

Technical University of Denmark



Snake venomics of monocled cobra (Naja kaouthia) and investigation of human IgG response against venom toxins

Laustsen, Andreas Hougaard; Gutiérrez, José María; Lohse, Brian; Rasmussen, Arne R.; Fernández, Julián; Milbo, Christina; Lomonte, Bruno

Published in: Toxicon

Link to article, DOI: 10.1016/j.toxicon.2015.03.001

Publication date: 2015

Document Version Peer reviewed version

Link back to DTU Orbit

Citation (APA):

Laustsen, A. H., Gutiérrez, J. M., Lohse, B., Rasmussen, A. R., Fernández, J., Milbo, C., & Lomonte, B. (2015). Snake venomics of monocled cobra (Naja kaouthia) and investigation of human IgG response against venom toxins. Toxicon, 99, 23-35. DOI: 10.1016/j.toxicon.2015.03.001

DTU Library Technical Information Center of Denmark

General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

1											
2	Snake venomics of monocled cobra (Naja kaouthia) and										
3	investigation of human IgG response against venom toxins										
4											
5	Andreas H. Laustsen ¹ , José María Gutiérrez ² , Brian Lohse ¹ , Arne R. Rasmussen ³ ,										
6	Julián Fernández ² , Christina Milbo ^{1,4} , Bruno Lomonte ²										
7											
8	¹ Department of Drug Design and Pharmacology, Faculty of Health and Medical Sciences,										
9	University of Copenhagen, Denmark										
10	² Instituto Clodomiro Picado, Facultad de Microbiología, Universidad de Costa Rica,										
11	San José, Costa Rica										
12	³ Royal Danish Academy of Fine Arts, School of Conservation, Denmark										
13	⁴ Department of Systems Biology, Technical University of Denmark, Denmark										
14											
15	Running title: Proteomics of monocled cobra venom										
16											
17	Keywords: Naja kaouthia; monocled cobra; snake venom: proteomics; toxicity; human										
18	IgG response; immunity										
19											
20 21 22 23 24 25 26 27 28	Address for correspondence:Dr Bruno LomonteProfessorInstituto Clodomiro PicadoFacultad de MicrobiologíaUniversidad de Costa RicaSan José, COSTA RICAbruno.lomonte@ucr.ac.cr										

29 Highlights

30		
31	•	The venom proteome of the monocled cobra, Naja kaouthia, is presented
32	•	Most abundant venom components are three-finger toxins (neurotoxins and
33		cytotoxins) and phospholipases A ₂
34	•	Toxicity (LD_{50}) screening identified three-finger toxins as the most medically
35		relevant of this venom
36	•	IgG response from a self-immunized human was significant against several toxins,
37		although weak against those with higher toxicity
38		

39 Abstract

The venom proteome of the monocled cobra, *Naja kaouthia*, from Thailand, was 40 characterized by RP-HPLC, SDS-PAGE, and MALDI-TOF-TOF analyses, yielding 38 41 different proteins that were either identified or assigned to families. Estimation of relative 42 protein abundances revealed that venom is dominated by three-finger toxins (77.5%; 43 including 24.3% cytotoxins and 53.2% neurotoxins) and phospholipases A₂ (13.5%). It also 44 contains lower proportions of components belonging to nerve growth factor, 45 ohanin/vespryn, cysteine-rich secretory protein, C-type lectin/lectin-like, nucleotidase, 46 47 phosphodiesterase, metalloproteinase, L-amino acid oxidase, cobra venom factor, and cytidyltransferase protein families. Small amounts of three nucleosides were also 48 evidenced: adenosine, guanosine, and inosine. The most relevant lethal components, 49 categorized by means of a 'toxicity score', were α -neurotoxins, followed by 50 cytotoxins/cardiotoxins. IgGs isolated from a person who had repeatedly self-immunized 51 with a variety of snake venoms were immunoprofiled by ELISA against all venom 52 fractions. Stronger responses against larger toxins, but lower against the most critical α -53 neurotoxins were obtained. As expected, no neutralization potential against N. kaouthia 54 55 venom was therefore detected. Combined, our results display a high level of venom complexity, unveil the most relevant toxins to be neutralized, and provide prospects of 56 discovering human IgGs with toxin neutralizing abilities through use of phage display 57 screening. 58

59 (199 words)

61 **1. Introduction**

Snakebite is a serious medical condition affecting a large number of people 62 63 worldwide; especially in impoverished rural areas of Asia, Africa and Latin America (Gutiérrez et al., 2006; Warrell, 2010a). Asia is the continent where the majority of these 64 bites take place, and also where most deaths occur (Alirol et al., 2010; Chippaux, 1998; 65 Kasturiratne et al., 2008). Currently there are 29 recognized extant species of terrestrial 66 cobras assigned to the genus Naja (Utez and Hošek, 2015). Of these, 11 species are found 67 in Asia, and 18 inhabit Africa (Utez and Hošek, 2015; Wallach et al., 2009). Among the 68 69 cobras, Naja kaouthia (monocled cobra) is widespread in southern Asia and responsible for a significant part of the bites recorded (Viravan et al., 1986; Warrell, 1995; Kulkaew et al., 70 2009). A study in Bangladesh discovered records of 764 snakebites during 1988-1989, of 71 which 34% of the 168 deaths (22% case fatality) were attributed to the cobras Naja naja 72 and *N. kaouthia* (Warrell, 2010b). *N. kaouthia* occurs from north-eastern India, Bangladesh 73 and Bhutan, across southern China, southward to northern Peninsular Malaysia (Prakash et 74 al., 2012; Wüster, 1996) and is common in most of its range (Stuart and Wogan, 2012). N. 75 kaouthia is adapted to a broad range of habitats, including agricultural land, human 76 77 settlements, and bigger cities. However, its natural habitat includes paddy fields, swamps, mangroves, grasslands, scrublands and forest (Stuart and Wogan, 2012). The diet of N. 78 kaouthia covers a broad range of animals from frogs, snakes, small birds to mammals (Cox 79 et al., 1998), and even fish (Kyi and Zug, 2003). N. kaouthia is assessed as Least Concern 80 at IUCN Red List of Threatened Species because of its large distribution, tolerance of a 81 broad range of modified habitats, and is reported to be in abundance (Stuart and Wogan, 82 83 2012). In Thailand it has been reported that many cases of hospitalization due to snakebite

were caused by *N. kaouthia* (Kulkeaw et al., 2009). Human envenomings by *N. kaothia* are
predominantly characterized by neuromuscular paralysis which in severe cases ends up in
respiratory paralysis, and by local tissue damage, i.e. swelling, necrosis and blistering
(Warrell, 1995; Wongtongkam et al., 2005).

Enzymatic activities and toxicity of the venom of N. kaouthia have previously been 88 studied (Pakmanee et al., 1998; Mukherjee and Maity, 2001; Das et al., 2013). Furthermore, 89 90 a qualitative proteome of the venom has been reported showing the presence of 61 proteins 91 belonging to 12 protein families (Kulkaew et al., 2007). The most prominent component is the long α -neurotoxin α -cobratoxin (Richard et al., 2013), constituting about 25% of the 92 93 venom (Kulkaew et al., 2009) and belonging to the three-finger toxin family, of which N. 94 kaouthia venom has several other members (Kulkaew et al., 2007). Other proteins that have 95 been studied include phospholipases A₂ (PLA₂s) (Joubert and Taljaard, 1980a; Reali et al., 2003; Doley and Mukherjee, 2003), L-amino acid oxidases (Tan and Swaminathan, 1992; 96 97 Sakurai et al., 2001), cardiotoxins (Joubert and Taljaard, 1980b; Fletcher et al., 1991; 98 Debnath et al., 2010; Jamunaa et al., 2012), and nerve growth factors (Kukhtina et al., 2001). However, a proteomic analysis of this venom that integrates an estimation of relative 99 protein abundances together with a detailed screening of the toxicity of its various 100 101 components, is pending. An integrated functional and proteomic characterization of N. 102 kaouthia venom is relevant not only for a deeper understanding of the composition of the 103 venom and its relationships to toxicity, but also to establish a basic platform for 104 antivenomic analyses.

Developing more potent antivenoms with better safety profiles is of medical relevance. Besides the traditional animal-derived antivenoms generated by immunization of horses with pools of snake venoms (Gutiérrez et al., 2011), various alternative approaches 108 have been pursued at the experimental level in order to generate more specific antibody-109 based therapies aimed at neutralizing the most important toxins in the venom. One approach has been to raise murine antibodies specific towards relevant N. kaouthia toxins 110 (Charpentier et al., 1990; Masathien et al., 1994). Another approach for identifying potent 111 antitoxins is phage display screening (Roncolato et al., 2015). By using this methodology, 112 inhibitors against N. kaouthia toxins have been identified for both PLA₂s (Chavanayarn et 113 114 al., 2012) and a-cobratoxin (Stewart et al., 2007; Kulkaew et al., 2009; Richard et al., 115 2013). These studies discovered inhibitors through screening of phage display libraries 116 based on heavy chain fragments obtained from llama. Although promising, it could, 117 however, be argued that llama-derived inhibitors, despite being much smaller than whole IgG antibodies and therefore likely to be less immunogenic, are still non-human. An 118 alternative to overcome this problem is based on modern antibody humanization procedures 119 120 (Safdari et al., 2013). In another study, inhibitors based on human single chain Fv (scFv) fragments from non-immunized donors were developed (Kulkaew et al., 2009). These scFv 121 fragments were not effective in rescuing mice injected with venom, which is likely to be 122 123 due to naïve origin of the gene library for these fragments, since none of the human donors 124 had been exposed to snake venom. Thus, the search for novel approaches to generate 125 human antibodies against cobra venom toxins remains a challenge.

With this study, we report for the first time a venomics-based quantitative estimation of the proteome and a full protein lethality profile for *N. kaouthia* venom, in order to lay the foundation for developing a recombinant antivenom by identifying the most relevant toxins present in this venom. Furthermore, we uncover the presence of human IgG antibodies specific to *N. kaouthia* venom in the serum obtained from a unique individual exposed to low doses of a wide variety of snake venoms over a period of more than 25 years. The binding ability of his antibodies, as well as of those from a person bitten twice
by the Southern Burrowing Asp, *Atractaspis bibronii*, was tested against the different
venom fractions from *N. kaouthia*, in comparison to a healthy volunteer unexposed to snake
venoms. Finally, the neutralizing ability against *N. kaouthia* venom of the purified human
IgG antibodies was evaluated in a mouse model.

137

138 2. Materials and Methods

139 2.1 Snake venom

Naja kaouthia venom was obtained from Latoxan SAS, Valence, France. The
venom is a pool collected from several specimens, originally from Thailand.

