

Scottish Agricultural Science Agency

A simplified multi-residue method for the rapid screening and confirmation of pesticides present in fruit and vegetable crude extracts using isocratic HPLC separation combined with electrospray tandem mass spectrometry.

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<u>AIM:</u> To develop a generic LCMSMS method that improves the analytical efficiency of pesticide multi-residue analysis.

Introduction

The application of atmospheric pressure ionisation LCMSMS techniques has facilitated the multi-residue analysis of crude extracts from fruit and vegetables. The enhanced selectivity of MSMS can significantly reduce potential interference from non-target substances and discriminate between co-eluting and isobaric analytes. The utility of electrospray MSMS detection in combination with isocratic HPLC separation was investigated to see if further efficiency gains could be made compared to gradient separations used previously in our laboratory

Experimental

<u> HPLC Isocratic Method – Agilent 1100 HPLC system</u>

Column: Hypersil C_{18} 3 μ m BDS (4.6 x 100mm I.D.)

Mobile Phase: Methanol: 10mM aqueous ammonium acetate (70:30v/v)

Flow Rate: 0.5mlmin⁻¹ (post-column split → approx. 20µlmin⁻¹ into ion source)

Temperature: 35°C Injection volume: 5µl or 10µl

<u>Mass Spectrometry Method – Micromass Quattro Ultima</u>

Acquisition: Electrospray ± ive ion mode

Multiple Reaction Monitoring (MRM)

Single Ion Recording (SIR) for 2-phenylphenol

Collision Gas: Argon approx. 1.4x10⁻³ mbar

Data System: MassLynx 3.4
Desolvation Gas: Nitrogen approx. 500 lhr

Nebuliser Gas: Nitrogen Nitrogen

Cone Gas: Nitrogen @ 80lhr-1

Desolvation Temp: 350°C Capillary voltage: 3kV Source Temp: 150°C

Analytical Procedure

Extraction: Homogenisation with ethyl acetate/sodium sulfate/sodium hydrogen carbonate followed by methanol

solvent exchange ($\equiv 0.4g$ sample per ml)

Clean-up: None required – Crude extract filtered (0.45 µm PTFE Acrodisc syringe filter)

Results and Discussion:

The various pesticide/commodity combinations that have been analysed using this method are shown in Table 1. The target reporting levels (RL) used for each pesticide/commodity combination, were consistent with the levels set for the relevant part of the 2001 UK pesticide residue surveillance programme. All of the pesticides, and any important metabolites, yielded ions characteristic of the molecular weight of the neutral molecule (M) i.e. [M+H]+, [M+Na]+ or [M+NH]+ in positive ion mode or intense [M-H]- ions in negative ion mode. Structurally diagnostic product-ions were generated for each compound following collision-induced dissociation (CID) of the selected precursor ion with the exception of 2-phenylphenol [M-H]- precursor ion which remained intact. MSMS parameters determined following optimisation experiments are listed in table 2.

Table 1. Mixtures of pesticides sought in each commodity and target reporting level (RL) $\,$

P es ti ci de	RL				Com	modity		
	(mg kg ' 1)	Apple	Grape	Kiwi	Lemon	Peach	Sp in ach	Strawbe rry
2.4-D (free acid)	0.05							
Aldicarb	0.05							
Aldicarb sulfone	0.05				- 1			
Aldicarb sulfoxide	0.05							
Azoxystrobin	0.05							
Bendiocarh	0.50	- 1		- 1				
Butocarboxim	0.20			- 1				
Butocarboxim sulfone	0.20			- 1	- 1			
Butocarboxim sulfoxide	0.20			- :	- 1			
Carbaryl	0.01			- 1	- 1			
Carbondazim	0.01			- :	- :	•	- 1	- 1
Carbo fu ran	0.05	•		- 1	- 1			•
Carbo furan 3-h ydro xy	0.05			- 1	- 1			
Dichlofluanid	0.05			- 1				
Die tho fen earh	0.05		•			•		
Ethiofenearh	0.20			- :				
Fenhexamid	0.05			- 1				
Furath io carb	0.05					•		
lm azal il	0.02			•				
Kres ox im-methyl	0.05		•				- 1	- 1
Methiocarh	0.20	•		- :		- :	•	•
Methiogarh sulfone	0.20			- 1				
Methiocarb sulfoxide	0.20			- 1				
Methomyl	0.05			- :		•		
Metolearb	0.05			- 1				
Myclobutanil	0.05			- 1				
O xam vl	0.05	•				•	•	•
2-phenylphenol	0.10			- 1				
Penconazole	0.05				•			
Propie on azole	0.05	•						•
Pymetrozine	0.05							
Pyrime thanil	0.05							
Tebuconazole	0.05	•						•
Thiabendazole	0.05							
Thiodicath	0.05	•	•	- 1		•		•
Thio phan atc-methyl	0.10							
Triflox vs tro bin	0.05			- 1				1

