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# Lifetimes and stabilities of familiar explosive molecular adduct complexes during ion mobility measurements<sup>†</sup>

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Trapped ion mobility spectrometry coupled to mass spectrometry (TIMS-MS) was utilized for the separation and identification of familiar explosives in complex mixtures. For the first time, molecular adduct complex lifetimes, relative stability, binding energies and candidate structures are reported for familiar explosives. Experimental and theoretical results showed that the adduct size and reactivity, complex binding energy and the explosive structure tailor the stability of the molecular adduct complex. The flexibility of TIMS to adapt the mobility separation as a function of the molecular adduct complex stability (*i.e.*, short or long IMS experiments/low or high IMS resolution) permits targeted measurements of explosives in complex mixtures with high confidence levels.

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## Introduction

Methods for the determination of trace levels of explosives and explosive related materials were developed rapidly and placed into service following several incidents in the 1980s involving catastrophic attacks with bombs on large civilian aircrafts.<sup>1,2</sup> The method chosen and distributed widely was ion mobility spectrometry (IMS) which was still in nascent stages of discovery concerning principles of ionization chemistry and best practices for measurements of ion mobility.<sup>3-7</sup> Nonetheless, embodiments of IMS were able to operate economically for onsite screening of hand-luggage at security check points of passengers and were distributed in airports worldwide. Measurements by the Explosive Trace Detectors (ETDs) with IMS depend upon the collection and vaporization of explosive residue, formation of molecular ions through chemical reactions in the gas phase, and their separation in a weak electric field as they drift in a bath gas.<sup>8</sup> A necessary requirement for an IMS measurement is that molecular ions formed from a substance should be distinctive and should have lifetimes sufficient to pass through the drift region with a characteristic mobility. This can be challenging with explosive molecular ions which may exhibit brief lifetimes and undergo reactions or decompositions either in the reaction region or in the drift region.<sup>9,10</sup> While sufficient understanding existed on the ionization chemistry and stability of ions in air at ambient pressure to justify the development of ETDs based on IMS, precise knowledge of the kinetics of ion decompositions and even the means to measure ion lifetimes in air at ambient pressure were developed only recently.

Explosive ions are formed in IMS based ETDs through chemical reactions where an explosive molecule, M, is electrostatically associated with a reactant or reagent ion, commonly Cl<sup>-</sup>, through ion-dipole or ion-induced dipole interactions.<sup>6,11,12</sup> The ions have thermal energies in the ion source of an IMS analyzer and ion and molecule associations are favorable without an energy barrier. Excess energy from the association can be lost by collisions, by reactions, and by dissociation of the explosives from the ions by the high collision frequency and abundance of small polar neutrals in the purified air of the IMS drift tube. Common reactions with explosives include hydrogen abstraction of protons that are acidic enough to be lost as HCl from an adduct [M + Cl]<sup>-</sup> and loss of NO<sub>3</sub><sup>-</sup> which appears to arise as a Cl<sup>-</sup> displacement reaction with a fracture in a weak carbon-oxygen bond.8 In other instances, the original adduct  $[M + Cl]^{-}$  has sufficient lifetime to pass through the drift region and reach the detector as an intact ion. In other instances, the ion may survive in the reaction region (~3 ms) and undergo reactions or dissociation in the drift region, appearing as a distortion in the baseline of the mobility spectrum.<sup>13</sup> Methods were described to extract

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kinetic information from baseline distortions and refined methods developed recently as a kinetic IMS instrument to obtain rate data for specific ions over a range of temperatures without interference from unwanted ion neutral interactions.<sup>14</sup> Reactions including the loss of NO<sub>3</sub><sup>-</sup> and Cl<sup>-</sup> from thermalized ions require energy which has been measured by the kinetic IMS method as 60-89 kJ mol<sup>-1</sup> and match favorably with *ab initio* calculations.<sup>9,10</sup> These reactions are dependent not only on temperature and moisture but also on the precursor ion. While commercial ETDs produce Cl<sup>-</sup> by dissociative electron capture in a beta emitter source, electrospray ionization (ESI) sources afford flexibility and convenience to form adducts from other anions by spiking the ESI starting solution with various salts.<sup>15,16</sup> For example, the measurement of multiple adduct forms of a targeted compound increases the identification confidence while reduces the probability of having interference from the sample matrix.

