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# Drift Tube Ion Mobility: How to Reconstruct Collision Cross Section Distributions from Arrival Time Distributions?

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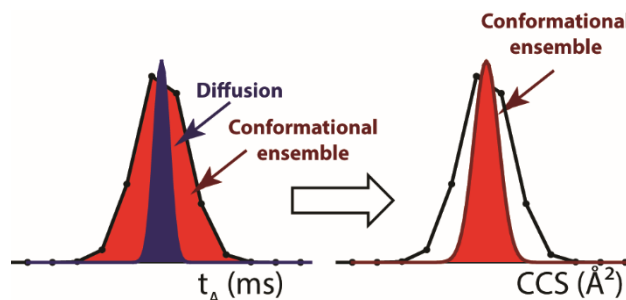
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**Abstract:** Ion mobility spectrometry allows one to determine ion collision cross sections, which are related to ion size and shape.

Collision cross sections (CCS) are usually discussed based on the peak center, yet the width of each peak contains further informa

tion on the distribution of collision cross sections of each conformational ensemble. Here, we analyze how to convert arrival time distributions (ATD) to CCS distributions (CCSD). With a calibration curve taking into account the CCS dependence of the time spent outside the mobility region, one can reconstruct CCS distributions with correct peak center values. However, the peak widths are incorrectly rendered because ion diffusion, which affects the peak width in the time domain, is irrelevant to collision cross sections. For drift tube ion mobility, we describe a new method, coined “FWHMstep”, using step-field experiment and processing the peak’s full width at half maximum to reconstruct CCSDs. The width of the CCS distributions helps to characterize the analyte’s structural heterogeneity, and/or its flexibility (i.e., the variety of ways the analyte ions can rearrange following electrospray into kinetically stable gas-phase conformations).



## INTRODUCTION

Ion mobility separates ions based on their electrophoretic mobility in a buffer gas under the influence of an electric field.<sup>1-3</sup> The mobility is proportional to the ion size, but also ion shape, and in that respect ion mobility spectrometry (IMS) is complementary to mass spectrometry. Recent technological advances have significantly improved the peak separation in ion mobility spectrometry. Resolving power values  $> 250$  has been achieved in linear drift tubes,<sup>4</sup>  $> 300$  in cyclic drift tubes,<sup>5</sup> up to 500 in field-assymmetric waveform IMS (FAIMS),<sup>6</sup> and a nominal resolution of  $\sim 340$  was obtained with traveling wave structures for lossless ion manipulation (SLIM).<sup>7</sup> The range of applications exploiting ion mobility as an additional dimension of separation is constantly growing, particularly towards clinical,<sup>8</sup> biological and environmental applications.<sup>9</sup>

A common instrumental configuration involves pulsing an ion packet into the mobility device, and measuring the time ions take to reach a detector situated after the mobility cell. The primary output is thus an arrival time distribution (ATD). Beyond separation and identification based on arrival time, ion mobility spectrometry is also increasingly used to characterize ion structural ensembles, via the collision cross section (CCS).<sup>1,10-14</sup> The CCS indicates the extent to which the ions are slowed down by collisions, and can be related to three-dimensional structural models by computation. Note that rigorously, one should call it the momentum transfer collision integral and use the symbol  $\Omega$ , but using the acronym CCS for “collision cross section” is however becoming common practice, and we will use it here for readability.

When the ATD peak width corresponds to the diffusion limit of the instrument, one generally assumes that a single structure is adopted, and it makes sense to discuss only the CCS of the center of the distribution. However, in electrospray IMS of large analytes such as peptides, proteins, and their noncovalent assemblies, peaks significantly broader than the diffusion limit are frequently observed. For example, large protein complexes can display peak widths up to 10% of the peak maximum.<sup>15</sup> Slicing the broad arrival time distributions by high-resolution IMS-IMS demonstrates that the underlying conformational ensemble actually consists of many distinct conformations having a continuum of CCS values.<sup>16,17</sup> The width of the distributions thus conveys information about the structural heterogeneity of each ensemble.<sup>18</sup>

The peak width can be represented in different ways. For example, for proteins, Bush's group presents the CCS of the intensity-weighted center of the distribution (50% of the cumulative distribution function), and error bars are drawn using the CCS values at 10% and 90% of the cumulative distribution function.<sup>19</sup> This simple representation however masks any particular shape of the arrival time distribution. Presenting ion mobility spectrometry results as CCS distributions (CCSD) is attractive when discussing shape, because it allows one to visually compare various molecular systems, compare results obtained on different instrumental platforms, or compare experimental and theoretical data. But despite graphs representing ion mobility results in the form of CCS distributions is becoming common in the literature, the procedure to change variables from arrival time to collision cross section is often left undiscussed. Here we clarify the different ways to change variables, and propose a new method, based on step-field drift tube experiments, to reconstruct CCS distributions that better reflect the ion conformational ensembles.

## EXPERIMENTAL METHODS

All ion mobility experiments were carried out in an Agilent 6560 drift tube IMS-Q-TOF instrument (Agilent Technologies, Santa Clara (CA), USA), with helium in the drift tube ( $p = 3.89$  Torr, measured using an Agilent CDG-500 capacitance diaphragm gauge) and nitrogen in the collision cell ( $p = 2.78 \times 10^{-5}$  Torr, measured using the instrument's Granville Phillips compact ion gauge). Arrival time distributions were extracted using IM-MS Browser B.07.01 and processed with Sigmaplot 12.5. The violin plots were generated using a MatLab script (see Supporting Information).

Samples included a mixture of hexakis-(fluoroalkoxy)phosphazines (TuneMix, Agilent Technologies), bradykinin (B3259,  $\geq 98\%$ , Sigma-Aldrich, Saint Quentin-Fallavier, France), bovine ubiquitin (U6253,  $\geq 98\%$ , Sigma-Aldrich), equine cytochrome *c* (C7752, assay  $\geq 95\%$ , Sigma-Aldrich), bovine serum albumin (A0281,  $\geq 99\%$ , Sigma-Aldrich), deoxyribonucleic acids (RPcartridge-gold purity, Eurogentec, Seraing, Belgium) and an artificial foldamer NO<sub>2</sub>-QX<sub>7</sub>-O-X<sub>7</sub>Q-NO<sub>2</sub> (provided by I. Huc, University of Bordeaux).<sup>20</sup> Tunemix was used as received. The foldamer was sprayed from 95/5 (v:v) CHCl<sub>3</sub>/MeOH. Native proteins were prepared in aqueous ammonium acetate (50-200 mM), and 0.5% sulfolane was added to ubiquitin to promote the

formation of 7+ ions. The polythymine mixture was sprayed from RNase free water. The higher-order nucleic acid structures (duplexes and quadruplexes) were sprayed from aqueous ammonium acetate (100 or 150 mM). The present article focuses on the data treatment, and the samples were selected mainly to illustrate different behaviors in terms of ion mobility peak width. More detail on sample preparation can be found in the supporting information.

