



أساسيات ISO 13485 للأجهزة الطبية



AGILE LEADERS
Training Center

29 Aug - 02 Sep 2027
الجبيل

أساسيات ISO 13485 للأجهزة الطبية

المرجع: 103600610_78172 التاريخ: 29 Aug - 02 Sep 2027 الموقع: الجبيل الرسوم: Euro 7500

Course Overview:

The ISO 13485 Foundation Training Course is designed to provide participants with a structured introduction to the Medical Devices Quality Management System MDQMS based on ISO 13485:2016 requirements. The course explains how organizations in the medical device sector establish and maintain a quality management system that ensures product safety, regulatory compliance, and consistent operational performance.

Participants will gain a clear understanding of the full medical device lifecycle, including design and development, production, distribution, installation, servicing, and post-market activities. The course introduces essential quality management system concepts such as documentation control, process-based management, risk-based thinking, internal audits, and continual improvement.

The training is aligned with internationally recognized certification pathways and prepares participants for the ISO 13485 foundation certification exam. It is suitable for professionals seeking to build foundational knowledge of regulated medical device environments and quality systems.

By the end of the course, participants will be able to understand how ISO 13485 supports regulatory compliance and improves product safety and quality in the global medical device industry.

Target Audience:

- Quality assurance and quality control professionals
- Regulatory affairs officers
- Medical device engineers and technicians
- Production and manufacturing staff
- Internal auditors and compliance officers
- Healthcare and laboratory professionals
- Entry-level quality managers
- Professionals entering the medical device industry

Targeted Organizational Departments:

- Quality assurance and quality control
- Regulatory affairs
- Research and development
- Manufacturing and production
- Supply chain and procurement
- Internal audit
- Engineering and product development
- Risk management

Targeted Industries:

- Medical device manufacturing
- Healthcare technology
- Pharmaceutical and biotechnology
- In vitro diagnostics industry
- Contract manufacturing organizations
- Hospital equipment manufacturing
- Laboratory equipment industry
- Healthcare service providers

Course Offerings:

By the end of this course, participants will be able to:

- Understand the structure and purpose of ISO 13485 in medical device quality systems
- Describe key principles of medical device quality management systems
- Identify ISO 13485 requirements and their application in organizations
- Understand documentation, process control, and compliance requirements
- Explain risk management principles in medical device operations
- Understand management responsibility and leadership roles in QMS
- Recognize internal audit and continual improvement processes
- Prepare for the ISO 13485 foundation exam
- Understand regulatory expectations in global medical device markets
- Apply ISO 13485 concepts in practical workplace scenarios

Training Methodology:

This training course is delivered using a structured and interactive approach that combines theoretical explanations with practical application. The learning experience includes instructor-led presentations supported by real-world examples from the medical device industry.

Participants will work through case studies that simulate actual quality management system challenges such as documentation control issues, supplier management problems, risk assessment scenarios, and non-conformance handling. These exercises help translate ISO 13485 requirements into practical workplace understanding.

Group discussions and guided activities are included to encourage engagement and reinforce key concepts. Participants are also given opportunities to analyze simplified quality system structures to understand how processes interact across an organization.

Knowledge checks, quizzes, and review questions are used throughout the course to strengthen understanding and prepare participants for the certification exam. Continuous trainer feedback ensures clarity and alignment with learning objectives.

Course Toolbox:

- ISO 13485 overview guide
- Medical device quality management system templates
- Risk management templates
- Document control and record keeping forms
- Internal audit checklists
- Case study workbook
- Sample quality manual and procedures structure
- Exam preparation questions
- Regulatory compliance checklist
- Glossary of medical device terms

Course Agenda:

Day 1: Introduction to Medical Device Quality Management Systems

- Introduction to ISO 13485 and medical device quality management systems Topic 1: •
- Evolution of ISO 13485:2016 and its regulatory context Topic 2: •
- Structure and scope of Medical Device Quality Management System MDQMS Topic 3: •
- Medical device lifecycle: design, production, distribution, and post-market Topic 4: •
- Regulatory environment for medical devices global overview Topic 5: •
- Key quality principles in regulated medical device industries Topic 6: •
- Introduction to risk-based thinking in medical device systems Topic 7: •
- Consolidation of MDQMS fundamentals and ISO 13485 core concepts Reflection & Review: •

Day 2: ISO 13485 Structure, Documentation, and Core Requirements

- ISO 13485 standard structure and clause breakdown Topic 1: •
- Quality management system documentation hierarchy Topic 2: •
- Document control and record management requirements Topic 3: •
- Quality manual development and system documentation Topic 4: •
- Management responsibility and leadership commitment Topic 5: •
- Resource management human resources, infrastructure, environment Topic 6: •
- Regulatory compliance and alignment with customer requirements Topic 7: •
- Understanding ISO 13485 system requirements and governance structure Reflection & Review: •

Day 3: Product Realization and Operational Processes

- Product realization lifecycle in medical device organizations Topic 1: •
- Planning of product realization and quality objectives Topic 2: •
- Design and development controls and verification activities Topic 3: •
- Purchasing and supplier evaluation processes Topic 4: •
- Production and service provision controls Topic 5: •
- Validation of processes and sterile product considerations Topic 6: •
- Monitoring and measurement equipment control Topic 7: •
- Application of operational controls in medical device environments Reflection & Review: •

