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Standard Test Methods for Lead in Gasoline by X-Ray Spectroscopy¹

This standard is issued under the fixed designation D 5059; 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.

This standard has been approved for use by agencies of the Department of Defense.

 ϵ^{1} Note—Warning notes were editorially moved into the standard text in August 2003.

1. Scope*

1.1 These test methods cover the determination of the total lead content of a gasoline within the following concentration ranges:

0.010 to 5.0 g Pb/US gal 0.012 to 6.0 g Pb/UK gal 0.0026 to 1.32 g Pb/L

- 1.1.1 Test Methods A and B cover the range of 0.10 to 5.0 g Pb/US gal. Test Method C covers the range of 0.010 to 0.50 g Pb/US gal.
 - 1.1.2 These test methods compensate for normal variation in gasoline composition and are independent of lead alkyl type.
 - 1.2 Test Method A (formerly in Test Method D2599)—Sections (formerly in withdrawn Test Method D 2599)—Sections 5-9. Test Method B (formerly in Test Method D2599)—Sections (formerly in withdrawn Test Method D 2599)—Sections 10-14. Test Method C (formerly in Test Method D3229)—Sections (formerly in withdrawn Test Method D 3229)—Sections 15-19.
- 1.3 The values stated in SI are to be regarded as the standard. For reporting purposes the values stated in grams per U.S. gallon are the preferred units in the United States. Note that in other countries, other units can be preferred.
- 1.4 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. For specific hazard statements, see Sections 5, 6, 11, and 18.

2. Referenced Documents

ASTM D5050 07

- 2.1 ASTM Standards: D2599Test Method for Lead in Gasoline by X-Ray Spectrometry² D3229Test Method for Low Levels of

 Lead in Gasoline by Wavelength

 Dispersive X-Ray Spectrometry
- D 3341 Test Method for Lead in Gasoline-Iodine Monochloride Method
- D 4057Practice for Manual Sampling of Petroleum and Petroleum Products Practice for Manual Sampling of Petroleum and Petroleum Products P
- <u>D 6299 Practice for Applying Statistical Quality Assurance Techniques to Evaluate Analytical Measurement System Performance</u>
- D 6792 Practice for Quality System in Petroleum Products and Lubricants Testing Laboratories

3. Summary of Test Method

3.1 There are three alternative test methods, as follows.

¹ These test methods are under the jurisdiction of Committee D02 on Petroleum Products and Lubricants and are the direct responsibility of Subcommittee D02.03 on Elemental Analysis.

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These test methods have been approved by the sponsoring committees and accepted by the cooperating organizations in accordance with established procedures.

2 Discontinued. See 1991

² 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, Vol 05.02-volume information, refer to the standard's Document Summary page on the ASTM website.

- 3.1.1 Test Method A (Bismuth Internal Standard Method High Concentration)—One volume of sample is mixed thoroughly with an equal volume of bismuth internal standard solution. The mixture is placed in the X-ray beam and the intensities of the lead L- α_1 radiation at 1.175 Å and the bismuth L- α_1 radiation at 1.144 Å are determined. The lead concentration of the sample is measured by comparing the ratio of gross counting rate at 1.175 Å with the gross counting rate at 1.144 Å to a previous prepared calibration curve of concentration versus the same ratios.
- 3.1.2 Test Method B (Scattered Tungsten Radiation Method)—The ratio of the net X-ray intensity of the lead L- α_1 radiation to the net intensity of the incoherently scattered tungsten L- α_1 radiation is obtained on a portion of the sample. The lead content is determined by multiplying this ratio by a calibration factor obtained with a standard lead solution of known concentration.
- 3.1.3 Test Method C (Bismuth Internal Standard Method, Low Concentration)—Twenty millilitres of sample is mixed thoroughly with two milliliters of bismuth internal standard solution. The mixture is placed in the X-ray beam of a spectrometer and the intensities of the lead L- α_1 radiation at 1.175 \mathring{A} , the bismuth L- α_1 radiation at 1.144 \mathring{A} , and a background at 1.194 \mathring{A} are determined. A blank, made with *iso*-octane and bismuth internal standard, is run using the same procedure. The lead concentration is measured by determining the ratio of the net counting rate at 1.175 \mathring{A} to the gross counting rate at 1.144 \mathring{A} for the sample, subtracting the comparable ratio found for the blank, and comparing to a previously prepared calibration curve of concentration versus the same ratios.

