
Tandem Mass Spectrometry Investigation of ADP-ribosylated Kemptide

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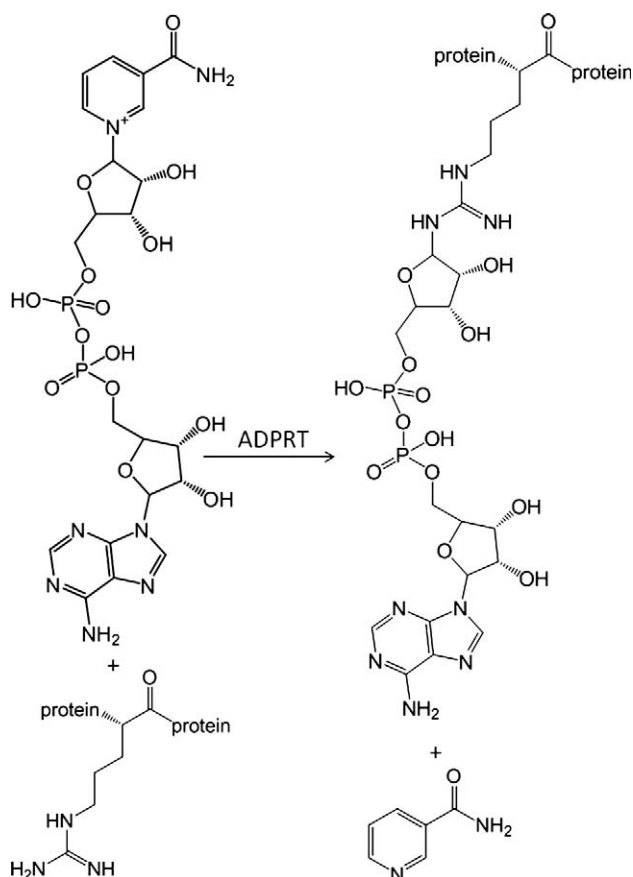
Bacterial adenosine diphosphate-ribosyltransferases (ADPRTs) are toxins that play a significant role in pathogenicity by inactivating host proteins through covalent addition of ADP-ribose. In this study we used ADP-ribosylated Kemptide (LRRASLG) as a standard to examine the effectiveness of three common tandem mass spectrometry fragmentation methods for assignment of amino acid sequence and site of modification. Fragmentation mechanisms investigated include low-energy collision-induced dissociation (CID), infrared multiphoton dissociation (IRMPD), and electron-capture dissociation (ECD); all were performed on a hybrid linear ion trap Fourier transform ion cyclotron resonance mass spectrometer. We show that ECD, but neither CID nor IRMPD, of ADP-ribosylated Kemptide produces tandem mass spectra that are interpretable with regard to amino acid sequence assignment and site of modification. Examination of CID and IRMPD tandem mass spectra of ADP-ribosylated Kemptide revealed that fragmentation was primarily focused to the ADP-ribose region, generating several potential diagnostic ions for use in discovery of ADP-ribosylated proteins. Because of the lower relative sensitivity of ECD during data-dependent acquisition to CID, we suggest a 2-fold strategy where CID and IRMPD are first used to detect ADP-ribosylated peptides, followed by sequence assignment and location of modification by ECD analysis. (J Am Soc Mass Spectrom 2009, 20, 477–483) © 2009 Published by Elsevier Inc. on behalf of American Society for Mass Spectrometry

Adenosine diphosphate-ribosyltransferases (ADPRTs) catalyze the transfer of ADP-ribose from β -NAD⁺ to target proteins (Scheme 1), commonly at arginine residues [1]. Members of the ADPRT family modify a diverse set of target proteins and share little sequence homology. The mammalian ADP-ribosyltransferase ART-1 modifies defensin-1, which ultimately blocks the antimicrobial and cytotoxic effects of the defensin HNP-1 [2]. Many ADPRTs found in bacteria have been implicated in pathogenicity through the inactivation of host proteins (e.g., *Vibrio cholera*, *Bordetella pertussis*, and *Streptococcus pyogenes* [1]). *S. pyogenes* produces the ADPRT toxin SpyA, which modifies vimentin, actin, and tropomyosin, all proteins involved in cytoskeletal structure [3]. ADP-ribosylation of these proteins prevents polymerization of the cytoskeleton, leading to collapse of intermediate filaments. Other ADPRT toxins are known to target proteins involved in signal transduction and regulatory functions [1]. Although there are numerous molecular assays demonstrating that ADP-ribosylation is an important step in pathogenicity, to date the exact site of ADP-ribosylation and number of modification targets remains unknown for many ADPRTs. A high-throughput

mass spectrometry assay designed for the identification of target proteins as well as sites of ADP-ribosylation would provide insight into mechanisms of infection.

