Centipedes subdue giant prey by blocking KCNQ channels

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Centipedes can subdue giant prey by using venom, which is metabolically expensive to synthesize and thus used frugally through efficiently disrupting essential physiological systems. Here, we show that a centipede (Scolopendra subspinipes mutilans, ∼3 g) can subdue a mouse (∼45 g) within 30 s. We found that this observation is largely due to a peptide toxin in the venom, SsTx, and further established that SsTx blocks KCNQ potassium channels to exert the lethal toxicity. We also demonstrated that a KCNQ opener, retigabine, neutralizes the toxicity of a centipede’s venom. The study indicates that centipedes’ venom has evolved to simultaneously disrupt cardiovascular, respiratory, muscular, and nervous systems by targeting the broadly distributed KCNQ channels, thus providing a therapeutic strategy for centipede envenomation.

Predators usually catch prey that are smaller than themselves because capture success is a function of relative prey/predator size. It declines as the ratio increases (1, 2), but there is anecdotal evidence that centipedes subdue prey much larger than themselves by means of venoms (3, 4). Venomous predators use potent venom to immobilize large prey as well as defend against predators (5). However, venom is metabolically expensive to synthesize. For example, the metabolic rate of snakes increased by an average of 11% during a 3-d study over which venom regeneration was measured (6). Thus, venoms are used frugally by effectively targeting essential physiological systems, such as nervous, respiratory, circulatory, and muscular systems, to ensure successful capture of prey (2, 7).

Snake venoms target circulatory or nervous systems to effectively capture prey (7), while spiders, scorpions, and snails often immobilize prey by using venoms that act on the nervous system (8, 9). Although some insights may be gleaned from recent work on centipede toxins (10–13), the underlying mechanism of how centipedes capture giant prey remains unknown.

Centipede envenomations occur frequently. In Hawaii, centipede bites were responsible for 11% of the cases that presented to hospital emergency departments from 2007 to 2011 (14). Severe clinical symptoms, such as vasospasm, acute hypertension, myocardial ischemia, and even death, have been observed following centipede envenomation (15–18). These symptoms point to cardiovascular disorders, and based on our knowledge of scorpion and spider toxins, ion channels are the likely targets. We screened centipede venom against various ion channels and found that a peptide toxin, SsTx, is extremely potent on all members of the KCNQ family of potassium ion channels. KCNQ is a family of multifunctional K+ channels that are involved in a diverse array of physiological functions, such as cardiac action potential repolarization, coronary circulation and reactive hyperemia, and cerebral neuron excitation (19, 20). KCNQ channels are also found in airway smooth muscle cells from rodent and human bronchioles (21). Based on the clinical symptoms caused by centipede bites and the role of KCNQ in essential physiological systems, we tested the following two hypotheses: (i) centipede venom blocks KCNQ channels to disrupt multiple essential physiological systems, such as cardiovascular, respiratory, and nervous systems, and thus (ii) KCNQ openers may neutralize toxicities caused by centipede envenomation.

Results

Centipedes Rapidly Subdue Mice. To test if centipedes are capable of rapidly subduing giant prey, golden head centipedes (Scolopendra subspinipes mutilans) with an individual body weight of ∼3 g were kept with a mouse (∼45 g) to observe their interactions. As illustrated in Fig. 1 d and Movie S1, a single centipede immediately attacked the mouse and subdued it within 30 s. The LD\textsubscript{50} values of the centipede crude venom in mice were 130 and 93 mg/kg after i.m. and i.p. injections, respectively (Fig. 1 g). Capture of giant prey has been reported for social foragers such as ants (22), spiders (23), or pseudoscorpions (24), in which predatory behavior is largely due to a peptide toxin in the venom, SsTx, and further established that SsTx blocks KCNQ potassium channels to exert the lethal toxicity. We also demonstrated that a KCNQ opener, retigabine, neutralizes the toxicity of a centipede’s venom. The study indicates that centipedes’ venom has evolved to simultaneously disrupt cardiovascular, respiratory, muscular, and nervous systems by targeting the broadly distributed KCNQ channels, thus providing a therapeutic strategy for centipede envenomation.

