



Standard Test Method for Determination of Diisopropyl Methylphosphonate, Ethyl Methylphosphonic Acid, Isopropyl Methylphosphonic Acid, Methylphosphonic Acid and Pinacolyl Methylphosphonic Acid in Soil by Pressurized Fluid Extraction and Analyzed by Liquid Chromatography/Tandem Mass Spectrometry¹

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1. Scope

1.1 This procedure covers the determination of Diisopropyl Methylphosphonate (DIMP), Ethyl Methylphosphonic Acid (EMPA), Isopropyl Methylphosphonic Acid (IMPA), Methylphosphonic Acid (MPA) and Pinacolyl Methylphosphonic Acid (PMPA), referred to collectively as organophosphonates (OPs) in this test method, in soil. This method is based upon solvent extraction of a soil by pressurized fluid extraction (PFE). The extract is filtered and analyzed by liquid chromatography/tandem mass spectrometry (LC/MS/MS). OPs are qualitatively and quantitatively determined by this method.

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

1.3 The Method Detection Limit² (MDL), electrospray ionization (ESI) mode and Reporting Range³ for the OPs are listed in [Table 1](#).

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.*

¹ This test method is under the jurisdiction of ASTM Committee E54 on Homeland Security Applications and is the direct responsibility of Subcommittee E54.03 on Decontamination.

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² The MDL is determined following the Code of Federal Regulations, 40 CFR Part 136, Appendix B utilizing solvent extraction of soil by PFE. A detailed process determining the MDL is explained in the reference and is beyond the scope of this Standard to be explained here.

³ Reporting range concentrations are calculated from [Table 4](#) concentrations assuming a 100 μ L injection of the lowest and highest level calibration standards with a 40 mL final extract volume of a 10 gram soil sample. Volume variations will change the reporting limit and ranges. The reporting limit (RL), lowest concentration of the reporting range, is calculated from the concentration of the Level 1 calibration standard as shown in [Table 4](#).

2. Referenced Documents

2.1 *ASTM Standards*:⁴

D653 [Terminology Relating to Soil, Rock, and Contained Fluids](#)

D1193 [Specification for Reagent Water](#)

D3694 [Practices for Preparation of Sample Containers and for Preservation of Organic Constituents](#)

D3740 [Practice for Minimum Requirements for Agencies Engaged in Testing and/or Inspection of Soil and Rock as Used in Engineering Design and Construction](#)

D3856 [Guide for Management Systems in Laboratories Engaged in Analysis of Water](#)

E2554 [Practice for Estimating and Monitoring the Uncertainty of Test Results of a Test Method Using Control Chart Techniques](#)

2.2 *Other Documents*:

EPA publication SW-846 [Test Methods for Evaluating Solid Waste, Physical/Chemical Methods](#)⁵

40 CFR Part 136 [The Code of Federal Regulations, Appendix B](#)⁶

3. Terminology

3.1 *Definitions*:

3.1.1 *analytical column, n*—the particles of the solid stationary phase fill the whole inside volume of a tube (column) that the mobile phase passes through using the pressure generated by the liquid chromatography system.

⁴ 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 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>

⁶ Available from U.S. Government Printing Office Superintendent of Documents, 732 N. Capitol St., NW, Mail Stop: SDE, Washington, DC 20401, <http://www.access.gpo.gov>.

TABLE 1 Method Detection Limit and Reporting Range

Analyte	ESI Mode	MDL (PPB)	Reporting Range (PPB)
Diisopropyl methylphosphonate	Positive	2.7	40-2000
Ethyl methylphosphonic acid	Negative	2.3	40-2000
Ethyl methylphosphonic acid	Positive	1.3	40-2000
Isopropyl methylphosphonic acid	Negative	5.7	40-2000
Isopropyl methylphosphonic acid	Positive	2.8	40-2000
Methylphosphonic acid	Positive	8.7	40-2000
Pinacolyl methylphosphonic acid	Negative	5.3	40-2000

3.1.2 *filter unit, n*—in this standard, a filter that is supported with an inert housing to the solvents as described in Section 7 of this standard.

3.1.3 *filtration device, n*—a device used to remove particles from the extract that may clog the liquid chromatography system. Described in section 7.3 of this standard.

3.1.4 *glass fiber filter, n*—A porous glass fiber material onto which solid particles present in the extraction fluid, which flows through it, are largely caught and retained, thus removing them from the extract.

3.1.5 *hypodermic syringe, n*—in this standard, a luer-lock-tipped glass syringe capable of holding a syringe-driven filter unit as described in section 7.3 of this standard.

3.1.6 *liquid chromatography (LC) system, n*—in this standard, a separation system using liquid as the mobile phase and a stationary phase packed into a column. The use of small particles packed inside a column and a high inlet pressure enables the separation of components in a mixture.

3.1.7 *organophosphonates (OPs), n*—in this test method, Diisopropyl Methylphosphonate (DIMP), Ethyl Methylphosphonic Acid (EMPA), Isopropyl Methylphosphonic Acid (IMPA), Methylphosphonic Acid (MPA) and Pinacolyl Methylphosphonic Acid (PMPA) collectively.

3.1.8 *pressurized fluid extraction, n*—the process of transferring the analytes of interest from the solid matrix, a soil, into the extraction solvent using pressure and elevated temperature.

3.1.9 *reporting range, n*—the quantitative concentration range for an analyte in this standard.

3.1.10 *tandem mass spectrometer, n*—an arrangement in which ions are subjected to two sequential stages of analysis according to the quotient mass/charge.

3.2 Abbreviations:

3.2.1 *DIMP*—diisopropyl methylphosphonate

3.2.2 *EMPA*—ethyl methylphosphonic acid

3.2.3 *IMPA*—isopropyl methylphosphonic acid

3.2.4 *LC*—liquid chromatography

3.2.5 *LCS/LCSD*—laboratory control spike/laboratory control spike duplicate

3.2.6 *mM*—millimolar, 1×10^{-3} moles/L

3.2.7 *MPA*—methylphosphonic acid

3.2.8 *MRM*—multiple reaction monitoring

3.2.9 *MS*—matrix spike

3.2.10 *NA*—not applicable

3.2.11 *ND*—non-detect

3.2.12 *PFE*—pressurized fluid extraction

3.2.13 *PMPA*—pinacolyl methylphosphonic acid

3.2.14 *PPB*—parts per billion

3.2.15 *QC*—quality control

3.2.16 *SD*—standard deviation

3.2.17 *SRM*—single reaction monitoring

3.2.18 *VOA*—volatile organic analysis

4. Summary of Test Method

4.1 For OPs soil analysis, samples are shipped to the lab between 0°C and 6°C. The samples are to be extracted, filtered and analyzed by LC/MS/MS within 7 days of collection.

4.2 The OPs and the surrogates (diisopropyl methylphosphonate-D₁₄, pinacolyl methylphosphonic acid-¹³C₆ and methylphosphonic acid-D₃) are identified by retention time and one SRM transition. The target analytes and surrogates are quantitated using the SRM transitions utilizing an external calibration. The final report issued for each sample lists the concentration of each organophosphonate target compound and each surrogate recovery.

5. Significance and Use

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

5.1.1 Due to the rapid development of newer instrumentation and column chemistries, changes to the analysis described in this standard are allowed as long as better or equivalent performance data result. Any modifications shall be documented and performance data generated. The user of the data generated by this Standard shall be made aware of these changes and given the performance data demonstrating better or equivalent performance.

5.2 Organophosphate pesticides affect the nervous system by disrupting the enzyme that regulates acetylcholine, a neurotransmitter. They were developed during the early 19th century, but their effects on insects, which were similar to their effects on humans, were discovered in 1932. Some are poisonous and were used as chemical weapon agents. Organophosphate pesticides are usually not persistent in the environment.^{7,8}

5.3 This test method is for the analysis of selected organophosphorous based pesticide degradation products.

5.4 This method has been investigated for use with various soils.

6. Interferences

6.1 Method interferences may be caused by contaminants in solvents, reagents, glassware, and other apparatus producing

⁷ Additional information about organophosphate pesticides is available on the Internet at <http://www.epa.gov> (2011).

⁸ Additional information about chemical weapon agents is available on the Internet at <http://www.opcw.org> (2011).

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.

