Packed Capillary Liquid Chromatography
Electrospray Mass Spectrometry and Tandem
Mass Spectrometry of Hydrolysed HT and HQ

By:
P.A. D'Agostino, L.R. Provost and J.R. Hancock

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Suffield Report No. 691

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and Tandem Mass Spectrometry of Hydrolysed HT and HQ

by

P. A. D’Agostino, L. R. Provost and J. R. Hancock
Packed capillary column liquid chromatography (LC) - electrospray mass spectrometry (ESI-MS) and tandem mass spectrometry (MS/MS) were used to identify munitions grade mustard hydrolysis products, including five longer chain diols, two partial hydrolysis products and three ether/thioether macrocycles. All ESI-MS data were collected under collisionally activated dissociation (CAD) conditions optimized to facilitate acquisition of both molecular and product ion information that could be used for structural identification purposes. Interpretation of the ESI-MS data enabled characterization of the diols resulting from hydrolysis of all three principal sulfur vesicants, bis(2-chloroethyl)sulfide (mustard or H), bis(2-chloroethylthio)ethane (sesquimustard or Q) and bis[(2-chloroethylthio)ethyl] ether (T), as well as novel products not previously associated with sulfur vesicant hydrolysis. The reported ESI-MS data could prove valuable for the identification of thiodiglycol and other sulfur vesicant hydrolysis products in samples collected by the Canadian Forces, during base cleanup operations or in support of United Nations Chemical Weapons Convention inspections.
Executive Summary


Introduction: The Canadian Forces (CF) may be called on to perform peacekeeping or battlefield operations in regions of the world where there is a significant threat of chemical/biological (CB) warfare agent use. To operate effectively in these theatres the CF must be able to identify the CB agent used. Mass spectrometry (MS), is a powerful analytical technique for the identification of both known and unknown compounds and DRE Suffield is currently investigating this instrumental technique in fulfilment of CF detection and identification requirements.

Results: Packed capillary column liquid chromatography (LC) - electrospray mass spectrometry (ESI-MS) and tandem mass spectrometry (MS/MS) were used to identify munitions grade mustard hydrolysis products, including five longer chain diols, two partial hydrolysis products and three ether/thioether macrocycles. All ESI-MS data were collected under collisionally activated dissociation (CAD) conditions optimized to facilitate acquisition of both molecular and product ion information that could be used for structural identification purposes. Interpretation of the ESI-MS data enabled characterization of the diols resulting from hydrolysis of all three principal sulfur vesicants, bis(2-chloroethyl)sulfide (mustard or H), bis(2-chloroethylthio)ethane (sesquimustard or Q) and bis[(2-chloroethylthio)ethyl] ether (T), as well as novel products not previously associated with sulfur vesicant hydrolysis.

Significance of Results: The CF may be deployed in regions of the world where there is a significant threat of chemical/biological warfare agent use. Identification of the CB agent is of importance since the results of such analyses would contribute to the development of strategic and political positions regarding future Canadian military operations and would facilitate the dissemination of technical advice to in-theatre field commanders and medical personnel.

Future Goals: The reported ESI-MS data could prove valuable for the identification of thiodiglycol and other sulfur vesicant hydrolysis products in samples collected by the Canadian Forces, during base cleanup operations or in support of Chemical Weapons Convention challenge inspections. Presence of longer chain diols, partial hydrolysis products or ether/thioether macrocycles formed following hydrolysis of longer chain sulfur vesicants would augment thiodiglycol detection, and greatly strengthen the argument for prior mustard presence in suspect samples.

The methods developed will be evaluated on aqueous field samples as part of a collaborative Canadian/American exercise organized under TTCP Group E, Action Group 42.
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INTRODUCTION

The use of the chemical warfare agent, mustard in the Iran/Iraq war (1,2), and threat of chemical weapons use in the Persian Gulf War emphasize the need for specific methods to detect and identify sulfur vesicants and their degradation products. Retrospective analysis of samples contaminated with chemical warfare agents collected by the Canadian Forces (CF) will continue to be an important means for the verification of allegations of use claims, find application during base cleanup operations, and be a critical component during future inspections in support of the Chemical Weapons Convention.

