

A homology model of the pore domain of a voltage-gated calcium channel is consistent with available SCAM data

Iva Bruhova and Boris S. Zhorov

Department of Biochemistry and Biomedical Sciences, McMaster University, Hamilton, Ontario L8N 3Z5, Canada

In the absence of x-ray structures of calcium channels, their homology models are used to rationalize experimental data and design new experiments. The modeling relies on sequence alignments between calcium and potassium channels. Zhen et al. (2005. *J. Gen. Physiol.* doi:10.1085/jgp.200509292) used the substituted cysteine accessibility method (SCAM) to identify pore-lining residues in the Ca_v2.1 channel and concluded that their data are inconsistent with the symmetric architecture of the pore domain and published sequence alignments between calcium and potassium channels. Here, we have built K_v1.2-based models of the Ca_v2.1 channel with 2-(trimethylammonium)ethyl methanethiosulfonate (MTSET)-modified engineered cysteines and used Monte Carlo energy minimizations to predict their energetically optimal orientations. We found that depending on the position of an engineered cysteine in S6 and S5 helices, the ammonium group in the long flexible MTSET-modified side chain can orient into the inner pore, an interface between domains (repeats), or an interface between S5 and S6 helices. Different local environments of equivalent positions in the four repeats can lead to different SCAM results. The reported current inhibition by MTSET generally decreases with the predicted distances between the ammonium nitrogen and the pore axis. A possible explanation for outliers of this correlation is suggested. Our calculations rationalize the SCAM data, validate one of several published sequence alignments between calcium and potassium channels, and suggest similar spatial dispositions of S5 and S6 helices in voltage-gated potassium and calcium channels.

INTRODUCTION

Voltage-gated Ca²⁺ channels are involved in many physiological functions. The channels are targets for drugs used to treat arrhythmias, hypertension, myocardial ischemia, chronic pain, neuronal degeneration, and other disorders (Hockerman et al., 1997). The pore-forming α_1 subunit of Ca²⁺ channels is formed by a single polypeptide chain that contains four homologous repeats (Hockerman et al., 1997). Each repeat includes six transmembrane helices: the voltage sensor (S1–S4), the outer helices S5, and the pore-lining helices S6. Ion selectivity is controlled by a ring of four glutamates (EEEE), which are located at the membrane-reentering P loops between S5s and S6s. Ca²⁺ and Na⁺ channels have evolved from K⁺ channels (Anderson and Greenberg, 2001), and the three families of channels are believed to have a similar folding of the pore-forming domains and transmembrane topology of S5s and S6s.

In the absence of x-ray structures of voltage-gated Ca²⁺ and Na⁺ channels, their homology models based on x-ray structures of K⁺ channels in the closed and open states (Doyle et al., 1998; Jiang et al., 2003; Long et al., 2005, 2007) are used to explain experimental data and suggest new experiments. Homology modeling relies on the sequence alignments of K⁺ channels

with Na⁺ and Ca²⁺ channels. Various sequence alignments have been proposed (Huber et al., 2000; Zhorov et al., 2001; Lipkind and Fozzard, 2003; Stary et al., 2008). The models of the pore-forming domain (S5–P–S6) based on these alignments have different patterns of exposure of residues to the inner pore. The substituted cysteine accessibility method (SCAM) is used to define the architecture of ion channels (Karlin and Akabas, 1998). SCAM data are usually interpreted based on the cysteine orientation concept, according to which the application of a methanethiosulfonate (MTS) reagent to a channel with an engineered cysteine in a pore-facing position results in the chemical modification of the cysteine. If the ammonium group of the MTS-modified cysteine is exposed to the permeation pathway, it is expected to inhibit the current. Orientation of the engineered cysteine toward the lipid bilayer or the protein interior is believed to suppress ionization of the thiol group and its reaction with an MTS reagent. The current in such channels is expected to be similar to control channels with the native residue in the respective position.

The location of the activation gate in the *Shaker* channel was predicted using SCAM (Liu et al., 1997), and major conclusions from this study were later confirmed

Correspondence to Boris S. Zhorov: zhorov@mcmaster.ca

Abbreviations used in this paper: MC, Monte Carlo; MCM, MC minimization; MTS, methanethiosulfonate; MTSET, 2-(trimethylammonium)ethyl MTS; SCAM, substituted cysteine accessibility method.

© 2010 Bruhova and Zhorov. This article is distributed under the terms of an Attribution–Noncommercial–Share Alike–No Mirror Sites license for the first six months after the publication date (see <http://www.rupress.org/terms>). After six months it is available under a Creative Commons License (Attribution–Noncommercial–Share Alike 3.0 Unported license, as described at <http://creativecommons.org/licenses/by-nc-sa/3.0/>).

by the x-ray structures of K^+ channels. More recently, SCAM was used to identify pore-lining residues in the $Ca_v2.1$ channel (Zhen et al., 2005). The authors of this meticulous study interpret their results as inconsistent with known sequence alignments between K^+ and Ca^{2+} channels and suggest an asymmetric architecture of the inner pore of $Ca_v2.1$. This conclusion sheds doubts on published homology models of Ca^{2+} channels. The above interpretation of the SCAM experiments is apparently based on the cysteine orientation concept, which does not take into consideration the conformational flexibility of long side chains of MTS-modified cysteines. Due to this flexibility, the exposure of the MTS ammonium groups to the permeation pathway and hence the current-inhibiting effects of MTS may not correlate with the orientation of the C^α - C^β vector to the pore axis. Such possibilities can be explored by homology modeling of the channel with MTS-modified cysteines.

Here, we have built 44 models of $Ca_v2.1$ with MTS-modified engineered cysteines and used Monte Carlo (MC) minimizations (MCMs) to predict energetically possible orientations of MTS-modified side chains in the channels. We found that the residual current observed upon 2-(trimethylammonium)ethyl MTS (MTSET) application (Zhen et al., 2005) correlates with the predicted distance between the ammonium nitrogen in the MTS-modified cysteine and the pore axis, but does not correlate with the orientation of the C^α - C^β vector of the cysteine. Our results support the alignment between K^+ and Ca^{2+} channels, which was used in our

previous modeling studies, and suggest a similar disposition of transmembrane helices in the pore-forming domains of voltage-gated K^+ and Ca^{2+} channels.

MATERIALS AND METHODS

Homology models of $Ca_v2.1$ (CAC1A_RABIT) with MTSET-modified engineered cysteines were built using the x-ray structure of $K_v1.2$ (Long et al., 2005) and sequence alignment shown in Table I. The models include the outer helices (S5s), P loops, and inner helices (S6s). The ascending limbs of P loops, including the selectivity filter residues, were built using the $Na_v1.4$ model (Tikhonov and Zhorov, 2005) as a template. Those parts of the channel, which are far from the inner pore, were not modeled. Repeats I–IV were arranged clockwise when viewed extracellularly (Dudley et al., 2000). We use the terms “previous repeat” and “next repeat” to designate sequential neighbors of a mutated repeat (e.g., neighbors of repeat I are next repeat II and previous repeat IV).

All calculations were performed using the ZMM program (<http://www.zmmsoft.com>). Non-bonded energy was calculated using the AMBER force field (Weiner et al., 1984, 1986) with a cutoff distance of 8 Å. Hydration energy was calculated using the implicit solvent method (Lazaridis and Karplus, 1999). Electrostatic interactions were calculated using the distance-dependent dielectric function. Ionizable residues, including those in the selectivity filter, were modeled in their neutral forms (Lazaridis and Karplus, 1999), except for the acidic residues in the cytoplasmic side of KcsA-based closed-channel models. MTS-modified cysteines (designated mC) are incorporated in the ZMM program as nonstandard amino acids. The atomic charges of the mC residues have been calculated by the semi-empirical method AM1 (Dewar et al., 1985) using MOPAC. The charge of +1 proton charge unit is distributed among the ammonium nitrogen and surrounding methylene and methyl groups. ZMM program with MTS-modified cysteines was previously used in theoretical studies of glutamate-gated ion channels (Tikhonov, 2007). The MCM

TABLE I
Sequence alignment^a and effects of MTSET on channels with engineered cysteines^b

Channel	Segment	No.	1	11	21		
KcsA	M1	o	23	ALHWRAGAA	TVLLVIVLLA	GSYLAVLAER	
K _v 1.2	S5	o	322	KASMRELGLL	IFFLFIGVIL	FSSAVYFAEA	
Ca _v 2.1	IS5	1o	220	MKAMIPLLQI	GLLLFFAILI	FAIIGLEFYM	
	IIS5	2o	608	LNSMKSIISL	<u>LFLFLFIVV</u>	FALLGMQLFG	
	IIIS5	3o	1380	VNSLKNVFN	<u>LIVYMLFMFI</u>	<u>FAVVAVQLFK</u>	
	IVS5	4o	1695	VQSFKALPYV	<u>CLLIAM</u> LFFI	YAIIGMQVFG	
	Pore-facing position			* *	* *		
KcsA	M2	i	86	1	11	21	31
K _v 1.2	S6	i	385	LWGRLVAVVV	MVAGITSFGL	VTAALATWV	GREQ
Ca _v 2.1	IS6	1i	336	IGGKIVGSLC	AIAGVLTIAL	PVPVIVSNFN	YFYH
	IIS6	2i	690	TWNWLYFIPL	IIIGSFEMLN	<u>LVLGVLSGEF</u>	AKER
	IIIS6	3i	1485	MVFSIYFIVL	TLFGNYILLN	<u>VFLAIVDNL</u>	ANAQ
	IVS6	4i	1785	MEMSIFVYVY	<u>FVVFPEFVN</u>	<u>IFVALIIITF</u>	QEQG
	IVS6	4i	1785	EFAYFYVSE	<u>IFLCSFLMLN</u>	<u>LEVAVIMDNF</u>	EYLT
Na _v 1.4	IVS6	4i	1565	SIGICFFCSY	IIISFLIVVN	MYIAIILENF	NVAT
Pore-facing position				* *	**	**	
Cytoplasm-facing position						**	*** *

Data for $Na_v1.4$ are from (Sunami et al., 2004).

