ACP-TX-I and ACP-TX-II, Two Novel Phospholipases A2 Isolated from Trans-Pecos Copperhead Agkistrodon contortrix pictigaster Venom: Biochemical and Functional Characterization

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Abstract: This work reports the purification and biochemical and functional characterization of ACP-TX-I and ACP-TX-II, two phospholipases A2 (PLA2) from Agkistrodon contortrix pictigaster venom. Both PLA2s were highly purified by a single chromatographic step on a C18 reverse phase HPLC column. Various peptide sequences from these two toxins showed similarity to those of other PLA2 toxins from viperid snake venoms. ACP-TX-I belongs to the catalytically inactive K49 PLA2 class, while ACP-TX-II is a D49 PLA2, and is enzymatically active. ACP-TX-I PLA2 is monomeric, which results in markedly diminished myotoxic and inflammatory activities when compared with dimeric K49 PLA2s, confirming the hypothesis that dimeric structure contributes heavily to the profound myotoxicity of the most active viperid K49 PLA2s. ACP-TX-II exhibits the main pharmacological actions reported for this protein family, including in vivo local myotoxicity, edema-forming activity, and in vitro cytotoxicity. ACP-TX-I PLA2 is cytotoxic to A549 lung carcinoma cells, indicating that cytotoxicity to these tumor cells does not require enzymatic activity.

Keywords: snake venom; Agkistrodon contortrix pictigaster; D49 PLA2; homologous K49 PLA2; myotoxin; edema-forming activity and cytotoxicity

Key Contribution: This study reports the first isolation and functional characterization of two basic PLA2 from A. c. pictigaster venom.

1. Introduction

Phospholipases A2 (PLA2), which hydrolyze 2-acyl ester bonds of 3-sn-phospholipids, releasing lysophospholipids and fatty acids, are widespread in snake venoms, facilitating the immobilization and digestion of prey [1,2]. Venom PLA2s belong to the secretory PLA2 (sPLA2) family (groups IA, elapids and IIA, viperids) [3,4]. PLA2s are small proteins (13–15 kDa) with 115–122 residues, and seven conserved disulfide bonds [5,6]. They are subdivided into two main subgroups: (1) D49 PLA2s, which have an aspartate residue at position 49, and typically have high catalytic activity [7,8], and (2) K49 PLA2s, with a lysine residue at position 49. These have little or no catalytic activity, but still induce various biological effects [9,10]. Both types of proteins show significant similarity in their three-dimensional structures,
although they exhibit different pharmacological actions, such as myotoxicity, neurotoxicity, anticoagulant activity, platelet aggregation inhibition/activation, hemolysis, edematogenicity, hypotension, bactericidal action, proinflammatory cytotoxicity, and antitumor activity [2,11].

A great number of sequences and crystal structures of viperid PLA\(_2\)s have been determined [11–13]. Despite their overarching similarities, sequence differences result in diverse biological functions [14]. Accordingly, characterization of individual toxins is important to better understand the pathophysiology of envenomation and to potentially improve therapeutic procedures [15].

\textit{Agkistrodon} is a genus of pit vipers ranging from the southern United States to northern Costa Rica [16]. Currently, this genus comprises four species: \textit{A. contortrix} (copperheads), \textit{A. piscivorius} (cottonmouth), \textit{A. bilineatus} (cantils), and \textit{A. taylori} (Taylor’s cantils) [17,18]. Copperheads comprise several subspecies: \textit{A. c. contortrix} (southern copperhead), \textit{A. c. latincinctus} (broad-banded copperhead), \textit{A. c. mokasen} (northern copperhead), \textit{A. c. phaeogaster} (Osage copperhead), and \textit{A. c. pictigaster} (Trans-Pecos copperhead). Subspecific taxonomy is based largely on gross morphology, color pattern, and scale counts [18]. Cottonmouths and copperheads are among the most common venomous snakes in the southeastern United States. Cottonmouths frequent streams, rivers, ponds, marshes, and swamps, whereas copperheads are found in deciduous hardwood forests with moist leaf litter, large logs, scattered rocks, and high levels of vegetative cover. These snakes account for \(\sim\)30\% of the non-lethal human envenomations in this region [19,20].

The Trans-pecos copperhead (\textit{A. c. pictagaster}) is found in western Texas, northern Chihuahua, and Coahuila (Mexico) [16]. Juveniles usually prey on invertebrates (spiders, millipedes, and insects), frogs, and small lizards, whereas adults primarily prey on vertebrates, including amphibians (salamanders and anurans), reptiles (lizards and snakes), birds, and small mammals (rodents) [16].