## THEORY

Combining the mobility equation (1) with the Mason-Schamp Equation (2) leads to equation (3), relating the time spent in the drift tube ( $t_D$ ) to the collision cross section ( $CCS$ ):

$$K_0 = K \frac{p}{p_0} \frac{T_0}{T} = \frac{v_D}{E} \frac{p}{p_0} \frac{T_0}{T} = \frac{L}{t_D E} \frac{p}{p_0} \frac{T_0}{T} \quad (1)$$

$$CCS = \frac{3}{16} \sqrt{\frac{2\pi}{\mu k_B T}} \frac{ze}{N_0} \frac{1}{K_0} \quad (2)$$

$$CCS = \frac{t_D}{L} \frac{3}{16} \sqrt{\frac{2\pi}{\mu k_B T}} \frac{zeE}{N_0} \frac{p_0}{p} \frac{T}{T_0} \quad (3)$$

$K$  is the ion mobility and  $K_0$  is the reduced ion mobility,  $p$  is the pressure,  $p_0$  is the standard pressure (760 Torr),  $T$  is the temperature,  $T_0$  is the standard temperature (273.15 K),  $N_0$  is the buffer gas number density at  $T_0$  and  $p_0$ ,  $L$  is the drift tube length,  $t_D$  is the drift time in the tube,  $E$  is the electric field across the drift tube,  $\mu$  is the reduced mass of the ion-gas interactions ( $\mu = m_{\text{gas}} \cdot m_{\text{ion}} / (m_{\text{gas}} + m_{\text{ion}})$ ),  $k_B$  is the Boltzmann constant,  $z$  is the absolute value of the ion nominal charge, and  $e$  is the charge of the electron.

**$t_0$  is not constant if collisions occur after the mobility region.** Equation (3) or analogues are commonly seen in ion mobility articles presenting collision cross section distributions.<sup>2,21</sup> If  $t_D$  is measured directly (for example using a grid at the end of the drift tube), Eq. (3) can be used. In commercial instruments, however, the measured ion transit time is the time difference between a pulsing gate in front of the tube and a detector further away from the rear of the tube. The arrival time  $t_A$  is the sum of the time spent in the drift cell ( $t_D$ ) and the time spent outside the drift cell ( $t_0$ ):

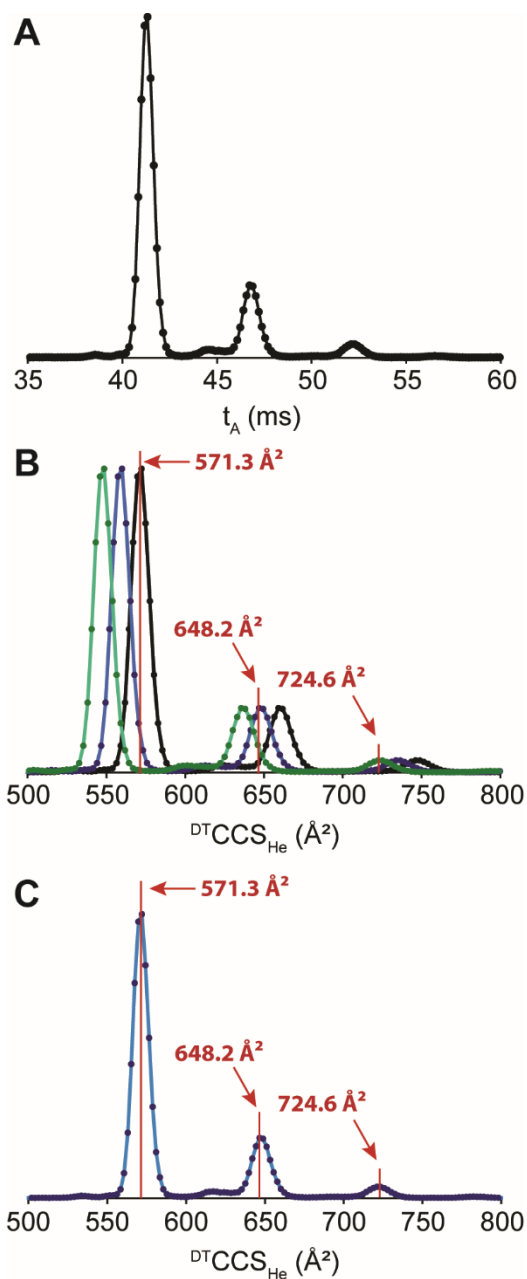
$$t_A = t_D + t_0 \quad (4)$$

A straightforward way to relate the experimental ATD to a CCS distribution is thus to combine Equations (3) and (4):

$$CCS = \frac{(t_A - t_0)}{L} \frac{3}{16} \sqrt{\frac{2\pi}{\mu k_B T} \frac{zeE}{N_0} \frac{p_0}{p} \frac{T}{T_0}} \quad (5)$$

Using Equation (5) to change variables from  $t_A$  to  $CCS$  implicitly assumes that  $t_0$  is independent of the  $CCS$ . This is true only for instruments that would transfer ions immediately to high vacuum, with effectively no further collisions prior to mass analysis. However it is not true for our instrument, which has a rear funnel beyond the drift zone where  $\Delta V$  is varied, and collision cell between the IMS and the mass analyzer. Collisions occur outside the drift tube as well, and the time spent outside the drift tube thus also depends on a collision cross section. The problem with assuming that  $t_0$  is constant is illustrated in Figure 1. Figure 1A shows the arrival time distribution of an organic foldamer molecule, containing three peaks. The collision cross section values obtained by the step-field method on each of the three peaks are 571.3, 648.2 and 724.6  $\text{\AA}^2$ . The  $t_0$  values are 5.85, 6.62 and 7.32 ms, respectively. There is no single “correct”  $t_0$  value which would be valid for all three conformational ensembles. So, using Eq. (5) leads to different  $CCS$  distributions depending on which peak was used to determine  $t_0$  (Fig. 1B). Each reconstruction is incorrectly representing the entire collision cross section distribution.

The difference is visually significant, so Equation (5) is not satisfactory for rendering  $CCS$  distributions containing multiple peaks. A safer option is to present the original arrival time distributions, annotate the  $CCS$  of the center of the peaks on the figure, and tabulate  $CCS$  values separately.<sup>22,23</sup>



**Figure 1.** **A)** Arrival time distribution of an organic foldamer ( $z = +2$ ) measured at  $\Delta V = 390.5$  V. **B)** Collision cross section distributions reconstructed assuming that  $t_0$  is constant: in black, using the  $t_0$  of the left peak, in blue, of the central peak and in green of the right peak. The red annotations represent the  $CCS$  values obtained by the step-field method. **C)** Reconstruction obtained by applying Equation (7). The correction factor  $a$  was determined from the peak on the left ( $571.3 \text{ \AA}^2$ ), yet all peaks are positioned correctly.



