



# Standard Test Methods for Solid Filling and Treating Compounds Used for Electrical Insulation<sup>1</sup>

This standard is issued under the fixed designation D 176; 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 These test methods cover physical and electrical tests for solid filling and treating compounds used for electrical insulation which are fusible to a liquid without significant chemical reaction. Compounds that are converted to the solid state by polymerization, condensation, or other chemical reaction are not included in these test methods.

1.2 These test methods are designed primarily for asphaltic or bituminous compounds, waxes, and fusible resins, or mixtures thereof, although some of these methods are applicable to semisolid types such as petrolatums. Special methods more suitable for hydrocarbon waxes are contained in Test Methods D 1168.

1.3 Adequate ventilation must be provided when these tests involve heating.

1.4 The test methods appear in the following sections:

Test Method	Sections
<i>Electrical Tests:</i>	
A-C Loss Characteristics and Permittivity (Dielectric Constant)	51-54
Dielectric Strength	42-45
Volume Resistivity-Temperature Characteristics	46-49
<i>Physical Tests:</i>	
Coefficient of Expansion or Contraction	22-41
Flash and Fire Points	9 and 10
Loss on Heating	11 and 12
Melting Point	5 and 6
Penetration	15 and 16
Softening Point	7 and 8
Specific Gravity	17-21
Viscosity	13 and 14

1.5 The values stated in SI units are to be regarded as the standard. The values given in parentheses are for information only.

1.6 *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.* For specific hazard statements, see **Note 4** and **Note 7**.

NOTE 1—There is no similar or equivalent IEC or ISO standard.

## 2. Referenced Documents

2.1 *ASTM Standards:*

- D 5 Test Method for Penetration of Bituminous Materials<sup>2</sup>
- D 6 Test Method for Loss on Heating of Oil and Asphaltic Compounds<sup>3</sup>
- D 70 Test Method for Specific Gravity of Semi-Solid Bituminous Materials<sup>2</sup>
- D 71 Test Method for Relative Density of Solid Pitch and Asphalt (Displacement Method)<sup>4</sup>
- D 88 Test Method for Saybolt Viscosity<sup>3</sup>
- D 92 Test Method for Flash and Fire Points by Cleveland Open Cup<sup>4</sup>
- D 127 Test Method for Drop Melting Point of Petroleum Wax, Including Petrolatum<sup>4</sup>
- D 149 Test Method for Dielectric Breakdown Voltage and Dielectric Strength of Solid Electrical Insulating Materials at Commercial Power Frequencies<sup>5</sup>
- D 150 Test Methods for AC Loss Characteristics and Permittivity (Dielectric Constant) of Solid Electrical Insulating Materials<sup>5</sup>
- D 257 Test Methods for D-C Resistance or Conductance of Insulating Materials<sup>5</sup>
- D 937 Test Method for Cone Penetration of Petrolatum<sup>4</sup>
- D 1168 Test Methods for Hydrocarbon Waxes Used for Electrical Insulation<sup>5</sup>
- D 1711 Terminology Relating to Electrical Insulation<sup>5</sup>
- E 28 Test Method for Softening Point by Ring-and-Ball Apparatus<sup>6</sup>
- E 102 Test Method for Saybolt Furol Viscosity of Bituminous Materials at High Temperatures<sup>3</sup>

## 3. Terminology

3.1 *Definitions:*

3.1.1 *dielectric strength, n*—the voltage gradient at which dielectric failure of the insulating material occurs under specific conditions of test.

<sup>1</sup> These methods of testing are under the jurisdiction of ASTM Committee D09 on Electrical and Electronic Insulating Materials and are the direct responsibility of Subcommittee D09.01 on Electrical Insulating Varnishes, Powders, and Encapsulating Compounds.

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<sup>2</sup> *Annual Book of ASTM Standards*, Vol 04.03.

<sup>3</sup> *Annual Book of ASTM Standards*, Vol 04.04.

