Functional and Structural Diversification of the Anguimorpha Lizard Venom System* 

Bryan G. Fry, Kelly Winter, Janette A. Norman, Kim Roelants, Rob J. A. Nabuurs, Matthias J. P. van Osch, Wouter M. Tewissse, Louise van der Weerd, Judith E. Mcnaughtan, Hang Fai Kwok, Holger Scheib, Laura Greisman, Elazar Kochva, Laurence J. Miller, Fan Gao, John Karas, Denis Scanlon, Feng Lin, Sanjaya Kuruppu, Chris Shaw, Lily Wong, and Wayne C. Hodgson

Venom has only recently been discovered to be a basal trait of the Anguimorpha lizards. Consequently, very little is known about the timings of toxin recruitment events, venom protein molecular evolution, or even the relative physical diversifications of the venom system itself. A multidisciplinary approach was used to examine the evolution across the full taxonomical range of this ~130 million-year-old clade. Analysis of cDNA libraries revealed complex venom transcriptomes. Most notably, three new cardiovascular peptide toxin types were discovered (celostatin, cholecystokin, and YY peptides). The latter two represent additional examples of convergent use of genes in toxic arsenals, both having previously been documented as components of frog skin defensive chemical secretions. Two other novel venom gland-overexpressed modified versions of other protein frameworks were also recovered from the libraries (epididymal secretory protein and ribonuclease). Lectin, hyaluronidase, and veficolin toxin types were sequenced for the first time from lizard venoms and shown to be homologous to the snake venom forms. In contrast, phylogenetic analyses demonstrated that the lizard natriuretic peptide toxins were recruited independently of the form in snake venoms. The latter two represent additional examples of convergent use of genes in toxic arsenals, both having previously been documented as components of frog skin defensive chemical secretions. Two other novel venom gland-overexpressed modified versions of other protein frameworks were also recovered from the libraries (epididymal secretory protein and ribonuclease). Lectin, hyaluronidase, and veficolin toxin types were sequenced for the first time from lizard venoms and shown to be homologous to the snake venom forms. In contrast, phylogenetic analyses demonstrated that the lizard natriuretic peptide toxins were recruited independently of the form in snake venoms. The de novo evolution of helokinestatin peptide toxin encoding domains within the lizard venom natriuretic gene was revealed to be exclusive to the helodermatid/anguid subclade. New isoforms were sequenced for cysteine-rich secretory protein, kallikrein, and phospholipase A2 toxins. Venom gland morphological analysis revealed extensive evolutionary tinkering. Anguid glands are characterized by thin capsules and mixed glands, serous at the bottom of the lobule and mucous toward the apex. Twice, independently this arrangement was segregated into specialized serous protein-secreting glands with thick capsules with the mucous lobules now distinct (Heloderma and the Lanthanotus/Varanus clade). The results obtained highlight the importance of utilizing evolution-based search strategies for biodiscovery and emphasize the largely untapped drug design and development potential of lizard venoms. Molecular & Cellular Proteomics 9:2369–2390, 2010.

Because of significant differences in anatomy of the venom delivery system and distant phylogenetic relatedness, it had been long assumed that the venom systems of snakes and helodermatid lizards were independently evolved (1–3). However, both lineages were recently revealed to be members of a clade (Toxicofera) that also included several lineages of other lizards recently shown to be venomous (4, 5). In contrast to the hypothesis of independent origins, this new perspective revealed that lizard and snake venom systems are homologous but highly differentiated descendants of an early evolved venom system in squamates. The basal condition was incipient glands on both the mandibular and maxillary regions with the reverse, resulting in the modern condition seen today. The lizard venom delivery system is less sophisticated than the high pressure injection mechanism of the front fanged advanced snakes, and the vast majority of the species pose trivial direct medical risks to human (with the exception of helodermatids and large varanids such as Varanus komodoensis). However, the effects of envenomation from medi-
cally important species such as Heloderma (for example) may be clinically complex with symptoms, including extreme pain, acute local swelling, nausea, fever, faintness, myocardial infarction, tachycardia, hypotension, and inhibition of blood coagulation (6–12).

It has been shown previously that venoms evolve via toxin recruitment events. In this process, a gene encoding for a normal body protein (typically one involved in key regulatory processes or bioactivity) is duplicated, and the copy is selectively expressed in the venom gland (13). Toxins are sourced at different times during the evolutionary history from a wide range of tissues, are of disparate scaffolds, and have diverse ancestral bioactivities. Functionally important toxin types are reinforced through adaptive evolution involving explosive duplication and diversification, creating a venom gland-specific multigene family. The likelihood for neofunctionalization is increased through random mutation, gene conversion, and unequal crossing over (14). Although the molecular scaffold of the ancestral protein is conserved, derived activities emerge through mutations of the surface chemistry (14–17). Because of the inherent chemical arms race between venomous animals and their prey, toxins have exquisite biological targeting with sometimes as little as a single amino acid change radically changing specificity or potency.

Previous work demonstrated that a core set of venom genes evolved in the common Toxicofera ancestor (4), and subsequent toxin recruitment events facilitated diversification into complex and varied venoms (18–20) (Fig. 1). The anguimorph lizards, of which the helodermatid lizards were long thought to be the only venomous members, are a very large assemblage with great diversity in body size and ecological niche occupied (21). To date, only the venoms of two extant helodermatid lizards and a small, taxonomically limited handful of varanid lizard species have been studied with the anguid lizard clade almost entirely ignored. Currently, seven protein families have been identified as recruited for use as toxins in lizard venoms (see Table I): helofensin (lethal toxin isoforms), exendin, CRiSP,1 kallikrein, natriuretic peptides, and type III phospholipase A2 (4, 20, 22). Two of these (CRiSP and kallikrein) are homologous to forms found in snake venoms (4, 18, 19, 23). An additional toxin type (helokinestatin peptides) are not the result of a recruitment event but rather are unique proline-rich derivatives within the ancestral gene (located upstream from the natriuretic peptide-encoding region) that are post-translationally cleaved off to form discrete bioactive

---

1 The abbreviations used are: CRiSP, cysteine-rich secretory protein; BLAST, basic local alignment search tool; RACE, rapid amplification of cDNA ends; CCK, cholecystokinin; T, tesla; PLA2, phospholipase A2; CRD, carbohydrate recognition domain; BNP, brain natriuretic peptide; CNP, C-type natriuretic peptide; VIP, vasoactive intestinal peptide.
peptides (22, 24). Helokinestatin peptides are thus a rare case of a single gene being mutated to additionally encode multiple products. Helokinestatin peptides are thus far known only from Heloderma venom, and the two varanid natriuretic precursor sequences reported to-date lack these derivatives (4, 20, 22).

As the venoms of Anguimorpha lizards have received less attention than those of the more medically important venomous snakes, it remains unknown to what extent the different toxin types have diversified and undergone structural and functional innovation. Detailed knowledge of the anguimorph toxins therefore is crucial for insight into the structure, function, and composition of the ancestral Toxicofera venom system and to address the question of how multiple lineages achieved evolutionary innovations. A thorough understanding of the molecular and functional diversity is also necessary to fully exploit the therapeutic potential of these venoms.

Transcriptome analyses by random sequencing of cDNA libraries has been proven to be an effective way of identifying novel isoforms of previously reported toxins as well as toxins of new protein classes (see Refs. 4, 18, 20, 22, 25, and 26), thus providing new insights into the molecular evolutionary origins. In this study, we have integrated multidisciplinary data to provide an in-depth investigation of anguimorph lizard venom molecular evolution to (a) characterize the venom transcriptomes of representative anguimorph lizards, (b) determine the timing of toxin recruitment events and patterns of toxin diversification, and (c) characterize changes in the venom gland structure and delivery architecture.

**MATERIALS AND METHODS**

**Species Examined**

Adult stage specimens of the following species were examined by either construction of a mandibular venom gland cDNA library (C), histology (H), or magnetic resonance imaging (M) as noted: Anguidae *Celestus warreni* (Haiti) (C, H, and M), * Gerrhonotus infernalis* (Texas) (C, H, and M), and *Ophisaurus apodus* (Greece) (C, H, and M); Heterodermatidae *Heloderma horridum horridum* (captive-bred offspring of founder stock from unknown geographical origin) (C and M); Lanhanotidae *Lanhanotus bomeensis* (Borneo) (M); Shinisauridae *Shinisaurus crocodilurus* (captive-bred offspring of founder stock from unknown geographical origin) (H); and Varanidae *Varanus acanthurus* (Newman, Western Australia) (C and M), *Varanus albigularis* (captive-bred offspring of founder stock from unknown geographical origin) (M), *Varanus eremius* (Giralia Station, Western Australia) (C), and *V. giganteus* (Sandstone, Western Australia) (C and M), *Varanus gilleni* (captive-bred offspring of founder stock from unknown geographical origin) (C and M), *Varanus glauerti* (Kununurra, Western Australia) (C), *Varanus gouldii* (Kununurra, Western Australia) (C and M), *Varanus indicus* (Alligator River, Northern Territory, Australia) (C), *V. komodoensis* (captive-bred offspring of founder stock from unknown geographical origin) (C, H, and M), * Varanus mertensi* (Kununurra, Western Australia) (C and M), *VaranusMitchelli* (Kununurra, Western Australia) (M), *Varanus panoptes panoptes* (Kununurra, Western Australia) (C and M), *Varanus prasinus* (captive-bred offspring of founder stock from unknown geographical origin) (M), *Varanus salvator* (Bali, Indonesia) (M), *Varanus scalaris* (Kununurra, Western Australia) (C and M), *Varanus tristis orientalis* (captive-bred offspring of founder stock from unknown geographical origin) (C and M), and *Varanus varius* (Mallacoota, Victoria, Australia) (C and H). Different individuals were used for each method.

