

RESEARCH ARTICLE

Mapping the tandem mass spectrometric characteristics of citrulline-containing peptides

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Rationale: Protein citrullination (deimination) is a post-translational modification of proteins converting arginine(s) into citrulline(s). “Overcitrullination” could be associated with severe pathological conditions. Mass spectrometric analysis of modified proteins is hindered by several problems. A comprehensive study of the fragmentation of deiminated peptides is not yet available. In this paper we have made an attempt to describe the characteristics of these processes, based on the studies of epitope model oligopeptides derived from clinically relevant proteins.

Methods: Solutions of purified model peptides containing either one or two citrulline residues as well as their native variants were injected directly into the electrospray source of a high accuracy and resolution quadrupole-time-of-flight instrument and were analysed by tandem mass spectrometry using low-energy collision-induced dissociation.

Results: Loss of isocyanic acid from citrulline residues is a preferred fragmentation route for deiminated peptides, which yields ornithine residues in the sequence. However, simultaneous detection of both the isocyanic acid loss and sequence fragments is often compromised. A preferential cleavage site was observed between citrulline and any other following amino acids yielding intensive complementary b- and y-type ions. Also, citrulline positioned at the C-termini displays a preferential cleavage N-terminal to this residue yielding characteristic γ_1 ions. These phenomena are described here for the first time and are referred to as the “citrulline effect”.

Conclusions: We found that the citrulline effect is very pronounced and could be used as a complementary tool for the confirmation of modification sites in addition to losses of isocyanic acids from the protonated molecules or from fragment ions. Low collision energy applied to peptide ions having partially mobile protons reveals the site of modification by generating specific and intensive fragments of the sequence. On the other hand, fragmenting precursor ions with mobile protons usually allows full sequence coverage, although citrulline-specific fragments may exhibit lower intensities compared to other fragments.

1 | INTRODUCTION

Protein citrullination (deimination) is a post-translational modification of proteins through which arginine residues (Arg, R) in certain sequentially well-defined positions are converted into citrulline

(Cit, X)¹ (Figure 1). Citrulline is a genetically non-coded amino acid that has a neutral side chain under physiological conditions. Deimination is catalysed by the peptidyl-arginine deiminase (PAD) enzymes and results in a monoisotopic mass increment of 0.9840 Da, isobaric with deamidation of asparagine and glutamine residues.

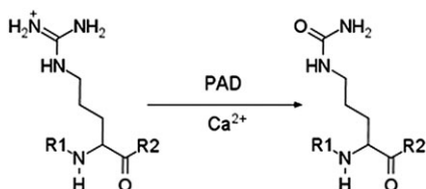


FIGURE 1 Protein deimination (citrullination) by the peptidyl arginine deiminase (PAD) enzymes altering physico chemical properties of the substrates

“Overcitrullination” is associated with several pathological conditions such as autoimmune diseases,² numerous neurodegenerative diseases,^{3–5} and even tumour growth.⁶ Nevertheless, only immunological (functional) or mass spectrometric (structural) methods are sensitive and specific enough for the detection of the modified species in complex biological samples due to the low abundance of deiminated proteins.⁷ Nowadays, immunodetection is used as a routine method to verify the presence of this post-translational modification (PTM) in proteins, e.g. for rheumatoid arthritis⁸ however, this approach is unable to determine the modification site. Locating the site of deimination is only possible with high resolution and high mass accuracy tandem mass spectrometry which lacks high specificity and therefore still needs improvement.⁹ In the last few years, several methods have been developed to improve the detection of deiminated polypeptides, including a specific and sensitive reaction with the modified residues¹⁰ or the enrichment of citrullinated peptides on beads with phenylglyoxal derivatives.^{11,12} Detection of modified proteins is a key step for the diagnosis, monitoring and staging of diseases associated with faulty deimination.

