



ISO 13485 specifies requirements for a quality management system where an organization needs to demonstrate its ability to provide Medical Devices and related services that consistently meet customer requirements and regulatory requirements applicable to medical devices and related services.

Organizations involved in medical device industry see ISO 13485 as the de facto standard towards regulatory compliance.

305-330-6337 | info@intiq.biz www.intiq.biz





ISO 13485 certification (also know as "registration"), is a third-party audit performed by a certification body such as MSECB who, upon verification that an organization is in compliance with the requirements of ISO 13485, will issue an ISO 13485 certificate. This certification is then maintained through regularly scheduled annual surveillance audits by the registrar, with re-certification of the Medical Devices Quality Management System performed on a triennial basis.

Benefits of implementing ISO 13485 Management System in your organization:

- Global Recognition
- Improved legal and regulatory or contractual requirements compliance
- ISO 13485 Medical Devices Quality Management System (QMS) is very close to the Food and Drug Administration's (FDA) QSR standards
- > Assistance in monitoring supply chain effectiveness
- Increased profit margins

- Improved product safety
- Increased Efficiency
- Proactive error detection and prevention
- Cost Savings
- More Effective Risk Management
- Increased likelihood of meeting Customer Requirements
- More Effective Risk Management
- Increased likelihood of meeting Customer Requirements

Benefits of ISO 13485 Certification to your customers:

- Quality ensured medical devices
- > Safe and effective medical devices
- Lower skepticism and increased confidence for endcustomer
- Boost of loyal customers and potential customers
- Increased profit margins



We help organizations to show commitment and competence with internationally recognized standards by providing this assurance through certification against rigorous, internationally recognized competence requirements. With a global coverage of over 150 countries worldwide, our mission is to provide our clients comprehensive services that inspire trust, continual improvement, demonstrate recognition, and benefit society as a whole.

To find out how you can obtain the ISO 13485 Certification, contact info@msecb.com

MSECB CERTIFICATION

YEAR 1 (INITIAL CERTIFICATION) -**AUDIT** INITIAL PRE-AUDIT **AUDIT PLAN** STAGE 1 AND 2 **CERTIFICATION PROCESS** (It is optional) Pre-audit must The audit plan has Non-conformities must be The Certificate will be issued be done at least 3 months closed at least 3 months after within 2 weeks after successful to be mutually before Certification Audit. agreed audit conclusions audit closing H YEAR 2 (1st SURVEILLANCE AUDIT) → YEAR 3 (2nd SURVEILLANCE AUDIT) SURVEILLANCE SURVEILLANCE **AUDIT PLAN AUDIT PLAN** AUDIT 1 **AUDIT 2** No longer than 12 No longer than 12 months months from the initial from the 1st surveillance certification audit audit

RE-CERTIFICATION AUDIT

Within two months before the triennial certificate expiration