Introduction to Medical Software

Providing a concise and accessible overview of the design, implementation, and management of medical software, this textbook will equip students with a solid understanding of critical considerations for both standalone medical software (software as a medical device/SaMD) and software that is integrated into hardware devices. It includes: practical discussion of key regulatory documents and industry standards, and how these translate into concrete considerations for medical software design; detailed coverage of the medical software lifecycle process; an accessible introduction to quality and risk management systems in the context of medical software; succinct coverage of essential topics in data science, machine learning, statistics, cybersecurity, software engineering, and healthcare, bringing readers up to speed; six cautionary real-world case studies which illustrate the dangers of improper or careless software processes. Accompanied by online resources for instructors, this is the ideal introduction for undergraduate students in biomedical engineering, electrical engineering and computer science, junior software engineers, and digital health entrepreneurs.

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Introduction to Medical Software
Foundations for Digital Health, Devices, and Diagnostics

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Contents

Preface xi
Acknowledgments xvii

Part I  Regulatory, Business, and Management Background

1 Introduction to Medical Software 3
Introduction 3
1.1 Medical Software and the Regulatory Process 3
1.2 The Medical Domain and Software 8
1.3 Emerging Challenges from the use of ML/AI 12
1.4 Conclusions 13
1.5 Summary 13
Recommended Reading and Resources 14

2 The FDA and Software 15
Introduction 15
2.1 The FDA and Medical Devices 15
2.2 A Review of Relevant FDA Documents 20
2.3 The FDA Regulatory Process 32
2.4 Regulatory Documents from Outside the United States 34
2.5 Implications of Regulatory Requirements for Software Design 35
2.6 The Future: Digital Health, AI, Self-Driving Cars? 37
2.7 Conclusions 39
2.8 Summary 40
Recommended Reading and Resources 40

3 Operating within a Healthcare System 44
Introduction 44
3.1 Software as Part of an Ecosystem 44
3.2 The Users: Patients, Doctors, and Everybody Else 46
3.3 Clinical Information Technology 49
3.4 Data Privacy 52
3.5 Cybersecurity 56
3.6 Conclusions 60
3.7 Summary 60
Recommended Reading and Resources 60
### 4 Quality Management Systems

**Introduction**

4.1 Introduction: The Organization Matters

4.2 Standards: ISO 9001, 90003, and 13485

4.3 The IMDRF QMS Document

4.4 Implications for Research and Scientific Software

4.5 General Remarks

4.6 Conclusions

4.7 Summary

**Recommended Reading and Resources**

### 5 Risk Management

**Introduction**

5.1 An Overview of Risk Management

5.2 Risk Analysis

5.3 Risk Evaluation

5.4 Risk Control

5.5 Risk Management and the Software Life Cycle

5.6 Conclusions

5.7 Summary

**Recommended Reading and Resources**

### 6 Taking an Idea to Market: Understanding the Product Journey

**Introduction**

6.1 Entrepreneurship: Should I Start a Company?

6.2 User-Centered Design: Consumer-Facing Medical Software Requires Finesse

6.3 Value Proposition: What Do You Bring to the Healthcare Conversation?

6.4 Intellectual Property

6.5 When to Raise Capital: Debt or Equity and What Does It All Mean?

6.6 Conclusions

6.7 Summary

**Recommended Reading and Resources**

### 7 Medical Software Applications and Growth Drivers

**Introduction**

7.1 Background and Definitions

7.2 The Technological Transformation of Healthcare

7.3 Healthcare Challenges

7.4 Promises of Healthcare Technology

7.5 Technology Growth Drivers

7.6 Healthcare Data

7.7 Technology in Clinical Trials

7.8 Regulatory Drivers
Contents

7.9 Industry Organizations 114
7.10 Remote Monitoring 116
7.11 Recommendations 117
7.12 Conclusions 117
7.13 Summary 118
Recommended Reading and Resources 118

