Non-covalent interactions of alkali metal cations with singly charged tryptic peptides

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The complexes formed by alkali metal cations (Cat\(^+ = \text{Li}^+, \text{Na}^+, \text{K}^+, \text{Rb}^+\)) and singly charged tryptic peptides were investigated by combining results from the low-energy collision-induced dissociation (CID) and ion mobility experiments with molecular dynamics and density functional theory calculations. The structure and reactivity of \([\text{M}+\text{H}+\text{Cat}]^{+}\) tryptic peptides is greatly influenced by charge repulsion as well as the ability of the peptide to solvate charge points. Charge separation between fragment ions occurs upon dissociation, i.e. \(b\) ions tend to be alkali metal cationised while \(y\) ions are protonated, suggesting the location of the cation towards the peptide N-terminus. The low-energy dissociation channels were found to be strongly dependant on the cation size. Complexes containing smaller cations (\(\text{Li}^+\) or \(\text{Na}^+\)) dissociate predominantly by sequence-specific cleavages, whereas the main process for complexes containing larger cations (\(\text{Rb}^+\)) is cation expulsion and formation of \([\text{M}+\text{H}]^{+}\). The obtained structural data might suggest a relationship between the peptide primary structure and the nature of the cation coordination shell. Peptides with a significant number of side chain carbonyl oxygens provide good charge solvation without the need for involving peptide bond carbonyl groups and thus forming a tight globular structure. However, due to the lack of the conformational flexibility which would allow effective solvation of both charges (the cation and the proton) peptides with seven or less amino acids are unable to form sufficiently abundant \([\text{M}+\text{H}+\text{Cat}]^{+}\) ion. Finally, the fact that \([\text{M}+\text{H}+\text{Cat}]^{+}\) peptides dissociate similarly as \([\text{M}+\text{H}]^{+}\) (via sequence-specific cleavages, however, with the additional formation of alkali metal cationised \(b\) ions) offers a way for generating the low-energy CID spectra of ‘singly charged’ tryptic peptides.

Keywords: structure–activity relationship; peptide structure; molecular modelling; collision-induced dissociation; alkali metals

Introduction

Over 30% of known proteins are metalloproteins.\(^{[1,2]}\) Protein–metal complexation as well as metal cation effects on structure is an important aspect in many fundamental biological processes.\(^{[3,4]}\) Although protein–metal complexes tend to favour divergent metals (e.g. catalytic centres) there is greater evidence for the important role of alkali metal ions, e.g. activation of enzymes, cellular metabolism, stabilisation of protein structure, etc.\(^{[5]}\) Another important aspect of protein/peptide–alkali metal interactions is in analytical applications, where fragmentation of these complexes within a mass spectrometer can provide useful information on the amino acid sequence of the peptide/protein.\(^{[6,7]}\)

Mass spectrometry provides a gas-phase environment for studying the structure and properties of protein/peptide–alkali metal complexes. It can be argued that studying these complexes in the gas phase has little biological relevance. However, many structural features may be the same in the gas phase as in the solution phase, and there are some evidence that the gas-phase structure of peptide fragments may be utilised for the solution-phase relevant studies of the peptide folding.\(^{[8,9]}\) Furthermore, observed gas-phase properties can be correlated to the biologically observed properties as well as enabling the development of controllable and predictable fragmentation methods.\(^{[6]}\)

So far, the majority of the studies have focused on singly charged alkali cationised peptides (some of the examples can be found in Refs. [6–18]). Collision-induced dissociation (CID) of the singly charged alkali metal cationised peptides led to the sequential loss of amino acids from the C-terminus, i.e. C-terminal sequencing,\(^{[6]}\) with the exception of the peptides with acidic residues.\(^{[7]}\) Only a few studies have considered the behaviour of the doubly charged protonated and sodiated non-tryptic peptides.\(^{[19,20]}\) Charge separation between \(b\) and \(y\) ions was observed. The \(y\) ions were mainly observed as sodiated species, in agreement with the location of the sodium towards C-terminus of the peptide.

