

Snake venomics of the South and Central American Bushmasters. Comparison of the toxin composition of *Lachesis muta* gathered from proteomic versus transcriptomic analysis

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ABSTRACT

We report the proteomic characterization of the venoms of two closely related pit vipers of the genus Lachesis, L. muta (South American Bushmaster) and L. stenophrys (Central American Bushmaster), and compare the toxin repertoire of the former revealed through a proteomic versus a transcriptomic approach. The protein composition of the venoms of Lachesis muta and L. stenophrys were analyzed by RP-HPLC, N-terminal sequencing, MALDI-TOF peptide mass fingerprinting and CID-MS/MS. Around 30-40 proteins of molecular masses in the range of 13-110 kDa and belonging, respectively, to only 8 and 7 toxin families were identified in L. muta and L. stenophrys venoms. In addition, both venoms contained a large number of bradykinin-potentiating peptides (BPP) and a C-type natriuretic peptide (C-NP). BPPs and C-NP comprised around 15% of the total venom proteins. In both species, the most abundant proteins were Zn²⁺-metalloproteinases (32–38%) and serine proteinases (25-31%), followed by PLA₂s (9-12%), galactose-specific C-type lectin (4-8%), L-amino acid oxidase (LAO, 3-5%), CRISP (1.8%; found in L. muta but not in L. stenophrys), and NGF (0.6%). On the other hand, only six L. muta venom-secreted proteins matched any of the previously reported 11 partial or full-length venom gland transcripts, and venom proteome and transcriptome depart in their relative abundances of different toxin families. As expected from their close phylogenetic relationship, the venoms of L. muta and L. stenophrys share (or contain highly similar) proteins, in particular BPPs, serine proteinases, a galactose-specific C-type lectin, and LAO. However, they dramatically depart in their respective PLA₂ complement. Intraspecific quantitative and qualitative differences in the expression of PLA₂ molecules were found when the venoms of five L. muta specimens (3 from Bolivia and 2 from Peru) and the venom of the same species purchased from Sigma were compared. These observations indicate that these class of toxins represents a rapidly-evolving

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gene family, and suggests that functional differences due to structural changes in PLA_{2s} molecules among these snakes may have been a hallmark during speciation and adaptation of diverging snake populations to new ecological niches, or competition for resources in existing ones. Our data may contribute to a deeper understanding of the biology and ecology of these snakes, and may also serve as a starting point for studying structure–function correlations of individual toxins.

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1. Introduction

Venom toxins likely evolved from proteins with a normal physiological function and appear to have been recruited into the venom proteome before the diversification of the advanced snakes, at the base of the Colubroidea radiation [1-4]. Given the central role that diet has played in the adaptive radiation of snakes [5], venoms represent the critical innovation that allowed advanced snakes to transition from a mechanical (constriction) to a chemical (venom) means of subduing and digesting prey larger than themselves, and as such, venom proteins have multiple functions including immobilizing, paralyzing, killing and digesting prey. Venoms produced by snakes of the family Viperidae (vipers and pit vipers) contain proteins that interfere with the coagulation cascade, the normal haemostatic system and tissue repair, and human envenomations are often characterized by clotting disorders, hypofibrinogenemia and local tissue necrosis [6,7]. In spite of the fact that viperid venoms may contain well over 100 protein components [8], venom proteins belong to only a few major protein families, including enzymes (serine proteinases, Zn²⁺-metalloproteinases, L-amino acid oxidase, group II PLA₂) and proteins without enzymatic activity (disintegrins, Ctype lectins, natriuretic peptides, myotoxins, CRISP toxins, nerve and vascular endothelium growth factors, cystatin and Kunitz-type proteinase inhibitors). However, snake venoms depart from each other in the composition and the relative abundance of toxins [1,2,8-14].

In addition to understanding how venoms evolve, characterization of the protein/peptide content of snake venoms also has a number of potential benefits for basic research, clinical diagnosis, development of new research tools and drugs of potential clinical use, and for antivenom production strategies [15]. Within- and between-species heterogeneity of venoms may also account for differences in the clinical symptoms observed in accidental envenomations. In order to explore the putative venom components, several laboratories have carried out transcriptomic analyses of the venom glands of viperid (Bitis gabonica [16], Bothrops insularis [17], Bothrops jararacussu [18], Bothrops jararaca [19], Agkistrodon acutus [20,21], Echis ocellatus [22], and Lachesis muta [23]), elapid (Oxyuramus scutellatus [24]), and colubrid (Philodryas olfersii [25]) snake species. Transcriptomic investigations provide catalogues of partial and full-length transcripts that are synthesized by the venom gland. Here, we report a proteomic analysis of L. muta venom, which complements the study of snake venom gene transcriptional activity (transcriptome) in the same species [23] by showing the relative abundance of the

various protein families that are actually secreted into the venoms.

Lachesis is a genus of venomous pit vipers widely distributed in remote, lowland tropical forested areas in Central and South America, and the only neo-tropical pit viper that lays eggs. Three species are currently recognized in this genus, *L. muta* (South American bushmaster), *L. stenophrys* (Central American bushmaster), and *L. melanocephala* (Black-headed bushmaster) (http://www.reptile-database.org) . The bushmasters are the largest of all pit vipers and the longest venomous snakes in the western hemisphere. Adults grow to an average of 2 to 2.5 m, although 3 m is not too unusual.

Lachesis melanocephala is endemic of the pacific versant of Costa Rica, where it is confined to the southwestern Osa Peninsula [26]. L. muta is found in South America in the equatorial forests east of the Andes, ranging from Colombia, eastern Ecuador, Peru, northern Bolivia, eastern and southern Venezuela, to Guyana, Surinam, French Guiana and much of northern Brazil. Lachesis stenophrys, is found in the Atlantic lowlands of southern Nicaragua, Costa Rica and the Atlantic and Pacific lowlands of central and eastern Panama. In South America it occurs in the Pacific lowlands of Colombia and northwestern Ecuador, the Caribbean coast of northwestern Colombia and inland along the Magdalena and Cauca river valleys.

Human envenoming by Lachesis (muta or stenophrys) are infrequent but rather severe and characterized by conspicuous local tissue damage (edema, hemorrhage and necrosis), nausea, coagulopathies, hypotension, shock and renal disturbances [27]. Brown [28] mentions a venom yield of 200-411 mg from L. muta and gives the following LD_{50%} values for mice: 1.5 mg/kg (i.v.), 1.6-6.2 mg/kg (i.p.), 6.0 mg/kg (s.c.). Paradoxically, although bites can be deadly, snake venoms also contain components of theraupeutic value. The venom of L. muta has attracted medical interest because its reported protective effect in rats subjected to high cytostatic doses, when administered at low (4 ng/ml) concentration in combination with Mg, Se, Zn (4 μ g/ml each) [29]. It has been also reported that administration of this formulation to nude mice that had developed tumors by inoculation of PANC-1 cells, inhibited tumor growth and angiogenesis, induced apoptosis, and modulated the activity of antioxidant enzymes [30]. Ongoing research from our laboratories, which will be published elsewhere, indicates that daily subcutaneous administration of L. muta venom (0.5 ml, 4 ng/ml) significantly (p<0.05) increased the survival of N-nitroso-N-methylureainduced tumor-bearing rats (103 days) compared to non-Lmtreated animals (66 days). Furthermore, the venom provoked

