1. Scope*

1.1 This fire-test-response test method covers a small-scale laboratory screening procedure for comparing the relative linear rate of burning or extent and time of burning, or both, of plastics in the form of bars, molded or cut from sheets, plates, or panels, and tested in the horizontal position.

Note 1—This test method, and test method A of IEC 60695-11-10 are technically equivalent.

Note 2—For additional information on materials which do not burn to the first reference mark by this test, see Test Method D 3801.

1.2 This test method was developed for polymeric materials used for parts in devices and appliances. The results are intended to serve as a preliminary indication of their acceptability with respect to flammability for a particular application. The final acceptance of the material is dependent upon its use in complete equipment that conforms with the standard applicable to such equipment.

1.3 The classification system described in the appendix is intended for quality assurance and the preselection of component materials for products.

1.4 This standard is used to measure and describe the response of materials, products, or assemblies to heat and flame under controlled conditions, but does not by itself incorporate all factors required for fire hazards or fire risk assessment of materials, products, or assemblies under actual fire conditions.

1.5 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 hazards statements, see 9.2.1.

2. Referenced Documents

2.1 ASTM Standards:

D 648 Test Method for Deflection Temperature of Plastics Under Flexural Load
D 883 Terminology Relating to Plastics
D 3801 Test Method for Measuring the Comparative Burning Characteristics of Solid Plastics in a Vertical Position
D 5025 Specification for a Laboratory Burner Used for Small-Scale Burning Tests on Plastic Materials
D 5207 Practice for Calibration of 20 and 125-mm Test Flames for Small-Scale Burning Tests on Plastic Materials
E 176 Terminology of Fire Standards
E 691 Practice for Conducting an Interlaboratory Study to Determine the Precision of a Test Method

2.2 IEC Standards:

IEC 60695-11-10 Fire Hazard Testing - Part 11-10 Test Flames - 50W Horizontal and Vertical Flame Test Methods

3. Terminology

3.1 Definitions:

3.1.1 Definitions used in this test method are in accordance with Terminology D 883, unless otherwise specified. For terms relating to fire, the definitions used in this test method are in accordance with Terminology E 176.

4. Summary of Test Method

4.1 A bar specimen of the material to be tested is supported horizontally at one end. The free end is exposed to a specified gas flame for 30 s. Time and extent of burning are measured and reported if the specimen does not burn 100 mm. An average burning rate is reported for a material if it burns to the 100 mm mark from the ignited end.
5. Significance and Use

5.1 Tests made on a material under conditions herein prescribed are of value in comparing the rate of burning or extent and time of burning characteristics, or both, of different materials, in controlling manufacturing processes, or as a measure of deterioration or change in these burning characteristics prior to or during use. Correlation with flammability under actual use conditions is not implied.

5.2 The rate of burning and other burning phenomena will be affected by such factors as density, pigments, any anisotropy of the material and the thickness of the specimen. Test data shall be compared only for specimens of similar thickness, whether comparisons are being made with the same or different materials. The rate of burning and other burning phenomena will vary with thickness.

5.3 It is feasible that sheet materials that have been stretched during processing will relax during burning and give erratic results unless they are first heated above their deflection temperature, in accordance with Test Method D 648, for a time sufficient to permit complete relaxation.

5.4 Burning tests require that certain variables be arbitrarily fixed, for example, specimen size, energy source and application time, and end points. Materials will be found that are unusually sensitive to one or more of the conditions chosen for this method leading to highly variable results. Additional burning characterization by other methods is highly desirable in such cases (see Note 2).

5.5 In this procedure, the specimens are subjected to specific laboratory test conditions. If different test conditions are substituted or the end-use conditions are changed, it will not always be possible by or from this test to predict changes in the fire-test-response characteristics measured. Therefore, the results are valid only for the fire-test-exposure conditions described in this procedure.

