Exploring the biochemical, haemostatic and toxinological aspects of mapanare dry-tail (*Porthidium lansbergii rozei*) snake’s venom causing human accidents in Eastern Venezuela

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Abstract. *Porthidium lansbergii rozei* (*P.l.rozei*) is a snake species belonging to the Venezuelan ophidian-fauna causing relatively frequent human accidents. This study has been developed to enrich the medical information about this snake’s accident, which is often handled with difficulties due to the ignorance about its toxic activities. This work has studied the biochemical and toxinological characteristics of the *P.l.rozei* venom. The objective was to determine its biochemical characteristics, including protein content determination, electrophoretic and *P.l.rozei* chromatographic profile. The toxinological and enzymatic functions, comprised lethality (LD₅₀), haemorrhagic (MHD), myotoxic, coagulant, oedematising, proteolytic, fibrinogenolytic, fibrinolytic, anticoagulant, and phospholipase activities and crude venom recognition by the Venezuelan polyvalent antivenom (PAOS). The venom induced haemorrhagic, myotoxic, anticoagulant, fibrinolytic, fibrinogenolytic, and phospholipase A₂ activities in mice; the LD₅₀ was not established even though the doses tested expand the different lethal doses reported in previous studies. The venom induced a dose-dependent oedema up to the dose of 4 μg, corresponding to 27.91%. In conclusion, the studied venom presented a haemostatic, toxinological, and enzymatic biochemical profile similar to the one described for specimens of the *Porthidium* genus. However, there were evident quantitative variations in their different activities, offering new evidence of the intra-species variations that exists in the different snake families. These variations must be accounted for appropriate clinical management.
INTRODUCTION

The accidents caused by venomous animals of several zoological groups (snakes, scorpions, and spiders), represent a global collective health problem especially associated to the tropical and subtropical zones (1). Venomous ophidic accident, however, is a neglected disease, despite its high incidence and severity, causing permanent functional sequelae that can progress in affected individuals and their death (2).

Within the most important American snake families, there have been described nine species of the Porthidium genus, which inhabit dry or wooded areas from southern Mexico to north-western South America including Venezuela, Ecuador and Colombia (3). In Venezuela, the Porthidium lansbergii species includes three subspecies: Porthidium lansbergii hutmanni, Porthidium lansbergii lansbergii and Porthidium lansbergii rozei (4, 5). The latter are distributed in the dry and wooded areas of Anzoátegui, Sucre, Bolívar,
Trujillo, Lara, Falcón, Aragua, Carabobo and Delta Amacuro states (Venezuela) (6).

A number of reports in the literature have documented different levels of variation in the composition of Porthidium venom (7, 8). These differentiations may be related to phylogenetic or taxonomic discrepancies, age, size, geographic location, diet, seasonal variations, variations associated with sex, among others, all these lead to divergences in the quantity and quality of the venoms. The venoms can differ, both at the macro and micro-variable levels, but also can share many components through wide taxonomic levels (9).

Regions with an abundant venomous herpetofauna, as Venezuela, must make efforts to identify snakes causing accidents in cattle and rural human populations. Their venoms must be extensively characterised, identifying the factors involved in the morbidity and mortality of envenomation, with the aim of create guidelines to advance the victim treatments and reinforce the phases of design, production and control of antivenoms, grounded on the accuracy of ophidism in the regions involved (10).

In Venezuela, information related to the Porthidium lansbergii subspecies -widely distributed in the national territory- is scarce. Although it is frequently known by rural doctors - not being a rare envenoming- its handling presents difficulties due to the ignorance of its toxic activities. This work has been developed to enrich the medical information of the type of accident. For this reason, in order to expand this knowledge, some biochemical characteristics and biological activities, as well as the recognition of the P. lansbergii rozei (P.l.rozei) venom by Venezuelan polyvalent (PAOS) antivenom serum were studied.

**MATERIALS AND METHODS**

**Reagents**

Bovine fibrinogen, thrombin and casein, agarose and creatine kinase (CK) determination Kit 520-C (Sigma-Aldrich Chemical Co., MO, USA), molecular weight markers for electrophoresis (SDS-PAGE) (Biorad, Philadelphia, USA), plastic Petri dishes (Falcon, USA), polyvalent anti-ophidic serum (PAOS) (Biotecfar S.A, Faculty of Pharmacy of the Central University of Venezuela).

**Experimental animals**

White male mice NIH strain, with weights pre-established for each test, purchased from the National Institute of Hygiene “Rafael Rangel” (Caracas, Venezuela) animal facility were used. They were kept in stainless steel cages with hydration and food supply *ad libitum*.

**Ethical statement**

All the experimental procedures related to the use of live animals were carried out by expert personnel, under the pertinent Venezuelan regulations, according to protocols of action approved by Ethical Committee of the Anatomical Institute of the Universidad Central de Venezuela, and the norms for the care and use of animal’s laboratory, published by the National Institute of Health (USA) (11).