Often liquid chromatography-tandem mass spectrometry (LC-MS/MS) methods are used to identify sites and protein targets of post-translational modifications (PTMs) in complex biological samples [4, 5]. Collision-induced dissociation (CID) is typically used as the main fragmentation method during LC-MS/MS analyses because the instrumental duty cycle is high and acquired peptide tandem mass spectra may be easily sequenced. This primarily results from the predictable peptide fragmentation pathways [6], allowing for identification of the parent proteins [7]. However, the presence of a PTM on a peptide can redirect CID fragmentation patterns such that sequence may not be assigned. Previously, Margarit et al. [8] demonstrated that CID of an ADP-ribosylated peptide did not result in “typical” peptide fragmentation, interfering with facile sequence interpretation. Additionally, CID analysis of SpyA ADP-ribosylated Vimentin [3] verified similar fragmentation trends, preventing modified peptide sequence assignment (S. M. Hengel, unpublished data). The inability to directly assign peptide sequences of ADP-ribosylated peptides using CID led us to undertake a detailed mass spectrometry fragmentation pattern analysis of a standard, ADP-ribosylated Kemptide (LRRASLG). Three common fragmentation meth-

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Scheme 1. ADPRTs catalyze the transfer of ADP-ribose to acceptor proteins from NAD⁺.

ods were investigated: (1) CID, (2) electron-capture dissociation (ECD), and (3) infrared multiphoton dissociation (IRMPD).

The goals of this study were to determine the most efficient technique for producing reliable and predictable fragment-ion spectra for the identification of ADP-ribosylated peptides and to determine whether diagnostic ions exist that might be used in future studies to identify new ADPRT targets. ADP-ribosylated Kemptide was chosen as a standard for examining which of the three methods—CID, ECD, and IRMPD—were best suited to (1) assign amino acid sequence and (2) produce ions diagnostic of ADP-ribosylation. Here we discuss the results of this comparison and how these methods may be used alone and in combination to identify ADP-ribosylated proteins in complex mixtures.

Experimental

ADP-Ribosylation of Peptide Standard

Reagents used for peptide modification were obtained from Sigma (St. Louis, MO, USA). A 5 mM solution of Kemptide (Leu-Arg-Arg-Ala-Ser-Leu-Gly, MW 771.4715) was modified as previously described [9], in the presence of 10 mM ADP-ribose, 20 mM DTE, and 100 mM phosphate buffer at 33 °C for 4 h. After incubation, the

extent of ADP-ribosylation was determined by comparing peak height and area of extracted ion chromatograms for modified and unmodified Kemptide precursor masses following separation on a C₁₈ column described in the following text.

Electrospray Ionization LTQ-FT-based Mass Spectrometry

Peptides were analyzed using electrospray ionization (ESI) in a linear ion trap Fourier transform ion cyclotron resonance cell mass spectrometer (LTQ-FT-ICR) (Thermo Electron Corp., San Jose, CA, USA). HPLC columns were packed in-house (0.75 μm i.d. × 11 cm; 100-Å Magic C18AQ; Michrom Bioresources, Auburn, CA, USA) and separations were performed with an inline Agilent 1100 binary HPLC pump (Agilent Technologies, Palo Alto, CA, USA) using a linear gradient of 5 to 35% ACN over 1 h. ESI voltage was applied with a liquid junction before the analytical column with a gold wire and a micro-tee junction [10]. All ions were measured in positive-ion mode, both precursor and fragment masses were detected in the FT mass analyzer. Source parameters were set as follows: ESI voltage, 2.5 kV; capillary temperature, 265 °C; capillary voltage, 13 V; tube lens, 35 V. For all fragmentation methods the resolution was set to 5.0e⁴ and 2.0e⁴ for MS and MS/MS, respectively. Automatic gain control was used to maintain constant ion populations at 5e⁵ and 2e⁵ for MS and MS/MS, respectively.