Gas chromatography (GC) (3,4) and mass spectrometry (MS) (5-14) have been used extensively for the identification of mustard and mustard related compounds with capillary column GC-MS being the most commonly employed technique for the detection of these compounds in environmental (10,12,15), biological (16-20) and decontamination (21) samples. Mustard destruction by hydrolysis or natural weathering in the environment results in the formation of thiodiglycol (22,23), a non-toxic compound that may be easily handled. However, some munitions grade mustard formulations contain only 50% to 80% mustard with most of the remaining content being other sulfur vesicants (9) which decompose to other products. GC-MS under both electron impact (EI) and ammonia chemical ionization (CI) conditions has been used for the mass spectrometric characterization of several hydrolysis products of longer chain sulfur vesicants commonly found in munitions grade mustard (14). Molecular ion information, critical for the confirmation of these hydrolysis products, was generally absent during EI analyses and ammonia (CI) was used to obtain complementary molecular ion information for the compounds or their trimethylsilyl (TMS) derivatives.

The hydrolysis products of sulfur vesicants would generally be associated with aqueous samples and development of capillary electrophoresis (CE) or liquid chromatography-mass
spectrometry (LC-MS) for their detection and identification would reduce sample handling and derivatization requirements. Direct aqueous MS analysis methods for mustard hydrolysis products have made use of either loop injection MS or LC-MS. A notable exception involved the use of micellar electrokinetic chromatography with UV detection for the detection of thiodiglycol, 1,4-dithiane, 1,4-thioxane and 2,2'-sulfinyldiethanol (24). Thermospray (25), atmospheric pressure chemical ionization (APCI) (26) and electrospray (27) mass spectrometric interfaces have all been used to facilitate the introduction and ionization of mustard hydrolysis products. In these cases, the investigations focussed on the analysis of thiodiglycol, half-mustard, thiodiglycol sulfone and thiodiglycol sulfoxide, compounds commonly associated with the degradation of mustard.

Munitions grade mustard formulations often contain additional sulfur vesicants, including Q, T and longer chain sulfur variants. In some cases the munitions were purposely developed to contain multiple vesicants, with HT and HQ being two munitions grade materials containing relatively crude mixtures of H and T and, H and Q, respectively (9). Mass spectrometric characterization of the additional degradation products that would arise following hydrolysis would be valuable during the analysis samples collected during CF operations, during the analysis of samples collected during base cleanups and in support of Chemical Weapons Convention compliance monitoring.

A packed capillary LC-ESI-MS method was developed to detect and identify thiodiglycol and the novel hydrolysis products of the longer chain sulfur vesicants contained in munitions grade HT and HQ samples. ESI-MS provided ample molecular ion information and structurally important product ion information were generated by promoting collisionally activated dissociation (CAD) in the ESI interface.
EXPERIMENTAL

Samples

Samples of HT and HQ munitions grade mustard formulations (2 mL) were hydrolysed in a 125 mL erlenmeyer flask with 50 mL water at 50°C overnight. Acetone was added to each sample to solubilize the remaining oil and each sample was stirred overnight at 50°C. Both hydrolysed samples cleared and the excess water was evaporated leaving a pale yellow oil for each sample. The oils were then distilled in a Kugelrohr oven at 220°C at 0.1 mm. Hydrolysed HT and HQ samples were dissolved in water at the 1 mg/mL level prior to LC-ESI-MS analysis.

Instrumental

All electrospray mass spectra were acquired using a Micromass Autospec-Q tandem mass spectrometer (Manchester, UK) equipped with the Mark II electrospray interface. The electrospray needle was operated at 7.6 kV and ions were accelerated into the mass spectrometer at 4 kV. Sampling cone voltages of 20 to 90 volts were utilized. Nitrogen (Very Dry, Liquid Carbonic Inc., Scarborough, Ont., Canada) bath gas was introduced into the interface (80°C) at a flow rate of 400 L/hr. Nitrogen nebulizer gas was introduced at a flow rate of 14 L/hr. The electrospray interface was pumped with both a rotary and a turbomolecular pump, which enabled maintenance of a 4x10^{-4} and 7x10^{-6} Pa within the source and analyzer regions of the instrument, respectively. LC-ESI-MS data were acquired in the continuum mode by scanning the magnetic sector from 340 to 50 u (7 sec/decade) or 600 to 100 u (7 sec/decade) with a resolution of 1000 (10% valley definition). Three to five scans were typically averaged to enhance the signal-to-noise ratio.
The product spectra of m/z 167 for 8-chloro-6-oxa-3-thia-1-octanol and 14-chloro-6,12-dioxo-3,9-dithia-1-tetradecanol and the product spectrum of m/z 271 for 14-chloro-6,12-dioxo-3,9-dithia-1-tetradecanol were obtained during LC-ESI-MS/MS analysis under a variety of quadrupole collisional cell conditions in an effort to enhance product ion production. The quadrupole collisional cell energy was varied from 10 and 55 eV and the argon pressure was varied between 4.2 x 10^{-5} and 1.4 x 10^{-4} Pa (near the cell). In the end, increasing the cell pressure and/or energy was effective only in reducing the transmission efficiency. Product ion relative intensities remained low under all conditions investigated with relative intensities remaining being between 1% and 10% of the precursor ion intensity. The quadrupole was operated at unit resolution and scanned from 200 to 50 u at 4 sec/scan or 300 to 50 u at 5 sec/scan.

