

1 **Factors to Consider Regarding Benefit-**  
2 **Risk in Medical Device Product**  
3 **Availability, Compliance, and**  
4 **Enforcement Decisions**

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6 **Draft Guidance for Industry and**  
7 **Food and Drug Administration Staff**

8 *DRAFT GUIDANCE*

9 **This draft guidance document is being distributed for comment purposes only.**

10 **Document issued on June 16, 2016.**

11 You should submit comments and suggestions regarding this draft document within 90 days of  
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13 Submit electronic comments to <http://www.regulations.gov>. Submit written comments to the  
14 Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane,  
15 rm. 1061, Rockville, MD 20852. Identify all comments with the docket number listed in the notice  
16 of availability that publishes in the *Federal Register*.

17 For questions about this document regarding CDRH-regulated devices, contact the Office of  
18 Compliance at 301-796-5900.



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

# Preface

36

37

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43

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73 **Draft Guidance for Industry and**  
74 **Food and Drug Administration Staff**

76 *This draft guidance, when finalized, will represent the current thinking of the Food and Drug*  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

81 **I. Introduction**

82 FDA has developed this draft guidance document to provide clarity for FDA staff and industry  
83 regarding the benefit and risk factors FDA may consider in prioritizing resources for compliance  
84 and enforcement efforts to maximize medical device quality<sup>1</sup> and patient safety. This draft  
85 guidance is not intended to limit FDA action; rather, it describes the general framework for  
86 medical device decision making in the product availability, compliance, and enforcement arenas.  
87 Product availability and other medical device compliance and enforcement decisions are generally  
88 fact-specific. However, FDA believes that explaining how we consider the factors listed in this  
89 draft guidance document will improve the consistency and transparency of these kinds of  
90 decisions. A common understanding of how FDA considers benefit and risk may better align  
91 industry's and FDA's focus on actions that maximize benefit to patients, improve medical device  
92 quality, and reduce risk to patients.

93  
94 This draft guidance, when finalized, is intended to provide a framework for FDA and stakeholders  
95 that sets forth overarching benefit-risk principles. FDA may consider the types of benefit-risk  
96 factors described in this draft guidance—including reliable patient preference information from a  
97 representative sample—on a case-by-case basis when determining the appropriate actions to take  
98 and to help ensure that informed and science-based decisions are made to the greatest extent

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<sup>1</sup> “Quality means the totality of features and characteristics that bear on the ability of a device to satisfy fitness-for-use, including safety and performance.” 21 CFR 820.3(s).

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99 practicable. Factors may be weighted differently for different types of decisions and as the  
100 timeframe allows. FDA intends to use pilots to help determine how to apply the benefit-risk  
101 framework described in this draft guidance.

102  
103 In addition, this draft guidance, when finalized, is intended to harmonize FDA’s approach to  
104 weighing benefits and risks for medical device product availability, compliance, and enforcement  
105 decisions with FDA’s benefit-risk framework for assessing medical device marketing and  
106 investigational device exemption (IDE) applications. The benefit-risk factors in this draft guidance  
107 also support assessment of medical devices with real world evidence. While the benefit-risk factors  
108 in this draft guidance are not identical to the other frameworks, this draft guidance builds upon  
109 FDA’s premarket review benefit-risk policy in an effort to improve consistency in our patient  
110 centered approach and decision making across the total product life cycle. This draft guidance is  
111 intended to complement, not supplant, FDA’s “[Guidance for Industry and Food and Drug  
112 Administration Staff - Factors to Consider When Making Benefit-Risk Determinations in Medical  
113 Device Premarket Approvals and De Novo Classifications.](#)”

114  
115 For the current edition of the FDA-recognized standard(s) referenced in this document, see the  
116 FDA Recognized Consensus Standards Database Web site at  
117 <http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/search.cfm>.

118  
119 FDA’s guidance documents, including this one, do not establish legally enforceable  
120 responsibilities. Instead, guidance documents describe the Agency’s current thinking on a topic and  
121 should be viewed only as recommendations, unless specific regulatory or statutory requirements  
122 are cited. The use of the word *should* in Agency guidance documents means that something is  
123 suggested or recommended, but not required.

## 124 **II. Scope**

125 The framework described in this draft guidance may be applicable to both industry and FDA  
126 decisions. The benefit-risk factors may be considered when device manufacturers evaluate  
127 appropriate responses to nonconforming product or regulatory compliance issues, such as  
128 determining whether to limit the availability of a medical device (e.g., a voluntary recall or market  
129 withdrawal). FDA may consider the benefit-risk factors during, for example, evaluation of device  
130 shortage situations, selection of the appropriate regulatory engagement mechanism following an  
131 inspection during which regulatory non-compliance was observed, evaluation of recalls, and  
132 consideration of petitions for variance from those sections of the Quality System (QS) regulation  
133 (21 CFR part 820) for which there were inspectional observations during a Premarket Approval  
134 (PMA) pre-approval inspection. Premarket submission review decisions, such as premarket  
135 notification (510(k)) substantial equivalence determinations, de novo classification, and PMA,  
136 Humanitarian Device Exemption (HDE) or IDE application approval decisions, are beyond the  
137 scope of this draft guidance.

138  
139 Because of the potentially direct effect on patients, medical device compliance and enforcement  
140 decisions that affect product availability should generally include consideration of specific factors.  
141 The factors described in this document can apply to many situations where the Agency or  
142 manufacturer has information that leads to quality, compliance, or other concerns regarding a  
143 medical device and considers taking action that could have a direct effect on the device’s

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144 availability. These situations may include information about new risks or about known risks  
145 occurring at greater than expected frequency or severity. To support a common understanding of  
146 other kinds of compliance and enforcement decision making, the factors in this draft guidance may  
147 also be considered when the Agency or manufacturer considers taking action that is unlikely to  
148 directly affect product availability but seeks to minimize risks to patients associated with  
149 manufacturer quality and regulatory compliance issues (e.g., issues in design, manufacturing, or  
150 reporting related to the device), while also considering the benefits patients may receive from the  
151 device. The intersection of this draft guidance with ISO 14971: *Medical devices – Application of*  
152 *risk management to medical devices*<sup>2</sup> is discussed in Appendix A.

153  
154 This draft guidance applies to both diagnostic and therapeutic medical devices subject to, and  
155 exempt from, premarket review. The scope of this draft guidance excludes medical devices  
156 regulated by FDA’s Center for Biologics Evaluation and Research (CBER); combination products,  
157 as defined in 21 CFR 3.2(e), for which CDRH is not the lead Center; and electronic products that  
158 are not devices as defined in section 201(h) of the Federal Food, Drug, and Cosmetic Act (FD&C  
159 Act), as regulated by CDRH under the Electronic Product Radiation Control (ERPC) provisions in  
160 the FD&C Act and implementing regulations (21 CFR Subchapter J–Radiological Health). This  
161 draft guidance also does not apply to products (e.g., drugs, biologics, dietary supplements, foods,  
162 tobacco products, or cosmetics) regulated by other FDA Centers.

163  
164 Guidance documents, including this draft guidance, are not binding, and the concepts and factors  
165 described herein generally explain how benefit-risk assessments can be made. This draft guidance  
166 does not preclude FDA from taking regulatory or other action in response to a violation of  
167 applicable law or regulation.

