Isolation and characterization of Bradykinin potentiating peptides from Agkistrodon bilineatus venom

Aisha Munawar, Anum Zahid, Amr Negm, Ahmed Akrem, Patrick Spencer and Christian Betzel

Abstract

Background: Snake venom is a source of many pharmacologically important molecules. Agkistrodon bilineatus commonly known as Cantil, is spread over Central America particularly in Mexico and Costa Rica. From the venom of Agkistrodon bilineatus we have isolated and characterised six hypotensive peptides, and two bradykinin inhibitor peptides. The IC-50 value of four synthesized peptides was studied, towards angiotensin converting enzyme, in order to study the structure-function relationship of these peptides.

Results: The purification of the peptides was carried out by size exclusion chromatography, followed by reverse phase chromatography. Sequences of all peptides were determined applying MALDI-TOF/TOF mass spectrometry. These hypotensive peptides bear homology to bradykinin potentiating peptides and venom vasodilator peptide. The peptide with m/z 1355.53 (M + H)\(^+\), and the corresponding sequence ZQWAQGRAPHPP, we identified for the first time. A precursor protein containing a fragment of this peptide was reported at genome level, (Uniprot ID P68515), in Bothrops insularis venom gland. These proline rich hypotensive peptides or bradykinin potentiating peptides are usually present in the venom of Crotalinae, and exhibit specificity in binding to the C domain of somatic angiotensin converting enzyme. Four of these hypotensive peptides, were selected and synthesized to obtain the required quantity to study their IC50 values in complex with the angiotensin converting enzyme. The peptide with the sequence ZLWPRPQIPP displayed the lowest IC50 value of 0.64 \(\mu\)M. The IC50 value of the peptide ZQWAQGRAPHPP was 3.63 \(\mu\)M.

Conclusion: The canonical snake venom BPPs classically display the IPP motif at the C-terminus. Our data suggest that the replacement of the highly conserved hydrophobic isoleucine by histidine does not affect the inhibitory activity, indicating that isoleucine is not mandatory to inhibit the angiotensin converting enzyme. The evaluation of IC 50 values show that the peptide with basic pI value exhibits a lower IC 50 value.

Keywords: Snake venom, Peptides, Angiotensin converting enzyme, Hypotension, MALDI/TOF-TOF, Size exclusion chromatography
stability of some venom peptides is a result of disulfide bond formation and posttranslational modification [9]. The most common type of the posttranslational modification observed in snake venom peptides is a pyroglutamate residue at the N-terminus [10]. Bradykinin potentiating peptides (BPP) are a good example in this respect. These peptides were isolated from the venom of Bothrops jararaca [11]. Captopril, which was the first orally active inhibitor of the angiotensin converting enzyme was designed based on the structure of BPPs isolated from Bothrops jararaca venom [12]. Since then numerous studies have been made to isolate and characterize BPPs from different snake venoms [13–18]. Efforts are being made to study the structure function relationship of this type of peptides in detail, and their possible modes of blood pressure lowering or vasodilatation. For example it was shown that the peptide Bi-PRO-10c [19] inhibits ACE, however argininosuccinate synthetase, present in the kidney cytosol is its primary target [15, 19–21].

Here we summarize the isolation and characterisation of five bradykinin potentiating peptides, one vasodilator peptide and three bradykinin inhibiting peptides from the venom of Agkistrodon bilineatus, generally known as Cantil. In order to determine their IC 50 values, the inhibitory activities of the synthetic analogues of four natural BPPs peptides towards angiotensin converting enzymes were also studied.

