AMMONIA AS A MODIFIER IN ION MOBILITY SPECTROMETRY: EFFECTS ON ION MOBILITIES AND POTENTIAL AS A SEPARATION TOOL

ROBERTO FERNÁNDEZ-MAESTRE\textsuperscript{1,1}, CHING WU, AND HERBERT H. HILL JR.\textsuperscript{3}

\textsuperscript{1}Universidad de Cartagena, Campus de Zaragocilla, Programa de Química, Cartagena, Colombia
\textsuperscript{2}Excellims Corporation, 20 Main Street, Acton, MA, USA,
\textsuperscript{3}Department of Chemistry, Washington State University, Pullman, WA 99163, USA,

(Received: July 20, 2012 - Accepted: December 4, 2012)

ABSTRACT

A commercial ammonia solution was introduced in the buffer gas of an ion mobility spectrometer and the mobilities of tetramethylammonium (TMA), tetraethylammonium (TEA), tetrapropylammonium (TPA), and tetrabutylammonium (TBA) ions, trizenylamine (TBAz), tributylamine (TBA), 2,4-dimethyl pyridine (2,4-lutidine), 2,6-di-tert-butyl pyridine (DTBP), serine, atenolol, and valinol decreased depending on their structures. Electrospray ionization-ion mobility spectrometry-quadrupole mass spectrometry was used in these experiments. Analyte ion mobilities decreased to different extents with the amount of ammonia introduced in the mobility spectrometer. When the amount of ammonia increased from 0.0 to 22 mmol m\textsuperscript{-3} (with a concomitant concentration of water of 106 mmol m\textsuperscript{-3}), percentage reductions in mobilities were: -4.8% (serine), -1.9% (reactant ions), -1.1% (TBA), -0.9% (TBAz), -0.5% (TEA), -0.4% (valinol and TBA), -0.3% (TEA, TMA, and 2,4-lutidine), and 0% (atenolol and DTBP). These selective variations in mobilities were due to formation of large analyte ion-ammonia-water clusters. The small change in mobility of tetralkylammonium ions, TBA, TBA, atenolol, and DTBP with the introduction of ammonia into the buffer gas was explained by steric hindrance of bulky substituents which shielded the positive charge of the ion from the attachment of ammonia and water molecules, and delocalized the positive charge. Ammonia in the buffer gas produced ion clusters with one or two ammonia molecules in compounds with little steric hindrance such as serine. At concentrations of ammonia of 4.4 mmol m\textsuperscript{-3} or higher (at 150°C), ligand-saturation with ammonia and water molecules occurred on the positive charge of some ions; when the positive charge saturated, no significant reductions in ion mobility occurred when increasing the concentration of ammonia in the buffer gas.

Keywords: Ion mobility spectrometry, gas modifier, ammonia, clustering, dopant

INTRODUCTION

IMS is an analytical technique that separates gas-phase ions moving in a drift tube under an applied electric field based on their size to charge ratios. IMS was introduced in 1970 by Karasek as detector for volatile organic molecules,\textsuperscript{1} and it became the analytical technique preferred for the analysis of explosives,\textsuperscript{2} illicit drugs,\textsuperscript{3,4} and chemical warfare degradation products.\textsuperscript{5,6} IMS has been used as a detector for liquid,\textsuperscript{7} gas,\textsuperscript{8} and supercritical fluid chromatography.\textsuperscript{9} IMS has also been applied to the determination of nicotine in tobacco,\textsuperscript{10} microorganisms,\textsuperscript{11} analysis of protein structures,\textsuperscript{12} study of the three-dimensional shape of polyatomic ions,\textsuperscript{13,14} and the separation of isomeric peptides.\textsuperscript{15} IMS has given an additional dimension of analysis to investigations by mass spectrometry; when coupled to mass spectrometry, IMS has been applied to metabolomics,\textsuperscript{16} and proteomics.\textsuperscript{17}

