

Access and Binding of Local Anesthetics in the Closed Sodium Channel^S

Iva Bruhova, Denis B. Tikhonov, and Boris S. Zhorov

Department of Biochemistry and Biomedical Sciences, McMaster University, Hamilton, Ontario, Canada (I.B., D.B.T., B.S.Z.); and Sechenov Institute of Evolutionary Physiology and Biochemistry, Russian Academy of Sciences, St. Petersburg, Russia (D.B.T.)

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ABSTRACT

Local anesthetics (LAs) are known to bind Na⁺ channels in the closed, open, and inactivated states and reach their binding sites via extracellular and intracellular access pathways. Despite intensive studies, no atomic-scale theory is available to explain the diverse experimental data on the LA actions. Here we attempt to contribute to this theory by simulating access and binding of LAs in the KcsA-based homology model of the closed Na⁺ channel. We used Monte Carlo minimizations to model the channel with representative local anesthetics *N*-(2,6-dimethylphenylcarbamoylmethyl)triethylammonium (QX-314), cocaine, and tetracaine. We found the nucleophilic central cavity to be a common binding region for the ammonium group of LAs, whose aromatic group can extend either along the pore axis (vertical binding mode) or to the III/IV domain interface (horizontal binding mode). The vertical mode was earlier predicted for the open channel, but only the horizontal mode is consistent with mutational data on the closed-channel block.

To explore hypothetical access pathways of the permanently charged QX-314, we pulled the ligand via the selectivity filter, the closed activation gate, and the III/IV domain interface. Only the last pathway, which leads to the horizontal binding mode, did not impose steric obstacles. The LA ammonium group mobility within the central cavity was more restricted in the vertical mode than in the horizontal mode. Therefore, occupation of the selectivity-filter DEKA locus by a Na⁺ ion destabilizes the vertical mode, thus favoring the horizontal mode. LA binding in the closed channel requires the resident Na⁺ ion to leave the nucleophilic central cavity through the selectivity filter, whereas the LA egress should be coupled with reoccupation of the cavity by Na⁺. This hypothesis on the coupled movement of Na⁺ and LA in the closed channel explains seemingly contradictory data on how the outer-pore mutations as well as tetrodotoxin and μ -conotoxin binding affect the ingress and egress of LAs.

Voltage-gated Na⁺ channels are involved in the generation of action potentials in excitable cells. Structural determinants of major physiological characteristics (i.e., ion selectivity, gating, and ligand binding) are localized in the α subunit (Catterall, 2000; Hille, 2001). It is composed of four homologous domains (I–IV) quasi-symmetrically arranged around the central pore (Sato et al., 2001). Each domain contains six transmembrane segments (S1–S6), a pore-forming region (P), and the intracellular N and C ends (Guy and Seetharamulu, 1986; Denac et al., 2000). The P region contains the selectiv-

ity filter formed by a circular motif of highly conserved Asp, Glu, Lys, and Ala residues called the DEKA locus (Terlau et al., 1991; Heinemann et al., 1992). The selectivity filter separates the outer pore, which is targeted by tetrodotoxin (TTX), saxitoxin, and conotoxins, from the inner pore, which is targeted by local anesthetics (LAs) and other small cationic drugs (Zhorov and Tikhonov, 2004). Na⁺ channels exist in the closed, open, fast-inactivated, and several slow-inactivated states. Structural features of the first three states are thought to be analogous to corresponding X-ray structures of K⁺ channels (Doyle et al., 1998; Zhou et al., 2001; Jiang et al., 2003a). A structural model of the outer pore in the slow-inactivated channel has been recently proposed (Tikhonov and Zhorov, 2007).

Blockade of Na⁺ channels by LAs is a well known and medically important phenomenon. The mechanisms of action and structure-function relationships of LAs have been intensively studied more than 30 years (Catterall, 2000; Ruetsch et al., 2001; Nau and Wang, 2004; Fozzard et al., 2005).

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ABBREVIATIONS: TTX, tetrodotoxin; LA, local anesthetic; QX-222, 2-[(2,6-dimethylphenyl)amino]-*N,N,N*-trimethyl-2-oxoethaniminium; QX-314, *N*-(2,6-dimethylphenylcarbamoylmethyl)triethylammonium; CTX, conotoxin; MCM, Monte Carlo minimization.

Inside the Na⁺ channel pore, LAs bind state-dependently. Various LAs, including etidocaine (Ragsdale et al., 1994), cocaine (Wright et al., 1998), and cocaethylene (Xu et al., 1994), block the closed Na⁺ channel. Three terms are used to describe this phenomenon: the closed-channel block, tonic block, or resting-channel block. In this article, we use the first term. Under high-frequency stimulation, the blocking effect increases. This phenomenon is called frequency-dependent or use-dependent block. Experiments on the state-dependent action of LAs demonstrated that LAs generally have a stronger potency in the open or inactivated channels compared with the closed channels (Hille, 1977; Chahine et al., 1992; Ragsdale et al., 1994; Xu et al., 1994; Wright et al., 1997). Two major hypotheses have been proposed to explain these effects. According to the “modulated receptor” hypothesis (Hille, 1977), LAs can bind to the closed, open, and inactivated channel states but with different affinity. According to the “guarded receptor” hypothesis (Starmer et al., 1984), LAs have a permanent affinity to their receptor, whereas the block depends on the frequency and duration of the channel gating, which imposes an energy barrier for the ingress and egress of the ligand.

Intensive mutational studies (Ragsdale et al., 1994; Wright et al., 1998; Li et al., 1999; Yarov-Yarovoy et al., 2001, 2002) demonstrated that the binding site for LAs is located in the inner pore, between the activation gate and the selectivity filter and involves residues in domains I, III, and IV. Residues in domain IV (see Methods for the residue labeling scheme), particularly Phe⁴¹¹⁵ and Tyr⁴¹²² were found critical for the use-dependent block by LAs (Ragsdale et al., 1994). It is noteworthy that mutations of these residues had qualitatively different effects on the closed-channel and use-dependent block. Closed-channel block by tetracaine requires a hydrophobic residue in position 4i15, whereas Tyr⁴¹²² is important for the use-dependent block (Li et al., 1999). In agreement with these results, use-dependent block by cocaine is sensitive to mutations F⁴ⁱ¹⁵C and Y⁴ⁱ²²C (O’Leary and Chahine, 2002). In case of the closed-channel block by cocaine, mutation Y⁴ⁱ²²K had no effect, whereas mutations F⁴ⁱ¹⁵K and N⁴ⁱ²⁰K decreased the cocaine potency by 2- and 3-fold, respectively (Wright et al., 1998). In contrast, the cocaine potency in the inactivated channel decreased in the same mutants, 6-, 21.3-, and 27.4-fold, respectively, indicating that cocaine blocks the channel state-dependently (Wright et al., 1998). The mechanisms responsible for qualitatively different effect of mutations on the closed-channel block and use-dependent block are unresolved.

The access of LAs to and from the open pore is conceptually clear. Upon channel activation, LAs enter through the open intracellular gate and move along the ion-conducting hydrophilic pathway to their binding site in the inner pore. Understanding of the hydrophilic pathway was advanced by the X-ray structures of K⁺ channels that have been crystallized in the closed (Doyle et al., 1998) and open states (Jiang et al., 2002a,b, 2003a,b; Long et al., 2005a,b). In the open conformation, the inner helices bend and form a wide vestibule, which is readily accessible for drugs from the cytoplasm.

In the closed state, the straight inner helices converge at the cytoplasmic side and obstruct ion permeation (Doyle et al., 1998). The bulky LA molecules are unlikely to pass through the closed gate. It was suggested that LAs reach the inner pore of the closed channel via a hydrophobic pathway

(Hille, 1977). Because lipid-soluble compounds can cross the membrane, they are active when applied both extra- and intracellularly (Nettleton and Wang, 1990). Quaternary analogs of LAs, such as QX-222 and QX-314, cannot cross the membrane and therefore are used as probes for exploring the access pathways into the channels. When applied intracellularly, the quaternary LAs can block only the open Na⁺ channels (Hille, 1977). Unlike neuronal (Na_v1.2) and skeletal-muscle (Na_v1.4) channels, cardiac (Na_v1.5) Na⁺ channels are blocked by quaternary LAs applied extracellularly (Alpert et al., 1989; Qu et al., 1995). The mutation F⁴ⁱ¹⁵A of Na_v1.5 effects both intra- and extracellular block by QX-314, suggesting that extracellular and intracellular LAs target the same site in the inner pore (Qu et al., 1995).

Several studies suggest that extracellularly applied LAs can reach their binding site through the outer pore. TTX binding to the outer pore of Na_v1.5 reduced the external QX-314 access and recovery after the use-dependent block (Qu et al., 1995). Mutations of Na_v1.4 also showed the involvement of the DEKA locus in the external access and binding of QX-314 (Sunami et al., 1997). Residue Tyr^{1p51}, which is critical for isoform-specific TTX action, affects the external access of QX-222 (Sunami et al., 2000). Mutation S^{4p49}L, one position upstream the DEKA locus, enhanced the external QX-314 access and dissociation (Sasaki et al., 2004). Cys substitutions of T^{3p48}, F^{3p49}, and S^{4p49} are accessible by the externally applied Cd²⁺ (Yamagishi et al., 1997) and sulfhydryl reagents (Yamagishi et al., 1997; Struyk and Cannon, 2002). All these data support a hypothesis that the outer pore of Na⁺ channels may be permeable to LAs and compounds of similar size.