Table 2. Optimum experimental parameters used for screening purposes.

PESTICIDE	RM M ⁸	Precursor ion assignment	MSMS transition	Cone Voltage (V) & Collision Ener gy (e V
2,4-D (free acid)	220	[M - H]	219 → 161	30.10
Aldicarh	190	[M+Na]	213 → 89	25.25
Aldicarh sulfone	222	[M+NH ₄]	240 → 86	25.20
Aldicarb sulfoxide	206	[M+H1	207 → 89	20.15
Az ox vs trob in	403	[M+H]	404 → 372	34.10
Bendiocarb	223	[M+H]	224 -> 167	30,15
Butoc arbox im	213	[M+H1	213 → 75	34.10
Butoc arbox im sulfone	222	[M+Na]	245 → 130	25,20
Butoc arbox im sulfoxide	206	[M+H1	207 → 75	30.15
Carbaryl	201	[M +H]	202 -> 145	20,15
Carbendazim	191	[M+H]	192 → 160	35,15
Carbofuran	221	[M+H]	222 → 165	27,15
Carbofuran 3-hydroxy	237	[M+H]	238 → 181	10,15
Dichlofluanid	332/334 ^b	[M+H]	333/335 → 224/226	20,15
Die tho fene arb	267	[M+H]	268 → 226	19,7
E thio fencarb	225	[M +H]	226 → 107	26,10
Fenhexamid	301	[M+H]	302 → 97	25,25
Furathioc arb	3 82	[M +H]	383 → 195	26,15
Imazalil	296	[M +H]	297 → 159	42,15
Isoprocarb	193	[M+H]	194 → 95	26,11
Kresoxim-methyl	313 b	[M+H]	314 → 206/222	23,10
Methiocarb	225	[M+H]	226 → 109	25,10
Methiocarb sulfone	257	[M +H]	258 → 226	25,10
Methiocarb sulfoxide	241	[M+H]	242 → 185	25,10
Me thom yl	1 84	[M+H]	185 → 128	27,10
Metolearb	165	[M+H]	166 -> 109	34,10
M yelobutanil	288	[M+H]	289 → 70	25,20
Oxamyl	241	[M +H]	242 → 72	21,20
2-phenylphenol Penconazole	170	[M - H]	169 (SIR)	21,n/a
	2 83	[M +H]	284 → 159	30,30
Propic onazole	341	[M+H]	342 → 159	25,30
Pymetrozine		[M +H]	218 → 105	24,28
Pyrime thanil Tehuconazole	199 307	[M+H] [M+H]	200 → 107 308 → 70	30,28 30,50
Thiahendazole	201			
Thiabendazole Thiodicarh	354	[M +H]	202 → 175	30,20 25,20
Thiogicare Thiophanate-methyl	342	[M +H]	355 → 88 343 → 151	25,20
Thiophanale-methyl Trifloxystrobin	342 408	[M+H] [M+H]	343 → 151	28,24 20,25
Relative Molecular		[M TI]	+02 7 : 60	20,23

^aRelative Molecular Mass ^b Sum of two characteristic transitions

The selectivity of MSMS and use of Multiple Reaction Monitoring (MRM) data acquisition procedures is demonstrated in figures 1a, and 1b, which show ion chromatograms obtained following analysis of nine pesticides sought in spinach and twelve of the twenty pesticides sought in kiwi-fruit.