When electrospray is used as the ionization source in IMS, the solvent molecules are stripped from the ions in a desolvation region by means of a heated counter-current flow of nitrogen buffer gas. This desolvation process produces single stable gas phase ions which are then pulsed by an ion gate into a drift region. The ions in the drift region are decelerated by repeated collisions against the buffer gas but are simultaneously accelerated by the electric field. This combination of decelerations and accelerations rapidly thermalizes the ions, and averages their velocities to a characteristic value that depends on their collision cross sections. These velocities can be used to calculate mobility constants, \( K \), a characteristic parameter of the ions:\textsuperscript{18}

\[ K = \frac{v}{E} = \frac{L^2}{V t_d} \]  

where \( v \) is the velocity of the ion in cm s\textsuperscript{-1}, \( E \) the electric field in the drift region in V cm\textsuperscript{-1}, \( L \) the distance in centimeters the ion travels from the ion gate to reach the detector, \( F \) the total voltage drop in volts in the drift region, and \( t_d \) the time the ion takes to travel the distance \( L \) in seconds. If the electric field is less than ~500 V cm\textsuperscript{-1} (at atmospheric pressure), \( v \) should be a linear function of \( E \).\textsuperscript{19} To compare values of \( K \) in different experimental conditions, ion mobilities must be normalized to standard conditions. This normalization yields reduced mobilities (in cm\textsuperscript{3}V\textsuperscript{-1}s\textsuperscript{-1}) which are characteristic of every compound in a given buffer gas:

\[ K_0 = \frac{K P}{760 T} \]  

where \( P \) is the pressure in Torr and \( T \) the temperature of the buffer gas in Kelvin.\textsuperscript{19}

Ion mobility spectrometry using buffer gas modifiers separates ions in a gas phase by exposing ions to a polar modifier.\textsuperscript{20,21} These modifiers enhance ion-molecule interaction forces. When polar modifiers are introduced into a buffer gas, they interact with the analytes ions depending on the ion structures and steric hindrance of the atoms bearing the charges; this interaction leads to the formation of clusters of different sizes that spend different times inside the drift tube. These clusters may be used to separate analytes that overlap in IMS. In a recent work, water did not significantly change the ion mobilities of analytes when it was introduced into the buffer gas;\textsuperscript{21} and it was uncertain if it was because of its size; the subject of this investigation was to investigate another low molecular weight modifier, ammonia, to compare the changes in ion mobilities with those of water when they are introduced into the buffer gas and demonstrate that the small changes in mobilities with small modifiers are due to their sizes; also, to determine the capabilities of ammonia in generating selective ion separations.

EXPERIMENTAL SECTION

Instrument. Experiments were carried out on an ion mobility spectrometer coupled to a quadrupole mass spectrometer through a 40-µm pinhole (Figure 1). The IMS instrument used an electrospray ionization source at atmospheric pressure. Routinely used operating conditions for this instrument were: sample flow, 3 µl min\textsuperscript{-1}; drift tube length, 25.0 cm; ionization region length, 7.5 cm; ESI voltage, 15.6 kV; voltage at first ring, 12.1 kV; voltage at the gate, 10.80 ± 0.01 kV; gate closure potential, ±40 V; gate pulse width, 0.1 ms; scan time, 35 ms; averages for every spectrum, 1000; pressure, 680-710 Torr; buffer gas, nitrogen; buffer gas flow, 0.93 liter min\textsuperscript{-1}; buffer gas temperature, 150 ± 2°C; ammonia flow rate, 0.17 to 0.75 µl min\textsuperscript{-1}.
recommend correcting reduced mobilities by comparing with K_{orthogonal} field stopped positive or negative ions approaching the IMS spectrometer are set to allow only ions with a specific mass or a range of masses to enter the mass spectrometer continuously, without pulsing the IMS, and are identified by the drift time in ms. SIM-IMS was performed to the analyte peaks and their clusters with ammonia. Analytes were identified by comparing their protonated molecules or clusters. Also, reduced mobilities of protonated ammonia, atenolol, serine, valinol, 2,4-dimethylpyridine (2,4-lutidine), 2,6-di-tert-butyl pyridine (DTBP), tributylamine (TBA), tributylphosphine oxide (TBPO), tetramethylammonium (TMA), tetraethylammonium (TEA), tetrapropylammonium (TPA), and tetrabutylammonium (TBA) chlorides (ACS reagent grade, ≥98% purity) were purchased from Sigma Aldrich Chemical Co. (Milwaukee, WI). These reagents were selected as analytes because they supply a series of ions with different molecular weights and steric properties needed to test the effects of size and steric hindrance on the change in mobilities of these molecules with the introduction of ammonia into the buffer gas. Ammonia was used here because in a previous study, water, a molecule of similar size, weight and polarity, was found to interact just slightly with several compounds. In this study, we investigate if the effects of ammonia in ion mobility are similar to those of water when they are introduced into the buffer gas of a mobility spectrometer to find generalizations on the effects of small modifiers.

