

Analysis of Pesticides in Kale Purchased from Local Grocers Using Triple-Quadrupole GC-MS/MS Analysis

1 Introduction

The introduction of pesticides to improve agricultural productivity has led to a marked change in how crops are grown. However, certain pesticides have been linked to a number of health issues. As such, regulatory agencies all over the world have placed restrictions on the types and levels of pesticides that can be used for agriculture. Accordingly, food samples need to be tested for pesticide content to meet regulatory requirements.

Triple-quadrupole gas chromatograph tandem mass spectrometry (GC-MS/MS) is widely used for pesticide analysis due to the sensitivity and specificity of the technique. The specificity offered by GC-MS/MS is excellent for analysis of samples with complex matrices. One such food example is kale. Kale frequently makes the Environmental Working Group's (EWG) "dirty dozen," a list of fruits and vegetables ranked according to high pesticide content[1]. This makes kale a good target for testing the capabilities of the JMS-TQ4000GC triple-quadrupole GC-MS/MS.

In this app note, we demonstrate the ability of the JMS-TQ4000GC to measure the pesticide content of kale samples purchased from local grocers. The sensitivity and specificity offered by selected reaction monitoring (SRM) will be used against a complex kale extract matrix, including matrix-matched standard samples.

2 Experimental

Organic and non-organic spinach samples were purchased from local grocers. Prior to extraction, unwashed leaves were homogenized using a food processor. Approximately 15 grams of homogenized sample was weighed out into a 50 mL plastic tube, followed the addition of 15 mL of acetonitrile (HPLC grade, Fisher Scientific). Samples were spiked with 3.75 μ L of 100 ppm chlorpyrifos-d10 (Cambridge Isotope Laboratories, PN# DLM-4360-1.2) as an internal standard to monitor recovery and then extracted using Restek Q-sep QuEChERS extraction salts (PN# 25851) according to the AOAC 2007.01 method[2]. Samples were then centrifuged for 10 minutes. One mL of the resulting supernatant was put through a dSPE cleanup step using Restek Q-sep QuEChERS dSPE Tubes (AOAC 2007.01 method, PN# 26125) by following the provided dSPE instruc-



The JEOL JMS-TQ4000GC.

tions. The dSPE supernatant was either used to create matrix-matched standards, or injected directly into the GC as a sample. For standard samples, the extracted matrix was spiked with stock pesticide standards (Restek PN# 34124 - 34129) to concentrations ranging from 0.1 to 100 ppb.

Samples were analyzed on the JMS-TQ4000GC using the parameters in [Table 1](#). Transitions and collision energies used for each compound are listed in supplementary [Table S1](#). Optimal product- and precursor-ion pairs and optimized collision energies for each pesticide were determined using built-in SRM optimization tools. Each standard and sample were run in triplicate with the exception of the one ppb standard samples, for which ten replicates were done to calculate the instrument detection limit (IDL) where possible.

3 Results and Discussion

Standard performance data is shown in [Figure 1](#), [Figure 2](#), and [Table 2](#). [Figure 1](#) shows SRM chromatograms for select pesticides at 5 ppb. Of the 54 pesticides measured, 51 were detected at 5 ppb or less, and instrument detection limits (IDL) of 45 pesticides were less than 1 ppb ([Table 2](#)). Calibration curves for selected pesticides are shown in [Figure 2](#). Good linearity ($R^2 > 0.99$) was observed for every pesticide within its dynamic range except for azoxystrobin ([Table 2](#)). Chlorinated pesticides, such as cypermethrin and chlordane, which are difficult to mea-

Table 1: GC-MS parameters.