Studies on the biochemical composition and toxic activities of copperhead venoms, including D49 PLA\(_2\) and homologous K49 PLA\(_2\), have been almost exclusively restricted to \textit{A. c. contortrix} and \textit{A. c. latincinctus}, [5,21–23]. In contrast, little information is available on \textit{A. c. pictigaster}. Partial characterizations of two disintegrins from this venom have been described by Lucena et al. [24]. A comparison of venom proteome variation in the genus \textit{Agkistrodon} found that \textit{A. c. pictigaster} venom contains ten protein families, dominated by PLA\(_2\)s (38.2\%) and metalloproteinases (30.2\%). The venom showed proteolytic, hemorrhagic, and myotoxic activities [25].

This work is the first report of two basic PLA\(_2\)s isolated from \textit{A. c. pictigaster} venom, with their identification and structural characterization found by biochemical and enzymatic experiments. Furthermore, we describe their biological activities and cytotoxic properties upon an A549 tumor cell line. The results of this study illuminate structure-function relationships of ACP-TX-I and ACP-TX-II PLA\(_2\), and improve our understanding of the chemistry of this venom.

2. Results

2.1. Purification and Biochemical Characterization of ACP-TX-I and ACP-TX-II

Chromatographic separation of \textit{A. c. pictigaster} venom by reversed phase high-performance liquid chromatography (RP-HPLC) on a C\(_{18}\) column resulted in 29 fractions, with two prominent peaks (16 and 18) eluting in more than 30\% acetonitrile (Figure 1). These peaks, representing about 30\% of total venom protein, were collected and screened for PLA\(_2\) activity. Fractions 16 and 18 were named ACP-TX-I and ACP-TX-II, respectively. Both ACP-TX-I and ACP-TX-II exhibited high purity when re-chromatographed using the same chromatography system, each showing only one peak (Figure S1). These peaks were also analyzed by SDS-PAGE, which manifested a single electrophoretic band with an \(Mr\) of approximately 14 kDa under reducing and non-reducing conditions (Figure 1 insert).

ESI-MS analysis demonstrated that the proteins were homogeneous, with molecular masses of 12,209.7 and 14,041.1 Da for ACP-TX-I and ACP-TX-II, respectively (Figure 2A,B).
2.2. Determination of the Amino Acid Sequences of ACP-TX-I and ACP-TX-II

In order to identify the purified proteins, they were digested with trypsin, and tryptic peptides were detected and characterized by mass spectrometry. Amino acid sequences of several tryptic peptides were obtained (Table 1). ACP-TX-I and ACP-TX-II shared 7 and 6 peptides with other viperid PLA\(_2\)s, respectively.

Assembly of partial protein sequences by similarity, using BLAST and multiple alignment, demonstrated that ACP-TX-I is a K49 PLA\(_2\) (Figure 3A), and is quite similar to well-characterized members of this family such as ACL PLA\(_2\) from A. c. lacintus, MjTX-II from Bothrops moojeni, BnSP-7 from B. neuwiedi pauloensis, etc. Sequenced peptides accounted for 64 amino acids, assuming that the number of residues is identical to that of homologous viperid toxins. This represented an estimated 53% of the protein (Figure 3A). ACP-TX-II shares conserved domain sequences common to catalytically active D49 PLA\(_2\)s. Peptide 1, having the sequence DATDRCCFVHDCCYQ/Q/K, contains an Asp (aspartic acid) residue that corresponds to position 49 in the complete amino acid sequence (Figure 3B).

Table 1. Tryptic peptides of ACP-TX-I and ACP-TX-II PLA\(_2\). Peptides were separated and sequenced by mass spectrometry. Molecular masses are monoisotopic.