***How to better estimate the contribution of  $t_0$  over the entire ATD?*** To reconstruct *CCS* distributions, one must account for the way  $t_0$  depends on the *CCS*. A convenient way is to empirically correlate arrival times ( $t_A$ ) and *CCS* values for a set of ions that represent those of interest. This can be done at a single field, and must be re-done for each instrumental setting that would change either  $t_D$  or  $t_0$ . Traveling wave IMS calibrations follow that procedure, by correlating arrival times ( $t_A$ ) with <sup>DT</sup>*CCS* values of calibrant ions in a given gas.<sup>24</sup>

For drift tubes, combining Eq. (3) and (4) with the expression of the electric field  $E$  as a function of the voltage difference between the entrance and the exit of the IMS cell ( $E = \Delta V/L$ ) gives:

$$t_A = t_0 + \left( CCS \cdot \frac{\sqrt{\mu}}{z} \right) \frac{16L^2}{3e} \sqrt{\frac{k_B T}{2\pi}} \frac{NT_0 p}{T p_0} \times \left( \frac{1}{\Delta V} \right) \quad (6)$$

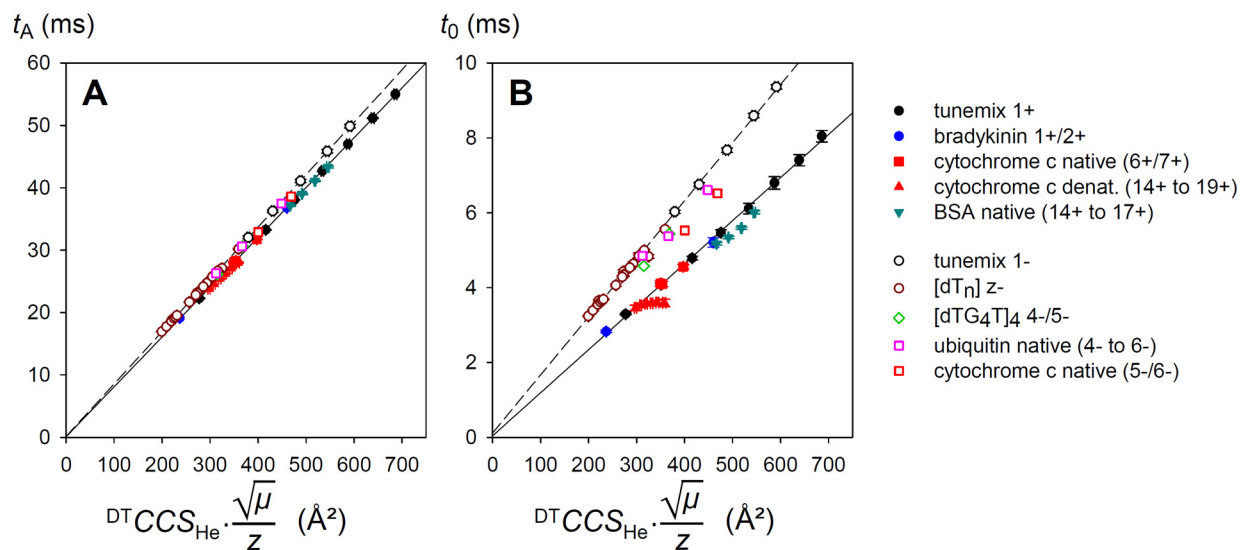
One can either use known <sup>DT</sup>*CCS* values of calibrants, or determine *CCS* values from step-field experiments. We use the latter approach. The step-field experiment consists in measuring the arrival time of the center of the ATD peak ( $t_A$ ) as a function of  $(1/\Delta V)$ , and fit the data by a linear equation to obtain the *CCS* from the slope, according to Equation (6).

In Equation (6),  $(CCS \cdot \sqrt{\mu}/z)$  is related to the system, and all other terms on the right hand side depend on the instrument (length, pressure, temperature and voltage).  $t_0$  could contain an instrument-dependent contribution (a fixed time necessary to travel through various ion optics to the analyzer) and an analyte-dependent contribution, which could itself be related to  $(CCS \cdot \sqrt{\mu}/z)$  if mainly collisions are responsible for slowing down the ions on their way to the analyzer.

At fixed drift voltage  $\Delta V$ , let's examine how  $t_A$  empirically correlates with <sup>DT</sup> $CCS_{He} \cdot \sqrt{\mu}/z$  for a variety of cations and anions, singly or multiply charged (Figure 2A). On first approximation, the correlation is linear and goes through (0,0). But on closer examination, the data segregate according to the analyte. Given that  $t_D$  is strictly proportional to <sup>DT</sup> $CCS_{He} \cdot \sqrt{\mu}/z$ , the segregation can only come from  $t_0$  (Figure 2B).

First, cations and anions follow a different trendline. Because our ultimate goal is to study fragile nucleic acid non-covalent complexes, we had lowered the collision hexapole entrance and delta voltages in the negative mode, to reduce collisional activation in the collision cell (the full list of instrumental parameters is provided in the Supporting Information). For this reason, our anions take longer to reach the mass analyzer. When using the same voltage absolute values in positive

ion mode, ion transmission was lost for large proteins. However, tunemix ions could be analyzed in both modes with the same absolute voltages, and the trends superimpose (see Supporting Information Figure S1).



**Figure 2.** **A)** Arrival time ( $t_A$ ) measured at  $\Delta V = 390.5$  V as a function of  $CCS \cdot \sqrt{\mu}/z$  determined from step-field experiments in helium drift tube, for various cations (filled symbols) and anions (empty symbols). See legend for the classification by compound class. The linear regressions were made using tunemix and bradykinin in positive mode, and tunemix and polythymine oligodeoxynucleotides in negative mode. The system-dependent deviation from the linear correlation comes from  $t_0$ , as expanded in panel **B)**. Error bars are standard deviations from three independent measurements. Note that ions were accelerated differently after the tube in positive and negative mode (IM Hex Entrance = 42 V vs. -32 V; IM Hex Delta = +8V vs. -3V, respectively).

More intriguing, even within a polarity, different molecular classes still segregate. The most striking example is that of denatured cytochrome *c* ions, which have lower  $t_0$  values than the trend line given by tunemix, bradykinin, and native cytochrome *c*. This is rationalized as follows. In our instrument  $t_0$  depends partly on collisions with nitrogen from the collision cell (the residence time in the collision cell is  $\sim 1$  ms,<sup>25</sup> and the rest of the time outside the drift tube is

mostly spent in the rear funnel), whereas  $t_D$  depends exclusively on collisions with helium in the drift tube. The correlation between nitrogen and helium reciprocal mobilities is not linear, and depends on the ion shape.<sup>10</sup> In particular, the correlation is very different for denatured proteins and for globular ions.<sup>26</sup> To verify this interpretation, we recorded positive ion mode mobility data with helium both in the drift tube and in the collision cell, and found that in those conditions, denatured cytochrome *c* data points were no longer outliers (supporting information Figure S2). Finally, in our experimental conditions, the CCS-independent part of  $t_0$  is negligible. Similar results were obtained by the manufacturer.<sup>27</sup>

In summary, a simple linear scaling using Equation (7) is a good approximation for changing variables between  $t_A$  and CCS for a given system:

$$CCS = a \cdot \frac{z}{\sqrt{\mu}} \times t_A \quad (7)$$

In the Agilent 6560 IMS-Q-TOF, changing variables using Equation (7) renders more realistic peak positions than if assuming constant  $t_0$  (see Figure 1C): all peak positions are rendered correctly. Again, if different gases are used in the drift tube and in the collision cell, the exact value of  $a$  will depend on the molecular nature of the ions, and not only on instrumental parameters. To determine  $a$ , knowing the CCS of the center of only one peak is sufficient. For unknowns, one will use the step-field method to determine that CCS. One can also use the CCS of a reference compound to lock the CCS scale. This is analogous to using a lock mass for internal mass calibration. Finally, note that Eq. (7) is valid only if  $z$  is the same for all species underlying the ATD. For example, its application is incorrect to deal with oligomers of different charges overlapping at a given  $m/z$ .