**cDNA Library Construction and Analysis**

To investigate the relationship between toxins and the encoding transcripts and also facilitate molecular evolutionary analyses, venom gland cDNA libraries were constructed from glands that had been surgically excised from living specimens under general anesthesia and immediately cryogenically preserved. RNA was harvested using the Qiagen RNAeasy Midi kit with subsequent selection of mRNAs using the Oligotex Midi kit. cDNA libraries were constructed using the Clontech Creator SMART cDNA Library Construction kit and transformed into One Shot Electrocompetent GeneHogs (Invitrogen). Isolation and sequencing of inserts was conducted at the Australian Genome Research Facility using BDtv3.1 chemistry with electrophoretic separation on an AB330xl. 384 colonies were sequenced, inserts were screened for vector sequences, and those parts were removed prior to analysis and identification.

Toxin sequences were identified by comparison of the translated DNA sequences with previously characterized toxins using a BLAST search of the Swiss-Prot/UniProt protein database (http://www.expasy.org/tools/blast/). In sequence alignments, signal peptides were ascertained utilizing the SignalP 3.0 Server (http://www.cbs.dtu.dk/services/SignalP/). Molecular phylogenetic analyses of toxin transcripts were conducted using the translated amino acid sequences. Comparative sequences from other venomous reptiles and outgroups were obtained through BLAST searching (http://www.expasy.org/tools/blast/) using representative toxin sequences. To minimize confusion, all sequences obtained in this study are referred to by their GenBank™ accession numbers (http://www.ncbi.nlm.nih.gov/sites/entrez?db=Nucleotide), and sequences from previous studies are referred to by their UniProt/Swiss-Prot accession numbers (http://www.expasy.org/cgi-bin/sprot-search-ful) except for the previously reported snake veficolin sequences, which are referred to by their GenBank accession numbers. Resultant sequence sets were aligned using the program ClustalX followed by visual inspection for errors. When presented as sequence alignments, the leader sequence is shown in lowercase, the prepro region is underlined, cysteines are highlighted in black, and functional residues are in bold. Data sets were analyzed using Bayesian inference implemented on MrBayes, version 3.0b4. The analysis was performed by running a minimum of $1 \times 10^6$ generations in four chains and saving every 100th tree. The log likelihood score of each saved tree was plotted against the number of generations to establish the point at which the log likelihood scores of the analysis reached their asymptote, and the posterior probabilities for clades were established by constructing a majority rule consensus tree for all trees generated after the completion of the burn-in phase. Sequence alignments can be obtained by e-mailing Dr. Bryan Grieg Fry.

**RT-PCR**

10-mg samples of lyophilized venom from each lizard species (H. suspectum and H. horridum) were dissolved separately in 1 ml of cell lysis/mRNA protection buffer supplied by Dynal Biotech. Polyadenylated mRNA was isolated by the use of magnetic oligo(dT) beads as described by the manufacturer (Dynal Biotech). mRNA was eluted in 20 μl of RNase-free water, and first strand cDNA synthesis for subsequent RACE reactions was performed using a SMART-RACE kit (Clontech) essentially as described by the manufacturer. Exendin RACE reactions were amplified using two sets of sense primers (S1, ATGAAAATCATCCTTTGGCTGTGTGTTT; S2, ATGAAAAGCATC-
CCTGTCACTGGCAATAGGCT) and antisense primers (AS1, CGAT-GGGCCCTTGGTGTAGT; AS2, TAACCGGATGCGGAGAGTGGT) for the cDNAs by thermostable polymerase (Invitrogen). These primers were complementary (S1/AS1) or partially complementary (S2/AS2) to a domain of nucleotide sequences in the 5’ and 3’ regions of the exendin-2 precursor cDNA cloned previously (H. suspectum cinctum (GenBank accession number EU790760). The PCR cycling procedure was as follows: an initial denaturation step for 1 min at 94 °C followed by 35 cycles consisting of denaturation for 30 s at 94 °C, primer annealing for 30 s at 62 °C (S1/AS1) or 58 °C (S2/AS2), and extension for 3 min at 72 °C. Gel electrophoresis of the PCR products from the venom library of each species was followed by further purification, cloning using a pGEM-T vector system (Promega Corp.), and subsequent sequencing using an ABI 3100 automated capillary sequencer. Kallikrein RACE reactions were amplified using two sets of sense primers (S1, ATCATTGGAGCCAGAATGCC; S2, ATCATGGAGCCAGAATGCC) and antisense primers (AS1, GGCCGGCTTGGGCTAGG; AS2, GAACAGGTGAAATAATGTTG) for the cDNAs by thermostable polymerase (Invitrogen). These primers were complementary to 5’ and 3’ regions of the precursor cDNA EU790962 cloned from H. suspectum cinctum in this study (S1/AS1) and backward translations of the protein sequence P43685 reported previously from H. horridum (S2/AS2) (34). The PCR cycling procedure was as follows: an initial denaturation step for 1 min at 94 °C followed by 35 cycles consisting of denaturation for 30 s at 94 °C, primer annealing for 30 s at 62 °C, and extension for 3 min at 72 °C. Gel electrophoresis of the PCR products from the venom library of each species was followed by further purification, cloning using a pGEM-T vector system (Promega Corp.), and subsequent sequencing using an ABI 3100 automated capillary sequencer.

Bioactivity Characterization of Cholecystokinin, Celestinogen, Goannatorytoxin, Helokininatin, Natriuretic, and YY Peptides

Peptide Synthesis—Peptides were synthesized on a CEM Liberty Peptide Synthesizer using Fmoc (N-(9-fluorenyl)methoxy carbonyl) solid-phase peptide chemistry. The peptides were cleaved from the solid-phase resin with TFA/H2O/triisopropylsilane/3,6-dioxa-1,8-octanedithiol (90:2.5:2.5:5) for 2 h. The crude peptides were isolated by ether precipitation, dissolved in 30:70 (v/v) ACN/H2O, and lyophilized. The crude linear peptides were reversed-phase HPLC-purified (Agilent 1100 HPLC System) before forming the single disulfide bond (in the natriuretic peptide only) with dipyridyldithiol (1 eq) in 100 mM ammonium acetate (peptide concentration, 1 mg/ml; 30 min). The pure cyclized natriuretic peptide was isolated by directly applying the ammonium acetate solution to a reversed-phase HPLC column, isolating pure peptide fractions, and lyophilization of the products. The identity of the pure cyclized peptide was confirmed by high resolution mass spectrometry on an Agilent Q-TOF 6510 LC/MS mass spectrometer.

Anesthetized Rat Studies—Male Sprague-Dawley rats were anesthetized with pentobarbital sodium (60–100 mg/kg intraperitoneally). A midline incision was made in the cervical region, and cannulas were inserted into the trachea, jugular vein, and carotid artery for artificial respiration (if required), administration of peptides, and measurement of blood pressure, respectively. Arterial blood pressure was recorded using a Gould P23 pressure transducer connected to a PowerLab system. All results were expressed as change in mean arterial pressure.

Guinea Pig Isolated Ileum Studies—Guinea pigs of either sex were killed by CO2 inhalation and exsanguination. The ileum was removed and cut into 2-cm segments. Segments of ileum were then mounted on wire tissue holders under 1-g resting tension in 5-ml organ baths containing physiological salt solution of the following composition: 118.4 mM NaCl, 4.7 mM KCl, 1.2 mM MgSO4, 1.2 mM KH2PO4, 25.0 mM NaHCO3, 11.1 mM glucose, 2.5 mM CaCl2 continuously bubbled with carbogen (95% O2, 5% CO2) and maintained at 34 °C. The preparation was allowed to equilibrate for 30 min. A cumulative concentration-response curve to bradykinin (1 nM–1 μM) using half-log unit increments was then obtained. The peptide (0.1 μM) was then added and left to equilibrate for 30 min. A cumulative concentration-response curve to bradykinin (1 nM–1 μM) was then repeated in the presence of the peptide as described above. Isometric contractions were measured via a Grass force displacement transducer (FT03) and recorded using a PowerLab system. Responses to bradykinin in the presence of the peptide were expressed as a percentage of the maximum response obtained from the initial bradykinin curve.

Venom Gland Structural Characterization

Histological sections were prepared from the intact head and the excised venom gland and delivery system. Whole heads were removed, and a cut was made to the underside to allow fast penetration of the fixative (10% neutral buffered formalin). After a minimum of 2 days, excess tissue was removed, and specimens were immersed in Kristensen’s decalcification solution and placed on a rotor for up to 3 weeks (depending on the size of the head). Before processing, the heads were bisected longitudinally for cutting transversely at 3 μm in two separate blocks. The processing schedule was as follows: 10% formalin, 2 h; absolute ethanol, 4 × 1 h; histolene, 3 × 1 h; paraffin
wax, 2 × 90 min. The sections were taken every 100 μm, and matching sections were stained with periodic acid-Schiff’s stain and Masson’s trichrome stain. In other specimens, the venom gland, venom duct, and adjacent bony and muscular tissues were excised and placed in decalcifying solution (Cal-Ex, Fisher) for 72–168 h. Each sample was dehydrated and cleared through a progressive ethanol series and Cyto-Sol (Fisher) prior to embedding in Paraplast (Fisher). Serial sections were cut at 10–12 μm. All species were sectioned in the frontal plane; when available, the contralateral venom gland and delivery system were sectioned either parasagittally or transversely. Sections were stained using a variant of Van Gieson’s stain, which provides clear distinction between connective tissue, muscle, and epithelium, or with hematoxylin and eosin.