| Peptides | Mass (Da) Expected | Amino Acid Sequence | Mass (Da) Calculated |
|----------|--------------------|---------------------|----------------------|
| **ACP-TX-I** | | | |
| 1 | 9.864.857 | GQ/KPK/QDATDR | 9.864.781 |
| 2 | 14.075.661 | DATDRCCFVHQ/K | 14.076.024 |
| 3 | 7.753.587 | VTGCDPK | 7.753.535 |
| 4 | 14.737.322 | AI/I/LCEEK/QNPCI/Q/K | 14.737.319 |
| 5 | 17.537.551 | MCECDK/QAVAI/CLC/IRE | 17.537.619 |
| 6 | 11.235.574 | ENL/IDTYNQ/Q/K | 11.235.509 |
| 7 | 9.845.405 | TYWK/QYPQ/K | 9.845.069 |
| **ACP-TX-II** | | | |
| 1 | 20.627.936 | DATDRCCFVHDCCYQ/G/K | 20.627.754 |
| 2 | 15.045.434 | CCFVHDCYQ/G/K | 15.045.356 |
| 3 | 22.618.772 | CCFVHDCYQ/G/Q/LTACSPQ/K | 22.619.149 |
| 4 | 17.687.631 | Q/KI/LCECDRAAAI/LCFS | 17.687.807 |
| 5 | 10.494.556 | DNI/I/I/LTYSQ/K | 10.494.666 |
| 6 | 9.845.087 | TYWKYPQ/K | 9.845.069 |
Figure 2. Molecular mass determination of ACP-TX-I (A) and ACP-TX-II (B) by nanoelectrospray tandem mass spectrometry using a Quadrupole Time-of-flight (Q-Tof) Ultima API mass spectrometer (MicroMass/Waters) with an output mass range of 6000–20,000 Da at a “resolution” of 0.1 Da/channel. Raw and deconvoluted electrospray mass spectra are shown (inserts).
Figure 3. ACP-TX-I (A) and ACP-TX-II (B) show significant similarity to K49 and D49 PLA2s, respectively (Edit Seq version 5.01© Program, DNASTAR Inc., Madison, WI, USA, 2001). ACL from A. c. laticinctus [26]; MjTX-II from B. moojeni [27]; BnSP-7 from B. neuwiedi pauloensis [28]; blK PLA2 from B. leucurus [29]; BbTX-II from B. brazili [6]; AP PLA2 from A. piscivorus [30]; APP PLA2 from A. p. piscivorus [31]; Pe PLA2 from Protobothrops elegans [32]; Ahp and BA2 PLA2 from A. halys pallas [33,34]. 

Hyphens indicate gaps generated by the alignment software.

2.3. Activity Measurements of ACP-TX-II

ACP-TX-I did not show PLA2 activity, but possessed a mass of ~14 kDa. ACP-TX-II displayed specific PLA2 activity of 29.31 ± 1.62 nmol/min (Figure 4A). The pH optimum was 8.0 (Figure 4B) and this protein was stable at temperatures from 35 to 40 °C (Figure 4D). At low concentrations, ACP-TX-II showed a sigmoidal relationship with temperature (Figure 4C) and a strict dependence on calcium ions (10 mM) for full activity. Substitution of Ca2+ with Mg2+, Mn2+, Cd2+, or Zn2+ (10 mM) significantly reduced enzyme activity (Figure 4E). Enzymatic activity of ACP-TX-II was abolished by EDTA and treatment with p-BPB. Incubation with crotapotins, F5 and F6, from C. d. collilineatus venom also diminished activity, while heparin did not significantly inhibit catalysis (Figure 4F).
isolation and characterization of two basic PLA2s, ACP-TX-I K49 and ACP-TX-II D49, the main components of this venom (Figure 1). Using C18 reversed phase HPLC, 29 peaks were obtained. The two most prominent peaks, 16 and 18, named ACP-TX-I and ACP-TX-II, respectively, were selected because of PLA2 activity and/or molecular mass in SDS-PAGE (Figure 1 insert). Multiple PLA2 isoforms in the same venom were derived from accelerated microevolution, in which high mutation rates in gene coding regions, mainly associated with amino acids exposed to the solvent, allowed development of new functions [6,14].

Figure 4. (A) PLA2 activity of A. c. pictigaster venom, ACP-TX-I and ACP-TX-II PLA2; (B) Effect of pH on the PLA2 activity of ACP-TX-II; (C) Effect of substrate concentration on the PLA2 activity of ACP-TX-II; (D) Effect of temperature on the PLA2 activity of ACP-TX-II; (E) Influence of ions (10 mM each) on PLA2 activity of ACP-TX-II; (F) Effect of heparin, EDTA crotapotins (F5 and F6) and chemical modification with BPB on PLA2 activity of ACP-TX-II. The results are the mean ± SEM of three determinations (* p < 0.05).

