Unit Operations of Particulate Solids

Theory and Practice



Enrique Ortega-Rivas



This page intentionally left blank



Unit Operations of Particulate Solids

Theory and Practice

Enrique Ortega-Rivas



This page intentionally left blank



CRC Press Taylor & Francis Group 6000 Broken Sound Parkway NW, Suite 300 Boca Raton, FL 33487-2742

© 2012 by Taylor & Francis Group, LLC CRC Press is an imprint of Taylor & Francis Group, an Informa business

No claim to original U.S. Government works

Printed in the United States of America on acid-free paper Version Date: 20110608

International Standard Book Number: 978-1-4398-4907-1 (Hardback)

This book contains information obtained from authentic and highly regarded sources. Reasonable efforts have been made to publish reliable data and information, but the author and publisher cannot assume responsibility for the validity of all materials or the consequences of their use. The authors and publishers have attempted to trace the copyright holders of all material reproduced in this publication and apologize to copyright holders if permission to publish in this form has not been obtained. If any copyright material has not been acknowledged please write and let us know so we may rectify in any future reprint.

Except as permitted under U.S. Copyright Law, no part of this book may be reprinted, reproduced, transmitted, or utilized in any form by any electronic, mechanical, or other means, now known or hereafter invented, including photocopying, microfilming, and recording, or in any information storage or retrieval system, without written permission from the publishers.

For permission to photocopy or use material electronically from this work, please access www.copyright.com (http://www.copyright.com/) or contact the Copyright Clearance Center, Inc. (CCC), 222 Rosewood Drive, Danvers, MA 01923, 978-750-8400. CCC is a not-for-profit organization that provides licenses and registration for a variety of users. For organizations that have been granted a photocopy license by the CCC, a separate system of payment has been arranged.

Trademark Notice: Product or corporate names may be trademarks or registered trademarks, and are used only for identification and explanation without intent to infringe.

Visit the Taylor & Francis Web site at http://www.taylorandfrancis.com

and the CRC Press Web site at http://www.crcpress.com

Contents

Part I		racterization of Particulate Systems Relation to Storage and Conveying		
1 Intr		n		
1.1				
1.2	Powde	er and Particle Technology	6	
1.3		olid State: Main Distinctive Properties		
	1.3.1	Primary Properties	8	
		1.3.1.1 Particle Size and Shape	9	
	1 0 0	1.3.1.2 Particle Density		
	1.3.2	Packing Properties		
		1.3.2.1 Bulk Density		
Pofe	roncoc	1.3.2.2 Other Packing Properties		
Kere	rences.			
2 Bull	k Solids	s: Properties and Characterization	31	
2.1	Introd	ductory Aspects	31	
2.2		ification of Powders		
	2.2.1	Jenike's Classification		
	2.2.2	Geldart's Classification		
2.3	Samp	ling	37	
2.4	Failur	e Properties	48	
	2.4.1	Introduction		
	2.4.2	Description of Failure Properties		
	2.4.3	Experimental Determinations		
	2.4.4	Other Important Properties	61	
2.5	Labor	ratory Exercise: Determination of Some		
		re Properties of Powders		
	2.5.1	Introduction		
	2.5.2	Instrument and Materials		
	2.5.3	Shear Tester Operation		
2 (2.5.4	Calculations and Report		
2.6		le Size Analysis		
	2.6.1	Introduction Definitions of Characteristic Linear Dimension		
	2.6.2			
	2.6.3	Types of Distributions	68	

		2.6.4	Measur	es of Central Tendency	69
		2.6.5		ation of Data	
		2.6.6		ory Techniques for Size Measurement	
			2.6.6.1	Microscopy	77
			2.6.6.2	Screening	
			2.6.6.3	Sedimentation	
			2.6.6.4	Centrifugation	82
			2.6.6.5	Light Diffraction	
			2.6.6.6	Particle Counters	
			2.6.6.7	Online Measurement Techniques	86
	2.7	Labora	atory Exe	ercise: Particle Size Analysis by	
				ods	89
		2.7.1	Introdu	ction	89
		2.7.2	Materia	lls and Instruments	90
		2.7.3	Accesso	ories	90
		2.7.4	Microso	copy Method	91
			2.7.4.1	Introduction	91
			2.7.4.2	Slide Preparation	
			2.7.4.3	Measurement of Particle Size	91
			2.7.4.4	Procedure	91
			2.7.4.5	Calculations and Report	
		2.7.5	Screeni	ng Method	92
			2.7.5.1	Introduction	
			2.7.5.2	Choice of Sieves	
			2.7.5.3	Procedure	
			2.7.5.4	Calculations and Report	
		2.7.6	Sedime	ntation Method	
			2.7.6.1	Introduction	
			2.7.6.2	Preparation of Sample	
			2.7.6.3	Procedure	
			2.7.6.4	Calculations and Report	
		2.7.7		nental Method	
			2.7.7.1	Introduction	
			2.7.7.2	Procedure	
				Calculations and Report	
	Refe	rences			97
2	Store	ago and	Convoy	ing of Bulk Solids	101
3	3.1			torage Alternatives	
	3.2			ige	
	3.3			gege	
	5.5	3.3.1	Conors	l Considerations	102
		3.3.2		of Silos	
		3.3.3		resses in Axisymmetrical Bins	
		3.3.4		resses in Other Types of Vessels	
		J.J. T	vvan St	icosco in Onier Types or vessels	112

Contents vii

	3.3.5	Natural	Discharge from Silos	112
		3.3.5.1	Flow Theories	
		3.3.5.2	Hopper Opening for Coarse Bulk Solids	
		3.3.5.3	Hopper Opening for Fine Bulk Solids	
	3.3.6	Assisted	l Discharge	123
		3.3.6.1	Types of Discharge	
		3.3.6.2	Passive Devices	
		3.3.6.3	Active Devices	
	3.3.7	Feeders	for Discharge Control	
		3.3.7.1	Feeders Description	127
		3.3.7.2	Charge and Power Calculations	128
		3.3.7.3	Discharge Control in Feeders	130
	3.3.8	Other D	esign Factors	130
3.4	Labor	atory Exe	rcise: Evaluation of Wall Loads in Silos	134
	3.4.1	Introdu	ction	134
	3.4.2	Equipm	ent and Materials	134
	3.4.3		ents and Apparatuses	
	3.4.4		ıre	
	3.4.5	Calcula	tions and Report	135
3.5	Bulk S	Solids Cor	nveying	135
	3.5.1		ction	
	3.5.2	Belt Cor	nveyors	136
	3.5.3	Chain C	Conveyors	143
		3.5.3.1	Scraper and Apron Conveyors	145
		3.5.3.2	Bucket Elevators	148
	3.5.4	Screw C	Conveyors	151
	3.5.5	Pneuma	itic Conveying	155
		3.5.5.1	Introduction: Types of Conveyors	
		3.5.5.2	Dense-Phase Systems	
		3.5.5.3	Dilute-Phase Systems	159
		3.5.5.4	Design and Selection of	
			Dilute-Phase Systems	163
3.6	Labor	atory Exe	rcise: Pneumatic Conveying	
			of Different Granules	171
	3.6.1	Introdu	ction	171
	3.6.2	Equipm	ent and Materials	171
	3.6.3	Procedu	ıre	171
	3.6.4	Calcula	tions and Report	172
Refer				
Part II	Bull	k Solids	s Processing	
4 Size	Reduct	ion		179

viii Contents

		4.1.1	Introductory Aspects	179
		4.1.2	Forces Involved in Size Reduction	
		4.1.3	Properties of Comminuted Materials	
	4.2	Energ	y Requirements in Comminution	182
		4.2.1	Rittniger's Law	
		4.2.2	Kick's Law	
		4.2.3	Bond's Law and Work Index	
	4.3	Size R	eduction Equipment	
		4.3.1	Classification	
		4.3.2	Characteristics	184
			4.3.2.1 Crushers	185
			4.3.2.2 Rollers Mills	188
			4.3.2.3 Hammer Mills	189
			4.3.2.4 Disc Attrition Mills	189
			4.3.2.5 Tumbling Mills	191
			4.3.2.6 Other Types of Mills	
		4.3.3	Operation of Equipment	
	4.4	Criter	ia for Selecting Size Reduction Processes	
		4.4.1	Characteristics of Raw Materials	
		4.4.2	Feeding and Discharge Control	201
		4.4.3	Moisture	202
		4.4.4	Heat Generation and Removal	202
		4.4.5	Summary of Criteria and Main Applications	203
	4.5	Labor	atory Exercise: Determination of	
		Reduc	ction Relations Using a Hammer Mill	204
		4.5.1	Introduction	204
		4.5.2	Equipment and Materials	204
		4.5.3	Instruments	205
		4.5.4	Screen Analysis	205
		4.5.5	Operation of the Hammer Mill	
		4.5.6	Calculations and Report	205
	Refer	ences		206
5			ement	
	5.1		luction: Agglomeration Processes	
	5.2		egation Fundamentals: Strength of Agglomerates	
	5.3		meration Methods	
			Tumbling Agglomeration	
		5.3.2	Pressure Agglomeration: Compaction	
		5.3.3	Equipment Operation Variables	
	5.4		ion Criteria for Agglomeration Methods	
		5.4.1	Raw Materials Characteristics	
		5.4.2	Product Properties	
		5.4.3	Technique Options	221
	5.5	Desig	n Aspects of Agglomeration Processes	222

	5.6	Appli	cations	224
	5.7		ratory Exercise: Comparing Methods for	
			ling Agglomeration of Powders	224
		5.7.1	Introduction	
		5.7.2	Equipment and Materials	
		5.7.3	Instruments	
		5.7.4	Measurement of Particle Size Distributions	
		5.7.5	Operation of the Drum Agglomerator	
		5.7.6	Operation of the Pan (Disc) Agglomerator	
		5.7.7	Friability Test	226
		5.7.8	Calculations and Report	
	Refe	ences .		
6	Mixi	ng		229
	6.1		duction	
	6.2	Blend	ing Mechanisms	230
	6.3		tical Approach of Mixing Processes	
		6.3.1	Sampling	
		6.3.2		
			Mixing Indexes and Mixing Rate	233
	6.4	Mixir	ng Equipment	
		6.4.1	Tumbling Mixers	
		6.4.2	Horizontal Trough Mixers	240
		6.4.3	Vertical Screw Mixers	241
	6.5	Desig	n and Selection Factors	241
	6.6	Labor	atory Exercise: Determination of Blending	
		Indice	es for Different Tumbling Mixers	244
		6.6.1	Introduction	244
		6.6.2	Equipment and Materials	244
		6.6.3	Instruments or Apparatuses	244
		6.6.4	Operation of the Mixers	245
		6.6.5	Calculations and Report	245
	Refe	rences .		246
7	Fluid		n	
	7.1	Theor	retical Fundamentals	
		7.1.1		
		7.1.2	Fluid Flow through Solids Beds	250
		7.1.3	Mechanism of Fluidization:	
			Aggregative and Particulate	
	7.2	Fluidi	ized Regimes	
		7.2.1	Pressure Drop-Velocity Relationship	254
		7.2.2	Incipient Fluidization and	
			Minimum Fluidizing Velocity	
		7.2.3	Heterogeneous Fluidization: Bubbling	261

x Contents

		7.2.4	Spouted Beds	264
	7.3	Appli	cations of Fluidization	
		7.3.1	Applications in the Petroleum and	
			Chemical Industries	268
		7.3.2	Fluidized-Bed Combustion	268
		7.3.3	Drying in Fluidized Beds	269
		7.3.4	Coating of Particles and Particulates	
	7.4	Labor	atory Exercise: Fluidized-Bed Coating of	
		Food 1	Particulates	276
		7.4.1	Introduction	
		7.4.2	Equipment and Materials	
		7.4.3	Instruments and Apparatuses	276
		7.4.4	Operation of the Fluidized Bed	276
		7.4.5	Evaluation of the Coating Thickness	277
		7.4.6	Friability Test	277
		7.4.7	Calculations and Report	
	Refer	ences	*	
Pa	rt III	Sep	paration Techniques for Particulate S	olids
8	Intro	ductor	y Aspects	283
	8.1	Differ	ent Mixtures Relevant in Industry	283
	8.2		fication of Separation Techniques	
	8.3	Specif	ic Techniques for Granular Materials	284
	Refer	ences		286
9	Solid	Mixtu	ıres	287
	9.1	Screen	ning	287
		9.1.1	Basic Principles	
		9.1.2	Design and Selection Criteria	
		9.1.3	Equipment Used	296
	9.2	Electr	omagnetic Separation	301
		9.2.1	Basic Principles	301
		9.2.2	Equipment and Applications	303
		9.2.3	Selection Criteria	306
	9.3	Electr	ostatic Separation	
		9.3.1	Basic Principles	307
		9.3.2	Equipment and Applications	
		9.3.3	Applications in Fine Particulate Systems	312
	9.4	Labor	atory Exercise: Efficiency of Separation on	
		Single	e-Stage Screening	
		9.4.1	Introduction	312
		9.4.2	Equipment and Materials	313

		9.4.3	Instruments or Apparatuses	313
		9.4.4	Screening Procedure	
		9.4.5	Calculations and Report	
	Refer	ences	*	316
10	Solid	l-Fluid	Systems	317
	10.1		uction: Simultaneous Flow of Fluids and Solids	
			Classification of Fluids	
			Dynamics of Particles Submerged in Fluids	
	10.2		tion Efficiency	
			Evaluation of Efficiency	
		10.2.2	Total Gravimetric Efficiency	325
			Partial Gravimetric Efficiency	
	10.0	10.2.4	Grade Efficiency and Cut Size	329
	10.3		Gas Separations	
			Introduction	
		10.3.2	Use of Cyclones	
			10.3.2.1 Description of the Process	
			10.3.2.2 Theoretical Aspects	
			10.3.2.4 Applications	
		10.3.3	Air Classifiers	
		10.5.5	10.3.3.1 Description of the Technique	
			10.3.3.2 Theoretical Aspects	
		10.3.3	Operation and Applications	
		10.3.4	Gas Filters	345
		10.0.1	10.3.4.1 Description of the Process	
			10.3.4.2 Operation Characteristics and Applications	
		10.3.5	Scrubbers	
			10.3.5.1 Process Description	
			10.3.5.2 Equipment and Applications	349
		10.3.6	Other Techniques	351
			10.3.6.1 Settling Chambers	351
			10.3.6.2 Electrostatic Separators	
	10.4	Solid-	Liquid Separation Techniques	
		10.4.1	Properties of Suspensions: Rheology and Flow	
			10.4.1.1 Laboratory Exercise:	
			Rheograms of Suspensions	359
		10.4.2	Pretreatment of Suspensions: Coagulation and	
			Flocculation	
		10.4.3	Selection of Specific Techniques	362
			10.4.3.1 Laboratory Exercise:	
			Settling Tests to Select a Proper Technique	
			10.4.3.2 Calculations and Report	366

xii Contents

10.4.4	Sedimer	ntation	367		
	10.4.4.1	Introduction	367		
	10.4.4.2	Free Settling	368		
		Hindered Settling			
		Sedimentation Rate Stages			
	10.4.4.5	Operating Principles: Design and			
		Selection Factors	370		
	10.4.4.6	Laboratory Exercise: Acceleration of			
		Settling Rates by Coagulants	377		
10.4.5		ıgation			
		Introduction: Centrifugal Techniques			
		Applications of Centrifugation			
		Centrifugal Clarification			
		Centrifugal Filtration			
		Fixed Wall Devices: Hydrocyclones			
		Centrifugation Equipment	388		
	10.4.5.7	Laboratory Exercise: Efficiency of			
		Hydrocyclone Separations			
10.4.6		n			
		Filtration Theory			
		Constant Pressure Filtration			
		Constant Rate Filtration			
		Filtration Media and Filter Aids			
		Filtration Equipment	403		
	10.4.6.6	Laboratory Exercise: Filtration Curves	100		
10.47	Od T	Using a Filter Press	408		
		echniques: Membrane Separations			
		ionless Scale-up of Equipment			
References			426		
Appendix 1: Inte	ernationa	ıl (SI) System of Units	429		
Appendix 2: Der	nsity and	Viscosity of Water at Different			
Ten	nperatur	es and 101.325 kPa	431		
		Viscosity of Air at Different			
Ten	nperatur	es and 101.325 kPa	433		
Appendix 4: Din	nensiona	l Analysis	435		
Appendix 5: Star	ndard Sc	reen Scales	437		
		for the Preparation of Laboratory Repo			
Appendix 7: Notation449					
Index			457		

Preface

The idea of writing this book was conceived many years ago when the author started teaching unit operations in the undergraduate program of chemical engineering at the Autonomous University of Chihuahua, even prior to attending the then Postgraduate School of Studies in Powder Technology at the University of Bradford. The importance, relevance and prevalence of the topic of particle technology remain up-to-date. Studies by the Rand Corporation in the 1980s identified substantial differences in the scale-up and start-up performance of plants processing powders versus those processing liquids or gases. Particularly, in 1985, the Rand Report surveyed 40 US plants and found out that start-up times were 200-300% of those predicted (compared to 20% for fluid-based plants) and only achieved approximately 50% of their designed throughput. By comparison, most fluid-based processes reached 90% of planned output over the same period. Some of the reasons described at the time of the report were related to an inadequate understanding of the behavior of particle systems, which is sensitive to process scale or process history in ways that would not be expected by engineers familiar with only liquid or gas systems.

The response to overcome these difficulties came by way of promoting research, programs of study, and other activities in the field of powder or particle technology worldwide. Substantial advancements have been achieved through all these initiatives but, apparently, many of the reported reasons for this unequal understanding of powder-based processes, as compared to fluid-based processes, remain up to today. This perception is implied by a current need of consultancy in the subject by different types of industries worldwide. In a recent seminar at the Autonomous University of Chihuahua, Richard Farnish, consulting engineer at the Wolfson Centre for Bulk Solids Handling Technology of the University of Greenwich, strongly supported this perception by sharing with the audience the many examples of his consulting activities in the subject for at least a decade.

An obvious outcome of the upsurge of academic activities related to particle technology that started in the 1980s but keep momentum to date is the production of scientific literature about this particular topic. Papers, book chapters, encyclopedia contributions, reports, reference books, and textbooks have been produced during these years. The contribution represented by this work intends to add to the literature on the subject by trying to provide a textbook aimed at training practicing engineers in bulk solids handling and processing who might be, apparently, on high demand for years to come. The idea of writing the book started realization of the need for complementary literature on the subject, as already mentioned. The great opportunity that the author had was time to learn from an enviable academic staff

xiv Preface

of premium quality and who had an undisputable pioneering role in establishing the discipline known as particle technology. This sparked in the author a vested interest and a great fascination for the subject. In presenting this work I took knowledge and inspiration directly from my PhD thesis supervisor, Lado Svarovsky, but had close contact with some of the other "big names" of the time: John Williams, Derek Geldart, Nayland Stanley-Wood, Arthur Hawkins, and a young Martin Rhodes, initiating his career focused on fluidization. Drawing an analogy, the senior academic staff of the University of Bradford at the time would resemble the pitching staff of the US baseball team, the Atlanta Braves of the 1990s, or the current tennis Spanish team competing for the Davis Cup.

The book deals with unit operations in chemical engineering involving handling and processing of particulate solids. In the literature, as well as in many higher-education study programs, unit operations have been classified as heat transfer and mass transfer unit operations with excellent textbooks written following such criteria. There are a number of unit operations, which have been left out of these mentioned classifications, related to mechanical operations and fluid flow phenomena, but mostly coinciding with the subjects studied by the engineering discipline of particle technology. Commonly, most traditional unit operations books cover in great detail those unit operations involving heat and mass transfer, leaving the operations dealing with particulate solids at a mere introductory level. Most of these books are normally written as textbooks. On the other hand, books classified in the disciplines of powder technology, particle technology, or bulk solids handling, cover in great detail all unit operations involving particulate solids and granular materials, but are generally written as reference books and not as textbooks.

This work is aimed at filling a gap in the topic of unit operations involving particulate solids as it is written as a textbook but may be considered a reference book also. It is presented in the chemical engineering unit operations fashion of many textbooks, but with the additional feature of including suggested laboratory experiments. It has been written for students, undergraduates and postgraduates, as well as for educators and practicing chemical engineers. Readers will find it useful because it represents a suitable textbook for a series of courses related to particle technology taught worldwide in universities and higher-education institutions.

The work is divided into three main themes: characterization of particulate systems and relation to storage and conveying, bulk solids processing, and separation techniques for particulate solids. In the first part of the book, primary and secondary properties of particles and particulate systems are reviewed and analyzed thoroughly, focusing on their characterization and the effects on selection and design of silos and conveyors. The main purpose is to provide the student with theoretical and practical tools to understand the behavior of powders and pulverized systems. The second part deals with the main industrial operations of dry solids processing in order to give

insight into the operation principles of the most important technologies to handle dry solids in bulk. The theoretical principles are coupled with laboratory exercises to provide information and skills to operate, optimize, and innovate particle processing technologies. The third segment of the book refers to applications of a very interesting subject, which sometimes has not been properly covered in higher education engineering programs: two-phase and multiphase flow. Many engineering processes deal with these types of systems but, since the topic may be considered part of fluid mechanics, rheology, or particle technology, it is often left out of educational programs. All the relevant systems in industrial processes combining two different components of the state of matter are described, and the technologies involved in separating phases by purely mechanical means, are studied in the final part of the book. The emphasis is, again, on providing balanced theoretical and practical components to learn and understand the operation of machinery and equipment to carry out relevant industrial operations such as centrifugation, filtration, and membrane separations.

This work was originally written in Spanish, but publishers in this language from Spain to Argentina and Mexico, did not find the project suitable or attractive. Some time passed, however, in the process of promoting the book with publishers in the Spanish language. The decision was thus made to translate the work into English and the response from publishers in English was prompt and positive. The whole project, from its origins in Spanish to its transformation into English has, therefore, taken time and involved extra work. Indeed, no man is an island, and many people at different times have contributed, voluntarily or involuntarily, to this project.

The author takes great pleasure in acknowledging the different participants in the project of writing this book. The main reason and inspiration for planning, developing, and concluding the work are, of course, the students of the last decades sitting at the course of Unit Operations I in the Chemical Engineering Program of the Autonomous University of Chihuahua. The first draft, from handwritten notes prepared to teach this undergraduate course, was created using first versions of the current word processors by Carmelita Gonzalez, then secretary of the postgraduate program in Food Technology of the Autonomous University of Chihuahua. Many students from both the undergraduate program in chemical engineering and the postgraduate program in food technology have prepared excellent laboratory reports at different times and some of them were used in preparing several of the laboratory exercises for the book. It would be impossible to mention all of them by name but their contribution is valued. Jocelyn Sagarnaga-Lopez and Hugo Omar Suarez-Martinez, MSc supervisees of the author and recent recipients of a State Award on Scientific Merit for their degree theses, wrote a chapter along with him on solid-liquid separations for a book on food processing and some aspects of the theoretical part of it were used in preparing the last chapter of this book. A useful hand in the task of translating the book and making sense of such translation was given by Israel Marquez, a student xvi Preface

in the chemical engineering program at the sister institution of the Chihuahua Institute of Technology.

Last, but not least, sincere and fondest appreciation is given to my wife Sylvia, my daughters Samantha and Christina, who suffer the inevitable reduction of time shared with them due to the extra investment of time for the book writing project.

Enrique Ortega-Rivas Chihuahua, Mexico

Author

Professor Enrique Ortega-Rivas holds an MSc in food process engineering from the University of Reading and a PhD in chemical engineering from the University of Bradford. He is currently a professor at the Autonomous University of Chihuahua, México, and has held a visiting scientist appointment at Food Science Australia and a visiting lectureship at Monash University, Australia. He was also a Fulbright scholar, acting as adjunct associate professor, at Washington State University. He has been awarded the status of national researcher, the maximum recognition that the Mexican government confers on academics based on their research achievements. He was shortlisted for the 2009 IChemE Innovation and Excellence Award in Food and Drink Processing and he is recipient of the 2010 Chihuahua Award in Technological Sciences. He has taught food process engineering in Chihuahua, Washington State and Monash, while other teaching topics have included unit operations involving particulate materials, as well as heat and mass transfer operations. His research interests include food engineering, particle technology, and solid-fluid separation techniques. He has focused his research efforts on the biological applications of hydrocyclones, in food powder processing, and in the employment of pulsed electric fields to pasteurize fluid foods.

Dr. Ortega-Rivas has served on many committees for international scientific events and performed peer reviews for national and international research funding organizations. He has also published numerous papers in international indexed journals, in addition to chapters in books and contributions to encyclopedias. He is the coauthor of the book *Food Powders: Physical Properties, Processing, and Functionality* published by Springer and editor of the book *Processing Effects on Safety and Quality of Foods* published by CRC Taylor & Francis. He is a member of the editorial boards of *Food and Bioprocess Technology: An International Journal, Food Engineering Reviews*, and *The Open Food Science Journal*. He reviews manuscripts for different international journals.

Part I

Characterization of Particulate Systems and Relation to Storage and Conveying

This page intentionally left blank



Introduction

1.1 Definitions of Unit Operations

The term unit operation has been used to describe a physical and/or mechanical procedure occurring parallel to a chemical reaction known as unit process, which happens in diverse materials processing industries. In order to understand this term properly, one should bear in mind that modernly structured industries were shaped during the great industrial revolution that started in England in the eighteenth century. With the invention of the steam engine, some industries developed and grew in complexity as other energy sources, such as oil and electricity, were incorporated (Valiente and Stivalet, 1980). In modern economics, there are four main components in economic activities of a nation or region:

- *Primary sector:* extraction of natural resources (agriculture, fishing, mining, etc.)
- Secondary sector: transformation of primary products (industry)
- Tertiary sector: services (commerce, banking, transportation, etc.)
- Quaternary sector: technological exploitation (research, design, and development)

The industrial sector needs to be further classified, but that would be a difficult task. A broad general categorization could be considered, however, as follows:

- Manufacturing industry (assembled goods, automotive industry, etc.)
- Construction industry (building construction, industrial construction, etc.)
- Materials processing industry

The materials processing industry receives diverse raw materials to be transformed directly from the primary sector, either for direct consumption as

finished products or for further transformation in some other types of industries. The materials processing industry may be divided into four categories:

- Chemical industry
- Pharmaceutical industry
- Food industry
- Metallurgy industry

In many industrial plants that process materials, a fundamental aspect of their operation is a chemical reaction known as unit process, such as oxygenation, hydrogenation, or polymerization. In order for a particular reaction to be carried out, a series of controlled steps to create optimal conditions are required. These steps or maneuvers are the physical and/or mechanical processes mentioned before (evaporation, distillation, pulverization, etc.) and are known as unit operations. The term unit operation arose from the need to standardize and systematize the teaching of chemical engineering as a discipline, due to the growth of the industry activity generated by the industrial revolution. A sort of tailor-made appropriate professional was required to operate chemical processing plants and promote a harmonic development of the materials processing industry. In 1887 George E. Davis proposed, in a famous series of 12 lectures at Manchester Technical School in England, the creation of a special career to cater for the growing chemical processing industry. Davis worked as an inspector for the Alkali Act of 1863, a very early piece of environmental legislation that required soda manufacturers to reduce emissions to the atmosphere. Davis also identified broad features common to all chemical factories, and he published his influential Handbook of Chemical Engineering (Davis, 1904) that roughly defined chemical engineering as a profession. His lectures were criticized for being common place know-how observed around operating practices used by British chemical industries at the time. Davis' ideas were, however, fundamental in initiating new thinking in the US chemical industry and sparking, eventually, the launching of chemical engineering study programs at several US universities. Chemical engineering courses, organized by Lewis M. Norton, were taught by 1888 at the Massachusetts Institute of Technology (MIT). In 1891, the Department of Chemistry at MIT granted seven Bachelor's degrees for Chemical Engineering, the first of their kind to be bestowed in the world. Almost at the same time, chemical engineering courses were offered at Pennsylvania State University in 1892, at Tulane University in 1894, and at Michigan University in 1898 (Valiente and Stivalet, 1980).

The pioneering courses of the chemical engineering degree programs consisted of deep knowledge of chemistry and physics complemented by courses of mechanical engineering and descriptive courses about equipment and important industry processes. As soon as the graduates began practicing, they realized about flaws and inconsistencies in their formation. The courses

Introduction 5

they received were descriptive but in practice they needed to make engineering. They needed to know how to design equipment, how to calculate the size of a new plant, and so on. This served to discuss the teaching of chemical engineering again in the schools. From these discussions, the concept of unit operation was originated, attributed to Arthur D. Little (Brown, 2005) who acted as president of the Inspecting Committee of the Department of Chemistry and Chemical Engineering at MIT. Such concepts may be defined textually as: "Any chemical process, at any scale, may be reduced to a coordinated series of what may be called unit operations, such as pulverize, dry, roast, crystallize, filter, evaporate, electrolyze, and so on. The number of these operations is not big and only a few of them is involved in a particular process" (Valiente and Stivalet, 1980, p. 35). Unit operations nowadays are diverse, since new types of industries have arisen, some consisting of very specific features. A classification of unit operations is not an easy task and some can be found in literature. The criteria used to categorize unit operations are varied and may include types of phases involved in a process, mass transfer or heat transfer possibility, governing force (e.g., physical or mechanical) in an operation, and so on.

In general terms, the book by Walker et al. (1937) could be recognized as the first formal text of chemical engineering and the original work of McCabe and Smith (1956) as the most widely recognized and used text for educational purposes. Also, *Perry's Chemical Engineers' Handbook* (Green and Perry, 2008) is considered as the classical reference chemical engineering. An inspection to all of these classical works would reveal that the most widely studied unit operations are those involving heat transfer such as evaporation, mass transfer, for example, extraction, and combined heat and mass transfer, for example, drying. This trend is reasonable since, as mentioned before, the first chemical engineers had to optimize processes of the main industries of the time (acids, alkalis, explosives, textiles, cellulose, paper, etc.) in which heat and mass transfer operations predominate. They had to evaluate the thermodynamic and kinetic constants required to be able to work with the immense number of fluids involved in those chemical processing industries.

As the first industrial processes grew more complex, the industry activity was multiplied and sophisticated and new and varied industries appeared. In such a changing reality, chemical engineers realized that, in practice, certain processes previous or subsequent to the unit process that were previously overlooked also had great importance in the overall efficiency of the operation. These previous and subsequent steps represent a series of physical operations that participate, mainly, in the storage and distribution of materials through a whole processing line. It was also observed that a series of operations not involving fluids, but granular solids instead, had been neglected for lacking the same response to heat or phase change, common in fluids in traditional operations (evaporation, condensation, drying, etc.). Furthermore, it was recognized that numerous granular solids and particulate materials intervene in a series of industrial processes, both as raw material

or as finished products. There was awareness that many of these processes, or part of them, involved bulk solids handling and processing of solids. The understanding of behavior of particulate materials and industrial powders, within chemical engineering, did not have the same degree of advancement showed by fluids. In such a way, a need emerged for adapting teaching practice and research activities in chemical engineering, as a way to contribute to the advancement of knowledge in powder handling and processing.

1.2 Powder and Particle Technology

The negligence in the study of the bulk solid state of matter was serious by the 1960s, and hence there was a critical need of growth based on research directions detected by an integrated interdisciplinary approach to problems in this subject. It has been argued that powder technology came out as an academic discipline in Germany during the 1960s (Tardos, 1995). Other European countries, such as the United Kingdom and the Netherlands followed suit and, like Germany, promoted work in academic groups of the recently created discipline. By the same time, Japan and Australia joined the trend, while in the United States, it was not until the 1980s that awareness about the lack of professional training in the area arose (Ennis et al., 1994). Directives of DuPont emphasized that 60% of the 3000 products their industries handled were in the form of particles and, therefore, their process engineers required additional and exhaustive training. Partly for this reason, the National Science Foundation assigned resources for the establishment of the discipline in the study plans of chemical engineering of some universities. Research efforts in powder technology initiated from the time the discipline was established. European and Asian professional association and societies recognized its importance since then. Some of these associations, such as the Institution of Chemical Engineers (IChemE) and the Society of Chemical Industry in the United Kingdom have included established research groups in the topic, and have organized meetings and conferences on a regular basis. In the United States it was only until 1992 when a division, known as the Particle Technology Forum, was formed within the American Institute of Chemical Engineers. In a more global context, diverse international associations organize conferences and congresses on powder technology with delegates from around the world attending to present the most advanced developments in the area. While in Europe, Japan, and Australia the discipline was known initially as powder technology, in the United States it was given the name of particle technology. Nowadays, this latter term is most commonly used.

Regardless of how it is denominated, powder or particle technology is a branch of engineering dealing with the systematic study of particulate materials in a broad sense, whether in dry form or suspended within some Introduction 7

fluid. For this reason, the interests of this discipline are numerous and comprise operations of characterization, storage, conveying, mixing, fluidization, classification, agglomeration, and so on, of powders and particulate systems. Since many biphasic and multiphasic fluids are handled in many materials processing industries, a series of separation techniques, such as filtration and centrifugation, are also relevant in the study of particle technology. Finely divided and pulverized solids possess certain characteristics that make them different from chunky solids and, so, they cannot be fully understood under the discipline of materials science. Also, although under certain conditions they may flow, they cannot be addressed either by classic rheology for their study and research. In such a way, this branch of engineering has been developed alongside other related branches and the discoveries brought by the study of particulate systems focusing on the particle technology principles has resulted in benefits for different industrial processes, including manufacturing of chemicals, pharmaceuticals, foodstuffs, ceramics, and so on.

The first book published in the powder technology field was possibly the one authored by Dallavelle (1943) in the early 1940s, which contained basically all of the topics related to particle technology previously mentioned. Orr (1966) published another text which widened the coverage of the topics in Dallavelle's book. Beddow (1981) wrote a third known text in particle technology, which has been extensively used for teaching purposes for many years. Some other books have been published in the general area of powder technology (Rhodes, 1990), as well as in related topics of particle size measurement (Allen, 1997), fluidization (Geldart, 1986), and separation techniques (Svarovsky, 2000). An excellent treatise on powders and bulk solids is the book by Schulze (2007), which includes properties, handling and flow in a single volume. As far as it could be surveyed, the focus of treating the topics of powder technology as unit operations has only been presented by Orr (1966).

1.3 The Solid State: Main Distinctive Properties

The properties of the solid state may be approached from different points of view. Properties of solids in large pieces differ from properties of particulate solids. The most important properties of pieces of solids include density, hardness, fragility, and tenacity (Brown, 2005). The density is defined as the mass by volume unit, while the hardness measures the resistance of the solids to be scratched. A solid hardness scale was proposed by Mohs (Brown, 2005) and can be listed as follows:

- 1. Talc
- 2. Gypsum
- 3. Calcite

- 4. Fluorite
- 5. Apatite
- 6. Orthoclase feldspar
- 7. Quartz
- 8. Topaz
- 9. Corundum
- 10. Diamond

In the above scale, each mineral scratches the previous. It is used to make reference to this chart to evaluate the hardness of some other known materials. For example, the hardness of ordinary glass is 5.8. The fragility is a property representing how easy a substance may be crumbled or broken by impact. It does not necessarily have a direct relation with hardness. Gypsum, for instance, is soft but not fragile. Finally, the tenacity is known as the property the metals and alloys present to resist collisions.

The molecular structure of solids determines some other features, such as shape of particles. Pieces of solids fracture following exfoliation planes determined by its inner molecular arrangement. For example, galena breaks into cubes, graphite into platelet shape, and magnetite into approximate rounded grains. It also depends on structure the resistance that certain material offers to slide onto another material, known as friction. The coefficient of friction is defined as the relation between the parallel force to the rubbing surface in the direction of the movement, and the perpendicular force to the rubbing surface and normal to the movement direction.

A material is considered a powder if it is composed of dry, discrete particles with a maximum dimension of less than 1000 µm (British Standards Institution, 1993), while a particle is defined by the *McGraw-Hill Dictionary of Scientific and Technical Terms* (2003, p. 1537) as "any relatively small subdivision of matter, ranging in diameter from a few angstroms to a few millimeters." These two definitions present a diffuse perspective upon which collection of individual entities may be identified as a powder or, simply, as a system of particles. In any case both groups of individual entities represent bulk materials of relevance in many industrial processes and efficient and reliable methods for characterizing them are needed in research, academia and industry. For convenience, properties of powder and particulate materials have been divided into primary properties (those inherent to the intimate composition of the material) and secondary properties (those relevant when considering the systems as assemblies of discrete particles whose internal surfaces interact with a gas, generally air).

1.3.1 Primary Properties

Particle characterization, that is, description of primary properties of powders in a particulate system, underlies all work in particle technology. Primary

particle properties such as particle shape and particle density, together with the primary properties of a fluid (viscosity and density), and also with the concentration and state of dispersion, govern the secondary properties such as settling velocity of particles, rehydration rate of powders, resistance of filter cakes, and so on. It could be argued that it is simpler, and more reliable, to measure the secondary properties directly without reference to the primary ones. Direct measurement of secondary properties can be done in practice, but the ultimate aim is to predict them from the primary ones, as when determining pipe resistance to flow from known relationships, feeding in data from primary properties of a given liquid (viscosity and density), as well as properties of a pipeline (roughness). As many relationships in powder technology are rather complex and often not yet available in many areas, particle properties are mainly used for qualitative assessment of the behavior of suspensions and powders, for example, as an equipment selection guide.

1.3.1.1 Particle Size and Shape

There are several single-particle characteristics that are very important to product properties (Davies, 1984). They include particle size, particle shape, surface, density, hardness, adsorption properties, and so on. From all these mentioned features, particle size is the most essential and important one. The term "size" of a powder or particulate material is very relative. It is often used to classify, categorize, or characterize a powder, but even the term powder is not clearly defined and the common convention considers that for a particulate material to be considered powder, its approximate median size (50% of the material is smaller than the median size and 50% is larger) should be less than 1 mm. It is also common practice to talk about "fine" and "coarse" powders; several attempts have been made at standardizing particle nomenclature in certain fields. For example, Table 1.1 shows the terms recommended by the British Pharmacopoeia referred to standard sieves apertures. Also, by convention, particle sizes may be expressed in different units depending on the size range involved. Coarse particles may be measured in centimeters or

TABLE 1.1Terms Recommended by the British Pharmacopoeia for Use with Powdered Materials

	BS Meshes		
Powder Type	All Passes	Not More Than 40% Passes	
Coarse	10	44	
Moderately coarse	22	60	
Moderately fine	44	85	
Fine	85	_	
Very fine	120	_	

millimeters, fine particles in terms of screen size, and very fine particles in micrometers or nanometers. However, due to recommendations of the International Organization for Standardization SI units have been adopted in many countries and, thus, particle size may be expressed in meters when doing engineering calculations, or in micrometers by virtue of the small range normally covered or when doing graphs.

The selection of a relevant characteristic particle size to start any sort of analysis or measurement often poses a problem. In practice, the particles forming a powder will rarely have a spherical shape. Many industrial powders are of mineral (metallic or nonmetallic) origin and have been derived from hard materials by any sort of size reduction process. In such a case, the comminuted particles resemble polyhedrons with nearly plane faces, 4-7 faces, and sharp edges and corners. The particles may be compact, with length, breadth, and thickness nearly equal, but, sometimes, they may be plate-like or needle-like. As particles get smaller, and by the influence of attrition due to handling, their edges may become smoother and, thus, they can be considered to be spherical. The term "diameter" is, therefore, often used to refer to the characteristic linear dimension. All these geometrical features of an important number of industrial powders, such as cement, clay, and chalk, are related to the intimate structure of their forming elements, whose arrangements are normally symmetrical with definite shapes such as cubes, octahedrons, and so on.

Considering the aspects mentioned above, expressing a single particle size is not simple when its shape is irregular. This case would be quite frequent in many applications, mostly when dealing with powders of truly organic origin. Irregular particles can be described by a number of sizes. There are three groups of definitions, as listed in Tables 1.2 through 1.4: equivalent sphere diameters, equivalent circle diameters, and statistical diameters, respectively. In the first group, the diameters of a sphere which would have the same property of the particle itself are found (e.g., the same volume, the same settling

TABLE 1.2A List of Definitions of "Equivalent Sphere Diameters"

Symbol	Name	Equivalent Property of a Sphere
$x_{\rm v}$	Volume diameter	Volume
$x_{\rm s}$	Surface diameter	Surface
$x_{\rm sv}$	Surface volume diameter	Surface-to-volume ratio
$x_{\rm d}$	Drag diameter	Resistance to motion in the same fluid at the same velocity
$x_{\rm f}$	Free-falling diameter	Free-falling speed in the same liquid at the same particle density
$x_{\rm st}$	Stokes' diameter	Free-falling speed if Stokes' law is used ($Re_p < 0.2$)
$x_{\rm A}$	Sieve diameter	Passing through the same square aperture

TABLE 1.3	
A List of Definitions of "Equivalent Circle Diameters"	

Symbol	Name	Equivalent Property of a Circle
$\overline{x_a}$	Projected area diameter	Projected area if the particle is resting in a stable position
x_{p}	Projected area diameter	Projected area if the particle is randomly orientated
$x_{\rm c}$	Perimeter diameter	Perimeter of the outline

velocity, etc.). In the second group, the diameters of a circle which would have the same property of the projected outline of the particle are considered (e.g., projected area or perimeter). The third group of sizes are obtained when a linear dimension is measured (usually by microscopy) parallel to a fixed direction. The most relative measurements of the diameters mentioned above would probably be the statistical diameters because they are practically determined by direct microscopy observation. Thus, for any given particle, Martin's and Feret's diameters could be radically different and, also, both different from a circle of equal perimeter or equal area (Figure 1.1). In practice, most of the equivalent diameters will be measured indirectly to a given number of particles taken from a representative sample and, therefore, it would be most practical to use a quick, less accurate measure on a large number of particles than a very accurate measure on very few particles. Also, it would be rather difficult to perceive the above-mentioned equivalence of the actual particles with an ideal sphericity. Furthermore, such equivalence would depend on the method employed to determine the size. For example, Figure 1.2 shows an approximate equivalence of an irregular particle depending on different equivalent properties of spheres.

Taking into account the concepts presented above, it is obvious that the measurement of particle size results dependent on the conventions involved in the particle size definition and also the physical principles employed in the determination process (Herdan, 1960). When different physical principles

TABLE 1.4A List of Definitions of "Statistical Diameters"

Symbol	Name	Dimension Measured
$\overline{x_{\mathrm{F}}}$	Feret's diameter	Distance between two tangents on opposite sides of the particle
$x_{\rm M}$	Martin's diameter	Length of the line which bisects the image of the particle
$x_{\rm SH}$	Shear diameter	Particle width obtained with an image shearing eyepiece
$x_{\rm CH}$	Maximum cord diameter	Maximum length of a line limited by the contour of the particle

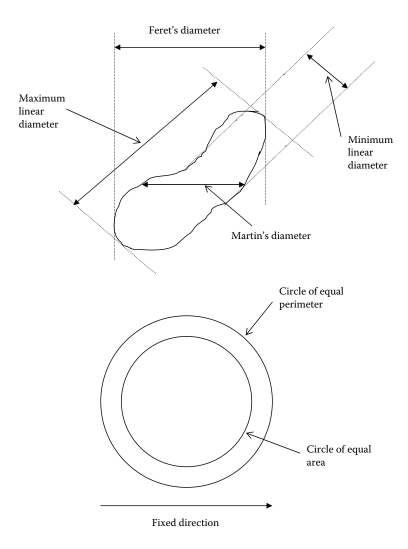


FIGURE 1.1
Methods used to measure the diameter of nonspherical particles.

are used in the particle size determination, it can hardly be assumed that they should give identical results. For this reason it is recommended to select a characteristic particle size to be measured accordingly to the property or the process which is under study. Thus, for example, in pneumatic conveying or gas cleaning, it is more relevant to use choose to determine the Stokes' diameter, as it represents the diameter of a sphere of the same density of the particle itself, which would fall in the gas at the same velocity as the real particle. In flow through packed or fluidized beds, on the other hand, it is the surface-to-volume diameter, that is, the diameter of a sphere having the same

Introduction 13

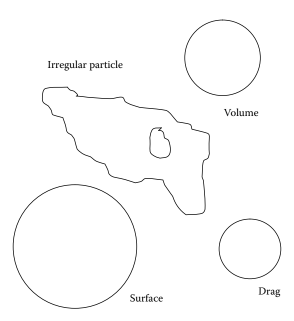


FIGURE 1.2 Equivalent spheres.

surface-to-volume ratio as the particle, which is more relevant to the aerodynamic process.

General definitions of particle shapes are listed in Table 1.5. It is obvious that such simple definitions are not enough to do the comparison of particle size measured by different methods or to incorporate it as parameters into equations where particle shapes are not the same (Herdan, 1960; Allen, 1997). Shape, in its broadest meaning, is very important in particle behavior and just looking at the particle shapes, with no attempts at quantification, can be beneficial. Shape can be used as a filter before size classification is performed. For example, as shown in Figure 1.3, all rough outlines could be eliminated by using the ratio: perimeter/convex perimeter, or all particles with an extreme elongation ratio.

The earliest methods of describing the shape of particle outlines used length L, breadth B, and thickness T in expressions such as the elongation ratio (L/B) and the flakiness ratio (B/T). The drawback with simple, one number shape measurements is the possibility of ambiguity; the same single number may be obtained from more than one shape. Nevertheless, a measurement of this type which has been successfully employed for many years is the so-called sphericity, $\Phi_{\rm s}$, defined by the relation

$$\Phi_{\rm s} = \frac{6V_{\rm p}}{x_{\rm p}S_{\rm p}} \tag{1.1}$$

General Delititions of Farticle Shape			
Shape Name	Shape Description		
Acicular	Needle shape		
Angular	Roughly polyhedral shape		
Crystalline	Freely developed geometric shape in a fluid medium		
Dentritic	Branched crystalline shape		
Fibrous	Regularly or irregular thread-like		
Flaky	Plate-like		
Granular	Approximately equidimensional irregular shape		
Irregular	Lacking any symmetry		
Modular	Rounded irregular shape		
Spherical	Global shape		

TABLE 1.5General Definitions of Particle Shape

where x_p is the equivalent diameter of particle, s_p is the surface area of one particle, and V_p is the volume of one particle. For spherical particles Φ_s equals unity, while for many crushed materials its value lies between 0.6 and 0.7. Since direct measurement of particle volume and surface is not possible, to evaluate such variables, a specific equivalent diameter should be used to perform the task indirectly. For example, when using the mean projected diameter x_a , as defined in Table 1.3, volume and surface of particles may be calculated using

$$V_{\rm p} = \alpha_{\rm v} x_{\rm p}^3 \tag{1.2}$$

and

$$s_{p} = \alpha_{s} x_{p}^{2} \tag{1.3}$$

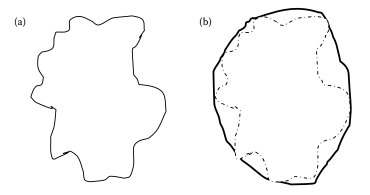


FIGURE 1.3
Relation between (a) perimeter and (b) convex perimeter of a particle.

Introduction 15

where α_v and α_s are the volume and surface factors respectively, and their numerical values are all dependent on the particle shape and the precise definition of the diameter (Parfitt and Sing, 1976). The projected diameter x_P is usually transferred into the volume diameter x_v of a sphere particle, as defined in Table 1.2, which is used as a comparison standard for the irregular particle size description, thus the sphere with the equivalent diameter has the same volume as the particle. The relationship between the projected and the equivalent diameters in terms of volume is expressed as follows:

$$x_{\rm v} = x_{\rm p} \left[\frac{6\alpha_{\rm v}}{\pi} \right]^{1/3} \tag{1.4}$$

where x_v is the equivalent diameter of the sphere of the same volume as the particle. When the mean particle surface area is known, the relationship between those two diameters is

$$x_{\rm v} = x_{\rm p} \left[\frac{\alpha_{\rm s}}{\pi} \right]^{1/2} \tag{1.5}$$

where all the variables have been previously defined.

Unambiguous shape representation involves collection and manipulation of a great deal of data. In view of this fact, consideration should be given to mechanical shape sorting before shape analysis. If shape is believed to be the cause of a particular problem or of powder behavior, then the use of sized and shaped-sorted material may provide the confirmation sought (Riley, 1968/1969; Shinohava, 1979). In some cases, however, this alternative route is not possible and, particularly in investigative work, detailed measure is necessary. Sebestyen (1959) suggested characterization of silhouettes by polar coordinates of their peripheries with the center of gravity of the figure as origin, as shown in Figure 1.4. When the R and θ readings are plotted, it is possible to represent the trace by a truncated harmonic series (Hatton, 1978). The value of the radius vector R as it is rotated about the origin is expressed as a function of the angle of rotation θ , in the truncated harmonic series of the form

$$R(\theta) = A_0 + \sum_{n=1}^{M} A_n \cos(n\theta - \varphi_n)$$
 (1.6)

where φ_n is the phase angle of the *n*th harmonic and $A_n = [(B_n)^2 + (C_n)^2]^{1/2}$. Each term of the harmonic series represents a particular shape and the silhouette is represented by different amplitudes and phases of these individual shapes (Figure 1.5). Clearly the system is not ideal because fine detail

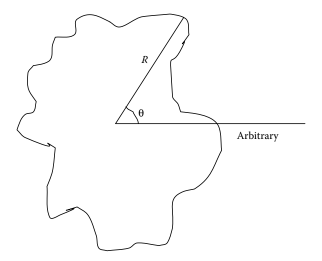


FIGURE 1.4Representation of polar coordinates in an irregular silhouette.

and protuberances produce difficulties of representation unless an inordinate large number of terms are used. Problems can also arise with silhouettes for reentrant features since R must remain single valued for all values of θ . The shapes can be reconstructed only by the use of both the amplitudes and the phases, but there are several ways of using the information in the representational truncated harmonic series. Attempts have been made at developing a series of combinations of harmonics to condense the information in the series (Luerkens et al., 1982), as well as using amplitudes of particular harmonics (Ehrlich and Weinberg, 1970; Holt, 1981).

An alternative approach to overcome shape evaluation difficulties with reentrant features is to represent any closed curve as a function of arc length by the accumulated change in direction of the curve (Gotoh and Finney, 1975). As represented in Figure 1.6, the outline is described essentially by taking a tangent round the shape and noting the shape in the angle. Reentrant

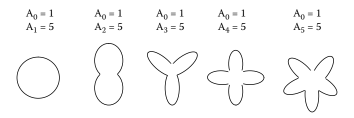


FIGURE 1.5 Shape of silhouettes at different amplitudes.

Introduction 17

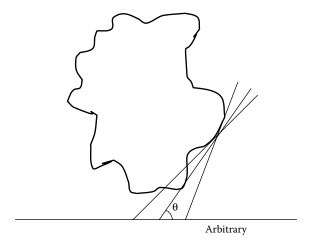


FIGURE 1.6 Description of a silhouette as a closed curve by its accumulated change in direction.

features are automatically dealt with and the outcome is again a harmonic series, but one which converges very slowly. The regenerate produced with a usefully small number of harmonic terms is not a closed figure, but this drawback has been overcome by a computational technique (Jones, 1983).

Detail of the surface of a particle, the roughness, can be observed by stereo microscopy, by scanning electron microscopy, and by confocal scanning optical microscopy, but a mathematical description depends on obtaining a satisfactory silhouette of the surface. This task is not easy, but considering that an image is available, the most promising approach would appear to be the one based on boundary-line analysis by fractal mathematics (Mandelbrot, 1977, 1982), which is concerned directly with the properties of the outline. In powder technology, the use of fractals was pioneered by Kaye (1981) among others, but the use of fractal mathematics is now appearing in many fields as well. When measuring lengths of irregular contours by stepping along them with a pair of dividers, as the step length s decreases, the estimated length s0 keeps on increasing without limit. In such a case, s1 is related to s2 by

$$L_{s} = K(s)^{1-D} (1.7)$$

or

$$\log L_{\rm s} = \log K + (1 - D)\log(s) \tag{1.8}$$

where *D* is known as the fractal dimension, being different for jagged and for smoothly rounded contours. The principle is, in general, applicable to any system where there is a series of estimates which tend toward infinity as the

resolution of the estimate improves. The emphasis is changed from the magnitude to the rate at which it is increasing toward infinity.

1.3.1.2 Particle Density

The density of a particle is defined as its total mass divided by its total volume. It is considered quite relevant for determining other particle properties such as bulk powder structure and particle size, and so it requires careful definition (Okuyama and Kousaka, 1991). Depending on how the total volume is measured, different definitions of particle density can be given the true particle density, the apparent particle density, and the effective (or aerodynamic) particle density. Since particles usually contain cracks, flaws, hollows, and closed pores, it follows that all these definitions may be clearly different. The true particle density represents the mass of the particle divided by its volume excluding open and closed pores, and is the density of the solid material of which the particle is made. For pure chemical substances, organic or inorganic, this is the density quoted in reference books of physical/chemical data. Since most inorganic materials consist of rigid particles, while most organic substances are normally soft, porous particles, true density of many biological powdered materials would be considerably low than those of mineral and metallic powders. Typical non metallic minerals would have true particle densities well over 2000 kg/m³, while some metallic powders can present true densities of the order of 7000 kg/m³. The apparent particle density is defined as the mass of a particle divided by its volume excluding only the open pores, and is measured by gas or liquid displacement methods such as liquid or air pycnometry. The effective particle density is referred as the mass of a particle divided by its volume including both open and closed pores. In this case, the volume is within an aerodynamic envelope as "seen" by a gas flowing past the particle and, as such, this density is of primary importance in applications involving flow round particles like in fluidization, sedimentation, or flow through packed beds.

Any of the three particle densities defined above should not be confused with bulk density of materials, which includes the voids between the particles in the volume measured. The different values of particle density can be also expressed in a dimensionless form, as relative density, or specific gravity, which is simply the ratio of the density of the particle to the density of water. It is easy to determine the mass of particles accurately but difficult to evaluate their volume because they have irregular shapes and voids between them. The apparent particle density, or if the particles have no closed pores also the true density, can be measured by fluid displacement methods, that is, pycnometry, which are in common use in industry. The displacement can be carried out using either a liquid or a gas, with the gas employed normally being air. Thus, the two known techniques to determine true or apparent density, when applicable, are liquid pycnometry and air pycnometry.

Liquid pycnometry can be used to determine particle density or fine and coarse materials depending on the volume of pycnometer bottle used. For fine powders, a pycnometer bottle of 50 mL volume is normally employed, while coarse materials may require larger calibrated containers. Figure 1.7 shows a schematic diagram of the sequence of events involved in measuring particle density using a liquid pycnometer. The particle density $\rho_{\rm s}$ is clearly the net weight of dry powder divided by the net volume of the powder, calculated from the volume of the bottle subtracting the volume of the added liquid, that is,

$$\rho_{\rm s} = \frac{(m_{\rm s} - m_{\rm 0})\rho}{(m_{\rm l} - m_{\rm 0}) - (m_{\rm sl} - m_{\rm s})}$$
(1.9)

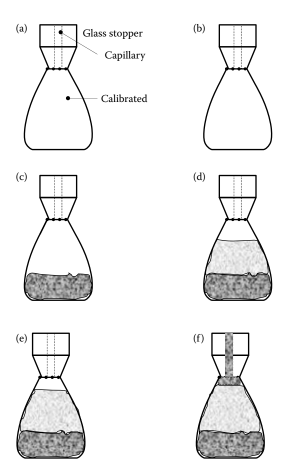


FIGURE 1.7 Descriptive diagram of density determination by liquid pycnometry: (a) description of pycnometer, (b) weighing, (c) filling to about 1/2 with powder, (d) adding liquid to almost full, (e) eliminating bubbles, and (f) topping and final weighing.

where m_s is the weight of the bottle filled with the powder, m_0 is the weight of the empty bottle, ρ is the density of the liquid, m_l is the weight of the bottle filled with the liquid, and m_{sl} is the weight of the bottle filled with both the solid and the liquid. Air bubbles adhering to particles and/or liquid absorbed by the particles can cause errors in density measurement. Therefore, a slowly particle absorbed liquid with low surface tension should be selected. Sometimes, when heating or boiling procedure is needed to do the gas evacuation, the liquid that has high boiling point and does not dissolve the particle should be used (Okuyama and Kousaka, 1991).

When the density of larger irregular shaped solid objects, such as compressed or aggregated bulk powders is needed, a method available to evaluate fruit or vegetable volumes might be used. A schematic diagram of a top-loading platform scale for volume and density measurement is shown in Figure 1.8. A beaker big enough to host the solid is partially filled with some kind of liquid which will not dissolve the solid. The weight of the beaker with the liquid in it is recorded and the solid object is completely immersed and suspended at the same time, using a string, so that it does not touch either the sides or bottom of the beaker. The total weight of this arrangement is recorded again, and the volume of the solid $V_{\rm s}$ can be calculated (Ma et al., 1997) by

$$V_{\rm s} = \frac{m_{\rm LCS} - m_{\rm LC}}{\rho} \tag{1.10}$$

where m_{LCS} is the weight of the container with liquid and submerged solid, m_{LC} is the weight of the container partially filled with liquid, and ρ is the density of the liquid.

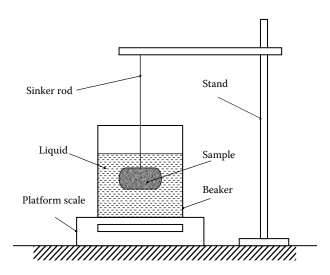


FIGURE 1.8Top-loading platform scale for density determination of irregular shaped objects.

Air pycnometry can be performed in an instrument which usually consists of two cylinders and two pistons, as shown in Figure 1.9. One is a reference cylinder, which is always empty, and the other has a facility for inserting a cup with the sample powder. With no sample present, the volume in each cylinder is the same so that, if the connecting valve is closed and one of the pistons is moved, the change must be duplicated by an identical stroke in the other as to maintain the same pressure on each side of the differential pressure indicator. Referring to Figure 1.9, if a sample is introduced in cylinder B, and the piston in cylinder A is advanced all the way to the stop, to equalize the pressures, the measuring piston will have to be moved by a smaller distance because of the extra volume occupied by the sample. The difference in the distance covered by the two pistons, which is proportional to the sample volume, can be calibrated to read directly in cubic centimeters, usually

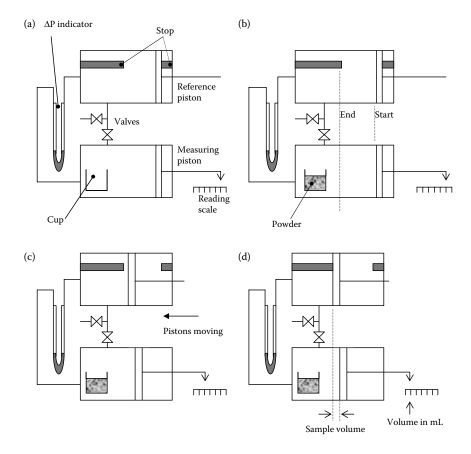


FIGURE 1.9Descriptive diagram of density determination by air pycnometry: (a) description of instrument, (b) filling of cup, (c) pistons displacement, and (d) reading.

with a digital counter. The method will measure the true particle density if the particles have no closed pores or the apparent particle density if there are any closed pores, because the volume measured normally excludes any open pores. If, however, the open pores are filled either by wax impregnation or by adding water, the method will also measure the envelope volume. By difference between the two volumes measured the open-pore volume will be obtained, and can be used as a measure of porosity. Materials that are not surface-active and incompressible, can be measured using room air and following the standard procedure that exposes the sample to air pressures from 1 to 2 bar. Compressible materials, however, would need to be tested at lower pressures (1/2 to 1 bar), while surface-reactive materials may be tested using helium or another inert gas, because they tend to absorb some or all of the air constituents. Considering all the aspects involved in the operation of air pycnometers, densities of most inorganic materials can be reliable determined using the standard procedure with normal air because their particles are normally hard and rigid.

The effective particle density is based on the average density within an aerodynamic envelope around it, including, therefore, any open or closed pores in the considered volume. One obvious way to measure the volume of the open pores is using a mercury porosimeter, but this is only suitable for coarse solids and the necessary equipment is very expensive. Alternative choices include the bed voidage method, the bed pressure drop method, and the sand displacement method. The bed voidage method (Abrahamsen and Geldart, 1980) is based on the observation that the minimum packed voidage is virtually the same for particles of similar size and particle shape. The procedure consists on pouring a small amount (0.2–0.25 kg) of a control powder of known particle effective density into a measuring cylinder and tapping it to its minimum volume, corresponding to the maximum bulk density. This procedure is repeated with the unknown powder, or several control powders in an ideal context. If the porosity after tapping is assumed to be the same for the control and unknown powders, then their effective densities are in the same ratio as their bulk densities. The bed pressure method is based on making measurements of bed pressure drop as a function of gas velocity at two voidages, when gas is passed through the bed of powder in the laminar flow regime. The bed is first fluidized to get it well mixed and then gently settled to its maximum voidage. Pressure drop is measured for at least four velocities; the bed is then tapped in order to reach as low voidage as possible and more measurements are made. Calling the set of measurements conditions 1 and 2 respectively, s the gradient of pressure drop, ρ_b the bulk density, and ρ_b the particle density, it can be shown (using the well-known Carman–Kozeny equation) that a basic relationship can be derived as follows:

$$\frac{S_1}{S_2} = \left(\frac{\rho_{b1}}{\rho_{b2}}\right) \left[\frac{(\rho_p - \rho_{b2})}{(\rho_p - \rho_{b1})}\right]^3$$
(1.11)

Introduction 23

where ρ_p is the only unknown variable and can be readily found by trial and error. The sand displacement method uses fine sand into which a known amount of coarse particles of the sample are mixed. The density of the sample is determined from the difference of the bulk density of the sand alone and that of the mixture. This method is sometimes used for density determinations of coarse bone particles, for example, and it gives lower density than that of the solid bone as measured by pycnometry. The use of effective particle density is largely restricted to fluidization and pneumatic conveying applications.

1.3.2 Packing Properties

As already stated, a powder is a complex form of solid material and it is made up of a very large number of individuals, each different from its neighbor. Individual, or inherent, properties have been already discussed in the previous section under the common term of primary properties. However, every time a particular powder sample is poured into a receptacle, the individual particles are located in different places from before and the structure of the powder is different. It is clear that it is not possible to predict quantitatively how a powder will behave from a knowledge of the measured properties of individual particles, and so direct measurement of bulk properties is, therefore, necessary. Because each repeated measurement on a sample will be upon a rearrangement of its population, there will be an inevitable scatter of readings. This difficulty has the consequence that the powder should be handled in as identical a manner as reasonably possible each time a measurement procedure is performed. Moreover, it would seem logical that there would be a development of standardized testing and characterization methods, but this has not happened.

1.3.2.1 Bulk Density

The measurement of the bulk density of powders in no exception to the general situation outlined above, but it is so fundamental to their storage, processing, and distribution that it does merit particular consideration. The bulk density of a powder is its mass divided by the bulk volume it occupies. The volume includes the spaces between particles and the envelope volume of the particles themselves. The spaces between particles are denoted as porosity or voidage, and can be defined as the volume of the voids within the bulk volume divided by the total bulk volume. Bulk density and porosity are related by

$$\rho_{\rm b} = \rho_{\rm s}(1 - \varepsilon) + \rho_{\rm g}\varepsilon \tag{1.12}$$

where ρ_b is the powder bulk density, ρ_s is the particle density, ϵ is the porosity, and ρ_g is the gas density.

The value of the bulk density depends on the state of the powder, particularly on its state of compaction. The bulk is a mixture of air (or other gas) and the solid particles, so that the bulk density can be anywhere between the density of the two phases involved. Thus, in the order of increasing bulk density, the following different definitions of this type of density that have been identified over the years are aerated bulk density, poured bulk density, tap density, and compacted bulk density. Each of these depends on the treatment to which the sample was subjected and, although there is a move toward standard procedures, these are far from universally adopted. There is still some confusion in the open literature in the sense of how these terms are interpreted. Some people consider the poured bulk density as loose bulk density, while others refer to it as apparent density. The actual meaning of aerated density can also be considered quite a confusing term. Strictly speaking, it should mean that the particles are separated from each other by a film of air and that they are, on the whole, not in direct contact with each other. Some authors, however, interpret the term as it would mean the bulk density after the powder has been aerated. Such an interpretation yields, in fact, the most loosely packed bulk density when, for cohesive materials, the strong inter-particle forces prevent the particles from rolling over each other. Considering this second interpretation, aerated and bulk densities could both be simply regarded as loose bulk density, and this approach is implied in many investigations when dealing with cohesive powders. For many fine and ultrafine powders, which are more likely cohesive in behavior, the terms more commonly used to express bulk density are loose bulk density, as poured and tapped bulk density, after vibration. Another way to express bulk density is in the form of a fraction of its particles' solid density, which is sometimes referred as the "theoretical density." This expression, as well as the use of porosity instead of density, enables and facilitates the unified treatment and meaningful comparisons of powders having considerably different particle densities.

The aerated bulk density is, in practical terms, the density when the powder is in its most loosely packed form. Such a form can be achieved by dropping a well-dispersed "cloud" of individual particles down into a measuring vessel. Alternatively, gas fluidization can sometimes be used with the gas turned slowly off and the particles allowed to settle quietly into place. The structure within the vessel is held by the cohesive forces between the particles, and can be extremely fragile. Leveling off the surface of the powder at the top of the vessel is difficult to achieve without causing particle movement leading to error, as some structure collapses. Determination of aerated bulk density can be performed using an apparatus such as the one illustrated in Figure 1.10. As shown, an assembly of a screen cover, a screen, a spacer ring, and a chute is attached to a mains-operated vibrator of variable amplitude. A stationary chute is aligned with the center of a preweighed 100 mL cup. The powder is poured through a vibrating sieve and allowed to fall a fixed height of 25 cm approximately through the stationary chute into the cylindrical cup.

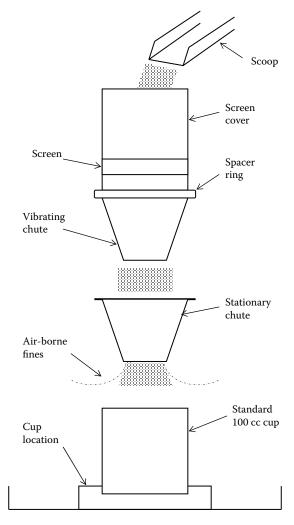


FIGURE 1.10Determination of aerated bulk density.

The amplitude of the vibration is set so that the powder will fill the cup in 20–30 s. The excess powder is skimmed from the top of the cup using the sharp edge of a knife or ruler, without disturbing or compacting, the loosely settled powder.

Poured density is widely used, but the measurement is often performed in a manner found suitable for the requirements of the individual company or industry. In some cases the volume occupied by a particular mass of powder is measured, but the elimination of operation judgment, and thus possible error, in any measurement is advisable. To achieve this, the use of a standard

volume and the measurement of the mass of powder to fill it are needed. Certain precautions which should be taken are clear, for example, the measuring vessel should be thick rather than thin, the powder should always be poured from the same height, and the possibility of bias in the filling should be made as small as possible. Although measuring of poured bulk density is far form standardized, many industries use a sawn-off funnel with a trap door or stop, to pour the powder through into the measuring container. Some decide on a standard volume, usually a measuring cylinder, to be used for the volume measurement and that varies from 50 to 1000 mL depending on the product and its size; 1 L is now accepted by many as preferred. It is also better to use a density cylinder with a 2:1 length-to-diameter ratio than the much slimmer measuring cylinder. Few people standardized the height of fall as well, realizing that the height will affect the compaction state of the powder. Finally, it is also the practice in some industries to allow the powder, after pouring it into the container, to settle to its stable density for about 10 min before scraping off the top.

The tap bulk density, as is implied by its name, is the bulk density of a powder which has been settled into a closer packing than existed in the poured state by tapping, jolting, or vibrating the measuring vessel. As with poured bulk density, the volume of a particular mass of powder may be observed, but it is generally better to measure the mass of powder in a fixed volume. Although many people in industry measure the tap density by tapping the sample manually, it is best to use a mechanical tapping device so that the conditions of sample preparation are more reproducible. An instrument useful to achieve such reproducibility is the Hosokawa powder characteristic tester, which has a standard cup (100 mL) and a cam-operated tapping device which moves the cup upward and drops it periodically (once in every 1.2 s). A cup extension piece has to be fitted and powder added during the sample preparation so that at no time the powder packs below the rim of the cup. After the tapping, excess powder is scraped from the rim of the cup and the bulk density is determined by weighing the cup.

The compacted bulk density would refer to the density of a powder compacted over and above what can be attained by tapping. There are no standard procedures for determining the compacted bulk density, other than the rather specialized soil compaction tests and density determinations, quoted in a British Standard for soils (British Standards Institution, 1975).

1.3.2.2 Other Packing Properties

Compressibility can be defined as the compaction of a powder by tapping or mechanical compression. These processes can occur either unintentionally as a result of handling or transporting, or intentionally as when producing tablets or agglomerates. In the materials processing industry, unintentional compression is normally undesirable, while operations aimed at obtaining defined shapes are usually required in some processes. The former will be Introduction 27

discussed in this section, whereas the latter will be included under the general scope of processing operations, in a subsequent part of this text.

Sone (1972) reported the following compressibility relationship, by tapping, for food powders:

$$\gamma_n = \frac{V_0 - V_n}{V_0} = \frac{abn}{1 + bn} \tag{1.13}$$

where γ_n is the volume reduction fraction, V_0 is the initial volume, V_n is the volume after n taps, and a and b are constants.

The applicability of Equation 1.13 was tested through its fit to the following linear form:

$$\frac{n}{\gamma_n} = \frac{1}{ab} + \frac{n}{a} \tag{1.14}$$

The constant a in Equations 1.13 and 1.14 represents the asymptotic level of the volume change or, in other words, the level obtained after a large number of tappings or a long time in vibration. The constant b is representative of the rate at which this compaction is achieved, that is, 1/b is the number of vibrations necessary to reach half of the asymptotic change. In general, this form of data presentation is very convenient for systems comparisons since it only involves two constants.

A very common undesirable aspect of compressibility is its negative influence on flowing capacity. Compression tests have been used widely in pharmaceutics, ceramics, metallurgy, civil engineering, and in the food powder field, as a simple and convenient technique to powder compressibility and flowability.

The pressure–density for powders in a compression test at low-pressure range can be described by the following equation (Barbosa-Canovas et al., 1987):

$$\frac{\rho(\sigma) - \rho_0}{\rho_0} = a + b \log \sigma \tag{1.15}$$

where $\rho(\sigma)$ is the bulk density under the applied normal stress σ , ρ_0 is the initial bulk density, and a and b are constants. The constant b represents, specifically, the compressibility of a given powder. Compression tests are useful in characterizing the flowability of powders because the interparticle forces that enable open structures in powder beds succumb under relatively low pressures. As shown in Equation 1.15, the constant b representing the change in bulk density by the applied stress is referred to as the powder compressibility. It has been found that b can be correlated with cohesion of a variety of powders and therefore could be a simple parameter to indicate

flowability changes (Peleg, 1977). Generally, the higher the compressibility the poorer the flowability, but if quantitative information about flowability is required, shear tests are necessary (Schubert, 1987).

One of the standard methods to evaluate the flowability of a particulate system is to calculate the Hausner ratio after tapping. The Hausner ratio is defined as the ratio of a powder system's initial (loose) bulk density to its tapped bulk density (i.e., the ratio of loose volume to tapped volume). It is easy to calculate the Hausner ratio and evaluate the flowability when the loose and tapped volumes of the test material are known. For a Hausner ratio of 1.0–1.1, the powder is classified as free flowing; 1.1–1.25, medium flowing; 1.25–1.4, difficult flowing; and >1.4, very difficult flowing (Hayes, 1987).

Adding small amounts of fine powders is often used to improve the flow properties of powdered materials in the chemical, pharmaceutical, and food industries. In this case, the fine particles coat the coarser particles of the main constituent and prevent them sticking together. Damp or sticky solids, which are difficult to handle, may be converted into free-flowing after the added fine powders absorb small quantities of liquids. Another way to improve flowability of food powders is by using the agglomeration process, which is accomplished by wetting the fine particles in an atmosphere of water or suitable solvent droplets, causing them to collide and stick together, and then drying the agglomerated material in an air stream. Apart from improving flowability, agglomerated powders may show better wettability and dispersibility in liquids, and tend to be dust-free (Hoseney, 1994). Agglomeration will be discussed further in Chapter 5 of this text.

Reconstitutability is the term used to describe the rate at which dried materials pick up and absorb water reverting to a condition which resembles the undried material, when put in contact with an excessive amount of this liquid (Masters, 1976). In the case of powdered dried biological materials, a number of properties may influence the overall reconstitution characteristics. For instance, wettability describes the capacity of the powder particles to absorb water on their surface, thus initiating reconstitution. Such a property depends largely on particle size. Small particles, representing a large surface area-to-mass ratio, may not be wetted individually. In fact, they may clump together sharing a wetted surface layer. This layer reduces the rate at which water penetrates into the particle clump. Increasing particle size and/ or agglomerating particles can reduce the incidence of clumping. The nature of the particle surface can also affect wettability. For example, the presence of free fat in the surface reduces wettability. The selective use of surface active agents, such as lecithin, can sometimes improve wettability in dried powders containing fat. Another important property is the sinkability, which describes the ability of the powder particles to sink quickly into the water. This depends mainly on the size and density of the particles. Larger, more dense particles sink more rapidly than finer, lighter ones. Particles with a high content of occluded air may be relatively large but exhibit poor sinkability because of their low density. Finally, dispersibility describes the ease Introduction 29

with which the powder may be distributed as single particles over the surface and throughout the bulk of the reconstituting water, while solubility refers to the rate and extent to which the components of the powder particles dissolve in the water. Dispersibility is reduced by clump formation and is improved when the sinkability is high, whereas solubility depends mainly on the chemical composition of the powder and its physical state.

References

- Abrahamsen, A. R. and Geldart, D. 1980. Behaviour of gas fluidized beds of fine powder. Part I: Homogeneous expansion. *Powder Technology* 26: 35–46.
- Allen, T. 1997. Particle Size Measurement. Vol. 1: Powder Sampling and Particle Size Measurement. London: Chapman & Hall.
- Barbosa-Canovas, G. V., Málave-López, J., and Peleg, M. 1987. Density and compressibility of selected food powders mixture. *Journal of Food Process Engineering* 10: 1–19.
- Beddow, J. K. 1981. *Particulate Science and Technology*. New York, NY: Chemical Publishing Co.
- British Standards Institution. 1975. BS 1377: 1975. *Methods of Tests for Soils for Civil Engineering Purposes*. London: BSI.
- British Standards Institution. 1993. BS 2955: 1993. *Glossary of Terms Related to Particle Technology*. London: BSI.
- Brown, G. G. 2005. *Unit Operations*. Delhi: CBS Publishers & Distributors.
- Dallavelle, J. M. 1943. *Micromeritics: The Technology of Fine Particles*. New York, NY: Pitman Publishing.
- Davies, R. 1984. Particle size measurement: Experimental techniques. In *Handbook of Powder Science and Technology*. Fayen, M. E. and Otten, L. (eds.). New York, NY: Van Nostrand Reinhold.
- Davis, G. E. 1904. A Handbook of Chemical Engineering. Manchester: Davis Bros.
- Ehrlich, R. and Weinberg, B. 1970. An exact method for characterization of grain shape. *Journal of Sedimentary Petrology* 40: 205–212.
- Ennis, B. J., Green, J., and Davies R. 1994. Particle technology: The legacy of neglect in the U.S. *Chemical Engineering Progress* 90: 32–43.
- Geldart, D. 1986. Gas Fluidization Technology. New York, NY: John Wiley & Sons.
- Gotoh, K. and Finney, J. L. 1975. Representation of size and shape of a single particle. *Powder Technology* 12: 125–130.
- Green, D. W. and R. H. Perry. 2008. *Perry's Chemical Engineer's Handbook*, 8th Ed. New York, NY: McGraw-Hill.
- Hayes, G. D. 1987. Food Engineering Data Handbook. New York, NY: John Wiley & Sons.
- Herdan, G. 1960. Small Particle Statistics. London: Butterworths.
- Hoseney, R. C. 1994. *Principles of Cereal Science and Technology*. St. Paul, MN: American Association of Cereal Chemists.
- Jones, S. 1983. The problem of closure in the Zahn–Roskies method of shape description. *Powder Technology* 34: 93–94.

- Kaye, B. H. 1981. Small Characterization of Fine Particles. New York, NY: John Wiley & Sons.
- Luerkens, D. W., Beddow, J. K., and Vetter, A. F. 1982. Morphological Fourier descriptors. *Powder Technology* 31: 209–215.
- Ma, L., Davis, D. C., Obaldo, L. G., and Barbosa-Canovas, G. V. 1997. Mass and spatial characterization of biological materials. In *Engineering Properties of Foods and Other Biological Materials*. Pullman, WA: Washington State University Publisher.
- Mandelbrot, B. B. 1977. Fractals, Form, Chance and Dimension. San Francisco, CA: Freeman.
- Mandelbrot, B. B. 1982. The Fractal Geometry of Nature. San Francisco, CA: Freeman.
- Masters K. 1976. Spray Drying Handbook. New York, NY: John Wiley & Sons.
- McCabe, W. L. and J. C. Smith. 1956. *Unit Operations of Chemical Engineering*. New York, NY: McGraw-Hill.
- McGraw-Hill Dictionary of Scientific and Technical Terms, 6th Ed. 2003. New York, NY: McGraw-Hill Professional.
- Okuyama, K. and Kousaka, Y. 1991. Particle density. In *Powder Technology Handbook*. Iinoya, K., Gotoh, K. and Higashitani, K. (eds.). New York, NY: Marcel Dekker. Orr, C. 1966. *Particulate Technology*. New York, NY: Macmillan.
- Parfitt, G. D. and Sing, K. S. W. 1976. *Characterization of Powder Surfaces*. New York, NY: Academic Press.
- Peleg, M. 1977. Flowability of food powders and methods for its evaluation—A review. *Journal of Food Process Engineering* 1: 303–328.
- Rhodes, M. J. 1990. Principles of Powder Technology. New York, NY: John Wiley & Sons.
- Riley, G. S. 1968/1969. An examination of the separation of differently shaped particles. *Powder Technology* 2: 315–319.
- Schubert, H. 1987. Food particle technology part I: Properties of particles and particulate food systems. *Journal of Food Engineering* 6: 1–32.
- Schulze, D. 2007. *Powders and Bulk Solids: Behavior, Characterization, Storage and Flow.* New York, NY: Springer.
- Sebestyen, G. S. 1959. On Pattern Recognition with Application to Silhouettes. DSc Thesis. Cambridge, MA: Massachusetts Institute of Technology.
- Shinohava, K. 1979. Mechanism of segregation of differently shaped particles in filling containers. *Industrial Engineering Chemical Process Design and Development* 18: 223–227
- Sone, T. 1972. Consistency of Foodstuffs. Dordrecht, The Netherlands: D. Reidel.
- Svarovsky, L. 2000. Solid–Liquid Separation. Oxford: Butterworth-Heinemann.
- Tardos, G. I. 1995. Development of a powder technology option at CCNY. *Chemical Engineering Education* (Summer 1995): 172–177, 181.
- Valiente, A. and R. P. Stivalet, 1980. El Ingeniero Químico, ¿Qué Hace? México City: Alahambra Mexicana.
- Walker, W. H., Lewis, W. K., and W. H. McAdams. 1937. *Principles of Chemical Engineering*. New York, NY: McGraw-Hill.

Bulk Solids: Properties and Characterization

2.1 Introductory Aspects

Taking into account all the sizes and shapes that may be found in solids, the most important from the standpoint of chemical engineering is the small particle. An understanding of the characteristics and features of minute solids populations is needed for designing processes and equipment dealing with streams containing such solids. As opposed to chunk solid pieces, finely divided solids can be referred to by denominations as varied as particulate solids, pulverized solids, granular materials, and so on. In process engineering literature (chemical engineering, mechanical engineering, chemistry, physics, materials science, etc.), a generic term having been used to describe a huge population of small particles forming a defined material, is simply "powder" or "industrial powder." Powders present some characteristics so distinctive that make them practically different, by comparison, with any other state of matter. Intrinsically, molecules forming a solid piece are joined by chemical bonds and interactions stronger than those forming liquid or gases. Externally, however, there are no interactions of the chemical bond type keeping together small particles forming a powder batch. It has been suggested that the individual entities within masses of particulate solids are so kept only by geometrical accommodation. In this chapter, the most important distinctive characteristics of industrial powders are described and discussed, in the context of their importance and relevance within the discipline of Powder Technology or Particle Technology.

Some of the distinctive characteristics of powders, that have made them subject of research efforts to understand their behavior are

- Powders are not solids, but may deform under compression.
- Powders are not liquids, but they may flow under certain circumstances.
- Powders are not gases, but they may be compressed up to a degree.

Considering the powders' features listed above, it becomes clear that even a definition of powder would be difficult to establish. A first approach could

be as follows: "A powder is a group of solid particles not filling completely the space they occupy; the space not filled by particles is filled by gas." This definition results somewhat inappropriate since nobody will call powder to a pile of rocks. In searching for a more appropriate definition, it has been observed for quite a while that dry powders possess many properties common to fluids, such as exerting pressure on container vessels and flowing through channels or orifices. There are, nonetheless, important differences such as

- A powder does not exert uniform pressure on all directions in confinement. The exerted pressure is minimal at the perpendicular direction of the applied pressure.
- An applied shear force on the surface of a mass of powder is transmitted through all the static mass, unless a fracture occurs.
- The density of a mass of powder varies depending on its degree of packaging; it increases if the powder is compacted by vibration, shaking, taping, and so on.

The above-listed characteristics originated, from the 1960s, a special focus of attention on the fluid-related properties that a powder may acquire under certain conditions. If powders would be able to follow some of the typical behavior of fluids, it would be possible to study them making use of modified fluid mechanics theoretical models. A suitable manner of making a powder behave as a fluid is to suspend its individual particles within a stream of gas. When any stable suspension of solid particles of a powder in a gas stream is achieved, it is said that the powder has been fluidized. A powder that is able to be fluidized would be most properly defined as: "Disperse diphasic system in which the disperse phase consists on particles of a finely divided solid and the continuous phase consists of a gas." The solid particles conform to a sort of mechanical network due to certain interparticle forces. The net can be expanded under some conditions, but will reassume its packed state under the influence of gravity. The packed, stationary state of a mass of powder presents mechanical resistance and some degree of elasticity. The interaction particle-gas is fundamental, and determines primarily the flowing capacity of the powder.

2.2 Classification of Powders

The different ways of classifying powders is closely related to the applications in which bulk properties play an important role. Since most applications in powder technology are relevant when considering powders as dispersion systems, they can be classified into different categories according to their particular

behavior in dynamic situations. For this reason, powders or dry particulate materials can be categorized according to their handling properties. The classification depends on the type of handling in question, with the cases of deaerated state and aerated state being most important. The classification of powders in the deaerated state is known as Jenike's classification, while the classification of powders in the aerated state is called Geldart's classification.

2.2.1 Jenike's Classification

In the deaerated state, powders can be classified using a shear cell tester or a compression tackiness tester. A precise definition of the flowability of a powder is only possible with several number of curves derived from a family of yield loci of the powder, measured with a shear cell. There are several designs of shear cells that are also used to measure other properties of powders, and so they will be discussed in detail in a further section, but basically they measure different shearing forces for corresponding normal forces on consolidated powder samples. The curve representing the relation between the maximum shear and normal stresses is called the yield locus. Each yield locus also gives one pair of values for the unconfined yield stress σ_c and major consolidation stress σ_1 . The ratio of the consolidation stress to the unconfined yield strength is called the flow factor ff. Jenike (1964) proposed a classification according to the position of one point of the failure function at a fixed value of the unconfined yield stress with respect to the flow factor line. Figure 2.1 shows a schematic representation of classification of powders following this criterion. As can be seen, at a fixed value of the unconfined yield strength of 22.3 N, the straight lines through the origin at a slope 1/ff would represent the categories of very cohesive, cohesive, easy flowing, and

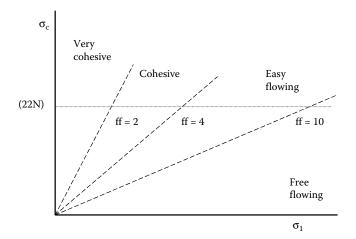


FIGURE 2.1Graphical representation of classification of powders according to Jenike.

Classification of Powders According to Jenike		
Type of Powder	ff Value at $\sigma_c = 22.3 \text{ N}$	
Very cohesive	Less than 2	
Cohesive	Between 2 and 4	
Easy flow	Between 4 and 10	
Free flow	More than 10	

TABLE 2.1Classification of Powders According to Jenike

free flowing, as represented in the mentioned figure, as well as in Table 2.1. In addition to Jenike's classification, Williams et al. (1970/1971) has also defined simple powders as those whose failure function is a straight line through the origin, providing that the slope of the line is less than 1/ff.

Another way of classifying powders is on the basis of the results obtained with a compression tackiness tester. This tester is a simplified version of the uniaxial compression test, on a cylindrical briquette of the powder, where the compression force at failure is a direct measure of the powder tackiness. The consolidation stresses used to form this briquette are generally greater than those used with the Jenike shear test. The total compression force is supposed to simulate pressures greater than that of the bottom of a 180 cm high pile of bags in storage. Tackiness values above 27 N usually indicate powders that can cause problems due to their sticky characteristics and the shape of a curve of tackiness value against initial compression load is also an indication of tackiness: nontacky powders give essentially a horizontal line while tacky products show the effect of increasing compression.

2.2.2 Geldart's Classification

With regard to the aerated state, a widely accepted classification of powders is the one proposed by Geldart (1973), which takes the two most important particle properties into account: particle size and particle density. Geldart's classification is basically derived from the behavior of powders when fluidizing by air at ambient conditions. In the fluidizing state, a bed of powder starts expanding when the fluidizing air reaches a critical velocity known as minimum fluidizing velocity. Considering the different ways of powders in expansion, Geldart suggested four types of powders. Figure 2.2 illustrates the four general manners in which different powders behave when subjected to these described conditions of fluidization. According to such figure, powders can be termed as A, B, C, and D type and may be described as follows. A powders are sometimes referred to as slightly cohesive or catalyst type and exhibit large bed expansion after minimum fluidization and before initiation of bubbling. The air velocity at which bubbling begins is known as the minimum bubbling velocity. Increasing this velocity will increase number and size of bubbles. B powders are known as sand like and present

35

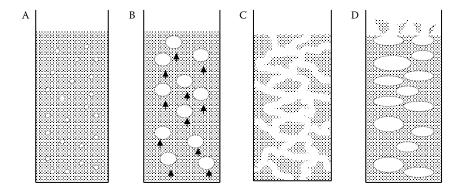


FIGURE 2.2 Behavior of powder beds when fluidized.

bubbling at the minimum fluidization velocity with a small bed expansion. In this type of powders the minimum fluidizing velocity equals the minimum bubbling velocity so that bubbles appear immediately with bed expansion. This bed expansion is said to be due to bubbling only. C powders are called cohesive and are difficult to fluidize at all. When the minimum expansion velocity is surpassed, these powders do not expand or form bubbles. The air is released through cracks forming irregular channel paths within their mass volume. They do not show any kind of expansion because the pressure drop through the bed is considerably lower than its weight over the cross-sectional area. D powders are simply known as large and can form stable spouted beds if the air is admitted only through a centrally positioned hole. Over the minimum fluidizing velocity these powders form large bubbles, which burst when reaching the surface so that air is released and spouting is observed. Geldart's classification can also be represented in a plot of particle fluid density difference versus particle size, as shown in Figure 2.3. As can be observed, D powders are extremely large to even be considered powders. It has been observed that their behavior is determined exclusively by hydrodynamics and that interparticle forces do not play a role.

The above-mentioned classification of powders may be useful even when fluidization as such is not of interest. For example, in the filling of containers and in the residence time needed in hoppers to avoid flooding out when the discharge valve is opened, the bed expansion and the rate of deaeration are of concern. There is a striking difference between A, B, and C powders in the way they behave in deaeration: A-type powders collapse at a constant rate, B-type powders deaerate almost instantaneously, while C-type powders collapse initially much faster than those in group A but may remain in a slightly aerated state for a considerable period, with the pressure at the bottom of the bed decaying very slowly. The deaeration rate can, therefore, also be used to characterize powders, but Geldart and Wong (1985) found this to be insufficiently sensitive and developed a standardized deaeration time test.

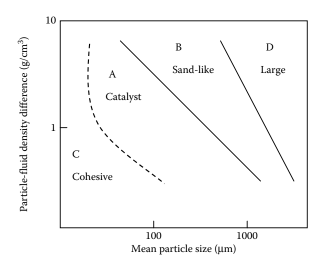


FIGURE 2.3Graphical representation of classification of powders according to Geldart.

Geldart's classification has been well accepted by powder technologists since its release. Some powders, however, may present behavior variations when certain conditions may change. For example, an A powder could show the behavior of a B powder if effective gravity would increase by using a gas with viscosity lower than air viscosity. Rietema et al. (1980) have suggested a generic classification by using dimensionless groups, which takes into account important variables such as cohesion, gas viscosity, and acceleration due to gravity. According to this approach, dimensionless groups may be used to represent the behavior limit between powders A and B, as well as between powders A and C. This approach does not consider, as previously mentioned, D systems of particles as powders. For the limit of behavior of powders A and B, theoretical principles discussed elsewhere (Rietema and Mutsers, 1973; Mutsers and Rietema, 1977) have shown that homogeneous bed expansion is possible when

$$FI < \left[\frac{150(1-\varepsilon)}{\varepsilon^2(3-2\varepsilon)}\right]^2 \tag{2.1}$$

where Fl is the fluidization number and ϵ is the porosity or voidage of the bed.

The limit of behavior between A and B powders is, therefore, represented by

$$Fl = \left\lceil \frac{150(1-\epsilon)}{\epsilon^2(3-2\epsilon)} \right\rceil^2$$
 (2.2)

In terms of the limit of behavior between A and C powders, a dimensionless Group known as cohesion number Coh, is defined as

$$Coh = \frac{C}{(\rho_s)(x)g}$$
 (2.3)

where C is a constant known as the constant of powder cohesion, ρ_s is the powder density, x is an equivalent diameter of the particle, and g is the acceleration due to gravity.

In the compacted (or consolidated) state the elasticity module of a powder has a value approximately equal to the value the cohesion constant. The fluidization number for a powder at its consolidated state can be, therefore, modified as

$$Fl^* = \frac{\rho_s^3 x^4 g^2}{\mu_g^2 C}$$
 (2.4)

where Fl* is the fluidization number for consolidated powders, and μ_g is the gas viscosity.

Equation 2.4 is, actually, the product of the inverse of the cohesion number and another dimensionless group known as the Archimedes number Ar, and thus

$$Fl^* = Ar(Coh)^{-1}$$
 (2.5)

The Archimedes number Ar is defined as

$$Ar = \frac{\rho_s^2 x^3 g}{\mu_g} \tag{2.6}$$

When values of Coh versus Ar are plotted, the resulting graph will show the behavior limits of powders A and B and powders A and C, as represented in Figure 2.4. As can be observed, the behavior limits of powders A and B are function of the bed voidage.

2.3 Sampling

Sampling is an important element of powder handling that demands careful scientific design and operation of the sampling systems. The general purpose of sampling is to collect a manageable mass of material which must be

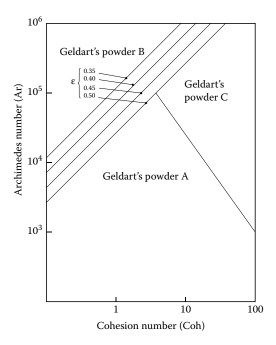


FIGURE 2.4 Dimensionless representation of Geldart's classification for powders A, B, and C.

representative of the total mass of the sampled powder. This action is achieved by taking many small samples from all parts of the total which, when combined, will represent this total with an acceptable degree of accuracy. All particles in the total must have the same probability of being included in the final sample, so all of them must be equally accessible. To satisfy these requirements, the following basic "golden rules" of sampling should be applied whenever possible:

- Sampling should be done preferably from a moving stream (for both powders and suspensions) but powder on a stopped belt can be sampled.
- A sample of the whole of the stream should be taken for many (equally spaced) periods of time, rather than part of the stream for the whole of the time.

The first rule recommends that the sample should be taken from flowing powder stream, such as discharging from a belt conveyor or a feeding flow from one storage container to another. Also, as the second rule mentions, the sampling process should continue for a long series of short time intervals (Masuda, 1997). It is very likely that the recombined, primary sample taken

from the whole will be too large for most powders tests. It will be, therefore, necessary to subdivide the original sample into secondary, or even tertiary, subsamples. This subdivision may be built into the primary sampler or it may be achieved with a separate sampling divider.

If a sampled material is perfectly homogeneous with respect to its properties, any fraction of the bulk would be exactly the same as far as those properties are concerned. For instance, if a certain powder is heterogeneous on size but homogeneous on density, it would be considered homogeneous if it is sampled to determine density. This is the reason why it is often easier to obtain a representative sample from liquid materials, where all their properties are, generally, homogeneous. For example, when sampling ionic solutions for component concentration, any variation in results would be normally attributed to experimental error. However, when the material is heterogeneous, as it virtually happens with industrial powders, it is expected to find difference in some measured properties. The source of these variations, in addition to variations caused by the assaying process, could be attributed to the fact that the smaller fragments of particulate material are themselves heterogeneous and to segregation of certain types of fragments due to handling of the bulk material (Gy, 1982). The probability of obtaining a perfect unbiased sample from the parent material is remote. If several samples are taken and they are representative, the expected variation may be estimated from statistical analysis (Allen, 1981).

The statistical theory of sampling considers a sample of size N (N specimens) selected randomly and independently. A random selection process must be such that every member of the population being sampled would have the same chance to be selected. The independency of specimens implies that the selection of one specimen does not influence the selection of another in the whole mass of the sampled powder. These requirements are not usually met in practice, either due to the two sources of variations previously mentioned or to the nature of the sampling technique, or both. For example, a perfectly blended batch of a binary mixture of powders may become segregated during the sampling process due to the nature of the mechanical handling system. The act of sampling can be considered a process that allows observing the realizations of a random variable whose statistical properties are determined by the inherent characteristics of the sampled population and the sampling process. Naturally, the apparent variability of the determined values can be greatly influenced by the sampling and measurement techniques. In statistics, a random variable is defined as a function which assigns real numbers to the outcomes of a random experiment. For particulate materials, the random experiment consists of: the selection of a fraction of a given volume of bulk by giving equal chance to all other fractions in the lot, further reduction of volume and selection of a fraction of the specimen for testing, and testing of a final fraction for different properties. The random variable being observed is defined by the sampling process. A change in the volume of the specimen in the sampling process will correspond to a

different experiment, and so it will result in the realization of a different random variable. When sampling from a well-mixed bulk of particulate material, it can be postulated that the random variables observed by choosing different specimen volumes would possess the same distribution function and the same mean, but different variances. The theoretical and experimental studies indicate that, keeping other things constant, reduction of specimen volume increases the variance. A practical relationship has been proposed by Gy (1982), which is useful in determining the minimum acceptable specimen weight as a function of the diameter of the coarsest fraction in the total mass of sampled powder. Such relationship can be represented as

$$M_{\rm s} \ge \frac{Cx^3}{\sigma^2} \tag{2.7}$$

where σ^2 is the variance of the tolerated sampling error, C is a constant characterizing the material to be sampled, x is the diameter of the coarsest fragment, and $M_{\rm s}$ is the weight of the specimen.

The arithmetic average of random independent observations of a normally distributed random variable is known to be the best estimator of the unknown population mean. The Central Limit Theorem (Kennedy and Neville, 1986) asserts that the arithmetic average of random independent observations of a random variable will be distributed normally when the sample size is infinitely large. Even for moderately sample sizes, the statistical behavior if the sample average is acceptably close to that of the Normal distribution. Also, by increasing the number of specimens (sample size), sample averages closer to the unknown value of the population mean are obtained. The Law of Large Numbers (Larson, 1995) establishes that when the sample size is infinitely large, the sample mean becomes equal to the population mean. In practice, however, small sample amounts are handled for obvious practical and economic reasons, and so it is practically impossible for the sample average to become equal to the unknown population mean. For this reason, it is used to establish an interval around the sample average, called a confidence interval, which will contain the unknown mean with a certain predetermined probability. Given the probability $(1-\alpha)$, the length of the confidence interval becomes a function of the population variance and the sample size. This relationship is derived from the following probability statement:

$$\operatorname{Prob}\left[\frac{\bar{X} - Z_{(1-\alpha)/2}\sigma}{\sqrt{N}} \le \mu \le \frac{\bar{X} + Z_{(1-\alpha)/2}\sigma}{\sqrt{N}}\right] = 1 - \alpha \tag{2.8}$$

where \overline{X} is the sample arithmetic average, σ is the population standard deviation, μ is the population mean and N is the sample size, and $Z_{(1-\alpha)/2}$ is the $(1-\alpha)/2$ percentile of standard Normal variable.

Bulk Solids 41

The length *L* of the confidence interval is

$$L = \frac{2Z_{(1-\alpha)/2}\sigma}{\sqrt{N}} \tag{2.9}$$

It is clear from Equation 2.9 that the length of the confidence interval is linearly proportional to the population standard deviation, and inversely related to the square root of the sample size. If σ were known, Equation 2.9 could be used to determine the minimum sample size required to obtain a confidence interval which will contain the unknown mean μ , with a $(1-\alpha)$ probability. An expression for the minimum sample size will, therefore, be

$$N \ge \frac{4Z_{(1-\alpha)/2}^2 \sigma^2}{L^2} \tag{2.10}$$

Since σ is usually unknown and the sampling process has normally the aim of estimating the standard deviation as well as the mean, Equation 2.10 has limited applicability but indicates, however, an important relationship between the sample size, the inherent variability of the sampled population, and the precision at which the mean μ is to be estimated. Such precision will be increased by reducing L and/or increasing (1- α), both resulting in larger sample sizes. Also, as indicated by the power of σ in Equation 2.10, its variability will increase the sample size quadratically. Reduction of σ is only possible by employing a different sampling method, further crushing or grinding the bulk to be sampled, or increasing the specimen volume. Considering that all these measures are not very practical for real purposes, an alternative would be the use of an estimator for σ , which is a well-known practice in statistics. A relationship for estimating the population standard deviation can be represented as follows:

$$S^{2} = \frac{1}{N-1} \sum_{i=1}^{N} (X_{i} - \bar{X})^{2}$$
 (2.11)

where *S* is the estimator for σ , and X_i is the assay value.

Since the variance is not known, but estimated by *S*, the following probability statement can be established:

$$\Pr{\text{ob}\left[\frac{\bar{X} - t_{(1-\alpha)/2, N-1} S}{\sqrt{N}} \le \mu \le \frac{\bar{X} + t_{(1-\alpha)/2, N-1} S}{\sqrt{N}}\right]} = 1 - \alpha \tag{2.12}$$

where S is the sample standard deviation, and $t_{(1-\alpha)/(2, N-1)}$ is the $(1-\alpha)/2$ percentile of the Student's t distribution with N-1 degrees of freedom.

The length of the confidence interval can be expressed by

$$L = \frac{2t_{(1-\alpha)/2, N-1}S}{\sqrt{N}} \tag{2.13}$$

An expression for the minimum sample size can be derived from Equation 2.13, similarly to the one derived from Equation 2.11, that is,

$$N \ge \frac{4t_{(1-\alpha)/2,N-1}^2 S^2}{L^2} \tag{2.14}$$

By inspecting Equations 2.11 and 2.14, it can be noticed that the values of S and t are not independent of N. Also, a sample of a certain size is needed, for example, N', to obtain S initially. This value will change as new specimens are sampled, assayed, and their values used in recalculating S. When the population variance is unknown, it is not possible to determine exactly the minimum sample size. An approach can be achieved by taking several samples and calculating successive approximations of N. To accomplish this, using a small sample size the estimator S is determined by means of Equation 2.11 and a new value for *N* is calculated with appropriate values of $t_{(1-\alpha)/(2, N-1)}$, S, and L using Equation 2.11. If this value is larger than the actual sample size, additional sampling is done and the procedure repeated. As the sample size increases the value of S will approach the unknown population standard deviation with decreasing error. However, the successive values of S as sample size increases may fluctuate around σ with decreasing error and, thus, the calculated values of N will also fluctuate. For this reason, it is advisable to increase the sample size by smallest possible increments, and continually calculate N until its value and the actual sample size are sufficiently close.

A practical approximation can be done considering the ratio L^2/S^2 in Equation 2.14 and estimating the population mean for a given confidence interval, such that its length be equal to a certain multiple of S. In such a manner, an initial determination of the sample size is possible. By rearranging terms in Equation 2.14, the following relationship can be obtained:

$$U \ge \frac{4t_{(1-\alpha)/2,N-1}^2}{N} \tag{2.15}$$

where $U = L^2/S^2$.

Equation 2.15 can be used to calculate different values of U determined by certain values of α and N. The value of the percentiles of Student's t distribution, are available in most of the standard textbooks on statistics. The results of such calculations for different confidence levels can be presented in graphical or tabular form. Table 2.2 lists U values for sample

TABLE 2.2 Sample Size as a Function of *U*

	$U = L^2/S^2$		
Sample Size N	$1-\alpha = 0.90$	$1-\alpha = 0.95$	$1-\alpha = 0.99$
2	79.63	323.09	8105.19
3	11.37	24.65	131.21
4	5.52	10.11	34.11
5	3.63	6.18	16.93
6	2.69	4.40	10.83
7	2.15	3.43	7.87
8	1.81	2.78	6.13
9	1.54	2.37	5.02
10	1.34	2.04	4.23
11	1.19	1.81	3.65
12	1.08	1.61	3.22
13	0.97	1.46	2.88
14	0.90	1.33	2.59
15	0.83	1.22	2.37
16	0.77	1.13	2.18
17	0.72	1.06	1.89
18	0.67	0.99	1.77
19	0.63	0.93	1.66
20	0.60	0.87	1.64
21	0.56	0.83	1.54
22	0.54	0.79	1.46
23	0.51	0.75	1.38
24	0.49	0.71	1.32
25	0.47	0.68	1.25
26	0.45	0.65	1.20
27	0.43	0.63	1.14
28	0.41	0.60	1.10
29	0.40	0.58	1.05
30	0.39	0.55	1.02
40	0.28	0.41	0.73
50	0.22	0.32	0.57
60	0.19	0.27	0.47
80	0.14	0.20	0.35
100	0.11	0.16	0.28
200	0.05	0.08	0.14
500	0.02	0.03	0.05

sizes from 2 to 500 for confidence levels of 0.90, 0.95, and 0.99. It can be used to read directly the required minimum sample size for a predetermined value of U. Table 2.2 shows that, in agreement with probability principles, as the confidence interval length L shortens with respect to the estimator of the standard deviation S, the unknown mean and standard deviation

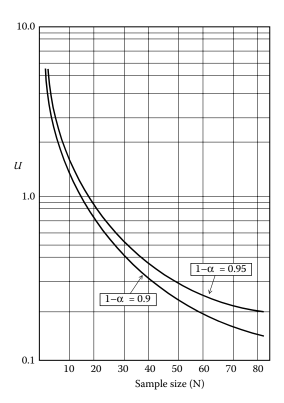


FIGURE 2.5 Relationship between U and L for two confidence intervals.

will be predicted with high certainty and so a large sample size N will be the most appropriate for such prediction. On the other hand, for a large value of L and a small value of S, the predictions of unknown mean and standard deviation will be very uncertain and the sample size N relatively low. A good approximation is to use a confidence interval length L equal to the standard deviation estimator S, and so the value of U would equal unity (Dodge and Romig, 1998). Figure 2.5 shows a plot of the tabulated values of Table 2.2. Considering all theoretical aspects of sampling discussed above, as well as field experience, it is recommended for practical purposes to use a sample size between 30 and 50 for most powdered and particulate materials.

Example 2.1

The size of a fine powder sample to be used in a process of manufacturing ceramics is to be determined. The interval *L* and the estimator *S* are approximately similar. A 95% confidence interval is considered appropriate for the sampling procedure.

SOLUTION

Since L and S are approximate equal, the value of U (L^2/S^2) will approach unity. Consequently, the interception of U=1 with the curve for $1-\alpha=0.95$ in Figure 2.5 will read the value of 18 directly. Alternatively, reading from Table 2.2 for U=0.99 and $1-\alpha=0.95$, the size sample is 18.

Sampling methods are normally designed or selected for collection of primary samples from large-scale processes. In many applications, samplers have to be designed specifically for a particular material and duty. Usually, there are two main stages in any sampling scheme. A gross sample, which represents the parent batch, is initially collected and, then, a representative laboratory sample is prepared from this gross sample. These two operations are generally performed by different techniques. Gross samples can range up to 50 kg or more, depending of the size of the bulk quantity, while laboratory samples rarely exceed 1 or 2 kg, or even less considering the wide array of state-of-the-art instrumental methods currently available for powder characterization. This small amount of sample may require further subdivision, as to use an appropriate number of specimens determined by the minimum sample size previously discussed.

Segregation is one of the main troubles encountered in the sampling of solid particles. In conical heaps of material, for instance, a large proportion of the coarser particles will generally be found in the lower levels of the pile. On the other hand, coarse particles tend to migrate toward the top of the contents of a container, especially after transportation. Any selected sampling method must take into account these irregularities. Grab sampling with scoops or shovels is considered a common alternative for obtaining bulk samples. A large number of random samples may be taken from a heap of the material, or regular samples (e.g., every *n*th shovelful) can be scooped into a sample container when the bulk material is transported manually from one location to another. Regular sampling is most often favored for materials packed in containers. A common sampling implement for this purpose is a piece of metal tubing with a sharp bottom edge and fixed to a long handle, known as sampling "thief." This sampler is pushed gently, firmly, and vertically into the full depth of the material, and then withdrawn for sample removal. When sampling from stored powders the samples have to be withdrawn from small spaces of equal volume within the total, in an effort to try to give a chance to every member of the population to be sampled. Since this is very difficult to achieve as not all of the material is equally accessible, sampling from stationary places is generally to be avoided.

For material flowing down chutes or from hoppers, a sample collector may be placed in the path of the outlet stream. Open-ended scoops or shovels are unsuitable in this case because they allow coarser particles to roll out of the sample and thus bias the sample toward the finer fractions. A ladle is the only recommended device for such sampling (Plowman, 1985) since it does not allow this rolling out to happen, unless overfilled. The falling stream is traversed with the laden to collect the sample, and if the whole of the stream

cannot be sampled in one pass, it is divided into several equal areas with the sample being collected from those incrementally. In agreement with the golden rules of sampling mentioned before, the collector should sample the whole of the outflow stream at regular time intervals.

Stopped belt sampling is considered the best manual sampling method. It is often taken as a reference method with others being compared against it. It follows the two golden rules of sampling in that it samples from a moving stream (with the movement momentarily stopped for taking the sample) and it takes the whole of the stream many times. It is carried out with a suitable profiled sampling frame which is inserted through the material on a stopped belt conveyor until it comes in contact with the conveyor belt over its full width. The material within the frame is then collected from the belt and it represents one increment in the manual sampling series.

At the laboratory level, gross samples can be subdivided into one or more smaller samples manually or mechanically. The most common manual method is the one known as "coning and quartering," which consists in mixing the sample and pile it into a conical heap, flatten it about one quarter of its original height, divide the flatten heap into four equal quarters, and reject two opposite quarters while remixing the remaining pair. This remaining amount is piled again into a new heap, and the procedure is repeated until the required sample quantity is obtained. As mentioned before, sampling methods have sometimes to be designed for specific purposes. Also, the many stages of a sampling scheme often required, makes sampling a technique which has to be carried out very carefully, and taking many samples is normally involved in the whole process.

Sample dividers are used to reduce the size of the primary sample and can be classified as intermittent and reciprocating cutters, as well as continuous and rotational dividers. The rotational dividers are usually considered more suitable than linear cutters. A common type of divider, shown in Figure 2.6, is a box with an open top divided into a number of compartments with their bottoms sloping toward opposite sides of the box. When a quantity of material is poured evenly into the hopper, it is split into equal portions. Thus, by successive repetitions of the procedure, the sample can be reduced to the minimum amount required to perform an assay. An alternative to reduce a sample to a given proportion in a single step can be performed with a sample reducer, such as the one illustrated in Figure 2.7. As can be seen, by pouring the sample through the hopper at the top of the inclined plane, a fixed reduction of the whole amount of sample can be obtained. A very efficient sample divider is the spinning riffler (Allen and Khan, 1970), which follows quite well both golden rules of sampling. As can be seen in Figure 2.8, such sampling splitter comprises a hopper mounted above a vibratory feeder, and a rotating carousel divided into many compartments. In operation, the sample is slowly conveyed from the hopper by the feeder onto sampling containers fixed to the compartments. The subsamples are collected into such containers, and the feed rate is controlled by varying the gap under the hopper, as

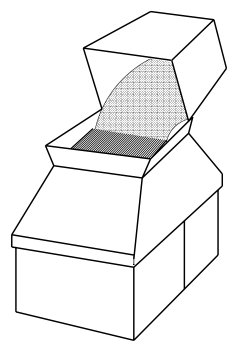


FIGURE 2.6 Sample divider or chute sample splitter.

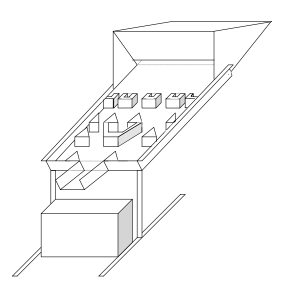


FIGURE 2.7 Diagram of a sample reducer.

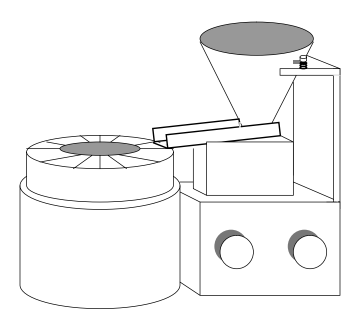


FIGURE 2.8 Schematic diagram of a rotary sample divider (spinning riffler).

well as the electromagnetic vibration of the feeder. Many commercial instruments based on this principle are available, covering sampling amounts as little as 25 g, or as large as 40 kg or powder, or even more.

There are many other types of samplers and sample dividers commercially available. Since sampling is a crucial step that may provide guidance about proper operation of an entire process, great care should be taken as to produce samples which will render results as reliable as possible. Due to some properties of powders and granular materials, such as those mentioned at the beginning of this chapter, in any specific design of sampler is advisable to verify whether it follows the so called golden rules of sampling for the particular application.

2.4 Failure Properties

2.4.1 Introduction

Powder flow is defined as the relative movement of a bulk of particles among neighboring particles or along the container wall surface (Peleg, 1977). The practical objective of powder flowability investigations is to provide both qualitative and quantitative knowledge of powder behavior, which can be

Bulk Solids 49

used in equipment design and in equipment performance prediction (Sutton, 1976). The flow characteristics of powders are of great importance in many problems encountered in bulk material handling processes in diverse types of industries, such as the agricultural, ceramic, chemical, food, mineral, mining, and pharmaceutical industries, because the ease of powder conveying, blending, and packaging depends on them (Chen, 1994). Practically, for all fine and very fine powders, the attractive forces between particles are large when compared with the weight of individual particles and so they are said to be cohesive and normally present flow problems. For example, such materials can fail to flow out of a container in which there is a hole, thousand of times larger than the diameter of the particle. To guarantee steady and reliable flow, it is crucial to accurately characterize the flow behavior of powders. The forces involved in powder flow are gravity, friction, cohesion (interparticle attraction), and adhesion (particle-wall attraction). The first requirement is to identify the properties which characterize the flowability of a particular material and to specify procedures for measuring them. The common belief that the flowability of a powder is a direct function of the angle of the repose is misleading and should be avoided because, as stated earlier, most industrial powders show different grades of cohesiveness. The angle of repose of a cohesive material is indeterminate, being dependent on the previous story of a given sample, and is irrelevant to the flow behavior of the material in any particular situation. The behavior of the material depends on the manner in which its strength varies with its state of compaction. For example, a slightly cohesive powder may be able to flow and be poured through a hole (forming a pile with a slope equal to the angle of repose), but after being compressed by simple tapping the flow would not develop due to the formation of a stable cohesive arch over the hole. A very small change in the bulk density of the powder would have, therefore, caused a dramatic change in its flowability. The important feature of a particulate material is the way its shear strength varies with the consolidating stress, and the properties used to identify and quantify such interactions are commonly known as the failure properties of a powder.

The requirement for getting powders to flow is for their strength to be less than the load put on them, that is, they must fail. The basic properties describing this condition are known as "failure properties" and they are the following: (a) the effective angle of internal friction, (b) the angle of wall friction, (c) the failure function, (d) the cohesion, and (e) the ultimate tensile strength. The failure properties take into account the state of compaction of the powder as this strongly affects its flowability unless the powder is cohesionless, like dry sand, and it gains no strength on compression. The time of consolidation can also have an effect on failure properties of powders. It is, therefore, important to test such properties under controlled conditions using sealed powder samples or air-conditioned rooms or enclosures. Also, time-consolidating samples must be tested to simulate storage conditions.