Another set of data suggests the involvement of the inner helices in the hydrophobic pathway of LAs. The mutation of Thr⁴ⁱ⁸ in Na_v1.5 to Val, which is a native residue in Na_v1.2, reduced block by external QX-314 (Qu et al., 1995). Mutation I⁴ⁱ¹¹A created an external pathway for LAs in the brain (Ragsdale et al., 1994) and muscle Na⁺ channels (Wang et al., 1998; Sunami et al., 2001). This mutation does not affect ion permeation or binding of toxins to the outer pore (Sunami et al., 2001). A μ -CTX mutant R13N binds to the outer pore of Na⁺ channels but does not fully block the current. External QX-222 increased block of the I⁴ⁱ¹¹A Na_v1.4 mutant in the presence of the μ -CTX mutant R13N, suggesting that the outer pore is not the access route for the LA (Sunami et al., 2001). Furthermore, the mutation I⁴ⁱ¹¹C in Na_v1.5 enhanced binding and facilitated escape of cocaethylene from the closed channel (O’Leary et al., 2003).

The coexistence of two pathways for charged LAs, one through the outer pore and another through the domain interface, was previously discussed (Lee et al., 2001). Based on the models of Na_v1.4 and Na_v1.5, we earlier hypothesized that the hydrophobic pathway goes through the “sidewalk” along the IIIP helix between the inner helices IIIS6 and IVS6 (Tikhonov and Zhorov, 2005a; Tikhonov et al., 2006). In this study, we elaborate this hypothesis by docking various ligands and computationally exploring various hypothetical access pathways.

In the absence of X-ray structures of voltage-gated Na⁺ channels, their homology models have been proposed to explain the effects of LAs (Lipkind and Fozzard, 2005; Tikhonov et al., 2006; Tikhonov and Zhorov, 2007). Unlike previous studies that were focused on explaining the use-

dependent block, the goal of this study was to explore access pathways and binding of LAs in the closed Na⁺ channel. We addressed the following questions. What are the binding modes of QX-314, cocaine, and tetracaine in the closed channel? What are the energetically possible ingress and egress routes for LAs in the closed channel? What is the impact of Na⁺ ions in the pore on the access and binding of LAs in the closed channel? Our study predicted that the LA molecules adopt different orientations in the closed and open channels. The energetically most favorable pathway for the LA molecule to and from the closed channel was the interface between domains III and IV, whereas the selectivity filter and the closed activation gate imposed large energy barriers for LAs. Finally, we propose that the occupancy of the pore with a Na⁺ ion affects both access and binding of LAs. These results provide a novel structural interpretation for a large body of experimental data.

Materials and Methods

Our model of the closed rNa_v1.5 included the outer helices, P loops, and inner helix, the backbones of which were built according to the X-ray structure of KcsA (Doyle et al., 1998). Extracellular linkers between the P-loops and transmembrane helices were not modeled. The ascending limbs of P-loops including the selectivity filter were taken from our earlier model (Tikhonov and Zhorov, 2005a). The sequences of P loops and inner helices of K⁺ and Na⁺ channels were aligned as proposed earlier (Zhorov and Tikhonov, 2004). The outer helices, which align unambiguously between Na⁺ and Ca²⁺ channels, were aligned with KcsA as proposed by Huber et al. (2000). Ionizable residues were treated as neutral except for the DEKA-locus ionizable residues, which were charged. The EEDD locus of the outer pore is proposed to bind four Na⁺ ions between domains (Tikhonov and Zhorov, 2007). To make the locus more flexible, we did not use here the interdomain Na⁺ bridges but neutralized the four acidic residues by four protons. Because the EEDD locus is more than 15 Å away from the binding site of LAs in the inner pore, this approximation would bias the LAs access through the selectivity filter, but is unlikely to affect the ligand binding in the inner pore. Cocaine and tetracaine models were protonated based on the data that protonated cocaine blocks Na⁺ channels (Nettleton and Wang, 1990). The DEKA locus was loaded either by an explicit water molecule or a Na⁺ ion, which were initially constrained, respectively, to the Asp and Lys or Asp and Glu side chains. In the subsequent Monte Carlo-energy minimization (MCM) trajectories, the constraints were removed, but neither the water molecule nor the Na⁺ ion moved away from the DEKA locus.

A universal scheme (Zhorov and Tikhonov, 2004) was used for residue labeling (Table 1). It includes the domain number (1–4), symbols “i,” “p,” or “o” for the inner helix, P loop, or the outer helix, and the relative number from the N end of a transmembrane helix or from the DEKA-locus positions 1p50, 2p50, etc. For example, Met²¹¹ designates methionine in the domain II inner helix, 11 positions downstream from its N end.

The MCM method (Li and Scheraga, 1987) was used to search for energetically favorable three-dimensional structures in the space of torsional angles, bond angles of ligands, and positions and orientations of roots of free molecules. α Carbons were constrained to the template structure by pins. A pin is a flat-bottom energy function that allows atoms to deviate penalty-free up to 1 Å from the template and imposes a penalty of 10 kcal mol⁻¹ Å⁻¹ for larger deviations.

The atomic charges of ligands were calculated by the AM1 method (Dewar et al., 1985) using the MOPAC program. Hydration energy was calculated by the implicit-solvent method (Lazaridis and Karplus, 1999). Hydration of the lipid-facing residues in the outer helices is unrealistic, but because these residues are not involved in the

binding of LAs and because the channel folding does not change in our model, the hydration of S5s has a minimal impact on our results.

Nonbonded interactions were calculated using the AMBER force field (Weiner et al., 1984, 1986). The ligands contained both heavy atoms and hydrogens. The protein was modeled in the united-atom approximation, but an all-atom model was also used to assess cation- π interactions of the ligand with aromatic residues. A cutoff of 8 Å was used for nonbonded interactions, but those involving ions and ionized groups were calculated without a cutoff. The homology model of the ligand-free closed channel was MC-minimized until 2000 consecutive energy minimizations did not decrease the energy of the apparent global minimum found. For docking of LAs, the multi-MCM protocol (Bruhova and Zhorov, 2007; Tikhonov and Zhorov, 2007) was employed as follows. First, 60,000 starting points with random positions, orientations, and conformations of the ligand were seeded within a cube having 14-Å edges. The cube center was placed in the cavity center. The size of the cube was chosen to ensure sampling in the central pore and in the domain interfaces. Each starting point was optimized in an MCM trajectory of 10 steps to remove steric overlaps with the channel. Three hundred lowest-energy structures found at this stage were further MC-minimized in longer trajectories to refine the protein-ligand complexes. These trajectories were terminated after 1000 consecutive steps did not decrease the energy. All the complexes within 10 kcal/mol from the apparent global minimum were analyzed.

Access pathways for LAs were explored by pulling QX-314 along a specific direction with the step of 0.5 Å (Tikhonov and Zhorov, 1998). At each position, the energy was MC-minimized. For QX-314 pulled through the selectivity filter or the closed activation gate, the pulling direction coincides with the pore axis. For the ligand pulled between domains III and IV, the pulling direction is parallel to helix IIIP. The ligand was pulled by the *para*-carbon of the benzene ring. The ligand flip-flop was prevented as described previously (Bruhova and Zhorov, 2007). For each pathway, three independent trajectories were calculated, and the ligand-channel energies at each position were averaged. All calculations were performed using the ZMM program (<http://www.zmmsoft.com>).

Results

LA Binding. Three structurally different LAs were chosen in our study: QX-314, which is a permanently charged derivative of lidocaine, cocaine with bulky fused rings, and tetracaine with an elongated hydrophobic tail attached to its benzene ring (Fig. 1). The LAs were docked in the closed Na_v1.5 model whose DEKA locus was loaded by either a water molecule (H₂O-DEKA model) or a Na⁺ ion (Na⁺-DEKA model).

LAs in the H₂O-DEKA Model. Figs. 2, A and B, show multiple randomly generated starting points. The multi-MCM of the starting points yielded ensembles of 50 most favorable complexes for each of the three compounds studied. In each complex, the ligand occurred in the inner pore. In most cases, the ammonium group was found in the central cavity, near the focus of P helices (Figs. 2C and 3A). The DEKA-locus side chains strongly contribute to the binding of all three LAs because the electrostatic attraction of the ammonium group to Asp^{1p50} and Glu^{2p50} overweighs the repulsion from Lys^{3p50}. Furthermore, carbonyl groups of residues in positions p47 through p50 of all four repeats also stabilize the ligand via electrostatic interactions.

The ligand orientation greatly varied between the complexes. All three LAs have an elongated shape. Let us define the long axis of a ligand as a line drawn between the ammonium nitrogen and the *para*-carbon of the benzene ring. Three energetically favorable binding modes of LAs inside

TABLE 1

Sequences of ion channels and effect of mutations in Na⁺ channels

In bold type are residues whose mutations are described under the footnotes in the rightmost column. Residues whose mutations affect closed-channel block are shaded. Residues whose mutations affect access of LAs are underlined. The standard segment name includes the domain number and symbols S5, S6, or P, for respective transmembrane segment or membrane-reentering P-loop. An alternative segment name, which is used in residue labels, includes the domain number and symbols o, p, and i for the outer helices, inner helices, and P loops, respectively. The absolute numbers (#) of the first residue in a segment are shown for the following sequences: Na_v1.2 (SCN2A_RAT), Na_v1.4 (SCN4A_RAT), and Na_v1.5 (SCN5A_RAT). Relative numbers of residues are shown above the aligned sequences. For the transmembrane segments, the relative numbers are counted from the beginnings of the respective KcsA helices as seen in the X-ray structure (Doyle et al., 1998). Relative numbers of residues in P loop are counted from the DEKA locus, the residues of which are assigned number 50 to avoid negative numbers for residues in P helices.

