An Optimal Method for the Analysis of Pesticides in a Variety of Matrices

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Introduction

The global agriculture industry uses over a thousand pesticides for the production of food and foodstuffs. Producers require pesticides to meet the increasing demand for reasonably priced food both in and out of season. This growing demand has increased the use of pesticides and expanded poor agricultural practices elevating risks in the food supply and environment. Analytical laboratories are then strained to evaluate and quantitate hundreds of pesticides in a wide range of matrices. Not only are laboratories faced with time constraints, but they also face matrix interferences that degrade their ability to accurately identify and quantitate the multitude of target pesticides.

The MassHunter Pesticide & Environmental Pollutant MRM Database (Rev. A.01.01) is the most comprehensive GC MRM Database on the market. With over 1000 compounds with at least 8 MRMs/compound, analysts have the ability to optimize their acquisition methods for their target compounds in a variety of matrices. The availability of multiple MRM transitions not only helps to address matrix interferences, but it also aids in accurately identifying compounds that may have several MRMs in common.

Matrix interferences have been a common complaint for MRM acquisitions in pesticides analysis. It has been seen that the usefulness of a given compound's MRMs can change depending on the matrix being measured, due to factors such as increased/decreased response (which changes the quant and qual ions). The ability to have multiple MRMs from which to choose aids in lab productivity, improved quant method generation, and achieving optimal analysis.

Experimental

Methodoloav

Table 1. 7890B GC Method Conditions

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The analysis was conducted on an Agilent 7890B GC and 7010 Series Triple Quadrupole GC/MS system. See Tables 1 – 3 for method parameters. The system was configured with a Multimode Inlet equipped with an ultra-inert liner (p/n: 5190-2293). The inlet was then connected to two HP-5ms UI columns (15 m \times 0.25 mm \times 0.25 µm; p/n: 19091S-431 UI) coupled to each other through a purged ultimate union (PUU) for the use of backflushing (see Figure 1).

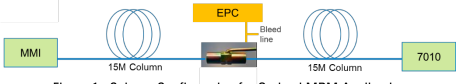


Figure 1. Column Configuration for Optimal MRM Application.					
	Figure 1	Column Con	figuration for	Optimal MRM Application	

4-mm Ultra Inert liner with wool

Table 3. 7010 MS/MS P	arameters
Electron Energy	70 eV
Tune	atunes.eihs.tune.xml
EM gain	10
MS1 & MS2 resolution	Wide
Collision Cell	1.5 mL/min N $_2$ & 2.25 mL/min He
Quant/Qual transitions	Matrix Optimized
Dwell times	Time Segment (TS) specific*
Source temperature	300 °C
Quad temperatures	150 °C

scan rate of ~5 scans/sec for the TS

Sample Prep

A selection of matrices were chosen over a variety of categories. Table 4 provides the chosen matrix per category and a quick look at the QuEChERS sample prep procedure that was followed.

injection mode	110t-splitless							
Injection volume	1 µL			Table 4, Matri	x Selection and Sample Pre	paration Used for Optimal MRM Application		
Inlet temperature	280 °C							
Carrier gas	He, constant	t flow 1.00 mL/	min (column 2 = 1.20 mL/min)	Category	Matrix	Sample Prep		
		60 °C	1 min	High Oil	Extra Virgin Olive Oil	3 g oil/7 mL water, EN salts (5982-5650), EMR-L (5982-1010), Polish Pouch (5982-0102), Dry step		
Oven program	40 °C/min	120 °C	0 min			3 g tea/7 mL water, EN salts, EN dSPE pigment		
	5 °C/min	310 °C	0 min	Difficult	Black loose Leaf Tea	(5982-5256)		
MS transfer line temperatur	nsfer line temperature 280 °C		High Pigment	Fresh Leaf Baby Spinach	10 g, EN salts, EN dSPE pigment (5982-5356)			
Table 2. PUU Backflush Settings*		High Starch	Jasmine Rice	3 g rice/7 mL water, EN salts, EN dSPE Fatty (5982-5156)				
Timing	Timing 1.5 min duration during post-run		High Water	Basic Cucumber	10 g, EN salts, EN dSPE General (5982-5056)			
Oven temperat	Oven temperature 310 °C		310 °C		Organic Honey	5 g honey/5 mL water, EN salts, EN dSPE		
Aux EPC press	Aux EPC pressure ~50 psi		High Sugar	organic noncy	General (5982-5056)			
Inlet pressure		~2 psi		High Acid	Navel Orange	10 g, EN salts, EN dSPE Fatty (5982-5156)		
			atory. A 1.5 min backflush nade for a 5 min backflush	Clean 15	Yellow Onion (not sweet)	10 g, EN salts, EN dSPE Fatty (5982-5156)		