This study is focused on doubly charged, protonated and alkali cationised, tryptic peptides. The main reason for this is that the vast majority of mass spectrometry-based proteomic studies uses tryptic digests of protein mixtures. Tryptic digests, despite significant efforts in desalting, always contain some proportion of salt adducts. A strategy utilising those adducts and increasing the information yield of proteomic experiment would be very useful. Furthermore, in doubly charged protonated and alkali cationised tryptic peptides, the charged arginine or lysine residue on the C-terminus may push the alkali cation towards the N-terminus (because of Coulombic repulsion). This is expected to drive the
charge separation between fragment ions, i.e. y ions are expected to be protonated, whereas b ions are alkali metal cationised. Here we address several issues of both ‘pure’ and ‘applied’ interest – in particular, how different alkali metal cations interact with the singly charged tryptic peptide backbone, how that interaction is reflected in their gas-phase fragmentation and how that can be utilised, e.g. for fast differentiation between b and y ions.

**Methods**

**Mass spectrometry and ion mobility**

All compounds used in the experiments were obtained from Sigma–Aldrich (Dorset, UK) without further purification. Peptide samples were made up to 10 µM in 50:50:1 water/methanol/acetic acid and introduced into the ESI source by direct infusion at a flow rate of 100 µl h⁻¹. The relative abundance of some [M+H+Cat]⁺⁺ species, especially in a case of rubidiated species, was increased by the addition of the alkali metal acetate to the sample solution.

CID experiments were performed on the 9.4 T Apex III Fourier-transform ion cyclotron resonance (FT-ICR) and the Esquire3000 quadrupole ion trap (IT) mass spectrometers (Bruker Daltonics, Coventry, UK). Both were equipped with Apollo electrospray ionisation sources. Argon (FT-ICR) or helium (IT) was employed for collisional cooling and as a collision gas. Mass selection and CID were carried out in the cell by using standard isolation and excitation procedures. On the IT, precursor ions were excited with a resonant RF. The excitation time was 40 ms and the RF amplitude was in the range of 0.44–0.75 V. On the FT-ICR instrument, the sustained off-resonance irradiation (SORI) CID was used for 150–250 ms, with activation 500–1000 Hz below resonance frequency.

Peptide–water interaction experiments were screened using gas-phase ion/molecule reactions within ICR cell. Experimental procedure similar to the gas-phase H/D exchange measurements (for more details about the gas-phase H/D exchange measurements, see Ref. [21]) followed by CID was used. Briefly, ions formed by ESI were transferred into ICR cell, mass selected and allowed to react (10–20 s) with the background pressure of H₂O. H₂O gas was introduced via a leak valve into the cell and the pressure used in the experiment was around ~8 × 10⁻⁸ Pa at the ambient temperature of 294 K. After reaction delay, SORI CID was employed and mass spectra were taken.

Ion mobility experiments were performed on the Synapt High Definition mass spectrometer (Waters Corp., Manchester, UK), described in detail in Ref. [22]. Ions were formed by ESI (the source temperature was 353 K and the capillary voltage was set to 3.2 kV) and injected into the mobility region. The bath gas in the T-Wave ion mobility cell was nitrogen, and the pressure was 0.53 mbar. The T-Wave was operated at 300 ms⁻¹ and linearly ramped in amplitude from 5 to 20 V. Arrival time distributions were recorded by synchronising the orthogonal acceleration time-of-flight mass spectrometric acquisitions with the gated release of ions from the transfer T-Wave. Two hundred mass spectra were acquired for each packet of ions. Instrument control and data acquisition were carried out using MassLynx (v4.1) software (Waters Corp.).

**Molecular modelling and theoretical cross sections**

All theoretically studied tryptic peptide structures are assumed to be protonated on the arginine side chain (the most basic site), and the alkali metal cation was placed close to the centre of the peptide backbone. In the molecular dynamics (MD) calculations, 0.5 fs integration step was used, all non-bonded interactions were calculated and peptide atomic charges were estimated via semi-empirical AM1 method. A combination of quenched dynamics and simulated annealing was used to sample the potential energy surface. Simulations at the 800 K for 30 ps were followed by slow cooling to the 200 K over 10 ps, and geometry optimisation using a steepest descent approach. For some complexes with K⁺ and especially with Rb⁺, we experienced expulsion of the cation during the simulations. Consequently, a part of dynamic simulations was carried out at 600 K. Two hundred structures were generated with this procedure for an each peptide variant. The structures within ~41 kJ mol⁻¹ (10 kcal mol⁻¹) from the lowest energy conformer were selected as the candidate group. The theoretical cross section of an each structure was calculated using the SIGMA ‘project approximation’ algorithm with atomic radii determined from the ion-helium Lennard-Jones parameters.[23,24] Average cross section was obtained from the candidate group. The conformational stability of the lowest energy structure (representative structure) was then tested with a 1 ns MD simulation at 300 K.