Channel	Segment		#	1	11	21			
KcsA	M1	o	14	ALHWRAAGAA	TVLLVIVLLA	GSYLAVLAER			
Na _v 1.5	IS5	1o	242	IQSVKKLADV	MVLTVFCLSV	FALIGLQLFM			
	IIS5	2o	825	GNSVGALGNL	TLVLAIIVFI	FAVVGMLQFVG			
	IIIS5	3o	1321	VGAIPSIMNV	LLVCLIFWLI	FSIMGVNLFSA			
	IVS5	4o	1644	MMSLPALFNI	GLLLFLVMFI	YSIFGMANFA			
				33	41	51			
KcsA	P	p	59	LITYPRAL	WWSVETATTV	GYGDLYPV			
Na _v 1.4	IP	1p	383	YDTFSWAF	LALFRLMTQD	<u>Y</u> WENLFQL	a		
	IIP	2p	738	MNDFHFSF	LIVFRILCGE	WIETMWDC	a		
	IIIP	3p	1220	YDNVGLGY	LSLLQVATFK	GWMDIMYA	a		
	IVP	4p	1512	FETFGNSI	ICLFEITTS	<u>A</u> GWDGLLNP	b		
Na _v 1.5	IP	1p	356	FDSFAWAF	LALFRLMTQD	<u>C</u> WERLYQQ	b		
	IIP	2p	884	MMDFHFAF	LIIFRILCGE	WIETMWDC			
	IIIP	3p	1404	FDNVGAGY	LALLQVATFK	GWMDIMYA			
	IVP	4p	1696	FQTFANSM	LCLFQITTS	<u>S</u> A GWDGLLSP	c		
Pore-facing position						**	***		
Sidewalk-facing position in IIIP						*	***		
				1	11	21	31		
KcsA	M2	i	86	LWGRVLAVVV	MVAGITSFGL	VTAALATWV	GREQ		
Na _v 1.2	IS6	1i	399	KTYMIFVVLV	IFLGSFYLIN	LILAVVAMAY	EEQN		
	IIS6	2i	957	TMCLTVFMMV	<u>M</u> VIGNLVVLN	LFLALLSSSF	SSDN		
	IIIS6	3i	1447	LYMYLYFVIF	IIFGSFFTLN	LFIGVIIDNF	NQQK		
	IVS6	4i	1750	SVGIFFVFSY	<u>I</u> IISFLVVVN	<u>M</u> YIAVILENF	SVAT		
Na _v 1.4	IS6	1i	415	KTYMIFVVI	IFLGSFYLIN	LILAVVAMAY	AEQN		
	IIS6	2i	770	AMCLTVFLMV	MVIGNLVVLN	LFLALLSSSF	SADS		
	IIIS6	3i	1262	LYMYLYFVIF	IIFGSFFTLN	LFIGVIIDNF	NQQK		
	IVS6	4i	1565	SIGICFFCSY	<u>I</u> IISFLVVVN	<u>M</u> YIAIILENF	NVAT		
Na _v 1.5	IS6	1i	388	KIYMIFFMLV	IFLGSFYLVN	LILAVVAMAY	EEQN		
	IIS6	2i	916	SLCLLVFLV	MVIGNLVVLN	LFLALLSSSF	SADN		
	IIIS6	3i	1446	LYMYIYFVVF	IIFGSFFTLN	LFIGVIIDNF	NQQK		
	IVS6	4i	1748	AVGILFFTTY	<u>I</u> IISFLVVVN	<u>M</u> YIAIILENF	SVAT		
Pore-facing position						*	*	**	*
Sidewalk-facing positions IIIS6							*	*	
Sidewalk-facing positions IVS6						*	**	*	

^a Mutations at the DEKA locus (D^{1p50}A, E^{2p50}A, and A^{4p50}D) allow external block by QX-222 and QX-314 and accelerate the recovery from block by internal QX-314 (Sunami et al., 1997). Mutation Y^{1p51}C enables the external block by QX-222 (Sunami et al., 2000).

^b Mutation C^{1p51}Y reduces the external QX-314 block (Sunami et al., 2000).

^c Mutation S^{4p49}L accelerates recovery of hNa_v1.5 from internally applied QX-314 and enhances tonic block by mexiletine and QX-314 (Sasaki et al., 2004).

^d Mutation Y¹¹¹⁷A increases the affinity of etidocaine to the closed channel (Yarov-Yarovoy et al., 2002).

^e Mutations M²¹¹A, N²¹⁵A, and V²¹⁸A increase closed-channel block by etidocaine (Yarov-Yarovoy et al., 2002).

^f Mutations L³¹⁹A, N³²⁰A, and I³²³A decrease affinity of inactivated channels to LAs but do not decrease the closed-channel block (Yarov-Yarovoy et al., 2001). Mutations L³¹⁹A and N³²⁰A increase the closed-channel block by lamotrigine and etidocaine. Mutation I³²³A increases the affinity for lamotrigine but not etidocaine to the closed channel.

^g Mutation I⁴¹¹A accelerates the channel recovery after use-dependent block and allows external QX-314 block (Ragsdale et al., 1994). Alanine mutations of Phe⁴¹⁵, Asn⁴²⁰, and Tyr⁴²² significantly decrease use-dependent block by etidocaine (Ragsdale et al., 1994). In the Na_v1.3 channel, closed-channel block requires a hydrophobic residue at position 415, whereas use-dependent block by tetracaine requires an aromatic residue 415 and Tyr⁴²² (Li et al., 1999).

^h Mutation of Leu³¹⁹ has no effect on closed-channel block by bupivacaine (Nau et al., 2003).

ⁱ Mutation C⁴¹⁸T allow the external QX-222 block (Sunami et al., 2000). Mutant I⁴¹¹A allows block by the external QX-314 (Wang et al., 1998) and QX-222 (Sunami et al., 2001). Benzocaine and etidocaine are less potent in the F⁴¹⁵A mutant (Wang et al., 1998). Lysine substitutions show that Tyr⁴²² is important for the use-dependent block, but not the closed-channel block by cocaine, whereas Phe⁴¹⁵ and Asn⁴²⁰ are important for both the use-dependent and closed-channel block (Wright et al., 1998).

^j T⁴¹⁸V reduces external QX-314 block; F⁴¹⁵A inhibits the binding of both external and internal QX-314 (Qu et al., 1995). I⁴¹¹C accelerates recovery and facilitates closed-state untrapping of cocaine (O'Leary et al., 2003). Mutation Y⁴²²C significantly reduces use-dependent block by cocaine (O'Leary and Chahine, 2002).

the channel can be categorized as vertical, horizontal, and angular. In the first two modes, the ligand long axis is either parallel (Fig. 2D) or perpendicular (Fig. 2F) to the pore axis. In the angular mode, the ligand long axis is at $45 \pm 10^\circ$ to the pore axis (Fig. 2E). QX-314 (Fig. 2) and cocaine (Fig. 3) can adopt all the three binding modes. Tetracaine (Fig. 4) adopts only the horizontal mode because its long hydrophobic tail does not fit in the closed pore.

Energetics of the binding modes is given in Table 2. The vertical mode is stabilized by rings of residues in positions i22 and i15. In the vertical mode, the hydrophobic bundle of residues i22 from all four repeats anchors the LA benzene ring (Figs. 2D and 3C). In particular, Phe²¹²² and Tyr⁴¹²² π -stack to opposite faces of the drug's benzene ring. In some cases, the ligand's carbonyl or the secondary amine group H-bonds with the OH group of Tyr⁴¹²². Residue Phe³¹²² at the bottom of the binding pocket interacts with the benzene ring in the edge-to-face manner (Burley and Petsko, 1985; Singh and Thornton, 1985). Ile¹¹²² also contributes to the bottom of the binding pocket, whereas Phe⁴¹¹⁵ is upper in the ligand side.

In the horizontal binding mode, the aromatic ring protrudes in the III/IV domain interface. Despite the fact that this orientation was not biased in our calculations, practically no low-energy complexes were found in which the aromatic ring of LAs were protruding in the other (I/II, II/III, or IV/I) domain interfaces. Only in the case of tetracaine were a few complexes found in which the aromatic ring protruded in the other interfaces, but their ligand-channel energy was weak. The horizontal binding mode is stabilized mainly by π -stacking interactions with Phe⁴¹¹⁵ (Table 2). Because of the horizontal ligand orientation, residues in the cytoplasmic halves of S6s practically do not contribute to the ligand binding. The long tail of tetracaine reaches Trp³⁰¹⁸ that contributes -0.7 kcal/mol to ligand-channel energy. In the an-

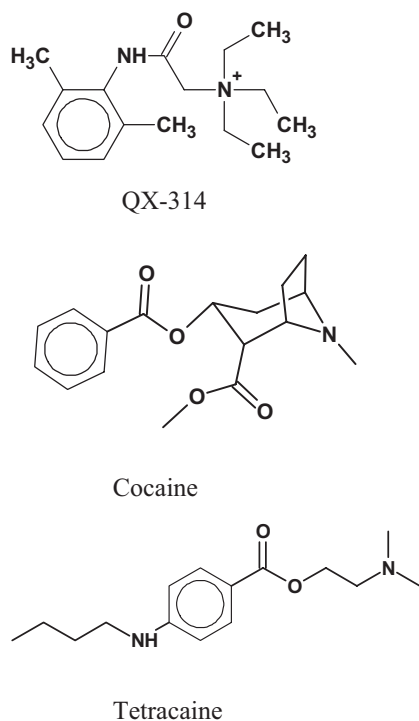


Fig. 1. Chemical structures of LAs that are considered in this study.

gular binding mode, which is intermediate between the vertical and horizontal modes, the benzene ring of the ligands binds between Phe⁴¹¹⁵ and Tyr⁴¹²².