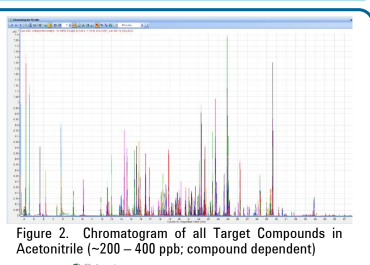
Matrix Optimized MRMs

Identification of Matrix Optimized MRM Transitions

Agilent Technologies offers the most comprehensive GC MRM Database for Pesticides and Environmental Pollutants (PN: G9250-60006). The MRM Database contains 1000+ compounds and up to 10 MRMs/compound. The all-inclusive database provides a surplus of MRMs to aid in accurate identification, utilize MRMs that fall within the ion ratio confidence limits, and avoid matrix interferences.

Across the globe there are a multitude of different applications and regulations that are followed. The P&EP MRM Database provides all of the material for users to identify the optimal MRMs for their specific analysis. In order to provide guidance on the optimal use of these MRMs, Agilent has begun to look at target compounds in a variety of matrices.

A total of 195 compounds were selected for the analysis. Each compound was analyzed in each of the 8 matrices and in acetonitrile (ACN). The top 5 MRM transitions for each target compound were selected based on response, ion ratio, and selectivity. From these, the top 3/4 MRMs were transferred to a matrix specific method for further optimal analysis.



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	A 8	C	D	E
1	Compound Name	CAS #	Target	My Target Compound List
2	1 Phenol	108-95-2	Target	
3	2 Dimefox	115-26-4	Target	Create New Target List
4	3 Dichlorobenzene, 1,2-	95-50-1	Target	
5	4 DBCP (Dibromo-3-chloropropane, 1,2-)	96-12-8	Target	Cours Courses Transmittee
6	5 Ethiolate	2941-55-1	Target	Save Current Target List
7	6 Methamidophos	10265-92-6	Target	
8	7 Dichlorvos	62-73-7	Target	Manage Target Lists
9	8 Trichlorfon	52-68-6	Target	
10	9 Disulfoton-sulfoxide	2497-07-6	Target	
11	10 Phthalide	87-41-2	Target	Add Compounds
12	11 EPTC	759-94-4	Target	
13	12 Mevinphos, Z-	338-45-4	Target	Remove Compounds
14	13 Mevinphos, E-	7786-34-7	Target	Remove compounds
15	14 Butylate	2008-41-5	Target	
16	15 Acephate	30560-19-1	Target	Import CAS Numbers
17	16 Acenaphthene-d10	15067-26-2	Target	
18	17 Heptenophos	23560-59-0	Target	
13	18 Omethoate	1113-02-6	Target	Build MRM Table
20	19 Thionazin	297-97-2	Target	
21	20 Propoxur	114-26-1	Target	Home
22	21 Demeton-S-methyl	919-86-8	Target	

Please note that due to the large amount of data collected, not all observations will be detailed in this poster. Contact the authors for further information

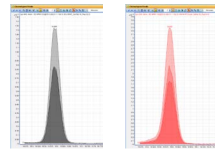
Changes in Quant (QQ) and Qual lons (Q1, Q2, ...)

The majority of pesticides analyzed indicated that the responses of the optimal MRM transitions often change.

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Solvent MRMs			Solvent MRMs Cucumber MRMs				
m/z	m/z (prod.)	CE	m/z	m∕z (prod.)	CE		
148	75.1	25	146	75.1	25		
111	75.1	10	146	111.1	15		
146	75.1	25	111	75.1	10		

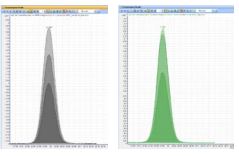
2. Dichlorohenzene in ACN 12. Dichlorohenzene in Cucumber



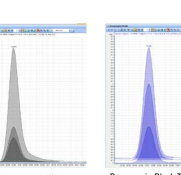
	Aldrin in ACN		Aldrin	in Yellow Onion	
Solvent MRMs			Yello	w Onion MRI	Ms
m/z	m/z (prod.)	CE	m/z	m/z (prod.)	CE
262.9	192.9	35	262.9	192.9	35
254.9	220	20	262.9	190.9	35
262.9	190.9	35	254.9	220	20

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	The start is the start of the start is the s
EPTC in ACN	EPTC in Organic Honey
LITCHIAGN	Li io in organic noney

