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Effective temperature of ions in traveling wave ion mobility spectrometry

Denis Morsa, Valérie Gabelica*, Edwin De Pauw

Physical Chemistry and Mass Spectrometry Laboratory, Department of Chemistry, University of Liège, Institut de Chimie Bat. B6c, B-4000 Liège, Belgium.

AUTHOR EMAIL ADDRESS: v.gabelica@ulg.ac.be

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ABSTRACT. Traveling wave ion mobility spectrometers (TW IMS) operate at significantly higher fields than drift tube ion mobility spectrometers. Here we measured the fragmentation of the fragile $p$-methoxybenzylpyridinium ion inside the TW ion mobility cell of the first-generation SYNAPT HDMS spectrometer. The ion’s vibrational internal energy was quantified by a vibrational effective temperature $T_{\text{eff,vib}}$, which is the mean temperature of the ions inside the cell that would result in the same fragmentation yield as observed experimentally. Significant fragmentation of the probe ion inside the TW IMS cell was detected, indicating that field heating of the ions takes place in TW IMS. For typical small molecule IMS conditions, $T_{\text{eff,vib}} = 555 \pm 2$ K. The variations of the effective temperature were studied as a function of the IMS parameters, and we found that $T_{\text{eff,vib}}$ decreases when the wave height decreases, when the pressure increases, or when the wave speed increases. The energy transfer efficiency of argon is higher than for He, N$_2$ or CO$_2$. $T_{\text{eff,vib}}$ being directly related to the ion speed inside the TW IMS, our results also provide new insight on the ion movement in TW IMS. We also discuss the influence of field heating of ions for calibration and structural studies in TW IMS.
INTRODUCTION

Ion mobility spectrometry (IMS) allows the separation of ions based on their three-dimensional structure. Whether used as a standalone separation method,\textsuperscript{1-3} or hyphenated to mass spectrometry, IMS has found numerous applications encompassing separation and structural studies of small molecules, polymers, proteins and nucleic acids, synthetic supramolecular assemblies and biologically relevant complexes.\textsuperscript{4-7}

Ion mobility separation is based on ion transport properties in gases, under the influence of an electric field.\textsuperscript{8,9} Different types of mobility cells exist, differing by the magnitudes of the electric field $E$ and of the gas number density $N$. Drift tube (DT) IMS cells operate at constant field, and at low $E/N$ ratios. The ions move in the field direction at a drift velocity $v$ that depends on their mobility $K$ as stated by Equation (1).

$$ v = KE = K_0 N_0 \cdot \frac{E}{N} $$

where $K_0$ is the ion reduced mobility defined at standard temperature ($T_0 = 273.16$ K) and pressure ($p_0 = 760$ Torr). At low $E/N$ conditions, the ion drift motion does not significantly disturb its thermal motion, and $K_0$ can be expressed quantitatively by the Mason-Schamp equation (Eq. 2),

$$ K_0 = \frac{3}{16} \frac{q}{N} \left( \frac{2\pi \mu k T}{\Omega} \right)^{1/2} \frac{1}{\Omega} $$

Where $q$ is the ion charge, $\mu$ is the reduced mass ($\mu = \frac{mM}{(m+M)}$, where $M$ is the mass of the gas and $m$ is the mass of the ion), $k$ is the Boltzmann constant, $T$ is the gas temperature, and $\Omega$ is the collision cross section.

Commercial IMS-MS instruments are currently based on other IMS methods involving higher fields, namely field-assisted waveform IMS (FAIMS),\textsuperscript{10} or traveling wave IMS (TW IMS).\textsuperscript{11,12} The use of higher $E/N$ ratios raises fundamental questions regarding the ion transport regime. Indeed, when this ratio increases, the ion drift velocity $v$ becomes significantly higher than the ion’s thermal velocity, and thus has to be accounted for in the ion transport equations.\textsuperscript{8} Assuming elastic collisions between the ion
and the gas, the Mason-Schamp equation can still be used provided that the gas temperature $T_{gas}$ is replaced by an effective temperature $T_{eff}$ (Eq. 3) that accounts for the field contribution:

$$T_{eff} = T_{gas} + T_{field} = T_{gas} + \frac{M}{3k}v^2$$

(Eq. 3)