2.4.2 Description of Failure Properties

The effective angle of internal friction δ is a measure of the friction between particles and depends on their size, shape, roughness, and hardness. The angle of wall friction ϕ is equivalent to the angle of friction between two solid surfaces except that one of the two surfaces is a powder. It describes the friction between the powder and the material of construction used to confine the powder, for example, a hopper wall. The wall friction causes some of the weight to be supported by the walls of a hopper. The failure function FF is a graph showing the relationship between unconfined yield stress (or the strength of a free surface of the powder) and the consolidating stress, and gives the strength of the cohesive material in the surface of an arch as a function of the stress under which the arch was formed. The cohesion C is, as mentioned earlier, is a function of interparticle attraction and is due to the effect of internal forces within the bulk that tend to prevent planar sliding of one internal surface of particles upon another. The ultimate tensile strength T of a powder compact is the most fundamental strength mechanism, representing the minimum force required to cause separation of the bulk structure without major complications of particle disturbances within the plane of failure.

There are several ways, direct or indirect, of testing the five failure properties defined above. Basically, all of them can be determined using a shear cell, but simplified or alternative procedures can be adopted when the aim is to monitor the flowability of the output from a process or compare a number of materials.

2.4.3 Experimental Determinations

There are basically three types of shear cells available for powder testing: (a) the Jenike shear cell, also known as the translational shear box; (b) the annular or ring shear cell, also called the rotational shear box; and (c) the rotational shear cell, which is a fixture of a powder rheometer.

The Jenike shear cell is circular in cross section, with an internal diameter of 95 mm. A vertical cross section through the cell is shown in Figure 2.9. It consists of a base, a ring which can slide horizontally over the base, and a cover. The Jenike test sequence simulates the changes in stresses acting on an element of materials flowing through bin. The ring and base are filled with the powder and a lid is placed in position. By means of a weight carrier, which hangs from a point at the center of the lid, a vertical compacting load can be applied to the powder sample. The lid carries a bracket with a projecting pin and a measured horizontal force is applied to such bracket, causing the ring and its content, as well as the lid, to move forward at a constant speed. The procedure to carry out a test on a powder consists of three steps: preconsolidation to ensure uniformity of samples, consolidation to reproduce flow with a given stress under steady-state conditions, and shearing to

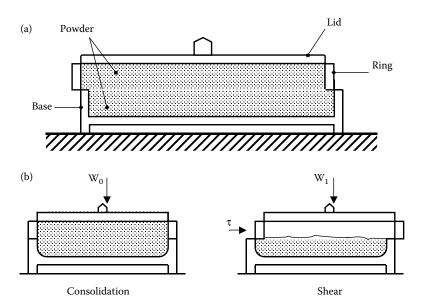


FIGURE 2.9 Diagram of Jenike's shear cell.

measure shear stress at failure. The preconsolidation consists in filling the assembly of base, ring and a packing mold ring over it, with a sample powder. A twisting top is placed on the sample and a force is applied to the top while giving it a number of oscillating twists. The force is released, the twisting top removed, and the powder surface is carefully scraped level with the upper half of the cell. In the consolidation step a shear cover is placed over the powder sample and a selected normal force is applied on it. A shear force is then continuously applied until it reaches a steady-state value indicating plastic flow. The shear force is then interrupted and the stem retracted. The shearing step comprises replacing the normal force of consolidation by a smaller force and reapplying the shearing force until the stress/strain peaks and falls off indicating failure plane in the sample and a point in the yield locus. The procedure is repeated with five or six different vertical loads, progressively smaller, applied to a set of identical samples and the shear force needed to initiate flow is found in each case. The forces are divided by the cross-sectional area of the cell to give stresses and the shear stress is plotted against the normal stress. The resulting graph is a yield locus (Figure 2.10), and it is a line that gives the stress conditions needed to produce flow for the powder when compacted to a fixed bulk density.

If the material being tested is cohesive, the yield locus is not a straight line and does not pass through the origin. It can be shown that the graph when extrapolated downward cuts the horizontal axis normally. As shown in Figure 2.10, the intercept *T* is the tensile strength of the powder compacts

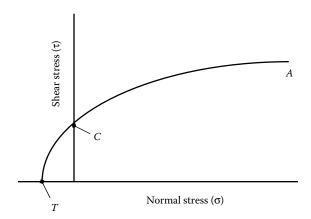


FIGURE 2.10 The Jenike yield locus.

tested and the intercept *C* is called the cohesion of the powder; the yield locus ends at the point A. The yield locus represents the results of a series of tests on samples which have the initial bulk density. More yield loci can be obtained by changing the sample preparation procedure and, in this way a family of yield loci can be obtained (Figure 2.11). This family of yield loci contains all the information needed to characterize the flowability of a particular material; it is not, however, in a convenient form. For many powders, yield locus curves can be described by the empirical Warren–Spring equation (Chasseray, 1994):



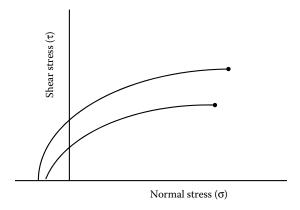


FIGURE 2.11
A family of yield loci.

where τ is the shear stress, C the material's cohesion, σ the normal stress, T the tensile stress, and n the shear index (1 < n < 2).

Apart from the cohesion and the tensile stress, the effective angle of internal friction can also be determined using the yield locus derived from testing a particular powder in a shear cell. The effective angle of internal friction is the resulting angle from drawing a tangent line to the Mohr circle inscribed in the yield locus, and passing through the origin (as shown in Figure 2.12).

The angle of wall friction can be measured by replacing the base of a Jenike shear cell by a plate of the material of which the hopper (or any sort of container) is to be made. The ring from the shear cell is placed on the plate and filled with powder and the lid is put in position. The shear force needed to maintain uniform displacement of the ring is found for different vertical loads on the lid. The slope of the graph of shear force against normal force gives the angle of friction between the particles and the wall, or angle of wall friction. This measure would complete the testing of a particulate material using only a Jenike shear cell.

The Jenike shear cell has been considered for long time the testing cell for establishing standard procedures in industrial applications and research. It has been recognized as one of the standards for testing bulk solids in the United States and in Europe, being especially focused on cohesive powders. The complexity of this method is such that errors due to poor technique can easily arise. A reference material has therefore been produced with which laboratories can verify both their equipment and experimental technique. The reference material consists of 3 kg of limestone powder packed in a polyethylene jar. It is accompanied by a certificate giving shear stress as a function of normal applied stress for four different powder compaction stresses.

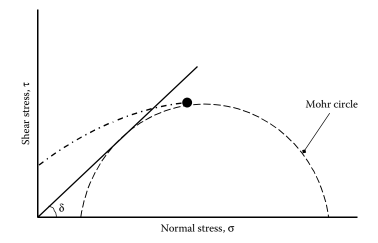


FIGURE 2.12Graphical determination of the effective angle of internal friction.

Although there are additional means to characterize solids flow properties, the Jenike method has been scrutinized and perfected over the years and remains the preferred approach. It is classified as a direct shear tester that is capable of providing information on a solid's "cohesive strength" as well as its "wall friction" properties. One of the major problems using the Jenike shear cell is that during operation, the shear force concentrates at the front of the shear cell. The shear force is nonuniformly applied to the sample. Similarly, the vertical force is also applied nonuniformly. At best, the results represent average stress conditions typically varying from a near-zero stress to the maximum applied (about two times larger than the average).

The annular shear cell was developed by Carr and Walker (1968) and has undergone a number of modifications. In this type of cell the shear stress is applied by rotating the top portion of an annular shear, as represented in Figure 2.13. These devices allow much larger shear distances to be covered both in sample preparation and testing, allowing a study of flow properties after failure. The original annular shear cells tended to give lower values for yield strength than the Jenike shear cell tester. The latest modifications to the ring cell tester have been made by Schulze (2007). In the Schulze ring shear tester the sample is placed in an outer circular channel. An angular lid attached to a crossbeam lies on top of the sample. Small bars are attached to the bottom side of the lid and the bottom of the cell to prevent the powder from sliding against the lid or the bottom. The movement of the cell with respect to the fixed lid causes the powder sample to shear. Load cells attached to tied rods measure the force needed to initiate this. To exert load on the sample, weights are hung from a crossbeam. This can be done during the shearing and consolidation of the powder sample. The cell can also be removed and time consolidation can be performed by placing weights on the sample outside of the test device. An automatic version of this measuring cell has been developed in

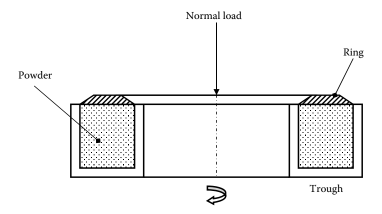


FIGURE 2.13 Diagram of an annular shear cell.

which a computer can automatically add loading to the sample and condition it. The instrument can be operated in a manual, semiautomatic, or totally automatic mode. Both, the original shear cells or the latest modified ones offer several important advantages over the Jenike-type shear cell. The area of shear is constant while the handling is easier, because consolidation and shear are quicker. After the sample is consolidated, a full locus can be generated without need of reconsolidating after each load. The consolidation process becomes more automated and uniform, eliminating much of the operator variability in the measurement process, characteristic of the Jenike-type tester.

Rotational shear cells, or powder rheometers, are relatively new testing instruments that can sensitively measure how flowability changes under a wide range of processing conditions, including various speeds, levels of entrapped air, and degrees of attrition. Powder rheometers also can condition the sample before testing, reducing the variability produced by differences in storage and handling. These instruments automatically rotate a blade through a cylindrical column of powder measuring the energy or force it needs to move through the sample and relating these measurements to various characteristics of the powder. The powder is placed in a circular vessel with a closed bottom. The blade is introduced into the powder and moves downward or upward in a helical motion, while the force on the blade shaft is recorded. The helical path along which the blade moves is dependant on the axial and rotational speeds, as well as in the direction of the blade rotation. The angle of approach the blade makes with the powder can be varied, so that the direction and angle of the measurement will allow one to measure compaction, shear, and slicing of the powder sample within the vessel. The torque data are treated to determine the largest theoretical torque exerted by the whole powder column on the rotor blade for any defined test condition. Therefore, powder shear can be measured under different shear conditions, including various downward compaction modes and upward expansion modes. The blade is interchangeable and different designs are used for different needs, including a rotating shear cell that can be used in a shallow column of powder. An arrangement as this is shown in Figure 2.14, and is called an open shear cell. As can be observed, the operations of this cell would be analogous to a rotational viscometer used in characterizing highly consistent non-Newtonian fluids. In congruence with this analogy, the open shear cell will measure both the force and the torque, and so these variables could be related to the shear stress and the normal stress to obtain the vield locus of a powder. Satisfactory results using rotational shear cells have been reported in different applications and with diverse industrial powders (Freeman and Cook, 2006; Freeman et al., 2009). Powder rheometers can provide simple and sensitive testing as they are relatively operator independent and vield quantitative values.

The angle of internal friction can be measured directly by the "grooved plate" method. The base of the Jenike shear cell is replaced by a metal plate in which a number of saw-toothed grooves are cut (Figure 2.15). These

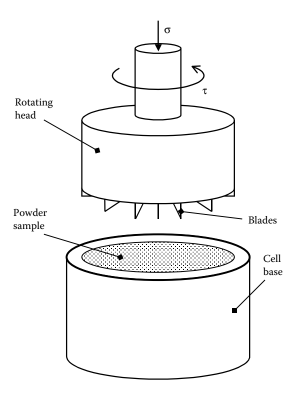


FIGURE 2.14 Diagram of rotating shear cell.

grooves are filled with the powder to be tested. The ring from the Jenike cell is then placed on the plate and filled with the powder and the lid placed into position. A load is placed on the lid and the ring is pushed across the grooves until the shear force settles out at a constant value which is measured, and this action is repeated for a number of different vertical loads. The graph of

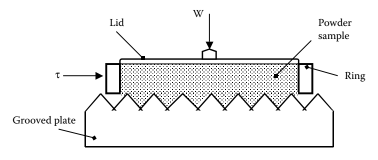


FIGURE 2.15The grooved plate method for measuring the angle of internal friction.

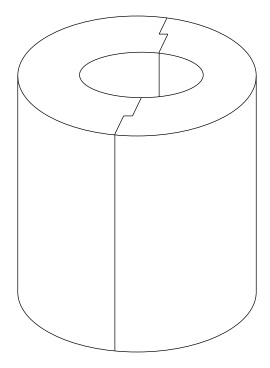


FIGURE 2.16Device for direct measurement of failure function.

shear force against normal force will be a straight line with its slope being the angle of internal friction of the powder.

For direct measurement of the failure function a split cylindrical die as shown in Figure 2.16 is used. The bore of the cylinder may conveniently be about 50 mm and its height should be just more than twice the bore. The cylinder is clamped so that the two halves cannot separate and it is filled with the powder to be tested, which is then scraped off level with the top face. By means of a plunger the specimen is subjected to a known consolidating stress. The plunger is then removed and the two halves of the split die are separated, leaving a free standing cylinder of the compacted powder. A plate is then placed on top of the specimen and an increasing vertical load is applied to it until the column collapses. The stress at which this occurs is the unconfined yield stress, that is, the stress that has to be applied to the free vertical surface on the column to cause failure. If this is repeated for a number of different compacting loads and the unconfined yield stress is plotted against the compacting stress, the failure function of the powder will be obtained.

A graph of a failure function of a powder is given in Figure 2.17. Although the results of this method can be used for monitoring or for comparison,

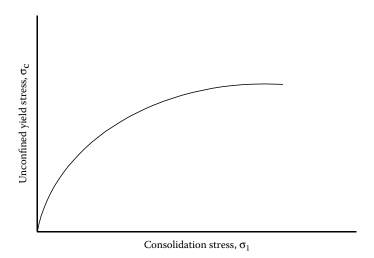


FIGURE 2.17 Failure function of a powder.

the failure function obtained will not be the same as that given by shear cell tests, due to the effect of die wall friction when forming the compact. A method of correcting for friction has been described by Williams et al. (1971). They proposed to compact the powder in a series of short increments and extrapolate results to estimate the unconfined yield stress for an infinite number of increments, that is, for a uniformly consolidated powder. Figure 2.18 shows a schematic diagram of the apparatus devised for the corrected method described. The cylinder is sectioned, not only into two halves, but also into 13 horizontal segments. The lower section at the base is 25.4 mm high, while the remaining 12 are half as high as this (12.7 mm high). The effect of the relationship length–diameter, used to adjust the values of the indirect method, is best evaluated using a single section at a time, that is, repeating the method of the complete cylinder section by section.

Two methods can be used for direct measurement of tensile strength. In the first, a mould of the same diameter as the Jenike cell is split across its diameter. The base of the cell is roughened by sticking sandpaper to it. The two halves are clamped together, the cell is filled with the powder, and then a lid is placed in position. The specimen is compacted by the application of a known vertical force to the lid and this, along with the clamp, is removed. The two halves of the cell, containing the specimen, rest on a base plate in which slots have been cut to form an air bearing. Air is introduced so that the cell can move horizontally without friction and the force needed to pull the two halves of the specimen are determined (Hong and Watanabe, 2000). Knowing the cross-sectional area of the specimen the tensile strength can be found. Measurements are made for a number of compacting loads and

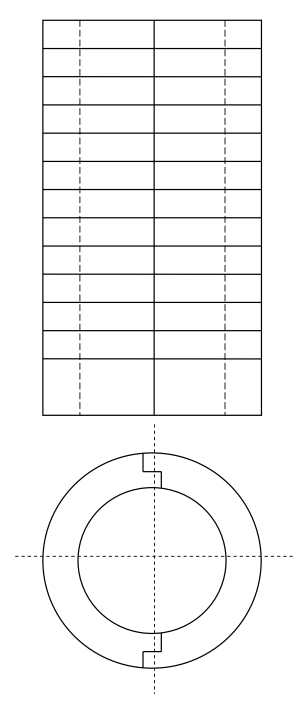


FIGURE 2.18 Scheme of the device for measuring a corrected failure function.

tensile strength is plotted against compacting stress. This method is quite difficult to perform properly, requiring careful attention to details. The second method is easier to use and gives results with less scatter. In this case a mould of the same diameter as the Jenike shear cell, and a lid which just fits inside it, are used. The base of the cell and the lower face of the lid are covered with sticking tape on which glue is spread. The cell is filled with the powder which is scraped off level with the top of the cell and the lid is placed in position. A compacting load is applied to the lid by means of a weight hanger and left in position until the glue has hardened. The lid is then attached through a tensile load cell to an electric motor, by which the lid is slowly lifted. The stress required to break the specimen is thus obtained. After failure, the lid and the base of the cell are examined and the result is accepted only if both are completely covered with powder, showing that tensile failure has occurred within the powder specimen and not at the surface. Figure 2.19 presents a diagram of these two methods for direct measuring of tensile strength.

The two methods described above for measuring the tensile strength of a lightly compacted powder do not give the same results, providing evidence that a bed of a particulate material will generally not be isotropic (i.e., its properties, including tensile strength, will be different in different directions). Since the second method is easier to use, while the compacting stress

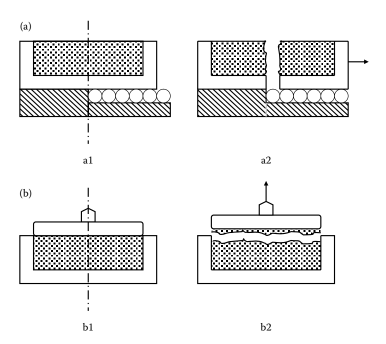


FIGURE 2.19 Methods for measuring tensile strength: (a) horizontal split cell, (b) lifting lid cell.

and the tensile strength are applied in the same direction, it is the mostly recommended for general characterization purposes.

2.4.4 Other Important Properties

As discussed previously, since most fine powders of importance in process engineering are cohesive, the use of characterization variables that do not take into account the state of consolidation have to be used with extreme care. However, there may be particulate systems of interest in different processing industries, which are free flowing, such as cereal grains, coarse sand, gravel, etc. For these types of free-flowing materials some characteristics angles can be used for calculations concerned with aspects of storage and transport. Some of these angles are the angle of repose, the angle of slide, the conveying angle, and the angle of spatula.

The angle of repose is defined as the angle of the free surface of a pile of powder to the horizontal plane. Depending on the conditions under which the pile has been poured and how the angle is measured, different values of the angle can be obtained for the same powder. Several types of angle of repose are used to assess powder flowability in an empirical manner; the most commonly employed are the poured and the drained angles of repose. The former is measured on a pile poured freely onto a flat surface while the latter is measured on the conical surface of powder in a flat-bottomed container when the powder has been allowed to flow out through an orifice in the base. A standard method for measuring the poured angle of repose has been reported for alumina (British Standards Institution, 1970). As shown in Figure 2.20, a funnel is cut short and adapted into a bridge. The lower opening of the funnel is located 52 mm from a metal plate on which concentric circles have been engraved. The center of the smallest circle is aligned with the center of the funnel. The powder, which is noncohesive, is discharged through the funnel until the poured pile just touches the lower opening of

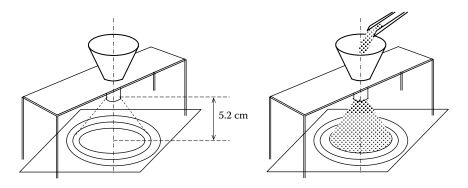


FIGURE 2.20Apparatus for measuring the poured angle of repose of a free-flowing powder.

the funnel. The concentric circles help in measuring easily the diameter of the poured pile and, since the height is fixed, by simple trigonometry the angle of repose is determined. Although, as mentioned, the method was developed for alumina, there is no reason why it may be applied for any other noncohesive, easily flowing, or free-flowing particulate system.

The drained angle of repose is always greater than the poured angle of repose, because in the first case the powder sliding or rolling down the slope is separating, while in the second it is converging. Although, as has been already discussed, the angle of repose could be considered meaningless in many practical applications, it can be used as a rough flowability indicator (Geldart et al., 1990). According to Carr (1976) angles of up to 35° indicate free flowability, angles of 35–45° indicate some cohesiveness, angles of 45–55° indicate cohesiveness (loss of free flowability), and angles 55° and above indicate very high cohesiveness and, therefore, very limited (or none) flowability.

The angle of slide is closely related to the drained angle of repose (Wilkinson et al., 1983). It is the minimum angle to the horizontal of a flat, inclined, surface that will allow bulk solid to flow from rest under its own weight. This angle is supposed to be useful in designing stationary chutes but its measurement has not been described as standardized. The value measured is expected to be highly influenced by the material of construction of the chute, the amount of material on it, and by humidity. It has been shown (Augenstein and Hogg, 1978) that the motion of individual particles strongly depends on the nature of the surface over which they flow: highly roughened surfaces cause shear within the flowing stream while smooth ones cause slip at the surface.

The conveying angle, also called the maximum angle of rise, is the angle at which a smooth belt conveyor can operate without the bulk powder running back down. There is a need for a standard test method to be developed for determination of this angle, to allow designers of belt conveying systems to have a reliable design criterion. A rough guide given in the literature (Wilkinson et al., 1983) establishes that the angle of rise used in such systems should not exceed about half the poured angle of repose.

The angle of spatula is measured by spooning out powder with a 22 mm wide flat-blade spatula that is inserted into the bulk powder parallel to the bottom of the container (Carr, 1965). The spatula is lifted straight up and out of the powder, keeping it parallel to a horizontal plane. The spatula is vibrated or tapped in a specified manner and the angle of the tangent to the powder surface at the edge of the spatula is measured, to the horizontal plane. The average angle of several measures is taken and, except for very free-flowing materials, its value is always greater than that of the poured angle of repose, since it is taken from a consolidated material. The angle of spatula is considered useful as a very simple and rough estimate of flowability; a value under 40° indicates that the material is free flowing.

Example 2.2

Table 2.3 shows values of unconfined yield stress versus consolidating stress for three powders: A, B, and C. Determine their corresponding flow factors, and classify them according to Jenike's classification.

SOLUTION

In order to determine then flow factor ff of each powder, the failure function FF has to be determined first. Then, according to Jenike's classification criterion, a straight line from the origin intercepting the failure function curve at a fixed value of 22.4 N for σ_c , would give a slope. The inverse value of the slope is the flow factor. The numerical value of ff for each powder would give its type according to limits given in Table 2.1.

Plotting the tabulated data in Table 2.3, Figure 2.21 is obtained, where the straight line from the origin intercepting the curve of powder C at the fixed value of 22.4 N for σ_c is exemplified. The slope for this straight line is 1.44, and its inverse value, that is, 0.69 is its flow factor ff. Following a similar procedure from powders A and B, values of 1.2 and 2.5, are obtained, respectively for ff.

A summary of the categorization of the studied powders is

Powder	Flow Factor (ff)	Type of Powder (Jenike's Classification)	
A	1.20	Very cohesive	
В	2.50	Cohesive	
С	0.69	Very cohesive	

TABLE 2.3 Example 2.2

		$\sigma_{c}(N)$	
$\sigma_1(N)$	A	В	С
0	0	0	0
10	8	5	16
20	16	10	28
30	23	14	38
40	29	17	48
50	34	20	57
60	38	22	64
70	42	24	70
80	44	26	72

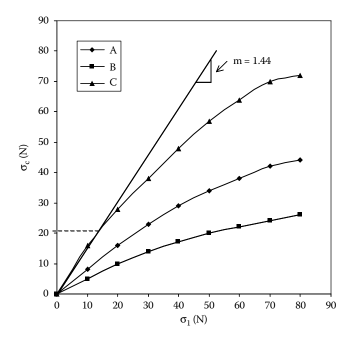


FIGURE 2.21 Graph for Example 2.2.

Example 2.3

A powder is characterized running an experiment in a Jenike's shear cell. The results are given in Table 2.4. Determine the failure properties that can be derived from the yield locus of the powder.

SOLUTION

According to discussion presented in Sections 2.4.2 and 2.4.3, the failure properties that can be read off directly from a yield locus are the cohesion C and the ultimate tensile strength T (Figure 2.10). Also, by drawing a Mohr circle and following the procedure described to modify the yield locus as illustrated in Figure 2.12,

TABLE 2.4 Example 2.3

τ (kPa)	σ (kPa)		
0.75	0.4		
1.05	0.9		
1.30	1.4		
1.50	2.0		
1.55	2.4		

the effective angel of internal friction can be found. The yield locus for the powder described in Table 2.4 is drawn and presented in Figure 2.22.

As indicated in Figure 2.22, the failure properties for the powder are C = 0.48 kPa, T = 0.3 kPa, and $\delta = 42^{\circ}$.

2.5 Laboratory Exercise: Determination of Some Failure Properties of Powders

2.5.1 Introduction

The technique of shear testing has traditionally been used to determine the shear behavior of consolidated powders. As discussed previously, the Jenike shear cell testing is generally accepted as an established method for measuring powder flowability and standards based on it have been issued, both in Europe and the United States. Some disadvantages of the Jenike shear cell have also been mentioned. The process is extremely time consuming and tedious and correct consolidation of sample is difficult to achieve. It can be hardly used for all types of bulk solids due to its limited shear

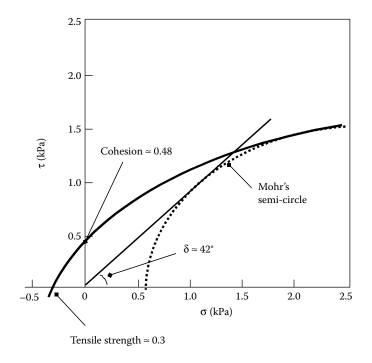


FIGURE 2.22 Graph for Example 2.3.

displacement, which is at most twice the thickness of the upper ring. Some bulk solids need a greater shear displacement to reach steady-state flow conditions. Perhaps the two critical practical disadvantages of the Jenike shear cell are its operator skillful dependence, and the extremely long time taken to perform a test under some specific conditions. Teunou and Fitzpatrick (2000) reported on an experimental stage of a project to take eight months to complete using one consolidating bench.

Alternatives to the Jenike shear cell are the annular shear cell and the rotational shear cell, also called the powder rheometer. It has been reported (Freeman and Cooke, 2006) that powder rheometers are reliable, fast, and give results comparable to standard methods based on the Jenike shear cell. The objective of the exercise is to compare the standard Jenike method for limestone CRM 116 powder with a method using a powder rheometer.

2.5.2 Instrument and Materials

A rotational shear cell instrument, such as the FT4® Powder Rheometer, equipped with a 48 mm rotational shear cell and a 30 mL shear measurement vessel. A batch of reference limestone powder (CRM 116), produced and sold by the Commission of European Communities.

2.5.3 Shear Tester Operation

Follow the directions given in the instrument manual for shear testing methodology. The routine procedure can be summarized as follows:

- Condition the sample using the test blade slicing downward through the powder followed by an upward traverse to lift the powder and drops it over the blade. This process helps to remove the effect of different sampling methodologies and powder storage times.
- Compact the sample by using a ventilated compaction piston (allowing entrained air to escape), with a force equal to that of the preshear normal stress.
- Shear the sample continuously until a steady-state flow is achieved, to complete the preshearing state.
- Reduced the normal stress, so that the sample is now over consolidated with respect to the normal stress now applied. Restart shearing and measure the point of incipient failure.
- For each preshear normal stress, take five measurements at the five normal stresses defined by the standard. Take also a measurement of shear stress at the preshear normal stress level, that is, at 3, 6, 9, and 15 kPa.
- The five measurements taken make up the yield loci for each preshear normal stress level.

2.5.4 Calculations and Report

- Obtain the yield loci by plotting shear stress versus normal stress at the different preshear normal stresses.
- Draw Mohr's circles on the graphs of the yield loci derived, to extrapolate the failure properties, such as the effective angle of internal friction. Determine the remaining failure properties that can be read from the graph.
- Perform a search to obtain the standard yield loci from Jenike's testing on CRM 116 limestone powder, in order to compare the results obtained with such standard.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out, as well as discussion on own data. Compare and discuss, as much as possible, agreements or discrepancies based on literature survey from studies and investigations on the subject.

2.6 Particle Size Analysis

2.6.1 Introduction

Particle size distribution measurement is a common method in any physical, mechanical, or chemical process because it is directly related to material behavior and/or physical properties of products. The bulk density, compressibility, and flowability of an industrial powder are highly dependent on particle size and its distribution. Segregation will take place in a freeflowing powder mixture because of the differences in particle sizes. There are many other properties of particulate systems that strongly depend on particle size, for example, activity of drugs, setting time of cement, and hiding power of pigments. Characterization of solid particles, which are in practice mostly irregular in shape, is usually done by analyzing particle size and its distribution. Other characteristic properties of the solid material may be included in the type of size measured; for example, Stoke's diameter combines size, density, and shape all in one parameter. Quantitative measure of particle shape can be obtained indirectly by analyzing two or more types of particle size and looking at different "shape coefficients" that relate to those sizes. There are many different types of instruments available for measuring particle size distribution but most of them would fall into four general methods: sieving, microscope counting techniques, sedimentation, and stream scanning.

2.6.2 Definitions of Characteristic Linear Dimension

In particle size measurement two most important decisions have to be made before a technique is to be selected for the analysis; these are concerned with the two variables measured, the type of particle size and the occurrence of such size. An irregular particle can be described by a number of sizes depending on dimension or property is measured. Particle size was previously discussed in Chapter 1 and definitions of equivalent sphere diameters, equivalent circle diameters, and statistical diameters have been provided (Tables 1.2, 1.3, and 1.4). Emphasizing what was already presented, is important to bear in mind that great care must be taken when making a selection of particle size, as an equivalent diameter, in order to choose the most relevant to the property or process which is to be controlled. For example, in solid-liquid separations where particle motion relative to the fluid is the governing mechanism for separation (gravity or centrifugal sedimentation, hydrocyclones) it is of course most relevant to use a method that measures the freefalling diameter or, more often, the Stoke's diameter. In filtration, on the other hand, it is the surface-volume diameter (measured, e.g., by permeametry), which is more relevant to the principle of separation in question.

The occurrence of amount of particle matter which belongs to specified size classes may be classified or arranged by diverse criteria as to obtain tables or graphs. In powder technology the use of graphs is convenient and customary for a number of reasons which will be mentioned and discussed as this topic is developed. For example, a particular size which is to be used as the main reference of a given material is easily read from a specific type of plot.

2.6.3 Types of Distributions

There are four different particle size distributions for a given particulate material, depending on the quantity measured: by number $f_N(x)$, by length $f_L(x)$, by surface $f_S(x)$, and by mass (or volume) $f_M(x)$. From all these, the second mentioned is not used in practice as the length of a particle by itself is not a complete definition of its dimensions. These distributions are related but conversions from one another is possible only in cases when the shape factor is constant, that is, when the particle shape is independent of the particle size. The following relationships show the basis of such conversions:

$$f_{\rm L}(x) = k_1 \cdot x \cdot f_{\rm N}(x) \tag{2.17}$$

$$f_{\rm S}(x) = k_2 \cdot x^2 \cdot f_{\rm N}(x)$$
 (2.18)

$$f_{\rm M}(x) = k_3 \cdot x^3 \cdot f_{\rm N}(x)$$
 (2.19)

Bulk Solids 69

where constants k_1 , k_2 , and k_3 contain a shape factor that may often be particle size dependent making an accurate conversion impossible without the full quantitative knowledge of its dependence on particle size. If the size of the particles does not vary with size, constants above can be easily found because, by definition of distribution frequency:

$$\int_{0}^{\infty} f(x) \, \mathrm{d}x = 1 \tag{2.20}$$

so the areas under the curve should be equal to 1.

The conversions mentioned above are to be avoided whenever possible, because of inherent errors in such procedures, by using a method that would give the desired type of distribution directly. Different methods give different types of distributions and the selection of a method should be done on the basis of both the particle size and the type of distribution required. In common processing applications, for example, solid–fluid separations, it is the size distribution by mass that is usually of interest, because the separation efficiency is based on gravimetric means. There are, however, cases such as liquid clarification where turbidity of the overflow is of importance, and size distribution by surface or even by number would be more relevant. Figure 2.23 shows the four types of distribution in a diagram.

2.6.4 Measures of Central Tendency

There is a great number of different average or mean sizes which can be defined for a given particle size distribution. The purpose of such measures

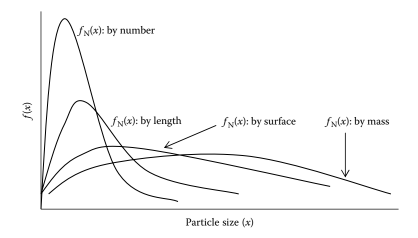


FIGURE 2.23 Four particle size distributions of a given particle population.

of central tendency is to represent a population of particles by a single figure. Although such representation will give no indication of the width of the distribution, it may sometimes provide a useful guide for process control. There are three most important measures of central tendency for a given size distribution. These are, as shown in Figure 2.24, the mode, the median, and the mean. The mode is the most commonly occurring size, that is, the size corresponding to the peak on the size distribution frequency curve. Some distributions may have more than one peak and those are commonly referred as multimodal. The median, or the 50% size, is the size at which half the particles are larger and half are smaller, that is, the size which splits the area under the distribution frequency curve into two halves. The median is most easily determined from the cumulative percentage curves, where it corresponds to 50%. There are many mean diameters that can be defined for a given particle size distribution with a definition of the general form:

$$g(\overline{x}) = \int_{0}^{\infty} g(x)f(x)dx$$
 (2.21)

where f(x) is the particle size distribution either by number, length, area, or mass, whichever may be of interest and g(x) is a certain function of particle

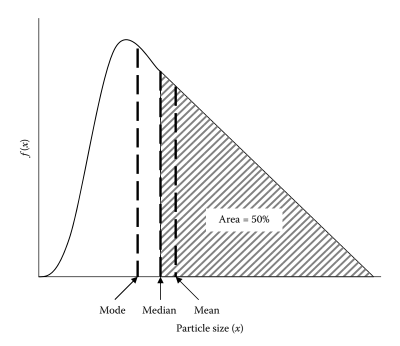


FIGURE 2.24 The mode, median, and mean of a particle size distribution.

Mean Diameters as a Function of $g(x)$			
Form of g(x)	Name of Mean Diameter, \bar{x}		
g(x) = x	Arithmetic mean, \overline{x}_a		
$g(x) = x^2$	Quadratic mean, \bar{x}_q		
$g(x) = x^3$	Cubic mean, \bar{x}_c		
$g(x) = \log x$	Geometric mean, \bar{x}_g		
g(x) = 1/x	Harmonic mean, \overline{x}_h		

TABLE 2.5 Mean Diameters as a Function of g(x)

size x, and depending on the form of this function there are different mean diameters \bar{x} as listed in Table 2.5.

Evaluation of the various means required for a given particle size distribution is based on Equation 2.21, which may also be written as

$$g(\overline{x}) = \int_{0}^{1} g(x) dF$$
 (2.22)

because

$$f(x) = \frac{\mathrm{d}F}{\mathrm{d}x} \tag{2.23}$$

If either f(x) or F(x) are available as analytical functions, the desired mean diameters are evaluated by integration following Equation 2.21. If, however, no analytical function is fitted and the particle size distribution is in a form of a graph or a table, evaluation of mean diameters can best be done graphically. Most particle size measurement techniques result in cumulative percentage F(x) and, thus, according to Equation 2.23 if F(x) is plotted against g(x) for a number of corresponding sizes, g(x) is then represented by the area under the curve with respect to the F(x) axis, as illustrated in Figure 2.25. The mean is evaluated from this area using the corresponding relation for g(x), as given in Table 2.5, being immaterial whether F(x) is plotted as percentage oversize or undersize. It is always best to represent a population of particles by the actual size distribution curve and only in cases when this is not possible or feasible, resort to using a single number, a measure of central tendency, for characterizing a particulate system. In such cases, care must be taken to select the type of mean size most relevant to the given application.

2.6.5 Presentation of Data

The common way of presenting particle size data of different powders is in form of a plot. Normally, the particle size axis is the "x" or horizontal and

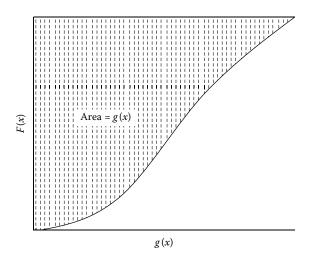


FIGURE 2.25 Evaluation of a mean \bar{x}_a from the cumulative percentage F(x).

the particle amount axis is the "y" or vertical. What is plotted on the particle size axis is a matter of which quantity used to represent size of individual particles suits better to a specific problem. The amount of particle matter, which belongs to specified size classes on the particle axis, may also be represented in several ways. Number of particles and mass of particles are the most common ones, but surface area and volume are used as well. In general, particle size distributions can be presented as frequencies f(x) or cumulative frequencies F(x), which are related to each other by the following equation:

$$f(x) = \frac{\mathrm{d}F(x)}{\mathrm{d}x} \tag{2.24}$$

The graphical representation of a particle size distribution is usually plotted in a cumulative form. In a typical cumulative plot, points are entered showing the amount of particulate material contributed by particles below or above a specified size. Hence, the curve presents a continuously rising or decreasing character. These oversize and undersize distributions, as illustrated in Figure 2.26, are simply related by

$$F(x)_{\text{oversize}} = 1 - F(x)_{\text{undersize}}$$
 (2.25)

A cumulative plot will, therefore, include a broad range of particle sizes. It is often convenient, however, to refer to a single characteristic size for the system. Many characteristic sizes have been proposed, most of them

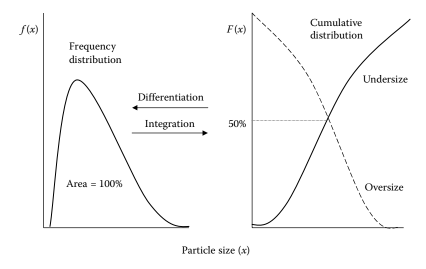


FIGURE 2.26 Relationship between frequency and cumulative distributions.

involving a mathematical formula. An important one, which can be read off any cumulative plot of the particle size data, is the median particle size. It is defined as that particle size for which the particle amount equals 50% of the total. If the particle amount is represented by number, such point is called the number median size. If mass is used as the measure of particle amount, this variable is known as the mass median size. The distinction between number and mass median is very important, since they differ generally by a considerable amount. Such difference means that number and mass cumulative plots do not agree for the same system of particles. The weight of a particle, which varies as the cube of its diameter, account for this mentioned disagreement. For practical purposes, it is reasonable to fit an analytical function to experimental particle size distribution data, and then handle this function mathematically in further treatment. It is, for example, very much easier to evaluate mean sizes from analytical functions than from experimental data. Several different distribution functions can be found in the literature. All of them should be treated as empirical equations as they very rarely have any theoretical relation to the process in which the particles were produced.

There are many different types of size distribution functions, either in cumulative or relative percentage frequency form. These include the normal distribution, log-normal distribution, and Rosin–Rammler functions (Allen, 1981); the Gates–Gaudin–Schuhmann, Bennett's form, Gaudin–Meloy, and modified Gaudin–Meloy functions (Bergstrom, 1966; Harris, 1968); the Roller and Svenson functions (Beke, 1964); the error function (Herdan, 1960); the modified beta function (Peleg et al., 1986; Peleg and Normand, 1986); and the Griffith and Johnson's S_B function (Yu and Standish, 1990).

The Gates-Gaudin-Schuhmann function is expressed as (Harris, 1968)

$$Y = \left[\frac{x}{k}\right]^m \tag{2.26}$$

where Y is the cumulative weight fraction under size x, k the characteristic size of the distribution, and m the measurement of the distribution spread, also called the Schuhmann slope.

The Rosin–Rammler function is widely used in particle size distribution characterization. It was introduced in comminution studies in 1933 (Harris, 1971/1972), but was also used to describe the particle size distribution of moon dust (Allen, 1981). Usually, it is a two-parameter function given as a cumulative percentage undersize (Svarovsky, 2000):

$$Y = 1 - \exp\left[-\left(\frac{x}{x_{\rm R}}\right)^n\right] \tag{2.27}$$

where Y is the weight fraction of material finer than size x, x_R is a constant giving a measure of the present particle size range, and n another constant characteristic of the analyzed material that gives a measure of the steepness of the cumulative curve. The x_R can easily be found from the plot in the Rosin–Rammler graph because it is the size corresponding to 100/e = 36.8%, and n is the slope of the line (Svarovsky, 2000). Lower values of n are associated with a more scattered distribution, while higher values of n will imply an increasingly uniform particle structure.

The modified Gaudin–Meloy function is expressed as (Bergstrom, 1966; Harris, 1968)

$$y = \left[1 - \left(1 - \frac{x}{x_0}\right)^r\right]^m \tag{2.28}$$

where Y is the cumulative weight fraction under size x, x_0 the parameter related to the maximum particle size, m is called the Schuhmann slope, and r is the ratio of x_0 to the size modulus.

Materials with a normal distribution of particle size are relatively rare and are found chiefly among the particulate produced by chemical processes like condensation or precipitation. The importance of this function, however, is that provides an idealized error distribution built upon the assumption that elementary errors or small causes combine at random to produce the observed effect (Herdan, 1960). The normal distribution function is given by

$$y = \frac{1}{\sigma\sqrt{2\pi}} \exp\left[-\frac{(x-a)^2}{2\sigma^2}\right]$$
 (2.29)

Bulk Solids 75

where y is the probability density, x the diameter of the particle, a the arithmetic mean, and σ the standard deviation. From the normal function, if the arithmetic mean a is zero, the probability of occurrence within the interval form the mean (zero) to the value x, that is, F(x), is given by the error function (Herdan, 1960):

$$F(x) = \frac{1}{\sigma\sqrt{2\pi}} \int_{0}^{x} \exp\left[-\frac{x^2}{2\sigma^2}\right] dx$$
 (2.30)

where σ and x have the same meaning as described in the normal function.

As descried by Herdan (1960), if a powder is obtained by comminution processes, such as milling, grinding, or crushing, its distribution appears to be governed very often by the log-normal law. Furthermore, the log-normal function is the most useful one among the different types of functions (Beddow and Meloy, 1980). It can be given in this form:

$$f(x) = \frac{1}{x \ln \sigma_g \sqrt{2\pi}} \exp\left[-\frac{(\ln x - \ln x_g)^2}{2 \ln^2 \sigma_g}\right]$$
(2.31)

where f(x) is the size distribution function for particle size x, x_g is the geometric mean of the distribution, and σ_g the geometric standard deviation of $\ln x$ (Peleg et al., 1986).

For many processes in which the population mode and spread vary independently and the size distributions have a finite range, the modified beta distribution function is more appropriate than the log-normal function because it has a finite range and the ability to describe symmetric as well as asymmetric distributions skewed to the right or left (Peleg and Normand, 1986). The modified beta distribution $f_{am}(x)$ is defined as

$$f_{am}(x) = \frac{x^{am}(1-x)^m}{\int_0^1 x^{am}(1-x)^m dx}$$
(2.32)

where *a* and *m* are constants, and *x* is the normalized length given by

$$x = \frac{x - x_{\min}}{x_{\max} - x_{\min}} \tag{2.33}$$

where x_{\min} and x_{\max} are the smallest and largest particle sizes, respectively, and therefore:

$$x_{\min} < x < x_{\max}$$
, $0 < x < 1$

2.6.6 Laboratory Techniques for Size Measurement

There is an abundance of methods available for measurement of particle size distributions and several textbooks, such as some referred in this chapter (Allen, 1981; Kaye, 1981), are available and review the field in great depth. Table 2.6 gives a schematic review of the methods available, size ranges covered and types of particle size and size distribution measured. This information is given in order to assist an engineer faced with the problem of selection of the best method for a given application. Only a preliminary selection can be attempted using Table 2.6, because it is impossible to list all the important factors influencing the choice, such as type of equivalent diameter required, quantity to be measured, size range, quantity of sample available, degree of automation required, and so on. It should be noted that many of these factors are interrelated and their relative importance varies in different applications. As mentioned in Section 2.6.1, most of the analytical techniques currently available for particle size measurement can be categorized into four main methods, namely, sieving, microscope counting techniques, sedimentation, and stream scanning. The most representative techniques following this classification will be reviewed in this section.

TABLE 2.6Analytical Techniques for Particle Size Measurement

	A	T (D: -1-	T (C:
Technique	Approximate Size Range (µm)	Size	Type of Size Distribution
Sieving			
Woven wire	37-4 000	$x_{\rm A}$	By mass
Electro formed	5–120	x_{A}	By mass
Microscopy			
Optical	0.8-150	$x_{\rm a}, x_{\rm F}, x_{\rm M}$	By number
Electron	0.001-5	$x_{\rm SH}, x_{\rm CH}$	
Gravity sedimentation			
Incremental	2-100	$x_{\rm st}, x_{\rm f}$	By mass
Cumulative	2-100	$x_{\rm st}, x_{\rm f}$	By mass
Centrifugal sedimentation			
Two layer-incremental	0.01-10	$x_{\rm st}, x_{\rm f}$	By mass
Cumulative			
Homogeneous-incremental			
Flow classification		$x_{\rm st}, x_{\rm f}$	
Gravity elutriation (dry)	5-100	$x_{\rm st}, x_{\rm f}$	By mass
Centrifugal elutriation (dry)	2–50	$x_{\rm st}, x_{\rm f}$	By mass
Impact separation (dry)	0.3-50	$x_{\rm st}, x_{\rm f}$	By mass or number
Cyclonic separation (wet or dry)	5-50	$x_{\rm st}, x_{\rm f}$	By mass
Particle counters			
Coulter principle (wet)	0.8-200	$x_{ m v}$	By number

2.6.6.1 Microscopy

Microscopy counting is the most direct method of particle size evaluation as the particles are actually counted using an optical microscope. Optical microscopy is used to perform such as task while electron microscopy is more often used to characterize shape and morphology of particles due to its range and resolution. Counting particles by the use of a microscope is quite tedious so the practical limit only reaches the value of 50 μ m, although it may be used for limit sizes of the order of 150 μ m (Table 2.6).

For microscope counting a sample is prepared by suspending particles in a liquid medium (normally with added dispersant) and put them on slide for examination. The recommended suspension concentration is, like in some other techniques, of about 0.1% by volume in order to avoid overlapping of particles. Once the slides have been prepared it is necessary to decide whether or not the particles should rest in their most stable position under gravity or should be held in the orientation in which they fell by coating the slides with some sticky substance. The counting is the done bearing in mind to important aspects, namely, first, a statistical diameter, such as the Martin's or Feret's diameter (Table 1.4), should be measured keeping the same direction for all counts in every slide and, second, at least six particles in every one of a hundred slides ought to be counted. Because of the nature of the technique, the data are normally collected in tabular form, but final graphical representation is recommended.

2.6.6.2 Screening

Screening, or sieving, is known as one of the most useful, simple, reproducible, and inexpensive methods of particle size analysis, and belongs to the techniques using the principle of geometry similarity. It is considered the only method giving a particle size distribution based on the mass of particles in each size range. The particle size is defined by the sieve aperture by which a particle may, or may not pass through. As presented in Table 2.6, all types of sieving cover a range from 5 μ m to 4 mm. This lower limit can be achieved using micromesh sieves, while the upper limit can be extended to the centimeter range by punched-plate sieves. The minimum applicable particle size range is limited by two main reasons: first, it is not possible to produce sieve cloth fine enough for it and, second, very small powders do not have strong enough gravity force to resist its tendency to adhere to one another and to the sieve cloth (Allen, 1981; Herdan, 1960).

A standard sieve series usually consists of a set of sieves with apertures covering a wide range from microns to centimeters. The sieve size is defined as the minimum square aperture through which the particles can pass. Sieves are often referred to by their mesh size, that is, the number of wires per linear inch. Mesh size and the wire diameter determine the aperture size. The ratio of aperture of a given sieve to the aperture of the next one in a

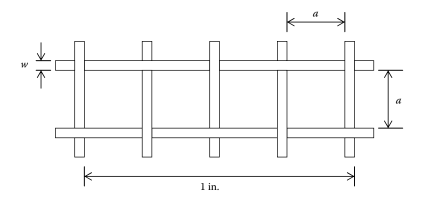


FIGURE 2.27 Dimensions of sieve on a screen.

sieve series is a constant. Mesh size M, screen aperture a, and wire diameter w, are related by the following equations (Figure 2.27):

$$M = \frac{1}{a + w} \tag{2.34}$$

$$a = \frac{1}{M} - w \tag{2.35}$$

Equations 2.34 and 2.35 allow for transformations between mesh number and aperture size. Both features of sieves are listed in standard screen scales. Standardized sieve apertures were first proposed by Rittinger (1867). Modern standards are based on either a $\sqrt{2}$ or $\sqrt[4]{2}$ progression. In the United States, the series of sieves with standard opening sizes are called "Tyler" sieves and the openings of successive sieves are based on a $\sqrt{2}$ progression starting at 45 µm.

Sieving analysis consists on stacking the sieves in an ascending order of aperture size, placing the material concerned on the top sieve, vibrating them by machine or hand for a fixed time, and determining the weight fraction retained on each sieve. Wet sieving method is useful for very fine powders or when the material is originally suspended in a liquid. It represents an excellent alternative for powders forming aggregates when dry sieving is used. There are two main forms in which the results of a sieve test can be presented: tabular and graphical. As previously discussed, graphical methods are preferred in particle size analysis as they provide with a simple way of identifying a representative size of the powder being analyzed. In sieving, like in all size measurement techniques, cumulative percentages of oversize or undersize material against particle size is plotted to obtain graphs of useful information for powder characterization. The convention would

command to use the sieve diameter x_A , as defined in Table 1.2, to be the particle size plotted on cumulative percentage graphs, but it is still customary to use mesh number instead. Sometimes the mesh number is plotted progressively, and so the normally obtained oversize graph, would present the typical rising character of an undersize graph, since mesh number gets larger as the particle size actually gets smaller.

Example 2.4

A sample of 200 g of an industrial powder is screened through a series of sieves and the results of the run are presented in Table 2.7. Elaborate a histogram, a frequency curve, and an oversize cumulative percentage graph, and determine the median size of the powder. Identify clearly such median size in the corresponding graph.

SOLUTION

Consulting the British Standards Screen Scale in Appendix 5, the nominal aperture sizes in micrometers are tabulated versus the weight retained. Then, the fractions retained on each of the sieves used in the test can be listed as percentage of the original test sample weight, and finally as the oversize cumulative percentages (i.e., running totals). The results of these calculations can be tabulated on a second table (Table 2.8).

The histogram can then be plotted as the limits of the size aperture versus the weight retained on each of the sieves. Plotting these two values, would give a histogram, as shown in Figure 2.28.

The vertical columns in the histogram extend between the various adjacent sieves used in the test. The points in the histogram may be plotted in between two sieve sizes. For example, for the peak fraction which passed through an 850 μ m sieve and was retained on a 600 μ m sieve, the mean particle size could be taken as the average of 850 and 600 = 725 μ m. By doing this procedure, the frequency curve in Figure 2.29 can be obtained.

TABLE 2.7 Example 2.4

Mesh Number (BS)	Weight Retained (g)		
10	2.2		
14	11.2		
18	41.2		
25	52.8		
36	37.0		
52	21.6		
72	11.8		
100	7.8		
150	5.4		
-150	9.0		

TABLE 2.8	
Example 2	.4

Mesh Number (BS)	Screen Aperture, x_A (μ m)	Weight Retained (g)	Cumulative Oversize Weight (g)	Cumulative Oversize Percentage
10	1700	2.2	2.2	1.1
14	1180	11.2	13.4	6.7
18	850	41.2	54.6	27.3
25	600	52.8	107.4	53.7
36	425	37.0	144.4	72.2
52	300	21.6	166.0	83.0
72	212	11.8	177.8	89.0
100	150	7.8	185.6	93.0
150	106	5.4	191.0	96.0
-150 (170)	90	9.0	200.0	100.0

Finally, the oversize cumulative percentage (last column in Table 2.8) versus the sieve aperture size would give the cumulative percentage graph (oversize) as shown in Figure 2.30.

A cumulative undersize graph would be obtained by subtracting from 100 each of the values of the cumulative oversize percentage column. The point at the 50% value of any of the cumulative percentage graphs is the median size. In the oversize cumulative percentage graph shown in Figure 2.30, the median size is marked as, approximately, $625 \, \mu m$.

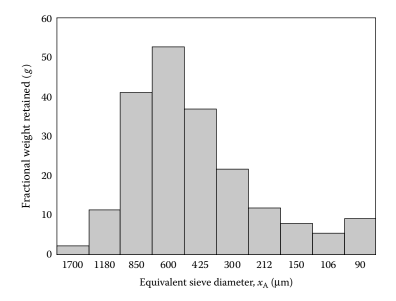


FIGURE 2.28 Graph for Example 2.4.

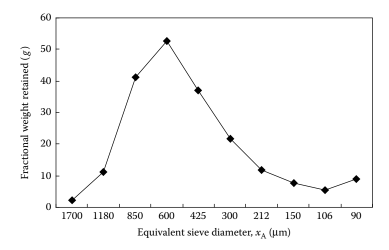


FIGURE 2.29 Graph for Example 2.4.

2.6.6.3 Sedimentation

The methods based on sedimentation are probably the most numerous and varied of all particle sizing techniques. They measure the Stokes' diameter of the particle and, therefore, measure the variable which is often most required to describe the behavior of the particles in suspension. There are some limitations with respect to particle shape in sedimentation analysis as particles that diverge grossly from spheres would not fall in a vertical pattern and, hence, the results when compared with other methods will tend to show

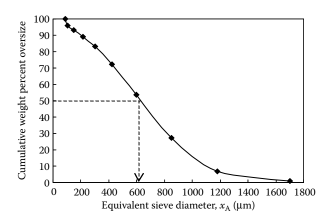


FIGURE 2.30 Graph for Example 2.4.

oversize material. The two most common methods of particle size measurement by sedimentation are gravimetric and centrifugal sedimentation.

Gravity sedimentation is normally carried out by using the "Andreasen" pipette technique. In its simplest form, a suspension of approximately 0.1% by volume of particles is placed in a measuring cylinder, and a pipette used to withdraw 10 mL from a fixed depth below the surface. Such an apparatus is shown in Figure 2.31. The concentration of samples are measured and compared with the initial one. Since all the particles will settle at their terminal velocities, the ratio of the two concentrations is the fractional weight of particles to have reached a depth of H cm. Excluded will be the particles with a diameter smaller than x_{st} (Table 1.2) as given in the relationship:

$$x_{\rm st} = \sqrt{\frac{18\mu H}{(\rho_{\rm s} - \rho)gt}} \tag{2.36}$$

where μ is the liquid viscosity, ρ_s is the particle density, ρ is the liquid density, g is the acceleration due to gravity, and t is the time.

From the fractional weight evaluated by Equation 2.36 the fraction undersize of a given powder can be determined. Allen (1981) has shown that variations in concentration within a settling suspension can be related to cumulative percentage undersize F(x) by the equation

$$F(x) = \frac{C(H,t)}{C(H,0)}$$
 (2.37)

where C(H, t) is the concentration in weight by volume at a depth H and time t, and C(H, 0) is the concentration in weight by volume at depth H and time zero. The timescale recommended for sample withdrawal (Allen, 1981) is the first sample at 1 min and then continuing in a geometric progression at a ratio equal to 2, that is, 2 min, 4 min, 8 min, and so on, as to obtain a smooth curve when plotting cumulative percentage undersize against Stokes' diameter because of the inherent skewed distribution of most powders.

2.6.6.4 Centrifugation

Centrifugal sedimentation can be used to extend the range of applicability of sedimentation to the submicron size. Centrifugal sedimentation can be carried out using a pipette centrifuge that consists of a bowl centrifuge equipped with a withdrawal device connected to a pipette. The principle of particle size evaluation is similar to that used in the Andreasen Pipette, except that the particle is subjected to centrifugal acceleration, instead of gravity force, giving thus the possibility of detecting particles well below the submicron range at reasonably short periods of time. A diagram of the apparatus is

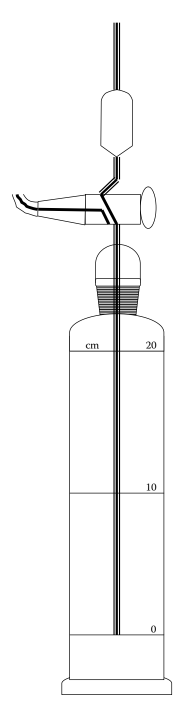


FIGURE 2.31 Andreasen's sedimentation pipette.

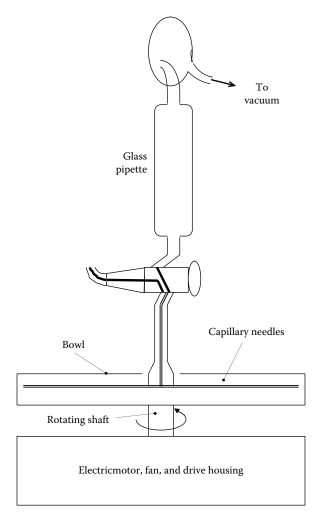


FIGURE 2.32 Diagram of pipette centrifuge.

given in Figure 2.32. As can be seen, it comprises a shallow bowl disc adapted to a hollow shaft equipped with six radially arranged narrow bore needles (a "spider"). The shaft is connected to a pipette by which samples are extracted following a sampling procedure similar to that used for the Andreasen Pipette technique. The modified expression of Stokes' law to account for the centrifugal field can be written as

$$x_1 = \sqrt{\frac{18\mu \ln(R/S)}{(\rho_s - \rho)\omega^2 t_1}}$$
 (2.38)

where x_1 is the largest particle present in the initial sample with-drawn at time t_1 , when the measurement radius is R and the surface is at radius S, while ω is the angular velocity. Allen (1981) presents a procedure for particle size evaluation, analogous to that of the Andresen pipette, using Equation 2.33.

Gravity and centrifugal sedimentation can be combined for the same sample in order to directly determine Stokes' diameter for a wide range of particle sizes. In such a way conversion are avoided and a mass distributions, applicable to processes where gravimetric efficiencies are relevant, can be properly derived. Ortega-Rivas and Svarovsky (1994) determined particle sizes distributions of fines powders using a combined Andreasen Pipette-pipette centrifuge method. They derive relations useful to model hydrocyclone separations, which were later employed to describe apple juice clarification.

Another alternative to speed up a sedimentation particle size measurement is by using an instrument known as photosedimentometer. It operates by exposing the stream flow of a suspension containing the particles to be sized, through a cell across which a light beam is passed. As each particle passes through the beam it blocks a proportion of this according to the cross-sectioned arch of the particle. The number of particles in preset size levels is, thus, easily recorded. Instruments like the photosedimentometer, using a light-blocking technique will, however, have problems in measuring particles whose refractive indices are close to those of the carrier liquid. In general terms, they run into problems with particles smaller than $2\,\mu m$ size.

2.6.6.5 Light Diffraction

Another way of taking advantage of light sources is the use of laser diffraction as a means of detecting particles both in liquid or air streams. The basic principle on which these instruments work is shown in Figure 2.33. In essence, the presence of particles in a light beam causes light diffraction. This action results in a set of light rings at various radii to the incident beam. Each set of diffraction rings is spaced radially at a distance fundamentally related to a specific particle diameter. The size range covered by instruments employing this principle is approximately 2–500 μ m. They are ideally suited to process control operations where results are required quickly with a minimum operator attention. Recent developments in the use of laser diffraction techniques have enabled particles in the submicron range to be analyzed.

2.6.6.6 Particle Counters

In instrumental particle counters, such as the well-known Coulter counter, the stream containing the particles is forced to pass through a flow

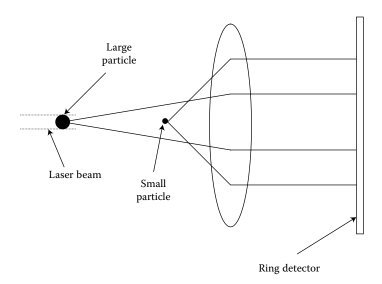


FIGURE 2.33 Principle of operation of a laser diffraction particle counter.

restriction which is subjected to an electrical field. The normal flow of electrical current between the poles is altered by the particles passing through the restriction. These electrical flow changes, function of size of particle passing, are registered as pulses, then counted and grouped according to size. These sort of counters will count and size particles over a range from 800 μm down to 0.5 μm and, as the basic response is related directly to particle volume, the instruments are for all practical purposes independent of problems associated with particle shape, color, or density, which affect most other methods.

2.6.6.7 Online Measurement Techniques

Automation of process control has created a need for continuous monitoring of particle size of particulate matter in process streams. Some online particle size analysis instrumentation has been developed recently to meet this need. It can initiate regulatory or shutdown signals in control systems. The basic requirements for such instrumentation are that it must operate automatically and continuously under preset instructions, and the response time from observations to readout must be so short as to be nearly instantaneous. This relatively new range of techniques follows the general pattern found in the whole subject of particle size measurement. Some just give a measure of central tendency (mean diameter), others give one or more points on the size distribution. They may be truly online in operating on the whole process stream, they may need a partial sample stream taken off the main

one, or they may merely be automated rapid response batch techniques. Online measurement is a buoyant area that is undergoing fast growth. Equipment may be broadly divided into two categories: stream scanning and field scanning. Stream scanning would include different techniques following the basic principle described in the previous section adapted online to various processes. Field scanning usually applies to concentrated systems in which some size dependent behavior of the bulk material is monitored and particle size deduced from theoretical or calibrated relationships. Ultrasonic attenuation, echo measurements, laser attenuation, online viscometry, electric noise correlation techniques, x-ray attenuation, and x-ray fluorescence are examples of such field scanning methods. Most relevant to solid-liquid separation problems are those field scanning instruments which deduce particle size from separation efficiency of some separational equipment using the concept of the analytical cut size. Within these methods, wet sieving machines and separation performance hydrocyclones can be mentioned.

Example 2.5

A sample of calcium carbonate is poured into water to make a test using an Andreasen Pipette. Dispersant is added at a concentration of 0.093% by weight. The density of the calcium carbonate is 2774 kg/m³. Fifteen samples are extracted starting at one minute and following a geometric progression at a 2 ratio. The initial height of the suspension in the Andreasen apparatus is 19.8 cm, and it is observed that every extraction decreases the level 0.4 cm. A container is weighed to determine the initial concentration of the powder in the suspension. Its initial weight (empty) is 1.31275 g, and after drying a 10 mL sample its final weight is 1.43655 (the sample contains the dispersant). Determine the powder median size using a cumulative percentage graph derived from data presented in Table 2.9.

SOLUTION

Since extractions are done following a 2 progression ratio, the sampling times would be at 1 min, 2 min, 4 min, 8 min, and so on until reaching the 15th sample. On the other hand, the initial level in the Andreasen container is 19.8 cm, then in the second extraction it would drop to 19.4 cm, in the third extraction would drop to 19.0 cm, and so on, until reaching the last extraction. In each extraction the fraction of particles having a definite size, function of height, and time as expressed in Equation 2.36 can be related to the initial concentration including all the range of particle sizes, according to Equation 2.37. For example, the fraction of particles having the diameter corresponding to 19.8 height and 1 min time (sample 1), can be calculated by weight difference between the final weight of the container and the initial weight of the container (bearing in mind that the weight of dispersant, which form a solution with the suspending water, has to be subtracted from each sample). Since the dispersant has 0.093% by weight, there will be 0.0093 g of dispersant in each 10 mL sample. Thus, the weight fraction of the first extracted sample

Example 2.5				
Sample	Initial Container Weight (g)	Final Container Weight (g)		
1	1.2913	1.4125		
2	1.2958	1.4152		
3	1.2868	1.4038		
4	1.2898	1.4022		
5	1.3118	1.4183		
6	1.3059	1.4032		
7	1.2924	1.3767		
8	1.3096	1.3792		
9	1.2935	1.3376		
10	1.2847	1.3224		
11	1.2854	1.3126		
12	1.3368	1.3544		
13	1.3393	1.3525		
14	1.3258	1.3366		
15	1.3075	1.3177		

TABLE 2.9 Example 2.5

(from Table 2.9) is = 1.4125–1.2913–0.0093(dispersant) = 0.1119 g. The original weight including all the range of sizes is determined following the same procedure from sample 1, from data given originally, that is, total sample weight = 1.43655–1.31275–0.0093 = 0.1145 g. The weight percentage of the first sample will, therefore, be $(0.1119/0.1145) \times 100 = 97.7\%$. Following this procedure for every sample the weight percentages for every fraction are calculated. For every calculated fraction, the corresponding Stokes diameter $x_{\rm st}$ is calculated as a function of height and time from the following version of Equation 2.36 (British Standards Institution, 1963):

$$x_{\rm st} = 175 \sqrt{\frac{\mu H}{\Delta \rho t}} \tag{2.39}$$

where μ is the viscosity in poises, H is the suspension height in cm, Δp is the difference in densities in g/cm³, and t is the sampling extraction time in min. For example, for the first sample, density, and viscosity (from physical properties tables) are 0.01 poise and 1.0 g/cm³, respectively, while the initial height is 19.8 cm and the sampling extraction time is 1 min. Substituting, thus, these values in Equation 2.39:

$$x_{\rm st} = 175 \sqrt{\frac{(0.01)(19.8)}{(1.744)(1)}} = 58.96 \ \mu \text{m}$$

Calculating, successively, weight fractions and Stokes diameters for all the samples as done for sample 1 above, Table 2.10 can be derived. By inspecting the time column of Table 2.10, it is implied that the experiment would have taken more that 11 days (16,384 min) for completion! In fact, the whole analysis was

TABLE 2.10 Example 2.5

Sample	t (min)	H (cm)	Fraction Concentration	Fraction Percentage	Stokes Diameter $x_{\rm st}$ (μ m)
1	1	19.8	0.1119	97.7	59.0
2	2	19.4	0.1101	96.2	41.3
3	4	19.0	0.1077	94.1	28.9
4	8	18.6	0.1031	90.0	20.2
5	16	18.2	0.0972	84.9	14.1
6	32	17.8	0.0880	76.8	9.9
7	64	17.4	0.0750	65.5	6.9
8	128	17.0	0.0603	52.7	4.8
9	256	16.6	0.0348	41.4	3.4
10	512	16.2	0.0284	37.8	2.4
11	1024	15.8	0.0179	24.7	1.7
12	2048	15.4	0.0083	10.8	1.2
13	4096	15.0	0.0039	4.9	0.8
14	8192	14.6	0.0015	1.7	0.6
15	16384	14.2	0.0009	1.1	0.4

carried out combining sedimentation and centrifugation techniques (Ortega-Rivas and Svarovsky, 1994), and so it was completed in a reasonable time.

Finally, from the tabulated data in Table 2.10 the Stokes diameter x_{st} can be plotted versus the cumulative weight undersize fraction F, as to obtain the graph shown in Figure 2.34.

From the above graph, the median size at the 50% value of $\it F$ is equal to 4.5 μm .

2.7 Laboratory Exercise: Particle Size Analysis by Different Methods

2.7.1 Introduction

Particle size analysis and measurement is an important operation in many industries. The stability, chemical reactivity, opacity, flowability, material strength, and some other properties of many materials are affected by the size distribution and characteristics of the particles within them. There are numerous techniques and instrumentation for all types of particle size analysis and characterization available. Some primary properties of particulate materials, mainly shape, will determine the way the particles are detected or

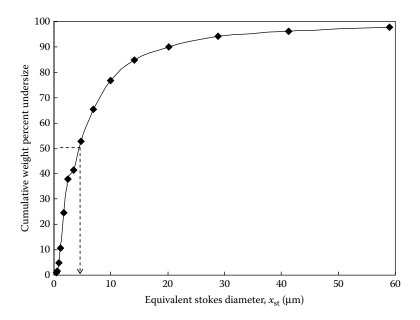


FIGURE 2.34 Graph for Example 2.5.

categorized by different particle size distribution measurement techniques. The objective of the exercise is to determine particle size distributions of industrial powders. At least two methods will be carried out with the same powder, in order to find out possible differences in the whole distribution as well as in the determination of the mean size.

2.7.2 Materials and Instruments

A sample of medium-sized dolomite with particle sizes ranging from approximately $8{\text -}80~\mu m$, and a second sample of coarse-sized dolomite having particles in a range about $40{\text -}400~\mu m$.

An optical microscope, a standard set of sieves, an Andreasen Pipette, and any versatile instrumental particle sizer, such as a laser diffraction equipment or an electrical sensing zone particle counter.

2.7.3 Accessories

A hairline eyepiece, an ocular micrometer or standard graticule, and a stage micrometer, provided for microscopy. Routine laboratory material and ancillary instruments to carry out gravimetric testing, that is, balance, oven, desiccator, measuring cylinders, stop watch, watch glasses, and so on.

2.7.4 Microscopy Method

2.7.4.1 Introduction

Microscopic examination is one of the oldest, most direct, and simplest means for determining the size and size distribution of fine particles. Microscopic examination is tedious but gives reliable and reproducible results if careful measurements are made. As indicated in Table 2.6, microscopy is applicable to particles sizes ranging from about 0.5 to 500 μm if an optical system is used.

2.7.4.2 Slide Preparation

Place a small sample of the powder onto a watch glass. Add a few drops of water containing a little teepol or other dispersing agent, except if the powder is water soluble, in which case an appropriate solvent should be used. Work the suspension with a small camel hair brush or glass rod. Add more or less liquid, so as to adjust the concentration of the suspension to give a representative selection of well-dispersed particles. Using the end of the brush or rod, transfer a drop of the suspension to a clean microscope slide. Cover with a slip taking care to avoid air bubbles.

2.7.4.3 Measurement of Particle Size

Once the particles are on the slide their diameters may be measured. However, since the particles are very likely to be irregular in shape, the meaning of "particle diameter" needs stating. Common equivalent diameters used in microscopy are the projected area diameter (Table 1.3) and statistical diameters, such as Feret's or Martin's diameters (Table 1.4). Feret's and Martin's diameter measurements must be taken in the same direction for all the particles. For the results to be statistically significant, measurements must be made on many particles. Counting of six particles in each of one hundred fields of view is often regarded as minimum.

2.7.4.4 **Procedure**

- Using the stage micrometer, calibrate the eyepiece for a particular object selected.
- Make up the sample as outlined in Section 2.7.1.4. Use coarse-sized dolomite.
- Using the calibrated eyepiece measure a selected statistical diameter (Feret's or Martin's diameter). Measure the diameters of 10 particles in, as many fields of view as possible, but not less than 10.

• Replace the calibrated hairline by the globe and circle graticule provided and with the stage micrometer, calibrate either the globe or the circles. Using the calibrated graticule measure the projected area diameter of 10 particles in, as many fields of view as possible, but not less than 10.

2.7.4.5 Calculations and Report

- Tabulate range of size versus the number of particles counted within the most convenient range of sizes selected.
- Follow a procedure similar to that of Example 2.4, in order to elaborate a histogram, a frequency curve, and an undersize cumulative percentage graph, and determine the median size of the powder (note that the particle size distribution obtained will be by number, while in Example 2.4 the particle size distribution obtained was by mass).
- Perform a literature search to find out particle size distributions of similar materials and compare type of graphs shown and median sizes reported.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out.

2.7.5 Screening Method

2.7.5.1 Introduction

In a size analysis by screening, material is successively passed over a series of sieves having progressively smaller openings. Particles are passed or retained on a particular aperture size of sieve surface. The object of the experiment is to determine the size distribution of coarse-sized dolomite and compare the results with those of the microscopy counting method.

2.7.5.2 Choice of Sieves

Weight a sample of 200 g of powder to be tested. Take a sieve of the top range of openings and a sieve of the bottom range of opening. By visual judgment and trial and error, work out a pair or distant sieve opening so that not more than about 5% of the sample weight should pass the finest sieve or be retained on the coarsest, that is, 10 g passing through the finest sieve and 10 g retained on the coarsest sieve. Once the terminal sieves have been decided upon, the intermediate sieves can then be chosen. For most purposes the ratio of diameters of openings in two successive sieves is: $1:\sqrt{2}$, for example, if the sieve range chosen is 16–60 mesh (BS Test Sieves Series), the choice to give a \bar{x} series would be: 16, 22, 30, 44, 60.

2.7.5.3 **Procedure**

- Weight a fresh sample of 200 g coarse-sized dolomite.
- Select the screens of the series determined by the procedure described above in Section 2.7.5.2.
- "Nest" the chosen sieves together with collection pan and lid.
- Place the weighed sample on the top sieve, assemble the nest on the vibratory unit, and vibrate for about 5 min. carefully remove any fines from the pan at the bottom of the nest. Replace the pan and vibrate for another 2 min. Remove the pan again and take of any fines therein. Repeat the procedure until no more fines are collected in the pan.
- Carefully disassemble the unit and weight the material left on each sieve, making note of the sieve aperture size for each sieve.

2.7.5.4 Calculations and Report

- Follow a procedure similar to that of Example 2.4, in order to elaborate a histogram, a frequency curve, and an undersize cumulative percentage graph, and determine the median size of the powder.
- Compare the results with those from the exercise of the microscopy method, and with results reported in the literature for similar material and same, or different, particle measurement techniques.
- Present a complete written report, according to scientific report guidelines either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out. Discuss on the comparisons made above.

2.7.6 Sedimentation Method

2.7.6.1 Introduction

Different sizes of particle settle at different terminal velocities, and this property can be used in several ways to determine the size distribution of a sample. In the Andreasen Pipette a sample of the material is dispersed in a liquid and allowed to settle from time zero. At intervals a sample of fluid is extract from a depth H below the surface. These samples are evaporated to dryness and weighed to give concentrations. This produces a set of concentrations C_t and the known initial concentration C_0 . At any time t the sample withdrawn will contain particles with terminal velocities equal to or less than H/t, at their original concentration, larger particles will have settled below the level of the pipette tip. The objective of the experiment is to determine the particle size distribution of dolomite powder, and compare

the results obtained with an instrumental method of particle size measurement.

2.7.6.2 Preparation of Sample

Use the medium-sized dolomite sample to prepare a suspension of, approximately, 0.5% by weight. Work out the weight of sample by using its density. For typical inert powders, 5–10 g in a liter of suspending medium will produce a 0.5–2% by weight suspension. Use as suspending medium a 0.1%, by weight, aqueous solution (distilled water) of sodium hexametaphosphate as dispersant. Prepare a set of weighing containers for gravimetric determination of sample concentration. Aluminum dishes, like those used for cup cakes, may be used for this purpose. It is advisable to prepare the dishes needed for each run by placing them in an oven at 100°C for about 2 h, and allow them to attain room temperature in a desiccator for another 2 h. The desiccator should contain only one layer of dishes. Weigh the dishes to 0.1 mg. Record their weights. Calculate gravimetrically the dispersant concentration, as well as the initial sample concentration, using previously prepared and weighed aluminum dishes.

2.7.6.3 Procedure

- Fill up the vessel of the Andreasen Pipette with the sample suspension. Agitate thoroughly. Start the stop watch.
- Extract samples operating the two-way stopcock to fill the 10 mL bulb by suctioning, and emptying the extracted sample by gravity or blowing on the corresponding weighed dish by turning to the second position of the stopcock.
- Continue sampling following a 2 progression ratio until the last programmed extraction.
- Take dishes with samples to an oven, in order to evaporate the suspending medium until complete dryness.
- Allow the samples to cool down and attain room temperature in a desiccator for, at least, 2 h.
- Weigh the dried and conditioned-to-room-temperature aluminum dishes. Record the final container weights.

2.7.6.4 Calculations and Report

- Follow the procedure described in solution of Example 2.5 to calculate, successively, weight fractions and Stokes diameters for all the samples.
- Construct a table similar to Table 2.10, with the collected data of the experiment.

- From the tabulated data draw a graph of the Stokes diameter x_{st} versus the cumulative weight undersize fraction F, as to obtain a plot like the one shown in Figure 2.34.
- From the graph obtained read the median size at the 50% value of *F*.

2.7.7 Instrumental Method

2.7.7.1 Introduction

Instrumental techniques for measurement of particle size distribution of powders have had a tremendous advancement in recent times. Numerous methods and procedures have been developed at a steady pace over the years, and there is the possibility of covering the wide spectrum from nanosystems, to ultrafine powders, and to coarse particulate assemblies. Many instruments offer nowadays quick, reliable results for a wide variety of powders and particulate systems, and for a number of applications. There is still, however, the need to understand the basic principles under which sophisticated instruments operate, as well as to resource to direct measurements under some circumstances. Some of the most modern instrumental techniques are based on an indirect measurement and carry out transformations among the different ways of expressing particles size distributions, that is, by number, surface, or mass. Sometimes it is advisable to avoid transformations because instruments assume a constant shape coefficient on such transformation, which is not necessarily the case, and overestimation or underestimations of size of certain particles may arise. Also, in very specific applications, or in cases of basic or applied research, is better to measure directly the most relevant particle size and particle size distribution. For example, if research is carried out in modeling of solid-liquid separations, a direct measurement of the Stokes equivalent diameter would be most appropriate. The aim of the exercise is to measure the particle size distribution of a sample of medium-sized dolomite, and compare the results with those of the Andreasen Pipette method.

2.7.7.2 **Procedure**

Use any instrumental technique for measurement of particle size distribution, to determine the particle size distribution of the medium-sized dolomite. Follow up the procedure described in the manual of the instrument.

2.7.7.3 Calculations and Report

Gather the information given by the instrument and compare graphs, median size, and any other details about the analysis provided by the instrument, with the corresponding values previously determined using the Andreasen Pipette method. Compare results, also, with data from a literature search and present a written report, including a thoughtful discussion on the comparative aspects mentioned.

Problem 1				
	σ _c (N)			
$\sigma_1(N)$	Powder 1	Powder 2		
10	5.98	4.00		
20	12.01	7.00		
30	17.00	9.67		
40	22.00	12.88		
50	28.11	16.50		

33.03

39.00

44.00

20.00

24.00

26.50

TABLE 2.11

60

70

80

PROBLEMS

- Table 2.11 shows values of the unconfined yield strength against consolidating stress for two industrial powders. Determine their cohesiveness.
- 2. Calculate the median size of a given grain flour whose results after a sieving test are given in Table 2.12.
- 3. A catalyst to be used in a series of processes has been analyzed. Results of such analysis are presented in Table 2.13. Derive the yield locus for the catalyst, including the appropriate Mohr circle, and report the failure properties that can be read from the drawn yield locus.
- 4. Tabulated data resulting from a screening test of three samples of flour are given in Table 2.14. Following a systematic procedure, determine the particle median size for every one of the flours.
- 5. A sample of 15 g carbonated salt with density of 2650 kg/m³ is poured into a liter of water to form a suspension. The prepared suspension was used to fill up an Andreasen Pipette vessel to the 20 cm mark. Suspension samples were taken at: 2, 8, 16, 32, 60, 120, 180, 240, 360, 480, 600, and 1440 min. The weight concentrations on

TABLE 2.12
Problem 2

r robiem z				
Mesh Number (Tyler)	Weight Retained (g)			
20	0.10			
40	10.0			
60	73.2			
80	76.2			
100	17.0			
120	16.5			
200	6.10			
Pan	0.90			

TABLE 2.13 Problem 3

Shear Stress (kPa)	Normal Stress (kPa)		
0.80	0.3		
1.60	1.0		
2.00	1.5		
2.25	2.2		
2.31	2.6		
2.32	2.8		

TABLE 2.14 Problem 4

	Weight Retained (g)		
Mesh Number (Tyler)	A	В	С
20	6.00	0.10	0.40
40	44.00	10.00	1.10
60	29.00	73.20	2.00
80	75.00	76.20	3.50
100	23.00	17.00	12.10
120	2.00	16.50	61.80
200	0.80	6.10	80.00
PLT	0.20	0.90	39.10

 $10~\mathrm{mL}$ of the every extracted sample were: $0.1465,\,0.0967,\,0.0883,\,0.0632,\,0.0386,\,0.0294,\,0.0178,\,0.0104,\,0.0073,\,0.0051,\,0.0030,\,\mathrm{and}\,0.0008~\mathrm{g}$. If every extracted sample drops the height level in the vessel $0.5~\mathrm{cm}$, evaluate the median size of the carbonated salt particles.

References

- Allen, T. 1981. Particle Size Measurement. London: Chapman & Hall.
- Allen, T. and Khan, A. A. 1970. Critical evaluation of powder sampling procedures. *Chemical Engineering* 238: CE108–CE112.
- Augenstein, D. A. and Hogg, R. 1978. An experimental study of the flow of dry powders over inclined surfaces. *Powder Technology* 19: 205–215.
- Beddow, J. K. and Meloy, J. K. 1980. *Testing and Characterization of Powders and Fine Particles*. London: Heyden and Son.
- Beke, B. 1964. *Principles of Comminution*. Budapest: Publishing House of the Hungarian Academy of Science.

- Bergstrom, B. H. 1966. Empirical modification of the Gaudin–Meloy equation. *AIME Transactions* 235: 45–45.
- British Standards Institution. 1963. BS 3406: 1963. Part 2: Methods for the Determination of Particle Size of Powders—Liquid Sedimentation Methods. London: BSI.
- British Standards Institution. 1970. BS 4140: 1970. Part 9: Determination of Angle of Repose. London: BSI.
- Carr, R. L. 1965. Evaluating flow properties of solids. *Chemical Engineering* (January 1965): 163–168.
- Carr, R. L. 1976. Powder and granule properties and mechanics. In Gas-Solids Handling in the Processing Industries, Marchello, J. M. and Gomezplata, A. (eds.). New York, NY: Marcel Dekker.
- Carr, R. L. and Walker, D. M. 1968. Annular shear cell for granular materials. Powder Technology 1: 369–373.
- Chasseray, P. 1994. Physical characteristics of grains and their byproducts. In *Primary Cereal Processing*, Godon, B. and Willm, C. (eds.). New York, NY: VCH Publishers.
- Chen, X. D. 1994. Mathematical analysis of powder discharge through longitudinal slits in a slowly rotating drum: Objective measurements of powder flowability. *Powder Technology* 21: 421–437.
- Dodge, H. F. and Romig, H. G. 1998. Sampling Inspection Tables. Single and Double Sampling. New York, NY: John Wiley and Sons.
- Freeman, R. and Cooke, J. 2006. Testing powders in process relevant ways. *Powder Handling & Processing* 18: 84–87.
- Freeman, R. E., Cooke, J. R., and Schneider, L. C. R. 2009. Measuring shear properties and normal stresses generated within a rotational shear cell for consolidated and non-consolidated powders. *Powder Technology* 190: 65–69.
- Geldart, D. 1973. Types of gas fluidization. *Powder Technology* 7: 285–292.
- Geldart, D. and Wong, A. C. Y. 1985. Fluidization of powders showing degrees of cohesiveness-II. Experiments on rates of de-aeration. *Chemical Engineering Science* 40: 653–661.
- Geldart, D., Mallet, M. F., and Rolfe, N. 1990. Assessing the flowability of powders using angle of repose. *Powder Handling and Processing* 2: 341–345.
- Gy, P. M. 1982. Sampling of Particulate Materials—Theory and Practice. Amsterdam: Elsevier.
- Harris, C. C. 1968. The application of size distribution equations to multi-event comminution processes. *AIME Transactions* 241: 343–358.
- Harris, C. C. 1971/1972. A multi-purpose Alyaudin–Rosin–Rammler–Weibull chart. *Powder Technology* 5: 39–42.
- Herdan, G. 1960. Small Particle Statistics. London: Butterworths.
- Hong, G.-H. and Watanabe, K. 2000. Powder bed tester. An instrument for measuring the powder yield locus. In: *Processing. Part IV*, Wöhlbier, R. H. (ed.). Clausthal-Zellerfeld, Germany: Trans Tech Publications.
- Jenike, A. W. 1964. *Storage and Flow of Solids*. Bulletin No. 123 of the Utah Engineer Experiment Station. Salt Lake City, UT: University of Utah.
- Kaye, B. H. 1981. Direct Characterization of Fine Particles. New York, NY: John Wiley and Sons.
- Kennedy, J. and Neville, A. M. 1986. *Basic Statistical Methods for Engineers and Scientists*. New York, NY: Harper & Row.
- Larson, H. J. 1995. Introduction to Probability. New York, NY: Addison-Wesley.

Bulk Solids 99

Masuda, H. 1997. Powder sampling. In *Powder Technology Handbook*, Gotoh, K., Masuda, H., and Higashitani, K. (eds.). New York, NY: Marcel Dekker.

- Mutsers, S. M. P. and Rietema, K. 1977. The effect of interparticle forces on the expansion of a homogeneous gas-fluidized bed. *Powder Technology* 18: 239–242.
- Ortega-Rivas, E. and Svarovsky, L. 1994. A direct combined method for measuring particle size distribution in obtaining grade efficiency curves. *Powder Handling and Processing* 6: 401–404.
- Peleg, M. 1977. Flowability of food powders and methods for its evaluation—A review. *Journal of Food Process Engineering* 1: 303–328.
- Peleg, M. and Normand, M. D. 1986. Simulation of size reduction and enlargement processes by a modified version of the beta distribution function. AIChE Journal 32: 1928–1930.
- Peleg, M., Normand, M. D., and Rosenau, J. R. 1986. A distribution function for particle populations having a finite size range and a mode independent of the spread. *Powder Technology* 46: 209–214.
- Plowman, P. M. 1985. Practical aspects of sampling. Bulk Solids Handling 5: 1259–1265.
- Rietema, K. and Mutsers, S. M. P. 1973. The effect of interparticle forces on the expansion of a homogeneous gas fluidized bed. *Proceedings International Symposium of Fluidization and Its Applications*. Tolouse, France.
- Rietema, K., Boonstra, J., Schenk, G., and Verkooijen, A. H. M. 1980. The interaction between gas and dispersed solids and its effect on handling operation of solids. *Proceedings of the European Symposium on Particle Technology*. Amsterdam, the Netherlands.
- Rittinger, P. R. 1867. Lehrbauch der Aufbereitungskunde. Berlin: Ernst and Korn.
- Schulze, D. 2007. *Powders and Bulk Solids: Behavior, Characterization, Storage and Flow.* New York, NY: Springer.
- Sutton, H. M. 1976. Flow properties of powders and the role of surface character. In *Characterization of Powder Surfaces*, Parfitt, G. D. and Sing, K. S. W. (eds.). New York, NY: Academic Press.
- Svarovsky, L. 2000. Solid–Liquid Separation. Oxford: Butterworth-Heinemann.
- Teunou, E. and Fitzpatrick, J.J. 2000. Effect of storage time and consolidation of food powder flowability. *Journal of Food Engineering* 43: 97–101.
- Wilkinson, H. N., Dufferll, C. H., Reed, A. R., and Bunting, J. 1983. *Bulk Solids Physical Property Guide*. Ascot, UK: British Material Handling Board.
- Williams, T. C., Birks, A. H., and Bhattacharya, D. 1970/1971. The direct measurement of the failure function of a cohesive powder. *Powder Technology* 4: 328–337.
- Yu, A. B. and Standish, N. 1990. A study of particle size distribution. *Powder Technology* 62: 101–118.

This page intentionally left blank



Storage and Conveying of Bulk Solids

3.1 Introduction: Storage Alternatives

In the materials processing industries there are some particulate systems, such as gravel or ores, which may be stored outdoors in large piles or heaps unprotected from the weather. When taking to process, such materials may be removed by dragline or tractor shovel and delivered to a conveyor. Since many bulk materials are able to deteriorate with outside ambient conditions such as moisture, there is a need to provide for protection to these piled particulate systems. Several alternatives that have been used, like domes and cylindrical structures with conical tops, are normally known as silos. Some of these structured forms of storage have been termed according to a particular application, such as corrugated-wall structures known as grain silos. The reclaiming systems for these types of silos are similar to those used in outdoors storage, that is, draglines, scraper reclaimers, bucket wheel reclaimers, and so on.

There are, however, numerous examples in many materials processing industries where different types of solids are to be stored in confinement, with the intention of being directly connected to processing lines by any sort of conveying system. The vessels used for confined storage of particulate solids are generally known as bins or silos. Difficulties of flow in particulate systems, when needed to be recovered from confined storage, is a well-known operative problem and diverse industries such as mining, chemical, and food industry have invested considerable time and money over the years trying to overcome such situation. Systematic study of solids flow behavior in bins and silos to ensure proper discharge, has been intensive since the formal starting of Powder, or Particle, Technology as a discipline. The pioneering studies of Andrew W. Jenike (Jenike, 1964), established the foundation of silo design and operation.

3.2 Outdoors Storage

Outdoor and structured storage systems are useful when huge amounts of materials need to be in inventory but direct connection to feed processing lines is somewhat difficult. For storage of thousands of tons of particulate materials, proper of some industries like the mining and metallurgy industry, this alternative seems to be the only feasible choice. Estimation of surface area for storage can be done by scaling-up from small samples of the particulate material, using a property like its angle of repose. Space allowance can be done by estimating the volume of the pile through aerial or ground surveys and multiplying by the bulk density of the material. If, as previously pointed out, environment outdoors conditions may affect the integrity of the stored piles of materials, an alternative to protect them would be the use of storage domes (Mitman, 1991; South, 1991).

Material reclaiming and feeding to processing lines can be carried out by manual or mechanized handling. The obvious trend is to use mechanized handling whenever feasible and possible. Apart from substantial savings when using mechanized handling for material reclaiming, additional advantages like efficiency in utilization of storage space, reduction of material wastage, reduction of operator fatigue, and improving control and rotation of stock, can be mentioned. Mechanized reclaiming systems include draglines, scraper reclaimers, bucket wheel reclaimers, and so on. The technology of stockpile storage is a bit outside of the scope of this book, but the reader may refer to valuable literature on the subject (McLean, 1990; Roberts and Teo, 1990; Walker, 1990; Wohlbier, 1977).

3.3 Confined Storage

Open-yard or structured storage may be only a part of a bulk solids handling facility. Due to processing requirements, there is normally the need of having an intermediate storage system which may be directly connected to a processing line. In some other cases, the storage capacity would not involve large tonnages and, therefore, confined storage would be an obvious choice. Furthermore, considering finely divided solids that are subjected to erosion by wind or rain, piling cannot be taken as a feasible manner of storage and, therefore, all these particulate systems may be most properly kept in containers or vessels known also as silos, but alternatively termed as bins and hoppers. In this case, a silo is tall and relatively small in diameter, while a bin is usually not so tall and fairly wide. A hopper is a small bin with a sloping bottom, for temporary storage of solids prior to be fed into a particular process. Normally, the lower part of a silo in a shape of a cone or inverted pyramid is also called a hopper. Generally, all these containers are loaded from the top and discharged from the bottom. A major problem in bin design is related to proper and controllable discharge.

3.3.1 General Considerations

The materials processing industries handle considerable amounts of powders and particulate materials every year. Silos, bins, and hoppers used to store these materials vary in capacity from a few kilos to multiton-capacity vessels. Start-up delays and ongoing inefficiencies are common in solids processing plants. An important cause of these problems is the improper design of bulk solids handling equipment. A six-year study of 40 solids processing plants in the United States and Canada (Merrow, 1988) revealed that 80% of these plants experienced solids handling problems. The study also found that these plants were slow in smoothing operation, with an average start-up time for some plants averaging 18 months. Once start-up began, performance around 40–50% of design was commonly observed.

The common flow problems in hoppers and silos can be summarized as follows: (a) no flow, (b) segregation, (c) flooding, and (d) structural failure. Lack of discharge in the no-flow situation can be attributed to the formation of a stable arch over the outlet, or a stable cavity called a "rathole" (Marinelli and Carson, 1992). With regard to segregation, many materials experience separation of fine and coarse particles (Carson et al., 1986) and such separation can seriously compromise the quality of the final product as well as the efficiency of the process. Flooding can be caused by the collapse of a rathole in a bin containing fine powder, resulting in uncontrollable flow of material, loss of product and clouds of dust (Royal and Carson, 1993), among other problems. Pertaining structural failure, each year over 1000 silos, bins and hoppers fail in North America alone. Most of these failures could have been prevented with proper and careful design, in which the loads imposed by the bulk solid being stored, had been well considered.

The design of bins, hoppers, and silos has never been given the attention it deserves. Approaches using properties such as angle of repose or angle of spatula in design considerations are ineffective, because the resulting values bear no relation to the design parameters needed to ensure reliable flow, mainly because particulate solids tend to compact or consolidate when stored. The attempt of trying to model bulk solids as fluids also leads to a bottleneck, due to the fact that flowing bulk solids generate shear stresses and are able to maintain these stresses even when their flow rate is changed dramatically. It is also improper to consider bulk solids as having viscosity since almost all bulk solids exhibit flow properties that are flow-rate independent. The systematic approach for designing powder handling and processing plants started in the mid-1950s by the pioneering work of Andrew W. Jenike. His concept was to model bulk solids using the principles of continuum mechanics. The resulting comprehensive theory (Jenike, 1964) describing the flow of bulk solids has been applied and perfected over the years, but is generally recognized worldwide as the only scientific guide to bulk solids flow.

The procedures for the design of a bulk solids handling plant are well established and follow four basic steps: (a) determination of the strength and flow properties of the bulk solids for the worst likely flow conditions expected to occur in practice; (b) calculation of the bin, stockpile, feeder, or chute geometry to give the desired capacity to provide a flow pattern with acceptable characteristics, in order to ensure that discharge is reliable and predictable; (c) estimation of the loadings on the bin and hopper walls and on the feeders and chutes under operating conditions; (d) design and detailing of the handling plant including the structure and equipment.

3.3.2 Types of Silos

As stated earlier, silo is recognized as the more generic term to refer to vessels for storage of particulate solids. A silo is a tall vessel with circular section and relatively small diameter ending in a slope bottom part, which is referred to as hopper. Since the early 1960s, mostly from the pioneering work of Jenike (1964), certain terms describing flow patterns started to become common in silo, bin and hopper design. The general theory pertaining to gravity flow of bulk solids has been documented through the years (Arnold et al., 1982; Roberts, 1988) and, from a standpoint of flow patterns there are basically three types of flow in symmetrical geometry: mass flow, funnel flow, and expanded flow.

In mass-flow bins (Figure 3.1a) the hopper is sufficiently steep and smooth to cause flow of all the solids in the bin without stagnant regions during discharge. The bulk solid is in motion at every point within the bin whenever material is drawn from the outlet. The flow is uniform and the bulk density of the feed is practically independent of the head of solids in the bin. Mass flow guarantees complete discharge of the bin contents at predictable flow rates. When properly designed, a mass-flow bin can remix the bulk of the solid during discharge even if segregation is promoted during filling. Mass-flow bins are classified according to the hopper shape and associated flow pattern. The main types of hoppers are: conical, wedge or chisel-shaped hoppers. Only conical hoppers operate with axisymmetrical flow, the remaining types tend to present a plane-flow pattern. In plane-flow bin, the slope to the vertical of the hopper is, on average, 8–10° larger than the corresponding value for axisymmetrical bins with conical hoppers. Therefore, they offer larger storage capacity for the same head room than the axisymmetrical bin, but this advantage may somewhat be offset by the long slotted opening needed, which can cause feeding problems. Figure 3.2 shows some hopper geometries indicating the type of flow. Mass-flow bins are generally recommended for cohesive materials, for materials which degrade with time, for fine powders, and for particulate systems which need to be prevented from segregation.

Funnel flow occurs when the hopper is not sufficiently steep and smooth to force the bulk solid to slide along the walls (Figure 3.1b). It is also

observed when the outlet of the bin is not fully effective, due to poor feeder or gate design. In a funnel-flow bin the stored material flows toward the outlet through a vertical channel forming within stagnant solids. The diameter of such channel approximates the largest dimension of the effective outlet. Flow out of this type of bin is generally erratic and gives rise to segregation problems. However, flow will continue until the level of the bulk solids in the bin drops an amount equal to the draw-down. At this level the bulk strength of the contained material is sufficient to sustain a stable rathole as illustrated in Figure 3.1b. Once the level defined by $H_{\rm D}$ in this figure is reached, there is no further flow and the material below this level represents dead storage. For complete discharge the bin opening needs to be at least equal to the critical rathole dimension, determined at the bottom of the bin corresponding to the bulk strength at this level. For many cohesive bulk solids, and for the normal consolidation heads

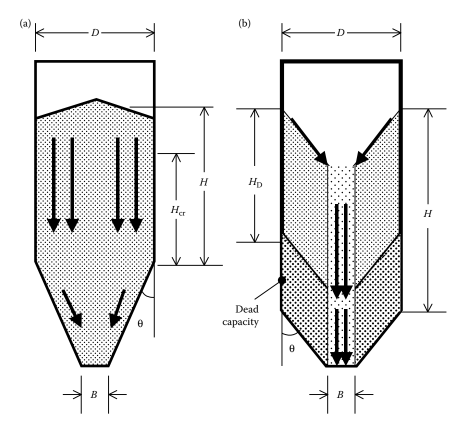


FIGURE 3.1 Types of flow patterns in hoppers. (a) Mass-flow. (b) Funnel-flow. D: silo diameter, H: average depth of material in cylindrical part, H_{cr} : critical depth to initiate flow in mass flow, H_D : dynamic flow depth in funnel flow, θ : hopper slope, B: bottom aperture dimension.

occurring in practice, ratholes measuring several meters high are often observed. This makes control of the product discharge rate quite difficult and funnel flow somewhat impractical. Funnel flow has the advantage of providing wear protection of the bin walls as the material flows against stationary material. However, it is a "first-in last-out" flow pattern which is unsatisfactory for fine bulk solids of low permeability. Such materials can aerate during discharge through the flow channel and this can give rise to flooding problems or uncontrolled discharge. Also, funnel-flow bins are

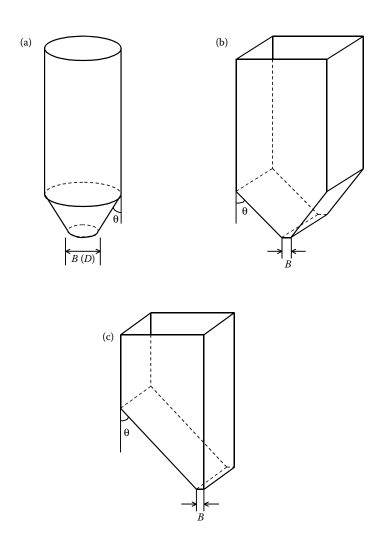


FIGURE 3.2 Common hopper geometries. (a) Axisymmetrical flow, (b) symmetrical plane flow, and (c) nonsymmetrical plane flow.

more prone to cause arching of cohesive solids than mass-flow bins and so they usually require larger outlets for dependable flow. Furthermore, this type of bin may cause segregation of solids and are unsuitable for solids that degrade with time in the stagnant regions. Cleanout of a funnel-flow bin is often uncertain because solid in the stagnant region may pack and cake. For all these reasons, funnel-flow bins are only suitable for coarse, free-flowing, or slightly cohesive, nondegrading solids when segregation is unimportant.

In expanded flow bins the hopper combines characteristics of mass flow and funnel flow, as illustrated in Figure 3.3. The higher part of the hopper operates in funnel flow while the lower operates in mass flow. The mass-flow outlet usually requires a smaller feeder than would be the case for funnel flow. The mass-flow hopper should expand the flow channel to a diagonal or diameter equal to or greater than the critical rathole diameter, thus eliminating the likelihood of ratholing. Funnel-flow bins provide the wall protection of funnel flow, along with the reliable discharge of mass flow. Expanded flow is ideal where large tonnages of bulk solids need to be stored and is particularly suitable for storing large quantities of bulk solids while maintaining acceptable head heights. The concept of expanded flow may be used to advantage in the case of bins or bunkers with multiple outlets. Expanded flow bins are recommended for the storage of large quantities of nondegrading solids. This design is also useful as a modification of

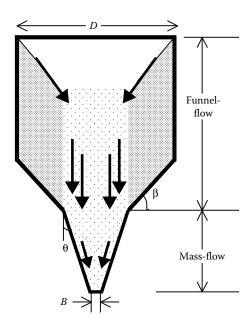


FIGURE 3.3 Expanded-flow hopper.

existing funnel-flow bins to correct erratic flow caused by arching, ratholing, or flushing.

3.3.3 Wall Stresses in Axisymmetrical Bins

The prediction of wall loads in bins is an important piece of information for their design. It is necessary to estimate the pressures at the wall which are generated when the bin is operated, in order to design the bin structure efficiently and economically. The approaches to the study of bin wall loads are varied and involve analytical and numerical techniques, such as finite-element analysis. Despite these varied approaches, it is clear that the loads are directly related to the flow pattern developed in the bin. The flow pattern in mass-flow bins is reasonably easy to predict but in funnel-flow bins such prediction becomes quite a difficult task. For this reason, unless there are compelling causes to do otherwise, bin shapes should be kept simple and symmetric.

Research relating to wall stresses dates back to the 1800s when Janssen (1895) published his now famous theory. More recent investigations (Walker, 1966; Jenike and Johanson, 1968, 1969; Walters, 1973; Clague, 1973) have shown that the solution of the problem of stress distributions in bins is extremely complex. However, most researchers agree that the loads acting on a bin wall are different during the initial stage of filling and during the stage of flowing in discharge, as represented in Figure 3.4. When bulk solids are charged into an empty bin, with the gate closed or the feeder at rest, the bulk solids settle as the solids head rises. In this settlement, the solids contract vertically in the cylindrical section and partially vertically in the hopper section. The major principal stress tends to align with the direction of contraction of the solids, forming which is termed as an active or peaked stress field. It is assumed that the solids are charged into the bin without significant impact to cause packing, and powders are charged at sufficiently low rate so that they deaerate. It is also assumed that the bin and feeder have been designed correctly for the solids to flow without obstruction. When the gate is fully opened or the feeder operates so that the solids start flowing out of the outlet, there is vertical expansion of the solids within the forming flow channel and the flowing mass of solids contract laterally. The major principal stresses within the flow channel tend to align with the lateral contractions and the stress field is said to be passive or arched.

The region of switch from active to passive stress field originates at the outlet of the bin, when the gate is opened or the feeder started, and rapidly moves upward into the bin as the solids are withdrawn. At the switch level a fairly large overpressure may be present, and it is assumed to travel upward with the switch at least to the level at which the channel intersects the vertical section of the bin. For a typical bin consisting of a hopper plus a cylindrical section above it, five stress fields have been recognized during the fill and discharge sequences: (a) in the cylindrical section during initial filling, where

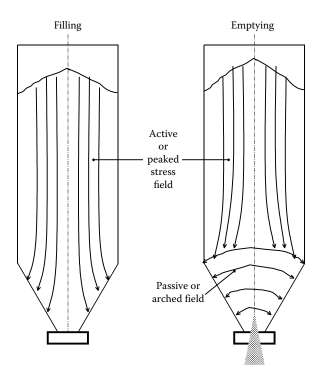


FIGURE 3.4
Stress distribution in silos.

the state of stress is peaked or active; (b) in the cylindrical section during emptying, where the state of stress is either peaked or changes to arched, depending on whether the switch level is assumed to be caught at the transition; (c) in the converging hopper section during filling, where the state of stress is assumed to be peaked; (d) in the converging hopper section during emptying, where the state of stress is assumed to be arched; (e) the switch field, the region in the bin where the peaked stress field established during initial filling is transformed into the arched stress field. This switch starts at the outlet of the hopper, if newly filled from completely empty, and then travels up very quickly as emptying continues, generally to become caught in the transition. Most of the researchers mentioned agree upon a wall pressure or stress distribution as shown in Figure 3.5.

The previously mentioned Janssen theory (Janssen, 1895) includes, possibly, the oldest reported attempt to calculate pressures in silos. Janssen derived an equation for the calculation of vertical and horizontal pressures and wall shear stresses. He assumed a vertical force balance at a slice element spanning the full cross section of a silo being filled with bulk solids, and determined the wall friction coefficient with a shear tester as well as the horizontal pressure ratio from pressure measurements in a model bin. He also assumed

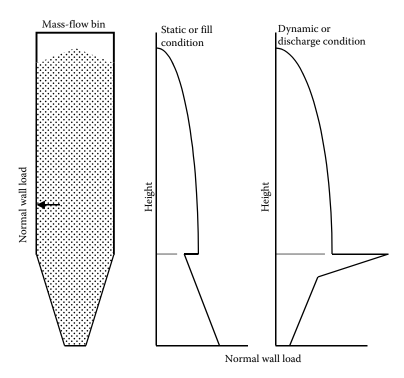


FIGURE 3.5 Wall load distribution in silo.

a constant vertical pressure across the cross section of the slice element and restricted his evaluation to vertical silo walls. The Janssen equation for the vertical pressure $p_{\rm v}$ on dependence of the depth h below the bulk solids top level reads as follows for a cylindrical silo:

$$p_{\rm v} = \frac{g\rho_{\rm b}D}{4\mu'K'} \left[1 - e^{-\left(\frac{4\mu'K'h}{D}\right)} \right]$$
 (3.1)

where g is the acceleration due to gravity, ρ_b is the bulk density of solids, D is the silo diameter, μ' is the sliding friction coefficient along the wall, and K' is the ratio of the horizontal to the vertical pressure, which can be expressed as

$$K' = \frac{1 - \sin \delta}{1 + \sin \delta} \tag{3.1b}$$

where δ is the angle of internal friction of solids.

The advantage of Janssen equation is the simplicity of an analytical equation and its general good agreement with pressure measurements in silos for

the state of filling. The disadvantages are its nonvalidity for the hopper section, its assumption of a constant vertical stress across the cross section, and its assumption of plastic equilibrium throughout the stress field in the silo. The increase in horizontal pressure occurring when discharge initiates cannot be explained by Janssen equation. This disadvantage can only be overcome by using finite-element methods, more sophisticated yield criteria and a very high degree of computational effort (Häußler and Eibl, 1984). Its simplicity and analytical solution have made Janssen equation the basis for the first standard for the calculation of loads in silos well over the years. This equation is still the most widely used analytical solution for the calculation of pressures in silos. However, the structural design of silos requires the incorporation of experience, measurement results in model bins and fullscale silos, as well as accepted safety margins for uncertainties. Major factors contributing to the loads in silos are the flow profile, the flow behavior, the interaction between wall material and bulk solids and the performance of feeders and discharge aids (Jenike, 1964). The loads in silos are influenced by many factors. Some of them are related to the bin structure, its material of construction and size. Many other factors, however, depend on the bulk solids flow properties, the design of outlet size, the type of feeder, the discharge aids, and the operating conditions. These factors are especially important for nonfree-flowing, cohesive bulk solids.

Example 3.1

Suppose that the powder described in Example 2.3 is stored in a silo 12 m high and 3 m diameter, while its bulk density is 850 kg/m³ and its friction coefficient with regard to the silo wall is 045. Calculate the static vertical and horizontal pressures exerted at the base of the cylindrical part of the silo.

SOLUTION

In Example 2.3 the calculated effective angle of internal friction was 42°. Using, therefore, Equation 3.1b and substituting such angle value, constant \mathcal{K}' is calculated as

$$K' = \frac{1 - \sin \delta}{1 + \sin \delta} = \frac{1 - \sin 42^{\circ}}{1 + \sin 42^{\circ}} = \frac{1 - 0.6691}{1 + 0.6691} = 0.1982 \approx 0.2$$

Having computed the value of K', all the variables may be substituted into Equation 3.1, that is,

$$p_{\rm v} = \frac{(9.81)(850)(3)}{4(0.45)(0.2)} \left[1 - e^{-\left(\frac{4\cdot0.45\cdot0.2\cdot12}{3}\right)} \right] \approx 53024 \text{ Pa}$$

Since K' was defined as the ratio of the horizontal to the vertical pressure: $p_h = K'(p_v)$, so that the horizontal pressure p_h is

$$p_h = (K')(p_v) = (0.2)(53024 \text{ Pa}) \approx 10605 \text{ Pa}$$

The results are, thus vertical static pressure $p_{\rm v} \sim 53$ kPa, horizontal static pressure $p_{\rm h} \sim 10.6$ kPa.

3.3.4 Wall Stresses in Other Types of Vessels

For the case of nonsymmetrical flow and some other complex flows like those presented in specific cases such as multioutlet silos, estimation of wall stresses is extremely difficult due to the absence of proximity of constant pressure on the vessels walls. Approaches for calculation in these types of vessels include modeling of a finite element for the interaction silo-solids (Mahmound, 1977) and a simplified theory consisting on describing flow of relatively small channels within a silo (Colijn and Peschl, 1981).

Adapting of silos has also been suggested to try to control pressures on the bins walls. A device for this purpose is known as the antidynamic discharge tube, used to minimize flow problems in tall grain silos. In this type of silo mass flow of grain occurs together with flow along the walls over a substantial height of the silo. This flow effect causes dynamic pressure generation of considerable higher magnitude (two- to threefold), than the static pressure developed in the silo after loading. It has been demonstrated (Thomson, 1997) that by using an antidynamic discharge tube is possible to control the flow pattern, so that funnel flow occurs without flow along the walls. The tube is located over the silo opening aligned to the geometric center and extended nearly all along the silo height. It contains perforations on its entire length to allow grain to enter the tube at different depth levels. On discharge the material flows through these orifices so that the solid only flows inside the tube, and over its top part, as shown in the diagram of Figure 3.6. Due to this flow pattern, the pressure exerted on the walls never exceeds the values of static pressure. A variation of the antidynamic discharge tube consists on a tube without perforations, place at a depth just below the half of the total silo height (Roberts, 1988b). Satisfactory results have been reported in silos with eccentric flow with multioutlet silos. Discharge tubes are only suitable for noncohesive, free-flowing solids such as grains and coarse sand. They should never be used with, even slightly, cohesive solids.

3.3.5 Natural Discharge from Silos

Only fluids can flow; bulk solids under gravity forces can fall, slide or roll, but against gravity, they must be lifted by mechanical means. Solids cannot be pumped by centrifugal or reciprocating pumps; for pumping, bulk solids ought to be suspended in liquids or gases. There is no satisfactory term to

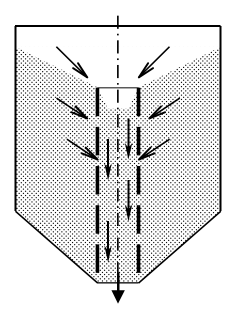


FIGURE 3.6 Principle of antidynamic discharge tube.

describe "flow" of bulk solids as they do not follow strict definitions of fluid behavior, since a fluid is considered to be a continuum in which there are no voids. For a fluid, when the rate of shear is linearly proportional to the shear stress, it is said to be Newtonian and the coefficient of proportionality is called the absolute viscosity. Any deviation from this definition makes the fluid non-Newtonian. Solids in suspension can be referred to as non-Newtonian mixtures to differentiate them from a number of non-Newtonian fluids which are a continuum or are perfectly homogeneous liquids. For all these reasons, bulk solids in suspension are occasionally referred to as "imperfect fluids."

3.3.5.1 Flow Theories

The gravity flow of bulk solids occurs under the pressure corresponding to the equivalent of a "static head" of the material. Such head would be caused by the height of a solid column in a bin, but in practice is often not available to produce the flow due to phenomena known as "arching" or "bridging." The velocity head at the discharge from the bin is usually a small fraction of the head, with the major part being consumed by the friction of the moving solids against the walls of the bin, as well as against like solids. Friction is the resistance which one body offers to the motion of a second body when the latter slides over the former. The friction force is tangent to the surfaces of

contact of the two bodies and always opposes motion. The coefficient of static friction for any two surfaces is the ratio of the limiting friction to the corresponding normal pressure. If a body rests on an inclined plane and if the angle of inclination of the plane to the horizontal is such that motion of the body impends, such angle is defined as the angle of repose. When two surfaces move relative to each other, the ratio of the friction developed to the normal pressure is called the coefficient of kinetic friction and is independent of the normal pressure. The coefficient of kinetic friction is also less than the coefficient of static friction and independent of the relative velocity of the rubbing surfaces. There is experimental evidence suggesting that the value of the kinetic friction coefficient increases as the velocity is decreased, and passes without discontinuity into that of static friction. All these principles would hold under conditions of a particular test, but must be modified in order to apply them to different conditions. It is, therefore, quite difficult to apply existing test data on a series of new tests because of the great variety of flowing conditions of bulk solids. The problem is particularly complicated since the properties of the flowing material depend on time and method of storage operation.

When granular solids are stored in an enclosed container, the lateral pressure exerted on the walls at any point is less than predicted from the head of the material above such point. There is usually friction between the wall and the solid particles, and the interlocking of these particles causes a frictional effect throughout the bulk solid mass. The frictional force at the wall tends to offset the weight of the solid and reduces the effect of the head of solids on the floor of the container. In an extreme case, such frictional force causes the mass of bulk solids to arch, or bridge, so that it would not fall even if the material below is discharged. For many granular solids, when the height of the solid bed reaches about three times the diameter of the bin, additional head of material shows virtually no effect on the pressure at the bin floor.

Solids tend to flow out of any opening near the bottom of a bin but are best discharged through an opening in the floor. The pressure at a side outlet is smaller than the vertical pressure at the same level and removal of solids from one side of a bin considerably increases the lateral pressure on the opposite side while the solids flow. When an outlet in the bottom of a bin containing free-flowing solids is opened, the material immediately above such opening begins to flow. A central column of solids moves downward without disturbing the material at the sides. Eventually lateral flow begins, starting from the top layer of solids and a conical depression forms in the surface of the mass of bulk solids being discharged. The material slides laterally into the central column moving at an angle approaching the angle of internal friction of the solids and the solids at the bin floor are the last to leave. If additional material is added at the top of the bin, at the same rate as the material leaving through the bottom outlet, the solids near the bin walls remain stagnant and do not discharge as long as flow persists. The rate of flow of granular solids by gravity through a circular opening in the bottom of a bin is dependent on the diameter of the opening as well as on the properties of the solid and is independent, within wide limits, on the head or height of the solids.

