

Application of Multiple Length Crosslinkers to the Characterization of Gaseous Protein Structure

Melanie Cheung See Kit¹, Ian K. Webb^{1,2*}

¹Department of Chemistry and Chemical Biology, Indiana University-Purdue University Indianapolis, Indianapolis, Indiana 46202, USA

²Center for Computational Biology and Bioinformatics, Indiana University School of Medicine, Indianapolis, Indiana 46202, USA

* Correspondence to: Ian K. Webb; e-mail: ikwebb@iu.edu

Keywords: Native mass spectrometry, Ion/Ion reactions, Ion mobility mass spectrometry, Crosslinking, Molecular Dynamics

Abstract

The speed, sensitivity, and tolerance of heterogeneity of native mass spectrometry, as well as the kinetic trapping of solution-like states during electrospray, makes mass spectrometry an attractive method to study protein structure. Increasing resolution of ion mobility measurements and mass resolving power and range are leading to the increase of the information content of intact protein measurements, and an expanded role of mass spectrometry in structural biology. Herein, a suite of different length noncovalent (sulfonate to positively charged side chain) crosslinkers was introduced via gas-phase ion/ion chemistry and used to determine distance restraints of kinetically trapped gas-phase structures of native-like cytochrome c ions. Electron capture dissociation allowed for the identification of crosslinked sites. Different length linkers resulted in distinct pairs of side chains being linked, supporting the ability of gas-phase crosslinking to be structurally specific. The gas-phase lengths of the crosslinkers were determined by conformational searches and density functional theory, allowing for the interpretation of the crosslinks as distance restraints. These distance restraints were used to model gas-phase structures with molecular dynamics simulations, revealing a mixture of structures with similar overall shape/size but distinct features, thereby illustrating the kinetic trapping of multiple native-like solution structures in the gas phase.

Introduction

Mass spectrometry (MS) has been used to study intact proteins since the advent of electrospray ionization (ESI) MS. The earliest experiments revealed two important fundamentals of the behavior of protein ions formed by ESI: (1) charge state distributions (CSDs) of proteins reflect their overall structure in solution, with higher CSDs reflecting unfolded structures and lower CSDs reflecting folded structures and (2) noncovalently bound protein complexes and porphyrin groups can be retained upon ionization.¹ Native mass spectrometry depends on the retention of important aspects of protein structure during ESI and transfer to the gas phase. Measurements from native MS, where ions are ionized in salty, aqueous conditions at physiological pH, have revealed that specific, noncovalent binding between proteins and ligands or proteins and other proteins can be retained in the gas phase.² Native MS has become a transformative technique for structural characterization of heterogeneous protein systems, allowing for intact proteins and protein complexes to be measured with speed, sensitivity, and selectivity³, thus expanding our knowledge of biological systems.⁴

Ion mobility mass spectrometry (IM/MS) measurements have shown that the collision cross sections (CCS) measured for globular proteins and protein complexes correlate strongly to X-ray and NMR structures.⁵⁻⁷ In addition, the number of unfolding domains for proteins and protein complexes in the gas phase is equal to the number of known folding domains in solution.⁸ Clearly, the overall size/shape and important inter-residue contacts⁹ can be maintained in the gas phase. Though the protein ions undergo a change in environment from solution to charged, evaporating droplets, to vacuum, it has been suggested that solution structures are “kinetically trapped,” and without significant activation, the equilibrium gas-phase unfolded structures are not observed.¹⁰⁻
¹¹ Kinetic trapping allows for the detection of folding intermediates which are not observable in solution,^{9, 12} a beautiful example of the versatility of structural MS measurements towards addressing the so-called “protein folding problem.”¹³

Not all aspects of protein solution structures are maintained upon ionization and transmission into the gas phase. Structures are known to compact in the gas phase. Compaction is the primary hypothesis behind why CCS measurements are systematically smaller than CCS values calculated by trajectory methods¹⁴ applied to molecular coordinates directly from high resolution solution structures. For example, molecular dynamics (MD) simulations predict that macromolecular rings will shrink into vacant cavities due to the lack of solvent.¹⁵ Charged side chains also collapse onto the protein surface for charge solvation by either carbonyl oxygens or acidic residues.¹⁵ Microsolvation studies of lysines in cytochrome c¹⁶ and ubiquitin¹⁷ with 18-crown-6 ether suggested that the effects of vacuum on charged side chains led to a reduction in CCS for folded proteins and an increase in CCS (i.e., destabilization) for unfolded proteins. These factors are combined with the possibility of pH changes, electrochemical processes, and changes in droplet temperature during electrospray.¹⁸⁻¹⁹

The gas-phase tools used to observe the preservation versus loss of solution-phase structure in native MS look at the averaged, overall structure that emerges from each charge state (e.g., gas-phase IR spectroscopy and IM).²⁰ These measurements do not provide localized structural details. For example, a single CCS can correlate to many possible solution structures.²¹ Although the native inter-residue contacts have been shown by molecular modeling to be maintained after electrospray,²² these details have been obtained only by molecular modeling approaches to which CCS and IM spectra are fit without additional experimental validation. The experimental measurements themselves do not elucidate these interactions, as CCS measurements, while useful, are a rotationally averaged measurement of the entire conformer/conformer family and cannot inform on localized three-dimensional structural information.

The combination of molecular modeling with experimentally distance restraints, from, for example, NMR,²³ single-molecule FRET,²⁴ and chemical crosslinking²⁵ and footprinting²⁶, has led to the successful determination of structures of natively folded proteins. This combination has

also been applied to proteins *in vacuo* by using radical-donor and radical-acceptor pairs in radical-directed dissociation (RDD)²⁷ as well as chemical crosslinking utilizing gas-phase reactions of oppositely charged ions²⁸⁻²⁹ (*i.e.*, ion/ion reactions). Previously, our group used the gaseous dianion of the sulfo-EGS (ethylene glycol bis(sulfosuccinimidyl succinate)) from negative mode electrospray ionization to provide distance constraints for the protein ubiquitin upon ion/ion reactions followed by electron capture dissociation.²⁸ Herein, we have expanded our panel of non-covalent crosslinkers ranging in size from 3.5 to ~20 Å to provide distance restraints used in gas-phase molecular dynamics simulations to characterize gas-phase structures of native-like cytochrome c ions. We expect that this development will be an important next step towards characterizing the changes in protein structure from solution to inside the mass spectrometer.

Materials and Methods

Materials and Sample Preparation. Cytochrome c from equine heart, ammonium acetate, potassium benzene-1,2-disulfonate (1,2-BDSA), 1,3-benzenedisulfonic acid disodium salt (1,3-BDSA), 1,5-naphthalenedisulfonic acid, disodium salt hydrate (1,5-NDSA), and pentadecafluoro-1-octanol (PFO) were obtained from Sigma-Aldrich (St. Louis, MO). Ethylene glycol bis(sulfosuccinimidyl succinate) (Sulfo-EGS), bis(sulfosuccinimidyl)suberate (BS3), bis(sulfosuccinimidyl) glutarate-d₀ (BS2G) disodium forms and Amberlite IR-120 (H) ion-exchange resin were purchased from ThermoFisher Scientific (Rockford, IL). Methanol, acetonitrile, and ammonium hydroxide were purchased from Fisher Scientific (Fairmont, NJ). Water was obtained from a Milli-Q Millipore A10 (Burlington, MA) water purification system at a resistivity of 18 MΩ or greater. The 7⁺ charge state of cytochrome c was prepared under aqueous native-like conditions at a final concentration of 10 μM in an aqueous 200 mM ammonium acetate solution, pH 7. The proton transfer reagent, PFO was electrosprayed at 1.25 mM in a 99/1 vol/vol mixture of methanol/ammonium hydroxide and the crosslinking reagents (1,2-BDSA, 1,3-BDSA and 1,5-NDSA) were first prepared at a concentration of 50mM in water and incubated with Amberlite cation exchange resin to facilitate ion exchange of Na⁺ and K⁺ and increase the relative

concentration of doubly-deprotonated form of the reagent. Following the ion exchange step, the stock solutions were diluted to a working concentration of 1mM using an 80/20 vol/vol mixture of acetonitrile/water. The remaining crosslinkers (Sulfo-EGS, BS3 and BS2G) were dissolved in acetonitrile at a final concentration of 1mM.

