

EPA Method 532.0 Determination of Phenylurea Compounds in Drinking Water by Solid Phase Extraction and High Performance Liquid Chromatography with UV Detection

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This is an Application Brief and does not contain a detailed Experimental section.

Abstract

This application brief demonstrates the determination of phenylurea compounds in drinking water by solid phase

extraction and High Performance Liquid Chromatography with UV detection.

Introduction

Phenylurea pesticides that were introduced worldwide in the 1950s are used commercially on a wide range of

food crops. Subsequently determined to possess a significant toxicological risk, the agricultural runoff of these

compounds may be found in drinking water supplies. The United States Environmental Protection Agency

(US EPA) requires that drinking water and raw surface water be monitored for the presence of phenylurea

pesticides and related compounds using EPA Method 532.0. The European Union (EU) regulation regarding

drinking water (EC Directive 98/83/EC), provides a general rule for pesticides and metabolites. This regulation

limits the maximum admissible concentration (MAC) at 0.1 ppb for each individual component, with the total

concentration not to exceed 0.5 ppb.

Experimental

HPLC Conditions

Instrument: Waters Alliance HPLC System with 2996 PDA

Detector

Eluent: A: 25 mM Phosphate, pH 2.4

B: Acetonitrile

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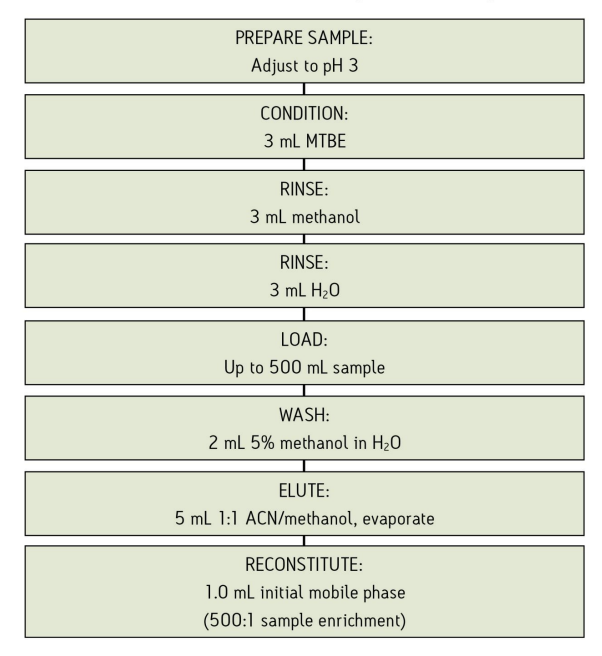
Column: SunFire C_{18} , 3.5 μ m 4.6 x 150 mm @ 30 $^{\circ}$ C

Injection: 20 µL

Flow rate:	1.5 mL/min
Detection:	PDA UV @ 245 nm
Data:	Waters Empower Software

Sample Preparation

Oasis® SPE Method for Pheylurea Pesticides Method for Oasis HLB Cartridge, 6 cc, 200 mg



Standard Preparation

Pipette 100 μ L of AccuStandard mix M-532 and 20 μ L mix M-532-SS into 880 μ L 1:1 water/acetonitrile (10 ppm analytes plus surrogates).

Eleuent Preparation

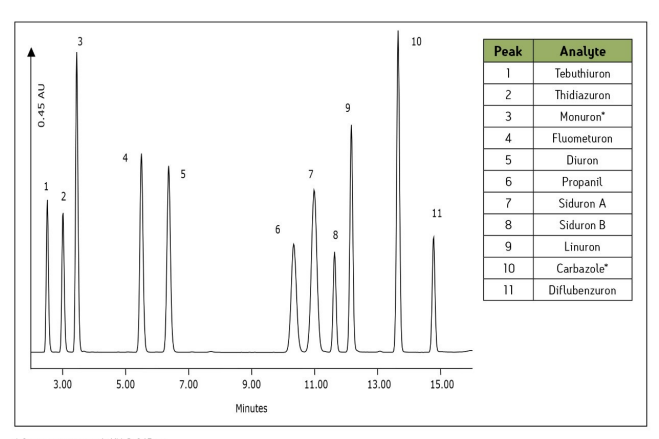
A: 25 mM phosphate

Dissolve 1.7 g of potassium dihydrogen phosphate (KH_2PO_4) and 850 μL phosphoric acid (H_3PO_4) in 100 mL water. Dilute to 1 L, then filter and degas. Verify that the pH is approximately 2.4.

B: Acetonitrile

Time	Flow	%A	%B	Curve
-	1.5	60	40	-
9.5	1.5	60	40	6
10.0	1.5	50	50	6
14.0	1.5	40	60	6
15.0	1.5	60	40	6

Eluent gradient.



^{*} Surrogate compounds UV @ 245 nm Standard chromatogram, 10 ppm each analyte.

Related Documents

- Multi-Residue Analysis of Priority Pollutants in Drinking and Surface Waters Using Solid-Phase Extraction 720001438EN https://www.waters.com/nextgen/us/en/library/application-notes/2007/multi-residue-analysis-of-priority-pollutants-in-drinking-and-surface-water-using-solid-phase-extraction-and-gc-tandem-quadrupole-ms-ms.html>
- Food and Environmental Residue Analysis 720002274EN
 https://www.waters.com/webassets/cms/library/docs/720002274en.pdf>
- Environmental Applications Book <u>720002123EN
 https://www.waters.com/webassets/cms/library/docs/720002123en.pdf>
 </u>

Featured Products

Alliance HPLC System https://www.waters.com/534293

Empower Chromatography Data System https://www.waters.com/10190609
720002727, August 2008
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