142

143 2.2 Venom separation by reverse-phase HPLC and SDS-PAGE

144 Following the 'snake venomics' analytical strategy (Calvete, 2011), crude venom was fractionated by a combination of RP-HPLC and SDS-PAGE separation steps. Venom 145 (2 mg) was dissolved in 200 µL of water containing 0.1% trifluoroacetic acid (TFA; 146 solution A) and separated by RP-HPLC (Agilent 1200) on a C₁₈ column (250 x 4.6 mm, 5 147 um particle; Teknokroma). Elution was carried out at 1 mL/min by applying a gradient 148 towards solution B (acetonitrile, containing 0.1% TFA): 0% B for 5 min, 0–15% B over 10 149 min, 15–45% B over 60 min, 45–70% B over 10 min, and 70% B over 9 min, as previously 150 151 described (Lomonte et al., 2014a). Manually collected fractions were dried in a vacuum centrifuge, redissolved in water, reduced with 5% β-mercaptoethanol at 100 °C for 5 min, 152 and further separated by SDS-PAGE in 12% gels (Bio-Rad). Proteins were stained with 153 colloidal Coomassie blue G-250, and gel images were acquired on a ChemiDoc[®] recorder 154 using ImageLab[®] software (Bio-Rad). 155

157 2.3 Protein identification by tandem mass spectrometry of tryptic peptides

Protein bands were excised from the polyacrylamide gels and subjected to reduction 158 (10 mM dithiothreitol), alkylation (50 mM iodoacetamide), and overnight in-gel digestion 159 with sequencing grade trypsin (Sigma), in 50 mM ammonium bicarbonate at 37°C. The 160 resulting tryptic peptides were extracted with 50% acetonitrile containing 1% TFA, and 161 162 analyzed by MALDI-TOF-TOF on an AB4800-Plus Proteomics Analyzer (Applied Biosystems). Peptides were mixed with an equal volume of saturated α -cyano-163 hydroxycinnamic acid (in 50% acetonitrile, 0.1% TFA), and spotted (1 µL) onto an Opti-164 TOF 384-well plate, dried, and analyzed in positive reflector mode. TOF spectra were 165 acquired using 1500 shots and a laser intensity of 3000. The ten most intense precursor ions 166 were automatically selected and their TOF/TOF fragmentation spectra were acquired using 167 500 shots at a laser intensity of 3900. External calibration in each run was performed with 168 CalMix[®] standards (ABSciex) spotted onto the same plate. For protein identification, 169 resulting spectra were searched against the UniProt/SwissProt database using ProteinPilot® 170 v.4 and the Paragon[®] algorithm (ABSciex) at $\geq 95\%$ confidence, or manually interpreted. 171 Few peptide sequences with lower confidence scores were manually searched using 172 BLAST (http://blast.ncbi.nlm.nih.gov) for protein similarity and protein family assignment. 173

174 RP-HPLC fractions corresponding to small molecules, eluting in the initial peaks of 175 the chromatogram, were analyzed by ESI-MS/MS on a Q-Trap[®] 3200 instrument (Applied 176 Biosystems). Samples (10 μ L) were loaded into metal-coated capillary tips (Proxeon) and 177 directly infused into a nano-ESI source (Protana) operated at 1300 V. Spectra were 178 acquired in positive Enhanced Resolution mode. nESI-MS performed in this instrument 179 was also used to determine the isotope-averaged mass of intact proteins in selected peaks from the RP-HPLC separation. For this purpose, mass spectra were acquired in Enhanced
Multicharge mode in the m/z range 700-1700, and deconvoluted with the aid of the
Analyst[®] v.1.5 software (ABSciex).

- 183
- 184 2.4 Relative protein abundance estimations

The relative abundances of the venom proteins identified were estimated by integrating the areas of their chromatographic peaks at 215 nm, using the ChemStation[®] software (Agilent), which correlates with peptide bond abundance (Calvete, 2011). For HPLC peaks containing several electrophoretic bands, their percentage distributions were assigned by densitometry, using ImageLab[®] (Bio-Rad).

190

191 2.5 Nucleoside and FAD analysis

192 The presence of selected nucleosides (adenosine, inosine, guanosine), and flavine adenine dinucleotide (FAD) was determined by spiking a sample of 1 mg of venom with 10 193 µg of each nucleoside or FAD, respectively, and separating it by reverse-phase HPLC as 194 195 described in section 2.2. If the nucleoside or FAD peak coincided with a peak already 196 present in a crude venom sample (as judged by the increment in the height of the peak), and if this venom peak showed an ESI-MS spectrum similar to that of the nucleoside or FAD, 197 198 the identity of venom component was judged to be the same as the nucleoside or FAD. Further confirmation of the molecular identities of the nucleosides was obtained by 199 200 acquiring their collision-induced dissociation MS/MS spectra in positive mode, using the 201 Enhanced Product Ion tool of the Analyst v1.5 in the QTrap 3200 mass spectrometer as 202 described (Laustsen et al., 2015). The abundance for nucleosides given in Table 1 is only indicative of an approximate value, due to differences in absorbance at 215 nm between 203

204 nucleosides and proteins.

205

206 2.6 In vitro enzymatic activities

207 2.6.1. Phospholipase A_2 activity

PLA₂ activity was assayed on the monodisperse synthetic chromogenic substrate 4nitro-3-octanoyloxybenzoic acid (NOBA) (Holzer and Mackessy, 1996). Twenty-five μ L containing various amounts of venom were mixed with 200 μ L of 10 mM Tris, 10 mM CaCl₂, 0.1 M NaCl, pH 8.0, and 25 μ L of NOBA to achieve a final substrate concentration of 0.32 mM. Plates were incubated at 37 °C for 60 min, and absorbances were recorded at 405 nm in a microplate reader. Samples were assayed in triplicates.

214

215 2.6.3 Proteinase activity

Variable amounts of venom (10 to 40 μ g) were added to 100 μ L of azocasein (10 mg/mL in 50 mM Tris–HCl, 0.15 M NaCl, 5 mM CaCl₂ buffer, pH 8.0), and incubated for 90 min at 37 °C. The reaction was stopped by addition of 200 μ L of 5% trichloroacetic acid, and after centrifugation (5 min, 13,000 rpm), 150 μ L of supernatants were mixed with 100 μ L of 0.5 M NaOH, and absorbances were recorded at 450 nm. The absorbance of azocasein incubated with distilled water alone was used as a blank, being subtracted from all readings (Wang et al. 2004). Samples were assayed in triplicates.

223

224 2.7 Toxicological profiling

225 2.7.1 Animals

In vivo assays were conducted in CD-1 mice, supplied by Instituto Clodomiro Picado, following protocols approved by the Institutional Committee for the Use and Care of Animals (CICUA), University of Costa Rica. Mice were housed in cages for groups of
4–8, and were provided food and water *ad libitum*.

- 230
- 231 2.7.2 Toxicity of crude venom and isolated toxins

The lethality of the whole venom and fractions or isolated toxins was tested by intravenous (i.v.) injection in groups of four mice (18–20 g body weight). Various amounts of venom or fractions/toxins were dissolved in phosphate-buffered saline (PBS; 0.12 M NaCl, 0.04 M sodium phosphate buffer, pH 7.2), and injected in the caudal vein, using an injection volume of 100 μ L. Then, deaths occurring within 24 hr were recorded. Median lethal dose (LD₅₀) was calculated by probits (Finney, 1971), using the BioStat[®] software (AnalySoft).

The toxicity of venom fractions was initially screened by selecting a dose based on 239 240 fraction abundance. The dose was selected to assess whether the fraction would score above or below 5 according to the Toxicity Score defined by Laustsen et al. (2015), which 241 represents the ratio of protein fraction abundance (%) in the venom divided by its estimated 242 LD₅₀ in CD-1 mice by i.v. injection. Fractions that were not lethal at this dose (yielding a 243 Toxicity Score < 5) were considered as having insignificant toxicity. All fractions that were 244 245 lethal at this level, and some fractions killing mice at a Toxicity Score level between 1-5 246 were further evaluated, and precise LD₅₀ values were further determined for them. Groups of 3-5 mice per dose were used in order to minimize the use of experimental animals. 247

248

249 2.8 Preparation of human IgGs and human serum

250 2.8.1 Donor 1

251	This donor has over a period of 25 year injected himself with sub-lethal amounts of
252	venom from the following snakes: Viridovipera (formerly Trimeresurus) stejnegeri,
253	Trimeresurus nebularis, T. macrops, T. popeorum, T. trigonocephalus, T. albolabris, T.
254	venustus, Bothriechis schlegelii, Crotalus enyo enyo, Naja kaouthia, N. naja, N. siamensis,
255	Agkistrodon contortrix, A. bilineatus, Bothrops asper, Crotalus oreganus, Crotalus
256	oreganus helleri, Cerastes cerastes, and Micrurus fulvius. In the last 5 years, this individual
257	has been injecting himself with venom from Naja kaouthia on an average of every 2 weeks.
258	Brian Lohse has a written consent from the Danish scientific ethical council (H-3-2013-
259	FSP60) to use the blood sample from this donor according to the law of the ethical
260	committee § 2.1. After having provided written and oral information about the project to
261	this donor, written consent was given for obtaining blood samples and for performing the
262	experiments included in the present work. The donor requested the inclusion of his name to
263	be acknowledged in publications.

265 2.8.2 Donor 2

Donor 2 is an entomologist, who was unfortunate to be bitten twice by the snake *Atractaspis bibronii* (Southern Burrowing Asp) during fieldwork in August 2013. After having provided written and oral information about the project to this donor, written consent was given for obtaining blood samples and for performing the experiments here included. The donor requested the inclusion of his name to be acknowledged in publications.

272

273 2.8.3 Blood sampling for obtaining IgGs

274	200 ml of blood was sampled from each donor in Vacuette [®] tubes of 4 mL,
275	containing Z Serum Clot Activator (Greiner bio-one). Every tube was gently tilted up and
276	down ten times as soon as it was filled with blood to ensure thorough mixing with the
277	coagulation factor. The tubes were kept at room temperature until full coagulation (30 min).
278	Then, tubes were centrifuged at 3000 g for 10 min, and the serum was transferred to
279	cryotubes and stored at -20 °C.

2.8.4 Protein A purification of IgG antibodies 281

After filtration of serum through a 0.45 µm membrane, the procedure accompanying 282 283 the Protein A Antibody Purification Kit (PURE1A, Sigma-Aldrich) was followed in order 284 to obtain human IgG antibodies in solution at physiological pH. IgG antibodies were stored at 4 °C. 285

286

287 2.8.5 Preparation of human serum from healthy volunteer

288 Blood was obtained from a healthy donor to serve as a negative control. After clotting, serum was separated by centrifugation, and stored at -20 °C. 289

290

291 2.9 Immunoreactivity of human IgGs against venom fractions by ELISA

Wells in MaxiSorp plates (NUNC, Roskilde, Denmark) were coated overnight with 292 293 0.6 µg of each HPLC venom fraction dissolved in 100 µL PBS. Then, wells were blocked by adding 100 µL PBS containing 2% (w:v) bovine serum albumin (BSA, Sigma) and 294 295 leaving the plates on a mixer at room temperature for 1 h. Plates were then washed five 296 times with PBS. A dilution of each human IgG pool in PBS + 2% BSA was prepared such that the concentration of IgG proteins was 86 µg/mL (as measured by their absorbance at 297

280 nm on a NanoDrop[®] 2000c instrument, Thermo Scientific). Then, 100 μL of these IgG 298 299 solutions were added to the wells, after which the plates were incubated for 2 h, followed by five additional washings with PBS. Then, 100 µL of a 1:2000 dilution of affinity-300 purified goat anti-human IgG (gamma-chain specific) antibodies conjugated to alkaline 301 phosphatase (Cappel, Organon Teknika) in PBS + 2% BSA was added to each well. The 302 plates were incubated for 2 h, and then washed five times with FALC buffer (0.05 M Tris, 303 0.15 M NaCl, 20 µM ZnCl₂, 1 mM MgCl₂, pH 7.4). Color development was achieved by 304 the addition of 100 μ L *p*-nitrophenyl phosphate (1 mg/mL in 9.7% v/v diethanolamine 305 buffer, pH 9.8). The absorbances at 405 nm were recorded (Multiskan FC, Thermo 306 307 Scientific) at several time intervals.