2.4. Pharmacological Activities of ACP-TX-I and ACP-TX-II

In vivo, ACP-TX-II PLA2 (20 and 50 µg), injected intramuscularly (IM), induced local myonecrosis, and time-course analysis showed a maximum increase in plasma CK 3 h after injection, returning to normal by 24 h (Figure 5B). In contrast, ACP-TX-I PLA2 showed no local myotoxic effect, even at high concentrations (Figure 5A). ACP-TX-I and ACP-TX-II PLA2 did not show systemic myotoxicity after intravenous (IV) injection (Figure S2).
ACP-TX-I PLA₂ had little edematogenic effect, since a 50-µg injection was necessary in order to observe 23.4% edema after 3 h (Figure 6A). In contrast, ACP-TX-II PLA₂ presented marked paw edema, with maximal activity (40.54%) 3 h after a 50-µg injection. Edema returned to normal levels after 24 h, showing dose-dependent activity.

ACP-TX-I was cytotoxic to cultured NIH/3T3 (non-tumor fibroblasts) and A549 (human lung carcinoma) cells treated with different concentrations of ACP-TX-I and ACP-TX-II PLA₂ (5–500 µg/mL) during a period of 24 h (Figure 7). ACP-TX-I PLA₂ cytotoxicity was dose-dependent on both cell types, causing a 50% decrease in cell viability at doses ≥20 µg/mL (Figure 7A). ACP-TX-II PLA₂ did not show toxicity in non-tumor cells used in this study, and 250–500 µg were required to decrease A549 tumor cell viability by 20 to 25% compared to controls, but the difference was not statistically significant (Figure 7B).

**Figure 5.** Myotoxic activity of ACP-TX-I PLA₂ (A) and ACP-TX-II PLA₂ (B) in mice. Time-course of the increments in plasma CK activity after an intramuscular injection of 20 and 50 µg of toxins compared to the injection of vehicle alone (PBS). Points represent means ± SD of four mice per group.

**Figure 6.** Edema-forming activity of ACP-TX-I PLA₂ (A) and ACP-TX-II PLA₂ (B) in mice. Induction of edema by toxins (10, 20 and 50 µg), injected SC in the footpads of mice. At various time intervals, the increase in footpad volume, compared to controls, was expressed as percent edema. Each point represents the mean ± SD of four animals per group.
Figure 7. In vitro cytotoxic activity of ACP-TX-I PLA₂ (A) and ACP-TX-II PLA₂ (B) in NIH/3T3 (non-tumor fibroblasts) and A549 (human lung carcinoma) cells. Cell viability (%) was estimated by neutral red uptake assay. Experiments were performed in triplicate (* p < 0.05).

3. Discussion

Crude venom of the Trans-Pecos copperhead, A. c. pictigaster, was fractionated by HPLC. Fractions of interest were analyzed by MS and screened for diverse bioactivities. This work reports isolation and characterization of two basic PLA₂s, ACP-TX-I K49 and ACP-TX-II D49, the main components of this venom (Figure 1). Using C₁₈ reversed phase HPLC, 29 peaks were obtained. The two most prominent peaks, 16 and 18, named ACP-TX-I and ACP-TX-II, respectively, were selected because of PLA₂ activity and/or molecular mass in SDS-PAGE (Figure 1 insert). Multiple PLA₂ isoforms in the same venom were derived from accelerated microevolution, in which high mutation rates in gene coding regions, mainly associated with amino acids exposed to the solvent, allowed development of new functions [6,14].

Both ACP-TX-I and ACP-TX-II were obtained in high purity. Re-chromatography using the same chromatography system showed only one peak for each fraction (Figure S1). On SDS-PAGE, a single band was seen with an Mr of approximately 14 kDa under reducing and non-reducing conditions (Figure 1 insert). Using this purification method, several other PLA₂ (Bp13 PLA₂, PhTX-I, -II, III, Bleu-PLA₂, Bbil-TX, BbTX-II, -III, etc.) from different venoms have been purified, showing that it is rapid and efficient for the purification of these proteins in one step [6–8,35–38].

Molecular masses determined for ACP-TX-I (12,209.7 Da) and ACP-TX-II (14,041.1 Da), obtained with ESI mass spectrometry (Figure 2), are very close to those of PLA₂s isolated from other snake venoms [26,34]. Partial amino acid sequencing of ACP-TX-I suggested that this protein probably has a Lys residue at position 49 (Figure 3A), based upon its high similarity to well characterized K49 PLA₂s such as ACL, MjTX-II, BnSP-7, bi-K, and BbTX-II from A. c. laticinctus, B. moojeni, B. neuwiedi pauloensis, B. leucurus, and B. brazili, [6,26–29]. In addition, ACP-TX-I showed negligible catalytic activity compared to crude venom and ACP-TX-II (Figure 4A).