With the recent development of trapped ion mobility spectrometry (TIMS), higher mobility resolution and the capability to interrogate and simultaneously measure the molecular ionneutral collision cross section (CCS) as a function of time after the molecular ion formation has permitted kinetic studies of molecular ion-neutral bath gas interactions at the millisecond to second time scale.<sup>17-22</sup> In the current study, the unique potential of TIMS to hold ions while interacting with bath gas molecules ("TIMS" thermostat) is utilized to study the stability and dissociation kinetics of familiar explosives with different adduct forms at the level of individual molecules. In particular, ion-neutral collision cross sections (CCS) are measured using TIMS for a series of familiar explosive standards in nitrogen as a bath gas and compared with traditional drift tube IMS measurements and theoretical calculations. TIMS-MS capability to separate and identify explosives from complex samples is also demonstrated. In addition, for the first time, molecular ion stability and lifetimes are reported for a series of familiar explosive molecular adducts.

## **Experimental section**

#### Chemicals

Individual standards of 2-methyl-1,3,5-trinitrobenzene (TNT), 1,3,5-trinitroperhydro-1,3,5-triazine (RDX), 3-nitrooxy-2,2-bis-(nitrooxymethyl)propyl nitrate (PETN) and octahydro-1,3,5,7tetranitro-1,3,5,7-tetrazocine (HMX) were obtained from Accu-Standard (New Haven, CT) and used as received. Ammonium chloride, ammonium formate, ammonium acetate and ammonium nitrate salts and chromatography grade water, methanol and acetonitrile solvents were obtained from Fisher Scientific (Suwanee, GA) and used as received. TNT, RDX and HMX were dissolved in 1:1 water: methanol : acetonitrile v/v ratio to a final concentration of 1  $\mu$ M. Each ammonium salt containing solution was prepared separately and added to each explosive solution to a final concentration of 10 mM of ammonium salt. An electrospray ionization source (ESI, Bruker Daltonics Inc., MA) was used for all analyses in negative ion mode. The sample purity was confirmed with sub ppm mass accuracy for each standard using ultra-high resolution mass spectrometry with a Solarix 7 T FT-ICR MS mass spectrometer (Bruker Daltonics Inc., Billerica, MA). A complex mixture of TNT + cappuccino was prepared by doping a standard cappuccino coffee solution with the TNT standard (1  $\mu$ M) to 100:1 v/v ratio; the complex mixture sample was diluted in 1:1:1 water: methanol: acetonitrile v/v ratio to a final concentration of 10 nM of the TNT standard.

#### **TIMS-MS operation**

Details regarding the TIMS operation and specifics compared to traditional IMS can be found elsewhere.<sup>17,19,21,23,24</sup> Briefly, mobility separation in TIMS is based on holding the ions stationary using an electric field against a moving gas. The separation in a TIMS device can be described by the center of the mass frame using the same principles as in a conventional IMS drift tube.<sup>25</sup> In traditional drift tube cells, mobility separation is related to the number of ion-neutral collisions (or drift time); analogously, the mobility separation in a TIMS device is related to the bath gas drift velocity, ion confinement and ion elution parameters. The mobility, *K*, of an ion in a TIMS cell is described by:

$$K = \frac{\nu_{\rm g}}{E} = \frac{A}{(V_{\rm elution} - V_{\rm base})} \tag{1}$$

where  $v_g$ , *E*,  $V_{elution}$  and  $V_{base}$  are the velocity of the gas, applied electric field, elution and base voltages, respectively. The constant *A* was determined using the reported mobilities of explosives.<sup>8,26</sup> In TIMS operation, multiple geometric isomers/conformers can be trapped simultaneously at different *E* values resulting from a voltage gradient applied across the IMS tunnel. After thermalization, trapped species are eluted by decreasing the electric field in stepwise decrements (referred to as the "ramp"). Each mobility-separated isomer/conformer eluting from the TIMS cell can be described by a characteristic voltage difference (*i.e.*,  $V_{elution} - V_{base}$ ). Eluted ions are then mass analyzed and detected by using a maXis impact Q-ToF mass spectrometer (Bruker Daltonics Inc., Billerica, MA).