Ion/ion Reactions and Ion Mobility Mass Spectrometry. The experiments were performed on a Synapt G2-Si High Definition Mass Spectrometer (Waters Corporation, Wilmslow, U.K.), equipped with an electron transfer dissociation (ETD) glow discharge source, a NanoLockspray source, an external electrospray voltage control module (GAA Custom Electronics LLC, Kennewick, WA) and an ExD cell (e-MSion, Corvallis, OR). Instrumental details for the gas-phase ion/ion reactions³⁰⁻³¹ and electron capture dissociation (ECD) using the ExD cell³² have been described previously. The protein was electrosprayed in positive ion mode using the nanolockspray source via borosilicate tips, pulled using a P-1000 micropipette puller (Sutter Instruments, Novato, CA) while the anionic reagents were infused through the reference sprayer at a flow rate of 1 μ L/min. The external voltage control module was used to synchronize the source polarity (+0.90 kV and -0.90 kV) and ion injection times (1s each) to sequentially introduce the anions and cations into the instrument. Cytochrome c charge states and reagent ions (either deprotonated PFO or doubly deprotonated crosslinker) were isolated in the quadrupole and stored in the trap cell with a trap traveling wave height of zero volts to allow for gas-phase ion/ion reactions of specific protein charge states to occur. The ion/ion products were then separated by ion mobility (IM) using nitrogen as the mobility gas at a flow rate of 10 mL/min while the Trap, He and Transfer cell flow rates were set to 10, 10 and 7 mL/min, respectively. This lower pressure regime was used to maintain conditions in the trap cell conducive to stable ion/ion product formation, as previously published.³¹ A ramping traveling wave velocity of 750 – 2400 m/s over a full IMS cycle and wave height of 40 V were used in the mobility cell. After IM separation, the ions were subjected to ECD fragmentation (using 3.5 eV electrons) in the ExD cell (detailed voltage

settings included in Table 1 of supplementary info) and supplemental activation in the transfer cell (Transfer CE of 30V) before mass analysis in the time-of-flight (TOF) mass spectrometer set to 'Resolution' mode. The experiments were all acquired in triplicate runs to ensure reproducibility.

Data Analysis. The IM peaks of the 1x crosslinked adduct or the 2x proton transfer product (by PFO) were selected, and the m/z data extracted using MassLynx v4.2 (Waters Corp.). The data was smoothed and centered and a lockmass correction (2134.1621 or 2650.4683 m/z corresponding to z_{18}^+ and z_{23}^+ fragments respectively) was applied. The processed data was exported as .mgf (Mascot Generic Format) files, and c and z fragments were annotated using LCMS spectator³³ (<https://omics.pnl.gov/software/lcmsspectator>), using a mass error tolerance of 20 ppm³⁴ and a raw intensity threshold of 1000 counts. In addition to an acetyl group at the N-terminus (42.01056 Da) and a Fe-containing heme group (616.17728 Da) covalently bound to Cys14 and Cys17 of cytochrome c, custom modifications were included at the N and C termini to identify the modified ECD fragments. These corresponded to 237.9606 Da (1,2-BDSA and 1,3-BDSA), 287.9762 Da (1,5-NDSA), 485.9886 Da (BS2G), 528.0356 Da (BS3) and 616.0153 Da (sulfo-EGS). The triplicate data was filtered to only include fragments observed in at least two out of three runs and manually verified before visualizing the sequence maps in ProSight Lite.³⁵

Molecular Modeling and CCS Calculations. The gas-phase optimized geometries of the anionic crosslinkers were used to determine the $N_{\zeta} - C_{\zeta}$ and $N_{\zeta} - N_{\zeta}$ distance (i.e., the lengths of restraints to use). First, the linkers containing rings were energy minimized in Avogadro³⁶ using default settings and used as is for geometry optimization with quantum mechanical methods due to the rigidity and planarity of the structures. BS2G, BS3, and sulfo-EGS conformers were searched with the Avogadro systematic rotor search. The twenty lowest energy conformers were used as starting structures for quantum mechanical geometry optimization. Geometries were optimized and zero-point corrected energies were calculated with density functional theory using the Becke, 3-parameter, Lee–Yang–Parr functional combinations³⁷⁻³⁸ and the basis set 6-31+G(d) in Gaussian 16.³⁹ Additionally, to account for the electrostatic bond (i.e. 'salt bridge' distance)

between the sulfonate sulfur and either the ammonium N_{ζ} of Lys or the guanidinium C_{ζ} of Arg, the energy-minimized structure of 1,5-NDSA was modeled with butyl ammonium and ethyl guanidinium respectively, resulting in distances of 3.2 Å and 4.2 Å. The sulfonate sulfur atom was used since all 3 oxygens are the same bond order and partial negative charge. Thus for the $N_{\zeta} - N_{\zeta}$ distance restraint, 6.4 Å (two Lys 'salt bridge' distances) was added to the crosslinker length while the $N_{\zeta} - C_{\zeta}$ restraint was calculated by adding the crosslinker length and 7.4 Å (sum of Arg and Lys 'salt bridge' distances, **Figure 1** and **Figure 5**). Gas-phase structures of cytochrome c were simulated using an existing protocol for gas-phase protein molecular dynamics (MD) that has been validated against experimental ion mobility measurements.^{15, 40} Briefly, five replicates of MD run for 100 ns in GROMACS 2019.4⁴¹ using the GROMOS 43a2 force field⁴² were simulated, incorporating distance restraints with a force constant of 100 kJ/mol Å².²⁷ Charged sites were determined from ECD of cytochrome c 7⁺ (*vide infra*) and acidic residues were protonated (neutral) for simulations. CCS_{He} values for MD models were calculated in IMoS software⁴³⁻⁴⁴ using the Trajectory Method (TM) algorithm with a helium drift gas and default Lennard-Jones parameters and protein charge set to 7. Helium was used as the ion/ion chemistry takes place in the helium-filled trap cell.

Results and Discussion

Electrostatic Crosslinking Using Various Reagent Length and Flexibility.

Crosslinking reagents were each reacted with cytochrome c 7⁺ in the gas phase, resulting in an electrostatic adduct formed between the two negatively charged sulfonate groups of the crosslinker and accessible, protonated amino acid residues on cytochrome c. The aromatic disulfonate salts (1,2-BDSA, 1,3-BDSA and 1,5-NDSA) provide a more rigid and shorter range of crosslinking distances while the more linear sulfo-NHS containing compounds (BS2G, BS3 and sulfo-EGS) are able to capture longer distances (**Figure 1**). Listed distances are measured between the two sulfonate sulfurs after gas-phase geometry optimization by averaging the distances for all structures within 2 kcal/mol of the lowest energy structure (zero-point corrected

energies). Of note, only sulfo-EGS was dynamic and adopted multiple structures spanning a range of crosslinking distances within 2 kcal/mol of the lowest energy conformer. The range of distances for BS3 and BS2G were less than an angstrom above or below the listed distance, which is likely within the experimental error of chemical crosslinking.⁴⁵ Since cytochrome c was crosslinked in the gas phase, the reagent structures were modeled in the absence of solvent to allow for charge solvation by the sulfonate groups to minimize charge repulsion while ignoring any solvent-mediated stabilization owing to charge and hydrogen bonds typically present in solution conditions.

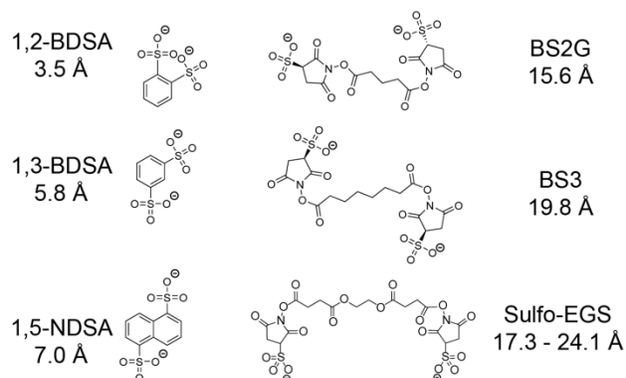


Figure 1. Panel of electrostatic crosslinking reagents with the respective spacer length between the two sulfonate groups calculated from geometry-optimized structures modeled in the gas phase.

