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1 **Peripheral Percutaneous Transluminal**
2 **Angioplasty (PTA) and Specialty**
3 **Catheters - Premarket Notification**
4 **(510(k)) Submissions**

5 **Draft Guidance for Industry and Food**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

7 ***DRAFT GUIDANCE***

8 **This guidance document is being distributed for comment purposes only.**

9 **Document issued on January 13, 2020.**

10 You should submit comments and suggestions regarding this draft document within 60 days of
11 publication in the *Federal Register* of the notice announcing the availability of the draft
12 guidance. Submit electronic comments to <https://www.regulations.gov>. Submit written
13 comments to Dockets Management Staff (HFA-305), Food and Drug Administration, 5630
14 Fishers Lane, rm. 1061, Rockville, MD 20852. Identify all comments with the docket number
15 listed in the notice of availability that publishes in the *Federal Register*.

16 For questions regarding this document, contact the Plaque Modification Devices Team in OHT2:
17 Office of Cardiovascular Devices/DHT2C: Division of Health Technology 2C at (301) 796-
18 6075.

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



24

Preface

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27 CDRH-Guidance@fda.hhs.gov to receive a copy of the guidance. Please include the document
28 number 16018 and complete title of the guidance in the request.

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Peripheral Percutaneous Transluminal Angioplasty (PTA) and Specialty Catheters – Premarket Notification (510(k)) 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Draft Guidance for Industry and Food and Drug Administration Staff

This draft guidance, when finalized, will represent the current thinking of the Food and Drug Administration (FDA or Agency) on this topic.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

This draft guidance document provides draft recommendations, including bench testing and coating characterizations for 510(k) submissions for peripheral percutaneous transluminal angioplasty (PTA) balloons and specialty catheters (e.g., infusion catheters, PTA balloon catheters for in-stent restenosis (ISR), scoring/cutting balloons). These devices are catheter-based devices intended to treat lesions in the peripheral vasculature. This document provides anatomy-specific testing recommendations and expands on FDA’s current thinking for testing of these devices. FDA is issuing this draft guidance to clarify FDA’s premarket submission recommendations for PTA catheters and specialty catheters and to promote consistency across submissions.

For the current edition of the FDA-recognized standards referenced in this document, see the FDA Recognized Consensus Standards database at <https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/search.cfm>. For more information regarding use of consensus standards in regulatory submissions, please refer to the FDA guidance titled “[Appropriate Use of Voluntary Consensus Standards in Premarket Submissions for Medical Devices](#).”¹

¹<https://www.fda.gov/regulatory-information/search-fda-guidance-documents/appropriate-use-voluntary-consensus-standards-premarket-submissions-medical-devices>

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79 This document supplements other FDA documents regarding the specific content requirements
80 of premarket submissions. You should also refer to 21 CFR 807.87 and FDA’s guidance,
81 [“Format for Traditional and Abbreviated 510\(k\)s.”](#)²

82
83 FDA's guidance documents, including this draft guidance, do not establish legally enforceable
84 responsibilities. Instead, guidances describe the Agency’s current thinking on a topic and should
85 be viewed only as recommendations, unless specific regulatory or statutory requirements are
86 cited. The use of the word *should* in Agency guidance means that something is suggested or
87 recommended, but not required.
88

89 II. Scope

90 The scope of this document is limited to class II PTA balloon catheters regulated under 21 CFR
91 870.1250 and class II specialty catheters regulated under 21 CFR 870.1210 and 21 CFR
92 870.1250 with product codes listed in the table below.

93 **Table 1: Device Types within the Scope of This Guidance.**

Regulation Number	Product Code	Device
870.1210	KRA	Continuous Flush Catheter
870.1250	DQY	Percutaneous Catheter
870.1250	LIT	Peripheral Transluminal Angioplasty Catheter
870.1250	PNO	Percutaneous Cutting/Scoring Catheter

94 In this guidance, PTA balloon catheters refer to standard peripheral angioplasty balloon
95 catheters. Specialty catheters can include but are not limited to the following 510(k) devices:
96 infusion catheters, balloon catheters with unique design characteristics (e.g., cutting/scoring),
97 and balloon catheters intended for specific indications (e.g., ISR, post-dilatation of stents).

98 III. Premarket Submission Recommendations

99 A. Device Description

100 We recommend you identify your device by the applicable regulation number and product code
101 indicated in Section II above and include the information described below.

- 102 • **Device components and mode of operation:** FDA recommends that you identify all
103 components and accessories included in the premarket submission, including packaging,
104 with a clear description of how the device is utilized to achieve the intended use in the
105 intended anatomy.

² <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/format-traditional-and-abbreviated-510ks-guidance-industry-and-fda->

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- 106 • **Photograph and engineering drawing(s) of the device:** FDA recommends that you
107 provide a photograph, as well as an engineering drawing with all dimensions, tolerances,
108 and components labeled, of the device. FDA recommends that you include this for each
109 device, accessory, or component included in the premarket submission.

- 110 • **Technological characteristics:** FDA recommends that you describe the technical and
111 performance specifications and include a brief description of the device design
112 requirements in the device description section of the premarket submission. The
113 specifications may include performance-related product measurement tolerances,
114 operating limitations, and any other functional, physical, and environmental
115 specifications of the device. We also recommend that you describe ranges and/or
116 accuracy of the specifications.

- 117 • **Materials:** FDA recommends that you provide a list of all components, their respective
118 material(s) of composition, and their patient-contacting classification (e.g., non-
119 contacting, indirect-contacting, or direct-contacting). For each component, you should
120 identify the generic material of construction and the unique material identifier.

B. Predicate Comparison

122 For devices reviewed under the 510(k) process, manufacturers must compare their new device to
123 a similar legally marketed predicate device to support its substantial equivalence (21 U.S.C.
124 360c(i); 21 CFR 807.87(f)). This comparison should provide information to show how your
125 device is similar to and different from the predicate. Side by side comparisons, whenever
126 possible, are desirable. See below for an example of how this information may be organized.
127 This table is not intended to represent an exhaustive list of comparative parameters; ensure you
128 provide all relevant device descriptive characteristics as outlined in the “Device Description”
129 section, above.

Description	Subject Device	Predicate Device (Kxxxxxx)
Indications for use		
Guidewire Compatibility		
Sheath Compatibility		
Catheter length		
Catheter Shaft Outer Diameter		
Balloon Lengths (if applicable)		
Balloon Diameters (if applicable)		
Nominal Pressure (if applicable)		
Rated Burst Pressure (if applicable)		

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Description	Subject Device	Predicate Device (Kxxxxxx)
Component Materials (list individually)		
Coating Material (if applicable)		
Coating Length (if applicable)		
Packaging Configuration		
Sterilization Method		

131
132
133

C. Biocompatibility

134 Significance: PTA balloon catheters and specialty catheters contain patient-contacting materials,
135 which, when used for their intended purpose (i.e., contact type and duration), may induce a
136 harmful biological response.

137 Recommendation: You should determine the biocompatibility of all patient-contacting materials
138 present in your device. If your device is identical in composition and processing methods to any
139 PTA balloon catheters or specialty catheters with a history of successful use, you may reference
140 previous testing experience or the literature, if appropriate. For some device materials, it may be
141 appropriate to provide either a reference to an FDA-recognized consensus standard or a letter of
142 authorization (LOA) for a device master file (MAF).

