line, 381 unequal molal latent heats, 382,385, 390 Binders for agglomeration, 354,355 Bingham liquids, 104, 105, 107-109,711 Biochemical glossary, 657 Blasius equation, friction factor, 69 Blenders, in line, 300-302 sketches, 302 power, 300 Blenders, powder, 301,302 Blowers, 143 application range, 149 two-lobe, 148 Boiling, 187. See also Reboilers Booklists bibliographies, 16 data collections, 16 encyclopedias, 16 equipment, 16 essential, for process design, 15 estimation of properties, 16 safety aspects, 16 special data collections, 17 Brine electrolysis, 648, 653, 656 Hooker cell, 648 mercury cell, 653 Briquetting, 354 gear, 358 integrated equipment, 358 product shapes, 362 rolls, 358 Brodie crystallizer-purifier, 545, 546 Bubblecap trays, 428,430-433 allowable vapor rate, 432 comparisons with other kinds, example, 431 Jersey Critical correlation, 432 liquid gradient, 433 Souders-Brown correlation, 43 Bubblepoint conditions, 375, 376 432 calculation diagram, 379 example, 378,379 Bubblepoint method, multicomponent distillation, 408 algorithm flowsketch, 406 Bucket elevators, 78, 86 Butadiene solubility, 420 vapor-liquid equilibria, 420 Butadiene sulfone manufacturing process, 35 Butanol/ethanol equilibria, 375 Butanol/water separation, 388 Butinediol reactor, 576

Buttner-Rosin pneumatic drver, 265 performance, 264 Butyl cresol purification, 547 Caprolactam hydrogenation. 576.577 Cartridge filters, 319 applications, 323 particle recovery range, 323 Catalyst bed support modes, 587 Catalysts effectiveness, 555,565-567 heterogeneous, 562 homogeneous, 562 industrial examples, 550-553 kinds, 563 organic reactions, 563 physical properties, solids, 564,565 pore tortuosity, 565 porosity, 564,565 selection basis, 562 surface area, 564, 565 Catalytic cracking reactors fluidized bed, 579,591,592 moving bed, 588 temperature and composition profiles, 595 transfer line type, 580, 581, 591 zeolite catalyst type, 591 Catalytic reformers, 572, 586 Catalyzed organic reactions, 563-564 Cement kilns, 576,590 Centrifugal compressors, 145 application range, 149, 150 cross section, 147 selection, 161 specification form, 66 specifications 152 Centrifugal pumps, 117-143,134,147 application range, 140,143 capacity-head range, 138 characteristic curves, 58 costs. 138 diffuser type, 139 double suction, 136, 139, 143 efficiency, 134, 137, 141, 144 glossary, 166 good qualities, 141, 142 impeller types, 140 parallel operation, 133, 147 seals, 137. 142 series operation, 133, 147 sinele suction. 136. 139. 143 ciscosity effect, 145 volute type, 139 Centrifuges, 321 data. 333 filtering types, 329 selection criteria, 307, 334 Chemical reactor control, 53-58 Chemical reactor operating patterns CSTR (continuous stirred tank reactor), 555,557 design basis, 549 material and energy balances, 555-559 non-flow, 557 packed bed, 559 PFR (plug flow reactor), 558 residence time distribution, 556 segregated flow, 561 Chemical reactors, 549-610 Chlorinated phenols separation, 383

Chloroform/acetone/MIBK equilibria. 416 Choking velocity, pneumatic conveying, 119 Chromatographic separations. 510,520 chromatograms, 520 economic data, 516 example, 125 mm column, 521 example, pinenes separation, 521 flowsketch 521 Circulating gas or solid reactions, 596 Clarifiers, 315, 320,338 performance, 320 Closed circuit grinding, 341,345, 351 Coal carbonization. 20 gasifier. 593 liquefaction. 609 slurry pipeline, 70 Coalescence, 613, 615 Codes and standards, 2.3 Coefficient of performance (COP), 224 Cokine, fluidized bed, 591 Colebrook equation, friction, factor, 94 Combustion gas turbine, 65, 68 arrangements, 67 performance calculation. 67 Compact exchangers, 178,194, 195 Compaction 351 Composition control, 40, 43 Composition profiles. See Temperature profiles, reactors Compressibility factor, 162 Compressibility, filter cake, 310, 311,316 cell measurements, 314 calculation example, 314 data. 316 Compression of gases, 153 efficiency, polytropic, 158, 159 friction. 156 ideal gases, 153, 155 isentropic, 155, 156 mixtures, 157 multistage, 161, 164 non-ideal, 157, 160 polytropic, 156 ratio, multistage, 161, 164 temoerature rise, 159, 162 thermodynamic diagram, 157 variable heat capacity, 157 Compression ratio, rule, 161 Compressors application ranges, 149, 150 characteristic curves. 59 control, 59,60 efficiencies, 158 operating ranges, 150 types, 145,147-149 Condensation, 187 Condensers configurations. 204 control, 46 design method, 205,206 heat transfer coefficients, 196 partial, model, 206 sizing example, 207 Conduction, thermal composite walls, 170 Fourier equation, 169 hollow cylinders, 170 metal walls, 182 Conductivity, thermal, data, 170

