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MORSE LABORATORIES, INC.

SOP# <u>Meth-101</u>

Revision <u>#2</u> Date <u>10/96</u>

DETERMINATION OF HEXAZINONE AND ITS METABOLITES IN SOIL

Reason for revision:

To reduce the initial sample size and amount of eluting solution in Section 4.2. The changes were made to provide a cleaner extract for instrumentation.

1.0 INTRODUCTION

This standard operating procedure (SOP) is intended for the determination of residues of hexazinone and hexazinone metabolites A (IN-T3937), B (IN-A3928), C (IN-T3935), A1 (IN-G3453), 1 (IN-JS472), and G3170 (IN-G3170) in soil.

2.0 EQUIVALENCE STATEMENT

During the conduct of this analysis, equivalent apparatus, solvents, glassware, or techniques (such as sample concentration) may be substituted for those specified in this method, except where otherwise noted. In the event an equivalent piece of equipment or an equivalent technique is used, its use will be documented in the study records, when appropriate.

3.0 MATERIALS

3.1 Apparatus and Equipment

Assorted laboratory glassware including: graduated cylinders, short stem glass funnels, pipers, volumetric flasks, evaporating flasks, microliter syringes.

Gas Chromatograph:

Hewlett-Packard 5890E gas chromatograph equipped with an N/P detector, an HP6890 Autosampler, and an

HP 3365II ChemStation

GC Column:

 $15~\mathrm{M} \times 0.53~\mathrm{mm}$ i.d. fused silica column cross bonded

with 0.5 μ m film thickness Rtx-35 .

Balances:

Analytical balance capable of weighing to ±0.1 mg

Top-loading balance capable of weighing to ±0.01 g

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Centrifuge:

IEC clinical centrifuge (International Equipment Co.,

Needham Hts., MA)

Centrifuge Tubes:

50 mL polyethylene or polypropylene centrifuge tubes

with screw-cap closures

Evaporators:

N-Evap, Models 112 and 115, attached to a nitrogen source (Organomation Associates, South Berlin, MA)

Turbo-Vap, 200 mL unit (Zymark Corp., Hopkinton,

MA)

Evaporator tubes:

Zymark 200 mL tubes (Zymark Corp., Hopkinton, MA)

Filter Paper:

Whatman, #541

Mixer:

Vortex Genie 2 (VWR Scientific, Bridgeport, NJ)

EDP Electronic Pipets

with suitable tips:

Rainin, Ridgefield, N.J.

Reservoirs (75 mL, 25 mL,

or 15 mL) and

adapters for use with

SPE cartridges:

Varian Analytical Instruments, Sunnyvale, CA

Solid Phase

Extraction Apparatus:

Vac Elut Mode SPS 24 (Varian Analytical Instruments,

Sunnyvale, CA), or

Visiprep™ Solid-Phase Extraction Vacuum Manifold

(Supelco, Inc., Bellefonte, PA)

Solid Phase

Extraction Columns:

SupelcleanTM EnviTM-Carb SPE Tubes, Custom, 12 mL Polypropylene Tubes with Teflon Frits packed with 1.5

gram of ENVI-CARB (Supleco, Inc., Bellefonte, PA)

C18 Mega Bond Elut, 6 cc/l gram (Varian Part #1225-

6001) (Varian, Inc., Harbor City, CA)

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Shaker:

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Burrell Wrist-action shaker (Burrell Corp., Pittsburgh,

PA)

Test Tubes:

 16×100 mm, borosilicate, silylated, with Teflon²-lined

screw caps

Ultrasonic Bath:

Branson Model 3200 Ultrasonic Bath (VWR Scientific)

pH meter:

Beckman φ34 pH meter (Beckman Instruments, Inc.,

Fullerton, CA)

3.2 Rengents

Acetone:

Pesticide residue quality

Dimethyldichiorosilane:

Supelco, Catalog No. 3-3009 (Supelco, Inc., Bellefonte,

PA)

Ethyl Acetate:

Pesticide residue quality

Glacial Acetic Acid:

Reagent grade

Hexane:

Pesticide residue quality

Methanol:

Pesticide residue quality

Potassium Phosphate,

Monobasic:

Reagent grade

Sodium Chloride:

Reagent grade

Toluene:

Pesticide residue quality

Water:

HPLC grade

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3.3 Primary Standards

Analytical grade, available from DuPont Agricultural Products Global Technology Division, E. I. duPont de Nemours and Company, Wilmington, DE

Anaivie	Standard Identification
Hexazinone	DPX-A3674
Metabolite A	IN-T3937
Metabolite B	IN-A3928
Metabolite C	IN-T3935
Metabolite Al	IN-G3453
Metabolite 1	IN-JS472
Metabolite G3170	IN-G3170

3.4 Reagent Preparation

3.4.1 3mM Acetic Acid

Add 173 μ L glacial acetic acid per one liter of HPLC grade water. Record the pH of this solution. This solution is stable for approximately two weeks at room temperature. Discard the solution when it starts showing signs of turbidity.

3.4.2 <u>0.1M KH,PO./0.5M NaCl Solution</u>

Weigh 13.6 g KH₂PO₄ and 29.2 g NaCl into a 1 liter volumetric flask. Dissolve the compounds with HPLC grade water. Bring to volume with HPLC grade water and mix well. Record the pH of this solution (~pH 4). This solution is stable for one month at room temperature. Discard the solution when it starts showing signs of turbidity.

3.4.3 Extraction Solution

Combine equal volumes of acetone and 0.1M KH₂PO₄/0.5M NaCl Solution. Mix well. Prepare fresh daily.

3.4.4 9:1 Acetone:3mM Acetic Acid

Mix 90 mL acetone with 10 mL 3mM acetic acid solution. Prepare fresh daily.

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3.4.5 9:1 Methanol:3mM Acetic Acid

Mix 90 mL methanol with 10 mL 3mM acetic acid solutions. Prepare fresh daily.

3.4.6 <u>Mixed solvent solution (20% acetone/50% ethyl acetate/30% toluene GC Dijuting Solution)</u>

In a 50 mL mixing cylinder add 10 mL acetone and 25 mL ethyl acetate. Dilute to volume with toluene. Mix well. Prepare fresh as needed.

3.5 Analytical Standard Preparation

3.5.1 Stock Standard Solutions

12.5 mg (corrected for purity) of each analytical standard is accurately weighed, quantitatively transferred to a 25 mL volumetric flask, and brought to volume with methanol to make individual stock standard solutions having a concentration of 500 μ g/mL. These stock standard solutions are to be stored at 1-8 °C when not in use. These solutions are stable for approximately six months when stored under these conditions.

3.5.2 Fortification (Spiking) Solutions

Typically, mixed standard solutions containing all seven analytes per standard solution are prepared by adding the appropriate amounts of standard solutions into a volumetric flask and diluting to volume with methanol. Store all spiking solutions in a refrigerator when not in use.

For preparation of various standard concentrations see form ML 471a in Appendix I.

3.5.3 Intermediate Standard Solutions

Typically, intermediate standard solutions containing all analytes are used to prepare the GC standards. The intermediate standard solutions and the GC standards are prepared in volumetric flasks and diluted with mixed solvent solution (20% acetone/50% ethyl acetate/30% toluene). Store these standard solutions in a refrigerator when not in use.

For preparation of various standard concentrations see form ML 472a in Appendix I.

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3.5.4 GC (Calibration) Standard Solutions

Typically, GC standards containing all seven analytes are prepared. Store GC standards in a refrigerator when not in use.

For preparation of various standard concentrations see form ML 472a in Appendix I.

4.0 ANALYTICAL PROCEDURE

4.1 Principle

The analytes are extracted from the soil sample with an aqueous buffer-acetone solution. The resulting extract is evaporated to the aqueous phase and passed through a disposable graphetized carbon SPE column. The analytes are eluted from the carbon column with acid-acetone solution. The cluate is evaporated to dryness and brought to the water phase for a second column cleanup using a C18 SPE column. The analytes are cluted from the column with acid-methanol. The cluate is evaporated to dryness and the residue dissolved in a suitable mixed solvent solution for gas chromatographic analysis using NP detection.

