4 Quality Management System

4.1 General Requirements

See Process Flow Chart FC 4.1-1

Alabama Specialty Products, Inc. has established, documented and implemented a Quality Management System (QMS) in accordance with the requirements of the ISO 9001:2008 Standard, it is not a stand-alone system, but is integrated within ASPI's operating discipline which encompasses the policies, requirements, and work processes of Environment, Health, Safety, Manufacturing, Human Resources and Quality.

Developed and endorsed by company management the QMS ensures that customers' receive quality, reliability and integrity in the products and services ASPI provides them and that customers' needs and requirements are met. The QMS calls for precise adherence to specifications, as well as legal and quality requirements.

Product quality is maintained through systems of standardization and process control. Service quality covers all aspects of customer transactions and is ensured by the function that is providing the service.

4.2 Documentation Requirements

4.2.1 General

The QMS documentation includes:

- A documented Quality Policy and objectives
- A Quality Manual, (requirements for the quality manual are contained in section 4.2.2)
- Documented procedures required by ISO 9001:2008 (i.e., sections 4.2.3, 4.2.4, 8.2.2, 8.3, 8.5.2 and 8.5.3)
- Documents (information and its supporting medium) needed by the organization to ensure the effective planning, operation and control of its processes
- Records required by ISO 9001:2008, (stating results achieved or providing evidence of activities performed)

4.2.2 Quality Manual

This Quality Manual has been prepared to describe Alabama Specialty Products, Inc. QMS. The scope and permissible exclusions of the QMS are described in section one of this manual. Each section of the manual references documented QMS procedures relating to the requirements outlined in that section. The Process Flow Chart, FC 4.1-1, provides a description of the interaction between the processes of the QMS system.

4.2.3 Control of Documents

All the QMS documents are controlled according to the Document Control Procedure, QP 4.2-1.

4.2.4 Control of Quality Records

Quality Records are maintained to provide evidence of conformity to requirements and the effective operation of the QMS. The records are maintained according to the Control of Documents and Quality Records, OP 4.2-1.

Sample Quality Manual Template

Janace E. Bubonia

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Implementing ISO 9000:2000 Matt Seaver, 2001 This text is aimed at the busy manager or proprietor who needs to implement ISO 9001 It consists of a commentary against each clause of ISO 9004 guidelines for performance improvements explaining the practical benefits of implementing the guidance that is given in the standard **Sample Quality manual for** AAC blocks manufacturing, 2020-03-20 This book Sample Quality manual for AAC blocks manufacturing useful for all AAC Block manufacturing companies It has contents such as Raw material selection and testing for AAC blocks manufacturing Process description and troubleshooting over AAC blocks manufacturing problems All testings of AAC blocks Marketing user Modern Aspects of Pharmaceutical Quality Assurance Minal Ghante, Manohar Potdar, Vidhya guide with best policies Bhusari, 2024-03-11 The pharmaceutical quality system ensures that the process performance is suitably achieved the product quality is regularly met improved opportunities are identified and evaluated and the knowledge is constantly expanded Auditing also plays a crucial role within the pharmaceutical industry It helps to assess and review quality to improve and build a better system for the benefit of companies This book aims to develop a tool that will substantially decrease the number of Inspectional Observations and Warning letters thus eliminating Import Alerts and Consent Decree This book targets the Pharmaceutical Industry and students of Pharmaceutical Quality Assurance so they can get in hand ready consolidated information on Pharmaceutical Quality guidelines Quality metrics and implementation of simplified SOP guidelines plant layouts to implement Quality metrics for Pharmaceutical Manufacturing systems in tablets capsules liquid orals and semi solid dosage forms The chapters cover the various aspects of Pharmaceutical Quality Assurance The selection of topics is mainly based on the requirements of Pharmaceutical regulatory guidelines of India the UK the USA Australia and South Africa Each chapter includes the abstract detailed explanation implementation guidelines flowcharts layouts and Standard Operating Procedure of quality metrics for the Pharmaceutical Manufacturing System ISO 9001:2000 Quality Management System Design Jay J. Schlickman, 2003 Provides a set of design rules for creating a quality management system that will naturally translate into successful ISO 9001 2000 certification The book identifies the key documentation components and supplies guidelines for outlining and writing the guality manual standard operating procedures work instructions forms and records Two case studies illustrate the upgrade and recertification of a corporation from ISO 9001 1994 to ISO 9001 2000 and the creation of a company s first quality management system. The author is an auditor certified by the ASQ ANSI registrar accreditation board Annotation copyrighted by Book News Inc Portland OR **Ouality Assurance** and Quality Improvement Handbook for Human Research Leslie M. Howes, Sarah A. White, Barbara E. Bierer, 2019-11-05 Helping human research protection program professionals create implement and evaluate quality assurance quality improvement programs Quality Assurance and Quality Improvement Handbook for Human Research is the first comprehensively designed instructional manual aimed at teaching human research protection program HRPP professionals

