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Standard Guide for Design and Evaluation of Primary Flexible Packaging for Medical Products¹

This standard is issued under the fixed designation F2097; 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 (ϵ) indicates an editorial change since the last revision or reapproval.

1. Scope

- 1.1 This guide provides directions for the design and evaluation of primary flexible packages for medical products. The package materials must be selected appropriately for manufacturing process, end use, and the product being packaged.
- 1.2 This guide provides a compendium of test methods, practices, and procedures. Specific individual test methods must be selected based on the pertinent characteristics of the specific product to be packaged and the purpose for testing, research and development, or compliance. Not all test methods will be applicable.
- 1.3 This guide does not address acceptability criteria, which need to be determined jointly by the package producer and the medical products manufacturer.
- 1.4 This guide does not assess the product to be packaged or the sterilization method to be used.
- 1.5 The units cited in the referenced standard should be used.
- 1.6 This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.

2. Referenced Documents

2.1 ASTM Standards:²

D374 Test Methods for Thickness of Solid Electrical Insulation (Metric) D0374_D0374M

D589 Test Method for Opacity of Paper (15° Diffuse Illuminant A, 89 % Reflectance Backing and Paper Backing) (Withdrawn 2010)³

D638 Test Method for Tensile Properties of Plastics

D645/D645M Test Method for Thickness of Paper and Paperboard (Withdrawn 2010)³

D685 Practice for Conditioning Paper and Paper Products for Testing

D726 Test Method for Resistance of Nonporous Paper to Passage of Air (Withdrawn 2009)³

D882 Test Method for Tensile Properties of Thin Plastic Sheeting

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² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

³ The last approved version of this historical standard is referenced on www.astm.org.



- D1003 Test Method for Haze and Luminous Transmittance of Transparent Plastics
- D1251 Test Method for Water Vapor Permeability of Packages by Cycle Method (Withdrawn 1999)³
- D1434 Test Method for Determining Gas Permeability Characteristics of Plastic Film and Sheeting
- D1709 Test Methods for Impact Resistance of Plastic Film by the Free-Falling Dart Method
- D1777 Test Method for Thickness of Textile Materials
- D1894 Test Method for Static and Kinetic Coefficients of Friction of Plastic Film and Sheeting
- D1922 Test Method for Propagation Tear Resistance of Plastic Film and Thin Sheeting by Pendulum Method
- D1938 Test Method for Tear-Propagation Resistance (Trouser Tear) of Plastic Film and Thin Sheeting by a Single-Tear Method
- D2019 Test Method for Dirt in Paper and Paperboard (Withdrawn 2010)³
- D2457 Test Method for Specular Gloss of Plastic Films and Solid Plastics
- D3078 Test Method for Determination of Leaks in Flexible Packaging by Bubble Emission
- D3079 Test Method for Water Vapor Transmission of Flexible Heat-Sealed Packages for Dry Products (Withdrawn 2018)³
- D3335 Test Method for Low Concentrations of Lead, Cadmium, and Cobalt in Paint by Atomic Absorption Spectroscopy
- D3420 Test Method for Pendulum Impact Resistance of Plastic Film
- D3718 Test Method for Low Concentrations of Chromium in Paint by Atomic Absorption Spectroscopy
- D3776 Test Methods for Mass Per Unit Area (Weight) of Fabric
- D3985 Test Method for Oxygen Gas Transmission Rate Through Plastic Film and Sheeting Using a Coulometric Sensor
- D4169 Practice for Performance Testing of Shipping Containers and Systems
- D4279 Test Methods for Water Vapor Transmission of Shipping Containers—Constant and Cycle Methods
- D4321 Test Method for Package Yield of Plastic Film
- D4332 Practice for Conditioning Containers, Packages, or Packaging Components for Testing
- D4754 Test Method for Two-Sided Liquid Extraction of Plastic Materials Using FDA Migration Cell
- D5264 Practice for Abrasion Resistance of Printed Materials by the Sutherland Rub Tester
- D7386 Practice for Performance Testing of Packages for Single Parcel Delivery Systems
- E167 Practice for Goniophotometry of Objects and Materials (Withdrawn 2005)³
- E171/E171M Practice for Conditioning and Testing Flexible Barrier Packaging
- E398 Test Method for Water Vapor Transmission Rate of Sheet Materials Using Dynamic Relative Humidity Measurement
- F17 Terminology Relating to Primary Barrier Packaging
- F88 Test Method for Seal Strength of Flexible Barrier Materials
- F99 Guide for Writing a Specification for Flexible Barrier Rollstock Materials
- F151 Test Method for Residual Solvents in Flexible Barrier Materials (Withdrawn 2004)³
- F372 Test Method for Water Vapor Transmission Rate of Flexible Barrier Materials Using an Infrared Detection Technique (Withdrawn 2009)³
- F392 Practice for Conditioning Flexible Barrier Materials for Flex Durability
- F748 Practice for Selecting Generic Biological Test Methods for Materials and Devices 2449baa772/astm-12097-20
- F813 Practice for Direct Contact Cell Culture Evaluation of Materials for Medical Devices
- F895 Test Method for Agar Diffusion Cell Culture Screening for Cytotoxicity
- F904 Test Method for Comparison of Bond Strength or Ply Adhesion of Similar Laminates Made from Flexible Materials
- F1140 Test Methods for Internal Pressurization Failure Resistance of Unrestrained Packages
- F1249 Test Method for Water Vapor Transmission Rate Through Plastic Film and Sheeting Using a Modulated Infrared Sensor
- F1306 Test Method for Slow Rate Penetration Resistance of Flexible Barrier Films and Laminates
- F1307 Test Method for Oxygen Transmission Rate Through Dry Packages Using a Coulometric Sensor
- F1443 Practice for Using 0.008-in. (0.203-mm) Aperture Reflectometers as Test Instruments for Measuring Visual Image Quality of Business Copy Images (Withdrawn 2017)³
- F1608 Test Method for Microbial Ranking of Porous Packaging Materials (Exposure Chamber Method)
- F1884 Test Methods for Determining Residual Solvents in Packaging Materials
- F1886 Test Method for Determining Integrity of Seals for Flexible Packaging by Visual Inspection
- F1921 Test Methods for Hot Seal Strength (Hot Tack) of Thermoplastic Polymers and Blends Comprising the Sealing Surfaces of Flexible Webs
- F1927 Test Method for Determination of Oxygen Gas Transmission Rate, Permeability and Permeance at Controlled Relative Humidity Through Barrier Materials Using a Coulometric Detector
- F1929 Test Method for Detecting Seal Leaks in Porous Medical Packaging by Dye Penetration
- F1980 Guide for Accelerated Aging of Sterile Barrier Systems for Medical Devices
- F2029 Practices for Making Laboratory Heat Seals for Determination of Heat Sealability of Flexible Barrier Materials as Measured by Seal Strength
- F2054 Test Method for Burst Testing of Flexible Package Seals Using Internal Air Pressurization Within Restraining Plates
- F2095 Test Methods for Pressure Decay Leak Test for Flexible Packages With and Without Restraining Plates
- F2096 Test Method for Detecting Gross Leaks in Packaging by Internal Pressurization (Bubble Test)