After the formation of a stable electrostatic complex in the trap cell, the reaction products and unmodified protein are separated by ion mobility (IM), with a drift time of 14.1 ms observed for the 1x crosslinked adduct formed using 1,3-BDSA (**Figure 2A**). The ion/ion reaction was tuned to favor the formation of the 1x crosslinked adduct while minimizing the addition of supplemental crosslinks, thus there is still some unmodified protein (drift time = 9.68 ms) observed after the reaction in the trap. The multiple (n) crosslinks species were not considered in our analyses since addition of n number of crosslinks effectively reduces the overall charge state by $-2 \times n$, thereby limiting the number of protonated sites available during electron capture dissociation (ECD) and hindering efficient fragmentation. The presence of excessive crosslinks can also potentially lock the protein structure into an irrelevant conformation that is different from the starting structure. The smaller peak (drift time = 11.8 ms) corresponds to [cytochrome c + 7H + \diamond]⁶⁺, representing a

“dead-end” link where the reagent is attached at only one protonated site using one of the two available sulfonate groups and the other sulfonate group has been protonated (i.e., neutralized) by proton transfer and is thus unreactive. Mobility-separated ion species are fragmented by ECD to identify the sites of modification on cytochrome c 7⁺. However, the overall sequence coverage was poor with the transfer CE voltage ‘off’ (at 4 V) (**Figure S1**). To improve sequence coverage and facilitate site annotation, ECD fragments are further subjected to supplemental activation in the transfer cell (transfer CE 30 V), which provides enough energy to disrupt the intramolecular interactions preventing the separation of the c and z fragment ions while minimizing the formation of collision induced dissociation (CID) products, which involve mobilization of protons and the possible loss of the crosslinkers. The energy in the transfer cell was carefully tuned to preserve the electrostatic complex as shown in the extracted mass spectrum of the IM peak (drift time = 14.10 ms), where the [cytochrome c + 7H + ♦]⁵⁺ complex (2519.26 *m/z*) is the major peak relative to the unmodified protein (2471.67 *m/z*). Efficient ECD is also demonstrated with two subsequent electron capture peaks [M + ♦]⁴⁺ and [M + ♦]³⁺ at 3149.07 and 4198.76 *m/z* respectively (**Figure 2B**). The IM and *m/z* plots for the other two aromatic disulfonate compounds are similar to 1,3-BDSA (**Figure S2**). Interestingly, for all three sulfo-NHS based crosslinkers, the additional energy in the transfer cell is sufficient to drive the formation of the covalent product [M + *]⁵⁺, corresponding to the loss of the two sulfo-NHS groups (**Figure S2**). However, since these covalent bonds are formed after ECD, the current implementation of the instrument platform prevents fragmentation of these gas-phase covalent crosslinked proteins. Thus, covalent crosslinking was not included in the analyses. Tuning the reactivity of the crosslinking chemistry and the instrument for more facile covalent crosslinking is currently in progress.

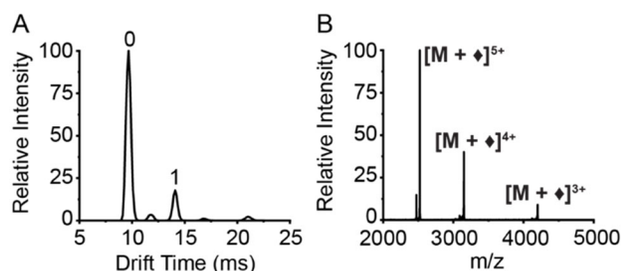


Figure 2. Gas-phase electrostatic reaction between [cytochrome c + 7H]⁷⁺ and [1,3-BDSA]²⁻. (A) Ion mobility spectrum following ion/ion reaction, with numbered labels corresponding to the number of electrostatic attachments. (B) Mass spectrum extracted from '1' electrostatic attachment IM peak (denoted by ♦).

A zoomed-in view of the 3200 – 4000 m/z range shows some annotated c and z fragments shifted by the additional mass of the 1,3-BDSA crosslinker in contrast to unmodified ions resulting from the cytochrome c 5⁺ precursor obtained from two consecutive proton transfer reactions from cytochrome c 7⁺ using PFO (**Figure 3**), which serves as a charge-reduced “control”, having the same net charge as the crosslinked cytochrome c 7⁺ without the modifications present. The unmodified c and z ions generated after the ion/ion reaction with PFO are also observed across the entire protein sequence, thereby demonstrating efficient ECD sequencing of cytochrome c (despite resulting charge prior to ECD equal to +5) (**Figure S3**). Therefore, changes in the fragmentation patterns between the charge reduced control and the crosslinked 7⁺ are likely due to the presence of the linker, aiding in the interpretation of the crosslinking ECD fragments results.

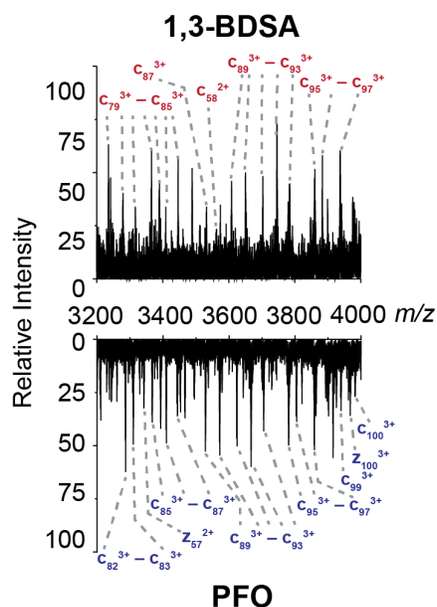


Figure 3. Extracted mass spectra from $[M + \text{♦}]^{5+}$ precursor ion comparing the c and z ions from crosslinking by 1,3-BDSA (top) and two consecutive proton transfer by PFO (bottom), with modified and unmodified fragments denoted in red and blue, respectively.

The protonated sites are first assigned to identify where electrostatic attachment by the sulfonate groups may occur. To do so, the ECD fragmentation data of unmodified cytochrome c 7^+ is analyzed to track where incremental increases in charge occur, as c and z fragments get larger, due to the presence of an ionizing proton. Similar strategies have been previously employed to determine the charge sites of ubiquitin after top-down fragmentation by ECD⁴⁶ and UVPD⁴⁷. Hence, the protonated sites for unmodified cytochrome c 7^+ include Lys13, Lys25, Arg38, Lys55, Lys73, Lys88 and Lys99 (**Figure S4**). Following cytochrome c crosslinking with the various disulfonate-containing compounds, the resulting c and z ions were analyzed and used in conjunction with the charge sites for cytochrome c 7^+ to determine the site(s) of electrostatic attachment. All the ECD experiments were run in triplicates, with the three datasets combined and filtered to only include reproducible ions (observed in $n \geq 2$). Both unmodified and crosslinked c and z ions were then manually validated and mapped onto the protein sequence to identify modified residues. Unmodified fragments (labeled in blue) are annotated without including any mass shift while modified fragments (shown in red) are mapped after including the mass increase due to the reagent (**Figure 4**). Both unmodified and modified fragments are helpful when

determining the position of the crosslinker. The smallest modified c and z ions ($[c + \diamond]$ and $[z + \diamond]$) that are observed, bracket the protein region being crosslinked since those require the added mass of the reagent to be assigned, indicating that a crosslinker must be present N-terminal to the smallest $[c + \diamond]$ ion and C-terminal to the smallest $[z + \diamond]$ ion. The amino acid sequence spanned by the crosslinker is also held together by the reagent, thereby resulting in an apparent silencing of fragmentation between the two crosslinked residues. Furthermore, the largest c and z unmodified fragments indicate that no crosslinking occurs N-terminal to the largest unmodified c ion and C-terminal to the largest unmodified z ion. Combining the location of the ionizing protons, the smallest modified c and z ions, the “silenced” regions, and the location of the largest unmodified c and z ions allows us to assign the location of the crosslinks. Notably, comparing the ECD pattern of the unmodified fragments of the crosslinked complex to that of the 2x proton transfer product using PFO shows that ECD sequence coverage is excellent in the absence of the crosslinker, especially in the Lys25 – Lys88 region where crosslinking is occurring (**Figure 4, Figure S3**). This provides additional confirmation that the gaps in sequencing are due to the presence of the crosslinker.