Significance

The work showed that centipede venom can cause disorders in cardiovascular, respiratory, and nervous systems. The cardiovascular toxicity of the venom comes mostly from a peptide toxin SsTx, which blocks the KCNQ family of potassium channels. Retigabine, a KCNQ channel opener, neutralizes centipede venom toxicity, and thus could be used to treat centipede envenomation.

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Data deposition: The NMR chemical shifts have been deposited in the Protein Data Bank, www.wwpdb.org (PDB ID code 5X03), and GenBank database (accession no. MG585384).

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Venom has evolved to disrupt multiple essential physiological systems. The prey is subdued by several individuals. A single spider, Zodarion cyrenaicum, can capture a giant ant within 10–20 min (25). By contrast, the capture efficiency of the centipede is much higher than that of the spider. This is possible because the centipedes’ venom has evolved to disrupt multiple essential physiological systems.

**Isolation and Identification of KCNQ Inhibitor from Centipede Venom.** We tracked the toxin that causes the lethal activity by using peptide purification coupled with toxicity experiments (Fig. 1B). Diluted crude venom of golden head centipedes was subjected to Sephadex G-50 gel filtration. The protein fraction containing SsTx is marked as a red arrow. (Inset) Lethality rate of mice recorded after i.p. and i.m. injection of crude venom and then fitted to a Hill equation (n = 3 animals per group, n = 3 groups per point). (C) Isolation of native SsTx (red arrow) from the pooled protein fraction by a C18 RP-HPLC column. (D) MALDI-TOF analysis and sequencing of SsTx, indicated by the 20-to-46 and 24-to-48 assignments. (E) NMR structural model of SsTx, with the electrostatic potential distribution shown in color (red = negative, blue = positive) on the right. The negatively charged surface and the location of R12 and K13 are shown on the NMR structure of SsTx. (F) Dose–response curves displaying the SsTx inhibition of KCNQ1, 2, 4, and 5. Data were fit with a Hill equation. The IC50 and slope factor values are 2.8 ± 0.5 μM and 1.74 ± 0.04 (n = 5 cells) for KCNQ1, 2.7 ± 0.4 μM and 0.99 ± 0.06 (n = 5 cells) for KCNQ2, 2.5 ± 0.4 μM and 1.18 ± 0.03 (n = 5 cells) for KCNQ4, and 2.7 ± 0.5 μM and 1.12 ± 0.06 (n = 5 cells) for KCNQ5, respectively. (G) The inhibitory percentage of KCNQ4 currents in the presence of 10 μM SsTx, recorded with increases in test pulses from a resting membrane potential of −80 mV (n = 3). (H) The half-maximum response of SsTx in different K+ concentrations recorded from KCNQ4-expressing HEK293T cells (n = 5 cells per bar). Average values represent means ± SEMs.

and C). A small peptide toxin [named here SmSp Spooky Toxin (SsTx); GenBank accession no. MG853584], with a molecular weight of 6,017.5 Da, was subsequently identified from the centipede venom (Fig. 1D). The primary sequence of mature SsTx was obtained by Edman degradation (Fig. 1D), and the sequence shows no resemblance to any known animal toxins by a search of nonredundant protein databases, via the BLAST server (https://blast.ncbi.nlm.nih.gov/Blast.cgi). Comparing the purified SsTx with the premature one, we found that the 76-amino acid peptide undergoes posttranslational modification by cleaving off a signal peptide of 23 amino acids in length and yielding the mature peptide of 53 amino acids (Fig. S1A). Partial reduction and MS indicated that SsTx contains two intramolecular disulfide bridges: Cys20-Cys46 and Cys24-Cys48 (Fig. 1D and Fig. S1 B–D). SsTx structure was solved by solution NMR analysis (Fig. 1E, PDB 5X05). The structure is compact, with amino acids in position 12 (arginine) and 13 (lysine), which form a positively charged surface (Fig. 1E). The C-terminal of the toxin is relatively flexible.