### 168 **III. Patient<sup>3</sup> Focused Benefit-Risk Assessments for Medical** 169 **Device Product Availability, Compliance, and** 170 **Enforcement Decisions**

171 FDA has authority to limit the availability of violative medical devices and to pursue other  
172 compliance and enforcement actions related to violative medical devices. FDA recognizes that, to  
173 achieve the Agency’s goal of protecting and promoting the public health, decisions regarding these  
174 actions should be made while focusing on the impact on patients. Failure to consider the short-term  
175 and long-term impact of non-compliance on the benefit-risk profile of the device and the benefit-  
176 risk tradeoffs of FDA’s decision options on the health and quality of life of patients could result in  
177 regulatory actions with unintended adverse effects (e.g., shortage of medically necessary devices).

178  
179 In certain situations involving risks of patient harm, FDA and industry, individually or  
180 collaboratively, can help maximize benefit and reduce risk to patients by assessing the situation,  
181 considering patients’ perspectives, evaluating any regulatory non-compliance or device

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<sup>2</sup> For the current edition of the FDA-recognized standard(s) referenced in this document, see the FDA Recognized Consensus Standards Database Web site at <http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/search.cfm>.

<sup>3</sup> Although this draft guidance focuses on patients, when relevant, the benefit-risk factors also take into account benefits or risks for non-patient users of medical devices, such as healthcare providers and caregivers.

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182 nonconformity in light of the benefit-risk profile of the device,<sup>4</sup> factoring in alternatives, where  
183 available, considering the benefit-risk tradeoffs for patients of each decision option, and  
184 determining the most appropriate next steps.

185

186 When FDA is working with a manufacturer to address a failure to comply with applicable statutes  
187 or regulations, observed unanticipated harm to patients or users, or identified device  
188 nonconformities, FDA strives to be clear with that manufacturer about the benefits and risks the  
189 Agency is considering. As with premarket review decisions, when making medical device product  
190 availability, compliance, and enforcement decisions informed by benefit-risk, FDA may consider  
191 relevant, reliable information relating to patient perspectives on what constitutes meaningful  
192 benefit, what constitutes risk, and what tradeoffs patients are willing to accept, if such information  
193 is available at the time of decision, as well as what alternatives are available. Before arriving at a  
194 decision, FDA may also consider the manufacturer’s approach to minimize harm or to mitigate the  
195 increased risks that result from regulatory non-compliance or nonconformity of the product, their  
196 compliance history, and the scope of the issue.

197

198 By providing greater clarity about the factors we consider, we intend to improve consistency and  
199 transparency and to better align industry’s and FDA’s focus on actions that maximize benefit to  
200 patients, improve medical device quality, and reduce risk to patients. In Appendices B, C, and D,  
201 draft benefit-risk assessment worksheets have been provided to support consideration of the factors  
202 by FDA staff and industry.

203

204 Note that as with premarket benefit-risk determinations made when evaluating marketing and IDE  
205 applications, benefit-risk assessments made in product availability, compliance, and enforcement  
206 contexts may change over time. For example, as the practice of medicine evolves, clinical  
207 experience increases, or additional treatment options become available to patients, a benefit-risk  
208 conclusion may change.

209 **IV. Description of Factors to Consider Regarding Benefit-**  
210 **Risk for Medical Device Product Availability,**  
211 **Compliance, and Enforcement Decisions**

212 In assessing benefit-risk factors for purposes of medical device product availability, compliance,  
213 and enforcement decisions, FDA considers relevant and reliable evidence and data available to the  
214 Agency at the time of a decision—including reliable patient preference information from a  
215 representative sample— on a case-by-case basis, to help ensure that informed and science-based  
216 decisions are made to the greatest extent practicable. FDA may use available evidence or request  
217 data to assess these factors, as appropriate. The benefit-risk assessments covered in this draft  
218 guidance document may compare the benefits and risks identified based on currently available

---

<sup>4</sup> “*Nonconformity* means the nonfulfillment of a specified requirement.” 21 CFR 820.3(q).

In the preamble to the final rule for the QS regulation, “FDA emphasizes that a ‘nonconformity’ may not always rise to the level of a product defect or failure, but a product defect or failure will typically constitute a nonconformity.” (61 FR 52610.)

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219 information to those benefits and risks that were identified during premarket review benefit-risk  
220 assessments (or early risk assessments documented as part of a manufacturer’s risk management  
221 process) in order to understand whether there has been a change in the benefit-risk analysis over  
222 time. We generally consider our device benefit-risk assessment along with data/information related  
223 to benefit-risk factors outlined in the draft guidance to reach our judgment about how to proceed in  
224 each situation.

225

226 **A. Factors for the Assessment of Medical Device Benefits**

227 When prioritizing compliance and enforcement efforts to maximize medical device quality and  
228 patient safety, FDA may assess the extent of benefit of a device by considering factors such as  
229 those listed below. The following factors, when relevant, should be considered in the aggregate.  
230 The factors may be considered early in the medical device product life cycle and reassessed as the  
231 device is used more widely.

232

233 Benefit, as described by the potential benefit factors, may change over time. The text below  
234 describes each factor for the purposes of this draft guidance and provides examples of how each  
235 factor may be considered.

236

237 **Type of benefit(s)** includes, but is not limited to, the medical device’s impact on patient health and  
238 clinical management. Examples include the effect of the device on patient treatment plans and  
239 quality of life; impact on survival; and how much the medical device can aid in improving patient  
240 function, preventing loss of function, or providing relief from the symptoms of the disease or  
241 condition that the medical device is intended to treat.

242

243 As a medical device is used, clinicians may find unanticipated ways to use the medical device and  
244 additional types of benefit. For example, a surgical tool may be cleared for use in hernia repair  
245 surgery. Surgeons may find additional uses for the surgical tool that may lead to clearance of new  
246 uses, thus increasing the types of benefit.

247

248 **Magnitude of benefit(s)** is the degree to which patients experience the treatment benefit or the  
249 effectiveness of the medical device. The change in patients’ conditions or the change in necessary  
250 clinical management may allow FDA to determine the magnitude of the benefit. Magnitude of  
251 benefit may be assessed against standards of care and expected performance and may change over  
252 time.

253

254 **Likelihood of patients experiencing one or more benefits** is the likelihood that the medical  
255 device will effectively treat or diagnose the patient’s disease or condition. A medical device may  
256 not provide effective treatment or diagnosis for all patients. One method of determining the  
257 likelihood of benefit, for a particular patient population, is to determine the number of patients  
258 treated effectively and divide this by the total number of patients treated.

259

260 In assessing benefit, FDA may consider whether there are subpopulations included in the  
261 indication for use that are more likely to retain expected benefits than the overall population. If  
262 the subgroups can be identified, the likelihood of those patients experiencing benefit from the  
263 device may increase. The benefit for a subpopulation may also be greater than for the population  
264 as a whole, and this greater benefit should be considered in the overall benefit-risk assessment.

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265

266 **Duration of effects** is how long the benefit can be expected to last for the patient. Curative  
267 treatments may be seen as providing higher benefit because of a longer duration of effect.

268

269 Knowledge of the duration of treatment effect may change as the medical device is used. For  
270 example, a medical device may have been approved with clinical endpoint data demonstrating  
271 effectiveness for 6 months. As the medical device is used, patients may experience significantly  
272 longer treatment effects than those described in the device labeling.

