

European Commission

EudraLex - Volume 4 Good manufacturing practice (GMP) Guidelines.

Volume 4 of "The rules governing medicinal products in the European Union" contains guidance for the interpretation of the principles and guidelines of good manufacturing practices for medicinal products for human and veterinary use laid down in Commission Directives 91/356/EEC, as amended by Directive 2003/94/EC, and 91/412/EEC respectively.

Introduction

- Introduction (7/02/2011)
- Commission Directive 2003/94/EC, of 8 October 2003, laying down the principles and guidelines of good manufacturing practice in respect of medicinal products for human use and investigational medicinal products for human use Replacement of Commission Directive 91/356/EC of 13 June 1991 to cover good manufacturing practice of investigational medicinal products.
- Commission Directive 91/412/EEC of 23 July 1991 laying down the principles and guidelines of good manufacturing practice for veterinary medicinal products.

Part I - Basic Requirements for Medicinal Products

- Chapter 1 Pharmaceutical Quality System (into operation since 31 January 2013)
- Chapter 2 Personnel (into operation since 16 February 2014).
- Chapter 3 Premise and Equipment (into operation since 1 March 2015)
 - · See transitional arrangement for toxicological evaluation on page 1 of Chapter 3
 - · Provious
- Chapter 4 Documentation (January 2011)

Eudralex Volume 4 Gmp Guide

John Geigert

Eudralex Volume 4 Gmp Guide:

Good Manufacturing Practice (GMP) Guidelines Mindy J. Allport-Settle, 2009-12 This title combines all of the human and veterinary Regulations Directives and guidance for medicinal products used by the pharmaceutical industry as their main source when manufacturing and distributing medicinal products in the European Union **EU Annex 11 Guide to** Computer Validation Compliance for the Worldwide Health Agency GMP Orlando Lopez, 2015-04-06 Good Manufacturing Practice GMP ensures medicinal products are produced consistently and controlled to the quality standards appropriate for their intended use and as required by product specifications or marketing authorization Annex 11 details the European Medicines Agency EMA GMP requirements for computer systems The purpose of Annex 11 is Nuclear Medicine and Molecular Imaging for Physicists Michael Ljungberg, 2022-03-22 This state of the art handbook the third and final in a series that provides medical physicists with a comprehensive overview into the field of nuclear medicine focuses on highlighting the production and application of radiopharmaceuticals With this the book also describes the chemical composition of these compounds as well as some of the main clinical applications where radiopharmaceuticals may be used Following an introduction to the field of radiopharmacy three chapters in this book are dedicated towards in depth descriptions of common radionuclides and radiopharmaceuticals used during diagnostic studies utilizing planar Single Photon Emission Computed Tomography SPECT imaging in addition to during Positron Emission Tomography PET imaging and finally radiotherapy These chapters are followed by those describing procedures relating to quality control and manufacturing good manufacturing practices also encompassing aspects such as environmental compliance Furthermore this volume illustrates how facilities handling these chemicals should be designed to comply with set regulations Like many pharmaceuticals the development of radiopharmaceuticals relies heavily on the use of mouse models Thus the translation of radiopharmaceuticals i e the process undertaken to assure that the functionality and safety of a newly developed drug is maintained also in a human context is covered in a later chapter This is followed by a chapter emphasising the importance of safe waste disposal and how to assure that these procedures meet the requirements set for the disposal of hazardous waste Several chapters have also been dedicated towards describing various medical procedures utilizing clinical nuclear medicine as a tool for diagnostics and therapeutics As physicists may be involved in clinical trials a chapter describing the procedures and regulations associated with these types of studies is included This is followed by a chapter focusing on patient safety and another on an imaging modality not based on ionizing radiation ultrasound Finally the last chapter of this book discusses future perspectives of the field of nuclear medicine This text will be an invaluable resource for libraries institutions and clinical and academic medical physicists searching for a complete account of what defines nuclear medicine The most comprehensive reference available providing a state of the art overview of the field of nuclear medicine Edited by a leader in the field with contributions from a team of experienced medical physicists chemists engineers scientists and clinical medical

