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Mass spectral signatures of complex post-translational modifications in proteins: a proof-of-principle based on X-ray irradiated vancomycin

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Mass spectral signatures of complex post-translational modifications in proteins: a proof-of-principle based on X-ray irradiated vancomycin

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ABSTRACT: Characterizing post-translational modifications (PTM) of proteins is of key relevance for the understanding of many biological processes, as these covalent modifications strongly influence or even determine protein function. Among the different analytical techniques available, mass spectrometry is attracting growing attention because recent instrumental and computational improvements have led to a massive rise of the number of PTM sites that can be identified and quantified. However, multiple PTM occurring at adjacent amino-acid residues can lead to complex and dense chemical patterns that are a challenge to characterize. By means of X-ray synchrotron radiation coupled to mass spectrometry, and through the test-case of the glycopeptide antibiotic vancomycin, we show that such a pattern has a unique and robust signature in terms of photon energy and molecular environment. This highlights the potential of this technique in proteomics and its value as a tool to understand the biological roles of PTM.

INTRODUCTION

Post-translational modifications (PTM) of proteins are essential in biology, since they tailor protein activity by cleavage and/or creation of covalent bonds. The removed and/or added chemical groups can be quite diverse: the most common PTM are phosphorylation, acetylation and glycosylation. They involve the attachment of a phosphoryl, an acetyl or a carbohydrate group, respectively, to an amino-acid side chain. Other important PTM lead to removal of a peptide from the protein N- or C-terminal side, for instance in collagen. Hydroxylation of prolines and lysines is another PTM, which for instance stabilizes the collagen triple-helix structure responsible for the specific mechanical properties of connective tissues such as skin, cartilage, nails and bones. PTM can also result from oxidative stress, aromatic and sulfur-containing amino-acids being especially prone to cross-linking or oxidation.1 For instance, these processes are crucial for the synthesis of glycopeptide antibiotics such as vancomycin. The latter is a last-resort drug against Gram-positive bacteria, blocking the renewal of their cell-wall by acting as a ligand for a precursor of the main cell-wall component. Vancomycin is naturally formed by bacteria but has also been synthesized.2 Once the seven-residue peptidic chain has been created, several PTM have to occur in the five tyrosine side-chains. They involve the addition of OH groups, cross-linking, the addition of chlorine atoms to two of these side-chains, and glycosylation of the central one (see Fig. 1).

The numerous and dense cross-linkages in the resultant peptide create a rigid binding pocket allowing for stereospecific recognition of the receptor via noncovalent binding. Identifying such a complex modification is extremely challenging, even for state-of-the-art methods such as capillary electrophoresis or mass spectrometry (MS) techniques.3 The latter have proven to be very powerful for deciphering the site and identity of PTM in proteins such as histones, which undergo a particularly large number of PTM.4 Generally, two approaches can be followed: top-down or bottom-up. In the first, gas-phase intact proteins are cleaved into fragments whose mass-over-charge (m/z) ratio allows identifying PTM. In the second, proteins undergo enzymatic digestion that produces peptides, which are identified thanks to MS and fragmentation techniques. Protein and peptide fragmentation is often performed by collision-induced dissociation (CID) using a rare gas or N2, and when coupled to tandem MS, CID has allowed identifying disulfide bridges in conotoxins.5 However, electron-capture or transfer dissociation are better suited for assigning labile PTM such as O-linked glycosylation or phosphorylation of serines or threonines.5 A combination of electron-transfer dissociation and CID has been applied for studying cyclotides containing numerous cysteine knots.6 VUV photoionization has recently been proposed for the identification of multiple PTM in histones.7,8 In 2019, photodissociation using a UV laser revealed mass spectral signatures of dityrosine cross-linking in peptide dimers.9
molecules. concentration with 1 % of formic acid to protonate the have been prepared in 50:50 (volume ratio) water/methanol at 50 purchased from Sigma-Aldrich as powders of over 80 % gas injected into the ion trap during the filling process. Trapping is facilitated by collisions with a helium buffer subsequently accumulated in a 3D radiofrequency ion Paul over-charge selected with a quadrupole mass-filter (QMF) and an ion funnel and guided into an octopole before being mass- the molecular ion beam is then focused into the source and transported into the vacuum chamber through a systems and synchrotron radiation. Briefly, protonated the growing field of PTM identification and quantitation in work extends the potential of mass spectrometry techniques in labile ones, can be identified with mass spectrometry. Our heavy-modified peptide, we prove that all PTM, not only complex and dense PTM. Using vancomycin as a model technique, gives robust mass spectral signatures of particularly soft X-ray photoionization, as a biomolecular fragmentation, and hence for the main fragments observed after photoabsorption. Arrows indicate on which side of the cleaved bond the charge is located. The groups #1 and 2 containing phenyl rings are encircled.