143 If you are unable to identify a legally marketed predicate device with similar location/ duration
144 of contact and intended use that uses the same materials as used in your device, we recommend
145 you conduct and provide a biocompatibility risk assessment. The assessment should explain the
146 relationship between the identified biocompatibility risks and potential mitigation strategies as
147 well as identify any knowledge gaps that remain. You should then identify any biocompatibility
148 testing or other evaluations that have been conducted to mitigate any remaining risks.

149 We recommend that you follow the FDA guidance, “[Use of International Standard ISO-10993-1,](#)
150 [‘Biological evaluation of medical devices – Part 1: Evaluation and testing within a risk](#)
151 [management process’](#)”³, which identifies the types of biocompatibility assessments that should
152 be considered and recommendations regarding how to conduct related tests.

153 Per ISO 10993-1: *Biological evaluation of medical devices – Part 1: Evaluation and testing*
154 *within a risk management process* and Attachment A of FDA’s guidance on ISO-10993-1, PTA
155 balloon catheters and specialty catheters are external-communicating devices in contact with
156 circulating blood for a limited contact duration. Therefore, the following endpoints should be
157 addressed in your biocompatibility evaluation:

³ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/use-international-standard-iso-10993-1-biological-evaluation-medical-devices-part-1-evaluation-and>

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- 158 • cytotoxicity;
- 159 • sensitization;
- 160 • irritation or intracutaneous reactivity;
- 161 • acute systemic toxicity;
- 162 • material-mediated pyrogenicity;
- 163 • hemocompatibility;
- 164 ○ direct and indirect hemolysis;
- 165 ○ SC5b-9 complement activation; and
- 166 ○ thrombogenicity.

167 Please note that a genotoxicity assessment may be requested if PTA balloon catheters or specialty
168 catheters contain novel patient-contacting materials that have not been previously evaluated for use
169 in contact with circulating blood in legally marketed medical devices.

170
171 If an animal study is being conducted in order to evaluate the safety or performance of your
172 device, you may consider evaluating your device for a thrombogenic response in this study in
173 lieu of a 4-hour canine study. If you choose this approach, you should capture information
174 comparable to the 4-hour canine study (e.g., anticoagulation regimen, activated clotting time
175 (ACT), thrombus formation on your device and the implanted vessel after use, including
176 pictures). If anticoagulation is used, you should discuss how this method relates to clinical
177 practice.

178 179 **D. Sterility**

180 Significance: PTA balloon catheters and specialty catheters come in contact with blood and
181 should be adequately sterilized to minimize infections and related complications.

182 Recommendation: For PTA balloon catheters and specialty catheters labeled as sterile, we
183 recommend that you provide information for the finished device in accordance with the FDA
184 guidance, “[Submission and Review of Sterility Information in Premarket Notification \(510\(k\)\)](#)
185 [Submissions for Devices Labeled as Sterile.](#)”⁴

186 Devices in contact with the cardiovascular system should meet Devices in contact with the
187 cardiovascular system should meet pyrogen limit specifications discussed in the FDA guidance,

⁴ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/submission-and-review-sterility-information-premarket-notification-510k-submissions-devices-labeled>

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188 [“Submission and Review of Sterility Information in Premarket Notification \(510\(k\)\)](#)
189 [Submissions for Devices Labeled as Sterile.”](#)⁵

190

191 **E. Pyrogenicity**

192 **Significance:** Pyrogenicity testing is used to assess the risk of febrile reaction due to gram-
193 negative bacterial endotoxins and/or chemicals that can leach from a medical device (e.g.,
194 material-mediated pyrogens).

195 **Recommendation:** To address the risks associated with the presence of bacterial endotoxins, PTA
196 balloon catheters and specialty catheters should meet pyrogen limit specifications by following
197 the recommendations outlined in the 510(k) Sterility Guidance. You should also follow the
198 recommendations in [“Guidance for Industry Pyrogen and Endotoxins Testing: Questions and](#)
199 [Answers.”](#)⁶ To address the risks associated with material-mediated endotoxins, you should
200 follow the recommendations in the FDA guidance, [“Use of International Standard ISO-10993,](#)
201 [‘Biological Evaluation of Medical Devices Part 1: Evaluation and Testing’.](#)”⁷

202

203 For devices intended to be labeled as “non-pyrogenic,” we recommend that both bacterial
204 endotoxin and material-mediated pyrogens be addressed. Devices in contact with the
205 cardiovascular system should meet pyrogen limit specifications discussed in the FDA guidance,
206 [“Submission and Review of Sterility Information in Premarket Notification \(510\(k\)\)](#)
207 [Submissions for Devices Labeled as Sterile.”](#)⁸

208

209 **F. Shelf-Life and Packaging**

210 **Significance:** Shelf-life testing is conducted to support the proposed expiration date through
211 evaluation of the package integrity for maintaining device sterility and/or evaluation of any
212 changes to device performance or functionality.

213 **Recommendation:** With respect to package integrity for maintaining device sterility for PTA
214 balloon catheters and specialty catheters, you should provide a description of the packaging,
215 including how it will maintain the device’s sterility, a description of the package integrity test
216 methods, and a summary of the package integrity test data, including the test, acceptance criteria,
217 results, and any deviations noted.

218

⁵ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/submission-and-review-sterility-information-premarket-notification-510k-submissions-devices-labeled>

⁶ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/guidance-industry-pyrogen-and-endotoxins-testing-questions-and-answers>

⁷ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/use-international-standard-iso-10993-1-biological-evaluation-medical-devices-part-1-evaluation-and>

⁸ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/submission-and-review-sterility-information-premarket-notification-510k-submissions-devices-labeled>

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219 After subjecting the full packaging configuration to simulated shipping (per ASTM D4169:
220 *Standard Practice for Performance Testing of Shipping Containers and Systems*) and climatic
221 conditioning (per ASTM D4332: *Standard Practice for Conditioning Containers, Packages, or*
222 *Packaging Components for Testing*), we recommend that you assess the packaging integrity and
223 strength of both the materials and seal of the sterile barrier. The integrity of the packaging
224 materials can be assessed using test methods such as the bubble leak test (per ASTM F2096:
225 *Standard Test Method for Detecting Gross Leaks in Packaging by Internal Pressurization*
226 *(Bubble Test)*) and burst testing (per ASTM F2054/F2054M: *Standard Test Method for Burst*
227 *Testing of Flexible Package Seals Using Internal Air Pressurization Within Restraining Plates*).
228 The integrity of the seals can also be assessed using numerous test methods, including a visual
229 assessment (per ASTM F1886/F1886M: *Standard Test Method for Determining Integrity of*
230 *Seals for Flexible Packaging by Visual Inspection*), the bubble leak test (per ASTM F2096:
231 *Standard Test Method for Detecting Gross Leaks in Packaging by Internal Pressurization*
232 *(Bubble Test)*), and the dye penetration test (per ASTM F1929: *Standard Test Method for*
233 *Detecting Seal Leaks in Porous Medical Packaging by Dye Penetration*). A seal strength
234 assessment (per ASTM F88/F88M: *Standard Test Method for Seal Strength of Flexible Barrier*
235 *Materials*) should also be conducted at baseline and after aging (accelerated with real-time
236 confirmatory testing) in order to ensure that the seals will not be compromised due to any force
237 exerted on the seal.

