QMS ISO 9001 Document Control Plan Template – Example

QMS ISO 9001 Document Control Plan

Reviewed by	Approved by		
	Reviewed by		

1. Purpose

This section defines the purpose of the Document Control Plan, which is to establish a procedure for the monagement, control, and revision of all documents relevant to the Quality Monagement System QAMS to ensure that they remain current aggressed, and properly maintained. The purpose is also to comply with 50 9001 requirements for document portion.

2. Scope

This section outlines the types of documents covered by the Occument Control flow, including policies, procedures, work instructions, forms, monutes, and external documents relevant to the QAG. The scope defines which deportments or processes are subject to document control and the applicability of the control system across the argenization.

Beach

3. Definitions

- Ecoument: Any wither, electronic, or digital information used within the QMS.
- Revision: A formal change made to an existing document.
- Obsolete Document: A document that is no longer valid for use and has been replaced or dronved.
- Document Owner: The individual responsible for ensuring that a document is up-todate, accurate, and compliant

4. Responsibilities

Defines the key roles in the document control process:

- Document Dwner, Responsible for droffing, revising, and maintaining the document
- Document Controller, Ensures that documents are properly stored, distributed, and updated
- Approving Authority: The individual(s) responsible for approving the document before it is released for use
- Users: Personnel who must follow the procedures defined in controlled documents

5. Document Control Procedure

5.1 Document Creation and Identification

All documents must be clearly identified with a unique document number, title, version number, taked date, and depotiment of function to which they pertain, both new document or form must tolow as thuckured noming control for traceability and control.

5.2 Document Review and Revision

Prior to approval, all documents must undergo a thorough review proces. This includes vertifying the accuracy, compliateness, and relevance of the content, ensuring compliance with applicable standards or regulatory requirements. The document should also be revised periodically to remain current.

5.3 Document Approval

Documents must be formally approved by the relevant authority before being released for use. Each approval must be documented with the name of the approver(s), date of approval, and signature (digital or handwritten). No document may be issued without this approval.

5.4 Document Distribution

Once approved, documents must be distributed to relevant personner. Distribution methods may include electronic dissemination (e.g., through a document management system) or physical copies. The distribution list must be maintained, ensuring only the latest version of the document is in use.

5.5 Document Revision

inherever a change is made to a document, a new revision number is assigned. The revision history should dearly slate the reason for the revision, date of the change, and the approving outhority. As superseded versions must be archived to avoid confusion.

5.6 Document Storage and Accessibility

All controlled documents must be securely stored, whether in a digital or physical farmat. The document control system must ensure that documents are easily accessible to authorized personnel and protected from damage, loss, or unauthorized changes. Secure backup procedures should be in place for electronic documents.

5.7 Document Retention and Archiving

There must be a defined referring period for each document type based on its relevance, requisitory requirements, or company policy. After this period, documents are archived or disposed of securely. Archived documents must remain accessible for auditing or reference purposes.

5.8 Obsolete Documents

Obsolists or superseded discurrents must be clearly marked and removed from active use to prevent additional use. They should be anothived or disposed of in accordance with the retention policy. Obsolists documents may only be accessed for historical purposes or audits.

Document Control Register

6.1 Register Content

The Document Control Register is a log that tracks all controlled documents, including document number. We service mumber, date of issue, owner, next review date, and status formers obsolete). If provides a centralized, up-to-date overview of all documents under control.

4.2 Review Schedule

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4.3 - Revenior Status

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7. Security and Appens Control

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8. Training and Awareness

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9. Auditing and Compliance

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Padhraic Smyth

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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 ISO 9001:2008 for Small Businesses Ray Tricker, 2014-01-21 Small businesses face many challenges today including the increasing demand by larger companies for ISO compliance Compliance is a challenging task for any organisation and can often be time consuming and costly particularly for small businesses who are unlikely to have quality assurance experts on the payroll However it is still possible to achieve compliance without the need for expensive consultancy or training that takes you out of the office Ray Tricker has already guided hundreds of businesses through the challenge and this the 5th edition of his life saving ISO guide has been rewritten and refined following 5 years field use of working with the standard The one area that an organisation particularly a small business always wants to know is how much is it going to cost to implement and operate a QMS compliant with ISO 9001 2008 and is it going to be worth the trouble Due to popular demand Edition 5 now includes a brand new chapter on the cost of implementing ISO 9001 2008 This edition provides Relevant examples that put the concepts and requirements of the standard into a real life context Down to earth explanations to help you determine what you need to work in compliance with and or achieve certification to ISO 9001 2008 An example of a complete generic Quality Management System consisting of a Quality Manual plus a whole host of Quality Processes Quality Procedures and Work Instructions Access to a free software copy of this generic QMS files available from the author to give you a starting point from which to develop your own documentation ISO 9001 2008 is the most widely followed quality management standard and the rewards can be great opening up new business opportunities as well as bringing real improvements to your processes and outputs Guide to Quality Management Systems for the Food Industry Ralph Early, 2012-12-06 Whenever I step into an aeroplane I cannot avoid considering the risks associated with flying Thoughts of mechanical failure pilot error and ter rorist action fill my mind I try to reassure myself with statistics which tell me there is greater chance of injury crossing the road The moment the plane takes off I am resigned to my fate placing faith