Part II Scientific and Technical Background

8 Mathematical Background 123
Introduction 123
8.1 Data-Based Decision-Making 123
8.2 A Brief Introduction to Probability Theory 125
8.3 An Introduction to Statistics 131
8.4 Machine Learning and Its Complications 137
8.5 Are These Results Statistically Significant? 139
8.6 Importance of Randomization in Clinical Trials 141
8.7 Conclusions 143
8.8 Summary 143
Recommended Reading and Resources 144

9 Topics in Software Engineering 147
Introduction 147
9.1 Software Engineering Overview 147
9.2 Software Life Cycles 148
9.3 Software Engineering with AI/ML Modules 152
9.4 Modern Computing Environments 157
9.5 Software Testing 160
9.6 Source-Code Management 163
9.7 Conclusions 167
9.8 Summary 167
Recommended Reading and Resources 168

Part III An Example Medical Software Life Cycle Process

10 The Overall Process 175
Introduction 175
10.1 Overview of the Software Life Cycle Process 175
10.2 The IEC 62304 Standard 176
10.3 The Life Cycle Process 178
10.4 Software Safety Classifications 182
10.5 Other Issues 183
10.6 Conclusions 186
10.7 Summary 186
Recommended Reading and Resources 187
# Contents

## 11 Identifying User Needs 189

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>189</td>
</tr>
<tr>
<td>11.1 Exploring the User’s Needs</td>
<td>190</td>
</tr>
<tr>
<td>11.2 Creating a Use Specification Document</td>
<td>195</td>
</tr>
<tr>
<td>11.3 Example: The Image-Guided Neuro-navigation Project</td>
<td>196</td>
</tr>
<tr>
<td>11.4 Conclusions</td>
<td>197</td>
</tr>
<tr>
<td>11.5 Summary</td>
<td>198</td>
</tr>
</tbody>
</table>

Recommended Reading and Resources                                        198

## 12 The System Requirements Specification 201

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>201</td>
</tr>
<tr>
<td>12.1 A Bridge from User Needs to Software Design</td>
<td>201</td>
</tr>
<tr>
<td>12.2 Regulatory Background</td>
<td>202</td>
</tr>
<tr>
<td>12.3 An Example System Requirements Template</td>
<td>203</td>
</tr>
<tr>
<td>12.4 Writing a Requirement</td>
<td>207</td>
</tr>
<tr>
<td>12.5 Risk Management and Requirements</td>
<td>208</td>
</tr>
<tr>
<td>12.6 Reviewing System Requirements</td>
<td>209</td>
</tr>
<tr>
<td>12.7 Example: The Image-Guided Neuro-navigation Project</td>
<td>210</td>
</tr>
<tr>
<td>12.8 An Annotated Outline</td>
<td>213</td>
</tr>
<tr>
<td>12.9 Conclusions</td>
<td>216</td>
</tr>
<tr>
<td>12.10 Summary</td>
<td>216</td>
</tr>
</tbody>
</table>

Recommended Reading and Resources                                        216

## 13 The Software Design Document 218

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>218</td>
</tr>
<tr>
<td>13.1 A Bridge from Requirements to Code</td>
<td>218</td>
</tr>
<tr>
<td>13.2 Regulatory Background</td>
<td>218</td>
</tr>
<tr>
<td>13.3 Writing the Software Design Document</td>
<td>220</td>
</tr>
<tr>
<td>13.4 Risk Management</td>
<td>225</td>
</tr>
<tr>
<td>13.5 User Interface Evaluation</td>
<td>227</td>
</tr>
<tr>
<td>13.6 Example: The Image-Guided Neuro-navigation Project</td>
<td>228</td>
</tr>
<tr>
<td>13.7 Conclusions</td>
<td>230</td>
</tr>
<tr>
<td>13.8 Summary</td>
<td>230</td>
</tr>
</tbody>
</table>

