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# **Cybersecurity in Medical Devices: Quality System Considerations and Content of Premarket Submissions Draft Guidance for Industry and Food and Drug Administration Staff**

***DRAFT GUIDANCE***

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**Document issued on April 8, 2022.**

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For questions about this document regarding CDRH-regulated devices, Suzanne Schwartz, Office of Strategic Partnerships and Technology Innovation at (301) 796-6937 or email [CyberMed@fda.hhs.gov](mailto:CyberMed@fda.hhs.gov). For questions about this document regarding CBER-regulated devices, contact the Office of Communication, Outreach, and Development (OCOD) at 1-800-835-4709 or 240-402-8010, or by email at [ocod@fda.hhs.gov](mailto:ocod@fda.hhs.gov).

**When final, this guidance will supersede Content of Premarket Submissions for Management of Cybersecurity in Medical Devices – Final Guidance, October 2, 2014**



U.S. Department of Health and Human Services  
Food and Drug Administration  
Center for Devices and Radiological Health  
Center for Biologics Evaluation and Research

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## **Preface**

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DRAFT

# Cybersecurity in Medical Devices: Quality System Considerations and Content of Premarket Submissions

## Draft Guidance for Industry and Food and Drug Administration Staff

*This draft guidance, when finalized, will represent the current thinking of the Food and Drug Administration (FDA or Agency) on this topic. It does not establish any rights for any person and is not binding on FDA or the public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations. To discuss an alternative approach, contact the FDA staff or Office responsible for this guidance as listed on the title page.*

### I. Introduction

With the increasing integration of wireless, Internet- and network- connected capabilities, portable media (e.g., USB or CD), and the frequent electronic exchange of medical device-related health information, the need for robust cybersecurity controls to ensure medical device safety and effectiveness has become more important.

In addition, cybersecurity threats to the healthcare sector have become more frequent and more severe, carrying increased potential for clinical impact. Cybersecurity incidents have rendered medical devices and hospital networks inoperable, disrupting the delivery of patient care across healthcare facilities in the U.S. and globally. Such cyber attacks and exploits may lead to patient harm as a result of clinical hazards, such as delay in diagnoses and/or treatment.

Increased connectivity has resulted in individual devices operating as single elements of larger medical device systems. These systems can include health care facility networks, other devices, and software update servers, among other interconnected components. Consequently, without adequate cybersecurity considerations across all aspects of these systems, a cybersecurity threat can compromise the safety and/or effectiveness of a device by compromising the functionality of any asset in the system. As a result, ensuring device safety and effectiveness includes adequate device cybersecurity, as well as its security as part of the larger system.

For the current edition of the FDA-recognized consensus standard(s) referenced in this document, see the [FDA Recognized Consensus Standards Database](https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/search.cfm).<sup>1</sup> For more information

<sup>1</sup> Available at <https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/search.cfm>.

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36 regarding use of consensus standards in regulatory submissions, please refer to the FDA  
37 guidance titled “[Appropriate Use of Voluntary Consensus Standards in Premarket Submissions](#)  
38 [for Medical Devices](#)”<sup>2</sup> and “[Standards Development and the Use of Standards in Regulatory](#)  
39 [Submissions Reviewed in the Center for Biologics Evaluation and Research](#).”<sup>3</sup>  
40

41 The contents of this document do not have the force of law and are not meant to bind the public  
42 in any way, unless specifically incorporated into a contract. This document is intended only to  
43 provide clarity to the public regarding existing requirements under the law. FDA’s guidance  
44 documents, including this draft guidance, should be viewed only as recommendations, unless  
45 specific regulatory or statutory requirements are cited. The use of the word *should* in Agency  
46 guidance means that something is suggested or recommended, but not required.

## 47 **II. Scope**

48 This guidance document is applicable to devices that contain software (including firmware)  
49 or programmable logic, as well as software as a medical device (SaMD). The guidance is  
50 not limited to devices that are network-enabled or contain other connected capabilities. This  
51 guidance describes recommendations regarding the cybersecurity information to be  
52 submitted for devices under the following premarket submission types<sup>4</sup>:  
53

- 54 • Premarket Notification (510(k)) submissions;
- 55 • De Novo requests;
- 56 • Premarket Approval Applications (PMAs) and PMA supplements;
- 57 • Product Development Protocols (PDPs);
- 58 • Investigational Device Exemption (IDE) submissions; and
- 59 • Humanitarian Device Exemption (HDE) submissions.
- 60

61 This guidance applies to all types of devices within the meaning of section 201(h) of the  
62 Federal Food, Drug, and Cosmetic Act (FD&C Act) whether or not they require a  
63 premarket submission. Therefore, the information in this guidance should also be  
64 considered for understanding FDA’s recommendations for devices for which a premarket  
65 submission is not required (e.g., for 510(k)-exempt devices).  
66

67 As IDE submissions have a different benefit-risk threshold and are not marketing authorizations,  
68 specific considerations for IDE submission documentation are provided in Appendix 3.  
69 Appendix 4 contains terminology used throughout the guidance.  
70

## 71 **III. Background**

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<sup>2</sup> Available at <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/appropriate-use-voluntary-consensus-standards-premarket-submissions-medical-devices>.

<sup>3</sup> Available at <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/standards-development-and-use-standards-regulatory-submissions-reviewed-center-biologics-evaluation>.

<sup>4</sup> Manufacturers should also consider applying the cybersecurity principles described in this guidance to the device constituent parts of other premarket submission types (e.g., Biologics License Applications (BLAs)) and to devices exempt from premarket review.

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72 FDA recognizes that medical device security is a shared responsibility among stakeholders  
73 throughout the use environment of the medical device system, including health care facilities,  
74 patients, health care providers, and manufacturers of medical devices. For the purposes of this  
75 guidance, the term “medical device system” includes the device and systems such as health care  
76 facility networks, other devices, and software update servers to which it is connected.

77  
78 Events across the healthcare sector have stressed the importance of cybersecurity to patient  
79 safety. The WannaCry<sup>5</sup> ransomware<sup>6</sup> affected hospital systems and medical devices across the  
80 globe. Vulnerabilities identified in commonly used third-party components, like URGENT/11<sup>7</sup>  
81 and SweynTooth<sup>8</sup>, have led to potential safety concerns across a broad range of devices and  
82 clinical specialties. In 2020, a ransomware attack on a German hospital highlighted the potential  
83 impacts due to delayed patient care when a cybersecurity attack forced patients to be diverted to  
84 another hospital<sup>9</sup>.

85  
86 The FDA issued a final cybersecurity guidance addressing premarket expectations in 2014  
87 “[Content of Premarket Submissions for Management of Cybersecurity in Medical Devices](#),” and  
88 the complementary guidance “[Postmarket Management of Cybersecurity in Medical Devices](#)”  
89 (“Postmarket Cybersecurity Guidance”)<sup>10</sup> in 2016. However, the rapidly evolving landscape, an  
90 increased understanding of emerging threats, and the need for capable deployment of mitigations  
91 throughout the total product lifecycle (TPLC) warrants an updated, iterative approach to device  
92 cybersecurity. The changes proposed since the 2014 guidance are intended to further emphasize  
93 the importance of ensuring that devices are designed securely, are designed to be capable of  
94 mitigating emerging cybersecurity risks throughout the TPLC, and to more clearly outline FDA’s  
95 recommendations for premarket submission information to address cybersecurity concerns.

96  
97 One way these TPLC considerations for devices can be achieved is through the implementation  
98 and adoption of a Secure Product Development Framework (SPDF). An SPDF is a set of  
99 processes that reduce the number and severity of vulnerabilities in products throughout the  
100 device lifecycle. Examples of such frameworks exist in many device sectors including the  
101 medical device sector. The recommendations contained in this guidance document, when  
102 finalized, are intended to supplement FDA’s “[Postmarket Management of Cybersecurity in](#)  
103 [Medical Devices](#),” “[Cybersecurity for Networked Medical Devices Containing Off-the-Shelf](#)

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<sup>5</sup> Additional information on the WannaCry Ransomware attack is available at: <https://h-isac.org/wannacry-ransomware-update/>.

<sup>6</sup> Ransomware is a type of malicious software, or malware, that infects a computer and restricts users’ access to it until a ransom is paid to unlock it.

<sup>7</sup> The FDA Safety Communication on the URGENT/11 vulnerabilities is available at: <https://www.fda.gov/medical-devices/safety-communications/urgent11-cybersecurity-vulnerabilities-widely-used-third-party-software-component-may-introduce>.

<sup>8</sup> The FDA Safety Communication on the SweynTooth vulnerabilities is available at: <https://www.fda.gov/medical-devices/safety-communications/sweyntooth-cybersecurity-vulnerabilities-may-affect-certain-medical-devices-fda-safety-communication>.

<sup>9</sup> Additional information on the German hospital ransomware attack is available at: <https://www.wired.co.uk/article/ransomware-hospital-death-germany>.

<sup>10</sup> See FDA’s guidance “[Postmarket Management of Cybersecurity in Medical Devices](#)” available at <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/postmarket-management-cybersecurity-medical-devices>.

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104 [\(OTS\) Software”<sup>11</sup>](#) and “[Guidance for the Content of Premarket Submissions for Software](#)  
105 [Contained in Medical Devices.”<sup>12</sup>](#) When finalized, this guidance will replace the final guidance  
106 “[Content of Premarket Submissions for Management of Cybersecurity in Medical Devices.”<sup>13</sup>](#)  
107

108 The recommendations in this guidance also generally align with or expand upon the  
109 recommendations in the Pre-Market Considerations for Medical Device Cybersecurity  
110 section of the International Medical Device Regulators Forum final guidance “Principles and  
111 Practices for Medical Device Cybersecurity,” issued March 2020.<sup>14</sup>

## 112 **IV. General Principles**

113 This section provides general principles for device cybersecurity relevant to device  
114 manufacturers. These principles, found throughout this guidance document, are important to  
115 the improvement of device cybersecurity and, when followed, are expected to have a positive  
116 impact on patient safety.

### 117 **A. Cybersecurity is Part of Device Safety and the Quality** 118 **System Regulations**

119 Device manufacturers must establish and follow quality systems to help ensure that their  
120 products consistently meet applicable requirements and specifications. These quality systems  
121 requirements are found in Quality System Regulation (QSR) in 21 CFR Part 820. Depending on  
122 the device, QS requirements may be relevant at the premarket stage, postmarket stage<sup>15</sup>, or both.  
123

124 In the premarket context, in order to demonstrate a reasonable assurance of safety and  
125 effectiveness for certain devices with cybersecurity risks, documentation outputs related to the  
126 requirements of the QSR may be one source of documentation to include as part of the premarket  
127

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<sup>11</sup> See FDA’s guidance “[Cybersecurity for Networked Medical Devices Containing Off-the-Shelf \(OTS\) Software](#)” available at <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/cybersecurity-networked-medical-devices-containing-shelf-ots-software>.

<sup>12</sup> See FDA’s guidance “[Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices](#)” available at <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/guidance-content-premarket-submissions-software-contained-medical-devices>.

<sup>13</sup> For the 2014 guidance on premarket submissions for management of cybersecurity, see FDA’s guidance “[Content of Premarket Submissions for Management of Cybersecurity in Medical Devices](#)” available at <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/content-premarket-submissions-management-cybersecurity-medical-devices-0>.

<sup>14</sup> See IMDRF Guidance “Principles and Practices for Medical Device Cybersecurity” available at <http://www.imdrf.org/docs/imdrf/final/technical/imdrf-tech-200318-pp-mdc-n60.pdf>.

<sup>15</sup> In the postmarket context, QSR design controls may also be important to ensure medical device cybersecurity and maintain medical device safety and effectiveness. FDA recommends that device manufacturers implement comprehensive cybersecurity risk management programs and documentation consistent with the QSR, including but not limited to complaint handling (21 CFR 820.198), quality audit (21 CFR 820.22), corrective and preventive action (21 CFR 820.100), software validation and risk analysis (21 CFR 820.30(g)) and servicing (21 CFR 820.200).



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128 submission<sup>16</sup> See also “[Guidance for the Content of Premarket Submissions for Software](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/guidance-content-premarket-submissions-software-contained-medical-devices)  
129 [Contained in Medical Devices](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/guidance-content-premarket-submissions-software-contained-medical-devices)” (available at [https://www.fda.gov/regulatory-information/search-](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/guidance-content-premarket-submissions-software-contained-medical-devices)  
130 [fda-guidance-documents/guidance-content-premarket-submissions-software-contained-medical-](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/guidance-content-premarket-submissions-software-contained-medical-devices)  
131 [devices](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/guidance-content-premarket-submissions-software-contained-medical-devices) ), hereafter “Premarket Software Guidance.” For example, 21 CFR 820.30(a) requires  
132 that for all classes of devices automated with software, a manufacturer must establish and  
133 maintain procedures to control the design of the device in order to ensure that specified design  
134 requirements are met (“QSR design controls”). As part of QSR design controls, a manufacturer  
135 must “establish and maintain procedures for validating the devices design,” which “shall include  
136 software validation and risk analysis, where appropriate.” 21 CFR 820.30(g). As part of the  
137 software validation and risk analysis required by 21 CFR 820.30(g), software device  
138 manufacturers may need to establish cybersecurity risk management and validation processes,  
139 where appropriate.

140 Software validation and risk analyses are key elements of cybersecurity analyses and  
141 demonstrating whether a connected device has a reasonable assurance of safety and  
142 effectiveness. FDA requires manufacturers to implement development processes that account  
143 for and address cybersecurity risks as part of design controls (21 CFR 820.30). For example,  
144 these processes should address the identification of security risks, the design requirements for  
145 how the risks will be controlled, and the evidence that the controls function as designed and  
146 are effective in their environment of use for ensuring adequate security.

#### **A Secure Product Development Framework (SPDF) may be one way to satisfy QSR requirements**

150 Cybersecurity threats have the potential to exploit one or more vulnerabilities that could lead  
151 to patient harm. The greater the number of vulnerabilities that exist and/or are identified over  
152 time in a system in which a device operates, the easier a threat can compromise the safety  
153 and effectiveness of the medical device. A Secure Product Development Framework (SPDF)  
154 is a set of processes that help reduce the number and severity of vulnerabilities in products.<sup>17</sup>  
155 An SPDF encompasses all aspects of a product’s lifecycle, including development, release,  
156 support, and decommission. Additionally, using SPDF processes during device design may  
157 prevent the need to re-engineer the device when connectivity-based features are added after  
158 marketing and distribution, or when vulnerabilities resulting in uncontrolled risks are  
159 discovered. An SPDF can be integrated with existing processes for product and software  
160 development, risk management, and the quality system at large.