**Results**

**Isolation of Bradykinin potentiating peptides from Agkistrodon bilineatus venom**

The size exclusion chromatogram (Fig. 1) shows the fractionation of crude Agkistrodon bilineatus venom. The inset of the figure shows the SDS-PAGE (Sodium dodecyl sulfate-polyacrylamide gel electrophoresis) of the size exclusion chromatogram peaks, which indicate that the later eluting peaks (from 7 to 10) contains molecules below 18 kDa. The fractions corresponding to Peaks 8 and 9 (Fig. 1) showed inhibition towards the angiotensin converting enzyme. The fractions under these peaks were further fractionated by reverse phase chromatography, on a C-18 column. The activity of the material corresponding to each peak was again tested for inhibitory activity towards the angiotensin converting enzyme and all peptides corresponding to the peaks 1 to 4 (Fig. 2) were found to inhibit the angiotensin converting enzyme. Matrix-assisted laser desorption/ionization time of flight mass spectrometry (MALDI-TOF-MS) demonstrated the presence of peptides between m/z 1019.51- 1370.70 (M + H)⁺. The sequence information of the peptides was obtained by MALDI-TOF/TOF mass spectrometry. Three bradykinin inhibitor peptides were isolated, which differ at their N-terminus. One of these inhibitory peptides has the sequence TPPAGPDVGPR. While the peptide with the molecular mass 1019.51 (M + H)⁺ with the sequence PPAGPDVGPRG lacks a threonine at its N-terminus and has a glycine at its C-terminus, which is not present in the other sequence. In addition to these bradykinin inhibitors six bradykinin potentiating peptides were identified, all of which are homologues to each other. Among these six BPP peptides, five have XPP residues at their C-terminus, a highly conserved motif for BPPs, while the ZQWAQGRAPHPP peptide with 1355.53 Da (M + H)⁺ molecular weight, has a histidine instead of the canonical isoleucine just before the two prolines. Secondly all these BPP peptides have a pyroglutamate, represented by Z (Table 1) at their N-terminus, except the PKVSPRWPPXPP peptide, m/z 1370.70 (M + H)⁺ . This peptide has an atypical N-terminus without glutamate or pyroglutamate.

![Fig. 1](image-url) Fractionation of crude Agkistrodon bilineatus venom applying a size exclusion column, Superdex-75, 16x60 mm, at pH 5. Inset of the figure shows a SDS-PAGE (15 % glycine, non reducing gel) of the fractions
The peptide sequencing was done in an automated mode, by applying Mascot Inhouse search. Although with MALDI-TOF/TOF, it is not possible to differentiate between isobaric leucine and isoleucine, and quasi-isobaric (Gln/Lys) residues, however previous studies, Edman degradation and cDNA, of similar BPPs, support the presence of the amino acid isoleucine instead of leucine, and Gln rather than Lys [16, 22–24]. Therefore at all places in the manuscript X refers to Ile/Leu.

Determination of IC 50 values

The inhibition of ACE by the bradykinin potentiating peptides was observed by measuring the fluorescence of the Dnp group of the ACE fluorescent substrate Abz-Phe-Arg-Lys (Dnp)-Pro-OH. Four BPPs were selected to study the IC 50 values (Table 2). The IC 50 values of these peptides have not been studied before. Peptide 6, with the sequence ZLWPRPQXPP showed the lowest IC 50 value 0.64 μM. The IC 50 value of peptide 7 (ZQWAQGRAPHPP) was 3.63 μM, peptide 5 with a sequence ZSAPGNEAXPP exhibited the highest IC 50 value of 30.00 μM among all the four peptides analysed. Peptide 3 with the sequence ZNWPHPQXPP, had an IC 50 value of 0.76 μM. All four peptides were able to inhibit the ACE activity, but to a different extent. This was also demonstrated by dose response curve (Fig. 3) and the calculation of IC 50 values (Table 2). The IC 50 values of the selected peptide inhibitors reflected a difference in the affinity towards the ACE. The order from stronger to weaker inhibition is found to be as

![Graph showing Absorbance at 280 nm vs Time (min) with peak fractions 8-9](Image)

Fig. 2 Further purification of peak fractions 8-9, shown before in Fig. 1, after filtering through a 3 kDa membrane, applying a C-18 column (100 X 4.6 mm)

Table 1 Peptides identified in *Agkistrodon bilineatus* venom

| RP fraction | Peptide identity | Observed Mass (M + H)+ | Mascot Score | Sequence | Homology with the peptide | Peptide Family |
|-------------|------------------|------------------------|--------------|----------|---------------------------|---------------|
| 1           | Peptide1         | 1019.51                | 18           | PPAGPDVGPR | Q27149: Lachesis muta muta | BPP and CNP   |
| 2           | Peptide2         | 1063.54                | 85           | TPPAGPDVGPR| P85025: Agkistrodon bilineatus | BPP and CNP |
| 1           | Peptide3         | 1063.54                | 24           | ZSAPGNEAXPP| P0C7K3: Crotalus viridis viridis | BPP |
| 3           | Peptide5         | 1190.58                | 62           | ZNWPHPQXPP | Q7T1M3: Bothrops jararacussu | BPP and CNP |
| 4           | Peptide6         | 1214.65                | 19           | ZLWPRPQXPP | P0C756: Crotalus atrox | BPP |
| 3           | Peptide7         | 1355.53                | 12           | ZQWAQGRAPHPP| P68515: Bothrops insularis | BPP and CNP |
| 4           | Peptide8         | 1370.70                | 58           | QGWWPRPGPXPP| P0C7R7: Bothrops alternatus | BPP |
| 4           | Peptide9         | 1370.70                | 21           | PKVSPRRWPPXPP| P84746: Bothrops jararacussu | venom vasodilator peptide |