6. Apparatus

6.1 Test Chamber, enclosed laboratory hood, or chamber free of induced or forced draft during test, having an inside volume of at least 0.5 m³. An enclosed laboratory hood with a heat-resistant glass window for observing the test and an exhaust fan for removing the products of combustion after the tests is recommended. The atmosphere in and around the test exhaust fan for removing the products of combustion after the heat-resistant glass window for observing the test and an inclined at 45° angle as illustrated in Fig. 1.

6.2 Test Fixture, a laboratory ring stand or test fixture equipped with a means of holding a 125 mm² wire gauze horizontal and a small clamp permitting the specimen to be held with its longitudinal axis horizontal and its transverse axis inclined at 45 ± 2° angle as illustrated in Fig. 1.

6.3 Laboratory Burner, constructed in accordance with Specification D 5025.

6.4 Gas Supply, a supply of technical-grade methane gas with suitable regulator and meter for uniform gas flow. Natural gas mixtures having an energy density of approximately 37 MJ/m³ have been found to provide similar results. However, technical-grade methane gas shall be used as the referee in cases of dispute.

6.5 Wire Gauze, 20-mesh (approximately 20 openings per 25 mm), made with 0.43 ± 0.03 mm diameter iron wire cut to approximately 125 mm², to sustain burning or glowing particles falling from the specimens.

6.6 Timing Device, accurate to 0.5 s.

6.7 Scale, graduated in millimetres.

6.8 Micrometer, accurate to 0.05 mm.

6.9 Conditioning Room or Chamber, capable of being maintained at 23 ± 2°C and 50 ± 5 % relative humidity.

6.10 Flexible Specimen Support Fixture, used to facilitate the testing of specimens that sag and touch the wire gauze. (See 9.4 and Fig. 2.)

7. Test Specimens

7.1 All test specimens shall be cut from a representative sample of the material (sheet or end products), or shall be cast or injection-, compression-, transfer- or pultrusion-molded to the necessary form. After any cutting operation, care shall be taken to remove all dust and any particles from the surface; cut edges shall be fine sanded to have a smooth finish. Unless otherwise agreed, fabrication of test specimens shall be in accordance with the specifications of the material being tested.

7.2 Specimens shall be 125 ± 5 mm long by 13.0 ± 0.5 mm wide, and provided in the minimum thickness and in the 3.0 (−0.0 +0.2) mm thickness. The 3.0 mm thick specimens are not necessary if the minimum thickness is greater than 3.0 mm, or the maximum thickness is less than 3.0 mm. The maximum thickness shall not exceed 13 mm. The maximum width shall not exceed 13.5 mm. The edges shall be smooth, and the radius on the corners shall not exceed 1.3 mm.

7.3 It is possible that the results of tests carried out on test specimens of different colors, thicknesses, densities, molecular masses, directions of anisotropy and types, or with different additives, fillers/reinforcements will be different.

7.3.1 Test specimens in the minimum and maximum densities, melt flows and level of fillers/reinforcements contents shall be considered representative of the range, if the results yield the same flame test classification. If the burning characteristics are not essentially the same for all specimens representing the range, the evaluation is to be limited only to the materials in the densities, melt flows, and fillers/reinforcements contents tested. Additional specimens in the intermediate densities, melt flows, and fillers/reinforcements contents are to be tested.

7.3.2 Uncolored test specimens and test specimens with the highest level of organic and inorganic pigment loading by weight are considered representative of the color range, if the test results are essentially the same. When certain pigments are
known to affect flammability characteristics, they are also to be tested. Specimens to be tested are those that:

(a) contain no coloring  
(b) contain the highest level of organic pigments  
(c) contain the highest level of inorganic pigments  
(d) contain pigments which are known to adversely affect flammability characteristics

8. Conditioning

8.1 Condition ten bar specimens for each material and thickness to be tested in accordance with Test Method D 618 at 23 ± 2°C and 50 ± 5% relative humidity for a minimum of 48 h. Once removed from the conditioning atmosphere test the specimens within 1 h.