**Snakes and venom**

The P.l.rozei specimens were captured in the surroundings of Cumaná city (Sucre state, Venezuela) (Fig. 1) in “Cerros de Caiguire” which are a physiographic unit located at the North of the city and the East of Manzanares River, from 5 to 160 m elevation, bordered by a coastal plain that separates the highlands to the south and east of the mentioned hills (12). The ecological area of the snake is located in a bio-region of xerophytic climate, with an average annual temperature of 28°C, amplitude between the maximum daytime temperature and the minimum nocturnal temperature of 10 to 12°C (amplitude of megathermic type), rainfall near 500 mm annual, with an average relative humidity of 50% or less. The flora corresponds to typical dry areas of plant formation, such as *Opuntia*, cactuses, spiny herbs, and trophophilous forests (13).
Venoms pools from three adult males of mapanare dry-tail (*Porthidium lansbergii rozei*) (*P.l.rozei*) were used. The snakes were kept in captivity in the Serpentarium of the Tropical Medicine Institute of the Universidad Central de Venezuela (Caracas, Venezuela). After extraction, the venom was centrifuged, lyophilized and stored at -30°C.

*Porthidium lansbergii hutmanni* pool venom used for chromatography was purchased from Bioreptilia C.A (Caracas, Venezuela). This venom was extracted from specimens captured in Margarita Island (Venezuela).

**Plasma**

Fresh human plasma was obtained from blood samples of researchers in this work, without alterations in haemostasis or taking anticoagulants. The sample was mixed with 3.8% sodium citrate in a 9:1 ratio. Plasma was obtained by centrifugation at 2000 rpm for 15 minutes.

**Electrophoresis in polyacrylamide gels (SDS-PAGE) of *P.l.rozei* crude venom**

To establish the electrophoretic profile with 25 μg of *Pl.rozei* venom, 12% SDS-PAGE was performed using Tris/glycine gels, under reducing and native conditions, following the Laemmli (1970) method (14). Additionally, 10% Tris/Tricine gels (15) were used to visualise the fibrinogenolytic venom activity. The gels were fixed and stained with silver stain and/or Coomassie blue.

**Fractionation of proteins by reverse phase high resolution liquid chromatography**

*Porthidium l. rozei* and *Pl. hutmanni* venoms were run in a liquid phase chromatography (RP-HPLC). Briefly, in 200 μL [0.1% trichloroacetic acid (TCA)] solution, 0.34 mg of venom was dissolved and chromatographed on a Simmetry® C-18 column (4.6 x 150 mm) using a high resolution chromatography (CLAR) Delta 1525 (Waters, Massachusetts, USA). The elution was carried out at a flow of 1mL/min, applying a gradient of solution B (acetonitrile in 0.1% TCA): 0% B for 5 min, 0 to 5% B for 10 min, 15 to 45% B for 60 min, 45 to 70% B for 10 min and 70% B for 9 min.

**Lethal Activity (LD50)**

The lethality of *P.l.rozei* venom was determined by the Spearman-Kärber method (16). Five mice per dose weighing 18–22 g were intraperitoneally injected with 200 μL of the serially diluted venom ranging (38.46 and 241.3 μg). Deaths were recorded during a 48 h period. Mice were observed up to 48 h after injection.
Coagulant activity

To determine the *P. l. rozei* venom coagulant activity, the method of Austen and Rhymes (17) was used. Bovine fibrinogen was handled as substrates to evaluate the thrombin-like activity, and fresh citrated plasma, to determine the procoagulant activity. Concisely, 0.3% fibrinogen solution (100 μL) diluted in 0.05 M Tris-HCl buffer, pH 7.4 (coagulation buffer) or citrated plasma was prepared and then incubated for 3 min at 37°C. At that point 100 μL of coagulation buffer, and 100 μL of thrombin solution (2.5 UI / mL) (positive control), or 100 μL of crude venom (1mg/mL) were added and mixed. Then, the coagulation time at 37°C was recorded. The results were expressed in seconds.

Anticoagulant activity

Aliquots of 0.2 mL of citrated human plasma (3.8% v/v) were distributed in glass tubes and incubated in a water bath for 5 min at 37°C. Then, 50 μL (40 μg) of *P. l. rozei* venom was added and further incubated for 10 min at 37°C. Finally, 50 μL of 0.25 M CaCl₂ was added and the coagulation time was determined. The negative control tubes contained plasma incubated only with PBS. The tests were carried out in duplicate and the results were expressed in seconds.

Fibrinogenolytic activity

The effect of the venom, on the fibrinogen chains using a solution of human fibrinogen (5 mg/mL) in 0.05M Tris-HCl buffer, pH 7.5 was evaluated by fibrinogen (Fg): *P. l. rozei* ratio of 100:1 (100μg Fg/1μg venom). A lane control containing only fibrinogen was used for the visualization of the band pattern corresponding to the α, β and γ chains of fibrinogen. The fibrinogen was incubated for 4 hours, at 37°C with the venom, initially in different substrate enzyme ratios, in order to select the appropriate proportion, later evaluating different incubation times, to determine the kinetics chains degradation by means of electrophoresis. Tris-Tricine system (15) 10% gels electrophoresis under reduced conditions, visualizing the protein bands with 0.5% Coomassie blue was performed.

