ภาคผนวก
มาตรฐาน ASTM D 792
Standard Test Methods for Density and Specific Gravity (Relative Density) of Plastics by Displacement

This standard is issued under the fixed designation D 792; 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 Department of Defense.

1. Scope

1.1 These test methods describe the determination of the specific gravity (relative density) and density of solid plastics in forms such as sheets, rods, tubes, or molded items.

1.2 Two test methods are described:

1.2.1 Test Method A—For testing solid plastics in water,

1.2.2 Test Method B—For testing solid plastics in liquids other than water.

1.3 The values stated in SI units are to be regarded as the 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.

Note 1—This standard is not equivalent to ISO 1183-1 Method A. This test method provides more guidelines on sample weight and dimension. ISO 1183-1 allows testing at an additional temperature of 27 ± 2°C.

2. Referenced Documents

2.1 ASTM Standards: 2

D 618 Practice for Conditioning Plastics for Testing
D 891 Test Methods for Specific Gravity, Apparent, of Liquid Industrial Chemicals
D 4968 Guide for Annual Review of Test Methods and Specifications for Plastics
D 6436 Guide for Reporting Properties for Plastics and Thermoplastic Elastomers

1 These test methods are under the jurisdiction of ASTM Committee D20 on Plastics and are the direct responsibility of Subcommittee D20.70 on Analytical Methods (Section D20.70.01). Current edition approved June 15, 2008. Published July 2008. Originally approved in 1944. Last previous edition approved in 2000 as D 792-00.

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.

E 1 Specification for ASTM Liquid-in-Glass Thermometers
E 12 Terminology Relating to Density and Specific Gravity of Solids, Liquids, and Gases
E 691 Practice for Conducting an Interlaboratory Study to Determine the Precision of a Test Method
IEEE/ASTM SI-10 Practice for Use of the International System of Units (SI) (the Modernized Metric System)

3. Terminology

3.1 General—The units, symbols, and abbreviations used in these test methods are in accordance with IEEE/ASTM SI-10.

3.2 Definitions:

3.2.1 specific gravity (relative density)—the ratio of the mass of a given volume of the impermeable portion of the material at 23°C to the mass of an equal volume of gas-free distilled or de-mineralized water at the same temperature; the form of expression shall be:

Specific gravity (relative density) 23/23°C
(or sp gr 23/23°C)

Note 2—This definition is essentially equivalent to the definition for apparent specific gravity and apparent density in Terminology E 12, because the small percentage difference introduced by not correcting for the buoyancy of air is insignificant for most purposes.

3.2.2 density—cubic metre of impermeable portion of the material at 23°C. The form of expression shall be:

D 313. kg/m³ (Notes 2-4)

Note 3—The SI unit of density, as defined in IEEE/ASTM SI-10, is kg/m³. To convert density in g/cm³ to density in kg/m³, multiply by 1000.

Note 4—To convert specific gravity 23/23°C to density 23°C, kg/m³, use the following equation:

\[ D_{313} \cdot 997.5 \]

Where 997.5 kg/m³ is the density of water at 23°C.

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

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4. Summary of Test Method

4.1 Determine the mass of a specimen of the solid plastic in air. It is then immersed in a liquid, its apparent mass upon immersion is determined, and its specific gravity (relative density) calculated.

5. Significance and Use

5.1 The specific gravity or density of a solid is a property that is conveniently measured to identify a material, to follow physical changes in a sample, to indicate degree of uniformity among different sampling units or specimens, or to indicate the average density of a large item.

5.2 Changes in density of a single material are due to localized differences in crystallinity, loss of plasticizer, absorption of solvent, or to other causes. It is possible that portions of a sample differ in density because of their differences in crystallinity, thermal history, porosity, and composition (types or proportions of resin, plasticizer, pigment, or filler).

5.3 Density is useful for calculating strength-weight and cost-weight ratios.

6. Sampling

6.1 The sampling units used for the determination of specific gravity (relative density) shall be representative of the quantity of product for which the data are required.