4.2 Sample Extraction

Note: Use disposable laboratory equipment whenever possible. All glassware used for the analysis must be meticulously cleaned prior to use.

- 1. Weigh a 5.0 g sample into a 50 mL centrifuge tube. Fortify at this point.

 Allow the spiking solvent to evaporate under a fume hood.
- 2. Add 20 mL extracting solution. Manually shake ten times. Shake for an additional 10 minutes using a wrist-action shaker at maximum speed.
- Centrifuge for 10 minutes at approximately 2500 rpm.
- 4. Decant the resulting extract through a filter into a silanized Zymark tube.
- 5. Repeat steps $4.2.2 \rightarrow 4.2.4$ one more time decanting the extract through the same filter into the same collection vessel. Rinse the filter with 2×10 mL acetone.

This is a stopping point. Refrigerate extracts for further processing.

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- 6. Evaporate the combined extract to the water phase (~15 mL) at 50 °C using the Turbo-Vap evaporator. Use a template to determine this endpoint.
- 7. Dilute the extract to ~50 mL with HPLC grade water. Mix well.

This is a stopping point. Refrigerate the extract for further processing.

Just prior to carbon SPE column cleanup, sonicate the extract for 5 minutes.
 Mix well.

4.3 Solid Phase Extraction (SPE)Purification

- 1. Pack the carbon column by gently pushing the column frit onto the carbon packing. Condition the column with one column volume of 9:1 acetone: 3mM acetic acid. Elute to dry and dry the column for approximately 30 seconds more with vacuum, adjusting the vacuum gauge to minimize loss of small amounts of carbon. Continue conditioning the column with two column volumes of water without allowing the column to go to dry. Discard all elutions.
- 2. Attach a reservoir to the cartridge and pass the sample extract through the cartridge. Use vacuum to facilitate sample elution. Do not allow the cartridge to go to dry. Discard this sample load. Remove the reservoir.
- 3. Rinse the sample container with one column volume (10-11 mL) of water and add rinsing to the column. Elute to dryness and dry for approximately 30 seconds more with vacuum. Adjust the vacuum gauge to minimize the loss of small amounts of carbon. Discard this wash.
- 4. Rinse the column with one column volume of hexane. Allow the hexane wash to go to dryness and dry for ~30 seconds more with vacuum. Adjust the vacuum gauge to minimize the loss of small amounts of carbon. Discard the hexane wash.
- 5. Elute the analytes into a 16 × 100 mm silanized glass test tube (marked at ~1 mL and ~5 mL) with 10 mL of 9:1 acetone:3mM acetic acid. Elute by gravity. However, vacuum may be used to start the elution.

Note: See Appendix II for detailed instructions for silylating glassware.

This is a stopping point. Refrigerate extracts for further processing.

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- 6. Evaporate the sample extract to ~1 mL at 40 50 °C using the N-Evap.
- 7. Wash the sides of the sample test tube with ~2 mL accrone and evaporate the extract to 0.2 to 0.5 mL.
- 8. Repeat washing the sides of the test tube with a second 2 mL portion of acetone.
- Take the extract to dryness.
- 10. Dissolve the residue in 200 μ L acetone. Briefly vortex the extract and sonicate for 5 minutes. Again, vortex briefly.
- 11. Add ~1 mL of water and evaporate to the water phase.
- 12. Add -4 mL HPLC water and continue the evaporation step to ensure removal of all traces of the acetone.
- 13. Adjust to ~5 mL with HPLC water.

This is a stopping point. Refrigerate the extract for further processing..

 Vortex. Sonicate for 5 minutes. Vortex. The extract is now ready for additional SPE column cleanup.

This is a stopping point. Refrigerate the extract for further processing.

4.4 Additional Column Cleanup with C18 SPE Cartridge:

Note: To ensure uniformity of the column packing in the cartridge, tap the cartridge on the lab bench, then gently push the column frit onto the column packing prior to use.

- 1. Pass 5 mL 9:1 (Methanol:3mM Acetic Acid) to the column. Elute to dryness and dry for 30 seconds more with vacuum. Continue conditioning with 2 × 5 mL HPLC water without allowing the column to go to dryness.
- 2. Load the aqueous extract from 4.3.14 onto the column and elute to ~1 cm above the column packing.

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- Rinse the sample test tube with ~2 mL HPLC water and add this rinsing to the column. Eiute just to dryness. Discard the sample load and wash.
- 4. Wash the column with 2 × 5 mL hexane. Elute the second 5 mL hexane wash to dryness. Continue drying the column for 30 seconds more with vacuum. Discard the hexane washes.
- 5. Elute the analytes into a 16 × 100 mm silanized glass test tube calibrated at 2.0 mL with 7 mL 9:1 Methanol:3mM Acetic Acid. A suitable reservoir may be used for this step to facilitate elution. Elute by gravity. However, vacuum may be used to start the elution.

This is a stopping point. Refrigerate the extract for further processing.

4.5 Concentration and Processing of Sample Eluates for GC Analysis:

- Evaporate the sample extract to ~1 mL at approximately 50 °C using the N-Evap.
- 2. Wash the sides of the sample test tube with ~1 mL methanol and evaporate the extract to 0.2 to 0.5 mL.
- 3. Repeat washing the sides of the test tube with a second 1 mL portion of methanol.
- 4. Take the extract to dryness. Determine the final volume of the extract at this point so that the concentration of the sample in the final extract is 2.5 g/mL.
- 5. Dissolve the residue in a volume of acetone equivalent to 20% of the final volume of the extract. Briefly vortex the extract and sonicate for 5 minutes. Again, vortex briefly.
- 6. Add a volume of ethyl acetate equivalent to 50% of the final volume of the extract and ensure complete solution of the analytes by vigorously vortexing the mixture.
- 7. Adjust the final volume of the sample extract with toluene. Sonicate for 5 minutes. Mix thoroughly for GC analysis. The final concentration of the sample in the extract is 1 mL = 2.5 g.

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4.6 GC Analysis

Note: The column and conditions stated in the method have been satisfactory for the matrix being analyzed. The specific column packing/coating, carrier gas, column temperature and flow rate listed are typical conditions for this analysis. Specific conditions used will be noted on each chromatographic run and will not otherwise be documented.

4.6.1 Operating Conditions

Instrument: HP5890E gas chromatograph equipped with an N/P detector,

electronic pressure controlled inlet (packed/purged), an HP6890

autosampler, and an HP 3365 II ChemStation.

Column:

15 M × 0.53 mm i.d. fused silica column crossbonded with

0.5 μm film thickness Rtx-35

Inlet Liner:

2 mm i.d. liner lightly packed with fused silica wool

Injection Volume:

 $2 \mu L$

Carrier Gas:

Heljum

Flow:

Column: Makeup: 5 mL/min. (constant flow)

15 mL/min.

Temperature:

Injector.

290 °C

Detector:

285 °C

Column:

Initial: 150 °C

Rate I:

25 °C/min. to 275 °C hold 2.2

minutes,

Rate 2:

10 °C/min.

Final:

285 °C hold for 1.00 minute .

4.6.2 Sample Analysis

Prepare a four-point standard curve by injecting constant volumes of combined analyte standard solutions prepared in a mixed solvent solution. Use constant volume injections for sample extracts as well. Sample responses not bracketed by the standard curve require dilution and reinjection.

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4.6.3 Calculations

Calculations for instrumental analysis are conducted using a validated software application to create a standard curve based on linear regression. These regression functions are used to calculate a best fit line (from a set of standard concentrations in μ g/mL versus peak area response) and to determine concentrations of the analyte found during sample analysis from the calculated best fit line.