The amount of solids discharged through an opening in the bottom of a bin, and the failure to restart the flow after intermission, depend on the bin design, shape, and the location of the opening, apart from the flowing properties of the solid. The flowing properties of the granular material include grain nature, size, moisture content, temperature, adhesion, cohesion, and, above all, time of consolidation at rest. There are very few solids that are free flowing and will restart flow after an extended period at rest. Examples of them include inert materials such as graded gravel and dry sand. Pressure distribution within a bin affects its design for strength, but does not enter into calculation of the solids flow from the hopper. It has been mentioned that the volume of bulk solids discharged is independent of the head above the orifice, due to the arching effect. Therefore, the design of bins or silos in terms of their ability to initiate flow without any aid, is based on solid mechanics theories considering only the hopper of the container.

The mass-flow and funnel-flow limits in silos are well known and have been used extensively in proper design. The limits for conical hoppers and plane hoppers depend on the hopper half-angle θ , the effective angle of internal friction δ and the wall friction angle ϕ . Once the wall friction angle and effective angle of internal friction have been determined by experimental means, the hopper half angle θ may be determined. In function form it can be expressed as

$$\theta = f(\varphi, \delta) \tag{3.2}$$

The bounds for conical and plane-flow hoppers are plotted for three values of δ in Figure 3.7. For the case of conical hoppers, it is recommended that the half-angle θ be chosen to be 3° less than the limiting value. For plane hoppers the bounds between mass-flow and funnel-flow are much less critical than for conical hoppers. In plane hoppers much larger hopper half-angles are possible, which means that the discharging bulk solids will undergo a significant change in direction as it moves from the cylindrical part to the hopper bottom.

Typically, flow in a hopper is based on the radial stress field theory, which does not take into account the influence of the surcharge head due to the cylinder on the flow pattern developed. It is, however, known that complete mass flow in a hopper is affected by the cylinder surcharge head. There is a minimum level $H_{\rm cr}$ (Figure 3.1), which is required to enforce mass flow in the hopper. For a common mass-flow bin this level is from about 0.75D to 1.0D. Benink (1989) identified a third flow region, apart from mass flow and funnel flow, which has been known as intermediate flow. The three flow regions suggested by this author are illustrated in Figure 3.8. According to Benink (1989) the surcharge head has a significant influence on the flow pattern generated.

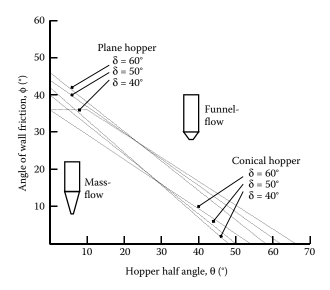


FIGURE 3.7 Limits for mass-flow in conical and plane-flow channels.

In the study, fundamental relationships for $H_{\rm cr}$ in terms of the different bulk solids and hopper geometrical features are derived. These relationships are used to predict the critical height $H_{\rm cr}$ at which the flow changes.

3.3.5.2 Hopper Opening for Coarse Bulk Solids

Coarse bulk solids are those in which the particle size range is such that the air permeability is sufficiently high to allow air to percolate easily through

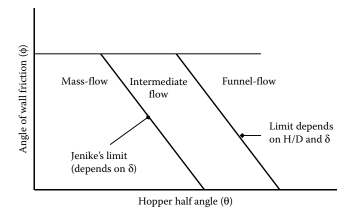


FIGURE 3.8 Flow regimes for plane-flow hopper defined by Benink (1989).

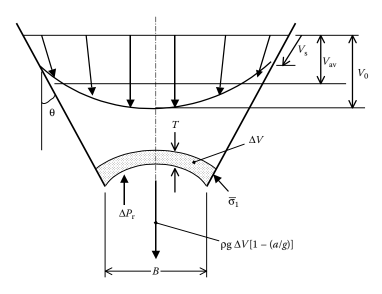


FIGURE 3.9 Flow channel and equilibrium of forces on the arch formed on a hopper.

the stored solids. According to the diagram shown in Figure 3.9, the accelerated flow of bulk solids in the region of the outlet of the hopper can be considered when the air pressure gradient Δp_r equals zero. Analyzing the forces it may be shown that

$$\overline{\sigma}_1 = \frac{\rho_b g B}{H(\theta)} \left[1 - \frac{a}{g} \right] \tag{3.3}$$

where $\bar{\sigma}_1$ is the stress acting in arch at angle 45°, ρ_b is the bulk density of solids, g is the acceleration due to gravity, B is the hopper opening, a is the acceleration of discharging bulk solids and $H(\theta)$ is a factor to account for variation in arch thickness, hopper half-angle, and hopper type, whether conical or plane. It has an approximate value of 2.2 for plane-flow hoppers and 2.4 for conical hoppers.

The minimum hopper opening to just prevent a cohesive arch from forming occurs when static equilibrium prevails, that is, when the acceleration of discharging bulk solids approaches zero, and thus, substituting a = 0 into Equation 3.3 and transposing:

$$B_{\min} = \frac{\bar{\sigma}_1 H(\theta)}{\rho_b g} \tag{3.4}$$

The hopper half-angle is chosen from the mass-flow limits as shown in Figure 3.7, while the condition for $\bar{\sigma}_1 = \sigma_c$ is obtained from the intersection

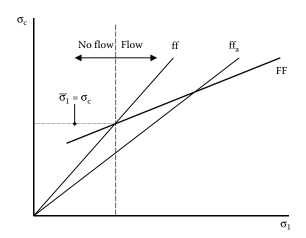


FIGURE 3.10 Design graphic for hopper geometry.

point of the flow factor ff line and failure function FF as illustrated in Figure 3.10. As discussed in the previous chapter, the failure function is a bulk solid parameter and represents bulk strength, while the flow factor is a flow channel parameter. The flow factor ff can be defined by the relationship:

$$ff = \frac{\overline{\sigma}_1}{\sigma_c} \tag{3.5}$$

The flow factor ff and function $H(\theta)$ are given as design curves (Jenike, 1964; Arnold et al., 1982; Roberts, 1988) like the one given in Figure 3.11. Following the work of Johanson (1965), it may be shown that the acceleration in Equation 3.4 may be expressed as

$$a = g \left[1 - \frac{ff}{ff_a} \right] \tag{3.6}$$

where ff is the critical flow factor based on the minimum arching dimension and ff_a is the actual flow factor based on the actual opening dimension. This actual flow factor may be represented as the relationship between stresses, according to Figure 3.10, by

$$ff_a = \frac{\sigma_1}{\sigma_c} \tag{3.7}$$

where σ_1 is the major consolidation pressure at outlet corresponding to dimension B.

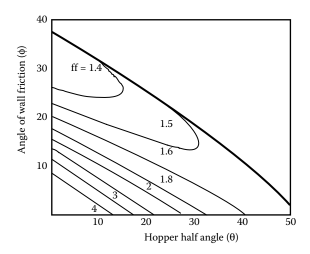


FIGURE 3.11 Flow factor graph for conical hopper and effective angle of internal friction, $\delta = 40^{\circ}$.

The acceleration has two components, that is,

$$a = a_{\rm c} + a_{\rm v} \tag{3.8}$$

where $a_{\rm c}$ is the convergence component due to flow channel and $a_{\rm v}$ is the component due to velocity increase as flow is initiated. This last component may be represented as a function by

$$a_{\rm v} = g \left[1 - \frac{\rm ff}{\rm ff_a} \right] - \frac{2V^2(m+1)\tan\theta}{B}$$
 (3.9)

where V is the discharge velocity and m is a constant which takes the value of zero for plane-flow hoppers and unity for axisymmetrical or conical hoppers.

Equation 3.9 shows that as the discharge velocity V increases, the component due to velocity increase as flow is initiated a_v tends to zero. Thus, an average terminal discharge velocity V_a is reached when $a_v = 0$, that is,

$$V_{\rm a} = \sqrt{\frac{Bg}{2(m+1)\tan\theta}} \left[1 - \frac{ff}{ff_{\rm a}} \right]$$
 (3.10)

The flow rate Q_0 may be calculated by the following equation:

$$Q_0 = \rho_b B^{(1+m)} L^{(1-m)} (\pi/4)^m V_a$$
(3.11)

where *L* is the length of slot for rectangular opening in plane-flow hoppers, in which case, *B* would represent the wide of slot. It can be easily seen that, for conical hoppers *B* would represent the diameter of the circular opening and, thus, Equation 3.10 above is applicable for both cases of plane-flow hoppers, either with square or slot opening, and axisymmetrical or conical hoppers.

For design purposes B and Q_0 can be plotted as a function of θ , as presented in Figure 3.12. In such a way, several options of hopper geometry can be explored. In many cases the flow rate determined above the unimpeded discharge will be well in excess of the plant requirements. For this reason feeders may be used to control discharge rate to any required value.

3.3.5.3 Hopper Opening for Fine Bulk Solids

For the case of fine powders, discharge from a hopper will be rather difficult due to the low permeability of the powder to the air flow. Flow rates much lower than those calculated by the method described above can be common in practice. The analysis of the flow process is much more complex and involves two-phase flow theory. The flow of fine powders has been studied over the years by several researchers such as McLean (1979), Arnold et al. (1982), and Arnold and Gu (1990). Larger hopper openings are required than for the equivalent coarse bulk solids and, sometimes, air permeation is needed to bias the pore pressure in the hopper in order to assist the discharge. Fine powders are prone to flooding and uncontrollable discharge, when allowed to aerate. For this reason extreme care should be taken in designing and installing any air permeation system. Care must also be exercised in ensuring that the interface between the hopper and feeder is correctly designed to prevent problems due to flooding.

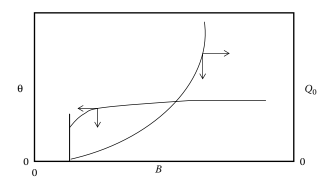


FIGURE 3.12 Hopper half-angle θ and mass flow rate versus hopper opening *B*.

Example 3.2

The powder characterized in Example 2.3 has the failure function given in Table 3.1 while its bulk density is 960 kg/m^3 . If the determined angle of wall friction is 18° , calculate the hopper opening and the discharge flow rate, considering that the term ff/ff_a can be considered negligible.

SOLUTION

The effective angle of internal friction determined in Example 2.3 was 42°, and so the design graph for 40° may be used for calculations. In such graph, the value of $\phi=18$ is prolonged until a distance 3° short from the limit line of funnel flow to read a value of ff = 1.5 (Figure 3.13). The inverse of ff will give the slope of the line to be drawn in the failure function graph, derived from Table 3.1. The slope of the line at a value of ff = 1.5 is 34°. Drawing a line of such slope in the graph of the failure function a value for $\bar{\sigma}_1$ would, therefore, be about 1180 N/m² as indicated in Figure 3.14.

Substituting, thus, values in Equation 3.4:

$$B_{\text{min}} = \frac{\overline{\sigma}_1 H(\theta)}{\rho_b g} = \frac{(1180)(2.4)}{(960)(9.81)} = 0.3007 \text{ m} \sim 30 \text{ cm}$$

To evaluate the discharge mass flow rate Q, the discharge velocity V_a needs to be determined. The hopper half-angle is read of from Figure 3.13 as 28°, so that $\tan (28^\circ) = 0.5317$. For conical hoppers m = 1, and ff/ff_a approaches zero. With all these figures and considerations, substituting into Equation 3.10:

$$V_{\rm a} = \sqrt{\frac{Bg}{2(m+1)\tan\theta} \left[1 - \frac{\rm ff}{\rm ff_a}\right]} = \sqrt{\frac{(0.3)(9.81)}{2(1+1)(0.5317)} \left[1 - 0\right]} = \sqrt{\frac{2.943}{1.8652}} = 1.176 \, \rm m/s$$

TABLE 3.1 Example 3.2

σ_1 (kPa)	σ _c (kPa)		
0.2	0.30		
0.4	0.51		
0.6	0.66		
0.8	0.81		
1.0	0.92		
1.2	1.03		
1.4	1.10		
1.6	1.16		
1.8	1.18		
2.0	1.20		

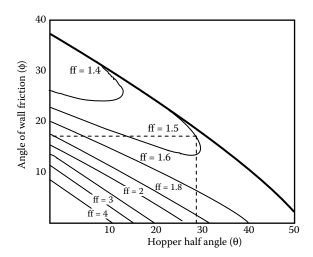


FIGURE 3.13 Flow factor graph (similar to Figure 3.11) showing the value of ff and the hopper half angle.

Equation 3.11 for a value of m=1 would reduce to the expression below. Substituting the derived value $V_{\rm a}$ above and the hopper opening dimension calculated previously:

$$Q_0 = \rho_b \left(\frac{\pi(B)^2}{4}\right) V_a = (960) \left(\frac{\pi(0.3)^2}{4}\right) (1.176) = 79.82 \text{ kg/s}$$

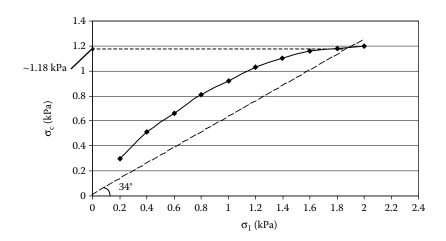


FIGURE 3.14 Failure function from Table 3.1 showing the critical stress defined by Equation 3.3.

The optimum hopper opening dimension would therefore, be 30 cm approximately and, with such an aperture, a discharge flow rate of about 80 kg/s can be obtained.

3.3.6 Assisted Discharge

3.3.6.1 Types of Discharge

The above-mentioned procedures to determine optimum hopper outlet diameter might be taken only as an estimate, because in real applications problems due to flow blockages, such as arching and ratholing, may occur. In order to ensure flow from bins, even after the hopper geometry has been determined following careful calculations such as those previously discussed, flow promotion may be necessary. Classification of flow promotion may be termed as passive and active involving energy. A third class of flow promotion may be considered the use of feeders, which are useful not only to promote flow, but also to control the flow rate.

3.3.6.2 Passive Devices

These types of devices for flow promotion do not require energy and are normally known as inserts. An insert is usually placed within the hopper section of a bin, with the purpose to expand the size of the active flow channel in a funnel flow bin, as to approach mass flow. Another aim of an insert is to relieve pressure at the outlet region. Inverted cones and pyramids have been used for years in this regard, but with limited success. Apart from the typical inverted cone or pyramid insert, the cone-in-cone insert or Binsert® (Johanson, 1982), has also been tested as a flow promoter. As shown in Figure 3.15, the Binsert is actually a sort of funnel inside an expanded flow hopper, or a hopper within another hopper. In this insert, the material flows through the inner hopper, as well as through the annulus between the inner and the outer hopper. It has proved very effective and has the added advantage of promoting not only flow, but also mixing or blending capability, provided the bin geometry and insert are correctly selected. In such a way, a completely uniform velocity with an absolute minimum degree of particle segregation can be achieved.

3.3.6.3 Active Devices

Vibrating hoppers by the use of electrical motors, pneumatic knockers, eccentric drives, or electromagnetic units, are one of the most important and versatile flow assisters. The application of mechanical vibrators to promote gravity flow of bulk solids has been studied in detail by Roberts et al. (1986). They developed a dynamic shear test apparatus, which permits the shear strength to be determined in the presence of vibrations which may be applied over a range of amplitudes and frequencies. The strength of bulk solids is

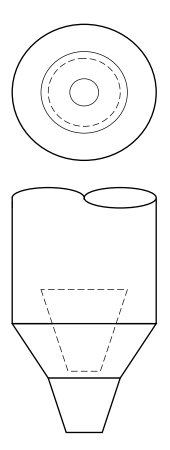


FIGURE 3.15
BINSERT® system.

measured in terms of the yield locus and Mohr diagram as shown in Figure 3.16a. This figure shows the two yield loci for vibration and no vibration conditions. The unconfined yield strengths σ_c and σ_{cf} in the mentioned figure give one point in the respective flow functions shown in Figure 3.16b. A family of yield loci, for at least three consolidation conditions, need to be obtained for the complete flow functions in Figure 3.16b to be determined.

The dynamic shear strength decays exponentially with increase in vibration velocity in agreement with the following relation:

$$\tau_{\rm f} = \tau_{\rm fo} \left[\frac{\tau_{\infty}}{\tau_{\rm fo}} + \left(1 - \frac{\tau_{\infty}}{\tau_{\rm fo}} \right) \exp \left\{ -\frac{2\pi X_{\rm r} f}{U} \right\} \right]$$
(3.12)

where f is the frequency, while τ_{∞} and U are constants depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid.

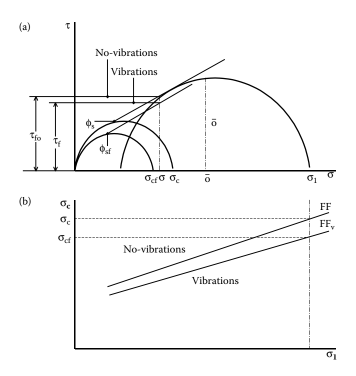


FIGURE 3.16Yield loci and flow functions in vibration and no-vibration situations.

Constant U in Equation 3.12 is the bulk solid vibration velocity constant. Experience has shown that U is independent of the consolidation pressure and applied normal pressure. Knowing the value of U for a particular bulk solid, the values of the relative amplitude X_r and the frequency f for maximum shear strength may be calculated from

$$X_{\rm r}f \ge 0.8\,U\tag{3.13}$$

where X_r is expressed in millimeters, f in Hertz, and U in mm/s. The best results are achieved by using higher frequencies and lower amplitudes.

The decay in shear strength with vibration velocity is similar to the decay in shear strength with voidage on the shear plane. These findings may seem to suggest that the vibration velocity is directly related to the voidage. Reliable gravity flow in bins depends, to a very significant extent, on the magnitude of the friction angle between the flowing bulk solids and the hopper wall. As previously stated, in hopper design the wall friction angle has a major influence on the flow pattern developed. Vibrations applied to the hopper wall can reduce wall friction to a marked extent. Dynamic wall shear tests

may be readily performed using the dynamic shear apparatus. The influence of mechanical vibration may induce flow by: (a) improving the mass-flow performance of an existing mass-flow bin by reducing the wall friction angle and critical arching dimension, (b) improving the funnel-flow performance of an existing funnel-flow bin by reducing the critical rathole dimension, (c) by lowering both the bulk strength and wall friction angle to convert an existing funnel-flow bin to a mass-flow bin.

Effective flow promotion depends on the ability of the stored bulk mass to transmit vibration energy to the region of the flow blockage. It is usual to install the vibrator on the hopper wall as to provide an immediate benefit through the resultant reduction in wall friction. If the flow blockage is in the form of an arch, vibration applied to the hopper wall at or near the outlet may cause the arch to fail and flow to occur. On the other hand, for stable ratholes, the vibration energy needs to be transmitted through the bulk mass in order to assist flow. The dynamic shear test apparatus, coupled with the theory of failure, provides information on the frequency and amplitude necessary to be applied at the zone of the flow blockage (Roberts et al., 1986). For best results, the higher modes of vibration should be used in flow promotion in order to create multiple tensile planes of failure as illustrated in Figure 3.17. Excitation frequencies in the order of 100 Hz, or higher, are necessary. Generally, the higher the frequency the higher the mode of vibration created in the bulk mass and, hence, the greater the number of failure zones. On the other hand, the vibration energy transmitted may have a higher attenuation at the higher excitation frequencies.

Air cannons or air blasters are also commonly used to promote gravity flow in bins. Terziovski and Arnold (1990) indicate that the sizing and placing of air blasters are normally carried out using techniques based on experience

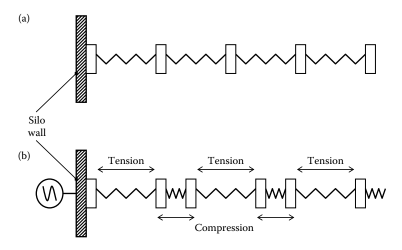


FIGURE 3.17 Formation of failure planes by vibration. (a) Static case and (b) dynamic case.

and guesswork. Based on their own research on the use of air blasters, they provide a procedure for the correct selection of these devices for efficient performance using the measured flow properties of the bulk solid and the bin geometry. In general, air blasters are used in a retrofit situation to correct an inadequate design where flow blockages occur. However, in a new design where there is some uncertainty in the variations in bulk strength of a bulk solid for the full range of operating conditions, it is a good practice to make provision in the design for future installation of air blasters, in case it would become necessary.

3.3.7 Feeders for Discharge Control

In designing hoppers for silos, the procedure described so far would consist on making calculations to determine optimum hopper slope and outlet opening in order to ensure flow. In case flow does not occur, a flow promotion device can be selected after careful study of conditions and factors. Once flow out of a bin is guaranteed, the next step to complete the proper design of the bulk storage plant would consist on controlling the flow rate, as to provide adequate feed to any given food powder process. In order to do so, the use of a feeder will become necessary.

3.3.7.1 Feeders Description

A feeder is a device used to control the flow of bulk solids from a bin. Any feeder must be selected to suit a particular bulk solid and the range of feed rates required. It is particularly important to design the hopper and feeder as an integral unit, in order to ensure that the flow from the hopper is fully developed with uniform draw of material from the entire hopper outlet. There are several types of feeders but the most common are the belt or apron feeder, the screw feeder, the vibratory feeder, and the star feeder. Careful considerations, such as those described above, should be taken in selecting a feeder for a particular application.

An apron feeder consists on a conveyor belt positioned just below a tapered opening of a hopper. Due to the geometry of the conveyor, apron feeders are particularly suitable for slot opening hoppers. Care must be taken that dead spots are not produced in the flow channel above the feeder belt. The capacity of a belt feeder can be increased by tapering the outlet in the horizontal and vertical planes. In order to ensure proper flow of nonfree-flowing solids along the front bin wall, a sloping striker plate at the front of the hopper may be necessary. Belts have been used successfully under slot openings as long as 30 m with a constant slot width of 205 mm. Provisions should be made for field adjustment of the space between the skirt and the belt to provide uniform flow along the entire length. Since the minimum distance between the skirt and the belt should allow the largest particles to pass under, very long belt feeders are limited to the finer solids.

A screw adapted to a hopper bin is known as a screw feeder. Screw feeders are quite useful in producing uniform feed rates for a variety of bilk solids. An important factor in this kind of feeder is the need for a variable pitch screw to produce a uniform draw of material across the entire hopper opening. For uniform flow to occur, the screw feeder opening-to-diameter ratio should not exceed a value of 6.

In a vibratory feeder, there is an inclined pan with different sort of motion located below a hopper opening. Vibratory feeders provide uniform flow along a slot opening of limited length. The distance between the feeder pan and the hopper is increased in the feed direction. Slot length is limited by the motion of the feeder. Because in long slots the upward component of motion is not relieved by the front opening, solids tend to pack. This can cause flow problems with sticky bulk materials, as well as large demand of power for free-flowing bulk solids. To overcome such difficulties, vibratory feeders and reciprocating-plate feeders are designed to feed across the slot. Although this type of feeder may require several drives to accommodate extreme width, the drives are small due to the short length of the feeder.

Star feeders may provide highly uniform withdrawal along a slot hopper opening. They normally comprise six paddles whose ends converse in a common point forming a radial arrangement, or a six-point star when looking at its cross section. When the star rotates, withdrawal of bulk solids is attained. A vertical section of at least one outlet width should be added above the feeder to ensure uniform discharge across the opening. Figure 3.18 presents schematic diagrams of the four types of feeders described, while Table 3.2 shows a preliminary guide for choosing a feeder.

3.3.7.2 Charge and Power Calculations

The estimation of feeders loads and drives powers requires knowledge of the stress fields generated in the hopper during the initial filling conditions, as well as during discharge. Under filling conditions, a peaked stress field is generated throughout the entire bin. As illustrated in Figure 3.19, once flow is initiated an arched stress field is generated in the hopper and a much greater proportion of the bin load is supported by the walls of the hopper. Consequently, the load acting on the feeder is reduced considerably. It is common for the load acting on the feeder under flow conditions to be in the order or 20% of the initial load. The arched stress field is quite stable and is maintained even if the flow is stopped. This means that once flow is initiated and then the feeder is stopped while the bin is still full, the arched stress field is retained and the load on the feeder remains at the reduced value. The load W on the feeder is given by

$$W = q \rho_b g L^{(1-m)} B^{(2+m)} \tag{3.14}$$

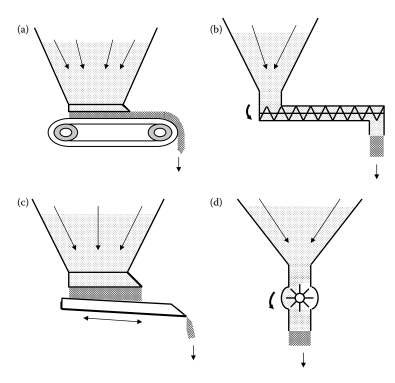


FIGURE 3.18
Types of feeders. (a) Apron feeder; (b) screw feeder; (c) vibratory feeder; and (d) star feeder.

where q is a nondimensional surcharge factor, ρ_b is the bulk density of solids, g is the acceleration due to gravity, L is the length of the slotted opening, B is the width of slot or diameter of circular opening, and m is the symmetry factor whose values are 0 for plane flow hopper and 1 for conical hopper. Equations for q are available in the literature (Manjunath and Roberts, 1986, 1986b).

TABLE 3.2Guide for Feeder Selection

Surac for recuer beleetion							
Material Characteristics	Type of Feeder						
Free flowing materials	Belt, oscillating or vibrating, screw, star						
Nonabrasive granular materials	Apron, bar flight, belt, oscillating or vibrating, screw, star						
Difficult to handle materials (abrasive, hot, etc.)	Apron, bar flight, belt, oscillating or vibrating, reciprocating						
Heavy materials, materials with lumps, highly abrasive	Apron, oscillating or vibrating, reciprocating						

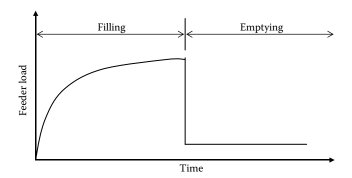


FIGURE 3.19
Load on a feeder at filling and emptying conditions.

3.3.7.3 Discharge Control in Feeders

The loads on feeders and the torque during start-up may be controlled by ensuring that an arched stress field fully or partially exists in the hopper prior to starting. This may be achieved by: (a) cushioning in the hopper, that is, leaving a quantity of material in the hopper as a buffer storage, (b) starting the feeder under the empty hopper before initiating filling, (c) raising the feeder up against the hopper bottom during filling and then lowering it to the operating conditions prior to starting; in this way an arched stress field may be partially established.

3.3.8 Other Design Factors

Previous sections have underlain, in a sort of timing sequence, the steps suggested for the optimum design of a confined storage system for bulk solids. Special emphasis has been placed on the hopper design, as it is considered the key issue for discharge control from bins. There are, however, several other factors influencing selection and design of bins and silos for bulk solids storage. Some of these factors are: surface or wall friction in relation to cohesion and adhesion, wear in bulk handling plant due to impact and abrasion, and additional physical characteristics of the bulk material.

Friction and their related properties, like cohesion and adhesion, have tremendous influence in the performance of hoppers, feeders, and chutes. For this reason, lining of surfaces in contact with the flowing solid may be considered to facilitate handling. It has been observed that friction depends on the interaction between the relevant properties of the bulk solid and lining surface, with external factors such as loading conditions and environmental features. Careful choice of lining material to achieve low fraction and wear is a matter of great relevance. Factors to be considered in the selection of lining materials include surface friction, adhesive characteristics,

resistance to impact, anticorrosive properties, method of attachment, and cost. It is advisable to carry out appropriate tests to determine the relevant flow properties of the bulk solid and the proposed lining material. The fundamental property of the solid to be considered would be the wall friction angle ϕ that can be determined as described in Section 2.4.3, or, alternatively, may be defined as follows:

$$\varphi = \tan^{-1} \left(\frac{\tau}{\sigma_{w}} \right) \tag{3.15}$$

where τ is the shear stress at wall surface and σ_w is the corresponding normal stress or pressure. The friction angle ϕ between the bulk solid and the boundary surface decreases as the normal pressure increases.

A common problem in bulk solids handling plants is the buildup of solids in surfaces of the discharging and conveying devices, due to cohesiveness and adhesiveness of the bulk solids being handled. If buildup is not controlled, blockages may eventually occur. In order to avoid buildup and blockages, it is necessary for the body forces generated in the bulk mass be sufficient to overcome the forces due to adhesion and shear. The body forces are normally those due to weight component of the bulk solid, but also may include inertia forces in dynamic systems such as in the case of belt conveyor discharge. The most suitable materials and the possibility of lining should be considered for feeders and chutes. In extreme cases, vibratory devices adapted to these components of the bulk solids handling plant, may be contemplated.

Wear of bulk solids handling plants is one of the major problems resulting from routine processing activity. Wear may result from impact or abrasion or, as is often the case, from a combination of both factors. Erosive type wear due to impact consists of plastic deformation and cutting wear. Excessive wear may occur in pipe bends of pneumatic conveying systems and at the discharge points of belt conveyors. Impact wear depends on factors such as the relative hardness of the particles and the condition of the wall surface. Abrasive or rubbing wear occurs in storage bins and silos, particularly in hoppers, under mass-flow conditions. On such conditions, the pressures in a hopper will vary significantly over the whole surface, with the maximum pressure occurring at the transition and then decreasing toward the outlet. The velocity of the solids adjacent to the hopper wall increases nonlinearly from the transition to the outlet. The magnitude of the velocity at a particular point on the hopper wall depends on the bin discharge rate and the bulk solid velocities are low, with pure sliding taking place. Abrasive wear also occurs in transfer chutes, feeders, belt conveyors, vibratory conveyors, screw conveyors, and, in general, in any mechanical device involving the motion of bulk solids. Roberts et al. (1990) have suggested the use of a nondimensional Relative Wear Number N_{WR} to allow for comparisons between the different bin and chute geometries. $N_{\rm WR}$ is defined as

$$N_{\rm WR} = \left(\frac{\sigma_{\rm w}}{\rho_{\rm b} g B}\right) \left(\frac{V_{\rm s}}{V_{\rm 0}}\right) \tan \varphi \tag{3.16}$$

where σ_w is the normal pressure at boundary, ρ_b is the solid bulk density, g is the acceleration due to gravity, B is the outlet dimension of hopper or chute width, V_s is the velocity at the sliding wall, V_0 is the sliding velocity at a reference location, and ϕ is the wall friction angle.

Some other characteristics of the bulk solids can affect design and flowing capacity in a bulk solids handling plant. In terms of particle size, bulk solids will flow better at a coarser range of particle sizes and with minimum presence of fines. Conditioning by agglomeration or any other process in order to minimize fines and improve flowability, is sometimes carried out in bulk solids handling plants. The alternative may be, however, time consuming and economically unfeasible. The moisture affects flow, with bulk solids flowing better at lower moisture content. Many inert industrial powders can absorb water up to a certain point, before presenting severe flow problems. Biological materials can be better water absorbents and may cause flow problems, unless a treatment, like drying, is included in their handling. The cost incurred in drying may be offset by improving flow, minimizing shipping costs, and controlling deterioration losses. Moisture content can also be controlled by substituting the air in the voidage of the bed of stored particles, by a dry stable gas such as nitrogen. Regarding temperature, the flow may become difficult at high temperatures, mostly in biological materials that may release sticky substances of low melting points. In this case, a cooling system may be necessary to maintain stored solids free from releasing substances. As with the drying alternative previously mentioned, consideration should be given to the possible compensation of the cost of cooling, in terms of flow improvement and some other advantages, such as quality preservation of the stored material. Storing time may also affect flowability as attrition and erosion may smooth sharp edges of polyhedral particles rounding them and facilitating their flow.

As a final part of design factors in a bulk solids handling plant, ancillary accessories like gates and level gauges may be taken into account. Gates are used to control release of flow from bins and hoppers to feeders or conveyors. There are several gate designs, such as the hand-slide gate, the lever-operated quadrant gate, and the rack-and-pinion gate. The hand-slide gate could be, in fact, quite difficult to operate by hand, and so it is restricted to low-capacity vessels and light bulk solids. The lever-operated quadrant gate is not used to control the flow, but rather to allow the free discharge of lumpy materials with a quick opening turn. The rack-and-pinion gate is the most sophisticated design of the three gates listed. It is considered a precision gate, which is usually tightly sealed against dust and dribble, and can be operated either

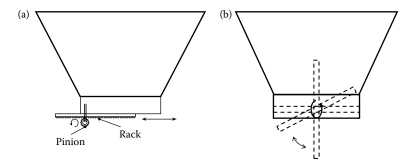


FIGURE 3.20
Gates for silos: (a) rack-and-pinion gate and (b) lever-operated quadrant gate.

manually (with minimum effort compared with the hand-slide gate), or adapted to the electric, pneumatic, or hydraulic mode. Figure 3.20 shows diagrams of the lever-operated quadrant gate and the rack-and-pinion gate.

Solids-level gauges, or controls, are used for warning about the level of solids within a bin or silo, or to protect conveyors from damage due to jamming if placed in transfer and discharge chutes. Solids-level controls can simply activate an audio or visual warning signal, or may be instrumentally adapted to conveying systems to start or stop automatically. Figure 3.21 illus-

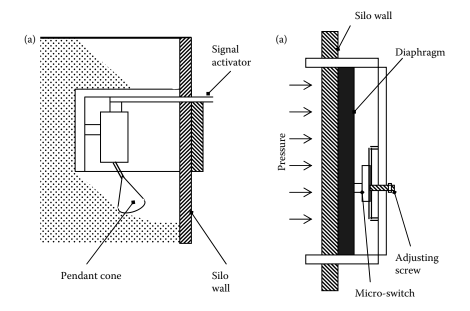


FIGURE 3.21Solids-level controls for silos: (a) pendant cone type and (b) diaphragm design.

trates two types of solids-level controls: the pendant cone level controller and the diaphragm level control unit.

3.4 Laboratory Exercise: Evaluation of Wall Loads in Silos

3.4.1 Introduction

Silo design has been a big concern since the end of the nineteenth century. The physical phenomenon arising from the interactions between confined solids and the walls of its storage structures is poorly understood. The starting point to design may be the calculation of pressure of silo walls at different depths in order to choose silo building materials, structural supports, and so on. The original theory of Janssen to estimate pressure on silo walls has been generally accepted and perfected over the years. This theory describes the friction between a layer of grain and the silo wall and establishes the static two-dimensional mathematical equilibrium of the vertical forces. From the established vertical forces, the vertical pressure, and thus the horizontal pressures, can be obtained. The objective of the experiment is to calculate pressure loads on a model grain silo at different heights as to corroborate the applicability of Janssen's equation.

3.4.2 Equipment and Materials

A small-scale metal silo (1–1.2 m height and 0.5–0.6 m diameter) can be used for the experiment. The material to be stored can be any kind of available grain, like wheat or barley.

3.4.3 Instruments and Apparatuses

Strain gauges or pressure cells and electronic wall pressure indicator.

3.4.4 Procedure

- Characterize the grain in terms of its main physical properties, such as bulk density and angle of repose.
- Fix the strain gauges at different (at least four) equally spaced distances along the height of the silo.
- Connect the gauges to the electronic pressure indicator.
- Fill up the silo to its complete height.
- Record the pressure at the different levels and at different times, for example, immediately after filling the silo, 1 h after filling, several hours after filling, and 24 h after filling.

3.4.5 Calculations and Report

- Make plots of recorded wall pressures at the different distances from bottom and at the different times.
- Calculate the wall pressures using the Janssen equation.
- Compare theoretical and experimental values.
- Perform a literature search to find out the result of measurements on similar materials stored in silos. Analyze and discuss these findings.
- Present a complete written report, according to scientific report guidelines either given by instructor or consulted in the literature. Include graphs, charts, and calculations made. Include, as an important part of the report, the discussion elaborated from the literature survey previously performed.

3.5 Bulk Solids Conveying

3.5.1 Introduction

Handling of particulate solids in processing industries is concerned with movement of materials in different cases such as from supply point to store or process, between stages during processes, or to packing and distribution. The movement of materials is a crucial activity which adds nothing to the value of the product, but can represent an added cost if not managed properly. For this reason, responsibility for material handling is normally vested in specialist handling engineers, and many manufacturing industries adopt this procedure. When a specific materials handling department is not provided, the responsibility for efficient handling of materials falls on the production manager and his/her staff. It is important, therefore, for production executives to have a sound knowledge of the fundamentals of good handling practice.

Broadly speaking, the many types of handling equipment comprise conveyors, elevators, cranes/hoists, trucks, and pneumatic systems. Motion of the material can be based on using gravity force or manual labor, as well as by employing power means. The method of motion provides a convenient basis for subdividing the types of conveyors into gravity or manually moved, such as chute, roller or skate-wheel conveyors; and power driven, such as roller, belt, slat, chain, vibratory, magnetic, screw, flight, and pneumatic conveyors.

The above categories include all the possibilities of materials handling within a processing plant, that is, either in bulk or packed. For bulk particulate or powdered materials, which fall within the scope of this book, a simpler and more convenient classification would comprise the following types of conveyors: belt, chain and screw, as well as pneumatic equipment. In the

literature all these forms of bulk material movement are known as conveyors and, thus, a proper categorization of the handling equipment for bulk particulate food solids would comprise the following groups: (a) belt conveyors; (b) chain conveyors: scraper conveyors, apron conveyors and bucket elevators; (c) screw conveyors; and (d) pneumatic conveyors: dense phase systems and dilute phase systems.

The above classification groups the different types of conveying systems in virtue of their operating principle. For example, regardless of whether a bulk material is being moved horizontally, inclined, or vertically, chain conveying can perform the duty based on the same principle of fixing an element (a paddle or a bucket) to a system of chains externally powered.

Bulk solids conveying represents one of those disciplines which have been neglected in terms of scientific research and development, when compared with other operations in particle technology directly related to processing, such as attrition or agglomeration. Conveying systems and supplies are normally provided by a large, capable manufacturing industry, which holds much of the engineering information in form of brochures, data sheets, and nomographs. Tables 3.3 and 3.4 present a general guide on conveying systems selection.

3.5.2 Belt Conveyors

The belt conveyor consists, essentially, of an endless belt operating between two or more pulleys, friction driven at one end and carried on an idler drum at the opposite end. Belt conveyors are used for movement of different types of bulk solids at distances varying from meters to kilometers and, thus, the belt and its load have to be supported on idlers on both conveying and return sections. Heavy-duty machines also have thick belts and these require larger diameter idler end drums. The construction of a belt conveying system requires a support structure, usually of steelwork. Belt conveyors have a high mechanical efficiency since, in larger installations, all the load is carried on antifriction bearings. Damage to the product in movement is slight because there is little or no relative motion, between the product being transported and the belt. The carrying capacity is high since relatively high speeds are possible when the whole system is operating. Bulk solid materials can be carried long distances mainly in horizontal plane, although there is a possibility of inclined conveying with certain limit to the angle of elevation. Inclination is usually limited to 15° or 20° and, in order to avoid excess spillage or "runback," special belts with corrugated sidewalks and lateral ribs are used for steeper inclination, up to a nominal 45°. A properly designed and maintained belt system has a long service life, but the initial cost is usually high, so installation is only advisable when amortization of the high initial cost can be assured.

The elements comprising a belt conveying system are: the belt, a drive system, some tension elements, the idlers, and appropriate loading and

TABLE 3.3Main Features of Different Conveying Systems

System		Diı	ecti	on		Free	que	ncy	1	Loca	tio	1	S	ervi	ce		Hei	ght		Ma	teri	al S	tate
	Vertical upward	Vertical downward	Inclined upward	Inclined downward	Horizontal	Continuous	Noncontinuous	Occasional	Spot	Path	Limited area	Unlimited area	Permanent	Temporary	Nondefined	Over head	Head level	Ground level	Underground	Packed	Bulk	Solid	Liquid
Conveyor			x	x	X	х			x	х			X				х	x	x	х	x	х	
Elevator	х	x				x		x	x				x			x				х	x	x	
Crane	x	x					x	x			x		x	x				x	x	x	x	x	
Truck					x		x			x		x			x			x		x	x	x	X
Pneumatic	X	x	x	x	x	x							X			X	x				x	X	X

TABLE 3.4Guide for Conveyor Selection

Function	Type of Conveyor
Horizontal transport of materials	Apron, belt, drag flight, screw, vibrating
Upward or downward inclined transport	Apron, belt, flight, forklift truck, screw, skip hoist
Elevation of materials	Bucket elevator, forklift truck, skip hoist, pneumatic
Combined path (horizontal and vertical)	Gravity-discharge bucket, pivoted bucket, pneumatic
Material distribution from bins and bunkers	Belt, flight, gravity discharge bucket, pneumatic, screw
Collecting materials from trucks and railroad cars	Car dumper, car shaker, mechanical shovel, pneumatic

discharge devices. Belts must be flexible enough to conform to the pulleys, wide enough to carry the quantity and type of material required, have sufficient strength to stand up under the expected load and operating tension, and posses a resistant surface. Stitched canvas, solid-woven balata, and rubber belts may be commonly used. Stitched canvas and woven belts are normally impregnated with a waterproofing material. A rubber belt is made of canvas or woven material impregnated and vulcanized with rubber and covered with a rubber sheet. Balata belts are similar to rubber belts as regards aging, but may be affected by temperatures over 50°C.

The drive should be at the discharge end of the belt and can be a conventional belt drive. The pulley must be large enough to provide proper contact surface with the belt to ensure a positive drive. Additional contact surface may be attained by using an idler pulley to provide more wrap contact of the belt. Pulley diameters must be large enough to keep from overflexing the belt. The take-up, which may be needed due to the stretch of the belt or to possible contraction and expansion because of changes in moisture and temperature, can be performed manually by adjusting screws or automatic by attaching a dead weight. Such adjustment can be carried out on the foot end pulley or on an idler pulley. The belt and its contents are supported by idler pulleys. Figure 3.22 illustrates the main types of idlers: troughed and flat. The straight pulley is used for the empty belt return and infrequently for carrying the load. It may be manufactured out of plain wood or light steel. Troughed idlers are normally the support of the whole belt, to increase the conveyor capacity or to confine powdered and particulate materials which would not stay on a flat belt.

Stretching of the belt can be controlled by using different types of tension devices. Two common designs are the gravity tensor and the sliding pulley illustrated in Figure 3.23. As previously mentioned, some conveyor belts incorporate an idler to snub back the belt under one pulley end to increase

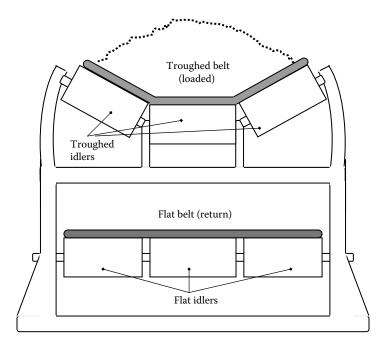


FIGURE 3.22 Idler pulleys for belt conveyors.

friction and minimize sliding of the belt over the pulley, and such arrangement can also be considered a tension device.

The material can be fed onto the belt by hand or by a mechanism providing continuous and steady flow. This may be a simple hopper with a gate valve or, if the material is not free flowing, coupled to an apron, screw, vibrating, or any other type of feeder. The material may be discharged over the end of the belt, by a diagonal scraper, by titling one or more of the idler pulleys, or by a tripper. As shown in Figure 3.24, a tripper consists of two idler pulleys mounted in such a way that cause the belt to take the shape of an S.

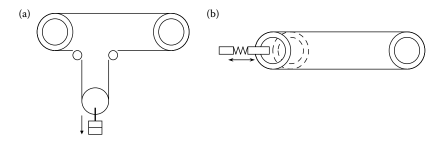


FIGURE 3.23 Tensors for belt conveyors: (a) gravity tensor and (b) sliding pulley.

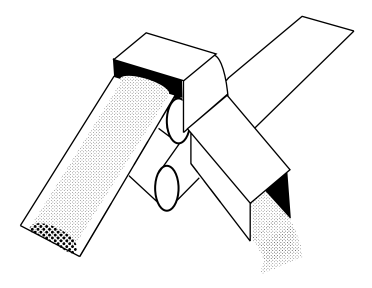


FIGURE 3.24 Tripper used for belt conveyor discharge.

The material is discharged over the top pulley and is caught by a chute which diverts it to either side of the belt, or to both. A short-belt conveyor, operating at right angles to the main belt, may replace the chute when it is required to move the material a considerable distance from the main belt. Table 3.5 lists power requirements for trippers. Discharge by tilting idlers may not be

TABLE 3.5Power for belt Conveyor Trippers

	Power Consumption (hp)				
t Width (cm)	Normal Idlers	Roller Idlers			
5	0.75	0.50			
5	1.00	0.75			
5	1.00	0.75			
7	1.50	1.25			
3	1.50	1.25			
)	1.75	1.25			
2	2.50	1.75			
1	3.00	2.50			
.7	4.00	3.00			
.0	5.00	3.25			
.0	6.00	5.00			
.4	7.00	6.00			
.0	6.00				

advisable since the material is discharged over a considerable length of the belt and there is an additional strain placed upon the belt due to twisting. An angle scraper is the simplest discharging device and provides satisfactory discharge for many materials. The various discharge systems are designed so that they may be controlled remotely.

A large belt conveyor system is an expensive an intricate facility which should be designed and installed under the supervision of specially trained conveyor engineers. During the last decades, the knowledge about belt conveyor design has been enriched and new design procedures have been proposed based on the capabilities offered by modern computer systems. Foote et al. (1988) examine the payoff of simulation and network analysis in conveyor system design, while Roberts (1994) presents an overview of technical and economical factors involved in conveyor design. Belt conveying design is far from standardized, and the following suggestions are only intended for preliminary approximations and calculations.

The horsepower required for movement of material by belt conveyors can be calculated by conventional engineering methods by considering the frictional resistance of the belt, the lift, and the frictional resistance of the different pulleys and trippers. However, the constants used in such procedure may vary with change in operating conditions. Also, flexing of the load and belt between supporting pulleys absorbs some energy. Power for standard installations can be consulted in tabulated data or, alternatively, calculated from the following equations:

Power to drive empty conveyor:

$$P = \frac{F(L + L_0)(0.06Wv)}{270}$$
 (3.17)

Power to convey material on level:

$$P = \frac{F(L + L_0)T}{270} \tag{3.18}$$

Power to lift material:

$$P = \frac{TH}{270} \tag{3.19}$$

Power to operate a tripper:

$$P = Yv + ZT (3.20)$$

where P is the power in horsepower, F is a friction factor taking values between 0.03 and 0.05, L is the conveyor length in meters, L_0 is a constant taking values between 30.5 and 45.7, W is the weight of all moving parts of the conveyor in kilograms per meter of overall length, V is the conveyor speed in V in the conveyor capacity in tons/V, and V is the height of lift in

Transmission Relation
1.875
1.500
1.250
1.125

TABLE 3.6Pulley Transmission Relations

meters. *Y* and *Z* are constants. The total power required is the sum of the powers calculated from Equations 3.17 through 3.20.

Belt conveyors have high initial capital so that their use may be well justified only for transportation at long distances, even in the range of kilometers. Consequently, their most common function is for handling heavy duty minerals, high tonnage rates and/or long distances, for example, coal, ore, and similar materials.

The belt has to have the appropriate tension prior to operating to prevent it from slipping over the pulleys. The loaded conveyor in operation is subjected to a combined tension comprising the initial tension and a tension equivalent to the consumed power. The latter can be considered a net tension and the total sum as the total tension. The initial tension depends on the drag due to the pulley arrangement while the net tension on the belt can be calculated as the relationship between the power in kg m/min and the belt speed in m/min. A variable known as the transmission relation can be defined between the total tension and the net tension. From this relation and the net tension, an overall tension on the belt can be found. This overall tension can be defined as a function of the belt width to determine the number of layers of the belt. A summary of all these criteria to calculate appropriate belt tension is presented in Tables 3.6 and 3.7.

The final part of the design and selection of conveying belts should consider characteristics of the solid, such as abrasiveness, moisture, temperature, stickiness, and adhesiveness. Testing should be performed for extreme,

TABLE 3.7 Appropriate Thickness of Belt

Belt Width (cm)	Number of Layers
30.5	3–4
45.7	3–5
61.0	4–7
76.2	5–8
91.4	6–9
106.7	6–10
122.0	7–12

rather than average conditions, to verify appropriate tension and other operating conditions of the conveyor.

Alternative designs to the conventional belt conveyors include steep-angle and vertical conveying systems with the surface worked into pockets formed by high sidewalls and scoop cleats, and hose-belt or tube conveyors in which the flat belt is shaped by means of concave idlers and guide rollers to overlap the belt edges (Bahke, 1991).

Example 3.3

Calculate the power to move a load of 850 tons/h of a granular material on a conveyor belt, 1 m wide and 100 m long. The conveyor operates at 150 m/min and its moving parts weight 5000 kg approximately. Constants F and L_0 have been evaluated as 0.04 and 38, respectively.

SOLUTION

The overall power will include the power to operate the empty conveyor and the power to move the load. Equation 3.17 is used to calculate the power of the empty conveyor while Equation 3.18 is used to calculate the power needed to actually convey the load. Constant 0.06 on those equations is expressed in ton min/kg h, and constant 270 is expressed in ton m/hp h. The remaining constants are dimensionless. Substituting values in Equation 3.17 for the empty conveyor:

$$P = \frac{0.04(100 + 38)[(0.06)(50)(150)]}{270} = 9.2 \text{ hp}$$

For the load, using Equation 3.18:

$$P = \frac{0.04(100 + 38)(850)}{270} = 17.4 \text{ hp}$$

The overall power will, therefore, be 26.6 hp.

3.5.3 Chain Conveyors

Compared to belt conveying systems, chain conveyors present a series of contrasting characteristics such as cheaper construction, noisy operation, slower movement, mechanic inefficiency, and no specialized skill required for their design. Chain conveyors are very versatile in design being more easily adaptable to different duties than belt conveyors. The main components of these types of conveying systems are the chains, the moving elements, and the drives. The types of chain available for conveying are extremely varied and a simple complete classification would be difficult to provide. Types of chain commonly used in chain conveyors are the malleable detachable, malleable pintle, steel, roller, and combination. The malleable detachable chain is, possibly, the most common and is normally used for light intermittent service.

(a)

Pintle chain, characterized by a pin which connects the links, is used for more rigorous service such as vertical grain elevators. Steel chain is used where high strength or good wearing qualities, or both, are needed. Roller chain is fitted with rollers or wheels to minimize friction and reduce wear. Combination chain is made in such a way that different features of the above three types are combined to provide certain definite performance characteristics. Chains of special alloys are available for operation in the presence of extreme or adverse factors such as heat, chemicals, abrasive substances, and so on. Some chain-link types are shown in Figure 3.25.

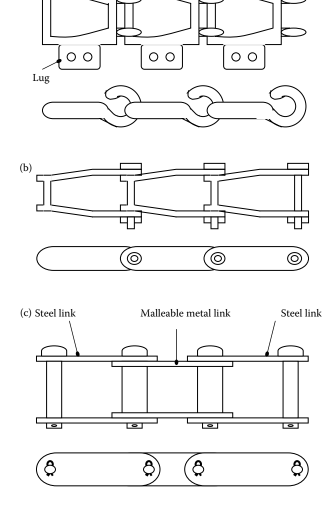


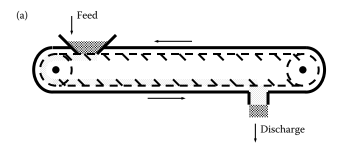
FIGURE 3.25
Links for chain conveyors: (a) malleable metal adapted with lugs, (b) steel, and (c) combined.

Depending upon how chain and moving elements are mounted for a specific duty, several types of chain conveyors can be obtained. Scraper conveyors will have the moving element, or flight, fitted perpendicular between two strands of roller chains, while apron conveyors will have the moving elements, or pans, overlapped and mounted on the same plane of roller chains forming a continuous "apron." When buckets are fixed to two strands of roller chains mounted on a steep-sloped or vertical structure, elevation of bulk solids can be possible in the so-called bucket elevators.

3.5.3.1 Scraper and Apron Conveyors

These conveying systems are used for granular, nonabrasive materials and are simple, cheap, easily constructed, and may operate at steep inclines. Power requirements are, however, high and wear may be excessive. Chain is made out of links, such as those illustrated in Figure 3.25, with lugs to which flights of various type are fastened. The links may be fitted with rollers for carrying the load and minimize frictional resistance. The simplest scraper conveyor is one made with "sawdust" chain, in which the links actually act as flights. As the name implies, it is used particularly for removing sawdust from sawing and milling operations, but it can be employed for many other industrial applications. Scraped conveyors with attached flights are designed in a variety of ways. The simplest of them is a single chain with flights operating in a wood or steel trough. Conveyors and elevators of this type are used extensively for moving the products of farming operations. Materials of large granular size are conveyed on the top of the conveyor. For small granular material, the lower flights are enclosed and the material is conveyed at this point. This allows greater capacities because the trough can operate full and at a higher speed, compared with topside movement. Also, the enclosed feature permits high-elevation angles. Discharge for both top and bottomrun conveying is usually at the head shaft. However, by putting gates in the lower trough, discharge for lower flight conveying can be made at any point of the conveyor. Similarly, top flight discharge can be facilitated by gates at the top trough and an open or skeleton lower trough.

Flight height, length, and spacing will depend on the expected duty of the conveyor. For small grains and other particulate materials, flat flights are recommended with a height of about 0.4 the flight length, and spaced at approximately one length. Flight speeds vary from 23 to 38 m/min. Low speeds should be used for materials of large granular size such as ear corn and walnuts. Small granular material, such as small grain and sesame seed, can be moved at higher speeds particularly if conveyed in the lower enclose portion of the conveyor. High speeds can damage the product and so, where practicable, capacity should be provided by large-size flights rather than by high speeds. The capacity of a scraper conveyor operating on the level can be assumed as 115% of the rectangular space between two flights when designed as suggested above. Figure 3.26 shows two scraper conveyor designs.



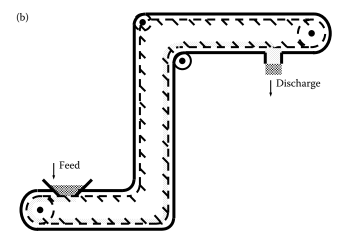


FIGURE 3.26 Scraper conveyors: (a) horizontal conveyor and (b) Z-type conveyor-elevator.

The main variables for selection and design of scraper conveyors are the capacity and the power. The capacity of horizontal scraper conveyors loaded with materials of approximately 800 kg/m^3 bulk density, has been calculated and is presented in Table 3.8. The theoretical horsepower requirement P for flight conveyors can be determined from the following equation:

$$P = \frac{[(2vL_cWF_c) + C(LF_m + H)]}{1400}$$
(3.21)

where v is the speed of conveyor in m/min, L_c is the horizontal projected length of conveyor in meters, W is the weight of flights and chain in kilograms per meter of overall length, F_c is the coefficient of friction for chain and flights, C is the capacity of material in kg/min, L is the horizontal projected length of loaded conveyor in meters, F_m is the coefficient of friction for the material, and H is the height of lift in meters.

The calculated horsepower must be adjusted to compensate for expected maximum capacity, starting friction, loss in the driving mechanization,

Capacity of Scraper Conve	y015-7-7-
Flight Size (cm)	Flight Weight (kg)
10×25.4	6.80
10×30.5	8.62
12.7×30.5	10.43
12.7×38.1	14.06
15.24×45.7	18.14
20.30×45.7	27.21
20.30×50.8	31.75
20.30×61.0	40.82
25.40×61.0	52.16

TABLE 3.8Capacity of Scraper Conveyors^{a,b,c}

^c Capacity of inclined conveyor compared with horizontal one:

Percent Capacity of Horizontal
77
55
33

variation in friction coefficients, type of power unit, and so on. The value of the coefficient F_c for chain dragging on metal can be taken as 0.33, while that for chain over wood can be approximated to 0.6. Friction coefficients for different materials are listed in Table 3.9.

For roller chains, F_c can be calculated from the following expression:

$$F_{\rm c} = X + \frac{d}{D} + \frac{2Y}{D} \tag{3.22}$$

TABLE 3.9Friction Coefficient for Various Materials Sliding on Metal

Material	$F_{ m m}$
Bituminous coal	0.590
Anthracite	0.330
Coke	0.355
Damp ashes	0.530
Dry sand	0.600
Limestone	0.585

 $^{^{\}rm a}~$ For $800~kg/m^3$ bulk density material and conveyor velocity of 30~m/min.

b For different density materials, capacity proportional to weight per cubic meter of material.

where X is a constant (0.33 for metal on nonlubricated metal, and 0.2 for metal on lubricated metal), d is the diameter of the rod on which the roller rotates, D the roller diameter, and Y is a constant whose value can be taken as 0.03 for general purposes.

For inclined conveyors, the power can be calculated from

$$P = WL_{c}(F_{m}\cos\alpha + \sin\alpha) + W_{1}L_{c}(F_{c}\cos\alpha + \sin\alpha) + WL_{c}(F_{m}\cos\alpha - \sin\alpha)$$
(3.23)

where α is the slope angle and W_1 is the load in kg per overall length of conveyor.

If the flights in the scraper conveyors are replaced with flat slats, steel plates or boards, a moving platform or apron is obtained and can be used for movement of sacked or large unit-sized materials. The typical design is a series of pans mounted between two strands of roller chain, with pans overlapping to eliminate dribble, and often equipped with end plates for deeper loads. Apron conveyors are not suitable for conveying of powdered and particulate materials. For such materials, their main application is feeding at controlled rates, with lump sizes large enough to minimize dribbling. Apron-feeder applications range from fairly light duties with light-gauge steel pans, up to extremely heavy duties requiring reinforced manganese steel pans with center supports. Capacity and power of apron conveyors can be found in tables and design graphs reported on text books (Badger and Banchero, 1955; Green and Perry, 2008).

3.5.3.2 Bucket Elevators

This type of conveying comprises high-capacity units primarily intended for bulk elevation of relatively free-flowing materials and may be considered a special adaptation of chain conveying. Bucket elevators are the simplest and most dependable equipment units for vertical lifting of different types of granular materials. They are available in a wide range of capacities and may operate either in the entire open or be totally enclosed. High efficiency in bucket elevators results from the absence of frictional loss from sliding of the material on the housing, and this feature distinguishes it from the vertical, or nearly vertical, scraper conveyor. The material carrying element of this sort of conveying is the bucket, which may be enclosed in a single housing called a leg, or two legs may be used. The return leg may be located some distance from the elevator leg. A single or double chain is used to attach the buckets. The more important considerations affecting the design and operation of bucket elevators are: (a) the physical properties of the conveyed material, (b) the shape and spacing of the buckets, (c) the speed at which the elevator is driven, (d) the method of loading the elevator, and (e) the method of discharging the elevator.

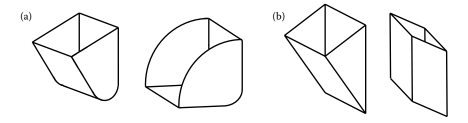


FIGURE 3.27 Buckets: (a) malleable metal for spaced bucket elevator and (b) steel for continuous bucket elevator.

Important physical properties of the material being elevated are particle size, lump size, moisture content, angle of repose, flowability, abrasiveness, friability, and so on. The design of the buckets has to do, principally, with capacity and ease of discharge. They may constructed out of malleable iron or steel and can be shaped with either sharp or round bottoms, as illustrated in Figure 3.27.

Mounting and spacing of the buckets will conform into a specific elevator design. Some typical bucket elevators are shown schematically in Figure 3.28. They may be fastened to the chain at the back (Figure 3.28a) or at the side if mounted in two chains (Figure 3.28b). Guides are sometimes used for two-chain installations, particularly in the up leg. Single-chain installations have no guides or supports between the head and foot wheels except for, possibly, an idler or two placed at strategic points to eliminate whip. The

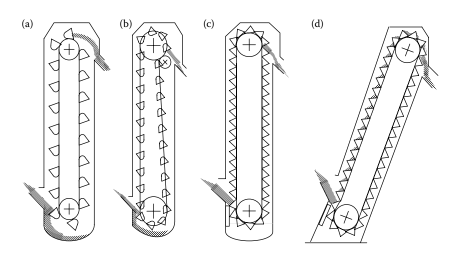


FIGURE 3.28Bucket elevators: (a) centrifugal-discharge-spaced buckets, (b) positive-discharge-spaced buckets, (c) continuous bucket, and (d) super capacity continuous bucket.

center spacing of buckets varies with their size, shape, speed, as well as head and foot wheel diameter. The buckets must be placed so that the centrifugal discharging grain does not hit the bucket ahead of the one discharging. For general purposes, the spacing will be from 2 to 3 times the projected width.

As shown diagrammatically in Figure 3.28, bucket elevators can be mainly loaded in three different ways. Spaced buckets receive part of the charge directly from a chute and part by scooping (Figure 3.28a), continuous buckets are filled as they pass through a loading leg with a feed spout above the tail wheel (Figure 3.28b) and, also, they can be loaded in a bottomless boot with a cleanout door (Figure 3.28c). In terms of discharge, spaced-bucket centrifugal-discharge elevators (Figure 3.28a) are the most common and are usually equipped with malleable-iron round-shaped buckets (Figure 3.27) spaced to prevent interference in loading or discharge. This type of elevator will handle almost any free-flowing fine or small-lump material. Speeds can be relatively high for fairly dense materials, but must be lowered considerably for aerated or low bulk-density materials to prevent fanning action. Spaced-bucket positive-discharge elevators (Figure 3.28b) are essentially the same as centrifugal-discharge units, except that the buckets are mounted on two strands of chain and are snubbed back under the head sprocket to invert them to allow positive discharge. These units are designed especially for materials which are sticky or tend to pack, and the slight impact of the chain seating on the snub sprocket, combined with complete bucket inversion, is generally sufficient to empty the buckets completely. Continuous-bucket elevators (Figure 3.28c) are generally used for larger-lump materials or for materials too difficult to handle with centrifugal-discharge elevators. Buckets used are of the steel type, such as those illustrated in Figure 3.27. They are closely spaced and the back of the preceding one serves as a discharge chute for the bucket which is dumping as it rounds the head pulley. Close bucket spacing reduces the speed at which the elevator operates to maintain capacities comparable with the spaced-bucket units. Gentle discharge is, therefore, promoted to help preventing excessive degradation and making this type of elevator suitable for handling finely pulverized or aerated materials. Super-capacity continuous-bucket elevators (Figure 3.28c) are designed for high lifts and large-lump material. They handle high tonnages and are usually operated at an incline to improve loading, as well as discharge conditions. Running speeds are low and, due to the heavy loads, bucket-supporting chain is usually guided on the elevating and return parts.

Bucket elevator horsepower for spaced buckets and digging boots can be calculated using the following equation:

$$P = \frac{TH}{152} \tag{3.24}$$

For continuous buckets with loading leg, the horsepower can be calculated by

$$P = \frac{TH}{167} \tag{3.25}$$

In Equations 3.24 and 3.25, *T* is the bucket capacity in tons/h and *H* is lift in meters. Both equations include normal drive losses, as well as loading pickup losses, and are applicable for vertical or slightly inclined lifts. For estimating purposes, tables and design graphs can be consulted in reference books, such as Green and Perry (2008).

3.5.4 Screw Conveyors

These systems are used to handle finely divided powders, damp materials, hot substances that may be chemically active, as well as granular materials of all types. Screw conveyors are also used for batch or continuous mixing, for feeding where a fairly accurate rate is required, and for conventional conveying, as well as elevating duties, particularly if the run is short. They operate on the principle of a rotating helical screw moving material in a trough or casing. Flights are made out of stainless steel, copper, brass, aluminum or cast iron, principally. They may be hard surfaced with Stellite or similar materials to resist highly abrasive materials. Although screw conveyors are simple and relatively inexpensive, power requirements are high and single sections are limited in length. The standard pitch screw has a pitch approximately equal to the diameter and is used on most horizontal installations and on inclines up to 20°. Half standard pitch screws may be used for inclines greater than 20°. Double-flight and triple-flight, variable-pitch and steepeddiameter screws are available for moving difficult materials and controlling feed rate. Ribbon screws are used for wet or sticky substances, while special cut flight and ribbon screws are used for mixing. Figure 3.29 shows main components of screw conveyors.

As previously mentioned, horizontal screw conveying is the predominant way of operation. Horizontal screw conveyors usually run in a U-shaped trough, with or without cover, depending upon the type of service and the characteristics of the material being moved. The screw is supported by brackets at different standard spacing. For elevating at a steep incline, a cylindrical housing would be required. The tube normally operates full with no brackets between the ends, since they would interfere with the movement of material. The material supports and guides the flight, so that satisfactory operation results if the conveyor is kept full. The use of hollow screws and pipes for circulating hot or cold fluids would permit that screw conveyors may be used for heating, cooling, and drying operations. It is possible to seal a screw conveyor from the outside atmosphere in order to operate outdoors without special protection. The conveyor can be completely sealed to operate in its own atmosphere at positive or negative pressure. The casing can also be

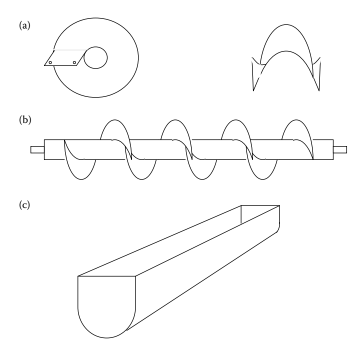


FIGURE 3.29 Screw conveyor components: (a) flight, (b) screw formed by mounting flights on an axle, and (c) trough.

insulated to maintain internal temperatures in areas or extreme ambient temperatures. The casing can be designed with a drop bottom for easy cleaning to avoid contamination when different materials are to be conveyed through the same system.

Screw conveyors are usually made up of standard sections coupled together, and so special attention should be given to bending stresses in the couplings. Hanger bearings supporting the flights can obstruct the flow of material when the trough is loaded above its level. Thus, with difficult materials, the load in the trough must be kept below this level. Alternatively, special hanger bearings which minimize obstruction should be selected. Since screw conveyors operate at relatively low rotational speeds, the fact that the outer edge of the flight may be moving at a relatively high linear speed is often neglected. This may create a wear problem, and if wear is too severe it can be reduced by the use of hard-surfaced edges, detachable hardened flight segments, rubber covering, or high-carbon steels. Screw conveyors are well suited as feeders or metering devices under bins or hoppers. The portion of the screw under the container is usually designed with half or third pitch, the balance of the screw full pitch. Thus, the main portion of the screw operates half or third full, while the metering portion of the screw operates full.

Concise data and formulae are normally not available for individual design problems and it is recommended to consult specialized engineers when designing and installing large screw conveying systems. Data that could be available to assist in selection and design are normally empirical in nature. Roberts (1999, 2000), presents analytical data to predict the performance of screw conveyors. The power requirement of a screw conveyor is a function of its length, elevation, type of hanger brackets, type of flights, the viscosity or internal resistance of the material, the coefficient of friction of the material on the flights and housing, and the weight of the material. Consideration must also be given to additional power needed to start a full screw, to free a jammed screw, or to operate with material which has a tendency to stick to the trough sides. The horsepower *P* required to drive a screw conveyor depends on the dimensions of the system and the characteristics of the material. A rough approximation for normal horizontal operation can be determined from the following relation:

$$P = \frac{CL\rho_b F}{4500} \tag{3.26}$$

where C is the capacity in m³/min, L is the conveyor length in meters, ρ_b is the apparent density of material in kg/m³, F is a factor depending on the type of material, as appearing in Table 3.10. If horsepower is less than 1, it should be doubled; if it ranges from 1 to 2, it should be multiplied by 1.5; if it ranges from 2 to 4, it should be multiplied by 1.25; and if it ranges from 4 to 5, it should be multiplied by 1.1. No correction is necessary for values above 5 hp.

Example 3.4

A measure of 200 ton/h of limestone is to be transported to a processing plant 4 m above its storing place. Two options need to be contemplated to move the material: a belt conveyor or a bucket elevator. The maximum slope for the belt conveyor is said to be 15° , its moving parts weight 1550 kg, and at maximum capacity the velocity is 24 m/min. For the belt conveyor, minimum values of the constants F and L_0 apply. Demonstrate which conveying system would suit best for the duty of transporting the limestone.

SOLUTION

The obvious choice of system would be the one consuming less power. For the belt conveyor, it would be necessary to calculate the power to operate it empty and, then, add the power to overcome the lifting of the material. The length of the belt can be calculated considering the following diagram:



Type a (F = 1.2) Light, Fine, Nonabrasive, Free-Flowing Materials ρ_b : 480 to 640 kg/m³	Type b ($F = 1.4-1.8$) Nonabrasive, Granular or Fines Mixed with Lumps ρ_b : up to 830 kg/m ³	Type c ($F = 2.0-2.5$) Non and Mildly Abrasive, Granular or Fines Mixed with Lumps ρ_b : $640-1200 \text{ kg/m}^3$	Type d ($F = 3.0$ –4.0) Mildly Abrasive or Abrasive, Fine, Granular or Fines with Lumps ρ_b : 830–1600 kg/m³
Barley	Cacao seeds	Borax	Bauxite
Corn flour	Coffee seeds	Brown coal	Bone meal
Cotton seed flour	Corn	Charcoal	Cement
Crushed coal	Corn meal	Cocoa	Chalk
Crushed limestone	Fine alum	Dehydrated milk	Cinders coal
Granular dried malt	Flakes of graphite	Fly ash	Clay
Malt	Hydrated lime	Granular moist malt	Dry sand
Rice	Jelly granules	Granulated cork	Feldspar
Wheat	Soy meal	Starch	Ground slate
Wheat flour		Icing sugar	Pebble lime
		Lumpy alum	Powdered gypsum
		Paper pulp	Powdered limestone
		Salt	Powdered sulfur
		Soap powder	Raw sugar
		Soda ash	Zinc oxide

TABLE 3.10Material Factors for Horizontal Screw Conveyors

By using simple trigonometry the length of the belt corresponds to a side of a rectangle triangle:

Soft coal

length =
$$\frac{4}{\sin 15^\circ}$$
 = $\frac{4}{0.2588}$ ~ 15.5 m

Using the belt length, the weight of the moving pieces of the conveyor are determined as 100 kg (i.e., 1550/15.5). Taking the minimum values of F and L_0 , as indicated above, substituting into Equation 3.17:

$$P = \frac{0.03(15.5 + 30.5)[(0.06)(100)(24)]}{270} = 0.736 \text{ hp}$$

To calculate the power for lifting the material, Equation 3.19 can be used as follows:

$$P = \frac{200(4)}{270} = 2.963 \text{ hp}$$

The total power to transport the material using a belt conveyor is, thus:

$$P_{\text{total}} = 0.736 + 2.963 = 3.699 \text{ hp}$$

For the bucket elevator, power for lifting of the material by spaced and continuous buckets respectively, are calculated by Equations 3.23 and 3.24 as

$$P = \frac{(200)(4)}{152} = 5.26 \text{ hp}$$

$$P = \frac{(200)(4)}{167} = 4.79 \,\mathrm{hp}$$

The best option is, therefore, the use of the belt conveyor, since it would consume less power that either of the two bucket elevator systems.

Example 3.5

A screw conveyor transports seeds of 500 kg/m³ in bulk density and consumes 0.45 hp in power. Calculate the capacity, in tons/h, if the screw is 3.5 m long.

SOLUTION

The capacity can be transposed from Equation 3.25 to give

$$C = \frac{P(4500)}{L\rho_b F}$$

The annotation in Equation 3.25 states that the power is to be multiplied by 2 when less than unity. The power consumed (0.45 hp) should, therefore, be divided by two to substitute in the equation above. The constant F can be consulted in Table 3.10 in which, according to the density of the seeds, the material would be Class a, and F would equal 1.2. The units of numeric constant in Equation 3.25 are kg m/hp min. Substituting values into the equation above:

$$C = \frac{(0.225)(4500)}{(3.5)(500)(1.2)} = 0.4821 \,\mathrm{m}^3/\mathrm{min}$$

Transforming using the density of the material and the equivalence between minutes and hours, the capacity is 14.46 ton/h.

3.5.5 Pneumatic Conveying

3.5.5.1 Introduction: Types of Conveyors

One of the most important bulk solids handling techniques in a solids handling plant is the movement of material suspended in a stream of air over horizontal, inclined or vertical surfaces, ranging from a few to several

hundred meters. This type of conveying is one of the most versatile, handling materials that range from fine powders through 6.35-mm pellets and bulk densities of 16 to more than 3200 kg/m³. As compared with previously discussed methods, pneumatic conveying offers the containment and flexibility of pipeline transport for bulk solids that, otherwise, will be exposed to direct contact with moving mechanical parts. One of the earliest recorded uses of pneumatic conveying (Reed and Bradley, 1991) was for unloading wheat from barges to flour mills at the end of the nineteenth century in London. Some other grains, as well as different cargo such as alumina, cement, and plastic resins are still unloaded using the same basic methods. Other common applications include unloading trucks, railcars, and barges, transferring materials to and from storage vessels, injecting solids into reactors and combustion chambers, and collecting fugitive dust by vacuum. Advantages in cleanliness and low contamination are reflected in the fact that if properly constructed and maintained, pneumatic systems can be virtually dust free. Vacuum systems offer the advantage that leakage is into the pipeline, so that even damaged or leaky operations do not result in product loss. Sealed systems can prevent most forms of contamination, and contact with moving mechanical components is minimal. Pressure systems prevent inward leakage and can use dry, inert gas for conveying in order to exclude oxygen and moisture to handle very sensitive biological materials.

Pneumatic conveying has some drawbacks such as high power consumption, limitations in overall distance and capacity, as well as severe wear of equipment and attrition or degradation of the materials being conveyed. In terms of consumption, taken on the basis of cost per unit weight per unit distance conveyed, pneumatic conveying is by far the most expensive method of moving materials. In many practical cases, however, higher cost is justified, because the alternatives are not practical due to layout limitations, containment and cleanliness, as well as low maintenance of this way of conveying. With regard to distance and capacity, pneumatic conveyors are generally restricted to about 300 tons/h and 1000 m, although not simultaneously. Pertaining wear and attrition, such problems normally arise only when systems are not properly designed and operated.

Despite the above-mentioned limitations, pneumatic conveying is an important and practical form of transporting bulk solids, with applications in numerous processing industries. The range of materials that can be handled is nearly unlimited, including powders and particulates of different types. The limitations on what can be conveyed depend more upon the physical nature of the material that on its generic classification. Particle size, hardness, resistance to damage, and cohesive properties are key factors in determining whether a material is suitable for this sort of conveying. Cohesive or sticky materials are often difficult to handle in a pneumatic conveyor. Moist substances that are wet enough to stick to the pipeline walls usually cannot be conveyed successfully. Materials with high oil or fat contents can also cause severe buildup in pipelines making conveying not practical.

Pneumatic conveying systems can be categorized in a number of ways depending on their function, as well as type and magnitude of operating pressure. The solids loading is a useful criterion to classify pneumatic conveyors, which can run over a wide range of conditions bounded on one end by gas alone with no entrained solids, and at the other end by a completely full pipe where the solids are plugging the line. Most industrial conveying systems operate somewhere in between these two extremes, being ranked broadly as either dense-phase or dilute-phase systems, depending upon the relative solids loading and velocity of the system. This is best described in a general state diagram, which is a plot of pressure per unit length of pipe as a function of conveying gas velocity, with constant solids flow rate. At higher velocities particles are generally suspended in the gas with low solids loading ratio, typically below 15, and termed as dilute-phase conveying. If the gas velocity is slowly decreased, the pressure required to convey a constant amount of solids also drops. After reaching a minimum, a further reduction in gas velocity results in an increase in pressure, as particles begin to fall out of suspension and interparticle collision increase. This region, with a solids loading ratio typically higher than 15 and the gas velocity below the saltation velocity, is that of dense-phase conveying. With many materials is difficult to establish a definite boundary separating dense-phase and dilute-phase regions, and conveying can occur over a continuous range from fully suspended to a slow moving bed. With other materials very distinct regions are observed and the conveying progresses in either a very stable or unstable way.

Dense-phase conveying, also termed "nonsuspension" conveying, is normally used to discharge particulate solids or to move materials over short distances. There are several types of equipment such as plug-phase conveyors, fluidized systems, blow tanks, and, more innovative, long-distance systems. Dilute-phase, or dispersed-phase conveyors, are more versatile in use and can be considered the typical pneumatic conveying systems as described in the literature. The most accepted classification of dilute-phase conveyors comprises: pressure, vacuum, combined, and closed-loop systems.

3.5.5.2 Dense-Phase Systems

The development of dense-phase conveying technology has broadened significantly the potential for using these systems to cover longer moving distances. Compared with conventional dilute-phase transport, these relatively high-pressure systems can be designed safely to operate at reduced conveying velocities which, in turn, lead to lower pipeline wear, smaller pipe sizes, and lower operation costs. Several types of dense-phase conveying have been identified as plug-phase conveyors, fluidized systems, blow tanks, and long-distance systems.

Plug-phase conveying basically involves the transportation of a limited batch of material per conveying cycle. The device functions by introducing pressurized air on top of a head of material contained in the pressure vessel. If the material is free flowing, it will flow through a valve and move into a short conveying line. When using this system, care should be taken to avoid surges of air, caused either by the tank emptying or by the air breaking through the product. This type of conveying system can be used to transport granular materials such as grains and other food particulates over relatively short distances, for example, up to 200 m. The method of air injection can have a significant impact on the overall performance and efficiency of the system. Compared with conventional conveying systems, both dense-phase and dilute-phase, plug-phase conveying offers some advantages such as use of lower average conveying velocities, and minimization of system erosion.

Particulate solids, contained in silos or bulk transportation vehicles, may be fluidized to assist discharge using a dense-phase pneumatic fluidizes system. Other applications are conveying of prefluidized, finely divided non free-flowing materials over short distances, such as from storage to the entrance of a main conveying system. In fluidized systems, air is introduced in sufficient quantity and at sufficient pressure to produce gravity flow of the material. A diagram of a fluidized system is given in Figure 3.30. Fluidizing is accomplished by means of a chamber in which air is passed through a bed, upon which the material to be conveyed rests. The beds used consist of porous ceramic tiles, sintered metal or plastic, fine wire-mesh, and similar materials. As air passes through the porous bed, each particle is surrounded

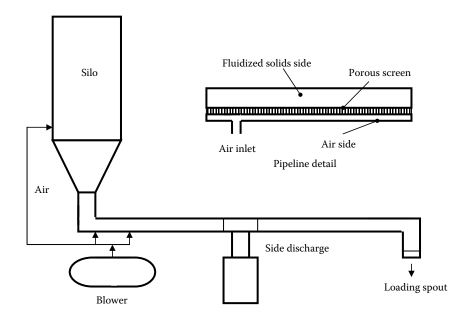


FIGURE 3.30 Dense-phase fluidized conveying system.

by a film of air. At the point of incipient fluidization, the material starts behaving as a free-flow powder or particulate. Fluidized solids may be forced through pipelines by air under pressure passed through a perforated flexible inner tube running the length of the pipeline. The fluidized material is then conveyed by establishing a pressure drop along the pipeline. Fluidized systems are the most reliable and economical method of conveying certain powders and bulk solids over distances ranging from few meters up to about 1 km. They take advantage of the fluidization and air-retention properties of the bulk solid (Mainwaring and Reed, 1987). The suitability of a given material to this mode of conveying, and the best type of feeder for these systems should be determined by test work.

A blow tank is a type of slow-motion, dense-phase pneumatic conveying that allows friable products to be conveyed with extremely low levels of product degradation or damage, because solids conveying velocities usually occur in the range of 0.25–2 m/s. Such features are very important for a wide variety of friable products, especially for those prone to attrition. Low-velocity systems operate at relatively high conveying pressures, for example, 200–500 kPa, and consequently must have reliable hardware. Also, due to the extremely high levels of concentration which occur during conveying, the subsequent operation conditions depend quite strongly on the nature and physical properties of the material being transported. It is important, for these reasons, that full-scale tests be exercised prior to the selection or design of such equipment.

One of the most interesting and innovative applications of dense-phase pneumatic conveying is the long-distance transportation. Conveying of fine and coarse materials over distances up to 3 or 4 km is possible with this system. This can be achieved by tandem arrangement of blow tanks. Longdistance systems operate at high pressures and particular attention must be given to the control of solids feeding into the pipeline. A manner of doing this is by using a cone-dosing system, in which the flow of solids is controlled by the movement of a double cone located near the blow tank outlet. The cone moves continuously up and down inside the vessel, but the stroke is adjusted by a proportional-integral controller, based on a conveying line back-pressure measuring signal and set point. The higher the conveying or operating pressure, the lower the cone set point inside the vessel. The flow of solids into the conveying pipeline is restricted on this manner, until a satisfactory conveying pressure is restored. Furthermore, the cone dosing system provides an additional and useful feature of mechanical agitation, as well as flow assistance of cohesive materials.

3.5.5.3 Dilute-Phase Systems

As stated earlier, dilute-phase conveying is the commonly employed method for transporting a wide variety of suspended solids using air flowing axially along a pipeline. The method is mainly characterized by the low solids to air ratio and by the fact that air and solids flow as a two-phase system inside a pipeline. The main components of dense-phase pneumatic conveying systems basically include gas movers, solids feeders, pipelines, and solids separators. Their location within the system may vary from type to type but their functions are the same. The gas mover provides an appropriate flow rate of gas required for the transport at the right velocity and pressure. The solids feeder introduces the solids particles at a controlled rate into the pipeline where they are mixed with the conveying gas. The pipeline consists of straight sections, both horizontal and vertical, connected together with bends. At the end of the trajectory, in any type of solid–gas separator, the solids are decelerated and recovered from the gas stream to be stored in a silo or fed into another unit. The gas, which is typically air, is generally released into the atmosphere. Secondary components comprise controls, safety equipment, and instrumentation.

By far, the most common device for moving gas in a pneumatic conveying system is the Roots-type, rotary lobe blower. Such blower is preferred since it provides an economical source of gas flow that meets the pressure or vacuum and flow requirements for the largest category of systems. An important feature of this type of blower is that it delivers a nearly constant volume over its operating pressure range. This is important because control of the gas in pneumatic conveying system is critical for stable operation. Singlestage blowers generate up to 200 kPa while double-stage blowers generate up to about 270 kPa. For low-pressure conveyors, centrifugal fans can be used as gas movers. High-speed, single-stage centrifugal fans deliver air at about 120 kPa and must be supplied with clean air if excessive abrasion of the light impellers is to be avoided. The slower-running, multistage, heavy-duty centrifugal fan can deliver up to 140 kPa and, being slower running, is relatively resistant to particle abrasion. Compressors deliver air at high pressures but require lubrication being, thus, potential contaminants of the solids in movement. They provide pressures up to 800 kPa so, with certain measures, they find applications in high-pressure systems, as well as in dense-phase conveying.

For proper operation of a dilute-phase, pneumatic conveying system the solids fed into the pipeline must be carefully controlled. Two common types of feeders are the venturi feeder and the rotary valve or star feeder, illustrated in Figure 3.31. The venturi feeder is only suitable for low-pressure systems. The rotary valve feeder, also known as star feeder as described previously in this chapter is widely used for feeding medium-pressure conveyors. This feeder is efficient and simple in principle, but requires careful design in order to minimize air leakage. Excessive air loss from feeders, wastes power, causes dust, and leads to system instability. Rotary feeders work well on free-flowing, nonabrasive powders and special types are available for more difficult materials.

The number and type of bends used in the pipeline may constitute a critical design problem. The difficulties that may arise by the use of bends are

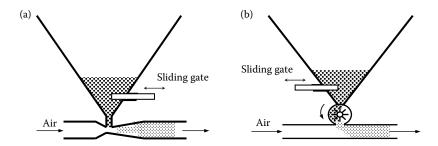


FIGURE 3.31 Feeders for dilute-phase conveying systems: (a) venture feeder and (b) star feeder.

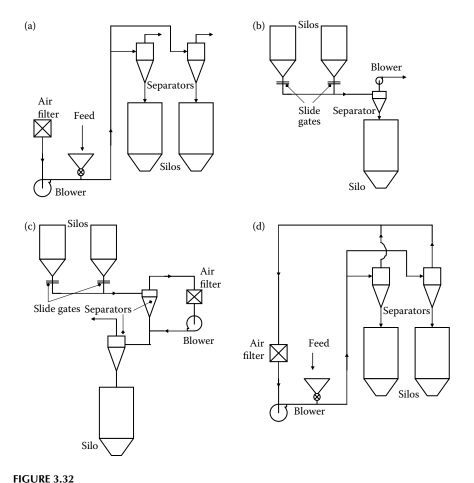
abrasive wear, product degradation, product buildup, and pressure loss. The latter is the problem primarily related to operation. In general terms, all the bend associated drawbacks are minimized by use of smooth, long-radius bends instead of short radius and blind tees types of bends. The system layout is also important. It is often more troublesome to have many bends in short distances than few bends in long distances. In the former case, the contribution of each bend will be more significant than in the latter. However, for systems handling nonabrasive, nondegradable materials, the number and type of elbow is not very critical and selection or design should probably be based on cost.

Separating the solids from the airstream when conveying concludes can be accomplished in a number of ways. For most of the cases, a single technique is insufficient to separate the conveyed material completely and satisfactory. The most common approach consists in using a cyclone followed by a bag filter. Both separating techniques will be discussed in a later chapter. The selection of a gas—solids separating system should be based on the material characteristics, degree of separation required, environmental regulations, concentration of solids, and cost.

Operating variables of dilute-phase conveying systems comprise gas (air) velocity, pressure drop, gas-solids ratio, and particle properties. Pertaining air velocity, most particulate materials may be conveyed satisfactorily at air speeds within the range of 15–25 m/s. Velocities above this may cause abrasion of tube bends and product damage may represent difficulties. At extremely low speeds, solids tend to settle out and block horizontal pipe runs. In terms of pressure drop, if air at high pressure is used, its correspondingly high initial energy will enable more conveying to be accomplished, per kilogram of air, than if low-pressure air is used. High-pressure systems are, however, proportionately more expensive than low-pressure ones. The maximum pressure recommended for general-purpose, dilute-phase conveying of solid particulate materials is about 170 kPa. With regard to solid–air ratio, for maximum efficiency this should be as high as possible, but without invading the range of dense-phase conveying. There is an upper

limit for this ratio for specific materials; exceeding it will cause blockage of the system due to saltation. Finally, material properties such as size, shape, density, and surface properties need to be carefully considered in operation and selection of dilute-phase conveying systems. Other important properties are friability, hygroscopicity, as well as susceptibility to impact, abrasion damage or oxidation.

Figure 3.32 shows the four main types of dilute-phase conveying systems previously mentioned. The pressure system, also called positive-pressure, or push system operates at super-atmospheric pressure and is used for delivery to several outlets from one inlet (Figure 3.32a). Although most applications of these systems lie within the scope of dilute-phase conveying, under certain arrangements they can also operate as high-pressure, dense-phase



Dilute-phase pneumatic conveying systems: (a) pressure system, (b) vacuum system, (c) combined system, and (d) closed-loop system.

conveyors. In general, pressure systems can hold higher capacities and longer conveying distances than negative-pressure systems. The vacuum, negativepressure or pull system, works at subatmospheric pressure and is used for delivery to one outlet from several inlets (Figure 3.32b). Vacuum systems are usually limited to shorter distances than positive ones and are more restrained to operate with dilute, low solids loading than pressure systems. When both features of pressure and vacuum systems are combined in a unit, the advantages of each can be exploited (Figure 3.32c). This arrangement consists of two sections: a pull/push system with a negative-pressure front end, followed by a positive-pressure loop. The benefit is that they capitalize on the ease of feeding into a vacuum and combine this with the higher capacity and longer conveying distance when using positive pressure. Recirculation of the conveying air, as in the closed-loop system (Figure 3.32), reduces contamination of product by the air and limits product dehydration. However, such systems are often difficult to control and an intercooler may be required to prevent the pump overheating the recirculated air.

3.5.5.4 Design and Selection of Dilute-Phase Systems

In contrast with those conveying methods previously discussed, pneumatic conveying can be perfectly identified as a case of two-phase flow, which is a topic well covered by fluid mechanics. Single-phase flow within a pipe can be described by basic equations that are considered the basic fundamentals to start calculations in order to design pneumatic conveying systems. However, while single-phase flow in a pipe is well understood, adding solids into the moving stream in the so-called two-phase flow complicates matters significantly. As solid particles are introduced in a moving stream of gas, the pressure drop in the line begins to increase because momentum is transferred to the particles to accelerate them to the conveying velocity. The total pressure drop consists of two components: the one due to the gas flow alone, and that required for transporting the particles. Apart from the gas velocity, some other variables such as the amount of solids in the pipeline and the velocity of the solids relative to the gas also affect the pressure drop.

The moving gas stream applies drag and lift to the particles. For particles to be conveyed in such gas stream, the velocity of the gas must be sufficiently high to stop particles settling out. In flow through horizontal pipes the minimum air velocity to stop particles settling to the bottom of the pipe is called the saltation velocity. The equivalent velocity for flow through vertical pipes is known as the choking velocity. The saltation velocity is a function of the density of the gas and the solids, as well as particle and pipeline diameter (Cabrejos and Klinzing, 1994). There is also a direct relationship between the saltation velocity and the solids loading ratio. Generally, saltation occurs at higher velocities when the solids loading ratio is also high.

In terms of designing equipment for pneumatic conveying, there is another type of velocity, that is, the minimum conveying velocity, used to describe the correlation of gas velocity to the behavior of solid particles inside a pipeline. This velocity is the lowest one necessary to prevent plugging the line in a given system for a given material. Some researchers have suggested the use of the saltation velocity with a safety factor, while others have developed empirical correlations. Some of these correlations, however, often predict widely differing velocities for the same set of conditions (Wypych, 1999).

The saltation velocity can be calculated based on the procedure presented by Zenz (1964), as follows:

Calculate the parameter x_p^* by the equation:

$$x_{\rm p}^* = \frac{x_{\rm p}}{\left[3(\mu_{\rm g})^2/4g\rho_{\rm g}(\rho_{\rm s} - \rho_{\rm g})\right]^{1/3}}$$
(3.27)

where x_p is the particle diameter, μ_g is the gas viscosity, ρ_g and ρ_s are the gas and solid densities, respectively. If there is a mixture of particle sizes, that is, a particle size distribution which is quite common in most industrial powders, calculate x_p^* for the largest and smallest particle.

From Figure 3.33 calculate u_{ss}^* from x_p^* and use the following relation to determine the minimum conveying velocity u_{ss} for a single particle:

$$u_{\rm ss} = 0.19(u_{\rm ss}^*) \left[\frac{4g\mu_{\rm g}(\rho_{\rm s} - \rho_{\rm g})}{3(\rho_{\rm g})^2} \right]^{1/3} (D_{\rm t})^{0.4}$$
 (3.28)

where $D_{\rm t}$ is the pipe diameter in mm, but taken as dimensionless in the above equation.

Estimate n, the gradient of the curve in Figure 3.33 at x_p^* . For a particle size distribution n is the slope of the line joining the values of u_{ss}^* for the largest and smallest particle.

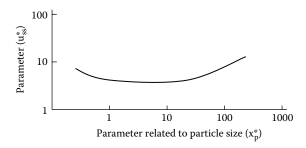


FIGURE 3.33 Minimum conveying velocity for a single particle for a pipe diameter D_t of 63.5 mm. (Adapted from Zenz, F. A. 1964. *Industrial and Engineering Chemistry Fundamentals* 3: 65–75.)

Transpose u_s , the saltation velocity from the following equations:

$$\frac{G_{\rm s}}{\rho_{\rm s}} = \frac{0.214 \, (n)^{1.5} (u_{\rm s} - u_{\rm ss})}{u_{\rm ss}} \tag{3.29}$$

for n > 0.068, or

$$\frac{G_{\rm s}}{\rho_{\rm s}} = \frac{0.0032 \left(u_{\rm s} - u_{\rm ss}\right)}{u_{\rm ss}} \tag{3.30}$$

for -0.11 < n < 0.068.

In Equations 3.28 and 3.29, G_s is the flux of solids. For particle size distributions, the largest particle size should be used in these equations in order to avoid underestimation of the saltation velocity.

Pressure drop is another key variable involved in design and operation of pneumatic conveying systems. Pressure drop requirement is closely associated with gas flow and solids rate. The same amount of solids can be conveyed in a line using a number of velocity and pressure drop combinations. Pressure drop in pneumatic conveying is determined by calculating potential energy, kinetic energy, and frictional energy changes in horizontal and vertical sections, as well as in bends in the conveying line separately. The estimation of the overall pressure drop results from summing up all the individual contributions, depending on the specific layout of a pneumatic conveying system. The detailed procedure is as follows:

Take the initial air velocity as 10% above the saltation velocity, to minimize risk of settling of coarse particles.

In horizontal sections the potential energy loss is zero, the kinetic energy gain $E_{\bf k}$ is given by

$$E_{\rm k} = G_{\rm s} u_{\rm p} \tag{3.31}$$

where G_s is the flux of solids and u_p is the particle velocity.

If particle has accelerated to its maximum value, the particle velocity is defined by

$$u_{\rm p} = u_0 - u_{\rm t} \tag{3.32}$$

where u_0 is the air velocity and u_t is the terminal settling velocity of the particle.

The behavior of particles settling in a fluid is well understood, and the settling velocity depends on the flow regime, that is,

$$u_{\rm t} = \frac{x_{\rm p}^2 (\rho_{\rm s} - \rho_{\rm g}) g}{18\mu_{\rm o}}$$
 (3.33)

for Re_p < 0.4, which is the well-known Stokes' law. Also:

$$u_{\rm t} = \left[\frac{4(\rho_{\rm s} - \rho_{\rm g})^2 g^2}{225\rho_{\rm g}\mu_{\rm g}} \right]^{1/3} (x_{\rm p})$$
 (3.34)

for $Re_p < 500$. And, finally:

$$u_{\rm t} = \left[\frac{3.1 x_{\rm p} (\rho_{\rm s} - \rho_{\rm g}) g}{\rho_{\rm g}} \right]^{1/2}$$
 (3.35)

for $Re_p < 2 \times 10^5$.

In Equations 3.32 through 3.34, Re_p refers to the particle Reynolds number, which is defined by

$$Re_{p} = \frac{x_{p}\rho_{g}u_{t}}{\mu_{g}}$$
 (3.36)

and all the variables have been previously defined.

The frictional energy loss due to the gas flow E_{fg} is given by

$$E_{\rm fg} = \frac{2f_{\rm g}\rho_{\rm g}u_0^2L}{D} \tag{3.37}$$

where $f_{\rm g}$ is a gas friction factor, L is the pipe length, and D is the pipe diameter. The gas friction factor $f_{\rm g}$, adopts values according to the system Reynolds number Re, that is,

$$f_{\rm g} = 0.0791(\text{Re})^{-0.25}$$
 (3.38)

for $3 \times 10^3 < \text{Re} < 10^5$. And

$$f_{\rm g} = 0.0008 + 0.0552(\text{Re})^{-0.237}$$
 (3.39)

for $10^5 < \text{Re} < 10^8$.

As indicated above, the Reynolds number in Equations 3.37 and 3.38 is that for the system, that is,

$$Re = \frac{D\rho_{\rm g}u_0}{\mu_{\rm g}} \tag{3.40}$$

The frictional energy loss due to the particle flow $E_{\rm fp}$ can be estimated by

$$E_{\rm fp} = \left(\frac{\pi}{8}\right) \left(\frac{f_{\rm p}}{f_{\rm g}}\right) \left(\frac{\rho_{\rm s}}{\rho_{\rm g}}\right)^{1/2} \left(\frac{G_{\rm s}}{G}\right) E_{\rm fg} \tag{3.41}$$

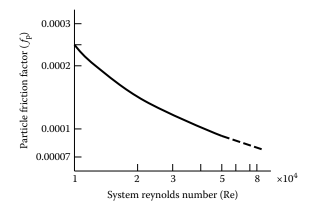


FIGURE 3.34Particle friction factor as a function of system Reynolds number, Re. (Adapted from Rose, H. E. and Barnacle, H. F. 1957. *Engineer, London* 203: 898, 939.)

where G is the flux of air and f_p is the particle friction factor, which can be determined graphically, as a function of Re, from Figure 3.34.

The frictional loss in flow round bends $E_{\rm fb}$ can be calculated from:

$$E_{\rm fb} = 2f_{\rm b} \left(\frac{G_{\rm s}}{G}\right) \left(\frac{u_0^3}{u_{\rm p}}\right) \rho_{\rm g} \tag{3.42}$$

where f_b equals 0.375, 0.188, or 0.125 for bend radius: pipe diameter ratios of 2, 4, or 6 or more, respectively.

For vertical sections, there will be normally no kinetic energy change. The frictional energy losses are calculated as for the horizontal sections. The potential energy gain E_p is given by

$$E_{\rm p} = \left(\frac{G_{\rm s}}{G}\right) \left(\frac{u_0}{u_{\rm p}}\right) \rho_{\rm s} g h \tag{3.43}$$

where h is the height of the section.

As stated earlier, the sum of all the energy losses described above, applying for a particular system layout, will give the total pressure drop through the conveyor. A practical approach for preliminary calculations in selection and design of pneumatic conveying systems is the use of nomographs, such as those presented by Green and Perry (2008).

Example 3.6

Calculate the power required to convey pneumatically 1500 kg/h of a flour of 1300 kg/m³ density, with the particle size distribution given in Table 3.11. The

Weight Retained (g)
0.10
10.00
73.20
76.20
17.00
16.50
6.10
0.90

TABLE 3.11 Example 3.6

distance of transportation will be 50 m, horizontally, and using piping 6.5 cm in diameter. The velocity of air available for the pneumatic conveyor is 10% above saltation velocity. The air density and viscosity can be taken as 1 kg/m³ and 2×10^{-5} kg/m s, respectively.

SOLUTION

From Table 3.11, and consulting US Standard Screen Series, the maximum and minimum particle sizes for the flour are:

 $x_p \text{ (maximum)} \sim 0.850 \text{ mm} = 8.5 \times 10^{-4} \text{m}$

 $x_{\rm p}$ (minimum) ~0.063 mm = 6.3 × 10⁻⁵m

The remaining relevant data are:

$$Q_s = 1500 \text{ kg/h} = 0.416 \text{ kg/s}$$

L = 50 m

 $D_t = 65 \text{ mm} = 0.065 \text{ m}$

 $\rho_s = 1300 \text{ kg/m}^3$

 $\rho_g = 1 \text{ kg/m}^3$

 $\mu_{g} = 2 \times 10^{-5} \text{ kg/ms}$

The example is solved in two steps: first, the saltation velocity is calculated, and second, the energy losses due to every aspect of the particles moving in the airstream inside the pipe are evaluated to estimate the total pressure drop needed for the particles to complete the conveying distance.

Saltation velocity:

Equation 3.26 is used to calculate the parameter x_p^* for the maximum and minimum particle size. For the maximum particle size:

$$x_{\rm p}^* = \frac{8.5 \times 10^{-4}}{\left[3(2 \times 10^{-5})^2 / 4(9.81)(1)(1300 - 1)\right]^{1/3}} = 29.66 \approx 30$$

From Figure 3.33, for $x_p^* = 30$, $u_{ss}^* \sim 6.0$. Thus, substituting values into Equation 3.27:

$$u_{\rm ss} = 0.19(6.0) \left[\frac{4(9.81)(2 \times 10^{-5})(1300 - 1)}{3(1)^2} \right]^{1/3} (65)^{0.4} \approx 4.2 \,\mathrm{m/s}$$

To evaluate the slope in Figure 3.33, and continue solving the example, the value of x_p^* minimum is needed, and it is calculated following the same procedure as above. Using, thus, Equation 3.26, a value of x_p^* for the minimum particle size $(6.3 \times 10^{-5} \text{ m})$ of about 2.2 is obtained.

Using the plot in Figure 3.33 the values y_2 and y_1 corresponding to values $x_2 = 30$ and $x_1 = 2$ are, approximately, 6 and 4, so the slope of the line joining the points for the maximum and minimum particle sizes is

$$n = \frac{6 - 4}{30 - 2} \approx 0.07$$

The slope n is, therefore, greater than 0.068, so that Equation 3.28 should be used to substitute values, as follows:

$$\frac{125.36}{1300} = \frac{0.214(0.07)^{1.5}(u_s - 4.2)}{4.2}$$

where the flux of solids G_s is calculated from the relationship between the mass flow rate and the cross-sectional area of the pipe, that is:

$$G_{\rm s} = \frac{Q}{A} = \frac{0.416 \text{ kg/s}}{0.0033 \text{ m}^2} = 125.36 \text{ kg/m}^2 \text{ s}$$

Transposing for u_s , the saltation velocity is

$$u_c = 106.4 \text{ m/s}$$

Adding 10% to the saltation velocity, the initial air velocity in the pneumatic conveying is calculated as

$$u_0 \approx 117 \text{ m/s}$$

The kinetic energy of the particles is defined by Equation 3.30, where the particle velocity u_p is a function of the terminal settling velocity u_t , depending itself on the settling regime, as explained in Section 3.5.5.4. Iterations are, therefore, needed. Assuming applicability of Equation 3.32 to calculate u_t , it is found that the value of the particle Reynolds number Re_p surpasses the limit of 0.4, so a second approach using Equation 3.33 is done, that is,

$$u_{t} = \left[\frac{4(1300 - 1)^{2}(9.81)^{2}}{225(1)(2 \times 10^{-5})} \right]^{1/3} (8.5 \times 10^{-4}) \approx 4.46 \text{ m/s}$$

verifying the applicability using Re_n:

$$Re_p = \frac{(8.5 \times 10^{-4})(4.46)(1)}{2 \times 10^{-5}} = 189.5$$

agreement is found with the limiting values of Re_p between 0.4 and 500, so the particle velocity is: $u_p = 117 - 4.46 = 112.54$, and the particle kinetic energy is

$$E_{\rm k} = (125.36)(112.54) \approx 14,108 \,{\rm N/m^2}$$

The frictional energy lost due to the gas flow is determined by Equation 3.36, where the value of the friction factor $f_{\rm g}$ depends on the system Reynolds number, that is,

Re =
$$\frac{(0.065)(117)(1)}{2 \times 10^{-5}}$$
 = 380, 250

using, therefore, Equation 3.38:

$$f_{\rm g} = 0.0008 + 0.0552(380250)^{-0.237} \approx 3.43 \times 10^{-3}$$

and the frictional energy lost due to the gas flow from Equation 3.36 is:

$$E_{\rm fg} = \frac{2(3.43 \times 10^{-3})(1)(117)^2(50)}{0.065} \approx 72,173 \,\text{N/m}^2$$

The frictional energy lost due to the particle flow can be estimated using Equation 3.40. In such equation, the gas flux may be calculated as the product of the gas velocity by its density. Since the air density equals unity, the gas flux is: $G = 117 \text{ kg/m}^2 \text{ s}$. Also, the particle friction factor f_p to be substituted into Equation 3.40 can be determined from the system Reynolds number Re as in Figure 3.34. In such figure, for Re = 380,250, $f_p = 0.00007$. Substituting, thus, all the values in Equation 3.40:

$$E_{\rm fp} = \left(\frac{\pi}{8}\right) \left(\frac{0.00007}{0.00343}\right) \left(\frac{1300}{1}\right)^2 \left(\frac{125.36}{117}\right) (72173) \approx 22365 \, \text{N/m}^2$$

The total pressure drop is the sum of all the above energy losses, that is,

$$\Delta P_{total} = E_k + E_{fg} + E_{fp} = 14108 + 72173 + 22365 = 108646 \approx 109 \text{ kN/m}^2$$

The power can be calculated as the product of the volumetric flow rate of air and the total pressure drop. The volumetric flow rate of air $Q_{\rm air}$ is the product of the air velocity and the cross-sectional area of the conveying pipe, that is, $Q_{\rm air} = (117)(0.0033) = 0.3861 \, {\rm m}^3/{\rm s}$. The power would, therefore, be

 $(\Delta P_{total})(Q_{air})=(108646N/m^2)(0.3861m^3/s)=41948.22\approx42000$ W Finally, the horsepower can be determined as

$$Power = \frac{41984.22 \,\text{W}}{735.4996 \,\text{hp/W}} = 57.08 \,\text{hp}$$

3.6 Laboratory Exercise: Pneumatic Conveying Characteristics of Different Granules

3.6.1 Introduction

Pneumatic conveying can be defined as the transport of solid materials, from one place to another, using a transporting gas. The materials can be moved through the pipe with air, but sometimes nitrogen is used for materials that could cause a chemical reaction with the air. One of the most important issues in pneumatic conveying is how to transport the materials with the lowest pressure drop and thus least amount of energy. The pressure drop behavior can be observed by preparing a state diagram. A state diagram is a plot of pressure difference versus the transport gas velocity at a fixed solid flow rate. The objective of the experiment is to derive a basic phase diagram using, at least, two different testing materials.

3.6.2 Equipment and Materials

A laboratory or pilot-scale conveying system comprising: hoppers, air compressor, rotary feeder, polyvinyl chloride (PVC) conveying pipe, control valves, pressure transducers, rotameter, and electronic weight indicator. Polypropylene granules of different size, as testing material.

3.6.3 Procedure

- Operate the compressor to introduce air into the conveying pipe connected to the feed hopper by means of the rotary feeder, by opening the air control valve.
- Drive granules into the conveying line by the feed hopper using the rotary feeder. Handle the valves in order to control solids and air flow rate.
- Control the air flow rate via the rotameter located after the air valve controlling the airstream from the compressor and before the rotary feeder.
- Allow the granules to get into a recycle solids hopper, feed an intermediate hopper by closing the valve after the recycle solids hopper to enable the mass solids flow rate to be measured using the electronic weight indicator adapted to the intermediate hopper.
- Carry out two to three runs varying the solids fed, and the air and solids flow rate by handling the control valves.
- Take readings of the air velocity, pressure at the different pressure transducers, and the weight of solids passing through the intermediate hopper.

3.6.4 Calculations and Report

- Tabulate data on air superficial velocity, solids mass flux, and average pressure gradient between pressure transducers points, for all the experimental runs.
- Derive a plot of superficial air velocity for different values of solids mass flux versus the pressure gradient, for the different granular solids studied. This plot represents the phase diagram for the conveying process.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out, as well as discussion on own data. Compare results, also, with data from a literature search and present a written report, including a thoughtful discussion on the comparative aspects mentioned.

PROBLEMS

- 1. A silo is to be designed to store granular material of 480 kg/m^3 in bulk density. The material is kept on the open forming conical heaps 7.5 m long and 2 m high. A metal cylinder, 2.5 m diameter and 10 m long, may be adapted to construct the silo. The maximum pressure on the base of the cylinder is 14 kPa and the friction coefficient μ' for the granular material is 0.5. Suggest, supported by calculations, whether the cylinder could be adapted to build a silo to store the granular material.
- 2. Predict the basic dimensions of a silo to store 32 tons of a particulate solid of 1600 kg/m^3 density. The material has uniformly sized particulates of 1 mm. The angle of repose of the material is 32° while its friction coefficient μ' is 0.48. The maximum pressure allowed at the bottom of the cylindrical part of the silo is 3.5 kPa.
- 3. Suggest the optimum outlet dimension of a conical hopper in a silo storing moist crushed limestone of 1350 kg/m³ density. The results of characterizing the limestone using in terms of failure properties are presented in Tables 3.12 through 3.14.
- 4. Calculate the saltation velocity for transporting grains with a limiting size of 2 mm and a density of 1300 kg/m³, through a pipe of 6.5 cm in diameter. Air density can be taken as 1 kg/m³ and air viscosity as 2×10^{-5} kg/m s.

TABLE 3.12Wall Friction Angle of Crushed Limestone for Three Moisture Contents (Table for Problem 3)

2.5%	3.8%	5.0%
29°	28°	28.5°

TABLE 3.13Effective Angle of Internal Friction of Crushed Limestone for Three Moisture Contents (Table for Problem 3)

2.5%	3.8%	5.0%
42°	40°	41.6°

TABLE 3.14Consolidating Stress against Unconfined Yield Stress of Crushed Limestone for Three Moisture Contents (Table for Problem 3)

σ ₁ (kPa)	σ _c (kPa)		
	2.5% Moisture	3.8% Moisture	5.0% Moisture
0.2	0.26	0.30	0.25
0.4	0.48	0.50	0.50
0.6	0.70	0.65	0.66
0.8	0.85	0.80	0.82
1.0	0.88	0.90	0.91
1.2	0.99	1.05	0.98
1.4	1.12	1.10	1.08
1.6	1.14	1.15	1.18
1.8	1.18	1.20	1.22
2.0	1.26	1.24	1.28

5. The pneumatic conveying system of the previous example consists on 50 m horizontal pipe, two 0.5 m radius bends, and 15 m vertical pipe. Calculate the overall pressure drop on the system if the initial air velocity is 10% over the saltation velocity.

References

- Arnold, P. C. and Gu, Z. H. 1990. The effect of permiability on the flow rate of bulk solids from mass-flow bins. *Powder Handling & Processing* 2: 229–238.
- Arnold, P. C., McLean, A. G., and Roberts, A. W. 1982. *Bulk Solids: Storage, Flow and Handling*. Newcastle, Australia: The University of Newcastle Research Associates (TUNRA).
- Badger, W. L. and Banchero, J. T. 1955. *Introduction to Chemical Engineering*. New York, NY: McGraw-Hill.
- Bahke, E. A. 1991. New developments of belt conveyor systems. *Bulk Solids Handling* 11: 135–145.
- Benink, E. J. 1989. Flow and Stress Analysis of Cohesionless Bulk Materials in Silos Related to Codes. PhD Thesis, Enschede, The Netherlands: The University of Twente.

- Cabrejos, F. J. and Klinzing, G. E. 1994. Minimum conveying velocity in horizontal pneumatic transport and the pickup and saltation mechanisms of solid particles. *Bulk Solids Handling* 14: 541–550.
- Carson, J. W., Royal, T. A., and Goodwill, D. J. 1986. Understanding and eliminating particle segregation problems. *Bulk Solids Handling* 6: 139–144.
- Clague, K. 1973. *The Effects of Stresses in Bunkers*. PhD Thesis. Nottingham: University of Nottingham.
- Colijn, H. and Peschl, A. S. Z. 1981. Non-symmetrical bin flow problems. *Bulk Solids Handling* 1: 377–384.
- Foote, B. L., Bobbie, L., Radidran, A., and Badiru, A. B. 1988. Simulation and network analysis payoff in conveyor system analysis. *Industrial Engineering* 20: 48–53.
- Green, D. W. and Perry, R. H. 2008. *Perry's Chemical Engineers' Handbook*, 8th Ed. New York, NY: McGraw-Hill.
- Häußler, U. and Eibl, J. 1984. Numerical investigations of discharging silos. *Journal of Engineering Mechanics* 110: 957–971.
- Janssen, H. A. 1895. Versuche über getreidedruck in silozellen. Zeitschrift des Vereins Deutscher Ingenieure 39: 1045–1049.
- Jenike, A. W. 1964. *Storage and Flow of Solids*. Bulletin No. 123 of the Utah Engineer Experiment Station. Salt Lake City UT: University of Utah.
- Jenike, A. W. and Johanson, J. R. 1968. Bin loads. *Journal of Structural Division, ASCE* 95: 1011–1015.
- Jenike, A. W. and Johanson, J. R. 1969. On the theory of bin loads. *Journal of Engineering for Industry, Transactions of the ASME Series B* 91: 339–342.
- Johanson, J. R. 1965. Method of calculating rate of discharge from hoppers and bins. *AIME Transactions* 232: 69–80.
- Johanson, J. R. 1982. Controlling flow patterns in bins by the use of an insert. *Bulk Solids Handling* 2: 495–498.
- Mahmound, M. H. 1977. Practical finite element modeling of silo-material interaction. *Proceedings of the Technical Program,* International Powder and Bulk Solids Handling & Processing Conference, Rosemont, IL, USA.
- Mainwaring, N. J. and Reed, A. R. 1987. Permeability and air retention characteristics of bulk solid materials in relation to modes of dense-phase pneumatic conveyor performance. *Bulk Solids Handling* 7: 415–425.
- Manjunath, K. S. and Roberts, A. W. 1986. Wall pressure-feeder load interactions in mass-flow hopper/feeder combinations: Part I. *Bulk Solids Handling* 6: 769–775.
- Manjunath, K. S. and Roberts, A. W. 1986b. Wall pressure-feeder load interactions in mass-flow hopper/feeder combinations: Part II. *Bulk Solids Handling* 6: 903–911.
- Marinelli, J. and Carson, J. W. 1992. Solve solids flow problems in bins, hoppers and feeders. *Chemical Engineering Progress* 88: 22–28.
- McLean, A. G. 1979. Flow Rates of Simple Bulk Solids from Mass-Flow Bins. PhD Thesis. Wollongong, Australia: University of Wollongong.
- McLean, A. G. 1990. Improved stockpile performance. *Bulk Solids Handling* 10: 19–22. Merrow, E. W. 1988. Estimating startup times for solids-processing plants. *Chemical Engineering* 89: 89–92.
- Mitman, E. T. 1991. Dome silo storage systems. Bulk Solids Handling 11: 71–72.
- Reed, A. R. and Bradley, M. S. A. 1991. Advances in the design of pneumatic conveying systems. A United Kingdom perspective. *Bulk Solids Handling* 11: 93–97.