^aShown alignments of S5 (Huber et al., 2000) and S6 (Zhorov et al., 2001) segments were used to build the $Ca_v2.1$ models.

^bBold-oblique and underlined characters indicate positions where MTSET inhibits channels with engineered cysteines ≥ 30 and $<30\%$, respectively (Zhen et al., 2005).

method (Li and Scheraga, 1987) was used to optimize the models. During energy minimizations, C^α atoms were constrained to corresponding positions of the template using pins. A pin is a flat-bottom energy function, which allows an atom to deviate penalty-free up to 1 Å from the template and imposes a penalty of 10 kcal mol⁻¹ Å⁻¹ for deviations >1 Å.

Each model was MC minimized until 2,000 consecutive minimizations did not update the apparent global minimum. Then, the multi-MCM protocol (Bruhova and Zhorov, 2007) was used to predict all low energy orientations of the MTS-modified cysteine. The side chain torsions of the ¹³C residue were sampled from 60,000 random starting points. Each starting point was optimized in an MCM trajectory of 10 steps. The top 1,000 low energy conformations were further MC minimized for 1,000 steps. All conformations in which the interaction energy between the ¹³C residue and the rest of the channel did not exceed 4 kcal/mol from the apparent global minimum were analyzed. No specific energy terms were used for cation- π interactions, which were accounted for due to partial negative charges at the aromatic carbons (Bruhova et al., 2008). Further details of methodology can be found elsewhere (Bruhova and Zhorov, 2007; Tikhonov and Zhorov, 2007; Bruhova et al., 2008).

To validate our methodology, we predicted orientations of ¹³C²²³ in a cysteine transpeptidase, Sortase B. Comparison of the lowest energy orientation and the x-ray structure (Zong et al., 2004) shows that the ammonium nitrogen of ¹³C²²³ in the model is just 0.66 Å away from the experimental position (Fig. S1). We use a labeling scheme that is universal for P loop channels (Zhorov and Tikhonov, 2004). A residue label includes the repeat number (1–4), which may be omitted when the label is pertinent to all four repeats, segment type (*o*, outer helix; *p*, P loop; *i*, inner helix), and the residue relative number in the segment (Table I).

Online supplemental material

Table S1 presents the experimental data on the current inhibition by MTSET (Zhen et al., 2005). The predicted mobility of ammonium groups of respective ¹³C residues is shown in Fig. S1, which demonstrates the validity of our methodology of predicting conformations of ¹³C residues using the x-ray structure of a cysteine transpeptidase. Figs. S2–S4 display conformations and local environments of residues ¹³Cⁱ²³, ¹³Cⁱ²⁴, ¹³Cⁱ²¹, and ¹³Cⁱ²⁵ in respective open channels, and Fig. S5 shows energetically most favorable orientations of ¹³Cⁱ²⁴ residues in the KcsA-based model of the closed Ca_v2.1 channel. Table S1 and Figs. S1–S5 are available at <http://www.jgp.org/cgi/content/full/jgp.200910288/DC1>.

RESULTS

SCAM data and the cysteine-orientation concept

In the K_v1.2 template, vectors C^α - C^β in positions *i15*, *i18*, *i19*, *i22*, and *i23* direct to the pore axis, in positions *i16*, *i17*, *i20*, *i21*, *i25*, and *i29* direct to a neighboring S6 or S5, and in positions *i24*, *i26*, *i27*, *i28*, *i30*, and *i31* direct to the cytoplasm (Fig. 1). Assuming generally similar 3-D dispositions of S5s and S6s in Ca_v2.1 and K_v1.2, and correctness of the alignment in Table I, the SCAM data (Zhen et al., 2005), which are shown in Table S1, can be divided into two categories. In agreement with the cysteine orientation concept, the first-category data show that MTSET inhibits channels Cⁱ¹⁵, Cⁱ¹⁹, Cⁱ²³, Cⁱ²¹⁸, and Cⁱ⁴¹⁸, in which respective C^α - C^β vectors direct to the pore axis. The second-category data, which disagree with the cysteine orientation concept, include three groups of observations: (1) MTSET does not inhibit channels Cⁱ¹¹⁸, Cⁱ³¹⁸, and Cⁱ²², despite the fact that respective vectors C^α - C^β point to the pore axis; (2) MTSET inhibits channels C^{2o10} and C^{4o10}, despite the fact that respective positions are far from the pore; and (3) MTSET inhibits channels Cⁱ¹⁶, Cⁱ²¹⁰, Cⁱ³¹⁰, Cⁱ³²¹, Cⁱ²⁴, and Cⁱ²¹⁵, despite the fact that respective vectors C^α - C^β direct away from the pore axis.

Below, we describe the Ca_v2.1 models and consider the SCAM data in view of orientations of C^α - C^β vectors in ¹³C residues relative to the pore axis. Unless otherwise mentioned, the described orientation of an ¹³C residue corresponds to the lowest energy conformation.

MTS-modified cysteines in the pore-facing positions of S6s

The current of ¹³Cⁱ¹⁵ channels is inhibited by MTSET by 39.0–65.5%. According to our calculations, the ammonium groups of ¹³Cⁱ¹⁵ occur in the pore, where they are stabilized by electrostatic interactions with the nucleophilic C termini of P helices (residues *p47–p49*) and with the EEEE locus and interact with residues at the same level (*i15*) and lower levels *i18* and *i19* (Fig. 2, A and B).

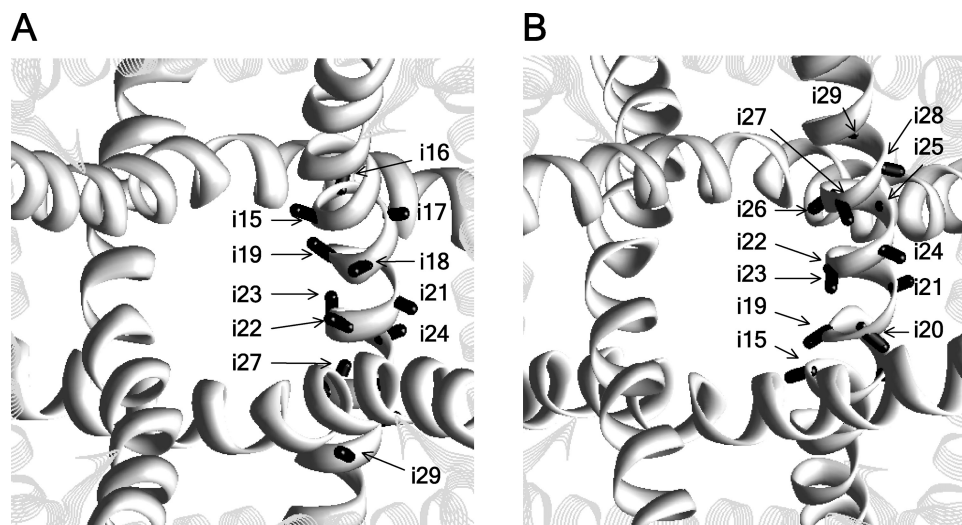


Figure 1. The extracellular (A) and cytoplasmic (B) views of the K_v1.2 x-ray structure, with the C^α - C^β bonds of positions *i15*–*i29* shown as sticks. The S5 and S6 helices are shown as strands and ribbons, respectively. The P loops are not shown for clarity.

MTSET inhibits channels C^{218} and C^{418} more than C^{118} and C^{318} . Calculations predict two orientations of mC^{118} side chains (Fig. 3). In the pore-facing orientations, the ammonium groups are attracted to the C ends of the P helices, the EEEE locus, and residues *i15* and *i19*. In the repeat interface orientation, the ammonium groups occur between S5 and next repeat S6. The energetically preferable orientations depend on the neighboring residues, which are distinct in different repeats (Table I). The pore orientation of mC^{218} is energetically most preferable. The side chain of mC^{418} is equally stable in the pore and interface IV/I, where it experiences cation- π interactions with F^{116} . The side chain of mC^{118} is most stable in interfaces I/II, where it interacts with L^{1010} , L^{2112} , and Y^{2116} . The side chain of mC^{318} is most stable in interface III/IV, where it experiences cation- π interactions with F^{3122} and F^{4112} and hydrophobic interactions with I^{3010} .

MTSET strongly inhibits C^{119} channels. The mC^{119} side chains are in the pore and orient either upward to the focus of P helices (Fig. 2 C) or downward (Fig. 2 D). The

latter orientation is more stable due to cation- π interactions with F^{2122} , F^{3122} , and F^{4122} . In both orientations, the mC^{119} ammonium groups are close to the pore axis.

Channels C^{122} are only weakly sensitive to MTSET, despite the fact that their respective C^α - C^β vectors direct to the pore axis. The mC^{122} side chains can adopt two orientations: to the pore, where they interact with a neighboring F^{122} , or to the repeat interfaces, where they interact with aromatic residues in positions *i12*, *i16*, or *i18* (Fig. 4). The pore-facing orientations of mC^{2122} , mC^{3122} , and mC^{4122} are energetically more preferable than the repeat interface orientations. The weak effect of MTSET on respective channels may be due to large hydrophobic residues *i19*, *i22*, and *i26* (Fig. 5) that would prevent the ionization of C^{122} and thus the reaction with MTSET.

MTSET inhibits the C^{123} channels by 43.6–87.7%. The ammonium groups of mC^{123} are oriented into the pore. For example, the pore orientation of mC^{1123} is stabilized by interactions with L^{1126} , F^{2122} , F^{4130} , F^{4122} , and I^{4126} (Fig. S2).

