

Designation: D3606-06<sup>€1</sup> Designation: D 3606 - 07

An American National Standard

# Standard Test Method for Determination of Benzene and Toluene in Finished Motor and Aviation Gasoline by Gas Chromatography<sup>1</sup>

This standard is issued under the fixed designation D 3606; 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.

e¹Note—Updated 1.4 editorially in November 2006.

## 1. Scope\*

- 1.1 This test method covers the determination of benzene and toluene in finished motor and aviation gasolines by gas chromatography.
- 1.2 Benzene can be determined between the levels of 0.1 and 5 volume % and toluene can be determined between the levels of 2 and 20 volume %.
- 1.3 The precision for this test method was determined using conventional gasoline as well as gasolines containing oxygenates (ethers such as methyl *tert*-butyl ether, ethyl *tert*-butyl ether and *tert*-amyl methyl ether).
  - 1.4Methanol may cause interference.
- 1.4 Methanol may cause interference. Appendix X1 provides an option for modifying the test method for analyzing samples containing ethanol.
  - 1.5 The values stated in SI units are to be regarded as the standard. The values given in parentheses are for information only.
- 1.6 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.

# 2. Referenced Documents (https://standa

2.1 ASTM Standards: <sup>2</sup>

D 4057 Practice for Manual Sampling of Petroleum and Petroleum Products

E 694 Specification for Laboratory Glass Volumetric Apparatus

E 969 Specification for Glass Volumetric (Transfer) Pipets

E 1044 Specification for Glass Serological Pipets (General Purpose and Kahn)

E 1293 Specification for Glass Measuring Pipets

# 3. Summary of Test Method

3.1 An internal standard, methyl ethyl ketone (MEK), is added to the sample which is then introduced into a gas chromatograph equipped with two columns connected in series. The sample passes first through a column packed with a nonpolar phase such as dimethylpolysiloxane (8.1.1) which separates the components according to 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 packed with a highly polar phase such as 1,2,3-tris(2-cyanoethoxy) propane (8.1.2) which separates the aromatic and nonaromatic compounds. The eluted components are detected by a thermal conductivity detector. The detector response is recorded, the peak areas are measured, and the concentration of each component is calculated with reference to the internal standard.

# 4. Significance and Use

4.1 Benzene is classed as a toxic material. A knowledge of the concentration of this compound can be an aid in evaluating the possible health hazard to persons handling and using the gasoline. This test method is not intended to evaluate such hazards.

<sup>&</sup>lt;sup>1</sup> This test method is under the jurisdiction of ASTM Committee D02 on Petroleum Products and Lubricants and is the direct responsibility of Subcommittee D02.04.0L on Gas Chromatography Methods.

Current edition approved May 1, 2006. Published June 2006. Originally approved in 1977. Last previous edition approved in 2004 as D3606-04a.

Current edition approved Nov. 1, 2007. Published January 2008. Originally approved in 1977. Last previous edition approved in 2006 as D 3606-06<sup>c1</sup>

<sup>&</sup>lt;sup>2</sup> 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.



## 5. Apparatus

- 5.1 *Chromatograph*—Any chromatographic instrument that has a backflush system and thermal conductivity detector, and that can be operated at the conditions given in Table 1, can be employed. Two backflush systems are shown. Fig. 1 is a pressure system and Fig. 2 is a switching valve system. Either one can be used.
  - 5.2 Columns:
- 5.2.1 Column A—One 0.8-m (2.5-ft) by 3.2-mm (1/8-in.) outside diameter stainless steel column packed with 10 mass % dimethylpolysiloxane (for example, OV-101) on Chromosorb W, 60 to 80 mesh.
- 5.2.2 *Column B*—One 4.6-m (15-ft) by 3.2-mm outside diameter stainless steel column packed with 20 mass % TCEP on Chromosorb P, 80 to 100 mesh.
- 5.3 Recorder, a strip chart recorder. An electronic integrating device or a computer capable of graphical presentation of the chromatogram. The electronic integrating device or computer must be capable of measuring 0.1 volume % MEK with satisfactory signal-to-noise. If a strip chart recorder is to be used, a 0 to 1-mV range recording potentiometer with a response time of 2 s or less and a maximum noise level of  $\pm 0.3$  % of full scale is recommended. The detector strip chart recorder combination must produce a 4-mm deflection for a 2- $\mu$ L sample containing 0.1 volume % MEK when operated at maximum sensitivity.
  - 5.4 Microsyringe, 5-µL capacity.
  - 5.5 Volumetric Pipets, Class A, 0.5, 1, 5, 10, 15, and 20-mL capacities (see Specification E 694 and E 969).
- 5.6 Measuring Pipets, 1 and 2-mL capacities calibrated in 0.01 mL; 5-mL calibrated in 0.1-mL, for use in dispensing volumes of benzene and toluene not covered by the volumetric pipets (see Specification E 1044 and E 1293) during preparation of standard samples (see 11.1).

