Sample Limited Characterization of a Novel Disulfide-Rich Venom Peptide Toxin from Terebrid Marine Snail *Terebra variegata*

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Abstract

Disulfide-rich peptide toxins found in the secretions of venomous organisms such as snakes, spiders, scorpions, leeches, and marine snails are highly efficient and effective tools for novel therapeutic drug development. Venom peptide toxins have been used extensively to characterize ion channels in the nervous system and platelet aggregation in haemostatic systems. A significant hurdle in characterizing disulfide-rich peptide toxins from venomous animals is obtaining significant quantities needed for sequence and structural analyses. Presented here is a strategy for the structural characterization of venom peptide toxins from sample limited (4 ng) specimens via direct mass spectrometry sequencing, chemical synthesis and NMR structure elucidation. Using this integrated approach, venom peptide Tv1 from *Terebra variegata* was discovered. Tv1 displays a unique fold not witnessed in prior snail neuropeptides. The novel structural features found for Tv1 suggest that the terebrid pool of peptide toxins may target different neuronal agents with varying specificities compared to previously characterized snail neuropeptides.

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Introduction

There is a need to discover new compounds that enhance the drug pipeline for treating ailments. The disulfide-rich protein toxins found in venomous organisms such as snakes, spiders, scorpions, leeches, and marine snails are highly efficient and effective for manipulating physiological pathways and for novel therapeutic and insecticidal drug development [1–4]. In an effort to identify novel disulfide compounds from venomous snails, we have investigated the Terebridae, a sister group to the more familiar cone snails. Novel peptide toxins from the Terebridae would enhance the pool of therapeutic compounds available for biomedical research. Venomous disulfide-rich compound includes ion channel and enzyme inhibitors, growth factors, and structural or ligand-binding domains [5]. The disulfide-rich peptide library produced by venomous organisms is estimated to contain well over 2 million compounds. The potential and promise of these compounds are evidenced by breakthroughs such as, angiotensin-converting enzyme (ACE)-inhibitor, Captopril [6] and more recently, ziconotide (Prialt), the first drug from a venomous marine snail, Conus magus, which is used to treat chronic pain in HIV and cancer patients [7]. Peptides, while currently not as bio-available as small molecule therapeutics, have advantages of higher target specificity and selectivity, and decreased toxicity. Despite their advantages, the use of peptides in drug development lags in the discovery and characterization stages. Obtaining workable quantities of venomous natural products has remained a significant challenge due to the large amount of material that has traditionally been needed to identify and structurally characterize peptidic compounds [8–12]. A sensitive and robust method for discovery and characterization of disulfide-rich peptides from sample limited venom sources would facilitate their application as biochemical tools and potential drug development targets [13–15]. Presented here is an integrated approach of liquid chromatography mass spectrometric sequencing, chemical peptide synthesis, and NMR for the rapid identification and characterization of sample limited bioactive disulfide-rich venom peptide toxins (Fig. 1). Structural characterization of *Terebra variegata* peptide toxin Tv1 was accomplished using a total venom sample of 4 ng, an amount that would previously make characterization of venom components almost impossible [16,17].

Tv1 is the first peptide structurally characterized from a terebrid snail. Terebrids are part of the Conoidean superfamily of predatory mollusks, which includes cone snails and turrids...
Tv1 is a novel twenty-one amino acid teretoxin peptide with a cysteine scaffold similar to the M-superfamily of cone snail neurotoxins, CC-C-C-C. Apart from the cysteine pattern, there is no sequence homology between Tv1, TR(I/L)CCGCYWNGSKDVCSQSCC, and known M superfamily conotoxins from Conus snails (Table 1). Conoidean venoms are a rich source of neuroactive disulfide-rich peptides, used to investigate cellular communication in the nervous system [17–22] and have diverse biomedical applications, including pain therapy [23,24]. In contrast to conotoxins, terebrid toxins, teretoxins, which were recently identified, [25,26] are an untapped resource for discovering novel neuropeptides. The sequence and structural characterization approach presented here significantly reduces the number of snails needed, while allowing for rapid characterization of venom components, in an effort to conserve natural resources (Fig. 1).

**Results and Discussion**

**Venom Extraction and De-Novo Sequencing of Tv1**

The primary sequence characterization of Tv1 was accomplished using charge-enhanced Electron Transfer Dissociation (ETD) in combination with traditional Collisional Activated Dissociation (CAD) on reduced and alkylated Terebra variegata venom [27]. The peptide toxins in the crude venom were reduced and subsequently alkylated using either iodoacetamide or N,N-dimethyl-2-chloro-ethylamine [27]. The latter converts every cysteine residue into a dimethyl lysine analogue, which increases the charge state of each cysteine-rich peptide toxin, resulting in excellent sequence coverage with ETD.