The possibility that $T_{eff}$ might be significantly higher than $T_{gas}$ when $E/N$ increases has important implications. First, the knowledge of its value is crucial for a rigorous interpretation of mobility measurements in terms of ion shapes via the collision cross sections. Intriguingly, recent reports show that TW IMS calibration of $\Omega$ against $t_d$ is not as universal as initially postulated. Bush et al. have shown that calibrating with denatured proteins leads to significant errors in the determination of collision cross sections of native proteins. Other recent reports show that the accuracy of the collision cross section calibration depends on wave height, and even more on the wave velocity. These findings could reflect that ion transport in TW IMS is not exactly the same as in DT IMS, and that high field effects on the ion transport might be involved.

The second implication is that translational energy can be converted into rotational and vibrational internal energy via inelastic collisions. This point has already been raised for TW IMS by Shvartsburg and Smith, and is also crucial for relating ion mobility data to the ion’s shape. Vibrational excitation can indeed cause modifications of ion geometries (and therefore of $\Omega$) because different conformational states of the molecule can become populated. At extreme cases, vibrational excitation could even lead to fragmentation of the ion inside the IMS cell, like in collision-induced dissociation (CID). Therefore, in order to devise guidelines for properly using TW IMS to infer structural information, it is crucial to assess whether field heating causes significant increases of the ion’s internal energies. Here we used the well-established survival yield method to determine the vibrational effective temperature of a small benzylpyridinium ion inside a commercial TW IMS instrument.
METHODS

**Chemicals.** The \( p \)-methoxybenzyl/pyridinium salt was synthesized by condensation of the \( p \)-methoxybenzyl chloride with pyridine. The recrystallized salt was solubilized with acetonitrile/water (50/50) and injected at \( 10^{-6} \) mol/L.

**Instrument.** All measurements were performed on a first-generation Synapt HDMS spectrometer (Waters, Manchester, U.K.) with samples introduced by electrospray. Pressure in the IMS was measured using both a pirani gauge and a gas-independent capacitance gauge (Oerlikon Leybold Vacumm, CERAVAC CTR 90) linked to a vacuum controller (Vacom, MVC-3 UHV Multichannel Vacuum Controller). These were connected to the same inlet. All pressures reported in this work are those measured with the capacitance gauge. Instrument default set up and a table listing the corresponding pressure values between Pirani and capacitance gauge are provided in Supporting Information S1 and S2, respectively.

**Calculation of the ion vibrational effective temperature.** We studied the fragmentation reaction of \( p \)-methoxybenzylpyridinium ions \((m/z = 200)\) which leads to the formation of \( p \)-methoxybenzyl cations \((m/z = 121)\) and neutral pyridine. All arrival time distributions were analyzed with Peakfit, Systat Software (v. 4.11) and peaks were fitted using an exponentially modified Gaussian (EMG). Peak areas related to the precursor ions \( A_{\text{precursor}} \) and to the fragment ions \( A_{\text{fragment}} \) were quantified and used to calculate fragmentation yields \((FY)\) and survival yields \((SY)\) as given by Equation 4.

\[
FY = \frac{A_{\text{fragment}}}{A_{\text{precursor}} + A_{\text{fragment}}} = 1 - SY
\]  

\( T_{\text{eff,vib}} \) is defined as the mean temperature of the ions that would result in the same survival yield as experimentally observed. Because numerous collisions with the gas are at the basis of IMS, we can safely assume that the ions inside the TW IMS undergo sufficiently frequent low energy collisions so
that the ion population can be characterized by a Boltzmann distribution at a temperature $T_{\text{eff,vib}}$ all along the cell. For each measurement, the temperatures were calculated from the survival yields according to a method previously described in the literature\textsuperscript{22-25} and detailed in Supporting Information S3-S5. All arrival time distributions were recorded at least three times for each set of experimental conditions, and processed independently for extracting $F_Y$, $S_Y$, and the drift time $t_d$. All $T_{\text{eff,vib}}$ data presented here report the mean values and standard deviations of at least three independent measurements and data treatments.

