A New Family of Small (4kDa) Neurotoxins from the Venoms of Spiders of the Genus Phoneutria

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INTRODUCTION

One of the most intriguing features of peptide toxins is that many of them have similar actions despite having dissimilar sequences and folds, whereas others have diverse effects even when having similar sequences or folds [1]. This poses the interesting question of which are the structural determinants of specific actions and the key amino acid residues (and their respective positions) that are crucial for the binding of a toxin to its target, such as an ion channel. One of the strategies to reveal the structural determinants of a certain action is to compare naturally occurring toxins with different structures but with similar effects. This comparison may provide clues as to the major characteristics required for a specific mode of action.

The venom of the very aggressive South American solitaire spider Phoneutria nigriventer (Keys.) has been extensively studied and contains a wealth of biologically active peptides and proteins. Many of them act on ion channels or membrane receptors of insects and mammals, and may serve as models for the development of new biologically active compounds with potential use in medicine or agriculture [2-4]. Excitatory symptoms such as salivation, lachrymation, priapism, convulsions, followed by flaccid or spastic paralysis and death occur following icv injection in mice [5]. The venom of Phoneutria nigriventer was fractionated into four groups of peptides with neurotoxic activity (denominated PhTx1, PhTx2, PhTx3 and PhTx4), that have different primary structures and pharmacological properties [6-10], revealing it as a rich source of bioactive peptides. PhTx1 was shown recently to have a voltage dependent inhibitory activity on Na+ channels [11]. Fraction PhTx2 is responsible for the venom’s prevailing excitatory symptoms by inhibiting Na+ channel inactivation. It is composed of 9 peptides, 8 of which with high similarity. Toxins PhTx2-2 and PhTx2-6 have been extensively studied and were shown to inhibit Na+ channel inactivation [12, 13] Fraction PhTx3 was shown to be a complex mixture of at least six different toxins with diverse pharmacological properties [9]. Most of these had their primary activity characterized. PhTx3-1, was shown to inhibit the A-type K+ current that is associated with Ca2+ oscillation and pacemaker activity [14]. PhTx3-3 is a Ca2+ channel inhibitor, with higher affinity to P/Q- and R-type channels [15-17], PhTx3-4, also referred to as α-photonetoxin IIIA, inhibits N and P/Q-type Ca2+ channels with high affinity, and L-type Ca2+ channels with lower affinity [18, 19]. PhTx3-6 inhibits Ca2+ channels with higher affinity for N- and R-type channels [20]. PhTx3-2 was shown to inhibit L-type Ca2+ channels, but possible activities on other types of Ca2+ channels have not been investigated yet [21]. PhTx4 is active on insects and has much weaker toxic effects on mice [10]. All these toxins are peptides with molecular masses of 3.5 to 9 kDa. Most of these toxins had their sequences determined chemically, and subsequently confirmed (in a few cases with small corrections) by the analysis of clones from cDNA libraries constructed using the venom gland of the spider [13, 14, 21-24].

Recently it has been demonstrated the presence of a highly complex pool of smaller (up to 2kDa) peptides that provoke contractions in the smooth muscle of guinea pig ileum. The amino acid sequences of 15 isoforms were determined by tandem mass spectrometry [25]. All of these molecules, which are structurally related to the tachykinin family of neuro-hormone peptides, possessed N-terminal residues, and low similarity with other neurotoxins. Three toxins caused moderate inhibition of L-type Ca2+ channels. The structure of toxin PRTx27C3 was modeled and compared with toxin ADO1. The importance of four residues is suggested.
pyroglutamate residues and exhibited evidence of other post-translational modifications such as proteolysis and C-terminal amidation.

In the present work we now describe another new family of small (4kDa) toxins obtained from Phoneutria nigriventer and from the venoms of the two related species of spiders Phoneutria reidyi and Phoneutria keyserlingi. These small proteins caused spastic paralysis and death when injected ivc in mice, and all but one have inhibitory effects on L-type currents of GH3 cells. Their primary structures show that they are unrelated to previously described toxins. We propose a molecular model for the most active of these toxins, PRTx27C3, and based on comparison with the sequence and structure of other toxins, the importance of the amino acid residues R6, K14, F26 and K18 is speculated.