- Roberts, A. W. 1988. *Modern Concepts in the Design and Engineering of Bulk Solids Handling Systems*. Newcastle, Australia: The University of Newcastle Research Associates (TUNRA).
- Roberts, A. W. 1988b. Some aspects of grain silo wall pressure research-influence of moisture content on loads generated and control of pressures in tall multi-outlet silos. *Proceedings of the 13th International Powder and Bulk Solids Conference*, Chicago.
- Roberts, A. W. 1994. Advances in the design of mechanical conveyors. *Bulk Solids Handling* 14: 255–281.
- Roberts, A. W. 1999. The influence of granular vortex motion in the volumetric performance of enclosed screw conveyors. *Powder Technology* 104: 56–67.
- Roberts, A. W. 2000. Predicting the performance of enclosed screw conveyors. *From Powder to Bulk Conference*. IMechE, London, June 2000.
- Roberts, A. W. and Teo, L. H. 1990. Design considerations for maximum reclaim capacity of conical stockpiles. *Bulk Solids Handling* 10: 9–15.
- Roberts, A. W., Ooms, M., and Scott, O. J. 1986. Influence of vibrations on the strength and boundary friction characteristics of bulk solids and the effect on bin design. *Bulk Solids Handling* 6: 161–169.
- Roberts, A. W., Ooms, M., and Wiche, S. J. 1990. Concepts of boundary friction, adhesion and wear in bulk solids handling operations. *Bulk Solids Handling* 10: 189–198.
- Rose, H. E. and Barnacle, H. E. 1957. Flow of suspensions of non-cohesive spherical particles in pipes. *Engineer, London* 203: 898, 939.
- Royal, T. A. and Carson, J. W. 1993. How to avoid flooding in powder handling systems. *Powder Handling & Processing* 5: 63–67.
- South, D. B. 1991. Curved bin walls. *Bulk Solids Handling* 11: 75–76.
- Terziovski, M. and Arnold, P. C. 1990. On the effective sizing and placement of air blasters. *Bulk Solids Handling* 10: 181–185.
- Thomson, F. M. 1997. Storage and flow of particulate solids. In: *Handbook of Powder Science and Technology*, Fayed, M. E. and Otten, L. (eds.). New York, NY: Chapman & Hall.
- Walker, D. M. 1966. An approximate theory for pressure and arching in hoppers. *Chemical Engineering Science* 21: 975–978.
- Walker, K. 1990. Pendular bucket wheel stacker reclaimer system. *Bulk Solids Handling* 10: 25–26.
- Walters, J. K. 1973. A theoretical analysis of stresses in silos with vertical walls. *Chemical Engineering Science* 28: 13–15.
- Wohlbier, R. H. 1977. Stacking, Blending & Reclaiming of Bulk Materials. Zurich: Trans Tech Publications.
- Wypych, P. W. 1999. The ins and outs of pneumatic conveying. *International Symposium on Reliable Flow of Particulate Solids III*. Porsgrunn, Norway, August 11–13, 1999.
- Zenz, F. A. 1964. Conveyability of materials of mixed particle size. *Industrial and Engineering Chemistry Fundamentals* 3: 65–75.

This page intentionally left blank



Part II Bulk Solids Processing

This page intentionally left blank



Size Reduction

4.1 Fundamental Principles of Comminution

4.1.1 Introductory Aspects

In many applications of the material processing industry, frequently size reduction of solid materials for different purposes is necessary. For example, size reduction may be a specific requirement of a process or may aid other operations such as expression or extraction. Comminution is the generic term used for size reduction, although such a term includes operations such as crushing or pulverizing. The reduction mechanism consists of deforming the solid piece until it breaks or tears. Breaking of hard materials along cracks or defects in their structures is accomplished by applying diverse forces.

The objective of comminution is to produce small particles from larger ones. Smaller particles are the desired product either because of their large surface or because of their shape, size, and number. The energy efficiency of the operation can be related to the new surface formed by the reduction in size. The geometric characteristics of particles, both alone and in mixtures, are important in evaluating a product from comminution. In an actual process, a given unit does not yield a uniform product, whether the feed is uniformly sized or not. The product normally consists of a mixture of particles, which may contain a wide variety of sizes and even shapes. In comminuted products, the term "diameter" is generally used to describe a characteristic dimension related to particle size. As described in Chapter 2, the shape of an individual particle is conveniently expressed in terms of the sphericity Φ_s , which is independent of particle size. For spherical particles, Φ_s equals unity, while for many crushed materials, its value lies between 0.6 and 0.7.

The different types of particle size distributions have been discussed extensively in Chapter 2. No single distribution applies equally well to all comminuted products, particularly in the range of coarser particle sizes. For finer particles, however, the most commonly found distribution follows a log-normal function (Herdan, 1960), which is the most useful one among the different types of functions (Beddow and Meloy, 1980).

4.1.2 Forces Involved in Size Reduction

As previously mentioned, in comminution of solid materials, the reduction mechanism consists of deforming the solid piece until it breaks or tears and such breaking may be achieved by applying diverse forces. The types of forces commonly used in processing industries are compressive, impact, attrition or shear, and cutting. In a comminution operation, more than one type of force acts, usually. Compressive forces are used for coarse crushing of hard materials. Coarse crushing implies reduction to a size of about 3 mm. Impact forces can be regarded as general purpose forces and may be associated with coarse, medium, and fine grinding of a variety of materials. Shear or attrition forces are applied in fine pulverization, when size of products can reach the micrometer range. Sometimes a term referred as ultrafine grinding is associated with processes in which the submicron range of particles is attained. Finally, cutting gives a definite particle size and may even produce a definite shape.

A good number of industrial powders have been produced by crushing of large size pieces. For example, metallic and nonmetallic minerals are extracted from the earth crust in the form of rocks and boulders, but in their final presentation, in products such as carbonates, limestone, or silicates, their appearance is that of fine powders containing particles whose equivalent diameters could well be in the order of micron fractions. It is not possible to reduce the size of a solid material in a single operation and with a single equipment from its macroscopic range of size (e.g., rocks) to its final form of fine powder. The processes of size reduction are carried out, consequently, in stages or defined ranges of reduction. Regularly, the size reduction within the limits of the macroscopic range of objects is named coarse reduction, which applied in the mesoscopic range is known as intermediate reduction, while the one corresponding to the microscopic range is defined as fine reduction. The kinds of forces commonly used in size reduction operations correspond, usually, to such specificity of ranges. Thus, predominating forces in coarse reduction are compressive forces, in intermediate reduction are impact forces, and in fine reduction are shearing forces. Table 4.1 summarizes the types of forces commonly used in size reduction processes.

Cutting forces produce exact and defined sizes and may, even, produce exact shapes. The cutting forces are used for soft materials in operations such

TABLE 4.1Types of Forces Used in Size Reduction Equipment

Force	Principle	Example of Equipment
Compressive	Nutcracker	Crushing rolls
Impact	Hammer	Hammer mill
Attrition	File	Disc attrition mill
Cut	Scissors	Rotary knife cutter

as dicing of vegetables, slicing of fruits, or mincing of meat. As such, the operations based in cutting forces do not generate products with distribution of sizes, and hence the theoretical principles of these differ from the processes typically considered comminution. The operations based on exact reduction by cutting are, therefore, out of the scope of this book, as they may constitute a number of relevant unit operations in more specific processes, for example, in food process engineering.

4.1.3 Properties of Comminuted Materials

The mechanical forces involved in comminution previously discussed are used to reduce the magnitude of the size of diverse hard materials either from inorganic origin (e.g., rocks and minerals) or from organic origin (e.g., grains and oilseeds). For a good number of these materials, independent of their origin, comminuted particles obtained after any size reduction operation will resemble polyhedrons with nearly plane faces and sharp edges and corners. The number of major faces may vary, but will be usually between 4 and 7. A compact grain with several nearly equal faces can be considered as spherical so the term diameter is normally used to describe particle size of these comminuted products. Due to impact forces, but particularly to friction during the processes of reduction as well as in manipulation and transportation afterwards, the angles are softened and the polyhedron shape tends to a spherical shape, and so the sphericity of crushed materials tends to unity, as previously mentioned.

The predictable shape of the crushed particles described in the previous paragraph has to do with the molecular structure of the diverse solid materials subjected to size reduction. The predominant key components of the crystal units which form the solid matrix of materials, both organic and inorganic, are silicon and carbon, respectively. As it is known, such elements belong to the same group in the periodic table, and their principal compounds are on a fundamental atomic structure of tetrahedral shape. When applying mechanical forces tending to reduce the size, the materials will fracture, therefore, forming polyhedral fractions such as the smithereens formed in shattered glass. An ideal size reduction pattern to achieve a high reduction ratio of hard brittle materials could be obtained firstly by compressing, then by using impact force, and finally by shearing or rubbing. Therefore, hard brittle materials would successfully produce powders by using different forces in a comminution operation, whereas tough ductile materials such as meat can only be satisfactorily reduced in size by applying cut forces. As previously stated, cutting is considered a process totally different from comminution because its operating principles are quite different to those governing the size reduction of hard materials.

In any particular comminution operation of hard materials, more than one type of the previously described forces is actually present. Regardless of the uniformity of the feed material, the product always consists of a mixture of

particles covering a wide range of sizes. Some size reduction equipment is designed to control the size of the largest particles in its products, but the fine sizes are not under such control. In spite of the hardness of the comminuted materials, the above-mentioned shape of produced particles would be subjected to attrition due to interparticle and particle-equipment contacts within the dynamics of the operation. Thus, particle angles will smooth gradually with the consequent production of fines. In actual practice, any feed material will possess an original particle size distribution, while the obtained product will end with a new particle size distribution having a whole range finer than the feed distribution. A consequence of the progressive reduction to accomplish high relations of reduction is the excessive accumulation of fines. The fines are presented both by initial edge breaking as for extreme friction in the fine reduction stages. For this reason, the crushed materials increase the number of their particles following an exponential or logarithmic progression and not a linear progression. As the size of the particles is reduced, each mother particle produces a greater number of particles derived from it. The concentration of fines is, normally, uncontrollable and undesirable.

A product specification will commonly require a finished product not to contain particles greater than (or smaller than, depending on the application) some specified size. In comminution practice, particle size is often referred as screen aperture size (x_A , as defined in Chapter 2). The reduction ratio, defined as the relation between average size of feed and average size of product, can be used as an estimate of the performance of a comminution operation. The values for average size of feed and product depend on the method of measurement but the true arithmetic mean, obtained from screen analyses on samples of the feed and product streams, is commonly used for this purpose. Reduction ratios depend on the specific type of equipment. As a general rule the coarser the reduction, the smaller the ratio. For example, coarse crushers have size reduction ratios below 8:1, while fine grinders may present ratios as high as 100:1. However, large reduction ratios, as those obtained when dividing relatively large solid lumps to ultrafine powders, are normally attained by several stages using diverse crushing and grinding machines. A good example of this is the overall milling of wheat grain into fine flour, in which crushing rolls in series of decreasing diameters are employed.

4.2 Energy Requirements in Comminution

As previously discussed, in the breakdown of hard and brittle solid materials, two stages of breakage are recognized: (a) initial fracture along existing fissures within the structure of the material and (b) formation of new fissures or crack tips followed by fracture along these fissures. It is also accepted that only a small percentage of the energy supplied to the grinding equipment is

Size Reduction 183

actually used in the breakdown operation. Figures of less than 2% efficiency have been quoted (Richardson et al., 2002) and, thus, grinding is a very inefficient process, perhaps the more inefficient of the traditional unit operations. Much of the input energy is lost in deforming the particles within their elastic limits and through interparticle friction. A large amount of this wasted energy is released as heat which, in turn, may be responsible for heat damage of thermal-sensitive materials.

Theoretical considerations suggest that the energy required to produce a small change in the size of unit mass of material can be expressed as a power function of the size of the material, that is,

$$\frac{\mathrm{d}E}{\mathrm{d}x} = -\frac{K}{x^n} \tag{4.1}$$

where dE is the change in energy, dx is the change in size, K is a constant, and x is the particle size.

Equation 4.1 is often referred as the general law of comminution and has been used by a number of workers to derive more specific laws depending on the application.

4.2.1 Rittniger's Law

Rittinger (1867) considered that for the grinding of solids, the energy required should be proportional to the new surface produced and gave to the power *n* the value of 2, obtaining thus the so called Rittinger's law by integration of Equation 4.1:

$$E = K \left[\frac{1}{x_2} - \frac{1}{x_1} \right] \tag{4.2}$$

where E is the energy per unit mass required for the production of a new surface by reduction, K is called Rittinger's constant and is determined for a particular equipment and material, x_1 is the average initial feed size, and x_2 is the average final product size. Rittinger's law has been found to hold better for fine grinding, where a large increase in surface results.

4.2.2 Kick's Law

Kick (1885) reckoned that the energy required for a given size reduction was proportional to the size reduction ratio and took the value of the power n as 1. In such a way, by integrating Equation 4.1, the following relation, known as Kick's law is obtained:

$$E = K \left[\ln \frac{x_1}{x_2} \right] \tag{4.3}$$

where x_1/x_2 is the size reduction ratio. Kick's law has been found to hold more accurately for coarser crushing where most of the energy is used in causing fracture along existing cracks.

4.2.3 Bond's Law and Work Index

A third version of the comminution law is the one attributed to Bond (1952, 1963), who considered that the work necessary for reduction was inversely proportional to the square root of the size produced. In Bond's consideration, n takes the value of 3/2, giving the following version (Bond's law) also by integration of Equation 4.1:

$$E = 2K \left[\frac{1}{\sqrt{x_2}} - \frac{1}{\sqrt{x_1}} \right] \tag{4.4}$$

When X_1 and X_2 are measured in micrometers and E in kWh/ton, $K = 5E_i$, where E_i is the Bond Work Index, defined as the energy required to reduce a unit mass of material from an infinite particle size to a size such that 80% passes a 100 μ m sieve. The Bond's work index is obtained from laboratory crushing tests on the feed material. Bond's law holds reasonably well for a variety of materials undergoing coarse, medium, and fine size reduction.

4.3 Size Reduction Equipment

4.3.1 Classification

As has been previously discussed, size reduction is a unit operation widely used in a number of processing industries. Many types of equipment are used in size reduction operations. In a broad sense, size reduction machines may be classified as crushers used mainly for coarse reduction, grinders employed principally in intermediate and fine reduction, ultrafine grinders utilized in ultrafine reduction, and cutting machines used for exact reduction (McCabe et al., 2005). An equipment is generally known as crusher when it performs coarse reduction and mill when used for all other applications. The above-mentioned classification includes several categories of each type of machine; so, in total, a number of approximately 20 different designs are recognized in varied comminution processes. Table 4.2 presents a broad classification of size reduction equipment.

4.3.2 Characteristics

The list shown in Table 4.2 is not exhaustive, since there is a diversity of size reduction equipment and machinery used in a wide range of processing

TABLE 4.2Size Reduction Machines Used in Food Process Engineering

Range of Reduction	Generic Name of Equipment	Type of Equipment
Coarse	Crushers	Jaw crushers Gyratory crushers Crushing rolls
Intermediate	Grinders	Roller mills Hammer mills Tumbling mills Disc attrition mills
Fine	Ultrafine grinders	Hammer mills with internal classification Fluid-energy mills Agitation mills

industries. Some size reduction units have very specific applications. The most common size reduction equipment will be reviewed as follows.

4.3.2.1 Crushers

The size reduction machines known as crushers are slow-speed units used for coarse reduction of large volumes of solids into pieces. The main types of crushers in use are jaw crushers, gyratory crushers, and roll crushers, which comprise smooth- as well as toothed-roll designs. Apart from the toothed-roll crushers, the three remaining crushing machines operate by compression and can break large lumps of extremely hard materials, as in the primary and secondary reduction of metallic and nonmetallic minerals. Toothed-roll crushers perform a dual function of tearing and crushing the feed, and are used for softer materials such as coal, gypsum, limestone, clays, and so on.

In jaw crushers, the crushing elements are two very strong plates called jaws. One of them is vertical and is fixed to the machine frame, while the second is articulated to the equipment so that it can present a sway movement over the fixed one. The three possible forms of articulation constitute the three most popular models known as Dodge, Blake, and Denver crushers, which are illustrated in Figure 4.1. As can be observed, in every case, feeding is carried out by the upper section, the solids go through a narrowing thus the compression to perform the fracture increases, and discharging finishes the operation through the lower section. It can also be appreciated that the Dodge model will give more uniformity in product size as the lower opening is fixed but it will have a greater risk of jamming, which contrast to the opposite operative way of the Blake model. The Denver model, on the other hand, combines the advantages and disadvantages of the previously described models, since the displacement of the mobile jaw is developed in two planes (from left to right and up and down). In all of the models, the active faces of the jaws are flat or slightly bulged, and they may be provided

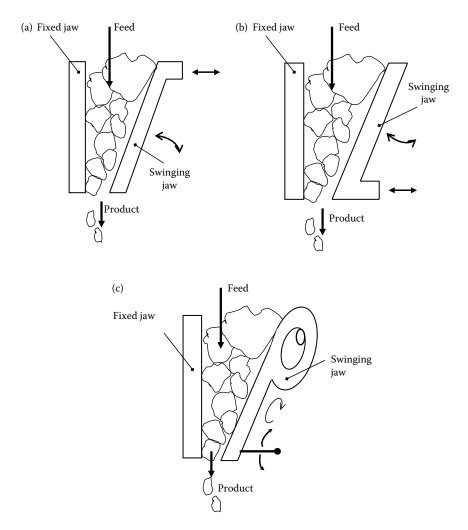


FIGURE 4.1Jaw crushers: (a) Dodge model, (b) Blake model, and (c) Denver model.

with shallow grooves. All jaw crushers are large size units with large capacities and upper openings of up to 3 m^2 of section. They are able to produce up to 600 ton/h of crushed materials.

Rotary crushers consist of a funnel-shaped housing of very resistant interior surface, inside which a conical crushing head moves following a gyroscopic pattern. The crushing head is less tapered than the housing, and mounted in such way that the opening is bigger in the upper section of the equipment and smaller in the lower section. The movement of the crushing head is not a simple rotation, but the consequence of the axis translation adopting the extreme positions pointed in Figure 4.2. The head executes a

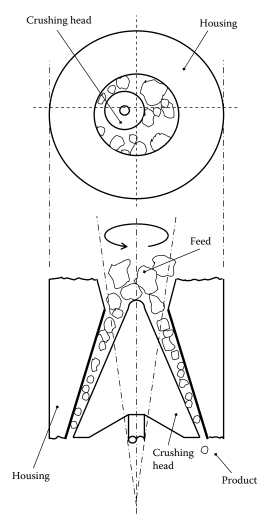


FIGURE 4.2 Gyratory crusher diagram.

full rotation every 300–500 gyroscopic translations. The crushing head is mounted on a heavy shaft pivoted at the top of the machine. An eccentric drives the bottom end of the shaft; so at any point of the periphery of the housing the bottom of the head moves towards and then away from, the stationary wall. Feed is loaded on a side of the crusher, while in the opposite end, the crushing goes on, as illustrated in Figure 4.2. Solids caught in the V-shaped space between the crushing head and the stationary housing are broken and rebroken, until they pass out from the bottom to the discharge port. The size of the product may be varied between certain limits raising or lowering the position of the crushing head regarding the housing.

4.3.2.2 Rollers Mills

As previously mentioned, depending on the level of reduction roller mills may be used for coarse and intermediate grinding mainly, although there are some applications in fine reduction like in the grinding of cereals. In these types of equipment, two or more heavy steel cylinders revolve toward each other, and hence fed particles are nipped and pulled through. The nipped particles are subjected to compressive force causing the reduction in size. In some designs, differential speed is maintained as to exert also shearing forces on the particles. The roller surface can be smooth or can carry corrugations, breaker bars or teeth, as a manner of increase friction, and facilitate trapping of particles between the rolls. Toothed-roll crushers can be mounted in pairs, such as the smooth-roll crushers, or with only one roll working against a stationary curved breaker plate. Toothed-roll crushers are much more versatile than smooth-roll crushers but have the limitation that they cannot handle very hard solids. They operate by compression, impact, and shear and not by compression alone, as do smooth-roll crushers. In order to avoid breakings and deformations of the mounting axes of the rollers, at least one of them is adapted over a slide and is kept in position by strong springs

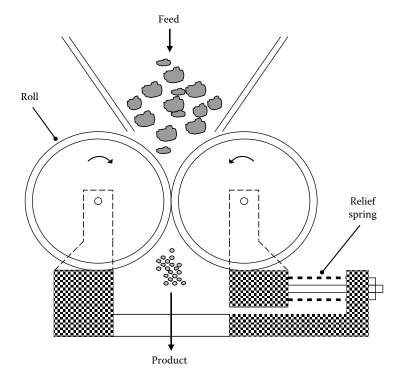


FIGURE 4.3 Diagram of crushing rolls.

which act over the bearings and allow it to retreat lightly to give way to some coarse or unbreakable piece, or in case of jamming in the loading zone. Figure 4.3 shows a diagram of a typical roller mill.

When used as crushers, roller mills have diameters of up to 2 m with lengths of up to 1 m and operate at speeds between 50 and 300 rev/min. In optimal operation, they only produce low relations of reduction in the order of 3 or 4:1, meaning that the feed reduces its average size a fourth of its original size as a maximum. The diameter and spacing between the rollers may be modified in a great range, which allows considerable variations in the average sizes of feeds and products. This characteristic, along with the low installation cost, allows the roller mills to be employed in a diversity of applications. For example, as previously mentioned, they are widely used in diverse stages of cereal grinding, mainly wheat. They are also used in order to reduce the size of diverse minerals such as calcite and kaolin, and for ceramic materials such as magnetite, zircon sand, silicon carbide, and so on.

4.3.2.3 Hammer Mills

Figure 4.4 shows a hammer mill, an equipment that contains a high-speed rotor turning inside a cylindrical case. The rotor carries a collar bearing a number of hammers around its periphery. By the rotating action, the hammers swing through a circular path inside the casing containing a toughened breaker plate. Feed passes into the action zone with the hammers driving the material against the breaker plate and forcing it to pass through a bottom mounted screen by gravity when the particles attain a proper size. Reduction is mainly due to impact forces, although under choking conditions, attrition forces can also play a part in such reduction. The hammers may be replaced by knives, or any other devices, as to give the mill the possibility of handling tough, ductile, or fibrous materials. The hammer mill is a very versatile piece of equipment, which gives high reduction ratios and may handle a wide variety of materials from hard and abrasive to fibrous and sticky. Hammer mills provide good performance, high capacity, and need little space, apart from attaining the highest reduction relations for single comminution units. Their disadvantages are the elevated noise and vibration they cause, and the accelerated wearing of their hammers and hardened walls.

The applications of hammer mills are numerous and cover diverse industrial processes. They are capable of grinding almost anything: tough fibrous solids such as bark or leather, steel turnings, soft wet pastes, sticky clay, hard rock, and so on.

4.3.2.4 Disc Attrition Mills

These types of mills, as those illustrated in Figure 4.5, make use of shear forces for size reduction, mainly in the fine size range of particles. There are several basic designs of attrition mills. The single disc mill (Figure 4.5a) has

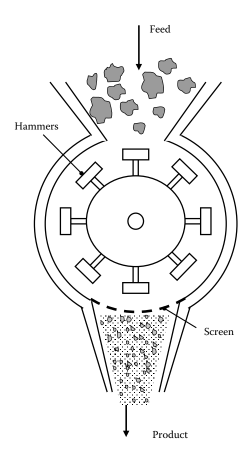


FIGURE 4.4 A hammer mill.

a high-speed rotating grooved disc leaving a narrow gap with its stationary casing. Intense shearing action results in comminution of the feed. The opening between the discs is adjustable between certain limits, and at least one of them, as in the case of the roller mills, is mounted on a sliding axis with tense spring action in order to avoid jamming. In the double-disc mill (Figure 4.5b), the casing contains two spinning discs, which rotate in opposite directions giving a greater degree of shear compared with the single-disc mill. The pindisc mill carries pins or pegs on the rotating elements. In this case, impact forces also play an important role in particle size reduction. The Buhr mill (Figure 4.5c), which is the older type of attrition mill originally used in flour milling, consists of two circular stones mounted on a vertical axis. The upper stone is normally fixed and has a feed entry port, while the lower stone rotates. The product is discharged over the edge of the lower stone. The discs are generally made of iron or steel for materials of a certain toughness, or stone for softer materials. The rubbing face of the discs includes different

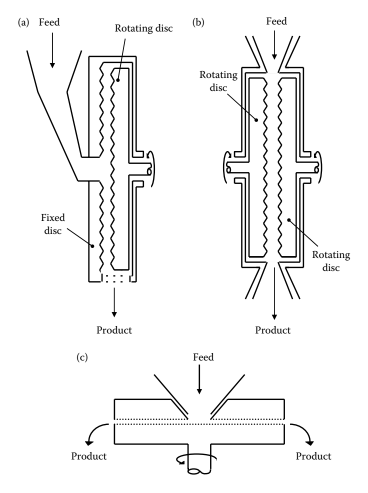


FIGURE 4.5 Disc attrition mills: (a) single-disc mill, (b) double-disc mill, and (c) Buhr mill.

patterns of grooves, corrugations, or teeth, and thus they can carry out a variety of operations including grinding, cracking, granulation, and shredding. Due to the high speed of disc rotation, a considerable amount of heat may be generated, requiring, sometimes, of cooling with water or refrigerated brine. Applications of disc mills are numerous, including dry grinding of wheat, wet grinding of corn for separation of gluten, sugar crushing, cocoa pulverization, and production of fish flour.

4.3.2.5 Tumbling Mills

A tumbling mill is used in many industries for fine grinding. It basically consists of a horizontal slow-speed rotating cylinder partially filled with either balls or rods. The cylinder shell is usually of steel, lined with carbon steel plate, porcelain, silica rock, or rubber. The balls are normally made of steel or flint stones, while the rods are usually manufactured by high-carbon steel. The reduction mechanism is carried out as follows: as the cylinder rotates, the grinding medium is lifted up the sides of the cylinder and drop on to the material being comminuted, which fills the void spaces between the medium. The grinding medium components also tumble over each other, exerting a shearing action on the feed material. This combination of impact and shearing forces brings about a very effective size reduction. Since a tumbling mill basically operates in a batch manner, different designs have been developed to make the process continuous.

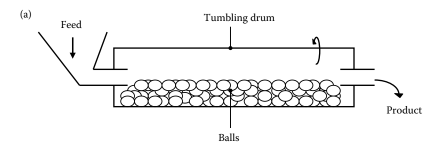
As illustrated in Figure 4.6a, in a trunnion overflow mill, the raw material is fed in through a hollow trunnion at one end of the mill and the ground product overflows at the opposite end. Putting slotted transverse partitions in a tube mill converts it into a compartment mill (Figure 4.6b). One compartment may contain large balls, a second one small balls, and a third one pebbles, achieving thus a segregation of the grinding media with the consequent rationalization of energy. A very efficient way of segregating the grinding medium is the use of the conical ball mill shown in Figure 4.6c. While the feed solid enters from the left into the primary grinding zone where the diameter of the shell is a maximum, the comminuted product leaves through the cone at the right end where the diameter of the shell is a minimum. As the shell rotates, the large balls move toward the point of maximum diameter, and the small balls migrate toward the discharge outlet. Therefore, the initial breaking of feed particles is performed by the largest balls dropping the greatest distance, whereas the final reduction of small particles is done by small balls dropping a smaller distance. In such an arrangement, the efficiency of the milling operation is greatly increased.

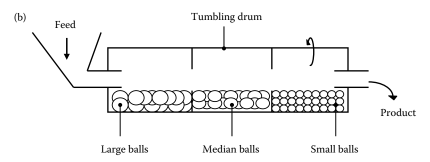
Applications of ball mills are varied and may be used in calcite and kaolin grinding, as well as in grinding of diverse ceramic materials employed in processes of the refractory industry.

4.3.2.6 Other Types of Mills

For ultrafine grinding, some designs previously described may be used at smaller scales. For example, high-speed hammer mills provided with internal or external classification can handle a variety of very fine dry powders with great efficiency. Another alternative for grinding of ultrafine dry powders would be the fluid-energy or jet mills. Agitation mills are used for ultrafine wet grinding.

A hammer mill for ultrafine grinding with intermediate classification is known as the Mikro-Atomizer. In this model, a set of swing hammers is mounted between two rotor discs in the same fashion used in conventional hammer mills. In addition to the hammers, the rotor shaft carries two fans in





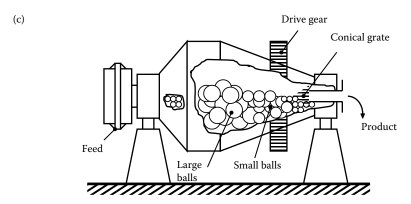


FIGURE 4.6 Tumbling mills: (a) trunnion overflow mill, (b) compartment mill, and (c) conical mill.

order to draw air in the direction of an outlet channel as to carry the produced fine particles within the discharged air stream. There are short radial vanes located on the rotor discs for separating oversize particle from those of acceptable size. Due to the high rotation speed in the mill, coarse particles are concentrated along the wall of the chamber because of centrifugal force acting on them. The air stream tends to carry the fine particles inward from the grinding zone toward the shaft, while the separator vanes tend to throw particles outward away from the shaft. The passing of a given particle

between the separator vanes and out to be discharged depends on which force predominates: the drag exerted by the air or the centrifugal force exerted by the vanes. Finer particles are carried through and discharged, whereas coarser particles return to be reduced even more in the grinding chamber. The particle size of the product may be varied changing the rotor speed and the size or number of the separator fins.

In fluid-energy (jet) mills, the particles of the solid are suspended in a stream of gas and driven along a circular or elliptical duct at high speed. The size reduction of the particles is accomplished, in part, by particle–particle impact and friction. The particles pass through a classifier nozzle when they reach a determined size. The suspending gas is, usually, compressed air or superheated steam, admitted at a pressure of approximately 6900 Pa. Solids are fed near the bottom of the loop through a Venturi injector. Classification of ground particles takes place at the upper bend of the loop. The gas flows around this bend at high-speed projecting coarser particles toward the outer wall and congregating the fines in the vicinity of the inner wall. A discharge opening against the inner wall at this upper part of the loop leads to cyclone separator and a bag collector. Fluid-energy mills are highly efficient when the fed particles are no larger than 150 μm .

Agitation mills are vessels of capacities between 4 and 1000 L, which contain a liquid with crushing elements such as balls or sand grains in suspension. The solid to be ground is fed by the upper section and some kind of agitation is applied, while the product is taken by the lower section through a sieve.

Cutting machines that are similar to grinding equipments and produce fine systems with uniform size distributions are known as rotary knife cutters. They are used for materials that are too tenacious or too resilient to be broken by compression, impact, or attrition such as rubbers, plastics, or leaves of vegetables and spices. The rotary knife cutters have a very similar design to the hammer mills, since they have a horizontal rotor that spins between 200 and 900 rev/min inside a cylindrical chamber. The rotor has mounted from 2 to 12 hardened steel or stellite knives that pass leaving a very narrow clearing in front of a series of stationary knives. Feed is introduced by the upper section and the product passes through a sieve in a lower opening in order to be collected and withdrawn. The cutting elements are exchangeable according to the specific needs of a process.

Example 4.1

A crusher is fed with limestone having particles of 2 cm median equivalent diameter and discharges a product consisting of particles of 0.5 cm median equivalent diameter. The equipment operates at a capacity of 1.2×10^4 kg/h consuming a power of 10 hp. If the requirements of the process demand a finer size of the product (0.4 cm of median equivalent diameter) decreasing the capacity to 1.0×10^4 kg/h, determine the theoretical power consumption under the new conditions.

SOLUTION

The feed particles can be considered very coarse, and so Kick's law would be the most appropriate. The starting and final sizes, in average, are of 0.02 and 0.005 m, respectively, while the power is equivalent to 7.46 kW. It is known, also, that the energy of operation is derived from the relation between power and flow rate, that is, E = P/Q, therefore, the constant K for Equation 4.3 may be calculated as follows:

$$K = \frac{\frac{7.46}{12000}}{\ln\left(\frac{0.02}{0.005}\right)} = 4.48 \times 10^{-4} \text{ kWh/kg}$$

Since the new conditions are not significantly different, the constant *K* may be used in the same relationship to find for the new conditions the power as

$$P = (10000)(0.000448) \ln \left(\frac{0.02}{0.004} \right) = 7.21 \text{ kW}$$

Thus, the theoretical power for the new conditions of the comminution operation will be 7.2 kW approximately.

4.3.3 Operation of Equipment

The diversity of designs of the above-described machinery implies that their operating variables differ quite considerably. While energy requirements are generally governed by the comminution laws previously discussed, as illustrated in Example 4.1, some other features such as capacity and rotational velocity are particular for each unit and may be related, in some way, to the predominant force performing the reduction action. Two types of equipment with critical particularities of operation would be roller mills and tumbling mills.

In crushing rolls, the angle formed by the tangents to the roll faces at the point of contact between a particle and the rolls is called the angle of nip. It is an important variable for specifying the size of a pair of crushing rolls for a specific duty and is found as follows. Figure 4.7 shows a pair of rolls and a spherical particle just being gripped between them. The radii of rolls and particle are R and r, respectively. The clearance between the rolls is 2d. Line AB passes through the centers of the left-hand roll and the particle, as well as through point C, which is the point of contact between the roll and the particle. As shown in Figure 4.7, if α is the angle between line AB and the horizontal, line OE is a tangent to the roll at point C and it makes the same angle α with the vertical. Neglecting gravity, two forces act at point C: the tangential frictional force F_{tr} having a vertical component F_{tr} sin α . Force F_{tr} is related to force F_{tr} through the coefficient of friction μ , so $F_{tr} = \mu' F_{tr}$. Force F_{tr} sin α tends to expel

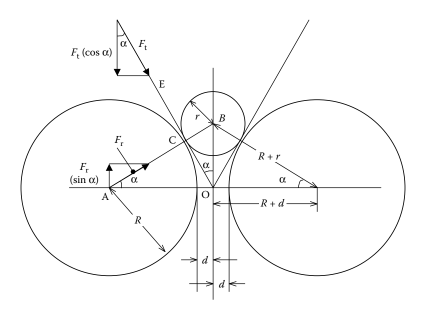


FIGURE 4.7 Diagram of forces and angle of nip in crushing rolls.

the particle from the rolls, while force $\mu T_r \cos \alpha$ tends to pull it into the rolls. If the particle is to be crushed:

$$F_r \mu' \cos \alpha \ge F_r \sin \alpha$$
 (4.5)

or

$$\mu' \ge \tan \alpha$$
 (4.6)

When $\mu' = \tan \alpha$, the angle α is half the angle of nip. A simple relationship exists between the radius of the rolls, the size of the feed, and the gap between the rolls. Thus, from Figure 4.7,

$$\cos \alpha = \frac{R+d}{R+r} \tag{4.7}$$

The largest particles in the product would have a diameter 2*d* and Equation 4.7 provides a relationship between the roll diameter and the size reduction that can be expected in the mill.

From Figure 4.7, it follows that 2R will represent the diameter of the roll D_r , 2r will be the diameter of the feed D_f (when the feeding pieces have a proper sphericity Φ_s), and 2d will be considered the product diameter D_p . Considering these definitions, the theoretical volumetric capacity Q of crushing

Size Reduction 197

rolls is the volume of the continuous ribbon of product discharged from the rolls and can be given by

$$Q = \frac{ND_{\rm r}D_{\rm p}L}{60} \tag{4.8}$$

where *N* is the roll speed in rpm, and *L* is the length of face in meters.

Knowing the bulk density of the discharge stream, the approximate mass flow rate may be estimated. In practice, the actual capacity is found to lie between 0.1 and 0.3 of the theoretical one.

The load of balls in a tumbling mill should be such that when the mill is stopped, the balls occupy somewhat more than one-half the volume of the mill. In operation, the balls are picked up by the mill wall and carried nearly to the top where they fall to the bottom to repeat the process. Centrifugal force maintains the balls in contact with the wall and with each other during the upward trajectory. While they keep in contact with the wall, the balls exercise some grinding action by slipping and rolling over each other. Most of the grinding occurs, however, at the zone of impact where the free-falling balls strike the bottom part of the mill. The faster the mill rotates, the farther the balls are carried up inside the wall and the greater the power consumption. The added power is profitable used because when the balls are carried at the higher point, they will have a greater impact on the bottom and would perform a better reduction capacity. When the speed is too high, however, the balls are carried over and the mill would be practically centrifuging the balls, impeding them to fall. The speed at which centrifuging occurs is called the critical speed and little or no grinding is carried out when the mill operates at this, or higher, velocity. Operating speeds must be well calculated in order not to be considerably smaller than the critical speed, because little grinding action will occur, or considerably higher than the critical speed, because centrifuging will cancel the grinding capacity of the mill.

The speed at which the outermost balls lose contact with the wall of the mill depends on the balance between gravitational and centrifugal forces. Referring to the diagram in Figure 4.8, considering the ball at point A on the periphery of the mill, the radii of the mill and the ball will be R and r, respectively. The center of the ball is, thus, R-r from the axis of the mill and the radius AO form the angle α with the vertical. Two forces act on the ball: the force of gravity mg, where m is the mass of the ball, and the centrifugal force $mu^2/(R-r)$, where u is the peripheral speed of the center of the ball. The centripetal component of the force of gravity is $mg(\cos \alpha)$, which opposes the centrifugal force. As long as the centrifugal force exceeds the centripetal one, the particle will not loss contact with the wall. As the angle α decreases, however, the centripetal force increases. If the speed does not exceed the critical value, a point is reached where the opposing forces are equal and the

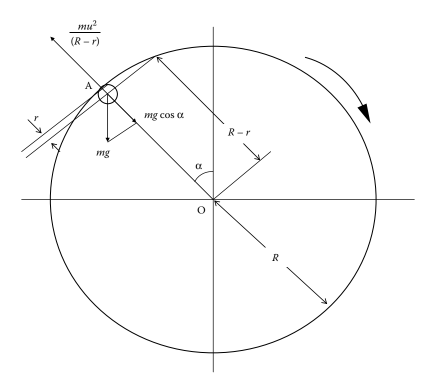


FIGURE 4.8 Diagram of forces on balls in a ball mill.

particle is nearly to fall. The angle at which this occurs is found by equating the centrifugal and centripetal forces, that is,

$$mg\cos\alpha = \frac{mu^2}{(R-r)}\tag{4.9}$$

Transposing for $\cos \alpha$, Equation 4.9 transforms to

$$\cos \alpha = \frac{u^2}{(R-r)g} \tag{4.10}$$

The speed u is related to the speed of rotation by the equation

$$u = 2\pi N(R - r) \tag{4.11}$$

and, thus, Equation 4.11 can be written as

$$\cos \alpha = \frac{4\pi^2 N^2 (R - r)}{g} \tag{4.12}$$

Size Reduction 199

At the critical speed, $\alpha = 0$, consequently $\cos \alpha = 1$, and N becomes the critical speed N_c . With all these considerations, Equation 4.12 transforms into

$$N_{\rm c} = \frac{1}{2\pi} \sqrt{\frac{g}{(R-r)}}$$
 (4.13)

Rod mills can produce 5–200 ton/h of material reduced up to about 1 mm sizes, while ball mills can give 1–50 ton/h of fine powder with 70–90% sizes in the range of 70 μ m. The total energy requirement for a typical rod mill grinding different hard materials is about 5 hp h/ton, whereas for a ball mill, it is approximately 20 hp h/ton. Tube mills and compartment mills normally need somewhat more power than these figures. As the product becomes finer, the capacity of a given mill diminishes and the energy requirement increases.

Example 4.2

A roller mill is used to obtain 4900 kg/h of ground bentonite with 960 kg/m³ bulk density. The rollers spin both at 1.5 m/s and are 1.22 m long with 7.6 cm diameter. Empirical measurements have determined that the mill operates at 30% efficiency. Calculate the relation of reduction obtained in the operation if the friction coefficient is 0.52.

SOLUTION

The relation of reduction is obtained as the relation between equivalent mean diameters of the feed and the product. In order to obtain the median size of the product, Equation 4.8 may be used to transpose the product diameter $D_{\rm p}$, that is,

$$D_{\rm p} = \frac{(C)(60)}{ND_{\rm r}L}$$

It is known that the peripheral speed v and the rotation speed N are related by: $v = \pi D_r N$, and, therefore, substituting in such relation the rotation speed can be calculated as 188.47 rev/min. On the other hand, since the equations are theoretical, process theoretical capacity is determined as: (4900 kg/h)/0.3 = 16333 kg/h. Using the bulk density, mass flow rate could be transformed into volumetric flow rate, as $(16,333 \text{ kg/h})/960 \text{ kg/m}^3 = 17.01 \text{ m}^3/\text{h}$. Furthermore, considering the equivalence between seconds and hours, the volumetric capacity of $0.0047 \text{ m}^3/\text{s}$ would be taken to substitute in the above transposed equation. Thus, the product diameter D_p is obtained as

$$D_{\rm p} = \frac{(0.0047)(60)}{(188.47)(0.152)(1.22)} = 0.00806 \,\mathrm{m} = 0.8 \,\mathrm{cm}$$

In order to obtain the average median size of the feed, Equation 4.7 could be used, since, as it was previously mentioned, $2r = D_f$. By using the friction coefficient μ' ,

according to Equation 4.6 it could be assumed that: $\mu' = \tan \alpha$ and the arc tangent of the friction coefficient would give the value of the angle α . By this procedure the value of the angle: $\alpha = 27.47^{\circ}$. Knowing this value, $\cos \alpha$ can be also known as 0.8872. From Equation 4.7, the radius of the feed particle r is calculated as

$$r = \frac{R+d}{\cos\alpha} - R = \frac{0.076 + 0.004}{0.8872} - 0.038 = 0.014 \text{ m}$$

Since $D_a = 2r$, $D_a = 0.028 \text{ m} = 2.8 \text{ cm}$.

Finally, the relation of reduction would, therefore, be

Relation of reduction =
$$\frac{D_a}{D_p} = \frac{2.8}{0.8} = 3.5$$

Thus, the relation of reduction is 3.5:1, approximately.

Example 4.3

Calculate the critical speed in revolution/minute, of a ball mill with an internal diameter of 1200 mm loaded with balls of 70 mm diameter.

SOLUTION

Expressed in meters, the radius of the mill and the balls will be 0.6 and 0.035 m, respectively. Therefore, a direct substitution in the Equation 4.13 will lead to

$$N_c = \frac{1}{2\pi} \sqrt{\frac{9.81}{(0.6 - 0.035)}} = 0.6625 \text{ 1/s} = 39.75 \text{ rev/min.}$$

4.4 Criteria for Selecting Size Reduction Processes

In deciding how to crush or grind a solid material, process engineers should consider factors such as the size distributions of feed and product, the hardness and mechanical structure of the feed, the moisture, and the temperature sensitivity of the feed. Regarding the size distributions of the materials, each type of crusher or grinder is intended for a certain size of feed and product. It is usually possible to exercise some control over the size of feed, but sometimes it must be taken as it comes. As there is an upper limit on the size that can be accepted by a machine without jamming, if there is an oversized material, a guard screen is needed to keep too large pieces out of the crusher or grinder. In the case of too much undersized material, prescreening the feed can cut the amount which goes through the equipment. For a small-scale operation, such a cut is important as it decreases the capacity required; in a large-scale equipment, though, the undersized particles simply pass

through the throat, where there is always ample room, and so removing them does not greatly affect capacity. Although many criteria and factors may be taken into consideration, paramount factors in designing comminution processes would be

- Characteristics of feed material: mechanical and structural; related to hardness, abrasiveness, size, shape, and so on
- · Feed and discharge control
- Moisture
- Heat removal or supply

4.4.1 Characteristics of Raw Materials

Regarding the main features of the feed material, the pieces or particles must be adequate in size and shape in order to avoid jamming, entrapment, or general malfunction of the equipment used to accomplish the size reduction. Prior knowledge of the particular structure of the feed material can indicate the type of force most likely to be used in performing the size reduction. If the material is friable or has a crystalline structure, fracture may occur easily along cleavage planes, with larger particles fracturing more easily than smaller ones. In these cases, crushing using compressive forces would be recommended. When few cleavage planes are present, and new crack tips have to be formed, impact and shear forces may be more advisable. There are some solid materials having fibrous structures and hence they are not easily reduced in size by compression or impact. In such cases, shredding or cutting may need forces to perform the desired size reduction.

One of the major factors that govern the choice and design of size reduction machines is the hardness of the material to be processed. The greater the hardness, the highest energy consumption and the longer the time in the action zone of the reducing forces. Normally, very hard materials are also very abrasive; so the parts of the equipment in contact with them must be made of special materials. As a general guidance, the hardness is defined in accordance with Mohs' scale, which is divided into 10 grades of hardness (Chapter 1). As a rule of thumb, using Mohs' scale, any material would be considered soft when having a value between 1 and 3, medium-hard if presenting a value between 3.5 and 5, and hard when showing a value between 5 and 10.

4.4.2 Feeding and Discharge Control

In most comminution processes, maintaining continuous flow of feed and product is a requirement. For this purpose, a pretreatment of the feed by sieving or even by grinding (e.g., coarse crushing or breaking before passing to an intermediate reduction mill) is necessary. The discharge rate should

match the feed rate as much as possible, which is normally guaranteed by the assistance of the force of gravity. Sometimes, however, the use of some sort of conveying system to aid either feed or raw materials or discharge of finished products could be necessary. Some comminution equipment may require a very specific discharge system, such as the peripheral discharge devices adapted to certain tumbling or friction mills. In some ultrafine grinding equipment the fine size of the product requires of a fluid, such as air or water, to assist product collection. A suspension stream is, inevitably, created so that a number of separation devices to recover the dry solids, such as cyclones or bag filters, would be required.

4.4.3 Moisture

The presence of moisture can be either beneficial or inconvenient in comminution processes. It is well known that safety problems are caused by dust formation arising during the dry milling of many solid materials. It is known, for example, that a concentration of fine inert powders over 30 mg/L requires a spark of 100 mJ to cause an explosion. A spark of this magnitude may be caused by an electric failure or by metal-metal accidental collisions, common in handling tools. The presence of small quantities of water has been found useful in the suppression of dust. In applications where the presence of moisture is acceptable, water sprays are often used to reduce dust formation. Some other applications allow for large quantities of water to be introduced in the size reduction process; wet milling of corn is a good example of this. On the other hand, in many cases feed moisture content in excess of 2–3% can lead to clogging of the mill with the consequent effect on throughput and efficiency. Agglomeration can also be caused by moisture, being undesirable when a free-flowing powder is needed to control feed rate.

4.4.4 Heat Generation and Removal

As has been stated earlier, comminution is, possibly, the most inefficient unit operation in the material processing industry. The excessive friction presented in most size reduction machines causes increase in heat which can lead to a considerable increase in temperature of the material being processed. There are many materials of biological and organic origin that are normally heat sensitive, and may cause degradation reactions, release of sticky substances, and many other undesirable phenomena. Knowledge of the chemical stability of heat-sensitive materials, as well as some relevant physical properties such as their melting temperatures, should be taken into account before deciding the type of machine used for their size reduction. Many crushing and grinding machinery and equipment may, therefore, be equipped with cooling devices such as jackets, coils, and so on.

TABLE 4.3Preliminary Guide for Selecting Size Reduction Equipment

•		O	1 1		
Equipment	Max. Feed Size (mm)	Min. Prod. Size (mm)	Capacity (ton/day)	Hardness Limit (Mohs)	Applications Examples
Jaw crusher	1500	150	<1->103	9	Metallic and nonmetallic minerals
Gyratory crusher	2000	300	>103	9	Metallic and nonmetallic minerals
Roller mill	30	1	$1 - > 10^3$	7.5	Cereals, vegetables, calcite, kaolin
Hammer mill	40	0.01	<1 - <103	4	Phosphates, pigments, dried fruits
Disc attrition mill	12	0.07	<1 – 103	3	Cellulose, asbestos, rubber
Ball mill	4	0.3	$10 - > 10^3$	9	Calcite, kaolin, ceramics
Fluid-energy mill	30	0.001	<1 - 102	9	Ceramics, pesticides, pigments
Agitation mill	0.075	0.005	<1 – 10 ²	5	Silicon, titanium dioxide, ceramics

4.4.5 Summary of Criteria and Main Applications

As previously discussed, one of the determinant factors for the right selection of size reduction equipment is consideration of the hardness of the material to be processed. The second factor of great relevance is the required capacity, either fixed or subjected to variations by process requirements. Taking these two main considerations along with a thorough analysis of as many other factors as necessary, flowcharts can be devised to select and locate the most suitable comminution units for a size reduction plant. The combination of different criteria may be summarized in charts, like the one given in Table 4.3, which includes some typical applications of the principal size reduction equipment.

Example 4.4

Calcite rocks of 2350 kg/m³ density are comminuted in crushing rolls, having 1 m diameter and 40 cm length. The faces of the rollers are separated by 1.25 cm in the narrowest point and operate at a rotation speed of 75 rpm. The angle of nip has been determined as 30°. Determine the maximum permissible size of fed particles and the capacity in ton/h, considering that it only represents 12% of the theoretical.

SOLUTION

As in the case of Example 4.2, in order to calculate the maximum permissible size of feed, the transposed form of Equation 4.7 could be used, considering that the angle α represents half the angle of nip. Thus: $\cos \alpha = \cos(15^\circ) = 0.9659$. On the other hand, the radius of the roller, R, is 0.5 m, while the dimension d represents half the distance between rollers, that is, 0.00625 m. Substituting, then in the transposed form of Equation 4.7,

$$r = \frac{0.5 + 0.00625}{0.9659} - 0.5 = 0.024 \text{ m}$$
, therefore: Da = $2r = 0.048 \text{ m} \sim 5 \text{ cm}$

Equation 4.8 can be used to compute the theoretical volumetric flow rate. Substituting all the terms,

$$Q = \frac{(75)(1)(0.0125)(0.38)}{60} = 0.00625 \text{ m}^3/\text{s}$$

Since real capacity only represents 12% of theoretical, in reality the capacity would only be $0.00075~\text{m}^3/\text{s}$. Also, the answer is required in ton/h, so using the density of the material in order to transform to mass flow: $(0.00071~\text{m}^3/\text{s})$ (2350 kg/m³) = 1.7625~kg/s. Finally, transforming to kg/h and then to ton/h, the required capacity of 6.345~ton/h is obtained. Thus, the results are maximum permissible size of fed particles in average $\approx 5~\text{cm}$; optimum capacity $\approx 6.3~\text{ton/h}$.

4.5 Laboratory Exercise: Determination of Reduction Relations Using a Hammer Mill

4.5.1 Introduction

It has been argued that hammer mills constitute the most efficient comminution machines available for different size reduction purposes. Their relations of reduction are, therefore, considered among the top within the range of size reduction equipment. The object of this experiment is to study the factors affecting the performance of a hammer mill using the reduction ratio obtained as criteria.

4.5.2 Equipment and Materials

Laboratory-scale hammer mills are available from a number of equipment manufacturers. Any laboratory-scale unit would be suitable for the laboratory exercise. Suggested raw materials are granulated sugar and granulated salt.

Size Reduction 205

4.5.3 Instruments

Standard set of sieves as particle size analyzers.

4.5.4 Screen Analysis

Measure the particle size distributions of feed materials and final products performing a screen analysis, and determine the mean particle sieve diameter as described in Chapter 2 (Section 2.7).

4.5.5 Operation of the Hammer Mill

- Select an appropriate amount of a raw material according to mill size and screen it through a set of sieves so as to determine the size of distribution and average size, according to the procedure mentioned in the previous section.
- Select one of the screen plates of the mill, perhaps one with the finest holes, and screw it into position. Start the motor and read the wattmeter over a short period. This reading will give a measure of the power consumption with the machine running idle.
- Introduce a weighed quantity of the material, according to specific instruction of the equipment, and keep a constant rate sufficient to maintain the ammeter reading constant, so as to guarantee efficient use of energy in the run.
- Collect the product for the entire run and carry out a screen analysis
 of the product, similar to the one carried out before for the feed
 material.
- Repeat the experiment using the second feed material.
- Determine the reduction relation ratio for the two experiments.

4.5.6 Calculations and Report

- Compare the reduction relations for the tested materials and elaborate a discussion on significant differences, if any, of these two materials.
- Perform a literature search to find out whether similar materials comminuted in different size reduction equipment, would present different reduction relations. Analyze and discuss on these findings.
- Present a complete written report, according to scientific report guidelines, either given by the instructor or consulted in the literature. Include graphs, charts, and calculations carried out. Include, as an important part of the report, the discussion elaborated from the literature survey previously performed.

PROBLEMS

- 1. 150 ton/day of pyrite of 2 mm average size need to be milled to an average size of 0.1 mm, using any of three ball mills of 12.5, 25, and 50 cm in diameter, respectively, all with balls of 5 cm of diameter as the pulverizing medium. Which mill would be more adequate to perform the duty knowing that, everything included, 10 kW/rpm are approximately needed to operate? The value of K has been reported as 2.5×10^{-8} kW h m/kg, and the process is 30% efficient.
- 2. 120 ton/h of clay having an average size of 101.6 mm (4 in.) is going to be reduced to an average size of 3.175 mm (1/8 of an in.). If the work index E_i has been estimated as 16.4, calculate the power necessary to crush the material, as well as the power needed to crush it even more, and hence the final size would be of 1 mm average size.
- 3. A material is fed to a pair of crushing rollers, each one rotating at the same speed of $1.25 \, \text{m/s}$. Each roller is $1.5 \, \text{m}$ long with $7.5 \, \text{cm}$ diameter. The product has a density of $1050 \, \text{kg/m}^3$ and is discharged at a flow of $5000 \, \text{kg/h}$. If the measured flow is 30% of the theoretical value, calculate the relation of reduction of the process.

References

Beddow, J. K. and Meloy, J. K. 1980. *Testing and Characterization of Powders and Fine Particles*. London: Heyden and Son.

Bond, F. C. 1952. The third theory of comminution. AIME Transaction 193: 484-494.

Bond, F. C. 1963. Some recent advances in grinding theory and practice. *British Chemical Engineering* 8: 631–634.

Herdan, G. 1960. Small Particle Statistics. London: Butterworths.

Kick, F. 1885. Das Gesetz der Proportionalen Widerstande und Sein Enwendung. Leibzig: Artuz Felix.

McCabe, W. L., Smith, J. C., and Harriot, P. 2005. *Unit Operations in Chemical Engineering*, 7th Ed. New York, NY: McGraw-Hill.

Richardson, J. F., Harker, J. H., and Backhurst, J. R. 2002. Coulson & Richardson's Chemical Engineering, Vol. 2. Oxford: Butterworth-Heinemann.

Rittinger, P. R. 1867. Lehrbauch der Aufbereitungskunde. Berlin: Ernst and Korn.

Size Enlargement

5.1 Introduction: Agglomeration Processes

Size enlargement operations are used in the materials processing industry with different aims, such as improving handling and flow, reducing losses, producing specific shapes, boosting appearance, and so on. Processes involving any kind of size enlargement receive diverse denominations including compaction, granulation, encapsulation, pelletizing, agglomeration, and sintering. Some of these terms may represent synonyms referring to the same operation but applied in different industries, while others define operations employed in very specific processes, such as sintering in metallurgy. A generic classification could divide size enlargement operations into two main categories: those operations in which the enlargement objective is to control functional properties such as rehydration, and those in which the enlargement objective is to produce defined forms such as tablets or pellets. In a broad sense the former could be defined as tumbling agglomeration, whereas the latter could be referred to as pressure compaction.

5.2 Aggregation Fundamentals: Strength of Agglomerates

Agglomeration may be defined as the process of bonding individual particles in a random way, in order to end with an aggregate considerably larger in size and with a porous structure. As previously mentioned, agglomeration comprises a variety of operations in which particles are grouped for different purposes (Green and Perry, 2008). Figure 5.1 shows some common mechanisms of agglomerate binding that originate in coordination systems, much as those observed in crystallization arrangements (Pietsch, 1991). Each particle interacts with neighboring particles, while points of interaction may be kept together by physical contact or by attractive forces due to the proximity of such points. The number of interaction sites of each particle within the agglomerate is known as coordination number. In reality, agglomerates

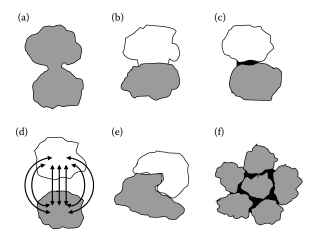


FIGURE 5.1 Different binding mechanisms in agglomeration: (a) partial melting sinter bridges, (b) chemical reaction hardening binders, (c) liquid bridges hardening binders, (d) molecular and liketype of forces, (e) interlocking bonds, and (f) capillary forces.

consist of a huge number of particles, which makes it difficult to estimate their coordination number. Indirect measurement of the coordination number can be made as a function of other properties of the agglomerate. In regular packs of monosized spherical particles the coordination number k and the porosity or void volume ϵ , are related by

$$k\varepsilon \approx \pi$$
 (5.1)

Equation 5.1 gives an appropriate approximation of coordination numbers for ideal agglomerate structures. Table 5.1 lists coordination numbers calculated from Equation 5.1 and compares them with coordination numbers of ideal structures, like some illustrated in Figure 5.2.

TABLE 5.1Geometric Arrangement, Porosity, and Coordination Number of Packings of Monosized Particles

	Coordination			
Geometric Arrangement	Porosity (ε)	Number (π/ϵ)	k	
Cubic	0.476	6.59	6	
Orthorhombic	0.395	7.95	8	
Tetragonal-spheroidal	0.302	10.39	10	
Rhombohedral (pyramidal)	0.260	12.08	12	
Rhombohedral (hexagonal)	0.260	12.08	12	

Size Enlargement 209

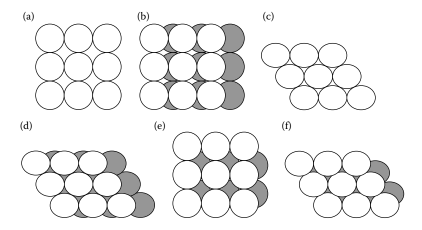


FIGURE 5.2 Packings of mono-sized spherical particles: (a) cubic, (b) and (c) orthorhombic, (d) tetragonal-spheroidal, (e) rhombohedral (pyramidal), (f) rhombohedral (hexagonal).

A generic relationship to describe the tensile strength of agglomerates σ_t held together by binding mechanisms acting at the coordination points is

$$\sigma_{t} = \frac{1-\varepsilon}{\pi} k \frac{\sum_{i=1}^{n} A_{i}(x,...)}{x^{2}}$$
 (5.2)

where A_i is the adhesive force due to a particular binding mechanism and x is the representative size of the particles forming the agglomerate.

Substituting Equation 5.1 into Equation 5.2, the following relation is obtained:

$$\sigma_{t} = \frac{1 - \varepsilon}{\varepsilon} \frac{\sum_{i=1}^{n} A_{i}(x, ...)}{x^{2}}$$
(5.3)

A further simplification results because many binding mechanisms are a function of the representative particle size *x* and thus

$$\sigma_{t} = \frac{1 - \varepsilon}{\varepsilon} \frac{\sum_{i=1}^{n} A_{i}(x, ...)}{x}$$
(5.4)

The ellipses within parentheses in Equations 5.2 through 5.4 indicate that A_i is also a function of other, unknown, parameters.

When liquid bridges have formed at the coordination points, A_i depends on the bridge volume and the wetting characteristics represented by the wetting angle. There are models available for predicting adhesion forces of various types (Pietsch, 1991), but A_i might be of different magnitude at each of the many coordination points, due to roughness or microscope structure of particulates forming the agglomerates.

The representative particle size most appropriate to describe the agglomeration process is the surface equivalent diameter, $x_{\rm sv}$ (Table 2.5), because porosity is surface dependant. As described in Chapter 2, such diameter is the size of a spherical particle which, if the powder consisted of only these particles, would have the same specific surface area as the actual sample. When determining the specific surface area, methods must be chosen that only measure the outer particle surface excluding the accessible inner surface due to open particle porosity. One of these recommended methods is permeability, also discussed in Chapter 2.

From the previous paragraphs, it can be gathered that the strength of agglomerate structures held together by bonding mechanisms is highly dependent on porosity and particle size, or, more properly, specific surface area. The relationship would be inversely proportional in both cases, that is, higher strength at lower porosities and lower surface areas. Tumbling agglomeration is often aided by adding small quantities of liquids, generally water. Depending on the degree of moisture, three states of agglomerate stability can be identified: pendular state when moisture is located only in the contact points of particles, funicular state when a layer of moisture covers the surface of particles, and capillary state when moisture fills the voids of a bed of particles (Green and Perry, 2008). Agglomerates that are completely filled with liquid obtain strength from the negative capillary pressure in the structure. When liquid bridges help maintaining the stability of an agglomerate, its strength can be calculated by

$$\sigma_{t} = c \left[\frac{1 - \varepsilon}{\varepsilon} \right] \left[\frac{\alpha}{x_{s} [f(\delta)]} \right]$$
 (5.5)

where c is a correction factor, α is the surface tension of the liquid, x_s is the surface equivalent diameter of the particle, and $f(\delta)$ is a constant related to the angle of contact of particles. Reported values of constant c vary between 2.8 and 8.0, depending on the degree of moisture trapped by agglomerates with the maximum value of 8.0 corresponding to the capillary state. Regarding values of $f(\delta)$, they also correlate to moisture content of agglomerates with a maximum value of unity corresponding to the capillary state.

For high-pressure agglomeration and the effect of matrix binders general formulas have not yet been developed. It can be considered, however, that the effects of variables would follow the trend described before, with porosity, particle surface, contact area, and adhesion, all playing an important role.

For nonmetallic powders, the following equation can be used to evaluate the needed applied pressure *p* to agglomerate:

$$\log p = mV_{\rm R} + b \tag{5.6}$$

where V_R represents the relationship V/V_s , being V the compacted volume at a given pressure and V_s the volume of the solid material to be compacted; m and b are correlation constants.

Example 5.1

Particles of $0.1~\mu m$ diameter are going to be agglomerated by mobile liquid bridges mechanism using water as the bonding agent. Calculate the strength of the agglomerate when the voids of the aggregates are totally wet. The porosity of the agglomerates is 0.65 and the surface tension of water is 72.8 dyne/cm.

SOLUTION

The agglomerate corresponds to the capillary state, with the voids completely wet. As previously mentioned, the values for such a case for c and $f(\delta)$ on Equation 5.5 are 8.0 and 1.0, respectively.

Substituting, thus, directly in Equation 5.5:

$$\sigma_{\rm t} = 8.0 \left\lceil \frac{(1 - 0.65)}{(0.65)} \right\rceil \left\lceil \frac{72.8}{(1 \times 10^{-5})(1)} \right\rceil = 3.92 \times 10^6 \text{ dyne/cm}^2$$

The strength of the agglomerate is, therefore, 3.92×10^6 dyne/cm².

Example 5.2

Calculate the volume of a tablet made of compressed ceramic powder by pressing it in a 54 cm^3 cavity applying a pressure of 316 MPa. The process of pressurization followed the model described by Equation 5.6, and the correlation constants m and b were determined as -4.402 and 4.1157, respectively.

SOLUTION

Substituting values directly on Equation 5.6, and then transposing V_R from such equation, the volumes relationship would be obtained as

$$\log(316) = -4.402(V_R) + 4.1157$$

$$V_{\rm R} = \frac{(2.4997 - 4.1157)}{-4.402} = 0.367$$

 $V_{\rm R}$ represents the volume of 1 compressed tablet of volume V, and so the solid before compression would be $V_{\rm s}$. Assuming that the uncompressed solid occupies the total volume of the press cavity (54 cm³), the volume of the tablet can be calculated as

$$V = V_{\rm R}V_{\rm s} = (0.367)(5.4 \times 10^{-5}) \,{\rm m}^3 = 1.98 \times 10^{-5} \,{\rm m}^3$$

Thus, the tablet volume would be 20 cm³ approximately.

5.3 Agglomeration Methods

With few exceptions, agglomeration methods can be classified into two groups: tumble/growth agglomeration and pressure agglomeration. Also, agglomerates can be obtained using binders or in a binderless manner. The tumble/growth method produces agglomerates of approximate spherical shape by buildup during tumbling of fine particulate solids, the resulting granules are at first weak and require binders to facilitate formation, and posttreatment is needed to reach final and permanent strength. On the other hand, products from pressure agglomeration are made from particulate materials of different sizes, and they are formed without the need of binders or posttreatment acquiring immediate strength.

5.3.1 Tumbling Agglomeration

The mechanism of tumble/growth agglomeration is illustrated in Figure 5.3. As shown, the overall growth process is complex and involves both disintegration of weaker bonds and reagglomeration by abrasion transfer and coalescence of larger units (Cardew and Oliver, 1985). Coalescence occurs at the contact point when, at impact, a binding mechanism develops which is stronger than the separating forces. Additional growth of the agglomerate may proceed by further coalescence, or by layering, or both. The most important and effective separation force counteracting on the bonding mechanism is the weight of the solid particle. For particles below 10 µm approximately, the natural attraction forces, such as molecular, magnetic, and electrostatic, become significantly larger than the separation forces due to particle mass and external influences. In such a way, natural agglomeration occurs. The mechanism of tumble/growth agglomeration is similar to that of natural agglomeration. The particles to be agglomerated are larger, however, and so the particle-to-particle adhesion would need to be increased by addition of binders, such as water or other more viscous liquids, depending on the properties of particles being agglomerated and the required strength of the agglomerate structure. On the other hand, the collision probability may be enhanced by providing higher particle concentration.

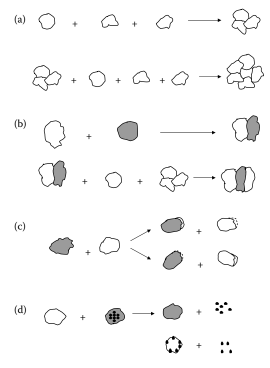


FIGURE 5.3 Kinetics of tumble/growth agglomeration: (a) nucleation, (b) random coalescence, (c) abrasion transfer, and (d) crushing and layering.

The conditions needed for tumbling agglomeration can be provided by inclined discs, rotating drums, any kind of powder mixer, and fluidized beds (Figure 5.4). In general terms, any equipment or environment creating random movement is suitable for carrying out tumbling agglomeration. In certain applications, very simple tumbling motions, such as on the slope of storage piles or on other inclined surfaces, are sufficient for the formation of crude agglomerates. The most difficult task of tumble/growth agglomeration is to form stable nuclei due to the presence of few coordination points in small agglomerates. Also, since the mass of particles and nuclei are small, their kinetic energy is not high enough to cause microscopic deformation at the contact points that enhances bonding. Recirculation of undersized fines provides nuclei to which feed particles adhere more easily to form agglomerates. In the whole process, tumble/growth agglomeration renders first weak agglomerates known as green products. These wet agglomerates are temporarily bonded by surface tension and capillary forces of the liquid binder. This is the reason why, in most cases, tumble/growth agglomeration requires some sort of posttreatment. Drying and heating, cooling, screening, adjustment of product characteristics by crushing, rescreening, conditioning, and recirculation of undersize material, are some of the processes which have

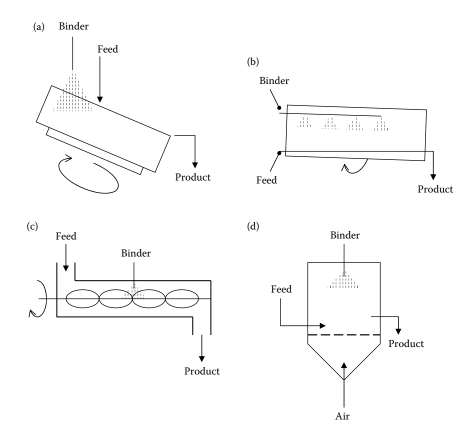


FIGURE 5.4 Equipment for tumble/growth agglomeration: (a) inclined rotating disc, (b) inclined rotating drum, (c) ribbon powder blender, and (d) fluidized bed.

been used as posttreatment in tumble/growth agglomeration. Sometimes, a large percentage of recycle must be rewetted for agglomeration and needs to be processed again, causing economical burden to this technology (Pietsch, 1983). For larger size or mass of the particles being agglomerated by tumble/growth methods, the forces trying to separate newly created bonds during growth become significant, until further size enlargement by tumbling is not possible, even if strong binders are added. There is, therefore, a definite limitation on the coarseness of a particle size distribution, being in the range of x_s between 200 and 300 μ m.

5.3.2 Pressure Agglomeration: Compaction

In contrast to tumbling agglomeration where no external forces are applied, in pressure agglomeration pressure forces act on a confined mass of particulate solids, which is then shaped and densified (Engelleitner, 1994). Pressure

agglomeration is normally carried out in two stages. The first one comprises a force rearrangement of particles due to a little applied pressure while a second step consists on a steep pressure rise during which brittle particles break and malleable particles deform plastically (Pietsch, 1994). The mechanism of pressure agglomeration is illustrated in Figure 5.5. There are two important phenomena that may limit the speed of compaction and, therefore, the capacity of the equipment: compressed air in the pores and elastic spring-back. Both can cause cracking and weakening which, in turn, may lead to destruction of the pressure-agglomerated products. The effect of these two phenomena can be reduced if the maximum pressure is maintained for some time, known as dwell time prior to its release.

Pressure agglomeration can be performed employing a low-, a medium-, or a high-pressure mode. When utilizing low- or medium-pressure agglomeration, relatively uniform agglomerates can be obtained. Under these conditions, the porosity of the feed material is changed but no change in particle size or shape occurs. The feed mixture is often prepared with fine particles and binders giving, thus, a sticky mass which may be formed by forcing it through holes in differently shaped screens or perforated dies. Agglomeration and shaping are, therefore, due to pressure forcing the material through the holes, as well as by frictional forces. High-pressure agglomeration is characterized by a large degree of densification, resulting in low product porosity. Typically, the products from high-pressure agglomeration feature high strength immediately after discharge from the equipment. To increase strength a bit further, addition of small amounts of binders or use of post-treatment methods are possible. High-pressure agglomeration is considered

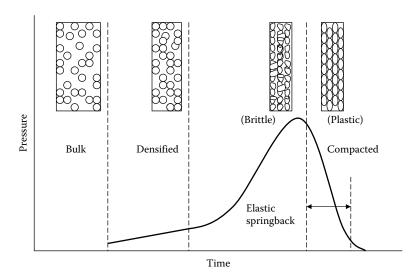


FIGURE 5.5 Mechanism of pressure agglomeration.

a versatile technique by which particulate material of any kind and size, that is, from nanometers to centimeters, can be successfully processed under certain conditions.

Pressure agglomeration can be carried out in different types of equipment. Generally, low- and medium-pressure agglomeration is achieved in extruders including the screen extruder, the screw extruder, and the intermeshing gears extruder. On the other hand, high-pressure agglomeration is performed in presses such as the punch-and-die press, the compacting roller press, and the briquetting roller press. Low- and medium-pressure agglomeration yield relatively uniform agglomerates of elongated spaghetti-like or cylindrical shape, whereas high-pressure agglomeration produces pillow- or almond-like shapes. Figure 5.6 presents equipment used for low- and medium-pressure agglomeration, while Figure 5.7 illustrates some common machinery for high-pressure agglomeration.

5.3.3 Equipment Operation Variables

On tumbling agglomeration, an important factor in operation would be the quantity of bonding liquid to add to the process in order to obtain resistant and functional agglomerates. On the other hand, due to agglomerates being formed as a consequence of gentile particle contact in the tumbling process, rotation speed of the agglomerating units would also be crucial. Optimum rotation speed should guarantee sufficient time to promote interparticle contact, but not so much time as to risk agglomerate stability.

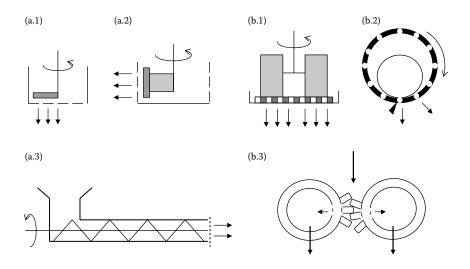


FIGURE 5.6 Equipment used for (a) low- and (b) medium-pressure agglomeration: (a.1) screen extruder, (a.2) basket extruder, (a.3) cylindrical-die screw extruder, (b.1) flat-die extruder, (b.2) cylindrical-die extruder, (b.3) intermeshing-gears extruder.

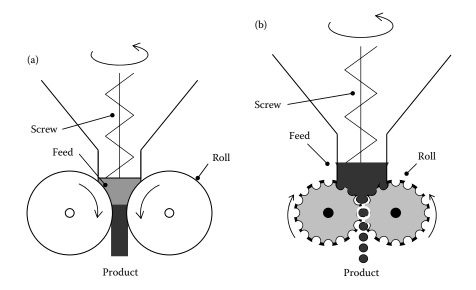


FIGURE 5.7 Equipment used for high-pressure agglomeration: (a) compacting roller press, (b) briquetting roller press.

Appropriate liquid content, expressed as fraction *X*, for an optimum agglomeration process, can be represented by the following relation:

$$X = \frac{1}{\left[1 + (1 - \varepsilon)\rho_{s}/(\varepsilon)(\rho)\right]}$$
 (5.7)

where ϵ is the porosity of the material, ρ_s is the solid density, and ρ is the liquid density.

In terms of optimum rotation speed of tumbling agglomeration units, normally rotating drums, this can be calculated by

$$N = \frac{42.3}{\sqrt{D}}$$
 (5.8)

where N is the rotational speed in rev/min, and D is the diameter of the drum in m.

For rotating discs used in agglomeration purposes, the power *P* and the mass flow rate *Q* can be calculated by the following equations:

$$P = k_1(D)^2 (5.9)$$

$$Q = k_2(D)^2 (5.10)$$

where k_1 and k_2 are empirical constants, while D is the disc diameter.

Example 5.3

A weight of 1600 kg of a 1230 kg/m 3 granular solid need to be agglomerated in a rotating drum, 3 m long and 1.5 m diameter. Water is used as the agglomerating aid and 1/3 of the drum capacity is recommended for optimum operation. Calculate the moisture percentage necessary to run the operation.

SOLUTION

The moisture fraction can be determined by substituting in Equation 5.7, while porosity e relates volume of air within the bed with the total volume. It is, therefore, required to calculate the volume of the solids bed, as a third of the volume of the drum, that is,

$$V_{\text{total}} = (\pi)(r)^2(h) = (3.1416)(0.75)^2 \,\text{m}^2(3) \,\text{m} = 5.3 \,\text{m}^3$$

$$V_{\text{effective}} = \frac{V_{\text{total}}}{3} = \frac{5.3 \,\text{m}^3}{3} = 1.767 \,\text{m}^3$$

On the other hand, the volume of solids occupying a third of the drum volume may be determined from its mass and its density, that is,

$$V_{\text{solid}} = \frac{m}{\rho_{\text{s}}} = \frac{1600 \text{ kg}}{1230 \text{ kg/m}^3} = 1.3008 \approx 1.3 \text{ m}^3$$

The air volume within the bed of particles would, therefore, be

$$V_{\text{air}} = V_{\text{effective}} - V_{\text{solid}} = 1.767 - 1.3 = 0.467 \,\text{m}^3$$

while the porosity is

$$\varepsilon = \frac{V_{\text{air}}}{V_{\text{effective}}} = \frac{0.467 \,\text{m}^3}{1.767 \,\text{m}^3} = 0.2643$$

Finally, using the value of porosity calculated above, all terms for substitution in Equation 5.7 are available and, thus

$$X = \frac{1}{\frac{1 + (1 - 0.2643)(1230) \text{ kg/m}^3}{(0.2643)(1000 \text{ kg/m}^3)}} = 0.1238$$

And, so, adding 12.38% moisture would give the best possible results in the agglomeration process.

5.4 Selection Criteria for Agglomeration Methods

There is a large variety of techniques and equipment available to carry out agglomeration duties in the materials processing industries. Some guidelines are given in the literature or can be provided by manufacturers. Table 5.2 summarizes some of the preliminary considerations to start a selection process of an agglomeration method by a practicing engineer. In general terms, features of the feed, the product, and the method are most important to be considered in the selection process (Pietsch, 1991).

5.4.1 Raw Materials Characteristics

The first characteristics of feed to be considered are particle size and particle size distribution. A limit in the range of few hundred micrometers defines the applicability of methods using growth mechanisms based on coalescence in moving beds of particles. With regards to larger particles, which may also refer to seed agglomerates, they can only be incorporated if an adequate amount of binder or enough small particles are present. Since small particles embed in larger ones, the strength of the agglomerate is caused by the matrix of fine powder in this case. In general terms, is difficult to agglomerate narrow particle-size distributions or monosized particles. Adding a binder can cause relatively large particles to agglomerate. It may be more economical, however, to crush larger particles in order to obtain material suitable for growth agglomeration. This is particularly crucial when it is desirable a highporosity product. Pressure agglomeration is more suitable for larger feed size particles, for example, sand-like material or particles up to 20–30 mm large. Since the external forces acting upon the mass result in particle disintegration or deformation, the upper limit of feed particle size is determined more by restrictions of the feeder than in the ability to agglomerate. In most cases consolidation occurs in a short period, and so considerable amount of air must be removed during compaction in order to obtain sound agglomerates. There is an increasing resistance to flow with decreasing particle size

TABLE 5.2General Guidelines for Selecting Agglomeration Processes

Feed Variables	Product Variables	Method Variables	
Size and size distribution	Size and shape of aggregate	Batch or continuous	
Moisture content	Green agglomerate resistance	Capacity	
Material characteristics	Cured agglomerate resistance	Dry or moist operation	
Material specific features	Porosity and related features	Simultaneous process	
Secondary properties		Energy	
Binding characteristics		Costs	

due to the smaller pore radii. For this reason, fine bulk solids of about 150 μm or less, can only be agglomerated by pressure methods if certain preconditions, such as low speed and dwell time, are established.

Free moisture can play an important role in growth agglomeration by coalescence. In such a case, moisture provides the binder or prevailing binder mechanism. The maximum volume of liquid must not be more than 95% of anticipated agglomerate porosity. Wet agglomeration is sensitive to this limit because a small excess of moisture will cause the entire mass to turn to mud. For the case of fluidized-bed agglomerators, which may also act as driers, the moisture content is less critical because it has to be high enough to make the feed pumpable. In pressure agglomeration moisture must be kept low being, sometimes, a precondition a completely dry feed. Due to the extreme compression forces in high-pressure agglomeration, crushing, rearrangement, and deformation of the solid take place and result in a considerable reduction of porosity. Excess water is either squeezed out or remains in the mass as an incompressible component, with the effect of resultant low strength of the agglomerate.

Material characteristics such as chemical properties, particle density, brittleness, elasticity, plasticity, wettability, and abrasivity, all play important roles in the choice of an agglomeration method. Particular chemical characteristics may be necessary to bring about the required chemical bonding, or may be incompatible with certain conditions of a method, such as addition of water or other liquids in most growth agglomeration techniques. Density of the feed particles determines particle weight and other field forces which may be counteractive to addition by coalescence. Brittleness, elasticity, plasticity, and abrasivity are most important for pressure agglomeration and of less concern for growth methods. Wettability, on the other hand, is paramount for all agglomeration methods using surface tension and capillary forces in the growth regime. Wetting of particle surfaces is a requirement for green strength.

Bulk properties, such as bulk density and flowability, as well as temperature, can be adjusted prior to agglomeration in order to improve size enlargement. High bulk density and unacceptable flowability are sometimes correlated by using two agglomeration methods in series. To reduce negative impact of these two factors, some sort of preconditioning should be necessary. For example, in some compacting processes fine feed powders are preagglomerated to reduce the compaction stroke and improve the flow of feed into a die. This increases the speed of the turret for rotary table tabletting machines. At the same time, this technique avoids segregation of the feed mix by stabilizing the blend in a granular form. Finally, the binding features of a given material must be considered for a possible agglomeration process. The binding characteristics should be assessed in order to decide whether agglomeration can be carried out binderless, due to the presence of an inherent binder in the material, or with the addition of binders such as water and some other liquids.

5.4.2 Product Properties

Some properties such as shape, dimensions, and particle size distribution of the agglomerated product, also influence proper selection of a suitable method. Agglomerated products are normally expected to improve in properties such as free flowing characteristics or dust-free features. Granular, free-flowing, dust-free products can be manufactured using almost all methods of size enlargement. The task of narrowing down the size distribution of the discharge is done by screening out under- and oversized components. While the fines are recirculated to the agglomeration unit, oversized particles are crushed and either rescreened or directly recirculated with the fines. Granular products can also be obtained by crushing and screening large agglomerates using criteria such as product porosity, product density, solubility, or reactivity.

The shape of the product is another important property. Spherical products are often desirable from an agglomeration operation, and such shape can be obtained using all growth agglomeration methods. Contrastingly, spherical products cannot normally be obtained with high-pressure agglomeration, unless extremely accurate feed control can be established. By using some types of pressure agglomeration equipment, such as tabletting machines, approximations to the spherical shape such as pillow-, lens-, or almond-shaped compacts can be obtained.

In terms of strength, such property is relevant for the final product, but also plays a role during the size enlargement operation. In growth agglomeration green agglomerates are formed first and then must be cured to obtain permanent bonding. A weak state could exist when the binding mechanism of the green agglomerate disappears before the permanent, cured bond sets in. Unless large amount of matrix binders are used, or agglomerates are cured at extremely high temperatures or by some chemical reaction, growth agglomeration products will be normally weaker than pressure agglomeration ones.

Pertaining porosity, different strength levels develop primarily because agglomerates growing by coalescence feature higher porosity than those from pressure agglomeration. Materials which may disperse easily, and are only agglomerated to improve handling of the intermediate product, should just have enough strength to survive their short existence. In some other cases a large specific surface is more important than high density and strength. Normally, an increase in external forces acting on the particulate matter during size enlargement will cause porosity and related characteristics to decrease, and density and strength to increase.

5.4.3 Technique Options

Agglomeration processes can be carried out in a batch or a continuous manner, depending on specific requirements and applications. Batch modes are

generally low in capacity, but are characterized by a better control than that exercised in a continuous process. Most large-volume applications operate in a continuous form, but may be accompanied by significant variations in quality. In growth agglomeration uncontrolled buildup must be removed, whereas in pressure agglomeration worn parts must be replaced. Most of the growth methods are wet processes using bind liquids for forming green agglomerates, while, normally, high-pressure techniques are operated as dry processes. Agglomeration can be sometimes carried out simultaneously with some other process. Simultaneous processing happens in mixer–granulators, granulator–driers, or even mixer–granulator–driers. Mixers are often also granulators in which both processes occur in different zones. In fluidized-bed granulators, however, agglomeration and drying can take place simultaneously.

Agglomeration shows potential for diverse applications such as recycling of wastes containing valuable ingredients or disposal of particulate wastes without value in an environmentally safe and acceptable way. These applications cannot always find economic justification as they must be typically performed in compliance with legislation. Finely divided particulate material is often released to the environs in many dry-operated processes. Often, these fines are precipitated or removed by pollution control devices, but recontamination of the environment is an obvious concern and is normally regulated. In these cases, agglomeration methods can be employed to obtain a size-enlarged material in order to handle and dispose it in a convenient way. Since, in most cases, one of the reasons for size enlargement is improvement of material handling, an agglomeration facility must be located near the particulate solid source. A suitable method must consider, therefore, the availability and cost of utilities and ancillary devices, such as binders or energy sources when wet granulation may need these requirements. Sometimes, since disposal may represent the main aim of the process, the same task may be accomplished using roller presses for dry compaction, and granulation by crushing and screening.

5.5 Design Aspects of Agglomeration Processes

Agglomeration processes consist on varied operations within a complete system aimed at obtaining a desired product. Mixing and screening are two common operations which are part of an agglomeration process. There are some other varied posttreatments, such as those previously mentioned that are used, mainly, to give strength to the agglomerate.

In tumble/growth agglomeration, if more than one feed powder is treated, the components must be metered and premixed. Homogenization may also be necessary due to the risk of selective agglomeration as the particulates

fed would involve many different sizes. During mixing, some of the liquid or dry binders could be added. It is also possible to feed all or part of the recycle into the mixer. Aeration of the premixed material is an important factor, so particles will be loose and able to move randomly, in order to pick up binder and agglomerate upon impact. A metered addition to the agglomerating equipment improves and accelerates agglomerate growth by seeding the charge. This is because recycle, despite its representing undersize product, consists largely of somewhat preagglomerated material. Control of the growth mechanism also requires addition of some of the liquid or dry binders in the agglomeration unit. Tumbling equipment produces green agglomerates that are better bonded by liquids. The agglomerate sizes and shapes are very varied ranging within wide limits. Sometimes it is possible to screen the green agglomerates and feed only a narrow particle size distribution to the posttreatment stage. The moist recycle should be sent directly to the agglomeration unit. Green agglomerates are often weak and sticky, so they tend to blind screen quite easily. For this reason, separation of over- or undersized material at this step may be avoided. The discharge from the posttreatment may also be screened as to remove fines which may be formed by abrasion and breakage or, contrastingly, in order to retain oversized agglomerates which may have developed by secondary agglomeration of the still-moist and sticky green agglomerates. Oversized agglomerates could then be crushed to obtain a recycle which is normally dried, and so it should be returned directly to the mixer and incorporated back to the process. In this way, tumble/growth agglomeration is an efficient process because recycling is doing continuously and losses are minimal. The problem of attrition and fines production is normally presented in handling and distribution of agglomerated products after being released from the agglomeration process. Such problem is severe in food products and has been thoroughly discussed in books dealing with the subject (Barbosa-Canovas et al., 2005; Onwulata, 2005).

With regard to pressure agglomeration, posttreatment is normally needed only for the case of low- and medium-pressure agglomeration. These methods typically require liquid binders to ensure easy formability. High-pressure agglomeration, on the other hand, does not include posttreatment and, in most applications, only dry additives may be added. In contrast to tumble/growth agglomeration which requires fine particulates as well as dispersion and aeration features, pressure agglomeration operates well using particles of wide size distributions and without aeration. In fact, aeration of the feed prior to agglomeration must be avoided to facilitate the operation, and the maximum particle size that can be handled increases with increasing pressure. Large particles do not segregate and are easily incorporated during the forming of the agglomerate under pressure. When high forces are applied as previously mentioned, brittle disintegration and plastic deformation occur. Also, a considerable volume reduction takes place with densification ratios as high as 1:5. Agglomerate strength increase with higher pressures during

densification and forming. Knives could be used to cut extrudates and diverse types of separators may be used to break strings of briquettes into single units. Pressure agglomeration can also be used to obtain granulate products. In this case, a separator is used as a prebreaker. The product is obtained between the two decks of double-deck screens. The oversized material is crushed and rescreened, while the undersized one is recirculated. Multiple step crushing and screening operations may be employed in order to improve the yield and obtain cleaner granular products.

5.6 Applications

Agglomeration applications are numerous and comprise a range of industries. For example, in metallurgical processes, sintering is used for obtaining agglomerates of different metallic minerals such as aluminum, copper, iron, gold, and silver. Pertaining nonmetals, diverse products such as salt, cement, asbestos, and ceramics, are agglomerated in some way. In the chemical industry, detergents, fertilizers, oxides, sulfites, and some other products are commercialized in some sort of agglomerate or compact form. The pharmaceutical industry produces many and diverse products as pellets, tablets, capsules, granules, and so on. Specific shapes of different sorts are extensively obtained by extrusion in industries as diverse the grain processing industry and the polymers industry. There are many other important applications of size enlargement in the food and feed industries. Feed and pet foods are compacted in different shapes and forms by compaction, extrusion, and so on. In the food industry, tumbling agglomeration is used in the food processing to improve reconstitutability of a number of products including flours, cocoa powder, instant coffee, dried milk, sugar, sweeteners, fruit beverages powders, instant soups, and diverse spices.

5.7 Laboratory Exercise: Comparing Methods for Tumbling Agglomeration of Powders

5.7.1 Introduction

Perhaps the most critical property of powder agglomerates is their ability to withstand postprocessing handling. Most of the agglomerates produced by tumbling and related techniques are friable, hence attrition represents a real problem. The objective of the exercise is to compare two agglomeration methods in terms of the strength of the aggregates they produce.

5.7.2 Equipment and Materials

Horizontal rotating drum and rotating tilting pan, at laboratory or pilot plant scale. Feed materials: fine chemicals such as limestone powder or silica flour. Binding agent: water.

5.7.3 Instruments

Light-scattering particle size distribution analyzer, and friability tester.

5.7.4 Measurement of Particle Size Distributions

Determine the particle size distributions of feed and product for the compared methods, preferably by light scattering, following principles and procedures described in Chapter 2.

5.7.5 Operation of the Drum Agglomerator

- Fill the drum with the recommended load, normally 5–15 kg, according to size and characteristics of the equipment.
- Start the rotating cycle at a chosen constant speed, again in terms of equipment capability. Typical rotation velocities are 10, 15, or 20 rpm.
- Operate the spraying system to moisten the load being tumbled.
- Run the experiment for 20–30 min, as a maximum. If the drum design allows, extract samples every 5 or 10 min to verify visually the progress of the agglomeration process.
- Stop the drum and collect two samples. Allow them to dry, either on an open tray or using any type of drier. Use one sample to determine the particle size distribution, and the other to run the friability test.

5.7.6 Operation of the Pan (Disc) Agglomerator

- Operate the pan at the recommended speed and continuously add the feed material with the water spaying functioning. Laboratory pan agglomerators run at variable speeds but generally in the range of 15–25 rpm.
- Keep adding feed at a rate required for the green agglomerates to start forming and maturing until reaching a size enough to be discharged.
- Collect two samples, like in the drum agglomerator experiment and follow the same procedure, that is, use the first sample to determine the particle size distribution and the second one to run the friability test.

5.7.7 Friability Test

- Weight the sample and introduce it in the drum of the tester.
- Run the tester, according to specific instructions of the apparatus. A number of applications give satisfactory results at 20–25 rpm for 20–30 min.
- Unload the drum (if not done automatically) and weigh the processed sample.

5.7.8 Calculations and Report

- Measure the particle size distribution of the agglomerates from the two methods being tested. Make calculations to determine which method worked best in terms of producing larger agglomerates.
- Pertaining strength of agglomerates, from friability testing determine friability as the difference of weighed sample before and after the test. Express the friability in percentage of weight loss.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out. Include analysis from own data and, as much as possible, discussion based on literature survey from studies and investigations on the subject.

PROBLEMS

- 1. Calculate the strength of an agglomerate of particles that may be considered spherical in shape with a mean equivalent diameter of 10.4 mm. The union strength per contact point is 2.4×10^{-7} kg, and the agglomerate is geometrically arranged in a simple cubic manner.
- 2. Determine the tensile strength of a fraction of detergent particles agglomerate retained in a 200 mesh (Tyler) sieve. The agglomerate packings tend to adopt an orthorhombic geometry and the bonds stabilize in a pendular state. Water has been used as the bonding agent, and an estimate of the constant related to the angle of contact of particles represents a third of its maximum value.
- 3. Compute the rate, in kg/h, of a 12 in. (30.48 cm) pan agglomerator. The empirical constant of the relationship describing the agglomerator operation has been reported as 55.8 kg/s m².

References

Barbosa-Cánovas, G. V., Ortega-Rivas, E., Juliano, P., and Yan, H. 2005. Food Powders: Physical Properties, Processing, and Functionality. New York, NY: Kluwer Academic/Plenum Publishers.

Cardew, P. T. and Oliver, R. 1985. Kinetics and mechanics in multiphase agglomeration systems. Notes of course on agglomeration fundamentals. *4th International Symposium on Agglomeration*. Toronto University, Waterloo, Ontario, Canada.

- Engelleitner, W. H. 1994. *Method Comparison*. Notes of Course on Briquetting, Pelletizing, Extrusion, and Fluid Bed/Spray Granulation. The Center for Professional Advancement. Chicago, IL.
- Green, D.W. and Perry, R.H. 2008. *Perry's Chemical Engineers' Handbook*, 8th Ed. New York, NY: McGraw-Hill.
- Onwulata, C. W. 2005. *Encapsulated and Powdered Foods*. Boca Raton, FL, USA: CRC Taylor & Francis.
- Pietsch, W. 1983. Low-energy production of granular NPK fertilizers by compaction-granulation. *Proceedings of Fertilizer '83*, pp. 467–479. London: British Sulphur Corp.
- Pietsch, W. 1991. Size Enlargement by Agglomeration. Chichester, UK: John Wiley & Sons.
- Pietsch, W. 1994. Parameters to be considered during the selection, design, and operation of agglomeration systems. *Preprints of 1st International Particle Technology Forum, Part I,* pp. 248–257. New York, NY: AIChE.

This page intentionally left blank



Mixing

6.1 Introduction

The unit operation in which two or more materials are interspersed in space with one another is one of the oldest and yet one of the least understood of the unit operations of process engineering. Mixing is used in the materials processing industry with the main objective of reducing differences in properties such as concentration and color, between different parts of a system. Since the components being mixed can exist in any of the three states of matter, a number of mixing possibilities arise. The mixing cases involving a fluid, for example, liquid–liquid and solid–liquid, are most frequently encountered and so they have been extensively studied. Despite the importance of the mixing of granular and particulate materials in many processing areas, fundamental work of real value to either designers or users of solids mixing equipment is still relatively sparse. It is through studies in very specific fields, such as powder technology and multiphase flow that important advances in understanding of mixing of solids have been made.

Mixing is more difficult to define and evaluate with powders and particulates than it is with fluids, but some quantitative measures of dry solids mixing may aid in evaluating mixer performance. In actual practice, however, the proof of any mixing operation efficiency lies in the properties of the mixed material it produces. A significant proportion of research efforts are directed toward the development of new and novel mixing devices for finely divided solids. It is difficult, however, to generalize or standardize solids mixing operations for diverse equipment designs and specific applications. There are neither many mathematical models, nor numerous scaling and design procedures developed for solid–solid mixing operations. Relationships between mixing operation variables and mixing quality indices are, therefore, difficult to derive for the case of solids blending (Niranjan and de Alwis, 1993; Niranjan, 1995).

6.2 Blending Mechanisms

Three mechanisms have been recognized in solids mixing: convection, diffusion, and shear. In any particular process one or more of these three basic mechanisms may be responsible for the course of the operation. In convective mixing masses or group of particles transfer from one location to another, in diffusion mixing individual particles are distributed over a surface developed within the mixture, while in shear mixing groups of particles are mixed through the formation of slipping planes developed within the mass of the mixture. Shear mixing is sometimes considered as part of a convective mechanism. Pure diffusion, when feasible, is highly effective producing very intimate mixtures at the level of individual particles but at an exceedingly slow rate. Pure convection, on the other hand, is much more rapid but tends to be less effective, leading to a final mixture that may still exhibit poor mixing characteristics on a fine scale. These features of diffusion and convective mixing mechanisms suggest that an effective operation may be achieved by combination of both, in order to take advantage of the speed of convection and the effectiveness of diffusion.

Compared with fluid mixing, in which diffusion can be normally regarded as spontaneous, particulate systems will only mix as a result of mechanical agitation provided by shaking, tumbling, vibration, or any other mechanical mean. Mechanical agitation will provide conditions for the particles to change their relative positions either collectively or individually. The movement of particles during a mixing operation, however, can also result in another mechanism which may retard, or even reverse, the mixing process and is known as segregation. Since two or more different powders would consist of particles differing in physical properties, particularly size and/or density, solids mixing would be normally accompanied by a tendency to unmix. Thus, in any mixing operation mixing and demixing may occur concurrently and the intimacy of the resulting mix depends on the predominance of the former mechanism over the latter. Apart from the properties already mentioned, surface properties, flow characteristics, friability, moisture content, and tendency to cluster or agglomerate may also influence the tendency to segregate. The closer the feed materials are in size, shape, and density, the easier the mixing operation and the more intimacy of the final mix. A particular powdered solids mixing process can be considered analogous to a chemical reaction trying to reach an equilibrium state. Once the mixing and demixing mechanism reach a state of equilibrium, the condition of the final mix is determined and further mixing will not produce a better result.

When particles having significant different densities are mixed, it can be observed that the denser particles tend to settle to the bottom of the mixture to lower, presumably, the total potential energy of the system. On the other hand, if coarse and fine particles are set in motion, the fines tend to segregate

to the bottom by a possible percolation mechanism, in which the fines can pass through the interstices between larger particles. Williams (1968/1969) also showed that a single large particle placed on a bed of smaller particles will, on vibrating the bed, tend to rise toward the bed top, even if its density is greater than that of the finer material. An explanation to this is that the large solid particle, which will be generally denser than the loosely packed bed, causes a compaction of the bed immediately beneath it. In consequence, the freedom of movement of such large particle, in response to vibration, will be restricted to the lateral and upward directions, and the net result will be a tendency to rise.

Due to unusual behavior in the mixing and segregation phenomena in solids blending, such as that discussed above, segregation theories would only be applicable to specific situations. A general theory of segregation, regardless of the particular circumstances in which the operation take place, has not yet been offered to explain the segregation phenomena in particulate systems. In any blending operation the mixing and demixing mechanisms will be acting simultaneously. The participation of each of these two sets of mechanisms will be dictated by the environment and the tendency of each component to segregate out of the system. The importance of segregation on the degree of homogeneity achieved in solids mixing cannot be overemphasized. Any tendency for segregation to occur must be recognized when selecting solids mixing equipment. Segregation in a mixture of dry solids is readily detected by the use of a heap test. A well-mixed sample of the solids is poured through a funnel so as to form a conical heap. Samples taken from the central core and from the outside edge of the cone should have essentially the same compositions if segregation is not to be a problem. When the two samples have significantly different compositions, it can be assumed that segregation would be very likely to occur unless a very careful choice of equipment is made. It is generally accepted that the efficiency of a mixing process must be related to both the flow properties of the components, and to the selection or design of the mixer.

6.3 Statistical Approach of Mixing Processes

In the mixing of particulate solid materials the probability of getting an orderly arrangement of particles, which would represent the perfect mixing, is virtually zero. In practical systems the best mixture attainable is that in which there is a random distribution of the ingredients. An ideal random distribution of two solid components in equal proportions would resemble a chess board, that is, white and black squares in a perfect alternate pattern. In practice, however, a perfectly random mixture is commonly defined as one in which the probability of finding a particle of a constituent of the mixture

is the same for all its points. Over the years many workers have attempted to establish criteria for the completeness and degree of mixture. In order to accomplish this, very frequent sampling of the mix is usually required and, tending to be statistical in nature, such an exercise is often of more interest to mathematicians than to process engineers. Thus, in practical mixing applications, an ideal mixture may be regarded as the one produced at minimum cost and which satisfies the product specifications at the point of use.

6.3.1 Sampling

Sampling is a crucial step in the mixing process because any form of control of mixing operations involves sampling procedures. The sample must be representative of the mixture and postsampling handling must not alter it. As sampling has a statistical aspect, sampling procedures following a pure mathematician approach are not completely practical in industrial situations. The confidence that can be placed by any results obtained from the sampling and analysis of a mixture is greatly influenced by several factors including the method of sampling, the number of samples, the size of the sample, and the location in the bulk material from which the sample is taken. If sampling is not performed carefully, every mixture determination could be considered meaningless and mixing quality would be doubtful. Due to the crucial aspect of sampling in the different aspects of particle technology, some texts (Barbosa-Cánovas et al., 2005) include complete chapters on sampling. Suggestions for taking a representative sample when dealing with solid-solid mixing processes are practically the same given for sampling in characterization of particulate solids (Chapter 2). For the specific case of solid-solid mixing, Harnby (1985) has recommended collecting samples from the outflow of a mixer, in the moving stream, instead of taking them from a static mass inside the mixer. In such a way, the possibility of bias in sample retrieval is minimized.

It can be demonstrated by statistical means, that the larger the number of samples the more reliable the results. For example, using statistical theory of sampling, it is stated that a most representative sample would approach an infinite number of samples. In other words, the only way of including every member of a population being sampled is taking this whole population as a sample. Such an approach would be, of course, unfeasible and unreasonable for practical purposes in mixing of industrial powders. As discussed and described in Chapter 2 (Section 2.3), an optimum number of samples for many applications in particle technology would be between 20 and 50, so as to obtain representative results. The size of the sample is also important. If a simple particle is drawn from the mixture no mixing is evident. In contrast, if the whole mixture were to be analyzed, provided the ingredients were present in the correct proportions, complete homogeneity would appear to be achieved. As both of these extremes are impractical and unreliable, the recommended sample volume, often called the scale of scrutiny or characteristic

sample size, falls between them and it is defined as the size of sample that may be taken to correspond with the product usage. In animal feed manufacture, for instance, feed contains carbohydrates and proteins, balanced with added nutrients. In a particular feed, an animal must receive the correct balance of components. Provided that the required quantities of the necessary ingredients are present in the food consumed at each feed, intimate mixing is not essential. Thus, the volume of sample that would give such a balance would be the useful one, regardless of its perfection in statistical terms.

6.3.2 Blending Quality: Mixing Indexes and Mixing Rate

Considering some previously discussed aspects of the mixing process, it can be gathered that solid—solid mixing is a complicated task not easily described by mathematical modeling. Mixture quality results from several complex mechanisms operating in parallel which are hard to follow and fit to a particular model. Dankwertz (1952) has defined the scale and intensity of segregation as the quantities necessary to characterize a mixture. The scale of segregation is a description of unmixed components, while the intensity of segregation is a measure of the standard deviation of composition from the mean, taken over all points in the mixture. In practice is difficult to determine these parameters, since they require concentration data from a large number of points within the system. They provide, however, a sound theoretical basis for assessing mixture quality. Establishing general criteria for mixing indices is a difficult task, due to the segregating trend of the solids mixing process. On the other hand, as previously discussed, an endpoint of a mixing process depends on many factors, such as the scale of scrutiny.

The degree of uniformity of a mixed product may be measured by analysis of a number of spot samples. Powder mixers act on two or more separate materials to intermingle them. Once a material is randomly distributed through another, mixing may be considered to be complete. Based on these concepts, the well-known statistical parameters mean and standard deviation of component concentration can be used to characterize the state of a mixture. If spot samples are taken at random from a mixture and analyzed, the standard deviation of the analyses s about the average value of the fraction of a specific powder \bar{x} is estimated by the following relation:

$$s = \sqrt{\frac{\sum_{i=1}^{N} (x_i - \overline{x})^2}{N - 1}}$$
 (6.1)

where x_i is every measured value of fraction of one powder and N is the number of samples.

The standard deviation value on its own may be meaningless, unless it can be checked against limiting values of either complete segregation s_0 , or complete randomization s_r . The minimum standard deviation attainable with

any mixture is s_r and it represents the best possible mixture. Furthermore, if a mixture is stochastically ordered, s_r would equal zero. Based on these limiting values of standard deviations, Lacey (1954) defined a mixing index M_1 as follows:

$$M_1 = \frac{s_0^2 - s^2}{s_0^2 - s_r^2} \tag{6.2}$$

The numerator on Equation 6.2 would be an indicator of how much mixing has occurred, while the denominator would show how much mixing can occur. In practice, however, the values of s, even for a very poor mixture, lie much closer to s_r than to s_0 . Poole et al. (1964) suggested an alternative mixing index, that is,

$$M_2 = \frac{s}{s_r} \tag{6.3}$$

Equation 6.3 clearly indicates that for efficient mixing or increasing randomization the inverse of M_2 would approach unity. The values of s_0 and s can be determined theoretically. These values would be dependent on the number of components and their size distributions. Simple expressions can be derived for two-component systems, based on their fractions. For a binary mixture of two components of known volume an expression for s_r would be

$$s_{\rm r} = \sqrt{\frac{X_{\rm A}(1 - X_{\rm A})}{n}}$$
 (6.4)

where X_A is the weight fraction for a component, while n is the number of particles in the sample.

For a multicomponent particulate mixture Poole et al. (1964) demonstrated that

$$s_{\rm r}^2 = \frac{pq}{\left\lceil w/(q(\sum f_{\rm a} w_{\rm a})_{\rm p} + p(\sum f_{\rm a} w_{\rm a})_{\rm q}) \right\rceil}$$
(6.5)

where p and q are the proportions by weight of components within a total sample weight w, and f_a is the size fraction of one component of average weight w_a in a particle size range. For a given component in a multicomponent and multisized particulate system, Stange (1963) presented an expression for s_{rr} as follows:

$$s_{\rm r}^2 = \frac{p^2}{w} \left\{ \left[\frac{1-p}{p} \right]^2 \cdot p\left(\sum f_{\rm a} w_{\rm a}\right)_{\rm p} + q\left(\sum f_{\rm a} w_{\rm a}\right)_{\rm q} + r\left(\sum f_{\rm a} w_{\rm a}\right)_{\rm r} + \cdots \right\}$$
(6.6)

Equations 6.2 and 6.3 can be used to calculate mixing indices defined by Equation 6.1. Another suggestion for the characterization of degree of homogeneity in mixing of powders has been reviewed by Boss (1986), with the degree of mixing M_3 defined as

$$M_3 = 1 - \frac{s}{s_0} \tag{6.7}$$

Some other mixing indices have been reviewed by Fan and Wang (1975). McCabe et al. (2005) presented the following relationship to evaluate mixing time *t* for solids blending:

$$t = \frac{1}{k} \ln \frac{1 - 1/\sqrt{n}}{1 - 1/M_2} \tag{6.8}$$

where k is a constant and n is the number of particles in a spot sample. Equation 6.8 can be used to calculate the time required for any required degree of mixing, provided k is known and segregating forces are not active.

Mixing times should not be very long due to the, unavoidable, segregation nature of most particulate solids mixtures. Instead of improving efficiency, long mixing times often result in poor blending characteristics. A graph of degree of mixing versus time is recommended to select the proper mixing time quantitatively. A plot of this sort may look like the one shown in Figure 6.1. Most cases of mixing of powders will attain maximum degree of homogeneity in less than 15 min, when the proper type of machine and the working capacity have been chosen.

Example 6.1

Table 6.1 refers to a mixing process of sand and clay particles used to obtain a refractory material. Calculate the time needed for the mixing index to reach a value of 0.9. In every run the mixer is fed with a batch of 254 g clay poured over 300 g sand. Average number of particles in each sample is 120.

SOLUTION

To estimate a mixing time using Equation 6.8 the value of constant k should be determined empirically. This determined constant will be, then, used to calculate an approximate time for the mixing process. In this case two experimental runs are reported so k will be determined twice and an average value of it will be used for an estimate of time. From data in Table 6.1 and using Equation 6.1 values of s are calculated as

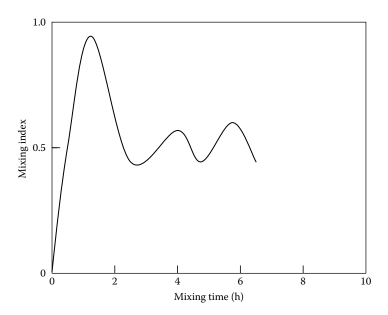


FIGURE 6.1 Representative plot of mixing time versus mixing index for solids.

$$s_1 = 0.1603$$

$$s_2 = 0.0992$$

Pertaining s_{rr} as defined by Equation 6.4, considering X_A as sand fraction according to Table 6.1, and substituting such value:

$$s_{\rm r} = \sqrt{\frac{0.5415(0.4585)}{120}} = 0.0455$$

TABLE 6.1 Example 6.1

Run 1	Time (s)	Sand Fraction									
		0.64	0.66	0.49	0.32	0.33	0.68	0.64	0.52	0.32	0.26
		0.74	0.77	0.49	0.33	0.31	0.63	0.70	0.54	0.35	0.32
		0.73	0.67	0.64	0.48	0.38	0.61	0.58	0.38	0.33	0.32
		0.62	0.87	0.49	0.44	0.35	0.70	0.60	0.52	0.39	0.36
2	87	0.50	0.48	0.49	0.42	0.33	0.54	0.67	0.53	0.35	0.36
		0.60	0.65	0.46	0.43	0.37	0.62	0.63	0.49	0.47	0.49
		0.60	0.62	0.58	0.59	0.48	0.55	0.46	0.34	0.38	0.37
		0.56	0.63	0.52	0.39	0.49	0.60	0.58	0.45	0.45	0.32
		0.59	0.48	0.50	0.52	0.32	0.63	0.59	0.47	0.39	0.36

Using, then, Equation 6.3 values of the mixing index M_2 are calculated for each experimental run as

$$M_{2-1} = \frac{0.1603}{0.0455} = 3.5321$$

 $M_{2-2} = \frac{0.0992}{0.0455} = 2.1802$

Using the corresponding values of the mixing index M_2 calculated above, and substituting in Equation 6.8 for each experimental run:

$$k_1 = \frac{1}{45} \ln \frac{1 - \left[\frac{1}{\sqrt{120}} \right]}{1 - (\frac{1}{3}.5231)} = 0.0053$$

$$k_2 = \frac{1}{87} \ln \frac{1 - \left[\frac{1}{\sqrt{120}} \right]}{1 - (\frac{1}{2}.1802)} = 0.0059$$

With average value of k = 0.0056 and the required optimum for the mixing index $(1/M_2) = 0.9$, substituting again in Equation 6.8:

$$t = \frac{1}{0.0056} \ln \frac{1 - 0.0913}{1 - 0.9} = 394.07 \,\mathrm{s}$$

Consequently, an optimum mixing time would be 394 s, approximately.

Example 6.2

In a mixing operation, 150 g wheat are blended with 250 g sorghum. Samples with an estimate number of 1500 particles are taken periodically to verify the progress of the operation. Calculate the optimum mixing time according to data reported in Table 6.2.

SOLUTION

This example follows the same direction as the previous one. There is not a preestablished ideal mixing index to be attained in this case, however, and so the

TABLE 6.2 Example 6.2

Run	Time (min)	Wheat Fraction
1	1.0	0.52 0.56 0.38 0.33 0.59 0.48 0.28 0.22 0.62 0.60 0.68 0.51 0.55 0.54 0.30 0.33 0.50 0.28 0.38 0.53
2	1.5	0.44 0.47 0.55 0.50 0.51 0.40 0.41 0.46 0.55 0.60 0.58 0.55 0.67 0.63 0.50 0.69 0.49 0.51 0.60 0.67
3	2.0	0.50 0.53 0.48 0.45 0.54 0.47 0.55 0.50 0.40 0.42 0.51 0.44 0.41 0.41 0.42 0.57 0.43 0.42 0.50 0.51

mixing index will be averaged just like the *k* value in Equation 6.8 will be also averaged from the experimental data given in Table 6.2. Therefore, using Equation 6.1, the three values for *s* from Table 6.2 are

$$s_1 = 0.1343$$

 $s_2 = 0.0855$
 $s_3 = 0.0529$

The value of s_{rr} considering X_A the fraction of wheat as indicated in Table 6.2 is, thus

$$s_{\rm r} = \sqrt{\frac{0.375(0.625)}{1500}} = 0.0125$$

Mixing indices M_2 corresponding to each experiment are then calculated from Equation 6.3 as

$$M_{2-1} = \frac{0.1343}{0.0125} = 10.744$$

$$M_{2-2} = \frac{0.0855}{0.0125} = 6.840$$

$$M_{2-3} = \frac{0.0529}{0.0125} = 4.232$$

Also, corresponding k values transposed from Equation 6.8 are

$$k_{1} = \frac{1}{1} \ln \frac{1 - \left[\frac{1}{\sqrt{1500}} \right]}{1 - \left[\frac{1}{10.744} \right]} = 0.0715$$

$$k_{2} = \frac{1}{1.5} \ln \frac{1 - \left[\frac{1}{\sqrt{1500}} \right]}{1 - \left[\frac{1}{6.840} \right]} = 0.0879$$

$$k_{3} = \frac{1}{2} \ln \frac{1 - \left[\frac{1}{\sqrt{1500}} \right]}{1 - \left[\frac{1}{4.232} \right]} = 0.1217$$

Using average values of k and M_2 and substituting into the same Equation 6.8, an approximate mixing time is calculated as follows:

$$t = \frac{1}{0.0937} \ln \frac{1 - \left[\frac{1}{\sqrt{1500}} \right]}{1 - \left[\frac{1}{7.282} \right]} = 1.29 \,\text{min}$$

The calculated mixing time of about 1.3 min was obtained for an averaged $1/M_2$ value very low, that is $(1/M_2) \approx 0.14$, considering that the ideal value should approach unity. Since the estimates of k and mixing indices cannot be taken

outside of the experimental data, the obvious conclusion is that experimental runs were all short in time, and more experimentation is needed at more prolonged times. From the trend shown in determination of k and M_2 values, it can be implied that an equilibrium mixing–segregation is far from being established within the reported times.