Note 1—Other volume dispensing equipment capable of delivering the specified volumes within the stated tolerance limits may be used as an alternative to the requirements stated in 5.5 and 5.6.

- 5.7 Flasks, volumetric, 25 and 100-mL capacity.
- 5.8 Vibrator, electric.
- 5.9 Vacuum Source.
- 5.10 Evaporator, vacuum, rotary.
- 5.11 *Flask*, boiling, round-bottom, short-neck, with <sup>24</sup>/<sub>40</sub> standard taper joint, 500-mL capacity. Suitable for use with evaporator (5.10).
  - 5.12 Lamp, infrared.
  - 5.13 Burets, automatic, with integral reservoir, 25-mL capacity.

#### 6. Materials

- 6.1 Carrier Gas—Helium, 99.99 % pure. (Warning—Compressed gas under high pressure.)
- 6.2 Support—Crushed firebrick, acid-washed, 60 to 80-mesh and 80 to 100-mesh.

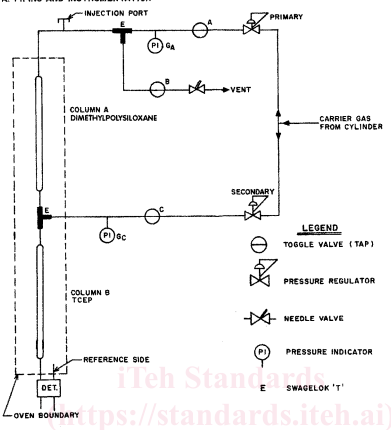
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**TABLE 1 Instrument Parameters** 

Detector	thermal conductivity
Columns:	two, stainless steel
Length, m	(A) 0.8; (B) 4.6
Outside diameter, mm	3.2
Stationary phase	(A) dimethylpolysiloxane, 10 mass %
	(B) TCEP, 20 mass %
Support	(A) Chromosorb W, 60 to 80-mesh
	(B) Chromosorb P, 80 to 100-mesh
Reference column	Any column or restriction may be
	used.
Temperature:	
Sample inlet system, °C	200
Detector, °C	200
Column, °C	145
Carrier Gas:	helium
Linear Gas Rate, cm/s	6
Volume flow rate, cm <sup>3</sup> /min	approximately 30
Column head pressure, kPa (psi)	approximately 200 (30)
Recorder range, mV	0 to 1
Chart speed, cm/min	1
Sample size, µL	2
Total cycle time, min	8
Backflush, min	approximately 0.75 <sup>A</sup>

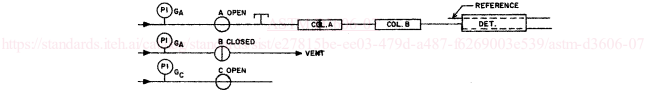
<sup>&</sup>lt;sup>A</sup> This back flush time must be determined for each column system.