8.2 Conduct testing in a laboratory atmosphere of 15 to 35°C and 45 to 75% relative humidity.

9. Procedure

9.1 Prepare at least ten bar specimens. After measuring and recording the specimen thickness, mark each specimen with two lines perpendicular to the longitudinal axis of the bar, 25 ± 1 and 100 ± 1 mm from the end that is to be ignited.
9.2 Conduct the burning test in a chamber, enclosure, or laboratory hood free of induced or forced draft.

**Note** 7—**Warning:** Products of combustion may be toxic. An enclosed laboratory hood and an exhaust fan for removing the products of combustion after the tests are recommended. The exhaust fan is turned off during the test and turned on immediately following the test in order to remove products of combustion.

9.3 Clamp the specimen at the end farthest from the 25 mm reference mark, in a support with its longitudinal axis horizontal and its transverse axis inclined at 45 ± 2° as illustrated in Fig. 1. Clamp the wire gauze horizontally beneath the specimen, with a distance of 10 ± 1 mm between the lowest edge of the specimen and the wire gauze, and with the free end of the specimen even with the edge of the gauze. Any material remaining on the wire gauze from the previous test must be burned off or a new section of wire gauze used for each test.

9.4 If the specimen sags at its free end during the initial setup and is not able to maintain the distance of 10 ± 1 mm as specified in 9.2, the flexible specimen support fixture illustrated in Fig. 2 shall be used. Position the support fixture under the specimen with the small extending portion of the support fixture at least 20 mm from the free end of the specimen. Provide enough clearance at the clamped end of the specimen so that the support fixture can be moved freely sideward. As the flame front progresses along the specimen, withdraw the support fixture at the same approximate rate, preventing the flame front from contacting the flexible specimen support fixture, so that there is no effect on the test flame or on the burning of the specimen.

9.5 With the central axis of the burner tube in the vertical position, place the burner remote from the specimen, ignite, and adjust it to produce a blue flame 20 mm high. Adjust the gas supply and the air ports of the burner until a 20-mm yellow-tipped blue flame is produced, and then increase the air supply until the yellow tip just disappears. Measure the height of the flame. If the flame height is not 20 ± 2 mm, adjust the burner gas supply to give the proper flame height. Once the flame has been properly set to a height of 20 ± 2 mm wait for at least 5 min to allow the burner conditions to reach equilibrium.

**Note** 8—See Practice D 5207 for recommended back pressure and flow rate for the gas supply and calibration procedure for the 20 mm flame.

9.6 Place the burner so that the test flame impinges on the free end of the test specimen to a depth of approximately 6 mm starting the timing device simultaneously. The central axis of the burner tube is to be in the same vertical plane as the longitudinal bottom edge of the specimen and inclined toward the end of the specimen at an angle of approximately 45 ± 2 degrees to the horizontal. See Fig. 1. Apply the flame for 30 ± 1 s without changing its position. If the test specimen shrinks from the applied flame without ignition, the material is not suitable for evaluation by these test methods. Excessive distortion of the specimen during the test will invalidate the results. Withdraw the test flame sufficiently from the specimen (see Note 9) so that there is no effect on the specimen after 30 ± 1 s or as soon as the flame front of the specimen reaches the 25 mm mark (if less than 30 s). Restart the timing device when the flame front reaches the 25 mm reference mark.

**Note** 9—Withdrawing the burner a distance of 150 mm from the specimen has been found satisfactory.

9.7 If the specimen continues to burn, with a flame or glowing combustion (visible glow without flame), after removal of the test flame, record the elapsed time (t), in seconds, for the flame front to travel from the 25 mm reference mark to the 100 mm reference mark and record the burned length (L), as 75 mm. If the flame front passes the 25 mm reference mark but does not reach the 100 mm reference mark, record the elapsed time (t), in seconds, and the burned length (L), in millimetres between the 25 mm reference mark and where the flame front stopped.

9.8 Repeat the test procedure (9.1-9.7) until three specimens have burned to or beyond the 100 mm reference mark, or ten specimens have been tested.