308

309 2.10 Neutralization studies with human IgGs

After purification according to section 2.8.4, the human IgGs were kept refrigerated. Six months later, 3 mL of IgGs were dialyzed against 2 L of PBS overnight and lyophilized. The IgGs were reconstituted in 500 μ L PBS and the protein concentration was measured on NanoDrop to be 14 mg/ml. IgG was mixed with venom at a ratio of 78 mg IgG/mg venom and incubated at 37 °C for 30 min. Aliquots of 120 μ L of this mixture, containing 4 LD₅₀s of venom, were then injected i.v. into a group of four mice (18-20 g) and deaths occurring within 24 hr were recorded.

317

318 **3.0 Results and Discussion**

319 *3.1 Venomics*

A proteomic characterization of *N. kaouthia* venom coupled with an estimation of its protein relative abundances was, with this study, carried out for the first time. Twenty322 eight peaks from the venom were resolved by RP-HPLC. The first three fractions eluting 323 from the column were non-peptidic, while the remaining 25 peaks were further resolved into 50 protein bands by SDS-PAGE separation (Fig.1). In-gel digestion and MALDI-TOF-324 TOF analysis of these yielded a total of 38 identified proteins, while only 4 remained 325 unknown. Altogether, the unidentified venom components represent less than 1% of the 326 327 total venom protein content, while the rest were either identified or assigned to protein 328 families (Table 1). A previous qualitative proteomic profile was reported for this venom, 329 describing matches to 61 orthologous proteins classified into 12 groups (Kulkaew et al., 330 2007). Of these, oxoglutarate dehydrogenase complex was identified on the basis of a 331 single peptide that matched such enzyme from a prokaryote (Burkholderia cenocepacia), 332 while serum albumin likely corresponds to contaminating traces derived from slight trauma that may occur during venom collection (see for example Lomonte et al., 2014b). In 333 334 addition, the previous study considered cardiotoxins and cytotoxins as two different protein groups, in spite of these two names being used to refer to the same type of toxins, 335 structurally classified within the three-finger toxin family (Kini and Doley, 2010). Our 336 337 results are concordant with the remaining eight protein families detected by Kulkaew et al. 338 (2007), but further expand the proteomic profile of N. kaouthia venom with the addition of members belonging to the nerve growth factor, ohanin-like/vespryn, C-type lectin/lectin-339 340 like, nucleotidase, phosphodiesterase, L-amino acid oxidase, and cytidyltransferase protein families (Table 1 and Fig.2), and moreover, providing a quantitative estimation of their 341 342 relative abundances in the venom.

Peaks 1 to 3 were analyzed by direct infusion using nESI-MS/MS, since they did not show proteins by gel electrophoresis. Peaks 2 and 3 had molecular masses of 268 Da and 283 Da, respectively, and the nucleoside analysis by HPLC (Fig.3) revealed their identities as adenosine and guanosine, respectively. Also, a trace of inosine between peaks
2 and 3 was detected by the nucleoside analysis by HPLC (Fig.3).

All peptidic peaks were tested in vivo for acute toxicity, except for peaks 23 and 26, 348 which yielded very low amounts of proteins (Table 2), and peaks 24, 27, and 28, which 349 correspond to metalloproteinases, enzymes known to be inactivated by the RP-HPLC 350 separation procedure. In Fig.2, the overall protein composition of the venom of N. kaouthia, 351 352 expressed as percentage of total protein and nucleoside content, is represented. Two main 353 families predominate in the venom: Three-finger toxins (3FTx; 77.5%, of which 24.3% are 354 cytotoxins, the rest being neurotoxins) and PLA₂s (PLA₂; 13.5%). In partial agreement with 355 previous findings, the major single component of the three-finger toxins was determined to 356 be alpha-elapitoxin-Nk2a (α -cobratoxin), a long-chain neurotoxin. However, in our results, 357 this toxin represented 32.3% of the whole venom, which is significantly higher than the reported values of 23-25% (Kulkaew et al., 2009). This increase might reflect either 358 359 geographical variations in venom composition or differences in methodology. As other α neurotoxins, alpha-elapitoxin-Nk2a is able to bind to nicotinic receptors at the motor end-360 plate of muscle fibers, thus generating a flaccid paralysis leading to respiratory failure and 361 death (Ultsintong et al., 2009; Alkondon et al., 1990). This type of neurotoxin is 362 responsible for the life-threatening neuromuscular paralysis in human victims of neurotoxic 363 cobra bites (Warrell, 1995). 364

An interesting finding, discovered by the lack of high acute toxicity in the rodent model (Table 2), and reinforced by the difference in human IgG binding in the ELISA assays (Fig.6) was that the protein identified in peak 10 as alpha-elapitoxin-Nk2a appears to correspond to a closely related homolog. In order to validate this, both peak 10 and peak 6 369 (known to contain alpha-elapitoxin-Nk2a) were analyzed by nESI-MS/MS (Fig.5), 370 according to which their masses were 7619 and 7826 Da, respectively (alpha-elapitoxin-7831 371 Nk2a has reported of Da according а mass to: http://www.uniprot.org/uniprot/P01391). Thus, peak 10 should be a homolog of alpha-372 elapitoxin-Nk2a sharing peptide sequences (Table 1), but different by 207 Da in total mass. 373 374 Since the toxicity of this protein is markedly lower in comparison to alpha-elapitoxin-Nk2a, 375 further studies to determine its complete amino acid sequence and bioactivities could be of 376 interest, as it represents nearly 10% of the venom proteome.

377 In addition to α -neurotoxins, other representatives of the three-finger toxin family 378 detected in N. kaouthia venom proteome are the so-called 'cytotoxins' (Table 1). These 379 toxins are known to interact with and disrupt the integrity of plasma membrane of various 380 cell types, thus inducing irreversible cell damage (Konshina et al., 2012). These cytotoxins are responsible for the local necrotic effect characteristic of envenomings by N. kaouthia 381 (Warrell, 1995). Also, PLA₂ is commonly an abundant component in the majority of elapid 382 383 venom proteomes characterized so far, and it was detected in our proteomic analysis in N. 384 kaouthia venom. PLA2 activity was confirmed for N. kaouthia venom in vitro, although it 385 was significantly lower than for *M. nigrocinctus*, a Central American elapid species 386 (Fig.4A). Elapid PLA₂s have been shown to exert a number of toxicological effects, such as 387 myonecrosis (Bhat and Gowda, 1989; Harris, 1991), anticoagulation (Kini, 2005; Doley 388 and Mukherjee, 2003), inhibition of platelet aggregation (Clemetson et al., 2007), and 389 edema (Wang and Teng, 1990). In the case of N. kaouthia venom, PLA₂s might contribute to the local inflammatory and necrotizing effects observed in human patients. Furthermore, 390 391 cardiotoxins and PLA₂s act synergistically to induce cell membrane damage (Harvey et al., 392 1983), thus potentiating the tissue damaging effect of the venom. Altogether, a high proportion of the components of *N. kaouthia* venom belong to these two toxin families, i.e. three-finger toxins and PLA₂s. This further supports the concept that high diversity within just a few toxin families may be a general theme within elapid venoms (Calvete, 2013; Laustsen et al., 2015). Such idea is supported also by findings of Vonk et al. (2013) on the genome of the elapid *Ophiophagus hannah* (king cobra), in which the three-finger toxins family was found to be highly diversified.

399 In amounts each representing <2% of the total venom content, the venom also 400 contains members of the nerve growth factor (NGF), Ohanin/vespryn (OHA), Cysteine-rich secretory protein (CRISP), C-type lectin/lectin-like (CTL), nucleotidase (NUCL), 401 402 phosphodiesterase (PDE), metalloproteinase (MP), L-amino oxidase (LAO), Cobra venom 403 factor (CVF), and Cytidyltransferase (CTT) families. The low content of proteinases is 404 further supported by the negligible proteinase activity of the venom upon a general 405 substrate such as azocasein (Fig.4B). Also, regarding non-peptidic material, small amounts were identified to be adenosine, inosine, and guanosine, which have also been detected in 406 407 other snake venoms (Aird, 2002; Laustsen et al., 2015).

408 The results of toxicity testing of the fractions are summarized in Table 2. Unsurprisingly, the lethality of the venom stems from the presence of potent α -neurotoxins 409 and cytotoxins, with PLA_{2} s playing a secondary role leading to local tissue damage at the 410 411 bite wound. A Toxicity Score is shown in Table 2, which takes into account both potency 412 and abundance. Toxins with a higher score, as estimated in mice, are expected to be of 413 higher medical relevance. These are abundant and/or very potent, while toxins with a low 414 score are less medically relevant. Based on this score, it is clear that alpha-elapitoxin-Nk2a 415 is by far the toxin that is most clinically relevant to neutralize, with a value of 326. Some toxins have very low LD₅₀s (particularly fractions 4 and 5), but due to their lower 416

417 abundance their Toxicity Scores are much lower. Others have intermediate LD_{50} potencies, 418 but become relevant due to their high abundance (e.g. fraction 16). The cytotoxins present in the venom act by a different mechanism than the α -neurotoxins, providing N. kaouthia 419 venom with a dual strategy for prey subduction. It is interesting to note that the 420 accumulated Toxicity Scores for all toxins amount to the same value as the Toxicity Score 421 422 for the whole venom. This is not the case for the venom from the related elapid, 423 Dendroaspis polylepis, which shows evidence of synergism, and where the accumulated 424 Toxicity Scores is considerably lower than the Toxicity Score of whole venom (indicating 425 synergism between the toxins and not simply additive effect) (Laustsen et al., 2015). One 426 interpretation of this finding could be that N. kaouthia venom does not exert significant 427 synergism, but instead relies on two complementary strategies for acute toxicity.