Quantification was carried out from interpolation against calibration data generated using matrix-matched standards that covered the analyte concentration range of interest. The use of matrix-matched standards was necessary to compensate for signal

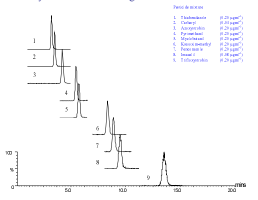


Figure 1a. Ion chromatograms of nine pesticides sought in spinach. Time-scheduled data acquisition sequence required 2 sets of three, 1 set of 2 and 1 single MRM channels.

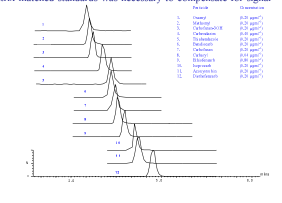


Figure 1b. Ion chromatograms of twelve pesticides (from 20) sought in kiwi fruit. 1 set of 5, 1 set of 4 and 1 set of 2 and a single MRM channels used.

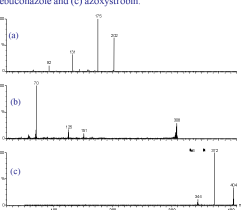
suppression observed in matrix compared to the response in pure solvent. The concentration range of matrix-matched standards spanned 0.005- $0.8~\mu gml^{-1}$. Correlation coefficients (r) greater than 0.99 were achieved routinely, following calibration of each pesticide.

The % recovery of each pesticide from organic produce that had been fortified at concentrations equivalent to the reporting level and at 4 times the reporting level (in triplicate at both levels) was determined. Mean recoveries greater than 60% were obtained for all the pesticides examined with the majority of recoveries better than 70%, although there were a few exceptions. Factors such as analyte instability in solution, extraction protocol and possible matrix effects however, resulted in inferior or irregular recovery of methiocarb and metabolites (peach/kiwi fruit), thiophanate-methyl (lemon/kiwi fruit), 2,4-D (lemon) and ethiofencarb (kiwi fruit) even though matrix-matched calibration of these analytes was readily achieved. Consequently, data associated with these compounds was considered as qualitative only. Although mean recoveries of aldicarb, butocarboxim and their metabolites were in the range 54-96%, analysis of these compounds was troublesome. Improvement are anticipated following refinements in sample preparation procedures. Table 3 contains example recovery data achieved for 16 pesticides sought in peach.

Table 3. Example recovery and precision data obtained for pesticides sought in peach following multiresidue analysis of fortified organic produce.

of fortified organic produce.							
P es ti cid e	Fortification Levels	% Mean	RSD	n	Ran ge		
	(m gk g ^{·1})	Recovery1					
Azoxystro bi n	0.20 & 0.05	78	9.7	6	70-86		
Bendio carb	0.20 & 0.05	79	8.7	6	70-86		
Carbaryl	0.04 & 0.01	78	7.7	6	68-87		
Dic hlofluanid	0.20 & 0.05	84	9.7	5	60-93		
Fenhexamid	0.20 & 0.05	79	5.9	6	72 - 84		
lma zal il	0.08 & 0.02	77	7.5	6	67-87		
Methiocarb	0.80 & 0.20	71	10.1	5	59-78		
Methiocarb sulfone	0.80 & 0.20	64	17.5	5	55-76		
Methiocarb sulfoxide	0.80 & 0.20	96	43.2	5	41-147		
Myclobutanil	0.20 & 0.05	77	11.4	6	67-86		
Penconazole	0.20 & 0.05	77	12.1	6	66-88		
Propiconazole	0.20 & 0.05	80	9.9	6	71-91		
Pyme troz ine	0.20 & 0.05	73	6.1	6	66-79		
Pyrime tha nil	0.20 & 0.05	84	8.4	6	77-88		
Tebuconazole	0.20 & 0.05	78	11.3	6	68-89		
Thiabendazole	0.20 & 0.05	78	9.5	6	68-86		

Figure 2. Examples of ESIMSMS mass spectra containing precursor → production-ion transitions used for screening and confirmation purposes. (a) thiabendazole, (b) tebuconazole and (c) azoxystrobin.