6.1.1 If it is known or suspected that the sample consists of two or more layers or sections having different specific gravities, either complete finished parts or complete cross sections of the parts or shapes shall be used as the specimens, or separate specimens shall be taken and tested from each layer. The specific gravity (relative density) of the total part shall not be obtained by adding the specific gravity of the layers, unless relative percentages of the layers are taken into account.

7. Conditioning

7.1 Conditioning—Condition the test specimens at 23 ± 2°C and 50 ± 5% relative humidity for not less than 40 h prior to test in accordance with Procedure A of Practice D 618, unless otherwise specified by the contract or relevant material specifications. In cases of disagreement, the tolerances shall be ±1°C and ±2% relative humidity.

7.2 Test Conditions—Conduct tests in the standard laboratory atmosphere of 23 ± 2°C and 50 ± 5% relative humidity, unless otherwise specified in this specification or by the contract or relevant material specification. In cases of disagreement, the tolerances shall be ±1°C and ±2% relative humidity.

TEST METHOD A FOR TESTING SOLID PLASTICS IN WATER (SPECIMENS 1 TO 50 g)

8. Scope

8.1 This test method involves weighing a one-piece specimen of 1 to 50 g in water, using a sinker with plastics that are lighter than water. This test method is suitable for plastics that are wet by, but otherwise not affected by water.

9. Apparatus

9.1 Analytical Balance—A balance with a precision of 0.1 mg or better is required for materials having densities less than 1.00 g/cm³ and sample weights less than 10 grams. For all other materials and sample weights, a balance with precision of 1 mg or better is acceptable (see Note 5). The balance shall be equipped with a stationary support for the immersion vessel above the balance pan ("pan straddle").

NOTE 5—The balance shall provide the precision that all materials tested have three significant figures on density. In case that materials with different densities are tested on one single balance, use the balance that provides at least three significant figures for all materials concerned.

NOTE 6—To assure that the balance meets the performance requirements, check on zero point and sensitivity frequently and perform periodic calibration.

9.2 Sample Holder, corrosion-resistant (for example, wire, gemholder, etc.).

9.3 Sinker—A sinker for use with specimens of plastics that have specific gravities less than 1.00. The sinker shall: (1) be corrosion-resistant; (2) have a specific gravity of not less than 7.0; (3) have smooth surfaces and a regular shape; and (4) be slightly heavier than necessary to sink the specimen. The sinker shall have an opening to facilitate attachment to the specimen and sample holder.

9.4 Immersion Vessel—A beaker or other wide-mouthed vessel for holding the water and immersed specimen.

9.5 Thermometer—A thermometer readable to 0.1°C or better.

10. Materials

10.1 Water—The water shall be substantially air-free and distilled or de-mineralized water.

NOTE 7—Air in water can be removed by boiling and cooling the water, or by shaking the water under vacuum in a heavy-walled vacuum flask. (Warning—Use gloves and shielding.) If the water does not wet the specimen, add a few drops of a wetting agent into the water. If this solution does not wet the specimen, Method B shall be used.

11. Test Specimen

11.1 The test specimen shall be a single piece of material with a size and shape suitable for the testing apparatus, provided that its volume shall be not less than 1 cm³ and its surface and edges shall be made smooth. The thickness of the specimen shall be at least 1 mm for each 1 g of weight. A specimen weighing 1 to 5 g was found to be convenient, but specimens up to approximately 50 g are also acceptable (see Note 8). Care shall be taken in cutting specimens to avoid changes in density resulting from compressive stresses or frictional heating.

NOTE 8—Specifications for certain plastics require a particular method of specimen preparation and should be consulted if applicable.

11.2 The specimen shall be free from oil, grease, and other foreign matter.

12. Procedure

12.1 Measure and record the water temperature.
12.2 Weigh the specimen in air. Weigh to the nearest 0.1 mg for specimens of mass 1 to 10 g and density less than 1.00 g/cm³. Weigh to the nearest 1 mg for other specimens.