In a TIMS device, the total analysis time can be described as:

Total IMS time = 
$$T_{\text{trap}} + (V_{\text{elution}}/V_{\text{ramp}}) \times T_{\text{ramp}} + \text{ToF}$$
  
=  $T_{\text{o}} + (V_{\text{elut}}/V_{\text{ramp}}) \times T_{\text{ramp}}$  (2)

where  $T_{\rm trap}$  is the thermalization/trapping time, ToF is the time after the mobility separation, and  $V_{\rm ramp}$  and  $T_{\rm ramp}$  are the voltage range and time required to vary the electric field, respectively. The elution voltage can be experimentally determined by varying the ramp time for a constant ramp voltage. This procedure also determines the time ions spend outside the separation region  $T_{\rm o}$  (*e.g.*, ion trapping and time-of-flight).

The TIMS funnel is controlled using in-house software, written in National Instruments Lab VIEW, and synchronized

with the maXis Impact Q-ToF acquisition program.<sup>17,23</sup> TIMS separation was performed using nitrogen as a bath gas at *ca*. 300 K and typical pressures at the entrance and back regions of the TIMS analyzer were  $P_1 = 2.6$  and  $P_2 = 1.0$  mbar, respectively (see more details in ref. 19). The same RF (2040 kHz and 200–350 V<sub>pp</sub>) was applied to all electrodes including the entrance funnel, the mobility separating section, and the exit funnel. At all times, the axial electric field was kept under the low field limit (E/p < 10 V cm<sup>-1</sup> torr<sup>-1</sup>) throughout the TIMS and no significant ion heating is produced by the RF confinement.

Mobility values (*K*) were correlated with CCS ( $\Omega$ ) using the equation:

$$\Omega = \frac{(18\pi)^{1/2}}{16} \frac{ze}{\left(k_{\rm B}T\right)^{1/2}} \left[\frac{1}{m_{\rm I}} + \frac{1}{m_{b}}\right]^{1/2} \frac{1}{K} \frac{760}{P} \frac{T}{273.15} \frac{1}{N^{*}}$$
(3)

where *ze* is the charge of the ion,  $k_{\rm B}$  is the Boltzmann constant,  $N^*$  is the number density under standard temperature and pressure conditions, and  $m_{\rm I}$  and  $m_{\rm b}$  refer to the masses of the ion and bath gas, respectively.<sup>25</sup>

The analysis of the molecular adduct decomposition was considered as a first order reaction. The molecular adduct abundance at a given time is defined by the equation:

$$I = I_0 \exp(-kt) \tag{4}$$

where *k* is the decomposition rate ( $k = 1/t_d$ ),  $t_d$  is the lifetime of the molecular adduct complex, and  $I_o$  is the initial abundance.

#### Theoretical calculations

Geometries and binding energies of candidate structures were optimized at the DFT/B3LYP/6-31+g(d) level using Gaussian 09 software.<sup>27</sup> Vibrational frequencies were calculated to guarantee that the optimized structures correspond to a real minima in the energy space, and zero-point energy corrections were applied to calculate the relative stability. Partial atomic charges were calculated using the Merz-Singh-Kollman scheme constrained to the molecular dipole moment.<sup>28,29</sup> Theoretical ion-neutral collision cross sections were calculated using the trajectory method (TM) in MOBCAL version for nitrogen<sup>30,31</sup> with a bath gas at *ca.* 300 K. It should be noted that the MOBCAL version for nitrogen was used assuming the similarity of the molecules to those used to develop the Lennard-Jones potential at 300 K in ref. 30 and 31; for other molecules, alternative methods may be more accurate (see ref. 32). All optimized geometries and MOBCAL input files can be found in the ESI.†