**Estimation of the average ion speed inside the TW IMS.** As will be discussed further below, both $T_{\text{eff}}$ (Eq. 4) and $T_{\text{eff,vib}}$ are related to the ion speed $v$. In TW IMS, the ion speed is different from the experimentally accessible apparent drift velocity $\overline{v_{d(TW)}}$, which is equal to the tube length ($L = 185$ mm) divided by the corrected drift time $\tau$ (see Supporting Information S4). The mathematical description of traveling wave ion guides given by Shvartsburg and Smith\textsuperscript{20} allows to relate the average ion speed $v$ to the apparent drift velocity $\overline{v_{d(TW)}}$ and the wave speed $s$. For the Synapt HDMS, $v$ was estimated by using Equation 5 (see derivation in Supporting Information S6).

$$v^2 = \frac{2 \overline{v_{d(TW)}} s^2}{s + \overline{v_{d(TW)}}}$$ (5)
RESULTS AND DISCUSSION

**Fragmentation before, after, and inside the TW IMS**

Product ion spectra of the para-methoxybenzylpyridinium ion \( (M^+, m/z = 200) \) recorded in the ion mobility mode revealed significant fragmentation into para-methoxybenzyl \( (F^+, m/z = 121) \), even when recorded at the lowest possible trap and transfer collision energies. A typical product ion spectrum is shown in Figure 1(a). The fragments \( F^+ \) detected are due to \( M^+ \) fragmentation after the quadrupole and before the TOF pusher. This can be in the trap T-wave, in front of the IMS cell, inside the IMS cell, and/or in the transfer T-wave. As the goal is to characterize ion heating inside the TW IMS cell, we will use the arrival time distribution of the fragments to extract the contribution due to fragments formed inside the TW IMS.

The arrival time of an ion is equal to the time elapsed between its entrance in the TW IMS cell and the moment it reaches the TOF pusher. The ATD of fragments formed in the source and selected in the quadrupole is shown in Figure 1(b). This ATD is therefore representative of fragments formed before the TW IMS. Figure 1(c) shows the ATD of precursor ions selected in the quadrupole and detected as precursors. The two ATDs of Figures 1(b) and 1(c) give the key to interpret the ATD of the fragments formed after selection of the precursor \( M^+ \) in the quadrupole. Fragments formed before the TW IMS have the same ATD as selected fragments in Figure 1(b), whereas fragments formed after the TW IMS have the same ATD as selected precursors in Figure 1(c) because they travelled inside the TW IMS as precursor ions.

A typical experimental ATD of fragments formed from the precursors selected in the quadrupole, i.e. the ATD reconstructed for the \( m/z = 121 \), is shown in Figure 1(d). This ATD shows three contributions. The one at earliest arrival times (in green) corresponds to fragments formed before the TW IMS, for example in the trap region. The one at latest arrival times (in blue) corresponds to fragments formed...
after the TW IMS, for example in the transfer region. The third contribution (in red) is constituted by fragments arriving at intermediate times. It corresponds to fragments that spent part of their time in the TW IMS as precursor ions and part of their time as fragment ions, and therefore it corresponds to fragments formed inside the TW IMS.

In order to determine the fragmentation yields in the different regions of the mass spectrometer, the ion counts from the different contributions were integrated. The area of the precursor ion ($A_M$) was integrated from the total ATD at $m/z = 200$. To separate the different contributions in each fragment ion ATD, the curves corresponding to fragments formed purely before IMS and to those formed purely after IMS were fitted to extract the peak shapes of these contributions. Then, the experimental ATD of the fragment formed from the selected precursor ion (Figure 1(d)) was fitted with two peaks having the shapes determined from Figures 1(b) and 1(c). Only the peak height was allowed to vary. These fitted peaks were integrated to give $A(F,\text{before})$ and $A(F,\text{after})$, the peak areas corresponding to the fragment formed before and after the TW IMS cell, respectively. Finally, these two fitted peaks were subtracted from the experimental data to obtain the ion counts corresponding to fragments formed inside the TW IMS (red contribution in Figure 1(d)). Integrating this contribution gave $A(F,\text{inside})$.

