

RESEARCH ARTICLE

Characterization of Tyrosine Nitration and Cysteine Nitrosylation Modifications by Metastable Atom-Activation Dissociation Mass Spectrometry

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Abstract

The fragmentation behavior of nitrated and S-nitrosylated peptides were studied using collision induced dissociation (CID) and metastable atom-activated dissociation mass spectrometry (MAD-MS). Various charge states, such as 1+, 2+, 3+, 2-, of modified and unmodified peptides were exposed to a beam of high kinetic energy helium (He) metastable atoms resulting in extensive backbone fragmentation with significant retention of the post-translation modifications (PTMs). Whereas the high electron affinity of the nitrotyrosine moiety quenches radical chemistry and fragmentation in electron capture dissociation (ECD) and electron transfer dissociation (ETD), MAD does produce numerous backbone cleavages in the vicinity of the modification. Fragment ions of nitrosylated cysteine modifications typically exhibit more abundant neutral losses than nitrated tyrosine modifications because of the extremely labile nature of the nitrosylated cysteine residues. However, compared with CID, MAD produced between 66% and 86% more fragment ions, which preserved the labile -NO modification. MAD was also able to differentiate I/L residues in the modified peptides. MAD is able to induce radical ion chemistry even in the presence of strong radical traps and therefore offers unique advantages to ECD, ETD, and CID for determination of PTMs such as nitrated and Snitrosylated peptides.

Key words: Posttranslational modifications, PTMs, Nitration, Nitrosylation, Activation techniques, Fragmentation techniques, Electron transfer dissociation, ETD, Electron capture dissociation, ECD, Metastable atoms, Penning ionization, Peptide sequencing

Introduction

E lucidating how a PTM can alter a protein's structure and biological function is an integral part of proteomics. The

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ability to identify and localize these modifications, in particular nitration and S-nitrosylation, can aid in biomarker discoveries for neurological diseases [1, 2], cardiovascular diseases [3, 4], lung diseases [5, 6], and diabetes [7]. Tyrosine and cysteine amino acid residues are highly susceptible to oxidation by one or more forms of reactive nitrogen species such as nitric oxide radicals [8]. For example, reaction of a nitrogen dioxide radical with tyrosine can produce a covalently bound $-NO_2$ to the tyrosine side chain thereby producing 3-nitrotyrosine [9]. Cysteine nitro-

Received: 23 September 2010 Revised: 13 November 2010 Accepted: 14 November 2010 Published online: 29 January 2011 sylation can occur through a reaction with nitric oxide radicals resulting in the addition of -NO [10]. Developing rapid and accurate mass spectrometry (MS)-based methods for direct detection of both nitration and S-nitrosylation can aid in the understanding of how the reactive nitrogen species plays a role in certain diseases and can provide new diagnostic tools.

MS has become a leading tool in proteomics because of its ability to sequence peptides and proteins and characterize PTMs. The most widely used MS techniques for studying nitrotyrosine and nitrosylated cysteine PTMs in peptides and proteins are matrix assisted laser desorption/ionization timeof-flight MS (MALDI-TOF) and electrospray ionization (ESI) MS/MS [9]. UV-MALDI has been noted to lead to a high degree of photodegradation or in-source decay of nitrated or S-nitrosylated peptides, thereby rendering the modified peptides weak or absent [9, 11, 12]. The dissociation method traditionally employed in ESI MS/MS is CID, and CID tends to preferentially cleave side chain PTMs instead of the peptide backbone. The preference for side-chain cleavages leads to poor sequencing information and PTM characterization. Biotin-switching, in which the labile nitrosylated cysteines are derivatized to a more stable biotinylated cysteine, was developed to combat these issues [13]. However, the precise control of the chemical reactions has proven to be difficult and can result in false positives [12, 14]. Methods such as precursor ion scanning of the nitrotyrosine immonium ions [15] and selected reaction monitoring combined with UV absorption [16] have both shown to be promising techniques to characterize nitrotyrosine when analyzing complex samples.

Alternative types of peptide dissociation methods that utilize radical chemistry have been recently introduced to provide both accurate sequencing and PTM information. ECD, first introduced in 1998 by McLafferty and coworkers, has proven to effectively fragment the peptide backbone in Fourier transform ion cyclotron resonance mass spectrometers (FT-ICR-MS) [17]. This method preferentially cleaves the more energetic peptide backbone N-Ca bonds resulting in mainly c-/z-type ions, in contrast to the "slowheating" dissociation methods such as CID and IRMPD [18]. The latter activation methods tend to cleave the lower energy C-N bonds and produce b-/y-type ions. ETD, introduced by Hunt and co-workers, uses ion-ion reactions to transfer electrons from a radical anion to polycations resulting in ECD like fragmentation [19]. Successful PTM site determinations of phosphorylation [19, 20], sulfation [21, 22], and glycosylation [23–25], among others, have been reported for ECD and ETD. However, traditional ECD and ETD are not applicable to singly protonated or negatively charged precursor ions such as the more acidic peptides. Recently, new techniques have been developed to apply these two methods, NETD and niECD, to negative ions [26-28].