12.3 If necessary, attach to the balance a piece of fine wire sufficiently long to reach from the hook above the pan to the support for the immersion vessel. In this case attach the specimen to the wire such that it is suspended about 25 mm above the vessel support.

Note 9—If a wire is used, weigh the specimen in air after hanging from the wire. In this case, record the mass of the specimen, \( a = (\text{mass of specimen + wire, in air}) \) - (mass of wire in air).

12.4 Mount the immersion vessel on the support, and completely immerse the suspended specimen (and sinkers, if used) in water (see 10.1) at a temperature of 23 ± 2°C. The vessel must not touch sample holder or specimen. Remove any bubbles adhering to the specimen, sample holder, or sinker, by rubbing them with a wire. Pay particular attention to holes in the specimen and sinker. If the bubbles are not removed by this method or if bubbles are continuously formed (as from dissolved gases), the use of vacuum is recommended (see Note 10). Determine the mass of the suspended specimen to the required precision (see 12.2) (see Note 11). Record this apparent mass as \( b \) (the mass of the specimen, sinker, if used, and the partially immersed wire in liquid). Unless otherwise specified, weigh rapidly in order to minimize absorption of water by the specimen.

Note 10—Some specimens may contain absorbed or dissolved gases, or irregularities which tend to trap air bubbles; any of these may affect the density values obtained. In such cases, the immersed specimen may be subjected to vacuum in a separate vessel until evolution of bubbles has substantially ceased before weighing (see Test Method B). It must also be demonstrated that the use of this technique leads to results of the required degree of precision.

Note 11—It may be necessary to change the sensitivity adjustment of the balance to overcome the damping effect of the immersed specimen.

12.5 Weigh the sample holder (and sinker, if used) in water with immersion to the same depth as used in the previous step (Notes 12 and 13). Record this weight as \( w \) (mass of the sample holder in liquid).

Note 12—If a wire is used, it is convenient to mark the level of immersion by means of a shallow notch filed in the wire. The finer the wire, the greater the tolerance is permitted in adjusting the level of immersion between weighings. With wire Avg No. 36 or finer, disregard its degrees of immersion and, if no sinker is used, use the mass of the wire in air as \( w \).

Note 13—If the wire is used and is left attached to the balance arm during a series of determinations, determine the mass \( a \) with the aid of a

tare on the other arm of the balance or as in Note 9. In such cases, care must be taken that the change of mass of the wire (for example, from visible water) between readings does not exceed the desired precision.

12.6 Repeat the procedure for the required number of specimens. Two specimens per sample are recommended. Determine acceptability of number of replicate test specimens by comparing results with precision data given in Tables 1 and 2. Use additional specimens if desired.

13. Calculation

13.1 Calculate the specific gravity of the plastic as follows:

\[
\text{sp gr} \ 23/23°C = a(a + w - b)
\]

where:

\( a \) = apparent mass of specimen, without wire or sinker, in air,

\( b \) = apparent mass of specimen (and of sinker, if used) completely immersed and of the wire partially immersed in liquid, and

\( w \) = apparent mass of totally immersed sinker (if used) and of partially immersed wire.

13.2 Calculate the density of the plastic as follows:

\[
D(\text{conversion to } 23°C, \text{ kg/m}^3) = \text{sp gr } 23/23°C \times 997.5
\]

13.3 If the temperature of the water is different than 23°C, use the density of water listed in Table 3 directly, or use the following equations to calculate the density of water at testing temperature:

\[
M = \Delta D/\Delta t
\]

\[
D(\text{conversion to } 23°C) = \text{sp gr } 23/23°C \times 997.5
\]

where:

\( M \) = slope,

\( \Delta D \) = difference between the lowest and highest temperature tolerance for the standard density of water (\( D @ 21°C \) – \( D @ 25°C \)),

\( \Delta t \) = difference between the highest and lowest temperature tolerance recommended, (21°C–25°C),

\( t_a \) = temperature of air, and

\( t_w \) = temperature of water.

