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sites in peptides using soft X-ray action spectroscopy†

Mapping the electronic transitions of protonation

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Near-edge X-ray absorption mass spectrometry (NEXAMS) around the nitrogen and oxygen K-edges was employed on gas-phase peptides to probe the electronic transitions related to their protonation sites, namely at basic side chains, the N-terminus and the amide oxygen. The experimental results are supported by replica exchange molecular dynamics and density-functional theory and restricted open-shell configuration with single calculations to attribute the transitions responsible for the experimentally observed resonances. We studied five tailor-made glycine-based pentapeptides, where we identified the signature of the protonation site of N-terminal proline, histidine, lysine and arginine, at 406 eV, corresponding to N 1s $\rightarrow \sigma^*(NH_x^+)$ (x = 2 or 3) transitions, depending on the peptides. We compared the spectra of pentaglycine and triglycine to evaluate the sensitivity of NEXAMS to protomers. Separate resonances have been identified to distinguish two protomers in triglycine, the protonation site at the Nterminus at 406 eV and the protonation site at the amide oxygen characterized by a transition at 403.1 eV.

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Introduction

In the biological medium, the charge states of molecules such as peptides, proteins, and nucleic acids play a significant role as they influence conformation and reactivity. Within the physiological pH range, an important mechanism altering the charge state is the protonation and deprotonation of specific chemical groups. To improve our understanding of biological processes and the structure-function relationship, we need to localize the (de)protonation sites in biomolecules. Experimentally, protonation sites in peptides have been studied using different techniques over the last few decades mainly by gas-phase basicity studies¹⁻⁴ in order to evaluate proton affinities of chemical groups. According to the proton affinities of the different sites, protonation in amino acids and small peptides occurs preferably at the nitrogen site of the most basic side chains (arginine, histidine, and lysine with proton affinities of 250.2, 231.8, and 237.1 kcal mol^{-1} , respectively⁵) and at the N-terminus primary amine -NH₂. However, in certain cases, for example, when the number of protons exceeds the number of basic residues, protonation can also occur on the peptide backbone. Action spectroscopy techniques, and in particular infrared (IR) spectroscopy, are powerful tools for the determination of molecular structures in the gas phase that can be used to assess the protonation site of molecules as big as small peptides.6,7

Protonated triglycine has been studied theoretically and experimentally by many groups mostly to understand its structure and sites of protonation that remains controversial. In her theoretical study on glycine oligomers, Chung-Phillips concluded that the most stable conformer of triglycine is protonated at an amide

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oxygen in the backbone.8 Wu and McMahon employed infrared multiphoton dissociation (IRMPD) to determine the structure and protonation site of triglycine. They showed that, under their experimental conditions, two distinct protomers of triglycine with different protonation sites were responsible for the recorded spectrum, one protonated at the N-terminus of the tripeptide and the other protonated on the first amine oxygen. The same conclusion was drawn by Grégoire and co-workers in a study in which they combined resonant IRMPD of protonated triglycine and Car-Parrinello molecular dynamics calculations.7 However, in a recent study, Li and co-workers found that the protonation site of triglycine is the N-terminal nitrogen rather than the amine oxygen, which they found to be at least 0.14 kcal mol⁻¹ more stable. ¹⁰ The authors also calculated the X-ray absorption spectra at the carbon, nitrogen, and oxygen K-edges of triglycine protomers with different conformations. They observed clear differences between the linear and folded conformers of triglycine, with resonances unique to some conformers and with the strongest effects being observed at the nitrogen and oxygen edges. This theoretically demonstrated that near-edge X-ray absorption fine structure (NEXAFS) spectroscopy can be sensitive to the different protonation sites of a peptide. Furthermore, this method has already been applied to determine the position of hydrogen atoms and the protonation state of biomolecules in the liquid phase and in crystals. 11,12

Over the last few decades, soft X-ray spectroscopy has become an important technique for studying the electronic structure of biomolecular systems, both in the condensed13,14 and gas phases. 15-17 In fact, soft X-ray excitation is element specific, so tuning the photon energy to a resonance allows site-selective excitation of a molecule to specific unoccupied molecular orbitals. Very recently, Wang and co-workers demonstrated using near-edge X-ray absorption mass spectrometry (NEXAMS) that the electronic structure of gas-phase oligonucleotides is influenced by the protonation site. 18 In a previous study, Otero and Urquhart observed that peptides in the solid phase show differences in their N 1s NEXAFS spectra depending on the protonation state of the amine group. 19 For amine-protonated species, they observed a feature at a high energy of around 406 eV, well above the resonance around 402 eV characterizing an unprotonated amine group.

To go further and map the different electronic transitions responsible for the protonation site in peptides, we studied tailormade glycine-based pentapeptides. Glycine is the simplest amino acid with a single hydrogen as its side chain and has been studied extensively.^{2,20-26} Therefore, the study of peptides made from glycines and one basic amino acid at the C-terminus, or a proline at the N-terminus, allowed us to avoid probing effects from side chains' atoms that are not at the actual protonation site, while keeping the system simple for comparison with calculations. To probe the electronic structures of the different protonation sites in peptides, we studied five different glycinebased protonated pentapeptides by soft X-ray action spectroscopy. As protonation occurs preferentially at nitrogen sites, we present here a study at the nitrogen K-edge of model peptides with known protonation sites: [GGGGG + H]⁺, [PGGGG + H]⁺, $[GGGGH + H]^+$, $[GGGGR + H]^+$, and $[GGGGK + H]^+$ (noted G_5 , PG_4 , G₄H, G₄R, and G₄K, respectively, in the following), where G

$$H_3N^{\downarrow}$$
 NH_2 NH_2 NH_3 NH_3 NH_3 NH_3 NH_2 NH_2 NH_2 NH_3 NH_2 $NH_$

Fig. 1 Chemical structure of protonated pentaglycine (top) and the chemical structures of the last amino acid of G₄R, G₄H, G₄K, and the first amino acid of the sequence PG₄, from left to right (bottom).

stands for glycine, P for proline, H for histidine, R for arginine, and K for lysine, as well as protonated triglycine [GGG + H]⁺, denoted G₃ in the following. The chemical structures of G₅ and of the amino acids that make up the other pentapeptide sequences are shown in Fig. 1. G₅ and PG₄ are both N-terminal-protonated but with primary and secondary amines, respectively. Histidine, arginine, and lysine are basic residues, and the protonation occurs on their side chains. The study of triglycine allows us to test the theory of Li and co-workers. 10

Methods

All custom peptides were purchased from ProteoGenix (Schiltigheim, France), at >95% purity and used without further purification. The sample solutions were prepared at 30 µM concentration in 1:1 water/methanol with 1% of formic acid to ensure the protonation of the peptides.

The experiments have been performed during two experimental campaigns, one at the BESSY II synchrotron (HZB, Berlin, Germany) and one at the PETRA III synchrotron (Deutsches Elektronen-Synchrotron DESY, Germany) on two different apparatus with a similar principle. The results for protonated G₄H, G₄R, and PG₄ have been obtained by coupling our home-built tandem mass spectrometer to the P04 soft X-ray beamline of PETRA III. This setup relies on a high-fluence electrospray ionization (ESI) source to introduce the ions into the gas phase. The electrosprayed ions are transported through a heated capillary into the first pumping stage which hosts an ion funnel that collects and focuses the ions. They continue through a radio frequency (RF) octupole ion guide to a quadrupole mass filter where the ions of interest are selected according to their mass-to-charge (m/z) ratio. Then, a second octupole and a set of Einzel lenses guide and focus the ions into an RF 3D Paul ion trap where they are accumulated prior to irradiation with photons. To allow efficient trapping of the ions, their kinetic energy is cooled down to room temperature by collisions with helium buffer gas. All cationic products are eventually extracted from the trap and analyzed in a reflectron time-of-flight (Re-TOF) mass spectrometer ($m/\Delta m = 1800$). The duration of the synchrotron light exposure, typically 2 seconds, is controlled with an

optomechanical shutter. At the N K-edge, the energy bandwidth was set to 250 meV. The photon energy scans were performed by steps of 200 meV across the N K-edge (397-412 eV). A photodiode placed downstream of the ion trap was used to record the photon flux. The NEXAMS spectra obtained during this beamtime were normalized to photon flux and precursor ion intensity to remove the contribution of the ESI source fluctuations. To account for all sources of background ions, the data acquisition was divided into repeated cycles of three mass scans. First, a mass spectrum of the ESI only was recorded to measure the precursor ion intensity and fragments originating from energetic collisions with the buffer gas. Second, the mass spectrum resulting from irradiation of the trapped molecules was recorded. For the third spectrum, the ESI is switched off and the mass spectrum resulting from the irradiation of the residual gas was recorded. The mass spectrum resulting from the ESI and the one resulting from the irradiation of the residual gas were subtracted from the mass spectrum resulting from the irradiation of the trapped molecules. To obtain the final mass spectrum, a series of a hundred cycles was accumulated to average over long-term fluctuations in the source. Therefore, in the final subtracted mass spectrum, the precursor ion peak of each molecule appears negative due to this procedure.