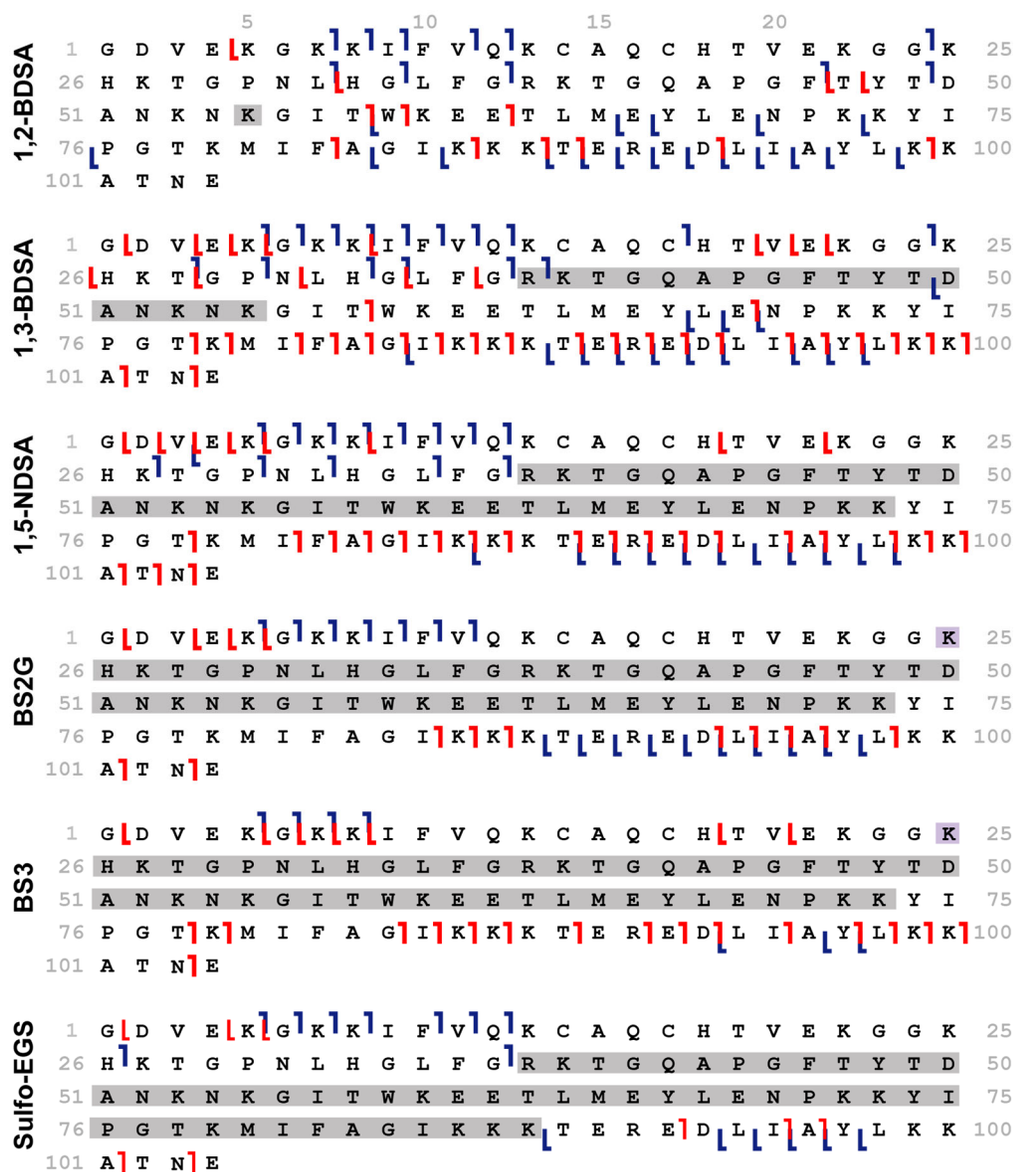


Figure 4. Sequence ladders generated from the ECD fragmentation of the ion/ion crosslinking product of cytochrome c 7⁺. Modified fragments including the added mass of the reagent are labeled in red while unmodified ions are denoted in blue. Crosslinked regions are highlighted in gray.

For the cytochrome c 7⁺ and 1,2-BDSA adduct, the largest unmodified ions are c₄₉ and z₄₆ while the smallest modified fragments are [c₅₈ + ♦] and [z₅₇ + ♦], suggesting that the crosslinker is present in this Tyr48 – Thr58 region. However, the only protonated site for electrostatic attachment is Lys55 (highlighted in gray), implying that 1,2-BDSA is only bound at that one site. This is not surprising considering that the sulfonate groups on 1,2-BDSA are next to each other (spacer length = 3.9 Å) and are likely unable to bridge the distance between Lys55 and its closest

accessible protonated site. The remaining reagents are all able to crosslink a pair of residues, with different spacer lengths providing a useful range of crosslinking distances that can be detected. In the case of 1,3-BDSA (spacer length = 5.8 Å), the reagent crosslinks Arg38 and Lys55 since the largest unmodified c and z ions are c_{38} and z_{55} while the smallest modified fragments are $[c_{58} + \blacklozenge]$ and $[z_{68} + \blacklozenge]$, with almost complete silencing of sequence coverage between those fragments. With a larger spacer length of 7.0 Å, 1,5-NDSA forms a Arg38 – Lys73 crosslink, given the largest unmodified ions are c_{37} and z_{18} while the smallest modified fragments are $[c_{78} + \blacklozenge]$ and $[z_{83} + \blacklozenge]$. The region between Arg38 and Lys73 shows no sequence coverage, but for the charge reduced only (via PFO) version, there is significant fragmentation in this region, providing additional evidence of the crosslink's location. Reaction with BS2G results in c_{11} and z_{16} as the largest unmodified ions and smallest modified ones as $[c_{85} + \blacklozenge]$ and $[z_{99} + \blacklozenge]$. The largest unmodified ions observed for BS3 are c_8 and z_{11} while $[c_{78} + \blacklozenge]$ and $[z_{84} + \blacklozenge]$ are the smallest modified fragments. Despite BS2G and BS3 having different spacer lengths of 15.6 Å and 19.8 Å respectively, they both react at Lys25 and Lys73. Finally, with its largest unmodified ions being c_{37} and z_{16} and smallest modified fragments being $[c_{92} + \blacklozenge]$ and $[z_{99} + \blacklozenge]$, sulfo-EGS binds at Arg38 and Lys88, indicating that these residues are within 17.3 Å and 24.1 Å apart at the ends of the side chains. Using a panel of crosslinkers with distinct size and flexibility allows the capture of multiple residue pairs used to map the structure of cytochrome c 7⁺ and, it provides insights into the proximity and orientation of the protonated sites in the gas phase.

Molecular Dynamics Models Generated Using Experimental Gas-phase Crosslinking Data as Distance Restraints. The various modified sites identified experimentally (Arg38 – Lys55, Arg38 – Lys73, Arg38 – Lys88 and Lys25 – Lys73) restrain distances between crosslinked residues and guide MD simulations of cytochrome c in the gas phase. The spacer length of the crosslinker (sulfonate sulfur to sulfonate sulfur) predicted using the DFT geometry optimized gas-phase structures (**Figure 1**) is added to the 'salt bridge' (i.e. gas-phase electrostatic bond) distance to obtain the Arg C_ζ – Lys N_ζ and Lys N_ζ – Lys N_ζ distances used as restraints for

the crosslinked pairs. Analyses of large chemical crosslinking datasets have shown that using the $N_{\zeta} - N_{\zeta}$ straight line distances accurately recapitulated condensed phase structures, especially when compared to the more traditional $C_{\alpha} - C_{\alpha}$ distance restraint.^{48, 49} **Figure 5** illustrates how the gas-phase electrostatic bond distance was calculated for the interaction from the sulfonate sulfur (S) to the guanidinium carbon (C_{ζ}) of Arg (4.2 Å) and that between the sulfonate sulfur (S) and nitrogen (N_{ζ}) of Lys (3.2 Å). Thus, for the Arg $C_{\zeta} - Lys N_{\zeta}$ restraint, a total noncovalent bond distance of 7.4 Å was added to the reagent spacer length while 6.4 Å was used in the case of Lys $N_{\zeta} - Lys N_{\zeta}$. This approach accounts for the flexibility of the side chains and is an improved approximation for the crosslinked (i.e., restraint) distance, resulting in a distance restraint range of 24.7 – 31.5 Å for sulfo-EGS.

