Design and Implementation of Medical Instrumentation

Gary H. Harding ECRI

WITHIN our society is a technical subculture that has brought about a significant increase in the quality, as well as complexity, of medical care. Society's demands for quality medical care, and the common belief that "more is better," have spurred the development of medical instrumentation. Industry, medical, legal, academic, and governmental sectors have, in either cooperation or conflict, been responsible for the development of sophisticated medical instrumentation, for example, imaging that can twist the body inside out, put it back together, and provide a high-resolution digital picture with patient information skillfully and artfully printed on it. On the other hand, in some cases a "calibrated eyeball" and a pair of tweezers are sufficient diagnostic and therapeutic tools to solve the patient's problem.

Medical personnel responsible for patient care and for the instrumentation necessary to provide that care are often technologically subverted. Nurses are spending more time with instrumentation and less time with the patient. The progression to "he can't be turning blue because the ventilator is working and the monitor is not alarming" is not off in the future—it's happened...and the patient is blue!

The engineer, particularly the biomedical engineer, is in a unique position to assist in the growth and safe use of medical instrumentation. If he becomes familiar with the mechanisms of instrument failure and success, and methodically reviews each new design with these in mind, the result will be safer, more effective, and more useful instrumentation.

PERFORMANCE PROBLEMS

The failure of instrumentation to perform its intended function can be the result of component failure, design flaws (electrical, mechanical, or software), interference (from patient, equipment, or environment), or operator error. Component failure

Current production and quality control test procedures among manufacturers are such that component failure leading to instrument malfunction is more common than it should be. As experience and automation make manufacturing and quality control test procedures more efficient, the incidence of component failure should decrease substantially.

Electrical, mechanical, and software design flaws that go undetected by the manufacturer, but are discovered by the user, are usually the result of poor design coordination, a deficiency in the structure of the manufacturer's prototype design and evaluation procedures, or a lack of understanding of clinical need and limits.

Often, too little thought is given to assessing the riskbenefit ratio of selected performance and control features. Improper instrument design almost ensures that, at some time, the device will be misused. An example is the inclusion of an unmarked "alarms off" position at the lower adjust-

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ment of an alarm limit control. A user not familiar with the subtleties of operating that device may think that such an adjustment will result in an alarm setting at a value just lower than the last marking. Even if the function of this control is clearly defined in the operator's manual, it is not inherently obvious to the user. Thus, if misapplication or misuse does occur, some responsibility rests with the designer.

The use of microprocessors has made the job of developing and implementing "glich-free" instrumentation that much more difficult. Most medical device manufacturers have limited experience in the quality control methods necessary to test computerized instrumentation. In addition, completely testing such systems (as is often done with seronautic or spacecraft instrumentation) is quite expensive; it can be difficult to decide which procedures are really necessary for inclusion in specific medical instrumentation quality control procedures.

The flowchart shown in Fig. 1 illustrates a model for the design, evaluation, and implementation of a medical instrument. This model should be used only as a guide, and changes (additions and deletions) to fit the actual need of the particular instrument under development should be made as needed.

It is imperative that personnel from engineering, clinical testing, and marketing specify the task definition. Many errors, which would otherwise be detected later in development or after the product is introduced into the field, can be discovered and dealt with, perhaps more cost-effectively, if users, designers, evaluators, and business and marketing personnel interact at the outset. Such interaction can reduce later costs (for example, modification or retrofitting), resulting in a lower product development expenditure by the manufacturer, a lower acquisition cost for the user, and an overall reduction in expenditures for health care. A classic mistake by manufacturers has been to rely heavily on a single consultant. If his advice fails to meet practical and business needs, the instrument may be both clinically and financially unsuccessful.

Although it is not uncommon for biomedical engineers to control prototype development because of their experience with engineering design and knowledge of physiological systems, it is a mistake to exclude others from this process. Critical decisions involving trade-offs and implementation are made during prototype development; an aggressive opendoor policy to promote discussion among engineers, clinicians (users), and consultants can result in a consensus and informed decisions. Periodic review of prototype status by the consulting and developing groups, including a discussion of issues and solutions, may eliminate what might otherwise become serious future problems.

