

## BRIEFING

**<1381> Assessment of Elastomeric Components Used in Injectable Pharmaceutical Product Packaging/Delivery Systems**, *PF* 43(3) [May–June 2017]. The previous proposal for this chapter, published in *PF* 43(3) under a different title, *Elastomeric Evaluation of Elastomeric Components Used in Pharmaceutical Packaging/Delivery Systems* was canceled and an updated version is being proposed. The Packaging and Distribution Expert Committee is enacting this new general chapter to support the planned revisions to *Elastomeric Closures for Injections* (381), which are also proposed in this issue of *PF*. This new chapter:

1. Describes elastomeric components and their materials of construction for use in pharmaceutical product packaging/delivery systems.
2. Provides a high-level introduction to elastomer chemistry, manufacturing technology, and the post-processing of components.
3. Designates and expands on baseline requirements.
4. Discusses identification testing.
5. Describes requirements and responsibilities.
6. Provides a summary of required and recommended test methods.

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**Add the following:**

**^<1381> ASSESSEMENT OF ELASTOMERIC COMPONENTS  
USED IN INJECTABLE PHARMACEUTICAL PRODUCT  
PACKAGING/DELIVERY SYSTEMS**

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#### 1. INTRODUCTION

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Risk to drug product quality may exist when elastomeric components come into direct or indirect contact with pharmaceutical products. Elastomeric components used in pharmaceutical packaging/delivery systems must be proven suitable for their intended use based on aspects of protection, compatibility, and performance.

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Tests and specifications applicable to elastomeric components used in packaging/delivery systems for injectables are referenced in conjunction with *Injections and Implanted Drug Products* (1).

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*Elastomeric Closures for Injections* (381) presents a body of test procedures and acceptance criteria for elastomeric components employed in injectable product packaging/delivery systems. The biological reactivity tests and physicochemical tests provide baseline information on a component's suitability. The test methods and acceptance criteria are detailed in their description but lack an explanation as to their relevance to the composition of the elastomers. Due to the large and diverse nature of the pharmaceutical marketplace, it may not be intuitive to stakeholders as to the proper use and application of (381). Therefore, a primary purpose of this chapter is to communicate the key concepts that form the foundation of (381) and to establish and clarify its application and applicability.

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Beyond the baseline requirements provided in (381), elastomers will need to be qualified for intended use commensurate with the level of risk to drug product quality. These evaluations would encompass studies for extractables and leachables. Recommendations for conducting these studies are found in *Assessment of Extractables Associated with Pharmaceutical Packaging/Delivery Systems* (1663) and *Assessment of Drug Product Leachables Associated with Pharmaceutical Packaging/Delivery Systems* (1664).

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Finally, elastomeric component functional suitability as part of the finished product packaging/delivery system must be demonstrated. Relevant information is found in *Elastomeric Component Functional Suitability in Parenteral Product Packaging/Delivery Systems* (382) and *Assessment of*

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57 *Elastomeric Component Functional Suitability in Parenteral Product*  
58 *Packaging/Delivery Systems*(1382).

59 **2. SCOPE**

60 This chapter serves as a supplement to the elastomeric component  
61 baseline requirements in (381). This chapter seeks to 1) describe elastomeric  
62 components and their materials of construction for use in pharmaceutical  
63 packaging/delivery systems; 2) provide a high-level introduction to  
64 elastomer chemistry, manufacturing technology, and the post-processing of  
65 components; 3) designate and expand on baseline requirements; 4) discuss  
66 identification testing; 5) discuss test requirements and responsibilities; and  
67 6) provide a summary of required and recommended test methods.

68 Elastomeric components utilized for injectable pharmaceutical products  
69 within the scope of this chapter include, but are not limited to, those used  
70 for vials and bottles (stoppers and cap liners), prefilled syringes (plungers,  
71 needle shields, and tip caps), cartridges (plungers and seal liners), flexible  
72 bags (injection ports), and blow-fill-seal containers (cap liners). Also within  
73 the scope are elastomeric components of systems or packages that are  
74 intended for transient product storage and/or delivery intended for specific  
75 pharmaceutical products (e.g., the elastomeric components of an infusion  
76 set or a single-use syringe included as part of a co-packaged combination  
77 product or linked by way of labeling for use with a specific pharmaceutical  
78 product). Components of similar systems intended for general product use  
79 are out of the scope of this chapter. All elastomeric components in direct or  
80 indirect contact with the pharmaceutical product are within the scope.