We examined SsTx on TRPV1, TRPV2, BK Ca, hERG, voltage-gated sodium channels, voltage-gated potassium channels, and voltage-gated calcium channels (Fig. S1 E–I and O) and found that SsTx exhibits potent inhibitory effect on KCNQ4, with an IC50 of 2.5 ± 0.4 μM (Fig. S1 J and K). SsTx also inhibited other KCNQ channels with IC50 values of 2.8, 2.7, and 2.7 μM for the homotetrameric complexes of KCNQ1, KCNQ2, and KCNQ5, respectively (Fig. 1F and Fig. S1 L–P). Further analyses confirmed that SsTx blockade becomes progressively less when the membrane potential becomes more depolarized, and that higher K+ concentrations increase the IC50 values (Fig. 1 G and H), consistent with a model in which SsTx binds to the outer pore domain of KCNQ channels (26). Since sequences of the outer pores of members of the KCNQ family exhibit high similarity, SsTx can inhibit different KCNQ channels and thus affect multiple physiological processes.

**Direct Interaction Between SsTx and KCNQ Channels.** Given that KCNQ4 is important in the regulation of pulmonary vascular tone, and blocking KCNQ4 increases the isometric tension of interlobular arteries (27), we hypothesized that SsTx may contribute to the vasospasm by inhibiting the KCNQ4 channel as reported in clinical cases of centipede envenomation (17). We therefore first investigated the interaction between SsTx and KCNQ4. The onset of channel inhibition upon perfusion of 10 μM SsTx is fast, with a time constant of 0.086 ± 0.011 s, and the recovery of channel current after washout of the toxin is relatively slow, with a time constant of 0.265 ± 0.007 s (Fig. S2 A and B), which is consistent with an IC50 of 2.5 μM. Alanine substitution was performed to identify residues important for SsTx’s affinity on KCNQ4. Alanine substitution to six positively charged residues (K4A, K10A, K11A, R12A, K13A, and K45A) reduces the toxin’s inhibitory effect on KCNQ4 (Fig. 2 A and B). These mutant toxins preserved WT-like structural features (Fig. S2C), suggesting that the reduced inhibition is caused by a direct interaction between SsTx and KCNQ4. In particular, alanine substitution at positions 12 and 13 increased IC50 values to 104.7 and 117.5 μM, respectively, while that of the WT is 2.5 μM (Fig. 2C). These two residues in combination with other positively charged amino acids in the N-terminus form a positively charged surface in SsTx (Fig. 1E). Collectively, we concluded that this positively charged surface constitutes the bioactive binding surface for SsTx to target KCNQ channels. Consistent with this conclusion, conservative replacement of lysine with arginine (K13R) or vice versa (R12K) has minimal influence on SsTx inhibition of KCNQ channels (Fig. S2D).

Next, we explored the toxin-binding sites on KCNQ channels. Since toxin binding is reduced under depolarizing membrane potentials and under higher intracellular K+ concentration (Fig. 1 G and H), similar to the “knock off” effect reported by earlier
studies on charybdotoxin (26), we postulated that SsTx interacts with the pore domain of KCNQ channels. This hypothesis is consistent with SsTx's positively charged bioactive surface. We made point mutations to the residues located on the extracellular side outer-pore region as well as the extracellular loops of the KCNQ4 α subunit. We found that D266 in the turret and D288 in the P-loop region are crucial in SsTx-induced inhibition (Fig. 2D). In the presence of 10 μM SsTx, ~99 and 97% inhibitory activity was lost in KCNQ4_D266G and KCNQ4_D288A, respectively (Fig. 2E). As illustrated in Fig. 2F, the IC_{50} of SsTx on WT KCNQ4 is 2.5 μM, and the value becomes 108.2 ± 6.3 μM for KCNQ4_D266G and 218.8 μM for KCNQ4_D288A. In addition, mutations at both positions did not affect voltage activation of KCNQ4 (Fig. S2E), suggesting that these residues directly affect SsTx-KCNQ4 interaction. In a structural homology model of KCNQ4, D266 and D288 form a negatively charged surface near the outer-pore region that could interact with SsTx (Fig. S3B).