In this article, we report the proof-of-principle that soft X-ray photoionization, as a biomolecular fragmentation technique, gives robust mass spectral signatures of particularly complex and dense PTM. Using vancomycin as a model heavily-modified peptide, we prove that all PTM, not only labile ones, can be identified with mass spectrometry. Our work extends the potential of mass spectrometry techniques in the growing field of PTM identification and quantitation in proteins.

EXPERIMENTAL

Vancomycin hydrochloride and Ac₂SOAc²A have been purchased from Sigma-Aldrich as powders of over 80 % purity, and used without further purification. Solutions have been prepared in 50:50 (volume ratio) water/methanol at 50 µM concentration with 1 % of formic acid to protonate the molecules.

A home built tandem mass spectrometer, described in detail elsewhere, has been used to record mass spectra of the photoproducts from the interaction between molecular systems and synchrotron radiation. Briefly, protonated molecular systems are produced with an electrospray ion source and transported into the vacuum chamber through a heated capillary. The molecular ion beam is then focused into an ion funnel and guided into an octopole before being mass-over-charge selected with a quadrupole mass-filter (QMF) and subsequently accumulated in a 3D radiofrequency ion Paul trap. Trapping is facilitated by collisions with a helium buffer gas injected into the ion trap during the filling process. Molecular ions have been irradiated by X-ray photons at the U49-2 PGM-1 beamline of the BESSY II synchrotron (Helmholtz-Zentrum Berlin). Photon beam exposure of the trap content, typically during 300 to 1000 ms, is controlled with a mechanical shutter in order to guarantee that more than 90 % of the product cations result from the absorption of a single photon. To do so, the irradiation time is tuned to induce a depletion of the precursor ion below 10 %. Since the absorption of multiple photons is a sequential process at these fluxes (10¹²-¹³ s⁻¹), the absorption events are independent, thus a probability p for absorbing one photon gives the probability p² for two photons. Neglecting the absorption of more than two photons, we obtain p²+p < 0.1 and thus p < 0.09. Precursor ions and cationic fragments are then extracted from the trap, analyzed by a time-of-flight reflectron mass spectrometer, and detected by microchannel plates. Mass spectra of the non-irradiated trap content (beam-off) and irradiated residual gas are recorded as well, the latter allowing to spot background peaks due to photoionization of residual gas molecules. Then, the beam-on mass spectrum is subtracted from the beam-off one, and the resulting spectrum shows the precursor ion depletion with a negative intensity. Assuming that absorption of one photon leads to ionization and/or fragmentation of the precursor ion, this depletion (area under the peak) is proportional to the total photo-absorption yield. All relative yields have been obtained by calculating the area under each peak, normalizing by the precursor ion depletion, by the total yield of all cationic species formed by photoabsorption, and by detection efficiency.

