Structural basis for voltage-sensor trapping of the cardiac sodium channel by a deathstalker scorpion toxin

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Voltage-gated sodium (Na\textsubscript{v}) channels initiate action potentials in excitable cells, and their function is altered by potent gating-modifier toxins. The α-toxin LqhIII from the deathstalker scorpion inhibits fast inactivation of cardiac Na\textsubscript{v}1.5 channels with IC\textsubscript{50} = 11.4 nM. Here we reveal the structure of LqhIII bound to Na\textsubscript{v}1.5 at 3.3 Å resolution by cryo-EM. LqhIII anchors on top of voltage-sensing domain IV, wedged between the S1-S2 and S3-S4 linkers, which traps the gating charges of the S4 segment in a unique intermediate-activated state stabilized by four ion-pairs. This conformational change is propagated inward to weaken binding of the fast inactivation gate and favor opening the activation gate. However, these changes do not permit Na\textsuperscript{+} permeation, revealing why LqhIII slows inactivation of Na\textsubscript{v} channels but does not open them. Our results provide important insights into the structural basis for gating-modifier toxin binding, voltage-sensor trapping, and fast inactivation of Na\textsubscript{v} channels.

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Eukaryotic voltage-gated sodium (Na\(_V\)) channels generate the inward sodium current that is responsible for initiating and propagating action potentials in nerve and muscle. The sodium current is terminated within 1–2 ms by fast inactivation. A wide variety of neurotoxins bind to six distinct receptor sites on Na\(_V\) channels and modify their function. α-Scorpion toxins and sea anemone toxins bind to Neurotoxin Receptor Site 3, dramatically inhibit fast inactivation of Na\(_V\) channels, and cause prolonged and/or repetitive action potentials. Scorpions utilize these toxins in their venoms to immobilize prey by inducing paralysis and causing cardiac arrhythmia. Because of their high affinity and specificity, scorpion toxins are used widely to study the structure and function of Na\(_V\) channels, α-Scorpion toxins bind to the voltage sensor (VS) in domain IV (DIV), which is important for triggering fast inactivation. Therefore, structures of the high-affinity complexes of α-scorpion toxins and Na\(_V\) channels will provide critical information for understanding the structural basis for toxin binding, voltage-sensor trapping, and fast inactivation.

Eukaryotic Na\(_V\) channels contain four homologous, nonidentical domains composed of six transmembrane segments (S1–S6), organized into a voltage-sensing module (VS, S1–S4) and a pore module (PM, S5–S6) with two intervening pore helices (P1 and P2). The S4 segments contain four to eight repeats of a positively charged residue (usually Arg) flanked by two hydrophobic residues. These positively charged residues serve as gating charges, moving outward upon depolarization to initiate the process of activation. Chemical labeling and voltage clamp fluorometry suggest that DIV-IV and DIV-VS are primarily responsible for activation of the channel, whereas DIII-IV induces fast inactivation. A triple hydrophobic motif, Ile-Phe-Met (IFM), located in the DIII-DIV linker, serves as the fast inactivation gate. Mutation of the IFM motif can completely abolish fast inactivation.

Determination of the structures of prokaryotic and eukaryotic α-Scorpion channels has remarkably enriched our understanding of their structure and function. Those structures revealed that Na\(_V\) channels share similar key structural features. The central pore is formed by the four PMs with the four VSs arranged in a pseudosymmetric square array on their periphery. The four homologous domains are organized in a domain-swapped manner, in which each VS interacts most closely with the PM of the neighboring domain. The four S6 segments come together at their intracellular ends to form the activation gate. Intriguingly, in the structures of mammalian Na\(_V\)s, the IFM motif binds in a receptor site formed by the DIII S4–S5 linker and the intracellular ends of the S5 and S6 segments in DIV, which suggests a local allosteric mechanism for fast inactivation of the pore by closing the intracellular activation gate.

The α-scorpion toxins bind to DIV-VS in its resting state, trap it in an intermediate-activated conformation, and inhibit fast inactivation, providing an attractive target for studying the coupling of DIV-VS to pore opening and fast inactivation. Strong depolarization can reverse voltage-sensor trapping and drive the α-scorpion toxin off its receptor site, providing direct evidence for a toxin-induced conformation of the VS. The cryo-EM structure of the α-scorpion toxin AaHII was resolved bound to two different sites on a nonfunctional chimera of the cockroach sodium channel Na\(_V\)Pas, which contained 132 amino acid residues of the DIV-VS of the human neuronal sodium channel Na\(_V\)1.7 embedded within 1449 residues of Na\(_V\)Pas. These results revealed structures of AaHII bound to the VSs in both DIII and DIV but did not resolve whether AaHII bound to either of these sites was functionally active in the chimera. Therefore, the precise structural mechanism by which α-scorpion toxin binds to the DIV-VS in a native sodium channel and blocks fast inactivation still remains elusive.

LqhIII from the “deathstalker scorpion” Leirus quinquestratus hebraeus (also known as the Israeli yellow scorpion and the North African striped scorpion) is classified as an α-scorpion toxin and shares the common αββ scaffold containing four pairs of Cys residues that form disulfide bonds. Most scorpion toxins paralyze prey by targeting the sodium channels in nerve and skeletal muscle specifically. In contrast, LqhIII binds with the highest affinity to the human cardiac sodium channel, with an estimated EC\(_{50}\) of 2.5 nM. It prevents fast inactivation efficiently, and it dissociates at an extremely slow rate, making it exceptionally potent.