14. Report

14.1 Report the following information:

### TABLE 1 Test Method A Specific Gravity Tested in Water

<table>
<thead>
<tr>
<th>Material</th>
<th>Mean</th>
<th>( S_e^d )</th>
<th>( S_e^c )</th>
<th>( c^d )</th>
<th>( R^d )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polypropylene</td>
<td>0.9007</td>
<td>0.00196</td>
<td>0.00297</td>
<td>0.00555</td>
<td>0.00841</td>
</tr>
<tr>
<td>Cellulose Acetate Butyrate</td>
<td>1.1973</td>
<td>0.00232</td>
<td>0.00304</td>
<td>0.00657</td>
<td>0.00960</td>
</tr>
<tr>
<td>Polystyrene Sulfide</td>
<td>1.1708</td>
<td>0.00540</td>
<td>0.00738</td>
<td>0.01528</td>
<td>0.02089</td>
</tr>
<tr>
<td>Thermoset</td>
<td>1.3136</td>
<td>0.00271</td>
<td>0.00313</td>
<td>0.00876</td>
<td>0.02171</td>
</tr>
<tr>
<td>Polyvinyl Chloride</td>
<td>1.3396</td>
<td>0.00243</td>
<td>0.00615</td>
<td>0.00888</td>
<td>0.01947</td>
</tr>
</tbody>
</table>

\( S_e^d \) = within laboratory standard deviation for the individual material. It is obtained by pooling the within-laboratory standard deviations of the test results from all of the participating laboratories.

\( S_e^c \) = between-laboratories reproducibility, expressed as standard deviation: \( S_e = (S_e^d)^2 + (S_e^c)^2 \)

\( c^d \) = within laboratory critical interval between two test results = 2.8 \times S_e^d

\( R^d \) = between-laboratory critical interval between two test results = 2.8 \times S_e^c
### TABLE 2 Test Method B Specific Gravity Tested in Liquids Other Than Water

<table>
<thead>
<tr>
<th>Material</th>
<th>Mean</th>
<th>$S_a$</th>
<th>$S_a^2$</th>
<th>$S_b$</th>
<th>$R$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polypropylene</td>
<td>0.9023</td>
<td>0.00139</td>
<td>0.00239</td>
<td>0.00393</td>
<td>0.00669</td>
</tr>
<tr>
<td>LDPE</td>
<td>0.9215</td>
<td>0.00109</td>
<td>0.00193</td>
<td>0.00308</td>
<td>0.00546</td>
</tr>
<tr>
<td>HDPE</td>
<td>0.9678</td>
<td>0.00126</td>
<td>0.00189</td>
<td>0.00336</td>
<td>0.00529</td>
</tr>
<tr>
<td>Thermoset</td>
<td>1.3130</td>
<td>0.00160</td>
<td>0.00217</td>
<td>0.00453</td>
<td>0.00688</td>
</tr>
</tbody>
</table>

$S_a$ = within laboratory standard deviation for the individual material. It is obtained by pooling the within-laboratory standard deviations of the test results from all of the participating laboratories.

$S_b$ = between-laboratories reproducibility, expressed as standard deviation: $S_b = \sqrt{\frac{\sum (s_i^2)}{n}}$

$C = \frac{S_b}{S_a}$ = between-laboratory critical interval between two test results $= 2.8\times S_a$.

$R = \frac{S_a}{S_a}$ = between-laboratory critical interval between two test results $= 2.8\times S_a$.