Fibrinolytic activity

Fibrinolytic activity of *P. l. rozei* venom was calculated by the fibrin plate method as described by Marsh and Arocha-Piñango (18). Briefly, fibrin plates were prepared using 3-em diameter Petri dishes: 1.5mL of a 0.1% plasminogen-rich fibrinogen in imidazol saline buffer, pH 7.4 was coagulated by adding 75 mL bovine thrombin (10 U/mL, in 0.025M CaCl₂). The mixture was incubated at room temperature for 30 min. Then, 10 μL sample (1 mg/mL) was applied over the fibrin film, and after 24 h incubation at 37°C the diameter of the lysed areas was measured. Fibrinolytic activity was expressed as the lysed area per microgram of protein (mm²/μg). As positive control, 10 μL (1 mg/mL) of *Bothrops colombiensis* crude venom and 12 μL of saline solution as negative control were used.

Oedematising activity

For the activity of oedema formation, the method of Yamakawa et al. (19) was used. A group of five mice, weighing 18-20 gr, were injected into the footpad of the mice right lower limb with venom serial dilutions (2 to 32μg / 50μL diluted in saline solution). The plantar pad of the left lower limb was injected with 50 μL of saline solution; one hour after the injection, the animals were sacrificed by ether inhalation and then both limbs were cut with a scalpel at the tarsal joint level. Both extremities were weighed in the analytical balance, and the oedema was determined, as the percentage of weight increase, in the right extremity with respect to the left. The minimum oedematising dose (MOD) was defined as the least amount of venom producing an oedema higher than 30% after one hour of exposure. The percentages (%) of oedema were plotted against the *P. l. rozei* doses.
Determination of minimal haemorrhagic dose (MHD)

To determine the MHD for the *P.l.rozei* venom, Kondo et al. (20) method was used. Serial venom doses (2.5 to 40 μg) were dissolved in 100 μL of isotonic saline; 0.1mL of each dilution was intradermal injected into the depilated abdominal skin of mice. The mouse was sacrificed and the skin removed after 2 h. The haemorrhagic diameter on the skin was measured and the MHD determined. The MHD is defined as the amount of venom protein that causes a 10 mm haemorrhagic spot, determining the diameter of the developed lesion using the equation:

\[ D = 2\sqrt{A/\pi} \]

Caseinolytic activity

To determine the caseinolytic activity of the *P.l.rozei* venom, a casein zymography was carried out according to the method of Da Silveira et al. (21). Concisely, 40 μg *P.l.rozei* crude venom was diluted in sample buffer (4X) and run on 12% SDS-polyacrylamide gels, copolymerized with 2 mg/mL casein, under non-reducing conditions, at 4°C and 20 mA. After electrophoresis, the gels were washed twice for 30 min with 2.5% Triton X-100 to remove the SDS and then incubated in the zymography incubation buffer (0.05 M Tris-HCl, pH 7.3, 0.2 M NaCl, 0.001 M CaCl₂, 0.001 M MgCl₂) at 37°C for 18 h. The gels were stained with 0.25% Coomassie brilliant blue R-250 in acetic acid: ethanol: water solution (7:30:63 v/v) and then kept in the same solution. The presence of caseinolytic activity was defined as light bands on a dark background.

Myotoxic activity

The ability of the *P.l.rozei* venom inducing skeletal muscle necrosis was evaluated in mice (18-20 g). A single dose of 40 μg/50 μL of venom was injected into the gastrocnemius muscle. The dose was selected on the basis of previous studies on Viperidae snake venoms (22). As a negative control, the mice received injections of 50 μL of saline solution. After 3 hours of injection, blood was collected by cardiac puncture, for enzymatic creatine kinase (CK) serum determination using a commercial kit assay (Sigma Kit 520-C, Sigma Chemical Co.) The CK activity was expressed in international units per litre (IU/L). Once the blood sample was taken, the mice were sacrificed by cervical dislocation, and the extremity inoculated with the studied venom was macroscopically examined, comparing it with the extremity of the negative control mouse.

Zymography of phospholipase A2 (PLA₂)

In order to identify and estimate the PLA₂s molecular weights, a *P.l.rozei* venom sample using a zymography was analysed. Briefly, the venom sample (10 μg) was subjected to a 12% SDS-PAGE on an under non-reducing conditions, at 60 V and 4°C. The gel was washed for 1 hour in 500 mM Tris-HCl, pH 7.4, containing 2.0% (v/v) Triton X-100 and incubated for another hour, in the same solution, but with 1.0% triton X-100 (v/v). After removing the SDS residues, the gel was washed a third time for 30 minutes in 50 mM Tris-HCl, pH 7.4, containing 140 mM NaCl and 2.5 mM CaCl₂. It was then incubated for 14 hours at room temperature on a 1.0% (w/v) agarose gel, prepared in 50 mM Tris-HCl, pH 7.4, containing 140 mM NaCl and 2.5 mM CaCl₂ and 2.0 % of egg yolk. The light zones indicated the PLA₂ presence.

Recognition of the *P.l.rozei* venom by polyvalent antivenom serum (PAOS)

The venom toxins identification was carried out by the technique of double immunodiffusion in agarose gels (23) for the detection of anti-*P.l.rozei* antibodies in the polyvalent antivenom serum (PAOS). This antivenom neutralizes at least 2 mg of bothropic venom by 1 mL of serum. Briefly, slides were pre-treated with 1% agarose and then covered with 3.5 mL of 2% agarose. Once the gel was solidified, the wells were drilled equidistant and distributed, one central and 6 peripherals. In the central and
peripheral wells there were placed 10 μL of PAOS and 10 μL of 2.5, 10 to 40 μg dilutions of Pl.rozei venom, respectively. It was incubated at 37°C for 24 hours. Once the precipitation bands were detected, the gel was washed with 0.9% saline and stained with Coomassie Blue.