Based on these areas, the fragmentation yields ($FY$) of the parent ion before, inside, and after the IMS were calculated using adequate expressions of Equation 4 (see detailed equations in Supporting Information S7). To prove that the peak assignment and the data treatment are fundamentally correct, we determined the three fragmentation yields as a function of the trap collision energy, which should only affect $FY_{\text{before}}$, and as a function of the transfer collision energy, which should only affect $FY_{\text{after}}$. The results in Figures 2(a) and 2(b) show that only the expected fragmentation yields are affected, thereby validating our approach.
Figure 1. Mass spectra and arrival time distributions (ATDs) recorded on the Synapt HDMS with the following settings: bias = 13V, CE1/CE2 = 2V, wave height = 13V, wave speed = 600 m/s, N2 pressure = 0.516 mbar. (a) Product ion spectrum of the para-methoxybenzylpyridinium ion M$^+$ ($m/z = 200$). The para-methoxybenzyl fragment ion F$^+$ ($m/z = 121$) is detected. (b) ATD of F$^+$ when selecting $m/z = 121$ in the quadrupole. (c) ATD of M$^+$ when selecting $m/z = 200$ in the quadrupole. (d) ATD of F$^+$ ($m/z = 121$) when selecting the precursor ion at $m/z = 200$ in the quadrupole. The contribution shown in green is similar to the ATD of F$^+$ in (b), the one shown in blue is similar to the ATD of M$^+$ in (c). The remaining contribution obtained by subtraction is shown in red.
**Figure 2.** Effect of strategic potential differences on the fragmentation occurring before, inside and after the mobility cell. (a) Influence of the trap collision energy (bias = 13V, Transfer CE = 2V, wave height = 13V, wave speed = 600 m/s, N$_2$ pressure = 0.516 mbar. (b) Influence of the transfer collision energy (same conditions as above, Trap CE = 2 V). (c) Influence of the bias voltage. (Trap/Transfer CE = 2 V).
In a similar experiment, we tested the effect of the bias voltage, which is the acceleration voltage between the last lens before the IMS and the IMS itself, to determine whether increasing the bias was causing fragmentation before and/or inside the IMS. The results in Figure 2(c) reveal that the bias voltage has a dramatic influence on the fragmentation contribution before the IMS. The role of the bias voltage is to accelerate the ions from the lower pressure region of the trap to the higher pressure region of the IMS. It is already known that the bias must be kept low to reduce ion fragmentation, but this is detrimental to the ion transmission. Typical bias voltages recommended for intact protein analysis are 20-60 V.\textsuperscript{13} With our fragile probe ion, fragmentation is detected already at a bias voltage as low as 10 V.

Moreover, Figure 2(c) shows that fragmentation occurs mostly before the IMS. This observation is consistent with activation taking place in front of the IMS cell and imparted by collisions with the gas leaking out from the front and end apertures of the mobility cell. At bias voltages between 10 V and 19 V, although the fragmentation before the IMS is abundant, the fragmentation inside the IMS remains low. This suggests that precursor ions are significantly heated just before the IMS cell, and then cooled down once entering the IMS cell. All subsequent measurements aimed at investigating the effect of the TW IMS cell parameters on the effective vibrational temperature of the ions were carried out at a bias voltage of 10 V in order to minimize the contribution to fragmentation due to the bias acceleration.

**On the interpretation of \( T_{\text{eff,vib}} \)**

As a starting point, we determined the ion temperature using a set of IMS parameters typically recommended by the manufacturer (wave height = 9 V, wave speed = 600 m/s, IMS gas: N\textsubscript{2} pirani reading = 0.55 mbar, i.e. \( p = 0.516 \) mbar as measured with the capacitance gauge). Other acceleration voltages were kept low (Bias = 10 V, trap collision energy = 2 V and transfer collision energy = 2 V). These values were selected to ensure the best compromise between transmission and separation of analyzed species. For these conditions, we found \( T_{\text{eff,vib}} = 555 \pm 2 \) K. If we assume a bath gas temperature
$T_{\text{gas}} = 300 \text{ K}$, this corresponds to a field contribution to ion heating $T_{\text{field}} = 255 \pm 2 \text{ K}$, which indicates a significant field heating of ions.