A selective loss of isocyanic acid (HNCO) has been reported for a few citrulline-containing model peptides using collision-induced dissociation (CID)¹³ (Figure 2A). Creese et al. utilised the loss of HNCO for selecting deiminated peptides in a complex sample and

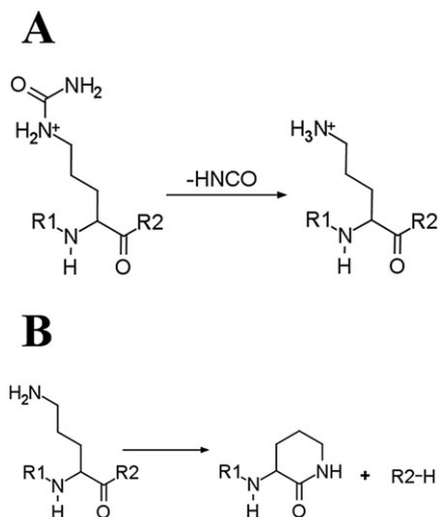


FIGURE 2 Proposed mechanism for citrulline effect leading to facile cleavage C-terminal to citrulline residues for protonated peptides in the gas phase. First step: loss of isocyanic acid (HNCO) characteristic to citrulline residues, leading to an ornithine residue (A).⁸ Second step: cleavage of the Orn-Aaa bond (B)¹²

further fragmenting them with electron transfer dissociation (ETD) to obtain a good sequence coverage.¹⁴

However, the behaviour of citrulline-containing peptides upon CID has not been a target for systematic investigation so far. Our study is therefore focused on unravelling the major characteristics of fragmentation of citrullinated peptides. To obtain clinically relevant results we selected protein epitopes which play role in rheumatoid arthritis (RA).^{15–18} RA is a systemic autoimmune disease in which citrullinated autoantigens are generated and are recognised by antibodies, resulting in a chronic inflammatory condition, destroying joints and causing severe pain to the patients.¹⁹ For these studies several oligopeptide variants were synthesised and analysed to gain insight into the major characteristics of CID fragmentation of deiminated peptides.

2 | EXPERIMENTAL

2.1 | Materials

Formic acid, distilled water and acetonitrile for MS analyses were from VWR (Radnor, PA, USA). Leu-enkephalin was used as a reference material for accurate mass measurements.

2.2 | Methods

Model peptides GER, TRGRS, VERHQS, GPRVVER and GYRARPAK were selected from the epitope regions of fibrin, filaggrin and collagen, which are substrates of PADs and are involved in rheumatoid arthritis.^{15–18} These model peptides represent a variety of amino acids, e.g. basic, acidic, aromatic and neutral side chains. All peptides were synthesised with free N- and C-termini by a Syro2000 peptide synthesiser using Fmoc-^tBu strategy. Native Arg-containing peptides were prepared along with their Cit-containing variants by systematic replacement of Arg to Cit. Ala-containing peptides were also synthesised as control compounds by substituting all arginines with Ala. A full list of the peptides is shown in Table 1.

Peptides were purified by preparative reversed-phase high-performance liquid chromatography (RP-HPLC) using a Luna C18 column (250 × 10 mm, 5 μm, 100 Å; Phenomenex) with gradient elution (eluent A: 0.1% (v/v) trifluoroacetic acid (TFA) in water; eluent B: 0.1% (v/v) TFA, 20% (v/v) water in acetonitrile; 0 min: 2% B, 2 min: 2% B, 30 min: 97% B). Peptides were freeze-dried and subsequently analysed by tandem mass spectrometry with a high mass accuracy and resolution QTOF Premier hybrid mass spectrometer (Waters) equipped with electrospray ionisation (ESI) source. Samples were directly injected into the ESI source from a solvent mixture of acetonitrile/water (1:1, v/v) with 0.1% formic acid content at a flow rate of 10 μL/min.

3 | RESULTS

This research is focused on the collision-induced dissociation (CID) of oligopeptides containing citrulline residues to identify their preferred fragmentation routes. For this, fragmentation characteristics were

