



Standard Test Methods for Comparison of Waterborne Petroleum Oils by Gas Chromatography¹

This standard is issued under the fixed designation D3328; 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 comparison of petroleum oils recovered from water or beaches with oils from suspect sources by means of gas chromatography (**1, 2, 3**).² Such oils include distillate fuel, lubricating oil, and crude oil. The test method described is for capillary column analyses using either single detection (flame ionization) or dual detection (flame ionization and flame photometric) for sulfur containing species.

1.2 This test method provides high resolution for critical examination of fine structure that is resistant to weathering. The flame-photometric detection for sulfur components is an adjunct, not a substitute, for flame-ionization detection in the identification of waterborne petroleum oils (**4-12**). For this reason, flame photometric detection is optional.

1.3 *This standard does not purport to address 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

2.1 ASTM Standards:³

D1129 Terminology Relating to Water

D1193 Specification for Reagent Water

D2549 Test Method for Separation of Representative Aromatics and Nonaromatics Fractions of High-Boiling Oils by Elution Chromatography

D3325 Practice for Preservation of Waterborne Oil Samples

D3326 Practice for Preparation of Samples for Identification of Waterborne Oils

¹ These test methods are under the jurisdiction of ASTM Committee D19 on Water and are the direct responsibility of Subcommittee D19.06 on Methods for Analysis for Organic Substances in Water.

Current edition approved Feb. 15, 2013. Published March 2013. Originally approved in 1974. Last previous edition approved in 2006 as D3328 – 06. DOI: 10.1520/D3328-06R13.

² The boldface numbers in parentheses refer to the references at the end of these test methods.

³ 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.

D3415 Practice for Identification of Waterborne Oils

D4489 Practices for Sampling of Waterborne Oils

D5739 Practice for Oil Spill Source Identification by Gas Chromatography and Positive Ion Electron Impact Low Resolution Mass Spectrometry

E355 Practice for Gas Chromatography Terms and Relationships

3. Terminology

3.1 *Definitions*—For definitions of terms used in this test method, refer to Practice **D3415**, Terminology **D1129**, and Practice **E355**.

4. Significance and Use

4.1 Identification of a recovered oil is determined by comparison with known oils, selected because of their possible relationship to the particular recovered oil. The known oils are collected from suspected sources. Samples of such known oils must be collected and submitted along with the unknown for analysis. At present, identification of the source of an unknown oil by itself cannot be made (for example, from a library of known oils).

4.2 The use of a flame-photometric detector in addition to the flame-ionization detector provides a second, independent profile of the same oil, that is, significantly more information is available from a single analysis with dual detection.

4.3 Many close similarities (within uncertainties of sampling and analysis) will be needed to establish identity beyond a reasonable doubt. The analyses described will distinguish many, but not all samples. For cases in which this method does not clearly identify a pair of samples, and for important cases where additional comparisons are needed to strengthen conclusions, other analyses will be required (refer to Practice **D3415**). In particular, Practice **D5739** is useful for such cases.

5. Interferences

5.1 Compounds that have the same retention time as petroleum hydrocarbons will interfere in the comparison of the unknown with known oils. This is particularly true if animal fat or vegetable oil, naturally occurring hydrocarbons, or spill-treatment chemicals are present in relatively large amounts. Independent analysis, for example, infrared spectroscopy, will

establish the presence of these contaminants if their presence is suspected. Animal or vegetable oils can be removed effectively by Test Method **D2549** or by Practices **D3326** (Method D).

NOTE 1—Test Method **D2549** will also remove the aromatic fraction.

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 shall conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society.⁴

6.2 Unless otherwise indicated references to water shall be understood to mean reagent water that meets the purity specifications of Type I or Type II water presented in Specification **D1193**.

6.3 *Air*—For use with the flame-ionization and flame-photometric detectors; may be obtained using a laboratory pure air generator, or from a zero grade tank supply.

6.4 *Carrier Gas*—High-purity grade helium is used as carrier gas.

6.5 *Cyclohexane*—High-purity (HPLC-grade). For sample preparation and for use in reference standards.