**MATERIALS AND METHODS**

**Venoms**

Male and female specimens of the spiders Phoneutria nigriventer (Keys.) and Phoneutria keyserlingi were collected in the region of Santa Barbara and Mariana, respectively, both in the state of Minas Gerais (Brazil), and kept in the arachnidarium of the Fundação Ezequiel Dias (Belo Horizonte, Brazil). Venom from the live adult spiders was obtained by electrical stimulation of the fangs (5-12 μl/spider, 160 mg/ml), transferred to siliconized glass tubes in ice, diluted with the same volume of distilled water and centrifuged at 4000 x g to remove insoluble materials and cellular debris. The supernatant was lyophilized and stored at –18 ºC. The venom of the spider Phoneutria reidyi, which was collected from specimens captured in the Amazon region near the hydro-electric reservoirs at Tucurui (Pará), Samuel (Roraima) and Balbina (Amazonas), was a generous gift from the Butantan Institute (São Paulo, Brazil).

**Purification of Venom Proteins**

The venoms of all three species were processed in the same manner. Aliquots of 25-30 mg of lyophilized venom were dissolved in 2 ml of aqueous 0.1% TFA and centrifuged at 4000 x g for 10 min to remove insoluble materials. The brownish yellow supernatant was applied to a preparative reversed-phase column (2.2 x 25 cm Vydac C4) and subjected to ion-exchange HPLC were then dissolved in 2 ml of 10 mM sodium phosphate buffer pH 6.1, and subjected to ion-exchange FPLC on a column (6.4 mm x 30 mm) of Resource™ S (Amersham Pharmacia Biotech) equilibrated in the same buffer. After application of the sample, the column was initially washed with the starting buffer for 10 min and then eluted with a gradient of 0-0.5 M NaCl in the same buffer at a flow rate of 1 ml/min over a period of 45 min. In one experiment the material obtained from the reversed-phase step was fractionated on an anion exchange HPLC column (4.1 mm x 30 cm) of Synchronax AX-300 using a linear gradient of 0-0.5 M NaCl in 10 mM Tris-HCl buffer pH 8.6 at a flow rate of 1 ml/min. The fractions collected were desalted and further purified by reversed-phase HPLC on an analytical column (4.6 mm x 25 cm Vydac C18 218TP54) using a linear gradient of 0-40% acetonitrile in 0.1% (v/v) TFA over 60 min. The homogeneity of all fractions obtained was examined by mass spectrometry and propionic acid/urea PAGE using the method described by Chettibi and Lawrence [26]. The gel mixture contained 22.5 g of acrylamide monomer, 0.8 g of bis-acrylamide, 36 g of urea and 2 ml of propionic acid made up to 100 ml (final volume) with distilled water. To 60 ml of this solution was added 400 μl of 10% (w/v) ammonium persulphate solution and 50 μl of TEMED. Gels were run with 2% acetic acid in both anode and cathode compartments. Samples were prepared by addition of an equal volume of 30% sucrose solution containing neutral red marker dye. After running gels were stained with 0.1% Coomassie blue in methanol/water/acetic acid (50/50/7), and destained in methanol/acetic acid/water (5/7/100).

**Mass Spectrometry Analyses**

ES-Q-TOF mass spectrometry analyses were carried out using a Q-TOF Micro™ (Micromass, UK) equipped with an electrospray ionization source operated in the positive mode. Capillary voltage was 3000 V and sample cone voltages were 40-60 V. Mass spectrometer calibrations were made using sodium iodide with cesium iodide in 2000 Da range. Samples diluted in 50% acetonitrile/ 0.1% TFA were introduced using a syringe pump with flow rates of 5-10 μl/min. The spectrum used was the result from 20 scans (2.4 s) combined. Original data (m/z) were treated (base line subtraction, smoothing and centering) and transformed into a mass (Da) spectrum. Collision induced dissociation (MS/MS) was carried out using argon and collision energies in the range 30-45 V. Data analyses were carried out with MassLynx® 3.5 software.