Retail samples of each commodity were extracted and screened for the presence of residues using the isocratic LCMSMS procedure. Confirmation of residues detected at or above the reporting level following initial screening experiments was achieved using an alternative precursor \rightarrow product-ion transition and the same isocratic method. This was not possible for carbaryl or carbendazim, since alternative transitions were not of sufficient intensity under prevailing conditions, or for 2-phenylphenol. In the case of carbendazim and 2-phenylphenol, alternative HPLC methods were used to achieve confirmation of any residues, whereas confirmation of carbaryl residues was achieved using SIR and the original isocratic method.

Examples of product-ion mass spectra containing transitions used for screening and confirmation experiments are shown in figure 2.

Table 4 contains example LCMS/MS parameters used for confirmation purposes. Typical screening and confirmation results obtained for pesticide residues detected in various samples are detailed in table 5. Maximum Residue Levels (MRL) specified by the (i) CODEX Alimentarius Commission or (ii) in the UK Statutory Instrument for Pesticide Maximum Residue levels in Crops, Food and Feeding Stuffs applicable to the pesticide/commodity combinations are also shown in table 5.

Table 4. LCMS/MS methods used for confirmation of residues detected in samples.

Pesticide	Screen method	Confirmation Method		
Azoxystrobin	m/z 404 → m/z 372	m/z 404 → m/z 344		
Carbaryl	m/z 202 → m/z 145	SIR m/z 202		
Imazalil	m/z 297 → m/z 159	m/z 297 → m/z 69		
Myclobutanil	m/z 289 → m/z 70	m/z 291 → m/z 70		
2-phenylphenol	SIR m/z 169	SIR m/z 169 ^b		
Tebuconazole	m/z 308 → m/z 70	m/z 308 → m/z 125		
Thiabendazole	m/z 202 → m/z 175	m/z 202→ m/z 131		

^a Isocratic acetonitrile:water 50:50 v/v, C₁₈ Elite column 100 x 4.6mm x 5μn ^b Isocratic acetonitrile:water 70:30 v/v, C₁₈ Elite column 100 x 4.6mm x 5μn

Table 5. Example of correlation between screening and confirmation measurements of various pesticide residues detected in samples.

Analyte	Fortification (mgkg ⁻¹)	Commodity (MRL mgkg ⁻¹)	Residue Level ^a (mgkg ⁻¹)		% Recovery ^b	
	((·	Screen	Confirmation	Screen	Confirmation
Azoxystrobin	0.20	Grape (2.0)	0.20	0.20	72	74
Carbaryl	0.04	Kiwi Fruit (10.0)	0.10	0.10	84	68
Imazalil	0.20	Lemon (5.0)	1.40	1.50	78	83
Myclobutanil	0.20	Strawberry (1.0)	0.13	0.13	73	74
2-phenyl phenol	0.40	Lemon (10.0)	2.10	2.10	88	78
Tebuconazole	0.20	Peach (1.0)	0.13	0.13	81	86
Thiabendazole	0.20	Lemon (5.0)	2.00	2.20	73	83

a Not corrected for recover

Once the utility of isocratic LCMSMS in this application area had been established, it was then possible to assess the impact of this experimental approach upon the efficiency of analytical procedures carried out in our laboratory. This was achieved by comparing overall analysis times of the isocratic LCMSMS method with gradient LCMSMS methods used in our laboratory. It could take up to 20 minutes for gradient equilibration between each run, which compromised the benefits achieved from the direct analysis of crude extract. This was eliminated by the use of isocratic separation. In addition, the frequency of adaptation of gradient methods or the need for development of 'customised' gradients was significantly reduced with regard to the pesticides involved in this study.

CONCLUSIONS:

- A method has been developed that combines isocratic HPLC separation and tandem mass spectrometry for the quantitative and qualitative determination of pesticide multi-residues in crude extracts of a variety of fruit and vegetables.
- This experimental approach has provided significant efficiency gains of at least 25% against gradient LCMSMS methods used previously.

b Single spike corresponding to 4 times the reporting level included in analysis batch