personnel Includes the latest practical research in the field in addition to explaining fundamental theory and the field s history A Comprehensive and Practical Guide to Clinical Trials Delva Shamley, Brenda Wright, 2017-06-07 A Comprehensive and Practical Guide to Clinical Trials provides an overview of the entire process of clinical research in one thorough and easy to read handbook that offers those involved in clinical research a clear understanding of how the components of a study are related It focuses on the practical aspects of the preparation and execution of a clinical trial and offers tools and resources to help the entire team understand how their responsibilities tie together with the tasks and duties of other members This allows for better planning and prioritization and can lead to more effective and successful clinical trials With practical examples checklists and forms this book is a useful guide for planning and conducting clinical trials from beginning to end Describes the entire clinical trial management process from start to finish in a step by step guide Provides best practice elements including case studies practical examples activities and checklists Data Integrity and Data Governance R. D. McDowall, 2018-11-09 This book provides practical and detailed advice on how to implement data governance and data integrity for regulated analytical laboratories working in the pharmaceutical and allied industries

Laboratory Control System Operations in a GMP Environment David M. Bliesner, 2020-04-27 Develop an understanding of FDA and global regulatory agency requirements for Laboratory Control System LCS operations In Laboratory Control System Operations in a GMP Environment readers are given the guidance they need to implement a CGMP compliant Laboratory Control System LCS that fits within Global Regulatory guidelines Using the Quality Systems Approach regulatory agencies like the FDA and the European Medicine Agency have developed a scheme of systems for auditing pharmaceutical manufacturing facilities which includes evaluating the LCS In this guide readers learn the fundamental rules for operating a CGMP compliant Laboratory Control System Designed to help leaders meet regulatory standards and operate more efficiently the text includes chapters that cover Laboratory Equipment Qualification and Calibration Laboratory Facilities Method Validation and Method Transfer Laboratory Computer Systems Laboratory Investigations as well as Data Governance and Data Integrity The text also includes chapters related to Laboratory Managerial and Administrative Systems Laboratory Documentation Practices and Standard Operating Procedures and General Laboratory Compliance Practices Additionally a chapter outlining Stability Program operations is included in the text In addition to these topics it includes LCS information and tools such as End of chapter templates checklists and LCS guidance to help you follow the required standards Electronic versions of each tool so users can use them outside of the text An In depth understanding of what is required by the FDA and other globally significant regulatory authorities for GMP compliant systems For quality assurance professionals working within the pharmaceutical or biopharma industries this text provides the insight and tools necessary to implement government defined regulations WHO Expert Committee on Specifications for Pharmaceutical Preparations WHO Expert Committee on Specifications for Pharmaceutical

Preparations, World Health Organization, 2014 The Expert Committee on Specifications for Pharmaceutical Preparations works towards clear independent and practical standards and guidelines for the quality assurance of medicines Standards are developed by the Committee through worldwide consultation and an international consensus building process The following new guidelines were adopted and recommended for use in addition to 20 monographs and general texts for inclusion in The International Pharmacopoeia and 11 new International Chemical Reference Substances The International Pharmacopoeia updating mechanism for the section on radiopharmaceuticals WHO good manufacturing practices for pharmaceutical products main principles Model quality assurance system for procurement agencies Assessment tool based on the model quality assurance system for procurement agencies aide memoire for inspection Guidelines on submission of documentation for pregualification of finished pharmaceutical products approved by stringent regulatory authorities and Guidelines on submission of documentation for a multisource generic finished pharmaceutical product quality part Challenge of CMC Regulatory Compliance for Biopharmaceuticals John Geigert, 2023-06-15 Each year for the past three years there have been about 50 new molecular medicines approved by the United States Food Addresses current FDA and EMA requirements and expectations for CMC regulatory compliance Now includes CMC regulatory compliance for the new gene based biopharmaceuticals The Transformation of Environmental Law and Governance Sindico, Francesco, Switzer, Stephanie, Qin, Tianbao, 2021-10-19 This cutting edge book considers the functional inseparability of risk and innovation within the context of environmental law and governance Analysing both hard and soft innovation the book argues that approaches to socio ecological risk require innovation in order for society and the environment to become more resilient

Quality assurance of pharmaceuticals: a compendium of guidelines and related materials. Volume 2. Good manufacturing practices and inspection World Health Organization, 2024-01-31 The GMP Compendium for Medical Products is a valuable resource for manufacturers regulators and other stakeholders involved in producing and distributing medical products It covers various topics from quality management systems to personnel hygiene equipment validation and complaint handling The guidance provided is based on the latest scientific and technical knowledge and considers the evolving regulatory landscape and the challenges faced by the industry

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