Research Article

Chemical Modifications of PhTX-I Myotoxin from Porthidium hyoprora Snake Venom: Effects on Structural, Enzymatic, and Pharmacological Properties

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We recently described the isolation of a basic PLA2 (PhTX-I) from Porthidium hyoprora snake venom. This toxin exhibits high catalytic activity, induces in vivo myotoxicity, moderates footpad edema, and causes in vitro neuromuscular blockade. Here, we describe the chemical modifications of specific amino acid residues (His, Tyr, Lys, and Trp), performed in PhTX-I, to study their effects on the structural, enzymatic, and pharmacological properties of this myotoxin. After chemical treatment, a single His, 4 Tyr, 7 Lys, and one Trp residues were modified. The secondary structure of the protein remained unchanged as measured by circular dichroism; however other results indicated the critical role played by Lys and Tyr residues in myotoxic, neurotoxic activities and mainly in the cytotoxicity displayed by PhTX-I. His residue and therefore catalytic activity of PhTX-I are relevant for edematogenic, neurotoxic, and myotoxic effects, but not for its cytotoxic activity. This dissociation observed between enzymatic activity and some pharmacological effects suggests that other molecular regions distinct from the catalytic site may also play a role in the toxic activities exerted by this myotoxin. Our observations supported the hypothesis that both the catalytic sites as the hypothetical pharmacological sites are relevant to the pharmacological profile of PhTX-I.

1. Introduction

Phospholipase A2 (PLA2; EC 3.1.1.4) enzymes catalyse the hydrolysis of acyl ester bond of 1,2-diacyl-3-sn-phosphoglycerides at the sn-2 position, with a requirement for a Ca2+ [1]. PLA2 enzymes from snake venom are quite fascinating from both biological and structural point of view. Despite their structure being conserved, they exhibit a wide range of pharmacological activities, including neurotoxicity [2], myotoxicity [3, 4], and cardiotoxicity [5] as well as anticoagulant, hemolytic [6], antiplatelet [7], hypotensive [8], hemorrhagic [9], and edema inducing effects [10]. However, not all the PLA2 enzymes induce all these pharmacological effects. In general, an individual PLA2 exhibits one or more specific pharmacological effects [11, 12].

Within the family Viperidae, two distinct types of venom PLA2 molecules have been described, all of them sharing a high degree of homology both in primary and three-dimensional structure [13]: the “classical” PLA2, that present an invariant Asp49 residue that plays a key role in catalysis; and the “PLA2 homologues,” devoid of enzymatic activity, that present the substitution of Asp49 by Lys49 or, less frequently, by Ser, Arg, Gln, or Asn [14, 15]. Despite their difference in catalysis, both the Asp49 and the Lys49 proteins are
able to induce various pharmacological effects [16]. Thus the structure-function relationship among this group of proteins is subtle and complicated [1]. In some cases, the pharmacological effects result from their enzymatic activities, probably through the action of the products of hydrolysis, lysophospholipids, and fatty acids, that alter cell membrane shape and permeability [17, 18] but for many of them, the pharmacological effects are independent of their enzymatic activities, such as Lys49 PLA2 myotoxins, which lack hydrolytic activity and therefore act via another mechanism, which is only partially understood. A site close to the C-terminus, comprising a variable combination of basic and hydrophobic amino acids, has been identified as being responsible for toxicity [19].

In a previous work, we showed that Porthidium hyopora snake venom is a rich source of PLA2 enzymes. Additionally, we purified a myotoxic PLA2 (PhTX-I) to homogeneity in reverse-phase HPLC, which constitutes of a single polypeptidic chain, has a molecular mass of 14,249 Da, and whose amino acid sequence exhibits high identity with other polypeptidic chain, has a molecular mass of 14,249 Da, and whose amino acid sequence exhibits high identity with other myotoxic Asp49 PLA2 [20]. We also demonstrated that PhTX-I (20 μg/mL) caused edema, in vivo creatine kinase release, C2C12 skeletal muscle myoblasts cytotoxicity, and neuromuscular blockade of chick biventer cervicis muscle preparations. However, it is still unknown whether these pharmacological effects were mediated by the phospholipase catalytic activity of PhTX-I or not. One of the strategies employed for the elucidation of the relationship between catalytic activity and pharmacological effects of PLA2 is based on the chemical modification of specific residues in these enzymes [21]. Using this approach, a dissociation of pharmacological effects and enzymatic activity for various PLA2 has been observed, suggesting the presence of separate enzymatic and pharmacological active site(s) contained within their amino acid sequences [22, 23]. In the present study, we investigated the effects of chemical modifications of specific amino acid residues (His, Tyr, Lys, and Trp), performed in PhTX-I, on their enzymatic, structural, and pharmacological properties.