6.2 All reagents and solvents shall be of pesticide residue purity or higher to minimize interference problems.

6.3 Matrix interferences may be caused by contaminants that are co-extracted from the sample. The extent of matrix interferences can vary considerably from sample source depending on variations of the sample matrix.

7. Apparatus

7.1 LC/MS/MS System:

7.1.1 *Liquid Chromatography (LC) System*^{9,10}—A complete LC system is required in order to analyze samples. A LC system that is capable of performing at the flows, pressures, controlled temperatures, sample volumes, and requirements of the standard shall be used.

7.1.2 *Analytical Column*^{11,10}—A column that achieves adequate resolution shall be used. The retention times and order of elution may change depending on the column used and need to be monitored. A reverse-phase analytical column that combines the desirable characteristics of a reversed-phase HPLC column with the ability to separate polar compounds was used to develop this test method. MPA elutes early in the chromatograph, at approximately 2 minutes, which is just beyond the instrument void volume of 1.5 minutes. A column is required that elutes MPA after the instrument void volume.

7.1.3 *Tandem Mass Spectrometer (MS/MS) System*^{12,10}—A MS/MS system capable of multiple reaction monitoring (MRM) analysis or any system that is capable of performing at the requirements in this standard shall be used.

7.2 Pressurized Fluid Extraction Device (PFE)^{13,10}:

7.2.1 A PFE system was used for this test method with appropriately-sized extraction cells. Cells are available that will accommodate the 10 g sample sizes used in this test method. Cells shall be made of stainless steel or other material capable of withstanding the pressure requirements (≥ 2000 psi) necessary for this procedure. A pressurized fluid extraction device shall be used that can meet the necessary requirements in this test method.

⁹ A Waters Acquity UPLC H-Class System was used to develop this test method and generate the precision and bias data presented in Section 16. The sole source of supply known to the committee at this time is Waters Corporation, Milford, MA 01757.

¹⁰ If you are aware of alternative suppliers, please provide this information to ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the responsible technical committee,¹ which you may attend.

¹¹ A Waters-Atlantis® dC18, 150 mm \times 2.1 mm, 3 μ m particle size, was used to develop this test method and generate the precision and bias data presented in Section 16. The sole source of supply known to the committee at this time is Waters Corporation, Milford, MA 01757.

¹² A Waters Quattro micro™ API mass spectrometer was used to develop this test method and generate the precision and bias data presented in Section 16. The sole source of supply known to the committee at this time is Waters Corporation, Milford, MA 01757.

¹³ A Dionex Accelerated Solvent Extraction (ASE® 200) system with appropriately sized extraction cells was used to develop this test method and generate the precision and bias data presented in Section 16. The sole source of supply known to the committee at this time is Dionex Corporation, Sunnyvale, CA 94088.

7.2.2 *Glass Fiber Filters*.^{14,10}

7.2.3 *Amber VOA Vials*—40 mL for sample extracts and 60 mL for PFE.

7.3 Filtration Device:

7.3.1 *Hypodermic Syringe*—A luer-lock tip glass syringe capable of holding a syringe driven filter unit.

7.3.1.1 A 50 mL Lock Tip Glass Syringe size is recommended since a 40 mL sample extract may result.

7.3.2 *Filter Unit*^{15,10}—Filter units of polyvinylidene fluoride (PVDF) were used to filter the PFE extracts.

7.3.2.1 *Discussion*—A filter unit shall be used that meets the requirements of the test method.

8. Reagents and Materials

8.1 *Purity of Reagents*—High Performance Liquid Chromatography (HPLC) pesticide residue analysis and spectrophotometry grade chemicals shall be used in all tests. Unless indicated otherwise, it is intended that all reagents shall conform to the Committee on Analytical Reagents of the American Chemical Society.¹⁶ Other reagent grades may be used provided they are first determined to be of sufficiently high purity to permit their use without affecting the accuracy of the measurements.

8.2 *Purity of Water*—Unless otherwise indicated, references to water shall mean reagent water conforming to ASTM Type I of Specification D1193. It must be demonstrated that this water does not contain contaminants at concentrations sufficient to interfere with the analysis.

8.3 *Gases*—Nitrogen (purity $\geq 97\%$) and Argon (purity $\geq 99.999\%$).

8.4 Acetonitrile (CH₃CN, CAS # 75-05-8).

8.5 2-Propanol (C₃H₈O, CAS # 67-63-0).

8.6 Methanol (CH₃OH, CAS # 67-56-1).

8.7 Formic Acid (HCO₂H, $\geq 95\%$, CAS # 64-18-6).

8.8 Diisopropyl Methylphosphonate (C₇H₁₇O₃P, DIMP, CAS # 1445-75-6).

8.9 Ethyl Methylphosphonic Acid (C₃H₉O₃P, EMPA, CAS # 1832-53-7).

8.10 Isopropyl Methylphosphonic Acid (C₄H₁₁O₃P, IMPA, CAS # 1832-54-8).

8.11 Methylphosphonic Acid (CH₃O₃P, MPA, CAS # 993-13-5).

¹⁴ Whatman Glass Fiber Filters 19.8 mm, Part # 047017, specially designed for the PFE system,¹³ were used to develop this test method and generate the precision and bias data presented in Section 16. The sole source of supply known to the committee at this time is Dionex Corporation, Sunnyvale, CA 94088.

¹⁵ Millex®-GV Syringe Driven Filter Units PVDF 0.22 μ m (Catalog # SLGV033NS) were used to develop this test method and generate the precision and bias data presented in Section 16. The sole source of supply known to the committee at this time is Millipore Corporation.

¹⁶ *Reagent Chemicals, American Chemical Society Specifications*, American Chemical Society, Washington, D.C. For Suggestions on the testing of reagents not listed by the American Chemical Society, see *Annual Standards for Laboratory Chemicals*, BDH Ltd., Poole, Dorset, U.K., and the *United States Pharmacopeia and National Formulators*, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.

8.12 Pinacolyl Methylphosphonic Acid ($C_7H_{17}O_3P$, PMPA, CAS # 616-52-4).

8.13 Diisopropyl Methylphosphonate- D_{14} ($C_7H_3D_{14}O_3P$, DIMP- D_{14} , Unlabeled CAS # 1445-75-6).

8.13.1 DIMP- D_{14} represents deuterium labeled diisopropyl methylphosphonate where the two isopropyl moieties contain all 2H .

8.14 Methylphosphonic Acid- D_3 ($CH_2D_3O_3P$, MPA- D_3 , Unlabeled CAS # 993-13-5).

8.14.1 MPA- D_3 represents deuterium labeled methylphosphonic acid where the methyl moiety contains all 2H .

8.15 Pinacolyl Methylphosphonic Acid- $^{13}C_6$ ($C_7H_{17}O_3P$, PMPA- $^{13}C_6$, Unlabeled CAS # 616-52-4).

8.15.1 PMPA- $^{13}C_6$ represents ^{13}C labeled pinacolyl methylphosphonic acid where all the trimethylpropyl carbon atoms are uniformly labeled ^{13}C .

8.16 Ottawa Sand (CAS # 14808-60-7) or equivalent.

8.17 Drying Agent.^{17,10}

9. Hazards

9.1 Normal laboratory safety applies to this method. Analysts shall wear safety glasses, gloves, and lab coats when working in the lab. Analysts shall review the Material Safety Data Sheets (MSDS) for all reagents used in this test method and shall be fully trained to perform this test method.

10. Glassware Washing, Sampling and Preservation

10.1 *Glassware Washing*—All glassware is washed in hot tap water with a detergent and rinsed in hot water conforming to ASTM Type I of Specification **D1193**. The glassware is then dried and heated in an oven at 250°C for 15 to 30 minutes. All glassware is subsequently cleaned with acetone and methanol, respectively.

10.2 *Sampling*—Grab samples must be collected in pre-cleaned glass jars with polytetrafluoroethylene (PTFE) lined caps demonstrated to be free of interferences. This test method requires at least a 10 g sample size per analysis. A 100 g sample amount should be collected to allow for quality control samples and re-analysis. Field blanks are needed to follow conventional sampling practices.

10.3 *Preservation*—Store samples between 0°C and 6°C from the time of collection until analysis. Analyze the samples within 7 days of collection. If the samples are above 6°C when received or during storage or not analyzed within 7 days of collection, the data are qualified estimated and noted in the case narrative that accompanies the data.