packed beds, 587.599~601 Control valves, 129-131 Convection. forced. 186 Convection: natural, 186 equations, 189 Conveyors, mechanical, 76-88 Cooling towers, 280-285 approach to equilibrium, 234 bid data required, 284 kinds of fill 282 performance curves, 284 sizing example, 281,282 sketches, 283 types, comparison, 280,282,285 water loss, 285 Cost indexes, inflation, 669 Costs of equipment, 663 alphabetical index of eauinment. 664 distillation tower example: 663 inflation indexes, 669 installed cost multipliers, 668, 669 purchased and installed cost, example, 663 Cracking furnace reactor, 574 Cracking, gas oil, 33 Cryogenics, 229 Crystal growth rate, 531,533 Crystallization conditions for crystal formation. 528 crystal growth rate, 533 design example, 524 ideal stirred tank model, 533-539 melt type, 523,543-548 MSMPR model, 533-539 size distribution, 525 solution type, 523 Crystallization data literature, 535 Crystallization equipment, 537-543 APV-Kestner, 542,543 batch. stirred and cooled. 539. 540 Brennan-Koppers, 545,547,548 Brodie, 545,546 circulating coolers, 539, 540 circulating evaporators, 539, 540-542 direct refrigeration, 540, 541,543 DTB (draft tube baffle), 540,542 Escher-Wyss, 542,543 Kureha, 545, 547 MWB (Metallwerk Buchs) process, 543, 545 Oslo, 541,543 scraped jacketed pipe, 538, 540 sugar vacuum pan, 540,541 Swenson-Walker, 538,540 TNO, 545,547 Tsukushima. 542,543 twinned, Nyvlt, 541,543 Crystal size distribution, 525,530,532,539 distribution, from seeded tank, 534 from commercial equipment, 532 Crushing. See also Size reduction product size distribution, 337 CSTR (continuous stirred tank reactor), 555.557 comparison, complex reactions, 569 comparison with PFR, 571 comparison with segregated flow, 561 Cumene synthesis, 56,57 Cyclone separators, 616 calculation, example, 621 dimensions, 620

drum with tangential inlet, 620 performance, 621 pressure drop, 617 Diaphragm pump, 142, 143 Deentrainment commercial equipment, 619 cyclone dimensions, 620 cyclones, 616,620,621 empty drums, 615,617 wire mesh nads. 615.616.618.620 Densities, bulk, data, 677-679 Desalination, 632 performance, 640 Design basis, 12 questionnaire 13, 14 Detergent manufacture, 35 Dewpoint conditions, 376 calculation, 378 Dichlorbenzene purification by crystallization, 547 Diffusion equimolal counter, 399 stagnant film, 399 Dimensionless groups heat transfer, 182 mixing, 290,291 table, 188 Direct contact heat transfer, 185 Dispersion model, 560-562 first order reactions, 561 second order reactions, 562 Distillation, 371-457 batch, 390 binary, 379 column assembly, 371 flash, 375 multicomponent, 393 Rayleigh, 378 Distillation, petroleum, 37 Distillation, simple. See Rayleigh Dodecylbenzene sulfonate reactor, 576 Dorr classifier, 338 Double pipe exchangers, 195 Droplet sizes, 613 range, 631 Drum dryers, 254,255 performance, 261-263 sketches, 260 system, 260 Drums, 611 capacity, partially full, 626, 627 compressor surge, 612,613 design example, 628 dimensions, 611 gas-liquid separators, 613, 620 holdup, 612 liquid-liquid separators, 612 reflux, 612 Dryers batch, 241,247,248 belt. 242.252.253 belt conveyor, calculation, 239 classification. 237. 241. 242. 244 continuous tray, 242,247,251 costs, 240 drum, 254, 255, 260-263 evaporation rates, 243 fluidized bed, 262-272 paddle and ribbon, 249 pan, 250,251 performance, comparative, 243

pneumatic conveying, 237,255,263-267 products, 240,245 residence time distribution, 245 rotary, 247,251, 254,256-259 specifications, 240,246 spray, 237,268-279 tumbler, 249, 250 tunnel, calculation example, 238 vacuum, 248 Dryers, pilot plant sizes fluidized bed. 237 pneumatic conveying, 237 rotary, 237 spray, 237,278 Wyssmont tray, 237, 251 Drying rate, 234,237 constant period, 235, 237 data, 235,236 falling period, 235,237 pilot plant testing, 237, 251, 278 DTB (draft tube baffle) crystallizer, 540, 542 Dualflow trays, 426 Economic analyses, 4 list of published cases, 4, 5 optimum efficiency, Linnhoff, 6 waste heat recovery, 11 Effectiveness, catalyst, 555 ammonia synthesis, 567 sulfur dioxide oxidation, 567 Thiele modulus, 566 Effectiveness, heat transfer. See F-method Efficiency extraction equipment, 487 sieve tray extractors, 483 Efficiency, compression isentropic, 155 polytropic, 159 volumetric, 161 Efficiency, distillation trays, 397, 439ff Efficiency, packed towers, 436,437 442, 449,452-456 Efficiency, tray, 439 AIChE method, 439 Bakowski method, 442,451 Chu et al. method, 442,451 data in terms of vapor factor, F, 448 F factor, 429,448 McFarland et al. method, 439,442,451 O'Connell method, 439,450,451 survey of data, 444-447 vapor rate and, 448,449 Electrochemical synthesis, 645 cell types, 648,653,654 energy requirements, 653 fuel cells, 646, 655 overvoltage, 654 reactions, 646 Electrodialysis, 508, 517 equipment, 513, 514 performance, brackish water, 515 Emissivity of gases, 220 Enerev balance in fluid flow. 92 mechanical 93 units, example, 94 Engineering manhours for projects, 2 Enthalpy residual, 158, 159 Enthalpy-concentration chart, 524 application example, 529 construction example, 390 distillation diagram, 391

distillation equations, 380 some salt solutions. 529,530 Entropy residual, 158, 159 Enzymes, 650 examples, 650 production, 658 Erlang distribution, residence time, 558 Equation of state, gases, 91 density calculation, 91 Ethanol/acetic acid separation, 385 Ethanol/butanol equilibria, 375 Ethanol/isopropanol/water separation, 421 Ethanol/water separation distillation. 387, 394, 424 reverse osmosis, 642 Ethylene manufacture process, 40 purification, 40,47 specifications, 40 Ethvlene reactor, 573.574.588.593 circulating sand, 593 flame reactor, 573 Evaporators, 208-211 backward and forward feed, 211 heat transfer coefficients, 211 sketches, 209-211 thermal economy, 210 Extended surfaces, heat transfer, 188 calculation example, 193 sketches, 178 Extraction equipment, 476-490 centrifugal, 484,487,490 comparison of types, 477 mixer-settlers, 477, 479 packed towers, 478,480,482,485-487 performance comparison, 478 pulsed towers, 481,483,487 RDC (rotating disk contactor), 482, 487 reciprocating trays, 481 rotary, agitated, 482,485-487 sieve tray towers, 480,486,487 spray towers, 478,480,486 Extraction, liquid-liquid, 459 countercurrent, 466,470 464.468 crosscurrent, dispersed phase selection, 476 eauilibria. 459-463 extract rekux, 468,470,472 immiscible solvents, 464, 469 minimum reflux, 467,469 minimum solvent/feed ratio, 468,471 minimum stages, 467,469 model, 469 multicomnonent, 473-476 single stage, 463; 468 staee reauirements. 463-472 412, 417-422 Extractive distillation, additive selection, 412, 414 ethanol/isopropanol/water process, 421 examples of processes, 417 isoprene recovery, 36,417,422 McCabe-Thiele diagram, 422 methylcyclohexane/toluene/phenol process, 421 vapor liquid equilibria, 416, 418-420 Extrusion, 358 ring, 358,366 ring applications, 365 screw, 360, 366