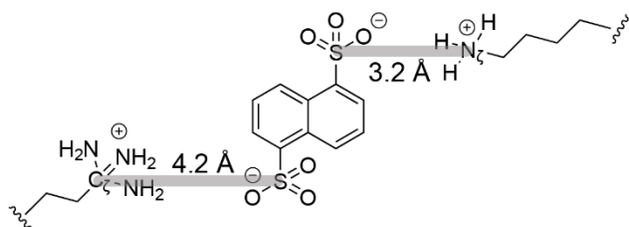


Figure 5. The $S - C_{\zeta}$ and $S - N_{\zeta}$ gas-phase electrostatic bond distances are measured between sulfonate sulfur and either ethylene guanidinium C_{ζ} (4.2 Å) or butylene ammonium N_{ζ} (3.2 Å) respectively (highlighted in gray).

Table 1. Summary of Measured Crosslinking Distances and CCS_{He} Values Calculated for X-ray Structure (PDB ID 1HRC) and MD Models.¹

Model	1,3-BDSA 38C _ζ -55N _ζ (Å)	1,5-NDSA 38C _ζ -73N _ζ (Å)	Sulfo-EGS 38C _ζ -88N _ζ (Å)	BS2G/ BS3 25N _ζ -73N _ζ (Å)	CCS_{He} (Å ²)
1HRC (X-ray)	17.4	26.2	25.6	33.4	1310
1	14.5	21.5	23.0	30.3	1187
2	14.3	16.5	23.5	23.9	1227
3	14.3	23.2	22.9	25.3	1166
4	14.8	19.9	23.1	24.1	1186
5	12.4	21.7	23.4	25.5	1169
Restraint distance (Å)	13.2	14.4	24.7	22.0 - 26.2	

Final structures from five replicate gas-phase MD runs of 100 ns restrained by the experimentally determined crosslinking distances were simulated using the X-ray structure of cytochrome c (PDB ID: 1HRC) as a starting structure. **Table 1** summarizes the measured distances between the crosslinked pairs as well as the collision cross-section values (CCS_{He}) calculated using the Trajectory Method algorithm and default Lennard-Jones parameters in IMoS. Calculations were run in helium since the ion/ion reaction takes place in the trap cell with helium as the drift gas. All of the distances for the X-ray structure are larger than the calculated restraint distance as is the computed CCS_{He} value of 1310 Å² compared to 1247 Å² published previously for native cytochrome c 7⁺ using a drift tube ion mobility instrument with He as the drift gas by Shelimov et al.⁵⁰ **Figure 6A** also shows the X-ray structure of cytochrome c and with the side chains oriented outwards and none of the measured distances below the calculated restraint

¹ Numbers bolded in blue represent experimental distances measured using X-ray and MD structures that are smaller than the calculated restraint distance.

distances. These data correlate with the compaction of solution structure upon transfer to the gas phase, a phenomenon that has been thoroughly studied.⁵¹⁻⁵²

The calculated CCS_{He} for all the MD models are within 6.5 % of the experimental value (1247 Å²) obtained by Shelimov et al. While the CCS_{He} values provide a coarser approach to study protein structure in the gas phase, the experimental crosslinking distances are useful in gaining residue-level structural details. None of the MD models satisfy the restraint distances for all four crosslinking pairs, suggesting a degree of dynamicity of the side chain residues in the gas phase and the presence of multiple co-existing structures that are being captured using gas-phase crosslinking. Of the five restrained structures we modeled, model 5 had distances within calculated crosslinker lengths for three out of four restrained distances (the Arg38 – Lys55 (1,3-BDSA), Arg38 - Lys88 (Sulfo-EGS), and Lys25 – Lys73 (BS2G and BS3)). Although all five of the Arg38 – Lys73 distances were overlength for our calculation of the 1,5-NDSA distance, the Arg38 – Lys73 distance for model 2 is only 2.1 Å larger than the 1,5-NDSA distance, showing that the structure that 1,5-NDSA is crosslinking is likely similar but non-identical to model 2. Models 2, 3, and 4 had two out of four distances within the crosslinker distances. (Arg38 – Lys88 and Lys25 – Lys73), while model 1 had only one distance (Arg38 – Lys88) within the crosslinker distance. This shows that structures similar to model 5 (and perhaps model 2) were more often sampled by our crosslinkers, and structures similar to model 1 were least sampled by our crosslinkers. Structures similar to model 5 may be sampled most often because they are the most populated structure family; however, additional MD runs are needed to confirm whether or not this hypothesis is statistically rigorous.

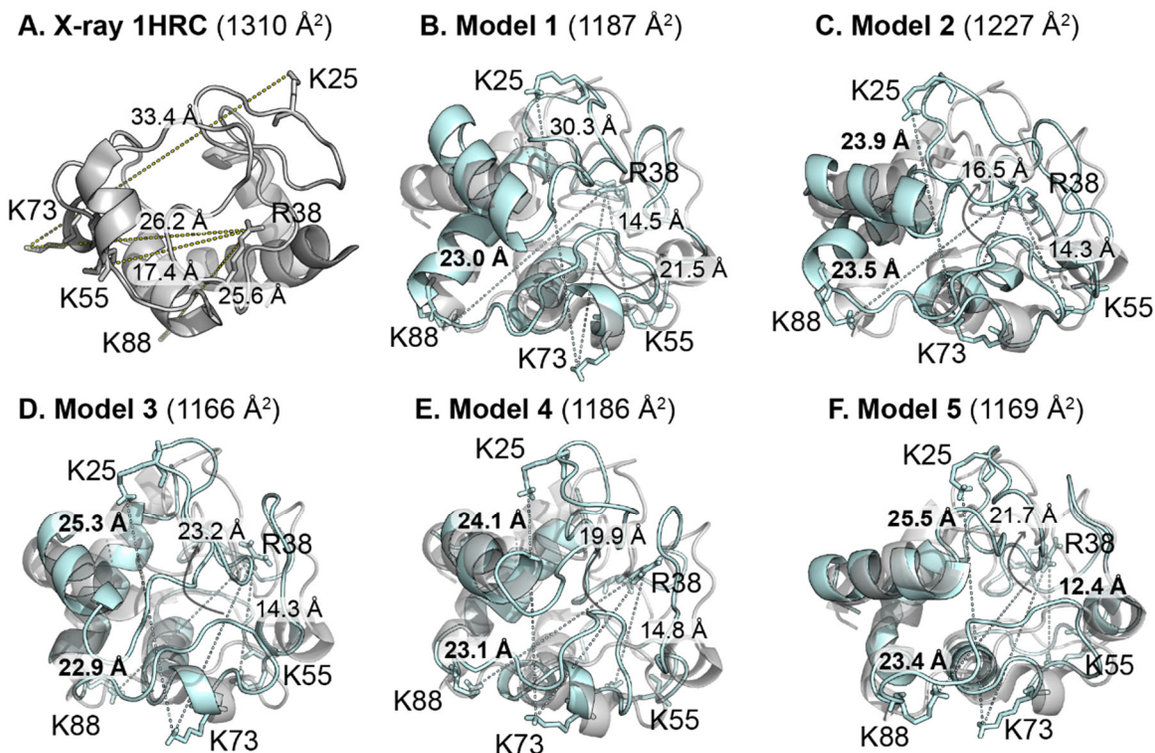


Figure 6. Modified side chains displayed using cytochrome c (A) x-ray structure only and (B – F) MD models 1 – 5 (cyan) aligned to the x-ray structure (gray). Dotted lines represent the C_ζ – N_ζ or N_ζ – N_ζ distances measured for the different crosslinked residue pairs, with measured distances within the calculated restraint distance highlighted in bold.