Combining complementary dissociation strategies CAD and ETD facilitates de novo sequencing, as each dissociation strategy has specific fragmentation characteristics, which can be used to confirm the corresponding ETD and CAD spectra (Fig. 1). This is advantageous as peptide toxin sequences can be obtained directly from crude venom without prior fractionation and using only mass spectrometry [27]. This feature is particularly important for sequencing peptide toxins from animals producing minimal amounts of venom, such as terebrids, where only nanogram quantities are extracted from each snail specimen. Four nanograms of crude Terebra variegata salivary gland contents were extracted with 30% acetonitrile/water in 0.1% trifluoroacetic acid. An aliquot of the crude venom was analyzed using LC-MS on an Orbitrap XL mass spectrometer to determine the number of teretoxins present. LC-MS analysis was repeated after reduction of the crude venom to determine the number of cysteine residues present in each component. At this point, Tv1 was selected for further analysis due to its similarity in molecular weight and number of cysteines to M superfamily conotoxins. A reduced version of Tv1 was dissociated using CAD (Fig. 1b, and Figure S1 in File S1) and the dimethyl lysine analogue version of the teretoxin was dissociated using ETD (Fig. 1b and Figure S2 in File S1). The ETD MS/MS spectrum resulted in almost 100% sequence coverage. The sequence assignment by ETD could be supported with the CAD spectrum on the reduced toxin (Fig. 1b and Figure S1 in File S1). There were two places of ambiguity in the Tv1 peptide sequence: (1) the inherent uncertainty of isoleucine (Ile) and leucine (Leu) at the third position, which cannot be differentiated with the mass spectrometer used for this analysis, and (2) the order of the first two amino acids, Thr, Arg or Arg, Thr. The second ambiguity was resolved based on the absence of the c1 ion, and the resulting Tv1 sequence was determined as TR(I/L)CCGCYWNGSKDVCSQSCC. However, the Ile/Leu ambiguity at position 3 could not be resolved as there was not sufficient crude extract remaining after MS characterization.

![Figure 1. Discovery and characterization of disulfide-rich Tv1 teretoxin. An integrated approach for characterizing sample limited disulfide peptidic natural products was applied: a) RP-HPLC on-line separation of Terebra variegata venom (highlighted is the region where the sequenced peptide eluted). b) ETD and CAD MS/MS analysis recorded on the native peptide to determine the sequence of Tv1 peptide. c) RP-HPLC of linear and folded versions of chemically synthesized Tv1 peptide. d) MS/MS spectrum of partially reduced and alkylated Tv1 used to determine the disulfide connectivity. e) NMR solution structure of folded Tv1 peptide. f) Bioactivity of 20 μM of Tv1 (−) versus normal saline (NS) (−) solution injected into polychaete worms. Y-axis indicates level of activity, where 3 is the normal activity of the NS injected worm, 1 and 2 indicate decreased activity due to partial paralysis. doi:10.1371/journal.pone.0094122.g001](https://www.plosone.org/doi/10.1371/journal.pone.0094122.g001)
Table 1. Comparison of Teretoxin Tv1 and known M superfamily Conotoxins.

| Teretoxin | Sequence | Target(s) | Reference |
|-----------|----------|-----------|-----------|
| Tv1       | TRICGCYWNQSKDVCSQSCC | unknown | This work |
| MtrIIIe   | VCCPFGGACHELCYCCD* | unknown | [23,33,49] |
| TxxIIa    | CCSSDVCDFHPSCTECG* | unknown | [50,51] |
| MtrIIIa   | GCSCGFGACFGCCVDCCV | unknown | [50] |
| TxxIIb    | CCPPVAICNMGCIKPC* | unknown | [31,50] |
| Reg12a    | GECCOOQWCGOCTDOCC | unknown | [52] |
| Tx3f      | RCCSDFPCIDCRCA* | unknown | [31] |
| Qc3.1     | ACCDPDWDGACGYSGCC | unknown | [23] |
| BullIIA   | VTCRCCGKRGECGRWCRDHSSRC* | Na1,2,1,3,1,4 | [53] |
| BullIIb   | VGERCCNKNGRGGRWCRDHSSRC* | Na1,2,1,3,1,4,1,5,1,7 | [53] |
| BullIC    | IVDRCCNNGKRGGRSRRWCRDHSSRC** | Na1,2,1,3,1,4,1,7 | [53] |
| MIIIA     | ZGCNHVPNPGSGRWCRDHACCC* | TTX-r | [54] |
| KIIIA     | CCGC5KWRHDSRC* | TTX-r | [55] |
| PI1IA     | ZRLCCGFOKSCSRSROQCKOHCC | Na1,1,4 | [38] |
| GI1A      | RDRC10KCKIDROQCKQOQEC* | Na1,2,1,4,1,6–1,8 | [56] |
| GI1B      | RDRC10KCKIDROQCKQOQEC* | Na1,2,1,4,1,6–1,8 | [56] |
| GI1C      | RDRC10KCKIDROQCKQOQEC* | Na1,2,1,4,1,6–1,8 | [56] |
| CnIIA     | GRCDVPNACGRWCRHACCC* | TTX-r | [54] |
| CnIIb     | ZGCCEGMPLLTRFWRQNNARCROQ | TTX-r | [54] |
| SmIIA     | ZRCNGRRGCSSWRCDHSRCC | TTX-r, TTX-s | [57] |
| SIIA      | ZNCNGGCCSSWKWRCDHARCC* | TTX-s, TTX-s | [58] |
| CIIA      | GRCEGPNQGSCSSRWCDDHARCC* | TTX-r | [59] |

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To confirm Tv1 de novo assignment, the peptide was chemically synthesized using Fmoc Solid Phase Peptide Synthesis (SPPS). Due to inherent uncertainty in the amino acid sequence at position 3, both Ile3 and Leu3 containing peptides were synthesized and subjected to the same chemical derivatization strategy. The two synthesized Tv1 teretoxins both resulted in an MS/MS spectrum identical to the native peptide, confirming the de novo sequence assignment in which TR is the order of the first two amino acids, but with remaining I/L uncertainty at position 3 due to the mass spectrometer used [Fig. 2 and Figure S2 in File S1]. The novel peptide teretoxin, Tv1 from venomous marine snail Teretia variigata was thus identified and its sequence was determined.