Fragmentation Methods

Reported tandem mass spectra for both modified and unmodified Kemptide are from doubly charged precursor masses 657.27 and 386.74 *m/z*, respectively; tandem mass spectra were acquired in the FT-ICR cell during LC-MS/MS analysis. Fragment ion spectra acquired with CID were produced using 35% collision energy and a 2.0 Da isolation window. ECD tandem mass spectra were acquired using 5 eV fragmentation energy for 70 ms with a delay of 0.3 ms and a 3.0 Da isolation window. IRMPD was performed by irradiating precursor ions with photons produced by a CO₂ laser [Synrad firestar series V20, model FSV20SFB; 75 Watts (10.2–10.8 μm)] with a pulse duration of 50 ms, a power of 60%, and an isolation window of 5.0 Da.

Limit of Detection

To determine the limit of detection of ADP-ribosylated Kemptide in a complex mixture, a bovine serum albumin (BSA) peptide standard was used to dilute the analyte (Michrom Bioresources). BSA was maintained at a concentration of 5 pmol/μL and serial dilutions of the analyte were initiated with a 1:1 ratio of BSA to ADP-ribosylated Kemptide. ADP-ribosylated Kemptide concentrations ranged from 2.5 pmol/μL to 250 amol/

μL in 5-fold serial dilution increments (i.e., 1:1; 1:10; 1:50; 1:100...). Samples were analyzed using data-dependent LC-MS/MS conditions described earlier, selecting the five most intense peaks observed in the MS spectrum for MS/MS, acquiring both CID and IRMPD data for each dilution.

Results and Discussion

Collision Induced Dissociation (CID)

The process of conducting CID combined with automated, data-dependent ion selection provides a fast duty cycle that is well suited for LC-based analysis of mixtures and is therefore commonly used in proteomics [7]. However, the CID-generated tandem mass spectrum of ADP-ribosylated Kemptide contains none of the observed *b* or *y* ions in the unmodified Kemptide CID tandem mass spectrum (Figure 1a) making sequence and site of ADP-ribosylation assignment impossible (Figure 1b). Close examination of the ADP-ribosylated Kemptide CID tandem mass spectrum revealed several ions not accountable by Kemptide peptide backbone fragmentation (Figure 1b). Based on accurate mass measurements, the known structure of ADP-ribose, and Kemptide amino acid sequence, most of these ions appear to be formed through fragmentation of the ADP-ribose region.