238 With respect to evaluating the effects of aging on device performance or functionality, shelf-life
239 studies should evaluate the critical physical and mechanical properties of the device that are
240 required to ensure it will perform adequately and consistently during the entire proposed shelf
241 life. To evaluate device functionality after aging, we recommend that you assess each of the
242 bench tests described in Section III.G and repeat all tests that evaluate design components or
243 characteristics that may be affected by aging. A rationale should be provided for any deviations
244 from the methods used for the baseline testing (e.g., smaller sample size, different device sizes
245 assessed, omitted testing).

246 For PTA balloon catheters and specialty catheters that are provided sterile and/or have a
247 proposed expiration date, we recommend that you provide a summary of the test methods used
248 for your shelf-life testing, results and the conclusions drawn from your results. If you use devices
249 subject to accelerated aging for shelf life testing, we recommend that you specify the way in
250 which the devices were aged. We recommend that you age your devices as per the currently
251 FDA-recognized version of ASTM F1980: *Standard Guide for Accelerated Aging of Sterile*
252 *Barrier Systems for Medical Devices* and specify the environmental parameters established to
253 attain the expiration date. For devices or components containing polymeric materials, you should
254 plan to conduct testing on real-time aged samples to confirm that the accelerated aging is
255 reflective of real-time aging. This testing should be conducted in parallel with 510(k) review and
256 clearance with results documented to file in the device's design history file in accordance with
257 the provisions of 21 CFR 820.30 (i.e., the test reports do not need to be submitted to FDA).

258
259

260 **G. Non-Clinical Performance Testing**

261 **(1) Standard Performance Testing for PTA and Specialty**
262 **Catheters**

263 Non-clinical performance testing is recommended for PTA and specialty catheters in order to
264 fully characterize the device and also ensure that the devices can perform as intended. The
265 testing recommended below should be conducted on the finished product that was subjected to
266 all manufacturing processes, including sterilization. Otherwise, a discussion of the differences
267 between the test article and finished product should be discussed and justified.

268
269 For information on recommended content and format of test reports for the testing described in
270 this section, refer to FDA’s guidance, “[Recommended Content and Format of Non-Clinical](#)
271 [Bench Performance Testing Information in Premarket Submissions.](#)”⁹

272
273 Please note that the recommendations provided in ISO 10555-1: *Intravascular Catheters –*
274 *Sterile and Single-Use Intravascular Catheters – Part 1: General Requirements* and ISO 10555-
275 *4: Sterile and Single-Use Intravascular Catheters – Part 4: Balloon Dilatation Catheters* are
276 directly applicable to PTA catheters and many specialty catheters. Therefore, the testing and
277 methods recommended in these standards should be followed, or a rationale for deviating from
278 these methods should be provided. However, these standards may not include all testing
279 recommended by FDA or may not be specific enough regarding the type of recommended
280 testing. Therefore, the recommendations described below, which augment these consensus
281 standards, should also be followed.

282
283 **a. Dimensional Verification**

284 Significance: Accurate device dimensions help the physician to select the proper product and
285 accessory device sizes. They may also affect the operator’s ability to track the catheter to and
286 across lesions.

287 Recommendation: We recommend that you provide dimensional specifications and tolerances as
288 well as data to verify that these specifications are met for your device as manufactured. At a
289 minimum, we recommend that you measure and report catheter effective length, shaft inner and
290 outer diameter, and crossing profile. For balloon catheters, the balloon outer diameter and length
291 should also be characterized, as described in ISO 10555-4.

292 The crossing profile is defined as the maximum diameter found between the proximal end of the
293 balloon and the distal tip of the catheter. Testing should address potential differences in crossing
294 profile that may exist in the circumferential direction. For these situations, we recommend that
295 you evaluate the crossing profile of your catheter along different longitudinal paths (e.g., rotating
296 the test sample 90° for measurements). We recommend that you report the crossing profile in the

⁹ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/recommended-content-and-format-non-clinical-bench-performance-testing-information-premarket>

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297 instructions for use, the outside package labeling, or both. We recommend using the methods
298 described in ASTM F2081-06: *Standard Guide for Characterization and Presentation of the*
299 *Dimensional Attributes of Vascular Stents* or their equivalents. If pass/fail testing is employed,
300 such as “go/no go” gauges, a rationale should be provided to support this method and the size of
301 these aids.

302 The crossing profile data should be used to support the labeled introducer sheath compatibility.
303 Since the size of commercially-available introducer sheaths vary, introducer sheath compatibility
304 testing alone is not sufficient to support a labeled sheath compatibility. If you are labeling your
305 device with a smaller introducer sheath compatibility than your crossing profile data, a scientific
306 rationale should be provided. Inner diameter data should be used to support the labeled guide
307 wire compatibility. Pass/fail sheath compatibility and guidewire compatibility testing can be also
308 conducted with the simulated use assessment as described in Section III.G(1)b, but should be
309 considered supporting information.

310 **b. Simulated Use**

311 Significance: The recommended instructions for use and techniques for preparation, insertion,
312 tracking, deployment, retraction, and removal, if properly followed, should safely and reliably
313 deploy the balloon to the intended location without adversely affecting the device.

314 Recommendation: We recommend that you conduct testing to demonstrate that the balloon
315 catheter can be safely and reliably prepared, inserted, tracked, deployed, retracted, and removed
316 using the recommended techniques, accessory devices, and instructions for use, without damage
317 to the device. We recommend that this simulated use testing be performed by tracking the device
318 through an *in vitro* fixture that mimics *in vivo* physiologic and anatomic conditions (e.g., a
319 tortuous path in a 37 °C aqueous environment) to the length that would enter a patient in clinical
320 use. The clinical basis and rationale for the model used should be provided. In general, FDA
321 recommends a three-dimensional model, including a worst-case entry angle, with a sufficient
322 number of curves. The length, diameters, number of curves, and radii of curvature should be
323 representative of worst-case anatomy for which the device is intended. An engineering drawing,
324 with all dimensions labeled, and images of the model should be provided.

325 We recommend that you conduct testing with accessory devices that would be used in a typical
326 clinical procedure (e.g., introducer) using worst-case sizes (e.g., smallest inner-diameter
327 introducer sheath per labeled compatibility). You should report any abnormality or difficulty
328 observed during the simulated procedure as well as any damage observed to the PTA catheter or
329 any of the accessory devices.

330 We recommend that you measure and report the diameter and axial location of the largest
331 deflated balloon profile, including the inner member or wire. This information can be used to
332 determine the extreme dimensions of compatible accessory devices (i.e., minimum internal
333 diameter), which should be identified in the labeling. Determining the insertion/retraction forces
334 may also be informative as this may assist in supporting the specifications used for device tensile
335 testing.

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336 It may be possible to combine the simulated use testing with coating integrity testing (see
337 Section III.G(1)l) and/or particulate evaluation (see Section III.G(1)m), but you should take care
338 to ensure that only minimal additional handling of the sample is required for the coating integrity
339 evaluation such that particulates are neither lost nor generated.

340 **c. Balloon Rated Burst Pressure**

341 Significance: The rated burst pressure (RBP) is the pressure at which 99.9% of balloons can
342 survive with 95% confidence. Failure of a balloon to maintain integrity at the RBP could result
343 in device failure or vessel damage.

344 Recommendation: We recommend that you follow ISO 10555-4, Annex A, when conducting this
345 testing. In addition to what is described in this standard, the following should be taken into
346 consideration.

