



Standard Test Method for Analysis of Styrene by Capillary Gas Chromatography¹

This standard is issued under the fixed designation D 5135; 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 (ε) indicates an editorial change since the last revision or reapproval.

1. Scope *

1.1 This test method covers the determination of the impurities typically found in styrene and uses those values to determine the purity of styrene by difference by gas chromatography. Typical impurities in commercial styrene are found in Table 1.

1.2 This test method is applicable to impurity concentrations in the range of 0.001 to 1.00 % and for styrene purities of 99 % or higher when using the internal standard procedure.

1.3 The following applies to all specified limits in this standard: for purposes of determining conformance with this standard, an observed value or a calculated value shall be rounded off “to the nearest unit” in the last right-hand digit used in expressing the specification limit, in accordance with the rounding-off method of Practice E 29.

1.4 *This standard does not purport to address all 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 a specific hazard statement, see Section 9.

2. Referenced Documents

2.1 ASTM Standards:

D 3437 Practice for Sampling and Handling Liquid Cyclic Products²

D 4307 Practice for Preparation of Liquid Blends for Use as Analytical Standards³

D 4790 Terminology of Aromatic Hydrocarbons and Related Chemicals²

E 29 Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications⁴

E 355 Practice for Gas Chromatography Terms and Relationships⁵

E 691 Practice for Conducting an Interlaboratory Test Pro-

TABLE 1 Impurities Known or Suggested to be Present in Commercial Styrene

Ethylbenzene
<i>o</i> -xylene
<i>m</i> -xylene
<i>p</i> -xylene
Isopropylbenzene
<i>n</i> -propylbenzene
<i>m</i> -ethyltoluene
<i>p</i> -ethyltoluene
α -methylstyrene
<i>m</i> -vinyltoluene
<i>p</i> -vinyltoluene
Phenylacetylene

gram to Determine the Precision of Test Methods⁴

E 1510 Practice for Installing Fused Silica Open Tubular Capillary Columns in Gas Chromatographs⁵

2.2 Other Document

OSHA Regulations—29CFR paragraphs 1910.1000 and 1910.1200⁶

3. Terminology

3.1 See Terminology D 4790 for definitions of terms used in this test method.

4. Summary of Test Method

4.1 In this test method, the chromatogram peak area for each impurity is compared to the peak area of the internal standard (*n*-heptane or other suitable known) added to the sample. From the response factors of these impurities relative to that of the internal standard and the amount of internal standard added, the concentration of the impurities are calculated. The styrene content is obtained by subtracting the total amount of all impurities from 100.00.

5. Significance and Use

5.1 This test method is designed to obtain styrene purity on the basis of impurities normally present in styrene and may be used for final product inspections and process control.

5.2 This test method will detect the impurities shown in Table 1, non-aromatic hydrocarbons containing ten carbons or

¹ This test method is under the jurisdiction of ASTM Committee D16 on Aromatic Hydrocarbons and Related Chemicals and is the direct responsibility of Subcommittee D16.07 on Styrene, Ethylbenzene, and C₉ and C₁₀ Aromatic Hydrocarbons.

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² Annual Book of ASTM Standards, Vol 06.04.

³ Annual Book of ASTM Standards, Vol 05.02.

⁴ Annual Book of ASTM Standards, Vol 14.02.

⁵ Annual Book of ASTM Standards, Vol 03.06.

⁶ Available from U.S. Government Printing Office, Superintendent of Documents, 732 North Capitol St., NW, Mail Stop SDE, Washington, DC 20401.

less, and others where specific impurity standards are available. Absolute purity cannot be determined if unknown impurities are present.

6. Interferences

6.1 The internal standard chosen must be satisfactorily resolved from any impurity as well as the product peak. A peak will be satisfactorily resolved from a neighboring peak if the distance from the valley to the baseline between the two peaks is not greater than 50 % of the peak height of the smaller of the two peaks.