**Note** 10—For classification purposes, if only one specimen does not comply with the criteria, test an additional set of specimens. See X1.5.

10. Calculation

10.1 Calculate the linear burning rate (V), in millimetres per minute, for each specimen where the flame front reaches the 100 mm reference mark using the equation:

\[ V = 60L/t \]

where:

- \( L \) = the burned length, in millimetres, as defined in 9.7; and
- \( t \) = the time, in seconds, as defined in 9.7.

**Note** 11—If the flame front reached the 100-mm reference mark, \( L = 75 \).

**Note** 12—The SI units of the linear burning rate is metre per second. In practice, the unit millimetre per minute is used.

**Note** 13—It is acceptable to report the results in cm/min by using the method prescribed in 10.1 and then dividing the obtained rate by ten.

10.2 Calculate the average linear burning rate or classify the material in accordance with the appendix.

11. Report

11.1 Include the following in the complete report:

11.1.1 **Material Identification**—Include generic description, manufacturer, commercial designation, lot number, and color.

11.1.2 The thickness, as measured with a micrometer to the nearest 0.1 mm, of the test specimen.

11.1.3 The nominal apparent density (rigid cellular materials only).

11.1.4 The direction of any anisotropy relative to the test specimen dimensions.

11.1.5 Conditioning treatment.

11.1.6 Any prior treatment before testing, other than cutting, trimming and conditioning.

11.1.7 Whether or not the specimen continued to burn (with or without visible flame) after application of test flame.

11.1.8 Whether or not the flame front reached the 25 and 100 mm reference marks.

11.1.9 For specimens with which the flame front does not reach or pass the 25 mm reference mark, a statement that indicates the flame front did not reach or pass the 25 mm reference mark. Do not report an elapsed time (t) and burned length (L).
11.1.10 For specimens with which the flame front passed the 25 mm reference mark but did not reach the 100 mm reference mark, the elapsed time \((t)\) and burned length \((L)\).

11.1.11 If a specimen does not burn to the 100 mm mark because of dripping, flowing, or falling burning particles, the report must so indicate.

11.1.12 If a specimen is reignited by burning material on the gauze, the report must so state.

11.1.13 For specimens with which the flame front reached the 100 mm reference mark, the average linear burning rate, \((V)\).

11.1.14 Whether the flexible specimen support fixture was used.

11.1.15 The caveat contained in 1.4 herein shall be incorporated in its entirety in the test report issued.

11.1.16 Optional—Flame classification as determined from the appendix.

12. Precision and Bias

12.1 Table 1 is based on a round robin completed in 1987\(^8\) in accordance with Practice E 691, involving three self-supporting materials tested by eleven laboratories. Each laboratory conditioned, at 23°C and 50 % relative humidity, the specimens that it tested. Each laboratory conducted the tests in a laboratory hood with the hood exhaust essentially turned off. All three materials were classified by the test as possessing an average burning rate. Each test result consisted of an average linear burning rate determined from three specimens. Each laboratory obtained three test results for each material.

12.2 Table 2 is based on a round robin completed in 1986\(^9\) in accordance with Practice 691, involving four materials that required use of the flexible specimen support fixture and tested by six different laboratories. For each material, all samples were provided by one source. The individual specimens were cut and distributed by one laboratory. Each laboratory conditioned, at 23°C and 50 % relative humidity, the specimens that it tested. Each laboratory obtained three test results for each material.

