

Determination of Limonene in Recycled HDPE Pellets

Application Note

AN0019

INTRODUCTION

Over four and a half million tonnes of plastic is produced annually in the UK alone, with over 85% of this going to landfill⁽¹⁾. Over 45% of plastic bottles are sent to recycling with the number expected to significantly increase in the coming years. Recycled plastics have a variety of uses from new bottles/ food trays to underground draining products and soft furnishing materials⁽¹⁾.

During the recycling process, clear and mixed colour plastics are separated before they are granulated and re-separated. The granulated plastics are then thoroughly cleaned and dried before being tested in quality control and sent to be manufactured. The plastics are turned into high density polyethylene (HDPE) pellets.

It is vital that the pellets produced from the recycling are deodorised from contaminants and flavour compounds especially when they are manufactured into food packaging.

Limonene is a cyclic monoterpene and is the major component of the oil in citrus fruits. Limonene contaminates recycling materials; during the recycling process, high levels of limonene in the pellets must be removed during deodorisation ⁽²⁾. During quality control testing, the pellets must have a limonene concentration of less than 20ppm.

This application note demonstrates the extraction of limonene from the HDPE pellet with excellent limit of detection using the highly sensitive SCION 436 GC with single quad (SQ) mass spectrometer (MS).

EXPERIMENT

Table 1 details the analytical parameters used throughout this application.

Table 1. Analytical conditions of the GC-MS

Conditions		
Injector	S/SL at 250°C	
Columns	30m x 0.25mm x 1.4µm (Scion624-MS)	
Oven Conditions	55°C (hold 1 minute)	
	220°C at 20°C/min (hold 1 minute)	
Carrier	Helium 1mL/min	
Transfer Line	250°C	
Ion Source	200°C	
MS	Full Scan 42m/z to 350m/z	
Peak Identification	Limonene (Ions 93/68)	

Limonene calibration standards were prepared in Carbon Disulphide (CS_2) at a range of Oppm, 5ppm, 10ppm, 20ppm and 40ppm.

0.1g HDPE pellets were extracted using 1mL of CS₂ and incubated at room temperature for sixty minutes. The extracts were then injected into the GC using a 8400 autosampler. The HDPE pellets were analysed in triplicate.

A full scan method was used to identify m/z 93, the quantifier ion of limonene with m/z 68 as a qualifier ion. The total ion chromatogram (TIC) was subjected to a NIST library search for confirmation.

RESULTS

The limonene was identified with a retention time of 7.265 minutes using the full scan method and injection of an analytical standard. Figure 1 shows the TIC of limonene and the associated spectra obtained via the full scan method.

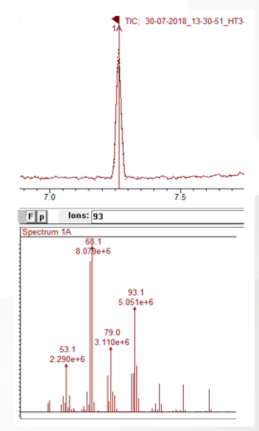


Figure 1. Limonene TIC and spectra





A NIST library search of the limonene peak gave a spectral match of 91%. Figure 2 shows the NIST library search, generated through the SCION Mass Spectrometry Work Station (MSWS) software.

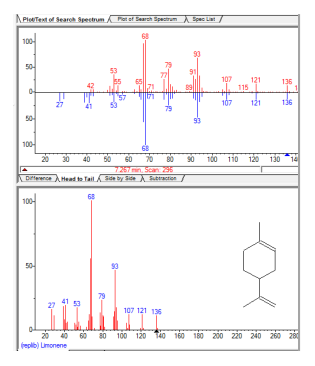


Figure 2. NIST library search; 91% match

After accurate identification of limonene, via the NIST library, the calibration standards were analysed in triplicate. A blank sample was also analysed for quality control purposes. Figure 3 shows the calibration curve generated from the triplicate injections, using the average of the replicates.

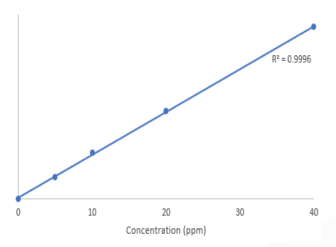


Figure 3. Calibration curve of limonene

The calibration showed excellent linearity with an R^2 value of 0.9996. The HDPE pellets were then also analysed in triplicate (three separate samples prepared). Retention time and extraction ion 93 were used for confirmation of the limonene. Figure 4 details the TIC and TIC for ion 93.

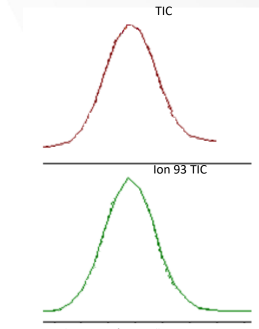


Figure 4. TIC and Ion 93 TIC of HDPE pellet

The calibration curves were used to calculate the concentration of limonene in the samples. Table 2 details the peak area and calculated concentration of each injection as well as the average peak area, average concentration and RSD% of the analysis.

Injection	Peak Area	Concentration (ug/100mg)
1	5.933E ⁺⁷	0.001
2	5.953E ⁺⁷	0.001
3	5.955E ⁺⁷	0.001
Ave.	5.947E ⁺⁷	0.001
%RSD	0.20	0

As shown in table 2, the %RSD for the peak area was exceptional, at only 0.20%, with the %RSD of calculated concentration being zero. This shows the robustness of the SCION 436 GC with SQ MS. The concentration of limonene in the HDPE pellets was $0.01\mu g/g$ (0.01ppm), well below the 20ppm limit allowed to be present in recycled plastics. This extremely low presence of limonene in the pellets demonstrate that the recycled plastics were successfully deodorised during the milling process.





CONCLUSION

The high sensitivity and robustness of the SCION GC-MS coupled with a simple sample extraction made analysis of the HDPE pellets quick and easy, generating confident, reproducible results. The analysed HDPE pellets contained 0.01ppm of limonene. The limonene concentration was significantly lower than the 20ppm limit clearly showing that the pellets have been deodorised, thus passing quality control.

REFERENCES

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[2] Welle, F. (2005). Develop A Food Grade HDPE Recycling Process. Oxon: The Waste & Resources Action Programme, p.4.

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