Magnetic resonance imaging (MRI) was used to examine the three-dimensional shape and internal anatomy of the venom glands from individual specimens of each species. For all except Heloderma, formalin-ethanol-fixed heads were first submerged in Fomblin (Solvay Solexis) and placed under vacuum to prevent air artifacts. Depending on head size, imaging was performed on either 9.4-Tesla (T) (small/medium) or 17-T (large) vertical 89-mm-bore systems (Bruker BioSpin, Rheinstetten, Germany) with a Bruker Micro2.5 gradient system of 1 T/m and a transmit/receive birdcage radiofrequency coil with a diameter of 10–30 mm. Bruker ParaVision 4.0 software was used for image acquisition. Anatomical images were acquired using a three-dimensional gradient echo sequence. The field of view and matrix were varied to fit the individual samples, resulting in voxel sizes between 403 and 703 μm3. Imaging parameters were as follows: echo time, 8 ms; repetition time, 40 ms; flip angle, 20°; 4–8 averages; total scan time, between 3 and 9 h/sample depending on size and resolution. A Philips Achieva, 1.5-T clinical MRI scanner (Philips Healthcare, Best, The Netherlands) using a surface coil with a diameter of 47 mm was used to scan the preserved heads of H. horridum horridum (CAS34292) from the California Academy of Science. Several scout images were performed, and the coil was repositioned to obtain maximum signal to noise in the anatomical area to be imaged. A three-dimensional three-dimensional shape and internal anatomy of the venom glands from individual specimens of each species. For all except Heloderma, formalin-ethanol-fixed heads were first submerged in Fomblin (Solvay Solexis) and placed under vacuum to prevent air artifacts. Depending on head size, imaging was performed on either 9.4-Tesla (T) (small/medium) or 17-T (large) vertical 89-mm-bore systems (Bruker BioSpin, Rheinstetten, Germany) with a Bruker Micro2.5 gradient system of 1 T/m and a transmit/receive birdcage radiofrequency coil with a diameter of 10–30 mm. Bruker ParaVision 4.0 software was used for image acquisition. Anatomical images were acquired using a three-dimensional gradient echo sequence. The field of view and matrix were varied to fit the individual samples, resulting in voxel sizes between 403 and 703 μm3. Imaging parameters were as follows: echo time, 8 ms; repetition time, 40 ms; flip angle, 20°; 4–8 averages; total scan time, between 3 and 9 h/sample depending on size and resolution. A Philips Achieva, 1.5-T clinical MRI scanner (Philips Healthcare, Best, The Netherlands) using a surface coil with a diameter of 47 mm was used to scan the preserved heads of H. horridum horridum (CAS34292) from the California Academy of Science. Several scout images were performed, and the coil was repositioned to obtain maximum signal to noise in the anatomical area to be imaged. A three-dimensional gradient echo sequence was acquired with the following parameters: echo train length, 11; repetition time/echo time/flip angle, 1000 ms/65 ms/90°; field of view, 100 mm; 220 slices; acquisition voxel size, 0.2 × 0.2 × 0.2 mm3; number of signal averages, 1; scan time, 65 min (1:05:27). Images with different angulations were reconstructed afterward on a Vitrea work station (Vital Images Inc., Minnetonka, MN). Image segmentation of the glands was performed manually in Amira 4.1 (Mercury Computer Systems), and three-dimensional surface renderings were generated.

RESULTS

Toxin Transcript Types Recovered—BLAST analyses of transcripts recovered from the venom gland cDNA libraries identified seven protein types with sequence homology to previously characterized toxicoferan toxins (Table I). Three toxin types previously only sequenced from snake venoms were recovered in this study: lectin (recovered from O. apodus and V. indicus), hyaluronidase (recovered from H. suspectum cinctum and G. infernalis), and veficolin (V. komodoensis). CRiSP, kalliirein, natriuretic peptide, and PLAr transcripts were sequenced for the first time from anguid lizards, and additional isoforms were sequenced from the helodermatid and varanid libraries. Nerve growth factor was sequenced for the first time from Heloderma. Similar to the multidomain-encoding transcripts from Heloderma, homologous anguid natriuretic mRNA full-length transcripts also were multido- main-, multiproduct-encoding (natriuretic and helokinestatin peptides). However, the characteristic helokinestatin proline-rich domains remained consistently absent from the varanid natriuretic transcripts (Fig. 2A). Although exendin-3 and -4 are present in large amounts in both Heloderma species and easily purified by reversed-phase HPLC, in contrast, exendin-1 and -2 peaks were only visible in H. suspectum. Specific primer-directed PCR studies showed strong levels of exendin-1 in H. suspectum, but trace amounts were detected in H. horridum (Fig. 3A). In contrast, exendin-2 was also strongly detected in H. suspectum but entirely absent from H. horridum (Fig. 3B).

Toxin Molecular Evolution—The CRiSP toxins showed differential variation in structural and functional domains, particularly in the residues that guide bioactivity (Fig. 4). The cysteine pattern and other structural residues were highly conserved as was the PR1 tetrad functional domain. However, the PR1 binding site displayed a much lower level of conservation. Similarly, the ion channel-binding residues in the CRD were also quite variable. In two variants, newly evolved cysteines were present: one in the C. warreni isoform GU441473 upstream from the first ancestral cysteine and the other one in between the second and third ancestral cysteines of the two isoforms recovered from V. indicus (GU441469 and GU441470). Both V. indicus isoforms were also truncated, lacking the last six cysteines.

Newly evolved cysteines were also shown to be present in two different kalliirein variants: one type from the anguid clade and two others from the varanids (Fig. 5). In a novel form of the kalliirein toxins recovered from both the C. warreni and G. infernalis libraries, there was not only a newly derived cysteine at alignment position 219 but also a deletion of the last ancestral cysteine. In the variant recovered from both the V. acanthurus and V. scalaris libraries, a new cysteine was evolved at alignment position 124, resulting in an odd number of cysteines.

A long C-terminal tail was revealed in the protein translations of the H. suspectum PLA2 mRNA sequences in this study that had not been seen in either of the reported H. suspectum sequences previously determined by protein sequencing (P80003 and P16354) (Fig. 6). The translated forms of the full-length mRNA sequences corresponding to the region at which the protein sequences stop (Gly for P80003 and GR for P16354) are rich in dibasic proteolytic sites (GRKR-SQKRRK for EU790967 and GRKR-SQKRRK for EU790968) (Fig. 6), common recognition sites for proteolytic removal of segments in precursors from both vertebrates (27) and invertebrates (28). Therefore, P16354 may represent an incompletely proteolyzed product relative to P80003. The C-terminal tail of EU790968 is much longer than that of EU790967, which terminates with the proteolysis motif. The full-length sequences from the anguids and varanids revealed that this RK proteolytic cleavage motif is ubiquitously present in anguimorph isoforms and thus represents a basal condition (Fig. 6).
Diversification of Anguimorpha Lizard Venom System

Phylogenetic analysis of the lizard venom natriuretic peptides showed them to be the result of an independent toxin recruitment event relative to the natriuretic peptides from snakes venoms (Fig. 7). Each lineage formed a monophyletic group that was reciprocally monophyletic, all lizard toxins clustering together and all snake toxins clustering together but with the two toxin types being phylogenetically distinct from each other. Consistent with the lack of atrial natriuretic peptide in the reptile physiology (29), the lizard venom types were nested within the BNP clade. In contrast, as reported previously (30), the snake forms were all nested within the CNP clade (Fig. 7). The ASDEN sequence that links the Heloderma post-translationally cleaved helokinestatin peptide isoforms encoded by the multidomain, multiproduct natriuretic transcript (22) was only partially conserved between variants in G. infernalis (AAEEN, AEEN, and AEGEEN), respectively) but not at all conserved in the same proline-rich region of the C. warreni transcript (Fig. 2A).