81 **3. DESCRIPTION OF ELASTOMERIC COMPONENTS IN PACKAGING SYSTEMS**

82 Certain components used in pharmaceutical packaging/delivery systems  
83 must have elastic properties for the system to function properly. Elastomers  
84 are a unique family of polymers with properties including the ability to  
85 recover from being stretched or deformed beyond their original state. This  
86 allows components to be flexible, maintain a seal, and be able to reseal after  
87 puncturing. In the following sections, typical elastomer materials,  
88 compositions, and physical attributes are summarized. Various compounding  
89 ingredients, including curing systems, are required to produce optimal  
90 elastomeric performance. In thermoset elastomers, the physical attributes  
91 will depend on polymer cross-linking, achieved during the vulcanization  
92 process. The cured polymer will produce a chemical reaction and by-  
93 products that will impact the chemical makeup of the elastomeric  
94 formulation. The compounding ingredients, reaction by-products, and post-  
95 processing effects, such as sterilization, will influence the outcome of the  
96 component's chemical characterization.

97 Given the complex nature of packaging systems and their manufacturing  
98 and development processes, multiple testing procedures are needed to  
99 establish their suitability for use with a specific pharmaceutical product. The  
100 logical manufacturing and development process for packaged drug  
101 products—starting with the packaging system’s materials of construction,  
102 continuing with the packaging system itself, and ending with the packaged  
103 drug product—forms the basis of the following three-stage approach to  
104 packaging/delivery systems qualification:

- 105 • **Component screening:** The baseline requirements for biological  
106 reactivity and physicochemical testing described in this chapter.
- 107 • **Controlled extraction studies:** Studies as described in (1663) to  
108 create extractables profile(s) of particular pharmaceutical  
109 packaging/delivery systems, packaging components, or materials of  
110 construction.
- 111 • **Pharmaceutical product assessment:** Actual-case measurement of  
112 confirmed leachables in the pharmaceutical product in the  
113 packaging/delivery system intended for the commercial market. (For  
114 additional information, see (1664)).

115 Assessment of elastomeric component functionality is performed within the  
116 context of the intended product packaging system. (For additional  
117 information, see (382) and (1382).)

#### 118 4. ELASTOMERIC COMPONENTS—MATERIALS OF CONSTRUCTION

##### 119 4.1 Thermoset and Thermoplastic Elastomeric Components

###### 120 4.1.1 THERMOSET COMPOSITION (TYPICAL)

121 A thermoset elastomer is a polymer system in which the elastomeric  
122 properties are derived from chemical cross-linking that is irreversible. This  
123 cross-linking is created between a curative (cross-linking agent) and a  
124 polymer (elastomer) when the materials are subjected to heat and pressure.  
125 In thermoset elastomers de-cross-linking is not possible without destruction  
126 of the material. Other ingredients that typically are part of the thermoset  
127 rubber formulation are shown in [Table 1](#).

128 **Table 1. Thermoset Elastomeric Components: Typical Rubber**  
129 **Ingredients**

| Ingredient                       | Function  |
|----------------------------------|---|
| Polymers (elastomers)            | Provide elastic properties after curing             |
| Curatives (cross-linking agents) | Form cross-links to provide elasticity and strength |

| Ingredient                | Function  |
|---------------------------|---|
| Fillers/extenders         | Impart hardness, modulus/deformation, strength, reinforcement     |
| Processing aids           | Impart flexibility, fatigue resistance, mold flow                 |
| Antioxidants/antiozonants | Provide stabilization, i.e., protection against UV, oxygen, ozone |
| Plasticizers              | Aid processing by providing flexibility, hardness                 |
| Dyes/coloring materials   | Impart color  |

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4.1.2 POLYMER TYPES AND ATTRIBUTES

Typical polymers used in elastomeric components for packaging/delivery systems for injections, along with characteristic physical attributes and component examples, are shown in [Table 2](#).