### TABLE 3 Standard Density of Water

<table>
<thead>
<tr>
<th>°C</th>
<th>$\rho_{\text{kg/m}^3}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>21</td>
<td>997.9948</td>
</tr>
<tr>
<td>22</td>
<td>997.7720</td>
</tr>
<tr>
<td>23</td>
<td>997.5412</td>
</tr>
<tr>
<td>24</td>
<td>997.2994</td>
</tr>
<tr>
<td>25</td>
<td>997.0480</td>
</tr>
</tbody>
</table>


The leading figure decreases by 1.

14.1.1 Complete identification of the material or product tested, including method of specimen preparation and conditioning.

14.1.2 Average specific gravity (relative density) for all specimens from a sampling unit corrected to 23.0°C (Table 3) are reported as sp gr 23/23°C = $\rho$, or average density reported as $D_{23} = \rho$ kg/m³.

Note 14—Reporting density in g/cm³ is also acceptable provided that it is agreed upon by the users.

14.1.3 A measure of the degree of variation of specific gravity or density within the sampling unit such as the standard deviation and number of determinations on a homogeneous material or the averages plus these measures of dispersion on different layers or areas of a nonhomogeneous product.

14.1.4 Report the temperature of the water.

14.1.5 Report the density and specific gravity with three significant figures.

14.1.6 Any evidence of porosity of the material or specimen.

14.1.7 The method of test (that is, Method A of Test Method D 792), and 

14.1.8 Date of test.

15. Precision and Bias

15.1 See Section 23.

### TEST METHOD B FOR TESTING SOLID PLASTICS IN LIQUIDS OTHER THAN WATER (SPECIMENS 1 TO 50 g)

16. Scope

16.1 Test Method B uses a liquid other than water for testing one-piece specimens, 1 to 50 g, of plastics that are affected by water or are lighter than water.

17. Apparatus

17.1 The apparatus shall include the balance, wire, and immersion vessel of Section 8, and, optionally, the following:

17.2 Pycometer with Thermometer—A 25-mL specific gravity bottle with thermometer, or

17.3 Pycometer—A pycometer of the Weld type, preferably with a capacity of about 25 mL and an external cap over the stopper.

17.4 Thermometer—A thermometer having ten divisions per degree Celsius over a temperature range of not less than 5°C or 10°F above and below the standard temperature, and having an ice point for calibration. A thermometer short enough to be handled inside the balance case will be found convenient. ASTM Thermometer 23°C (see Specification E 1) and Anschütz-type thermometers have been found satisfactory for this purpose.

17.5 Constant-Temperature Bath—An appropriate constant-temperature bath adjusted to maintain a temperature of 23 ± 0.1°C.

18. Materials

18.1 Immersion Liquid—The liquid used shall not dissolve, swell, or otherwise affect the specimen, but shall wet it and shall have a specific gravity less than that of the specimen. In addition, the immersion liquid shall be non-hygrosopic, has a low vapor pressure, a low viscosity, and a high flash point, and shall leave little or no waxy or tarry residue on evaporation. A narrow cut distilled from kerosine meets these requirements for many plastics. The specific gravity 23/23°C of the immersion liquid shall be determined shortly before and after each use in this method to a precision of at least 0.1% relative, unless it has been established experimentally in the particular application that a lesser frequency of determination also provides the desired precision.

Note 15—For the determination of the specific gravity of the liquid, the use of a standard plummet of known volume or of Method A, C, or D of Test Methods D 891, using the modifications required to give specific gravity 23/23°C instead of specific gravity 60/60°F, is recommended. One suggested procedure is the following:
If a constant-temperature water bath is not available, determine the mass of the clean, dry pycnometer with thermometer to the nearest 0.1 mg on an analytical balance. Fill the pycnometer with water (10.1) cooler than 23°C. Insert the thermometer-stopper, causing excess water to be expelled through the side arm. Permit the filled bottle to warm in air until the thermometer reads 23.0°C. Remove the drop of water at the tip of the side arm with a bit of filter paper, taking care not to draw any liquid from within the capillary, place the cap over the side arm, wipe the outside carefully, and determine the mass of the filled bottle again to the nearest 0.2 mg. Empty the pycnometer, dry, and fill with immersion liquid. Determine the mass with the liquid in the same manner as was done with the water. Calculate the specific gravity 23/23°C of the liquid, \( d \), as follows:

\[
d = (b - e)/(w - e)
\]

where:
- \( e \) = apparent mass of empty pycnometer,
- \( w \) = apparent mass of pycnometer filled with water at 23.0°C, and
- \( b \) = apparent mass of pycnometer filled with liquid at 23.0°C.