**Bioassays**

The toxicity of each fraction was assayed by icv injection in albino mice (18-22 g). The animal assays were kept to a minimum and were carried out in accordance with ethical procedures of the Fundação Ezequiel Dias. The lyophilized samples were dissolved in 0.15 M saline, containing 0.25 mg/ml of BSA for icv injection of 5 μl samples, as described by Rezende et al. [7]. The appearance of neurotoxic symptoms (excitation, salivation, lachrymation, priapism, spastic or flaccid paralysis, scratching, tail elevation) or the death of animals was observed after injection. Control animals received physiological saline alone. All assays were carried out in duplicates.

**S-Reduction and Alkylation**

The proteins (20-200 nmol) were S-reduced and alkylated with 4-vinyl pyridine essentially as described by Hensch [27]. The material was dissolved in 1 ml of 6 M gua-
The temperature was set to 300 °K by means of the Berendcules [30] in a cubic box with periodic boundary conditions. The toxin was solvated by 1054 SPC water molecules in the GROMACS package [28], using the force field Gromos Molecular dynamics simulations were carried out with the GROMACS package [28]. One by one according to the primary structure of toxin A model for toxin PRTx27C3 was built by taking the model from the PDB and using the GROMOS force field. Molecular Modeling BLAST programs. SWISS-PROT/TREMBL data bases using the FASTA 3 and the BLAST programs. The amino acid sequence of the various proteins were compared with the sequences of other related proteins in the SWISS-PROT/TREMBL data bases using the FASTA 3 and BLAST programs.

**Determination of Amino Acid Sequences**

Samples (4-10 nmol) of the S-pyridyl-ethylated proteins were dissolved in 1 ml of 0.1 M ammonium bicarbonate pH 7.9 and digested with trypsin for 3.5 h using 2% (w/w) enzyme/substrate. After lyophilisation the peptides produced were separated by reversed-phase HPLC on a column (4.6 mm x 25 cm) of Vydac C18 (small pore, 201SP54) by using an extended gradient of 0 to 50% acetonitrile in 0.1% trifluoroacetic acid for 180 min at a flow rate of 1 ml/min. The collected proteins were lyophilized.

**Sequence Comparisons**

The amino acid sequence of the various proteins were compared with the sequences of other related proteins in the SWISS-PROT/TREMBL data bases using the FASTA 3 and BLAST programs.

**Molecular Modeling**

A model for toxin PRTx27C3 was built by taking the toxin ADO1 as a reference, whose residues were replaced one by one according to the primary structure of toxin PRTx27C3. The resulting structure was used as an initial configuration for the molecular dynamics simulation.

Molecular dynamics simulations were carried out with the GROMACS package [28], using the force field Gromos 96 [29]. The toxin was solvated by 1054 SPC water molecules [30] in a cubic box with periodic boundary conditions. The temperature was set to 300 °K by means of the Berendsen scheme [31] of coupling to an external bath with coupling constant of 0.1 ps. A constant pressure of 1 bar was applied also by means of the Berendsen algorithm [31]. Before the actual simulation, a steepest descend method was employed to remove close contacts followed by a 400 ps position restrained molecular dynamics to equilibrate the water molecules around the peptide, which was kept fixed. Finally, the actual simulation was performed for 5ns with a time step of 1 fs.

**Protein Nomenclature**

Each protein / peptide purified and studied during this work is described by a code. For example the species of Phoneutria from which it originated is indicated by PN= Phoneutria nigriventer; PR= Phoneutria reidyi, or PK= Phoneutria keyserlingi. The first number indicates the number of the peak eluted from the preparative reversed-phase HPLC on Vydac C4 that contained the protein. The following letter C indicates that the second step was cationic exchange FPLC (Resource S, pH 6.1). The subsequent number indicates the peak number during the second step.