2. Material and Methods

2.1. Reagents. 2,4′-Dibromoacetophenone (BPB), 4-nitrobenzenesulfonyl fluoride (NBSF), 4-nitrobenzenesulfonyl chloride (NPSC), 4-nitro-3-(octanoyloxy) benzoic acid, and other reagents were from Sigma Chemical Co. (St. Louis, MO, USA); acetic anhydride (AA) was from Merck.

2.2. Purification of PhTX-I. The PhTX-I PLA2 from Porthidium hyopora venom was purified by reverse phase HPLC [20]. Briefly, 5 mg of whole venom was dissolved in 200 μL of buffer A (0.1% TFA) and centrifuged at 4500 g; the supernatant was then applied to a μ-Bondapak C18 column (0.78 × 30 cm; Waters 991-PDA system), previously equilibrated with buffer A for 15 min. The elution of the protein was then conducted using a linear gradient (0%–100%, v/v) of buffer B (66.5% Acetonitrile in buffer A) at a constant flow rate of 1.0 mL/min. The chromatographic run was monitored at 280 nm of absorbance, and after elution the fraction was lyophilized and stored at –40°C.

2.3. Chemical Modifications. Modification of His residues with 2,4′-Dibromoacetophenone (BPB) was carried out as previously described [21]. Briefly, 3 mg of PhTX-I were dissolved in 1 mL of 0.1 M Tris-HCl containing 0.7 mM EDTA (pH 8.0) and 150 μL of BPB (1.5 mg/mL, in ethanol), and the mixture incubated for 24 h at 25°C. Modification of Lys residues with acetic anhydride (AA) was performed at a protein: reagent molar ratio of 1: 50 [21]. PhTX-I (3 mg) was dissolved in 1.5 mL of 0.2 M Tris-HCl buffer at pH 8.0, and 10 μL of AA was added and the mixture was incubated for 1 h at 25°C. Tyr residues were modified by treatment with 4-nitrobenzenesulphonyl fluoride (NBSF) as previously described [24]. Briefly, 1 μmol of PhTX-I (10 μmol of Tyr) was dissolved in 14 mL of 0.1 M Tris-HCl (pH 8.0) and incubated with 10 μmol, of NBSF for 20 h at 25°C. Modification of Trp residues was performed according to Takasaki et al. [25]. Briefly, 9 mg of PhTX-I were dissolved in 4 mL 50% acetic acid containing 1 mg of 2-nitrobenzenesulfonyl chloride (NPSC) and incubated for 1 h at 25°C. In all cases, excess reagent was removed by ultrafiltration through a Millipore’s Amicon Ultra-15 membrane and washed with distilled water, followed by lyophilization.

2.4. Amino Acid Analysis. Amino acid analysis was performed on a Pico-Tag Analyzer (Waters Systems) as described by Heinrikson and Meredith [26]. Native PhTX-I PLA2, and their modified derived samples (30 μg) were hydrolyzed at 105°C for 24 h, in 6 M HCl (Pierce sequencing grade) containing 1% phenol (w/v). The hydrolysates were reacted with 20 μL of derivatization solution (ethanol: triethylamine: water: phenylisothiocyanate, 7:1:1:1, v/v) for 1 h at room temperature, after which the PTC amino acids were identified and quantified by HPLC, by comparing their retention times and peak areas with those from a standard amino acid mixture.

2.5. Electrophoresis. Native PhTX-I PLA2, and their modified derivatives were examined by Tricine SDS-PAGE in a discontinuous gel and buffer system, under reducing and nonreducing conditions [27]. The molecular mass markers used were (in kDa): phosphorylase B—97.0, albumin—66.0, ovalbumin—45.0, carbonic anhydrase—30.0, soybean trypsin inhibitor—20.1, and α-lactalbumin—14.4.

2.6. Mass Spectrometry. An aliquot (4.5 μL) of the modified proteins was injected in C18 (100 μm × 100 mm) RP-UPLC (nanoAcquity UPLC, Waters) coupled with nanoelectrospray tandem mass spectrometry on a Q-Tof Ultima API mass spectrometer (MicroMass/Waters) at a flow rate of 600 nL/min. The gradient was 0%–50% acetonitrile in 0.1% formic acid over 45 min. The instrument was operated in MS continuum mode, and the data acquisition was from m/z 100–3000 at a scan rate of 1 s and an interscan delay of 0.1 s. The spectra were accumulated over about 300 scans and the multiple charged data by the mass spectrometer on
the m/z scale were converted to the mass (molecular weight) scale using maximum entropy-based software supplied with Masslynx 4.1 software package. The processing parameters were output mass range 6.000–20.000 Da at a “resolution” of 0.1 Da/channel; the simulated isotope pattern model was used with the spectrum blur width parameter set to 0.2 Da; the minimum intensity ratios between successive peaks were 20% (left and right). The deconvoluted spectrum was then smoothed (2 x 3 channels, Savitzky Golay smooth) and the mass centroid values obtained using 80% of the peak top and a minimum peak width at half height of 4 channels.

