Jingzhaotoxin-I, a novel spider neurotoxin preferentially inhibiting cardiac sodium channel inactivation

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Running title: JZTX-I, a cardiac sodium channel inactivation inhibitor

The amino acid sequence of jingzhaotoxin-I and the nucleic acid sequence of jingzhaotoxin-I cDNA have been accessed in Swiss Protein database under Swiss-Prot # P83974 and AJ854060, respectively. (http://srs6.ebi.ac.uk/srsbin/wgetz?-page+srsq2+- noSession)
SUMMARY

Jingzhaotoxin-I (JZTX-I), a 33-residue polypeptide, is derived from the Chinese tarantula Chilobrachys jingzhao venom, based on its ability to evidently increase the strength and the rate of vertebrate heart-beats. The toxin has three disulfide bonds with the linkage of I-IV, II-V, III-VI, which is a typical pattern found in inhibitor cystine knot molecules. Its cDNA, determined by rapid amplification of 3’ and 5’ cDNA ends, encoded 62-residue precursor with a small pro-region of 8 residues. Whole-cell configuration indicated that JZTX-I was a novel neurotoxin preferentially inhibiting cardiac sodium channel inactivation by binding to receptor site 3. Although JZTX-I also exhibits the interaction with channel isoforms expressing in mammalian and insect sensory neurons, its affinity for tetrodotoxin-resistant subtype in mammalian cardiac myocytes (IC$_{50}$ 31.6 nM) is about 30-fold higher than for tetrodotoxin-sensitive subtypes in latter tissues. Unaffecting outward delay-rectified potassium channels expressing in Xenopus Laevis oocytes and tetrodotoxin-resistant sodium channels in mammal sensory neurons, it hopefully represents a potent ligand to discriminate cardiac sodium channels from neuronal tetrodotoxin-resistant isoforms. Furthermore, different from any reported spider toxins, the toxin neither modifies the current-voltage relationships nor shifts the steady-state inactivation of sodium channels. JZTX-I therefore defines a new subclass of spider sodium channel toxins. JZTX-I is a $\alpha$-like toxin first reported from spider venoms. The result provides an important witness for a convergent functional evolution between spider and other animal venoms.
INTRODUCTION

Voltage-gated sodium channels (VGSCs\(^1\)) are integral plasma membrane proteins composed of a pore-forming \(\alpha\) subunit (260 kDa) associated with up to four auxiliary \(\beta\) subunits (21-23 kDa) (1, 2). VGSCs play a vital role in the initiation and propagation of exciting signals on most excitable tissues. Similar to the \textit{shaker}\,potassium channel, the three-dimensional structure of sodium channel is a bell-shaped molecule (3, 4). From mammals over ten mammalian subtypes (Na\(_{1.1}-1.9\), and Na\(_x\)) exhibiting relatively similar pharmacological properties in different expression systems have been identified and characterized. Sequence analysis demonstrates that they have been highly conserved during evolution. Furthermore, to be beneficial for catching their prey, the properties of sodium channels in many animals even have evolved geographically in a coevolutionary arms race with their toxic prey (5, 6).

Many polypeptides targeting VGSCs have been identified in many kinds of animal venoms, such as those from scorpions, spiders, snakes and marine animals. With different critical residues and distinct functional characterization, these naturally occurring toxins are proved to be important tools to distinguish the different subtypes and disclose the function-structure relationships of VGSCs. Over six neuronal receptor sites are suggested to elucidate such relationships (7). Based on the analysis of precursor organization and gene structure combined with a three-dimensional fold, Zhu \textit{et al.} suggested that inhibitor cystine knot (ICK) peptides from animals shared a common evolutionary origin (8). Compared with scorpion and snake toxins, spider peptides are shorter ones containing around 35 residues with three/four disulfide

\(^1\)The abbreviations used are:

VGSC (Na\(_x\)), voltage-gated sodium channel; K\(_v\), voltage-gated potassium channel; TTX, tetrodotoxin; TTX-R, TTX-resistant; TTX-S, TTX-sensitive; DRG, dorsal root ganglion; RACE, rapid amplification of cDNA ends; ICK, inhibitor cystine knot; HPLC, high pressure liquid chromatography; MALDI-TOF, matrix-assisted laser desorption/ionization-time of flight; TCEP, Tris (2-carboxyethyl) phosphine.
bonds. Despite distinct amino acid compositions, most spider peptides are also found to share a clear homological ICK fold determined structurally by NMR and homology modeling techniques (9). The majority of them, such as δ-atracotoxins (δ-ACTXs) and μ-agatoxins, share a common mode of inhibiting inactivation kinetics of sodium currents on vertebrate or/and insect VGSCs by binding to receptor site 3 (10-12). Functional convergence widely occurs among animal toxins with different origins during evolution (8, 14). Many significant evidences appear in scorpions, snakes and marine animals, but similar evidence still remains to be found in spiders. The remarkable functional difference between spider site-3 toxins and other animal toxins is that the former evidently depress current amplitudes while the latter enhance current amplitudes evoked under whole-cell patch-clamp recording (10-12). Spider site-3 toxins all shows no effects on tetrodotoxin-resistant (TTX-R) sodium channels, such as Na,1.5 and Na,1.8-1.9. New emerging ligands will play an important role in elucidating their subtle difference. Moreover, They are proved to be potential valuable pharmaceutics or insecticides (13).