11. Preparation of LC/MS/MS

11.1 *LC Operating Conditions Used to Develop This Test Method*⁹:

¹⁷ Varian – Chem Tube – Hydromatrix®, 1kg (Part # 198003) was used to develop this test method and generate the precision and bias data presented in Section 16 by recommendation of the PFE manufacturer. The sole source of supply known to the committee at this time is Agilent Technologies, Inc., 5301 Stevens Creek Blvd., Santa Clara, CA 95051. (Note: Some drying agents have been shown to clog PFE transfer lines.)

TABLE 2 Gradient Conditions for Liquid Chromatography

Time (min)	Flow (μL/min)	Percent CH_3CN	Percent Water	Percent 2% Formic Acid in Water
0	300	0	95	5
4	300	0	95	5
5	300	45	50	5
9	300	45	50	5
10	300	95	0	5
13	300	95	0	5
14	300	0	95	5
20	300	0	95	5

11.1.1 Injection volumes of all calibration standards and samples are 100 μL and are composed of primarily water. The first sample analyzed after the calibration curve is a blank to ensure there is no carry-over. The gradient conditions for the liquid chromatograph are shown in **Table 2**.

11.1.2 *Temperatures*—Column, 30°C; Sample compartment, 15°C.

11.1.3 *Wash and Purge Solvent*—60% Acetonitrile/40% 2-Propanol, Pre- and Post Inject Wash Solvent: 6 Seconds.

11.1.4 Specific instrument manufacturer wash and purge specifications shall be followed in order to eliminate sample carry-over in the analysis.

11.2 Mass Spectrometer Parameters¹²:

11.2.1 To acquire the maximum number of data points per SRM channel while maintaining adequate sensitivity, the tune parameters shall be optimized according to the instrument. Each peak requires at least 10 scans per peak for adequate quantitation. This test method contains five target compounds and three surrogates which are in different SRM experiment windows in order to optimize the number of scans and sensitivity. Variable parameters regarding retention times, SRM transitions, and cone and collision energies are shown in **Table 3**. Mass spectrometer parameters used in the development of this method are listed below:

The instrument is set in the Electrospray source setting.
 Capillary Voltage: 3.5 kV
 Cone: Variable depending on analyte (**Table 3**)
 Extractor: 2 V
 RF Lens: 0.2 V
 Source Temperature: 120 °C
 Desolvation Temperature: 300 °C
 Desolvation Gas Flow: 700 L/hr
 Cone Gas Flow: 25 L/hr
 Low Mass Resolution 1: 14.0
 High Mass Resolution 1: 14.0
 Ion Energy 1: 0.8 V
 Entrance Energy: -1 V
 Collision Energy: Variable depending on analyte (**Table 3**)
 Exit Energy: 2 V
 Low Mass Resolution 2: 14
 High Mass resolution 2: 14
 Ion Energy 2: 1.0 V
 Multiplier: 650 V
 Gas Cell Pirani Gauge: 0.60 Pa
 Inter-Channel Delay: 0.02 s
 Inter-Scan Delay: 0.1 s if acquiring in one ESI mode, 0.4 s if acquiring in both.
 Repeats: 1
 Span: 0 Daltons
 Dwell: 0.1 s

TABLE 3 Retention Times, SRM Transitions, and Analyte-Specific Mass Spectrometer Parameters

Analyte	ESI Mode	Retention Time (min)	SRM Mass Transition (Parent >Product)	Cone Voltage (Volts)	Collision Energy (eV)
Diisopropyl methylphosphonate	Positive	8.8	181.2>139.1	25	6
Ethyl methylphosphonic acid	Negative	3.6	123.0>94.9	30	12
Ethyl methylphosphonic acid	Positive	3.6	125.0>96.9	25	10
Isopropyl methylphosphonic acid	Negative	7.5	137.0>94.9	28	13
Isopropyl methylphosphonic acid	Positive	7.5	139.1>96.9	25	7
Methylphosphonic acid	Positive	2.0	96.9>78.8	45	15
Pinacolyl methylphosphonic acid	Negative	8.6	179.1>94.9	35	18
DIMP-D ₁₄ (Surrogate)	Positive	8.8	195.2>147.1	23	7
PMPA- ¹³ C ₆ (Surrogate)	Negative	8.6	185.1>94.9	35	18
MPA-D ₃ (Surrogate)	Positive	2.0	99.9>81.8	40	15

12. Calibration and Standardization

12.1 The mass spectrometer shall be calibrated per manufacturer specifications before analysis. In order to obtain valid and accurate analytical values within the confidence limits, the following procedures shall be followed when performing the test method.

12.2 *Calibration and Standardization*—To calibrate the instrument, analyze eight calibration standards containing the eight concentration levels of the organophosphonates and surrogates prior to analysis as shown in [Table 4](#). A calibration stock standard solution is prepared from standard materials or purchased as certified solutions. Stock standard solution A (Level 8) containing the organophosphonates, diisopropyl methylphosphonate-D₁₄, pinacolyl methylphosphonic acid-¹³C₆ and methylphosphonic acid-D₃ is prepared at Level 8 concentration and aliquots of that solution are diluted to prepare Levels 1 through 7. The following steps will produce standards with the concentration values shown in [Table 4](#). The analyst is responsible for recording initial component weights carefully when working with pure materials and correctly carrying the weights through the dilution calculations. Calibration standards are not filtered.

12.2.1 Prepare stock standard solution A (Level 8) by adding to a 50 mL volumetric flask individual methanol solutions of the following: 250 µL of 100 µg/mL solutions of DIMP, EMPA, IMPA, MPA, PMPA, MPA-D₃ and PMPA-¹³C₆, and 25 µL of 1000 µg/mL of DIMP-D₁₄ and then dilute to 50 mL with water. The preparation of the Level 8 standard can be accomplished using different volumes and concentrations of stock solutions as is accustomed in the individual laboratory. Depending on stock concentrations prepared, the solubility at that concentration shall be ensured.

12.2.2 Aliquots of Solution A are then diluted with water to prepare the desired calibration levels in 2 mL amber glass LC vials at concentrations shown in [Table 4](#), calibration standards are not filtered. The calibration standard vials shall be used within 24 hours to ensure optimum results. Stock calibration standard solutions are replaced every 14 days if not previously discarded for quality control failure.

12.2.3 Inject each calibration standard and obtain its chromatogram. External calibration curves are generated from the calibration standards monitoring the SRM transition of each analyte. Calibration software is utilized to conduct the quantitation of the target analytes and surrogates. The SRM transition of each analyte is used for quantitation and confirmation. The

use of SRM transitions gives additional confirmation than by the selective ion monitoring technique because the parent ion is isolated and fragmented to the product ion.

12.2.4 The calibration software manual shall be consulted to use the software correctly. The quantitation method is set as an external calibration using the peak areas in ppb units. Concentrations may be calculated using the data system software to generate linear regression or quadratic calibration curves. Forcing the calibration curve through the origin is not recommended. Each calibration point used to generate the curve shall have a calculated percent deviation less than 30% from the generated curve. Refer to sections [12.2.4.1](#) and [12.2.4.2](#) to determine if linear or quadratic calibration curves may be used.

12.2.4.1 Linear calibration may be used if the coefficient of determination, r^2 , is >0.98 for the analyte. The point of origin is excluded, and a fit weighting of 1/X is used in order to give more emphasis to the lower concentrations. If one of the calibration standards other than the high or low point causes the r^2 of the curve to be <0.98, this point shall be re-injected or a new calibration curve shall be regenerated. If the low or high point is excluded, or both, minimally a five point curve is acceptable, the reporting range shall be modified to reflect this change.

12.2.4.2 Quadratic calibration may be used if the coefficient of determination, r^2 , is >0.99 for the analyte. The point of origin is excluded, and a fit weighting of 1/X is used in order to give more emphasis to the lower concentrations. If one of the calibration standards, other than the high or low, causes the curve to be <0.99, this point shall be re-injected or a new calibration curve shall be regenerated. If the low or high point is excluded, or both, a six point curve is acceptable using a quadratic fit. An initial eight point curve over the calibration range is suggested in the event that the low or high point must be excluded to obtain a coefficient of determination >0.99. In this event, the reporting range shall be modified to reflect this change.