2.7. Circular Dichroism. Circular dichroism (CD) spectra of native PhTX-I PLA2 and their modified derivatives were recorded with a JASCO model J-720-ORD 306 spectropolarimeter equipped with a thermoelectric sample temperature controller (Peltier system) following standard procedures previously described [28]. After centrifugation at 4000 g for 5 min, samples (1–4 μM protein in 10 mM sodium phosphate, pH 8) were transferred to a 10-mm path-length quartz cuvette. Circular dichroism spectra in the wavelength range 260 to 200 nm were collected, using a bandwidth of 0.1 nm and a response time of 1 s. Data collection was performed at 25 °C with 50 nm/min scanning speed. At least ten scans were accumulated for each sample, and all spectra were corrected by subtraction of buffer blanks. The estimation of secondary structure elements was performed using the CDNN Deconvolution software (version 2.1), and Origin 7.5 (OriginLab) was used for graphics and analysis.

2.8. PLA2 Activity. PLA2 activity was measured using the assay described by Cho and Kezdy [29] and Holzer and Mackessy [30] modified for 96-well plates. The standard assay mixture contained 200 μL of buffer (10 mM Tris–HCl, 10 mM CaCl2, and 100 mM NaCl, pH 8.0), 20 μL of substrate 4-nitro-3-(octanoyloxy) benzoic acid (3 mM), 20 μL of water, and 20 μL of PhTX-I PLA2 or their modified derivatives (1 mg/mL) in a final volume of 260 μL. After adding proteins (20 μg) the mixture was incubated for up to 40 min at 37 °C, measuring absorbance at intervals of 10 min. The enzyme activity, expressed as the initial velocity of the reaction (V0), was calculated based on the increase of absorbance after 20 min. All assays were done in triplicate, and the absorbances at 425 nm were measured with a VersaMax 190 multiwell plate reader (Molecular Devices, Sunnyvale, CA).

2.9. Inhibition. The inhibitory effects of EDTA or low molecular weight heparin from porcine intestinal (Mr 6,000 Da) on pharmacological and enzymatic activities of PhTX-I were assessed by incubating the enzyme with 1 mM solution of this chelating agent or a heparin:toxin molar ratio of 2:1 for 30 min at 37 °C. The inhibition of PLA2 activity of PhTX-I by crotapotins F2 and F3 from Crotalus durissus colubritae also was evaluated by incubating the two proteins (1:1, w/w) for 30 min at 37 °C and then assaying the residual enzyme activity.

2.10. Chick Biventer Cervicis Muscle Preparation (BCP). Animals were anesthetized with halothane and sacrificed by exsanguination. The biventer cervicis muscles were removed and mounted under a tension of 0.5 g, in a 5 mL organ bath (automatic organ multiple-bath LE01 Letica Scientific Instruments. Barcelona, Spain) at 37 °C containing aerated (95% O2-5% CO2) Krebs solution (pH 7.5) of the following composition (mM): NaCl 118.70, KCl 4.70, CaCl2 1.88, KH2PO4 1.17, MgSO4 1.17, NaHCO3 25.00, and glucose 11.65. Contracture to exogenously applied acetylcholine (ACH; 110 μM for 60 s) and KCl (20 mM for 130 s) was obtained in the absence of field stimulation, before and after the addition of a single dose of PhTX-I PLA2 (1.4 μM) or their modified derivatives (1.4 μM). A bipolar platinum ring electrode was placed around the tendon, which runs the nerve trunk supplying the muscle. Indirect stimulation was performed with a (MAIN BOX LE 12404 Panlab s.l. Powerlab AD Instruments Barcelona, Spain) stimulator (0.1 Hz, 0.2 ms, 3-4 V). Muscle contractions and contractures were isometrically recorded by force-displacement transducers (Model MLT0201 Force transducer 5 mg–25 g Panlab s.l. AD Instruments Pty Ltd. Spain) connected to a PowerLab/4SP (OUAD Bridge AD Instruments, Spain).

2.11. Myotoxic Activity. Groups of five Swiss mice (18–20 g) received an intramuscular injection (i.m.) of 20 μg of PhTX-I PLA2 or their modified derivatives dissolved in 100 μL of PBS, in the gastrocnemius. A control group received 100 μL of PBS (0.12 M NaCl, 0.04 M sodium phosphate, pH 7.2). Three hours after injection, blood was collected from the tail into heparinized capillary tubes, and the plasma creatine kinase (CK; EC 2.7.3.2) activity was determined by a kinetic assay (Sigma 47-UV). Activity was expressed in U/L, one unit defined as the phosphorylation of 1 μmol of creatine/min at 25 °C.