Here we report the isolation, cDNA sequence determination, and functional characterization of a novel sodium channel toxin, Jingzhaotoxin-I (JZTX-I), from the venom of Chinese tarantula Chilobrachys jingzhao (Araneae: Theraphosidae: Chilobrachys) (15). The toxin is composed of 33 residues stabilized by three disulfide bridges (I-IV, II-V and III-VI) assigned by partial reduction, sequencing and multi-enzymatic digestion. JZTX-I shows no effect on neuronal TTX-R VGSCs and Kv1 channels, but it inhibits channel inactivation of neuronal TTX-S subtypes and cardiac TTX-R subtypes. To the best of our knowledge, JZTX-I is a α-like toxin first reported to date from spider venoms. It provides an important witness for studying the convergent and divergent functional evolution of animal venoms.

**EXPERIMENTAL PROCEDURES**

*Toxin purification and sequencing*
JZTX-I was fractionated from Chinese tarantula Chilobrachys jingzhao venom, using a combination of ion-exchange chromatography and reverse-phase high pressure liquid chromatography (HPLC) as previously described (13). Lyophilized venom (10 mg in 2 ml in distilled water) was applied to a Waters protein-Pak CM 8 H column (5 mm × 50 mm) initially equilibrated with 0.2 M sodium phosphate buffer, pH 6.25 (buffer A). Then the column was eluted with a linear gradient (see Fig. 1A) at a flow rate of 3.0 ml/min. The fraction of interest collected was applied to a Vydac C18 (C4) analytical reverse-phase HPLC column (218TP54, 4.6 × 250 mm) and eluted at a flow rate of 0.7 ml/min by a linear gradient of acetonitrile containing 0.1% v/v trifluoroacetic acid (see Fig. 1B). The molecular mass and purity of toxin were determined by matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry. The full amino acid sequence was obtained from a single sequencing run on an Applied Precise™ 491-A protein sequencer by automated Edman degradation.

Assignment of the disulfide bridges of JZTX-I
Native peptide (0.1 mg) was partially reduced in 20 µl citrate buffer (1 M, pH 3.0) containing 6 M guanidine-HCl and 0.05 M Tris (2-carboxyethyl) phosphine (TCEP) at 40 °C for 10 min. The fractions monitored at 280 nm were separated on a C18 reverse-phase HPLC column with a linear gradient elution (25-50% acetonitrile in 40 min). The masses of all fractions collected were determined by MALDI-TOF MS. The intermediate with free thiols were lyophilized and then alkalized by adding 100 µl of 0.5 M iodoacetamide (pH 8.3). The alkalized peptide was desalted by reverse-phase HPLC and then submitted to an Applied Biosystems Model 491 gas-phase sequencer. The Edman degradation was performed with a normal automatic cycle program. Concerning the protease digestion strategy, native JZTX-I (0.1 mg) was dissolved in 0.2 ml Tris-HCl (0.2 M, pH7.5) buffer containing trypsin (4 µg), chymotrypsin (4 µg) and V₈ protease (4 µg). The mixture was incubated at 37 °C for 16 h. Then, the
masses of enzymolytic products were analyzed using MALDI-TOF mass spectrometry.

Identification of JZTX-I cDNA

The full-length of JZTX-I cDNA was obtained using rapid amplification of cDNA ends (RACE) methods as described previously (27). First, according to the manufacture’s instruction, the total RNA was extracted from 0.1 g fresh venom glands of female spiders using TRIzol reagent kit. 5 µg RNA was taken to convert mRNA into cDNA using the Superscript II reverse transcriptase with an universal oligo(dT)-containing adapter primer (5’-GGCCACGCGTCGACTAGTAC (dT)17-3’). The cDNA was then used as template for PCR amplification in 3’-RACE. A degenerate primer 1 (5’-GC(A/T/C/G)AA(C/T)TT(T/C)GC(A/C/G/T)TG(T/C)AA (G/A)AT(A/C/T)-3’) was designed corresponding to the N-terminal residues (18ANFACKI24) of mature JZTX-I. The partial cDNA of mature toxin was amplified by PCR technique using primer 1. Second, based on the partial cDNA sequence of JZTX-I determined by 3’-RACE, the anti-sense primers were designed and synthesized for 5’-RACE as follows: the gene specific primer 2 (5’-GGCCTAAGGGCTCCAGATACA-3’). With the strategy described by the RACE kit supplier, the 5’-end cDNA of JZTX-I was amplified using its gene-specific primer 2. Amplified products in both 3’- and 5’-RACE were precipitated and cloned into the pGEM-T easy vector for sequencing. DNA sequencing was performed by Bioasia Inc.. Nucleic acid sequences were analyzed using the software of DNAclub (by Xiongfong Chen) and DNAmam (by Nynnon biosoft).