12.2.5 The retention time window of the SRM transitions shall be within 5% of the retention time of the analyte in a midpoint calibration standard. A midpoint calibration standard is defined at or between Levels 4-6 in [Table 4](#) in this test method. If this is not the case, re-analyze the calibration curve to determine if there was a shift in retention time during the analysis and re-inject the sample. If the retention time is still incorrect in the sample, refer to the analyte as an unknown.

TABLE 4 Concentrations of Calibration Standards (PPB)

Analyte/Surrogate	LV 1	LV 2	LV 3	LV 4	LV 5	LV 6	LV 7	LV 8
Diisopropyl methylphosphonate	10	25	50	100	200	300	400	500
Ethyl methylphosphonic acid	10	25	50	100	200	300	400	500
Isopropyl methylphosphonic acid	10	25	50	100	200	300	400	500
Methylphosphonic acid	10	25	50	100	200	300	400	500
Pinacolyl methylphosphonic acid	10	25	50	100	200	300	400	500
DIMP-D ₁₄ (Surrogate)	10	25	50	100	200	300	400	500
PMPA- ¹³ C ₆ (Surrogate)	10	25	50	100	200	300	400	500
MPA-D ₃ (Surrogate)	10	25	50	100	200	300	400	500

12.2.5.1 The chromatographic peak shape for EMPA in the Nebraska soil was poor compared to the other soils tested. A blank soil, “unspiked soil”, and matrix spike soils were analyzed for each soil type for quality control purposes. The EMPA peak shape in the calibration curve and Nebraska soil is shown in Fig. X1.1 in both the ESI positive and negative modes. Monitoring the SRM transition for EMPA in both the positive and negative electrospray modes resulted in similar chromatographic peak shape in the matrix spike sample. A comparison to an unspiked soil shall be made by the analyst to determine presence or absence of the target analyte in soils where chromatographic peak shape may be an issue. Data for EMPA and IMPA are collected in both the electrospray positive and negative modes providing more information for an analyst to make such a decision in those cases. The PMPA, DIMP, IMPA and MPA chromatographic peak shapes were shown to be less affected by the various matrices tested.

12.2.6 A midpoint calibration check standard shall be analyzed at the end of each batch of 20 samples or within 24 hours after the initial calibration curve was generated. This end calibration check shall be the same calibration standard that was used to generate the initial curve. The results from the end calibration check standard shall have a percent deviation less than 35% from the calculated concentration for the target analytes and surrogates. If the results are not within these criteria, the problem shall be corrected and either all samples in the batch shall be re-analyzed against a new calibration curve or the affected results shall be qualified with an indication that they do not fall within the performance criteria of the test method. If the analyst inspects the vial containing the end calibration check standard and notices that the sample evaporated affecting the concentration, a new end calibration check standard shall be made and analyzed. If this new end calibration check standard has a percent deviation less than 35% from the calculated concentration for the target analytes and surrogates, the results shall be reported unqualified if all other quality control parameters are acceptable.

12.3 If a laboratory has not performed the test before or if there has been a major change in the measurement system, for example: new analyst or new instrument, perform a precision and bias study to demonstrate laboratory capability and verify that all technicians are adequately trained and follow relevant safety procedures.

12.3.1 Analyze at least four replicates of a sample containing the target compounds and surrogates at a concentration between 200 and 800 ppb in Ottawa sand. This test method was tested at 400 ppb. Each replicate shall be taken through the

complete analytical test method including any sample preservation and pretreatment steps.

12.3.2 Calculate the mean (average) percent recovery and relative standard deviation (RSD) of the four values and compare to the acceptable ranges of the quality control (QC) acceptance criteria for the Initial Demonstration of Performance in Table 5.

12.3.3 This study shall be repeated until the single operator precision and mean recovery are within the limits in Table 5.

12.3.4 The QC acceptance criteria for the Initial Demonstration of Performance in Table 5 are preliminary until a collaborative study is conducted. Single-laboratory data is shown in the Precision and Bias Section. The analyst shall be aware that the performance data generated from single-laboratory data tend to be significantly tighter than those generated from multi-laboratory data. The laboratory shall generate its own in-house QC acceptance criteria which meet or exceed the criteria in this test method. References on how to generate QC acceptance criteria are ASTM Standard E2554 or Method 8000B in EPA publication SW-846.

12.4 Surrogate Spiking Solution:

12.4.1 A surrogate standard solution containing MPA-D₃, PMPA-¹³C₆ and DIMP-D₁₄ is added to each 10 g soil sample. A stock surrogate spiking solution is prepared in methanol at 40 ppm for MPA-D₃, PMPA-¹³C₆ and DIMP-D₁₄. The surrogates are added to each sample to achieve a concentration of 400 ppb (that is, 100 µL of a 40 ppm methanol solution containing MPA-D₃, PMPA-¹³C₆ and DIMP-D₁₄ is added to a 10 g soil sample). The result obtained for the surrogate recovery shall fall within the limits of Table 5. If the limits are not met, the affected results shall be qualified with an indication that they do not fall within the performance criteria of the test method.

12.5 Method Blank:

12.5.1 Analyze a sample blank, Ottawa sand, with each batch of 20 or fewer samples. The concentration of target analytes found in the blank must be three times below the reporting limit. If the concentration of target analytes is found above this level, analysis of samples is halted until the contamination is eliminated, and a blank shows no contamination at or above this level or the results shall be qualified with an indication that there is a blank contamination and report the concentration found in the blank sample.

12.6 Laboratory Control Sample (LCS):

12.6.1 To ensure that the test method is in control, analyze a LCS prepared with the OPs and surrogates at a concentration

TABLE 5 Quality Control Acceptance Criteria (Test Concentration at 400 ppb)

Analyte	ESI Mode	Initial Demonstration of Performance			Lab Control Sample	
		Recovery (%)		Precision	Recovery (%)	
		Lower Limit	Upper Limit	Maximum % RSD	Lower Limit	Upper Limit
Diisopropyl methylphosphonate	+	70	130	30	70	130
Ethyl methylphosphonic acid	+	70	130	30	70	130
Ethyl methylphosphonic acid	-	70	130	30	70	130
Isopropyl methylphosphonic acid	+	70	130	30	70	130
Isopropyl methylphosphonic acid	-	70	130	30	70	130
Methylphosphonic acid	+	50	150	40	50	150
Pinacolyl methylphosphonic acid	-	70	130	30	70	130
DIMP-D ₁₄ (Surrogate)	+	70	130	30	70	130
PMPA- ¹³ C ₆ (Surrogate)	-	70	130	30	70	130
MPA-D ₃ (Surrogate)	+	50	150	40	50	150

between 200 and 800 ppb. This test method was tested at 400 ppb. The LCS is prepared following the analytical method and analyzed with each batch of 20 samples or less. Prepare a stock matrix spiking solution in methanol containing EMPA, MPA, IMPA, DIMP and PMPA each at 40 ppm. An Ottawa sand sample is spiked with the matrix spiking solution to achieve a concentration of 400 ppb (that is, 100 µL of a 40 ppm methanol solution containing of DIMP, EMPA, IMPA, MPA and PMPA is added to a 10 g soil sample). The results obtained for the LCS shall fall within the limits in [Table 5](#).

12.6.2 If the results are not within these limits, analysis of samples is halted until the problem is corrected, and either all samples in the batch shall be re-analyzed or the results shall be qualified with an indication that they do not fall within the performance criteria of the test method.

12.7 Matrix Spike (MS):

12.7.1 To check for interferences in the specific matrix being tested, perform a MS on at least one sample from each batch of ten or fewer samples. This is accomplished by spiking the sample with a known concentration of OPs and following the analytical method. Prepare a stock matrix spiking solution in methanol containing EMPA, MPA, IMPA, DIMP and PMPA each at 40 ppm. Spiking 100 µL of this stock solution into 10 g of soil to yield a concentration of 400 ppb for EMPA, MPA, IMPA, PMPA and DIMP in the soil.

12.7.2 If the spiked concentration plus the background concentration exceeds that of the Level 8 calibration standard, the sample shall be diluted to a level near the midpoint of the calibration curve.

12.7.3 Calculate the percent recovery of the spike (P) using [Eq 1](#):

$$P = 100 \frac{|A(V_s + V) - BV_s|}{CV} \quad (1)$$

where:

- A = concentration found in spiked sample,
- B = concentration found in unspiked sample,
- C = concentration of analyte in spiking solution,
- V_s = volume of sample used,
- V = volume of spiking solution added, and
- P = percent recovery.