6.4 Mixing Equipment

In general terms, mixers for dry solids have nothing to do with mixers involving a liquid phase. According to the mixing mechanisms previously discussed, solids mixers can be classified into two groups: segregating mixers and nonsegregating mixers. The former operate mainly by a diffusive mechanism while the latter practically involve a convective mechanism. Segregating mixers are normally nonimpeller-type units, such as tumbler mixers, whereas nonsegregating mixers may include screws, blades and ploughs, in their designs, and examples of them include horizontal trough mixers and vertical screw mixers. Solids mixing can be carried out satisfactorily in equipment units serving other purposes in the materials processing industries. An example of this is the use of fluidized beds to perform solids mixing duties. Mixing times required in fluidized beds are significantly lower than those required in conventional powder mixers. Van Deemter (1985) discussed different mixing mechanisms prevailing in fluidized beds.

6.4.1 Tumbling Mixers

These mixing units operate by the principle of promoting blending by contact of masses of solids inside a revolving vessel. The vessels take various forms, such as those illustrated in Figure 6.2, and may be fitted with baffles or stays to improve their performance. The shells rotate at variable speeds (up to 100 rpm), and with working capacities around 50% to 60% of the total. They are manufactured using a wide variety of materials, including stainless steel. This type of equipment is best suitable for gentle blending of powders with similar physical characteristics. Segregation can represent a problem if particles vary, particularly in size and shape. Rotation speed may be fixed between 50% and 80% of the critical rotation speed $N_{\rm cr}$ given by the following relationship:

$$N_{\rm cr} = \frac{0.498}{\sqrt{R_{\rm max}}} \tag{6.9}$$

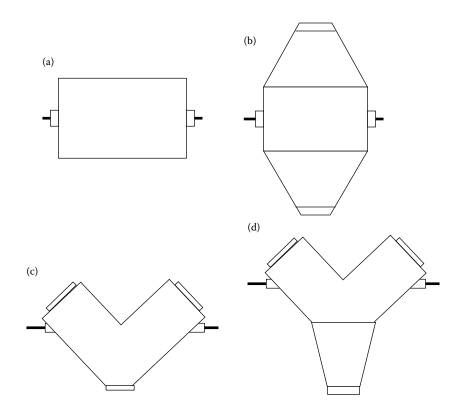


FIGURE 6.2 Tumbler mixers used in food powder blending: (a) horizontal cylinder, (b) double cone, (c) *V*-cone, and (d) *Y*-cone.

6.4.2 Horizontal Trough Mixers

Horizontal trough mixers consist of a semicylindrical horizontal vessel in which one or more rotating devices are located. For simple operations single or twin screw conveyors are appropriate and one passage through such a system may be good enough. For more demanding duties a ribbon mixer, like the one shown in Figure 6.3, may be used. A typical design of a ribbon mixer will consist of two counteracting ribbons mounted on the same shaft. One moves the solids slowly in one direction while the other moves it quickly in the opposite direction. There is a resultant movement of solids in one direction, and so the equipment can be used as a continuous mixer. Some other types of ribbon mixers operate on a batch basis. In these designs troughs may be closed, so as to minimize dust hazard, or may be jacketed to allow temperature control. Due to small clearance between the ribbon and the trough wall, this kind of mixers can cause particle damage and may consume high amounts of power.

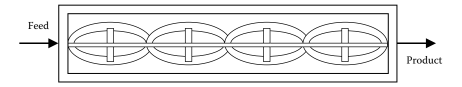


FIGURE 6.3 Plain view of an open ribbon mixer.

6.4.3 Vertical Screw Mixers

In vertical screw mixers a rotating vertical screw is located in a cylindrical or cone-shaped vessel. The screw may be mounted centrally in the vessel or may rotate or orbit around the central axis of the vessel near the wall. Such mixers are schematically shown in Figure 6.4a and b, respectively. The latter arrangement is more effective and stagnant layers near the wall are eliminated. Vertical screw mixers are quick, efficient, and particularly useful for mixing small quantities of additives into large masses of material.

6.5 Design and Selection Factors

Before selecting solids-mixing equipment, a careful study ought to be made of several performance characteristics. As previously stated, mixing of

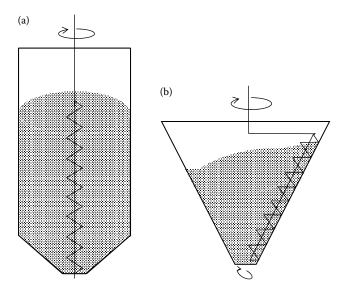


FIGURE 6.4 Vertical screw mixers: (a) central screw and (b) orbiting screw.

powdered solids is a complex operation and mathematical modeling can be hardly used. Many factors affect the operation of solids blending, and so process features such as mixing homogeneity and time, loading and discharging arrangements, power consumption, equipment wear, and so on need to be analyzed and properly weighed up in order to take the most convenient decision.

As has already been mentioned, blending uniformity and mixing rate are best evaluated using design graphs, bearing in mind the tendency to segregate and considering that long mixing times tend to worsen, rather than improving, efficiency. Solids mixing can be a batch or continuous operation. In batch mode, proper mixing design will produce the desired blend in few minutes. Determination of residence time in continuous operation is a more difficult task but, considering the main properties of solids blending, such times also tend to be short, in the order of few minutes or even seconds. The ribbon-type mixer is often used for continuous mixing, although it is also employed for batch mixing. Continuous mixing should be considered an option only if a single formulation can be run for an extended period, or when the fluctuations of the outgoing product are within process requirements. When any of these factors are compromised, the batch mode of operation would be preferred as to assure the most attainable mixing uniformity.

Loading and discharge are also important aspects, more critical in continuous operation mode. The total handling system must be considered in order to obtain optimum charging and discharging conditions. This would include the efficient use of weight hoppers and surge bins, minor-ingredient premixing, location of discharge gates, and any other ancillary device used to aid the continuity of the process.

Power requirements are not a major concern when choosing solids mixers since other considerations usually predominate. Sufficient power must be provided in order to handle the maximum needs, as well as to prevent changes during the mixing operation. When materials and operating conditions are subjected to variation, enough power should be made available for the heaviest bulk-density materials and for the extreme conditions of operation. If the loaded equipment is to be started from rest, there should be sufficient power for this. When speed variations may be desirable, this should be taken into account for power requirements planning.

The ease, frequency, and thoroughness of cleaning are crucial when batches of different nature are to be mixed alternatively in the same equipment. Plain tumbling vessels are easy to clean, provided that adequate openings are available. Areas which could be difficult to clean are seals or stuffing boxes, crevices at baffle support, any corners, and discharge arrangements. If cleaning between different batches is time consuming, several small mixers should be considered instead of a large single unit.

Dust formation should be avoided for safety reasons and when loss of dust may significantly affect batch composition. When dust release to the surrounding atmosphere cannot be avoided, the use of solid–gas separation

devices, such as cyclones or bag filters, should to be considered as part of the design. Minimization of dust formation can be achieved by using less dusty but equally satisfactory batch ingredients, by employing palletized forms of extremely dusty materials, by proper venting so as to enable filtering or displaced air rather than unregulated loss of dust-laden air, or by addition of liquids if tolerable. Water addition in small quantities would be effective in minimizing dust on discharge from the mixer, and it will also render the batch less dusty in subsequent handling stages. Water or any other liquid should be directed into the batch material instead of onto the bare surface of the mixer, since this could cause buildup. Spraying by using a nozzle would be the considered the most convenient way of incorporations moisture into the mixing batch. Nozzle spray pressure should be sufficient to penetrate the batch, but not so high as to cause heavy splashing. The liquid should be added to the well-mixed batch, particularly when premature addition of liquid could impair the adequacy of blending. Also, both time of addition and time of application are important and should be carefully considered. Quantity of added moisture should be carefully considered because an excessive amount may cause the solids being blended to deliver a product showing a paste-like consistency.

Equipment wear could be taken as a crucial issue especially with abrasive materials. Abrasion-resistant coatings, such as rubber coating, special alloys, or platings, need to be considered in these cases. Any internal agitator device may wear even when operating at slow speed. When high abrasive materials are to be mixed, the benefits of an agglomerated-breaking device must be weighed up against potential contamination and maintenance costs.

The capacity would be an important factor when the mixed batches vary considerably in size from time to time. There are some features of the mixing operation which are not flexible in terms of capacity. For example, certain agitation devices in tumbler mixers do not function properly unless a given capacity is kept. In general, the effect of percentage of mixer volume occupied by the batch on the adequacy of mixing should be considered when changing from recommended operating volumes are planned.

Dry powder mixers should be selected or designed for a particular operation, firstly by analyzing and giving careful examinations to the applicable areas discussed above. Mixer selection should also involve consideration of the place of the mixer within the overall process. Possible integration of many steps within the overall processing of solids in a particular application deserves scrutiny at this time. If there is no machinery available including all the necessary requirements, consideration should be given to modifications in order to obtain the most desirable features combination of the equipment.

Pilot tests are relevant in the final decision of selection or design of any specific mixing process. In general larger pilot plant units would provide more reliable data for performance prediction of large-scale powder mixing plants. With geometrically similar tumblers, if the speeds are adjusted to

give comparable motion and the mixer volume fraction occupied by the load is the same, scale-up of results will be straightforward. The presence of internal rotating devices leads to difficulties in scaling-up clearances, blade area to mixture volume, and sizes as well as speeds of the rotating devices. The actual materials to be processes in the industrial operation should be used if possible in the scale-up procedure. If substitute materials need to be used, they should have the same mixing characteristics. Differences in materials of construction between the pilot and the production unit should be considered, since these may have a bearing on caking, abrasion, and some other adverse effects.

6.6 Laboratory Exercise: Determination of Blending Indices for Different Tumbling Mixers

6.6.1 Introduction

Solid-solid mixing processes normally start by pouring known quantities or two or more solids into a mixing unit. The overall proportion of any component is known by the relation between the mass of that component and the total mass of all components (i.e., the solids fraction). If two components are mixed in equal proportion, it would be expected that uniformity of mixing can be verified by sampling, and when all extracted samples present proportions of 0.5 for any of the components, a perfect mixing had been achieved. It is unrealistic, however, to expect all the samples to have an exact proportion of one component: some samples will have more, some less, but the overall mean average of one component as proportion, would tend to equal the original proportion due to the solids fractions combined initially. Most of the proposed models to determined mixing indices as an indicative of blending homogeneity are based on experimental determination of sample's mean values and variances. The objective of the experiment is to determine mixing indices for a solids mixing operation using two types of tumbling equipment.

6.6.2 Equipment and Materials

Two tumbling mixers (e.g., double cone mixer and V-cone mixer). Two batches of differently colored ballotini beads with a size range 0.8–1.2 mm each and the same density.

6.6.3 Instruments or Apparatuses

Set of standard sieves and sieve shaker as well as analytical balance.

6.6.4 Operation of the Mixers

- Standardize the size of each solid sample by using a 1 mm sieve, as to separate the fine fraction of Sample A and the coarse fraction of Sample B.
- Adjust the speed of rotation of one of the mixers.
- Weight the same quantity of each standardized solid, for example, 500 g, and pour them into the mixer.
- Start the mixing operation at constant speed and record the time (t = 0 min).
- Stop the mixer after two minutes (t = 2 min).
- Collect five 30 g spot samples from different parts of the mixer.
- Measure and record the weights.
- Sieve them using the 1 mm screen and collect the fine and coarse fractions of the sample.
- Record the weights of the fractions separated in the previous step.
- Make an estimate of the number of particles in the sample, either by counting (using the bear eye or a microscope) or relating properties like minimum weight detected with particles able to be counted.
- Continue the mixing operation for 2 more minutes and stop at this third time (t = 4 min). Repeat the procedure for the t = 2 min sample.
- Continue, progressively, taking samples and repeating procedures up to complete 10 min mixing.
- Repeat the experiment for the second mixer.

6.6.5 Calculations and Report

- The recorded weight of the fractions separated from the spot samples will give an estimate of the proportion of one selected component. By determining these proportions, elaborate a table such as Tables 6.1 or 6.2.
- With the estimate of the number of particles in the spot samples, make a development like the one given in the solution of Examples 6.1 and 6.2. Calculate values of *k* using Equation 6.8.
- Elaborate plots like the one shown in Figure 6.1 for both tumbling methods.
- Recalculate an ideal time for the two tested processes, to reach a value of $1/M_2$ of 0.95.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature.

1 Toblem 2											
Run	Time (s)	Flavor Aid Fraction									
1	75	0.95	0.25	1.04	0.15	0.88	0.92	0.18	0.35	0.20	0.48
2	100	0.58	0.81	0.84	0.38	0.78	0.46	0.50	0.60	0.27	0.68
3	125	0.67	0.50	0.59	0.45	0.72	0.75	0.57	0.66	0.29	0.70
4	250	0.61	0.58	0.63	0.62	0.54	0.59	0.68	0.62	0.60	0.63
5	500	0.62	0.61	0.60	0.59	0.61	0.62	0.60	0.60	0.58	0.60

TABLE 6.3 Problem 2

Include graphs, charts, and calculations carried out. Include analysis from own data and, as much as possible, discussion based on literature survey from studies and investigations on the subject.

PROBLEMS

- 1. A granular product from the ceramics industry is mixed with 12% (by weight) of a light colorant. After 3 min mixing, 12 spot samples are taken randomly and analyzed for color uniformity. The instrument relates color concentration with its proportion in each sample, giving the following results for colorant concentration in the 12 samples taken: 11.28, 10.88, 9.94, 11.34, 12.18, 11.23, 12.91, 10.92, 10.22, 11.96, 11.87, and 11.75. Calculate the mixing index.
- 2. A screw mixer is used to incorporate a flavor aid to a jelly. The product is packed in 500 g packets and the allowed range for the flavor aid is 0.57% to 0.63% per packet. Ten samples with an approximate of 6000 particles each are taken at the different times given in Table 6.3. Determine if sampling demonstrates that the mixing requirement is accomplished at the maximum 500 s treatment, and calculate mixing indices for each run. Make comments on the efficiency of the mixing process, based on the obtained results.

References

Barbosa-Cánovas, G. V., Ortega-Rivas, E., Juliano, P., and Yan, H. 2005. Food Powders: Physical Properties, Processing, and Functionality. New York, NY: Kluwer Academic/Plenum Publishers.

Boss, J. 1986. Evaluation of the homogeneity degree of a mixture. *Bulk Solids Handling* 6: 1207–1210.

Dankwertz, P. V. 1952. The definition and measurement of some characteristics of mixing. *Applied Science Research* 3A: 279–281.

Fan, L. T. and Wang, R. H. 1975. On mixing indices. Powder Technology 11: 27–32.

Mixing 247

Harnby, N. 1985. The selection of powder mixers. In *Mixing in the Process Industries*, Harnby, N., Edwards, M. F., and Nienow, A. W. (eds.). London: Butterworths.

- Lacey, P. M. C. 1954. Developments on the theory of particle mixing. *Journal of Applied Chemistry* 4: 257–268.
- McCabe, W. L., Smith, J. C., and Harriot, P. 2005. *Unit Operations in Chemical Engineering*, 7th Ed. New York, NY: McGraw-Hill.
- Niranjan, K. 1995. An appraisal of the characteristics of food mixing. In *Food Process Design and Evaluation*, Singh, R. K. (ed.). Lancaster, PA, USA: Technomics.
- Niranjan, K. and de Alwis, A. A. 1993. Agitation and agitator design. In: *Encyclopedia of Food Science, Food Technology and Nutrition*, Vol. 1, Macrae, R., Robinson, R. K. and Sadler, M. (eds.). London: Academic Press.
- Poole, K. R., Taylor, R. F., and Wall, G. P. 1964. Mixing powders to fine scale homogeneity: studies of batch mixing. *Transactions of the Institution of Chemical Engineers* 42: T305–T315.
- Stange, K. 1963. Die mischgute einer Zufallsmischung aus drei und mehr Komponenten. Chemie Ingenieur Techik 35: 580–582.
- Van Deemter, J. J. 1985. Mixing. In *Fluidization*, 2nd Ed. Davidson, J. F., Clift, R. and Harrison, D. (eds.). London: Academic Press.
- Williams, J. C. 1968/1969. The mixing of dry powders. *Powder Technology* 2: 13–20.

This page intentionally left blank



7.1 Theoretical Fundamentals

Particulate solids, when packed within a container, form an arrangement allowing for the void space between particles to be filled by a gas, normally air. Such arrangement has generally been referred to as a bed, and the interactions between the solids particles and the voids filled with air are related by an important property known as porosity. Any particular fluid flowing at low velocities through a porous bed of solid particles, for example, in a packed tower, does not cause the particles to move. The fluid passes through the small, tortuous channels, losing pressure energy. The pressure drop in a stationary bed of solids can be described by fundamental fluid mechanics theory. However, if the fluid velocity is steadily increased a point is eventually reached at which the particles no longer remain stationary but are fluidized under the influence of the fluid. Fluidization is a relevant property of particulate solids since, when fluidized, particulate solids within a fluid stream could behave somewhat as fluids and, therefore, they may be studied under the scope of fluid mechanics. Some fluid-like properties that fluidized bed present are, for example:

- The surface of the bed becomes horizontal and, if disturbed, returns quickly to horizontal.
- Like in a liquid, the pressure increases with depth.
- Objects of lower density float on the surface of the fluidized bed, whereas objects of higher density sink to the bottom of the fluidized bed.
- If a hole is opened on the wall of the containing vessel of the fluidized bed, particles will pour out as a jet, similar to water from a puncture in a water container.
- Bubbles can rise through the fluidized bed and burst in the surface as in a liquid. Moreover, the bubble shapes and rise velocities are similar to those of bubbles in liquids.

The state of being fluidized, called "fluidization" as already done, turns out to be advantageous in numerous operations of the materials processing industry run, normally, by practicing chemical engineers.

7.1.1 Bulk Density and Porosity of Beds

When a powder just fills a vessel of known volume V and the mass of the powder is m then the bulk density of the powder is m/V. However, if the vessel is tapped in most cases the powder will settle leaving top empty space, so adding more powder would be necessary to end with the vessel completely filled. If the mass now filling the vessel is m' then the bulk density is m'/V > m/V. Clearly, this change in density just described has been caused by the influence of the fraction of volume not occupied by a particle, known as porosity. The bulk density is, therefore, the mass of particles that occupies a unit volume of a bed, while porosity or voidage is defined as the volume of the voids within the bed divided by the total volume of the bed. These two properties are in fact related via the particle density in that, for a unit volume of the bulk powder, there must be the following mass balance:

$$\rho_{\rm b} = \rho_{\rm s}(1 - \varepsilon) + \rho_{\rm a}\varepsilon \tag{7.1}$$

where ρ_b is the powder bulk density, ρ_s is the particle density, ϵ is the porosity, and ρ_a is the air density. As the air density is small relative to the powder density, it can be neglected and the porosity can thus be calculated simply as

$$\varepsilon = \frac{(\rho_{\rm s} - \rho_{\rm b})}{\rho_{\rm s}} \tag{7.2}$$

Equation 7.2 gives the porosity or voidage of the powder and whether or not this includes the pores within the particles depends on the definition of particle density used in such evaluation.

7.1.2 Fluid Flow through Solids Beds

Fluidization can be considered a special case of fluid-solids flow with a rather complex behavior (Jackson, 2000). The fluid flow resistance through the voids in a bed of solids is the resultant of the total drag of all the particles in the bed (McCabe et al., 2005). Depending on the particle's Reynolds number, laminar flow, turbulent flow, form drag, separation, and wake formation may occur. Similar to the drag of a single solid particle, there is no sharp transition between laminar and turbulent flow like that occurring in flow through conduits of constant cross section. Calculations of the total pressure drop through a bed of solids can be based on estimates of the total drag of the fluid on the solid boundaries of the tortuous channels through the bed of

particles (Zenz and Othmer, 1960). This can be done by assuming that the actual channels may be effectively replaced by a set of identical parallel conduits, each of variable cross section, and that the mean hydraulic radius of the channels is appropriate to account for the variations in channel cross-sectional size and shape. The total drag per unit area of channel wall would be the sum of viscous drag forces and inertial forces, while a direct determination of the total surface area of the bed would be the sum of the individual surface area of all particles within the bed. Since this latter approach would be unpractical and unattainable, total surface area could be derived from estimation of the relation surface volume for the solids in the bed. The volume fraction of solids particles in a bed can be defined as $1-\varepsilon$, being ε the porosity previously defined by Equation 7.2. The bed porosity can easily evaluated *in situ* by filling the bed with a liquid, draining, and comparing the volume of the liquid to the total volume of the bed as calculated from its depth and cross section.

The volumetric average velocity of the fluid through the channels within the bed u and the velocity of the fluid stream just before entering to the first layer of solids u_0 are related by the porosity. The usual configuration of a packed bed is a large cylindrical vessel with a supporting grid some distance above the bottom of the tower. The velocity u_0 is known as the superficial velocity or empty-tower velocity, and u is the upward or downward velocity in the open section below the grid or in the empty tower above the bed. The average free cross section of all the channels has the same ratio to the total cross section of the empty tower, while the volume of the channels has the same ratio to the total volume of the bed of solids. The above-mentioned relation between u and u_0 can be represented by

$$u = \frac{u_0}{\varepsilon} \tag{7.3}$$

7.1.3 Mechanism of Fluidization: Aggregative and Particulate

The effect of flowing of a fluid through a bed of particles may cause three effects as a function of the fluid velocity: the bed may expand a little but remain fixed at lower fluid velocities, the particles may become freely supported in the fluid al higher velocities, and the particles may become totally suspended by the fluid stream and may be transported within it as in pneumatic transport. If the fluid flow rate is increased above the minimum required to produce a fluidized bed, this will continue to expand so that the average distance between particles will become greater, or the fluid excess will pass through the bed in the form of rising pockets of gas giving origin essentially to a two-phase system. From the earliest applications of fluidized beds it was noticed that liquid-fluidized beds presented a smooth appearance expending progressively as the fluid flow rate increased, while most

gas-fluidized beds were typically heterogeneous containing the rising pockets of gas previously mentioned practically free from particles. These gas pockets were referred to as "bubbles" and the terms "particulate" and "aggregative" were introduced to describe the two types of fluidized behavior. Generally, particulate fluidization occurs with solid–liquid systems and with solid–gas systems when the particles are very fine and only over a limited range of velocities. Aggregative fluidization occurs with all other solid–gas systems and, sometimes, with solid–liquid systems for high-density solids. The growth rate of a small bubble will generally be greater in solid–gas systems that in solid–liquid systems.

The type of fluidization occurring in a process will be related to the maximum stable size of bubble that can exist within the fluidized bed. If the circulation velocity of gas within the bubble exceeds the terminal velocity of particles, these will be sucked in at the rear of the bubble, which will thus tend to be destroyed. Since the bubble rise velocity will increase with bubble volume, being substantially independent of the properties of the fluidized bed, the maximum stable bubble size will increase with the terminal falling velocity of the particles in the fluid. If this size exceeds by a large amount compared to the particle diameter, the bubble will appear obvious and the fluidization will be aggregative. On the other hand, when the bubble size is of the same order of the particle diameter, such bubble can no longer be distinguished from the interstitial gas and the fluidization will appear to be particulate. On this basis, most solid—gas systems will give rise to aggregative fluidization whereas most solid—liquid systems will originate particulate fluidization.

An empirical criterion to distinguish the type of fluidization governing a process was proposed quite early by Wilhelm and Kwauk (1948). They suggested the use of the Froude number, that is $(Fr = u_{mf}^2/xg)$ as the criterion for the type of fluidization observed. In general, aggregative fluidization occurs at values above unit and particulate fluidization at values below unity. In the Froude number u_{mf} denotes the minimum fluidization velocity and x a particle equivalent diameter. Within the context of qualitative distinctions between particulate and aggregative fluidization, research efforts have been focused for a while into three avenues. One has consisted of lengthy mathematical analyses in which the bed has been treated as a homogeneous mass regardless of its physical details. A second one has sought empirical correlations of associated heat and mass transfer, as well as bed expansion and other properties necessary in equipment design. The third avenue has directed research in studying the lack homogeneities occurring in the more widely applicable aggregatively or gas-fluidized beds, that is, investigating fundamentally the bubble formation process from experimental and hydrodynamic viewpoints. The first of these directions was to distinguish between gas- and liquid-fluidized beds; the second provided necessary data sufficient for first-generation plant designs; and the third produced basic data on bubble shape, velocity of rise, effect on particle mixing, and related observations.

7.2 Fluidized Regimes

An understanding of the fluidized regimes within the bed should include criteria to distinguish the limits from the fixed to the fluidized, and to the suspended (transport) state of particles in a stream of fluid. The relation between the superficial velocity u of the fluid, calculated from the whole cross section of the containing vessel, and the pressure gradient is shown in Figure 7.1. If the flow is streamline a straight line of slope unity is obtained at low velocities. At the fluidizing point the pressure gradient begins to fall because the porosity of the bed increases, and the fall continues until the velocity is sufficient for transport of the solids to take place. The pressure gradient then starts to increase again because the frictional drag of the fluid at the walls of the tube starts to become significant. A minimum fluidizing velocity will mark a transition from the fixed to the fluidized state. At the minimum fluidizing velocity it may be expected that flow relations for the fixed bed would be applicable to the fluidized bed. At the opposite limit when a bed is fully expanded so that its voidage approaches unity, it consists effectively of single isolated particles suspended in the fluid, and thus any relationship describing a fluidized bed should extrapolate at unit voidage to one applicable to an individual particle. At intermediate conditions, the particulate fluidized bed is hydrodynamically similar to a sedimenting suspension.

In a fluidized bed the particles undergo no net movement as they are supported by the upward fluid flow, whereas in a sedimenting suspension the particles are moving continuously downward and the only net movement of

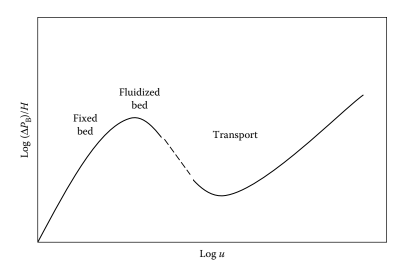


FIGURE 7.1 Pressure gradient in bed as a function of fluid velocity.

fluid is that due to the displacement of fluid by the particles being settled. This suggests that similar velocity–voidage relationships might exist for sedimenting suspensions and particulate-fluidized beds.

In fluidized beds there are favorable conditions for rapid heat and mass transfer between the solids and the fluid. Very rapid mixing of the solids generally occurs so that the coefficients for the transfer of heat to boundary surfaces are quite high. Fluidized beds are, therefore, used both as heat exchangers and chemical reactors particularly where close control of temperature is required and where large amounts of heat must be added to, or removed from the system. There are, of course, many other relevant applications of fluidized beds in the materials processing industry. A proper knowledge of the flow patterns of both fluid and particles is, therefore, necessary.

7.2.1 Pressure Drop-Velocity Relationship

When a fluid passes vertically through a bed of particles the pressure drop $\Delta P_{\rm B}$ will initially increase as the velocity u is increased, as shown in Figure 7.2. The relation between pressure drop and velocity will be that applicable to a fixed bed. For fine particles, the increasing straight line in Figure 7.2 has been properly described by a relationship known as the Carman–Kozeny equation:

$$u = \frac{\varepsilon^3}{5(1-\varepsilon)^2} \frac{\Delta P_{\rm B}}{S^2 \mu H}$$
 (7.4)

where S is the specific surface of the solid, μ the fluid viscosity, and H the bed depth.

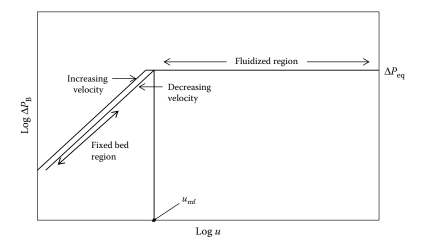


FIGURE 7.2 Ideal pressure drop–velocity curve.

As the velocity reaches a point in which the frictional pressure drop equals the buoyant weight per unit area of particles (ΔP_{eq}), any further increase in velocity must result in a slight upward movement of the particles forming the bed. The particles will rearrange so that the resistance to fluid flow will decrease and, in general, the voidage of the bed will increase. In an idealized system, the pressure drop ΔP_{eq} will remain constant, as represented by the plateau in Figure 7.2. Further increases in velocity will result in continuing expansion of the bed until, eventually, any further separation of the particles will only be possible if they become physically separated from one another and become free to move within the fluid. At this stage the bed is just fluidized, is said to be at the "point of incipient fluidization," and the superficial fluid velocity there is known as the "minimum fluidizing velocity" u_{mf} . If the fluid velocity is further increased the pressure drop over the bed remains constant, but if the fluid velocity is progressively reduced the pressure remains constant at the point of incipient fluidization. Further reduction of velocity is accompanied by a progressive decrease in pressure drop and the pressure drop-velocity curve will generally be lower than the one obtained while the velocity was being increased, since in the absence of vibration the bed voidage will remain approximately at the value ε_{mf} corresponding to a bed at the point of incipient fluidization.

The behavior described above is an idealized one and the deviation from it, gives a good indication of the factors influencing the performance of the bed. Deviations from ideal behavior can be due to a number of reasons. For example, as the velocity approaches the minimum fluidizing velocity some normal expansion will occur before the pressure drop has reached the buoyant weight per unit area of bed. This effect would be most marked when the bed is initially highly consolidated. In addition, local variations in the permeability of a randomly packed bed would tend to cause a gradual transition between the two sections of the curve. Also, the tendency of the particles to interlock with one another may cause partial bridging, particularly in smalldiameter beds, with the results that frictional forces are exerted by the walls of the bed container. It is, therefore, possible for pressure drops in excess of the theoretical value to be obtained, so that the curve will pass through a point of maximum pressure drop, exhibiting a small characteristic "hump," as shown in Figure 7.3. Furthermore, nonuniformity in the structure of the bed will result in preferential flow of fluid in particular areas, resulting in possible coexistence of fixed and fluidized regions within the bed. The bed may appear to be well fluidized, but part of its weight may still be borne by the bed distributor, resulting in a pressure drop below its expected value. The minimum velocity at which the bed weight is fully supported by the fluid appears as point u_{fs} in Figure 7.4.

The form of the curve of pressure drop against velocity for the fixed and fluidized beds should provide considerable amount of information on the structure of the bed. Deviations from the idealized behavior are attributable to intraparticle forces and maldistribution of the fluid in the bed. For practical

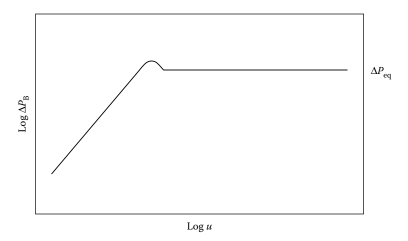


FIGURE 7.3Pressure drop over fixed and fluidized beds as a function of fluid velocity.

purposes, the minimum fluidized velocity is defined as shown in Figure 7.4 as the point of intersection of the pressure drop lines for the fixed and fluidized regimes.

7.2.2 Incipient Fluidization and Minimum Fluidizing Velocity

Due to the complexity of interaction of fluid-solid systems is not possible to predict precisely the behavior of fluidized-bed operations. With solid-gas

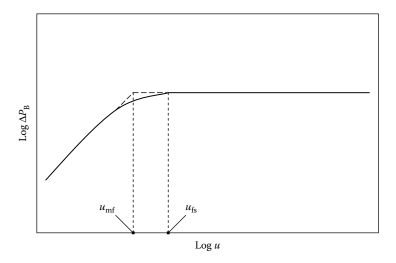


FIGURE 7.4 Effect of presence of nonfluidized regions in bed.

systems there is a very marked contrast in the behavior of solid materials: some of them will fluidize extremely well, while some others will be virtually incapable of being fluidized. The following properties of the solid and the fluid will be determinant in achieving well-fluidized systems: (a) low particle density, (b) small particle size, (c) small particle size range, (d) particle shape approaching spherical, and (e) high fluid density. The properties of the solids identified as the more influential in making it capable of giving a well-fluidized system are small particle size and low density. Unfortunately, those same properties have been observed to make the initiation of fluidization difficult. If surface forces between the particles are significant, they will be greater with small particles because of their high specific surface. If the particles possess low density, the gravitational forces tending to pull them apart will be small. Thus, small low-density particles can cause serious channeling even though they may fluidize pretty well once they have been brought into suspension.

In an idealized system the minimum fluidizing velocity is that at which the bed suddenly changes from a fixed to a fluidized state. In practice, however, there may be a large transitional region so that the minimum fluidizing velocity will be difficult to determine. The problem of definition generally becomes greater for particles of a wide size range. The minimum fluidized velocity can be determined graphically or can be calculated analytically. In the graphical method, that determination can be conveniently done by using the plot of frictional pressure drop against velocity. If separate lines are drawn through the points for the fixed and fluidized regions, and the points in the intermediate region are ignored, the point of intersection of these two lines will give a reproducible value for the minimum fluidized velocity, just as shown in Figure 7.4.

An approximate analytical calculation of the minimum fluidizing velocity can be carried out by using an expression for the relation between pressure drop and superficial velocity for a fixed bed and putting the pressure drop equal to the buoyant weight of the particles. The pressure drop through the bed is then equal to its apparent weight per unit area, and the porosity at the onset of fluidization is the maximum that the fixed bed can attain. Previous knowledge of the voidage of the bed at the minimum fluidizing velocity (ϵ_{mf}) would be necessary, but this would depend on the shape and size range of the particles. A value of about 0.4 would be appropriate for approximately spherical particles. Taking into account the above considerations, an expression for the pressure drop over the fluidized bed at incipient fluidization will be given by

$$\Delta P_{\rm B} = \frac{mg}{A} = (\rho_{\rm s} - \rho_{\rm f})(1 - \varepsilon)Hg \tag{7.5}$$

where m is the mass of solids in the bed, g is the acceleration due to gravity, A is the cross-sectional area of the bed, ρ_s is the solids density, and ρ_f is the fluid density.

If the bed expands the product $(1 - \varepsilon)$ will remain constant. Using the values of each term proper for the conditions of incipient fluidization:

$$\Delta P_{\rm B} = (\rho_{\rm s} - \rho_{\rm f})(1 - \varepsilon_{\rm mf})H_{\rm mf}g \tag{7.6}$$

For fine particles the pressure drop-velocity relationship will be given by the Carman–Kozeny equation, which will take the following form for incipient fluidization:

$$u_{\rm mf} = \frac{\varepsilon_{\rm mf}^3}{5(1 - \varepsilon_{\rm mf})^2} \frac{\Delta P_{\rm B}}{S^2 \mu H_{\rm mf}} \tag{7.7}$$

Substituting Equation 7.6 into Equation 7.7, the following expression results:

$$u_{\rm mf} = \frac{\varepsilon_{\rm mf}^3}{5(1 - \varepsilon_{\rm mf})} \frac{(\rho_{\rm s} - \rho_{\rm f})g}{S^2 \mathfrak{u}}$$
 (7.8)

For uniform spherical particles S = 6/x, where x is an equivalent particle diameter. Also, taking $\varepsilon_{\rm mf} = 0.4$, Equation 7.8 would transform into

$$u_{\rm mf} = 0.00059 \frac{x^2 (\rho_{\rm s} - \rho_{\rm f})g}{\mu}$$
 (7.9)

For larger particles, a more general equation such as the Blake, Carman, or Ergun (1952) equation must be used for the pressure drop through the fixed bed. Using the Ergun equation:

$$\frac{\Delta P_{\rm B}}{H_{\rm mf}} = 150 \frac{(1 - \varepsilon_{\rm mf})^2}{\varepsilon_{\rm mf}^3} \frac{\mu u_{\rm mf}}{x^2} + 1.75 \frac{(1 - \varepsilon_{\rm mf})}{\varepsilon_{\rm mf}^3} \frac{\rho_{\rm f} u_{\rm mf}^2}{x}$$
(7.10)

Substituting for $\Delta P_{\rm B}/H_{\rm mf}$ from Equation 7.6 and multiplying both sides by $\rho_{\rm f} x^3/\mu^2 (1 - \varepsilon_{\rm mf})$, the following expression will result:

$$\frac{\rho_{\rm f}(\rho_{\rm s} - \rho_{\rm f})gx^3}{\mu^2} = 150 \frac{1 - \varepsilon_{\rm mf}}{\varepsilon_{\rm mf}^3} \frac{u_{\rm mf}x\rho_{\rm f}}{\mu} + \frac{1.75}{\varepsilon_{\rm mf}^3} \left(\frac{u_{\rm mf}x\rho_{\rm f}}{\mu}\right)^2 \tag{7.11}$$

Identifying the following groups:

$$\frac{\rho_f(\rho_s - \rho_f)gx^3}{\mu^2} = Ga \quad \text{and} \quad \frac{u_{mf}x\rho_f}{\mu} = Re_{mf}$$

where Ga is the Galileo number and Re_{mf} is the Reynolds number at incipient fluidization point.

Considering the above-defined dimensionless numbers, Equation 7.11 would modify to

$$Ga = 150 \frac{1 - \varepsilon_{mf}}{\varepsilon_{mf}^{3}} Re_{mf} + 1.75 \frac{1}{\varepsilon_{mf}^{3}} Re_{mf}^{2}$$
 (7.12)

Taking $\varepsilon_{\rm mf}$ = 0.4 and transposing

$$Re_{mf}^2 + 51.4 Re_{mf} - 0.0366 Ga = 0 (7.13)$$

Thus,

$$Re_{mf} = 25.7 \left[\sqrt{1 + 5.53 \times 10^{-5} Ga} - 1 \right]$$
 (7.14)

Equation 7.14 can be used for nonspherical particles if the diameter of the sphere with the same specific surface as the particles is used. If the pressure drop through the bed is a significant proportion of the total pressure and the fluid is compressible, the velocity of the fluid will increase as it passes through the bed. The top of the bed will thus tend to fluidize at a lower flow rate of fluid than the bottom.

The minimum fluidizing velocity $u_{\rm mf}$ can be expressed in terms of the free falling velocity (terminal velocity) $u_{\rm t}$ of the particles in the fluid. The Ergun equation relates the Galileo number Ga to the Reynolds number at the incipient fluidization point $\epsilon_{\rm mf}$. The Reynolds number involving the terminal velocity of the particle in the fluid (Re_t = $u_{\rm t} x \rho_{\rm f}/\mu$), or particle Reynolds number, can also be expressed in terms of the Galileo number but no single expression covers the full range of Reynolds numbers and so it is most satisfactory to cover each of the three zones separately, as to give the following three expressions of Ga as a function of Re_t:

$$Ga = 18 Re_{t} (Ga < 3.6)$$
 (7.15)

$$Ga = 18 Re_{t} + 2.7 Re_{t}^{1.687} (3.6 < Ga < 10^{5})$$
(7.16)

$$Ga = \frac{1}{3} Re_t^2 (Ga > 10^5)$$
 (7.17)

It is, thus, possible for any value of ε_{mf} to calculate the relation Re_t/Re_{mf} as a function of Ga, as shown in Figure 7.5.

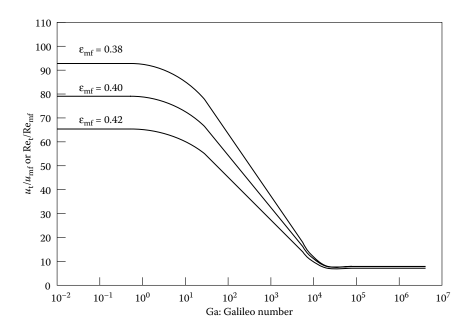


FIGURE 7.5Ratio of terminal falling to minimum fluidization velocity as a function of Galileo number.

Example 7.1

A bed of fractioned pulverized coal 65-mesh (Tyler) is fluidized by air at room-temperature conditions. The height of the static bed is 1.80 m. If the coal particles may be considered spherical, while their density is 1350 kg/m³, calculate the pressure drop required for fluidization.

SOLUTION

Consulting a Tyler Standard Screen Scale Table, it can be read the value of the screen equivalent diameter as 0.208 mm, so the pulverized coal may be considered a fine powder. Since the particles are taken as spherical, the ideal porosity ϵ = 0.4 can be used for calculation. Finally, from any reference book (or appendix of some engineering textbook) listing physical properties of materials, at approximately 25°C the density of air is of the order of 1.2 kg/m³. Substituting directly into Equation 7.5:

$$\Delta P_{\rm B} = (1350 - 1.2)(1 - 0.4)(1.8)(9.81) = 14,290.26 \text{ N/m}^2 \approx 14.3 \text{ kPa}$$

Example 7.2

A bed of 100 kg dried peas is to be fluidized in chilled air at -5° C as a part of a process. The peas are 5 mm in size and their density approaches 1000 kg/m³.

Calculate the air velocity, the pressure drop through the bed if the cross-sectional area is 0.3 m², and the bed height at a voidage 20% above the minimum.

SOLUTION

The peas can be considered spherical and homogeneous in size. The ideal voidage $\epsilon=0.4$, can therefore, be used. Consulting any reference book (or appendix of some engineering textbook) listing physical properties of materials, at approximately -5° C the density of air is in the order of 1.3 kg/m³ while its viscosity is around 1.6×10^{-5} Pa s. The particulates are, obviously, large and so, the minimum fluidization velocity should be transposed from the expression of Re_{fm} in Equation 7.14:

$$u_{mf} = \frac{25.7(\mu_g) \left[\sqrt{1 + 5.53 \times 10^{-5} Ga} - 1 \right]}{x \rho_f}$$
$$= \frac{25.7(1.6 \times 10^{-5} \left[\sqrt{1 + 5.53 \times 10^{-5} Ga} - 1 \right]}{0.005(1.3)}$$

 $u_{\rm mf} = 0.9692 \text{ m/s} \approx 1 \text{ m/s}.$

The pressure drop through the bed can be estimated substituting directly into the first right-hand term of Equation 7.5, that is;

$$\Delta P_{\rm B} = \frac{(100)(9.81)}{0.3} = 3270 \,\text{Pa}$$

The porosity at 20% above the minimum would be 0.48 and, thus, transposing for the height $H_{\rm mf}$ in Equation 7.6:

$$H_{\rm mf} = \frac{3270}{(1 - 0.48)(1000 - 1.3)(9.81)} = 0.64186 \,\mathrm{m} \approx 0.64 \,\mathrm{m}$$

7.2.3 Heterogeneous Fluidization: Bubbling

Bubbles or particle-free voids may appear in a fluidized bed once the minimum fluidization velocity has been surpassed. Bubbles in a gas-fluidized bed migrate upward, just like bubbles in a boiling liquid. Figure 7.6 shows the typical pattern followed by bubbles in many cases of fine particulate solids fluidized by a gas. At superficial velocities above $u_{\rm mfr}$, fluidization may be either bubbling or nonbubbling. Some combinations of fluid and particles give rise to "only bubbling fluidization" while some other combinations give origin to "only non-bubbling fluidization." Gas-fluidized systems will present only bubbling fluidization or nonbubbling fluidization starting at $u_{\rm mfr}$ followed by bubble fluidization as fluidizing velocity increases. Nonbubbling fluidization is also known as particulate or homogeneous fluidization,

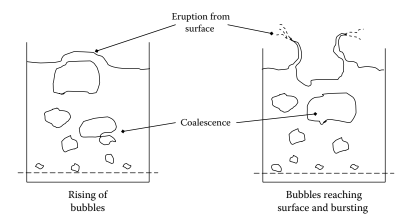


FIGURE 7.6Behavior of bubbles for a typical fluidized Group B powder (Geldart's classification).

whereas bubbling fluidization is often referred to as aggregative or heterogeneous fluidization.

The fluidization properties of a powder in air may be predicted by establishing in which group of the Geldart's classification (Chapter 2, Section 2.2.2) such powder lies. Since the range of gas velocities over which nonbubbling fluidization occurs in Group A powders is small, bubbling fluidization is the most commonly observed type in gas fluidized systems commercially in use. The superficial gas velocity at which bubbles first appear in a fluidized bed is known as the minimum bubbling velocity $u_{\rm mb}$. Abrahamsen and Geldart (1980) correlated the maximum values of $u_{\rm mb}$ with gas and particle properties using the following equation:

$$u_{\rm mb} = 2.07(e)^{0.716F} \left(\frac{\chi \rho_{\rm g}^{0.06}}{\mu^{0.347}} \right)$$
 (7.18)

where F is the fraction of powder less than 45 μ m.

In most solid–gas systems bubbling occurs when the gas velocity is increased well over the minimum fluidizing velocity. The simplest description of the expansion of a gas-fluidized bed comes from the two-phase theory of fluidization (Rhodes, 1998), originally attributed to Toomey and Johnstone (1952). This theory suggests that the bubbling fluidized bed is composed of two phases: the bubbling phase and the particulate phase (also known as the emulsion phase), and that all the gas in excess of that required to fluidize the system will pass through the bed in the form of bubbles.

The lower and upper limits of gas velocity at which a uniform particulate phase can exist are the minimum fluidizing velocity and the minimum bubbling velocity, respectively. It is quite difficult to define the latter precisely, since it will depend critically on the previous history of the bed, on the nature

of the gas distributor, and on the presence of obstructions within the bed. The ratio $u_{\rm mb}/u_{\rm mf}$ does give a measure of the degree to which the bed can be expanded, tending to be relatively high for fine low-density solids and high-density gas. There is a minimum stable size of bubbles that can exist within the particulate phase. Smaller bubbles tend to shrink and become assimilated in the dense phase, while large bubbles tend to grow at the expense of the gas in the continuous phase. If a bubble larger than the critical size is injected into an expanded bed, this will initially expand by an amount equal to the volume of injected gas. However, when the bubble breaks the surface, the bed level will fall back to a value below that before injection was effected. The transference of gas between dense and bubble phases within a fluidized bed is an important factor, which must be considered in designing fluidized beds as reactors.

Powders of small particle size and low density are difficult to fluidize initially, but once they reach the fluidized state they can give origin to beds that can be expanded over a wide range before bubbling occurs. The formation of large bubbles in gas-fluidized systems is often prevented by introducing standard tower packings into the bed to produce what is known as a packed-fluidized bed.

The bed expansion can be expressed in terms of the fraction occupied by bubbles ε_{B} :

$$\varepsilon_{\rm B} = \frac{H - H_{\rm mf}}{H} = \frac{Q - Q_{\rm mf}}{Au_{\rm B}} = \frac{u - u_{\rm mf}}{u_{\rm B}}$$
(7.19)

where H is the bed depth at u and $H_{\rm mf}$ is the bed depth at $u_{\rm mf}$; Q is the actual gas flow rate to the fluid bed and $Q_{\rm mf}$ is the gas flow rate at incipient fluidization; A is the bed section area; $u_{\rm B}$ is the mean rise velocity of a bubble in the bed.

The voidage of the emulsion phase is taken to be that at minimum fluidization ϵ_{mf} . The mean bed voidage is then given by

$$(1 - \varepsilon) = (1 - \varepsilon_{\rm B})(1 - \varepsilon_{\rm mf}) \tag{7.20}$$

In practice, however, the two-phase flow theory overestimates the volume of gas passing through the bed as bubbles, that is, the visible bubble flow rate (Rhodes, 1998) and better estimates of bed expansion may be obtained by replacing ($Q-Q_{\rm mf}$) in Equation 7.19 with the following relationship for the bubble flow rate:

$$Q_{\rm B} = YA(u - u_{\rm mf}) \tag{7.21}$$

where: 0.8 < Y < 1.0 for Group A powders (Geldart's classification) 0.6 < Y < 0.8 for Group B powders (Geldart's classification) 0.2 < Y < 0.6 for Group D powders (Geldart's classification)

Strictly, the above equations should be written in terms of $u_{\rm mb}$ rather than $u_{\rm mf}$, and $Q_{\rm mb}$ rather than $Q_{\rm mf}$, so that they are valid for both Group A and Group B powders. Practically, however, it makes little difference, since both $u_{\rm mb}$ and $u_{\rm mf}$ are usually much smaller than the superficial fluidizing velocity u and, therefore: $(u-u_{\rm mf})\approx (u-u_{\rm mb})$.

7.2.4 Spouted Beds

The spouted bed is a variant of the fluidized bed in which, instead of distributing the upward-moving gas evenly across the solids bed by a gas distributor, the gas enters as a jet at the center of the conical base of the chamber. Figure 7.7 shows a schematic diagram of a spouted bed. The solid particles are dispersed into the jet of gas, the spout, and the entrance of the fluidizing gas. In operation, a systematic cyclic movement of the solids is established: the particles travel upward in the axial jet zone and downward in the

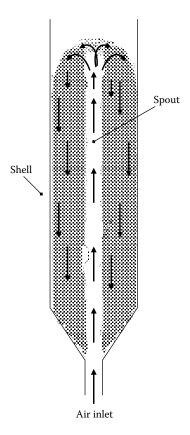


FIGURE 7.7 Schematic diagram of a spouted bed.

annular dense moving bed. The spouted-bed technique permits satisfactory agitation of solid materials that are too coarse and uniform in size for good fluidization. Spouted beds were originally designed for drying of granular materials, but they have proved to be of interest for a variety of processes including blending of polymer chips, coating of tablets, granulation of fertilizers, carbonization of coal, and pyrolysis of shale.

Spouted beds feature special advantages compared with normal fluidized beds when the particles, or rather particulates, to be fluidized cover sizes over the few millimeters range. If such is the case, in conventional fluidized beds the fluidization may be sluggish and solids mixing, as well as gasparticle effective contact, would be difficult.

Unlike fluidization spouting displays a pressure gradient dP/dZ (being Z the vertical distance from gas inlet orifice), which is not uniform along the bed height; it is small near the base and increase to a maximum value at the bed surface. The pressure drop arises out of two parallel resistances: that of the spout in which dilute phase transport of particles is occurring and that of the annulus which is a downward-moving packed bed with countercurrent flow of gas. The respective pressure drop gradients at various bed levels are in approximate balance, except in the vicinity of the gas inlet. At the top of a deep bed, the pressure gradient approaches that needed to support the solids and fluidize the material. When the gas velocity in the annulus becomes similar to the fluidization velocity, the spouted bed reaches the limit of its stability and such condition corresponds to the maximum spoutable bed depth. The pressure drop over an operating spouted bed is lower by about one-third for a similar fluidized bed, since the chamber wall support part of the weight of the particles. However, the pressure drop required to initiate spouting is considerable higher (Epstein and Mathur, 1982), which must be considered as an important factor when selecting the gas supply source. Similar to the minimum fluidizing gas velocity in conventional fluidized beds, there is a minimum gas velocity below which a spouted bed cannot be sustained. The value of this gas velocity is, however, dependent on both solid and fluid properties, along with bed geometry. Furthermore, it should be noted that there is a maximum spoutable bed depth beyond which the spouted bed may become a normal fluidized bed. The value of this depth is, also, dependent on solid and fluid properties, as well as on bed geometry.

The regime of stable spouting is critically dependent on certain conditions and, unless they are satisfied, the particles movement will become random leading to a state of aggregative fluidization and eventually to slugging if air flow is increased. The main variables involved are particle size, particle size distribution, gas-inlet diameter, column diameter, cone angle, gas flow rate, and bed height. The conditions for stable spouting are less critical for coarser particles and for wider columns. The minimum particle diameter for which the limits of stability become wide enough to be of practical interest appears to be in the 1–2 mm size range. Uniformity of particle size favors spouting

stability because the lower permeability of a bed containing a range of sizes would tend to distribute the gas more effectively rather than produce a jet action. The tolerance for nonuniformity in size will increase with increasing average particle size.

As previously mentioned, the vertical pressure gradient in a spouting bed increases from zero at the base to a maximum near the top of the bed. A total spouting pressure drop $\Delta P_{\rm SB}$ at the maximum spoutable bed depth $H_{\rm sm}$ can be estimated by the following relation:

$$\Delta P_{\rm SB} = \frac{2H_{\rm sm}\rho_{\rm s}(1-\epsilon_0)}{\pi} \tag{7.22}$$

where ε_0 is the voidage for the loosely packed bed.

The minimum gas velocity at which a bed will remain in the spouting state depends on solid and fluid properties, as well as on bed geometry. Unlike the minimum fluidization velocity, the minimum spouting velocity $u_{\rm ms}$ for a given material decreases with bed height and with increasing column diameter. In general, for a bed near its maximum spoutable depth, the minimum spoutable velocity is similar to the minimum fluidizing velocity of the material. Since the maximum spoutable bed depth in large diameter columns is much deeper than would normally be used in practice, the gas requirement for spouting in large columns would be lower than that for fluidization. The following correlation to estimate the minimum spouting velocity $u_{\rm ms}$, was originally proposed by Mathur and Gishler (1955) for cylindrical columns with short conical base:

$$u_{\rm ms} = \left(\frac{x}{D}\right) \left(\frac{D_{\rm i}}{D}\right)^{1/3} \sqrt{\frac{2gH(\rho_{\rm s} - \rho_{\rm f})}{\rho_{\rm f}}}$$
(7.23)

where x is an equivalent particle diameter, D is the column diameter, D_i is the diameter of gas inlet orifice, H is the bed depth, ρ_s is the particle density, and ρ_f is the fluid density.

At a depth just above the maximum spoutable, a spouted bed changes into a fluidized state. Therefore, if the minimum fluidization velocity for a solid material is known, its maximum spoutable bed depth in a given column can be estimated by substituting $u_{\rm mf}$ for $u_{\rm ms}$ in the minimum spouting velocity equation. A general expression for the maximum spoutable bed depth is given as follows:

$$H_{\rm sm} = K(x)^{n-1/m} (7.24)$$

where K is an empirical constant, while n and m are empirical exponents.

The spout diameter d_s for a wide range of materials, column diameter, orifice diameter, and bed depth, can be estimated by

$$d_{\rm s} = 14.5(0.115\log D - 0.192)\sqrt{\frac{G}{\rho_{\rm s}}}$$
 (7.25)

where *D* is the column diameter, and *G* is the gas mass flow rate.

Example 7.3

A batch of rapeseed is fluidized in a spouted bed at a pressure drop through the bed of 315 Pa. The column diameter of the bed is 15 cm while the air inlet diameter is 1.25 cm. The granules of rapeseed are 1.8 mm in diameter and their density is 1100 kg/m³. Calculate the minimum spoutable velocity for rapeseed.

SOLUTION

The rapeseed granules are practically spherical and relatively large. Thus, the voidage of the loosely packed bed can be taken as 0.4. The air density, consulted in tables of physical properties is around 1.2 kg/m³ at ambient temperature. Transposing the height of the bed from Equation 7.22:

$$H = \frac{(315)(3.1416)}{2(1100)(1-0.4)} = 0.7496 \approx 0.75 \,\mathrm{m}$$

Using this obtained bed height, along with the available data, substituting into Equation 7.23:

$$u_{ms} = \left(\frac{0.0018}{0.15}\right) \left(\frac{0.0125}{0.15}\right)^{1/3} \sqrt{\frac{2(9.81)(1100 - 1)}{1.2}} = 0.7496 \approx 0.75 \,\mathrm{m}$$

7.3 Applications of Fluidization

The use of fluidization processing was developed mainly by the petroleum and chemical industries for operations where the very high heat transfer coefficients and the high degree of uniformity of temperature within the bed enabled the development of processes, which would otherwise be impracticable. Fluidization technologies are nowadays used quite extensively in many industries where it is desirable to bring about intimate contact between small solid particles and a gas stream.

7.3.1 Applications in the Petroleum and Chemical Industries

The distillation of crude oil in the petroleum industry produce large surplus of high boiling materials. Cracking has been used to convert these materials into compound of lower molecular weight and lower boiling point, particularly petroleum spirit. The cracking was initially introduced in the industry by using a fixed catalyst, but local variations of temperature in the process led to a relatively inefficient operation. Also, the deposition of carbon on the surface of the catalyst particles created the need of taking the catalyst bed out of service periodically, in order to burn off the deposited carbon. Many of these difficulties are eliminated by the use of a fluidized catalyst, since it is possible to continuously remove the catalyst from the reaction vessel and to supply regenerated catalyst to the plant. The high heat transfer coefficients for fluidized systems account for the very uniform temperatures within the reactors, and make it possible to control conditions very closely.

Fluidized catalysts are also used in the synthesis of high-grade fuels from mixtures of carbon monoxide and hydrogen obtained either by coal carbonization or from partial oxidation of methane. An important application in the chemical industry is the oxidation of naphthalene to phthalic anhydride, which is an important reagent in the chemical industry used for large-scale production of plasticizers for plastics. It is also used in the chemical industry for the production of certain dyes.

7.3.2 Fluidized-Bed Combustion

A relevant application of fluidization that has attracted considerable interest for some time is fluidized-bed combustion. The combustible material is held in a fluidized bed of inert material while the air for combustion is the fluidizing gas. The system is being developed for steam raising on a very large scale, and it has been used for generation of electricity and for incineration of domestic refuse.

The particular features of fluidized combustion of coal that have risen interest are its suitability for use with very low-grade coals including those with high ash contents and sulfur concentrations, as well as its possibility of producing low concentrations of sulfur dioxide in the stack gases. This situation is due to the very much lower temperatures in the fluidized bed than those obtained in conventional grate-type furnaces, and the possibility of reacting of the sulfur in the coal with limestone or dolomite to enable its discharge as part of the ash.

The bed material normally consists of sand or ash, of particle size between 500 and 1500 μm . These materials gradually become replaced by ash from the coal and additives used for the sulfur removal. Ash is continuously removed from the bottom of the bed and, in addition, there is a considerable carryover by elutriation and the flyash is collected using cyclone separators. Coal has a lower density than the bed material and, therefore, tends to float,

but in a vigorously bubbling bed it can become well mixed with the remainder of the material and the degree of mixing determines the number of feed points required.

One of the major advantages of fluidized-bed combustion of coal is that it is possible to absorb the sulfur dioxide formed. Generally, limestone or dolomite is added and it breaks up in the bed to yield calcium oxide or magnesium and calcium oxide, which then react with the sulfur dioxide. It is possible to regenerate the solid in a separate reactor using gas consisting of hydrogen and carbon monoxide. There is some evidence that the reactivity of the limestone or dolomite is improved by the addition of chloride, but its use is normally avoided because it causes corrosion. Fluidized-bed combustion decrease pollution levels compared to conventional grate-type furnaces, because less oxides of nitrogen are formed.

7.3.3 Drying in Fluidized Beds

The principle of fluidized-bed dryers is that heated air is forced up through a bed of solids under such conditions that such solids are suspended in the air. The heated air acts both as fluidizing and drying medium. As a result, continuity of the drying process is possible and drying time may be shortened. While fluidized-bed drying has many advantages, particularly the continuity of operation just mentioned, it must be borne in mind that it has certain special features that may render it difficult to operate. For example, polydisperse materials may not be dried uniformly, stability of hydrodynamic conditions must be ensured, and dust must be trapped. The applicability of drying in a fluidized bed largely depends on the design of the equipment. It is accordingly very important to select the construction of drying equipment best suited to the aggregate state of the material and its physicochemical properties, as well as to the scale of production. A number of different designs of fluidized-bed dryers have given successful industrial service in recent years. Many new designs are undergoing laboratory test and field trials.

There is a great variety of designs of fluidized-bed dryers, and several criteria are used to classify them. From a technological standpoint they are divided into two groups: dryers for granular materials and dryers for pastes, solutions, suspensions and molten materials. With regard to operating conditions, they can be classified into continuous, semicontinuous and batch dryers. In terms of construction, they comprise single and multichamber types. As the advantage of fluidized-bed drying is giving continuity to a traditional batch process, continuous single- or multichamber operation for particulate materials is a regular option of operation in many applications.

One of the simplest and most widely used designs of fluidized-bed dryers is the continuously operated dryer in a cylindrical single chamber. As shown in Figure 7.8, wet material is continuously delivered from a hopper to the

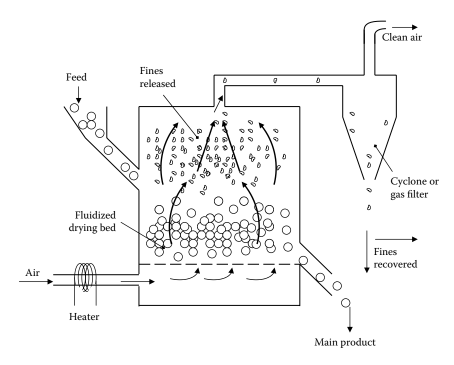


FIGURE 7.8 Diagram of single-chamber fluidized-bed dryer.

drying chamber which contains a fluidized bed of material. Hot gas from a furnace is mixed with air in the mixing chamber and delivered by a fan to the space beneath the grid. Dry product is discharged through the outlet located immediately above the grid on the opposite side of the feed hopper. The dust-laden exhaust gas passes to the cyclone and continues to the bag filter to be completely cleaned and returned to the atmosphere. Fines from the bottom outlets of cyclone and filter are combined with the dry product. Single-chamber dryers usually operate with shallow beds of materials (300–400 mm), high outputs (500–1000 kg/h moisture for 1 m² of grid), and low furnace gas consumption (up to 12 kg per kg of moisture).

Multichamber dryers contain a number of perforated shelves on which mesh or cloth is stretched, and in this way the flow of hot gas is uniformly distributed through the bed of material. They consist of a closed rectangular chamber divided into three sections one above another (Figure 7.9). The two upper sections are drying zones and the lower is a cooling zone. The grids slope at an angle of 2–3° to the horizontal. Wet material is delivered to the top chamber, it is gradually dried and passes to the second chamber and then to the cooling zone to be finally discharged through a flap valve. Multichamber dryers equipped with a mixer are used to ensure more uniform fluidization of very wet or finely divided materials.

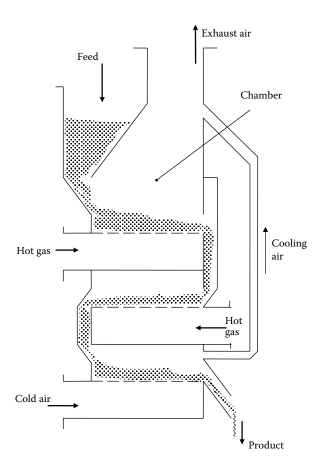


FIGURE 7.9 Multichamber fluidized-bed dryer.

Design equations used to determine drying time and dimensions of the drying equipment are required in order to develop methods of drying diverse materials in a fluidized bed. These equations should be derived on the basis of theoretical analysis of a physical model of the process, as well as on experimental determinations of the influence of the most important factors on its kinetics. Drying is a complex heat and mass transfer process including a number of additional effects which may cause undesirable characteristics in the product, such as shrinkage or case hardening. Drying in a fluidized bed has special features: the particles of moist material receive heat from a flow of gas, which acts both as a heat transfer and a fluidizing medium. The gas bulk flow rate is limited because the gas velocity in the drying chamber must be limited to avoid excessive entrainment of material. Difficulties could also arise because motion of individual particles of material in the drying zone is indeterminate and they may rotate, change in size,

and agglomerate. Furthermore, a proportion of the gas phase passes through the bed in the form of bubbles.

In fluidized-bed drying, normally the limiting kinetic factor is the process of external transfer of surface moisture from the material to the surrounding medium. In such a case, temperature gradients and variations in moisture content within the material are usually small and, thus, the material temperature is assumed constant and equal to the wet bulb temperature. The mechanism of heat transfer of the process is, therefore, considered to be convective. Under these circumstances it is assumed that the quantity of heat supplied determines the quantity of moisture evaporated. In drying processes by convective mechanism, at least two drying periods are recognized: a constant rate period and a falling rate period. The duration of the constant rate period is generally calculated from an energy balance using heat transfer equations. The most difficult problem is to determine the duration of the falling rate period. In practice, experimental drying curves are often used to determine the critical moisture content of the material. The tests are carried out on a small model which must reproduce the drying conditions such as temperature, air humidity, air velocity, and so on. During this falling rate period, the temperatures of the material and drying agent rise at all points in the fluidized bed. The distribution of the quantity of heat between evaporating moisture and heating the wet material is a function of the kinetics of internal heat and mass transfer. Equations that include the constant and falling rate periods have been proposed. For example, for drying grains in a spouted bed, an expression describing isothermal diffusion within grains is

$$\frac{w - w_{\rm s}}{w_1 - w_{\rm s}} = \frac{6}{\pi^2} \sum_{n=1}^{\infty} \frac{1}{n^2} \exp\left(-\frac{n^2 \pi^2 x^2}{9}\right)$$
(7.26)

where

$$x = \frac{S_{\rm p}}{V_{\rm p}} \sqrt{Dt} \tag{7.26b}$$

In Equations 7.26 and 7.26b w is the moisture content of the material at time t, $w_{\rm s}$ is the surface moisture content of the material, $w_{\rm l}$ is the initial moisture content of the material, $S_{\rm p}$ is the surface area of particles, $V_{\rm p}$ is the particle volume, and D is the diffusion coefficient.

7.3.4 Coating of Particles and Particulates

Coating or encapsulation of particles, granules, or particulates can be carried out by a number of unit operations. Some of the operations used for mixing or dry powders can be equally applied for coating. Equipment that can be used for the purpose of mixing powders/coating particulates includes

Fluidization 273

rotating drums, ribbon mixers, and fluidized beds (Ortega-Rivas, 2005). Fluidized-bed coating was developed in the 1950s, and it is normally known as "Wurster process" (Arshady, 1993). The application of the fluidized-bed technique for the coating of particles aims to manufacture a product of uniform quality and morphology, which is basically determined by the actual physical state of the droplets (Eichler, 1989). Droplet formation, contact, spreading, coalescence, and evaporation, are occurring almost simultaneously during the process (Jones, 1985). The nozzles typically used in fluidized-bed coating are either binary or pneumatic; liquid is supplied at a low pressure and is sheared into droplets by air. Droplet size and distribution are more controllable with this type of nozzle than with a hydraulic one, especially at low liquid flow rates (Filková and Mujumdar, 1995). Furthermore, the air used for atomization also contributes to evaporation of the coating solvent, increasing the droplet viscosity and inhibiting spreading and coalescence on contact with the core material (Jones, 1985). There have been several designs of fluidized-bed configurations over the years. The conventional top-spray method (Figure 7.10) evolved from fluidized-bed dryers. The particles are accelerated from the product container past the nozzle, and so they are sprayed counter-currently by the stream of atomized droplets. The coated particles travel through the coating zone into the expansion chamber, falling back into the product container and continue cycling throughout the process (Jones, 1988). In the bottom-spray method (Figure 7.11) the particles are recycled through the coating zone in seconds, as in the top-spray technique, but the fluidization pattern is much more controlled. The droplets and particles travel cocurrently, avoiding almost completely premature droplet evaporation. The film-forming droplets can spread out at lowest viscosity, producing a very dense film (Mehta and Jones, 1985). A relatively new approach to coating is the tangential-spray method, with a rotary base, as shown in Figure 7.12. A combination of three forces occurs during fluidization, which has been described as spiralling helix. A centrifugal force by a rotating disc causes the product to move toward the wall of the chamber, the air velocity through an annular gap provides acceleration upward, while gravity cascades the product inwards and toward the disc once again. Beneath the surface of a rapidly tumbling bed, a nozzle is positioned to spray the coating tangentially to and concurrently with the flow of particles (Jones, 1988).

A way of evaluating coating efficiency is by determining the deposited coating mass m_c , and relating it to the coating solution dry matter content DM. Dewettinck and Huyghbaert (1998) have suggested the following expression for the evaluation of coating efficiency E_c :

$$E_{\rm c} = \frac{m_{\rm c}}{m_{\rm cs} \cdot {\rm DM}} \tag{7.27}$$

where m_{cs} is the mass of the coating solution.

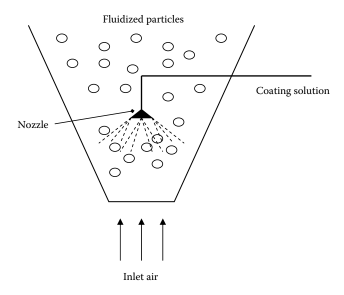


FIGURE 7.10 Diagram of top-spray fluidized-bed equipment.

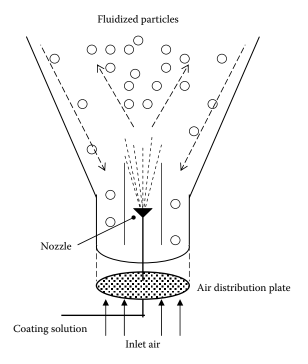


FIGURE 7.11 Diagram of the bottom-spray (Wurster) fluidized-bed coater.

Fluidization 275

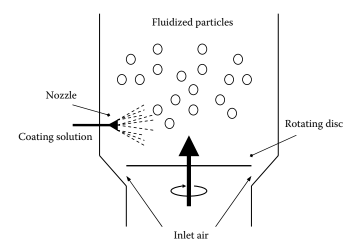


FIGURE 7.12 Diagram of tangential-spray, rotor, fluidized-bed coater.

Coating thickness $d_{\rm c}$ is sometimes difficult to measure conveniently due to the fine size of some powders and to the unevenness of the surface of some granular materials. A reasonably estimate of the coating thickness for spherical microcapsules can be calculated using the following expression (Dewettinck, 1977):

$$d_{c} = \left\{ \left[\left(\frac{\rho_{s}}{\rho_{c}} \right) \cdot \left(\frac{m_{s} + m_{c}}{m_{s}} - 1 \right) + 1 \right]^{1/3} - 1 \right\} \cdot \left\{ \frac{x}{2} \right\}$$
 (7.28)

where ρ_s is the density of the core material, ρ_c is the density of the coating material, m_s is the mass of the core material, m_c is the mass of the coating material, and x is an equivalent diameter of the core material.

An important advantage of fluidized-bed coating is its ability to minimize attrition of the particles being processed. Attrition represents one of the major problems in food powder processing with many adverse effects such as segregation of fines from the bed structure, degradation of product quality, changes in bulk flow behavior, and reduction of reconstitutability of the product (Ortega-Rivas, 2005). The segregated fines can be released to the surrounding atmosphere causing dust emissions and constituting a safety problem. Among all the problems associated with dust emissions, such as health hazards, abrasion damage to equipment, impaired visibility, and so on, the risk of dust explosion is one of the greatest concerns in many dry powder operations, such as industrial grain and cereal processing (Barbosa-Canovas et al., 2005).

Reduction of attrition can be also used as an indicative of the coating efficiency. A manner to determine degree of attrition in coated materials can be done by performing a friability test, which is widely used in different agglomerated and compacted food and pharmaceutical products. A friability tester consists, simply, of a rotating disc or drum on which materials are subjected to tumbling at different times and rotation speeds. Friability is determined as the difference of weighed sample before and after the test. The tumbling action in a friability tester replicates typical handling situations, to predict how well coated products will withstand handling during transportation and storage.

7.4 Laboratory Exercise: Fluidized-Bed Coating of Food Particulates

7.4.1 Introduction

Coating with sweet coverings is used in product development in the food industry, particularly in obtaining ready-to-eat breakfast cereals. Different edible coatings in cereals are applied in order to obtain advantages such as to enhance flavor, protect the food particulate from oxygen and moisture, or improve quality and functionality. A well applied and thick enough coat will be determinant in accomplishing these attributes and, also, will provide resistance to attrition or erosion due to handling and distribution. The exercise is aimed at testing different variables in coating food granules and verifying uniformity, thickness, and firmness of the applied coat.

7.4.2 Equipment and Materials

A batch fluidized bed of the type of a Glatt[®] laboratory unit. Unsweetened puffed cereal, such as puffed wheat or puffed rice. Ingredients to prepare the coating syrup: sugar, cocoa, and water.

7.4.3 Instruments and Apparatuses

Optical microscope, analytical balance, friability tester.

7.4.4 Operation of the Fluidized Bed

- Prepare 2–3 L of syrup mixing 80 g cocoa and 430 g sugar in 400 mL water.
- Load the fluidizing unit with a batch of 400–500 g of the particulates.

Fluidization 277

• Operate the unit at ambient conditions and at an air velocity enough to fluidize the particulates (1–3 m/s).

- Spray the syrup, once the particulates are completely suspended, at a rate of about 15 mL/min.
- Run the process for 10, 15, and 20 min.
- Stop the fluidizing unit and discharge the product.
- Dry the coated granules placing them on a tray at room temperature, or proceeding to a mild drying in a tray drier.

7.4.5 Evaluation of the Coating Thickness

- Cut out, very carefully, several thin slices of several granules coated at different times.
- Observe them at the microscope.
- Estimate the coat thickness by using an appropriate eyepiece graticulate.

7.4.6 Friability Test

- Weigh the sample and introduce it in the drum of the tester.
- Run the tester, according to specific instructions of the apparatus. A number of applications give satisfactory results at 20–25 rpm for 20–30 min.
- Unload the drum (if not done automatically) and weigh the processed sample.

7.4.7 Calculations and Report

- Determine the coating efficiencies, using Equation 7.27, for the three different samples. Record observations for further analysis.
- Compare the observed coat thicknesses with calculated thicknesses using Equation 7.28, for the three treatments of samples.
- From friability testing determine friability as the difference of weighed samples at the different conditions before and after the test. Express the friability in weight loss percentage.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out. Include analysis from own data and, as much as possible, discussion based on literature survey from studies and investigations on the subject.

PROBLEMS

- 1. The approach velocity of a gas ($\rho = 1.3 \text{ kg/m}^3$, $\mu = 1.65 \times 10^{-5} \text{ kg/m s}$) flowing through a bed of spherical particles is 0.032 m/s. If the bed is 2.9 m thick, determine the pressure drop developed.
- 2. A reactor contains catalyst particles shaped into cylinders 1.25 cm diameter and 2 cm high. A gas having a density of 1.28 kg/m³ and a viscosity of 1.73×10^{-5} Pa s flows at a steady rate through the bed, which has a voidage of 0.4. If the superficial velocity established is 0.6 m/s and the pressure drop through the bed is 2.5 kPa, calculate the bed height.
- 3. Anthracite coal of 1500 kg/m³ density is to be fluidized with air at 140 kPa in a vessel 3 m in diameter. 48 tons of coal, ground to an average diameter of 70-mesh (US Standards) are placed in the vessel, which is filled to a height of 2 m. Calculate the static void fraction and the minimum void fraction at height for fluidization.
- 4. Calculate the air velocity required to fluidize 50 kg of a polymer consisting on spherical particles of 3 mm diameter and 700 kg/m³ density. The fluidized medium is air of 1 kg/m³ density and 1.6×10^{-5} kg/m s viscosity. Estimate also the pressure drop if the particles are contained in a 40 cm diameter vessel.

References

Abrahamsen, A. R. and Geldart, D. 1980. Behaviour of gas fluidized beds of fine powder. Part I: Homogeneous expansion. *Powder Technology* 26: 35–46.

Arshady, R. 1993. Microcapsules for food. Journal of Microencapsulation 10: 413-435.

Barbosa-Canovas, G. V., Ortega-Rivas, E., Juliano, P., and Yan, H. 2005. *Food Powders: Physical Properties, Processing, and Functionality*. New York, NY: Kluwer Academic/Plenum Publishers.

Dewettinck, K. 1977. Fluidized Bed Coating in Food Technology: Process and Product Quality. PhD Thesis. Ghent, Belgium: University of Ghent.

Dewettinck, K. and Huyghbaert, A. 1998. Top-spray fluidized bed coating: effect of process variables on coating efficiency. *Lebensmittel-Wissenchaft und Technologie* 31: 568–575.

Eichler, K. 1989. Principles of Fluid-bed Processing. Binzen, Germany: Glatt GmbH.

Epstein, N. and Mathur, K. B. 1982. Applications of spouted beds. In *Handbook of Multiphase Systems*, Hetsroni, G. (ed.). New York, NY: McGraw-Hill.

Ergun, S. 1952. Fluid flow through packed columns. *Chemical Engineering Progress* 48: 89–94.

Filková, I. and Mujumdar, A. S. 1995. Industrial spray drying systems. In *Handbook of Industrial Drying*, Vol. 1, Mujumdar, A. S. (ed.) New York, NY: Marcel Dekker.

Jackson, R. 2000. *The Dynamics of Fluidized Particles*. Cambridge, UK: Cambridge University Press.

Jones, D. M. 1985. Factors to consider in fluid-bed processing. *Pharmaceutical Technology* 9: 50–62.

Fluidization 279

Jones, D. M. 1988. Controlling particle size and release properties. In *Flavor Encapsulation*, Risch, S. J. and Reineccius, G. A. (eds.) Washington, DC: American Chemical Society.

- Mathur, K. B. and Gishler, P. E. 1955. A technique for contacting gases with coarse solid particles. *AIChE Journal* 1: 157–164.
- McCabe, W. L., Smith, J. C., and Harriot, P. 2005. *Unit Operations in Chemical Engineering*. 7th Ed. New York, NY: McGraw-Hill.
- Mehta, A. M. and Jones, D. M. 1985. Coated pellets under the microscope. *Pharmaceutical Technology* 9: 52–60.
- Ortega-Rivas, E. 2005. Handling and processing of food powders and particulates. In *Encapsulated and Powdered Foods*, Onwulata, C. (ed.) Boca Raton FL, USA: CRC Taylor & Francis.
- Rhodes, M. 1998. Introduction to Particle Technology. Chichester, UK: John Wiley & Sons.
- Toomey, R. D. and Johnstone, H. F. 1952. Gaseous fluidization of solid particles. *Chemical Engineering Progress* 48: 220–237.
- Wilhelm, R. H. and Kwauk, M. 1948. Fluidisation of solid particles. *Chemical Engineering Progress* 44: 201–218.
- Zenz, F. A. and Othmer, D. F. 1960. *Fluidization and Fluid-Particle Systems*. New York, NY: Reinhold.

This page intentionally left blank



Part III

Separation Techniques for Particulate Solids

This page intentionally left blank



Introductory Aspects

8.1 Different Mixtures Relevant in Industry

In the material processing industry a great number of raw materials and products formed by several components are treated. Not only mixtures involving phases of the same state of matter, but also two-phase and multiphase systems are normally encountered in a number of processes. Many mixtures or multiphase systems often need to be purified or depurate in order to obtain products, as free as possible, from impurities. For example, caustic soda and salt are treated in evaporators, an important number of food products are dehydrated, some ceramic and colorants are filtered, while sugars and fruit juices may be centrifuged. Depuration or separation of mixture components can be carried out in a number of ways, and technologies to do so involve the use of some kind of energy. The ways to separate based on heat or mass transfer constitute a series of well-studied unit operations, while the ones known as mechanical separations were somewhat neglected in higher-education programs for a long time (Svarovsky, 2000). A common problem facing process engineers in day-to-day practice in different materials processing industries is selecting, optimizing, or operating techniques involving the separation of one or more components of a mixture or a complex fluid.

8.2 Classification of Separation Techniques

A number of unit operations have as fundamental purpose the separation of two or more components within a system. Mixtures and multiphase fluids previously mentioned are varied in the materials processing industries and include solutions, suspensions, dispersions, and solid blends. The components within these systems may include any of the three common states of matter (i.e., solid, liquid, or gas), and so different combinations are possible. There are many criteria used to determine the most suitable strategy to

perform a specific separation. Some factors that could be considered to define or select a method of separation are

- Differences in physical properties of the compounds
- Differences in chemical properties of the compounds
- · Possibility of mass transfer
- Possibility of phase changes in some of the compounds
- Use of natural forces to achieve the separation
- Use of external forces to achieve the separation

Analyzing briefly the above-listed criteria, it is evident that classification of separation techniques is not an easy task, and that it is complicated to even consider a unified categorization. There are a number of separation techniques bearing little or no interest for certain industries, while there are techniques forming a minimal part of a more general unit operation. For example, magnetic separation is a separation technique encompassed by a unit operation of great importance in food engineering: the cleaning of raw materials. Independently of referring to a general interest classification or one of interest exclusively for a branch of engineering, two commonly used criteria to propose a classification of separation techniques are

- Classification of the separation techniques according to those involving phase change or mass transfer from one phase to another, known as diffusional operations, and those that are useful in the separation of solid particles or drops of a liquid and that are generally based in the application of an external physical force, known as mechanical separations.
- Classification based on the separation techniques according to the different combinations possible due to the states of matter (phases) found in any bicomponent or multicomponent system.

Two possible classifications of separation techniques based on the above criteria are given in Tables 8.1 and 8.2.

8.3 Specific Techniques for Granular Materials

Several separation techniques listed in the previous section are used to separate finely divided solids within a bicomponent system (Rushton et al., 2000). A thorough understanding of particle technology principles will therefore be useful not only to design storage and conveying systems (Part I) or

TABLE 8.1Classification of Separation Techniques Based on Diffusion Properties

- <u></u>	•		
Diffusional Operations	Mechanical Separations		
Phase equilibrium	Sieving		
Distillation	Sedimentation		
Leaching and extraction	Centrifugation		
Operations of multicomponents by stages	Filtration		
Diffusion and mass transfer between stages			
Gas absorption			
Humidification operations			
Solids drying			

optimally operate processing equipments (Part II), but also in order to have a greater chance of success in selecting, designing, or operating a system that involves separation techniques were one of the phases is a powder or particulate solid (Cheremisinoff, 1995). In this context, only some of the previously listed techniques, for example, liquid–liquid and liquid–gas systems would be particularly related to particle technology. Diffusional operations related to heat and/or mass transfer are based on principles out of the scope of particle technology and are widely addressed in classical chemical engineering texts. This final part of the book will be, therefore, dedicated to the study of operations known as mechanical separations, mechanical–physical separation processes, or physical separations. The two kinds of two-phase systems that will be studied are the solid–solid and solid–fluid mixtures. The usual way to separate the components of a solid–solid system is in dry, making use of differences in properties such as size, shape, density, as well as electric or magnetic characteristics. For the case of the solid–fluid mixtures

TABLE 8.2Classification of Separation Techniques Based on Involved Phases

	1	1		
Liquid-Liquid Systems	Liquid–Solid Systems	Liquids with Dissolved Solids	Solid-Gas Systems	Solid Mixtures
Distillation	Sedimentation	Separation by membranes	Cyclones separation	Sieving
Liquid extraction	Centrifugation	Hydrometallurgical extraction	Gas filtration	Electromagnetic separation
Decantation	Hydrocyclone separation	Evaporation	Gravitational methods	Electrostatic separation
Ionic exchange	Filtration	Crystallization of solutions	Scrubbing	
Dieresis and electro dialysis	Drying of wet solids	Flotation		

the most common case is when a fluid is a liquid, usually water, and techniques of great industrial importance, such as filtration and centrifugation are included.

References

Cheremisinoff, P. N. 1995. Solids/Liquids Separation. Boca Raton, FL: CRC Press.
 Rushton, A., Ward, A. S., and Holdich, R. G. 2000. Solid-Liquid Filtration and Separation Technology. Weinhem, Germany: Wiley-VCH.
 Svarovsky, L. (ed.) 2000. Solid-Liquid Separation. Oxford: Butterworth-Heinemann.

Solid Mixtures

9.1 Screening

9.1.1 Basic Principles

Screening is a separation technique of a mixture of various sizes of solids particles into several fractions, based on size difference. It consists of forcing the mixture through a screen of a specific size aperture. Small-capacity plane screens are often called sieves. By vibrating or oscillating a screen, particles smaller than a given aperture pass through this being thus separated from the remaining mixture. Screens are made from metal bars, perforated or slotted metal plates, woven wire cloth, or fabric, such as silk bolting cloth. Metals used include steel, stainless steel, bronze, copper, nickel, and monel. The screen surface may be plane (horizontal or inclined) or it may be cylindrical. The size aperture of the screens, range from about 0.1 mm to approximately 250 mm, with exceptional cases in which the aperture may be as large as 460 mm.

Screening has two main applications: laboratory technique for particle size analysis, and industrial operation for fractionation and classification of particulate solids. Although screen aperture, defined as the space between the individual wires of a wire mesh screen, is the preferred terminology for screening operations, the former designation of mesh number, defined as the number of wires per lineal inch, is still widely adapted. In both applications, shape and size of particles may influence separation and efficiency. A combined shape–size effect may cause blinding or clogging of the screen openings, and is particularly likely to occur when particles have sizes very close in dimension to the screen aperture. In general terms, there would be three defined possibilities of a given particle facing a screen aperture: (a) the particle being too large in relation to the aperture so that it would be easily retained, (b) the particle being too small in relation to the aperture so that it would go through easily, (c) the particle having a critical dimension so that it would be trapped and promote blinding or clogging of the screen surface. This mentioned critical dimension has been defined when the particle has an approximate size of 1.1D (Figure 9.1), which happens if the angle θ between the particle and the screen aperture is less than tan⁻¹ μ, μ being the

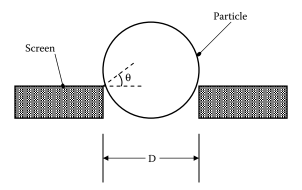


FIGURE 9.1Relation between particle size and sieve aperture to cause clogging.

coefficient of friction between the particle and the screen material. The extreme case of blinding produces complete clogging of the screen with a consequent damage that would impair separation and operation efficiency. Damaged screens should, therefore, be repaired or replaced immediately. It has been demonstrated that clogging is affected by the size of the screen aperture and the particle shape (Beddow, 1980). It has also been reported that particle shape has a significant effect in efficiency for circular and rectangular screen apertures, but only a minor effect for square screen apertures (Nakayima et al., 1978).

The material passing through a given screen is termed undersize, fines or minus (-) material, while the material retained in a given size screen is called oversize, coarse or plus (+) material. Either stream may be the desired (product) stream or the undesired (reject) stream. The objective of a screening operation is to separate a feed stream into two fractions, an underflow that is passed through the screen and an overflow that is rejected by the screen. An ideal screen would sharply separate the feed in such a way that the smallest particle in the overflow would be just larger than the largest particle in the underflow. Such an ideal separation would define a cut diameter D_{pcl} which would represent the point of separation between the fractions, as indicated in Figure 9.2a. For an ideal operation, a plot of screen opening against cumulative fraction retained would have the shape shown in Figure 9.2b. As can be seen, the largest particle of the underflow has the same size of the smallest particle of the overflow. In practice, however, the plot would have the shape illustrated in Figure 9.2c, in which there is an overlap. The underflow has an appreciable amount of particles larger than the desired cut diameter while the overflow has particles smaller than such desired cut diameter. It has been observed that the overlap is small when particles are spherical (or close to a spherical shape) and is larger when particles are needle like, fibrous, or tend to agglomerate. The main problems

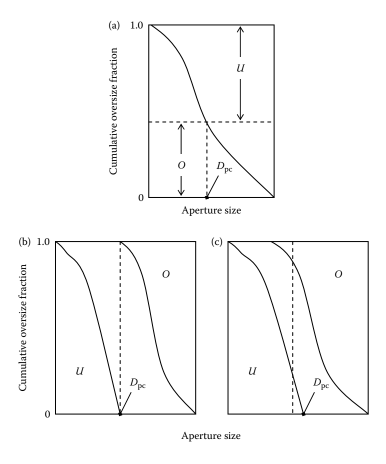


FIGURE 9.2 Diagram of cumulative frequency of oversized particles for: (a) feed, (b) ideal separation, and (c) real separation.

encountered in screening result from sample stickiness, sieve blockage, and agglomeration. According to Strumpf (1986), all these problems increase exponentially as the screen aperture decreases.

9.1.2 Design and Selection Criteria

The efficiency of a screening operation may be evaluated by simple mass balances. Let F be the mass flow rate of feed, O the mass flow rate of coarse fraction, and U the mass flow rate of fines; also, let X_F be the coarse mass fraction in the feed, X_O the coarse mass fraction in the overflow and X_U the coarse mass fraction in the underflow. Furthermore, the fines fractions in the feed, the overflow and the underflow would be $1 - X_F$, $1 - X_O$, and $1 - X_U$,

respectively. Since the total of the material fed to the screen must leave either as overflow or underflow:

$$F = O + U \tag{9.1}$$

The coarse fraction in the feed must also leave in the two streams, and so

$$FX_{\rm F} = OX_{\rm O} + UX_{\rm H} \tag{9.2}$$

Elimination of U from Equations 9.1 and 9.2 gives

$$\frac{O}{F} = \frac{(X_{\rm F} - X_{\rm U})}{(X_{\rm O} - X_{\rm U})} \tag{9.3}$$

Similarly, elimination of O gives

$$\frac{U}{F} = \frac{(X_{\rm O} - X_{\rm F})}{(X_{\rm O} - X_{\rm U})} \tag{9.4}$$

The effectiveness of a screen is a measure of how well it performs the separation of coarse and fine fractions. If the screen functioned perfectly, all of material O would be in the overflow and all of material U would be in the underflow. A manner of determining screen efficiency would be by calculating the ratio of oversized material O that is actually in the overflow to the amount of material O entering with the feed, that is,

$$E_{\rm O} = \frac{OX_{\rm O}}{FX_{\rm F}} \tag{9.5}$$

Similarly, considering the fine material:

$$E_{\rm U} = \frac{U(1 - X_{\rm U})}{F(1 - X_{\rm E})} \tag{9.6}$$

An overall combined efficiency may be defined as the product of Equations 9.5 and 9.6, and it may be denoted simply as (Foust et al., 1960)

$$E = \frac{OUX_{\rm O}(1 - X_{\rm U})}{F^2 X_{\rm F}(1 - X_{\rm F})}$$
(9.7)

Substituting Equations 9.3 and 9.4 into Equation 9.7 gives

$$E = \frac{(X_{\rm F} - X_{\rm U})(X_{\rm O} - X_{\rm F})X_{\rm O}(1 - X_{\rm U})}{(X_{\rm O} - X_{\rm U})^2(1 - X_{\rm F})X_{\rm F}}$$
(9.8)

TABLE 9.1 Example 9.1

Sieve Aperture	Cumulative Fraction			
Size (µm)	Feed	Coarse	Fine	
850	0.02	0.06	_	
710	0.12	0.28	_	
600	0.26	0.52	0.02	
500	0.45	0.78	0.08	
425	0.68	0.90	0.24	
355	0.82	0.96	0.46	
300	0.90	1.00	0.68	
212	0.98	_	0.84	

Equation 9.8 is an alternative expression to evaluate screen efficiency without involving the streams and only using the fractions.

Example 9.1

Data on a screening operation is presented in Table 9.1. Particle size distributions of feed, overflow, and underflow are given as cumulative frequency. The screen used for separation has an aperture size of 460 μ m, and 1000 kg/h of feed are processed obtaining 650 kg/h of overflow. Calculate the efficiency of the operation.

SOLUTION

Having the particle size distributions of the three streams, as well as two of their flow rates, Equation 9.8 can be used to evaluate efficiency. Fractions $X_{\rm F}$, $X_{\rm O}$, and $X_{\rm U}$ can be determined from a plot of equivalent sieve diameter versus cumulative frequencies of the feed, overflow, and underflow, at the cut diameter $D_{\rm pc}$. Plotting data from Table 9.1, the graph in Figure 9.3 is obtained.

Consulting values on this graph: $X_O = 0.14$, $X_F = 0.58$, and $X_U = 0.85$. From a simple mass balance from Equation 9.1 the flow rates are: F = 1000 kg/h, O = 650 kg/h, U = 350 kg/h, and thus, substituting values in Equation 9.7:

$$E = \frac{(650)(350)(0.86)(1 - 0.15)}{(1000)^2(0.58)(1 - 0.58)} = 0.6827$$

The overall efficiency of the operation is, therefore, 68%, approximately.

Example 9.2

The fine fraction of a catalyst retained on a 7-mesh screen (Tyler) is to be separated to be used in a melting process. The particle size analysis of fractions obtained is given in Table 9.2. Calculate the efficiency of the screening operation.

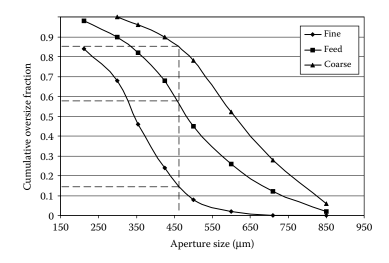


FIGURE 9.3 Example 9.1.

SOLUTION

Consulting the Tyler Standard Screen Scale in Appendix 5, a Table of equivalent sieve diameter as a function of cumulative frequency is derived. From such a table, a plot similar to the one derived in the previous example is obtained. The plot is shown in Figure 9.4, and values of $X_{\rm O}$, $X_{\rm F}$, and $X_{\rm U}$, at the cut diameter $D_{\rm pc}$ of 7-mesh (2.794 mm from the Tyler Standard Screen Scale) are 0.68, 0.2, and 0, respectively.

TABLE 9.2 Example 9.2

	Cumulative Fraction			
Mesh	Feed	Coarse	Fine	
4	0.00	0.30	_	
6	0.07	0.56	_	
8	0.43	0.84	0.00	
10	0.85	0.92	0.20	
14	0.97	0.96	0.58	
20	0.99	1.00	0.83	
28	1.00	_	0.91	
35	_	_	0.94	
65	_	_	0.96	
Pan	_	_	1.00	

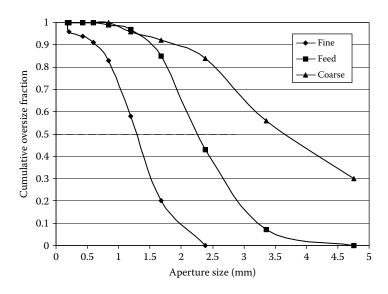


FIGURE 9.4 Example 9.2.

The value of $X_U = 0$ would foresee a high screen efficiency. Substituting values in Equation 9.8:

$$E = \frac{(0.2 - 0)(0.68 - 2.0)(0.68)}{(0.68)^2(0.2)(1 - 0.2)} = 0.8823$$

The overall efficiency of the operation, as expected, is high of about 88%.

Although accurate relationships are not available for estimating operating characteristics in screen operation, certain fundamentals apply and may be used as guidelines when running and designing a screening process. Efficiency of separation, along with capacity, is the most important variable involved in industrial screening. Capacity and effectiveness are opposing factors as maximum effectiveness is related to small capacity, while large capacity is only attainable at the expense of efficiency. Industrial screening is normally operated at high capacities, so that efficiency is expected to be lower when compared with laboratory screening, which is operated at a lower scale. Efficiencies between 85% and 95% are considered reasonable for both scales of screening.

The capacity of a screen is measured by the mass of material that can be fed per unit time to a unit area of screen, and can be simply controlled by varying the feed rate to the equipment. The effectiveness obtained for a given capacity is dependent on the specific nature of the screening operation. The chance of passing through the screen of an undersized particle is a function of the number of times the particle strikes the screen surface, as well as its probability of passage in a single contact. If a screen is overloaded, the number of contacts

is small and the chance of passing on contact is reduced by particle interference. The improvement of effectiveness obtained at the expense of reduced capacity is a result of more contacts per particle and better chances for passing through the screen aperture on each contact. A particle would have an ideal opportunity of passage when striking the surface perpendicularly, which would only be possible if it were oriented with its minimum dimensions parallel to the screen surface. Additional conditions would be no interference by other particles, as well as not sticking to, or wedging into, the screen surface. None of these conditions apply to actual screening, but this ideal situation can be used as a basis for estimating the effect of mesh size and wire dimensions on screen performance. If the width of a screen would be negligible in comparison with the size of the openings, the wires would not interfere with particle passage and, practically, the entire screen surface would be active. In such a case, the probability of passage of a striking particle would approach unity. In actual screening the diameter of the wire, or the fraction of the surface not constituting openings, is significant and the solid meshes strongly affect screen performance, especially by retarding the passage of particles nearly as large as the screen openings. When large and small particles are present, the large ones tend to segregate in a layer next to the screen preventing the smaller particles from reaching the screen surface. All these factors tend to reduce capacity and lower efficiency.

Based on the information given and discussed above, a processing engineer should be able to evaluate capacity and efficiency for a particular screening industrial operation. In order to avoid problems and come up with the most suiTable choice for a process involving screening, as important as these mentioned characteristics, other details can be equally important. Structural supports will be under varying conditions and so attention must be given to adequately sizing them, not only for the loads to be carried out but for the deflection and vibratory conditions that could prevail in operation. Feed and product chutes also deserve careful consideration. The feed to the screen must be delivered so as to cause minimal amount of abrasion or disturbance of the bed of material on the screen. The trajectory of material being discharged from feed conveyors or other units of equipment must be considered, and the force of the falling material directed against an abrasive-resistant wear plate, or a dead-bed of the material itself. The screening operation should be considered within the context of a whole process flow sheet. A description of the unit operations immediately preceding and following is quite relevant. Details of the equipment adjacent to the screening step must be carefully considered. For example, there is little point in dry screening a dusty material if the succeeding stages are wet. The screening operation involved should also describe the methods used for controlling feed rate, product collection, required screen efficiency, number and size of products, and so on.

Several relevant properties of the material being separated are to be known or determined to properly select or design appropriate screening equipment. Some of these include: particle size distribution, particle shape, bulk density,

moisture content, abrasiveness, and corrosiveness. The particle size distribution is essential to correctly size the screening unit, as well as to specify the type of screen to be installed. As previously mentioned, particle shape is determinant in promoting, or avoiding, blinding or clogging of screen units. Long or splinter-like, round or oval, or cubic particles will have slightly different screening characteristics, and may have a great influence on choice of cloth opening. Bulk density permits the determination of the volume of flow, and a measure of the load to be carried by the screen. Moisture content may cause difficulties due to stickiness. Information on moisture content, along with data on the process following screening, will allow selection of dry, damp, or wet screening. In some dry-screening applications, when moisture content is low, the choice of a proper screen medium will eliminate problems. Predrying of materials using heated screen cloths, or going wet by adding water sprays, are all equally effective in damp screening. The abrasive characteristics of the material have great influence on choice of materials of construction, as well as in selection of methods of loading, collecting, and transporting products. Corrosive features will also influence materials of construction used in screen frames, media, chutes, feedboxes, and other elements of the system. This property is relevant for choosing between dry and wet screening.

Example 9.3

A granular material has been analyzed giving the results presented in Table 9.3. The product from which the sample was taken is separated using a screen of 1 mm opening, and the fine fraction is to be used for a specific process. The screen is fed at a rate of 8660 kg/h, obtaining 1960 kg/h of coarse material and, practically, no particles smaller than 1 mm. Since the fraction of interest is the fine one, calculate the efficiency of the operation, expressed as separation of fines.

SOLUTION

In this case, tabulated data include only the particle size distribution of feed. The retained fraction is given, rather than the cumulative fraction. Following a procedure similar to that in Example 2.4, a plot of cumulative frequency can be obtained. Since $D_{\rm pc}$ is 1 mm, the feed fraction $X_{\rm F}$ can be read directly from Table 9.3, as

TABLE 9.3 Example 9.3

1			
Particle Size Range	Weight (g)		
Larger than 2 mm	0.0		
Between 2 and 1 mm	19.8		
Between 1 and 0.5 mm	58.2		
Between 0.5 and 0.1 mm	10.2		
Smaller than 0.1 mm	11.8		

0.198. From a mass balance using Equation 9.1, U = 6700 kg/h. Since there are no fines within the coarse stream, fractions X_O and X_U are 1.0 and 0, respectively. Substituting into Equation 9.6:

$$E_{\rm U} = \frac{(6700)(1-0)}{(8660)(1-0.198)} = 0.9646$$

The efficiency, expressed in terms of separation of fines is, thus, 96.5% approximately.

9.1.3 Equipment Used

Screening as a unit operation may be carried out in different types of equipment. Three types are more common: grizzlies (bar screens); shaking, vibratory or oscillating screens; and revolving screens or trommels. Grizzlies are used for screening larger particles (pieces greater than 25 mm). They consist of a set of parallel bars, spaced to the desired separation. The bars are often wedge shaped to minimize clogging. They may be used horizontally or inclined at angles up to 60°. Vibrating grizzlies are available, the feed material passing over the screening surface in a series of jerks.

The machines simply termed as screens can consist of many types: vibrating, shaking, reciprocating, and oscillating, to name only a few. Industrial screening equipment is used over a wide range of particle sizes. For small particles woven cloth or wire screen are used, while for larger sizes perforated metal plates or grids can be employed. Dynamic screening equipment can be classified according to the type of motion used to shake up and transport the material on the screen. Vibrating screens are used as standard practice when large capacity and high efficiency are required. The capacity, particularly in the finer sizes, is greater by far when compared with any other of the screen types, so that they are normally preferred when efficiency of the screen is the governing factor of a given process. Other advantages of vibrating screens include accuracy of sizing, increased capacity per unit area, low maintenance costs, and saving in installation space. The screens used in this type of equipment can be mounted horizontally or inclined and their operating range of frequencies is high (1000–7000 Hz).

Shaking screens consist on a rectangular frame, holding wire cloth or perforated plate, slightly inclined and suspended by loose rods or supported from a base frame by flexible flat springs. The frame is driven with a reciprocating motion, and the material to be screened is fed at the upper end advancing by the forward stroke of the screen. As the feed advances on the screen surface, finer particles pass through the screen openings. Shaking screens may be conventionally used for both screening and conveying of a number of materials. Their advantages include the low headroom and power requirement, while some disadvantages are the high maintenance cost of the screen and the supporting structure due to vibration, and the low capacity compared to inclined high-speed vibrating screens.

Reciprocating screens are operated with a shaking motion, long stroke, at frequencies between 20 and 200 Hz. They produce size fractions removing the coarse material first and the fines last. These machines contain several decks of screens, one above the other, held in a box or casing. The coarsest screen is located at the top while the finest is mounted at the bottom. Suitable discharge ducts allow proper removal of the several fractions. Figure 9.5 shows a diagram of a reciprocating screen. The feed is dropped on the top screen, and screens and casing are gyrated in order to sift the particles through the screen openings. In the vertically gyrated design the casing is inclined at an angle of 16-30° to the horizontal plane. The gyrations are vertically about the horizontal axis. The screens are rectangular shaped and fairly long, with typical dimensions of 0.46×1.2 m to 1.5-4.3 m. The speed of gyration and the amplitude of throw are adjustable, and also is the angle of tilt. A particular combination of speed and throw would normally give the maximum yield of required product from a given speed. Satisfactory separation into fine fractions usually requires an angle not more than 20°. A horizontally gyrated screen design contains rectangular slightly inclined screens, which are gyrated at the feed end. The discharge end reciprocates but does not gyrate. This combination of motions stratifies the feed, so that fine particles travel downward on the screen surface, where they are pushed through the opening on the screen by the larger particles moving on top. The screen surface is often double, as shown in Figure 9.5. Between the two screen surfaces rubber balls are positioned and held in separate compartments. As the screen operates, the rubber balls strike the screen surfaces, freeing the opening of any material that would tend to plug them.

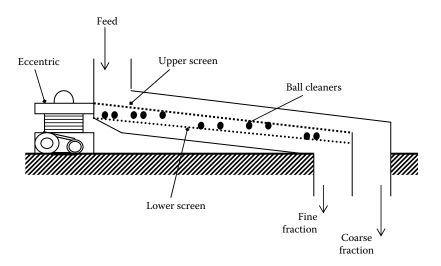


FIGURE 9.5 Horizontally gyrated reciprocating screen.

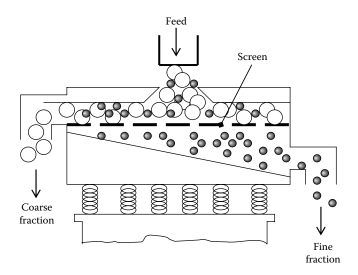


FIGURE 9.6 Diagram of oscillating round screen.

Oscillating screens usually operate at gentle speeds, between 300 and 400 oscillations/min, and with a long linear stroke. The motion of the screen is of an orbital nature, practically on the plane to the screen deck, and may be transmitted by high-velocity eccentrics to the sieves. Vibration is accomplished by eccentric weights on the upper and lower ends of the motiongenerator shaft. Rotation of the top weight creates vibration in the horizontal plane, which causes the material to move across the screen cloth to the periphery. The lower weight acts to tilt the machine, causing vibration in the vertical and tangential planes. The angle of lead given the lower weight with relation to the upper weight provides variable control of the spiral screening pattern. Speed and spiral pattern of material travel over the screen cloth can be set by the operator for maximum throughput. Oscillating screen units are box-like machines, either round or square, with a series of screen nested atop one another. The sieves are made of wire mesh or perforated screen decks to separate relatively dense, dry, undersized materials, from less dense oversized materials. A schematic diagram of an oscillating round screen is given in Figure 9.6.

Reels or trommels are revolving cylindrical screens mounted almost horizontally, with one extreme opened. Their screening surface may consist of wire mesh or perforated sheet. Hexagonal cross sections are also used since these lead to agitation which aids the separation of fine material. Figure 9.7 shows a simple design or revolving screen. The feed material, which travels through the openings increasing in size along the drum, is discharged by a chute on the lower extreme of the equipment. Several arrangements of

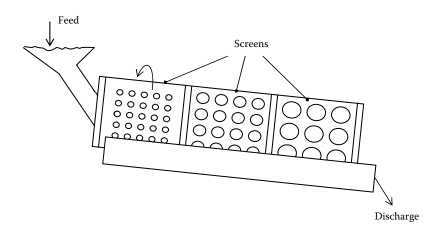


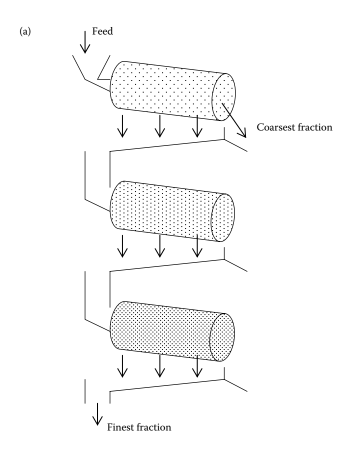
FIGURE 9.7 Single drum revolving screen.

trommels can be made to accomplish separation according to specific process requirements. Figure 9.8 presents two of them. The simplest case is when the drum perforations are uniform all over the surface. In such a case, to separate different fractions of a given material, the drums are mounted in series. Two alternatives may arise, in that the drum with the larger perforations is located as the first or the last of the series. When the largest perforations drum is the first of the series, the most convenient arrangement is to place the drums on top of the other (Figure 9.8a), so as to send the finest fraction of material to the following screen and successively. On the other hand, if the largest perforations drum is the last of the series the coarsest fraction advance to the next drum, and a more convenient arrangement will be to have them mounted one right after the other as a series-consecutive drum screen. Some other arrangements include the concentric drum arrangement illustrated in Figure 9.8b. This particular mounting has the advantage of compactness but, since the feed enters at the center, the smallest screen area will handle the complete feed load while the smaller fines fractions will be progressively processed into the larger screen areas.

The capacity of a trommel increases with increasing speed of rotation until a critical speed is achieved. At speeds greater than this, the material does not cascade over the surface but is carried round and centrifugal force and separation is seriously impaired. The critical speed of a trommel is given by

$$N = \frac{42.3}{(D)^{1/2}} \tag{9.9}$$

where N is the number of revolutions of the trommel per minute, and D is the diameter of the trommel in meters.



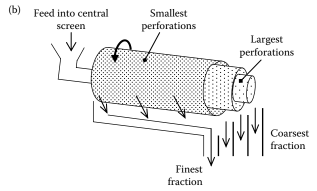


FIGURE 9.8 Arrangements for revolving screens: (a) parallel type and (b) concentric drum.

9.2 Electromagnetic Separation

9.2.1 Basic Principles

A particle of a given material exposed to a magnetic field will become magnetized to some extent and may act as a magnetic dipole. When plotting the intensity of the applied magnetic field *H* and the magnetization induced in the particle M, the resulting graphs represent the three ways in which magnetized materials may behave (Kolm, 1975). The materials thus classified, as illustrated in Figure 9.9, are: ferromagnetic, paramagnetic (magnetically attracted), and diamagnetic. The relationship M/H is known as the susceptibility. Paramagnetic substances have positive susceptibilities, and induced magnetization increases the magnetic-flux density within the substance. Diamagnetic materials have negative susceptibilities, and an induced field in this case cancels part of the magnetic-field intensity. Permeability μ , which is often used somewhat imprecisely in referring to ferromagnetic substances, is the ratio of the magnetic-flux density to the magnetic-field density. Magnetization of different materials is dependent directly on the degree of magnetic susceptibility and the applied magnetic field intensity. As can be seen in Figure 9.9, ferromagnetic materials quickly become magnetically saturated so

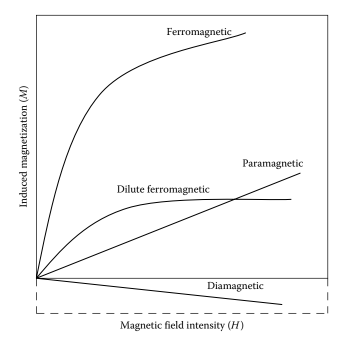
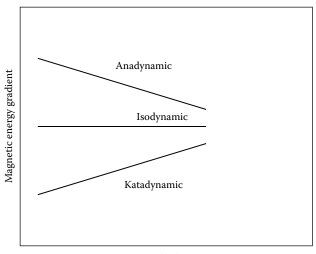


FIGURE 9.9 Magnetic behavior of materials.

that an increase in the intensity of the magnetic field bears no further effect after a certain point. Paramagnetic materials, on the other hand, follow a direct proportion for the magnetic-flux density and the magnetic-field intensity. Some of these substances cannot be, virtually, saturated.

Three types of magnetic fields can be identified: isodynamic, katadynamic, and anadynamic. As can be observed in Figure 9.10, in an isodynamic magnetic region the gradient of magnetic energy does not vary with an increase in the magnetic-field intensity. This type of magnetization is useful to concentrate particles according to slight differences in susceptibility. A katadynamic region is characterized by an increase in the gradient of magnetic energy directly proportional to the magnetic-field intensity. In this region the higher gradients of magnetic energy, along with the higher intensity of the magnetic field, may be applied to a particle. The region of anadynamic field, in which the energy gradient is inversely proportional to the magnetic-field intensity, is particularly useful to separate particles of low paramagnetic susceptibility (Sun, 1976).

Magnetic separation processes work by establishing a specific equilibrium among magnetic forces, particle–particle forces, and inertial forces acting on the particle. All these forces compete in a given operation, and so the role of a design exercise consists on allowing interaction to act in such a way that separation of particles may become feasible. Feed on a magnetic separator can be divided into two or more streams. The "tailings" are the magnetic product, the "mags" are the magnetic concentrate, and the "middlings" are a product lower in magnetization that can also be obtained.



Magnetic filed intensity

FIGURE 9.10 Types of magnetic fields.

9.2.2 Equipment and Applications

Magnetic separators may be classified into conventional separators and high-gradient separators. Table 9.4 lists the main types of equipment available to perform electromagnetic separations. Conventional processes are based on concentrating the magnetic material while the desired product may be either part: the magnetic or the nonmagnetic. Typical applications of conventional separators include removal of small iron pieces (wire, rivets, diverse iron fractions) from varied materials and mineral beneficiation of highly magnetic minerals. Some applications of conventional magnetic separators are given in Table 9.5. There are three main designs of conventional magnetic separators: the grate separator, the drum separator, and the belt separator. The drum separator, whose schematic diagram is shown in Figure 9.11, is probably the equipment most widely used.

High-gradient magnetic separation may be considered an alternative to flotation and extraction for treatment of fine solids suspended in water (Gaudin, 1974), which would categorize it as a solid–liquid separation technique. There are, however, some applications for dry separations. A widely used high-gradient magnetic separator is the "Ferrofilter Franz." A schematic diagram of this type of equipment is given in Figure 9.12. The equipment comprises a solenoid magnet, a magnetic return circuit, a stainless-steel filtration ribbon, and a large voidage magnetic matrix. The unit is operated discontinuously and the feed flow is parallel to the magnetic field. Although

TABLE 9.4Electromagnetic Common Separators and Their Main Characteristics

		Approximate Value of Effective Magnetic Parameters			
Туре	Involved Forces	Strength Range (cm)	Field	Force ^a (dyn/cm ²)	
			Gradient (Gauss/cm)	Fe ₃ O ₄	CuO
Grate	Gravity	1	500	3×10^{5}	5
Drum	Gravity, hydrodynamic drag	5	500	3×10^5	5
Belt	Gravity	1–10	100-1000	550×10^4	0.2 - 20
Davis tube	Gravity, hydrodynamic drag	1	4000	2×10^6	3×10^2
Frantz: isodynamci	Gravity	1	2000	1×10^6	4×10^2
Frantz: ferrofilter	Hydrodynamic drag	0.1	1×10^5	1×10^8	2×10^4
Jones	Hydrodynamic drag	0.1	2×10^5	1×10^8	8×10^4
Kolm- Marston	Hydrodynamic drag	0.001	2×10^6	1×10^9	8×10^5

^a For a particle whose diameter is equal or smaller than the range of magnetic force.

TABLE 9.5

Typical Applications of Magnetic Separators

Iron Pieces Removal

Food processing

- Manufacture of Chemical products
- Mineral beneficiation
- Manufacture of pharmaceuticals
- Cooling fluids treatment

Mineral Beneficiation

Iron recovery

- From solid waste
- From heavy media

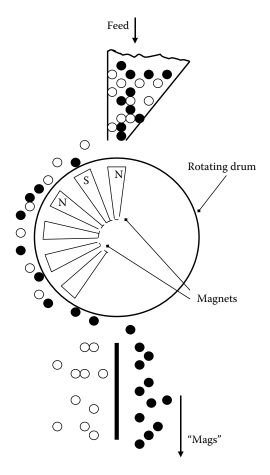


FIGURE 9.11 Conventional drum magnetic separator.