Cell preparation

Rat DRG neurons were acutely dissociated and maintained in a short-term primary culture according to the procedures adapted from ref. (16). Briefly, 30-day adult Sprague-Dawley rats of either sex, in adherence with protocols approved by the
Hunan Normal University Animal Care and Use Committee, were killed by decapitation without anesthetization, the dorsal root ganglia were removed quickly from the spinal cord, and then they were transferred into Dulbecco's modified eagle's medium (DMEM) containing trypsin (0.5 g/L, type III), collagenase (1.0 g/L, type IA) and DNase (0.1 g/L, type III) to incubate at 34 °C for 30 min. Trypsin inhibitor (1.5 g/L, type II-S) was used to terminate enzyme treatment. The DRG cells were transferred into 35 mm culture dishes (corning, Sigma) containing 95% DMEM, 5% newborn calf serum, hypoxanthine aminopterin thymidine supplement and penicillin-streptomycin, and then incubated in the CO2 incubator (5% CO2, 95% air, 37 °C) for 1-4 h before patch-clamp experiment.

Single ventricular cardiomyocytes were enzymatically dissociated from adult rats according to the procedures adapted from ref. (17). Briefly, Sprague-Dawley rats (about 250 g) of either sex were killed by decapitation without anesthetization, the heart was rapidly removed and rinsed in ice-cold Tyrode’s solution containing (in mM): NaCl 143.0, KCl 5.4, NaH2PO4 0.3, MgCl2 0.5, glucose 10.0, HEPES 5.0, CaCl2 1.8 at pH7.2. Then the heart was mounted on a Langendorff apparatus for retrograde perfusion via the aorta with non-recirculating Ca2+-free Tyrode’s solution bubbled at 37 °C by 95% O2 and 5% CO2. After 10 min, perfusate was switched to a Ca2+-free Tyrode’s solution supplemented with 0.3% collagenase IA and 0.7% BSA, and the hearts were perfused in a recirculated mode for 5 min. After the enzymic solution was replaced by KB buffer containing (in mM): L-glutamic 70.0, KCl 25.0, taurine 20.0, KH2PO4 10.0, MgCl2 3 EGTA 0.5, glucose 10.0, HEPES 10.0 at pH7.4, the partially digested hearts were cut, minced, and gently triturated with a pipette in the KB buffer at 37°C for 10 min. The single cells were obtained after undigested tissues filtered through 200-µm nylon mesh. All cells were used within 8 h of isolation.

Cotton bollworm central nerve ganglion neurons were acutely dissociated and maintained in a short-term primary culture as described previously (18). Briefly, 10
day-old cotton bollworms were killed in 75% alcohol and washed in saline, containing (in mM): NaCl 90, KCl 6, MgCl₂ 2, HEPES 10, D-glucose 140 at pH 6.6. After wiped off the enteron, the central nerve ganglia were removed quickly. Then the nerve ganglia torn were incubated at room temperature (20-25 °C) in enzyme solution for 20 min, containing (in mM) NaCl 90, KCl 6, HEPES 10, D-glucose 25, D-mannitol 115, trypsin 0.3%, collagenase (IV) 0.15% at 6.6. Their enzymatic digestion was stopped in 35 mm culture dishes containing TC-100 (Gibico) 45%, DMEM (Sigma) 45%, fetal bovine serum 10%, D-glucose 100 mM, glutamine 0.6 mM, glutathione and penicillin-streptomycin at pH 6.6. After the cells in the ganglia were gently dispersed using suction tube, they were incubated in CO₂ incubator (5% CO₂, 95% air, 28 °C) for 2-3 h before patch-clamp experiment.

Whole-cell recording.
Sodium currents were recorded from experimental cells using whole-cell patch clamp technique at room temperature (22-25 °C). Recording pipettes (2-3 µm diameter) were made from borosilicate glass capillary tubing and its resistances were 1.0-2.0 MΩ when filled with internal solution contained (in mM): CsF 135, NaCl 10, HEPES 5 at pH 7.0. The external bathing solution contained (in mM): NaCl 30, CsCl 5, D-glucose 25, MgCl₂ 1, CaCl₂ 1.8, HEPES 5, triethanolamine chloride 20, tetramethylammonium chloride 70 at pH 7.4. After establishing the whole-cell recording configuration, the resting potential was held at -80 mV for at least 4 min to allow adequate equilibration between the micropipette solution and the cell interior. Ionic currents were filtered at 10 kHz and sampled at 3 kHz on EPC-9/10 patch clamp amplifier (HEKA, Germany). The P/4 protocol was used to subtracted linear capacitive and leakage currents. Experimental data was acquired and analyzed by the program pulse+pulsefit8.0 (HEKA, Germany). The needed concentrations of toxin dissolved in external solution were applied onto the surface of experimental cells by low-pressure injection with a microinjector (IM-5B, Narishige). Concerning DRG
neurons containing TTX-S and TTX-R sodium channels, the cells with diameters of 30-40 µm were chosen for the experiments, for larger DRG cells (>30 µm) tended to express TTX-S VGSCs while smaller ones (<10 µm) tended to express TTX-R VGSCs (19). TTX-R currents were separated from total currents using 0.2 µM TTX blocking TTX-S channels completely.