In contrast to various homologous K49 PLA₂s, ACP-TX-I migrated as a monomer in SDS-PAGE when analyzed under reducing and non-reducing conditions (Figure 1 insert). Recently, it was demonstrated that SDS induces oligomerization of the Lys49 PLA₂, BPI, from Proteobothrops flavoviridis [39], although under the same conditions, ACP-TX-I behaves as a monomer. K49 PLA₂s isolated from three cottonmouths (A. p. piscivorus, A. p. leucostoma, and A. p. conanti) also behaved as monomers [14].

Partial amino acid sequencing of ACP-TX-II PLA₂ showed that it belongs to the D49 PLA₂ family, with an Asp residue at position 49 (Figure 3B). The catalytic site formed by H48, D49, Y52, and D89 is conserved, as reflected by the high enzymatic activity of the toxin (Figure 4A). Comparison of the amino acid sequence of ACP-TX-II shows a high degree of homology with other myotoxic D49 PLA₂ from viperid venoms (Figure 3B). It was not possible to determine the amino acid sequence
corresponding to the Ca\(^{2+}\)-binding loop, comprising of residues Y24 to G35, and containing glycine residues at positions 26, 30, 32, and 33, and cysteine residues at positions 27 and 29 [40]. This domain is important to maintain Ca\(^{2+}\) in the correct position for a nucleophilic attack on the substrate.

Optimum enzymatic activity of ACP-TX-II PLA\(_2\) occurred at pH 8 and 37 °C (Figure 4B,D), and Ca\(^{2+}\) was an obligatory co-factor [38,41]. Ca\(^{2+}\) replacement with other divalent ions (Mg\(^{2+}\), Mn\(^{2+}\), Cd\(^{2+}\), and Zn\(^{2+}\)) resulted in a loss of catalytic activity (Figure 4E) [42]. Since ACP-TX-II requires Ca\(^{2+}\) for activity, chelators such as EDTA inhibited the enzymatic activity (Figure 4F). Histidine residue alkylation also abolished enzymatic activity of ACP-TX-II PLA\(_2\) (Figure 4F), although alkylation does not completely disrupt the 3D structure of PLA\(_2\) enzymes and their capacity to bind phospholipids, but it could modify their ability to interact with specific ligands or proteins [43].

Crotapotin is the acidic moiety of crotoxin that specifically chaperones the basic PLA\(_2\) subunit, inhibiting its catalytic activity [44]. F5 and F6 crotapotins from C. d. collilineatus inhibited the enzymatic activity of ACP-TX-II PLA\(_2\) by approximately 55% (Figure 4F). These results are consistent with PhTX-II PLA\(_2\) from Porthidium hyoprora, which was inhibited ~60% by F2 and F3 crotapotins from C. d. collilineatus [8]. These results suggest that crotapotins may bind to Agkistrodon PLA\(_2\), much like how they bind to crotaline PLA\(_2\)s.

Although snake venom PLA\(_2\)s exhibit diverse pharmacological activities [2], myotoxicity is one of the most common effects [45]. Necrosis induced in vivo in skeletal musculature by intramuscular injection or ex vivo by incubation with differentiated skeletal muscle cells [46] is evidenced by increased plasma CK levels. ACP-TX-II PLA\(_2\) increased serum CK levels when injected intramuscularly (Figure 5B). Local myotoxicity is a characteristic of viperid envenomations, in which PLA\(_2\)s affect predominantly muscles near the injection site [47]. This is consistent with clinical examinations of Agkistrodon bites in the United States, where the main clinical manifestations are local effects that are in some cases associated with permanent dysfunction, while systemic effects are generally absent [19].

ACP-TX-II PLA\(_2\) showed no systemic myotoxicity when injected intravenously. CK levels were similar to those of controls (Figure S2). Perhaps it lacks specificity and attaches to tissues at the site of injection, consistent with the hypothesis that myotoxins may act locally or systemically, proposed by Gutierrez and Ownby [46] in order to explain the pharmacological specificity of venom PLA\(_2\)s. Myotoxic PLA\(_2\)b bind predominantly to different cell types, as well as to muscle fibers, and are quickly sequestered after injection. On the other hand, systemic myotoxic PLA\(_2\)s, such as PLA\(_2\)s F6 and F6a from Crotalus durissus collilineatus [47], have high selectivity for skeletal muscle fibers and do not attach to other cells. This specificity allows systemic myotoxins to spread beyond the injection site, reaching the bloodstream and distant muscle cells and causing rhabdomyolysis.