### ***How to account for diffusion and extract the intrinsic width of the CCS distributions?***

Although the previous reconstruction properly renders the peak positions, it does not properly render the width of the CCSDs. Part of the ATD peak width is due to diffusion, and part is due to the conformational ensemble having a distribution of CCS values prior to IMS. We want to extract the latter, which is an intrinsic property of the conformational ensemble contributing to a given peak.

Each peak in the experimental ATD is fitted by a Gaussian curve. We thus here define a conformational ensemble as a collection of analyte conformations that have a Gaussian distribution of collision cross sections. Here  $FWHM_{\text{conf}}$  will denote the contribution of the conformational ensemble to the experimental peak width in the time domain. At each voltage, we obtain the peak center  $t_A$  and the experimental full width at half maximum of the arrival time distribution ( $FWHM_{\text{exp}}$ ). Let's exploit the latter.

If all ions in the ensemble have the same CCS value, the  $FWHM$  is due mainly to diffusion:

$$FWHM_{\text{diff}} = 4t_D \sqrt{\ln(2)} \sqrt{\frac{k_B T}{ze \Delta V}} \quad (8)$$

Diffusion occurs both inside the drift tube (contributing to the width of  $t_D$ ) and outside the drift tube (contributing to the width of  $t_0$ ). Both contribute to the width of the  $t_A$  distribution (i.e., of the ATD). Each can be computed if voltages are known. This is easy for the contribution inside the drift tube. The contribution to the peak width due to phenomena happening outside the drift tube ( $FWHM_{t_0}$ ) is more difficult to account for: it includes contributions from the ion injection gate<sup>28,29</sup> (here,  $t_{\text{gate}} = 100 \mu\text{s}$  in all experiments), from diffusion outside the IMS, and possibly from electric fields outside the IMS. Assuming that the experimental ATD is a convolution of several Gaussian contributions,  $FWHM_{\text{exp}}$  is given by:

$$FWHM_{\text{exp}} = \sqrt{FWHM_{\text{conf}}^2 + FWHM_{\text{diff in IMS}}^2 + FWHM_{t_0}^2} \quad (9)$$

The sum of the latter two contributions can be determined by measuring the  $FWHM_{\text{exp}}$  of a monomorphic compound ( $FWHM_{\text{conf}} = 0$ ) having the same charge and drift time as the analyte of interest. This approach can be applied to traveling wave ion mobility as well,<sup>30</sup> the only difference being the necessity to replace Eq. (7) by an equation appropriate to TWIMS to link the arrival time and CCS variables.<sup>24,31</sup>

In drift tubes, one can also use the step-field experiment to obtain the contribution of  $FWHM_{\text{conf}}$ , alleviating for the need for reference compounds. The first step is to subtract the in-IMS contribution (easy to compute from Eq. (8) in DTIMS) from the experimental FWHM, to calculate  $FWHM_{\text{step}}$ , which is thus the convolution of the contributions of the analyte and of peak broadening outside the drift region (Eq. 10):

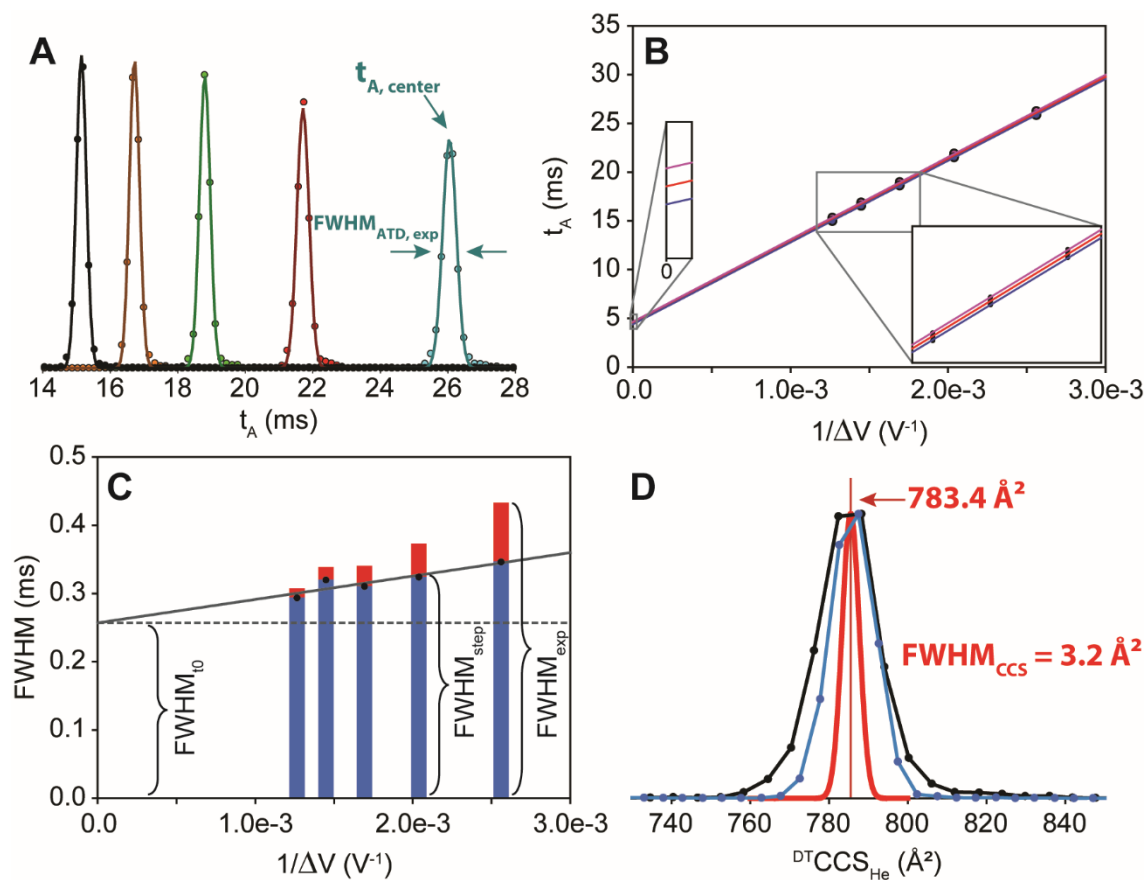
$$FWHM_{step} = \sqrt{FWHM_{exp}^2 - FWHM_{diff\ in\ IMS}^2} = \sqrt{FWHM_{conf}^2 + FWHM_{t_0}^2} \quad (10)$$

The contribution outside the IMS is by definition independent of the drift tube voltage  $\Delta V$ . Thus, any change in  $FWHM_{step}$  along the step-field experiment is related to the analyte's conformational heterogeneity.