347 We recommend that you conduct this testing on complete catheters or subassemblies in which
348 the balloon is mounted on the catheter shaft. We recommend that you conduct testing on the
349 longest length of every balloon diameter and the shortest length of both the smallest diameter
350 and largest diameter. **Table 2** illustrates the recommended test matrix.

351 **Table 2: Balloon Sizes Recommended for RBP Testing (Example).**

Balloon Diameter (mm)	Balloon Length (mm)				
	40	60	80	100	120
4.0	X				X
4.5					X
5.0					X
5.5					X
6.0	X				X

352 We recommend that you test balloons that are not constrained by any test fixture, such as tubing,
353 and that you inflate the balloons incrementally until failure. We recommend that you record as
354 test failures any loss of:

- 355 • integrity of the balloon, such as a rupture or leak; or
- 356 • pressure due to failure of the balloon, shaft, or seals.

357 We recommend that you record the pressure at which the device failed and the failure mode
358 (e.g., longitudinal tear, circumferential tear, pinhole). A discussion and rationale should be
359 provided for the failure mode observed. We also recommend that you calculate RBP as the
360 pressure at which 99.9% of the balloons will survive with 95% confidence based on statistical
361 analysis of the test data. The lower tolerance limit determined from this analysis should be
362 reported and be used to support the RBP specified in the device labeling.

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363 **d. Balloon Fatigue (Repeat Balloon Inflations)**

364 Significance: Balloons on PTA catheters are often inflated multiple times during clinical use.
365 Failure of the balloon to withstand multiple inflations could lead to device failure or vessel
366 damage.

367 Recommendation: We recommend that you follow ISO 10555-4, Annex B, when conducting this
368 testing, unless otherwise specified below. In addition to what is described in this standard, the
369 following should be taken into consideration.

370 We recommend that you determine the repeatability, to 20 inflations, of successful balloon
371 inflation to the RBP. We recommend that you test device sizes according to the “four corners”
372 paradigm:

- 373 • largest diameter/longest length;
- 374 • largest diameter/shortest length;
- 375 • smallest diameter/longest length; and
- 376 • smallest diameter/shortest length.

377 **Table 3** illustrates the recommended test matrix.

378 **Table 3: Example of “Four Corners” Test Matrix.**

Balloon Diameter (mm)	Balloon Length (mm)				
	40	60	80	100	120
4.0	X				X
4.5					
5.0					
5.5					
6.0	X				X

379 We recommend that you test balloons that are not constrained by any test fixture such as tubing
380 and that you inflate the balloons incrementally until they reach the RBP. For each sample, we
381 recommend that you hold the RBP for 30 seconds (or the time specified in the instructions for
382 use), deflate the balloon, and inflate it again to the RBP, for a total of 20 cycles (or provide a
383 scientific rationale to support the number of cycles used as worst-case). 20 cycles are
384 recommended in order to ensure that testing is worst-case and provides a sufficient safety margin
385 for clinical use, as PTA catheters and other specialty catheters may be inflated multiple times.
386 Please note that ISO 10555-4 recommends inflation for 10 cycles. If fewer cycles are used for
387 this testing than the FDA-recommended 20 cycles, we recommend that you provide a clinical
388 rationale to support the methods used. We recommend that you report any loss of pressure,
389 whether due to failure of the balloon, shaft, or proximal or distal seals, as a test failure. We
390 recommend that you record all failure modes and that your results demonstrate that 90% of the
391 balloons will survive the test with at least 95% confidence.

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e. Balloon Compliance (Diameter vs. Pressure)

394 Significance: The diameter of a deployed PTA balloon varies with inflation pressure. A
395 compliance chart in the labeling that relates balloon diameter to balloon pressure guides selection
396 of catheter size to fit the target vasculature site. Incorrect selection of catheter size may lead to
397 device failure or vessel damage.

398 Recommendation: We recommend that you follow ISO 10555-4, Annex D, when conducting this
399 testing. In addition to what is described in this standard, the following should be taken into
400 consideration.

401 We recommend that you test balloon sizes, as illustrated in Table 2, and that you test multiple
402 product lots. We recommend that you include data showing inflation pressure versus balloon
403 diameter over the full range of recommended inflation diameters and report the results in the
404 instructions for use, the outside package labeling, or both. A graphical or tabular presentation
405 (i.e., a compliance chart) should be included in the labeling. We recommend that you identify the
406 nominal inflation pressure and RBP. The compliance chart may include pressures up to (but not
407 exceeding) 25% above the RBP, if you provide data and statistics demonstrating that 99% of the
408 balloons will not fail at the listed pressure with 95% confidence. We also recommend that you
409 describe if and how you performed any data rounding and show all instances, if applicable.
410 Compliance charts should not be normalized (i.e., modified in any way in order to ensure that the
411 nominal diameter is exactly achieved at the labeled nominal pressure) or calculated based on
412 limited testing. Table 4 shows an example of compliance chart for a balloon with 4.0 mm to 6.0
413 mm diameters, with a nominal pressure of 9 atm and varying RBPs.

414

Table 4: Balloon Compliance Chart Example.

Pressure (atm)	Balloon Nominal Diameter (mm) (X = balloon diameter at the given pressure)				
	4.0	4.5	5.0	5.5	6.0
9.0*	X	X	X	X	X
10.0	X	X	X	X	X
11.0	X	X	X	X	X
12.0	X	X	X	X	X
13.0	X	X	X	X	X
14.0	X	X	X	X**	X**
15.0	X	X	X**	X	X
16.0	X**	X**	X	X	X

415

*Nominal; **RBP

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416 **f. Balloon Inflation and Deflation Time**

417 Significance: Balloons occlude the target vessel and obstruct blood flow while inflated. Inflation
418 and deflation times affect occlusion time. Excessively slow inflation or deflation of a balloon
419 could lead to prolonged lack of blood flow and damage to downstream tissues. Both inflation and
420 deflation time are pertinent to evaluate, as both of these attributes may affect device performance
421 and may result in prolonged lack of blood flow and damage to downstream tissues.

422 Recommendation: We recommend that you follow ISO 10555-4, Annex C, for deflation time
423 testing. In addition to what is described in this standard, the following should be taken into
424 consideration when conducting balloon inflation and deflation time testing.

425 We recommend that you demonstrate, using techniques recommended in your instruction
426 manual, that the balloon inflates and deflates within acceptable times and provide the clinical
427 basis for your acceptance criteria. We recommend that you test the largest diameter at the longest
428 balloon length and evaluate which other sizes to test based on your risk analysis. We also
429 recommend you specify the balloon deflation times in your labeling.

430 **g. Catheter Bond Strength**

431 Significance: Failure of bonds in the catheter could lead to device failure, vessel damage, and/or
432 embolic risk due to device remnants within the vasculature.

