

Handbook of Process Chromatography

Development, Manufacturing, Validation and Economics



Lars Hagel, Günter Jagschies and Gail Sofer



William S. Levine

Handbook Of Process Chromatography Second Edition Development Manufacturing Validation And Economics: Handbook of Process Chromatography Gunter Jagschies, Gail K. Sofer, Lars Hagel, 2007-12-08 This book will update the original edition published in 1997 Since the publication of the first edition the biotechnology and biologics industries have gained extensive knowledge and experience in downstream processing using chromatography and other technologies associated with recovery and purification unit operations. This book will tie that experience together for the next generation of readers Updates include sources and productivity types of products made today experiences in clinical and licensed products economics current status of validation illustrations and tables automated column packing automated systemsNew topics include the use of disposables multiproduct versus dedicated production design principles for chromatography media and filters ultrafiltration principles and optimization risk assessments characterization studies design space platform technologies process analytical technologies PATs biogenerics comparability assessments Key Features new approaches to process optimization use of patform technologies applying risk assessment to process design **Handbook of Biogeneric Therapeutic Proteins** Sarfaraz Niazi,2024-04-15 Since 1972 which marks the invention of recombinant engineering more than 500 therapeutic proteins have been approved for clinical use Today biological drugs constitute almost 70% of all new drugs and have a biological origin The first edition of this book dealt with biosimilars and this edition i e the second edition focuses on new drugs yet limits to therapeutic proteins Newer technologies for drug development represent the updated topics in the book and include repur posing AI driven identification of newer designs novel expression systems manufacturing using these systems rapidly changing regulatory pathways and legal hurdles This edition discusses how to identify develop manufacture and take multibillion dollar products to market within the shortest possible time Features Complete and thorough coverage of the regulatory and technological challenges of developing generic therapeutic proteins Comprehensive discovery to market newer technologies regulatory planning and IP hurdles are included that are not found elsewhere Expanded volume that must be in the hands of every company interested in biological drugs including the mRNA based biopharmaceutical companies fast appearing on the market Discusses how to identify develop manufacture and take multibillion dollar products to market in the shortest possible time Renowned author and entrepreneur in the field of drug Process Architecture in Biomanufacturing Facility Design Jeffery Odum, Michael C. discovery and production Flickinger, 2017-11-03 Essential information for architects designers engineers equipment suppliers and other professionals who are working in or entering the biopharmaceutical manufacturing field Biomanufacturing facilities that are designed and built today are radically different than in the past The vital information and knowledge needed to design and construct these increasingly sophisticated biopharmaceutical manufacturing facilities is difficult to find in published literature and it s rarely

taught in architecture or design schools This is the first book for architects and designers that fills this void Process Architecture in Biomanufacturing Facility Design provides information on design principles of biopharmaceutical

manufacturing facilities that support emerging innovative processes and technologies use state of the art equipment are energy efficient and sustainable and meet regulatory requirements Relying on their many years of hands on design and operations experience the authors emphasize concepts and practical approaches toward design construction and operation of biomanufacturing facilities including product process facility relationships closed systems and single use equipment aseptic manufacturing considerations design of biocontainment facility and process based laboratory and sustainability considerations as well as an outlook on the facility of the future Provides guidelines for meeting licensing and regulatory requirements for biomanufacturing facilities in the U S A and WHO especially in emerging global markets in India China Latin America and the Asia Pacific regions Focuses on innovative design and equipment to speed construction and time to market increase energy efficiency and reduce footprint construction and operational costs as well as the financial risks associated with construction of a new facility prior to the approval of the manufactured products by regulatory agencies Includes many diagrams that clarify the design approach Process Architecture in Biomanufacturing Facility Design is an ideal text for professionals involved in the design of facilities for manufacturing of biopharmaceuticals and vaccines biotechnology and life science industry including architects and designers of industrial facilities construction equipment vendors and mechanical engineers It is also recommended for university instructors advanced undergraduates and graduate students in architecture industrial engineering mechanical engineering industrial design and industrial interior design Technology Z F Cui, H S Muralidhara, 2010-09-23 Membrane technology is a rapidly developing area with key growth accross the process sector including biotech separation and biomedical applications e g haemodialysis artificial lungs through to large scale industrial applications in the water and waste water processing and the food and drink industries As processes mature and the cost of membranes continues to dramatically reduce so their applications and use are set to expand Process engineers need access to the latest information in this area to assist with their daily work and to help to develop and apply new and ever more efficient liquid processing solutions This book covers the latest technologies and applications with contributions from leading figures in the field Throughout the emphasis is on delivering solutions to practitioners Real world case studies and data from leading organizations including Cargill Lilly Microbach ITT mean this book delivers the latest solutions as well as a critical working reference to filtration and separation professionals Covers the latest technologies and applications in this fast moving bioprocessing sector Presents a wide range of case studies that ensure readers benefit from the hard won experience of others saving time money and effort World class author team headed up by the Chair of Chemical Engineering at Oxford University UK and the VP of Plant Operations and Process Technology at Cargill Corp the food services company and largest privately owned company in the US **Process Scale Purification of Antibodies** Uwe Gottschalk, 2017-03-07 Promoting a continued and much needed renaissance in biopharmaceutical manufacturing this book covers the different strategies and assembles top tier technology experts to address the challenges of antibody purification