161  
162 Using an SPDF is one approach to help ensure that QSR requirements are met. Because of its  
163 benefits in helping comply with QSRs and cybersecurity, FDA encourages manufacturers to  
164 use an SPDF, but other approaches might also satisfy QSR requirements.

---

<sup>16</sup> This guidance and its recommendations are not intended to suggest that FDA will evaluate an applicant’s compliance with the QSR as part of its premarket submission in our determination of a device’s substantial equivalence, as this is not a requirement for premarket submissions under section 513 of the FD&C Act. This guidance is intended to explain how FDA evaluates the performance of device cybersecurity and the cybersecurity outputs of activities that are part and parcel of QSR compliance, and explain how the QSR can be leveraged to demonstrate these performance outputs

<sup>17</sup> While the SPDF terminology has not been used in prior FDA guidance, the concepts around secure product development and risk management align with expectations in the Quality System and Labeling Regulations. As cybersecurity continues to evolve, FDA continues to align its terminology to reflect best practices.

165 **B. Designing for Security**

166 FDA will assess the adequacy of the device’s security based on the device’s ability to provide  
167 and implement the security objectives below throughout the system architecture.

168 **Security Objectives:**

- 169 • Authenticity, which includes integrity;
- 170 • Authorization;
- 171 • Availability;
- 172 • Confidentiality; and
- 173 • Secure and timely updatability and patchability.

174  
175 Premarket submissions should include information that describes how the above security  
176 objectives are addressed by and integrated into the device design. The extent to which security  
177 requirements, architecture, supply chain, and implementation are needed to meet these objectives  
178 will depend on:

- 179 • the device’s intended use and indications for use;
- 180 • the presence and functionality of its electronic data interfaces;
- 181 • its intended and actual environment of use;
- 182 • the type of cybersecurity vulnerabilities present;
- 183 • the exploitability of the vulnerabilities; and
- 184 • the risk of patient harm due to vulnerability exploitation.

185  
186  
187 SPDF processes aim to reduce the number and severity of vulnerabilities and thereby reduce the  
188 exploitability of a device and the associated risk of patient harm. Because exploitation of known  
189 vulnerabilities or weak cybersecurity controls should be considered reasonably foreseeable  
190 failure modes for systems, these factors should be addressed in the device design. The benefit of  
191 following an SPDF is that a device is more likely to be secure by design, such that the device is  
192 designed from the outset to be secure within its system and/or network of use.

193 **C. Transparency**

194 A lack of cybersecurity information, such as information necessary to integrate the device into  
195 the use environment, as well as information needed by users to maintain the device’s  
196 cybersecurity over the device lifecycle, has the potential to affect the safety and effectiveness of  
197 a device. In order to address these concerns, it is important for device users to have access to  
198 information pertaining to the device’s cybersecurity controls, potential risks, and other relevant  
199 information. For example:

- 200 • insufficient information pertaining to whether a device has undisclosed cybersecurity  
201 vulnerabilities or risks may be relevant to determining whether a device’s safety or  
202 effectiveness could be degraded;
- 203 • user manuals that do not include sufficient information to explain how to securely  
204 configure or update the device may limit the ability of end users to appropriately manage  
205 and protect the device; and/or

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- 207       • a failure to disclose all of the communication interfaces or third-party software could fail  
208       to convey potential sources of risks.

209  
210 This information and other relevant information is important in helping understand a device’s  
211 cybersecurity, the threats that it may be exposed to, and how those threats may be prevented or  
212 mitigated. Without it, cybersecurity risks could be undisclosed, inappropriately identified, or  
213 inappropriately responded to, among other potential impacts, which could lead to compromises  
214 in device safety and effectiveness.

215  
216 FDA believes that the cybersecurity information discussed in this guidance is important for the  
217 safe and effective use of interconnected devices and should be included in device labeling, as  
218 discussed below in Section VI.

## 219       **D. Submission Documentation**

220 Device cybersecurity design and documentation is expected to scale with the cybersecurity risk  
221 of that device. Manufacturers should take into account the larger system in which the device may  
222 be used. For example, a cybersecurity risk assessment performed on a simple, non-connected  
223 thermometer may conclude that the risks are limited, and therefore such a device needs only a  
224 limited security architecture (i.e., addressing only device hardware and software) and few  
225 security controls based on the technical characteristics and design of the device. However, if a  
226 thermometer is used in a safety-critical control loop, or is connected to networks or other  
227 devices, then the cybersecurity risks for the device are considered to be greater and more  
228 substantial design controls and documentation should be submitted in the premarket submission  
229 in order to demonstrate reasonable assurance of safety and effectiveness.

230  
231 Cybersecurity risks evolve over time and as a result, the effectiveness of cybersecurity controls  
232 may degrade as new risks, threats, and attack methods emerge. As cybersecurity is part of device  
233 safety and effectiveness, cybersecurity controls should take into consideration the intended and  
234 actual use environment (see section IV). In the 510(k) context, FDA evaluates the cybersecurity  
235 information submitted and the protections the cybersecurity controls provide in demonstrating  
236 substantial equivalence.<sup>18</sup> See section 513(i) of the FD&C Act and 21 CFR 807.100(b)(2)(ii)(B).

237  
238 In addition, inadequate cybersecurity controls may cause a device to be misbranded under  
239 section 502(f) of the FD&C Act because its labeling does not bear adequate directions for use or  
240 under section 502(j) of the FD&C Act because it is dangerous to health when used in the manner  
241 recommended or suggested in the labeling, among other possible violations.

242  
243 The cybersecurity information being recommended to be included in submissions as detailed in  
244 this guidance is based on risks due to cybersecurity, not on any other criteria or level of  
245 risk/concern established in a separate FDA guidance (e.g., the software risk criteria in the  
246 Premarket Software Guidance). For example, a device that is determined to have a greater  
247 software risk may only have a small cybersecurity risk due to how the device is designed.  
248 Likewise, a device with a smaller software risk may have a significant cybersecurity risk.

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<sup>18</sup> For more information, please refer to the guidance titled, “The 510(k) Program: Evaluating Substantial Equivalence in Premarket Notifications [510(k)]” regarding the substantial equivalence review standard.

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249 Therefore, the recommendations in this guidance regarding information to be submitted to the  
250 FDA are intended to address the cybersecurity risk, as assessed by the cybersecurity risk  
251 assessment, and are expected to scale based on the cybersecurity risk. The premarket submission  
252 documentation recommendations throughout this guidance apply to all premarket submissions  
253 and are intended to be used to support FDA’s assessment of a device’s safety and effectiveness.  
254

## 255 **V. Using an SPDF to Manage Cybersecurity Risks**

256 The documentation recommended in this guidance is based on FDA’s experience evaluating the  
257 safety and effectiveness of devices with cybersecurity vulnerabilities. However, sponsors may  
258 use alternative approaches and provide different documentation so long as their approach and  
259 documentation satisfies premarket submission requirements in applicable statutory provisions  
260 and regulations. The increasingly interconnected nature of medical devices has demonstrated the  
261 importance of addressing cybersecurity risks associated with device connectivity in device  
262 design because of the effects on safety and effectiveness.<sup>19</sup> Cybersecurity risks that are  
263 introduced by threats directly to the medical device or to the larger medical device system can be  
264 reasonably controlled through using an SPDF.  
265

266 The primary goal of using an SPDF is to manufacture and maintain safe and effective devices.  
267 From a security context, these are also trustworthy and resilient devices. These devices can then  
268 be managed (e.g., installed, configured, updated, review of device logs) through the device  
269 design and associated labeling by the device manufacturers and/or users (e.g., patients, health  
270 care facilities). For health care facilities, these devices may also be managed within their own  
271 cybersecurity risk management frameworks, such as the National Institute of Standards and  
272 Technology Framework for Improving Critical Infrastructure Cybersecurity, generally referred to  
273 as the NIST Cybersecurity Framework or NIST CSF.  
274

275 FDA recommends that manufacturers use device design processes such as those described in the  
276 QSR to support secure product development and maintenance. Other frameworks that satisfy the  
277 QSR and align with FDA’s recommendations for using an SPDF already exist and may be used,  
278 such as the medical device-specific framework that can be found in the Medical Device and  
279 Health IT Joint Security Plan (JSP).<sup>20</sup> Frameworks from other sectors may also comply with the  
280 QSR, like the framework provided in ANSI/ISA 62443-4-1: 2018 Security for industrial  
281 automation and control systems Part 4-1: Product security development life-cycle  
282 requirements.<sup>21</sup>  
283

284 The following subsections provide recommendations for using SPDF processes which FDA  
285 believes provide important considerations for the development of devices that are safe and  
286 effective, how these processes can complement the QSR, and the documentation FDA  
287 recommends manufacturers provide for review as part of premarket submissions. The

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<sup>19</sup> Addressing cybersecurity risks is in addition to addressing other risks, including software, biocompatibility, sterilization, and electromagnetic compatibility, among others.

<sup>20</sup> Medical Device and Health IT Joint Security Plan (JSP) is available at <https://healthsectorcouncil.org/the-joint-security-plan/>.

<sup>21</sup> ANSI/ISA-62443-4-1: 2018 *Security for industrial automation and control systems Part 4-1: Product security development life-cycle requirements* outlines a secure product development lifecycle similar to that of the JSP.

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288 information in these sections do not represent a complete SPDF. In addition, FDA does not  
289 recommend that manufacturers discontinue existing, effective processes.  
290

## 291 **A. Security Risk Management**

292  
293 To fully account for cybersecurity risks in devices, the safety and security risks of each device  
294 should be assessed within the context of the larger system in which the device operates. In the  
295 context of cybersecurity, security risk management processes are critical because, given the  
296 evolving nature of cybersecurity threats and risks, no device is, or can be, completely secure.  
297 Security risk management should be part of a manufacturer's quality system. Specifically, the  
298 QSR requires, among other things, that manufacturers' processes address design (21 CFR  
299 820.30), validation of the production processes (21 CFR 820.70), and corrective or preventive  
300 actions (21 CFR 820.100). These processes entail the technical, personnel, and management  
301 practices, among others, that manufacturers use to manage potential risks to their devices and  
302 ensure that their devices remain safe and effective, which includes security.  
303

304 The process for performing security risk management is a distinct process from performing  
305 safety risk management as described in ISO 14971:2019. This is due to the scope of possible  
306 harm and the risk assessment factors in the context of security may be different than those in  
307 the context of safety. Also, while safety risk management focuses on physical injury or  
308 damage to property or the environment, security risk management may include not only risks  
309 that can result in patient harm but also those risks that are outside of FDA's assessment of  
310 safety and effectiveness such as those related to business or reputational risks.  
311

312 Effective security risk management also addresses that cybersecurity-related failures do not  
313 occur in a probabilistic manner where an assessment for the likelihood of occurrence for a  
314 particular risk could be estimated based on historical data or modeling. This non-probabilistic  
315 approach is not the fundamental approach described in safety risk management under ISO  
316 14971:2019. Instead, security risk assessment processes focus on exploitability, or the ability  
317 to exploit vulnerabilities present within a device and/or system. Additional discussion on  
318 exploitability assessments for the security risk assessment can be found in the FDA's  
319 Postmarket Cybersecurity Guidance.<sup>22</sup> Exploitability for a cybersecurity risk during a  
320 premarket assessment may be different compared to a risk assessment performed for a  
321 postmarket vulnerability. For example, some of the exploitability factors discussed in the  
322 guidance (e.g., Exploit Code Maturity, Remediation Level, Report Confidence<sup>23</sup>) may not be  
323 applicable to unreleased software. In these instances, a premarket exploitability assessment  
324 could either assume a worst-case assessment and implement appropriate controls, or provide a  
325 justification for a reasonable exploitability assessment of the risk throughout the total product  
326 lifecycle and how the risk is controlled.  
327

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<sup>22</sup> See Footnote 10.

<sup>23</sup> These factors of exploitability are from the Common Vulnerability Scoring System (CVSS) Version 3.0 as identified in the Postmarket Cybersecurity Guidance. Additional information on CVSS is available at <https://www.first.org/cvss/>.

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328 FDA recommends that manufacturers establish a security risk management process that  
329 encompasses design controls (21 CFR 820.30), validation of production processes (21 CFR  
330 820.70), and corrective and preventive actions (21 CFR 820.100) to ensure both safety and  
331 security risks are adequately addressed. For completeness in performing risk analyses under 21  
332 CFR 820.30(g), FDA recommends that device manufacturers conduct both a safety risk  
333 assessment per ISO 14971:2019 and a separate, accompanying security risk assessment to  
334 ensure a more comprehensive identification and management of patient safety risks. The scope  
335 and objective of a security risk management process, in conjunction with other SPDF processes  
336 (e.g., security testing), is to expose how threats, through vulnerabilities, can manifest patient  
337 harm and other potential risks. These processes should also ensure that risk control measures  
338 for one type of risk assessment do not inadvertently introduce new risks in the other. AAMI  
339 TIR57:2016 details how the security and safety risk management processes should interface to  
340 ensure all risks are adequately assessed.<sup>24</sup>

341  
342 Known vulnerabilities should be mitigated in the design of the device. For marketed devices, if  
343 comprehensive design mitigations are not possible, compensating controls should be  
344 considered. All devices, when any known vulnerabilities are only partially mitigated or  
345 unmitigated by the device design, they should be assessed as reasonably foreseeable risks in  
346 the risk assessment and be assessed for additional control measures or risk transfer to the  
347 user/operator, or, if necessary, the patient. Risk transfer, if appropriate, should only occur when  
348 all relevant risk information is known, assessed, and appropriately communicated to users and  
349 includes risks inherited from the supply chain as well as how risk transfer will be handled  
350 when the device/system reaches end of support and end of life and whether or how the user is  
351 able to take on that role (e.g., if the user may be a patient).

352  
353 Specific security risk management documentation where FDA has recommendations regarding  
354 their scope and/or content are discussed in the subsections below. The documentation FDA  
355 recommends manufacturers provide in their premarket submissions is summarized in the  
356 Security Risk Management Documentation below (Section V.A.4.).

#### **1. Threat Modeling**

357  
358  
359 Threat modeling includes a process for identifying security objectives, risks, and  
360 vulnerabilities across the system, and then defining countermeasures to prevent, or mitigate the  
361 effects of, threats to the system throughout its lifecycle. It is foundational for optimizing  
362 system, product, network, application, and connection security when applied appropriately and  
363 comprehensively.