Prototypes must be evaluated in steps, from least to greatest risk and from least to greatest difficulty within each step. It is unwise and unsafe to jump directly from prototype development to actual clinical trial. Safety, performance, and human factors characteristics must first be explored in laboratory and simulated environment evaluations. Such testing can be expensive, and may be abbreviated for simple devices or when substantially equivalent devices have already been tested. (Considerable discussion over whether two devices are "substantially equivalent" is often warranted.) Decisions to modify evaluation procedures should be made by the interdisciplinary group during task definition, so that such decisions can be legitimate and informed.

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List of Journals Indexed in Index Medicus National Library of Medicine (U.S.),1972 Issues for 1977 1979 include also Special List journals being indexed in cooperation with other institutions Citations from these journals appear in other MEDLARS bibliographies and in MEDLING but not in Index medicus Medical Instrumentation ,1979 Medical Devices and Human Engineering Joseph D. Bronzino,Donald R. Peterson,2018-10-08 Known as the bible of biomedical engineering The Biomedical Engineering Handbook Fourth Edition sets the standard against which all other references of this nature are measured As such it has served as a major resource for both skilled professionals and novices to biomedical engineering Medical Devices and Human Engineering the second volume of the handbook presents material from respected scientists with diverse backgrounds in biomedical sensors medical instrumentation and devices human performance engineering rehabilitation engineering and clinical engineering More than three dozen specific topics are examined including optical sensors implantable cardiac pacemakers electrosurgical devices blood glucose monitoring human computer interaction design orthopedic prosthetics clinical engineering program indicators and virtual instruments in health care The material is presented in a systematic manner and has been updated to reflect the latest applications and research findings

Managing Medical Devices within a Regulatory Framework Beth Ann Fiedler, 2016-09-10 Managing Medical Devices within a Regulatory Framework helps administrators designers manufacturers clinical engineers and biomedical support staff to navigate worldwide regulation carefully consider the parameters for medical equipment patient safety anticipate problems with equipment and efficiently manage medical device acquisition budgets throughout the total product life cycle This contributed book contains perspectives from industry professionals and academics providing a comprehensive look at health technology management HTM best practices for medical records management interoperability between and among devices outside of healthcare and the dynamics of implementation of new devices Various chapters advise on how to achieve patient confidentiality compliance for medical devices and their software discuss legal issues surrounding device use in the hospital environment of care the impact of device failures on patient safety methods to advance skillsets for HTM professionals and resources to assess digital technology. The authors bring forth relevant challenges and demonstrate how management can foster increased clinical and non clinical collaboration to enhance patient outcomes and the bottom line by translating the regulatory impact on operational requirements Covers compliance with FDA and CE regulations plus EU directives for service and maintenance of medical devices Provides operational and clinical practice recommendations in regard to regulatory changes for risk management Discusses best practices for equipment procurement and maintenance Provides guidance on dealing with the challenge of medical records management and compliance with patient confidentiality using information from medical devices Medical Equipment Maintenance Binseng Wang, 2012 Presents the basic elements of medical equipment maintenance and management required of healthcare leaders responsible for managing or