Updates existing topics and adds new ones that include purification of antibodies produced in novel production systems novel separation technologies novel antibody formats and alternative scaffolds and strategies for ton scale manufacturing Presents new and updated discussions of different purification technologies focusing on how they can address the capacity crunch in antibody purification Emphasizes antibodies and innovative chromatography methods for processing Martin's Physical Pharmacy and Pharmaceutical Sciences Patrick J. Sinko, 2023-02-08 Consistently revised and updated for more than 60 years to reflect the most current research and practice Martin's Physical Pharmacy and Pharmaceutical Sciences 8th Edition is the original and most comprehensive text available on the physical chemical and biological principles that underlie pharmacology and the pharmaceutical sciences An ideal resource for PharmD and pharmacy students worldwide teachers researchers or industrial pharmaceutical scientists this 8th Edition has been thoroughly revised enhanced and reorganized to provide readers with a clear consistent learning experience that puts essential principles and concepts in a practical approachable context Updated content reflects the latest developments and perspectives across the full spectrum of physical pharmacy and a new full color design makes it easier than ever to discover distinguish and understand information providing users the most robust support available for applying the elements of biology physics and chemistry in work or study

Advances in Vinegar Production Argyro Bekatorou, 2019-09-11 In industrial vinegar production there are three main types of methods involved the slow handcrafted traditional method Orleans or French method and the rapid submerged and generator methods The current trend is to fuse traditional techniques with state of the art technologies and a variety of approaches have been developed to increase fermentation efficiency and reduce cost and fermentation time This book reports on all the recent innovations in vinegar production and compares them to the traditional submerged fermentation systems The new trends on raw materials substrate pretreatment strategies alcoholic fermentation and acetitification systems are also reviewed Biopharmaceutical Processing Gunter Jagschies, Eva Lindskog, Karol Lacki, Parrish M. Galliher, 2018-01-18 Biopharmaceutical Processing Development Design and Implementation of Manufacturing Processes covers bioprocessing from cell line development to bulk drug substances The methods and strategies described are essential learning for every scientist engineer or manager in the biopharmaceutical and vaccines industry. The integrity of the bioprocess ultimately determines the quality of the product in the biotherapeutics arena and this book covers every stage including all technologies related to downstream purification and upstream processing fields Economic considerations are included throughout with recommendations for lowering costs and improving efficiencies Designed for quick reference and easy accessibility of facts calculations and guidelines this book is an essential tool for industrial scientists and managers in the biopharmaceutical industry Offers a comprehensive go to reference for daily work decisions Covers both upstream and downstream processes Includes case studies that emphasize financial outcomes Presents summaries decision grids graphs and overviews for quick reference Chemical Engineering Design Ray Sinnott, Gavin Towler, 2019-05-26 Chemical

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Chemical Engineering Design Gavin Towler, Ray Sinnott, 2012-01-25 Chemical Engineering Design Second Edition deals with the application of chemical engineering principles to the design of chemical processes and equipment Revised throughout this edition has been specifically developed for the U S market It provides the latest US codes and standards including API ASME and ISA design codes and ANSI standards It contains new discussions of conceptual plant design flowsheet development and revamp design extended coverage of capital cost estimation process costing and economics and new chapters on equipment selection reactor design and solids handling processes A rigorous pedagogy assists learning with detailed worked examples end of chapter exercises plus supporting data and Excel spreadsheet calculations plus over 150 Patent References for downloading from the companion website Extensive instructor resources including 1170 lecture slides and a fully worked solutions manual are available to adopting instructors This text is designed for chemical and biochemical engineering students senior undergraduate year plus appropriate for capstone design courses where taken plus graduates and lecturers tutors and professionals in industry chemical process biochemical pharmaceutical petrochemical sectors New to this edition Revised organization into Part I Process Design and Part II Plant Design The broad themes of Part I are flowsheet development economic analysis safety and environmental impact and optimization Part II contains chapters on equipment design and selection that can be used as supplements to a lecture course or as essential references for students or practicing engineers working on design projects New discussion of conceptual plant design flowsheet development and revamp design Significantly increased coverage of capital cost estimation process costing and economics New chapters on equipment selection reactor design and solids handling processes New sections on fermentation adsorption membrane separations ion exchange and chromatography Increased coverage of batch processing food pharmaceutical and biological processes All equipment chapters in Part II revised and updated with current information Updated throughout for latest US codes and standards including API ASME and ISA design codes and ANSI standards Additional worked examples and homework problems The most complete and up to date coverage of equipment selection 108 realistic commercial design projects from diverse industries A rigorous pedagogy assists learning with detailed worked examples end of chapter