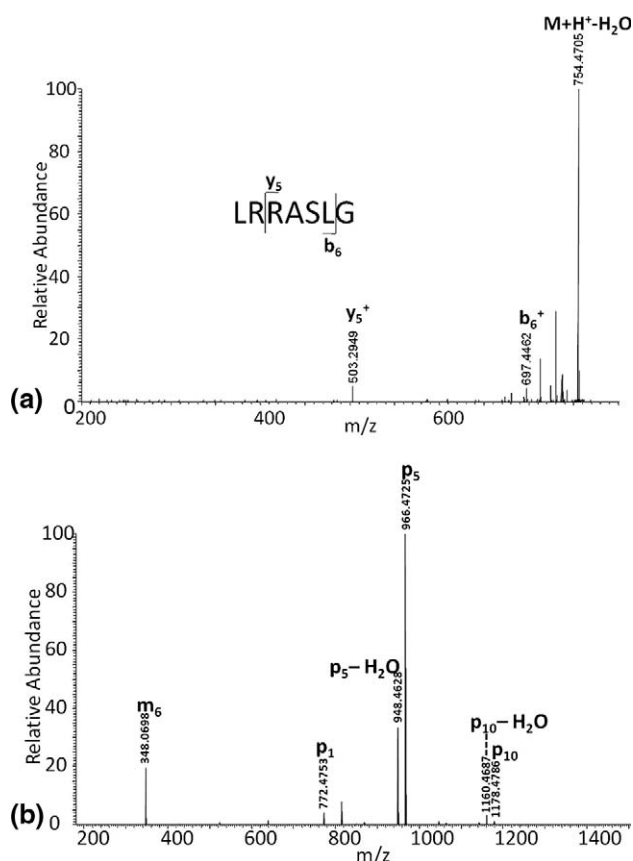
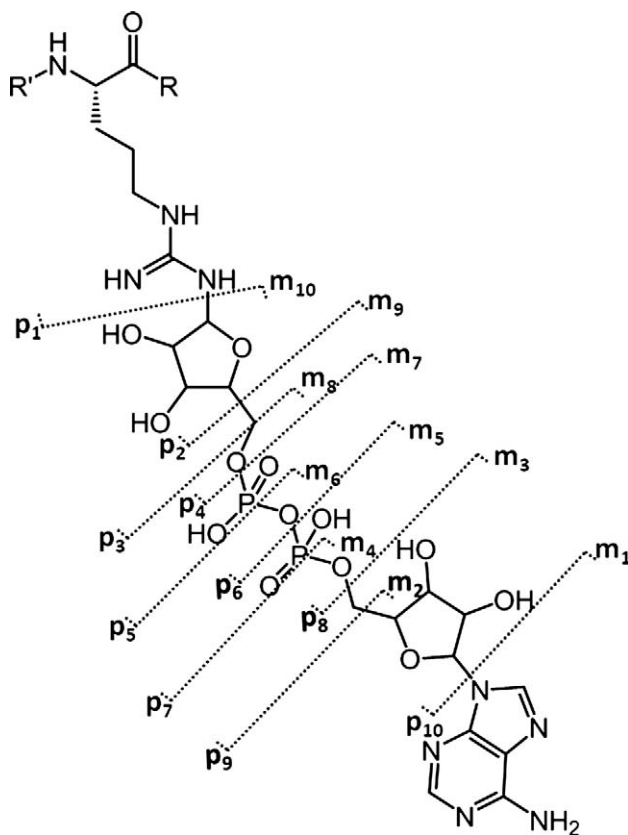


Figure 1. Collision-induced dissociation tandem mass spectra of (a) Kemptide and (b) ADP-ribosylated Kemptide.



Scheme 2. ADP-ribose fragmentation scheme: fragment ions containing the intact peptide backbone plus a portion of the modification are labeled p_1 through p_{10} . Fragment ions independent of the peptide sequence, only containing the ADP-ribose modification, are labeled m_1 through m_{10} .

ADP-ribose contains ten single bonds, excluding cross-ring fragmentation, which can yield fragment ions. To keep track of fragment ions occurring on the ADP-ribose backbone, we suggest the nomenclature shown in Scheme 2. Fragment ions with charge retained on the ADP-ribose group, but with loss of the peptide, are labeled $m_1 \dots m_{10}$ for modification. Fragment ions that contain part of the ADP-ribose region plus the intact peptide backbone are labeled $p_1 \dots p_{10}$. We do not show any cross-ring fragmentation sites labeled in Scheme 2 because none was observed in our studies. However, we realize that such fragments might be formed by some future fragmentation method or combination of current methods. For these potential fragment ions we propose the use of a variation on the carbohydrate fragmentation nomenclature proposed by Domon and Costello [11]. In place of the subscript number for single bond fragments, rings are assigned a Roman numeral (i.e., I for the ribose closest to the peptide, then II, etc.), and cross-ring fragments are noted using numbers indicating which cross-ring bonds are broken. Similar to the single bond fragments along ADP-ribose, *m* and *p* will be used to designate on which side the charge is observed. For example, a fragmentation event at the ribose ring, adjacent to the guanine

functional group, between bonds zero and two would be designated as $^{0,2}p_1$ or $^{0,2}m_1$, depending on which side the charge is observed. The ADP-ribosylated Kemptide CID tandem mass spectrum contains an observed ion at m/z 348, labeled m_6 (Scheme 2), which results from fragmentation of the APP-ribose pyrophosphate bond. This fragmentation generates two types of ions explained with structures where charge is retained on both sides of the cleavage site. The measured mass difference for the ion labeled m_6 and that of the putative structural assignment is calculated to be 1.7 ppm, strongly suggesting the assignment is correct. We also note that previously reported results of CID ADP-ribosylated peptides [8] yielded similar fragment ions localized between ribose and the phosphate group proximal to the peptide backbone, again suggesting that the ADP-ribose modification “directs” the CID fragmentation process of ADP-ribosylated peptides.

