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응용 자료

Novel Extraction Techniques with ACQUITY UPLC with 2D Technology: Part I Pesticides Screening in Drinking Water

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Abstract

This application demonstrated the effectiveness of two extraction process, single stage captive and triple stage captive for pesticides screening by 2D LC ToF in drinking water.

Benefits

- Fast 30-minute extraction protocol
- · Trace level detection at parts per trillion
- · 90 second homogenization

Introduction

Many countries around the world have strict regulatory guide lines for drinking water quality. To satisfy legislative requirements, analytical methods have been developed to monitor a wide range of contaminants at trace levels using analytical techniques such as gas chromatography/mass spectrometry (GC-MS) or liquid chromatography/tandem quadrupole mass spectrometry (LC-MS/MS).

Trace level analysis at ppt (part per trillion) constitute the bulk of the work load for the majority of testing laboratories worldwide. Current analytical techniques use a combination of extraction procedures, often requiring an enrichment process and accurate detection for any given target analyte. As such, large sample volumes are usually extracted using various manual extraction methods (i.e., solid-phase extraction (SPE), liquid-liquid, etc.) and are concentrated into a smaller volume. As an example, a typical extraction method usually starts with a 500 mL of sample and ending up with a final volume of a 100 μ L (5000:1 enrichment ratio). If higher sensitivity is required, the only alternative left is to process larger sample volume but will require an increase in time and manual labor.

In recent years, efforts are now being diverted to investigate effective screening methods with high resolution Time-of-Flight (ToF) instruments and with the capability of reaching sub ppb (part per billion) levels. With current single chromatography separation setup and the inherent low sensitivity of ToF instrument compared to tandem quadrupole MS, this demand is quite difficult to achieve. As such, a new analytical strategy is needed to reach those goals. This application will discuss the performance of 2D LC-QToF setup for the analysis of pesticides residues in drinking water at sub ppb level. With an enrichment factor of 20:1 from a

rapid fractionation sample preparation protocol using two mixed mode sorbents, the gap between method and instrument limits of quantitation (LoQ) can be eliminated with large volume injection. Furthermore, by using an At-column dilution 2D LC configuration, 100% organic solvent extracts can be injected directly, thus eliminating all evaporation and reconstitution steps from any sample preparation protocol.¹⁻⁴

Experimental

Two MRM transitions, quantification and confirmation, for each pesticide were selected and optimized. The MRM conditions are listed in Table 1.

For this application, finding the optimum extraction and chromatographic condition for this multi-residue analysis posed a significant challenge. The chromatographic conditions were tested on several trapping chemistries (Oasis HLB, XBridge C_{18} , and XBridge C_{8}) and separation chemistries (BEH C_{18}) The loading (low pH, high pH, and neutral pH) and eluting mobile phase (MeOH + 0.5% formic acid and ACN + 0.5% formic acid) were also optimized using an automated 6x6 process.

All pesticides standards were purchased from Sigma Aldrich. The extraction process was performed on preconditioned reversed-phase sorbent Oasis HLB SPE Cartridge, 6 cc, 150 mg, (p/n: 186003365) for the captive extraction or a dual mixed-mode Oasis MCX SPE Cartridge, 6 cc, 150 mg (p/n: 186000256) and MAX SPE Cartridge, 6 cc, 150 mg, (p/n: 186000369) for the screening extraction.

Chromatography and MS/MS conditions

Loading conditions

Column:	Oasis HLB Direct Connect HP, 20 μ m, 2.1 \times 30				
	mm (p/n: 186005231)				
Loading:	MilliQ water (pH 7, no additives)				
Flow rate:	2 mL/min				
At-column dilution:	5% (0.1 mL/min loading pump and 2 mL/min				
	diluting pump)				

UPLC conditions

UPLC system:	ACQUITY UPLC with 2D Technology configured for "Trap and Elute" with At-column dilution				
Runtime:	10 min				
Column:	ACQUITY UPLC BEH C_{18} , 1.7 μ m, 2.1 \times 50 mm (p/n: 176000863)				
Column temp.:	60 °C				
Mobile phase A:	Water + 0.5% formic acid				
Mobile phase B:	Acetonitrile + 0.5% formic acid				
Elution:	5-minute linear gradient from 5% (B) to 95% (B)				
Flow rate:	0.500 mL/min (Elution pump)				
Injection volume:	100 μL				
MS conditions					
MS system:	Xevo TQ-S				
Ionization mode:	ESI positive				
Capillary voltage:	3.0 kV				
Cone voltage:	90.0 V				
Source temp.:	150 °C				
Desolvation temp.:	550 °C				