The effective vibrational temperature should be proportional to the relative kinetic energy of collision between the ion and the gas $E_{\text{kin,rel}}$, which is the maximum amount of kinetic energy that can be transformed into ion internal energy per collision.\textsuperscript{19} $E_{\text{kin,rel}}$ is given by Equation 6, where $v$ is the ion velocity in the laboratory frame of reference, and $\mu$ is the reduced mass.

$$T_{\text{eff,vib}} \propto E_{\text{kin,rel}} = \frac{1}{2} \mu \, v^2$$

In the traveling wave ion guide, the ions alternatively move forward and backwards depending on whether they are on the front side or on the rear side of a wave, respectively. The average $T_{\text{eff,vib}}$ therefore depends on the ions’ average speed $v$ (the speed is the absolute value of the velocity), independently of whether the ions move forward or backwards, because the relevant parameter is their relative collision energy compared to the gas, which is considered as static on average. The value of $T_{\text{eff,vib}}$ is thus an indicator of the ions speed $v$ inside the TW cell.
Influence of the wave height and pressure on the ion temperature

In steady state conditions the ion velocity $v$ depends on $E/N$ (Equation 1). Experimental variables affecting the $E/N$ ratio will therefore be discussed jointly. The gas number density $N$ is directly proportional to the gas pressure $p$, while the average electric field intensity $E$ is directly proportional to the wave height. The influence of the gas pressure on the ion effective temperature at constant wave height is illustrated in Figure 3(a). All the other instrumental parameters remaining unchanged, the ion temperature decreases from 706 K to 555 K when the pressure increases from 0.275 mbar to 0.516 mbar. The influence of the wave height on the ion effective temperature, at three different pressures, is illustrated on Figure 3(b). $T_{\text{eff,vib}}$ increases when the wave height increases. The offset between the curves is due to the pressure effect.

As explained in Eq. (6), $T_{\text{eff,vib}}$ should be proportional to the ion speed squared $v^2$, which should be proportional to $E^2/N^2$ (Eq. 1). To test this hypothesis, we estimated $v^2$ as described in Equation (5) and Supporting Information S6. The plot of $T_{\text{eff,vib}}$ as a function of $v^2$ is shown on Figure 3(c). The experimental points reported on Figure 3(b) are now more closely bunched together, although a slight offset persists, which can be due to the approximations behind the estimation of $v$ from the drift times.

Overall, changing either the wave height or the gas pressure has a similar effect on the ion effective temperature. The analyst willing to obtain the lowest effective temperature should therefore tune either the wave height or pressure so as to diminish the ion velocity $v$. At constant wave speed $s$, this means tuning to lower the apparent drift velocity $v_{\text{drift}}(TW)$, and therefore to increase the apparent drift time $t_d$ (see Supporting Information S8).
Figure 3: Influence of pressure, wave height and ion drift speed on $T_{\text{eff,vib}}$. Conditions were set as follows: bias = 10V, trap/transfer CE = 2V, wave speed = 600 m/s, gas = N$_2$. (a) Evolution of $T_{\text{eff,vib}}$ as a function of gas pressure at a wave height = 9 V. (b) Evolution of $T_{\text{eff,vib}}$ as a function of wave height at three different pressures (0.275 mbar, 0.4 mbar and 0.516 mbar). (c) Evolution of $T_{\text{eff,vib}}$ with ion drift speed squared $v^2$, estimated using Equation (5).
Influence of the wave speed on the ion temperature

