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Philip A. Geis

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Pharmaceutical Microbiological Quality Assurance and Control David Roesti, Marcel Goverde, 2020-01-02 Relying on practical examples from the authors experience this book provides a thorough and modern approach to controlling and monitoring microbial contaminations during the manufacturing of non sterile pharmaceuticals Offers a comprehensive guidance for non sterile pharmaceuticals microbiological QA QC Presents the latest developments in both regulatory expectations and technical advancements Provides guidance on statistical tools for risk assessment and trending of microbiological data Describes strategy and practical examples from the authors experience in globalized pharmaceutical Compounding Sterile Preparations Ryan Forrey, Lindsey Amerine, Angela W. companies and expert networks Yaniv, 2023-11-13 The latest edition of Compounding Sterile Preparations by Ryan A Forrey Lindsey B Amerine and Angela W Yaniv reflects the latest advancements in the field providing you with an indispensable resource to navigate the complex landscape of sterile compounding New in this Edition Updated Standards All chapters have undergone extensive revisions to align with the most recent literature and the revised USP standards USP Chapter Now includes information on radiopharmaceutical compounding in USP Chapter Expanded Knowledge Base Two brand new chapters covering Allergenic Extracts and Corrective and Preventative Action CAPA Plans Microbial Limit and Bioburden Tests Lucia Clontz, 2008-10-14 In recent years the field of pharmaceutical microbiology has experienced numerous technological advances accompanied by the publication of new and harmonized compendial methods It is therefore imperative for those who are responsible for monitoring the microbial quality of pharmaceutical biopharmaceutical products to keep abreast of the latest changes Microbial Limit and Bioburden Tests Validation Approaches and Global Requirements guides readers through the various microbiological methods listed in the compendia with easy to follow diagrams and approaches to validations of such test methodologies Includes New and Updated Material Now in its second edition this work is the culmination of research and discussions with technical experts as well as USP and FDA representatives on various topics of interest to the pharmaceutical microbiologist and those responsible for the microbial quality of products materials equipment and manufacturing facilities New in this edition is an entire chapter dedicated to the topic of biofilms and their impact on pharmaceutical and biopharmaceutical operations The subject of rapid methods in microbiology has been expanded and includes a discussion on the validation of alternative microbiological methods and a case study on microbial identification in support of a product contamination investigation Substantially updated and revised this book assists readers in understanding the fundamental issues associated with pharmaceutical microbiology and provides them with tools to create effective microbial contamination control and microbial testing programs for the areas under their responsibility

Cosmetic Microbiology Philip A. Geis,2020-12-06 This updated edition provides research scientists microbiologists process engineers and plant managers with an authoritative resource on basic microbiology manufacturing hygiene and

product preservation It offers a contemporary global perspective on the dynamics affecting the industry including concerns about preservatives natural ingredients small manufacturing resistant microbes and susceptible populations Professional researchers in the cosmetic as well as the pharmaceutical industry will find this an indispensable textbook for in house training that improves the delivery of information essential to the development and manufacturing of safe high quality Specification of Drug Substances and Products Christopher M. Riley, K. Lien Nguyen, 2024-09-22 Specification of Drug Substances and Drug Products is a fully comprehensive reference on Specification Setting for Pharmaceuticals There have been several recent developments in the ICH Guidelines which were not captured in previous editions notably the new guideline on Development of Analytical Procedure and the revisions to the validation guidelines and the specification quidelines This edition contains chapters discussing the unique requirements for the universal critical quality attributes as well as the specific tests required to characterize and control different types of products ranging in complexity from small molecules in immediate release oral dosage forms to complex products such as drug antibody conjugates and mRNA based products This substantially expanded revision of the 2nd edition will serve as practical comprehensive reference for scientists managers educators and consultants involved in the development and regulation of pharmaceutical products Presents critical assessment potential impact and application of the recent revisions to ICH guidelines on method validation O2 as well as the latest guideline on Analytical Method Development Q14 and the special regional requirements in non ICH regions Addresses comprehensive treatment of the development and validation of analytical methodologies used in the analysis control and specification of a variety of different types of dosage forms ranging from traditional oral solid dosage forms to proteins nRNA based drugs vaccines and gene therapy This book will also address drug device combination products such as digital drug delivery systems transdermal systems and inhalation products Presents detailed treatment of latest statistical approaches including new approaches to the treatment of validation data method specification setting and shelf life prediction based on Sterility, Sterilisation and Sterility Assurance for Pharmaceuticals Tim Sandle, 2025-08-01 Sterility stability data Sterilisation and Sterility Assurance for Pharmaceuticals Technology Validation and Current Regulations Second Edition is an in depth guide to the world of pharmaceutical sterilization This new edition has been updated to reflect the latest standards and regulations ensuring alignment with current practices It explores emerging methods and techniques complemented by new case studies that provide practical examples Readers will gain comprehensive knowledge about sterilization s critical role in healthcare and pharmaceutical manufacturing highlighting the importance of controlling microbial challenges to ensure product safety and patient well being The book discusses sterility sterilization methods such as gamma radiation e beam dry heat steam gas vapor filtration and new techniques like X ray sterilization liquid phase sterilization ultraviolet light supercritical gases and sterilization assurance governance It covers biopharmaceutical manufacturing processes including aseptic filling container and packaging design and cleanroom environments This edition is essential for professionals in