6.6 *Hydrogen*—For use with the flame-ionization and flame-photometric detectors; may be obtained using a hydrogen generator, or from a prepurified grade tank supply.

6.7 *Methylene Chloride*—For use in reference standards and glassware cleaning.

6.8 *Normal Alkane Standards*—Normal alkanes, decane through hexatriacontane, for use as reference compounds.

6.9 *Thiophene*—For use in optimization of flame-photometric detector.

7. Reference Standards

7.1 *Normal Paraffinic Hydrocarbons*—Prepared mixtures of approximately decane to hexatriacontane, or selected individual normal paraffins, are run under normal analysis conditions to determine retention times of compounds.

7.2 *Resolution Mixture*—Equal mixtures of *n*-heptadecane, *n*-octadecane, pristane and phytane in solution. See **Annex A1** for details (**A1.2.1**).

8. Sampling

8.1 Collect a representative sample in accordance with Practice **D4489**.

8.2 If the sample is not to be analyzed within 1 week, it should be preserved in accordance with Practice **D3325** because of the possibility of bacterial decomposition of normal paraffins in the sample.

8.3 The sample should be prepared for analysis in accordance with Practices **D3326**, because of the great variety of materials and circumstances associated with collecting petroleum oils from the environment. For heavier oils, a procedure to deasphalt the oil may be necessary.

9. Summary of Test Method

9.1 This test method uses a gas chromatographic capillary column system for the separation of petroleum hydrocarbons. The effluent of the column may be detected with a flame-ionization detector, or it may be split (1 + 2) between a flame ionization and a flame-photometric detector. The flame photometric detector is equipped with a narrow bandpass interference filter for spectral isolation of the sulfur emission at 394 nm. The relative peak size of each component (as indicated by retention time) of recovered oil is compared visually with the relative peak size of each component (of like retention time) of the suspected source.

NOTE 2—This dual detector method is based on the early work done by Kahn (**13**), Garza (**4**), and Adlard (**7**).

9.2 In this test method, elution of characteristic hydrocarbons occurs generally in order of increasing boiling point.

10. Apparatus

10.1 *Chromatographic Column*—Fused silica capillary column with bonded phase SE-30 or equivalent, 30 m by 0.32 mm inside diameter (0.1 μ m film thickness).

NOTE 3—Other columns, providing equivalent or better resolution may be substituted (see **Annex A1**), but the analysis time will be increased with longer columns.

10.2 *Gas Chromatograph*—A commercial or custom designed gas chromatograph with heated injection and detector zones and a column oven capable of being programmed from 75°C to at least 325°C for heavier oils (higher boiling than gasoline, jet fuels, etc.).

10.2.1 For light distillate fuels, the chromatograph must be capable of programming from 50°C and also be capable of maintaining isothermal control at 50°C.

10.2.2 *Carrier Gas Pressure Regulator* is substituted pressure regulator for the mass flow controllers to give more precise rates in the low flow ranges (1 to 5 mL/min).

10.2.3 *Injection Port*—The use of glass injector inserts that can be replaced or cleaned frequently, or both, will prolong the useful life of the column (**3**).

10.2.4 *Detectors*—A hydrogen-flame ionization detector is always used for analyses. A flame-photometric detector with a 394 nm bandpass filter is used for dual detection (**9**, **10**, **11**, **12**).

10.2.5 *Carrier Gas Makeup* is required at the effluent of the column with a temperature independent mass flow controller.

10.2.6 *Effluent Splitter*—An effluent splitter with a split ratio of 1 + 2 (FID/FPD) is required for dual detection.

10.2.7 *Bleeder for Reference Compound*—A device for in-line bleed of a reference compound (thiophene and cyclohexane) into the carrier flow for detector optimization is required, when using a flame-photometric detector.

10.2.8 *Recorder*, or an integrator or computer data handling system capable of acquiring data at a rate compatible with the

⁴ "Reagent Chemicals, American Chemical Society Specifications," Am. Chemical Soc., Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, see "Reagent Chemicals and Standards," by Joseph Rosin, D. Van Nostrand Co., Inc., New York, NY, and the "United States Pharmacopeia."