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Structural signatures of the class III lasso peptide BI-32169 and the branched-cyclic topoisomers using trapped ion mobility spectrometry–mass spectrometry and tandem mass spectrometry

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Abstract

Lasso peptides are a class of bioactive ribosomally synthesized and post-translationally modified peptides (RiPPs) characterized by a mechanically interlocked topology, where the C-terminal tail of the peptide is threaded and trapped within an N-terminal macrolactam ring. BI-32169 is a class III lasso peptide containing one disulfide bond that further stabilizes the lasso structure. In contrast to its branched-cyclic analog, BI-32169 has higher stability and is known to exert a potent inhibitory activity against the human glucagon receptor. In the present work, tandem mass spectrometry, using collision-induced dissociation (CID) and electron capture dissociation (ECD), and trapped ion mobility spectrometry-mass spectrometry (TIMS-MS) experiments were carried out to evidence specific structural signatures of the two topologies. CID experiments showed similar fragmentation patterns for the two topoisomers, where a part of the C-terminal tail remains covalently linked to the macrolactam ring by the disulfide bond, which cannot clearly constitute a signature of the lasso topology. ECD experiments of BI-32169 showed an increase of hydrogen migration events in the loop region when compared with those of its branched-cyclic topoisomer evidencing specific structural signatures for the lasso topology. The high mobility resolving power of TIMS resulted in the identification of multiple conformations for the protonated species but did not allow the clear differentiation of the two topologies in mixture. When in complex with cesium metal ions, a reduced number of conformations led to a clear identification of the two structures. Experiments reducing and alkylating the disulfide bond of BI-32169 showed that the lasso structure is preserved and heat stable and the associated conformational changes provide new insights about the role of the disulfide bond in the inhibitory activity against the human glucagon receptor.

Keywords BI-32169 \cdot Lasso topologies \cdot Branched-cyclic peptides \cdot Collision-induced dissociation \cdot Electron capture dissociation \cdot Trapped ion mobility spectrometry–mass spectrometry

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Introduction

Lasso peptides are a unique class of ribosomally synthesized and post-translationally modified peptides (RiPPs) exhibiting a fascinating mechanically interlocked structure [1]. All the discovered lasso peptides share an N-terminal macrolactam ring, generated by an isopeptide bond between the α -amino group and the side chain carboxyl group of a glutamate or aspartate residue, through which the C-terminal part is threaded (see Electronic Supplementary Material (ESM) Fig. S1). This threaded fold is predominantly stabilized by steric interactions and can also be assisted by the presence of disulfide bonds (ESM Fig. S1) [2-4]. Sterically demanding amino acid side chains (e.g., Phe, Trp, or Tyr), called plugs, are needed to trap the threaded tail within the macrolactam ring and therefore stabilize the entropically disfavored lasso topology (ESM Fig. S1). The lasso peptide family is divided into four classes depending on the presence of one (class III) or two (class I) interlinked disulfide bonds and one (class IV) handcuff disulfide bond while the class II lasso peptides have no disulfide bond [5] (ESM Fig. S1). The highly compact structures of lasso peptides confer a great resistance to chemical and proteolytic degradation as well as, in many cases, against thermally induced unthreading [6–9]. The extraordinary mechanically interlocked topology of lasso peptides encompasses a large panel of functions, such as enzyme inhibitory, receptor antagonistic, antimicrobial, or antiviral properties [1, 10]. A limitation of the biological activity of lasso peptides is the unthreading of the C-terminal tail, a trend reported for several lasso peptides, yielding their corresponding branched-cyclic topoisomers [7–9, 11].