In general, the horizontal binding mode has more preferable electrostatic energy than the vertical one because of a stronger attraction of the ligand's ammonium group to the nucleophilic C-ends of the P-helices (Table 2). This is compensated by a larger dehydration cost of the horizontal mode versus the vertical. Interactions with the S6s are more important for the vertical mode. Contributions of the key binding determinants in IVS6 to the vertical and horizontal modes are different: in the vertical mode, Tyr⁴¹²² contributes more than Phe⁴¹¹⁵, but only Phe⁴¹¹⁵ contributes significantly to the horizontal mode. The above characteristics of the binding modes are similar for the three LA docked in this study, suggesting that other LAs may bind similarly in the closed Na⁺ channel.

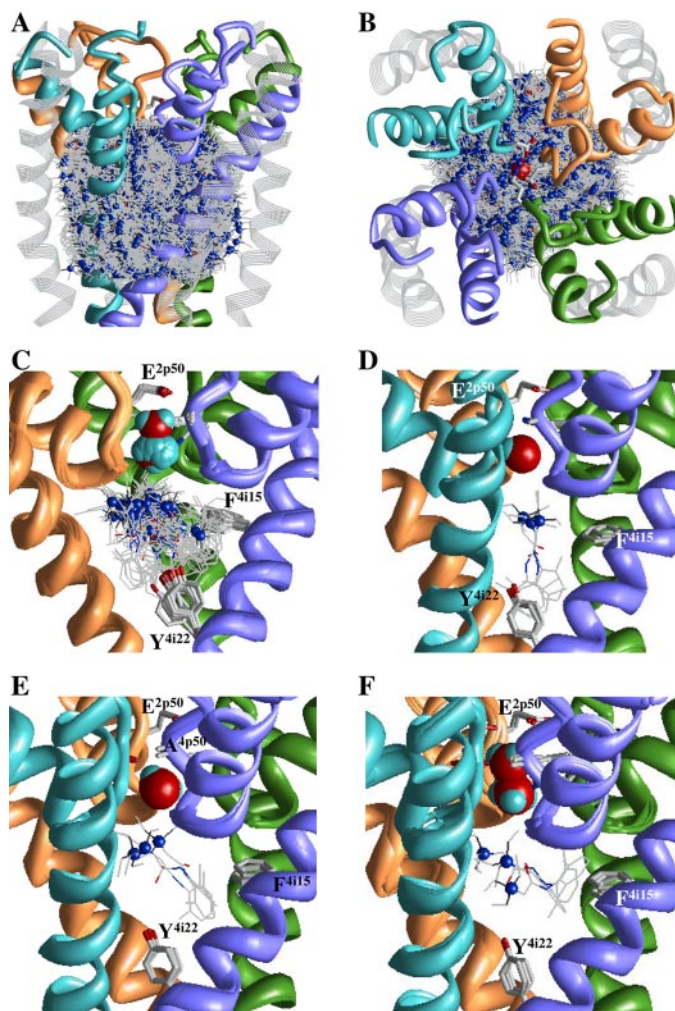


Fig. 2. Searching for energetically favorable binding modes of QX-314 in the closed Na_v1.5 with a water molecule in the selectivity filter. The inner helices and P-loops of domains I, II, III, and IV are colored cyan, orange, green, and violet, respectively. The outer helices are shown as gray strands. The water molecule is space-filled. The side chains of residues in the DEKA locus as well as conserved Phe⁴¹¹⁵ and Tyr⁴¹²² are shown as sticks. The ligand is wire-frame, with the ammonium nitrogen shown as a blue sphere. A and B, side and top views at the superposition of starting points. For clarity, only 6000 of the 60,000 starting points are shown. C, fifty lowest-energy binding modes. D, E, and F, superposition of the lowest energy vertical, angular, and horizontal binding modes, respectively. For clarity the outer helices are not shown in C-F.

LAs in the Na⁺-DEKA Model. Earlier, we predicted that the binding mode of lidocaine in the open Na⁺ channel depends dramatically on whether a Na⁺ ion or a water molecule populates the DEKA locus (Tikhonov et al., 2006; Tikhonov and Zhorov, 2007). In this study, we placed a Na⁺ ion in outer pore of the closed Na_v1.5 in a position equivalent to that of the K⁺ ion position 4 in the outer pore of K⁺ channels (Tikhonov and Zhorov, 2007). In this position, the Na⁺ ion counterbalances the excessive negative charge in the DEKA locus, which becomes less attractive for the pore-bound cations than in the H₂O-DEKA model.

Cocaine and QX-314 were docked by the multi-MCM method. In the majority of the 50 lowest-energy structures collected for each ligand, the ammonium group is displaced from the selectivity filter in the cytoplasmic direction due to electrostatic repulsion from the Na⁺ ion in the DEKA locus. In the vertical binding mode, the repulsion energy between the ammonium group and the Na⁺ ion is as strong as 13.0

kcal/mol. Because of the limited space in the central cavity, the drugs had a tendency to adopt the horizontal binding mode with the aromatic ring protruding in the III/IV domain interface (Fig. 3B). In this mode, the ligand ammonium group and the Na⁺ ion are more distant from each other than in the vertical mode. As a consequence, the repulsion between the Na⁺ ion and the ligand in the horizontal binding mode varies from 1.5 to 8 kcal/mol, which is much weaker than in the vertical mode (Table 2). In the lowest-energy complex, the cocaine ester group is attracted by the Na⁺ ion, whereas the benzene ring π -stacks with Phe⁴¹¹⁵ (Fig. 3D). In some complexes, the Na⁺ ion relocated from the starting position to the extracellular face of the DEKA locus (Fig. 3D), the level equivalent to position 2 in K⁺ channels (Tikhonov and Zhorov, 2007). This relocation decreased the Na⁺-ligand repulsion.

Thus, in the Na⁺-DEKA and H₂O-DEKA models, the LAs adopt the horizontal and vertical binding modes, respectively, and the LA binding energy significantly depends on the occupancy of the DEKA locus.

The above docking experiments of our study have been performed with the united-atom model of the channel, which is insensitive to cation- π interactions because atomic charges at aromatic carbons are close to zero. To assess a possibility of cation- π interactions, we created and MC-minimized an all-atom model of the closed channel and docked QX-314 in the vertical and horizontal modes with constraints imposing either π -stacking or cation- π interactions between QX-314 and Phe⁴¹¹⁵. After MC-minimizing the complexes, constraints were removed and the second round of MC minimizations provided structures shown in Supplemental Fig. S1. The structures with cation- π interactions were found less prefer-

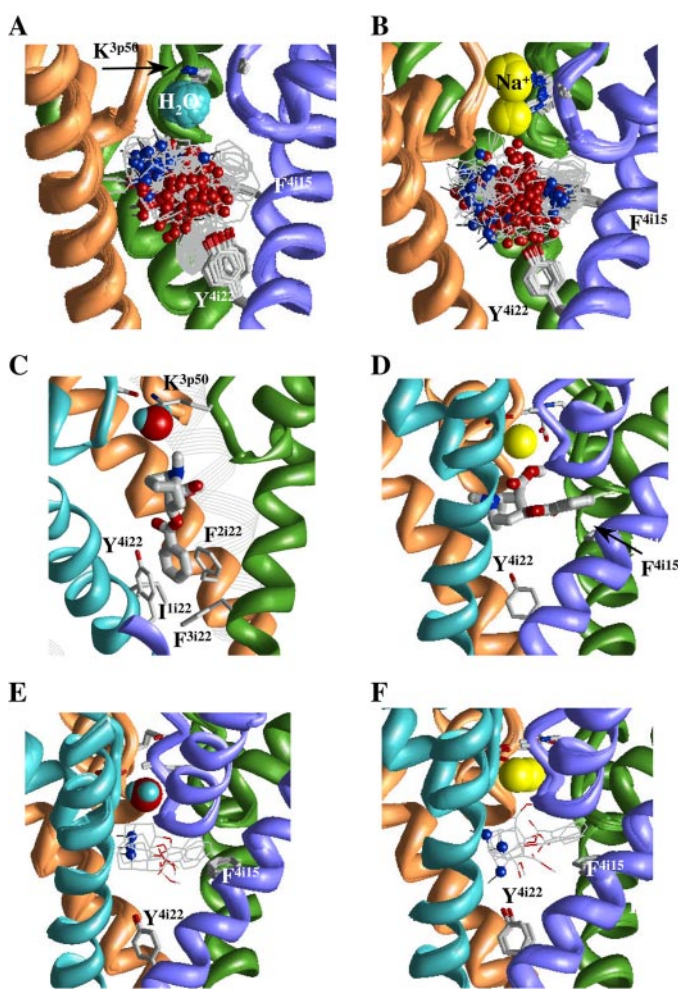


Fig. 3. Searching for energetically favorable binding modes of cocaine in the closed Na_v1.5 with a water molecule (A, C, and E) or Na⁺ ion (B, D, and F) in the DEKA locus. For clarity the outer helices are not shown. A and B, fifty lowest-energy binding modes in the H₂O-DEKA and Na⁺-DEKA models, respectively. C, a vertical binding mode of cocaine. The ligand's aromatic group interacts with three aromatic residues in positions i22. D, the energetically most favorable binding mode of cocaine in the Na⁺-DEKA model. The oxygen atoms of cocaine approach the Na⁺ ion, whereas the aromatic ring of the drug extends to the sidewalk between helices IIIS6 and IVS6. E and F, the superposition of the energetically best horizontal binding modes.

