Standard Test Method for Separation of Representative Aromatics and Nonaromatics Fractions of High-Boiling Oils by Elution Chromatography

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1. Scope

1.1 This test method covers the separation and determination of representative aromatics and nonaromatics fractions from hydrocarbon mixtures that boil between 232 and 538°C (450 and 1000°F). Alternative procedures are provided for the separation of 2 g or 10 g of hydrocarbon mixture.

NOTE 1—Some components may not be eluted from the chromatographic column for some types of samples under the conditions used in this method.

NOTE 2—Test Method D 2007 is an alternative method of separating high-boiling oils into polar compounds, aromatics, and saturates fractions.

1.2 An alternative procedure is provided to handle samples boiling below 232°C (450°F), but whose 5 % point is above 178°C (350°F) as determined by Test Method D 2887. This procedure is given in Annex A1.

1.3 The values stated in acceptable SI units are to be regarded as the standard. The values given in parentheses are provided for information purposes only.

1.4 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:
D 2007 Test Method for Characteristic Groups in Rubber Extender and Processing Oils and Other Petroleum-Derived Oils by the Clay-Gel Adsorption Chromatographic Method
D 2425 Test Method for Hydrocarbon Types in Middle Distillates by Mass Spectrometry
D 2786 Test Method for Hydrocarbon Types Analysis of Gas-Oil Saturate Fractions by High Ionizing Voltage Mass Spectrometry
D 2887 Test Method for Boiling Range Distribution of Petroleum Fractions by Gas Chromatography
D 3239 Test Method for Aromatic Types Analysis of Gas-Oil Aromatic Fractions by High Ionizing Voltage Mass Spectrometry

3. Terminology

3.1 Descriptions of Terms Specific to This Standard:
3.1.1 aromatics fraction—the portion of the sample desorbed with the polar eluants. The aromatics fraction may contain aromatics, condensed naphthenic-aromatics, aromatic olefins, and compounds containing sulfur, nitrogen, and oxygen atoms.
3.1.2 nonaromatics fraction—the portion of the sample eluted with n-pentane. The nonaromatics fraction is a mixture of paraffinic and naphthenic hydrocarbons if the sample is a straight-run material. If the sample is a cracked stock, the nonaromatics fraction will also contain aliphatic and cyclic olefins.

4. Summary of Test Method

4.1 A weighed amount of sample is charged to the top of a glass chromatographic column packed with activated bauxite and silica gel. n-Pentane is added to the column to elute the nonaromatics. When all of the nonaromatics are eluted, the aromatics fraction is eluted by additions of diethyl ether, chloroform, and ethyl alcohol.

4.2 The solvents are completely removed by evaporation, and the residues are weighed and calculated as the aromatics and nonaromatics fractions of the sample.

5. Significance and Use

5.1 The determination of compound types by mass spectrometry requires, in some instances, a preliminary separation of the petroleum sample into representative aromatics and nonaromatics fractions, as in Test Methods D 2425, D 2786, and D 3239. This test method provides a suitable separation technique for this application.

6. Apparatus

6.1 Chromatographic Columns, as shown in Fig. 1. Different chromatographic columns are provided for the analysis of 2 and 10-g samples.
6.2 Beakers, 100, 250, and 600-mL, inverted-rim type.
6.3 Steam Bath.
6.4 Electric Vibrator, for packing column.
6.5 Weighing Bottles or Erlenmeyer Flasks, 25 and 50 mL.

7. Reagents and Materials

This test method is under the jurisdiction of ASTM Committee D-2 on Petroleum Products and Lubricants and is the direct responsibility of Subcommittee D02.04 on Hydrocarbon Analysis.


1 Annual Book of ASTM Standards, Vol 05.02.

Beakers available from Kontes Glass Co., Vineland, NJ, by ordering "Anti-Creep" beakers and referring to Drawing No. 9413-A.
7.1 Purity of Reagents—Reagent grade chemicals shall be used in this test. Unless otherwise indicated, it is intended that all reagents shall conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society, where such specifications are available. Other grades may be used, provided it is first ascertained that the reagent is of sufficiently high purity to permit its use without lessening the accuracy of the determination.

7.2 Bauxite,\textsuperscript{5} 20 to 60-mesh. Before use, activate the bauxite by heating at 538°C (1000°F) for 16 h. Transfer the activated material to an airtight container while still hot and protect thereafter from atmospheric moisture.