The equation used for the least squares fit is:

$$y = mx + b$$

where,

y = peak area response

 $x = \mu g/mL$ found for peak of interest

m = slope

b = y-intercept

The calculations for ppb analyte found and percent recovery (for fortified samples) for each of the seven analytes are:

1. ppb analyte =

where:

 $\mu g/mL$ analyte is calculated by linear regression based on peak area response.

1000 = conversion factor from μ g to ng

g sample = amount of sample analyzed

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mL solvent = volume of extraction solvent

mL aliquot = volume of sample extract taken through the procedure

mL final volume = volume of final extract submitted to GC

GC dil. fact. = the magnitude of dilution required to

the magnitude of dilution required to bracket the response of the sample within the standard curve responses. When the sample requires no dilution, the GC dilution factor = 1

2. The percent recovery for fortified control samples is calculated as follows:

% Recovery = $\frac{ppb}{ppb}$ found in fortified control - ppb found in control \times 100

5.0 REFERENCES

- 1. Dupont Report No. AMR 3888-95 (Draft), "Analysis of Hexazinone and Metabolites in Soil and Groundwater using GC/MS," received 11/22/95.
- 2. Dupont Report No. AMR 2896-94
- 3. Morse Laboratories, Inc. SOP# Meth-93, Revision #4, "Determination of Hexazinone and its Metabolites in Water"

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Silylation of Glassware

Purpose:

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Silylation is a process used to chemically treat glassware in order to prevent or minimize binding of analyte residues to the glass surface.

Caution:

DO NOT ALLOW DIMETHYLDICHLOROSILANE TO COME IN CONTACT WITH WATER. CHLORINE GAS AND HYDROGEN CHLORIDE GAS WILL BE PRODUCED.

THIS PROCEDURE MUST BE DONE UNDER A FUME HOOD. THE TECHNICIAN MUST WEAR HEAVY LATEX GLOVES.

Procedure:

1. Prepare 100 mL of a 5% (v/v) solution of dimethyldichlorosilane (DMDCS) in hexane.

To a glass stoppered glass container (approximately 200 mL volume) add 95 mL hexane. Slowly add 5 mL DMDCS. Stopper and invert to mix.

Larger volumes can be prepared using the proportions discussed above, however attempt to prepare amounts that will be nearly totally used to avoid disposal of excess solution.

- 2. Pour a small amount of the DMDCS solution into the glassware to be treated. Rotate the glassware to thoroughly coat the inside surfaces. Pour excess solution into the next piece of glassware to be treated.
- 3. Allow the treated glassware to dry (approximately 20 minutes). Rinse with deionized water, then acctone. Again allow to dry.

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- 4. Glassware is now ready for use.
 - Note: Any glassware that is cleaned with a brush after it has been silylanized, must be resilylanized.
 - Store pure DMDCS at room temperature.
 - 5% solutions of DMDCS in hexane are stable for 5 days when stored well-stoppered at room temperature.

SOP Prepared by: <u>Frances Brookev</u>

<u>Kevin Clark</u>



DuPont Report No. 2292

ANALYTICAL METHOD FOR THE DETERMINATION OF HEXAZINONE AND METABOLITES OF INTEREST IN SOIL AND WATER USING ELECTROSPRAY-LC/MS/MS

F. W. Brill, Tom Gardner

1.0 SUMMARY

A residue method is described for the extraction, purification, and quantitation of hexazinone (DPX-A3674) and metabolites A (IN-T3937), A-1 (IN-A3453), B (IN-A3928), C (IN-T3935), I (IN-IS472), and G3170 (IN-G3170) in soil and water...

Water samples (20 mL) were filtered, concentrated, and purified using solid phase extraction cartridges (Envi-Carb and C13). The instrumental analysis for detection and quantitation of the analytes was accomplished by LCMS/MS with an electrospray interface in multiple reaction monitoring (MRM).

Soil samples (5g) were extracted with 0.1M KH.PO./0.5M NaCl solution, filtered, concentrated, and purified using solid phase extraction cartridges (Envi-Carb and C18). The instrumental analysis for detection and quantitation of the analytes was accomplished by LC/MS/MS with an electrospray interface in multiple reaction monitoring (MRM).

The limit of quantitation (LOQ) for water was 0.100 ppb for hexazimone and all 6 metabolites. The LOQ for soil was 2.0 ppb for hexazimone and all 6 metabolites. Test samples were fortified at the LOQ and 5 \times LOQ for each analyte in both soil and water to validate the method

To be completed

2.0 INTRODUCTION

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Hexazinone (DPX-A3674) is the active ingredient in Velpar® herbicides registered for postemergence control of many annual and biennial weeds. Analyte structures, chemical names, DuPont code numbers, and Chemical Abstracts registry numbers are provided in Figure 1.

In this method, water samples are acidified, filtered, purified and concentrated by SPE. A combination of an Envi-Carb and a C18 SPE step are used to remove most matrix and substances that may interfere with the insurmental analysis...

The purified extracts were analyzed by ESI-LC/MS/MS using positive mode multiple reaction monitoring (MRM) for hexazinone and metabolites A, A-1, B, C, 1, and G3170. Quantitation was based on the integration of a single MRM transition response. This analysis quantitatively detects hexazinone and all 6 metabolites, the LOQ for each determined to be 0.1 ppb in water and 2.0 ppb in soil.

To be completed

3.0 MATERIALS

Equivalent equipment and materials may be substituted unless otherwise specified (see section 5.3); note any specifications in the following descriptions before making substitutions. The equivalency/suitability of any substitution should be verified with acceptable control and fortification recovery data. The materials are listed in order of first appearance in the method.

3.1 Equipment

Standards Preparation:

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- Mettler AE 163 analytical balance (Mettler instrument Corp., Hightstown, N.J.)
- · Pipers/pipertors and tips suitable for preparation of standards and sample fortifications
- Assorted flasks, beakers and volumetric cylinders

Samele Extraction:

- Mettler PM460 top-loading analytical balance (Mettler Instrument Corp.)
- · 250-mL high density polypropylene centringe tubes (Nalgene)
- RapidVap[®] Evaporation System (Labconco Corp., Kansas City, Mo.)
- Tissumizer homogenizer Model FTD-1810 (Tekmar Company, Cincinnati, Ohio)
- 0.45-µm filter tmit, P.N. 150-0045 (Nalge Company, Rochester, NY)
- Sorvall® RC5C centrifuge with SS34 rotor (DuPont Company Sorvall® Products, Wilmington, Del.) or IEC HN-SII centrifuge (Damon/IEC Division, Needham, Mass.).
- Buchner Funnel
- Filter Paper, Whatman #5 (Cat. No. 1005 070) to fit Buchner Funnel

Sample Purification:

- C18 Mega Bond Elur[®], 20cc/lg, P.N. 1225-6001 (Varian, Harbor City, CA.)
- Supelclean⁷⁴ Envi-Caro SPE column, 60mL/1.5g, CUSTOM (Supelco Inc., Bellefonte, Pa.)
- 15-ml. cap. reservoir, empty, P.N. 1213-1010 (Varian); Luer stopcock, P.N. 1213-1005 (Varian); Bond Elut[®] adapters, P.N. 1213-1005 (Varian)
- N-Evap Analytical Evaporator, Model 111 (Becton Dickinson & Co., Franklin Lakes, N.J.)
- 2-µm, 13-mm AcroDisc 13CR PTFE syringe filter, P.N. 4422 (Gelman Sciences, Ann Arbor, Mich.)
- 4.5-um, 25-mm AcroDisc 13CR PTFE syringe filter, P.N: 4419 (Gelman Sciences, Ann Arbor, Mich.)
- Vacuum manifold, P.N. 5-7030 (Supelco Inc., Bellefonte, Pa.)
- 3200R-4 Bransonic Ultrasonic Cleaner (Bransonic Ultrasonics Corp., Danbury, Conn.).