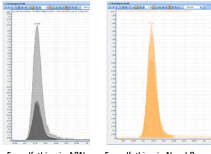
Solvent MRMs			Organ	ic Honey MR	Ms
m/z	m/z (prod.)	CE	m/z	m/z (prod.)	CE
128	86	5	128	86	5
132	90	5	132	90	5
132	62	10	189.1	128	5



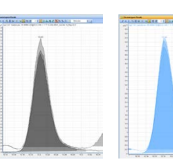
tra	ns-Chlordane in ACN	trans-Ch	lordane in Spinach		
Solvent MRMs			Sp	inach MRMs	;
m/z	m/z (prod.)	CE	m/z	m/z (prod.)	CE
271.8	236.9	15	372.8	265.9	25
372.8	265.9	25	271.7	236.9	15
374.8	265.8	15	374.8	265.8	15



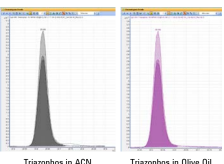
	Propoxur in ACN		Prop	oxur in Black Tea	
Solvent MRMs			Bla	ck Tea MRM	S
m/z	m/z (prod.)	CE	m/z	m/z (prod.)	CE
110	63	25	110	63	25
152	110	10	110	64	15
110	92	10	152	110	10



Fensulfothion in ACN Fensulfothion in Navel Orange						
Solvent MRMs			Orange MR	Ms		
m/z (prod.)	CE	m/z	m/z (prod.)	CE		
96.8	20	140	125	10		
125	10	156	141	10		
141	10	291.8	156	15		
	Ivent MRMs m/z (prod.) 96.8 125	Ivent MRMs m/z (prod.) CE 96.8 20 125 10	Ivent MRMs Navel m/z (prod.) CE m/z 96.8 20 1400 125 10 1560	Ivent MRMs Navel Orange MR m/z (prod.) CE m/z m/z (prod.) 96.8 20 140 125 125 10 156 141		



	Carbofuran in ACN		Carbofuran in Jasmine Rice						
So	lvent MRMs		Jasmine Rice MI						
m/z	m/z (prod.)	CE	m/z	m/z (prod.)					
149.1	77.1	30	151	136.1					
164.2	149.1	10	150	134					
164.2	103.1	25	136	77					



	mazophos in Acr	Thazophos in Olive Oli							
So	lvent MRMs	Olive Oil MRMs							
m/z	m/z (prod.)	CE	m/z	m/z (prod.)	(
161.2	134.2	5	161.2	134.2					
161.2	106.1	10	161.2	106.1	1				
257	162.1	5	161.2	91	1				

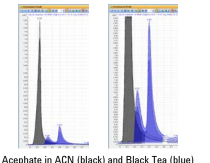




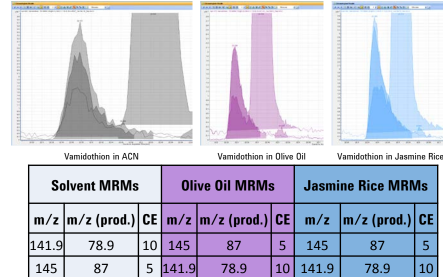
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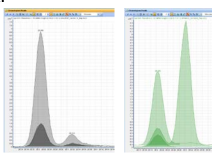
Matrix Interferences Are Real

A compound's MRMs were seen to be interfered with by the matrix.



So	lvent MRMs	Black Tea MRMs						
m/z	m/z (prod.)	CE	m/z	m/z (prod.)	CE			
136	94	15	142	96	5			
78.9	47	10	78.9	47	10			
142	96	5	124.9	47	15			





Phonothrin II in ACN

	Filehouinin II III Auf	N								
So	lvent MRMs		Spinach MRMs							
m/z	m/z (prod.)	CE	m/z	m/z (prod.)	CE					
122.9	81.1	5	122.9	81.1	5					
182.9	168.1	10	122.9	79.1	20					
182.9	153.1	15	182.9	168.1	10					

Results

5 108.9 78.9 5 108.9 78.9

108.9 78.9

The 7010 Series Triple Quadrupole GC/MS system can confirm pesticide residues at the low ppb level even in the most complex extracts. The calibration standards were prepared at concentrations ranging from 0.12 pg/ μ L to 50 pg/ μ L; for 90% of compounds a calibration curve with a R² \geq 0.990 was produced. All analyzed pesticides obtained a %RSD of repeated measurements of \leq 30%, and 90% have a LOQ \leq 1.5 pg/µL. A representative selection of compounds and their calculated values are shown for Organic Honey and Baby Spinach compared to ACN solvent.