**Evolutionary tree of spider sodium channel toxins.**

Phylogeny of spider sodium channel toxins was constructed as described (14). In our study, we focused on the spider toxins from the species in the Family Mygalomorphae. Two groups of well-known spider toxins (µ-agatoxin I-VI and δ-paluIT 1-4) were chosen as the typical represents from the family Araneomorphae in order to outline the phylogeny clearly (12, 20). Multiple sequences of spider toxins were edited using the Bioedit Sequence Alignment Editor software, and then aligned and refined manually using Clustal W1.8 program (21). A pairwise distance matrix was calculated on the basis of the proportions of different amino acids. The matrix was then used to construct trees by the neighbor-joining (NJ) method (MEGA2.1) (22). The reliability of branching patterns was assessed using 1000 bootstrap replications.

**RESULTS**

**Purification and sequence analysis of JZTX-I**

The fraction of interest (JZTX-I) was purified using a combination method of ion-exchange HPLC and reverse-phase HPLC (Fig. 1), based on its ability to evidently increase the strength and the rate of vertebrate heart-beats (figure not shown). The molecular mass of naturally occurring toxin was determined to be 3675.64 Da by MALDI-TOF mass spectrometry. Its full amino acid sequence was performed by N-terminal Edman degradation and found to be composed of 33 residues including six cysteines (Fig. 3A). Six cysteines were assayed to form three disulfide bridges by the molecular mass of alkylized sample, which increased (58 × 6)
Da. The calculated molecular mass (3675.4 Da) corresponding to the primary sequence was consistent with the measured mass, suggesting that the C-terminal residue (Pro$^{33}$) was not amidated. JZTX-I was a novel tarantula toxin exhibiting limited sequence similarity with any reported peptide but 50.0% with JZTX-III. Despite the significant sequence divergence, sequence alignment indicated that all six cysteines in JZTX-I were strictly conserved at similar positions in most spider peptides adopting a typical ICK fold, such as HWTX-I, ProTx-I and HNTX-I.

**Assignment of the disulfide bridges of JZTX-I**

Fig. 2A showed the typical reverse-phase HPLC separation of the partially reduced mixture of JZTX-I by TECP. As MALDI-TOF mass spectrometry analysis pointed out, only one intermediate was obtained and further resolved to contain one disulfide bridge in peak II, whereas peaks I and III represented intact peptide and completely reduced peptide, respectively, because their molecular masses increased 0 (peak I), 4 (peak II) and 6 Da (peak III) compared to that of native peptide, respectively. Then peak II was collected and alkalized immediately with iodoacetamide followed by further purification using reverse-phase HPLC. Molecular mass determination and sequencing indicated that the free thiols of the fraction of interest had been alkylated. The sequencing results showed that signals of Pth-CM-Cys signals were observed except at the 16$^{th}$ and 29$^{th}$ cycles, positively supporting that the remaining disulfide bridge was cross-linked by Cys$^{16}$-Cys$^{29}$. In Fig. 2B, when JZTX-I was exposed to multi-enzymes (trypsin, chramotrypsin and V$_8$ protease) in the buffering solution, a series of smaller enzymolytic fragments were produced and their molecular masses were measured by MALDI-TOF mass spectrometry. A following reverse-phase HPLC separation of the enzymolytic mixture could produce a 2176 Da fragment, which was sequenced to be A1-W6 (containing C2), G12-F20 (C16 and C17) and L28-W31 (C29) as described in the inset of Fig. 2B. So, two disulfide bridges should be paired among C2, C16, C17 and C29. Since a disulfide bridge between C16 and C29 had been
determined above, we concluded that the second was cross-linked between C2 and C17. Accordingly, by process of elimination, the third disulfide bond is between C9 and C22.

Therefore, the disulfide linkage of JZTX-I was homologous to that for ICK peptides from spider venoms, such as ProTxs, HWTX-IV and HNTX-I, as well as conotoxins (δ-, μ-, κ-, and ω-), although the positions of six cysteines were not conserved between them (13, 23-25).

Cloning and sequencing JZTX-I cDNA

The full-length cDNA sequence of JZTX-I was completed by overlaying two fragments resulting from 3’- and 5’-RACE. As shown in Fig. 3B, the oligonucleotide sequence of the cDNA was a 383-bp bond found to comprise 5’-untranslated region (UTR), open reading frame and 3’-UTR. The open reading frame encoded a 63-residue peptide corresponding to the JZTX-I precursor, which contained a signal peptide of 21 residues, a pro-peptide of 8 residues and a mature peptide of 33 residues. The prepro-regions, composed of signal peptide and pro-peptide, are a hydrophobic peptide common to all spider toxins. Furthermore, its homology is an important proof that spider toxins can be grouped into different superfamilies to analyze their evolutionary relationship. The prepro-peptide of JZTX-I precursor showed limited sequence identity with those of other reported precursors but 66.7% and 57.1% with JZTX-III and GsMTx-4 (16, 26), respectively. Different from most spider sodium channel toxins, JZTX-I had no extra amino acid residues G or G+R/K at its C-terminus, known to allow “post-modification” α-amidation at the C-terminal residue (27), also implying that the C-terminal residue of mature toxin is not amidated. A polyadenylation signal (AATAAA) emerged in the 3’-UTR at position 17 upstream of the poly(A).