364  
365 With respect to security risk management, and in order to identify appropriate security risks  
366 and controls for the system, FDA recommends that threat modeling be performed to inform  
367 and support the risk analysis activities. As part of the risk assessment, FDA recommends threat  
368 modeling be performed throughout the design process and be inclusive of all system elements.

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<sup>24</sup> AAMI TIR57:2016 Principles for medical device security—Risk management describes the security risk management process and how the security risk management process should have links into the safety risk management process and vice versa.



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The threat model should:

- identify system risks and mitigations as well as inform the pre- and post-mitigation risks considered as part of the security risk assessment;
- state any assumptions about the system or environment of use (e.g. hospital networks are inherently hostile, therefore manufacturers are recommended to assume that an adversary controls the network with the ability to alter, drop, and replay packets); and
- capture cybersecurity risks introduced through the supply chain, manufacturing, deployment, interoperability with other devices, maintenance/update activities, and decommission activities that might otherwise be overlooked in a traditional safety risk assessment processes.

FDA recommends that premarket submissions include threat modeling documentation to demonstrate how the risks assessed and controls implemented for the system address questions of safety and effectiveness. There are a number of methodologies and/or combinations of methods for threat modeling that manufacturers may choose to use. Rationale for the methodology(ies) selected should be provided with the threat modeling documentation. Additional recommendations on how threat modeling documentation should be submitted to FDA are discussed in Section V.B. below.

Threat modeling activities can be performed and/or reviewed during design reviews. FDA recommends that threat modeling documentation include sufficient information on threat modeling activities performed by the manufacturer to assess and review the security features built into the device such that they holistically evaluate the device and the system in which the device operates, for the safety and effectiveness of the system.

## **2. Third-Party Software Components**

As discussed in the FDA guidances “[Off-The-Shelf \(OTS\) Software Use in Medical Devices](#)”<sup>25</sup> and “[Cybersecurity for Networked Medical Devices Containing Off-the-Shelf \(OTS\) Software](#),”<sup>26</sup> medical devices commonly include third-party software components including off-the-shelf and open source software. When these components are incorporated, security risks of the software components become factors of the overall medical device system risk management processes and documentation.

As part of demonstrating compliance with quality system design controls under 21 CFR 820.30(g), and to support supply chain risk management processes, all software, including that developed by the device manufacturer (“proprietary software”) and obtained from third parties should be assessed for cybersecurity risk and that risk should be addressed. Accordingly, device

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<sup>25</sup> See FDA guidance [Off-The-Shelf \(OTS\) Software Use in Medical Devices](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/shelf-software-use-medical-devices) available at: <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/shelf-software-use-medical-devices>.

<sup>26</sup> See FDA guidance [Cybersecurity for Networked Medical Devices Containing Off-the-Shelf \(OTS\) Software](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/cybersecurity-networked-medical-devices-containing-shelf-ots-software) available at: <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/cybersecurity-networked-medical-devices-containing-shelf-ots-software>.

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407 manufacturers are expected to document all software components<sup>27</sup> of a device and to mitigate  
408 risks associated with these software components.

409  
410 In addition, under 21 CFR 820.50, manufacturers must put in place processes and controls to  
411 ensure that their suppliers conform to the manufacturer’s requirements. Such information is  
412 documented in the Design History File, required by 21 CFR 820.30(j), and Design Master  
413 Record, required by 21 CFR 820.181. This documentation demonstrates the device’s overall  
414 compliance with the QSR, as well as that the third-party components meet specifications  
415 established for the device. Security risk assessments that include analyses and considerations of  
416 cybersecurity risks that may exist in or be introduced by third-party software and the software  
417 supply chain may help demonstrate that manufacturers have adequately ensured such  
418 compliance and documented such history.

419  
420 As part of configuration management, device manufacturers should have custodial control of  
421 source code through source code escrow and source code backups.<sup>28</sup> While source code is not  
422 provided in premarket submissions, if this control is not available based on the terms in supplier  
423 agreements, the manufacturer should include in premarket submissions a plan of how the third-  
424 party software component could be updated or replaced should support for the software end. The  
425 device manufacturer is also expected to provide to users whatever information is necessary to  
426 allow users to manage risks associated with the device.

427  
428 One tool to help manage supply chain risk as well as clearly identify and track the software  
429 incorporated into a device is a Software Bill of Materials (SBOM), as described below.

#### **(a) Software Bill of Materials**

430  
431  
432  
433 A Software Bill of Materials (SBOM) can aid in the management of cybersecurity risks that exist  
434 throughout the software stack. A robust SBOM includes both the device manufacturer-  
435 developed components and third-party components (including purchased/licensed software and  
436 open-source software), and the upstream software dependencies that are required/depended upon  
437 by proprietary, purchased/licensed, and open-source software. An SBOM helps facilitate risk  
438 management processes by providing a mechanism to identify devices that might be affected by  
439 vulnerabilities in the software components, both during development (when software is being  
440 chosen as a component) and after it has been placed into the market throughout all other phases  
441 of a product’s life.<sup>29</sup>

442  
443 Because vulnerability management is a critical part of a device’s security risk management  
444 processes, an SBOM or an equivalent capability should be maintained as part of the device’s  
445 configuration management, be regularly updated to reflect any changes to the software in

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<sup>27</sup> The use of “component” in this guidance is consistent with the definition in 21 CFR 820.3.

<sup>28</sup> While some suppliers may not grant access to source code, manufacturers may consider adding to their purchasing controls acquisition of the source code should the purchased software reach end of support or end of life from the supplier earlier than the intended end of support or end of life of the medical device.

<sup>29</sup> For additional information see the Department of Commerce National Telecommunications and Information Administration’s multi-stakeholder process for software transparency.

<https://www.ntia.doc.gov/SoftwareTransparency>



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446 marketed devices, and should support 21 CFR 820.30(j) (Design History File) and 820.181  
447 (Design Master Record) documentation.

448  
449 To assist FDA’s assessment of the device risks and associated impacts on safety and  
450 effectiveness related to cybersecurity, FDA recommends that premarket submissions include  
451 SBOM documentation as outlined below. SBOMs can also be an important tool for transparency  
452 with users of potential risks as part of labeling as addressed later in Section VI

#### 453 454 **(b) Documentation Supporting Software Bill of Materials**

455  
456 FDA’s guidance documents “[Off-The-Shelf \(OTS\) Software Use in Medical Devices](#)”<sup>30</sup> and  
457 “[Cybersecurity for Networked Medical Devices Containing Off-the-Shelf \(OTS\) Software](#)”<sup>31</sup>  
458 describe information that should be provided in premarket submissions for software components  
459 for which a manufacturer cannot claim complete control of the software lifecycle. In addition to  
460 the information recommended in those guidances, for each OTS component, the following  
461 should also be provided in a machine-readable format in premarket submissions.

- 462  
463 A. The asset(s) where the software component resides;  
464 B. The software component name;  
465 C. The software component version;  
466 D. The software component manufacturer;  
467 E. The software level of support provided through monitoring and maintenance from  
468 the software component manufacturer;  
469 F. The software component’s end-of-support date; and  
470 G. Any known vulnerabilities.<sup>32</sup>

471  
472 Industry-accepted formats of SBOMs can be used to provide this information to FDA; however,  
473 if any of the above elements are not captured in such an SBOM, we recommend that those items  
474 also be provided, typically as an addendum, to FDA for the purposes of supporting premarket  
475 submission review. Additional examples of the type of information to include in a SBOM can be  
476 found in the Joint Security Plan - Appendix G (“Example Customer Security Documentation”)<sup>33</sup>  
477 and Sections 2.3.17 and 2.3.18 of the Manufacturer Disclosure Statement for Medical Device  
478 Security (referred to as MDS2 or MDS<sup>2</sup>)<sup>34</sup>.

479  
480 As part of the premarket submission, manufacturers should also describe how the known  
481 vulnerabilities (item (G) above) were discovered to demonstrate whether the assessment methods

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<sup>30</sup> See FDA guidance Off-The-Shelf (OTS) Software Use in Medical Devices available at:

<https://www.fda.gov/regulatory-information/search-fda-guidance-documents/shelf-software-use-medical-devices>.

<sup>31</sup> See FDA guidance Cybersecurity for Networked Medical Devices Containing Off-the-Shelf (OTS) Software available at: <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/cybersecurity-networked-medical-devices-containing-shelf-ots-software>.

<sup>32</sup> Known vulnerabilities are vulnerabilities that are published in the public National Vulnerability Database (NVD) or similar software vulnerability and/or weakness database. NVD is available at <https://nvd.nist.gov/vuln/full-listing>

<sup>33</sup> Medical Device and Health IT Joint Security Plan (JSP) is available at <https://healthsectorcouncil.org/the-joint-security-plan/>.

<sup>34</sup> The Manufacturer Disclosure Statement for Medical Device Security is available at <https://www.nema.org/standards/view/manufacturer-disclosure-statement-for-medical-device-security>.

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482 were sufficiently robust. For third-party components with known vulnerabilities, device  
483 manufacturers should provide in premarket submissions:

- 484
- 485 • A safety and security risk assessment of each known vulnerability; and
- 486 • Details of applicable safety and security risk controls to address the vulnerability. If risk  
487 controls include compensating controls, those should be described in an appropriate level  
488 of detail

489

490 For additional information and discussion regarding proprietary and third-party components, see  
491 section V.B.2., Security Architecture Views, below.

### 492 **3. Security Assessment of Unresolved Anomalies**

493

494 FDA's [Premarket Software Guidance](#), recommends that device manufacturers provide a list of  
495 software anomalies (e.g., bugs or defects) that exist in a product at the time of submission. For  
496 each of these anomalies, FDA recommends that device manufacturers conduct an assessment of  
497 the anomaly's impact on safety and effectiveness, and consult the Premarket Software Guidance  
498 to assess the associated documentation recommended for inclusion in such device's premarket  
499 submission.

500

501 Some anomalies discovered during development or testing may have security implications and  
502 may also be considered vulnerabilities. As a part of ensuring a complete security risk assessment  
503 under 21 CFR Part 820.30(g), the assessment for impacts to safety and effectiveness may include  
504 an assessment for the potential security impacts of anomalies. The assessment should also  
505 include consideration of any present Common Weakness Enumeration (CWE) categories.<sup>35</sup>  
506 For example, a clinical user may inadvertently reveal the presence of a previously unknown  
507 software anomaly during normal use, where the impact of the anomaly might occur sporadically  
508 and be assessed to be acceptable from a software risk perspective. Conversely, a threat might  
509 seek out these types of anomalies, and identify means to exploit them in order to manifest the  
510 anomaly's impact continuously, which could significantly impact the acceptability of the risk  
511 when compared to an anomaly assessment that didn't include security considerations.

512

513 The criteria and rationales for addressing the resulting anomalies with security impacts should be  
514 provided as part of the security risk assessment documentation in the premarket submission.

### 515 **4. Security Risk Management Documentation**

516

517 To help demonstrate the safety and effectiveness of the device, manufacturers should provide the  
518 outputs of their security risk management processes in their premarket submissions, including  
519 their security risk management plan and security risk management report. A plan and report such

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<sup>35</sup> Examples of SW91 defect classification mapped to CWE can be found in Annex D of AAMI's SW91 Classification of Defects in Health Software. Additional information on CWE categories can be found at <https://cwe.mitre.org/>.

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520 as those described in AAMI TIR57,<sup>36</sup> inclusive of the system threat modeling, SBOM and  
521 associated documentation, and unresolved anomaly assessment(s) described above, should be  
522 sufficient to support the security risk management process aspect of demonstrating a reasonable  
523 assurance of safety and effectiveness.<sup>37</sup>

524

525 The security risk management report should:

526 • summarize the risk evaluation methods and processes, detail the security risk assessment,  
527 and detail the risk mitigation activities undertaken as part of a manufacturer's risk  
528 management processes; and

529 • provide traceability between the security risks, controls and the testing reports that  
530 ensure the device is reasonably secure.

531

## 532 **5. TPLC Security Risk Management**

533

534 Cybersecurity risks may continue to be identified throughout the device's TPLC. Manufacturers  
535 should ensure they have appropriate resources to identify, assess, and mitigate cybersecurity  
536 vulnerabilities as they are identified throughout the supported device lifecycle.

537

538 As part of using an SPDF, manufacturers should update their security risk management report as  
539 new information becomes available, such as when new threats, vulnerabilities, assets, or adverse  
540 impacts are discovered during development and after the device is released. When maintained  
541 throughout the device lifecycle, this documentation (e.g., threat modeling) can be used to quickly  
542 identify vulnerability impacts once a device is released and to support timely Corrective and  
543 Preventive Action (CAPA) activities described in 21 CFR 820.100.

544

545 Over the service life of a device, FDA recommends that the risk management documentation  
546 account for any differences in the risk management for fielded devices (e.g., marketed devices or  
547 devices no longer marketed but still in use). For example, if an update is not applied  
548 automatically for all fielded devices, then there will likely be different risk profiles for differing  
549 software configurations of the device. FDA recommends that vulnerabilities be assessed for any  
550 differing impacts for all fielded versions to ensure patient risks are being accurately assessed.  
551 Additional information as to whether a new premarket submission (e.g., PMA, PMA supplement,  
552 or 510(k)) or 21 CFR Part 806 reporting is needed based on postmarket vulnerabilities and  
553 general postmarket cybersecurity risk management are discussed in the Postmarket  
554 Cybersecurity Guidance.<sup>38</sup>

555

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<sup>36</sup> Details on the content for security risk management plans and reports beyond those specifically identified can be found in AAMI TIR57 Principles for medical device security—Risk management.

<sup>37</sup> While security architecture is likely captured as a component of the security risk management process, it is discussed separately for the purposes of this guidance due to the level of detail recommended to be provided by manufacturers in order to facilitate FDA review of the safety and effectiveness of the device.

<sup>38</sup> See Footnote 6.

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556 To demonstrate the effectiveness of a manufacturer’s processes, FDA recommends that a  
557 manufacturer track and record the measures and metrics below<sup>39</sup>, and report them in premarket  
558 submissions and PMA annual reports (21 CFR 814.84), when available.<sup>40</sup> Selecting appropriate  
559 measures and metrics for the processes that define an SPDF is important to ensure that device  
560 design appropriately addresses cybersecurity in compliance with QSR. At a minimum, FDA  
561 recommends tracking the following measures and metrics:

- 562
- 563 • Percentage of identified vulnerabilities that are updated or patched (defect density).
  - 564 • Time from vulnerability identification to when it is updated or patched.
  - 565 • Time from when an update or patch is available to complete implementation in devices  
566 deployed in the field.