Recommended Reading and Resources                                        231

## 14 Software Construction, Testing, and Verification 232

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>232</td>
</tr>
<tr>
<td>14.1 Two Concrete Steps of the Software Life Cycle</td>
<td>232</td>
</tr>
<tr>
<td>14.2 Software Construction</td>
<td>232</td>
</tr>
<tr>
<td>14.3 Software Testing and Verification</td>
<td>235</td>
</tr>
<tr>
<td>14.4 Creating a Testing and Verification Plan</td>
<td>237</td>
</tr>
<tr>
<td>14.5 Conclusions</td>
<td>238</td>
</tr>
<tr>
<td>14.6 Summary</td>
<td>239</td>
</tr>
</tbody>
</table>

Recommended Reading and Resources                                        239
CONTENTS

15 Software Validation 241
Introduction 241
  15.1 Regulatory Background 242
  15.2 Human Subject Studies and Clinical Trials 247
  15.3 Designing a Validation Strategy 249
  15.4 Conclusions 252
  15.5 Summary 252
Recommended Reading and Resources 253

16 Deployment, Maintenance, and Decommissioning 256
Introduction 256
  16.1 Deployment 256
  16.2 Maintenance 257
  16.3 Decommissioning 258
  16.4 Conclusions 259
  16.5 Summary 259
Recommended Reading and Resources 259

Part IV Case Studies

17 Therac-25: Software that Killed 263
Introduction 263
  17.1 Radiation Therapy 263
  17.2 The Therac Accidents 264
  17.3 Regulatory Reforms and Future Considerations 266

18 Mars Climate Orbiter: Lost without a Trace 268
Introduction 268
  18.1 Outer Space Exploration 268
  18.2 The Mars Orbiter Mission 269
  18.3 Fault Investigation 270
  18.4 Lessons Learned 273

19 HealthCare.gov: The Failed Launch of a Critical Website 274
Introduction 274
  19.1 Background 274
  19.2 The Launch of the Website 276
  19.3 The Post-Failure Investigation 277

20 The 2020 Iowa Caucus App: An Unreliable App that Caused National Embarrassment 280
Introduction 280
  20.1 The US Presidential Election Primary Process 280
  20.2 The 2020 Iowa Caucus 281
Contents

20.3 The Software Engineering Aspects 283
20.4 Conclusions 284

21 The Boeing 737 MAX Disasters: Using Software to Fix Hardware Problems 286
Introduction 286
21.1 Background 286
21.2 Disaster Strikes 289
21.3 Disaster Aftermath 290

22 The Averted Y2K Crisis: Successful Crisis and Risk Management 293
Introduction 293
22.1 The Y2K Problem 293
22.2 Preparations 294
22.3 The Aftermath 296

References 299
Index 316
Preface

The goal of this book is to provide in one brief and accessible volume a survey of the critical material involved in the design, implementation, and management of medical software for both standalone software (“software as a medical device – SaMD”) and software that is part of a physical medical device. One will find more detailed treatments of many of the topics covered in this book in specialized books that focus on some of the topics we cover (e.g. software engineering, systems engineering, probability theory, machine learning). Depth was not our goal; this book is explicitly designed to provide a broad survey.

Our hope is to familiarize the reader with the span of topics he or she may need in entering this field and to provide pointers to more specialized publications as this becomes necessary. For example, most computer scientists have very limited exposure to statistical decision theory, and we think that even the cursory coverage in this book will at least enable them to understand “what they do not know” and seek help as opposed to being ignorant of this entire field and attempting to reinvent the wheel in an amateurish manner!

An emerging challenge in medical software is the increasing use of big data and artificial intelligence/machine learning (AI/ML) techniques. This places an even greater stress on proper software design and management. Given that these are “black box” methods, in which the human understanding of what actually is going on is limited, a proper software quality process will be even more critical in creating reliable software tools. We introduce this topic in Section 1.3. In that section we also provide pointers to the other sections of the book in which we discuss issues related to the use of AI/ML methods.

