

The book was found

ISO 14971:2000, Medical Devices -- Application Of Risk Management To Medical Devices



Synopsis

This International Standard specifies a procedure by which a manufacturer can identify the hazards associated with medical devices and their accessories, including in vitro diagnostic medical devices, estimate and evaluate the risks, control these risks and monitor the effectiveness of the control. The requirements of this International Standard are applicable to all stages of the life cycle of a medical device. This International Standard does not apply to clinical judgements relating to the use of a medical device. It does not specify acceptable risk levels. This International Standard does not require that the manufacturer has a formal quality system in place. However, risk management can be an integral part of a quality system (see, for example, Table G.1).

Book Information

Paperback: 40 pages

Publisher: Multiple. Distributed through American National Standards Institute (ANSI) (August 23, 2007)

Language: English

ASIN: B000XYTKUY

Product Dimensions: 8.2 x 0.1 x 10.5 inches

Shipping Weight: 4 ounces (View shipping rates and policies)

Average Customer Review: Be the first to review this item

Best Sellers Rank: #5,277,854 in Books (See Top 100 in Books) #83 in [Books > Engineering & Transportation > Engineering > Reference > American National Standards Institute \(ANSI\)](#)

Publications #1629 in [Books > Business & Money > Insurance > Risk Management](#) #35147 in [Books > Science & Math > Technology](#)

[Download to continue reading...](#)

ISO 14971:2000, Medical devices -- Application of risk management to medical devices ISO 14971:2007, Medical devices - Application of risk management to medical devices ISO/TS 22004:2005, Food safety management systems - Guidance on the application of ISO 22000:2005 ISO/IEC 31010:2009, Risk management - Risk assessment techniques ISO/IEC 20000-2:2012, Information technology - Service management - Part 2: Guidance on the application of service management systems Forensic Assessment of Violence Risk: A Guide for Risk Assessment and Risk Management ISO 13485:2016, Third Edition: Medical devices - Quality management systems - Requirements for regulatory purposes ISO 13485:2003, Medical devices - Quality management systems - Requirements for regulatory purposes ISO 12100:2010, Safety of machinery - General

principles for design - Risk assessment and risk reduction ISO/IEC 27002:2005, Information technology - Security techniques - Code of practice for information security management (Redesignation of ISO/IEC 17799:2005) Hedging Currency Exposures: Currency Risk Management (Risk Management Series) Making Enterprise Risk Management Pay Off: How Leading Companies Implement Risk Management Fundamentals of Risk Management: Understanding, evaluating and implementing effective risk management Security Risk Management: Building an Information Security Risk Management Program from the Ground Up ISO 31000:2009, Risk management - Principles and guidelines ISO Guide 73:2009, Risk management - Vocabulary ISO/IEC 27005:2011, Information technology - Security techniques - Information security risk management ISO 14155:2011, Clinical investigation of medical devices for human subjects - Good clinical practice ISO 11135:2014, Second Edition: Sterilization of health-care products - Ethylene oxide - Requirements for the development, validation and routine control of a sterilization process for medical devices ISO/TS 20022-3:2004, Financial services - UNiversal Financial Industry message scheme - Part 3: ISO 20022 modelling guidelines

[Contact Us](#)

[DMCA](#)

[Privacy](#)

[FAQ & Help](#)