Effects of JZTX-I on sodium currents.
TTX-S and TTX-R VGSCs co-express in adult rat DRG neurons (1), whereas only TTX-S type is distributed in cotton bollworm central nerve ganglion neurons (18). TTX-R VGSCs are the primary type in adult rat ventricular myocytes, although Maier et al. suggested that some brain TTX-S subtypes were situated in its transverse tubules (28). The sodium channels expressing in these tissues have been defined as different isoforms based on their divergent amino acid sequences. Generally, TTX-R sodium currents activated and inactivated more slowly than TTX-S types under whole-cell configuration. JZTX-I at 1 µM in the bath solution was resistant to control TTX-R sodium currents in rat DRG neurons (Fig. 4A, n=5), but sensitive to TTX-R currents in ventricular myocytes and TTX-S currents in sensory neurons (Fig. 4B-D). JZTX-I evidently inhibited channel inactivation with distinct affinity on three tested tissues without changing peak current amplitudes or the time of current peak. Similar result was detected after application of β-pompilidotoxin (β-PMTX), which was considered to selectively interfere with sodium channel inactivation process without affecting the activation (29). Then, the efficiency of the toxin was assayed by measuring the I_{5ms}/I_{peak} ratio which gave an estimate of the probability for the channel not to be inactivated after 5 ms. JZTX-I at 1 µM inhibited channel inactivation of TTX-S currents in rat DRG neurons by 51.2 ± 8.0% (Fig. 4B, n=20), whereas a equal inhibition of TTX-R currents was achieved in rat cardiac myocytes exposed to the toxin only at 50 nM (Fig. 4C, n=5). The toxin at 1 µM could slow the decay of TTX-S sodium currents in cotton bollworm neurons by 52.4 ± 6.8% (Fig. 4D, n=11). All the inhibitions above were in a time-dependant and concentrate-dependant manner, and their IC_{50} values were assessed to be 0.13 µM, 31.6 nM and 0.76 µM, respectively (Fig. 4E). Furthermore, we checked the ion component of the inward currents maintained at the ending of depolarizing pulse. Because they could be eliminated in both rat DRG and insect neurons by TTX (0.2 µM) completely in Fig. 4B (n=8), ion currents were still conducted through sodium channels. The fact demonstrated that JZTX-I didn’t change the ion selectivity of sodium channels.
Fig. 5 showed the families of superimposed sodium currents in rat DRG neurons, rat cardiac myocytes and cotton bollworm neurons. The control current-voltage relationships indicated that their profound properties corresponding to divergent sequences of α-subunit represented some subtle difference in the thresholds of initial activation and the reverse potentials. After about 2 min of JZTX-I application, re-examination revealed that inhibiting channel inactivation was detected on three tested tissues at the given depolarizing pulses between the threshold and reverse potential. Interestingly, different greatly from both spider excitatory toxins (e.g. δ-ACTXs) and depressant toxins (e.g. HWTX-IV), the novel toxin neither shifted the current-voltage relationships nor changed control peak currents amplitudes (11, 13).

Previous studies have suggested that parallel to inhibiting inactivation kinetics, spider toxins also shifted the steady-state inactivation of sodium channels to a more negative potential (11, 17). To further assess the profound actions of JZTX-I on sodium channels, we also investigated the effect of the tarantula toxin on steady-state inactivation by employing a standard two-pulse protocol detailed in the illustration of Fig. 6. Under control condition, the putative midpoint of steady-state inactivation was around –57.1 mV in adult rat DRG neurons and cardiac myocytes, whereas it was –39.1 mV in cotton bollworm neurons. Unexpectedly, JZTX-I treatment did not evidently shift the steady-state inactivation in three experimental tissues, although channel inactivation inhibition was remained at test pulses similar to other spider excitatory toxins (e.g. δ-ACTXs) (10, 11).

Effects of JZTX-I on voltage-gated potassium channels (VGPCs)

Spider venoms contain many potassium channel toxins with high affinity besides sodium channel toxins. Although they are known to be potent specific for their respective receptors, Middleton et al. reported ProTx I at < 1 µM, a sodium channel toxins from the tarantula *Therixopelma pruriens* venom, also partially blocked Kv1.2 channel expressed in oocytes (23). Here we further checked the toxin on three
different VGPC isoforms (Kv1.1-1.3 channel) expressed in *Xenopus Laevis* oocytes using the two-electrode voltage clamp technique as described previously (30). However, no effects were detected after application of JZTX-I at 1 µM (Fig. 7, n=4).

**DISCUSSION**

Sodium channel toxins from venomous animals have been shown to lead to new insecticides and pharmaceuticals. To elucidate their structure-function relationship enhances our understanding of the properties of sodium channel proteins. In this work, we have isolated and characterized a novel spider toxin, jingzhao toxin-I (JZTX-I), from the Chinese tarantula *C. jingzhao* venom (15). Containing a conserved disulfide connectivity (I-IV, II-V and III-VI) and conforming exactly to the ICK definition as described by Escoubas et al. (31), JZTX-I represents a typical inhibitor cystine knot (ICK) structure, although it exhibits significant sequence divergence with other ICK peptides (24, 32). JZTX-I is a preferential cardiac sodium channel neurotoxin (IC$_{50}$ 31.6 nM) unaffected by Kv1.1-1.3 channels and neuronal TTX-R VGSCs. Its potential modulating site is reasonably assumed to receptor site 3.