In this work, we elucidate the molecular mechanisms of voltage-sensor trapping and block of fast inactivation by α-scorpion toxins in the context of a functional native toxin–receptor complex by determining the cryo-EM structure of rat cardiac sodium channel Na\(_V\)1.5 in complex with the α-scorpion toxin LqhIII at 3.3 Å resolution. Our experiments provide important insights into the structural basis for gating-modifier toxin interaction, voltage-sensor trapping, electromechanical coupling in the VS, and fast inactivation of the pore.

**Results**

**Voltage-sensor trapping of Na\(_V\)1.5 by LqhIII.** For our structural studies, we took advantage of the fully functional core construct of the rat cardiac sodium channel Na\(_V\)1.5 (rNa\(_V\)1.5c), which can be isolated with high yield and high stability. Expression of rNa\(_V\)1.5c in the human embryonic kidney cell line HEK293S GnTT– and recording from single cells in whole-cell patch clamp mode (see “Methods”) yields inward sodium currents that activate rapidly and inactivate within 6 ms (Fig. 1a, black trace; the inward current is plotted as a negative quantity by convention). Perfusion of increasing concentrations of LqhIII progressively slows the fast inactivation process and makes it incomplete (Fig. 1a, colored traces). We measured the sodium current remaining 6 ms after the depolarizing pulse as a metric of LqhIII action (Fig. 1a, dotted line), because the unmodified sodium current has declined to nearly zero by this time, whereas substantial toxin-modified sodium current remains. The EC\(_{50}\) value for the increase in sodium current remaining at 6 ms following the stimulus is 11.4 nM (Fig. 1a). This effect of LqhIII and other α-scorpion toxins is achieved by trapping the VS in DIV of sodium channels in a conformation that allows sodium channel activation but prevents coupling to fast inactivation. Voltage-sensor trapping develops slowly and progressively over more than 20 min, with a half-time of 11.3 min at 100 nM (Fig. 1b). As expected from previous work, strong depolarizing pulses to +100 mV cause dissociation of the toxin and loss of its blocking effect on fast inactivation (Fig. 1c). The molecular mechanism for this long-lasting voltage-dependent block of fast inactivation of Na\(_V\)1.5 sodium currents by LqhIII is unknown.

**Structure determination of rNa\(_V\)1.5c/LqhIII complex by cryo-EM.** We analyzed the structural basis for the potent voltage-sensor trapping effects of LqhIII by cryogenic electron microscopy (cryo-EM). LqhIII was incubated with purified rNa\(_V\)1.5c for 30 min. The regulatory proteins FGF12b and calmodulin were added to stabilize the isolated protein, and the toxin/channel complex was further purified by size-exclusion chromatography (SEC). A symmetric peak of the toxin/channel complex was collected from the second SEC run (Supplementary Fig. 1a, b). Detailed descriptions of protein expression, purification, cryo-EM imaging, and data processing are presented in “Methods.” Cryo-EM data were collected on a Titan-Krios electron microscope and processed using RELION (Supplementary Fig. 1c, d; Supplementary Fig. 2a–c). A 3D reconstruction map...
of the rNav1.5c/LqhIII complex was obtained at an overall resolution of 3.3 Å, based on the Fourier Shell Correlation (FSC) between independently refined half-maps (Fig. 2a). Strong density specifically localized near the extracellular side of DIV-VS shows that there is only one LqhIII molecule bound to rNav1.5C (Fig. 2b, purple), as expected from previous biochemical studies of scorpion toxin binding to sodium channels 9. The local resolution for the PM core region is ∼3.0–3.5 Å, whereas the four peripheral VSS have local resolutions of ∼3.5–4.0 Å (Fig. 2c). The resolution for the toxin is lower than the channel protein (∼4.0–5.0 Å, Fig. 2c). However, the interacting surface of the toxin that binds to DIV-VS has a resolution of ∼3.5–4.0 Å for the amino acid side chains that form the complex, as they are tightly bound (Supplementary Fig. 2d, e). The 3D structure of the tightly disulfide-crosslinked toxin is well-known from previous studies (Supplementary Fig. 3a) 29, allowing it to be accurately fit into the observed density. No significant density was observed at high resolution for the C-terminal domain (CTD), FGF12b, or calmodulin (Fig. 2b), indicating that these components of the purified protein complex are mobile.

The overall structure and LqhIII binding site. The high-resolution cryo-EM density map allowed us to build an atomic model for the rNav1.5C/LqhIII complex (Fig. 3a, b). The overall structure of the rNav1.5C/LqhIII complex is very similar to our previous apo-rNav1.5C structure 21, with a minimum RMSD of 0.78 Å over 1164 residues. However, local conformational differences give many important insights. The structure of LqhIII is rigidly locked by disulfide bonds, except for the β2β3 loop and C-terminal region, which are highly flexible in solution as revealed by nuclear magnetic resonance analyses (Fig. 3c). Remarkably, LqhIII uses these two flexible regions to bind to the extracellular side of DIV-VS by wedging its β2β3 loop and C-terminus into the aqueous cleft formed by the S1–S2 and S3–S4 helical hairpins (Fig. 3b). These features are in close agreement with previous molecular-mapping studies of neurotoxin receptor site 311 and with the structure of the AaHII/NaVPas-NaV1.7 chimera 26 (see “Discussion”). The toxin may attack Neurotoxin Receptor Site 3 in the DIV-VS using its most flexible regions to allow it to dock in a stepwise manner that results in a tight induced-fit complex.