To investigate the influence of the wave speed $s$ on $T_{\text{eff,vib}}$, the pressure was kept constant and the effect of the wave speed was studied at different wave heights. The results on Figure 4(a) show that $T_{\text{eff,vib}}$ decreases when $s$ increases. This means that the average ion speed $v$ decreases when $s$ increases. When $v$ is estimated from $\overline{v_d(TW)}$ and $s$ using Equation (5), $v$ indeed decreases when $s$ increases. Figure 4(b) shows the correlation between $T_{\text{eff,vib}}$ and $v^2$. Even if the alignment of the data points is not perfect given the experimental error, this figure confirms that $v$ estimated using Eq. (5) is on first approximation a good predictor of the ion effective temperature. Supporting Information S9 shows additional data as a function of the wave height, with wave speeds ranging from 400 to 1500 m/s.

**Figure 4:** (a) Influence of wave speed $s$ on $T_{\text{eff,vib}}$. Conditions were set as follows: bias = 10 V, trap/transfer CE = 2V, N$_2$ pressure = 0.516 mbar. Four series of wave height were studied: 9 V, 11 V, 13 V and 15 V. Wave speed was varied between 400 m/s and 1000 m/s. (b) Correlation between the ion apparent speed squared $v^2$ and $T_{\text{eff,vib}}$. 
The fact that the ion speed $v$ changes with the wave speed $s$ reveals an interesting aspect of the ion movement in TW IMS. In effect, the wave speed $s$ affects neither the electric field $E$ nor the gas number density $N$, so it should not affect the $E/N$ ratio. If the ions could reach the steady state drift velocity $v = KE$ at every time point in the TW IMS, $v$ should actually be independent of $s$. The fact that $v$ decreases when $s$ increases therefore indicates that the ions do not always have time to reach their steady state drift velocity $KE$. In other words, the ions cannot instantaneously accommodate for the changes of electric field intensity and/or direction. Therefore, in TW IMS, instead of Equation (1) one should take into account that:

$$v \leq KE$$

(7)

Higher wave speeds induce a larger inequality, because the ions have less time to keep up with the field changes. This is important for the understanding of the ion movement in TW IMS, but also for extracting ion mobilities $K$ from the apparent drift velocities $\overline{v_d(TW)}$. The equations expressing the $\overline{v_d(TW)}$ as a function of $KE$ for TW IMS$^{20}$ currently assume that $v$ is equal to $KE$ at all times. Our results suggest that this is not the case, and that the deviation increases as the wave speed increases.
**Influence of the gas (He, N₂, Ar, CO₂) on ion heating**

Nitrogen is the most often used gas in TW IMS, because it allows good transmission and separation of most analytes. Helium is however typically used in drift tube IMS for structural analysis by comparison between experimental and theoretical collision cross sections. Finally, argon and carbon dioxide can be used to enhance separation of some analytes.²⁶-²⁹ We investigated the effect of the nature of the gas on the ion field heating in TW IMS.

Figure 5(a-b) shows the evolution of $T_{\text{eff,vib}}$ as a function of the wave height. Not all four gases could be compared at the same pressure and the same wave speed. In conditions adequate for N₂, Ar and CO₂ ($s = 600$ m/s, $p = 0.275$ mbar, Figure 5(a)), the very high mobility of ions in helium induces a coalescence of the ATD peaks that makes the contribution inside the IMS impossible to quantify. Helium was therefore compared to nitrogen at $p = 0.516$ mbar and $s = 1500$ m/s in Figure 5(b). Note also that the bias had to be increased to 24 V in He, compared to 10 V in the other gases. We checked that this did not influence $FY_{\text{inside}}$.

We can see from these figures that, all IMS instrumental parameters (wave height, wave velocity and pressure) being equal, ion heating ranks as follows: He > Ar > N₂ > CO₂. It seems counterintuitive that helium induces more heating than heavier gases, but the nature of the gas greatly affects the ion mobility $K_0$, both because via the reduced mass $\mu$ and the collision cross section $\Omega$ (see Equation 2). When the ion temperatures are compared at a given ion speed squared $v^2$ (Figure 5c), the ranking becomes Ar > CO₂ > N₂ > He.

