Detection and Classification of Explosive Compounds in Presence of Interferents Utilizing Laser Ion Mobility Spectrometry

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Abstract
Ion Mobility Spectrometers (IMS) are frequently used to screen airline passengers and their luggage for explosives and explosive by-products. Traditionally, in IMS technology different kinds of molecules are distinguished on account of their different ion mobilities in a weak electric field. In this paper we report on methods of resolution enhancement applicable to IMS drift time spectra. We find that a robust classification of trace explosives can be obtained by algorithmic resolution enhancement of the raw drift time spectra and by an additional parameterisation of the desorption function of the sampled explosives.

Introduction to IMS
Ion Mobility Spectrometry (IMS) is a trace-detection technology which is based on mobility measurement of charged particles and ions under atmospheric pressure conditions [1]. The charge is applied to the particles either by radioactive radiation, electron attachment, field-, photo- or electrospray ionisation. After ionisation, the particles or molecules undergo an acceleration process under the influence of a linear electrical drift field towards an ion collector electrode. Different kinds of molecules can be distinguished on account of their characteristic arrival times at the collector electrode, which depends on the molecular size and/or cross-sections. Important applications of IMS are in the fields of analytical chemistry [2] or explosives detection [3].

LIMS Hardware Instrumentation
The work presented here was performed with a laser ion mobility spectrometer (LIMS) prototype developed at EADS-IW [4]. Here, in contrast to classical IMS systems, a laser, instead of a radioactive $^{63}$Ni source is used for the ionisation of analyte, dopant and calibrant substances as this is a very selective and soft ionisation method [5]. The laser is a fixed wavelength UV laser based on a flash-lamp pumped Nd:YAG crystal with a wavelength of 266 nm. The studies were performed with a laser power of 20 mW and a pulse duration time of 5 ns. A schematic view of the instrument is shown in Fig. 1.

![Fig.1: Schematic design of the LIMS drift tube; (b) stacked-design drift tube assembled within the LIMS drift tube package [6].](image)

The drift tube has a drift length of 10.6 cm consisting of ceramic drift plates with integrated electrode patterns. An electrical field of $E = 248 \text{ V/cm}$ was applied for the measurements. We have used a rectangular drift tube design as it allows for a relatively long interaction path of the UV laser beam with the analyte molecules in the entrance region of the LIMS drift tube. Pressurized synthetic air with a flow rate of 100 sccm was used to generate the carrier and drift gas flows. Throughout the experiments, the inlet and drift tube temperatures were set at 150°C and 40°C, respectively. The desorber heater temperature was kept constant and varied between 80°C and 280°C, depending on the type of explosive.
Explosive standard solutions were procured from Restec company. The explosive compounds were solved in acetonitrile or methanol with a concentration of 1000 µg/1000 µl. For measurement, the explosive/interferent solutions were applied to sampling swabs. A syringe device was used for dosing the explosives in the 1 to 500 ng range. After preparation, the swabs were heated at a constant temperature for 60 s to allow the evaporation of the explosives and the beginning depletion of the swabs to be followed in sufficient detail.

Results
Spectral analysis of several explosive compounds was performed. The explosives investigated included substances with aromatic (2,4-DNT, 2,6-DNT, 2,4,6-TNT, DNB, TNB) and non-aromatic ring structures (Tetryl, HMX) as well as others with aliphatic backbone structures (PETN, GTN and TATP). All of them were detectable with LIMS with detection limits ranging from 0.5 ng (TNT) to approximately 20 ng (GTN).

In total, we investigated 21 interferent compounds for their LIMS response. These substances belonged to five different chemical groups which are all known to interfere with the detection of TNT, RDX & HMX, PETN & EGDN using conventional radioactive IMS [7]. The five groups are (1) acidic fog (nitrate phenols) from air pollution, (2) tobacco by-products, (3) synthetic fragrances used in perfumes and scents, (4) insecticides and fertilizers and (5) vasodilators in drugs for heart diseases. The 21 studied compounds were chosen because of their similarity to explosives with regard to hydrocarbon backbone structure and functional side groups. The suspected contaminants have at least one nitro functional group, with the exception of methyl parathion. The nitro functional group is the primary electron-withdrawing group present in most explosive compounds. Six of these compounds could not be detected with LIMS. Those that could be detected, exhibited detection limits in the µg range and thus were detectable with sensitivities at least ten times lower than those of the corresponding explosives. Additionally, these detectable interferents caused no problems regarding PLS classification as they differed in their spectral characteristics from the corresponding explosives.

Algorithmic System and Spectral Features
All spectra presented here were acquired in negative ion mode. The data acquisition frequency was 50 kHz. The scan period was set at 100 ms. Most of the results presented here were achieved by offline analysis with a Matlab based spectral analysis system.

Currently our resolution in raw spectra is around R ~ 30. The inherently low spectral resolution in ion mobility spectrometers - at least as compared to mass spectrometers - requires additional measures to classify explosives and to distinguish them from their corresponding interferents. This can be done using phenomenological methods or by extracting specific spectral features from the drift time spectra. In the first method, the entire spectra are used for classification, whereas in the second specific spectral features such as peak positions, peak shape etc. are compared with features from a database. The main substance-specific features in the drift time spectrum - the molecule-specific drift time and the corresponding K0 value - can be estimated up to a certain resolution which is often insufficient for classification. In order to ameliorate this situation we used a deconvolution algorithm proposed by Lucy [8]. With this deconvolution, our base instrumental resolution of R ~ 30 could be improved to R ~ 50. Fig. 2 shows a drift time spectrum of 2,4-Dinitrophenol and 2,4,6-TNT before and after deconvolution. The improvement in spectral resolution is immediately visible.