The results on protonated G₃, G₅, and G₄K have been obtained at the UE52_PGM Ion Trap beamline of the BESSY II synchrotron, as described elsewhere. ^{27,28} Briefly, the instrument is equipped, similar to the setup described above, with an ESI source, a quadrupole mass filter, a cryogenically cooled linear ion trap and a Re-TOF mass spectrometer. For G4K, the temperature of the trap was set at 290 K. Contrary to side-chain protonation, backbone protonation is more sensitive to a peptide's conformation. In an attempt to reduce the number of conformers in the trap and limit them to the lowest energy conformers, the temperature of the trap was set to 8 K for G₃ and G₅. The N K-edge spectra were recorded by steps of 100 meV and 210 meV bandwidth, across the range 397-410 eV (only to 408 eV for G4K). Measurements at the O K-edge for G5 and G3 were additionally performed to assess the effect of a potential amide oxygen protonation. Here, the photon-energy bandwidth was 200 meV and the photon energy was ramped in 100 meV steps between 528 and 545 eV. A photodiode located behind the interaction region was used to record the photon flux for data normalization. Note that the design of the ion trap instrument limits the detection of the irradiation products to a small m/zwindow. The extraction parameters were set to observe fragments over a mass range of about m/z 50–150 for G_4K and over a mass range of m/z 25–70 for the measurements of G_5 and G_3 . The NEXAMS spectra, or total ion yield (TIY) spectra, can be assumed to be good approximations of the X-ray absorption cross section, and thus comparable to the calculated DFT/ROCIS oscillator strengths. Indeed, the partial ion yields for individual fragments are the results of the convolution of the absorption cross section of the molecule with the fragments' branching ratio. By summing up, the contribution of all fragments into a "total ion yield" spectrum, we approach the result of an absorption spectrum.²⁹ Nevertheless, disagreement in spectral intensity could arise in the case of detecting multiple ions from the same photoabsorption event or due to discrepancies in the overall detection efficiency of ions. NEXAMS, or TIY NEXAFS, has been extensively used and compared with X-ray absorption calculations. 17,18,30-35

Theoretical calculations

Structural candidates for G₃ and the different pentapeptides were obtained with different structure search techniques. For G₃, a random structure search was performed, where all dihedral angles of the molecule were allowed to rotate with random angles. The search was conducted using the GenSec package. 36,37 We sampled 600 structures starting from the three different protonation states labelled GGGH1, GGGH2 and GGGH3 (see Fig. 5). These structures were fully optimized with the PBE038 exchange-correlation functional including the pairwise TS-vdW corrections³⁹ with the FHIaims program package.40 The final optimized structures were ranked by their total energy, and we selected conformers lying on the lowest 1 eV range for each protonation state. In total, we obtained 10 GGGH1, 10 GGGH2, and 5 GGGH3 conformers. This search strategy was targeted at finding low-temperature, enthalpically stabilized conformers. We subsequently calculated in silico X-ray absorption spectra using the density functional theory (DFT) combined with the restricted-open-shell configuration interaction with singles excitations (ROCIS) for the lowest energy conformer of each protonation state. 41 The calculated X-ray absorption spectra of GGGH3 do not reproduce the experiment at all, the results for this conformer will not be discussed further in this paper. Apart from the fact that only a few conformations were found in the ab initio based GenSec structure search performed in this work, there is evidence in the literature that only a few conformers contribute to the spectroscopic signals of small glycine peptides, as shown in the study by Wu and Mc Mahon.9 In addition, the extensive conformer search performed by Hudgins and co-workers found essentially only one stable conformation in the case of glycine pentapeptides and smaller glycines.42

A different strategy was adopted for the pentapeptides, targeted at finding higher temperature stable conformers. We conducted molecular dynamics simulations and DFT/ROCIS calculations for the pentapeptides using the GROMACS program (version 2018.8) and the ORCA electronic structure package, 43 respectively. To efficiently sample the potential energy surface (PES), exponentially distributed replica exchange molecular dynamics (REMD) simulations with different temperatures ranging from 300 K to 648 K were performed based on the Amber ff14SB force field44 in GROMACS. The conformers generated by the REMD simulation were represented by the values of their dihedral angles for the backbone and side chains. Calculating these descriptors for each snapshot of the REMD simulation resulted in a feature dataset of the REMD trajectory. These features were used to cluster conformers with similar structures using the probabilistic analysis of molecular motifs (PAMM)^{45,46} such that we could identify recurring molecular conformations of different metastable states sampled during REMD. PAMM is specifically designed to work in relatively high-dimensional spaces and can partition the underlying probability distribution function into modes corresponding to different conformers. The lowest REMD

energy conformation of each identified cluster was chosen as the representative conformer of that cluster and considered for the calculation of the absorption spectra, as discussed by Kotobi and co-workers. 47 A Boltzmann weighting factor based on the REMD energy differences between the representative conformers and normalized by the cluster population was employed to estimate the statistical relevance of the various metastable states and to obtain weighted average spectra considering a canonical ensemble of the conformers. Conformers, from REMD, having a weighting factor less than 2% of the contribution were assumed to be negligible. All theoretical absorption spectra presented in this paper were obtained using the DFT/ROCIS approach, employing the TZVP Ahlrichs basis set⁴⁸ in combination with the B3LYP exchange-correlation functional.⁴⁹ The optimized conformers were used to calculate transition energies and dipole moments for carbon, nitrogen and oxygen K-edges. We calculated the average absorption spectra by weighting the contribution of different conformers, using the previously calculated weighting factors. The final absorption spectra were then broadened, using a pseudo-Voigt profile with a full width at half-maximum (FWHM) of 0.6 eV.⁵⁰ The shift in the energy of the theoretical spectra is done by matching the first resonance of the theoretical spectra to the corresponding one in the experimental spectra, representing π^*_{CONH} electronic transitions in the peptide backbone, as explained later. 51,52 This shift is due to the B3LYP functional underestimating the core excitation energies. 53,54 Such a method has already been used successfully to compare DFT calculations with soft X-ray absorption spectra. 15,17,23,55

Results and discussion

Photodissociation mass spectra analysis

Fig. 2(a)-(f) show the mass spectra measured for peptides G₃ (m/z 190), $G_5 (m/z 304)$, $PG_4 (m/z 344)$, $G_4H (m/z 384)$, $G_4R (m/z 403)$, and G_4K (m/z 375) at the photon energy of 401.6 eV (π *(CONH) resonance, as discussed later). The photodissociation mass spectra over the full mass range for the six peptides are shown in the ESI† (Fig. S1–S6). As can be seen in Fig. S1–S6 (ESI†), for all peptides, no intact photoionized precursor ions have been observed. Fragments are labelled according to the peptide fragment nomenclature established by Roepstorff, Fohlman, and Biemann. 56,57 A table summarizing the different fragments is shown in ESI.†

Triglycine and pentaglycine are composed of the same residues, which reflects in their similar mass spectra (Fig. 2(a) and (b)) with fragments stemming from the backbone. m/z 28 corresponds to a fragment that can be attributed to C₂H₄⁺ or CH₂N⁺, m/z 29 is attributed to the CHO⁺ fragment. The immonium ion of glycine is observed at m/z 30 and at m/z 31 is the fragment CH_3O^+ . m/z 42 corresponds to $C_2H_4N^+$, m/z 43 to CHNO⁺, m/z 45 to $COOH^{+}$, and m/z 56 to $C_2NOH_2^{+}$. These fragments have also been reported for neutral glycine under VUV ionization at 21.21 eV by Lago and co-workers⁵⁸ and by Majer and co-workers using electron-impact ionization.⁵⁹

In the photodissociation mass spectrum of PG_4 (Fig. 2(c)), fewer fragments are observed. The most prominent one is the

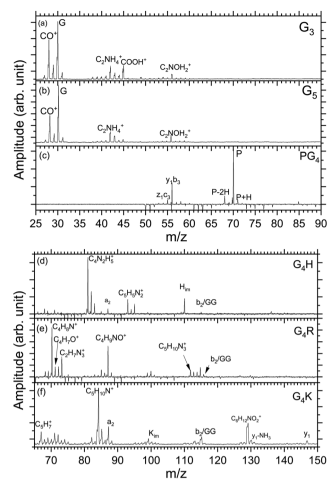


Fig. 2 Low mass region of the photodissociation mass spectra of all investigated protonated peptides measured at 401.6 eV photon energy. The mass range changes between the two sets of mass spectra to show the main fragments produced by photodissociation. Peaks of negative intensity are due to the background subtraction

fragment at m/z 70, which is the immonium ion of the proline amino-acid residue, that is also produced by collision-induced dissociation (CID) as observed by Zhang and co-workers. 60 The other two fragments at m/z 68 and m/z 71 are related to proline as well. The two fragments at m/z 56 and m/z 55 are attributed to internal backbone fragments, C2NOH2+ and C2NOH+, respectively.