2.12. Edema-Forming Activity. The ability of PhTX-I PLA2 and their modified derivatives to induce edema was studied in groups of five Swiss mice (18–20 g). Fifty μL of phosphate-buffered saline (PBS; 0.12 M NaCl, 0.04 M sodium phosphate, pH 7.2) with toxins (1 μg/paw) were injected in the subplantar region of the right footpad. The left footpad received 50 μL of PBS, as a control. The paw volume was evaluated plethysmographically (Model 7140 Plethysmometer, Ugo Basile, USA), immediately before the injection (basal) and after 1 h. Edema-forming activity was expressed as the percentage of the increase in volume of the right footpad in comparison to the left foot pad (control). The equation for calculation of the percentage of edema in toxins injected paw was

\[
\%\text{edema} = \left( \frac{T_x \times 100}{T_0} \right) - 100, \tag{1}
\]

where \( T_x \) is the edema (volume) measured at each time interval and \( T_0 \) is the volume of the paw (intact, zero time before toxins injection). The percentage of edema calculated was subtracted from the matched values at each time point in the saline injected hind paw (control).
Table 1: Amino acid composition of native PhTX-I and their chemically modified derivatives.

| Amino acid | PhTX-I native | PhTX-I modified | PhTX-I modified | PhTX-I modified | PhTX-I modified |
|------------|---------------|----------------|----------------|----------------|----------------|
| Asp        | 11            | 10.92 (11)     | 10.9 (11)      | 10.72 (11)     | 10.56 (11)     |
| Glu        | 6             | 5.98 (6)       | 6.12 (6)       | 5.92 (6)       | 6.37 (6)       |
| Ser        | 3             | 3.37 (3)       | 3.15 (3)       | 3.21 (3)       | 3.03 (3)       |
| Gly        | 11            | 11.43 (11)     | 11.25 (11)     | 11.39 (11)     | 11.13 (11)     |
| His        | 2             | 1.74 (2)       | 1.94 (2)       | 1.6 (2)        | 0.65 (1)       |
| Arg        | 7             | 7.62 (7)       | 6.56 (7)       | 7.15 (7)       | 7.42 (7)       |
| Thr        | 6             | 6.34 (6)       | 6.09 (6)       | 6.35 (6)       | 6.34 (6)       |
| Ala        | 5             | 4.81 (5)       | 5.29 (5)       | 5.34 (5)       | 5.44 (5)       |
| Pro        | 6             | 5.82 (6)       | 6.01 (6)       | 6.03 (6)       | 5.74 (6)       |
| Tyr        | 10            | 10.24 (10)     | 9.79 (10)      | 5.94 (6)       | 10.21 (10)     |
| Val        | 3             | 3.33 (3)       | 3.34 (3)       | 3.49 (3)       | 3.49 (3)       |
| Met        | 1             | 1.47 (1)       | 1.17 (1)       | 1.18 (1)       | 1.45 (1)       |
| Cys        | 14            | 13.96 (14)     | 13.98 (14)     | 13.74 (14)     | 13.69 (14)     |
| Ile        | 2             | 2.47 (2)       | 2.35 (2)       | 2.27 (2)       | 2.22 (2)       |
| Leu        | 8             | 8.29 (8)       | 8.44 (8)       | 8.12 (8)       | 8.01 (8)       |
| Phe        | 4             | 3.58 (4)       | 4.49 (4)       | 4.74 (4)       | 4.21 (4)       |
| Lys        | 17            | 10.02 (10)     | 16.93 (17)     | 17.35 (17)     | 17.2 (17)      |
| Trp        | 3             | ND             | ND             | ND             | ND             |

*aAmino acid sequence. Amino acid analysis of chemical modifications with bAA (Lys), cNPSC (Trp), dNBSF (Tyr), and eBPB (His). ND: nondetermined.

2.13. Cytotoxic Activity

2.13.1. Maintenance of NG97 and NCIH-3T3 Cell Culture. Cytotoxicity was assayed on NG97 and NCIH-3T3 cells, which were grown in plastic flasks (25 cm²) with RPMI 1640 medium (Cultilab, Campinas, SP, Brazil), supplemented with 2% L-glutamine, 120 μg/mL garamycin, and 13% inactivated fetal bovine serum (complete medium). The cultures were incubated at 37°C in an atmosphere containing 5% CO₂. Medium was changed every 48 h, and when the culture reached confluence, the subculture was performed by treatment with trypsin and versene (Adolfo Lutz, São Paulo, SP, Brazil). Variable amounts of native PhTX-I and chemically modified derivatives were diluted in assay medium and added to cells in 96-well plates. Experiments were carried out in triplicate.