RESULTS AND DISCUSSION

The soft X-ray photoabsorption spectra of doubly-protonated vancomycin [V+2H]²⁺ are shown in Fig. 2. In the spectrum at 100 eV, we mainly observe the same species as we have previously observed after photoabsorption in the VUV range. The peak at m/z 483.8 corresponds to non-dissociative ionization (NDI) of the precursor ion, a minor process compared to fragmentation following ionization. The yields of the complementary Y₀²⁺ and B₀²⁺ fragments are significantly higher (see Fig. 2). Both are due to cleavage of the glycosidic bond (linking the carbohydrate group to the rest of the molecule) after ionization. X-ray photoionization thus allows identifying the mass of the carbohydrate involved in glycosylation. The most intense peaks are assigned to the O²⁺ and c₁²⁺ fragments of the pseudo-peptidic backbone (see Fig. 1). Note that the peak corresponding to the latter (at m/z 144) can also be attributed to B₁²⁺, formed by cleavage of the glycosidic bond within the carbohydrate moiety. The relative yield of NDI and large fragments is smaller than in the case of photons in the 14-30 eV range, which can be explained by a rise of the vibrational energy transferred by the photon to the molecular system with photon energy, as it has previously been shown in the soft X-ray range. With increasing photon energy, a more striking difference develops in the mass spectra. Whereas VUV photoabsorption solely leads to fragmentation into peaks at m/z 100, 118, 127, 144, 149, a multitude of additional peaks with m/z < 300 are induced by soft X-ray photoabsorption (see Fig. 2). In the following, we will show that these peaks can be assigned to internal fragments whose formation requires at least two bond cleavages (see below). The total relative yield of these internal fragments increases with photon energy up to 300 eV, and is compensated by the fall of non-dissociative ionization and all
of singly-protonated vancomycin reported after CID at 4 keV kinetic energy on argon, but are not mentioned and their exact mass is not possible to obtain from the figure.\textsuperscript{16} Since these high-energy collisions induce ionization, like X-rays, this seems to be consistent with these peaks coming from fragments being formed after ionization. In Fig. 3, one can notice that for soft X-ray photoabsorption, groups of peaks of similar intensity appear: they are separated by one mass unit, indicating singly-charged fragments. All precursor ions containing isotopes were present in the trap after \( m/z \) selection by the QMF (see the experimental section), but even considering the maximum number of carbon and chlorine atoms possible (22 carbons or 16 carbons and 2 chlorines) for the fragment corresponding to the peak of highest mass (270 amu), the natural abundance of \( ^{13}\text{C} \) and \( ^{37}\text{Cl} \) cannot explain these patterns. Thus, they are probably due to extensive H scrambling, which has also been found to occur in photoionized 3-aminophenol,\textsuperscript{17} an aromatic molecule containing one phenyl ring with one OH group. Indeed, these groups of peaks are typical of mass spectra of neutral H-rich molecules containing aromatic rings,\textsuperscript{e.g.} ionized by electron impact,\textsuperscript{18} ion impact at keV or MeV energy,\textsuperscript{19–22} or by absorption of X-ray photons.\textsuperscript{23,24} A good example is ionization of lorazepam, a molecule containing two aromatic rings substituted by one chlorine atom as in vancomycin: groups of peaks separated by 1 mass unit are also observed in the same mass range.\textsuperscript{18} Ionization of small carbohydrates (such as that of vancomycin) also yields such groups of peaks, but in a much lower mass range, typically below 100 amu.\textsuperscript{25–29} Therefore, we have calculated the mass of potential fragments originating from the groups containing aromatic rings in vancomycin. The results are shown in Scheme 1, where structures of fragments accounting for all groups of peaks seen in the mass spectra are proposed. In Fig. 3, we include vertical bars at the \( m/z \) of these singly-charged fragments with the maximum number of H atoms (leaving the bond orders unchanged), and without any H atom left. These bars mark off each group of peaks. To confirm that the natural abundance of \( ^{13}\text{C} \) and \( ^{37}\text{Cl} \) isotopes does not account for the observed peaks, we also show the simulated isotopic pattern of each fragment in Fig. S1 (see the Supporting Information). It demonstrates that extensive H atom scrambling occurs for these fragments. Interestingly, their formation can be explained by only three pathways, two of them involving the central group containing the three phenyl rings (noted 1 in Fig. 1) and accounting for most of the groups of peaks. The remaining groups are assigned to fragments formed by a third pathway, starting with separation of the biphenyl group bound to the C-terminus of the pseudo-peptidic backbone (noted 2 in Fig. 1). In addition, each aromatic ring can be identified separately. Thus, these groups of peaks assigned to internal fragments can be considered as a mass spectral signature of the complex PTM pattern of vancomycin. Internal fragments are not often exploited in MS of biopolymers, because of the intricate data analysis required. Here, the latter is made easier by the following features: the peaks fall at integer \( m/z \) values, are easily separated even with a modest resolving power, and are structured in distinct groups of masses separated by 1 amu. Another important point is the influence of photon energy on this mass spectral signature: it can be deduced from Fig. 3. Despite the large energy range (100-531.5 eV) covering C, N and O K-edges, the relative intensity of these peaks varies only very slightly, the overall shape of the groups of peaks
remaining remarkably stable. This is very important if they are used as mass spectral signature for identifying PTM. Consider also that we have shown in the previous paragraph that the total yield of these fragments increases with photon energy from 100 to 300 eV because of the increase in vibrational energy transferred to vancomycin after ionization. Therefore, this behavior might be traced to a high potential energy barrier for creating the largest fragments shown in Scheme 1, due to the need to cleave several bonds, and lower barriers separating the subsequent smaller fragments.

