



Designation: D4895 – 18 (Reapproved 2023)

Standard Specification for Polytetrafluoroethylene (PTFE) Resin Produced From Dispersion¹

This standard is issued under the fixed designation D4895; 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.

This standard has been approved for use by agencies of the U.S. Department of Defense.

1. Scope

1.1 This specification² covers polytetrafluoroethylene (PTFE) prepared by coagulation of a dispersion. These PTFE resins are homopolymers of tetrafluoroethylene or modified homopolymers containing not more than 1 % by weight of other fluoromonomers. The materials covered herein do not include mixtures of PTFE with additives such as colors, fillers, or plasticizers; nor do they include reprocessed or reground resin or any fabricated articles because the properties of such materials have been irreversibly changed when they were fibrillated or sintered.

1.2 The values stated in SI units as detailed in [IEEE/ASTM SI-10](#) are to be regarded as standard. The values given in parentheses are for information only.

1.3 The following safety hazards caveat pertains only to the Specimen Preparation Section, Section 9, and the Test Methods Section, Section 10, of this specification: *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, health, and environmental practices and determine the applicability of regulatory limitations prior to use.* See Warning note in 9.1.1 for a specific hazards statement.

NOTE 1—Information in this specification is technically equivalent to related information in ISO 20568-1 and ISO 20568-2.

1.4 *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.*

¹ This specification is under the jurisdiction of ASTM Committee D20 on Plastics and is the direct responsibility of Subcommittee D20.15 on Thermoplastic Materials.

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² Specifications for other forms of polytetrafluoroethylene are found in Specifications D4441 and D4894.

2. Referenced Documents

2.1 ASTM Standards:³

- D618 Practice for Conditioning Plastics for Testing
- D638 Test Method for Tensile Properties of Plastics
- D792 Test Methods for Density and Specific Gravity (Relative Density) of Plastics by Displacement
- D883 Terminology Relating to Plastics
- D1708 Test Method for Tensile Properties of Plastics by Use of Microtensile Specimens
- D1895 Test Methods for Apparent Density, Bulk Factor, and Pourability of Plastic Materials
- D3892 Practice for Packaging/Packing of Plastics
- D4052 Test Method for Density, Relative Density, and API Gravity of Liquids by Digital Density Meter
- D4441 Specification for Aqueous Dispersions of Polytetrafluoroethylene
- D4591 Test Method for Determining Temperatures and Heats of Transitions of Fluoropolymers by Differential Scanning Calorimetry
- D4894 Specification for Polytetrafluoroethylene (PTFE) Granular Molding and Ram Extrusion Materials
- E11 Specification for Woven Wire Test Sieve Cloth and Test Sieves
- E29 Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications
- E177 Practice for Use of the Terms Precision and Bias in ASTM Test Methods
- IEEE/ASTM SI-10 Use of the International System of Units (SI): The Modern Metric System

2.2 ISO Standards:⁴

- ISO 20568-1 Plastics Fluoropolymer Dispersions and Molding and Extrusion Materials—Part 1: Designation and Specification
- ISO 20568-2 Plastics Fluoropolymer Dispersions and Molding and Extrusion Materials—Part 2: Preparation of

³ 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.

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

*A Summary of Changes section appears at the end of this standard

Test Specimens and Determination of Properties
ISO 13322-2 Particle size analysis—Image analysis
methods—Part 2: Dynamic image analysis methods

3. Terminology

3.1 Definitions—The definitions given in Terminology D883 are applicable to this specification.

3.2 Definitions of Terms Specific to This Standard:

3.2.1 bulk density, *n*—the mass in grams per litre of resin measured under the conditions of the test.

3.2.2 extended specific gravity (ESG), *n*—the specific gravity of a specimen of PTFE material molded as described in this specification and sintered (see 3.2.7) for an extended period of time, compared to the sintering time for the measurement of SSG (see 3.2.8), using the appropriate sintering schedule given in this specification.

3.2.3 lot, *n*—one production run or a uniform blend of two or more production runs.

3.2.4 preforming, *vb*—compacting powdered PTFE material under pressure in a mold to produce a solid object, called a preform, that is capable of being handled. Molding and compaction are terms used interchangeably with preforming for PTFE.

3.2.5 reground resin, *n*—resin produced by grinding PTFE material that has been preformed but has never been sintered.

3.2.6 reprocessed resin, *n*—resin produced by grinding PTFE material that has been preformed and sintered.

3.2.7 sintering, *n*—as it applies to PTFE, a thermal treatment during which the PTFE is melted and recrystallized by cooling with coalescence occurring during the treatment.

3.2.8 standard specific gravity (SSG), *n*—the specific gravity of a specimen of PTFE material molded as described in this specification and sintered using the appropriate sintering schedule given in this specification.

