

Designation: F2981 – 15

Standard Test Method for Verifying Nonporous Flexible Barrier Material Resistance to the Passage of Air¹

This standard is issued under the fixed designation F2981; 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 test method is to be used to verify a specific material design property. Some flexible barrier materials are designed to have a resistance to the passage of air through the membrane structure. These materials are characterized as nonporous. This test method provides a means to verify this property by challenging a material with a given volume of air under pressure over a specific time period.

1.2 This test method is not intended to measure the diffusion properties of a material nor to identify or quantify the presence of pinhole damage to the design that may result in leaks.

1.3 *Units*—The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

1.4 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:²

- E171 Practice for Conditioning and Testing Flexible Barrier Packaging
- 2.2 ISO Standards:³
- ISO 11607-1 Packaging for Terminally Sterilized Medical Devices—Part 1: Requirements for Materials, Sterile Barrier Systems and Packaging Systems
- ISO 5636-5 Paper and Board—Determination of Air Permeance and Air Resistance (Medium Range)—Part 5: Gurley Method

2.3 *TAPPI Standard*:⁴ TAPPI Test Method T 460 om-06 Air Resistance of Paper (Gurley Method)

3. Significance and Use

3.1 This material challenge is presented in ISO-11607-1 Annex C as a normative test method to demonstrate that a material is nonporous and satisfies the microbial barrier requirements.

4. Apparatus

4.1 *Gurley Cylinder-type Densometer*, or equivalent apparatus compliant with ISO 5636-5.

4.1.1 *Air Volume and Pressure*, as standardized in TAPPI T-460 and ISO 5636-5.

4.2 Clock or Timer.

4.3 Dimensional Scale, in 0.1 millimeter increments.

5. Conditioning

5.1 Conditioning of samples will depend on the material under evaluation. If conditioning before testing is appropriate, normal and desirable, then condition the test specimens following ASTM E171.

6. Preparation and Procedure

6.1 Measure and cut sample of material to be tested approximately 50 mm \times 50 mm square. Other cut sizes of samples can be used if easier to manipulate and position in holder without damage, wrinkling or introducing leaks.

6.2 Prior to inserting the material, ensure that the column is raised in the ready position. Loosen the wheel on the bottom of densometer and place the web between the clamps. By inserting the material samples with the smoother side facing upward toward the cylinder, the risk of leaking around the clamp is minimized.

6.3 Rotate the wheel to the right to tighten until the material is secure. (See Fig. 1.)

¹ This test method is under the jurisdiction of ASTM Committee F02 on Flexible Barrier Packaging and is the direct responsibility of Subcommittee F02.20 on Physical Properties.

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

⁴ Available from Technical Association of the Pulp and Paper Industry (TAPPI), 15 Technology Parkway South, Norcross, GA 30092, http://www.tappi.org.