There have been only a few reports applying ECD and ETD to nitrated and S-nitrosylated peptides. It has been

shown that in ECD and ETD, backbone fragmentation is completely inhibited in peptides containing specific tags with electron affinities (EA) larger than 1 eV. Such tags are termed electron predators, and examples include 3-nitrobenzyl and 3,5-dinitrobenzyl moieties [29]. Beauchamp and co-workers proposed that the propagation of the captured electron can relax through space or through bond (π) to high EA predators (tags), which competes with the traditional electron transfer to the amide π^* orbital. A hydrogen atom from a separate site on the peptide can then be transferred to the electron predator radicals due to their high H-atom affinity. Tureček, on the other hand, proposes intermolecular H-atom transfer from the aminoketyl radical of the chargereduced species to the nitrobenzyl moieties instead of direct electron sequestering [30]. In this mechanism, backbone fragmentation is inhibited based on the high H-atom affinity of the nitrobenzyl groups. Regardless of the exact mechanism, the presence of nitrobenzyl moieties inhibits traditional ECD and ETD N-Cα backbone cleavage. Recently, nitrotyrosine residues have been shown to hinder backbone cleavage during ECD because of the electron predator or Hatom trap effect [31]. These results demonstrate that ECD is not the method of choice to characterize nitrated tyrosine residues of 2+ and 3+ charge states of peptides [31]. However, top-down analysis of multiply charged, larger nitrotyrosine-containing proteins using ECD has shown promise when combined with other "slow heating" methods to identify PTM sites [32].

Few studies have concentrated on direct MS detection of cysteine nitrosylation, probably due to its labile nature. S-nitrosylated bovine insulin was explored with ETD resulting in only a select number of backbone cleavages and fragment ions retaining the modification, thereby hindering site determination [22]. The major dissociation pathway observed in ECD was charge reduction (electron transfer without fragmentation) with and without losses of the NO [22]. ETD was therefore unable to identify the location of the PTM.

Recently, a new type of dissociation method using metastable atoms as the electron vehicle or potential energy source has been explored in RF ion trapping instruments, which we term metastable atom-activated dissociation (MAD). The interaction of isolated precursor ions with a high kinetic energy beam of argon or helium metastable atoms produces a high degree of peptide backbone cleavages resulting in a-, b-, c-, x-, y-, and z-type ions while retaining PTMs [33]. To date, comprehensive dissociation has been demonstrated to fragment multiply charged cations and anions [33–35], 1+ cations [33, 36, 37], phosphorylated cations [33, 35, 37], disulfide bonds [35], to cleave the amide ring structure of proline [33] and to differentiate isoleucine from leucine [33, 36].

Unlike ECD and ETD, there is evidence for two competing dissociation mechanisms responsible for the type and degree of fragment ions observed in MAD. The first is Penning ionization, which has been well characterized with neutral precursor molecules and occurs when the potential energy of the metastable atom is greater than the ionization potential of the neutral [38] or biomolecular ion. It has been well documented that metastable atoms are attracted to regions of high electron density such as the lone pairs of electrons on carbonyl oxygen atoms. Penning ionization at these sites leads to the generation of radicals at the lone pair [39]. The second mechanism observed in MAD is a similar mechanism proposed for ECD and ETD and is referred to as charge reduction. The charge reduction mechanism occurs when an electron from the metastable atom is transferred to the polycation of interest. The specific electron capture mechanism in ECD and ETD of peptides and proteins is still in debate [40-43]. However, all mechanisms agree that the electron transfer process generates a radical bio-ion that can rearrange, propagate along the peptide backbone, and subsequently fragment.

In this work, we investigated the ability of He MAD to characterize nitrated and S-nitrosylated peptides. We conducted a comparison of MAD and CID on the 1+, 2+, 3+, and 2- charge states of two nitrated peptides, DRVnYIHPF and GPLEnYGFAKGPLAK, and two S-nitrosylated peptides, VFDARDnCRSAQ and EMFTYInCNHIK, as well as the respective unmodified peptides. S-nitrosylated peptides were extremely labile and often difficult to isolate. However, when combining the MAD fragmentation information from different precursor ion charge states, modification site location was achieved. We demonstrate that MAD is a viable alternative for the direct analysis of nitrated and Snitrosylated peptides and provides radical-like fragmentation where ETD and ECD fail to deliver. The unique chemistry and structural information obtained through MAD can complement CID, ETD, and ECD methods of analysis.

Experimental

Instrumentation

All experiments were performed on a modified Bruker EsquireLC QIT MS (Bruker Daltonics, Bremen, Germany) as described previously [33]. An Ion Tech FAB gun (P50 PSU; Teddington, UK) was used as the metastable atom source as reported elsewhere [33, 35]. A deflection electrode consisting of two flat-ended 4-40 screws mounted within a PEEK (polyether ether ketone) sleeve was mounted on the bottom of the FAB gun to prevent ions and free electrons from entering the trap. The electrodes were positioned 1 cm downstream of the exit orifice of the FAB gun, orthogonal to the FAB beam, with an inter-electrode spacing of 5 mm. One electrode was grounded and the other was biased to +800 V using a high voltage power supply (EL 03R 15 L; Glassman High Voltage Inc., High Bridge, NJ, USA). For pulsed operation of the FAB gun, the anode power supply was replaced with a 10 kV high voltage amplifier (ANT 10B10; Matsusada Precision Inc., Shiga, Japan). A trigger was taken from the Bruker EsquireLC at pin 15 of connector

J522 and fed to a function/arbitrary waveform generator (33250A; Agilent, Santa Clara, CA, USA), which generated a square pulse with a set pulse width and delay time depending on precursor mass. Complete pulse operation and generation has been previously described [33]. The FAB gun was placed approximately 2 cm above a 4 mm hole in the ring electrode, which enabled the metastable atoms to traverse to the center of the QIT and interact with the isolated biomolecular ions of interest. Background ions were observed in both positive and negative modes due to Penning ionization of residual pump oils, which could be actively removed from the trap by employing a trapping voltage corresponding to a LMCO value above m/z 200 for positive mode and m/z 300 for negative mode. Before use each day, the FAB gun was turned on for 30 min to "burn" away residual contaminants such as pump oil. This helped decrease background ion signals.