**Table 2. Typical Elastomers for Thermoset Elastomeric Components**

| Elastomer   | Physical Attributes   | Typical Components  |
|---|---|---|
| Isobutylene/isoprene copolymer (butyl) (IIR);<br>Brominated isobutylene isoprene (BIIR);<br>Chlorinated isobutylene isoprene (CIIR);<br>Brominated isobutylene para methylstyrene terpolymer (BIMS) | Gas barrier;<br>Aging resistance  | Stoppers, plungers, lined seals   |
| Natural polyisoprene (NR); <sup>a</sup> Synthetic polyisoprene (IR)   | Good coring and reseal behavior;<br>Abrasion resistance;<br>Higher gas permeability | Stoppers, plungers for single-use syringes, septa, needle shields, tip caps |
| Styrene butadiene rubber (SBR)  | Higher gas permeability   | Needle shields, tip caps  |
| Ethylene propylene diene monomer rubber (EPDM)  | Chemical resistance;<br>Aging resistance  | O-rings   |
| Acrylonitrile butadiene rubber (nitrile) (NBR)  | Chemical resistance (e.g., mineral oils)  | Stoppers, plungers, gaskets, O-rings  |

| Elastomer   | Physical Attributes | Typical Components                           |
|---|---------------------|--|
| Polychloroprene (neoprene) (CR); Epichlorohydrin (ECO/CO) | Chemical resistance | O-rings, gaskets                             |
| Polysiloxane (VMQ,PMQ,PVMQ)                               | Heat resistance     | Stoppers, plungers, tubing, gaskets, O-rings |

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<sup>a</sup> NR must be labeled appropriately per CFR 21:<http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm070929.pdf>

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#### 4.2 Thermoplastic Composition (Typical)

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A thermoplastic elastomer (TPE) is a polymer system with a different nature. Elasticity is incorporated in a different way (see below), and, more importantly, thermoplastic elastomers have a thermal behavior that is comparable to plastics. Thermoplastic elastomers when heated lose their elasticity and become deformable like a plastic. Cooled down again, they regain their elasticity. This cycle can be repeated. Unlike thermoset elastomers, the elastic behavior of thermoplastic elastomers is reversible.

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TPEs, like thermoset elastomers, owe their elasticity to a polymer network that has a certain degree of structure. However, for thermoset elastomers, the structure occurs through chemical cross-linking with covalent bonding, whereas TPEs use a different mechanism. For example, in styrenic block copolymers, hard polystyrene blocks serve as physical cross-links in a three-dimensional network of softer chains of a different polymer such as polyisoprene or ethylene/butylene copolymer.

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In a different class of thermoplastic rubber named thermoplastic vulcanizates, a hard plastic phase and a soft elastomeric phase are present. The hard phase can use material such as polyolefin, while the soft phase is a thermoset rubber phase with a high degree of chemical cross-linking. The curing of the rubber phase is done in a process known as "dynamic vulcanization". During this process, chemical cross-linking takes place during the mixing of the elastomer with the vulcanization system and other ingredients, such as polyolefin. Additional ingredients that may be present in TPEs are fillers, antioxidants, antiozonants, and plasticizers.

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#### 4.3 Surface Coatings and Treatments

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After manufacturing, elastomeric components may not fulfill all the properties that are required for their application. An additional coating or surface treatment may be necessary. The most common surface treatment is siliconization, which is used to overcome the inherent tackiness of

169 components and to provide lubricity. Tackiness will negatively impact  
 170 component processing (e.g., sterilization, machinability at filling) and may  
 171 introduce permanent component deformation. Chlorination is a less  
 172 frequently used surface treatment. In this treatment, the component surface  
 173 is exposed to chlorine, resulting in non-tacky and somewhat lubricious parts.  
 174 In order to provide non-tackiness and lubricity, but at the same time reduce  
 175 extractable levels (i.e., barrier effect), components may be covered, on part  
 176 of their surface or on their entire surface, with lubricious barrier materials  
 177 that are applied either by coating or by film lamination. Polymeric coatings  
 178 and surface treatments are shown in [Table 3](#).