Alignment of individual gas-phase MD model with the x-ray structure of cytochrome c provides a visual assessment of how dynamic the crosslinked amino acid residues are (**Figure 6**). Overall, the secondary structure features overlap well between the condensed phase (gray) and gas-phase (cyan) structures, such as α -helices spanning regions such as Thr49 – Lys55, Asn69 – Ile75 and Lys87 – Glu108 (C-terminus), albeit with slight shifts in relative positions. The modified residues Lys55, Lys73 and Lys88 are part of the α -helices while Lys25 and Arg38 are in random coil regions. The decreased overlap for the random coil regions demonstrates compaction in the gas-phase as well as the trapping of kinetically frozen solution structures during the electrospray process. Thus, despite marginal changes in the calculated CCS_{He} values, our restrained MD models show that cytochrome c undergoes rearrangements at the residue level, which are uniquely probed by gas-phase crosslinking. Moreover, the use of a multipass traveling wave cyclic IM (cIM) device to achieve higher resolution separation has led to the detection of new IM features within a broad IM peak distribution for cytochrome c 7⁺, likely due to various,

highly similar conformers present within that broadened IM peak.⁵³ Thus, our gas-phase crosslinking study using a Synapt G2-Si instrument provides evidence for multiple co-existing conformers in the gas-phase when combined with MD calculations, recapitulating the published results obtained for cytochrome c 7⁺ after multiple passes in a cIM. However, we note that a fuller exploration of the solution phase conformational landscape through longer simulations or replica exchange methods⁵⁴⁻⁵⁵ is necessary for gas-phase *de novo* structural determination.

Finally, we recognize that our models are dependent on our choice of a starting structure, which is the crystal structure of cytochrome c. If we performed a full gas-phase conformational search, our results may have been different, showing evidence for non-native structures. However, rigorous modeling of the energetics of ions in a Synapt G2-Si, the same instrument that we used for this study, showed that protein ion temperatures without applying collision induced unfolding were on the order of much less than 400 K (e.g., for native-like myoglobin 8⁺ ions, temperatures were near 330 K),⁵⁶ which would not support an unfolding and refolding mechanism, also consistent with MD of the transition of folded proteins from the droplet to the gas phase.⁵⁷ Since the gas phase lacks solvent, and thus refolding back into the native structure can be considered highly unlikely,¹¹ approaches such as gas phase simulated annealing MD performed at much higher temperatures might be expected to give artifactual results for well-folded proteins. Indeed, increasing the temperature (i.e., through collisional activation) of compact cytochrome c structures in the gas phase does produce extended states that compact down to the original CCS values, with the compact structures displaying a “permanent” increase in CCS.⁵⁸⁻⁵⁹ With these considerations, the most accurate modeling of gaseous structures of well-folded proteins would need to keep temperatures low enough that the gas phase rearrangement barriers are not reached. Additionally, using the crystal structure of folded proteins, followed by relaxation in the gas phase, accurately reproduces experimental observables.^{15, 40}

Conclusions

We have demonstrated, for the first time, the use of an ensemble of different length gas-phase noncovalent crosslinkers via ion/ion reactions to provide insight into the nature of gas-phase structures of a cooperatively folded protein. Crosslinking was performed prior to ion mobility, allowing for ion/ion reactions products to be separated and identifications made based on ECD with mild supplemental activation. We only used c and z ions to determine noncovalent linking identifications to prevent misidentifications due to mobilization of the sulfonate-lysine bonds, though previous work has shown this bond has covalent-like strength in the gas phase.⁶⁰ The most abundant native-like ion from cytochrome c, the 7⁺ charge state, yielded different identifications of crosslinks with increasing linker length, evidence that the crosslinks are sensitive to 3-D structure. When used to restrain gas-phase molecular dynamics simulations, the crosslink identifications predicted structures that gave calculated CCS_{He} in reasonable agreement with experimental values, giving evidence that cytochrome c native-like ions remain compact in the gas phase. This work, along with mobility measurements and infrared spectroscopy²⁰, clearly supports the kinetic trapping of solution-like structures during gentle transfer to the gas phase.

Interestingly, and importantly, the combination of ion/ion reaction-derived distance restraints and molecular dynamics did not converge on a single structure that satisfied all criteria. This correlates with observations based on tandem IM measurements from the Clemmer Laboratory, where they observed that the arrival time distribution of ubiquitin ions was composed of many non-interconverting structures,⁶¹ which, again, was recapitulated with cyclic tandem ion mobility experiments with cytochrome c.⁶² Instead, we predict structures that are similar in CCS, but are not identical. Our structures are not exhaustive; there are several individual conformations that we did not determine that are still supported by our measurements. Thus, in the future, a more thorough conformational search is warranted. However, our modeled structures do show preservation of the overall protein fold, where the “breathing” motion of less ordered regions and compaction of side chains to form new hydrogen bonds/salt bridges account for the differences

in structure. Therefore, in the future, we envision the application of this combined experimental and MD method to follow changes in solution structure due to unfolding conditions, ligand and metal binding, etc., and determine their effects on gas-phase structure and whether non-native structures are retained in the gas phase.

Supporting Information

ECD sequence coverage of crosslinked cytochrome c without supplemental activation, IM and m/z spectra of the rest of the crosslinking ion/ion reactions, ECD sequence coverage of charge reduced cytochrome c via ion/ion reaction with PFO, ECD sequence coverage of unmodified cytochrome c 7^+ showing the assignment of ionizing protons via increase in fragment ion charge state.

Acknowledgements

Portions of this work were funded by the National Institutes of Health under R21GM134408 and the Indiana University-Purdue University Indianapolis School of Science. We would like to acknowledge Dr. Amber Rolland and Professor James Prell from the University of Oregon for assistance with the implementation of gas phase molecular dynamics.