Preparation of Peptides

The S-nitrosylated and non-nitrosylated forms of two endothelial nitric oxide synthase (eNOS) peptides (194-204; 205-215) and non-nitrated angiotensin II were provided by Protea Biosciences Inc. (Morgantown, WV, USA). Nitrated GPLEnYGFAKGPLAK was purchased from Alta Bioscience (Birmingham, UK). Trimethylnitrate (TMN), AccuGENE Tris-Borate buffer, methanol (HPLC grade), and glacial acetic acid were purchased from Sigma-Aldrich (St. Louis, MO, USA). All peptides were reconstituted in a 1:1 mixture of methanol and water with either 1% acetic acid or ammonium hydroxide (depending on desired charge state) to provide solutions between 20 and 40 µM of the individual peptides. Ultrahigh-purity helium and argon (Airgas, Parkersburg, WV, USA) were further purified on-line with a noble gas purifier (HP2; VICI, Houston, TX, USA). Angiotensin II was nitrated by incubating for 1 h at 37°C with 11 mM TNM in 5 mM Tris-Borate (pH 8.3), as described by Lee et al. [44]. Samples were desalted by using C₁₈ SpinTips Sample Prep Kit, using protocol 2 (Protea Biosciences Inc., Morgantown, WV, USA).

Method

All peptides were directly injected using an electronic syringe pump (BM-1000; Protea Bioscience Inc., Morgantown, WV, USA) at a flow rate of 200 μ L/h. Isolation windows varied between m/z 1 and 6 depending on the labile nature of the modified precursor ion. Ion accumulation was set to 15–25 ms to reach an ion target of 20,000. Precursor ion intensity ranged from $8 \times 10^4 - 4 \times 10^5$ AU depending on peptide and precursor charge state. All fragment ion identifications and assignments were manually determined based on predicted fragmentation patterns and were within $\pm 0.8 \, m/z$ of the expected product ions. Peaks were only assigned if the intensities were at least three times the signal-to-noise ratio. The 2+ and 3+ ions were identified according

to the expected m/z value and the presence of an isotope envelope peak at m/z + 0.5 and m/z + 0.3, respectively.

CID

All cationic and anionic peptides were fragmented using the "SmartFrag" option in the Bruker EsquireLS NT 4.5 software, which applied a sweeping collisional energy starting at 20% and ending at 150% of a set value. The fragmentation time was set to 20 MS in all experiments. The collisional energy was set to 0.4–1.2 V depending on precursor mass and compound stability. CID MS/MS data was acquired in 1 min intervals, and consist of approximately 200 averaged spectra.

MAD

1+, 2+, 3+, and 2– precursor ions were exposed to a beam of high kinetic energy He metastable atoms with the FAB anode powered at 6 kV. A pulsed metastable atom beam was applied for 75–150 MS depending on the charge state of the precursor ion. The pressure in the vacuum chamber, outside the ion trap, was kept at 1.8 mbar (uncorrected) with helium leaking out of the trap and an additional 1.0 mbar (uncorrected) of the noble gas from the metastable atom source. MAD MS/MS spectra were acquired in ~2 min intervals and consist of between 300 and 450 averaged spectra. MAD MS/MS data were background subtracted using a spectrum obtained with identical conditions as above, but with the ESI source off.

Results and Discussion

Tyrosine Nitrated Peptides

Figure 1(a–f) show MAD-MS spectra of the 1+, 2+, and 3+ charge states of nitrated angiotensin II (DRVnYIHPF) and the synthetic peptide, GPLEnYGFAKGPLAK. A lower case 'n' indicates the amino acid directly to the right is modified. Both peptides had a precursor ion mass increase of 45 Da relative to the unmodified peptides, confirming the addition of an –NO₂ group. Peptide sequence inserts in Figure 1 omit those fragment ions that have undergone a neutral loss of the PTM modification.

Combining the fragmentation products for the 1+, 2+, and 3+ charge states of angiotensin II (Figure 1(a-c), respectively) a total of 26 distinct a-, b-, c-, x-, y-, and z-type fragment ions out of 42 possible backbone cleavages are observed. At least one backbone cleavage is observed between each amino acid residue for the 1+ and 2+ charge states resulting in complete sequence coverage. MAD of the 1+ precursor ion (Figure 1(a) yielded 21 total fragment ions with seven 2+ product ions and a dominant Penning ionized species, [M + H]^{2+*}. The production of 2+ product ions from 1+ precursor ions is direct evidence for Penning ionization (PI) [33, 38] and is unique to MAD in comparison to CID,

ECD, and ETD. The small insert in Figure 1(a) is the isotopic distribution of the PI product ion species showing that the most intense peak is the 1+ species. Figure 1(a) of the 1+ precursor shows 8 C–N, 6 C $_{\alpha}$ –C, and only 2 N–C $_{\alpha}$ bond cleavages. The figure also shows that the nitration modification remains mostly intact during exposure to He metastable atoms, which enables PTM site determination. In total, seven different product ion-types undergo a neutral loss of the PTM modification (indicated by * in the spectrum), however five of these seven ions are also observed with the modification intact. A neutral loss of 18 Da (–H₂O) from both the [M+H]^{2+*} and the [M+H-NO₂]^{2+*} species was observed.