<sup>4</sup> *Annual Book of ASTM Standards*, Vol 05.01.

<sup>5</sup> *Annual Book of ASTM Standards*, Vol 10.01.

<sup>6</sup> *Annual Book of ASTM Standards*, Vol 06.03.

3.1.2 For definitions of other terms relating to electrical insulation see Terminology **D 1711**.

3.2 *Definitions of Terms Specific to This Standard:*

3.2.1 *loss on heating, n— of filling or treating compound*, the change in weight of a compound when heated under prescribed conditions at a standard temperature for a specified time.

3.2.2 *melting point, n— of filling or treating compound*, the temperature at which the compound becomes sufficiently fluid to drop from the thermometer used in making the determination under prescribed conditions.

3.2.3 *penetration, n— of filling or treating compound*, the distance traveled by a standard needle (or cone) as it pierces a specimen under specified conditions of load, time and temperature.

3.2.4 *softening point, n— of filling or treating compound*, the temperature at which the central portion of a disk of the compound held within a horizontal ring of specified dimensions has sagged or flowed downward a distance of 25 mm (1 in.) under the weight of a 10-mm (3/8-in.) diameter steel ball as the sample is heated at a prescribed rate in a water or glycerin bath.

**4. Sampling and Conditioning**

4.1 Due to the diverse nature of the compounds and the various forms and packages commercially available, no standard methods of sampling have been established. When the

sample is in the form of cakes or ingots, a representative sample may usually be secured by breaking or cutting a transverse section from the middle of the cake or ingot. When the material is shipped in pails or drums, a sample may be removed with a clean knife, hatchet, auger or other cutting tool, discarding the top 50 or 75 mm (2 or 3 in.) of the compound. Melting of the compound should be avoided unless it can be poured directly into the testing container. A melting and pouring temperature of 50°C (90°F) above the softening point is recommended for filling testing containers with asphaltic compounds. Take care not to overheat the compound nor to entrap air.

4.2 With certain materials that tend to entrap gasses due to high viscosity at pouring temperatures, or to froth on heating, it may be necessary to degas the material prior to testing in order that consistent results may be secured (unless the particular test includes such procedure). If degassing is required, it shall be performed by heating the material in a vacuum oven. The temperature and vacuum shall be high enough, and the time long enough to ensure driving off the mechanically entrapped gasses, but should tend to decompose as little as possible. A temperature 50°C (90°F) higher than the softening point of the compound, an absolute pressure of 7 to 21 kPa (1 to 3 psi), and a time of 30 to 45 min are recommended for asphaltic compounds. The sample shall then be poured into the testing container.

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**PHYSICAL TESTS**

**MELTING POINT**

**5. Significance and Use**

5.1 The melting point is useful in selecting a filling or treating compound that will not flow at the operating temperature of the device in which it will be used. It is also essential that it shall not be so high as to injure the insulation at the time of pouring. This test method may be used for specification, classification, and for control of product uniformity.

**6. Procedure**

6.1 Determine the melting point of petrolatums, waxes, and similar compounds of a relatively sharp melting point by Test Method **D 127**.

NOTE 2—This method should not be used for asphalts and other types with a prolonged melting range.

**SOFTENING POINT**

**7. Significance and Use**

7.1 The softening point is useful in selecting a filling or treating compound that will not flow at the operating temperature of the device in which it is used. It is also an indication of the pouring temperature, which should not be so high as to injure the insulation of a device. This test method is used, when

the compound has no definite melting point, for purposes of specification, classification, and control of product uniformity.

**8. Procedure**

8.1 Determine the softening point in accordance with Test Method **E 28**.

**FLASH AND FIRE POINTS**

**9. Significance and Use**

9.1 The flash and fire points must be high enough so that the possibility of an explosion or fire is at a minimum when the compounds are being heated and poured. A flash point at least 35°C (63°F) above the pouring temperature is usually considered necessary for safe operations. An unusually low flash point for a given compound indicates a mixture or contamination with a volatile material. This test method is useful for purposes of specification, classification, and control of product uniformity.