BI-32169 is a class III lasso peptide produced by *Streptomyces* sp. that exerts a potent inhibitory activity against the human glucagon receptor [4, 12]. To date, BI-32169

remains the only member of class III. BI-32169 features a nine residue macrolactam ring, generated by an isopeptide bond between the α -amino group of Gly1 and the side chain carboxyl group of Asp9 (highlighted in green in Fig. 1), a loop of five residues (highlighted in blue) located above the ring, and a C-terminal tail of five residues (highlighted in orange) located below the ring. Its C-terminal part is sterically entrapped in the macrolactam ring by the Trp13 and Trp17 residues located on each side of the ring (highlighted in red in Fig. 1). Additionally, the C-terminal Cys19 residue forms a disulfide bond with Cys6, which is located within the macrolactam ring, resulting in a bicyclic primary structure (Fig. 1). Nuclear magnetic resonance (NMR) was employed to unambiguously characterize the threading of the C-terminal tail through the macrolactam ring [4]. It has been proposed that the Pro12 and Pro16 residues of BI-32169 introduce two kinks resulting in a twisted S-like structure, in which the first kink, involving the Ile10-Trp13 residues, can be characterized as a type I β -turn. This sets the two cysteine residues in close proximity allowing the formation of the disulfide bond. However, NMR approaches require relatively large amounts of samples, and the analyses of mixtures with the branchedcyclic topoisomer are not straightforward. MS-based approaches have been also applied as a tentative to elucidate lasso structures in the gas phase. Cleavage within the Cterminal region of BI-32169 upon collision-induced dissociation (CID) [4, 13] showed associated b_i and y_i product ions where a part of the C-terminal tail remains covalently linked to the macrolactam ring by the disulfide bond. However, these cross-linked product ions do not clearly constitute a signature of the lasso topology, as this type of fragment could also arise from its branched-cyclic topoisomer. Electron capture/transfer dissociation (EC/TD) [13, 14] and ion mobility spectrometrymass spectrometry (IMS-MS) [15, 16] serve as an efficient

Fig. 1 Sequences and schematics of the BI-32169 and its branchedcyclic analog. The macrolactam rings are colored in green, the loop residues in blue, the plugs in red, and the C-terminal tail in orange. The disulfide bonds are shown by black lines



alternative strategy for the identification of the lasso and branched-cyclic topologies.

In the present work, BI-32169 and its branched-cyclic topoisomer (Fig. 1) were investigated using tandem mass spectrometry (CID and ECD) and trapped ion mobility spectrometry–mass spectrometry (TIMS-MS). The CID and ECD patterns as well as the TIMS-MS profiles of the two topologies of BI-32169 were compared for the first time to evidence specific structural signatures for lasso peptides containing disulfide bonds. In addition, the disulfide bond of BI-32169 was reduced/alkylated in order to derive information on the conformational changes in the absence of the disulfide bond, and the lasso peptide was exposed to different temperatures in order to determine if the lasso structure is maintained in absence of the disulfide bond.

Experimental section

Peptides and sample preparation

Details on BI-32169 production have been reported previously [4, 12]. Briefly, Streptomyces sp. cells were grown in GYM medium at 28 °C for 5 days. BI-32169 was purified from the mycelium by extraction using methanol. The resulting extracts were directly subjected to reversed-phase high-performance liquid chromatography (RP-HPLC) for purification. The branched-cyclic peptide of BI-32169 was obtained by solidphase synthesis from Genepep (St Jean de Védas, France). The reduction of the disulfide bond of both peptides was carried out with 1 mM of tris(2-carboxyethyl)phosphine (TCEP) at 50 °C for 15 min. The alkylation was performed using 10 mM of N-ethylmaleimide (NEM) at room temperature for 15 min. Solutions were prepared at a final concentration of 5 µM in 10 mM ammonium acetate (NH₄Ac). To investigate the thermal stability of the BI-32169 and reduced/ alkylated BI-32169, a solution of 10 µM of lasso peptide was incubated at 25 °C, 50 °C, 75 °C, and 95 °C for 3 h. The same procedure was applied to the branched-cyclic analog of BI-32169 and the corresponding reduced/alkylated peptide as a control. Samples were subsequently analyzed via TIMS-MS. Low-concentration Tuning Mix calibration standard (TuneMix, G24221A) was purchased from Agilent Technologies (Santa Clara, CA). Details of the Tuning Mix structures are described elsewhere [17].