433 Recommendation: We recommend that you test the bond strength at all locations where
434 adhesives, thermal fusion, or other joining methods are used for bonding components of the
435 catheter. Multiple bonds/joints that are located in close proximity should not be tested together
436 unless they are physically joined at the same location. Prior to evaluating tensile strength, we
437 recommend you precondition catheters by tracking through a tortuous path fixture (as described
438 in Section III.G(1)b). We recommend that the testing demonstrate that all joints/bonds can
439 withstand tensile forces greater than those that may be experienced during clinical use. As such,
440 we also recommend that you provide the clinical basis (e.g., literature, retraction forces) for your
441 bond strength acceptance criteria. As discussed above, insertion and retraction force assessments
442 during simulated use testing may also be used to support your bond strength acceptance criteria.
443 Comparative testing involving a legally marketed predicate device that has a history of safe use
444 is also appropriate. Please note that the values identified in ISO 10555-1: *Intravascular*
445 *Catheters – Sterile and Single-Use Intravascular Catheters – Part 1: General Requirements*
446 alone should not be used to rationalize your acceptance criteria, as the clinical relevance of these
447 criteria have not been established for peripheral interventional applications. The test
448 method/protocol for this testing should clearly describe the methods utilized, including the
449 portions of the device that were fixed into each clamp and the pull rate.

450 **h. Tip Pull Test**

451 Significance: Failure of bonds in the distal tip could lead to device failure, vessel damage, and/or
452 embolic risk due to device remnants within the vasculature.

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453 Recommendation: For devices with one or more joints in the distal tip (e.g., spring or nose-cone
454 tips), we recommend evaluating the tensile force that will separate the distal tip from the
455 catheter. Prior to testing, we recommend that you precondition catheters prior to tip pull testing
456 by tracking through a tortuous path fixture (as described in Section III.G(1)b). Please note that
457 this testing should be conducted on all tips that are joined or bonded to the catheter by any
458 means, regardless of tip length. If the tip is not long enough to be gripped for tensile testing,
459 modifications to the manufacturing (i.e., longer tip joined by same method) or test methods (i.e.,
460 alternate or modified grip) should be employed.

461 **i. Flexibility and Kink Test**

462 Significance: Catheters may be subjected to tight angulations in tortuous vasculature during use.
463 Inability to withstand flexural forces that are typical of clinical use could lead to device failure or
464 vessel damage.

465 Recommendation: We recommend that you conduct testing which demonstrates that the catheter
466 will not kink at a bend radius that is appropriate for the intended anatomy. For example, we
467 recommend that you consider wrapping the catheter around a series of mandrels with
468 successively smaller radii until the catheter kinks, the lumen collapses, or the device shows no
469 kinking at a radius smaller than what could be considered worst-case for the intended anatomy.
470 This testing should be conducted along the full length, or representative portions, of the catheter
471 without the use of a guidewire as this would indicate a worst-case scenario (or a rationale should
472 be provided if a guidewire is used). We also recommend you provide the clinical basis for your
473 acceptance criterion. This could include literature or testing demonstrating the proposed criterion
474 is appropriate in representative angulations for the intended anatomy. Assessment of the kink
475 resistance of your device during simulated use alone is not considered a worst-case assessment as
476 it does not challenge the device to failure. This should be considered supporting information.

477 **j. Torque Strength**

478 Significance: Catheters may be subjected to torsional forces during use. Even non-fixed wire
479 catheters could be subject to torsional forces if the tip is inadvertently caught on a stent, calcified
480 lesion, etc. Inability to withstand torsional forces that are typical of clinical use could lead to
481 device failure or vessel damage.

482 Recommendation: We recommend that you assess the ability of the catheter to withstand
483 torsional forces when the distal tip is not free to rotate by rotating the proximal end of the
484 catheter until failure. We recommend that you precondition catheters prior to evaluating torque
485 strength by tracking through a tortuous path fixture, as described in Section III.G(1)b. We also
486 recommend that you test the torque strength of the catheter in the simulated-use fixture by
487 tracking through the fixture and then clamping the distal end and rotating the proximal end. We
488 recommend that you report the number of rotations to failure and the failure mode for each
489 sample tested. Alternatively, it may be possible to test the device to a specific number of turns
490 (i.e., not to failure) if the pre-determined acceptance criterion is established as worst-case
491 compared to clinical use.

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492 **k. Radiopacity**

493 Significance: Insufficient radiopacity may impede safe and reliable delivery of the balloon to the
494 intended location as it will not be clearly visible during use.

495 Recommendation: We recommend that you demonstrate that the radiopaque markers/materials
496 on the balloon catheter can be seen under typical fluoroscopic methods. We recommend that you
497 provide a qualitative or quantitative measure of radiopacity, wherein the balloon catheter is
498 visible using real-time and plain film x-ray. It is acceptable to provide images from animal
499 studies, *in vitro* phantoms, or equivalent models in order to support the visibility/radiopacity of
500 your device. If these data are leveraged from animal or bench testing, please provide a reference
501 in the submission to where the images can be located. The methods described in *ASTM F640-12*:
502 *Standard Test Methods for Determining Radiopacity for Medical Use* are generally considered
503 acceptable.

504 **l. Coating Integrity**

505 Significance: Coatings are intended to improve the performance of the device. Delamination or
506 degradation of a coating may lessen its benefit or otherwise negatively impact its clinical
507 performance and patient safety, e.g., causing embolization downstream.

508 Recommendation: Coating integrity testing should be conducted if your device has any coating
509 along the length of the catheter and/or on the balloon portion of the device. We recommend that
510 you address the aspects described below for any coatings applied to the surfaces of your product.

511 *Coating Description*

512 We recommend that you describe the clinical purpose and intended function of the coating, such
513 as enhanced radiopacity, thromboresistance, or lubricity. We also recommend that you describe
514 the physical structure of the coating, such as coating thickness, and indicate its chemical
515 identification.

516 *Test Samples*

517 You should conduct all testing on the finished product that was subjected to all manufacturing
518 processes, including sterilization. You should provide a scientific or statistical justification for
519 the sample size for each test. We recommend that you implement a sampling plan to examine
520 multiple lots of product (≥ 3) to assess both inter- and intra-lot variability. You should perform
521 testing on the extremes (i.e., “four corners”) and an appropriate intermediate size for the entire
522 product matrix proposed, as depicted in Table 5.

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Table 5: Example of “Four Corners Plus Intermediate” Test Matrix.

Balloon Diameter (mm)	Balloon Length (mm)				
	40	60	80	100	120
4.0	X				X
4.5					
5.0			X		
5.5					
6.0	X				X

525 It may be possible to combine coating integrity testing and particulate evaluation (Section
526 III.G(1)m) with simulated use testing (Section III.G(1)b), but you should take care to ensure that
527 only minimal additional handling of the sample is required for the coating integrity evaluation
528 such that particulates are neither lost nor generated.

529 *Interpretation of Data*

530 Coating integrity is considered a characterization test. While acceptance criteria do not need to
531 be included in the premarket submission, descriptions of visualization criteria for the assessment
532 (e.g., no voids, no cracks) should be provided. Furthermore, you should provide an interpretation
533 of the analysis.

534 Test reports should include a detailed discussion of the morphology of the coated surfaces. If
535 numerous defects are observed, quantifying defects using microscopy may be helpful. This may
536 include counting the number of total defects per unit area or measuring the total representative
537 defect area. You should support your discussion with representative color images, including any
538 areas with observed defects, at a sufficient magnification to characterize the defects. Multiple
539 magnifications may be warranted to visualize and adequately characterize the product. If the
540 coating is difficult to visualize (e.g., clear hydrophilic coating), measures should be taken in
541 order to ensure proper visualization (e.g., dyeing). The discussion of acceptable coating integrity
542 should include a justification that the number, size, and/or total area of defects observed will not
543 impact clinical performance or safety. Side-by-side testing with a predicate device may be
544 helpful to support substantial equivalence for 510(k) devices.