TABLE I

| Toxin type          | Species                  | GenBank accession no.                                   | Bioactivity                                                                 |
|---------------------|--------------------------|--------------------------------------------------------|-----------------------------------------------------------------------------|
| CRiSP               | All                      | EU790958, GU441462, GU441463, GU441464, GU441465, GU441466, GU441467, GU441468, GU441469, GU441470, GU441471, GU441472, GU441473 | Blockage of ryanodine receptors and potassium channels, producing lethargy, paralysis, and hypothermia (57–61) |
| Helokinestatin      | C. warreni, G. infernalis| GU441510, GU441511                                       | Bradykinin inhibition (22, 24); bioactivity determined for new variants from anguid venoms in this study (Fig. 2D). |
| Hyaluronidase       | G. infernalis, H. suspectum | EU790961, GU441476                                      | Venom spreading factor (42)                                                  |
| Kallikrein          | All                      | EU790962, GU441477, GU441478, GU441479, GU441480, GU441481, GU441482, GU441483, GU441484, GU441485, GU441486, GU441487, GU441488, GU441489, GU441490, GU441491, GU441492, GU441493, GU441494, GU441495, GU441496, GU441497, GU441498, GU441499, GU441500, GU441501, GU441502, GU441503, GU441504, HM437246 | Release of bradykinin from kinogen (34, 62–65); a derivative form also cleaves fibrinogen (66) |
| Lectin              | O. apodus, V. indicus    | GU441505, GU441506                                       | Snake venom forms have been shown to have various modes of action upon platelet aggregation (67, 68) |
| Natriuretic peptide | C. warreni, G. infernalis, H. suspectum, V. glauerti, V. scalaris | EU790965, GU441507, GU441508, GU441509, GU441510, GU441511 | Hypotension induction leading to loss of consciousness; mediated through the binding of GC-A, resulting in the relaxation of cardiac smooth muscle (4, 16, 20); structure-function studies in this study revealed residues crucial for bioactivity (Fig. 2, B and C) |
| Nerve growth factor | H. suspectum cinctum     | EU790966                                                | Uncharacterized                                                             |
| PLA,                | C. warreni, H. suspectum, V. gilleni, V. glauerti, V. scalaris, V. tristis | EU790967, EU790968, GU441524, GU441525, GU441526, GU441527, GU441528. | Inhibition of epinephrine-induced platelet aggregation (4, 69)               |
| Veficolin           | V. komodoensis           | HM641898                                                | Uncharacterized                                                             |

2374 Molecular & Cellular Proteomics 9.11
FIG. 2. A, alignment of natriuretic peptide/helokinestatin-encoding precursors from the mandibular venom glands of G. infernalis (GU441510) (1), C. warreni (GU441511) (2), H. suspectum circum (EU790965) (3), V. scalaris (GU441508) (4), V. glauerti (GU441509) (5), V. varius (Q2XXL8) (6), V. komodoensis (B6CJV0) (7), and V. glauerti (GU441509) (8). Bounding boxes surround post-translationally cleaved bioactive peptides, functionally important prolines are highlighted in magenta, cysteines are in black, and two typically invariant natriuretic peptide intraring residues (Asp7 and Ile9) are in green. The signal sequence is in lowercase.

B, Effect of post-translationally cleaved bioactive natriuretic peptide isoforms on (B) mean arterial pressure or (B*) endothelium denuded aortic ring. C, structure-function relationships of a natriuretic peptide variant from G. infernalis that lacks the two invariant amino acids Asp7 (7-D) and Ile9 (9-I) on (C) mean arterial pressure or (C*) endothelium denuded aortic ring. D, Effect of helokinestatin peptides (0.1 mM; n = 5) on cumulative concentration-response curves to bradykinin (1 nM–0.3 mM); peptide sequence tested follows each taxon name. NP, natriuretic peptide; PE, phenylephrine. In panels B and D, number before each taxon identifies in A the precursor of the peptide being tested.
the smallest sized cleavage fragment with full cardiovascular activity and receptor CCK-A binding activity was an octapeptide domain seen in normal body cholecystokinin peptides homologous to YY peptides normally expressed in the intestine (Fig. 10A). Bioactivity testing revealed a potent unique triphasic action, rapid biphasic hypertension followed by prolonged hypotension (Fig. 10, B and C). The third type, celestopoxin, recovered from C. warreni, did not show any significant homology to any known protein type (E < 10) (Fig. 11). Two isoforms (GU441474 and GU441475) that differed only in a 12-amino acid insertion at alignment positions 135–147 were recovered. Both peptides had an arginine-rich cleavage motif to liberate the bioactive domain from the propeptide (Fig. 11). Contained within the bioactive domain was a sequence repeat (ALGVVGGLPVPK and LPGVAGGLPVPK) that differed crucially in the formation of a proline bracket missing in the first repeat that has a Leu residue in place of the Pro residue. Celestopoxin analog testing revealed that this internal proline-bracketed motif LPGVAGGLPVPK is responsible for full activity with the activity of this fragment being equipotent to the activity of the full-length peptide in producing hypotension, whereas the ALGVVGGLPVPK fragment was inactive. Neither adrenoreceptor antagonists nor 5-hydroxytryptamine antagonists could prevent or reverse the response.

In addition to the three peptide types characterized above, also recovered were mutant versions of the epididymal secretory protein and ribonuclease molecular scaffolds (Table II), each displaying evidence of the typical toxin-characteristic duplication and diversification (Figs. 12 and 13). The epididymal secretory protein sequences recovered from the libraries of three varanids were notable in being significantly longer than corresponding normal body proteins, including an additional eight cysteines (Fig. 12). Within these venom gland...
isoforms, two variant classes were present that differed in a deletion in alignment positions 115–157 that removes four of the 20 cysteines (Fig. 12) with potential resultant consequences for structure and function. The ribonuclease sequences from *G. infernalis* lack two of the ancestral cysteines present in the normal body form (Fig. 13). Two of these isoforms had a frameshift mutation near the C terminus, resulting not only in slightly truncated sequences and the loss of cysteines, but also in the loss of PR1 tetrad (magenta), PR1 structurally important residues (green), PR1 substrate binding site (dark blue), CRD possible ion channel blocking sites (light blue), cysteines (black), and otherwise unassigned conserved surface histidines (red). The signal sequence is in lowercase. < and > designate incomplete N and C termini of the precursor, respectively. * designates the N terminus of a post-translationally processed protein sequence.

Fig. 4. Sequence alignment of CRiSP toxins from mandibular venom glands (f–16) and related toxins from maxillary venom glands of snakes (17 and 18). 1, Varanus giganteus (GU441466); 2, V. acanthurus (Q2XXR0); 3, *H. horridum* (Q91055); 4, *G. infernalis* (GU441472); 5, *H. suspectum cinctum* (EU790958); 6, *V. varius* (Q2XXP1); 7, *V. komodoensis* (B6CJU4); 8, *V. gilleni* (GU441467); 9, *V. panoptes* (GU441464); 10, *V. gouldii* (GU441465); 11, *V. mertensi* (GU441463); 12, *O. apodus* (GU441462); 13, *C. warreni* (GU441473); 14, *V. indicus* (GU441469); 15, *V. tristis* (GU441468); 16, *V. scalaris* (GU441471); 17, *T. stejnegeri* (P60623); 18, *P. australis* (Q8AVA4). Highlighted sequences correspond to PR1 tetrad (magenta), PR1 structurally important residues (green), PR1 substrate binding site (dark blue), CRD possible ion channel blocking sites (light blue), cysteines (black), and otherwise unassigned conserved surface histidines (red). The signal sequence is in lowercase. < and > designate incomplete N and C termini of the precursor, respectively. * designates the N terminus of a post-translationally processed protein sequence.
two of the ancestral cysteines but also producing two new cysteines (Fig. 13).

Structural Variation of Venom System—The combination of MRI and histological analysis revealed extensive variations in the relative size and shape of the mandibular venom glands, the central lumina, and the associated duct system (Figs. 14 and 15). Differences were notable in both the number and physical orientation of the gland compartments as well as the relative encapsulation of the glands. The number of discernible compartments ranged from three discernible compart-

**Fig. 5.** Sequence alignment of kalilikerin toxins from mandibular venom glands. 1. *H. horridum* (P43685) sequenced previously (34); 2. *H. horridum* sequenced in this study (HM437246); 3. *H. suspectum* (EU790962); 4. *G. infernalis* (GU441495); 5. *C. warreni* (GU441497); 6. *C. warreni* (GU441498); 7. *V. glauerti* (GU441481); 8. *V. gilleni* (GU441480); 9. *V. gouldii* (GU441483); 10. *V. panoptes* (GU441490); 11. *V. mertensi* (GU441484); 12. *V. glauerti* (GU441487); 13. *V. tristis* (GU441488); 14. *V. indicus* (GU441482); 15. *V. gouldii* (GU441489); 16. *V. gouldii* (GU441492); 17. *M. mitchelli* (Q2XXN0); 18. *V. komodoensis* (B6CJU6); 19. *V. scalaris* (VSCA_CL3Contig1); 20. *V. scalaris* (GU441478); 21. *V. scalaris* (GU441479); 22. *C. warreni* (B6CJU5); 23. *G. infernalis* (GU441496); 24. *C. warreni* (GU441473). Cysteines are highlighted in black, and the signal sequence is shown in lowercase. The prepro region is underlined. < designates an incomplete N terminus of the precursor. * designates the N terminus of a post-translationally processed protein sequence.
ments in *Lanthanotus* (Fig. 15D), six compartments in both *H. horridum* and *H. suspectum* (Fig. 15, A and B), and six compartments in all *Varanus* examined in this study in gland arrangements homologous to that previously shown for *V. komodoensis* (19) to one compartment per tooth for up to 16 compartments in the anguid lizards as shown for *Celestus*, *Gerrhonotus*, and *Ophisaurus* (posteriorly located were other compartments that secreted only mucus and thus were not included in the count) (Figs. 14 and 15F).

The anguid lizards all had mixed glands with a serous portion occupying the bottom of the glands and a mucous part above it, leading to the duct, which opens at the base of the teeth, as characteristic for dental glands with the entire arrangement covered by a single thin capsule (Fig. 14C and 15F).

*Molecular Composition of Anguimorpha Venoms*—In this study, sequencing of a myriad of toxin types (Table II) from all cDNA libraries indicated the common presence of complex venom transcriptomes in Anguimorpha lizards. Multiple transcripts of the majority of toxins were recovered from individual cDNA libraries with sequences thus supported by overlapping clones. Isoforms were recovered with significant variations, including changes in key functional residues and changes in the number and spacing of cysteine residues, modifications typically associated with neofunctionalization (evolution of novel bioactivities), a pattern consistent with accelerated diversification in toxin multigene families as observed in snake venoms (13, 14, 18). The kallikrein toxins in particular showed explosive gene duplication and diversification with extensive duplication events preceding speciation.

