

Application News

Gas Chromatography Mass Spectrometry Liquid Chromatography Mass Spectrometry

No. **M293**

Analysis of Toxic Chemical Substances Adsorbed on Microplastics

The possibility that toxic chemical substances adsorbed on microplastics (MPs) in the environment may impact the ecosystem by desorbing from the MPs, migrating to the bodies of living organisms, and concentrating in those organisms has been pointed out.

The Shimadzu Group has been involved in evaluations of the adsorption characteristics of chemical substances on MPs as part of analysis of MPs $^{(1),\,(2)}$.

Here, we introduce an example of an evaluation of the MPs adsorption characteristics of polycyclic aromatic hydrocarbons (PAHs) and per- and polyfluoroalkyl substances (PFAS), which are known to have toxicity and bioaccumulation properties. Using commercially-available particulate polypropylene (PP), polystyrene (PS), and polyethylene (PE) as MPs samples, the amount of adsorption of PAHs and PFAS in water on the various plastics was evaluated quantitatively by GC-MS/MS and LC-MS/MS.

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MPs Samples

The following three types of plastic particles were used as the MPs samples.

- 1) Polypropylene (PP)
- 2) Polystyrene (PS)
- 3) Polyethylene (PE)

The long and short diameters of each type of particle were calculated from stereoscopic microscope images to confirm that the size of the materials used was less than 5 mm, which is the definition of MPs. Table 1 shows the appearance and microscopic images of the above-mentioned plastic particles 1) to 3), together with the average size of 10 particles sampled at random. The following instruments were used in the measurements.

- Microscope digital camera system : Moticam 1080
- Stereoscopic microscope : STZ-171-TP

Test Method

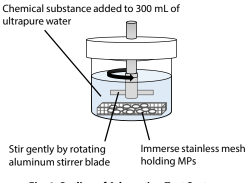
First, an adsorption test of PAHs and PFAS (per- and polyfluoroalkyl substances)^(*) on the MPs samples was conducted. Fig. 1 shows an outline of the adsorption test system. The MPs were immersed in water, to which PAHs or PFAS had been added, and the water was stirred gently for 24 h to promote adsorption. The amounts of added to MPs were 100 ng of PAHs and 8 ng of PFAS in 300 mL of ultrapure water.

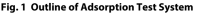
Fig. 2 shows an outline of the test flow. After the adsorption test, the MPs were removed from the test system and dried. With some of the samples, ultrasonic extraction by hexane was used as a pretreatment for the PAHs, and ultrasonic extraction by methanol was used as a pretreatment for the PFAS. The extracts obtained here were injected into the GC-MS/MS and LC-MS/MS, respectively, for quantitative analysis of the PAHs and PFAS.

(*) For analogs, refer to EPA Method 537.1 and EPA Method 533.

Material		РР	PS	PE	
Appearance					
Microscopic image					
Size [mm]	Long diameter	4.899±0.151	3.191±0.149	4.331±0.194	
	Short diameter	3.606±0.151	2.888±0.144	3.564±0.125	

Table 1 MPs Samples Used in Experiment





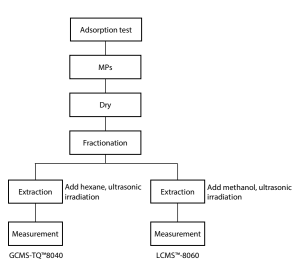


Fig. 2 Outline of Test Flow

Measurement of PAHs by GC-MS/MS

In measurement of the PAHs, a GCMS-TQ8040 triple quadrupole gas chromatograph mass spectrometer (Fig. 3) was used. Table 2 shows the measurement conditions of the GCMS-TQ8040.

The measurements were carried out in the high selectivity MRM mode. Based on the calibration curve linearity and peak S/N ratio, the lower limit of quantitation of the compounds was assumed to be 0.5 to 2 ng/mL. Fig. 4 shows the MS chromatogram of the PAHs standard solution (2 ng/mL each). Quantitative calculations were performed by the internal standard method.