Day 4: Measurement, Risk Management, and Continuous Improvement

- Measurement, analysis, and performance evaluation Topic 1: •
- Internal audit planning and execution Topic 2: •
- Control of nonconforming products and deviation handling Topic 3: •
- Corrective and preventive actions CAPA system Topic 4: •
- Risk management principles in ISO 13485 systems Topic 5: •
- Post-market surveillance and complaint handling Topic 6: •
- Continuous improvement and process optimization Topic 7: •
- Strengthening quality performance and improvement mechanisms Reflection & Review: •

Day 5: Certification Preparation and ISO 13485 Exam Readiness

- Overview of ISO 13485 Foundation certification exam structure Topic 1: •
- Competency domains and exam framework MDQMS principles Topic 2: •
- Key ISO 13485 clauses revision and consolidation Topic 3: •
- Sample exam questions and practice scenarios Topic 4: •
- Common mistakes and exam strategies Topic 5: •
- Certification pathways Foundation to Auditor and Implementer levels Topic 6: •
- Final revision and readiness assessment Topic 7: •
- Full course integration and exam preparedness evaluation Reflection & Review: •

FAQ:

What specific qualifications or prerequisites are needed for participants before enrolling in the course?

No formal prerequisites are required. However, basic knowledge of quality systems or exposure to healthcare or manufacturing environments is helpful.

How long is each day's session, and what is the total course duration?

Each session typically lasts 4-5 hours per day, including discussions and activities. The total course duration is five days.

What is the main challenge participants usually face in ISO 13485 Foundation training?

Most participants initially find it challenging to understand how regulatory requirements connect to practical quality system processes across the medical device lifecycle.

How This Course is Different from Other ISO 13485 Courses:

This training provides a clear and structured introduction to medical device quality management systems without requiring prior advanced technical knowledge. It focuses on building foundational understanding of ISO 13485 and how it is applied in real organizational environments.

Unlike advanced implementation or auditing courses, this program emphasizes clarity, progressive learning, and practical interpretation of requirements. It helps participants understand how a quality management system operates across departments and processes in the medical device industry.

The course also prepares participants for the ISO 13485 foundation certification exam by combining structured learning with practical examples and review exercises. This ensures participants not only understand the standard but are also able to apply it in real-world scenarios.

It is ideal for professionals entering regulated industries or seeking a strong foundation before progressing to advanced certification levels.

فئات الدورات التدريبية

الدورات التدريبية في مجال البيئة والاستدامة	دورات إدارة الجودة وتطوير العمليات
دورات إدارة و تحليل البيانات ودورات علم البيانات	دورات إدارة و تطوير الموارد البشرية
دورات الذهن السيرياني ودورات تقنية المعلومات	دورات الاتصال الجماهيري و السياسات والعلاقات العامة
دورات التدريب القانوني والهشتريات والتعاقدات	دورات التسويق وإدارة علاقات العملاء وإدارة المبيعات
دورات الحوكمة وإدارة المخاطر والامتثال	دورات السكرتارية و إدارة المكاتب
دورات الصحة والسلامة والذهن المهني	دورات الصيانة ودورات المجالات الهندسية المتنوعة
دورات المحاسبة و التهويل و دورات الإدارة المالية	دورات المهارات الشخصية وتطوير الذات
دورات في مجالات القيادة والإدارة	دورات معتمدة من قبل هيئات دولية
دورات مكتب إدارة المشاريع وإدارة المشاريع الرشيقية	



مدن التدريب

أهستردام هولندا	أكرا غانا	أثينا اليونان
الدار البيضاء المغرب	الجبيل المملكة العربية السعودية	استنبول تركيا
القاهرة مصر	الرياض المملكة العربية السعودية	الدوحة قطر
باريس فرنسا	الهناوة مملكة البحرين	الكويت الكويت
براغ جمهورية التشيك	بانكوك تايلند	باكو أذربيجان
بورتو البرتغال	برلين ألمانيا	برشلونة إسبانيا
تورنتو كندا	تيليسي جورجيا	بوكيت تايلاند
جوهانسبرغ جنوب افريقيا	جنيف سويسرا	جاكرتا جمهورية اندونيسيا
زنجار تنزانيا	روما إيطاليا	حلي الإمارات العربية المتحدة
سيول كوريا الجنوبية	سنغافورة سنغافورة	سان دييغو الولايات المتحدة الأمريكية
طرابزون تركيا	شيكاغو الولايات المتحدة الأمريكية	شرم الشيخ مصر
عمان المملكة الأردنية الهاشمية	طوكيو اليابان	طشقند أوزبكستان
فيينا النمسا	فرانكفورت ألمانيا	عن بعد منصة زووم

مدن التدريب

لانكاوي ماليزيا	كيب تاون جنوب افريقيا	كوالالمبور ماليزيا
هاربيا اسبانيا	لندن المملكة المتحدة	لشبونة البرتغال
هونتر سويسرا	مسقط سلطنة عمان	مدريد اسبانيا
نيروبي كينيا	هيونج المانيا	ميلان ايطاليا
		نيويورك الولايات المتحدة الأمريكية