273

274 **Patient preference on benefit** is the value that patients place on use of the medical device.  
275 Faced with a severe or chronic disease, a patient may highly value the benefit provided by a  
276 medical device in light of the specific condition that patient has. For example, patients dying of  
277 congestive heart failure may highly value a medical device that extends their lives for a few  
278 months. Patients with less severe or chronic diseases may or may not place the same value on a  
279 device with a short-term benefit.

280

281 **Benefit factors for healthcare professionals or caregivers** include the benefit that healthcare  
282 professionals or caregivers experience by improving the way they care for patients, whether this  
283 directly improves patient outcomes or improves clinical practice. FDA recognizes that certain  
284 devices, such as surgical tools that allow different techniques or devices that positively affect  
285 ongoing patient management, may improve the benefit profile.

286

287 **Medical necessity** should be considered if a medical device provides benefits or addresses needs  
288 unmet by other medical devices or therapies. Benefit considerations should include an  
289 assessment of whether another medical device or therapy could be used in substitution, and the  
290 availability of that other medical device or therapy.

291

292

## **B. Factors for the Assessment of Medical Device Risks**

293

294 When prioritizing compliance and enforcement efforts to maximize medical device quality and  
295 patient safety, FDA may assess the risk that a medical device will cause patient direct or indirect  
296 harm by considering factors such as those listed below. The following factors, when relevant,  
297 should be considered in aggregate. Each factor may be considered early in the medical device  
298 product life cycle and reassessed as the device is used more widely. Changes in risk should be  
299 noted in the manufacturer's risk management documentation. Changes in risk may occur due to,  
300 among other things, observed unanticipated harm to patients exposed to the device or to device  
301 users, changes in the medical device use environment, identified medical device nonconformities,  
302 and issues related to the design or manufacturing of the device. It should be noted that all devices  
303 have some level of anticipated risk, even without device nonconformities or regulatory non-  
304 compliance.

305

306 Medical device nonconformities may directly increase risk or introduce new risks. Failure to  
307 comply with applicable statutes or regulations also may be a negative indicator of a manufacturer's  
308 ability to consistently manufacture high quality medical devices, even if a device made by such a  
309 manufacturer still performs as expected. Postmarket data may also show that risk is higher than  
310 anticipated, even in the absence of a medical device nonconformity or regulatory non-compliance.  
Therefore, the risk factors listed below take into account considerations related to nonconforming

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311 devices, failure to comply with applicable statutes or regulations, and harm potentially unrelated to  
312 compliance with legal requirements or device nonconformities.

313

314 Risk, as described by the potential risk factors, may change over time. The text below describes  
315 each risk factor for purposes of this draft guidance and provides examples of how the risk factor  
316 may be considered.

317

318 **Risk severity** is categorized into three levels and includes a duration component. The three  
319 levels are medical device-related deaths or serious injuries, medical device-related non-serious  
320 adverse events, and medical device-related events without reported harm.

321

322 Medical device-related deaths and serious injuries include those events (including  
323 procedure related complications) that may have been or were attributed to the use of the  
324 medical device and that cause or contribute to a death or injury or illness that is life-  
325 threatening, results in permanent impairment or damage to the body, or requires medical or  
326 surgical intervention to prevent permanent harm to the body.

327

328 Medical device-related non-serious adverse events include those events (including  
329 procedure related complications) that may have been or were attributed to the use of the  
330 medical device and that cause or contribute to minor, temporary or medically reversible  
331 injuries that do not meet the criteria for classification as a medical device-related serious  
332 injury.

333

334 Medical device-related events without reported harm can include medical device  
335 nonconformities which have no related harm, medical device malfunctions which have no  
336 related harm, procedure related complications with no related harm, and instances where a  
337 nonconformity or regulatory noncompliance was observed at the medical device  
338 manufacturing facility and no defective devices were released to the market. A medical  
339 device nonconformity can include the failure of a medical device to meet its performance  
340 specifications even though the device still performs adequately to meet the needs of a  
341 given patient.

342

343 Duration of harm to patient - Depending on circumstance, medical devices can cause harm  
344 to patients that is temporary, repeated but reversible, or permanent.

345

346 **Likelihood of risk** considers three risk factors related to the potential number of patients at risk of  
347 experiencing harm: the likelihood that a medical device will have problems, the likelihood of a  
348 patient experiencing harm, and the total number of patients exposed.

349

350 Likelihood of medical device nonconformity is the likelihood that the medical device will  
351 exhibit a specific failure mode or defect. Regulatory non-compliance may increase the  
352 likelihood of a medical device nonconformity. One method of calculating the likelihood of  
353 medical device nonconformity is to identify the number of nonconforming medical devices  
354 and divide by the total number of medical devices manufactured, under the same conditions.

355

356 Likelihood of a harmful event given exposure to a nonconforming device is the proportion of  
357 the intended population treated with or diagnosed by the nonconforming medical device that

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358 would be expected to experience a harmful event if exposed to a nonconforming device. One  
359 method to calculate this likelihood is to take the number of patients treated with a  
360 nonconforming medical device and harmed and to divide by the total number of patients treated  
361 with nonconforming devices, over a similar time period, if reliable data exist.

362  
363 The likelihood of a harmful event given exposure to a nonconforming device should be  
364 compared to the likelihood of a harmful event given exposure to a conforming device.

365  
366 Number of patients exposed is the number of patients exposed to a nonconforming medical  
367 device or to a medical device manufactured by a noncompliant manufacturer.

368  
369 **Nonconforming product risks** include whether nonconforming product has been distributed and  
370 if so, how many nonconforming devices are on the market.

371  
372 **Duration of exposure to population** is the length of time between initial patient exposure to  
373 the device with the identified risk of harm and the point at which the risk of harm is  
374 successfully addressed.

375  
376 **False-positive or false-negative results** are important risk factors for diagnostics. If a diagnostic  
377 medical device gives a false-positive result, the patient might, for example, be incorrectly  
378 diagnosed with a serious disease and receive an unnecessary treatment, incurring all the risks that  
379 accompany that treatment. If a diagnostic medical device gives a false-negative result, the patient  
380 might not be diagnosed with the correct disease or condition and might not receive an effective  
381 treatment (thereby missing out on the benefits that treatment would confer). The risks associated  
382 with false positives and false negatives can be multifold, but are considered by FDA in light of  
383 probable risks.

384  
385 **Patient tolerance of risk** is the concern that patients have regarding harm or potential harm caused  
386 by the device. Patient tolerance of risk may take into account both the patients' willingness and  
387 unwillingness to use a nonconforming medical device, to use a device manufactured by a non-  
388 compliant manufacturer, or to tolerate harm (both probable and actual). Risk tolerance varies  
389 among patients, and affects individual patients' decisions as to whether risks associated with the  
390 medical device's technology are acceptable in exchange for the benefit. Risk tolerance may also  
391 vary with risk severity (e.g., there may be special subpopulations in which risk severity is higher).  
392 Patients may not understand device-related risks for all types of devices (e.g., lack of FDA review,  
393 certain diagnostics). For prescription devices, a patient's assessment of risk would be appropriately  
394 informed by information from his or her clinician.