RESULTS

Gel electrophoresis (SDS-PAGE) of Pl.rozei venom

The 12.0% SDS-PAGE of Pl.rozei venom was run under reducing and non-reducing conditions. The gel under non-reducing conditions revealed in the first group two (2) bands located between ~66.2 to 45.0 kDa molecular weights (MW). The second group was between ~45.0 to 31.0 kDa MW and had three (3) bands. Under reducing conditions the first group had three (3) bands between ~45.0 to 31.0 kDa MW, and two bands with lower MW: one (1) band found between 31.0 to 21.5 kDa; while the other bands were between MW of 21.5 to 14.4 kDa (Fig. 2).

Venom fractionation by reverse phase (HPLC)

Fig. 3 shows the comparative chromatograms corresponding to Pl.rozei and Pl.hutmanni venoms run in a Simmetry® C-18 column. There were 28 signals, two of them were very intense and corresponded to 21 and 82 min in the venom of Pl.rozei. The chromatograms differed mainly in the second intense signal (close to 82 min) of P. I. hutmannii, and the acetonitrile signal of Pl.rozei at 87 min.

Coagulant activity

The evaluated bovine fibrinogen and fresh citrated plasma were uncoagulable using 100 μg/venom dose; the positive plasma controls gave an average of 19.85 ± 1.2 s, and the positive control for fibrinogen gave a mean of 18.75 ± 0.9 s. The results are shown including the mean ± standard deviation with n = 5 (Table I).

Anticoagulant activity

The Pl.rozei venom in vitro presented anticoagulant activity, showing the fresh citrated plasma uncoagulable at 300 s with a dose of 40 μg of venom. On the other hand, the negative plasma control gave an average of 25.0 ± 1 s. The results are shown including the mean ± standard deviation with n = 5 (Table I).

Fibrinogenolytic activity

Fig. 4 shows the Pl.rozei venom fibrinogenolytic activity, at a fibrinogen (Fg): Pl.rozei ratio of 100:1 (100μg Fg/1μg venom); whole degradation of the Aα chains as well as soft degradation of the Bβ chains was observed. No apparent alteration to the γ chains, even after 24 h of incubation, was noticed. The degradation of the Aα chains occurred between 1 to 30 min. In contrast, the degradation of the Bβ chains was between 5 min to 2 h. This fibrinogenolytic activity was completely inhibited by metalloprotease inhibitor (EDTA), while the serine proteinase inhibitor (trypsin or trypsin-like...
Biochemical and toxinological characteristics of *P. l. rozei* venom

**Fig. 3.** Chromatograms of *Porthidium l. rozei* (A) and *P. lansbergii hutmanni* (B) venoms (0.34 mg) comparison by Simmetry® C-18 column by reverse phase high resolution liquid chromatography. Elution: Buffer B: 0-70% in 94 min. Existence of several forms and quantities of proteins in both subspecies has stimulated considerations as possible functional venom differences among them.

**TABLE I**

COMPARISON OF THE *P. lansbergii rozei* VENOM (CUMANÁ, SUCRE STATE, VENEZUELA) ENZYMATIC ACTIVITIES WITH DIFFERENT *Porthidium* VENOM SPECIES PREVIOUS REPORTS

<table>
<thead>
<tr>
<th>Species</th>
<th>MOD (μg/mouse)</th>
<th>Coagulant</th>
<th>Anticoagulant</th>
<th>Fibrinolytic (mm&lt;sup&gt;2&lt;/sup&gt;/μg)</th>
<th>PLA&lt;sub&gt;2&lt;/sub&gt;</th>
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<tr>
<td><em>Porthidium lansbergii hutmanni</em></td>
<td>1.03</td>
<td>AA</td>
<td>NE</td>
<td>6.8</td>
<td>NE</td>
<td>Girón et al (2011)&lt;sup&gt;10&lt;/sup&gt;</td>
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<td>(Margarita Island, Venezuela)</td>
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<tr>
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<td>NE</td>
<td>NE</td>
<td>NE</td>
<td>AM PA</td>
<td>Ramos et al (2010)&lt;sup&gt;16&lt;/sup&gt;</td>
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<tr>
<td><em>Porthidium lansbergii lansbergii</em></td>
<td>AM</td>
<td>AA</td>
<td>PA</td>
<td>NE</td>
<td>AM PA</td>
<td>Jiménez-Charris et al (2015)&lt;sup&gt;27&lt;/sup&gt;</td>
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<td>(Atlantic, Colombia)</td>
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<td><em>Porthidium Nasutum</em></td>
<td>1.4 ± 0.1</td>
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<td>Otero et al (2002)&lt;sup&gt;16&lt;/sup&gt;</td>
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<td>NE</td>
<td>AA</td>
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<td>NE</td>
<td>NE</td>
<td>Bogarin et al (2000)&lt;sup&gt;15&lt;/sup&gt;</td>
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<tr>
<td><em>Porthidium Nasutum</em></td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
<td>AM PA</td>
<td>Gutiérrez et al (2013)&lt;sup&gt;47&lt;/sup&gt;</td>
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<td>NE</td>
<td>AM PA</td>
<td>Gutiérrez et al (2013)&lt;sup&gt;47&lt;/sup&gt;</td>
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<td><em>Porthidium lansbergii hutmanni</em></td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
<td>NE</td>
<td>Pineda et al (2008)&lt;sup&gt;26&lt;/sup&gt;</td>
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<tr>
<td><em>Porthidium lansbergii rozei.</em></td>
<td>AA</td>
<td>AA</td>
<td>PA</td>
<td>16.9 ± 2.8</td>
<td>PA</td>
<td>This study</td>
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<td>(Sucre state, Venezuela)</td>
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</table>