Figure 1(b) is a MAD spectrum of the 2+ precursor ions. MAD resulted in 25 product ions and cleavage between every amino acid. Of note in Figure 1(b) is the neutral loss of a hydrogen radical from the charge-reduced ions, $[M + 2H]^{+\bullet}$, which is more abundant than the PI product species, [M + 2H]^{3+•}. Again the most dominant fragmentation pathway was cleavage of the C-N bonds, with nine total b-/y-type ions. However, a large number of a- and c-type ions were also observed. The neutral loss of 29 Da from the side chain of the z₄ ion produced the w₄ ion and enables the definitive identification of an isoleucine residue instead of the more ambiguous, isobaric isoleucine residue. Isoleucine undergoes a neutral loss of •CH2CH3 (29 Da), whereas leucine undergoes a neutral loss of •CH(CH₃)₂ (43 Da) [45]. The majority of the product ions, 16 out of a possible 18, contain the nitrated tyrosine modification. Neutral losses of 18 Da ($-H_2O$) were observed from the $[M + 2H]^+$ species and the precursor ion, $[M + 2H]^{2+}$, respectively. Figure 1(c) is a fragmentation spectrum of the 3+ precursor ions using MAD. In total, only 12 product ions were observed and were mostly C-N bond cleavages. The most intense product ion peaks observed are the $[M + 2H]^{2+}$ species, presumably from a hydrogen radical loss from the charge-reduced product ion, and the neutral loss of H_2O from the $[M + 3H]^{2+\bullet}$ species. The minimal amount of backbone cleavages in the electron transfer from the metastable atom to the polyprotonated precursor is consistent with observations in ETD, although MAD is inhibited to a smaller extent than ETD [31]. The nitration modification was retained in all cases when dissociating the 3+ precursor. The hydrogen radical loss from the charge-reduced species, $[M + 3H]^{2+\bullet}$, is the most intense peak within the charge-reduced isotope envelope in Figure 1(c); in fact this is the case for all positively charged peptides in this study. However, when evaluating the isotopic distribution of the charge-reduced species, electron transfer is still apparent by the observation of the $[M + 3H]^{2+\bullet}$ peak (Figure 1(c)). A mass isolation window of m/z 1 was possible in this case because of the stabile nature of the nitrated peptides studied. The smaller isolation window reduced the contribution of ¹³C to the intensity of the charge-reduced peak. Hydrogen radical neutral loss from the charge-reduced species is also noted in ECD [31, 46]. In each of the three charge states studied, the nitration modification is unambiguously identified to be bound to the fourth amino acid residue.

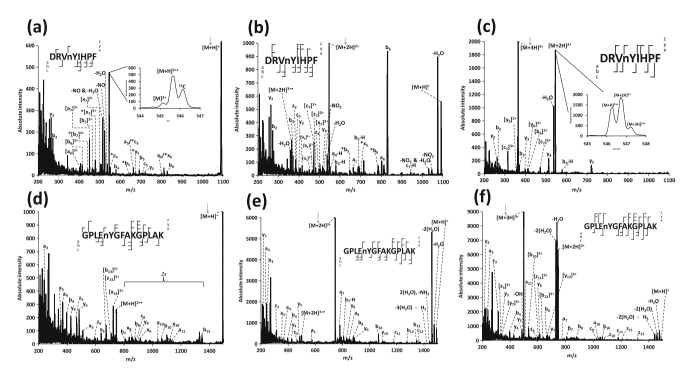


Figure 1. He MAD spectra of nitrated angiotensin II (DRVnYIHPF): (a) $[M + H]^+$, (b) $[M + 2H]^{2+}$, (c) $[M + 3H]^{3+}$, and He MAD spectra of nitrated GPLEnYGFAKGPLAK: (d) $[M + H]^+$, (e) $[M + 2H]^{2+}$, (f) $[M + 3H]^{3+}$. The precursor is indicated by an arrow and the neutral loss of the modification is indicated by an asterisk. Fragments that do not retain the modification are omitted from the peptides sequence inserts

Figure 1(d-f) show comprehensive backbone cleavage for the 1+, 2+, and 3+ precursor ions of the synthetic peptide GPLEnYGFAKGPLAK. A total of 42 distinct a-, b-, c-, x-, y-, and z-type fragment ions, out of a total of 78, are produced and complete peptide sequencing is achieved when combining different charge state product ions. MAD of the 1+ precursor ion (Figure 1(d)) produced 25 product ions as well as three 2+ product ions and the PI product ion species, [M + H]^{2+•}. The major fragmentation pathway tends to be the C_{α} -C bonds with 11 total a-/x-type ions. An additional nine C-N bond cleavages and four N-C_α bond cleavages were also observed. The observation of the w₃ ion, a neutral loss of 43 Da from the z_3 ion, allows for the differentiation of leucine from isoleucine at position 12. Out of 10 possible fragment ions containing the nitrated tyrosine, all 10 retained the modification thereby allowing for PTM site determination. The most notable feature in the MAD spectrum of the 2+ species shown in Figure 1(e) are the neutral losses from the charge-reduced species $[M + 2H]^{+ \bullet}$. The four neutral losses are 18, 36, 53, and 70 Da, which correspond to $[M + 2H - H_2O]^{+\bullet}$, $[M + 2H - 2(H_2O)]^{+\bullet}$, $[M + 2H - 2(H_2O)]^{+\bullet}$ $-NH_3^{+\bullet}$, and $[M + 2H - 3(H_2O) - NH_3]^{+\bullet}$, respectively. The first two ions (loss of 18 and 36 Da) are more abundant than the [M + H]⁺ species which corresponds to a neutral loss of a hydrogen radical from the charge-reduced species, [M + 2H]^{+•}. Similar neutral losses are also observed when dissociating the same nitrated peptide through ECD [31]. For MAD of the 2+ precursor, a total of 24 product ions are

produced with backbone cleavage between every amino acid residue with the exception of the first two, as illustrated in Figure 1(e). Again, the w₃ ion was observed, thereby enabling the 12th residue to be confirmed to be leucine. Similar to the 1+ precursor for this peptide (Figure 1(d)), no neutral losses associated with the loss of the nitrated PTM were observed and 11 fragment ions containing the nitration modification were observed. The major fragmentation pathway observed was cleavage of the C–N bonds, producing 13 b-/y-type ions.