In order to achieve a more accurate description of dissociation energies, representative structures were further optimised at the B3LYP/STO-3G level of theory. Single point B3LYP calculations with the 6-311G(d,p) basis set (for all atoms except the Rb cation) were used to obtain final energies. All MD simulations were performed using HyperChem (Waterloo, Ontario, Canada) with AMBER 99 force field, whereas quantum mechanics density functional theory calculations were performed using the GAUSSIAN 03.[25]

**Results**

Figure 1 represents an example of the effect of the sodium cation on the dissociation process. By considering the sequence of [Glu¹]-fibrinopeptide B (GluFib, EGVNDNEEGFFSAR), [M + H]⁺ is expected to produce fragments associated with the selective cleavage of the amide bond, C-terminal to the acid residue upon collisional excitation; the so-called ‘aspartic acid effect’ (i.e. y6, y7, y9 or y13). This is indeed observed in experiments where fragmentation result from the initial excitation of the whole desorbed ion population during the MALDI process.[26] However, if the molecular ion is selected and subjected to the low-energy CID, the main fragmentation path is the loss of water (Fig. 1(a)). Further excitation of [M + H – 18]⁺ produces fragments y6 and y9 which can be attributed to the aspartic acid effect. Observed fragment ions suggest that the loss of water is from the Glu¹ residue. Similar behaviour was observed for singly charged (no basic residues, proton on N-terminal nitrogen) peptides containing Glu at the N-terminus (Scheme 1(a)). In our case the ionising proton is most likely to be located on the arginine side chain suggesting that the neutral N-terminal Glu can exhibit expulsion of water, (Scheme 1(b)).

The addition of the sodium cation to [M + H]⁺ and subsequent collisional excitation of [M + H + Na]²⁺ produces y ions connected with the aspartic acid effect (Fig. 1(b)). However, the corresponding b ions also appear in the spectrum as [b8 – H + Na]⁺ suggesting charge separation due to the Coulombic repulsion during low-energy CID. Formation of [y13 + Na]²⁺ appears to be in contrast with latter, however, that is not the case as it will be explained in one of the following sections.
Scheme 1. Loss of water from N-terminal Glu residue.

To provide greater insight into the binding of the metal ion, ion mobility experiments and MD simulations were performed. Figure 2 illustrates assigned structures of [M + H]$^+$ and [M + H + Na]$^{2+}$ of GluFib. Ion mobility results for protonated GluFib indicate fairly wide distribution of populations (Fig. 2(a)) with maxima corresponding to the drift time of 8.87 and 9.46 ms. The structure with the larger cross section (corresponds to larger drift time) is depicted in Fig. 2(b). MD simulations suggest that the central part of the backbone folds into a stable loop solvating the Arg side chain making the C-terminus fairly rigid. The N-terminus, on the other hand, remains relatively flexible which might explain relative wide cross-section distribution. It is important to note that Glu$^1$ side chain is not sterically hindered or involved in any non-covalent interactions and can be easily activated. The remaining part of the population is represented by a more compact globular structure stabilised with multiple hydrogen bonds and capped with Arg side chain (Fig. 2(c)). Here, the Glu$^1$ side chain is involved in hydrogen bonding which most likely hinders activation.

The arrival time distribution of [M + H + Na]$^{2+}$ yielded a single population with the drift time of 4.83 ms (spectrum not shown). MD simulations suggest that the sodiated/protonated GluFib is influenced by Columbic repulsion between the protonated guanidinium group and the sodium cation causing Na$^+$ placement towards the N-terminus (Fig. 2(d)). The presence of the sodium cation causes rigidification of the N-terminus due to multiple electrostatic interactions of carbonyl oxygen atoms with the sodium cation. A number of side chain carbonyl oxygens (Asp, Glu, Asn residues) provide a fairly good charge solvation shell (five ligands of which four are from side chains) without the need for involving much of the peptide bond carbonyl groups.