TABLE 1 Sequence and analytical data of the peptides

Sequence	Origin	Calculated monoisotopic molecular mass / Da	Measured monoisotopic molecular mass / Da	Mass accuracy / ppm
GER	collagen (multiple repetition)	360.1757	360.1755	-0.6
GEX	Cit-analogue	361.1597	361.1598	0.3
GEA	Ala-analogue	275.1117	275.1118	0.4
TRGRS	filaggrin(311–315)	575.3140	575.3148	1.4
TRGXS	Cit-analogue	576.2980	576.2978	-0.3
TXGRS	Cit-analogue	576.2980	576.2981	0.2
TXGXS	Cit-analogue	577.2820	577.2813	-1.2
TAGAS	Ala-analogue	405.1860	405.1863	0.7
VERHQS	fibrin α 40–45	754.3722	754.3738	2.1
VEXHQS	Cit-analogue	755.3562	755.3563	0.1
VEAHQS	Ala-analogue	669.3082	669.3073	-1.3
GPRVVER	fibrin α 36–42	811.4664	811.4693	3.6
GPRVVEX	Cit-analogue	812.4504	812.4526	2.7
GPXVVER	Cit-analogue	812.4504	812.4489	-1.8
GPXVVEX	Cit-analogue	813.4345	813.4353	1.0
GPAVVEA	Ala-analogue	641.3384	641.3383	-0.2
GYRARPAK	fibrin β 70–77	917.5195	917.5220	2.7
GYRAXPAK	Cit-analogue	918.5035	918.5075	4.4
GYXARPAK	Cit-analogue	918.5035	918.5067	3.5
GYXAXPAK	Cit-analogue	919.4876	919.4907	3.4
GYAAAPAK	Ala-analogue	747.3915	747.3922	0.9

compared by using oligopeptides and their substituted analogues of 3–8 amino acid residues containing one or two arginines. In these compounds, Arg residues were replaced either by Cit or by Ala (Table 1).

Major fragmentation characteristics of the peptides can be well interpreted by the mobile proton theory,²⁰ which we find useful for Cit-peptides as well. Based on this, peptides could be categorised as having either (1) no mobile protons (where the number of arginines is equal to or greater than the number of ionising protons within the sequence), (2) partially mobile protons (where the number of ionising protons is greater than that of arginines but fewer than that of or equal to the overall basic residues and *N*-terminus), or (3) mobile protons (where the number of ionising protons is greater than that of the overall basic sites).²¹ It should also be noted that this categorisation is only for practical purposes and sometimes lower charge states are missing from the single-stage MS spectra. This is the case for large non-tryptic peptides or the ones containing several slightly basic amino acids (His, Lys, Cit). Protons of peptide ions with low charge states in this category sometimes behave as if they were non-mobile ones (e.g. the singly protonated GPXVVEX peptide, data not shown).

We found that charge-state distribution in single-stage MS is altered and shifted to lower values by substituting Arg residues with Cit ones, suggesting the neutral characteristic of citrulline under ESI (Figure S1, supporting information). This is in agreement with previous studies.⁹ In case of the pentapeptides, however, higher charge states were not necessarily absent, only less intensive, indicating that Cit could be partially charged under MS conditions at least for a limited number of peptides. On the other hand, Cit could be considered a neutral residue due to lack of proton sequestration ability in our MS/MS experiments.

Charge states of model peptides having no mobile protons usually tend to produce fragments corresponding to the neutral loss of isocyanic acid as base peak or as the second most intensive peak at low collision energies (Figure S2, supporting information). HNC loss of citrulline residues yields ornithines in the sequence. High intensity of HNC loss can be explained by the proton sequestration of arginine residues disabling low-energy protons to migrate through the sequence inducing alternate decomposition pathways. Enhanced collision energy has a beneficial impact on generating sequence ions but along with hardly identifiable combinations of complex neutral losses. This latter feature was especially pronounced in the case of the threonine-containing pentapeptide series. It is, however, important to note that the intensities of precursor ions having no mobile protons are usually low amongst peptides derived from enzymatic cleavage of proteins.

Parent ions having partially mobile protons also show highly abundant loss of HNC, often being base peaks at low energies (Figure 3C, Figure S3B, supporting information). In these cases, the neutral loss of isocyanic acid was accompanied by highly abundant $b_x\text{-}y_z$ products originated from a preferential cleavage site that is C-terminal to Cit residues (Figures 3B, 3C, 3D). Preferential cleavage of the Cit–Aaa bond of peptides in the gas phase is described here for the first time and termed the citrulline effect. In our opinion, cleavage of the Cit–Aaa bond is facilitated by the formation of ornithine (Orn) residues during the neutral loss of HNC from citrulline. The following mechanism could explain the process: the peptide containing citrulline(s) is first protonated on one of its basic sites. After collision-induced excitation, the proton migrates to the δ -NH-group initiating the dissociation process. The reaction leads to a protonated peptidyl-Orn residue (Figure 2A).¹³ In the second