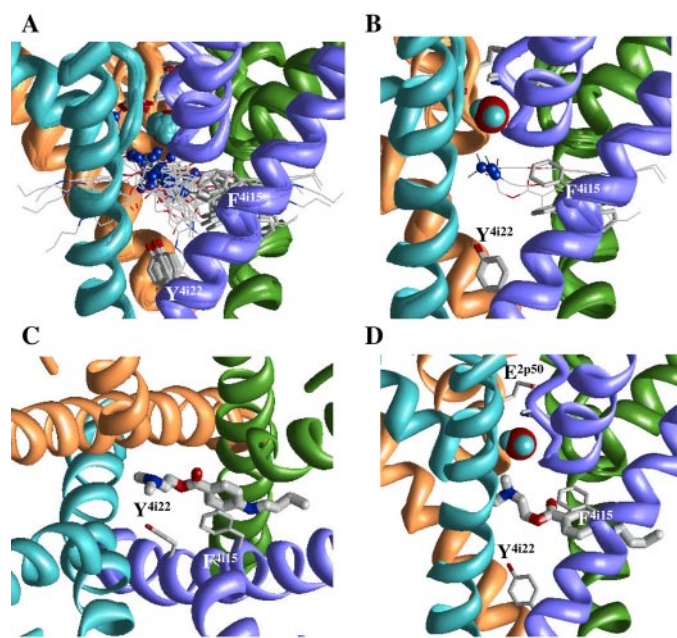


Fig. 4. Searching for energetically favorable binding modes of tetracaine in the closed Na_v1.5 channel with a water molecule in the DEKA locus. A, fifty lowest-energy complexes. In most of the complexes, the drug's aromatic ring protrudes in the III/IV domain interface. B, the superposition of three lowest-energy complexes representing the horizontal binding mode. C and D, the side and top views of the lowest-energy complex representing the horizontal binding mode. Note π -stacking of the drug's aromatic ring with Phe⁴¹¹⁵.

able than those with stacking interactions (Supplemental Table S1).

Access Pathways of QX-314. The extracellular access of LAs to the closed Na⁺ channels is discussed in several studies (Ragsdale et al., 1994; Qu et al., 1995; Sunami et al., 1997; Sunami et al., 2001) but it remains unclear whether LAs pass between the III/IV domain interface or through the selectivity filter. To address this, we pulled QX-314 through these two hypothetical pathways (Figs. 5 and 6). As a control, the ligand was also pulled through the closed activation gate (Fig. 5A). The vertical binding mode was chosen as a starting position for pulling QX-314 through the selectivity filter and the closed gate, whereas the horizontal mode was used as the starting point for pulling the ligand via the III/IV interface. None of the computed profiles have ligand-receptor energy better than in the starting point (Figs. 5C and 7). This agrees with the notion that the inner pore is the energetically preferable binding site for LAs.

Pulling QX-314 through the closed gate (Fig. 5A) imposes a van der Waals energy barrier of 60 kcal/mol (Fig. 5C), indicating that the closed gate is impassible. Two peaks of the energy profile at positions -10 and -17 Å are determined by repulsion between the ligand's ammonium group and residues in positions i22 and i26, which face the pore axis. Thus, as expected, the closed activated gate does not allow QX-314 to reach the central cavity from the cytoplasm.

Pulling QX-314 through the selectivity filter not loaded with a water molecule or a Na⁺ ion (Fig. 5B) results in the energy barrier of 33 kcal/mol, which is caused by steric clashes and unfavorable desolvation (Fig. 5C). Residues in positions p49 and p50 form the tightest rings. In particular, Lys^{3p50}, Gly^{2p49}, and Ser^{4p49} contribute to the large energy barrier. Residues in positions p51, p52, and p53 do not impose energy barriers but attract the ligand.

Pulling QX-314 through the III/IV domain interface (Fig. 6) did not cause high-energy barriers (Fig. 7). Residues of the IIIP-helix interact with the ligand all along the pathway. The ligand-channel energy weakens as the ligand leaves the central cavity and loses electrostatic attractions from the DEKA

locus and nucleophilic C ends of P helices. Residues Thr⁴ⁱ⁸ and Ile⁴ⁱ¹¹ contribute only weakly to the drug-channel energy at the drug-binding site (position 0), but their contributions increase along the pathway. The most significant interaction of QX-314 with Ile⁴ⁱ¹¹ corresponds to the rightmost part of the profile, indicating that Ile⁴ⁱ¹¹ affects the drug access much stronger than the drug binding in the inner pore. Residue Thr⁴ⁱ⁸, which is far from the pore axis, faces the III/IV interface. The experimentally observed effect of mutations in this position on the external drug access can be explained by direct interaction with the ligand entering the domain interface. It should be noted that both Thr⁴ⁱ⁸ and Ile⁴ⁱ¹¹ interact with the drug at the rightmost part of the pathway (Fig. 7) and concertedly attract the ligand at this part where the ligand-channel attraction energy is rather small, whereas the desolvation cost increases. Therefore changes in the size and hydrophobicity in positions 4i8 and 4i11 can significantly affect access of ligands through the domain interface. Thus, the computed energy profile through the III/IV domain interface agrees with the hydrophobic access pathway concept.

Comparison of the energy profiles calculated in the H₂O-DEKA and Na⁺-DEKA models show that the Na⁺ ion significantly destabilizes QX-314 in position 0 and along the access pathway (Fig. 7B). In position 0, the difference between the drug-channel energies of the two models is 5 kcal/mol. The difference in positions 3 to 6 is much larger because in this narrow part of the pathway the Na⁺ ion and the ligand's ammonium nitrogen approach each other as close as 6 Å. Besides this electrostatic repulsion, the Na⁺ ion does not significantly affect the ligand trajectory and contributions of individual residues to the ligand-channel energy.

Residue Phe⁴ⁱ¹⁵ plays a key role in the access pathway of QX-314 (Figs. 6 and 7D). At position 0, Phe⁴ⁱ¹⁵ faces the pore and π -stacks with the benzene ring of QX-314. As QX-314 is pulled between domains III and IV, Phe⁴ⁱ¹⁵ reorients to let QX-314 pass. The reorientation is associated with an energy barrier at positions 2.5 to 3.5 Å. The barrier is much higher in the Na⁺-DEKA model than in the H₂O-DEKA model.

TABLE 2

Energetically favorable binding modes of LAs in the closed Na_v1.5 with the DEKA locus loaded by either H₂O or Na⁺

Shown are Ligand-receptor energy, its components, and contributions (kcal/mol) from individual segment and specific residues. For cocaine and QX-314, the lowest-energy structure for each binding mode is characterized. The entropy contributions were not taken into account, which is a standard approximation in homology modeling. These contributions would partially compensate the large enthalpy of ligand-channel interactions. The Na⁺-DEKA model energy is not shown because it is much weaker than in the H₂O-DEKA model. The Na⁺-DEKA model with tetracaine was not explored.

Ligand-Channel Energy and Its Components	Cocaine			QX-314			Tetracaine Horizontal H ₂ O
	Vertical H ₂ O	Horizontal		Vertical H ₂ O	Horizontal		
		H ₂ O	Na ⁺		H ₂ O	Na ⁺	
	<i>kcal/mol</i>						
Total	−40.6	−38.2	−36.2	−39.3	−39.0	−31.3	−35.2
van der Waals	−24.4	−26.1	−31.5	−23.3	−21.9	−22.7	−27.1
Desolvation	13.2	23.3	28.1	8.7	14.1	14.2	14.4
Electrostatics	−29.4	−35.5	−32.8	−24.8	−30.8	−22.7	−22.4
Segments ^a							
P loops	−19.9	−23.3	−30.7	−18.8	−24.6	−25.0	−20.7
All S6s	−19.8	−14.9	−10.8	−19.7	−13.6	−14.1	−12.3
Key determinants of action ^b							
Phe ⁴ⁱ¹⁵	−0.2	−3.9	−2.9	−1.0	−2.2	−2.4	−2.1
Tyr ⁴ⁱ²²	−3.2	−0.6	−0.4	−3.5	−0.4	−0.6	0.0

^a The energy values include both the side-chain and backbone contributions to ligand binding.

^b The energy values include only the side-chain contributions to ligand binding.

Discussion

Understanding molecular determinants of state-dependent action of LAs, which are widely used clinically and in experiments, are of obvious importance. Residues Phe⁴ⁱ¹⁵ and Tyr⁴ⁱ²² have long been known as critical determinants of the use-dependent block. LAs in the vertical but not in horizontal mode can simultaneously bind to both Phe⁴ⁱ¹⁵ and Tyr⁴ⁱ²², which are separated by two helical turns of IVS6. Several structural models of Na⁺ channels visualized the vertical binding mode and analyzed contacts of LAs with LA-sensing residues (Lipkind and Fozzard, 2000, 2005; Fozzard et al., 2005; Tikhonov et al., 2006; McNulty et al., 2007; Tikhonov and Zhorov, 2007).