In the spectrum of G_4H , shown in Fig. 2(d), most of the ions observed result from fragmentation taking place at the side chain of histidine. The spectrum shows at m/z 110 the immonium ion of histidine (C5H8N3+), as observed in the CID experiment, 61 accompanied by an ammonia loss (-NH3) at m/ z 93 and an additional loss of HCN at m/z 83. The fragments at m/z 82 and 81 result from further hydrogen losses, the latter being the dominant fragment. The peak at m/z 87 is attributed to the a_2 fragment. The fragment at m/z 115 can be attributed to the internal backbone fragment GG or the b₂ fragment which have the same elemental composition.

For the mass spectrum of G_4R , presented in Fig. 2(e), some fragments remain unassigned and the others are mainly due to

fragmentation at the protonated arginine side chain. We do not observe the immonium ion of arginine at m/z 129 but we do observe the related ions at m/z 112, 87, 70. Other fragments at m/z 71, 72, 73, 88 and 85 are attributed to the arginine residue, and the fragments' elemental compositions are given in Table S1 (ESI†). Only the fragments at m/z 70, 71, 72, 88, and 112 have been reported in CID experiments. ⁴⁵ Note that the intense peak at m/z 87 most likely also has a contribution from

the a_2 ion. GG/b_2 fragments are also observed at m/z 115.

In the spectrum of G_4K (Fig. 2(f)), we were not able to clearly identify either the fragments around m/z 70 or those around m/zz 100. These fragments are not background and were therefore included in the analysis. We observed the y_1 fragment at m/z147 and an $-NH_2$ loss from the y₁ fragment at m/z 131. m/z 67 and m/z 84 attributed to $C_5H_7^+$ and $C_5H_{10}N^+$ ions from the lysine side chain have been also observed in the CID.⁶⁰ Other lysine-related fragments are observed at m/z 129, m/z 112, and m/z 101, the latter being the immonium ion of lysine. m/z 87 is the a_2 fragment from the peptide. The fragment at m/z 115 can be attributed to the internal backbone fragment GG or the b₂ fragment for G₄H and G₄R. As the amino-acid sequences of G₄K and G₄H are the same for the first four residues, it is expected to observe a2 and b2 for all peptides under study except PG4. The internal fragment GG should also be a common fragment of all peptides. The fact that it is not observed in PG4 (see Fig. S2, ESI†) suggests the fragment m/z is most likely due to a b₂ fragment rather than GG in G₄H, G₄R and G₄K.

To summarize, in the mass spectra of G_4K , G_4H , and G_4R , the majority of the ions are attributed to fragments from the side chain of the basic residues in the peptides, lysine, histidine, and arginine, respectively. PG_4 , G_5 , and G_3 do not contain a basic side chain, thus the fragments observed arise only from backbone dissociation. Most of the observed fragments were also reported in collision-induced dissociation (CID) experiments 60 suggesting that, in our case, internal conversion followed by intramolecular vibrational redistribution of internal energy is the main process following photoexcitation/photoionization and Auger decay, eventually leading to statistical dissociation of the peptides.

Different protonation sites on amino groups in peptides

The total ion yields (TIY) of each molecule were calculated by summing the areas under the peaks of all detected fragments in the photo-induced mass spectra, for each photon energy over the scanned energy range. No specific fragmentation channels could be identified over the scanned photon energies for the different studied peptides, as shown in the different partial ion yield spectra in the ESI† (Fig. S7–S12).

[GGGGG + H]^{\dagger}. Fig. 3(a) presents the experimental TIY and calculated absorption spectra of G_5 at the N K-edge. The final DFT/ROCIS spectrum is the convolution of the two most populated conformers according to the REMD simulations that have been performed, with a weighting factor of 0.52 for the first conformer (conformer 1), shown in Fig. S13 (ESI \dagger), and a weighting factor of 0.47 for the second conformer (conformer 2), shown in Fig. S15 (ESI \dagger). Conformers 1 and 2 are very similar, conformer 1 being more compact, with the protonated

N-terminus making hydrogen bonds with the oxygen atoms of the backbone and of the carboxylic group at the C-terminus of the peptides. Our calculated structures and the DFT/ROCIS spectrum are in agreement with those obtained by Li and coworkers. Fig. S14 and S16 (ESI†) show the respective calculated DFT spectra of conformers 1 and 2 with the different molecular orbitals responsible for the main transitions.

The first resonance A at 401.6 eV is attributed to N 1s $\rightarrow \pi^*(\text{CONH})$ transitions within peptide bonds according to the DFT/ROCIS transition lines. This has already been reported for peptides in previous papers. This has already been reported for peptides in previous papers. The origin of resonance B at 402.6 eV is controversial. For gas-phase diglycine, Feyer and coworkers attributed this feature to an N 1s $\rightarrow \pi^*(\text{N-C}\alpha)$ transition, as well as Dörner and co-workers for protonated methionine-enkephalin. Zubavichus and co-workers proposed that this feature is an N 1s $\rightarrow \pi^*(\text{CONH})$ transition, that is energetically shifted because of potentially different conformations existing in their studied homopolypeptides. Using time-dependent density functional theory (TDDFT) to calculate NEXAFS spectra of α -helix and β -turn models based on glycines, Buckley and Besley attributed the first two narrow resonances at the N K-edge to N 1s $\rightarrow \pi^*(\text{CONH})$,

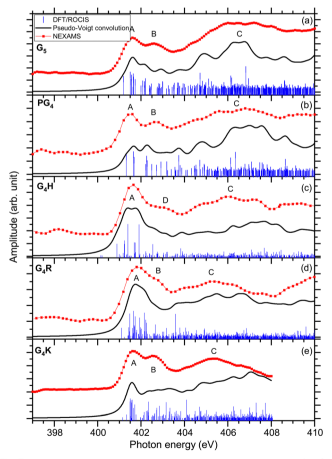


Fig. 3 The experimental nitrogen K-edge total ion yield spectra for $G_5, PG_4,\, G_4H,\, G_4R,\, \text{and}\,\, G_4K,\, \text{are}$ shown with red dotted lines. In blue, the DFT/ROCIS X-ray absorption lines are displayed for each molecule. The absorption lines were shifted by $\sim\!13$ eV and were broadened using a Pseudo-Voigt profile with a 0.6 eV FWHM represented in black.

with the second band associated with π^* orbitals in a neighbouring residue.⁵⁰ Their computed spectra also suggest that this second feature is highly susceptible to the conformation of peptide, being more pronounced in an α -helix than in a β -turn. In the energy range of resonance B, the analysis of the lowest unoccupied molecular orbitals (LUMOs) of the conformers of G₅ (LUMO+1 of conformer 1 and LUMO+3 of conformer 2) leads to transitions from the N 1s level in one peptide bond to unoccupied $\pi^*(CONH)$ orbitals in neighbouring peptide bonds. We finally attribute, according to our calculation and in agreement with the work of Buckley and Besley, 50 resonance B to N_i 1s $\rightarrow \pi^*(CON_iH)$ where N_i and N_i are different nitrogen atoms in the peptide backbone.

The DFT/ROCIS calculations show two major transitions in the energy region of 404-408 eV, one resonance at 404.8 eV and one at 406.6 eV, that are contributing in the experimental spectrum to one broad spectral feature, noted C. The study of the molecular orbitals of the two main conformers of G₅ (LUMO+1 and LUMO+2 of conformer 1, LUMO of conformer 2) assigned the first resonance to an N 1s $\rightarrow \pi^*(CO)$ transition in the carboxylic group of the C-terminus, where the DFT/ROCIS calculations show that the transition is due to the excitation from the nitrogen atoms of the N-terminus to the carboxylic group. Such a transition occurs thanks to the folded conformation of G₅ and the spatial proximity of the N and C-termini. The peak at 406.6 eV is assigned to transitions located at the protonated N-terminus, N 1s \rightarrow $\sigma^*(NH_3^+)$ with the density of molecular orbitals at the N-terminus (LUMO+5 of conformer 1 and conformer 2). Similar transitions at 406.8 and 406.5 eV have been previously reported by Otero and Urquhart in solid-state NEXAFS spectra of zwitterionic glycine and glycine hydrochloride, respectively, and assigned to both N 1s \rightarrow $\sigma^*(NH)$ and N 1s $\rightarrow \sigma^*(CN)$.

