

Designation: D8086 - 20

Standard Test Method for Determination of Methanol and Ethanol in Electrical Insulating Liquids of Petroleum Origin by Headspace (HS)-Gas Chromatography (GC) Using Mass Spectrometry (MS) or Flame Ionization Detection (FID)¹

This standard is issued under the fixed designation D8086; 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 determination of byproducts of cellulosic materials degradation found in electrical insulation systems that are immersed in insulating liquid. Such materials include paper, pressboard, wood and cotton materials. This test method allows the analysis of methanol and ethanol from the sample matrix by headspace GC-MS or GC-FID.

1.2 This test method has been used to test for methanol and ethanol in mineral insulating liquids and less flammable electrical insulating liquids of mineral origin as defined in D3487 and D5222 respectively. Currently, this method is not a practical application for ester liquids.

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

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

D923 Practices for Sampling Electrical Insulating Liquids

D3487 Specification for Mineral Insulating Oil Used in Electrical Apparatus

- D3612 Test Method for Analysis of Gases Dissolved in Electrical Insulating Oil by Gas Chromatography
- D5222 Specification for High Fire-Point Mineral Electrical Insulating Oils
- D5837 Test Method for Furanic Compounds in Electrical Insulating Liquids by High-Performance Liquid Chromatography (HPLC)

3. Terminology

3.1 Definitions:

3.1.1 *extract ion mass spectrum, n*—a record that shows a specific mass-to-charge ratio (m/z) extracted from a mass spectrum.

3.1.2 mass spectrum, n—a record that shows the relative number of ions of various mass that are produced when a given substance is processed in a mass spectrometer.

4. Summary of Test Method

4.1 Analysis of methanol (CH₃OH) and ethanol (C₂H₅OH) in electrical insulating liquids consists of bringing an insulating liquid sample in contact with a gas phase (headspace) in a closed vessel. The dissolved species contained in the insulating liquid are then equilibrated between the two phases in contact under controlled conditions (according to Henry's law). At equilibrium, the headspace is over-pressurized with a carrier gas and then the content of a loop is filled by the depressurization of the headspace against the ambient atmospheric pressure (see Note 1).

Note 1—Other headspace principles may also be used but need to be verified and the analytical performance may be somewhat different than listed.

4.2 The gases contained in the loop or in the syringe are introduced into a gas chromatograph.

4.3 Methanol and ethanol in the test specimen are quantified using calibration curves.

¹This test method is under the jurisdiction of ASTM Committee D27 on Electrical Insulating Liquids and Gases and is the direct responsibility of Subcommittee D27.03 on Analytical Tests.

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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.

5. Significance and Use

5.1 Methanol and ethanol are generated by the degradation of cellulosic materials used in the solid insulation systems of electrical equipment. More particularly, methanol comes from the depolymerization of cellulosic materials.^{3,4,5,6}

5.2 Methanol and ethanol, which are soluble in an insulating liquid to an appreciable degree, will proportionally migrate to that liquid after being produced from the cellulose.

5.3 High concentrations or unusual increases in the concentrations of methanol or ethanol, or both, in an insulating liquid may indicate cellulose degradation from aging or incipient fault conditions. Testing for these alcohols may be used to complement dissolved gas-in-oil analysis and furanic compounds as performed in accordance with Test Methods D3612 and D5837 respectively.

6. Interferences

6.1 Vessels used for this test need to be prepared with solvents containing no trace of methanol and ethanol. Additionally, solvents that can break down into these alcohols must not be used.

7. Apparatus

7.1 Analytical balance capable of weighing to the nearest 0.0001g.

7.2 Headspace sampler either equipped with an injection loop and a transfer line or direct injection with gas-tight syringe connected to the injection port of the gas chromatograph. The sampler must be capable of equilibrating the species of interest in a specific time. The required equilibration time can be minimized by mixing the sample during the equilibration period and this can be achieved by using a sampler equipped with mechanical shaking. A direct injection headspace vial system may also be used.

7.3 Gas chromatograph equipped with a mass spectrometer as described in Table 1 or equipped with FID detector as described in Table 2 (see Note 2).

Note 2—This method was developed with He as the carrier gas. Other carrier gases may also be used with this method but must be verified. Analytical performance may be somewhat different than that listed in this method.