The close interactions of the C-terminus and the β2β3 loop of LqhIII with rNav1.5C are illustrated in Fig. 3b, d. At the C-terminus of LqhIII, Glu63 interacts with the Asn329-linked glycans from DI-PM, and Lys64 dips into the aqueous cleft and interacts with Gln1615 (Fig. 3b, d). The end of the β2β3 loop inserts into the DIV-VS cleft and partially unwinds the last helical turn of the S3 segment. Mutagenesis studies mapping Neurotoxin Receptor Site 3 revealed a negatively charged residue in the extracellular S3-S4 linker that is conserved among Nav channels and is critical for α-scorpion toxin binding 4,9. In agreement with those studies, the conserved negatively charged residue Asp1612 at an equivalent position in NaV1.5 mediates this crucial interaction with the bound toxin. His43 and His15 wrap around Asp1612 like pincers forming a hydrogen bond (∼2.5 Å) and a potential salt bridge (∼4.0 Å), respectively (Fig. 3d). Moreover, we note that the backbone carbonyl of His43 engages the backbone carbonyl of Thr1608 at a distance of 2.8–3.5 Å, which may contribute to the affinity or specificity of interactions with the β2β3 loop (Fig. 3d) 30.
**Fig. 2** Cryo-EM structure of the rNaV1.5C/LqhIII complex. 

- **a** The FSC between independently refined half-maps for rNaV1.5C/LqhIII reconstruction.
- **b** Overall cryo-EM reconstruction (side view, left; top view, right) of the rNaV1.5C/LqhIII complex. NaV1.5C and LqhIII colored in grey and purple, respectively.
- **c** Local resolution (side view, left; top view, right) of the EM map colored from blue to red representing resolution from high to low (sidebar).

**Fig. 3** Overall structure of rNaV1.5C/LqhIII complex and LqhIII binding site. 

- **a** Cartoon representation of the overall structure of the NaV1.5C/LqhIII. LqhIII, DIV-VS and DIII-DIV linker were colored in purple, blue and orange, respectively. The same color scheme is applied hereafter in this paper unless specified otherwise. The glycosyl moieties are shown in sticks colored in yellow. The black dash square is indicated for panel (b).
- **b** Zoom-in view of LqhIII binding to the DIV-VS. The black dash square indicated for panel (d).
- **c** The NMR structure of the LqhIII (PDB code: 1FH3) indicating the flexibility of the β2β3 loop and C-terminus.
- **d** Detailed interactions between LqhIII and DIV-VS. Key residues shown in sticks were labeled. Interaction surfaces of the DIV-VS (blue) and the LqhIII (purple). Key residues for the interaction shown in yellow shading and embedded sticks.
An intermediate-activated state of DIV-VS trapped by LqhIII. Fast inactivation of Na\textsubscript{V} channels requires activation of DIV-VS\textsuperscript{9,10,12,13}. Because there is no membrane potential during solubilization and purification, the VSs of published Na\textsubscript{V} structures are usually in partially or fully activated states. In our apo-rNa\textsubscript{V,1.5C} structure, four of the six gating charges of DIV-VS pointed outward on the extracellular side of the hydrophobic constriction site (HCS), as expected for an activated state\textsuperscript{21}. As a result, the fast inactivation gate in the apo-rNa\textsubscript{V,1.5C} structure binds tightly in a hydrophobic pocket next to the activation gate\textsuperscript{21}.

A hallmark feature of the action of α-scorpion toxins is strongly voltage-dependent dissociation from their receptor site, which correlates with the voltage dependence of activation of sodium channels (Fig. 1\textsuperscript{9,24,25}). The structure of the rNa\textsubscript{V,1.5C}/LqhIII toxin complex reveals the molecular basis for this important aspect of scorpion toxin action. In the complex of the toxin with the partially activated state of the DIV-VS, the positive charge of the ε-amino group of K\textsubscript{64} on LqhIII interacts with the same negatively charged side chain in the ENC that interacts with R1 and R2 in the activated conformation of the VS (Fig. 4a, b). In light of these structures, it seems likely that outward movement of the S4 segment during activation of the DIV-VS creates a clash with K\textsubscript{64} and causes toxin dissociation by both electrostatic repulsion and steric hindrance. This potent combination of electrostatic repulsion and physical clash is sufficient to overcome the high binding energy of the α-scorpion toxins.

Superposition of the fully activated state (grey) and toxin-induced intermediate-activated state (blue) of the DIV-VS revealed a remarkable conformational difference (Fig. 4c). From S1 through most of S3 there is little or no structural change, whereas the final two helical turns of S3 and the entire S4 segment undergo dramatic conformational shifts. Notably, Gly1607 serves as a pivot point for S3 rotation, and the rotation of upper S3, in turn, moves S4 downward ~11 Å, such that R1 and R2 in the intermediate-activated state are approximately in the positions of R3 and R4 in the fully activated state (Fig. 4c). This toxin-induced conformational change in the S3–S4 linker is further documented by our fit to the cryo-EM density, which is illustrated in Supplementary Fig. 3c–e. At the intracellular end of S4, an elbow-like bend is formed between S4 and the S4–S5 linker, which pushes the S4–S5 linker inward ~4.6 Å inward at its N-terminal end (Fig. 4c). Intriguingly, our previous resting-state structure of Na\textsubscript{V}\textsubscript{Ab} showed that a similar elbow pushes the S4–S5 linker and its connection to S4 strikingly inward and twists this segment in order to close the intracellular activation gate\textsuperscript{31}. This conformational change in the S4–S5 linker is further supported by the close fit of our structural model to the cryo-EM density (Supplementary Fig. 4a).