MAD of the 3+ species, Figure 1(f), produced 28 different product ions and two charge-reduced species, $[M + 3H]^{2+\bullet}$ and $[M + 2H]^{+\bullet}$. The $[M + 2H]^{+\bullet}$ product presumably results from two subsequent charge reduction reactions. The most intense peaks within the charge-reduced isotopic distributions correspond to the neutral loss of a hydrogen radical producing the $[M + 2H]^{2+}$ and $[M + H]^{+}$ species. Again, neutral losses of 18 and 36 Da were observed for both the $[M + 3H]^{2+\bullet}$ and $[M + 2H]^{+\bullet}$ species and a neutral loss of 53 Da from the $[M + 2H]^{+\bullet}$ species. No neutral losses of the modification were observed, so 13 different PTMcontaining fragment ions were available for nitration site determination. The w₃ ion is again observed, thereby enabling the 12th residue to be confirmed to be leucine. The majority of the backbone cleavages produced b-/ytype ions as similarly noted for MAD of the 2+ species. Again, using MAD, characterization of the nitration modification was unambiguously assigned to the 5th amino acid, tyrosine.

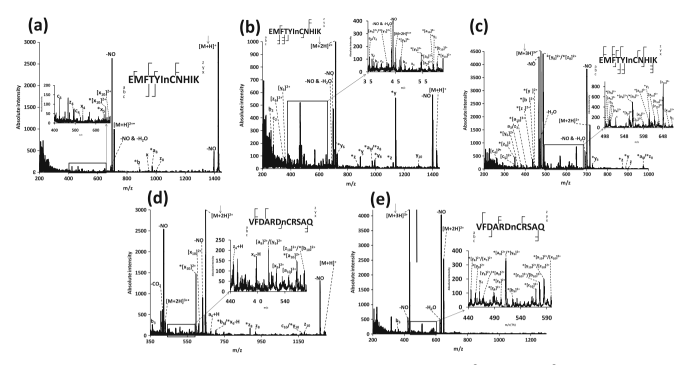


Figure 2. He MAD spectra of S-nitrosylated EMFTYInCNHIK: (a) $[M + H]^+$, (b) $[M + 2H]^{2+}$, (c) $[M + 3H]^{3+}$, and He MAD spectra of S-nitrosylated VFDARDnCRSAQ: (d) $[M + 2H]^{2+}$, (e) $[M + 3H]^{3+}$. The precursor is indicated by an arrow and the neutral loss of the modification is indicated by an asterisk. Fragments that do not retain the modification are omitted from the peptides sequence inserts

Cysteine Nitrosylated Peptides

Figure 2(a–e) show MAD spectra of the 1+, 2+, and 3+ charge states of the S-nitrosylated peptide EMFTYInCNHIK (205-215 of eNOS) and the 2+ and 3+ charge states of the Snitrosylated peptide VFDARDnCRSAQ (194-204 of eNOS). Isolation of the latter 1+ peptide was not possible because of the labile nature of this PTM. The precursor mass of each of the peptides had an increase of 29 Da relative to the unmodified peptides, confirming the addition of an -NO group. Again, the fragments with a neutral loss of the PTM modification were omitted from the peptide sequence inserts in Figure 2. Due to the extremely labile nature of the S-nitrosylation [47, 48], more gentle MS parameters and wider m/z windows of 5-6 were required to isolate the precursor ions of the modified peptides. A higher percentage of S-nitrosylated neutral losses (30 Da) were detected relative to nitrated tyrosine containing peptides. The disruption of the S-NO bond has been observed to easily break under the most modest gas phase conditions [47].

When combining the product ions observed for the 1+, 2+, and 3+ charge states of the EMFTYInCNHIK peptide, MAD produced 37, out of a possible 60, distinct fragment ions (Figure 2(a), (b), and (c)). MAD of the 1+ precursor, illustrated in Figure 2(a), produced 11 fragment ions and an intense PI product species, $[M + H]^{2+\bullet}$, with neutral losses of the S-nitrosylation from both the precursor ion and the PI product ion. Forty five percent of the backbone cleavages in Figure 2(a) were between the C_{α} –C bonds producing a-/x-type ions and 36% cleaved the N– C_{α} bond (c-/z-type ions). Of the six detected fragments that contained the modified cysteine

residue, only three retained the S-nitrosylation group. Snitrosylation PTM site determination was therefore difficult or unambiguous with the [M + H]⁺ precursor. More comprehensive dissociation was achieved for the 2+ precursor of the peptide (Figure 2(b)), producing 30 product ion types. However, 16 of the 19 fragment ions that could potentially give modification site information did not retain the Snitrosylation, thereby hindering PTM location. For MAD of the 2+ precursor (Figure 2(b)), complete peptide sequencing was achieved but the site of the PTM could not be unambiguously assigned. For the 2+ precursor 37% of the fragment ions cleaved the C-N bonds, with the majority being y-type ions. $N-C_{\alpha}$ and $C_{\alpha}-C$ bond cleavages are equally abundant at about 31% each. MAD of the 3+ precursor ion also achieved complete sequencing information by producing 35 different product ions (including those which lost the Snitrosylation) as shown in Figure 2(c). In contrast to the dissociation of the 1+ and 2+ species, 10 different product ions retained the labile S-nitrosylation, thereby successfully determining the modification site on the cysteine as opposed to the tyrosine or threonine residues. A larger percentage of product ions retain the C-terminus than the N-terminus because of the two basic amino acid residues at the C-terminus.

