

Quantitative Analysis of Microplastics Using the Pyroprobe with GC-MS

Application Note

Environmental

Author:

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Abstract

This application note demonstrates quantitative analysis of microplastics by pyrolyzing the sample into a GC-MS.

Introduction

Over 300 million tons of plastics are being manufactured every year and the scale of production is growing at a faster pace¹. Studies showed that >10% of plastics eventually entered and contaminated the ocean^{2,3}. Beside the traditional topics on cleaning macro plastic pollution in the sea, a growing attention on microplastics in sea water with <5 mm diameter size has been rising⁴. Many of the existing scientific surveys on the microplastics in ocean were using FT-IR and Raman techniques. Automation was recently achieved by optic method⁵ to improve testing speed. However, such technique could only provide limited information on polymer type and size and is not capable in offering an accurate weight percentage of the pollutants. PY-GC-MS technique was introduced for this purpose⁶.

In the previous publication⁷, Pyroprobe was shown to identify microplastics in the sea water over ppm concentration when coupling to a mainstream GC-MS. This application note is a continued study to further quantify the microplastics concentration.

Experimental Setup

Five different plastics, which include PP, PET, PVC, PS and PE, were mixed at a random ratio. The mixing sample, 100 μ g, was weighed and loaded into a DISC tube to mimic a complex sample concentrated from sea water. This simulant was then analyzed by a CDS 6150 Pyroprobe with a GC-MS. The system parameters are listed below:

Experimental Parameters

6150 Pyroprobe	GC-MS:
DISC: 700°C	Column: 5% phenyl (30m x 0.25mm)
Ramp Rate: 20°C per ms	Carrier: Helium 1.00 mL/min,
DISC Interface: 300°C	50:1 split
Transfer Line: 300°C	Injector: 360°C
Valve Oven: 300°C	Oven: 40°C for 2 minutes
	12°C/min to 320°C (10min)
	Ion Source: 230°C
	Mass Range: 35-600amu

Results and Discussion

Figure 1 is the pyrogram of the plastic mixture sample. All five polymers were all correctly identified. Table 1 listed each of the identified polymer with its marked peak. Figure 2 further displayed all the marked peaks for each polymer.

After polymer identification, a calibration curve was drawn on polystyrene only to demonstrate the feasibility of mass quantification on each component. The calibration curve was composed of four points.



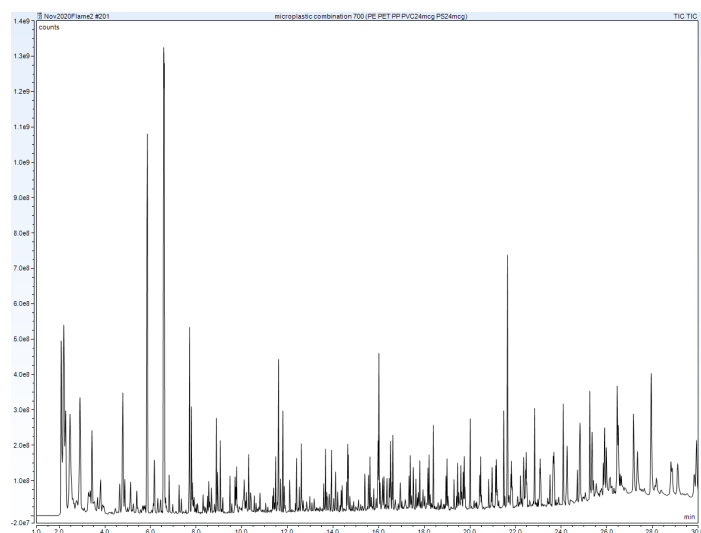
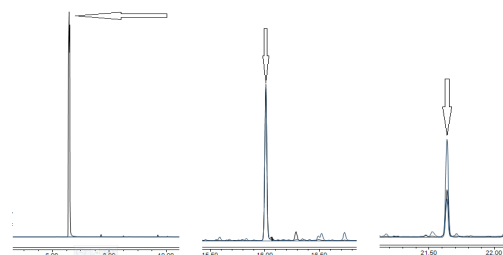


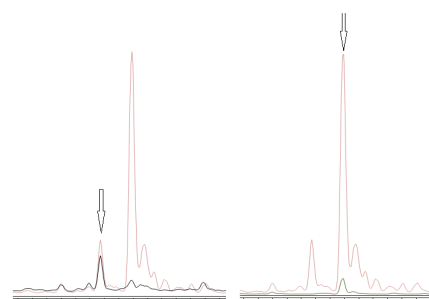
Figure 1. Pyrogram of mixed plastics, containing polypropylene, polyethylene terephthalate, polyvinyl chloride, polystyrene, and polyethylene.

