



# Standard Specification for Nitrile Examination Gloves for Medical Application<sup>1</sup>

This standard is issued under the fixed designation D6319; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon ( $\epsilon$ ) indicates an editorial change since the last revision or reapproval.

## 1. Scope

1.1 This specification covers certain requirements for nitrile rubber gloves used in conducting medical examinations and diagnostic and therapeutic procedures.

1.2 This specification covers nitrile rubber examination gloves that fit either hand, paired gloves, and gloves by size. It also provides for packaged sterile or nonsterile or bulk non-sterile nitrile rubber examination gloves.

1.3 This specification is similar to that of Specification **D3578** for rubber examination gloves.

1.4 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

## 2. Referenced Documents

### 2.1 ASTM Standards:<sup>2</sup>

**D412** Test Methods for Vulcanized Rubber and Thermoplastic Elastomers—Tension

**D573** Test Method for Rubber—Deterioration in an Air Oven

**D3578** Specification for Rubber Examination Gloves

**D3767** Practice for Rubber—Measurement of Dimensions

**D5151** Test Method for Detection of Holes in Medical Gloves

**D6124** Test Method for Residual Powder on Medical Gloves

### 2.2 ISO Standard:

**ISO 2859** Sampling Procedures and Tables for Inspection by Attributes<sup>3</sup>

### 2.3 Other Documents:

<sup>1</sup> This specification is under the jurisdiction of ASTM Committee **D11** on Rubber and is the direct responsibility of Subcommittee **D11.40** on Consumer Rubber Products.

Current edition approved May 1, 2010. Published June 2010. Originally approved in 1999. Last previous edition approved in 2005 as D6319 – 00a (2005)<sup>e1</sup>. DOI: 10.1520/D6319-10.

<sup>2</sup> For referenced ASTM standards, visit the ASTM website, [www.astm.org](http://www.astm.org), or contact ASTM Customer Service at [service@astm.org](mailto:service@astm.org). For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

<sup>3</sup> Available from American National Standards Institute, 25 W. 43rd St., 4th Floor, New York, NY 10036.

## U.S. Pharmacopeia<sup>4</sup>

## 3. Significance and Use

3.1 The specification is intended as a referee procedure for evaluating the performance and safety of nitrile rubber examination gloves. The safe and proper use of nitrile rubber examination gloves is beyond the scope of this specification.

## 4. Material

4.1 Any nitrile rubber polymer compound may be used that permits the glove to meet the requirements of this specification.

4.2 A lubricant that meets the current requirements of the **U.S. Pharmacopeia** for absorbable dusting powder may be applied to the glove. Other lubricants may be used if their safety and efficacy have been previously established.

4.3 The inside and outside surface of the nitrile rubber examination gloves shall be free of talc.

## 5. Sampling

5.1 For referee purposes, gloves shall be sampled from finished product, after sterilization when labeled sterile, and inspected in accordance with **ISO 2859**. The inspection levels and acceptable quality levels (AQL) shall conform to those specified in **Table 1**, or as agreed upon between the purchaser and the seller, if the latter is more comprehensive.

## 6. Performance Requirements

6.1 Gloves, sampled in accordance with Section **5**, shall meet the following referee performance requirements:

6.1.1 Product comply with requirements for sterility when tested in accordance with **7.2** when labeled sterile.

6.1.2 Shall comply with freedom from holes when tested in accordance with **7.3**.

6.1.3 Have consistent physical dimensions in accordance with **7.4**.

6.1.4 Have acceptable physical property characteristics in accordance with **7.5**.

6.1.5 Have a powder residue limit of 2.0 mg in accordance with **7.6**.

6.1.6 Have a recommended maximum powder limit of 10 mg/dm<sup>2</sup> in accordance with **7.7**.

<sup>4</sup> U. S. Pharmacopeia, latest edition, Mack Publishing Co., Easton, PA 19175.

**TABLE 1 Performance Requirements**

Characteristic	Related Defects	Inspection Level	AQL
Sterility	fails sterility	A	N/A
Freedom from holes	holes	G-1	2.5
Dimensions	width, length, and thickness	S-2	4.0
Physical properties	before aging, after accelerated aging	S-2	4.0
Powder-free Residue	exceeds maximum limit	N=5	N/A
Powder Amount	exceeds recommended maximum limit	N=2	N/A

<sup>A</sup>See U.S. Pharmacopeia.