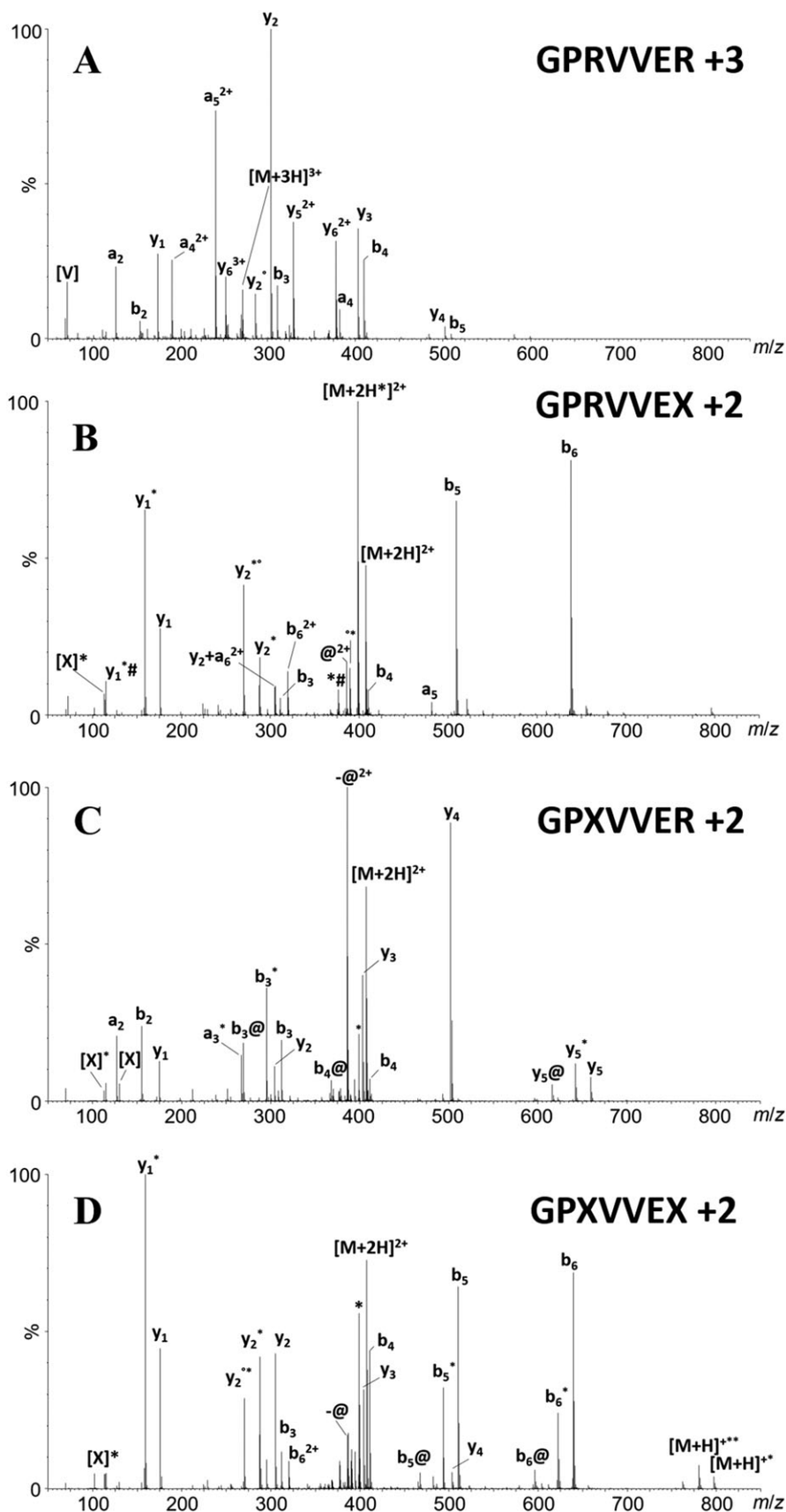


FIGURE 3 Tandem mass spectra of the heptapeptide model series. Triply protonated GPRVVER fragmented at a collision energy of 7.5 eV (A), doubly protonated GPRVVEX at 10.0 eV (B), doubly protonated GPXVVER at 10.0 eV (C) and doubly protonated GPXVVEX at 7.5 eV (D). HNCO (@), NH_3 (*), H_2O (°) and CO_2 (#) side chain losses are also indicated. Charge states and collision energies were tuned that the protonated molecules, the sequence ions and isocyanic acid loss could be detected with a reasonable yield. It should be noted that the +2 ion of peptide GPRVVER at a collision energy of 10.0 eV remained intact due to the proton sequestration of Arg. Cit-containing peptides produce abundant peaks corresponding to Cit-Aaa cleavages due to the citrulline effect with charge retention on the fragment with the larger proton affinity

step, a favourable, six-membered lactam ring is formed (Figure 2B) from Orn. Facile cleavage of residues C-terminal to Orn within peptides in the gas phase has been previously described by McGee and McLuckey.²²