#### A. PIPING AND INSTRUMENTATION

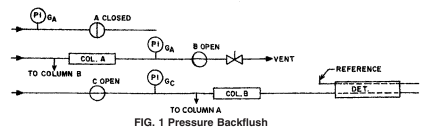


## B. FLOW SWITCHING SYSTEM

## I. FORWARD FLOW



## 2. BACKFLUSH

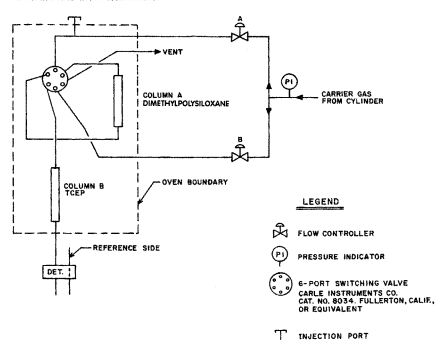


- 6.3 Liquid Phases—1,2,3-Tris(2-cyanoethoxy) propane (TCEP) and methyl silicone.<sup>3</sup>
- 6.4 Solvents:
- 6.4.1 *Methanol*, reagent grade. (**Warning**—Flammable. Vapor harmful. Can be fatal or cause blindness if swallowed or inhaled.)
  - 6.4.2 *Chloroform*, reagent grade. (Warning—Can be fatal if swallowed. Harmful if inhaled.)
- 6.4.3 *Methylene Chloride*, for cleaning columns. (**Warning**—Harmful if inhaled. High concentrations can cause unconsciousness or death.)

<sup>&</sup>lt;sup>3</sup> Packed column liquid phases such as OV 101 are considered to be of the dimethylpolysiloxane type. Other equivalent phases can also be used. Consult with the column manufacturer or phase supplier for information.

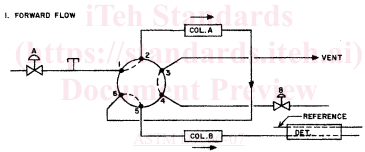


#### A. PIPING AND INSTRUMENTATION



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#### B. FLOW SWITCHING SYSTEM



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## 2. BACKFLUSH

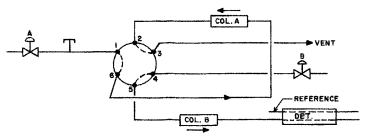


FIG. 2 Valve Backflush

- 6.4.4 Acetone, for cleaning columns. (Warning—Extremely flammable. Vapors can cause flash fires.)
- 6.5 Internal Standard:
- 6.5.1 Methyl Ethyl Ketone (MEK), 99.5% minimum purity. (Warning—Flammable. Vapor can be harmful.)
- 6.6 Calibration Standards:
- 6.6.1 *Benzene*, 99 + mol %. (**Warning**—Poison. Carcinogen. Harmful or fatal if swallowed. Extremely flammable. Vapors can cause flash fires.)
  - 6.6.2 *Isooctane* (2,2,4-trimethyl pentane), 99<sup>+</sup> mol %. (Warning —Extremely flammable. Harmful if inhaled.)
  - 6.6.3 *Toluene*, (Warning—Flammable. Vapor harmful.)
  - 6.6.4 *n-Nonane*, 99 + mol %. (Warning—Flammable. Vapor harmful.)

## 7. Sampling

7.1 Gasoline—(Warning —Extremely flammable. Vapors harmful if inhaled.) Samples to be analyzed by this test method shall



be obtained using the procedures outlined in Practice D 4057.

## 8. Preparation of Column Packings

- 8.1 Prepare two packing materials (one packing material consists of 10 mass % dimethylpolysiloxane on Chromosorb W; the other, 20 mass % TCEP on Chromosorb P) in accordance with the following procedures:
- 8.1.1 Dimethylpolysiloxane Packing—Weigh 45 g of the Chromosorb W, 60 to 80 mesh and pour into the 500-mL flask (5.11). Dissolve 5 g of the dimethylpolysiloxane in approximately 50 mL of chloroform. (**Warning**—Can be fatal if swallowed. Harmful if inhaled.) Pour the methyl silicone-chloroform solution into the flask containing the Chromosorb W. Attach the flask to the evaporator (5.10), connect the vacuum, and start the motor. Turn on the infrared lamp and allow the packing to mix thoroughly until dry.
- 8.1.2 1,2,3-Tris(2-cyanoethoxy) Propane (TCEP) Packing —Weigh 80 g of Chromosorb P, 80 to 100 mesh and pour into the 500-mL flask (5.11). Dissolve 20 g of TCEP in 200 mL of methanol and pour into the flask containing the Chromosorb P. Attach the flask to the evaporator (5.10), connect the vacuum, and start the motor. Turn on the infrared lamp and allow the packing to mix thoroughly until dry. (Do not heat the packing over 180°C.)