ECD experiments

ECD experiments were carried out on a Solarix 7-T FT-ICR mass spectrometer (Bruker, Billerica, MA) equipped with a nanoESI source operating in the positive ion mode. Sample aliquots (10 μ L) were loaded in a pulled-tip capillary to the MS inlet. The ESI high voltage, capillary exit, and skimmers I

and II were set to 1500 V, 200 V, 5 V, and 16 V, respectively. ECD experiments were performed with a heated hollow cathode operating at a current of 1.6 A. Electrons emitted during 0.2 s were injected into the ICR cell with a 1.5-V bias and 16-V ECD lens.

Native TIMS-MS and CID experiments

Details regarding the TIMS operation can be found elsewhere [18, 19]. Briefly, ion mobility experiments were performed on a custom built nanoESI-TIMS coupled to an Impact Q-TOF mass spectrometer (Bruker, Billerica, MA; ESM Fig. S2) [18]. The TIMS unit is controlled using a custom software in LabView (National Instruments) synchronized with the MS platform controls [19]. Sample aliquots (10 μ L) were loaded in a pulled-tip capillary biased at 700-1500 V relative to the MS inlet. TIMS separation is based on holding the ions stationary using an electric field (E) against a moving buffer gas (ESM Fig. S2) [20]. In TIMS operation, multiple isomers/ conformers are trapped simultaneously at different E values resulting from a voltage gradient applied across the IMS tunnel region (ESM Fig. S2). Mobility-selected ions are then eluted from the TIMS analyzer region by decreasing the electric field (ESM Fig. S2). TIMS separation was carried out using nitrogen (N_2) as buffer gas at room temperature (T). The v_{g} is set by the pressure difference between the funnel entrance (P1 = 2.6 mbar) and exit (P2 = 1.1 mbar; ESM Fig. S2). An rf voltage of 250 V_{pp} at 880 kHz was applied to all electrodes. Separations were performed using a voltage ramp (V_{ramp}) of -250 to -100 V, deflector voltage (V_{def}) of 60 V, and base voltage (V_{out}) of 60 V. TIMS separation depends on the gas flow velocity (v_g) , elution voltage $(V_{elution})$, ramp time (t_{ramp}) , and base voltage (V_{out}) [18, 20]. The reduced mobility, K_0 , is defined by:

$$K_0 = \frac{v_{\rm g}}{E} \cong \frac{A}{(V_{elution} - V_{out})} \tag{1}$$

The constant *A* was determined using known reduced mobilities of calibration standards (Tuning Mix) [17]. The elution voltage was determined experimentally by varying the ramp time for a constant ramp voltage as describe elsewhere [17]. The measured mobilities were converted into collision cross sections (CCS, $Å^2$) using the Mason–Schamp equation:

$$\Omega = \frac{(18\pi)^{1/2}}{16} \frac{q}{(k_B T)^{1/2}} \left(\frac{1}{m} + \frac{1}{M}\right)^{1/2} \frac{1}{N} \times \frac{1}{K}$$
(2)

where q is the ion charge, $k_{\rm B}$ is the Boltzmann constant, N is the gas number density, m is the ion mass, and M is the gas molecule mass [20].

Collision-induced dissociation (CID) experiments were performed in the collision cell located after the TIMS analyzer (ESM Fig. S2). The mass-selected $[M+2H]^{2+}$ ions were fragmented using nitrogen as collision gas at a collision energy of 26 V. The mobility resolving power (*R*) and the resolution (*r*) are defined as $R = \Omega/w$ and $r = 1.18 \times (\Omega_2 - \Omega_1)/(w_1 + w_2)$, where *w* is the full peak width at half maximum (FWHM).