7.3 Chloroform (Warning—Toxic. May be fatal if swallowed. See Annex A2.1.).

7.4 Cleaning Solution—Chromic-sulfuric acid (Warning—Causes severe burns. A recognized carcinogen, strong oxidizer, contact with organic material may cause fire. See Annex A2.2.).

7.5 Diethyl Ether, anhydrous, (Warning—Extremely flammable.). The ethyl ether used in this test method should be free of peroxides as determined by the procedure in “Reagent Chemical, American Chemical Society Specifications.”

7.6 Ethyl Alcohol, denatured, conforming to Formula 2B of the U.S. Bureau of Internal Revenue (Warning—Flammable.).

7.7 Pressuring Gas, dry air or nitrogen, delivered to the top of the column at a regulated gage pressure of 0 to 2 psi (13.8 kPa) (Warning—Compressed gas.).

7.8 n-Pentane, commercial grade, aromatic-free. Some samples of waxy stocks may not dissolve completely in n-pentane, in which case cyclohexane, commercial grade, aromatic-free, may be substituted for n-pentane (Warning—Extremely flammable liquid.).

7.9 Silica Gel,\textsuperscript{6} 100 to 200-mesh.


\textsuperscript{5} Bauxite available from Porocel Corp., Little Rock, AR.

\textsuperscript{6} Silica gel available from W.R. Grace and Co., Davison Chemical Div., Baltimore, MD 21203, by specifying Code 923.
8. Procedure

Note 3—The procedural details differ depending on the initial boiling point of the sample. If the 5% point is above 178°C (350°F), but below 232°C (450°F) use procedure described in Annex A1. If above 232°C continue as written depending on amount of sample to be analyzed. Instructions specific for 2-g samples are given in 8.4.1 to 8.4.13, and instructions specific for 10-g samples are given in 8.5.1 and 8.5.8.

8.1 Select the appropriate column, depending on whether 2 or 10 g of sample are to be analyzed. Clean the column with chromic-sulfuric acid. (Warning—Causes severe burns. See Annex A2.2.) followed by distilled or demineralized water, acetone, and dry air or nitrogen.

8.2 Introduce a small plug of glass wool into the column, pressing it firmly into the lower end to prevent the flow of silica gel from the column.

8.3 Clamp the column in a vertical position. Add small increments of silica gel, while vibrating the column along its length, until the tightly packed silica gel extends to the lower mark on the chromatographic column. Continue to vibrate the column and add bauxite until the bauxite layer extends to the upper mark on the chromatographic column. Vibrate the column for an additional 3 min after filling is completed.

8.4 If 2 g of sample are to be analyzed, continue as in 8.4.1, otherwise continue as in 8.5.

8.4.1 If the sample is viscous, warm it with intermittent mixing or shaking until it is completely fluid. Transfer a representative sample (approximately 2 g) to a 25-mL weighing bottle or flask. Determine the weight of the sample to the nearest 1 mg by weighing the flask before and after transfer. Add 10 mL of n-pentane. (Warning—Extremely flammable liquid.) to the flask and dissolve the sample. If the sample does not dissolve completely in cold n-pentane, warm it in warm water or over a steam bath. If the sample does not dissolve in warm n-pentane, take a fresh sample and substitute cyclohexane for the n-pentane.

8.4.2 Add 10 mL of n-pentane to the top of the column to prewet the adsorbent. When the liquid level reaches the top of the bauxite bed, transfer the sample solution from the weighing flask to the top of the column. Rinse the flask with three successive 3-mL washes of n-pentane. Add each wash to the top of the column. Then rinse the walls of the column bulb with two 3-mL portions of n-pentane, allowing the liquid level to reach the top of the bauxite bed before adding the next portion. Finally add 35 mL of n-pentane to the column bulb.

8.4.3 Place a 50-mL graduate beneath the column to collect the eluate. The elution rate should be approximately 1 mL/min.

8.4.4 When the n-pentane level reaches the top of the bauxite bed, add 80 mL of diethyl ether. (Warning—Extremely flammable.) Connect the pressuring gas to the top of the column and adjust the pressure to maintain an elution rate of 1 to 2 mL/min.