3.2.9 strained specific gravity (strained SG), *n*—the specific gravity of a specimen of PTFE material molded, sintered, and strained as described in this specification.

3.2.10 stretching void index (SVI), *n*—a measure of the change in specific gravity of PTFE material which has been subjected to tensile strain as described in this specification.

3.2.11 thermal instability index (TII), *n*—a measure of the decrease in molecular weight of PTFE material which has been heated for a prolonged period of time.

3.2.12 unstrained specific gravity (USG), *n*—the specific gravity, prior to straining, of a specimen of PTFE material used in the Stretching Void Index Test (see 10.9) of this specification.

4. Classification

4.1 This specification covers the following types of PTFE:

4.1.1 Type I and Type II—Resin produced from dispersion. Each type of resin has the same requirements for bulk density, particle size, water content, melting peak temperature, tensile, and elongation. Each type of resin is divided into grades in accordance with standard specific gravity (SSG), Thermal Stability Index (TII), and Stretching Void Index (SVI). Grades are divided into classes according to extrusion pressure.

NOTE 2—See Tables 1 and 2 for details about grades and classes.

4.2 A line callout system is used to specify materials in this specification. The system uses predefined cells to refer to specific aspects of this specification, as illustrated as follows:

Specification				
Standard Number Block	Type	Grade	Class	Special Notes
Example: Specification D4895 - XX	I	2	C	

For this example, the line callout would be Specification D4895 - XX, I2C, and would specify a coagulated dispersion form of polytetrafluoroethylene that has all of the properties listed for that type, grade, and class in the appropriate specified properties or tables, or both, in the specification identified. A comma is used as the separator between the standard number and the type. Separators are not needed between the type, grade, and class.⁵

5. Mechanical Properties

5.1 The resins covered by this specification shall be in accordance with the requirements prescribed in Tables 1 and 2, when tested by the procedures specified herein.

6. Other Requirements

6.1 The resin shall be uniform and shall contain no additives or foreign material.

6.2 The color of the material as shipped by the supplier shall be natural white.

⁵ See the Form and Style for ASTM Standards manual, available from ASTM Headquarters.

TABLE 1 Detail Requirements for all Types,^A Grades and Classes

Type	Bulk Density, g/L	Particle Size Average Diameter, μm	Water Content, max, %	Melting Peak Temperature, °C		Tensile Strength, min, MPa	Elongation at Break, min, %
				Initial	Second		
I	550 ± 150	500 ± 200	0.04	^B	327 ± 10	19	200
II	550 ± 150	1050 ± 350	0.04	^B	327 ± 10	19	200

^AThe types, grades, and classes are not the same as those in previous editions of Specification D4895.

^BGreater than 5.0°C above the second melting peak temperature.

TABLE 2 Detail Requirements for All Types,^A Grades and Classes

Type	Grade	Class	Standard Specific Gravity		Extrusion Pressure, MPa	Thermal Instability Index, max	Stretching Void Index, max
			min	max			
I	1	A	2.14	2.18	5 to <15 ^B	50	NA ^C
		B	2.14	2.18	15 to <55 ^D	50	NA ^C
		C	2.14	2.18	15 to <75 ^E	50	NA ^C
	2	A	2.17	2.25	5 to <15 ^B	50	NA ^C
		B	2.17	2.25	15 to <55 ^D	50	NA ^C
		C	2.17	2.25	15 to <75 ^E	50	NA ^C
	3	C	2.15	2.19	15 to <75 ^E	15	200
		D	2.15	2.19	15 to <65 ^E	15	100
		E	2.15	2.19	15 to <65 ^E	50	200
	4	B	2.14	2.16	15 to <55 ^D	15	50
		A	2.14	2.25	5 to <15 ^B	50	NA ^C
	II	1	A	2.14	2.25	5 to <15 ^B	50

^AThe types, grades, and classes are not the same as those in previous editions of Specification D4895.

^BTested at a reduction ratio of 100:1 (reduction ratio is the ratio of the cross-sectional area of the preform to the cross-sectional area of the die).

^CNot applicable.

^DTested at a reduction ratio of 400:1.

^ETested at a reduction ratio of 1600:1.

6.3 For purposes of determining conformance, all specified limits for this classification system are absolute limits, as defined in Practice E29.

6.3.1 With the absolute method, an observed value is not rounded, but is to be compared directly with the limiting value. Example: In Table 2 Type I, Grade 4, Class B, under Specific Gravity, 2.14 shall be considered as 2.140000 and 2.16 shall be considered 2.160000.

7. Sampling

7.1 Sampling shall be statistically adequate to satisfy the requirements in Section 11.

8. Number of Tests

8.1 Lot inspection shall include tests for bulk density, particle size, and extrusion pressure. Periodic tests shall consist of all the tests specified in Tables 1 and 2 and shall be made at least once per year.