F2203 Test Method for Linear Measurement Using Precision Steel Rule

F2217 Practice for Coating/Adhesive Weight Determination

F2227 Test Method for Non-Destructive Detection of Leaks in Non-sealed and Empty Packaging Trays by CO₂ Tracer Gas Method

F2228 Test Method for Non-Destructive Detection of Leaks in Packaging Which Incorporates Porous Barrier Material by CO₂
Tracer Gas Method

F2250 Practice for Evaluation of Chemical Resistance of Printed Inks and Coatings on Flexible Packaging Materials

F2251 Test Method for Thickness Measurement of Flexible Packaging Material

F2252 Practice for Evaluating Ink or Coating Adhesion to Flexible Packaging Materials Using Tape

F2227 Test Method for Non-Destructive Detection of Leaks in Non-sealed and Empty Packaging Trays by CO₂ Tracer Gas Method

F2228 Test Method for Non-Destructive Detection of Leaks in Packaging Which Incorporates Porous Barrier Material by CO₂
Tracer Gas Method

F2338 Test Method for Nondestructive Detection of Leaks in Packages by Vacuum Decay Method

F2391 Test Method for Measuring Package and Seal Integrity Using Helium as the Tracer Gas

F2475 Guide for Biocompatibility Evaluation of Medical Device Packaging Materials

F2476 Test Method for the Determination of Carbon Dioxide Gas Transmission Rate (CO₂TR) Through Barrier Materials Using an Infrared Detector

F2559 Guide for Writing a Specification for Sterilizable Peel Pouches

F2622 Test Method for Oxygen Gas Transmission Rate Through Plastic Film and Sheeting Using Various Sensors

F2638 Test Method for Using Aerosol Filtration for Measuring the Performance of Porous Packaging Materials as a Surrogate Microbial Barrier

F2714 Test Method for Oxygen Headspace Analysis of Packages Using Fluorescent Decay

F2824 Test Method for Mechanical Seal Strength Testing for Round Cups and Bowl Containers with Flexible Peelable Lids

F2825 Practice for Climatic Stressing of Packaging Systems for Single Parcel Delivery

F2981 Test Method for Verifying Nonporous Flexible Barrier Material Resistance to the Passage of Air

F3004 Test Method for Evaluation of Seal Quality and Integrity Using Airborne Ultrasound

F3039 Test Method for Detecting Leaks in Nonporous Packaging or Flexible Barrier Materials by Dye Penetration

F3136 Test Method for Oxygen Gas Transmission Rate through Plastic Film and Sheeting using a Dynamic Accumulation Method

F3169 Test Method for Leak Detection in Blister Packaging by Vacuum Deflection Method by Laser Measurement

F3287 Test Method for Nondestructive Detection of Leaks in Packages by Mass Extraction Method

F3299 Test Method for Water Vapor Transmission Rate Through Plastic Film and Sheeting Using an Electrolytic Detection Sensor (Coulometric P₂O₅ Sensor)

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F3300 Test Method for Abrasion Resistance of Flexible Packaging Films Using a Reciprocating Weighted Stylus

2.2 EN/ISO Standards:⁴

EN 868/1ISO 187 Annex C Gurley, Schopper, Dye Penetration Paper, Board and Pulps—Standard Atmosphere for Conditioning and Testing and Procedure for Monitoring the Atmosphere and Conditioning of Samples

ISO 2556 Plastics—Determination of Gas Transmission Rate of Films and Thin Sheets Under Atmospheric Pressure— Manometric Method

ISO 5636-5 Paper and Board—Determination of Air Permeance (Medium Range)—Part 5: Gurley Method

ISO 10993 Biological Evaluation of Medical Devices

ISO 11607-1 Packaging for Terminally Sterilized Medical Devices, Annex C

ISO 15105-1 Plastics—Film and Sheeting—Determination of Gas Transmission Rate—Part 1: Differential-Pressure Method

ISO 15105–2 Plastics—Film and Sheeting—Determination of Gas Transmission Rate—Part 2: Equal-Pressure Method

2.3 Military Specification:⁵

Mil Spec 36954C Bacterial Filtration Efficiency

2.4 TAPPI Standards:⁶

TAPPI T 402 Standard Conditioning and Testing Atmospheres for Paper, Board, Pulp Handsheets, and Related Products

TAPPI T 404T 404 Tensile Breaking Strength and Elongation of Paper and Paperboard

TAPPI T 425 Opacity of Paper (15/D Geometry, Illuminant A/2 Degrees, 89% Reflectance Backing and Paper Backing)

TAPPI T 437T 437 Dirt in Paper and Paperboard

TAPPI T 460T 460 Air Resistance of Paper (Gurley Method)

TAPPI T 494T 494 Tensile Breaking Properties of Paper and Paperboard (Using Constant Rate of Elongation Apparatus)

⁴ Available from International Organization for Standardization (ISO), 1 rue de Varembé, Case postale 56, CH-1211, Geneva 20, Switzerland, http://www.iso.ch.

⁵ Available from Standardization Documents Order Desk, Bldg. 4 Section D, 700 Robbins Ave., Philadelphia, PA 19111-5094, Attn: NPOPS.