Further examination into the origin of other prominent ions in Figure 1b (i.e., the base peak) suggests that such ions are complementary fragments to the ion at 348 m/z and an additional fragment near the adenine portion of the ADP-ribose group (fragments labeled p_5 , p_{10} ; Scheme 2). Unlike the ion at 348 m/z , which appears to be charged fragments of ADP-ribose without peptide attached, ions p_5 and p_{10} contain a mass equal to that of Kemptide plus a residual mass from ADP-ribose fragmentation. Charge states of this type of fragment ion are equivalent to the initial charge state minus the loss of the protonated ADP-ribose fragment. Because it is not a neutral loss, often seen in CID with other post-translational modifications like phosphorylation [5], the charge state of the observed fragment ions may vary. Therefore, we predict that CID spectral complexity of ADP-ribose modified peptides will increase directly with amino acid number and charge state of the precursor peptide.

Although the CID tandem mass spectrum of ADP-ribosylated Kemptide contains ions that may be diagnostic of ADP-ribose, the spectrum does not contain any information on peptide sequence, as seen in the associated CID tandem mass spectrum of Kemptide (Figure 1a). In an attempt to understand what single tandem mass spectrometry method, or combination of methods, provides full interpretation of peptide sequence, site of modification, and generates potential diagnostic ions, we investigated the fragmentation of Kemptide and ADP-ribosylated Kemptide by ECD and IRMPD.

Electron Capture Dissociation (ECD)

Compared with the CID tandem mass spectrum (Figure 1a), a more complete ion series is observed with ECD fragmentation on the unmodified Kemptide standard (Figure 2a). During ECD one charge on the peptide is neutralized due to the capture of an electron. As a result, peptides that are initially doubly charged retain a charge on only one of the fragment ion products; thus

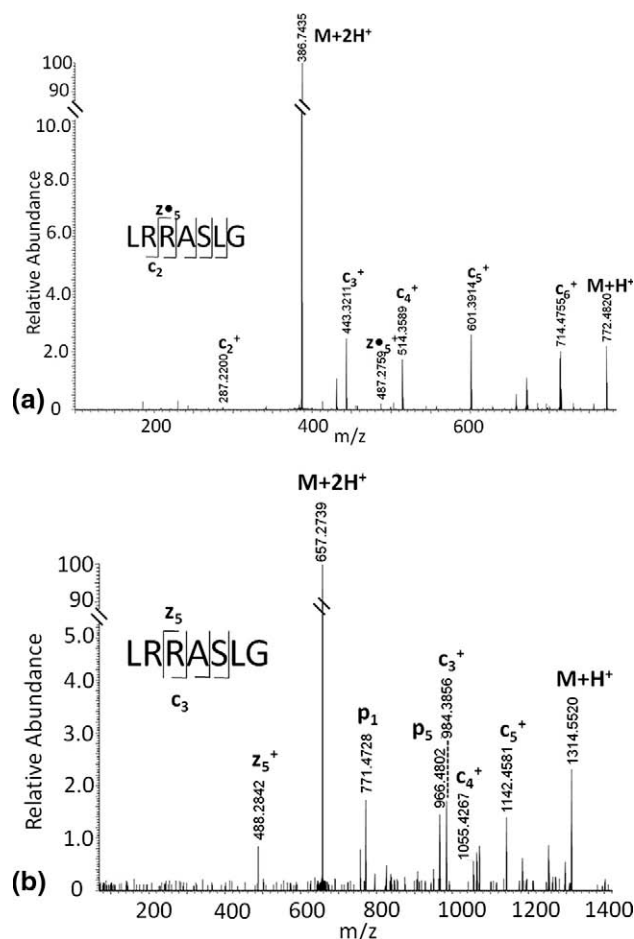


Figure 2. Electron capture dissociation tandem mass spectra of (a) Kemptide and (b) ADP-ribosylated Kemptide.

both complete c or z ion series may not be detected (Figure 2a). Therefore, larger peptides with higher charge states should have a greater probability to produce both c and z ions upon ECD fragmentation, increasing the opportunity for assigning post-translational modification sites.