Abbreviations used: Z, pyroglutamate; BPP, Bradykinin potentiating peptide; CNP, C-type natriuretic peptide. X, Isoleucine/Leucine
follows: ZLWPQXPP > ZNWPHPQXPP > ZQWAQG RAPHPP > ZSAPGNEAXPP. The strongest inhibitors, ZLWPQXPP and ZNWPHPQXPP, both have PQXPP at the C terminus.

Discussion

Hypertension is a result of complex interplay of genetic and environmental factors [25], and involves an intricate pathophysiology, with angiotensin converting enzyme playing a key role in the regulation of blood pressure [26]. The first bradykinin potentiating peptide, isolated from the Bothrops jararaca venom became the precursor of today's most commonly used antihypertensive drugs like captopril or lisinopril [11, 12]. After the discovery of the first BPP, homologous of these proline rich peptides have been isolated also from other snake venoms, particularly from the members of Crotalinae subfamily. The BPPs isolated from Agkistrodon bilineatus show homology with BPPs isolated from other snake venoms (Table 1). A NCBI data base search showed that the peptide 7 with m/z 1355.53 (M + H)⁺ and a corresponding sequence ZQWAQGRAPHPP was identified for the first time at the peptide level. Previously the sequence of the precursor protein of this peptide was reported at the transcript level in the venoms of Bothrops insularis and Bothrops jararaca [27, 28]. This peptide lacks the typical IPP motif at its C-terminus, usually present in the snake venom BPPs. However the low IC 50 value (3.63 μM) of this peptide towards ACE suggests that the IPP motif at the C-terminus is not important for the inhibitory activity of snake venom BPPs towards ACE. The presence of a histidine residue next to the two prolines at the C-terminus indicates the potential involvement of this residue in binding to the zinc ion of the angiotensin converting enzyme, thereby inhibiting its activity. Previous studies report the hypotensive effect and ACE inhibitory ability, of similar BPPs by employing classical pharmacological assays or studying inhibition constants of BPPs over ACE. Since various methods were used for the assay, it is difficult to compare inhibitory ability of BPPs, isolated in this study, with other BPPs having different sequences. Ki value of Bj-PRO-10C was recently reported to be 0.20 μM [24]. BPP-XIe had a Ki value of 0.084 μM and that of BPP-AP was 0.035 μM [18]. The sequence alignment (Fig. 4) shows the homology of peptide 7 with other BPP and CNP type peptides. The same residues are highlighted in yellow colour. It can be inferred from Fig. 4, that in addition to lacking a usual IPP sequence at the C-terminus, peptide 7 also has a unique N-terminus, having a sequence QQWA. The sequence QQWA, which has been highlighted in grey colour, appears several times in the precursor, always before a mature BPP. The sequence analysis of the precursor proteins and the mature peptides (Fig. 4) [14, 27, 29–31], gives a clue that QQWA must be a cleavage site, that might have been misinterpreted by the processing mechanism along the maturation of peptide 7. Also the missing IPP C-terminus in peptide 7 (ZQWAQGRAPHPP) is exactly after HPP on the precursor sequence (P68515, Fig. 4) indicating that the novel peptide 7 is a different processing product of a canonical BPP precursor.