F, tray capacity factor, 429,448

F-method, heat transfer, 173, 175-177 example, 180 formulas. 179 Fans, 143 application range, 149 blade shape, 152,153 controls, 151 efficiency, 151, 153 laws of, 151 performance, 151,152 Feed tray location, distillation, 397 Kirkbride equation, 397 Feedback control, 39,41 Feeders, granular solids, 45, 83, 87,88 Fenske minimum trays, 382, 395 Fermentation, 648 characteristics, 649 eauinment sketch, 659 flowsketches, 658 operating conditions, 650,652-654 process types, 650 products, commercial, 657 reactors, 654,659,660 Film coefficients, heat transfer, 180 data, 186 Filter cakes compressibility, 310, 311, 314,316 permeability, 315 porosity, 315,316, 318,319 resistivity, 313-315,317, 319 Filter media, 313 porosities and permeabilities, 315 Filters, pressure, 319 commercial sizes, 323 Filtration constant pressure, 306 constant rate, 306 data sheet, testing, 322 example, with centrifugal pump, 311 laboratory testing, 317,318, 321 scaleup, 317,318 SCFT concept, 317 test data, example, 310 theory, 306 Filtration equipment application and performance, 320, 321, 330 belt, 319,325,328 double drum, 319,327 horizontal rotary, 319, 325,327 Kelly, 307,319,323, 324 leaf, 319,323 plate and frame, 319, 323, 324, 331 rotary disk, 307,319, 327 rotary drum. 319. 326-328 Sparkler, 324 Sweetland, 319,323-325 Vallez. 319.323-325 Fired heater reactor, 574, 575 Fired heaters, 211 box size, rule, 213 design example, 217-220 design procedure, 214-217 efficiency. 213-214 heat fluxes and temperatures, 213 peak temperatures, 214 sketches, 212 Fittings, pipe, resistances, 95, 98-100 Fixed bed solid catalysis, 596 Flame reactor, 573 Flash conditions, 375-377 example, 378

Ks dependent on composition, 377 Flight conveyors, 81 Flocculants, 309 Flocculation, 351 Flooding, packed towers, 433,441 liquid-liquid, 485 Flow control fluids, 58,59 solids, 43,44 Flow number, agitation, 290 limiting values, 293 Flowsheets example descriptions, 33-37 examples, 20,26-30 mechanical (P & I), 21-25 process, 19 process, checklist, 20 process descriptions, list, 33 symbols, 21-25 Pluidization, 120-126 bed expansion, 121,126 characteristics, 123 freeboard in vessel, 126 kinds of particles, 124 minimum bubbling rate, 121, 122, 126 minimum rate, 121, 122, 126 regimes, 123 TDH (transport disengagement height), 121,122 vessel dimensions, 125 viscosity, 126 Fluidized bed agglomeration, 362 performance data, 368 sketches, 369 spouted bed, 363, 368,369 Fluidized bed catalysis. 601-604 Fluidized bed dryers, 262-265, 267, 268 gas velocity, 267 performance, batch, 270 performance, continuous, 271.272 sizing, example, 272 sketches, 268,269 thermal efficiency, 267 Fluidized bed processes, 577 Fluidized bed reactors, 579,580,582, 591, 592 control, 55, 58 ebullating beds, 593 mechanism. 580 multistage, '593 noncatalytic, 582, 593 operating data, 579 Fluidized beds, non-catalyzed solids, 595 Foam separation, 635 data, 644 equipment, 644 Fouling factors, heat transfer, 180 data, 183, 184, 186 Fourier equation, 169 Fractionation. See Distillation Fractionator conntrol lower ends, 49,50 upper ends, 51,52 Freeze drying, 639 cvcle lengths. 639 products; 646 Friction, 93 Friction factor, 92 Colebrook equation, 94 granular beds, 117 non-Newtonian fluids, 109 Rounds equation, 94

Schacham equation, 94 von Karman equation, 95 Froth flotation, 636,638 equipment, 645 performance, 638 Fuel cells, 646 characteristics, 655, 656 Fugacity coefficient, 372 Gas dispersion, 296,297,299,300 design example, 301 flooding, 299 mass transfer coefficients, 299 Gas flow in pipe lines, 109 adiabatic, 110-112 isentropic, 109 isothermal, 110, 112 non-ideal, 111, 113 Gas-liquid reactions, 595 with solid catalysts 604.605.607.609 Gas-liquid-solid'reactions, 609 Gas-liquid separators, 613 deentrainers, wire mesh, 615,616 droplet sizes, 613 drum with tangential inlet, 620 empty drums, 615 entrainment, 613,617 example, empty drum, 618 example, sieve tray, 617 kev dimensions, 617,618 sieve tray correlation, 618 Gas oil cracking, 33 Gas separation hydrogen enrichment, 633 membrane processes, 633 Gas-solid flow. See also Pneumatic conveying choking velocity 119 pressure drop 119.120 Gas treating plant 36, 36 Gauges of plates, sheets and wires, 676 Gear pumps, 134,141,143 GHSV, 549 Gilhland correlation, trays, 397 Molokhanov equation, 397 Globulation, 351 Glossary biochemical, 657 centrifugal pumps, 166 Gradient, liquid, bubblecap trays, 433 Graesser extractor, 484 Granular beds. See Packed beds Granular materials bulk densities, 78-79,677-679 angle of inclination, 78-79 angle of repose, 78-79,677-679 Granulators, 352 applications, 355 capacity and power needs, 356 pans, 352 performance on cement, 356 products, 355 rotating dish, 353,355,359 rotating drum, 353, 355 size stratification. 355 tumbling, moisture requirement, 356 Gravitational constant, 91, 92 Hand correlation, liquid-liquid equilibria, 463,465 Heads, vessel design example, 628