7.4 VF-624ms capillary column (60 m \times 0.25 mm diameter with a film thickness of 1.4 μ m) or DB-624 (60 m \times 0.53 mm

TABLE 1 Instrumental Conditions for MS Detection

Test SpecimenLoop volume0.5 mLShakingPowerMaximum LevelTemperaturesSample90 °CInjection loop150 °CTransfer line175 °CPressureVial over-pressure138 kPaTimesEquilibration40 minPressurization0.2 minLoop fill0.12 minLoop equilibration0.25 minInjection6 minDirect Injection HeadspaceSystem parametersTest Specimen2.5 mLOven Temperature90 °COven Temperature90 °COven ParametersShaking for 40 minSyringe Temperature100 °CGas Chromatograph parameters275 °C at 138 kPaSplit ratio5 : 1Column60m VF-624msOven Temp Initial40 °C for 10 minRamp 140 to 275 °C at 20 °C/minHold275 °C for 33.25 minDetectorIonization energy 70 eVMass SpectrometerIonization energy 70 eVInterface at 280 °Cm/z = 30-300 in TIC mode at 0.35 scans/s					
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m/z = 30-300 in TIC mode at 0.35 scans/s		Interface at 280	Interface at 280 °C		
scans/s		m/z = 30-300 in	m/z = 30-300 in TIC mode at 0.35		
		scans/s			

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TABLE 2 Instrumental Conditions for FID Detection

Headspace Sampler parameters				
Test Specimen	Loop volume	0.5 mL		
Shaking	Power	Maximum Level		
Temperatures	Sample	90 °C		
	Injection loop	150 °C		
	Transfer line	175 °C		
Pressure	Vial over-pressure	138 kPa		
Times	Equilibration	40 min		
	Pressurization	0.2 min		
	Loop fill 12 pe 1 2 /astm	0.12 min		
	Loop equilibration	0.25 min		
	Injection	6 min		

Direct Injection Headspace System parameters				
Test Specimen	2.5 mL			
Oven Temperature	90 °C	90 °C		
Oven Parameters	Shaking for	Shaking for 40 min		
Syringe Temperature	100 °C	100 °C		
Gas Chromatograph parameters				
He carrier gas flow:	11 mL/min fo	11 mL/min for 20 min		
Injector Splitless	110 °C meg	110 °C megabore direct		
Column	DB-624	DB-624		
	60 m × 0.53	60 m \times 0.53 mm ID 3 μ m film thickness		
Oven Temp Initial	35 °C for 10	35 °C for 10 min		
Ramp 1	35 to 250 °C	35 to 250 °C at 5 °C/min		
Hold	250 °C for 5	250 °C for 5 min		
Detector				
FID	Temperature	300 °C		
	Hydrogen	40 mL/min		
	Air	400 mL/min		
	He Makeup	5 mL/min		

ID with a film thickness of 3 μ m) for the separation of methanol and ethanol. Other columns have been found to be suitable (see Note 3).

Note 3—Columns that give adequate peak separation may also be used with this method but must be verified. Analytical performance may be somewhat different than that listed in this method.

³ Jalbert, J., Gilbert, R., Tétreault, P., Morin, B., Lessard- Déziel, D., "Identification of a chemical indicator of the rupture of 1,4-β-glycosidic bonds of cellulose in an oil-impregnated insulating paper system," *Cellulose*, 14:295-309, 2007.

⁴ Gilbert, R., Jalbert, J., Tétreault, P., Morin, B., and Denos, Y., "Kinetics of the production of chain-end groups and methanol from the depolymerization of cellulose during the ageing of paper/oil systems," Part 1: Standard wood kraft insulation, *Cellulose*, 16: 327-338, 2009.

⁵ Gilbert, R., Jalbert, J., Duchesne, S., Tétreault, P., Morin, B., and Denos, Y., "Kinetics of the production of chain-end groups and methanol from the depolymerization of cellulose during the ageing of paper/oil systems," Part 2: Thermallyupgraded insulating papers, *Cellulose*, 17: 253-269, 2010.

⁶ Jalbert, J., Rodriguez-Celis, E., Duchesne, S., Morin, B., Ryadi, M., and Gilbert, R., "Kinetics of the production of chain-end groups and methanol from the depolymerization of cellulose during the ageing of paper/oil systems," Part 3: extension of the study under temperature conditions over 120 °C, *Cellulose*, 22: 829-848, 2015.

7.5 Headspace glass vials of 20 mL nominal capacity. The same batch of vials must be used for calibration and the analysis of test specimens from samples (see Note 4).

Note 4—Other vessel volumes may also be used but the analytical performance would have to be verified and may be somewhat different than that specified in the method.