LC/MS Analysis:

The following HPLC/MS system was used for this method including equipment necessary to divert the flow going into the mass spectrometer to waste and of post-column splitting of the HPLC flow.

- Micro-Mass Quartro II triple sector quadrupole instrument with API source/interface configured for ESI operation.
- MassLynx data acquisition software running under Windows NT.
- Waters Model 1100 HPLC system including pump module, autosampler, degassing module, column compartment module, (Waters Corp., Milford, Mass.)
- Electrically-actuated, high-pressure 6-port switching valve with 1/16 in. fittings, #EC6W, used for effluent diversion from MS (4-port valve would be adequate, #EC4W) (Valco Instrument Ca., Inc., Houston, TX)
- High-pressure stainless steel 1/16 in fitting tee, #ZT1, for use in post-column flow splitting. Split ratio adjusted by altering restriction on waste-side of tee by altering length of capillary tubing. (Valco Instrument Co., Inc.)
- Zorbax[®] 4.6 mm id × 250 mm Rx-C3, 5-µm particle size, P.N. 380967-901 (MAC-MOD Analytical, Inc., Chadds Ford, Pa.) Substitute columns may give sub-optimal performance.

3.2 Reagents and Standards

Reagents:

OmniSolv³ glass distilled acctone, hexane, methanol

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- Baker Analyzed Reagent porassium phosphate, dibasic, crystal, P.N. 3252-01, KH₂PO₄ (J.T. Baker, Phillipsburg, N.J.)
- Baker Analyzed Reagent' sodium chloride, crystal, P.N. 3252-01, KH,PO₄ (J.T. Baker, Phillipsburg, N.J.)
- GR glacial acetic acid (EM Science)
- All water was Milli-Q[®] distilled, deionized water (Millipore Corp., Bedford, Mass.).

 <u>Standards</u>: All standards were synthesized by DuPont Agricultural Products, Wilmington, Del. Standards should be greater than 95% purity
 - DPX-A3674-213 (hexazinone)
 - IN-T3937-3 (membolite A)
 - IN-G3453-2 (membolite A-1)
 - IN-A3923-4 (membolite B)
 - IN-T3935-3 (metabolite C).
 - IN JS472-2 (metabolite 1)
 - IN G3170-2 (metabolite G3170)

3.3 Safety and Health

No unusually hazardous materials are used in this method. All appropriate material safety data sheets should be read and followed, and proper personal protective equipment should be used.

4.0 METHODS

4.1 Principles of Analytical Method

4.1.1 Sample Extraction

The 0.1M KH_PO/0.5M NaCl soil extraction solution yielded high recoveries and has been demonstrated as an effective extraction agent for hexazinone and its metabolites. Water is not extracted, it is directly filtered, concentrated, and purified on the SPE columns.

4.1.2 Extract Purification

Carbon phase (Envi-Carb) and Reverse phase (C18), Bond Elut[®] SPE procedures are used to provide extensive extract purification. The water and the soil extracts are passed onto and retained on Envi-Carb columns. The column is then washed with water and hexane to remove most of the matrix material. Elution is with an acetonet3 mM acetic acid (90:10 v:v) solution. The acetone in the elutae is removed and the sample is brought up in water and loaded onto a C18 column. The C18 is then washed with water and hexane, following which it is eluted with methanol. The sample is blown down to dryness through a series of procedures designed to minimize loss of analyte on the container walls and is brought up to a final volume in water, for a final filtration and LCMS analysis.

4.1.3 LC/MS Analysis

The method utilizes electrospray ionization (ESI) and is operated in positive ion mode using Multiple Reaction Monitoring, (MRM) on the Micromass instrumentation for all of the analytes. A single transition was monitored for each analyte: [MH+1] fragmenting to the [M-cyclobexane moiety]. Selection of these transitions was based upon the mass spectra generated during the method development process with the instrument in scanning mode. The spectra generated by ESI-LCMS for the analytes are shown in Figure 2. The base ion. [MH]*, for each analyte is selected for quantitation. The sensitivities of each analyte varied, the G3170 showing the least sensitivity in the positive ESI SRM mode, the hexazimone showing the most.

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DuPont Report No. 2292

1.2 Analytical Procedure

12.1 Glassware and Equipment Cleaning Procedures

Disposable labware are used whenever possible in this method. Reusable labware, which includes the evaporation vessels and volumetries for standard solutions, are cleaned by washing with a laboratory grade detergent followed by tap water rinses (3) and distilled water rinses (3). A final actione rinse may be used to remove the residual water and promote drying. Certain items of glassware must be treated silanized to prevent absorption of analyses onto the glasss.

4.2.2 Preparation and Stability of Reagent Solutions

0.1M KH.PO 10.5M NaCl Solution: Add 13.6g KH.PO, and 29.2g NaCl into a 1 liter volumetric flask. Dissolve the compounds with HPLC or Milli-Q water. Bring to volume and mix well. Record the pH of the solution (should be -4). This solution is stable for 1 month, discard earlier if it show signs of turbidity or bacteria growth. Scale volume as necessary.

3 mM Acetic Acid: Add 173 µL of glacial acetic acid per 1 L of HPLC or Milli-Q water. The solution is stable for 1 month. Discard earlier if it shows signs of turbidity or bacteria growth.

1/10 diluted Glacial Acetic Acid. Prepare a 1:10 dilution of glacial acetic acid by placing 9 mL of HPLC or Milli-Q water into a 15 mL plastic centrifuge tube or any other suitable container with a cap. Add 1 mL of glacial acetic acid, mix carefully. Cap when not in use.

Extraction solution for Soil; is prepared by combining accome and the 0.1M KH,PO,0.5M NaCl Solution in a 90:10 v/v ratio. This solution is prepared daily as needed, however will be stable for I month.

90:10 Acetone:3mM Acetic Acid Solution: Mix 90 mL of acetone with 10 mL of 3 mM Acetic Acid solution. This solution is used for SPE column conditioning and elution. Scale volume as needed, stable for 1 month.

90:10 Methanol:3mM Acetic Acid Solution: Mix 90 mL of methanol with 10 mL of 3 mM Acetic Acid solution. This solution is used for SPE column conditioning and elution. Scale volume as needed, stable for 1 month.

HPLC Aqueous Mobile Phase: A 0.01M acetic acid solution for use as mobile phase is prepared by filling a 1 L volumentic flask with distilled, desonized water, adding 600 µL of concentrated acetic acid, gently agitating the mixture, and diluting to volume in distilled, desonized water. This solution should be prepared as needed or sooner if turbidity or bacterial growth is observed.

4.2.3 Stock Solutions Preparation

If possible, standards with a purity greater than 95% are to be used. Individual 100±2-µg/mL, stock standards solutions for hexazinone and the 6 listed metabolites were prepared by weighing 5±0.1 mg of each standard in a separate, taxed 50-mL volumetric flask and diluting to volume in acctone. Sample weights were determined to 3 significant figures. The analytical balance must provide a weight precision to 3 significant figures, the amount and volume has to be adjusted to meet this criteria. For example, increase the amount to 10.0±0.2 mg and use a 100-mL volumeric flask. Clearly label each stock solution with date prepared, analyte, and concentration. Stock solutions are stored under refrigeration (4±2 °C) and must be replaced at least every 120 days or when approximately half-volume. Keep stock solutions stoppered to restrict solvent evaporation.

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4.2.4 Intermediate Standard and Fortification Solution Preparation

Water/Soil Intermediate Standard: An Intermediate 1.0 µg/mL mixed standard solution is made from the 7 individual 100 µg/mL stock solutions. Place 0.5 mL of each of the 7 individual 100 µg/mL stock solutions into a 50 mL volumetric flask using a piper. Place the volumetric onto the N-Evap and pass nitrogen through the flask to blow off all of the accetone, remove immediately. Add HPLC or Milli-Q water and fill to the 50 mL line. Vortex, sonicate for 5 min.

