

# Standard Test Method for Determination of Aldicarb, Aldicarb Sulfone, Aldicarb Sulfoxide, Carbofuran, Methomyl, Oxamyl, and Thiofanox in Water by Liquid Chromatography/Tandem Mass Spectrometry (LC/MS/MS)<sup>1</sup>

This standard is issued under the fixed designation D7645; 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 ( $\varepsilon$ ) indicates an editorial change since the last revision or reapproval.

# 1. Scope

1.1 This procedure covers the determination of aldicarb, aldicarb sulfone, aldicarb sulfoxide, carbofuran, methomyl, oxamyl, and thiofanox (referred to collectively as carbamates in this test method) in water by direct injection using liquid chromatography (LC) and detected with tandem mass spectrometry (MS/MS). These analytes are qualitatively and quantitatively determined by this test method. This test method adheres to multiple reaction monitoring (MRM) mass spectrometry.

1.2 The Detection Verification Level (DVL) and Reporting Range for the carbamates are listed in Table 1.

1.2.1 The DVL is required to be at a concentration at least 3 times below the Reporting Limit (RL) and have a signal/noise ratio greater than 3:1. Fig. 1 displays the signal/noise ratios of the primary single reaction monitoring (SRM) transitions, and Fig. 2 displays the confirmatory SRM transitions at the DVLs for the carbamates.

1.2.2 The reporting limit is the concentration of the Level 1 calibration standard as shown in Table 2 for the carbamates.

1.3 *Units*—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 and health practices and determine the applicability of regulatory limitations prior to use.

# 2. Referenced Documents

- 2.1 ASTM Standards:<sup>2</sup>
- D1129 Terminology Relating to Water
- D1193 Specification for Reagent Water
- D2777 Practice for Determination of Precision and Bias of Applicable Test Methods of Committee D19 on Water
- D3694 Practices for Preparation of Sample Containers and for Preservation of Organic Constituents
- D3856 Guide for Management Systems in Laboratories Engaged in Analysis of Water
- D4841 Practice for Estimation of Holding Time for Water Samples Containing Organic and Inorganic Constituents
- D5847 Practice for Writing Quality Control Specifications for Standard Test Methods for Water Analysis
- E2554 Practice for Estimating and Monitoring the Uncertainty of Test Results of a Test Method Using Control Chart Techniques
- 2.2 Other Documents:<sup>3</sup>
- EPA Publication SW-846 Test Methods for Evaluating Solid Waste, Physical/Chemical Methods
- EPA Method 531 Measurement of *N*-Methyl Carbamoyloximes and *N*-Methyl Carbamates in Drinking Water by Direct Aqueous Injection HPLC with Post Column Derivatization
- EPA Method 531.2 Measurement of *N*-Methylcarbamoyloximes and *N*-Methylcarbamates in Water by Direct Aqueous Injection HPLC with Postcolumn Derivatization

<sup>&</sup>lt;sup>1</sup> This test method 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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<sup>&</sup>lt;sup>2</sup> 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.

<sup>&</sup>lt;sup>3</sup> Available from National Technical Information Service (NTIS), U.S. Department of Commerce, 5285 Port Royal Road, Springfield, VA, 22161 or at http:// www.epa.gov/epawaste/hazard/testmethods/index.htm.

#### TABLE 1 Detection Verification Level and Reporting Range

Analyte	DVL (ng/L)	Reporting Range (µg/L)
Aldicarb	250	1-100
Aldicarb Sulfone	250	1-100
Aldicarb Sulfoxide	250	1-100
Carbofuran	250	1-100
Methomyl	250	1-100
Oxamyl	250	1-100
Thiofanox	250	1-100

EPA Method 538 Determination of Selected Organic Contaminants in Drinking Water by Direct Aqueous Injection-Liquid Chromatography/Tandem Mass Spectrometry (DAI-LC/MS/MS)

#### 3. Terminology

3.1 Definitions:

3.1.1 For definitions of terms used in this standard, refer to Terminology D1129.

3.2 Definitions of Terms Specific to This Standard:

3.2.1 *carbamates, n*—in this test method, aldicarb, aldicarb sulfone, aldicarb sulfoxide, carbofuran, methomyl, oxamyl, and thiofanox collectively.

3.2.2 *detection verification level, DVL, n*—a concentration that has a signal/noise ratio greater than 3:1 and is at least 3 times below the Reporting Limit (RL).

3.2.3 *independent reference material, IRM, n*—a material of known purity and concentration obtained either from the National Institute of Standards and Technology (NIST) or other reputable supplier. The IRM shall be obtained from a different lot of material than is used for calibration.