In terms of therapeutic targets, based on their Toxicity Score, the most relevant 428 429 toxins in N. kaouthia venom that should be neutralized by an effective antivenom are (in order of importance): alpha-elapitoxin-Nk2a (P01391), cobrotoxin-c (P59276) and 430 cobrotoxin-b (P59275), and the homolog of three-finger toxin from Naja atra (E2IU03), 431 432 while the cytotoxins are of less importance in terms of overall lethality, although they are 433 likely to play a relevant role in local necrosis, and hence should be neutralized by antivenoms. In addition, neutralization of PLA₂s should also be considered a priority in 434 435 order to limit local tissue damage from envenomings. Thus, antivenomic studies with N. kaouthia venom should establish whether antivenoms are able to bind these toxins. 436

437

438 3.2 Immunoreactivity of human IgGs against venom fractions

The isolated human IgGs from the unique individual who had self-injected small
amounts of venom from various different snakes, including *N. kaouthia*, showed significant

441 binding ability to isolated venom fractions, clearly above the background established by 442 serum of a healthy donor, or by IgGs from the donor bitten twice by a different snake 443 species (Fig.6). It is evident that the antibody response of the individual exposed to N. kaouthia venom was higher against larger toxins present in later eluting peaks, while the 444 response was limited against the most important α -neurotoxins (peaks 4, 5, and 6), which 445 could be due to the low immunogenicity of small toxins (Fernández et al., 2011). Therefore, 446 447 it was not surprising that IgGs were unable to neutralize the lethality of whole venom when 448 mice were challenged with 4 LD_{50} s mixed with IgGs at a level of 0.013 mg toxin per mg of 449 IgGs. Nevertheless, it could be particularly interesting to further investigate these 450 antibodies via phage display screening. In llamas, it has been shown that using naïve libraries for phage display screening did not yield high affinity inhibitors to snake toxins, 451 while excellent inhibitors of alpha-elapitoxin-Nk2a were discovered this way, when a 452 453 library from an immunized llama was used (Richard et al., 2013). Kulkeaw et al. (2009) were the first to report drug discovery effort using a human phage display library. However, 454 they similarly did not find strong inhibitors of snake toxins, most likely given the fact that 455 456 their library was naïve. Therefore, using the antibody genes from the unique donor to 457 construct a phage display library, and subsequently screening this library, has the potential to yield toxin inhibitors of therapeutic value. Further studies of the donor's immune 458 459 response against toxins from other snake species are warranted and may help direct phage display screening efforts towards the generation of new antibodies of human origin. 460

461

462 **4.0 Concluding remarks and outlook**

463 Differing from previous proteomic studies on *N. kaouthia* venom, the present work 464 combined the cataloguing of venom proteins with an estimation of their relative abundances, together with an assessment of their lethal toxicity for mice. This approach uncovered that the most abundant group of proteins was three-finger toxins, followed by PLA₂s. Based on an evaluation of the accumulation of Toxicity Scores and knowledge about toxin functions, the venom seems to have a dual strategy for prey subduction elicited by α -neurotoxins and cytotoxins, between which pharmacological interaction is speculated only to be of additive value rather than synergism.

ELISA immunoprofiling of human IgGs isolated from a human donor, who had repeatedly injected himself with snake venom from various different species, revealed presence of antibodies with high binding ability to several of the isolated venom fractions, although not against the medically most relevant toxins of *N. kaouthia*. Still, the presence of these antibodies opens the prospect of discovering inhibitors of therapeutic value.

476

477 Acknowledgments

The authors thank Steve Ludwin and Thomas Pape for donation of blood samples 478 for this study. We thank Dr. Steven D. Aird (Okinawa Institute of Science and Technology, 479 480 Japan) for fruitful discussions about nucleosides present in snake venom. We further thank Jens Kringelum (Technical University of Denmark), Mikael Engmark (Technical 481 University of Denmark), and Alexandra Bak Jakobsen (Denmark) for fruitful scientific 482 483 discussions. We thank the Department of Drug Design and Pharmacology, University of Copenhagen, and Instituto Clodomiro Picado, Universidad de Costa Rica, for supporting 484 485 the research. Finally, we are grateful to the following foundations for financial support: Drug Research Academy (University of Copenhagen), Dansk Tennis Fond Oticon Fonden, 486 487 Knud Højgaards Fond, Rudolph Als Fondet, Henry Shaws Legat, Læge Johannes Nicolai Krigsgaard of Hustru Else Krogsgaards Mindelegat for Medicinsk Forskning og 488

Medicinske Studenter ved Københavns Universitet, Lundbeckfonden, Torben of Alice
Frimodts Fond, Frants Allings Legat, Christian og Ottilia Brorsons Rejselegat for Yngre
Videnskabsmænd og -kvinder, and Fonden for Lægevidenskabens Fremme.

492

493 **Conflicts of interest statement**

The authors declare that there are no conflicts of interest related to this study. Sources that provided financial support were not involved in the collection, analysis, or interpretation of data, nor in writing the report and submitting it for publication.

498 **References**

- Aird, S.D., 2002. Ophidian envenomation strategies and the role of purines. Toxicon 40,
 335–393.
- Alirol, E., Sharma, S.K., Bawaskar, H.S., Kuch, U., Chappuis, F., 2010. Snake bite in South
 Asia: a review. PLoS Negl. Trop. Dis. 4, e603.
- Alkondon, M., Albuquerque, E.X., 1990. Alpha cobratoxin blocks the nicotinic
 acetylcholine receptor in rat hippocampal neurons. Eur. J. Pharmacol. 191, 505–
 506.
- Bhat, M.K., Gowda, T.V., 1989. Purification and characterization of a myotoxic
 phospholipase A₂ from Indian cobra (*Naja naja naja*) venom. Toxicon 27, 861–873.
- Calvete, J.J., 2011. Proteomic tools against the neglected pathology of snake bite
 envenoming. Expert Rev. Proteomics 8, 739–758.
- Calvete, J.J., 2013. Snake venomics: from inventory of toxins to biology. Toxicon 75, 44–
 72.
- Charpentier, I., Pillet, L., Karlsson, E., Couderc, J., Ménez, A., 1990. Recognition of the
 acetylcholine receptor binding site of a long-chain neurotoxin by toxin-specific
 monoclonal antibodies. J. Mol. Recognition 3, 74–81.
- Chavanayarn, C., Thanongsaksrikul, J., Thueng-In, K., Bangphoomi, K., Sookrung, N.,
 Chaicumpa, W., 2012. Humanized single-domain antibodies (VH/VHH) that bound
 specifically to *Naja kaouthia* phospholipase A₂ and neutralized the enzymatic
 activity. Toxins 4, 554–567.
- Chippaux, J.P., 1998. Snake-bites: appraisal of the global situation. Bull. World Health
 Organ. 76, 515–524.

- 521 Chuman, Y., Nobuhisa, I., Ogawa, T., Deshimaru, M., Chijiwa, T., Tan, N.H., Fukumaki,
 522 Y., Shimohigashi, Y., Ducancel, F., Boulain, J.C, Ménez, A., Ohno, M., 2000.
- 523

- Regional and accelerated molecular evolution in group I snake venom gland phospholipase A_2 isozymes. Toxicon 38, 449–462.
- Clemetson, K.J., Lu, Q., Clemetson, J.M., 2007. Snake venom proteins affecting platelets
 and their applications to anti-thrombotic research. Curr. Pharm. Des. 13, 2887–
 2892.
- Cox, M., Van Dijk, P.P., Nabhitabhata, J., Thirahupt, K., 1998. A photographic guide to
 snakes and other reptiles of Peninsular Malaysia, Singapore and Thailand. Ralph
 Curtis Books, Sanibel Island, Florida, 144 pp.
- Das, D., Urs, N., Hiremath, V., Vishwanath, B.S., Doley, R., 2013. Biochemical and
 biological characterization of *Naja kaouthia* venom from North-East India and its
 neutralization by polyvalent antivenom. J. Venom Res. 4, 31–38.
- Debnath, A., Saha, A., Gomes, A., Biswas, S., Chakrabarti, P., Giri, B., Biswas, A.K., Das
 Gupta, S., Gomes, A., 2010. A lethal cardiotoxic-cytotoxic protein from the Indian
 monocellate cobra (*Naja kaouthia*) venom. Toxicon 56, 569–579.
- Doley, R., Mukherjee, A.K., 2003. Purification and characterization of an anticoagulant
 phospholipase A₂ from Indian monocled cobra (*Naja kaouthia*) venom. Toxicon 41,
 81–91.
- Fernández, J., Alape-Girón, A., Angulo, Y., Sanz, L., Gutiérrez, J.M., Calvete, J.J.,
 Lomonte, B., 2011. Venomic and antivenomic analyses of the Central American coral
 snake, *Micrurus nigrocinctus* (Elapidae). J. Proteome Res. 10, 1816–1827.
- Finney, D.J., 1971. Statistical Methods in Biological Assay. London: Charles Griffin and
 Company Ltd.

- Fletcher, J.E., Jiang, M.S., Gong, Q.H., Yudkowsky, M.L., Wieland, S.J., 1991. Effects of a
 cardiotoxin from *Naja naja kaouthia* venom on skeletal-muscle Involvement of
 calcium-induced calcium release, sodium-ion currents and phospholipase A₂ and
 phospholipase C. Toxicon 29, 1489–1500.
- Gutiérrez, J.M., Theakston, R.D.G., Warrell, D.A., 2006. Confronting the neglected
 problem of snake bite envenoming: The need for a global partnership. PLoS Med 3,
 e150.
- Gutiérrez, J.M., León, G., Lomonte, B., Angulo, Y., 2011. Antivenoms for snakebite
 envenomings. Inflamm. Allergy Drug Targets 10, 369-380.
- Harris, J.B., 1991. Phospholipases in snake venoms and their effects on nerve and muscle. *In:* Snake Toxins (Harvey, A.L., Ed.), pp.91-129. New York, Pergamon Press.
- Harvey, A.L., Hider, R.C., Khader, F., 1983. Effect of phospholipase A on actions of cobra
 venom cardiotoxins on erythrocytes and skeletal muscle. Biochim. Biophys. Acta
 728, 215–221.
- Holzer, M., Mackessy, S.P., 1996. An aqueous endpoint assay of snake venom
 phospholipase A₂. Toxicon 34, 1149–1155.
- Jamunaa, A., Vejayan, J., Halijah, I., Sharifah, S.H., Ambu, S., 2012. Cytotoxicity of
 Southeast Asian snake venoms. J. Ven. Animals Toxins incl. Trop. Dis. 18, 150–
 156.
- Joubert, F.J., Taljaard, N., 1980a. Purification, some properties and amino-acid sequences
 of two phospholipases A (CM-II and CM-III) from *Naja naja kaouthia* venom. Eur.
 J. Biochem. 112, 493–499.