All LC separations were performed with an Applied Biosystems Model 140B dual syringe pump (Foster City, CA) equipped with a Zorbax 150 mm x 0.32 mm i.d. C_{18} SB (5 μm) packed fused-silica capillary column and a Rheodyne 8125 (Cotati, CA) injector with a 5 μL sample loop. The following solvent compositions were prepared for sample introduction: Solvent A (0.1% trifluoroacetic acid (TFA) in water) and Solvent B (0.1% TFA in acetonitrile/water, 95:5). Chromatographic separations were performed using a 1% to 75%B gradient over 30 minutes. In order to minimize dead volume effects and ensure reproducible mixing, the mobile phase was delivered at 200 μL/min and split prior to the injector such that the flow through the column was 5 μL/min.

RESULTS AND DISCUSSION

HQ and HT munitions grade mustard formulations, containing predominately mustard (H), and sesquimustard (Q), and mustard and bis[(2-chloroethylthio)ethyl] ether (T), respectively (9), were hydrolysed in an attempt to characterize the principal hydrolysis products associated
with these samples. Mustard hydrolysis results in the production of the partial hydrolysis product, hemisulfur mustard, which converts to the full hydrolysis product, thiodiglycol. Detection of thiodiglycol in suspect samples could provide evidence for the prior presence of mustard, but the presence of only thiodiglycol may not provide the conclusive evidence required to prove prior mustard presence. Detection of longer chain diols, partial hydrolysis products or ether/thioether macrocycles formed following hydrolysis of longer chain sulfur vesicants could provide much needed additional evidence to verify prior mustard presence.

Figure 1 illustrates typical total-ion-current chromatograms obtained following LC-ESI-MS analysis of the two hydrolysis samples. Thiodiglycol was the principal diol detected in both samples. Five additional diols related to Q, T and longer chain sulfur vesicants were also detected along with two partial hydrolysis products and three ether/thioether macrocycles (refer to Table I). The higher mass diols were separated by LC, but in general some peak tailing occurs during GC separation of diols. With more active or poorly conditioned capillary columns the quality of chromatography deteriorates significantly and detection of these hydrolysis products, including thiodiglycol, would be unreliable. The activity of GC columns becomes less critical during the analysis of the TMS derivatives of the hydrolysis products, as these compounds are less polar than the underderivatized compounds (14). Use of LC for the separation of sulfur vesicant hydrolysis products prior to MS characterization offers a number of potential advantages over GC. Most samples containing sulfur vesicant hydrolysis products would be aqueous in nature and these samples could then be analysed directly without the need for additional sample handling and/or derivatization. In addition, thermal degradation, a distinct possibility with increasing diol mass, would be minimized during LC-ESI-MS analysis.