Sample preparation and introduction. Analytes were prepared at a concentration of 50 μM in ESI solution (47.5 % methanol; 47.5 % water; 5 % acetic acid). Liquid samples or blank solutions (ESI solution) were infused continuously by electrospray ionization using 250 μl syringes (Hamilton, Reno, NV) at a flow rate of 3 μl min^{-1} into a 30 cm long, 100 μM ID capillary (Polymer Micro Technologies, Phoenix, AZ). This capillary was connected to a 50 μm ID silica capillary through a stainless steel union (Valco, Houston, TX). The end tip of this capillary was centered at a target screen, placed at the entrance of the mobility spectrometer. The target screen was made out of 2-mm stainless steel mesh and it had a 0.5-cm round hole in the center. A high voltage of 15.6 kV (for a 3.5 kV bias between the capillary tip and the target screen at the first ring) was applied to the stainless steel union to electrospray positive ions. Different syringes and capillaries were used for every compound, whenever it was possible, to prevent cross contamination between the analytes.

Ammonia introduction. Ammonia modifier was introduced into the buffer gas at concentrations of 0.0, 0.19, 0.89, 4.4, and 22 mmol m^{-3}. Ammonia was introduced with gas tight syringes (pumped by a KD Scientific pump, model 210) to avoid leaking. The syringes were connected to 10-cm-long, 50-μm ID silica capillary and this capillary was coupled to a T-junction into the buffer gas line before the buffer gas heater (Figure 1). The introduction of ammonia before the buffer gas heater provided a longer path to obtain a homogeneous mixture of ammonia with the buffer gas. To help vaporize the modifier, the temperature of the T-junction was increased to approximately 150 °C using a heating tape (OMEGA Engineering, Stamford, CT). Ammonia flow was countercurrent to the drift of ions to increase the interactions with them. The introduction of ammonia at the end of the drift tube allows a mixture with the buffer gas that is more homogeneous than that obtained when it is introduced with the carrier gas as in experiments when IMS was introduced in the 70's.

Identification of compounds. All analytes were detected as protonated mono- or di-cationic species, [M+H]^+ or [M+2H]^2+, or their clusters with ammonia. Analytes were identified by comparison of their m/z ratios in mass spectrometry to the molecular weight of their protonated molecules or clusters. Also, reduced mobilities of protonated ammonia and ammonium were compared to those from literature for further identification. Additionally, SIM-IMS was performed to the analyte peaks and their clusters to identify the peaks in the IMS spectrum; when SIM-IMS is applied, only one peak appears in the IMS spectrum.

Calibration. To account for errors in measuring instrumental parameters, Eiceman et al. recommend correcting reduced mobilities by comparing with standards:

\[
K_{o (unknown)} / J_d (unknown) = t_d (standard) / K_{o (standard)}
\]

where \(K_o\) is the reduced mobility in cm^2V^{-1}sc^{-1} and \(t_d\) the drift time in ms. A recently proposed IMS calibration method was used. This method uses DTBP as the chemical standard to calibrate the instrument by replacing their drift time and mobility value in Equation 1. This method also uses 2,4-lutidine to determine the presence of contamination in the buffer gas.

