



Designation: D4128 – 18 (Reapproved 2026)

Standard Guide for Identification and Quantitation of Organic Compounds in Water by Combined Gas Chromatography and Electron Impact Mass Spectrometry¹

This standard is issued under the fixed designation D4128; 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 guide covers the identification and quantitation of organic compounds by gas chromatography/mass spectrometry (GC-MS) (electron impact) that are present or extracted from water and are capable of passing through a gas chromatograph without alteration. This guide can be used to provide tentative identifications of volatile and semi-volatile organics, but is restricted to (a) compounds for which reference spectra can be obtained and (b) compounds that can be separated by gas chromatography (GC). These restrictions are imposed on the guide, but are not a limitation of the technique. The guide is written for analysis using automated data acquisition and handling.

1.2 Guidelines have been included for quantitation using ASTM Test Methods [D3871](#), [D3973](#), and other GC-MS volatile/semivolatile procedures used for environmental analysis². The actual detection limits for each component must be determined in each laboratory. Actual detection amounts will vary with the complexity of the matrix, the kind and condition of the GC-MS system, the sample preparation technique chosen, and the application of cleanup techniques to the sample extract, if any. Lower levels of detection can be achieved using modern sensitive instruments or with selected ion monitoring (SIM). To determine the interlaboratory detection estimate (IDE) and the interlaboratory quantitation estimate (IQE), follow Practices [D6091](#) and [D6512](#).

1.3 The guide is applicable to the identification of many organic constituents of natural and treated waters. It includes all modes of sample introduction, including injection of organic extracts, direct aqueous injection, and purge and trap techniques.

1.4 The guide is applicable to capillary column gas chromatography.

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

1.6 *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.*

1.7 *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:³

- [D1066 Practice for Sampling Steam](#)
- [D1129 Terminology Relating to Water](#)
- [D1192 Guide for Equipment for Sampling Water and Steam in Closed Conduits](#) (Withdrawn 2003)⁴
- [D1193 Specification for Reagent Water](#)
- [D2908 Practice for Measuring Volatile Organic Matter in Water by Aqueous-Injection Gas Chromatography](#)
- [D3370 Practices for Sampling Water from Flowing Process Streams](#)
- [D3694 Practices for Preparation of Sample Containers and for Preservation of Organic Constituents](#)
- [D3871 Test Method for Purgeable Organic Compounds in Water Using Headspace Sampling](#)
- [D3973 Test Method for Low-Molecular Weight Halogenated Hydrocarbons in Water](#)
- [D5175 Test Method for Organohalide Pesticides and Polychlorinated Biphenyls in Water by Microextraction and](#)

¹ This guide is under the jurisdiction of ASTM Committee [D19](#) on Water and is the direct responsibility of Subcommittee [D19.06](#) on Methods for Analysis for Organic Substances in Water

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² EPA Methods 624 and 8260C (volatiles) and EPA Methods 625 and 8270D (semivolatiles) are suitable for quantitation.

³ For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at www.astm.org/contact. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

⁴ The last approved version of this historical standard is referenced on www.astm.org.

Gas Chromatography

D5316 Test Method for 1,2-Dibromoethane and 1,2-Dibromo-3-Chloropropane in Water by Microextraction and Gas Chromatography

D5317 Test Method for Determination of Chlorinated Organic Acid Compounds in Water by Gas Chromatography with an Electron Capture Detector

D5789 Practice for Writing Quality Control Specifications for Standard Test Methods for Organic Constituents (Withdrawn 2002)⁴

D6091 Practice for 99 %/95 % Interlaboratory Detection Estimate (IDE) for Analytical Methods with Negligible Calibration Error (Withdrawn 2023)⁴

D6512 Practice for Interlaboratory Quantitation Estimate (Withdrawn 2023)⁴

E260 Practice for Packed Column Gas Chromatography

E355 Practice for Gas Chromatography Terms and Relationships

2.2 EPA Documents:

EPA Method 524.2–1995 Measurement of Purgeable Organic Compounds in Water by Capillary Column Gas Chromatography/Mass Spectrometry⁵

EPA Method 624–1984 Purgeables⁵

EPA Method 625–1984 Base/Neutrals and Acids⁵

SW-846 Method 8270D Semivolatile Organic Compounds by Gas Chromatography (GC-MS)^{6,7}

SW-846 Method 8260C Volatile Organic Compounds by Gas Chromatography (GC-MS)^{6,7}

3. Terminology

3.1 Definitions:

3.1.1 For definitions of terms relating to water used in this standard, refer to Terminology **D1129**. For definitions of terms relating to gas chromatography used in this standard, refer to Practice **E355**.