To further validate interactions between the positively charged residues on SsTx and the negatively charged residues of KCNQ4, we applied thermodynamic mutant cycle analysis (28–30) to experimentally investigate the molecular interactions between SsTx and KCNQ4 (Fig. 3A–C). The interaction between the residue pair is considered energetically significant when the calculated interaction Ln(Ω) is much larger than the other pairs (31). As illustrated in Fig. 3A–C, residue K13 from SsTx and D288 from KCNQ4 has a Ln(Ω) of 3.12, while the Ln(Ω) values are much smaller between residues K4, K10, K11, R12, or K45 on SsTx and residue D266 on KCNQ4. Similarly, residue R12 of SsTx and residue D288 of KCNQ4 have a Ln(Ω) value of 2.53, while the Ln(Ω) values for residues K4, K10, K11, K13, or K45 on SsTx and residue D288 on KCNQ4 are all much smaller (Fig. 3E and F and Fig. S2F). These results indicate that residues R12 and K13 on SsTx functionally interact with residues D288 and D266 on KCNQ4, respectively.

We would like to point out that Ln(Ω) values calculated based on IC_{50} measurements cannot be interpreted simply as a measure of physical interactions. In addition, the Hill slope is 1.18 ± 0.03 for inhibition of KCNQ4 by SsTx (Fig. 1F), and thus multiple SsTx molecules could interact with positive synergy to inhibit the channel. Consistent with this interpretation, docking of SsTx onto a structural model of KCNQ4 shows that multiple SsTx can be accommodated simultaneously (Fig. 3E and Fig. S2G). Regardless of these complications, D266 and D288 are highly conserved in all members of the KCNQ family (Fig. S2H) and thus could form the same “hot spots” in the pore region of all KCNQ channels. The negatively charged surface patch is also complementary to the positively charged surface on the toxin. Overall, our results indicate that the molecular interaction between SsTx and KCNQ channels is formed by at least two specific interacting residue pairs as well as van der Waals and electrostatic interactions contributed by K4, K10, K11, and K45.

**SsTx Impairs Cardiovascular Systems.** We subsequently investigated the in vivo effects of SsTx and centipede crude venom on cardiovascular systems. Following 5- and 10-mg/mL administration of the venom, the constriction force of the coronary artery increased by ~1.6 (80%) and ~2.1 (105%) g, respectively, compared with ~2.2 g (110%) induced by the positive control, saturated phenylephrine (Fig. 4D). To determine if SsTx is the key component of the venom that causes vessel disorder, we compared the vessel activity of crude venom both in the presence and absence (CV-SsTx free or CV-Sf) of SsTx. As shown in Fig. 4B, CV-Sf (crude venom without SsTx) has only a small effect on arterial tension. Compared with 10 mg/mL crude venom-induced
vasoconstriction, which was ~105%, 10 mg/mL CV-Sf only increases the vasoconstriction by ~7.5%, indicating that SsTx is the main active component in the venom that exerts vessel activity. As expected, 1 and 5 μM SsTx increased the constriction force of the coronary artery by ∼50.3 and 105%, respectively (Fig. 4E). Additionally, 0.5 mg/kg SsTx induced acute hypertension in mice (Fig. 5A). Intravenous injection of crude venom induced acute hypertension in mice (Fig. 5A). Additionally, 0.5 mg/kg SsTx induced vasoconstriction, which further caused a significant increase in blood pressure (increased by 30.4 ± 5.7% in systolic blood pressure and by 40 ± 6.3% in diastolic blood pressure) in mice (Fig. 5B). SsTx also caused vasospasm and acute hypertension in Macaca monkeys while atropine and retigabine inhibited the effects (Fig. 5C and D). Intravenous injection of retigabine (1 mg/kg) reversed hypertension induced by SsTx (0.1 mg/kg) as the blood pressure was reduced from ~196 mmHg to a normal range (Fig. 5D). Following i.v. injection of SsTx, the mice kept twitching and shaking before death (Movie S2), yielding an LD50 of ~0.85 mg/kg and the elimination half-life of 4.5 h (Fig. S6C). This lethal activity is likely due to coronary-induced myocardial ischemia, because we observed an inverted T wave in monkeys (Fig. 5E) after 5 min of i.v. injection of SsTx. The inverted T wave is a typical syndrome for myocardial ischemia (32). The inverted T wave was also reversed by retigabine (Fig. 5E). Thus, SsTx-induced coronary spasm plays a key role in myocardial ischemia, which usually progresses to heart failure if ischemic symptoms are prolonged (33). Unlike scorpion toxins that inhibit BKCa currents to induce vasospasm (34, 35), centipedes may use a different strategy by inhibiting KCNQ4 (Fig. S1O). We estimate that one bite from a centipede likely injects 15–30 μL crude venom, which has a concentration of ~400 mg peptides per milliliter, is sufficient to subdue a small rodent (Movie S1). Although crude venoms from certain snakes and snails have higher lethal activity than that of centipedes (36, 37), it is not clear how much volume is injected in each attack.