4. Significance and Use

- 4.1 These test methods determine the concentration of lead (from alkyl addition) in gasoline. These alkyl additives improve the antiknock properties.
- 4.2 Test Method C is used to ensure compliance of trace lead as required by federal regulations for lead-free gasoline (40 CFR part 80).

TEST METHOD A (BISMUTH INTERNAL STANDARD)

5. Apparatus

5.1 X-ray Spectrometer, capable of measuring radiations mentioned in 3.1.1 and of being operated under the following instrumental conditions or other giving equivalent results:

Tube Voltage Tube Current Analyzing Crystal Optical Path

Detector

50 kV 20 to 45 mA Lithium Fluoride (LiF) Air, Helium

(Warning—Compressed gas under pressure)
Proportional or Scintillation

Note 1—The X-ray spectrometer and manner of use should comply with the regulations governing the use of ionizing radiation or recommendations of the International Commission of Radiological Protection, or both.

6. Reagents and Materials

- 6.1 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, it is intended that all reagents conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society where such specifications are available.³ 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 determinations.
 - 6.2 Hydrocarbon-Soluble Bismuth.

Note 2—Bismuth 2-Ethylhexoate has been found suitable to use. Other bismuth containing materials that are hydrocarbon-soluble may also be used when they are certified to conform to 6.1.

6.3 Bismuth Internal Standard Solution —Dilute the hydrocarbon-soluble bismuth with a suitable solvent. If bismuth 2-ethylhexoate is used, add 2-ethylhexanoic acid as a stabilizer (see Note 3) to obtain a solution containing the following:

3.00 g Bi/US gal at 15.5°C (60°F) or 3.60 g Bi/UK gal at 15.5°C (60°F) or 0.793 g Bi/L at 15°C

Note 3—Some stability difficulties have been experienced with bismuth 2-ethylhexoate internal standard solution. If the standard is blended to contain 5 % 2-ethylhexanoic acid, the standard has been found to last almost indefinitely. The 2-ethylhexanoic acid stabilizes *iso*-octane, toluene, and benzene solutions of the bismuth 2-ethylhexoate which are otherwise stable for only a day or two. Normal octanoic acid does *not* stabilize solution.

- 6.4 *Iso-octane*. (Warning—Extremely flammable.)
- 6.5 Solvent, capable of dissolving the bismuth internal standard. Mixed xylenes and dodecane have been found suitable to use.

³ Discontinued. See 1991 Annual Book of ASTM Standards, Vol 05.03.

³ 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.



- 6.6 Hydrocarbon-Soluble Lead—Either tetraethyllead (TEL) or a lead-containing compound (for example, lead naphthenate) with a certifiable lead concentration.
- 6.7 Lead (Pb) Standard Solution —Dissolve tetraethyllead (TEL) (Warning—TEL is toxic by ingestion), lead naphthenate (see Note 4), or other suitable lead containing compound in iso-octane (Warning-Extremely flammable), toluene, or a mixture of these two solvents. This standard solution shall contain an accurately known lead concentration of approximately the following magnitude:

5 g Pb/US gal at 15.5°C (60°F) or 6 g Pb/UK gal at 15.5°C (60°F) or 1.3 g Pb/L at 15.5°C

6.7.1 Keep the standard solution refrigerated when not in use.

Note 4—A lead naphthenate solution of same lead concentration has also proven satisfactory as a calibration material. Concentrated TEL is not used to make up standard solutions. The concentrated solution is too acutely toxic to be handled safely under normal laboratory conditions.