567 Averages of the above measures should be provided if multiple vulnerabilities are identified and  
568 addressed. These averages may be provided over multiple time frames based on volume or in  
569 response to process or procedure changes to increase efficiencies of these measures over time.

## 570 **B. Security Architecture**

571

572 Manufacturers are responsible for identifying cybersecurity risks in their devices and the systems  
573 in which they expect those devices to operate, and implementing the appropriate controls to  
574 mitigate those risks. These risks may include those introduced by device reliance on hospital  
575 networks, cloud infrastructure, or “other functions” (as defined in FDA’s guidance “Multiple  
576 Function Device Products: Policy and Considerations), for example.<sup>41</sup> FDA recommends that all  
577 medical devices provide and enforce the security objectives in Section IV, above, but recognizes  
578 that implementations to address the security objectives may vary.

579

580 A security architecture, like a system architecture, defines the system and all end-to-end  
581 connections into and/or out of the system. A security architecture definition process<sup>42</sup> includes  
582 both high-level definitions of the devices and/or systems that interact, and detailed information  
583 on the implementations for how those interactions occur and are secured. It contains information  
584 that demonstrates that the risks considered during the risk management process are adequately  
585 controlled, which, in turn, supports the demonstration of the safety and effectiveness of the  
586 medical device system.

587

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<sup>39</sup> The measures and metrics provided are examples; alternative or additional measures and metrics may also be considered and reported.

<sup>40</sup> If a manufacturer has not released prior products or the premarket submission does not pertain to a marketed product (e.g., PMA supplement), FDA acknowledges that these measures and metrics might not be available, but recommends that manufacturers include these as part of their risk management plan and SPDF processes.

<sup>41</sup> See FDA Guidance “[Multiple Function Device Products: Policy and Considerations](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/multiple-function-device-products-policy-and-considerations)” available at: <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/multiple-function-device-products-policy-and-considerations>.

<sup>42</sup> NIST 800-160v1, Systems Security Engineering states that security architecture definition process generates a set of representative security views of the system architecture to inform the selection of an appropriate security architecture. The process also ascertains vulnerability and susceptibility to disruptions, hazards, and threats.

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588 Under 21 CFR 820.30(b), a manufacturer must establish and maintain plans that describe or  
589 reference the design and development activities and define responsibility for implementation.  
590 Such plans must be reviewed, updated, and approved as design and development evolves. 21  
591 CFR 820.30(b). Under 21 CFR 820.30(c), a manufacturer must establish and maintain  
592 procedures to ensure that the design requirements relating to a device are appropriate and address  
593 the intended use of the device, including the needs of the user and patient. Under 21 CFR  
594 820.30(d), a manufacturer must establish and maintain procedures for defining and documenting  
595 design output in terms that allow an adequate evaluation of conformance to design input  
596 requirements. 21 CFR 820.30(d) also states that design output procedures shall contain or make  
597 reference to acceptance criteria and shall ensure that those design outputs that are essential for  
598 the proper functioning of the device are identified.

599  
600 FDA recommends that these plans and procedures include design processes, design  
601 requirements, and acceptance criteria for the security architecture of the device such that they  
602 holistically address the cybersecurity considerations for the device and the system in which the  
603 device operates.

604  
605 FDA recommends that premarket submissions include documentation on the security  
606 architecture as discussed throughout this section. The objective in providing security architecture  
607 information in premarket submissions is to provide to the FDA the security context and trust-  
608 boundaries of the system in terms of the interfaces, interconnections, and interactions with  
609 external entities that the system has. The details of these elements enable the identification of the  
610 parts of the system through which attacks might be executed. Thus, as a whole, these details help  
611 to provide a sufficient understanding of the system such that FDA can evaluate adequacy of the  
612 architecture itself as it relates to safety and effectiveness.

613  
614 Analysis of the entire system should be performed to understand the full environment and  
615 context in which the device is expected to operate. The security architecture should include a  
616 consideration of system-level risks, including but not limited to risks related to the supply chain  
617 (e.g., to ensure the device remains free of malware, or vulnerabilities inherited from upstream  
618 dependencies such as third-party software, among others), design, production, and deployment  
619 (i.e., into a connected/networked environment).

620  
621 FDA recommends that this architecture information take the form of “views,” discussed in more  
622 detail in the following sub-sections and Appendix 2, and that these views be provided during  
623 premarket submissions to demonstrate safety and effectiveness. If the documentation identified  
624 in this section already exists in other risk management documentation, FDA does not expect  
625 manufacturers to separate out this information into new document(s); such documentation can be  
626 provided and the submission can reference the relevant sections.

627  
628 Throughout this section, FDA outlines the recommended security controls and recommendations  
629 on how to document the resultant security architecture in premarket submissions through specific  
630 Security Architecture Views.

### 631 **1. Implementation of Security Controls**

632

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633 FDA considers the way in which a device addresses cybersecurity risks and the way in which  
634 the device responds when exposed to cybersecurity threats as functions of the device design.  
635 Effective cybersecurity relies upon security being “built in” to a device, and not “bolted on”  
636 after the device is designed. FDA recommends that device manufacturers’ design processes  
637 include design inputs for cybersecurity controls.<sup>43</sup> Under 21 CFR 820.30(c), a manufacturer  
638 must establish and maintain procedures to ensure that the design requirements relating to a  
639 device are appropriate and address the intended use of the device, including the needs of the  
640 user and patient. Under 21 CFR 820.30(d), a manufacturer must establish and maintain  
641 procedures for defining and documenting design output in terms that allow an adequate  
642 evaluation of conformance to design input requirements. These output procedures shall contain  
643 or make reference to acceptance criteria and shall ensure that those design outputs that are  
644 essential for the proper functioning of the device are identified.

645  
646 FDA recommends that these procedures include design requirements and acceptance criteria  
647 for the security features built into the device such that they holistically address the  
648 cybersecurity considerations for the device and the system in which the device operates.

649  
650 Security controls allow manufacturers to achieve the security objectives outlined in Section IV  
651 above and are an integral part of an SPDF. FDA recommends that an adequate set of security  
652 controls will include, but not necessarily be limited to, controls from the following categories:

- 653
- 654 • Authentication;
  - 655 • Authorization;
  - 656 • Cryptography;
  - 657 • Code, Data, and Execution Integrity;
  - 658 • Confidentiality;
  - 659 • Event Detection and Logging;
  - 660 • Resiliency and Recovery; and
  - 661 • Updatability and Patchability.
- 662

663 For each of the security control categories above, specific control recommendations and  
664 implementation guidance for consideration to avoid common pitfalls are detailed in Appendix 1.

665  
666 Implementation of the controls should be applied across the system architecture using risk-based  
667 determinations associated with the subject connections and devices. Without adequate security  
668 controls across the system, which include management, technical, and operational controls, there  
669 is no reasonable assurance of safety and effectiveness. Additionally, deficiencies in the design of  
670 selected security controls or the implementation of those controls can have dramatic impacts on a  
671 system’s ability to demonstrate or maintain its safety and effectiveness.

672

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<sup>43</sup> There are useful frameworks to use in the generation of these design inputs including the OWASP Security by design principles, AAMI/ISA-62443-4-1, as well as medical device specific frameworks including the Hippocratic Oath for Connected Medical Devices, and Building Code for Medical Device Software Security. For a specific implementation of the OWASP Security by design principles, see the Medical Device and Health IT Joint Security Plan (JSP).



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673 FDA recommends the requirements and acceptance criteria for each of the above categories be  
674 provided in premarket submissions to demonstrate safety and effectiveness. Manufacturers  
675 should submit documentation in their premarket submissions demonstrating that the security  
676 controls for the categories above and further detailed in Appendix 1 have (1) been  
677 implemented, and (2) been tested in order to validate that they were effectively implemented  
678 (see Cybersecurity Testing section, V.C, below).

679  
680 Premarket documentation submitted by manufacturers may include the demonstration of  
681 comparable or additional security controls that may not be described in Appendix 1. If using  
682 alternate controls that are not described in this document, manufacturers should provide  
683 documentation and tracing of specific design features and security controls to demonstrate that  
684 they provide appropriate levels of safety and effectiveness. As cybersecurity design controls are  
685 established early in the development phase, FDA recommends that device manufacturers utilize  
686 the FDA Q-submission process to discuss with the agency design considerations for  
687 cybersecurity risk management throughout the device lifecycle.<sup>44</sup> Additional information on  
688 premarket documentation recommendations for design controls are discussed in the Security  
689 Architecture Views section below.

## **2. Security Architecture Views**

690  
691 In addition to the design control requirements (i.e., 21 CFR 820.30(b), 21 CFR 820.30(c), 21  
692 CFR 820.30(d), and 21 CFR 820.30(g)) outlined above for Security Architecture, 21 CFR  
693 820.100 requires that manufacturers establish policies, procedures, and other plans as appropriate  
694 to identify and respond to issues in devices. FDA recommends manufacturers develop and  
695 maintain security architecture view documentation as a part of the process for the design,  
696 development and maintenance of the system. If corrective and preventive actions are identified,  
697 these views can be used to help identify impacted functionality and solutions that address the  
698 risks.

699  
700 FDA recommends that premarket submissions include the architecture views described in this  
701 section. These architecture views can contribute to the demonstration of safety and effectiveness  
702 in premarket submissions by illustrating how the controls to address cybersecurity risks have  
703 been applied to the system.

704  
705 The security architecture may be expressed at different levels of abstraction and with different  
706 scopes or views.<sup>45</sup> The number and extent of the architecture views provided in the submission  
707 will be dependent on the attack surface(s) identified through threat modeling and risk  
708 assessments for the device. These views can therefore be an effective way to communicate the  
709 threat model to FDA and will naturally scale the documentation provided with the cybersecurity  
710 risk of the device.

711

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<sup>44</sup>For more information, see FDA’s guidance entitled “[Request for Feedback on Medical Device Submissions: The Q-Submission Program](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/requests-feedback-and-meetings-medical-device-submissions-q-submission-program),” available at <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/requests-feedback-and-meetings-medical-device-submissions-q-submission-program>.

<sup>45</sup> Architecture view is defined by NIST 800-160v1 as “A work product expressing the architecture of a system from the perspective of specific system concerns.”

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712 FDA recommends providing, at minimum, the following types of views in premarket  
713 submissions:

- 714 • Global System View;
- 715 • Multi-Patient Harm View;
- 716 • Updateability/Patchability View; and
- 717 • Security Use Case View(s).

718  
719 Documenting these views should include both diagrams and explanatory text. These diagrams  
720 and explanatory text should contain sufficient details to permit an understanding of how the  
721 assets within the system function holistically within the associated implementation details. For  
722 the security architecture views, manufacturers should consider the information outlined in  
723 Appendix 2 when determining the level of detail to include in premarket submissions.

- 724  
725 These security architecture views should:
- 726 • Identify security-relevant system elements and their interfaces;
  - 727 • Define security context, domains, boundaries, and external interfaces of the system;
  - 728 • Align the architecture with (a) the system security objectives and requirements, (b)  
729 security design characteristics; and
  - 730 • Establish traceability of architecture elements to user and system security requirements.

731  
732 The extent of these security views in a premarket submission is expected to vary based on the  
733 architecture and potential cybersecurity risk posed to the device. For example, systems with  
734 network and/or cloud access would be expected to have more Security Use Case Views than a  
735 system that only has a USB interface.

736

#### 737 (a) Global System View

738  
739 A global system view should describe the overall system, including the device itself and all  
740 internal and external connections. For interconnected and networked devices, this view should  
741 identify all interconnected elements, including any software update infrastructure(s), health care  
742 facility network impacts, intermediary connections or devices, cloud connections, etc.

743  
744 Depending on the complexity of the system, it may not be feasible to include all data flow  
745 specifics in a singular global system view. Additional views can be provided that detail the  
746 communication specifics as identified in Appendix 2 and do not need to be duplicated if captured  
747 in one of the other types of views detailed below.

748

#### 749 (b) Multi-Patient Harm View

750  
751 When devices are capable of connecting (wired or wirelessly) to another medical or non-  
752 medical product, to a network, or to the Internet, there is the possibility that multiple devices  
753 can be compromised simultaneously. Because of that connectivity, if a device is compromised,  
754 or if a non-device function (i.e., any function that does not fall within section 201(h) of the



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755 FD&C Act) that could impact the device function is compromised, the device may introduce a  
756 safety risk to patients through security risk. This may change the device’s intended use. For  
757 example, a non-device function could be hacked to perform a device function and ultimately  
758 harm patients.

759  
760 Depending on the device risk and use environment, a multiple-device compromise may have  
761 severe impacts for multiple patients, either through impact to the device itself and/or to health  
762 care facility operations (e.g., multiparameter bedside monitors all restarting at once, leaving all  
763 monitors connected to the same network no longer monitoring patient vitals and staffing levels  
764 not able to monitor all patient vitals).

765  
766 FDA recommends that manufacturers address how their device(s) and system(s) defend against  
767 and/or respond to attacks with the potential to harm multiple patients in a multi-patient harm  
768 view. This view should include the information outlined in Appendix 2. These risks, once  
769 identified, may also need to be assessed differently in the accompanying cybersecurity risk  
770 assessment due to the different nature of the risk.

771

#### 772 (c) Updatability and Patchability View

773

774 With the need to provide timely, reliable updates to devices in order to address emerging  
775 cybersecurity risks throughout the total product lifecycle of the device, FDA recommends  
776 manufacturers provide an updateability and patchability view. This view should describe the  
777 end-to-end process that permits software updates and patches to be provided (deployed) to the  
778 device, and should include detailed information as outlined in Appendix 2.

779

780 For example, if a device manufacturer intends to push software from a software update server to  
781 an in-clinic cardiac implant programmer, “end-to-end” means the path from the update server to  
782 the in-clinic programmer. The software update path will likely include traversing technology that  
783 the device manufacturer does not control, and therefore the design should provide for the  
784 protection of the end-to-end path and take into account any additional cybersecurity risk created  
785 or posed by those non-manufacturer-controlled technologies.