179 **Table 3. Surface Coatings, Films, and Treatments for Elastomeric**  
 180 **Components**

| Coating or Treatment   | Physicochemical Effect            | Typical Components            |
|--|-----------------------------------|-------------------------------|
| Silicone: silicone oil or emulsion, cross-linked silicone oil  | Non-tackiness, lubricity          | All components                |
| Chlorination   | Non-tackiness                     | Small, thin-walled components |
| Parylene   | Barrier, non-tackiness, lubricity | Stoppers, plungers            |
| Fluoropolymer coating  | Barrier, non-tackiness, lubricity | Stoppers, plungers            |
| Fluoropolymer lamination [ethylene tetrafluoroethylene (ETFE), fluorinated ethylene propylene (FEP)] | Barrier, non-tackiness, lubricity | Stoppers, plungers            |

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**4.4 Compounds of Concern**

183 Elastomeric components are made of various materials of construction.  
 184 Some of these materials may raise quality or safety concerns. It is  
 185 recommended that the user evaluate the presence of such materials in  
 186 components, either by direct use or by use in the manufacturing processes  
 187 of the materials of construction. An overview of such materials, together  
 188 with the associated concerns, is shown in [Table 4](#).

189 **Table 4. Elastomeric Components: Compounds of Concern**

| Compound of Concern   | Source   | Risk  |
|---|--|---|
| Latex   | Associated with compounds containing dry natural rubber or derivatives               | Associated with anaphylaxis in individuals allergic to natural rubber latex proteins                                  |
| Materials of animal origin  | Fatty acids and their metal salts used as processing aids or slip agents in polymers | Potential sources of transmissible spongiform encephalopathies (TSEs) in pharmaceutical products before manufacturing |
| 2-Mercapto-benzothiazole (MBT)  | Vulcanization accelerator used in the production of rubber                           | Potential carcinogen  |
| N-nitrosamines  | Associated with the use of certain secondary amines in the cure system               | Potential carcinogens   |
| Phthalates [e.g., orthophthalates such as bis(2-ethylhexyl) phthalate (DEHP)] | Used as a plasticizer or softener in polymers  | Associated with developmental, reproductive, and endocrine health effects   |
| Polycyclic aromatic hydrocarbons (PAHs)                                       | Associated with carbon black (colorant or reinforcing agent)                         | Potential carcinogens   |

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191 **Latex:** Contact with elastomers or packaging/delivery systems containing  
192 natural rubber latex, dry natural rubber, or synthetic derivatives of natural  
193 rubber latex has been associated with allergic reactions, including  
194 anaphylaxis, in individuals allergic to natural rubber latex proteins. There is  
195 no one threshold level of exposure that can be considered safe; factors  
196 include the exposure route and the immune status of the individual.  
197 Information regarding suitable labeling can be found in the FDA Guidance for  
198 *Industry Recommendations for Labeling Medical Products to Inform Users*  
199 *that the Product or Product Container is not Made with Natural Rubber*  
200 *Latex* (December 2, 2014).

201 **Materials of animal origin:** Pharmaceutical products having raw  
202 materials derived from animal sources have a risk of transmissible  
203 spongiform encephalopathy (TSEs). Cattle infected with bovine spongiform  
204 encephalopathy (BSE) can transmit to humans in the form of Creutzfeldt-  
205 Jakob disease (CJD) and the variant (vCJD). This disorder is caused by a

206 prion (protein) that causes the brain to be porous or spongy. Manufacturers  
207 should use materials from “non TSE-relevant animal species” or non-animal  
208 origin where possible to minimize the risk. When tallow is derived from TSE-  
209 relevant starting material, a rigorous manufacturing process should  
210 eliminate or reduce or any TSE risk.

211 **2-Mercapto-benzothiazole (MBT):** 2-Mercapto-benzothiazole (MBT) is  
212 used as a non-volatile vulcanization accelerator in the production of rubber.  
213 MBT reacts with zinc oxide and sulfur to cross-link rubber. It can be bonded  
214 in the vulcanizate or may be present as a substance incorporated in the  
215 polymer. The amount of free MBT in the final product will be dependent on  
216 the vulcanization conditions.

217 **N-nitrosamines:** N-nitrosamines are organic nitrogen-containing  
218 compounds that, under certain reaction conditions, are yielded as reaction  
219 products of nitrosating agents (e.g., certain oxides of nitrogen and  
220 nitrosatable secondary amines). Most secondary amines, but not all, may  
221 lead to carcinogens. There are 6 N-nitrosamines of concern that are listed  
222 in [Orally Inhaled and Nasal Drug Products \(1664.1\)](#).