When combining the dissociation of both the 2+ and 3+ precursors of the VFDARRnCRSAQ peptide (Figure 2(d) and 2(e), at least one cleavage between each amino acid is observed, resulting in 21 distinct product ions in total. Cleavage of the C_{α} –C bond is the main backbone fragmentation observed when dissociating the 2+ species (Figure 2 (d)) with nine a-/x-type ions or 42% of the total product

ions. The main dissociation pathway observed is the neutral loss of 30 Da from both the PI, $[M+2H]^{3+\bullet}$, and the $[M+2H]^{+\bullet}$ species, corresponding to the loss of NO from the modification site. Figure 2(e) shows the dissociation of the 3+ species with 19 evenly distributed backbone cleavages. The two main peaks are the $[M+2H]^{2+}$ species (the charge-reduced species with a hydrogen radical loss) and the neutral loss of the NO from the $[M+3H]^{2+\bullet}$ species. For the 2+ and 3+ precursor ions, less than half of the product ions containing the nitrosylated cysteine residue retained the modification.

Effect of Modification on Peptide Backbone Cleavage

There is evidence that PTMs can have an adverse effect on the fragmentation efficiency of some electron-induced dissociation methods. Cooper and Creese show that phosphorylation modifications can interact with basic residues decreasing the amount of ECD backbone cleavages in 2+ species [49]. Therefore, determining how a certain modification affects fragmentation efficiencies and type of fragment ions observed needs to be further explored. Jones et al. provide evidence that ECD of nitrotyrosine-containing peptides severely inhibits N-Cα backbone cleavages [31]. In fact, for 2+ species, little backbone dissociation—except for small neutral losses from the charge-reduced species—is observed. More cleavages are noted for ECD of the 3+ species, most likely due to columbic repulsion extending the peptide conformation when not in close proximity to the modification [31]. This phenomenon is explained in two separate studies by Sohn et al. and Tureček, which have shown that ECD and ETD backbone fragmentation is

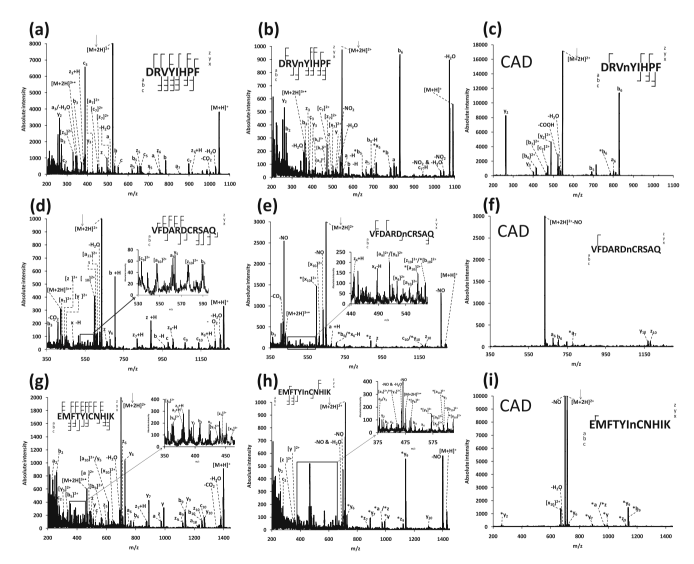


Figure 3. Comparison of the 2+ charged peptides: (a) MAD of non-nitrated DRVYIHPF, (b) MAD of nitrated DRVnYIHPF, (c) CID of nitrated DRVnYIHPF, (d) MAD of non-nitrosylated VFDARDCRSAQ, (e) MAD of S-nitrosylated VFDARDnCRSAQ, (f) CID of S-nitrosylated VFDARDnCRSAQ, (g) MAD of non-nitrosylated EMFTYICNHIK, (h) MAD of S-nitrosylated EMFTYInCNHIK, and (i) CID of S-nitrosylated EMFTYInCNHIK. The precursor is indicated by an arrow and the neutral loss of the modification is indicated by an asterisk. Fragments that do not retain the modification are omitted from the peptides sequence inserts

completely inhibited when high EA (>1 eV) nitrobenzyl moieties are present [29, 30]. S-nitrosylation has also been shown to hinder the efficiency of ETD. ETD of S-nitrosylated insulin chain B was investigated by Mikesh et al. with very little backbone cleavage and PTM retainment observed [22]. Mikesh et al. concluded that the neutral loss of the NO from the charge-reduced species could act as its own proton transfer reagent, which ultimately decreases fragmentation yields.

Comparison of MAD and CID spectra of the nitrated and S-nitrosylated peptides and their unmodified equivalents in the 2+ charge state are shown in Figure 3. MAD of the modified peptides produced enough dissociation to confer complete sequencing information, which has not been demonstrated for ECD and ETD. Comparing the amount of backbone cleavages observed with MAD and CID (see Supplemental Figure 1 and Supplemental Table 1), we note that CID alone is unable to obtain enough information to sequence the peptides and determine unambiguously the modification site locations. In fact, MAD produces almost two times more fragment ion types than CID that contain modification information. Figure 4 and Supplemental Table 2 compares the total number and types of ions for the spectra in Figure 3 produced for the 2+ charge state of the unmodified and modified peptides. We observe a slight decrease in the number of backbone cleavage types when either modification is present (nitrosylated cysteine or nitrated tyrosine), with reductions between 5% and 42%. Nonetheless, due to the ample dissociation of the modified peptides, these reductions do not undermine the ability to sequence the peptides independent of the PTM.