395  
396 **Risk factors for healthcare professionals or caregivers** may be considered when the risk may  
397 have an adverse impact on the clinician or caregiver.

398  
399 **C. Additional Benefit-Risk Factors to Consider When Making**  
400 **Product Availability, Compliance, and Enforcement Decisions**

401 In addition to the benefit-risk factors described above, FDA may consider additional important  
402 benefit-risk factors related to product availability, compliance, and enforcement decisions, such as

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403 those listed below. The text below describes additional factors. Section V provides examples of  
404 how all the factors may be considered in specific situations.

405

406 **Uncertainty** is an important factor, since at any point in the total product life cycle, there is never  
407 100% certainty regarding the safety, effectiveness, or quality of a device. However, the degree of  
408 certainty of the benefits and risks of a device is a factor FDA considers when making benefit-risk  
409 assessments.

410

411 **Mitigations** are actions taken by the manufacturer, by FDA, or by other stakeholders to recover  
412 benefit, to limit risk from nonconforming product, to address underlying QS problems, or to limit  
413 harm. Mitigations could address, among other considerations, as applicable: clinical practice; use  
414 errors; unmet medical needs; the use environment; user population; user skill level; clinical  
415 understanding in assessing risk; current expectations in clinical use; any changes in medical  
416 practice, e.g., standard of care, that could increase risk; and use in emergency/crisis situations.

417

418 **Detectability** refers to whether a nonconformity could be identified, either by the  
419 manufacturer or by the user. A nonconformity which can be identified prior to use of the  
420 device may harm fewer patients than a nonconformity which is not identified prior to use. A  
421 detected nonconforming device may still cause patient harm (e.g., a mislabeled orthopedic  
422 implant may cause a delay in surgery). Time between exposure to a nonconforming device and  
423 symptoms can increase the frequency of harm because it can take longer to determine the  
424 cause of the harm, making it likely that patients will be exposed to the device in the  
425 intervening time.

426

427 **Failure mode** is the specific method or type of failure. The failure mode may be used to identify  
428 the cause of the nonconformance including whether the nonconformance is related to  
429 manufacturing, design, use conditions, or environment.

430

431 **Scope of the device issue** should be evaluated to assess whether the risks identified are potentially  
432 inherent to similar devices of this type (i.e., whether the risk is specific to a single device, a single  
433 manufacturer, or is industry wide).

434

435 **Patient impact** is the impact on the health and quality of life of patients if a particular compliance  
436 or enforcement action is, or is not, taken or if the device relevant to the nonconformity or  
437 regulatory non-compliance is not available. FDA and, where appropriate, industry should consider  
438 whether patients are better off if the device is or is not available.

439

440 **Preference for availability** relates to both the patient and the caregiver. FDA and industry, where  
441 appropriate, should understand whether patients and caregivers would prefer to have access to the  
442 device relevant to the nonconformity or regulatory non-compliance and whether patients and  
443 caregivers adequately understand related benefits and risks.

444

445 **Nature of violations/Nonconforming product** may include whether the violation was systemic or  
446 non-systemic in nature as well as the extent of any product nonconformity.

447

448 **Firm compliance history** may include the manufacturer's regulatory history and initiative in  
449 identifying and correcting issues, the repetitiveness of such issues, and the manufacturer's

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450 communication with FDA. When considering the firm’s compliance history, FDA may determine  
451 that it is appropriate to provide prior notice to the manufacturer as to what is required, what  
452 violations appear to exist, and, in the case of violations of regulatory significance, that failure to  
453 comply may result in the initiation of enforcement action.

454 **V. How FDA Considers Benefit-Risk in Patient Focused**  
455 **Medical Device Product Availability, Compliance, and**  
456 **Enforcement Decisions**

457 FDA may use a benefit-risk assessment to help the Agency make informed appropriate decisions.  
458 An FDA benefit-risk assessment for medical device product availability, compliance, and  
459 enforcement decisions begins with the existence of certain events, such as a recall, variance  
460 petition, safety signal, or medical device nonconformity, that may lead FDA to take regulatory  
461 action.

462  
463 FDA initiates a benefit-risk assessment by evaluating available benefit information on the  
464 applicable medical device and assessing the benefit information by considering the relevant benefit  
465 factors described in Section IV and in Appendix B – Worksheets for Benefit Assessments. Some  
466 potential sources of benefit information include literature, prior premarket submissions, clinical  
467 studies, registries, patient input, knowledgeable clinicians, and risk management documentation  
468 voluntarily supplied by the manufacturer.

469  
470 FDA would next assess the available risk information on the medical device and assesses the risk  
471 information by considering the relevant risk factors described in Section IV and Appendix C –  
472 Worksheets for Risk Assessments. Some potential sources of risk information include medical  
473 device reports (MDRs), inspection reports, literature, prior premarket submissions, clinical studies,  
474 registries, patient input, knowledgeable clinicians and risk management documentation voluntarily  
475 supplied by the manufacturer.

476  
477 FDA would complete the benefit-risk assessment by considering any factors from Appendix D that  
478 are relevant for assessing decision options.

479  
480 When appropriate, FDA would use the outcome of a benefit-risk assessment to inform decisions  
481 related to product availability. The types of product availability decisions where this may be useful  
482 include:

- 483
- 484 • When should a firm’s recall strategy appropriately include a correction instead of a  
485 removal?
  - 486 • What actions, if any, may FDA take when continued access to a nonconforming device or a  
487 device manufactured by a firm with regulatory compliance issues might be needed during a  
488 shortage situation?
  - 489 • When is it in the best interest of the public health to grant a variance from certain QS  
490 regulation requirements for QS issues identified during a PMA pre-approval inspection?
  - 491 • When might FDA exercise enforcement discretion and not take immediate action against a  
492 company for marketing a device with a significant change or modification prior to  
493 obtaining clearance, as required by 21 CFR 807.81(a)(3)?

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494  
495 Before making a decision that is likely to affect product availability, FDA may also consider the  
496 impact on the patient if the device is available or not available, whether the issue affects a single  
497 manufacturer or the whole industry, and patient or caregiver preference for availability. Specific  
498 benefit-risk assessments should be viewed in the larger context that includes consideration of the  
499 additional factors described in Section IV.C, but generally, if the benefit-risk assessment indicates  
500 high benefit to patients with little risk, FDA may be more likely to decide that it is appropriate for  
501 patients to have access to a nonconforming device while the long-term corrective action is taken if  
502 alternative treatments are not available. Alternatively, if the benefit-risk assessment indicates low  
503 benefit to patients with high risk, FDA would be more likely take action to limit product  
504 availability.

505  
506 In addition to compliance and enforcement decisions that potentially have a direct effect on  
507 product availability, when appropriate, FDA may use the outcome of a benefit-risk assessment to  
508 inform other decisions related to compliance and enforcement. Examples of the other types of  
509 compliance and enforcement decisions where this may be useful include:

- 510
- 511 • Is a manufacturer’s proposed correction strategy adequate given the benefit-risk
  - 512 assessments and mitigation activities?
  - 513 • Upon observing a violation, when might FDA send a Warning Letter or Untitled Letter and
  - 514 when would it be appropriate to take an alternative, more informal approach?
- 515

516 When making compliance and enforcement decisions that are unlikely to directly affect product  
517 availability, FDA may also consider whether regulatory non-compliance increases risk of harm to  
518 patients, whether taking (or not taking) a contemplated compliance or enforcement action would  
519 impact patients, the manufacturer’s regulatory history, and steps taken by the manufacturer to  
520 address the situation. Specific benefit-risk analyses will again need to be viewed in context, but  
521 generally, if FDA’s benefit-risk assessment indicates high benefit to patients with little risk, FDA  
522 may decide to work with the manufacturer to address the underlying issue without initiating a  
523 formal compliance or enforcement action. If FDA’s benefit-risk assessment indicates low benefit  
524 to patients with high risk, FDA would likely take formal compliance or enforcement action to  
525 address the problem.