786

#### 787 (d) Security Use Case Views

788

789 In addition to the views identified above, security use case views should also be provided.  
790 Security use cases should be included for all system functionality through which a security  
791 compromise could impact the safety or effectiveness of the device. These security use cases  
792 should cover various operational states of elements in the system (e.g., power on, standby,  
793 transition states, etc.) and assess clinical functionality states of the system (e.g., programming,  
794 alarming, delivering therapy, send/receive data, reporting diagnostic results, etc.).

795

796 The number of security use cases that should be assessed will scale with the cybersecurity  
797 complexity and risk of the device. Each view should include detailed information as outlined in  
798 Appendix 2. For use cases identified that share the same security assessment, the associated

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799 diagrams and explanatory text can describe the multiple use cases covered by the view in lieu of  
800 providing duplicative information in multiple places. For example, programming commands and  
801 sending/receiving device data may share the same communication protocol and therefore may  
802 not exhibit differences between the security views for both scenarios, despite having different  
803 clinical risk assessments.  
804

## 805 **C. Cybersecurity Testing**

806  
807 As with other areas of product development, testing is used to demonstrate the effectiveness of  
808 design controls. While software development and cybersecurity are closely related disciplines,  
809 cybersecurity controls require testing beyond standard software verification and validation  
810 activities to demonstrate the effectiveness of the controls in a proper security context to therefore  
811 demonstrate that the device has a reasonable assurance of safety and effectiveness.  
812

813 Under 21 CFR 820.30(f), a manufacturer must establish and maintain procedures for verifying  
814 the device design. Such verification shall confirm that the design output meets the design input  
815 requirements. Under 21 CFR 820.30(g), a manufacturer must establish and maintain procedures  
816 for validating its device design. Such design validation shall include software validation and risk  
817 analysis, where appropriate. FDA recommends verification and validation include sufficient  
818 testing performed by the manufacturer on the cybersecurity of the system through which the  
819 manufacturer verifies and validates their inputs and outputs, as appropriate.  
820

821 Security testing documentation and any associated reports or assessments should be submitted in  
822 the premarket submission. FDA recommends that the following types of testing, among others,  
823 be provided in the submission:  
824

- 825 a. Security requirements
- 826 ○ Manufacturers should provide evidence that each design input requirement was
  - 827 implemented successfully.
  - 828 ○ Manufacturers should provide evidence of their boundary analysis and rationale
  - 829 for their boundary assumptions.
- 830
- 831 b. Threat mitigation
- 832 ○ Manufacturers should provide details and evidence of testing that demonstrates
  - 833 effective risk control measures according to the threat models provided in the
  - 834 system, use case, and call-flow views.
  - 835 ○ Manufacturers should ensure the adequacy of each cybersecurity risk control
  - 836 (e.g., security effectiveness in enforcing the specified security policy,
  - 837 performance for maximum traffic conditions, stability and reliability, as
  - 838 appropriate).
- 839
- 840 c. Vulnerability Testing (such as section 9.4 of ANSI/ISA 62443-4-1)

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- 841 ○ Manufacturers should provide details and evidence<sup>46</sup> of the following testing  
842 pertaining to known vulnerabilities:  
843     ▪ Abuse case, malformed, and unexpected inputs,  
844         • Robustness  
845         • Fuzz testing  
846     ▪ Attack surface analysis,  
847     ▪ Vulnerability chaining,  
848     ▪ Closed box testing of known vulnerability scanning,  
849     ▪ Software composition analysis of binary executable files, and  
850     ▪ Static and dynamic code analysis, including testing for credentials that are  
851         “hardcoded,” default, easily-guessed, and easily compromised.
- 852 d. Penetration testing
- 853     ○ The testing should identify and characterize security-related issues via tests that  
854         focus on discovering and exploiting security vulnerabilities in the product.  
855         Penetration test reports should be provided and include the following elements:  
856         ▪ Independence and technical expertise of testers,  
857         ▪ Scope of testing,  
858         ▪ Duration of testing,  
859         ▪ Testing methods employed, and  
860         ▪ Test results, findings, and observations.

861 Device manufacturers should indicate in the test reports where the testing was performed, and  
862 what level of independence those responsible for testing devices have from the developers  
863 responsible for designing devices. In some cases, it may be necessary to use third parties to  
864 ensure an appropriate level of independence between the two groups, such that vulnerabilities or  
865 other issues revealed during testing are appropriately addressed. For any third party test reports,  
866 manufacturers should provide the original third party report. For all testing, manufacturers  
867 should provide their assessment of any findings including rationales for not implementing or  
868 deferring any findings to future releases.  
869

870

871 As identified in Sections V.A.2. and V.A.3. above, vulnerabilities and anomalies identified  
872 during testing should be assessed for their security impacts as part of the security risk  
873 management process. In non-security software testing, a benefit analysis of a discovered defect  
874 may lead to the conclusion that an anomaly does not need to be fixed, as its impact on system  
875 functionality may be small or unlikely. Conversely, in security testing, the exploitability of an  
876 anomaly may necessitate that it is mitigated because of the greater, and different type of, harm  
877 that it could facilitate.

878

879 For issues that will be addressed in future releases (i.e., remediation deferred for a future  
880 software release because current risk was assessed to be acceptable), the plans for those releases  
881 should be detailed in the premarket submission to include the vulnerabilities that future software  
882 releases will address, anticipated timelines for release, whether devices released in the interim  
883 will receive those updates, and how long it will take the update to reach the devices.

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<sup>46</sup> For any testing tools or software used, the details provided may include, but may not be limited to, the name of the tool, version information as applicable, and any settings or configuration options for the tools used.

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884 There are numerous authoritative resources for outlining security testing that may partially fulfill  
885 the testing outlined above.<sup>47</sup> FDA recommends that cybersecurity testing should occur  
886 throughout the SPDF. Security testing early in development can ensure that security issues are  
887 addressed prior to impacting release timelines and can prevent the need to redesign or re-  
888 engineer the device. After release, cybersecurity testing should be performed at regular intervals  
889 (e.g., annually) to ensure that potential vulnerabilities are identified and able to be addressed  
890 prior to their ability to be exploited.  
891

## 892 **VI. Cybersecurity Transparency**

893  
894 In order for users to manage security risks in devices, either by an end user or within a larger  
895 risk management framework like the NIST CSF, transparency is critical to ensure safe and  
896 effective use and integration of devices and systems. This transparency can be conveyed  
897 through both labeling and the establishment of vulnerability management plans. However,  
898 different types of users (e.g., manufacturers, servicers, patients, etc.) will have different  
899 abilities to take on a mitigation role, and the need for actions to ensure continued cybersecurity  
900 should be appropriate for the type of user.

### 901 **A. Labeling Recommendations for Devices with** 902 **Cybersecurity Risks**

903  
904 FDA regulates device labeling in several ways. For example, section 502(f) of the FD&C Act  
905 requires that labeling include adequate directions for use. Under section 502(a)(1) of the FD&C  
906 Act, a medical device is deemed misbranded if its labeling is false or misleading in any  
907 particular.  
908

909 For devices with cybersecurity risks, informing users of relevant security information may be an  
910 effective way to comply with labeling requirements relating to such risks. FDA also believes that  
911 informing users of security information through labeling may be an important part of QSR  
912 design controls to help mitigate cybersecurity risks and help ensure the continued safety and  
913 effectiveness of the device. Therefore, when drafting labeling for inclusion in a premarket  
914 submission, a manufacturer should consider all applicable labeling requirements and how  
915 informing users through labeling may be an effective way to manage cybersecurity risks and/or  
916 to ensure the safe and effective use of the device. Any risks transferred to the user should be  
917 detailed and considered for inclusion as tasks during usability testing (e.g., human factors  
918 testing<sup>48</sup>) to ensure that the type of user has the capability to take appropriate actions to manage  
919 those risks-.

---

<sup>47</sup> The following standards may partially meet the security testing recommendations in ANSI/UL 2900 Software Cybersecurity for Network-Connectable Products and ANSI/ISA-62443-4-1-2018 Security for industrial automation and control systems Part 4-1: Product security development life-cycle requirements. Additional standards may also meet or partially meet the testing recommendations outlined in this section.

<sup>48</sup> See FDA Guidance “[Applying Human Factors and Usability Engineering to Medical Devices](https://www.fda.gov/regulatory-information/search-fda-guidance-documents/applying-human-factors-and-usability-engineering-medical-devices)” available at: <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/applying-human-factors-and-usability-engineering-medical-devices>

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920  
921 The recommendations below aim to communicate to users relevant device security information  
922 that may enable their own ongoing security posture, thereby helping ensure a device remains safe  
923 and effective throughout its lifecycle. The depth of detail, the exact location in the labeling for  
924 specific types of information (e.g., operator’s manual, security implementation guide), and the  
925 method to provide this information should account for the intended user of the information.  
926 Instructions to manage cybersecurity risks should be understandable to the intended audience,  
927 which might include patients or caregivers with limited technical knowledge. The manufacturer  
928 may wish to employ methods to ensure certain information is available only to the user, and if it  
929 does so through an online portal, should provide an up-to-date link.<sup>49</sup>

930  
931 FDA recommends the following be included in labeling to communicate relevant security  
932 information to users.<sup>50</sup>

- 933
- 934 1. Device instructions and product specifications related to recommended  
935 cybersecurity controls appropriate for the intended use environment (e.g., anti-  
936 malware software, use of a firewall, password requirements).  
937
  - 938 2. Sufficiently detailed diagrams for users that allow recommended cybersecurity  
939 controls to be implemented.  
940
  - 941 3. A list of network ports and other interfaces that are expected to receive and/or  
942 send data. This list should include a description of port functionality and indicate  
943 whether the ports are incoming, outgoing, or both, along with approved  
944 destination end-points.  
945
  - 946 4. Specific guidance to users regarding supporting infrastructure requirements so  
947 that the device can operate as intended (e.g., minimum networking requirements,  
948 supported encryption interfaces).  
949
  - 950 5. A SBOM as specified in Section V.A.2.b or in accordance with an industry  
951 accepted format to effectively manage their assets, to understand the potential  
952 impact of identified vulnerabilities to the device (and the connected system), and  
953 to deploy countermeasures to maintain the device’s safety and effectiveness.  
954 Manufacturers should provide or make available SBOM information to users on a  
955 continuous basis. If an online portal is used, an up-to-date link should be  
956 provided. The SBOM should be in a machine readable format.  
957
  - 958 6. A description of systematic procedures for users to download version-identifiable  
959 manufacturer-authorized software and firmware, including a description of how  
960 users will know when software is available.

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<sup>49</sup> For more information regarding FDA’s policy on labeling changes and submission requirements, manufacturers can use the FDA Guidance Search Tool to identify relevant guidance documents for their product and submission type. <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/>.

<sup>50</sup> See IEC TR 80001-2-2 and IEC TR 80001-2-8 and IEC TR 80001-2-9 for further labeling information for compliance with these standards.

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7. A description of how the design enables the device to respond when anomalous conditions are detected (i.e., security events) in order to maintain safety and effectiveness. This should include notification to the user and logging of relevant information. Security event types could be configuration changes, network anomalies, login attempts, or anomalous traffic (e.g., send requests to unknown entities).
  8. A high-level description of the device features that protect critical functionality (e.g., backup mode, disabling ports/communications, etc.).
  9. A description of backup and restore features and procedures to restore authenticated configurations.
  10. A description of the methods for retention and recovery of device configuration by an authenticated authorized user.
  11. A description of the secure configuration of shipped devices, a discussion of the risk tradeoffs that might have been made about hardening options implemented by the device manufacturer, and instructions for user-configurable changes. Secure configurations may include end point protections such as anti-malware, firewall/firewall rules, allow lists, deny lists, security event parameters, logging parameters, and physical security detection, among others.
  12. Where appropriate for the intended use environment, a description of how forensic evidence is captured, including but not limited to any log files kept for a security event. Log file descriptions should include how and where the log file is located, stored, recycled, archived, and how it could be consumed by automated analysis software (e.g., Intrusion Detection System, IDS).
  13. Where appropriate, technical instructions to permit secure network deployment and servicing, and instructions for users on how to respond upon detection of a cybersecurity vulnerability or incident.
  14. Information, if known or anticipated, concerning device cybersecurity end of support and end of life. At the end of support, a manufacturer may no longer be able to reasonably provide security patches or software updates. If the device remains in service following the end of support, the manufacturer should have a pre-established and pre-communicated process for transferring the risks highlighting that the cybersecurity risks for end-users can be expected to increase over time.
  15. Information on securely decommissioning devices by sanitizing the product of sensitive, confidential, and proprietary data and software.

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1006 A revision-controlled, Manufacturer Disclosure Statement for Medical Device Security (MDS2)  
1007 and Customer Security Documentation as outlined in the HSCC Joint Security Plan (JSP) may  
1008 address a number of the above recommendations.

## 1009 **B. Vulnerability Management Plans**

1010  
1011 Recognizing that cybersecurity risks evolve as technology evolves throughout a device's TPLC,  
1012 FDA recommends that manufacturers establish a plan for how they will identify and  
1013 communicate vulnerabilities that are identified after releasing the device with users. This plan  
1014 can also support risk management processes in accordance with 21 CFR 820.30(g) and corrective  
1015 and preventive action processes in accordance with 21 CFR 820.100.

1016  
1017 FDA recommends that manufacturers submit their vulnerability communication plans as part of  
1018 their premarket submissions so that FDA can assess whether the manufacturer has sufficiently  
1019 addressed how to maintain the safety and effectiveness of the device after marketing  
1020 authorization is achieved.

1021  
1022 Vulnerability communication plans should include the following elements:

- 1023 a) Personnel responsible;  
1024 b) Sources, methods, and frequency for monitoring for and identifying vulnerabilities (e.g.,  
1025 researchers, NIST NVD, third-party software manufacturers, etc.);  
1026 c) Periodic security testing to test identified vulnerability impact;  
1027 d) Timeline to develop and release patches;  
1028 e) Update processes;  
1029 f) Patching capability (i.e., rate at which update can be delivered to devices);  
1030 g) Description of their coordinated vulnerability disclosure process; and  
1031 h) Description of how manufacturer intends to communicate forthcoming remediations,  
1032 patches, and updates to customers.

1033  
1034 Additional recommendations on coordinated vulnerability disclosure plans may be found in  
1035 FDA's Postmarket Cybersecurity Guidance.<sup>51</sup>

1036

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<sup>51</sup> See Footnote 10.