**Electrophysiology**

GH3 cells were maintained in DMEM (Sigma Chemical Co., MO, USA) supplemented with Fetal Bovine Serum (Cultilab, Campinas, Brazil) 10%, penicillin/streptomycin 1% (Sigma Chemical Co., MO, USA) and incubated at 37 °C, CO2 5%. The cells were detached with a trypsin/EDTA solution and plated in Petri dishes for electrophysiological measurements. Ionic currents were recorded from GH3 cells by using the whole cell patch clamp technique. The data were acquired either with a HEKA/EPC9 amplifier controlled by Pulse 8.11 software (HEKA Elektronik Gmbh, Lambrecht, Germany), sampled at 10 kHz and filtered 3.33 kHz or with an Axopatch 200B amplifier controlled by Pclamp6 software (Axon Instruments, Union City, USA), sampled at 10 kHz and filtered at 5 kHz. The voltage protocol used to record the currents was: from a holding potential of -80 mV were applied 100 ms pulses from -100 mV to 50 mV (10 mV increments). To analyze the current as a function of the time, the test pulse was set at 0 mV, where the maximal currents were observed (not shown). The protocol was repeated each 5 s. Patch pipettes were pulled from soft glass capillaries with a tip resistance of 2-4 MΩ. The cells were sealed in a bath solution containing (in mM): NaCl 130, KCl 5, CaCl2 2, MgCl2 1, HEPES 10, Glucose 10, pH 7.4 with NaOH. The currents were recorded under continuous perfusion of a control solution containing (in mM): NaCl 100, BaCl2 25, HEPES 10, Glucose 10, KCl 5, Tetrodotoxin 0.0003, pH 7.4 with NaOH. Ba2+ was substituted for Ca2+ as the charge carrier because it is conducted better and prevents the Ca2+-dependent inactivation, thus facilitating the measurement of the currents. The pipettes were filled with a solution containing (in mM): CsCl 140, EGTA 5, HEPES 10, ATP-Mg 4, pH 7.2 with CsOH. The toxins were diluted to final concentrations of 200 nM or 1 μM in the control solution containing cytochrome C 4 μM to prevent adsorption [17]. Data were analyzed with PulseFit 8.11 software (HEKA Elektronik Gmbh, Lambrecht, Germany), Clampfit (Axon Instruments, Union City, USA) and SigmaPlot 5 (Jandel Inc., USA), being presented as mean ± SEM unless stated otherwise.

**RESULTS AND DISCUSSION**

The proteins described in this study were partially purified from the majority of the other venom components by subjecting the crude venoms of Phoneutria nigriventer, Phoneutria reidyi and Phoneutria keyserlingi to an initial step of reversed-phase HPLC on preparative columns of Vydac C4, by using extended gradients of acetonitrile in aqueous 0.1% TFA. The peaks collected at concentrations of 36-38% acetonitrile were numbered as PN26, PN27, PR27 and PK32, as shown in Fig. (1). After lyophilization, the contents of each fraction were examined by propionic acid/urea PAGE and by Q-TOF mass spectroscopy, which revealed that the major components in each fraction had a mass of approximately 4 kDa, but were contaminated with other larger proteins.
Figure 1. Profiles of the reversed-phase HPLC chromatography. Comparison of the preparative reversed-phase HPLC (Vydac C4) profiles obtained for the crude venoms of PN, *Phoneutria nigriventer*; PR, *Phoneutria reidi* and PK, *Phoneutria keyserlingi*. The amount of protein injected onto the column was 30mg for PN and PR and 15mg in the case of PK. The presence of peptides or proteins in the eluate was detected by measuring the UV absorption at 214nm. The void volume was 55 ml. All other experimental details are given in the text. The arrows indicate the peaks which were collected and further purified by ion-exchange FPLC.

