



Designation: D7515 – 19 (Reapproved 2023)

Standard Test Method for Purity of 1,3-Propanediol (Gas Chromatographic Method)¹

This standard is issued under the fixed designation D7515; 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 describes the gas chromatographic determination of purity for 1,3-propanediol (PDO). This test method was originally developed to determine the purity of 1,3-propanediol used for the application as the freeze point depressant base fluid in formulated PDO engine coolants. Use of the method for purity of PDO for other applications may be viable.

1.2 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

1.2.1 *Exception*—Inch-pound units (psi) are used in [Table 1](#), Pressure Program, Options A and B.

1.3 Review the current Material Safety Data Sheets (MSDS) for detailed information concerning toxicity, first aid procedures, and safety precautions.

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, health, and environmental practices and determine the applicability of regulatory limitations prior to use.*

1.5 *This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.*

2. Referenced Documents

2.1 ASTM Standards:²

¹ This test method is under the jurisdiction of ASTM Committee D15 on Engine Coolants and Related Fluids and is the direct responsibility of Subcommittee D15.04 on Chemical Properties.

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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* volume information, refer to the standard's Document Summary page on the ASTM website.

[D1123 Test Methods for Water in Engine Coolant Concentrate by the Karl Fischer Reagent Method](#)

[E177 Practice for Use of the Terms Precision and Bias in ASTM Test Methods](#)

[E300 Practice for Sampling Industrial Chemicals](#)

[E691 Practice for Conducting an Interlaboratory Study to Determine the Precision of a Test Method](#)

[E2409 Test Method for Glycol Impurities in Mono-, Di-, Tri- and Tetraethylene Glycol and in Mono- and Dipropylene Glycol \(Gas Chromatographic Method\)](#)

3. Summary of Test Method

3.1 The sample is analyzed by a temperature-programmed gas chromatograph, equipped with a capillary column and flame ionization detector (FID), and quantification is performed by direct area normalization.

3.2 Additionally, the use of a reference sample using Ethylene, Propylene, or Dipropylene Glycol (EG, PG or DPG) in 1,3-PDO (minimum purity 99.5 %) should be used as a performance check (see [Section 8](#)).

NOTE 1—The application of this reference sample is also used to demonstrate the separation of commonly used glycols (EG, PG and DPG) in engine coolants, from PDO. Solutions of EG, PG, or DPG in concentrations of 0.1 to not more than 1 % may be used.

4. Significance and Use

4.1 Knowledge of an approved method is required to establish whether the product meets the requirements of its specifications. The use of glycols in the reference sample is not intended to suggest the presence of glycol (EG, PG and DPG) impurities, but to demonstrate and quantify the separation of commonly used Engine Coolant glycols from PDO.

5. Apparatus

5.1 *Gas Chromatograph(s)*—provided with a sample splitter or on-column injection, flame ionization detector and temperature-programming facilities. The instrument must be suitable for analysis according to the operating instructions given in [Table 1](#). To account for differences among laboratory equipment, the two most common column choices are listed.

TABLE 1 Typical Operating Parameters for the GC Analysis of PDO

Column ^A	Option A	Option B	Option C ^B
Type	Capillary	Capillary	Capillary
Material	Fused Silica	PEG	PEG
Length × I.D.	10 m × 0.1 mm	30 m × 0.25 mm	30 m × 0.32 mm
Stationary Phase	DB-5	ZB-Wax	ZB-Wax
Film Thickness	0.17 µm	0.25 µm	1 µm
Detector System			
Type	FID	FID	FID
Sensitivity	The ratio of the signal to the noise level must be at least 2:1 at a concentration of 5 mg/kg glycols in PDO	The ratio of the signal to the noise level must be at least 2:1 at a concentration of 5 mg/kg glycols in PDO	The ratio of the signal to the noise level must be at least 2:1 at a concentration of 5 mg/kg glycols in PDO
Temperatures			
Column Oven			
Initial	0.5 min at 35 °C	0 min at 50 °C	0 min at 100 °C retention time 0.5 min
Ramp 1	35 °C to 85 °C at 50 °C/min	50 °C to 200 °C at 15 °C/min	100 °C to 180 °C at 15 °C/min; 5.83 min
Ramp 2	85 °C to 325 °C at 100 °C/min	200 °C to 250 °C at 40 °C/min	180 °C to 225 °C at 30 °C/min; 0 min
Ramp 3	2 min at 325 °C	17 min at 250 °C	Hold 3.7 min at 225 °C
Detector	325 °C	250 °C	250 °C
Carrier Gas	Helium	Helium	Helium or Nitrogen
Calibration	This method employs straight area normalization so no calibration is required	This method employs straight area normalization so no calibration is required	Calibration
Injected Volume	0.1 µL	0.2 µL	1.0 µL
Pressure Program	0.5 min at 30 psi 30 psi to 100 psi at 100 psi/min 8 min at 100 psi Gas saver on at 0.5 min	Pressure: 13.2 psi at 50 °C Flow: 1.1 mL/min Velocity: 28 cm/s	Constant flow at 5 mL/min
Split Ratio	1:250 or appropriate split ratio to allow adequate sensitivity as defined under Detector System	1:18 or appropriate split ratio to allow adequate sensitivity as defined under Detector System (only if split injection technique is used)	Inlet temperature 235 °C Split flow 300 mL/min Split ratio 60

^A The columns are available commercially. Some column suppliers market alternative stationary phases. The chromatogram obtained must be identical, with regard to separation of PDO and other glycol components, to those illustrated in Fig. A1.1 and Fig. A1.2.

^B Option C instrument method is for reference only; due to instrument operational variations, temperature ramp, gas flow, and split flow/ratio may need adjustment to achieve separation of analyzed glycols.

NOTE 2—Other column suppliers market alternative stationary phases, therefore, it is permissible to use a different column from an alternative supplier. However, the chromatogram obtained must be identical, with regard to separation of PDO and other glycol components, to those illustrated in Fig. A1.1 and Fig. A1.2.

5.1.1 *Columns*—The analytical column used must completely separate EG, PG or DPG from PDO. Fig. A1.1 and Fig. A1.2 show examples of chromatograms conforming to the requirements.

5.2 *Digital Integration Equipment*—A computer with data collection software.

5.3 *Analytical Balance*, readability 0.1 mg, calibrated. Calibrate and verify at regular intervals.

5.4 *Crimp Top Vials*, 1 mL and 5 mL.

5.5 *Crimper/De-capper*, for capping and de-capping the vials.

5.6 *Micro Syringes*, 5 µL or 10 µL.

5.7 *Bottles*, 100 mL, with screw cap.

6. Reagents and Materials

6.1 *Purity of Reagents*—Unless otherwise indicated, it is intended that all reagents shall conform to the specifications of the Committee on Analytical Reagents of the American Chemi-

cal 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 determination.

6.2 Reagents:

6.2.1 *1,3-Propanediol (PDO)*, minimum purity 99.5 % mass (m/m).

6.2.2 *Ethylene Glycol (EG)*, minimum purity 99.5 % mass (m/m).

6.2.3 *Propylene Glycol (PG)*, minimum purity 99.5 % mass (m/m).

6.2.4 *Dipropylene Glycol (DPG)*, minimum purity 99.0 % mass (m/m).

6.3 *Water or Methanol*, HPLC grade.

6.4 *Internal Reference Sample*.

7. Sampling, Test Specimens and Test Units

7.1 Follow the relevant instructions for sampling as given in Practice E300.

³ ACS Reagent Chemicals, Specifications and Procedures for Reagents and Standard-Grade Reference Materials, 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.