Synthesis, Characterization and Oxidative Folding of Tv1
Tv1 was chemically synthesized using Fmoc Solid Phase Peptide Synthesis without prior knowledge of disulfide connectivity, and all cysteine residues were protected with a standard side chain protecting group, trityl. The synthesized Tv1 peptide was purified to 99% using RP-HPLC (Figure S3 in File S1) and its mass was confirmed by MALDI mass spectrometry. The expected average mass of the linear peptide [M+H] 2316.3 m/z was observed (Figure S4 in File S1). Tv1 was folded using thiol-assisted air oxidation reaction and purified by HPLC (Figure S3 in File S1). The mass of folded Tv1, 2310.7 m/z, is reduced by 6 Da confirming that three intramolecular disulfide bonds had been formed (Figure S5 in File S1). No differences were observed in synthesis and folding kinetics of the two Tv1 peptides with Ile or Leu at the third amino acid position, and both versions of Tv1 were used to determine the disulfide connectivity of the peptide.

Disulfide Mapping of Tv1
Due to the limited number of enzymatic cleavage sites and the presence of adjacent cysteine residues, traditional strategies for disulfide mapping such as tryptic digestion and acid hydrolysis were difficult to implement. The disulfide connectivity of Tv1 was independently determined by MS/MS mapping and NMR spectroscopy. For MS/MS mapping, a partial reduction and dual alkylation protocol was applied using reducing agent TCEP-HCl (Tris(2-carboxyethyl)phosphine hydrochloride) and alkylating agents NEM (N-ethylmaleimide) and IAM (iodoacetamide) [28–30]. Chemically synthesized and oxidatively folded Tv1 peptide was first partially reduced at pH 3.0 with TCEP, and subsequently alkylated with NEM. The partially alkylated Tv1-NEM peptide was reduced further with TCEP at pH 7.0 and alkylated again with IAM this time. Dual NEM/IAM alkylation resulted in Tv1 peptide species that were labeled with 2, 4 or 6 NEM and IAM groups (Fig. 3 and Figure S6 in File S1). The location of NEM and IAM modifications in each of the six partially reduced species was determined by matching the MS/MS b- and y- series ions to theoretical patterns. All six partially reduced species provided complementary, non-contradictory information and the resulting connectivity was determined to be Cys4–Cys20, Cys5–Cys21 and Cys7–Cys16 (Fig. 3 and Table S1 in File S1). Tv1 has a cysteine scaffold similar to M Superfamily conotoxins (Table 1), but it displays a disulfide pattern previously unknown in native cone snail peptides [23,31–35] with a disulfide bridge formed between the two middle cysteines (Cys7 and Cys16) and a parallel 2-disulfide bond “staple” (Cys4–Cys20 and Cys5–Cys21) linking the N- and C-termini of the peptide.
Characterization of Tv1 by NMR Solution Structure

To confirm the disulfide bond connectivity derived from MS, the solution structure of Tv1 was derived using standard homonuclear proton NMR techniques on unlabeled folded synthetic peptide. Initial NMR experiments were carried out on Tv1 peptide containing either Ile or Leu at the third position. The spectra were very similar for both versions of the peptide, with the only differences found in the signals of the Ile3 and Leu3 residues themselves, indicating that both peptides adopt the same structure. Subsequent NMR structural calculations were carried out using only the Ile3 form of the peptide.

Proton assignments were obtained from 2D NOESY and TOCSY spectra, and carbon chemical shifts were assigned with the help of a natural-abundance $^{13}$C-HSQC spectrum (Table S2 in File S1). Initial structure calculation runs were carried out using only manually assigned Nuclear Overhauser Effect (NOE) distance restraints. Disulfide connectivities were then determined based on proximity of cysteine residues in the 10 lowest-energy structures and were in agreement with the disulfide bond pattern derived by MS analysis.

Direct evidence for the Cys7–Cys16 disulfide bond was also observed from NOE crosspeaks between the H$^\alpha$ of Cys16 and the H$^\beta$ of Cys7, as well as a number of connectivities to the two flanking residues, Gly6 and Tyr8 (Figure S7 in File S1). The disulfide linkages were determined, additional distance restraints for the disulfide bonds were introduced. The final round of structure calculations was run with covalent bonds between the connected cysteine side chains. A bundle of the 10 lowest-energy structures was obtained with an RMSD of 0.42 Å for backbone and 0.74 Å for all heavy atoms (Figure S8 and Table S3 in File S1). A query for other peptide structures similar to the determined Tv1 structure using the software packages VAST [36] or PDBeFOLD [37] did not yield any hits, indicating that the structure of Tv1 represents a new fold for small peptide toxins.

The most prominent and best-defined structural feature is a $\beta$-hairpin from Cys7 to Cys16 that is clamped together by the disulfide bond formed between these two residues (Fig. 1 and Figure S9 in File S1). The remainder of the peptide wraps around the side chain aromatic ring of Tyr8, in particular through the formation of hydrophobic interactions with the side chains of Ile3 and Val15. The N- and C-terminal loops are clamped together in an antiparallel way through the Cys4–Cys20, Cys5–Cys21 double-disulfide bond arrangement, giving the whole teretoxin the shape of a flattened ellipsoid (Figure S8 in File S1). Coordinates for the Tv1 structures have been submitted to the PDB and are available under accession code 2mix.

