



Designation: F2026 – 14

# Standard Specification for Polyetheretherketone (PEEK) Polymers for Surgical Implant Applications<sup>1</sup>

This standard is issued under the fixed designation F2026; 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 polyetheretherketone (PEEK) polymer in virgin forms as supplied by a vendor (pellets, powder, fabricated forms, and so forth). It provides requirements and associated test methods for these thermoplastics when they are to be used in the manufacture of intracorporeal devices such as surgical implants or components of surgical or dental devices.

1.2 The properties included in this specification are those applicable for PEEK polymers only. Indicated properties are for fabricated forms. Materials or forms containing colorants, fillers, processing aids, or other additives, as well as polymer blends which contain PEEK, or reclaimed materials, are not covered by this specification.

1.3 This specification is designed to recommend physical, chemical, and biological test methods to establish a reasonable level of confidence concerning the performance of virgin PEEK polymers for use in medical implant devices.

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

1.5 When evaluating material in accordance with this specification, hazardous materials, operations, and equipment may be involved. *This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.*

## 2. Referenced Documents

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

<sup>1</sup> This specification is under the jurisdiction of ASTM Committee F04 on Medical and Surgical Materials and Devices and is the direct responsibility of Subcommittee F04.11 on Polymeric Materials.

Current edition approved March 1, 2014. Published May 2014. Originally approved in 2000. Last previous edition approved in 2012 as F2026 – 12. DOI: 10.1520/F2026-14.

<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.

D256 Test Methods for Determining the Izod Pendulum Impact Resistance of Plastics

D638 Test Method for Tensile Properties of Plastics

D648 Test Method for Deflection Temperature of Plastics Under Flexural Load in the Edgewise Position

D695 Test Method for Compressive Properties of Rigid Plastics

D790 Test Methods for Flexural Properties of Unreinforced and Reinforced Plastics and Electrical Insulating Materials

D792 Test Methods for Density and Specific Gravity (Relative Density) of Plastics by Displacement

D1505 Test Method for Density of Plastics by the Density-Gradient Technique

D3418 Test Method for Transition Temperatures and Enthalpies of Fusion and Crystallization of Polymers by Differential Scanning Calorimetry

D4000 Classification System for Specifying Plastic Materials

F748 Practice for Selecting Generic Biological Test Methods for Materials and Devices

### 2.2 ISO Standards:<sup>3</sup>

ISO 178 Plastics—Determination of Flexural Properties

ISO 180 Plastics—Determination of Izod Impact Strength

ISO 527 Plastics—Determination of Tensile Properties—Part 1: General Principles

ISO 1183 Plastics—Methods for Determining the Density of Non-cellular Plastics—Part 2: Density Gradient Column Method

ISO 10993 Biological Evaluation of Medical Devices, Parts 1-12

ISO 13485 Medical Devices—Quality Management Systems—Requirements for Regulatory Purposes

### 2.3 Other Documents:

United States Pharmacopeia, Vol. XXI, or latest edition<sup>4</sup>

## 3. Terminology

### 3.1 Definitions of Terms Specific to This Standard:

<sup>3</sup> Available from American National Standards Institute (ANSI), 25 W. 43rd St., 4th Floor, New York, NY 10036, <http://www.ansi.org>.

<sup>4</sup> Available from U.S. Pharmacopeia (USP), 12601 Twinbrook Pkwy., Rockville, MD 20852-1790, <http://www.usp.org>.

3.1.1 *fabricated forms, n*—those items into which the virgin forms may be converted. These include shapes and forms produced by means of machining, extruding, and compression molding virgin forms into a subsequent entity (for example, fibers, tubes, rods, slabs, sheets, film, or complex shaped parts and devices).

3.1.2 *formulated compound, n*—materials, parts, or devices fabricated from virgin forms in such a way as to contain intentional or unintentional adjuvant substances.