223 **Phthalates:** Phthalates are a group of chemicals (orthophthalates and  
224 terephthalates) used in many polymer products to make plastics more  
225 flexible or as polymer-solvating agents. Phthalate esters are prepared by the  
226 esterification of phthalic anhydride with alcohols (C1/C2-C13). They are  
227 broadly divided into two distinct main groups with very different  
228 applications, toxicological properties, classification, and legal requirements.  
229 Orthophthalates used in food packaging are being reviewed by the FDA due  
230 to concerns for reproductive and endocrine health effects. More information  
231 can be found in the US *Federal Register* proposed rules (Vol. 83 No. 220,  
232 November 14, 2018).

233 **Polycyclic aromatic hydrocarbons:** Polycyclic aromatic hydrocarbons  
234 (PAHs), also known as polynuclear aromatic hydrocarbons (PNAs), are  
235 organic compounds with two or more fused aromatic (benzene) rings and  
236 are found as contaminants in oils used in elastomeric formulations and in  
237 carbon black used as a colorant or reinforcing agent. Carbon black is a  
238 material produced by the incomplete combustion of heavy petroleum  
239 products (i.e., ethylene cracking or coal tar). There are different types of  
240 carbon black and those with lower surface-area-to-volume ratios will have  
241 fewer PAHs absorb onto the surface during the manufacturing process. The  
242 PAHs are firmly bound to the carbon black surface under normal handling  
243 and use. They can only be extracted from the surface of the carbon black  
244 under rigorous laboratory conditions with strong solvents at elevated  
245 temperatures. There are 17 PAHs of concern that are listed in [Orally Inhaled  
246 and Nasal Drug Products \(1664.1\)](#).

247 5. ELASTOMERIC COMPONENTS—MANUFACTURING TECHNOLOGY AND STERILIZATION PROCEDURES

248 5.1 Generic Manufacturing

249 5.1.1 THERMOSET ELASTOMERS

250 The overall process of manufacturing thermoset elastomers consists of a  
251 rubber compound held in a heated mold under pressure. During this process,  
252 the elastomer is “cured” by undergoing a chemical reaction resulting in an  
253 elastic polymer network. The basic steps in manufacturing such components  
254 for pharmaceutical use are as follows:

- 255 • **Weighing of rubber ingredients:** Portions of the various  
256 ingredients are weighed according to instructions that reflect the  
257 rubber compound formulation.
- 258 • **Mixing:** The weighed portions of the various materials of  
259 construction are homogeneously mixed.
- 260 • **Preforming:** The mixed rubber is brought into a physical shape  
261 that allows easy handling in the subsequent step.
- 262 • **Molding:** Elasticity is introduced and the components are shaped  
263 by the curing reaction. The products are not individually shaped but  
264 are attached to a web.
- 265 • **Die-trimming:** The components are separated from the web by  
266 die-trimming.
- 267 • **Washing, lubrication, and drying:** The components are brought  
268 into their final state of microbiological and particulate cleanliness  
269 before packing. Lubrication, most often in the form of siliconization,  
270 typically is combined with washing and drying.
- 271 • **Packing:** The components are packed in suitable packaging  
272 material.

273 5.1.2 THERMOPLASTIC ELASTOMERS

274 TPEs are processed like plastic materials. They are injected, as a hot  
275 mixture, into a cooled mold. Unlike thermoset elastomers, a thermoplastic  
276 elastomer does not involve curing agents during molding. TPE components  
277 are not attached to a web, so they do not need die-trimming. Co-injection of  
278 a plastic material with a TPE is a way to create a two-material component,  
279 where the TPE part has a sealing and/or resealing function.

280 5.2 Sterilization Procedures

281 Elastomeric components for injections undergo sterilization as individual  
282 components prior to the filling process and then may be sterilized a second  
283 time as part of an assembled packaging system after filling. Sterilization of  
284 an individual component may be done using ethylene oxide, ionizing  
285 radiation, or steam; method selection is dependent on the elastomeric

286 formulation. For example, depending on the irradiation dose, some  
287 elastomeric formulations may not withstand ionizing radiation. Sterilization  
288 of the filled packaging system is usually done using steam. The desired  
289 outcome is a sterile component with no change to its critical parameters  
290 such as material chemical profile (extractables), functional performance, or  
291 drug product compatibility. For guidance on sterilization procedures, refer to  
292 the following suite of chapters: *Sterilization of Compendial*  
293 *Articles* (1229), *Steam Sterilization by Direct Contact* (1229.1), *Gaseous*  
294 *Sterilization* (1229.7), and *Radiation Sterilization* (1229.10).