## 7. Referee Test Methods

7.1 The following tests shall be conducted to ensure the requirements of Section 6, as prescribed in Table 1:

7.2 *Sterility Test*—Testing for sterility shall be conducted in accordance with the latest edition of the U.S. Pharmacopeia.

7.3 *Freedom from Holes*—Testing for freedom from holes shall be conducted in accordance with Test Method D5151.

7.4 *Physical Dimensions Test*:

7.4.1 The gloves shall comply with the dimension requirements prescribed in Table 2.

7.4.2 The length shall be expressed in millimetres as measured from the tip of the middle finger to the outside edge of the cuff.

7.4.3 The width of the palm shall be expressed in millimetres as measured at a level between the base of the index finger and the base of the thumb. Values of width per size other than listed shall meet the stated tolerance specified in Table 2.

7.4.4 The minimum thickness shall be expressed in millimetres as specified in Table 2 when using a dial or digital micrometer that meets requirements described in Test Methods D412 and Practice D3767, and in the locations indicated in Fig. 1. For referee tests, cutting the glove is necessary to obtain single-thickness measurements. (See Practice D3767 for more information.)

7.5 *Physical Requirements Test*:

7.5.1 Before and after accelerated aging, the gloves shall conform to the physical requirements specified in Table 3. Tests shall be conducted in accordance with Test Methods D412. Die C is recommended.

7.5.2 *Accelerated Aging*—The gloves shall be aged in accordance with Test Method D573. Test the gloves in accordance with either one of the following methods:

7.5.2.1 After being subjected to a temperature of  $70 \pm 2^\circ\text{C}$  for  $166 \pm 2$  h, the tensile strength and ultimate elongation shall not be less than the values specified in Table 3. This method shall be the conditions for referee tests.

7.5.2.2 After being subjected to a temperature of  $100 \pm 2^\circ\text{C}$  for  $22 \pm 0.3$  h, the tensile strength and ultimate elongation shall not be less than the values specified in Table 3.

7.6 *Powder Free Gloves*—Determine the powder residue using Test Method D6124.

7.7 *Powdered Gloves*:

7.7.1 Determine the recommended maximum powder limit using Test Method D6124 for powdered gloves.

7.7.2 Determine the square decimetres for the glove size as in the paragraph on determining the square decimetres of glove size in Specification D3578.

## 8. Acceptance

8.1 Gloves will be considered to meet the referee performance requirements when test results conform to the requirements prescribed in Table 1.

8.2 Retests or reinspections are permissible under the provision of the U.S. Pharmacopeia and ISO 2859.

## 9. Packaging and Package Marking

9.1 *Sterile Packaging*:

9.1.1 The unit of packaging shall normally be one glove or one pair of gloves.

9.1.2 A glove or pair of gloves, normally, shall be enclosed in an inner wallet or wrapper. The wrapper shall be of sufficient size when opened to provide a field for glove-donning purposes.

9.1.3 The glove or pair of gloves, and accompanying wrapper if utilized, shall be totally enclosed in an outer package that will allow sterilization of the product.

9.1.4 The outer package shall have a method of closure sufficient to ensure the sterility of the product until opened or damaged.

9.1.5 The outer package shall have sufficient strength and integrity to withstand normal transportation and storage within the intermediate or shipping cartons, or both.

9.1.6 The method of closure of the outer package shall be such that prior opening will be detectable by the user.

**TABLE 2 Dimensions and Tolerances**

NOTE—Sizing that falls within the tolerance overlaps between two sizes may be labeled as a size range including both sizes, for example, small/medium and medium/large.

Designation	Size							Tolerance, mm
	6	6 ½	7	7 ½	8	8 ½	9	
Width by size	75	83	89	95	102	108	114	±6
Width by		x-small 70	small 80	Unisize 85	medium 95	large 110	X-large 120	±10
Length		220	220	230	230	230	230	min
Thickness, mm:								
finger				0.05				min
palm				0.05				min