	Agilent 7890B GC	JMS-TQ4000GC
Column	ZB5-MSplus 30.0 m, 0.25 mm i.d., 0.20 μm (Phenomenex, Torrence, CA)	Ion Source Temp. 250 °C Interface Temp. 300 °C Ionization Mode EI+, 70 eV, 100 μA Measurement Mode SRM, High Target Cycle Time \approx 330 ms Channel Time 20 - 100 ms Relative EM Voltage 900 V Collision Gas N ₂ , 10 %
Inlet Liner	4 mm single taper w/wool on bottom (Phenomenex, Torrence, CA)	
Inlet Temp.	250 °C	
Carrier Gas Type, Flow	He, 1.000 mL/min	
Mode	Pulsed Splitless	Oven Program
Pulsed Press., Time	206.84 kP, 0.55 min	80 °C (0.75 min) → 35 °C/min → 190 °C → 5 °C/min → 240 °C → 20 °C/min → 330 °C (4 min)
Purge Flow	30 mL/min, 1.0 min	
Septum Purge Flow	3.0 mL/min	
Injection Volume	2.0 μL	

sure using liquid chromatography, were easily measured using the TQ4000GC.

Data for spinach samples purchased from local grocers are shown in [Figure 3](#) and [Table 3](#). Overall, very few pesticides were observed, even in the non-organic samples. This was surprising, because kale is usually high on the EWG's "dirty dozen" list. Most pesticides found were unquantifiable due to their extremely low concentration. Figure 3 shows the quantitative and qualifier ions for permethrin measured in Brand B non-organic kale. Even though the concentration of permethrin was quite low (< 1 ppb, [Table 3](#)), its presence was still detectable due to the sensitivity of the TQ4000GC and SRM analysis.

4 Conclusions

The performance of the JMS-TQ4000GC triple quadrupole mass spectrometer was tested for measuring pesticides in extracted kale samples. Of the 54 pesticides measured, 51 were observed at 5 ppb or less in matrix-matched standard samples, and 45 had IDLs < 1 ppb. Calculated linearity and IDLs also showed good performance. Chlorinated pesticides, which can be difficult to measure using other chromatographic methods, were easily measured using GC-MS.

Additionally, kale samples were purchased from local grocers and analyzed using methods mentioned above. A small variety of pesticides were observed, however, all concentrations were less than respective action limits set by the EPA. Because of the low overall concentrations of pesticides detected on the samples, no conclusion about organic/non-organic pesticide comparison could be reached. Although at very low concentration (< 1 ppb), permethrin was still detectable. Due to its sensitivity and selectivity, the JMS-TQ4000GC could be a valuable addition to any food lab conducting pesticide testing.

References

- [1] *Dirty Dozen™ Fruits and Vegetables with the Most Pesticides*. URL: <https://www.ewg.org/foodnews/dirty-dozen.php>.
- [2] AOAC Official Methods of Analysis. *Pesticide Residues in Foods by Acetonitrile Extraction and Partitioning with Magnesium Sulfate*. 2007.01.