A distinguishing feature of ECD is the retention of peptide side-chain modifications [12–14]. Predictably, we see in our ECD data of ADP-ribosylated Kemptide (Figure 2b) that ADP-ribose is retained on the arginine side chain. Additionally, the ECD tandem mass spectrum of ADP-ribosylated Kemptide contains three of six predicted c ions and one of six z peptide fragment ions within 10 ppm mass accuracy of the calculated theoretical values. As a result, peptide sequence and the site of ADP-ribosylation can be elucidated from the ECD tandem mass spectrum. The Kemptide amino acid sequence contains two arginine residues and, using ECD, we were able to localize the site of modification to the second amino acid, i.e., LR^{*}RASLG, where R^{*} = arginine ADP-ribosylated on the guanidinium amino group. In contrast to the CID tandem mass spectrum, only two observed ions of lower intensity correlate to fragmentation of the ADP-ribose group in the ECD tandem mass spectrum (fragment p_1 and p_5 , Figure 2b).

This indicates that, unlike CID of ADP-ribosylated Kemptide, a minimal amount of the ECD energy is directed to ADP ribose and the majority of energy is directed to fragmentation of the peptide N—C_α amide backbone in a predictable manner. Thus, ECD is a reliable fragmentation method that can provide information for peptide sequencing and site assignment of ADP-ribosylated Kemptide.

Infrared Multiphoton Dissociation (IRMPD)

As expected, the IRMPD fragment ion spectrum of the unmodified Kemptide peptide standard contains *b* and *y* ions, as well as a diagnostic immonium ion (Figure 3a). Despite the fact that both CID and IRMPD generate fragment ions through an increase in vibrational energy, there is a noticeable difference between the unmodified Kemptide CID and IRMPD tandem mass spectra. One likely source of this variation is the time-of-flight effect observed for CID fragment ions generated in the LTQ and subsequently transmitted to the ICR cell, as opposed to IRMPD fragment ions that are both generated and measured in the ICR cell [15]. Compared to the unmodified Kemptide ECD tandem mass spectrum, the observed IRMPD ion series is less complete—peptide sequencing of this short peptide would be challenging. Implementation of IRMPD frag-

mentation on ADP-ribosylated Kemptide produced a tandem mass spectrum with only one fragment ion, *y*₅, corresponding to peptide backbone fragmentation (Figure 3b). Upon further analysis, the IRMPD tandem mass spectrum base peak, 136 *m/z*, is not a result of peptide backbone fragmentation, but instead is the complement of fragment ion *p*₁₀ (Scheme 2) found in the CID spectrum. This ion, *m*₁, appears to result from the loss of adenine from ADP-ribose, is independent of peptide sequence, and confirmed again using <1 ppm mass error.

Distinct from the ADP-ribosylated Kemptide ECD spectrum and the unmodified Kemptide IRMPD spectrum, the precursor ion is no longer observed when using IRMPD to fragment ADP-ribosylated Kemptide. This may be due to the ease in which the ADP-ribose group fragments, as seen in the CID tandem mass spectrum, and an increase of photon absorptivity [16], making the fragmentation more efficient for an ADP-ribose group when compared with a peptide backbone.

