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# Standard Test Method for Distillation of Road Tars<sup>1</sup>

This standard is issued under the fixed designation D 20; 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 ( $\epsilon$ ) indicates an editorial change since the last revision or reapproval.

## 1. Scope

1.1 This test method covers the distillation of road tars.

1.2 This standard does not purport to address all of the safety problems, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

#### 2. Referenced Documents

2.1 ASTM Standards:

E 1 Specification for ASTM Thermometers<sup>2</sup>

#### 3. Summary of Test Method

3.1 A 100-g sample is distilled at a controlled rate from a 300-mL flask into tared receivers. The masses of distillate fractions at a series of specified temperatures and of residue at the maximum specified temperature are determined. If desired, the residue and distillates may be used for further testing.

#### 4. Significance and Use

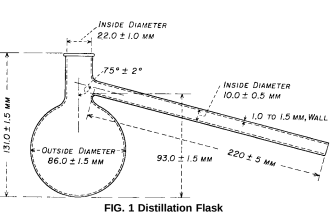
4.1 The distillation test separates tar into fractions according to a series of specified temperatures.

#### 5. Apparatus

5.1 *Flask*—A side-arm distillation flask, as shown in Fig. 1, conforming to the following dimensions:

Diameter of bulb, outside, mm Diameter of neck, inside, mm Diameter of side-arm, inside, mm Height of flask, outside, mm Vertical distance, bottom of bulb, outside, to hori-	$\begin{array}{c} 86.0\pm1.5\\ 22.0\pm1.0\\ 10.0\pm0.5\\ 131.0\pm1.5\\ 93.0\pm1.5\end{array}$
zontal tangent at side-arm, inside, mm Length of side-arm, mm Angle of side-arm, deg Thickness of side-arm wall, mm	220 ± 5 75 ± 2 1.0 to 1.5

5.2 *Condenser Tube*— A tapered glass condenser, as shown in Fig. 2 and Fig. 3, having the following dimensions:



utside diameter of small end, mm	12.5 ±
utside diameter of large end, mm	28.5 ± 3

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12.0 - 1.0
$28.5\pm3.0$
$360 \pm 4$
$100 \pm 5$

5.3 *Source of Heat*— A heat source consisting of a bunsen or meker-type burner (Note 1) or an electric heater. The electric heater<sup>3</sup> shall have an output variable of 750 W and an upper refractory with dimensions as shown in Fig. 4. The temperature of the heater shall be controlled by a variable transformer or rheostat suitable for the voltage used, and shall be fitted with a clamp for mounting on a vertical support rod.

Note 1—An artificial gas model used with natural gas has been found to give a uniform and easily controlled source of heat.

5.4 *Flask Shield and Cover for Flame Distillation*—A steel shield (preferably stainless) lined with 3-mm Transite board, non-asbestos, with two-part cover made from 6-mm Transite board of the form and dimensions shown in Fig. 5.

5.5 Flask Shield and Cover for Electric-Heater Distillation—A steel shield (preferably stainless) fitted with mica windows, and a cover of the same construction and dimensions as those for flame distillation (5.4) except for the height of the shield (see Fig. 6).

5.6 Gauze for Flame Distillation—Two sheets of 1.0-mm opening wire gauze made of 0.56-mm diameter nickel-chromium wire measuring a minimum of 125 mm in diameter or (125 by 125-mm) square.

5.7 Burner Chimney for Flame Distillation—Construct a cylindrical metal shield approximately 100 mm high, 95 to 105

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<sup>&</sup>lt;sup>2</sup> Annual Book of ASTM Standards, Vol 14.03.

<sup>&</sup>lt;sup>3</sup> The Precision Ful-Kontrol 750-W heater with built-in variable transformer control, available from Precision Scientific Co., Chicago, IL, has been found satisfactory. This heater is only available for 115 V. 50/60 Hz.

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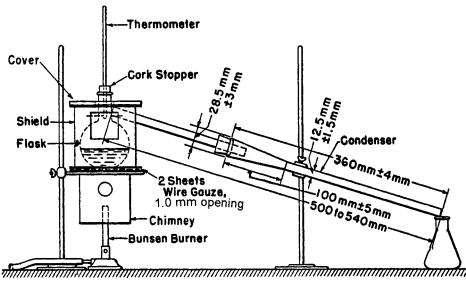


FIG. 2 Apparatus Assembly for Flame Distillation

mm in diameter, and having a peephole 25 mm in diameter centered about 32 mm below the ring support. The top of the shield shall be flanged to permit its being suspended from the ring support.