8.4.5 Collect 50 mL of n-pentane eluate in the graduate. Rinse the tip of the column with 1 to 2 mL of n-pentane, adding this to the 50 mL in the graduate (Note 5). Label the 50-mL graduate as n-pentane eluate.

8.4.6 When the ether level reaches the top of the bauxite bed, release the gas pressure and add 100 mL of chloroform. (Warning—Toxic. May be fatal if swallowed.) to the top of the column. Reconnect the gas pressuring system and continue the elution. When 80 mL of eluate have been collected in the graduate, rinse the column tip with 1 mL of ether and add the rinse to the 100-mL graduate. Change the receiver to a 250-mL graduate. Label the 100-mL graduate as ether-eluted fraction.

8.4.7 When the chloroform level reaches the top of the bauxite bed, release the gas pressure and add 75 mL of ethyl alcohol. (Warning—Flammable liquid.) Reconnect the gas pressuring system and continue the elution until the alcohol level reaches the top of the bauxite bed. Release the gas pressure. Rinse the column tip with 1 mL of chloroform adding this to the graduate. Label the 250-mL graduate as chloroform-alcohol-eluted fraction.

8.4.8 Weigh a 100-mL inverted-rim beaker to the nearest 1 mg. Quantitatively transfer the n-pentane eluate to this beaker and allow the n-pentane to evaporate at room temperature. Cyclohexane, if used as the elution solvent, is evaporated on a steam bath. Evaporation is accelerated in both cases by directing a controlled stream of dry nitrogen downward onto the surface of the liquid.

8.4.9 When all the solvent appears to be evaporated, stop the nitrogen flow, allow the beaker to come to room temperature, and dry the outside of the beaker to remove any condensed moisture. Reweigh the beaker to the nearest 1 mg.

8.4.10 Repeat the evaporation step for 5 min periods until the weight loss between successive evaporation is less than 0.20 mg. Heat from a steam bath is generally required during the final evaporation steps to remove completely the elution solvent. The weight of the residue in the beaker is the quantity of the nonaromatics fraction.

8.4.11 Weigh a 250-mL inverted-rim beaker to the nearest 1 mg. Quantitatively transfer the chloroform-alcohol-eluted fraction to this beaker and evaporate on a steam bath with a controlled stream of dry nitrogen directed downward onto the surface of the liquid. When the solvent is evaporated, remove the beaker from the steam bath, cool to room temperature, and add quantitatively the ether-eluted fraction. Evaporate the ether at room temperature as described in 8.4.8, 8.4.9, and 8.4.10. Determine the weight of the residue (aromatics fraction) to the nearest 1 mg.

8.4.12 The weight of the aromatics plus the nonaromatics fraction recovered must equal at least 95% of the sample charged. If 95% recovery is not obtained, repeat the test. Recoveries greater than 100% indicates incomplete removal.
of solvent or the condensation of moisture in the beakers.

8.4.13 Transfer the aromatics and nonaromatics fractions into suitable size vials for storage pending further analysis.

8.5 If 10 g of sample are to be analyzed, continue as in 8.5.1.

8.5.1 Warm the sample with intermittent mixing or shaking until it is completely fluid. Transfer a representative sample (approximately 8 to 10 g) to a 50-mL weighing bottle or flask. Determine the weight of the sample to the nearest 1 mg by weighing the flask before and after sample transfer. Add 20 mL of n-pentane to the flask and dissolve the sample. If the sample does not dissolve completely in cold n-pentane, warm it in warm water or over a steam bath. If the sample does not dissolve in warm n-pentane, take a fresh sample and substitute cyclohexane for n-pentane.

8.5.2 Add 45 mL of n-pentane to the top of the prepacked large column to prewet the adsorbent. When the n-pentane level reaches the top of the bauxite bed, transfer the sample solution from the weighing flask to the top of the column. Rinse the flask with three successive 3-mL washes of n-pentane. Add each wash to the top of the column. Then rinse the walls of the column bulb with two 3-mL portions of n-pentane, allowing the level of each portion to reach the top of the bauxite bed before adding the next portion. Finally add 70 mL of n-pentane to the column bulb.

8.5.3 Place a 200-mL graduate beneath the column to collect the eluate. The elution rate should be approximately 3 mL/min.