Enhanced collision energy results in the appearance of other informative sequence fragments, as well as their 'ornithinated' variants by HNCO loss while the peaks corresponding to the citrulline effect remain very intensive. We also found (Figure S3, supporting information) that the proline effect was inferior to the citrulline effect, similarly to previously described results for ornithine-containing peptides.²² The only exception was the peptide GYXARPAK, in which a more preferred cleavage at the YX peptide bond was observed leading to the intensive peak pair of b_2 and a_2 fragment ions (Figure S3, supporting information).

If Cit is in an $(n-1)$ position relative to the C-terminus, the intensity of the y_1 ion will also be increased. The pentapeptide model set based on the TRGRS peptide sequence is shown in Figure S4 (supporting information) as an example. Y_1 ions are characteristic to tryptic peptides and confirm the identity of C-terminal Arg or Lys. These important fragment ions are also present in the spectra of peptides containing C-terminal Cit with high intensities and are generally accompanied by NH_3 loss (Figures 3B and 3D).

Peptide charge states having mobile protons also display the loss of HNCO; however, backbone fragmentation becomes a competitive pathway. The former process is therefore less pronounced. Fragment ions show losses of HNCO as well. In these cases, complementary b/y fragment ions corresponding to the citrulline effect may not have significantly higher intensities than other ones. Sequence fragments which are not present in the tandem mass spectra of charge states

possessing partially mobile protons could also appear for alternative charge states having mobile protons.

The most intensive fragment ions of the peptides and ions with HNCO loss are summarised in Table 2.

4 | DISCUSSION

Although several methods have been developed for the analysis of citrullination, a comprehensive study of fragmentation characteristics at the peptide level is still not available. In this work, we identified basic principles connected to the MS/MS behaviour of peptides containing citrulline(s) to improve MS-based detection of deimination in terms of specificity, reliability and speed in the future.

Model peptides GER, TRGRS, VERHQS, GPRVVER, GYRARPAK and all their possible deiminated variants were synthesised by solid-phase peptide synthesis (Table 1). Control peptides were also synthesised by systematic replacement of Cit residues by Ala. The peptides selected correspond to the epitope regions of proteins (collagen, filaggrin, fibrin) which are targets of immunorecognition in rheumatoid arthritic joint tissues. Some of the peptides mimic cleavage results by trypsin or LysC (Arg or Lys residues at the C-termini).

Our results were in good agreement with the mobile proton hypothesis, described previously for general interpretation of peptide fragmentation by tandem mass spectrometry.²⁰

We observed that peptide charge states having partially mobile protons provide a pronounced intensity of fragments corresponding to the Cit-Aaa cleavage at low-energy CID leading to b- and/or y-type