5.8 *Receivers*—Erlenmeyer flasks or beakers having a nominal capacity of 50 to 125 mL, and tared to the nearest 0.1 g.

5.9 Balance and Masses, accurate to 0.1 g.

5.10 *Thermometer*— An ASTM High Distillation Thermometer having a range from -2 to  $+400^{\circ}$ C and conforming to the requirements for Thermometer 8C as prescribed in Specification E 1.

#### 6. Preparation of Sample

6.1 Thoroughly stir or otherwise mix the sample immedi-

ately before removing the portion for testing, to ensure that such portion will be representative of the sample. If warming is necessary, take care to avoid loss of volatile material.

#### 7. Preparation of Apparatus

7.1 Assemble the apparatus as follows (see Fig. 2 and Fig. 3):

7.1.1 Suspend the burner chimney by its flange from the support ring, place the specified two sheets of flat wire gauze on the burner chimney, and place the flask shield on the upper sheet of gauze. In case the electric heater is used, attach the heater to a vertical support so that at the end of the distillation it can be lowered at least 150 mm. Place the upper refractory on the heater with the larger opening facing upwards. Set the flask shield on the upper refractory.

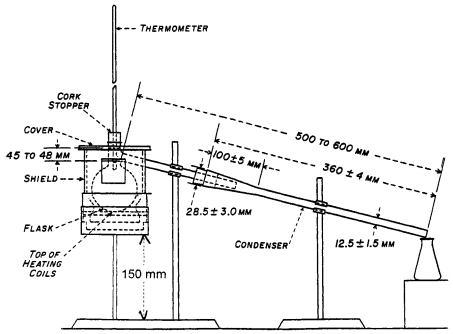


FIG. 3 Apparatus Assembly for Electric Heater Distillation

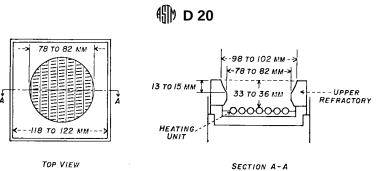


FIG. 4 Upper Part of Electric Heater

#### 8. Procedure

7.1.2 Inset the thermometer through a new and rolled select quality regular-length cork of suitable size. Position the thermometer and cork in the neck of the flask so that the cork fits tightly and the bottom of the cork is from 23 to 28 mm above the lowest point of the junction between the side-arm and the neck of the flask. Then adjust the thermometer in the cork so that the top of the bulb is level with the lowest point of the juncture between the side-arm and the neck of the flask. Align the stem of the thermometer on the axis of the bulb through the neck of the flask.

7.1.3 When using the flame distillation apparatus, place the flask in the flask shield with its bulb resting on the gauze. Position the burner so that it is directly beneath the point where the bulb of the flask contacts the gauze. If the electric heater is used, place the flask in the flask shield and support the flask so that its bottom is between 4 and 7 mm above the heating coils.

7.1.4 Connect the condenser tube to the side-arm of the flask with a tight cork joint, having the side-arm project 30 to 50 mm through the cork. The distance from the neck of the flask to the outlet end of the condenser tube shall be between 500 and 600 mm. Support the condenser tube in a position such that it is in alignment with the side-arm of the flask, and the thermometer is vertical. Place the shield cover over the flask shield around the neck of the flask.

8.1 Weigh the flask (along with the thermometer and cork) to the nearest 0.1 g and then weigh  $100.0 \pm 0.1$  g of the sample into the flask.

8.2 Apply heat to the flask so that the first drop of distillate (oil or water) falls from the end of the condenser tube in 5 to 15 min. Within 2 min after the first drop, adjust the rate of distillation, and subsequently maintain the rate, so that from 50 to 70 drops per min fall from the end of the condenser. Warm the condenser tube whenever necessary to prevent accumulation of solid distillates in the tube.

8.3 Collect the distillate fractions in tared receivers, changing receivers as the thermometer indicates the maximum temperature, corrected as described in 8.4, for each specified fraction. The following fractions are usually specified.

Up to 170°C
170 to 235°C
235 to 270°C
270 to 300°C
Residue at 300°C

Fractions at other temperatures, such as 170 to 200°C and 200 to 235°C, or 300 to 355°C and residue at 355°C, are sometimes required.