567	Joubert, F.J., Taljaard, N., 1980b. The complete primary structures of three cytotoxins
568	(CM-6, CM-7 and CM-7A) from Naja naja kaouthia (Siamese cobra) snake venom.
569	Toxicon 18, 455–467.
570	Karlsson, E., 1973. Chemistry of some potent animal toxins. Experientia 29, 1319–1327.
571	Kasturiratne, A., Wickremasinghe, R., de Silva, N., Gunawardena, N.K., Pathmeswaran,
572	A., Premaratna, R., Savioli, L., Lalloo, D.G., de Silva, H.J., 2008. The global burden
573	of snakebite: a literature analysis and modelling based on regional estimates of
574	envenoming and deaths. PLoS Med. 5, e218.
575	Kini, R.M., Doley, R., 2010. Structure, function and evolution of three-finger toxins: mini
576	proteins with multiple targets. Toxicon 56, 855-867.
577	Kini, R.M., 2005. Structure-function relationships and mechanism of anticoagulant
578	phospholipase A_2 enzymes from snake venoms. Toxicon 45, 1147–1161.
579	Konshina, A.G., Dubovskii, P.V., Efremov, R.G., 2012. Structure and dynamics of
580	cardiotoxins. Curr. Protein Pept. Sci. 13, 570-584.
581	Kukhtina, V.V., Tsetlin, V.I., Utkin, Y.N., Inozemtseva, L.S., Grivennikov, I.A., 2001.
582	Two forms of nerve growth factor from cobra venom prevent the death of PC12
583	cells in serum-free medium. J. Nat. Toxins 10, 9–16.
584	Kulkeaw, K., Chaicumpa, W., Sakolvaree, Y., Tongtawe, P., Tapchaisri, P., 2007.
585	Proteome and immunome of the venom of the Thai cobra, Naja kaouthia. Toxicon
586	49, 1026–1041.
587	Kulkeaw, K., Sakolvaree, Y., Srimanote, P., Tongtawe, P., Maneewatch, S., Sookrung, N.,
588	Tungtrongchitr, A., Tapchaisri, P., Kurazono, H., Chaicumpa, W., 2009. Human
589	monoclonal ScFv neutralize lethal Thai cobra, Naja kaouthia, neurotoxin. J.
590	Proteomics 72, 270–282.

- Kyi, S.W., Zug, G.R., 2003. Unusual foraging behaviour of *Naja kaouthia* at the Moyingye
 Wetlands Bird Sanctuary, Myanmar. Hamadryad 27, 265–266.
- Laustsen, A.H., Lomonte, B., Lohse, B., Fernández, J., Gutiérrez, J.M., 2015. Unveiling the
 the nature of black mamba (*Dendroaspis polylepis*) venom through venomics and
 antivenom immunoprofiling: identification of key toxin targets for antivenom
 development. J. Proteomics (submitted).
- Lomonte, B., Tsai, W.C., Ureña-Díaz, J.M., Sanz, L., Mora-Obando, D., Sánchez, E.E.,
 Fry, B.G., Gutiérrez, J.M., Gibbs, H.L., Calvete, J.J., 2014a. Venomics of New
 World pit vipers: genus-wide comparisons of venom proteomes across *Agkistrodon*.
 J. Proteomics 96, 103-116.
- Lomonte, B., Pla, D., Sasa, M., Tsai, W.C., Solórzano, A., Ureña-Díaz, J.M., FernándezMontes, M.L., Mora-Obando, D., Sanz, L., Gutiérrez, J.M., Calvete, J.J., 2014b.
 Two color morphs of the pelagic yellow-bellied sea snake, *Pelamis platura*, from
 different locations of Costa Rica: snake venomics, toxicity, and neutralization by
 antivenom. J. Proteomics 103, 137–152.
- Masathien, C., Billings, P., Ratananbanangkoon, K., 1994. Production and characterization
 of monoclonal antibodies neutralizing the postsynaptic neurotoxin 3 from *Naja kaouthia* venom. J. Nat. Toxins 3, 155–163.
- Meng, Q.X., Wang, W.Y/, Lu, Q.M/, Jin, Y., Wei, J.F., Zhu, S.W., Xiong, Y.L., 2002. A
 novel short neurotoxin, cobrotoxin c, from monocellate cobra (*Naja kaouthia*)
 venom: isolation and purification, primary and secondary structure determination,
 and tertiary structure modeling. Comp. Biochem. Physiol. 132C, 113–121.

- 613 Mukherjee, A.K., Maity, C.R., 2001. Biochemical composition, lethality and 614 pathophysiology of venom from two cobras - Naja naja and N. kaouthia. Comp. 615 Biochem. Physiol. B 131, 125–132.
- Pakmanee, N., Khow, O., Kumsap, W., Omori-Satoh, T., Chanhome, L., Sriprapat, S., 616 Sitprija, V., 1998. Envenomation of mice by Thai cobra (*Naja kaouthia*) venom: 617 tolerable venom concentration and exposure time. Toxicon 36, 809-812. 618
- 619 Prakash, S., Mishra, A.K., Raziuddin, M., 2012. A new record of cream coloured morph of 620 Naja kaouthia Lesson, 1831 (Reptilia, Serpentes, Elapidae) from Hazaribag, 621 Jharkhand, India. Biodiversity J. 3, 153–155.
- Reali, M., Serafim, F.G., da Cruz-Hofling, M.A., Fontana, M.D., 2003. Neurotoxic and 622 myotoxic actions of Naja naja kaouthia venom on skeletal muscle in vitro. Toxicon 623 41, 657-665. 624
- 625 Richard, G., Meyers, A.J., McLean, M.D., Arbabi-Ghahroudi, M., MacKenzie, R., Hall, J.C., 2013. In vivo neutralization of alpha-cobratoxin with high-affinity llama 626 single-domain antibodies (V_H Hs) and a V_H H-Fc antibody. PLoS One 8, e69495. 627
- 628 Roncolato, E.C., Campos, L.B., Pessenda, G., Costa e Silva, L., Furtado, G.P., Barbosa, 629 J.E., 2015. Phage display as a novel promising antivenom therapy: a review. Toxicon 93, 79-84.

- Safdari, Y., Farajnia, S., Asgharzadeh, M., Khalili, M., 2013. Antibody humanization 631 methods - a review and update. Biotech. Gen. Eng. Rev. 29, 175-186. 632
- 633 Sakurai, Y., Takatsuka, H., Yoshioka, A., Matsui, T., Suzuki, M., Titani, K., Fujimura, Y.,
- 2001. Inhibition of human platelet aggregation by L-amino acid oxidase purified 634 from Naja naja kaouthia venom. Toxicon 39, 827-1833. 635

636	Stewart, C.S., MacKenzie, C.R., Hall, J.C., 2007. Isolation, characterization and
637	pentamerization of alpha-cobrotoxin specific single-domain antibodies from a naive
638	phage display library: preliminary findings for antivenom development. Toxicon 49,
639	699–709.
640	Stuart, B., Wogan, G., 2012. Naja kaouthia. In: The IUCN Red List of Threatened Species.
641	Version 2014.3. http://www.iucnredlist.org. Downloaded on 19 January 19, 2015.
642	Tan, N.H., Swaminathan, S., 1992. Purification and properties of the L-amino acid oxidase
643	from monocellate cobra Naja naja kaouthia venom. Int. J. Biochem. 24, 967–973.
644	Uetz, P., Hošek, J., 2015. The Reptile Database, http://www.reptile-database.org. Accessed
645	January 26, 2015.
646	Utsintong, M., Talley, T.T., Taylor, P.W., Olson, A.J., Vajragupta, O., 2009. Virtual
647	screening against α-Cobratoxin. J. Biomol. Screening 14, 1109–1118.
648	Viravan, C., Veeravat, U., Warrell, M.J., Theakston, R.G.D., Warrell, D.A., 1986. ELISA
649	confirmation of acute and past envenoming by the monocellate Thai cobra (Naja
	· · · · · · · · · · · · · · · · · · ·
650	kaouthia). Am. J. Trop. Med. Hyg. 35, 173–181.
650 651	
	kaouthia). Am. J. Trop. Med. Hyg. 35, 173–181.
651	<i>kaouthia</i>). Am. J. Trop. Med. Hyg. 35, 173–181. Vonk, F.J., Casewell, N.R., Henkel, C.V., Heimberg, A.M., Jansen, H.J., McCleary, R.J.R.,
651 652	 <i>kaouthia</i>). Am. J. Trop. Med. Hyg. 35, 173–181. Vonk, F.J., Casewell, N.R., Henkel, C.V., Heimberg, A.M., Jansen, H.J., McCleary, R.J.R., Kerkkamp, H.M., Vos, R.A., Guerreiro, I., Calvete, J.J., Wüster, W., Woods, A.E.,
651 652 653	 <i>kaouthia</i>). Am. J. Trop. Med. Hyg. 35, 173–181. Vonk, F.J., Casewell, N.R., Henkel, C.V., Heimberg, A.M., Jansen, H.J., McCleary, R.J.R., Kerkkamp, H.M., Vos, R.A., Guerreiro, I., Calvete, J.J., Wüster, W., Woods, A.E., Logan, J.M., Harrison, R.A., Castoe, T.A., de Koning, A.P., Pollock, D.D., Yandell,
651 652 653 654	 <i>kaouthia</i>). Am. J. Trop. Med. Hyg. 35, 173–181. Vonk, F.J., Casewell, N.R., Henkel, C.V., Heimberg, A.M., Jansen, H.J., McCleary, R.J.R., Kerkkamp, H.M., Vos, R.A., Guerreiro, I., Calvete, J.J., Wüster, W., Woods, A.E., Logan, J.M., Harrison, R.A., Castoe, T.A., de Koning, A.P., Pollock, D.D., Yandell, M., Calderon, D., Renjifo, C., Currier, R.B., Salgado, D., Pla, D., Sanz, L., Hyder,
 651 652 653 654 655 	 <i>kaouthia</i>). Am. J. Trop. Med. Hyg. 35, 173–181. Vonk, F.J., Casewell, N.R., Henkel, C.V., Heimberg, A.M., Jansen, H.J., McCleary, R.J.R., Kerkkamp, H.M., Vos, R.A., Guerreiro, I., Calvete, J.J., Wüster, W., Woods, A.E., Logan, J.M., Harrison, R.A., Castoe, T.A., de Koning, A.P., Pollock, D.D., Yandell, M., Calderon, D., Renjifo, C., Currier, R.B., Salgado, D., Pla, D., Sanz, L., Hyder, A.S., Ribeiro, J.M., Arntzen, J.W., van den Thillart, G.E., Boetzer, M., Pirovano,

- Wallach, V., Wüster, W., Broadley, D.G., 2009. In praise of subgenera: taxonomic status of
 cobras of the genus *Naja Laurenti* (Serpentes: Elapidae). Zootaxa 2236, 26–36.
- Wang, J.P., Teng, C.M., 1990. Comparison of the enzymatic and edema-producing
 activities of two venom phospholipase A₂ enzymes. Eur. J. Pharmacol. 190, 347–
 354.
- Wang, W.J., Shih, C.H., Huang, T.F., 2004. A novel P-I class metalloproteinase with broad
 substrate-cleaving activity, agkislysin, from *Agkistrodon acutus* venom. Biochem.
 Biophys. Res. Comm. 324, 224–230.
- Warrell, D.A., 1995. Clinical toxicology of snakebite in Africa and the Middle East /
 Arabian peninsula. In: Handbook of Clinical Toxicology of Animal Venoms and
 Poisons (Meier, J., White, J., Eds.), pp 433–492. CRC Press.
- 670 Warrell, D.A., 2010. Snake bite. Lancet 375, 77–88.
- Warrell, D.A., 2010b. Guidelines for the management of snake-bites. World Health
 Organization, 152 pp.
- Wongtongkam, N., Wilde, H., Sitthi-Amorn, C., Ratanabanangkoon, K., 2005. A study of
 Thai cobra (*Naja kaouthia*) bites in Thailand. Mil. Med. 170, 336–341.
- Wüster, W., 1996. Taxonomic changes and toxinology: Systematic revisions of the asiatic
 cobras (*Naja naja* species complex). Toxicon 34, 399–406.