⁶ Available from Technical Association of the Pulp and Paper Industry (TAPPI), 15 Technology Parkway South, Norcross, GA 30092, http://www.tappi.org.



TAPPI T 519 Diffuse Opacity of Paper (D/O Paper Backing)

TAPPI T 536T 536 Resistance of Paper to Passage of Air (High Pressure Gurley Method)

TAPPI T 547 T 547 Air Permeance of Paper and Paperboard (Sheffield Method)

2.5 ISTA Procedures:⁷

ISTA 3A Packaged Products for Parcel Delivery System Shipments 70 kg (150 lb) or Less (standard, small, flat, or elongated)

ISTA 3E Unitized Loads of Same Product

ISTA 4AB Packaged-Products for Shipment in Known Distribution Channels

ISTA 6-FEDEX-A FedEx Procedures for Testing Packaged Products Weighing Up to 150 lbs

3. Terminology

- 3.1 Definitions of Terms Specific to This Standard:
- 3.1.1 *barrier requirements, n*—the need to promote or inhibit moisture, gas, or light, or a combination thereof, while maintaining necessary levels of sterility.
- 3.1.2 <u>durability conditioning</u> requirements, n—material properties relevant to the ability of the package to protect the product. exposure to specific temperature, humidity, and time conditions to simulate particular field conditions, mimic the effects of aging, or to minimize the variation in test results.
- 3.1.3 distribution simulation, n—conditioning or stressing of the packaging system, so that its ability to withstand the hazards of handling, distribution, and storage can subsequently be evaluated.
- 3.1.4 <u>integrity and seal material and package performance attributes and characteristics requirements</u>, n—<u>material properties relevant to</u> the ability of the package to <u>prevent inadvertent escape of contents or entrance of outside substances protect the product</u> while preserving intended opening for use features.
- 3.1.5 package performance, n—the ability of the packaging system, including the sterile barrier system and protective packaging, to withstand the hazards of handling, distribution, and storage distribution and storage as approximated via distribution simulation.
- 3.1.6 *printing printed ink requirements*, *n*—the printed ink properties needed to ensure physical and chemical resistance to degradation.
 - 3.1.7 processing requirements, n—the material characteristics needed to ensure the consistent and reliable production of the package.
 - 3.1.8 safety requirements, n—safeguard product against contamination and deleterious health effects.
 - 3.1.9 *visibility and appearance requirements, n*—the desired package aesthetics needed to permit or inhibit viewing of the product or to enhance product presentation.
 - 3.2 For other terms used in this guide, see Terminology F17.

4. Significance and Use

- 4.1 This design and evaluation guide describes multiple categories for evaluating flexible medical packages and packaging materials. These include safety, barrier properties, durability, package and seal barrier, material and package performance attributes and characteristics, package integrity, visibility and appearance, processing, printing ink properties, and package performance-printed ink, distribution simulation, and conditioning.
- 4.2 The intent of this design and evaluation guide is to evaluate all cited categories and select those that are applicable. Once the product has been characterized and the sterilization methodology has been defined, there are numerous sets of requirements for any specific package. This design and evaluation guide provides an avenue for assessing these requirements and choosing test methods for both evaluating the package design and monitoring package compliance.

⁷ Available from International Safe Transit Association (ISTA), 1400 Abbot Rd., Suite 160, East Lansing, MI 48823-1900, http://www.ista.org.

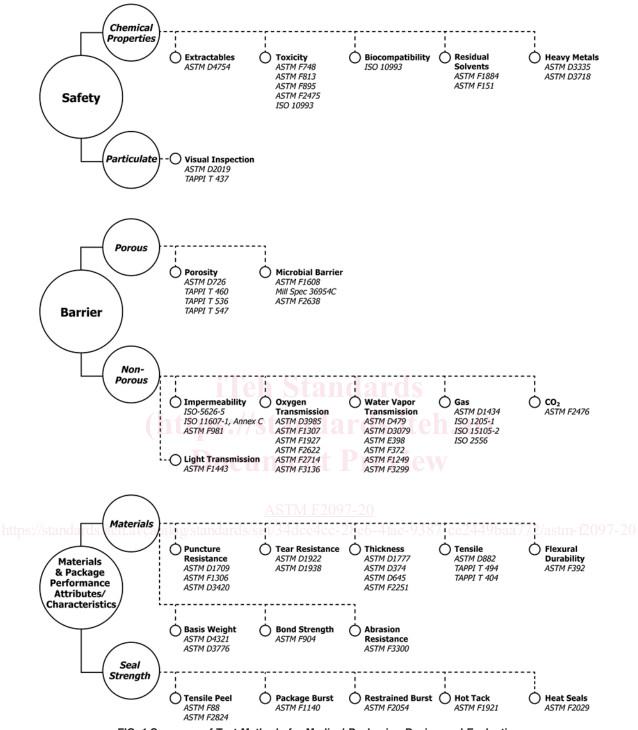


FIG. 1 Summary of Test Methods for Medical Packaging Design and Evaluation

Note 1—Many of the standards included in this guide are consensus standards that are recognized by the United States Food and Drug Administration (FDA). Selection and use of a U.S. FDA recognized consensus standard is voluntary and the sole responsibility of the user in determining its applicability. For further information, consult the U.S. FDA Medical Device Standards—Standards and Conformity Assessment_Program at http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/Standards/default.htm.https://www.fda.gov/medicaldevices/device-advice-comprehensive-regulatoryassistance/standards-and-conformity-assessment-program

4.3 Product characterization shall include mass or weight, geometry (length and width, height, and shape) and product composition.