545 We recommend that you address the aspects described below for any coatings applied to the
546 surfaces of your product.

547 *Baseline Coating Integrity*

548 We recommend that you conduct a visual assessment of the coating integrity on all appropriate
549 surfaces of the final catheter to establish a baseline for comparison to coating characteristics after
550 testing performed after simulated use. If the coating is present on the balloon surface, unfolding
551 or partially inflating the device may be necessary to characterize coating at different locations.
552 We recommend that you appropriately quantify characteristics such as continuity and voids in
553 the coating, as described above.

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554 *Simulated Use Coating Integrity*

555 We recommend that you evaluate the coating integrity via visual assessment after simulated use.
556 Catheters should be tracked through an aqueous, tortuous path fixture (as described in Section
557 III.G(1)b) and then expanded in the aqueous medium to the maximum labeled diameter
558 described in the instructions for use prior to visual inspection.

559 We recommend you test coating integrity under the worst-case conditions of use. For example,
560 for balloons intended for ISR or post-deployment stent expansion, we recommend that you
561 evaluate the coating integrity after tracking the device through a tortuous path fixture and
562 inflating to the largest labeled diameter within a stent which has been deployed in the mock
563 vessel.

564 *Functional Testing*

565 We recommend you demonstrate that the coating can achieve its intended function. For example,
566 if a coating is intended to provide lubricity to the catheter, it may be helpful to demonstrate that
567 the frictional forces are decreased or at least equivalent to similar products with similar coatings.
568 For this type of assessment, we recommend that you characterize the drag force of the coating
569 (e.g., pinch test) after the samples are prepared per the instructions for use.

570 **m. Particulate Evaluation (Coated Devices Only)**

571 Significance: Particulate matter can be generated by the manufacturing process, environment, or
572 from the breakdown of any coating (e.g., hydrophilic coating) on the catheter or from the device
573 packaging. If particles are introduced in the bloodstream during an angioplasty procedure, they
574 may present an embolic risk to the patient. Measurement of the total quantity and size of
575 particulates a device may generate is an indication of embolic risk. Due to lower embolic risks of
576 peripheral devices as compared to other vasculatures, if the coating and substrate are not novel
577 and coating integrity testing has been conducted with acceptable results, a particulate evaluation
578 may not be needed. However, this testing should be conducted if these factors have not been met,
579 or to further support the coating integrity of your device.

580 Recommendation: We recommend that you measure the total quantity and size of the particulates
581 generated during the simulated use of your device, addressing the aspects described below.

582 *Test Samples*

583 We recommend conducting all testing on the finished product that was subjected to all
584 manufacturing processes, including sterilization. A scientific or statistical justification for the
585 sample size should be provided. We recommend that you implement a sampling plan to examine
586 multiple lots of product (≥ 3) to assess both inter- and intra-lot variability. You should perform
587 testing on the extremes and an appropriate intermediate size for the entire product matrix
588 proposed (i.e., “four corners” and intermediate size matrix; see Table 5).

589 It may be possible to combine the particulate evaluation and simulated use coating integrity
590 testing (Section III.G(1)l) with simulated use testing (Section III.G(1)b), but you should take care

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591 to ensure that only minimal additional handling of the sample is required for the coating integrity
592 evaluation such that particulates are neither lost nor generated.

593 *Interpretation of Data*

594 Particulate testing should be conducted as part of your design verification testing and should not
595 be for characterization only. A rationale for the criteria used as well as a discussion of the results
596 should also be provided. The discussion of acceptable particulate evaluation should include a
597 justification that the number and size of particulates will not impact safety or clinical
598 performance. This may include a reference to any applicable standards or the use of side-by-side
599 testing with a legally marketed device (e.g., predicate device) demonstrating equivalent results.

600 *Test Methods*

601 We recommend that you evaluate particulate generated by the entire PTA system, including
602 accessory devices expected to be used during a clinical procedure. Catheters should be tracked
603 through an aqueous, tortuous path fixture (as described in Section III.G(1)b) and then expanded
604 in an aqueous medium to the maximum labeled diameter described in the instructions for use
605 prior to visual inspection. When deployed, the balloon should be in direct contact with the
606 simulated vessel without the use of other coatings, lubricants, sheaths, or protective wraps
607 between the balloon and the simulated vessel. To ensure measurement of the total number of
608 particulates that could be potentially introduced into the bloodstream, the catheter should be
609 inserted into the test fixture to the extent at which it would be inserted in clinical use. The total
610 number of particulates, including those from the catheter and accessory devices, should be
611 reported in each of three size ranges: $\geq 10\mu\text{m}$, $\geq 25\mu\text{m}$, and at the largest size for which validation
612 yields $\geq 75\%$ recovery. At a minimum, the largest size should be $\geq 50\mu\text{m}$. Appropriate precautions
613 should be taken to ensure that the particles are suspended during sampling for particle counting
614 and sizing to minimize artifacts from the test system.

615 We recommend that you perform particulate evaluation under the worst-case conditions of use.
616 For example, for balloons intended for ISR or post-deployment stent expansion, we recommend
617 that you evaluate the quantity and sizes of particulates generated from tracking the device
618 through the tortuous path fixture (as described in Section III.G(1)b) and inflating to the largest
619 labeled diameter within a stent which has been deployed in the mock vessel.

620 *Method Validation*

621 You should describe and validate particle counting and sizing methods. Validation should be
622 conducted using particulate standards of known quantity and size. They should be introduced
623 into your model and counting apparatus in a similar manner as the device would be introduced
624 clinically. The percent recovery, or accuracy, should be determined and meet the criteria
625 described above. For a system to be considered validated, $\geq 90\%$ recovery should be
626 demonstrated for the $\geq 10\mu\text{m}$ and $\geq 25\mu\text{m}$ size ranges. Please note that recovery rates well above
627 100% would not be considered valid.

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629 *Acceptance Criteria*

630 Particulate testing should be conducted as part of your design verification testing and should not
631 be for characterization only. Therefore, specific criteria should be established, justified, and met.
632 If large amounts of particulates are shed, it may be important to demonstrate comparability to a
633 legally-marketed predicate device used in the same target vasculature or provide evidence of
634 safety through your animal studies (with appropriate downstream assessments). A scientific
635 rationale should be provided to support the particulate acceptance criteria that is used.

636 *Particulate Chemical Identification*

637 Particulate matter can be generated from numerous sources, including the manufacturing process
638 and/or environment contamination, from the breakdown of any coating on the catheter, or from
639 the device packaging. It is important to establish that a significant number of particulates are not
640 being introduced from other unintended sources, as described above, which may present an
641 embolic risk. Therefore, if a large amount of particulates are shed from your device, it may be
642 pertinent to conduct additional analysis, such as a chemical characterization of the particulates,
643 in order to determine their source. For this testing, FDA recommends that you perform chemical
644 identification of representative particulate populations and report the results in relative amounts
645 (percentages). Chemical characterization of captured particulates for identity can be
646 accomplished through a variety of methods including energy-dispersive x-ray spectroscopy
647 (EDX), Fourier transform infrared (FTIR) spectroscopy, Raman spectroscopy, mass
648 spectroscopy, or diffraction techniques.

