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Research Article

Organic compositional analysis of propellant powders using monolithic material sorption extraction (MSSE)-a feasibility study

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Abstract

The application of monolithic material sorption extraction, specifically MonoTrap™, to the extraction of organic gunshot residue (OGSR) compounds from unburnt propellant powders is described. Four different MonoTraps were assessed for their capability to extract OGSR compounds from two different ammunition types. Extracts were analysed using gas chromatography-mass spectrometry. Results indicated that the carbon disc was suitable for the extraction of OGSR compounds from unburnt propellant powders. Quantities for major compounds were comparable to methanol extractions. The method was successfully used to detect a wide range of OGSR compounds, including nitrotoluenes, nitroglycerin, diphenylamines and centralites and is expected to be applicable to a wide range of ammunition types.

Introduction

The analysis of additives in smokeless propellant powder is often an integral part of investigations of improvised explosive devices (IEDs) and the evaluation of organic gunshot residue (OGSR) [1,2]. Recently, several methods for the characterisation of OGSR compounds in smokeless propellant powders have been developed [3-5]. Both solvent extraction and solid-phase microextraction (SPME) have been employed for this purpose.

Monolithic material, which is used for sorptive extraction, consists of rigid macroporous polymers prepared by bulk polymerisation in a closed mould [6]. The material is created by polymerisation of a monomer mixture with a porogenic solvent, forming a highly porous polymer bed [7]. Inorganic monolithic material, which is used for MonoTrap™, mainly consists of porous high-purity silica [6,8]. Monolithic material is also widely used as a stationary phase in liquid chromatography [7,9-11]. Other applications include its use as a column for anion-exchange chromatography [12], in tube solid-phase extraction (SPE) [9], in needle extraction [10,13] and stir bar/stir rod sorptive extraction (SBSE/SRSE) [7,14,15]. MonoTraps are employed for a wide range of biological applications, including the extraction of flavour and aroma compounds [16], the characterisation of odorants [17], and the determination of plant hormones [18]. It has wide uses in separation science and in (bio) catalysis [9], and is used for sample preparation in drug and pharmaceutical analysis [11].

MonoTrap™ has a large surface area (150 m²/gram or more), which is provided by the network of interconnected pores in the silica frame [7,8,19]. Due to the network of pores, with sizes in the low micrometer range, this material possesses very good
permeability. Consequently, high analyte migration rates are achieved [7,8,11,19]. This results in significantly faster extraction times for stir bars coated with monolithic materials (up to a few hours [7,14]) than for stir bars coated with a thick layer of a material, such as polydimethylsiloxane (PDMS) (up to 72 hours [20]). MonoTrap™ is suitable for non-polar to polar compounds [19] and it requires only 200 μL of solvent to perform the extraction, which is both environmentally and economically advantageous [4]. MonoTrap™ is available in a silica type and in a type in which the silica frame contains activated carbon that acts as an adsorptive medium. This type of MonoTrap™ has octadecyl-groups conjugated to its frame [8,11]. Both the silica and the carbon type are available in disc and rod configurations [11].

The application of monolithic material sorption extraction (MMSE), more specifically MonoTrap™, to the extraction of OGSR compounds commonly found in smokeless propellant powders and GSR samples is examined for the first time in this study. The ability to extract a wide range of OGSR compounds is demonstrated using a standard solution. The selection of the compounds included in the standard is based on previous work, in which relevant compounds associated with propellant are highlighted [4,21]. A comprehensive list of organic compounds associated with propellant and GSR can be found elsewhere [22]. The capability of MonoTrap™ to extract OGSR compounds from unburnt ammunition propellant is evaluated. Different types (i.e. silica and carbon) and different configurations (i.e. discs and rods) are compared, and the ability of these materials to recover OGSR compounds present in propellant powder, and thus potentially in OGSR samples, is discussed.