Homologous K49 PLA\(_2\)s, despite lacking catalytic activity, also cause myonecrosis when injected intramuscularly in mice [48]. Basic/hydrophobic amino acid residues located in the C-terminal region are considered one of the structural determinants for K49 myotoxicity [49,50]. To exert myotoxicity, these toxins function as obligate dimers [11]. According to this model, catalytically inactive K49 dimers undergo structural rearrangement when a membrane fatty acid enters the hydrophobic channel of one of the monomers. This reorientation aligns the C-terminal regions of the two monomers in the same plane and facilitates membrane destabilization when specific hydrophobic amino acids (Leu121, Phe125) penetrate the membrane. Once the membrane becomes disorganized, cells lose ionic control, resulting in cell death [11].

Unlike most Bothrops K49 PLA\(_2\)s, ACP-TX-I does not display detectable local myotoxicity in mice (Figure 5A). Its lack of myotoxicity may reflect its monomeric structure (Figure 1 insert), preventing it from adopting different oligomeric configurations, appropriate to the physicochemical environment. These data corroborate findings with MjTX-I, which is a monomeric K49 PLA\(_2\) in solution isolated from Bothrops moojeni venom, and which shows markedly decreased myotoxic activity [51].

ACP-TX-I and ACP-TX-II induce edema, but even at high concentrations, ACP-TX-I does not reach 30% edema (Figure 6A). The marked edema induced by ACP-TX-II is likely due to phospholipid hydrolysis. This possibility is supported by studies showing that chemical modification of D49 PLA\(_2\)s
with p-BPB, which nullifies catalytic activity, dramatically reduces edematogenic activity of MT-III PLA2 from B. asper as well as other D49 PLA2s [52,53].

In addition to in vivo myotoxic and edematogenic activities, ACP-TX-I and ACP-TX-II were assayed for cytotoxicity in vitro on NIH/3T3 human fibroblasts and A549 lung cancer cells. ACP-TX-I showed high cytotoxicity in both cell lines (Figure 7A). On the other hand, even at concentrations as high as 500 µg/mL, ACP-TX-II had no inhibitory effect on NIH/3T3 fibroblasts. However, the A549 lung cancer cell line was sensitive to this myotoxin, losing ~30% viability after 24 h of incubation (Figure 7B).

The exact molecular mechanism by which snake venom PLA2 decrease cell viability is unknown. Some authors have proposed that cytotoxic activity on tumor cells is associated with apoptosis induction [54], and propose that PLA2 activity accelerates the rate of phospholipid renewal, which induces membrane changes that occur during apoptosis [55]. However, other mechanisms have been proposed: CC-PLA2-1 and CC-PLA2-2 from Cerastes cerastes inhibit cancerous cell adhesion and migration, as well as angiogenesis [56]. Crotoxin B interferes with signaling at epidermal growth factor receptors [57]. Bothrops myotoxins promote fatty acid-dependent lysis by interacting with a receptor able to activate intracellular lipase [58], etc. A K49 PLA2 from Protobothrops flavoviridis induces cell death by caspase-independent apoptosis, accompanied by rapid plasma membrane disruption in human leukemia cells. Some homologous PLA2s, such as MTX-II, exert cytotoxicity regardless of catalytic activity [59]. It is noteworthy that the cytotoxic mechanism depends on activities exerted by molecular regions other than the catalytic site.

4. Materials and Methods

4.1. Venom and Reagents

Venom was obtained from the National Natural Toxins Research Center (Texas A&M University-Kingsville). Solvents and reagents used were HPLC grade, sequence grade, or high purity, obtained from Sigma, Aldrich Chemicals, Merck and BioRad.

4.2. Purification of ACP-TX-I and ACP-TX-II

PLA2 enzymes, ACP-TX-I and ACP-TX-II, from A. c. pictigaster venom were isolated by RP-HPLC, following Huancahuire-Vega et al. [7]. A measure of 20 mg of whole venom was dissolved in 250 µL of 0.1% TFA (buffer A) and centrifuged at 4500 g. Supernatant was then applied to an analytical RP-HPLC µ-Bondapak C18 column (0.78 × 30 cm; Waters 991-PDA system Milford, MA., USA), and equilibrated with buffer A for 15 min. Protein elution employed a linear gradient (0–100%, v/v) of 66.5% acetonitrile in 0.1% TFA (buffer B) at a flow rate of 1.0 mL/min. Elution was monitored at 280 nm and PLA2 activity was assayed in each fraction. Active PLA2 fractions (ACP-TX-I and II) were collected, lyophilized, and used for subsequent biochemicalfunctional characterization.