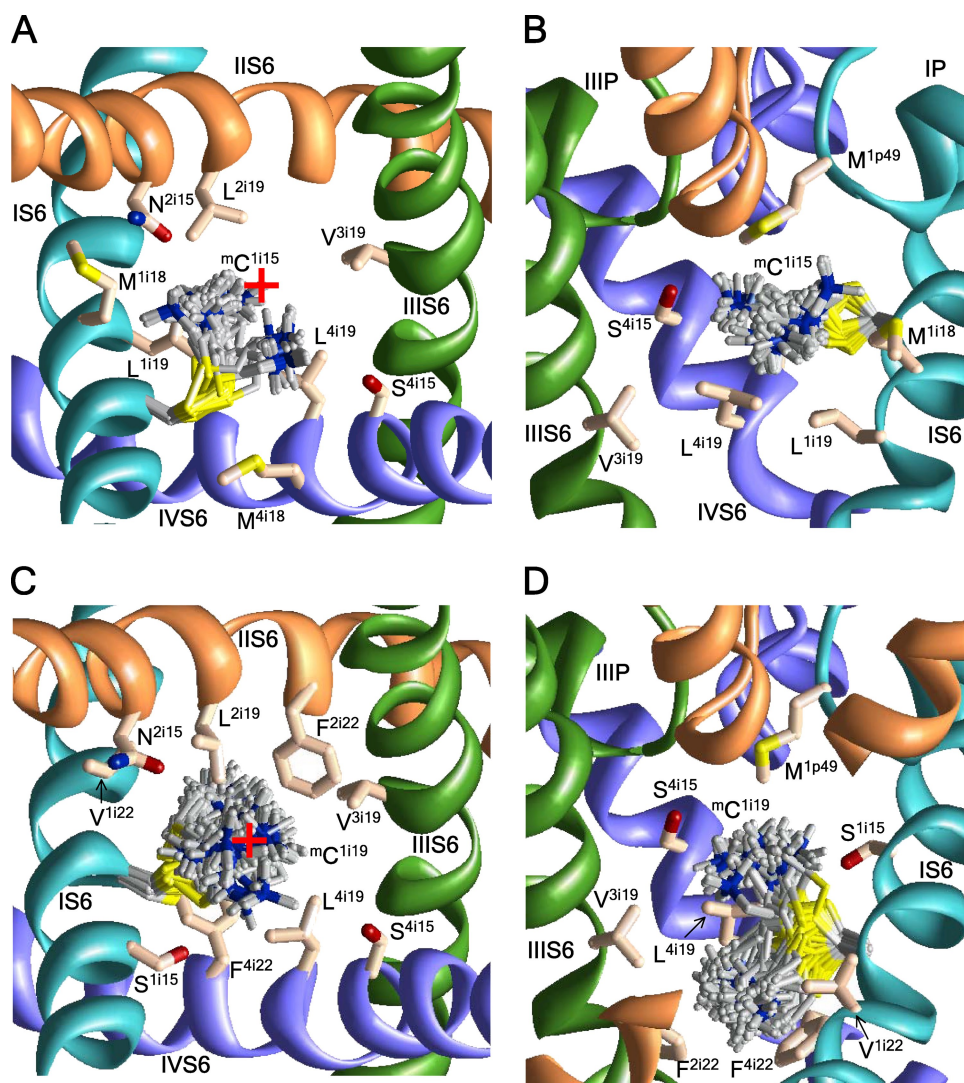


Figure 2. The extracellular and side views of mC^{115} (A and B) and mC^{119} (C and D) in the open $Ca_v2.1$ channel. The side chains of the mC residues in different conformations within 4 kcal/mol from the apparent global minima are superimposed and shown as gray sticks with blue nitrogen and yellow sulfur atoms. Native residues are shown in the lowest energy conformation as pale orange sticks with red oxygens, blue nitrogens, and yellow sulfur atoms. The P loops and S6s in repeats I, II, III, and IV are cyan, orange, green, and violet, respectively. For clarity, P loops in A and C, IIS6 in B, and S5s in A–D are not shown. The ammonium group of mC^{115} is inside the pore (A) between levels *i15* and *i18* (B). The ammonium group of mC^{119} is close to the pore axis (C) approaching either the focus of P helices or level *i22* (D). The red cross at A and C indicates the pore axis.

S6 positions that do not face the pore

The mC^{i16} side chains have two orientations (Fig. 6 A), with the ammonium group either approaching the pore or being away from the pore. In either orientation, the ammonium groups are rather far from the pore axis.

Cysteines at positions *i20* substitute native asparagines that are highly conserved in Ca^{2+} and Na^+ channels. Mutants C^{i120} and C^{i420} are nonfunctional. MTSET inhibits channels C^{i20} and C^{i320} by 56.2 and 42.0%, respectively. The side chains of mC^{i20} and mC^{i320} can adopt three orientations. In the energetically most preferable orientations, the ammonium groups face S5s (Fig. 6 B), whereas orientation to the pore is less preferable.

The C^α - C^β vectors of C^{i21} s direct to S5s. The C^{i421} mutant is not functional. MTSET does not inhibit channels C^{i121} and C^{i2121} , but it inhibits the C^{i321} channel by $\sim 40\%$. The mC^{i121} and mC^{i2121} side chains fit between the mutated repeat S5 and the next repeat S6 (Fig. S3). Cation- π interactions with F^{i430} and electrostatic interactions with Q^{i109} stabilize the ammonium group of mC^{i121}

in interface IV/I. Similar interactions with F^{i130} and E^{i129} stabilize the ammonium group of mC^{i2121} in interface I/II. The side chain of mC^{i321} orients into the pore, where it experiences cation- π interactions with F^{i318} and F^{i322} .

MTSET inhibits channels C^{i24} by 73.6–100%, despite the fact that vectors C^α - C^β direct toward cytoplasm rather than to the pore axis, (Fig. 1 B). In the most preferable conformations, cation- π interactions with F^{i130} , F^{i3130} , and F^{i430} attract the ammonium groups of mC^{i2124} , mC^{i4124} , and mC^{i1124} , respectively, toward the pore. However, in these orientations, the ammonium nitrogen is as far as 6–8 Å from the pore axis. In the alternative orientations, toward S5, the ammonium group is even farther from the pore axis (Fig. S2 B). The ammonium group of mC^{i3124} lacks an aromatic partner (note L^{i2130} vs. F^{i130} , F^{i3130} , and F^{i430} , Table I) and faces away from the pore in the most preferable orientation. Thus, our open $Ca_v2.1$ models are inconsistent with the strong current inhibition by mC^{i24} s. Possible explanations of this fact are provided in the Discussion.

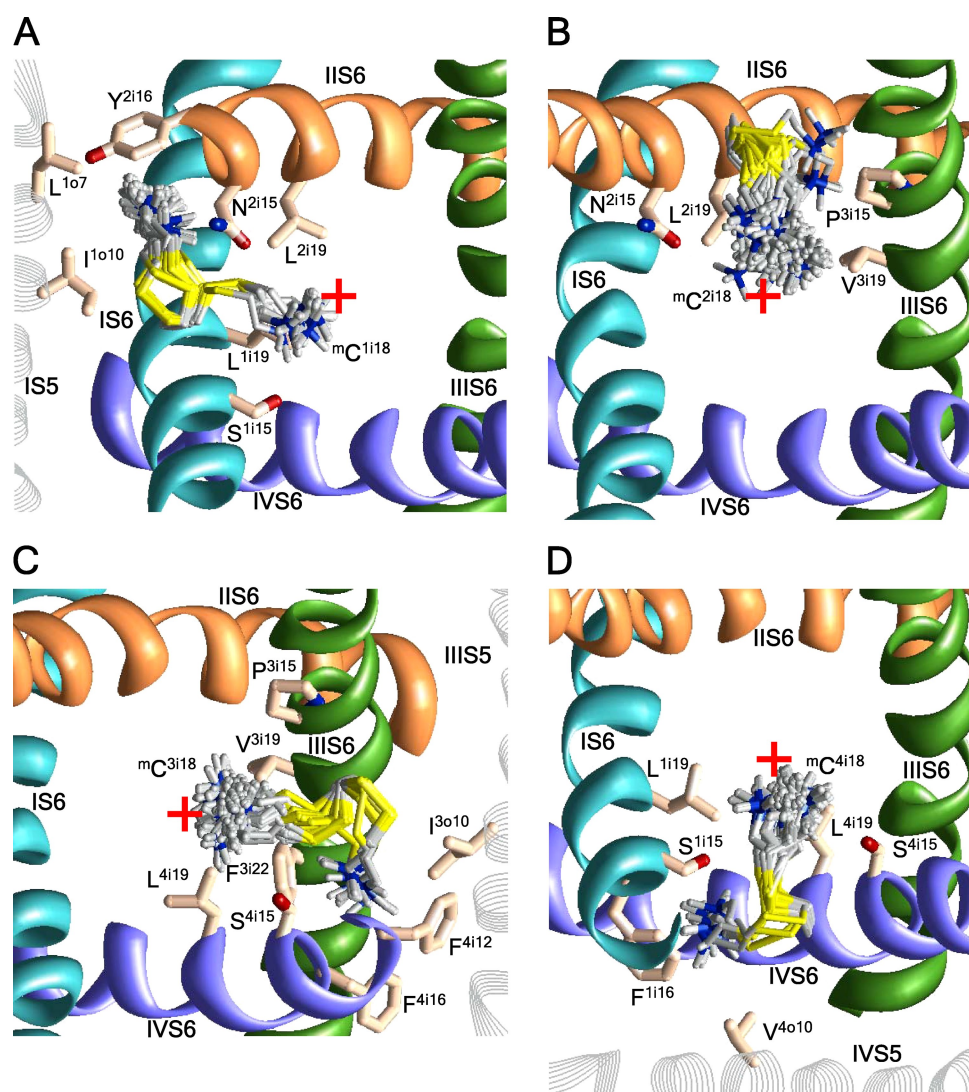


Figure 3. The extracellular views of various orientations of mC^{i18} residues in $Ca_v2.1$. For clarity, the P loops are not shown. The red cross indicates the pore axis. (A) Cation- π interactions with Y^{2116} stabilize the repeat interface orientation mC^{i118} ; the pore orientations have higher energy. (B) In the most preferable conformations, mC^{i218} is oriented in the pore. (C) Cation- π interactions with F^{412} and F^{322} stabilize orientation of mC^{i318} in the repeat interface; the pore orientations are less preferable. (D) Both pore and repeat interface orientations of mC^{i418} are energetically favorable.