Although Tv1 has a cysteine scaffold similar to M Superfamily conotoxins (Table 1), the observed differences in fold and disulfide bonding pattern are not surprising given the lack of sequence homology to M superfamily conotoxins. It is interesting to note that the loop between C3 and C4 is at least 4 amino acid residues longer in Tv1 compared to any of the other listed conotoxins, which is in agreement with the fact that these are the residues involved in the formation of the $\beta$-hairpin that is lacking in conotoxins. No $\beta$-strand structures are found in any of the other M-superfamily conotoxins (Fig. 4) [28–32]. To our knowledge, the only other example of Tv1's disulfide bond pattern was found in an active non-native minor refolding product for $\mu$-conotoxin PIIIA [38]. The 3D structures of both the native and non-natively refolded forms of PIIIA are very different from the structure of
Tv1 as they contain segments of α-helix and no β-sheet structure. The difference between Tv1 and the PI-IIIa refolded product with the same disulfide bridge arrangement further supports the conclusion that Tv1 is representative of a new and unique group of peptide toxins.

Figure 3. Disulfide mapping of Tv1. a) RP-HPLC fractionation of all differentially folded peaks (on HPLC gradient of 20% to 35% buffer B (80% Acetonitrile, 0.1% TFA) in buffer A (water, 0.1% TFA) over 75 min. b) MS-MS spectrum of partially reduced peptide peaks showing representative alkylated b and y ions. c) Schematic representation of Tv1 with predicted disulfide bridges. doi:10.1371/journal.pone.0094122.g003

Figure 4. Structural comparison of Tv1 teretoxin and M-superfamily conotoxins. Comparison of the NMR structure of Tv1 with that of M superfamily conotoxins SmIIIA, mr3e, and KIIIA reveals significant structural differences between Tv1 and these conotoxins despite all having the same cysteine scaffold CC-C-C-CC. All structures are shown in cartoon representation with disulfide bonds highlighted in yellow. All figures were prepared using PyMol (www.pymol.org). Conotoxin structural references are as follows: MrIIIe [33] SmIIIA [61] and KIIIA [62]. doi:10.1371/journal.pone.0094122.g004
isms.
product peptide libraries produced by various venomous organ-
discovery and characterization of the vast disulfide-rich natural
integrated approach presented here can significantly enhance
mass spectrometry sequencing strategy. The sample limited
shell size [40], highlighting the need for an efficient microscale
venomous marine conoidean snails, are smaller than 8 mm in
an abundance of gastropods, including most species of the
and used as tools in neuroscience and as drug leads. In addition,
compared to disulfide-rich peptide toxins previously characterized
different mechanisms of action and different target specificities
structural features of Tv1 suggest that teretoxins will reveal
vestiges of the discovery and characterization of Tv1 allows for the
unequivocal analysis of minuscule amounts of sample. Four
nanograms of venom sample was used to elucidate the amino
acid sequence of Tv1. Such a minute amount of sample would
previously be inaccessible for natural product discovery from
venomous organisms. There are over 400 different species of
terebrids, each predicted to express between 50–200 different
peptide toxins in their venom. With an integrated approach of
mass spectrometry, chemical peptide synthesis, and NMR as
applied to Tv1, it will be possible to access the substantial quantity
of venom extract exhibited nAChR inhibitory activity [39]. Inhibi-
tion of nicotinic receptors at the neuromuscular junction can result
in paralysis. Further experiments are underway to determine the
specific molecular target of Tv1.

Conclusion
A novel teretoxin, Tv1, from Terebra variegata was structurally
identified and found to be biologically active. The strategy applied
to the discovery and characterization of Tv1 allows for the
unambiguous analysis of minuscule amounts of sample. Four
nanograms of venom sample was used to elucidate the amino
acid sequence of Tv1. Such a minute amount of sample would
previously be inaccessible for natural product discovery from
venomous organisms. There are over 400 different species of
terebrids, each predicted to express between 50–200 different
peptide toxins in their venom. With an integrated approach of
mass spectrometry, chemical peptide synthesis, and NMR as
applied to Tv1, it will be possible to access the substantial quantity
of disulfide-rich peptides (>80,000) in the Terebridae. The novel
structural features of Tv1 suggest that teretoxins will reveal
different mechanisms of action and different target specificities
compared to disulfide-rich peptide toxins previously characterized
and used as tools in neuroscience and as drug leads. In addition,
an abundance of gastropods, including most species of the
venomous marine conoidean snails, are smaller than 8 mm in
shell size [40], highlighting the need for an efficient microscale
mass spectrometry sequencing strategy. The sample limited
integrated approach presented here can significantly enhance
discovery and characterization of the vast disulfide-rich natural
product peptide libraries produced by various venomous organ-
isms.