2.13.2. Cellular Viability by Neutral Red Uptake Assay. This assay was done according to the method described by Borenfreund and Borrero [31]. After a 4 h incubation with serum-free medium containing 50 μg of neutral red/mL the cells were washed quickly with PBS and then 0.1 mL of a solution of 1% (v/v) acetic acid: 50% (v/v) ethanol was added to each well to extract the dye. After shaking for 10 minutes on a microtitre plate shaker, the absorbance was read at 540 nm. The cell death was determined in comparison of the absorbance obtained from nontreated cells.

2.14. Statistical Analyses. Results were reported as mean ± SEM. The significance of differences among means was assessed by analysis of variance followed by Dunnett’s test, when several experimental groups were compared with the control group. Differences were considered statistically significant if P < 0.05.

3. Results

Table 1 shows the result of analysis of amino acid composition of the modified proteins of PhTX-I PLA₂ compared to the amino acid sequence of native PhTX-I. After chemical treatment a single His, 4 Tyr, and 7 Lys residues were modified by BPB, NBSF, and AA, respectively. With the methodology employed it was not possible to determine the changes in Trp residues. In all these cases the exact position of the groups modified is unknown. The homogeneity of the modified proteins was evaluated by SDS-PAGE, as shown in Figure 1. Both native PhTX-I and modified forms migrated similarly by electrophoresis, indicating the presence of a single band electrophoretic for each protein, demonstrating the high homogeneity of the samples as well as no detectable change in the apparent molecular weight of the modified forms of the PhTX-I.

Molecular mass values of the modified proteins were determined by ESI mass spectrometry. Figures 2(a), 2(b), 2(c), and 2(d) show the mass spectrum of modified forms of PhTX-I, by BPB, AA, NPSC, and NBSF, respectively; each peak represents the protein carrying a different number of charges (protons). Figures inserted in each spectrum show the deconvolution of the spectra obtained of the modified proteins. The modified protein in His, Lys, Trp, and Tyr residues had molecular mass of 14440.7, 14537.9, 14470.1, and 15068.9 Da, respectively.

To determine whether the chemical modifications in PhTX-I caused changes in protein secondary structure, far-UV circular dichroism (CD) was used. Figure 3 shows...
CD spectra of native PhTX-I and their modified forms which demonstrate two negative bands of similar magnitude (−11,000 to −10,500 deg·cm²·dmol⁻¹) at 208 and 222 nm and a positive one at −190 nm (data not shown), indicating a consistent content of α-helical structures. The exception was NBSF modified PhTX-I that had lower signal at 222 nm than the unmodified protein. Based on the CDNN program analysis of the native PhTX-I spectrum, the contents of α-helices, β-sheets, and β-turns were 32%, 18%, and 17%, respectively. The contents of secondary structure of PhTX-I chemically modified by BPB, NPSC, or AA were very similar to native PhTX-I, with no significant changes in the CD spectrum between them, suggesting that secondary structure of protein remains practically unchanged. However, NBSF treatment resulted in a significant change (−10,500 to −9,000 deg·cm²·dmol⁻¹) in molar ellipticity of the negative band at 222 nm, which by CDNN software analysis indicates an altered content of α-helices, β-sheets, and β-turns (30%, 19% and 18%, resp.).

The catalytic activity of native PhTX-I and their modifications were studied using the chromogenic substrate 4-nitro-3-[(octanoyloxy) benzoic acid. The catalytic activity of native PhTX-I was almost completely abolished by BPB, but only partially reduced after modification of Tyr or Lys residues; NPSC did not cause a significant decrease in this activity (Figure 4). The incubation of native PhTX-I with crotopotins F2 and F3 from C. d. collinetaus and EDTA diminished the activity; heparin did not significantly inhibit the catalytic activity (Figure 4).

Figure 5 shows the graphical representation of the blockade of the contractile response in the neuromuscular transmission (BCP) of native PhTX-I and modified derivatives. The native PhTX-I at a concentration of 1.4 μM blocked the indirectly evoked contractions reaching 50% of the block in about 20 min. This activity was markedly diminished for all the PhTX-I chemically modified derivatives, except for modification of Trp where no significant change in force of contraction was produced by the modification.

Figure 6 shows the effect of the different chemical modifications on the myotoxic activities of PhTX-I, by time-course measurement of plasma levels CK after intramuscular injection of proteins. It can be seen that even when BPB and AA were the most effective reagents altering this activity, most of the treatments had some effect. Three hours after treatment started, the myotoxic effect was reduced by 85%, 80%, and 72% by treatment with AA, BPB, and NBSF, respectively. EDTA and heparin inhibited around 74% and 50% of this activity.