MOD: Minimum oedematising dose; PLA<sub>2</sub>: Phospholipase A2; NE: Not evaluated; AM: Another methodology; PA: Present activity; AA: Absent activity.
inhibitor benzamidine) had no effect (data not shown). Coagulant activity of fibrinogen (pre-treated with *Pl. rozei* venom [100:1]) by thrombin resulted in the formation of fine fibrin threads after 280 s. Additionally, fibronectin degradation of fibronectin (a molecule present in the fibrinogen used) can be observed after 5 minutes of incubation. The degradation products are in the molecular mass range of 97.4 and 66.2 KDa.

**Fibrinolytic activity**

The fibrinolytic activity, determined on fibrin plates and expressed as specific activity (mm²/μg), is shown in Fig. 5. It was evident that there was no lysis area in the negative control plate, whereas in the positive control plate the lysis area occupied the entire plate. In the plate with *Pl. rozei* crude venom, a delimited area of lysis was observed. The fibrinolytic activity of the venom was 16.9 ± 2.8 mm²/μg. The results are shown including the mean ± standard deviation with n = 3 (Table I).

![Fig. 4. Fibrinogenolytic 10% (Tris/Tricine) SDS-PAGE.](image-url)

Fig. 4. Fibrinogenolytic 10% (Tris/Tricine) SDS-PAGE. The fibrinogen was incubated with *Pl. rozei*, to 100:1 ratio. Lanes 1) molecular weight marker; 2-5) incubation of 1, 5, 15 and 30 min; 6-9): 1, 2, 4 and 24 h; 10) control of fibrinogen. The arrow shows products of fibronectin degradation.

![Fig. 5. Fibrinolytic activity of the *Pl. rozei* venom.](image-url)

Fig. 5. Fibrinolytic activity of the *Pl. rozei* venom. Fibrin plates with: A) negative control (0.85% NaCl solution); B) 10 μg of *Pl. rozei* venom; C) positive control (10 μg of *Bothrops colombiensis* venom).
Caseinolytic activity

The zymogram corresponding to the *P. l. rozei* venom showed no caseinolytic activity, since no casein digestion bands were observed, in contrast to the *B. colombiensis* venom positive control, in which, at least four clear bands corresponding to the substrate digestion zones could be observed (Fig. 6).

Myotoxic activity

In the evaluation of the myotoxic activity, an increase of the creatine kinase enzymatic activity was observed. In the negative mouse control, it was 166 U/L and in those inoculated with 40 μg of venom, it was 1555 ± 572 U/L (Table II). In addition, macroscopic muscle damage was observed (data not shown).

Zymography of phospholipase A₂ (PLA₂)

In this study, the presence of PLA₂ was analysed by an egg yolk zymogram. The two venoms tested show clear zones in the region near 15 kDa. The same amount of venom (10 μg) was used in both cases. The *B. colombiensis* crude venom represented the assay positive control (Fig. 7).

**Porthidium l. rozei** venom recognition by polyvalent antivenom serum (PAOS)

The polyvalent antivenom serum (PAOS) was able to recognize the *P. l. rozei* venom, which was evidenced as bands of precipitation in the agarose gel (23), denoting the presence of anti-*P. l. rozei* antibodies in the serum (Fig. 8).

**Lethal dose fifty (LD₅₀)**

*Porthidium l. rozei* venom was not lethal to mice at a maximum dose of 241.3 μg/mouse, therefore, it was not possible to determine the lethal dose 50. In this group, an animal showed signs of neurotoxicity manifested as paroxysmal tachycardia, tachypnea, tonic-clonic movements and sphincter relaxation. These manifestations lasted seconds and occurred during the first 15 min post inoculation (Table II).

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![Fig. 6. Caseinolytic activity (casein zymogram). A: molecular weight markers; B) *P. lansbergii rozei* venom; C) *Bothrops colombiensis* venom (positive control).](image)

![Fig. 7. Phospholipases A₂ 12% SDS-PAGE zymogram on 1% agarose gel and 2% egg yolk. A) molecular weight markers; B) 10 μg of *B. colombiensis* venom; C) 10 μg of *Plansbergii rozei* venom.](image)
Minimal Haemorrhagic Dose (MHD)

The MHD for crude venom was 13.87 ± 3.20 μg/mouse. These lesions increased their intensity and diameter in a dose-dependent manner (Table II).