It is well known that effects of mutations on the block of the open/inactivated and closed channels are different (Hille, 1977; Chahine et al., 1992; Ragsdale et al., 1994; Xu et al., 1994; Wright et al., 1997; Nau et al., 2003), but molecular determinants of this difference were unclear. Here we systematically simulated binding of structurally different LAs in

the closed-channel model. The chosen LAs contain an ammonium group and a benzene ring but essentially vary in size and chemical structure. Because the central cavity in the closed channel is rather small, it was unclear how these ligands interact with ligand-sensing residues. Furthermore, because occupation of the DEKA locus in the closed state is unknown, we populated the locus by either a water molecule or a Na⁺ ion. Unbiased docking using a powerful multi-MCM search resulted in ensembles of complexes in which QX-314 and cocaine adopted vertical, horizontal, or intermediate (angular) binding modes. In contrast, only the horizontal binding mode was found for tetracaine, because the vertical mode of this elongated drug does not fit in the confined central cavity of the closed channel. The horizontal binding mode of tetracaine was also recently proposed by another group (Scheib et al., 2006).

Ragsdale and coauthors (Li et al., 1999) systematically studied state-dependent action of tetracaine on Na_v1.3 and its mutants and demonstrated that frequency-dependent

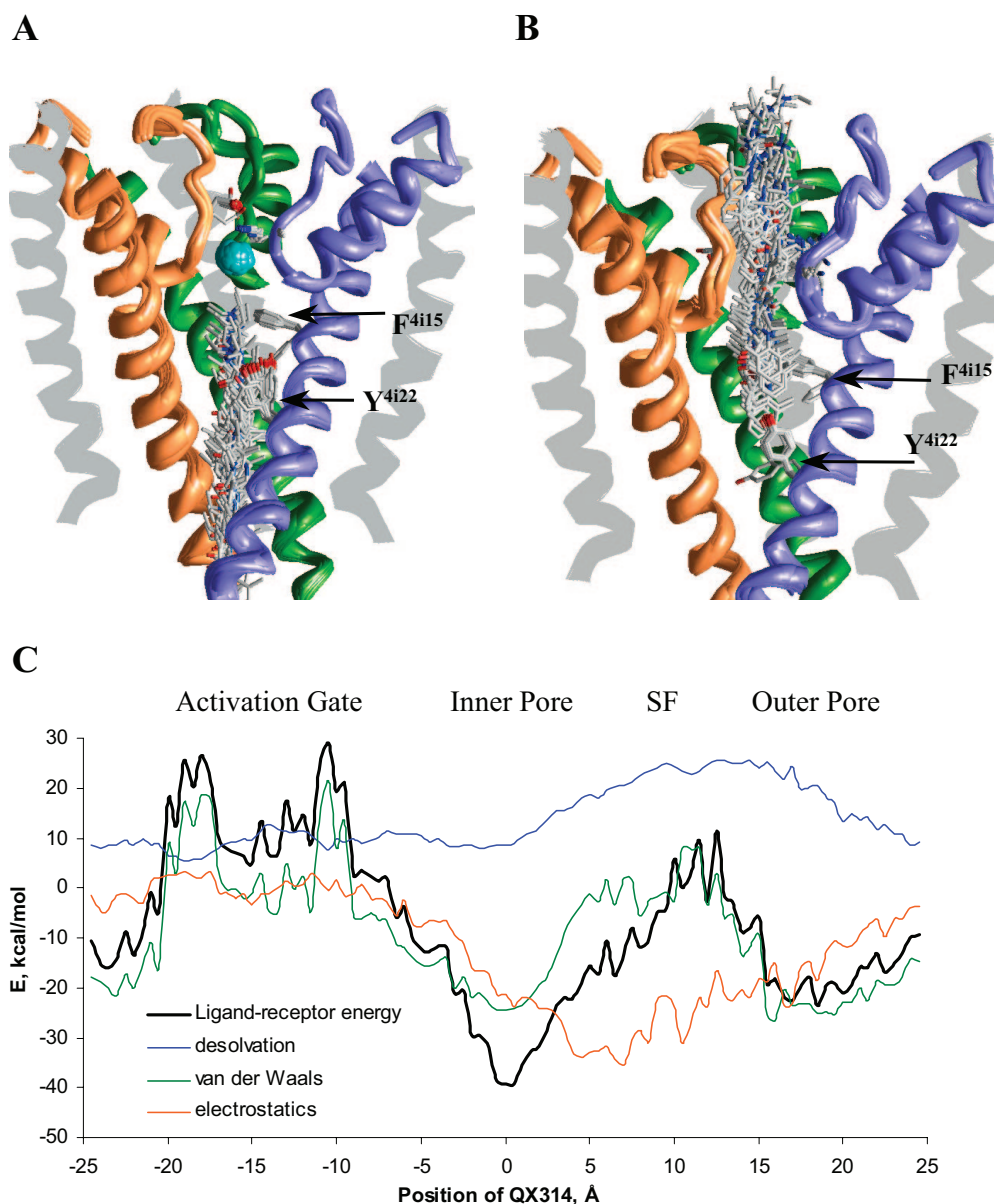


Fig. 5. Exploring hypothetical access pathways of QX-314. A and B, superposition of the MC-minimized structures obtained for QX-314 pulled out of the pore through the closed activation gate (A) and the selectivity filter (B). QX-314, DEKA residues, Phe⁴ⁱ¹⁵ and Tyr⁴ⁱ²² are shown as sticks. For clarity, domain I is not shown. C, the MC-minimized energy profiles showing the total drug-channel energy and its components. From the starting position 0, which corresponds to the lowest-energy vertical binding mode of QX-314 (Fig. 2D), the ligand was pulled at 0.5-Å steps either through the selectivity filter (positions 0 to 25) or through the activation gate (positions 0 to -25).

block requires an aromatic residue (Phe, Tyr, or Trp) in position 4i15, while the closed-channel block requires a hydrophobic residue in this position. Furthermore, the frequency-dependent block required Tyr⁴ⁱ²², while the resting-channel block was possible with Ser, Ala, Cys, Ile, or Phe in this position. These data support our conclusion that tetracaine blocks the closed channel in the horizontal mode. In-

deed, Phe⁴ⁱ¹⁵ contributes -2.1 kcal/mol, whereas Tyr⁴ⁱ²² did not have a noticeable contribution (Table 2).

Docking of QX-314 and cocaine did not predict a single binding mode. However, experimental data suggest that the horizontal binding mode in the closed channel is preferable for these drugs and possibly other LAs. Indeed, mutation Y⁴ⁱ²²K had no effect on the closed-channel block of rNa_v1.4 by cocaine, whereas mutations F⁴ⁱ¹⁵K lowered the LA affinity 2-fold (Wright et al., 1998). The hNa_v1.5 mutation I⁴ⁱ¹¹C weakly affects the use-dependent block by cocaine, whereas Y⁴ⁱ²²C had the strongest effects (O'Leary and Chahine, 2002). Thus, residues in the C-terminal half of IVS6 affect predominantly the use-dependent block, whereas residues at the N-terminal half affect mainly the closed-channel block. Mutation N⁴ⁱ²⁰K also decreased cocaine binding 3-fold (Wright et al., 1998), but this residue does not face the pore in our model and probably affects the ligand binding allosterically. Indeed, mutations of Asn⁴ⁱ²⁰ affect slow inactivation (Chen et al., 2006). Thus, experimental data support the notion that LAs bind to the closed channel in the horizontal mode, but they block the open channel in the vertical mode.

A recent study (Ahern et al., 2008) provides additional

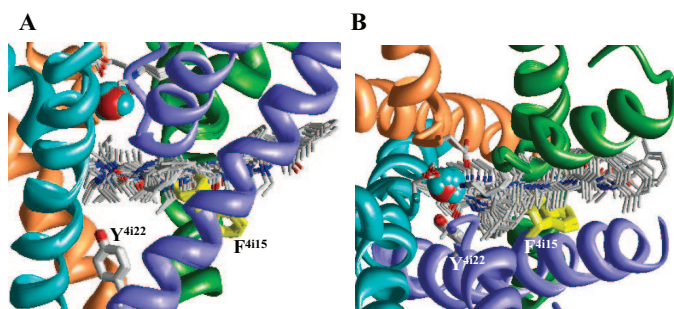


Fig. 6. Pulling QX-314 through the sidewalk between segments IIIP, IIIS6, and IVS6. A and B, the side and top views of the superposition of QX-314 pulled from the starting position 0, which corresponds to the horizontal binding mode found from random sampling (Fig. 2 F).