Table 1 – Assignment of the reverse-phase fractions of *Lachesis muta* venom (Santa Cruz de la Sierra, Bolivia), isolated as in Fig. 1, to protein families by N-terminal Edman sequencing, mass spectrometry, and collision-induced fragmentation by nESI-MS/MS of selected peptide ions from in-gel digested protein bands (separated by SDS-PAGE as in Fig. 2)

HPLC fraction	N-terminal sequencing	Isotope-averaged molecular	l Peptide ion		Peptide ion		MS/MS-derived sequence	Protein family
Lm-		111255	m/z	Z				
1	N.D.	384.0		2	Ac-TPPAGPD(+41)	[Q27J49]		
		465.6		2	TPPAGPDVGP(+91)	BPP-C-NP precursor		
		581.8		2	DHHAVGGGGGGGGGA(+91)			
2	N.D.	480.6		2	ZKKWPPGH	[Q27J49]		
3	N.D.	532.1		2	TPPAGPDVGPR	[Q27J49]		
		482.0		2	PPAGPDVGPR			
		768.5		1	AGPDVGPR			
		697.4		1	GPDVGPR			
4 5		543.5		1	DVGPR			
4, 5	n.p.	700 6		0		[007140]		
6	N.D.	702.6		2	ZKKWPPGHHIPP	[Q27]49]		
7	ND	/64.1		1	ZKKWPP	[007]40]		
/	N.D.	597.2		2	ZKKWPPGHHI	[Q27]49]		
		549.0		2				
0	ND	302.9 705 5		2		[027]40]		
0	N.D.	620.2		2 2		[027]49]		
5 10	N.D.	622.1		2		[027]49]		
10	N.D.	544.1		2		[Q27]49]		
		945 5		2				
		1032 5		1	ZKWDPPPIS			
11	ND	868 7		3	ZKWDPPPISPPLIKPHES	[027]49]		
**	11.2.	000.7		5	PAGGTTA			
		845 1		3	ZKWDPPPISPPLLKPHE			
		01012		0	SPAGGTT			
		758.7		3	ZKWDPPPISPPLLKPHESPAG			
		653.0		3	ZKWDPPPISPPLLKPHE			
		611.5		3	ZKWDPPPISPPLLKPH			
		688.7		3	PPPISPPLLKPHESPAGGTTA			
		578.6		3	PPPISPPLLKPHESPAG			
12	N.D.	715.8		3	ZKWDPPPISPPLLKPHESP	[Q27J49]		
		679.3		2	ZKWDPPPISPPL			
13	GDGCFGLKLDRIGSMSGLGC	1983.4				C-NP [Q27J49]		
	GCFGLKLDRIGSMSGLGC	1811.6						
14	Blocked	12041, 12154				unknown		
15	Blocked	16 kDa 💵	556.2	2	NPNPVPTGCR	Nerve growth factor		
			632.1	2	IDAACVCISR			
			682.1	2	ALTMEGNQASWR			
16	HLLQFGDLINKIARRNGIS-	13932.6	649.6	2	HLLQFGDLINK	PLA ₂		
	-YYGFYGCYCGL		605.3	3	EICECDRDAAICFR			
17	HLLQFGDLINKIARRNGIS-	13916.3	649.1	2	HLLQFGDLINK	PLA ₂		
	-YYGFYG		605.3	3	EICECDRDAAICFR			
			490.7	2	EICECDR			
			548.7	2	DNLDTYDNK			
			752.7	2	CCFVHDCCYGK			
			512.7	2	YWFFHPK			
40			507.9	2	GRPQDATDR			
18		14009 6				DI A		
	-GCICGLGGQGKPQDA	14008.0				CDISD		
10		24003.9 20 kDa				2 chain for proteiness		
19	ALCENTINEUROPACIONSCIENCE	50 KDa				2-chain ser-proteinase		
	SADEDZEZBKAEIU	24683.8						
20	IICCDECNINEHRSI	24005.0 28 kDa	647 5	2	XNXXDYFVCP	Ser-proteinase		
20	HOOPEGIAIIAFLIKOF	20 KDa	763.8	2	IIGGDECNINEHR	Ser proteniase		
	VEGGDECNINEHRSI	26 kDa 🗖	608.1	2	KVPNKDEETR	Ser-proteinase [O27I47]		
	. Jobbonn Brindl	20 1124	714.9	2	SLPSSPPSVGSVCR	Ser proteinase [Q2/)T/]		
			773.4	2	VEGGDECNINEHR			
	SVDFDSESPRKPEIO	24 kDa 🗖	,, 5.1	2		CRISP		
		-						

Table 1 (d	continued)					
HPLC fraction	N-terminal sequencing	Isotope-averaged molecular	l Pepti ion	de 1	MS/MS-derived sequence	Protein family
Lm-		111855	m/z	Z		
21	VFGGDECNINEHRSL	27 kDa 🗖				Ser-proteinase [Q27J47]
22	M: NNCPQDWLPMNG	28 kDa ■/				Gal-lectin [Q9PSM4]
	m: VIGGDECNINEHR	14 kDa ↓ 29 kDa ▼				Ser-proteinase [P33589]
23	FLVALY M: VIGGDECNINEHRFLVALYDG- -LSGTFLCG	30 kDa 🔻				Ser-proteinase [P33589] Venombin A
	m: VFGGDECNINEHRSLVVLFNS- -SGFLCAGT	30 kDa 🔻				Ser-proteinase [Q27J47]
23–29	NNCPQDWLPMNGL CYKIFD	28 kDa 🗖	639.6	2	LWNDQVCESK	Gal-lectin [Q9PSM4]
		14 kDa 🔻	620.9	2	DFSWEWTDR	
			644.3	2	SCTDYLTWDK	
			005.8 796.9	2		
			639.3	2	YGESLEIAEYISDYHK	
24	IVGGDECNINEHRFL VALYDP	30 kDa ▼	00010	5		Serine proteinase
25	VIGGDECNINEHRF LVALYD	30 kDa 🔻				Serine proteinase [P33589]
26, 27	VIGGDECNINEHRSL VALYD	38 kDa 🔻	575.5	2	NVKFDDEQR	Serine proteinase Venombin A
			756.9	2	VIGGDECNINEHR	(S35689)
			867.3	2	VLCAGVLEGGIDTCNR	
	VECCDECNINELID	21 kDa 🔻	772.8	2	SLPSNPPSEDSVCR	Soring protoinage (D84026)
	SLVVLFN	SI KDa	711.5	2		Planning proteinase (r64030)
00		21 hDa 🔻	/11.5	2	AIYPEFGLPATSR	Plasminogen-activator
28	VFGGDEGNINEHRSLVV	31 KDa	791.4	2	(261 2)PSSPPSVCSVCR	Serine proteinase
28–38		29 kDa 🗖	, 11.0	2	(201.2)10011010000	
		14 kDa 🔻	459.8	3	GHCYKPFNEPK	C-type lectin-like
29	ADDRNPLGECFRETD YEEFLEIAK	58 kDa ■▼	583.1	2	KFWEDDGIR	L-amino acid oxidase
			641.0	2	SAGQLYEESLGK	
			744.1	2	ETDYEEFLEIAK	
30	(NFPPYEANIMRV)	26 kDa ■▼	781.3	2	VHEIVNTLNGFYR	PI-metalloproteinase(P22796)
			590.1	2	TFGEWRER	Hemorrhagic factor LHFII
			869 5	2	VIELVAVADHCMETK	
31	Blocked	26 kDa ■▼	781.3	2	VHEIVNTLNGFYR	PI-metalloproteinase (P22796)
			590.1	2	TFGEWRER	Hemorrhagic factor LHFII
			641.1	2	NSVGIVQDHSPK	C C
			869.5	2	YIELVVVADHG <u>m</u> FTK	
32, 33	Blocked	37 kDa ■▼	518.8	2	SVGIVQDYR	PIII-Metalloproteinase
			672.2	3	LTPGSQCADGECCDQCR	
			6/1.3	2	YIELVLLADHK	DI motolloprotoinago
		20 KDa	701.5	2	TECEWDED	(P22796)
34	Blocked	58 kDa ■▼	752.6	2	XFCEFNNFPCR	PIII-Metalloproteinase
51	biocheu	50 KDu	650.0	2	YVEXVVVADHR	
35	Blocked	27 kDa ■▼	577.6	2	ZVVTAEQQR	PI-Metalloproteinase
			604.6	2	QGAQCAEGLCCDQCR	
			530.3	2	XACEPQDVK	
			627.2	2	PQCXXQQXPR	
		_	635.6	2	LYCFPSSPATK	
36	Blocked	110 kDa =	761.9	2	SAAADTXEAFADWR	(PIII-Metalloproteinase) ₂
		49 kDa 🔻	594.9	3	(581.5)VVVADHN(467.0)	
						(continued on next page)