	A representative selection of compounds and their calculated values are shown for organic noney and baby opinach compared to Acit solvent.														
	SOLVENT					ORGANIC HONEY					SPINACH				
CMPD	%RSD	IDL _{RSD} (pg)	MDL (pg∕µL)	iLOQ (pg∕µL)	%Error	%RSD	IDL _{RSD} (pg)	MDL (pg∕µL)	iLOQ (pg∕µL)	%Error	%RSD	IDL _{RSD} (pg)	MDL (pg∕µL)	iLOQ (pg∕µL)	%Error
Ethoprophos	11.13	0.386	0.408	1.477	5.72	8.72	0.303	0.293	1.061	3.11	8.25	0.286	0.295	1.068	3.18
BHC-alpha	9.38	0.325	0.343	1.241	5.51	7.83	0.271	0.261	0.943	4.01	7.94	0.275	0.276	0.997	0.16
Dazomet	11.15	0.386	0.412	1.492	6.81	4.38	0.152	0.152	0.552	0.45	9.10	0.315	0.322	1.165	2.22
BHC-beta	9.27	0.321	0.340	1.229	5.80	17.19	0.596	0.541	1.959	9.10	8.76	0.304	0.303	1.096	0.24
Aminocarb	19.89	0.690	0.737	2.665	6.75	8.40	0.291	0.285	1.032	2.09	9.76	0.339	0.362	1.310	6.91
Phenanthrene-D10	7.68	0.266	0.280	1.014	5.19	6.59	0.229	0.217	0.786	4.92	8.95	0.311	0.312	1.129	0.49
Diazinon	9.63	0.333	0.352	1.274	5.69	7.33	0.254	0.238	0.862	6.15	5.78	0.200	0.201	0.728	0.46
2,4-D butyl ester	15.67	0.541	0.576	2.083	6.35	8.09	0.280	0.260	0.940	7.08	19.66	0.679	0.687	2.485	1.13
Chlorpyrifos-methyl	9.96	0.345	0.364	1.316	5.37	7.76	0.269	0.252	0.910	6.42	7.04	0.244	0.244	0.884	0.20
Triadimefon	12.71	0.440	0.464	1.680	5.41	4.26	0.148	0.137	0.497	6.97	10.17	0.352	0.363	1.313	2.95
Heptachlor endo-epoxide	9.57	0.662	0.698	2.526	5.41	7.75	0.536	0.493	1.783	8.13	7.17	0.496	0.490	1.774	1.25
Flurenol-butyl	9.09	0.311	0.328	1.185	5.47	6.85	0.234	0.219	0.793	6.32	18.80	0.642	0.650	2.350	1.13
Chlordane-cis	8.35	0.290	0.305	1.103	5.26	13.08	0.453	0.411	1.486	9.42	21.67	0.751	0.750	2.713	0.14
DDT-o,p'	4.42	0.153	0.162	0.585	5.31	8.78	0.305	0.270	0.977	11.35	23.04	0.799	0.785	2.839	1.84
Hexazinone	11.71	0.407	0.431	1.558	5.67	4.91	0.171	0.157	0.569	7.85	7.40	0.257	0.263	0.951	2.10
Azinphos-ethyl	9.01	0.312	0.329	1.191	5.45	13.77	0.477	0.436	1.579	8.53	16.08	0.557	0.564	2.042	1.25
Permethrin, (1R)-trans-	10.89	0.377	0.396	1.431	5.05	10.25	0.354	0.335	1.211	5.57	22.36	0.773	0.793	2.870	2.56

Conclusions

Matrix interferences are a common complaint for MRM acquisitions in pesticides analysis and can alter a target compound's

- Changes in Quant (Q0) and Qualifier ion (Q1, Q2, ...) responses are the most common. These changes merely affect the relative abundances of the MRMs which plays a part in method development for optimum quantitative data analysis.
- The availability of multiple MRMs per compound allows a user to discriminate among compounds with similar transitions, and to select MRMs that fulfill desired ion ratio confidence limits.
- The main challenges come from extremely large matrix interferences, which are encountered more often in complex matrices such as Loose Leaf Black Tea or Spinach. The number of usable MRMs for a given target compound can be reduced, and the shift in RT can push a target out of a Time Segment. In these cases great care must be exercised to produce accurate results for all analytes.

Matrix Optimized MRM Transitions aid in lab productivity, improved quant method generation, and optimal analysis!

Agilent's MassHunter P&EP MRM Database (Rev. A.01.01) is the most comprehensive GC MRM Database on the market. With the evolving market and demand for matrix optimized transitions, Agilent will add to the value of their MRM Database with the addition of 7800 matrix optimized transitions to help provide customers with their optimal pesticides analysis!