## 1037 **Appendix 1. Security Control Categories and Associated** 1038 **Recommendations**

1039  
1040 The following sections provide detailed descriptions of each of the security control categories  
1041 introduced in Section V.B.1. as well as specific recommendations for security controls and their  
1042 implementation to avoid common pitfalls.  
1043

### 1044 **A. Authentication**

1045 There are generally two types of authentication controls—information and entities—and a  
1046 properly-secured system is able to prove the existence of both.  
1047

1048 Authentication of *information*<sup>52</sup> exists where the device and the system in which it operates is  
1049 able to prove that information originated at a known and trusted source, and that the information  
1050 has not been altered in transit between the original source and the point at which authenticity is  
1051 verified. It is important to note that while authenticity implies that data is accurate and has been  
1052 safeguarded from unauthorized user modification (i.e., integrity), integrity alone does not  
1053 provide assurance that the data is real and came from a trusted source. Therefore, for the  
1054 purposes of this guidance, authentication is discussed as a larger security objective over integrity.  
1055

1056 Authentication of *entities* exists where a device and the system in which it operates is able to  
1057 prove the identity of an endpoint (whether hardware and/or software) from which it is sending  
1058 and/or receiving information, or authorized user/operator at that endpoint.  
1059

1060 As part of normal operations within a secure system, devices are expected to verify the  
1061 authenticity of information from external entities, as well as prove the authenticity of information  
1062 that they generate. A system that appropriately accounts for authenticity will evaluate and ensure  
1063 authenticity for: (1) information at rest (stored); (2) information in transit (transmitted); (3) entity  
1064 authentication of communication endpoints, whether those endpoints consist of software or  
1065 hardware; (4) software binaries; (5) integrity of the execution state of currently running software;  
1066 and (6) any other appropriate parts of the system where a manufacturer’s threat model and/or risk  
1067 analyses reveal the need for it.  
1068

1069 On a technical level, the strength of a device’s authentication scheme is defined by the amount of  
1070 effort, including time, that an unauthorized party would need to expend to identify the  
1071 decomposition of the authentication scheme. For example, this could be the time and resources  
1072 necessary to determine the correct “output” of a cryptographic function from which a  
1073 cryptographically-based authentication scheme is built and which an unauthorized party could  
1074 use to bypass the authentication scheme and gain access to the system.  
1075

1076 When choosing an authentication scheme, manufacturers should keep in mind the following  
1077 generally applicable characteristics of different types of schemes. Implicit authentication

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<sup>52</sup> For the purposes of this control, “information” includes the software/firmware itself, as well as input and output data.



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1078 schemes, based solely on non-cryptographic interfaces, handshakes, and/or protocols, are  
1079 inherently weak because, once they are reverse-engineered, an unauthorized user can easily  
1080 emulate the correct behavior and appear to be authorized. Cryptographic authentication protocols  
1081 are generally superior, but they need careful design choices and implementation practices to  
1082 achieve their full strength. In addition, these schemes are still limited by the confidentiality of the  
1083 cryptographic keys needed to interact with the scheme, and by the integrity of the devices that  
1084 hold or otherwise leverage those keys (see the cryptography subsection below). Therefore, for  
1085 device operations where non-authenticated behavior could lead to harm, devices should  
1086 implement additional, non-routine signals of intent based on physical actions, such as a  
1087 momentary switch, to authorize the command/session.  
1088

1089 The following list provides additional recommendations for the implementation of authentication  
1090 schemes:  
1091

- 1092 • Use cryptographically strong<sup>53</sup> authentication, where the authentication functionality  
1093 resides on the device, to authenticate personnel, messages, commands updates, and as  
1094 applicable, all other communication pathways. Hardware-based security solutions should  
1095 be considered and employed when possible;
- 1096 • Authenticate external connections at a frequency commensurate with the associated risks.  
1097 For example, if a device connects to an offsite server, then the device and the server  
1098 should mutually authenticate each session and limit the duration of the session, even if  
1099 the connection is initiated over one or more existing trusted channels;
- 1100 • Use appropriate user authentication (e.g., multi-factor authentication to permit privileged  
1101 device access to system administrators, service technicians, or maintenance personnel,  
1102 among others, as needed);
- 1103 • Require authentication, and permission in certain instances, before permitting software or  
1104 firmware updates, including those updates affecting the operating system, applications,  
1105 and anti-malware functionality;
- 1106 • Strengthen password protections. Do not use passwords that are hardcoded, default,  
1107 easily-guessed, or easily compromised (e.g., passwords that are the same for each device;  
1108 unchangeable; can persist as default; difficult to change; and/or vulnerable to public  
1109 disclosure);
- 1110 • Implement anti-replay measures in critical communications such as potentially harmful  
1111 commands. This can be accomplished with the use of cryptographic nonces (an arbitrary  
1112 number used only once in a cryptographic communication);
- 1113 • Provide mechanisms for verifying the authenticity of information originating from the  
1114 device, such as telemetry. This is especially important for data that, if spoofed or  
1115 otherwise modified, could result in patient harm, such as the link between a continuous  
1116 glucose monitor (CGM) system and an automated insulin pump;
- 1117 • Do not rely on cyclic redundancy checks (CRCs) as security controls. CRCs do not  
1118 provide integrity or authentication protections in a security environment. While CRCs are  
1119 an error detecting code and provide integrity protection against environmental factors  
1120 (e.g., noise or EMC), they do not provide protections against an intentional or malicious  
1121 actor; and

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<sup>53</sup> See the definition of security strength in Appendix 4, Terminology.

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- 1122       • Consider how the device and/or system should respond in event of authentication  
1123       failure(s).

## 1124       **B. Authorization**

1125       For the purposes of this guidance, authorization is the right or permission that is granted to a  
1126       system entity (e.g., a device, server, or software function) to access a system resource. More  
1127       specifically, as a defensive measure, an authorization scheme enforces privileges, i.e. “rights,”  
1128       associated with authenticated sessions, identities and/or roles. These privileges either permit  
1129       allowed behavior, or refuse disallowed behavior in order to ensure that system resources are only  
1130       accessed in accepted ways, by accepted parties.

1131  
1132       Within an adequately designed authorization scheme, the principle of least privileges<sup>54</sup> should be  
1133       applied to users, system functions, and others, to only allow those entities the levels of system  
1134       access necessary to perform a specific function.

1135  
1136       For example, in a situation in which a malicious actor has gained access to a credential  
1137       associated with patient privileges, that malicious actor should not be able to access device  
1138       resources or functionality reserved for the manufacturer or for the health care provider, such as  
1139       device maintenance routines or the ability to change medication dosage amounts.

1140  
1141       While authentication schemes based on cryptographically-proven designs are generally  
1142       considered more robust and are therefore preferred, meaningful authorization checks can be  
1143       performed based on other compelling evidence (e.g., benefit/risk assessment in accordance with  
1144       Section 6.5 of AAMI TIR57 and associated supporting justification and as evidenced through  
1145       security testing). For example, a medical device programmer that is capable of Near-Field  
1146       Communications (NFC) could have elevated privileges that are granted based on a signal of  
1147       intent<sup>55</sup> over NFC that cannot physically be produced by another unauthorized device over  
1148       Radio-Frequency (RF) (e.g., a home monitor).

1149  
1150       The following list provides recommended design implementations for an authorization scheme:

- 1151       • Limit authorized access to devices through the authentication of users (e.g., user ID and  
1152       password, smartcard, biometric, certificates, or other appropriate authentication method);
- 1153       • Use automatic timed methods to terminate sessions within the system where appropriate  
1154       for the use environment;
- 1155       • Employ an authorization model that incorporates the principle of least privileges by  
1156       differentiating privileges based on the user role (e.g., caregiver, patient, health care  
1157       provider, system administrator) or device functions; and
- 1158       • Design devices to “deny by default” (i.e., that which is not expressly permitted by a  
1159       device is denied by default). For example, the device should generally reject all  
1160       unauthorized connections (e.g., incoming TCP, USB, Bluetooth, serial connections).  
1161       Ignoring requests is one form of denying authorization.

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<sup>54</sup> CNSI 4009-2015 defines least privilege as “The principle that a security architecture should be designed so that each entity (e.g., user, asset) is granted the minimum system resources and authorizations that the entity needs to perform its function.”

<sup>55</sup> Signal of intent in this use is specific to the implementation of NFC communications.

1162 **C. Cryptography**

1163 Cryptographic algorithms and protocols are recommended to be implemented to achieve the  
1164 secure by design objectives outlined in Section IV. While high-quality, standardized  
1165 cryptographic algorithms and protocols are readily available, several commercial products that  
1166 include cryptographic protections have been shown to have exploitable vulnerabilities due to  
1167 improper configurations and/or implementations.

1168  
1169 While other sections of this guidance reference cryptographic controls, the following  
1170 recommendations are specifically related to the selection and implementation of the underlying  
1171 cryptographic scheme used by a device and the larger system in which it operates:  
1172

- 1173 • Select industry-standard cryptographic algorithms and protocols, and select appropriate  
1174 key generation, distribution, management and protection, as well as robust nonce  
1175 mechanisms.
- 1176 • Use current NIST recommended standards for cryptography (e.g., FIPS 140-2<sup>56</sup>, NIST  
1177 Suite B<sup>57</sup>), or equivalent-strength cryptographic protection that are expected to be  
1178 considered cryptographically strong throughout the service life of the device.
- 1179 • Design a system architecture and implement security controls to prevent a situation where  
1180 the full compromise of any single device can result in the ability to reveal keys for other  
1181 devices.
  - 1182 ○ For example, avoid using master-keys stored on device, or key derivation  
1183 algorithms based solely on device identifiers or other readily discoverable  
1184 information.
  - 1185 ○ Avoid using device serial numbers as keys or as part of keys. Device serial  
1186 numbers may be disclosed by patients seeking additional information on their  
1187 device or might be disclosed during a device recall to identify affected products  
1188 and should be avoided as part of the key generation process. Public-key  
1189 cryptography can be employed to help meet this objective.
- 1190 • Implement cryptographic protocols that permit negotiated parameters/versions such that  
1191 the most recent, secure configurations are used, unless otherwise necessary.
- 1192 • Do not allow downgrades, or version rollbacks, unless absolutely necessary for safety  
1193 reasons. Downgrades can allow attackers to exploit prior, less protected versions and  
1194 should be avoided.

1195 **D. Code, Data, and Execution Integrity**

1196 Many cybersecurity incidents are caused, at their root, by the violation of some form of device  
1197 integrity. This includes the violation of stored code, stored and operational data, or execution  
1198 state. The following recommendations are provided to address each of these categories.  
1199

- 1200 • **Code Integrity**

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<sup>56</sup> NIST FIPS 140-2 Cryptographic Module Validation Program available at:

<https://csrc.nist.gov/Projects/Cryptographic-Module-Validation-Program/Standards>

<sup>57</sup> NIST FIPS 140-2 Suite B available at: <https://csrc.nist.gov/CSRC/media/projects/cryptographic-module-validation-program/documents/security-policies/140sp2851.pdf>

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- Authenticate firmware and software. Verify authentication tags (e.g., signatures, message authentication codes (MACs)) of software/firmware content, version numbers, and other metadata. The version numbers intended to be installed should themselves be signed or have MACs. Devices should be electronically and visibly identifiable (e.g., Unique device identifier (UDI), model number, serial number);
  - Allow installation of cryptographically authenticated firmware and software updates, and do not allow installation where such cryptographic authentication either is absent or fails. Use cryptographically signed updates to help prevent any unauthorized reductions in the level of protection (downgrade or rollback attacks) by ensuring that the new update represents an authorized version change.
    - One possible approach for authorized downgrades would be to sign new metadata for downgrade requests which, by definition, only happen in exceptional circumstances;
  - Ensure that the authenticity of software, firmware, and configuration are validated prior to execution, e.g., “allow-listing”<sup>58</sup> based on digital signatures;
  - Disable or otherwise restrict unauthorized access to all test and debug ports (e.g., JTAG, UART) prior to delivering products; and
  - Employ tamper evident seals on device enclosures and their sensitive communication ports to help verify physical integrity.
- **Data Integrity**
    - Verify the integrity of all incoming data, ensuring that it is not modified in transit or at rest. Cryptographic authentication schemes verify integrity, but do not verify validity;
    - Validate that all data originating from external sources is well-formed and compliant with the expected protocol or specification. Additionally, as appropriate, validate data ranges to ensure they fall within safe limits; and
    - Protect the integrity of data necessary to ensure the safety and effectiveness of the device, e.g., critical configuration settings such as energy output.
  - **Execution Integrity**
    - Use industry-accepted best practices to maintain and verify integrity of code while it is being executed on the device. For example, Host-based Intrusion Detection/Prevention Systems (HIDS/HIPS) can be used to accomplish this goal; and
    - Carefully design and review all code that handles the parsing of external data using automated (e.g., static and dynamic analyses) and manual (i.e., code review) methods.

## 1238 E. Confidentiality

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<sup>58</sup> For the purposes of this guidance, “allow-list” means “a list of discrete entities, such as hosts or applications that are known to be benign and are approved for use within an organization and/or information system.” This term is leveraged from definition of “whitelist” in NIST SP 800-128.

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1239 Manufacturers should ensure support for the confidentiality<sup>59</sup> of any/all data whose disclosure  
1240 could lead to patient harm (e.g., through the unauthorized use of otherwise valid credentials, lack  
1241 of encryption). Loss of confidentiality of credentials could be used by a threat-actor to effect  
1242 multi-patient harm. Lack of encryption to protect sensitive information and or data at rest and in  
1243 transit can expose this information to misuse that can lead to patient harm. For example,  
1244 confidentiality is required in the handling and storage of cryptographic keys used for  
1245 authentication because disclosure could lead to unauthorized use/abuse of device functionality.  
1246

1247 The proper implementation of authorization and authentication schemes as described in Sections  
1248 (a) and (b) of this appendix (Appendix 1 – Security Control Categories and Associated  
1249 Recommendations) will generally assure confidentiality. However, manufacturers should  
1250 evaluate and assess whether this is the case during their threat modeling and other risk  
1251 management activities and make any appropriate changes to their systems to ensure appropriate  
1252 confidentiality controls are in place.