**SsTx Causes Disorders of Nervous and Respiratory Systems.** Neuronal KCNQ channels (including KCNQ2, 3, and 5) are the molecular correlates of the M-current (19). The importance of M-channels is illustrated by the fact that a 25% decrease in M-current leads to neonatal epilepsy in humans (38) and that a disruption of kcnq2 in mice is lethal (39). Since our study shows that SsTx acts as a potent inhibitor of KCNQ2, KCNQ3, and KCNQ5 (Fig. 1F), we would like to test whether SsTx could cause seizure or severe nervous disorder in small animals. Since SsTx does not cross the blood–brain barrier, the risk of seizure in human after envenomation is less of a concern. However, it could be relevant in small animals where a centipede bite could go through the skulls. After...
potential generation (39) and the release of neurotransmitter (40), the direct M-channel blockers are severely neurotoxic. To better understand the electrophysiological changes in hippocampal neurons after SsTx application, we conducted patch clamp recordings on brain slices of mice. At the concentration of 10 μM, SsTx evoked a spiking frequency of ~20 Hz, much higher than the untreated neurons (~12 Hz) (Fig. 6 A and B). Acetylcholine concentration in mice hippocampi increased by ~63.6% and ~81.8% after the administration of 1 μL of 10 μM SsTx and 100 μM linopirdine (a well-known inhibitor of KCNQ channels), respectively (Fig. 6C). As expected, retigabine inhibited the SsTx-induced increase in spiking frequency and acetylcholine secretion (Fig. 6 A–C), as well as seizure in mice (Movie S3).

Given KCNQ’s important function in the respiratory system (21), effects of the crude venom and SsTx on rat respiratory functions were subsequently investigated (Fig. 6 D–H). Following the i.v. injection of SsTx, the respiratory rate was significantly decreased while respiratory amplitude was increased. One-and-one-half hours after administration of 2 mg/kg SsTx, the respiratory rate was reduced by ~75% and respiratory amplitude increased by ~150% (Fig. 6 D–F). In vitro experiments showed that crude venom (Fig. 6G) or SsTx (Fig. 6H) can induce bronchial ring contraction and that the contraction can be reduced by retigabine. Our results show that SsTx and the crude venom cause disorders not only on the cardiovascular system but also on nervous and respiratory systems.

Discussion
Although venom is a potent weapon for prey capture, its regeneration requires an increased metabolic rate over a long duration. Thus, strategies of venom optimization likely have evolved to minimize venom expenditure by venomous animals (7). Targeting of multiple essential physiological systems by centipede venom is an example of such venom optimization. The centipede toxin SsTx simultaneously disrupts several essential systems, including respiratory, cardiovascular, and perhaps the muscular system, by blocking multifunctional KCNQ channels. SsTx causes cardiovascular toxic symptoms including hypertension, myocardial

![Fig. 5. SsTx causes disorders in the cardiovascular system. (A) Blood pressure of mice, including SBP and DBP recorded 10 min after i.v. injection of saline (n = 6 animals per bar) or CV (30 mg/kg, n = 6 animals per bar), *P < 0.05. (B) Blood pressure of mice, including SBP and DBP recorded 10 min after i.v. injection of saline (n = 10 animals per bar) or SsTx (0.5 mg/kg, n = 10 animals per bar), *P < 0.05. (C) Blood pressure of monkeys, including SBP (in red) and DBP (in blue) recorded every 3 min before and after i.v. injection of SsTx (n = 3, 0.1 mg/kg, solid line) and atropine. (D) Blood pressure of monkeys, including SBP (in red) and DBP (in blue) recorded every 3 min before and after i.v. injection of SsTx (n = 3, 0.1 mg/kg; solid line) and RTG (1 mg/kg). (E) Representative three-lead surface ECG of a monkey recorded 10 min after i.v. injection of saline (n = 3) or SsTx (n = 3, 0.01 mg/kg) and followed by a rescue with RTG (1 mg/kg) after SsTx (0.01 mg/kg) injection (n = 3). CV, crude venom; DBP, diastolic blood pressure; RTG, retigabine; SBP, systolic blood pressure.