Regarding the edema-inducing effect, after alkylation with BPB, PhTX-I lost around 70% of its activity one hour after treatment, while modification of Lys and Tyr residues caused only a partial decrease of this activity (55% and 40%, resp.); however, NPSC did not inhibit this activity, as can be seen in Figure 7.

Native PhTX-I was cytotoxic to NIH-3T3 fibroblast cell line and to a lesser extent for the NG97 cell line derived from a human astrocytoma grade III (Figure 8). The cytotoxic activity of PhTX-I was independent of enzymatic activity, since BPB-treated PhTX-I was able to produce cytotoxicity in both cell types. After treatment with NPSC the cytotoxic activity was partially reduced. On the other hand, acetylation and sulfonylation of Lys and Tyr residues, respectively, reduced strongly the cytotoxic activity in both cell types.

### 4. Discussion

We recently described the isolation of a basic PLA₂ (PhTX-I) from *P. hyoprora* using reverse phase HPLC [20]. This toxin exhibits high catalytic activity, shares various structural similarities with other "bothropic" PLA₂, and has the conserved and essential Asp residue in position 49. PhTX-I induces *in vivo* myotoxicity, moderates footpad edema (at concentrations up to 10 μg/mL and 0.5 μg/mL, resp.), and causes *in vitro* neuromuscular blockade in chick biventer cervicis muscle preparations (at concentrations of 1.4 μM). Here, we have described the chemical modifications of specific amino acid residues (His, Tyr, Lys, and Trp), performed in PhTX-I and how theses modifications affected the structural, enzymatic, and pharmacological properties of this myotoxin.

A first important observation was that after chemical treatment a single His, 4 Tyr, 7 Lys, and one Trp residues were modified by BPB, NBSF, AA, and NPSC, respectively (Table 1). These results were confirmed by mass spectrometry (Figure 2). The mass of native PhTX-I, 14249.22 Da [20], after treatment with NBSF (15068.96 Da) increased 819.74 Da (Figure 2(d)), demonstrating modification of fourth Tyr residues (up to 187.16 Da in each residue corresponding to reagent NBS that would be incorporated). Similarly, only three Tyr residues (7, 70, and 77) with the highest exposed surface areas in the notoxin were modified by NBSF, suggesting that the seven remaining residues are “buried” within the molecule [32]. Acetylation of Lys residues of basic PLA₂ myotoxins three (PrTX-I, -III and BnSP-7) caused a complete loss of basicity, being demonstrated by electrophoretic analysis [22, 33]; however AA-treated PhTX-I (14537.9 Da) (Figure 2(b)) shows that only seven Lys residues were modified, due to which there is an increase of 288.60 Da, equivalent to seven times the mass of acetyl radical (42 Da) incorporated into
the amino group of K residues, showing similar behavior to the native protein in electrophoresis gel (Figure 1).

Similarly, an increase of 221.39 Da in NPSC-treated PhTX-I (Figure 2(c)) indicated a modification of a single Trp residue. By analogy with other results using PrTX-I and -III from *B. pirajai* [22], we would expect that, with the three Trp residues present in the structure of PhTX-I, this reagent should modify the residue with a larger area of exposure; it would be more easily attacked by the NPSC. Alkylation of His by BPB has been widely used to assess the role of enzymatic activity in the pharmacological actions of PLA$_2$ [22, 23, 33–36]. PhTX alkylated with BPB had a molecular mass of 14440.7 Da (Figure 2(a)), which confirmed the modification of only one residue of His. His48 is a highly conserved amino acid residue in PLA$_2$, which has a vital role in catalysis [37]. Since the enzymatic activity of the PhTX-I was almost completely abolished after this modification, His48 was likely the residue modified, because this amino acid is part of the catalytic triad of this protein family.

Examination of native PhTX-I by CD spectroscopy indicated that the predominant secondary structure of this PLA$_2$ consisted of alpha-helices (Figure 3), in agreement with the results obtained for others as PLA$_2$ from Taiwan cobra [38], PLA2A from *C. d. ruruima* [39], and BnIV from *B. newidii* [40]. The secondary structure of PhTX-I chemically modified derivatives did not alter significantly after modifications as evidenced by the CD spectra, which exhibited almost the same profile as that of native toxin. Only NBSF-treated toxin revealed detectable changes in secondary structure composition when compared to native toxin (Figure 3). This difference could be due to partial helix unfolding but the small change suggested that such unfolding was minimal. Kini [1] described that the alkylation by BPB does not affect the three-dimensional structure of PLA$_2$ or its ability to bind phospholipids, but may alter the ability to interact with specific proteins or ligands. Thus, it is suggested that the chemical modifications performed in this study mainly affected the specific residues involved in such modifications and did not result in drastic conformational changes in the molecule.