Methods
Venom Extraction and De-Novo Sequencing
Four Terebra variegata snails were collected by MH from the Las
Perlas Archipelago in Panama on an expedition with the RV
URRACA and glands were dissected. Scientific research permits
were provided by the Smithsonian Tropical Research Institute
Permit Office (STRI-SPO) and The Panama Aquatic Resources
Authority (ARAP). Pooled crude venom contents from all 4
specimens were extracted with aid of light sonication in 30%
acetonitrile/water acidified with 0.1% trifluoroacetic acid and
lysine dimethyl analog versions of toxins) with 0.1 M acetic acid
and gradient eluted into the LTQ XL as described above. To
obtain higher quality MS/MS spectra in this second analysis, 12
columns as described above. The column was rinsed for 10 min
based on this initial screen optimal charge states for subsequent
targeted ETD analysis were chosen. For subsequent analysis either
reduced, iodoacetamide alkylated crude venom were pressure loaded onto a self-packed
PicoFrit column (New Objective) with integrated emitter tip as
described above and eluted using the same gradient as above into
the LTQ Orbitrap XL with ETD. Instrument was operated in a
data dependent mode cycling through a full scan (300–2,000 m/z, single scan) followed by 8 MS/MS
scans on the 4 most abundant ions from immediate preceding full scan. Next, aliquots of reduced and iodoacetamide
alkylated crude venom were pressure loaded onto a self-packed
PicoFrit column (New Objective) with integrated emitter tip as
described above and eluted using the same gradient as above into
the LTQ Orbitrap XL with ETD. Instrument was operated in a
data dependent mode cycling through a full scan (300–2,000 m/z, single scan) followed by 8 MS/MS scans on the 4 most abundant
ions from immediate preceding full scan. The instrument was
programmed to acquire first 4 CAD and then 4 ETD spectra on the 4 most abundant ions. The cations were isolated with a 2-Da
mass window and set on a dynamic exclusion list for 30 seconds
after they were first selected for MS/MS. Target value for MS/MS
was set to 10^4 cations and 10^6 anions of fluoranthene. For
CAD an activation time of 30 ms was chosen. For ETD the ion/
ion reaction time was set to 100 ms. In both cases 4 scans were
averaged per MS/MS spectrum. In a majority of cases, complete
sequence elucidation after this initial analysis was not possible, but
based on this initial screen optimal charge states for subsequent
targeted ETD analysis were chosen. For subsequent analysis either
reduced, iodoacetamide alkylated, or dimethyl lysine analog
versions of toxin were pressure loaded onto the self-packed
columns as described above. The column was rinsed for 10 min
(for reduced or iodoacetamide alkylated toxins) or for 20 min (for
lysine dimethyl analog versions of toxins) with 0.1 M acetic acid
and gradient eluted into the LTQ XL as described above. To
obtain higher quality MS/MS spectra in this second analysis, 12
scans were averaged and the data acquired in high resolution with
7,500 resolution setting. The spectra were manually interpreted
(Figures S1 and S2 in File S1).

Synthesis and Purification of Tv1
Tv1 peptide was synthesized by microwave assisted Fmoc SPPS
on a CEM Liberty synthesizer using standard side chain protection.
Standard Fmoc cystine with side chain protecting group Trityl (Fmoc-Cys (Trt)-OH) was used for all cysteine
residues in the Tv1 peptide. Following treatment of peptide resin with Reagent K [92.5% TFA (Trifluoroacetic acid), 2.5% TIS

Table 2. Tv1 bioactivity in polychaete worms.

| Activity | % Time spent |
|----------|--------------|
| Tv1 (20 μM) | NSS |
| at 0 | 0.995 | 1.45 |
| Between 0 and 1 | 74.47 | 0 |
| Between 1 and 2 | 19.98 | 0 |
| Between 2 and 3 | 4.55 | 98.55 |

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Oxidative Folding of Tv1

A one-step thiol-assisted oxidation was used to prepare folded Tv1 peptide. The linear peptide (29 μM) was incubated in 0.1 M Tris – HCl, 0.1 M NaCl, 100 μM EDTA, 1 mM GSH, 1 mM GSSG, pH 7.5. The folding reaction was terminated by acidification with 8% formic acid at 15 min, 30 min, 1, 2, 3, 4, and 24 h and the folding yield monitored using UHPLC. A preparative scale folding reaction was then conducted at an optimized time of 2 h, and the folded peptide was purified using X-Bridge semipreparative column (as mentioned earlier). Elution was carried out at 5 mL/min with 15% buffer B and 85% buffer A for the first 5 min, then increasing buffer B to 35% in 45 min. The purity was confirmed using Acquity UHPLC column (Figure S3 in File S1), and the molecular mass of the oxidized peptide was confirmed by MALDI-TOF (Waters, Micromass, CH3CN matrix) (Figure S4 in File S1).

Disulfide Mapping by Tandem MS/MS

A partial reduction and dual alkylation strategy was employed using reducing agent TCEP (Tris carboxy ethyl-phosphine hydrochloride, Thermo-Fisher) and alkylating agents NEM (N-ethylmaleimide, Sigma) and IAM (iodoacetamide, Sigma). Conditions for the partial reduction and alkylation of folded peptides were optimized modifying previous protocols [30,41]. Briefly, 9 μl of folded peptide (1 mM) was dissolved in 0.1 M citrate buffer, pH 3.0 and 1 μl of TCEP (20 mM in 0.1 M citrate buffer, pH 3.0) was added to it. Reduction was carried out at 4°C for 90 min and was terminated by injecting onto an RP-HPLC column [X-Bridge analytical column (5×150 mm, 5 μm particle size, Waters Corporation, Milford, MA, USA)] and eluted with 20% buffer B in buffer A for 5 min, then increasing buffer B to 35% in 75 min. Six partially reduced peaks (Figure S6 in File S1) were each collected directly into microcentrifuge tubes containing 100 μl NEM (200 mM in 0.1 M citrate buffer pH 3) and incubated at 37°C for 2 h in the dark. Partially reduced and alkylated reactions were then lyophilized and desalted to remove unreacted NEM using UHPLC. The desalted fractions were incubated with 100 nM TCEP at 55°C for 1 h to completely reduce the NEM labeled peptide and alkylated with 275 μM IAM for 2 h at 37°C in dark. The alkylation steps resulted in peptide species that were labeled with 2, 4 or 6 NEM or IAM groups. The labeled peptide was purified using X-Bridge semipreparative column (as mentioned earlier). Elution was carried out at 5 mL/min with 30% buffer B (80% Acetonitrile, 0.1% TFA) and 70% buffer A (0.1% TFA) for the first 5 min, then increasing buffer B to 35% in 45 min. HPLC buffer composition remained the same all through the work. Purity of collected peptide was confirmed by RP-UHPLC using an Acquity UPLC (BEH 300 C18 1.7 μm, Waters Corporation) column and eluted using a linear gradient from 0% to 75% buffer B in 3.5 min (Figure S3 in File S1). The identity of synthesized peptide was confirmed by molecular mass measurement of purified peptide using MALDI-TOF (Waters, Micromass, CH3CN matrix) (Figure S4 in File S1).