Fig. 3 GCMS-TQ[™]8040 Triple Quadrupole Gas Chromatograph Mass Spectrometer

Table 2 GCMS-TQ8040 Measurement Conditions GC DB-5ms (Agilent Technologies/J&W) Column $(30 \text{ m} \times 0.25 \text{ mm} \text{ l.D.}, 0.25 \text{ }\mu\text{m})$ 60 °C (1 min) – 15 °C/min – 200 °C (0 min) – 8 °C/min – 320 °C (10 min) Column oven temp. program Injection mode **Splitless** 300 °C Vaporizing chamber temp. Injection volume 2 µL Carrier gas Helium (1.0 mL/min) MS FI Ionization method 70 V Ionization voltage 300 °C Interface temp. Measurement mode MRM Measured ions (m/z) Compound (target substance) Quantitative ion Reference ion Compound (internal standard) Quantitative ion Reference ion Naphthalene 128>128 128>102 Naphthalene-d₈ 136>136 136>108 Acenaphthylene 152>152 152>151 Acenaphthylene-d₈ 160>160 160>158 Acenaphthene- d_{10} Acenaphthene 153>153 153>152 162>162 162>160 Fluorene 165>165 165>164 Fluorene-d₁₀ 174>174 174>171 Anthracene-d₁₀ Anthracene 178>178 178>152 188>188 188>160 Phenanthrene 178>178 178>152 Phenanthrene-d₁₀ 188>188 188>160 Fluoranthene 202>202 202>200 Fluoranthene- d_{10} 212>212 212>208 Pyrene 202>202 202>200 Pyrene- d_{10} 212>212 212>208 Benz[a]anthracene-d₁₂ Benzo[a]anthracene 240>240 240>236 228>228 228>226 Chrvsene 228>228 228>226 Chrvsene- d_{12} 240>240 240>236 Benzo[b]fluoranthene 252>252 252>250 Benzo[b]fluoranthene-d₁₂ 264>264 264>260 Benzo[k]fluoranthene Benzo[k]fluoranthene-d₁₂ 252>252 252>250 264>264 264>236 Benzo[a]pyrene 252>250 Benzo[a]pyrene-d₁₂ 264>264 264>260 252>252 Benzo[ghi]perylene 276>276 276>274 Benzo[ghi]perylene-d₁₂ 288>288 288>284 Indeno[1,2,3-cd]pyrene Indeno[1,2,3-cd]pyrene-d₁₂ 288>284 276>276 276>274 288>288 Dibenzo[a,h]anthracene 278>276 Dibenzo[a,h]anthracene-d₁₄ 278>278 292>292 292>288 Hexachlorobenzene-13C6 292>292 292>292 (x100,000) 1.25

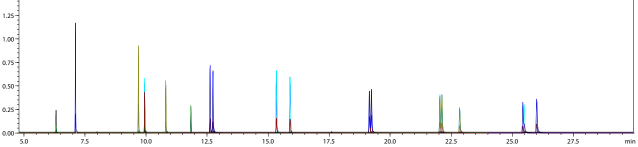


Fig. 4 MRM Chromatogram of PAHs Standard Solution (2 ng/mL Each) (Only Target Substances)

Measurement of PFAS by LC-MS/MS

The instrument used in measurements of the PFAS was a LCMS-8060 triple quadrupole liquid chromatograph mass spectrometer (Fig. 5). Table 3 shows the measurement conditions.

The measurements were carried out in the high selectivity MRM mode. Based on the calibration curve linearity and peak S/N ratio, the lower limit of quantitation of the compounds was assumed to be 0.01 to 0.5 ng/mL. Fig. 6 shows the MS chromatogram of the PFAS standard solution (0.5 ng/mL each). Quantitative calculations were performed by the internal standard method.



Fig. 5 LCMS[™]-8060 Triple Quadrupole Liquid Chromatograph Mass Spectrometer

Table 3 LCMS-8060 Measurement Conditions

Instruments	LC : Nexera™ X2 MS : LCMS-8060 triple quadrupole liquid chromatograph mass spectrometer							
LC								
Column	Inertsil ODS-SP (GL Sciences) (150 mm × 2.1 mm l.D., 3 μm)							
Column temp	40 °C							
Injection volume	10 μL							
Mobile phase A	10 mmol/L ammonium acetate aqueous solution							
Mobile phase B	Acetonitrile							
Mobile phase flow rate	0.2 mL/min							
Gradient	B conc. 0% (0 min) $ ightarrow$ 10% (1 min) $ ightarrow$ 40% (7 min) $ ightarrow$ 95% (31-38 min) $ ightarrow$ 10% (47 min)							
MS								
Ionization method	ESI							
Polarity	Negative							
Measurement mode	MRM							
MRM transition								
Compound (target substance)	Quantitative ion	Reference ion	Compound (internal standard)	Quantitative ion	Reference ion			
PFHxA	313>269	313>119	PFHxA- ¹³ C₅	318>273	318>121			
PFHpA	363>319	363>169	PFHpA- ¹³ C ₄	367>322	367>169			
PFOA	413>369	413>169	PFOA- ¹³ C ₈	421>376	421>172			
PFNA	463>419	463>219	PFNA- ¹³ C ₉	472>427	472>223			
PFDA	513>469	513>219	PFDA- ¹³ C ₆	519>474	519>219			
PFUnDA	563>519	563>269	PFUnDA- ¹³ C ₇	570>525	570>274			
PFDoDA	613>569	613>269	PFDoDA-13C2	615>570	615>169			
PFTrDA	663>619	663>169	-	-	-			
PFTeDA	713>669	713>169	PFTeDA- ¹³ C ₂	715>670	715>169			
PFHxDA	813>769	813>169	PFHxDA- ¹³ C ₂	815>770	815>169			
PFOcDA	913>869	913>169	-	-	-			
PFBS	299>80	299>99	PFBS- ¹³ C ₃	302>80	302>99			
PFPeS	349>80	349>99	-	-	-			
PFHxS	399>80	399>99	PFHxS- ¹³ C ₃	402>80	402>99			
PFHpS	449>80	449>99	-	-	-			
PFOS	499>80	499>99	PFOS- ¹³ C ₈	507>80	507>99			
PFNS	549>80	549>99	-	-	-			
PFDS	599>80	599>99	-	-	-			
PFDoS	699>80	699>99	-	-	-			