## **F. Event Detection and Logging**

1253  
1254 Event detection and logging are critical capabilities that should be present in a device and the  
1255 larger system in which it operates in order to ensure that suspected and successful attempts to  
1256 compromise a medical device may be identified and tracked. These event detection capabilities  
1257 and logs should include storage capabilities, if possible, so that forensic discovery may later be  
1258 performed.  
1259

1260 While many of the following recommendations are tailored for workstations, the concepts  
1261 presented below also apply to embedded computing devices. Manufacturers should consider  
1262 these items for all devices:  
1263

- 1264 • Implement design features that allow for security compromises and suspected  
1265 compromise attempts to be detected, recognized, logged, timed, and acted upon during  
1266 normal use. Acting upon security events should consider the benefit/risk assessment in  
1267 accordance with Section 6.5 of AAMI TIR57 in determining whether it is appropriate to  
1268 affect standard device functionality during a security event.
- 1269 • Ensure the design enables forensic evidence capture.<sup>60</sup> The design should include  
1270 mechanisms to create and store log files off the device to track security events.  
1271 Documentation should include how and where log files are located, stored, recycled,  
1272 archived, and how they could be consumed by automated analysis software (e.g.,

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<sup>59</sup>For the purposes of this guidance, loss of confidential protected health information (PHI) is not considered patient harm. Although protecting the confidentiality of PHI is beyond the scope of this document, it should be noted that manufacturers and other entities, depending on the facts and circumstances, may be obligated to protect the confidentiality, integrity and availability of PHI throughout the product lifecycle, in accordance with applicable federal and state laws, including the Health Insurance Portability and Accountability Act (HIPAA). For more information on HIPAA, please visit <https://www.hhs.gov/hipaa/for-professionals/security/laws-regulations/index.html>

<sup>60</sup> Forensic evidence capture is a necessary part of digital forensics. [NIST SP 800-86](#)

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- 1273 Intrusion Detection System (IDS)). Examples of security events include, but are not  
1274 limited to, configuration changes, network anomalies, login attempts, and anomalous  
1275 traffic (e.g., sending requests to unknown entities).
- 1276 • Design devices such that the potential impact of vulnerabilities is limited by specifying a  
1277 secure configuration. Secure configurations may include endpoint protections, such as  
1278 anti-malware, firewall/firewall rules, allow-listing, defining security event parameters,  
1279 logging parameters, and/or physical security detection.
  - 1280 • Design devices such that they may integrate and/or leverage antivirus/anti-malware  
1281 protection capabilities. These capabilities may vary depending on the type of device and  
1282 the software and hardware components it contains:
    - 1283 ○ For devices that leverage Windows Operating System:
      - 1284 ■ Antivirus/anti-malware is recommended on the device. Manufacturers are  
1285 recommended to qualify multiple options to support user preferences for  
1286 different options, especially if the device is used in health care facility  
1287 environments.
    - 1288 ○ For devices that leverage other Commercial Operating Systems (i.e., Ubuntu,  
1289 Unix, Linux, Apple, Android, etc.)
      - 1290 ■ Antivirus/anti-malware may be recommended based on the environment  
1291 and associated risks of the device. Different operating systems will likely  
1292 follow a case-by-case determination based on network exposure and risk.
    - 1293 ○ For devices that leverage Embedded Operating Systems (i.e., Real-Time  
1294 Operating Systems, Windows embedded, etc.)
      - 1295 ■ Antivirus/anti-malware is generally not needed unless a particular risk or  
1296 threat is identified that would not be addressed by other expected security  
1297 controls.
  - 1298 • Design devices to enable software configuration management and permit tracking and  
1299 control of software changes to be electronically obtainable (i.e., machine readable) by  
1300 authorized users.
  - 1301 • Design devices to facilitate the performance of variant analyses such that the same  
1302 vulnerabilities can be identified across device models and product lines.
  - 1303 • Design devices to notify users when malfunctions, including those potentially related to a  
1304 cybersecurity breach, are detected.
  - 1305 • Consider designing devices such that they are able to produce a SBOM in a machine  
1306 readable<sup>61</sup> format.

## 1307 **G. Resiliency and Recovery**

1308 Devices should be designed to be resilient to possible cybersecurity incident scenarios (also  
1309 known as “cyber-resiliency”). Cyber-resiliency capabilities are important for medical devices  
1310 because they provide a safety margin against unknown future vulnerabilities.

1311  
1312 The following recommendations are intended to help designers achieve cyber-resiliency:  
1313

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<sup>61</sup> Recommendation 2.2 from the Health Care Industry and Cybersecurity Task Force (HCIC TF) Report on Improving Cybersecurity in the Health Care Industry available here:  
<https://www.phe.gov/Preparedness/planning/CyberTF/Documents/report2017.pdf>

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- 1314 • Implement features that protect critical functionality and data, even when the device has  
1315 been partially compromised. For example, process isolation, virtualization techniques,  
1316 and hardware-backed trusted execution environments all provide mechanisms to  
1317 potentially contain the impact of a successful exploitation of a device.
- 1318 • Design devices to provide methods for retention and recovery of trusted default device  
1319 configuration by an authenticated, authorized user.
- 1320 • Design devices to specify the level of resilience, or independent ability to function, that  
1321 any component of the system possesses when its communication capabilities with the rest  
1322 of the system are disrupted, including disruption of significant duration.
- 1323 • Design devices to be resilient to possible cybersecurity incident scenarios such as  
1324 network outages, Denial of Service,<sup>62</sup> excessive bandwidth usage by other products,  
1325 disrupted quality of service<sup>63</sup> (QoS), and/or excessive jitter<sup>64</sup> (i.e., a variation in the delay  
1326 of received packets).

## 1327 **H. Firmware and Software Updates**

1328 Devices should be capable of being updated in a secure and timely manner to maintain safety and  
1329 effectiveness throughout the product's lifecycle. Despite best efforts, undiscovered, exploitable  
1330 vulnerabilities may exist in devices after they are marketed. This is especially true over the  
1331 device's service life, as threats evolve over time and exploit methods change, and become more  
1332 sophisticated.

1333  
1334 FDA recommends that manufacturers should not only build in the ability for devices to be  
1335 updated, but that manufacturers also plan for the rapid testing, evaluation, and patching of  
1336 devices deployed in the field. The following recommendations can help to achieve this:

- 1337  
1338 • Design devices to anticipate the need for software and firmware patches and updates to  
1339 address future cybersecurity vulnerabilities. This will likely necessitate the need for  
1340 additional storage space and processing resources.
- 1341 • Consider update process reliability and how update process works in event of  
1342 communication interruption or failure. This should include both considerations for  
1343 hardware impacts (timing specifics of interruptions) and which phase of the update  
1344 process the interruption or failure occurs.
- 1345 • Consider cybersecurity patches and updates that are independent of regular feature update  
1346 cycles.
- 1347 • Implement processes, technologies, security architectures, and exercises to facilitate the  
1348 rapid verification, validation, and distribution of patches and updates.
- 1349 • Preserve and maintain full build environments and virtual machines, regression test  
1350 suites, engineering development kits, emulators, debuggers, and other related tools that  
1351 were used to develop and test the original product to ensure updates and patches may be  
1352 applied safely and in a timely manner.

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<sup>62</sup> Denial of Service is an attack that prevents or impairs the authorized use of the information system, resources, or services.

<sup>63</sup> From CNSSI 4009 Committee on National Security Systems (CNSS) Glossary.

<sup>64</sup> From NIST SP 800-127 Guide to Securing WiMAX Wireless Communications.

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- Maintain necessary third-party licenses throughout the supported lifespan of the device. Develop contingency plans for the possibility that a third-party company goes out of business or stops supporting a licensed product. Modular designs should be considered such that third-party solutions could be readily replaced.

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1358 **Appendix 2. Submission Documentation for Security**  
1359 **Architecture Flows**

1360  
1361 In premarket submissions, FDA recommends that manufacturers provide detailed information for  
1362 the views identified in Section V.B.2. Methods for providing the views and the expectations for  
1363 the level of detail to provide are discussed in the sections below. In addition to diagrams and  
1364 explanatory text, call-flow views can be provided to convey some of the information details  
1365 expected to be addressed in the architecture views.

1366 **A. Call-Flow Diagrams**

1367  
1368 A call-flow view is a diagram with explanatory text that describes the sequence of process or  
1369 protocol steps in explicit detail. For each of the views, manufacturers may provide call-flow  
1370 information to detail the communications included in the associated use case.

1371  
1372 Call-flow views should provide specific protocol details of the communication pathways  
1373 between parts of the system, to include authentication or authorization procedures and session  
1374 management techniques. These views should be sufficiently detailed such that engineers and  
1375 reviewers should be able to logically and easily follow data, code, and commands from any asset  
1376 (e.g., a manufacturer server) to any other associated asset (e.g., a medical device), while possibly  
1377 crossing intermediate assets (e.g., application). The call-flow views may also include items from  
1378 the information details identified below for the views identified in Section V.B.2. if the  
1379 information is better represented or conveyed through a call-flow view.

1380 **B. Information Details for an Architecture View**

1381  
1382 For each view described in Section V.B.2., manufacturers should provide a system-level  
1383 description and analysis inclusive of end-to-end security analyses of all the communications in  
1384 the system regardless of intended use. This should include detailed diagrams and traces for all  
1385 communication paths as described below. Security-relevant analysis requires the ability to  
1386 construct and follow a detailed trace for important communication paths, which describes how  
1387 data, code, and commands are protected between any two assets in the device's system. This  
1388 analysis can also help identify the software that should be included in the SBOM for each device.

1389  
1390 The FDA recommends that security architecture views should include at least the following:

- 1391
- 1392 a. Detailed diagrams and supporting explanatory text that identify all manufacturer  
1393 and network assets of the system in which the device will operate, including but  
1394 not limited to:
    - 1395
    - 1396 i. Device hardware itself (including assessments for any commercial  
1397 platforms);

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- 1398 ii. Applications, hardware, and/or other supporting assets that directly  
1399 interact with the targeted device, such as configuration,  
1400 installation/upgrade, and data transfer applications;
- 1401 iii. Health care facility-operated assets;
- 1402 iv. Communications/networking assets; and
- 1403 v. Manufacturer-controlled assets, including any servers that interact with  
1404 external entities (e.g., a service that collects and redistributes device data,  
1405 or a firmware update server).
- 1406
- 1407 b. For every communication path that exists between any two assets in the security  
1408 use case view (and/or explanatory text), including indirect connections when there  
1409 is at least one intermediate asset (e.g., an app), the following details should be  
1410 provided:
- 1411 i. A list of the communication interfaces and paths, including  
1412 communication paths (e.g., between two assets through an intermediary),  
1413 including any unused interfaces;
- 1414 ii. An indication of whether the path is used for data, code, and/or  
1415 commands, and type of data/information/code being transferred;
- 1416 iii. Protocol name(s), version number(s), and ports/channels/frequencies;
- 1417 iv. Detailed descriptions of the primary and all available functionality for  
1418 each system asset, including assessment of any functionality that is built in  
1419 but not currently used or enabled (e.g., dormant application functionality  
1420 or ports), including assurance that this functionality cannot be activated  
1421 and/or misused;
- 1422 v. Access control models or features (if any) for every asset (such as  
1423 privileges, user accounts/groups, passwords);
- 1424 vi. Users' roles and levels of responsibility if they interact with the assets and  
1425 communication channels.
- 1426 vii. Any "handoff" sequences from one communication path to another (e.g.,  
1427 from asset to asset, network to network, or Bluetooth to Wi-Fi), and how  
1428 the data, code, and/or commands are secured/protected during handoff  
1429 (i.e., how is their integrity/authenticity assured);
- 1430 viii. Explanations of intended behavior in unusual/erroneous/unexpected  
1431 circumstances (e.g., termination of a connection in the middle of a data  
1432 transfer);
- 1433 ix. Authentication mechanism (if any), including the algorithm name/version  
1434 (if available), "strength" indicators (e.g., key bit length, number of  
1435 computational rounds) and mode of operation (if applicable);
- 1436 x. Descriptions of the cryptographic method used and the type and level of  
1437 cryptographic key usage and their style of use throughout the system (e.g.,  
1438 one-time use, key length, the standard employed, symmetric or otherwise).  
1439 Descriptions should also include details of cryptographic protection for  
1440 firmware and software updates;
- 1441 xi. Detailed analyses by cryptography experts if a cryptography algorithm is  
1442 proprietary, or a proprietary modification of a standard algorithm;

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- 1443           xii.    For each authenticator created, a list of where it is verified, and how  
1444           verification credentials (e.g., certificates, asymmetric keys, or shared keys)  
1445           are distributed to both endpoints;  
1446           xiii.   A precise, detailed list of how each type of credential (e.g., password, key)  
1447           is generated, stored, configured, transferred, and maintained, including  
1448           both manufacturer- and health care facility-controlled assets (e.g., key  
1449           management and public key infrastructure (PKI));  
1450           xiv.    Identity management<sup>65</sup> (if any), including how identities are  
1451           managed/transferred and configured (e.g., from manufacturer to  
1452           programmer and from programmer to device);  
1453           xv.    If communication sessions are used or supported, a detailed explanation of  
1454           how sessions are established, maintained, and broken down, including but  
1455           not limited to assurances of security properties such as uniqueness,  
1456           unpredictability, time-stamping, and verification of session identifiers;  
1457           xvi.   Precise links between diagram elements (or explanatory text), associated  
1458           hazards and controls, and testing;  
1459           xvii.   Explanations or links to the evidence that may be used to justify security  
1460           claims and any assumptions; and  
1461           xviii.   Traceability to the SBOM described in section V.B.2, above, for  
1462           proprietary and third-party code.  
1463

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<sup>65</sup> For the purposes of this guidance, “identity management” means the process that governs the authentication and authorization of users to devices and assets.

1464 **Appendix 3. Submission Documentation for Investigational**  
1465 **Device Exemptions**

1466  
1467 FDA acknowledges the need to balance innovation and security in designs especially during  
1468 clinical trials. In order to ensure security is addressed early in the device design, FDA has  
1469 identified a subset of the documentation recommended throughout this guidance to submit with  
1470 IDE applications.

1471  
1472 Under 21 CFR 812.25, manufacturers must provide an investigational plan as a part of their IDE  
1473 application. For devices within the scope of this guidance, FDA recommends that this  
1474 investigational plan include information on the cybersecurity of the subject device.

1475  
1476 Specifically, FDA recommends the following documentation be included as part of IDE  
1477 applications:

- 1478 • Inclusion of cybersecurity risks as part of Informed Consent Form (21 CFR 50.25(a)(2)  
1479 and 21 CFR 812.25(g));
- 1480 • Global, Multi-patient and Updateability/Patchability views (21 CFR 812.25(c), (d))
- 1481 • Security Use case views for functionality with safety risks (e.g., implant programming)  
1482 (21 CFR 812.25(c), (d));
- 1483 • Software Bill of Materials (21 CFR 812.25(c), (d)); and
- 1484 • General Labeling – Connectivity and associated general cybersecurity risks,  
1485 updateability/process (21 CFR 812.25(f)).

