

## Application Note

No. GC-13-ADI-013

GCMS-TQ8040

### Determination of chemical contaminants in marine fish by GC-MS/MS using QuEChERS as an extraction method

#### □ Introduction

As Persistent Organic Pollutants (POPs) are environmental contaminants with a strong potential for bioaccumulation, these chemicals are expected to be present in farm animals and food products of animal origin.<sup>[1]</sup> In recent years, the occurrence of various POPs, in the marine environment has received much attention with regards to their effects on human health. Especially in India, marine water is contaminated due to uncontrolled disposal from industries and domestic waste. Hence the fish obtained from it has high risk of contamination.

Evidently, it becomes necessary to have sensitive, accurate, reliable, reproducible and fast analytical method to quantify these POPs in marine fish at ppb levels.

Fish is a complex matrix and hence requires selective extraction and extensive cleanup such as QuEChERS (Quick Easy Cheap Effective Rugged Safe) to ensure trace level detection with adequate precision and accuracy. In this study, Shimadzu GCMS-TQ8040 was used in Multiple Reaction Monitoring (MRM) mode to analyze POPs like Organo-Chlorine Pesticides (OCP), Polycyclic Aromatic Hydrocarbons (PAH) and Poly-Chlorinated Biphenyls (PCB).

Prawns sample purchased from local market was extracted, spiked and analyzed for obtaining LOD, LOQ, precision and recovery.

#### □ Experimental

Extraction of pesticides from prawns:

Extraction of pesticides was done using modified AOAC QuEChERS method, as given below<sup>[2]</sup>.

Remove the shell of prawns. Take 10 g of finely grounded prawns sample.  
Add 10 mL of water. Homogenize the sample and Keep it for 30 min.

Add 10 mL of acetonitrile containing 0.5 % acetic acid.

Immediately, add QuEChERS salts containing 6 g MgSO<sub>4</sub> + 1.5 g NaAcetate (Restek® Corporation, Catalog # 26238)

Shake and vortex for 1 min.

Centrifuge at 5000 rpm for 5 min.

Allow to stand for 1 min for phase separation

Draw 6 mL from upper extract of acetonitrile for further cleanup

Transfer the extract to dSPE (dispersive Solid Phase Extraction) tube containing 1200 mg anhydrous MgSO<sub>4</sub>, 400 mg PSA (Primary secondary amines) and 400 mg C18 (Restek® Corporation, Catalog # 26221). Vortex for 2 min.

Centrifuge the mixture at 5000 rpm for 5 min.

Collect the supernatant and filter through a 0.2 µm PTFE membrane filter

Inject 1.0 µL of the clean extract into GCMS-TQ8040

**GC-MS/MS Analytical Conditions**

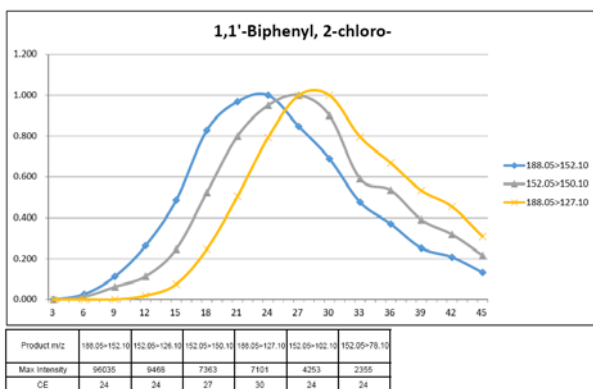
The analysis was carried out on Shimadzu GCMS-TQ8040 as per the conditions given in Table 1.