## 9. Preparation of Column

- 9.1 Cleaning Column—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 drain beaker under the outlet end of the tubing. Pour about 50 mL of methylene chloride (Warning—Harmful if inhaled. High concentrations can cause unconsciousness or death) into the funnel and allow it to drain through the steel tubing and into the drain beaker. Repeat the washing procedure with 50 mL of acetone. (Warning—Harmful if inhaled. High concentrations can cause unconsciousness or death.) Remove the funnel and attach the steel tubing to an air line, using vinyl tubing to make the connection. Remove all solvent from the steel tubing by blowing filtered, oil-free air through or pulling a vacuum.
- 9.2 Packing Columns—Preform Columns A and B separately to fit the chromatograph. Pack the 0.8-m tubing (Column A) with the dimethylpolysiloxane packing (8.1.1) and the 4.6-m tubing (Column B) with the TCEP packing (8.1.2) using the following procedure. 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 (1/4-in.) of packing and insert a glass wool plug in this end of the column.

# 10. Configuration of Apparatus and Establishment of Conditions

10.1 Conditioning Column—Install Columns A and B as shown in Fig. 1 or Fig. 2 in accordance with the system preferred (5.1). Do not connect the exit end of Column B to the detector until the columns have been conditioned. Pass helium gas through the column at approximately 40 cm<sup>3</sup>/min. Condition the column at the listed temperatures for the specified time periods.

Temperature, °C Standards/Sist/eZ / 813 be-ee03-479d-a48 /	Hours at Temperature
50	1/2
100	1/2
150	1
170	3

- 10.2 Assembly—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 systems for leaks.
  - 10.3 Flow Rate Adjustment:
  - 10.3.1 Column System Setup for Pressure Backflushing (Fig. 1):
- 10.3.1.1 Open Tap A and B and close C; set the primary pressure regulator to give the desired flow (Table 1) through the column system (at an approximate gage pressure of 205 kPa (30 psi)). Measure the flow rate at the detector vent, sample side. Observe the pressure on gage  $G_C$ .
- 10.3.1.2 Close Tap A and open B and C. The pressure reading on gage  $G_A$  should fall to zero immediately. If not, open the needle valve until the pressure falls to zero.
- 10.3.1.3 Close Tap B. Adjust the secondary pressure regulator until the reading of gage  $G_C$  is 3.5 to 7 kPa (0.5 to 1 psi) higher than observed in 10.3.1.1.
- 10.3.1.4 Open Tap B and adjust the backflush vent control needle valve until the pressure recorded on  $G_A$  approximates a gage pressure of 14 to 28 kPa (2 to 4 psi).
  - 10.3.1.5 Forward Flow—Open Taps A and C and close Tap B (Fig. 1 B1).
- 10.3.1.6 Backflush—Close Tap A and open Tap B. (There should be no baseline shift on switching from  $forward\ flow$  to backflush. If there is a baseline shift increase the secondary pressure slightly.) (Fig. 1)
  - 10.3.2 Column System Setup for Valve Backflushing (Fig. 2):
- 10.3.2.1 Set the valve in the *forward flow* mode (Fig. 2 B1), and adjust flow control A to give the desired flow (Table 1). Measure the flow rate at the detector vent, sample side.
  - 10.3.2.2 Set the valve in the backflush position (Fig. 2 B2), measure the flow rate at the detector vent, sample side. If the flow

has changed, adjust flow control B to obtain the correct flow. (Flows should match to within  $\pm 1 \text{ cm}^3/\text{min}$ ).