## **Results and discussion**

A prerequisite for good analytical IMS performance is the ability to separate and identify molecular species with high reproducibility. The IMS resolution of hand held IMS instruments (*e.g.*, ETDs) is commonly  $R_{IMS} = 20$  or below; however, laboratory research IMS instruments using drift tube IMS

designs can routinely reach  $R_{IMS} = 80-100.^{33-37}$  Recently, we have reported the advantages of TIMS technology to achieve higher mobility resolution  $(R_{IMS} = 150-250)$ .<sup>19,20</sup> Different from other IMS forms (e.g., field asymmetric IMS,<sup>38</sup> differential mobility spectrometer,<sup>39-41</sup> segmented quadrupole drift cell,<sup>42</sup> cylindrical drift tubes,<sup>43</sup> and traveling wave ion guide<sup>44</sup>), TIMS mobility resolution varies with the size, mass and charge of the molecule of interest; that is, different trapping conditions are required to compensate for molecular ion diffusion and for coulombic repulsion of molecular ions during the trapping and elution steps. In practice, this translates into a lower mobility resolution for high mobility and low mass-to-charge ratio species when compared to previously reported values during fast TIMS mobility scans (see Fig. 1 for common explosives). One alternative to increase the TIMS mobility resolution is to reduce the ramp speed which results in higher IMS resolution. For example, a high mobility resolution of  $R_{\text{TIMS}} >$ 120 can be achieved for the analysis of explosives which results in a 3-5 fold increase in resolution when compared to commercially available ETD instruments.

The high mobility resolution of a TIMS device provides great potential for the analysis of explosives in complex mix-



Fig. 1 Typical IMS projection spectra for (a) TNT and (b) PETN using ESI-TIMS-MS.



**Fig. 2** (a) 2D IMS-MS contour plot of a complex mixture (cappuccino + TNT); (b) inset in the m/z = 224-229 range, and (c) IMS projection plots of m/z = 226 for the complex mixture and a TNT standard.

tures when coupled to mass spectrometry (see Fig. 2). That is, the ability to separate common interferences, to increase peak capacity, and to reduce chemical noise using orthogonal separations permits better identification of explosives using accurate CCS (<5% accuracy using external calibration) and m/z

measurements (in the example presented, mass resolution was  $R_{\text{TOF}}$  = 30–40k). Nevertheless, when internal calibrants are used for CCS determination in a TIMS device over a narrower CCS range the accuracy is better than a few percent. When compared to other hyphenated MS techniques for the analysis of familiar explosives,<sup>15,28,45–51</sup> TIMS-MS provides higher throughput, dynamic range and reduced analysis time. While an increase in peak capacity is observed during TIMS-MS analysis, the most challenging part involves the identification of compounds from the 2D IMS-MS plots. If standards are available for the a priori selected target (see Fig. 2c), the identification can be achieved by direct correlation of the IMS and MS data. It should be noted that additional IMS-MS/MS can further increase the identification capabilities. Another alternative is the coupling of TIMS to ultrahigh resolution MS analyzers (see the example in ref. 52); however, it should be noted that TIMS-TOF-MS operates with much shorter acquisition times.

While TIMS-MS provides high confidence for the analysis of common explosives, one way to further improve the confidence level is to simultaneously measure different molecular adducts.<sup>15,16</sup> That is, each measured molecular adduct form provides a two point identification (*i.e.*, CCS and m/z). Multiple molecular adducts can be formed during ESI of explosives by spiking the ESI starting solution with various salts (see the example in Fig. 3). In practice, this translates into a CCS and m/z shift for each adduct form, thus increasing the confidence level (see more details in Table 1). Compound identification from complex mixtures is typically challenged by the existence of molecular interferences in the IMS or MS domain. The use of multiple IMS and MS identification points from multiple adduct forms of a targeted compound increases the identification confidence while reducing the probability interferences from the sample matrix. In addition, since TIMS permits the measurement of CCS using first principles, the identification can be complemented with theoretical calculations; this approach can be very useful for the case of molecular adduct complexes that can exist as multiple conformations in the gas phase (see the example in ref. 52). Table 1 summarizes theoretical and experimental CCS of all the molecular adduct complexes observed (all structures are provided in the ESI,† see Fig. S1<sup> $\dagger$ </sup>). A K<sub>o</sub> absolute error of less than 0.5% was observed in TIMS replicate measurements. Close inspection shows that a good agreement is observed between the theoretical and TIMS experimental values (<5% difference). The largest difference between K<sub>o</sub> values measured by using TIMS and literature values can be attributed to the sample introduction (see ref. 8). For example, K<sub>o</sub> values of 1.45, 1.48 and 1.54 have been reported for TNT  $[M - H]^-$  for sample introduction by desorption, ESI, and vapor (membrane), respectively.