Oedematising activity

An oedema developed in the footpad after a subcutaneous injection of *P.l.rozei* venom, but a minimum oedematogenic dose (30%), with the increased dose until 4 μg was not reached, corresponding to 27.91% increase in the weight of the inoculated limb. Then, the percentage of induced oedema began to decline. With this dose, it was not possible to determine the minimum oedematising dose (MOD), defined as the amount of venom that produces an oedema of 30% after one hour of exposure (Table I). In addition, necrosis was observed (data not reported).

**TABLE II**

COMPARISON OF THE TOXINOLOGICAL *P.l.rozei* (CUMANÁ, SUCRE STATE, VENEZUELA) VENOM ACTIVITIES WITH DIFFERENT *Porthidium* VENOM SPECIES PREVIOUS REPORTS

<table>
<thead>
<tr>
<th>Species</th>
<th>LD$_{50}$ (μg/mouse)</th>
<th>MHD (μg/mouse)</th>
<th>Myotoxic CK (U/L)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Porthidium lansbergii hutmani</em> (Margarita Island, Venezuela)</td>
<td>50 (20-320)</td>
<td>0.98</td>
<td>NE</td>
<td>Girón <em>et al.</em> (2011)</td>
</tr>
<tr>
<td><em>Porthidium lansbergii rozei</em> (Falcón state, Venezuela)</td>
<td>NE</td>
<td>AM</td>
<td>AM</td>
<td>Ramos <em>et al.</em> (2010)</td>
</tr>
<tr>
<td><em>Porthidium Nasutum</em> (Antioquia, Colombia)</td>
<td>109 (82-131)</td>
<td>0.96 ± 0.2</td>
<td>105.000 ± 12.000</td>
<td>Otero <em>et al.</em> (2002)</td>
</tr>
<tr>
<td><em>Porthidium nasutum</em> (Costa Rica)</td>
<td>216</td>
<td>0.5</td>
<td>NE</td>
<td>Bogarin <em>et al.</em> (2000)</td>
</tr>
<tr>
<td><em>Porthidium Nasutum</em> (Costa Rica)</td>
<td>178 (149-290)</td>
<td>1.6 ± 0.3</td>
<td>1389 ± 604</td>
<td>Gutiérrez <em>et al.</em> (2013)</td>
</tr>
<tr>
<td><em>Porthidium Ophryomegas</em> (Pacific and Costa Rica Center)</td>
<td>149 (117-186)</td>
<td>1.1 ± 0.1</td>
<td>2251 ± 1330</td>
<td>Gutiérrez <em>et al.</em> (2013)</td>
</tr>
<tr>
<td><em>Porthidium lansbergii hutmani</em> (Margarita Island, Venezuela)</td>
<td>94.6</td>
<td>25 ± 0.8</td>
<td>NE</td>
<td>Pineda <em>et al.</em> (2008)</td>
</tr>
<tr>
<td><em>Porthidium lansbergii rozei</em> (Sucre state, Venezuela)</td>
<td>Indeterminate in the used range (38.46 to 241.3)</td>
<td>13.87 ± 3.2</td>
<td>1555 ± 572</td>
<td>This study</td>
</tr>
</tbody>
</table>

LD$_{50}$: Lethal dose fifty; MHD: Minimal haemorrhagic dose; NE: Not evaluated; AM: another methodology.

Fig. 8. Recognition of the *P.l.rozei* venom by the polyvalent antivenom serum (PAOS). A) 10 μL of PAOS; B) positive control: *B. colombiensis* (10 μL/40 μg) venom; C) *P.l.rozei* (10 μL/2.5 μg) venom; D) *P.l.rozei* (10 μL/10 μg) venom; E) *P.l.rozei* (10 μL/40 μg) venom F) 10 μL of saline solution; G) empty well.
shown), which increased in intensity directly proportional to the increase in the dose (16 μg) of venom evaluated.

**DISCUSSION**

Snake bites are of a major collective health importance in Venezuela as a cause of morbidity and mortality. *Porthidium* spp., are responsible for several envenomation cases in East and West Venezuela. Antivenom immunotherapy is the only specific treatment available for envenomed patients. In general, more than 4000 annual patients showed signs of envenomation and were treated with antivenom immunotherapy in the last years, most of them without diagnosis of the species involved in the envenomation. Peasants of both sexes were the most frequently affected. The overall lethality was nearly 2%. Lethality and medical hospitalization were significantly high in the differently treated patients (1).

There are not publications of *Pl.rozei* clinical cases of envenomation in the country, for which specific clinical information is lacking. The oral communication of rural physicians, who have attended some cases, refer that patients present the appearance of moderate pain at the site of the bite, accompanied by bleeding through the imprint of the fangs, and presence of slight oedema, erythema and generalised ecchymosis covering at least two segments of the affected limb. The patients have successfully responded to the treatment with the polyvalent anti-ophidic serum used in the country (PAOS, Biotecfar, S.A, Venezuela).