678 Figure legends

679

Figure 1: Separation of *N. kaouthia* (**A**) venom proteins by RP-HPLC (**C**), followed by SDS-PAGE (**B**). Two mg of venom were fractionated on a C_{18} column and eluted with an acetonitrile gradient (dashed line), as described in Methods. Fractions were further separated by SDS-PAGE under reducing conditions. Molecular weight markers (Mw) are indicated in kDa. Coomassie-stained bands were excised, in-gel digested with trypsin, and subjected to MALDI-TOF/TOF analysis for assignment to protein families, as shown in Table 1.

687

Figure 2: Composition of *N. kaouthia* venom according to protein families, expressed as
percentages of the total protein content. 3FTx: three-finger toxin; PLA₂: phospholipase A₂;
NGF: nerve growth factor; OHA: Ohanin/vespryn; CRISP: Cysteine-rich secretory protein;
CTL: C-type lectin/lectin-like; NUCL: nucleotidase; PDE: phosphodiesterase; MP:
metalloproteinase; LAO: L-amino oxidase; CVF: Cobra venom factor; CTT:
Cytidyltransferase; UNK: Unidentified proteins; NP: non-protein components. A division
between cytotoxins (CYT) and neurotoxins (NTX) is given for the three-finger toxins.

695

Figure 3: Presence of selected nucleosides and FAD in *Naja kaouthia* venom shown by spiking crude venom with 10 μ g of nucleosides (adenosine, inosine, guanosine) or FAD and separating the venom components by reverse-phase HPLC. If a the peak of a nucleoside coincides with the peak of a venom component, and if mass determination yielded the same mass for the venom component as calculated for the nucleoside, the venom component was judged to consist of the corresponding nucleoside. *N. kaouthia* venom contains small
amounts of adenosine, guanosine, and traces of inosine, but no FAD.

703

Figure 4: (A) Comparison of the phospholipase A₂ activity between the venoms of *Naja kaouthia* and *Micrurus nigrocinctus*. *N. kaouthia* displays some enzymatic activity, although significantly less than *M. nigrocinctus*. **(B)** Comparison of the proteolytic activity between the venoms of *N. kaouthia*, *M. nigrocinctus*, and *Bothrops asper*, evaluated on azocasein. *N. kaouthia* shows negligible proteinase activity.

709

Figure 5: Electrospray ionization-mass spectrometry analysis of *N. kaouthia* venom fractions 6 and 10 (see Fig.1 and Table 1). Samples were analyzed in enhanced multicharge positive mode as described in Methods. Fraction 6 (Nk-6) presented a homogeneous multi-charged series (**A**) which deconvoluted to an isotope-averaged mass (M_{av}) of 7826.0 \pm 0.8 Da (**B**). The multi-charged series of fraction 10 (Nk-10) (**C**) showed a main protein with M_{av} 7619.0 \pm 1.3 Da and a less abundant protein of 7656.0 \pm 0.7 Da (**D**), thus demonstrating difference between fractions 6 and 10.

717

Figure 6: ELISA-based immunoprofiling of human antibodies (**Donor 1 IgG**: protein Apurified IgG antibodies from Donor 1, **Donor 2**: protein A-purified IgG antibodies from Donor 2, **Human negative:** normal human serum from healthy volunteer, never bitten by a snake) to the fractions of *Naja kaouthia* venom separated by RP-HPLC (see Materials and Methods for details). Toxins with the highest Toxicity Score (4, 5, and 6) are marked with a skull symbol.

Peak %		1 0.2	2 0.4	3 0.5	4 1.2	5a 0.3	5b 0.4	5c 3.3				6a 13.5			
	,	2	4	S	2		4	c. C			ίλ	∞.	9		
Mass (kDa)▼			I	I	13	67	54	10			13	10	10		
Peptide ion	m/z		ı	ı	2945.2	1475.7	ı	1728.7 1014.5 1316.6	1453.6 2945.2	1758.8	1315.5 2241.9	1547.7 2184.9 1315.5 2241.9	1315.5 2241.9	2187.9 1011.5	
lon	z	1	1	1	-	-				1					
MS/MS-derived or N-terminal (Nt) sequence		non-peptidic	non-peptidic (Adenosine)	non-peptidic (Guanosine)	XECHNBBSSBTPTTTGCSGGETNCYK	SXDXXBBWEEBS(R ^{Io})		XECHNBBSSBAPTTK BWWSDHR VBPGVNXNCCR	(N ^{da})GXEXNCCTTDR XECHNBBSSBTPTTTGCSGGETNCYK	XECHNBBSSBTPTTK	TWCDAFCSXR TGVDXBCCSTDNCNPFPTR	RVDXGCAATCPTVK TGVDXB <u>C</u> CSTDNCNPFPTR TWCDAFCSXR TGVDXBCCSTDNCNPFPTR	TWCDAFCSXR TGVDXBCCSTDNCNPFPTR	SXFGVTTEDCPDGBNXCFK WH(M ^{ax})XVPGR	
Conf			ı	ı	99	66	ı	96 66	66 66	99	99 99	66 66 66	66 66	99 82.8	
Š			ī	ī	25	11	ī	15 8	12 28	13	13 16	13 14 24	13 21	23 8	
Protein family; ~ related protein *		unknown	non-peptidic (Nucleosides)	non-peptidic (Nucleosides)	3FTx three-finger toxin <i>Naja atra</i> , ∼E2IU03	Cytidyltransferase choline-phosphate cytidylyltransferase A isoform X2 <i>Python bivittatus</i> , ~XP_007440684	unknown	3 FTx cobrotoxin-c <i>Naja kaouthia</i> , ∼PS9276	three-finger toxin Naja atra, ~E21U03	cobrotoxin-b <i>Naja kaouthia</i> , ~P59275	3FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ~P01391	3FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ~P01391	3 FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ∼P01391	muscarinic toxin-like prot.2 <i>Naja kaouthia</i> , ~P82463	

Table 1: Assignment of the RP-HPLC isolated fractions of *Naja kaouthia* venom to protein families by MALDI-TOF-TOF of selected peptide ions from in-gel trypsin-digested protein bands.

	13b	13 a	12b	12a	116	11 a	10			9b		9a
	S	2.5	2.0	2.1	2.0	4.9	10.2			0.4		0.3
	10	22	10	15	10	15	10			10		10
1842.6	2829.3 1087.4 1798.8	1842.6	1610.5 1842.6 1697.6 1157.5 1826.6 2356.0	1842.6	1826.6 1669.5 1232.5 1697.6 1842.6 900.4	1842.7	1315.5 2241.9	1629.6	2241.9 1315.5	1177.5 2186.8	962.4	1415.6 1363.6 2603.1 2260.9
-		-		-		-		1			1	
SWWDFADYGCYCGR	TCPAGBNXCYBMFMVSNBTVPVBR YVCCNTDR NSXXV(K ^{eb})YVCCNTDR	SWWDFADYGCYCGR	GGNNACAAAVCDCDR SWWDFADYGCYCGR TYSYECSBGTXTCK XSGCWPYFK CCBVHDNCYNEAEK XAAXCFAGAPYNNNNYNXDXK	SWWDFADYGCYCGR	CCBVHDNCYNEAEK GDNDACAAAVCDCDR NMXBCTVPNR TYSYECSBGTXTCK SWWDFADYGCYCGR CWPYFK	SWWDFADYGCYCGR	TWCDAFCSXR TGVDXBCCSTDNCNPFPTR	XTCXNCPEMFCGK	TGVDXBCCSTDNCNPFPTR TWCDAFCSXR	TXCYNHXTR TSETTEXCPDSWYFCYK	BYFFETK	CBNPNPEPSGCR AXTMEGNBASWR EDHPVHNXGEHSVCDSVSAWVTK GXDSSHWNSYCTETDTFXK
99	66 66	99	66 66 66 66	66	99 99 99 99 98,2	99	66 66	99	99 98.8	66 66	99	66 66 66
14	14 10 15	9	20 19 14 20 22	11	22 20 13 16 9	10	13 15	14	11 10	7 17	11	10 18 29 27
Phospholipase A ₂	3FTx cytotoxin 3 <i>Naja kaouthia</i> , ∼P01446	Phospholipase A2 acidic PLA2 - D <i>Naja sputatrix</i> , ~Q91900	Phospholipase A2 acidic PLA2 - 1 <i>Naja atra</i> , ~P00598	Phospholipase A ₂ acidic PLA ₂ - 1 <i>Naja sputatrix</i> , ~Q91900	Phospholipase A2 acidic PLA2 - 1 <i>Naja kaouthia</i> , ∼P00596	Phospholipase A ₂ acidic PLA ₂ - 2 <i>Naja sagittifera</i> , ~P60044	3FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ~P01391	weak neurotoxin NNAM2 <i>Naja atra</i> , ~Q9YGI4	alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ~P01391	3FTx muscarinic toxin-like prot.3 <i>Naja kaouthia</i> , ∞P8?464	NGF Naja sputatrix, ~Q5YF89	Nerve growth factor NGF <i>Naja kaouthia</i> , ∼P61899

19	18b	18a		17b	17a	16b	16a		15	14b	14a	
0.2	4.6	0.8		1.2	0.5	3.7	5.0		2.4	1.1	1.3	
10	10	13		10	12	10	21		Ξ	10	25	
ı	1087.4 948.4 924.4 2918.3 1771.8	1315.5	1798.8	1771.8 2918.2 1087.4	2141.9 2270.0 930.4 913.4 1810.8 1495.6	2861.3 1412.6 1568.7 1798.7 1087.4	1087.5	1420.6	1412.6 1798.8 1568.7 1087.4	2829.3 1087.4 1798.8	1087.4	
		-	1				-	1			1	
	YVCCNTDR GCXDVCPK MFMVATPK GCXDVCPBSSXXVBYVCCNTDRCN SSXXV(K ^{ca})YVCCNTDR	TWCDAFCSXR	NSXXV(K ^{ca})YVCCNTDR	SSXXV(K ^{ca})YVCCNTDR GCXDVCPBSSXXVBYVCCNTDRCN YVCCNTDR	ADVTFDSNTAFESXVVSPDK ADVTFDSNTAFESXVVSPDBK EWAVGXAGK SPPGNWBK TVENVGVSBVAPDNPER FDGSPCVXGSPGFR	TCPAGBNXCYBMFMMSDXTXPVBR MFMMSDXTXPVK MFMMSDXTXPVBR NSXXV(K°°)YVCCNTDR YVCCNTDR	YVCCNTDR	CHNTBXPFXYK	MFMMSDXTXPVK NSXXV(K ⁴⁸)YVCCNTDR MFMMSDXTXPVBR YVCCNTDR	TCPAGBNXCYBMFMVSNBTVPVBR YVCCNTDR NSXXV(K ^{en})YVCCNTDR	YVCCNTDR	
ı	66 66 66 66	94.7	99	99 83,4 99	66 66 66 66 66	99 99 99 99	man	66	66 66 66	66 66	91.6	
ı	$10 \\ 10 \\ 11 \\ 14$	8	16	14 7	15 17 13 10 19	11 11 11 10	man	16	11 15 10	10 12 15	Ţ	
unknown	3FTx cytotoxin 8 <i>Naja atra</i> , ∼Q91124	3FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ∼P01391	cytotoxin 3a Naja atra, ~Q98959	3FTx cytotoxin 4N <i>Naja atra</i> , ~Q9W6W9	Ohanin/vespryn thaicobrin <i>Naja kaouthia</i> , ∼P82885	3FTx three-finger toxin <i>Naja atra</i> , ∼E2IU04	3FTx three-finger toxin <i>Naja atra</i> , ~E2IU04	cytotoxin 5V Naja atra, ~Q9W716	3FTx cytotoxin VC-1 <i>Naja oxiana</i> , ∼Q9PS33	3FTx cytotoxin 2 <i>Naja kaouthia</i> , ~P01445	acidic PLA ₂ - D <i>Naja sputatrix</i> , ~Q91900 3FTx cytotoxin VC-1 <i>Naja oxiana</i> , ~Q9PS33	