649 Chemical identification of representative particulate material should be performed with
650 justification for the method and sample analyzed. The sample should be sufficiently large in
651 order to ensure that the particulates assessed are representative of the particulates that would be
652 generated during the deployment of the device. The method used should be capable and
653 sufficient for chemical identification. Specific details regarding the capture and analysis (e.g.,
654 how the samples were filtered, color images of the filters, how the samples were chosen, details
655 regarding the number of particulates analyzed as compared to the total particulates filtered) of
656 the particulates should be provided.

657 There are certain instances when providing additional supporting analyses may allow for reduced
658 (e.g., smaller sample size, fewer particulates analyzed) or omitted chemical identification testing.
659 Supporting analyses could include any or all of the following:

- 660 • particulate quantitation studies with the uncoated balloon catheter manufactured in the
661 identical way as the coated device but including potential inclusion of a “dummy”
662 coating process, demonstrating sufficiently low amounts of particulates;
- 663 • a discussion regarding the potential interactions of your coating, including all
664 components, with the catheter materials and their potential to introduce some of the
665 catheter extractables/leachables into the particulates;

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- 666 • representative color images of the particulates captured on the entire filter demonstrating
667 no concerning information (e.g., unexpected appearance);
- 668 • a risk assessment regarding potential contaminants and the coating chemical
669 compositions;
- 670 • a discussion of the animal studies data indicating no concerning downstream or embolic
671 events; and
- 672 • a discussion and references to any historical clinical data indicating no concerning
673 embolic events.

(2) Additional Tests for Catheters Intended for Infusion of Contrast Media or Other Fluids

a. Catheter Body Burst Pressure

677 Significance: The catheter body should be designed to withstand pressures typically needed to
678 achieve contrast media flow rates used in clinical practice. Inability to withstand pressures that
679 are typical of clinical use could lead to device failure or vessel damage.

680 Recommendation: We recommend that you determine the maximum pressure that the catheter
681 body can withstand during injection. We recommend you conduct the testing under clinical use
682 conditions (i.e., including use of a syringe, automatic injector). The contrast medium or fluid
683 should be representative of worst-case clinical conditions. We also recommend you provide the
684 clinical basis for your acceptance criteria.

b. Infusion Flow Rate

686 Significance: The catheter should be designed to achieve clinically acceptable contrast media
687 flow rates. Inability to achieve acceptable flow rates could lead to user error and adverse clinical
688 consequences.

689 Recommendation: We recommend that you conduct testing that demonstrates that the catheter is
690 capable of achieving clinically acceptable contrast media flow rates. We recommend that testing
691 be conducted at maximum catheter burst pressures (as identified in Section III.G(2)a) as well as
692 pressures typical of clinical use. We recommend that you report the maximum flow rate in the
693 device labeling. We also recommend you provide the clinical basis for your acceptance criteria.

(3) Additional Tests for Catheters Intended for In-Stent Restenosis (ISR) Use or for Stent Expansion following Stent Deployment

697 If you label a PTA catheter for ISR use or for stent expansion immediately following stent
698 deployment (for purposes of securing the stent to the vessel wall and ensuring that the stent is
699 completely deployed), we recommend you conduct balloon rated burst pressure and fatigue

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700 testing within an expanded stent (see Sections III.G(1)c and III.G(1)d). If the balloon has a
701 coating on it, we also recommend conducting coating integrity and particulates testing in a
702 simulated use model that includes an expanded stent (see Sections III.G(1)l and III.G(1)m).

703 **(4) Additional Tests for Scoring/Cutting Balloons**

704 Scoring and cutting balloons concentrate the dilating forces along the scoring elements or
705 atherotomes. Due to the additional design features, scoring and cutting balloons have additional
706 considerations beyond a standard PTA catheter.

707 **a. Scoring/Cutting Mechanism Securement**

708 Significance: Detachment of the scoring/cutting mechanism(s), whether wire or atherotomes,
709 could result in device failure, vessel damage, and/or embolic risk due to device remnants within
710 the vasculature.

711 Recommendations: We recommend that you determine the force (e.g., tensile, shear) at which
712 the bonding of the scoring/cutting mechanism fails. We recommend you provide the clinical
713 basis for your test method and acceptance criteria based on the type and level of risk.

714 **b. Scoring/Cutting Performance**

715 Significance: The scoring/cutting mechanism of the device introduces additional risks, such as
716 vascular damage, as compared to a standard PTA catheter. Failure to achieve adequate scoring or
717 cutting could lead to the device not performing as intended.

718 Recommendations: We recommend that you demonstrate that the device can score a lesion, as
719 intended. Performance of your device should be evaluated in a calcified bench model, animal
720 model with calcified lesions, cadaveric model, and/or clinical study and compared to a legally
721 marketed predicate device. We encourage you to contact the FDA early to discuss the proposed
722 model to evaluate the scoring/cutting performance (see FDA guidance “[Requests for Feedback
and Meetings for 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).”)¹⁰

724 **c. Substantially-Equivalent Safety Outcomes (Demonstration of 725 No Added Risks)**

726 Significance: If a scoring/cutting balloon catheter has novel technological characteristics (i.e.,
727 scoring/cutting mechanism that is different from the standard scoring wire or cutting atherotomes
728 of the predicate device), additional safety questions may arise, such as added risk of vessel
729 dissection or perforation.

730 Recommendations: If different technological characteristics as compared to the predicate are
731 used to achieve the intended function, we recommend that you assess whether the safety
732 outcomes (i.e., scoring depth, perforation/dissection rate) of your device are substantially

¹⁰ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/requests-feedback-and-meetings-medical-device-submissions-q-submission-program>

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733 equivalent to those of the identified predicate, using the predicate device as the control in an
734 animal model and/or clinical study.

735

736 H. Animal Safety and Performance Testing

737 Significance: Animal testing is generally recommended to evaluate the *in vivo* safety of some
738 specialty catheters and potentially some PTA balloon catheters, particularly for new designs,
739 significant device modifications, and new indications for use. An example of this is for a scoring
740 balloon with a new cutting mechanism.

741
742 Recommendation: Animal testing of PTA balloon catheters and specialty catheters should
743 address factors that cannot be evaluated through bench tests or in a clinical study. The study
744 design and endpoints should be based upon the mechanism of action of the device and mitigation
745 of risk.

746
747 FDA supports the principles of the “3Rs,” to reduce, refine, and replace animal use in testing
748 when feasible. You should consider the best practices for the development, conduct and
749 presentation of these animal studies while incorporating modern animal care and use strategies.

750
751 We encourage manufacturers to take advantage of the Q-Submission Program to ensure that the
752 animal study protocol addresses safety concerns and contains elements which are appropriate for
753 a regulatory submission (e.g., the study should be performed under Good Laboratory Practice
754 (GLP) regulations as stated in 21 CFR 58 at an animal study facility with appropriate licensure
755 and accreditations).¹¹ In addition, if you are proposing to use a non-animal testing method that
756 you believe is suitable, adequate, validated, and feasible, we recommend that you discuss the
757 proposal using the Q-Submission Program. We will consider if such an alternative method could
758 be assessed for equivalency to an animal test method. For details on the Q-Submission Program,
759 please refer to the guidance “[Requests for Feedback and Meetings for Medical Device
760 Submissions: The Q-Submission Program](#).”¹²

761
762 For devices with notable dissimilarity from legally-marketed PTA devices (e.g., new indications,
763 designs, technology), we recommend that you conduct animal testing to confirm safety of the
764 procedure, to evaluate the functional characteristics of the device design, and to assess the
765 performance of the PTA or specialty catheter.