4.3. Electrophoresis

Molecular masses of ACP-TX-I and II were determined under reducing and non-reducing conditions using Tricine SDS-PAGE in a discontinuous gel and buffer system [60]. Lysozyme, soybean trypsin inhibitor, carbonic anhydrase, ovalbumin, albumin, and phosphorylase B were used as molecular mass markers.

4.4. Determination of Molecular Masses of the Purified Proteins by Mass Spectrometry

A measure of 4.5 µL aliquots of ACP-TX-I and ACP-TX-II PLA2 was injected using a C18 (100 µm × 100 mm) ultra-high-performance reversed phase column (nanoAcquity UPLC, Waters) coupled with nanoelectrospray tandem mass spectrometry on a Quadrupole Time-of-flight (Q-Tof) Ultima API mass spectrometer (MicroMass/Waters Milford, MA., USA) at a flow rate of 600 nL/min. The spectrometer was operated in MS continuum mode, and data acquisition was from m/z 100–3000 at a scan rate of 1 s and an interscan delay of 0.1 s. The gradient used was 0–50% acetonitrile in 0.1%
formic acid over 45 min. A number of m/z mass spectra were accumulated over about 300 scans, and data were converted to molecular masses using maximum-entropy-based software from Masslynx 4.1 (Waters, Milford, MA., USA, 2005). Output masses ranged from 6000–20,000 Da at a 0.1 Da/channel “resolution.” The simulated isotope pattern model was used with the spectrum blur width parameter set to 0.2 Da, and minimum intensity ratios between successive peaks were 20% (left and right). After the smoothing of deconvoluted spectra, mass centroid values were obtained using 80% of the peak top and a minimum peak width at half the height of 4 channels [8].

4.5. Analysis of Tryptic Digests

Prior to trypsin addition (Promega sequencing grade modified), ACP-TX-I and II PLA$_2$ were reduced (DTT 5 mM for 25 min to 56 °C) and alkylated (iodoacetamide, 14 mM for 30 min). After trypsin addition (20 ng/µL in Ambic 0.05 M), samples were incubated 16 h at 37 °C. The reaction was stopped with 0.4% formic acid and samples were centrifuged at 2500 rpm for 10 min. Pellets were discarded and supernatants were dried in a Speed Vac. Peptides were separated by C$_{18}$ reverse phase chromatography (100 µm × 100 mm) (nanoAcquity UPLC, Waters, Milford, MA., USA) coupled with nanoelectrospray tandem mass spectrometry on a Q-Tof Ultima API mass spectrometer (MicroMass/Waters) at a flow rate of 600 nL/min. In order to select ions of interest, an ESI/MS mass spectrum (TOF MS mode) was acquired for each HPLC fraction before performing tandem mass spectrometry, over the mass range of 100–2000 m/z. Then, these ions were fragmented in the collision cell (TOF MS/MS mode). Raw data files from LC-MS/MS runs were processed using Masslynx 4.1 (Waters) and analyzed using the MASCOT search engine, version 2.3 (Matrix Science Ltd. London, UK) against the snake database, using the following parameters: trypsin as the enzyme, fragment mass tolerance of ±0.1 Da, peptide mass tolerance of ±0.1 Da, and oxidation as a variable modification for methionine. Sequence alignments of ACP-TX-I and ACP-TX-II with K49 and D49 PLA$_2$s, respectively, were made using ClustalW in Edit Seq 5.01 © DNASTAR. (Madison, WL, USA, 2001).

4.6. PLA$_2$ Activity

PLA$_2$ activity was assayed as described by Holzer and Mackessy [61], and adapted for 96-well plates. The final volume of the standard assay mixture was 260 µL and contained 20 µL of substrate 4-nitro-3-(octanoyloxy) benzoic acid (3 mM), 200 µL of buffer (10 mM Tris–HCl, 10 mM CaCl$_2$, and 100 mM NaCl, pH 8.0), 20 µL of water, and 20 µL of ACP-TX-I or ACP-TX-II (1 mg/mL). The mixture was incubated at 37 °C for 40 min, measuring absorbances at intervals of 10 min. The initial velocity (Vo) was calculated based on the value of absorbance after 20 min of reaction. ACP-TX-II was chosen by studying kinetic parameters. Different substrate concentrations (40, 20, 10, 5, 2.5, 1.0, 0.5, 0.3, 0.2, and 0.1 mM) incubated in in Tris–HCl buffer, pH 8.0 at 37 °C, were used. The optimal temperature was determined by incubating the enzyme at different temperatures. Similarly, buffers of different pHs (4–10) were used in order to determine the optimal pH. All assays were done in triplicate and absorbances at 425 nm were measured with a VersaMax 190 multiwell plate reader (Molecular Devices, Sunnyvale, CA, USA).