This is an introductory book. One can and should follow the material here with further study, using both original regulatory materials, industry standards, and more advanced books. Our goal can be summarized by the phrase “to convert unknown unknowns to known unknowns.” Our goal is to make our reader aware of important material he or she is not as familiar with as one should be, and to pursue further study to acquire such knowledge.

This is not a programming book. Our goal is to describe the enabling activities that support programmers in producing high-quality software in the context of medical applications. We are less concerned by questions such as ‘How should we code?’ Our focus, rather, is on answering higher-level questions such as ‘How do we decide what we need to code?’ and ‘How should the process be organized?’ There is a wealth of material available that describes the actual coding process, and, therefore, we chose not to duplicate this type of description here.
This is also not a handbook for navigating the regulatory process. Our goal in writing this book was not to create a guide that one can follow to bring a product to market. We aim rather to explain how the regulatory process(es) affects the process of creating software for medical purposes. This will enable both a junior programmer entering this field and, perhaps, a high-level manager supervising a medical software project to understand why things are done the way they are. The regulatory documents and the associated industry standards capture much of what the medical device/software community has learned about this process over the past 20–30 years and constitute primary sources in this field.

Much of the content of this book is an expanded version of handouts prepared for the needs of the class Medical Software Design (BENG 406b) taught at Yale University over the period 2017–2021 by two of the authors (X. Papademetris and A. Quraishi). In our experience, university graduates are badly equipped by their coursework for careers in this general area, as many of the topics that are of critical importance to the medical device/medical software industry are almost never covered in undergraduate curricula.

The Structure of the Book
This book consists of four parts. The first two provide necessary background on the topic. They describe the environment that medical software lives in, be it scientific, regulatory, clinical, managerial, or financial. The third part describes the actual process of design, implementation, and testing of medical software, and the last part presents six case studies in which (for the most part) failure to follow the principles described earlier in the book led to expensive disasters and even deaths.

Part I: Regulatory, Business, and Management Background
This goal of this part of the book is to provide the necessary regulatory and business/management background for medical software. It is subdivided into the following chapters: Chapter 1: Introduction to Medical Software provides a brief overview of the entire field of medical software. Chapter 2: The FDA and Software focuses on the regulatory aspects of medical software, with a particular emphasis on the regulations and guidance issued by the US Food and Drug Administration (FDA). While this may initially appear to be a purely US-centric approach, it is worth pointing out that the most recent guidance documents from the FDA are based on the work of an international body, the International Medical Devices Regulator Forum (IMDRF), which represents an attempt at establishing international consensus in this field. Furthermore, Section 2.4 within this chapter discusses regulations from the EU and China and other countries.

Chapter 3: Operating within a Healthcare System has two goals. The first is to briefly describe the confusing and complex world of healthcare systems. The second goal is to introduce core clinical information technology standards and to discuss data privacy issues and related legislation, such as HIPAA (US) and GDPR (EU). The chapter concludes with a discussion of the critical topic of cybersecurity.
A quality management system is a core regulatory requirement for the development of safe and high-quality medical software (and any other product). This is described in ISO 9001 [131] and its more specialized derivatives (e.g. ISO 13485 [132]) for software and medical devices. We discuss this topic in Chapter 4: Quality Management Systems. Skill and technical competence alone do not suffice; one must perform the work in a properly managed organization whose culture is centered on quality. Chapter 5: Risk Management provides a description of this important topic, which is also a core regulatory requirement. Our bases here are the appropriate international standards, in particular ISO 14971 [133].

Medical software development happens in a business environment. Chapter 6: Taking an Idea to Market: Understanding the Product Journey is a guided tour of this area with a particular focus on startups and entrepreneurship. We cover here issues related to intellectual property, fundraising, and marketing. Even if the reader is not particularly interested in starting his or her own company, these are important issues to understand as they also drive management decisions in established companies.