526 **VI. Examples Demonstrating Benefit-Risk Assessments for**  
527 **Medical Devices**

528 The examples below are hypothetical or simplified real-world situations, and are offered only for  
529 illustrative purposes; i.e., no example is a complete treatment of the benefit-risk issues associated  
530 with any actual FDA decision. The decisions described in these examples are not predictive of  
531 future FDA decisions; rather, they are hypothetical outcomes and are intended only to demonstrate  
532 how FDA considers the factors described in this draft guidance, including how we assess benefits  
533 and risks during product availability, compliance, and enforcement decisions. Similar scenarios  
534 may result in different outcomes depending on the circumstances.  
535

536

## **A. Examples Related to Product Availability Decisions**

537

### **Example 1: Recall and shortage**

538

539 Background: An implantable coated device was developed which reduced thrombosis by more than  
540 80%. There were three field complaints for a malfunction in the device's first few months of wide  
541 scale commercial use. This malfunction represented an anticipated failure mode that occurred more  
542 frequently than expected. During these events associated with the malfunction, blood loss  
543 occurred, but no serious injuries occurred. The manufacturer submitted MDRs for these events.  
544

545

546 Removal of the product from the field would have resulted in the cancellation of hundreds of  
547 surgeries. However, the company recognized that it had product in the field with a postmarket  
548 quality nonconformity requiring a correction or removal, which must be reported to FDA under 21  
549 CFR 806.10. The company proposed to send a communication to the field alerting users to the risk  
550 related to the nonconformity and to continue monitoring the events in the field to better understand  
551 how best to address the issue in the long term.

551

552 Benefits: The patient population for this device includes those patients at elevated risk of  
553 thrombosis. As noted above, this device reduces thrombosis by more than 80%. The likelihood of  
554 the benefit was high. A reduction in thrombosis has significant impact on patient outcomes. The  
555 magnitude of the benefit was high. There were no other comparable treatment options.  
556

557

558 Risks: For the different patient subpopulations that may be treated with the device, FDA  
559 considered the risks of additional blood loss and increased associated surgery time should a device  
560 with this nonconformity be used. Three malfunctions with no serious adverse events had been  
561 reported. The severity of the risk was low. The manufacturer shared information indicating that  
562 3000 devices had been implanted. The likelihood of the risk appears low.

563

564 Patient tolerance for risk and perspective on benefit: Patients appreciate the benefit of limiting  
565 thrombosis. Thrombosis is a concern for many patients and caregivers using these types of devices.

566

567 Uncertainty: FDA considered the uncertainty of the adverse event rate. It was unclear if the adverse  
568 event rate would increase. There were 300 patients in a clinical trial, and there had been 3000  
569 devices implanted in the first few months of wide scale commercial use. As experience with the  
570 device increases, if the number of adverse events and the number of implantations are accurately  
571 tracked, the uncertainty regarding the adverse event rate would decrease.

571

572 Mitigation: FDA reviewed the manufacturer's risk management information, including how this  
573 malfunction can be addressed during surgery to minimize the impact on the patient. FDA also  
574 reviewed the proposed communication to physicians explaining the issue.  
575

576

577 Patient impact: FDA considered the impact on patients if the device was not available in the  
578 marketplace, which included delayed surgeries or treatment with a less beneficial device.

578

579 Decision: FDA found the benefits to be high and the risks to be low in this situation. The  
580 manufacturer shared highly detailed information with FDA, which allowed FDA to better  
581 understand the malfunction rate and mitigation methods. After conducting a Health Hazard

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582 Evaluation, FDA classified this recall as a Class II<sup>5</sup> recall and agreed that the proposed  
583 communication to the device users was in the best interest of public health. FDA and the  
584 manufacturer continued actively monitoring the situation to determine the most appropriate long-  
585 term solution.  
586

587 **Example 2: Evaluation of a variance petition**  
588

589 Background: A drug delivery system was developed that included a safety feature not available  
590 with other medical devices. This system is programmable to automatically suspend drug delivery  
591 when it detects that a predefined threshold has been reached. FDA noted inspectional observations  
592 for deviations from the QS regulation during a PMA pre-approval inspection of the drug delivery  
593 system manufacturer’s facility. The manufacturer petitioned for a variance under section 520(f)(2)  
594 of the FD&C Act (21 U.S.C. 360j(f)(2)) and 21 CFR 820.1(e) from those sections of the QS  
595 regulation for which there were inspectional observations.  
596

597 Benefits: The medical device had a safety feature to stop drug delivery not available on medical  
598 devices already on the market. The unique safety feature stopped drug delivery when the medical  
599 device detected that continued delivery of the drug was no longer indicated and could be harmful.  
600 The magnitude and likelihood of benefit is high.  
601

602 Risks: Several observations of non-compliance with the QS regulation were identified during the  
603 pre-approval inspection. Specifically, the manufacturer did not have a well-functioning CAPA  
604 (corrective and preventive action) system, and several processes lacked documented procedures.  
605 The CAPA system observations did not have a direct impact on patient safety. There was no  
606 indication that nonconforming devices had been released. FDA determined that the severity and  
607 likelihood of risk related to the observations of non-compliance were low in this case, although this  
608 does not mean that non-compliance with CAPA regulations is generally low risk.  
609

610 Patient tolerance for risk and perspective on benefit: Data collected during clinical trials show that  
611 patients and caregivers highly valued this unique safety feature, as it greatly decreased overdose  
612 related fears.  
613

614 Mitigation: As part of the variance, the manufacturer agreed to resolve all of the QS violations by a  
615 set date, and to proactively contact all of the users of the medical device every 90 days to collect  
616 information about the medical device and any malfunctions that might have occurred. The  
617 manufacturer also agreed to investigate all complaints and provide quarterly reports detailing the  
618 results of its surveillance program related to the device to FDA.  
619

620 Decision: FDA agreed that the proposed variance plan provided methods and controls that satisfied  
621 FDA’s concerns in the areas where the QS violations were identified and that were sufficient to

---

<sup>5</sup> “Recall classification means the numerical designation, i.e., I, II, or III, assigned by the Food and Drug Administration to a particular product recall to indicate the relative degree of health hazard presented by the product being recalled.” 21 CFR 7.3(m).

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622 assure that the device will be safe and effective. Given the need in the patient community for a  
623 medical device with the unique safety feature, FDA determined that granting the variance was in  
624 the best interest of the public health.

625

626 **Example 3: Continued access to nonconforming product**

627

628 Background: A biological indicator used in monitoring hospital steam sterilization was not  
629 performing as expected in the field. The manufacturer initiated a voluntary recall, which it reported  
630 to FDA under 21 CFR 806.10. During its internal investigation of this postmarket quality  
631 nonconformity, the manufacturer determined that the source of the problem was with the  
632 manufacturing line and identified which lots were and were not impacted. FDA classified this  
633 recall as Class II.