Table 1 (continued)							
HPLC fraction		N-terminal sequencing	Isotope-average molecular	ed Pepti ion	de 1	MS/MS-derived sequence	Protein family
			mass				
Lm-				m/z	Ζ		
37	Blocked		27 kDa ■▼	627.2	2	PQCXXQQXPR	PI-Metalloproteinase
				635.6	2	LYCFPSSPATK	
38	Blocked		48 kDa ■▼	657.1	2	YXEXVVVADHR	PIII-Metalloproteinase
				504.8	2	(182.6)YNQPSK	

X, Ile or Leu; Z, pyrrolidone carboxylic acid; Ac-, N-acetyl; *m*, methionine sulphoxide. Unless other stated, for MS/MS analyses, cysteine residues were carbamidomethylated. Molecular masses of native proteins were determined by electrospray-ionization (\pm 0.02%) or MALDI-TOF (*****) (\pm 0.2%) mass spectrometry. Apparent molecular mass determined by SDS-PAGE of non-reduced (\blacksquare) and reduced (\blacktriangledown) samples; n.p., non-peptidic material found. M and m, denote mayor and minor products co-eluting in the same HPLC fraction. Previously reported proteins are identified by their databank accession codes.

50–70% cell proliferation inhibition of cultured human neoplasia-derived cell lines MDA-MB-231 and MCF-7 (breast cancer), PANC-1 (pancreatic carcinoma), WM35 and HT168 (melanoma), and U937 (histiocytic lymphoma).

Otero and co-workers [31] have reported quantitative differences in toxic and enzymatic activities along with subtle variations in the electrophoretic patterns of L. muta and L. stenophrys venoms from Brazil, Colombia, and Costa Rica, although experimental envenomation by these venoms induced a qualitatively similar pathophysiological profile. Here, we report the proteomic characterization of the toxin composition of L. muta and L. stenophrys venoms, and compare the toxin repertoire of the former revealed through a proteomic versus a transcriptomic approach. Venom toxin composition provides a comprehensible catalogue of the venom-secreted proteins, which may contribute to a deeper understanding of the biology and ecology of the snake, the biological effects of the venom, and may also serve as a starting point for studying structure-function correlations of individual toxins.

2. Experimental section

2.1. Isolation and relative quantitation of venom proteins

Venom of L. stenophrys (Central American Bushmaster) was pooled from specimens collected in Costa Rica and kept at the serpentarium of the Instituto Clodomiro Picado, University of Costa Rica in San José. L. muta (South American Bushmaster) venom samples were obtained from 3 wild caught specimens and were kindly provided by SICAE' s.r.l., a snakes farm located in the nature reserve Potrerillos del Guendá (Santa Cruz de la Sierra, Bolivia, P.O. Box 1615) (www.sicae-online. com). Venoms from 2 specimens of L. muta from Peru (Iquitos, Departamento de Loreto or Cenepa, Mamayaque) were also analyzed. A further sample of L. muta venom was purchased from Sigma-Aldrich (Alcobendas, Madrid, Spain; catalog no. V7376). The source of this venom was not provided by the vendor. For reverse-phase HPLC separations, 2-5 mg of crude, lyophillized venom of L. muta were dissolved in 100 µl of 0.05% trifluoroacetic acid (TFA) and 5% acetonitrile, and insoluble material was removed by centrifugation in an Eppendorff centrifuge at 13,000 ×g for 10 min at room temperature.

Proteins in the soluble material were separated using an ETTAN™ LC HPLC system (Amersham Biosciences) and a Lichrosphere RP100 C_{18} column (250 × 4 mm, 5 μ m particle size) eluted at 1 ml/min with a linear gradient of 0.1% TFA in water (solution A) and acetonitrile (solution B) (5% B for 10 min, followed by 5-15% B over 20 min, 15-45% B over 120 min, and 45-70% B over 20 min). Protein detection was at 215 nm and peaks were collected manually and dried in a Speed-Vac (Savant). Given that the wavelength of absorbance for a peptide bond is 190-230 nm, protein detection at 215 nm allows to estimate the relative abundances (expressed as percentage of the total venom proteins) of the different protein families from the relation of the sum of the areas of the reverse-phase chromatographic peaks containing proteins from the same family to the total area of venom protein peaks in the reverse-phase chromatogram. In a strict sense, and according to the Lambert-Beer law, the calculated relative amounts correspond to the "% of total peptide bonds in the sample", which is a good estimate of the % by weight (gr/100gr) of a particular venom component.

2.2. Characterization of HPLC-isolated proteins

Isolated protein fractions were subjected to N-terminal sequence analysis (using a Procise instrument, Applied Biosystems, Foster City, CA, USA) following the manufacturer's instructions. Amino acid sequence similarity searches were performed against a non-redundant protein sequence databank (comprising all non-redundant GenBank CDS translations+RefSeq Proteins+PDB+SwissProt+PIR+PRF) (http:// www.ncbi.nlm.nih.gov/BLAST/blastcgihelp.shtml#protein_data bases) using the BLAST program [32] implemented in the WU-BLAST2 search engine at http://www.bork.embl-heidelberg.de. The molecular masses of the purified proteins were determined by SDS-PAGE (on 12-15% polyacrylamide gels) and by electrospray ionization (ESI) mass spectrometry using an Applied Biosystems QTrap™ 2000 mass spectrometer [33] operated in Enhanced Multiple Charge mode in the range m/z 600-1200.