If a constant-temperature water bath is available, a pycnometer without a thermometer may be used (compare 30.2).

Note 16—One standard object which has been found satisfactory for this purpose is the Reimann Thermometer Plummet. These are normally supplied calibrated for measurements at temperatures other than 23/23°C, so that recalibration is not necessary for the purposes of these methods.

19. Test Specimen
19.1 See Section 11.

20. Procedure
20.1 The procedure shall be similar to Section 12, except for the choice of immersion liquid, and the temperature during the immersed weighing (12.3) shall be 23 ± 0.5°C.

21. Calculation
21.1 The calculations shall be similar to Section 13, except that \( d \), the specific gravity 23/23°C of the liquid, shall be placed in the numerator: (see 13.1)

\[
Sp \, gr \, 23/23°C = (a \times d)/(a + w - h)
\]

22. Report
22.1 See Section 14.

23. Precision and Bias
23.1 Tables 1 and 2 are based on an interlaboratory study conducted in 1985 in accordance with Practice E 691, involving 5 materials tested with Test Method A by six laboratories or four materials tested with Test Method B by six laboratories. Each test result was based on two individual determinations and each laboratory obtained four test results for each material. (Warning—The explanations of \( r \) and \( R \) are only intended to present a meaningful way of considering the approximate precision of these test methods. The data of Tables 1 and 2 should not be applied to acceptance or rejection of materials, as these data apply only to the materials tested in the round robin and are unlikely to be rigorously representative of other lots, formulations, conditions, materials, or laboratories. Users of this test method should apply the principles outlined in Practice E 691 to generate data specific to the materials and laboratory (or between specific laboratories). The principles of 23.2-23.2.3 would then be valid for such data.)

23.2 Concept of \( r \) and \( R \) in Tables 1 and 2—If \( S_r \) and \( S_R \) have been calculated from a large enough body of data, and for test results that were averages from 4 test results for each material, then:

23.2.1 Repeatability—Two test results obtained within one laboratory shall be judged not equivalent if they differ by more than the \( r \) value for that material. The concept \( r \) is the interval representing the critical difference between two test results for the same material, obtained by the same operator using the same equipment on the same day in the same laboratory.

23.2.2 Reproducibility—Two test results obtained by different laboratories shall be judged not equivalent if they differ by more than the \( R \) value for that material. The concept \( R \) is the interval representing the critical difference between two test results for the same material, obtained by different operators using different equipment in different laboratories.

23.2.3 Any judgment in accordance with 23.2.1 or 23.2.2 would have an approximate 95% (0.95) probability of being correct.

23.3 There are no recognized standards by which to estimate bias of this test method.

24. Keywords
24.1 density; relative density; specific gravity

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4 Supporting data are available from ASTM Headquarters. Request RR:D20-1133.
SUMMARY OF CHANGES

Committee D20 has identified the location of selected changes to this standard since the last issue (D 792 - 00) that may impact the use of this standard. (June 15, 2008)

1. Deleted references to D 1898, Old Note 1, Note 6, and rearranged the order of Notes.
2. Revised Note 1, ISO Statement.
4. Re-defined specific gravity in by changing “unit volume” to “given volume.” Removed “equal density of.”
6. Clarified definition of ΔD and deleted “...” in equation in 13.3.
7. Changed significant figures from four to three to harmonize with Guide D 6436 in 14.1.5.
8. Added Note 5 and Note 14.
9. Editorial changes, including the removal of permissive languages from the text.

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