634

635 The manufacturer had no history of regulatory noncompliance. It opened a CAPA item to address  
636 the root cause of the problem and notified FDA that the long-term correction would result in a  
637 decrease in the volume of biological indicators available to hospitals. The decrease in volume was  
638 projected to last for 18 months. Within a few months, FDA received notification from multiple  
639 sources that surgeries were being delayed due to the lack of biological indicators. The  
640 manufacturer provided information regarding the level of certainty for successful completion of a  
641 sterilization cycle when using the nonconforming biological indicators in accordance with  
642 proposed modified labeling. After consulting with FDA, the manufacturer determined that the  
643 proposed labeling change was one that would require submission of a new 510(k) under 21 CFR  
644 807.81(a)(3).

645

646 Benefits: The manufacturer provided information on the benefit of using the nonconforming  
647 biological indicators according to the modified labeling. While the benefit had decreased from the  
648 anticipated benefit considered for conforming biological indicators during premarket review, the  
649 benefit for this use of the nonconforming biological indicators remained high and included an  
650 assurance of sterility and a reduction in surgical delays.

651

652 Risks: FDA considered the risks associated with use of nonconforming biological indicators. FDA  
653 received no reports of infection or injury related to the biological indicator or the hospital steam  
654 sterilizers for the time that the nonconforming biological indicator was in the field. FDA also  
655 recognized that a properly maintained and operated sterilizer is expected to result in effective  
656 sterilization cycles; the biological indicators provide confirmation. Based on the data and  
657 information available to FDA, the likelihood of risk of harm to patients was assessed to be low if  
658 the nonconforming biological indicators were used in accordance with the proposed modified  
659 labeling.

660

661 Mitigation: In this situation, there was no mitigation that could render the benefit-risk profile of the  
662 nonconforming biological indicators sufficiently positive to justify the continued use of the  
663 nonconforming device, without some additional mitigation step. The manufacturer's proposal to  
664 modify labeling for the nonconforming devices mitigated potential harm to patients.

665

666 Patient tolerance for risk and perspective on benefit: Patient perspective on risks associated with  
667 the nonconforming biological indicators was not readily available. Contact with hospitals indicated  
668 that they were seeking FDA's assistance on how best to manage the shortage of biological

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669 indicators needed to monitor steam sterilization cycles while still protecting their patients from the  
670 potential use of non-sterile devices.

671

672 Patient impact: FDA considered the impact on patient health and quality of life if the  
673 nonconforming biological indicators were not available, which included delayed surgeries or  
674 prioritization of critical surgeries over other surgeries as a result of rationing biological indicators.

675

676 Decision: In this situation, where alternatives were not readily available, FDA worked with the  
677 manufacturer to identify data that supported use of the nonconforming biological indicators with  
678 the proposed modified labeling. FDA concluded that it would not take action against the  
679 manufacturer for marketing the nonconforming biological indicators with that labeling  
680 modification while the manufacturer worked to implement its long-term correction and while the  
681 decreased volume of conforming biological indicators continued. FDA determined that this course  
682 of action would provide the most beneficial option for patients compared to other options.  
683 Consequently, the company was able to provide the marketplace with a sufficient volume of  
684 biological indicators while correcting the underlying problem.

685

686 **B. Examples Related to Compliance and Enforcement**  
687 **Decisions**

688 **Example 1: Evaluating whether to send a Warning Letter or take an alternative**  
689 **approach**

690

691 Background: During an inspection of an aesthetic device manufacturer’s facility, FDA  
692 investigators observed, among other things, that the firm did not maintain adequate complaint files.  
693 Noted deficiencies in the complaint system included a backlog of complaints related to the device  
694 that the manufacturer had not evaluated to determine if an MDR or investigation was necessary  
695 and pending complaint investigations that remained unresolved after more than six months without  
696 explanation. The manufacturer did not submit a response to the Form FDA 483 (FDA 483), List of  
697 Inspectional Observations issued at the close of the inspection.

698

699 Benefits: Reported clinical studies demonstrated that some patients treated saw long-term aesthetic  
700 improvement. The magnitude and likelihood of benefit for the device was assessed to be moderate.  
701 However, the device was not medically necessary, and there was no evidence that it provided a  
702 unique treatment effect or benefit compared to similar devices on the market.

703

704 Risks: The likelihood and severity of risk for similar devices was low. During the inspection,  
705 however, the FDA investigator’s review of a sample of the complaints received for the device  
706 indicated that some patients had experienced adverse events of varying severity. Based on the  
707 information in those complaints, it was unclear if those adverse events may have been caused by  
708 use of the device. Without additional information, FDA could not determine the likelihood of risk  
709 for the device.

710

711 Patient tolerance for risk and perspective on benefit: There was information indicating that patients  
712 preferred similar devices on the market.

713

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714 Uncertainty: The significant deficiencies in the manufacturer’s complaint system created  
715 uncertainty about the risks for this device: the manufacturer could not provide complete  
716 information regarding the number of complaints involving adverse events and the failure to timely  
717 and adequately evaluate complaints may have allowed malfunctions, defects, or other  
718 nonconformities to go undetected.

719

720 Patient impact: FDA determined that there would be no substantial negative impact for patients if it  
721 issued a Warning Letter. Patients or healthcare professionals reluctant to choose a device for which  
722 a Warning Letter has been issued would have alternative products available, and there was not a  
723 strong patient preference for the device.

724

725 Firm compliance history: Some of FDA’s observations related to the manufacturer’s complaint  
726 handling system were repeat observations that had been noted during the previous inspection.

727

728 Decision: After thorough evaluation, FDA decided to issue a Warning Letter to the firm and to  
729 investigate further whether the device may have caused adverse events. Although the device  
730 provided a moderate benefit, that benefit was available to patients through alternatives, and there  
731 was significant uncertainty regarding the likelihood of risk for the device. In addition, the failure to  
732 correct previously noted deficiencies in its complaint system and the failure to respond to the FDA  
733 483 indicated that less formal communications with the firm might be ineffective for achieving  
734 compliance and minimizing risk to patients. If, after gathering further information regarding  
735 adverse events, FDA determined that the device presented a higher risk to patients, the Agency  
736 would consider taking additional action, including action to limit availability of the device.

737

738 **Example 2: Evaluation of potential actions following an inspection with**  
739 **observed Quality System deficiencies**

740

741 Background: FDA’s inspection of a manufacturing facility for a spinal fixation system intended for  
742 posterior, non-cervical pedicle fixation resulted in the issuance of an FDA 483, which noted,  
743 among other things, two complaint records that lacked evaluations to determine if an MDR was  
744 required to be filed, a CAPA record with no documentation of an investigation, and deficiencies in  
745 a process validation. This was FDA’s first inspection of the facility, and some deficiencies were  
746 more significant than others, although none of the deficiencies were significant enough to warrant  
747 a Warning Letter. FDA conducted a benefit-risk analysis as part of its evaluation of whether to  
748 issue an Untitled Letter or to engage with the firm in a less formal manner, such as in a regulatory  
749 meeting.