2.3. In-gel enzymatic digestion and mass fingerprinting

Protein bands of interest were excised from a Coomassie Brilliant Blue-stained SDS-PAGE and subjected to automated reduction with DTT and alkylation with iodoacetamide, and digestion with sequencing grade bovine pancreas trypsin (Roche) using a ProGest™ digestor (Genomic Solutions) following the manufacturer's instructions. The tryptic peptide mixtures were dried in a Speed-Vac and redissolved in 5 µl of 70% acetonitrile and 0.1% TFA. Digests (0.65 µl) were spotted onto a MALDI-TOF sample holder, mixed with an equal volume of a saturated solution of α-cyano-4-hydroxycinnamic acid (Sigma) in 50% acetonitrile containing 0.1% TFA, dried, and analyzed with an Applied Biosystems Voyager-DE Pro MALDI-TOF mass spectrometer, operated in delayed extraction and reflector modes. A tryptic peptide mixture of Cratylia floribunda seed lectin (SwissProt accession code P81517) prepared and previously characterized in our laboratory was used as mass calibration standard (mass range, 450-3300 Da).

CID-MS/MS 24

For peptide sequencing, the protein digest mixture was loaded in a nanospray capillary column and subjected to electrospray ionization mass spectrometric analysis using a QTrap 2000 mass spectrometer (Applied Biosystems) [33] equipped with a nanoelectrospray source (Protana, Denmark). Doubly- or triplycharged ions of selected peptides from the MALDI-TOF mass fingerprint spectra were analyzed in Enhanced Resolution MS mode and the monoisotopic ions were fragmented using the Enhanced Product Ion tool with Q0 trapping. Enhanced Resolution was performed at 250 amu/s across the entire mass range. Settings for MS/MS experiments were as follows: Q1-unit resolution; Q1-to-Q2 collision energy — 30-40 eV; Q3 entry barrier — 8 V; LIT (linear ion trap) Q3 fill time — 250 ms; and Q3 scan rate — 1000 amu/s. CID spectra were interpreted manually or using a licensed version of the MASCOT program (http:// www.matrixscience.com) against a private database containing 927 viperid protein sequences deposited in the SwissProt/ TrEMBL database (Knowledgebase Release 12 of July 2007; http://us.expasy.org/sprot/). MS/MS mass tolerance was set to ±0.6 Da. Carbamidomethyl cysteine and oxidation of methione were fixed and variable modifications, respectively.

2.5. Variation in venom composition between Lachesis taxa

We used similarity coefficients to estimate the similarity of venom proteins between taxa. These coefficients are similar to the bandsharing coefficients used to compare individual genetic profiles based on multilocus DNA fingerprints [34]. We defined the Protein Similarity Coefficient (PSC) between two species "a" and "b" in the following way: $PSC_{ab} = [2 \times (no.$ of proteins shared between a and b)/(total number of distinct proteins in a+total number of distinct proteins in b)]×100. We judged two proteins (listed in Tables 1 and 2) as being different when they met one or more of these criteria: 1) Had different N-terminal sequences and/or distinct internal peptides sequences (derived from MS/MS data) corresponding to homologous regions; 2) had different peptide mass fingerprints; 3) were of different sizes (judged by MALDI-TOF MS or SDS-PAGE). For these comparisons, two proteins were judged to differ in size if they differed by more than our estimate of the 95% confidence interval for particular sizing techniques (0.01% for ESI-QTrap MS; 0.4% for MALDI-TOF MS-derived masses, and

+1.4 kDa for SDS-PAGE-determined masses); or 4) eluted in different reverse-phase HPLC peaks. We emphasize that these measures will give only minimum estimates of the similarities between the venom profiles. We suspect that a number of the proteins that we judge to be the same using the above criteria would be found to differ at one or more of these criteria if more complete information were available.

Results and discussion 3.

3.1. Characterization of bushmaster venom proteomes

The crude venoms of L. muta and L. stenophrys were fractionated by reverse-phase HPLC (Figs. 1 and 3), followed by analysis of each chromatographic fraction by SDS-PAGE (Figs. 2 and 4) and N-terminal sequencing. Molecular masses of purified proteins were determined by ESI-MS or MALDI-TOF mass spectrometry (Tables 1 and 2). Fig. 5A displays an example of electrospray-ionization mass spectrum of the 25 kDa protein isolated in fraction 18 (Figs. 1 and 2) and identified as a member of the CRISP family (Table 1). Protein fractions showing single electrophoretic band, molecular mass, and N-terminal sequence were straightforwardly assigned by BLAST analysis (http://www.ncbi.nlm.nih.gov/ BLAST) to a known protein family, indicating that representative members of most snake venom toxin families are present amongst the 927 viperid protein sequences deposited to date in the SwissProt/TrEMBL database. Protein fractions showing heterogeneous or blocked N-termini were analyzed by SDS-PAGE and the bands of interest are subjected to automated reduction, carbamidomethylation, and in-gel tryptic digestion in a ProGest digestor (Genomic Solutions). The resulting tryptic peptides are then analyzed by MALDI-TOF mass fingerprinting followed by amino acid sequence determination of selected doubly- and triply-charged peptide ions by collision-induced dissociation tandem mass spectrometry. Product ion spectra were manually or using either the online form of the MASCOT program (searching against the nonredundant MSDB database) or a licensed version of this program against a private snake venom database comprising 927 sequence entries (212 in SwissProt, 715 in TrEMBL) plus the previously assigned peptide ion sequences from snake venomics projects carried out in our laboratory [9-14]. Fig. 5B illustrates the *de novo* sequencing of a doubly-charged tryptic peptide ion (m/z 774.1) from protein Lm29 (Figs. 1 and 2) identified as an L-amino acid oxidase (Table 1). The outlined snake venomics approach allowed us to assign unambiguously all the isolated venom toxins representing more than 0.05% of the total venom proteins (i.e. less than 50 ng in 100 μ g of venom proteins) to known protein families (Tables 1 and 2).

Supporting the view that venom proteomes are mainly composed of toxins belonging to a few protein families [4,9-14,16-24], the proteins found in the venoms of L. muta and L. stenophrys cluster, respectively, in 8 and 7 different families (bradykinin-potentiating peptides, NGF, PLA2, serine proteinase, cysteine-rich secretory proteins (CRISP; only found in L. muta), C-type lectins, L-amino acid oxidase (LAO), and Zn²⁺dependent metalloproteinases) (Fig. 6), whose relative abundances are listed in Table 3.