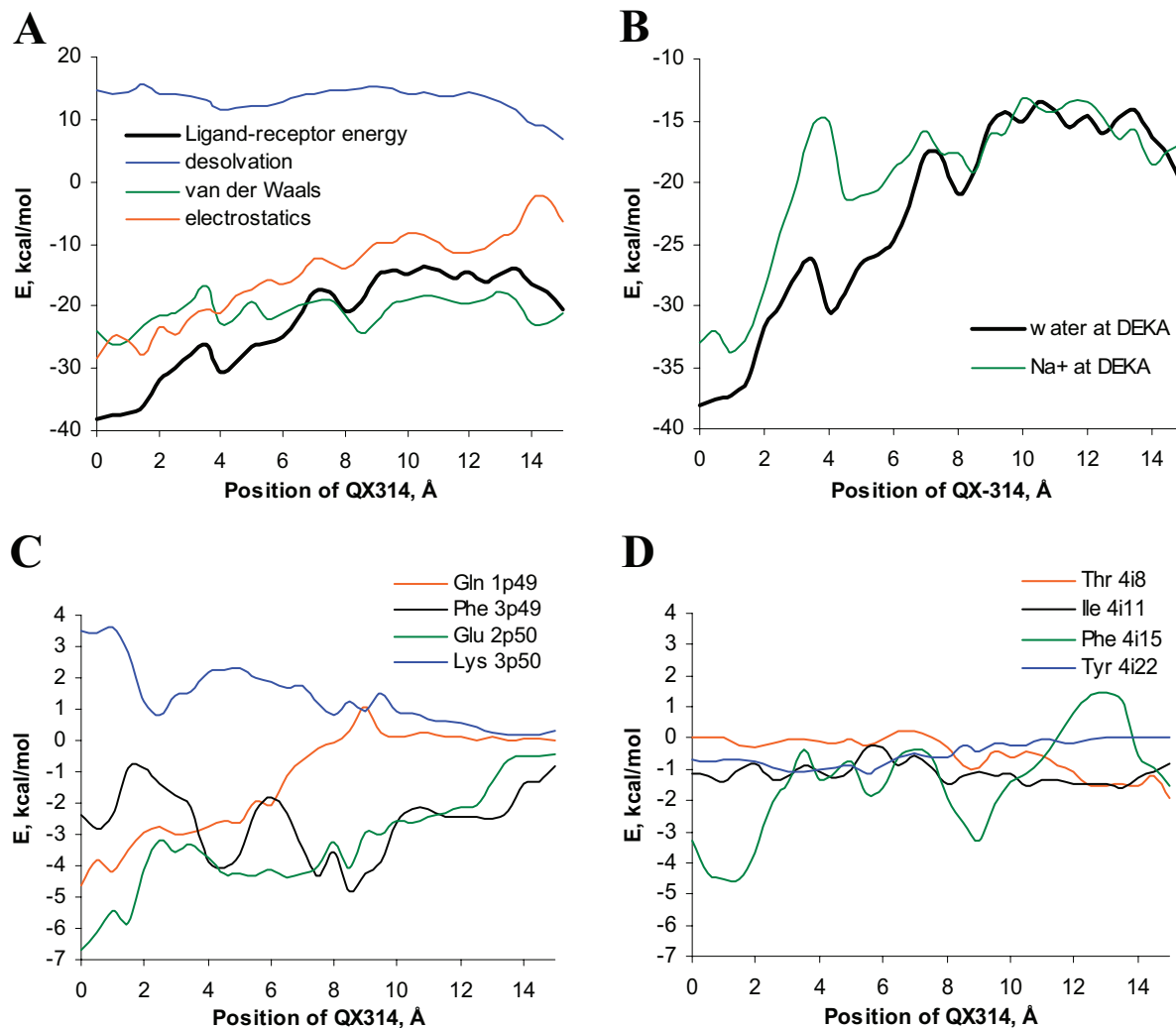


Fig. 7. The MC-minimized energy profile of QX-314 pulled through the sidewalk. A, the energy profile with total drug-channel energy and its components. Position 0 corresponds to the lowest-energy drug-channel complex, which was found in the multi-MCM search and which represents the horizontal binding mode (Fig. 2 F). The ligand was pulled at 0.5 Å steps. B, the energy profile of QX-314 pulled through the III-IV interface of the H₂O-DEKA and Na⁺-DEKA models. C and D, most significant energy contributions to the energy profile.

support for our model. The authors used a series of fluorinated derivatives of aromatic amino acids (Santarelli et al., 2007a,b) to systematically reduce the negative electrostatic potential at aromatic carbons of Phe⁴ⁱ¹⁵ and Tyr⁴ⁱ²². The disruption of π -cation interaction of Phe⁴ⁱ¹⁵ abolished the use-dependent block by lidocaine without affecting the closed-channel block. Fluorination of Tyr⁴ⁱ²² had no effect. This result strongly suggests that π -cation interactions takes place only at Phe⁴ⁱ¹⁵ and only in the use-dependent block. In our models of the use-dependent block (Tikhonov et al., 2006; Tikhonov and Zhorov, 2007) the amino-groups of LAs are located near the Phe⁴ⁱ¹⁵ and can form the π -cation contact. Contrary to this, in the horizontal binding mode (that we suggest to correspond to the closed channel block) Phe⁴ⁱ¹⁵ interacts with the LAs' aromatic rings rather than with the ammonium groups. Thus, despite the united-atom approximation, which is used in our model, is unable to reveal the π -cation interactions, our models agree with experimental data (Ahern et al., 2008).

The vertical binding mode, which probably corresponds to the use-dependent block, and the horizontal binding mode, which is proposed here for the closed-channel block, have similar structural features. In both binding modes, the amino-group of LAs is located below the selectivity filter in the region, which is attractive for cationic particles because of the cooperative effect of the P helices macrodipoles. However, in the horizontal mode the aromatic moiety of the drugs faces the domain interface and the channel lumen is not completely occluded especially by ligands with a small ammonium group (lidocaine). This raises a question about the mechanism of channel block. A recent study (McNulty et al., 2007) suggested that for asymmetrically bound LAs, the block is caused by a combination of electrostatic and hydrophobic effects. This mechanism is plausible and fully agrees with our models. Moreover, effects of charge and hydrophobicity on the action of sodium channel ligands were the subjects of our recent studies (Tikhonov and Zhorov, 2005b; Wang et al., 2006, 2007a,b). Sodium channel activators like batrachotoxin and veratridine bind in the sodium channel to the site, which significantly overlaps with the LA binding site. These drugs do not completely occlude the channel. The charged groups of these drugs do not face the pore. Rather, the drugs expose hydrophilic groups to the remaining lumen. As a result, the channel is conducting despite the presence of relatively large molecules in the pore. Introduction of positively charged residues in position 2i15 does not abolish channel conductance but inverses batrachotoxin action from activation to block.

LA Access through the III/IV Domain Interface. Previous studies proposed two possible access pathways of LAs to the inner pore of the closed Na⁺ channel: through the selectivity filter and through the domain interface (see Introduction). Residues, which when mutated affect the LA access, face either the pore axis or the domain interface (Table 1). Except for residues at the P-loop turn, which can face both pathways, all other positions face either the first or the second pathway. Both pathways were proposed to coexist (Lee et al., 2001). However, positive values of van der Waals energy at positions 10 to 13 of the pathway through the selectivity filter (Fig. 5C) indicate steric clashes of the LA molecule with the P-loop residues, which face the pore axis. In contrast, van der Waals energy component is negative all along the access pathway through the sidewalk (Fig. 7A). The

existence of this pathway readily explains mutational data indicating the involvement of residues in positions 4i8, 4i11, 4i15, and 4p49 in the extracellular access (Table 1). All these residues face the sidewalk and interact with QX-314 according to the energy profiles (Fig. 7, C and D). In particular, the Na_v1.5 mutation T⁴ⁱ⁸V, which increases the sidewalk hydrophobicity, renders the channel less sensitive to the extracellular block by LAs (Qu et al., 1995). Mutation I⁴ⁱ¹¹A in the muscle and brain Na⁺ channels makes them sensitive to extracellularly applied quaternary LAs (Ragsdale et al., 1994; Wang et al., 1998; Sunami et al., 2001). Furthermore, mutation I⁴ⁱ¹¹C facilitates escape of the trapped cocaethylene from Na_v1.5 (O'Leary et al., 2003). Our models explain the effects of these mutations: the bulky Ile⁴ⁱ¹¹ in the wild type Na_v1.4 prevents LAs to enter the pore, whereas smaller Ala⁴ⁱ¹¹ enables LAs to pass. A naturally occurring hNa_v1.5 mutation S^{4p49}L accelerates recovery from the block by internally applied QX-314, enhances the closed-channel block by mexiletine and QX-314, and attenuates the use-dependent block by mexiletine (Sasaki et al., 2004). The S^{4p49}L mutation was suggested to effect the LA access through the selectivity filter (Sasaki et al., 2004), but in our model S^{4p49} faces the sidewalk and contributes to LA binding.

The extracellular access of drugs is not an exclusive feature of Na⁺ channels. Mutations of various residues that face the III/IV domain interface affect binding of drugs to L-type Ca²⁺ channel (Hockerman et al., 1997). Permanently charged benzothiazepines (Hering et al., 1993; Seydl et al., 1993; Kurokawa et al., 1997) and dihydropyridines (Kass et al., 1991; Kwan et al., 1995) block the L-type Ca²⁺ channel from the extracellular side. Dihydropyridines were proposed to access Ca²⁺ channels via the III/IV domain interface (Yamaguchi et al., 2003). The recent X-ray structures of KcsA with hydrophobic cations show long hydrophobic moieties bound in the interface between S6, S5, and P helices, which is suggested to provide an alternative access pathway for drugs to the inner pore (Lenaus and Gross, 2008).