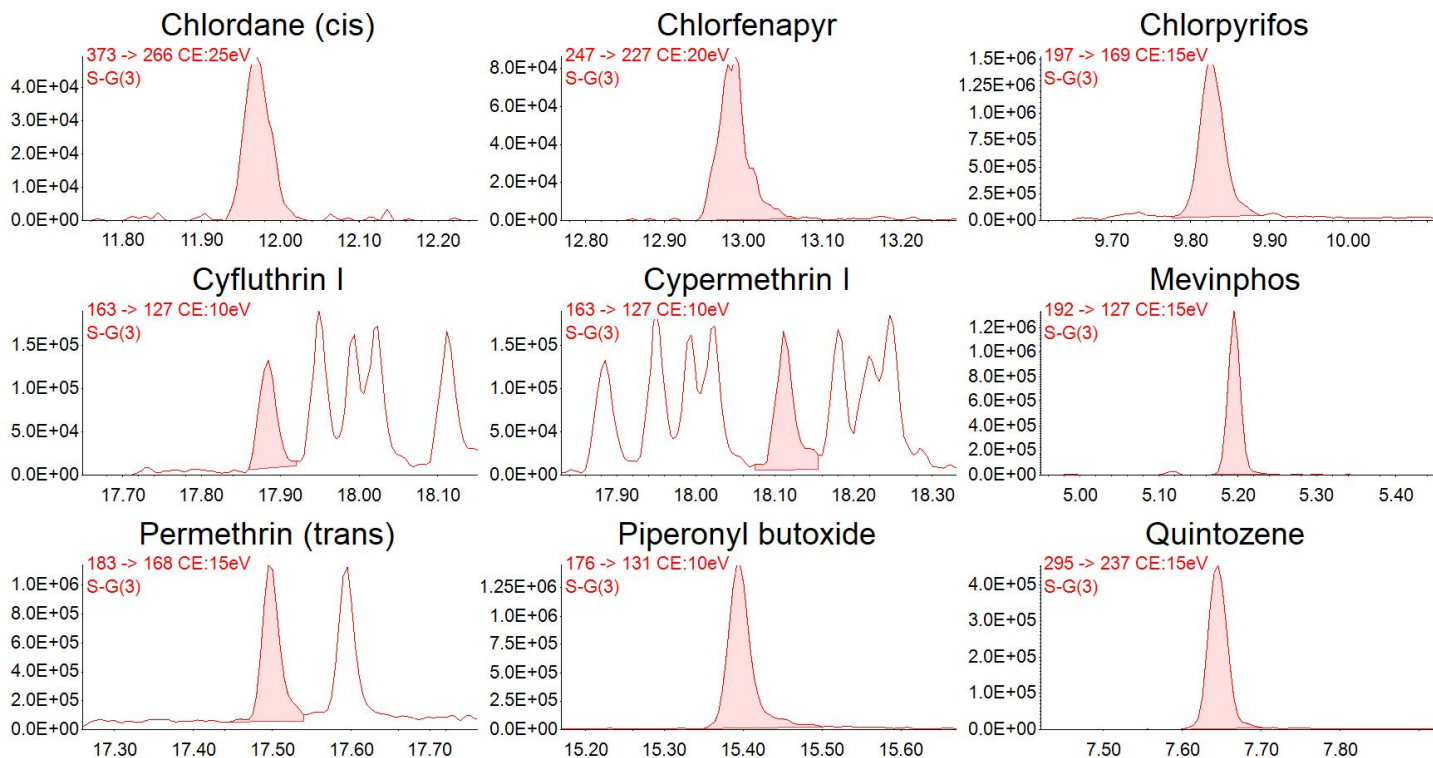


Figure 1: Selected SRM Chromatograms of a 1 ppb matrix-matched standard sample.