Table 2 – Assignment of the reverse-phase fractions of Lachesis stenophrys venom (Costa Rica), isolated as in Fig. 3, to protein families by N-terminal Edman sequencing, mass spectrometry, and collision-induced fragmentation by nESI-MS/MS of selected peptide ions from in-gel digested protein bands (separated by SDS-PAGE as in Fig. 4)

HPLC fraction	N-terminal sequencing	Isotope- averaged	Pepti ion	de	MS/MS-derived sequence	Protein family
Ls-		molecular mass	m/z	Z		
1	N.D.	480.6		2	ZKKWPPGH	[Q27J49]
2	N.D.	532.4 482.1		2 2	TPPAGPDVGPR PPAGPDVGPR	[Q27J49]
3, 4	n.p.	700.0		0		[007140]
5	N.D.	702.8		2	ZKKWPPGHHIPP	[Q27]49]
		549.0		2	ZKKWPPGHH	
		530.1		2	WPPGHHIPP(+41)	
6	N.D.	639.1		2	ZEWPPGHHIPP	[Q27J49]
		622.1		2	PRPQIPPLVVQ	
7	N.D.	748.4		2	SHKGWPPRPQIPP	[Q27J49]
		572.9		2	GWPPRPQIPP	
0	ND	649.8		2	WPPRPQIPPLV	[007140]
ŏ	N.D.	623.1 542.4		2		[Q27]49]
9	GDGCEGI KI DRIGSMSGI GC	1983 1		Z	WITKIQIII	C-NP [027149]
10	Blocked	16 kDa ■▼	556.2	2	NPNPVPTGCR	Nerve growth factor
			632.1	2	IDAACVCISR	0
			682.1	2	ALTMEGNQASWR	
11	N.D.	15 kDa 🔻	649.6	2	HLLQFGDLIDK	PLA ₂
12	HLLQFGDLIDKIAGR	14052	649.7	2	HLLQFGDLIDK	PLA ₂
13	HLLQFGDLIDKIAGR	13898	649.7	2	HLLQFGDLIDK	PLA ₂
			753.3	2	CCFVHDCCYGK	
14		07 hp.	605.4	3	EICECDRDAAICFR	
14	IIGGDECNINEHRFL	37 KDa - *	64/./ 762.9	2	XNXXDYEVCK	Serine proteinase
		13 kDa ▼	/05.0	Z	IIGGDEGNINEAK	
15	(V/I)(V/I)GGDECNINEHRFL	15 kDa 34 kDa ■▼	647.7	2	XNXXDYEVCR	Serine proteinase
			763.8	2	IIGGDECNINEHR	I
			621.9	2	TGXWGXR	
16	(V/I)(V/I)GGDECNINEHRFL	32 kDa 🗖	756.8	2	VIGGDECNINEHR	Serine proteinase
			683.4	2	(292.2)PEFGLPATSR	
		_	715.3	2	SXPSSPPSVGSVCR	
17	M: NNCPQDWLPMNGLCY	28 kDa	670.6	3	NNCPQDWLPMNGLCYK	Gal-lectin [Q9PSM4]
		14 kDa 🔨	639.6	3	YGESLEIAEYISDYHK	
			786.8	2	EFGVELVSLIGYR	
			730.8	2		
			765.4	2	GOAEVWIGLWDKK	
			657.7	2	AWEDAEMFCR	
			621.1	2	DFSWEWTDR	
			685.3	2	KDFSWEWTDR	
		_	800.8	2	KYKPGCHLASFHR	
	m: VIGGDECNINEHRFL	34 kDa 🔻				Serine proteinase
18	IVGGDECNINEHRFL	34+26 kDa ▼	756.9	2	IVGGDECNINEHR	Serine proteinase
10	ND		711.5	2	AIYPEFGLPATSR	Carrier and the second
19	N.D.	29 KDa - V	690.4 715.2	2	(306.4)PEFGLPAISK	Serine proteinase
20	VEGGDECININEARTE	55+24 KDa	647.7	2	XNXXDYEVCR	Serine proteinase
21	VIGGDECNINEHRSLVALYD	38 kDa ■▼	575.5	2	NVKFDDEOR	Ser-proteinase Venombin A
		50 112 4	756.9	2	VIGGDECNINEHR	(S35689)
			867.3	2	VLCAGVLEGGIDTCNR	
			710.9	2	SLMNIYLGMHNK	
			772.8	2	SLPSNPPSEDSVCR	
	VFGGDECNINEHRSLVVLFN	31 kDa ■▼	773.9	2	VFGGDECNINEHR	Ser-proteinase (P84036)
			711.5	2	AIYPEFGLPATSR	Plasminogen-activator
22	VFGGDECNINEHRSLVV	32 kDa 🔻	715.3	2	SLPSSPPSVGSVCR	Serine proteinase
			64/./	2	XNXXDYEVCK	
			772.0	2	VEGGDECNINEHP	
	VVGGDECNINEHR	31 kDa ▼	749.9	2	VVGGDECNINEHR	Serine proteinase

Table 2 (continued)						
HPLC fraction	N-terminal sequencing	Isotope- averaged	Peptide ion		MS/MS-derived sequence	Protein family
Ls-		molecular mass	m/z	Z		
23	N.D.	32 kDa 🔻	488.2	2	ETYPNVPR	Serine proteinase
			618.8	2	XNXXDYAVCR	
			744.6	2	(261.2)PSSPPSVGSVCR	
			791.8	2	CANXNXXDYXVCR	
			824.8	2	NDTEWDKDXMXXR	
24,25	ADDRNPLGECFRETDYEEFL	58 kDa 🗖 🗸	583.1	2	KFWEDDGIR	L-amino acid oxidase
			641.0	2	SAGQLYEESLGK	
			744.0	2	ETDYEEFLEIAK	
26	N.D.	110+52 kDa ■▼	532.3	2	YNGNXNTXR	PIII-metalloproteinase
27	N.D.	52 kDa 🗖 🗸	532.3	2	YNGNXNTXR	PIII-metalloproteinase
		27 kDa▼	605.3	2	DYYEMFXTK	PI-metalloproteinase
			640.8	2	NSVGXVQDHSPK	
		15 kDa▼	621.3	2	DFSWEWTDR	Gal-lectin [Q9PSM4]
			786.9	2	EFCVELVSTGYR	
28–31	Blocked	27 kDa▼	802.6	2	VHEXVNTXNVFYR	PI-metalloproteinase
			605.3	2	DYYEMFXTK	~ (P22796)
			540.3	2	TFGEWRER	
			640.8	2	NSVGXVQDHSPK	
			861.5	2	YXEXVVVADHGMFTK	
			532.8	2	YNGNXNTXR	
			683.4	3	TLLIAVTMAHELGHNLGMK	
29	Blocked	52 kDa 🗖 🗸	526.7	2	GNYYGYCR	PIII-metalloproteinase
			801.3	2	MYEXANTVNDXYR	
			684.8	3	XTVKPEAGYTXNAFGEWR	
30	Blocked	110 kDa ■ 48 kDa ▼	635.8	2	XYCFPSSPATK	(PIII-Metalloproteinase) ₂
31	Blocked	52 kDa ■▼	506.2	2	FTSAGNVCR	PIII-metalloproteinase
			650.2	2	YVEXVVVADHR	1
			752.4	2	XFEFNNFPCR	
32	N.D.	110 kDa 🗖	752.4	2	XFEFNNFPCR	(PIII-Metalloproteinase) ₂
		52 kDa 🔻	506.2	2	FTSAGNVCR	, i i i i i i i i i i i i i i i i i i i

X, lle or Leu; Z, pyrrolidone carboxylic acid; Ac-, N-acetyl; *m*, methionine sulphoxide. Unless other stated, for MS/MS analyses, cysteine residues were carbamidomethylated. Molecular masses of native proteins were determined by electrospray-ionization (\pm 0.02%) or MALDI-TOF (*) (\pm 0.2%) mass spectrometry. Apparent molecular mass determined by SDS-PAGE of non-reduced (\blacksquare) and reduced (\blacktriangledown) samples; n.p., non-peptidic material found. M and m, denote mayor and minor products co-eluting in the same HPLC fraction. Previously reported proteins are identified by their databank accession codes.