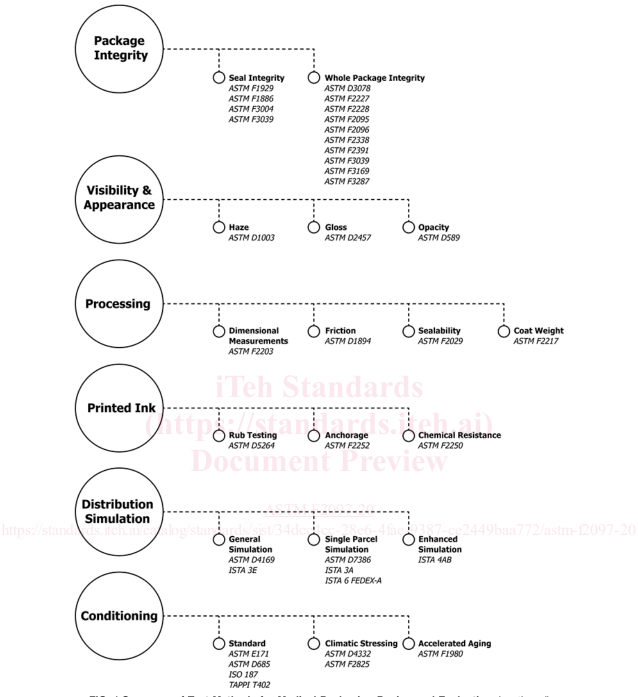


FIG. 1 Summary of Test Methods for Medical Packaging Design and Evaluation (continued)

- 4.4 All categories must be considered for applicability.
- 4.5 The Summary of Test Methods for Medical Packaging Design and Evaluation (Fig. 1) provides a compact graphical presentation of the test methods referenced in this guide.
- 4.6 Test Description and Applicability (see Table 1):
- 4.6.1 Table 1 lists the test methods commonly used to evaluate flexible medical packaging. The test methods are used in two phases.

TABLE 1 Test Description and Applicability Table

			· · · · ·	
Test Category	Test Method	<u>Usage</u>	Description	Applicability
		afety Requirements		
Future etilal		Chemical Properties	This took mother than been all the	
Extractibles Usage	ASTM D4754	This test method covers the use of	This test method has been applied to a variety of migrant/polymer systems in	
R&D evaluation		the FDA migration	contact with numerous foods and food	
		cell in the extraction	simulants. Though most of the migrants	
		of components and	examined were radiolabeled, the use of	
		permits	the FDA cell has been validated for mi-	
		quantitation of indi-	gration studies of unlabeled styrene from	
		vidual migrants from	polystyrene.	
		plastic	This test method has been shown to yield reproducible results under the con-	
		materials by suitable extracting liquids,	ditions for migration tests requested by	
		including liquid	the FDA. However, if the data is to be	
		foods and food-	submitted to the FDA, it is suggested	
		stimulating solvents.	that their guidelines by consulted.	
		This test method		
		provides a two-	Because it employs two-sided extraction,	
		sided, liquid extrac-	this test method may not be suitable for	
		tion test for plastic	multilayered plastics intended for single-	
		materials that can	sided food contact use. The size of the FDA migration cell as	
		be formed into film, sheet, or disks.	described may preclude its use in deter-	
		ones, or dions.	mining total nonvolatile extractives in	
			some cases.	
Safety/Chemical	ASTM D4754	R&D	This test method covers the use of the	This test method has been applied to a
Properties/			FDA migration	variety of migrant/polymer systems in
Extractables			cell in the extraction of components and	contact with numerous foods and food
			permits	simulants. Though most of the migrants
			quantitation of individual migrants from plastic	examined were radiolabeled, the use of the FDA cell has been validated for mi-
			materials by suitable extracting liquids,	gration studies of unlabeled styrene from
			including liquid	polystyrene.
			foods and food-stimulating solvents. This	This test method has been shown to
			test method	yield reproducible results under the con-
			provides a two-sided, liquid extraction	ditions for migration tests requested by
			test for plastic	the FDA. However, if the data is to be
			materials that can be formed into film, sheet, or disks.	submitted to the FDA, it is suggested that their guidelines by consulted.
			Shoot, or disho.	that their galacimes by consulted.
				Because it employs two-sided extraction,
				this test method may not be suitable for
				multilayered plastics intended for single-
				sided food contact use. 097-20
				The size of the FDA migration cell as
				described may preclude its use in determining total nonvolatile extractives in
				some cases.
Toxity	ASTM F748	This practice recom-	The biocompatibility of materials used in	
Usage		mends generic bio-	single-component or multicomponent	
R&D evaluation		logical test methods	medical devices for human use depends	
		for materials and	to a large degree on the particular nature	
		devices according to	of the end-use application. It is not pos-	
		end-use applica- tions. Tests include	sible to specify a set of biocompatibility test methods which will be necessary	
		those performed on	and sufficient to establish biocompatibility	
			for all materials and applications.	
		materials, end		
		materials, end products, and ex-	While chemical testing for extractable	
		products, and ex- tracts. Rationale and comments on	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary	
		products, and ex- tracts. Rationale and comments on current state of the	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing	
		products, and ex- tracts. Rationale and comments on current state of the art are included for	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice.	
		products, and ex- tracts. Rationale and comments on current state of the art are included for all test pro-	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of	
		products, and ex- tracts. Rationale and comments on current state of the art are included for all test pro- cedures described.	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rap-	
		products, and ex- tracts. Rationale and comments on current state of the art are included for all test pro- cedures described. Biological evaluation	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods	
		products, and ex- tracts. Rationale and comments on current state of the art are included for all test pro- cedures described. Biological evaluation of materials	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by	
		products, and ex- tracts. Rationale and comments on current state of the art are included for all test pro- cedures described. Biological evaluation of materials and devices, and	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by necessity only a guideline. These test	
		products, and extracts. Rationale and comments on current state of the art are included for all test procedures described. Biological evaluation of materials and devices, and related subjects	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by necessity only a guideline. These test protocols are intended to apply to materi-	
		products, and ex- tracts. Rationale and comments on current state of the art are included for all test pro- cedures described. Biological evaluation of materials and devices, and related subjects such as pyrogen	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by necessity only a guideline. These test	
		products, and extracts. Rationale and comments on current state of the art are included for all test procedures described. Biological evaluation of materials and devices, and related subjects	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by necessity only a guideline. These test protocols are intended to apply to materials and medical devices for human appli-	
		products, and extracts. Rationale and comments on current state of the art are included for all test procedures described. Biological evaluation of materials and devices, and related subjects such as pyrogen testing and batch	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by necessity only a guideline. These test protocols are intended to apply to materials and medical devices for human appli-	
		products, and ex- tracts. Rationale and comments on current state of the art are included for all test pro- cedures described. Biological evaluation of materials and devices, and related subjects such as pyrogen testing and batch testing of production	While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by necessity only a guideline. These test protocols are intended to apply to materials and medical devices for human appli-	