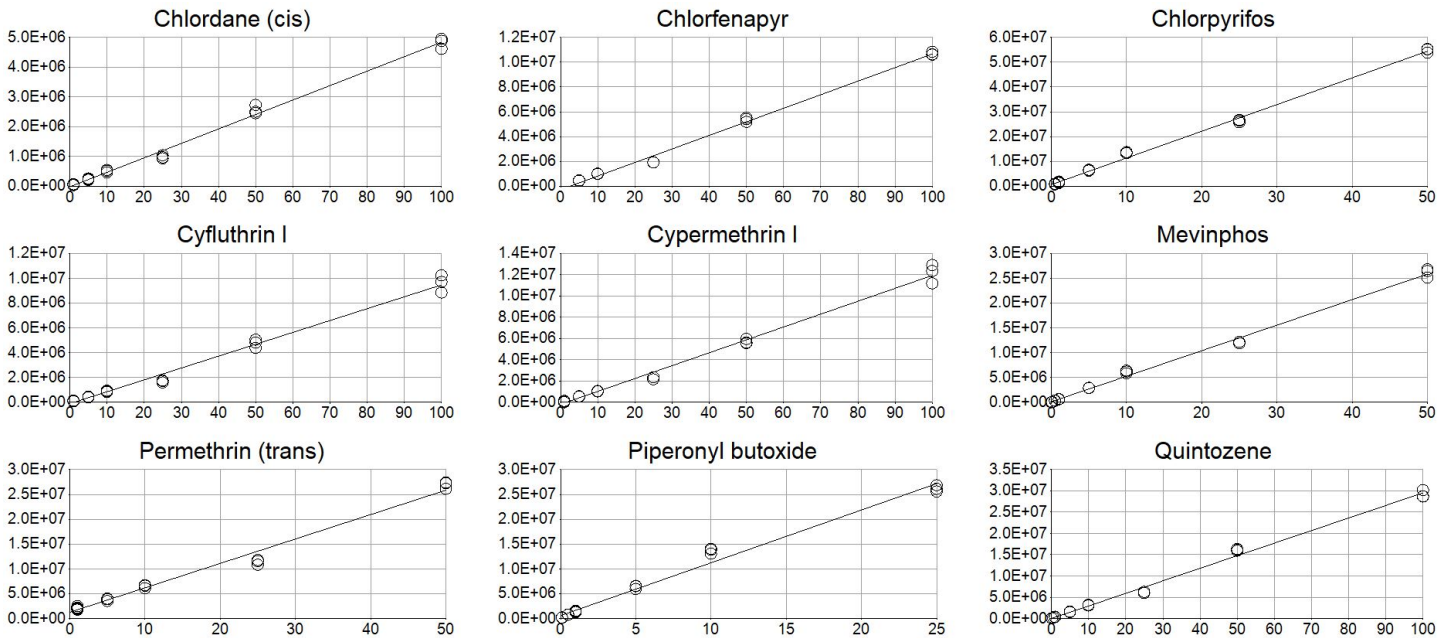


Figure 2: Selected calibrations curves for matrix-matched standard samples.

Table 2: Range, linearity, instrument detection limit (IDL), and CV of pesticide standards measured in spinach matrix.

Compound	Range (ppb)	Linearity (R^2)	IDL (ppb)	CV (%)
Acephate	25 - 100	0.9912	N/A	N/A
Aldicarb (decomp)	0.5 - 100	0.9984	0.32	11.3
Azoxystrobin	1 - 100	0.9898	0.69	24.5
Bifenazate	0.5 - 100	0.9931	0.48	16.8
Bifenthrin	0.1 - 100	0.9978	0.18	6.3
Boscalid	0.5 - 100	0.9966	0.55	19.4
Carbaril	1-100	0.9946	0.32	11.3
Carbofuran	1-100	0.9944	0.44	15.8
Chlorantraniliprole	5-100	0.9928	N/A	N/A
Chlordane	1-100	0.9974	0.52	18.5
Chlorfenapyr	1-100	0.9975	0.40	14.0
Chlorpyrifos	0.5 - 100	0.9984	0.23	8.1
Chlorpyrifos-d10	0.5 - 100	0.9983	0.25	8.7
Cinerin	5-100	0.9955	N/A	N/A
Clofentezine	0.5-100	0.9989	0.36	12.9
Coumaphos	1-100	0.9900	0.75	26.4
Cyfluthrin	1-100	0.9952	0.46	16.2
Cypermethrin	1-100	0.9943	0.54	19.3
Diazinone	0.5-100	0.9986	0.11	4.0
Dichlorvos	0.5-100	0.9933	0.19	6.6
Dimethoate	1-100	0.9967	0.37	13.1
Dimethomorph	1-100	0.9941	0.48	17.0
Ethoprophos	0.5-100	0.9983	0.22	7.7
Etofenprox	0.5-100	0.9990	0.22	7.6
Etoxazole	0.5-100	0.9981	0.31	11.0
Fenoxy carb	1-100	0.9960	0.44	15.8
Fipronil	0.5-100	0.9958	0.43	15.2
Fludioxonil	1-100	0.9940	0.58	20.5
Imazalil	1-100	0.9924	0.46	16.3
Jasmolin	10-100	0.9957	N/A	N/A
Kresoxim-methyl	0.5-100	0.9984	0.26	9.3
Malathion	0.5-100	0.9980	0.19	6.6
Metalaxyll	1-100	0.9974	0.31	11.0
Methiocarb	1-100	0.9967	0.35	12.3
Methomyl	1-100	0.9983	0.31	10.8
Methyl Parathion	5-100	0.9983	N/A	N/A
Mevinphos	0.5-100	0.9989	0.14	4.9
Myclobutanil	0.5-100	0.9965	0.25	9.0
Oxamyl	25-100	0.9922	N/A	N/A
Paclobutrazol	0.5-100	0.9941	0.36	12.7
Permethrin	1-100	0.9948	0.42	14.7
Phosmet	1-100	0.9943	0.97	34.4
Piperonyl butoxide	0.5-100	0.9965	0.22	7.7
Prallethrin	1-100	0.9960	0.23	8.3
Propiconazole	1-100	0.9969	0.45	15.8
Propoxur	0.5-100	0.9980	0.33	11.8
Pyrethrin	50-100	0.9959	N/A	N/A
Pyridaben	0.5-100	0.9987	0.36	12.8
Quintozene	1-100	0.9976	0.20	7.0
Spiromesifen	5-100	0.9922	N/A	N/A
Spiroxamine	0.5-100	0.9974	0.20	7.0
Tebuconazole	0.5-100	0.9963	0.25	9.0
Thiamethoxam	5-100	0.9933	N/A	N/A
Trifloxystrobin	1-100	0.9960	0.26	9.1