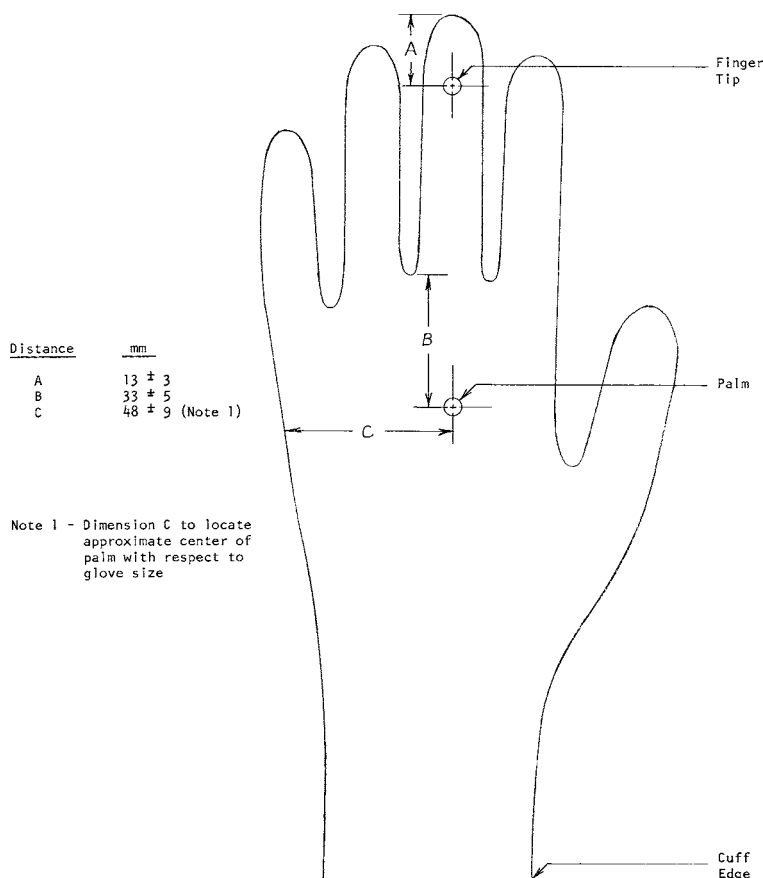


FIG. 1 Location of Thickness Measurements

TABLE 3 Physical Requirements

Before Aging		After Accelerated Aging	
Tensile Strength	Ultimate Elongation	Tensile Strength	Ultimate Elongation
14 MPa, min	500 % min	14 MPa min	400 % min

9.1.7 None of the packaging material shall contain any material likely to impair the quality and use of the gloves.

9.1.8 Intermediate cartons and shipping cases shall be of sufficient strength to maintain the quality and sterility of the product during normal transportation and storage.

#### 9.2 Nonsterile and Bulk Packaging:

9.2.1 The gloves shall be enclosed in an outer package that has sufficient strength to withstand normal transportation and storage within the cartons or shipping cases, or both.

9.2.2 None of the packaging material shall contain any material likely to impair the quality and use of the gloves.

9.2.3 Cartons and shipping cases shall be of sufficient strength to maintain the quality of the product during normal transportation and storage.

#### 9.3 Package Marking:

9.3.1 Sterile packages shall bear markings for the contents to include the glove size, instructions for opening, the legend “sterile,” and a manufacturing lot number.

9.3.2 Nonsterile and bulk packages shall bear markings for the contents to include the glove size and a manufacturing lot number.

9.3.3 The outermost case shall be labeled with the glove size and a manufacturing lot number. Sterile product cases shall also be marked with the legend “sterile.”

9.3.4 All levels of packaging shall conform to all appropriate government labeling regulations.

### 10. Keywords

10.1 examination gloves; nitrile; rubber

*ASTM International takes no position respecting the validity of any patent rights asserted in connection with any item mentioned in this standard. Users of this standard are expressly advised that determination of the validity of any such patent rights, and the risk of infringement of such rights, are entirely their own responsibility.*

*This standard is subject to revision at any time by the responsible technical committee and must be reviewed every five years and if not revised, either reapproved or withdrawn. Your comments are invited either for revision of this standard or for additional standards and should be addressed to ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the responsible technical committee, which you may attend. If you feel that your comments have not received a fair hearing you should make your views known to the ASTM Committee on Standards, at the address shown below.*

*This standard is copyrighted by ASTM International, 100 Barr Harbor Drive, PO Box C700, West Conshohocken, PA 19428-2959, United States. Individual reprints (single or multiple copies) of this standard may be obtained by contacting ASTM at the above address or at 610-832-9585 (phone), 610-832-9555 (fax), or [service@astm.org](mailto:service@astm.org) (e-mail); or through the ASTM website ([www.astm.org](http://www.astm.org)). Permission rights to photocopy the standard may also be secured from the ASTM website ([www.astm.org/COPYRIGHT/](http://www.astm.org/COPYRIGHT/)).*