Results and discussion

CID fragmentation pattern of BI-32169

The CID spectra of the doubly protonated species of the class III lasso peptide BI-32169 (m/z 1018.9) and its branchedcyclic topoisomer are illustrated in Fig. 2a, b, respectively. The CID spectrum of BI-32169 displayed a complicated fragmentation pattern due to the presence of the disulfide bond (Fig. 2a). The disulfide bond linking the C-terminus to the macrolactam ring prevents the formation of classic b_i/v_i series. However, cross-linked product ions from b_i and v_i ions, denoted as $[(b_i)^{-}(v_i)]$, were observed. These fragments result from two bond cleavages in the C-terminal part, where a part of the C-terminal tail remains covalently linked to the macrolactam ring by the disulfide bond as illustrated in Fig. 2a. Such cross-linked species were also previously observed for the class I lasso peptides sviceucin and siamycin I which contain two disulfide bonds [13]. The present cross-linked product ions are different from the mechanically interlocked fragments, denoted as $[(b_i)^*(y_i)]$, previously observed for class II lasso peptides, where b_i and y_i ions remain associated through steric interactions imparted by bulky residues, which entrap the C-terminal tail inside the macrolactam ring [13].



Fig. 2 CID spectra of the doubly protonated species of **a** BI-32169 and **b** its branched-cyclic topoisomer (m/z 1018.9). Typical cross-linked product ions are highlighted in red and labeled on the peptide cartoons (right of

each panel). The macrolactam rings, the loop residues, the plugs, the Cterminal tails, and the disulfide bonds are highlighted in green, blue, red, orange, and black, respectively

The main cross-linked product ions observed for BI-32169 were $[(b_9)^-(y_9)]^{2+}$, $[(b_{10})^-(y_2)]$, $[(b_{10})^-(y_4)]$, and $[(b_{10})^-(y_5)]$, associated with their complementary internal fragment ions $(b_{10}y_{10})_1$, $(b_{17}y_9)_7$, $(b_{15}y_9)_5$, and $(b_{14}y_9)_4$, respectively. The presence of a disulfide bond in the lasso structure could also explain the formation of many internal product ions (Fig. 2a).

The CID spectrum of the doubly protonated species of the branched-cyclic analog of BI-32169 exhibited a similar fragmentation pattern as compared with the lasso structure (Fig. 2b). The same cross-linked product ions and internal fragments were observed for the two topologies. Contrary to class II lasso peptides, for which lasso-specific mechanically interlocked fragments are observed, the CID experiment of BI-32169 is less informative as the cross-linked products occur for the branched-cyclic analog of BI-32169 as well (Fig. 2). Thus, CID experiments are not suitable to determine structural signatures for the lasso topology of class I and III lasso peptides containing disulfide bonds.

ECD fragmentation pattern of BI-32169

We previously demonstrated the utility of ETD for identifying class II lasso peptide topologies as an alternative fragmentation approach to overcome the limited structural information provided by CID experiments [13]. The class III lasso peptide BI-32169 and its branched-cyclic analog displayed very close fragmentation patterns in ECD for the doubly protonated species (Fig. 3). Both topologies presented the charge-reduced $[M+2H]^{+}$ ions (m/z 2037.9) as the most abundant species. ECD has been showed to be very powerful for biomolecules containing disulfide bonds as the capture of a single electron can induce a dissociation of both backbone and disulfide bonds [21, 22]. In fact, c'_i/z_j^{\bullet} series were observed, consisting of c'_{12} to c'_{18} (except c'_{15}) together with their complementary z_7 to z_1 (except z_4), that involve the cleavage of the disulfide bond between Cys6 and Cys19 concomitant with backbone cleavages within the C-terminal part of both topologies (Fig. 3). As previously mentioned for class II lasso peptides [13, 14], significantly different relative abundance was observed for the fragments only comprising the macrolactam ring. Indeed, the branched-cyclic analog of BI-32169 exhibited abundant c'_9 and z_{10} fragments (Fig. 3b) while z_{10} and c'_9 were much smaller and absent, respectively, for the lasso structure (Fig. 3a). This difference probably means that the two complementary c'_9 and z_{10} fragments remained sterically interlocked in the lasso topology.