The PhTX-I PLA$_2$ is a Ca$^{2+}$-dependent enzyme, with maximum activity at pH 8 and 40°C, reaching $V_{\text{max}}$ and $K_{\text{m}}$ of 11.76 nmoles/min and 1.96 mM, respectively [20]. Heparin slightly decreased enzymatic activity of PhTX-I, (25%) (Figure 4); similarly, this polyanionic compound resulted acting as negative allosteric modulator of PLA$_2$ *C. d. cascadella*.
Wavelength (nm)
- 10500
- 9000
- 7500
- 6000
- 4500
- 3000
- 1500
0

Figure 3: Far-UV circular dichroism spectra of native PhTX-I and their chemically modified derivatives. Native PhTX-I (continuous lines) and their chemically modified derivatives (symbols): NBSF-treated PhTX-I (circles), BPB-treated PhTX-I (inverted triangles), NPSC-treated PhTX-I (open triangles), and AA-treated PhTX-I (squares).

Figure 4: PLA₂ activity of native PhTX-I PLA₂ and their chemically modified derivatives with BPB, AA, NBSF, NPSC and inhibitory effect of heparin, EDTA, and crotapotins (F2 and F3) upon 4-nitro-3-(octanoyloxy) benzoic acid substrate *(P < 0.05).

Figure 5: Neuromuscular blockade of chick biventer cervicis preparations incubated at 37°C with native PhTX-I and their chemically modified derivatives with BPB, AA, NBSF, and NPSC, all in concentration 1.4 µM. The points are the mean ± SEM of six experiments. *(P < 0.05) compared to the twitch-tension before toxin addition.

Figure 6: Increments in plasma CK activity after intramuscular injection of native PhTX-I their chemically modified derivatives with BPB, AA, NBSF, and NPSC (all 20 µg/100 µL) and incubation of PhTX-I (20 µg) with heparin and EDTA. Controls were injected with 100 µL of PBS. Three hours after injection, blood was collected, and serum levels were measured. Values are mean ± SEM of five mice at each time point.

On the other hand, EDTA greatly decreased catalytic activity of PhTX-I (88%), as β-bungarotoxin and notexin, which were inhibited by EDTA even in the presence of an excess of Ca²⁺ [42]. The F2 and F3 crotapotins from C. d. collilineatus significantly inhibited the enzymatic activity of PhTX-I at approximately 55% (Figure 4), in agreement with BjIV PLA₂ of B. jararacussu, which was inhibited by 50% in its catalytic activity by F7, F3, and F4 crotapotins from C. d. terrificus, C. d. collilineatus, and C. d. cascavella [43]. These results suggest that crotapotins can bind to bothropics PLA₂ in a manner similar to that of crotalics PLA₂, and raise the possibility that bothropic venoms may contain crotapotin-like proteins which inhibit the catalytic activity of PLA₂.

Acetylation of Lys residues significantly reduced the enzymatic activity; however, a residual activity was detected,
corresponding to 26% (Figure 4), similarly to MT-III (B. asper) and MT-I (B. godmani) [21]. The mode of specific acetylation is not clear, but there is some evidence of reduction of the calcium-binding capacity, thereby reducing the enzymatic activity of PLA2 [44]. In contrast, both myotoxic and cytotoxic effects were totally abolished, whereas a residual edematogenic effect remained (Figures 6, 7, and 8). These observations agree with previous studies in which the Edema (%)

His48 is a highly conserved residue in PLA2, which has an important role in catalysis [37]. Alkylation of His by BPB has been widely used to assess the role of enzymatic activity in the pharmacological actions of PLA2 [21, 35, 36, 49]. Here, alkylation of His at the active site of PhTX-I markedly abolished enzymatic activity (<4% residual activity) (Figure 4). Others have reported residual enzymatic activity following alkylation of His48, as βBuTX and nontoxin from N. n. nigricollis and N. n. atra PLA2, with values close to that found for BPB-PhTX-I (5%–8% of residual activity) [34]. Myotoxic, neurotoxic, and edema-forming activities of PhTX-I, were drastically reduced by this modification (Figures 5, 6, and 7); EDTA by treatment also affected myotoxicity and edematogenic activity (Figures 6 and 7), strengthening the hypothesis that phospholipid enzymatic hydrolysis is involved in these effects. Similarly, neurotoxic and myotoxic activities were inhibited almost completely after alkylation of His48 of PLA2 Basp-III (B. asper), PrTX-III (B. pirajai), BthTX-II (B. jararacussu), Cdc-9, and Cdc-10 (C. d. cumanesia) [21, 22, 35, 49], showing that these pharmacological effects are dependent on the catalytic activity. Since alkylation of the active site His48 completely abolished the catalytic activity and strongly attenuated these three pharmacological effects, Kini [1] suggested the hypothesis that PLA2 activity potentiates these pharmacological effects induced by Bothrops and Crotalus myotoxins.