formulas, partially full, 627 thickness, formulas, 625 types, 627 Hearth reactors. 578,590 Heat exchangers. 169-229 control. 44-47 types, 188 Heat transfer coefficients film convection and radiation. 189 equations, 189-192 Heat transfer coefficients, overall, 170 air coolers, 196-198 condensers, 196 data. 183-185 range of values, 180 Heat transfer, direct contact. 185 Heat transfer, fluidized beds, 589, 592 data. 606-608 experimental work survey, 608 horizontal tubes, 609 submerged coils, 606 vertical tubes, 606,607, 609 walls, 606, 608 Heat transfer media, 172 Heat transfer. packed beds. 578.600.601 at the wall 599 between particles and fluids, 599 overall coefficient, 603,606 thermal conductivity, 600, 601 Heat transfer, reactors, 582 between particle and fluid, 587 fixed beds. 578 fluidized beds, 589,592,606-609 immersed coils, 597 jacketed vessels, 597 overall coefficients, 602,603 stirred tanks, 586, 594, 598, 599 walls. 587.589.599.606 Heat transfer, units of quantities, 188 Heaters, fired. See Fired heaters Heptane/toluene/MEK separation, 424 HETP (height equivalent to a theoretical plate), 442 correlations, 448 data, 453-456 Hickman still, molecular distillation, 427 HIGEE fractionator, 399 Homogeneous gas reactions, 592 HTU (height of a transfer unit), 401,404, 422 Cornell et al. correlation, 449 data, 452, 454, 456 liquid-liquid extraction, 484 Humid volume, 234 Humidity, 231 relative. 231 Hydrochloric acid electrolysis, 656 Hydrocyclones, 307,320, 339 liquid, 613 performance graph, 340 sizing example, 341 sketches. 330 Hydrofining reactor, 587 Hydroformer, fluidized bed, 591.592 Hydrogenation reactor, 576 Impellers, agitation kinds, 288-290 location, 288 size. 288 sizing example, 293, 294 Impellers, centrifugal pumps, 140

Information sources, 2 Installation cost factors, 668, 669 Insulation, 219 economic thickness, 220,224 high temperature, 222,223 low temperature, 221, 222 medium temperatures, 222 Ion exchange, 506-508 desien practices 506.508 equilibria 497.499, 500 equipment, 508,512,513 membranes, 517 operating practices, 506, 508 properties of materials, 507 selectivity example, 503 selectivity scales, anions and cations, 507 Ion exchange equipment, 508,512,517 continuous processes, 515, 519,520 fixed bed arrangements, 512 performance, Uranium recovery, 515 sizing example, 513 Isoprene recovery, 36,417,420,422 solubility, 420 vapor-liquid equilibria, 420 Isotope separation, thermal diffusion, 649-651 Jet compressors, 143 K-chart, 372 Kelly filter, 307,319, 323, 324 Kenics blender, 302 Knudsen diffusivity, 564 Kremser-Brown formula, 398,399,466 Laminar flow, 92 non-Newtonian, 108, 109 Leaching, 470-473, 491-493 battery, 493 Bollman bucket type, 491 Bonotto tower, 491 continuous equipment, 491 equipment, 488,491,492 example, calculation, 472 Hansa-Muehle bucket type, 491 Hildebrandt tower, 491 settling tanks, 492 Level control, 43,44 Lewis-Matheson method, distillation. 404 LHSV. 549 Linde trays, 430 Liquid-liquid equilibria, 459-463 distribution coefficient, 460 Hand correlation, 463, 465 Janecke coordinates, 460.461 tielines, 460 Liquid-liquid extraction, 459-493. See also Extraction control, 50,53-55 Liquid-liquid reactions, 595 Liauid seal ring compressors 149. 153 heating effect, 153' specifications, 155 Lockhart-Martinelli correlation, 115, 116 McCabe Delta-L law, crystallization, 533 calculation example, 534 McCabe-Thiele diagram, distillation, 381 Magnesium sulfate/water diagram, 526, 529

Mass transfer coefficient, 399.442 data, 454,455 gas dispersion, 299 Material and energy balances, 3-5 distillation, 391, 401, 405 Maximum mixedness, 560 volume ratio to segregated flow, 560 Zwieterings equations, 560 Melt crystallization. 543-548 Brenhan-Koppers purifier, 545,547, 548 Brodie crystallizer-purifier, 545, 547 Kureha purifier. 545.547 multistage, 543 MWB process, 543,545 Phillips process, 544-546 Schildknecht column, 543,544,546 TNO bouncing ball process 545.547 Melt purification. See 'Melt crystallization Membranes, 631,632,641 applications, 632 cellulose acetate, 635 equipment configurations, 632 gas permeation.633.644 hollow fiber, 632,633,641,643 performance, 646 Permasep. 641 olate and frame. 638 Prism, 633,643 properties. 635.636 structures. 632 tubular. 638.639 types, 635, 636 Merkel diagram. See Enthalpyconcentration chart MESH equations, 405 Methanol synthesis. 580.585 Methanol/water separation, 390 Methylcyclohexane/toluene/phenol separation, 421 Methylethylketone/water equilibria, 42 Microbial processes. See Fermentation Mixers. See **also** Agitation 423 blend time, 290 dimensionless groups, 290 gas dispersion, 296-301 in line type, 300,301 powders and pastes, 301,303,304 power number, 290-292 quality characterization, 290-292 suspension of solids, 295-299 tank desien. 287.288 Moisture c&tent, critical, 237 Molecular distillation, 425-427 equipment sketches, 427 Hickman still, 427 operating conditions, 426 rate of evaporation, 426 Molecular sieves applications, 496 capacity decline with use, 506 properties, 496. 497 Molokhanov equation, distillation. 397 Motors applications, 62 relative costs, 61 types, 61 MSMPR crystallization model. 533-539 MTBE (methyl tertiary butyl ether), 597 MTZ (mass transfer zone), adsorption, 500,501