7. Apparatus

7.1 *Gas Chromatograph*—Any instrument having a flame ionization detector that can be operated at the conditions given in Table 2. The system shall have sufficient sensitivity to obtain

TABLE 2 Typical Instrument Parameters

Carrier gas	helium
Carrier gas flow rate at 110°C, mL/min	1.2
Detector	flame ionization
Detector temperature, °C	240
Injection port temperature, °C	230
Hydrogen flow rate, mL/min	30
Air flow rate, mL/min	275
Make up gas	nitrogen
Make up gas flow rate, mL/min	23
Split flow, mL/min	150
Column	60 m × 0.32 mm ID × 0.5 µm bonded polyethylene glycol- fused silica capillary
Column temperature, °C	110
Chart speed, cm/min	1
Sample size, µL	0.6

a minimum peak height response for 0.001 weight % impurity of twice the height of the background noise.

7.2 *Column*—A capillary column containing a stationary phase of cross-linked polyethylene glycol has been found to be satisfactory. Any column can be used that is capable of resolving all significant impurities from styrene and from the internal standard.

7.3 *Recorder*—Electronic integration, with tangent capabilities is recommended.

7.4 *100-mL Volumetric Flask*.

7.5 *Microsyringes*, assorted volumes.

8. Reagents and Materials

8.1 *Purity of Reagent*—Reagent grade chemicals shall be used in all tests. 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,⁷ where such specifications are available.

8.2 *Carrier Gas*—Chromatographic grade helium is recommended (minimum purity of 99.95 mole %).

8.3 *Hydrogen and Air*, for FID detector.

8.4 *n-Heptane*, 99.0 % minimum purity, or other internal standard, such as *n*-octane, previously analyzed to be free of compounds coeluting with impurities in the sample.

8.5 *Styrene*, the highest purity available, but not less than 99.7 %.

8.6 *Pure Compounds*, for calibration, shall be those compounds that are typically present in commercial styrene. These should be at least 99 % pure as they are to be used for determining response factors.

9. Hazards

9.1 Consult the latest OSHA regulations, suppliers' Material Safety Data Sheets, and local regulations for all materials used in this procedure.

10. Sampling

10.1 Sample the material in accordance with Practice D 3437.

11. Preparation of Apparatus

11.1 Follow manufacturer's instructions for mounting and conditioning the column into the chromatograph and adjusting the instrument conditions described in Table 2, allowing sufficient time for the equipment to reach equilibrium. See Practices E 355 and E 1510 for additional information on gas chromatographic practices and terminology.

12. Calibration

12.1 Prepare a calibration mixture containing approximately 99.5 weight % styrene and the expected significant impurities at their expected concentration (see Practice D 4307). Weigh all components to the accuracy required to calculate the concentration of each to the nearest 0.001 %.

12.2 With a microsyringe, add 50 µL of internal standard to a 100-mL volumetric flask about three-fourths full of calibration mixture. Mix well. Add calibration mixture to mark and again mix well. If *n*-heptane is used as the internal standard, using a density of 0.684 for *n*-heptane and 0.906 for styrene, this solution will contain 0.0377 weight % *n*-heptane.

12.3 Also prepare a sample of the styrene used for the calibration blend with and without *n*-heptane to determine the concentration of existing impurities and interfering compounds with internal standard. If impurities in the styrene emerge with the chosen internal standard, an alternate internal standard must be used.

12.4 Inject an appropriate amount of sample into the chromatograph and obtain a chromatogram.

12.5 Measure the areas of all peaks, including the internal standard, except the styrene peak.

12.6 Calculate the response factors for each impurity relative to the internal standard as follows:

$$RF_i = \frac{C_i}{(C_s) \left(\frac{A_i}{A_{si}} - \frac{A_b}{A_{sb}} \right)} \quad (1)$$

where:

⁷ *Reagent Chemicals, American Chemical Society Specifications*, American Chemical Society, Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, see *Analar Standards for Laboratory Chemicals*, BDH Ltd., Poole, Dorset, U.K., and the *United States Pharmacopeia and National Formulary*, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.