12.7.4 The percent recovery of the spike shall fall within the limits in [Table 6](#). If the percent recovery is not within these limits, a matrix interference may be present in the selected

sample, a matrix suppression or enhancement of the response or extraction efficiency of the analyte, or both, may be poor in the soil matrix. The results shall be qualified with an indication that they do not fall within the performance criteria of the test method. The recoveries of OPs in the matrix spike samples are required for all data generated and shall accompany the analytical results due to the variation in recoveries in the various soil matrices as shown in Precision and Accuracy Section 16. It has been demonstrated that in certain soil types, primarily clay, recoveries are low or, for MPA, less than the reporting limit (see Section 16).

12.7.4.1 Various extraction solvents and procedures were studied. The extraction procedures included PFE, sonication and tumbling. The solvents included water, methanol and acetonitrile in various combinations. The adjustment of pH was also investigated and included the use of ammonium hydroxide, acetic acid and sodium hydroxide. Water was shown to produce the overall best results in these studies.

12.7.5 The matrix spike/matrix spike duplicate (MS/MSD) limits in [Table 6](#) are preliminary until a collaborative study is completed. Matrix spike recovery data for six different soils is included in the Precision and Accuracy Section 16. The matrix spike recovery data is variable amongst the soils tested. The matrix variation between different soils may tend to generate significantly wider control limits than those generated by a single laboratory in one soil matrix. It is recommended that the laboratory generate its own in-house QC acceptance criteria which meet or exceed the criteria shown in [Table 6](#) in this test method.

12.7.5.1 The laboratory shall generate its own in-house QC acceptance criteria after the analysis of 15–20 matrix spike samples of a particular soil matrix. References on how to generate QC acceptance criteria are ASTM Standard [E2554](#) or Method 8000B in EPA publication SW-846.

12.8 Duplicate:

12.8.1 To check the precision of sample analyses, analyze a sample in duplicate with each batch of 10 or fewer samples. If the sample contains the analyte at a level greater than 5 times the detection limit of the method, the sample and duplicate may be analyzed unspiked; otherwise, an MSD shall be used.

12.8.2 Calculate the relative percent difference (RPD) between the duplicate values (or MS/MSD values) as shown in [Eq 2](#). Compare to the RPD limit in [Table 6](#).

TABLE 6 MS/MSD Quality Control Acceptance Criteria

Analyte/Surrogate	ESI Mode	Test Conc. (ppb)	MS/MSD		
			Recovery (%)		Precision
			Lower Limit	Upper Limit	Maximum RPD (%)
Diisopropyl methylphosphonate	+	400	30	130	30
Ethyl methylphosphonic acid	+	400	30	130	30
Ethyl methylphosphonic acid	-	400	30	130	30
Isopropyl methylphosphonic acid	+	400	30	130	30
Isopropyl methylphosphonic acid	-	400	30	130	30
Methylphosphonic acid	+	400	30	130	30
Pinacolyl methylphosphonic acid	-	400	30	130	30
DIMP-D _{1,4} (Surrogate)	+	400	30	130	30
PMPA- ¹³ C ₆ (Surrogate)	-	400	30	130	30
MPA-D ₃ (Surrogate)	+	400	30	130	30

$$RPD = \frac{|MSR - MSDR|}{(MSR + MSDR) \div 2} \times 100 \quad (2)$$

where:

RPD = relative percent difference,
MSR = matrix spike recovery, and
MSDR = matrix spike duplicate recovery.

12.8.3 If the result exceeds the precision limit, the batch shall be re-analyzed or the results shall be qualified with an indication that they do not fall within the performance criteria of the test method.

13. Pressurized Fluid Extraction Procedure

13.1 Mix the soil or sediment sample thoroughly, especially composite samples. Note the overall appearance of the sample; for example, how much water or liquid phase is present and whether foreign objects such as sticks, leaves, rocks, etc., are present. It is important to consult the client on how the samples should be processed. Decant and discard any water layer if the client wants only the solid portion analyzed; alternatively, if the client requires the analysis of both phases, then pour the liquid layer into a separate container, measure and conduct the appropriate extraction procedure. Prior to weighing, discard foreign objects, unless instructed otherwise by the client.

13.2 Three cell sizes applicable to this test method are available for the PFE System: 11, 22 and 33 mL. The 33 mL cell equals the volume of the largest Soxhlet thimble commonly used for this test method. In general, when choosing a cell size, select the smallest cell that holds enough sample to produce accurate extraction results. The 11 mL cell holds approximately 10 g, the 22 mL cell holds approximately 20 g, and the 33 mL cell holds approximately 30 g, all dry mass. Take into account any drying agent needed, which increases sample volume. When preparing the sample, make sure that the drying agent and sample are thoroughly mixed.

13.3 Weigh out samples into crucibles or evaporating dishes depending on anticipated contaminant levels and take into consideration any action levels or detection limits required by the client. Sample sizes are as follows: 5 g or less for high level concentrations of target analytes, 10 g for medium level, and 30 g for low level analysis, based upon a dry basis. This analysis is based upon a 10 gram sample size. Also, take into account the water content of the samples to determine the

amount of diatomaceous earth required to dry them sufficiently and not overload the extraction cell, refer to section 13.6. Be sure to include all QA/QC samples such as method blanks, laboratory control samples and matrix spike samples.

13.4 Spike each soil sample with a 100 µL of a 40 ppm methanol solution containing the surrogates.

13.5 For each matrix spike and LCS/LCSD, spike each sample with a 100 µL of a 40 ppm methanol solution containing the OPs.

13.6 All matrices shall be mixed with a drying agent¹⁷ before being loaded into the cells. The drying agent recommended by the PFE manufacturer was used in this test method. It dries samples quickly, provides a cleaner transfer of the mixtures to the cell, extracts well and prevents clogging of the frit in the end cap of the extraction cell, which normally occurs when sodium sulfate is used to dry samples. If the sample appears dry, use 4 g sample to 1 g diatomaceous earth. If the sample appears wet, use 4 g sample to 2 g diatomaceous earth. If the sample is a liquid, use 5 g sample to 3 g diatomaceous earth. Mix the sample with diatomaceous earth thoroughly in a small mortar or evaporating dish. Add diatomaceous earth and stir the mixture until a sandy texture is observed.

13.7 To prepare a 10 g sample, collect 22 or 33 mL PFE cells with appropriately sized caps.

NOTE 1—If the soil sample with the drying agent can fit into the smaller sized cell without packing it down, the smaller sized cell should be chosen. Do not pack the soil down into the cell, this will prevent effective extraction. Hand-tighten the main body of the extraction cell with a cell cap and insert a disposable glass fiber filter at the bottom of the cap. Place the prepared sample into each cell.

13.8 Fill any void volume in the cell with the drying agent or Ottawa sand. Assemble each extraction cell by hand-tightening the cell caps on each end. Do not use a wrench or other tool to tighten the cap. If the extraction cells are packed tightly, an over-pressurized condition can cause the system to shut down. Prior to using the cell caps, verify that the white O-rings are in place and in good condition. Check the polyether ether ketone (PEEK) seals inside the caps and replace if necessary.

13.9 Load the cells in numerical order and hang the cells vertically in the tray slots from their top caps. The bottom cap contains the glass fiber filter.

13.10 Load the rinse tubes into the rinse slots.

13.11 For each sample setup, load a labeled 60 mL collection vial into the corresponding vial tray position. The label or any markings shall be between 34 and 78 millimetres from the top of the collection vial or the solvent sensor will return an error when trying to read the solvent level in the vial, and the PFE will move onto the next row of the sequence. Prepare a method on the PFE using the following conditions (These parameters are based on the PFE system to develop this test method):

Pressure: 6.9×10^6 Pa
 Temperature: 50°C
 Preheat Time: 5 minutes
 Heat Time: 5 minutes
 Static Time: 5 minutes
 Flush Volume: 40%
 Purge Time: 60 seconds
 Static Cycles: 2
 Solvent: Water

13.12 If the type of solvent or solvent mixture in any of the bottles has changed or the PFE system has not been used recently, the lines shall be rinsed by pressing the “rinse” button on the control panel before use.