Reconstructing the CCS distribution means finding not only the CCS of the center of the peak, but also the CCS of the ions that drifted faster (smaller CCS) or slower (larger CCS) than the center of the peak. We will illustrate the method with the arrival time distributions of the G-quadruplex [dTG<sub>4</sub>T]<sub>4</sub><sup>5-</sup> (Figure 3). Figure 3B shows the linear regression as a function of  $1/V$  for the arrival time of the peak center  $t_{A,center}$ , which according to the usual step-field procedure (Eq. (6)) gives  $CCS_{peak\ center}$ . We do the same for ions with arrival times at half-maximum of the peak, using  $(t_{A,center} \pm FWHM_{step}/2)$ . The contributions outside the IMS contribute to the intercept, but not to the slope. Thus the change of slopes of these two new linear regressions is due only to the contribution of the different CCS values across the conformational ensemble. In other words, the two new CCS values obtained from the two new slopes are equal to  $CCS_{peak\ center} \pm FWHM_{CCS}/2$ . We thus obtain  $FWHM_{CCS}$ , the width of the CCS distribution, and can construct a Gaussian CCS distribution that contains only the contributions of the conformational heterogeneity of the ensemble (Figure 3D).

The procedure explained above is easier to understand intuitively. However it is faster, mathematically equivalent, and more accurate (lower error propagation) to fit directly the  $FWHM_{step}$  values as a function of  $1/\Delta V$ . Also, the residuals and the quality of the fit are easier to assess visually. The  $FWHM_{CCS}$  is directly obtained from the slope of  $FWHM_{step}$  as a function of  $1/\Delta V$  (see Fig. 3C), following Eq. (11), itself derived from Eq. (6):

$$FWHM_{step} = FWHM_{t_0} + \left( FWHM_{CCS} \cdot \frac{\sqrt{\mu}}{z} \right) \frac{16L^2}{3e} \sqrt{\frac{k_B T}{2\pi}} \frac{NT_0 p}{Tp_0} \times \left( \frac{1}{\Delta V} \right) \quad (11)$$



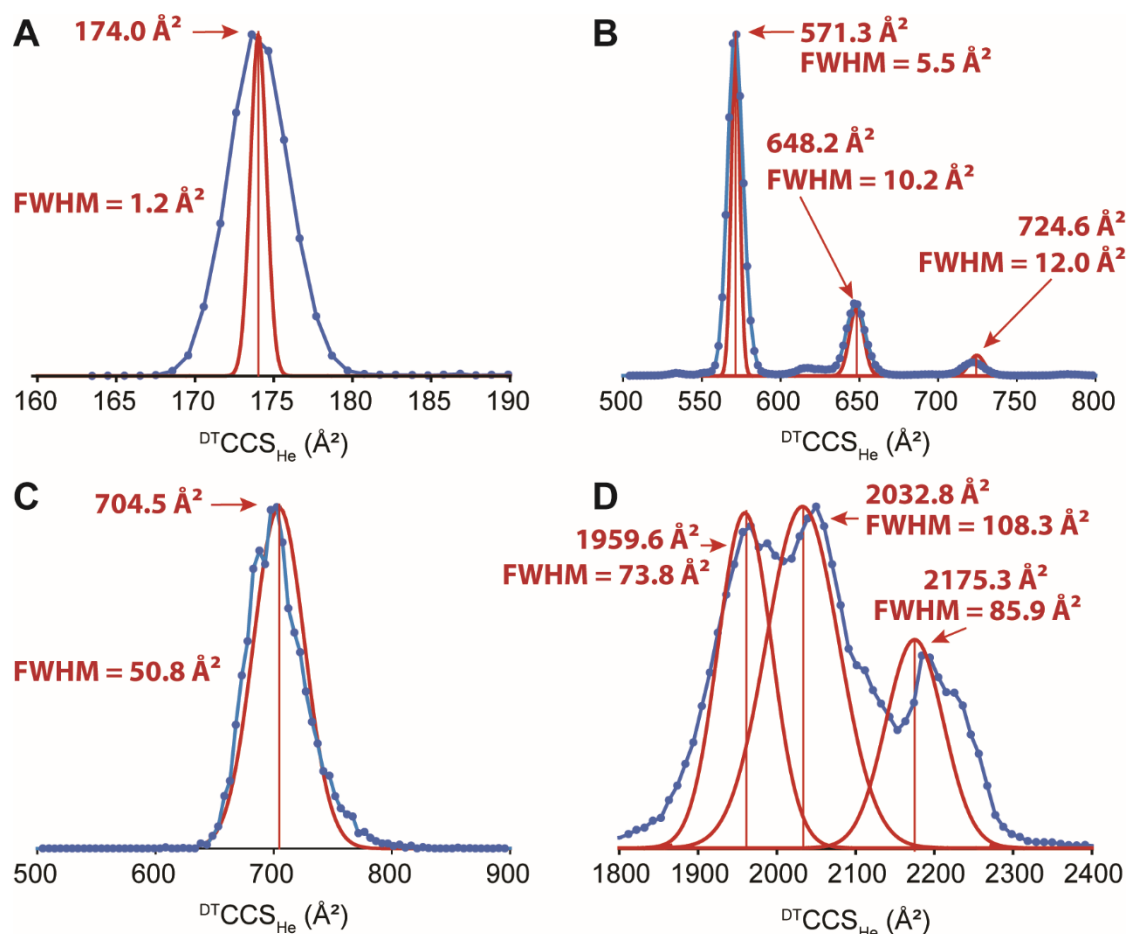
**Figure 3.** **A)** Arrival time distribution for the DNA tetramolecular G-quadruplex  $[dTG_4T]_4^{5-}$  at five different  $\Delta V$  values (390, 490, 590, 690 and 790 V, in cyan, red, green, brown and black, respectively). The Gaussian curves were obtained by fitting the experimental ATDs, and were normalized based on their area. **B)** Arrival time versus  $1/\Delta V$  for the peak center and  $t_{A, \text{center}} \pm \frac{1}{2} FWHM_{\text{step}}$ . Inserts show the fitting of the three series of points and their respective intercept ( $t_0$ ). **C)** How the experimental peak width at half maximum ( $FWHM_{\text{exp}}$ , full bar) and  $FWHM_{\text{step}}$  (blue portion of the bar) vary with  $1/\Delta V$ . The solid line shows the linear fit from which  $FWHM_{\text{CCS}}$  is obtained from the slope, and the  $FWHM_{t_0}$  is obtained from the intercept. **D)** Reconstructed CCS distributions. Black: data points converted using Equation (5) with  $\Delta V = 390.5$  V (reconstruction at all drift voltages, see Supporting Figure S3A). Blue: data points converted using Equation (7) with  $\Delta V = 390.5$  V (reconstruction at all drift voltages, see Supporting Figure S3B). Red: Gaussian curve reconstructed with the “FWHMstep” method.