Thiodiglycol has been recently analysed by ESI-MS following loop injection (27) and by LC-APCI-MS (26) with both authors reporting similar spectra. The mass spectra contained the protonated adduct at m/z 123 and a base ion at m/z 105 due to loss of H₂O from the (M+H)+ ion.
Additional molecular ion evidence was provided by the presence of a \((M+Na)^+\) adduct ion during ESI-MS analysis. ESI-MS conditions, in particular the sampling cone voltage, can have a considerable impact on mass spectrometric content. For this reason, thiodiglycol standards were analysed under a variety of conditions during LC-ESI-MS method development. Sampling cone voltage was varied between 20 and 90 volts with the effects tabulated in Table II. In general, the relative intensity of the product ion due to loss of \(H_2O\) increased with increasing sampling cone voltage, with 40 volts being selected as a reasonable compromise between molecular ion and product ion content. Concentration also has an effect on the nature of the ESI-MS data acquired. At higher concentrations (approximately 0.5 to 0.8 mg/mL), such as those encountered during the analysis of the HT and HQ hydrolysis samples, the thiodiglycol mass spectra contained slightly higher \((M+H)^+\) relative intensity and a significant \((2M+H)^+\) dimer at \(m/z\) 245. Minor adduct ions due to \((M+NH_4)^+, (M+Na)^+, (M+CH_3CN+H)^+\) and \((2M+Na)^+\) were also observed at \(m/z\) 140, \(m/z\) 145, \(m/z\) 164 and \(m/z\) 267, respectively. Figure 2a) illustrates typical ESI-MS data obtained for thiodiglycol at higher concentrations with a sampling cone voltage of 40 volts.

EI and ammonia CI-MS data have been reported for the two longer chain diols, 3,6-dithia-1,8-octanediol and 6-oxa-3,9-dithia-1,11-undecanediol, that result from the hydrolysis of Q and T, respectively (14). ESI-MS has not been previously used for the identification of these important hydrolysis products (Peak numbers 2 and 3) and was used to characterize these and three novel higher mass diols (Peak numbers 5, 6 and 7). Figures 1b), 1c), 2a), 2b) and 2c) illustrate typical ESI-MS data obtained for the longer chain diols: 3,6-dithia-1,8-octanediol; 6-oxa-3,9-dithia-1,11-undecanediol; 3,6,9-trithia-1,11-undecanediol; 6-oxa-3,9,12-trithia-1,14-tetradecanediol; and 6,12-dioxo-3,9,15-trithia-1,17-heptadecanediol (or 6,15-dioxa-3,9,12-trithia-1,17-heptadecanediol), respectively.

Diol ESI-MS data were rich in both molecular ion and product ion content, enabling structural identification of these hydrolysis products. The ESI-MS information obtained for 6-
Oxa-3,9-dithia-1,11-undecanediol, the hydrolysis product of T, was typical of that acquired. The mass spectrum contained a significant (M+H)+ ion at m/z 227, a (2M+H)+ ion at m/z 453, and a sodium adduct at m/z 249. A significant product ion due to loss of H₂O was observed at m/z 209 along with product ions at m/z 181, m/z 149 and m/z 105 due to (M+H–H₂O–C₃H₆)+, (M+H–H₂O–SC₂H₄)+ and (M+H–H₂O–SC₂H₄–OC₂H₄)+, respectively (Figure 1c). The protonated 1,4-thioxane ring at m/z 105 was significant for all the diols and represented the lowest mass product ion recorded above m/z 50.

Ions representing sequential loss of either SC₂H₄ or OC₂H₄ from higher mass ions enabled assignment of S and O positioning for all the diols except chromatographic peak number 7. In this case (Figure 2c) molecular mass was determined to be 330 u based on the presence of (M+H)+, (M+NH₄)+ and (M+Na)+ ions at m/z 331, m/z 348 and m/z 353, respectively. The ESI mass spectrum contained significant product ions at m/z 313, m/z 285, m/z 209, m/z 181, m/z 149 and m/z 105 that could be attributed to (M+H–H₂O)+, (M+H–H₂O–C₂H₄)+, (M+H–H₂O–SC₂H₄OC₂H₄)+, (M+H–H₂O–SC₂H₄OC₂H₄–C₂H₄)+, (M+H–H₂O–SC₂H₄OC₂H₄–SC₂H₄)+, (M+H–H₂O–SC₂H₄OC₂H₄–SC₂H₄–OC₂H₄)+, respectively. The order of S and O at positions 12 and 15 could not be determined since a product ion due to neutral loss of OC₂H₄ or SC₂H₄ from the (M+H–H₂O)+ was not recorded. In this case one of two possible hydrolysis products, 6,12-dioxo-3,9,15-trithia-1,17-heptadecanediol or 6,15-dioxo-3,9,12-trithia-1,17-heptadecanediol, was postulated. This ambiguity was not resolved through the use of LC-ESI-MS/MS.