**10. Procedure**

10.1 Determine the flash and fire points of all compounds in accordance with Test Method **D 92**.

10.2 In the case of certain compounds containing chlorine, the flash may be indefinite and no fire point may exist. Report this fact.

## LOSS ON HEATING

### 11. Significance and Use

11.1 Loss on heating includes loss of moisture and volatile constituents less any weight gain due to oxidization. It is useful for control of product uniformity and as an indication of pot or tank life if the test is performed at the appropriate temperature. This test method shall not be used to compare compounds of different basic chemical compositions.

### 12. Procedure

12.1 Determine the loss on heating of asphaltic and certain other types of compounds by Test Method **D 6**.

NOTE 3—The reproducibility of this test method may be poor due to insufficient control of the air circulation over the specimens and to weight gain from oxidation of some compounds. With certain compounds it may be desirable to conduct the test at a lower temperature than the specified temperature of 163°C (325°F).

NOTE 4—**Caution:** When compounds of low flash point and high volatility are tested, the oven shall have low-temperature heating elements and a safety door latch to relieve pressure in case of an explosion.

## VISCOSITY

### 13. Significance and Use

13.1 The Saybolt viscosity is nearly proportional to the kinematic viscosity of filling and treating compounds and hence, it is an indication of whether or not the material will flow readily under its own weight at a prescribed temperature. It is also satisfactory for control of product uniformity and for specification purposes.

### 14. Procedure

14.1 For waxes, petrolatums, and other low-viscosity-type compounds determine the viscosity as Saybolt Universal viscosity by Test Method **D 88**. The standard temperatures for testing are: 21, 38, 54, or 99°C (70, 100, 130, or 210°F).

14.2 For asphaltic and other high-viscosity compounds, determine the Saybolt Furol viscosity. The standard temperatures for testing Furol viscosity are: 25, 38, 50, 60, 82, and 99°C (77, 100, 122, 140, 180, 210°F).

14.3 For higher temperatures, special techniques and thermometers are required. The standard temperatures are 121, 149, 177, 204, and 232°C (250, 300, 350, 400, 450°F). In these cases determine the viscosity by Test Method **E 102**.

NOTE 5—For testing waxes and petrolatums, the standard temperature for comparison purposes is 99°C (210°F), and Saybolt Universal viscosity is used. For estimation of the properties of asphaltic and other compounds of high viscosity, it is desirable to measure the viscosity at a number of standard temperatures above the softening point. A curve may be plotted on log-log paper and the temperature at which the Saybolt Furol viscosity is 470 s may be determined. This viscosity corresponds approximately to a kinematic viscosity of 1000 centistokes, and is a viscosity at which the compound may conveniently be poured from the container. With potting compounds, it is also desirable to know the temperature at which the Saybolt Furol viscosity is 100 s, since this viscosity is low enough for production potting operations.

## PENETRATION

### 15. Significance and Use

15.1 Penetration is an indication of the softness or indentability of a compound. Penetration values are used as a basis for classification, specification, and control of product uniformity.

### 16. Procedure

16.1 Determine penetration in accordance with Test Method **D 5**. This test method is applicable to all compounds except very soft materials and petrolatums. Unless specified otherwise, the standard conditions of test are:

	Weight, g	Time, s
At 25°C (77°F)	100	5

Other standard conditions are:

	Weight, g	Time, s
At 0°C (32°F)	200	60
At 46°C (115°F)	50	5

16.2 For very soft materials, such as petrolatums, use Test Method **D 937**.

## SPECIFIC GRAVITY

### 17. Significance and Use

17.1 Specific gravity is useful for indicating product uniformity and for calculating the weight of a given volume of material. In some instances it is useful in estimating the amount of mineral fillers in a compound. If specific gravity is known at several temperatures, the coefficient of expansion may be calculated. If the specific gravity of a compound is determined before and after degassing, the volume of entrapped gases may be calculated.