3.3 Acronyms:

3.3.1 CCC, n-Continuing Calibration Check

3.3.2 IC, n-Initial Calibration

3.3.3 *LC*, *n*—Liquid Chromatography

3.3.4 *LCS/LCSD*, *n*—Laboratory Control Sample/ Laboratory Control Sample Duplicate

3.3.5 MeOH, n-Methanol

3.3.6 *mM*, *n*—millimolar,  $1 \times 10^{-3}$  moles/L

3.3.7 MRM, n-Multiple Reaction Monitoring

3.3.8 MS/MSD, n-Matrix Spike/Matrix Spike Duplicate

3.3.9 NA, adj—Not Available

3.3.10 ND, n-non-detect

3.3.11 *P&A*, *n*—Precision and Accuracy

3.3.12 PPB, n-parts per billion

3.3.13 PPT, *n*—parts per trillion

3.3.14 QA, adj-Quality Assurance

3.3.15 QC, adj-Quality Control

3.3.16 RL, n-Reporting Limit

3.3.17 RSD, n-Relative Standard Deviation

3.3.18 *RT*, *n*—Retention Time

3.3.19 SDS, n-Safety Data Sheets

- 3.3.20 SRM, n-Single Reaction Monitoring
- 3.3.21 SS, *n*—Surrogate Standard
- 3.3.22 TC, n—Target Compound

3.3.23  $\mu M$ , *n*—micromolar, 1 × 10<sup>-6</sup> moles/L

3.3.24 VOA, n-Volatile Organic Analysis

## 4. Summary of Test Method

4.1 This is a performance-based method, and modifications are allowed to improve performance.

4.2 For carbamate analysis, samples are shipped to the lab acidified between  $0^{\circ}$ C and  $6^{\circ}$ C and analyzed within 14 days of collection. In the lab, the samples are spiked with surrogates, filtered using a syringe driven filter unit, and analyzed directly by LC/MS/MS.

4.3 The carbamates, methomyl- ${}^{13}C_2$ ,  ${}^{15}N$  (surrogate) and carbofuran- ${}^{13}C_6$  (surrogate) are identified by retention time and two SRM transitions. The target analytes and surrogate are quantitated using the primary SRM transitions utilizing an external calibration. The final report issued for each sample lists the concentration of carbamates and the surrogate recoveries.

## 5. Significance and Use

5.1 This test method has been developed by U.S. EPA Region 5 Chicago Regional Laboratory (CRL).

5.2 The *N*-methyl carbamate (NMC) pesticides: aldicarb, carbofuran, methomyl, oxamyl, and thiofanox have been identified by EPA as working through a common mechanism. These affect the nervous system by reducing the ability of enzymes. Enzyme inhibition was the primary toxicological effect of regulatory concern to EPA in assessing the NMC's food, drinking water, and residential risks. In most of the country, NMC residues in drinking water sources are at levels that are not likely to contribute substantially to the multi-pathway cumulative exposure. Shallow private wells extending through highly permeable soils into shallow, acidic ground water represent what the EPA believes to be the most vulnerable drinking water. Aldicarb sulfone and aldicarb sulfoxide are breakdown products of aldicarb and should also be monitored due to their toxicological effects.<sup>4</sup>

5.3 This test method has been investigated for use with reagent, surface, and drinking water for the selected carbamates: aldicarb, aldicarb sulfone, aldicarb sulfoxide, carbofuran, methomyl, oxamyl, and thiofanox.

#### 6. Interferences

6.1 Method interferences may be caused by contaminants in solvents, reagents, glassware, and other apparatus producing discrete artifacts or elevated baselines. All of these materials are demonstrated to be free from interferences by analyzing laboratory reagent blanks under the same conditions as samples.

<sup>&</sup>lt;sup>4</sup> Additional information about Carbamate pesticides area available from United States Environmental Protection Agency (EPA), http://www.epa.gov.

FIG. 1 Example Primary SRM Chromatograms Signal/Noise Ratios

7.75

8.00

8.25

8.50

8.75

9.00

9.25

S/N:PtP=11.82

5.75

6.00

100 5/N:PtF=11.02 5.68 4:5.76 5.92 Aldicarb Sulfoxide, 250 ppt, S/N = 12

6.50

6.75

7.00

7.25

7.50

6.25

1: MRM of 2 Channels ES+

9.50

207.1 > 131.9

1.19e4 - Time

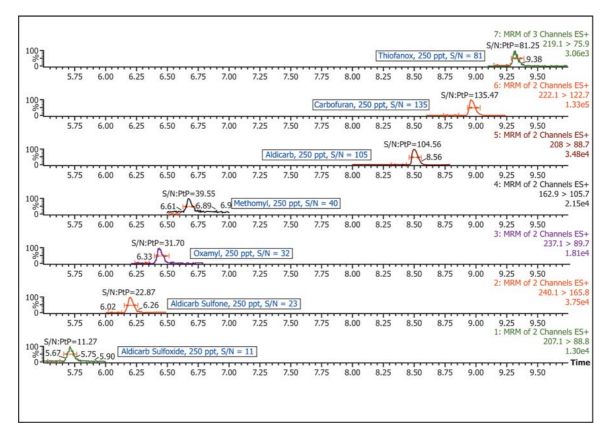
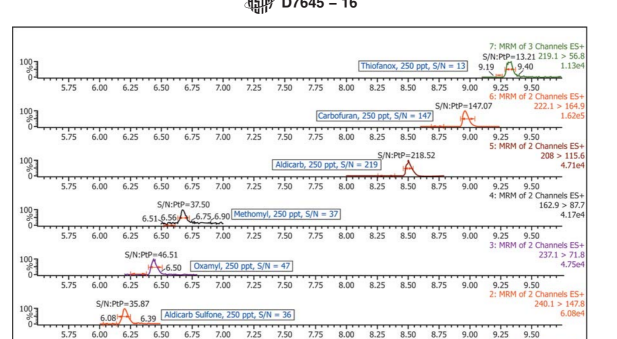


FIG. 2 Example Confirmatory SRM Chromatograms Signal/Noise Ratios



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