In addition to new isoforms of previously isolated and characterized toxin types, we discovered three new peptide toxin types that we have named cholecystoxin (from *V. varius*), celestoxin (from *C. warreni*), and goannatyrotoxin (from *V. glauerti*). Of particular note is that two of our new peptide toxin
types (cholecystokinin and goannatyrotoxin) represent additional cases of convergence for the selection of a gene type for use as a toxin. Cholecystokinin and YY peptides have both been previously characterized as bioactive components in frog skin chemical arsenals. The cholecystokinin-derived frog peptides such as caerulein are potent ligands of CCK receptors that interfere with the physiological pathways of endogenous CCK and gastrin and cause acute pancreatitis, vomiting, diarrhea, decreased blood pressure, changes in thermoregulation, and inhibition of exploratory and feeding behaviors (31). A similar level of potency was observed for the lizard venom cholecystokins. Endocrine effects of very low concentrations (subnanomolar) of circulating CCK are critical for nutrient homeostasis in the mammal, stimulating exocrine pancreatic secretion, gallbladder emptying, and bowel motility as well as inducing satiety. Higher concentrations of frog skin CCK-like peptides (caerulein) have been shown to stimulate pancreatic acinar cell basolateral exocytosis of zymogens central to the development of the rodent model of pancreatitis. It is interesting that CCK-like peptides secreted via the exocrine route can exhibit the toxic effects previously only attributed to a pharmacologic animal model. In contrast, although the goannatyrotoxin has a triphasic effect upon the cardiovascular system (biphasic hypertension followed by...
prolonged hypotension), the frog skin YY peptides have broad spectrum antibiotic activity (32, 33). The hypotensive celestoxins did not display significant homology to any known peptide type. We also recovered from the libraries two protein types (epididymal secretory protein and ribonuclease) normally expressed in myriad non-salivary tissues and with a diversity of ancestral bioactivities (Table III).

The significant differences between the *H. suspectum cinctum* kallikrein toxins obtained in this study and the previously reported kallikrein toxin from *H. horridum* venom (P43685) (34) must be noted. This is particularly in stark contrast with the high degree of similarity of our *H. suspectum cinctum* toxins with other anguimorph lizard kallikrein toxins (Fig. 5). Our sequence alignments show that the tertiary structure determined by cysteine bonds is likely to be affected if the previously reported sequence P43685 is indeed correct. P43685 was reported as having 13 cysteines (missing the second ancestral cysteine with two new cysteines in atypical positions). Conspicuously, the last cysteine of the *H. horridum* sequence is absent from all other lizard toxin sequences previously reported or recovered in this study but is instead a characteristic of snake venom isoforms. Another significant difference between the helodermatid sequences is an insertion in P43685 at positions 126–132 that is not present in any of the clones from this study or any of the previously reported varanid lizard kallikrein toxins. These sequence differences are markedly higher than analogous sequence differences in CRiSP, exendin, lethal toxin 1, or PLA2 between both Heloderma species. RT-PCR investigations revealed that primers designed using the reported *H. horridum* sequence P43685 did not react with *H. horridum* cDNA, whereas conversely, primers designed utilizing our *H. suspectum* sequence EU790962 from this study reacted strongly with the *H. horridum* cDNA library (Fig. 3C). Sequencing of the kallikrein isoform encoded in our *H. horridum* cDNA library (GenBank accession number HM437246) revealed it to be very similar to our *H. suspectum* sequence although contrasting starkly with the reported P43685 sequence. Thus, we conclude that the

**Fig. 8. Bayesian phylogenetic reconstruction of veficolin toxins and related non-toxin ficolin sequences.**

**Table II.** Novel toxin types recovered from Anguimorpha lizard mandibular venom gland cDNA libraries

| Toxin type       | Species               | GenBank accession no.                                                                 | Bioactivity                                      |
|------------------|-----------------------|---------------------------------------------------------------------------------------|-------------------------------------------------|
| Celestoxin       | *C. warreni*          | GU441474, GU441475                                                                    | Shown in this study to be hypotensive (Fig. 11)  |
| Cholecystoxin    | *V. varius*           | GU441461                                                                              | Shown in this study to bind CCK-A receptor (Fig. 9)|
| Epididymal secretory protein | *V. gouldii, V. indicus, V. komodoensis* | GU441519, GU441520, GU441521, GU441522, GU441523                                  | Uncharacterized                                  |
| Goannatyrotoxin  | *V. eremius*          | GU441529                                                                              | Shown in this study to have a hypertensive/hypotensive triphasic effect (Fig. 10, B and C) |
| Ribonuclease     | *G. infernalis*       | GU441512, GU441513, GU441514, GU441515, GU441516, GU441517, GU441518,                 | Uncharacterized                                  |
previously reported H. horridum sequence P43685 (34) contains multiple errors. The impact of this is not limited to interpretations of lizard venom molecular evolution but also decimates the central conclusion of a study that examined convergent mutation in kallikrein toxins between helodermatid lizards and shrews and in particular relied upon the questionable insert at alignment position 126–132 as a key element (35).

Toxin Structure-Function Relationships—Our functional testing of cholecystokinin, celestoxin, goannatyrotoxin, helokinestatin, and natriuretic peptide analogs demonstrated that all these peptide toxins target the vascular smooth muscle. We were able to demonstrate that the cholecystokinin peptide activity is mediated by an eight-amino acid region with a post-translationally modified tyrosine (sulfated tyrosine) essential for activity, the same structure-function relationship as that of the mammalian endogenous body form (36, 37) (Fig. 9). Proline brackets also guided the activities of helokinestatin (Fig. 2D), goannatyrotoxin (Fig. 10), and celestoxin (Fig. 11) peptides.

Residue replacement studies of a novel natriuretic peptide from G. infernalis allowed us to demonstrate that both the aspartate at ring-position 7 and isoleucine at ring-position 9 contributed equally but both must be present for full activity in the endothelium-denuded aortic rings (Fig. 2C). The differential effect upon the mean arterial pressure of anesthetized animals, with the isoleucine 9 mutant producing a much greater effect than the aspartate 7 indicating action perhaps upon different pathways or receptor subtypes in the circulating systems. However, the variable bioactivity of other iso-

![FIG. 9. A, sequence alignment of precursors of V. varius cholecystokinin (GU441461) (1) from the mandibular venom gland, the frog skin defensive skin toxins from Litoria xanthomera (P56264) (2) and Phyllomedusa sauvagei (Q7LCZ4) (3), and the body peptides from Python molurus (Q8JHB9) (4) and Homo sapiens (P06307) (5). A box indicates a minimal bioactive processed peptide (both the long (22-amino acid) and short (eight-amino acid) forms of V. varius cholecystokinin (GU441461) are shown). Y with a black background represents the key bioactive post-translationally modified sulfatyrosine, * designates the only bioactive peptide domains sequenced to date. B, relative binding of cholecystokinin receptor A of the two variants highlighted in boxes in A (one is 22 amino acids, and one is eight amino acids), a variant of CCK8 with a phosphotyrosine instead of the sulfatyrosine, and human CCK8 with a sulfatyrosine.

![FIG. 10. A, sequence alignment of precursors of V. eremius goannatyrotoxin (GU441529) (1), the frog skin antimicrobial peptide P80952 from Phyllomedusa bicolor (2), and the related non-toxin, ordinary body peptides from H. sapiens (P10082) (3), Anguilla japonica (Q7CL2) (4), and Gallus gallus (P28673) (5). A box indicates a post-translationally processed bioactive peptide. The second Y with a black background is in amide form, * designates the N and C termini of a post-translationally processed peptide sequence. B and C, effect of intravenous injection of V. eremius goannatyrotoxin (GU441529). P1, phase 1; P2, phase 2; P3, phase 3; MAP, mean arterial pressure.
forms that contain these two invariant residues (GU441511 from *C. warreni* and GU441510 from *G. infernalis*) yet were only weakly active demonstrates that bioactivity is mediated by additional residues (Fig. 2B). The *C. warreni* isoform that is lessened in bioactivity contains the intraloop functional residues but does possess prolines in its C-tail (Fig. 2B). C-tail prolines have been shown previously to be responsible for the lessened activity of snake venom isoforms (16) and thus are the likely reason for the lowered activity of this isoform. In this isoform, either intraloop invariant residues are conserved or the likely reason for the lowered activity of this isoform. In this isoform, the proline-kinked tail is the more likely cause. Thus, the reduced activity in GU441509 from *V. glauerti* may be due to the otherwise invariant valine (non-polar) at ring position 12 being changed to threonine (polar).

Using our previously described methodology (18), molecular modeling studies revealed that several features are conserved in the CRiSPs investigated in this study (Fig. 4). Noteworthy are the conserved cysteine pattern and the tetrad in the PR1 domain, which was speculated to form part of an active site of unknown function (38). A third conserved feature is the structurally important residues, despite the extensive amount of cysteines, postulated by Guo et al. (38); albeit, we did not find all of the proposed amino acids conserved.

The suspected binding site in the CRiSP PR1 domain (38) is less conserved and is expected to affect substrate specificity if the proteins would indeed expose enzymatic activity. Of the three sequence patches lining up to form the postulated binding site, the first is the most conserved. After four highly variable positions, the following sequence motif could be deduced: [R][T/V][V/V][G/K][V/I/L][G][E][N][F/Y][S] (square brackets enclose conserved, functional residues). It has to be noted that where more than one residue occurs in a certain position the second and third amino acids are present in one case only. The second sequence stretch accounting for the putative substrate binding site (Fig. 4, alignment positions 130–138) could also be put into a sequence motif (although this motif is less conserved than the first motif), [G/P/A]XX[P/Q]XX[M/V/K]X, with variants such as alanine in position 2 in the snake toxin representatives Trimersurus stejnegeri (P60623) and Pseudechis australis (Q84VA4), glutamine in position 5 in the helodermatid sequences, and at position 8 valine in *T. stejnegeri* and *P. australis* and lysine in *V. tristis*. The third patch of amino acids to participate in substrate binding is variable (alignment 179–185).