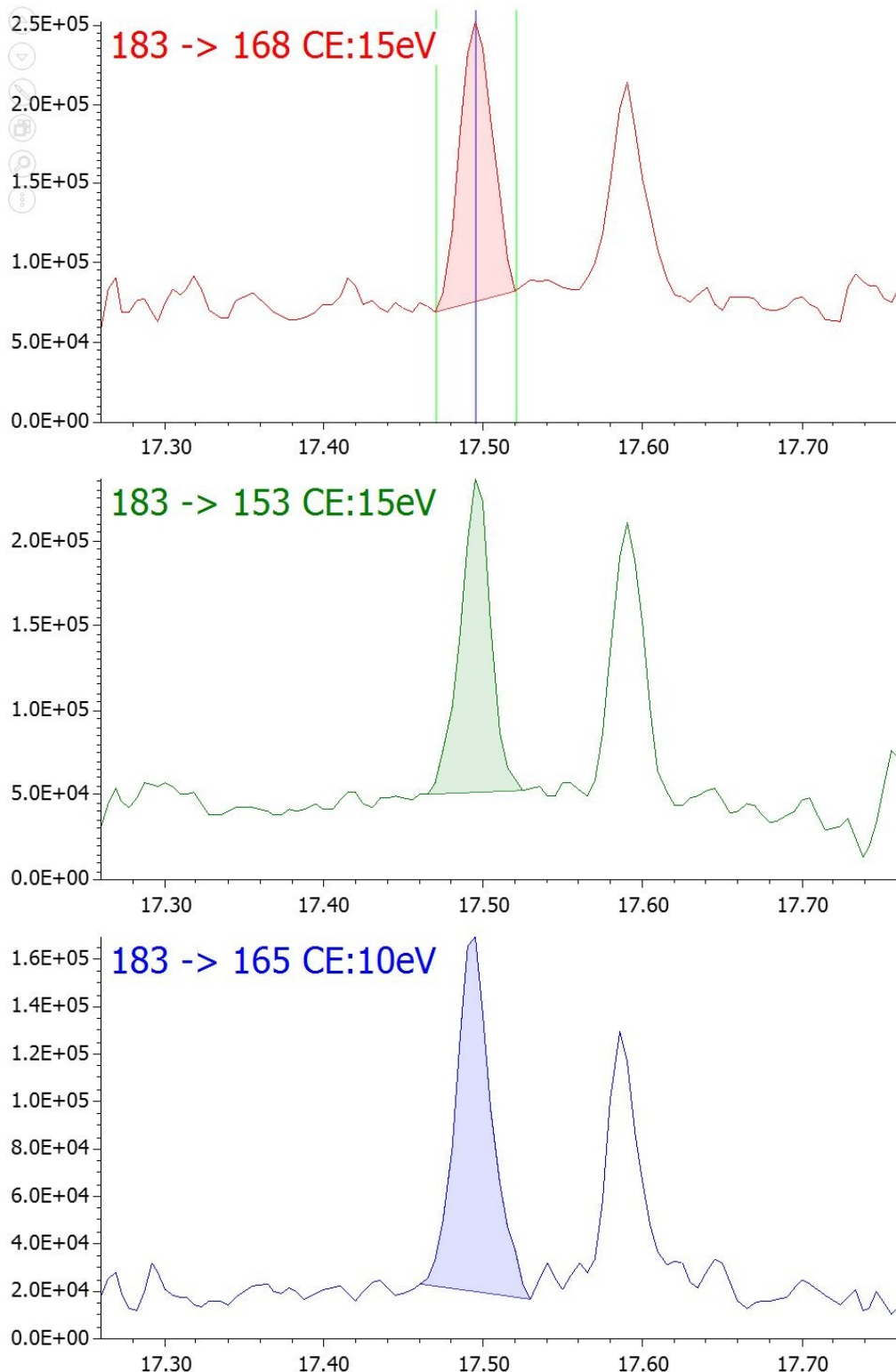


Figure 3: SRM Chromatogram of trans-permethrin found in Brand B organic-labeled spinach. The top chromatogram is the quantitative ion; qualifier ions are shown in the middle and bottom chromatograms.

Table 3: Pesticides detected (ppb) and sample recoveries (%) for spinach samples.

Compound	Regular		Organic	
	Brand A	Brand B	Brand C	Brand D
Azoxystrobin	< 0.1	82.8		
Bifenthrin	1.17	< 0.1		
Clofentazine	< 0.1	< 0.1	< 0.5	< 0.1
Cyfluthrin	7.05			
Dimethoate	< 0.1	1.93	1.10	1.50
Dimethomorph		4.00	2.50	
Etofenprox		t	t	
Myclobutanil		0.473	0.423	
Paclobutrazole		< 0.5	< 0.5	t
Permethrin	~1	< 0.1	< 0.1	t
Tebuconazole	< 0.1	0.277	< 0.1	
Thiamethoxam	2.64			
Recoveries	94.3	83.3	86.5	69.9

Table S1: SRM transitions and collision energies for each pesticide.

	Quantitative Ion		Qualifier Ion 1		Qualifier Ion 2	
	Transition	CE (eV)	Transition	CE (eV)	Transition	CE (eV)
Acephate	136 → 94	15	125 → 79	10	125 → 47	20
Aldicarb (decomposed)	115 → 68	5	115 → 100	5	115 → 69	5
Azoxystrobin	344 → 156	30	388 → 345	20	388 → 300	20
Bifenazate	258 → 196	15	300 → 196	20	300 → 258	10
Bifenthrin	181 → 165	30	181 → 166	10	181 → 164	40
Boscalid	140 → 112	10	140 → 76	25	142 → 114	15
Captan	149 → 105	5	149 → 79	15	149 → 77	25