Vectors $C^{\alpha}-C^{\beta}$ in positions *i25* direct away from the pore (Fig. 1). MTSET weakly inhibits channels C^{1i25} , C^{3i25} , and C^{4i25} , but it strongly inhibits channel C^{2i25} . Aromatic residues *i16* stabilize the orientations of mC^{1i25} , mC^{3i25} , and mC^{4i25} away from the pore (Fig. S4), whereas large residues L^{1i26} , L^{3i26} , and V^{4i26} preclude orientations into the pore. In contrast, cation- π interactions with F^{2i22} stabilize the orientation of mC^{2i25} to the pore, which is not precluded by small A^{2i26} .

C^{2i29} is the only channel in which MTSET increases the current. In our model, the mC^{2i29} ammonium group is oriented into the II/III interface and binds between the side chains of N^{3o9} , N^{3o6} , and N^{3i20} . Position *i29* is four helical turns closer to the cytoplasm than position *i14*, where the gating-hinge glycine is located in K^+ channels. Superposition of the x-ray structures of K^+ channels in the open and closed conformations shows that position *i29* shifts significantly between these structures. We suggest that the electrostatic attraction of the mC^{2i29} ammonium group to N^{3o9} , N^{3o6} , and N^{3i20} stabilizes the

open-gate conformation of the pore domain. This can explain the unique characteristics of the mC^{2i29} channel.

MTS-modified cysteines in S5s

Despite the fact that S5 helices do not line the pore, MTSET inhibits channels C^{2o10} and C^{4o10} (but not C^{3o10}). Experimental data for the C^{1o10} channel are unavailable. In the energetically preferable conformations, the ammonium groups of mC^{2o10} and mC^{4o10} extend between S6s and approach the pore (Fig. 7, A and C), whereas the orientation of mC^{3o10} to the repeat interface is stabilized by cation- π interactions with F^{3i14} , F^{3i17} , F^{3i18} , and Y^{3o14} (Fig. 7 B). MTSET inhibits C^{3o15} , C^{4o12} , and C^{4o17} channels by ~ 30 – 40% . In our models, the side chain of mC^{4o17} extends toward the pore, whereas the side chains of mC^{3o15} and mC^{4o12} face away from the pore.

Currents correlate with the distance of mC_N^+ from the pore axis

The above results provide multiple examples showing that the mC side chains adopt essentially different orientations

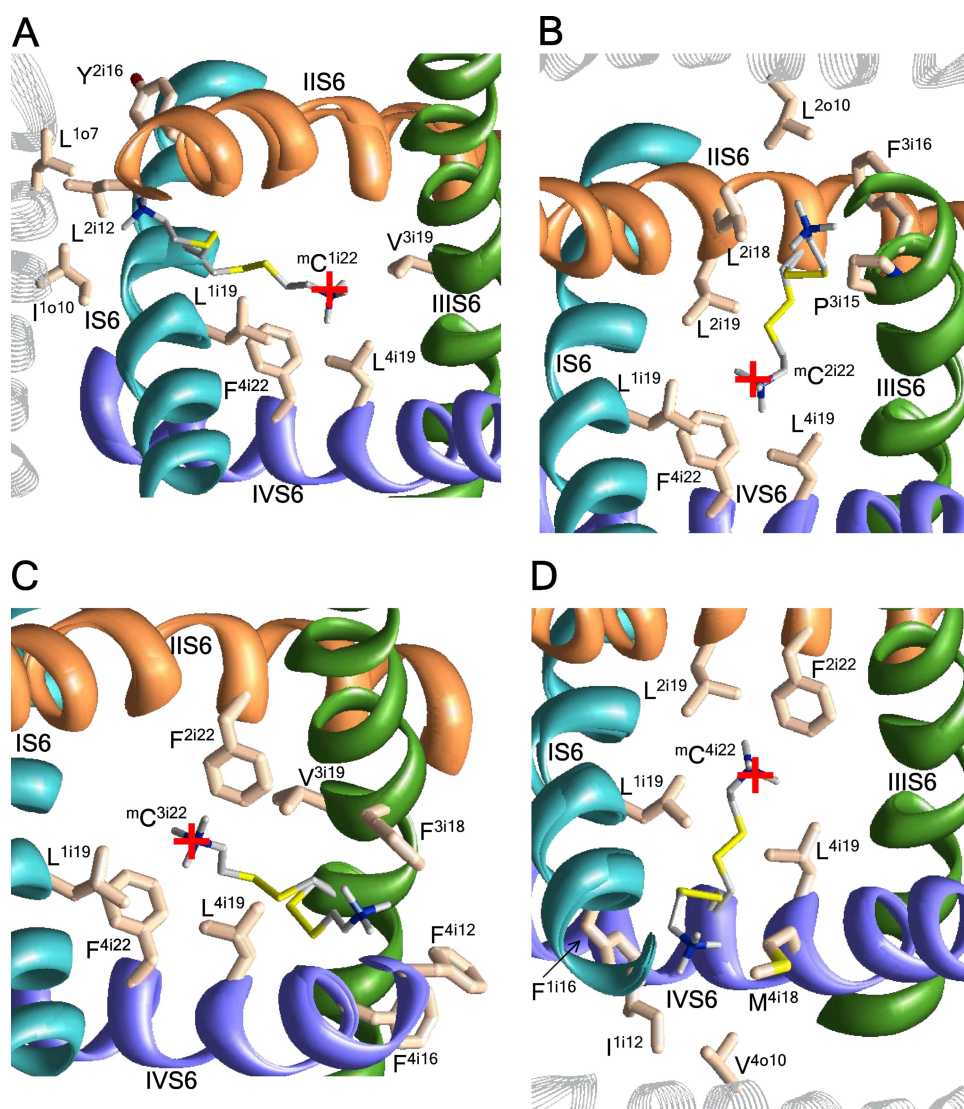


Figure 4. The extracellular views of mC^{1i22} (A), mC^{2i22} (B), mC^{3i22} (C), and mC^{4i22} (D). The red cross indicates the pore axis. P loops are not shown for clarity. Orientations of the mC^{i22} side chains in the pore are energetically more preferable than repeat interface orientations.

relative to the pore. Electrostatic and cation- π interactions between the ${}^m\text{C}$ ammonium group and its surrounding residues stabilize these particular orientations. Importantly, the MTSET potency correlates with the distance between the pore axis and the N^+ atom of the respective ${}^m\text{C}$ residue (Fig. 8 A). The current inhibition decreases with the distance, approaching the level of $\sim 20\%$ at distances $>16 \text{ \AA}$. The inhibition of the channels at large distances cannot be explained by electrostatic repulsion between the MTSET ammonium group and permeating cations; rather, it reflects the fact that MTSET inhibits by $19 \pm 5.9\%$ the “control channel” in which the α_1 subunit lacks both native and engineered cysteines (Zhen et al., 2005). In Fig. 8 B, the energetically preferable orientations of ${}^m\text{C}$ residues in the pore are shown by the N^+ atoms, which are colored according to the MTSET effect on the respective channels. Substantial current inhibition is usually observed when the N^+ atom (colored yellow) is inside the pore, and weak inhibition is usually observed when the N^+ atom (colored blue) is outside the pore. In contrast, the C^β atom position of the engineered cysteine does not correlate with the current inhibition by MTSET (Fig. 8, C and D).

DISCUSSION

In the absence of x-ray structures of Na^+ and Ca^{2+} channels, their homology models based on x-ray structures of K^+ channels are used to interpret data from mutational, electrophysiological, and ligand-binding experiments. The most crucial step in homology modeling is the sequence alignment. Because the sequence similarity between Ca^{2+} and Na^+ channels is rather high, alignment between these channels is unambiguous (Zhorov and Tikhonov, 2004). In contrast, due to poor sequence similarity between K^+ channels on one hand and Ca^{2+} and Na^+ channels on the other hand, there is no consensus alignment of S5 and S6 segments between these channels. In particular, the proposed alignments for S6s (Huber et al., 2000; Zhorov et al., 2001; Lipkind and Fozzard, 2003; Shafrir et al., 2008; Stary et al., 2008) differ in positions of asparagines that are highly conserved in every repeat of eukaryotic Ca^{2+} and Na^+ channels and are present in the homotetrameric bacterial channel NaChBac (Ren et al., 2001). In the alignment proposed by Lipkind and Fozzard (2003), these asparagines do not appear in matching positions of the four