In contrast, cytotoxic activity upon NG97 and NCIH-3T3 cells not was affected by His modification (Figure 8), suggesting that enzymatic activity is not required for this effects and that there are other molecular regions involved in cellular membrane perturbation. Our results agree with MTX-I and II PLA2 from B. brazili, which displayed cytotoxic activity against Jurkat lines independently of catalytic activity [50]. Some authors propose that cytotoxic activity on tumor cell lines is associated with apoptosis induction, considering the fact that PLA2 enzymes have been proposed to play a role in mediating apoptosis in various models, including cell lines [51]. The PLA2 activity is proposed to accelerate turnover of phospholipids, which may influence membrane changes that occur during apoptosis [52]. We suggested important role of Lys residues in cytotoxic effect, because PhTX-I treatment with AA abolished this activity.

Studies have been directed trying to understand the mechanisms involved in the inflammatory response induced by myotoxic PLA2 from several snake venoms [53–55]. However, the relationship between enzymatic activity and edema is contradictory [56]. It is assumed that myotoxic and edematogenic activities can be induced by different structural domains in these PLA2, or that a partial overlapping of these domains occurs [55, 57].

Residual enzymatic activity after sulfonylation of Tyr residues was 38% (Figure 4). Tyr52 and Tyr73 are part of the catalytic site of PLA2; giving structural support to stabilize the catalytic system [37], changes in this system would affect the enzymatic activity. Tyr-modified PhTX-I decreased myotoxic and neurotoxic activities more than the enzymatic activity (Figures 4, 5, and 6), once again indicating the dissociation between enzymatic and pharmacological activities.
Cytotoxic activity of PhTX-I also was reduced drastically after modification by NBSF (Figure 8), indicating that Tyr residues would also be involved in this process. Zhao et al. [58] observed that myotoxic PLA

2 have a set of Tyr residues located at the C-terminal region of the molecule. These Tyr may contribute to the hydrophobic-cationic combination proposed to play a role in myotoxicity and cytotoxicity [19, 46, 59]. PhTX-I show a Tyr residue in C-terminal region; alterations of this acid amino in this region of the molecule are causing reduction of toxicity of PhTX-I. However, the influence of conformational changes induced by NBSF in these effects cannot be discarded (Figure 3).

NPSC-treated PhTX-I did not significantly decrease enzymatic activity (Figure 4), suggesting that the modified Trp residue is not related to the catalytic system. In the same way, this modification showed no changes as compared to native PhTX-I, in edematogenic and cytotoxic activities (Figures 7 and 8); the myotoxic effect was minimally affected (Figure 6). In this sense Trp residues of PhTX-I have little or no direct action on the muscle. The Trp modifications of PhTX-I also maintained the action upon blockade of the contractile response in chick biventer cervicis muscle preparation (Figure 5). In contrast, modified Trp affected only the neurotoxic effect caused by MjTX PLA2-II [60], indicating the relevant role of this residue in this activity and suggesting that chemical modification could be interfering with the stability of the interaction between the monomers of this dimeric toxin, since Trp77 helps to maintain the homodimeric interaction. This suggests that the shift from dimeric to monomeric form of myotoxin may reduce the ability to affect the plasma membrane [60]. Because PhTX-I is a monomeric toxin, modified Trp would not affect the pharmacological activity of this toxin.

5. Conclusions

Our results indicate the critical role played by Lys and Tyr residues in myotoxic, neurotoxic activities and mainly in the cytotoxicity displayed by PhTX-I. His residue and, therefore, catalytic activity of PhTX-I are relevant for edematogenic, neurotoxic, and myotoxic effects, but not for its cytotoxic activity. Our observations supported the existence of pharmacological sites, distinct from the catalytic site, that contribute to the development of toxicity of these toxins and the hypothesis that the catalytic activity would potentiate the myotoxic and neurotoxic effects induced by snake venom PLA

2. Finally, although a partial dissociation is shown, both the catalytic sites as the hypothetical pharmacological sites are relevant to the pharmacological profile of PhTX-I.

Abbreviations

PLA

2: Phospholipase A

2

PhTX-I: Porthidium hyoprora myotoxin-I
EDTA: Ethylenediaminetetraacetic acid
BPB: 2,4'-Dibromoacetophenone
NBSF: 4-Nitrobenzenesulfonil fluoride
AA: Acetic anhydride
NPSC: 2-Nitrobenzenesulfonyl chloride
CD: Circular dichroism
PBS: Phosphate-buffer saline
CK: Creatine kinase.

Ethical Approval

The animals and research protocols used in this study followed the guidelines of the Ethical Committee for use of animals of ECAE-IB-UNICAMP SP, Brazil (protocol number...
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