Standard Test Method for Benzene in Hydrocarbon Solvents by Gas Chromatography

This standard is issued under the fixed designation D 4367; 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 (e) indicates an editorial change since the last revision or reapproval.

1. Scope

1.1 This test method covers the determination by gas chromatography of benzene at levels from 0.01 to 1 volume % in hydrocarbon solvents.

Note 1—For benzene levels lower than 0.01 volume %, use Test Method D 6229.

1.2 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.

1.3 For hazard information and guidance, see the supplier’s Material Safety Data Sheet. For specific hazard statements, see Section 7.

2. Referenced Documents

2.1 ASTM Standards:

D 3606 Test Method for the Determination of Benzene and Toluene in Finished Motor and Aviation Gasoline by Gas Chromatography

D 6229 Test Method for Trace Benzene in Hydrocarbon Solvents by Capillary Gas Chromatography

E 260 Practice for Packed Column Gas Chromatography

E 300 Practice for Sampling Industrial Chemicals

3. Summary of Test Method

3.1 An internal standard, methyl ethyl ketone (MEK), is added to the material and then introduced into a gas chromatograph equipped with two columns connected in series. The specimen passes first through a column packed with the nonpolar phase, methyl silicone, which separates the components by boiling point. After octane has eluted, the flow through the nonpolar column is reversed, flushing out the components heavier than octane. The octane and lighter components then pass through a column with the highly polar phase, 1,2,3-tris(2-cyanoethoxy)propane, that separates the aromatic and nonaromatic compounds. The eluted components are detected by a conventional detector and recorded on a strip chart. The peak areas are measured and the concentration of each component is calculated by reference to the internal standard.

4. Significance and Use

4.1 Benzene is classed as a toxic and carcinogenic material. A knowledge of the concentration of this compound may be an aid in evaluating the possible health hazards to persons handling and using hydrocarbon solvents, but this test method is not intended to evaluate such hazards.

5. Apparatus

5.1 Chromatograph—Any gas chromatographic instrument that has a backflush system and flame ionization detector and that can be operated at the conditions given in Table 1. The detector-recorder combination must produce a 4-mm deflection for a 1-µL specimen containing 0.05 volume % MEK when operated at maximum sensitivity.

5.2 Columns, one 0.8-m (2.5-ft) length of 3.2-mm (¼-in.) outside diameter stainless steel tubing and one 4.6-m (15-ft) length of 3.2-mm (¼-in.) outside diameter stainless steel tubing.

5.3 Recorder, Strip Chart—Potentiometer with a full-scale deflection of 1 mV, a full-scale response time of 2 s or less, and a maximum noise level of ±0.3 % of full scale.

5.4 Microsyringe, 5-µL capacity.

5.5 Pipets, measuring 1 and 2 mL, graduated in 0.01 mL; 5, 10, and 20-mL capacity.

5.6 Flasks, volumetric, 25 and 100-mL capacity.

5.7 Vibrator, electric.

5.8 Vacuum Source.

5.9 Evaporator, vacuum, rotary.

5.10 Flask, boiling, round-bottom, short-neck, with 24/40 T joint, 500-mL capacity. Suitable for use with the evaporator (see 5.9).

5.11 Lamp, infrared.

5.12 Burets, automatic, with integral reservoir, 25-mL capacity.

Note 2—Suppliers of stationary phases and supports can be found in Research Report RR:D01-1038, available from ASTM Headquarters.

6. Reagents and Materials

6.1 Purity of Reagents—Reagent grade chemicals shall be...
7. Hazards

7.1 Many hydrocarbon solvents are flammable and hazardous; use special precautions when handling them. Of the reagents used in this procedure, methanol, chloroform, methylene chloride, acetone, methyl ethyl ketone, benzene (see 6.11.1), and n-nonane are hazardous.

7.2 Benzene is volatile and highly flammable. Exercise care to prevent accidental ignition. Benzene is also carcinogenic and toxic; acute or chronic poisoning may result from inhalation of benzene vapor, absorption of benzene through the skin, or drinking benzene.

8. Sampling

8.1 Take samples of solvents to be analyzed by this test method using the procedures described in Practice E 300.

9. Preparation of Columns

9.1 Column Packing Preparation—Prepare the two packing materials, one containing 10 % methyl silicone and the other 25 % TCEP, as follows:

9.1.1 Weigh 45 g of the acid-washed calcined diatomite support 60 to 80 mesh, into a 500-mL flask (see 5.10). Dissolve 5 g of the methyl silicone in approximately 50 mL of chloroform. (Warning—See Note 3.) Pour the methyl silicone–chloroform solution into the flask containing the support. Attach the flask to the evaporator (see 5.9), connect the vacuum, and start the motor. Turn on the infrared lamp and allow the packing to mix thoroughly until dry.