13.13 If the PFE system is run under method control, it will extract cells in numerical order. It will inject each extract into the corresponding receiving vial with the same number until all the cell slots have been loaded and extracted or until it cannot load two cells in a row. If it is run under schedule control, the PFE system will inject the extract(s) of each vial into the corresponding receiving vial(s) designated in the schedule.

13.14 The PFE system extract is then filtered using the filtration device described in section 7.3.

13.15 Begin sample analysis after a passing calibration curve is generated as described in Section 12. An order of analysis may be method blank(s), laboratory control sample(s), sample(s), duplicate(s), and matrix spike sample(s) followed by an end calibration check standard.

14. Calculation or Interpretation of Results

14.1 For quantitative analysis of MPA-D₃, PMPA-¹³C₆, DIMP-D₁₄, DIMP, EMPA, IMPA, MPA and PMPA, the SRM transitions are identified by comparison of retention times in the sample to those of the standards. External calibration curves are used to calculate the amount of target analytes and surrogates. Calculate the concentration in µg/kg (ppb) for each analyte. Organophosphates are reported if present at or above the reporting limit. If the concentration of the analyte is determined to be above the calibration range, the sample is diluted with reagent water to obtain a concentration near the midpoint of the calibration range and re-analyzed.

14.2 Data for EMPA and IMPA are collected both in ESI positive and ESI negative modes. Both ESI positive and ESI negative quantitative results shall be reported for all samples including method blanks, laboratory control samples and matrix spikes.

14.3 DIMP and PMPA may be overlapping in this test method. If there is overlap encountered due to poor chromatography which may be caused by a variation in elution solvent

concentrations or column degradation this is not a concern if the analyte specific SRM transitions are employed. DIMP and PMPA both have parent ion of 181.2 m/z, and both will produce the same product fragment under certain conditions. In ESI negative mode, PMPA will produce a parent ion of 179.1 m/z, but DIMP will not. Therefore, the optimum transition for PMPA is in the ESI negative mode using 179.1>94.9. In ESI positive mode, the DIMP parent ion 181.2 m/z will produce a 139.1 m/z product ion by loss of one isopropyl moiety, whereas PMPA is not able to lose that fragment. Therefore, the optimum SRM transition for DIMP is in the ESI positive mode using 181.2>139.1.

15. Report

15.1 Determine the results in units of µg/kg (ppb) in a soil sample. Calculate the concentration in the sample using the linear or quadratic calibration curve generated. All data that do not meet the specifications in the test method shall be appropriately qualified.

16. Precision and Bias

16.1 The determination of precision and bias was conducted by the United States Environmental Protection Agency (US EPA), Chicago Regional Laboratory (CRL) in a single-laboratory study. A multi-laboratory validation is being planned. The goal of the test method will be to generate multi-laboratory participants within the next 5 years to enable a full validation study to produce a research report.

16.2 This test method was used by the US EPA CRL on Ottawa Sand, Nebraska, Georgia, and 4 ASTM reference soils (CH-1, ML-1, CL-1 and SP-1).¹⁸ The characterization data for the Nebraska and Georgia soils are in Appendix Table 1. ASTM reference soil CH-1 is Fat Clay (CH)- Vicksburg Buckshot Clay, ASTM reference soil ML-1 is Silt (ML)-Vicksburg Silt, ASTM reference soil CL-1 is Lean Clay (CL)-Annapolis Clay and ASTM reference soil SP-1 is Sand (SP)-Frederick Sand. The samples were spiked with target OPs and surrogates to obtain a 400 ppb concentration of each as described in Section 12. Tables 7 and 8 contain the recoveries for the surrogates and Tables 9-15 contain the recoveries for the OPs spiked at 400 ppb. Tables 16-22 contain the OPs recovery data for different soil matrices spiked at 1600 ppb.

16.3 Recoveries for some of the organophosphonates were poor in certain soils.

16.3.1 MPA was the poorest performer in clay soils. There was no recovery for MPA in Georgia, Nebraska and CL-1 soils at the 400 ppb or 1600 ppb spike level except for a 12.3 % recovery for one of the Georgia 1600 ppb spiked samples. MPA spiked at 400 ppb in the CH-1 soil type resulted in no recovery and average recovery of 12.4% recovery at the 1600 ppb spike level.

16.3.2 EMPA recovery was poor in CL-1 spiked soils averaging ~23% recovery.

16.3.3 IMPA recovery was low in Nebraska, CH-1 and CL-1 soil types with an average recovery in those soils of ~27%.

¹⁸ Reference to the ASTM soils and soil reports can be found at: <http://www.durhamgeo.com/downloads/ASTM%20Soil%20Reports.html> (2011)

TABLE 7 Surrogate Recovery Data

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes											
	Sand			Georgia			Nebraska			ASTM Soil CL-1		
	MPA-D ₃ (ESI +)	PMPA- ¹³ C ₆ (ESI -)	DIMP-D ₁₄ (ESI +)	MPA-D ₃ (ESI +)	PMPA- ¹³ C ₆ (ESI -)	DIMP-D ₁₄ (ESI +)	MPA-D ₃ (ESI +)	PMPA- ¹³ C ₆ (ESI -)	DIMP-D ₁₄ (ESI +)	MPA-D ₃ (ESI +)	PMPA- ¹³ C ₆ (ESI -)	DIMP-D ₁₄ (ESI +)
Method Blank	608.0	384.2	460.4	<RL	370.0	364.0	< RL	237.6	240.0	<RL	91.7	190.8
1	505.6	385.8	473.5	<RL	362.5	346.9	< RL	237.2	281.0	<RL	58.2	134.8
2	493.1	377.6	403.1	<RL	354.6	364.3	< RL	239.6	284.2	<RL	58.4	149.9
3	495.1	386.5	424.6	<RL	350.5	356.3	< RL	240.0	289.2	<RL	43.6	130.9
4	453.0	375.5	427.5	<RL	349.9	355.4	< RL	226.2	265.3	<RL	27.6	189.6
5	227.7	363.2	413.6	<RL	347.9	350.9	< RL	210.6	260.0	<RL	66.8	165.6
6	457.4	371.4	436.3	<RL	355.8	335.1	< RL	225.3	264.6	<RL	72.3	155.5
7	468.2	361.2	402.7	<RL	362.8	358.5	< RL	237.3	303.2	<RL	32.0	130.3
Average Recovery:	463.5	375.7	430.2	-	356.8	353.9	-	231.7	273.4	-	56.3	155.9
Average % Recovery:	115.9	93.9	107.6	-	89.2	88.5	-	57.9	68.4	-	14.1	39.0
Standard Deviation:	107.1	9.9	25.7	-	7.7	9.6	-	10.3	19.8	-	21.4	24.5
% Relative SD	23.1	2.6	6.0	-	2.2	2.7	-	4.5	7.2	-	37.9	15.7

TABLE 8 Surrogate Recovery Data Continued

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes								
	ASTM Soil SP-1			ASTM Soil ML-1			ASTM Soil CH-1		
	MPA-D ₃ (ESI +)	PMPA- ¹³ C ₆ (ESI -)	DIMP-D ₁₄ (ESI +)	MPA-D ₃ (ESI +)	PMPA- ¹³ C ₆ (ESI -)	DIMP-D ₁₄ (ESI +)	MPA-D ₃ (ESI +)	PMPA- ¹³ C ₆ (ESI -)	DIMP-D ₁₄ (ESI +)
Method Blank	109.6	380.4	395.4	403.5	387.3	391.9	107.2	138.4	101.6
1	105.0	353.9	385.2	295.2	382.9	392.2	85.2	218.8	183.2
2	135.8	356.5	380.6	362.4	386.0	385.8	60.4	68.0	67.6
3	132.4	373.5	386.0	364.0	393.8	400.0	16.0	84.0	41.6
4	95.4	363.7	423.2	340.5	392.8	396.8	60.4	149.6	102.0
5	87.8	372.2	398.2	347.4	394.0	389.2	18.4	62.4	50.0
6	85.8	382.8	388.0	333.3	397.5	392.2	89.2	290.0	231.2
7	99.0	361.3	373.8	359.4	404.8	391.2	62.8	285.2	232.0
Average Recovery:	106.3	368.1	391.3	350.7	392.4	392.4	62.5	162.1	126.2
Average % Recovery:	26.6	92.0	97.8	87.7	98.1	98.1	15.6	40.5	31.5
Standard Deviation:	18.9	10.8	15.0	30.9	7.0	4.4	32.4	92.8	78.4
% Relative SD	17.8	2.9	3.8	8.8	1.8	1.1	51.9	57.3	62.2