3.2 Definitions of Terms Specific to This Standard:

3.2.1 *characteristic ion, n*—usually the primary ion in the mass spectrum used to measure response for quantitation purposes. When there are interferences in the mass chromatogram of a primary ion, a secondary characteristic ion must be used for quantitation.

3.2.2 *confirmed identification, n*—in order to confirm a tentative identification, both the GC retention time and the mass spectrum of a compound shall uniquely match those of a reference compound as demonstrated by co-injection of the authentic standard with the tentatively identified compound.

3.2.3 *mass chromatogram, n*—(see *Discussion*)—a limited mass RGC, or mass chromatogram, represents the intensities of ion currents for only those ions having particular mass to charge ratios. It is a means of quickly scanning a complex RGC plot to locate peaks which could be specific compounds or

types of compounds. However, a complete mass spectrum is required for tentative identification.

3.2.3.1 *Discussion*—There are several synonyms in current use for mass chromatogram. These include: mass fragmentogram, extracted ion current profile, and limited mass reconstructed gas chromatogram.

3.2.4 *match, n*—two criteria must be satisfied to verify a comparison of a sample component to a standard match: (1) elution of the sample component at the same retention time as the standard component as shown by co-injection or standard addition, and (2) correspondence of the sample component and the standard component mass spectrum. If co-elution of interfering components prohibits accurate assignment of the sample component retention time from the total ion chromatogram, the retention time should be assigned by using extracted ion current profiles for ions unique to the component of interest. To meet the second criteria, all ions present in the authentic mass spectra at a relative intensity greater than 10 % (whereas the most abundant ion in the spectrum equals 100 %) must be present in the sample spectrum; the relative intensities of these ions must agree within ± 20 % between the standard and sample spectra. (As an example, for an ion with an abundance of 50 % in the standard spectra, the corresponding sample abundance must be between 30 % and 70 %.) However, there may be additional peaks in the sample mass spectrum caused by co-eluting interfering components that are not present in the reference mass spectrum.

3.2.5 *reconstructed gas chromatogram (RGC), n*—(see *Discussion*)—an RGC is the computer output representing either the summed intensities of all scanned ion intensities or a sample of the total current in the ion beam for each spectrum scan plotted against the corresponding spectrum number. Generally, it can be correlated with a flame ionization detector gas chromatogram.

3.2.5.1 *Discussion*—There are many synonyms in common use for RGC. These include: total ionization plot, total ionization current trace, reconstructed ion chromatogram, total ion current profile, and total ion chromatogram.

3.2.6 *reference compounds, n*—these are authentic materials used to obtain mass spectra, gas chromatographic retention data, and response factors. The operator can prepare the standards or they can be prepared commercially. Quality control solutions should be prepared independently from the calibration solutions. Quantitation methods may also require surrogate spiking solutions to determine extraction efficiency.

3.2.7 *semi-volatile organic compound, n*—an organic compound that can be separated from water by extraction, either liquid/liquid or solid phase, undergo volume adjustment, and be injected onto a GC. The compounds must elute from the column within its temperature range without alteration of the structure of the compound.

3.2.8 *tentative identification, n*—all identifications are considered tentative until confirmed by co-injection of an authentic reference compound showing identical retention time and similar mass spectra. (Tentative identification based on library matches only are subjected to false positives.)

⁵ Available from United States Environmental Protection Agency (EPA), William Jefferson Clinton Bldg., 1200 Pennsylvania Ave., NW, Washington, DC 20460, <http://www.epa.gov>.

⁶ Available from National Technical Information Service (NTIS), 5301 Shawnee Rd., Alexandria, VA 22312, <http://www.ntis.gov>.

⁷ SW 846 can be found online at <https://www.epa.gov/hw-sw846>.