8.2 The tests listed in Tables 1 and 2, as they apply, are sufficient to establish conformity of a material to this specification. One set of test specimens as prescribed in Section 9 shall be considered sufficient for testing each sample. The average of the results for the specimens tested shall conform to the requirements of this specification.

9. Specimen Preparation

9.1 Test Disks for Tensile Properties:

9.1.1 Use the die shown in Fig. 1 for the molding of test disks (see Note 2). Place flat aluminum disks, 0.1 to 0.4 mm (0.004 in. to 0.016 in.) thick and 76 mm (3 in.) in diameter, on both sides of the resin. The test resin shall be near ambient temperature prior to molding (see Note 3). (**Warning**—PTFE resins can evolve small quantities of gaseous products when heated above 204°C (400°F). Some of these gases are harmful. Consequently, exhaust ventilation must be used whenever these resins are heated above this temperature, as they are during the sintering operations that are a part of this specification. Since the temperature of burning tobacco exceeds 204°C (400°F), those working with PTFE resins shall ensure that tobacco is not contaminated.)

NOTE 3—For maximum precision, these weighing and preforming operations shall be carried out at 23 ± 2°C (73.4 ± 3.6°F) (the “near ambient” temperature referred to herein). These operations shall not be performed at temperatures below 21°C (70°F) due to the crystalline transition that occurs in PTFE in this temperature region which leads to possible cracks in sintered specimens and differences in specimen density (as well as changes in other physical properties). Problems caused by the effect of temperature on the specific gravity or density of PTFE shall be minimized when the measurement is made using immersion procedures if a sensitive thermometer (for example, one reading ± 0.1°C) is used in the liquid and the temperature is adjusted to be at least 22°C.

9.1.2 Screen 14.5 g of PTFE resin through a No. 10 sieve into the die. Adjust the lower plug height to allow the resin in the die can be leveled by drawing a straightedge in contact with the top of the die across the top of the die cavity. Insert the die in a suitable hydraulic press and apply pressure gradually (see Note 4) until a pressure of 14 MPa (2030 psi) is attained. Hold this pressure for 3 min. Remove the disk from the die. Write the sample identification number on the preform using an appropriate marker that will not affect the PTFE during sintering.

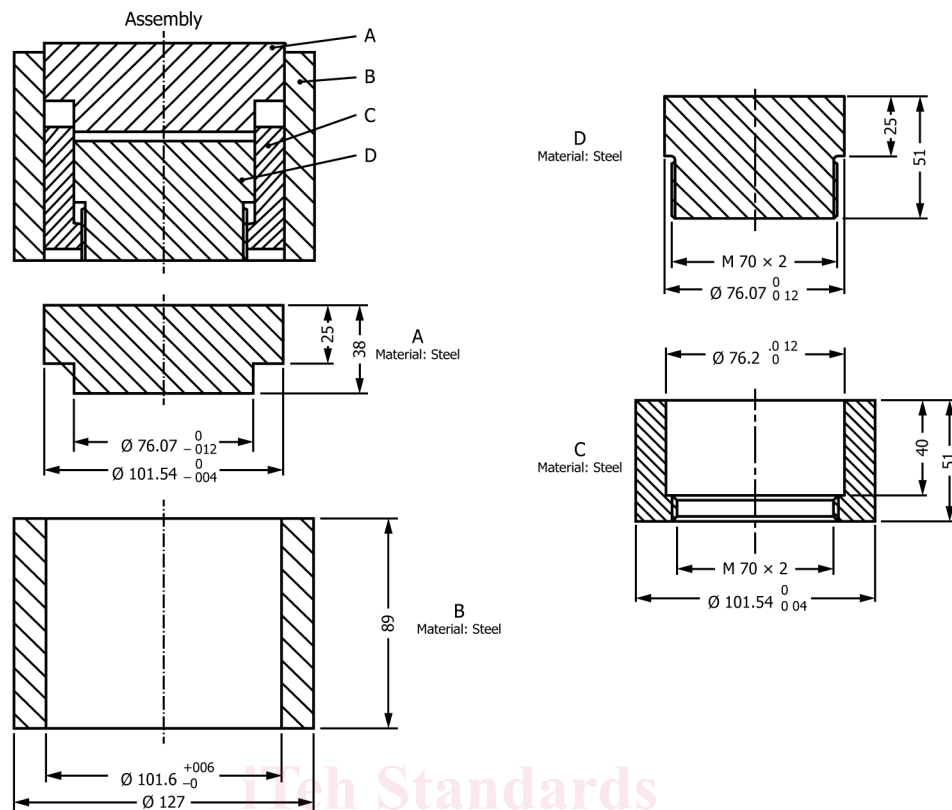
NOTE 4—As a guide, increasing the pressure at a rate of 3.5 MPa (500 psi)/min is suggested until the desired maximum pressure is attained.