Water Fortification and Standard Solution: A 50 ng/ml. water fortification and standard solution is prepared from the mixed 1.0-µg/ml. Intermediate Standard. Remove the Intermediate Standard from the refrigerator and aillow it to reach room temperature. Then shake by hand to insure standard consistency before making dilutions. Place 2.5 ml. of the 1.0 µg/ml. intermediate stock solution into a 50 ml. volumentic flask. Fill up to the line with water and mix. Label the solution with the date prepared, analytes, and concentration. Store the solution in the refrigerator. It is stable for 90 days.

Soil Fortification and Standard Solution: A 200 ng/mL water fortification and standard solution is prepared from the mixed 1.0-µg/mL Intermediate Standard. Remove the Intermediate Standard from the refrigerator and allow it to reach room temperature. Then shake by hand to insure standard consistency before making dilutions. Place 10.0 mL of the 1.0 µg/mL intermediate stock solution into a 50 mL volumetric flask. Fill up to the line with water and mix. Label the solution with the date prepared, analyses, and concentration. Store the solution in the refrigerator. It is stable for 90 days.

4.2.5 LC Calibration Standard Solution Set Preparation

Water: The calibration standard set is made from the 50 ng/mL Water Fortification and Standard solution. Remove the Water Fortification and Standard solution from the regigerator and allow it to reach room temperature. Then shake by hand to insure standard consistency before making dilutions. Refer to the table below. Place the specified aliquot of the 50.0 µg/mL Water Fortification and Standard solution into a 10 mL volumetric flask. Fill to the line with HPLC or Milli-Q* water. Label as calibration standard solutions with date prepared, analytes, and concentration. Store these solutions in the refrigerator. They are stable for 90days.

<u>Identifier</u>	Aliquot of 50 ne/mL Std.(uL)	<u>Final Vol.</u>	Standard Concentration (ng/mL)
WSI	2000	10 m L	10
WS2	1000	10 mL	5.0
WS3	400	10 mL	2.0
WS4	134	10 mL	0.67

Clearly label the calibration solutions with the date prepared, analytes, and concentration. Autosampler vials should be filled to approximately 2/3 capacity and capped to minimize solvent evaporation. The 10 mL volumetric flasks of standards will be stable for 2 weeks. Use fresh aliquots from them for each analysis set.

Soil: The calibration standard set is made from the 200 ng/mL Soil Fortification and Standard solution. Remove the Soil Fortification and Standard solution from the refrigerator and allow it to reach room temperature. Then shake by hand to insure standard consistency before making dilutions. Refer to the table below. Place the specified aliquot of the 200.0 µg/mL Soil Fortification and Standard solution into a 10 mL volumente flask. Fill to the line with HPLC or Milli-Q⁹ water. Label as calibration standard solutions with date prepared, analytes, and concentration. Store these solutions in the refrigerator. They are stable for 90days.

Identifier	Aliquot of 200 ng/mL Std.(uL)	Final Vol.	Standard Concentration (ne/mL)
SS1	2500	10 mL	50.0

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SSZ	1250	<u>1m</u> 01	25.0
SS3	500	10 mL	10.0
SS4	167	10 m L	3.33

Clearly label the calibration solutions with the date prepared, analytes, and concentration. Autosampler vials should be filled to approximately 2/3 capacity and capped to minimize solvent evaporation. The 10 ml. volumetric flasks of standards will be stable for 2 weeks. Use fresh aliquots from them for each analysis set.

4.2.5 Source and Characterization of Samples

4.2.7 Storage and Preprocessing of Samples

4.2.3 Sample Fortification Procedure

Water: All water test samples are fortified with hexazimone and the 6 listed metabolites from the 50 ng/mL Water Fortification and Standard solutions. The appropriate fortification volume is drawn into a piper/pipertor and placed into 20 mL of water. The water is then mixed. Refer to the table below for the volumes of Water Fortification and Standard solution to use for making the LCQ and 5 x LOQ fortifications.

Soil; All soil test samples are fortified with hexazinone and the 6 listed metabolites from the 200 ng/ml. Soil Fortification and Standard solutions. The appropriate fortification volume is drawn into a piper/pipettor and placed into 5g of soil. The soil is allowed to stand for 10 minutes. Refer to the table below for the volumes of Soil Fortification and Standard solution to use for making the LOQ and 5 x LOQ fortifications.

<u>Matrix</u>	Amount to	Volume Fortification Soln.	Volume Fortification Soln.	Levels: all analytes LOQ
	<u>Fortify)</u>	Low Fortification (LOO)	High Fornification (LOQ)	and 5 x LOO
Water	20 mL	40 μL	200 μL	0.1 ppb,, 0.5 ppb
Soil	5 g	50	250	2.0 ppb,. 10 ppb

4.2.2a Concentration and Purification Procedures for Water

Solution Requirements/Sample (extraction + purificate	ion):	
Milli-Q water	50	mL
methanol	10	$\mathbf{m}\mathbf{L}$
hexane	10	mL
90:10 acetone: 3 mM Acetic Acid	10	mL
90:10 methanol:3mM Acetic Acid	10	шL

SAMPLE CONCENTRATION AND PURIFICATION

- 1. Remove samples from refrigerator or freezer and allow to warm to room temperature.
- 2. Measure 20 mL of the water sample and place it into a 250 mL centrifuge bottle.
- Fortify test samples if required.
- Add 30 mL of Milli-Q water and then acidify by adding 200 μL of the 1/10 dijuted glacial acetic acid solution. Cap and briefly vottex mix.

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ENVI-CARB BOND ELUT® CLEANUP

- Place the 1.5 g Envi-Caro Bond Eluts³ into the vacuum manifold and condition them by placing 10 mL of 90:10 acctone:3 mM acetic acid solution into the tubes and adjusting the flow to a fast drip (-10 mL/minute). Using a gentle vacuum to pull through all of the liquid, let the column dry for 30 seconds after the last of the liquid has exited. Load and condition the column with two 10 mL volumes of water. Do not let the liquid level drop below the top of the sorbent after the second water load has passed through. Do not allow the nibe to dry.
- Load the water sample onto the column, in several portions if necessary due to column
 reservoir capacity. Turn on the vacuum and allow the sample to pass through the
 column at a slow drip. Do not allow the column to go dry. Do not collect the liquid.
- Place 5 mL of water into the empty 50 mL centrifuge tube, vortex. Place this rinse onto
 the Bond Elut and allow to pass through at a slow drip. Pull all the water through, allow
 to dry, passing air through for about 30 seconds after last drop has come off.
- Place 5 mL of hexane into the Bond Elist and allow to pass through at a slow drip.
 Allow to dry, passing air through for about 30 seconds after last drop has come off.
- Open the vacuum manifold and place plastic 15 mL centrifuge tubes into the manifold in order to collect the cluste.
- 6. Place 10 mL of 90:10 acctone:3 mM accide acid solution into the Bond Elut tube and allow to pass through at a slow drip. Pull all of the liquid through into the collection tubes. Break the vacuum and remove the tubes containing the cluate.
- 7. Place sample on an N-Evap and evaporate to approximately 1 mL at 40-50 deg.
- Wash the sides of the sample test tube with approximately 2 mal of acctone and
 evaporate the extract to 0.2 to 0.5 mL. Repeat this washing with a second 2 mL volume
 of acctone. Evaporate the sample to dryness
- 9. Add 200 µL, of accione to the tube, vortex and sonicate for 5 minutes, vortex again. Add approximately 1 mL of water and evaporate to the water phase. Add approximately 4 mL of HPLC water and continue the evaporation step to ensure removal of all traces of accione This is a very critical step, the presence of any accione will prevent the polar analytes from being retained on the C18 column in later steps.)
- 10. Adjust to approximately 5 mL with HPLC water. Vortex, sonicate for 5 minutes, vortex
- Proceed directly to the C18 SPE clean-up. This is a stopping point. Samples can be placed into the refrigerator for storage for up to 1 week.