Potential Diagnostic Ions of ADP-ribosylated Peptides

Both IRMPD and CID fragmentation techniques generated ions independent of the Kemptide amino acid sequence, i.e., 136 and 348 *m/z*, respectively, yet specific to ADP-ribose. One of these ions, 348 *m/z*, was previously observed using CID to study an ADP-ribosylated actin peptide [8]. In addition, CID tandem mass spectrometry generated fragment ions that are indicative of an ADP-ribose modification and dependent on peptide sequence. Ions such as these are potentially diagnostic of ADP-ribose modifications on a peptide and may be useful in discovery-based proteomic screens that seek to identify targets of ADPRTs. However, because neither CID nor IRMPD produced fragment ions that lead to peptide sequence assignment, precursor ions identified containing ADP-ribose in an initial CID or IRMPD screening analysis would need to be further subjected to ECD fragmentation. There are two primary benefits of using CID over ECD or IRMPD in the initial discovery of ADP-ribosylated peptide precursor ion masses: (1) ions that are dependent on and independent of peptide precursor ion mass to charge ratios are produced, providing multiple fragment ions indicative of ADP-ribosylated peptides, thus reducing the false discovery rate; and (2) duty cycle is high, when acquired in an LTQ, relative to IRMPD and ECD, providing better analytical sampling for detection of ADP-ribosylated peptides. However, in this study all fragment ions were measured in the FT-ICR cell to generate high confidence structure assignments. When analyzing samples with a large dynamic range where sensitivity is crucial to detection, IRMPD should provide two advantages: (1) the IRMPD-produced diagnostic ion at 136 *m/z* is the base peak of the tandem mass spectrum and (2) ions are

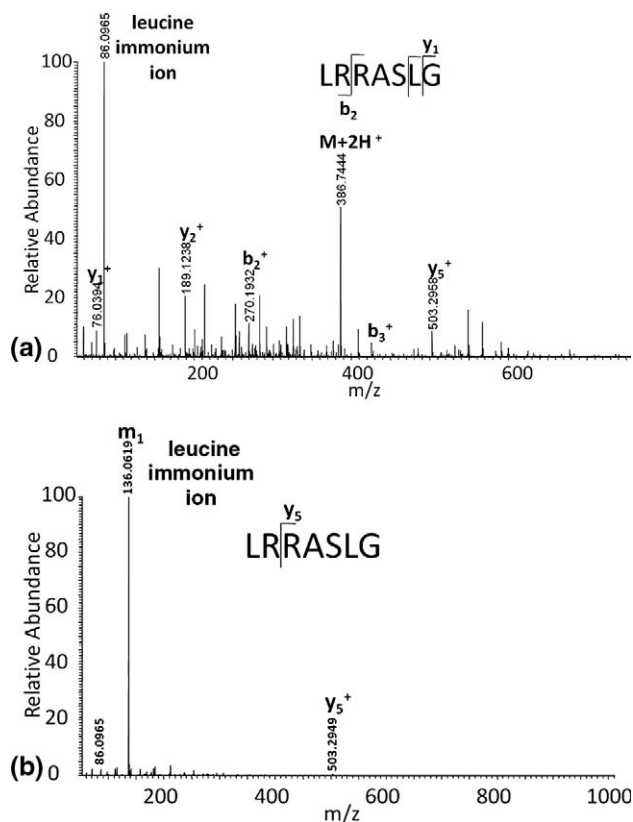


Figure 3. Infrared multiphoton dissociation tandem mass spectra of (a) Kemptide and (b) ADP-ribosylated Kemptide.

measured at high mass accuracy in the ICR cell. Both CID and IRMPD have unique benefits when searching for diagnostic ions in tandem mass spectra. Provided sample availability and sufficient instrument access, we suggest performing both CID and IRMPD LC-MS/MS experiments measuring fragment masses in the ICR cell generating high mass accuracy data. If CID fragments were generated and measured in the LTQ while IRMPD fragmentation occurs in the FT-ICR, then both data could be acquired in a single run with minimal reduction in duty cycle. However, compared with measuring fragment ions in the ICR cell, the loss of mass accuracy for CID fragment ions acquired in the LTQ would be substantial. Given that no neutral loss trigger is possible with the observed type of fragment ions, a discovery-based screen designed to detect ADP-ribosylated peptides would need to be conducted in two LC-MS/MS experiments. First, a typical data-dependent experiment would be performed using CID, and/or IRMPD and, second, an ECD experiment using an inclusion list based on the first experiment would be completed. When MS-driver software advances to allow on-the-fly switching from CID/IRMPD to ECD with this type of diagnostic ion, then the discovery and identification of ADP-ribosylated peptides could be conducted in a single LC-MS/MS experiment.