NMR Solution Structure Derivation

Samples for Nuclear Magnetic Resonance (NMR) studies were prepared by dissolving lyophilized oxidized peptide into either a 9:1 ratio of H2O:D2O or in 100% D2O at a concentration of ~260 μM. Sample pH was adjusted to pH 6.0 and 50 μM trimethylsilyl propanoic acid was added as a reference compound. Spectral acquisition was carried out at 5°C on a Varian Inova 600 MHz NMR spectrometer equipped with a cryogenically cooled HCN probe. 2D-TOCSY spectra were acquired in 90% H2O (64 scans with 1024×512 points) and D2O (48 scans with 1024×512 points) with spin-lock times of 60 ms, whereas 2D-NOESY spectra were acquired with mixing times of 200 ms in H2O with a matrix size of 1024×512 points in 64 scans and in D2O with a similar matrix size in 112 scans. A natural abundance 1H1C HSQC correlation spectrum was also acquired in 1024 scans with a matrix size of 1024×80 points. Acquired spectra were processed with NMRPipe [42] and analyzed with CCPNmr Analysis [43]. Proton assignments were made by comparison of TOCSY and NOESY spectra, and used to assign the 13C chemical shifts through the HSQC spectrum (Table S2 in File S1). Distance restraints for structural calculations were obtained from 2 NOESY spectra: one spectrum collected in 90% H2O with a mixing time of 200 ms and another spectrum collected in D2O with a mixing time of 400 ms. Manual assignment was performed for about 90% of all NOE cross-peaks. Dihedral angle restraints were derived from the assigned chemical shifts using the software TALOS+ [44]. Structural calculations were carried out with ARIA2 [45]/CNS [46]. Initial structural calculations were conducted using only the manual NOE assignments (with distances automatically calibrated by the software based on peak volumes) and without any restraints for the disulfide bonds. An ARIA2 run consisting of 8 iterations of simulated annealing followed by explicit water refinement of the 10 lowest-energy structures was performed. Disulfide connectivities were determined based on this initial structural bundle and included in the final round of structural calculations as covalent bonds. Eight iterations were performed in ARIA with 20 structures generated per iteration and the 7 lowest energy structures of each iteration forming the structural ensemble for the next iteration. For this run, NOE assignments were allowed to be automatically adjusted by the software during the calculation. For the last iteration, 100 structures were generated, and ten lowest-energy solutions were subjected to explicit solvent refinement. The quality of the final structural bundle was assessed by Procheck NMR (Table S3 in File S1) [47].

Bioactivity Assay on Polychaete Worms

Nereis virens (polychaete worms) (<3.5 g) were maintained at 4°C in salt water prior to injection. Worms were laid on a Styrofoam block and injected with an insulin syringe. A modified needle cap was used to maintain a consistent 1.5 mm depth of needle puncture. Worms were injected between the 5th and 7th segments on the ventral anterior end with the aim of targeting the ventral nerve cord. Control worms were injected with 12 μl of 0.9% normal saline solution (NSS). Lyophilized samples of Tv1 toxin were resuspended in NSS and diluted to a concentration of 20 μM which was confirmed using a ThermoScientific Nanodrop 2000c spectrophotometer. At least three worms were used for each trial-one saline control and two injected with Tv1 toxin for a total of 3 trials. After injection, the worms were transferred to side-by-side containers filled with room temperature salt water. The worm movements were then recorded by video in order to evaluate the effects of the toxin over a period of at least 3.5 hours. Varying temperature elicits different phenotypic responses from N. virens.
water at 4°C has an anesthetizing effect, while room temperature prompts a vigorous response [40].

Taking the worms from 4°C water and immersing them in room temperature water (25°C) immediately after injection stimulates an active response from the worms, making it easier to observe the paralysis (or partial paralysis) effects induced by a novel toxin in the first minutes after injection. Worms were observed every other day for two weeks after injection. The recorded videos were analyzed blind by two independent observers; the activity of each worm over a small time interval was noted on a scale of 1–3, with 3 being extremely active and 1 being inactive. Each video was analyzed and annotated three times by each observer. Observations for each worm began at the time of injection (Table S4 in File S1). The numerical results of the two independent viewings were averaged together if results were similar, if two viewers disagreed, a third observer was employed to review the disputed section of video. Results of relative activity vs. time were plotted to quantify the phenotypic effects of the toxin and saline injection on N. virens (Fig. 1f). The data was also used to calculate what percentage of time each worm spent inactive (a rating of 1), active (an averaged rating of 1.5–2), and extremely active (rating 2.5–3) (Table 2).