Except for the absence of a CRISP molecule in the venom of *L*. stenophrys, the two Lachesis species investigated show very similar overall venom toxin compositions (Table 3). However, comparison of the chromatographic separations of the venom proteins from *L*. muta and *L*. stenophrys (Figs. 1 and 3) and the tryptic peptide mass fingerprints of their individual protein bands (Tables 1 and 2), evidenced both, a number of very similar (or identical) proteins but also toxins from the same family showing a large degree of structural divergence.

Identical L. muta and L. stenophrys venom components include a number of bradykinin-potentiating peptides (BPPs) and a C-type natriuretic peptide released from the 239-aminoacid precursor protein Q27J49 (Fig. 7). BPPs found in fractions Ls7, Lm9/Ls6, Lm6/Ls5, and Lm10/Ls8 have been previously identified by MALDI-TOF MS in the crude venom of a specimen kept in captivity at the serpentarium of the Fundação Ezequiel Dias (Belo Horizonte, Brazil) [35], indicating that expression of these peptides appear not to exhibit geographical variation. BPPs have been described as snake venom inhibitors of the

angiotensin-converting enzyme, a dipeptidylcarboxypeptidase expressed in endothelial, epithelial and neuroepithelial cells, which converts inactive angiotensin I into the potent vasoconstrictor angiotensin II, and degrades bradykinin into bradykinin (1-7) or bradykinin (1-5) [36]. BPPs prevent the hypertensive effect of the angiotensin II and potentiate the hypotensive effect of the circulating bradykinin. C-natriuretic peptides elicit natriuretic, diuretic, and vasorelaxant activities. Lachesis protein Q27J49 encodes BPPs and a C-natriuretic peptide, combining in one precursor molecule two kinds of vasoactive molecules. Vasodilatation and hypotension contribute synergistically to overall venom toxicity evoking the rapid diffusion of toxic substances in the circulatory system and a hypotensive shock, which is a major cause of death of the prey or victim induced by viper snake bites. The BPPs were also essential for the development of the first commercial ACE inhibitor, captopril, for the treatment of human hypertension [37]. Although L. muta and L. stenophrys express similar relative amounts of BPPs into their venoms, each snake showed



Fig. 1 – Reverse-phase HPLC separation of the Lachesis muta venom proteins. Two milligrams of Lachesis muta venom (Santa Cruz de la Sierra, Bolivia) were applied to a Lichrosphere RP100 C_{18} column, which was then developed with the following chromatographic conditions: isocratically (5% B) for 10 min, followed by 5–15% B for 20 min, 15–45% B for 120 min, and 45–70% B for 20 min. Fractions were collected manually and characterized by N-terminal sequencing, ESI mass spectrometry, tryptic peptide mass fingerprinting, and CID-MS/MS of selected doubly- or triply-charged peptide ions. The results are shown in Table 1.

distinct complements of Q27J49-derived peptides, suggesting that the processing steps required to form the mature BPPs are overlapping though not identical in the two Lachesis species. However, whether the occurrence of distinct sets of BPPs reflects an evolutionary adaptation or merely a neutral consequence of speciation deserves further detailed investigations.



Fig. 3 – Reverse-phase HPLC separation of the venom proteins of *Lachesis stenophrys*. Two milligrams of *L. stenophrys* venom (Costa Rica) were applied to a Lichrosphere RP100 C_{18} column, which was then developed as in Fig. 1. Fractions were collected manually and characterized by N-terminal sequencing, ESI mass spectrometry, tryptic peptide mass fingerprinting, and CID-MS/MS of selected doubly- or triply-charged peptide ions. The results are shown in Table 2.

Through the pathophysiological consequences of the presence of large amounts of BPPs in Lachesis venoms deserve further and detailed consideration, the large content of BPPs in the two Lachesis venoms investigated may be associated with the conspicuous hypotension of very rapid onset which characterizes bushmaster envenomation cases, an effect that is likely to contribute to hemodynamic complications leading to cardiovascular shock [27].



Fig. 2–SDS-PAGE of reverse-phase separated fractions from the venom of *Lachesis muta* (Santa Cruz de la Sierra, Bolivia). SDS-PAGE showing the protein composition of the reverse-phase HPLC separated venom protein fractions run under non-reduced (panel A) and reduced (panel B) conditions. Molecular mass markers (in kDa) are indicated at the left of each gel. Protein bands were excised and characterized by mass fingerprinting and CID-MS/MS. The results are shown in Table 1.



Std 10 11 12 13 14 15 16 17 18 19 20 21 Std 22 23 24 25 26 27 28 29 30 31 32

Fig. 4–SDS-PAGE of reverse-phase separated fractions from the venom of Lachesis stenophrys (Costa Rica). SDS-PAGE showing the protein composition of the reverse-phase HPLC separated venom fractions (see Fig. 3) run under non-reduced (panel A) and reduced (panel B) conditions. Molecular mass markers (in kDa) are indicated at the left of each gel. Protein bands were excised and characterized by mass fingerprinting and CID-MS/MS. The results are shown in Table 2.

Furthermore, the low molecular mass of these peptides is likely to confer them with very low antigenicity, with the consequent implications for antivenom development. It would be relevant to assess whether antivenoms are effective at binding and neutralizing BPPs present in Lachesis venoms.

Other very similar or identical proteins in *L. muta* and *L. stenophrys* venoms, based on their similar chromatographic



Fig. 5 – Mass spectrometric characterization of isolated proteins and tryptic peptides. (A) Electrospray-ionization mass spectrum of the major protein isolated in HPLC fraction 18 (Figs. 1 and 2, Table 1). From the series of ions $(M+16H)^{16+}-(M+21H)^{21+}$ an isotope-averaged molecular mass of 24683.9±2.1 Da, and was identified by N-terminal sequencing as a member of the CRISP family. (B) MS/MS spectrum of the doubly-charged tryptic parent ion of m/z 744.1 (encircled) from the 58 kDa protein isolated in fraction Lm29 of Lachesis muta venom HPLC separation (Figs. 1 and 2, Table 1). Ions of the major sequence-specific y-ion series and of a minor series of the complementing b-ions, from which the sequence (231.2) DYEEFXEXAK sequence tag was deduced, are labelled. This sequence is present as ETDYEEFLEIAK in the N-terminal sequence of the 58 kDa parent protein identified as an L-amino acid oxidase (Table 1).