9.1.3 Place the sintering oven in a laboratory hood (or equip it with an adequate exhaust system) and sinter the preforms in accordance with Table 3, Procedure A (see Note 5).

NOTE 5—Although the rate of heat application is not critical, the cooling cycle is most important and the conditions cited in this procedure must be followed very closely. If they are not followed, the crystallinity of the disks and the resulting physical properties will be markedly changed. Therefore, the use of a programmed oven is recommended for the most precise sintering cycle control and the hood window shall be left down during the entire sintering procedure, the latter being an important safety consideration.

9.2 Test Specimens for Standard Specific Gravity and Thermal Instability Index:

9.2.1 A cylindrical preforming mold, 29-mm (1.14-in.) internal diameter by at least 76 mm (3 in.) deep, is used to prepare the preforms. Clearance shall be sufficient to ensure escape of air during pressing. Place flat aluminum foil disks, normally 0.13 mm (0.005 in.) thick and 29 mm (1.14 in.) in diameter on both sides of the resin. The test resin shall be near ambient temperature prior to molding (see Note 3).



NOTE 1—All dimensions are in millimetres.

FIG. 1 Mold Assembly for the Preparation of Specimens for the Determination of Tensile Properties

TABLE 3 Sintering Procedures

	A	B
Initial temperature, °C (°F)	290 (554)	290 (554)
Rate of heating, °C/h (°F/h)	120 ± 10 (216 ± 18)	120 ± 10 (216 ± 18)
Hold temperature, °C (°F)	380 ± 6 (716 ± 10)	380 ± 6 (716 ± 10)
Hold time, min	30 + 2, -0	360 ± 5
Cooling rate, °C/h (°F/h)	60 ± 5 (108 ± 9)	60 ± 5 (108 ± 9)
Second hold temperature, °C (°F)	294 ± 6 (561 ± 10)	294 ± 6 (561 ± 10)
Second hold time, min	24 + 0.5, -0	24 + 0.5, -0
Period to room temperature, min	≥30	≥30

9.2.2 Weigh out 12.0 ± 0.1 g of resin and place it in the die. Screen resins through a No. 10 sieve. Compacted resins shall be broken up by hand-shaking cold resin in a half-filled sealed glass container. Condition the resin in the sealed glass container in a freezer or dry-ice chest. After breaking up resin lumps, allow the sealed container to equilibrate to near ambient temperature. Then screen and weigh the 12.0 ± 0.1 -g sample. Insert the die in a suitable hydraulic press and apply pressure gradually (see Note 4) until a pressure of 14 MPa (2030 psi) is attained. Hold this pressure for 2 min. Remove the preform from the die. Write the sample identification number on the preform using an appropriate marker that will not effect the PTFE during sintering.

9.2.3 Sinter the preforms in accordance with Table 3 (see Note 5).

9.2.3.1 For SSG specimens use Procedure A.

9.2.3.2 For ESG specimens use Procedure B.

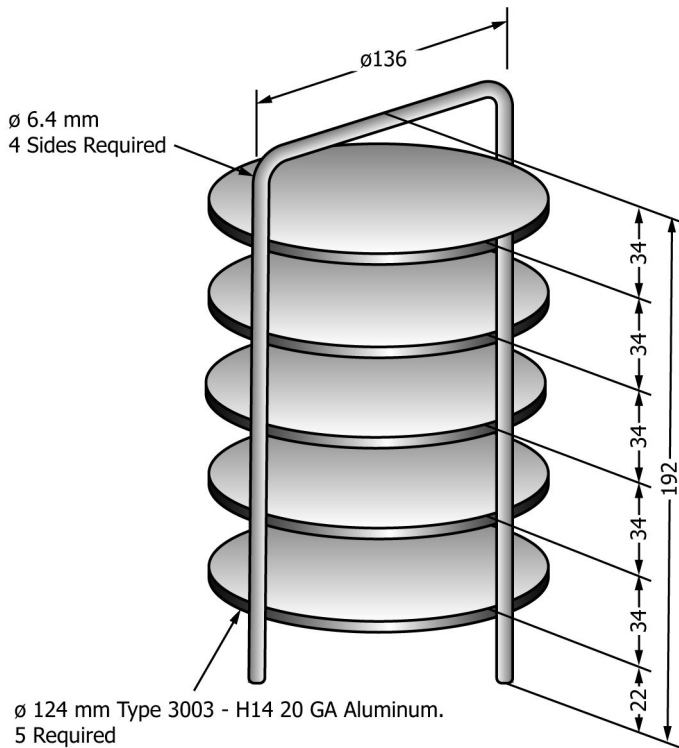
NOTE 6—Improved precision in SSG and ESG test results has been obtained with the use of an upright, cylindrical oven and an aluminum sintering rack. The cylindrical oven has an inside diameter of 140 mm (5.5 in.) and an inside depth of 203 mm (8 in.) plus additional depth to accommodate a 51-mm (2-in.) thick cover, and is equipped with suitable heaters and controllers to sinter specimens in accordance with the procedures in Table 3. The rack, as shown in Fig. 2, allows preforms to be placed symmetrically in the center region of the oven. Place six preforms on each of the middle oven rack shelves (if six or fewer preforms are to be sintered, place them on the middle rack, filling in with “dummies” as needed). Place “dummies” on the top and bottom shelves. Specimens must be spaced evenly in a circle on each shelf, with none of them touching. An oven load must be no less than 18 pieces including “dummies.” “Dummies” are defined as normal 12-g specimens that have previously been through the sintering cycle. “Dummies” must only be used for an additional two or three thermal cycles, due to eventual loss of thermal stability and physical form.