in pilots who are highly qualified and superbly trained for the task of delivering me safely to my destination To be a passenger in an aeroplane is to express faith in the systems used by the airline It is to express a faith in the quality of the airline's organisation and the people who work within it The same is true of surgery Thoughts of mortality are difficult to avoid when facing the surgeon s knife However faith in the surgeon s training and skill faith in the anaesthetist and theatre technicians faith in the efficient resources and quality of the hospital all help to convince that there is little need to worry Apart from flying and surgery there are many facets of life which entail risk but knowing the risks we willingly place our confidence in others to deliver us safely In the consumption of food however few of us consider the risks Everyday if we are fortunate we eat food Food sustains and gives us pleasure Food supports our social interactions Essentials of Quality with Cases and Experiential Exercises Victor E. 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Clements, 1999 a management system **Tietz** Textbook of Clinical Chemistry and Molecular Diagnostics - E-Book Carl A. Burtis, Edward R. Ashwood, David E. Bruns, 2011-12-16 As the definitive reference for clinical chemistry Tietz Textbook of Clinical Chemistry and Molecular Diagnostics 5th Edition offers the most current and authoritative guidance on selecting performing and evaluating results of new and established laboratory tests Up to date encyclopedic coverage details everything you need to know including analytical criteria for the medical usefulness of laboratory procedures new approaches for establishing reference ranges variables that affect tests and results the impact of modern analytical tools on lab management and costs and applications of statistical methods In addition to updated content throughout this two color edition also features a new chapter on hemostasis and the latest advances in molecular diagnostics Section on Molecular Diagnostics and Genetics contains nine expanded chapters that focus on emerging issues and techniques written by experts in field including Y M Dennis Lo Rossa W K Chiu Carl Wittwer Noriko Kusukawa Cindy Vnencak Jones Thomas Williams Victor Weedn Malek Kamoun Howard Baum Angela Caliendo Aaron Bossler Gwendolyn McMillin and Kojo S J Elenitoba Johnson Highly respected author team includes three editors who are well known in the clinical chemistry world Reference values in the appendix give you one location for comparing and evaluating test results NEW Two color design throughout highlights important features illustrations and content for a quick reference NEW Chapter on hemostasis provides you with all the information you need to accurately conduct this type of clinical testing NEW Six associate editors lend even more expertise and insight to the reference NEW Reorganized chapters ensure that only the most current information is included ISO 9000 Quality Management System Design Jav J. Schlickman, 1998 Integrating ISO 14001 Into a Quality Management System Marilyn R. Block, I. Robert Marash, 2002 This book encourages critical thinking about the similarities between ISO 14001 and the ISO 9001 QS 9000 quality standards and shares approaches used by various organizations through examples of documentation that have withstood the scrutiny of registrars Understanding and Implementing ISO 9000:2000 David L. Goetsch, Stanley Davis, 2002 As organizations that compete in the global marketplace seek to achieve peak performance worldclass quality and personnel improvement interest in ISO 9000 certification has gained worldwide momentum This practical resource and guide uses a step by step model to understanding the ISO 9000 2000 series of standards and explains how to use them in an organization Provides a detailed summary of similarities and differences between the earlier version and ISO 9000 2000 Features ISO INFO boxes throughout that show how material being discussed applies to an actual work setting Offers a Serialized Case Study after every chapter that applies the information from a given chapter to how it was used in a real setting in which ISO 9000 registration was pursued the story unfolds with each new chapter showing the problems strategies mistakes and achievements of a company going through the registration process Ideal as a step by step how to guide for use in organizations that are interested in ISO 9000 registration The Certified Quality Manager Handbook Duke Okes, Russ Westcott, 2001 Designed as a study aid for those preparing to take the Certified Quality Manager Examination administered by the American Society for Quality ASQ this book provides a thorough understanding of the principles terms and concepts of quality management The new second edition contains practical examples from many different industries and organizations including manufacturing health care government education and the service industries Implementing ISO 9001:2000 Tom Taormina, Keith Brewer, 2002 ISO 9001 2000 A start to finish implementation guide to the newly updated ISO standard from two of the world's leading quality practitioners. The realist's guide to ISO 9001 2000 internal audits choosing registrars ensuring customer focus measuring results and much more Dell's Asset Recovery Business ARB A start to finish ISO 9001 case study from the insiders who made it happen Moving ISO 9001 from cost center to profit center To maintain their ISO 9001 certifications organizations worldwide must upgrade to the brand new ISO 9001 2000 standard In this book two of the world's leading quality practitioners present the most realistic business focused quide to ISO 9001 2000

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achieving registration is spent on fees for consultants to help explain the rules and prepare for assessment For the cover price of The Quality Systems Manual you could buy yourself about one hour of a consultant s time Practical Support for ISO 9001 Software Project Documentation Susan K. Land, John W. Walz, 2006-10-13 This book addresses how to meet the specific documentation requirements in support of the ISO 9001 software process definition documentation and improvement which is an integral part of every software engineering effort Provides a set of templates that support the documentation required for basic software project control and management The book provides specific support for organizations that are **Understanding and Implementing ISO 9000 and ISO Standards** pursuing software process improvement efforts David L. Goetsch, Stanley Davis, 1998 Appropriate for courses based on the ISO 9000 series of quality standards as a supplemental text in courses dealing with Quality Management and Total Quality Management TQM and as a hands on guide for use in private sector organizations This practical teaching resource how to guide provides a step by step model for understanding the ISO 9000 family of standards and implementing the ISO 9001 standard in a total quality environment Interest in ISO 9000 certification is gaining momentum worldwide as organizations that compete in the global marketplace seek to achieve consistent peak performance world class quality and continual improvement of their processes products and management systems

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