Our present model predicts that LAs access the closed Na⁺ channel via III/IV domain interface. However, because mutations of the outer-pore residues and toxin binding in the outer pore are also known to influence the LA access into the closed channel, the challenge is to explain these experimental observations in view of our model. We propose here that the above influence is mediated by Na⁺ ions that can occupy two nucleophilic sites, one in the DEKA locus and another in the cavity center. The antagonism between the Na⁺ ion in the DEKA locus and LA binding in the open channel was described before (Tikhonov and Zhorov, 2007). Our present calculations show that occupation of the DEKA locus by a Na⁺ ion also affects LA binding in the closed channel (Table 2) and hinders the LAs access to the pore via the III/IV domain interface (Fig. 7B). In view of these data, mutations in the ascending limbs, which are likely to affect Na⁺ binding in the DEKA locus, should affect access of LAs through the domain interface. For example, mutations D^{1p50}A and E^{2p50}A at the DEKA locus make the Na_v1.4 channel sensitive to the external QX-222 (Sunami et al., 1997). The authors of this study proposed a straightforward interpretation of the experiments, according to which substitutions of large residues in the outer pore with small residues can provide a path for the external QX-222. We do not rule out this interpretation, but in view of our calculations, the above mutations would de-

stabilize Na⁺ in the DEKA locus and hence facilitate the LAs access through the domain interface.

In the X-ray structure of Kv1.2, the voltage-sensor domain does not block the sidewalk, but to enter it a ligand should interact with lipids and/or an auxiliary subunit of the channel. In the absence of experimental data, we cannot consider structural details of these interactions. A weakly hydrated tetraalkylammonium cation such as QX-314 has more chances to pass by hydrophobic residues of the sidewalk than strongly hydrated inorganic or ammonium cations. For example, energy calculations suggest that trialkylammonium-methanethiosulfonate reagents, which are used in the substituted cysteine-accessibility method, can reach those levels in the pore of ion channels, which are closed for hydrated inorganic ions (Tikhonov et al., 2004; Tikhonov and Zhorov, 2007).

The facts that TTX reduces the rates of extracellular QX-314 block and recovery from the use-dependent block support a hypothesis that LA access their binding site through the selectivity filter (Qu et al., 1995). Here, we propose an alternative explanation for these observations. It involves a Na⁺ ion in the central cavity (Fig. 8) at position, which is equivalent to the K⁺ binding site 5 seen in X-ray structures of K⁺ channels. Because a Na⁺ ion and LAs are obvious competitors for the nucleophilic cavity center, the Na⁺ ion should leave this site to allow LA binding (Fig. 8A). However, the

closed channel, which is blocked by TTX, does not provide the escape route for the Na⁺ ion. Residing the cavity, the Na⁺ ion prevents QX-314 access in the TTX-blocked closed channel (Fig. 8B). The TTX-induced reduction of the channel recovery from the use-dependent block can be also explained. In the absence of TTX, the escape of LAs through the sidewalk releases the nucleophilic site in the cavity center, and this site will be occupied by the incoming Na⁺ ion. In the TTX-blocked closed channel, the Na⁺ ion cannot access the cavity and replace the LA molecule. It would be an energetically nonpreferable situation if a LA molecule left the nucleophilic cavity center, and this region is not reoccupied by a Na⁺ ion.

In contrast to TTX, the μ -CTX mutant R13N binds to the outer pore of Na⁺ channels without completely blocking the Na⁺ current (Sunami et al., 2001). In the Na_v1.4 mutant I⁴¹¹A, externally applied QX-222 reaches the inner pore of the toxin-bound channel and further blocks the current. In view of our model, the incomplete block of the closed channel by the peptide toxin does not prevent exchange of Na⁺ and LAs in the central cavity (Fig. 8C). Thus, comparison of the effects of TTX and the μ -CTX mutant on the LA ingress and egress in the closed channel suggests that movement of inorganic and organic cations is coupled in the Na⁺ channel.

The residues whose mutations affect access of LAs in the closed channel are clustered in two groups: one group is localized at the sidewalk and another in the outer pore including the selectivity filter. The hypothesis of the coupled movement of organic and inorganic ions explains why mutations in both clusters affect the drug access through the sidewalk. However, it is difficult to explain how mutations of S6 residues could affect the drug access via the selectivity filter. Indeed, because the access-controlling S6 residues are not buried inside the protein, they are unlikely to allosterically affect the access via the outer pore. Furthermore, the energy profile at Fig. 7C suggests steric hindrances for an LA molecule at the selectivity-filter level, but even in the absence of the hindrances, the coupled-movement hypothesis suggests that the access to the closed channel through the selectivity filter is unlikely. Indeed, an LA molecule approaching the central cavity through the selectivity filter would encounter an electrostatic repulsion from the Na⁺ ion residing in the cavity center. The Na⁺ ion cannot escape through the closed activation gate, or through the hydrophobic sidewalks or through the outer pore, which is blocked by the approaching LA molecule. Therefore, displacement of the resident Na⁺ ion by the LA molecule coming through the selectivity filter is hardly possible.

P-loop channels have nucleophilic sites in the outer pore, the selectivity filter, and the central cavity. Numerous experimental data show that the interaction between cations at nucleophilic sites leads to the coupled movement in the multi-ion pores. For example, K⁺ ions are thought to move through the selectivity filter of K⁺ channels by sequentially occupying sites 1/3 and 2/4 (Zhou and MacKinnon, 2003). A recent experimental and theoretical study of the tetrabutylammonium-blocked KcsA shows that the position of the organic cation in the cavity depends on the selectivity-filter occupancy by K⁺ ions (Faraldo-Gómez et al., 2007). Here we applied the concept of the coupled movement to explain seemingly contradictory experimental data on access and binding of LAs in Na⁺ channels.

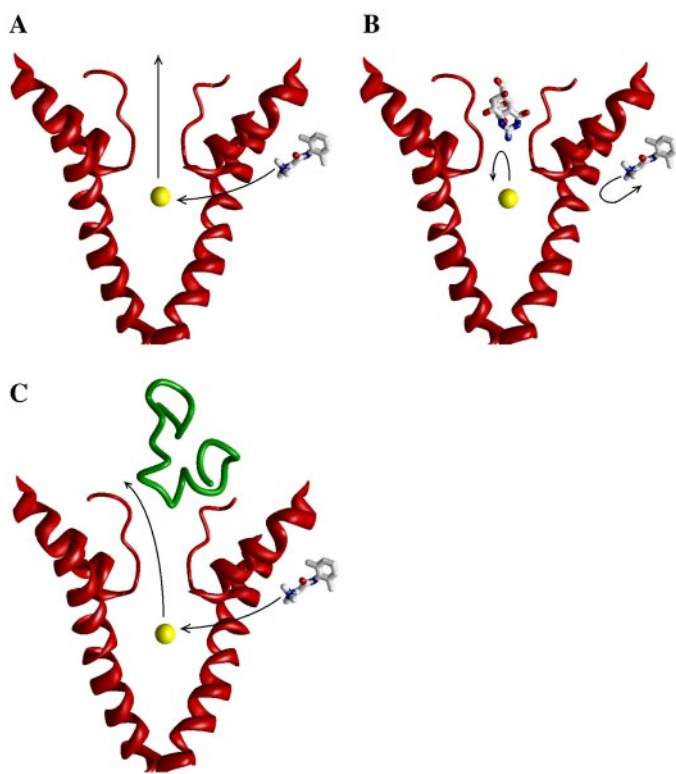


Fig. 8. A scheme of coupled movement of Na⁺ and a LA molecule in the closed Na⁺ channel. A, an LA molecule accessing the inner pore via the sidewalk repels the resident Na⁺ ion, which can move away via the unblocked outer pore and release the nucleophilic central cavity for binding of the LA ammonium group. B, when the outer pore is blocked by TTX, the Na⁺ ion in the central cavity lacks a hydrophilic escape route, remains in the central cavity, and prevents the LAs access. C, the μ -CTX mutant (green ribbon), binds in the outer pore but does not completely block the Na⁺ current. The Na⁺ ion can escape through the incompletely blocked outer pore and free the nucleophilic central cavity for the LAs binding. See Discussion for more details.

Modeling Limitations. In this work, we used a rather simple method of energy calculations. The atom-atom energy terms included van der Waals interactions, implicit solvent, and Coulomb's electrostatics but lacked an explicit term for cation- π interactions. We also neglected the entropy component of free energy. Despite the limited precision of coarse-grained methods, they are widely used for homology modeling of proteins, which, like sodium channels, have a limited sequence similarity with X-ray templates. Employment of high-precision time-consuming methods in such cases seems impractical because an assumption that the model and the template have similar backbone geometry influences results to a greater extent than approximations of energy calculations. To compensate these limitations, we generated a large number of starting points, which cover virtually all ligand-binding scenarios (Fig. 2, A and B). Subsequent MC-minimizations yielded ensembles of energetically possible binding scenarios, which include different structures (Figs. 2–4). Analysis of these ensembles allowed us to find particular binding modes, which agree with experimental data. For example, the vertical binding mode of QX-314 in the H₂O-DEKA model of the closed channel corresponds to the lowest-energy structure (Table 2), but the horizontal binding mode, which has a slightly higher energy, is consistent with a large number of experimental data. Taking into account a limited precision of energy calculations in the homology model, we concluded that the horizontal binding mode corresponds to the closed-channel block.

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Address correspondence to: Boris S. Zhorov, Department of Biochemistry and Biomedical Sciences, McMaster University, Hamilton, Ontario, Canada. E-mail: zhorov@mcmaster.ca