Test Category	Test Method	Usage	Description	Applicability
Safety/Chemical Properties/ Toxicity - Biocompatibility	ASTM F748	R&D	This practice recommends generic biological test methods for materials and devices according to end-use applications. Tests include those performed on materials, end products, and extracts. Rationale and comments on current state of the art are included for all test procedures described. Biological evaluation of materials and devices, and related subjects such as pyrogen testing and batch testing of production lots are also discussed.	The biocompatibility of materials used in single-component or multicomponent medical devices for human use depends to a large degree on the particular nature of the end-use application. It is not possible to specify a set of biocompatibility test methods which will be necessary and sufficient to establish biocompatibility for all materials and applications. While chemical testing for extractable additives and residual monomers or residues from processing aids is necessary for most implant materials, such testing is not included as a part of this practice. The reader is cautioned that the area of materials biocompatibility testing is a rapidly evolving field, and improved methods are evolving rapidly, so this practice is by necessity only a guideline. These test protocols are intended to apply to materials and medical devices for human application.
Toxity Usage R&D evaluation	ASTM F813	This practice describes a reference method of direct contact cell culture testing that may be used in evaluating the cytotoxic potential of materials for use in the construction of medical materials and devices. This practice may be used either directly to evaluate materials or as a reference against which other cytotoxicity test methods	This practice tends to be used less frequently due to the risk of inducing a response from mechanical damage due to direct placement of the sample onto the cell layer. This practice may be suitable for products which have leachates that are not able to diffuse through agar and are not too heavy.	
Safety/Chemical Properties/ Toxicity - Biocompatibility	ASTM F813 itch.ai/catalog/s	may be compared: R&D AST tandards/sist/34d	This practice describes a reference method of direct contact cell culture testing that may be used in evaluating the cytotoxic potential of materials for use in the construction of medical materials and devices. This practice may be used either directly to evaluate materials or as a reference against which other cytotoxicity test methods may be compared.	This practice tends to be used less frequently due to the risk of inducing a response from mechanical damage due to direct placement of the sample onto the cell layer. This practice may be suitable for products which have leachates that are not able to diffuse through agar and are not too heavy.

4.6.1.1 Package Design: Characterization of the Materials and Evaluation of the Resultant Package—This is referred to as "R&D Evaluation" in Table 1. Testing during this phase is characterized by the generation of quantitative data on the performance of the component materials and the package assembly. These test methods are lengthy, making them inappropriate for the manufacturing environment where rapid response is required for process control. Often, they are expensive and require specialized equipment not readily available at a medical packaging or device manufacturing facility.

4.6.1.2 Package Compliance: Routine Monitoring of Adherence to Specifications—This is referred to as "Compliance Testing" in Table 1. Testing during this phase must be rapid, inexpensive, and readily implemented in a manufacturing environment. The objective is not to develop design data, but to ensure that the design specifications are being met. These test methods do not necessarily make direct measurements of critical values, but detect variations in material, process, or product that are indicative of all critical characteristics.

		TABL	E 1 Continued	
Test Category	Test Method	<u>Usage</u>	Description	Applicability
Test Category Toxicity Usage R&D evaluation Safety/Chemical Properties/ Toxicity - Biocompatibility https://standards.it	ASTM F895	The agar diffusion assay is an indirect contact test in which the test material is placed onto an agar layer that protects the cells. This test method is commonly used to evaluate the response of small samples that have at least one flat surface such as elastomeric closures.	This is one of a series of reference test methods for the assessment of cytotoxic potential, employing different techniques. Assessment of cytotoxicity is one of several tests employed in determining the biological response to a material, as recommended in Practice F748. This test method is appropriate for materials in a variety of shapes and for materials that are not necessarily sterile. This test method would be appropriate in situations where the amount of material is limited. For example, small devices or powders could be placed on the agar and the presence of a zone of inhibition of cell growth could be examined. While the agar layer can act as a cushion to protect the cells from the specimen, there may be materials which are sufficiently heavy to compress the agar and prevent diffusion or to cause mechanical damage to the cells. This test method is not appropriate for leachables that are not water soluble because they may not diffuse through agar or agarose and thus not be detected. This test method would not be appropriate for these materials. The L 929 cell line was chosen because it has a significant history of use in assays of this type. This is not intended to imply that its use is preferred; only that the L 929 is an established cell line, well characterized and readily available, that has demonstrated reproducible results in several laboratories. The agar diffusion assay is an indirect contact test in which the test material is placed onto an agar layer that protects the cells. This test method is commonly used to evaluate the response of small samples that have at least one flat surface such as elastomeric closures.	This is one of a series of reference test methods for the assessment of cytotoxic potential, employing different techniques. Assessment of cytotoxic potential, employing different techniques. Assessment of cytotoxicity is one of several tests employed in determining the biological response to a material, as recommended in Practice F748. This test method is appropriate for materials in a variety of shapes and for materials that are not necessarily sterile. This test method would be appropriate in situations where the amount of material is limited. For example, small devices or powders could be placed on the agar and the presence of a zone of inhibition of cell growth could be examined. While the agar layer can act as a cushion to protect the cells from the specimen, there may be materials which are sufficiently heavy to compress the agar and prevent diffusion or to cause mechanical damage to the cells. This test method is not appropriate for leachables that are not water soluble because they may not diffuse through agar or agarose and thus not be detected. This test method would not be appropriate for these materials. The L 929 cell line was chosen because it has a significant history of use in assays of this type. This is not intended to imply that its use is preferred; only that the L 929 is an established cell line, well characterized and readily available, that has demonstrated reproducible results in several laboratories.