295 **Ethylene oxide sterilization:** Ethylene oxide (EtO) can be used to  
296 sterilize elastomeric components, but the method has the drawback of  
297 requiring an outgassing time period to allow the levels of residual  
298 compounds to fall below the regulatory limits. Most EtO sterilization  
299 processes involve three different stages, which are preconditioning,  
300 sterilization, and degassing. The preconditioning stage includes a dwell time,  
301 under controlled temperature and humidity, which may decrease the  
302 resistance of microorganisms to inactivation and reduce the sterilization  
303 cycle time. Sterilization is completed by the introduction of EtO gas followed  
304 by an aeration phase that removes residual EtO. The aeration phase also  
305 allows time for removal of the common EtO degradants, ethylene  
306 chlorohydrin and ethylene glycol, from the elastomer. The degassing time  
307 required depends on factors such as the composition and size of the  
308 elastomeric part.

309 **Ionizing radiation sterilization:** Ionizing radiation can use either  
310 electron beam (e-beam) or gamma radiation. Cobalt 60 is a frequently used  
311 source of gamma rays, which are very penetrating. An e-beam produces  
312 accelerated electrons, which do not have the ability to penetrate materials to  
313 the same depth achieved by gamma rays. The energy provided by these two  
314 methods is sufficient to deliver a sterilizing dose but is also capable of  
315 exciting and dissociating polymer bonds. The free radicals produced within  
316 polymer structures initiate a series of complex chemical reactions (e.g.,  
317 chain scission or cross-linking) that may continue for a period of time after  
318 irradiation is completed. Inhibitors and stabilizers can be added to polymer  
319 formulations and are designed either to absorb energy or react with the free  
320 radicals. Even so, not all elastomeric formulations are deemed to be suitable  
321 for radiation sterilization.

322 **Steam sterilization:** Steam sterilization is typically performed in an  
323 autoclave under saturated steam conditions. Commonly used cycles are  
324 121°–122° for 30–60 min. The components are dried following sterilization,  
325 not only to remove surface water but also to remove residual moisture that  
326 has entered the matrix of the elastomer. Drying procedures need to be

327 optimized based on a number of factors including the length of the  
328 sterilization cycle, the elastomeric formulation, and the elastomer size and  
329 shape. Particular care should be taken when the elastomeric components are  
330 to be used to seal a lyophilized or powder-filled product, as residual moisture  
331 in the component can migrate into the formulation over time.

#### 332 6. IDENTIFICATION TESTS

333 Components are made of a wide variety of elastomeric materials and  
334 optional polymeric coatings. For this reason, it is beyond the scope  
335 of (381) to specify identification tests that encompass all possible component  
336 presentations. However, it is the responsibility of the component supplier  
337 and the drug product manufacturer to verify the component's elastomeric  
338 formulation and any coating or laminate material used according to suitable  
339 identification tests. Examples of some of the analytical test methodologies  
340 that may be used include specific gravity, percentage of ash analysis, sulfur  
341 content determination, Fourier-transform infrared spectroscopy–attenuated  
342 total reflectance (FTIR–ATR) test, chromatography of an extract, UV  
343 absorption spectrophotometry of an extract, or infrared absorption  
344 spectrophotometry of a pyrolysate.

#### 345 7. TEST REQUIREMENTS AND RESPONSIBILITIES

346 Elastomeric closures provided by a supplier to a drug product  
347 manufacturer may demonstrate compliance with specific (e.g., biological  
348 and/or physicochemical) specifications as required by the drug product  
349 manufacturer in accordance with requirements established by the  
350 manufacturer for storage and/or delivery of a unique drug product.  
351 Processing such as sterilization may be performed on finished components  
352 by the supplier prior to delivery to the drug product manufacturer, or by the  
353 drug product manufacturer before use, as agreed upon by both the supplier  
354 and the drug product manufacturer.

355 For elastomeric closures that are subsequently processed or sterilized by  
356 the drug product manufacturer after receipt from a supplier, the drug  
357 product manufacturer is responsible for ensuring that the closures are  
358 compliant with appropriate specifications required for storage and/or  
359 delivery of a unique drug product after the application of such processing  
360 and/or sterilization conditions (i.e., in their ready-to-use state). This is of  
361 particular importance if closures are processed under conditions that may  
362 impact the biological, physicochemical, or functionality characteristics of the  
363 closure.