C18 BOND ELUT CLEANUP/FINAL PREPARATION

- Place the C13 Bond Eiut[®] tubes onto the vacuum manifold and condition them by placing 5 mL of methanol:3mM Acetic Acid (90:10 v/v) solution into each tube and adjusting the flow to a fast drip. Drain the tube and allow to dry for 30 seconds more under vacuum. In the same manner condition next with 10 mL of water, in two 5 mL increments, waiting until the first 5 mL is completely below the first before adding the second 5 mL. Stop the flow when the level of the water drops slightly below the top of the sorbent in the bottom tube. Do not let the tubes dry.
- Place the Envi-Carb charte solution (step 11 ENVI-CARB cleanup above) into the C13 Bond Elur[®] nibe. Apply vacuum, open the stopcock and load the cluate onto the column with a very slow drip. Stop when the level of the liquid is 1 cm above the top of the sorbent. Discard the aqueous effluent.
- 3. Backrinse the original Envi-Carb cluate container by placing 2.0 mL of water into it vortex mixing for a few seconds. Pour this rinsate into the C13 cartridge. Allow this (rinse)water wash to pass through the C13 and discard. Allow all of the water to be drawn out of the tuce. Immediately shut off vacuum, discard load volume

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- 4. Place 5.0 mL of hexane into the C18 carridge. Allow the hexane to pass through at a moderate drip. After the hexane has completely passed through the column, add another 5.0 mL of hexane and allow that to completely pass through the column as well. Allow the vacuum to pull air through the carridge until there is no more dripping.
- Break vacuum and place a new 15 mL plastic centrifuge tube into the vacuum manifold under the C18 mises to collect the cluate.
- Place 10 mL of methanol:3 mM Acetic Acid (90:10 v/v) solution into the column.
 Allow the cluate to pass through at a moderate drip and collect. Discard the C18 column.
- When elution is finished, place the tube containing the extract onto the N-Evap and blow it down to approximately 1 mL
- Wash the sides of the tube with approximately 1 mL methanol and blow down to
 approximately 0.2 t 0.5 mL. Repeat with another 1 mL wash and blow down to dryness.
 Immediately stop the nitrogen flow and remove the tube from the N-Evap.
- Add exactly 2.0 mL of HPLC or Milli-Q water to the nine containing the evaporated C18 cluate. Vortex mix for 5 seconds and place into the ultrasonic bath. Allow the sample to remain in the bath (set at 30 to 45 degrees) for 5 to 10 minutes
- Filter approximately 1.0 mL of this final volume solution through a 0.2 micron AcroDisc filter into an appropriate autosampler vial. Cap.
- This sample is now ready for LC/MS/MS analysis. It can be stored for up to 3 weeks in a refrigerator at 4 °C.

4.2.9b Concentration and Purification Procedures for Soil

Solution Requirements/Sample (extraction + purificatio	n):	
Milli- Q water	50	mL
Extraction solution	20	mL
methanol	10	шL
hexane	10	шĽ
90:10 acetone: 3 mM Acetic Acid	10	mL
90:10 methanol:3mM Acetic Acid	10	mL

SAMPLE CONCENTRATION AND PURIFICATION

SAMPLE PREPARATION AND EXTRACTION

- 1. Remove samples from refrigerator or freezer and allow to warm to room temperature.
- 2. Weigh out a 5 g sample of the soil and place it into a 50 mL centrifuge bottle.
- Fortify samples if required.
- Add 20 mL of 0.1M KH.PO./0.5M NaCl extraction solution. Manually shake 10 times.
 Place onto a wrist-action shaker and shake for 10 minutes at maximum speed.
- 5. Centrifuge the sample for 10 minutes at approximately 2500 tpm.
- Attach a 20 mL syringe to a 0.45 μm Acrodisc filter. Decant the supernatent into the 2 syringe. Filter into a 50 mL centrifuge tube. Place tube immediately onto an N-Evap at 40 to 50 °C and begin evaporation.
- 7. Repeat steps 4 through 5. When there is sufficient room in the 50 mL centrifuge tube (on the N-Evap) filter the second supernatent into it. Rinse syringe with 2 x 10 mL volumes of accione and place accione into the 50 mL centrifuge tube containing the filtrate.
- Evaporate the combined extract/rinse to the water phase (approximately 15 mL).
- Add HPLC water until the volume is approximately 50 mL
- 10. Just prior to proceeding to the SPE cleanup sonicate extract for 5 min. and vortex mix.

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ENVI-CARB BOND ELUT® CLEANUP

- Place the 1.5 g Envi-Carb Bond Eluts² into the vacuum manifold and condition them by
 placing 10 mL of 90:10 acctone:3 mM acetic acid solution into the tubes and adjusting
 the flow to a fast drip (-10 mL/minute). Using a gentle vacuum to pull through all of the
 liquid, tet the column dry for 30 seconds after the last of the liquid has exited. Load and
 condition the column with two 10 mL volumes of water. Do not let the liquid level drop
 below the top of the sorbent after the second water load has passed through. Do not
 allow the tube to dry.
- Load the soil extract onto the column, in several portions if necessary due to column
 reservoir capacity. Turn on the vacuum and allow the sample to pass through the
 column at a slow drip. Do not allow the column to go dry. Do not collect the liquid.
- Place 5 mL of water into the empty 50 mL centritige tube, vortex. Place this rinse onto
 the Bond Elut and allow to pass through at a slow drip. Pull all the water through, allow
 to dry, passing air through for about 30 seconds after last drop has come off.
- Place 5 mL of hexane into the Bond Elut and allow to pass through at a slow drip.
 Allow to dry, passing air through for about 30 seconds after last drop has come off.
- Open the vacuum manifold and place plastic 15 mL centrifuge tubes into the manifold in order to collect the cluate.
- 10. Place 10 mL of 90:10 acctone:3 mM acetic acid solution into the Bond Elut tube and allow to pass through at a slow drip. Pull all of the liquid through into the collection tubes. Break the vacuum and remove the tubes containing the chuste.
- 11. Place sample on an N-Evap and evaporate to approximately 1 mL at 40-50 deg.
- 12. Wash the sides of the sample test tube with approximately 2 mal of acetone and evaporate the extract to 0.2 to 0.5 mL. Repeat this washing with a second 2 mL volume of acetone. Evaporate the sample to dryness
- 13. Add 200 µL of acctone to the tube, vortex and sonicate for 5 minutes, vortex again. Add approximately 1 mL of water and evaporate to the water phase. Add approximately 4 mL of HPLC water and continue the evaporation step to ensure removal of all traces of acctone This is a very critical step, the presence of any acctone will prevent the polar analytes from being retained on the C18 column in later steps.)
- 10. Adjust to approximately 5 mL with HPLC water. Vortex, sonicate for 5 minutes, vortex
- Proceed directly to the C18 SPE clean-up. This is a stopping point. Samples can be
 placed into the refrigerator for storage for up to 1 week.