9.2.4 Remove all flash from each specimen so that no air bubbles will cling to the edges when the specimen is immersed in the solution for weighing during the standard specific gravity and thermal instability index tests. It is recommended for this section and during testing that cotton gloves be worn while handling test specimens.

9.3 Test Disks for Stretching Void Index (SVI):

9.3.1 Mold the disk as in 9.1.1.

9.3.2 Screen 29 g of PTFE resin through a 2.00-mm (No. 10) sieve into the die. Adjust the lower plug to allow the resin to be leveled by drawing a straightedge in contact with the top



NOTE 1—Aluminum plates tack welded to rods.
 NOTE 2—All dimensions are in millimetres.

FIG. 2 Sintering Rack for SSG Specimens

of the die across the top of the die cavity. Insert the die in a suitable hydraulic press and apply pressure gradually (see Note 4) until a pressure of 7 MPa (1015 psi) is attained. Hold this pressure for 2 min, then increase the pressure to 14 MPa (2030 psi) and hold for an additional 2 min. Remove the disk from the die. Write the sample identification number on the preform using an appropriate marker that will not effect the PTFE during sintering.

9.3.3 Sinter the preforms in accordance with Table 3, Procedure A (see Note 5).

9.3.4 Remove all flash from those portions of these specimens that will be used for determination of specific gravities so that no air bubbles will cling to their edges when the specimens are immersed in liquid during these tests. It is recommended that cotton gloves be worn while handling test specimens.

9.4 Conditioning Test Specimens:

9.4.1 For tests of tensile properties and all tests requiring the measurement of specific gravities, condition the test specimens in general accordance with Procedure A of Practice D618, with the following deviations therefrom: (1) the aging period shall be a minimum of 4 h immediately prior to testing, (2) the laboratory temperature shall be $23 \pm 2^\circ\text{C}$ ($73.4 \pm 3.6^\circ\text{F}$), and (3) there shall be no requirement respecting humidity. The other tests require no conditioning of the molded test specimens.

9.5 Test Conditions:

9.5.1 Tests shall be conducted at the standard laboratory temperature of $23 \pm 2^\circ\text{C}$ ($73.4 \pm 3.6^\circ\text{F}$), unless otherwise specified in the test methods or in this specification. This

deviation from the standard laboratory temperature is made because of the necessity for maintaining test temperatures above approximately 21°C (70°F). See Note 3 for additional details. Since these resins do not absorb water, the maintenance of constant humidity during testing is not required.

10. Test Methods

10.1 Melting Characteristics by Thermal Analysis:

10.1.1 Significance and Use—For PTFE resins that have been melted prior to use, the melting peak temperature characteristics of a resin provide important information about the thermal history of the material. Melting peak temperatures (see Fig. 3) are used to determine conformance of a resin to the melting peak temperature requirements in Table 1 of this specification.

10.1.2 Apparatus—Use apparatus described in Test Method D4591.

10.1.3 Procedure—Measure melting peak temperatures in accordance with procedures given in Test Method D4591. An initial melting peak temperature above the melting peak temperature obtained on the second and subsequent melting (defined as the second melting peak temperature) indicates that the resin was not melted before the test. The second melting peak temperature occurs at about 327°C (621°F). The difference between the initial and second melting peak temperatures is greater than 5°C (9°F). If peak temperatures are difficult to discern from the curves (that is, because the peaks are rounded rather than pointed) straight lines should be drawn tangent to the sides of the peak. These lines intersect at the peak temperature. Where more than one peak occurs during the initial melting test, the presence of any peak corresponding to the second melting peak temperature indicates the presence of some previously melted material.