Table 1. Analytical conditions

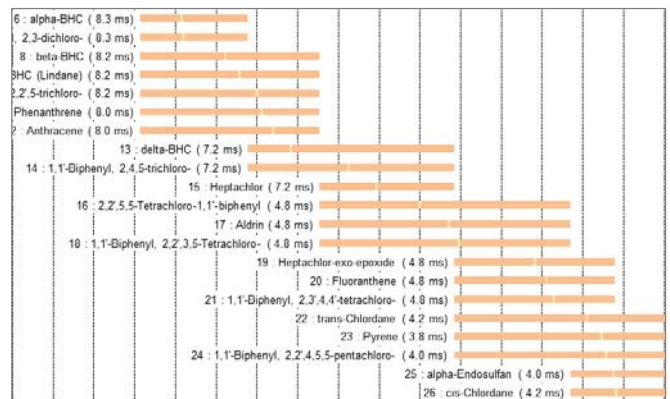
<b>Chromatographic parameters</b>			
• Column	: Rxi-5Sil MS (30 m length, 0.25 mm I.D , df=0.25 µm)		
• Injection Mode	: Splitless		
• Sampling Time	: 2.00 min		
• Split Ratio	: 5.0		
• Carrier Gas	: Helium		
• Flow Control Mode	: Linear Velocity		
• Linear Velocity	: 40.2 cm/sec		
• Column Flow	: 1.2 mL/min		
• Injection Volume	: 1.0 µL		
• Injection Type	: High Pressure Injection (250.0 kPa hold for 1.50 min)		
• Total Program Time	: 45.87 min		
• Column Temp. Program	: Rate (°C /min)	Temperature (°C)	Hold time (min)
		70.0	2.00
	25.00	150.0	0.00
	3.00	200.0	0.00
	8.00	280.0	15.00
<b>Mass Spectrometry parameters</b>			
• Ion Source Temp.	: 230.0 °C		
• Interface Temp.	: 280.0 °C		
• Ionization Mode	: EI (Electron Ionization)		
• Acquisition Mode	: MRM		

**MRM method development**

Individual mixtures of OCP, PAH and PCB standards were procured from Restek®. For analyzing OCP, the MRM transitions existing in Smart Database were used. For PAH, they were obtained from existing application data and imported in Smart Database. In case of PCB, MRMs were optimized. For optimization, about 1 ppm standard mixture of PCB was analyzed using scan mode. Retention times of individual components were identified and precursor ions were selected. Using selected precursor ion, product ion scan was performed with different Collision Energies (CE). For each component of PCB, MRM transitions with appropriate CEs were determined (Refer Figure 1). All the above steps were simplified with the help of Smart MRM optimization tool. These optimized MRMs for PCB were then registered to Smart Database containing transitions of OCP and PAH. From this, the final method with optimum segments (Refer Figure 2) and minimum three MRM transitions per compound was generated. The mixture of OCP, PAH and PCB was analyzed using the created method and the MRM chromatogram for the same is shown in Figure 3.



**Figure 1. CE Optimization using Smart MRM optimization tool**



**Figure 2. Optimum segmented method created using Smart Database**

**Results and Discussion**

Prawns sample was extracted to prepare matrix blank, which was spiked with various concentration levels of POPs to prepare matrix match or post-extraction spike linearity. Using this linearity parameters like LOD, LOQ, precision were studied. Against this linearity, pre-extraction spike was analyzed to study the recoveries.