C18 BOND ELUT CLEANUP/FINAL PREPARATION

- Place the C18 Bond Elur® rubes onto the vacuum manifold and condition them by placing 5 mL of methanol:3mM Acetic Acid (90:10 v/v) solution into each tube and adjusting the flow to a fast drip. Drain the tube and allow to dry for 30 seconds more under vacuum. In the same manner condition next with 10 mL of water, in two 5 mL increments, waiting until the first 5 mL is completely below the first before adding the second 5 mL. Stop the flow when the level of the water drops slightly below the top of the sorbent in the bottom tube. Do not let the tubes dry.
- Place the Envi-Carb cluate solution (step 11 ENVI-CARB cleanup above) into the C18
 Bond Elur[®] tube. Apply vacuum, open the stopcock and load the cluate onto the column
 with a very slow drip. Stop when the level of the liquid is 1 cm above the top of the
 sorbent. Discard the aqueous effluent.
- Backrinse the original Envi-Carb cluste container by placing 2.0 mL of water into it vortex mixing for a few seconds. Pour this rinsate into the C18 cartridge. Allow this

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(rinse)water wash to pass through the C18 and discard. Allow all of the water to be drawn out of the tube. Immediately shut off vacuum, discard load volume

- 4. Place 5.0 mL of hexane into the C1S cartridge. Allow the hexane to pass through at a moderate drip. After the hexane has completely passed through the column, add another 5.0 mL of hexane and allow that to completely pass through the column as well. Allow the vacuum to pull air through the cartridge until there is no more dripping.
- Break vacuum and place a new 15 mL plastic centrifuge tube into the vacuum manifold under the C13 tubes to collect the cluste.
- Place 10 mL of methanol:3 mM Acetic Acid (90:10 v/v) solution into the column.
 Allow the cluste to pass through at a moderate drip and collect. Discard the C18 column.
- When elution is finished, place the tube containing the extract onto the N-Evap and blow it down to approximately 1 mL
- 11. Wash the sides of the tube with approximately 1 mL methanol and blow down to approximately 0.2 t 0.5 mL. Repeat with another 1 mL wash and blow down to dryness. Immediately stop the nitrogen flow and remove the tube from the N-Evap.
- 12. Add exactly 2.0 mL of HPLC or Milli-Q water to the tube containing the evaporated C18 cluate. Vortex mix for 5 seconds and place into the ultrasonic bath. Allow the sample to remain in the bath (set at 30 to 45 degrees) for 5 to 10 minutes
- Filter approximately 1.0 mL of this final volume solution through a 0.2 micron AcroDisc filter into an appropriate autosampler vial. Cap.
- This sample is now ready for LCMS/MS analysis. It can be stored for up to 3 weeks in a refrigerator at 4 °C.

4.3 Instrumentation

4.3.1 Description

This method requires an LCMS/MS instrument equipped with an electrospray interface for the analysis of water and soil. An autosampler is also required for the unattended analysis of multiple samples and calibration solutions. The analytical data in this method was obtained from a Micromass Quattro II LC/MS equipped with the HP Series 1100 Modular HPLC System which includes a dual channel pump, autosampler, vacuum degasser, temperature-controlled column compartment, and a UV-Vis detector (not required for this method). The HPLC and Mass Spectrometer operating parameters for these systems are given in section 4.3.2.

The chromatographic column and mobile phases specified provide good peak shapes and resolution of the analyses. This allows sufficient time to switch mass spectrometric acquisition parameters thereby allowing better optimization of each channel. The HPLC solvent program includes a period of high-organic solvent flow to purge matrix materials from the column and allows sufficient re-equilibration time between runs. Do not alter these periods. In general, instrument response is good to excellent for the analyses on the positive ESI SRM channels. Always use the <u>combination</u> of HPLC conditions and mass spectrometer parameters specified for each matrix.

4.3.2 Operating Conditions: Micromass Quattro II with HP Series 1100 HPLC

ANALYTICAL CONDITIONS FOR SOIL AND WATER

High Performance Liquid Chromatography (HPLC)

Mobile Phase A:

Aqueous 0.01 M acetic acid

Mobile Phase B:

Acetonimie

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Mobile Phase Program:	Time(mi	n) %A	<u>%B</u>	
•	0	100	0	
•	3	90	10	
•	10	50	50	
	15.0	25	75	
	15.1	5	95	
	20.0	5	95	
	20.1	0	100	
	30.0	0	100	
On-column Flow Rate:	1.0 mL min,	split –5:1, D	ivert 0-6.9,	15.5-30 min
Injection Volume:	50 µL			
Column:	Zorbax RX-(column attac		.6 mm id, 5	μm; with guard
Cohumn Temperature:	30 °C	464		
Mass Spectrometer (MS)				
Ionization Mode:	Positive ESI			4
HV Lens	0.26 KV			
Capillary Voltage:	4.0 KV			
Cone Voltage	32 Volts			
Collision Voltage	18			
General parameters Collision Gas Pressure MS1 LM/HM Resolution MS2 LM/RM Resolution Source Temperature:	Positive ESI 1.7-2.0x10e- both 10 both 12 85 °C			
Acquisition Functions	set as shown	below:		
1 ESP+	Parent	<u>Dauehter</u>	De	vell
Start 7.0min End 9.5	171.1	71.1		2 sec
Int Chan. Delay: 0.02 sec				
2 ESP+	Parent	Daughter		<u>vell</u>
Start 9.5min	255.3	157.1		2 sec
End 12.0	267_2	171.0	1.0	
Int Chan, Delay; 0.02 sec	269.3	171.0	. 0.1	2
3 ESP+	Parent	Daughter	_ D	well
Start 12.0min	239.2	157.0		08
End 17.0	253.2	171.1		.08
Int Chan, Delay: 0.02 sec				

Approximate Retention Times of the 7 Analytes

Analyte	Approx. Retention Time (min)	Acquisition Function: Transition
membolite G3170	8.39	1: 171.1 -71.0
membolite A	10.63	2: 269.3-171.0
membolite A-1	11.62	2: 269.3-171.0

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membolite I	11.27	2: 267.2-171.0
membolite C	10.12	2: 255.3-157.0
hexazinone	14.46	3: 253.2-171.0
membolite B	13.62	3: 239.2-157.0

Alternate Mass Spectrometer (MS) Conditions: Confirmation Channels

If co-cluting matrix components interfere with any of the analytes on their primary positive electrospray MS/MS channels, use the following electrospray conditions and alternate channels to both confirm that these co-cluting peaks are in fact interferences and to quantitate the analytes. Alternate channels can be determined from examining the full scan spectra or the daughter ion spectra of each analyte shown in Figure 2. The alternate electrospray parameters (increased collision gas energy) specified will also significantly reduce or eliminate the interfering peaks on the original channels. Re analyze samples using the alternate channels along with the originals for comparison.

Ionization Mode:	<u>Positive ESI</u>
HV Lens	0.1 KV
Capillary Voltage:	3.6 KV
Cone Voltage	25 Volts
Collision Voltage	30
General parameters	Positive ESI
General varameters Collision Gas Pressure	Positive ESI 1.7-2.0x10e-3
Collision Gas Pressure	1.7-2.0x10e-3
Collision Gas Pressure MS1 LM/HM Resolution	1.7-2.0x10e-3 both 8

Acquisition Functions

set for selected alternate MRM transitions:

4.3.3 Calibration Procedures

Instrumentation performance was initially evaluated for calibration, response, linearity by the following procedures:

- The LCMSMS instrument was used in positive electrospray mode to optimize the [M-cyclohexane moiery]" peak while still showing the base peak [MH]". For the negative electrospray compounds, the instrument was tuned to maximize the [M-1]" peak.
- Full-scan, on-column analysis of a ≥ 4-µg/mL calibration solution containing each of the
 analytes was performed to verify appropriate m/z responses and abundance ratios.
- Analysis of calibration solutions (see section 4.2.5 for preparation) and construction of a
 5-point calibration curve was performed to verify linearity (R² ≥ 0.98) over the analytical
 range and evaluate detector response (low calibration solution response ≥ 5/1 signal-tonoise).

Routine manufacturer instrument calibration and maintenance procedures should be followed as necessary to optimize performance. It is also accessary to ensure that the electrospray interface components are clean. A dirty source can cause drift in instrument response.

43.4 Sample Analysis

Initially, at least 1, and preferably 2 runs of a matrix-containing sample should be made to equilibrate the ESI-LCMS system prior to running a sample set unless instrument performance allows otherwise.

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Calibration solutions and samples should be alternately analyzed through the analysis set so test samples can be quantified using the average response of bracketing standard (i.e., standard, sample, standard, sample, standard, etc.) If instrument performance is stable it is permissible to run 2 samples between each standard, provided a standard precedes the first sample of the set and follows the last sample of the set. At a minimum, a check sample consisting of a control fortified at the LOQ and processed through the method should be run with every sample set.