overseeing this function It will enable these individuals to understand their professional responsibilities as well as what they should expect from their supervised staff and how to measure and benchmark staff performance **Medical Devices and Systems** Joseph D. Bronzino, 2006-04-19 Over the last century medicine has come out of the black bag and emerged as one of the most dynamic and advanced fields of development in science and technology Today biomedical engineering plays a critical role in patient diagnosis care and rehabilitation More than ever biomedical engineers face the challenge of making Index of NLM Serial Titles National Library of Medicine (U.S.),1981 A keyword listing of serial titles currently received by the National Library of Medicine Selected References on Hospital Equipment United States. Health Care Facilities Service. Office of Equipment Consultation, 1973 Sterilisation of Biomaterials and Medical Devices Sophie Lerouge, Anne Simmons, 2012-09-27 The effective sterilisation of any material or device to be implanted in or used in close contact with the human body is essential for the elimination of harmful agents such as bacteria Sterilisation of biomaterials and medical devices reviews established and commonly used technologies alongside new and emerging processes Following an introduction to the key concepts and challenges involved in sterilisation the sterilisation of biomaterials and medical devices using steam and dry heat ionising radiation and ethylene oxide is reviewed A range of non traditional sterilisation techniques such as hydrogen peroxide gas plasma ozone and steam formaldehyde is then discussed together with research in sterilisation and decontamination of surfaces by plasma discharges Sterilisation techniques for polymers drug device products and tissue allografts are then reviewed together with antimicrobial coatings for self sterilisation and the challenge presented by prions and endotoxins in the sterilisation of reusable medical devices The book concludes with a discussion of future trends in the sterilisation of biomaterials and medical devices With its distinguished editors and expert team of international contributors Sterilisation of biomaterials and medical devices is an essential reference for all materials scientists engineers and researchers within the medical devices industry It also provides a thorough overview for academics and clinicians working in this area Reviews established and commonly used technologies alongside new and emerging processes Introduces and reviews the key concepts and challenges involved in sterilisation Discusses future trends in the sterilisation of biomaterials and medical devices The Design and Manufacture of Medical Devices J. Paulo Davim, J. Paulo Davim, 2012-10-16 Medical devices play an important role in the field of medical and health technology and encompass a wide range of health care products Directive 2007 47 EC defines a medical device as any instrument apparatus appliance software material or other article whether used alone or in combination including the software intended by its manufacturer to be used specifically for diagnostic and or the apeutic purposes and necessary for its proper application intended by the manufacturer to be used for human beings The design and manufacture of medical devices brings together a range of articles and case studies dealing with medical device R devices for specific applications such as spinal and craniofacial implants and other issues related to medical devices such as precision machining and integrated telemedicine systems Contains articles on

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sequenced to follow the complete lifecycle of open source medical technologies. The information provided is eminently practical as it is supported by real cases of study in which collaboration among medical professionals engineers and technicians patients and patient associations policy makers regulatory bodies and citizens has proven beneficial. The book is also supported by an online infrastructure UBORA through which open source medical devices can be collaboratively developed and shared for the democratization of medical technology and for promoting accessible biomedical engineering education. Current Catalog National Library of Medicine (U.S.),1979 First multi year cumulation covers six years 1965.

World Congress on Medical Physics and Biomedical Engineering 2018 Lenka Lhotska, Lucie Sukupova, Igor Lacković, Geoffrey S. Ibbott, 2018-05-29 This book vol 2 presents the proceedings of the IUPESM World Congress on Biomedical Engineering and Medical Physics a triennially organized joint meeting of medical physicists biomedical engineers and adjoining health care professionals Besides the purely scientific and technological topics the 2018 Congress will also focus on other aspects of professional involvement in health care such as education and training accreditation and certification health technology assessment and patient safety. The IUPESM meeting is an important forum for medical physicists and biomedical engineers in medicine and healthcare learn and share knowledge and discuss the latest research outcomes and technological advancements as well as new ideas in both medical physics and biomedical engineering field

Applications of Polyurethanes in Medical Devices Ajay Padsalgikar, 2022-05-19 Applications of Polyurethanes in Medical Devices provides detailed coverage of polyurethane PU chemistry processing and preparation for performant medical devices Polyurethanes have found many uses in medical applications due to their biocompatibility biostability physical properties surface polarity and the ability to suit the field of application This book enables the reader to understand polyurethane and how this valuable material can be used in medical devices Sections cover the chemistry structure and properties of polyurethane with in depth sections examining raw materials reaction chemistry synthesis techniques reaction kinetics material microstructure and structure property relationships Subsequent chapters demonstrate how polyurethane can be utilized in medical device applications examining biological properties rheology and processing before methodical coverage explains how polyurethane may be used for each category of medical device Finally future directions and safety and environmental aspects are covered Bridges the gap between polyurethane chemistry processing and preparation for cutting edge medical device applications Includes in depth coverage of polyurethane covering raw materials chemistry synthesis techniques reaction kinetics properties and microstructural analysis Takes a valuable and practical approach addressing manufacturing issues and using testing and modeling to solve problems encountered in processing **Medical Instrument Design and Development** Claudio Becchetti, Alessandro Neri, 2013-07-29 This book explains all of the stages involved in developing medical devices from concept to medical approval including system engineering bioinstrumentation design signal processing electronics software and ICT with Cloud and e Health development Medical Instrument Design and Development