Figure 2. Profiles of the cation exchange FPLC chromatography. Cation exchange FPLC chromatography of the fractions PN27, PN26A0, PK32 and PR27 obtained from the preparative reversed-phase HPLC. In the case of PN26A0, the fraction PN26 was subjected to an additional step of anion exchange FPLC on Synchropak AX-300 before being applied to the cation exchanger. The fractions were dissolved in 2ml of 10mM Na phosphate buffer pH 6.1 and applied to a column of Resource S equilibrated in the same buffer. After washing with the starting buffer for 10min the column was then eluted with a gradient of 0-0.5M NaCl in the same buffer at a flow rate of 1ml/min over a period of 45 min. The dotted line indicates the gradient concentration of NaCl. The arrows indicate the peaks which were selected for desalting/further purification by reversed-phase HPLC on analytical columns of Vydac C18.

The proteins in each of the fractions PN27, PR27 and PK32 were then subjected to ion exchange FPLC on Resource S at pH 6.1, using a gradient of 0-0.5 M NaCl, which gave a good resolution of the mixtures, as depicted in Fig. (2). The small 4 kDa proteins of interest were located in the peaks PN27C4, PR27C3 and PK32C4, all of which were eluted from the ion exchanger at a concentration of 0.2 M NaCl. A slightly different procedure was used for the material in PN26 which was initially fractionated on an anion exchange HPLC column of Synchropak AX-300 at pH 8.6, and eluted with a gradient of 0-0.5 M NaCl. The majority of the protein in the mixture failed to bind to the column and was collected in the first peak PN26A0. The contents of this fraction were desalted, lyophilized and applied to FPLC on the Resource S cation exchanger at pH 6.1. This step yielded three peaks, the third of which, PN26A0C3, contained the 4 kDa protein. Each of the four fractions PN26A0C3, PN27C4, PR27C3 and PK32C4 obtained from the ion exchange chromatographies were then desalted and further purified by reversed-phase FPLC on an analytical column of Vydac18.
All four fractions gave a single sharp band when examined by propionic acid/urea PAGE (not shown). The homogeneity of the proteins in the fractions PN26A0C3, PR27C3 and PK32C4 was confirmed by the results of Q-TOF mass spectroscopy, which indicated that their respective molecular masses were 4058.07 ± 0.45, 4043.64 ± 0.22 and 3996.78 ± 0.03. In the case of PN27C4, two molecular mass values of 4057.34 ± 0.25 and 4023.45 ± 0.14 were obtained, which suggested the presence of two very similar isoforms, and this was subsequently confirmed by amino acid sequencing which revealed the presence of phenylalanine and leucine in the cycle corresponding to position 7.

The amino acid sequences of these toxins, determined by automated sequencing of the intact native proteins, the S-pyridyl-ethylated forms, were confirmed by sequencing the peptides derived from them by digestion with trypsin. The pyridyl-ethylated forms, were confirmed by sequencing the automated sequencing of the intact native proteins, the S-terminally inserted to align the 6th cysteine residue. The shaded residues are equal to the corresponding amino acid in PRTx27C3.

Although other proteins of Phoneutria venom also produce spastic paralysis [5], the toxins reported in the present paper are not related to them. The alignment between the primary sequences on Fig. (3) shows that there is a significant identity between the assassin bug peptide and the toxins studied in this present work. Due to the scarce yield of our material, preliminary electrophysiological assays were carried out on L-type Ca²⁺ channels of GH3 cells. These channels have important physiological role in the control of blood pressure, heart beating, endocrine secretion, and modulation of neurotransmitter release. Fast inactivating LVA and non-inactivating HVA Ba²⁺ currents, which permeate through Ca²⁺ channels, have been reported in GH3 cells [38]. Although Ca,2,1 (P/Q-type) encoding mRNA has been detected, nearly all HVA currents are carried through dihydropyridine-sensitive L-type channels [17, 39]. GH3 cells also express Na⁺ and K⁺ channels [14, 40]. In contrast with what could be expected from the mice symptoms, there was no significant effect of these neurotoxins on either Na⁺ or K⁺ currents of these cells (not shown).