9.1.2 Weigh 75 g of acid-washed pink diatomaceous earth, 80 to 100 mesh, into a 500-mL flask (see 5.10). Dissolve 25 g of TCEP in 200 mL of methanol and pour into the flask containing the support. Attach the flask to the evaporator (see 5.9), connect the vacuum, and start the motor. Turn on the infrared lamp and allow the packing to mix thoroughly until dry, but do not heat the packing above 180°C.

9.2 Column Preparation:

9.2.1 Clean the stainless steel tubing as follows: Attach a metal funnel to one end of the steel tubing. Hold or mount the stainless steel tubing in an upright position and place a beaker under the outlet end of the tubing. Pour about 50 mL of methylene chloride into the funnel and allow it to drain through the steel tubing into the beaker. Repeat the washing with 50 mL of acetone. Remove the funnel and connect the steel tubing to an air line, by means of vinyl tubing. Remove all solvent from the steel tubing by blowing filtered, oil-free air through or applying a vacuum.

9.2.2 Pack the 0.8-m (2.5-ft) tubing (Column A) with the methyl silicone packing (see 9.1.1) and the 4.6-m (15-ft) tubing (Column B) with the TCEP packing (see 9.1.2) as follows: Preform Columns A and B separately to fit the chromatographic instrument. Close one end of each tubing with a small, glass wool plug and connect this end to a vacuum source by means of a glass-wool packed tube. To the other end connect a small polyethylene funnel by means of a short length of vinyl tubing. Start the vacuum and pour the appropriate packing into the funnel until the column is full. While filling each column, vibrate the column with the electric vibrator to settle the packing. Remove the funnel and shut off the vacuum source. Remove the top 6 mm (¼ in.) of packing and insert a glass wool plug in this end of the column.

9.3 Prepacked columns conforming to specifications listed in Table 1, and in 5.2, 9.1, and 9.2 may be obtained from any reputable chromatography supply company.
10. Preparation of Chromatographic Apparatus and Establishment of Conditions

10.1 Column Conditioning—Join Columns A and B as shown in Fig. 1. Connect the inlet of Column A to the injection port of the chromatograph. Pass helium gas through the column at approximately 40 mL/min. Condition the columns in accordance with the following time-temperature schedule.

<table>
<thead>
<tr>
<th>Temperature, °C</th>
<th>Time, h</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>0.5</td>
</tr>
<tr>
<td>100</td>
<td>0.5</td>
</tr>
<tr>
<td>150</td>
<td>1</td>
</tr>
<tr>
<td>170</td>
<td>3</td>
</tr>
</tbody>
</table>

10.2 Connect the outlet of Column B to the detector port. Adjust the operating conditions to those listed in Table 1, but do not turn on the detector circuits. Check the system for leaks.

10.3 Adjust the flow rate as follows:

10.3.1 Set the value in the forward flow mode (Fig. 2(a)) and adjust Flow Controller A to give the required flow rate (Table 1). Measure the flow rate at the detector vent, specimen side.

10.3.2 Set the valve in the backflush position (Fig. 2(b)) and measure the flow rate at the detector vent, specimen side. If the rate has changed, adjust Flow Controller B to obtain the required flow rate to within ±1 mL/min.

10.3.3 Turn on the detector circuit. Change the valve from forward flow to the backflush position several times and observe the baseline. There should be no baseline shift or drift after the initial peak resulting from the pressure surge with the valve change. If there is a baseline shift, slightly increase or decrease flow with Controller B to balance the baseline. (A persistent drift indicates leaks somewhere in the system.)

10.4 Determine time before backflushing, which varies for each column system and must be determined experimentally as follows:

10.4.1 Prepare a mixture of 5 volume % isooctane in n-nonane. Using the injection technique described in 11.3 and with the system in the forward flow mode, inject 1 µL of the isooctane–n-nonane mixture. Allow the chromatogram to run until the n-nonane has eluted and the recorder pen has returned to baseline. Measure the time in seconds from the injection until the recorder pen returns to baseline between the isooctane and n-nonane peaks. At this point all of the isooctane but essentially none of the n-nonane should have eluted. One half of the measured time approximates the time to backflush and should be from 30 to 120 s.

10.4.2 Repeat the run, including the injection, but switching the system to the backflush mode at the determined backflush time. This should result in a chromatogram of isooctane with little or no n-nonane evident.

10.4.3 If necessary, make additional runs, adjusting the time to backflush until a chromatogram of all the isooctane and little or none of the n-nonane is obtained. This established backflush time, including the actual valve operations, must be used in all subsequent calibrations and analyses.

11. Calibration and Standardization

11.1 Standard Solutions—Prepare seven standard solutions covering the range of 0 to 1 volume % benzene as follows: For each standard, measure the volume of benzene listed below into a 100-mL volumetric flask. Dilute to volume with isooctane, with all components and glassware at normal room temperature, and mix thoroughly.