4.4 Calculations

4.4.1 Methods

The ppm found and percent of applied recoveries are determined from calculations which compare the peak area responses of each malyte to the response of bracketing analytical standards. Use <u>only</u> the primary MRM channel for quantitation. <u>Do not use the Total Ion Chromatogram</u>. Peak area response values should be entered in an EXCEL® spreadsheet template designed to perform the appropriate calculations.

The response factor (RF) is calculated for each analyte in all calibration standard analyses by dividing the peak area by the analyte concentration. One may use either the average response factor of the entire set of standards if the response has been stable over the period of analysis, or one may use the average of the closest two standards which bracket the sample being quantitated if instrument drift has been noted. In either case, valid recovery data are generated if the percent relative standard deviation (%RSD) for the peak area responses of the standards used is less than or equal to 20% Values for the amount (ppm) detected in samples are determined by comparing the peak area of each sample analyte detected with the average response factor and correcting for aliquots, weight of sample, and final volume.

The percent of applied recoveries for fortified samples are calculated by dividing ppm found by ppm applied and multiplying by 100.

C (conc. .of sample μ g/mL) = Area(sample)/RF(avg.)

ppb (sample) = C/W * (1/AF) * FV

W = weight of sample (g, 1 mL = 1g)

AF = aliquot fraction (vol. aliquot/total vol.)

FV = total volume of final analysis solution (mL)

Sample Calculations (Bracketing Standards): For a hypothetical analyte:

20 mL (+20g) water sample fortified at 0.100 pph, aliquot fraction = 1, Final Volume = 2 mL

Response of 0.67 ng/mL Std = 700 area counts Response of 2.0 ng/mL Std = 2200 area counts Response of 0.100 ppb fortification = 1200 area counts

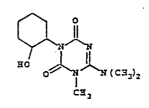
RF(avg.)= [(2200/2.0 ng/mL) + (700/0.67 ng/mL)]/2 = 1072(mL/ng)Conc. sample = 1200/1072(mL/ng) = 1.119 ng/mLAF = 20mL/20mL = 1.0

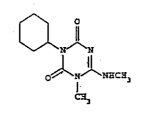
ppb sample = [1.119 (ag/mL)/20g] * (1/1) * 2 mL = 0.1119 ng/g = 0.1119 ppb recovery = 0.1119/0.1000 * 100% = 112%

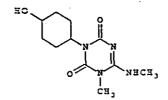
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FIGURE 1. CHEMICAL STRUCTURES AND NAMES

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Hexazinone
DuPort Code No.: DPX-A3674

Chemical Name: S-Triazine-2,4-(1H,3H)-cione, 3-cyclohexyl-6-cimethylamino -1-methyl

CAS Registry No.: 51235-04-2

Metabolite A <u>DuPost Code No.</u>: T3937

Chemical Name: S-Triazine-2,4-(1H,3H)-dione, 6-dimethylamino-3-(4-hydroxycyclo- hexyl)-1-methyl

CAS Registry No.: none

Metabolite A1 DiPont Code No.: G3453 ~

Operical Name S-Trissine 24(1H,3-1)-done, 3(trans-2-ydroxycyddrexy)-1-metryl-8-dmetrylamino-

CAS Registry No.: none

Metabolite B <u>DuPort Code No.:</u> IN-A3928

Chemical Name: S-Triazine-2,4-(1H,3H)-dione, 3-cyclohexyl-1-methyl-6-(methylamino)-

CAS Recistry No.: 56611-54-2

Metabolite C

DiPort Code No.: IN-T3935

Chemical Name: S-Triazine-2,4-(1H,3H)-dione, 3-(4-hydroxycydohexyl)-1-methyl-6 (methylarrino)-

CAS Registry No.: 72585-88-7

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FIGURE 1 (CONTINUED) CHEMICAL STRUCTURES AND NAMES

Metabolite C1 DuPart Code No.: IN-G3454

Chemical Name: S-Triazine-2,4-(1H,3H)-cione, 3-(2-hydroxycyclohexyl)-1-methyl-6- (methylamino)-

CAS Fedistry No: none

Metabolite D DuPont Code No.: IN-82838

Chemical Name: S-Triazine-24,6 (1H,3+(5-)-trione, 1-oycidrex/H-methyl

CAS Registry No.: none

Metabolite E DuPort Code No: IN-T3936

Chemical Name: S-Triazine-2,4,6-(1H,3H,5H)trione, 1-(4 hydroxycyclonexyl)-3-methyl

CAS Becisty No.: none

Metabolite F

DiPort Carle No.: IN L3221

Chemical Name: S-Triazine-2,4-(1H,3H)-dione, 3-cyclohexyl-6-arrino-1-methyl-

CAS Recisty No.: rone

Metabolite G DuPort Code No.: IN-T4916

Chemical Name: S-Triazine-2,4-(1H,3H)-dione, 3-cyclohexyl-6-methylamino

CAS Peristry No.: none

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FIGURE 1.(CONTINUED) CHEMICAL STRUCTURES AND NAMES

Metabolite H Di Port Code No.: IN-M3111

Openical Name: S-Trizzine-2,4-(1H,3H)-dione, 3 cyclonexy-6-dimetrylamino CAS Registry.No.: none

Metabolite 1 D. Part Orde No: INUS 472

Chemical Native S-Triacine-2,4-(1H,2H)-Clone, 6-(dimetrylatrino) -1-metryl-3-(4-macyclohexyl)

CAS Registry No: none

Metabolite G3170 D. Prot Corie No.: IN-G3170

Chemical Name: S-Triazine-2,4-(1H,3H)-done,1-methyl-6-dimethyl amino

CAS Pecistry No.: none

Metabolite G3170 n-glucoside D_Pmt Code No.: IN-NC333

Chemical Name: S-Triazine-2,4-(1H,3H)-done, 1-methyl-6-dimethylamino n-glucoside

CAS Fedistry No.: none

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Modifications to Dupont-2549 to confirm by LC/MS/MS

If sample splitting is needed to allow confirmation by LC/MS/MS, the 16x100 mm silanized glass tube used in step 4.4.5 must be calibrated at 1.0 mL as well as 2.0 mL.

Follow the method (Met-101, revision 2) through step 4.5.1. At step 4.5.1 insert at end. Adjust to 2.0 mL with methanol, vortex to mix, sonicate 5 minutes. Readjust to 2.0 mL with methanol and vortex. Remove 1 mL for LC/MS/MS if required.

Evaporate to dryness. Continue with method DuPont-2292 (soil and water LC/MS/MS method) final preparation step 4.2.9: C18 Bond Elut Cleanup/Final Preparation, step 8.

Standards for LC/MS/MS analysis may be prepared from standard solutions used for GC/NPD analysis by evaporating the organic solvent and redissolving the residue in water with mixing and sonication.



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PROTOCOL DEVIATION		
Deviation Number:1 Date of Occurrence:05/06/99		
CAL Study Number:008-036 Sponsor Study Number:DuPont-2547		
DESCRIPTION OF DEVIATION		
Step 4.2.9a, C18, Step 9 of DuPont-2292, located in Protocol Appendix 2, states that exactly 2.0 mL of HPLC water should be added to the tube containing the evaporated C18 eluate. Instead, for the reagent blank, sample controls, and spikes A-E, the volume to be added to the tube was calculated as described in step 4.5.4 of DuPont-2549 (Protocol Appendix 1), so that the concentration of the sample in the final extract was 2.5 g/mL.		
ACTIONS TAKEN		
i.e., amendment issued, SOP revision, etc		
Deviation issued.		
Recorded By/Date: Shaoshi sheng 5-17-99		
IMPACT ON THE STUDY		
No negative impact on the study.		
Study Director Signature Study Director Signature 5-17-99 Date		
Study Director Signature Date CAL QAU Review GW 5 19 99		
February 12, 1998/		