![Fig.2: The effect of deconvolution to the drift time spectrum of 2,4-DNP and 2,4,6-TNT. The broad feature at about 33 ms corresponds to the unresolved dimers of both compounds.](image-url)
Deconvolution also improves the statistical (phenomenological) classification with a Partial Least Square (PLS) algorithm [9]. Fig. 3 shows a series of PLS classification trained spectra of TNT and those of the traceable nitrated phenols from our interferent list.

![Figure 3: Demonstration of PLS classification with TNT against acidic fog components.](image)

The left template shows randomly chosen test spectra that are background reduced, calibrated and deconvoluted. These give input to the PLS classification which was trained with a disjunct training data set. The right template shows the corresponding classification vector. The 20 TNT test spectra are correctly classified except for two of them.

### Desorption Features

Even after an algorithmic resolution enhancement, the measurement of the peak positions is often insufficient to identify and distinguish different analytes. For example, TNT and TNB have a very similar chemical structure and almost identical $K_0$ values ($K_0,\text{TNT} = 1.54$, $K_0,\text{TNB} = 1.53$) and thus are not resolvable with our present spectral resolution. Desorption features may be used to distinguish substances with non-resolvable drift time values, which is demonstrated in Fig. 4.

![Figure 4: The normalised desorption histories of TNT and TNB. TNT and TNB can be distinguished on the basis of their desorption curves. Start and end of desorption, 60 s later, can be clearly seen.](image)

(a) Derivatives

There are several extractable parameters of the desorption functions available for classification. One of the seemingly obvious approaches, namely the use of first derivatives to gain distinguishing features, fails within the desired fast classification time of 10 seconds. This is due to the fact that during the first seconds of desorption, the rise in intensity shows a collective behaviour in all spectral components, caused by a change in the air flow and the availability of dopant and consecutive reactant ions. The
numerical estimation of curvature values and the necessary calculation of second derivatives appeared to be too noisy.

(b) Mathematical modelling
The direct parametrisation with a modeling function yields good results, but is computationally expensive and requires a non-linear optimisation algorithm that is demanding on real time applications due to the unknown number of iterations needed for convergence. This type of algorithm imperatively requires a set of start parameters that lie near the solution to guarantee convergence. We found that a fit with an asymmetric Gauss function with four parameters gives satisfactory results, as shown in Fig. 5. Desorption curves can be characterized with 4 parameters (features). We also tested fits with a log-normal function with inferior outcomes. Parameterisation of this kind needs also extended measurements (about 50 s) and is therefore not suitable for short measurement cycles.

\[
G_1 = \frac{1}{1 + \exp(- (x - \mu) / \sigma_1)}
\]
\[
G_2 = \frac{1}{1 + \exp(- (x - \mu) / \sigma_2)}
\]
\[
G(x, \sigma, \mu, f) = f \cdot (G_1 - G_2)
\]

![Fig. 5: Fit of an asymmetric Gauss function to the desorption curve of TNB.](image)

(c) Rise times and extrapolated desorption times
To avoid the computational and algorithmic demanding non-linear optimisation, a third method was tested. We estimated the total desorption time by linearly extrapolating the decaying slope of the logarithm of the desorption function. Unfortunately, this method also requires relatively long measurements of around 60 s to gain reasonable results for our set of explosive compounds. It may be more effective for fast desorbing substances. The estimated desorption time is defined as the time at which the intensity is equal to 0.05, e.g. 5 percent of the peak intensity. Fig. 6 shows the normalized desorption history of TNT and the linear extrapolation. The fall times are a promising feature to distinguish substances with identical drift times.

![Fig. 6: Linear extension of the log of the desorption curve of TNT.](image)
Conclusions
Spectral analysis of several major explosive compounds with aromatic (2,4-DNT, 2,6-DNT, 2,4,6-TNT, DNB, TNB), non-aromatic ring structures (Tetryl, HMX) and aliphatic hydrocarbon backbone structures (PETN, GTN and TATP) was performed. All of them could be detected with LIMS with detection limits ranging from 0.5 ng for TNT up to about 20ng for GTN.

The sensitivity of the LIMS traceable interferents was at least ten times lower than for explosives, with detection limits in the µg range. Six out of 21 interferent compounds could not be detected at all, and therefore did not pose any problem for explosive detection. Additionally, the remaining interferents caused no problems regarding PLS classification as their spectral characteristics are distinguishable from those of the explosives.

Algorithmic enhancement of the LIMS base instrumental resolution has been achieved. Algorithmic enhancement improved the resolution of drift time peaks which were not separated in the raw spectra. Deconvolution also improved the resolution in statistical classification (PLS).

Analysis of desorption curves allows for the extraction of additional spectral features. These can be used to distinguish substances with similar drift times, e.g. TNB and TNT. The most promising way of gaining such features was the estimation of the rise and decay times from the observed desorption curves.

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References: