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Medical Devices Seeram Ramakrishna, Lingling Tian, Charlene Wang, Susan Liao, Wee Eong Teo, 2015-08-18 Medical Devices and Regulations Standards and Practices will shed light on the importance of regulations and standards among all stakeholders bioengineering designers biomaterial scientists and researchers to enable development of future medical devices Based on the authors practical experience this book provides a concise practical guide on key issues and processes in developing new medical devices to meet international regulatory requirements and standards Provides readers with a global perspective on medical device regulations Concise and comprehensive information on how to design medical devices to ensure they meet regulations and standards Includes a useful case study demonstrating the design and approval process

Using Event-B for Critical Device Software Systems Neeraj Kumar Singh, 2013-06-19 Defining a new development life cycle methodology together with a set of associated techniques and tools to develop highly critical systems using formal techniques this book adopts a rigorous safety assessment approach explored via several layers from requirements analysis to automatic source code generation This is assessed and evaluated via a standard case study the cardiac pacemaker Additionally a formalisation of an Electrocardiogram ECG is used to identify anomalies in order to improve existing medical protocols This allows the key issue that formal methods are not currently integrated into established critical systems development processes to be discussed in a highly effective and informative way Using Event B for Critical Device Software Systems serves as a valuable resource for researchers and students of formal methods The assessment of critical systems development is applicable to all industries but engineers and physicians from the health domain will find the cardiac pacemaker case study of particular value The FDA and Worldwide Quality System Requirements Guidebook for Medical <u>Devices</u> Amiram Daniel, Ed Kimmelman, 2008-02-21 This new and expanded second edition maintains the organizational approach of the first and includes the requirements and guidance contained in the Quality System Regulation QSReg the ISO 13485 2003 standard the ISO TR 14969 2004 guidance document and as appropriate a number of the FDA and Global Harmonization Task Force GHTF guidance documents This second edition also addresses a number of additional topics such as the incorporation of risk management into the medical device organization s QMS QMS issues related to combination products the key process interactions within a QMS effective presentation of and advocacy for a QMS during FDA inspections and third party assessments and future FDA compliance and standards activities The organization of the guidebook is based on the order of the requirements in the QSReg For each substantive requirement section there is A verbatim statement of the QSReg requirement A description of the comparable requirement in ISO 13485 2003 focusing on any additions to or differences from the requirements contained in the QSReg Excerpts of the FDA responses to relevant comment groups contained in the Preamble to the QSReg Excerpts from various FDA guidance documents related to quality management systems A description of the relevant guidance contained in ISO TR 14969 2004 focusing on any additions to or differences

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