- 10.3.2.3 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 valve kick that results from the pressure surge. If there is a baseline shift, increase or decrease flow control *B* slightly to balance the baseline. (A persistent drift could indicate leaks somewhere in the system.)
- 10.4 Determine Time to Backflush—The time to backflush will vary for each column system and must be determined experimentally as follows. Prepare a mixture of 5 volume % isooctane in n-nonane. Using the injection technique described in 11.4 and with the preferred system (10.3) 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 detector signal has returned to baseline. Measure the time in seconds, from the injection until the detector signal 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 time determined should approximate the "time to backflush" and should be from 30 to 60 s. Repeat the run, including the injection, but switching the system to the backflush mode at the predetermined "time to backflush." This should result in a chromatogram of isooctane with little or no n-nonane visible. If necessary, make additional runs, adjusting the "time to backflush" until this condition of all the isooctane and little or no n-nonane is attained. The "time to backflush" so established, including the actual valve operations, must be used in all subsequent calibrations and analyses.

### 11. Calibration and Standardization

11.1 Standard Samples—Prepare seven standard samples covering the range 0 to 5 volume % benzene and 0 to 20 volume % toluene as follows: For each standard, measure the volume of benzene and of toluene listed below into a 100-mL volumetric flask. Dilute to volume with *iso*octane (2,2,4-trimethyl pentane), with all components and glassware at ambient temperature.

Benzene		Toluene	
Volume %	mL	Volume %	mL
5	5.0	20	20.0
2.5	2.5	15	15.0
1.25	1.25	10	10.0
0.67	0.67	5	5.0
0.33	0.33	12 n n 2 r 2.5 C	2.50
0.12	0.12		1.0
0.06	0.06	0.5	0.50

- 11.2 Calibration Blends—Accurately measure 1.0 mL of MEK into a 25-mL volumetric flask, and fill to the mark with the first standard sample (11.1). Continue doing this until all blends have been prepared.
  - Note 2—Commercially prepared calibration standards may be used, including those that are pre-mixed with the MEK internal standard.
- 11.3 *Chromatographic Analysis*—Chromatograph each of the calibration blends using the conditions established in 10.4 using the following injection technique:
  - 11.4 Injection of Sample:
- 11.4.1 Use of an automatic liquid sample injection system is highly recommended. If manual injections are to be made, the injection technique in 11.4.2 is necessary so that sharp symmetrical peaks will be obtained.
- 11.4.2 Flush the 5- $\mu$ L microsyringe at least three times with the sample mixture and then fill with about 3 $\mu$ L of the sample. (Avoid including any air bubbles in the syringe.) Slowly eject the sample until 2.0  $\mu$ L remains in the syringe; wipe the needle with tissue and draw back the plunger to admit 1 to 2  $\mu$ L of air into the syringe. Insert the needle of the syringe through the septum cap of the chromatograph and push until the barrel of the syringe is resting against the septum cap; then push the plunger to the hilt and remove the syringe immediately from the chromatograph.
- 11.5 *Calibration*—Measure the area of both aromatic peaks and of the internal standard peak as directed in 12.4. Calculate the ratio of the benzene peak area to the MEK peak area. Plot the concentration of benzene versus the ratio. Make the same calculation and plot for toluene. See Fig. 3 for an example. This 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 should be linear.
- Note 3—Calibrations using computer-based chromatography systems are an acceptable alternative to the calibration procedure specified in 11.5.
- Note 4—If the calibration has been shown to be linear, a least squares calculation may be performed to calculate a calibration factor. The precision statement in Section 15 was developed from data obtained from calibration plots and may not apply if calibration factors are used.

### 12. Procedure

- 12.1 Preparation of Sample—Accurately measure 1.0 mL of MEK into a 25-mL volumetric flask. Fill to the mark with the sample to be tested and mix well.
- 12.2 *Chromatographic Analysis*—Chromatograph the sample, using the conditions established in 10.4 "time to backflush" and the injection technique described in 11.4. The valves must be turned to *backflush* mode at the time determined in 10.4 so that undesirable components do not enter Column B.
- 12.3 *Interpretation of Chromatogram* Identify on the chromatogram the benzene, toluene, and the internal standard MEK peaks from the retention times of the standards.