Multicomponent distillation, 393 absorption factor method, 398 azeotropic. 420-426 bubblepoint (BP) method, 406-409 computer program references. 404 concentration profiles, 394 distribution of non-kevs. 395 Edmister method, 398,399 extractive, 412, 417-422 feed tray location, 397 free variables, number of. 395 Lewis-Matheson method; 404 MESH eauations. 405-407 molecular, 425-427 nomenclature. 405 number of theoretical trays, 397 packed towers, 433-439 petroleum, 411-415 reflux, minimum, 397 reflux, operating, 397 SC (simultaneous correction) method, 408-411 sequencing of columns, 393, 394 short cut design example, 396 SR (sum rates) method, 407-409 Thiele-Geddes method, 404,407 trays, minimum number, 395 tray towers, 426-433 Wang-Henke method, 406-409 Multicomponent extraction, 473-476 calculation procedure, 474, 477 example, 476 material balance, 474 Multipass heat exchangers, 173,175-177 Multiple feeds and products, distillation, 386 MWB (Metallwerk Buchs) crystallization process, 543,545 Naphthalene purification, crystallization, 547 Naphthali-Sandholm method, 404 algorithm flowsketch, 411 Nitric acid reactor, 576 Nitrogen fixation, 574,578,588 Nitrotoluene isomers separation, 544 Noncatalytic reactions with solids, 595 Non-Newtonian liquids, 100, 103-109 Bineham. 104.105.107-109 dilatant, 103, 104 laminar flow, 108,109 pressure drop in lines, 106, 109 pseudoplastic, 103, 104 rheopectic, 104,105 slurries, 71 thixotropic, 104-106 viscoelastic, 105, 106 Notation, 672 NPSH, pumps, 133,146 centrifugal pumps, 146 positive displacement pumps, 134, 135 various pumps, 144 NRTL equation, 475 NTU (number of transfer units), 401, 402, 404 air-water interaction, 277 Octane/toluene/phenol equilibria, 416 Olefin production, 33,40 Orifices, flow through, 95 Osmosis, 631 equation, 631

Osmotic pressure calculation example, 633 concentration effect, 632 equation, 631 maxima in industrial operations, 631 molecular weight effect, 632 Packed beds, 117-119 friction factor, 117 permeability, 117 porosity, 177 supports in vessels, 574, 587 thermal conductivity, 587, 599-601 two-phase flow, 118, 119 Packed bed reactors, 559,572 support of catalyst bed, 587 Packed towers, 433 allowable Row rates, 438,440, 441 efficiency, 436,437,442-456 flooding, 433,438 internals, sketches, 434 kinds of packing, 433,435 liquid distribution, 434, 439 liquid holdup, 439 packing factors, 438 packing size selection. 433 performance comparison, example, 441, 442 pressure drop, 437-440,443 structured packings, 435,437 Packed towers, air/water interaction, 277-280 HTU data, 280 NTU, 277 tower heights, 279 Packed towers, extraction, 478-485 capacity, 484,487 efficiency, 487 flooding, 485 sizing example, 486 Packed towers, separations in, 398 absorption or stripping, 401, 402 absorption example, 403 distillation, 401 distillation example, 402 equimolal counterdiffusion, 399 mass transfer coefficients, 399 mechanism, diagrams, 400 stagnant film diffusion, 399 Packing factors, 438 Pall rings, 435 capacity and pressure drop, 440,443 Parametric pumping, 639,640, 642 cycles, 648 data, 647 schematic, 647 Partially miscible liquid distillation, 388 Particle size classification, air, 337, 338 classification, wet, 338,339 distribution, 337 351-369 enlargement, measurement with sieves, 336 range, 631 reduction, 339-350 surface average, 124 Paste blenders. 301. 303 Pebble heater, 575, 578, 588 Peclet number, 558,559 Peristaltic pump, 142, 143 Permsep membranes, 641

flow rates, 631

Petroleum distillation, 48, 411-415 design data, 415 flowsketch of crude distillation, 414 linear velocities, 415 overflash, 415 pressure drop, 415 pseudocomponent mixtures, 412,413 refinery block diagram, 412 stripping steam usage, 415 TBP (true boiling point) curve, 409, 412.413 tray requirements, 415 Petroleum products compositions, 680 Petroleum properties correlations, 413 Petroleum refinery flowsketch, 26 PFR (plug flow reactor), 55,558 comparison with CSTR, complex reactions, 569 volume ratio to CSTR, 571 Phase diagrams nitrotoluene isomers, 544 salt solutions, 526 use of, example, 528 Phenol by the chlorbenzene process, 34 Phosgene synthesis, 594 Phthalic anhydride synthesis, 593 PID (proportional-integral-derivative) controllers, 41, 42 Pinenes Separation, chromatographic, 521 Pipe fittings resistances, 95, 98-100 Pipelines networks, 98, 100-102 non-Newtonian liquids, 106 optimum economic size, 102,103 Piping dimensions, 675 schedule number, 129 Piston pump. See Positive displacement Plate and frame filters, 319 performance, 331 sizes, commercial, 323 Plate exchangers, 189,195 Plates, fractionating. See Trays Pneumatic conveying, 71-76 dense phase transfer, 74 equipment, 72,73,75 flow rates, 72 operating conditions, 73 pilot plant, 74 power requirements, 72,74,77 oressure drop, 76.77 Pneumatic conveying dryers, 255 performance, 263,264 pilot plant size, 237 sizing example, 266,277 sketches 265 Podbielniak extractor. 478. 484. 487 Poiseuille equation, 94 Polyethylene reactor, 573, 574 Polytropic head, 156 Positive displacement pumps application range, 143 characteristics, 134, 143 discharge curves, 135, 141 efficiency, 135 steam consumption, 135 Powder blenders, 301,303 Power law behavior, non-Newtonian, 71 Power number, mixing, 290-292 limiting values, 293 Power requirement for pipelines, 98