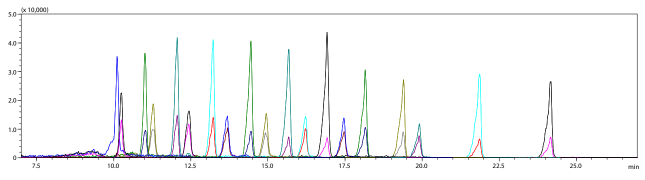


Fig. 6 MRM Chromatogram of PFAS Standard Solution (0.5 ng/mL Each) (Only Target Substances)

Analysis Results

Fig. 7 shows the results of the analysis of the PAHs by GC-MS/MS, and Fig. 8 shows the results of the analysis of the PFAS by LC-MS/MS. Adsorption on the MPs was confirmed for all PAHs substances and for some PFAS substances. Adsorption of the PAHs on PP and PE tended to be large. However, the amount of adsorption of the PFAS tended to differ for each chemical substance. Based on these results, because the adsorption characteristics on MPs differed depending on the chemical substance, it is thought that some are easily affected by the MPs material, in other words, its molecular structure and others are not so.

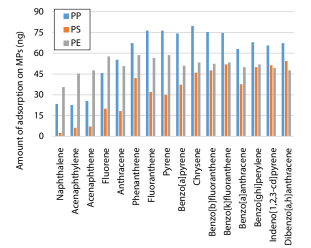


Fig. 7 GC-MS/MS Analysis Results: PAHs

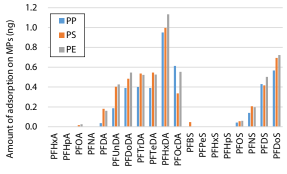


Fig. 8 LC-MS/MS Analysis Results: PFAS

Conclusion

In order to clarify the adsorption characteristics of chemical substances on MPs, the amount of adsorption of PAHs and PFAS on MPs in water was evaluated quantitatively by using GC-MS/MS and LC-MS/MS.

It was found that adsorption characteristics on MPs differed for each chemical substance. In addition, it was also suggested that the hydrophobicity of the individual chemical substances is a critical factor for their transfer ratio. Adsorption of chemical substances on MPs is considered to depend mainly on their hydrophobic interaction. Therefore, we examined the correlation between Log Kow^(*1) and Log D^(*2), which show the hydrophobicity of chemical substances, and chemical substances transfer ratio to MPs. The transfer ratio to MPs was defined as the value obtained by dividing the amount of PHAs or PFAS adsorbed on the MPs by the amount of those substances added to water. Fig. 9 shows the relationship between the hydrophobicity of the chemical substances and the transfer ratio to the MPs. In order from the top, this figure shows the transfer ratios to PP, PS, and PE. In these results, the transfer ratio showed a tendency to increase with higher values of Log Kow and Log D, independent of the type of MPs.

- (*1) Log Kow : octanol/water partition coefficient
- (*2) Log D : Log Kow is a partition coefficient of a chemical substance which considers the influence of electrolytic dissociation (ionization) in water. Here, this coefficient was calculated for water having pH 6.4 at the time of the test.

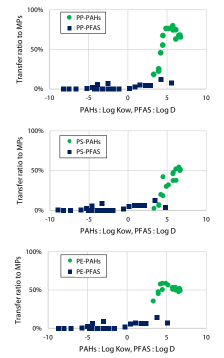


Fig. 9 Hydrophobicity of Chemical Substances and Transfer Ratios to MPs (Top: PP, Middle: PS, Bottom: PE)

<References>

- (1) Makoto Yasojima, Haruna Mizuka, Takaki Mine, Hiroaki Takemori, Shinji Takeuchi, Yoshihide Yasui, "Adsorption Characteristics of Chemical Substances on Microplastics," Proceedings of the 22nd Symposium of the Japan Society on Water Environment (2019)
- (2) Makoto Yasojima, Haruna Mizuka, Takaki Mine, Hiroaki Takemori, "Existence of Unknown Chemical Substances Adsorbed on Microplastics Immersed in Rivers and Adsorption Characteristics of Chemical Substances on Microplastics," Proceedings of the 56th Environmental Engineering Research Forum (2019)

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