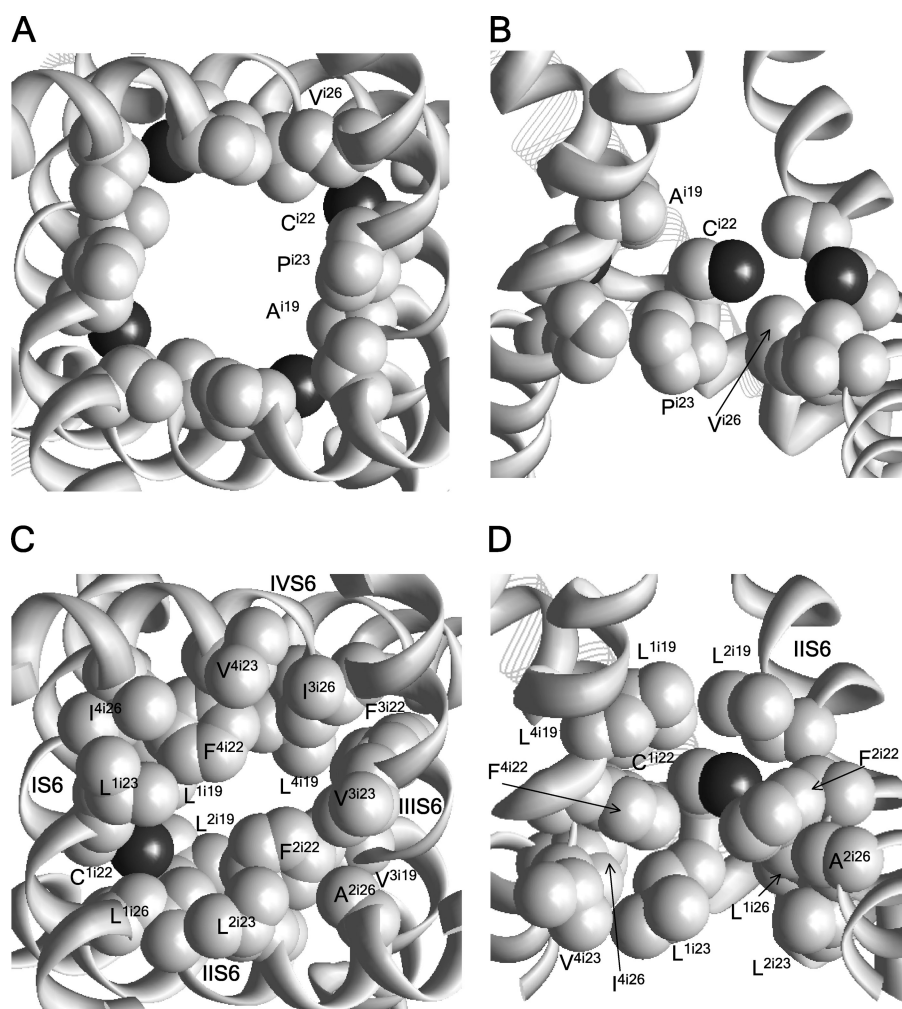


Figure 5. The cytoplasmic (A and C) and side (B and D) views of the environment for C^{122} in the *Shaker* (A and B) and $\text{Ca}_v2.1$ (C and D) channels. Side chains in positions *i19*, *i22*, *i23*, and *i26* are space-filled with gray carbon and black sulfur atoms.

repeats. K_v1.2-based models of the Ca_v1.2 (Stary et al., 2008) and NaChBac (Shafrir et al., 2008) have been built with the alignment in which an insertion is introduced at the conserved asparagines.

Intensive studies identified residues that, when mutated, affect the action of ligands targeting the pore of voltage-gated Ca²⁺ (Hockerman et al., 1997) and Na⁺ channels (Catterall et al., 2005). Homology models of these channels were used to visualize the binding sites and propose atomic mechanisms of various drugs, including benzothiazepines (Tikhonov and Zhorov, 2008), dihydropyridines (Zhorov et al., 2001; Lipkind and Fozzard, 2003; Cosconati et al., 2007; Tikhonov and Zhorov, 2009), phenylalkylamines (Lipkind and Fozzard, 2003; Cheng et al., 2009), local anesthetics (Lipkind and Fozzard, 2005; Tikhonov and Zhorov, 2007; Bruhova et al., 2008), steroidal sodium channel activators (Wang et al., 2006), and pyrethroid insecticides (O'Reilly et al., 2006; Du et al., 2009).

The above models have been built using different templates and different alignments. How sensitive are these results to the choice of the template and alignment?

In a recent study (Cheng et al., 2009), several Ca_v1.2 models based on different open-channel templates (KvAP, MthK, and K_v1.2) and different alignments were compared in terms of interactions with a flexible phenylalkylamine ligand devapamil. The alignment, which we use here (Table I), and all three templates were found to be consistent with the ligand–channel contacts known from experiments. The reason for such promiscuity of the modeled ligand–channel interactions to the choice of the open-channel template is the flexibility of both devapamil and side chains of devapamil-interacting tyrosines. This flexibility compensated rather small differences in the templates, all of which have similar patterns of the pore-facing residues. However, a shift of the S6 alignment between K⁺ and Ca²⁺ channels by just one position resulted in such dramatic reorientation of devapamil-interacting residues in the models that it was not possible to establish critical devapamil–Ca_v1.2 contacts known from experiments.

Because ¹³C residues can be considered as long, flexible tethered ligands, predicted contacts of the ¹³C

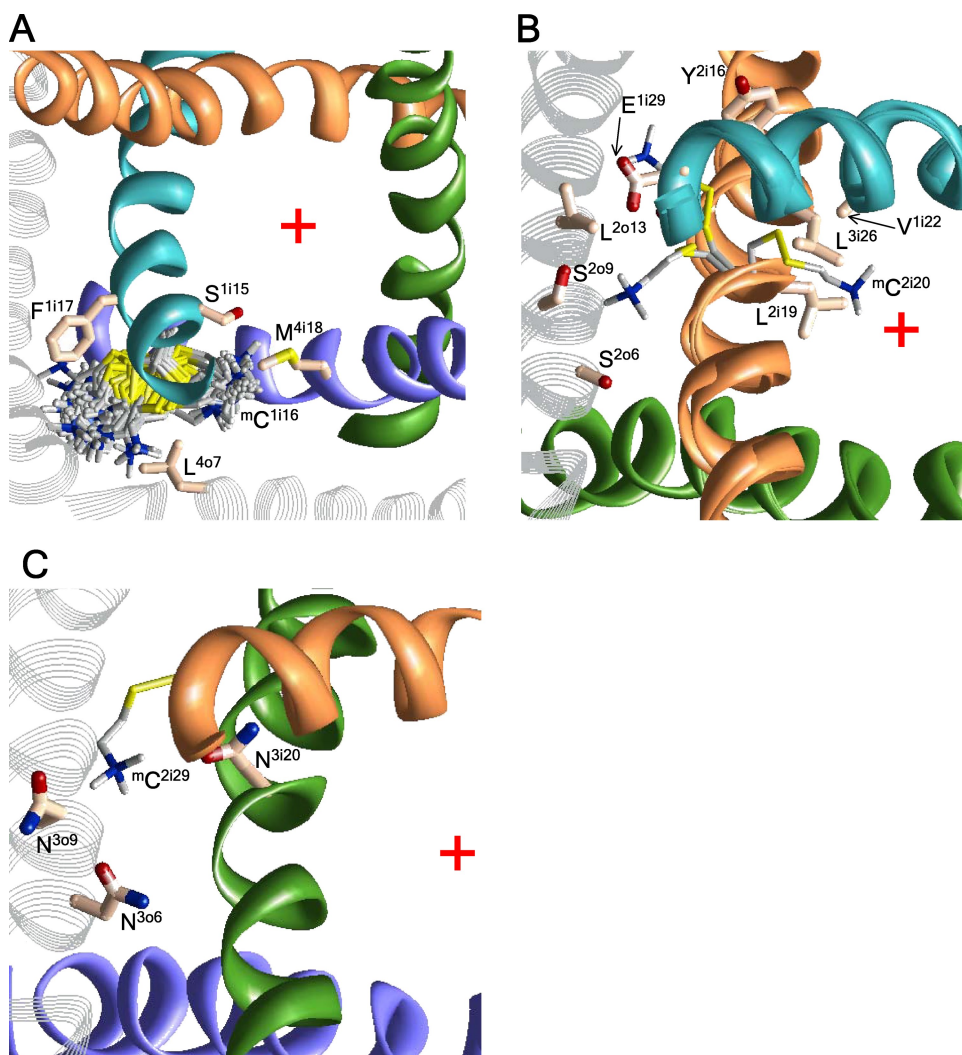


Figure 6. (A) The extracellular view of ¹³C¹¹¹⁶ that orients either along IVS6 or toward IS5 and IVS5. (B) The cytoplasmic view of three possible orientations of ¹³C²¹²⁰. (C) The cytoplasmic view of ¹³C²¹²⁹ interacting with N³⁰⁶, N³⁰⁹, and N³¹²⁰. The red cross indicates the pore axis.

ammonium groups should also be highly sensitive to the sequence alignment, but rather insensitive to the choice of the open-channel x-ray template. A single-position shift in the alignment would turn a pore-directing vector $C^\alpha-C^\beta$ away from the pore and vice versa. For example, C^{i15} channels are sensitive to MTSET, whereas C^{i17} channels are not (Zhen et al., 2005). In agreement with these data and the cysteine orientation concept, our models built using alignment shown in Table I (Zhorov et al., 2001) have vectors $C^\alpha-C^\beta$ in positions $i15$ and $i17$, which direct to the pore axis and away from it, respectively. Models built using other alignments (Huber et al., 2000; Lipkind and Fozzard, 2003) are unlikely to provide a correlation with experimental SCAM data (Zhen et al., 2005). Guy and coworkers (Durell and Guy, 2001; Stary et al., 2008) proposed an S6 alignment, which is similar to that shown in Table I, but it has an insertion at position $i20$ of each repeat of the Ca^{2+} channel. A model built with this alignment does not resolve

the problem of the pore-away orientation of mC^{i24} residues in the outlying channels (Fig. 8), whereas $C^\alpha-C^\beta$ vectors in residues downstream of positions $i20$ orient differently than in our models, which are based on our alignment (Table I) and which explain the SCAM data.

Despite different alignments underlying these models, and different details of predicted ligand-channel interactions, the above studies agree that the x-ray structures of K^+ channels provide reasonable templates for the homology modeling of Na^+ and Ca^{2+} channels. This opinion was undermined by the interpretation of SCAM experiments with $Ca_v2.1$, which suggests that Ca^{2+} and K^+ channels have different patterns of pore-lining residues and questions the symmetric arrangement of the Ca^{2+} channel repeats around the pore axis (Zhen et al., 2005).