A clustal w sequence alignment of the BPPs isolated in this work is shown in Fig. 5. All these peptides have two prolines at their C-terminus, and the preceding residue in all peptides is an isoleucine, except in peptide 7, which has a histidine at this position. Four peptides (P3, P5, P6, P7) have a pyroglutamate at their N-terminus, while P8 has a glutamine and P9 has a proline instead. The presence of pyroglutamate modification at the N-terminus indicates the natural resistance to degradation to these type of peptides. Studies have shown that the presence of PP residues at the C-terminus of the peptides confers resistance to hydrolysis by peptidases [32]. The analysis of crystal structure of Ang II in complex with sACE, showed that the proline, which is the seventh residue of Ang II, prevents hydrolysis of the peptide by ACE, thus inhibits it [33], and hence could serve as a negative

![Inhibition curves](image)

**Fig. 3** Dose response curves of the synthesized peptides, ACE activity Vs log of inhibitor concentration. The N-terminus of all the synthesized peptides have a pyroglutamate modification, while C-terminus is a free carboxylic acid.
feedback mechanism in the regulation of Ang II. Further the study of crystal complex of C-domain of somatic s-ACE and BPPb showed that the two C-terminal prolines of the peptide strongly interacts with the primary binding site, through the formation of hydrogen bonds with multiple residues [33]. Therefore the presence of several proline residues in BPPs, has equipped these peptides with inherent resistance to hydrolysis. The data obtained in the present work and literature studies show that isoleucine is usually present next to the two C-terminus prolines [14–16, 18, 20, 34–36]. The results of the IC 50 values obtained for the synthesized peptides (Table 2) illustrate that while peptides 5, 6 and 7 show low IC 50 values, peptide 3 has a much higher value. As shown in Table 2 peptide 3 has the lowest pI value among all the four synthesized peptides. Further, sequence analysis of these four peptides (Fig. 5) shows that in peptide 3 two polar residues asparagine and glutamic acid are present next to each other, while in the other three peptides the charged amino acid residues are flanked by either proline, glycine, or tryptophan. The low pI value and the presence of the two charged residues together, might be responsible for the high IC 50 value of this peptide, as these factors might hinder the appropriate binding of this peptide within the active site of ACE. In addition to these BPPs, three bradykinin inhibitory peptides were also isolated and characterized. Peptide 2 was previously reported by Graham, et. al [37] in Agkistrodon bilineatus venom, indicating that these peptides have been preserved as an essential component of snake venoms, over the process of evolution. The same peptide 2 was recently reported from the venom of Agkistrodon bilineatus howardgloydi [38]. A systematic study of BPPs, in terms of their sequence, physical properties and molecular targets might lead to a complete understanding of the mechanism of hypertension induced by these peptides, and will support the design of new drugs with more efficacy, specificity and lower side effects.

**Conclusion**

Snake venoms, are a source of many molecules with biomedical importance. The BPPs isolated and characterized within this work show homology with BPPs already isolated from other snake venoms. However, an extensive data base search revealed that peptide 7 with the sequence ZQWAQGRAPHPP has been identified for the first time. All previous descriptions of this peptide were inferred from transcripts, with no evidence of secretion of the gene product in the venom. The analysis of IC 50 values (Table 2) for ACE, applying four
synthetic replica of these BPPs, provided further insights into the structure-function relationship of these peptides. The low IC 50 value of the synthetic peptide ZQWAQGRAPHPP indicates that the presence isoleucine in the IPP sequence is not mandatory to inhibit ACE. Furthermore the calculated pl values of these peptides suggest that the peptides with basic pl value show a lower IC 50 value. Secondly the sequence analysis of the BPPs isolated and described in this work and other studies [13, 14, 18, 34], show that these peptides usually have a pyroglumate modification at their N-terminus, and have a high ratio of proline residues. These two factors provide inherent stability to these peptides towards degradation, particularly in venom glands, which contain various types of enzymes, as well as in the prey’s blood. Further, it can be concluded that the presence of peptide 2 (bradykinin inhibitor peptide) reinforces the idea that these peptides are conserved in the venom of *Agkistrodon bilineatus* over the process of evolution and that these molecules, so far found in many venoms, must play a key role in predation.

**Experimental**

**Materials**

The snake venom of *Agkistrodon bilineatus bilineatus*, was obtained from Venom Supplies, Australia, and was placed at -20° till further use. The venom was collected from the snakes bred in the farms of Venom Supplies, Australia. All the solvents used were of HPLC grade and were obtained from Merck. The substrate Abz-Phe-Arg-Lys (Dnp)-Pro-OH for angiotensin converting enzyme was obtained from BACHEM. Angiotensin converting enzyme from rabbit lung (A6778 - 0.25 UN), was obtained from Sigma. The synthesized peptides ZLWPRPQIPP, ZQWAQGRAPHPP, ZSAPGNEAIPP, ZNWPHPQIPP were purchased from China Peptide company, Shanghai, China. The peptides were 95 % pure, and were kept at -20° till use. The N-terminus of these synthetic peptide is pyroglutamate (represented by Z), while the C-terminus is an amino acid with free carboxylic group.