Test Category	Test Method	<u>Usage</u>	Description	Applicability
Texicity Usage R&D evaluation	ASTM F2475	This guide provides information to determine the appropriate testing for biocompatibility of packaging materials used to contain a medical	This method applies to packaging for medical devices.	
Safety/Chemical Properties/ Toxicity - Biocompatibility	<u>ASTM F2475</u>	device. R&D Evaluation	This guide provides information to determine the appropriate testing for biocompatibility of packaging materials used to contain a medical device.	This method applies to packaging for medical devices.
Biocompatibility Usage R&D-evaluation	ISO-10993	This entails a series of standards for evaluating the biocompatibility of a medical device prior to clinical study. Part 1 of the Standard uses an approach to test selection thatis very similar to the Tripartite Guidance.	When selecting the appropriate tests for biological evaluation of a medical device, one must consider the chemical characteristics of device materials and the nature, degree, frequency, and duration of its exposure to the body. Note: FDA has made several modifications to the testing required by ISO 10993—Part 1.	
Safety/Chemical Properties/ Toxicity - Biocompatibility	<u>ISO 10993</u>	R&D Evaluation ITCh S DS://Sta	This entails a series of standards for evaluating the biocompatibility of a medical device prior to clinical study. Part 1 of the standard uses an approach to test selection that is very similar to the Tripartite Guidance.	When selecting the appropriate tests biological evaluation of a medical devione must consider the chemical charateristics of device materials and the nature, degree, frequency, and duratic of its exposure to the body. Note: FD/has made several modifications to the testing required by ISO 10993–Part 1.
Retained solvents Usage R&D evaluation Compliance testing	ASTM F1884 ds.iteh.ai/catalog/	This test method covers determination of the amount of residual solvents released from within a packaging	This test method does not address the determination of total retained solvents in a packaging material. Techniques such as multiple headspace extraction can be employed to this end. For purposes of verifying the identity of or identifying unknown volatile	
		material contained in a sealed vial under a given set of time and temperature conditions and is a recommended alternative for Test Method F151. This test method covers a procedure for quantifying volatile compounds whose identity has been established, and are retained in packaging materials.	compounds, the analyst is encouraged to incorporate techniques such as gas chromatography/mass spectroscopy, gas chromatography/infrared spectroscopy, or other suitable techniques in conjunction with this test method. This is an off-line head space analysis. It is sensitive to technique and sampling equipment resulting in large variations (-25 %) between laboratories. It is a simplified version of Test Method F151 providing about the same level of accuracy. This method differs from Test Method F151 in that it specifies certain conditions. Test Method F1884, for example, specifies a pre-heat condition of 90°C for 20 min. Test Method F151 defines a procedure for determining optimum heating time and temperature conditions for the preheat. Because solvents will escape from surface wraps on a roll of film, this test	
			sace wraps on a roll of film, this test should be performed immediately after manufacturing to provide an indication of solvent levels in the inner wraps of the roll of film.	

4.6.2 It is important to note that no individual test method is entirely predictive of final package performance. Filled packages must be evaluated under conditions of use.

		TABL	E 1 Continued	
Test Category	Test Method	<u>Usage</u>	Description	Applicability
Safety/Chemical Properties/ Retained Solvents	ASTM F1884	R&D Evaluation Compliance Testing	These test methods cover determination of the amount of residual solvents released from within a packaging material contained in a sealed vial under a given set of time and temperature conditions and is a recommended alternative for Test Method F151. These test methods cover a procedure for quantifying volatile compounds whose identity has been established, and are retained in packaging materials.	These test methods do not address the determination of total retained solvents in a packaging material. Techniques such as multiple headspace extraction can be employed to this end. For purposes of verifying the identity of or identifying unknown volatile compounds, the analyst is encouraged to incorporate techniques such as gas chromatography/mass spectroscopy, gas chromatography/mass spectroscopy, gas chromatography/infrared spectroscopy, conther suitable techniques in conjunction with this test method. This is an off-line head space analysis. It is sensitive to technique and sampling equipment resulting in large variations (~25 %) between laboratories. It is a simplified version of Test Method F151 providing about the same level of accuracy. This method differs from Test Method F151 in that it specifies certain conditions. Test Methods F1884, for example, specifies a pre-heat condition of 90 °C for 20 min. Test Method F151 defines a procedure for determining optimum heating time and temperature conditions for the preheat. Because solvents will escape from surface wraps on a roll of film, this test should be performed immediately after manufacturing to provide an indication of solvent levels in the inner wraps of the roll of film.
Retained solvents Usage R&D evaluation Compliance testing	ASTM F151	This test method provides an index for comparing the level of solvents retained in flexible barrier materials of the same construction, which result from easting,	This method is essentially identical to Test Method F1884 except for a complicated determination of the optimum heating time and temperature for the films in the head space container. There is no improvement in the interlaboratory variation. All other comments under Test Method F1884 apply equally to Test Method F151.	
		coating, printing, or laminat- ing operations. This test method does not yield abso- lute quantitative measurements of solvents retained in flexible barrier mate- rials.		
Safety/Chemical Properties/ Retained Solvents	ASTM F151	R&D Evaluation Compliance Testing	This test method provides an index for comparing the level of solvents retained in flexible barrier materials of the same construction, which result from casting, coating, printing, or laminating operations. This test method does not yield absolute quantitative measurements of solvents retained in flexible barrier materials.	This method is essentially identical to Test Methods F1884 except for a complicated determination of the optimum heating time and temperature for the films in the head space container. There is no improvement in the interlaboratory variation. All other comments under Test Method F1884 apply equally to Test Method F151.

4.6.2.1 Distribution simulation is intended to provide a standardized, uniform, and repeatable basis of conditioning or stressing a package system so that the packaging system's ability to withstand routine distribution can subsequently be evaluated. The damage-producing motions, forces, conditions, and sequences of transport environments are simulated within a laboratory setting. The intended test sequences and intensity levels can be adjusted to address the specifics of the distribution cycle.