10.2 Bulk Density:

10.2.1 Significance and Use—Bulk density gives an indication of how a resin performs during the filling of processing equipment. PTFE resins tend to compact during shipment and storage. Because of this tendency to pack under small amounts of compression or shear, Test Method D1895 is not applicable

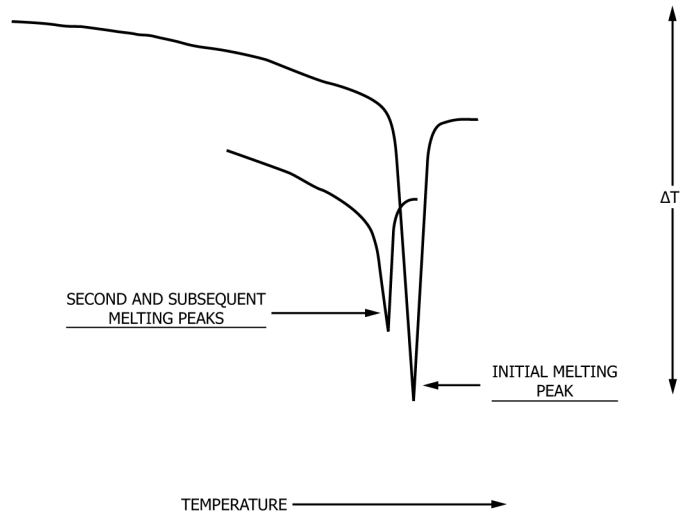


FIG. 3 Melting Characteristics by Thermal Analysis

to these resins. The procedure given in 10.2.2 through 10.2.5 must be used to measure this property.

10.2.2 Apparatus:

10.2.2.1 Funnel—A funnel arrangement as shown in Fig. 4.

10.2.2.2 Feeder⁶—A feeder with a No. 8 wire screen placed over approximately the top two thirds of the trough. The funnel shall be mounted permanently in the feeder outlet.

10.2.2.3 Controller⁷

10.2.2.4 Volumetric Cup and Cup Stand (see Fig. 5)—The volumetric cup shall be calibrated initially to 250 mL by filling it with distilled water, placing a planar glass plate on top, drying the outside of the cup, and weighing. The net weight shall be 250 ± 0.5 g. The top and bottom faces of the volumetric cup and the cup stand shall be machined plane and parallel.

10.2.2.5 Leveling Device—The leveler (see Fig. 6) shall be affixed permanently to the table and adjusted so that the sawtooth edge of the leveler blade passes within 0.8 mm (0.031 in.) of the top of the volumetric cup.

10.2.2.6 Work Surface—The work surface for holding the volumetric cup and leveler shall be essentially free from vibration. The feeder, therefore, must be mounted on an adjoining table or wall bracket.

10.2.2.7 Balance—Balance, having an extended beam, with a capacity of 500 g and a sensitivity of 0.1 g, or equivalent.

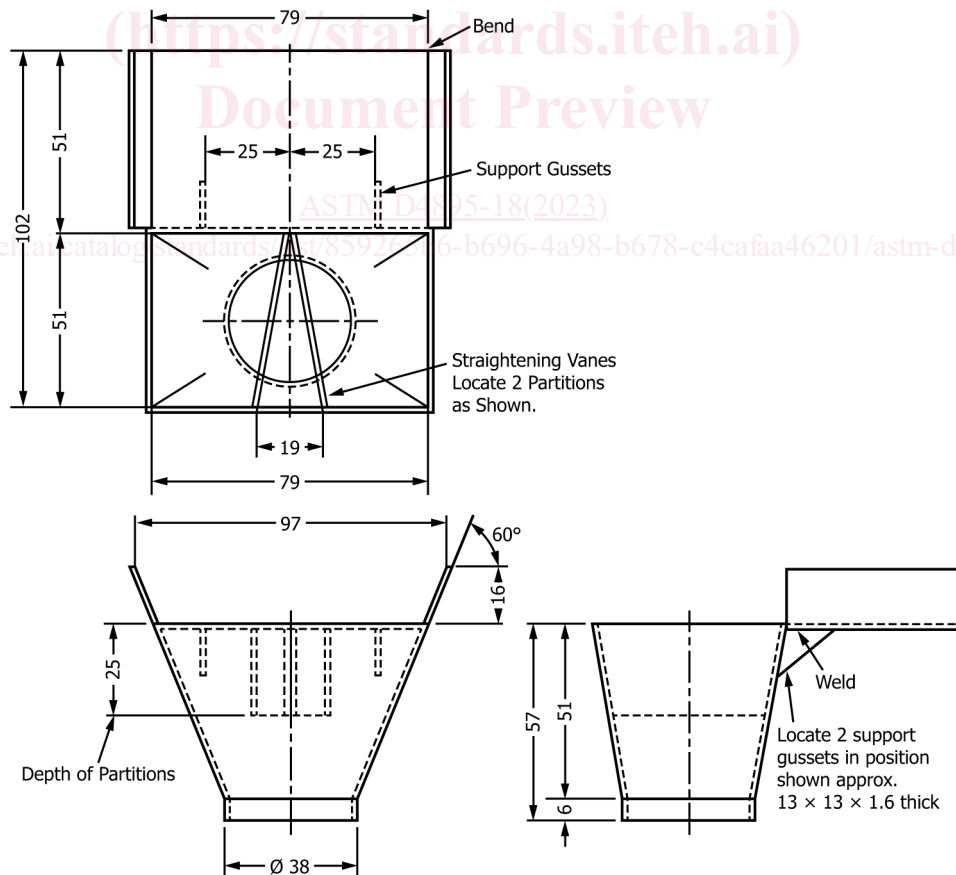
10.2.3 Procedure—Place the clean, dry volumetric cup on the extended beam of the balance and adjust the tare to zero. Select about 500 mL of the resin to be tested and place it on the feeder screen. Put the cup in the cup stand and place the assembly such that the distance of free-polymer fall from the feeder outlet to the top rim of the cup shall be 39 ± 3 mm (1.5 ± 0.012 in.). Increased fall causes packing in the cup and higher bulk density values. Set the controller so that the cup is filled in 20 to 30 s. Pour the sample on the vibrating screen and fill the cup so that the resin forms a mound and overflows. Let the resin settle for about 15 s and then gently push the cup and its stand beneath the leveler. Exercise care to avoid agitation of the resin and cup before leveling. Weigh the resin to the nearest 0.1 g.