References

1. Hamdy, O. M.; Julian, R. R., Reflections on charge state distributions, protein structure, and the mystical mechanism of electrospray ionization. *J Am Soc Mass Spectrom.* **2012**, *23*, 1-6.
2. Loo, J. A., Studying noncovalent protein complexes by electrospray ionization mass spectrometry. *Mass Spectrometry Reviews.* **1997**, *16*, 1-23.
3. McLafferty, F. W., Tandem mass spectrometry. *Science.* **1981**, *214*, 280-287.
4. Leney, A. C.; Heck, A. J. R., Native Mass Spectrometry: What is in the Name? *J. Am. Soc. Mass Spectrom.* **2017**, *28*, 5-13.
5. Butler, K. E.; Takinami, Y.; Rainczuk, A.; Baker, E. S.; Roberts, B. R., Utilizing Ion Mobility-Mass Spectrometry to Investigate the Unfolding Pathway of Cu/Zn Superoxide Dismutase. *Front Chem.* **2021**, *9*, 614595.
6. Ruotolo, B. T.; Benesch, J. L.; Sandercock, A. M.; Hyung, S. J.; Robinson, C. V., Ion mobility-mass spectrometry analysis of large protein complexes. *Nat Protoc.* **2008**, *3*, 1139-1152.
7. Ruotolo, B. T.; Robinson, C. V., Aspects of native proteins are retained in vacuum. *Curr Opin Chem Biol.* **2006**, *10*, 402-8.
8. Zhong, Y.; Han, L.; Ruotolo, B. T., Collisional and Coulombic unfolding of gas-phase proteins: high correlation to their domain structures in solution. *Angew Chem Int Ed Engl.* **2014**, *53*, 9209-9212.
9. Wagner, N. D.; Russell, D. H., Defining Noncovalent Ubiquitin Homodimer Interfacial Interactions through Comparisons with Covalently Linked Diubiquitin. *J Am Chem Soc.* **2016**, *138*, 16588-16591.
10. Wyttenbach, T.; Bowers, M. T., Structural stability from solution to the gas phase: native solution structure of ubiquitin survives analysis in a solvent-free ion mobility-mass spectrometry environment. *J. Phys. Chem. B.* **2011**, *115*, 12266-12275.
11. Clemmer, D. E.; Russell, D. H.; Williams, E. R., Characterizing the Conformationome: Toward a Structural Understanding of the Proteome. *Acc Chem Res.* **2017**, *50*, 556-560.
12. El-Baba, T. J.; Woodall, D. W.; Raab, S. A.; Fuller, D. R.; Laganowsky, A.; Russell, D. H.; Clemmer, D. E., Melting Proteins: Evidence for Multiple Stable Structures upon Thermal Denaturation of Native Ubiquitin from Ion Mobility Spectrometry-Mass Spectrometry Measurements. *J Am Chem Soc.* **2017**, *139*, 6306-6309.
13. Dill, K. A.; MacCallum, J. L., The protein-folding problem, 50 years on. *Science.* **2012**, *338*, 1042-6.
14. Mesleh, M. F.; Hunter, J. M.; Shvartsburg, A. A.; Schatz, G. C.; Jarrold, M. F., Structural information from ion mobility measurements: Effects of the long-range potential. *Journal of Physical Chemistry.* **1996**, *100*, 16082-16086.
15. Rolland, A. D.; Prell, J. S., Computational Insights into Compaction of Gas-Phase Protein and Protein Complex Ions in Native Ion Mobility-Mass Spectrometry. *Trends Analyt Chem.* **2019**, *116*, 282-291.
16. Warnke, S.; von Helden, G.; Pagel, K., Protein structure in the gas phase: the influence of side-chain microsolvation. *J Am Chem Soc.* **2013**, *135*, 1177-80.
17. Goth, M.; Lermyte, F.; Schmitt, X. J.; Warnke, S.; von Helden, G.; Sobott, F.; Pagel, K., Gas-phase microsolvation of ubiquitin: investigation of crown ether complexation sites using ion mobility-mass spectrometry. *Analyst.* **2016**, *141*, 5502-10.

18. Konermann, L.; Silva, E. A.; Sogbein, O. F., Electrochemically induced pH changes resulting in protein unfolding in the ion source of an electrospray mass spectrometer. *Anal. Chem.* **2001**, *73*, 4836-44.
19. Soleilhac, A.; Dagany, X.; Dugourd, P.; Girod, M.; Antoine, R., Correlating droplet size with temperature changes in electrospray source by optical methods. *Anal. Chem.* **2015**, *87*, 8210-7.
20. Seo, J.; Hoffmann, W.; Warnke, S.; Bowers, M. T.; Pagel, K.; von Helden, G., Retention of Native Protein Structures in the Absence of Solvent: A Coupled Ion Mobility and Spectroscopic Study. *Angew Chem Int Ed Engl.* **2016**, *55*, 14173-14176.
21. Bakhtiari, M.; Konermann, L., Protein Ions Generated by Native Electrospray Ionization: Comparison of Gas Phase, Solution, and Crystal Structures. *J. Phys. Chem. B.* **2019**, *123*, 1784-1796.
22. Bleiholder, C.; Liu, F. C., Structure Relaxation Approximation (SRA) for Elucidation of Protein Structures from Ion Mobility Measurements. *J. Phys. Chem. B.* **2019**, *123*, 2756-2769.
23. Cavalli, A.; Salvatella, X.; Dobson, C. M.; Vendruscolo, M., Protein structure determination from NMR chemical shifts. *Proceedings of the National Academy of Sciences.* **2007**, *104*, 9615-9620.
24. Uphoff, S.; Holden, S. J.; Le Reste, L.; Periz, J.; van de Linde, S.; Heilemann, M.; Kapanidis, A. N., Monitoring multiple distances within a single molecule using switchable FRET. *Nature Methods.* **2010**, *7*, 831-836.
25. Brodie, N. I.; Popov, K. I.; Petrotchenko, E. V.; Dokholyan, N. V.; Borchers, C. H., Solving protein structures using short-distance cross-linking constraints as a guide for discrete molecular dynamics simulations. *Science Advances.* **2017**, *3*, e1700479.
26. Khaje, N. A.; Eletsy, A.; Biehn, S. E.; Mobley, C. K.; Rogals, M. J.; Kim, Y.; Mishra, S. K.; Doerksen, R. J.; Lindert, S.; Prestegard, J. H.; Sharp, J. S., Validated determination of NRG1 Ig-like domain structure by mass spectrometry coupled with computational modeling. *Communications Biology.* **2022**, *5*, 452.
27. Ly, T.; Julian, R. R., Elucidating the tertiary structure of protein ions in vacuo with site specific photoinitiated radical reactions. *J Am Chem Soc.* **2010**, *132*, 8602-9.
28. Cheung See Kit, M.; Carvalho, V. V.; Vilseck, J. Z.; Webb, I. K., Gas-Phase Ion/Ion Chemistry for Structurally Sensitive Probes of Gaseous Protein Ion Structure: Electrostatic and Electrostatic to Covalent Cross-Linking. *Int J Mass Spectrom.* **2021**, *463*, 116549-116559.
29. Webb, I. K.; Mentinova, M.; McGee, W. M.; McLuckey, S. A., Gas-phase intramolecular protein crosslinking via ion/ion reactions: ubiquitin and a homobifunctional sulfo-NHS ester. *J Am Soc Mass Spectrom.* **2013**, *24*, 733-43.
30. Webb, I. K.; Morrison, L. J.; Brown, J., Dueling electrospray implemented on a traveling-wave ion mobility/time-of-flight mass spectrometer: Towards a gas-phase workbench for structural biology. *Int. J. Mass spectrom.* **2019**, *444*, 116177.
31. Carvalho, V. V.; Cheung See Kit, M.; Webb, I., An Ion Mobility and Gas-Phase Covalent Labeling Study of the Structure and Reactivity of Gaseous Ubiquitin Ions Electro sprayed from Aqueous and Denaturing Solutions. *Journal of the American Society for Mass Spectrometry.* **2020**.
32. Williams, J. P.; Morrison, L. J.; Brown, J. M.; Beckman, J. S.; Voinov, V. G.; Lermyte, F., Top-Down Characterization of Denatured Proteins and Native Protein Complexes

Using Electron Capture Dissociation Implemented within a Modified Ion Mobility-Mass Spectrometer. *Anal. Chem.* **2020**, *92*, 3674-3681.

33. Park, J.; Piehowski, P. D.; Wilkins, C.; Zhou, M.; Mendoza, J.; Fujimoto, G. M.; Gibbons, B. C.; Shaw, J. B.; Shen, Y.; Shukla, A. K.; Moore, R. J.; Liu, T.; Petyuk, V. A.; Tolić, N.; Paša-Tolić, L.; Smith, R. D.; Payne, S. H.; Kim, S., Informed-Proteomics: open-source software package for top-down proteomics. *Nature Methods.* **2017**, *14*, 909-914.

34. Donnelly, D. P.; Rawlins, C. M.; DeHart, C. J.; Fornelli, L.; Schachner, L. F.; Lin, Z.; Lippens, J. L.; Aluri, K. C.; Sarin, R.; Chen, B.; Lantz, C.; Jung, W.; Johnson, K. R.; Koller, A.; Wolff, J. J.; Campuzano, I. D. G.; Auclair, J. R.; Ivanov, A. R.; Whitelegge, J. P.; Paša-Tolić, L.; Chamot-Rooke, J.; Danis, P. O.; Smith, L. M.; Tsybin, Y. O.; Loo, J. A.; Ge, Y.; Kelleher, N. L.; Agar, J. N., Best practices and benchmarks for intact protein analysis for top-down mass spectrometry. *Nature Methods.* **2019**, *16*, 587-594.

35. Fellers, R. T.; Greer, J. B.; Early, B. P.; Yu, X.; LeDuc, R. D.; Kelleher, N. L.; Thomas, P. M., ProSight Lite: graphical software to analyze top-down mass spectrometry data. *Proteomics.* **2015**, *15*, 1235-8.