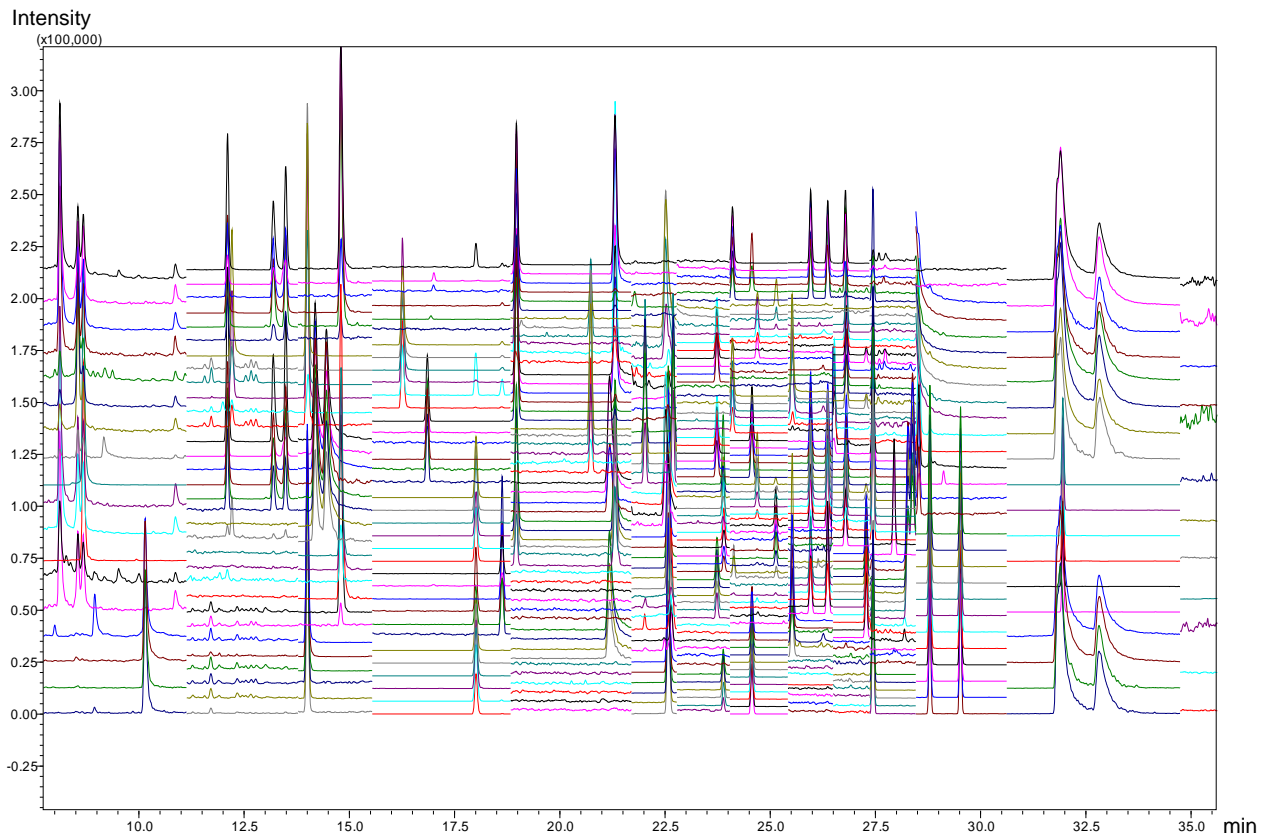


Figure 3. MRM Chromatogram for 10 ppb POPs mixture in prawns matrix

**Summary of the results**

Relative Standard Deviation (% RSD) for 5 ppb standard solution (n=5) was less than 15 % for all components. Calibration plot of matrix match standards ranging from 1 ppb to 50 ppb concentration level showed linear response with  $r^2$  more than 0.995. Recoveries for the prawns sample spiked with 5 ppb standard mixture were in the range of 70 to 130 %. The statistical data obtained for PAH, PCB and OC is shown in Table 2, 3 and 4 respectively.

Table 2. Quantitation results for PAH

ID	PAHs	Retention time (min)	Target MRM (m/z)	$r^2$	S/N at 5 ppb at LOQ level	% RSD at LOQ level (n=5)	% Recovery at LOQ
1	Naphthalene	5.41	128.10>128.10	0.9993	2261.7	3.97	117
2	Acenaphthylene	8.09	152.10>152.10	0.9994	349.37	5.46	94
3	Acenaphthene	8.53	153.10>153.10	0.9999	16.05	1.74	104
4	Fluorene	10.11	166.10>166.10	0.9996	98.46	8.96	96
5	Phenanthrene	14.12	178.10>178.10	1.0000	22.50	3.00	111
6	Anthracene	14.39	178.10>178.10	1.0000	31.09	5.13	95
7	Fluoranthene	21.23	202.10>202.10	1.0000	112.84	5.22	107
8	Pyrene	22.59	202.10>202.10	1.0000	55.95	7.10	102
9	Benzo[a]anthracene	28.44	228.10>228.10	0.9950	17.20	4.25	124
10	Chrysene	28.44	228.10>228.10	0.9948	21.66	4.47	131
11	Benzo[k]fluoranthene	32.00	252.10>252.10	0.9999	999.8	4.80	85
12	Benzo[b]fluoranthene	32.01	252.10>252.10	0.9998	303.42	7.99	91
13	Benzo[a]pyrene	32.97	252.10>252.10	0.9993	88.39	9.18	79
14	Indeno[1,2,3-cd]pyrene	37.37	276.10>276.00	0.9996	22.22	7.19	70
15	Dibenz(a,h)anthracene	37.45	278.10>278.10	0.9948	36.15	12.28	94
16	Benzo[g,h,i]Perylene	38.42	276.10>276.10	0.9996	918.40	9.98	70