As with class II lasso peptides [13, 14], ECD spectra of BI-32169 displayed an increase of hydrogen migration events near the loop region (Ile10-Asn14) when compared with those of the branched-cyclic topoisomer, as illustrated in Fig. 3c. In fact, BI-32169 showed fragment ions involving extensive hydrogen migration within c_{12} to c_{14} and their complementary z '7 to z'_5 product ions (highlighted in red in Fig. 3) while these fragments were lower in relative intensity for the branchedcyclic topology, indicating that the hydrogen migration events in the Ile10-Asn14 region occur less frequently in the absence of a lasso structure (Fig. 3c). Contrary to CID, where BI-32169 and the branched-cyclic topoisomer present similar fragmentation patterns, ECD enables specific structural signatures for the class III lasso peptide BI-32169 through the presence of extensive hydrogen migrations. As a consequence, the combination of the present results with a previous study [13] enables us to firmly propose that ECD can be used to characterize lasso topologies regardless of the lasso peptide class and the size of the loop and to differentiate lasso peptides from their unthreaded branched-cyclic topoisomers. However, the differences in the hydrogen migration events were less pronounced as previously observed for class II lasso peptides [13]. This could be explained by the presence of the disulfide bond that links the macrolactam ring to the C-terminus, which imposes additional constraints for the branched-cyclic topoisomer of BI-32169 by fixing the C-terminal part in closer proximity to the macrolactam ring. This feature facilitates hydrogen migration events more strongly than the completely unstructured C-terminal part in the branched-cyclic analogs of class II lasso peptides.

Native nESI-TIMS-MS analysis of BI-32169

We demonstrated the utility of the ECD approach to evidence specific structural signatures for lasso peptides containing disulfide bonds. However, the less pronounced differences in the hydrogen migration events combined with the limited application for the case of mixtures caused us to consider TIMS-MS as a complementary strategy. TIMS-MS was proved very effective in characterizing class II lasso peptides and in differentiating them from their unthreaded branched-cyclic topoisomers [15, 16]. Native TIMS spectra corresponding to the doubly protonated species of BI-32169 and its branchedcyclic topoisomer are presented in Fig. 4a. The collision cross sections (CCS) and resolving power (R) metrics are listed in Table S1 (see ESM). Previous traveling wave IMS (TWIMS) experiments showed a single broad arrival time distribution $(R \sim 15)$ for the doubly protonated species of BI-32169 (ESM Fig. S3) [23]. By contrast, the high mobility resolving power $(R \sim 70-200)$ achieved by using TIMS at fast scan rates (Sr = 0.3 V/ms) resulted in the identification of multiple IMS bands for the two topoisomers, providing additional information on the conformational spaces adopted during native conditions, as previously shown for the lasso topology of BI-32169 [15] (Fig. 4a). In addition, the multiple IMS bands of BI-32169 cover a relatively large CCS region as compared with those of the class II lasso peptides [15], suggesting that the disulfide bond prevents the peptide from collapsing to more compact structures. We previously demonstrated that the branchedcyclic topologies adopt significantly more extended structures



Fig. 3 ECD spectra of the doubly protonated species of **a** BI-32169 and **b** the branched-cyclic topoisomer (m/z 1018.9). Typical hydrogen migration events are highlighted in red and labeled on the peptide cartoons (right of each panel). **c** Bar plot showing the hydrogen migration events of BI-32169 (blue bars) and the branched-cyclic topoisomer (red bars)

obtained by comparison between the experimental and theoretical isotope patterns. The macrolactam rings, the loop residues, the plugs, the Cterminal tails, and the disulfide bonds are highlighted in green, blue, red, orange, and black, respectively

The use of metalation, especially the doubly cesiated spe-

cies, has been shown to be very powerful in reducing the

conformational space (i.e., number of IMS bands) via intra-

molecular complexation as well as in separating lasso and

branched-cyclic topologies (see examples in a previous report

than the class II lasso peptides, permitting a clear separation of the two topoisomers in native TIMS-MS using fast scan rates (Sr = 0.3-0.56 V/ms) [15]. In the case of the branched-cyclic analog of BI-32169, the multiple conformations were just slightly more extended than the lasso topology, suggesting that the disulfide bond prevents the peptide from extending of the unthreaded C-terminal part (Fig. 4a). As a consequence, the lasso and branched-cyclic topoisomers of BI-32169 were not clearly separated for the doubly protonated species.