#### 364 8. SUMMARY OF (381) PHYSICOCHEMICAL TESTS

##### 365 8.1 Test Samples

366 Elastomeric components are comprised of multiple materials that have  
367 unique attributes for each configuration, application, and change resulting  
368 from any treatment that occurs after formation (post-processing). This can  
369 be a challenge for establishing elastomeric baselines due to multiple  
370 combinations of materials and post-processing treatments by the component  
371 manufacturer and the drug product manufacturer.

372 Test samples described in (381) are to mimic the finished, processed  
373 component intended for the final product and its packaging/delivery system.  
374 It is the drug product manufacturer who is ultimately responsible for the  
375 quality and safety of the finished pharmaceutical product. Therefore, the  
376 drug product manufacturer is responsible for ensuring component  
377 conformance to the baseline requirements of (381).

## 378 8.2 Physicochemical Tests

379 **Appearance (turbidity/opalescence):** This is a nonspecific test for all  
380 the extractable species in a rubber formulation that are not soluble in an  
381 aqueous solution. A high turbidity is the indication of a high extractable  
382 potential. Species promoting turbidity have numerous origins in a rubber  
383 formulation, including fatty-acid derivatives, residues of curing systems, and  
384 oligomers from the elastomer.

385 **Color:** This is a nonspecific test indicative of the presence of extractable  
386 species in a rubber formulation that have the capacity of attributing color to  
387 an aqueous solution. Species that cause color may have several origins in a  
388 rubber formulation. Aqueous solutions are common in pharmaceutical  
389 packaging/delivery systems.

390 **Acidity or alkalinity:** This is a nonspecific test indicative of the acidic,  
391 basic, or buffering power of the aqueous extractables from the rubber  
392 formulation. High values in the acidity/alkalinity test may need to be  
393 evaluated in conjunction with the specifics of a drug solvent vehicle and  
394 anticipated specification of the drug product for pH.

395 **Absorbance:** This UV spectrum test of an aqueous extract from a rubber  
396 formulation is indicative of the unsaturated or aromatic character of the  
397 chemical species extracted. Unsaturated compounds in the extracts may  
398 originate from many raw materials and additives of a rubber formulation  
399 such as antioxidants and curing or dyeing agents.

400 **Reducing substances:** This is a nonspecific test. Extracted species from  
401 a rubber formulation with potential reducing power may originate from most  
402 raw materials of a rubber formulation (polymers, curing systems,  
403 preservatives, antioxidants, etc.).

404 **Volatile sulfides:** This test is specific for rubber formulations containing  
405 sulfur. Sulfur and sulfur precursors are often used as components of curing  
406 systems for rubber.

407 **Ammonium:** This test is specific for rubber formulations with nitrogen-  
408 containing raw materials. Ammonium ions can be generated during the  
409 curing process. Thiurams and thiazoles are examples of nitrogen-containing  
410 curing systems used.

411

### 8.3 Biocompatibility Tests

412 In vitro biocompatibility testing is performed as described in general test  
413 chapter *Biological Reactivity Tests, In Vitro* (87). Materials that do not meet  
414 the requirements of the in vitro test may require performance of appropriate  
415 in vivo tests according to the procedures set forth in the general test  
416 chapter *Biological Reactivity Tests, In Vivo* (88).

417

### 8.4 Extractable Elements Method

418 Knowledge of the potential extractable elements present in an elastomeric  
419 component is important in establishing the component's suitability.  
420 Knowledge of the elements that are likely to be present and the  
421 concentrations at which they may be observed provides information to  
422 determine potential product quality risk. Elastomeric components can vary  
423 widely in terms of their intentionally and unintentionally added elements and  
424 their potential use. Because of this, it is challenging to provide universally  
425 effective and efficient tests methodologies, lists of target elements, and  
426 reporting requirements. It is the material user's responsibility to evaluate  
427 the need for extractable elements testing and, if such testing is necessary,  
428 to establish and justify the means by which testing is accomplished, taking  
429 into account target elements, extraction conditions, extract analysis, and  
430 reporting requirements.