The general biochemical features of JZTX-I precursor read from the cDNA sequence are similar to most spider toxin precursors, which are predicted to comprise a signal peptide, an intervening pro-peptide and a mature peptide (8, 27). The structural organization suggests that JZTX-I should be matured through a post-translational cleavage during the course of secretion where the prepropeptide can be removed at an endoproteolytic site anterior to mature peptide. Different from JZTX-III precursor having an uncommon signal site (-X-Ser-) (16), JZTX-I precursor contains one site (-X-Arg-) common to most animal toxin precursors. Generally, the similarities of the prepropeptide sequences are important criteria for defining the superfamilies for naturally occurring toxins. The toxins in the same superfamily always share conserved regions in the prepro-region, particularly in the signal peptide. Since their prepro-peptides exhibit 51.7% sequence identity each other, JZTX-I should belong to
the same superfamily as JZTX-III in the tarantula C. jingzhao. Interestingly, further analysis indicated that the pro-region length in both JZTX-I (8 residues) and JZTX-III (5 residues) precursors are much smaller than the lengths (over 25 residues) in other cDNAs of spider toxins.

Animal toxins can produce a common evolutionary event in order to be beneficial for catching their prey and defending against their predators (34). Analysis of evolution not only exhibits the evolutionary trace of animal toxins but also can predict their profound functional characterizations. Here we exhibit the phylogeny of spider sodium channel toxins based on their amino acid sequences. As shown in Fig. 8, spider sodium channel toxins are divergent in two evolutionary routes. The tarantula toxins from the family Theraphosidae with large body size are clustered in the second branch and preferentially inhibit mammalian sodium channels, whereas the toxins from the family Hexathelidae with small body size are clustered into the first branch, suggesting that they derive from different ancestor peptides. Interestingly, our studies also suggested clearly that the ancestor peptides for Araneomorphae toxins (δ-paluITs and µ-agatoxins) are more closely related to Hexathelidae toxins than to Theraphosidae toxins (δ-ACTXs), for they are together clustered in the first branch. The putative suggestion is further supported by the evidence that the disulfide connectivity (I-V, II-VI, III-VII and IV-VIII) is popular in both of them (35), whereas the disulfide bridge arrangement (I-IV, II-V and III-VI) is conserved in tarantula toxins (13, 16, 24). JZTX-I from tarantula venom is also clustered in the second branch of the phylogeny. Under whole-cell configuration, the characterizations of JZTX-I on sodium channels are significantly different from those of other tarantula toxins, such as HNTXs and ProTxs, but similar to those of spider toxins from the species in Hexathelidae and Araneomorphae. The novel toxin evidently inhibited fast inactivation of TTX-R VGSCs expressed in cardiac myocytes and TTX-S VGSCs expressed in vertebrate and insect sensory neurons, and belong to the excitatory group (32). Intriguingly, unlike δ-ACTXs reducing peak current amplitudes or scorpion
α-mammal toxins increasing peak current amplitudes (11, 36), JZTX-I altered the properties of sodium channels in a similar manner to scorpion α-like toxins and β-PMTX (33, 36). Their treatment shows no any effect on peak current amplitudes, I-V curves or steady-state inactivation. JZTX-I is a α-like toxin reported firstly from spider venoms and can define a new subclass of spider excitatory toxins. It will be an important evidence witnessing the divergent and convergent evolution between spider toxins and other animal toxins.