We do not doubt the experimental observations of the SCAM study, but we show here that interpretation of these observations requires analysis of some factors, which were apparently not considered in the original

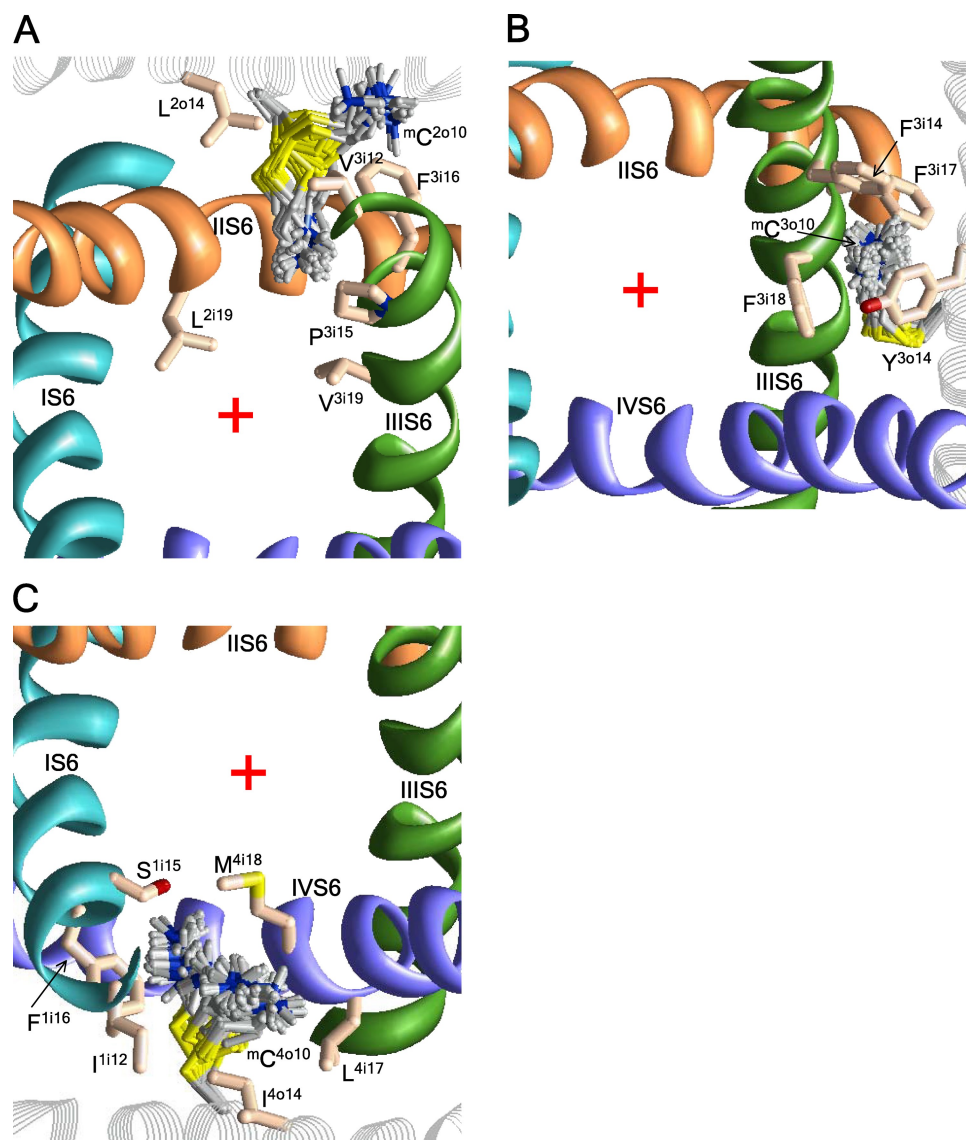


Figure 7. The extracellular view of possible orientations of mC residues in S5s. mC^{2010} (A) and mC^{4010} (C) can extend their ammonium groups toward the pore. (B) mC^{3010} is stabilized inside the repeat interface by cation- π interaction with F^{318} and Y^{3014} . The red cross indicates the pore axis.

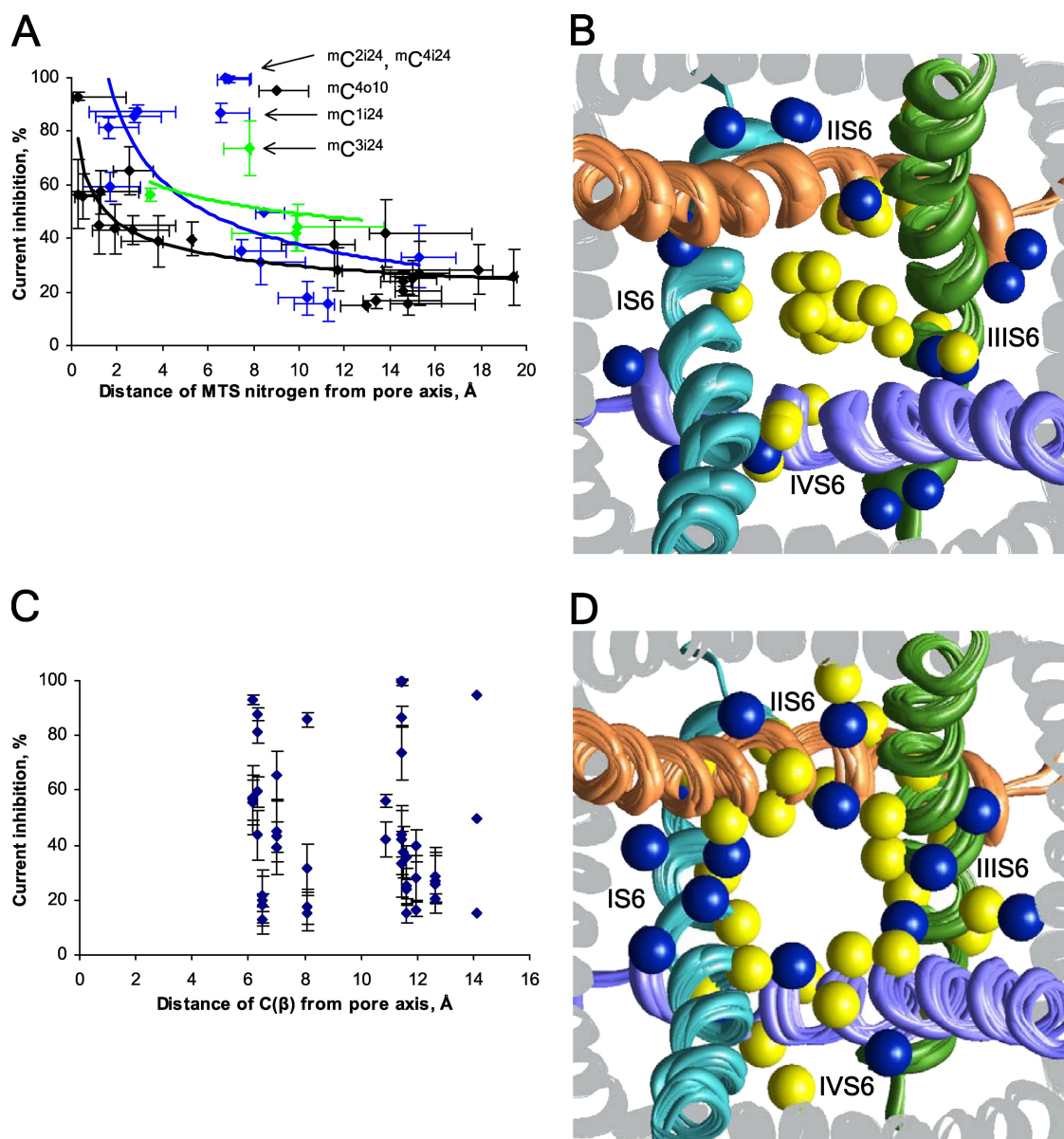


Figure 8. The residual current upon MTSET application correlates with the distance of the MTS atom $N^{+}\text{-}^{13}\text{C}$ (A and B), but not atom $C^{\beta}\text{-}^{13}\text{C}$ (C and D) from the pore axis. (A and C) The experimental values of the current inhibition with standard deviations (Zhen et al., 2005) are plotted against the predicted distances of atoms $N^{+}\text{-}^{13}\text{C}$ (A) or C^{β} (C) from the pore axis. Data are shown for channels with engineered cysteines in positions *i15–i21*, *i23–i25*, and *o10*. Black dots represent the apparent global minima of channels in which all minimum energy conformations of ^{13}C side chains are unambiguously oriented in respect to the pore (e.g., inside the pore for channels mC^{i15} or outside the pore for channels mC^{i17}). Blue dots represent the apparent global minima of the channels in which the ^{13}C side chain adopts low energy conformations with distinct orientation in respect to the pore (e.g., channels mC^{i18}). A green dot represents a local minimum (within 2 kcal/mol from the apparent global minimum) of a channel in which the ^{13}C side chain adopts conformations with distinct orientation in respect to the pore (e.g., channels mC^{i16}). Horizontal lines show the $N^{+}\text{-}^{13}\text{C}$ atom mobility in conformations within 2 kcal/mol from the apparent global minimum (Table S1). Note a smooth decrease of the current inhibition with increase of the distance between the MTS nitrogen and the pore axis. The current inhibition of $\sim 20\%$ at distances >16 Å corresponds to MTSET block of the “control channel,” in which eight native cysteines in the α_1 subunit have been replaced with alanines and no engineered cysteines have been introduced (Zhen et al., 2005). (B and D) The extracellular view of $\text{Ca}_v2.1$, with atoms $^{13}\text{C}\text{-}N^{+}$ (B) and $^{13}\text{C}\text{-}C^{\beta}$ (D) shown as spheres. P loops are omitted for clarity. Yellow and blue spheres represent the respective atoms in the channels, which are inhibited by MTSET by >30 and $\leq 30\%$, respectively. (B) In most of the channels, which are strongly inhibited by MTSET, the yellow-colored ammonium nitrogen ($^{13}\text{C}\text{-}N^{+}$) is located either close to the pore axis or at the inner surface of the pore, whereas in the channels, which are weakly inhibited by MTSET, the blue-colored ammonium nitrogen is not inside the pore. (D) Location of β carbons does not correlate with the level of current inhibition by MTSET. Both yellow and blue spheres are randomly distributed at different sides of the inner helices.

study (Zhen et al., 2005). These factors are conformational flexibility of long side chains of ^{13}C residues and their interaction with neighboring residues. In this study, we used the K_v1.2-based models of Ca_v2.1, which are based on the alignment shown in Table I. We reasoned that if our results explain the SCAM data, this supports the alignment as well as the generally similar spatial disposition of S5s and S6s in K⁺ and Ca²⁺ channels.