Table 3. Quantitation results for PCB

ID	PCBs	Retention time (min)	Target MRM (m/z)	r <sup>2</sup>	S/N at 5 ppb at LOQ level	% RSD at LOQ level (n=5)	% Recovery at LOQ
1	1,1'-Biphenyl, 2-chloro-	8.67	188.05>152.10	0.9998	3181.75	3.25	95
2	1,1'-Biphenyl, 2,3-dichloro-	12.19	222.00>152.10	0.9999	691.78	1.48	92
3	1,1'-Biphenyl, 2,2',5-trichloro-	13.99	255.95>186.00	0.9999	1516.47	1.01	93
4	1,1'-Biphenyl, 2,4,5-trichloro-	16.25	255.95>186.00	0.9999	582.39	2.17	85
5	2,2',5,5-Tetrachloro-1,1'-biphenyl	18.00	289.90>220.00	0.9999	1326.79	2.16	86
6	1,1'-Biphenyl, 2,2',3,5-Tetrachloro-	18.96	289.90>220.00	0.9998	597.52	4.12	87
7	1,1'-Biphenyl, 2,3',4,4'-tetrachloro-	21.29	289.90>219.90	1.0000	117.87	2.57	80
8	1,1'-Biphenyl, 2,2',4,5,5-pentachloro-	22.57	325.90>255.90	0.9999	1487.66	5.86	73
9	1,1'-Biphenyl, 2,2',3,4,5-pentachloro-	23.73	325.90>255.90	0.9998	263.70	6.44	75
10	2,3,3',4,6'-Pentachloro-1,1'-biphenyl	24.09	325.90>255.80	0.9997	303.94	1.99	80
11	2,2',3,5,5',6'-Hexachloro-1,1'-biphenyl	24.56	359.85>289.80	0.9999	635.77	4.58	70

Table 4. Quantitation results for OC

ID	OCs	Retention time (min)	Target MRM (m/z)	r <sup>2</sup>	S/N at 5 ppb (LOQ level)	% RSD at LOQ level (n=5)	% Recovery at LOQ
1	alpha-BHC	12.15	180.90>144.90	0.9999	740.26	4.36	106
2	beta-BHC	13.23	180.90>144.90	0.9999	424.32	1.78	113
3	gamma-BHC (Lindane)	13.57	180.90>144.90	0.9999	484.21	3.37	106
4	delta-BHC	14.83	180.90>144.90	0.9997	132.15	4.84	100
5	Heptachlor	16.92	271.80>236.90	1.0000	376.67	7.29	98
6	Aldrin	18.71	262.90>193.00	0.9997	62.23	3.17	78
7	Heptachlor-exo-epoxide	20.82	352.80>262.90	0.9998	132.20	6.57	114
8	trans-Chlordane	22.10	372.80>263.90	0.9999	424.26	7.84	90
9	cis-Chlordane	22.69	372.80>263.90	1.0000	388.00	7.73	82
10	alpha-Endosulfan	22.74	194.90>160.00	0.9999	11.86	11.71	97
11	Dieldrin*	23.95	262.90>193.00	0.9997	162.75	10.58	78
12	p,p'-DDE	23.99	246.00>176.00	1.0000	205.29	3.00	86
13	Endrin*	24.78	262.90>191.00	1.0000	184.28	8.87	107
14	beta-Endosulfan*	25.22	194.90>160.00	0.9988	24.94	11.52	107
15	p,p'-DDD	25.63	235.00>165.00	1.0000	365.71	4.30	109
16	Endosulfan sulfate	26.61	271.80>236.90	0.9988	183.35	11.75	95
17	p,p'-DDT	26.86	235.00>165.00	1.0000	101.05	5.24	103

Note \* : For these compounds LOQ was 10 ppb

#### □ Conclusion

Shimadzu GCMS-TQ8040 with Smart MRM feature was able to optimize MRM transitions with ease.

New Smart Database tool creates method with optimum segments leading to increased dwell-time, which resulted in achieving high sensitivity for trace level quantitation of POPs in complex matrix like prawns.

The MRM method developed for POPs can be used for screening of pesticides in various marine fish products. For 90 % of the POPs, LOQ of 5 ppb was achieved.

#### □ References

1. Weiss, J., Paepke, O. and Bergman, A. 2005. A worldwide survey of polychlorinated dibenzo-p-dioxins, dibenzofurans, and related contaminants in butter. *Ambio* 34(8):22-30.
2. Pesticide Residue in Foods by Acetonitrile Extraction and Partitioning with Magnesium Sulfate (AOAC Official Method 2007.01), (2007), 06.