Materials and Methods

Solvents and standards

Camphor, carbazole, diphenylamine (DPA), 4-nitrodiphenylamine (4-NDPA), 2,4-dinitrodiphenylamine (2,4-DNDPA), ethylphenylamine (EPA), dimethyl phthalate (DMP), diethyl phthalate (DEP), dibutyl phthalate (DBP), diisobutyl phthalate (DIBP), 3-nitrotoluene (3-NT), 2,3-dinitrotoluene (2,3-DNT), 2,4-dinitrotoluene (2,4-DNT), 2,6-dinitrotoluene (2,6-DNT), 3,4-dinitrotoluene (3,4-DNT), and triacetin were purchased from Sigma Aldrich (Bellefonte, PA, USA). 2-Nitrodiphenylamine (2-NDPA) was obtained from LGC Standards (Middlesex, UK). Methyl centralite (MC) was purchased from Santa Cruz Biotechnology (Dallas, TX, USA). Analytical grade methanol (Sigma Aldrich, UK) was used for the preparation of the standard. MonoTrap desorption was performed with analytical grade methanol or dichloromethane (DCM) (Fisher Scientific, Loughborough, UK).

Acquisition of propellant

Hodgdon HP-38 propellant for self-loading ammunition was obtained from the Wellington Rifle and Pistol Club (Skipton, Yorkshire, UK). Nottinghamshire police provided. 223 Magtech Rem tactical rounds, of which the bullet was pulled using a kinetic hammer in order to collect the unburnt propellant.

MonoTrap™ extractions

MonoTrap™ adsorptives (silica disc, silica rod, carbon disc, and carbon rod) were purchased from GL Science Inc. (Tokyo, Japan). The extractions were carried out by placing the MonoTrap™ directly onto 100 mg propellant powder or onto the standard solution. The vial was then placed in an oven (Nabertherm) at 80°C for 3 hours as per the manufacturer’s recommendations.

Analyte desorption was accomplished by solvent extraction of the MonoTrap™ with 200 μL of methanol or DCM. After 5 minutes of sonication at ambient temperature, the MonoTrap™ was removed and as much of the solvent as possible was transferred to a GC vial (Chromacol ltd., Herts, UK) containing a 150 μL glass insert with polymeric
feet (Agilent Technologies, Santa Clara, CA, USA) [8]. Sequential desorption of a single MonoTrap™ was also attempted by performing the desorption procedure with DCM, and subsequently repeating it using methanol.

**Methanol extractions of propellant powder**

Methanol extractions were performed by adding 2 mL of methanol to 100 mg propellant powder. The vials were sonicated for 1 hour [4], after which the supernatant was centrifuged for 15 minutes and then filtered through a 45 μm PTFE filter.

**GC-MS instrumentation and conditions**

Chromatographic analysis, optimised from the method reported by Dalby & Birkett [4], was performed on an Agilent 6890N Network GC system, equipped with a J&W Scientific HP5-MS UI (30 m x 0.25 mm x 0.25 μm) column. Sample introduction was performed using an autosampler (Agilent 7683 series). Splitless injection and a solvent delay of 1.8 minutes were used for all samples. The following chromatographic conditions were used: initial oven temperature of 50°C, rising by 10°C/min to 100°C, second ramp of 5°C/min to 180°C and held for 2.5 min, third ramp of 30°C/min to 200°C and held for 2.5 min, and a fourth ramp of 30°C/min to 300°C, which was held for 2 min. The total run time was 32 minutes. The flowrate of the carrier gas (helium) was maintained at 1.2 mL/min. Methanol blanks were run in between samples. Statistical processing of the data was performed on the obtained peak areas.

The GC was coupled with an Agilent 5975B Inert MSD system using electron ionisation (EI). In full scan mode, masses were scanned from m/z 40 to 500. Mass spectra for recorded peaks were further evaluated using the NIST database (MS search programme Version 2.0, NIST, MSS ltd. Manchester, England).

**Results and Discussion**

**Extraction and separation of OGSR compounds**

The ability of the MonoTraps to extract a wide range of OGSR compounds was demonstrated by the extraction of a standard solution. Good chromatographic separation was achieved for most compounds (Figure 1). Only the dinitrotoluenes (4 and 5, and 6 and 7) are not baseline resolved, and carbazole (10) and methyl centralite (11) co-eluted. All compounds, however, could be identified using mass spectral data. A comparison of the results of all four MonoTrap™ types is shown in figure 2.