RESULTS AND DISCUSSIONS

Table 1 summarizes the percentage reduction in mobilities (%ΔK_o) for the analytes with the introduction of ammonia in the buffer gas. %ΔK_o was defined as the percentage difference between \(K_o\) in N_2-only buffer gas and \(K_o\) when ammonia modifier was introduced into the buffer gas at a given concentration. When ammonia concentration was increased from 0.0 to 22 mmol m^{-3} at 150 °C, %ΔK_o values were: -1.9% (reactant ions), -0.9% (TBA), -1.1% (TBA), 0.89% (TBA), and -0.4% (TBA).
-4.8% (serine), -0.4% (valinol), -0.4% TBA, -0.5% (TEA), -0.3% (TEA, TMA, and 2,4-lutidine), and 0% (atenolol and DTBP) (Table 1). In Table 1, %\( \Delta K \) values were not statistically different for TBA and TBzA. %\( \Delta K \) values for TBA, TMA, TPA, 2,4-lutidine, atenolol, DTBP, and valinol ions, were not statistically different from 0. Only differences greater than 0.5% were considered significant; this value was calculated from the maximum relative standard deviation of the drift times, 0.07 ms.

For selected analytes with nitrobenzene (at 0.95 mmol m\(^{-3}\)), \( ^{28} \) ethyl lactate (at 1.7 mmol m\(^{-3}\)), \( ^{22} \) 2-butanol (at 6.8 mmol m\(^{-3}\)), \( ^{22} \) a-trifluoromethyl benzyl alcohol (at 2.3 mmol m\(^{-3}\)), \( ^{22} \) and methyl 2-chloropropionate (at 0.93 mmol m\(^{-3}\)) \(^{22} \) modifiers in the buffer gas, the average %\( \Delta K \) values were: valinol (-20%), 2,4-lutidine (-13%), and serine (-22%). These %\( \Delta K \) values are large compared to those of the same analytes in water or ammonia/water at much larger modifier concentrations (Table 1). This result indicates that to obtain large changes in \( K \) values large modifiers must be used.

Table 1. %\( \Delta K \) values when ammonia and moisture were introduced into the buffer gas. Percent reduction in mobility, %\( \Delta K \), for selected compounds at a concentration of 176 mmol m\(^{-3}\) of water in the buffer gas (third column) \(^{1} \) and a mixture of water at 106 mmol m\(^{-3}\) and ammonia at 22 mmol m\(^{-3}\) (fourth column). %\( \Delta K \) values for water at 7.0 mmol m\(^{-3}\) in the buffer gas were 0.0 indicating a small influence of moisture in ion mobilities. Differences of less than 0.5 units in %\( \Delta K \), were not significant and may arise from the standard deviation of the drift time measurements in this work (0.07 ms); for the same reason, %\( \Delta K \) values of -0.5% or less were considered zero.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Molecular weight, amu</th>
<th>Water(^1)</th>
<th>Ammonia/water</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reactant ions</td>
<td>19, 37, 55, 73, 91...</td>
<td>-1.1</td>
<td>-1.9</td>
</tr>
<tr>
<td>2,4-lutidine</td>
<td>107.1</td>
<td>-1.2</td>
<td>-0.3</td>
</tr>
<tr>
<td>DTBP</td>
<td>191.3</td>
<td>-0.6</td>
<td>0</td>
</tr>
<tr>
<td>TBA ions</td>
<td>242.5</td>
<td>0</td>
<td>-0.4</td>
</tr>
<tr>
<td>TEA ions</td>
<td>130.3</td>
<td>-0.8</td>
<td>-0.5</td>
</tr>
<tr>
<td>TMA ions</td>
<td>74.2</td>
<td>-1.5</td>
<td>-0.3</td>
</tr>
<tr>
<td>TPA ions</td>
<td>186.4</td>
<td>-0.6</td>
<td>-0.2</td>
</tr>
<tr>
<td>TBzA</td>
<td>287.4</td>
<td>0</td>
<td>-0.9</td>
</tr>
<tr>
<td>Serine</td>
<td>105.1</td>
<td>-4.7</td>
<td>-4.8</td>
</tr>
<tr>
<td>Valinol</td>
<td>103.2</td>
<td>-1.1</td>
<td>-0.4</td>
</tr>
<tr>
<td>Atenolol</td>
<td>266</td>
<td>-</td>
<td>0</td>
</tr>
</tbody>
</table>