7.6 Crimping system, including crimp head and decapper head.

7.7 Perforated aluminum caps.

7.8 Polytetrafluoroethylene (PTFE) faced butyl septa for headspace vials.

7.9 30 mL or 50 mL glass syringes equipped with three-way plastic stopcocks for sampling.

7.10 The instrumental conditions for the analysis by GC-MS are given in Table 1 and a total ion and extract ion (m/z = 31) mass spectrums obtained under these conditions are given in Fig. 1. The system must be capable of sufficiently



FIG. 1 Typical Chromatograms a) Total ions and b) extracted ion (m/z = 31) mass spectra

separating the component gases and measuring from low $\mu g/kg$ (ppb) to mg/kg (ppm) levels. The detection limits obtained by one laboratory with a 0.5 mL injection loop and capillary column is 23 and 26 $\mu g/kg$ (ppb) using a signal/noise = 3 for methanol and ethanol respectively (see Note 5).

Note 5—The detection limits were obtained from the analysis of methanol and ethanol at a concentration of 40 μ g/kg. These results were obtained with a headspace sampler coupled with a gas chromatograph of one commercial source; other devices can be used but the analytical performance may be somewhat different than that specified in the method.

7.11 The instrumental conditions for the analysis by FID are given in Table 2 and the chromatogram obtained under these conditions is provided in Fig. 2. The detection limits obtained by one laboratory with a 2.5 mL direct injection is $30 \mu g/kg$ for both methanol and ethanol.

8. Reagents and Materials

8.1 Carrier gas with purity at least 99.999 %.

8.2 Methanol and ethanol - HPLC grade.

8.2.1 10-µL, 50-µL, 100-µL, 250-µL and 100-mL gastight syringes.

8.3 Volumetric dispenser.

9. Sampling, Test Specimens, and Test Units

9.1 Obtain insulating liquid samples in accordance with sampling procedures listed in Practices D923 for the use of glass syringes. Since methanol and ethanol are volatile, sample bottles are not to be used.

10. Preparation of Apparatus

10.1 *Headspace (HS) Vial Preparation:* 10.1.1 Prepare a series of vials and the appropriated caps equipped with PTFE-faced butyl septa. Ensure that the lined side is turned towards the inside of the vial. The use of the vials is detailed in Sections 11 and 12.

11. Calibration and Standardization

11.1 Preparation of Calibration Standards in Insulating Liquid:

11.1.1 The signal is calibrated by injecting a series of dilutions prepared from a primary standard of methanol and ethanol in an insulating liquid that is the same as the test

specimens being tested. The primary standard is prepared by diluting a weighed 10 μ L of each, methanol and ethanol in 100 mL of insulating liquid that does not contain any detectable amounts of methanol and ethanol according to this method. This will produce a final concentration of about 100 mg/kg (ppm) each. These exact weights may not be possible, and thus the actual values are to be recorded in order to determine the actual concentration of the primary standard. Primary standard is to be prepared and stored in a 100-mL glass gastight syringe with a luer lock septum tip.

11.1.2 Alternatively, the primary standard can be prepared by adding 1 mL of each alcohol to 498 mL of toluene. This will produce a concentration of about 1800 mg/kg (ppm). Next, add 1 mL of the primary standard to 999 mL of insulating liquid. This will produce a final concentration of about 1800 μ g/kg (ppb). The actual concentrations shall be calculated with the actual weights of the volumes used.

11.1.3 Stock calibration standards are to be made with each insulating liquid that will be tested.

11.1.4 Regardless of the injection mode (HS sampler or direct HS sampler), prepare the calibration standards with the appropriate volume. Directly prepare the standards in 20 mL HS vials of which the final liquid volume will be 10 mL. Crimp the vials using aluminum caps equipped with PTFE-faced butyl septa using a standard crimper head. Add a volume of primary standard in the vial through the septum using an array of gastight syringes. The added volume is to be weighed in order to know the accurate concentration of the calibration standards. An internal standard could be used with the MS detection method (see Note 6).

Note 6—An internal standard is a substance that is spiked in a constant amount to calibration standards and samples. The internal standard solution should be prepared in an insulating liquid that is the same as the test specimens being tested. A laboratory spiked 5 μ L of ethanol-d6 (CD₃CD₂OD) solution 500 μ g/kg as internal standard. This prepared solution is stable and performs ideally when used in conjunction with mass spectrometry detection. For quantification, the response (*rsp*) of the alcohols is divided by the response of the internal standard.

11.1.5 Place the vials onto the headspace sampler rack and begin the analysis using the instrumental conditions given in Table 1 or Table 2.

11.2 Standardization:



FIG. 2 FID Typical Chromatogram