Partially open intracellular activation gate. Rapid voltage-dependent activation is one of the signature functions of Na\textsubscript{V} channels\textsuperscript{12}. A full understanding of the mechanism of activation requires structural information on Na\textsubscript{V} channels in different states. Side-by-side comparison of our previous activated-state structure with our current LqhIII-bound intermediate-activated structure of rNa\textsubscript{V,1.5C} reveals key steps in the coupling of the conformational changes of the DIV-VS to the intracellular activation gate (Figs. 4c and 5). To reach the intermediate-activated state (blue) from the fully activated state (grey), S4 of DIV-VS moves inward ~11 Å and pushes the DIV S4–S5 linker inward ~4.6 Å through the formation of an elbow-like bend (Figs. 4c and 5a). The shifted DIV S4–S5 linker engages DIV-S6 through Ser1655 (Figs. 4c and 5c). As a result, DIV-S6 shifts toward the pore axis, and this movement pushes DI-S6 away from the center of the orifice (Fig. 5a, b and Supplementary Fig. 4b, c). Meanwhile, the DI S4–S5 linker moves outward away from the activation gate, which shifts both DI-S6 and DII-S6 away from the
pore axis and contributes to a more open conformation. The shifts of the four S6 segments result in an enlarged activation gate with a van der Waals diameter of 6.6 Å (Fig. 5b and Supplementary Fig. 4b, c), which is ∼1 Å larger than the activation gate in apo-rNaV1.5C, but ∼2 Å smaller than the expected orifice of the activation gate of fully open rNaV1.5C when modeled using the open state of NaVAb21,32. Unambiguous density shows Tyr1769 in D IV-S6 in an outward-pointing conformation making the opening of the activation gate larger, whereas it is pointed inward in the fully activated structure (Fig. 5b). Together, these conformational movements result in a wider opening in the activation gate in the toxin-induced intermediate-activated state, which could in principle be sufficient for sodium conductance.

Loosely bound fast inactivation gate. In activated NaV structures, the IFM motif of the fast inactivation gate binds to a hydrophobic pocket formed by DIII S4–S5 linker and the intracellular ends of DIV-S5 and DIV-S619–21. In the intermediate-activated rNaV1.5C/LqhIII structure, weak, yet consistent, density for the IFM motif suggests that the conformational changes of the VS and activation gate have in turn altered the conformation of apo-rNaV1.5C structure (Supplementary Fig. 4b, c). Considering that the two proteins were purified following the same procedures, it is likely that the same set of lipid and detergent molecules would be available for binding. Therefore, we believe the larger lipid or detergent molecule is able to bind in the lumen of the activation gate of the intermediate-activated structure because of its larger diameter rather than because of a change in lipid or detergent concentrations between the two protein preparations.

Fig. 4 Conformational Change of DIV-VS. a, b Structures of the activated NaV1.5 DIV-VS and intermediate-activated NaV1.5 DIV-VS were colored in grey and blue, respectively. Gating charges (grey or blue), ENC (red), HCS (yellow), and INC (red) were shown in sticks. The shift of each gating charge was indicated by black dashed lines. c Superposition of NaV1.5 DIV-VS between activated and intermediate-activated state. Red arrows indicate the conformational changes. d Superposition of the intermediate-activated NaV1.5 DIV-VS and resting-state NaVAb-VS.
the bound IFM motif and made it less stable (Fig. 5d, e, Supplementary Fig. 4b, c). Evidently, the movement of DIV-S6 squeezes the binding pocket, and the shift of DIII-S6 destabilizes binding of the IFM motif. We observed a small shift in the position of the IFM motif in the intermediate-activated structure, as well as weaker density for the Met side chain, consistent with greater mobility and partial dissociation (Fig. 5d, e). These changes are illustrated most clearly in Fig. 5e by comparing cryo-EM density contoured at 4σ, 5σ, and 6σ in the presence and absence of LqhIII. We propose that α-scorpion toxins inhibit fast inactivation by both trapping the DIV-VS in an intermediate-activated conformation and altering the shape of the IFM binding pocket, which causes slowed IFM binding, more rapid dissociation of the IFM motif, and release of the fast inactivation gate. These effects are responsible for the slowed rate of fast inactivation and the incomplete extent of fast inactivation that are hallmarks of the action of LqhIII and other α-scorpion toxins (Fig. 1).