According to the current study, the venom of *Pl.rozei*, a snake geographically located in a neighbourhood of Cumaná, Sucre state, Venezuela (Fig. 1) was able to induce experimental haemorrhagic, myotoxic, anticoagulant, fibrinolytic, fibrinogenolytic, phospholipase A₂ activities, aspects that are in agreement with the great majority of snakes that belong to this genus (24). However, there were evident quantitative variations when comparing the results reported in previous evaluations for the same venom snakes (Tables I and II). In the *Pl.rozei* LD₅₀ venom estimation, limited signs of intoxication in the mice were observed, and none of the different experimental groups died during the time of venom exposure. The doses tested (38.46 to 241.3 μg / mouse) include all the LD₅₀ s reported in previous studies with *Porthidium* genus species from different geographical regions. As reported by Girón et al. (8) the LD₅₀ was 50μg, while Pineda et al. (7) obtained a LD₅₀ of 94.6 μg / mouse for the same species and Jiménez-Charris et al. (25) obtained a LD₅₀ of 164 μg / mouse for *P. langsbergii*. Otero et al. (26) reported a LD₅₀ for *P. nasutum* of 109 μg / mouse (Table II). These differences among different studies may be due to the intraspecies venom variability with snakes of different origin (27). The results obtained in this research suggest that *Pl.rozei* is a less potent venom than those evaluated by other authors.

The comparative chromatograms corresponding to *Pl.rozei* and *Pl.hutmanni* venoms run in a Simmetry® C-18 column were different in the second intense signal (close to 82 min) of *P. l. hutmanni* and in the acetonitrile signal of *Pl.rozei* at 87 min. The intra-signal ratio showed that some peptides were found in a greater proportion in *Pl.hutmanni*, which could result in a greater toxicity of the *Pl.hutmanni* venom, as compared to the *Pl.rozei*, due to the larger number of components, probably toxins in that venom. The objective of this comparison was to show that both venoms, from the same genus, are different in their chromatographic profiles and therefore probably in their toxic action. This interspecies venom variability surely originates different clinical manifestations in the envenomed victims.

The haemorrhagic effect is mainly due to the zinc-dependent metalloproteases actions, which function at microvasculature level, destroying the capillary vessels and causing the erythrocytes and plasma exit into the interstitial space, producing selec-
tive extracellular matrix degradation components, such as different forms of collagen (type IV, type VI, fibrillar, etc.), laminin and fibronectin, these weakens the endothelial basement membrane, followed by its distension and rupture by hemodynamic forces (28, 29).

In the haemorrhagic activity evaluation, which is one of the characteristic effects of *Porthidium* venoms (7, 8, 26, 30, 31), it was observed that the *P.l.rozei* venom exhibits a 13.87 ± 3.2 μg/mouse MHD, which is high when compared with previous reports for this genus (Table II), in which *P.l. hutmanni* (8) reported a 0.98 μg/mouse MHD, *Porthidium nasutum* from Costa Rica (32) obtained a 0.5 μg/mouse MHD, while Gutiérrez et al. (33) reported a 1.6 μg/mouse MHD. In Colombia, the same species (26) presented a 0.96 μg/mouse MHD. All these data suggest that the venom currently studied has a low haemorrhagic activity as well as a low concentration of metalloprotease-like enzymes, in comparison with these venoms.

On the other hand, when comparing the haemorrhage area obtained with 10 μg of the venom under study (68.7 mm²) with *P.l. rozei* (32.5 mm²) evaluated by Ramos et al. (31), it was evident that *P.l.rozei* (Cumaná, Sucre state) had higher haemorrhagic action.

Referring to the current *P.l.rozei*, an increase of CK showed its important myotoxic activity; this result compared with previous *Porthidium* venom studies (Table II) demonstrated that *P.lansbergii* (25), *Porthidium nasutum* and *Porthidium ophryomegas* (33) had the same comportment. Myotoxins and other peptides operating in concert with sodium channels, increasing sodium influx and causing myofibril necrosis are the result of venom toxins (usually basies) action, but sometimes, comprising molecules with acidic phospholipase A₂ (PLA₂) activity, inducing calcium influx, results in permanent cell damage (35).

In the oedematising activity evaluation using 2 to 32μg doses, after one hour of exposure, there was no 30% increase of the right extremity with respect to the left (negative control), which did not allow to calculate the MOD. Nevertheless, an oedematising action was observed. It can be deduced that the *P.l.rozei* venom is not as rich in inflammatory substances, including phospholipases A₂ as other *Porthidium* venoms. The association between oedema and the inflammatory response is obvious and it appears to be species-specific (36). These toxins can induce necrosis, oedema, and inflammation, degrading extracellular matrix components and weakening the regeneration of affected skeletal muscle (37, 38).

Additionally, the *P.l.rozei* venom showed no coagulant activity on fresh citrated plasma or in bovine fibrinogen, which corresponds to the results obtained in previous studies (Table I).

In the evaluation of the anticoagulant activity, the *P.l.rozei* venom showed this in vitro action on fresh citrated plasma (uncoagulable at 300 s). This result was compatible with the one obtained for *P.l. lansbergii* (25) (Table I). Anticoagulant enzymes including protein C activators, factor IX/X, factor Xa and thrombin inhibitors, phospholipases A₂ (PLA₂) have also been described in snake venoms. These proteases inhibit coagulation by enzymatic or non-enzymatic mechanisms (39). The components responsible for the *P.l.rozei* venom anticoagulant activity remains to be identified.