431 **Target elements:** Relevant elements that should be tested for include  
432 those that have been intentionally added to the elastomer (e.g. zinc) and  
433 those of toxicological concern. Some of the relevant elements for elastomeric  
434 components for injections include antimony, arsenic, cadmium, cobalt,  
435 copper, lead, lithium, mercury, nickel, vanadium, and zinc.

436 Elastomeric components used in injectable packaging systems do not  
437 dissolve under the conditions of use. Rather, substances derived from the  
438 component accumulate in the packaged articles by the process of leaching  
439 (extraction). Thus, the appropriate and relevant sample-preparation process  
440 for assessing elements in an elastomeric component is extraction, as  
441 opposed to complete digestion. A screen for relevant extractable elements  
442 can be achieved based on a rigorous extraction method to determine the  
443 potential for elements to extract. The results of the screening study can then

444 provide the basis for targeting elements for simulated or intended use  
445 studies. Screening is an important first step to ensure proper elements and  
446 sensitivities are realized. Trace elemental analysis can be challenging and  
447 the extraction method is critical. The following method has been verified  
448 based on recovery of critical elements related to elastomer components.

449 **Extraction solution:** Prepare a solution of a mixture of acids with gold  
450 (Au) to stabilize mercury (Hg) in the following ratio: 0.2 N nitric acid (HNO<sub>3</sub>),  
451 0.05 N hydrochloric acid (HCl), and 200 ppb gold (Au). Prepare the solution  
452 in a volume sufficient to prepare all standards, blanks, spikes, and  
453 extractions. Care should be taken to use high-purity reagents.

454 **Analysis:** Place whole, uncut components equivalent to 1 g/2.5 mL of  
455 the *Extraction solution* into a suitable plastic container and record the  
456 weight. Prepare two extraction blank solutions (one for spiking) using a  
457 container of the same type as that used for the samples, omitting the  
458 closures. Seal the containers and place in an oven at 70°. Remove  
459 containers after 24 h and allow to cool. Analyze within 48 h. Extracts, spikes,  
460 and blanks are to be analyzed by inductively coupled plasma–mass  
461 spectrometry (ICP–MS) and/or inductively coupled plasma–optical emission  
462 spectroscopy (ICP–OES). Refer to [Elemental Impurities—  
463 Procedures \(233\)](#) for analytical procedures and system suitability.

464 **Extraction recovery:** Prepare a 10 µg/mL solution of antimony (Sb),  
465 arsenic (As), cadmium (Cd), cobalt (Co), copper (Cu), lead (Pb), lithium (Li),  
466 mercury (Hg), nickel (Ni), vanadium (V), and zinc (Zn) in *Extraction  
467 solution* [0.2 N nitric acid (HNO<sub>3</sub>), 0.05 N hydrochloric acid (HCl), and 200  
468 ppb gold (Au)]. Using a suitable pipet, spike one of the blank extraction  
469 solutions with the appropriate volume of the 10-µg/mL solution to obtain a  
470 concentration of 0.05 µg/g.

471 **Analysis and reporting threshold:** Instrumentation and methods are  
472 those specified in [Elemental Impurities—Procedures \(233\)](#) and include an  
473 inductively coupled plasma–atomic emission spectrometer and an inductively  
474 coupled plasma–mass spectrometer (see [Plasma Spectrochemistry \(730\)](#)), as  
475 directed. The reporting threshold is 0.05 µg/g converted to µg/component.  
476 Calculate and report results based on the original sample size. [NOTE—  
477 Appropriate measures must be taken to correct for matrix-induced  
478 interferences (e.g., argon chloride interference with arsenic  
479 determinations).]

480 **Calculations:**

481 
$$\text{percent recovery} = \frac{(d \times 100)}{S} \text{ percent recovery} = \frac{(d \times 100)}{S}$$
  
482

*d* = amount of element detected (µg)

*s* = amount of element spiked ( $\mu\text{g}$ )

483  $\mu\text{g/g component} = (a \times t) / w$   $\mu\text{g/g component} = (a \times t) / w$

*a* = concentration of extract ( $\mu\text{g/mL}$ )

*t* = total extract, corrected for average blank (mL)

*w* = sample weight (g)

484  $\mu\text{g/component} = c \times s$   $\mu\text{g/component} = c \times s$

*c* = element concentration in extract ( $\mu\text{g/g}$ )

*s* = total weight of component (g)

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