TABLE 9 Organophosphates Recovery Data in Ottawa Sand (400 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes in Ottawa Sand							
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)	
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL	
1	419.5	411.7	375.2	417.0	378.8	374.6	424.1	
2	407.5	421.4	377.1	415.4	374.5	371.3	357.5	
3	423.7	379.0	387.2	415.6	383.2	387.0	367.9	
4	401.0	359.7	364.7	367.2	366.3	372.4	365.7	
5	160.6	335.2	347.3	361.9	364.2	361.7	354.6	
6	381.5	361.0	363.8	381.8	372.8	368.3	365.2	
7	380.7	377.5	351.2	374.7	359.0	360.5	342.7	
Average Recovery:	367.8	377.9	366.7	390.5	371.3	370.8	368.2	
Average % Recovery:	92.0	94.5	91.7	97.6	92.8	92.7	92.1	
Standard Deviation:	92.9	30.2	14.3	24.6	8.6	8.9	26.1	
% Relative SD	25.3	8.0	3.9	6.3	2.3	2.4	7.1	

16.3.4 Poor recovery for PMPA was found with CH-1 soil at ~33% and CL-1 soil at ~17%.

16.3.5 Poor recovery for DIMP was found with CH-1 soil at ~23% and CL-1 soil at ~38%.

17. Keywords

17.1 liquid chromatography; mass spectrometry; organo-phosphonates; pressurized fluid extraction; soil

TABLE 10 Organophosphates Recovery Data in Georgia Soil (400 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes in Georgia Soil						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	<RL	360.8	331.7	297.4	329.8	360.8	346.9
2	<RL	375.8	331.1	284.4	326.7	363.4	361.2
3	<RL	379.8	342.1	296.0	328.7	366.8	355.9
4	<RL	360.0	335.4	280.1	323.5	372.7	358.3
5	<RL	383.4	337.1	286.8	328.3	356.6	352.1
6	<RL	335.5	302.0	248.2	302.2	332.2	336.4
7	<RL	394.2	359.1	289.1	343.4	383.6	349.6
Average Recovery:	-	369.9	334.1	283.1	326.1	362.3	351.5
Average % Recovery:	-	92.5	83.5	70.8	81.5	90.6	87.9
Standard Deviation:	-	19.5	17.1	16.6	12.3	15.9	8.3
% Relative SD	-	5.3	5.1	5.9	3.8	4.4	2.4

TABLE 11 Organophosphates Recovery Data in Nebraska Soil (400 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes in Nebraska Soil						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	< RL	307.6	218.4	138.4	91.0	240.1	281.3
2	< RL	328.4	213.2	136.0	87.4	240.0	267.8
3	< RL	306.1	216.4	145.2	92.6	242.6	279.4
4	< RL	287.2	208.3	146.3	91.8	232.3	253.8
5	< RL	294.9	218.4	163.1	103.4	260.2	293.9
6	< RL	295.5	206.9	139.6	91.5	225.8	256.4
7	< RL	292.6	212.0	142.1	87.6	235.0	288.3
Average Recovery:	-	301.8	213.4	144.4	92.2	239.4	274.4
Average % Recovery:	-	75.4	53.3	36.1	23.0	59.9	68.6
Standard Deviation:	-	13.8	4.6	9.0	5.3	10.8	15.5
% Relative SD	-	4.6	2.2	6.3	5.8	4.5	5.6

TABLE 12 Organophosphates Recovery Data in ASTM Soil CH-1 (400 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes in ASTM Reference Soil CH-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	<RL	183.2	147.2	160.4	123.6	147.2	111.6
2	<RL	57.2	39.6	51.6	30.8	34.4	27.2
3	<RL	106.4	84.0	94.0	71.2	87.2	49.2
4	<RL	144.4	121.6	130.4	101.6	117.6	71.6
5	<RL	91.2	69.6	78.8	59.2	69.6	45.6
6	<RL	316.4	264.8	277.6	221.2	270.8	228.8
7	<RL	318.0	272.8	285.6	227.6	280.8	218.0
Average Recovery:	-	173.8	142.8	154.1	119.3	143.9	107.4
Average % Recovery:	-	43.5	35.7	38.5	29.8	36.0	26.9
Standard Deviation:	-	105.7	92.8	93.9	77.7	96.8	83.6
% Relative SD	-	60.8	65.0	61.0	65.1	67.3	77.8

TABLE 13 Organophosphates Recovery Data in ASTM Soil CL-1 (400 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes in ASTM Reference Soil CL-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	<RL	114.2	100.8	120.9	98.8	86.4	185.7
2	<RL	70.2	64.6	71.8	55.6	40.8	121.7
3	<RL	64.1	56.6	69.3	52.0	40.5	143.7
4	<RL	113.7	82.1	111.3	78.6	62.2	204.2
5	<RL	86.0	64.0	84.1	61.0	49.5	149.8
6	<RL	133.4	102.0	142.7	95.2	85.3	184.8
7	<RL	83.1	58.5	81.3	53.3	43.3	142.4
Average Recovery:	-	95.0	75.5	97.3	70.7	58.3	161.8
Average % Recovery:	-	23.7	18.9	24.3	17.7	14.6	40.4
Standard Deviation:	-	25.8	19.5	27.9	20.1	20.3	29.9
% Relative SD	-	27.1	25.8	28.7	28.5	34.8	18.5

TABLE 14 Organophosphates Recovery Data in ASTM Soil ML-1 (400 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes in ASTM Reference Soil ML-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	343.0	456.7	391.8	411.9	373.6	403.6	416.8
2	291.8	464.8	388.9	425.9	368.2	409.7	415.1
3	339.1	463.9	399.7	409.4	367.2	413.5	416.1
4	328.7	473.2	399.4	414.1	375.8	413.9	414.4
5	358.6	469.7	401.1	411.2	375.0	419.6	409.1
6	337.8	477.7	413.1	419.8	394.3	428.3	420.6
7	289.2	474.7	416.0	418.8	394.0	433.0	414.5
Average Recovery:	326.9	468.7	401.4	415.9	378.3	417.4	415.2
Average % Recovery:	81.7	117.2	100.4	104.0	94.6	104.3	103.8
Standard Deviation:	26.4	7.3	10.0	5.9	11.3	10.4	3.4
% Relative SD	8.1	1.6	2.5	1.4	3.0	2.5	0.8

TABLE 15 Organophosphates Recovery Data in ASTM Soil SP-1 (400 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 400 ppb Spikes in ASTM Reference Soil SP-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	47.4 [†]	37.7 ^{A,B}	<RL	<RL	<RL	<RL
1	87.7	423.4	391.6	370.2	327.6	375.6	392.0
2	119.4	437.8	420.6	360.6	338.2	376.5	390.4
3	116.1	435.6	404.1	380.8	352.4	392.3	401.7
4	74.7	439.2	389.1	384.0	336.3	389.8	447.7
5	65.9	441.6	384.7	359.0	316.4	390.6	415.2
6	65.6	489.5	411.2	412.0	362.2	411.0	414.2
7	75.0	459.0	416.9	364.0	333.0	385.4	387.2
Average Recovery:	86.3	446.6	402.6	375.8	338.0	388.7	406.9
Average % Recovery:	21.6	111.7	100.7	94.0	84.5	97.2	101.7
Standard Deviation:	22.7	21.6	14.3	18.7	15.3	11.9	21.2
% Relative SD	26.3	4.8	3.6	5.0	4.5	3.1	5.2

^A Blank concentrations are not subtracted from sample results, EMPA not present in Method Blank associated with 1600 ppb spiked samples for soil SP-1.

^B Below Reporting Limit.