We conclude this part with a more forward-looking chapter. Chapter 7: Medical Software Applications and Growth Drivers provides a review of the current state of the industry and, in particular, describes the excitement and emerging applications based on the use of AI/ML techniques in the area of digital health.

Part II: Scientific and Technical Background
Our aim in this part is to fill in common gaps in background knowledge of undergraduate students (and practitioners) in two important areas. 

Chapter 8: Mathematical Background is a fast-paced tour of probability, statistics, and machine learning. Anyone involved in the design of medical software (especially as this touches upon diagnosis and treatment) needs to have some basic background in topics such as detection theory and to understand terms such as false positive and false negative. Most computer scientists, in our experience, have minimal exposure to these topics.

Chapter 9: Topics in Software Engineering introduces basic material from software engineering for those of our readers coming from medical, engineering, and management backgrounds as opposed to computer science. Here, we discuss topics such as software life cycles, software testing, and source code management. All of these are critical in the design and implementation of medical software. We particularly highlight Section 9.3, where we provide a description of the challenges and techniques involved in the use of AI/ML modules within medical software.

Part III: An Example Medical Software Life Cycle Process
This practical application of software life cycles is the heart of this book. Here, we provide a simplified recipe for designing, implementing, testing, and distributing medical software to illustrate the concepts discussed in IEC 62304 [119]. The topic is introduced in Chapter 10: The Overall Process.
Preface

Next, Chapter 11: Identifying User Needs discusses the beginning of a project. This is where we identify what needs to be done and begin to plan how to do it. In this chapter, we also present a sample project, “The Image Guided Neuro-Navigation Project,” which we use to anchor our description of the various components of the process in the subsequent chapters.

Chapter 12: The System Requirements Specification discusses the critical process of creating the system requirement document, which is the master plan or anchor for the whole process. Next, Chapter 13: The Software Design Document discusses (for the first time) the actual, concrete process of designing software. Chapter 14: Software Construction, Testing, and Verification provides a brief description of software implementation and testing.

Chapter 15: Software Validation describes the validation process, which includes a brief description of clinical trials and issues related to the evaluation of AI/ML modules. This part of the book concludes with Chapter 16: Deployment, Maintenance, and Decommissioning, a short chapter that discusses these last steps in the software life cycle.

Part IV: Case Studies

This last part of the book consists of six case studies that relate to either software failures or the averting (in the case of the Y2K story) of crises that would have been caused by software issues that were successfully remedied in time. We use these to illustrate and reinforce some of the lessons presented earlier in the book.

Chapter 17: Therac-25 presents the famous case of the Therac-25. These accidents, which were caused by software bugs in a radiation treatment machine, resulted in serious injuries and deaths in a number of cases in the 1980s. They also resulted in the FDA taking an active interest in medical device software. Much of the regulation in this area derives directly or indirectly from the Therac-25 disasters.

Chapter 18: Mars Climate Orbiter describes the loss of the Mars Climate Orbiter. This is a classic failure of integration testing. Chapter 19: HealthCare.gov: The Failed Launch of a Critical Website describes the failed launch of the federal health insurance marketplace web page. During the creation of this web page, the planned software process was not followed, resulting in confusion and overruns. Chapter 20: The 2020 Iowa Caucus App presents the case of the 2020 Democratic Iowa Caucus App. This is both a case of management failure, effectively trying to do things “on the cheap” in a mission-critical setting, and a case of lack of awareness of the users and their environment.

Chapter 21: The Boeing 737 MAX Disasters describes the ongoing saga involving the design of this airplane. This case, which has some strong parallels to the Therac-25, is an example of an attempt to cut corners and use software to fix/rectify fundamental hardware problems, and using shortcuts and obfuscation to get around regulatory processes. Two 737 MAX airplanes crashed, resulting in hundreds of fatalities.