pharmaceuticals healthcare and medical device manufacturing providing the knowledge needed to comply with current standards and regulations Includes nine new chapters with many new case studies Offers coverage on the most current standards and regulations Provides full coverage of novel sterilization methods Acoustics of Layered Media II Leonid M. Brekhovskikh, Oleg A. Godin, 2013-04-17 This is the sequel to our book Acoustics of Layered Media I Plane and Quasi Plane Waves Springer Ser Wave Phenom Vol 5 Taken together these two monographs present a systematic exposition of the theory of sound propagation in inhomogeneous media which starts from first principles and includes recent results More advanced topics are considered in this second volume Although the theory of wave beams and fields of localized sources is more sophisticated than the theory of quasi plane waves it embraces a much wider range of interesting problems that are also important for applications We exploit the results of Acoustics of Layered Media I as long as it is expedient to consider sound fields as a superposition of plane or quasi plane waves However the knowledgeable reader will view this book as self contained Similar topics have been treated in the book by L M Brekhovskikh Waves in Layered Media the English version of the second edition of which was published by Academic Press in 1980 Since Waves in Layered Media became very popular we have tried here to retain its spirit However the majority of this text is devoted to new material which reflects the significant progress of the theory during recent years In particular acoustic fields in a moving fluid are considered and much attention is paid to sound propagation in range dependent environments which is currently on the leading edge of research The Pharmacy Technician, 7e Perspective Press, 2020-01-15 Endorsed by the American Pharmacists Association activities APhA The Pharmacy Technician 7e is a valuable tool for pharmacy technician students This applied accessible book is a practical text for understanding the principles career concepts and pharmacy skills needed to be a successful pharmacy technician It offers clear concise information to help students learn the material and pass the national certification exams the Pharmacy Technician Certification Exam PTCE and the Exam for Certification of Pharmacy Technicians ExCPT This book was designed to be accompanied by The Pharmacy Technician Workbook Certification Review 7e to help prepare for the certification exams This textbook aligns with the Fifth Edition of the American Society of Health System Pharmacists ASHP Model Curriculum for Pharmacy Technician Education and Training Programs and the 2020 content outline for the Pharmacy Technician Certification Examination PTCE **Pharmaceutical Microbiology** Tim Sandle, 2015-10-09 Pharmaceutical Microbiology Essentials for Quality Assurance and Quality Control presents that latest information on protecting pharmaceutical and healthcare products from spoilage by microorganisms and protecting patients and consumers With both sterile and non sterile products the effects can range from discoloration to the potential for fatality The book provides an overview of the function of the pharmaceutical microbiologist and what they need to know from regulatory filing and GMP to laboratory design and management and compendia tests and risk assessment tools and techniques These key aspects are discussed through a series of dedicated chapters with topics covering auditing validation data analysis bioburden toxins

microbial identification culture media and contamination control Contains the applications of pharmaceutical microbiology in sterile and non sterile products Presents the practical aspects of pharmaceutical microbiology testing Provides contamination control risks and remediation strategies along with rapid microbiological methods Includes bioburden endotoxin and specific microbial risks Highlights relevant case studies and risk assessment scenarios Conceptual Development of Industrial Biotechnology for Commercial Production of Vaccines and Biopharmaceuticals Basanta Kumara Behera, 2023-03-07 Conceptual Development of Industrial Biotechnology for Commercial Production of Biopharmaceuticals and Vaccines provides insights on how to bring sustainability into biologic drug production. The cumulative facts and figures within in the book are helpful to promoters in monitoring value chain transfer process of super quality biologics for better return in profits In addition this is a useful reference for students researchers and scientists in biotechnology pharmaceutical science medical sciences and the R D division of biotechnology based industries Conceptual development of biotechnology has taken new avenues with the integration of medical sciences physical science and engineering hence this is a timely source The current global market for vaccines especially COVID 19 is tremendous Bivalent oral polio vaccine diphtheria tetanus containing and measles containing vaccines have a high demand internationally and recombinant DNA technology and protein engineering are helpful in the production of quality bio products Informs how biotechnology and pharmaceutical industries act as central pillars for the stable production of value added biological drugs and vaccines from genetically engineered suitable vectors like microbe or cell lines from animals mammals or plants Highlights various traditional and modern techniques used for improvising the quality of suitable vectors to produce biologic drugs and vaccines under GMP manufacturing facilities Provides updated information on the latest microchip based bioreactors disposable bag bioreactors and animal systems as bioreactors to produce biologic drugs like Smart Biomolecules next generation therapeutics Bio similar drugs Bio betters and antibody drug conjugates Explains how the closed bioreactors with proper mechanical amendments are used for vaccine The Certified Pharmaceutical GMP Professional Handbook Mark Allen Durivage, 2016-05-23 The purpose of production this handbook is to assist individuals for the Certified Pharmaceutical Good Manufacturing Practices Professional CPGP examination and provide a reference for the practitioner The second edition reflects the Body of Knowledge which was updated in 2015 This edition has also incorporated additional information including updated references The updates reflect the current trends and expectations of the evolving pharmaceutical industry driven by consumer expectations and regulatory oversight This handbook covers compliance with good manufacturing practices GMPs as regulated and guided by national and international agencies for the pharmaceutical industry It covers finished human and veterinary drugs and biologics and combination devices as well as their component raw materials including active pharmaceutical ingredients APIs and excipients and packaging and labeling operations The Practice of Pharmacy ,1917 The ASQ Certified Pharmaceutical GMP Professional Handbook Mark Allen Durivage, 2024-09-30 The ASO Certified Pharmaceutical GMP Professional

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