3.1.3 *virgin forms, n*—the initially delivered form of the polymer as synthesized from its monomers prior to any processing or fabrication into a medical device. The provided resin is typically in the form of pellets, granules, or powder and is the material from which fibers, tubes, rods, slabs, sheets, films, or specific parts and devices are fabricated.

#### 4. Classification

4.1 The PEEK polymer in the scope of this specification is a pure semicrystalline homopolymer consisting of phenylene rings connected by ether (E) and carbonyl (or ketone, K) groups along the polymer chain (see [Appendix X1](#)). Its polymeric structure is defined by the repeating unit EEK.

4.2 Types of PEEK plastics, molding, and extrusion grades are described in Classification System [D4000](#).

#### 5. Properties

5.1 The properties listed below shall be considered in selecting material(s) in accordance with the specific end-use requirements.

5.2 The infrared spectrum<sup>5</sup> of these materials is characteristic of their molecular repeating units. A representative spectrum is listed in [Appendix X3](#). The PEEK polymer shall yield an infrared spectrum, which exhibits major bands only at the wavelengths listed for a standard reference spectrum of that material.

5.2.1 The infrared spectrum, as used in this specification, is to identify the specific type of poly aryl ether ketone (PAEK) present and does not necessarily indicate an acceptable degree of material purity.

5.2.2 The presence of additional bands in the sample's infrared spectrum compared to that of the reference material may indicate a different PAEK or impurities, or both.

5.3 The physical and chemical property requirements for the virgin polymer are listed in [Table 1](#). If additional characteristics are necessary because of a specific application, the procedures referenced in [Section 2](#) are recommended, or as agreed upon between the vendor and the purchaser.

5.4 The viscosity requirements will vary depending upon the grade and test method. The method and requirements shall be agreed upon between the vendor and the purchaser.

5.5 The chemical, physical, and mechanical properties of fabricated forms are related to the processes utilized in producing the fabricated form (for example, molding, machining, sterilization, and so forth). Additionally, the properties necessary for a particular device to perform properly will vary from one device type to another. [Table 2](#) lists some typical properties of non-sterilized fabricated forms.

5.6 Test specimens shall be fabricated (machined, injection molded, and so forth) from the virgin polymer, or finished part, in such a way as to effectively represent the material characteristics of the non-sterilized finished part.

5.6.1 As with any material, some characteristics may be altered by the processing techniques (for example, molding, extrusion, machining, assembly, and sterilization) required for the production of a specific part or device. Therefore, properties of fabricated forms of these polymers should be evaluated using test methods which are appropriate to ensure safety and efficacy as agreed upon by the vendor, purchaser, and regulating bodies.

5.7 Tests and test procedures shall be such as to ensure a high level of control and characterization of the virgin polymer as received from the supplier. The test methods referenced in [Section 2](#) may be appropriate (Test Methods [D648](#) and [D695](#)).

<sup>5</sup> Silverstein, R. M., Bassler, G. C., and Morrill, T. C., *Spectroscopic Identification of Organic Compounds*, 5th ed., John Wiley & Sons, New York, NY.

**TABLE 1 Required Properties of Virgin Resin**

Parameter	Method	Requirement
Glass transition temperature, $T_g$ (°C)	DSC, <sup>4</sup> 20°K/min, sealed sample, $T_g$ taken on second reheat, <a href="#">D3418</a>	125 - 165
Melt temperature, $T_m$ (°C)	DSC, 20°K/min, sealed sample, $T_m$ taken as max point on reheat endotherm, <a href="#">D3418</a>	320 - 360
Recrystallization temperature, $T_c$ (°C)	DSC, 20°K/min, sealed sample, $T_c$ taken as max point on cooling exotherm, <a href="#">D3418</a>	260 - 320
Viscosity	As agreed per <a href="#">5.4</a>	As agreed per <a href="#">5.4</a>
Infrared spectrum	As agreed per <a href="#">5.2</a>	As agreed per <a href="#">5.2</a>
Total heavy metals as lead, max, %	US Pharmacopeia, Test 231	<0.1

<sup>4</sup> Differential Scanning Calorimetry (DSC).