36. Hanwell, M. D.; Curtis, D. E.; Lonie, D. C.; Vandermeersch, T.; Zurek, E.; Hutchison, G. R., Avogadro: an advanced semantic chemical editor, visualization, and analysis platform. *Journal of Cheminformatics.* **2012**, *4*, 17.

37. Lee, C.; Yang, W.; Parr, R. G., Development of the Colle-Salvetti correlation-energy formula into a functional of the electron density. *Physical review B.* **1988**, *37*, 785.

38. Becke, A., Density-functional thermochemistry. III. The role of exact exchange (1993) *J. Chem. Phys.* **98**, 5648.

39. Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Petersson, G. A.; Nakatsuji, H.; Li, X.; Caricato, M.; Marenich, A. V.; Bloino, J.; Janesko, B. G.; Gomperts, R.; Mennucci, B.; Hratchian, H. P.; Ortiz, J. V.; Izmaylov, A. F.; Sonnenberg, J. L.; Williams; Ding, F.; Lipparini, F.; Egidi, F.; Goings, J.; Peng, B.; Petrone, A.; Henderson, T.; Ranasinghe, D.; Zakrzewski, V. G.; Gao, J.; Rega, N.; Zheng, G.; Liang, W.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; Throssell, K.; Montgomery Jr., J. A.; Peralta, J. E.; Ogliaro, F.; Bearpark, M. J.; Heyd, J. J.; Brothers, E. N.; Kudin, K. N.; Staroverov, V. N.; Keith, T. A.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A. P.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Millam, J. M.; Klene, M.; Adamo, C.; Cammi, R.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Farkas, O.; Foresman, J. B.; Fox, D. J. *Gaussian 16 Rev. C.01*, Wallingford, CT, 2016.

40. Rolland, A. D.; Biberic, L. S.; Prell, J. S., Investigation of Charge-State-Dependent Compaction of Protein Ions with Native Ion Mobility–Mass Spectrometry and Theory. *J. Am. Soc. Mass Spectrom.* **2022**, *33*, 369-381.

41. Abraham, M. J.; Murtola, T.; Schulz, R.; Páll, S.; Smith, J. C.; Hess, B.; Lindahl, E., GROMACS: High performance molecular simulations through multi-level parallelism from laptops to supercomputers. *SoftwareX.* **2015**, *1-2*, 19-25.

42. Schuler, L. D.; Van Gunsteren, W. F., On the choice of dihedral angle potential energy functions for n-alkanes. *Molecular Simulation.* **2000**, *25*, 301-319.

43. Larriba, C.; Hogan Jr, C. J., Free molecular collision cross section calculation methods for nanoparticles and complex ions with energy accommodation. *Journal of Computational Physics.* **2013**, *251*, 344-363.

44. Larriba, C.; Hogan Jr, C. J., Ion mobilities in diatomic gases: measurement versus prediction with non-specular scattering models. *The Journal of Physical Chemistry A*. **2013**, *117*, 3887-3901.
45. Merkle, E. D.; Rysavy, S.; Kahraman, A.; Hafen, R. P.; Daggett, V.; Adkins, J. N., Distance restraints from crosslinking mass spectrometry: mining a molecular dynamics simulation database to evaluate lysine-lysine distances. *Protein Sci*. **2014**, *23*, 747-59.
46. Kit, M. C. S.; Carvalho, V. V.; Vilseck, J. Z.; Webb, I. K., Gas-phase ion/ion chemistry for structurally sensitive probes of gaseous protein ion structure: Electrostatic and electrostatic to covalent cross-linking. *International Journal of Mass Spectrometry*. **2021**, *463*, 116549.
47. Morrison, L. J.; Brodbelt, J. S., Charge site assignment in native proteins by ultraviolet photodissociation (UVPD) mass spectrometry. *Analyst*. **2016**, *141*, 166-176.
48. Gong, Z.; Ye, S.-X.; Tang, C., Tightening the Crosslinking Distance Restraints for Better Resolution of Protein Structure and Dynamics. *Structure*. **2020**, *28*, 1160-1167.e3.
49. Degiacomi, M. T.; Schmidt, C.; Baldwin, A. J.; Benesch, J. L. P., Accommodating Protein Dynamics in the Modeling of Chemical Crosslinks. *Structure*. **2017**, *25*, 1751-1757.e5.
50. Shelimov, K. B.; Jarrold, M. F., Conformations, unfolding, and refolding of apomyoglobin in vacuum: An activation barrier for gas-phase protein folding. *J. Am. Chem. Soc.* **1997**, *119*, 2987-2994.
51. Hall, Z.; Politis, A.; Bush, M. F.; Smith, L. J.; Robinson, C. V., Charge-state dependent compaction and dissociation of protein complexes: insights from ion mobility and molecular dynamics. *J Am Chem Soc*. **2012**, *134*, 3429-38.
52. Pacholarz, K. J.; Porrini, M.; Garlish, R. A.; Burnley, R. J.; Taylor, R. J.; Henry, A. J.; Barran, P. E., Dynamics of Intact Immunoglobulin G Explored by Drift-Tube Ion-Mobility Mass Spectrometry and Molecular Modeling. *Angewandte Chemie International Edition*. **2014**, *53*, 7765-7769.
53. Eldrid, C.; Ujma, J.; Kalfas, S.; Tomczyk, N.; Giles, K.; Morris, M.; Thalassinos, K., Gas Phase Stability of Protein Ions in a Cyclic Ion Mobility Spectrometry Traveling Wave Device. *Analytical Chemistry*. **2019**, *91*, 7554-7561.
54. Sugita, Y.; Okamoto, Y., Replica-exchange molecular dynamics method for protein folding. *Chemical physics letters*. **1999**, *314*, 141-151.
55. Rhee, Y. M.; Pande, V. S., Multiplexed-Replica Exchange Molecular Dynamics Method for Protein Folding Simulation. *Biophysical Journal*. **2003**, *84*, 775-786.
56. Donor, M. T.; Shepherd, S. O.; Prell, J. S., Rapid Determination of Activation Energies for Gas-Phase Protein Unfolding and Dissociation in a Q-IM-ToF Mass Spectrometer. *J Am Soc Mass Spectrom*. **2020**, *31*, 602-610.
57. McAllister, R. G.; Metwally, H.; Sun, Y.; Konermann, L., Release of Native-like Gaseous Proteins from Electrospray Droplets via the Charged Residue Mechanism: Insights from Molecular Dynamics Simulations. *J Am Chem Soc*. **2015**, *137*, 12667-76.
58. Eldrid, C.; Ben-Younis, A.; Ujma, J.; Britt, H.; Cragolini, T.; Kalfas, S.; Cooper-Shepherd, D.; Tomczyk, N.; Giles, K.; Morris, M.; Akter, R.; Raleigh, D.; Thalassinos, K., Cyclic Ion Mobility–Collision Activation Experiments Elucidate Protein Behavior in the Gas Phase. *Journal of the American Society for Mass Spectrometry*. **2021**, *32*, 1545-1552.

59. Borotto, N. B.; Osho, K. E.; Richards, T. K.; Graham, K. A., Collision-Induced Unfolding of Native-like Protein Ions Within a Trapped Ion Mobility Spectrometry Device. *Journal of the American Society for Mass Spectrometry*. **2022**, *33*, 83-89.
60. Cheung See Kit, M.; Shepherd, S. O.; Prell, J. S.; Webb, I. K., Experimental Determination of Activation Energies for Covalent Bond Formation via Ion/Ion Reactions and Competing Processes. *J Am Soc Mass Spectrom*. **2021**, *32*, 2313-2321.
61. Koeniger, S. L.; Merenbloom, S. I.; Clemmer, D. E., Evidence for many resolvable structures within conformation types of electrosprayed ubiquitin ions. *J. Phys. Chem. B*. **2006**, *110*, 7017-21.
62. Eldrid, C.; Ujma, J.; Kalfas, S.; Tomczyk, N.; Giles, K.; Morris, M.; Thalassinos, K., Gas Phase Stability of Protein Ions in a Cyclic Ion Mobility Spectrometry Traveling Wave Device. *Anal. Chem*. **2019**, *91*, 7554-7561.