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Implementing Quality in Laboratory Policies and Processes Donnell R. Christian Jr., Stephanie Drilling, 2009-11-24 In order to gain accreditation every laboratory must have a superior quality assurance program The keys to a successful program are the operational and technical manuals and associated documents which define the program and its various components Written by experts with global experience in setting up laboratories Implementing Quality in Laboratory Policies

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Manual of Food Quality Control Food and Agriculture Organization of the United Nations,1993-01-01 Water Quality Manual: Hydrologic and physical aspects of the environment Earl C. Shirley, Richard B. Howell, Kenneth D. Kerri,1976 Developing an ISO 13485-Certified Quality Management System Ilkka Juuso,2022-03-20 Developing an ISO 13485 Certified Quality Management System An Implementation Guide for the Medical Device Industry details the lessons learned from a real world project focusing on building an ISO 13485 2016 Quality Management System QMS from scratch and then

having it officially certified It is a practical guide to building or improving your existing QMS with tried and tested solutions The book takes a hands on approach first teaching the top 25 lessons to know before starting to develop a QMS and then walking you through the process of writing the quality manual and the standard operating procedures training the staff on the QMS organizing an internal audit executing a management review and finally passing the necessary external audits and obtaining certification It helps you to progress from one task to the next and provides all the essential information to accomplish each task as quickly and efficiently as possible It does not attempt to replicate the standard but instead drills into the standard to expose the core of each section of the standard and reorganize its contents into a practical workflow for developing maintaining and improving a Lean QMS The book includes a wealth of real world experience both from the author s personal dive into quality management and from the experiences of other companies in the field and provides handy checklists for ensuring key documents and processes are fit for use the emphasis here is to help ensure you have considered all relevant aspects In addition the book is not intended as a cheat sheet for the standard or as a review of the standard that only adds lengthy commentary on each of the clauses Instead the book fixes easy misunderstandings regarding QMS provides insight into why the various clauses are written the way they are and provides a great base to both understanding ISO 13485 QMS and developing your own QMS The book is intended to serve both experts and novices audiences it provides special insight on the most crucial and effective aspects of QMS Food Industry Quality Control Systems Mark Clute, 2008-10-22 With more than 100 tables spreadsheets and program outlines Food Industry Quality Control Systems features all of the systems needed for food manufacturing facilities to assure food quality and safety This book outlines the role of management in overall quality control and the role of HACCP within a company It explains the basics of quality control programs including pest control programs lot coding organizational charts specifications supplier certification and sanitation programs Readers can visit the CRC website to download supporting programs in a plug and play format and adaptable forms with specific examples and resources that enable companies to customize forms to fit their own needs **Ouality Procedures for** Hardware and Software D.J. Smith, J.S. Edge, 2012-12-06 How to Use This Book The primary purpose of this book is to assist small companies involved in both hardware and software to devise and evolve their own quality systems There are a number of national and now international standards which outline the activities for which procedures and records need to be specified They are described and compared in Chapter 2 and the subsequent guidance in the book is intended to assist in meeting them Although at first sight the operations of a hardware equipment developer may seem very different from those of a software house the basic requirements of a quality system such as the BS 5750 and ISO 1987 series of documents are the same For this reason the same standard can be called for in both areas and it will be seen in Part 2 that suitable procedures can be derived to meet both types of operation Quality standards BS 5750 AQAP ISO 9000 series distinguish between companies carrying out on the one hand both design and manufacturing fixed functions and on the other hand those