solvents retained in flexible barrier mate-



Test Category	Test Method	<u>Usage</u>	Description	Applicability
Heavy metals Usage R&D evaluation	ASTM D3335	This test method covers the determination of lead contents between 0.01 and 5%, cadmium contents between 50 and 150 ppm (mg/kg), and cobalt contents between 50 and 2000 ppm (mg/kg) present in the nonvolatile portion of liquid coating or contained in dried films by means of atomic	Higher levels of all three elements can be determined by this test method, provided that appropriate dilutions and adjustments in specimen size and reagent quantities are made. This test method is not applicable to the determination of lead in samples containing antimony pigments (low recoveries are obtained). If lead is present in the sample to be analyzed in the form of an organic lead compound at a concentration greater than 0.1 %, small losses of lead may occur, resulting in slightly poorer precision.	
Safety/Chemical Properties/ Heavy Metals	<u>ASTM D3335</u>	absorption- R&D Evaluation	This test method covers the determination of lead contents between 0.01 and 5 %, cadmium contents between 50 and 150 ppm (mg/kg), and cobalt contents between 50 and 2000 ppm (mg/kg) present in the nonvolatile portion of liquid coating or contained in dried films by means of atomic absorption.	Higher levels of all three elements can be determined by this test method, provided that appropriate dilutions and adjustments in specimen size and reagen quantities are made. This test method in not applicable to the determination of lead in samples containing antimony piments (low recoveries are obtained). If lead is present in the sample to be analyzed in the form of an organic lead corpound at a concentration greater than 0.1 %, small losses of lead may occur, resulting in slightly poorer precision.
Heavy metals Usage R&D evaluation	ASTM D3718 (htt	This test method eovers the determination of the content of chromium (including chromium oxide) in the range from 0.005 to 1.0 % present in the solids of liquid coatings or in dried films obtained from previously coated sub-	Higher concentrations of chromium can be determined by this test method provided that appropriate dilutions and adjustments in specimen size and reagent quantities are made:	
		strates by means of atomic absorption.		
Safety/Chemical Properties/ Heavy Metals	<u>ASTM D3718</u>	R&D Evaluation	This test method covers the determination of the content of chromium (including chromium oxide) in the range from 0.005 to 1.0 % present in the solids of liquid coatings or in dried films obtained from previously coated substrates by means of atomic absorption.	Higher concentrations of chromium can be determined by this test method pro- vided that appropriate dilutions and ad- justments in specimen size and reagen quantities are made.
Particulate			27saile of atomic absorption.	

- 4.6.2.2 Performance testing must be conducted to confirm product protection and sterile package integrity in design validation. Confirmation is by means of product testing and package integrity testing. The specific tests used for evaluation will be dependent upon the product and the sterile barrier system. Consideration should be given to evaluating the effects of environmental extremes.
- 4.7 Once the design of the package and/or packaging materials has been determined, it may be appropriate to create a package and/or material specification. Guides F99 or F2559 may provide useful guidance.

Safety/Particulater ASTM D2019 ASTM D201	Test Category				
Safety Particulate/ Image: Interest of the second of the s		Test Method	<u>Usage</u>	Description	Applicability
intended-for-the-rum mencial estimation of-dirt in paper-or paper-board in terms of-equivalent black area. This test method is easitified such as personal or the estimation of visual inspection ASTM D2019 AST	Visual inspection Usage Compliance testing	TAPPI T 437	suited for the visual estimation of dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard is defined as any foreign matter embedded in the sheet, which, when examined by reflected, not transmitted, light has a contrasting eolor to the rest of the surface and has an equivalent black area of 0.04 mm² or	characteristics other than dirt. Frequently used for estimation of gels, fisheyes, ink	
eharacteristics and It may be entirely inadequate when norwisual effects such as grittiness of dirt are of importance. This is ASTM's version of TAPPI T 437 This test method is intended for the numerical estimation of visual characteristics other than dirt. Frequently used for estimation of gels, sisheyes, in splashes, and other visual estimation of hir in paper or paperboard in terms of dirt are of importance. This is ASTM's version of TAPPI in test method is suited for the visual estimation of hir in paper or paperboard in terms of dirt are of importance. This is ASTM's version of TAPPI in the test method is suited for the visual estimation of hir in paper or paperboard in terms of dirt are of importance. This is ASTM's version of TAPPI in the test method is suited for the visual estimation of hir in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of the visual estimation of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suitable of the visual estimation of dirt in paper or paperboard in the suit	Visual-inspection Usage Compliance testing	ASTM D2019	intended for the nu- merical estimation of clirt in paper or paperboard in terms of equivalent black area. This test method is satisfae-	characteristics other than dirt. Frequently used for estimation of gels, fisheyes, ink splashes, and other visual defects.	
the TAPPI Dirt Estimation Chart. R&D Evaluation This test method is intended for the numerical estimation of dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard in terms of equivalent black area of 0.04 mm² or over. This test method can be used to size characteristics other than dirt. Frequently used for estimation of gels, fisheyes, ink splashes, and other visual estimation of dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperbo			characteristics and it may be entirely inadequate when nonvisual effects such as gritiness of dirt are of importance. This is ASTM's version of TAPPI		
ASTM D2019 R&D Evaluation This test method is intended for the numerical estimation of dirt in paper or paperboard in terms of equivalent black area. This test method is satisfactory only for the estimation of visual characteristics and it may be entirely inadequate when nonvisual effects such as grittiness of dirt are of importance. This is ASTM's version of TAPPI T 437 TAPPI T 437 Compliance testing This test method can be used to size characteristics other than dirt. Frequently used for estimation of gels, fisheyes, ink splashes, and other visual estimation Chart. This test method can be used to size characteristics other than dirt. Frequently used for estimation of gels, fisheyes, ink splashes, and other visual estimation of dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard is defined as any foreign matter embedded in the sheet, which, when examined by reflected, not transmitted, light has a contrasting color to the rest of the surface and has an equivalent black area of 0.04 mm² or over.			the TAPPI Dirt Esti-		
estimation of dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard is defined as any foreign matter embedded in the sheet, which, when examined by reflected, not transmitted, light has a contrasting color to the rest of the surface and has an equivalent black area of 0.04 mm ² or over.	Safety/Particulate/ Visual inspection	ASTM D2019		merical estimation of dirt in paper or paperboard in terms of equivalent black area. This test method is satisfac- tory only for the	characteristics other than dirt. Frequently used for estimation of gels, fisheyes, ink splashes, and other visual
Barrier Requirements				may be entirely inadequate when nonvisual effects such as grittiness of dirt are of importance. This is ASTM's version of TAPPI T 437. It refers to the TAPPI Dirt Estima-	
		<u>TAPPI T 437</u>	Compliance testing	may be entirely inadequate when nonvisual effects such as grittiness of dirt are of importance. This is ASTM's version of TAPPI T 437. It refers to the TAPPI Dirt Estimation Chart. This test method is suited for the visual estimation of dirt in paper or paperboard in terms of equivalent black area. Dirt in paper or paperboard is defined as any foreign matter embedded in the sheet, which, when examined by reflected, not transmitted, light has a contrasting color to the rest of the surface and has an equivalent black area of 0.04 mm² or	characteristics other than dirt. Frequently used for estimation of gels, fisheyes, ink splashes, and other visua