Fig. 4 TIMS spectra of **a** the doubly protonated species and **b** the doubly cesiated species of BI-32169 (blue traces) and its branched-cyclic topoisomer (red traces). The dashed line represents the TIMS profile of both topoisomers in mixture. Schemes highlight the macrolactam rings in green, the loop residues in blue, the plugs in red, and the C-terminal tail in orange. The disulfide bonds are represented by black lines. The *R*, *r*, and *Sr* values are given

resolving power (*R*) metrics are listed in Table S1 (see ESM). Inspection of Fig. 4b shows that the doubly cesiated species resulted in better mobility separation ($r \sim 1.7$) at fast scan rates between BI-32169 and its branched-cyclic topoisomer. In addition, the use of cesium metal ions permitted to considerably reduce the number of conformations for both topoisomers (Fig. 4b). Contrary to the class II lasso peptides, the doubly cesiated species of BI-32169 displayed three distinct IMS bands at similar CCS, suggesting that the disulfide bond imposes strong constraints to the structure making the lasso topology of BI-32169 very rigid; even the introduction of a metal ion with a large ionic radius barely extended the lasso structure. In the case of the doubly cesiated species of the branched-cyclic analog of BI-32169, three distinct IMS bands, including two compact minor conformations, were observed indicating that the disulfide bond prevents the peptide from completely extending the unthreaded C-terminal part, even when introducing a metal ion with a large ionic radius.

The disulfide bonds of the two BI-32169 topoisomers were reduced and alkylated in order to assess the conformational changes of the loss of the disulfide bond on the lasso and branched-cyclic structures. The effect of the reduction and alkylation of the disulfide bonds of BI-32169 and its branched-cyclic topoisomer is presented in Fig. 5a, and the measured collision cross sections (CCS) and resolving power (R) metrics are listed in Table S1 (see ESM). The reduced/ alkylated BI-32169 and the branched-cyclic analog exhibited different mobility profiles, indicating that the lasso structure is preserved even when losing the disulfide bond. The high mobility resolving power ($R \sim 75-235$) also resulted in the identification of multiple conformations for the two reduced and alkylated topoisomers. However, the major IMS band of the reduced/alkylated BI-32169 was observed at significantly higher CCS ($\sim + 30 \text{ Å}^2$) as compared with the major IMS band of the doubly protonated species of BI-32169 (Figs. 4a and 5a). In fact, a low CCS difference ($\sim + 8 \text{ Å}^2$) was obtained by comparing the major IMS band of the doubly protonated and cesiated species of BI-32169. This means that the loss of the disulfide bond makes the C-terminal tail of the reduced/ alkylated BI-32169 more flexible and induces an expansion of the lasso structure. The corresponding branched-cyclic peptide exhibited two major conformations, for which more extended structures were observed as compared with the main conformations of the branched-cyclic analog of BI-32169 (Figs. 4a and 5a). This indicates an unfolding of the free Cterminal tail due to the absence of the disulfide bond. Nevertheless, this trend was less pronounced for the main compact structures when compared with that for the lasso structure, as reflected by the smaller CCS difference ($\sim +$ 15 $Å^2$) between the branched-cyclic analog of BI-32169 and its reduced/alkylated form, even adopting an overall topology close to the reduced/alkylated BI-32169. This suggests that the free C-terminal tail of the reduced/alkylated branchedcyclic analog of BI-32169 can also be folded probably by making intramolecular interactions between the flexible Cterminal tail and the macrolactam ring.