Five residues in the CRiSP CRD were brought forward as possible interaction partners with K+ channels (Fig. 4). In *T. stejnegeri*, these residues were Tyr150, Cys212, Thr213, Arg214, and Glu215 as well as the C-terminally located Phe244. Whereas Tyr150 was supposed to only interact with the ion channel, the remaining residues were thought to be involved in the actual blocking of this protein. In *H. suspectum*, position 125 was a histidine rather than a tyrosine, the latter of which was observed in all other proteins studied. Not surprisingly, the cysteine in position 212 is conserved throughout all sequences because cysteines are predominantly involved in maintaining the three-dimensional structure of the proteins. The three residues following this cysteine are not conserved in the cysteine active site of unknown function (38). A third conserved feature is the structurally important residues, despite the extensive amount of cysteines, postulated by Guo et al. (38); albeit, we did not find all of the proposed amino acids conserved.

Using our previously described methodology (18), molecular modeling studies revealed that several features are conserved in the CRiSPs investigated in this study (Fig. 4). Noteworthy are the conserved cysteine pattern and the tetrad in the PR1 domain, which was speculated to form part of an active site of unknown function (38). A third conserved feature is the structurally important residues, despite the extensive amount of cysteines, postulated by Guo et al. (38); albeit, we did not find all of the proposed amino acids conserved.

The suspected binding site in the CRiSP PR1 domain (38) is less conserved and is expected to affect substrate specificity if the proteins would indeed expose enzymatic activity. Of the three sequence patches lining up to form the postulated binding site, the first is the most conserved. After four highly variable positions, the following sequence motif could be deduced: [R][T/V][V/V][G/K][V/I/L][G][E][N][F/Y][S] (square brackets enclose conserved, functional residues). It has to be noted that where more than one residue occurs in a certain position the second and third amino acids are present in one case only. The second sequence stretch accounting for the putative substrate binding site (Fig. 4, alignment positions 130–138) could also be put into a sequence motif (although this motif is less conserved than the first motif), [G/P/A]XX[P/Q]XX[M/V/K]X, with variants such as alanine in position 2 in the snake toxin representatives Trimersurus stejnegeri (P60623) and Pseudechis australis (Q84VA4), glutamine in position 5 in the helodermatid sequences, and at position 8 valine in *T. stejnegeri* and *P. australis* and lysine in *V. tristis*. The third patch of amino acids to participate in substrate binding is variable (alignment 179–185).

Five residues in the CRiSP CRD were brought forward as possible interaction partners with K+ channels (Fig. 4). In *T. stejnegeri*, these residues were Tyr150, Cys212, Thr213, Arg214, and Glu215 as well as the C-terminally located Phe244. Whereas Tyr150 was supposed to only interact with the ion channel, the remaining residues were thought to be involved in the actual blocking of this protein. In *H. suspectum*, position 125 was a histidine rather than a tyrosine, the latter of which was observed in all other proteins studied. Not surprisingly, the cysteine in position 212 is conserved throughout all sequences because cysteines are predominantly involved in maintaining the three-dimensional structure of the proteins. The three residues following this cysteine are not conserved in the cysteine active site of unknown function (38). A third conserved feature is the structurally important residues, despite the extensive amount of cysteines, postulated by Guo et al. (38); albeit, we did not find all of the proposed amino acids conserved.

Using our previously described methodology (18), molecular modeling studies revealed that several features are conserved in the CRiSPs investigated in this study (Fig. 4). Noteworthy are the conserved cysteine pattern and the tetrad in the PR1 domain, which was speculated to form part of an active site of unknown function (38). A third conserved feature is the structurally important residues, despite the extensive amount of cysteines, postulated by Guo et al. (38); albeit, we did not find all of the proposed amino acids conserved.

The suspected binding site in the CRiSP PR1 domain (38) is less conserved and is expected to affect substrate specificity if the proteins would indeed expose enzymatic activity. Of the three sequence patches lining up to form the postulated binding site, the first is the most conserved. After four highly variable positions, the following sequence motif could be deduced: [R][T/V][V/V][G/K][V/I/L][G][E][N][F/Y][S] (square brackets enclose conserved, functional residues). It has to be noted that where more than one residue occurs in a certain position the second and third amino acids are present in one case only. The second sequence stretch accounting for the putative substrate binding site (Fig. 4, alignment positions 130–138) could also be put into a sequence motif (although this motif is less conserved than the first motif), [G/P/A]XX[P/Q]XX[M/V/K]X, with variants such as alanine in position 2 in the snake toxin representatives Trimersurus stejnegeri (P60623) and Pseudechis australis (Q84VA4), glutamine in position 5 in the helodermatid sequences, and at position 8 valine in *T. stejnegeri* and *P. australis* and lysine in *V. tristis*. The third patch of amino acids to participate in substrate binding is variable (alignment 179–185).

Five residues in the CRiSP CRD were brought forward as possible interaction partners with K+ channels (Fig. 4). In *T. stejnegeri*, these residues were Tyr150, Cys212, Thr213, Arg214, and Glu215 as well as the C-terminally located Phe244. Whereas Tyr150 was supposed to only interact with the ion channel, the remaining residues were thought to be involved in the actual blocking of this protein. In *H. suspectum*, position 125 was a histidine rather than a tyrosine, the latter of which was observed in all other proteins studied. Not surprisingly, the cysteine in position 212 is conserved throughout all sequences because cysteines are predominantly involved in maintaining the three-dimensional structure of the proteins. The three residues following this cysteine are not conserved in the cysteine active site of unknown function (38). A third conserved feature is the structurally important residues, despite the extensive amount of cysteines, postulated by Guo et al. (38); albeit, we did not find all of the proposed amino acids conserved.

Using our previously described methodology (18), molecular modeling studies revealed that several features are conserved in the CRiSPs investigated in this study (Fig. 4). Noteworthy are the conserved cysteine pattern and the tetrad in the PR1 domain, which was speculated to form part of an active site of unknown function (38). A third conserved feature is the structurally important residues, despite the extensive amount of cysteines, postulated by Guo et al. (38); albeit, we did not find all of the proposed amino acids conserved.
V. varius and G. infernalis. The same applied for the residues in position Phe\(^244\) in T. stejnegeri, although all residues in this position contain a quite bulky side chain. There are two possible interpretations for these findings. 1) The residues under investigation are not involved in the blocking of ion channels, i.e., K\(^{+}\) channels. 2) Different CRiSPs interact with different ion channels, which contain different selectivity filters. The latter fact cannot be ruled out completely because it is known that e.g. the selectivity filters of ion channels may consist of both basic and acidic amino acids as well as uncharged amino acids (see Ref. 39). Thus, from the nature of amino acid modifications in this region as well, it can be inferred that, despite the high degree of overall sequence conservation between the CRiSP toxins of both helodermatid species as compared with CRiSPs from other anguimorph species, these toxins may interact with different ion channel isoforms.

Utilizing molecular modeling techniques as previously described by us (18), we also observed that the anguimorph CRiSPs contained three conserved surface histidines (His\(^82\), His\(^140\), and His\(^323\)). All three histidines are at least partially exposed to solvent, which is of particular interest with regard
to their side chain chemistry. This is of particular interest if one takes into account the pK\textsubscript{a} value of the \(\delta\)-NH in the imidazole side chain. The side chain pK\textsubscript{a} in free histidine is 6.02. However, this value may vary in protein environment. Because the venom pH is acidic (e.g. pH 5 for Bothrops jararaca (40)), it can be assumed that in venom most of these solvent-exposed histidines are protonated. When an anguimorph lizard (or snake for that matter) bites, venom is transferred into the bloodstream of the prey, and the venom proteins are thus exposed to a physiological pH of 7.3. At this pH, the surface histidine side chains are mostly deprotonated, which could affect the functionality and possibly the toxicity of CRiSPs after administration.

In addition to changes in functional residues, neofunctionalisation may also be facilitated through changes in tertiary
structure as a result of variations in the pattern and number of cysteines. Such changes were evident in CRiSP (Figure 4), kallikrein (Figure 5), epididymal secretory protein sequences (Figure 12), and ribonuclease (Figure 13). In both of the CRiSP variants with newly evolved cysteines (GU441473 from C. warreni, and GU441468 & GU441470 V. indicus), this results in an odd-number of cysteines, thus potentially resulting in dimerisation. Further, the V. indicus sequences have a truncation leading to the loss of six cysteines. In the novel anguid kallikrein toxins GU441496 from C. warreni and GU441502 from G. infernalis have deletions ancestral cysteines combined with the evolution of new cysteines. The variants recovered from V. acanthurus (Q2XXN1) and V. scalaris (GU441486) may also form a dimer as a result of a newly evolved cysteine resulting in an odd-number of cysteines. The venom epididymal secretory protein sequences recovered from the Varanus libraries have additional eight cysteines present. In addition, an additional mutation deleted four of the twenty cysteines. Such variation was present in two species (V. indicus and V. komodoensis), both of which also possessed an isoform that did not have this deletion, indicating that this mutation occurred prior to speciation from their common ancestor in one of the isoforms present at that time. The basal type ribonuclease toxin sequences (recovered from both C. warreni and G. infernalis) have deletions of two ancestral cysteines. The two isoforms of a derived variant recovered from the G. infernalis library have a frame-shift mutation near the C-terminal, resulting not only in slight truncated sequences and the loss of two of the ancestral cysteines, but also producing two new cysteines.