![Fig. 6. SsTx interrupts nervous and respiratory systems. (A) Response of CA1 pyramidal neurons after application of 40 pA depolarizing current injected into the cell when challenged sequentially with 10 μM SsTx, 100 μM LNP, and the mixture of 10 μM SsTx and 40 μM RTG. Before applications of each compound, the brain slices were perfused by bath solution for 10 min to ensure that the firing rates return to the level before compound administration. (B) Summary of spiking frequency in the presence of bath solution (n = 7), 10 μM SsTx (n = 7), 100 μM LNP (n = 7), and the mixture of 10 μM SsTx and 40 μM RTG (n = 4). *P < 0.05. (C) Acetylcholine concentrations quantified 30 min after microinjection of 1 μL of saline (n = 6), 20 μM SsTx (n = 6), 100 μM LNP (n = 6), or the mixture of 10 μM SsTx and 40 μM RTG (n = 3) into the hippocampus of mice. *P < 0.05. (D) Representative respiratory traces on rats were recorded before and after 2 mg/kg SsTx administration by i.v. injection. The changes in rats’ respiratory rates (E) and amplitudes (F) when challenged with i.v. injected SsTx (n = 3 animals per point, 2 mg/kg). Representative vascular contractility of bronchial ring contraction when challenged sequentially with (G) CV and RTG (n = 3 rats) and (H) SsTx and RTG (n = 3 rats). CV, crude venom; LNP, linopirdine; RTG, retigabine.](image-url)
ischemia, vasospasm, and twitching (Figs. 4 and 5); neurotoxic symptoms such as seizure (Fig. 6 A and B); and respiratory disorders such as lowered respiratory rate (Fig. 6 D and E).

These symptoms are clinically relevant because they have been observed in patients following centipede envenomation (17, 41). Removing SsTx from the crude venom significantly decreased the toxicity on the cardiovascular system (Fig. 4 A and B), demonstrating that the single venom component SsTx plays a crucial role in the prey-capture strategy.

Among case reports, tissue necrosis is a very common symptom caused by centipede envenomation. Indeed, local injection of SsTx in the back skin of mice caused significant tissue necrosis (Fig. S5). This local effect of SsTx is likely caused by spasm of capillary vessels. In comparison with other reported centipede crude venom activities, such as platelet-aggregating and phospholipase A2 activities (42), SsTx also seems to be a predominant factor for local symptoms, especially tissue necrosis.

Severe clinical cardiovascular symptoms, even death, have been reported following centipede bites, yet no effective therapeutic interventions are available. In this study, we illustrate that KCNQ channels are the likely targets for centipede venom and that the KCNQ opener retigabine, which is approved for treating epilepsy, could be used to treat centipede envenomation. Our experiments at the molecular, cellular, organ, and animal levels showed that retigabine reverses the effect of either crude centipede venom or purified SsTx. In conclusion, our work reveals that the centipede has evolved a simple yet powerful strategy to disable multiple essential physiological systems in prey and shows that an equally simple strategy could be effective in neutralizing the venom’s toxicity.

Materials and Methods

Animal procedures were performed in compliance with the Institutional Animal Care and Use Committee of Laboratory Animals of Kunming Institute of Zoology, Chinese Academy of Sciences. For the video recording of the centipede’s bite, the mice were put into a 2-L conical flask to adapt to the environment. The centipede was delivered into the conical flask after the mice were stabilized. Purification, NMR, tissue isolation, cell culture, and electrophysiological measurements were performed by using standard approaches. See SI Materials and Methods for a detailed description.

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