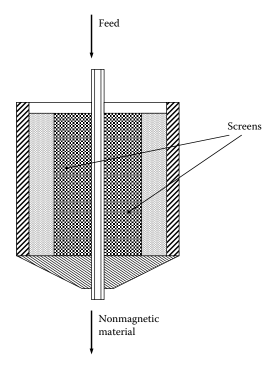


FIGURE 9.12 High-gradient magnetic separator.

this equipment is exclusively used in the kaolin industry for slurry separations, several designs based on it have been developed for dry separation applications. The high-gradient magnetic dry separators can be fed by gravity or within an airstream, and they can handle materials as fine as 325-mesh in size. An example of application of this type of equipment is the separation of paramagnetic impurities from finely pulverized dry minerals (Iannicelli, 1976).

Some other types of high-gradient magnetic separators include the carrousel continuous separator and the superconductive separator. The carrousel-type separator consists on a wheel containing cells where iron fiber matrices are fixed. As the wheel turns, the cells pass continuously near a highly magnetized field region where feed is introduced. The magnetic particles are trapped inside the matrices, and when the cell goes far from the magnetized region the particles are removed by washing. The superconductive separator is based on the low electric resistance showed by some materials cooled at certain temperature. These materials are known as superconductors, and their resistance may be reestablished by applying a specific electrical current or by placing the material in a magnetic field with intensity over a critical value. The magnetic field can, thus, act as a solenoid

for the superconductive material, without elevating the temperature more than a Joule (Watson, 1979).

9.2.3 Selection Criteria

High-gradient magnetic separations are very sensitive to process variables. When a paramagnetic particle is trapped by a ferromagnetic wire and exposed to a magnetic-field intensity H, it is interesting to know the relation among the particle radius b, the wire radius a, and the force exerted by the wire on the particle. Theoretical calculations suggest that the magnetic strength depends critically on the radii of particle and wire, as shown in Figure 9.13 (Kolm, 1975). The attractive force between the particle and the wire reaches a maximum when the relation a/b attains a value of 2.69. A system in which particle and wire radii are adjusted to a maximum magnetic strength value is known as a fixed system, and is effective only on certain ranges of particle size.

Kolm (1975) has studied the effect of increasing particle radius for the case of cupric oxide exposed to magnetic fields, and some data are reported as a plot in Figure 9.14. The Figure shows the plot of the relation of the magnetic force to the added opposite (gravity and drag) forces against particle radius. Clearly, a particle within the size range of $100~\mu m$ is the most appropriate for

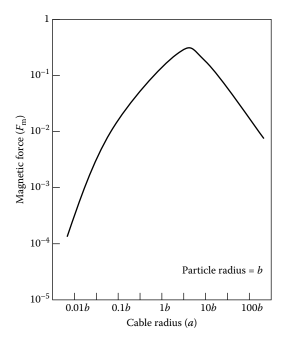


FIGURE 9.13Plot of magnetic force versus wire diameter of a ferromagnetic wire.

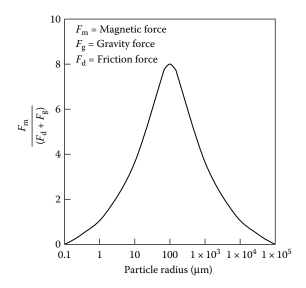


FIGURE 9.14 Plot of fixed system for high-gradient magnetic separation of CuO.

the fitted system and separation of particles far from that size range will become difficult.

9.3 Electrostatic Separation

9.3.1 Basic Principles

Electrostatics may be defined as the study of stationary electrical charges, under conditions in which positive and negative parts are separated from each other. Electrostatic separation is accomplished by use of forces acting on charged or polarized bodies in an electrical field. Several methods can be used for charging, and include conductive induction, contact electrification, and ion bombardment. Conductive induction and ion bombardment are used to separate conductive materials from nonconductive materials. Examples of conductive materials include ilmenite, rutilium, galena, pyrite, and specularite. Among nonconductive materials quartz, zirconium, monazite, and diamonds can be mentioned. Contact electrification is usually employed to separate nonconductive materials, such as phosphoric rock and quartz, based on their degree of conductivity.

Conductive induction principle is illustrated in Figure 9.15. A charged body moves toward an isolated metal object B. Since similar charges reject while different charges attract, negative charges on object A induce electrons

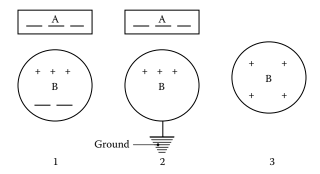


FIGURE 9.15
Mechanism of conductive induction.

on object B so that they move to the other side of B, leaving a net positive charge on the side of object B closest to object A. If object B is grounded, nonbond electrons flow out of earth leaving a positive net charge on the place shown. If object B is taken away, the positive charge distributes uniformly over the surface of the metallic object. In contact electrification a substance is rubbed against the other so that an excess of negative charges is fixed to one of the surfaces. A positive charge is developed on the body deficient in electrons, while the body attracting electrons results negatively charged. Table 9.6 shows the effects on rubbing different materials against carbon and pyrite (Anderson et al., 1979). The method of ion bombardment is used to charge materials of dissimilar electrical properties. Conductive and nonconductive particles get in touch with a grounded conducting surface. Both particles are bombarded by ions of atmospheric gases generated by an electrical corona discharge from a high-voltage electrode. When ion bombardment ceases, the conductor particle loses its acquired charge to ground very rapidly and experiences an opposite electrostatic force tending

TABLE 9.6Electric Charges Developed in Carbon and Pyrite Particles Sliding Inside Pipes

Pipe Material	Charge Magnitude (10 ⁻⁸) C/g		
	Carbon	Pyrite	
Polyvinyl chloride	(+) 2.00	(+) 1.40	
Galvanized iron	(+) 0.40	(-) 0.47	
Vinyl pipe	(+) 0.22	(-) 0.09	
Copper	(+) 2.00	(-) 0.35	
Plexiglass	(+) 2.20	(-) 2.30	
Aluminum	(+) 0.50	(-) 0.31	

to repel it from the conducting surface. The nonconducting particle side kept away from the conducting surface becomes coated with ions of charge opposite in electrical polarity to that of the surface, and it experiences an electrostatic force tending to hold it to the surface. If the electrostatic force is larger than the force of gravity or other forces tending to separate the nonconducting particle from the conducting surface, the particle is held in contact with the surface and is said to be "pinned."

Particles can also be charged by comminution. It has been demonstrated that, in the millimeter range, charging by comminution does not produce particles with sufficient charge as to be influenced by an electrical field (Lawver, 1969). Within the micrometer range, however, it has been found that the resulting charge is high enough for the particles to show an electrical response, rather than a gravitational one, in an electric field. For homogeneous substances, comminution will produce particles somewhat balanced in terms of electrical charges, so that electrostatic separation would not be a feasible option. Particles differing in chemical composition are satisfactorily charged by comminution, and electrostatic separation should give appropriate results.

9.3.2 Equipment and Applications

Table 9.7 gives a list of industrial applications of electrostatic separation (Lawver, 1969). The different types of equipment make use of either direct or alternating current sources in the range of $3 \times 10^3 - 3 \times 10^6$ V/m. The high-tension sources used in commercial electrostatic separators cover a range as high as 250×10^3 V, but the power used is really lower, with a maximum value of 0.5 kW/h per ton of feed. Direct current is used to produce charge by conductive induction and ion bombardment, while alternating current is used to produce charge by polarization.

TABLE 9.7Typical Industrial Applications of Electrostatic Separators

Minerals Combination	Surface Treatment	Usual Charge Mechanism
Hematite-quartz	Drying	Ion bombardment
Limenite-rutile	Washing and drying	Ion bombardment
Zirconium-ilmenite	Drying	Ion bombardment
Cassiterite-scheelite	Drying	Ion bombardment
		Conductive induction
Feldspar-quartz	Drying	Contact electrification
Halite-silvanite	Heating to 340°C	Contact electrification
Pyrite-carbon	Drying	Ion bombardment
•		Conductive induction
Carbon-chalkboard	Moisture control	Conductive induction
Diamond-silicon	Washing and drying	Conductive induction

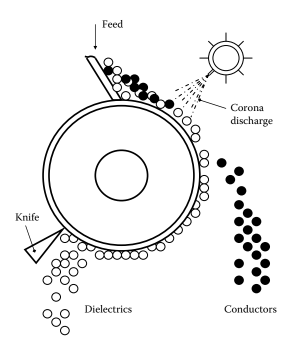


FIGURE 9.16Corona discharge electrostatic separator.

A common equipment making use of ion bombardment mechanism of charging particles is the one known as corona discharger. Figure 9.16 shows a diagram of a corona discharge unit, simply known as high-tension separator (Gaudin, 1971). As particles enter to the high-tension machine and pass through the corona they are charged. The amount of charge is function of the particle radius, the relaxing time, the particle permeability, and the residence time. Conductive particles share their charge quickly with the rotating drum before being projected away from it. Nonconductive particles are attracted to the drum surface and are removed by a scraper or knife. In this manner, a defined separation between conductive and nonconductive materials is carried out. Corona discharge units can handle up to 136 kg of material/12.7 mm of rotor, with particles 0.2–1.0 mm in diameter. As particle size increases, capacity decreases significantly.

In corona discharge units a criterion to discriminate conductors from nonconductors is the relaxing time T_r , defined as the time required for a unit of initial charge density P_0 to decrease in a 1/e proportion from its original value, that is,

$$P = P_0 e^{-\sigma t/\varepsilon} \tag{9.10}$$

where P is the charge density at time t, σ the conductivity, and ϵ the permeability.

Relaxing time for pyrite is approximately 10⁻³ s, while that for quartz is about 10⁶ s. Quartz is, therefore, considered a nonconductor due to its long relaxing time, while pyrite is considered a conductor because of its short relaxing time.

Apart from the corona discharge unit, two common types of electrostatic separators are the plate separator and the screen–plate separator, illustrated in Figure 9.17. Induction is used in both units to charge the feed

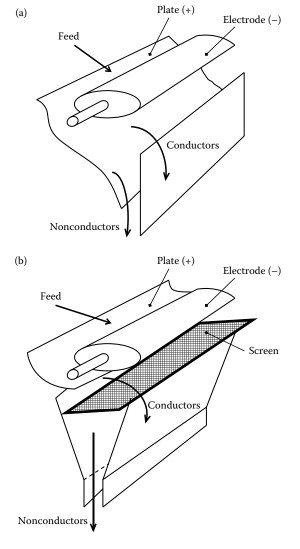


FIGURE 9.17 Electrostatic separators: (a) plate type and (b) screen–plate type.

particles. Efficiency of separation is lower for plate and screen–plate separators than that for corona discharge separators. For this reason, these types of units are mounted in series, with as much as five separation stages (Anonymous, 1976).

9.3.3 Applications in Fine Particulate Systems

In order to attain satisfactory results when using electrostatic separations for finely pulverized materials, some alternatives have been suggested. It has been reported a fluidized bed design specially modified to avoid drag of particles within the gas stream (Kiewlet et al., 1978). It was found that a combination of low gas flow and mechanical vibration may fluidize efficiently a bed of fine particles minimizing the drag problem. Results for a synthetic mixture of glass and iron powders, sized between 15 and 55 μm , are given. Effects of temperature, moisture, and voltage were investigated for a mixture 50/50, by weight, of the mentioned materials. Three important conclusions were found: vibration along with fluidizing action may extend the use of electrostatic separation to mixtures of fine powders difficult to fluidize by gas stream only; vibration does not minimize the extracting effect, since recovering efficiencies well over 60% were observed for the steel component; temperature and voltage are the most determinant variables of the operation.

Another method to separate fine particles by electrostatic separation is the case of a dilute phase closed electrostatic circle (Inculet et al., 1977), developed to eliminate the drag normally associated when handling particles below 40 μ m in size. Due to the electric field influence, particles are deflected from the main stream trajectory and are extracted by central collectors. Particles are suspended in a gas stream, introduced to the loop by gas inductors consisting of two concentric rotating cylinders. The corona charging sections include high-voltage corona wires, while the electrostatic separation section is located within a segment of the loop 1.2 m long, in which the high-voltage electrodes produce a horizontal electric field. The feed system consists on a fluidized bed mounted laterally to the continuous loop.

9.4 Laboratory Exercise: Efficiency of Separation on Single-Stage Screening

9.4.1 Introduction

Screening is a separation of materials on the basis of size as a means of preparing a product for subsequent operation. Screening is performed

using equipment with one or two sieving surfaces. A single sieve provides two products. The product from the top is called the oversized or plus product while the one which passes through the screen is the undersized or minus product. Material passing one sieve surface but remaining on the next one in a double deck screen is an intermediate product. During ideal screening, the plus product contains only particles greater than the size of openings of the screen surface while minus product consists of particles smaller than the opening. Efficiency of screening, measured as the quantity of misplaced material in any of the streams, can be affected by many factors. Clogging is a serious problem that may have a tremendous impact in efficiency. Apart from particle size and shape, that may directly cause clogging, some other characteristics of the feed material, such as its moisture content, may also cause clogging diminishing efficiency of the separation process. The objective of the experiment is to evaluate the effect of moisture content of the material fed to a screening process, in clogging and efficiency of the separation.

9.4.2 Equipment and Materials

US standard test sieves series, or laboratory size oscillating round screen unit. Two batches of 5–10 kg of coarse sand, and fine sand.

9.4.3 Instruments or Apparatuses

Granatary scale and moisture balance.

9.4.4 Screening Procedure

- Mix thoroughly two equal parts of coarse and fine sand. If using standard test sieves, combine 100 g of coarse sand with 100 g of fine sand. If using screen equipment, prepare a sample according to the size of the unit. Separate a sample to determine the moisture content using the moisture balance.
- Use, for any of the cases above, a 0.355 mm sieve (45-mesh), and establish it as the cut diameter $D_{\rm pc}$. Operate the testing setup for the recommended time accordingly.
- Collect the underflow and overflow samples, weigh them, and analyze them for their particle size distributions using the set of sieves required and following the procedure given in Section 2.7. (Laboratory exercise: particle size analysis by different methods.)
- Collect data and elaborate table and plots, like those derived for Examples 9.1 and 9.2.

- Prepare a second sample to run another separation experiment.
 Moisten it slightly with an atomizer and separate a portion to determine the moisture content.
- Repeat the experiment for the dry sample.
- Prepare a third sample to run another separation experiment. Moisten it a bit more than the previous one with an atomizer and separate a portion to determine the moisture content.
- Repeat the experiment for the first moisten sample.

9.4.5 Calculations and Report

- Determine the overall separation efficiency for the three experiments run above, following procedure described in Example 9.2.
- Analyze the results and find out whether a correlation between separation efficiency and moisture content can be established.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out, as well as discussion on own data. Compare and discuss, as much as possible, agreements or discrepancies based on literature survey from studies and investigations on the subject.

TABLE 9.8 Problem 1

	Cun	Cumulative Fraction		
Mesh	Feed	Coarse	Fine	
4	0.0000	0.00	0.00	
6	0.0251	0.08	0.00	
8	0.1501	0.52	0.00	
10	0.4708	0.91	0.20	
14	0.7278	0.96	0.60	
20	0.8868	0.99	0.85	
28	0.9406	1.00	0.93	
35	0.9616	_	0.97	
48	0.9718	_	1.00	
65	0.9795	_	_	
100	0.9853	_	_	
150	0.9894	_	_	
200	0.9925	_	_	

TABLE 9.9 Problem 2

Mesh	Cumulative Fraction			
	Feed	Coarse	Fine	
20	0.75	2.00	0.05	
40	50.80	76.25	5.05	
60	86.92	95.95	41.65	
80	96.32	98.50	79.75	
100	97.96	99.25	88.25	
120	98.96	99.80	96.50	
200			99.55	
Pan	100.0	100.0	100.0	

PROBLEMS

- 1. A mixture of crushed stone with the particle size distribution given in Table 9.8 is sieved through a 10-mesh screen (Tyler). Calculate the overall efficiency of the operation.
- 2. Determine the efficiency of a screening operation of a catalyst treated to increase reaction kinetics of a specific chemical process. Data from screening of the streams resulting from the operation are presented in Table 9.9. The separation operation was carried out using a 0.425 mm sieve.
- 3. A blend of crystals is to be separated intro three fractions: a coarse one retained on a 8-mesh sieve, an intermediate one retained on a 14-mesh sieve, and a fine one passing through the 14-mesh sieve. The results from the screen analyses are listed in Table 9.10. Calculate the efficiency for each sieve in the fractionation of the crystals.

TABLE 9.10 Problem 3

		Differential Analysis (Fraction)			
Mesh	Feed	Coarse	Intermediate	Fine	
3	0.035	0.14			
4	0.150	0.50	0.042		
6	0.275	0.24	0.358		
8	0.235	0.08	0.308	0.200	
19	0.160	0.04	0.183	0.267	
14	0.091		0.102	0.202	
20	0.034		0.007	0.196	
28	0.013			0.089	
35	0.007			0.046	
48	0.000				

References

- Anderson, J. M., Parobek, L., Bergougnou, M. A., and Inculet, I. I. 1979. Electrostatic separation of coal macerals. *IEEE Transactions on Industry Applications* IA-15: 291–293.
- Anonymous. 1976. Electrostatic separation system. *Australian Mining* (December 1976): 27–29.
- Beddow, J. K. 1980. *Particulate Science and Technology*. New York, NY: Chemical Publishing Company.
- Foust, A. S., Wenzel, L. A., Clump, C. W., Maus, L., and Anderson, L. B. 1960. *Principles of Unit Operations*. New York, NY: John Wiley & Sons.
- Gaudin, A. M. 1974. Progress in magnetic separation using high intensity, high gradient separators. *Mining Congress International* (Jan, 1974): 18–21.
- Gaudin, A. M. 1971. Principles of electrical processing. *Minerals Science and Engineering* (April 1971): 46–57.
- Iannicelli, J. 1976. New developments in magnetic separation. *IEEE Transactions on Magnetics* 12: 436–443.
- Inculet, I. I., Bergougnou, M. A., and Brown, J. D. 1977. Electrostatic separation of particles below 40 μμ in a dilute phase continuous loop. *IEEE Transactions on Industry Applications* IA-13: 370–373.
- Kiewlet, C. W., Bergougnou, M. A., Brown, J. D., and Inculet, I. I. 1978. Electrostatic separation of fine particles in vibrated fluidized beds. *IEEE Transactions on Industry Applications* IA-14: 526–530.
- Kolm, H. H. 1975. Tackling the sixth state. *The Physics Teacher* 13: 73–79.
- Lawver, J. E. 1969. State of the art of electrostatic separation of minerals. *Journal of the Electrochemical Society* 116: 57C-60C.
- Nakajima, Y. N., Whiten, W. J., and Withe, M. R. 1978. Method for measurement of particle size distribution by sieves. *Transactions of the Institute of Mining and Metallurgy* 87: C194–203.
- Strumpf, D. M. 1986. Selected particle size determination techniques. *Manufacturing Confectioner* 66: 111–114.
- Sun, J. J. 1976. Magnetic barrier—A new concept in magnetic separation. *IEEE Transactions on Magnetics* 12: 483–485.
- Watson, J. H. P. 1979. Aplications and improvements in high gradient magnetic separation. *Filtration and Separation* 16: 70–72, 74–75, 78.

10

Solid-Fluid Systems

10.1 Introduction: Simultaneous Flow of Fluids and Solids

Practically, in most operations relevant to process engineering numerous fluids are handled. Fluids of all sorts constitute raw materials and finished products in industries as diverse as the chemical industry, the biotechnology industry, the mining industry, and the food industry. Therefore, the study of fluid mechanics is a fundamental support for practicing engineers in the materials processing industry. Consequently, the subject is relevant in programs of study of chemical engineering, mechanical engineering, biosystems engineering, and related engineering disciplines. Not all the fluids handled in the mentioned industries are pure fluids; as previously discussed, suspensions, emulsions, dispersions, and many other two-phase and multiphase systems are also encountered in a number of industrial processes. Finely divided solids in a continuous liquid phase represent, for example, common suspensions and slurries in diverse operations that need to be handled and processed in different manners. A deep understanding of two-phase and multiphase systems is not normally covered in basic courses of study of fluid mechanics and, yet, many flow problems that a process engineer will face in day-to-day duties involve some type of nonpure fluid. Slurries and suspensions may behave as non-Newtonian fluids even at lower solids concentrations. Their study may be considered a specific topic of fluid mechanics known as flow through porous media or flow past immersed bodies. It may also be considered a specific topic of rheology or of particle technology. As titled in this chapter, simultaneous flow of fluids and solids presents an insight of the main characteristics of suspensions behavior. Particle-fluid interactions and rheology of suspensions are of utmost importance for an understanding of the phenomena governing relevant solid-fluid separations in materials processing industries.

10.1.1 Classification of Fluids

According to rheology, which embraces the study of flow behavior in very general way, there are two main types of flow: viscous and elastic. The first

occurs in fluids, whereas the second is common to solids. An intermediate behavior of flow in between is also found. Thus, perhaps the most useful classification that, in terms of flow behavior, can be made is by means of a spectrum extending from elastic solids on one extreme to viscous flow on the other. Because the laws governing solids flow differ totally from those describing fluids flow, viscous and viscoelastic flow are normally studied together. Table 10.1 presents a summary of the main types of fluids.

Newtonian theory provides the simplest cases of viscous behavior, in which the stresses are related to the velocity gradients existing at time of the observation. For an incompressible fluid in a simple shear, such theory may be stated, according to Figure 10.1, by the first function in Table 10.1. In the most general sense, any fluid response not explainable by Newtonian theory may be termed non-Newtonian. In practice, however, non-Newtonian fluids, as opposed to Newtonian, are those in which the viscosity is not a constant value, but is a function of the imposed shear rate. The dispersions resultant of dissolving solids in liquids, are normally of non-Newtonian characteristics. In the same manner, the suspensions of finely divided solids into continuous liquid phases are usually non-Newtonian.

A wide variety of nonlinear relationships between stress and shear rate have been used. Possibly, the most common one known as the power-law, also appearing in Table 10.1, representing the shear thickening and shear thinning fluids as described there. If the exponent n is less than one, the material flows more easily the faster it is sheared, and the apparent viscosity

TABLE 10.1Classification of Fluids

Fluid	Characteristic Function	Examples
I. Newtonian	$\tau = \mu \dot{\gamma}$	Air, water, steam, all gases, milk, vegetable oil, honey
II. Non-Newtonian		
1. Time independent		
(a) Bingham plastic	$\tau = \tau_{\rm o} + \mu_{\rm o} \; \dot{\gamma}$	Toothpaste, peanut butter, butter, potter's clay, mustard, mayonnaise
(b) Shear thickening (dilatant)	$ \tau = K \dot{\gamma}^n \\ n > 1 $	Quick sand, thick starch solutions, wet beach sand, fine powders in suspension
(c) Shear thinning (pseudo plastic)	$ \tau = K \dot{\gamma}^n \\ n < 1 $	Paper pulp, paint, apple sauce, banana puree, orange juice concentrate
2. Time dependent nonelastic		
(a) Rheopectic	No unique	Bentonite clay
(b) Thixotropic	No unique	Paints, printing inks, tomato ketchup, oil drilling mud
3. Viscoelastics	No unique	Saliva, nearly all-biological fluids, concentrated tomato soup, bread dough, many polymeric solutions

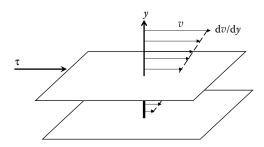


FIGURE 10.1 Definition of viscosity: τ = shear stress; v = fluid velocity; $dv/dy = \dot{\gamma}$ (shear rate).

decreases with increasing shear rate. If, on the other hand the exponent n is greater than one, the apparent viscosity increases with the increasing shear rate. Because they follow the power law, dilatant and pseudo-plastic fluids are also known as power-law fluids. Another case of flow behavior can be considered when a solid like material does not flow initially at all (or hardly at all), until some critical stress have been reached, and then it flows in a Newtonian manner. Such a behavior characterizes the so-called Bingham plastic fluids, as can be described by the second equation on Table 10.1. Graphically, all the above-mentioned models can be represented as illustrated in Figure 10.2.

There is a further approach to describe the flow behavior of systems in which the viscosity varies with time as well as with shear rate. A material

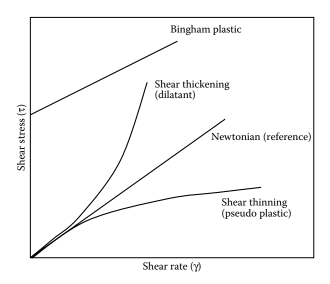


FIGURE 10.2 Rheograms for different types of fluids.

whose viscosity thins out with time is said to be thixotropic, while one whose viscosity thickens with time is said to be rheopectic. Finally, viscoelastic behavior refers to wide variety of effects that are somewhere between solid and fluid responses. With viscoelastics, the τ versus $\dot{\gamma}$ diagram only covers part of the whole behavior, and so transient experiments are needed to characterize their elastic properties.

By far, the most understood behaviors are those of Newtonian fluids and time-independent non-Newtonian suspensions. A series of flow equations and charts have been developed in order to predict their flow characteristics. For other types of non-Newtonians the flow equations, if they can be developed at all, are much more complicated. However, under certain assumptions, for example, steady-state flow without acceleration (flow in straight pipes without nozzles, bends, orifices, etc.), these fluids can often be treated as time independent too.

All the above-mentioned models have been usually developed for the nonturbulent region. There is a dearth of information about non-Newtonian behavior at the turbulent level, mainly perhaps due to the fact that most of the practical applications (pumping, flow and mixing of slurries, etc.) lie in laminar flow regime.

10.1.2 Dynamics of Particles Submerged in Fluids

If a particle moves relative to the fluid in which it is suspended, the force acting on the particle is the gravity force $F_{\rm g}$, while the forces opposing the motion are the buoyant force $F_{\rm b}$ and the drag force $F_{\rm D}$. The resultant of forces acting on the particle may be represented by

$$\Sigma F = F_{\rm g} - F_{\rm b} - F_{\rm D} \tag{10.1}$$

The external gravity force is defined as

$$F_{\rm g} = mg \tag{10.2}$$

where m is the mass of the particle and g the acceleration due to gravity.

The buoyancy force equals the mass of the displaced fluid due to the acceleration of the particle, based on the Archimedes principle, that is,

$$F_{\rm b} = \frac{m\rho \, g}{\rho_{\rm s}} = V_{\rm s} \rho \, g \tag{10.3}$$

where ρ is the fluid density, ρ_s is the particle density, and V_s is the volume of the particle.

The drag force is conventionally expressed as

$$F_{\rm D} = C_{\rm D} \left[\frac{u^2}{2} \right] \rho A_{\rm p} \tag{10.4}$$

where u is the particle–fluid relative velocity, ρ is the fluid density, A_p is the area of the particle projected in direction of the motion, and C_D is a coefficient of proportionality known as the drag coefficient.

The resultant force in Equation 10.1 equals the acceleration as the particle accelerates on its way down. Since acceleration du/dt is due to the mass of the particle, the resultant force can also be expressed as

$$m\frac{\mathrm{d}u}{\mathrm{d}t} = F_{\mathrm{g}} - F_{\mathrm{b}} - F_{\mathrm{D}} \tag{10.5}$$

Substituting Equations 10.2 through 10.4 into Equation 10.5 and transposing for acceleration the following relation is obtained:

$$\frac{\mathrm{d}u}{\mathrm{d}t} = g \left[\frac{\rho_{\mathrm{s}} - \rho}{\rho_{\mathrm{s}}} \right] - \frac{C_{\mathrm{D}}u^{2}\rho A_{\mathrm{p}}}{2m}$$
 (10.6)

In settling under influence of gravity g is constant while the drag always increases with velocity. Equation 10.6 shows that the acceleration decreases with time and approaches zero. The particle quickly reaches constant velocity, which is the maximum attainable, and which is called the terminal velocity. The equation for terminal velocity u_t is found, for settling under gravity, by taking du/dt = 0 and transposing for the particle–fluid relative velocity from Equation 10.6, so as to obtain:

$$u_{\rm t} = \left\lceil \frac{2g(\rho_{\rm s} - \rho)m}{A_{\rm p}\rho_{\rm s}C_{\rm D}\rho} \right\rceil^{1/2} \tag{10.7}$$

For spherical particles $m = (\pi x^3 \rho_s)/6$, and $A_p = (\pi x^4)/4$, being x an equivalent diameter. Substituting these values into Equation 10.7:

$$u_{\rm t} = \left\lceil \frac{4(\rho_{\rm s} - \rho)gx}{3C_{\rm D}\rho} \right\rceil^{1/2} \tag{10.8}$$

Assuming that the drag force is due to the inertia of the fluid, C_D would be constant and dimensional analysis shows that C_D is generally a function of the particle Reynolds number Re_p . The form of the function depends on the regime of the flow. This relationship for rigid spherical particles is shown in

Figure 10.3. At low Reynolds number under laminar flow conditions when viscous forces prevail, C_D can be determined theoretically form Navier–Stokes equations and the solution is known as Stokes' law and represented by

$$F_{\rm D} = 3\pi\mu ux \tag{10.9}$$

where μ is the viscosity of the fluid.

This is an approximation, which gives the best results for $\text{Re}_p \to 0$; the upper limit of its validity depends on the error that can be accepted. The usually quoted limit for the Stokes region of $\text{Re}_p = 0.2$ is based on an error of about 2% in the terminal settling velocity. Equations 10.4 and 10.9, combined with the definition of the particle Reynolds number, that is, $\text{Re}_p = (xu\rho)/\mu$, give another form of Stokes' law as follows:

$$C_{\rm D} = \frac{24}{{\rm Re}_{\rm p}} \left({\rm Re}_{\rm p} < 0.2 \right)$$
 (10.10)

shown in Figure 10.3 as a straight line. For $Re_p > 1000$ the flow is fully turbulent with inertial forces prevailing, and C_D becomes constant and equal to 0.44 (the Newton region). The region in between $Re_p = 0.2$ –1000 is known as the transition region, and C_D is either described in a graph or by one or more empirical equations. A generally accepted value for this region can be taken as $C_D = 18/(Re_p)^{0.6}$.

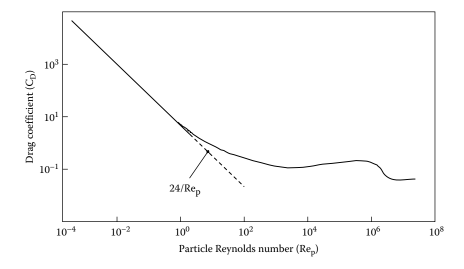


FIGURE 10.3 Drag coefficient versus particle Reynolds number for spherical particles.

Substituting Equation 10.10 into Equation 10.8, for the Stokes region, the terminal settling velocity is represented by

$$u_{\rm t} = \frac{x^2(\rho_{\rm s} - \rho)g}{18\mu} \tag{10.11}$$

For the transition region, substituting $C_D = 18/(\text{Re}_p)^{0.6}$ into Equation 10.8, the solution gives

$$u_{\rm t} = \left[\frac{x^{1.6} (\rho_{\rm s} - \rho) g}{13.875 \mu^{0.6} \rho^{0.4}} \right]^{1/1.4}$$
 (10.12)

Finally, substituting $C_D = 0.44$, for the Newton region the following relation is obtained:

$$u_{\rm t} = 1.74 \left[\frac{x(\rho_{\rm s} - \rho)g}{\rho} \right]^{1/2}$$
 (10.13)

Direct quantitative use of Equation 10.8 is not feasible in practice because numerical values of $C_{\rm D}$ should be required, and $C_{\rm D}$ depends on $u_{\rm t}$. Since particles settle on definite ranges according to Figure 10.3, the selection of any of the Equations 10.11, 10.12, or 10.13 ought to be determined for any given particle size and settling conditions. If the terminal velocity of a particle of known diameter is required, the particle Reynolds number is unknown and a choice of equation cannot be made. For example, if Equation 10.11 is to be used to determine the terminal velocity of a given particle size, Re_p should not exceed the limiting value of 0.2, but the calculated value of $u_{\rm t}$ will be used to verify whether that limiting Re_p value was not exceeded. A method has been suggested (McCabe et al., 2005) to avoid the use of successive approximations when determining terminal settling velocities of particles in a fluid.

In many solid–fluid systems concerned with separation of fine particles, which are all most difficult to separate, the Reynolds numbers are low, often less than 0.2, due to the low values of x and u. Therefore, in many applications in industry, only the Stokes region has to be considered. Variations in particle shape can be accounted for by obtaining separate curves of C_D against Re_p for each shape. Graphs of this sort are presented in classical text and reference books (Green and Perry, 2008; McCabe et al., 2005). The curves for shapes differing greatly from spherical shape, for example, disks and other plate-like particles, apply only to a specified orientation of the particle. In the free motion of nonspherical particles through a fluid the orientation is constantly changing, and such change consumes energy. The effective drag on the particle increases, and C_D is greater than

for the motion of the fluid past a stationary particle. As a result, the terminal velocity is less than would be predicted from curves for particles with a fixed orientation. For very small particles, the effect of the Brownian movement has to be taken into account. Brownian movement is a random motion imparted to the particle by collisions between the particles and the molecules of the surrounding fluid. The effect becomes appreciable at a particle size of approximately $2\,\mu m$ and predominates over the force of gravity with a particle size of 0.1 μm or less. The random movement of the particle tends to suppress the effect of gravity and the settling does not occur.

Example 10.1

Oil drops 20 μ m in diameter are to be settled in air at a temperature of 37.8°C and a pressure of 101.3 kPa. The density of the oil is 900 kg/m³. Calculate the terminal sedimentation velocity of the oil drops.

SOLUTION

The density of the oil drops is very high compared to that of air, and so, they will settle in the same way a solid particle would do. At the given conditions of air, consulting values on Appendix 3, the density and viscosity of air are 1.137 kg/m³, and 1.9×10^{-5} kg/m s, respectively. Supposing that settling would occur in the Stokes region, substituting values into Equation 10.11:

$$u_{\rm t} = \frac{(2 \times 10^{-5})^2 \text{m}^2 (900 - 1.137) \text{ kg/m}^3 (9.81) \text{ m/s}^2}{18(0.000019) \text{ kg/ms}} = 0.0103 \text{ m/s}$$

In order to verify whether the appropriate selling regime was chosen, the particle Reynolds number is determined as

$$Re_{p} = \frac{(2 \times 10^{-5}) m (1.137) \text{ kg/m}^{3} (0.0103) \text{ m/s}}{(0.000019) \text{ kg/ms}} = 0.0123$$

The value of Re_p is below the limiting figure of 0.2, and so the first approach was correct and the terminal velocity of oil drops through air is $u_t = 0.0103$ m/s.

10.2 Separation Efficiency

10.2.1 Evaluation of Efficiency

The imperfection of separation generally observed in any device performing a solid–fluid separation operation, can be characterized evaluating some

kind of separation efficiency. The problem for expressing such efficiency is particularly relevant in some areas of particle technology such as cleaning of gases, solid—liquid separations, and particle classification. In this section, several procedures and criteria to evaluate separation efficiency are discussed. There are some differences according to the specific application, but basic rules and principles are common to a number of solid—fluid separation techniques. Due to the complexity of most of the solid—fluid systems different efficiencies can be defined, such as the total gravimetric efficiency derived from a mass balance, or several partial efficiencies referred to specific components in a mixture. A very useful concept for separators in which no physical barrier exists to retain particles is the grade efficiency related to the cut size or cut point.

10.2.2 Total Gravimetric Efficiency

The first and most obvious definition of separation efficiency is simply the overall mass recovery as a fraction of the feed flow rate. In solid–gas separations the general purpose is to eliminate fine solids from a gas stream, although there is also the case of separating gas trapped in a bed of granular solids. In terms of solid–liquid separations, the efficiency can be related not only to the quantity of solids retained in the concentrated stream, but also to the quantity of solids leaving the clarified stream. In solid classification, a way of evaluating efficiency is by looking at the misplaced material in a specified fraction, that is, fines in coarse fraction or coarse material in fines fraction. All these cases can be generally expressed as a mass balance. According to Figure 10.4, the overall efficiency equivalent to mass recovery or retention of particles is defined as

$$E_{\rm t} = \frac{M_{\rm c}}{M} \tag{10.14}$$

where all the components are as defined in the mentioned figure. If there is no accumulation of solids in the separator, then

$$M = M_{\rm c} + M_{\rm f} \tag{10.15}$$

and there is a choice of three possible combinations of the material streams for the total efficiency testing. It can be shown (Trawinski, 1977) that if all the operating conditions are equal, the most accurate estimation of the local efficiency comes from the two leaving streams.

The total, or overall efficiency E_t , can also be calculated from gravimetric analyses of particle size distributions of the solids in all the streams, because the mass balance may also be referred to any given fraction of particles of the

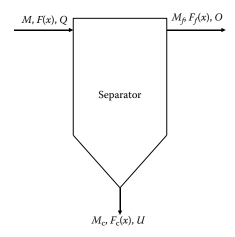


FIGURE 10.4

Diagram of mass balance in a separator: M: mass flow rate of solids in the feed, M_c : mass flow rate of separated solids, M_f : mass flow rate of nonseparated solids, F(x): cumulative percentage oversize of feed solids, $F_c(x)$: cumulative percentage oversize of separated solids, $F_f(x)$: cumulative percentage oversize of nonseparated solids, Q: volumetric flow rate of feed suspension, U: volumetric flow rate of underflow suspension, O: volumetric flow rate of overflow suspension.

total particle size distribution. This type of efficiency can be represented mathematically by

$$dF(x) = E_{t} dF_{c}(x) + (1 - E_{t}) dF_{f}(x)$$
(10.16)

A disadvantage of this way of expressing efficiency is due to the fact that errors in measuring particle size distributions are usually greater than those involved in measuring mass flow rates.

In some solid–liquid separation equipment, such as hydrocyclones, disk self-cleaning centrifuges, or settling tanks, when the underflow keeps significant amount of liquid the total efficiency defined in Equation 10.14 includes a flow partition effect. This effect refers to the fact that, since these devices divide the feed flow into two streams, some solids would follow any of the streams simply by inertia. The ratio of solids partition can be related to the partition coefficient described by the volumetric underflow (U) rate and the volumetric feed rate (F or Q). The relation U/Q is known as underflowsto-throughput ratio R_f . The underflow-to-throughput ratio describes a sort of "guaranteed efficiency" regardless of whether the separation device is operating properly or not. It is sometime desirable to subtract the effect of the solids partition by referring to a "net" efficiency. Numerous definitions for a net efficiency have been reported (Van Ebbenhorst Tengbergen and

Rietema, 1961) but, possibly, the most widely employed is that known as the "reduced efficiency" (Kelsall, 1966) defined by

$$E_t' = \frac{E_t - R_f}{1 - R_f} \tag{10.17}$$

where R_f is the underflow-to-throughput ratio U/Q.

Equation 10.17 satisfies the basic requirements for a definition of efficiency as it takes the extreme values of zero for null separation and unity for complete separation. It is important to note that the reduced efficiency is equivalent to the "clarification number" sometimes employed in liquid filtration (Purchas, 1971).

On occasion, either for mathematical convenience or for interest in the remaining fines in the overflow, a net efficiency of fines E_f is defined as

$$E_{\rm f} = \frac{M_{\rm f}}{M} \tag{10.18}$$

or else, in terms of total efficiency E_t :

$$E_{\rm f} = 1 - E_{\rm t} \tag{10.19}$$

For some applications, in which the fines mentioned represent the product, for example, powder classification or mineral beneficiation, the net efficiency of fines is referred to as the confusing term of "recovery."

An alternative way of expressing efficiency is by means of the degree of separation α , defined as the relation between the total efficiency and the fines efficiency, that is,

$$\alpha = \frac{E_{\rm t}}{1 - E_{\rm t}} \tag{10.20}$$

Some other types of gravimetric efficiency have been reported in the literature. Although some of them would be quite specific, a general knowledge would be important due to the complexity of solid–fluid separation techniques. Svarovsky (1979) has given an exhaustive review of the general topic of separation efficiency in process engineering.

10.2.3 Partial Gravimetric Efficiency

The total efficiency defined by Equation 10.14 includes all particle sizes present in the feed solids. If only a narrow range of particle sizes is of interest,

another efficiency of separation particular to that range can be defined. A mathematical expression of such partial efficiency is

$$E_{\rm p} = \left(\frac{M_{\rm c}}{M}\right)_{x_1/x_2} \tag{10.21}$$

where x_1 and x_2 represent the particle size limits of a definite range.

A typical example of partial efficiency is the so-called "hygienic efficiency" used as a criterion to verify ambient air quality, which includes particles in the range of $0.5–5\,\mu m$ considered harmful for the human respiratory system.

Two usual definitions of partial efficiency are the "rejection efficiency" and the "acceptance efficiency" or recovery index. The rejection efficiency, relevant in classification of powders, is considered as the mass of coarse product greater than the cut size (the limiting size dividing a particle size distribution into fines and coarse fractions) related to the mass of feed solids greater than the same cut size. According to this, in Equation 10.21 x_1 would represent the cut size (x_{50}) and x_2 would be the maximum particle size in the feed. The rejection efficiency E_r can be expressed as a function of cumulative oversize (coarse) percentage F_c and the cumulative percentage of coarse fraction in the feed F_c as

$$E_{\rm r} = E_{\rm t} \left[\frac{F_{\rm c}(x_{50})}{F(x_{50})} \right]$$
 (10.22)

With regard to the acceptance efficiency, it may be considered opposite to the rejection efficiency in the sense that the undersized particles are those considered as the reference for defining such efficiency. In this case, x_1 in Equation 10.21 would represent zero, while x_2 would be taken as the cut point. The acceptance efficiency E_a can be expressed as a function of cumulative undersized (fines) percentage F_f and the cumulative percentage of fines fraction in the feed F_f as

$$F_{\rm a} = (1 - E_{\rm t}) \left[\frac{1 - F_{\rm f}(x_{50})}{1 - F(x_{50})} \right]$$
 (10.23)

A special case of acceptance efficiency is the one known as screen or sieve efficiency E_s , which assumes a perfectly formed screen so that the underflow is free from particles larger than the equivalent screen diameter x_A . According to this criterion, Equation 10.23 can be reexpressed as

$$E_{\rm s} = \frac{F_{\rm c}(x_{\rm A}) - F(x_{\rm A})}{F_{\rm c}(x_{\rm A})[1 - F(x_{\rm A})]}$$
(10.24)

Considering that in agreement with the integrated version of Equation 10.16: $F_f(x_A) = 0$, and $E_t = [F(X_A)]/[F_c(X_A)]$, the screen efficiency can be measured without knowing the total efficiency avoiding, thus, the need to measure the solids flow rate. If the recovery index is determined for any particle size, and not only for the cut size, a curve known as probability cumulative partition curve is obtained. Such curve is used in determining the recovery index in classification of powders.

10.2.4 Grade Efficiency and Cut Size

If the particle size range on Equation 10.21 becomes infinitesimal, the obtained efficiency corresponds to a single particle size x and it is known as the grade efficiency, defined by

$$G(x) = \left(\frac{M_{\rm c}}{M}\right)_x \tag{10.25}$$

The grade efficiency has become a very useful definition, since most industrial powders consist of an infinite number of differently sized particles. Thus, a single particle size really corresponds to a range of particles having almost similar sizes. Therefore, the grade efficiency of most separation equipments is a continuous function of x. This function is seldom expressed analytically but graphically. An S-shaped curve is usually obtained for separators in which inertial or gravity body forces perform the separation.

As the value of the grade efficiency has the character of probability, plotting the probability for any given size fraction against particle size would give a curve as shown in Figure 10.5a. This sort of curve is known as a grade efficiency curve, or Tromp curve, and is derived from screen analysis data on the feed, overflow and underflow streams. It can be shown (Trawinsky, 1977) that the grade efficiency curve is, really, derived from a step-wise calculation drawing a line through the midpoints of size intervals. The derivation of the grade efficiency curve can be carried out graphically using tabulated data, or by computing means. The basic equations for deriving the grade efficiency curve, one for every stream involved in solid–fluid separations, are worked out from the definition in Equation 10.25 and the mass balance shown in Equation 10.16 as follows:

$$G(x) = E_{t} \left[\frac{\mathrm{d}F_{c}(x)}{\mathrm{d}F(x)} \right]$$
 (10.26)

$$G(x) = 1 - (1 - E_{t}) \left[\frac{dF_{t}(x)}{dF(x)} \right]$$
 (10.27)

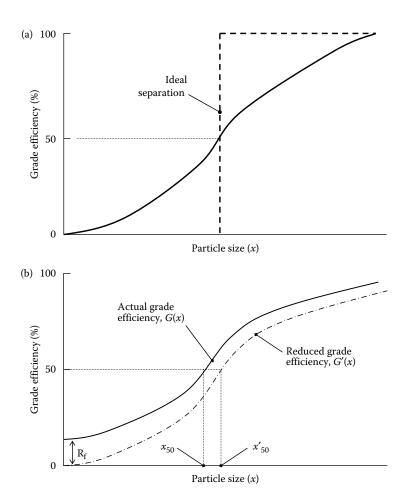


FIGURE 10.5
Grade efficiency curve: (a) without taking into account dead flux effect, and (b) taking into account dead flux effect.

$$\frac{1}{G(x)} = 1 + \left[\frac{1}{E_{t}} - 1\right] \left[\frac{dF_{f}(x)}{dF_{c}(x)}\right]$$
(10.28)

According to Svarovsky (2000) the combination giving best results is that using both leaving streams in a solid–fluid separation process, that is, over-flow and underflow, as represented by Equation 10.28.

Grade efficiency curves derived for practical applications normally do not pass through the origin. This can be explained bearing in mind that, as has been previously described, some separation devices are flow dividers, and so the underflow always contains a certain quantity of very fine particles which simply follow the flow, and are split in the same ratio as the liquid. In order

to account for the partition effect (also called "dead flux") of some solid-fluid separators, the reduced grade efficiency, similar to the reduced efficiency represented by Equation 10.17, can then be obtained as

$$G'(x) = \frac{G(x) - R_{\rm f}}{1 - R_{\rm f}}$$
 (10.29)

As can be seen in Figure 10.5b, the correspondent curve to this definition does cross the origin.

Solid–fluid separators are among those equipments whose separation performance is highly dependent on particle size. Thus, according to this, an overall efficiency of separation is not the best way to evaluate their performance. For this reason, it is advisable to express their efficiency for each size of particle in the range measured. As experience has shown, the only single number, which in some way gives the separation power of a solid–fluid separator, is the cut size. This cut size or cut point (x_c , x_{50}) is the particle size at which the grade efficiency curve shows a value of 50%. In other words, particle with such a size will have a probability of 50% of being separated or, most practically, all particles above the cut size will be generally going to the underflow, whereas those below the cut size will be normally reported to the overflow.

As can be concluded, the cut size is derived from a grade efficiency curve. As the calculation and plotting of a full grade efficiency curve is a time consuming task, alternative ways of determining the cut size have been proposed (Trawinski, 1976, 1977). The main drawback of these methods is the need of plotting limited data, and so the advantageous feature is greatly reduced. On the other hand, depending on specific applications, empirical equations of regression models can be consulted to have a rough estimate of cut size in defined equipment units. Doheim et al. (1985), for example, present a model for prediction of cut size in hydrocyclones using empirical equations.

The sharpness of separation can be related to the shape of the curve in a number of manners. Knowing the grade efficiency curve, comparisons of classification sharpness of different equipments can be made by plotting G(x) against dimensionless size relation x/x_{50} , and comparing the resultant normalized curves. To define the curve steepness, a ratio of two sizes corresponding to two different percentages on the grade efficiency curve on either side, and 50% equidistant, can be used. This parameter is called the sharpness index and is represented by

$$H_{25/75} = \left(\frac{x_{25}}{x_{75}}\right) \tag{10.30}$$

It was previously stated that the grade efficiency curve does not go through the origin. The cut size as has been defined is derived from this curve. If, in order to practically assess the performance of solid–fluid separators, a reduced grade efficiency curve is used, the particle size which gives 50% efficiency in such a curve is called the reduced cut size and is represented by x'_{50} .

The maximum attainable efficiency related to particle size would be that minimum particle size with 100% probability of being reported to the underflow. Graphically, by extrapolating the end part of the curve to the horizontal axis, such size will be obtained. It has been proved that in practice, the maximum of the efficiency is around 98%, and the minimum size corresponding to this efficiency is represented by x_{98} and known as the approximate limit of separation.

As a final note is important to bear in mind that solid–fluid separations are governed by mechanical forces, such as the gravity force, and so it is important that the screen analysis data on the relevant streams used to derive a grade efficiency curve had been measured avoiding transformations. The use of a particle sizer that measures the relevant particle size directly, that is, any method measuring directly the Stokes diameter $x_{\rm st}$, should be preferred over a method relying on transforming size distributions from number to surface, or surface to mass, and so on.

Example 10.2

A hydrocyclone, 44 mm in diameter, is used to separate a suspension of limestone in water. The density of the limestone is 2885 kg/m³ and the gravimetric separation efficiency was 28.34% while the partition coefficient was determined as 11.62%. Derive a grade efficiency curve to find out the reduced cut size, from the data given in Table 10.2.

TABLE 10.2 Example 10.2

Particle Size (µm)	F _c (%)	$F_{\rm f}$ (%)
15	20.06	69.35
20	25.62	79.73
25	31.51	86.81
30	37.60	91.57
35	43.75	94.72
40	49.83	96.77
45	55.73	98.08
50	61.35	98.89
55	66.63	99.38
60	71.50	99.67
65	75.92	99.83
70	79.88	99.92
75	83.38	99.96
80	86.42	99.98

SOLUTION

Table 10.2 presents the particle size distributions for the overflow and underflow through the hydrocyclone. Therefore, Equation 10.28 would be the appropriate to derive the grade efficiency. Since the separation was carried out in a hydrocyclone, and this type of device normally presents a dead flux effect previously described, Equation 10.29 should be used to derive the reduced grade efficiency. Carrying out the proper computations using the tabulated data and the equations mentioned, Table 10.3 is obtained.

Plotting the data in Table 10.3, the following graph (see Figure 10.6) can be drawn.

Finally, taking the reading from the above graph in the 50% point of the G'(x) axis an approximate reduced cut size $x'_{50} = 40 \ \mu m$ is obtained.

10.3 Solid-Gas Separations

10.3.1 Introduction

Two-phase systems of fine dispersed solids in a continuous gas stream are quite common in many processing industries dealing with dry solids operations. In diverse processes such as the dry milling of grains, some combustion reactions, or the grinding of raw materials in the cement industry, fine solids may be released to the surrounding air as part of their normal operation. Tiny solid particles (dust) produced in many processes may become airborne and carried out by air currents. There are many problems associated with dust emissions, such as health hazards, abrasion damage to equipment, impaired

TABLE 10.3 Example 10.2

Particle Size (µm)	G(x) (%)	G'(x) (%)
15	12.92	2.68
20	18.87	9.33
25	26.42	17.77
30	35.44	27.85
35	45.54	39.13
40	56.02	50.85
45	66.09	62.10
50	75.05	72.12
55	82.47	80.41
60	88.23	86.84
65	92.44	91.55
70	95.36	94.82
75	97.29	96.98
80	98.51	98.33

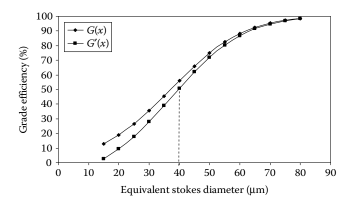


FIGURE 10.6 Example 10.2.

visibility, unpleasant odors, material loss, and risk of dust explosions. Another common situation of two-phase (solid–gas) system is when such system results from a specific operation, like in pneumatic conveying or in spray drying. In all these described cases, it becomes necessary to separate the dispersed or suspended solids from the airstream (or any other gas).

Two types of solids dispersoids causing problems in diverse processes have been recognized: dust consisting on particles larger than 1 μm , and fume formed by particles smaller than 1 μm . Fumes are also defined as particles that, once settled, do not normally return to an airstream, while dusts can be considered particles that even after settling may become airborne again as an effect of an air draft or blast.

A number of solid–gas separation techniques, such as cyclonic separation, gas filtration, or scrubbing, can be used to purify dust-laden gases or recover particulate solids from streams carrying them.

10.3.2 Use of Cyclones

An inert force commonly employed in separating solids particles from a gas stream is the centrifugal force. Cyclones and air classifiers represent techniques that make use centrifugal force as a means of performing separation. Air classification may be considered an analytical technique with some industrial applications, while cyclonic separation constitutes one of the most useful industrial operations to treat dust-laden air and gases in a number of processes and applications.

10.3.2.1 Description of the Process

Cyclones are by far the most common type of gas-solids separation device used in diverse industrial processes. They have no moving parts, are

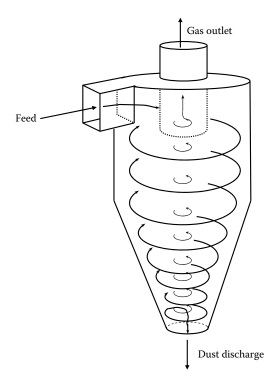


FIGURE 10.7 Schematic diagram of a cyclone.

inexpensive compared to other separation devices, can be used at high temperatures, produce a dry product, have low energy consumption, and are extremely reliable. Cyclone operation obeys fundamental physical laws and there are certain limitations in particle size to be successfully captured in a given unit. However, high-efficiency cyclones can be designed to collect a great proportion of any industrial dust, down to the smaller particle sizes and approaching the borderline between dust and fumes.

As illustrated in Figure 10.7, a cyclone consists of a vertical cylinder with a conical bottom, a tangential inlet near the top, and outlets at the top and the bottom, respectively. The top outlet pipe protrudes into the conical part of the cyclone in order to produce a vortex when a dust-laden gas (normally air) is pumped tangentially into the cyclone body. Such a vortex develops centrifugal force and, because the particles are much denser than the gas, they are projected outward to the wall flowing downward in a thin layer along this in a helical path. They are eventually collected at the bottom of the cyclone and separated. The inlet gas stream flows downward in an annular vortex, reverses itself as it finds a reduction in the rotation space due to the conical shape, creates an upward inner vortex in the center of the cyclone, and then exits through the top of the cyclone. In an ideal operation in the upward flow

there is only gas while in the downward flow there are all the particles fed with the stream. Cyclone diameters range in size from less than 0.05 m to 10 m, feed concentrations cover values from 0.1 kg/m³ to approximately 50 kg/m³, while gas inlet velocities may be in the order of 15–35 m/s.

10.3.2.2 Theoretical Aspects

A cyclone is in fact a settling device in which a strong centrifugal force, acting radially, operates instead of the relatively weak gravity force, acting vertically. Due to the small range of particles involved in cyclone separation (the smallest particle that can be separated is about 5 μ m), it is considered that Stokes law primarily governs the settling process. Substituting, thus, the expression for centrifugal acceleration $g = \omega^2 r$ into Equation 10.11, the resultant equation is

$$v_{\rm r} = \frac{x^2 (\rho_{\rm s} - \rho_{\rm g}) \omega^2 r}{18 \mu_{\rm o}}$$
 (10.31)

where v_r is the terminal velocity of the particle covering the radius r, x is the equivalent particle diameter, ρ_s the density of the particle, ρ_g the density of the gas, and μ_g the viscosity of the gas.

Also, the centrifugal acceleration is a function of the tangential component of the velocity according to the following expression:

$$\omega = \frac{v_{\text{tan}}}{r} \tag{10.32}$$

where v_{tan} is the tangential velocity of the particle in the radius r. Substituting Equation 10.32 into Equation 10.31:

$$v_{\rm r} = \frac{x^2(\rho_{\rm s} - \rho_{\rm g})v_{\rm tan}^2}{18\mu_{\rm g}r}$$
 (10.33)

Multiplying Equation 10.33 by g/g, the resultant equation gives

$$v_{t} = \left[\frac{x^{2} (\rho_{s} - \rho_{g}) g}{18 \mu_{g}} \right] \frac{v_{tan}^{2}}{gr} = (u_{t}) \frac{v_{tan}^{2}}{gr}$$
(10.34)

where u_t is the terminal settling velocity defined by Equation 10.11. As can be implied, and according to Equation 10.34, the higher the terminal velocity, the easier it is to "settle" a particle within a cyclone.

10.3.2.3 Operating Variables

For a given particle size, the terminal velocity is a maximum in the inner vortex, where *r* is small, so that the finest particles separated from the gas are eliminated in the inner vortex. These particles migrate through the outer vortex to the wall of the cyclone and drop passing the bottom outlet. Smaller particles that do not have time to reach the wall are retained by the air, and carried to the top outlet. Although the chance of a particle for separation decreases with the square of the particle diameter, the fate of a particle depends also on its position in the cross section of the entering stream and on its trajectory in the cyclone. Thus, the separation according to size is not sharp. For cyclones the cut diameter or cut size, can be defined as that diameter for which one-half of the inlet particles, by mass, are separated while the other half are retained by the gas. The cut size is a very useful variable to determine separation efficiency of a cyclone. Since a given powder to be separated in a cyclone would have an extremely fine half of its distribution, such half may not be easily separated using conventional pressure drops. Therefore, it is advisable to make it the cut size to coincide with the mean size of a powder particle size distribution to guarantee separation of the coarse part of such distribution, as the fine one may unattainable due to the small range involved. According to this criterion, the cut size would represent the size to be captured inside the cyclone to guarantee a 50% collection of the fed particles. Substituting such size in Equation 10.34 would fix a velocity to carry out the desired separation. Since in practice evaluation of radial velocity is difficult as it depends on the terminal gravitational velocity, the tangential velocity, and the radial and axial position of the cyclone, empirical relations alternative to Equation 10.34 have been suggested. An alternative equation that has been used for this purpose is the following:

$$v_{\rm t} = \frac{x^2(\rho_{\rm s} - \rho_{\rm g})b_1}{18\mu_{\rm g}r^n}$$
 (10.35)

where b_1 and n are empirical constants.

Experience and theory have shown that there are certain relationships among cyclones dimensions that should be observed for efficient cyclone performance (Geldart, 1986), and which are generally related to the cyclone diameter. There are several different standard cyclone "designs" and a very common is called the "Stairmand" design, whose dimensions are shown in Figure 10.8. Using standard geometries of cyclones is much easier to predict effects on variables changes and scale-up calculations are greatly reduced. Such calculations may be carried out by means of dimensionless relationships. Selection and operation of cyclones can be described by the relationship between the pressure drop and the flow rate, and the relationship

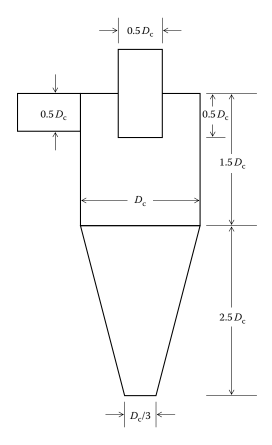


FIGURE 10.8 Dimensions of standard Stairmand cyclone.

between separation efficiency and flow rate (Svarovsky, 1981). The fundamental equations to apply the dimensionless scale-up model are

$$Eu = \frac{2\Delta P}{\rho_g v^2} = k \tag{10.36}$$

$$Stk_{50} = \frac{4Qx_{50}^2\rho_s}{18\mu_g\pi D_c^3} = k$$
 (10.37)

where Eu is the Euler number for a given pressure drop ΔP of a gas of density $\rho_{\rm g}$ flowing through the cyclone at a velocity v. Stk₅₀ is the Stokes number for a cut size x_{50} of a solid of density $\rho_{\rm s}$ suspended in a gas of viscosity $\mu_{\rm g}$ passing through the cyclone of diameter $D_{\rm c}$ at a volumetric flow rate Q. The cyclone body velocity v is the characteristic velocity that can be defined in various

ways, but the simplest one is based on the cross section of the cylindrical body so that

$$v = \frac{4Q}{\pi D_c^2}$$
 (10.38)

As can be implied by inspecting Equations 10.36 through 10.38, for given values of ΔP , ρ_s , ρ_g , and μ_g : $X_{50} \sim (D_c)^{0.5}$, whereas for a given value of X_{50} : $\Delta P \sim (Q)^{2/3}$; $\Delta P \sim (D_c)^2$; and $Q \sim (D_c)^3$.

When designing cyclone systems, it is essential for optimum results to have full process data available, as there are a number of variable factors that will determine the final performance.

Cyclone inlet velocity not only affects efficiency of separation but also reflects in pressure loss and possible erosion. Gas viscosity has an important effect on particle efficiency, and so it is advisable to check its dependency with temperature and consider those cases in which a gas different from air is involved in the process. Smaller cyclone diameters increase overall efficiency, but also promote erosion. In addition to this, it is sometimes necessary to consider possible attrition of solids in the cyclone, which will result in production of fines and considerable losses. Erosion occurs primarily where the particles first impact the cyclone wall, but also occur at the bottom of cyclones too short to accommodate the length of the naturally occurring vortex.

In order to remove as much as the dust from a gas stream as possible, cyclones are often placed in series with each other to increase the overall collection efficiency relative to a single unit. Although this is theoretically true, the efficiency is increased at the expense of pressure drop, and so extreme care should be taken to consider whether great efficiency will render a costly process. There is also a great complexity when trying to improve collection efficiency using many small cyclones in parallel to replace a large cyclone. A disadvantage of parallel cyclones is that equal gas distribution to each cyclone can be difficult to achieve. If this occurs, collection efficiency is reduced with the advantages of parallel cyclones being greatly negated.

There is an increasing trend toward using cyclones for final cleaning for environmental purposes in processes where it is becoming economically impractical to use other methods. There are many applications where this can be done subject to certain limitations, by the use of one, two, or even three stages of separation. In many cases a first-stage cyclone can be used to deal with high inlet loadings and will produce excellent results at the expense of reasonable pressure loss. The addition of a second stage to deal with the first-stage losses can then often achieve required results, but where further cleaning is necessary a third-stage cyclone may provide the answer. The use of a third-stage cyclone invariably means that a high degree of clean-up is necessary and, therefore, the third stage should give the best possible efficiency and be capable of maintaining this efficiency for a long time. The use of several stages to try to improve efficiency or remove very fines particles

may become impractical by employing only cyclones. This difficulty can be alleviated by combining methods so that, for environmental purposes when certain particles need to be removed regardless of their fineness, bag filters may be coupled with cyclones.

10.3.2.4 Applications

As mentioned before, cyclones are extensively used in the processing industry to reduce particle load to safe levels in dry milling, as well as in classification of particles in closed-circuit grinding operations. They are also employed in recovering fines from spray drying and fluidized bed drying processes. Another important application is in pneumatic conveying of diverse food products, such as grains and flours.

Example 9.3

Calculate the cut size for particles suspended in air and treated in a cyclone 1.48 m in diameter operating at a pressure drop of 1200 Pa. The density of the particles id 2.6 g/cm³, the density of the suspending air is 1.2 kg/m³, while its viscosity is 1.825×10^{-5} N s/m². The pressure drop coefficient is 320 and the Stokes number is 0.00012.

SOLUTION

As can be deducted, the pressure drop coefficient refers to the Euler number. Substituting, thus, Equation 10.38 into Equation 10.36 and transposing for *Q*, and substituting values on the resulting equation, that is,

$$Q = \sqrt{\frac{\Delta P \pi^2 D_c^4}{8 \rho_g \text{Eu}}} = \sqrt{\frac{(1200) \text{ kg/s}^2 m (\pi)^2 (1.48)^4 m^4}{8 (1.2) \text{ kg/m}^3 (320)}} = 4.3 \text{ m}^3/\text{s}$$

Using the above flow rate, and transposing for the cut size in Equation 10.37, it is substituted along with the other values to obtain the cut size as

$$x_{50} = \sqrt{\frac{\text{Stk}_{50}(18)\mu\pi D_c^3}{4Qp_s}} = \sqrt{\frac{0.00012(18)(1.825 \times 10^{-5}) \text{ kg/ms}(\pi)1.48^3 \text{ m}^3}{4(4.3)\text{m}^3/\text{s}(2600) \text{ kg/m}^3}}$$

The cut size is, therefore: $x_{50} = 2.99 \times 10^{-6} \text{ m} \sim 3 \text{ }\mu\text{m}$.

10.3.3 Air Classifiers

10.3.3.1 Description of the Technique

Air classification is a method of separating powdery, granular, or fibrous materials in accordance with the settling velocity and combined with the influence of particle size, particle density, and particle shape. The procedure of winnowing or aspiration is a traditional way of separating chaff from grain after threshing and is one of the simplest forms of air classification. Ideally, the separation effect of an air classifier should be such that all particles that exceed the cut point are transported into the coarse fraction and the smaller particles are transported to the fines fraction. In this sense, air classification basically consists of dividing particle size distributions of given powders and, as such, is a technique commonly used in combination with size reduction equipment, normally to eliminate fines that may affect properties like wettability and dispersibility. The major interest in air classification is that it provides a means for separating small particles, in a dry manner, which cannot be readily achieved by sieving, that is, below 50 μm . Ortega-Rivas and Svarovsky (2000) report a successful sharp split of the particle size distribution of calcium carbonate, into a fine fraction with mean particle size as fine as 6 μm .

10.3.3.2 Theoretical Aspects

The mode of operation of a typical air classifier is as follows. The inlet air is mixed with the material being separated. The feed particles are subjected to a centrifugal force operating from a revolving rotor and a drag force produced by the air current, which moves in a spiral direction toward the central shaft. As previously mentioned, the separation is based on differential mass, density, and shape. The larger and more dense particles are influenced by the mass-dependent centrifugal forces and move toward the outside of the chamber, where they are removed by a screw conveyor or some alternative means. The smaller, lighter particles are more subjected to the frictional forces of the air current moving, therefore, with the airstream and leaving from the center of the classifying chamber to be separated by a cyclone from the airstream. The relative magnitude of these two forces can be changed by altering the rotational speed of the disc and the air velocity. By varying either of these, there will be an effect on the cut size. A common design of air classifier consists of a rotating wheel with zigzag channels over its surface, each of them comprising six components. A diagram of this type of classifier is shown in Figure 10.9.

Classifiers with the facility to change rotational speed and air velocity independently will be very flexible in operating terms. Although the separation mainly takes place within the classifying chamber, some preliminary removal of the coarsest particles may be achieved outside the chamber. The disc or turbine can be mounted on a horizontal or vertical axis. The latter produces a centrifugal force in the horizontal plane, favoring high throughputs but low precision in cut size (Fedoc, 1993). Air classifiers are categorized by reference to a number of factors such as the presence or absence of a rotor, the drag force on the air, the relative velocity and direction of the air and particles, the use of directional devices such as vanes or cones, and the

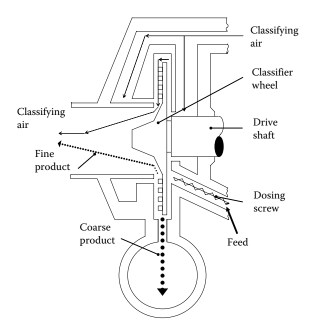


FIGURE 10.9 Diagram of an air classifier.

location of the fan and fines collection devices. Other important features include the capacity of the classifier and the energy utilization. A comprehensive treatise of classifier types has been presented by Klumpar et al. (1986).

The obvious method of characterization of separation capability of air classifiers is by using the cut size concept. Ideally, all particles below the cut size would end up in the fines stream while all particles above the cut size would follow the coarse stream. However, there will be always misplaced material, that is, a small fraction of particles smaller than the cut size would be in the coarse stream and an equally small proportion of particles larger than the cut size would appear in the fines stream. The extent of the overlap due to misplaced material, as well as the cut size, can be determined by measuring the particle size distributions of both streams, and presenting their data as a weight frequency distribution. The yields of fines $Y_{\rm f}$ and coarse $Y_{\rm c}$ streams need to be identified. When they are equal, the point of overlap gives the cut size. When they are not equal, which is most likely, the frequency distribution for the fines stream must be multiplied by the yield for the fines stream, while the yield for the coarse stream must be multiplied by the point of intersection of these curves.

In air classifiers a solid-state element in a rotor is radially affected by two forces: a centrifugal force toward the outside and the frictional force of the air toward the inside. Since particles quite fine are of interest, assuming Stokes law applicability, the resulting equilibrium between the two mentioned forces for a certain grain size would be the so-called cut size. Particles larger than the cut size are centrifugally extracted as oversized toward the outside, while smaller particles are carried inside by the airstream. Assuming a constant air throughput, the fineness of the separation in the classifier depends on the peripheral speed of the particle, which in turn is in conformity with the peripheral speed of the rotating wheel. With the speed remaining constant, an increasing air throughput changes the cut point within the coarser range. In principle, any cut point can be attained by combining two matching values of speed and air throughput.

By equating the above-mentioned forces when they are in equilibrium, a relation for the cut size x_{50} based on Stokes law can be derived as follows:

$$x_{50}^2 = \frac{18\mu_{\rm g}v_t r}{v_{\rm tan}^2(\rho_{\rm s} - \rho_{\rm g})}$$
 (10.39)

where $v_{\rm t}$ is the radial speed of air, r is the radius of the classifier wheel, and $v_{\rm tan}$ is the peripheral speed of particle. Equation 10.39 is useful in predicting how airflow and rotational speed may influence the cut size. Basically, it predicts that increasing the airflow rate would increase the cut size, whereas increasing the rotational speed would decrease this cut size.

10.3.3 Operation and Applications

Due to the complicated flow conditions in air classifiers, the classifying results with the given values of speed and air throughputs cannot be predetermined. For this reason, the assignment of cut point and operating data of the classifier are determined by experiments with a calibration curve. The various particle sizes of a given material of known distribution are used to plot the calibration plot. These particle sizes are separated under certain conditions of speed and air throughput. The coarse fraction is weighed at the same time. The particle size corresponding to the coarse grain proportion and legible from the fineness characteristics is the cut point according to the operating conditions of the individual classifier, also known as analytical cut point. For inert materials with densities of the order of 2600 kg/m³, cut points between 1 and 100 µm are obtained with a "normal" air throughput (0.5–3 kg/h) and by varying the speed between 2500 and 20,000 rpm. Figure 10.10 shows a calibration curve for these variables. Within these ranges, the volumetric flow rate of air V is related to the peripheral velocity N by the following expression:

$$V = 55 - \frac{N}{1000} \tag{10.40}$$

where V is expressed in m^3/h and N in rpm.

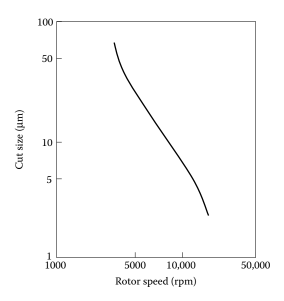


FIGURE 10.10Calibration curve for an air classifier.

For materials different to the one used for obtaining the graph, if properties are not radically different it can be shown that the equation below will give possibility of evaluating a new cut size:

$$\frac{(x_{50})_1}{(x_{50})_2} = \sqrt{\frac{\rho_2}{\rho_1}} \tag{10.41}$$

where $(x_{50})_1$ and ρ_1 are the characteristics of the material being evaluated, while $(x_{50})_2$ and ρ_2 correspond to the calibration curve material (calcium carbonate for Figure 10.10).

Air classification is used in a number of industries in important applications. For example, it is used in the pharmaceutical industry to determine the absorbing capacity of drugs and active substances, which is a function of their fineness. In grain processing, an application is the wheat flour fractionation to separate the coarse, low protein fraction, from the fine, high protein fraction. Other food-related applications include classification of confectionery products, soy flour, potato granules, rice flour, lactose, and oleaginous fruits, removal of shells or hulls from disintegrated peanuts, cottonseed, rapeseed, or cocoa beans, preparation of oat-bran, and separation of gossypol from cottonseed protein. Air classifiers are widely used in many comminution processes in the mining, chemical, and other types of industries, for separation of specific fraction for diverse purposes and applications.

10.3.4 Gas Filters

10.3.4.1 Description of the Process

Gas-solids separations can be performed using filtration means. Gas filters are used for final particulate removal in many processes of the chemical and related industries. These filters can capture particles much smaller in diameter than a cyclone so they are commonly placed downstream of a cyclone in diverse applications. A gas filter generally consists of a porous fabric, which can be woven to conform to the shape of a cylinder or may be supported in a frame. The former is called a bag filter while the latter is known as an envelope filter. The main difference of both designs is in the way solids are accumulated. In bag filters dust may accumulate inside, whereas in envelope filters it would form a cake outside. The filtering arrangements are placed in a matrix so that their total area will result in a low gas velocity through the bags and, therefore, a low pressure drop through the filter. Gas velocities through the filtering media are of the order of 0.005-0.02 m/s. Particulate loadings to these filters generally lie in the range of 0.2-250 g/cm³. Diagram of the two main types of gas filters are given in Figures 10.11 and 10.12. Filters used in gas-solids separations may be woven or felted fabrics of natural or synthetic fibers. There are tables listing properties of filtering media to determine whether they are suitable for applications at diverse conditions, such as high temperatures, corrosive and chemical resistance, and so on (Green and Perry, 2008). Table 10.4 presents properties of some fabrics and fibers used in air filtration, while Table 10.5 gives resistance and permeability factors for some air filtration fabrics.

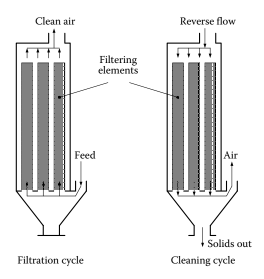


FIGURE 10.11Bag filter with internal accumulation of solids.

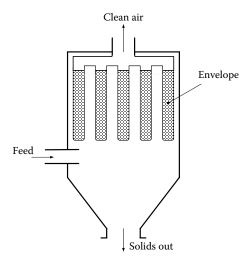


FIGURE 10.12Bag filter with external accumulation of solids.

Filtrations may also be made with granular solids in the form of stationary or moving beds. Many other types of materials porous in nature, or capable of providing a screening effect after weaving or fabrication, may be found suitable for certain filtration applications.

In gas filtration, the collected particles build up on the surface of the filtering medium and cause a gradual increase in the pressure drop through the

TABLE 10.4Physical Properties of Some Fabrics and Fibers Used in Air Filtration

Fiber	Weight (g/m²)	Thickness (mm)	Rupture Force (N/mm)	Elongation (% to Rupture)	Air Permeability $[(m^3/s)/m^2 a$ $\Delta p_f = 0.12 \text{ kPa}]$	K_{c}
Wool	859.3	3.43			0.138	1.03
Wool	788.7	3.28			0.151	0.93
Orlon	405.5	1.14	11.38	18	0.102 – 0.127	1.11-1.39
Orlon	665.9	2.24	14.89	18	0.076-0.102	1.39-1.85
Orlon	892.8	3.18	19.26	60	0.051 - 0.102	1.39-2.78
Acrylic	665.9	1.91	17.51	22	0.076-0.102	1.39-1.85
Dynel	892.8	3.18	10.51	80	0.152-0.203	0.70 - 0.93
Dacron	665.9	2.03	21.89	22	0.076-0.102	1.39-1.85
Dacron	368.3	6.35	3.50	150	1.016-1.143	0.11 – 0.14
Dacron	892.8	3.18	30.65	80	0.102 – 0.152	0.93 - 1.39
Nylon	892.8	3.18	17.51	100	0.152-0.203	0.70 - 0.93
Arnel	892.8	3.18	10.51	80	0.152-0.203	0.70 – 0.93
Teflon	580.3	1.35			0.419	0.34

TABLE 10.5	
Resistance and Permeability Factors for So	me Fabrics

Fabric	Weight (g/m²)	Thread Diameter (mm)	Pore Size (mm)	Air Permeability [(m ³ /s)/m ² a $\Delta p_{\rm f} = 0.12~{\rm kPa}$]	$K_{\rm c}$
Twilled cotton	196.4	0.254	0.254	2.396	0.09
Shiny cotton	255.9	0.229	0.178	0.523	0.27
Shiny cotton	306.2	0.254	0.127	0.163	0.88
Shiny cotton	379.4	0.279	0.102	0.127	1.12
Lana	427.8	0.356		0.427	0.33
Spun glass		0.762		0.009	1.60
Dacron	498.5			0.005	0.29
Teflon	323.6			0.102	1.39

filter. After a certain limiting pressure drop is reached, the bags are cleaned by pulsing gas back through the filter to remove the solids buildup on the bag. As described above, standard commercial types of filters use a bag or an envelope, the latter being actually a retainer for the filter cloth. The bag filters are, in fact, elongated cylinders that may be opened at both ends, to allow a cleaning cycle by using air jets blowing into the top. On the other hand, envelope filters, with a frame for support and with large flat surfaces exposed, do not present the cleaning capabilities of the bag filters, which can be given more shaking or can be collapsed without risking any harm. When air is blown in reverse direction to filtration flow for cleaning purposes, such cleaning can be made more effective. The envelope filters are, therefore, best used on dusts that are easily shaken or removed from the cloth surface. The dust is removed in the envelope type by beating the screen supporting the filter and by shaking or rocking the frame by mechanical means.

The use of a blow ring in some bag- or cylinder-type mechanical filters allows for continuous operation while performing the cleaning duty. Dust is collected on the inside while the blow ring travels up and down along the outer surface of the bag. The blow ring has an inside slot used to blow gas against the bag wall. The ring is tight enough to partially collapse the bag in order to break the dust cake, and provide a close seal so the gas blown is fully delivered through the filter when blowing back. Dust may be collected on the outside of filter tubes or bags if a support is provided inside to prevent the filter collapsing. Sometimes, it is possible to remove the cake from the outer wall by periodically using jet-compressed air from inside the filter, in order to produce a shock to break the cake from the outer wall whence it can be settled into the bin. The envelope- or frame-type filter collects the dust on the outer wall of the filter, as expected, since the outer wall is easier to get to in this arrangement.

10.3.4.2 Operation Characteristics and Applications

The operating variables of gas filtration are the resistance to flow, the permeability of air to the filtering medium, and the resistance due to particle accumulation. With regard to the resistance to flow, the pressure drop across the filtering medium ΔP_f can be represented by

$$\Delta P_{\rm f} = K_{\rm c} \mu_{\rm g} V_{\rm f} \tag{10.42}$$

where K_c is a constant depending on the filtering medium and V_f is the gas superficial velocity through the filtering medium.

The resistance to the layer of particles accumulated during the filtration cycle can be calculated determining a variable known as cake resistance factor K_1 :

$$K_1 = \frac{\Delta P_{\rm c}}{V_{\rm f}w} \tag{10.43}$$

where ΔP_c is the pressure drop through the powder layer and w is the powder mass flow rate approaching the filter. Table 10.6 presents some values of the factor K_1 .

As previously mentioned, bag filters have, practically, the same applications of cyclones, being normally coupled with these in order to remove the finest tails of particle size distributions of diverse industrial powders. They can be used in recovery of fines fractions in spray drying and pneumatic conveying. They are also widely used in industries and many other places for purification of ambient air. Recovered materials by filtration are varied and include granite powder, fumes from combustion processes, fine dust released in many industrial applications, and so on.

10.3.5 Scrubbers

Wet collectors, also called scrubbers, refer to a type of separating device in which a liquid is used to aid the collection of dusts, fumes, and mists in a

TABLE 10.6Cake Resistance Factors for Some Powders

	K_1 for Particle Smaller than					
Powder	850 µm	106 µm	90 µm	45 μm	20 µm	
Granite	1.58	2.20			19.8	
Stone	0.96		6.30			
Wood			6.30			
Resin		0.62			25.2	
Oates	1.58		9.60	11.0		
Maize	0.62		3.78	8.80		

number of industrial operations. There are numerous designs of scrubbers commercially available that have been devised over the years, but they operate basically on the same principle in which solid particles may be often scrubbed from a gas stream by spraying a jet of a liquid, usually water, into that stream. Depending on their design and operating conditions, particle scrubbers can be adapted to collect fine or coarse particles that are intercepted by the droplets of the water spray and are removed by the scrubber in the form of slurry. The predominant mechanism of particle entrapment by the liquid appears to be inertial deposition. Flow-line interception is only a minor mechanism in the collection of the finer dust particles by liquid droplets of the sizes encountered in scrubbers. Diffusion is indicated to be a relatively minor mechanism for the particles larger than 0.1 µm that are of principal concern. Although scrubbers are normally more efficient than cyclones, they have the disadvantage of collecting the solids wet instead of dry. If this feature is unacceptable or impractical for processing reasons, the solids must be separated from the liquid. In such a way, the gas-solids collection problem would be replaced by a solid-liquid separation difficulty.

10.3.5.1 Process Description

The predominant separation mechanism in scrubbing is impact due to the inertia of a particle with density relatively higher than that of the liquid. In such a way, the relative velocity between the particle and the liquid droplet is an important factor and the process may depend on the acceleration of, both, the particle and the droplet in relation to the point of impact. The process may be controlled by varying the droplet velocity, in order for the particle to collapse with it. The efficiency of the scrubbers is normally expressed as the fraction of particles collected from a volume of gas, determined by the projected area of the receiving body. To achieve this, a separation number Se (Maas, 1979), may be defined as

$$Se = \frac{u_t V_0}{g_1 D_b} \tag{10.44}$$

where $u_{\rm t}$ is the Stokes terminal velocity, V_0 is the mean gas velocity, $g_{\rm L}$ is the local gravity acceleration, and $D_{\rm b}$ is the liquid droplet diameter. The efficiency of separation η can be correlated to the separation number Se using graphs like the one shown in Figure 10.13.

10.3.5.2 Equipment and Applications

A typical device, known as spray cyclonic scrubber, consists on a cylindrical chamber with a conical bottom as illustrated in Figure 10.14. The scrubber

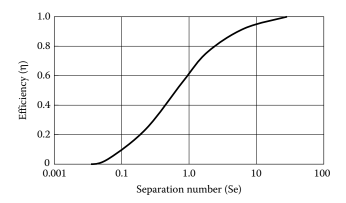


FIGURE 10.13 Efficiency of separation versus separation number for spherical droplets in scrubbers.

operation is countercurrent with the feed entering by a tangential port at the bottom and traveling upward through a water layer formed by the spray falling on the swirl vanes mounted on paddles. After the gas current passes through the water layers toward the internal center of the vanes, it changes direction crossing the paddles. An upper vane is used to separate the

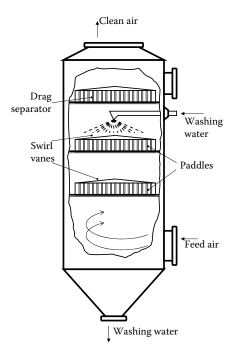


FIGURE 10.14 Diagram of spray cyclone scrubber.

remaining droplets and allow the air to come out clean from the top outlet, as shown in Figure 10.14. This equipment combines the cyclonic action with the scrubbing of the liquid.

Other types of equipment used for scrubbing include plate towers, self-induced spray scrubbers, venturi scrubbers, mechanical scrubbers, and fiberbed scrubbers. The principle of operation of all these units is, practically, similar. Their operating principles can be consulted in reference textbooks (Green and Perry, 2008).

Gas scrubbers have the same sort of applications of the previously described solid–gas separators. They are particularly useful in the separation of fumes from combustion reactions and to eliminate mists in production of different acids in many industrial processes.

10.3.6 Other Techniques

10.3.6.1 Settling Chambers

Gravity collectors are generally built in the form of long, empty, horizontal, rectangular chambers with an inlet at one end and an outlet at the side or top of the other end (Figure 10.15). Inside that chamber the gas velocity is reduced to enable dust to settle out by the action of gravity. Its simplicity lends it to almost any type of construction. Practically, however, its industrial utility is limited to removing particles larger than 43 mm diameter. For removing smaller particles, the required chamber size is generally excessive. By assuming a low degree of turbulence relative to the settling velocity of the dust particle in question, the performance of a gravity settling chamber expressed as efficiency of separation η may be given by

$$\eta = \frac{u_t L_s}{H_s V_s} \tag{10.45}$$

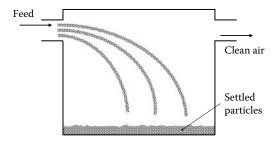


FIGURE 10.15 Dust-settling chamber.

where u_t is the terminal settling velocity of the particle, L_s is the length of the chamber, H_s is the height of the chamber, and V_s is the superficial velocity of the gas through the chamber.

Since the pressure drop through a settling chamber is small, and because low gas velocities are used, the chamber is not subject to abrasion and may therefore be used as a precleaner to remove very coarse particles and thus minimize abrasion on subsequent equipment.

10.3.6.2 Electrostatic Separators

This type of equipment separate solids, or liquids, from a gas stream by passing it through a strong, high-voltage field produced between two electrodes of opposite polarity. The field imposes a charge on the particles so that they migrate toward the collecting electrode. The particles are usually removed from the electrode by periodic rapping. The advantage of an electrostatic precipitator is that it can collect solids of very small size in a dry form. Electrostatic precipitators are generally large units of equipment, because collection efficiency is proportional to the area of the collecting electrodes. For this reason, capital costs are very high, although operating costs are low and may justify the use of a precipitator instead of a filter for some particular applications (Svarovsky, 1981).

Electrostatic separators can be selected according to the following fundamental equation:

$$\eta = 1 - e^{-w(A/V)} \tag{10.46}$$

where η is the efficiency of separation, w is the migration velocity, A is the area of the collecting electrodes, and V is the volumetric flow rate of gas.

Theoretically, the migration velocity w can be evaluated from

$$w = \frac{aE_0 E_p}{6\pi\mu_g} \tag{10.47}$$

where a is the radius of the dust particle, E_0 is the strength of the electrical field where the particle is charged, E_p is the strength of the field in the settling point, and μ_g is the gas viscosity.

Electrostatic precipitators have been used successfully to eliminate fine dusts in many types of waste gases. They have been employed in settling of fine powders of lead oxide, arsenic, and foundry gases. Other applications include settling of dust from gases produced in cement ovens, and in the recovery phosphorous pentoxide from gases produced in phosphorous ovens.

10.4 Solid-Liquid Separation Techniques

As previously mentioned, solid-liquid separations constitute an important number of operations in diverse processing industries. Solid-liquid separation is an important industrial process used for recovery of solids from suspensions and/or purification of liquids. Dense, concentrated, and dilute suspensions handled in many of these industries need to be depurated or purified by separating phases, and the desired product can be the clarified liquid or the dried solid. The different theoretical aspects needed to understand the operating principles of all the mechanical operations employed in separating components in two-phase and multiphase systems can be facilitated by, first, dealing with the general aspects of properties of suspensions. Properties and pretreatment of suspensions will be studied in the first part of this section, prior to get involved in the study if the different solid-liquid separation techniques. Many slurries and suspensions treated in the materials processing industry behave as non-Newtonian liquids but the current theory describing the solid-liquid separation process normally applies to Newtonian suspensions only. The reviewing part dealing with properties of suspensions will therefore include a component dealing with non-Newtonian suspensions.

10.4.1 Properties of Suspensions: Rheology and Flow

The rheological properties of suspension have been studied since the beginning of last century. The first research was done by Einstein (1906, 1911), in his classical study of the viscosity of dilute suspension of rigid spheres. His approach is purely hydrodynamic, and his model consists of an isolated sphere situated in a simple shear flow field in an infinite fluid. A number of workers have been extending Einstein's analysis (Frisch and Simha, 1956). Three approaches have been presented: the theoretical basis of viscosity computation, the effect of particle texture and shape, and the effect of concentration. Some relationships have been developed for correction of viscosity of suspensions in which nearest-neighbor interactions cannot be neglected. Guth and Simha (1936) considered the first-order effect of spheres interacting with one another. They found that

$$\frac{\mu}{\mu_o} = 1 + 2.5\phi + 14.1\phi^2 \tag{10.48}$$

where μ is the viscosity of the suspension, μ_o is the viscosity of the pure solvent, and \emptyset is the volume fraction of spheres in the suspension. A later correction gave a value of 12.6 for the last constant (Simha, 1952).

For more concentrated systems other approaches have been given. Several authors have made experimental studies of the effects of concentration on the

viscosity of suspension of spheres. The data they have proposed are considerably scattered. Thomas (1963) has tried to find the sources of scatter in the data. He found that it is caused in large part by variations in particle size. For small particle sizes (diameters <1–10 μ m) colloidal forces become important and the viscosity begins to increase as particle size decreases. The viscosity is also shear rate dependent in this case. For particles larger than 1–10 μ m, the particle Reynolds number becomes significant. The inertial effects results in an increase in relative viscosity with increasing particle size. Thomas has developed a unique curve, by eliminating diameter and shear rate effects. For low-concentration data, such a graph can be represented by

$$\frac{\mu}{\mu_o} = 1 + 2.5\emptyset + 11.4\emptyset^2 \tag{10.49}$$

Equations 10.48 and 10.49 would be used for correcting the viscosity of concentrated suspensions following a Newtonian behavior.

Many concentrated suspensions follow non-Newtonian behavior and considerations ought to be made as the viscosity is not a constant anymore, and such fluids should be characterized properly to use their parameters of characterization instead of viscosity in calculations using some dimensionless groups. Most of the theory developed for non-Newtonian fluid flow through tubes, apply to laminar flow round smooth tubes. The most studied non-Newtonian fluids are the power-law fluids and there are some relationships available for pressure-loss-flow rate in purely viscous and viscoelastics flows principally. Relations for thixotropic and rheopectic systems appear not to be available.

Under the assumptions that purely viscous behavior prevails and that no slip occurs at the tube wall, the power-law equation for laminar flow is in the form (Geankoplis, 2003):

$$\tau_{\rm w} = \frac{D\Delta P}{4L} = K' \left(\frac{8v}{D}\right)^{n'} \tag{10.50}$$

where τ_w is the shearing stress at the wall of the tube, D is the tube diameter, ΔP is the pressure drop, L is the length of the tube, v the mean velocity, n' is the slope of the line when the data are plotted on logarithmic coordinates. For n'=1, the fluid is Newtonian; for n'<1, pseudo plastic, or Bingham plastic if the curve does not go through the origin; and for n'>1, dilatant. The term K' is the consistency index; as the name suggest, the larger its value the thicker or more viscous the fluid.

It may be shown (Dodge and Metzner, 1959) that the constant K' may be related to the analogous power-law constant K (Table 10.1) as follows:

$$K' = K \left(\frac{3n'+1}{4n'}\right)^n \left(\frac{8v}{D}\right)^{n-n'}$$
 (10.51)

if the fluid obeys the power law, n = n' and

$$K' = K \left(\frac{3n+1}{4n}\right)^n \tag{10.51b}$$

Since Equation 10.50 rigorously portrays the laminar flow behavior of the fluid (provided n' and K' are evaluated at the correct shear stress), it may be used to define a Reynolds number applicable to all purely viscous fluids under laminar flow conditions. This dimensionless group can be derived simply by the substitution of $(D\Delta P/4L)$ from Equation 10.50 into the usual definition of the fanning friction factor, that is,

$$f = \frac{D\Delta P/4L}{\rho v^2/2} \tag{10.52}$$

such substitution leads to

$$f = \frac{16\gamma}{D^{n'}v^{2-n'}\rho}$$
 (10.53)

where $\gamma = K'8^{n'-1}$ and all the remaining components as already defined.

By letting f = 16/Re as for Newtonian fluids in laminar flow, the above-mentioned generalized Reynolds number can be obtained as

$$Re^* = \frac{D^{n'}v^{2-n'}\rho}{\gamma} \tag{10.54}$$

If the equation is desired in terms of *K* instead of *K'*, Equation 10.51b may be substituted into Equation 10.54 and

$$Re^* = \frac{D^n v^{2-n} \rho}{K8^{n-1} (3n+1/4n)^n}$$
 (10.55)

For Newtonian fluids n' = 1 and $K' = \mu$, so γ reduces to μ and Re* in Equation 10.55 transforms to the familiar $Dv\rho/\mu$ showing that this traditional dimensionless group is merely a special restricted form of the more general described here.

By far, most of the study of the flow behavior of non-Newtonian suspensions has been devoted to monosized and monoshaped systems, mainly of spherical particle geometry. In reality, however, solid–liquid mixtures usually consist of dissolved solids with a variety of particle sizes, shapes, and concentrations. The available theory for the behavior of non-Newtonian

suspensions is, therefore, of a limited application, and when dealing with real systems, empirical corrections are necessary to fully explain the suspension properties. Studies of some real applications such as slurry transport are also encountered in the literature. The slurry to be transported may have settling or nonsettling characteristics. Sand in water, salt in benzene and clay in drilling mud are typical examples of settling slurries, while paper pulp is a good example of nonsettling slurry. The law governing the settling rate of particles at low Reynolds numbers is the well known Stokes law. Settling phenomena of slurries in pipelines, as well as modifications of original Stokes law for flocculated suspensions and particles of odd shapes, have been studied by Thomas (1961, 1962). Metzner and Whitlock (1958) have found non-Newtonian viscosity effects at high volume fractions in the laminar regime. Such effects are mainly of the shear thinning type if the particles are large, although shear thickening is sometimes the type. The effects of shear rate and particle diameter in concentrated suspensions have been also investigated. For monodispersed suspension the relative viscosity decreases with increasing shear rate for concentrations less than about 0.5, and increases with increasing particle size. The turbulent regime is sometimes preferred to transport slurries within pipe in order to prevent settling, and to provide high throughputs. In this case, the viscosity of the suspension has been normally used to correlate the turbulence friction factors (Shaver and Merrill, 1959).

For the case of settling of non-Newtonian suspensions, it has been reported (Ortega-Rivas and Svarovsky, 1993) that the Stokes number Stk_{50} can also be expressed in terms of the parameters of characterization of non-Newtonian suspensions, using a procedure similar to that described above for the Reynolds number. A generalized Stokes number (Stk^*)₅₀ for settling of power-law suspensions can be expressed as follows:

$$Stk_{50}^{*}(r) = \frac{x_{50}^{n'+1}(r)(\rho_{s} - \rho)v^{2-n'}}{18K(3)^{n-1}[2n + 1/3n]^{n}D_{c}}$$
(10.56)

where $x_{50}(r)$ represents the "reduced" cut size including the dead flux effect previously described, ρ_s is the solid density, ρ is the liquid density, v is the superficial characteristic velocity of the separator, and D_c is the characteristic linear dimension of the equipment.

Non-Newtonian fluids in turbulent flow generally show lower friction factors and, consequently, lower pressure drops than do Newtonian fluids at corresponding Reynolds number. A detailed analysis of non-Newtonian flow at turbulent regime began with two papers published by Dodge and Metzner (1959) and Shaver and Merrill (1959). Dodge and Metzner used solutions of carboxymethylcellulose and carbopal (a B. F. Goodrich soluble thickener), as well as clay suspensions. They presented a viscous correlation in terms of parameters from shear-stress—shear-rate data and concluded that

except for the carboxymethylcellulose, the behavior of this material could be correlated with the following generalization of the Newtonian friction factor:

$$\sqrt{\frac{1}{f}} = A_{1n} \log \left[\text{Re}^* (f)^{1 - 0.5n'} \right] + C'_n$$
 (10.57)

where A_{1n} and C'_n are parameters dependent on the flow behavior index n. While the choice of power-law equation in the derivation requires that the Reynolds number used in Equation 10.57 be the power-law special case of Equation 10.54, it has been shown (Dodge and Metzner, 1959) that any errors due to this approximation are less important under turbulent flow conditions than in the laminar region.

The utility of Equation 10.57 and of Reynolds number outside the laminar region has been shown diagrammatically in the above-mentioned paper. They found excellent agreement between experimental and extrapolated data. Furthermore, deviations from smooth curves were no greater for those fluids, which did not obey the power law than for those that did. The parameters A_{1n} and C'_n in Equation 10.57 must be evaluated empirically, just as in the simpler case of Newtonian behavior. An alternative way of presenting the Dodge and Metzner relation above described based on the conventional Reynolds number and an apparent viscosity, evaluated at the wall, has also been proposed (Shaver and Merrill, 1959). Velocity profiles of certain non-Newtonian suspensions have been compared with Newtonian profiles, and are somewhat similar except at high solid concentrations, where they tend to be steeper and laminar like (Bogue and Metzner, 1963). Correlations in terms of viscous parameters have also been reported (Brodkleg et al., 1961). Other works showing non-Newtonian velocity profiles are reported by Seyer and Metzner (1969) and Nicodemo et al. (1969). All these reviews emphasize the need of additional parameters to completely characterize materials in turbulent flow. This can be done either by using a time constant from independent rheological measurements, or by means of molecular properties such as molecular weight, concentration, and the nature of the monomer units. If a variety of materials are going to be treated, the most useful correlating variable is an elastic property. Metzner and Park (1964) used a shear stresses ratio evaluated at the wall shear stress while Seyer and Metzner (1967) used the Deborah number.

Example 10.4

Table 10.7 shows results from testing a suspension of calcium carbonate in a rotational viscometer. Identify the fluid behavior of the suspension and the corresponding parameters of characterization of such fluid.

SOLUTION

In order to identify the behavior of the suspension fluid, the data in Table 10.7 can be plotted to verify resemblance to the typical fluids represented in Figure 10.2. The parameters of characterization can be determined from a logarithmic plot, where *n* would represent the flow behavior index and the intercept of the line on the *y*-axis the fluid consistency index. Plotting data from Table 10.7, Figure 10.16 is obtained.

Comparing the above plot with those in Figure 10.2, it can be concluded that the suspension is a shear thickening or dilatant fluid. Transforming the plot in Figure 10.16 to a logarithmic plot, Figure 10.17 is obtained.

The mathematical function fitting plot in Figure 10.17 is

$$\log \tau = n \log(\dot{\gamma}) + \log K$$

The slope of the line in Figure 10.17 is 1.3953, while at the interception of the line, the values of log τ and log $\dot{\gamma}$ are –2.4485 and 1.6232, respectively. Using logarithms laws, the value for K can be transposed as

$$-2.4485 = \log K + 1.3953(1.6232)$$

$$-2.4885 = \log K + 2.2684$$

$$\log K = -2.4485 - 2.2648$$

$$\log K = -4.7133$$

$$K = 1.9 \times 10^{-5}$$

The values of the characterization parameters of the fluid are: $n \sim 1.39$, $K \sim 0.000019$ kg/m s²⁻ⁿ.

TABLE 10.7 Example 10.4

_	
Shear Rate (1/s)	Shear Stress (Pa)
42.21	0.0268
59.70	0.0504
84.48	0.0670
119.40	0.0892
168.95	0.1394
238.80	0.2144
337.31	0.3082
477.60	0.5896
674.61	0.8040
995.20	1.7152
1349.22	2.4120
1910.40	4.5560

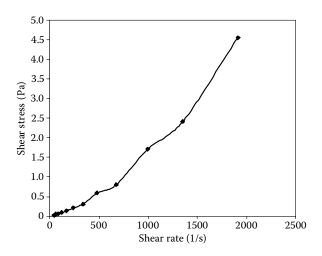


FIGURE 10.16 Example 10.4.

10.4.1.1 Laboratory Exercise: Rheograms of Suspensions

10.4.1.1.1 Introduction

The rheological parameters of non-Newtonian fluids can be estimated by using any type of coaxial viscometer. In these instruments a bob or spindle rotates within a test fluid contained into a cup. The torque necessary to overcome the viscous resistance is measured while the degree to which the spring

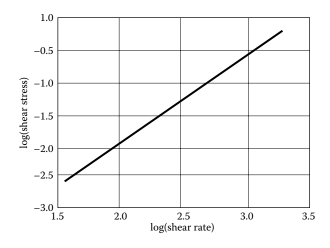


FIGURE 10.17 Example 10.4.

is wound is detected by a rotational transducer, which is proportional to the viscosity of the test fluid. From these measurements data are generated to estimate the shear stress versus shear rate relationship of the fluid. The graph of shear stress against shear rate is called a flow behavior curve or rheogram, and is used to determine the behavior of a testing fluid. The objective of the exercise is to determine rheograms for power-law suspensions using a coaxial cylinder viscometer.

10.4.1.1.2 Instruments and Materials

Any rotational viscometer, such as the Brookfield® model DV, or the Anton-Paar® Rheolab QC, can be used. The suggested test fluids of the pseudoplastic type can be paper pulp or orange juice concentrate, while the dilatant type can be concentrated starch or limestone suspensions.

10.4.1.1.3 Procedure

- Turn on the instrument and attach, carefully the appropriate spindle to the viscometer shaft avoiding any side thrust.
- Insert the spindle into the test fluid up to the immersion groove cut in the spindle shaft.
- Set the desired spindle speed by rotating the spindle control knob or by any control system for that end in the specific instrument.
- Run the test and record the shear stress data at various spindle speeds or shear rates.
- Record the generated data for all the fluid tests run.

10.4.1.1.4 Calculations and Report

- Plot the rheograms for all the test fluids, from the shape of the curves decide which type of behavior was followed for every fluid.
- Plot log shear stress against log shear rate curves for the graphs derived above. Estimate their power-law parameters, as done for Example 10.4. Write the specific power-law equations for each fluid.
- Present a complete written report, according to scientific report guidelines, either given by an instructor or consulted in the literature. Include graphs, charts, and calculations carried out, as well as discussion on own data. Compare the obtained power-law parameters from data based on literature survey from studies and investigations on the subject, and discuss the results accordingly.

10.4.2 Pretreatment of Suspensions: Coagulation and Flocculation

An important aspect of solid–liquid separation is the pretreatment or conditioning of the feed stream. Several treatments can be applied to the feed suspension including coagulation, flocculation, pH adjustment, freezing,

heating, or crystallization. Coagulation and flocculation could be considered the most widely used methods for conditioning the feed going into several solid–liquid separation techniques.

Coagulation and flocculation are often used interchangeably since both processes conduce to increase in particle size but there is, however, a subtle difference between these two terms. Coagulation is the process that brings together particles into contact to form agglomerates. The suspension is added with chemical hydrolyzing coagulants, such as alum, ferric salts, or lime, in order to produce agglomerates up to 1 mm in size. Some coagulants neutralize the surface charges on the primary particles, while others suppress the double layer or some even combine with particles through hydrogen bridging or complex formation. Flocculation, on the other hand, makes use of agents usually in the form of synthetic polyelectrolytes of high molecular weight, which interconnect and enmesh the particles into huge flocs up to 10 mm in size. Flocculation agents underwent rapid development in the past, which led to remarkable performance of many of the separation equipment in different industries. Flocculating agents are relatively expensive so that correct dosage is critical, not only in terms of cost but also because overdosing may inhibit the flocculation process by coating completely with the subsequent restabilization of the suspension. It can also cause operating problems, such as blinding of filter media or mud-balling and under-drain constriction in sand filters. Optimum dosage corresponds to the situation of having about one-half of the area of the particles covered with polymer.

In-line mixers or paddle-type flocculators are normally placed in front of some types of separation equipment, such as gravity clarifiers. The aim of using these mixers or flocculators is to assist the natural process of bringing together particles and polyelectrolytes by Brownian motion, increasing particle collision by orthokinetic flocculation through the motion of the fluid and velocity gradients in the flow. The rate of flocculation in clarifiers is also increased by recycling of the flocs, as to increase the rate of particle—particle collisions through the increase in solids concentration.

The specific type of floc needed depends on the particular solid–liquid separation that such flocs are supposed to enhance. Rotary vacuum filters require, for example, evenly sized, small, strong flocs, with capture of ultrafine particles within the floc to prevent cloth blinding. The flocs should not be subjected to sedimentation in the vat or breakage by the agitator, and should not be likely to cause localized air breakthrough, cake collapse, shrinkage, or cracking, in the dewatering stage. Large and loosely packed flocs are required in filtration operations preceded by a gravity selling step, such as belt presses. The resulting free-draining sediment can then be subjected to a controlled breakdown over a period of time conducing, ultimately, to a complete collapse of the cake due to its mechanical squeezing between the belts. In gravity thickening, large and relatively fragile flocs are necessary as to allow higher settling rates and fast collapse in the compression zone.

The optimum flocculant type and dosage depend on many factors, such as solid concentration, particle size distribution, surface chemistry, electrolyte content, and pH value. The flocculant selection and the optimization of its dosage therefore require extensive experimentation with only some general guidance as the ionic change or molecular weight taken as initial criterion. Anionic flocculants, for example, are known to perform well for coal slurries as the relatively high levels of calcium ions and hydrogen bonds, provide salt linkages for anchoring polymer anionic groups to the coal surface.

As the process of selecting flocculant type and dosage are a function on many factors, such as those mentioned above, the critical assessment of all of them for a particular choice would be a rather complex matter. A quick test to determine the optimum flocculant type and dosage consists on letting to settle samples of the suspension to be treated in one liter measuring cylinders, and add different doses of the potential flocculant to each of them. After an initial vigorous shaking for 1 min, gentle mixing is carried out for 5 min, and then the samples are left in repose until settling is visually evident. Two aliquots of the clarified liquid are extracted taking care of not disturbing the precipitate. The first aliquot is filtered through a Whatman[®] filter paper No. 2, while the second one is kept untreated. Both aliquots are assayed for turbidity, suspended solids, pH, and any other variable relevant to clarity of the decanted. When the results obtained from both aliquots are approximately similar, the optimum type and concentration of flocculant or coagulant have been determined.

10.4.3 Selection of Specific Techniques

The great number of applications of solid–liquid separation techniques in many industries, along with the diversity of equipment available for use in few or many of these applications, has made the process of choosing solid–liquid separation techniques and equipment a difficult one. Selecting a solid–liquid separator for a particular duty faces complexity because of the interaction of conflicting demands and criteria involved in the process. Numerous attempts have been made to devise selection charts, concise guidelines, and any other type of material to help in the decision of selecting processing equipment for a particular solid–liquid separation application. A selection system, and not just some form of superchart, would be required to take an account of the many factors implicit in the complicated task of separators selection in the materials processing industry.

Purchas (1981) has suggested a selection system consisting of three selection steps to produce a shorthand profile of the application from a combination of desk study to define the required duty, and some simple laboratory testing to characterize the separation properties of the solid–liquid mixture. The profile is expressed as a sort of code formed from a group of alphabetic letters (e.g., bdh-BEG-K), and is used in a fourth step to select suitable

TABLE 10.8Classification of Solid–Liquid Separation Equipment Based on Suitability for Duty and Slurry Type

	Suitable for Duty	Slurry Characteristics		
Equipment	Specification	Settling	Filtration	
Batch pressure filters with vertical elements	a, b, c D f, g, h, i	A, B D, E F, G	I, J	
Batch pressure filters with horizontal elements	b, c D g, h	A, B D, E F, G	J, K	
Continuous vacuum filter with bottom-fed drum	a, b, c E f, g, h, i	A, B D, E F, G, H	I, J, K	
Deep bed filters	a, b E F	A D F		
Filter press	a, b, c D f, g, h, i	A(B) D, E F, G, H	I, J	
Worm screen filtering centrifuge	A E G	C E H	K, L	
Pendulum filtering centrifuge	b, c D g, h	A, B, C D, E G, H	J, I, L	
Gravity settlers	a, b, c d, e f, g, h	B, C F F, G		
Hydrocyclones	a, b E f, g, h	B, C F F, G		
Microfiltration and ultrafiltration	a, b(c) d(e) F	A D F	I	
Screw and hydraulic press	a, b d, e G	A D, E H	I, J	
Tubular and bowl centrifuges	b(c) D f, g	A, B D, E F		
Disc centrifuge	a, b, c d, e f, g	A, B D, E F, G		

Source: Adapted from Purchas, D. B. 1981. Solid/Liquid Separation Technology. Croydon, UK: Uplands Press.

potentially suitable types of equipment from their corresponding application profiles, as listed in Table 10.8. The detailed steps are the following:

The first step is used to define the separation duty in terms of three criteria, namely scale, mode of operation, and objective. The mode of operation can be batch or continuous, while the objective may be referred to the clarification of the liquid phase or the recovery of the settling solids. If the objective is solid recovery, three further criteria are introduced, depending on whether the solids are to be discharged without further treatment, after washing or after drying on place. The procedure is summarized in Figure 10.18a, from which an appropriate letter (a–i) is assigned for each of the duty criteria.

The second step consists of running a simple test to identify the settling characteristic of the slurry. A sample is allowed to settle in a one liter

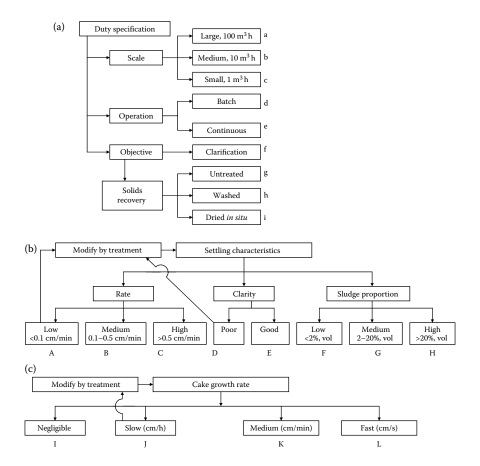


FIGURE 10.18

Sequence of steps in selecting a solid–liquid separation technique: (a) definition of basic characteristics of the process, (b) coding sludge settling characteristics, and (c) coding filtering characteristics.

measuring cylinder from about an hour and observations are made for the rate of settling, the clarity of the supernatant layer, and the depth of settling sludge relatively to the total sample depth. The observations are expressed as the corresponding code letters (A–H) giving in the summarized diagram shown in Figure 10.18b. When slurry is found to have poor settling characteristics, pretreatment adding coagulants or flocculants should be considered, and the process should be repeated.

The third step quantifies the filtration characteristic of the slurry from a second simple test, in which the speed of cake formation is roughly assessed. Using a laboratory Buchner funnel or any other simple suction filter, a small sample of about 200 mL of slurry is filtered and the time is recorded. The observed time and the cake thickness provide a basis to identify the filtration characteristic and assign a code letter (I–L) from those listed in Figure 10.18c. The possibility of pretreatment could be contemplated, again, if considered appropriate.

The fourth and final step is used to compare the separation duty profile derived from the three previous steps, with the equipment application profiles listed in Table 10.8 as to identify the most likely equipment for the solid–liquid separation duty.

10.4.3.1 Laboratory Exercise: Settling Tests to Select a Proper Technique

10.4.3.1.1 *Introduction*

Solid–liquid separations fall into two general categories: those involving clarification, and those dealing with cake formation and separation. Clarification is, generally, an operation to be carried out in equipment other than filters and involving, often, centrifugal forces. Cake formation or recovery of solids is a typical operation that may be performed in any type of filtering device. The objective of the experiment is to determine the suitability of a filter or a centrifugal device to carry out a separation task at laboratory scale, focusing on the important factor of the concentration of solids.

10.4.3.1.2 Instruments and Materials

The routine laboratory material and glassware to carry out gravimetric testing, such as analytical balance, oven, desiccator, and sample containers, would be sufficient for the experiments. There will be need of six to eight one liter measuring cylinders, also. Any type of laboratory centrifuge and a setup to perform filtering consisting of a filter flask, a Buchner funnel, and available vacuum, will complete the supplies needed. Suspensions of a fine powder, such as limestone, in water prepared at various concentrations with narrow range, but including both the dilute and concentrated suspension category, that is, 6%, 8%, 10%, 12%, and 14% solids, by volume.

10.4.3.1.3 Procedure

• Proceed as described in the previous section, by consulting first the chart in Figure 10.18a and assign the appropriate letters code.

As a first approach, consider clarification for the most dilute suspensions and recovery for the concentrated suspensions. The choice of batch or continuous operation may be arbitrary but, again, the most dilute suspensions tend to be treated in continuous equipment while the most concentrated tend to be directed to batch equipment.

- Carry out the first suggested in the previous section testing, that is, allow samples to settle in a one liter measuring cylinder from about an hour and observe the rate of settling, the clarity of the supernatant layer, and the depth of settling sludge relatively to the total sample depth. Assign the corresponding code letters using chart in Figure 10.18b.
- Take on the second suggested test by using a laboratory Buchner funnel to filter a small sample of about 200 mL of slurry. By using the filtration time and the cake thickness as criteria, identify the filtration characteristic and assign a code letter from those listed in Figure 10.18c.
- Identify the most likely equipment from the profiles listed in Table 10.8 for the solid–liquid separation duty.
- From all the choices made, use the results that may considered two options, one filtering option and another centrifugation-based option, and run tests in a laboratory centrifuge, and in the filtering setup. Compare the quality of the supernatant for both techniques, and choose the one that provides the lowest concentration of suspended solids. If the concentration of suspended solids is somewhat similar for both techniques, use discriminative criteria, such as the cost of the operation, the time in completing the task, and the labor-demanding feature of each technique. A rough estimate of the cost can be worked out by looking at the power of the electric motors of the centrifuge and the vacuum pump used in filtration, in relation to the time they operated to complete the task.

10.4.3.2 Calculations and Report

- Present all calculations, if done, for the discriminative criteria of cost, convenience, and so on, made in choosing one technique over another.
- Write a complete written report, according to scientific report guidelines, either given by an instructor or consulted in the literature. Include graphs, charts, and calculations carried out. Justify the choices made, as scientifically as possible.

10.4.4 Sedimentation

10.4.4.1 Introduction

Sedimentation can be defined as a unit operation to perform separation of a suspension into a supernatant clear fluid and dense slurry containing higher concentration of solids. In a more convenient manner, it should be established that the settling of solids of a suspension in sedimentation is due to gravity force, and industrial sedimentation can be described as the gravitational settling of solids suspended in liquids. The uses of sedimentation in industry fall into the categories of solid-liquid separations, solid-solid separations as in particle classification, and other operations such as mass transfer or washing. In solid-liquid separations the solids are removed from the liquid because any, or both, of the phases are more valuable separated or because they have to be separated before disposal. When the primary purpose is to produce the solids as highly concentrated slurry the process is called thickening, whereas if the purpose is to clarify the liquid the process is referred to as clarification. Usually, the feed concentration to a thickener is higher that that to a clarifier. Some types of equipment can accomplish both thickening and clarification in a single stage, provided that they are correctly designed and operated.