1. Ammonia and the reactant ions

The ion mobility spectrum of the electrosprayed solvent is shown in Figure 2a without introducing ammonia in the buffer gas. IMS spectra were obtained by pulsing the ion gate and setting the mass spectrometer to let all ions pass (radiofrequency-IMS mode). In this spectrum, the main reactant ion peak was observed at a drift time of 13.81 ms which corresponded to a reduced mobility value of 2.50 cm\(^2\) V\(^{-1}\) s\(^{-1}\). When the ion gate was held open to obtain mass spectra (MS mode), the main ions peaks were (H\(_2\)O)\(_2\)H\(^+\) at m/z 19, 37, 55, 73, and 91 as shown in Figure 2b; these peaks might have small contributions of ammonia ions, (NH\(_3\))\(_2\)H\(^+\), at m/z 18, 35 etc. The MS spectrum shows that the instrument was clean because no significant peaks appeared above 91 amu; it was important to control contamination because contaminants cluster to ion analytes ions and change their mobilities. \(^{21} \) These water clusters combined into one mobility peak in the IMS spectrum at 13.81 ms due to the equilibria (H\(_2\)O)\(_2\)H\(^+\) ↔ (H\(_2\)O)\(_{con}\)H\(^+\) + mH\(_2\)O.

19…

Figure 2. IMS (a) and MS spectra (b) of the ESI solvent in pure nitrogen.

The reactant ion peak, (H\(_2\)O)\(_2\)H\(^+\), showed at 13.81 ms (2.50 cm\(^2\) V\(^{-1}\) s\(^{-1}\)) in (a); the MS spectrum shows that the instrument was clean.

Figures 3a and 3b show the spectra of the reactant ions when 22 mmol m\(^{-3}\) of ammonia was added to the buffer gas; even at the maximum concentration of ammonia used, predominant reactant ions were water clusters at 37, 55, 73, 91, and 109 amu due to the high concentration of water in the mixture introduced in the buffer gas and the presence of water cluster ions from the ESI solution; however, these peaks might have contributions of ammonia clusters at 35, 52, 69, and 86 amu; water concentration was higher because the modifier was 29% NH\(_3\) in water; water concentrations of 0.90, 4.2, 21, and 106 mmol m\(^{-3}\) were concomitant to ammonia concentrations of 0.19, 0.89, 4.4, and 22 mmol m\(^{-3}\), respectively; in the IMS spectra in Figure 3a, the major reactant ion occurred at a drift time of 14.08 ms which corresponded to cluster ions of water and ammonia, with a reduced mobility of 2.45 cm\(^2\) V\(^{-1}\) s\(^{-1}\), (H\(_2\)O)\(_2\)(NH\(_3\))\(_n\)H\(^+\).

Figure 3. IMS (a) and MS spectra (b) of the ESI solvent (reactant ions) when 22 mmol m\(^{-3}\) of ammonia was added to the buffer gas. The reactant ion peak, (H\(_2\)O)\(_2\)H\(^+\), appeared at 14.08 ms in (a). The IMS peaks broadened and their mobilities decreased (from 2.50 in Figure 2 to 2.45 cm\(^2\) V\(^{-1}\) s\(^{-1}\)) due to interactions with the modifier mixture of ammonia and water.

2. Drift region equilibria

The ion mobility peak of serine was observed at 18.47 ms when a 50 \( \mu \)M solution of the amino acid was electrosprayed into the IMS-MS, with no modifier introduced into the buffer gas (Figure 4a). When ammonia was added to the buffer gas at concentrations of 0.19 and 0.89, mmol m\(^{-3}\) (with concomitant water concentrations of 0.90 and 4.2 mmol m\(^{-3}\)) the mobility of serine ions decreased from 1.856, to 1.782, and 1.760 cm\(^2\) V\(^{-1}\) s\(^{-1}\), respectively (Figures 4a-4c). Figure 4d is the mass spectrum of serine at 0.89 mmol m\(^{-3}\).
clusters of Ser(NH\(_3\))H\(^+\) and Ser(NH\(_3\))\(_2\)H\(^+\). It is noteworthy that ammonia was more efficient than water in clustering to serine, contrary to what happened when clustering to reactant ions (Figure 2). Further investigation is needed to explain this finding. Even if the mass spectra showed two ion clusters and the protonated molecular ion, the ion mobility spectrum contained a single peak for serine. One single peak means that clustering-declustering reactions reached equilibrium in the buffer gas at rates rapid enough to yield only one ion mobility peak with a weighted average of the mobilities of the individual ions;analyte ions traveled across the drift tube rapidly interconverting into the analyte-modifier clusters Ser(NH\(_3\))H\(^+\) and Ser(NH\(_3\))\(_2\)H\(^+\), back and forth. Therefore, the following equilibria may have occurred in the drift region between the protonated molecules of serine, ammonia, water, and serine-ammonia-water clusters (only ammonia molecules are included):  