Although the GluFib example illustrates how attachment of the sodium cation alters the gas-phase structure and the low-energy CID process, the effects of different alkali metal cations and peptide primary structure implications need to be determined.

**Effect of different alkali metal cations**

Figure 3 shows the CID fragment ion mass spectra of the alkali metal cationised osteocalcin fragments 7–19 (GAPVPYPDPLEPR). The lithiated and sodiated precursor ions dissociate exclusively by producing b and y type fragment ions associated with the aspartic acid effect. Potassiated osteocalcin dissociates predominantly by producing b and y ions, however, with additional metal ion expulsion (observed on the IT data, see Supporting Information). For rubidiated peptides, the loss of the alkali metal cation becomes even more pronounced. Similar, in a way consistent, behaviour was observed for cationised des-Arg-bradykinins, where Cs$^+$ adducts (in contrast to Na$^+$ and K$^+$) produced very simple post-source decay spectra dominated by metal ion signal.[15] These results suggest that the dissociation process is strongly dependant on cation size, with small alkali metal ions Li$^+$ and Na$^+$ complexing more strongly than the larger K$^+$ and Rb$^+$, in agreement with previous observations on similar systems.[17,29]

According to the ion mobility measurements and the theoretical modelling data, conformation of the osteocalcin does not change significantly with cation size (Table 1). Osteocalcin forms a tight globular structure with the predominant interaction between the alkali metal cation and peptide bond carbonyl groups. Charge solvation within the tight loop is a preferred binding site for all alkali metal cations (Fig. 3). Due to the larger ionic radius, K$^+$ and Rb$^+$ have five ligands involved in the coordination shell, whereas Li$^+$ and Na$^+$ have four. Li$^+$ with the smallest ionic binds most strongly to the osteocalcin, whereas larger cations bind less strongly (Table 1). Interestingly, estimated magnitude of K$^+$ and Rb$^+$ bond dissociation energies (100–150 kJ mol$^{-1}$) lies around
threshold for Asp effect, which is 100 – 130 kJ mol$^{-1}$ for unhindered Asp side chain. Thus, this comparison could explain why for rubidiated peptides metal expulsion becomes a dominant process.

Additionally, we wanted to check is there a change in the system ([M+H+Cat]$^{2+}$) behaviour as a consequence of the interaction with water medium. Interaction with water could induce either further stabilisation or a change in the peptide conformation (e.g. for some amino acids and peptides it is shown that one molecule of water may alter the relative stability of the conformers$^{[31–33]}$) and which may influence the reaction path of the molecule, e.g. the CID process. Hydration effects were screened using gas-phase ion/molecule reactions within the ICR cell, i.e. selected peptide ions were allowed to react with background pressure of H$_2$O and subsequently subjected to SORI CID. The SORI CID spectra produced were identical to those obtained without interaction with water, suggesting that the initial conformation is not significantly changed at least not in a way that will affect the dissociation process.

**Influence of peptide primary structure**

The results from a series of experiments (similar to those illustrated) performed on tryptic-like peptides are summarised in Table 2. The presented data set allows several conclusions and implications to be made. In general, charge separation between fragment ions (b ions are alkali metal cationised, whereas y ions are protonated) occurs upon fragmentation. Peptides with seven or fewer amino acids are unable to form sufficiently abundant [M+H+Cat]$^{2+}$ ions. For shorter peptide chain lengths, the lack of conformational flexibility which allows effective solvation of both charge points by carbonyl oxygen’s make charge–charge repulsion main reason why [M+H+Cat]$^{2+}$ ions are not formed. This observation is consistent with recent theoretical calculations on a series of oligoglycine ions which found that the formation of the [M + H + Na]$^{2+}$ complex is endothermic for peptides with five or less glycines.$^{[17]}$

Structural data obtained for GluFib and osteocalcin suggest a possible relationship between the primary peptide structure and the nature of the sodium coordination shell formed. Peptides with a significant number of side chain carbonyl oxygens (e.g. GluFib) can provide a good charge solvation shell without the need for involving much of the peptide bond carbonyl groups and thus necessitating the formation of a tight globular structure with the predominant interaction between alkali metal cation and peptide bond carbonyl groups (e.g. osteocalcin). However, a significant number of side chain carbonyl oxygen cannot exclude formation of tight structure due to effective salvation/neutralisation of the cation charge and thus allowing closer positioning of protonated C-terminus.