Interpreting SCAM experiments is not straightforward. Several factors should be taken into consideration (Karlin and Akabas, 1998). MTS reagents, such as MTSET, react with water-accessible ionized cysteines to form a covalent bond. If a cysteine is exposed to the lipid bilayer or buried inside the protein, the ionization of the thiol group is suppressed. It is assumed that the MTS reagent covalently bound to the engineered cysteine and exposed to the pore decreases the current. The current may be unaffected because of two causes. First, the reaction does not proceed due to hydrophobic environment, steric constraints, lack of ionized Cys residues, or other grounds. Second, the reaction proceeds, but the MTS-modified cysteine does not affect ion permeation.

A brief application of an MTS reagent may result in incomplete chemical modification of cysteines (Liu et al., 1997), but prolonged exposure increases the probability of disulfide formation, even with partially buried cysteines. The prolonged Ca_v2.1 exposure to MTSET suggests that cysteines in different sides of S6s and S5s were modified (Zhen et al., 2005), despite the fact that some positions do not face the ion permeation pathway.

Besides the kinetic effects, other factors should be considered to interpret SCAM data. An MTS-modified cysteine has a long flexible side chain: in the all-trans conformation, the distance between atoms C $^{\alpha}$ and N⁺ is 8.4 Å. Prediction of energetically optimal conformations of an ^{13}C residue can be considered as docking of a tethered ligand to the channel. The energetically optimal position of the ^{13}C ammonium group depends on interactions with neighboring residues, among which electrostatic attractions (including those with the nucleophilic C ends of P helices) and cation- π interactions play the major role. In some positions where vector C $^{\alpha}$ -C $^{\beta}$ directs to the pore axis, the ^{13}C ammonium group does not bind in the pore. And in some positions where vector C $^{\alpha}$ -C $^{\beta}$ directs away from the axis, the ammonium group can reach the pore through the repeat interface or by wrapping around the mutated helix. Thus, the exposure of the ^{13}C ammonium group to the pore may not correlate with the angle between the C $^{\alpha}$ -C $^{\beta}$ vector and the vector drawn from the C $^{\alpha}$ atom to the pore axis.

Our calculations predict that the reported current inhibition by MTSET generally decreases with the predicted distances between the ammonium nitrogen and the pore axis (Fig. 8, A and B). This trend is important. First, it shows that interpretation of the SCAM data is possible in gradual rather than discrete terms. Second,

it supports the underlying sequence alignment between Ca²⁺ and K⁺ channels (Table I). Third, it implies the fourfold symmetry of transmembrane helices in the pore-forming domain of Ca²⁺ channels and a similar disposition of S5s and S6s in K⁺ and Ca²⁺ channels. Fourth, it shows that significant block is observed only when the ammonium group occurs in the pore, but partial inhibition is possible when the ammonium group is rather far from the pore axis. This is in agreement with the single-channel recordings, which demonstrate that MTSET decreases the current amplitude (Lu et al., 1999).

In homotetrameric K⁺ channels, a single mutation to Cys yields four identical potential targets for the reaction with an MTS reagent. Linking the subunits in a single polypeptide chain allowed the expression of channels with one, two, three, or four cysteines at a given S6 position (Lu et al., 1999). MTSET application to the channels with one, two, and three cysteines in position *i18* inhibited the current by 24, 55, and 80%, respectively. MTSET application to channels with one, two, and three engineered cysteines in position *i22* resulted in the current inhibition by 51, 80, and 89%, respectively (Lu et al., 1999). Thus, the introduction of one positive charge into the open pore of the K⁺ channel does not fully inhibit current. Furthermore, protonation at the selectivity filter of Ca²⁺ channels reduces single-channel conductance, but it does not produce a complete channel block (Prod'homme et al., 1987; Pietrobon et al., 1989). The above observations are consistent with the fact that the ^{13}C residues at pore-facing positions, such as *i15* and *i19*, reduce the current but do not completely inhibit it. On the other hand, incomplete inhibition could indicate that the MTS application did not modify 100% of the respective channels.

Fig. 8 A was obtained using data for 40 channels. In 36 cases, the data point represents the energetically most preferable conformation of the respective ^{13}C residue, and in four channels, green points represent local minima with the energies up to 2 kcal/mol above the apparent global minima. Thus, only 10% of the data points represent the less populated conformations. The fact that our method yields a poorer correlation when only the apparent global minima were considered is understandable in view of the limited precision of the homology modeling. Because of these limitations, quantitative analysis is difficult, particularly when an ^{13}C side chain has two distinct groups of conformers, one group inside the pore (the ammonium nitrogen within 4 Å from the pore axis) and another away from the pore. Two distinct groups of conformers were observed, e.g., for $^{13}\text{C}^{i16}$ and $^{13}\text{C}^{i18}$ (Figs. 6 A and 3).

In Fig. 8, we did not include the Cⁱ²² channels, which are not inhibited by MTSET. Respective C $^{\alpha}$ -C $^{\beta}$ vectors face the pore, and in the most preferable conformations of $^{13}\text{C}^{i22}$ residues, the N⁺ atoms occur inside the pore. There are local minima with the N⁺ atoms beyond

the pore, and the Cⁱ²² channels could be represented by green points in Fig. 8 A. However, we believe that Cⁱ²² channels are insensitive to MTSET because large hydrophobic residues in the pore-facing positions *i19*, *i22*, and *i26* (Table I) create a highly hydrophobic environment that precludes the reaction with MTSET. In Na⁺ channels, Yⁱ⁴²² was proposed to face the pore and interact with local anesthetics (Tikhonov and Zhorov, 2007; Bruhova et al., 2008). However, the Yⁱ⁴²²C mutant is insensitive to MTS ethylammonium or MTSET (Sunami et al., 2004). A possible cause is the hydrophobic environment of Cⁱ⁴²² created by hydrophobic residues in the pore-facing positions *i19*, *i22*, and *i26*. In contrast to Ca²⁺ and Na⁺ channels, MTSET inhibits the Cⁱ²² mutants of *Shaker* (Liu et al., 1997), Kir2.1 (Lu et al., 1999), Kir6.2 (Phillips et al., 2003), and K_{Ca}3.1 (Klein et al., 2007). Position *i22* of the *Shaker* is surrounded by Aⁱ¹⁹, Cⁱ²², and Vⁱ²⁶, which provide a favorable environment for reaction with MTSET (Fig. 5). The ring *i22* of Ca_v2.1 is unique in terms of the completely hydrophobic environment at its own level and the levels of the pore-facing residues *i19* and *i26* above and below the ring, respectively. The correlation in Fig. 8 A suggests a common mechanism of current inhibition by MTSET, but we cannot rule out that incomplete block of some channels also results from slow reaction with MTSET.

The correlation in Fig. 8 A has five prominent outliers, which correspond to channels C^{4o10}, Cⁱ¹²⁴, C²ⁱ²⁴, C³ⁱ²⁴, and C⁴ⁱ²⁴. These channels are strongly blocked by MTSET, despite the fact that corresponding ^mC_N⁺ atoms are 6.5–9 Å from the pore axis. Vectors C^α–C^β in positions *i24* are close to the cytoplasm in the x-ray structures of K_v1.2, KcsA, and the closed-K_v1.2 model (Pathak et al., 2007), suggesting that MTSET could attack Cⁱ²⁴s not from the open pore, but from the cytoplasm. The strong current inhibition in channels ^mCⁱ²⁴ may arise from stabilization of the closed-channel conformation. To explore this possibility, we have built KcsA-based models of the ^mCⁱ²⁴ mutants of Ca_v2.1 and sought for possible contacts of the ^mCⁱ²⁴ ammonium group with nearby acidic residues, which were modeled in the ionized forms (Fig. S5). MC minimizations with distance constraints biasing inter-repeat salt bridges yielded low energy structures with the following salt bridges: ^mCⁱ¹²⁴–D⁴ⁱ²⁸, ^mC²ⁱ²⁴–E^{Li29}, ^mC³ⁱ²⁴–D²ⁱ²⁸, and ^mC⁴ⁱ²⁴–E³ⁱ³². These remained stable in subsequent MC minimizations without the constraints. These salt bridges could be formed upon MTSET application to the hyperpolarized membrane and preclude channel opening upon membrane depolarization. However, in the absence of an x-ray structure of a closed voltage-gated channel, we cannot rule out other possible mechanisms of closed-channel stabilization, e.g., interaction of ^mCⁱ²⁴ residues with the β subunit or cytoplasmic segments of the α₁ subunit.

Experiments with fluorinated aromatic residues (Santarelli et al., 2007; Ahern et al., 2008; Xiu et al.,

2009) proved the long-proposed role of cation–π interactions in ligand receptor recognition. Despite the fact that the AMBER force field lacks a specific energy term for cation–π interactions, these interactions can be detected in structures where the ammonium group is attracted to partial negative charges of aromatic carbons (Bruhova et al., 2008). Such structures were earlier predicted for complexes of Na⁺ channels with local anesthetics (Fozzard et al., 2005; Lipkind and Fozzard, 2005; Tikhonov and Zhorov, 2007). Here, we found many structures in which the ^mC ammonium groups are attracted to aromatic residues via cation–π interactions. These interactions were particularly important in stabilizing the ammonium groups of ^mC^{3o10}, ^mCⁱ¹¹⁸, and ^mC³ⁱ¹⁸ in the repeat interfaces, as well as ^mCⁱ¹⁹ in the pore.

According to our models, the repeat interfaces would provide the access paths for MTSET to engineered cysteines in those positions of S5s and S6s that do not face the pore. Furthermore, the ammonium group of an ^mC residue can extend through a repeat interface into the pore and decrease the current. Interesting examples are ^mC^{2o10} and ^mC^{4o10}, whose ammonium groups can approach the pore only through the repeat interface. This prediction is consistent with our studies, which suggest that the III/IV repeat interface provides the extracellular access route for local anesthetics into Na⁺ channels (Bruhova et al., 2008) as well as benzothiazepines (Tikhonov and Zhorov, 2008) and dihydropyridines (Tikhonov and Zhorov, 2009) in Ca²⁺ channels.