**LqhIII does not open the intracellular activation gate.** Our static cryo-EM view of the intracellular activation gate of apo-rNav1.5C suggests that it is partially open compared to our tightly closed resting-state model of rNav1.5C based on the NaVAb resting state21,31. However, it does not appear to be open enough to conduct hydrated Na⁺32,33. To test this hypothesis, we used molecular dynamics methods similar to those we previously applied to the NaVAb structure32,33 in order to investigate the effect of LqhIII on pore hydration and dilation of the intracellular activation gate (Fig. 6). The inner pore of rNav1.5C is depicted...
Fig. 6 Molecular dynamics analysis of hydration and Na\(^{+}\) permeation through the rNa\(_{1.5}\)/LqhIII complex. a Side view of rNa\(_{1.5}\) (orange ribbons; domains II and IV) from MD simulations highlighting Na\(^{+}\) ions (blue spheres), the water-occupied volume within a cylinder of radius 8.5 Å (red surface), and the protein-occupied volume within a cylinder of radius 12 Å (colorless surface). In this snapshot, the protein cavity (outlined in a black rectangle) at the intracellular activation gate (ICAG; purple shaded region, \(-2.8\) nm < z < \(-1.5\) nm) is dehydrated. The QuickSurf representation in VMD was used for surfaces. b Average hydration along the pore-axis for simulations of rNa\(_{1.5}\)/LqhIII (red line) with and (black line) without the toxin. Overall pore hydration was unchanged whether or not the toxin is included in simulations. Shading corresponds to the standard error of the mean (s.e.m.). c Molecular representations of the gate containing \(N_{\text{water}} = 3\) (left) or 15 (right) water molecules. d Average probability distribution of \(N_{\text{water}}\) in the gate. The gate is more likely to contain 10 or more water molecules when the toxin is present. Data points and shading represent mean and s.e.m. e Bottom (intracellular) view of the activation gate for cryo-EM rNa\(_{1.5}\) (PDB ID: 6UZ3; cyan), cryo-EM rNa\(_{1.5}\)/LqhIII (yellow), and an example conformation from the most sampled basin in MD (magenta) are superimposed. The distances \(d_1\) and \(d_2\) between opposing helices are shown schematically (see “Methods”). f Free energy of \(d_1\) vs. \(d_2\) computed from MD simulations. The reference cryo-EM structures of (cyan+) rNa\(_{1.5}\) and (yellow+) rNa\(_{1.5}\)/LqhIII are indicated. Contour lines are shown every 0.5 kcal/mol from 0 to 4 kcal/mol. In the simulations, the intracellular activation gate often contracts and adopts an asymmetric conformation with \(d_2 < d_1\), with the symmetric conformation observed by cryo-EM corresponding to a metastable state 0.5 to 1 kcal/mol higher in free energy. Time frames spread across 30 independent simulations of rNa\(_{1.5}\) with or without LqhIII were used for analyses (\(n = 29,026\) frames for rNa\(_{1.5}\) and \(n = 29,899\) frames for rNa\(_{1.5}\)/LqhIII).
with $d_x$ deviating by up to 4 Å from the toxin-bound cryo-EM structure (Fig. 6f). Overall, asymmetric conformations in which $d_x < d_i$ are slightly favored (by 0.5–1.0 kcal/mol) relative to the symmetric cryo-EM conformation of rNaV1.5C/LqhIII. The average values of both $d_i$ and $d_x$ undergo small but significant increases relative to simulations without LqhIII, as does the average number of water molecules in the gate (Supplementary Fig. 5a). Although the pore remains predominantly dewetted irrespective of the presence of LqhIII, fluctuations leading to more dilated conformations of the activation gate are correlated with larger hydration numbers (Supplementary Fig. 6a), with transient occurrences of 10 or more water molecules connecting the central cavity to the intracellular environment (Fig. 6c, d). However, the activation gate is not significantly more likely to be wetted in simulations of rNaV1.5C with LqhIII than in simulations without LqhIII (Supplementary Fig. 5b), consistent with the fact that toxin binding does not open the gate sufficiently for passage of Na$^+$. Nevertheless, the analysis of fluctuations in diameter and hydration of the intracellular activation gate provides an initial suggestion that LqhIII binding may facilitate the transition to the open state of NaV1.5, which requires activation of the VSs in domains I–III for completion.