It is interesting to note that [b$_3$ − H + Na]$^{+}$ is the smallest b ion observed (Table 2). Despite, in some cases (PPGFSPFR and ADSGEGDFAEGGGVR), a prominent cleavage at the C-terminal side of the second amino acid residue, the corresponding [b$_3$ − H $^{+}$]...
Figure 3. SORI CID spectra together with low-energy conformations of doubly charged protonated/alkali metal cationised osteocalcin: (a) [M + H + Li]^{2+}, (b) [M + H + Na]^{2+}, (c) [M + H + K]^{2+} and (d) [M + H + Rb]^{2+}.

Analytical implications

In general, during tandem MS [M+H+Cat]^{2+} peptides exhibit behaviour like [M + H]^{+} species, i.e. sequence-specific cleavages, however, with the additional formation of the alkali metal cationised b ions (Table 1). The dissociation of [M+H+Cat]^{2+} ions offers a way to generate CID spectra of 'singly charged' tryptic peptides. This is especially useful in ESI which generally tends to form multiply charged ions. Furthermore, by combing information from CID spectra of singly (here [M+H+Cat]^{2+}) and doubly charged precursor ions in the same experiment, the benefits of the charge remote and the charge-directed cleavage can be exploited and possibly improve the identification process without increasing the workload. Slightly different coupling, spectra of peptides cationised with different alkali metal cations (as in Fig. 3), will allow us to distinguish between classes of ions (b and y) without any prior knowledge of the peptide sequence. When overlapping those spectra, y ions have the same mass whereas b ions have the mass shift in accordance with mass difference between corresponding cationising metals. The fact that in some cases large y ions tend to retain a cation can reduce the utility of the proposed experiment. However, only to a limited extend, cause y ions with retained cation will be doubly charged, thus shifting in half mass difference between corresponding cationising metals. However, with ESI as a method of choice for generating [M+H+Cat]^{2+} ions there could be some difficulties with obtaining high abundance of the specific metal cationised ion of interest. The presence of small amounts of metal salts in the electrospray
solution may significantly reduce a peptide ion abundance due to both the suppression of ionisation and the formation of multiple species having metal adducts. However, new methods for the controlled generation of a metal cationised peptide ions are emerging such as cation switching ion/ion reactions.\[34\]

**Conclusions**

The work presented here reveals some insights into the non-covalent interactions of tryptic peptides and alkali metal cations in the gas phase. Important findings can be summarised as follows:

The structure and reactivity of [M+H+Cat]2+ tryptic peptides are greatly influenced by charge repulsion as well as the ability of the peptide to solvate cations. Charge separation between fragment ions occurs upon dissociation, i.e. b ions tend to be alkali metal cationised whereas y ions are protonated, suggesting the location of the cation towards the peptide N-terminus.

Low-energy dissociation process is strongly dependant on the cation size. Smaller cation (Li+ or Na+) complexes dissociate by sequence-specific cleavages whereas complexes with larger cations (Rb+ or Na2+) dissociate by cation expulsion which is consistent with calculated bond dissociation energies.

Peptides with a significant number of side chain carbonyl oxygen’s can provide a good charge solvation without need for involving much of the peptide bond carbonyl groups and thus forming of a tight globular structure. [M+H+Cat]2+ peptides dissociate in similar manner as [M+H]+, i.e. following charge-remote dissociation pathways, offering a way to generate a low-energy CID spectrum of the ‘singly charged’ tryptic peptides.

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**Supporting information**

Supporting information may be found in the online version of this article.

**References**


**Table 2.** Fragment ions observed in low-energy CID spectra of [M+H]+ and [M+H+Na2]+

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Species</th>
<th>Fragment ions</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>b ions</td>
</tr>
<tr>
<td>AFLDASK</td>
<td>[M+H]+, [M+H+Na]+, [M+H+Na2]+</td>
<td>b7, b9, b2, b4, b5, b7, b8, b2, b3, b4, b5, b7, b8, a</td>
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<td>AFLNASR</td>
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<td>PPGFSFPR</td>
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<td>b7, b9, b2, b4, b5, b7, b8, a</td>
</tr>
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*a* [b]–[H + Na]+.  
*b* [y6 + Na]2+.  

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