TABLE 16 Organophosphates Recovery Data in Ottawa Sand (1600 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 1600 ppb Spikes in Ottawa Sand						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	1136.0	1467.6	1542.8	1540.8	1576.8	1545.6	1504.4
2	1056.8	1537.6	1578.0	1618.4	1636.0	1596.0	1580.0
3	1192.0	1493.2	1524.0	1545.6	1536.0	1541.2	1550.0
4	1015.6	1477.2	1473.6	1564.4	1504.8	1488.8	1510.4
5	1058.0	1471.2	1504.0	1553.6	1505.2	1534.0	1545.2
Average Recovery:	1091.7	1489.4	1524.5	1564.6	1551.8	1541.1	1538.0
Average % Recovery:	68.2	93.1	95.3	97.8	97.0	96.3	96.1
Standard Deviation:	71.0	28.7	39.4	31.4	55.6	38.1	31.0
% Relative SD	6.5	1.9	2.6	2.0	3.6	2.5	2.0

TABLE 17 Organophosphates Recovery Data in Georgia Soil (1600 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 1600 ppb Spikes in Georgia Soil						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	<RL	1524.0	1396.8	1422.0	1307.6	1415.2	1614.0
2	<RL	1574.8	1377.6	1412.4	1316.4	1403.2	1587.2
3	<RL	1596.0	1421.2	1462.8	1326.8	1425.6	1621.2
4	61.6	1550.0	1326.0	1364.8	1252.8	1307.2	1607.2
5	<RL	1484.4	1273.6	1311.2	1232.4	1326.0	1578.0
Average Recovery:	12.3	1545.8	1359.0	1394.6	1287.2	1375.4	1601.5
Average % Recovery:	0.8	96.6	84.9	87.2	80.5	86.0	100.1
Standard Deviation:	-	43.7	59.2	58.2	41.9	54.7	18.3
% Relative SD	-	2.8	4.4	4.2	3.3	4.0	1.1

TABLE 18 Organophosphates Recovery Data in Nebraska Soil (1600 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 1600 ppb Spikes in Nebraska Soil						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	<RL	1979.6	1126.8	401.2	211.6	972.8	1326.0
2	<RL	1966.0	1065.2	410.0	213.6	953.6	1283.6
3	<RL	1925.6	1073.2	452.0	214.0	913.2	1274.4
4	<RL	2004.4	1066.8	444.4	215.6	923.2	1340.4
5	<RL	1964.4	999.2	396.8	181.2	823.6	1236.0
Average Recovery:	-	1968.0	1066.2	420.9	207.2	917.3	1292.1
Average % Recovery:	-	123.0	66.6	26.3	13.0	57.3	80.8
Standard Deviation:	-	28.6	45.3	25.5	14.6	57.5	41.9
% Relative SD	-	1.5	4.2	6.1	7.0	6.3	3.2

TABLE 19 Organophosphates Recovery Data in ASTM Soil CH-1 (1600 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 1600 ppb Spikes in ASTM Reference Soil CH-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	134.4	1600.8	975.6	1412.8	794.4	969.2	744.0
2	80.4	1060.8	588.4	927.6	499.2	613.2	232.8
3	189.6	730.4	390.8	647.6	355.6	424.8	156.4
4	358.4	359.6	180.0	302.0	167.6	190.0	260.4
5	226.4	296.0	138.8	234.4	124.8	140.0	84.0
Average Recovery:	197.8	809.5	454.7	704.9	388.3	467.4	295.5
Average % Recovery:	12.4	50.6	28.4	44.1	24.3	29.2	18.5
Standard Deviation:	105.5	538.9	342.3	484.2	272.2	338.8	260.0
% Relative SD	53.3	66.6	75.3	68.7	70.1	72.5	88.0

TABLE 20 Organophosphates Recovery Data in ASTM Soil CL-1 (1600 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 1600 ppb Spikes in ASTM Reference Soil CL-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	<RL	390.8	380.4	361.2	380.8	326.0	561.2
2	<RL	336.8	314.8	287.6	315.6	256.0	524.4
3	<RL	427.6	398.8	349.6	390.4	316.0	541.2
4	<RL	579.2	549.2	490.8	530.4	458.0	671.6
5	<RL	322.0	320.8	282.0	289.2	213.2	545.6
Average Recovery:	-	411.3	392.8	354.2	381.3	313.8	568.8
Average % Recovery:	-	25.7	24.6	22.1	23.8	19.6	35.6
Standard Deviation:	-	103.0	94.8	84.2	93.7	92.7	58.9
% Relative SD	-	25.0	24.1	23.8	24.6	29.6	10.4

TABLE 21 Organophosphates Recovery Data in ASTM Soil ML-1 (1600 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 1600 ppb Spikes in ASTM Reference Soil ML-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	887.6	1644.0	1458.8	1552.4	1356.4	1395.2	1414.8
2	1022.4	1694.0	1439.6	1550.4	1322.0	1349.2	1598.8
3	845.6	1669.6	1406.8	1537.2	1232.4	1305.2	1503.2
4	1017.2	1566.0	1378.0	1440.4	1202.0	1266.4	1331.2
5	842.4	1615.6	1302.8	1481.2	1173.2	1263.6	1446.4
Average Recovery:	923.0	1637.8	1397.2	1512.3	1257.2	1315.9	1458.9
Average % Recovery:	57.7	102.4	87.3	94.5	78.6	82.2	91.2
Standard Deviation:	90.1	49.6	61.2	49.5	78.7	56.3	99.9
% Relative SD	9.8	3.0	4.4	3.3	6.3	4.3	6.8

TABLE 22 Organophosphates Recovery Data in ASTM Soil SP-1 (1600 ppb Spike)

Precision and Accuracy Samples	Measured ppb from 1600 ppb Spikes in ASTM Reference Soil SP-1						
	MPA (ESI +)	EMPA (ESI +)	EMPA (ESI -)	IMPA (ESI +)	IMPA (ESI -)	PMPA (ESI -)	DIMP (ESI +)
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL
1	1307.6	1604.8	1657.2	1574.0	1574.0	1466.0	1540.4
2	1288.4	1631.2	1702.0	1610.8	1638.8	1439.6	1585.2
3	1222.0	1653.2	1610.0	1584.0	1588.8	1438.0	1490.4
4	1194.8	1576.4	1570.8	1502.4	1525.6	1392.0	1451.2
5	1279.2	1673.6	1658.8	1590.4	1410.0	1466.0	1604.8
Average Recovery:	1258.4	1627.8	1639.8	1572.3	1547.4	1440.3	1534.4
Average % Recovery:	78.7	101.7	102.5	98.3	96.7	90.0	95.9
Standard Deviation:	47.8	38.5	50.5	41.3	86.8	30.2	64.1
% Relative SD	3.8	2.4	3.1	2.6	5.6	2.1	4.2

APPENDIX

(Nonmandatory Information)

TABLE X1.1 Characterization Data for GA and NE Soils

Properties	GA Soil	NE Soil
Sand	46%	6%
Silt	22%	60%
Clay	32%	34%
pH	5.0	5.6
Total Organic Carbon	0.2%	2.1%

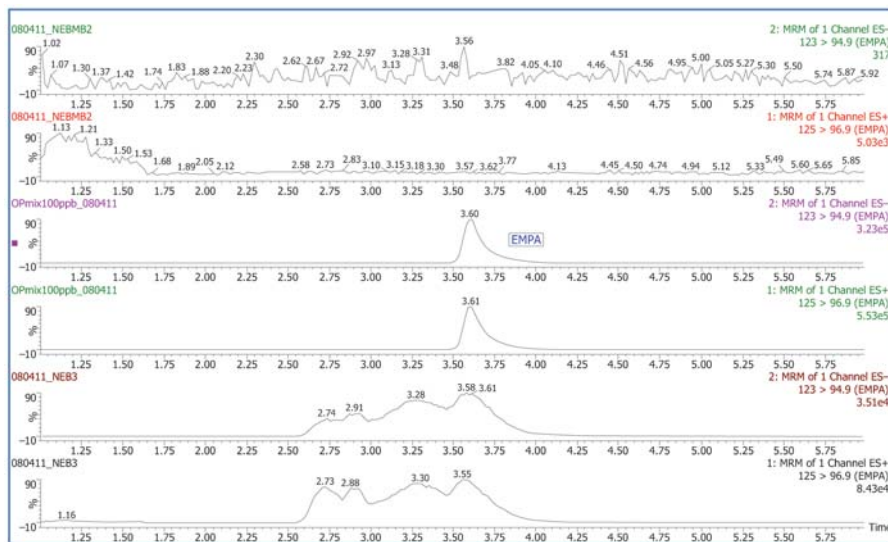


FIG. X1.1 EMPA Chromatograph Comparisons

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