Carbaril	144 → 115	20	144 → 116	15	115 → 89	20
Carbofuran	164 → 103	30	164 → 149	10	149 → 77	25
Chlorantraniliprole	384 → 357	20	386 → 359	15	112 → 76	15
Chlordane (cis)	373 → 266	25	375 → 266	25	371 → 264	30
Chlordane (trans)	373 → 266	25	375 → 266	25	371 → 264	30
Chlорfenapyr	247 → 227	20	247 → 200	30	328 → 247	25
Chlorpyrifos	197 → 169	15	199 → 171	20	314 → 258	15
Chlorpyrifos-d10	200 → 172	20	260 → 167	25	260 → 139	30
Cinerin I	123 → 81	10	107 → 91	10	123 → 79	20
Cinerin II	107 → 91	10	121 → 93	5	167 → 107	5
Clofentezine	137 → 102	10	137 → 75	25	139 → 102	15
Coumaphos	226 → 163	15	226 → 198	10	362 → 226	15
Cyfluthrin I	163 → 127	10	163 → 91	15	165 → 91	15
Cyfluthrin II	163 → 127	10	163 → 91	15	165 → 91	15
Cyfluthrin III	163 → 127	10	163 → 91	15	165 → 91	15
Cyfluthrin IV	163 → 127	5	163 → 91	15	165 → 91	15
Cypermethrin I	163 → 127	10	163 → 91	15	165 → 91	15
Cypermethrin II	163 → 127	10	163 → 91	15	165 → 91	15
Cypermethrin III	163 → 127	10	163 → 91	15	165 → 91	15
Cypermethrin IV	163 → 127	10	163 → 91	15	165 → 91	15
Diazinone	304 → 179	15	179 → 121	35	179 → 137	20
Dichlorvos	109 → 79	5	220 → 185	5	145 → 109	10
Dimethoate	93 → 63	10	125 → 79	10	87 → 42	10
Dimethomorph I	301 → 165	10	303 → 165	10	301 → 273	10