10.2.4 Calculation—Calculate the bulk density as follows:

$$\text{grams of resin} \times 4 = \text{bulk density (grams per litre)}$$

⁶ A “Vibra-Flow” Feeder, Type FT01A, available from FMC Corp., Material Handling Division, FMC Building, Homer City, PA 15748, has been found satisfactory for this purpose.

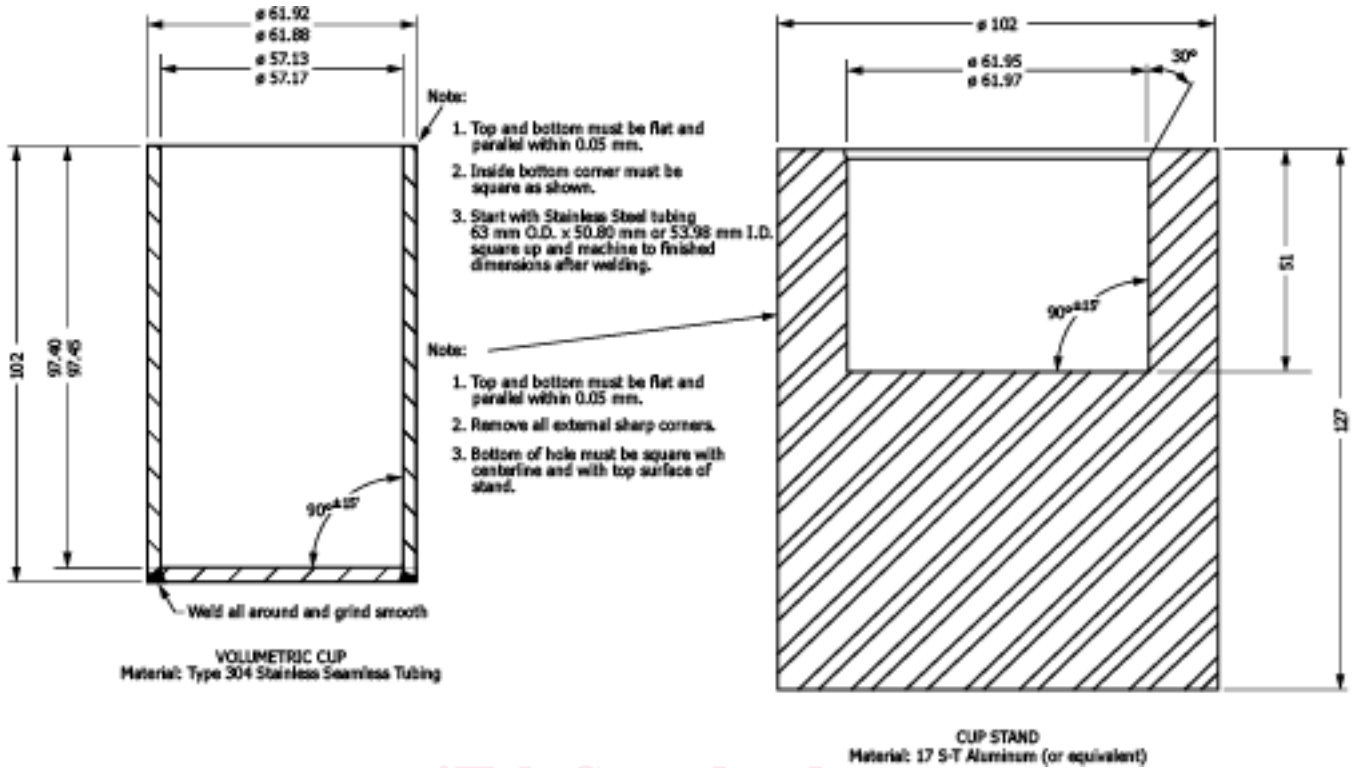
⁷ A “Syntron” controller, Type SCR1B, available from FMC Corp., address as shown in Footnote 10, has been found satisfactory for this purpose.