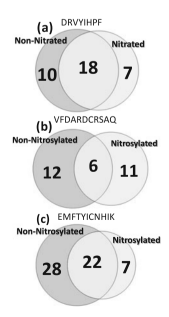


Figure 4. Venn diagrams comparing the number of fragmentation ion types produced after He MAD of the modified and unmodified peptides: (a) [DRVYIHPF+2H]²⁺, (b) [VFDARDCRSAQ+2H]²⁺, and (c) [EMFTYICNHIK+2H]²⁺

A direct comparison of MAD of the non-nitrated and nitrated species of the peptide GPLEYGFAKGPLAK was unattainable; however, we are still able to make some conclusions based on the MAD spectra of the nitrated form of this peptide species (Figure 1(d), (e), and (f)). Regardless of precursor charge state, extensive backbone cleavage is observed, including cleavage in and around the vicinity of the modification. When taking into account those fragments that undergo a neutral loss of the modification, all of the modified peptides studied contain at least two and as many as five backbone cleavage ion types directly adjacent to the modified residue. In contrast, fragmentation is inhibited near the site of nitration or S- nitrosylation in ECD and ETD [22, 31]. MAD fragmentation appears to not suffer the same inhibitions as ECD and ETD. There are two possible explanations for this contrasting behavior between MAD and ETD/ECD. The first explanation relates to the unique nature of metastable atom chemistry. Metastable atoms at the molecular level behave like positive ions and are known to be attracted to regions of high electron density or polar groups [50, 51]. In MAD, radicals can be directly generated anywhere along the peptide backbone where there is a polar group or lone pair of electrons, such as carbonyl oxygen atoms [33]. In ECD and ETD, radical migration must instead occur from original sites of protonation through long-range interactions with the carbonyl oxygen atoms along the backbone producing aminoketyl radicals. Whether the radical or hydrogen atoms are transferred first in ETD and ECD, these techniques are apparently more easily hindered by high electron- or hydrogen-affinity groups than MAD [30].

The second explanation for the difference between MAD and ETD/ECD fragmentation behavior may be due to high levels of secondary reactions in MAD. It was shown in our previous work that longer exposure times and more energetic He metastable atom exposure causes an increase in the relative abundance of low mass ions at the expense of high mass ions, most likely due to secondary reactions [33]. Although primary metastable atom reactions with the oxygen atom lone pairs of the -NO or -NO₂ modifications could result in radical sequestering without fragmentation, a second metastable atom interaction with the same peptide at a different location could induce the observed dissociation.

Figure 4 and Supplemental Table 2 compare the different amount and types of fragment ions produced for the modified and unmodified peptides DRVYIHPF, VFDARDCRSAQ, and EMFTYICNHIK after He MAD exposure. Even though, in Figure 3, we observe backbone cleavage adjacent to the modified residues, a greater abundance of these types of cleavages are observed with the unmodified peptides. This is especially noted in Figure 4(a) and (c), which have the greater reduction of overall cleavages. These two observations may suggest that additional sites of lone pairs of electrons, such as those found on the –NO or –NO₂ groups of the PTMs, in fact compete for the initial metastable atom interaction.

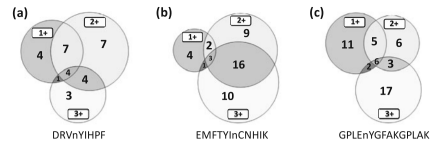


Figure 5. Venn diagrams comparing the number of fragment ion types produced by He MAD of the 1+, 2+, and 3+ charge states of the modified peptides: (a) DRVnYIHPF, (b) EMFTYInCNHIK, and (c) GPLEnYGFAKGPLAK

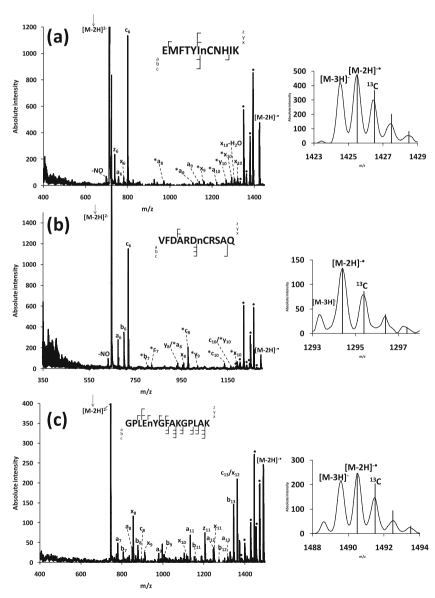


Figure 6. He MAD spectra of doubly-deprotonated S-nitrosylated: (a) EMFTYInCNHIK, (b) VFDARDnCRSAQ, and (c) GPLEnYGFAKGPLAK. The small inserts compare the actual and expected (vertical lines) isotopic distribution. The precursor is indicated by an arrow, neutral losses from the charge-reduced species are indicated by a dot, and the neutral loss of modification is indicated by an asterisk. Fragments that do not retain the modification are omitted from the peptides sequence inserts

One overall trend observed for the three peptides in Figure 3 is that MAD of the non-modified peptides produced a higher percentage of N–C $_{\alpha}$ backbone cleavages compared to the modified (nitrated and S-nitrosylated combined) peptides. The modified peptides typically produced higher percentages of C $_{\alpha}$ –C and C–N bond cleavages. A comprehensive analysis of a larger pool of peptides with varying types of modifications will need to be assessed before confirming trends in fragmentation behavior in MAD.

Effect of Charge State on Peptide Backbone Cleavage

The amount and type of backbone cleavages observed through MAD of the 1+, 2+, and 3+ charge states of the modified peptides, DRVnYIHPF, GPLEnYGFAKGPLAK, and EMFTYInCNHIK, and the 2+ and 3+ charge state of VFDARDnCRSAQ are compared in Figure 5 and Supplemental Table 3. The most striking observation is the vast difference in product ion types observed from the different precursor ion charge states. When comparing the three different charge states examined for a given peptide, only 13%, 6%, and 16% (Figure 5(a), (b), and (c)), respectively) of the total product ions observed are common to all charge states. This leads us to infer that the types of fragment ions produced by MAD are highly dependent on the charge state. There are at least three reasons for the differences in fragmentation types observed for MAD of the same peptide in different charge states. (1) A difference in mechanism, e.g., electron capture versus Penning ionization; (2) a difference in degree, e.g., electron capture of the 3+ precursor is more exothermic than electron capture of the 2+ precursor; or (3) a difference in the conformation of the peptide prior to interaction with the metastable atoms.