![Figure 3](image_url)

**Figure 3.** Zoom into the region of the mass spectra of [V+2H]^{2+} (see Fig. 2) where the internal fragments appear. Each of them is indicated by a letter above the corresponding group of peaks. Vertical bars correspond to the m/z of each singly-charged fragment shown in Scheme 1 with the maximum number of H atoms (leaving the bond orders unchanged), and without any H atom left. The structure of these fragments is given in Scheme 1.

**Scheme 1.** Proposed structure corresponding to the internal fragments of [V+2H]^{2+} (see Fig. 3). The bond cleavages responsible for their formation are highlighted.

- A: [K+H]^{+}
- B: [A+H]^{+}
- C: [T+H]^{+}
- D: [M+H]^{+}
- E: [L+H]^{+}
- F: [K+H]^{+}
- G: [K+H]^{+}
- H: [R+H]^{+}
- I: [V+H]^{+}
- J: [V+H]^{+}

The next question is to know whether the mass spectral signature of the PTM pattern of vancomycin is sensitive to a non-covalently bound molecular environment. To tackle this question, we have studied the vancomycin dimer as well as complexes between vancomycin and Ac-K{sup +}P{sup +}A (K is lysine and A alanine), a peptidic model of its receptor, abbreviated R in the following. The chemical structure of the receptor can be found in Fig. S2. In all cases, absorption of one X-ray photon in the 100-531.5 eV range leads to ionization and fragmentation of the precursor molecular system. For instance, Fig. 4 shows that at 100 eV, the [V+R+2H]^{3+} complex dissociates and subsequent intramolecular fragmentation of vancomycin and R occurs. Fragments of the latter have been identified by comparison with the mass spectra of [R{sup +}+H]^{+} (see Fig. S2). The same fragments as for VUV photons are observed, but large fragments are less abundant, as in the case of isolated vancomycin (cf. Fig. 2). Small fragments rise with photon energy from 100 to 300 eV, the total fragmentation yield slightly increasing for both vancomycin and R. Interestingly, the fragmentation yield of vancomycin is compensated by the falling yield of [R+H]^{+} and vice versa, which is unexpected. In our previous study, we have attributed the formation of [R+H]^{+} to a proton transfer between ionized vancomycin and neutral R. Therefore, the decrease in the [R+H]^{+} yield while vancomycin fragmentation increases might indicate that proton transfer becomes progressively less likely as photon energy increases. This might be due to more vibrational excitation as the X-ray photon energy rises from 100 to 300 eV, leading to the noncovalent complex dissociating on a faster timescale and quenching proton transfer. This is plausible because MacAleese et al. have reported that proton transfer can take up to hundreds of microseconds in an ionized peptide radical cation. These results show that photon energies over 300 eV give the highest yield of vancomycin fragments. Among the latter, and for all noncovalent systems studied here ([V+R+2H]^{3+}, [V+2R+2H]^{3+}, and [V+3H]^{3+}), we observe the same internal fragments as for isolated vancomycin, their relative yield increasing with photon energy up to 300 eV (see Fig. 4 and S2-S4). We can also notice that whatever the noncovalent system, the overall relative yield of the internal fragments of vancomycin is lower than in the case of isolated vancomycin. This is consistent with our hypothesis of fragmentation in the ground-state due to vibrational energy transfer, this energy being redistributed into more degrees of freedom when the system gets larger. Furthermore, if we focus on the pattern made by the peaks attributed to these internal fragments, we observe a high similarity whatever the molecular environment (cf. Fig. 5). This indicates that the mass spectral signature is robust and can be used to identify the groups responsible for the fragments, and thus the corresponding post-translational modifications in proteins.
CONCLUSIONS

In this contribution, we have shown that X-ray photoabsorption of a highly modified tyrosine-rich peptide coupled to mass spectrometric analysis of the resulting fragments gives a signature that can be employed to characterize particularly complex and dense PTM, especially those resulting from oxidative stress. Indeed, this mass spectral signature is robust with respect not only to the X-ray photon energy over a large range (100-531.5 eV), but also to the molecular environment of the peptide. This proof-of-principle expands the potential of mass spectrometry techniques in proteomics. Further work is now required to test the applicability of this method to other peptides, proteins and a wider range of PTM.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Simulated mass spectra of singly-charged internal fragments of vancomycin shown in Scheme 1; Photoabsorption mass spectra [R₂+H]⁺, [V₂+3H]³⁺ and [V+R₂+2H]²⁺; Total yield of internal fragments of vancomycin in different environments (PDF)

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All authors have given approval to the final version of the manuscript.

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Gas-phase X-ray photoabsorption coupled to mass spectrometry of a heavily-modified peptide provides a robust signature of its post-translational modifications.