[PGGGG + H]⁺. The NEXAMS spectrum of protonated PG₄, presented in Fig. 3(b), is similar to the spectrum of protonated G₅. The DFT-calculated structure of the main conformer (weighting factor of 0.97), shown in Fig. S17 (ESI†), is folded with the N-terminal nitrogen forming hydrogen bonds to the oxygen atoms of the C-terminus and the backbone. The proline group stands out from the folded system. Fig. S18 (ESI†) shows the respective calculated DFT spectrum of PG4 with the different molecular orbitals responsible for the main transitions.

The first resonance A at 401.6 eV is attributed to N 1s \rightarrow $\pi^*(CONH)$ transitions in a peptide bond according to the DFT/ ROCIS transition lines (LUMO). In the energy range of resonance B, the DFT/ROCIS calculations show a resonance at 402.1 eV and one at 403.7 eV, involving the molecular orbitals LUMO+1 and LUMO+3, that can both be attributed to electronic transitions from one peptide bond to a neighboring peptide bond. However, experimentally, only a resonance at 402.6 eV has been observed. We therefore attribute resonance B to N_i 1s $\rightarrow \pi^*(CON_iH)$ transitions to neighboring peptide bonds, as discussed for G5 and consistent with our observations for the different protonated pentapeptides in this study.

Feature C is composed of different electronic transitions which explain the broadness of the peak. The DFT/ROCIS result gives a resonance at 404.7 eV mainly due to an N 1s $\rightarrow \sigma^*(NH)$ transition from the nitrogen atoms in the backbone, whereas at

406.5 V, the calculated molecular orbital (LUMO+7) shows most of the density around the proline residue and describes an N 1s $\rightarrow \sigma^*(CH)$ electronic transition due to the excitation of the protonated secondary amine nitrogen. Note that, for the immonium ion (m/z 70), shown in Fig. S8 (ESI†), the intensity of feature C is lower than for the other fragments contributing to the total ion yield spectrum.

 $[GGGGH + H]^{+}$. The experimental NEXAMS spectrum of G_4H , shown in Fig. 3(c), also presents three main features. For G₄H, only the conformer with a weighting factor of 0.953 was studied, the other calculated conformers have a weighting factor below 0.01 and thus were not considered in the final analysis. The DFToptimized geometry of G₄H is shown in Fig. S19 (ESI†). The structure is mostly folded around the backbone of the peptide. The protonated nitrogen of the histidine side chain forms hydrogen bonds with the oxygen atoms of the backbone. In contrast to G5, the N-terminus does not form any hydrogen bonds with the backbone which leads to G₄H being less compact than G₅. Fig. S20s (ESI[†]) shows the respective calculated DFT spectra of G₄H with the different molecular orbitals responsible for the main transitions.

The first resonance A is located at 401.6 eV and can be partly attributed to N 1s $\rightarrow \pi^*(CONH)$ transitions in the peptide according to the analysis of the DFT/ROCIS transition lines at 401.2 eV. Additionally, the main resonance peak energetically overlaps with N 1s $\rightarrow \pi^*(C=N)$ electronic transitions in the imidazole ring of the histidine side chain. The calculations give the transition lines at 400.8 eV, 401.4 eV, and 401.9 eV due to the double bond C=N of the histidine imidazole ring (LUMO) and the transition lines at 401.2 eV, involving LUMO+1 and LUMO+2, stemming from the N 1s $\rightarrow \pi^*(CONH)$ peptide bond transitions. These additional histidine-related transitions contribute to the increased intensity and broadening of resonance A in comparison with the other peptides. In contrast to the other studied pentapeptides, neither experimentally nor theoretically, does the absorption spectrum of protonated G4H show a resonance B at 402.6 eV, attributed to N_i 1s $\rightarrow \pi^*(CON_iH)$ transition in neighboring peptide bonds. Resonance B at 402.6 eV is only distinguishable in the partial ion yield spectrum of the immonium ion (m/z 110) (Fig. S9, ESI†) is. This could be attributed to the fact that the absorption in the histidine side chain in resonance A does not lead to the production of the immonium ion (similar to what was discussed for the proline immonium ion in PG₄); thus, resonance A is relatively smaller in this fragment and resonance B can be observed. It is worth noting here that the PIY of G₄H is dominated by the ion yield of fragment m/z 81 that involves H_3 CN loss from the histidine side chain. This fragment shows a very high yield at resonance A, which most probably correlates with the N 1s $\rightarrow \pi^*(C=N)$ electronic transitions within the histidine side chain.

Resonance D at 403.1 eV is found, experimentally, at significantly higher energy than resonance B at 402.6 eV for the other protonated pentapeptides. The DFT/ROCIS calculation gives a resonance at 402.5 eV due to N 1s \rightarrow $\sigma^*(CH)$ and N 1s \rightarrow $\sigma^*(NH)$ electronic transitions (LUMO+6) in the histidine side chain where the protonation site is located. For G₄H, the

resonance at 403.1 eV is the signature of the protonation site which explains the low intensity of broad feature C compared to the intensity of feature C in G₅. The broad spectral feature located in C between 404 and 408 eV remains unclear as it is due to different electronic transitions from the backbone of the protonated peptides. The DFT/ROCIS transitions are all of low intensity and cannot be attributed to one resonance because the molecular orbitals are highly delocalized. This feature has a contribution from LUMO+13 and LUMO+19 for the transition lines at 407.1 eV and from LUMO+34 for the transition lines around 407.9 eV. The analysis of the LUMO+13 and LUMO+19 orbitals shows that feature C includes contributions from σ^* transitions from the backbone and the side chain.

[GGGGR + H]⁺. In Fig. 3(d), the experimental NEXAMS spectrum of G₄R at the nitrogen K-edge is compared with the result of the DFT/ROCIS simulations. The final DFT/ROCIS spectrum is the convolution of the two most populated conformers according to the REMD simulations that have been performed, with a weighting factor of 0.63 for the first conformer (conformer 1, shown in Fig. S21, ESI†) and a weighting factor of 0.28 for the second conformer (conformer 2, shown in Fig. S23, ESI†). The calculated structures are folded around both the side chain and the C-terminus. The N-terminus of both conformers does not form hydrogen bonds with either the C-terminus or the backbone of the side chain. Instead, the guanidine group, where the proton is located, forms a hydrogen bond network with the carboxylic group and backbone. For conformer 1, the nitrogen atom with the protonation site forms only one hydrogen bond with an oxygen atom of the backbone, whereas the one from conformer 2 forms a hydrogen bond with a nitrogen atom of the first glycine as well as one with an oxygen atom. Conformer 2 establishes more hydrogen bonds than conformer 1, which leads to a more folded structure. Fig. S22 and S24 (ESI†) show the respective calculated DFT spectra of conformers 1 and 2 with the different molecular orbitals responsible for the main transitions.

In the experimental spectrum, the first two resonances are not well resolved compared to the other studied peptides. The first resonance A at 401.6 eV has been attributed to the peptide bond resonance N 1s $\rightarrow \pi^*(CONH)$ arising from transitions to the first LUMOs of both conformers (LUMO to LUMO+6 for conformer 1 and from LUMO to LUMO+2 for conformer 2).

According to our DFT/ROCIS calculations, two resonances are responsible for the second feature B at 402.6 eV: N_i 1s \rightarrow $\pi^*(CON_iH)$ (LUMO of conformer 1) to a neighbouring peptide bond and N 1s $\rightarrow \pi^*(C=N)$ (LUMO+4 of conformer 2) in the arginine side chain. In the case of zwitterionic arginine, Zubavichus and co-workers observed a π^* resonance located around 401.4 eV that they attributed to the guanidine group of arginine, whereas Leinweber and co-workers observed the π^* resonance of arginine at 401.7 eV. 13,14 Moreover, Stewart-Ornstein and co-workers attributed the resonance at 402.9 eV of a Sub6 solid sample, an arginine-containing antimicrobial peptide, to a $\pi^*(C=N)$ transition in the arginine residue.⁶⁴ In the case of protonated G₄R, our calculations show that this resonance is shifted higher in energy to lie between the two