28	27	26c	26b		26a	25c	25b	25a	24d	24c
0.9	0.3	0.3	0.1		0.1	0.1	0.3	0.1	0.7	0.2
44	46	55	70		250	61	108	139	40	57
2713.2	1603.8 1235.7	2673.3 2922.5 2517.2	2030.0 1386.7 1875.1	2517.2	1339.7 1718.9	1449.8 2421.3 1719.0 1459.8	1434.7 1244.7 2106.1 1634.7 1518.9 1355.7	1434.7 1244.7 2106.1	2354.1 1471.6 1912.6 1012.4 1430.7 2689.0 1099.5	1430.8 1099.5 2689.1 1471.7 1430.7
-				1		<u> </u>				
DDCDXPEXCTGBSAECPTDVFBR	VYEMVNAXNTMYR VTXDXFGEWR	RSPXEECFBBNDYEEFXEXAR FGXBXNEFFBENENAWYYXNNXR SPXEECFBBNDYEEFXEXAR	FFYXDGNENFHVSXTAR BXDXFVHDFPR XXXNXPXNABSXPXTVR	SPXEECFBBNDYEEFXEXAR	EGWYVNMGPMR TFVTADYVXVCSTSR	VVSXNVXCTECR FHECNXGNXXCDAVVYNNXR ETPVXSNPGPYXEFR YXGYXNVXFDDK	DFYTFDSEAXVK SP(N ^{db})NXWVEER RPDFSTXYXEEPDTTGHK YCSGGTHGYDNEFK XWNYFHSTXXPK AATYFWPGSEVK	DFYTFDSEAXVK SP(N ^{da})NXWVEER RPDFSTXYXEEPDTTGHK	CPTXTNBCXAXXGPHFTVSPK TAPAFBFSSCSXR (N ^{db})GHPCBNNBGYCY(N ^{db})GK GCFDXNMR YXEFYVXVDNR HDCDXPEXCTGBSAECPTDSXBR DYBEYXXR	YXEFYVXVDNR DYBEYXXR HDCDXPEXCTGBSAECPTDSXBR TAPAFBFSSCSXR YXEFYVXVDNR
99	66 66	66 66	66 66	99	66 66	99 99 93,5	86 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	66 66	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	99 99 99
11	11 12	8 10 17	12 10 13	20	9 15	8 9 7	16 15 20 14 9	9 8	25 11 12 12 12	15 10 14 15
Metalloproteinase	Metalloproteinase mocarhagin, <i>Naja mossambica</i> , ~Q10749	L-amino acid oxidase L-amino-acid oxidase <i>Naja oxiana</i> , ∼P0D191	Cobra venom factor CVF <i>Naja kaouthia</i> , Q91132	L-amino-acid oxidase <i>Naja oxiana</i> , ~P0DI91	L-amino acid oxidase L-amino-acid oxidase <i>Naja atra</i> , ~A8QL58	5'-nucleotidase ecto-5'-nucleotidase 1 <i>Micrurus fulvius</i> , ~U3FYP9	Phosphodiesterase phosphodiesterase 1 <i>Micrurus fulvius</i> , ~U3FAB3	Phosphodiesterase phosphodiesterase 1 <i>Micrurus fulvius</i> , ~U3FAB3	Metalloproteinase kaouthiagin N <i>aja kaouthia</i> , ∼P82942	Metalloproteinase kaouthiagin <i>Naja kaouthia</i> , P82942

interpreted MS/MS spectrum.	with the following abbreviations: ^{ox} : oxidized; ^{da} : deamidated; ^{ca} : carbamyl; ^{am:} amide; ^{na:} Na cation; ^{al:} ammonia loss; ^{dl:} delta ^{H2C2} ; man: manually	Possible, although unconfirmed/ambiguous amino acid modifications suggested by the automated identification software are shown in parentheses.	determined by nESI-MS of selected RP-HPLC peaks, in Da. X: Leu/Ile; B: Lys/Gln; Z: pyroglutamate (2-oxo-pyrrolidone carboxylic acid).	of ProteinPilot [®] (ABSciex) $\mathbf{\nabla}$: estimated mass by SDS-PAGE under reducing conditions, in kDa.	* Cysteine residues are carbamidomethylated, unless underlined. Confidence (Conf) and Score (Sc) values are calculated by the Paragon algorithm	1207.6 1 ATXDXFGEWR 1807.9 1 YXEFY(M ^d)VVDNXMYR 2983.4 1 AABDDCDXPEXCTGBSAECPTDVFBR
	^{ca} : carbamyl; ^{am:} amide; ^{na:}	ations suggested by the aut	. X: Leu/Ile; B: Lys/Gln;	GE under reducing conditi	d. Confidence (Conf) and S	MYR JBSAECPTDVFBR
	Na cat	omated	Z: pyı	ions, ir	score (S	66 66
	ion; ^{al:}	lidenti	rogluta	n kDa.	šc) valu	10 12 10
	ammonia loss; ^{di:} delta ^{H2C2} ; man: manually	fication software are shown in parentheses,	mate (2-oxo-pyrrolidone carboxylic acid).	∇ : observed isotope-averaged masses as	ies are calculated by the Paragon algorithm	cobrin <i>Naja kaouthia</i> , ~Q9PVK7

Ŷ	∞	7	6	S	4	Whole venom	Peak
0.7 (3:4 mix ²)	8.0	0.9	32.3	4.0 (3:4:33 mix ²)	1.2		%
Nerve growth factor (43%) NGF <i>Naja kaouthia</i> , ~P61899 NGF <i>Naja sputatrix</i> , ~Q5YF89 3FTx (57%) muscarinic toxin-like prot.3 <i>Naja kaouthia</i> , ~P82464 alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ~P01391 weak neurotoxin NNAM2 <i>Naja atra</i> , ~Q9YGI4	3FTx weak tryptophan-containing neurotoxin, <i>Naja kaouthia</i> , ∼P82935	3FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ∼P01391 muscarinic toxin-like prot.2 <i>Naja kaouthia</i> , ~P82463	3FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ∼P01391	Cytidyltransferase (7.5%) choline-phosphate cytidylyltransferase A isoform X2 <i>Python bivittatus</i> , ~XP_007440684 Unknown (10%) 3FTx (82.5%) cobrotoxin-c <i>Naja kaouthia</i> , ~P59276 three-finger toxin <i>Naja atra</i> , ~E2IU03 cobrotoxin-b <i>Naja kaouthia</i> , ~P59275	3FTx three-finger toxin <i>Naja atra</i> , ∼E2IU03		Protein family
<u>~</u>	~2	0.36 (0.17-0.72)	0.10 (0.02-0.18)	<mark>0.07</mark> (<mark>0.001-0.19</mark>)	0.24 (0.12-0.41)	0.24 (0.12-0.41)	LD ₅₀ (95 % C.L.)
			0.1 [Karlsson, 1973]	80 (cobrotoxin- c), 400 (cobrotoxin-b) [Meng et al., 2002]		0.115*	Reported LD ₅₀ (mg/kg)
	<mark>~4</mark>	2.5	326	41.3	15.4	423.2	Toxicity Score ¹ % / LD ₅₀ (kg/mg)

 Table 2: Median lethal doses
 (LD₅₀s) of Naja kaouthina venom and the RP-HPLC isolated fractions

39

19	18	17	16	15	14	13	12	11	10
0.2	5.4 (4:23 mix ²)	1.7 (5:12 mix ²)	8.7	2.4	2.4 (27:23 mix ²)	7.5 (1:2 mix ²)	4.1 (51:49 mix ²)	6.9 (49:20 mix ²)	10.2
Unknown	3FTx (15%) alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ∼P01391 3FTx (85%) cytotoxin 8 <i>Naja atra</i> , ~Q91124	Ohanin/vespryn (29%) thaicobrin <i>Naja kaouthia</i> , ~P82885 3FTx (71%) cytotoxin 4N <i>Naja atra</i> , ~Q9W6W9 cytotoxin 3a <i>Naja atra</i> , ~Q98959	3FTx three-finger toxin <i>Naja atra</i> , ∼E2IU04	3FTx cytotoxin VC-1 <i>Naja oxiana</i> , ~Q9PS33	3FTx cytotoxin VC-1 <i>Naja oxiana</i> , ~Q9PS33 (54%) cytotoxin 2 <i>Naja kaouthia</i> , ~P01445 (46%)	Phospholipase A ₂ (33%) acidic PLA ₂ - D <i>Naja sputatrix</i> , ~Q91900 3FTx (66%) cytotoxin 3 <i>Naja kaouthia</i> , ~P01446	Phospholipase A ₂ acidic PLA ₂ - 1 <i>Naja sputatrix</i> , ~Q9I900 (51%) acidic PLA ₂ - 1 <i>Naja atra</i> , ~P00598 (49%)	Phospholipase A ₂ acidic PLA ₂ - 2 <i>Naja sagittifera</i> , ~P60044 (71%) acidic PLA ₂ - 1 <i>Naja kaouthia</i> , ~P00596 (29%)	3FTx alpha-elapitoxin-Nk2a <i>Naja kaouthia</i> , ~P01391
>0.25	<mark>1.00</mark> (<mark>0.43-2.93</mark>)	1.36 (0.62-2.03)	<mark>0.75</mark> (<mark>0.30-1.66</mark>)	<mark>1-2.5</mark>	<mark>1.60</mark> (<mark>1.21-2.28</mark>)	0.9 (0.09-1.29)	3.74 (2.17-8.75)	<mark>5.38</mark> (<mark>2.01-55.94</mark>)	>10
					1.2 (cytotoxin 2) [Joubert and Taljaard, 1980b]			10 [Chuman et al., 2000]	
Δ	<mark>5.4</mark>	1.25	<mark>11.6</mark>	< <mark>2.5</mark>	1.57	8.3	1.1	<mark>1.3</mark>	<u>^</u>