In the excitatory group, JZTX-I is the unique agent reported to date to alter the properties of Na\textsubscript{v}1.5 channels. It seems that JZTX-I has the ability to modulate much more VGSC subtypes than other spider toxins. δ-ACTX-Hv1a from Australian funnel-web spider venom is the most famous spider toxins. It inhibits the inactivation of sodium channels in vertebrate and insect sensory neurons (10, 11). But like other spider excitatory toxins, no similar description has been reported on sodium channels expressed in cardiac myocytes and skeletal muscles. Therefore, JZTX-I hopefully represents a useful ligand to elucidate the common properties among the subtypes of VGSCs and even discriminate Na\textsubscript{v}1.5 from other two TTX-R VGSCs (Na\textsubscript{v}1.8-1.9). Then, what is the underlying mechanism for the selectivity. It is well known that receptor site 3 is a short peptide composed of 13-15 residues situated at the DIV S3-S4 extracellular loop of α-subunit protein. Channel determinants demonstrate that three charged residues (E1613, E1616 and K1617), especially E1613, play a key role in binding peptide toxins. Despite low sequence similarity and different three-dimensional foldings with AaHIII (site 3 toxin), three residues (K3, R5 and D15) in δ-ACTX-Hv1a are assumed to interact structurally with them, respectively (35). However, the key residues in depressant group are determined to be K27 and R29 located at the loop IV of HNTX-IV scaffold (32). Such different allocation in two groups of spider toxins may contribute to better match their respective receptor sites. Both K27 and R29 are missing in JZTX-I at the corresponding positions, but three charged residues (K8, E11 and K13) are also clustered in the putative structure of
JZTX-I similar to δ-ACTX-Hv1a (Fig.3A). Further analysis of receptor site 3 indicates that E1613 and K1617 in neuronal TTX-S VGSCs are also strictly conserved in cardiac isoforms (D1612 and K1616), but they are accordingly substituted with uncharged residues (E/D→A/T) or anionic residues (K→E) in neuronal TTX-R subtypes (Na,1.8 and Na,1.9) (see Table 1). Moreover, the acidity of E1613 (pI=4.6) is weaker than that of D1612 (pI=4.3). Rogers et al. reported that E1613D significantly increased the affinity of ATX from 76 to 13 nM of the EC50 (37). These facts may partially explain that JZTX-I has a 30-fold affinity with cardiac VGSCs than with neuronal TTX-S isoforms and is preferential to Na,1.5 among three identified TTX-R VGSCs. For D1612 in cardiac subtype also conserves in skeletal muscle subtype (see Table 1), JZTX-I must be sensitive to Na,1.4. The hypotheses has been supported by the indirect result that the toxin significantly strengthened the normal contractions of mouse diaphragm induced by direct electrical stimulus, in which d-tubocurarine at high concentration was used to block the neuromuscular transmission completely.

In summary, in the present study a novel sodium channel toxin has been isolated and characterized from the tarantula *Chilobrachys jingzhao*. The toxin composed of 33 residues exhibits very low sequence homology with other peptides. It will be a novel useful tool to disclose the structure-functional relationship of receptor site 3 on sodium channel isoforms. Moreover, JZTX-I is the first reported α-like toxin from spider venoms, and will contribute to our understanding of the interaction between spiders and other venomous animals during evolution.

**ACKNOWLEDGEMENTS**

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isolation of cotton bollworm central nerve ganglion neurons.

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FIGURE LEGEND

Fig. 1. Purification and identification of JZTX-I.

(A), Ion-exchange HPLC chromatograph of the crude venom. An asterisk indicated the fraction with the retention time of 47.3 min containing the component of interest. (B), Analytical reverse-phase HPLC profile of the interest fraction on C18/C4 column. The linear gradients of buffer B (NaCl or acetonitrile) were indicated with a dashed line in (A) and (C).

Fig. 2. Identification of the disulfide bridge pattern of JZTX-I.

(A), Partial reduction of JZTX-I by TCEP was fractionated by reverse-phase HPLC. Three chromatographic peaks contained intact peptide (labeled I), partial reduced intermediate (labeled II) and complete reduced peptide (labeled III), respectively. (B), MALDI-TOF mass spectrometry analysis of multi-enzymatic peptide by trypsin, V₈ protease and chymotrypsin. A fragment with the mass of 2176 Da was chosen to be sequenced as detailed in the inset, in which solid line indicated the disulfide bridge pair determined by TCEP, whereas dash lines were the undetermined disulfide bridge pairs.

Fig. 3. Amino acid sequence and cDNA sequence of JZTX-I.

(A), comparison of the amino acid sequences of seven sodium channel toxins from tarantula venoms. JZTX-I and JZTX-III were isolated from *Chilobrachys jingzhao* (15, 16), ProTx-I and ProTx-II were from *Thrixopelma pruriens* (23), HNTX-I and HNTX-IV were from *Ornithoctonus hainana* (24, 32), and HWTX-IV were from *O. Huwenia* (13). The toxins had an amidated C-terminus indicated with an asterisk. The identical residues are shaded in black. The conserved disulfide bridge pattern of these toxins is indicated under their sequences. (B), the oligo-nucleotide sequence of JZTX-I cDNA. The amino acid composition of the precursor reading from the cDNA is suggested below the nucleotide sequence. The potential endoproteolytic sites are
pointed out with down arrows. The sequence of mature peptide was underlined by solid line.

**Fig. 4. Effects of JZTX-I on TTX-S and TTX-R sodium currents.**

All current traces were elicited by a 50 ms depolarizing potential of −10 mV from a holding potential −80 mV. (A), (B), (C) and (D) represented the typical time-dependent inactivation inhibition of JZTX-I on TTX-R currents and TTX-S currents in rat DRG neurons, TTX-R currents in rat cardiac myocytes and TTX-S currents in cotton bollworm neurons, respectively. The remaining currents at the end of depolarizing pulse could be further blocked by 200 nM TTX completely. JZTX-I at different concentrations failed to change the amplitudes of peak currents recorded in three checked tissues. (E), The concentration-dependent inhibition of sodium channel inactivation by JZTX-I. Every point (mean ± S.E.) comes from 3-20 separated experimental cells. These points were fitted according to Boltzmann equation: Inhibition% = 100/[1+exp(C-IC_{50})k], in which IC_{50} was the concentration of JZTX-I at half-maximal inhibition, k was the slope factor, and C was the concentration of JZTX-I.