Fig. 6–Proteomics and transcriptomics of Lachesis venoms. Comparison of the protein composition of the venoms of Lachesis *muta* (A) and Lachesis stenophrys (C) determined using a proteomic (this work) and a transcriptomic (panel B) approach [23]. BPP, bradykinin-potentiating/C-natriuretic peptide; NGF, nerve growth factor; LAO, L-amino acid oxidase; PLA₂, phospholipase A₂; SVMP, snake venom metalloproteinase; svVEGF, snake venom vascular endothelial growth factor; CRISP, cysteine-rich secretory protein; Gal-lectin, galactose-specific lectin; Ser-Prot, serine proteinase; 3FTx, three-finger toxin; DPP, dipeptidylpeptidase.

retention time and molecular masses and by sharing tryptic ions include the nerve growth factor isolated in fractions Lm15 and Ls10; the serine proteinase Q27J47 isolated in fractions Lm20 and Ls14/15, as well as the thrombin-like enzime venombin A [S35689] and the plasminogen-activating proteinase P84036, both eluted in HPLC fractions Lm26 and Ls21; the galactose-specific lectin Q9PSM4; the L-amino acid oxidase characterized in fraction Lm29 and Ls24; and the major PI-metalloproteinase, hemorrhagic factor LHFII [P22796].

Based on phylogenetic hypothesis, published morphological and behavioral differences, and the allopatric distributions of distinctive population groups, Zamudio and Green elevated *L. muta* and *L. stenophrys* to species level in 1997 [38]. They estimated that the Central and South American forms diverged 18–6 Mya, perhaps due to the uplifting of the Andes. As judged from the protein chemical and mass spectrometric data listed in Tables 1 and 2, each Lachesis venom may contain 24–26 different gene products. Using a similarity coefficient (PSC), we estimate that *L. muta* and *L. stenophrys* share only 8 proteins. Such a low figure (PSC=30–32%) highlight the rapid structural diversification of venom toxins of closely related congeneric taxa.

Table 3 - Overview of the relative occurrence of proteins of
the different toxin families in the venoms of Lachesis muta
and Lachesis stenophrys

Protein family	% of total venom proteins					
	Lachesis muta	Lachesis stenophrys				
BPP/C-NP	14.7	14.6				
Nerve growth factor	0.6	0.4				
PLA ₂	8.7	12.3				
CRISP	1.8	-				
Serine proteinase	31.2	25.6				
Gal-lectin/C-type lectin-like	7.9	3.6				
L-amino acid oxidase	2.7	5.3				
Zn ²⁺ -metalloproteinase	31.9	38.2				

3.2. Proteomic vs. transcriptomic of L. muta venom

Comparison of the protein composition of the venom of L. muta determined using a proteomic (this work) and a transcriptomic approach [23] shows clear differences, both in the relative occurrence of protein families (expressed as percentages of the total HPLC-separated proteins) (Fig. 6) and in the identity of the polypeptides of each protein family. It is worth to notice that only 6 venom proteins matched any of the previously reported 11 partial or full-length venom gland transcripts [23]. On the other hand, 16 L. muta venom components correspond to proteins not reported in the databasedeposited transcriptome of the same species [23]. This set of novel proteins comprise both minor components, i.e. the 17 kDa nerve growth factor (Lm15) and the C-type lectin(s) spread in fractions 28-38, and relatively abundant proteins, such as all the venom-secreted PLA₂ molecules (Lm16-18), the CRISP molecule found in venom fractions Lm18-20, several serine proteinases (Lm20, Lm24, and Lm28), the single 52 kDa L-amino acid oxidase found in fraction Lm29, and a number of snake venom metalloproteinases of the PI and PIII classes of the reprolysin family (Lm32-38) (Table 1). The low degree of venom composition accordance between the proteomic and the transcriptomic approaches has been also reported for B. gabonica [13], and clearly indicate that the cDNA library lacked many transcripts encoding venom-expressed proteins. On the other hand, in some cases the lack of correspondence between the proteomic and the transcriptomic data may be due to the unavailability of the cDNA-deduced protein sequences in the public-accessible databases. Hence, Junqueirade-Azevedo et al. [23] reported the cloning of two almost identical cDNA clusters coding for an L-amino acid oxidases (comprising 3.7% of the total toxin-coding ESTs), which matched LAOs from other Viperidae species over the entire 2705 bp extension and which may correspond to the L. muta LAO identified earlier by Sanchez and Magalhães [39]. Similarly, Junqueira-de-Azevedo et al. [23] also found clusters coding for single forms of nerve growth factor (NGF; 0.3% of total toxin-coding ESTs) and snake venom vascular

60 10 20 30 40 50 MFVSRLAASG LLLLALLAVS LDGKPVQQWS HKGWPPRPQI PPLVVQQWSQ KPWPPGHHIP |---- Ls7 |-Lm8 * 100 70 80 90 110 120 PVVVQEWPPG HHIPPLVVQQ WSQKKWPPGH HIPPLVVQKW DPPPISPPLL KPHESPAGGT ---Lm9,Ls6------Lm10,Ls8----Lm6,Ls5 --Lm8----Lm11 130 140 150 160 170 180 TALREELSLG PEAALDTPPA GPDVGPRGSK APAAPHRLPK SKGASATSAA SRPMRDLRTD |---Lm3,Ls2---| 190 200 210 220 230 239 GKQARQNWGR MMNPDHHAVG GGGGGGGARR LKGLAKKRVG DGCFGLKLDR IGSMSGLGC |-----Lm1-----| -----Lm13.Ls9-----

Fig. 7 – Bradykinin-potentiating peptides and C-natriuretic peptides. Mapping of the bradykinin-potentiating peptides and the C-natriuretic peptide found in the venoms of L. muta (Lm-) and L. stenophrys (Ls-) onto the cDNA-deduced amino acid sequence of the precursor protein Q27 J49. Peptides shared by both venoms are in boldface and underlined, while those distinctly expressed in L. muta and L. stenophrys are displayed in italics and in boldface on a gray background, respectively. Venom fractions in which the peptides were recovered are identified by their numbering in Tables 1 and 2. Asterisks indicated that the peptide isolated from the venom contained N-terminal pyroglutamic acid.

endothelial growth factor (svVEGF) in their reported *L. muta* transcriptome, although their sequences are not available in the non-redundant SwissProt/TrEML and NCBI databanks. However, in those venoms in which LAO and growth factors have been identified, these toxins appear to be expressed as single components, suggesting that they may represent products of single copy genes, and thus that the proteomic and the transcriptomic approach may match the same venom components.

Transcripts encoding putative secreted toxin classes [23], which were not be found in our proteomic analysis include three finger-like toxins [Q27J50], ohanin-like protein [Q27J48], svVEGF, and dipeptidyl peptidase. In addition, neither L. muta nor L. stenophrys venom contained detectable amounts of lachesin, a medium-size disintegrin [P31990] isolated from lyophilized venom of L. muta of non-reported origin purchased from Miami Serpentarium Laboratories (Salt Lake City, UT) [40]. The occurrence of non-venom-secreted toxins suggests that these messengers could exhibit an individual or a temporal expression pattern over the life time of the snake. Sex-based individual variation of snake venom proteome among B. jararaca siblings have been reported [41]. In addition, ontogenetic and geographical variations have been noticed in the venom proteomes of other snakes, i.e. Crotalus viridis viridis [42], C.v. oreganus [43,44], Bothrox atrox [45,46], and Bothrops asper [46], and might represent a common phenomenon in many other species (see below). Alternatively, the nonvenom-secreted or very low abundance (<0.05% of the total venom proteins), toxins may play a hitherto unrecognized physiological function in the venom gland, or may simply represent a hidden repertoire of orphan molecules which may eventually become functional for the adaptation of snakes to changing ecological niches and prey habits. Clearly, although further work is needed to clarify this point, overall, our results emphasize the relevance of detailed proteomic studies for a thorough characterization of the venom composition.