NOTE 7—Air or nitrogen pressure may be applied to the top of the column as necessary to accomplish and maintain a satisfactory elution rate. Three to five pounds of pressure generally is sufficient. If the correct pressure setting is known from previous runs, gas pressure can be applied after addition of the last increment of n-pentane. Otherwise, gas pressure should be applied when n-pentane begins to elute from the column and should be adjusted to give a flow rate of approximately 3 mL/min.

8.5.4 When the n-pentane level reaches the top of the bauxite bed, add 100 mL of diethyl ether. Connect the pressuring gas to the top of the column and adjust the pressure to maintain an elution rate of 3 to 5 mL/min.

8.5.5 Collect 130 mL of eluate in the graduate. Rinse the tip of the column with 1 to 2 mL of n-pentane, adding this to the 130 mL in the graduate. Change the receiver to a 100-mL graduate (Note 8). Label the 200-mL graduate as n-pentane eluted fraction.

NOTE 8—The n-pentane will have reached the absorbent bed before the required volume of eluate has been collected in the 200-mL receiver. Continue collection in this receiver after the addition of ether until the proper volume has been collected before changing to the 100-mL graduate.

8.5.6 When the ether level reaches the top of the bauxite bed, release the gas pressure and add 100 mL of chloroform to the top of the column. Reconnect the gas pressuring system and continue the elution until the alcohol level reaches the top of the bauxite bed. Release the gas pressure. Rinse the column tip with 1 mL of chloroform adding this to the graduate. Label the 500-mL graduate as chloroform-alcohol-eluted fraction.

8.5.7 When the chloroform level reaches the top of the bauxite bed, release the gas pressure and add 175 mL of ethyl alcohol. Reconnect the gas pressuring systems and continue the elution until the alcohol level reaches the top of the bauxite bed. Release the gas pressure. Rinse the column tip with 1 mL of chloroform adding this to the graduate. Label the 500-mL graduate as chloroform-alcohol-eluted fraction.

8.5.8 Weigh a 250-mL inverted rim beaker to the nearest 1 mg. Quantitatively transfer the n-pentane eluate to this beaker and evaporate the solvent on a steam bath. Evaporation can be accelerated by directing a controlled stream of dry nitrogen downward onto the surface of the liquid. Complete the workup of the nonaromatics fraction as described in 8.4.9 and 8.4.10.

8.5.9 Weigh a 600-mL inverted rim beaker to the nearest 1 mg (Note 9). Complete the workup of the aromatics fraction as described in 8.4.11, 8.4.12, and 8.4.13.

NOTE 9—The 600-mL inverted-rim beakers from some sources can exceed the capacity of the standard analytical balance, in which case a 250-mL inverted rim beaker can be used, and the chloroform-alcohol-eluted fraction evaporated in increments.

9. Calculation

9.1 Calculate the percentage of the aromatics fraction and the nonaromatics fraction as follows:

- Aromatics fraction, wt % = \[\frac{A}{A + B}\] \times 100 (1)
- Nonaromatics fraction, wt % = \[\frac{B}{A + B}\] \times 100 (2)

where:
- \(A\) = weight of aromatics fraction recovered, and
- \(B\) = weight of nonaromatics fraction recovered.

10. Precision and Bias

10.1 The following criteria should be used for judging the acceptability of results (95 % probability):

10.1.1 Repeatability—The difference between successive test results, obtained by the same operator with the same apparatus under constant operating conditions on identical test material would, in the long run, and in the normal and correct operation of the test method, exceed the following values only in one case in twenty: 0.4 weight % for 10 g of sample; and 1.4 weight % for 2 g of sample.

10.1.2 Reproducibility—The difference between two, single and independent results, obtained by different operators working in different laboratories on identical test material would, in the long run, and in the normal and correct operation of the test method, exceed the following values only in one case in twenty: 1.6 weight % for 10 g of sample; and 1.5 weight % for 2 g of sample.

NOTE 10—The procedure for analyzing 2 g of sample gives recoveries of aromatics fractions that are on average 0.35 weight % lower than the procedure for analyzing 10 g of sample.

10.2 Bias—Bias cannot be determined because there are no reference materials suitable for determining the bias in this test method.

NOTE 11—The precision of the procedure in Annex A1 has not been determined.
ANNEXES
(Mandatory Information)

A1. LOWER BOILING SAMPLE PROCEDURE

A1.1 Scope
   A1.1.1 This procedure covers the separation and determination of representative aromatics and nonaromatics from hydrocarbon mixtures whose 5 % boiling point is below 232°C (450°F), but above 178°C (350°F).