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however developing new successful biological drugs presents many challenges including labor intensive production processes tighter regulatory controls and increased market competition This book reviews the current state of the science offering readers a single resource that sets forth the fundamentals as well as tested and proven development strategies for biological drugs Moreover the book prepares readers for the challenges that typically arise during drug development offering straightforward solutions to improve their ability to pass through all the regulatory hurdles and deliver new drug products to the market Biological Drug Products begins with general considerations for the development of any biological drug product and then explores the strategies and challenges involved in the development of specific types of biologics Divided into five parts the book examines Part 1 General Aspects Part 2 Proteins and Peptides Part 3 Vaccines Part 4 Novel Biologics Part 5 Product Administration Delivery Each chapter has been prepared by one or more leading experts in biological drug development Contributions are based on a comprehensive review and analysis of the current literature as well as the authors first hand experience developing and testing new drugs References at the end of each chapter serve as a gateway to original research papers and reviews in the field By incorporating lessons learned and future directions for research Biological Drug Products enables pharmaceutical scientists and students to improve their success rate in developing new biologics to treat a broad range of human diseases Handbook of Therapeutic Antibodies Stefan Dübel, Janice M. Reichert, 2014-08-04 Still the most comprehensive reference source on the development production and therapeutic application of antibodies this second edition is thoroughly updated and now has 30% more content Volume 1 covers selection and engineering strategies for new antibodies while the second volume presents novel therapeutic concepts and antibodies in clinical study as well as their potential Volumes 3 and 4 feature detailed and specific information about each antibody approved for therapeutic purposes including clinical data This unique handbook concludes with a compendium of marketed monoclonal antibodies and an extensive index Beyond providing current knowledge the authors discuss emerging technologies future developments and intellectual property issues such that this handbook meets the needs of academic researchers decision makers in industry and healthcare professionals in the clinic Validation of Chromatography Data Systems Robert D McDowall, 2016-11-23 Guiding chromatographers working in regulated industries and helping them to validate their chromatography data systems to meet data integrity business and regulatory needs This book is a detailed look at the life cycle and documented evidence required to ensure a system is fit for purpose throughout the lifecycle Initially providing the regulatory data integrity and system life cycle requirements for computerised system validation the book then develops into a guide on planning specifying managing risk configuring and testing a chromatography data system before release This is followed by operational aspects such as training integration and IT support and finally retirement All areas are discussed in detail with case studies and practical examples provided as appropriate The book has been carefully written and is right up to date including recently released FDA data integrity guidance It provides detailed guidance on good practice and expands on the first edition making

it an invaluable addition to a chromatographer s book shelf Method Validation in Pharmaceutical Analysis Joachim Ermer, Phil W. Nethercote, 2014-08-27 This second edition of a global bestseller has been completely redesigned and extensively rewritten to take into account the new Quality by Design QbD and lifecycle concepts in pharmaceutical manufacturing As in the first edition the fundamental requirements for analytical method validation are covered but the second edition describes how these are applied systematically throughout the entire analytical lifecycle ObD principles require adoption of a systematic approach to development and validation that begin with predefined objectives For analytical methods these predefined objectives are established as an Analytical Target Profile ATP The book chapters are aligned with recently introduced standards and guidelines for manufacturing processes validation and follow the three stages of the analytical lifecycle Method Design Method Performance Qualification and Continued Method Performance Verification Case studies and examples from the pharmaceutical industry illustrate the concepts and guidelines presented and the standards and regulations from the US FDA European EMA and global ICH regulatory authorities are considered throughout The undisputed gold standard in the field EPA Publications Bibliography Quarterly Abstract Bulletin United States. Environmental Protection Agency, 1995-10 Biopharmaceutical Manufacturing Gary Gilleskie, Charles Rutter, Becky McCuen, 2021-09-07 Biopharmaceuticals medicines made by or from living organisms including cells from living organisms are extremely effective in treating a broad range of diseases Their importance to human health has grown significantly over the years as more biopharmaceutical products have entered the market and now the biggest selling drugs in the world are biopharmaceuticals Biopharmaceutical Manufacturing Principles Processes and Practices provides concise comprehensive and up to date coverage of biopharmaceutical manufacturing Written in a clear and informal style the content has been influenced by the authors substantial industry experience and teaching expertise That expertise enables the authors to address the many questions posed over the years both by university students and professionals with experience in the field Consequently the book will appeal both to undergraduate or graduate students using it as a textbook and specialized industry practitioners seeking to understand the big picture of biopharmaceutical manufacturing This book Fossil Energy Update Handbook of Near-Infrared Analysis Donald A. Burns, Emil W. Ciurczak, 2007-09-07 Fast inexpensive and easy to ,1978 use near infrared NIR spectroscopy can be used to analyze small samples of virtually any composition The Handbook of Near Infrared Analysis Third Edition explains how to perform accurate as well as time and cost effective analyses across a growing spectrum of disciplines Presenting nearly 50% new and re

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