Test Category	Test Method	<u>Usage</u>	Description	Applicability
Porosity Usage R&D evaluation Gompliance testing	ASTM D726	This test method is applicable in papers that permit the passage of up to 25 mL of air/0.785 in.2 in 15 s.	This test method cannot be used in those cases where the paper cannot be clamped securely against surface and edge leakage, such as, crepe or corrugated papers. For testing porous and semiporous paper, refer to TAPPLT 460, and T 536, respectively. Since the three test methods (D726, T 460, and T 436) do not give the same results, it is recommended that a specific method be agreed upon in specifications covering paper between the seller and the purchaser, and that the test method be chosen to conform to the principle range.	
Barrier/Porous/ Porosity	ASTM D726	R&D Evaluation Compliance Testing	This test method is applicable in papers that permit the passage of up to 25 mL of air/0.785 in. ² in 15 s.	This test method cannot be used in those cases where the paper cannot be clamped securely against surface and edge leakage, such as, crepe or corrugated papers. For testing porous and semiporous paper, refer to TAPPI T 46 and T 536, respectively. Since the three test methods (D726, T 460, and T 436) do not give the same results, it is recommended that a specific method be agreed upon in specifications covering paper between the seller and the purchaser, and that the test method be chosen to conform to the principle range.
Porosity Usage	TAPPLT 460	This test method references the use	The pressure differential used in this test method is 1.22 kPa. The recommended	
R&D evaluation		of a Gurley densometer that	range of time measured is from 5 to	
Compliance testing		measures the	1800 sec/100-mL cylinder displacement. For more impermeable materials the time	
		amount of time re-	requirements become so excessive that	
		quired for a certain volume	other techniques are preferable. Since this test method measures air passage	
		of air (100 cm ³) to	through the specimen, as well as, leak-	
		pass through a test specimen of a	age across the surface, it is unsuitable for rough-surfaced materials that can not	
		given area. The air	be securely clamped in the mechanism	
		pressure is	and may allow significant surface and	
		generated by a gravity-loaded cylin- der that captures an	edge leakage. For measurement of materials at higher pressure (3 kPa) refer to TAPPLT 536.	
		air volume within a chamber using a liquid seal. The	To measure materials at pressures up to 9.85 kPa, TAPPI T 547 references the use of a Sheffield tester which measures	
		pressurized volume of air is directed to	the amount of air passing through a ma- terial of a given area over a specific time	
		the clamping gasket ring, which	period.	
		holds the test speci- men. Air that		
		passes through an area of the test		
		specimen of 6.4 cm ² (1 in. ²) es-		
		capes to atmo-		
		sphere through the		
		holes in the down- stream clamping		
		ou oum olumping		



			E i Continueu	
Test_Category	Test Method	<u>Usage</u>	Description	Applicability
Barrier/Porous/ Porosity	<u>TAPPI T 460</u>	R&D Evaluation Compliance Testing	This test method references the use of a Gurley densometer that measures the amount of time required for a certain volume of air (100 cm³) to pass through a test specimen of a given area. The air pressure is generated by a gravity-loaded cylinder that captures an air volume within a chamber using a liquid seal. The pressurized volume of air is directed to the clamping gasket ring, which holds the test specimen. Air that passes through an area of the test specimen of 6.4 cm² (1 in.²) escapes to atmosphere through the holes in the downstream clamping plate.	The pressure differential used in this temethod is 1.22 kPa. The recommended range of time measured is from 5 to 1800 s/100-mL cylinder displacement. For more impermeable materials the timerquirements become so excessive that other techniques are preferable. Since this test method measures air passage through the specimen, as well as, leakage across the surface, it is unsuitable for rough-surfaced materials that cannot be securely clamped in the mechanism and may allow significant surface and edge leakage. For measurement of materials at higher pressure (3 kPa) refer to TAPPI T 536. To measure materials at pressures up to 9.85 kPa, TAPPI T 547 references the use of a Sheffield tester which measure the amount of air passing through a material of a given area over a specific time period.
Porosity Usage R&D evaluation Compliance testing	TAPPLT 536	10 mL of air to pass	This test method cannot be used in those cases where the paper cannot be clamped securely against surface and edge leakage, such as, crepe or corrugated papers. Since the three test methods (D726, T 460, and T 536) do not give the same results, it is recommended that a specific test method be agreed upon in specifications covering paper between the seller and the purchaser, and that the test method be chosen to conform to the principle range.	
Barrier/Porous/ Porosity https://standards	TAPPIT 536 s.iteh.ai/catalog/s	through: R&D Evaluation Compliance Testing tandards/sist/34c	This test method measures the amount of time required for a certain volume of air to pass through a test specimen of a given size. This test method measures at a higher pressure differential (3 kPa) and is recommended for papers that require 10 or more seconds for 10 mL of air to pass through.	This test method cannot be used in those cases where the paper cannot be clamped securely against surface and edge leakage, such as, crepe or corrugated papers. Since the three test methods (D726, T 460, and T 536) do not give the same results, it is recommend that a specific test method be agreed upon in specifications covering paper between the seller and the purchaser, and that the test method be chosen to conform to the principle range.
Poresity Usage R&D evaluation Compliance testing	TAPPI T 547	This method is used to measure the air permeance of a circular area of paper using a pressure differential of approximately 10 kPa (1.5 psig).	In order to accommodate a wide range of paper products, rubber clamping plates are available for five commonly used orifice diameters: 9.5 mm (0.375 in.), 19.1 mm (0.75 in.), 38.1 mm (1.5 in.), 57.2 mm (2.25in.), and 76.2 mm (3.00 in.). The air flow range for this method is 0 to 3348 mL/min (0 to 400 Sheffield units). Instruments are available with either variable flowmeters (glass tubes with internal tapers and floats) or electronic mass flowmeters.	