who only manufacture to specifications In practice the lesser requirements those applying to manufacture to fixed specifications are common to both levels of standard and the additional controls pertaining to design are added to obtain the higher standard Chapter 2 explains the differences in detail Malaria Microscopy Quality Assurance Manual World Health Organization, 2009 Early diagnosis and prompt effective treatment is the basis for the management of malaria and key to reducing malaria mortality and morbidity. An acceptable microscopy service is one that is cost effective provides results that are consistently accurate and timely enough to have a direct impact on treatment This requires a comprehensive and active quality assurance QA program This manual outlines a hierarchical structure based on retraining validation and the development of competency standards designed to ensure the quality of diagnosis necessary for a successful malaria program while remaining within the financial and personnel resources likely to be available. The mode of implementation of the QA system outlined in this manual will vary according to the organization of the national laboratory services dealing with malaria which may fall under the national malaria control program or under a separate laboratory structure working closely with the malaria program Procedures in Phlebotomy - E-Book John C. Flynn, 2023-08-12 Featuring concise easy to read coverage Procedures in Phlebotomy 5th Edition provides clear guidelines for performing both basic and special phlebotomy techniques It describes proper procedures for venipuncture special collection procedures and pediatric and geriatric considerations and addresses essential topics such as infection control OSHA guidelines and anatomy and physiology It also discusses professional issues such as interpersonal communication department management total quality and medical legal topics This edition includes enhanced infection control and prevention protocols updated healthcare law coverage expanded review questions and an updated practice test to prepare you for classroom and certification success Step by step illustrated procedures clearly outline essential phlebotomy competencies and techniques Essentials approach focuses on the need to know basics required to practice phlebotomy Case scenarios woven throughout the chapters help you continually apply content to practice Review questions in each chapter test your ability to recall information and apply critical thinking skills answers and rationales are included in the appendices Mock certification exam with 150 multiple choice questions and answers provides valuable practice for the credentialing test Safety Summary focuses on key points to ensure safe practice Competency checklists facilitate peer instructor and self evaluation on clinical procedures progress Key terms are listed at the beginning of each chapter highlighted in the text and defined in a back of book glossary NEW Protocols for enhanced infection control and prevention address COVID 19 and other emerging pathogens UPDATED Healthcare law coverage includes up to date content on the latest clinical guidelines EXPANDED and REVISED Review questions and a practice test help prepare you for classroom and certification success NEW COVID 19 content included throughout where relevant

Water safety plan manual World Health Organization, 2023-02-16 Understanding Quality Assurance in Construction H.W. Chung, 2002-01-04 The ISO 9000 family of quality standards has been adopted world wide as a framework for building

better relationships between suppliers and customers Originally a manufacturing industry concern quality is now acknowledged to be a key issue for the construction sector whose clients increasingly demand quality certification This book explains the concepts and practice of quality assurance and management in construction Clearly written and well illustrated with plenty of sample quality system documents and other pro forma this book will make the daunting task of developing implementing and managing a quality system a great deal easier for contractors This is practical guide for building and construction contractors and sub contractors project managers and other construction professionals Also for undergraduate and postgraduate students of building construction management and project management How to Establish a Document Control System for Compliance with ISO 9001:2015, ISO 13485:2016, and FDA Requirements Stephanie L. Skipper, 2015-10-14 This book explains the requirements for compliance with FDA regulations and ISO standards 9001 13485 for documented information controls and presents a methodology for compliance The document control system DCS or documented information control system DICS is the foundation of a quality management system It is the first quality system element that must be implemented because the establishment and control of documented processes and information in a quality controlled environment is dependent on the ability to proactively manage access to documents and the movement of documents through the document life cycle A well developed document control system benefits business by Improving knowledge retention and knowledge transfer within and across business units Improving access to knowledge based information Improving employee performance by providing standardized processes and communicating clear expectations Improving customer communication and satisfaction by providing documented information from which common understanding can be achieved Providing traceability of activities and documentation throughout the organization Improving organization of and access to documents and data Sample documents are included in the appendixes of this book to help clarify explanations and a full set of formatted procedures and document templates are available for download to get you off to an even faster start This book provides a process based approach that can be used for controlling all forms of documented information that are required to be managed under the quality management system

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