In snake venoms there are compounds with the ability to activate the fibrinolytic system, they can be classified as α or β fibrinogenases, depending on their activity preferentially on one or another fibrinogen chain, and they belong to both metalloprotease and serine protease enzymes (40). In the present study, the fibrinolytic activity (mm²/μg) was found, that the *P.l.rozei* venom induced a lysis area larger as the reported for *P.l. hutmanni* (8) (Table I). This fibrinolytic activity suggests that several venoms contain serine or metalloproteases acting directly on fibrin. These proteases are of great interest since they could have clinical relevance as thrombolytic agents (41).
Biochemical and toxinological characteristics of *P. l. rozei* venom

Fibrinogen, a dimeric plasma glycoprotein of 340 kDa constituted of six polypeptide chains (α, β, γ), contributes in the coagulation structure, and is transformed by thrombin to fibrin monomers (49). This protein has been employed for the estimation of the possible fibrinogenolytic influences of natural toxins, and for the confirmation of anticoagulant properties of Viperidae venoms (27, 28). A variety of these proteins have been utilised in preclinical medical investigation, to test their probable therapeutic potential as anticoagulants or auxiliary in thrombolytic therapy (51).

Regarding to the fibrinogenolytic activity, the degradation of the fibrinogen chains (α, β and γ) by the venom of *P. l. rozei* occurred as follows: the α chain was completely degraded at 30 min, whereas the degradation of the β chain was observed at 2 hours. The degradation of γ chain by the *P. l. rozei* venom was not observed. To date there have been only a few reports of fibrinogenolytic enzymes of snake venoms acting on the γ chains (42-45).

In addition to the fibrinogenolytic degradation, fibronectin degradation was observed. This adhesive protein is a gel band with the highest molecular weight, and its degradation from 5 min of incubation until 24 hours was observed. Lucena et al. (46), by an immunoblotting tests, identified this band as contaminating protein in the commercial fibrinogen used in the assays. This activity on fibronectin demonstrates the *P. l. rozei* venom effects on extracellular matrix components, and therefore, corroborates the presence of metalloproteases in this venom and its haemorrhagic activity.

On the other hand, in the evaluation of the caseinolytic activity, the zymogram corresponding to the *P. l. rozei* venom did not show casein digestion bands, demonstrating that this venom lacks caseinolytic activity, unlike the *B. colombiensis* positive control, which had at least four digestion bands corresponding to high molecular weights areas.

Phospholipases (PLA₂) are calcium-dependent enzymes that catalyse the hydrolysis of the sn-2 ester bond of phosphoglycerides leading to the formation of free fatty acids and lysophospholipids. In addition to its role in the digestion of prey, their activities is associated with neurotoxicity, myotoxicity, inhibition of coagulation and platelet aggregation, as well as inflammation, hypotension, haemolysis, haemorrhage and contribute to the development of oedema (33, 34). Here, regarding the phospholipase A₂ activity, clear areas were evident in the agarose-egg yolk gel. The two venoms tested showed clear zones close to 15 kDa, which corresponds to the PLA₂ activity on the lecithin present in the gel. Although equal amounts of venom were placed, the light areas patterns with 10 μg were different. The *B. colombiensis* venom presented a greater area, a fact that makes us think that *P. l. rozei* venom has a lower enzyme concentration or less potent enzymes than the *B. colombiensis* venom. Phospholipase A₂ activity has been reported in other *Porthidium* species, for instance, the *P. l. lansbergii* venom (25), besides *P. nasutum* and *P. ophryomelas* (33). With respect to the recognition of the *P. l. rozei* venom by polyvalent antivenom serum (PAOS), the agarose gel double immunodiffusion assays showed precipitation bands or immune complexes, which were observed as a continuous line of immunoprecipitation showing complete identity.

The recognition pattern presented in this work suggests an immunological identity between the components of the *P. l. rozei* venom and the toxins of *Crotalus, Bothrops* and *P. hutmanni*.

Finally, in the literature, cases of ophidian accident by *Porthidium* snakes are limited albeit accidents are not uncommon. Cornejo et al. (49) reported a case produced by *P. hutmanni*, in Nueva Esparta state (Venezuela) with local manifestations that included sudden pain and local bite bleeding, as well as oedema, erythema and burnt sensation.
In conclusion, *P. l. rozei* venom has been biochemical and toxinologically studied and the toxinological and enzymatic functions, which included lethality (LD$_{50}$), haemorrhagic (MHD), myotoxic, coagulant, oedematising, proteolytic, fibrinogenolytic, fibrinolytic, anticoagulant, phospholiphasic activities and the crude venom recognition by the Venezuelan polyvalent antivenom (PAOS) were described. Here is proposed, according regional physicians that antivenom is a definite, complete, and effective treatment for *P. l. rozei* envenomations, since administration of PAOS antivenom conduct to ample clinical recuperation, lacking progression to multiple organ dysfunction syndromes. Therefore, antivenom efficiently act toward the acute symptoms and can prevent envenoming development (48).

The venom presented a haemostatic, toxinological and enzymatic biochemical profile comparable to the showed for specimens of the *Porthidium* genus, although with palpable quantitative disparities in their diverse activities. The diagnosis and treatment of *Porthidium* bites is on occasion challenging for clinicians because adequate information has not been offered in clinical practice. There are no fixed diagnostic markers in this practice; therefore, conclusive diagnosis of snake-venom envenoming involves confident identification of the snake and surveillance of the clinical manifestations of envenomation (10).

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