$$\text{serine} + \text{ammonia} \rightarrow \text{serine-ammonia clusters}$$

This clustering with ammonia occurred due to the formation of hydrogen bonds between the electronegative nitrogen atom in ammonia and the partially positive charge on the amine hydrogen atoms of serine; clustering with water occurs through the formation of a hydrogen bond with the oxygen atom in water; clustering with ammonia and water stabilized the positive charge by sharing it with the nitrogen atom. As a result of these equilibria, the ion mobility of serine increased with increasing concentrations of ammonia in the buffer gas due to the increasing size of the cluster. The intensity of every ion species in the mass spectrometer was an indication of the relative concentration of ammonia in the buffer gas. At 0.89 mmol m\(^{-3}\), the intensity of the peaks decreased in the order: (Ser)H\(^+\), Ser(NH\(_3\))H\(^+\), Ser(NH\(_3\))\(_2\)H\(^+\) indicating a similar decrease in these cluster ion concentrations in the buffer gas; a concentration of (Ser)H\(^+\) larger than those of the clusters together with the absence of clusters with more than two ammonia molecules indicates that the buffer gas was not saturated with ammonia or that ammonia did not effectively clustered to serine. The clusters formed in the electrospray ion source, and they were in equilibria with ammonia and water molecules while traveling the drift tube because serine’s drift time increased as ammonia concentration increased, and the IMS peaks were well defined. The maximum number of ammonia molecules clustering with serine was two; this number is smaller than the eleven water molecules that attached to a similar ion, valinol, in a previous experiment maybe because the concentration of water was approximately 40 times larger before\(^{21}\) and water occupies part of the available places for attachment. In the experiments with water, %DK\(_v\) values at 7.0 mmol m\(^{-3}\) of water in the buffer gas were 0.0 indicating a small influence of moisture in ion mobilities and that the ion mobility reductions observed when ammonia was introduced into the buffer might be especially due to ammonia clustering. This is not contradictory with water clusters present in figures 2-4; because ammonia is a stronger base than water, ammonia clusters with serine better than water even if water was present at high concentrations as is demonstrated by the water clusters observed in figures 2-4.
3. Effects of ion structure on clustering

%ΔK<sub>v</sub> values appeared to depend on the size of the substituents on the ion charge. %ΔK<sub>v</sub> values were small or negligible for tetraalkylammonium, 2,4-lutidine, and DTBP ions which can be explained by steric hindrance on the charge; tetraalkylammonium ions have four bulky alkyl substituents that hinder the attachment of ammonia to the positive nitrogen as has been demonstrated before. The 3D models of TMA, DTBP, and 2,4-lutidine illustrate the steric hindrance in these ions (Figure 5). Figure 6a shows only a single peak for every tetraalkylammonium ion in the IMS spectrum of a mixture of these compounds when 22 mmol m<sup>-3</sup> of ammonia were introduced into the buffer gas and clusters with ammonia or water were absent; also, the mass spectrum of tetraalkylammonium ions exhibits only single peaks for these compounds up to 22 mmol m<sup>-3</sup> of ammonia in the buffer gas at 150 °C (Figure 6b) indicating the absence of ammonia or water clusters. Tetraalkylammonium ions appeared at 15.99 ms (TMA), 19.54 ms (TEA), 23.97 ms (TPA) and 28.35 ms (TBA) in the IMS spectrum and at m/z 75.2 (TMA), 131.2 (TEA), 186.4 (TPA) and 243.5 (TBA) in the mass spectrum. In DTBP, the hindrance was produced by the two tert-butyl groups and in 2,4-lutidine by the two methyl groups; in both compounds the charge is on a bulky aromatic ring and surrounded by alkyl groups (Figure 5).