TABLE 2 Fragmentation of the peptides

Sequence	Detected N-terminal sequential fragments	Detected C-terminal sequential fragments	Loss of HNCO detected
GER	b_2	Y_1, Y_2	-
GEX	b_2	$Y_1, (Y_2)$	MH^+, Y_1
GEA	b_2	$Y_1, (Y_2)$	-
TRGRS	a_1, b_2, b_3	Y_1, Y_2, Y_3	-
TRGXS	a_1, b_2, b_3, b_4	Y_1	(b_4)
TXGRS	a_1, b_2	Y_1, Y_3, Y_4	$b_2, (Y_4)$
TXGXS	b_2, b_3, b_4	$(Y_1), Y_2, Y_3, (Y_4)$	$MH^+ (2\times), b_2, b_4 (2\times), Y_3$
TAGAS	b_2, b_3, b_4	Y_1, Y_2, Y_3, Y_4	-
VERHQS	$a_1, b_2, b_3, b_4, (b_5)$	Y_1, Y_2, Y_3, Y_4, Y_5	-
VEXHQS	a_1, b_2, b_3, b_4, b_5	$Y_2, Y_3, Y_4, (Y_5)$	MH^+, b_3, b_4, b_5, Y_4
VEAHQS	$a_1, b_2, (b_3), (b_4), (b_5)$	$(Y_1), Y_2, (Y_3), Y_4, Y_5$	-
GPRVVER	$b_2, b_3, b_4, (b_5)$	Y_1, Y_2, Y_3, Y_4	-
GPRVVEV	$(b_2), b_3, b_4, b_5, b_6$	$Y_1, Y_2, (Y_3)$	MH^{2+}
GPXVVER	b_2, b_3, b_4	Y_1, Y_2, Y_3, Y_4, Y_5	b_3, b_4, Y_5
GPXVVEV	$(b_2), b_3, b_4, b_5, b_6$	Y_1, Y_2, Y_3, Y_4	$MH^{2+}, MH^{2+(2\times)}, (b_4), b_5, b_6, (Y_3)$
GPAVVEA	$(b_2), b_3, b_4, b_5, b_6$	Y_2, Y_3, Y_4	-
GYRARPAK	$a_2, b_3, (b_4), (b_5), (b_6)$	$Y_1, Y_2, Y_3, (Y_5)$	-
GYRAXPAK	$a_2, (b_3), (b_4), b_5, b_7$	Y_1, Y_2, Y_3	MH^{2+}, b_5, b_7
GYXARPAK	$a_2, b_3, (b_4), b_5, b_7$	$Y_1, (Y_2), Y_3, Y_4, Y_5, Y_6, Y_7^{2+}$	$MH^+, (b_3), (b_4), (b_5), (b_7), Y_6$
GYXAXPAK	$a_2, b_3, b_4, b_5, (b_7)$	$Y_1, (Y_2), Y_3, Y_4, Y_5, Y_6$	$MH^{2+} (2\times), (b_3), (b_4), (b_5), (Y_4), Y_5, (Y_6), (b_7)$
GYAAAPAK	$b_2, b_3, b_4, (b_5), (b_6)$	$Y_1, Y_2, Y_3, Y_4, Y_5, (Y_6)$	-

Fragments are generated from partially mobile parent ions. Fragments between 1 and 5% of the base peak intensity are in brackets. Peptides containing two citrulline residues show a more intensive fragmentation due to the citrulline effect.

ions. This feature is described here for the first time and it may also be observed for precursor ions with fully mobile protons. A similar phenomenon was described previously²² for the Orn–Aaa amide bond in Orn-containing peptides. We suppose that the isocyanic acid loss from Cit residues, resulting in a newly generated Orn residue in the sequence using CID, explains the facile Cit–Aaa bond cleavage (Figure 2).

Loss of isocyanic acid is also a very selective reaction of citrulline-containing peptides, which can be observed from parent ions and for fragment ions as well. Sequence coverage could also be increased – to a limited extent – by raising collision energies without losing valuable peaks corresponding to the citrulline effect. In our experiments, peptides were also analysed on an ion trap instrument and the compounds showed essentially the same fragmentation profiles indicating the robustness of the method. An example for low-energy CID tandem mass spectra acquired on an ion trap is shown on Figure S5 (supporting information).

Our results indicated that y_1 ions of Cit are abundantly present due to the cleavage at Aaa–Cit, if Cit is located at the C-terminus (Figures 3B and 3D). This is also an important feature, especially given the fact that the first MS-grade mutant enzyme capable of cleaving at the C-terminus of citrulline has been developed recently.²³

Based on our experiments, the major factors effecting favoured fragmentation routes for deiminated peptides could be outlined. These pathways, such as the neutral loss of isocyanic acid, conventional sequential fragmentation and specific sequential fragmentation pathways (e.g. citrulline effect), could be elucidated by the mobile proton theory. Increasing proton mobility decreases the intensities of peaks corresponding to the loss of HNCO as in the case of other types of neutral losses. On the other hand, the citrulline effect may remain intensive in the case of higher proton mobilities or at elevated collision energies. Thus, utilisation of the citrulline effect along with the loss of HNCO could greatly contribute to a more reliable identification and sequencing of deiminated peptides.

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SUPPORTING INFORMATION

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