Fig. (4) shows the effect of PRTx27C3 (200 nM and 1 μM) on Ba²⁺ currents. In the presence of PRTx27C3 (200 nM) non-inactivating currents were inhibited by 18% and there was a significant decrease (50%) of the non-inactivating currents, followed by a partial recovery, when the final concentration was 1 μM. Fig. (4A) and Fig. (4B) show the time-course of the toxin effect at 200 nM and 1 μM concentrations, respectively (n=3). The maximal inhibition was observed about 20 s after the addition, and the effect was not reversed upon washing. The rate of inhibition by the toxin was not affected by the frequency of stimulation (not shown).

In the presence of PNTx27C4 (1 μM) Ba²⁺ currents were inhibited by 16%. Fig. (4C) shows the time-course of PNTx27C4 effect (n=3). Similar to PRTx27C3, the maximal effect of PNTx27C4 was reached few seconds after the addition, and no immediate washout was observed. Fig. (4D) shows that in the presence of PNTx26A0C3 (1 μM) there

![Figure 3](image-url)
was a slow and small inhibition (15%) of the Ba$^{2+}$ current ($n=3$), and no immediate washout was observed. Finally, Ba$^{2+}$ currents were not significantly modified in the presence of PKTx32C4 (not shown).

Of the toxins with significant degree of identity with PRTx27C3, only two had their structures resolved so far: ADO1 [41] and Agelenin [42], PDB codes 1LMR and 2E2S, respectively. They display common features: both have compact core structures stabilized with three disulphide bridges, four loops and a short two stranded antiparallel sheet near the C-terminal. Both belong to the inhibitory cysteine knot (ICK) structural family. Albeit agelenin is derived from a spider, it has characteristics that make it less suitable for our modeling: (i) contrary to ADO1, it is not active in mammalian Ca$^{2+}$ channels in its native form [42], (ii) its sequence contains less identical residues than ADO1, as shown in Fig. (3), (iii) the amino acid residues reported to be important for agelenin activity are predominantly absent from PRTx27C3 [42]. Therefore, we used the structure of ADO1 as the initial template for modeling PRTx27C3, as described under Material and Methods. The modeled structure is depicted in Fig. (5). The Ramachandran plot statistics of this model, not considering the glycine residues, showed 61% of the dihedral angles laying in the most favorable regions, and 39% in the additional favorable regions. No residue was found in the disallowed region (not shown). Bernard et al. [41], by comparing the three-dimensional structure of ADO1 with already known structures of other Ca$^{2+}$ channels toxins, have observed conserved positioning of the amino acid residues R8, K11, K17, R31 and F27/Y28, and suggested that they might be significant for ADO1 activity. Fig. (3) shows that, of these proposed amino acids, R8 is well conserved in equivalent positions in the present family of peptides. The automatic superimposition of the model of PRTx27C3 and ADO1 structure, using Swiss-Pdb Viewer fitting routine, reveals close proximity of large stretches of the toxin backbones, mainly on the intermediate loop of ADO1, as shown in Fig. (5). This is the most conserved region of these molecules, as shown in Fig. (3). Interestingly, this structural alignment shows that three of the residues considered important for ADO1 activity, namely R8, K17 and F27, closely superimpose with the amino acid residues R6, K14 and F26 of PRTx27C3, respectively. Moreover, R6 and K14 are conserved in all toxins described in the present work. F26 is present only in PRTx27C3, and is replaced by less bulky hydrophobic residues in the other toxins, which have lower affinities. These amino acid residues are shown in Fig. (5), where they are displayed as sticks. One feature called our attention: Lys21 of ADO1 and Lys18 of PRTx27C3 protrude from the molecule at the same position. This residue is also present in PNTx27C4 and PNTx26An0C3, but is absent from PKTx32C4, which is devoid of the inhibitory activity on L-type Ba$^{2+}$ currents. There is some controversy on the effects of ADO1. Whilst it