**TABLE 1 Average Linear Burning Rate for Specimens Tested Without Flexible Specimen Support Fixture**

<table>
<thead>
<tr>
<th>Material</th>
<th>Nominal Specimen Thickness, mm</th>
<th>Average</th>
<th>(S_r^A)</th>
<th>(S_r^B)</th>
<th>(C_r)</th>
<th>(D_r^D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyethylene (PE)</td>
<td>3.0</td>
<td>15.2</td>
<td>0.7</td>
<td>1.3</td>
<td>1.9</td>
<td>3.7</td>
</tr>
<tr>
<td>ABS</td>
<td>3.2</td>
<td>27.9</td>
<td>2.1</td>
<td>4.1</td>
<td>5.7</td>
<td>11.5</td>
</tr>
<tr>
<td>Acrylic</td>
<td>3.0</td>
<td>29.7</td>
<td>1.7</td>
<td>2.2</td>
<td>4.9</td>
<td>6.1</td>
</tr>
</tbody>
</table>

\(^A\) \(S_r\) is the within-laboratory standard deviation for the indicated material. It is obtained by pooling the within-laboratory standard deviations of the test results from all of the participating laboratories:

\[
S_r = \left[ \sum (s_j^2) \right]^{1/2}
\]

\(^B\) \(S_r\) is the between-laboratories reproducibility, expressed as stated deviation:

\[
S_r = [(S_r^1)^2 + (S_r^2)^2 + \ldots + (S_r^n)^2]^{1/2}
\]

where: \(S_r\) = the standard deviation of laboratory means.

\(^C\) \(r\) is the within-laboratory critical interval between two test results = \(2.8 \times S_r\).

\(^D\) \(R\) is the between-laboratories critical interval between two test results = \(2.8 \times S_r\).

**TABLE 2 Average Linear Burning Rate for Specimens Tested With Flexible Specimen Support Fixture**

<table>
<thead>
<tr>
<th>Material</th>
<th>Nominal Specimen Thickness, mm</th>
<th>Average</th>
<th>(S_r^A)</th>
<th>(S_r^B)</th>
<th>(C_r)</th>
<th>(D_r^D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyurethane (PUR)</td>
<td>0.1</td>
<td>41.6</td>
<td>1.0</td>
<td>14.4</td>
<td>30.6</td>
<td>40.4</td>
</tr>
<tr>
<td>Polyurethane (PUR)</td>
<td>0.8</td>
<td>60.9</td>
<td>10.9</td>
<td>26.6</td>
<td>29.8</td>
<td>74.4</td>
</tr>
<tr>
<td>Polyurethane (PUR)</td>
<td>0.4</td>
<td>82.3</td>
<td>10.6</td>
<td>26.6</td>
<td>29.8</td>
<td>74.4</td>
</tr>
<tr>
<td>Polyethylene terephthalate (PET)</td>
<td>0.1</td>
<td>192.0</td>
<td>32.2</td>
<td>123.7</td>
<td>(E)</td>
<td>(E)</td>
</tr>
</tbody>
</table>

\(^A\) \(S_r\) = within-laboratory standard deviation for the indicated material. It is obtained by pooling the within-laboratory standard deviations of the test results from all of the participating laboratories, as follows:

\[
S_r = \left[ \sum (s_j^2) + (s_2^2) + \ldots + (s_n^2) \right]^{1/2}
\]

\(^B\) \(S_r\) = between-laboratories reproducibility, expressed as stated deviation, as follows:

\[
S_r = [(S_r^1)^2 + (S_r^2)^2 + (S_r^n)^2]^{1/2}
\]

where \(S_r\) = the standard deviation of laboratory means.

\(^C\) \(r\) = within-laboratory critical interval between two test results = \(2.8 \times S_r\).

\(^D\) \(R\) = between-laboratories critical interval between two test results = \(2.8 \times S_r\).

\(^E\) The number of laboratories in the interlaboratory study reporting a linear burning rate was too small to establish a between-laboratory standard deviation.

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\(^8\) Supporting data for Table 1 are available from ASTM Headquarters. Request RR: D-20-1149.

\(^9\) Supporting data for Table 2 are available from ASTM Headquarters. Request RR: D-20-1146.
and therefore shall not be used as a referee test method for
these two characteristics in case of dispute. Due to the rarity of
materials which consistently produce this result, a numerical
precision and bias statement for this type of test result is not
being actively pursued at this time.