Thermal stability of BI-32169

As a consequence of their compact and interlocked structures, many lasso peptides exhibit an outstanding stability toward Fig. 5 TIMS spectra (Sr = 0.3 V/ ms) of the doubly protonated species of the reduced/alkylated BI-32169 (blue traces) and the corresponding branched-cyclic peptide (red traces) at a 25 °C, b 50 °C, c 75 °C, and d 95 °C. Schemes highlight the macrolactam rings in green, the loop in blue, the plugs in red, and the C-terminal tail in orange. The alkylating reagents (NEM) are represented in yellow



thermal unthreading [7, 11, 25–27]. However, several lasso peptides have been reported to unfold at high temperature into their branched-cyclic topoisomers [7, 8, 26, 28]. These findings encouraged us to study the thermostability of BI-32169 and the reduced/alkylated form of BI-32169. The effect of the temperature on the conformational space of BI-32169 and BI-32169 reduced/alkylated together with their respective branched-cyclic topoisomers is illustrated in ESM Fig. S4 and in Fig. 5, respectively. The TIMS experiments carried out on the branched-cyclic peptides served as a control of the unfolding of the lasso topologies. Similar TIMS profiles were observed when BI-32169 was subjected to solution temperatures of 25 °C, 50 °C, 75 °C, and 95 °C for 3 h (ESM Fig. S4). This suggests that BI-32169 can withstand prolonged exposure to high temperature. Because of the strong heat stability of BI-32169, the reduced/alkylated form of BI-32169,

which has been demonstrated to be less rigid, was also subjected to the same treatment (solution temperatures of 25 °C, 50 °C, 75 °C, and 95 °C for 3 h, Fig. 5). As for BI-32169, the reduced/alkylated form displayed similar TIMS profiles under these conditions, indicating that the lasso structure is preserved and therefore heat stable. This suggests that the two bulky residues Trp13 and Trp17, which are located above and below the ring respectively, are sufficient to maintain the lasso topology even at highly elevated temperatures.

Conclusion

Herein, mass spectrometry experiments were performed for the first time on a lasso peptide containing a disulfide bond together with the corresponding branched-cyclic topoisomer to find specific structural signatures of the two topologies. The CID experiments showed similar fragmentation pattern for the two topologies, where a part of the C-terminal tail remains covalently linked to the macrolactam ring by the disulfide bond. As these cross-linked product ions also occurred for the branched-cyclic topology, CID did not revealed specific signatures of the lasso topology of BI-32169. That is, CID experiments are only effective for class II lasso peptides in which the loop region is five residues or longer.

ECD experiments of BI-32169 showed a larger extent of hydrogen migration near the loop region, when compared with those of the branched-cyclic topoisomer, providing evidence of specific structural signatures for the lasso topology. Note that the differences in the hydrogen migration events of BI-32169 and the branched-cyclic topoisomer are less pronounced, when compared with those of the class II lasso peptides, due to the presence of the disulfide bond which imposes additional constraints to the two topologies. These results suggest that ECD can be efficient to characterize lasso peptides regardless of the class and loop region size and to differentiate the lasso topologies from their unthreaded branched-cyclic topoisomers.

The potential of native nESI-TIMS-MS for identifying the class III lasso peptide BI-32169 from the branched-cyclic topoisomers is illustrated for the doubly protonated species. The lasso and branched-cyclic topologies of BI-32169 are clearly separated when complexed with cesium metal ions due to the reduction on the conformational space, and not when in complex with sodium and potassium. The present TIMS-MS data on lasso peptide containing disulfide bond combined with a previous work on class II lasso peptides (without disulfide bond) [24] allow to strongly propose that the metal cesium ions can be used to clearly separate any lasso peptide from its branched-cyclic topoisomers regardless of the class and loop size. We also showed the thermally stable behavior of BI-32169 and evidenced conformational changes by reducing and alkylating the disulfide bond of BI-32169. This provides new insights about the presence of the disulfide bond in the lasso structure which probably has a crucial role in the inhibitory activity against the human glucagon receptor.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no competing interests.

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