Table 1. Mixed Plastic Marker Peaks

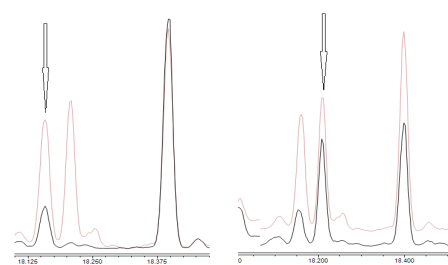
Polymer	Marker Peaks	Retention times	Ions
Polypropylene (PP)	trimer	5.86	70, 126
	tetramer	8.86, 8.90	69, 111
	pentamer	11.58, 11.77	69, 111
Polyethylene terephthalate (PET)	vinyl benzoate	9.66	77, 105
	benzoic acid	10.24	105, 122
Polyvinyl chloride (PVC)	HCl	2.08	36
	naphthalene	10.38	128
Polystyrene (PS)	monomer	6.58	104
	dimer	16.01	208
	trimer	21.64	312
Polyethylene (PE)	C10 diene	7.8	55, 81
	C10 alkene	7.69	55, 140
	C20 diene	18.16	55, 109
	C20 alkene	18.21	55, 111



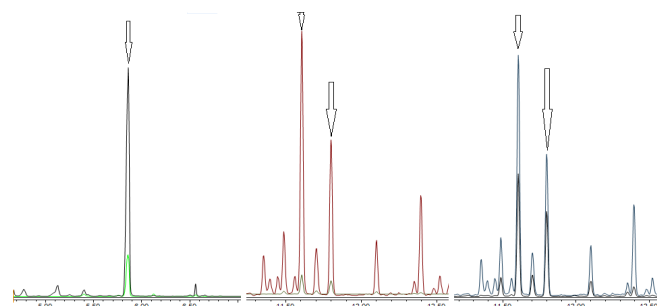
PS: monomer, dimer, trimer



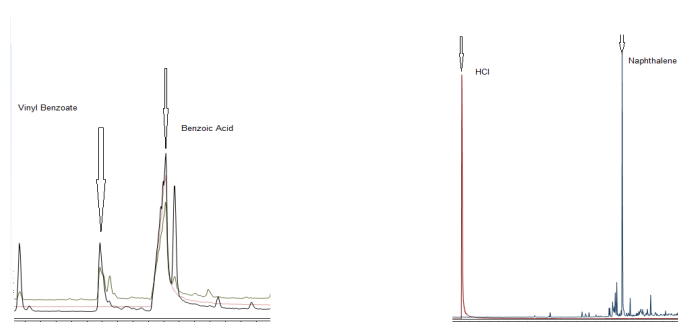
PE: C10 diene and alkene.



PE: C20 diene and alkene



PP: trimer, tetramer, pentamer.



PET: vinyl benzoate, benzoic acid

PVC: HCl, naphthalene

Figure 3. Extracted ion chromatograms for marker peaks in mixed plastics sample.

To make the calibration curve, a calibration stock solution was first prepared by dissolving polystyrene in toluene, resulting in a final polystyrene concentration of $11 \mu\text{g}/\mu\text{L}$. Aliquots of 0.2, 0.4, 0.6, and 0.8 μL stock solution with an absolute mass of polystyrene at 2.2, 4.4, 6.6, and 8.8 μg respectively were added to 4 Drop-In-Sample-Chamber (DISC) tubes individually. Each calibration sample was pyrolyzed at the same condition, and areas of styrene monomer were plotted against the absolute mass of polystyrene. The calibration curve was then fitted with a single order polynomial in Figure 4 and the R^2 of data fitting was 0.997, which showed excellent linearity.

The area of styrene in the mixed plastic sample was calculated by integrating its monomer peak. This area was plugged into the calibration line in Figure 4 to calculate the polystyrene mass as 7.69 μg .

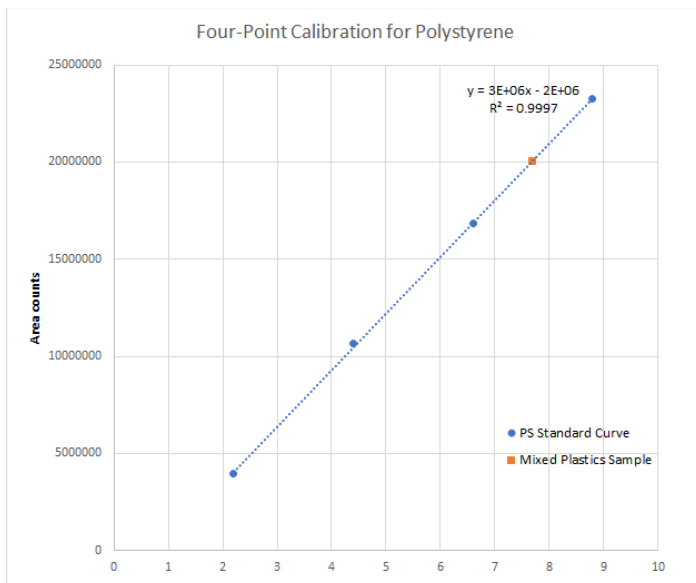


Figure 4. Polystyrene calibration.

Conclusions:

A mixture of 5 different polymers that mimics a complex environmental sample from sea water were analyzed quantitatively by Py-GC-MS method. The CDS Pyroprobe was demonstrated as a useful tool in investigating the pollution caused by microplastics.

References

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- (3) Jambeck, Jenna R., et al. "Plastic waste inputs from land into the ocean." *Science* 347.6223 (2015): 768-771
- (4) Barnes, David KA, et al. "Accumulation and fragmentation of plastic debris in global environments." *Philosophical Transactions of the Royal Society B: Biological Sciences* 364.1526 (2009): 1985-1998
- (5) K  ppler, Andrea, et al. "Identification of microplastics by FTIR and Raman microscopy: a novel silicon filter substrate opens the important spectral range below 1300 cm⁻¹ for FTIR transmission measurements." *Analytical and bioanalytical chemistry* 407.22 (2015): 6791-6801
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- (7) Sam, Karen, Detection of Plastic Pollution by the Pyroprobe, CDS App Note #214