![Figure 1: Chromatogram silica disc extract of standard 1: 1. EPA, 2. Camphor, 3. 3-NT, 4. DMP, 5.2,6-DNT, 6. 2,3-DNT, 7. 2,4-DNT, 8. DEP, 9. DPA, 10. Carbazole, 11. MC, 12. DIBP, 13. DBP, 14. 2-NDPA, 15. 4-NDPA, 16. 2,4-DNDPA.](image)
The discs produced a higher peak area than the rods for all OGSR compounds, except for DBP that resulted in a greater peak area in the extract of the silica rod than carbon disc. This could be due to the limited surface contact the rods had with the sample; less than a quarter of the rod was in direct contact with the liquid standard during extraction, compared to half the disc. Submerging the rod to increase the surface area in contact with the sample resulted in very poor extraction, as indicated by the manufacturer’s information [8]. Both discs produced comparable peak areas for the first eight compounds shown in Figure 1. The latter eight compounds clearly produced higher peak areas using the silica disc. Despite the fact that the extraction using discs resulted in greater quantities detected than the rods for almost every compound, the silica MonoTraps extracted more compounds (a total of 16 and 14 compounds for the disc and rod respectively) than the carbon MonoTraps (a total of 13 and 11 compounds for the disc and rod respectively). These results demonstrate that the carbon rod is the least effective overall. The silica rod produced comparable values to the carbon rod but it enabled the extraction of more compounds than both of the carbon MonoTraps. The most effective extraction was achieved with the silica disc.

**Propellant powder analysis**

**Methanol extractions:** Figure 3 shows the chromatograms of the methanol extractions of the Hodgdon HP-38 (Figure 3a) and Magtech .223 (Figure 3b) propellant. The chromatogram of Hodgdon HP-38 shows all known OGSR compounds detected. Due to a greater difference in obtained peak areas, only the three major compounds are visible in the chromatogram of the Magtech propellant. The full list of compounds detected in both propellants, including the standard deviations of the average peak areas are shown in table 1.

The results show that both propellant powders have a different composition as a different combination of OGSR compounds has been detected, with varying relative abundances between the compounds. Good qualitative repeatability was obtained for each propellant, and a good chromatographic separation was observed for all detected compounds. Furthermore, low standard deviations were obtained for Hodgdon HP-38 propellant samples in general, however, the values for the Magtech propellant were lower and resulted in relatively high standard deviations. This is possibly due to the fact that the Magtech propellant appears to have a coating, which might inhibit the release of OGSR compounds [4].

Four compounds were detected in addition to the OGSR compounds that were included in the standard, namely nitroglycerin (NG) and ethyl centralite (EC) (both known OGSR compounds), akardite II (AKII) and N,N-diphenylformamide (DPF). AKII is a stabiliser that may be used as an additive in propellant powder and is thus also associated with OGSR materials [21-25]. DPF has only been mentioned in conjunction with OGSR in the literature on a few occasions, but its function in propellant powder was not reported [26,27].
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**Figure 3:** Methanol extractions of 100 mg propellant: a) Hodgdon HP-38, b) Magtech .223 (1. NG, 2. DPA, 3. EC, 4. DBP, 5. 2-NDPA, 6. AKII, 7. 4-NDPA).

**Table 1:** Mean peak areas (PA) and standard deviations for methanol extractions of two propellant powders.

<table>
<thead>
<tr>
<th>Compounds</th>
<th>RTs (min)</th>
<th>Hodgdon HP-38</th>
<th>Magtech .223</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean PA (n = 3)</td>
<td>Std dev (%)</td>
<td>Mean PA (n = 3)</td>
</tr>
<tr>
<td>NG</td>
<td>12.6</td>
<td>1.96E+08</td>
<td>4.46</td>
</tr>
<tr>
<td>DPA</td>
<td>18.3</td>
<td>1.15E+08</td>
<td>3.27</td>
</tr>
<tr>
<td>DIBP</td>
<td>23.8</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>EC</td>
<td>24.3</td>
<td>3.01E+06</td>
<td>5.63</td>
</tr>
<tr>
<td>DBP</td>
<td>25.6</td>
<td>9.06E+06</td>
<td>4.11</td>
</tr>
<tr>
<td>2-NDPA</td>
<td>25.7</td>
<td>1.95E+06</td>
<td>26.56</td>
</tr>
<tr>
<td>AKII</td>
<td>27.0</td>
<td>8.52E+05</td>
<td>8.46</td>
</tr>
<tr>
<td>4-NDPA</td>
<td>29.1</td>
<td>3.16E+06</td>
<td>6.35</td>
</tr>
<tr>
<td>DPF</td>
<td>21.8</td>
<td>1.14E+05</td>
<td>31.58</td>
</tr>
</tbody>
</table>

**MMSE extraction of propellant extracts:** Figure 4 shows the results of the extraction of Hodgdon HP-38 and Magtech .223 propellants.

The MonoTrap™ extractions produced similar results to the methanol extractions of the propellant powders, particularly for the major compounds in each powder. DPA and NG respectively were the major compounds extracted from Hodgdon HP-38 propellant. Their quantities were much higher than the amounts of the other four OGSR compounds present (Figure 4a), which were all extracted using the carbon disc and to a lesser degree with the silica rod. In contrast to the results obtained from the standard extraction, the silica disc extracted one compound, 4-NDPA, less than the carbon disc. The only OGSR compound that was not extracted using the MonoTraps, was AKII.

NG, DPA and DBP were clearly present as major compounds in the Magtech propellant (Figure 4b), and similar peak areas were obtained. The achieved peak areas of DIBP and 4-NDPA varied according to MonoTrap™ type, and 4-NDPA was not detected in the carbon rod extracts. EC was the only compound that was not detected. The overall amounts extracted from the Magtech propellant were lower for all MonoTrap™ types, which could be due to the coating of the propellant grains [4].

For both propellant powders, the most effective extraction was achieved using the carbon disc. When comparing the results achieved with the extraction of the carbon...
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It shows that the average peak areas obtained per compound are about one order of magnitude lower for the carbon disc. Despite this fact, only two compounds, one per powder, were not detected using this method. It should be noted that AKII was not included in the standard and thus it is unknown whether this compound is suitable for extraction with the developed MMSE method. EC may not have been detected in the .ăr Magtech propellant extracts as a result of the coating these powder grains appear to have. Lower overall yields were detected for the Magtech propellant compared to the Hodgdon propellant for both methanol extractions as well as MMSE.

**MMSE extraction of propellant-DCM desorption:** An initial attempt to improve the detection of these compounds was made by using DCM for the desorption of the MonoTraps following extraction of the Hodgdon propellant. This powder was selected as it contained the most OGSR compounds. The carbon rod was not tested due to its poor performance.

Desorption with DCM resulted in greater quantities detected for almost all compounds (Figure 5a). Subsequent methanol desorption, however, still resulted in relatively high yields of OGSR compounds (Figure 5b). Despite higher yields detected after DCM desorption, 4-NDPA was only detected in the carbon disc.

The improved desorption efficiency achieved with DCM resulted in the detection of peak areas of NG and EC obtained with the carbon disc that are comparable to the methanol extraction (Table 2). When combining the peak areas of the DCM and methanol fractions, the amounts for DPA and DBP are also comparable. The peak area values for 2-NDPA are approximately half of those produced by the methanol
extraction, and only 4-NDPA remains one order of magnitude below the methanol extraction value. These results suggest that the extraction of propellant powder using the carbon disc and sequential desorption is a viable nearly solvent-free alternative to methanol extraction in the analysis of the major propellant compounds and even some of the minor compounds. Further optimisation of the desorption method is ongoing and could potentially lead to improvement beyond that obtained for methanol extractions, given the fact that NG and EC already resulted in slightly better yields.

**Conclusion**

A method for the extraction of OGSR compounds from propellant powders, which enabled the effective separation and detection of a wide range of known OGSR compounds, has been presented. The major advantage of this method is that it requires
only 200 μL of solvent to perform the extraction, which is both environmentally and economically advantageous. The results have clearly demonstrated that extraction of propellant powder using the carbon disc is a viable alternative to methanol extraction. Similar amounts were detected for the major propellant compounds using both methods. Minor OGSR compounds were also extracted and it is suspected that further optimisation of the desorption method would significantly improve their detection. The results from the optimisation work have shown that multiple solvents may be required to desorb all compounds.

Further work by the authors is focusing on the improvement of the desorption method in order to ensure that all extracted compounds are recovered from the MonoTrap materials. Due to its capability to extract a wide range of OGSR compounds, including nitrotoluenes, diphenylamines and centralites, this method is expected to be applicable to a much wider range of ammunition types than was initially tested here.

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