Three ether/thioether macrocycles, two of which had been previously detected during GC-MS analysis of aqueous samples containing mustard hydrolysis products, were also identified during LC-ESI-MS analysis (10). The molecular mass for all three ether/thioether macrocycles (Figure 4) could be determined by the presence of a intense (M+H)+ ion and adduct ions due to (M+NH₄)+ and (M+Na)+. In one case (Figure 3a)), an adduct ion possibly due to
(M+Na+CH₃CN)+ was observed at m/z 272. The ring structures resisted fragmentation and few product ions were observed. Loss of H₂O from the (M+H)+ ion, a significant fragmentation pathway for diol, was absent and consistent with the ether/thioether macrocyclic structures postulated. All three ether/thioether macrocycles, 1,7-dioxoa-4,10-dithiacyclooctadecane, 1,7,13-trioxa-4,10,16-trithiacyclooctadecane (or another isomer) and 1,7-dioxoa-4,10,13-trithiacyclodecanedecane could be expected to form by ring closure (with loss of H₂O) of the detected diols, 6-oxa-3,9-dithia-1,11-undecanediol, 6-oxa-3,9,12-trithia-1,14-tetradecanediol and 6,12-dioxo-3,9,15-trithia-1,17-heptadecanediol (or 6,15-dioxo-3,9,12-trithia-1,17-heptadecanediol), respectively.

Two partial hydrolysis products, 8-chloro-6-oxa-3-thia-1-octanol and 14-chloro-6,12-dioxo-3,9-dithia-1-tetradecanol, the former having been previously characterized during GC-MS analysis (14), were identified following interpretation of acquired LC-ESI-MS and LC-ESI-MS/MS data. The ESI mass spectrum for 8-chloro-6-oxa-3-thia-1-octanol (Figure 5a)) contained characteristic chlorine isotopic clusters and (M+H)+ and (2M+H)+ ions at m/z 185 and 369, respectively. Product ions, due to loss of H₂O from the hydroxyl terminal of the (M+H)+ ion, at m/z 167 and the protonated 1,4-thioxane ion (m/z 105) were also observed. The 1,4-thioxane ion could arise from either loss of HOC₂H₄Cl from the (M+H)+ ion or loss of H₂O followed by loss of C₂H₃Cl. The latter possibility was discounted following LC-ESI-MS/MS analysis since the m/z 105 ion was not observed in the product spectrum of m/z 167. The only products of m/z 167 (Rel. Int. 100%) were at m/z 107 (Rel. Int. 5%), m/z 87 (Rel. Int. 2%) and m/z 63 (Rel. Int. 1%), indicating loss of SC₂H₄, HOC₂H₃Cl and 1,4-thioxane, respectively, from the (M+H⁻H₂O)+ precursor ion. For these product ions to be detected the m/z 167 must contain -OC₂H₄Cl at the chloro terminal in the partial hydrolysis product. Positions 3 and 6 must therefore contain S and O, respectively.

The ESI-MS data acquired for 14-chloro-6,12-dioxo-3,9-dithia-1-tetradecanol (Figure
5b)) contained (M+H)^+, (M+NH4)^+ and (M+Na)^+ ions at m/z 289, m/z 306 and m/z 311, respectively, that indicated a molecular mass of 288 u. The molecular ion adduct isotopic clusters were consistent with the presence of chlorine. Product ions at m/z 271, m/z 243, m/z 209, m/z 181, m/z 167 and m/z 105 were likely due to (M+H−H2O)^+, (M+H−H2O−C2H4)^+, (M+H−HOC2H4Cl−C2H4)^+, (M+H−H2O−SC2H4OC2H4)^+, (M+H−HOC2H4Cl−SC2H4OC2H4)^+, respectively. Interpretation of the ESI-MS data indicated a compound with following structure:

\[
\text{HOC}_2\text{H}_4−X_3−\text{C}_2\text{H}_4−X_6−\text{C}_2\text{H}_4−X_9−\text{C}_2\text{H}_4−\text{O}−\text{C}_2\text{H}_4\text{Cl},
\]

[where if \( X_3 \) is S then \( X_6 \) is O (or vice versa) based on loss of 1,4-thioxane from m/z 271, and if \( X_6 \) is O then \( X_9 \) is S (or vice versa) based on loss of 1,4-thioxane from m/z 209].