NOTE 1—Funnel Material: type 304 Stainless Steel 16 Gage (1.6-mm thickness).

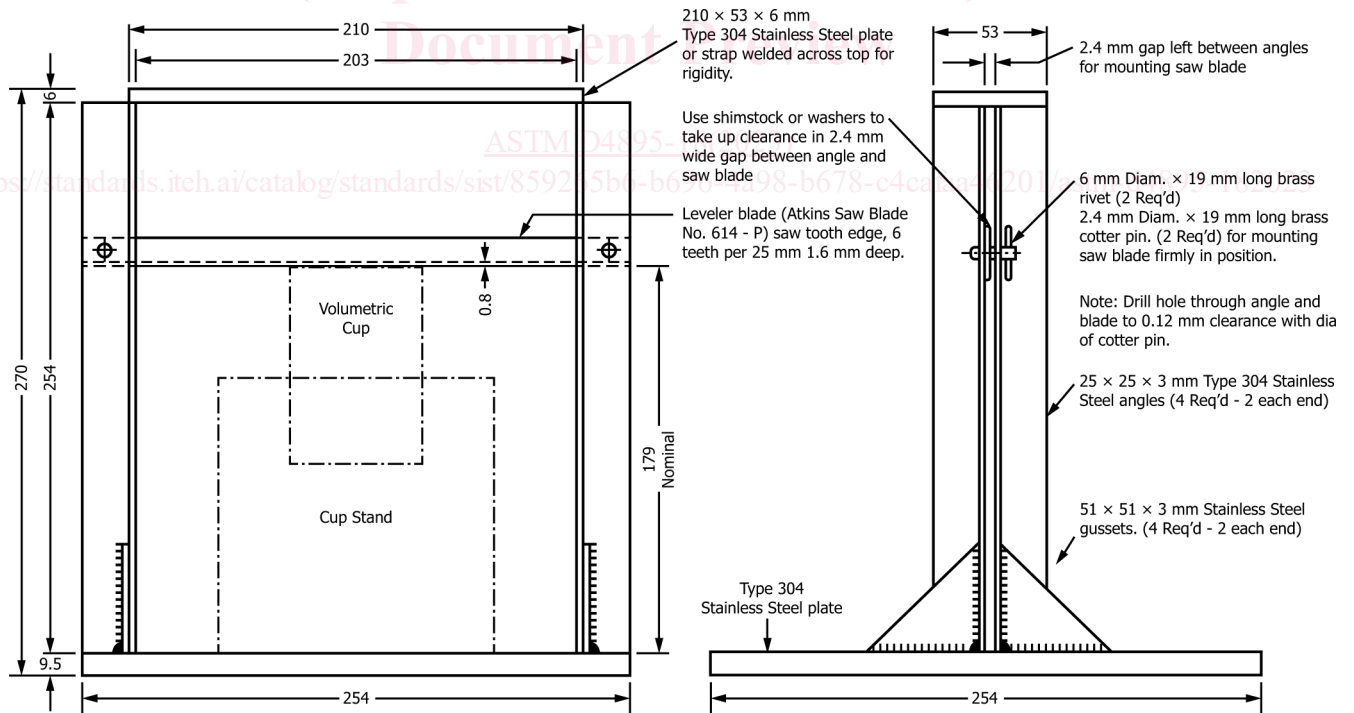
NOTE 2—All dimensions are in millimetres.

FIG. 4 Details of the Funnel Used for the Determination of Bulk Density



NOTE 1—All dimensions are in millimetres.

FIG. 5 Volumetric Cup and Cup Stand for the Determination of Bulk Density



NOTE 1—Base plate must be flat and parallel. Saw blade, when mounted, must be square to and parallel with base plate within 0.13 mm from end to end. Height of saw blade must have 0.8 mm or less clearance between blade and assembled cup and cup stand (as indicated by phantom lines). Welded construction where indicated. Material: as noted.

NOTE 2—All dimensions are in millimetres.

FIG. 6 Leveler Stand for the Determination of Bulk Density