**Discussion**

We determined the structure of rat NaV1.5C in complex with the α-scorpion toxin LqhIII by single-particle cryo-EM. Biochemical and biophysical studies support only a single neurotoxin receptor site 3 per sodium channel located in the VS in DIV, at which α-scorpion toxins, sea anemone toxins, and related gating-modifier toxins bind$^{9,10}$. Consistent with this expectation from functional studies, we found a single molecule of LqhIII bound to the VS in DIV. The toxin binds at the extracellular end of the aqueous cleft formed by the S1–S2 and S3–S4 helical hairpins in the VS through its β2β3 loop and its C-terminal. Many conserved amino acid residues that are important for α-scorpion toxin binding and its functional effects on sodium channels are located in key positions in the toxin–receptor-binding interface (Fig. 3e). These results provide convincing evidence that we have correctly identified the pharmacologically important Neurotoxin Receptor Site 3 on the native cardiac sodium channel and defined its mode of toxin binding at high resolution. α-Scorpion toxins bind to Neurotoxin Receptor Site 3 in a voltage-dependent manner, with high-affinity binding to the resting state$^{3,24,25}$. Depolarization reduces toxin affinity and causes toxin dissociation$^{9,24,25}$. The bound toxin prevents the normal outward movement of the gating charges in DIV-S4, as measured directly from gating currents$^{10}$. The outward movement of the DIV-S4 segment correlates with fast inactivation, as measured by voltage clamp fluorometry with specifically labeled S4 residues$^{12,13}$. These studies highlight the importance of DIV-S4 and neurotoxin receptor site 3 in triggering fast inactivation. LqhIII prefers to bind to the resting state of rNaV1.5C, and strong depolarization causes dissociation in biochemical and electrophysiological studies using nM concentrations of toxin (Fig. 1)$^{9,24,25}$. In spite of that, the binding affinity of LqhIII for rNaV1.5C is high enough to overcome this opposing electrostatic energy gradient when high concentrations of rNaV1.5C and LqhIII are used to drive the binding interaction and induce gating charge transfer of DIV-VS into the intermediate-activated state. The S4 segment of the intermediate-activated DIV-VS has shifted two helical turns inward when compared with that of activated DIV-VS; however, it is not as deeply inward as in the resting state of the NaV1.5-AB-VS, which suggests the toxin-induced intermediate-activated state is indeed a normal intermediate state in the function of the VS. To account for the functional effects of the toxin, the toxin-modified VS must allow activation and pore opening, even while impairing fast inactivation. Trapping the DIV-VS in an intermediate-activated state that does not trigger fast inactivation, but nevertheless is permissive for activation driven by the other three VSs, would satisfy this mechanistic requirement. Thus, we propose that the toxin-induced intermediate-activated state–defined here is indeed the voltage-sensor trapped state that is responsible for the gating-modifier properties of the α-scorpion toxins. The Binding of LqhIII traps the VS in a unique conformation. The S4 segment is in $\beta_{10}$ helical conformations from R1 to R4, with two gating charges on each side of the HCS (Fig. 4). This conformation places the four gating charges in a straight line. R1 and R2 interact strongly with two negatively charged side chains of the ENC, whereas R3 and R4 interact strongly with two negatively charged side chains in the INC. These four ion-pair interactions bridging the HCS are unique in VS function. No other position of the primary gating charges can simultaneously form four ion-pair interactions bridging the HCS. We propose that this unique, toxin-stabilized conformation of the gating charges is the key voltage-sensor-trapped state that allows the deathstalker scorpion toxin to paralyze and kill its prey. The movement of the S4 segment implied by comparison of the structures of apo-NaV1.3C and LqhIII/rNaV1.5C fits closely with the sliding-helix model of voltage sensing$^{31,38,39}$. Compared to the activated state, in the toxin-induced intermediate-activated state, the S4 segment moves inward ~11 Å and rotates slightly as $\beta_{10}$ helix is converted to alpha-helix on the intracellular side of the HCS. The gating charges exchange ion-pair partners from the ENC to the INC; however, they do not move as far inward as observed in the resting state. Thus, the toxin-induced conformational change in the VS exactly follows the proposed voltage-driven movement of the S4 segment based on the sliding helix voltage-sensing model and the structures of the activated and resting states of NaV1.5$^{31}$.

Our complex structure provides an excellent model for investigating the coupling between gating charge transfer and fast inactivation. The cryo-EM structure of AaaIII/NaV1.7-DIV-VS chimera suggested that R5 and K6 of NaV1.7-DIV-VS were stabilized by interaction with the NaV1.7-CTD. By contrast, in our fully functional LqhIII/rNaV1.5C structure, the CTD was not observed. In fact, the CTD’s also were not observed in the high-resolution structures of human NaV1.2, 1.4, and 1.7 channels either$^{19–21,40}$. These results suggest that the CTDs of native mammalian NaV channels are disordered and/or mobile and differ substantially from the cockroach CTD in the Na Vpas-NaV1.7 chimera, whose amino acid sequence is not similar to the CTD’s of mammalian NaV channels. Based on this comparison, it seems likely that the CTD plays a secondary role or a regulatory role in fast inactivation in mammalian NaV channels, which may include transient interactions with the potential R5 and R6 gating charges of DIV-S4. Our structure gives additional insights into the structural basis for coupling of the conformational change of the intermediate-activated DIV-VS to fast inactivation. The inward movement of the DIV-S4 segment and its gating charges from their activated positions propagates a voltage-driven conformational change inward to the DIV-S4–S5 linker, forming an elbow that angles the S4–S5 linker into the cytosol. This substantial movement twists the S4–S5 linker and disturbs the conformation of the binding pocket for the IFM motif, thereby destabilizing the fast-inactivated state. In contrast, the forward coupling of VS movement to fast inactivation likely involves outward movement of the S4 segment, unbending the elbow, untwisting the S4–S5 linker, and opening the receptor site for binding of the IFM motif of the fast inactivation gate. Binding of LqhIII would oppose this series of conformational events that lead to fast inactivation. Compared to the resting state, the outward movement of the DIV-S4 segment in the partially activated VS and the resulting loosening of the elbow bend in the DIV.
S4–S5 linker also partially opens the activation gate through the coupled movement of the nearby intracellular end of DI-S6 away from the pore axis. Our molecular dynamics analyses show that this partially open activation gate structure is still functionally closed with respect to Na⁺ conductance. Nevertheless, this conformational change may be an essential permissive movement of DIV that allows subsequent activation of the DI–DIV VSs to drive pore opening with only partial activation of the DIV VS in its toxin-induced intermediate-activated state. This coupling among domains may occur through the domain-swapped organization that places the DIV-VS adjacent to the DI-PM. Consistent with this idea, the DI-S6 segment is moved away from the axis of the pore in the toxin-induced deactivated state, similar to its movement during the pore opening of Naᵥ₁.7. In this position, it would promote activation and pore opening of DIV–DIV activation gate residues to give the fully open channel.