peptide bond resonances at 402 eV. The overlap with this second transition explains why the resonance here is not clearly separated by one eV from the first peptide bond resonance, as some transition lines from the DFT/ROCIS calculations are related to the $\pi^*(C=N)$ resonance. Interestingly, resonance B is more apparent in the PIY of the fragment m/z 73 (C₂H₇N₃⁺) that is formed from the intact guanidine group of the arginine side chain. It can therefore be understood that N 1s \rightarrow $\pi^*(C=N)$ in the arginine would likely result in a C=N bond cleavage explaining why these transitions are not observed in the PIY of m/z 73 (Fig. S10, ESI†) and thus better resolving the separation of resonances A and B. The DFT/ROCIS calculations show a transition at 403.6 eV that can possibly be responsible for the small feature before the broad one between 404 and 408 eV. This transition is attributed to N 1s $\rightarrow \sigma^*(NH_2^+)$ transitions from the arginine side chain due to the additional proton for conformer 1 (LUMO+9). For conformer 2, it is more ambiguous as there are two main transition lines at 403.96 eV and 404.1 eV that are responsible for this small feature. The transition at 403.96 eV is due to the LUMO+2 and is attributed to N 1s $\rightarrow \pi^*(CO)$ transitions, whereas at 404.1 eV, the molecular orbital (LUMO+13) shows most of the density around the arginine side chain, with some of the density in the backbone as well, and describes an N 1s $\rightarrow \sigma^*(NH)$ transition due to an electronic excitation in the protonated side chain. Therefore, we concluded that the fingerprint of the protonation site is the first resonance of feature C, at 404 eV. We observed a broad feature, noted C between 404 eV and 408 eV which is less intense than feature C of the other protonated peptides in this study relative to feature A. As the spectra are normalized on the intensity of peak A, this effect could be explained by a higher intensity of peak A in the case of G_4R because of the additional $\pi^*(C=N)$ transition from the arginine side chain that can be found between the resonances A and B. The DFT/ROCIS calculations show that different electronic excitations are responsible for feature C with very low amplitude, which arises mainly from highly delocalized molecular orbitals. Comparing with the NEXAMS spectra of the other pentapeptides of this study, we conclude that the feature is mainly due to σ^* transitions from the backbone.

[GGGGK + H]⁺. The REMD calculations of the conformers of G₄K revealed two main structures responsible for the experimental spectrum, which we will hereinafter refer to as conformer 1 and conformer 2, with a weighting factor of 0.45 each. The structures, shown in Fig. S25 and S27 (ESI†), are mostly folded, with the protonated amino group of the lysine side chain forming hydrogen bonds with oxygen atoms along the peptide backbone and at the C-terminus. Fig. S26 and S28 (ESI†) show the respective calculated DFT spectra of conformers 1 and 2 with the different molecular orbitals responsible for the main transitions.

The experimental spectrum of G_4K presented in Fig. 3(e) shows the same three main features as the other peptides studied. The first peak A is the main peptide bond resonance N 1s $\rightarrow \pi^*(CONH)$ at 401.6 eV (LUMO+1 of conformer 1 and LUMO+2 of conformer 2). This is consistent with our results obtained for the other peptides. As for protonated G₄R, the second feature at 402.6 eV is due to different contributions. The

DFT/ROCIS calculations of conformer 1 show that the peak at 402.4 eV only originates from N_i 1s $\rightarrow \pi^*(CON_iH)$ transitions to neighbouring peptide bonds (LUMO+5). On the other hand, the results of the DFT/ROCIS simulation for conformer 2 show that the resonance at 402.6 eV has major contributions from two different electronic transitions. The first one at 402.3 eV is attributed to N 1s $\rightarrow \pi^*(COOH)$ (LUMO+5) in the C-terminus but with the excitation coming from the nitrogen of the fourth glycine rather than the nitrogen of the C-terminus connected to the lysine residue. The second electronic transition is N 1s \rightarrow $\sigma^*(NH)$ in the protonated lysine side chain at 403.1 eV (LUMO+3). The third feature is due to N 1s $\rightarrow \sigma^*$ (CH) electronic transitions from the backbone (LUMO+8 of conformer 1) around 404.5 eV, and N 1s $\rightarrow \sigma^*(NH)$ from the backbone and the protonation site (LUMO+3 of conformer 1, LUMO+4 and LUMO+3, of conformer 2) at 407 eV. Relative to the yield of the N 1s $\rightarrow \pi^*(CONH)$ resonance, the signature of the protonation site in G₄K is less intense than for G₅.

The study of these five protonated model pentapeptides at the N K-edge revealed some features that are common to any protonated peptide. The main results of the study are summarized in Table 1. As was reported previously, all peptides show a strong resonance related to electronic transitions in the peptide bonds, always located at 401.6 eV. 13,23,26 In the NEXAMS spectra, all peptides show a resonance at 402.6 eV that we attribute to transitions from a nitrogen 1s level in one peptide bond to π*(CONH) antibonding orbitals in a neighbouring peptide bond, except for protonated G₄H, which has a feature centred at 403.1 eV due to N 1s $\rightarrow \sigma^*(CH)$ transitions in the histidine side chain where the protonation site is located.

For protonated G_4R , the $\pi^*(C=N)$ transition from the arginine side chain contributes to the resonance at 401.6 eV and the one at 402.6 eV, which results in the resonance appearing broader than for the other peptides. Similarly, G₄H shows $\pi^*(C=N)$ transitions in the histidine's imidazole ring which contributes to the resonance at 401.6 eV. The NEXAMS spectrum of every peptide shows a broad feature between 404 and 408 eV with different intensities relative to the N 1s $\rightarrow \pi^*(CONH)$ resonance. The comparison of the experimental spectra with DFT/ROCIS calculations shows that the feature comes from σ^* resonances from the backbone of the peptide with a major contribution from $\sigma^*(NH)$ resonances located at the protonated nitrogen site of these peptides, both from the basic side chain and the N-terminus. In PG₄ and G₄H, we observed that the immonium ion exhibits a lower intensity of feature C, which can be explained by the resonant excitation in the peptide bond leading to more immonium ion production compared to the absorption of the photon in the side chain, around 406 eV. Interestingly, we did not observe any resonance from the unprotonated N-terminus in the experimental total ion yield spectra in the cases of G₄H, G₄R, and G₄K. Furthermore, for peptides with a protonated side chain, we observed that resonance A is more intense than resonance C as compared to the peptides with a protonation site on the N-terminus. The additional transitions in the side chains energetically close to the resonances A and B lead to resonance A being more intense than for G₅ or PG₄. We could

Table 1 Experimental energy position (exp) and proposed assignment of the spectral feature observed in the total nitrogen K-edge NEXAMS spectra of the different peptides as well as the energies of the DFT/ROCIS absorption lines (calc)

Land Control			
Peak	Exp (eV)	Calc (eV)	Assignment
$\overline{G_5}$			
A	401.6	401.6	N 1s $\rightarrow \pi^*(CONH)$
В	402.6	402.1	$N_i 1s \rightarrow \pi^*(CON_iH)$
С	404-408	404.8	N 1s $\rightarrow \pi^*(CO)$
		406.6	$N 1s \rightarrow \sigma^*(NH_3^+)$
PG_4			
A	401.6	401.6	$N 1s \rightarrow \pi^*(CONH)$
В	402.6	402.1	$N_i 1s \rightarrow \pi^*(CON_iH)$
		403.7	, ,
С	404-408	404.7	N 1s $\rightarrow \sigma^*(NH)$
		406.5	N 1s $\rightarrow \sigma^*(CH)$
G_4R			
A	401.6	401.7	N 1s $\rightarrow \pi^*(CONH)$
В	402.6	402.6	$N_i 1s \rightarrow \pi^*(CON_jH)$
			$N 1s \rightarrow \pi^*(C=N)$
C	404-408	403.6	$N 1s \rightarrow \sigma^*(NH_2^+)$
C	404 400	403.96	N 1s $\rightarrow \pi^*(CO)$
		404.1	$N 1s \rightarrow \sigma^*(NH)$
G_4H			
A	401.6	400.8	N 1s $\rightarrow \pi^*(CONH)$
А	401.0	401.4	$N 1s \rightarrow \pi^*(C = N)$
		401.4	N 15 → h (C—N)
D	403.1	402.5	$N 1s \rightarrow \sigma^*(NH)$
С	404-408	407.1	N 1s $\rightarrow \sigma^*$
		407.9	
G_4K			
A	401.6	401.6	N 1s $\rightarrow \pi^*(CONH)$
В	402.6	402.4	$N_i 1s \rightarrow \pi^*(CON_iH)$
		402.3	N 1s $\rightarrow \pi^*(COOH)$
		403.1	$N 1s \rightarrow \sigma^*(NH)$
С	404-408		$N 1s \rightarrow \sigma^*(NH)$
			N 1s $\rightarrow \sigma^*(CH)$

observe differences between G₄H, G₄R, and G₄K NEXAMS spectra in the shape and the intensity of the resonances A and B. The total ion yield spectrum of G4H does not exhibit any resonance at 402.6 eV but one at 403.1 eV, resonance B appears as a shoulder on resonance A for G₄R, while for G₄K resonance B is almost as intense as resonance A. We discussed that these transitions are better resolved, or absent, in the PIY of certain fragments, especially when comparing e.g. immonium ion and fragments involving C=N bond break. These differences allow distinguishing them from each other. As shown in the previous section, the high intensity of resonance C is due to the protonation site located on the N-terminus. For neutral diglycine in the gas phase, the feature observed at 406 eV is weaker and shows about the same intensity as the resonance at 401.6 eV, 22 which agrees with our conclusion.