The measurement of multiple adduct forms of familiar explosives depends on the probability of forming the molecular adduct complex and its relative stability. During ESI ion formation, changes in the relative salt content can be used to preferentially target the formation of an adduct form as a way to avoid potential CCS and/or m/z interference. In addition,



**Fig. 3** Typical TIMS spectra for (a) HMX and (b) PETN as a function of the adduct form. Distances between the molecules and the adducts are shown.

the relative stability of the molecular ion complex during the TIMS-MS measurements will provide the best adduct candidate for effective detection. Explosives present different affinities for each molecular adduct complex. For example, TNT presents very low affinity to form a molecular adduct; however, HMX, RDX and PETN form a variety of complexes  $(e.g., [M + Cl]^{-}, [M + HCOOH - H]^{-}, [M + CH_3COOH - H]^{-}$ and  $[M + NO_3]^-$ ). Inspection of the molecular adduct lifetimes shows that the larger the adduct size the lower the complex stability (see Fig. 4 and Table 2). For example, PETN [M]<sup>-</sup> shows the largest lifetime (400 ms) when compared to the other molecular adducts  $[M + Cl]^-$  (85 ms),  $[M + HCOOC - H]^-$ (92 ms), and  $[M + NO_3]^-$  (85 ms). Moreover, the explosive structure influences the probability of forming molecular adducts. For example, HMX presents larger binding energy and longer lifetimes  $(\sim 3-4\times)$  for the molecular adduct forms when compared with RDX and PETN (see Table 2). Inspection of the HMX complex optimized geometries shows that the multiple coordination between the HMX molecule and the adduct favors the stability of the complex. That is, if the charge is protected, TIMS-MS experiments show no ion loss in up to two seconds of trapping (e.g.,  $m/z = 301 \text{ C}_3\text{N}_3(\text{CF}_3)_3 \text{ [M]}^-$  from the Agilent tuning mix,<sup>53</sup> Fig. 4a). Moreover, if the charge is exposed (e.g., TNT [M - H]<sup>-</sup>), ions can undergo charge neutralization via charge transfer with the bath gas molecules (e.g., proton transfer). In the case of the molecular adduct, the reactive nature of the adduct ion and the probability to collide with a bath gas molecule increase the chances for decomposition of the molecular adduct complex by transferring the charge carrying adduct to a bath gas molecule (e.g., decomposition by adduct transfer). That is, TIMS-MS experiments suggest that the collision rate and bath gas composition (or impurities) can be the defining factors for the observation of the molecular adduct complex. Although we cannot establish the mechanism for the molecular adduct complex decomposition, preliminary results suggest that the electrostatic nature of the complex can be lost by the interaction with a third partner (bath gas molecule), a short life complex formation, followed by the detachment of the adduct from the molecular complex.

During TIMS analysis, a short analysis time will increase the probability to observe a molecular adduct complex; however, a slower electric field ramp speed will provide higher mobility separations but longer measurement times. That is, high resolution TIMS separation can be limited by the molecular adduct complex lifetime and initial population (or abundance). Moreover, this observation can be extrapolated to the case of traditional drift tube IMS measurements in that long drift times will reduce the probability to observe a molecular complex ion form. In any IMS separation, since the number of collision defines the mobility resolution, the probability to observe a molecular adduct complex at high IMS resolution is limited by its stability and the composition of the bath gas.

## Conclusions

The analytical capabilities of TIMS-MS for the separation and identification of familiar explosives have been demonstrated. In particular, a three to five fold increase in mobility resolution

	Ionic form	m/z	TIMS experimental		_	
Compound			$K_{\rm o} \left( {\rm cm}^2  {\rm V}^{-1}  {\rm s}^{-1} \right)$	$CCS(Å^2)$	Reported $K_{\rm o}$ (cm <sup>2</sup> V <sup>-1</sup> s <sup>-1</sup> )	Theoretical CCS (Å <sup>2</sup> )
TNT	[M – H] <sup>–</sup>	226.010	1.48	143	1.48	136
$RDX + NH_4Cl$	$[M + Cl]^{-}$	257.003	1.44	147	1.44*	149
$RDX + NH_4NO_3$	$[M + NO_3]^-$	284.022	1.36	154	1.35*	152
$HMX + NH_4Cl$	$\left[M + Cl\right]^{-2}$	331.015	1.29	161	1.25	162
$HMX + HCO_2$	$[M + HCOOH - H]^{-}$	341.044	1.28	162	_	161
HMX + $NH_4C_2H_3O_2$	$[M + CH_3COOH - H]^-$	355.059	1.23	169	_	169
$HMX + NH_4NO_3$	$[M + NO_3]^-$	358.034	1.23	167	_	165
PETN	[M*]	316.013	1.37	152	_	151
$PETN + NH_4Cl$	$[M + Cl]^-$	350.982	1.17	178	1.20	182
$PETN + HCO_2$	$[M + HCOOH - H]^{-}$	361.011	1.14	182	_	179
$PETN + NH_4NO_3$	$[M + NO_3]^-$	378.001	1.11	187	1.14	188