RF_i = response factor relative to the internal standard,
 A_{si} = area of internal standard in calibration mixture,
 A_i = area of impurity peak in calibration mixture,
 A_{sb} = area of internal standard in styrene used in making calibration mixture,
 A_b = area of impurity in styrene used to make calibration mixture,
 C_s = weight percent internal standard in calibration mixture, and
 C_i = weight percent impurity in calibration mixture.

where:
 A_i = area of impurity,
 A_s = area of internal standard,
 RF_i = response factor for impurity, relative to the internal standard, and
 C_s = concentration of internal standard, in weight percent.

14.3 Calculate the styrene content by subtracting the sum of the impurities from 100.00. Styrene weight percent = 100.00 – (sum of impurities).

13. Procedure

13.1 Establish stable instrument operation at the prescribed or selected operating conditions. Reference should be made to instructions provided by the manufacturer of the chromatograph.

13.2 Prepare sample as described in 12.2.

13.3 Inject appropriate amount of sample into the chromatograph and obtain the chromatogram. A typical chromatogram is shown in Fig. 1.

14. Calculation

14.1 Measure the areas of all peaks, including the internal standard, except the styrene peak.

14.2 Calculate the weight percent of the individual impurities, C_i , as follows:

$$C_i = \frac{(A_i) (RF_i) (C_s)}{A_s} \quad (2)$$

15. Report

15.1 Report the concentration of impurities to the nearest 0.001 % and the styrene content to the nearest 0.01 %.

16. Precision and Bias

16.1 *Precision*—The following criteria should be used to judge the acceptability (95 % probability level) of results obtained by this test method (see Practice D 691). The criteria were derived from a round robin among six laboratories. The data were run on two days using different operators.

16.2 *Intermediate Precision*—Results in the same laboratory should not be considered suspect unless they differ by more than the normal amount shown in Table 3 and Table 4.

16.3 *Reproducibility*—The results by each of two laboratories should not be considered suspect unless they differ by more than the amount shown in Table 3 and Table 4.

16.4 *Bias*—Bias of this test method for the analysis of styrene has not been determined because no acceptable reference material is available.

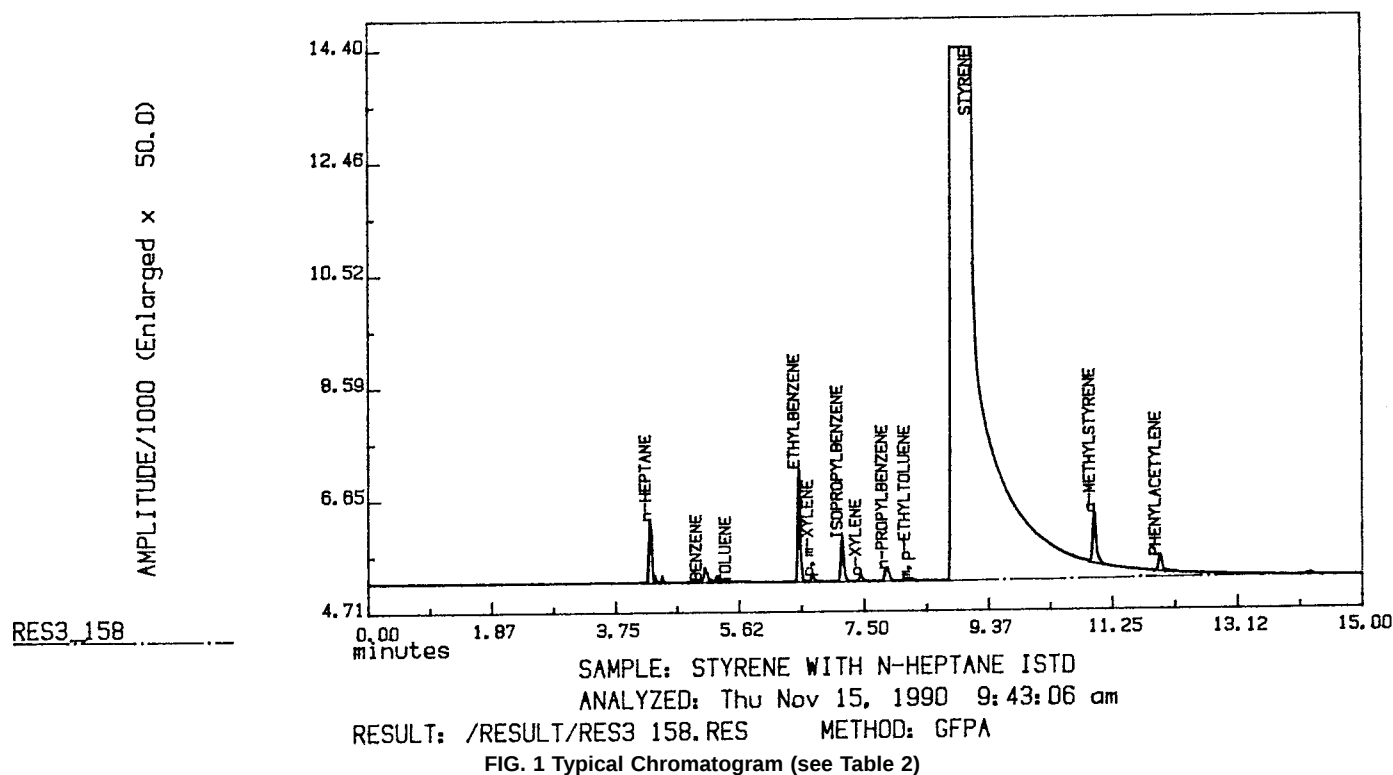


TABLE 3 Precision for Styrene and Impurities at Stated Levels

Component	Concentration, weight %	Intermediate Precision, %	Reproducibility, %
Styrene	99.74	0.017	0.054
α -methylstyrene	0.028	0.0001	0.004
Isopropylbenzene	0.008	0.001	0.001
<i>n</i> -propylbenzene	0.004	0.0003	0.001
<i>m</i> - and <i>p</i> -ethyltoluene	0.014	0.001	0.005
<i>p</i> , <i>m</i> -xylene	0.125	0.005	0.007
<i>o</i> -xylene	0.030	0.001	0.042

TABLE 4 Precision for High Purity Styrene and Impurities at Stated Levels

Component	Concentration, weight %	Intermediate Precision, %	Reproducibility, %
Styrene	99.96	0.024	0.033
Ethylbenzene	0.014	0.003	0.004
α -methylstyrene	0.007	0.002	0.003

17. Quality Control

17.1 *Quality Control*—Confirm the performance of the instrument or the test method procedure by regularly analyzing QC samples.

17.2 The following practice has been found to be effective. A sufficient quantity (5 gal) process sample of styrene is collected and stored in a tightly sealed, stainless steel metal can. This container is equipped with a stainless steel safety faucet, a pressure-relief cap and an internal flame arrestor. The 5 gal can is kept in a refrigerator. Samples are withdrawn from this can once a week and transferred to a quart glass amber bottle that is fitted with a poly-seal cap. The material used for the quality control sample is taken from the glass bottle, which is not kept refrigerated.

18. Keywords

18.1 analysis by gas chromatography; impurities in styrene; purity of styrene; styrene; styrene monomer

SUMMARY OF CHANGES

Committee D16 has identified the following selected changes to this standard since the last issue (D 5135 - 95) that may impact the use of this standard.

- (1) Section 6 (Interferences) was added.
- (2) Section 17 (Quality Control) was added.

- (3) Several editorial revisions were made to conform to D16 Editorial Guidelines for gas chromatographic test methods.

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