Protonation at the amide oxygen: the case of triglycine

Rodriquez and co-workers investigated the proton migration in protonated triglycine using density functional theory. 65 In this study, they found that protonation of the N-terminal nitrogen atom is higher in free energy than the different isomers

resulting from protonation at the carbonyl oxygen of the N-terminal residue. However, in a recent study, Li and co-workers10 calculated that the protonation site of triglycine is the N-terminal nitrogen rather than the amine oxygen, that they found to be at least 0.61 kcal mol⁻¹ more stable than the other protonation sites possible for triglycine. They suggested that experimental studies using soft X-ray spectroscopy could provide a way to distinguish the various G₃ isomers, since NEXAFS spectra should show unique peaks at nitrogen and oxygen K-edges for the different protomers.

Using protonated pentaglycine, we were able to establish the N K-edge fingerprint of an N-terminal protonated peptide. Thus, comparing the NEXAMS spectra of G₅ and G₃ should show whether only conformers of G₃ with the protonation site at the N-terminal nitrogen are present under our experimental conditions. As protonation of an amine oxygen is expected to influence the NEXAMS spectra at the oxygen K-edge, we present in the following results for both, the N and O K-edges. The carbon K-edge spectra of G₅ and G₃ have also been recorded, but as they exhibit only little differences, with a main resonance at 288.3 eV for both protonated peptides which come from C 1s $\rightarrow \pi^*(CONH)$ transitions from the backbone, they are only shown in the ESI† (Fig. S33 to S35) and will not be discussed further in this manuscript. The partial ion yield spectra at the oxygen K-edge are shown in the ESI† (Fig. S36 and S37).

Comparison of the experimental spectra of G₅ and G₃

In this section, we first compare the experimental NEXAMS spectra of protonated pentaglycine and triglycine. We performed DFT/ROCIS calculations on different isomers and possible protonation sites of triglycine to interpret the results of the experimental spectra.

As shown in the previous section and in previous theoretical and experimental studies, 3,9 the protonation site of protonated pentaglycine is located at the N-terminus, i.e. on the amide nitrogen of the peptide. We have shown in the previous section that the electronic transition due to the protonation site of pentaglycine is located in the broad feature centred at 406 eV. The protonation site is responsible for the relatively high intensity of this feature as compared to the main N 1s \rightarrow $\pi^*(CONH)$ resonance. Thus, we can use this resonance to map the difference between the two glycine-based peptides.

At the N K-edge, see Fig. 4, we observed clear differences and similarities. For both protonated peptides, a resonance is located at 401.6 eV and is due to an N 1s $\rightarrow \pi^*(CONH)$ electronic transition in the peptide bonds blue-shifted by 0.1 eV for G₃ compared to G₅. The resonance at 402.6 eV is not observed for protonated triglycine but we observed a feature at 403.1 eV that has not been observed for protonated pentaglycine. For G_4H , a resonance is observed at 403.1 eV for N 1s \rightarrow σ^* (NH). Note that in the NEXAMS spectrum of solid triglycine, no resonance between the one at 401.5 eV and the broad peak centred at 406 eV has been observed.²² In the work of Gordon and co-workers22 as well as of Zubavichus and co-workers21 on solid-state triglycine, they attributed the feature at 406 eV to $\sigma^*(NH)$ transitions from the amide group that is protonated in

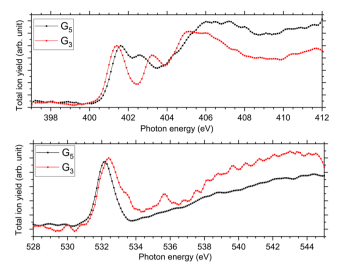


Fig. 4 Experimental NEXAMS spectra at the N K-edge (top) and at the O K-edge (bottom) of G₅ in black and G₃ in red.

the zwitterionic form, which is consistent with our conclusion on the protonation site of peptides. This suggests that the peak we observe at 403.1 eV in the N K-edge spectrum of G₃ could be due to an electronic transition from a different protonation site than the one of pentaglycine. This being said, both protonated peptides exhibit this broad feature between 404 and 408 eV that we attributed previously to excitations to the $\sigma^*(NH)$ and $\sigma^*(CH)$ orbitals, with an intense contribution from an $-NH_3^+$ protonated amino group. This suggests, in contrast, that triglycine would also be protonated at its N-terminus. Thus, from the experimental spectra only, we can highlight differences but cannot conclude whether triglycine has one or two protonation sites, as the experimental NEXAMS spectrum could suggest two different ones. A comparison with DFT/ROCIS calculations is shown in the next section for a better assignment of the different resonances.

At the O K-edge, see Fig. 4, both spectra show an intense resonance centred at 532.2 eV, blue-shifted by 0.2 eV for G₅ compared to G3. This feature has previously been observed in protonated peptides^{33,63,66} and proteins^{67,68} as well as for gasphase diglycine by Feyer and co-workers.²⁶ This resonance is attributed to an O 1s $\rightarrow \pi^*(CONH)$ transition in the peptide bonds. In their study, Gordon and co-workers also attributed the resonance at 532.2 eV of solid diglycine and triglycine to an O 1s $\rightarrow \pi^*(CONH)$ resonance with a small contribution from the O 1s $\rightarrow \pi^*(COO^-)$ electronic transition.²² G₃ exhibits a shoulder at 533.4 eV that is not observed in the case of G₅ and was not observed for gas-phase diglycine.26 Dörner and coworkers attributed this transition to an O 1s $\rightarrow \pi^*(COOH)$ transition in the carboxylic group which was observed at 532.5 eV for leucine-enkephalin. 15

In the case of protonated triglycine, a second resonance is observed at 535.6 eV that is not observed in the experimental spectrum of G5. This resonance has been attributed to an O 1s $\rightarrow \pi^*(C=0)$ transition in the carboxylic group by Feyer and co-workers for gas-phase glycine and diglycine.26 The spectrum

of protonated triglycine at the oxygen K-edge does not exhibit any feature that at first glance would be associated with the protonation site.

To more deeply analyse the NEXAMS spectra for triglycine and understand better our experimental observation and the differences with pentaglycine, we compared the result of the NEXAMS experiment of protonated triglycine with DFT/ROCIS calculations.

N K-edge

In the case of protonated triglycine, a random structure search was performed in order to get different structures of this protonated peptide. Theoretically, the structure search has been performed at 0 K to probe the lowest-energy conformers of triglycine that should compare with our experimental conditions, where the ions in the trap were cryogenically cooled down to 8 K. The result of the conformer search and geometry optimization gives two main protomers, noted GGGH1 and GGGH2. The calculated structures of GGGH1 and GGGH2 are shown in Fig. 5. The first protomer GGGH1 has a more folded structure, with hydrogen bonds between the protonated N-terminus and the carboxyl oxygen atoms. In contrast, the GGGH2 structure is more linear, with the proton bound to the first amide oxygen forms a hydrogen bond with the terminal nitrogen.

In the theoretical study of Li and co-workers, they found that the most stable conformer of the traditional N-terminal protonation site of triglycine is 2.53 kcal mol⁻¹ more stable than the most stable one of the O-protonated conformers at the highly accurate coupled cluster singles and doubles (CCSD) level of theory. 10 Thus, it is expected that the experimental spectrum will be reproduced by mainly the calculated spectrum of GGGH1.

Fig. 6 presents the results of the DFT/ROCIS calculations for the absorption spectra of both isomers at the nitrogen K-edge.