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**Figure 4.** Effects of the purified toxins on L-type Ba$^{2+}$ currents. (A) Time course of the effect of PRTx27C3 (200nM) in the steady-state L-type Ba$^{2+}$ currents (mean $\pm$ SEM, $n=3$) recorded at 0mV. (B) Same as (A), but with 1μM of PRTx27C3. (C) Time course of the effect of PNTx27C4 (1μM) in the steady-state L-type Ba$^{2+}$ currents (mean $\pm$ SEM, $n=3$) recorded at 0mV. (D) Time course of the effect of PNTx26A0C3 (1μM) in the steady-state L-type Ba$^{2+}$ currents (mean $\pm$ SEM, $n=3$) recorded at 0mV. In all graphs the solid line shows the time interval for which the toxin was applied and the inserts show representative traces recorded at 0mV in the absence and in the presence of the toxin under investigation.
is reported as an L-type Ca\textsuperscript{2+} channel inhibitor on the PDB data bank, it is assumed in the literature as having no significant effect on L-type Ca\textsuperscript{2+} currents for its similarity with Ptu1 toxin, which is devoid of this effect. However, to our knowledge, while effect on P/Q-type mammalian Ca\textsuperscript{2+} channels has been shown, no direct determination of the effect of ADO1 on L-type channels has been reported in the literature. On the other hand, we cannot exclude the possibility that the toxins described in the present work have more prominent effect on other types of Ca\textsuperscript{2+} channels. The limited amount of pure material prevented us from testing this possibility.

Lysine residues have been shown to be crucial for inhibition K\textsuperscript{+} and Ca\textsuperscript{2+} channels in other toxins. Furthermore, the motif –C-C-K– seen in these toxins is also present in other 13 toxins, including the L-type calcium channel blockers calciceptin and FS2, toxins from the black mamba snake. Some of these toxins act on other ion channels, notably K\textsuperscript{+} channels. It has been pointed out the presence of conserved amino acid residues on Ca\textsuperscript{2+} and K\textsuperscript{+} channels, which explained the similar effects of gramotoxin and hanatoxin, respectively, on these channels [43]. As seen in both our model of PRTx27C3 and in the structure of ADO1, this motif places the lysine residue in a fixed position, whereas its side chain is very exposed and with high flexibility. Interestingly, a similar sequence (-C-C-R-) exists in \omega-conotoxin GVIA. Alanine substitution showed that the arginine residue of this sequence (Arg17) is important for the activity of the toxin, but not for its binding [44]. We suggest that the motif CCK is important for the activity of the present toxins.

It should be noted that there is some controversy over the exact taxonomic position of *Phoneutria keyserlingi*. Eickstedt [45] distinguished *Phoneutria nigriventer* from *Phoneutria keyserlingi* by the morphological characteristics of larger epigynal lateral guides and less curved embolus of the latter species. However Simo and Brescovit [46] consider that these differences only represent intraspecific variability, and suggest that *Phoneutria keyserlingi* should be regarded as a synonym of *Phoneutria nigriventer*. The results of this present biochemical investigation in which the amino acid sequences of three isoforms of proteins isolated from the venom of PN can be compared with single isoforms from PK.
and PR, show that whilst there are strong similarities between them, they are distinct molecules.

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ABBREVIATIONS

ATP = Adenosine 5’-triphosphate

EDTA = Ethylenediaminetetraacetic acid

EGTA = [Ethylene bis-(oxyethylenenitrilo)]tetraacetic acid

ICV = Intracerebroventricular

PAGE = Polyacrylamide gel electrophoresis

PK = Phoneutria keyserlingi

PN = Phoneutria nigriventer

PR = Phoneutria reidyi

SEM = Standard error of the mean

SPC = Simple Point Charge

PAGE = Polyacrylamide gel electrophoresis

UV = Ultra violet

REFERENCES