Table S1: (continued)

	Quantitative Ion		Qualifier Ion 1		Qualifier Ion 2	
	Transition	CE (eV)	Transition	CE (eV)	Transition	CE (eV)
Dimethomorph II	301 → 165	10	303 → 165	10	301 → 273	10
Ethoprophos	158 → 97	15	158 → 81	15	158 → 114	10
Etofenprox	163 → 107	25	163 → 135	10	163 → 77	35
Etoxazole	141 → 113	15	300 → 270	25	204 → 176	10
Fenoxy carb	116 → 88	5	255 → 186	15	186 → 157	15
Fipronil	367 → 213	30	351 → 255	15	369 → 215	35
Fludioxonil	248 → 182	15	248 → 154	15	154 → 127	10
Imazalil	215 → 173	10	173 → 145	20	173 → 109	30
Jasmolin I	123 → 81	10	123 → 79	20	164 → 109	10
Jasmolin II	107 → 91	10	121 → 77	20	121 → 93	10
Kresoxim-methyl	116 → 89	15	131 → 89	25	116 → 63	25
Malathion	127 → 99	10	173 → 99	10	173 → 117	10
Metalaxy l	160 → 130	20	160 → 144	20	160 → 145	10
Methiocarb	168 → 109	15	168 → 153	10	153 → 109	10
Methomyl	105 → 88	5	105 → 58	10	88 → 44	5
Methyl Parathion	263 → 109	15	109 → 79	10	125 → 79	10
Mevinphos	192 → 127	15	127 → 109	10	127 → 79	20
Myclobutanil	179 → 125	15	150 → 123	20	181 → 127	15
Naled	145 → 109	10	109 → 79	5	147 → 109	10
Oxamyl	162 → 115	10	162 → 145	5	145 → 111	10
Paclobutrazol	236 → 125	10	125 → 89	25	236 → 132	20
Permethrin (cis)	183 → 168	15	183 → 153	15	183 → 165	10
Permethrin (trans)	183 → 168	15	183 → 153	15	183 → 165	10
Phosmet	160 → 133	15	160 → 77	20	160 → 105	20
Piperonyl butoxide	176 → 131	10	176 → 145	15	176 → 161	10
Prallethrin	123 → 81	5	134 → 91	10	123 → 95	5
Propiconazole I	173 → 145	20	173 → 109	30	175 → 147	20
Propiconazole II	173 → 145	20	173 → 109	30	175 → 147	20
Propoxur	110 → 63	25	152 → 110	15	110 → 64	20
Pyrethrin I	123 → 81	10	105 → 77	20	123 → 79	15
Pyrethrin II	160 → 117	15	107 → 91	10	133 → 105	10
Pyridaben	147 → 117	20	147 → 115	35	147 → 119	10
Quintozene	295 → 237	15	249 → 214	15	249 → 179	35
Spiromesifen	272 → 254	5	272 → 209	15	272 → 231	15
Spiroxamine I	100 → 72	10	100 → 58	10	100 → 41	20
Spiroxamine II	100 → 72	10	100 → 58	10	100 → 41	20
Tebuconazole	250 → 125	20	125 → 89	20	125 → 99	20
Thiamethoxam	212 → 139	10	247 → 212	5	247 → 182	10
Trifloxystrobin	116 → 89	15	131 → 89	30	131 → 116	20