Fig. S29 and S31 (ESI†) show the respective calculated DFT spectrum of GGGH1 and GGGH2 at the N K-edge with the different molecular orbitals responsible for the main transitions. At first glance, the two calculated spectra are showing strong differences, especially in the region from 402 eV to 408 eV and neither spectrum reproduces the experimental spectrum well overall. As for the other peptides, the DFT/ROCIS transition lines of the first resonance at 401.6 eV stem from N 1s $\rightarrow \pi^*(CONH)$ transitions in the peptide bonds for both protomers. According to our calculations, in the case of GGGH1, the resonance at 403.4 eV

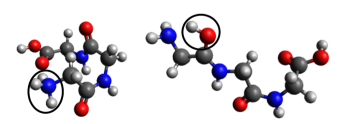


Fig. 5 Calculated structures for GGGH1 (left) and GGGH2 (right) (B3LYP/ ZORA-DEF2-TZVP). The dark circles highlight the position of the proto-

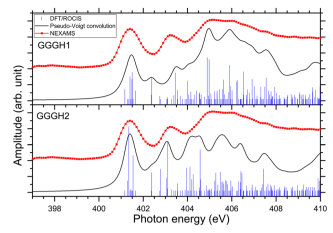


Fig. 6 Experimental nitrogen K-edge ion yield spectrum in red of G3 as well as DFT/ROCIS spectrum for the calculated structure GGGH1 (shifted +12.07 eV) and for the calculated structure GGGH2 (shifted +12.56 eV). The absorption lines were broadened using a pseudo-Voigt profile with a 0.6 eV FWHM represented in black

is attributed to a transition from N 1s to $\sigma^*(CH)$ orbitals (LUMO+4) in the backbone of triglycine, whereas in the case of GGGH2, it is an N 1s $\rightarrow \sigma^*(NH)$ transition in the unprotonated terminal-NH₂ (LUMO+2). Interestingly, the resonance calculated for the GGGH2 conformer is more intense than it is in the case of GGGH1 and reproduces better the peak observed at this energy in the experimental spectrum. The broad feature centred at 406 eV is due to many transitions as described previously for the protonated pentapeptides. For GGGH1, the DFT/ROCIS calculations reproduce well the broad feature, and the analysis of the different electronic transitions shows that, for this isomer, the feature is mainly due to N 1s $\rightarrow \sigma^*(CH)$ transitions from the backbone and N 1s $\rightarrow \sigma^*(NH)$ transitions from the protonated terminal-NH₃⁺. In the case of GGGH2, the calculated spectrum does not reproduce the high intensity of the feature between 404 and 408 eV. The main transition calculated at 404.8 eV (LUMO+10) is highly delocalized and can be due to N 1s → $\sigma^*(CH)$ or N 1s $\rightarrow \sigma^*(NH)$ transitions from the backbone. The main resonances observed for G3 at the nitrogen and oxygen Kedges are summarized in Table 2. Ultimately, the experimental spectrum cannot be reproduced with sufficient confidence by either of the two calculated isomers, since only the spectrum of GGGH2 can explain the resonance at 403.1 eV while only the spectrum of GGGH1 reproduces the broader feature centred at 405 eV. As in the work of Wu and McMahon, the comparison of the experimental spectrum with the calculated ones suggests that both isomers might be present under our experimental conditions and contribute to the measured NEXAMS spectrum. In this experiment only the linear ion trap was cooled down to 8 K, which was conducted at the Ion Trap end-station. The other parts of the setup were at room temperature, such that we can expect to have different stable isomers of protonated triglycine transferred and cooled down in the trap, as explained by Li and co-workers¹⁰ and already shown for biomolecular ions by Warnke and co-workers using ion mobility spectrometry combined with a cryogenic ion trap.69

Table 2 Experimental energy position (exp) and proposed assignment of the spectral feature observed in the total nitrogen K-edge and oxygen Kedge NEXAMS spectra of G_3 as well as the energies of the DFT/ROCIS absorption lines (calc)

Exp (eV)	Calc (eV)	Assignment
$\overline{G_3}$		
401.6	401.5	N 1s $\rightarrow \pi^*(CONH)$
403.2	403.1	$N 1s \rightarrow \sigma^*(NH_2)$
404-408	404.8	N 1s $\rightarrow \sigma^*(CH)$
		$N 1s \rightarrow \sigma^*(NH_3^+)$
532.3	532.4	O 1s $\rightarrow \pi^*(CONH)$
535.5	534.7	O 1s $\rightarrow \pi^*(CO)$
	535.6	O 1s $\rightarrow \sigma^*(OH)$

We have therefore attempted to estimate the relative populations of the two isomers present in our experiment by mixing the contribution of both isomers into a new calculated spectrum and by comparing the peak intensities with the experimental spectrum. The two resonances that we considered as markers for the comparison are the one at 403.1 eV to obtain the contribution of GGGH2 to the experimental spectrum and the broad feature centred at 405.2 eV to obtain the contribution of GGGH1, both of which are related to the protonation state of the N-terminus. To do this, we normalized the experimental spectrum and the calculated spectra to the first resonance, which is the peptide bond resonance and is similar in both calculated spectra. After normalization, we compared, for different isomer mixing ratios, the relative intensity between the calculated and experimental peak height for both resonances. The results are shown in Fig. 7. On the basis of the spectral comparison, we concluded that our experimental spectrum arises from a relative population of N-protonated triglycine of 30% with 70% of O-protonated triglycine. This ratio is the closest one to the experimental spectrum with 90% of the relative intensity for both resonances.

The result of the isomer mixing on the calculated absorption spectra is shown in Fig. 9(a). This is in very good agreement with the results of Voss and co-workers who performed IR-IR double resonances of protonated triglycine in an ion trap at 10 K and concluded with a relative population of 35%

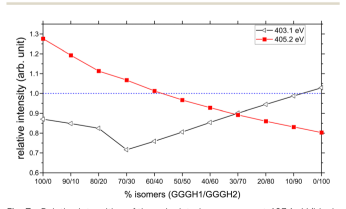


Fig. 7 Relative intensities of the calculated resonances at 403.1 eV (black, σ^* (NH) of the unprotonated – NH₂ group) and at 405.2 eV (red, σ^* (CH) from the backbone), compared to the experimental ones, as a function of the isomer mixing ratio GGGH1 and GGGH2. The dashed blue line represents a relative intensity of 1.

N-protonated and 65% O-protonated. 70 The authors also point out that this population is more representative of the isomers present at 300 K originating from the ESI source rather than the 0 K population distribution.

O K-edge

Fig. 8 shows the NEXAMS spectrum obtained at the oxygen Kedge and its comparison with DFT/ROCIS calculations for the two main isomers of protonated triglycine. Fig. S30 and S32 (ESI†) show the respective calculated DFT spectrum of GGGH1 and GGGH2 at the O K-edge with the different molecular orbitals responsible for the main transitions. The different resonances are summarized in Table 2. For GGGH1, the first resonance at 532.4 eV is attributed to O 1s $\rightarrow \pi^*(CONH)$ transitions (LUMO and LUMO+2) in the peptide bond. The other main transition line at 533 eV, which is shown in the calculated spectrum, can be attributed to O 1s $\rightarrow \pi^*(CO)$ (LUMO+1) in the carboxylic group starting from the oxygen atom of the -C=O group. In contrast, in the case of GGGH2, we observed only the first resonance related to the O 1s \rightarrow $\pi^*(CONH)$ transition. For GGGH1, the Pseudo-Voigt convolution of the DFT/ROCIS spectrum results in a much broader peak, closer to the experimental result.

For both calculated spectra, the second feature includes two resonances, at 534.7 eV and 535.7 eV for GGGH1 (534.9 eV and 535.9 eV for GGGH2), but only one was observed experimentally, centred at 535.6 eV. In the case of GGGH1, the first resonance at 534.7 eV is due to an O 1s $\rightarrow \pi^*(CO)$ transition to the LUMO+1 in the carboxylic group but starting from the oxygen atom of the -OH group, and the second is attributed to O 1s $\rightarrow \sigma^*(OH)$ transitions (LUMO+5) in the carboxylic group of protonated triglycine. For GGGH2, the first resonance can be attributed to the peptide bond resonance O 1s $\rightarrow \pi^*(CONH)$ in the molecular orbitals LUMO and LUMO+1, whereas the resonance at 535.9 eV is due to O 1s \rightarrow $\sigma^*(OH)$ transitions (LUMO+8) at the C-terminus of the peptide, similar to GGGH1.