17.2 Displacement tests are used to determine the specific gravity of both untreated and degassed compounds. Conventional methods are used for the solid state, and plummet displacement for the liquid state. The values obtained may then be used to compute the approximate coefficient of cubical expansion by Test Method **C** (see Sections **34-36**).

## WATER DISPLACEMENT METHODS

### 18. Procedure

18.1 Determine the specific gravity by Test Method **D 71** or Test Method **D 70**.

## PLUMMET DISPLACEMENT METHOD

### 19. Scope

19.1 The specific gravity of the material at the desired temperature is calculated from the weight of the compound displaced by a calibrated aluminum plummet.

### 20. Apparatus

20.1 *Balance*—An analytical balance equipped with pan straddle.

20.2 *Plummet*—An aluminum plummet of suitable shape weighing 5 to 10 g.

20.3 *Beaker*—A 400-mL heat-resistant glass beaker wrapped with a suitable thermal insulation.

20.4 *Thermometer*—A thermometer of suitable range.

20.5 *Wire*—Two pieces of fine copper wire.

## 21. Procedure

21.1 *Calibration of Plummet*—Make the following weight determinations of the plummet to the nearest 1 mg as follows:

$$a - b = \text{weight of water displacement in grams at } 25^{\circ}\text{C (77}^{\circ}\text{F)} \quad (1)$$

where:

$a$  = weight in air, g, and

$b$  = weight suspended in water, g, at  $25^{\circ}\text{C (77}^{\circ}\text{F)}$ .

21.2 Correct the value of the plummet displacement ( $D_p$ ) in terms of grams of water at  $25^{\circ}\text{C (77}^{\circ}\text{F)}$  to the pouring temperature,  $t_p$ , in degrees Celsius, by means of the following equation

$$D_p = 0.000076(t_p - 25)(a - b) + (a - b) \quad (2)$$

NOTE 6—The factor 0.000076 is the coefficient of cubical expansion per degree Celsius.

21.3 *Testing of the Sample*—Carefully melt the sample in the beaker and raise the temperature to approximately  $15^{\circ}\text{C (27}^{\circ}\text{F)}$  above the desired test temperature. Place the beaker on the straddle and suspend the plummet in the compound by the fine copper wire. (The weight of the wire should be tared.)

21.4 Balance the scales approximately and at the same time stir the sample slowly, using the thermometer as a stirring rod. When the sample has cooled to the desired temperature, rapidly complete the weighing.

21.5 *Calculation of Specific Gravity,  $t_p/25^{\circ}\text{C}$* —Calculate the specific gravity as follows:

$$\text{Sp gr, } t_p/25^{\circ}\text{C} = (W_a - W_c)/D_p \quad (3)$$

where:

$W_a$  = weight of plummet in air, g, and

$W_c$  = weight of plummet in compound, g.

## COEFFICIENT OF EXPANSION OR CONTRACTION

### 22. Scope

22.1 The following four test methods are included:

22.1.1 *Test Methods A and B*—Methods A and B for true coefficient of expansion are intended for use only where the uniformity of the material under test justifies a high degree of precision. Test Method A is suitable for testing low-viscosity types such as waxes and petrolatums. Test Method B is suitable for testing asphalts and high-viscosity materials, also for opaque materials that may give difficulty in reading the glass scale of Test Method A.

22.1.2 *Test Methods C and E*—Test Methods C and E are intended for faster testing where high precision may not be justified. These test methods may be used for determining either true or effective coefficient of expansion but are not used as referee test methods.

### 23. Significance and Use

23.1 Coefficient of expansion is useful in computing the amount of void space that will remain in a device filled with compound after the compound has cooled to the ambient temperature. It also is one indication of the thermal shock resistance of a compound.

23.2 The effective coefficient of expansion is determined on materials that have not been degassed just prior to test. It is important for many purposes to know the effective coefficient of the material as received or after heating to the maximum temperature of application. Consistent results, however, may only be obtained with gas-free compounds.