4.7. Inhibition and Chemical Modifications

The effects of EDTA and low molecular weight heparin (Mr 6.000 Da) on enzymatic activity of ACP-TX-II PLA$_2$ were performed by incubating the enzyme with a 1 mM solution of EDTA or a molar ratio of 2:1 (heparin:toxin) at 37 °C for 30 min. Similarly, the effect of crotapotins F5 and F6 (1 mg/mL) from Crotalus durissus collilineatus upon enzymatic activity of ACP-TX-II was evaluated under the same conditions. Additionally, modification of His residues of ACP-TX-II with p-bromophenacyl bromide (BPB) (1.5 mg/mL in ethanol) was performed [43].
4.8. Myotoxic Activity

Different amounts (20 and 50 µg) of ACP-TX-I and II PLA\textsubscript{2} dissolved in 100 µL of PBS were injected IM or IV, in groups of four Swiss mice (18–20 g). The control group received 100 µL of PBS. Blood was collected from the tail into the heparinized capillary tubes at different intervals (1, 3, 6, 9, and 24 h), and 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, where one unit is defined as the phosphorylation of 1 mmol of creatine/min at 25 °C.

4.9. Edema-Forming Activity

Fifty µL of phosphate-buffered saline (PBS; 0.12 M NaCl, 0.04 M sodium phosphate, pH 7.2) with ACP-TX-I and II PLA\textsubscript{2} (10, 20, and 50 µg/paw) were injected into the subplantar region of the right footpad of five Swiss mice (18–20 g). The left footpad received 50 µL of PBS, as a control. Immediately before inoculation (basal) and at different time intervals (1, 3, 6, 9, and 24 h) paw volume was evaluated by plethysmography (Model 7140 Pletysmometer Ugo Basile, VA, Italy). Edema-forming activity was expressed as the percentage increase in volume of the right foot pad in comparison to the left foot pad (control). The percentage of edema in toxin-inoculated paws was calculated with the equation: % edema = [(T_x \times 100)/T_0] – 100. T_0 is the paw volume before toxin injection. T_x is the edema (volume) measured at each time interval. The percentage of edema calculated was subtracted from the matched values at each time point in the saline injected hind paw (control) [62].

4.10. Cytotoxic Activity

Cytotoxic activity was assayed on NIH 3T3 fibroblasts (ATCC®CCL-1658™) and A549 lung cancer (ATCC®CCL-185™) cells, grown in plastic flasks (25 cm\textsuperscript{2}) with RPMI 1640 medium (Cultilab, Campinas, SP, Brazil), was supplemented with 2% L-glutamine, 120 µg/mL garamycin, and 13% inactivated fetal bovine serum (complete medium). Cultures were incubated at 37 °C in an atmosphere containing 5% CO\textsubscript{2}. The medium was changed every 48 h, and when the culture reached confluence, subculturing was performed by treatment with trypsin and versene (Adolfo Lutz, São Paulo, SP, Brazil). Variable amounts of both ACP-TX-I and II were diluted in the assay medium and added to cells in 96-well plates. Experiments were carried out in triplicate. Cellular viability was assayed by the neutral red uptake assay of Ates et al. [63]. After treatment with toxins, the medium was removed and the culture was washed with PBS. For each well, 0.2 mL RPMI medium containing 50 µg/mL Neutral Red dye was added. The plate was incubated for 3 h at 37 °C to capture dye by viable cell lysosomes. After incubation, the medium containing the dye was removed and the wells rapidly washed with formalin-calcium to remove unincorporated dye from the cells. 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 min on a microtitre plate shaker, absorbance was read at 540 nm. Cell viability was expressed as percentages compared to control and untreated cells.

4.11. Statistical Analyses

Results were reported as mean ± SEM. Dunnett’s test was used to determine the significance of differences among means by analysis of variance when several experimental groups were compared with the control group. Differences were considered statistically significant if p < 0.05.

Supplementary Materials: The following are available online at http://www.mdpi.com/2072-6651/11/11/661/s1, Figure S1: Re-chromatography on an analytical RP-HPLC C18 analytical column of ACP-TX-I and ACP-TX-II. Figure S2: ACP-TX-II produces local myotoxicity when injected intramuscularly, but little systemic myotoxicity when injected intravenously, whereas ACP-TX-I and crude venom injected intravenously produce no systemic myotoxicity.

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Ethical statement: 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 and international law and policies.

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