offers a comprehensive theoretical background with extensive use of diagrams graphics and tables around 400 throughout the book The book explains how the theory is translated into industrial medical products using a market sold Electrocardiograph disclosed in its design by the Gamma Cardio Soft manufacturer The sequence of the chapters reflects the product development lifecycle Each chapter is focused on a specific University course and is divided into two sections theory and implementation The theory sections explain the main concepts and principles which remain valid across technological evolutions of medical instrumentation The Implementation sections show how the theory is translated into a medical product The Electrocardiograph ECG or EKG is used as an example as it is a suitable device to explore to fully understand medical instrumentation since it is sufficiently simple but encompasses all the main areas involved in developing medical electronic equipment Key Features Introduces a system level approach to product design Covers topics such as bioinstrumentation signal processing information theory electronics software firmware telemedicine e Health and medical device certification Explains how to use theory to implement a market product using ECG as an example Examines the design and applications of main medical instruments Details the additional know how required for product implementation business context system design project management intellectual property rights product life cycle etc Includes an accompanying website with the design of the certified ECG product www gammacardiosoft it book Discloses the details of a marketed ECG Product from Gamma Cardio Soft compliant with the ANSI standard AAMI EC 11 under open licenses GNU GPL Creative Common This book is written for biomedical engineering courses upper level undergraduate and graduate students and for engineers interested in medical instrumentation device design with a comprehensive and interdisciplinary system perspective

Healthcare and Medical Devices Jay Kalra and Nancy J. Lightner, 2022-07-24 Healthcare and Medical Devices
Proceedings of the 13th International Conference on Applied Human Factors and Ergonomics AHFE 2022 July 24 28 2022
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Congress on Medical Physics and Biomedical Engineering May 26-31, 2012, Beijing, China Mian Long, 2013-02-11
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Table of Contents Medical Instrumentation Journal

- 1. Understanding the eBook Medical Instrumentation Journal
 - The Rise of Digital Reading Medical Instrumentation Journal
 - Advantages of eBooks Over Traditional Books
- 2. Identifying Medical Instrumentation Journal
 - Exploring Different Genres
 - o Considering Fiction vs. Non-Fiction
 - Determining Your Reading Goals
- 3. Choosing the Right eBook Platform
 - Popular eBook Platforms
 - Features to Look for in an Medical Instrumentation Journal
 - User-Friendly Interface
- 4. Exploring eBook Recommendations from Medical Instrumentation Journal
 - Personalized Recommendations

- Medical Instrumentation Journal User Reviews and Ratings
- Medical Instrumentation Journal and Bestseller Lists
- 5. Accessing Medical Instrumentation Journal Free and Paid eBooks
 - Medical Instrumentation Journal Public Domain eBooks
 - Medical Instrumentation Journal eBook Subscription Services
 - Medical Instrumentation Journal Budget-Friendly Options
- 6. Navigating Medical Instrumentation Journal eBook Formats
 - ∘ ePub, PDF, MOBI, and More
 - Medical Instrumentation Journal Compatibility with Devices
 - Medical Instrumentation Journal Enhanced eBook Features
- 7. Enhancing Your Reading Experience
 - Adjustable Fonts and Text Sizes of Medical Instrumentation Journal
 - Highlighting and Note-Taking Medical Instrumentation Journal
 - Interactive Elements Medical Instrumentation Journal
- 8. Staying Engaged with Medical Instrumentation Journal
 - o Joining Online Reading Communities
 - Participating in Virtual Book Clubs
 - Following Authors and Publishers Medical Instrumentation Journal
- 9. Balancing eBooks and Physical Books Medical Instrumentation Journal
 - Benefits of a Digital Library
 - Creating a Diverse Reading Collection Medical Instrumentation Journal
- 10. Overcoming Reading Challenges
 - Dealing with Digital Eye Strain
 - Minimizing Distractions
 - Managing Screen Time
- 11. Cultivating a Reading Routine Medical Instrumentation Journal
 - Setting Reading Goals Medical Instrumentation Journal
 - Carving Out Dedicated Reading Time
- 12. Sourcing Reliable Information of Medical Instrumentation Journal
 - Fact-Checking eBook Content of Medical Instrumentation Journal

- Distinguishing Credible Sources
- 13. Promoting Lifelong Learning
 - Utilizing eBooks for Skill Development
 - Exploring Educational eBooks
- 14. Embracing eBook Trends
 - Integration of Multimedia Elements
 - Interactive and Gamified eBooks

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