As explained above, the ion internal temperature is expected to depend on the relative kinetic energy $E_{\text{kin,rel}}$, which is the maximum kinetic energy transferrable into internal energy. From the estimated ion speed $v$ and the reduced mass of the system, the relative kinetic energy per collision can be estimated using Equation (6). The correlation between $T_{\text{eff,vib}}$ and $E_{\text{kin,rel}}$ is shown in Figure 5(d). This figure gathers
all the datapoints collected throughout this study, because we have shown before that the calculation of \( v \) using Equation (5) provides a good estimate that correlates well with \( T_{\text{eff,vib}} \) for a given gas. This graph shows that the effect of the reduced mass on \( E_{\text{kin,rel}} \) suffices to explain the ranking between CO\(_2\), N\(_2\) and He. In contrast, the argon data deviate significantly from those obtained in the three other gases. Collisions with argon at a given \( E_{\text{kin,rel}} \) convert kinetic energy into internal energy more efficiently.

Several reasons can explain differences of energy transfer efficiency between gases. Monoatomic gases are more efficient for ion heating than polyatomic gases because the latter can take away part of the total energy as vibrational and rotational energy.\(^{30}\) This explains that CO\(_2\) is causing less ion heating than argon even though it is slightly heavier. Argon is also causing more ion heating than helium, although both are monoatomic gases. This could be due to the higher polarizability of argon, which would either favor the formation of a sticky complex in which the total energy can be redistributed,\(^{19}\) and/or induce some argon acceleration towards the ion just before the collision.\(^{31}\) Note also that a higher actual \( E_{\text{kin,rel}} \) than predicted from \( v \) is expected, because the relevant kinetic energy for energy transfer it is the kinetic energy just before the collision and not the average kinetic energy. The magnitude of that deviation can also depend on the target gas. Overall, given the experimental errors of our study, only argon showed a significantly enhanced ion heating compared to helium, nitrogen and CO\(_2\). Xenon would be expected to induce even higher ion heating.
Figure 5: Influence of the nature of the gas on ion vibrational effective temperature $T_{\text{eff,vib}}$. (a) Comparison between nitrogen (N$_2$), carbon dioxide (CO$_2$) and argon (Ar) as a function of wave height. Gas pressure = 0.275 mbar, wave speed = 600 m/s, bias = 10 V. (b) Comparison between helium (He) and nitrogen (N$_2$) as a function of wave height. Gas pressure = 0.516 mbar, wave speed = 1500 m/s, bias = 24 V (He) – 10 V (N$_2$) (c) Correlation between the drift velocity squared $v^2$ and $T_{\text{eff,vib}}$ for N$_2$, CO$_2$, Ar and He. (d) Correlation between $E_{\text{kin,rel}}$ (Eq. 6) and $T_{\text{eff,vib}}$ for N$_2$, CO$_2$, Ar and He.
CONCLUSIONS

Our results demonstrate a significant field heating of the p-methoxybenzylpyrdinium ion (m = 200 Da, z = 1) inside the TW IMS. We obtained vibrational effective temperatures $T_{\text{eff},\text{vib}}$ ranging between 500 K and 800 K. The temperature can be decreased by lowering the wave height, increasing the gas pressure, or increasing the wave velocity. However, to obtain reasonable ion mobility separation in this mass range on a first-generation Synapt HDMS mass spectrometer, the effective temperature cannot be decreased below 500 K. Preliminary results on a second-generation Synapt G2 HDMS mass spectrometer, operated at higher pressure and wave height than the Synapt G1 HDMS, show effective temperatures ranging between 600K and 700K (manuscript in preparation), suggesting that this is a general feature of travelling wave ion mobility spectrometers.

The fact that $T_{\text{eff},\text{vib}}$ is higher than the gas temperature has several implications:

(1) The evolution of $T_{\text{eff},\text{vib}}$ with the wave height, gas pressure and wave speed highlights that $T_{\text{eff},\text{vib}}$ is related to $v$ (the ion speed in the laboratory frame of reference, see Eq. 15). In that respect, the study of $T_{\text{eff},\text{vib}}$ also reveals important aspects of the ion motion in TW IMS. The wave speed $s$ affects $T_{\text{eff},\text{vib}}$, and thus the drift speed $v$ of ions, although it has no impact on $K$ or $E$. This reveals that ions cannot instantaneously accommodate for the changes of electric field induced by the waves.