## TEST METHOD A—USING GLASS FLASK

### 24. Apparatus

24.1 *Flask*—A glass flask<sup>7</sup> holding approximately 250 mL to the zero mark, and graduated for 25 mL in 0.1-mL divisions, the neck of the flask being 10 mm in internal diameter.

24.2 *Oil Bath*—For heating the sample, a cylindrical oil bath approximately 25.4 cm (10-in.) inside diameter and 50 cm (20 in.) in inside depth with a false bottom 2.5 cm (1 in.) from the bottom and provision for circulating and heating the oil.

24.3 *Metal Collar*—Lead or iron collars for use on the neck of the flask during test to prevent oil currents of the bath from moving the flask.

### 25. Calibration

25.1 The capacity of the flask at the zero point and several points on the scale, shall be determined by filling the flask with distilled water at a known temperature and weighing.

### 26. Procedure

26.1 Maintain the flask under a vacuum of 640 mm (25 in.) Hg at a temperature  $50^{\circ}\text{C (90}^{\circ}\text{F)}$  higher than the softening point (ring and ball method, as determined in accordance with Section 8) while filling, and for approximately 30 min after filling is completed. Fill flask to within the last millilitre marked on the neck when held at the maximum test temperature and slowly cooled to room temperature (10 to 12 h). Before starting the test, examine the flask for the presence of cavities or irregular contraction of the compound. Some compounds, after cooling below the liquid state, tend to stick to the sides of the neck of the flask. In such cases, it is necessary to gradually warm the neck and flow the compound to meet the rest, after which the flask shall be placed in the bath for several hours to ensure temperature equilibrium.

26.2 With the compound satisfactorily placed in the flask at the lowest temperature, read the height of the column in the neck and then slowly heat the bath. Take readings at  $5^{\circ}\text{C (9}^{\circ}\text{F)}$  intervals, holding the bath as constant as possible at each point until no more expansion occurs at that point. Repeat the procedure for each point until maximum temperature is reached.

<sup>7</sup> A Pyrex or quartz glass flask is satisfactory for this purpose.

26.3 *Precautions*—During the test, take temperature readings at top and bottom of the bath to detect any variation. Make readings of the expansion of the compound at intervals long enough to ensure uniform temperature distribution and complete movement of the compound. Until complete liquefaction, the interval shall be 3 to 4 h; after liquefaction, it may be reduced to 30 min.

## 27. Calculation

27.1 After securing the readings over the temperature range desired, plot a curve from the temperature and expansion readings from which the coefficient of expansion shall be calculated, as follows:

$$E = [(V_1 - V)/(T_1 - T)V] + C \quad (4)$$

where:

- $E$  = coefficient of expansion ( $1/T$ ) of the compound,
- $V$  = original volume occupied by the compound,  $L$ ,
- $V_1$  = volume at higher temperature occupied by the compound,  $L$ ,
- $T$  = original temperature,
- $T_1$  = higher temperature, and
- $C$  = coefficient of cubical expansion of the glass container. This may be taken as three times the linear coefficient of expansion.

27.2 The coefficient of expansion shall be calculated for three temperature ranges, as follows:

27.2.1 From the minimum temperature at which the measurement was made to 10°C (18°F) below the softening point. This is intended to give the average coefficient for the solid condition.

27.2.2 From 5°C (9°F) above the softening point to 50°C (90°F) above the softening point. This is intended to give the average coefficient for the liquid condition.

27.2.3 From the minimum temperature at which a measurement was made to 50°C (90°F) above the softening point.

## 28. Report

28.1 Report the following information:

28.2 Type of cell used, copy of the volume-temperature curve, temperature ranges as defined in 27.2, and

28.3 Coefficient of expansion corresponding to each of the three temperature ranges.

### TEST METHOD B—USING METALLIC CELL

## 29. Apparatus

29.1 *Metal Cell*—A cell made of steel, consisting of four parts: a cylinder about 64 mm (2.5 in.) in internal diameter having a rigid bottom, a metallic gasket, and a cover to which a steel capillary tube is attached. The cell shall have an internal volume of approximately 250 mL. A metallic cell that has been found suitable is described in Annex A1.