If the 1,4-thioxane ring at m/z 105 ion includes the O from the hydroxyl portion of the compound then \( X_3 \) must be a S. This would require \( X_6 \) to be O, which in turn means that \( X_9 \) would be S.

Evidence to support this assumption was provided during LC-ESI-MS/MS acquisition of the product ions of m/z 167 and m/z 271. The product spectrum for m/z 167 was virtually indistinguishable from that obtained during analysis of 8-chloro-6-oxa-3-thia-1-octanol, which was consistent with a \(-\text{C}_2\text{H}_4−\text{S}−\text{C}_2\text{H}_4−\text{O}−\text{C}_2\text{H}_4\text{Cl}\) structure at the chloro terminal. Position 9 (\( X_9 \)) would be occupied with an S atom. The product spectrum of m/z 271 (Rel. Int. 100%), contained a product ion at m/z 243 (Rel. Int. 8%) due to loss of C2H4 and three additional product ions at m/z 211, m/z 167 and m/z 107 (all with 1% to 2% Rel. Int.), due to sequential losses of SC2H4, OC2H4 and SC2H4 from the (M+H−H2O)^+ precursor ion, respectively. This would suggest the following structure, \text{HO}−\text{C}_2\text{H}_4−\text{S}−\text{C}_2\text{H}_4−\text{O}−\text{C}_2\text{H}_4−\text{SC}_2\text{H}_4−, at the hydroxyl terminal, where S occupies positions 3 (\( X_3 \)) and 9 (\( X_9 \)) and O occupies position 6 (\( X_6 \)). The structure of this partial hydrolysis product was determined to be 14-chloro-6,12-dioxa-3,9-dithia-1-tetradecanol based on this evidence.
CONCLUSIONS

Packed capillary column LC-ESI-MS and LC-ESI-MS/MS was used to characterize thiodiglycol and ten related longer chain diol, partial hydrolysis and ether/thioether macrocyclic compounds formed following hydrolysis of munitions grade HT and HQ samples. ESI-MS data were collected with a sampling cone voltage that promoted collisionally activated dissociation, with the resultant mass spectra being rich in both molecular and product ion information.

LC-ESI-MS has been demonstrated for higher mass sulfur vesicant hydrolysis product analysis, extending the range of analytical options available to the researcher confronted with the identification of chemical warfare agents or their decomposition products. This technique is an attractive alternative to GC-MS for the analysis of aqueous samples containing the hydrolysis products of sulfur vesicants since the aqueous samples may be analysed directly with little risk of thermal decomposition and without the need for additional sample handling or derivatization. The reported ESI-MS data could prove valuable for the verification of thiodiglycol and other sulfur vesicant hydrolysis products in samples collected by the CF during operations, in support of base cleanups or during Chemical Weapons Convention challenge inspections. Presence of longer chain diols, partial hydrolysis products or ether/thioether macrocycles formed following hydrolysis of longer chain sulfur vesicants would augment thiodiglycol detection, and greatly strengthen the argument for prior mustard presence.
REFERENCES


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**Table I:** Compounds Identified Following Packed Capillary LC-ESI-MS Analysis