Timing of Toxin Recruitment Events and of Functional Diversification—Particularly notable was our identification of two additional toxin types (lectin and hyaluronidase) that share an ancient homology with toxins from snake venoms and thus form part of the core Toxicofera arsenal (Fig. 1). We were also able to determine the timing of diversification of the crucial tripeptide lectin toxin functional motif. Previously, based upon snake toxin studies, it was hypothesized that the EPN variation, which mediates antiplatelet activity through the utilization of the mannose binding site (41), was basal to all snake venoms, whereas the QPD variant, which utilizes the galactose binding site (41), was more recently derived within snake venoms (18). However, both motifs were recovered in this study (O. apodus and V. indicus, respectively) (Fig. 16). Therefore, as both motifs are basal to all toxicoferan venoms, it is unclear which is the ancestral state and which was an early derivation. Although hyaluronidase activity has been reported previously for Heloderma venom (42), lack of sequence data prevented determination of molecular identity and thus the relative relationship to the snake toxin forms, which themselves have only been recently sequenced (18, 43). Our sequencing of lizard venom hyaluronidase from both H. suspectum and G. infernalis allowed for the determination that not only was this a basal lizard venom toxin type but that it was indeed a toxin basal to all Toxicofera venoms (Fig. 1).

It had been previously unclear whether the exendin peptides were founded by a recruitment of a glucagon-like or VIP-like peptide (22). The relative presence of exendin peptides in Heloderma venoms was investigated through primer-directed sequencing with the aim to determine the relative timing of diversification between the ancestral glucagon-derived exendin-3 and -4, which are less toxic (22), and the more recently derived exendin-1 and -2, which contain the more potent VIP-like domain. It was shown that exendin-1 is present in only very low levels in H. horridum, whereas exendin-2 is absent entirely. Our results confirm a glucagon peptide ancestry of the exendin peptides. The molecular evolution of the VIP domain precedes the H. horridum/suspectum speciation event, but the duplication of exendin-1 to additionally produce exendin-2 postdates speciation (Fig. 1). The high levels of exendin-1 in H. suspectum but not H. horridum may be linked to the significant ecological niche occupation and size difference between the two species.

For the natriuretic peptides, we were able to elucidate the timing of domain mutations that additionally encode for the post-translationally liberated proline-rich helokinestatin peptides; the helokinestatins are a derivation that occurred basally to the anguid/helodermatid clade (Figs. 1, 2, and 7). We were also able to resolve the relationship to snake venom forms; lizard and snake toxin forms are in distinct clades separated by ordinary body forms and thus represent independent toxin recruitment events (Fig. 7). However, as with

---

**Table III**

| Toxin                          | Tissue type for normal secretion | Bioactivity of ancestral protein                                                                 |
|--------------------------------|----------------------------------|--------------------------------------------------------------------------------------------------|
| Celestinoxin                   | No match                         | Unknown                                                                                          |
| Cholecystokinin (cholecystokinin) | Expressed in brain, lung, testis, and throughout the length of the small intestine; in the brain, expressed predominantly in the optic tectum and brain stem | Myriad including induction of gastric secretion, stimulation of pancreatic secretion, increase of blood circulation and water secretion in the stomach and intestine, and stimulation of smooth muscle contraction |
| Epididymal secretory protein   | Epididymis                       | Unknown                                                                                          |
| Goannatyrasin (YY peptide)     | Intestine                        | Unknown                                                                                          |
| Ribonuclease                    | Expressed predominantly in the pancreas | Inhibits exocrine pancreatic secretion, has a vasoconstrictory action, and inhibits jejunal and colonic mobility |
the convergent recruitment of PLA2 genes for use as toxins in
snake and lizard venoms (13, 19, 44), different types of natri-
uretic body peptides were recruited for toxic mutation. The
lizard toxins are within the BNP clade, whereas the snake
toxins are within the CNP clade.

Phylogenetic structure exists within the snake toxins both in
the presence of derived proline-rich molecules and in the
presence of a derived C-terminal tail. Within the tail-less
snake venom isoforms, the gene has also been convergently
mutated to encode for additional, post-translationally
 cleaved proline-rich peptides within the viperid snakes: pit
viper natriuretic genes additionally encode for bradykinin-
potentiating peptides (see Ref.45), whereas the
Echis
genus of true vipers encodes metalloprotease-inhibiting pep-
tides (46). Although the snake venom natriuretic peptides
are founded by a tail-less CNP gene, some forms possess a
derived C-terminal tail and form a subclade within the main
tail-less clade and thus represent a recent derivation (Fig.
7). While the tail-less snake natriuretic toxins bind the gua-
nylyl cyclase B receptor, the derived forms with C-terminal
tails instead bind the guanylyl cyclase A receptor (16). Thus
the derived C-terminal tail possessing snake natriuretic tox-
ins are convergent both in tail and targeting with the C-
terminal tail possessing, guanylyl cyclase A binding lizard
natriuretic toxins.

Derivations of Venom System—We have shown previously
that in snake species that have developed secondary forms
of prey capture (e.g., constricting) or switched to feeding on eggs
the reptile venom system undergoes rapid degeneration char-
acterized by significant atrophying of the glands, reduction in
fang length, and accumulated deleterious mutations in the
genes encoding for the venom proteins (18, 47, 48). This is a
consequence of selection pressure against the bioenergetic
cost of protein production (49). In the lizard
V. komodoensis,
the robust glands and high venom yield thus argued for active use
of the venom system (20). Therefore, as the Anguimorpha
glands are not only retained but continue to diversify, this ar-
gues for active evolution operating under selection pressure.

Previous anatomical studies of
Heloderma
mandibular venom glands used traditional dissection and histological
methods and reported a single ducted gland for
H. horridum
and variably three to five compartments for the
H. suspectum
venom gland (1, 3, 50–53). By using MRI, here we found that
there instead are six compartments in both with each com-
partment with its own duct that terminates at the base of the
grooved teeth (Fig. 14, A and B). In contrast, although the
varanid lizards also have independently derived six compart-
ments, the compartment structures are distinct from those of
the helodermatids (compare Fig. 14, A and B, with Ref. 20).
and the varanid compartment ducts terminate between successive bladelike teeth as noted previously for V. komodoensis (20).

The architecture of the mandibular venom glands of the Anguimorpha shows two modes (Figs. 14 and 15): a mixed sero/mucous arrangement (anguids) or an encapsulated protein gland (helodermatids and varanids). In the Anguids, the gland has one compartment per tooth. Extensive intralumen drainage channeling was evident, but the major lumen is unstructured. In contrast are the independently derived encapsulated forms in Heloderma and in the Lanthanotus/Varanus clade. Segregation of the protein- and mucus-secreting regions and encapsulation of the protein glands occurred independently twice with the Heloderma and the Lanthanotus/Varanus arrangements representing independent derivations of the system (Figs. 1, 14, and 15). In both cases, posterior compartments were fused, thus increasing lumen storage space. Extensive intralumen drainage channeling in both cases feeds into highly structured lumens.

Although the adaptive significance and evolutionary polarity of the Anguimorh lizard venom gland variation are unclear, the glands are evidently more complex than reported previously (Fig. 14) and emphasize the notion that substantial evolutionary innovation of any aspect of the venom system can occur even among closely related species in this genus. To understand the evolutionary pressures responsible for the variation in gland structure, we need more information on the ecology, particularly feeding, on the one hand and on the function of the secretion epithelium and venom delivery on the other. Examination of variation in venom gland architecture and development across the group and beyond might provide further insights into the structure variability (increase or decrease in compartmentalization) and, together with differences in venom composition, explain the ecological and evolutionary significance of the entire venom apparatus. The retention of the ancestral maxillary and mandibular gland arrangement in O. apodus (Fig. 15) is particularly intriguing.

Conclusion—Our major findings are of a multidisciplinary nature with uses ranging from evolutionary biology to protein evolution and highlight the underutilization of lizard venom as bioresources in drug design and development. Collectively, these findings strongly support the notion that during the long evolutionary history of the 130 million-year-old Anguimorpha lizard radiation (54) there has been extensive modification of the venom system. All variables appear to change independently, ranging from the biochemical variation and specialization of the venoms to the glandular morphology. We still know little about the venom macroevolution and the potential role in the ecology/evolution of the animals themselves. We do not know enough of composition, the role of individual venom components in prey capture, or in vitro activities. Furthermore, the toxin types recovered in this study should not be considered as the full suite because the relatively limited sampling only recovered the dominant forms. More extensive sampling will no doubt recover novel isoforms of types identified to date as well as entirely new toxin classes. In addition, the relationships of transcriptomes to proteomes may reveal translational variances as has been documented for snakes (see Ref. 55).