**Fig. 5. Effects of JZTX-I on the current-voltage relationships of sodium channels.**

Family of both TTX-S and TTX-R current traces were induced by 50 ms depolarizing steps to various potentials from a holding potential of −80 mV. Test pulses ranged from −80 mV to +50 mV. The I-V curves showed the relationships of between current traces before (control) and after adding JZTX-I at different concentrations in rat DRG neurons (A, B), rat cardiac myocytes (C) and cotton bollworm neurons (D), respectively. In (B), (C) and (D), I_{5ms} was shown as the current inactivated at 5 ms.

**Fig. 6. Effects of JZTX-I on the steady-state inactivation of sodium channels.**

In cotton bollworm neurons (A) and (B), sodium currents were induced by a 50 ms...
depolarizing potential of +20 mV for various prepulse potentials for 1 s ranging from –90 mV to 0 mV. In rat DRG neurons (C) and cardiac myocytes (D), sodium currents were induced by a 50 ms depolarizing potential of –10 mV from various prepulse potentials for 1 s which ranged from –130 mV to –30 mV with a 10 mV increment. The data dots shown as the ratio of $I_{\text{test}}$ to $I_{\text{max}}$ were fitted according to the Boltzmann equation: $I_{\text{test}}/I_{\text{max}}=1/(1+\exp((V-V_{1/2})/k))$, where $V$ was the prepulse potential, $V_{1/2}$ pointed out further by dash lines was the voltage at which $I$ was 0.5 $I_{\text{max}}$, and $k$ was the slop factor. After the treatment of JZTX-I at 1 µM, 50 nM and 1 µM, the $V_{1/2}$ values were shifted only by +2.0 mV (-59.0 mV → -57.0 mV), +2.8 mV (-55.3 mV → -58.1 mV) and -3.7 mV (-39.1 mV → -42.8 mV) in rat DRG neurons, cardiac myocytes and cotton bollworm neurons, respectively.

**Fig. 7. Effects of JZTX-I on VGPCs expressed in Xenopus Laevis oocytes.**

$K_{v}1.1$ (A), $K_{v}1.2$ (B) and $K_{v}1.3$ (C) current traces were evoked by a 500 ms depolarization to +10 mV from a holding potential of –90 mV. No evident changes of the currents were detected in the absence and presence of 1 µM JZTX-I.

**Fig. 8. Unrooted phylogenetic tree of spider sodium channel toxins.**

The numbers on the branches were the bootstrap percentages supporting a given partition. The main characteristic functions of these spider toxins were shown on the right.
### Table 1

**Analysis of amino acid sequences of receptor site 3 on different sodium channel subtypes.** The emerging positions of functional residues are shaded in grey.

<table>
<thead>
<tr>
<th>DIV-S3</th>
<th>Extracellular loop</th>
<th>DIV-S4</th>
</tr>
</thead>
<tbody>
<tr>
<td>(DRG neuron) TTX-S Na_v.1</td>
<td>SIVGMFLAELIE--KYFVSPTLFRVIRLAR</td>
<td></td>
</tr>
<tr>
<td>(skeletal muscle) Na_v.1.4</td>
<td>SIVGLALSDLIQ--KYFVSPTLFRVIRLAR</td>
<td></td>
</tr>
<tr>
<td>(cardiac myocyte) Na_v.1.5</td>
<td>SIVGTVLSDLIIQ--KYFFSPTRLFRVIRLAR</td>
<td></td>
</tr>
<tr>
<td>(DRG neuron) TTX-R Na_v.1.8</td>
<td>SIGSLIFSAILKSLVNYFSPTLFRVIRLAR</td>
<td></td>
</tr>
<tr>
<td>(DRG neuron) TTX-R Na_v.1.9</td>
<td>SIVSTMISTLENQEHIPFPPTLFRVIRLAR</td>
<td></td>
</tr>
</tbody>
</table>
Fig. 1
Fig. 2
Fig. 3
Fig. 4

A. DRG TTX-R currents

B. DRG TTX-S currents

C. Cardiac TTX-R currents

D. Insect TTX-S currents

E. Percent of \( I_{\text{to}} \) to \( I_{\text{peak}} \) vs. log[JZTX-I], M

- DRG neurons
- Cardiac myocytes
- Insect neurons
Fig. 5

(A) Rat DRG TTX-S currents

(B) Rat DRG neurons

(C) Rat cardiac myocytes

(D) Insect neurons

Fig. 5
Fig. 6
Fig. 7
Fig. 8

(Excitatory toxins)
Inhibiting channel activation kinetics

(Depressant toxins)
Inhibiting channel inactivation kinetics

δ-paluT2
µ-agatoxin-III
δ-paluT4
µ-agatoxin-II
µ-agatoxin-I
µ-agatoxin-VI
µ-agatoxin-IV
µ-agatoxin-V
δ-paluT3
δ-paluT1
Magi 4
δ-ACTX-Hv1b
δ-MSTX-Mb1a
δ-AcTx-Ar1
δ-AcTx-Hv1a
JZTX-I
JZTX-III
HNTX-I
HNTX-III
ProTx-II
ProTx-I
HWTX-IV
HNTX-IV
HNTX-V

0.1
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