Supporting Information

File S1 This file contains Figure S1–Figure S9 and Table S1–Table S4. Figure S1, CAD of native Tv1. MS/MS spectrum recorded on a [M+2H]2+ ion after reduction of cysteine residues. The sequence is given above the spectrum and observed b, a and y-type fragment ions are labeled in the spectrum. Observed peptide backbone cleavage is indicated in the sequence above with 1 and 2 for N- and C-terminal fragment ions, respectively. Doubly charged fragment ions are labeled with +2. The neutral loss of water from the precursor ion is shown as [M+2H]2+–H2O, but neutral losses of fragment ions are not labeled. The spectrum was recorded at a resolution of 7500 at m/z 400 and all fragment ions have a mass accuracy of better than 5 ppm. Figure S2, ETD of native (black) and synthetic Tv1 (blue), MS/MS spectrum recorded on a [M+6H]6+ ion after conversion of cysteine residues to dimethyl lysine analogs. The sequence is given above the spectrum and observed c and z-type fragment ions are indicated in the sequence with 1 and 2, respectively. Doubly charged fragment ions of type c and z are labeled with +2, triply charged ions are of type c and z are indicated with *, z-type fragment ions that resulted from cleavage at cysteine with subsequent loss of the cysteine side chain are denoted in italic and charge reduced species are labeled in the spectrum with #. The spectrum was recorded at a resolution of 7500 at m/z 400 and all fragment ions have a mass accuracy of better than 5 ppm. Figure S3, RP-UHPLC chromatograms of Tv1 linear and oxidized peptide at 214nm. During a pilot folding reaction, over 90% of the linear Tv1 peptide fully oxidized and showed a peak at 1.36 minute in comparison to 1.83 minute linear Tv1 peak at the gradient of 0–75% buffer B (30% acetonitrile, 0.1% TFA) in buffer A (0.1% TFA) within two hours. Figure S4, Analysis of linear Tv1 peptide by MALDI-TOF mass spectrometry. MALDI-TOF spectrum of Tv1 peptide using z-Cyano-4-hydroxycinnamic acid matrix. Figure S5, Analysis of oxidized Tv1 peptide by MALDI-TOF mass spectrometry, MALDI-TOF spectrum of Tv1 peptide using z-Cyano-4-hydroxycinnamic acid matrix. Figure S6, RP-UHPLC analysis of partially reduced Tv1 peptide. On a UPLC (BEH 300 C18 1.7 μm, Waters Corporation, Milford, MA, USA) column, a linear gradient of 0–75% buffer B (30% acetonitrile, 0.1% TFA) in buffer A (0.1% TFA) over 6 minutes and peaks were assigned by their degree of reduction. Labels indicate how many disulfides are present. Figure S7, NOE contacts confirming the C7–C16 disulfide bond. An overlay of the 1H2H fingerprint region of the NOESY (in black) and TOCSY (in blue) spectra shows NOE crosspeaks linking Cys 16 and Cys 7 as well as contacts in the residues flanking the C7–C16 disulfide bond (C7-S17, Y8-C16, G6-C16, G6-N18). Figure S8, Bundle of the 10 lowest energy structures of Tv1 after explicit water refinement. The 10 lowest energy structures are shown in stick representation, displaying the tight convergence found in the final structural bundle. Figure S9, Structure of Tv1. An overlay of a cartoon representation and a stick model of the lowest-energy structure of Tv1 shows the β-sheet character of the peptide and reveals the important role of the Tyr β side chain in the formation of a small hydrophobic core. Table S1, Predicted and observed of b and y ions of differentially alkylated peptides by auto and targeted MS/MS analysis. Table S2, Chemical shift assignments of Tv1 (in ppm). Table S3, Structural statistics for the final 10 models of Tv1. Table S4, Sample raw data of Tv1 bioactivity in polychaete worms. (DOC)

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Author Contributions

Conceived and designed the experiments: PA MH. Performed the experiments: PA AG MB BU VR JQ PM. Analyzed the data: PA AG MB BU JQ PM BC SP. Contributed reagents/materials/analysis tools: MH BU SP BC. Wrote the paper: PA MH BU SP BC.