(2) Collisions in the TW IMS are inelastic, i.e. they result in a transfer of relative translation energy into internal energy of the analyte. The collision regime is therefore different in TW IMS than in DT IMS. In DT IMS, ion motion in the direction of the electric field is negligible compared to thermal motion and the ions are at the gas temperature. In TW IMS, the field contribution to ion velocity becomes significant compared to the thermal contribution. This also implies that the ion transport
regime is different in TW IMS than in DT IMS, with a switch to “high field” conditions caused by the higher $E/N$ ratios at stake in the traveling wave ion guide.

(3) Stating the obvious is to say that, if collisions are inelastic, it means that the colliding partners do not behave as hard spheres. This is important in the interpretation of the collision cross section $\Omega$ itself, which is actually a momentum transfer collision integral. When the ion-gas collision model deviates from the hard spheres limit, ion-gas interaction potentials come into play, and the collision cross section itself becomes dependent on the temperature.$^{32}$ In other words, $\Omega$ should become $\Omega(T_{\text{eff}})$ if field heating is significant.

(4) The fact that $T_{\text{eff,vib}}$ is higher than the gas temperature also implies that the effective temperature in the Mason-Schamp equation, $T_{\text{eff}}$ (Eq. 2), is higher than the gas temperature. Therefore, the expected dependence of $T_{\text{eff}}$ with the ion mass and charge has also important implications on the collision cross section calibration. Indeed, the usual calibration procedure consists in measuring the drift time $t_d$ in TW cells for ions of known $\Omega$. The collision cross section is corrected to account for both $q$ and $\mu$. However, the fact that the ion mobility depends on $T_{\text{eff}}$, and that $T_{\text{eff}}$ could change from ion to ion, is not currently taken into account in the calibration procedure. This may explain why calibration with denatured proteins (high charge-to-mass ratios) is not reliable for native proteins (low charge-to-mass ratios).$^{15}$ they do not have the same effective temperature. Future work will be devoted to take the effective temperature into account for collision cross section calibration of TW IMS. Meanwhile, it is advisable to calibrate TW IMS with ions of similar masses and charges as the analyte.

An important question is how the effective temperature changes with the nature of the ion. In the present study, we investigated the field heating of a single analyte ($m = 200$ and $z = 1$) and demonstrate a strong correlation between $T_{\text{eff,vib}}$ and $v$. This last parameter is related to the ion mobility $K$ (Eq. 1) which contains all the ion characteristics required for comparison with other molecules (electric charge
\( q \), reduced mass \( \mu \), and collision cross section \( \Omega \). Therefore, based on Equation (2), we can anticipate that larger ions should, at equivalent charge, be less heated than small entities as their progression in the gaseous medium is slower. However the mobility increases with the charge, and therefore multiply charged ions of large molecules could still be significantly heated. Further experiments need to be carried out in order to confirm these trends.

To conclude, the use of TW IMS for structural studies must be considered with care, because the ion heating could in some instances be significant enough to alter ion structures while they are crossing the mobility cell. This is a disadvantage if the goal is to preserve the solution-phase structures, and if these are unstable at higher internal temperatures. Relatively fragile small molecules (like the benzylpyridium ion) or non-covalent complexes observed at relatively high \( z/\Omega \) ratios are more likely to be concerned than large native protein complexes, typically analyzed at relatively low charge states compared to their size. On the other hand, field heating of the ions could also be exploited by modulating the T-wave instrumental parameters, as a convenient way of changing the ion temperature and exploring conformational dynamics.

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Supporting Information Available. Correspondence between Pirani and capacitance gauge pressure measurements, details of vibrationnal effective temperature calculation, derivation of Equation (5), detailed calculation of \( FY \) before, inside and after IMS, additional data as described in text, are available at http://pubs.acs.org.
REFERENCES