## RESULTS

Figure 3 illustrates the reconstruction for the DNA tetramolecular G-quadruplex  $[\text{dTG}_4\text{T}]_4^{5-}$ , which has a narrow arrival time distribution (resolving power  $(t_A/\text{FWHM}_{t_A}) \approx 56$  at 390.5 V). When reconstructed with Equation (7), the apparent  $\text{CCS}/\text{FWHM}_{\text{CCS}}$  is naturally identical. Nevertheless, it is still unduly broad. Figure 3C shows how  $\text{FWHM}_{\text{exp}}$  and  $\text{FWHM}_{\text{step}}$  change with  $1/\Delta V$  for  $[\text{dTG}_4\text{T}]_4^{5-}$ . We can see that the diffusion inside IMS contributed significantly to the peak width in the time domain (red bar). Besides, peak broadening outside the IMS is not negligible either ( $\text{FWHM}_{t_0} = 257 \pm 17 \mu\text{s}$ ). When the inherent width of the CCS distribution of the conformational ensemble is obtained from the  $\text{FWHM}_{\text{step}}$  method, the relative  $\text{FWHM}_{\text{CCS}}$  is 0.7% of the center CCS (the equivalent “resolution” would be over 150).

The narrow distribution is attributed to the rigidity of the parallel G-quadruplex stem.<sup>13</sup> Other examples of narrow CCS distributions are shown in Figures 4A (tunemix,  $m = 922$  and  $z = 1$ ) and 4B (the foldamer of Figure 1). The blue distributions are reconstructed using Eq. (7), with the proportionality factor  $a$  determined from the center of the most intense peak, and the red distributions are Gaussian curves reconstructed with the “ $\text{FWHM}_{\text{step}}$ ” method. The linear regressions on the peak widths are shown in Supporting Information Figure S4. The relative peak widths at half maximum ( $\text{CCS}/\text{FWHM}_{\text{CCS}}$ ) are 0.7 % for the tunemix ion 922, and 1.0 % for the foldamer main peak.

Fig. 4C illustrates the processing of a broad ATD, recorded with the DNA duplex  $[\text{dCGCGGGCCCGCG}]_2^{5-}$ . Duplexes are less rigid than parallel G-quadruplexes such as  $[\text{dTG}_4\text{T}]_4^{5-}$ . They collapse into a variety of structures by forming extra phosphate-phosphate hydrogen bonds upon desolvation and declustering, and the various collapsed structures do not interconvert after desolvation.<sup>32</sup> The CCS distribution reconstructed with the “ $\text{FWHM}_{\text{step}}$ ” method is as broad as the one reconstructed using Eq. (7). Its relative width at half-maximum is 7.2 %. One may argue that the absolute contribution of the diffusion differs on both ends of the peaks when the CCS values differ so much. However, the relative contribution of the diffusion to the peak width is very small (see supporting Figure S4), thus this is a very minor source of error.



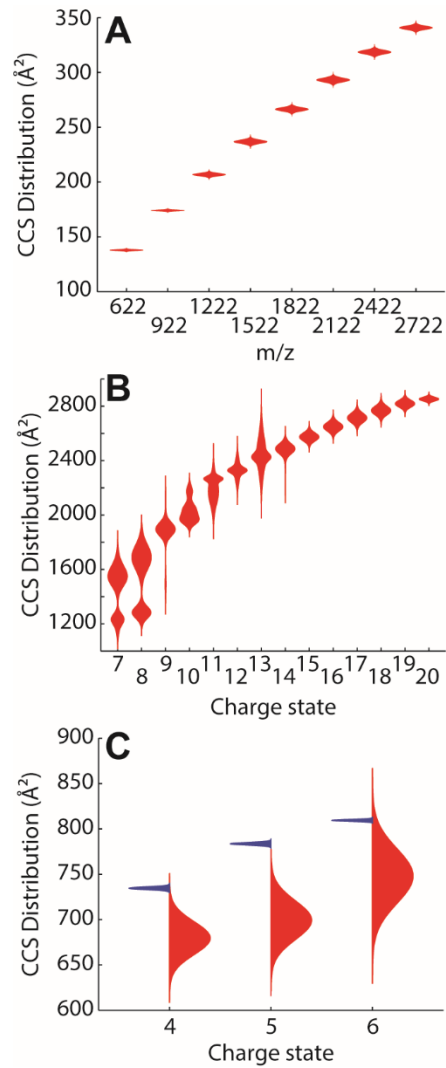
**Figure 4.** Examples of reconstructed CCS distributions. Blue: data points converted using Equation (7). Red: Gaussian curves reconstructed with the “FWHMstep” method. **A)** Tunemix ion at  $m = 922$  and  $z = 1$ . **B)** Foldamer  $[\text{NO}_2\text{-QX}_7\text{-O-X}_7\text{Q-NO}_2]^{2+}$ . **C)** DNA duplex  $[\text{dCGCGGGCCCGCG}]_2^{5-}$ . **D)** denatured  $[\text{cytochrome } c]^{10+}$ .

Finally, Fig. 4D illustrates the output of the “FWHMstep” method on a more complex arrival time distribution ( $[\text{cytochrome } c]^{10+}$  produced from denaturing solution conditions). The experimental arrival time distribution at our lowest drift voltage ( $\Delta V = 390.5$  V) shows four peaks, the one at the longest arrival time being a shoulder of the penultimate peak. However, the two slowest peaks are not resolved at every drift voltage, so we could not apply the FWHMstep method to all four peaks, and thus fitted the last two peaks by a single Gaussian function. This illustrates one limitation of our method: the same peaks must be resolved at all drift voltages of the step-field experiment. The three reconstructed Gaussian curves have different widths, and



this provides information on the polydispersity of each resolvable ensemble. We also see that the positions of the peaks obtained by applying Eq. (7) on the first peak (blue curve) are not the same as when using the step-field method. As we saw in Figure 2, different protein conformations have different  $t_0$  dependences on the helium CCS. The step-field reconstructions (in red) are the correct peak positions.

The reconstructed CCS distributions can be used to build violin plots (Figure 5), which are useful to visually compare peak centers and peak widths simultaneously. The violin plots show a distribution along the vertical axis, for different datasets on the horizontal axis. When only one series is shown, the distribution is presented with its mirror image to create the violin. For example, the evolution of CCS as a function of size is often used to determine if the structure grows following an isotropic or linear law, hinting at spherical or rod-like structures, respectively.<sup>33</sup> Traditional plots show points corresponding to the CCS of the peak center. Figure 5A illustrates this for tunemix ions. The peak width is useful to determine the goodness of a fit, and whether a data point is an outlier or not. Another frequent data visualization is the CCS as a function of the charge state. Such graphs usually display dots.<sup>2</sup> Figure 5B shows the violin plot of denatured cytochrome *c* (as the sum of the reconstructed Gaussian curves), which adds visual information on the relative abundances of each ensemble, and on their intrinsic polymorphism. Finally, asymmetric violin plots (one distribution towards the left, one towards the right) facilitate comparisons between two systems or conditions. Figure 5C shows how two DNA structures with the same size (24 nucleobases) compare. The violin plot highlights simultaneously the difference in CCS values, the charge dependence, and the width of the distribution.