750

751 Benefits: This firm’s particular spinal fixation system had unique features that made it less invasive  
752 and therefore associated with a shorter surgical time than other devices of its type. Clinical studies  
753 included in the premarket submission for the device demonstrated patient benefits, including  
754 quicker recovery and reduced postsurgical pain. The magnitude and likelihood of the benefit for  
755 this device were assessed to be high.

756

757 Risks: The two complaints that lacked an evaluation for whether an MDR must be submitted did  
758 not involve a death or serious injury, and searches of FDA’s Manufacturer and User Facility  
759 Device Experience (MAUDE) database revealed no MDRs reporting that the device may have  
760 caused or contributed to a death or serious injury. The likelihood of risk to patients was assessed to

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761 be low. FDA reviewed the firm’s nonconformance data during and after the inspection and  
762 determined that it was within the expected parameters for the device. There was no indication that  
763 nonconforming product had been released.

764

765 Patient tolerance for risk and perspective on benefit: FDA considered patient preference. Patients  
766 expressed a strong preference for this spinal fixation system because of the reduction in pain and  
767 recovery time.

768

769 Mitigation: The firm’s responses to the FDA 483 issued at the end of the inspection indicated the  
770 firm’s identification and early implementation of voluntary corrective actions that appeared to be  
771 significant steps to achieve compliance.

772

773 Nature of violations/Nonconforming product: In addition, the inspection did not reveal evidence of  
774 widespread QS deficiencies or nonconformities that were attributed to other failures in the Quality  
775 System.

776

777 Firm compliance history: This manufacturer had no history of regulatory non-compliance.

778

779 Decision: Since, among other things, the firm’s nonconformance data was within the expected  
780 parameters for the device, FDA determined that there was low risk to patients associated with the  
781 inspectional observations. After careful consideration of all available information, FDA pursued a  
782 regulatory meeting with the firm instead of issuing an Untitled Letter to address the manufacturer’s  
783 inspectional deficiencies. FDA decided that, in this lower risk situation, a regulatory meeting  
784 would be the most efficient means of achieving compliance, as it would engage the manufacturer  
785 in a dialogue on its proposed corrections/corrective actions. If the manufacturer fails to progress  
786 toward voluntary compliance in a timely manner, then FDA may consider conducting a follow-up  
787 inspection, issuing a Warning or Untitled Letter, or other consequences.

788

789 **Appendix A - Intersection of this Draft Guidance with ISO**  
790 **14971: *Medical devices – Application of risk management to***  
791 ***medical devices***

792  
793 ISO 14971 provides medical device manufacturers with a framework to systematically manage the  
794 risks to people, property and the environment associated with the use of medical devices.  
795 Specifically, the standard describes a process through which the medical device manufacturer can  
796 identify hazards associated with a medical device, estimate and evaluate the risks associated with  
797 these hazards, control these risks, and monitor the effectiveness of those controls throughout the  
798 product’s life cycle. Implementing this standard requires the user to make decisions on the  
799 acceptability of individual risks, and overall residual risk for a medical device throughout its life  
800 cycle.

801  
802 ISO 14971 is an FDA-recognized standard, and assuring conformity with this standard may help  
803 device manufacturers meet the requirements specified in the design controls section (21 CFR  
804 820.30) and other sections of 21 CFR Part 820. Both ISO 14971 and 21 CFR Part 820 take a total  
805 life cycle approach to management of risks associated with medical devices, and expect that  
806 manufacturers will incorporate postmarket data into their device risk management process,  
807 including new and changes to existing risks identified after the device is on the market.

808  
809 This draft guidance document provides a benefit-risk framework for FDA and stakeholders  
810 regarding use of benefit-risk information in medical device product availability, compliance, and  
811 enforcement decisions. Good documentation of risk management decisions by manufacturers may  
812 help to streamline these decisions for both FDA and manufacturers, produce outcomes for patients  
813 that deliver the most benefit for the least amount of risk, and provide a reasonable assurance of  
814 safety and effectiveness.

815 **Appendix B - Worksheets for Benefit Assessments**

816

817 The following worksheet identifies factors that may be considered in the assessment of benefit for product  
818 availability, compliance and enforcement decisions across the total product life cycle.  
819

<b>Anticipated benefit</b>	<b>Initial assessment during design and testing</b>	<b>Current assessment</b>
<b>Type of benefit(s)</b>	What is the medical device’s anticipated impact on clinical management and patient health? What benefits were initially anticipated? Was a clinical trial conducted? What benefits were expected based on similar devices?	What is the medical device’s impact on clinical management and patient health? Does the marketed product achieve the anticipated benefits? Have additional benefits been observed?
<b>Magnitude of benefit(s)</b>	For each benefit assessed: What was the medical device’s originally anticipated impact on patient health and clinical management? What was the originally anticipated effect of the device on patient management and quality of life, likelihood of survival, improving patient function, preventing loss of function, or providing relief from the symptoms of the disease or condition? What was the anticipated magnitude of each treatment effect? What scale is used to directly measure the anticipated benefit? How did the anticipated benefit rank on that scale? Is the device life supporting or life sustaining?	For each benefit assessed: What is the medical device’s impact on patient health and clinical management? Is the effect of the device on patient management and quality of life, likelihood of survival, improving patient function, preventing loss of function, or providing relief from the symptoms of the disease or condition as anticipated? Did the magnitude of each treatment effect increase or decrease? For each benefit assessed, does real world data demonstrate the same rate of successful diagnosis or treatment? Has the benefit rank on that scale increased or decreased over time? Has real world practice led to new benefits?

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Anticipated benefit	Initial assessment during design and testing	Current assessment
<b>Likelihood of patients experiencing one or more benefits</b>	<p>What proportion of patients was expected to benefit from the device?            Did the original labeling indicate which patients will experience a benefit?            How did the benefits assessed vary across subpopulations?            Was there a variation in public health benefit for different populations?</p>	<p>Using real world data or other data collection, what proportion of patients have been observed to benefit from the device?            Has the likelihood of a patient within a subpopulation experiencing benefit changed?            Has there been a change in variation of benefits across sub-populations?            Has use of the medical device exposed a variation in public health benefit for different populations?</p>
<b>Duration of effects</b>	<p>Does the device cure a disease or provide a temporary treatment?            Could the duration, if relevant, of each treatment effect, be determined?            If so, what was it?</p>	<p>Is the duration of effect consistent with the anticipated duration of effect?            Were there assumptions that proved to be inaccurate?</p>
<b>Patient preference on benefit</b>	<p>What is the severity of the disease state?            Is this a chronic disease?            If chronic, can the illness be managed with other treatments or therapies?            How long do patients live with the disease?            Even if the benefit is in a small portion of the population, do those patients who would experience the benefit value it?            Is the duration of the benefit achieved of value to patients?            How much do patients value this treatment?            Does the treatment improve overall quality of life?            Are the benefits of the medical device well understood?            Is communication regarding change in benefit realistic?</p>	<p>What is the severity of the disease state?            Is this a chronic disease?            If chronic, can the illness be managed with other treatments or therapies?            How long do patients live with the disease?            Even if the benefit is in a small portion of the population, do those patients who would experience the benefit value it?            Is the duration of the benefit achieved of value to patients?            How much do patients value this treatment?            Does the treatment improve overall quality of life?            Are the benefits of the medical device well understood?            Is communication regarding change in benefit realistic?</p>
<b>Benefit factors for healthcare professionals or caregivers</b>	<p>Were there anticipated benefits to healthcare professionals or caregivers?</p>	<p>Does real world experience change the understanding of benefits to professionals or caregivers?</p>