A1.2 Summary of Method
   A1.2.1 A Kuderna-Danish apparatus is used to evaporate solvents from the aromatic and nonaromatic fractions.

A1.3 Significance and Use
   A1.3.1 This procedure extends the range of this test method to separate the samples whose boiling range is specified in Test Methods D 2425, D 2786, and D 3239, all of which refer to this method to provide fractions for analyses.

A1.4 Apparatus
   A1.4.1 Kuderna-Danish Evaporator:
   A1.4.1.1 250-mL Flask, with top female standard taper 24/40 and bottom male standard taper 24/12 with glass hooks for retaining springs.
   A1.4.1.2 Macro Snyder Distillation Column, 3 ball, with male standard taper 24/40.
   A1.4.1.3 Conical Weighing Bottles, with female standard taper 24/12, 30-mL capacity with glass hooks for retaining springs.

A1.5 Procedure
   A1.5.1 The 10-g chromatographic column is used. Clean column with chromic-sulfuric acid, distilled or demineralized water, acetone, and dry air or nitrogen.
   A1.5.2 Same as 8.2.
   A1.5.3 Same as 8.3.
   A1.5.4 Same as 8.5.1
   A1.5.5 Same as 8.5.2.
   A1.5.6 Same as 8.5.3.
   A1.5.7 Same as 8.5.4.
   A1.5.8 Same as 8.5.5.
   A1.5.9 Same as 8.5.6.
   A1.5.10 Same as 8.5.7.
   A1.5.11 Weigh a 30-mL conical weighing bottle, to which a boiling chip is added, to the nearest 1 mg. Attach the weighing bottle to the 250-mL flask with the retaining springs. Quantitatively transfer the n-pentane eluate to the flask. Attach the Snyder distillation column to the flask and evaporate on a steam bath. Evaporation of most of the n-pentane is complete when balls in the Snyder distillation column stop moving.
   A1.5.12 Separate weighing bottle containing the concentrated pentane solution from the flask and weigh it after it has cooled to room temperature.
   A1.5.13 Gently swirl weighing bottle on the hot steam bath surface while directing a gentle stream of nitrogen to the bottle (Note A1.1). After 3 min, cool to room temperature and weigh.
   NOTE A1.1—Use nitrogen rate of approximately 100 mL/min. Do not direct nitrogen flow on liquid, but rather along inside of weighing bottle.
   A1.5.14 Repeat step A1.5.13 at 2-min intervals until weight loss between successive evaporations is less than 50 mg. The weight of the residue in the weighing bottle is the nonaromatic fraction.
   A1.5.15 Quantitatively transfer the chloroform-alcohol-eluted fraction from A1.5.10 to a 600-mL beaker and evaporate off the solvent on a steam bath using a stream of nitrogen to facilitate the evaporation rate. Allow to cool to room temperature.
   A1.5.16 Weigh a 30-mL conical weighing bottle and attach to Kuderna-Danish apparatus as described in A1.5.11. Transfer the ether-eluted fraction A1.5.9 into the beaker containing the residue from A1.5.15. Quantitatively transfer this mixture into the flask and evaporate on the steam bath as described in A1.5.11.
   A1.5.17 Complete the workup of the aromatic fraction as described in A1.5.12, A1.5.13, and A1.5.14.
   A1.5.18 Same as 8.4.12.
   A1.5.19 Same as 8.4.13.

A1.6 Calculation
   A1.6.1 Same as 9.1.

A1.7 Results
   A1.7.1 Two samples representing aromatic and nonaromatic concentrates from a middle distillate with an initial boiling point 149°C (300°F) were subjected to the evaporation procedure. Interlaboratory testing on the preceding samples was done by five laboratories, in duplicate, at two different times. Recoveries of 97 to 102 % with less than 3 % solvent remaining were obtained in the last study.

A1.8 Precision and Bias
   A1.8.1 There are no interlaboratory test data to establish a statistical statement of precision for the procedure in Annex A1 of Test Method D 2549.
   A1.8.2 There are no interlaboratory test data to establish a statistical statement of bias for the procedure in Annex A1 of Test method D 2549.
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