3.3. Intraspecific variation in venom-secreted PLA₂ molecules

The PLA₂ molecules isolated in fractions 16–18 (Fig. 1, Table 1) display high N-terminal sequence similarity to the hemolytic and platelet aggregation inhibitory Lm-PLA₂-I and Lm-PLA₂-II characterized by Fully et al. [47] from L. muta venom provided by Sigma or Fundação Ezequiel Dias (FUNED, Brazil), to LMPA1 (P84651) from the same source, and to two basic neurotoxic Asp49 PLA₂ molecules (LmTX-I and LmTX-II) isolated from L. muta venom purchased from Sigma [48,49] (Table 4). In particular, the N-terminal sequence of the PLA₂ isolated in fraction Lm18 (Table 1) seems to be identical to that of LM-PLA2-II, except for the striking lack of the serine residue at position 16. This residue is absolutely conserved in the structures of all known myotoxic PLA₂ molecules [50], strongly suggested that a gap at position 16 may represent a sequencing or a typographical error. Regardless of that, none of the other PLA₂ isoenzymes reported in the literature could be matched to any of the PLA2 molecules found in the venom of L. muta from the nature reserve Potrerillos del Guendá (Santa Cruz de la Sierra, Bolivia) sampled here (Table 4). This prompted us to investigate if the lack of identity could be due to geographic variations of *L. muta* venoms. To this end, the venoms of 5 specimens (3 from Bolivia and 2 from Peru) and the venom purchased from Sigma (unknown origin) were compared. Noteworthy, the 6 reverse-phase HPLC separations were essentially superimposable, except for quantitative and qualitative differences in PLA₂ expression (Fig. 8, Table 4). Thus, each venom exhibited a distinct combination, and/or concentration, of the same three PLA₂ molecules (Lm16, Lm17, and Lm18) listed in Table 1. The most abundant PLA₂ molecules in Bolivian specimens were Lm16 and Lm18, whereas Lm17 was the predominant $\ensuremath{\text{PLA}}_2$ in Peruvian L. muta venoms (Fig. 8). The venom purchased from Sigma (of non-declared origin) displayed the "Bolivian PLA₂



Bolivia-1 corresponds to the venom analyzed in detailed in Table 1. Numbers in parentheses indicate the reverse-phase HPLC fraction of Figs. 1 and 7 containing the corresponding PLA₂ molecules. N-terminal sequences and, when available, the molecular mass of PLA₂ proteins characterized previously from venoms of *L. muta* specimens kept in captivity at the Fundação Ezequiel Dias (Belo Horizonte, Brazil) (PA2_LACMU, LM-PLA2-I, and LM-PLA2-II) or isolated from *L. muta* venom purchased from Sigma (LmTX-I and LmTX-II), are displayed.

signature" (Lm16+Lm18) but departed from the Bolivian and Peruvian venom-secreted PLA₂ profiles in expressing a unique PLA₂ molecule (labelled 16a in Fig. 8) with identical N-terminal sequence and very similar isotope-averaged molecular mass (13723 Da) as myotoxic PLA₂ molecules from *B. jararacussu* (AAO27453; 13714 Da) [18], *B. pirajai* (1QLL_A; 13744 Da), and *B. asper* (P24605; 13,725 Da). As a whole, these results point out to a high degree of intraspecific variability in the expression of phospholipases A₂ in *L. muta* venoms. Intraspecific variability of PLA₂ loci has been reported in other viperid (Vipera palestinae [51]) and crotalid (*B. asper* [52]; *Trimeresurus flavoviridis* [53]) species, and this phenomenon is often linked to differences in diet among populations [53,54].

Snake venom phospholipase A₂ genes are members of a large, rapidly-evolving multigene family with many diverse functions. Positive Darwinian selection is common in group-II viperid snake venom PLA₂ genes and is associated with the evolution of new toxin functions and speciation events [55]. Adaptive evolution of group-I phospholipases in elapids is also associated with speciation events [56], suggesting adaptation of the phospholipase arsenal to novel prey species after niche shifts.

3.4. Concluding remarks

Reports of envenomations by species of Lachesis are scarce, probably due to the fact that these species are distributed in remote tropical rainforests. The recorded cases [27,57-60] are characterized by local and systemic manifestations typical of viperid venoms. Local effects include pain, edema, and hemorrhage, with the development of blisters. Systemic manifestations include coagulopathy and widespread bleeding. In some cases, a 'parasympathomimetic-like' effect has been described, with bradicardia, abdominal colics, nausea and vomiting [60]. These latter effects have been described in South American cases, but not in those occurred in Central America. Our detailed proteomic analysis of the venoms of Lachesis species supports the hypothesis that snake venom proteomes are composed of proteins belonging to only a few toxin families exhibiting structural divergence and distinct relative abundances in even closely related species. However, there does not appear to be a simple relationship between venom composition and the presence or absence of the described 'parasympathomimetic-like' effect. Because species-specific effects of venom components are largely unknown, it is difficult to assign a functional role unequivocally to the variation we observed in Lachesis venoms. Subtle functional differences in some venoms components, which may confer distinct pharmacological activities to proteins from the same family but distinctly expressed in L. muta and L. stenophrys, may be responsible for speciesspecific effects. Our proteomic analysis may serve as a starting point for studying structure-function correlations of individual toxins aiming at the development of new research tools and drugs of potential clinical use [61-63]. It is also worth to notice that though intraspecific variation in venom toxins may inform us about evolutionary processes acting at the species or population level, it represents also a source of



Fig. 8–Intraspecific variation in PLA2 isoenzyme venom composition. Panels A–F, details of the reverse-phase HPLC chromatograms of the venoms from *Lachesis muta* from Bolivia (A–C), Peru (D and E), and purchased from Sigma (F). Numbers refer to the reverse-phase HPLC separation shown in Fig. 1. Table 3 displays the N-terminal sequences and molecular masses of the PLA₂ molecules from the different sources. The peak 16a marked with asterisk in F corresponds to a PLA₂ molecule specifically found in the venom sold by Sigma.

Table 4 - Comparison and occurrence of PLA2 molecules characterized in the venom proteomes of Lachesis muta

concern in antivenom production strategies. Broad spectrum antivenoms may thus be prepared using pooled venoms. On the other hand, as knowledge of a particular group increases, its categorisation may need to be re-assessed. At this respect, the occurrence of intraspecific toxin composition variation in separated snake populations, as revealed in this and other works, suggests that *polytypic* species with a number of subspecies or races are more widespread than previously thought. Along with analysis of morphological traits, a detailed proteomic characterization of snake venoms may aid in establishing a taxonomic subdivision of snake species.

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