766
767 For scoring balloons, we strongly recommend animal testing to demonstrate equivalent safety
768 outcomes for all scoring/cutting devices, as compared to their predicate, , especially when the
769 technological characteristics differ. We recommend that you evaluate these devices in an

¹¹ See also FDA Guidance “General Considerations for Animal Studies for Cardiovascular Devices”
(<https://www.fda.gov/regulatory-information/search-fda-guidance-documents/general-considerations-animal-studies-cardiovascular-devices-guidance-industry-and-fda-staff>).

¹²<https://www.fda.gov/regulatory-information/search-fda-guidance-documents/requests-feedback-and-meetings-medical-device-submissions-q-submission-program>

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770 appropriate animal model that closely approximates the intended use of the device in humans and
771 that you provide a supporting rationale for the chosen animal model in your submission. The
772 predicate device should be used as a control in these studies. We strongly recommend that these
773 studies be conducted in accordance with 21 CFR part 58 or explain why the noncompliance
774 would not impact the validity of the study data provided to support a substantial equivalence
775 determination.
776

777 **I. Clinical Performance Testing**

778 Clinical evidence is generally unnecessary for most PTA balloon and specialty catheters;
779 however, such testing may be requested in situations such as the following:

- 780 • indications for use dissimilar from legally marketed devices of the same type (e.g.,
781 treatment of specific diseases or lesion types);
- 782 • new technology (i.e., technology different from that used in legally marketed devices of
783 the same type); and
- 784 • cases where engineering and/or animal testing raise issues that warrant further evaluation
785 with clinical evidence.

786 If a clinical study is needed to demonstrate substantial equivalence, i.e., conducted prior to
787 obtaining 510(k) clearance of the device, the study should generally be conducted under the
788 Investigational Device Exemptions (IDE) regulation, 21 CFR 812. Generally, we believe PTA
789 balloon catheters and specialty catheters addressed by this guidance document are significant risk
790 devices subject to all requirements of 21 CFR part 812. Please see the FDA guidance,
791 “[Significant Risk and Nonsignificant Risk Medical Device Studies](#).”¹³ In addition to the
792 requirements of 21 CFR part 812, sponsors of such trials must comply with the regulations
793 governing institutional review boards (21 CFR part 56) and informed consent (21 CFR part 50).
794 When data from clinical investigations conducted outside the United States are submitted to
795 FDA for PTA and specialty catheters, the requirements of 21 CFR 812.28 may apply.¹⁴ 21 CFR
796 812.28 outlines the conditions for FDA acceptance of clinical data from investigations conducted
797 outside the US when submitted to support premarket submissions. For more information, see the
798 FDA guidance, “[Acceptance of Clinical Data to Support Medical Device Applications and
799 Submissions: Frequently Asked Questions](#).”¹⁵

¹³ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/significant-risk-and-nonsignificant-risk-medical-device-studies>

¹⁴ Applies to data from clinical investigations that began on or after February 21, 2019 and are submitted to support a premarket submission, including IDEs, PMAs, and 510(k)s.

¹⁵ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/acceptance-clinical-data-support-medical-device-applications-and-submissions-frequently-asked>

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800 In some cases, “real-world data” (RWD) may be used to support expansion of the indication for
801 a device for which 510(k) clearance has already been obtained. Whether the collection of RWD
802 for a legally-marketed device requires an IDE depends on the particular facts of the
803 situation. Specifically, if a cleared device is being used in the normal course of medical practice,
804 an IDE would likely not be required. For additional information regarding this topic, please refer
805 to the FDA Guidance entitled “[Use of Real-World Evidence to Support Regulatory Decision-
806 Making for Medical Devices.](#)”¹⁶
807

808 **J. Labeling**

809 The regulatory submission must include proposed labeling in sufficient detail to satisfy the
810 requirements of 21 CFR 807.87(e) for premarket notification and 21 CFR 814.20(b)(10) for
811 premarket approval submissions. Labeling for PTA balloon catheters and specialty balloons
812 should include all applicable information, including indications, contraindications, warnings,
813 product information, a summary of the clinical data (if applicable), and directions for use.

814 As prescription devices, PTA balloon and specialty catheters are exempt from having adequate
815 directions for lay use under section 502(f)(1) of the Federal Food, Drug, and Cosmetic Act
816 (FD&C Act) (21 U.S.C. 352(f)(1)) as long as the conditions in 21 CFR 801.109 are met. For
817 instance, labeling must include adequate information for practitioner use of the device, including
818 indications, effects, routes, methods, frequency and duration of administration and any relevant
819 hazards, contraindications, side effects and precautions. (21 CFR 801.109(d)).

820 **K. Modifications**

821 In accordance with 21 CFR 807.81(a)(3), a device change or modification “that could
822 significantly affect the safety or effectiveness of the device” or represents “a major change or
823 modification in the intended use of the device” requires a new 510(k). The changes or
824 modifications listed below would likely require submission of a new 510(k). Note that this list is
825 not exhaustive but provides examples of modifications that are likely to require submission of a
826 new 510(k). For additional details, please see FDA guidance “[Deciding When to Submit a 510\(k\)
827 for a Change to an Existing Device.](#)”¹⁷
828

829 Such changes or modifications include:

- 830 • Change in device dimensions: FDA considers this change to be a modification in design.
831 FDA has determined that this change could significantly affect safety and effectiveness of
832 the device as it may alter the device performance. Thus, if dimensional changes are not
833 within the range that was previously cleared, testing may be needed to support the
834 change. The magnitude and criticality of the modified dimension should be considered.

¹⁶ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/use-real-world-evidence-support-regulatory-decision-making-medical-devices>

¹⁷ <https://www.fda.gov/regulatory-information/search-fda-guidance-documents/deciding-when-submit-510k-change-existing-device>

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835 • Change to indirect or direct blood contacting components: FDA considers this change to
836 be a modification in material. FDA has determined that this change could significantly
837 affect safety and effectiveness of the device by altering engineering attributes and/or
838 introducing different types or quantities of residual chemicals, which could result in a toxic
839 response. Therefore, a change in the material could impact device performance and
840 biocompatibility, which could impact patient safety.

841 • Change in sterilization technique: FDA considers this to be a significant change. FDA has
842 determined that this change could significantly affect safety and effectiveness of the
843 device as it could impact device sterility and biocompatibility. For example, changes to
844 an ethylene oxide sterilization process may leave increased ethylene oxide residuals.
845 Additionally, changes in sterilization may unintentionally affect device materials, which
846 could consequently affect the safety and/or performance of the device.

847 Examples of changes or modifications in the indications for use of the device that would likely
848 require a new 510(k) are:

- 849 • a change in specific lesion characteristics (e.g., chronic total occlusion, ISR); and
- 850 • claims in improvement of outcomes in other technologies (e.g., pre-treatment with
851 scoring balloons improves outcomes of drug-coated balloons).

852 We believe that the following modifications will likely not require submission of a new 510(k):

- 853 • Minor changes in packaging: A minor change in packaging (e.g., replacing hardcopy
854 instructions for use with an electronic version, update to the expiration date) is not
855 expected to impact device safety and performance.
- 856 • Increase in shelf-life: An increase in device shelf-life is not expected to impact device
857 safety and performance as long as the testing protocol has been previously reviewed and
858 accepted in a prior submission. Additionally, the test results should fall within the
859 acceptance criteria previously found to be acceptable.

860