Power supply, 9 from a hot gas stream, 12 generation with steam, 11 Pressure control. 42.44.51.52.59.60 Pressure drop cyclone separators, 617 gas-solid flow, 119-120 granular beds, 117 heat exchanger example, 193, 194 heat exchangers, 188 non-Newtonian flow, 106-109 wire mesh pads, 616 Pressure drop, piplines, 92 chart method, 96 two-phase flow, 116 typical values, 95 Pressure vessel code, ASME, 625 Prilling, 361.362 equipment size, 367 flowsketch, 366 operating, data. 367 products of, 367 size distribution, 362 Prism membrane separation process, 633 643 Propeller pump, 143 Proprietary equipment, 1 Propylene oxide synthesis, 34 Pseudoplasticity, 71, 103, 104 Psvchrometric chart. 232 application example, 234 wide temperature range, 233 Pulsed extractors, 478,482,487 amplitude of pulse, 484 frequency, 484 hole size, 484 interfacial tension effect, 484 packing size 484 pulsing modes, 481 Pumps, 131-144 application ranges, 143, 150 characteristic curves, 134, 137 characteristics. 144 control, 55,58,59 dimensionless groups, 131 efficiency, 144 efficiency, Moody's formula, 131, 132 glossary; 166 oarallel oneration. 133. 147 performance, 144,150 performance, dimensionless, 131. 132 performance, typical, 144 seals, 137,142 selection criteria, 140, 143, 144 series operation, 133, 147 q, distillation feed condition, 381 q-tine, McCabe-Thiele diagram, 381 Questionnaires, vendors, index, 727 Raoult's law, 371 Rates, chemical reactions, 549 basic equations, 554 constant pressure, 554 constant volume, 554 integrals of equations, 556 Langmuir-Hinshelwood mechanism, 554law of mass action, 554 plug flow reactor, 555

simultaneous reactions, 554 solid catalyzed, 554

stirred tanks, 555 temperature effects, 553,554 Rate equations, chemical basic, 554 integrals of, 556 plug flow reactor, 555 stirred tanks. 555 Rayleigh distillation, 378 multicomponent, 379 Raymond flash dryer, 265 performance, 263 Raymond mill, 345 RDC (rotatine disk contactor). 482,487 capacity, 487, 489 design example, 488 formulas, 489 Reactors. chemical. 567-582 classification. 568 ebbulating bed. 593 fermentacon, 654.659.660 fired heater, 574,575 fixed bed, 572 flame, 573 fluidized bed, 577,591,592 gas-liquid, 570,571 immiscible liquids, 570, 571 mixed. 570 moving bed, 574,588,589 pebble heater, 574,575,578 rotary kiln, 575,576, 590 stirred tanks. 567-569 stirred tanks, batch and continuous, 569 stirred tanks, typical proportions, 568 tubular flow, 569 Reactors, fementation, 654,659,660 Reboilers, 206-208 control. 47 design example, 209 guide to selection, 208 sketches. 175 Reciprocating pumps. See Positive displacement pumps Reciprocating compressors, 146, 148, 149 sizes of commerical equipment, 153, 154 Redler conveyors, 85 sizing calculation, 88 Reflux, distillation, 371 minimum, ratio, 382,387,397 optimum, ratio, 382,387 Reflux drums, 612 holdup, 612 sketch, typical, 613 Watkins method for sizing, 612 Reforming, platinum catalyst, 572 Refractories, properties, 223 Refrigerants, 226 characteristics, 15 data, 227 freons, 228 Refrigeration, 224 ammonia absorption, 226,227,229 cascades, 226 circuits, 225 compression, 224-227 thermodynamic diagrams, 225 Regenerators, 188 Relative volatility, 374 correlation example, 375 Residence time distribution, 556,558,559 commercial and pilot equipment, 559 CSTR battery, 558,559 dispersion model, 560

drvers. 245 Erlang distribution, 556 impulse response, 556 laminar flow 561 Peclet number, 558,559 PFR. 558.559 variance. 558 Residual enthalpy and entropy, 158, 159 Reverse osmosis. See Osmosis. Desalination ethanol/water separation, 642 Reynolds number, 92 non-Newtonian flow. 108.109 Rheopectic liquids, 104, 105 Ripple travs. 426 Roll presses, 358 commercial sizes, 364 product list, 363 Rotary compressors, 149 performance data, 154, 155 Rotary drvers, 247 design details, 256 performance, 257-259 scaleup, 256 sketches. 254.255 tray type, 251 Rotarv kiln reactors. 575, 590 Roughness factor, pipelines, 94 Rounds equation, friction factor, 94 RRS (Rosin-Rammler-Sperling) equation, 525,530-532 Rules of thumb compressors and vacuum pumps, xiii convevors for particulate solids, xiii cooling towers, xiii crystallization from solution, xiv disintegration, xiv distillation and gas absorption, xiv drivers and power recovery equipment, XV drying of solids, x evaporators. xv extraction. liauid-liouid, xv filtration, xvi, fluidization of particles with gases, xvi heat exchangers, xvi insulation xvi mixing and agitation, xvii particle size enlargement, xvii piping, xvii reactors, xvii refrigeration, xviii size separation of particles, xviii utilities, common specifications, xviii vessels (drums), xviii vessels (pressure), xviii vessels (storage), xix Safety factors, 6,7 equipment sizing, table, 7 Safety, plant, 7 checklist about chemical reactions, 8 checklist of startup and shutdown, 8,9 potential hazards, 8 Salicylic acid purification, 646 Salt cake furnace, 590 Sand cracking reactor, 593 Sand filters, 118,322 (simultaneous correction) method, SC 404, 408, 410 algorithm flowsketch, 411 SCFT concept, filtration, 317