Sedimentation can be used for size separation, that is, classification of solids, being one of the simplest ways to remove coarse or dense solids from a feed suspension. Successive decantation in a batch system produces closely controlled size fractions of the product. Classification by sedimentation, however, does not produce a sharp separation.

Apart from the above-described applications, sedimentation is also used for other purposes. Relative motion of particles and liquid increases the mass-transfer coefficient. This motion is particularly useful for solvent extraction in immiscible liquid–liquid systems. Another important commercial application of sedimentation is in continuous countercurrent washing, where a series of continuous thickeners is used in a countercurrent mode in conjunction with returning of slurry, with the purpose of removing mother liquor or washing soluble substances from the solids.

As settling of solids in sedimentation is caused by the effect of gravity, difference of density between the solids and the suspended liquid is a necessary prerequisite. Also, it may be implied that the governing equations describing the process will be those derived in Section 10.1.2, that is, the Stokes law equation (Equation 10.11) and the related equations for sedimentation in the different settling regime regions (Equations 10.12 and 10.13). Predicting the settling behavior of particles within a liquid in real sedimentation system is, however, rather complicated because there are many factors involved in the process. Among these factors, the distance from the boundaries of the container and from other particles, can be mentioned as critical in affecting settling. Both of these mentioned factors are directly related to the concentrations of solids of the feed stream taken to a given

sedimentation unit. It has been suggested that if the relation of the diameter of particle to the diameter of the sedimentation vessel is over 1:200 or if the solids concentration by volume is less than 0.2%, any given particle is at sufficient distance from the boundaries of the vessel or from other particles so that its fall is not affected by them. In this case the process is called free settling. Contrastingly, when the motion of the particle is impeded by other particles, which will happen if the particles are near each other even they may not actually be colliding, the process is called hindered settling. The drag coefficient in hindered settling is greater than that in free settling.

10.4.4.2 Free Settling

As previously mentioned, the theory described in Section 10.1.2 including discussion on Equations 10.11 through 10.13 and the need of successive approximations in conjunction with Figure 10.3 to employ them properly, should be applied for calculation of industrial sedimentation within the free-settling regime. There are several cases in which such approximation is possible, for example, in measurement of particle size distribution by sedimentation or in particle classification. The approach, however, tends to be theoretical, since complete noninterference of particles settling within a liquid could be only possible for the case of these (the particles) being solitary. Experience has shown that in many real applications of industrial sedimentation correcting factors, mainly for suspension concentration, are needed in order to minimize calculation errors.

10.4.4.3 Hindered Settling

As described above, in concentrated suspensions particles get closer together and interfere with each other. If the particles are not disturbed uniformly, the overall effect is a net increase in settling velocity since the return flow caused by volume displacement predominates in particle-sparse regions. This is the well-known effect of cluster formation which is significant only in nearly monosized dispersions. With most practical widely dispersed suspensions clusters do not survive long enough to affect the settling behavior and, as the return flow is more uniformly distributed, the settling rate steadily declines with increasing concentration. This phenomenon of hindered settling and can be theoretically approached in three different manners: as a Stokes' law correction by introduction of a multiplying factor, by adopting effective fluid properties for the suspension different from these of the pure fluid, and by determination of bed expansion with a modified version of the well-known Carman–Kozeny equation. All the approaches can be shown to yield essentially identical results.

Svarovsky (1984) reviewed some important correlations accounting for the hindered settling effect and demonstrated that their differences are minimal.

According to this, the simple Richardson and Zaki equation is an obvious choice in practice. Such relation can be expressed as

$$\frac{u}{u_{+}} = (1 - C)^{4.65} \tag{10.58}$$

where u is the settling velocity at concentration C and $u_{\rm s}$ is the settling velocity of a single particle. The above relationship applies only to free, particulate separation unaffected by coagulation or flocculation and where all the particles are of uniform density.

The relationship above applies only to free, particulate separation unaffected by coagulation or flocculation, and where all particles are of uniform density.

10.4.4.4 Sedimentation Rate Stages

To analyze sedimentation in greater detail, the events occurring in a small-scale experiment conducted batch-wise as shown in Figure 10.19, can be observed. Particles in a narrow range will settle with about the same velocity. When this occurs, a demarcation line is observed between the supernatant clear liquid (zone A) and the slurry (zone B) as the process continues. The velocity at which this demarcation line descends through the column indicates the progress of the sedimentation process. The particles near the bottom of the cylinder pile up, forming a concentrated sludge (zone D) whose weight increases as the particles settle from zone B. As the upper interface approaches the sludge buildup on the bottom of the container, the slurry appears more uniform as a heavy sludge (zone D), the settling zone B disappears, and the process from then on consists only of the continuation of the slow compaction of the solids in zone D.

By measuring the interface height and solids concentrations in the dilute and concentrated suspensions, a graphic representation of the sedimentation

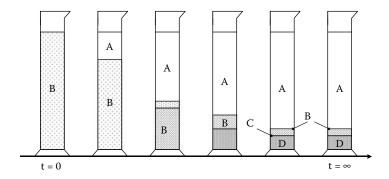


FIGURE 10.19

Progressive settling in a measuring cylinder: (A) clear liquid, (B) sludge at initial concentration, (C) transition zone, and (D) thick sludge at compression zone.

process can be prepared as shown in Figure 10.20. The plot shows the difference in interface height plotted against time, which is proportional to the rate of settling as well as to concentration.

As follows from an earlier discussion on spheres falling through a fluid medium, sedimentation is faster in liquids having low viscosities. Thus, sedimentation rates are higher at elevated temperatures. As also mentioned, an increase in the process rate may be achieved by increasing particle sizes through the use of coagulation or agglomeration, or by adding an electrolyte in the case of colloidal suspensions.

10.4.4.5 Operating Principles: Design and Selection Factors

Sedimentation equipment can be divided into batch-operated settling tanks and continuously operated thickeners or clarifiers. The operation of the former is very simple and their use has recently diminished. They are still used, however, when small quantities of liquids are to be treated. Most sedimentation processes operate in continuous units.

The conventional thickeners are constructed of steel in sizes less than 25 m diameter, or concrete in larger tanks that usually range up to 100 m diameter. The floor is often sloped toward the underflow discharge in the center. The tanks usually include a raking mechanism, which turns slowly around the center column in order to promote solids consolidation in the compression zone while aiding the underflow discharge. The rake arms are driven by fixed connections or dragged by cables or chains suspended from a drive arm that is rigidly connected to the drive mechanism. The rake arms are also connected to the bottom of the central column by a special arm hinge that allows both horizontal and vertical movements. This arrangement lifts the rakes automatically if the torque becomes excessive. The drive arm can be attached below the suspension level or, if scaling is a problem, above the basin. Figure 10.21 shows a diagram of a conventional circular section thickener.

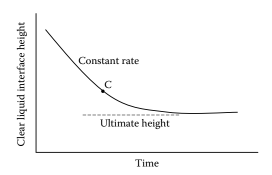


FIGURE 10.20 Sedimentation rate graph.

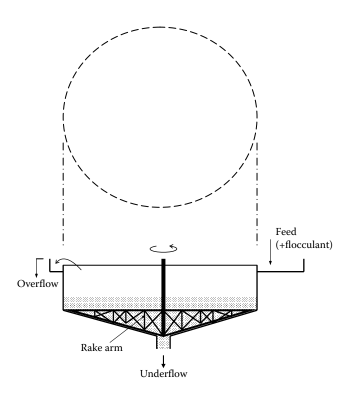


FIGURE 10.21 Diagram of circular section gravity thickener.

In order to increase capacity some modifications have been attempted in thickeners and clarifiers. Some examples of alternative designs in thickeners are shown in Figure 10.22.

In the circular-basin continuous thickener, illustrated in Figure 10.22a, after treatment with flocculant the feed stream enters the central feed well which dissipates the stream's kinetic energy and disperses gently into the thickener. The feed finds its height in the basin where its density matches the density of the inside suspension and spreads out at that level. Solids concentration increases downward in an operating thickener giving stability to the process. The settling solids and some liquid move downward. The amount of the latter depends on the underflow withdrawal rate. Most of the liquid moves upwards into the overflow, and is collected in a trough around the periphery of the basin. A typical thickener has three operating layers: clarification, zone settling, and compression. Frequently, the feed is contained in the zone settling layer that theoretically eliminates the need for the clarification zone because the particles would not escape through the interface. However, in practice, the clarification zone provides a buffer for fluctuations in the feed and the sludge level. The most important dimensions of a thickener

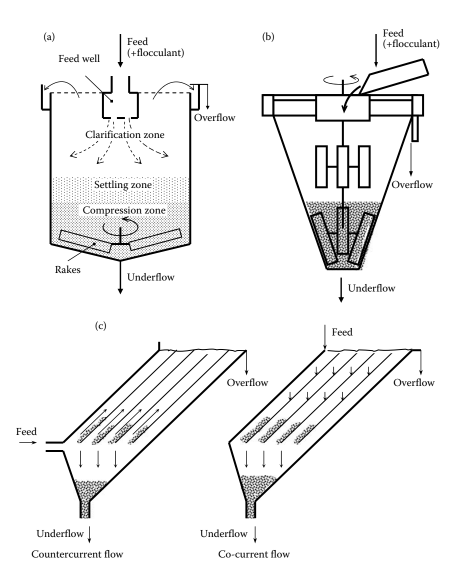


FIGURE 10.22

Alternative designs for gravity thickeners: (a) circular-basin continuous thickener, (b) deep-cone thickener, and (c) lamella thickener.

are pool area and depth. The pool area is chosen to be the largest of the three layer requirements. In most cases, only the zone settling and compression layer requirements need to be considered. However, if the clarity of the overflow is critical, the clarification zone may need the largest area. As to the pool depth, only the compression layer has a depth requirement because the

concentration of the solids in the underflow is largely determined by the time detention and, sometimes, by the static pressure. Thickness of the other two layers is governed only by practical considerations. The deep-cone thickener (Figure 10.22b), based on the deep-cone vessel used for the processing of coal and metallurgical ores, is equipped with a slow-turning stirring mechanism, which enhances flocculation in the upper part and acts as a rake in the lower section. The unit is used for densification of froth-flotation tailings at overflow rates from 6.5-10 m/h to a final discharge containing 25-35% moisture by weight. Other commercial deep-cone thickeners are of particular advantage where the final underflow density is increased by the large static head above the discharge point, for example, with flocculated clays. Another development is the Swedish Lamella thickener shown in Figure 10.22c. It consists of a number of inclined plates stacked closely together. In the countercurrent design, the flocculated feed enters the stack from a side feed box. The flow moves upward between the plates while the solids settle onto the plate surfaces and slide down into the sludge hopper underneath where they are further consolidating by vibration or raking. In theory, the effective settling area is the sum of the horizontal projected areas of all plates. In practice, however, only about 50% of the area is utilized. When treating sticky sludges, the whole lamella pack can be vibrated intermittently continuously to assist the sliding motion of the solids down the plates. In some instances, the plates are corrugated instead of flat, or they are replaced by tube bundles becoming what is known as a tube settler.

Thickeners are designed by the traditional Coe and Clevenger, or Talmage and Fitch procedures (Fitch, 1975), which use batch-settling data to evaluate the whole concentration range right up to the underflow concentration, even though a batch-settling test cannot possibly simulate the continuous-thickener process. The design procedures are based on plotting total flux versus concentration, as shown in Figure 10.23a. The concentration of the solids continuously increases from the feed value $C_{\rm f}$, to the underflow concentration $C_{\rm u}$. The total flux plot goes through a minimum (the critical flux $G_{\rm c}$) on which the design area A is based, according to a mass balance, on the following relation:

$$A = \frac{QC_{\rm f}}{G_{\rm c}} \tag{10.59}$$

where Q is the mass flow rate of particles by volume.

The total flux plot depends not only on the settling characteristics of the solids, but also on the selected underflow concentration because total flux includes the transport flux, or dead flux, contributed by movement of particles downward as underflow is withdrawn. The plot can be made independent of $C_{\rm u}$ by subtracting the transport flux contribution from $G_{\rm c'}$ and plotting the resultant settling flux $G_{\rm s'}$, as done in Figure 10.23b. A line drawn tangentially

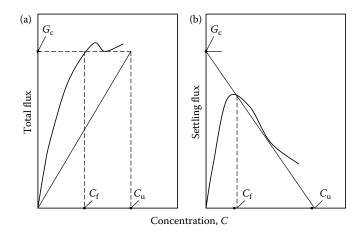


FIGURE 10.23

Design curves for thickeners: (a) total flux versus concentration, and (b) settling flux versus concentration.

from the selected C_u value to the undersize of the settling flux curve will intercept the *y*-axis at the same G_c value.

The settling flux curve can be constructed directly from batch settling tests carried out at several different concentrations in the zone settling regime (Coe and Clevenger method), or from just one test curve (Talmage and Fitch method) by converting the u = f(C) data into settling flux by the formula:

$$G_{\rm s} = uC \tag{10.60}$$

The Coe and Clevenger test overestimates the critical flux leading to underdesign of thickener area, while the Talmage and Fitch procedure underestimates it leading to overdesign. The latter procedure is less laborious because it only requires one settling test.

The thickener area can also be determined using a mass balance of the whole sedimentation process. Referring to the diagram in Figure 10.24, let *F*

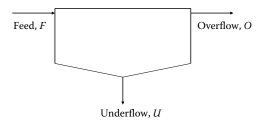


FIGURE 10.24
Mass balance on sedimentation unit.

be the mass flow rate of solids in the feed, U the mass flow rate of solids in the underflow, O the mass flow rate of solids in the overflow, X_F the fraction of solids in the feed, X_U the fraction of solids in the underflow, $1 - X_F$ the fraction of liquid in the feed, and $1 - X_U$ the fraction of liquid in the underflow.

The mass balances of liquid and solids in the process can be represented by

$$F(1 - X_{\rm F}) = O + (1 - X_{\rm U}) \tag{10.61}$$

$$F(X_{\rm F}) = U(X_{\rm U}) \tag{10.62}$$

It can be demonstrated that

$$O = \frac{F(X_{\rm U} - X_{\rm F})}{X_{\rm U}} \tag{10.63}$$

For the limiting conditions of producing a clear overflow, the terminal settling velocity of the particles u_t should be equal to the ascending velocity of the liquid u_1 , so that

$$u_{\rm t} = u_{\uparrow} = \frac{\text{Flowrate (volumetric)}}{\text{Area}} = \frac{O}{\rho A}$$
 (10.64)

Substituting Equation 10.64 into Equation 10.63, transposing for A and calling it minimum area A_{\min} :

$$A_{\min} = \frac{F(X_{\mathrm{U}} - X_{\mathrm{F}})}{X_{\mathrm{U}} u_{\mathrm{t}} \rho} \tag{10.65}$$

With regard to clarifiers, the conventional one-pass unit employs horizontal flow in circular or rectangular vessels. Figure 10.25 shows a schematic diagram of a rectangular basin with feed at one end and overflow at the other. The feed can be preflocculated in an orthokinetic flocculator that may form part of the unit. Settled solids are pushed to a discharge trench by paddles or blades on a chain mechanism or suspended from a traveling bridge. Circular basin clarifiers are most commonly fed through a centrally located feed well; the overflow is led into a trough around the periphery of the basin. The bottom gently slopes to the center and the settled solids are pushed down the slope by a number of motor-driven scraper blades that revolve slowly around a vertical center shaft. Like thickeners, circular clarifiers can be stacked in multitray arrangements to save space.

Clarifier performance depends on area, which is determined by the flocculation nature of the feed suspension. When the overflow clarity is

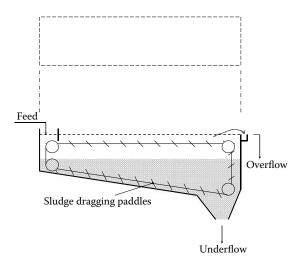


FIGURE 10.25 Diagram of rectangular-basin clarifier.

independent of overflow rate and depends only on detention time, the required time is determined by simple laboratory testing of residual solid concentrations in the supernatant versus detention time under the conditions of mild shear. This determination is sometimes called the second-order test procedure because the flocculation process follows a second-order reaction rate. In most cases, clarifying performance depends on detention time and overflow rate. Tests are conducted in a vertical tube that is as long as the expected depth of the clarifier, under the ideal assumption that a vertical element of a suspension, which has been clarified, maintains its shape as it moves across the tank.

If the suspension is nonflocculant, or if flocculation takes place prior to settling, the overflow clarity is independent of detention time and depends only on the overflow rate *Q* according to the relation:

$$Q_{\text{max}} = A(u_{\text{t}}) \tag{10.66}$$

in this case the settling velocity u_t is expressed by

$$u_{t} = \frac{H}{t} \tag{10.67}$$

where u_t is the maximum particle settling velocity that yields a satisfactory clarity in a simple laboratory sedimentation test, H is the height of the laboratory container, and t is the time in which the supernatant liquid becomes clear.

For nonflocculant suspensions, gravity clarifiers are sometimes used as for solids classification. The theoretical grade efficiency curve G(x) of the clarifier can be predicted, assuming laminar flow and no end effects, by

$$G(x) = \frac{A(u_{\rm t})}{Q} \tag{10.68}$$

where u_t is a function of particle size and can be found from Equation 10.11 if Stokes law applies.

Example 10.5

A 25% solids concentrated suspension contains particles of approximate spherical shape of an insoluble food powder. The mean size of the particles is 50 μm and their density is 2002 kg/m³. Calculate the settling velocity of such particles in water at room temperature.

SOLUTION

Due to the size and density of particles, a reasonably assumption would be settling within the Stokes law region. Properties of water can be consulted in Appendix 2. Using, thus, Equation 10.11, the terminal settling velocity is

$$u_{\rm t} = \frac{(5 \times 10^{-5})^2 \text{ m}^2 (1002) \text{ kg/m}^3 (9.81) \text{ m/s}^2}{18(0.001) \text{ kg/m s}} = 0.00136 \text{ m/s}$$

Verifying the applicability of the chosen sedimentation region, by calculating the particle Reynolds number:

$$Re_{\rm p} \, = \frac{(5 \times 10^{-5}) \, m \, (1000) \, kg/m^3 (0.00136) \, m/s}{0.001 \, kg/m \, s} \, = \, 0.06$$

The approximation was correct and the settling time is valid. Finally, due to the high concentration of solids, correction due to this factor is in order. Transposing for the corrected settling velocity from Equation 10.58 and substituting values, the real settling velocity is

$$u = u_{\rm t}(1 - C)^{4.65} = (0.00136)(1 - 0.25)^{4.65} = 0.000357$$
 m/s

10.4.4.6 Laboratory Exercise: Acceleration of Settling Rates by Coagulants

10.4.4.6.1 Introduction

Coagulation and flocculation are used to increase the effective particle size, thereby improving sedimentation settling rates. Coagulation produces

particles up to 1 mm in diameter, while flocculation renders flocs up to 1 cm in size. The objective of the exercise is to verify the advantage of adding a coagulant to a concentrated suspension, by comparing settling rates with and without the coagulant addition.

10.4.4.6.2 Instruments and Materials

Five or six 1-L measuring cylinders are required. A concentrated suspension of a fine powder, for example, limestone or starch in water, is also necessary. Any known coagulant agent, such as $Al_2(SO_4)_3$ or $Fe_2(SO_4)_3$.

10.4.4.6.3 Procedure

- Shake the suspension well and allow it to settle in the one-liter measuring cylinder.
- Record the clear liquid interfase height related to time, until it stabilizes. Make replicates to give statistical meaning to the exercise.
- Repeat the procedure adding Al₂(SO₄)₃ or Fe₂(SO₄)₃ as the coagulant agent.

10.4.4.6.4 Calculations and Report

- Construct graphs of rate of sedimentation (Figure 10.20) for both sedimentation processes.
- Locate the point of inflexion (critical point) for each graph.
- If critical points appear distant in both graphs, run a statistical test to determine whether they are significantly different.
- Write a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature. Include graphs, charts, and calculations carried out.

10.4.5 Centrifugation

10.4.5.1 Introduction: Centrifugal Techniques

As previously discussed, apart from gravity, centrifugal force can also be used to separate solids in suspension. Centrifugal sedimentation is a technique in which the suspension is subjected to a centrifugal field, to cause the solid particles to move radially apart from the center of rotation and lead to separation. A significant density difference between solid and liquid is a necessary prerequisite in sedimentation of any kind. Centrifugal sedimentation can be carried out in two types of equipment: rotated wall devices and fixed wall devices. Rotating wall devices are also known as centrifuges. The most popular fixed wall device is the hydrocyclone. The term "centrifugation" has been traditionally used to imply the operation of centrifugal sedimentation in rotating wall devices.

10.4.5.2 Applications of Centrifugation

Centrifugation presents varied and diverse applications in different processing industries. Many types of separations can be carried out through centrifugation, as long as a significant density difference between solid and liquid exists. The most common applications of centrifugation in processing industries are: (a) liquid–liquid separations, (b) centrifugal clarification, (c) sludge separation, and (d) centrifugal filtration.

Within the context of particle technology, the most relevant application of centrifugation is centrifugal clarification since it follows theoretical principles of dynamics of particles submerged in fluids as discussed in this book. Those principles are used to describe centrifugal clarification as a unit operation.

10.4.5.3 Centrifugal Clarification

Centrifugal clarification is the term used to describe the removal of small quantities of insoluble solids from a fluid by centrifugal means. If a dilute suspension containing solids with a greater density than the liquid is fed to a rotating cylindrical bowl, the solids will move toward the bowl wall. If an outlet is provided for the liquid near the center of rotation, then those particles of solid which reach the bowl wall will remain in the bowl. Those particles that do not reach the bowl wall will be carried out in the liquid. The fraction remaining in the bowl and the fraction passing out in the liquid will be controlled by the feed rate, that is, the dwell time, in the bowl. If a solid particle of diameter *x* moves radially in a liquid within a rotating bowl, at its terminal velocity under laminar flow conditions the radial velocity of the particle will be represented by Equation 10.31. The time required for a particle to travel an elemental radial distance, dR, is

$$dt = \frac{dR}{v_r} = \frac{18\mu}{\omega^2(\rho_s - \rho)x^2} \cdot \frac{dR}{R}$$
 (10.69)

Assuming that half of all those particles present in the feed with a particular diameter x_c are removed during their transit through the bowl, those particles with diameters greater than x_c will be mostly removed from the liquid, whereas those particles with diameters smaller than x_c will be likely to remain in the liquid. In this context x_c as here defined is known as the "cut-point" or "critical" diameter.

If clarification takes place in a simple cylindrical centrifuge with cross section as shown in Figure 10.26, all particles of diameter x_c contained in the outer half of the cross-sectional area of the ring of liquid will reach the bowl wall and will be removed from the liquid. The maximum distance that a particle in this zone has to travel to reach the bowl wall is $R_2 - [(R_1^2 + R_2^2)/2]^{1/2}$,

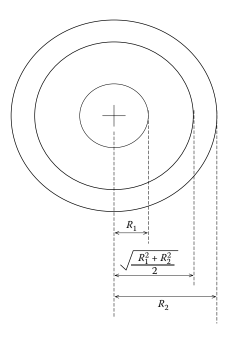


FIGURE 10.26

Schematic diagram of radial separation zones in the cross section of a centrifugal clarifier.

as indicated in Figure 10.26. The time required for a particle of diameter x_c to travel this distance is

$$t = \frac{18\mu}{\omega^2(\rho_s - \rho)x_c^2} \int_{[(R_1^2 + R_2^2)/2]^{1/2}}^{R_2} \frac{dR}{R}$$
 (10.70)

Integrating Equation 10.70 and substituting limits the following relation is obtained:

$$t = \frac{18\mu \ln\left(R_2/\left[(R_1^2 + R_2^2)/2\right]^{1/2}\right)}{\omega^2(\rho_s - \rho)x_c^2}$$
(10.71)

The minimum residence time for a particle in the bowl is V/Q, where V is the volume of liquid held in the bowl at any time and Q is the volumetric flow rate of liquid through the bowl. Thus, for a particle of diameter x_c to be separated out

$$\frac{V}{Q} = \frac{18\mu \ln\left(R_2/\left[(R_1^2 + R_2^2)/2\right]^{1/2}\right)}{\omega^2(\rho_s - \rho)x_c^2}$$
(10.72)

Equation 10.72 may be written in the form:

$$Q = 2 \left[\frac{g(\rho_{s} - \rho)x_{c}^{2}}{18\mu} \right] \cdot \left[\frac{\omega^{2}V}{2g\ln\left(R_{2}/\left[(R_{1}^{2} + R_{2}^{2})/2\right]^{1/2}\right)} \right]$$
(10.73)

The first term of Equation 10.73 includes the previously discussed expression for Stokes' law, that is, Equation 10.11. Thus, another way of expressing Equation 10.73 is as follows:

$$Q = 2u_{\rm g}\Sigma \tag{10.74}$$

where $u_{\rm g}$ is the terminal settling velocity of a particle of diameter $x_{\rm c}$ in a gravitational field and Σ is a characteristic parameter of any given centrifuge; equivalent to the area of a gravity settling tank with similar settling characteristics to the centrifuge, that is, one which will remove half of all particles of diameter $x_{\rm c}$. Different values of Σ are given in the literature (McCabe et al., 2005). For a simple cylindrical bowl centrifuge:

$$\Sigma \approx \frac{\pi \omega^2 b (3R_2^2 + R_1^2)}{2g} \tag{10.75}$$

where *b* is the height of the bowl. Also, for a disc-bowl centrifuge:

$$\Sigma = \frac{2\pi\omega^2(S-1)(R_x^3 - R_y^3)}{3g\tan\Omega}$$
 (10.76)

where *S* is the number of discs in stack, R_x and R_y are the outer and inner radii of stack, respectively, and Ω is the conical half-angle of discs (Trowbridge, 1962).

10.4.5.4 Centrifugal Filtration

The term describes the separation of solids from a liquid by filtration for the case of the filtrate flow being induced by centrifugal means. The general principle of the filtering centrifugal is shown in Figure 10.27. As can be observed, a suspension is fed into a rotating bowl with a perforated wall, which is lined with a suitable filter medium. The solids are forced onto the bowl wall giving rise to a filter cake through which the filtrate passes, under the influence of centrifugal force. The filtrate makes its pass through the filter medium and perforated wall also and, finally, into an outer casing and out a discharge port. Wash liquid may be sprayed through the solids to remove

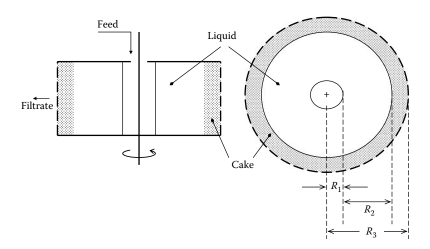


FIGURE 10.27 Diagram of simple centrifugal filter.

soluble material and, then, the cake is spun as dry as possible by rotating the bowl at a higher speed than during the operating run. The motor is shut off and, with the bowl slowly turning, the solids are discharged by scraping them out with an unloader knife, which peels the cake from the filter medium and drops it through an opening in the bowl floor. The filter medium is rinsed clean, or substituted if needed, the motor turned on, and the cycle repeated.

Direct comparison of pressure filtration and centrifugal filtration reveals certain differences. For example, in the latter both the centrifugal force and the filtering area increase with increase in radius. Centrifugal force acts on the filtrate passing through the cake and on the cake itself, aiding to the hydraulic pressure head. For a relatively simple arrangement of a centrifugal filter, assuming incompressible cake and neglecting kinetic energy changes in the filtrate, the rate of flow filtrate through it may be expressed by

$$Q = \frac{\rho \omega^2 (R_3^2 - R_1^2)}{2\mu \left[\alpha M_c / A_a A_l + \frac{R_m}{A_m} \right]}$$
(10.77)

where ρ is the density of the liquid, ω is the rotation speed, R_3 is the radius of the inner bowl surface, R_1 is the radius of the inner surface of the liquid ring, μ is the filtrate viscosity, α is the specific cake resistance, M_c is the mass of the solid cake in the bowl, A_a is the arithmetic mean of the cake area, A_1 is the logarithmic mean of the cake area, R_m is the resistance of the filter medium, and A_m is the area of the filter medium.

Equation 10.77 is applicable only for cakes of uniform thickness. In many practical cases cakes are thicker near the bottom than at the top of bowl. Such systems will give a filtration rate 5–20% greater than a cake of constant thickness of similar volume and permeability (Green and Perry, 2008).

Example 10.6

A viscous suspension contains particles of a solid of 1461 kg/m³ density, and is to be clarified by centrifugation. The liquid has a density of 801 kg/m³ and a viscosity of 100 cp. A clarifying centrifuge with bowl 44.5 diameter and 197 mm height is used. During operation an annular air core of 14.23 mm diameter is formed. Calculate the cut point for a capacity of 169.92 L/h, when the centrifuge runs at 23,000 rpm speed.

SOLUTION

To facilitate calculation, it is recommended to transform all the data to SI units. The viscosity is 0.1 kg/m s in SI units. Pertaining geometrical features, according to diagram in Figure 10.26: $R_2 = 0.0225$ m, $R_1 = 0.00761$ m, and b = 0.197 m. Finally, the capacity is 4.7245×10^{-5} m³/s, and the rotation speed is 2410 rad/s (2410 L/s), from the relation $\omega = (2\pi N)/60$, where N is rotational speed in rpm. With all the data in SI units, from Equation 10.75, the value of Σ is

$$\Sigma = \frac{3.14(2410)^2(1/s^2)(0.197)\,m\Big[3(0.02225)^2 + (0.00716)^2\Big]\,m^2}{2(9.81)\,m/s^2} = 281.5\,m^2$$

Finally, transposing for the cut point x_c from Equation 10.74 and solving:

$$x_c = \sqrt{\frac{18(0.1) \text{ kg/m s}(4.725 \times 10^{-5}) \text{ m}^3/\text{s}}{2(9.81) \text{ m/s}^2(1461 - 801) \text{ kg/m}^3}} = 4.83 \times 10^{-6} \text{ m}$$

Thus, the cut point is 4.8 µm, approximately.

10.4.5.5 Fixed Wall Devices: Hydrocyclones

The use of hydrocyclones is another possibility to separate suspended solids from a liquid taking advantage of the centrifugal force. The operating principle of a hydrocyclone is similar to that of a cyclone (Figure 10.7), differing only in the geometrical proportions due to the difference in magnitudes of properties between gases and liquids. Since liquids possess densities and viscosities proportionally much higher than gases, in hydrocyclones Δp is much narrower than in cyclones. Also, for the same reason of properties of phases, the viscous forces opposing settling of particles within a liquid are much greater than those within a gas. Hydrocyclone units have, therefore, proportionally much smaller diameters than cyclones. In such a manner the liquid flow, proportionally more dense and viscous than a gas flow, would

give origin to a vortex within the hydrocyclone with centrifugal force sufficient to separate solid particles.

Similar to cyclones and centrifuges, hydrocyclones may be evaluated in terms of separation efficiency by means of the cut size or cut point (x_c or x_{50}). The criteria to define and interpret efficiency, along with definitions and usefulness of the cut point concept, have been discussed in Section 10.2. Numerous analytical expressions to calculate the cut point for hydrocyclones have been suggested (Ortega-Rivas, 1989), but the most convenient way of evaluating it is, arguably, using the grade efficiency curve also discussed in Section 10.2. Due to the effect of concentration of suspensions, than will be normally much higher for hydrocyclones than for cyclones, the grade efficiency curve should be derived taking into account the dead flux effect as to obtain reduced grade efficiency curves (Section 10.2.4) like the example given in Figure 10.5 from calculations using Equation 10.29.

Since hydrocyclones do not have any rotating parts and the vortex action to produce centrifugal force is obtained by pumping the feed suspension tangentially into the cono-cylindrical body, the literature is full of studies of the effects of the relative geometric proportions on pressure drop or capacity and separation efficiency. Using this information hydrocyclone geometry could be selected to obtain an optimum performance in terms of cut size. In this sense, possibly the best opportunity to predict hydrocyclone performance is the use of a dimensionless scale-up model well described elsewhere (Ortega-Rivas and Svarovsky, 1993). Three dimensionless groups can be used to describe hydrocyclone operation and performance: the Euler number Eu, the Reynolds number Re, and the Stokes number Stk₅₀. For best application of the relationships among dimensionless groups, certain proportions must be unchanged. Such proportions are generally reported as a function of the diameter of the hydrocyclone. There are several different standard hydrocyclone designs in which proportions remain the same regardless of size. One of the most efficient designs for separation is called the Rietema (1961) cyclone, whose proportions are illustrated in Figure 10.28.

The Euler number, which is a pressure loss factor, is defined as the limit of the maximum characteristic velocity v obtained by a certain pressure drop ΔP across the hydrocyclone. It can be expressed as

$$Eu = \frac{2\Delta P}{\rho v^2} \tag{10.78}$$

where ρ is the liquid density and v is the superficial velocity in the hydrocyclone body.

The Reynolds number defines flow features of the system and, in the case of hydrocyclones, the characteristic dimension may be taken as the cyclone body diameter D_c :

$$Re = \frac{D_c v \rho}{\mu} \tag{10.79}$$

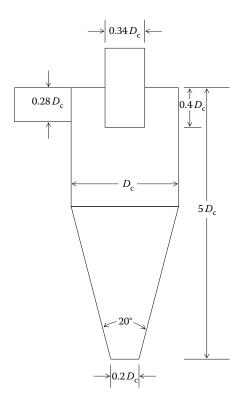


FIGURE 10.28 Rietema's optimum standard geometry hydrocyclone.

where ρ is the liquid density and μ is the average viscosity.

The Stokes number may be derived from basic fluid mechanics theory and is defined as follows:

$$Stk_{50} = \frac{x_{50}^2(\rho_s - \rho)v}{18\mu D_c}$$
 (10.80)

where x_{50} is the cut size, ρ_s is the solids density, and D_c is the hydrocyclone diameter.

It can be demonstrated (Svarovsky, 1984) that, similar to the case of cyclones, the most appropriate characteristic velocity to describe hydrocyclone operation is the superficial velocity in the hydrocyclone body described by Equation 10.38.

Owing to the effect of the suspension concentration that has an influence in the dead flux effect, sufficiently discussed in previous sections, the concentration as a fraction C and the underflow-to-throughput ratio R_f are

included in the dimensionless scale-up of hydrocyclone operation as additional dimensionless groups. The underflow-to-throughput is defined as

$$R_{\rm f} = \frac{U}{Q} \tag{10.81}$$

where *U* is the underflow rate and *Q* is the feed rate.

The dimensional analysis gives basic relationships between the above-mentioned dimensionless groups, which are useful to describe hydrocyclone operation and are the basis for scale-up calculations aimed at adapting results from laboratory experimentation to an industrial scale for a number of processes in diverse industries. For example, for feed suspensions up to 10% by volume Medronho and Svarovsky (1984) proposed the following relations, for hydrocyclones following Rietema's optimum proportions and treating inert solids suspensions:

$$Stk_{50}(r)Eu = 0.047 \left[ln \left(\frac{1}{R_f} \right) \right]^{0.74} e^{(8.96C)}$$
 (10.82)

Eu = 71(Re)^{-0.116}
$$\left(\frac{D_i}{D_c}\right)^{-1.3}$$
 e^(2.12C) (10.83)

$$R_{\rm f} = 1218 \left(\frac{D_{\rm u}}{D_{\rm c}}\right)^{-4.75} ({\rm Eu})^{-0.30}$$
 (10.84)

where $D_{\rm i}$, $D_{\rm c'}$ and $D_{\rm u}$ are the inlet, body, and underflow diameters of the hydrocyclone, respectively. The term (r) accompanying Stk₅₀ is included to indicate that the "reduced" grade efficiency was used to determine the "reduced" cut size x_{50} , in order to account for the dead flux effect.

For concentrations higher than 10% by volume, many practical slurries show non-Newtonian behavior and it can be shown (Ortega-Rivas and Svarovsky, 1993) that Reynolds and Stokes numbers can be reexpressed to consider such behavior. The correlations derived under this mentioned consideration are the following:

$$Stk_{50}^{*}(r)Eu = 0.006 \left[ln \left(\frac{1}{R_f} \right) \right]^{2.37} e^{(6.84C)}$$
 (10.85)

$$Eu = 1686(Re^*)^{-0.035} e^{(-3.39C)}$$
 (10.86)

$$R_{\rm f} = 32.8 \left(\frac{D_{\rm u}}{D_{\rm c}}\right)^{1.53} ({\rm Re}^*)^{-0.34} e^{(3.70C)}$$
(10.87)

where Stk_{50}^* and Re^* are the "generalized" Stokes and Reynolds numbers, meaning that they include the parameters of characterization of non-Newtonian suspensions, that is, the fluid consistency index K' and the flow behavior index n, instead of the medium viscosity (Ortega-Rivas and Svarovsky, 1993). The term generalized is used to imply that, for Newtonian suspensions, Stokes and Reynolds numbers above would reduce to the common forms normally found in the literature. The term (r) accompanies, again, $(Stk_{50})^*$ for the same reasons already given.

Example 10.7

Determine the cut size obtained when treating 1 L/s of a suspension of inert solids in water, with concentration of 5% by volume, in a Rietema's standard geometry hydrocyclone 1 in. in diameter and 5 mm in underflow opening. The solids density is 2150 kg/m³ and the available pressure drop of operation is 2×10^5 Pa.

SOLUTION

The low-suspension concentration will allow use of the properties of the suspending medium in calculations. Reasonable values for density and viscosity of water at approximate room temperature are 1000 kg/m^3 and 0.001 kg/m s, respectively. The feed rate F(Q) is $0.001 \text{ m}^3/\text{s}$ and the pressure drop $2 \times 10^5 \text{kg/s}^2$ m, both expressed in fundamental SI units. The characteristic velocity within the hydrocyclone can be calculated from Equation 10.38:

$$v = \frac{4(0.001) \text{m}^3/\text{s}}{3.1416(0.0254)^2 \text{m}^2} = 1.97 \text{m/s}$$

Using the above-calculated velocity, Eu can be calculated from Equation 10.78, that is,

$$Eu = \frac{2(2 \times 10^5) \text{ kg/s}^2 \text{ m}}{(1000) \text{kg/m}^3 (1.97) \text{ m/s}} = 102.7$$

once having the value of Eu, since the suspension concentration is low, Equation 10.84 can be used to calculate $R_{\rm f}$ as

$$R_{\rm f} = 1218 \left(\frac{0.005 \,\text{m}}{0.0254 \,\text{m}} \right)^{4.75} (102.7)^{-0.3} = 0.1346$$

the calculated values of Eu and R_i , can then be substituted into Equation 10.82 to transpose for the Stokes number and calculating it, that is,

$$Stk_{50}(r) = \frac{\left[\ln(1/0.1346)\right]^{0.74} \exp\left[8.96(0.05)\right]}{102.7} = 0.0019$$

Finally, from the original definition of the Stokes number given in Equation 10.80, and transposing for x_{50} :

$$x_{50} = \left(\frac{(0.0019)(18)(0.001)\text{kg/ms} (0.0254)\text{m}}{(2150 - 1000)\text{kg/m}^3(1.97)\text{m/s}}\right) = 1.55 \times 10^5 \text{ m}$$

The cut point is: $x_{50} \sim 15 \, \mu \text{m}$.

10.4.5.6 Centrifugation Equipment

Generally, the main types of equipment to carry out centrifugation duties in industry coincide in a way with the classification of centrifugation modes described in this book. Regardless of the criteria used to categorize centrifugation equipment, it is reasonably to classify centrifuges into: centrifugal clarifiers (tubular and disc bowl) used not only for clarification but for liquid–liquid separation, sludge or decanting centrifuges, and filtering centrifuges. Regarding hydrocyclones, apart from Rietema's geometry already defined there are some other designs available like the Bradley's hydrocyclone (Bradley, 1965), but geometric proportions really do not vary much. The alternatives in terms of hydrocyclone manufacturing relate to arrangements in series to provide the best possible efficiency at the highest attainable capacity.

Tubular and disc-bowl centrifuges are designs normally used for clarification. Figure 10.29 illustrates a tubular-bowl centrifuge consisting on a long and narrow cylindrical bowl rotating at high speed in an outer stationary casing. The feed is introduced through a stationary pipe at the bottom of the bowl, and quickly accelerated to the bowl speed by means of vanes or baffles. The two liquids are removed from the annular layers formed through a circular weir system, as shown in Figure 10.29. The control over the neutral zone radius is exercised by fitting rings with different internal diameters to the dense phase outlet. The solids capacity of a tubular bowl machine is seldom more than 2–5 kg and for economical operation the solids content of the feed should normally not exceed about 1% by weight. Disc-bowl centrifuges (Figure 10.30) comprise a relatively shallow, wide cylindrical bowl, rotating at moderate speed in a stationary casing. The bowl contains closely spaced metal cones, or discs, which rotate with the bowl and are placed one above the other with fixed clearance between them. The discs have one or more sets of matching holes in order to form channels through which the feed material flows. Under the influence of centrifugal force the dense phase, traveling toward the bowl wall, streams down the undersides of the discs while the light phase, displaced toward the center, flows over the upper faces of the discs. The liquids are thus divided into thin layers and the distance any drop of the liquid must travel to get caught up the appropriate outgoing stream is very small. The separated

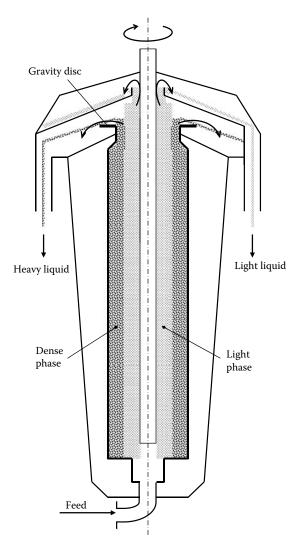


FIGURE 10.29 Diagram of tubular-bowl centrifuge.

liquids are removed by a weir system and, like in the case of tubular centrifuges, different gravity discs on the dense phase outlet need to be used to control the neutral zone position. Disc-bowl centrifuges have solids capacities in the range of 2–20 kg and are usually only suited to clarifying feeds with less than a few percent by weight of solids. Both, tubular and disc-bowl centrifuges, are useful for removing small traces of solids to produce clear liquids, especially if the solids are gelatinous in nature and unsuited to filtration.

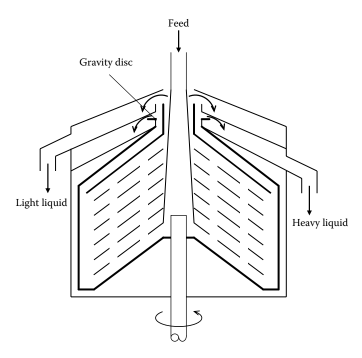


FIGURE 10.30 Diagram of disc-bowl centrifuge.

The nozzle-discharge (self-cleaning) and the valve-discharge (self-opening) centrifuges are sophisticated pieces of equipment designed to give continuity to the solid-liquid centrifugation process. A diagram of a nozzle-discharge centrifuge is given in Figure 10.31. As can be seen this centrifuges have a biconical bowl with a number of holes (approximately 0.3 cm in diameter) spaced around it at its largest diameter. The solids removed from the liquid are continuously discharged, in the form of thick slurry, into an outer casing. Feeds containing up to 25% solids can be handled in this type of clarifiers. The valve-discharge centrifuge has valves fitted in the solids discharge ports in the bowl which can be opened at desired intervals to get rid of the solids. Such valves may be controlled by timers or, alternatively, be automatically opened by hydrostatic control.

In order to handle concentrated suspensions, basket centrifuges, screening centrifuges, and decanter centrifuges, among others, have been employed. Decanter or scroll centrifuges are conventionally used for the continuous separation of solids from liquids. They consist of a solid horizontal bowl tapered at one end and enclosed within a cylinder (Figure 10.32). Thick slurry is pumped into the bowl along the central axis, and it is projected against the cylinder wall near the tapered section. A screw, revolving at a higher speed close to the bowl wall, conveys the solids along the tapered surface to the

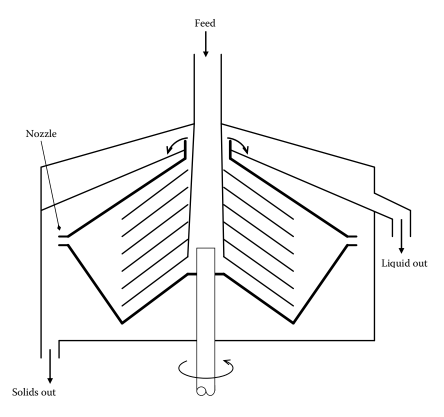


FIGURE 10.31 Diagram of nozzle-discharge centrifuge.

point of discharge and allows the inner layer of liquid to be drained out through a second discharge channel.

As previously mentioned, industrial applications of hydrocylone technology can only be feasible if using them in series. Since cut size is a direct function of hydrocyclone diameter, the best efficiency for a number of useful applications would be given by small units. On the other hand, it is known that capacity is a direct function of hydrocyclone diameter, and so larger units will deliver higher flow rates. Therefore, it would be necessary to reach a compromise between efficiency and capacity for industrial operation. In this sense, a basic small geometry may be kept to cut as fine particles as possible, while series arrangements may be required to render industrial capacities. A number of arrangements have been experienced, but probably the most efficient is the radial (spider) arrangement with the hydrocyclones axes horizontally place like spokes in a wheel lying down. As shown in Figure 10.33, one "wheel" or hydrocyclone deck accommodates a number of "spokes" or hydrocyclone bodies. For cyclone cavities with a cylindrical diameter of

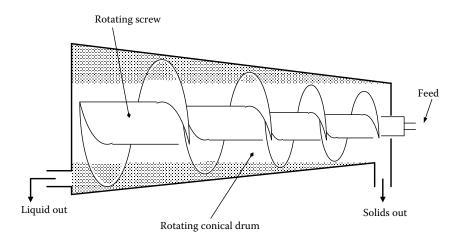


FIGURE 10.32 Principle of conveyor-bowl centrifuge.

10 mm the diameter of the deck is 300 mm. The whole cyclone deck can be molded by injection in one piece. The feed zone is the annular space closest to the periphery. The overflow is through peripheral screw-in-vortex finders. The underflow leaves via the central circular space where the small apex openings enter.

10.4.5.7 Laboratory Exercise: Efficiency of Hydrocyclone Separations

10.4.5.7.1 Introduction

Hydrocyclones can compete with centrifuges in clarification duties in diverse applications of the materials processing industries. Advantages of hydrocyclone technology are: low cost, small space requirement, no moving parts, easy cleaning, and no need to use any aid for separation. Hydrocyclones are easily manufactured and modified and have been well tested in thickening, clarification, classification, and other operations in many industries. The best way to define efficiency of hydrocyclones is by determining the cut size at given conditions. The aim of the exercise is to evaluate the cut size in separating a suspension using a hydrocyclone.

10.4.5.7.2 Equipment and Materials

A hydrocyclone test rig comprising a sump, a small hydrocyclone unit (10–20 mm in diameter), recirculating piping system, valves, pressure gauge, and a centrifugal pump, can be used for the experiment. Routine laboratory material and glassware to carry out gravimetric testing, such as analytical balance, oven, desiccator, and sample containers, will be

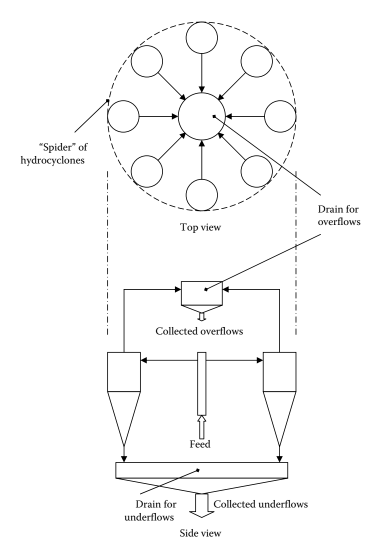


FIGURE 10.33
Diagram of radial (spider) arrangement of hydrocyclones for industrial scale.

sufficient for the test. The suggested material is fine limestone powder (median size 5–10 $\mu m)$ to prepare a dilute suspension in water.

10.4.5.7.3 Instruments and Apparatuses

An instrumental particle size analyzer, such as a laser diffraction or electro sensing zone counter, will be appropriate options for measuring particle size distributions. A rotational viscometer will be used to measure apparent viscosity if the suspension results are non-Newtonian.

10.4.5.7.4 Procedure

- Prepare a suspension with a concentration of about 5% by volume of fine calcium carbonate or similar powder in water.
- Run a test following the procedure described in Section 10.4.1.1. Determine whether the suspension is Newtonian or not. If the suspension is non-Newtonian, derive the rheograms to determine parameters of characterization of the fluid.
- Determine the particle size distribution of feed using an instrumental particle sizer.
- Operate the hydrocyclone rig by filling the tank with the suspension. Start the pump with the recirculating valve fully open. Allow the suspension to recirculate for a few minutes to stabilize.
- Determine the flow rate of feed.
- Open fully the valve for feed of the hydrocyclone. Slowly close the
 recirculating valve until all the feed stream goes to the cyclone. With
 the recirculating valve fully closed and the hydrocyclone feeding
 valve fully open, the maximum pressure gauge reading will be the
 maximum pressure drop attainable. Register such pressure when is
 as stable as possible.
- Determine the flow rate of leaving streams from the hydrocyclone.
- Carefully take a sample of the overflow and underflow simultaneously. Make two replicates of sampling.
- Determine gravimetrically the total efficiency of the separation process.
- Determine the particle size distribution of overflow and underflow streams with an instrumental particle sizer.

10.4.5.7.5 Calculations and Report

- Following a procedure like the one described in Example 9.2, derive the reduced grade efficiency curve for the separation process.
- From the curve, read off the cut size attained in the test.
- Present a complete written report, according to scientific report guidelines, either given by an instructor or consulted in the literature. Include graphs, charts, and calculations carried out. Include analysis from own data and, as much as possible, discussion based on literature survey from studies and investigations on the subject.

10.4.6 Filtration

Filtration may be defined as the unit operation in which the insoluble solid component of a solid-liquid suspension is separated from the liquid component by passing the suspension through a porous barrier which retains the solid particles on its upstream surface, or within its structure, or both. The solid–liquid suspension is known as the feed slurry or prefilt, the liquid component that passes through the membrane is called the filtrate and the barrier itself is referred to as the filter medium. The separated solids are known as the filter cake, once they form a detectable layer covering the upstream surface of the medium. The flow of filtrate may be caused by several means. Pressure and vacuum are two conventional ways of driving the suspension across the medium. Gravity and centrifugal forces may also be used for suspension medium crossing. In general terms, filtration theory applies to cases where cake buildup occurs.

10.4.6.1 Filtration Theory

In the initial stages of filtration the first particles of solid to encounter the filter medium become enmeshed in it, reducing its open surface area and increasing the resistance it offers to the flow of the filtrate. As filtration proceeds a layer of solids builds up on the upstream face of the medium and this layer, or cake, increase in thickness with time. Once formed, this cake in fact becomes the primary filtering medium. Filtrate passing through the filter encounters three types of resistance: a first resistance offered by channels of the filter itself, a second one because of the filter medium presence, and a third one due to the filter cake. The total pressure drop across the filter is equivalent to the sum of the pressure drops resulting from these three resistances. Usually, the pressure drop due to the channels of the filter is neglected in calculations. If $-\Delta P$ is the total pressure drop across the filter and $-\Delta P_{\rm c}$ and $-\Delta P_{\rm m}$ the pressure drops across the cake and medium respectively, then:

$$-\Delta P = -\Delta P_{\rm c} - \Delta P_{\rm m} \tag{10.88}$$

The pressure drop across the filter cake may be related to the filtrate flow by the expression (McCabe et al., 2005):

$$-\Delta P_{\rm c} = \frac{\alpha \mu w V}{A^2} \left[\frac{\mathrm{d}V}{\mathrm{d}t} \right] \tag{10.89}$$

where α is the specific resistance of the cake, μ is the viscosity of filtrate, w is the mass of solids deposited on the medium per unit volume of filtrate, V is the volume of filtrate, and A is the filter area normal to the direction of filtrate flow.

Alpha, α , physically represents the pressure drop necessary to give unit superficial velocity of filtrate of unit viscosity through a cake containing unit mass of solid per unit filter area. It is related to the properties of the cake by

$$\alpha = \frac{k(1-\varepsilon)S_0^2}{\varepsilon^3 \rho_o} \tag{10.90}$$

where k is a constant, ε is the porosity of the cake, S_0 is the specific surface area of the solid particles in the cake, and ρ_s is the solids density.

If a cake is composed of rigid nondeformable solid particles α is independent of $-\Delta P_c$ and does not vary throughout the depth of the cake, and is known as incompressible cake. However, if the cake contains nonrigid, deformable solid particles or agglomerates of particles the resistance to flow will depend on the pressure drop and will vary throughout the depth of the cake. In this case the cake is called compressible and an average value of the specific resistance for the entire cake must be used in Equation 10.89. This average specific resistance must be measured experimentally for any particular slurry.

By analogy with Equation 10.89 the filter medium resistance may be defined by the following relation:

$$-\Delta P_{\rm m} = \frac{R_{\rm m}\mu}{A} \left[\frac{\mathrm{d}V}{\mathrm{d}t} \right] \tag{10.91}$$

where $R_{\rm m}$ is a constant known as the filter medium resistance.

It is reasonable to assume that $R_{\rm m}$ is constant during any filtration cycle and that it includes the resistance to filtrate flow offered by the filter channels. This being the case, Equations 10.88, 10.89, and 10.91 can be combined to give

$$\frac{\mathrm{d}V}{\mathrm{d}t} = \frac{A(-\Delta P)}{\mu \left[\alpha w V/A + R_{\mathrm{m}}\right]} \tag{10.92}$$

Equation 10.92 is a general expression for the filtrate flow rate.

10.4.6.2 Constant Pressure Filtration

When the pressure drop is maintained constant, Equation 10.92 may be integrated thus

$$\int_{0}^{t} dt = \frac{\mu}{A(-\Delta P)} \left[\frac{\alpha w}{A} \int_{0}^{V} V \, dV + R_{\rm m} \int_{0}^{V} dV \right]$$
(10.93)

or, substituting limits and transposing for time *t*:

$$t = \frac{\mu}{(-\Delta P)} \left[\frac{\alpha w}{2} \left(\frac{V}{A} \right)^2 + R_{\rm m} \left(\frac{V}{A} \right) \right]$$
 (10.94)

Equation 10.94 is a general expression for the filtration time during constant pressure filtration. In order to use it, values of α and R_m must

be determined experimentally. This can be done by rewriting it in the following form:

$$\frac{\mathrm{d}t}{\mathrm{d}V} = KV + B \tag{10.95}$$

where

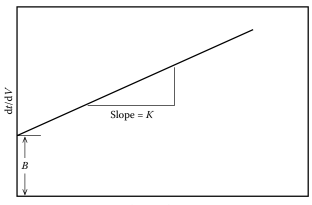
$$K = \left[\frac{\alpha w \mu}{A^2(-\Delta P)}\right] \tag{10.96}$$

and

$$B = \left\lceil \frac{R_{\rm m} \mu}{A(-\Delta P)} \right\rceil \tag{10.97}$$

As can be gathered, Equation 10.95 represents a straight line if dt/dV is plotted against V. Therefore, if a constant pressure filtration is carried out and values of V for different values of t are recorded, a graph of dt/dV versus V can be constructed as shown in Figure 10.34. The slope of this line is K and the intercept on the ordinate when V = 0 is B. Thus, by using a graph resulting of plotting dt/dV vs V, values of α and $R_{\rm m}$ can be directly determined from Equations 10.96 and 10.97, respectively.

For incompressible cakes Equation 10.94 can be used directly at different pressures. However, for compressible cakes, the relationship between α and $-\Delta P$ needs to be determined experimentally by performing filtration runs at different constant pressures. Empirical equations may be fitted to



Filtrate volume, V

FIGURE 10.34Graph of results from a constant pressure filtration test.

the results obtained. Two such equations have been suggested (McCabe et al., 2005):

$$\alpha = \alpha_0 (-\Delta P)^s \tag{10.98}$$

$$\alpha = \alpha_0' [1 - \beta(-\Delta P)^{s'}] \tag{10.99}$$

where α_0 , α'_0 , s, and β are empirical constants. Constant s is the compressibility coefficient of the cake taking a value of zero for incompressible sludges and positive for compressible ones. Having determined values for α_0 and s then Equations 10.98 and 10.99 can be used for constant pressure filtration calculations at different pressures when the cake is compressible.

Example 10.8

Table 10.9 presents data from a filtration run of slurry containing 23.47 kg/m³ of solids. The run was carried out at a constant pressure of 338 kPa in a plate-and-frame filter press with an effective filtration area of 0.0439 m². Calculate the constants α and $R_{\rm m}$.

SOLUTION

From data in Table 10.9 values of Δt and ΔV are worked out by progressive differentiation. Using those values a column of $\Delta t/\Delta V$ is derived and tabulated against a column of average volume $[(V_1 + V_2)/2]$. From these data a second table (Table 10.10) is obtained.

From Table 10.10 a plot of $\Delta t/\Delta V$ versus V will be constructed, as shown in Figure 10.35.

TABLE 10.9 Example 10.8

-			
Filtration Time (s)	Filtrate Volume (m³)		
9.4	0.000498		
9.5	0.001000		
16.3	0.001501		
24.6	0.002000		
34.7	0.002498		
46.1	0.003002		
59.0	0.003506		
73.6	0.004004		
89.4	0.004502		
107.3	0.005009		

TABLE	10.10
Examp	le 10.8

t (s)	V (m ³)	Δt	ΔV	$\Delta t/\Delta V$	$[(V_1 + V_2)/2]$
0	0				
9.4	0.000498	4.4	0.000498	8835	0.000249
9.5	0.001000	5.1	0.000502	10159	0.000749
16.3	0.001501	6.8	0.000501	13573	0.001250
24.6	0.002000	8.3	0.000499	16633	0.001750
34.7	0.002498	10.1	0.000498	20281	0.002249
46.1	0.003002	11.4	0.000504	22619	0.002750
59.0	0.003506	12.9	0.000504	25595	0.003254
73.6	0.004004	14.6	0.000498	29317	0.003755
89.4	0.004502	15.8	0.000498	31727	0.004253
107.3	0.005009	17.9	0.000507	35306	0.004755

The graph in Figure 10.35 is a straight line with a correlation coefficient 0.997. The slope of such line (K) is 6×10^6 s/m⁶, while the intercept with the y-axis (b) is 6352.4 s/m³. Transposing for α and R_m from Equations 10.96 and 10.97 respectively, the values of such constants are calculated as

$$\alpha = \frac{(6\times10^6)\text{s/m}^6(0.0439)^2\,\text{m}^4(338000)\text{kg/s}^2\,\text{m}}{(23.47)\text{kg/m}^3(0.001)\,\text{kg/ms}} = 1.665267\times10^{11}\text{m/kg}$$

$$R_m = \frac{(6352.4)\text{s/m}^3(0.0439)\text{m}^2(338000)\text{kg/s}^2\,\text{m}}{(0.001)\text{kg/ms}} = 9.4258\times10^{10}\,\text{L/m}$$

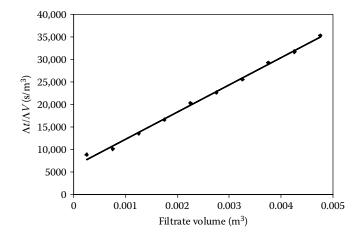


FIGURE 10.35 Example 10.8.

10.4.6.3 Constant Rate Filtration

If filtration is carried out at constant rate, then

$$\frac{\mathrm{d}V}{\mathrm{d}t} = k = \frac{V}{t} \tag{10.100}$$

Equation 10.100 may be rewritten as follows:

$$-\Delta P = \left[\frac{\mu \alpha w V}{A^2 t}\right] V + \left[\frac{\mu V R_{\rm m}}{A t}\right]$$
 (10.101)

or

$$-\Delta P = K'V + B' \tag{10.102}$$

where

$$K' = \left[\frac{\mu \alpha w V}{A^2 t}\right] \tag{10.103}$$

and

$$B' = \left\lceil \frac{\mu V R_{\rm m}}{At} \right\rceil \tag{10.104}$$

Once again it can be seen that Equation 10.102 represents a straight line if $-\Delta P$ is plotted against V. The slope of the line is K' and the intercept to the y-axis when V=0 is B'. Thus, for incompressible cakes α and $R_{\rm m}$ can again be determined by experimental means.

For compressible cakes, the relationship between α and $-\Delta P_{\rm c}$ must, again, be experimentally determined. If a relationship of the form shown in Equation 10.98 is assumed to apply, then Equation 10.89 may be modified to

$$(-\Delta P_{\rm c})^{1-s} = [-(\Delta P - \Delta P_{\rm m})]^{1-s} = \frac{\mu \alpha_0 w V}{A^2} \frac{V}{t}$$
 (10.105)

which, in turn may be written as

$$[-(\Delta P - \Delta P_{\rm m})]^{1-s} = K''t$$
 (10.106)

where

$$K'' = \frac{\mu \alpha_0 w}{A^2} \left\lceil \frac{V}{t} \right\rceil^2 \tag{10.107}$$

If it is assumed that $-\Delta P_{\rm m}$ is constant throughout a constant-rate filtration then by plotting t versus $-\Delta P$ and passing a smooth curve through the points and extrapolating the curve to the y-axis, an approximate value for $-\Delta P_{\rm m}$ can be obtained. If $-(\Delta P - \Delta P_{\rm m})$ is then plotted against t on log-log paper and a straight line is obtained, the slope of this line is (1-s). Thus, s can be calculated and K'' and α_0 can be derived from Equations 10.106 and 10.107. When the first log-log plot of $-(\Delta P - \Delta P_{\rm m})$ is not a straight line; further approximations for $-\Delta P_{\rm m}$ need to be made (McCabe et al., 2005).

Example 10.9

Calculate the volume obtained, and the time required in reaching such volume, in a constant-rate filtration process keeping a rate of 0.05 m³/min until the pressure drop reaches 250 kN/m², according to data in Table 10.11.

SOLUTION

For constant rate filtration, plotting directly $-\Delta P$ versus V, the graph shown in Figure 10.36 is obtained.

The correlation parameters of the slope line represented in Figure 10.36 are: K' = 490 and B' = 15.5, with a correlation coefficient of 0.9971. The equation representing the particular straight line in Figure 10.36 is, thus

$$-\Delta P = 490V + 15.5$$

Substituting the value of 250 kPa in the above equation and transposing for V, the volume at that maximum attained pressure is: V = 0.4878 m³.

Since time in Table 10.11 shows the same linear trend than pressure drop and volume, for the pressure drop of 250 kPa the approximate time is t = 8.92-9 min.

TABLE 10.11 Example 10.9

1			
t (min)	$-\Delta P (kN/m^2)$	V (m ³)	
0	0	0.00	
1	40	0.05	
2	65	0.10	
3	90	0.15	
4	110	0.20	
5	140	0.25	

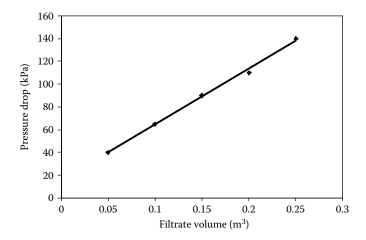


FIGURE 10.36 Example 10.9.

10.4.6.4 Filtration Media and Filter Aids

Filtration equipment is adapted with consumable materials for every particular duty. The main consumables in diverse filtration operations are the filtering medium and the filter aids. Filter media are selected, primarily, in function of the pore size that will be appropriate for retention of the solids in suspension. Other important functions of the filtering medium are the promotion of cake formation and the support of cake once formed. The medium should offer minimum resistance to flow accordingly to the need of a rapid formation of the filter cake. It must be strong enough to support the cake and retain its strength under the extreme conditions that may occur during operation. The surface characteristics of the filtering medium should facilitate cake removal. It must be nontoxic and chemically compatible with materials being filtered, and also it should be cost effective. Filter media may be rigid or flexible. Rigid media may be loose or packed. Examples of loose rigid media are sand, gravel, diatomaceous earths, and charcoal. Packed or fixed rigid media include porous carbon, porcelain, fused alumina, perforated metal plates, or rigid wire mesh. Flexible media may comprise woven fabrics of cotton, silk, wool, and jute. Flexible media made out of synthetics include nylon, polypropylene, polythene, polyvinylchloride, and copolymers. Glass fiber and flexible metal meshes are also used as filter media, as well as nonwoven materials, such as cotton fibers, wool fibers, and paper pulp. These materials may be available in preformed pads of various shapes. Table 10.12 summarizes different filtering materials along with their main features. Some other characteristics, such as the woven pattern and the geometric dimensions of the fibers, are considered selection criteria and can be presented in tabular form (Clark, 1990).

TABLE 10.12Physical Properties of Fibers Used in Materials for Filtering Media

Fiber	Maximum Temperature (°C)	Specific Gravity	Water Absorbance (%)	Elongation (% to Rupture)	Durability
Acetate	99	1.30	9–14	30–50	Poor
Acrylic	135-149	1.14-1.17	3–5	25-70	Good
Cotton	93	1.55	16–22	5–10	Regular
Spun glass	288-315	2.50-2.55	Up to 0.3	2–5	Poor
Nylon	107-121	1.14	6.5-8.3	30–70	Excellent
Polyester	149	1.38	0.04 - 0.08	10-50	Excellent
Polythene					
Low density	66–74	0.92	0.01	20-80	Good
High density	93-110	0.92	0.01	10-45	Good
Polypropylene	121	0.91	0.01-0.1	15-35	Good
PVC	66–71	1.38			Regular
Rayon	99	1.50-1.54	20-27	6-40	Poor
Saran	71-82	1.7	0.1-1.0	15-30	Regular
Wool	82–93	1.3	16–18	25–35	Regular

Another way of facilitating the filtration process, mainly when solids to be filtered are finely divided or are of a slimy highly compressible nature, is the use of filter aids. For this described type of solids the filtering medium tends to block quickly shortening, thus, filter cycles considerably. To overcome this difficulty and obtain reasonably filtering cycles, materials referred to as filter aids are often used. They consist of comparatively large, inert, noncompressible solid particles of different shapes. They may be applied by mixing with the feed slurry or they may be suspended in clear liquid, often some filtrate from a previous run, which is then passed through the filter in order to build a precoat of the filer aid on the medium. The filter aid forms then a rigid lattice structure on the medium and provides numerous channels for the filtrate to flow, slowing the plugging of the medium and prolonging filtration cycles. Examples of commonly used filter aids are kieselguhr and diatomaceous earths. Paper pulp, carbon, fuller's earth, and some other materials can be used as filter aids. As with filtering media, filter aids are available in a wide range of grades to suit particular applications.

10.4.6.5 Filtration Equipment

The most common types of filters used in industrial processing are pressure filters such filter presses, leaf filters, and cartridge filters, as well as vacuum filters like the rotary drum vacuum filter, and the rotary vacuum disc filter. Most filtration units produce a cake, which at long cycles may become reasonably dry at the expense of declining the flow rates. Keeping constant

rate filtration is costly as the pressure drop must be increased during the cycle.

A commonly used design of filter press is the plate-and-frame filter press. The components of a plate-and-frame filter press are the frames and the two types of grooved plates, filter plate and wash plate, shown in Figure 10.37a. The grooved plates covered on both sides with filter medium and alternatively assembled with frames, as illustrated in Figure 10.37b, are mounted in a rack. The whole assembly can be squeezed tightly together by a screw driven manually or by a hydraulic or pneumatic mechanism, to form a liquid tight unit (Figure 10.38). Both plates and frames are provided with openings

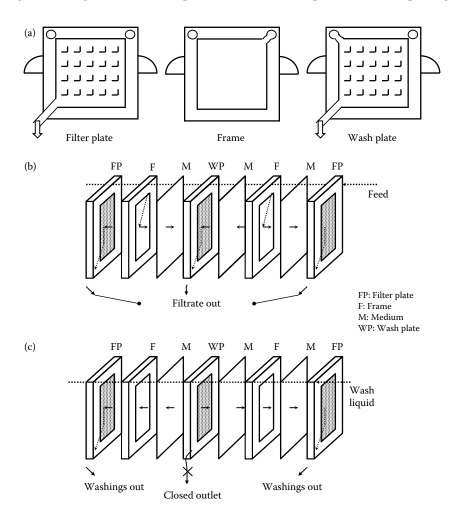


FIGURE 10.37

Description and arrangements of filter press components: (a) detail of plates and frames, (b) arrangement for the filtration cycle, and (c) arrangement for the washing cycle.

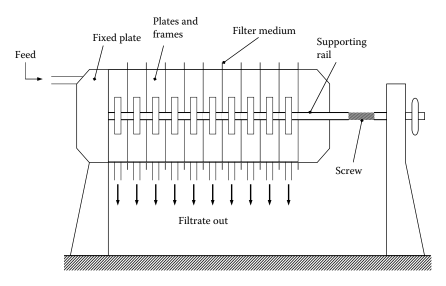


FIGURE 10.38Schematic diagram of a plate-and-frame filter press assembled and mounted.

at one corner, so that when the press is closed these openings form a channel through which the feed is introduced. Slurry is fed into the frames and the cake builds up in the hollow center of the frames. The filtrate passes through the medium and to the grooved surfaces of the filter plates to be removed via an outlet channel in each plate. Filtration continues until the flow of filtrate drops below a practical level, or when accumulation of cake in the frames elevates pressure to an extremely high level. Washing of the cake can be carried out by replacing the flow of feed slurry with wash liquid or by the use of special wash plates mounted in the arrangement so that every second plate is a wash plate. On running a filtration cycle these wash plates act as filter plates, while during washing their outlets are closed and the wash liquid is introduced to their surfaces through a special inlet channel. The flow path for both filtration and washing cycles using wash plates is illustrated in Figure 10.37b and c, respectively.

Another alternative of pressure filtration are pressure-vessel filters that make use of filter leaves as the basic filter element. A filter leaf consists on a wire mesh screen or grooved drainage plate over which the filter medium is stretched. The leaf may be suspended from the top or supported from the bottom or center. The supporting element is usually hollow and forms an outlet channel for the filtrate. Several designs of pressure-vessel filters with filter leaves elements are commercially available. Two of them are the horizontal-leaf filter and the vertical-tank vertical-leaf filter, both shown in Figure 10.39. Vertical-tank vertical-leaf filters (Figure 10.39a) are the cheapest of the pressure leaf-filters and have the lowest volume-to-area ratio. Cake discharge is accomplished by sluicing, either manually or by vibration. Horizontal-leaf

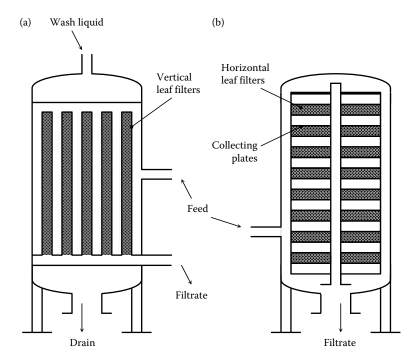


FIGURE 10.39 Pressure-vessel leaf filters: (a) vertical-tank, vertical-leaf filter, and (b) horizontal-leaf filter.

filters (Figure 10.39b), in either a horizontal or vertical tank, are advantageous when the flow is intermittent. Filtration takes place only on the upper size of the horizontal leaves, and so filtration area is limited compared with other pressure-vessel designs. Horizontal-leaf filters are most suitable when thorough cake washing is necessary. Cake discharge may be accomplished by rotation of the leaves.

In vacuum filters a subatmospheric pressure is maintained downstream of the medium while an atmospheric one is kept upstream. Because the pressure drop across the filter is limited to one atmosphere, they are not suited to batch operations. Some types of leaf filters, tube filters, or edge filters can be operated under vacuum, but continuous vacuum filters are far most common. Since the upstream pressure is atmospheric, cake discharge is facilitated thus favoring continuous operation. The rotary drum vacuum filter is, perhaps, the most common type of vacuum equipment used in an important number of applications. As can be seen in Figure 10.40a, vacuum is applied to the inside of a revolving drum covered with reinforced wire mesh and filtering cloth or medium. The surface of the drum consists of a number of shallow compartments formed between dividing strips running the length of the drum. Each compartment is connected by one or more pipelines to an

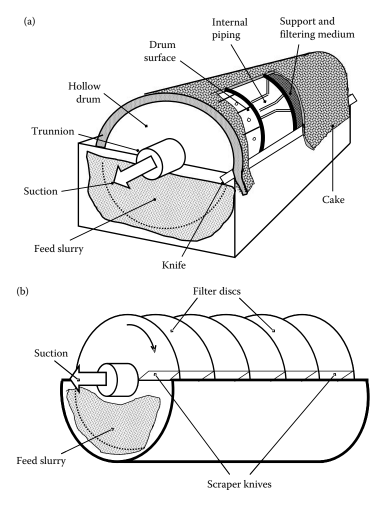


FIGURE 10.40 Equipment for vacuum filtration: (a) continuous rotary drum, and (b) rotary disc.

automatic rotary valve situated centrally at one end of the drum. Filtering medium covers the entire drum surface and is supported by perforated plates, grids, or wire mesh to provide drainage space between the medium and the floor of each shallow compartment. The drum dips into slurry and rotates, a vacuum is drawn on its surface and filtrate flows through the medium and out through the drain pipe from the compartment and is directed to the filtrate receiver by means of the valve. A layer of cake builds up on the outer surface of the medium and the cake is sucked free from the filtrate as the drum turns becoming dryer near the end of the turn. Prior to be submerged again in the slurry tank in completing a drum revolution, a knife removes the cake in the form of a continuous layer. Rotary vacuum

precoat filters may produce filtrates which would contain 0.1% or less suspended solids. Rotary vacuum disc filters (Figure 10.40b) consist of a number of circular filter leaves mounted on a horizontal axis about which they rotate. Each disc is fitted with a cake removal device and is divided into sectors, each of which has an individual outlet to the central shaft. These outlets form a continuous channel through which the filtrate flows from all sectors at the same angle. All the channels terminate in a rotary valve like the one used in drum filters. In operation each disc operates as a drum filter, with the cycle being controlled by the rotary valve.

10.4.6.6 Laboratory Exercise: Filtration Curves Using a Filter Press

10.4.6.6.1 Introduction

Filtration is a unit operation with diverse applications in the materials processing industries. The mode normally used in industrial filtration is cake filtration in which solids build up on the surface of a filtering medium, its accumulation known as cake. The factor used to make pass the filtrate through the cake and the medium can be pressure force or vacuum force. Pressure is used in an important number of equipment. The pressure drop is maintained constant in the filtration mode referred to as constant pressure filtration. In this mode of filtration the flow rates decline as the filtration cycle progresses. Another mode of filtration is constant rate filtration, in which the flow rates do not decline for a given cycle. Constant pressure filtration is, normally, more convenient because constant rate filtration is costly as the pressure drop must be increased during the cycle. A common piece of filtration equipment in numerous processing applications is the plate-andframe filter press. Any type of pressure-vessel filter normally requires filter aids to facilitate cake formation and a trouble-free operation. Some types of biological slurries, such as nonclarified fruit juices, tend to form a cake relatively easy without need of filter aids due to the pectinaceous nature of the suspended solids.

The objective of the experiment is to run a plate-and-frame filter press in constant pressure mode in order to determine the filtration plot describing a filtration cycle, and calculate the empirical constants of cake resistance and filter medium resistance.

10.4.6.6.2 Equipment and Materials

A laboratory size filter press rig comprising tank, pipeline, centrifugal pump, filter plates, valves, and pressure gauges. Domestic juice extractor and utensils of a food processing plant, such as knives, apple corer, containers, and so on. Routine laboratory material and glassware to carry out gravimetric testing, such as analytical balance, oven, desiccator, and sample containers, will be sufficient for the test. A batch of second-grade apples suitable for juice extraction is the suggested raw material.

10.4.6.6.3 Procedure

- Use a batch of apple sufficient to prepare about 10 L of nonclarified juice.
- Weigh the apples, wash them, core them, and peel them.
- Extract the juice until obtaining the approximate 10-L sample.
- Take an aliquot of the juice sample and determine, gravimetrically, its total solids content.
- Mount the filter press arrangement checking for leaks.
- Fill up the filter tank, record the volume.
- Open the filter feeding and the recirculating valves.
- Recirculate the suspension for 10 min.
- Stabilize the pressure gauge reading at 150 kPa, approximately.
- Open the valve of filtrate and close the valve of recirculation simultaneously.
- Take filtrate samples for 10 min, until the volume in the tank comes to a minimum.
- Verify the total volume of filtrate obtained.
- Drain the remaining liquid and clean the rig rigorously.

10.4.6.6.4 Calculations and Report

- Analyze the data collected following the procedure of Example 10.8, that is, derive a table like Table 10.10.
- Plot the data from the table obtained, and verify if they fit a straight line. If the linearity is confirmed, obtain the equation fitting the data from the regression model and determine the parameters.
- Calculate α and R_m from the obtained values of the slope K and the intercept B.
- Present a complete written report, according to scientific report guidelines, either given by instructor or consulted in the literature.
 Include graphs, charts, and calculations carried out. Include analysis from own data and, as much as possible, discussion based on literature survey from studies and investigations on the subject.

10.4.7 Other Techniques: Membrane Separations

Membrane separations are techniques used industrially for removal of solutes and emulsified substances from solutions by application of pressure onto a very thin layer of a substance with microscopic pores, known as a membrane. Membrane separation processes include reverse osmosis (RO), ultrafiltration (UF), microfiltration (MF), dialysis, electrodialaysis, gas

separation, and pervaporation. RO, UF, MF, and electrodialysis have been widely used commercially (Girard and Fukumoto, 2000). A suitable manner of classifying membrane separation techniques is by referring to their limiting size of retention. RO separates sizes from 0.0001 to 0.001 μm , UF range of separation is from 0.001 to 0.01 μm , and MF membranes have pore sizes 0.01–10 μm .

Due to the small size range of pores involved in membranes, an alternative convenient way of referring to separating capability is by means of a molecular weight rating, expressed in terms of a rejection coefficient against a species of specific molecular weight (Chen et al., 2004). Ideally, the membranes will have a sharply defined molecular weight cutoff (MWCO). Such an ideal membrane will retain all species greater than the MWCO but will allow all smaller ones to pass. Membranes are available in a number of increments in molecular weight cutoff ranges from 1000 up to 100,000 Daltons (Da). For example, MWCO range of RO is below 300 Da while that for UF membranes is between 300 and 300,000 Da.

Membrane separations were originated as a consequence of the need of desalting sea water using alternatives to the thermal processes based on evaporation. Sea water was successfully desalinated by a technique known as reverse osmosis in the 1958 (Sourirajan, 1970). The principle of solutes separation by reverse osmosis is illustrated in Figure 10.41. The natural solvent transfer through a semipermeable membrane originates a pressure known as osmotic pressure. By inverting such natural process, the solutes concentrate even more in the solution containing them in higher proportion, while they dilute to a minimum in the solution containing them in the lower proportion. Using membrane separations, phases can be separated from

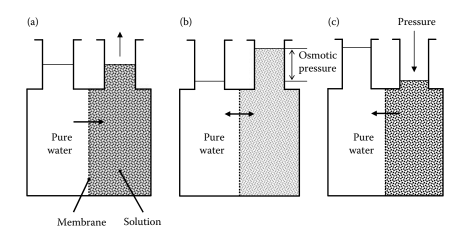


FIGURE 10.41
Principle of reverse osmosis. (a) Osmotic flow, (b) osmotic equilibrium, and (c) reverse osmosis.

two-phase and multiphase systems without the need of any phase state change, that is, without requiring thermal energy. The osmotic pressure π for dilute solutions can be established as a function of pressure and temperature, using fundamental thermodynamics, as follows:

$$\pi = MRT \tag{10.108}$$

where M is the molar concentration of the solution, R the universal gas constant, and T the absolute temperature.

UF is often compared with RO although the mechanism of separation is quite different. This difference is illustrated in Figure 10.42. As can be seen, in RO a rejection is based on electrostatic repulsion due to formation of a pure layer of water over the membrane, and the virtual charges on this layer reject charges of ionic free species of salt solutions (Figure 10.42a). Simultaneously, by a complex mechanism of sorption, diffusion, and desorption, pure water passes through the membrane performing the separation

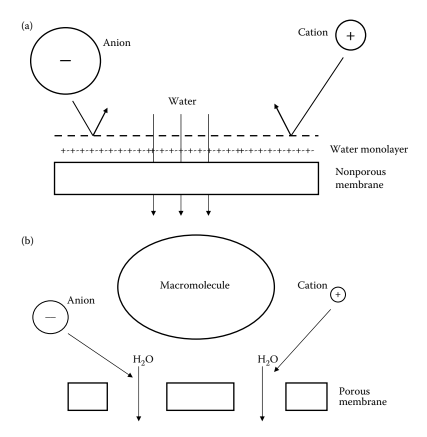


FIGURE 10.42
Rejection mechanisms for (a) reverse osmosis and (b) ultrafiltration.

process. UF membranes, on the other hand, are porous in nature, with a rigid and highly voided structure, and function in a manner analogous to a screen or sieve (Figure 10.42b). The pore network is randomly distributed, with pores passing directly through the membrane. The separating ability is based primarily on particle size, wherein particles and molecules larger than the largest pore are completely retained, whereas species smaller than the smallest pore are totally permeated. In general terms, the mechanism of separation in RO is known as salt rejection, whereas that in UF is called organic rejection. As described, UF can be considered as an extension of conventional filtration, with its separating ability ranging to the molecular level. The operating mode of the equipment is, however, different. As shown in Figure 10.43, in UF the fluid moves continuously across the membrane surface. The arrangement is known as cross-flow filtration and is used as means of sweeping the membrane surface to control the buildup of foulants and particulate matter.

Membrane separation processes operate in a continuous manner and so, unlike conventional filtration, the accumulated solids do not form a moist cake but, rather, dense slurry that is discharged like in centrifugation or hydrocyclone separation. This stream of concentrated solids is simply

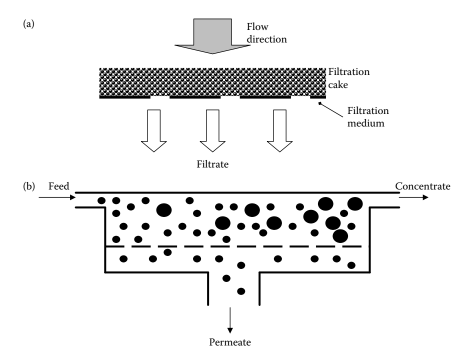


FIGURE 10.43

Modes of flow in filtration: (a) dead-end flow as in conventional cake filtration, and (b) cross-flow as in membrane separations.

known as concentrated while the clarified stream is called permeate. A schematic flow diagram of UF and some other membrane separations is shown in Figure 10.44. The flow rate *Q* through a membrane separation process may be represented by

$$Q = kA(\Delta P - \Delta \pi) \tag{10.109}$$

where k is a membrane permeability coefficient, A is the membrane superficial area, ΔP is the pressure drop, and $\Delta \pi$ the difference in osmotic pressure between the feed and the permeate. Equation 10.109 is more suitable for RO applications, due to the osmotic pressure exerted by solutes in such applications. For UF applications $\Delta \pi$ is negligible in relation to ΔP and the membrane thickness causes an effect in the process, and so it should be included as a variable. Under these considerations, Equation 10.109 can be reexpressed as

$$Q = \left[\frac{kA}{\delta}\right] \Delta P \tag{10.110}$$

where δ is the membrane thickness.

The pressure drop ΔP in Equation 10.110 is, really, an average pressure or pressure gradient due to the cross-flow arrangement of membrane separation processes. Referring to Figure 10.40, a feed stream containing differently size solids is pumped across the membrane surface at a velocity determined by the feed side pressure gradient ($P_1 - P_2$). This gradient, known as hydrodynamic pressure gradient, causes the continuous movement of fluid across the membrane that is referred to as cross-flow. As the feed stream flows across the membrane surface, smaller particles may be able to pass through the membrane and exit in the permeate stream (Figure 10.43) at a pressure P_3 , which is usually atmospheric. The rate of permeate flow is generally reported as flux, that is, flow rate per unit area of membrane. The driving force for

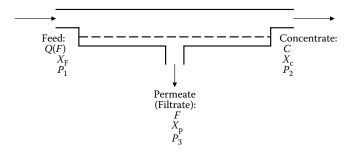


FIGURE 10.44 Mass balance in a membrane separation process.

permeate flow is also a pressure gradient, but is not the hydrodynamic pressure gradient defined above. Instead, is the pressure gradient that exits through the membrane, from feed size to permeate size, at each point along the membrane surface. This pressure gradient is known as the transmembrane pressure gradient or simply transmembrane pressure (TMP). Clearly, TMP varies along the membrane surface being a maximum at the inlet and a minimum at the outlet. An average TMP, according to the diagram in Figure 10.40, can be defined as

$$TMP = \frac{P_1 + P_2}{2} - P_3 \tag{10.111}$$

The permeate pressure P_3 is negligible as compared with the pressure gradient between feed and concentrate and, thus

$$TMP = \frac{P_1 + P_2}{2} \tag{10.112}$$

A way of expressing efficiency of a membrane separation process is by determining a recovery percentage as the relation of the permeate flow rate to the feed flow rate (Figure 10.44), that is,

$$%$$
Recovery = $\frac{F}{Q}(100)$ (10.113)

Recovery percentages depend on the feed concentration, being higher at lower concentrations of feed solids. A fraction of liquid, approximately equal to that of solids, should remain in the concentrate to make it flow. Therefore, as illustrated in Figure 10.45, the most diluted the feed, the higher the percentage of recovery.

Membranes for UF and similar processes are made out of a very thin film with microperforations of 0.1–1 μm size. Such tiny perforation on quite a thin film would be easily damaged by the applied pressures needed to perform separation in actual operation. The ultrathin skin or film is, therefore, supported on a relatively thick (100–200 μm thick) substructure. The retention is performed over the thin film, on which the pore sizes may vary due to the manufacturing technique. Two common membrane structures are the plane membrane and the hollow fiber membrane, illustrated in Figure 10.46.

Over the years, four generations of membrane materials have been appeared in the market. The first generation comprised cellulosic materials, the second included polymeric materials, the third was mainly from inorganic materials, and the fourth is based on carbon fibers. Conventional nomenclature of UF membranes includes two prefix letters (e.g., UM, PM, and XM) that refer to different polymers, and two last digits which indicate

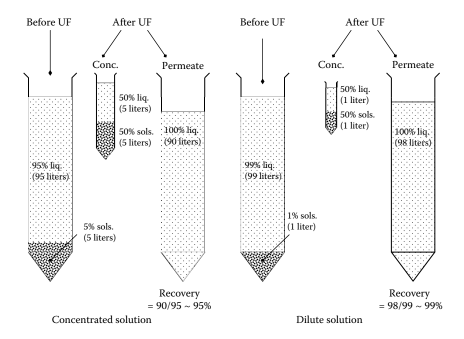


FIGURE 10.45Recovery percentages in membrane separation processes.

the nominal molecular weight cutoff. For instance, a commercial range of UF membranes would cover from UM05 to XM50, which would mean retention of macromolecules from about 5000 Da (UM05) to, approximately, 50,000 Da (XM50). Since pore sizes may vary due to material used and the manufacturing procedure, for the same nominal size, the actual pore size may be different. For example, Table 10.13 lists some types of membranes, classified according to the described nomenclature. As can be seen, for the same MWCO of 10,000 Da, two different membranes have different pore sizes.

TABLE 10.13Membranes Nomenclature

Membrane	MWCO (Da)	Approximate Pore Size (Å)
UM05	5000	21
UM10	10,000	30
PM10	10,000	38
PM30	30,000	47
XM50	50,000	66
XM100A	100,000	110
XM300	300,000	480

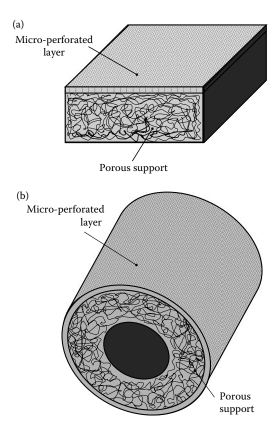


FIGURE 10.46
Common structures of membranes: (a) plane membrane and (b) hollow fiber membrane.

To be incorporated into an industrial installation, membranes are configured into modules and such modules mounted in structures comprising pipelines, valves, pressure gauges, and all the gadgetry that will constitute membrane separation equipment. Several arrangements or geometries of membranes are in use, including flat sheet membranes assembled similar in construction to a plate-and-frame filter press (Figure 10.47a), and the spiral-wound flat sheet membranes (Figure 10.47b) consisting of two flat sheets of membrane sandwiching a porous support medium, and wrapped with a plastic spacer around a central tube.

UF may be integrated in processing lines in more than one way. The simplest technique is the batch configuration illustrated in Figure 10.48a. In this mode, an initial volume of liquid is circulated through the UF system, and permeate is continuously removed until a final volume is achieved. On the other hand, recirculating topped-off batch is a variation shown in Figure 10.48b. This arrangement is more practical because it does not require the use

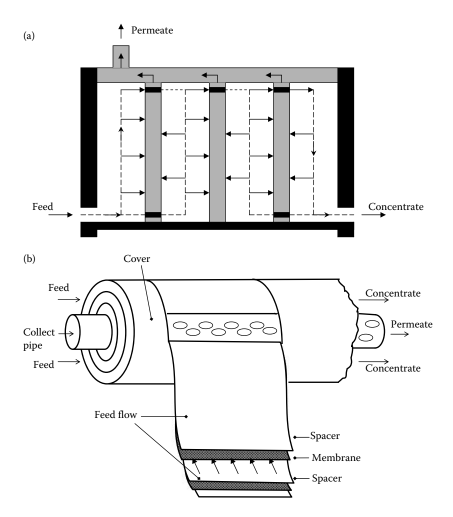


FIGURE 10.47 Membrane modules for assembly into equipment: (a) plate-and-frame arrangement and (b) spiral-wound arrangement.

of a large feed tank and allows the specification of a smaller prefilter prior to the system.

Example 10.10

Calculate the pressure drop needed to maintain a water flow rate of 15 mL/cm² min in a process of desalting sea water by RO at room temperature. The water contains 35 g/L of solids and the coefficient of permeability of the membrane can be taken as 1.125×10^{-4} cm/mm Hg min.

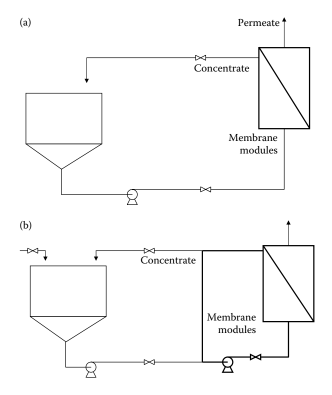


FIGURE 10.48

Diagram of membrane separation operation modes: (a) standard batch configuration and (b) topped-off batch system with recirculation loop.

SOLUTION

The osmotic pressure of the solution can be calculated using Equation 10.108, where molarity of salt is approximately 0.6 and at room temperature (298 K) the value of the universal gas constant is R = 0.08205 l atm/K mol. Substituting, thus, into Equation 10.108:

$$\pi = (0.6)(0.08205)(298) = 14.67 \text{ atm} \approx 11172 \text{ mm Hg}$$

The water flow rate is, indeed, a flux of water since it includes the area. Flux can be defined as: J = Q/A. On the other hand, since the osmotic pressure of the permeate approaches zero, $\Delta \pi$ is, simply, the osmotic pressure π , calculated above. Taking into account these arguments, transposing for ΔP in Equation 10.109, and substituting values:

$$\Delta P = \frac{J}{k} + \pi = \frac{15 \text{ cm}^3/\text{cm}^2 \text{ min}}{0.0001125 \text{ cm/mmHg min}} + 11172 \text{ mmHg} = 1.4451 \times 105 \text{ mmHg}$$

The pressure drop needed to maintain the required capacity is, thus: $\Delta P \sim 19,400 \text{ kPa}.$

10.4.8 Dimensionless Scale-up of Equipment

Optimization, adaptation, and innovation of industrial operations involving solid—liquid separations are based, like many other industrial processes, in laboratory experimentation followed by pilot plant testing and finishing with scale-up of equipment to establish a production line. Selection and sizing of equipment for solid—liquid separations is still a complex process requiring expert knowledge and expertise, and supported on testing on the actual system to be processed. Fundamentals play, also, an important role in the overall exercise of scaling-up procedures and, along with the expertise and testing, should render satisfactory results as to adjust theoretical calculations with real processing. This section reviews some basic theoretical principles useful for preliminary selection and sizing of solid—liquid separation equipment, in the most frequently employed categories.

Sizing of dynamics separators whose separation principle is governed by gravity or centrifugal forces may be easily scaled using dimensionless groups, providing the feed suspension is nonflocculating or the flocculation is completed prior to feeding of the suspension to the separator. The most relevant dimensionless groups for separators using gravitational or centrifugal forces that have been already described, discussed, and utilized in this book are the Reynolds number Re relating the inertial and viscous forces in a pocket of fluid, the Euler number Eu describing the resistance of a separator to flow, the Stokes number Stk relating the inertial and viscous forces on a particle, and the Froude number Fr relating the inertial and gravitational forces in a particle. In a form, more generic than the one used in this book, the Stokes number is represented by

$$Stk = \frac{\tau v}{D}$$
 (10.114)

An expression for the Froude number is

$$Fr = \frac{gD}{v^2} \tag{10.115}$$

In the Stokes number relationship τ is the particle relaxation time defined as

$$\tau = \frac{x^2(\rho_s - \rho)}{18\mu}$$
 (10.116)

In Equations 10.114 through 10.116 the components have already been defined, that is, v is the characteristic velocity of flow in the separator, D is

the characteristic linear dimension of the separator, g is the acceleration due to gravity, x is the equivalent Stokes diameter, ρ_s is the solids density, ρ is the liquid density, and μ is the liquid viscosity.

In gravity settling tanks, assuming low feed solids concentration and plug flow through a vessel of plan area *A*, the following relationship for its grade efficiency can be derived:

$$G(x) = Stk \cdot Fr \tag{10.117}$$

where the characteristic velocity is: v = Q/A, while Q is the volumetric flow rate and D is the height of the settling tank.

If *x* is the cut size x_{50} , then G(x) = 0.5, and the above equation becomes

$$Q = 2u_{\rm g}A \tag{10.118}$$

where $u_{\rm g}$ is the terminal settling velocity under gravity, of the cut size x_{50} from Stokes law:

$$u_{\rm g} = \frac{x_{50}^2(\rho_{\rm s} - \rho)g}{18\mu} \tag{10.119}$$

Equation 10.119 represents the very best performance of a settling tank. Such performance should be considered theoretical or ideal, and so it may not be totally applicable due to nonideal distribution of the flow, end effects, turbulence, and some others effects. However, it can be used as a preliminary approach to select the minimum area of the tank required to obtain a given cut size.

The sedimentation process in centrifugal devices is somewhat similar to sedimentation under gravity, except that the settling is radial and not parallel. A plug-model for the flow through the wall of the centrifuge leads to an exponential equation for the grade efficiency. For the cut size x_{50} , this mentioned equation would reduce to Equation 10.118 but with the area A replaced by a factor Σ . Such an equation is, in fact, Equation 10.74, the socalled Sigma theory equation. As mentioned when presenting such equation, factor Σ has a dimension of area and is calculated for the geometry of a particular centrifuge, so that each type of centrifuge has its own equation. Extensive work on the Sigma theory has shown that, while experimental results differ from calculated values when different centrifuges are considered, the scale-up between centrifuges of the same type is fairly reliable. This is based on a simple application of Equation 10.74 when a laboratory experiment with a small centrifuge 1 is compared with a larger scale operation on a larger centrifuge 2, of the same type and with the same slurry. For a case such as this, the following relation may apply:

$$\frac{Q_1}{\Sigma_1} = \frac{Q_2}{\Sigma_2} \tag{10.120}$$

When Equation 10.120 holds, the cut size and the overall separation efficiency, in both cases, should be the same. Equation 10.120 can be used for sizing centrifuges from laboratory tests, or for predicting capacities of existing machines.

Example 10.11

A centrifugal clarification system is used to eliminate settling solids from liquor resulting after a fermentation process. The solids have a density of 2650 kg/m³ and are suspended in water at a very low concentration. In laboratory tests it has been found that a satisfactory clarity with a sigma value of 220 m² and a capacity of 8×10^{-6} m³/s. If a larger centrifuge is used, with a sigma value of 2500 m² for large-scale operation: Which capacity will the new operation have? Which will be the effective cut size for that new capacity?

SOLUTION

Taking the values from laboratory testing as Σ_1 and Q_1 , substituting directly into Equation 10.120, and transposing for Q_2 :

$$Q_2 = \left(\frac{\Sigma_2}{\Sigma_1}\right) Q_1 = \frac{2500 \text{ m}^2}{220 \text{ m}^2} (8 \times 10^{-6}) \text{ m}^3/\text{s} = 9.09 \times 10^{-5} \text{ m}^3/\text{s}$$

The cut point can be obtained by substituting either set of values, that is, Q_1 and Σ_1 or Q_2 and Σ_2 , into Equation 10.74 and transposing for x_c . Using, thus, Q_1 and Σ_1 values and substituting in Equation 10.74, transposed for x_c :

$$x_{c} = \left[\frac{18(0.001) \text{ kg/ms } (8 \times 10^{-6}) \text{ m}^{3}/\text{s}}{2(1650) \text{ kg/m}^{3}(9.81) \text{ m/s}^{2}(220) \text{ m}^{2}} \right]^{1/2} = 1.42 \times 10^{-7} \text{m}$$

Therefore, the industrial capacity is 9×10^{-5} m³/s, approximately and the cut size is about 0.14 μ m.

For the case of centrifugal separation using hydrocyclones, at low solids concentrations (<1 or 2% by volume) the flow pattern in them is practically unaffected by the presence of particles, and so particle–particle interaction is negligible. The volume of the particles being separated into the underflow is small and the underflow-to-throughput ratio $R_{\rm f}$ is usually assumed to have no effect on the cut size x_{50} . The minimum effect of flow splitting can be easily compensated by using the reduced efficiency concept, as previously discussed. Dimensional analysis, coupled with fundamental theories of separation in hydrocyclones, gives two relationships between the three dimensionless groups Re, Eu, and Stk:

$$Stk'_{50}$$
 Eu = constant (10.121)

$$Eu = k(Re)^n \tag{10.122}$$

where k and n are empirical constants for a family of geometrically similar hydrocyclones, while Stk'_{50} is the Stokes number for the reduced cut size x'_{50} determined from the reduced grade efficiency G'(x) in Equation 10.29. The characteristic linear dimension is the diameter of the cylindrical part of the hydrocyclone and the characteristic velocity is the fluid linear velocity inside the cyclone defined by Equation 10.81. Constants k and k are determined for specific geometries of hydrocyclones, taking values as those given for Equations 10.78 through 10.80 and Equations 10.82 through 10.84. Scale-up calculations are particularly valuable for hydrocyclones, because efficiency and capacity are inversely proportional. Industrial operations normally require series of hydrocyclones, so that each unit will give the desired cut size while the complete arrangement will produce the needed capacity.

Example 10.12

It is required to clarify 8 L/s of the suspension in Example 10.4 using hydrocyclones of Rietma's optimum geometry. The concentration of the suspension is 15% by volume and its density has been measured as 1250 kg/m³. The maximum pumping capacity available for performing the separation is 315 kPa. Find out the optimum cyclone diameter and the number of units, if necessary, in order to obtain a cut size of 8 μ m. Small-scale experimentation has determined values of some dimensionless relationships as follows: $Stk_{50}Eu = 0.0625$, and Eu = 720. The density of the suspended solids is 2800 kg/m^3 .

SOLUTION

The suspension is highly concentrated, and resulted to be non-Newtonian (shear thickening type) as determined in solution of Example 10.4. The use of Equations 10.82 through 10.84 would be, therefore, most appropriate. The parameters of characterization n and K, from the rheogram of the suspension are: 1.39 and 2×10^{-5} kg/m s²⁻ⁿ, respectively.

Combining the definitions of Re* and Stk'_{50} (r) from Equations 10.55 and 10.56, with the definition of superficial velocity in the cyclone body (Equation 10.81), and considering the density of the suspension due to its high concentration, the following relations for the hydrocyclone diameter and the cut size are obtained:

$$\begin{split} D_c &= \sqrt[4]{\frac{(Q^2)(\text{Eu})(\rho_m)}{(1.23)(\Delta P)}} \\ \chi_{50}^{n'+1}(r) &= \frac{(\text{Stk}_{50}^*)(18)(3)^{n'-1}(K')\big[(2n+1)/(3n)\big]^{n'}(D_c)^{5-2n'}}{(1.27Q)^{2-n'}(\rho_s-\rho)} \end{split}$$

As previously stated, the cut size is directly proportional to the hydrocyclone diameter, while this (hydrocyclone diameter) is inversely proportional to the flow

rate. It would be, therefore, necessary to verify whether the required cut size could be obtained in a single hydrocyclone whose diameter would match the capacity needed. If both variables do not coincide (cut size and flow rate with a single unit) iterations would be needed until a number of units are found, each of them separating the required size, while processing the appropriate fraction of the total capacity. A first approximation would consist in calculating the hydrocyclone diameter, using the first equation derived above, which will handle the 8 L/s of slurry to be treated. Once having that diameter, with the second equation, the cut size that the single unit separates, will be known. If such diameter is coarser than 8 µm, the volumetric flow rate will be fractioned in equal parts to recalculate hydrocyclone diameter until a match, of number of units and cut size, is found. The first approximation leads to the following values:

$$D_{c} = \sqrt[4]{\frac{(0.008)^{2} \,\mathrm{m}^{6}/\mathrm{s}^{2}(720)(1250) \,\mathrm{kg/m}^{3}}{(1.23)(315000) \,\mathrm{kg/s}^{2} \,\mathrm{m}}} = 0.11 \,\mathrm{m}$$

and

$$x_{50}^* = 2.39 \sqrt[68.68 \times 10^{-5})(18)(1.53)(2 \times 10^{-5}) \frac{\text{kg/ms}^{2-n}(0.872)(0.007445)\text{m}^{5-2n}}{(0.0608)(\text{m}^3/\text{s})^{2-n}(1800)\text{kg/m}^3}}$$

$$x_{50}^* = 15 \ \mu m$$

So it is found that a 0.11 m diameter cyclone will separate particles much coarser than the required cut size. By performing iterations, if fractioning the rate 4 times, that is, when Q = 0.002 m³/s, a diameter of hydrocyclone $D_c = 0.055$ m and a cut size of 7.8×10^{-6} m are found. Therefore, the election would be the use of 4 units, 5.5 cm in diameter, to handle the capacity required and get the cut size needed.

Finally, for the case of filtration, conventional theory relates the pressure drop $-\Delta P$ to the flow rate Q through a porous bed by

$$\frac{Q}{A} = \frac{(-\Delta P)}{\mu R} \tag{10.123}$$

where R is known as the bed resistance. In cake filtration the bed resistance consists of the medium resistance in series with the resistance of the deposited cake so that the general filtration equation is written as

$$\frac{Q}{A} = \frac{(-\Delta P)}{\left\lceil \alpha \mu w V / A + (\mu R_{\rm m}) \right\rceil}$$
(10.124)

where $R_{\rm m}$ is the medium resistance.

The feed solids concentration has a profound effect on the performance of any cake filtration equipment. It affects the capacity and the cake resistance, as well

as the penetration of solids into the cloth. Thicker feed leads to improved performance of most filters, through higher capacity and lower cake resistance. The effects on solids yield can be demonstrated (Svarovsky, 1985) using the following equation:

$$Y = \left[\frac{2(-\Delta P)fw}{\rho \mu t_{\rm c}} \right]^{1/2} \tag{10.125}$$

where Y is the solids yield, that is, dry cake production in kg/m² s, f is the ratio of filtration and cycle times (submergence in continuous filters) and t_c is the cycle time. For the same cycle time (the same speed if the filter is continuous), if the concentration is increased by a factor of 4, production capacity is doubled. In other words, filtration area can be halved for the same capacity. For given operation conditions and submergence the dry cake production rate increases with the speed of rotation, accordingly to Equation 10.125, so the limiting factor is normally the cake thickness that can be successfully discharged by the method used in the filter.

PROBLEMS

- 1. Calculate the diameter of a cyclone, and the number of units if necessary, to recover particles of 6 μ m from dust-laden air. The density of the particles is 2600 kg/m³. The flow rate to be treated is 2.5 m³/s, and the maximum pressure drop attainable by the pumping capacity is 1200 N/m². Determined values of some dimensionless relationships by small-scale experimentation are the following: Stk₅₀ = 0.006, and Eu = 46.
- 2. Determine the free-settling time through a depth of 2 m of water at 20°C, for the particles listed in Table 10.14. All the particles may be considered spherical.
- 3. Calculate the rising velocity of an oil droplet $4\,\mu m$ in diameter, within an emulsion comprising an oily phase of $0.930\,g/cm^3$ density and an aqueous phase $1.036\,g/cm^3$ density. The viscosity of the aqueous phase is $1.6\times10^{-3}\,N\,s/m^2$.
- 4. A suspension of calcium carbonate in water is to be settled in a continuous sedimentation unit. The size of the particles is reasonably uniform of 50 μ m and the shape can be considered fairly spherical. The suspension has a temperature of 25°C and the specific gravity of the particles is 2.93. Calculate the concentration of the suspension if the settling velocity, determined experimentally, is 1.5×10^{-3} m/s.

TABLE 10.14
Problem 2

1 10010111 =		
Substance	Specific Gravity	Diameter (mm)
Galena	7.5	0.250
Carbon	1.3	6.000

TABLE 10.15 Problem 7

Produced Filtrate (kg)	Time (s)
0	0
0.25	7.0
0.50	20.0
0.75	35.0
1.00	53.0
1.25	76.0
1.50	102.0

- 5. Calculate the capacity of a centrifugal clarifier separating particles $2 \mu m$ in size and higher. The clarifier is 2 in diameter and 9 in height. The particles are suspended in water, have a density of 1620 kg/m^3 , and form a layer on the centrifuge wall 0.1 in thick. The rotational speed of the centrifuge is 15,000 rpm.
- 6. Estimate the cut point when treating 1 L/s of suspension in a hydrocyclone of 2 in diameter. The concentration of the suspension is 15% by volume, its density is 1250 kg/m³, and behaves as non-Newtonian with determined values of the flow behavior index of 1.28 and the fluid consistency index of 2.03×10^{-4} . The density of the suspended solids is 2800 kg/m^3 .
- 7. Table 10.15 was obtained during an experimental run in a laboratory filter operating at constant pressure. Calculate the values of specific resistances of cake and medium, and the time to produce 5 kg of filtrate under the following operation conditions: pressure 100 kN/m^2 , filtration area 0.05 m^2 , feed solids content 25 kg/m^3 , density of filtrate 1000 kg/m^3 , and viscosity of filtrate $8.84 \times 10^{-4} \text{ N s/m}^2$.
- 8. The values of specific cake resistance shown in Table 10.16 were calculated from the results of a number of experiments operating at different constant pressures. Determine the cake compression coefficient.

TABLE 10.16 Problem 8

α (m/kg)	$\Delta P (kN/m^2)$
1.10×10^{11}	46.2
1.44×10^{11}	117.2
1.68×10^{11}	194.5
1.79×10^{11}	318.6
2.01×10^{11}	338.5

References

- Bogue, D. C. and Metzner, A. B. 1963. Velocity profiles in turbulent pipe flow, Newtonian and non-Newtonian fluids. *Industrial and Engineering Chemistry Fundamentals* 2: 143–149.
- Bradley, D. 1965. The Hydrocyclone. Oxford: Pergamon Press.
- Brodkleg, R. S., Lee, J., and Chase, R. C. 1961. A generalised velocity distribution for non-Newtonian fluids. *AIChE Journal* 7: 392–393.
- Chen, W., Parma, F., Parkat, A., Elkin, A., and Sen, S. 2004. Selecting membrane filtration systems. *Chemical Engineering Progress* 100: 22–25.
- Clark, J. G. 1990. Select the right fabric for liquid–solid separation. *Chemical Engineering Progress* 86: 45–50.
- Dodge, D. W. and Metzner, A. B. 1959. Turbulent flow of non-Newtonian systems. *AIChE Journal* 5: 189–204.
- Doheim, M. A., Ibraheim, G. A., and Ahmed, A. A. 1985. Rapid estimation of corrected cut point in hydrocyclone classification units. *International Journal of Mineral Processing* 14: 149–159.
- Einstein, A. 1906. Eine neue bestimmung der moleküldimensionen. *Annalen der Physik* 324: 289–306.
- Einstein, A. 1911. Berichtigung zu meiner arbeit: eine neue bestimmung der moleküldimensionen. *Annalen der Physik* 339: 591–592.
- Fedoc, P. 1993. Air Classification, Encyclopedia of Food Science and Technology. London: Academic Press.
- Fitch, B. 1975. Current theory and thickening design, Parts 1–3. *Filtration and Separation* 12: 355–359, 480–488, 636–638.
- Frisch, H. L. and Simha, R. 1956. The viscosity of colloidal suspensions and macromolecular solutions. In *Rheology*, Vol. 1, Eirich, F. (ed.). Nueva York, NY: Academic Press
- Geankoplis, C. J. 2003. *Transport Processes and Separation Process Principles*. Upper Saddle River, NJ: Pearson Education Inc.
- Geldart, D. 1986. Gas Fluidization Technology. London: John Wiley and Sons.
- Girard, B. and Fukumoto, L. R. 2000. Membrane processing of fruit juices: A review. *Critical Reviews in Food Science and Nutrition* 40: 91–157.
- Green, D. W. and Perry, R. H. 2008. *Perry's Chemical Engineers' Handbook*, 8th Ed. New York, NY: McGraw-Hill.
- Guth, E. and Simha, R. 1936. Untersuchungen ubre die viskositat von suspensionen und losungen 3. Ubre die viskositat von kugel suspensionen. *Kolloid* 2 74: 266–275.
- Kelsall, D. F. 1966. The theory and applications of the hydrocyclone. In *Solid–Liquid Separation*, Poole, J. B. and Doyle, D. (eds.). London: Her Majesty Stationary Office (HMSO).
- Klumpar, I. V., Currier, F. N., and Ring, T. A. 1986. Air classifiers. *Chemical Engineering* 93: 77–92.
- Maas, J. H. 1979. Gas-solid separations. In *Handbook of Separation Techniques for Chemical Engineers*, Schweitzer, P.A. (ed.). New York, NY: McGraw-Hill.
- McCabe, W. L., Smith, J. C., and Harriot, P. 2005. *Unit Operations in Chemical Engineering*. 7th Ed. New York, NY: McGraw-Hill.

- Medronho, R. A. and Svarovsky, L. 1984. Test to verify hydrocyclone scale-up procedure. In *Proceedings of the 2nd International Conference on Hydrocyclones*, pp. 1–14. Cranfield, UK: BHRA The Fluid Engineering Centre.
- Metzner, A. B. and Park, M. G. 1964. Turbulent flow characteristics of viscoelastic fluids. *Journal of Fluid Mechanics* 20: 291–303.
- Metzner, A. B. and Whitlock, M. 1958. Flow behaviour of concentrated (dilatant) suspensions. *Transactions of the Society of Rheology* 2: 239–254.
- Nicodemo, L., Acierno, D., and Astarita, G. 1969. Velocity profiles in turbulent pipe flow of drag-reducing liquids. *Chemical Engineering Science* 24: 1241–1246.
- Ortega-Rivas, E. 1989. Dimensionless Scale-up of Hydrocyclones for Separation of Concentrated Suspensions. PhD Thesis Bradford, UK: University of Bradford.
- Ortega-Rivas, E. and Svarovsky, L. 1993. On the completion of a dimensinless scaleup model for hydrocyclone separation. *Fluid/Particle Separation Journal* 6: 104–109.
- Ortega-Rivas, E. and Svarovsky, L. 2000. Centrifugal air classification as a tool for narrowing the spread particle size distributions of powders. In *Processing Part I*, Wöhlbier, R. H. (ed.). Clausthal-Zellerfeld, Germany: Trans Tech Publications.
- Purchas, D. B. 1971. Industrial Filtration of Liquids. London: Leonard Hill Books.
- Purchas, D. B. 1981. Solid/Liquid Separation Technology. Croydon, UK: Uplands Press.
- Rietema, K. 1961. Performance and design of hydrocyclones. Parts I to IV. *Chemical Engineering Science* 15: 198–325.
- Seyer, F. A. and Metzner A. B. 1969. Turbulent phenomena in drag reducing systems. *AIChE Journal* 15: 426–434.
- Seyer, F. A. and Metzner, A. B. 1967. Turbulent flow properties of viscoelastic fluids. *Canadian Journal of Chemical Engineering* 45: 121–126.
- Shaver, R. G. and Merrill, E. W. 1959. Turbulent flow of pseudoplastic polymer solutions in straight cylindrical tubes. *AIChE Journal* 5: 181–188.
- Simha, R. 1952. A treatment of the viscosity of concentrated suspensions. *Journal of Applied Physics* 23: 1020–1024.
- Sourirajan, S. 1970. Reverse Osmosis. New York, NY: Academic Press.
- Svarovsky, L. 1979. The efficiency of separation processes. In *Progress in Filtration and Separation*, Wakeman, R. J. (ed.). Amsterdam: Elsevier Science.
- Svarovsky, L. 1981. Solid-Gas Separation. Amsterdam: Elsevier Science.
- Svarovsky, L. 1984. Hydrocyclones. London: Holt Reinhold and Winston.
- Svarovsky, L. 1985. *Solid–Liquid Separation Processes and Technology*. Amsterdam: Elsevier Science.
- Svarovsky, L. 2000. Efficiency of separation of particles from fluids. In *Solid–Liquid Separation*, Svarovsky, L. (ed.). London: Butterworths-Heinemann.
- Thomas, D. G. 1961. Transport characteristics of suspension II: Minimum transport velocity for flocculated suspension in horizontal pipes. *AIChE Journal* 7: 423–430.
- Thomas, D. G. 1962. Transport characteristics of suspension IV: Minimum transport velocity for large particles size suspension in round, horizontal pipes. *AIChE Journal* 8: 373–378.
- Thomas, D. G. 1963. Non-Newtonian suspensions. Part I: Physical properties and laminar transport characteristics. *Industrial and Engineering Chemistry* 55: 18–29.
- Trawinski, H. F. 1976. Die mathemat formulierung der Trump-kurve. *Aubereitungs-Technik* 17: 449–459.