29.2 *Oil Bath*—An oil bath as described in 24.2, Test Method A, with the exception that provision shall be made for supporting the metal cell.

## 30. Calibration

30.1 The cell shall be calibrated to determine its volume at various temperatures as follows:

30.1.1 Weigh the assembled cell to determine its tare weight.

30.1.2 Fill the cell with mercury until replacing the cover causes some to extrude through the capillary tubing. Record the weight of the cell and the mercury and note the temperature.

30.1.3 Place the cell in the oil bath in an inverted position. The capillary tubing should extend over the side of the oil bath in such a way that the extruded mercury may be caught in a beaker. The oil bath, which is several degrees above room temperature, will cause some of the mercury to be extruded from the capillary tube. When all expansion has taken place, weigh the mercury collected.

30.1.4 Adjust the oil bath for other test temperatures and note the amounts of mercury extruded. The weight of the mercury in the cell at any temperature is thus determined, and the volume may be calculated.

## 31. Procedure

31.1 While filling the cell, place it in an oil bath and maintain at a temperature 50°C (90°F) higher than the softening point of the compound (ring and ball method, as determined in accordance with Section 8). When the cell has been filled to within 6 mm (¼ in.) of the cover, place it in a vacuum oven and maintain at a vacuum of 640 mm (25 in.) Hg and a temperature 50°C (90°F) higher than the softening point of the compound for a period of not less than 30 min nor more than 45 min. At the end of this period slowly cool the cell to room temperature, and remove any irregularities in the surface of the compound.

31.2 Screw on the cover and re-weigh the cell and compound.

31.3 Pour sufficient mercury into the cell so that some is extruded when the cover is screwed down. Then weigh the cell again.

31.4 Invert the cell and place in the oil bath, and repeat the procedure prescribed in 30.1.3 and 30.1.4 for 5°C (9°F) intervals.

31.5 *Precautions*—Only clean, distilled mercury shall be used. During the test, take temperature readings at top and bottom of the bath to detect any variation. Readings of the expansion of the compound should be made at time intervals long enough to ensure uniform temperature distribution and complete movement of the compound. Until complete liquefaction of the compound the interval should be 3 to 4 h; after liquefaction, it may be reduced to 30 min.

NOTE 7—**Caution:** Mercury metal vapor poisoning has long been recognized as a hazard in industry. The maximum exposure limits are set by the American Conference of Governmental Industrial Hygienists.<sup>8</sup> The concentration of mercury vapor resulting from use of the above procedure can easily exceed these exposure limits. Mercury, being a liquid and quite heavy, will disintegrate into small droplets and seep into cracks and crevices in the floor if it is spilled. The increased area of exposure adds significantly to the mercury vapor concentration in the air. Mercury vapor concentration is easily monitored using commercially available sniffers. Spot checks shall be made periodically around operations where mercury

<sup>8</sup> American Conference of Governmental Hygienists, Building D-7, 6500 Glenway Drive, Cincinnati, OH 45211.

is exposed to the atmosphere. Thorough checks shall be made after spills. Emergency spill kits are available should the airborne concentration exceed the exposure limits. In addition, exercise care to keep the mercury from the hands. The use of rubber gloves is recommended for handling specimens in the above manner.

### 32. Calculation

32.1 After volumetric determinations have been made over the desired temperature range, plot a curve between volume and temperature readings from which the coefficient of expansion shall be calculated, as follows:

$$E = (V_1 - V)/(t_1 - T)V \quad (5)$$

where:

- $E$  = coefficient of expansion ( $1/T$ ) of the compound,
- $V$  = original volume occupied by the compound,  $L$ ,
- $V_1$  = volume at higher temperature occupied by the compound,  $L$ ,
- $T$  = original temperature, and
- $T_1$  = higher temperature.