References

1. Koh CY, Kini RM (2012) From snake venom toxins to therapeutics—cardiovascular examples. Toxicon 59: 497–506.
2. King GF, Hardy MC (2015) Spider-venom peptides: structure, pharmacology, and potential for control of insect pests. Annu Rev Entomol 59: 475–496.
3. Vetter I, Lewis RJ (2012) Therapeutic Potential of Cone Snail Venom Peptides (Conopeptides). Curr Top Med Chem 12: 1546–1552.
4. King GF (2011) Venoms as a platform for human drugs: translating toxins into therapeutics. Expert Opin Biol Ther 11: 1469–1484.
5. Check S, Krishna SS, Grishin NV (2006) Structural classification of small, disulfide-rich protein domains. J Mol Biol 359: 215–237.
6. Cushman DW, Oudejens MA (1991) History of the design of captothiol and related inhibitors of angiotensin converting enzyme. Hypertension 17: 589–592.
7. Miljanich GP (2004) Xiconotide: Neuronal Calcium Channel Blocker for Treating Severe Chronic Pain. Curr Med Chem 11: 3029–3040.
8. Prashanth JR, Lewis RJ, Dutertre S (2012) Towards an integrated venomics approach for accelerated conopeptide discovery. Toxicon 60: 470–477.
9. Dutertre S, Lewis RJ (2010) Use of venom peptides to probe ion channel structure and function. J Biol Chem 285: 13315–13320.
10. Favreau P, Laure Menin SM, Perret F, Cheneval O, Stocklin M, et al. (2006) Mass spectrometry strategies for venom mapping and peptide sequencing from crude venoms: Case applications with single arthropod species. Toxicon 47: 670–678.
11. Favreau P, Stocklin R (2009) Marine small venoms: use and trends in receptor and channel neuropharmacology. Curr Opinion Pharmacol 9: 594–601.
12. Escoubas P, Quinton L, Nicholson GM (2008) Venomics: unravelling the complexity of animal venoms with mass spectrometry. J Mass Spectrom 43: 279–295.
13. Vetter I, Davis JL, Rash LD, Anangi R, Mobli M, et al. (2011) Venomics: a new paradigm for natural products-based drug discovery. Amino Acids 40: 15–28.
14. Quinton L, Demure K, Dobson R, Gilles N, De Paute E (2007) New Method for Characterizing Highly Disulfide-Bridged Peptides in Complex Mixtures: Application to Toxin Identification from Crude research articles. J Proteome Res 6: 3216–3223.
15. Tao YL, Lu B, Cruz LJ, Yates JR III (2010) Proteomic analysis provides insights on venom processing in Conus textile. J Proteome Res 9: 2292–2301.
16. Bhattacharyya M, Gupta K, Gowd KH, Balaram P (2013) Rapid mass spectrometric determination of disulfide connectivity in peptides and proteins. Mol Biochem 9: 1340–1350.
17. Bhattacharya S, Kil Y, Ueberheide B, Chait BT, Tayo L, et al. (2012) Constrained de novo sequencing of conotoxins. J Proteome Res 11: 4191–4200.
18. Castelin M, Puilandre N, Kantor Y, Medica MV, Terryn Y, et al. (2012) Macroring of venom apparatus innovations in auger snails (Gastropoda; Conoidea; Terebridae). Mol Phylogenet Evol 64: 21–44.
19. Holford M, Puilandre N, Terryn Y, Cruaud C, Olivera B, et al. (2009) Evolution of the Toxoglossa venom apparatus as inferred by molecular phylogeny of the Terebridae. Mol Biol Evol 26: 15–25.
20. Olivera BM, Teichert RW (2007) Diversity of the neurotoxic Conus peptides. Mol Interv 7: 251–260.
21. Olivera BM (2006) Conus peptides: biodiversity-based discovery and exogenomics. J Biol Chem 281: 31173–31177.
22. Terrus H, Olivera BM (2009) Conus venoms: a rich source of novel ion channel-targeted peptides. Physiol Rev 89: 41–68.
23. Han Y-H, Wang Q, Jiang H, Liu L, Xiao C, et al. (2006) Characterization of novel M-superfamily conotoxins with new disulfide linkage. FEBS J 273: 4972–4982.
24. Balaj G (2008) Integrating the discovery pipeline for novel compounds targeting ion channels. Curr Opin Chem Biol 12: 441–447.
25. Imperial JS, Watkins M, Chen P, Hillyard DR, Cruz LJ, et al. (2003) The Phylogeny, Foregut Anatomy and Comparative Toxinology. 308B: 744–756.
26. Imperial JS, Kantor Y, Watkins M, Ei FMH, Stevenson B, et al. (2007) Venomous Auger Snail Hastula (Impages ) hectica (Linnaeus, 1758): Molecular Phylogeny, Foregut Anatomy and Comparative Toxicology. 308B: 744–756.
27. Ueberheide BM, Fenyo D, Alewood PF, Chait BT, (2009) Rapid sensitive
28. Nair SS, Nilsson CL, Emmett MR, Schaub TM, Gowd KH, et al. (2006) De Novo Sequencing and Disulfide Mapping of a Bromotryptophan-Containing Conotoxin by Fourier Transform Ion Cyclotron Resonance Mass Spectrometry. Anal Chem 78: 8082–8088.
29. Corpuz GP, Jacobsen RB, Jimenez EC, Watkins M, Walker C, et al. (2005) Determination of the M-conotoxin superfamily: characterization of novel peptides from molluscivorous Conus venoms. Biochemistry 44: 8176–8186.
30. Corpuz GP, Jacobsen RB, Jimenez EC, Watkins M, Walker C, et al. (2005) Definition of the M-conotoxin superfamily: characterization of novel peptides from molluscivorous Conus venoms. Biochemistry 44: 8176–8186.
31. Mora D, Franco A, Moller C, Mora D, Fields GB, et al. (2006) Hyperhydroxylation: anew strategy for neuronal targeting by venomous marine molluscs. In Muller E.G. Werner, editor. Progress in Molecular and Subcellular Biology. Springer: 83–99.
32. Bouchet P, Lozouet P, Sysoev AV (2009) An inordinate fondness for turrids. Zoomorphology. J Biol Chem 282: 30699–30706.
33. Wang Q, Jiang H, Han Y-H, Yuan D-D, Chi C-W (2008) Two different groups of signal sequence in M-superfamily conotoxins. Toxicon 51: 813–822.
34. Tietze AA, Tietze D, Ohlenenschläger O, Leipold E, Ullrich F, et al. (2012) Mol Interv 7: 251–260.
35. Bru¨nger AT, Adams PD, Wortz W, Delaware WL, Gros P, et al. (1998) Structure: A hybrid method for predicting protein backbone torsion angles from NMR chemical shifts. J Biol Chem NMR 44: 767–768.
36. Mora D, Franco A, Pisarewicz K, Moller C, Fields GB, et al. (2006) Structure and functional diversities among mu-conotoxins targeting TTX-resistant sodium channels. Biochemistry 45: 3723–3732.
37. Tietze AA, Tietze D, Ohlenenschläger O, Leipold E, Ullrich F, et al. (2012) Multidimensional spectral processing system based on UNIX pipes. J Biol Chem 287: 23026–23032.
38. Bouchet P, Lozouet P, Sysoev AV (2009) An inordinate fondness for turrids. Zoomorphology. J Biol Chem 282: 30699–30706.