Desolvation gas: 1100 L/hr

Cone gas: 50 L/hr

henyl Urea	MW	Cone (V)	Parent mass	Quant	CID	Qual	CID
duron	232.3	30	233.1	137.0	15	94.0	20
imefuron hlorobromouron	338.8 293.5	30	339.0 294.9	72.1 205.9	25 20	166.9 182.0	20 15
ifenoxurone	293.5	30	294.9	72.1	20	123.1	20
uometuron	232.2	30	233.0	72.1	20	46.1	15
nidiazuron	220.3	30	221.0	102.0	15	128.0	15
etobromuron	259.1	30	258.9	169.9	20	148.0	10
hloroxuron	290.7	30	291.0	72.0	20	164.0	15
nifensulfuron methyl	387.4	30	388.0	167.0	20	204.9	25
oproturon	206.3	30	207.2	72.1	15	165.0	20
onolinuron	214.6	30	215.1	126.1	15	148.1	10
ibenuron methyl	395.4	30	396.1	155.1	20	181.0	20
onuron	198.7	30	199.1	72.0	15	46.1	15
iuron	233.1	30	233.0	72.1	15	46.1	15
uturon	236.7	30	236.7	84.1	15	126.0	30
etsulfuron methyl	381.4	30 30	382.1	167.0 159.9	15 20	199.0	30 15
nuron hlortoluron	249.1 212.7	30	249.0 213.1	72.0	15	182.0 46.1	15
enuron	164.2	30	165.9	72.1	15	46.1	15
etoxuron	228.7	30	229.2	72.1	15	46.1	20
iazole	220.1	30	223.2	12.1	10	40.1	20
aconazole	705.6	30	705.1	392.1	30	432.1	30
uconazole	306.3	30	307.1	238.1	15	220.1	15
etoconazole	531.4	30	531.1	82.1	40	489.1	35
priconazole	349.3	30	350.1	127.0	30	281.1	15
osaconazole	700.8	30	701.3	683.2	30	127.0	60
avuconazole	437.5	30	438.0	224.0	20	215.0	20
fenoconazole	406.3	30	406.1	250.9	30	337.0	15
opiconazole	342.2	30	342.1	159.0	25	69.1	20
proconazole	291.8	30	292.1	70.0	15	125.0	25
othioconazole	344.3	30	344.1	326.0	10	189.0	20
buconazole	307.8	30	308.2	70.0	20	125.0	30
arbendazim	191.2	30	192.1	160.0	15	132.1	30
rganophosphorus							
hlorpyrifos	350.6	30	349.8	96.9	30	197.9	20
arathion methyl	263.2	30	263.9	125.0	20	231.9	15
zinphos methyl	317.3	20	318.0	132.0	15	125.0	20
crotophos	237.2	30	238.0	112.1	15	193.0	10
azinon	304.3	30	305.0	169.0	15	153.0	15
methoate	229.3	30	230.1	198.9	10 25	125.0	20 25
zinphos ethyl	345.4		346.1	96.9		137.0	
ichlorvos alathion	221.0 330.4	30	220.9 331.0	109.0	15 10	79.0 99.1	25 20
enitrothion	277.2	30	277.9	125.1	20	246.1	15
arathion	291.3	30	292.1	235.9	15	94.0	30
ropetamphos	281.3	30	282.1	138.0	20	156.0	15
evinphos	224.2	30	225.0	127.0	15	193.0	5
arbamate						1000	
ldicarb sulfoxide	206.3	15	207.1	89.0	15	132.0	5
xamyl	219.3	30	242.1	72.0	15	121.0	10
dicarb	190.3	30	213.1	89.1	15	116.0	10
ethiocarb sulfone	257.3	30	258.1	122.0	15	201.0	10
dicarb sulfone	222.3	30	223.1	86.1	15	148.0	10
minocarb	208.3	30	209.2	152.1	15	137.0	20
arbofuran	221.3	30	222.1	165.0	10	123.1	20
osulfocarb	251.4	30	252.1	91.0	15	128.1	10
ethiocarb	225.3	30	226.1	169.0	10	121.0	20
nobucarb	207.3	30	208.2	95.0	15	152.0	10
arbetamide	236.3	30	237.2	118.1	10	192.0	10
arbofuran-3-kto	235.2	30	236.2	179.0	10	161.0	15