**Figure 5.** Violin plots of CCS distributions reconstructed with the FWHMstep method. The y-axis is the CCS distribution of the ion labeled on the x-axis. For single series (A and B), the mirrored CCS distribution is represented. To visually compare two series (C), different distributions are plotted on the left and on the right. **A)** size dependence of CCS for tunemix in positive mode. **B)** Charge state dependence for denatured cytochrome *c* in positive mode. **C)** Comparison between two DNA systems containing 24 nucleobases: (red) the duplex [dCGCGGGCCCGCG]<sub>2</sub> and (blue) the G-quadruplex [dTG<sub>4</sub>T]<sub>4</sub>, for several charge states in negative mode.

## DISCUSSION

A significantly large  $FWHM_{CCS}$  means that the ion population under a given peak adopts a variety of gas-phase conformations, presenting a distribution of CCS values which is also Gaussian. A Gaussian distribution is the most probable distribution that can be assumed in the absence of resolvable peak shoulders in the ATD. If a shoulder is resolved by a peak fitting software at all fields of the step-field experiment, then it is treated as a separate ensemble. The necessity to obtain quality fits of the ATD can limit the applicability of our method to complex distributions. Peak shoulders are not necessarily resolvable at all fields. For complex ATDs, using Eq. (7) remains the back-up plan, despite not taking some sources of broadening into account. Complex ATD's are usually much broader than those of monomeric analytes would be, and we showed that in such cases (e.g., the duplex) the magnitude of the correction to the peak width offered by the  $FWHM_{step}$  method is small. Remember also that when different gases are used inside and outside the drift tube, the peak positions may not be rendered correctly. Another case of complex ATD consists of species with different charges  $z$  (for example,  $m/z$ -overlapped oligomers). Here, a different Gaussian CCSD must be reconstructed for each peak, using the appropriate  $z$  value. This is however difficult if the peaks overlap.

We also note that the Gaussian distribution of CCS values is valid only for systems that do not interconvert in the drift tube. If interconversion occurs on the time scale of the mobility separation, kinetic modeling is required to account for the shape of the arrival time distribution.<sup>34-36</sup> When inter-conversion is at least 10 times faster than the time scale of mobility separation, the entire ion population drifts with the same average time, and the distribution is as narrow as a diffusion-limited ATD would be.<sup>36</sup> Thus, a significantly broad peak indicates an intrinsic distribution of CCS values prior to entrance in the drift cell, and that these conformations do not interconvert at the temperature and time scale at which ion mobility separation is conducted.

Our procedure doesn't take into account the contribution of space charge. We initially thought that space charge was the reason why we don't find CCS distributions closer to dirac functions, even for the tunemix ions that show very narrow ATDs in super-resolution SLIM (structures for lossless ion manipulation) devices.<sup>37</sup> Our drift tube instrument being placed right after the source, we don't mass-select before IMS, thus broadening due to space charge could also occur

in the ion accumulation region. However, control experiments at lower concentrations and ion intensities led to the same peak widths. The narrowest peaks we obtain have a  $FWHM_{CCS} \leq 1\%$  of the peak center. This is narrower than with the highest resolving power obtainable in helium ( $\sim 56$  for a 5- charge state). We however have not uncovered yet why this error is always positive (not centered around zero-width), i.e. why for the presumably monomorphic tunemix ions the slopes  $\partial FWHM_{step}/\partial(1/\Delta V)$  are systematically positive and not equal to zero. Further investigation of the field effect on the peak width would be needed. In Figure S4, the standard error on the slope is rather constant throughout the dataset (15-30 ms/V<sup>-1</sup>), which translates into an error on the  $FWHM_{CCS}$  of 1.5 to 3 Å<sup>2</sup>. In relative terms, the error on the reconstructed width is substantial for narrow peaks only.

Although our procedure is not quantitatively perfect, it already better reflects the width of the CCS distribution, compared to Eq. (5) or (7). With these two equations, the reconstructed width depends on the resolving power and thus on the drift voltage (Supporting Information Figure S3). Also, in contrast with the procedure recently described by Kune et al.,<sup>30</sup> we make no assumption with regard to the number of conformations hidden under an ATD which is broader than if just diffusion-limited. Here, we treat each Gaussian ATD peak as an ensemble, each having a distribution of CCS values.

## CONCLUSION

In summary, we propose a new data processing method for step-field drift tube ion mobility experiments, to extract the width of the CCS distribution (CCSD) of each conformational ensemble, in addition to the average CCS values of each ensemble. The new “FWHMstep” method allows one to reconstruct CCSDs more faithfully, to visually compare different systems, different experimental conditions, or to compare experimental and theoretical CCS distributions instead of singular CCS values. For narrow peaks observed at the limits of our commercial drift tube’s resolving power in helium ( $\sim 56$ ), the reconstructed CCSDs had peak widths of less than 1% of the half-maximum. Thus, reconstructed distributions broader than 1% of the peak center indicate that ion populations consist of heterogeneous conformational ensembles, which do not interconvert on the time scale of the mobility separation. We also propose using violin plots of

the reconstructed collision cross section distributions to visualize trends in collision cross sections, both for the peak center and for the peak width.

## **ASSOCIATED CONTENT**

### **Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website:

Instrument tuning parameters, detailed sample preparation, list of  $t_A$ ,  $t_0$  and  $CCS$  values of Figure 2,  $t_0$  and  $CCS$  values in positive and negative mode with other collision cell voltages, and with helium in the collision cell, additional arrival time distributions and  $CCS$  distribution reconstructions, Matlab scripts to create the violin plots. (PDF)

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### **Author Contributions**

AM and VG conceived the research, SL and FR collected data, AM, SL, FR and VG treated the data and interpreted the results. AM and VG wrote the manuscript. All authors have given approval to the final version of the manuscript.