754 INDEX

Scatchard-Hildebrand equation, 374 Schacham equation, friction factor, 94 Scheibel extractor, 483,487 Schildknecht column, 543,544,546 Screens, separating, 335, 337 capacity, 335, 340 flat. 335 reciprocating, 335 revolving, 335 rotary sifter. 335 sketches. 338 Screw conveyors, 76,79.80 sizing calculation, 80 sizing data, 79 types of screws, 80 Screw pumps, 134, 141 performance, 148 Sedimentation 305-307 behavior. equipment, 319 equipment sketches. 320.338 Segregated flow, 560 volume ratio to CSTR, 561 volume ratio to maximum mixedness. 560 Separation, solid-liquid. See also Filtration chief mechanical means, 305 315.320 clarifying, comparison of equipment, 307 experimental selection routine, 308 flocculants. 309 slurry pretreatment, 309 thickening, 315,320 Separators, gas-liquid cyclones, 616 wire mesh, 616 Separators, liquid-liquid, 612 dispersed phase criteria. 612-615 droplet sizes, 612 example, calculation, 614 Settling rate, 70 Newton's equation, 614 Stokes' equation, 612, 614 Shale oil reactor, 589 Shell and tube heat exchangers, 195 design. 199 design procedure, 202 internals, 201 sketches, 178 TEMA classification, 200 tube counts, 203 Sherwood-Lobo flooding correlation, 441 Siemens-Martin furnace-regenerator, 590 Sieve tray extractors, 483 capacity, 484,487 diameters, 483, 487 efficiency. 483.487 pulsed, 478,483,487 sizing example, 486 Sieve trays, 428 assembly in a tower, 428 comparison with other types, example, 431 malfunctions, 431 operating ranges, 431 phenomena, 430 specifications, 429 Sintering, 363 orocess sketch. 359 She enlargement, 351-369 prilling, 361,362

products. 354 product shapes, 362 Size enlargement equipment, 351 disk granulators, 352.355 extruders, 358 fluidized bed, 362,368,369 paddle blender, 360 pin mixer. 361 roll presses, 358,363 rotating drum, 353.357 spouted bed, 362 tumblers, 351, 355 Size reduction, 339-350 application, various materials, 343, 344 operating ranges of equipment, 343 power requirement, example, 342 size distribution of product, 337 surface energy, 339 work index. 342 work required, 339.342 Size reduction equipment, 341 attrition mills, 343, 348, 351 ball mills, 342, 351 buhrstone. 343.352 colloid, 343, 352 disk attritor, 343, 352 gyratory, 343.352 hammer mills, 341.346.350.351 jaw crushers, 341,346, 349 jet mills, 350,351, 352 mikro-pulverizer, 350 pebble mills, 343 rod mills, 342,346,347 roll crushers, 341,346,349 roller mills, 343,345,348,351,352 squirrel cage disintegrator, 343,352 tube mills, 343,346,351 tumbling mills, 342 Slurry pretreatment, 309,315 Slurry transport, 69-71 critical velocity, 69 pressure drop, 69-71 Soave equation of state, 373 Sodium carbonate/water diagram, Sodium sulfate/water diagram, 529 530 Solubility of solids, 523 data. 525 phase diagrams, 526 supersaturation, 525,527,528 Souders-Brown correlation, 432 Space velocity, 549-553 Specification forms, index, 681 Specific speed, pumps, 131, 133, 136 Spouted beds, 362 Spray dryers, 268-279 atomizers, 274.276 operating variables, 277 particles sizes, 276 performance, 275 pilot plant performance, 278 pilot unit, 237, 278 product density, 276 product number 278 residence time, 270 residence time distribution, 245 sizing example, 279 sketches, 274 SR (sum rates) method, 407,409 algorithm flowsheet, 408 Standards and codes, 2, 3 Steam heaters, 45,46 Steam jet ejectors, 143, 162

arrangements. 163 performance profiles. 166 steam consumption, 165 theory, 166 Steam supply characteristics. 15 generation, 9, 10 power generation, 11 Steam turbines advantages. 62 data sheet. 66 efficiency. 63 steam requirement, calculation, 65 theoretical steam rates, 64 Steam, water properties, 673,674 Stirred tank, crystallization model, 533-539 applicability of the model, 536 data analysis, example, 537 multiple tanks, 536 performance, example, 538 product size distribution, 539 Stirred tank design, 287,288 baffles. 287 draft tubes, 287 impellers, 288-290,293,294 sketch, basic, 288 Stirred tank impellers descriptions. 288.290 location, 288 size, 288 size, calculation example, 293, 294 sketches. 289 speed. 288 Stokes eauation. 70.612.614 Storage tanks, 619 API standard sizes. 622 buried, 621 granular solids, 624 horizontal, 619 large sizes, 623 pressure, 621 supports, 621,624 vertical, 621 Structured packings, 437 Sublimation, 638 equipment. 639 products, 646 salicylic acid purification, 646 Sulfur dioxide oxidation reactors, 572, 578-580 reaction equilibria, 580 temperature profiles, 580 Sulfur isotope separation, 650 Supercooling, maximum, 527,528 Supersaturation, 525,527,528 crystal growth rate and, 533 data, 527 thermodynamic analysis, 531 Suspension of particles, 295, 296 data, 298 example calculation, 299 Suspensions, non-Newtonian behavior, 71 Sweetland filter, 319,323-325 Symbols, flowsheet, 21-25 Synthetic fuel reactors, 585, 594, 595 Tabletting machines, 356 Manestv. 356.358 product shapes, 362 356,358 Sharples, US Pharmacopeia specifications, 357