In previous work, the structure of a chimera of the cockroach sodium channel Naᵥ₁.7 with the AaHII toxin bound was determined by cryo-EM. The functional significance of this sodium channel in the cockroach is unknown, and this chimera containing a segment of the DIV-VS of human Naᵥ₁.7 was nonfunctional; therefore, it is difficult to precisely compare this prior work to the structures we present here. Unexpectedly, AaHII bound to the Naᵥ₁.7-VS chimera in two positions, one on the VS in DI of Naᵥ₁.7 and one on the DIV-VS contributed in part by Naᵥ₁.7, and it was not shown whether either of these sites was functional in the chimera. In contrast, we found only a single toxin binding site, as expected from previous structure-function studies. Neurotoxin Receptor Site 3 identified in our study is generally similar to the AaHII binding site found in DIV of the AaHII/Naᵥ₁.7-VS chimera26, but we found an important difference in the binding poses of the two toxins. Compared with AaHII bound to the Naᵥ₁.7-VS chimera, LqhIII bound to rNaᵥ₁.5C is rotated ~26° downward, further away from the glycan and DI of the channel (Supplementary Fig. 7). This difference may reflect an alteration in the receptor site within the receptor complex caused by artifactual constraints from the formation of the chimera. Alternatively, the structure of the functionally active LqhIII/rNaᵥ₁.5C complex described here may be characteristic of the cardiac sodium channel, which has numerous distinct features compared to neuronal sodium channels like Naᵥ₁.7. The exact position and composition of the glycan moiety adjacent to Neurotoxin Receptor Site 3 in rNaᵥ₁.5C may be one potentially important point of difference. Despite this difference in the position of the toxin bound, the overall structures of the two toxin-bound DIV-VSs are remarkably similar with an RMSD of 1.41 Å over 112 residues. This striking structural similarity indicates that the DIV-VSs in these two channel constructs are locked in a similar state by toxin binding at 0 mV.

Sea anemone toxins are not similar to scorpion toxins in amino acid sequence, yet they bind to Neurotoxin Receptor Site 3 and inhibit fast inactivation like α-scorpion toxins. Some sea anemone toxins, like Anthopleurin A and B, are highly active on cardiac sodium channels. It is likely that the sea anemone toxins trap the DIV-VS in a similar intermediate-activated conformation as we have observed here for LqhIII. β-Scorpion toxins are similar in structure to α-scorpion toxins. They bind to Neurotoxin Receptor Site 4, which is located in the DIV-VS in an analogous position to Neurotoxin Receptor Site 3 in the DIV-VS. They bind with high affinity to the activated state and trap the DIV- VS in its outward, activated position. This mode of voltage-sensor trapping enhances activation by shifting its voltage dependence to more negative membrane potentials. When they act synergistically in scorpion venom, β- and α-scorpion toxins negatively shift the voltage dependence of activation and block fast inactivation, respectively, resulting in persistently activated sodium channels, repetitive firing, depolarization block of neuromuscular transmission, and lethal arrhythmias in the heart. A large family of cysteine-knot toxins from spiders also act as gating modifiers of voltage-gated ion channels. They bind to the VS of Naᵥᵥ, Kᵥ, and Caᵥ channels in an analogous manner to α-scorpion toxins and either block or enhance activation of the VS. The spider toxins that enhance activation may also stabilize the unique voltage-sensor-trapped conformation of the VS that we have elucidated here, in which a linear configuration of the four primary gating charges bridges the HCS and interacts simultaneously with the ENC and INC in a high-affinity voltage-sensor-trapped complex. Thus, the toxin-bound state we have characterized here may have broad significance for VS trapping by a wide range of gating-modifier toxins from hundreds of species of spiders, scorpions, mollusks, and coelenterates, which all use this universal mechanism to immobilize their prey.

Methods

Electrophysiology: All experiments were performed at room temperature (21–24 °C) as described previously. Human HEK293S GnTI- cells (American Type Culture Collection Cat# CRL-3022) were maintained and infected on cell culture plates in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10% fetal bovine serum and glutamine/penicillin/streptomycin at 37 °C and 5% CO₂. Cells were held at -120 mV and 100-ms pulses were applied in 10 mV increments from -120 mV to +60 mV. A P/4 holding leak potential was set to -120 mV. An extracellular solution contained in mM: 140 NaCl, 2 CaCl₂, 2 MgCl₂, 10 HEPES, pH 7.4. Intracellular solution contained: 35 NaCl, 105 Gf, 10 EGTA, 10 HEPES, pH 7.4. Glass electrodes had a resistance of 1.5–3 MΩ. Currents resulting from applied pulses were filtered at 5 kHz with a low-pass Bessel filter and then digitized at 20 kHz. Data were acquired using an Axopatch 200B amplifier (Molecular Devices). Voltage commands were generated using Pulse 8.5 software (HEKA, Germany) and ITClan analog-to-digital interface (Instrutech, Port Washington, NY).

Protein expression and purification: Delineated expression and purification of rat Naᵥ₁.5C were described in our previous study. Briefly, Naᵥ₁.5C was expressed in HEK293S GnTI- cells (ATCC). The protein was extracted by 1% (w/v) n-dodecyl-β-maltopyranoside (DDM, Anatrace) and 0.2% (w/v) cholesteryl hemisuccinate Tris salt (CHS, Anatrace) in Buffer A containing 25 mM HEPES pH 7.4, 150 mM NaCl and 10% glycerol. After centrifugation, the supernatant was agitated with anti-Flag M2 antibody resin (Sigma). The mixture was then re-loaded to Buffer A supplemented with 0.06% glycol-dioisoside (GDN, Anatrace). Purified protein was then loaded onto a Superose-6 column (GE Healthcare) in 20 mM HEPES pH 7.4, 150 mM NaCl and 0.06% GDN, peak fractions were concentrated to ~1 mg/ml and mixed with 50 μM LqhIII (Latoxon Laboratory) and calmodulin overnight. The mixture was then re-loaded to Superose-6 column pre-equilibrated with buffer containing 25 mM imidazole pH = 6.0, 150 mM NaCl and 0.006% GDN. Finally, peak fractions were concentrated to 40 μl at 5 mg/ml.