28 0.9	27 0.3	0.5 26 (1:1:3 mix ²)	25 0.5 (4:1 mix ²)	24 1.2	23 0.3	(0.1 IIIX)	22 (6.1 min. ²)	
Metalloproteinase cobrin <i>Naja kaouthia</i> , ~Q9PVK7	Metalloproteinase mocarhagin, <i>Naja mossambica</i> , ~Q10749	L-amino acid oxidase (20%) L-amino-acid oxidase <i>Naja atra</i> , ~A8QL58 L-amino-acid oxidase <i>Naja oxiana</i> , ~P0DI91 Cobra venom factor (20%) CVF <i>Naja kaouthia</i> , Q91132 L-amino acid oxidase (60%) L-amino-acid oxidase <i>Naja oxiana</i> , ~P0DI91	Phosphodiesterase (80%) phosphodiesterase 1 <i>Micrurus fulvius</i> , ~U3FAB3 5'-nucleotidase (20%) ecto-5'-nucleotidase 1 <i>Micrurus fulvius</i> , ~U3FYP9	Metalloproteinase kaouthiagin <i>Naja kaouthia</i> , P82942	Nucleotidase endonuclease domain-containing 1 prot Micrurus fulvius, ~tr U3FCT9	C-type lectin/lectin-like (14%) LP-Pse-6 CTL Pseudonaja modesta, ~R4G314	Cysteine-rich secretory protein (86%) natrin-1 <i>Nata atra</i> ~07T1K6	C-type lectin/lectin-like LP-Pse-6 CTL <i>Pseudonaja modesta</i> , ~R4G314 Cysteine-rich secretory protein (86%) natrin-1 <i>Nata atra</i> ~07T1K6
			>0.25			>1.5		>0.5
<mark>N.t.</mark>	N.t.	N.t.	۵	N.t.	N.t.	\triangle		$\overline{\nabla}$

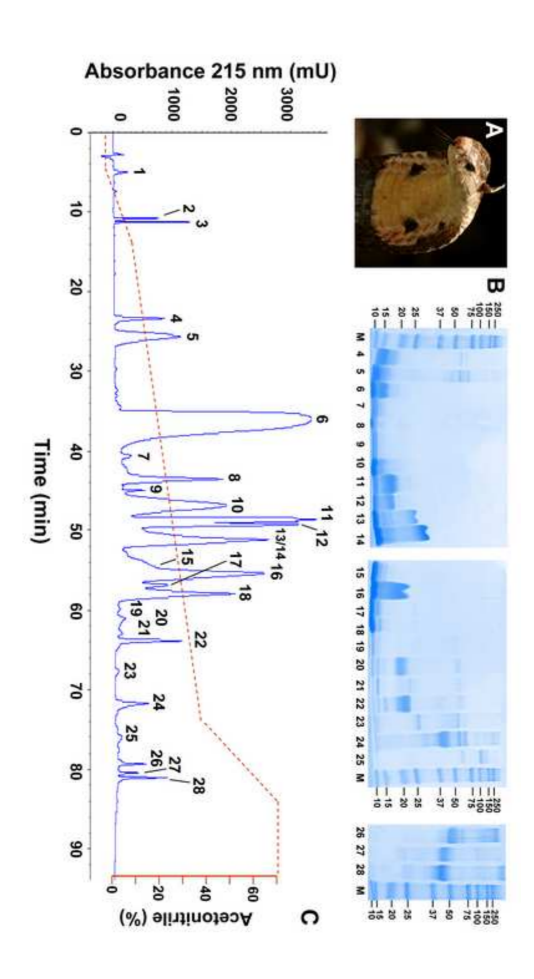
*http://snakedatabase.org/pages/LD50.php#legendAndDefinitions

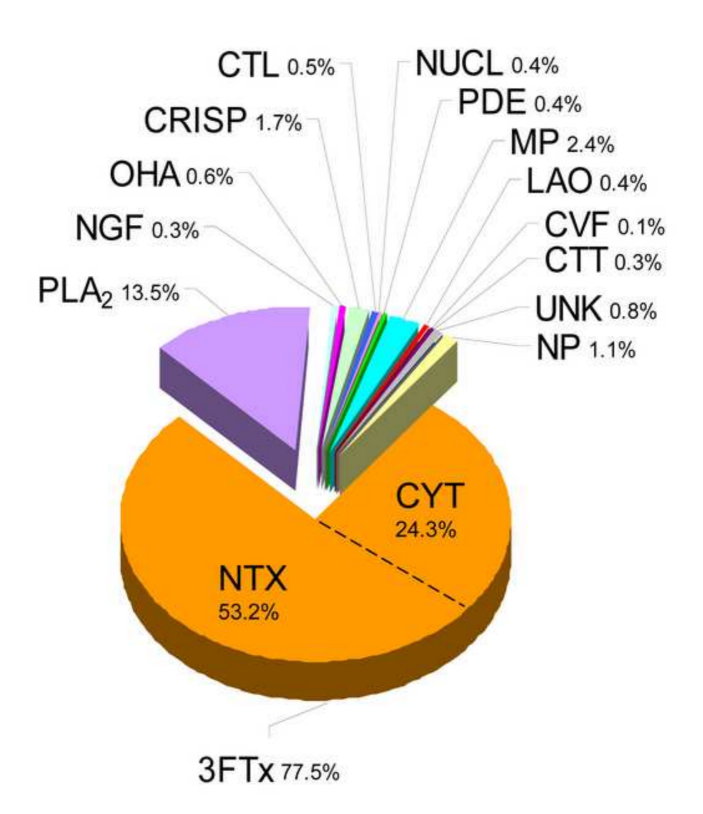
¹Toxicity Score was defined as the ratio of protein fraction abundance (%) in the venom divided by its estimated median lethal dose

 (LD_{50}) for CD-1 mice by i.v. injection.

indicated in the table. ² Mix indicates that the fraction did not contain a pure, isolated toxin, but instead a mixture of 2-4 different toxins in variable ratios

N.t. : not tested





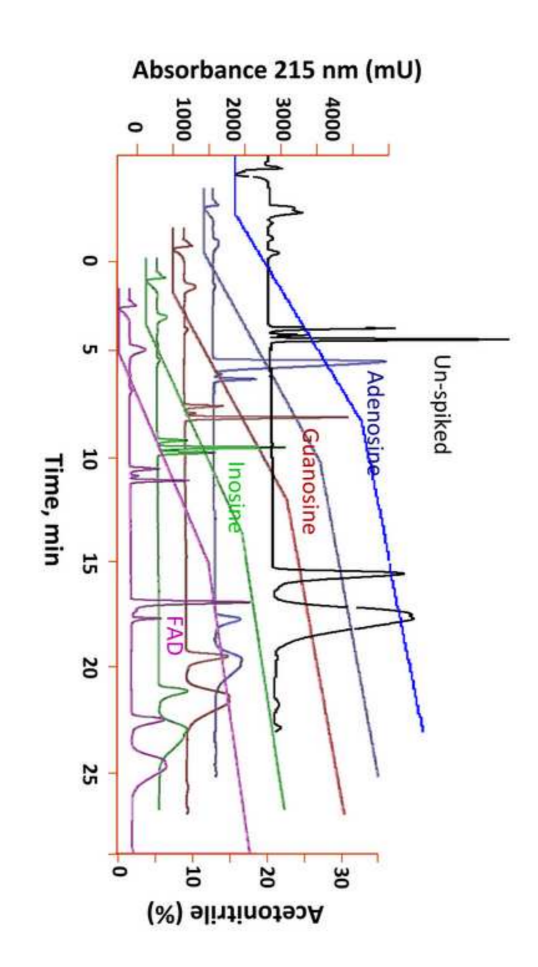
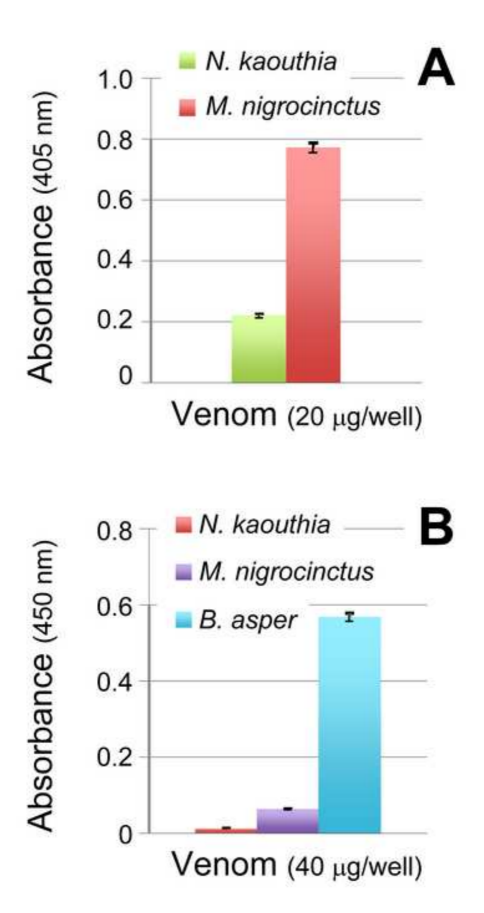


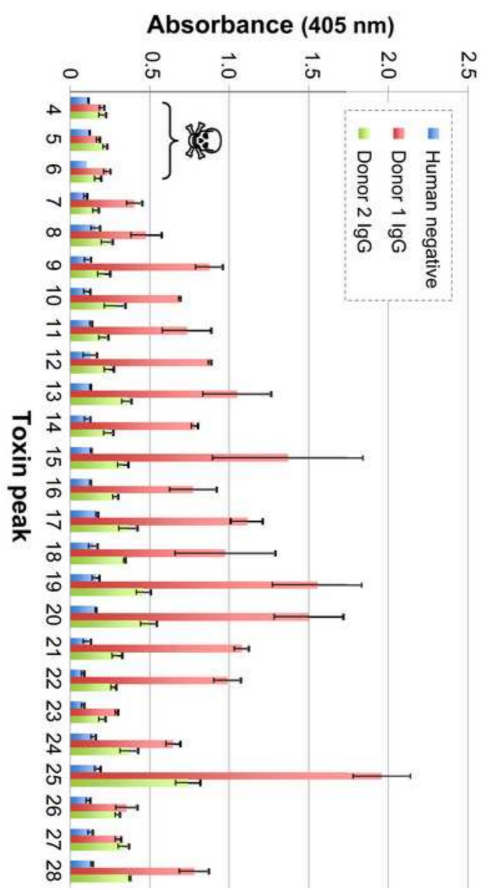
Figure 3 Click here to download high resolution image Figure 4 Click here to download high resolution image



Intensity (cps) 2.00e6 4.00e6 6.00e6 8.00e6 1.00e7 2.0e6 3.0e6 4.0e6 4.6e6 1.0e6 0.00 0.00 800 10+ 9+ 800 9+ 979.1 983.9 00 1000 958.1 953.4 962.5 1000 1124.2 1118.9 1089.4 1094.8 1100.0 1200 16.2 1200 6⁺ 1278.2 **6**+ 1305.3 1311.7 1277.3 1400 1400 m/z (Da) 1524.7 1582.1 1564.6 Nk-10 1566.3 NK-6 1600 1600 ဂ 5.0e7 1.0e8 1.4e8 3.0e7 5.9e7 1.0e7+ 2.0e7 4.0e7 5.0e7 0.0-7400 0.0 7600 7619.0 7656.0 7800 7826.0 7800 ٤ Nk-10 8000 8000 Nk-6

Figure 5 Click here to download high resolution image

Figure 6 Click here to download high resolution image



Graphical abstract Click here to download high resolution image