**Table 1** Experimental (TIMS), literature,<sup>8,26</sup> and theoretical mobility values of molecular adduct complexes from familiar explosives. Literature values used in the TIMS calibration are denoted with \*. A  $K_o$  error of less than 0.5% was observed in the TIMS replicate measurements



**Fig. 4** Relative abundance of familiar explosive molecular ions as a function of the trapping time: (a) TNT, (b) HMX, (c) RDX and (d) PETN. Note that for  $m/z = 301 \text{ C}_3 \text{N}_3 (\text{CF}_3)_3 \text{ [M]}^-$  no ion loss in up to 2 seconds of trapping is observed (a).

was observed for the TIMS analyzer when compared with commercial ETD IMS devices. The use of molecular adduct complexes increases the confidence level and permits the identification of familiar explosives using first principle CCS and m/z measurements. For the first time, lifetimes, relative stability, binding energies and candidate structures are reported for molecular adducts of familiar explosives. Inspection of the molecular adduct interaction with the residual bath gas showed three major trends: (i) molecular ions (*e.g.*,  $[M - H]^-$ ) are more stable than their molecular adduct counterparts (*e.g.*,  $[M + CI]^-$ ,  $[M + HCOOH - H]^-$ ,  $[M + CH_3COOH - H]^-$  and  $[M + NO_3]^-$ ), (ii) the stability of the chloride and nitrate

Compound	Ionic form	$t_{\rm d}~({\rm ms})$	k (s <sup>-1</sup> )	Binding energy (kcal mol <sup>-1</sup> )	Distance molecule- adduct (Å)
TNT	[M – H] <sup>–</sup>	520	1.92	_	
$RDX + NH_4Cl$	$[M + Cl]^{-}$	110	9.09	36.93	2.52
$RDX + NH_4NO_3$	$[M + NO_3]^-$	65	15.38	33.73	1.97
$HMX + NH_4Cl$	$[M + Cl]^{-1}$	296	3.38	47.47	2.52
$HMX + HCO_2$	[M + HCOOH − H] <sup>−</sup>	250	4.00	50.25	2.14
$HMX + NH_4C_2H_3O_2$	$[M + CH_3COOH - H]^-$	195	5.13	52.61	2.08
$HMX + NH_4NO_3$	$[M + NO_3]^-$	322	3.11	41.06	2.13
PETN	[M] <sup>-</sup>	400	2.50	_	
$PETN + NH_4Cl$	$[M + Cl]^{-}$	85	11.76	30.95	2.39
$PETN + HCO_2$	[M + HCOOH − H] <sup>−</sup>	92	10.87	33.57	2.10
$PETN + NH_4NO_3$	$[M + NO_3]^-$	85	11.76	28.50	2.28

Table 2 Lifetime ( $t_d$ ), decomposition constant (k), absolute binding energy and molecular-adduct distances of molecular adduct complexes from familiar explosives

adducts is higher than the formate and acetate adducts, and (iii) HMX forms the most stable molecular adduct complexes when compared with RDX and PETN. We interpret this relative stability as a consequence of the probability of decomposition and of charge exchange with the bath gas of the molecular adduct complexes. That is, the adduct size and reactivity, complex binding energy and the explosive structure define the stability of the molecular adduct complex. The TIMS flexibility to modify the mobility separation as a function of the molecular adduct stability (*i.e.*, short or long IMS experiments/low or high IMS resolution) permits targeted measurements of explosives in complex mixtures.

## Conflict of interest

The authors declare no competing financial interest.

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