<table>
<thead>
<tr>
<th>Peak No.</th>
<th>Mol. No.</th>
<th>Compound</th>
<th>Structure</th>
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<tbody>
<tr>
<td>1</td>
<td>122</td>
<td>Thiodiglycol</td>
<td>HO_2S_2OH</td>
</tr>
<tr>
<td>2</td>
<td>182</td>
<td>3,6-Dithia-1,8-octanediol</td>
<td>HO_2S_2OH</td>
</tr>
<tr>
<td>3</td>
<td>226</td>
<td>6-Oxa-3,9-dithia-1,11-undecanediol</td>
<td>HO_2S_2OH</td>
</tr>
<tr>
<td>4</td>
<td>184</td>
<td>8-Chloro-6-oxa-3-thia-1-octanol</td>
<td>HO_2S_2Cl</td>
</tr>
<tr>
<td>5</td>
<td>242</td>
<td>3,6,9-Trithia-1,11-undecanediol</td>
<td>HO_2S_2OH</td>
</tr>
<tr>
<td>6</td>
<td>286</td>
<td>6-Oxa-3,9,12-trithia-1,14-tetradecanediol</td>
<td>HO_2S_2OH</td>
</tr>
<tr>
<td>7</td>
<td>330</td>
<td>6,12-Dioxo-3,9,15-trithia-1,17-heptadecanediol or 6,15-Dioxo-3,9,12-trithia-1,17-heptadecanediol</td>
<td>HO_2S_2OH</td>
</tr>
<tr>
<td>8</td>
<td>208</td>
<td>1,7-Dioxo-4,10-dithiacyclododecane(^2)</td>
<td>One X=S, Other X=O</td>
</tr>
<tr>
<td>9</td>
<td>288</td>
<td>14-Chloro-6,12-dioxo-3,9-dithia-1-tetradecanol</td>
<td>HO_2S_2Cl</td>
</tr>
<tr>
<td>10</td>
<td>312</td>
<td>1,7,13-Trioxo-4,10,16-trithiacyclooctadecane or another isomer</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>268</td>
<td>1,7-Dioxo-4,10,13-trithiacyclopentadecane(^2)</td>
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</tr>
</tbody>
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1 Refer to Figure 1. 2 Probable isomer based on MS data obtained in a prior study (reference 14).
Table II: Typical m/z 123 and m/z 105 relative intensities obtained with different sampling cone voltages.

<table>
<thead>
<tr>
<th>Sampling Cone Voltage (volts)</th>
<th>Relative Intensity</th>
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<tr>
<td></td>
<td>m/z 123</td>
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<tr>
<td>20</td>
<td>83</td>
</tr>
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<td>40</td>
<td>40</td>
</tr>
<tr>
<td>60</td>
<td>31</td>
</tr>
<tr>
<td>90</td>
<td>7</td>
</tr>
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</table>
Figure 1: LC-ESI-MS total-ion-current (600 to 100 u) chromatograms obtained for a) HQ and b) HT hydrolysis samples (Compounds identified in Table I).
Figure 2: Typical ESI-MS data for a) thioglycol, b) 3,6-dithia-1,8-octanediol and c) 6-oxa-3,9-dithia-1,11-undecanediol obtained during LC-ESI-MS analysis of HT and HQ hydrolysis samples.
Figure 3: Typical ESI-MS data for a) 3,6,9-trithia-1,11-undecanediol, b) 6-oxa-3,9,12-trithia-1,14-tetradecanediol and c) 6,12-dioxo-3,9,15-trithia-1,17-heptadecanediol (or 6,15-dioxo-3,9,12-trithia-1,17-heptadecanediol) obtained during LC-ESI-MS analysis of HT and HQ hydrolysis samples.
Figure 4: Typical ESI-MS data for a) 1,7-dioxa-4,10-dithiacyclododecane, b) 1,7,13-trioxa-4,10,16-trithiacyclooctadecane (or another isomer) and c) 1,7-dioxa-4,10,13-trithiacyclopentadecane obtained during LC-ESI-MS analysis of HT and HQ hydrolysis samples.
Figure 5: Typical ESI-MS data for a) 8-chloro-6-oxa-3-thia-1-octanol and b) 14-chloro-6,12-dioxo-3,9-dithia-1-tetradecanol obtained during LC-ESI-MS analysis of HT and HQ hydrolysis samples.
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Packed capillary column liquid chromatography (LC)-electrospray mass spectrometry (ESI-MS) and tandem mass spectrometry (MS/MS) were used to identify munitions grade mustard hydrolysis products, including five longer chain diols, two partial hydrolysis products and three ether/thioether macrocycles. All ESI-MS data were collected under collisionally activated dissociation (CAD) conditions optimized to facilitate acquisition of both molecular and product ion information that could be useful for structural identification purposes. Interpretation of the ESI-MS data enabled characterization of the diols resulting from hydrolysis of all three principal sulfur vesicants, bis(2-chloroethyl)sulfide (mustard or H), bis(2-chloroethylthio)ethane (sesquimustard or Q) and bis[(2-chloroethylthio)ethyl]ether (T), as well as novel products not previously associated with sulfur vesicant hydrolysis. The reported ESI-MS data could prove valuable for the identification of thiodiglycol and other sulfur vesicant hydrolysis products in samples collected by the Canadian Forces, during base cleanup operations or in support of United Nations Chemical Weapons Convention inspections.

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