- Trawinski, H.F. 1977. Hydrocyclones. In *Solid/Liquid Separation Equipment Scale-up*, Purchas, D. B. (ed.). Croydon, UK: Uplands Press Ltd.
- Trowbridge, M. E. O' K. 1962. Problems in the scaling-up of centrifugal separation equipment. *Chemical Engineering* 40: 73–86.
- Van Ebbenhorst Tengbergen, H. J. and Rietema, K. 1961. Efficiency of phase separation. In *Cyclones in Industry*, Rietema, K. and Verver, C. G. (eds.). Amsterdam: Elsevier Science.

Appendix 1: International (SI) System of Units

The system is based on the following seven fundamental units: meter (m), kilogram (kg), second (s), ampere (A), Kelvin (K), candela (cd), and mole (mol).

In addition, the following derived units are relevant in engineering applications: Newton (N) = $kg \text{ m/s}^2$, joule (J) = N m, watt (W) = J/s.

The following multiples and submultiples of each unit are also relevant: 10^9 , giga (G); 10^6 , mega (M); 10^3 , kilo (k); 10^2 , hecto (h): 10, deca (da); 10^{-1} , deci (d); 10^{-2} , centi (c); 10^{-3} , milli (m); 10^{-6} micro (m); 10^{-9} , nano (n).

Useful Conversion Factors

Quantity	Non-SI Unit	Conversion Factor	SI Unit
Length	1 foot (ft)	= 0.3048	m
Mass	1 pound (lb)	≈ 0.454	kg
Volume	1 ft ³	$\approx 2.83 \times 10^{-2}$	m^3
	1 gallon (gal)	$\approx 3.78 \times 10^{-3}$	m^3
Specific volume	1 ft³/lb	$\approx 6.24 \times 10^{-2}$	m³/kg
Force	1 pound force (lb force)	≈ 4.4 5	N
Pressure, stress	1 lb force/in² (psi)	≈ 6.90	kN/m^2
	1 in. Hg	≈ 3.39	kN/m^2
	1 mm Hg (≈1.00 Torr)	≈ 0.133	kN/m^2
	1 ft H ₂ O	≈ 2.99	kN/m^2
	1 bar (bar)	$= 1 \times 10^{-2}$	kN/m^2
Density	1 ton/yd³	= 1328.94	kg/m³
	1 lb/ft³	≈ 16.02	kg/m³
	1 lb/gal	≈ 99.78	kg/m³
Dynamic viscosity	1 lb/ft s	≈ 1.49	kg/m s
	1 poise (P)	= 0.1	kg/m s
Energy	1 kWh	= 3.6	MJ
	1 erg (erg)	= 0.1	J
Power	1 horsepower (hp)	= 745.700	W

This page intentionally left blank



Appendix 2: Density and Viscosity of Water at Different Temperatures and 101.325 kPa

Temperature (°C)	Density (kg/m³)	Viscosity [(Pa s)10 ³]
0	999.87	1.7921
4	1000.00	1.5674
10	999.73	1.3077
20	998.23	1.0050
30	995.68	0.8007
40	992.25	0.6560

This page intentionally left blank



Appendix 3: Density and Viscosity of Air at Different Temperatures and 101.325 kPa

Temperature (°C)	Density (kg/m³)	Viscosity [(Pa s)10 ⁵]
-17.8	1.379	1.62
0	1.293	1.72
10	1.246	1.78
37.8	1.137	1.90
65.6	1.043	2.03
93.3	0.964	2.15
121.1	0.895	2.27
148.9	0.838	2.37
176.7	0.785	2.50
204.4	0.740	2.60
232.2	0.700	2.71
260.0	0.662	2.80

This page intentionally left blank



Appendix 4: Dimensional Analysis

Any physical quantity comprises a number giving its magnitude, and a unit representing its physical meaning. Dimensional analysis is an algebraic treatment of the symbols for units considered independently of magnitude. Dimensional analysis facilitates fitting experimental data to design equations where a completely mathematical treatment is not possible. It is a very useful tool for verifying consistency of units in equations, for converting units, and for scale-up of data obtained in physical models to predict the performance of full-scale equipment. As an example, derivation of dimensionless groups describing operation of fluid mixers is presented.

As the performance characteristics of fluid mixers involve a great number of variables, the use of dimensionless groups in this method is an obvious advantage. The basic concepts underlying dimensionless analysis and scale-up procedures have been used since a long time in classical mechanics principally. Perhaps, the most common method of dimensional analysis is that attributed to Lord Raleigh known as the Pi theorem. The basic steps of dimensional analysis can be found elsewhere, but it is known that the procedure is based on the combination of possible variables of the problem together into groups in order to find the least number of dimensionless groups. Then, by the use of physical arguments and experimental data, relationships between the dimensionless groups are derived.

In the particular case of fluids mixing, if linear dimensions such as the depth of liquid in the tank, the diameter of the tank, as well as the number, dimensions, and position of baffles are all in a definite geometrical ratio with the impeller diameter, then the power input to the agitator can be expressed as a function of the following variables:

$$P = f(\mu, g, \rho, N, D) \tag{A1}$$

where μ is the viscosity of the liquid, g is the acceleration due to gravity, ρ is the density of the liquid, N is the rotational speed of the impeller, and D is the diameter of the impeller.

Dimensional analysis indicates that Equation A1 may be reexpressed to include powers in all the variables as follows:

$$P = k(\mu^a g^b \rho^c N^d D^e)$$
 (A2)

where *k* is a constant.

If dimensions are given in terms of fundamental units, that is, mass *M*, length *L*, and time *T*, then by substituting such units of the variables, Equation A2 would transform into

$$\frac{ML^2}{T^3} = k \left[\left(\frac{M}{LT} \right)^a \left(\frac{L}{T^2} \right)^b \left(\frac{M}{L^3} \right)^c \left(\frac{1}{T} \right)^d (L)^e \right]$$
 (A3)

By equating fundamental units on both sides of Equation A3, and by algebraic means, common powers are grouped together to give

$$P = k \left[\left(D^5 N^3 \rho \right) \left(\frac{D^2 N \rho}{\mu} \right)^a \left(\frac{D N^2}{g} \right)^b \right]$$
 (A4)

where *k*, *a*, and *b* depend on the system and its geometry. Equation A4 can be rewritten as follows:

$$\frac{P}{D^5 N^3 \rho} = k \left[\left(\frac{D^2 N \rho}{\mu} \right)^a \left(\frac{D N^2}{g} \right)^b \right] \tag{A5}$$

Equation A5 is often referred to as the power equation. The term on the left-hand side of the equation is known as the dimensionless power number, the first term after the constant k on the right-hand side of the equation is the dimensionless Reynolds number Re, while the last term of the equation is the dimensionless Froude number Fr.

A number of workers have applied Equation A5 to liquid mixing using impeller-type agitators in vertical cylindrical tanks. The Reynolds number represents the ratio of applied forces to the viscous drag forces. The Froude number represents the ratio of applied forces to gravity forces. Vortex formation is a gravitational effect so when suppressed, for example, by using baffles in the mixing tanks, or at very low velocities represented by low values of Reynolds number (<300) the contribution of the Froude number in Equation A5 can be neglected. The power equation for these cases is, therefore, simply represented as

$$\frac{P}{D^5 N^3 \rho} = k \left(\frac{D^2 N}{\mu}\right)^a \tag{A6}$$

Plots for power number against Reynolds number on log—log coordinates, the so-called power curves, are available in the literature for particular mixer configurations. Power curves are independent of vessel size and are particularly useful in the scale-up of liquid mixers from pilot plant studies. It must be considered, however, that a given curve is only applicable to the geometric configuration for which it was developed.

Appendix 5: Standard Screen Scales

US Sieve Series and Tyler Equivalents (ASTM Standards)

Sieve Designation		Sieve C	pening	Wire D	iameter	Tyler
Standard (mm)	Alternate (in.)	mm	in.	mm	in.	Equivalent Designation
107.6	4.24	107.6	4.2400	6.400	0.2520	
101.6	4.00	101.6	4.0000	6.300	0.2480	
90.5	$3\frac{1}{2}$	90.5	3.5000	6.080	0.2394	
76.1	3	76.1	3.0000	5.800	0.2283	
64.0	2 ½	64.0	2.5000	5.500	0.2165	
53.8	2.12	53.8	2.1200	5.150	0.2028	
50.8	2	50.8	2.0000	5.050	0.1998	
45.3	$1\frac{3}{4}$	45.3	1.7500	4.850	0.1909	
38.1	$1\frac{1}{2}$	38.1	1.5000	4.590	0.1807	
32.0	$11\frac{1}{4}$	32.0	1.2500	4.230	0.1665	
26.9	1.06	26.9	1.0600	3.900	0.1535	1.050 in.
25.4	1	25.4	1.0000	3.800	0.1496	
22.6	7/8	22.6	0.8750	3.500	0.1378	0.883 in.
19.0	3/4	19.0	0.7500	3.300	0.1299	0.742 in.
16.0	5/8	16.0	0.6250	3.000	0.1181	0.624 in.
13.5	0.530	13.5	0.5300	2.750	0.1083	0.525 in.
12.7	1/2	12.7	0.5000	2.670	0.1051	
11.2	7/16	11.2	0.4380	2.450	0.0965	0.441 in.
9.51	3/8	9.51	0.3750	2.270	0.0894	0.371 in.
8.00	5/16	8.00	0.3120	2.070	0.0815	$2\frac{1}{2}$ mesh
6.73	0.265	6.73	0.2650	1.870	0.0736	3 mesh
6.35	1/4	6.35	0.2500	1.820	0.0717	
			Sieve no.			
5.66	3 ½	5.66	0.2230	1.680	0.0661	33½ mesh
4.76	4	4.76	0.1870	1.540	0.0606	4 mesh
4.00	5	4.00	0.1570	1.370	0.0539	5 mesh
3.36	6	3.36	0.1320	1.230	0.0484	6 mesh
2.83	7	2.83	0.1110	1.100	0.0430	7 mesh
2.38	8	2.38	0.0937	1.000	0.0394	8 mesh
2.00	10	2.00	0.0787	0.900	0.0354	9 mesh
1.68	12	1.68	0.0661	0.810	0.0319	10 mesh

continued

(continued)US Sieve Series and Tyler Equivalents (ASTM Standards)

Sieve Designation		Sieve C	pening	Wire D	iameter	Tyler
Standard (mm)	Alternate (in.)	mm	in.	mm	in.	Equivalent Designation
1.41	14	1.41	0.0555	0.725	0.0285	12 mesh
1.19	16	1.19	0.0469	0.650	0.0256	14 mesh
1.00	18	1.00	0.0394	0.580	0.0228	16 mesh
Micrometers						
841	20	0.841	0.0331	0.510	0.0201	20 mesh
707	25	0.707	0.0278	0.450	0.0177	24 mesh
595	30	0.595	0.0234	0.390	0.0154	28 mesh
500	35	0.500	0.0197	0.340	0.0134	32 mesh
420	40	0.420	0.0165	0.290	0.0114	35 mesh
354	45	0.354	0.0139	0.247	0.0097	42 mesh
297	50	0.297	0.0117	0.215	0.0085	48 mesh
250	60	0.250	0.0098	0.180	0.0071	60 mesh
210	70	0.210	0.0083	0.152	0.0060	65 mesh
177	80	0.177	0.0070	0.131	0.0052	80 mesh
149	100	0.149	0.0059	0.110	0.0043	100 mesh
125	120	0.125	0.0049	0.091	0.0036	115 mesh
105	140	0.105	0.0041	0.076	0.0030	150 mesh
88	170	0.088	0.0035	0.064	0.0025	170 mesh
74	200	0.074	0.0029	0.053	0.0021	200 mesh
63	230	0.063	0.0025	0.044	0.0017	250 mesh
53	270	0.053	0.0021	0.037	0.0015	270 mesh
44	325	0.044	0.0017	0.030	0.0012	325 mesh
37	400	0.037	0.0015	0.025	0.0010	400 mesh

British Standard Sieves

Nominal Aperture Size (mm)	Nominal Wire Diameter (mm)	Nearest Mesh Number
16.0	3.15	
13.2	2.80	
11.2	2.50	
9.50	2.24	
8.00	2.00	
6.70	1.80	
5.60	1.60	3
4.75	1.60	31/2
4.00	1.40	4

continued

(continued)British Standard Sieves

Nominal Aperture Size (mm)	Nominal Wire Diameter (mm)	Nearest Mesh Number		
3.35	1.25	5		
2.80	1.12	6		
2.36	1.00	7		
2.00	0.90	8		
1.70	0.80	10		
1.40	0.71	12		
1.18	0.63	14		
1.00	0.56	16		
Micrometers				
850	500	18		
710	450	22		
600	400	25		
500	315	30		
425	280	36		
355	224	44		
300	200	52		
250	160	60		
212	140	72		
180	125	85		
150	100	100		
125	90	120		
106	71	150		
90	63	170		
75	50	200		
63	45	240		
53	36	300		
45	32	350		
38	30	400		

This page intentionally left blank



Appendix 6: Guidelines for the Preparation of Laboratory Reports

The most common means of technical communication within and among organizations of all sorts, including industry, educational institutions, and professional societies, is the written technical report or article. In any professional activity, the writing and submission of reports is most certainly required. The technical report gives an insight into the writer's competences, abilities, and skills. It also reflects the writer's work habits and productivity. Regardless of the quality of the work reported, a poorly written report will reflect unfavorably on the writer.

Introductory Aspects

It is impossible to describe a standard format for the organization and contents of a given report. High-quality well-written reports contain, however, the following common components:

- Clearness of development
- Conciseness of expression
- Completeness of information
- Accuracy of details
- Correctness of form
- Convenience of layout

The details of format and content should be selected in full consideration of the reader's point of view and the purpose of the report. This criterion generally applies to classroom reports, as well as those expected to be produced after graduation in all types of arenas and scenarios.

The Importance of Neatness

From the days when typewriters were used, or even handwriting, the key characteristic of a written report has been its neatness and readability. In earlier days, decisions had to be made on whether to correct or amend parts of a document in the best possible way without sacrificing neatness, or rewrite a whole page or part. Nowadays, the ease of resourcing to the famous "copy-paste" practice, and some other similar alternatives, has created a trend to abuse of such routines so as to present bulky and apparently complete reports. Regardless of the audience to whom the document is written, reports are to be read by panels of experts in their fields of interest. Direct copy-paste inserts and some other editorial anomalies, in content as well as in format, will be detected by the trained eye. It is advisable to prepare figures and inserts with the text processor being used in writing, as well as to give uniformity of style to imported ideas from other texts.

Use of the English Language

The triumph of the Allies in World War II, along with a series of social and cultural events mainly in the twentieth century, has made English the undisputable recognized international language. Thus, writing skills in English are paramount since business, culture, and among all, science interchange, are made based on English written reports, articles, books, media presentations, and so on. Students in higher education programs all over the world may need to write technical reports in their own language, but after graduating, the importance of acquiring abilities to write in English increases significantly in order to gain access to more opportunities for professional development.

In technical and scientific reports it is good practice to use short, precise sentences and make sure that the text is written in sentences and not in phrases. Principles for punctuation and abbreviations should be used at all times. It is important to pay particular attention to spelling, sentence structure, agreement between subject and verb, and the use of consistent tense. It is also relevant to make sure that misinterpretation and unintended meanings are minimized. Once completed, the report is to be read thoroughly several times for assessing content and organization, as well as to perform a proof-reading. It is advisable to ask a classmate to read and criticize the report. Even the most professional authors usually have to rewrite parts of a written document.

Report Organization and Content

Title Page

The title page should be the first page of a formal report. The title should state briefly and completely the subject of the report. The name or organization or person for whom the report was prepared, the name of the individual or organization submitting the report, and the date should appear on the title page. Short or informal reports do not require a separate title page. In this instance, the title, the name of the writer, and the organization recipient of the report, should appear on the first page together with the abstract.

Table of Contents

Lengthy and formal reports require a table of contents, while short and informal reports do not. Most of the reports to be written by students will fall somewhere between these two extremes and, thus, the decision on whether or not to include a table of contents will be a matter of judgment. Whenever a table of contents will significantly clarify and demonstrate the continuity of the report, help the reader understanding the scope, or make it more convenient for the reader to find any point of interest, such a table should be included.

A table of contents should list, by title and page number, the divisions and subdivisions of the report in the order they appear, including all appendices. Tables and figures should be identified by number, title, and page in separate lists immediately following the table of contents. Page numbers, titles, and all the components of each item shown in the table of contents should be double-checked in a proofreading against the title, page, and so on, where the item occurs in the report.

Abstract

This section of the report is the most important individual section. The objective of the abstract is to provide the reader with a concise, overall statement of the high points of the report. Normally, the abstract should be no more than one written page. More than any other section of the report, the abstract should be prepared in consideration of the reader and the purpose of the own report. A reasonable assessment of the abstract will be, therefore, to read it from the reader's point of view and determine if is sufficiently clear and concise and conveys the appropriate information on the purpose, scope, and extent of the report. A good abstract will include a brief statement of the objectives of the study, experiment, or design, and results in the form on concise conclusions and/or recommendations for further work. Often, other information, such as important assumptions, alternatives analyzed, or procedures will be included depending on the purpose of the report and the reader's profile.

Introduction

The purpose of the introduction is to give the reader the minimum amount of information needed to understand the result, conclusions, and recommendations. Such information normally includes purpose (problem statement), scope, and nature of the study. A formal introduction may not be required for short reports where minimal background is required for understanding. A simple introductory statement will be sufficient in cases such as this.

Objectives

This section of the report should itemize in a brief, but clear manner, all the objectives of the work performed. No detailed description of each objective is required at this point and the best format is probably a simple listing of line items.

Background

The contents of this section, depends upon the nature of the work being reported. In general, theoretical principles related to the subject under study are to be explained and briefly discussed. A literature survey, as current as possible but balanced with classical references if appropriate, is a fundamental part of this section. Any derivations and equations used in the development of the work reported should be included in this part of the report. If alternate approaches were performed, a justification should be also given in this section. Assumptions made in the development of the work, should be stated and sufficiently supported.

Materials and Methods

A description of the equipment, instruments, and apparatuses used to perform the work should be included in this section. Raw materials used, routine procedures employed, and glassware along with consumables utilized, may be simply listed. Quite often, neat schematic diagrams of equipment and instruments will greatly enhance the clarity of the written text. The experimental methods should be clearly described, with reference given to standard methods like those reported in analytical chemists handbooks. The aim is for the reader to understand fully how the raw data were collected.

Results and Interpretation of Data

The result obtained from the work should be presented in this section. It is important to point out that these results are not the raw data collected from the methods described in the previous section, but the final values resulting from the computational analysis based on the raw data collected. When

possible, the results should be presented in tabular or graphical form. Table and/or figures cannot constitute, however, this section on their own. Some explanatory text must be incorporated to lead the reader through the results. Whenever possible, a statistical analysis on the data collected should be performed and the results included in this section.

Discussion

This section of the reports should include a critical analysis of the results obtained. It is appropriate at this stage to discuss items such as accuracy of the data, limitations imposed by the assumptions, comparisons with results available in the literature, operating problems due to availability of equipment, instruments, and infrastructure, and so on. Compatible comparisons with results in the literature are important in this section, so that few or several references will be appropriate.

Conclusions

The logical deduction of conclusions from the results should be clearly demonstrated. An itemization of conclusions from the work performed may be an appropriate format for this section.

Recommendations

A list of specific suggestions on how to improve the quality of the work reported is appropriate after the section of conclusions. These recommendations should address items such as experimental design, equipment modifications, sample collection systems, need for further work, and so on. Conclusions and Recommendations may conform into a single section.

References, Literature Cited

Use of cited references is an important part of a technical report. Cited literature is a basic foundation for the reader to realize that the report makes sense within the state-of-the-art of the related discipline. For this reason, normally all the literature is cited in a technical report and, as such, the proper denominations should be References or Literature Cited. Bibliography is usually not an appropriate title for this section. Some textbooks and encyclopedias often refer to suggested readings or bibliography, that have not been cited in the text, for the reader to get further information on some topics discussed. In this section, each reference cited in the body of the report should be listed in detail, so that an interested reader can find the article, book, or report. There are several references styles used in scientific literature, such as the American Chemical Society style or the Harvard style. As long as the references listed are consistent, any scientific style can be used.

Appendices

The body of the report should contain only those data, equations, drawings, or calculations, which pertain directly to results of the work. This information is often the result of much detailed data, computations, derivations, and so on, and it is usually desirable to place the latter in appendices to the main report. Specific examples of information that should be placed in appendices are raw data from experiments, detailed derivation of an equation, example computations, statistical tables of data interpretation such as analyses of variance, and so on. It is good practice to include sufficiently detailed information in appendices to allow the reader to check computations or conclusions of the report.

Illustrative Aids

Tables, charts, graphs, drawings, and photographs are valuable aids for presenting technical information and should be considered an integral part of the report. As such, they must be planned and prepared in rough form as the text itself is planned and developed. Illustrative aids, while based on the text, should be as nearly self-explanatory as possible, and reliance upon the text for clarification of illustrations should be kept to a minimum. Every illustration should be accompanied by a caption, which makes clear the nature of the contents and the meaning or significance of the information. Tables and figures should be identified by number and title. Reference to a particular table or figure in the text should be properly accomplished as, for example: Table 4 or Figure 5.

Tables

Tables should be prepared so that the arrangement maximizes clarity. Only relevant data should be included. Titles should be clear and complete, and placed at the top of the tables. It is permissible to use footnotes, which apply to specific portions of the table, immediately below it. Column headings should clearly identify the data in the column and include units in which the numbers are expressed. Columns and rows should be placed to facilitate ease of reading.

Drawings and Diagrams

It is important to make sure that all drawings and diagrams have a figure number and an appropriate caption or title. All pertinent parts of these figures should be clearly labeled and dimensioned. Overcrowding of drawings is to be avoided, so neatness and readability are satisfactory. As stated earlier, crude paste of imported figures is noticeable, while it demerits the quality of the report. It is preferable to make use of the available tool for graphics in the word processor, or to insert figures prepared in a compatible graphics elaboration system.

Charts and Graphs

Data, numerical results, and some other information should be presented in charts and graphs when possible. This type of presentation is usually superior to a tabular presentation because trends, comparisons, and relationships are more easily identified. Charts and graphs should be presented on a grid. The scale of the ordinate and the abscissa should be clear. It is recommended practice to measure the independent variable on the abscissa and the dependent variable on the ordinate. Both axes should be clearly labeled with the variable name, symbol, and units. If more than one curve is plotted, the conditions for each one should be identified. It is usually best to use different symbols and/or different types of lines for different curves. Colored lines or markers are not recommended because of problems in reproduction of reports. It is advisable to plot measured data at discrete points and reserve the use of continuous curves for theoretical or best-fit relations. Bar graphs are very useful in presenting data for which the relationship between variables is discontinuous.

Equations

Formulas and equations are often a very important component of a technical report. It is essential that all variables, exponents, parameter, and so on, be clearly defined immediately following the equation. The units of each component of an equation should be stated. Use of SI units is preferred. If the formula is written so that it is valid for any consistent set of units, then the fundamental dimension, for example, length or force, of each symbol should be listed. For lengthy reports including many equations, a list of nomenclature placed normally right after the list of tables and figures or right before the references, is recommended.

This page intentionally left blank



Appendix 7: Notation

Dimensions given in terms of mass, M; length, L; time, T; and temperature, θ

а	Arithmetic mean, screen aperture, acceleration (L/T^2) , radius of dust particle (L)
$a_{\rm c}$	Accelerations convergence component due to flow channel in silo (L/T^2)
$a_{\rm v}$	Acceleration component due to velocity increase as flow is initiated in silo (L/T^2)
A	Area (L^2), area of the collecting electrodes in Equation 10.46 (L^2)
$A_{\rm a}$	Arithmetic mean of cake area in filtering centrifuge (L^2)
A_1	Logarithmic mean of cake area in filtering centrifuge (L^2)
$A_{\rm m}$	Area of filter medium in filtering centrifuge (L^2)
A_{p}	Projected area of particle (L^2)
Ar	Archimedes number
A_{i}	Adhesive force due to particular binding mechanism (ML/T^2)
A_{1n}	Parameter dependent on the flow behavior index in Equation 10.57
b	Constant, height of bowl of tubular centrifuge (L)
b_1	Empirical constant
В	Hopper opening (<i>L</i>), intercept with <i>y</i> -axis in plot of constant pressure filtration run (T/L^3)
B'	Intercept with y -axis in plot of constant rate filtration run (M/LT^2)
C	Cohesion (M/L^2) , capacity (M/T) , volume fraction of solids in suspension
$C_{\rm f}$	Volume fraction of solids in feed
$C_{\rm u}$	Volume fraction of solids in underflow
Coh	Cohesion number
C_{D}	Drag coefficient
C_n	Parameter dependent on the flow behavior index in Equation 10.57
d	Diameter of rod supporting rollers in chain conveyors (L), radius of product in roller mills (L)
$d_{\rm c}$	Coat thickness (L)
$d_{\rm s}$	Spout diameter in spouted bed (<i>L</i>)
D	Fractal dimension, roller diameter in chain conveyors (L), diameter (L), diffusion coefficient (L^2/T)
D_c	Cyclone and hydrocyclone diameter (L)
$D_{\rm f}$	Diameter of feed in roller mills (<i>L</i>)
D_{i}	Diameter of inlet gas orifice in spouted beds (<i>L</i>), inlet diameter of hydrocyclone (<i>L</i>)
$D_{\rm o}$	Overflow pipe diameter of hydrocyclone (<i>L</i>)
D_{p}	Liquid droplet diameter (L)
DM	Dry matter content
$D_{\rm p}$	Diameter of product in roller mills (<i>L</i>)

continued

(continued)

(continue)	<u> </u>
$D_{\rm pc}$	Cut diameter (L)
$D_{\rm r}$	Diameter of roll in roller mills (L)
D_{t}	Pipe diameter (L)
$D_{\rm u}$	Underflow diameter of hydrocyclone (L)
Ε	Energy (ML^2/T^2), overall screen efficiency
$E_{\rm a}$	Acceptance efficiency
$E_{\rm c}$	Coating efficiency
E_{fb}	Friction energy due to flow round bends expressed as pressure drop in pneumatic conveying (M/LT^2)
$E_{ m fg}$	Friction energy due to gas flow expressed as pressure drop in pneumatic conveying (M/LT^2)
E_{fp}	Friction energy due to particle flow expressed as pressure drop in pneumatic conveying (M/LT^2)
$E_{\rm f}$	Net efficiency of fines
$E_{\rm i}$	Bond work index (ML^2/T^2)
$E_{\mathbf{k}}$	Kinetic energy expressed as pressure drop in pneumatic conveying (M/LT^2)
$E_{\rm O}$	Oversize screen efficiency
$E_{\rm p}$	Potential energy expressed as pressure drop in pneumatic conveying (M/LT^2) , partial efficiency, strength of electrical field in settling point (V^a/T)
$E_{\rm r}$	Rejection efficiency
$E_{\rm s}$	Screen efficiency
E_{t}	Total efficiency
$E_{ m t}^{'}$	Reduced total efficiency
Eu	Euler number
E_{U}	Undersize screen efficiency
E_0	Strength of electrical field (V^a/L)
f	Frequency (dimensionless) frequency $(1/T)$, fanning friction factor, ratio of filtration to cycle time in Equation 10.125
$f_{\rm a}$	Size fraction of one component of average weight $w_{\rm a}$
$f_{\rm b}$	Bend friction factor
f_{g}	Gas friction factor
$f_{\rm p}$	Particle friction factor
ff	Flow factor
ff_a	Actual flow factor based on the actual opening dimension
$f_{am}(x)$	Modified beta distribution given by Equation 2.27
$f_{\rm pv}$	Purely viscous fanning friction factor
f(x)	Frequency or occurrence related to size
$f_{\rm L}(x)$	Size distribution function by length
$f_{\rm M}(x)$	Size distribution function by mass (volume)
$f_{\rm N}(x)$	Size distribution function by number
$f_{\rm S}(x)$	Size distribution function by surface
$f(\delta)$	Constant related to angle of contact of particles in agglomerates
F	Mass flow rate of solids in feed (M/T) , force (ML/T^2) , cumulative percentage of
	coarse fraction in the feed

(continue	1)
$F_{\rm c}$	Coefficient of friction for chain and flights, cumulative oversize percentage
$F_{\rm b}$	Buoyancy force (ML/T^2)
$F_{ m f}$	Cumulative percentage of fines fraction in the feed
$F_{\rm g}$	Gravity force (ML/T^2)
$F_{\mathbf{m}}$	Coefficient of friction for the material
$F_{\rm r}$	Radial force (ML/T^2)
F_{t}	Tangential force (ML/T^2)
FF	Failure function
FF_{v}	Failure function at vibration conditions
F(x)	Cumulative frequency
$F_{ m D}$	Drag force (ML/T^2)
F(x)	Cumulative percentage oversize of feed solids
Fl	Fluidization number
Fl*	Fluidization number for powder at consolidated state
8	Acceleration due to gravity (L/T^2)
g _L	Local gravity acceleration (L/T^2)
G(x)	Function of particle size
G	Gas mass flow rate (M/T) , gas flux (M/L^2T)
$G_{\rm c}$	Critical flux (M/L^2T)
$G_{\rm s}$	Solids flux (M/L^2T)
G(x)	Grade efficiency
G'(x)	Reduced grade efficiency
h	Height (L)
H	Height or depth (<i>L</i>)
H_{mf}	Height of bed at minimum fluidization velocity (<i>L</i>)
$H_{\rm sm}$	Maximum spoutable bed depth (<i>L</i>)
$H_{\rm s}$	Height of gravity settling chamber
$H_{25/75}$	Sharpness index
$H(\theta)$	Factor to account for variation in arch thickness, hopper angle, and hopper type
J	Flux through membrane (M/L^2T)
k	Constant, coordination number
k_1, k_2	constants
K	Constant, correlation constant of the power law, slope of line in plot of constant pressure filtration run (T/L^6)
$K_{\rm c}$	Constant depending on the filtering medium in air filtration
K_1	Cake resistance factor in air filtration
K'	Ratio of horizontal to vertical pressure in silo, fluid consistency index (MT^n/L^2), slope of line in plot of constant rate filtration run (M/L^4T^2)
K"	Constant in constant rate filtration defined by Equation 10.107
L	Length of confidence interval, length (L)
$L_{\rm c}$	Horizontal projected length of conveyor (L)
$L_{\rm s}$	Estimated length (<i>L</i>), length of gravity settling chamber (<i>L</i>)
L_0	Constant in Equations 3.17 and 3.18
	continue

(continued	1)
m	Measure of distribution slope (Schuhmann slope), mass (M)
$m_{\rm c}$	Deposited coating mass (M)
$m_{\rm cs}$	Mass of coating solvent (M)
m_0	Weight of empty pycnometer (<i>M</i>)
m_1	Weight of pycnometer filled with liquid (<i>M</i>)
$m_{\rm s}$	Weight of pycnometer filled with solid (<i>M</i>)
$m_{\rm sl}$	Weight of pycnometer filled with liquid and solid (<i>M</i>)
$m_{\rm LC}$	Weight of container partially filled with liquid (M)
$m_{\rm LCS}$	Weight of container with liquid and submerged solid (M)
M	Mesh size, mass flows rate of solids in suspension (M/T) , molar concentration
$M_{\rm c}$	Mass flows rate of separated solids (M/T) , the mass of solid cake in bowl of filtering centrifuge (M)
$M_{ m f}$	Mass flows rate of unseparated solids (M/T)
$M_{\rm s}$	Weight of specimen (M)
M_1	Mixing index
M_2	Mixing index
M_3	Mixing index
N	Rotation speed $(1/T)$, sample size
$N_{\rm cr}$	Critical rotation speed $(1/T)$
$N_{ m WR}$	Relative wear number
n	Power, shear index, number of particles in sample, slope
n'	Flow behavior index
0	Mass flow rate of solids in overflow (M/T)
p	Proportion by weight of component within a total sample weight \boldsymbol{w}
$p_{\rm h}$	Horizontal pressure in silo wall (M/LT^2)
$p_{ m v}$	Vertical pressure in silo wall (M/LT^2)
P	Pressure (M/LT^2), power (ML^2/T^3), charge density at a given time in Equation 9.10 (C^b/L^3)
P_0	Initial charge density in Equation 9.10 (C^b/L^3)
Q	Proportion by weight of component within a total sample weight (M), volumetric flow rate (L^3/T)
$Q_{\rm B}$	Bubble flow rate (L^3/T)
$Q_{\rm mf}$	Gas flow rate at incipient fluidization (L^3/T)
Q_0	Discharge mass flow rate from silo (M/T)
r	Ratio of x_0 to size modulus, radius of particle in roller mills, radius of feed particle in roller mills (L), radius of ball in tumbling mills (L), radius (L)
R	Radius (L), radius of roll in roller mills (L), radius of drum in tumbling mills (L), radius of rotation (L), universal gas constant, bed resistance in Equation 10.123 (1/ L)
$R(\theta)$	Radius vector rotated about the origin
Re	Reynolds number
Re_{mf}	Reynolds number at incipient fluidization point
Re _p	Particle Reynolds number

Particle Reynolds number for the terminal velocity of the particle in the fluid
Reynolds number for power-law fluids
Underflow-to-throughput ratio
Resistance of filter medium $(1/L)$
Outer radii of stack of discs in disc-bowl centrifuge (L)
Inner radii of stack of discs in disc-bowl centrifuge (L)
Step length (L), gradient of pressure drop (M/LT^2), standard deviation of the analyses about the average value of the fraction of a specific powder, specific surface of solid (L^2), empirical constant, compressibility coefficient in Equation 10.98
Surface area of particle (L^2)
Estimate of standard deviation of fraction of specific powder at complete randomization
Estimate of standard deviation of fraction of specific powder at complete segregation
Empirical constant
Surface (L^2), population standard deviation, number of discs in stack in disc-bowl centrifuge
Separation number
Surface area of particle (L^2)
Specific surface area of the solid particles (L^2)
Stokes number
Stokes number including the reduced cut size
Stokes number for power-law fluids
Stokes number for power-law fluids including the reduced cut size
Temperature (θ), tensile stress (M/LT^2), conveyor capacity (M/T)
Time (<i>T</i>)
Cycle time in filtration (T)
Settling time (<i>T</i>)
Relaxing time (<i>T</i>)
Linear velocity in vertical direction (L/T) , upwards or downwards velocity in th open section below the grid or in empty tower above bed (L/T) , fluid–particle relative velocity (L/T) , settling velocity at concentration C (L/T)
Minimum bubbling velocity (L/T)
Minimum fluidization velocity (L/T)
Minimum spoutable velocity (L/T)
Saltation velocity (L/T)
Minimum conveying velocity for a single particle (L/T)
Terminal settling velocity of particles (L/T) , channel velocity calculated from the cross section (L/T)
Air velocity in pneumatic conveying (L/T), superficial velocity or empty-tower velocity (L/T)
Parameter related to conveying velocity in pneumatic conveying (Figure 3.33, Equation 3.27)
continue

(Contir	ided)
U	Estimator of sample size, constant in Equation 3.12 (L/T), mass flow rate of solids in underflow (M/T), volumetric flow rate of underflow (L^3/T)
v	Linear velocity in horizontal direction (L/T)
v_{g}	Terminal settling velocity under gravity (L/T)
$v_{\rm i}$	Inlet tangential velocity (L/T)
$v_{\rm o}$	Maximum tangential velocity (L/T)
$v_{\rm r}$	Radial settling velocity (L/T)
v_{t}	Radial speed of air (L/T)
$v_{\rm tan}$	Tangential velocity (L/T)
V	Volume (L^3), compacted volume of solids at a given pressure (L^3), linear velocity (L/T), volumetric flow rate of air (L^3/T)
$V_{\rm a}$	Average terminal discharge velocity from silo (L/T)
$V_{ m f}$	Gas superficial velocity through the filtering medium in air filtration (L/T)
V_{p}	Volume of particle (L^3)
$V_{ m R}$	Relation of volumes (V/Vs)
$V_{ m s}$	Volume of the solid material to be compacted (L^3)
V_0	Sliding velocity at a reference location defined by Equation 3.16 (L/T), initial volume (L^3), mean gas velocity (L/T)
w	Wire diameter (L), weight of sample (M), moisture content at time t , powder mass flow rate (M/T), migration velocity in Equation 10.46 (L/T), mass of solids deposited on the medium per unit volume of filtrate (M/L^3)
$w_{\rm a}$	Average weight of sample (<i>M</i>)
$w_{\rm s}$	Surface moisture content
w_1	Initial moisture content
W	Load on feeder (M/LT^2), weight of all moving parts of the conveyor in kilograms per meter of overall length, weight of flights and chain in kilograms per meter of overall length
x	Particle size
x_a	Projected area diameter (L)
$x_{\rm A}$	Equivalent sieve diameter
\overline{x}_a	Arithmetic mean of particle size (L)
$x_{\rm c}$	Cut point (L)
χ_{F}	Equivalent Feret diameter (L)
$x_{\rm g}$	Geometric mean of particle distribution; mass median size of particle (L)
$x_{\rm i}$	Every measured value of fraction of one powder
$x_{\rm m}$	Mode of particle size distribution
$x_{\rm M}$	Equivalent Martin diameter (L)
$x_{\rm o}$	Maximum particle size in sample (L)
$x_{\rm p}$	Diameter of particle (L)
$x_{\rm r}$	Constant function of particles size range (L), relative amplitude (L)
$x_{\rm R}$	Constant giving a measure of the present particle size range (L),
$x_{\rm st}$	Equivalent Stokes diameter
$x_{\rm v}$	Equivalent diameter of sphere having the same volume of particle (<i>L</i>)
x_0	Parameter related to maximum particle size. Particle size with zero centrifugal efficiency (L)

(continue	<u>لا</u>
<i>x</i> ₅₀	Cut size (L)
x_{98}	Approximate particle size fully separated (<i>L</i>)
x_{p}^{*}	Parameter related to particle size in pneumatic conveying (Equation 3.26)
x' ₅₀	Reduced cut size (L)
X	Fraction
$X_{\rm F}$	Mass fraction of coarse particles in feed
$X_{\rm i}$	Assayed value
$X_{\rm O}$	Mass fraction of coarse particles in the overflow
$X_{\rm r}$	Value of particular amplitude defined in Equation 3.13 (L)
X_{U}	Mass fraction of coarse particles in the underflow
$ar{X}$	Sample arithmetic average
y	Probability density
Y	Cumulative weight fraction undersize, solids yield in Equation $10.125 (M/L^2T)$
$Z_{(1-\alpha)/2}$	$(1-\alpha)/2$ percentile of standard Normal variable
Greek lette	rs
α	Angle of repose, half the angle of nip in roller mills, surface tension of liquid (M/LT^2) , degree of separation, specific cake resistance (L/M)
$\alpha_{\rm s}$	Surface factor defined in Equation 1.3
$\alpha_{\rm v}$	Volume factor defined in Equation 1.2
α_0	Empirical constant in Equation 10.98
α_0'	Empirical constant in Equation 10.99
β	Empirical constant in Equation 10.99
β′	Parameter in Equation 3.50
γ	Viscosity coefficient for power-law fluids (M/LT^{2-n})
γ_n	Volume reduction fraction
γ	Shear rate $(1/T)$
δ	Effective angle of internal friction, membrane thickness (<i>L</i>)
ΔP	Pressure drop (M/LT^2) , pressure drop across fixed bed (M/LT^2)
$\Delta P_{ m B}$	Pressure drop across fluidized bed (M/LT²)
$\Delta P_{ m c}$	Pressure drop through powder layer in air filtration (M/LT^2)
$\Delta P_{\rm eq}$	Pressure drop equal to buoyant weight of particles per unit area of bed (M/LT^2)
$\Delta P_{ m f}$	Pressure drop across the filtering medium of air filter (M/LT^2)
$\Delta P_{ m SB}$	Total spouting pressure drop at maximum spoutable bed depth (M/LT^2)
$-\Delta P$	Total pressure drop across the filter (M/LT^2)
$-\Delta P_c$	Pressure drops across the cake (M/LT^2)
$-\Delta P_{\rm m}$	Pressure drops across the medium (M/LT^2)
ε	Porosity or voidage, permeability in Equation 9.10
$\epsilon_{\scriptscriptstyle m B}$	Fraction of bed occupied by bubbles
$\epsilon_{ m mf}$	Voidage of bed at minimum fluidization velocity
η	Efficiency of separation in gravity settling chamber
θ	Hopper half-angle, hydrocyclone cone angle

	(001101111	,
$\begin{array}{lll} \mu_{\rm g} & {\rm Gas\ viscosity\ }(M/LT) \\ \mu_{\rm o} & {\rm Viscosity\ } of\ pure\ solvent\ }(M/LT) \\ \mu' & {\rm Sliding\ } friction\ coefficient\ in\ silo,\ friction\ coefficient\ in\ roller\ mills \\ \pi & {\rm Osmotic\ } pressure\ }(M/LT^2) \\ \rho_{\rm p} & {\rm Liquid\ } {\rm density\ }(M/L^3) \\ \rho_{\rm p} & {\rm Air\ } {\rm density\ }(M/L^3) \\ \rho_{\rm b} & {\rm Bulk\ } {\rm density\ }(M/L^3) \\ \rho_{\rm g} & {\rm Gas\ } {\rm density\ }(M/L^3) \\ \rho_{\rm g} & {\rm Gas\ } {\rm density\ }(M/L^3) \\ \rho_{\rm p} & {\rm Fluid\ } {\rm density\ }(M/L^3) \\ \rho_{\rm p} & {\rm Particl\ } {\rm density\ }(M/L^3) \\ \rho_{\rm p} & {\rm Particl\ } {\rm density\ }(M/L^3) \\ \rho_{\rm p} & {\rm Particl\ } {\rm density\ }(M/L^3) \\ \rho_{\rm p} & {\rm Solids\ } {\rm density\ }(M/L^3) \\ \rho_{\rm p} & {\rm Solids\ } {\rm density\ } {\rm under\ } {\rm applied\ } {\rm normal\ } {\rm stress\ } (M/L^3) \\ \sigma & {\rm Standard\ } {\rm deviation\ } \\ \sigma_{\rm c} & {\rm Unconfined\ } {\rm yield\ } {\rm strength\ } (M/LT^2) \\ \sigma_{\rm t} & {\rm Strength\ } {\rm o\ } {\rm agglomerates\ } (M/LT^2) \\ \sigma_{\rm t} & {\rm Strength\ } {\rm o\ } {\rm agglomerates\ } (M/LT^2) \\ \sigma_{\rm t} & {\rm Mayor\ } {\rm consolidation\ } {\rm stress\ } {\rm awll\ } {\rm surface\ } (M/LT^2) \\ \hline \Sigma & {\rm Operator\ } {\rm meaning\ } "{\rm algebraic\ } {\rm sum\ } {\rm o\ } f." {\rm characteristic\ } {\rm geometrical\ } {\rm features\ } {\rm o\ } {\rm centrifuge\ } {\rm $	μ	Liquid absolute viscosity (M/LT) , suspension viscosity (M/LT)
 μ_o Viscosity of pure solvent (M/LT) μ' Sliding friction coefficient in silo, friction coefficient in roller mills π Osmotic pressure (M/LT²) ρ Liquid density (M/L³) ρ_a Air density (M/L³) ρ_b Bulk density (M/L³) ρ_c Gas density (M/L³) ρ_m Slurry density (M/L³) ρ_m Slurry density (M/L³) ρ_p Particle density (M/L³) ρ_s Solids density (M/L³) ρ_s Solids density (M/L³) ρ_s Solids density (M/L³) σ Standard deviation σ_c Unconfined yield strength (M/LT²) σ_t Strength of agglomerates (M/LT²) σ_t Strength of agglomerates (M/LT²) σ_t Mayor consolidation stress (M/LT²) σ₁ Mayor consolidation stress (M/LT²) Σ Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge τ Shear stress (M/LT²), relaxation time in Equation 10.116 (T) τ_f Normal stress at vibration conditions (M/LT²) τ_m Shear stress at wall (M/LT²) τ_m Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12) ψ Wall friction angle φ Phase angle of the nth harmonic defined by Equation 1.6 Φ Sphericity ψ Volume fraction of spheres in suspension ω Angular velocity (1/T) 	μ_{a}	Air viscosity (M/LT)
 μ' Sliding friction coefficient in silo, friction coefficient in roller mills π Osmotic pressure (M/LT²) ρ Liquid density (M/L³) ρ_a Air density (M/L³) ρ_b Bulk density (M/L³) ρ_f Fluid density (M/L³) ρ_g Gas density (M/L³) ρ_m Slurry density (M/L³) ρ_p Particle density (M/L³) ρ_s Solids density (M/L³) ρ_s Solids density (M/L³) ρ(σ) Bulk density under applied normal stress (M/L³) σ Standard deviation σ_c Unconfined yield strength (M/LT²) σ_t Strength of agglomerates (M/LT²) σ_t Strength of agglomerates (M/LT²) σ_t Shear stress at wall surface (M/LT²) σ̄₁ Shear stress acting on arch at 45° (M/LT²) Σ Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge τ Shear stress (M/LT²), relaxation time in Equation 10.116 (T) τ_f Normal stress at vibration conditions (M/LT²) τ_m Shear stress at wall (M/LT²) τ_m Shear stress at wall (M/LT²) τ_m Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12) φ Wall friction angle φ Phase angle of the nth harmonic defined by Equation 1.6 Φ Sphericity Ø Volume fraction of spheres in suspension ω Angular velocity (1/T) 	$\mu_{\rm g}$	Gas viscosity (M/LT)
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	μ_{o}	Viscosity of pure solvent (M/LT)
$\begin{array}{lll} \rho_{a} & \text{Liquid density } (M/L^{3}) \\ \rho_{b} & \text{Bulk density } (M/L^{3}) \\ \rho_{f} & \text{Fluid density } (M/L^{3}) \\ \rho_{g} & \text{Gas density } (M/L^{3}) \\ \rho_{m} & \text{Slurry density } (M/L^{3}) \\ \rho_{p} & \text{Particle density } (M/L^{3}) \\ \rho_{p} & \text{Particle density } (M/L^{3}) \\ \rho_{p} & \text{Solids density } (M/L^{3}) \\ \rho_{s} & \text{Solids density } (M/L^{3}) \\ \sigma & \text{Standard deviation} \\ \sigma_{c} & \text{Unconfined yield strength } (M/LT^{2}) \\ \sigma_{t} & \text{Strength of agglomerates } (M/LT^{2}) \\ \sigma_{t} & \text{Normal stress at wall surface } (M/LT^{2}) \\ \sigma_{1} & \text{Mayor consolidation stress } (M/LT^{2}) \\ \hline \sigma_{1} & \text{Shear stress acting on arch at } 45^{\circ} (M/LT^{2}) \\ \hline \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \tau & \text{Shear stress } (M/LT^{2}), \text{ relaxation time in Equation } 10.116 \ (T) \\ \hline \tau_{t} & \text{Normal stress at vibration conditions } (M/LT^{2}) \\ \hline \tau_{w} & \text{Shear stress at wall } (M/LT^{2}) \\ \hline \tau_{w} & \text{Shear stress at wall } (M/LT^{2}) \\ \hline \phi & \text{Wall friction angle} \\ \phi & \text{Phase angle of the } n\text{th harmonic defined by Equation } 1.6 \\ \hline \Phi & \text{Sphericity} \\ \emptyset & \text{Volume fraction of spheres in suspension} \\ \phi & \text{Angular velocity } (1/T) \\ \hline \end{array} $	μ′	Sliding friction coefficient in silo, friction coefficient in roller mills
ρa Air density (M/L³) ρb Bulk density (M/L³) ρf Fluid density (M/L³) ρg Gas density (M/L³) ρm Slurry density (M/L³) ρp Particle density (M/L³) ρs Solids density (M/L³) ρs Solids density (M/L³) ρ (σ) Bulk density under applied normal stress (M/L³) σ Standard deviation σ _c Unconfined yield strength (M/LT²) σ _t Strength of agglomerates (M/LT²) σ _t Strength of agglomerates (M/LT²) σ _t Mayor consolidation stress (M/LT²) σ _t Mayor consolidation stress (M/LT²) σ _t Shear stress acting on arch at 45° (M/LT²) Σ Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge τ Shear stress (M/LT²), relaxation time in Equation 10.116 (T) τ _f Normal stress at vibration conditions (M/LT²) τ _m Shear stress at wall (M/LT²) τ _m Shear stress at wall (M/LT²) τ _m Shea	π	Osmotic pressure (M/LT^2)
$\begin{array}{lll} \rho_b & \text{Bulk density } (M/L^3) \\ \rho_f & \text{Fluid density } (M/L^3) \\ \rho_g & \text{Gas density } (M/L^3) \\ \rho_m & \text{Slurry density } (M/L^3) \\ \rho_p & \text{Particle density } (M/L^3) \\ \rho_s & \text{Solids density } (M/L^3) \\ \rho_s & \text{Solids density } (M/L^3) \\ \rho_{00} & \text{Bulk density under applied normal stress } (M/L^3) \\ \sigma & \text{Standard deviation} \\ \sigma_c & \text{Unconfined yield strength } (M/LT^2) \\ \sigma_t & \text{Strength of agglomerates } (M/LT^2) \\ \sigma_t & \text{Strength of agglomerates } (M/LT^2) \\ \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \hline \sigma_1 & \text{Shear stress acting on arch at } 45^{\circ} (M/LT^2) \\ \hline \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \tau & \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } 10.116 (T) \\ \hline \tau_f & \text{Normal stress at vibration conditions } (M/LT^2) \\ \hline \tau_w & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_w & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_w & \text{Shear stress at wall } (M/LT^2) \\ \hline \phi & \text{Wall friction angle} \\ \phi & \text{Phase angle of the } n\text{th harmonic defined by Equation } 1.6 \\ \hline \phi & \text{Sphericity} \\ \hline \mathcal{O} & \text{Volume fraction of spheres in suspension} \\ \hline \phi & \text{Angular velocity } (1/T) \\ \hline \end{array} $	ρ	Liquid density (M/L^3)
$\begin{array}{lll} \rho_f & & & & & & & \\ \rho_g & & & & & & \\ Gas & & & & & \\ Gas & & & & & \\ Cas & & & & \\ Cas & & & & \\ Cas & & \\ C$	ρ_{a}	Air density (M/L^3)
$\begin{array}{lll} \rho_g & \text{Gas density } (M/L^3) \\ \rho_m & \text{Slurry density } (M/L^3) \\ \rho_p & \text{Particle density } (M/L^3) \\ \rho_s & \text{Solids density } (M/L^3) \\ \rho(\sigma) & \text{Bulk density under applied normal stress } (M/L^3) \\ \sigma & \text{Standard deviation} \\ \sigma_c & \text{Unconfined yield strength } (M/LT^2) \\ \sigma_t & \text{Strength of agglomerates } (M/LT^2) \\ \sigma_w & \text{Normal stress at wall surface } (M/LT^2) \\ \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \hline \sigma_1 & \text{Shear stress acting on arch at } 45^\circ (M/LT^2) \\ \hline \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \\ \hline \tau & \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } 10.116 (T) \\ \hline \tau_f & \text{Normal stress at vibration conditions } (M/LT^2) \\ \hline \tau_w & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_w & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_{\infty} & \text{Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12)} \\ \phi & \text{Wall friction angle} \\ \phi & \text{Phase angle of the } n\text{th harmonic defined by Equation } 1.6 \\ \hline \Phi & \text{Sphericity} \\ \mathcal{O} & \text{Volume fraction of spheres in suspension} \\ \phi & \text{Angular velocity } (1/T) \\ \hline \end{array}$	ρ_{b}	Bulk density (M/L^3)
$\begin{array}{lll} \rho_m & \text{Slurry density } (M/L^3) \\ \rho_p & \text{Particle density } (M/L^3) \\ \rho_s & \text{Solids density } (M/L^3) \\ \rho(\sigma) & \text{Bulk density under applied normal stress } (M/L^3) \\ \sigma & \text{Standard deviation} \\ \sigma_c & \text{Unconfined yield strength } (M/LT^2) \\ \sigma_t & \text{Strength of agglomerates } (M/LT^2) \\ \sigma_w & \text{Normal stress at wall surface } (M/LT^2) \\ \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \hline \sigma_1 & \text{Shear stress acting on arch at } 45^\circ (M/LT^2) \\ \hline \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \\ \hline \tau & \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } 10.116 (T) \\ \hline \tau_f & \text{Normal stress at vibration conditions } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at operator of a given bulk solid } (\text{Equation } 3.12) \\ \phi & \text{Wall friction angle} \\ \phi & \text{Phase angle of the } n\text{th harmonic defined by Equation } 1.6 \\ \hline \phi & \text{Sphericity} \\ \mathcal{O} & \text{Volume fraction of spheres in suspension} \\ \phi & \text{Angular velocity } (1/T) \\ \hline \end{array}$	$\rho_{\rm f}$	Fluid density (M/L^3)
$\begin{array}{lll} \rho_m & \text{Slurry density } (M/L^3) \\ \rho_p & \text{Particle density } (M/L^3) \\ \rho_s & \text{Solids density } (M/L^3) \\ \rho(\sigma) & \text{Bulk density under applied normal stress } (M/L^3) \\ \sigma & \text{Standard deviation} \\ \sigma_c & \text{Unconfined yield strength } (M/LT^2) \\ \sigma_t & \text{Strength of agglomerates } (M/LT^2) \\ \sigma_w & \text{Normal stress at wall surface } (M/LT^2) \\ \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \hline \sigma_1 & \text{Shear stress acting on arch at } 45^\circ (M/LT^2) \\ \hline \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \tau & \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } 10.116 (T) \\ \hline \tau_f & \text{Normal stress at vibration conditions } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress } (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress } ($	ρ_{g}	Gas density (M/L^3)
 ρ_s Solids density (M/L³) ρ(σ) Bulk density under applied normal stress (M/L³) σ Standard deviation σ_c Unconfined yield strength (M/LT²) σ_t Strength of agglomerates (M/LT²) σ_w Normal stress at wall surface (M/LT²) σ₁ Mayor consolidation stress (M/LT²) σ̄₁ Shear stress acting on arch at 45° (M/LT²) Σ Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge τ Shear stress (M/LT²), relaxation time in Equation 10.116 (T) τ_f Normal stress at vibration conditions (M/LT²) τ_m Shear stress at wall (M/LT²) τ_m Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12) φ Wall friction angle φ Phase angle of the nth harmonic defined by Equation 1.6 Φ Sphericity Ø Volume fraction of spheres in suspension ω Angular velocity (1/T) 	-	Slurry density (M/L^3)
$\begin{array}{lll} \rho(\sigma) & \text{Bulk density under applied normal stress } (M/L^3) \\ \sigma & \text{Standard deviation} \\ \sigma_c & \text{Unconfined yield strength } (M/LT^2) \\ \sigma_t & \text{Strength of agglomerates } (M/LT^2) \\ \sigma_w & \text{Normal stress at wall surface } (M/LT^2) \\ \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \hline \sigma_1 & \text{Shear stress acting on arch at } 45^\circ (M/LT^2) \\ \hline \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge & \text{Toperator} & \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } & 10.116 \ (T) \\ \hline \tau_f & \text{Normal stress at vibration conditions } & (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at wall } & (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at wall } & (M/LT^2) \\ \hline \tau_{w} & \text{Shear stress at wall } & (M/LT^2) \\ \hline \end{array} \\ & \text{Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12)} \\ \phi & \text{Wall friction angle} \\ \phi & \text{Phase angle of the } & \text{nth harmonic defined by Equation } & 1.6 \\ \hline \Phi & \text{Sphericity} \\ \mathcal{O} & \text{Volume fraction of spheres in suspension} \\ \phi & \text{Angular velocity } & (1/T) \\ \hline \end{array}$	$\rho_{\rm p}$	Particle density (M/L^3)
	$ ho_{ m s}$	Solids density (M/L^3)
$\begin{array}{lll} \sigma_c & & & & & \\ \sigma_t & & & & \\ Strength of agglomerates (M/LT^2) \\ \sigma_w & & & & \\ Normal stress at wall surface (M/LT^2) \\ \sigma_1 & & & \\ Mayor consolidation stress (M/LT^2) \\ \overline{\sigma}_1 & & & \\ Shear stress acting on arch at 45^\circ (M/LT^2) \\ \hline{\Sigma} & & & \\ Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \\ \overline{\tau} & & \\ Shear stress (M/LT^2), relaxation time in Equation 10.116 (T) \\ \overline{\tau}_f & & \\ Normal stress at vibration conditions (M/LT^2) \\ \overline{\tau}_{w} & & \\ Shear stress at wall (M/LT^2) \\ \overline{\tau}_{w} & & \\ Shear stress at wall (M/LT^2) \\ \overline{\tau}_{v} & & \\ Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12) \\ \phi & & \\ Wall friction angle \\ \phi & & \\ Phase angle of the nth harmonic defined by Equation 1.6 \\ \Phi & \\ Sphericity \\ \mathcal{O} & & \\ Volume fraction of spheres in suspension \\ \phi & & \\ Angular velocity (1/T) \\ \end{array}$	$\rho(\sigma)$	Bulk density under applied normal stress (M/L^3)
$\begin{array}{lll} \sigma_t & Strength of agglomerates (\textit{M/LT}^2) \\ \sigma_w & Normal stress at wall surface (\textit{M/LT}^2) \\ \sigma_1 & Mayor consolidation stress (\textit{M/LT}^2) \\ \hline \sigma_1 & Shear stress acting on arch at 45^\circ (\textit{M/LT}^2) \Sigma & Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge $	σ	Standard deviation
$\begin{array}{lll} \sigma_w & \text{Normal stress at wall surface } (M/LT^2) \\ \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \hline \sigma_1 & \text{Shear stress acting on arch at } 45^\circ (M/LT^2) \\ \hline \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \tau & \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } 10.116 (T) \\ \hline \tau_f & \text{Normal stress at vibration conditions } (M/LT^2) \\ \hline \tau_{t0} & \text{Normal stress at no vibration conditions } (M/LT^2) \\ \hline \tau_w & \text{Shear stress at wall } (M/LT^2) \\ \hline \tau_{\infty} & \text{Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12)} \\ \phi & \text{Wall friction angle} \\ \phi & \text{Phase angle of the } n \text{th harmonic defined by Equation } 1.6 \\ \hline \Phi & \text{Sphericity} \\ \mathcal{O} & \text{Volume fraction of spheres in suspension} \\ \phi & \text{Angular velocity } (1/T) \\ \hline \end{array} $	$\sigma_{\rm c}$	Unconfined yield strength (M/LT^2)
$\begin{array}{lll} \sigma_1 & \text{Mayor consolidation stress } (M/LT^2) \\ \overline{\sigma}_1 & \text{Shear stress acting on arch at } 45^\circ (M/LT^2) \\ \Sigma & \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \tau & \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } 10.116 (T) \\ \tau_f & \text{Normal stress at vibration conditions } (M/LT^2) \\ \tau_{vo} & \text{Normal stress at no vibration conditions } (M/LT^2) \\ \tau_{vo} & \text{Shear stress at wall } (M/LT^2) \\ \tau_{vo} & \text{Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12)} \\ \phi & \text{Wall friction angle} \\ \phi & \text{Phase angle of the } n\text{th harmonic defined by Equation } 1.6 \\ \Phi & \text{Sphericity} \\ \mathcal{O} & \text{Volume fraction of spheres in suspension} \\ \omega & \text{Angular velocity } (1/T) \\ \end{array}$	$\sigma_{\scriptscriptstyle t}$	Strength of agglomerates (M/LT^2)
$ \overline{\sigma}_1 \qquad \text{Shear stress acting on arch at } 45^\circ (M/LT^2) $ $ \Sigma \qquad \text{Operator, meaning "algebraic sum of," characteristic geometrical features of a centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge \tau \qquad \text{Shear stress } (M/LT^2), \text{ relaxation time in Equation } 10.116 (T) \tau_f \qquad \text{Normal stress at vibration conditions } (M/LT^2) \tau_w \qquad \text{Shear stress at mo vibration conditions } (M/LT^2) \tau_w \qquad \text{Shear stress at wall } (M/LT^2) \tau_w \qquad \text{Shear stress at wall } (M/LT^2) \tau_w \qquad \text{Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12) } \phi \qquad \text{Wall friction angle } \phi \qquad \text{Phase angle of the } n \text{th harmonic defined by Equation } 1.6 \Phi \qquad \text{Sphericity } \emptyset \qquad \text{Volume fraction of spheres in suspension } \omega \qquad \text{Angular velocity } (1/T) $	$\sigma_{\rm w}$	Normal stress at wall surface (M/LT^2)
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	σ_1	Mayor consolidation stress (M/LT^2)
centrifuge equivalent to area of a gravity settling tank with similar settling characteristics of centrifuge τ Shear stress (M/LT^2) , relaxation time in Equation 10.116 (T) τ_f Normal stress at vibration conditions (M/LT^2) τ_{t0} Normal stress at no vibration conditions (M/LT^2) τ_{w} Shear stress at wall (M/LT^2) τ_{w} Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12) ϕ Wall friction angle ϕ Phase angle of the n th harmonic defined by Equation 1.6 Φ Sphericity \emptyset Volume fraction of spheres in suspension Φ Angular velocity $(1/T)$	$\overline{\sigma}_1$	Shear stress acting on arch at 45° (M/LT^2)
$\begin{array}{lll} \tau_{\rm f} & {\rm Normal~stress~at~vibration~conditions~(\it M/LT^2)} \\ \tau_{\rm f0} & {\rm Normal~stress~at~no~vibration~conditions~(\it M/LT^2)} \\ \tau_{\rm w} & {\rm Shear~stress~at~wall~(\it M/LT^2)} \\ \tau_{\rm \infty} & {\rm Constant~depending~on~the~consolidation~and~applied~normal~stress~pressure~during~shear,~of~a~given~bulk~solid~(Equation~3.12)} \\ \phi & {\rm Wall~friction~angle} \\ \phi & {\rm Phase~angle~of~the~}\it{n} th~harmonic~defined~by~Equation~1.6} \\ \Phi & {\rm Sphericity} \\ {\cal O} & {\rm Volume~fraction~of~spheres~in~suspension} \\ \omega & {\rm Angular~velocity~(1/\it{T})} \\ \end{array}$	Σ	centrifuge equivalent to area of a gravity settling tank with similar settling
$\begin{array}{lll} \tau_{\text{f0}} & & \text{Normal stress at no vibration conditions } (\textit{M/LT}^2) \\ \tau_{\text{w}} & & \text{Shear stress at wall } (\textit{M/LT}^2) \\ \tau_{\text{o}} & & \text{Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12)} \\ \phi & & \text{Wall friction angle} \\ \phi & & \text{Phase angle of the } \textit{n} \text{th harmonic defined by Equation 1.6} \\ \Phi & & \text{Sphericity} \\ \varnothing & & \text{Volume fraction of spheres in suspension} \\ \omega & & \text{Angular velocity } (1/T) \\ \end{array}$	τ	Shear stress (M/LT^2), relaxation time in Equation 10.116 (T)
$\begin{array}{lll} \tau_{w} & & \text{Shear stress at wall } (\textit{M/LT}^{2}) \\ \tau_{\omega} & & \text{Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12)} \\ \phi & & \text{Wall friction angle} \\ \phi & & \text{Phase angle of the } \textit{n} \text{th harmonic defined by Equation 1.6} \\ \Phi & & \text{Sphericity} \\ \emptyset & & \text{Volume fraction of spheres in suspension} \\ \omega & & \text{Angular velocity } (1/T) \\ \end{array}$	τ_{f}	Normal stress at vibration conditions (M/LT^2)
 Constant depending on the consolidation and applied normal stress pressure during shear, of a given bulk solid (Equation 3.12) ψ Wall friction angle φ Phase angle of the nth harmonic defined by Equation 1.6 Φ Sphericity Ø Volume fraction of spheres in suspension ω Angular velocity (1/T) 	$\tau_{\rm f0}$	Normal stress at no vibration conditions (M/LT ²)
during shear, of a given bulk solid (Equation 3.12) ϕ Wall friction angle ϕ Phase angle of the n th harmonic defined by Equation 1.6 Φ Sphericity \mathcal{O} Volume fraction of spheres in suspension Φ Angular velocity (1/ T)	$\tau_{\rm w}$	Shear stress at wall (M/LT^2)
φ Phase angle of the n th harmonic defined by Equation 1.6 $Φ$ Sphericity $Ø$ Volume fraction of spheres in suspension $Φ$ Angular velocity $Φ$	$ au_{\scriptscriptstyle\infty}$	
Φ Sphericity $Ø$ Volume fraction of spheres in suspension $Φ$ Angular velocity $Φ$ $Φ$ Angular velocity $Φ$	ф	Wall friction angle
\emptyset Volume fraction of spheres in suspension $ω$ Angular velocity $(1/T)$	φ	Phase angle of the <i>n</i> th harmonic defined by Equation 1.6
ω Angular velocity (1/ T)	Φ	Sphericity
	Ø	Volume fraction of spheres in suspension
Ω Conical half-angle of discs in disc-bowl centrifuge	ω	Angular velocity $(1/T)$
	Ω	Conical half-angle of discs in disc-bowl centrifuge

a Volt.

^b Coulomb.

A	Aggregative fluidization, 252. See also
Abrasian transfer 213	Bubbling—fluidization
Abrasion transfer, 213	Agitation mills, 192, 194, 203
Abrasiva wear 121	Air blasters. See Air cannons
Absolute vigosity, 112	Air cannons, 126
Absolute viscosity, 113	Air classification, 334, 340–341, 344
Acceptance efficiency, 328. See also	Air classifiers, 334, 340
Screening—efficiency	applications, 344
Acetate fiber, 403	calibration curve for, 344
Acrylic fiber, 403	categories, 341, 342
Active devices, 123	operation, 343–344
air cannons, 126	Alkali Act, 4
decay in shear strength, 125	Anadynamic magnetic field, 302
dynamic wall shear tests, 125–126	Andreasen Pipette-pipette centrifuge
effective flow promotion, 126	method, 85
failure plane formation, 126	Andreasen's sedimentation pipette, 83
yield loci and flow functions, 125	Angle of nip, 195, 196
Active stress field. See Peaked stress	Angle of repose, 49, 114. See Repose
field	angle
Aerated bulk density, 24, 25	drained, 62
Aeration, 223	poured, 61
Agglomerate. See also Green—products	Annular shear cell. <i>See</i> Ring shear cell
applied pressure calculation, 211	Antidynamic discharge tube, 112
stability, 210	principle, 113
strength of, 210, 211, 223–224	variation, 112
tensile strength of, 209	Apron feeder, 127, 129, 148
volume calculation, 211–212	Archimedes
Agglomeration, 28, 207	number, 37
applications, 224	principle, 320
binding mechanisms, 208	Arching phenomena, 113
compaction, 214	Assisted discharge
coordination number, 207	active devices, 123, 124–127
design aspects, 222–224	BINSERT® system, 124
equipment operation variables, 216	
method selection guidelines, 219	passive devices, 123
moisture, 220	types, 123
particle size, 214, 219, 226	Attraction forces, 212
product properties, 221	Attrition, 180, 275–276
raw materials characteristics, 219–220	n.
selection criteria for, 219–222	В
technique options, 221–222	Bag filter, 345, 346, 347, 348
tumbling, 212	Ball mill, 192, 203
Aggregation, 207. See also	critical speed, 200
Agglomeration	forces 198

Bar screens. See Grizzlies	phase, 262
Basket	in solid–gas systems, 262
centrifuge, 390	Bubbling fluidized bed
extruder, 216	bubbling phase, 262
Belt, 137	particulate phase, 262, 263
conveyor, 136, 138, 142	Bucket elevators, 148, 149
separator, 303	buckets, 149
thickness, 142	horsepower, 150–151
Belt conveyors, 136	spacing of buckets, 19–150
characteristics of solid, 142–143	Bulk, 24
components, 136, 137	density, 23, 24, 250
idler, 138–139	Bulk solid, 31
power for, 140–142, 143	failure properties, 48, 65
pulley transmission relations, 142	
	flow types, 104 handling plants 104, 131, 132
tension devices, 138, 139	handling plants, 104, 131, 132
thickness of, 142	particle size analysis, 67, 89
tripper, 139–140	powder classification, 32
Belt feeder. See Apron feeder Rinder 212, 220, 225, See also Water	sampling, 37
Binder, 212, 220, 225. See also Water	Bulk solid, storage, 101
Bingham plastic fluids, 318, 319	confined storage, 102
Bins, 102. See also Hopper; Silos	outdoors storage, 101
design considerations, 103	storage alternatives, 101
funnel flow, 104–107	wall loads in silos evaluation, 134
mass-flow, 104	Bulk solid conveying, 135
stress fields, 108–109	belt conveyors, 136–143
wall load prediction, 108	capacity calculation, 155
Binsert® system, 123, 124. See also	chain conveyors, 143–151
Passive devices	conveyer selection, 153–155
Blake crushers, 185, 186	pneumatic conveying, 155, 171
Blending	screw conveyors, 151–155
indices determination, 244–245	Buoyancy force, 320
mechanism, 230–231	
quality, 229, 233	C
uniformity, 242	
Bond Work Index, 184	Cake, 403, 408
Bond's law, 184	discharge, 405, 406
Breakage stages, 182	formation, 365
Bridging phenomena. See Arching	resistance factor, 348
phenomena	Capillary force aggregation, 208
Briquetting roller press, 217	Carman–Kozeny equation, 22, 254, 258
British standard sieves, 438–439	Cartridge filters, 403
Bubble, 249, 252	Central Limit Theorem, 40
air, 20	Centrifugal clarification, 379, 383
behavior, 262	capacity determination, 425
in gas-fluidized bed, 261	cut size, 421–422
powder effect, 34	disc-bowl centrifuges, 388, 389, 390
in solid–gas systems, 262	radial separation zones, 380
Bubbling	radial velocity, 379
fluidization, 261	Tubular centrifuges, 388, 389

Coarse, 288
bulk solids, 116
particles, 9, 10
reduction, 180, 182
Coating, 272
attrition, 275–276
efficiency evaluation, 273. See also
Attrition
equipments for, 272-273
thickness estimation, 275
Coefficient of friction, 8, 146
force, 196
between particle and screen,
287–288
screw conveyor power
requirement, 153
Cohesion
number, 37
powder, 50
Comminution, 75, 179, 309. See also Size
reduction
Bond's law, 184
energy requirements in, 182–183
feed pretreatment, 201–202
Kick's law, 183–184
moisture, 202
paramount factors, 201
reduction mechanism, 180
Rittniger's law, 183
Rosin–Rammler function, 74
screen aperture size, 182
work index, 184
Compacting roller press, 217
Compaction. See Pressure
agglomeration
Compartment mill, 193
Compressibility, 26–27
Compressive force, 180, 188
Concentrate, 413
magnetic, 302
test fluids, 360
Conductive induction, 307–308
by direct current, 309
Confidence interval, 40–41
length, 42
<i>U</i> and <i>L</i> , 44
Confined storage, 102
assisted discharge, 123
design factors, 130

Confined storage (Continued) feeders, 127 general considerations, 103 natural discharge, 112 silos types, 104 wall stresses, 108–112 Conical mill, 193 Coning and quartering, 46 Construction industry, 3 Contact electrification, 307, 308	Cyclone, 334, 335. See also Dilute-phase conveying; Hydrocyclone applications, 340 cut size, 337 diameter determination, 422–424 operating variables, 337–340 process description, 334–336 separation, 285 Stairmand design, 337–338 Cylindrical-die extruder, 216
Continuous mixing, 242	
ribbon-type mixer, 242	D
screw conveyors, 151	
Convective mixing, 230	Da. See Daltons (Da)
Conventional magnetic separation, 303	Daltons (Da), 410
Conversion factors, 429	Davis, George E., 4
Conveying angle, 62	Dead flux. See Partition effect
Conveyor	Dead-end flow filtration, 412
belt, 127, 136, 143	Decantation, 285, 367
features, 137	Decanter centrifuge, 390
selection guide, 138	Deep-cone thickener, 372, 373
Conveyor-bowl centrifuge, 392	Degree of separation, 327
Coordination number, 207	Dense-phase conveying, 157
of monosized particles, 208	blow tank system, 159
Corona discharge electrostatic	fluidized systems, 158–159
separator, 310	plug-phase conveying, 157–158
Cotton fiber, 403	Density, 7, 18, 429
Critical diameter. See Cut—point	aerated, 24
Critical speed, 197	aerated bulk, 24, 25
in revolution, 200	air, 433
trommel, 299	bulk, 24
Cross-flow filtration, 412, 413	compacted bulk, 26
Crushers, 184, 185–187	particle, 18
Crushing and layering, 213	poured, 25
Crushing rolls, 195	pycnometry, 19, 21–23
forces and angle of nip in, 196	solid, 24
Crystallization	tap bulk, 26, 28
agglomerate binding, 207	top-loading platform scale, 20
solutions, 285	water, 431
Cubic particle, 208, 209	Denver crushers, 185, 186
Cut	Diamagnetic materials, 301
diameter, 288	Diameter, 10
force, 180	equivalent circle, 11
point, 343, 379	equivalent sphere, 10
size, 331, 342, 337–388	mean, 71
Cutting	of nonspherical particles, 12
forces, 180–181	projected and equivalent, 15
machines 194	statistical 11

Ct 1 / 10	1 : 200
Stokes', 12	revolving screen, 299
surface-to-volume, 12	separator, 303
Diatomaceous earth, 403	Drying, 271, 285
Dieresis and electro dialysis, 285	in fluidized beds, 269–272
Diffusion, 349	periods, 272
isothermal, 272	Dust
mixing, 230	collection, 347
pure, 230	emission, 333
separation techniques, 285	explosion, 275. See also Attrition
Dilute phase closed electrostatic	formation, 242
circle, 312	settling chamber, 351
Dilute-phase conveying, 157, 159.	Dwell time, 215, 379
See also Cyclone	
air-solid two-phase flow, 159–160	E
blower, 160	L
closed-loop system, 162, 163	Economics, modern, 3
combined system, 162, 163	Electric charges, developed, 308
feeders, 160	Electromagnetic separation, 285, 301
operating variables of, 161–162	applications of, 304
pressure system, 162, 163	diamagnetic materials, 301
vacuum system, 162, 163	ferromagnetic materials, 301–302
Dilute-phase conveying system,	high-gradient magnetic separation
161–162	303, 307
Dimensional analysis, 321, 435	magnetic behavior, 301
power equation, 436	magnetic field types, 302
relationships between dimensionless	magnetic force vs. ferromagnetic
groups, 386, 421	wire diameter, 306
Disc agglomerator operation. See Pan	paramagnetic materials, 302
agglomerator operation	selection criteria, 306
Disc-bowl centrifuge, 381, 390	separators, 303
for clarification, 388	superconductors, 305
solids capacities, 389	Electrostatic, 307
Disperse diphasic system, 32	plate separators, 309, 311
Dispersed-phase conveying. See	screen–plate separators, 311
Dilute-phase conveying	separators, 352
Dispersibility, 28–29	Electrostatic separation, 285, 307
Distillation, 268, 285	applications, 309, 312
Dodge crushers, 185, 186	comminution, 309
Double cone, Tumbler mixers, 240	conductive induction, 307–308
Drag coefficient, 321	contact electrification, 307, 308
in hindered settling, 368	corona discharge electrostatic
vs. particle Reynolds number, 322	separator, 310
Drag force, 321	dilute phase closed electrostatic
Drum	circle, 312
agglomerator operation, 225	ion bombardment, 307, 308
concentric, 299, 300	polarization, 309
continuous rotary, 407	Empty-tower velocity. See Superficial
inclined rotating, 214	velocity
magnetic separator, 304	Emulsion phase, 262

Encapsulation. See Coating	Filtration, 285, 394–395, 408
Energy, 429	constant pressure, 396–399
efficiency, 179	constant rate, 400–402
requirements in comminution,	cross-flow, 412
182–183	curves using filter press, 408–409
sources, 3	dead-end flow, 412
vibration, 126	equipment, 403–408
Envelope filter, 345, 347	feed slurry, 395
Equivalent spheres, 13	resistance, 425
diameter, 10	theory, 395–396
Erosive wear, 131	Fine material, 120, 288. See also
Error function, 75	Powder
Evaporation, 285	Flat-die extruder, 216
	Flocculant. See Flocculation—agents
F	Flocculation, 361, 378
r	agents, 361, 362
Failure function (FF), 50, 118	clarifier performance, 375
critical stress, 122	second-order test procedure, 376
device scheme, 59	uses, 377
direct measurement, 57	Flocculators, 361
powders, 34, 57–58	Flocs, 361
Falling rate drying period, 272	Flotation, 285
Feed slurry, 395	Flow
Feeders, 127. See also Bulk solid—	angle of repose, 114
handling plants; Bulk solid,	coefficient of kinetic friction, 114
storage	granular material, 115
charge and power calculations, 128	partition effect, 326
discharge control in, 127, 130	promotion, 126, 123
flowability, 132	regions, 115
gates, 132, 133	theories, 113–116
level gauges, 133	Flow behavior curve. <i>See</i> Rheogram
load on, 130	Flow factor (ff), 33, 118
selection guide, 129	angle of internal friction, 119
solid buildup, 131	for conical hopper, 119
wall friction angle, 131	hopper half angle, 122
Feret's diameter, 11, 91	Flowability, 132
Ferrofilter Franz, 303	Fluid
Ferromagnetic materials, 301–302	classification of, 317–320
FF. See Failure function (FF)	
ff. See Flow factor (ff)	displacement methods, 18
	energy (jet) mills, 194 flow through solid beds, 250–251
Field scanning, 87 Filter	e
	mixing, 230
aids, 403	particle dynamics in, 320–324
cake, 395	Rheograms for, 319
filtration curves, 408–409	and solid flow, 317
medium, 395, 402–403	Fluid and solid separation, 324
presses, 403, 404, 405, 408–409	acceptance efficiency, 328
properties of fiber, 403	clarification number, 327
Filtrate, 395	cut size, 329, 331

degree of separation, 327	Reynolds number, 258, 259
drag coefficient vs. Reynolds	Fluidized regimes, 253
number, 322	bubbling fluidization, 261
evaluation, 324–325	Carman-Kozeny equation, 254
grade efficiency curve, 329, 330,	incipient fluidization and minimum
332–333	fluidizing velocity, 256
hygienic efficiency, 328	pressure drop-velocity relationship,
net efficiency, 326, 327	254–256
partial efficiency, 328	spouted beds, 264–267
partial gravimetric efficiency,	Fluidized-bed coating, 273
327–329	advantage, 275
partition effect, 331	bottom-spray, 274
sharpness index, 331	calculations and report, 277
total gravimetric efficiency, 325–327	equipments, 276
underflows to-throughput ratio, 326	food particulates, 276–277
Fluidization, 250	friability test, 277
applications, 267–269	tangential-spray, rotor, 275
bulk density, 250	thickness estimation, 275, 277
fluid flow through solid beds, 250–251	top-spray, 274
mass balance, 250	Fluidized-bed drying, 269
mechanism of, 251–253	design equation, 271, 272
porosity, 250	limiting kinetic factor, 272
spiralling helix, 273	multichamber, 270–271
void fraction calculation, 278	single-chamber, 270
Fluidization, point of incipient, 255	types, 269
Fluidized bed, 249	Flux
agglomeration equipment, 214	of air, 167
air velocity requirement, 278	dead, 331
bubble behavior, 262	design curves for thickeners 374
coating, 272	of solids, 165
combustion, 268–269	total, 373
determinants, 257	Force, 429
drying in, 269–272	centrifugal, 273
expansion, 263	compression, 34
fluid flow rate, 251–252	friction, 113–114
Galileo number, 258, 259	shear, 51, 53, 54
heat and mass transfer, 254	twisting top, 51
height calculation, 278	Fractal dimension, 17
nonfluidized regions in, 256	Fragility, 8
operation of, 276	Free-settling time determination, 424
packed, 263	Friability test, 276
in petroleum and chemical	fluidized-bed coating, 277
industries, 268	tumbling agglomeration, 226
porosity of, 250, 251	Friction, 8, 113
preferential flow, 255	coefficient, 8, 147
pressure drop, 254, 257, 258,	force, 113–114
260–261, 278	influence, 130
pressure gradient in, 253	Froude number, 252, 419, 436
properties, 249	Fuller's earth, 403

Gas absorption, 285 filtration, 285 fluidized systems, 261 separation, 409–410 Gas filters, 345 applications, 348 bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 Grain silos, 221 Grate separator, 303 Gravity, 320 Grant entrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Grab silos grantin, 409–410 Hammer mills, 189, 190. See also Rotary knife cutters reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-siloe gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization. See Bubbling—fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 104 furnel flow, 107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Handsner ation, 224-205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale ator, 348 impact wear, 131 with Mohs' scale, 201 Hadnesner atio, 28 Heterogeneous fluidization. See Rubbling—fluidization High-gradient magnetic separator, 305 Gealise, 32 Hardness, 7 glass, 8 impact wear, 13 with Mohs' scale ator, 348 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28		
absorption, 285 filtration, 409–410 for cultratine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hauser ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization. See Bubbling—fluidization. See Non-bubbling fluidization.	G	Grizzlies, 296
absorption, 285 filtration, 285 fluidized systems, 261 separation, 409–410 Gas filters, 345 applications, 348 bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 operation characteristics, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Gas filters, 345 Hammer mills, 189, 190. See also Rotary knife cutters reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separation, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 304 vibrating, 123 Horizontal	Gas	= = :
filtration, 285 fluidized systems, 261 separation, 409–410 Gas filters, 345 applications, 348 bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 prossure drop, 348 prossure drop, 348 prossure drop, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 82. See also Sedimentation thickener, 371, 372 Hammer mills, 189, 190. See also Rotary knife cutters reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization. See Bubbling—fluidization High-gradient magnetic separation, 303 separator, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 346, 347 operation relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separator, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homog	absorption, 285	
separation, 409–410 Gas filters, 345 applications, 348 bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Hammer mills, 189, 190. See also Rotary knife cutters reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hard-slide gate, 132		Gyratory Crusher, 167
Separation, 409–410 Gas filters, 345 applications, 348 bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 operation characteristics, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Hammer mills, 189, 190. See also Rotary knife cutters reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow factor, 118, 119 flow factor, 128 flow characteristic	fluidized systems, 261	***
applications, 348 bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates-Gaudin-Schuhmann function, 74 Gaudin-Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 knife cutters reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	separation, 409–410	Н
applications, 348 bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates-Gaudin–Schuhmann function, 74 Gaudin-Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 knife cutters reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	-	Hammer mills, 189, 190. See also Rotary
bag filter, 345, 346, 347, 348 blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 reduction relation determination, 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 192	applications, 348	
blow ring, 347 cake resistance factor, 348 envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates-Gaudin-Schuhmann function, 74 Gaudin-Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Day 124 204–205 for ultrafine grinding, 192 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separator, 303 separator, 305 High-pressure agglomeration, 215, 216 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Hatroness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Hatroness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Hatroness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separator, 305 High-gradient magneti		reduction relation determination,
envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates-Gaudin-Schuhmann function, 74 Gaudin-Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separation, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	blow ring, 347	204–205
envelope filter, 345, 347 fabric properties, 346, 347 operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Hand-slide gate, 132 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization. See Non-bubbling fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	cake resistance factor, 348	for ultrafine grinding, 192
fabric properties, 346, 347 operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates-Gaudin-Schuhmann function, 74 Gaudin-Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Hardness, 7 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separation, 303 separator, 305 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	envelope filter, 345, 347	
operation characteristics, 348 pressure drop, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 glass, 8 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separation, 303 separator, 305 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	fabric properties, 346, 347	
pressure drop, 348 process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 impact wear, 131 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization High-gradient magnetic separation, 303 separator, 303 separator, 303 separator, 305 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal		glass, 8
process description, 345 Gate, 132, 133 Gates—Gaudin—Schuhmann function, 74 Gaudin—Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 with Mohs' scale, 201 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separator, 303 separator, 305 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	pressure drop, 348	9
Gate, 132, 133 Gates-Gaudin-Schuhmann function, 74 Gaudin-Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Hausner ratio, 28 Heterogeneous fluidization. See Bubbling—fluidization High-gradient magnetic separation, 303 separator, 303 separator, 303 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization High-gradient magnetic separation, 303 separator, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization High-gradient magnetic separation, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization High-gradient magnetic separation, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization High-gradient magnetic separation, 303 separator, 305 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization Hopper, 102, 130. See also fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal		
function, 74 Gaudin–Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Geldart's classification, 33, 34 High-pressure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow factor, 118, 119 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal		
function, 74 Gaudin–Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Bubbling—fluidization High-gradient magnetic separation, 303 separator, 305 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization. See Non-bubbling fluidization, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization separation, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization. See Non-bubbling fluidization separator, 305 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization separator, 305 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization High-gradient magnetic separation, 303 separator, 305 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	Gates-Gaudin-Schuhmann	Heterogeneous fluidization. See
Gaudin–Meloy functions, modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 High-presaure agglomeration, 215, 216 Hollow fiber membrane, 414, 416 Homogeneous fluidization. See Non-bubbling fluidization Hopper, 102, 130. See also separation, 203 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	function, 74	
modified, 74 Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330,	Gaudin-Meloy functions,	
Geldart's classification, 33, 34 cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330,	modified, 74	
cohesion number, 37 dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330,	Geldart's classification, 33, 34	-
dimensionless representation, 38 limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330,	cohesion number, 37	
limit of behavior, 36–37 Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330,	dimensionless representation, 38	
Grab sampling, 45. See also Sampling Grade efficiency curve, 329, 330,	limit of behavior, 36–37	
Grade efficiency curve, 329, 330, 332–333 clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation thickener, 371, 372 Hopper, 102, 130. See also Bins; Silos coarse bulk solids, opening for, 116–120 design considerations, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	Grab sampling, 45. See also Sampling	
clarifier, 377 cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation Sedimentation tickener, 371, 372 coarse bulk solids, opening for, 103 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	Grade efficiency curve, 329, 330,	
cut size, 331 equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation Sedimentation thickener, 371, 372 Grain silos, 329 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	332–333	coarse bulk solids, opening for,
equations, 329 Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation Sedimentation thickener, 371, 372 expanded-flow, 107 fine bulk solids, opening for, 120–123 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	clarifier, 377	116–120
Grain silos, 101. See also Silos Granular material flow property, 115 products, 221 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation, 82. See also Sedimentation thickener, 371, 372 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	cut size, 331	design considerations, 103
Granular material flow property, 115 products, 221 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation, 82. See also Sedimentation thickener, 371, 372 flow channel and equilibrium of forces, 117 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	equations, 329	expanded-flow, 107
material flow property, 115 products, 221 flow factor, 118, 119 Grate separator, 303 flow in, 103, 105, 115 Gravity, 320 flow regimes, 116 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation, 82. See also Sedimentation Sedimentation Sedimentation Thickener, 371, 372 flow factor, 118, 119 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	Grain silos, 101. See also Silos	fine bulk solids, opening for, 120–123
products, 221 flow factor, 118, 119 Grate separator, 303 flow in, 103, 105, 115 Gravity, 320 flow regimes, 116 and centrifugal forces, 395 funnel flow, 104–107 and centrifugal sedimentation, 85 methods, 285 half-angle, 117, 120 sedimentation, 82. See also types, 104 Sedimentation vibrating, 123 thickener, 371, 372 Horizontal	Granular	flow channel and equilibrium of
Grate separator, 303 Gravity, 320 and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation, 82. See also Sedimentation thickener, 371, 372 flow in, 103, 105, 115 flow regimes, 116 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal		forces, 117
Gravity, 320 flow regimes, 116 and centrifugal forces, 395 funnel flow, 104–107 and centrifugal sedimentation, 85 geometries, 106, 118 methods, 285 half-angle, 117, 120 sedimentation, 82. See also types, 104 Sedimentation vibrating, 123 thickener, 371, 372 Horizontal	products, 221	flow factor, 118, 119
and centrifugal forces, 395 and centrifugal sedimentation, 85 methods, 285 sedimentation, 82. See also Sedimentation thickener, 371, 372 funnel flow, 104–107 geometries, 106, 118 half-angle, 117, 120 types, 104 vibrating, 123 Horizontal	Grate separator, 303	flow in, 103, 105, 115
and centrifugal sedimentation, 85 geometries, 106, 118 methods, 285 half-angle, 117, 120 sedimentation, 82. See also types, 104 Sedimentation vibrating, 123 thickener, 371, 372 Horizontal		flow regimes, 116
methods, 285 half-angle, 117, 120 sedimentation, 82. See also types, 104 Sedimentation vibrating, 123 thickener, 371, 372 Horizontal		funnel flow, 104–107
sedimentation, 82. See also types, 104 Sedimentation vibrating, 123 thickener, 371, 372 Horizontal	and centrifugal sedimentation, 85	geometries, 106, 118
Sedimentation vibrating, 123 thickener, 371, 372 Horizontal		half-angle, 117, 120
thickener, 371, 372 Horizontal		types, 104
		vibrating, 123
Green cylinder tumbler mixer 240	thickener, 371, 372	Horizontal
cylinder value of mixely 210	Green	cylinder tumbler mixer, 240
agglomerates, 223 trough mixers, 241–242	00	
products, 213 Humidification, 285		
Grinders, 185 Hydrocyclone, 383, 393. See also Cyclone		
Grinding, 183 bodies, 391	Grinding, 183	bodies, 391

Bradley's, 388	Isodynamic magnetic field, 302
cut point estimation, 425	Isotropic material, 60
cut size, 391	
deck, 391, 392	J
efficiency, 392–394	
Rietema's, 384–385	Janssen 110 111
separation, 285	equation, 110–111
underflow-to-throughput, 386	theory, 109
Hydrodynamic pressure gradient, 413	Jaw crushers, 185, 186
Hydrometallurgical extraction, 285	Jenike, Andrew W., 103
Hygienic efficiency, 328	Jenike shear cell, 50, 51 alternatives, 66
	disadvantages, 65, 66
T	problem, 54
I	wall friction angle, 53
IChemE. See Institution of Chemical	Jenike's classification, 33–34. See also
Engineers (IChemE)	Geldart's classification
Idler, 138–139	General Composition
Impact force, 180	T/
Imperfect fluids, 113. See also	K
Bulk solid	Katadynamic magnetic field, 302
Incipient fluidization point	Kick's law, 183–184
Galileo number, 258, 259	kieselguhr earth, 403
Reynolds number, 258, 259	Kinetic friction coefficient, 114
Inclined rotating disc, 214	
Inclined rotating drum, 214	L
Inserts. See Passive devices	
Institution of Chemical Engineers	Laboratory report preparation
(IChemE), 6	guidelines, 441
Instrumental	abstract, 443
particle counters, 85–86	appendices, 446
particle sizer, 90	background, 444
techniques, 95	charts and graphs, 447
Interlocking bond aggregation, 208	components, 441
Intermediate product. See Middling	conclusions, 445
stream Intermediate reduction, 180	discussion, 445
	drawings and diagrams, 446–447
Intermeshing-gears extruder, 216 Internal friction effective angle, 50, 53	equations, 447 illustrative aids, 446
conical hopper and, 119	introduction, 444
crushed limestone, 173	language, 442
grooved plate method, 55, 56	materials and methods, 444
hoppers dependency, 115	neatness, 442
Ion bombardment, 307, 308. See also	objectives, 444
Corona discharge electrostatic	recommendations, 445
separator	references, literature cited, 445
by direct current, 309	results and interpretation, 444–445
Ionic exchange, 285	tables, 443, 446
Irregular particle, 68	title page, 443
- *	* C

Lamella thickener, 372, 373	mass balance, 413
Laser diffraction particle counter, 86	MWCO, 410
Law of Large Numbers, 40	operation modes, 416-417, 418
Leaching and extraction, 285	permeate, 413
Leaf filters, 403, 405–406	pressure drop, 413, 417, 418–419
Length, 429	recovery, 414, 415
Level gauges, 133	MF. See Microfiltration (MF)
Lever-operated quadrant gate, 132, 133	Microfiltration (MF), 409
Light diffraction, 85	Microscopy counting, 77
Liquid bridges hardening binder,	calculations and report, 92
208, 210	particle size measurement, 91
Liquid extraction, 285	procedure, 91–92
Little, Arthur D., 5	slide preparation, 91
,	Middling stream, 302, 313
	Mikro-Atomizer, 193
M	Minimum bubbling velocity, 34, 262
Mag stream, 302	Minimum fluidizing velocity, 34, 253,
Magnetic	255, 256, 257
behavior, 301	as free falling velocity, 259
field types, 302	Galileo number, 259
separation, 302	ratio of terminal falling to, 260
separators, 303, 304	Reynolds number, 259
susceptibility, 301	Minimum spouting velocity, 266
Manufacturing industry, 3	Minus material, 288, 313
Martin's diameter, 11, 91	MIT. See Massachusetts Institute of
Mass, 429	Technology (MIT)
balance, 325, 326	Mixing, 229
median size, 73	blending, 230
Massachusetts Institute of	degree of, 235
Technology (MIT), 4	density impact on, 230–231
Materials processing industry, 3–4	equipment, 239
mixtures, 283	indexes, 233, 234, 246
particulate systems, 101	principle, 239
Maximum angle of rise. <i>See</i> Conveying	rate evaluation, 233, 242
angle	sampling, 232
Mean, 70	sand and clay, 235–237
diameters, 71	statistical approach of, 231–232
population, 40	time, 235–239
Mechanical separations, 285	Mixing equipment, 239
Mechanized reclaiming systems, 102	design and selection factors, 241–244
Median, 70	dust minimization, 243
particle size, 73	horizontal trough mixers, 240–241
Membrane, 409	ribbon mixer, 240
Membrane separation, 285, 409	tumbling mixers, 239
arrangements of, 416, 417	vertical screw mixers, 241
concentrate, 413	Mode, 70
cross-flow filtration, 412, 413	Modified beta function, 75
flow rate, 413	Modified Gaudin–Meloy functions, 74
hydrodynamic pressure gradient, 413	Mohs' scale, 201
,,	

Moisture, 202 agglomerate stability, 210 in agglomeration, 219, 220 bulk solid flow, 132 consolidating stress against unconfined yield stress, 173 in drying, 295 in electrostatic separators, 309 in fluidized-bed drying, 272 wall friction angle, 172–173 Molecular force aggregation, 208 Molecular weight cutoff (MWCO), 410 Monosized particles, 208–209 Multicomponent separation, 285	Organic rejection, 412. See also Ultrafiltration (UF) Orthorhombic particle, 208, 209 Oscillating screens, 296, 298 Osmotic equilibrium, 410 flow, 410 pressure, 410, 411. See also Reverse osmosis (RO) Outdoors storage, 101–102 Oversize particle, 288, 313 cumulative frequency of, 289
MWCO. See Molecular weight cutoff	P
(MWCO)	Packed fluidized bed, 263 Pan agglomerator operation, 225 Paramagnetic substances, 301
N. (Partial melting sinter bridge
Natural discharge, 112 flow theories, 113 hopper opening, 116, 120 Net efficiency, 326 of fines, 327 Newtonian fluid, 318 friction factor, 357 Non-bubbling fluidization, 261	aggregation, 208 Particle, 8 density, 18 particle outline shape, 13 perimeter and convex perimeter, 14 properties, 9 shape, 14, 15–16 single-particle characteristics, 9
Non-Newtonian fluid, 318 mixtures, 113 rheological parameters, 359 settling of, 356 suspensions, 354, 387 in turbulent flow, 356 velocity profiles of, 357	Particle counters, 85–86 Particle friction factor, 167 Particle size analysis, 67 accessories, 90 central tendency measures, 69–71 centrifugation, 82 characteristic linear dimension, 68
Nonsuspension conveying. See Dense-phase conveying, 17 Normal distribution function, 74, 75	data presentation, 71 by different methods, 89 distribution types, 68–69
Normal distribution function, 74–75 Notations, 449–456 Nozzle-discharge centrifuge, 390, 391 Nucleation, 213 Number median size, 73 Nylon fiber, 403	frequency and cumulative distributions, 73 instrumental method, 95 laboratory techniques, 76 light diffraction, 85 materials and instruments, 90 measurement, 76, 86 microscopy, 77, 91–92
0	particle counters, 85
Online measurement, 86–87 Open-yard storage, 102	screening, 77, 92–93 sedimentation, 81, 93–94

Particle size distribution, 69	Powder, 8, 32
instrumental techniques, 95	beds behavior, 35
mean, 70	characteristics, 31
measurement, 67	cohesion, 52
median, 70	flow, 48, 49, 120
mode, 70	fluid-related properties, 32
types, 68, 73	Geldart's classification, 34
Particle technology, 6	industrial, 10
Particle Technology Forum, 6	Jenike's classification, 33–34
Particle-free voids. See Bubble	and particle technology, 6, 7
Particulate	pressure–density, 27
fluidization, 252. See also Non-	standard sieves apertures, 9
bubbling fluidization	Powder failure properties, 48, 58
material, 49	
	angle of repose, 61
phase, 262–263	calculations and report, 67
Partition effect, 331	cohesion, 50
Passive devices, 123	corrected failure function
Peaked stress field, 108, 128	measurement, 59
Permeability, 301	description, 50
Permeate, 413	direct measurement, 57
Pervaporation, 410	experimental determinations,
Phase equilibrium, 285	50, 65
Photosedimentometer, 85	instrument and materials, 66
Plane membrane, 414, 416	Jenike yield locus, 52
Plate-and-frame membrane, 417	properties, 61
Plus material, 288, 313	shear cells, 50
Pneumatic conveying, 155, 171, 172–173	shear tester operation, 66
applications, 156, 159	tensile strength measurement, 60
conveyor types, 155	Warren–Spring equation, 52
dense-phase systems, 157–159	yield loci, 52
design and selection, 163	Powder rheometer, 66. See also
dilute-phase systems, 157, 159–163	Rotational shear cell
drawbacks, 156	Power, 429
frictional energy loss, 166–167	equation, 436
granules characteristics, 171–172	law fluids, 319, 354
minimum conveying velocity, 164	Preferential flow, 255
particle friction factor, 167	Prefilt. See Feed slurry
power requirement, 167-	Pressure, 429
pressure drop, 165	dilute-phase conveying system,
Point of incipient fluidization, 255	162, 163
Polyester fiber, 403	filter, 403
Polypropylene fiber, 403	vessel filters, 405, 406
Polythene fiber, 403	Pressure agglomeration, 214, 218
Polyvinyl chloride (PVC), 171	equipments, 216, 217, 221
Porosity, 23, 221	mechanism of, 215
beds, 50	speed limitation, 215
of monosized particles, 208	Pressure filtration. <i>See also</i> Centrifugal
Positive-pressure system, 162. See	filtration
Dilute-phase conveying system	
Diffute-priase conveying system	alternative, 405

comparison, 382	Relative Wear Number, 131–132
constant, 396–399, 408	Repose angle, 61
Process engineering literature, 31	drained, 62
Push system dilute-phase conveying.	poured, 61
See also Pressure—dilute-	Resultant force, 321
phase conveying system	Reverse osmosis (RO), 409, 410
PVC. See Polyvinyl chloride (PVC)	desalting, 410–411
Pycnometry, 18	rejection mechanisms, 411, 412
air, 21–23	retention limit, 410
liquid, 19	Revolving cylindrical screens, 298–300
1 '	Rheogram, 360
	for different fluids, 319
Q	of suspensions, 359
Quartz, 8, 311	Rheology, 317–318
Feldspar-quartz, 309	Rheopectic, 318, 320
	Rhombohedral particle, 208, 209
Hematite-quartz, 309	Ribbon
nonconductive materials, 307	
relaxing time for, 311	continuous, 197
	mixer, 240–241, 242
R	powder blender, 214
D 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	screws, 151
Rack-and-pinion gate, 132, 133	stainless-steel filtration, 303
Radial separation zones, 380	type mixer, 242
Radial velocity, 337, 379	Richardson and Zaki equation, 369
Raleigh, Lord, 435	Ring shear cell, 50
Random coalescence, 213	Rittniger's law, 183
Random mixture, 231	RO. See Reverse osmosis (RO)
Random variable, 39. See also Sampling	Rod mills, 199
sample size, 40	Roller and Svenson functions, 73
unknown population mean, 40	Rosin–Rammler functions, 73
Rathole, 103	Rotary
Rayon fiber, 403	crushers, 186
Reciprocating screens, 297	drum vacuum filter, 403, 406, 407
Reconstitutability, 28	knife cutters, 180, 194
attrition, 275	lobe blower, 160
tumbling agglomeration, 224	sample divider, 48
Reduced efficiency, 327. See also Net	tangential-spray method, 273
efficiency	vacuum disc filter, 403, 407
flow splitting effect compensation, 421	vacuum precoat filters, 407–408
Reduction Relation Determination	Rotary knife cutters, 194. See also
calculations and report, 205	Hammer mills
equipment and materials, 204	Rotary valve feeder. See Star valve
hammer mill operation, 205	feeder
instruments, 205	Rotational shear box. See Ring shear
introduction, 204	cell
screen analysis, 205	Rotational shear cell, 50, 55. See also
Reels, 298	Powder rheometer; Shear cell
Rejection efficiency, 328 Relative density, See Specific gravity	Instrument, 66
Relative density. See Specific gravity	Rubbing wear. See Abrasive wear

S	mesh size, 77
Salt rejection, 412. See also Reverse	procedure, 93, 313–314
osmosis (RO)	ratio of aperture, 77–78
Saltation velocity, 163, 164, 168	sieve choice, 92
Sample	sieve dimensions, 78
dividers, 46, 47, 48	sieving analysis, 78
reducer, 47	solid mixtures, 287
size, 43, 233	Screw conveyors, 151
Sampling, 37, 232	capacity calculation, 155
confidence interval, 40–41, 42	components, 152
coning and quartering, 46	material factors for, 154
grab, 45	power requirement, 153
methods, 45	Screw feeder, 127, 128, 129
population standard deviation	Scroll centrifuge, 390, 392
estimation, 41	Scrubbers, 348. See also Solid-liquid
random variable, 39	separation
rules, 38	applications, 351
segregation, 45	equipment, 349–351
statistical theory, 39	process description, 349
stopped belt, 46	separation number, 349
theory of, 232	separation vs. separation number
thief, 45	efficiency, 350
Saran fiber, 403	spray cyclone scrubber, 351
Scale of scrutiny, 232	Scrubbing, 285
Schuhmann slope, 74	Second-order test procedure, 376
Schulze ring shear tester, 54	Sedimentation, 81, 285, 367
Scraper conveyors, 145, 146	Andreasen's sedimentation
capacity of, 147	pipette, 83
material friction coefficient, 147	application of, 367
power requirement, 146, 148	calculations and report, 94–95
Screen, 296	centrifugal, 82, 378, 420
aperture size, 182	equipment principles, 370
British Standard sieves, 438–439	factors affecting settling, 367–368
extruder, 216	free settling, 368
Grizzlies, 296	gravity, 82, 371, 420
oscillating, 296, 298	hindered settling, 368
reciprocating, 297	mass balance on, 374–375
reels, 298	pipette centrifuge, 84
revolving cylindrical, 298–300	procedure, 94
shaking, 296	rate stages, 369–370
US sieve, 437–438	Richardson and Zaki equation, 369
vibratory, 296	sample preparation, 94
Screening, 77, 312–313	second-order test procedure, 376
calculations and report, 93, 314	settling rate acceleration, 377–378
centrifuge, 390	Segregation, 230, 239
clogging, 287–288	Separation
efficiency of, 289–296, 315	mechanism, 68
equipments, 296	number, 349
- *	techniques, 7, 284–285

Separation, approximate limit of, 332	stress distribution in, 109
Settling	types, 104
flux vs. concentration, 374	wall load, 110, 134–135
progressive, 369	Sinkability, 28
rate acceleration, 377–378	Size enlargement, 207. See also
test, 365–366	Agglomeration
velocity, 376	Size reduction, 179. See also
Shaking screens, 296	Comminution
Sharpness index, 331	Bond Work Index, 184
Shear	Bond's law, 184
angle of internal friction, 53	comminuted materials
comminution, 190	properties, 181
diameter, 11	energy requirements in, 182
mixing, 230	equipment selection guide, 203
on powder, 32	forces involved, 180-181
rate of, 113, 319	Kick's law, 183
strength, 124	processes selection criteria, 200-203
stress, 52	Rittniger's law, 183
testing, 65	Size reduction equipment, 184, 185, 206
thickening, 318, 319, 356	agitation mills, 194
thinning, 318, 356	angle of nip, 195
Shear cell	Ball mill, 192, 198, 200
annular, 54	characteristics, 184
Jenike shear cell, 50	classification, 184
rotational, 55, 56, 66	crushers, 185–187, 194–195
tester, 33	crushing rolls, 195-197, 203-204, 206
types, 50	cutting machines, 194
Shear thinning and thickening, 318	disc attrition mills, 189–191
non-Newtonian viscosity effects, 356	fluid-energy (jet) mills, 194
rheograms, 319	hammer mill, 189, 190, 192, 204–205
Sieves, 287	Mikro-atomizer, 193
choice of, 92	operation of, 195–199
dimensions of, 78	rod mills, 199
mesh size, 77	rollers mills, 188–189, 199–200
standard sieves apertures, 9	tumbling mills, 191–192, 197
Tyler sieves, 78	ultrafine grinding, 192
Sieving, 285. See also Screening	Slurry, 356. See also Decanter
Sigma theory equation, 420	centrifuge; Filter—presses;
Silhouettes	Vacuum filters
as closed curve, 17	Solid
at different amplitudes, 16	bulk density, 23
polar coordinates in irregular, 16	fluid separators, 331
Silos, 101. See also Bins; Hopper	free-flowing, 112, 115
design considerations, 103	hardness scale, 7–8
flow limits, 115	mixing, 239, 244
flow problems in, 103	mixtures separation, 287
grain, 101	packing properties, 23, 26
Janssen equation, 110	particle density, 18
natural discharge, 112	particle size and shape, 9
	particle office and offupe,

Calid (Continued)	massure dram 265 266
Solid (<i>Continued</i>) particles characterization, 67	pressure drop, 265, 266 stability, 266
primary properties, 8	Spray cyclone scrubber, 351
sampling, 232	Spun glass fiber, 403
in suspension, 113	Stairmand Cyclone design, 337–338
	Star feeder, 127, 128, 129
Solid–gas separation, 333 air classifiers, 334, 340–344	
analytical cut point, 343	Star valve feeder, 160, 161 Statistical diameters, 11, 91
bag filter, 345, 346, 347, 348	Stokes
blow ring, 347	diameter, 10, 12, 81, 85, 90, 334.
cake resistance factor, 348	See also Diameter
cyclones, 334–340	law, 84, 322, 367
devices, 242–243	number, 385, 386, 387, 419
electrostatic separators, 352	number relationship, 419
envelope filter, 345, 347	Stopped belt sampling, 46. See also
fabric properties, 346, 347	Sampling
gas filters, 345–38	Stream scanning, 87
scrubbers, 348–351	in size measurement, 76
separation vs. separation number	Stress, 429. See also Yield locus
efficiency, 350	arched, 128, 130
settling chambers, 351–352	compacting, 60
spray cyclone scrubber, 351	consolidation, 33, 34, 58, 173
Solid–liquid separation, 334, 353.	critical, 122
See also Sedimentation;	distribution in silos, 109
Settling—test;	peaked, 108, 128
Suspensions—rheograms of	powder compressibility, 247
cake formation, 365	shear, 51, 53, 103
clarification, 365	tensile, 53
colloidal forces, 354	wall, 108
dimensionless scale-up of	yield stress, 33, 57
equipment, 419	Structured storage. See Open-yard
equipment, 326, 363	storage
fanning friction factor, 355	Superconductors, 305
laminar flow, 354, 355	Superficial velocity, 251
Newtonian friction factor, 357	pressure drop, 395
non-Newtonian flow, 354, 356-357	Surface. See also Coefficient of friction
power-law constant, 354	active agents, 28
rheology and flow of, 353	angle of friction, 50
shear thinning and thickening, 356	area-to-mass ratio, 28
slurry, 356	bubble eruption, 262
Solubility, 29	conducting, 308
Specific gravity, 18	equivalent diameter, 210
Sphericity, 13	green strength, 220
Spinning riffler, 46, 48. See also	idler pulley, 138
Sample—dividers	moisture transfer, 272
Spiral-wound membrane, 417	of particles, 14
Spokes. See Hydrocyclone—bodies	Rittinger's constant, 183
Spouted bed, 264	screen, 287, 297
application, 265	for storage, 102

Transmembrane pressure gradient. See
Transmembrane pressure (TMP)
Transmission relations, 142
Tripper, 139–140
power to operate, 141
Trommels. See Reels
Tromp curve. See Grade efficiency
curve
Trunnion overflow mill, 192, 193
Tubular centrifuge, 388, 389
Tumbler mixer, 239
agitation device, 243
in food powder blending, 240
Tumbler mixers, Y-cone, 239, 240
capacity factor, 243
Tumbling
agglomeration, 210, 212–214
in friability tester, 276
green agglomerates, 223
mills, 185, 191–192, 197
mixers, 239, 240, 244–246
vessels, 242
Tumbling agglomeration, 212. <i>See also</i> Attrition
equipment for, 214
in food processing, 224
kinetics of, 213
method comparison, 224–226
quantity of bonding liquid, 216
Tumbling agglomeration of powders, 224
calculations and report, 226
drum agglomerator operation, 225
equipment and materials, 225
friability test, 226
instruments, 225
pan agglomerator operation, 225
particle size distribution
measurement, 225
Tyler sieves, 78. See also Screening
U
U
UF. See Ultrafiltration (UF)
Ultrafiltration (UF), 409
nomenclature of, 414-415
perforation, 414
in processing industry, 416
rejection mechanisms, 411, 412
retention limit, 410

Ultrafine	pycnometry, 19, 21
grinders, 185, 202	surface-to-volume diameter, 13
grinding, 180, 192	of voids, 23
particles in floc, 361	
powders, 24, 95, 182	W
Unconfined yield stress, 57. See also	VV
Failure function (FF)	Wall friction angle, 53, 131
stress against, 173	of crushed limestone of, 172–173
yield locus, 33	hopper half angle, 115
Underflows to-throughput ratio, 326	Wall stresses
Undersize particle, 288, 313	in axisymmetrical bins, 108–112
Unit operations, 3–6	in other vessel types, 112
heat transfer, 5	Warren–Spring equation, 52
traditional operations, 5	Water. See also Moisture;
Unit process, 4	Reconstitutability; Wetting
US sieve series, 437–438	absorbents, 132
	dust suppression, 202, 243
V	fiber water absorbance, 403
	layer in membrane, 411
Vacuum filters, 403, 406–408	sea water desalting, 410
Valve-discharge centrifuge, 390	spray cyclone scrubber, 350
V-cone tumbler mixers, 240	in ultrafine grinding, 202
Venturi feeder, 160, 161	Wet collectors. <i>See</i> Scrubbers
injector, 194	Wettability, 28
scrubbers, 351	in agglomeration method selection,
Vibratory. See also Abrasive wear	220
feeder, 127, 128, 129	Wetting, 28
screens, 296	Wheel. See Hydrocyclone—deck
Viscoelastic fluid, 318, 320	Wool fiber, 402
Viscometer, rotational, 360	Wurster process. See Fluidized-bed
Viscosity, 319. See also Shear—thinning	coating
absolute, 113	
air, 433	Y
dynamic, 429	V t
Geldart's classification, 36	Y-cone, tumbler mixers, 239, 240
shear rate dependence, 354	capacity factor, 243 Yield locus, 33
water, 431 Voidage. <i>See</i> Porosity	
mean bed, 263	angle of internal friction, 53
Volume, 429. See also Density	bulk solid strength, 124–125 failure properties, 64
calculation, 14	Jenike, 52
equivalent sphere diameters, 10	shear stress, 55
equivalent spriere diameters, 10	silear stress, oo