8.4 Do not change the position of the thermometer during

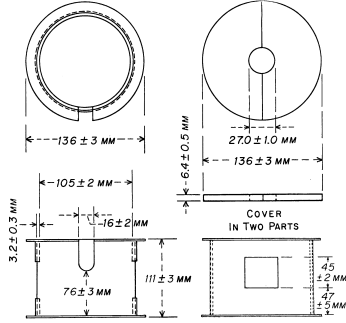


FIG. 5 Shield and Cover for Flame Distillation

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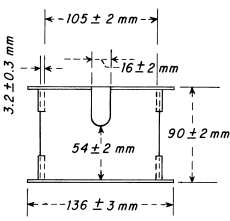


FIG. 6 Shield for Use with Electric Heater

the distillation. Make no correction for the emergent stem of the thermometer, but if the barometric pressure is outside the range of 756 to 765, adjust (but do not report) the temperature in accordance with Table 1.

8.5 When the maximum temperature specified for the test is indicated by the thermometer, immediately remove the flame and the flask-shield cover; or when the electric heater is used as a source of heat, immediately remove the flask shield cover and lower the heater at least 150 mm. Allow the apparatus to cool, for at least 5 min, or until no vapors are visible. Drain any oil remaining in the condenser tube into the receiver containing the last fraction.

8.6 Weigh the receivers containing the distillate fractions to the nearest 0.1 g. Weigh the flask (with the thermometer in place) and residue to the nearest 0.1 g.

8.7 Should the fraction to  $170^{\circ}$ C contain water, determine the water volume and calculate the new mass of oil distillate, assuming that 1 mL of water weighs 1 g. The amount of water contained in this fraction may be determined by either of the following methods.

TABLE 1 Adjustment of Distillation Test Temperatures for Barometric Pressure

	-					
Barometric Pressure, mm Hg	Fractionation Temperatures for Various Barometric Pressure Ranges, °C					
786 to 795	172	202	237	272	302	357
776 to 785	171	201	236	271	301	356
766 to 775	171	201	236	271	301	356
756 to 765	170	200	235	270	300	355
746 to 755	169	199	234	269	299	354
736 to 745	169	199	234	269	299	354
726 to 735	168	198	233	268	298	353
716 to 725	168	198	233	267	297	352
706 to 715	167	197	232	267	297	351
696 to 705	167	197	231	266	296	351
686 to 695	166	196	231	265	295	350
676 to 685	166	195	230	265	295	349
666 to 675	165	195	230	264	294	348
656 to 665	165	194	229	264	293	348
646 to 655	164	194	228	263	292	347
636 to 645	164	193	228	262	292	346
626 to 635	163	193	227	262	291	345
616 to 625	163	192	226	261	290	345
606 to 615	162	191	226	260	290	344
596 to 605	162	191	225	260	289	343

8.7.1 Transfer the fraction after weighing, to a tube or cylinder graduated in 0.1 mL. Rinse the receiver several times with toluene, adding the rinsings to the tube or cylinder containing the fraction or,

8.7.2 The fraction of 170°C may be collected in a tared graduated cylinder having a flared top. After weighing, add toluene which will result in a clear separation of the water and oil distillate.

8.8 If the residue from distillation is required for further testing, lower the thermometer until its bulb is in the liquid residue. If the residue is not completely fluid, heat it carefully to a temperature not exceeding  $150^{\circ}$ C by holding the bulb of the flask over a wire gauze heated by a gas burner or by immersion in a suitable bath whose temperature does not exceed  $150^{\circ}$ C. Incline the flask and rotate it so that the fluid residue will flow around the sides, and collect any oils that have condensed on the upper surfaces of the flask. Mix the contents of the flask until they are homogeneous. Allow the residue to cool to a temperature at which it can be readily poured from the flask without loss of volatile material and then pour it into the desired testing equipment or into a suitable receptacle. Cover the receptacle.

#### 9. Calculation

9.1 Convert the distillation results to a water-free basis, D, in %, as follows:

$$D = (F - W) \times [100/(100 - W)]$$
(1)

Second and Subsequent Fractions Including Residue:

$$D = F \times [100/(100 - W)]$$
(2)

where:

F = mass of the fraction or residue, g and

W = millilitres of water, expressed as grams, in the fraction distilling to 170°C.

## 10. Report

10.1 Report the following information:

10.1.1 The results of the distillation test as percentages to the nearest 0.1 %, based on the mass of water-free material. For road tars, it is customary to report the total distillate to a series of temperatures corresponding to the maxima specified for the fractions.

#### 11. Precision

11.1 The following criteria should be used for judging the acceptability of results at the 95 % probability level:

11.1.1 *Repeatability*— Duplicate results by the same operator should be considered suspect if the reported percentages differ by more than 1.2 absolute %.

11.1.2 *Reproducibility*— Results submitted by two laboratories should be considered suspect if the reported percentages differ by more than 3.9 absolute %.

Note 2—The precision limits given only apply to fractions having greater than 2.0% distillate.

#### 12. Keywords

12.1 distillation; road tar

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