32.2 Calculate the coefficient of expansion for the same three ranges as prescribed in Test Method A.

### 33. Report

33.1 Report the following information:

- 33.1.1 Type of cell used,
- 33.1.2 Copy of the volume-temperature curve,
- 33.1.3 Temperature ranges as defined in 27.2, and
- 33.1.4 Coefficient of expansion corresponding to each of the three temperature ranges.

## TEST METHOD C—SPECIFIC GRAVITY METHOD

### 34. Procedure

34.1 Determine the specific gravity of untreated or degassed compounds at two test temperatures by one or more of the procedures specified in Sections 17-21 applying to the state of the materials at the temperatures between which measurements are desired.

NOTE 8—When the temperature range includes the range over which the material changes from solid to liquid, a true coefficient of expansion cannot be calculated, although for practical purposes this may be done.

### 35. Calculation

35.1 From the temperature and specific gravity readings, calculate the coefficient of expansion as follows:

$$E = \text{sp gr at } T - \text{sp gr at } T_1 / (T_1 - T) \text{ sp gr at } T_1 \quad (6)$$

where:

- $E$  = coefficient of expansion ( $1/T$ ) of the compound,
- $T$  = initial temperature, and
- $T_1$  = higher temperature.

### 36. Report

36.1 Report the following information:

- 36.1.1 Method used,
- 36.1.2 Temperature ranges used, and
- 36.1.3 Coefficient of expansion over temperature ranges used.

## TEST METHOD D PYCNOMETER EXPANSION

### 37. Scope

37.1 This test method is another modification of the specific gravity method (Test Method C) and may also be applied to either untreated or degassed materials. This test method is applicable up to temperatures at which the extruded compound flows down the side of the flask and cannot be removed with sufficient precision for weighing.

### 38. Apparatus

38.1 *Flask and Pycnometer*—A 100-mL volumetric heat-resistant glass flask having the zero mark as near as possible to the bulb of the flask and having the neck of the flask cut off at the 100-mL point and ground square. A metal pycnometer may be used provided its coefficient of expansion is known and is applied in the calculation (Section 40).

38.2 *Oil Bath*—An oil bath which may consist of a tall-form heat-resistant glass beaker of sufficient size so that when the flask is supported about 1 in. from the bottom the oil level will reach at least to the zero mark of the flask.

38.3 *Metal Collar*—Lead or iron collars for use on the neck of the flask during heating to prevent oil currents of the bath from moving the flask.

### 39. Procedure

39.1 Allow the pycnometer to cool slowly to the lowest test temperature. During the cooling period keep the flask filled by adding more compound, and after equilibrium is reached, remove the excess material by passing a sharp, flat blade over the rim. Remove the flask from the bath and quickly weigh. Knowing the tare weight and volume of the flask, the specific gravity may be determined. For successively higher temperatures, it is only necessary to weigh the extruded portion. It is recommended that the extruded compound be cut off by tared single-edge razor blades which can be transferred directly to the balance pan. About 1½ h will generally be required to establish temperature equilibrium.

### 40. Calculation

40.1 From the temperature and weight readings calculate the coefficient of expansion as follows:

$$E = [(W - W_1)/W_1(T_1 - T)] - (WC/W_1) \quad (7)$$

where:

- $E$  = coefficient of expansion ( $1/T$ ),
- $W$  = initial weight of the compound in the flask, g,
- $W_1$  = weight of the compound in the flask at higher temperature, g,
- $T$  = initial temperature,
- $T_1$  = higher temperature, and
- $C$  = coefficient of cubical expansion of the flask.

### 41. Report

41.1 Report the following information:

- 41.1.1 Method used,
- 41.1.2 Temperature ranges used, and
- 41.1.3 Coefficient of expansion over temperature ranges used.