**Methods**

**Purification of Bradykinin potentiating peptides (BPP)**

The crude venom (50 mg) was dissolved in 1 ml of 100 mM ammonium acetate (pH 5), and centrifuged at 13000 rps. The undissolved material settled down, and the supernatant of crude venom solution was fractionated using a size exclusion column (Superdex-75, 16 x 60 mm) connected to an ÄKTA Purifier system (GE Healthcare). 100 mM ammonium acetate (pH 5) was used as the elution buffer. The fractionation was performed at a rate of 1 ml/min, and UV absorbance of the eluate was monitored at 220 and 280 nm. Fractions were collected and subjected to SDS-PAGE (15 % glycine gels) under non-reducing conditions. The gels were stained with Coomassie Blue. The peptide fractions (peak 8-9, Fig. 1) showing inhibitory activity towards the angiotensin converting enzyme were filtered through 3 kDa Amicon filter and further purified with Chromolith-C-18 column (100x 4.6 mm), using an Agilent 1200 system. Solvent A was 0.2 % formic acid in water and B was straight acetonitrile. A stepwise gradient 3-30 % B for 28 min, 30-40 % B for 9 min and 40-60 % B for 3 min and a flow rate of 2 ml/min was applied to isolate the peptides (Fig. 2). UV absorbance was monitored at 220 nm and 280 nm.

**Enzyme inhibition assay**

ACE activities in the presence of venom peptides were determined by a fluorescence energy transfer assay using Abz-Phe-Arg-Lys (Dnp)-Pro-OH as a substrate [39]. 1 mg of the substrate was weighed and dissolved in 1 ml DMSO. The exact concentration of the substrate was determined by taking four different volumes of the substrate stock solution, and constructing a standard curve spectrophotometrically at 365 nm, using the molar extinction coefficient of the Dnp group (ε356 = 17,300 M⁻¹ cm⁻¹), according to the Beer’s Lambert Law (A = εDnp x l x c). A stock solution of the enzyme was prepared by suspending 0.25 UN of ACE in 250 µl of the assay buffer (12.10 g Tris-base, 2.92 g NaCl and 1.36 mg ZnCl₂ in 1 l of deionised water, pH to 7.0 adjusted with HCl).

In order to determine the IC 50 values of the synthesized peptides, a stock solution (10 mM) of each was prepared in the assay buffer. From the stock solutions, six serial dilutions were prepared for each peptide. The concentrations of the six dilutions were as follows, 0.500 mM, 0.167 mM, 0.056 mM, 0.019 mM, 0.006 mM and 0.002 mM respectively. In the final assay, 85 µl of the assay buffer, 2 µl of the enzyme stock solution, and 5 µl of the dilute peptide solution were added. The mixture was incubated for 10 min at room temperature. The reaction was started by adding 10 µl of the working solution of the substrate (20 µl of the stock +280 µl of the assay buffer). The same procedure was adopted using 5 µl of venom fractions instead of the synthetic peptides. Fluorescence measurements were made at λex = 320 nm and at λem = 420 nm, for 5 min each. These experiments were repeated five times to ensure accuracy of the results.

**Matrix-assisted desorption/ionization time-of-flight mass spectrometry**

MALDI-TOF-TOF analyses were performed with a ultraflexXtreme instrument (Bruker Daltonics, Bremen, Germany). Samples were dried after reversed phase chromatography, dissolved in 30 % ACN, 0.1 % TFA in H2O and 1 µl of the solution was spotted on a MALDI target plate (MTP AnchorChip 384, Bruker Daltonics).
After drying 1 μl MALDI matrix (0.7 mg/ml Cyano-4-hydroxycinnamic acid (Bruker Daltonics) dissolved in 85 % ACN, 1 mM NH4H2PO4 and 0.1 % TFA dissolved in H2O) were spotted on the sample plate.