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## REFERENCES

- (1) Wyttenbach, T.; Bowers, M. T. *Mod. Mass Spectrom.* **2003**, *225*, 207-232.
- (2) Bohrer, B. C.; Merenbloom, S. I.; Koeniger, S. L.; Hilderbrand, A. E.; Clemmer, D. E. *Annu. Rev. Anal. Chem.* **2008**, *1*, 293-327.
- (3) Clemmer, D. E.; Jarrold, M. F. *J. Mass Spectrom.* **1997**, *32*, 577-592.
- (4) Groessl, M.; Graf, S.; Knochenmuss, R. *Analyst* **2015**, *140*, 6904-6911.
- (5) Merenbloom, S. I.; Glaskin, R. S.; Henson, Z. B.; Clemmer, D. E. *Anal. Chem.* **2009**, *81*, 1482-1487.
- (6) Shvartsburg, A. A.; Seim, T. A.; Danielson, W. F.; Norheim, R.; Moore, R. J.; Anderson, G. A.; Smith, R. D. *J. Am. Soc. Mass Spectrom.* **2013**, *24*, 109-114.
- (7) Deng, L.; Webb, I. K.; Garimella, S. V. B.; Hamid, A. M.; Zheng, X.; Norheim, R. V.; Prost, S. A.; Anderson, G. A.; Sandoval, J. A.; Baker, E. S.; Ibrahim, Y. M.; Smith, R. D. *Anal. Chem.* **2017**, *89*, 4628-4634.
- (8) Chouinard, C. D.; Wei, M. S.; Beekman, C. R.; Kemperman, R. H.; Yost, R. A. *Clin. Chem.* **2016**, *62*, 124-133.
- (9) Zheng, X.; Wojcik, R.; Zhang, X.; Ibrahim, Y. M.; Burnum-Johnson, K. E.; Orton, D. J.; Monroe, M. E.; Moore, R. J.; Smith, R. D.; Baker, E. S. *Annu. Rev. Anal. Chem.* **2017**, *10*, 71-92.
- (10) Bush, M. F.; Hall, Z.; Giles, K.; Hoyes, J.; Robinson, C. V.; Ruotolo, B. T. *Anal. Chem.* **2010**, *82*, 9557-9565.
- (11) Jurneczko, E.; Barran, P. E. *Analyst* **2011**, *136*, 20-28.
- (12) Marklund, E. G.; Degiacomi, M. T.; Robinson, C. V.; Baldwin, A. J.; Benesch, J. L. *Structure* **2015**, *23*, 791-799.
- (13) D'Atri, V.; Porrini, M.; Rosu, F.; Gabelica, V. *J. Mass. Spectrom.* **2015**, *50*, 711-726.
- (14) May, J. C.; Morris, C. B.; McLean, J. A. *Anal. Chem.* **2017**, *89*, 1032-1044.

- (15) Utrecht, C.; Rose, R. J.; van, D. E.; Lorenzen, K.; Heck, A. J. *Chem. Soc. Rev.* **2010**, *39*, 1633-1655.
- (16) Koeniger, S. L.; Merenbloom, S. I.; Clemmer, D. E. *J. Phys. Chem. B* **2006**, *110*, 7017-7021.
- (17) Allen, S. J.; Eaton, R. M.; Bush, M. F. *Anal. Chem.* **2017**, *89*, 7527-7534.
- (18) Maurer, M. M.; Donohoe, G. C.; Valentine, S. J. *Analyst* **2015**, *140*, 6782-6798.
- (19) Laszlo, K. J.; Munger, E. B.; Bush, M. F. *J. Phys. Chem. B* **2017**, *121*, 2759-2766.
- (20) Li, X.; Qi, T.; Srinivas, K.; Massip, S.; Maurizot, V.; Huc, I. *Org. Lett.* **2016**, *18*, 1044-1047.
- (21) Shi, L.; Holliday, A. E.; Shi, H.; Zhu, F.; Ewing, M. A.; Russell, D. H.; Clemmer, D. E. *J. Am. Chem. Soc.* **2014**, *136*, 12702-12711.
- (22) Ujma, J.; De Cecco, M.; Chepelin, O.; Levene, H.; Moffat, C.; Pike, S. J.; Lusby, P. J.; Barran, P. E. *Chem. Commun.* **2012**, *48*, 4423-4425.
- (23) Bleiholder, C.; Do, T. D.; Wu, C.; Economou, N. J.; Bernstein, S. S.; Buratto, S. K.; Shea, J. E.; Bowers, M. T. *J. Am. Chem. Soc.* **2013**, *135*, 16926-16937.
- (24) Ruotolo, B. T.; Benesch, J. L.; Sandercock, A. M.; Hyung, S. J.; Robinson, C. V. *Nat. Protoc.* **2008**, *3*, 1139-1152.
- (25) Ibrahim, Y. M.; Baker, E. S.; Danielson, W. F., 3rd; Norheim, R. V.; Prior, D. C.; Anderson, G. A.; Belov, M. E.; Smith, R. D. *Int. J. Mass Spectrom.* **2015**, *377*, 655-662.
- (26) Bleiholder, C.; Johnson, N. R.; Contreras, S.; Wyttenbach, T.; Bowers, M. T. *Anal. Chem.* **2015**, *87*, 7196-7203.
- (27) Stow, S. M.; Causon, T. J.; Zheng, X.; Kurulugama, R. T.; Mairinger, T.; May, J. C.; Rennie, E. E.; Baker, E. S.; Smith, R. D.; McLean, J. A.; Hann, S.; Fjeldsted, J. C. *Anal. Chem.* **2017**, *89*, 9048-9055.
- (28) Siems, W. F.; Wu, C.; Tarver, E. E.; Hill, H. H.; Larsen, P. R.; McMinn, D. G. *Anal. Chem.* **1994**, *66*, 4195-4201.

- (29) Wu, C.; Siems, W. F.; Asbury, G. R.; Hill, H. H. *Anal. Chem.* **1998**, *70*, 4929-4938.
- (30) Kune, C.; Far, J.; De Pauw, E. *Anal. Chem.* **2016**, *88*, 11639-11646.
- (31) Shvartsburg, A. A.; Smith, R. D. *Anal. Chem.* **2008**, *80*, 9689-9699.
- (32) Porrini, M.; Rosu, F.; Rabin, C.; Darre, L.; Gomez, H.; Orozco, M.; Gabelica, V. *ACS Cent. Sci.* **2017**, *3*, 454-461.
- (33) Bleiholder, C.; Dupuis, N. F.; Wyttenbach, T.; Bowers, M. T. *Nature Chem.* **2011**, *3*, 172-177.
- (34) Kinnear, B. S.; Hartings, M. R.; Jarrold, M. F. *J. Am. Chem. Soc.* **2001**, *123*, 5660-5667.
- (35) Kohtani, M.; Jones, T. C.; Sudha, R.; Jarrold, M. F. *J. Am. Chem. Soc.* **2006**, *128*, 7193-7197.
- (36) Poyer, S.; Comby-Zerbino, C.; Choi, C. M.; MacAleese, L.; Deo, C.; Bogliotti, N.; Xie, J.; Salpin, J. Y.; Dugourd, P.; Chirof, F. *Anal. Chem.* **2017**, *89*, 4230-4237.
- (37) Deng, L.; Ibrahim, Y. M.; Hamid, A. M.; Garimella, S. V.; Webb, I. K.; Zheng, X.; Prost, S. A.; Sandoval, J. A.; Norheim, R. V.; Anderson, G. A.; Tolmachev, A. V.; Baker, E. S.; Smith, R. D. *Anal. Chem.* **2016**, *88*, 8957-8964.