NOTE 14—Caution: The explanations of “r” and “R” given in 12.4-
12.4.3 are only intended to present a meaningful way of considering the
approximate precision of this test method. The data in Tables 1 and 2 shall
not be rigorously applied to acceptance or rejection of material, 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 shall apply the
principles outlined in Practice E 691 to generate data specific to their
materials and laboratory (or between specific laboratories). The principles
of 12.4-12.4.3 would then be valid for such data.

12.4 Concept of “r” and “R”—If \( S_r \) and \( S_R \) have been
calculated from a large enough body of data, and for test results
that were averages from testing three specimens for each test
result, then:

12.4.1 Repeatability, \( r \)—Two test results obtained within
one laboratory shall be judged not equivalent if they differ by
more than the “\( r \)” value for the material. “\( r \)” 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.

12.4.2 Reproducibility, \( R \)—Two test results obtained from
different laboratories shall be judged not equivalent if they
differ by more than the “\( R \)” value for the material. “\( R \)” 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.

12.4.3 Judgments in accordance with 12.4.1 and 12.4.2 have
an approximate 95% (0.95) probability of being correct.

12.5 Bias—There are no recognized standards on which to
base an estimate of bias for this test method.

13. Keywords

13.1 burning characteristics; combustion; extent of burning;
flammability; HB; horizontal burning rate; plastics; rate of
burning; small-scale burning test burning; time of burning

APPENDIX

(Nonmandatory Information)

X1. CLASSIFICATION SYSTEM FOR DETERMINING THE RELATIVE LINEAR RATE OF BURNING AND/OR EXTENT
AND TIME OF BURNING OF PLASTICS

X1.1 General

X1.1.1 This appendix covers a classification system for
characterizing the burning behavior of plastic materials, sup-
ported in a horizontal position, in response to a small-flame
ignition source. The use of a category designation code is
optional and is determined by examining the test results of
materials tested by this method. Each category code represents
a preferred range of performance levels that simplifies descrip-
tion in material designations or specifications and may assist
certification bodies to determine compliance with applicable
requirements.

X1.2 Category Designation—The behavior of specimens
shall be classified HB (HB = horizontal burning) if,

X1.2.1 There are no visible signs of combustion after the
ignition source is removed, or

X1.2.2 The flame front does not pass the 25 mm reference
mark, or

X1.2.3 The flame front passes the 25 mm reference mark
but does not reach the 100 mm reference mark, or

X1.2.4 The flame front reaches the 100 mm reference mark
and the linear burning rate does not exceed 40 mm/min for
specimens having a thickness between 3 and 13 mm or 75
mm/min for specimens having a thickness less than 3 mm.

X1.3 If only one specimen from the first set of specimens
does not comply with the criteria indicated, another set of
specimens is to be tested. All specimens from this second set
shall comply with the criteria indicated in order for the
material, of that thickness, to be classified as HB.

X1.4 If the linear burning rate does not exceed 40 mm/min
when tested in the 3.0 mm ± 0.2 mm thickness, the HB
category designation shall be extended to a 1.5 mm minimum
thickness.

X1.5 Recording the category designation in the test report
is optional.
SUMMARY OF CHANGES

This section identifies the location of selected changes to this test method. For the convenience of the user, Committee D20 has highlighted those changes that may impact the use of this test method. This section may also include descriptions of the changes or reasons for the changes, or both.

D 635 – 03:
(1) Revised 1.4 for E5 reference boilerplate.
(2) Added test specimen selection criteria in 7.3.
(3) Added additional Classification criteria in X1.4.
(4) Updated Figure 1.

D 635 – 97:
(1) Revisions were made to clarify the Scope.
(2) An IEC/ISO equivalency statement was added.
(3) Handling of specimens that are flexible was incorporated.
(4) Permissive language was removed.
(5) Specifications D 5207 for test-flame calibration was added.
(6) Calculations were updated.
(7) Precision and Bias information was clarified.
(8) The appendix was added.

D 635 – 98:
(1) Revised 12.3 concerning precision and bias for specimens that do not burn to the 100-mm reference mark.