Data acquisition was performed in positive ion mode using the flexControl software 3.3. The parameters were set as follows: ion source 1: 25 kV, ion source 2: 23.6 kV, lens: 7.5 kV. MS data were collected automatically using

Fig. 6 MALDI-TOF/TOF-MS of selected peptides. a. TPPAGPDVGPR [m/z 1063.55 (M + H)+]; b. ZNWPHPQIPP [m/z 1196.58 (M + H)+]; c. QGGWPRPGEIPP [m/z 1370.70 (M + H)+]. The figures show b and y ion series and the sequence annotation of the corresponding peptide.
autoXecute. Parameters were set as follows: laser power: 47 %; laser shots: 1000; movement, random walk with 100 shots per raster spot. Peaks were selected for LIFT measurement if they met the following criteria: signal to noise > 8, peak intensity > 300.

MS spectra were processed applying flexAnalysis (version 3.3, Bruker Daltonics). Further data analysis was performed using BioTools (version 3.2, Bruker Daltonics) and Mascot Inhouse Search. Mascot version 2.1.03 was used to analyse and search the spectra against the subset “other lobe-finned fish and tetrapod clade” of the Swissprot database. The precursor ion mass tolerance was set to 1 Da, the fragment ion mass tolerance was 0.5 Da.

The sequence annotation pictures of the representative peptides, prepared applying the Bruker software ProteinScape, version 3.0, are shown in Fig. 6.

Abbreviations

ACE: angiotensin converting enzyme; BPP: bradykinin potentiating peptides; Z: pyroglutamate; X: Isoleucine/leucine.

Competing interest

The authors declare no conflict of interest.

Authors’ contributions

AM carried out all the experiments and drafted the manuscript, AZ participated in the purification of peptides, AN and AM performed enzyme inhibition assays and performed the statistical analysis, AA participated in the purification of peptides and manuscript write up, PS participated in the design of purification protocol, write up of the manuscript and provided the venom. CB conceived of the study, and participated in its design and coordination and helped to draft the manuscript. All authors read and approved the final manuscript.

Acknowledgement

AM acknowledges the University of Hamburg, Germany for supporting a research visit. C.B., A.M. and P.S. acknowledge support by DAAD via PROBAL Biotechnology Centre, Instituto de Pesquisas Energeticas e Nucleares, Avenida Lineu Prestes, 2242 São Paulo, Brazil.

Received: 9 July 2015 Accepted: 5 January 2016
Published online: 14 January 2016

References

1. Meier J, Stocker KF. Biology and distribution of venomous snakes of medical importance and the composition of snake venom. In: Meier JWI, editor. Handbook of clinical toxicology of animal venoms and proteins. Boca Raton, Florida: CRC Press; 1995. p. 367–412.

2. Georgieva DAR, Betzel C. Proteome analysis of snake venom toxins: pharmacological insights. Expert Rev Proteomics. 2008;5(6):787–97.

3. Kang TS, Georgieva D, Genov N, Murakami MT, Sinha M, Kumar RP, et al. Enzymatic toxins from snake venom: structural characterization and mechanism of catalysis. FEBS J. 2011;278(23):4543–76.

4. Wang Y, Hong J, Liu X, Yang H, Liu R, Wu J, et al. Snake cathelicidin from Bungarus fasciatus is a potent peptide antibiotics. PLoS One. 2008;3(9):e3217.

5. Galverte JI. Venomics: digging into the evolution of venomous systems and learning to twist nature to fight pathology. J Proteome. 2009;72(2):1211–6.

6. Sanchez EE, Rodriguez-Acosta A, Palomar R, Lucena SE, Bashir S, Soto JG, et al. Colombistatin: a disintegrin isolated from the venom of the South American snake (Bothrops colombiensis) that effectively inhibits platelet aggregation and SK-Mel-28 cell adhesion. Arch Toxicol. 2009;83(5):271–9.

7. Zhang Y, Wu J, Yu G, Zhen C, Zhou X, Zhu S, et al. A novel natriuretic peptide from the cobra venom. Toxicon. 2011;57(1):134–40.

8. Earl ST, Maci PP, de Jersey J, Lavin MF, Dixon J. Drug development from Australian elapid snake venoms and the Venomics pipeline of candidates for haemostasis: Textilinin-1 (Q8008), Haempatch (Q8009) and CoVase (V0801). Toxicon. 2012;59(4):456–63.

9. Pimenta AM, De Lima ME. Small peptides, big world: biotechnological potential in neglected bioactive peptides from arthropod venoms. J Pept Sci. 2005;11(11):670–4.