Our final vignette, Chapter 22: The Averted Y2K Crisis, is a success story. This illustrates how proper planning and thorough work averted what might have become a serious problem caused by legacy computer software design.
Target Audience

Undergraduate students in computer science and engineering: The initial stimulus for writing this book was to address the needs of our undergraduate students in the Yale class Medical Software Design (BENG 406b).

Junior software engineers in the medical device industry: Our hope is that Part I of this book will provide this group with an understanding of the environment they find themselves in and some justification as to why things are done the way they are. Part II may also be profitable to those who lack some of the technical background. Their company will have a specific process for the actual design phase, so Part III may be a little less applicable, but may also, again, provide some explanations as to why their company’s process is what it is. One of the reviewers of this book also suggested that knowledge of this material will be helpful in the interview process when applying for jobs in the medical device industry.

Senior management and/or policymakers supervising and/or managing software projects: Such individuals need to be able to assess the quality of the underlying software and to account for the risks that aspects of this software pose to their own decision-making process. For these individuals, the primary utility of a book such as this one (especially Parts I and II) may be to enable them to ask the right questions of their subordinates, who are actually directly managing the software process. The case studies (Part IV) may also provide sobering reading.

Those interested in starting a digital health company: This new entrepreneur in digital health is entering the minefield of the medical world, which comes with regulatory constraints (e.g. the FDA) and environmental constraints (hospitals) that make the process significantly harder (at least at the beginning). Our hope is that this person will be able to quickly skim this book and get a sense of what their task will involve and how to seek appropriate help (e.g. consultants) as they begin this challenging (and potentially highly rewarding) task of setting up a company (e.g. quality management systems – see Chapter 4). Chapter 6 may provide them with a useful road-map for the business aspects of this task, including coverage of topics such as intellectual property, fundraising, and setting up human studies/clinical trials. Chapter 7 reviews the current state of the industry and provides examples of recent work that may serve as an inspiration for new entrepreneurs. Finally, reading through the software process part of the book (Part III) will at the very least make it plain that they must document their software designs before they start coding if they would like anything they produce to meet regulatory standards.

NOTES

1. These regulatory documents [64, 69, 70, 82, 89, 91, 92, 125–127, 188] and industry standards [1, 2, 119, 122–124, 131–134] form a good initial list.
2. Probably at the top of this list should be the book *Medical Device Software Verification, Validation, and Compliance* [261] by D.A. Vogel. Though slightly dated, it contains much valuable information.

3. In teaching a class on medical software we have consistently emphasized that “figuring out what to code is much, much, harder than actually coding.” We, obviously, recognize that, in certain cases, the actual coding process can be highly complex and requires significant expertise and experience! There are many other classes that provide training on the coding process itself.

4. The regulatory process differs from country to country and it is critical to ascertain what the requirements are for any particular country in which approval is sought. The good news is that there are ongoing efforts to harmonize these regulations through the work of the IMDRF.

5. In Chapters 11–13 there is a specific section that applies the material in that chapter to our fictional image-guided navigation software project presented in Chapter 10. The hope is that this provides students with concrete examples of how to go about completing class assignments. We also provide sample class assignments in these chapters, and also in Chapter 15. One of the authors was involved in the design and implementation of a research interface for the BrainLAB system [198], but he had no (and still does not have a) financial connection to the company. This was a purely scientific collaboration as part of an NIH-funded project.

6. When taking a class on medical software, many students expect the class to start at this point. By contrast, many of the regulations highlight the importance of quality management systems and risk management in addition to the software life cycle process.
Acknowledgments

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Ellie Gabriel, a Yale biomedical engineering undergraduate, wrote the initial drafts of the six case studies in Part IV of this book as part of a summer internship (2020) she did under the supervision of Dr. Papademetris. She also created the illustration for Figure 10.4. She, together with Maria Papademetris, did much of the initial proofreading for the text.

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