INDEX 755

Tall oil distillation, 36 TEMA classification, heat exchangers, 200 Temperature control, 42,45-47 Temperature difference, 172 logarithmic mean, 172 multipass exchangers, 173, 175-177 Temperature profiles, heat exchangers, 173 Temperature profiles, reactors ammonia synthesis, 582, 584 cement kiln, 590 cracking of petroleum, 595 endo- and exothermic processes, 584 jacketed tubular reactor, 584 methanol synthesis, 580 phosgene synthesis, 594 reactor with internal heat exchange, 584 sulfur dioxide oxidation, 580 visbreaking, 595 Thermal conductivity insulating materials, 222 packed beds, 587,599-601 Thermal diffusion separation, 642 cell sketch, 649 hydrocarbon isomers, 644,649 isotopes, 651 performance, 649 sulfur isotopes, 650 Thermosiphon reboilers, 207 calculation example, 193 Theta method, heat exchangers, 179, 181, 182 chart. 181 example calculation, 182 Thickeners, 315,320 performance, 307,320 sketches, 320, 338 Thiele-Geddes method, 404, 407 Thiele modulus, 566 Thixotrooic liauids. 104-106 Tielines, liquid-liquid equilibria, 460, 461 Hand correlation, 463,465 Ishida correlation, 463, 465 Othmer and Tobias correlation, 463, 465 Trays, fractionating assembly of sieve trays, 428 bubblecap, 428,430-433 capacity, F-factor, 429 capacity, Jersey Critical, 432 Souders-Brown, 432 capacity, cartridge, 428 design data sheet, 429 dualflow, 426 efficiency, 439-456 Linde, 430 ripple, 426 sieve, 428,429 turbogrid, 426 types, 426 valve. 429.430.432 Trickle reactors, 576, 607 Tridiagonal matrix, 407 Trommels, 335 Tube count table, heat exchangers, 203 Tubular heat exchangers sketches, 174, 175

TEMA classification, 200 tube count table, 203 Turbine pumps, 134, 139, 140, 142, 143 Turner equation, leaching, 466 Turbogrid trays, 426 Two-phase fluid flow, 111, 113-117 correlations, 115 granular beds, 118 homogeneous model, 113 Lockhart-Martinelli method, 115, 116 patterns, 114 pressure drop, calculation example, 116 segregated flow model, 114 void fraction, 116 Tyler sieves, 336 Ultrafiltration, 631 applications, 633 membranes, 637-639 Underwood minimum reflux binary, 387 multicomponent, 397 Units, conversion of, 671, 672 UNIQUAC equation, 475 Upflow fixed beds, 609 Uranium recovery, 515 Utilities, typical characteristics, 15 Vacuum drum filters, 319 air flow rates, 328 applications, 332 cycle design, 328 flowsketch, 326 laboratory test data, 312 minimum cake thickness, 328 operation, calculation example, 312, 313 operation modes, 326 performance, 307,333 sizes. commercial. 327 Vacuum filter, horizontal, 321 performance, 307 sizes, commercial, 327 sketches. 325 Vacuum pumps, 162 air leakage, 164, 165 operating ranges, 145 steam jet ejectors, 163, 165, 166 Vacuum systems air leakage, 164, 165 other gas leakage, 165 Vallez filter, 319,323-325 Valve trays, 429-432 comparison with other types, example, 431 diameter sizing chart, 432 sketches of valves, 430 Valves, 129 control, 129-131 friction in. 99 van der Waals equation of state, 111, 113 Vapor-liquid equilibria, 371 binary data, 376 binary x-y diagrams, 375 in presence of solvents, 416,418-420 Raoult's law, 371 relative volatility, 374

VER chart, 372 Vapor-liquid equilibrium data acetone/methanol, 416 acetone/water, 416,423 butadiene, 420 butane/2-butene in solvents, 419 butanol/water, 388 chloroform/acetone/MIBK, 416 cyclohexane/benzene in solvents, 418 ethane/butanel/pentane, 378 ethanol/acetic acid, 385 ethanol/butanol. 375 ethanol/water, 387,416 heptane/methylcylohexane in solvents, 418 isoprene, 420 methylethylketone/water, 423 octane/toluene/phenol, 416 Velocities in pipelines, typical, 95 Vendors questionnaires, list, 727 VER (vaporization equilibrium ratio), 371 chart, 372 Vessels, process ASME code. 625 design example 628 heads, 624,625 heads, types, 627,628 mechanical design, 621 pressure, 623 shells, 624, 625 temperature, 623 tensile strength, 623, 626 thickness formulas, 625 Visbreaker operation, 33 Viscosity units, 91 Volatility, relative, 374, 375 Wang-Henke method, 404,408 algorithm flowsketch, 406 Water cooling, typical conditions, 15 properties, 673,674 vapor pressure, 231 Wet bulb temperature, 234 WHSV. 549 Wilson equation, activity coefficients, 374 Wilson-Lobb-Hottel equation, 214 application, 219 Winkler process 582 Wire mesh deentrainers, 615 calculation example, 620 disengaging space, 616 kev dimensions, 618 k-values, 615 pressure drop, 616 tvnical installations, 616 Work index, size reduction, 342 Wyssmont dryer, 237,251 Xylenes separation by crystallization, 544, 545 Zippered belt conveyor, 84 sizing calculation, 88 Zwietering equation, maximum

mixedness, 560



Chemical Process Equipment Selection and Design

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Chemical Process Equipment is a guide to the selection and design of a wide range of chemical process equipment. Emphasis is placed on specific information concerning the process design and performance of equipment. To this end, attention is given to examples of successful applications, and a generous number of line sketches showing the functioning of equipment is included with many graphs and tables giving their actual performance. For coherence, brief reviews of pertinent theory, including numerical examples to illustrate the more involved procedures, are provided in key chapters.

Chemical process equipment is of two kinds: custom-designed and built or proprietary "off the shelf." Unlike custom-designed equipment, proprietary equipment is often developed without the benefit of much theory and is fitted to job requirements also without the benefit of much theory. The process design of proprietary equipment, as considered in this book, establishes its required performance and is then a process of selection from the manufacturer's offerings. Since more than one kind of equipment often is suitable for particular applications and may be available from several manufacturers, comparisons of equipment and typical applications are cited liberally.

Professor Walas, drawing upon his many years of experience in industry and academia, provides a wealth of valuable shortcut methods, rules of thumb, and design by analogy applications. References to sources of more accurate design procedures are cited whenever they are available. To illustrate the data essential to process design, a substantial number of equipment ratings forms and manufacturers' questionnaires have been collected. Because decisions often must be based on economic grounds, a short chapter on costs of equipment rounds out the book.

Chemical Process Equipment offers comprehensive and thoroughly practical coverage of this field. Focusing on real-world considerations, it will be of particular value to practicing engineers in chemical process and petroleum refining industries and students in chemical engineering design courses.



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