1486  
1487 FDA intends to review this information in the context of the overall benefit-risk assessment of  
1488 investigational devices as outlined in [Factors to Consider When Making Benefit-Risk](#)  
1489 [Determinations for Medical Device Investigational Device Exemptions](#).<sup>66</sup> Therefore, approval of  
1490 an IDE based on the documentation recommended above does not preclude the possibility of  
1491 future cybersecurity questions or concerns being raised during review of a subsequent marketing  
1492 application. This is, in part, due to the understanding that design changes may be needed and the  
1493 temporal nature of security. Security improvements will likely be needed between the time of  
1494 clinical trials and the device submitted for marketing authorization (e.g., operating system no  
1495 longer supported or nearing end of support, third party software updates, etc.).

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<sup>66</sup> See FDA Guidance “Factors to Consider When Making Benefit-Risk Determinations for Medical Device Investigational Device Exemptions” available at: <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/factors-consider-when-making-benefit-risk-determinations-medical-device-investigational-device>.

1497 **Appendix 4. Terminology**

1498 The terminology listed here are for the purposes of this guidance and are intended for use in the  
1499 context of assessing medical device cybersecurity. These terms are not intended to be applied in  
1500 any context beyond this guidance.

1501  
1502 **Asset** – anything that has value to an individual or an organization.<sup>67</sup>  
1503

1504 **Authentication** – the act of verifying the identity of a user, process, or device as a prerequisite to  
1505 allowing access to the device, its data, information, or systems, or provision of assurance that a  
1506 claimed characteristic of an entity is correct.<sup>68</sup>  
1507

1508 **Authenticity** – information, hardware, or software having the property of being genuine and  
1509 being able to be verified and trusted; confidence that the contents of a message originates from  
1510 the expected party and has not been modified during transmission or storage.<sup>69</sup>  
1511

1512 **Authorization** – the right or a permission that is granted to a system entity to access a system  
1513 resource.<sup>70</sup>  
1514

1515 **Availability** – the property of data, information, and information systems to be accessible and  
1516 usable on a timely basis in the expected manner (i.e., the assurance that information will be  
1517 available when needed).<sup>71</sup>  
1518

1519 **Compensating Controls** – a safeguard or countermeasure deployed, in lieu of, or in the absence  
1520 of controls designed in by a device manufacturer. These controls are external to the device  
1521 design, configurable in the field, employed by a user, and provide supplementary or comparable  
1522 cyber protection for a medical device.<sup>72</sup>  
1523

1524 **Confidentiality** – the property of data, information, or system structures to be accessible only to  
1525 authorized persons and entities and are processed at authorized times and in the authorized  
1526 manner, thereby helping ensure data and system security. Confidentiality provides the assurance

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<sup>67</sup> Definition is adapted from ISO/IEC 27032 Information technology — Security techniques — Guidelines for cybersecurity, clause 4.6.

<sup>68</sup> Definition is adapted from NIST FIPS 200 Minimum Security Requirements for Federal Information and Information Systems and from ISO/IEC 18014-2:2009(E) Information technology – Security techniques - Time-stamping Services - Part 2: Mechanisms producing independent tokens, clause 3.

<sup>69</sup> Adapted from NIST SP 800-53 Security and Privacy Controls for Federal Information Systems and Organizations: Authenticity is defined as “the property of being genuine and being able to be verified and trusted; confidence in the validity of a transmission, a message, or message originator. See Authentication.”

<sup>70</sup> Definition is adapted from CNSSI 4009-2015 Committee on National Security Systems (CNSS) Glossary.

<sup>71</sup> [ISO IEC 27000-2018, Clause 3.7: The property of being accessible and useful on demand by an authorized entity].

Definition is adapted from CNSSI 4009-2015 Committee on National Security Systems (CNSS) Glossary.

<sup>72</sup> Definition is adapted from NIST Special Publication “Assessing Security and Privacy Controls in Federal Information Systems and Organizations,” NIST SP 800-53A Rev. 4.

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1527 that no unauthorized users (i.e., only trusted users) have access to the data, information, or  
1528 system structures.<sup>73</sup>

1529  
1530 **Configuration** – the possible conditions, parameters, and specifications with which a device or  
1531 system component can be described or arranged.<sup>74</sup>

1532  
1533 **Configuration Management** - a collection of activities focused on establishing and maintaining  
1534 the integrity of information technology products and information systems, through control of  
1535 processes for initializing, changing, and monitoring the configurations of those products and  
1536 systems throughout the system development lifecycle.<sup>75</sup>

1537  
1538 **Cryptography** – the discipline that embodies the principles, means, and methods for providing  
1539 information security; including confidentiality, data integrity, non-repudiation, and  
1540 authenticity.<sup>76</sup>

1541  
1542 **Cybersecurity** – the process of preventing unauthorized access, modification, misuse or denial  
1543 of use, or the unauthorized use of information that is stored, accessed, or transferred from a  
1544 medical device to an external recipient.<sup>77</sup>

1545  
1546 **Decommission** – a process in the disposition process that includes proper identification,  
1547 authorization for disposition, and sanitization of the equipment, as well as removal of Patient  
1548 Health Information (PHI) or software, or both.<sup>78</sup>

1549  
1550 **Decryption** – is the cryptographic transformation of encrypted data (called “ciphertext”) into  
1551 non-encrypted form (called “plaintext”).<sup>79</sup>

1552  
1553 **Disposal** – a process to end the existence of a system asset or system for a specified intended  
1554 use, appropriately handle replaced or retired assets, and to properly attend to identified critical  
1555 disposal needs (e.g., per an agreement, per organizational policy, or for environmental, legal,  
1556 safety, security aspects).<sup>80</sup>

1557  
1558 **Encryption** – is the cryptographic transformation of data (called “plaintext”) into a form (called  
1559 “ciphertext”) that conceals the data’s original meaning to prevent it from being known or used.<sup>81</sup>

1560

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<sup>73</sup> Definition is adapted from ISO IEC 27000-2018, Clause 3.10: Property that information is not made available or disclosed to unauthorized individuals, entities, or processes.

<sup>74</sup> Adapted Definition is adapted from NIST SP 800-128 Guide for Security-Focused Configuration Management of Information Systems: Configuration is

<sup>75</sup> Definition is adapted from NIST SP 800-53 Rev. 4.

<sup>76</sup> Definition is adapted from CNSSI 4009-2015 (NIST SP 800-21 Second edition).

<sup>77</sup> Definition is adapted from ISO IEC 27032: 2012, Clause 4.20.

<sup>78</sup> Definition is adapted from Medical Device and Health IT Joint Security Plan (JSP). Available at <https://healthsectorcouncil.org/the-joint-security-plan/>.

<sup>79</sup> Definition is referenced from NIST SP 800-82 Guide to Industrial Control Systems (ICS) Security.

<sup>80</sup> Definition is adapted from 6.4.14.1 Disposal process purpose ISO/IEC/IEEE 12207:2017(E).

<sup>81</sup> Definition is referenced from NIST SP 800-82 Guide to Industrial Control Systems (ICS) Security.

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- 1561 **End of support** – a point beyond which the product manufacturer ceases to provide support,  
1562 which may include cybersecurity support, for a product or service.  
1563
- 1564 **Exploitability** – the feasibility or ease and technical means by which the vulnerability can be  
1565 exploited by a threat.<sup>82</sup>  
1566
- 1567 **Firmware** – software program or set of instructions programmed on the flash read-only memory  
1568 (ROM) of a hardware device. It provides the necessary instructions for how the device  
1569 communicates with the other computer hardware.<sup>83</sup>  
1570
- 1571 **Hardening** – a process intended to eliminate a means of attack by patching vulnerabilities and  
1572 turning off nonessential services.<sup>84</sup>  
1573
- 1574 **Hardware** –
- 1575
- 1576 **Integrity** – the property of data, information and software to be accurate and complete and have  
1577 not been improperly or maliciously modified.<sup>86</sup>  
1578
- 1579 **Lifecycle** – all phases in the life of a medical device, from initial conception to final  
1580 decommissioning and disposal.<sup>87</sup>  
1581
- 1582 **Malware** – software or firmware intended to perform an unauthorized process that will have  
1583 adverse impact on the confidentiality, integrity, or availability of an information system.<sup>88</sup>  
1584
- 1585 **Patch** – a “repair job” for a piece of programming; also known as a “fix”. A patch is the  
1586 immediate solution to an identified problem that is provided to users. The patch is not necessarily  
1587 the best solution for the problem, and the product developers often find a better solution to  
1588 provide when they package the product for its next release. A patch is usually developed and  
1589 distributed as a replacement for or an insertion in compiled code (that is, in a binary file or object  
1590 module). In many operating systems, a special program is provided to manage and track the  
1591 installation of patches.<sup>89</sup>  
1592
- 1593 **Patient harm** – injury or damage to the health of patients, including death.<sup>90</sup>  
1594
- 1595 **Programmable logic** – hardware that has undefined function at the time of manufacture and  
1596 must be programmed with software to function (e.g., Field-programmable gate array)

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<sup>82</sup> The definition is adapted from the Common Vulnerability Scoring System (CVSS) specification document (v3.1).

<sup>83</sup> Definition is adapted from NISTIR 8183. <https://nvlpubs.nist.gov/nistpubs/ir/2017/NIST.IR.8183.pdf>

<sup>84</sup> Definition is referenced from NIST SP 800-152.

<sup>85</sup> Definition is referenced from CNSSI 4009-2015 \_\_\_\_\_

<sup>86</sup> Definition is adapted from AAMI TIR 57 Clause 2.15.

<sup>87</sup> Definition is referenced from ANSI/AAMI/ISO 14971 Medical Devices – Application of Risk Management to Medical Devices, clause 2.7.

<sup>88</sup> Definition is referenced from NIST SP 800-53 Rev. 4.

<sup>89</sup> Definition is adapted from NIST SP 800-45 Version 2.

<sup>90</sup> Patient harm from cybersecurity risks is discussed at length throughout this guidance and the FDA Guidance “Postmarket Management of Cybersecurity in Medical Devices” issued December 2016. See Footnote 6.



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1597

1598 **Resilience** – the ability of an information system to continue to: (i) operate under adverse  
1599 conditions or stress, even if in a degraded or debilitated state, while maintaining essential  
1600 operational capabilities; and (ii) recover to an effective operational posture in a time frame  
1601 consistent with mission needs.<sup>91</sup>

1602

1603 **Secure Product Development Framework (SPDF)** - a set of processes that reduce the number  
1604 and severity of vulnerabilities in products. Additional information about an SPDF and its  
1605 implementation is discussed in Section IV.C. and throughout the guidance.

1606

1607 **Security Architecture** – a set of physical and logical security-relevant representations (i.e.,  
1608 views) of system architecture that conveys information about how the system is partitioned into  
1609 security domains and makes use of security-relevant elements to enforce security policies within  
1610 and between security domains based on how data and information must be protected. The  
1611 security architecture reflects security domains, the placement of security-relevant elements  
1612 within the security domains, the interconnections and trust relationships between the security-  
1613 relevant elements, and the behavior and interactions between the security-relevant elements.<sup>92</sup>

1614

1615 **Security Strength** –

1616

1617

1618

1619

1620 **Security Risk Management** – a process (or processes) that evaluates and controls threat-based  
1621 risks. For security risk management, this includes an evaluation of the impact of exploitation on  
1622 the device’s safety and effectiveness, the exploitability, and the severity of patient harm if exploited.

1623

1624 **Software Bill of Materials (SBOM)** – a list of software components that includes but is not  
1625 limited to commercial, open source, off-the-shelf, and custom software components. See Section  
1626 V.A.2 for a more complete description of an SBOM.

1627

1628 **System** – the combination of interacting elements or assets organized to achieve one or  
1629 more function.<sup>94</sup>

1630

1631 **Threat** – Threat is any circumstance or event with the potential to adversely impact the device,  
1632 organizational operations (including mission, functions, image, or reputation), organizational  
1633 assets, individuals, or other organizations through an information system via unauthorized  
1634 access, destruction, disclosure, modification of information, and/or denial of service. Threats  
1635 exercise vulnerabilities, which may impact the safety or effectiveness of the device.<sup>95</sup>

1636

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<sup>91</sup> As defined in NISTSP 800-53 Rev. 4 definition of Information System Resilience.

<sup>92</sup> Definition is referenced from NIST 800-160v1, Systems Security Engineering.

<sup>93</sup> Definition is referenced from NIST SP 800-108

<sup>94</sup> Definition is adapted from ISO/IEC/IEEE 12207:2017.

<sup>95</sup> Definition is adapted from NIST SP 800-53.

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1637 **Threat modeling** – a methodology for optimizing system, product, network, application, and  
1638 connection security by identifying objectives and vulnerabilities, and then defining  
1639 countermeasures to prevent, or mitigate the effects of, threats to the system.<sup>96</sup>  
1640

1641 **Trustworthy Device** – a medical device that: (1) is reasonably secure from cybersecurity  
1642 intrusion and misuse; (2) provides a reasonable level of availability and reliability; (3) is  
1643 reasonably suited to performing its intended functions; and (4) adheres to generally accepted  
1644 security procedures to support correct operation.<sup>97</sup>  
1645

1646 **Updatability and Patchability** – the ease and timeliness with which a device and related assets  
1647 can be changed for any reason (e.g., feature update, security patch, hardware replacement).  
1648

1649 **Update** –corrective, preventative, adaptive, or perfective modifications made to software of a  
1650 medical device.<sup>98</sup>  
1651

1652 **Vulnerability** - a weakness in an information system, system security procedure(s), internal  
1653 control(s), human behavior, or implementation that could be exploited.

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<sup>96</sup> Definition is adapted from CNSSI 4009-2015 (NIST SP 800-21 Second edition).

<sup>97</sup> Definition is adapted from NIST SP 800-32 Introduction to Public Key Technology and the Federal PKI Infrastructure.

<sup>98</sup> Definition is from IMDRF Guidance “Principles and Practices for Medical Device Cybersecurity” available at <http://www.imdrf.org/docs/imdrf/final/technical/imdrf-tech-200318-pp-mdc-n60.pdf>.