There is no observable trend between the precursor ion charge state and the product ion types observed in Figure 5. For example, MAD of the 3+ species of angiotensin II and the 1+ species of EMFTYInCNHIK produced the fewest fragment ion types of the three charge states observed for the specific peptide (Figure 5(a) and (b)), respectively). However, it was the 2+ precursor species that produced the fewest fragment types for the peptide GPLEnYGFAKG-PLAK (Figure 5(c)). For the three peptides studied, the 3+ precursor ions tend to produce more c-/z-type ions, whereas the 1+ precursors tend to produce more a-/x-type ions. The 2+ precursor ions typically produce more b-/y-type ions. These trends have not been validated against a larger database of precursor ions.

S-Nitrosylated and Nitrated Peptide Anions

Figure 6 shows the He MAD spectra of three peptides with a 2- charge state; two contain a S-nitrosylated modification (Figure 6(a) and (b)) and one contains a nitrated modification (Figure 6(c)). In all cases, the most abundant dissociation channels are neutral losses from the charge-reduced

species including H₂O (18 Da), CO₂ (44 Da), and either NO or NO₂ (30 or 46 Da, respectively). All the neutral losses observed from the charge-reduced species for the three peptides are listed in Table 1 and are indicated with a dot (•) within the spectra in Figure 6. Water and carbon dioxide groups are common neutral losses observed in other electron-induced dissociation methods on peptide anions, such as electron detachment dissociation (EDD) [52, 53] and negative electron transfer dissociation (nETD) [26, 54]. When dissociating protonated ions with MAD, the neutral loss of a hydrogen radical from the charge-reduced species is usually more abundant than the charge-reduced product ion. Deprotonated peptides also show signs of significant hydrogen radical loss, although not quite as abundant as when fragmenting cationic precursor ions. When comparing the ¹³ C envelope to the expected envelope (Figure 6(a-c)), neutral losses of hydrogen radicals are still apparent. Figure 6(a) and (b) show the He MAD spectra of two S-nitrosylated peptides, both containing a modification on the 7th residue. A variety of ion types are observed for each peptide, with a majority of a-/x-type ions for the peptide in Figure 6(a) and a majority of c- and y-type ions in Figure 6(b). In both cases, 70% of information required for sequencing is obtained through the production of 10 (Figure 6(a)) and 13 (Figure 6 (b)) distinct fragment ions, including a modification retainment percentage of 50% and 43%, respectively. In these two cases, the modification site can be unambiguously identified.

Figure 6(c) shows the MAD spectrum observed for the 2–precursors of the nitrated peptide GPLEnYGFAKGPLAK. Similar to the peptide in Figure 6(a,) a-, b-, and x-type ions are most common. Because of the more stable nature of the nitration modification for the peptide in 6c, all fragment ions retain the modification and enable PTM site determination. Furthermore, no observable trend can be noted based on ion

Table 1. List of the neutral losses observed from the charge-reduced species, $[M-2H]^{-}$, after He MAD of the 2– peptide ions

$[M-2H]^{2-}$ peptide ion	Neutral loss (Da)	Proposed loss
EMFTYInCNHIK	-30	-NO
	-45	-CO ₂ , -H
	-62	$-CO_2$, $-H_2O$
	-75	-NO, -CO ₂ , -H
	-92	$-NO, -CO_2, -H_2O$
	-107	-SNO, -CO ₂ , -H
VFDARDnCRSAQ	-18	$-H_2O$
	-30	-NO
	-45	−CO ₂ , −H
	-49	$-NO, -H_2O, -H$
	-62	$-CO_2$, $-H_2O$
	-75	-NO, -CO₂, −H
	-92	$-NO, -CO_2, -H_2O$
	-107	-SNO, -CO ₂ , -H
GPLEnYGFAKGPLAK	-18	$-H_2O$
	-36	$-H_2O$, $-H_2O$
	-45	−CO ₂ , −H
	-62	$-CO_2$, $-H_2O$
	-90	$-CO_2$, $-NO_2$
	-108	-CO ₂ , -NO ₂ , -H ₂ O

The neutral losses are indicated with a dot in the corresponding spectra

types produced, in contrast to EDD and nETD, which preferentially produce a-/x-type ions. That the fragmentation of EMFTYInCNHIK and VFDARDnCRSAQ appear to be directly adjacent to the highly electronegative nitrosylated cysteine residues supports the proposed mechanism of energy transfer as Penning ionization [33–35]. The site of preferential fragmentation seems to correlate to the PTM site rather than to the sites of deprotonation.

Conclusion

Extensive backbone cleavage of both nitrated and S-nitrosylated peptides were obtained after exposure to a high kinetic energy beam of He metastable atoms. MAD is less sensitive than ECD and ETD to the presence of nitrotyrosine, which can act as an electron predator and inhibit fragmentation in the latter two techniques. Complete peptide sequencing and PTM location is obtained for the majority of the peptides studied regardless of charge state or ion polarity. Amino acid side-chain cleavages enable L/I differentiation in several cases. Peptides containing nitrosylated cysteine residues prove to be highly labile, thereby making isolation prior to tandem analysis more difficult in general. However, we demonstrate that by combining multiple spectra of different charge states, accurate PTM site location can be achieved using MAD. Because of the unique fragmentation pathways offered through metastable atom chemistry, MAD is a complementary alternative to collisional or other electron-induced fragmentation methods.

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