<table>
<thead>
<tr>
<th>Volume %</th>
<th>mL</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td>0.25</td>
<td>0.25</td>
</tr>
<tr>
<td>0.10</td>
<td>0.10</td>
</tr>
<tr>
<td>0.05</td>
<td>0.05</td>
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<tr>
<td>0.01</td>
<td>0.01</td>
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<tr>
<td>0.005</td>
<td>0.005</td>
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</tbody>
</table>

11.2 Calibration Solutions—Accurately measure 0.5 mL of MEK into a 100-mL volumetric flask, fill to the mark with the
first standard solution (see 11.1), and mix thoroughly. Repeat with each of the other standard solutions.

11.3 Chromatographic Analysis—Using the conditions established in 10.3 and 10.4, chromatograph each of the calibration solutions after injecting them as follows: Flush the 5-µL microsyringe at least three times with the calibration solution and then fill with about 3 µL, avoiding inclusion of air bubbles in the syringe. Slowly eject the material until 1.0 µL remains in the syringe. Wipe the needle with a tissue and draw back the plunger to admit 1 µL of air into the syringe. Insert the needle of the syringe into the septum cap of the chromatograph and push through the septum until the barrel of the syringe is resting against the septum cap; then rapidly push the plunger to the hilt and immediately withdraw the needle from the injection port.

NOTE 4—This injection technique is necessary to obtain sharp symmetrical peaks.

11.4 Calibration—Measure the areas of the benzene and of MEK peaks by conventional means (Note 5). Calculate the ratio of the benzene peak area to the MEK peak area. Plot the concentration of benzene versus the ratio as in Fig. 3. The calibration must be done to ensure that the entire chromatographic system is operating properly and that the concentration of any one component has not exceeded the linear response range of any part of the system—column, detector, integrator, and other components. The calibration plot should be linear (Note 6). Determine the retention times for each component for future identification.

NOTE 5—The precision statement in Section 15 was developed from results obtained using electronic integrators or on-line computers. The precision statement may not apply if other methods of integration or peak area measurement are used.

NOTE 6—If the calibration is linear, a least-squares calculation may be performed to obtain a calibration factor. The precision statement in Section 15 was developed from results obtained from calibration plots and may not apply if calibration factors are used.

12. Procedure

12.1 Test Solution—Accurately measure 0.5 mL of MEK into a 100-mL volumetric flask. Fill to the mark with the material under test and mix well.

12.2 Chromatograph a specimen from the test solution using the conditions established in 10.3 and 10.4 and the injection technique described in 11.3.

NOTE 7—The valves must be turned to the backflush mode at the established backflush time so that undesirable components do not enter Column B.

12.3 Identify on the chromatogram the benzene and the internal standard MEK peaks from the retention times of the standards.

NOTE 8—The order of elution is nonaromatic hydrocarbons, benzene, MEK, and toluene when using the specified columns, as shown in Fig. 4.
12.4 Measure the areas under the benzene peak and under the MEK peak by conventional methods.

13. Calculation

13.1 Calculate the ratio of peak area of benzene to the peak area of MEK. Read from the calibration curve the volume % of benzene corresponding to the calculated peak ratio.

13.2 If the results are desired on a weight basis, convert to weight % as follows:

\[ \text{Benzene, weight \%} = \left( \frac{V}{D} \right) \times 0.8844 \]  

(1)

where:

- \( V \) = benzene, volume %, and
- \( D \) = relative density of sample at 15.6/15.6°C (60/60°F).

14. Report

14.1 Report the following information: benzene content in volume or weight % to the nearest 0.005 %.

15. Precision and Bias

15.1 Precision—The precision statements are based on an interlaboratory study in which analysts in each of six laboratories analyzed seven hydrocarbon solvent samples, including heptane, VM&P naphtha, mineral spirits, toluene, and aromatic solvent 100 on two different days. To each solvent, initially containing essentially no benzene, 0.1 to 0.5 volume % benzene was added. The within-laboratory standard deviation was found to be 0.0094 % absolute with 42 df and the between-laboratory standard deviation was 0.022 % absolute with 49 df. Based on these standard deviations, the following criteria should be used for judging the acceptability of results at the 95 % confidence level:

15.1.1 Repeatability—Two results, each the mean of duplicates, obtained by the same operator on different days should be considered suspect if they differ by more than 0.027 % absolute.

15.1.2 Reproducibility—Two results, each the mean of duplicates, obtained by operators in different laboratories should be considered suspect if they differ by more than 0.063 % absolute.

15.2 Bias—Bias can not be determined for this test method because there is no available material having an accepted reference value.

16. Keywords

16.1 benzene content; gas chromatography; hydrocarbon solvents

**SUMMARY OF CHANGES**

Committee D-1 has identified the location of selected changes to this standard since the last date of issue that may impact the use of this standard.

(1) References to specific suppliers of GC column stationary phases and supports have been removed from this test method and added to Research Report RR:D01-1038. A note regarding

Test Method D 6229 has been added to 1.1, and Test Method D 6229 has been added to 2.1.

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