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<b>Anticipated benefit</b>	<b>Initial assessment during design and testing</b>	<b>Current assessment</b>
<b>Medical necessity</b>	<p>Is the device essential to the survival of patients?            Are alternative treatments available?            What other therapies are available for this condition?            How effective are the alternative treatments?            How well-tolerated are the alternative therapies?</p>	<p>Is the device essential to the survival of patients?            Are alternative treatments available?            What other therapies are available for this condition?            How effective are the alternative treatments?            How well-tolerated are the alternative therapies?            How have treatment options changed since medical device development?</p>

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820 **Appendix C - Worksheets for Risk Assessments**

821

822 The following worksheet identifies factors that may be considered in the assessment of risk for  
823 product availability, compliance and enforcement decisions across the total product life cycle.  
824

Risk categories	Initial assessment during design and testing	Current assessment
<b>Factors Related to Risk Severity</b>		
<b>Medical device-related deaths and serious injuries</b>	What serious adverse events related to this medical device were known when FDA authorized the device for marketing? Were there any variations in serious adverse events among subpopulations?	Have medical device-related deaths or serious injuries occurred at expected severity? Are there unanticipated deaths or serious injuries? Were there any changes variations of serious adverse events among subpopulations?
<b>Medical device-related non-serious adverse events</b>	What non-serious adverse events related to this medical device were known at medical device clearance or approval? Were there any variations in temporary injury and medically reversible injuries among subpopulations?	Have temporary injuries related to the medical device occurred at expected severity? Have medical device-related injuries which could be medically reversed occurred at expected severity and frequency? Are there any unanticipated temporary injuries or medically reversible injuries? Were there any changes in variations in serious adverse events among subpopulations?
<b>Medical device-related events without reported harm</b>	What medical device malfunctions were anticipated when FDA authorized the device for marketing? Were there any variations in medical device events reported without harm among subpopulations?	Are there reports of medical device malfunctions? Are device malfunctions occurring at anticipated frequencies? Is the medical device malfunctioning in a manner that was not anticipated? Were there any changes in variations in medical device events reported without harm among subpopulations?

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<b>Duration of harm to patient</b>	How long does the harmful event last? Is the harmful event reversible? What type of intervention is required to address the harmful event?	Is the duration of harmful events longer than anticipated? Is the harmful event still reversible? Has the type of intervention required to address the harmful event changed?
<b><u>Risk factors related to Likelihood of Risk</u></b>		
<b>Likelihood of medical device nonconformity</b>	How frequently did the manufacturer anticipate this specific failure mode or defect would occur?	How frequently does this specific failure mode or defect occur? Has the rate of medical device failures increased? Has the mean time between failures decreased? How many medical devices are expected to have a problem?
<b>Likelihood of a harmful event given exposure to a nonconforming device</b>		What proportion of patients treated with or diagnosed by the nonconforming medical device is harmed?
<b>Number of patients exposed</b>		How many patients were exposed to nonconforming devices? How many patients were exposed to a device manufactured by a noncompliant manufacturer?
<b><u>Additional Risk Factors</u></b>		
<b>Nonconforming product risks</b>		Has nonconforming product been distributed? What is the number of units on the market and market share ?
<b>Duration of the exposure to the population</b>		How much time elapsed between initial exposure to a risk of harm and the point at which the risk of harm is successfully addressed? How long were affected populations exposed to the nonconforming device?

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<p><b>Risk from false-positive or false-negative results for diagnostics</b></p>	<p>What are the consequences of a false positive?          What are the consequences of a false negative?          Is this the only means of diagnosing the problem, or is it part of an overall diagnostic plan?</p>	<p>Have the consequences of diagnostic errors changed?          Have the practices related to diagnosing the problem changed?          Does this increase or decrease the risk?</p>
<p><b>Patient tolerance of risk</b></p>	<p>What level of concern do patients have regarding the risks?          Even if the risk is in a small portion of the population, do those patients who would experience the risk understand it?          Are patients willing to take the risk of this treatment to achieve the benefit?          How well are patients able to understand the risks of the treatment?</p>	<p>What level of concern do patients have regarding the risks?          Even if the risk is in a small portion of the population, do those patients who would experience the risk understand it?          Are patients willing to take the risk of this treatment to achieve the benefit?          How well are patients able to understand the risks of the treatment?</p>
<p><b>Risk Factors for healthcare professionals or caregivers</b></p>	<p>Are there risks to the healthcare professional or caregiver?          How significant are these risks?</p>	<p>Are there any changes in frequency or severity of risks for healthcare professionals and/or caregivers?          Do any changes in the frequency or severity of risk for the healthcare provider or caregiver impact the risks to the patient?</p>

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826 **Appendix D - Worksheet for assessing potential decisions**  
827 **based on the Benefit-Risk Assessment Outcome**

828  
829 The following worksheet identifies additional factors that may be considered for product  
830 availability, compliance and enforcement decisions at all phases of the total product life cycle.  
831

<b>Factors</b>	<b>Assessment Questions</b>
<b>Uncertainty</b>	What information does FDA have to assess benefit and risk? What is the quality of the information FDA is using (for example, MDRs, literature, registry or clinical trial data, limited case studies, etc.)? What is the uncertainty related to current understanding of benefits and risks?
<b>Mitigations</b>	Could you identify ways to mitigate the risks such as using product labeling, establishing education programs, etc.? What is the type of mitigation proposed? Is the intervention related to design, labeling, or training? Has the manufacturer corrected the cause of the nonconformity?
<b>Detectability</b>	Can the user easily recognize the hazard to avoid the harm? Can the problem with the medical device be corrected before use by the user?
<b>Failure Mode</b>	Has the manufacture identified the underlying cause? Has the firm submitted testing to the FDA? Has FDA conducted testing? What were the results?
<b>Scope of the device issue</b>	Are the risks identified potentially inherent to similar medical devices of this type (i.e., industry wide)?
<b>Patient impact</b>	What are the risks to patients if the device is not available? Are patients better off if the device is available? What are the risks to patients related to the inspectional observation or regulatory non-compliance? Does the observation or violation directly relate to product quality? Does the observed regulatory non-compliance raise concerns regarding the firm's ability to produce safe and effective medical devices?
<b>Preference for availability</b>	Would patients and caregivers prefer to have access to the device? Are the benefits and risks adequately understood?
<b>Nature of violations/Nonconforming product</b>	Was the violation systemic or non-systemic in nature? To what extent are the products nonconforming?

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**Firm compliance history**

Has the same or a similar inspectional observation or regulatory violation been observed at the manufacturer in the past 2 years? In the past 5 years? In the past 10 years?  
Does the firm have a history of regulatory compliance and high quality device production?  
Has the firm demonstrated chronic and systematic regulatory non-compliance over time?  
Is the regulatory non-compliance significant enough that FDA would take regulatory action?  
Was the harm anticipated in the firm risk management documentation?  
Was the harm reported to FDA by the firm quickly?  
Would providing notice to the firm assist in informing the firm of its legal responsibilities?

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