10. Munawar A, Trusch M, Georgieva D, Spencer P, Frochau V, Harder S, et al. Venom peptide analysis of Vipera ammodytes meridionalis (Viperinae) and Bothrops jararacussu (Crotalinae) demonstrates subfamily-specificity of the peptidome in the family Viperidae. Mol Biol Sys. 2011;72(2):3298–307.

11. Ondetti MA, Williams NJ, Sabo EF, Plussce J, Weaver ER, Kocy O. Angiotensin-converting enzyme inhibitors from the venom of Bothrops jararaca. Isolation, elucidation of structure, and synthesis. Biochemistry. 1971;10(22):4033–9.

12. Cushman DW, Ondetti MA. Design of angiotensin converting enzyme inhibitors. Nat Med. 1999;5(10):1110–3.

13. Hayashi MA, Camargo AC. The Bradykinin-potentiating peptides from venom gland and brain of Bothrops jararaca contain highly site specific inhibitors of the somatic angiotensin-converting enzyme. Toxicon. 2005;45(8):1163–70.

14. Wermelinger LS, Dutla DL, Oliveira-Canvalho AL, Soares MR, Bloch Jr C, Zingali RB. Fast analysis of low molecular mass compounds present in snake venom: identification of ten new pyroglutamate-containing peptides. Rapid Commun Mass Spectrom. 2005;19(12):1703–8.

15. Camargo AC, Lanzer D, Guerrero JR, Ferrareira SM, Bradykinin-potentiating peptides: beyond captoril. Toxicon. 2012;59(4):516–23.

16. Lopes DM, Junior NE, Costa PP, Martins PL, Santos CF, Carvalho ED, et al. A new structurally atypical bradykinin-potentiating peptide isolated from Crotalus durissus cascavela venom (South American rattlesnake). Toxicon. 2014;90:36–44.

17. Munawar A, Trusch M, Georgieva D, Hildebrand D, Kwiatkowski M, Behnken H, et al. Elapid snake venom analyses show the specificity of the peptide composition at the level of genera Naja and Notechis. Toxins. 2014;6(3):850.

18. Rioli V, Prezoto BC, Konno K, Melo RL, Klitzke CF, Ferro ES, et al. A novel bradykinin potentiating peptide isolated from Bothrops jararacussu venom using catalytically inactive oligopeptidase EP24. J Proteome. 2008;275(10):2442–54.

19. Guerrero JR, Lameu C, Oliveira EF, Klitzke CF, Melo RL, Linares E, et al. Argininosuccinate synthetase is a functional target for a snake venom anti-hypertensive peptide: role in arginine and nitric oxide production. J Biol Chem. 2009;284(30):20022–33.

20. Morais KL, Hayashi MA, Bruni FM, Lopes-Ferreira M, Camargo AC, Ulrich H, et al. Bj-PRO-5a, a natural angiotensin-converting enzyme inhibitor, promotes vasodilatation mediated by both bradykinin B2 and M1 muscarinic acetylcholine receptors. Biochem Pharmacol. 2011;81(6):736–42.

21. Paschoal JF, Yamauchi J, Miranda JR, Carretero G, Melo RL, Santos RA, et al. Insights into cardiovascular effects of proline-rich oligopeptide (Bj-PRO-10c) revealed by structure-activity analyses: dissociation of antihypertensive and bradycardic effects. Amino Acids. 2014;46(2):401–13.

22. Faria Ferreira LA, Calee MR A, Schrader M, Lebrun I, Habermehl G. Enzymatic toxins from snake venom: structural characterization and pharmacological insights. Expert Rev Proteomics. 2008;5(6):787–97.

23. Gomes CL, Konno K, Conceição IM, Lanzer D, Yamane Y, Prezoto BC, et al. Identification of novel bradykinin-potentiating...
peptides (BPPs) in the venom gland of a rattlesnake allowed the evaluation of the structure–function relationship of BPPs. Biochem Pharmacol. 2007;74(9):1350–60.

24. Kodama RT, Carvalho DC, Kuniyoshi AK, Kitano ES, Tashima AK, Barra BF et al. New proline-rich oligopeptides from the venom of African Adders: Insights into the hypertensive effect of the venoms. Biochimica et biophysica acta. 2015; doi:10.1016/j.bbagen.2015.02.005

25. Majumder K, Wu J. Molecular Targets of Antihypertensive Peptides: Understanding the Mechanisms of Action Based on the Pathophysiology of Hypertension. Int J Mol Sci. 2014;16(1):256–83.