However, we may safely assume that in common with snake venoms these lizard venoms possess a high potential for drug lead discovery as evidenced by the commercial success of an exendin-4 toxin analog (from Heloderma venom) in reaching the marketplace as the drug Byetta™ to treat type 2 diabetes. Underscoring their biodiscovery potential was our finding of three novel peptide types with potent and unique activities upon the cardiovascular system. Our structure–function studies provide data supporting the notion that anguimorph lizard venoms, although posing trivial human medical implications (with the exception of helodermatid species or large varanids such as V. komodoensis), provide interesting and unexploited templates for pharmacoeological studies. We would also like to emphasize that these species belong to an ever growing group of threatened species that may affect biodiversity and cause an important loss of genetic information. We therefore hope that this study will not only stimulate further research and provide a new understanding and realization of conservation efforts but will also encourage the utilization of this precious natural resource in a sustainable and ethical manner.

Acknowledgments—We express our deep gratitude to the naturalis Museum for allowing access to a L. borneensis specimen for MRI scanning. The California Academy of Sciences generously loaned us preserved specimens for magnetic resonance imaging studies; we are deeply grateful to Jens Vindum for facilitating this scientific exchange. We also thank Chip Cochran and Cale Morris for help in specimen collection.

REFERENCES

1. Kochva, E. (1974) Specialized glands of the lower jaw in anguimorphs, in Recherches Biologiques Contemporaines, Dedicated to the Memory of Dr. Manfred Gabe (1916–1973), L. Arvy (Ed.), Imprimerie Vagner, Nancy, France, 281–286
2. Kochva, E. (1978) Oral glands of the Reptilia, in Biology of the Reptilia (Gans, C., and Gans, K. A., eds) pp. 43–161, Academic Press, London
3. Saint Girons, H. (1988) The cephalic exocrine glands of the reptiles. I. Anatomical and histological data. Ann. Sci. Nat. Zool. 9, 221–255
4. Fry, B. G., Vidal, N., Norman, J. A., Vonk, F. J., Scheib, H., Ramjian, S. F., Kuruppu, S., Fung, K., Hedges, S. B., Richardson, M. K., Hodgson, W. C., Ignjatovic, V., Summerhayes, R., and Kochva, E. (2006) Early evolution of the venom system in lizards and snakes. Nature 439,
584–588
5. Vidal, N., and Hedges, S. B. (2005) The phylogeny of squamate reptiles (lizards, snakes, and amphisbaenians) inferred from nine nuclear protein-coding genes. C. R. Biol. 328, 1000–1008
6. Arano-Sánchez, D. (2008) Envenomation by a wild Guatemalan beaded lizard Heloderma horridum charlesbergi. Clin. Toxicol. 46, 897–899
7. Bock, D. D. (2005) Biology of Gila Monste: rs and Beaded Lizards, University of California Press, Berkeley, CA
8. Bogert, C. M., and de Campo, R. M. (1956) The Gila monster and its allies. The relationships, habits, and behavior of the lizards of the family Helodermatidae. Bull. Am. Mus. Nat. Hist. 109, 1–238
9. Bou-Abdou, C. F., and Kardassakis, D. G. (1988) Acute myocardial-infarction following a Gila monster (Heloderma suspectum cinctum) bite. West. J. Med. 148, 577–579
10. Cantrell, F. L. (2003) Envenomation by the Mexican beaded lizard: a case report. J. Toxicol. Clin. Toxicol. 41, 241–244
11. Hooker, K. R., Caravati, E. M., and Hartsell, S. C. (1994) Gila monster envenomation. Emerg. Med. 24, 731–735
12. Strimple, P. D., Tomassoni, A. J., Otten, E. J., and Banher, D. (1997) Report on envenomation by a Gila monster (Heloderma suspectum) with a discussion of venom apparatus, clinical findings, and treatment. Wildlife Environ. Med. 9, 147–170
13. Fry, B. G. (2005) From genome to “venome”: molecular origin and evolution of the snake venom proteome inferred from phylogenetic analysis of toxin sequences and related body proteins. Genome Res. 15, 403–420
14. Fry, B. G., Wüster, W., Kini, R. M., Brusic, V., Khan, A., Venkataraman, D., and Rooney, A. P. (2003) Molecular evolution and phylogeny of elapid snake venom three-finger toxins. J. Mol. Evol. 57, 110–129
15. Rodríguez de la Vega, R. C., Merino, E., Becerril, B., and Possani, L. D. (2003) Novel interactions between K+ channels and scorpion toxins. Trends Pharmacol. Sci. 24, 222–227
16. Fry, B. G., Wickramaratana, J. C., Lemme, S., Beuve, A., Garbers, D., Hodgson, W. C., and Aweeod, P. (2005) Novel natriuretic peptides from the venom of the inland taipan (Oxyuranus microlepidotus): isolation, characterization, and biological activity. Biochem. Biophys. Res. Commun. 325, 483–489
17. Mouhat, S., Jouirou, B., Mosbah, A., De Waard, M., and Sabatier, J. M. (2004) Diversity of folds in animal toxins acting on ion channels. Biochem. J. 378, 717–726
18. Fry, B. G., Scheib, H., van der Weerd, L., Young, B., McNaughtan, J., McHenry, C., Ferrara, T., Clausen, P., Scheib, H., Winter, K. L., Greisman, L. F., Nevalainen, T. J., Norman, J. A., Lewis, R. J., Norton, R. S., Renjifo, S. N., and Robertson, G. M. (2006) Modeling the pore structure of voltage-gated sodium channels in closed, open, and fast-inactivated states. Proc. Natl. Acad. Sci. U.S.A. 103, 10295–10299
19. Vouloudakis, I., Shi, Y., Nicolas, P., and Mor, A. (1996) Broad spectrum antibacterial activity of skin-PYY. FEBS Lett. 380, 237–240
20. Utsinsiharoen, P., Mackessy, S. P., Miller, R. A., and Tu, A. T. (1993) Complete primary structure and biochemical-properties of gilatoxin, a serine-protectase with kallikrein-like and angiotensin-degrading activities. J. Biol. Chem. 268, 21975–21983
21. Aminetzach, Y. T., Souli, J. R., Kong, C. Y., and Hoekstra, H. E. (2009) Convergent evolution of novel protein function in shrew and lizard venom. Curr. Biol. 19, 1101–1105
22. Miller, L. J., and Lybrand, T. P. (2002) Molecular basis of agonist binding to the type A cholecystokinin receptor. Pharmacol. Toxicol. 91, 282–285
23. Ondetti, M. A., Rubin, B., Engel, S. L., Plusec, J., and Sheehan, J. T. (1970) Cholecystokinin–pancreozymin—recent developments. Am. J. Dig. Dis. 15, 149–156
24. Guo, M., Teng, M., Niu, L., Liu, O., Huang, Q., and Hao, Q. (2005) Crystal structure of the cysteine-rich secretory protein stripes reveals that the cysteine-rich domain has a K+ channel inhibitor-like fold. J. Biol. Chem. 280, 12405–12412
25. Scheib, H., McClay, I., Guex, N., Clare, J. J., Blaney, F. E., Dale, T. J., Tate, S. N., and Robertson, G. M. (2006) Modeling the pore structure of voltage-gated sodium channels in closed, open, and fast-inactivated conformations reveals details of site 1 toxin and local anesthetic binding. J. Mol. Biol. Model. 12, 811–819
26. Sousa, J. R., Monteiro, R. Q., Castro, H. C., and Zingali, R. B. (2001) Proteolytic action of Bothrops jararaca venom upon its own constituents. Toxicon 39, 779–792
27. Drickamer, K. (1992) Engineering galactose-binding activity into a c-type mannose-binding protein. Nature 360, 183–186
28. Tu, A. T., and Hendon, R. R. (1983) Characterization of lizard venom hyaluronidase and evidence for its action as a spreading factor. Comp. Biochem. Physiol. B 76, 377–385
29. Harrison, R. A., Ibison, F., Willbram, D., and Wagstaff, S. C. (2007) Identification of cDNAs encoding viper venom hyaluronidases: cross-generic sequence conservation of full-length and unusually short variant transcripts. Gene 392, 22–33
30. Fry, B. G., and Wüster, W. (2004) Assembling an arsenal: Origin and evolution of the snake venom proteome inferred from phylogenetic analysis of toxin sequences. Mol. Biol. Evol. 21, 870–883
31. Soares, M. R., Oliveira-Carvalho, A. L., Wermeling, L. S., Zingali, R. B., Ho, P. L., Junqueira-de-Azevedo, I. L., and Diniz, M. R. (2005) Identification of novel bradykinin-potentiating peptides and C-type natriuretic peptide from Lachesis muta venom. Toxicon 46, 31–38
32. Wagstaff, S. C., Favreau, P., Chevalier, O., Laing, G. D., Wilkinson, M. C., Miller, R. L., Stöcklin, R., and Harrison, R. A. (2008) Molecular charac-
Diversification of Anguimorpha Lizard Venom System

Li, M., Fry, B. G., and Kini, R. M. (2005) Putting the brakes on snake venom evolution: the unique molecular evolutionary patterns of Aipysurus eydouxii (Marbled sea snake) phospholipase A2 toxins. *Biochem. Biophys. Res. Commun.* 365, 650–656

Morrissette, J., Elhayek, R., Possani, L., and Prestopino, G. (1994) The toxin helothermine affects potassium currents in newborn rat cerebellar granule cells. *J. Membr. Biol.* 139, 49–55

Ho, P. L., Soares, M. B., Yamane, T., and Raw, I. (1995) Reverse biology applied to *Micrurus corallinus*, a South American coral snake. *Toxicol. Toxin Rev.* 14, 327–337

Nikai, T., Imai, K., Sugihara, H., and Tu, A. T. (1988) Isolation and characterization of horridum toxin with arginine ester hydrolase activity from *Heloderma horridum* (beaded lizard) venom. *Toxicon* 26, 270–280