Testing Diagnostic Ions

Spiking ADP-ribosylated Kemptide into a complex mixture simulated an experiment designed to discover new targets of ADPRTs. These analyses also allowed us to test the proposed diagnostic ions and establish a limit of detection. Three proposed ions diagnostic of ADP-ribosylated peptides were investigated as marker ions for the purpose of rapid identification of unknown ADPRTs targets. These ions were selected based on intensity and observed frequency from MS/MS analyses of ADP-ribosylated Kemptide. Two of the ions, m_6 and p_5 (Scheme 2), are observed in CID tandem mass spectra and the third, m_1 , is observed when using IRMPD fragmentation.

Varying amounts of ADP-ribosylated Kemptide were spiked into a BSA protein digest (see Limit of Detection) for two reasons: (1) to determine a limit of detection and (2) to assess whether m_1 , m_6 , and p_5 fragment ions could be used to accurately and reproducibly detect the tandem mass spectra of ADP-ribosylated Kemptide in a mixture of non-ribosylated peptides from BSA. Data were acquired via a data-dependent ion selection scheme that selected the top five ions per precursor ion scan for tandem mass spectrometry (i.e., CID or IRMPD). A limit of detection was determined for both precursor ions and all three potential diagnostic fragment ions. In all cases, the limit of detection was determined to be femtomoles of ADP-ribosylated Kemptide in the presence of picomoles of the protein digest at a signal-to-noise ratio of 2:1.

Tandem mass spectra from the spiking experiments were sorted using PERL scripts that identified .dta files containing each of the three specified marker ions within 10 ppm mass accuracy. Precursor ion masses of the .dta files containing any one of these three diagnostic ions were then compared to the known ADP-ribosylated Kemptide mass. All observed precursor masses from the three PERL script lists that were outside a mass tolerance of 10 ppm were considered false positives. Using this measure, initial false-positive rates of each ion independent of the others were high: m_6 95% (CID), p_5 66% (CID), and m_1 92% (IRMPD). This is likely a reflection of the commonality of these ions in many tandem mass spectra. However, when precursor masses identified from .dta files selected by the three PERL scripts were used in combination rather than independently, the resulting overall false-positive rate was reduced to 0%. This same type of calculation changed the false-negative rate from 0 to 8%. This suggests that when trying to blindly screen tandem mass spectral datasets for .dta files containing ions diagnostic of ADP-ribose that one should include all three diagnostic ions to maintain a low false-positive rate. Using fragment ions diagnostic of ADP-ribose from both CID and IRMPD data, we were able to successfully identify in an automated manner (subsequently confirmed manually) .dta files of ADP-ribosylated Kemptide spiked into a digest of BSA. The same approach should be useful for discovery work in more complex whole cell lysates, although this may result in a higher false-positive rate.

Conclusions

Examination of three different fragmentation techniques for the purpose of understanding ADP-ribosylated Kemptide fragmentation patterns yielded tandem mass spectra with both unique and shared fragment ions. Although CID and IRMPD produced ions diagnostic of ADP-ribose useful for determining whether a parent mass contains ADP-ribose, the tandem mass spectra lack significant peptide backbone fragmentation, thus preventing identification of peptide sequence and modification site. Because ECD has been recognized as a technique that provides peptide backbone fragmentation while allowing for retention of post-translational modifications [13, 14], we investigated its utility for sequencing ADP-ribosylated peptides. Our data clearly demonstrated that ECD is the best fragmentation method for determining peptide sequence and the site of modification of ADP-ribosylated peptides. Electron-transfer dissociation (ETD) [17], a fragmentation method similar to ECD, may also provide sequence information for an ADP-ribosylated peptide, but has not been examined in this study. Disadvantages of ECD, however, include a longer duty cycle and low fragmentation efficiency relative to CID. Because of these drawbacks, we recommend first using CID and IRMPD fragmentation techniques to discover ADP-ribosylated peptides with diagnostic ions, followed by

targeted ECD analyses to decipher the location of the modification and the sequence of the peptide. This two-step assay will allow for a more thorough analysis when attempting to identify ADP-ribosylated targets compared to using only shotgun proteomic analysis with ECD alone.

Acknowledgments

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