![Figure 6. IMS (a) and MS spectra (b) of non-clustering compounds.](image1)

50-μM solutions of tetraalkylammonium ions, when 22 mmol m<sup>-3</sup> of ammonia were added to the buffer gas. Tetraalkylammonium ions have four bulky alkyl substituents that hinder clustering and only a single peak for every ion indicating the absence of ammonia clusters; tetraalkylammonium ions appeared at 15.99 ms (TMA), 19.54 ms (TEA), 23.97 ms (TPA).

![Figure 7. Effect of ammonia concentration in the buffer gas in ion mobility.](image2)

Although, size is proportional to mobility, atenolol (266.3 g mol<sup>-1</sup>) was unaffected by ammonia in the buffer gas while the ion mobility of another amine of similar weight, tribenzylylamine (287.4 g mol<sup>-1</sup>), decreased; this was because the size of atenolol is much smaller due to the formation of an intramolecular bridge that reduces atenolol size; this bridge also delocalizes the positive charge which decreases clustering and increase ion mobility.

In general, the changes in mobility with the introduction of modifier into the buffer gas were selective and dependent on the analyte structure. %ΔK<sub>v</sub> values decreased with molecular weight of the analyte; this trend may be due to the small effect on ion size when a molecule of ammonia clusters to large molecules. The selectivity in the reductions in ion mobilities might be applied to the separation of ions, such as valinol and serine; these ions overlap in the IMS spectra in N<sub>2</sub>-only buffer gas but they reach a separation of approximately 0.1 units of %ΔK<sub>v</sub> (or one ms) when ammonia is introduced in the buffer gas at a concentration of 0.89 mmol m<sup>-3</sup> enough to separate them with the resolution of the instrument (Figure 7).

4. Modifier Saturation

The sites on an ion to attach ammonia molecules are limited. A decrease in ion mobility change as a function of modifier concentration indicates that the positive charge of analytes is becoming saturated with ligands. Figure 7 shows a flattening of mobility values at 4.4 mmol m<sup>-3</sup> of ammonia in the buffer gas for reactant ions and those analytes that were affected by ammonia introduction such as serine; this flattening indicated saturation with ligands of the hydrogen atoms on the positive charge of the analyte, available for binding with modifier molecules; this saturation, due to high concentrations of ammonia in the buffer gas, would decrease the binding of additional modifier molecules of ammonia to the analytes, and a smaller decrease in mobility would be obtained with increasing concentrations of ammonia modifier. Overloading with ammonia was evident in the presence of clusters of serine with one and two ammonia molecules (Figure 4b).

CONCLUSIONS

The aim of this research was to study the addition of buffer gas modifiers in IMS to determine the capabilities of ammonia in generating selective ion separations and to compare the changes in ion mobilities using ammonia with those of water when introduced into the buffer gas. The ion mobilities of several compounds decreased, while other compounds were unaffected, when ammonia was introduced into the buffer gas of an ion mobility spectrometer. The degree of ion mobility reductions for these compounds was also different; the reductions in mobility were produced by the increase in collision cross sections of analytes due to the formation of clusters between the analyte ions and ammonia and water. The structure and size of the analytes determined the extent of clustering in most cases; the mobility of tetraalkylammonium ions and DTBP were less affected by the presence of ammonia modifier because the formation of clusters of these compounds with ammonia was prevented by steric hindrance and also because of their large size; the mobilities of large compounds, such as tribenzylylamine and tributylamine were less affected by the attachment of ammonia molecules when compared to smaller compounds such as serine, because the collision cross section of ions of large size is less affected by formation of clusters; on the contrary, the mobilities of small molecules such as serine and the reactant ions were largely affected by clustering with ammonia. These differences in the reductions in ion mobilities might be applied to the separation of compounds which overlap in the IMS spectra in N<sub>2</sub>-only buffer gas, such as valinol and serine. The change in mobility values as a function of modifier concentration was found to reach a limit at high concentrations of ammonia due to ligand saturation of the hydrogen atoms available for binding on the positive nitrogen of the analytes. Finally, more evidence was obtained that indicate that low molecular weight modifiers produce small changes in the mobilities of ions due to their small effect on the collision cross sections of ions. Some questions remained such as why valinol did not show large changes in ion mobility as those of serine, a compound of similar size and structure.

REFERENCES