Cryo-EM grid preparation and data collection: Three microliters of purified sample were applied to glow-discharged holey gold grids (UltraAstar foil, 300 mesh, R1.2/1.3), and blotted for 2.0–3.5 s at 100% humidity and 4 °C before being plunged frozen in liquid ethane cooled by liquid nitrogen using an FEI Mark IV Vitrobot. All data were acquired using a Titan Krios transmission electron microscope operated at 300 kV, a Gatan K2 Summit direct detector, and Gatan Quantum GIF energy filter with a slit width of 20 eV. A total of 4222 movie stacks were automatically collected using Leginon and cataloged with MotionCorr2. A total of 3805 micrographs with CTF fitted better than 6 Å were used for particle picking, and a total of 1,817,940 particles were automatically picked in RELION3.0. After several rounds of 2D classification, 882,608 good particles were selected and subjected to four classes of local alignment, 882,608 particles were further subjected to four classes of local alignment, and 1,522,468 were selected. Each of the last cround of 2D classification was subjected to the final RELION refinement. The final model was obtained using the CLC version of RELION. The final cryo-EM map of rNaᵥ₁.5C was used as an initial model. Each of the last five iterations was further subjected to four classes of local alignment and 3D classification with an angular search step at 3.75 Å. After
combining particles from the best 3D classes and removing duplicate particles, 570,843 particles were subjected to per-particle CTF estimation by GCTF followed by Bayesian polishing. The polished particles were selected to last round three class multi-reference 3D classification. The best class containing 267,595 particles was auto-refined and sharpened in Relion3.0. Local resolution was estimated by ResMap in Relion3.0. A diagram illustrating our data processing is presented in Supplementary Fig. 2.

Model building and refinement. The structures of rat rNaV1.5, (PDB code: 6LZ0) and LqhIII (PDB code: 1FHI) were fitted into the cryo-EM density map in Chimera52. The model was manually rebuilt in COOT54 and subsequently refined in Phenix35. The model vs. map FSC curve was calculated by Phenix.mtrage. Statistics for cryo-EM data collection and model refinement are summarized in Supplementary Table 1.

**Molecular dynamics model.** The cryo-EM structure of rNaV1.5, LqhIII lacking DI-DII-DIII linkers is composed of three chains which correspond to DI-DIV. The MODELER software (ver. 9.22) was used to insert missing residues and sidechains within the polypeptide chains, followed by quick refinement using MD with simulated annealing36. Neutral N- and C-termini were used for the three polypeptide chains in our refined model of rNaV1.5c. N-termini from chains DI and DIII-IV were acetylated, and a neutral amino terminus (–NH2) was used for DI. Neutral carbonyl groups (–COOH) were used for all C-termini. Disulfide bonds linking residues 327–432, 909–918, and 1730–1744 were included in our model of the channel as they were present in the cryo-EM structure; however, no glycans were added to the protein. Charged N- and C-termini were used for LqhIII and disulfide bonds linking residues 12–65, 16–37, 23–47, and 27–49 were included.

**Molecular dynamics simulations.** Molecular models of rNaV1.5c/ LqhIII with and without the toxin were prepared using the input generator, Membrane Builder from CHARMM-GUI57-61, from CHARMM-GUI (Io et al., 2009). The rNaV1.5c/ LqhIII model was embedded in a hydrated DMPC bilayer, with approximately 150 mM NaCl. The protein was translated and rotated for membrane embedding using PPM server52. The lipid bilayer was assembled using the replacement method, and solvent ions were added at random positions using a distance-based algorithm. A periodic rectangular cell with approximate dimensions of 14 × 14 × 13 nm was used, which comprised ~240,000 atoms. The CHARMM36 all-atom force field62-65 was used in conjunction with the TIP3P water model66. Non-bonded fixes for backbone carbonyl oxygen atoms with Na+ and lipid head groups with Na+ were imposed. Electrostatic interactions were calculated using the particle-mesh Ewald algorithm69,70 and chemical bonds were constrained using the LINCS algorithm71. The energy of the system was minimized with protein position restraints on the backbone (4000 kJ/mol/nm²) and side chains (2000 kJ/mol/nm²), as well as lipid position and dihedral restraints (1000 kJ/mol/nm²) using 5000 steps of steepest descent. The simulation systems were pre-equilibrated using multistep desalting. The number of water molecules in the gate was counted without the toxin were prepared using the input generator, Membrane Builder from CHARMM-GUI. The positions of all atoms were then rendered using Visual Molecular Dynamics75. Analyses were performed using MDTraj74 and molecular visualizations were rendered using Visual Molecular Dynamics77.

**Data availability.** Data supporting the findings of this paper are available from the corresponding authors on request. A reporting summary for this Article is available as a Supplementary Information file. Structural Data are available from the Protein Data Bank under EMDB entry ID EMD-2262 and PDB entry ID 7K1B. Source data are provided with this paper.

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