Note 5—When this lead standard solution is prepared with TEL, the lead concentration can be determined with Test Method D 3341.

6.8 *Toluene*. (Warning—Flammable. Vapor harmful.)

7. Calibration

- 7.1 Make dilutions of the lead (Pb) standard solution to give 0.10, 1.00, 2.00, 3.00, 4.00 and 5.00 g Pb/US gal at 15.5°C (60°F) or 0.10, 1.00, 2.50, 3.50, 5.00, and 6.00 g Pb/UK gal at 15.5°C (60°F) or 0.025, 0.264, 0.529, 0.793, 1.057, 1.322 g Pb/L at 15°C in toluene, iso-octane, or a mixture of these solvents.
 - 7.2 Allow the lead standards and bismuth internal standard solutions to come to room temperature.
- 7.3 Pipet accurately 10 mL of each standard into separate glass-stoppered bottles or flasks and add an equal, accurately measured volume of the bismuth internal standard solution to each one. Mix thoroughly.
- 7.4 Place one of these solutions in the sample cell using techniques consistent with good operating practice for the spectrometer employed. Place the cell in the instrument, allow the spectrometer atmosphere to reach equilibrium (if appropriate), and determine the counting rate at the lead L- α_1 line (1.175 Å) and at the bismuth L- α_1 line (1.144 Å).

Note 6—When possible, collect at least 100 000 counts at each line. When sensitivity or concentration, or both, makes it impractical to collect this many counts, the technique that allows the greatest statistical precision in the time allotted for each analysis should be used. Sample stability should also be considered in determining counting rate. Variation in counting rates should be observed and if the counting rate tends to go in one direction only, the sample is probably decomposing. If this occurs, shorter counting times should be used consistent with acceptable statistical precision.

7.5 Determine the ratio, R, for each standard as follows:

$$R = A/B \tag{1}$$

where:

A = counting rate at 1.175 Å, andg/standards/sist/bd30593a-a29f-490a-8a75-86d6048c35c7/astm-d5059-07

B = counting rate at 1.144 Å.

7.6 Plot a calibration curve relating R to the grams of lead per gallon.

Note 7—Many modern X-ray spectrometer instruments will plot and store the calibration curve, slope, and related information in the instrument computer system, as an alternative to hand-plotting this information.

8. Quality Control Checks

8.1 Confirm the calibration of the instrument each day it is in use by analyzing a quality control (QC) sample containing a quantifiable concentration of lead, that is, independent of the calibration curve. It is advisable to analyze additional QC samples as appropriate, such as at the beginning and end of a batch of samples or after a fixed number of samples, to ensure the quality of the results. Analysis of result(s) from these QC samples can be carried out using control chart techniques. 4 When the QC sample result causes the laboratory to be in an out-of-control situation, such as exceeding the laboratory's control limits, instrument re-calibration may be required. An ample supply of QC sample material shall be available for the intended period of use, and shall be homogeneous and stable under the anticipated storage conditions. If possible, the QC sample shall be representative of samples typically analyzed and the average and control limits of the QC sample shall be determined prior to monitoring the measurement process. The QC sample precision shall be checked against the ASTM method precision to ensure data quality. Further guidance on quality control can be found in Practices D 6299 and D 6792.

9. Procedure

- 9.1 Obtain sample in accordance with Practice D 4057.
- 9.2 Prepare the samples to be analyzed as described in 7.3 and 7.4 for the standard lead solutions and determine the ratio, R, as described in 7.5.

⁴ Annual Book of ASTM Standards, Vol 05.02.

ASTM MNL 7, Manual on Presentation of Data Control Chart Analysis, "Section 3, Control Charts for Individuals," 6th ed., ASTM International, W. Conshohocken.