enoxycarb	301.3	30	302.1	88.0	20	116.1	10
arbaryl	201.2	30	202.1	145.0	10	127.0	25
arbofuran-3-hydroxy ethiocarb sulfoxide	237.3 241.3	30 30	238.2	181.0 185.0	10 15	163.0	15 25
iazines	241.3	30	242.1	100.0	19.0	122.1	25
razines razine-desethyl-desisopropyl	145.6	30	146.1	79.0	15	104.0	15
opazine	229.7	30	230.2	146.0	20	188.0	15
netryn	227.3	30	228.2	186.0	20	96.0	25
rbutryn	241.4	30	242.2	186.0	20	91.0	25
ietazine	229.7	30	230.2	99.0	25	132.0	20
razine-desisopropyl 2 hydroxy	155.2	30	156.1	86.0	15	69.0	20
ometryn	241.4	30	242.2	158.0	25	200.0	20
razine-desethyl	187.6	30	188.1	146.0	15	79.0	25
rbuthylazine	229.7	30	230.2	174.0	15	96.0	25
netryn	213.3	30	214.1	124.1	20	96.0	20
mazine	201.7	30	202.2	132.0	20	124.1	15
	215.7	30	216.1	174.0	15	96.0	25
	173.6	30	174.1	132.0	20	96.0	15
razine desisopropyl							
razine desisopropyl thers orasulam	359.3	30	360.1	129.0	25	192.0	15
razine desisopropyl thers orasulam	359.3 256.1	30 30	360.1 256.1	129.0 189.9	25 15	192.0 172.9	15 20
razine desisopropyl thers orasulam opyzamide sulam	256.1 230.2		256.1 231.1	189.9 156.0	15 10		20 20
razine desisopropyl thers orasulam opyzamide sulam entazon	256.1 230.2 240.3	30 30 30	256.1 231.1 241.1	189.9 156.0 199.0	15 10 10	172.9 92.0 107.1	20 20 20
razine desisopropyl thers orasulam opyzamide sulam entazon ufenacet	256.1 230.2 240.3 363.3	30 30 30 30	256.1 231.1 241.1 364.1	189.9 156.0 199.0 152.0	15 10 10 20	172.9 92.0	20 20 20 10
trazine trazine desisopropyl thers orasulam ropyzamide sulam entazon ufenacet iflufenican	256.1 230.2 240.3 363.3 394.3	30 30 30 30 30	256.1 231.1 241.1 364.1 395.1	189.9 156.0 199.0 152.0 266.0	15 10 10 20 25	172.9 92.0 107.1 194.1 246.0	20 20 20 10 35
trazine desisopropyl thers orasulam opyzamide sulam entazon ufenacet iflufenican endimethalin	256.1 230.2 240.3 363.3 394.3 281.3	30 30 30 30 30 30	256.1 231.1 241.1 364.1 395.1 282.2	189.9 156.0 199.0 152.0 266.0 212.1	15 10 10 20 25 10	172.9 92.0 107.1 194.1 246.0 194.0	20 20 20 10 35 20
trazine desisopropyl thers orasulam ropyzamide sulam entazon ufenacet	256.1 230.2 240.3 363.3 394.3	30 30 30 30 30	256.1 231.1 241.1 364.1 395.1	189.9 156.0 199.0 152.0 266.0	15 10 10 20 25	172.9 92.0 107.1 194.1 246.0	20 20 20 10 35

- 2. Mallet, C.R., Botch-Jones, S., J. Anal. Toxicology, 1-11, 2016.
- 3. Mallet, C.R, Multi-dimensional Chromatography Compendium: Trap and Elute vs. At-column dilution, 720005339EN 2015.
- 4. Mallet, C.R., Analysis of Pharmaceuticals and Pesticides in Bottled, Tap and Surface Water Using the ACQUITY UPLC with 2D Technology, Waters Corporation, 720005167EN 2014.

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