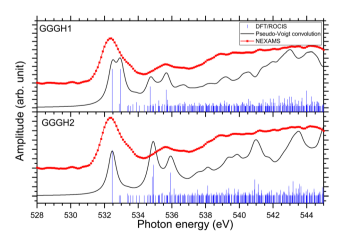


Fig. 8 Experimental oxygen K-edge ion yield spectrum in red of G3 as well as DFT/ROCIS spectrum (shifted +14.15 eV) for the calculated structure GGGH1 and for the calculated structure GGGH2 (shifted +14.76 eV). The absorption lines were broadened using a pseudo-Voigt profile with a 0.6 eV FWHM represented in black

At higher energy, the spectra are mainly constituted of excitations to σ^* orbitals. These results are in good agreement with the previous study of Feyer and co-workers for neutral gasphase diglycine.26

To confirm the results obtained at the nitrogen K-edge, we combined the two calculated spectra at the oxygen K-edge with the same factor, 30/70, for GGGH1 and GGGH2 as in the previous section. The resulting graph is shown in Fig. 9(b). Here, we achieve a good agreement with the experimental spectrum. We reproduced the different experimental resonances with the calculations. Despite the relative intensity of the resonance at 535 eV being higher for the calculated spectrum than the experimental spectrum, the attributed electronic transitions agree well with our results and the previous studies. 22,23,26

In this section, we have shown that there are clear differences between a protonated amide nitrogen and a protonated amine oxygen and that the difference is clearly detectable in the N K-edge NEXAMS spectra. When the protonation site is

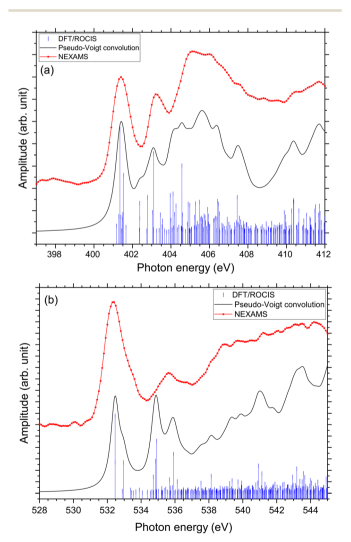


Fig. 9 Comparison of, in red, the experimental NEXAMS spectra of protonated triglycine at the N-K edge (a) and O K-edge (b) and, in black, the result of the convolution of the two calculated spectra GGGH1 and GGGH2 with 30% of GGGH1 and 70% of GGGH2.

located at the oxygen, we observed a resonance at 403.1 eV that is not observed for the case of a peptide protonated at a nitrogen, this resonance stems from an N 1s $\rightarrow \sigma^*(NH)$ transition in the unprotonated -NH2 N-terminus. However, our experimental spectrum still contains the signature of a protonated N-terminus with a broad and intense feature centred around 405 eV, which can be explained by our experimental conditions. In fact, during this experimental campaign, only the ion trap was held at 8 K, whereas the other parts of the setup were at room temperature, which allows different conformers to coexist in the trap. Moreover, as shown in the study of Wu and Mc Mahon,9 smaller peptides of glycine are more stable in a linear structure whereas peptides containing more than three glycines are more stable in a folded structure. It can be assumed that as triglycine is of an intermediate size, the folded structure and the linear structure can coexist. According to the simulations, protonated triglycine is more stable in a folded structure when the proton is on the N-terminus and the linear structure is the most stable with the proton on the first amide oxygen. By comparing the DFT/ROCIS calculations with the N K-edge experimental spectrum, we could confirm that the latter is representative of a mixing of protomers with a relative population of N-protonated triglycine of 30% with 70% of Oprotonated triglycine. At the oxygen K-edge, no striking difference appears between the two protomers on the calculated spectra, which indicates that the signature of the protonation site is best observed at the nitrogen K-edge. However, our measurement did not show most of the resonances expected by the theoretical work of Li and co-workers. 10 Their simulated NEXAFS spectra, at the nitrogen K-edge, did not show any feature around the energy of the peptide bond excitation, and we did not observe any π^* transition originating from the NH₂ terminus expected by Li and co-workers but a σ^* transition lying higher in energy than the expected transitions. They predicted many transitions in the region of 404-406 eV that we were not able to resolve experimentally. At the oxygen Kedge, we also observed striking differences between our experimental spectrum and their simulated one: we did not observe any peak at 531.3 eV or at 533.6 eV which they, respectively, attributed to an O 1s $\rightarrow \pi^*(C=O)$ transition only due to cyclic isomers or to linear isomers. One reason that could explain the differences between our results and their spectra can be the temperature used for the simulation, they calculated their spectra at a temperature of 498 K,10 while our ion trap was held at 8 K, which could result in different conformations of the protomers in the trap which would affect the absorption spectra. Overall, the NEXAMS technique has proven to be sensitive to the protonation site of peptides and we have shown that we can distinguish the different protomers present in the experimental setup and give an approximate ratio.

Conclusions

In conclusion, we have performed a NEXAMS study on five glycine-based protonated pentapeptides to investigate the effect

of the protonation site on the electronic structure and to record its spectroscopic fingerprint in the soft X-ray range. We have shown that the combination of DFT/ROCIS calculations and soft X-ray spectroscopy allows mapping the electronic transitions of the protonation site. The influence of a protonation site on a nitrogen atom has only a moderate effect on the entire spectrum of protonated pentapeptides. Experimentally, the NEXAMS technique does not allow for straightforward identification of the resonances imparted to the protonation site of a peptide, essentially owing to the numerous overlapping transitions to σ^* orbitals in these molecular systems and the intrinsic band broadening imparted to vibrational excitations in the core-hole excited state (Franck-Condon principle). However, we could observe that the high intensity of the feature centred at 406 eV was due to the protonation site and we discussed how certain changes in the spectral features, especially the shape and position of the resonances normally attributed to N 1s \rightarrow $\pi^*(CONH)$ and N_i 1s $\rightarrow \pi^*(CON_iH)$, can be attributed to the different basic residues in the peptides, e.g. by additional transitions to $\pi^*(C = N)$ and $\sigma^*(NH)$ in the side chain of histidine and arginine. We could also identify that the production of immonium ions of a residue is quenched when the absorption takes place on its side. This coincides with the increased yield of side-chain-related fragments under the same condition, which suggests that fast bond cleavage in the side chain is induced after localized photon absorption, prior to energy redistribution in the system. In addition, we are able to answer an open question by Dörner and co-workers¹⁵ concerning the presence of a peak at 399.2 eV in protonated methionine enkephalin [YGGFM + H]+ but not in leucine enkephalin [YGGFL + H]⁺. The authors proposed, based on literature and calculation, that this peak could be attributed to either an N 1s $\rightarrow \sigma^*(N-H)$ transition in an unprotonated terminal NH₂ group or an N 1s $\rightarrow \pi^*(C=N)$ transition in a peptide bond in an enol form. In our set of experiments, peptides G₄H, G₄R and G₄K have an unprotonated terminal NH2 group but do not show a prominent peak at 339.2 eV. The same is true for the calculations of Oprotonated triglycine that do not show a transition at this energy. The assignment of this band to N 1s $\rightarrow \sigma^*(N-H)$ can thus be excluded.

Moreover, we have demonstrated that the NEXAMS technique is sensitive to backbone protonation as demonstrated by studying protonated triglycine at the nitrogen K-edge and the oxygen K-edge. We have shown that a protonation site located at an amide oxygen leads to a resonance around 403 eV that is not present in the case of a peptide protonated at a nitrogen atom. However, the measurements did not reveal as striking differences between the protomers as expected from the work by Li and coworkers. 10 We have shown that the combination of DFT/ROCIS calculations and NEXAMS can help estimate the mixing ratio between the protomers observed during the experiment. This work should also be extended to the search for deprotonation site signatures in peptides and in other organic molecules where the information on the (de)protonation site(s) could be revealed using NEXAMS. Additionally, it could be tested experimentally if signatures of conformational structures such as α -helix and β -turn can

be resolved in the N_i 1s $\rightarrow \pi^*(CON_iH)$ transitions as proposed theoretically by Buckley and Besley.⁵⁰ With this work, we can foresee that NEXAMS is perfectly suited to follow ultrafast proton transfer/migration in biomolecules, for example in pump-probe studies at X-ray free-electron laser facilities, where a pump pulse would trigger the proton transfer process and a probe X-ray pulse would interrogate the position of that proton on the molecule. In addition, NEXAMS will allow us to follow the protonation/hydrogenation change in nanohydrated peptides and locate the water molecules on the biomolecules. Finally, due to the ionizing properties of soft X-ray radiation, NEXAMS experiments do not suffer from the limitations of dissociationbased spectroscopy techniques with respect to "stable" molecules, Mⁿ⁺, that would not break, but produce a minima intact photoionized precursor ions, $M^{(n+1)+\bullet}$, from which an absorption spectrum can be retrieved.⁷¹ In these cases, NEXAMS is a valuable complementary technique to e.g. IRMPD.

Author contributions

L. S. and S. B. did the conceptualization and supervision. J. L. and L. S. did the formal analysis. A. K. and D. M. did the theoretical calculations under the supervision of R. M. and M. R. J. L., K. S., C. M., L. S., I. U., A. K., F. T., S. B. performed the experiment at the P04 beamline of Petra III. J. L., L. S., I. U., B. O., V. Z., T. L., K. H. and M. T. participated in the experiment at BESSY II. All authors participated to review and editing the original draft written by J. L.

Conflicts of interest

There are no conflicts to declare.

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