26. Zhuo JL, Ferraò FM, Zheng Y, Li XC. New frontiers in the intrarenal Renin-Angiotensin system: a critical review of classical and new paradigms. Front Endocrinol. 2013;4:166.

27. Junqueira-de-Azevedo Ide L, Ho PL. A survey of gene expression and diversity in the venom glands of the pitviper snake Bothrops insularis through the generation of expressed sequence tags (ESTs). Gene. 2002;299(1-2):279–91.

28. Murayama N, Hayashi MA, Ohi H, Ferreira LA, Hermann WV, Saito H, et al. Cloning and sequence analysis of a Bothrops jararaca cDNA encoding a precursor of seven bradykinin-potentiating peptides and a C-type natriuretic peptide. Proc Natl Acad Sci U S A. 1997;94(4):1189–93.

29. Cortin A, Vieira CA, Giglio JR. Primary structure and biological activity of bradykinin potentiating peptides from Bothrops insularis snake venom. J Proteom Chem. 1990;2(2):221–7.

30. Souza GH, Catharino RR, Ila DR, Eberlin MN, Hyslop S. Peptide fingerprinting of snake venoms by direct infusion nano-electrospray ionization mass spectrometry: potential use in venom identification and taxonomy. J Mass Spectrom. 2008;43(5):594–9.

31. Cotton J, Hayashi MA, Cuniasse P, Vazeux G, Lanzer D, De Camargo AC, et al. Selective inhibition of the C-domain of angiotensin I converting enzyme by bradykinin potentiating peptides. Biochemistry. 2002;41(19):6065–71.

32. Lameu C, Ulrich H. Applications of Snake Venom Proline-Rich Oligopeptides (Bj-PROs) in Disease Conditions Resulting from Deficient Nitric Oxide Production. INTECH Open Access Publisher; Rijeka, Croatia 2013

33. Qassey G, Schwager SL, Sturrock ED, Isaac RE, Acharya KR. Molecular recognition and regulation of human angiotensin-I converting enzyme (ACE) activity by natural inhibitory peptides. Sci Rep. 2012;2:717.

34. Lanzer D, Konko K, Marques-Porto R, Vieira Portaro FC, Stocklin R, de Camargo AC M, et al. Identification of five new bradykinin potentiating peptides (BPPs) from Bothrops jararaca crude venom by using electrospray ionization tandem mass spectrometry after a two-step liquid chromatography. Peptides. 2004;25(7):1085–92.

35. Coutinho-Neto ACC, Souza GH, Zaqueo KD, Kayano AM, Silva RS, Zuliani JP, et al. ESI-MS/MS identification of a bradykinin-potentiating peptide from Amazon Bothrops atrox snake venom using a hybrid Qq-oToF mass spectrometer. Toxins. 2013;5(2):327–35.

36. Nunez AS, Roque AC. Studies on the molecular recognition between bioactive peptides and angiotensin-converting enzyme. J Mol Recognit. 2009;22(2):162–8.

37. Graham RL, Graham C, McClean S, Chen T, O’Rourke M, Hirst D, et al. Identification and functional analysis of a novel bradykinin inhibitory peptide in the venom of New World Crotalinae pit vipers. Biochem Biophys Res Commun. 2005;338(3):878–82.

38. Lomonte B, Tsai WC, Ureña-Díaz JM, Sanz LM-OD, Sánchez EE, Fry BGGJ, et al. Venomics of New World pit vipers: genus-wide comparisons of venom proteomes across Agkistrodon. J Proteome. 2014;16(96):103–16.

39. Carmona AK, Schwager SL, Juliano MA, Juliano L, Sturrock ED. A continuous flow ultra-performance liquid chromatography tandem mass spectrometry: potential use in venom identification and taxonomy. J Mass Spectrom. 2008;43(5):594–9.

40. Tashima AK, Zelanis A, Kitano ES, Lanzer D, Melo RL, Rioli V, et al. Peptidomics of three Bothrops snake venoms: insights into the molecular diversification of proteomes and peptidomes. Mol Cell Proteomics. 2012;11(11):1245–62.

41. Hayashi MA, Murbach AF, Lanzer D, Portaro FC, Prezotto BC, Fernandes BL, et al. The C-type natriuretic peptide precursor of snake brain contains highly specific inhibitors of the angiotensin-converting enzyme. J Neurochem. 2003;85(4):969–77.