In conclusion, here we used molecular modeling to reinterpret the results of the SCAM study of Ca_v2.1 (Zhen et al., 2005). We found that the residual current upon MTSET application does not correlate with the orientation of the C^α–C^β vector in the modified residue to the pore, but generally decreases with the distance between the pore axis and the N⁺ atom of the respective ^mC residue. Our models suggest that different local environments of equivalent positions in the four repeats lead to different SCAM results reported for such positions. Our study supports the sequence alignments between K⁺ and Ca²⁺ channels earlier proposed for S5s (Huber et al., 2000) and S6s (Zhorov et al., 2001), and suggests that the x-ray structure of K_v1.2 is a suitable template to model Ca²⁺ channels in the open-state conformation.

We thank Denis Tikhonov for helpful discussions.

This study was supported by a grant (MOP-53229) from the Canadian Institutes of Health Research (CIHR) to B.S. Zhorov. I. Bruhova is a recipient of the CIHR Canada Graduate Scholarships. Computations were made possible by the facilities of the Shared Hierarchical Academic Research Computing Network (SHARCNET; <http://www.sharcnet.ca>).

Lawrence G. Palmer served as editor.

Submitted: 2 July 2009

Accepted: 5 February 2010

REFERENCES

- Ahern, C.A., A.L. Eastwood, D.A. Dougherty, and R. Horn. 2008. Electrostatic contributions of aromatic residues in the local anesthetic receptor of voltage-gated sodium channels. *Circ. Res.* 102:86–94. doi:10.1161/CIRCRESAHA.107.160663
- Anderson, P.A., and R.M. Greenberg. 2001. Phylogeny of ion channels: clues to structure and function. *Comp. Biochem. Physiol. B Biochem. Mol. Biol.* 129:17–28. doi:10.1016/S1096-4959(01)00376-1
- Bruhova, I., and B.S. Zhorov. 2007. Monte Carlo-energy minimization of correolide in the Kv1.3 channel: possible role of potassium ion in ligand-receptor interactions. *BMC Struct. Biol.* 7:5. doi:10.1186/1472-6807-7-5
- Bruhova, I., D.B. Tikhonov, and B.S. Zhorov. 2008. Access and binding of local anesthetics in the closed sodium channel. *Mol. Pharmacol.* 74:1033–1045. doi:10.1124/mol.108.049759
- Catterall, W.A., A.L. Goldin, and S.G. Waxman. 2005. International Union of Pharmacology. XLVII. Nomenclature and structure-function relationships of voltage-gated sodium channels. *Pharmacol. Rev.* 57:397–409. doi:10.1124/pr.57.4.4
- Cheng, R.C., D.B. Tikhonov, and B.S. Zhorov. 2009. Structural model for phenylalkylamine binding to L-type calcium channels. *J. Biol. Chem.* 284:28332–28342. doi:10.1074/jbc.M109.027326
- Cosconati, S., L. Marinelli, A. Lavecchia, and E. Novellino. 2007. Characterizing the 1,4-dihydropyridines binding interactions in the L-type Ca²⁺ channel: model construction and docking calculations. *J. Med. Chem.* 50:1504–1513. doi:10.1021/jm061245a
- Dewar, M.J., E.G. Zoebisch, E.F. Healy, and J.J. Stewart. 1985. Development and use of quantum mechanical molecular models. 76. AM1: a new general purpose quantum mechanical molecular model. *J. Am. Chem. Soc.* 107:3902–3909. doi:10.1021/ja00299a024
- Doyle, D.A., J. Morais Cabral, R.A. Pfuetzner, A. Kuo, J.M. Gulbis, S.L. Cohen, B.T. Chait, and R. MacKinnon. 1998. The structure of the potassium channel: molecular basis of K⁺ conduction and selectivity. *Science*. 280:69–77. doi:10.1126/science.280.5360.69
- Du, Y., J.E. Lee, Y. Nomura, T. Zhang, B.S. Zhorov, and K. Dong. 2009. Identification of a cluster of residues in transmembrane segment 6 of domain III of the cockroach sodium channel essential for the action of pyrethroid insecticides. *Biochem. J.* 419:377–385. doi:10.1042/BJ20082082
- Dudley, S.C. Jr., N. Chang, J. Hall, G. Lipkind, H.A. Fozzard, and R.J. French. 2000. μ -Conotoxin GIIIA interactions with the voltage-gated Na⁺ channel predict a clockwise arrangement of the domains. *J. Gen. Physiol.* 116:679–690. doi:10.1085/jgp.116.5.679
- Durell, S.R., and H.R. Guy. 2001. A putative prokaryote voltage-gated Ca(2⁺) channel with only one 6TM motif per subunit. *Biochem. Biophys. Res. Commun.* 281:741–746. doi:10.1006/bbrc.2001.4408
- Fozzard, H.A., P.J. Lee, and G.M. Lipkind. 2005. Mechanism of local anesthetic drug action on voltage-gated sodium channels. *Curr. Pharm. Des.* 11:2671–2686. doi:10.2174/1381612054546833
- Hockerman, G.H., B.Z. Peterson, B.D. Johnson, and W.A. Catterall. 1997. Molecular determinants of drug binding and action on L-type calcium channels. *Annu. Rev. Pharmacol. Toxicol.* 37:361–396. doi:10.1146/annurev.pharmtox.37.1.361
- Huber, I., E. Wappl, A. Herzog, J. Mitterdorfer, H. Glossmann, T. Langer, and J. Striessnig. 2000. Conserved Ca²⁺-antagonist-binding properties and putative folding structure of a recombinant high-affinity dihydropyridine-binding domain. *Biochem. J.* 347:829–836. doi:10.1042/0264-6021:3470829
- Jiang, Y., A. Lee, J. Chen, V. Ruta, M. Cadene, B.T. Chait, and R. MacKinnon. 2003. X-ray structure of a voltage-dependent K⁺ channel. *Nature*. 423:33–41. doi:10.1038/nature01580
- Karlin, A., and M.H. Akabas. 1998. Substituted-cysteine accessibility method. *Methods Enzymol.* 293:123–145. doi:10.1016/S0076-6879(98)93011-7
- Klein, H., L. Garneau, U. Banderali, M. Simoes, L. Parent, and R. Sauvé. 2007. Structural determinants of the closed KCa3.1 channel pore in relation to channel gating: results from a substituted cysteine accessibility analysis. *J. Gen. Physiol.* 129:299–315. doi:10.1085/jgp.200609726
- Lazaridis, T., and M. Karplus. 1999. Effective energy function for proteins in solution. *Proteins*. 35:133–152. doi:10.1002/(SICI)1097-0134(19990501)35:2<133::AID-PROT1>3.0.CO;2-N
- Li, Z., and H.A. Scheraga. 1987. Monte Carlo-minimization approach to the multiple-minima problem in protein folding. *Proc. Natl. Acad. Sci. USA*. 84:6611–6615. doi:10.1073/pnas.84.19.6611
- Lipkind, G.M., and H.A. Fozzard. 2003. Molecular modeling of interactions of dihydropyridines and phenylalkylamines with the inner pore of the L-type Ca²⁺ channel. *Mol. Pharmacol.* 63:499–511. doi:10.1124/mol.63.3.499
- Lipkind, G.M., and H.A. Fozzard. 2005. Molecular modeling of local anesthetic drug binding by voltage-gated sodium channels. *Mol. Pharmacol.* 68:1611–1622.
- Liu, Y., M. Holmgren, M.E. Jurman, and G. Yellen. 1997. Gated access to the pore of a voltage-dependent K⁺ channel. *Neuron*. 19:175–184. doi:10.1016/S0896-6273(00)80357-8
- Long, S.B., E.B. Campbell, and R. MacKinnon. 2005. Crystal structure of a mammalian voltage-dependent Shaker family K⁺ channel. *Science*. 309:897–903. doi:10.1126/science.1116269
- Long, S.B., X. Tao, E.B. Campbell, and R. MacKinnon. 2007. Atomic structure of a voltage-dependent K⁺ channel in a lipid membrane-like environment. *Nature*. 450:376–382. doi:10.1038/nature06265
- Lu, T., B. Nguyen, X. Zhang, and J. Yang. 1999. Architecture of a K⁺ channel inner pore revealed by stoichiometric covalent modification. *Neuron*. 22:571–580. doi:10.1016/S0896-6273(00)80711-4
- O'Reilly, A.O., B.P. Khambay, M.S. Williamson, L.M. Field, B.A. Wallace, and T.G. Davies. 2006. Modelling insecticide-binding sites in the voltage-gated sodium channel. *Biochem. J.* 396:255–263. doi:10.1042/BJ20051925
- Pathak, M.M., V. Yarov-Yarovoy, G. Agarwal, B. Roux, P. Barth, S. Kohout, F. Tombola, and E.Y. Isacoff. 2007. Closing in on the resting state of the Shaker K(+) channel. *Neuron*. 56:124–140. doi:10.1016/j.neuron.2007.09.023
- Phillips, L.R., D. Enkvetchakul, and C.G. Nichols. 2003. Gating dependence of inner pore access in inward rectifier K(+) channels. *Neuron*. 37:953–962. doi:10.1016/S0896-6273(03)00155-7
- Pietrobon, D., B. Prod'hom, and P. Hess. 1989. Interactions of protons with single open L-type calcium channels. pH dependence of proton-induced current fluctuations with Cs⁺, K⁺, and Na⁺ as permeant ions. *J. Gen. Physiol.* 94:1–21. doi:10.1085/jgp.94.1.1
- Prod'hom, B., D. Pietrobon, and P. Hess. 1987. Direct measurement of proton transfer rates to a group controlling the dihydropyridine-sensitive Ca²⁺ channel. *Nature*. 329:243–246. doi:10.1038/329243a0
- Ren, D., B. Navarro, H. Xu, L. Yue, Q. Shi, and D.E. Clapham. 2001. A prokaryotic voltage-gated sodium channel. *Science*. 294:2372–2375. doi:10.1126/science.1065635
- Santarelli, V.P., A.L. Eastwood, D.A. Dougherty, R. Horn, and C.A. Ahern. 2007. A cation- π interaction discriminates among sodium channels that are either sensitive or resistant to tetrodotoxin block. *J. Biol. Chem.* 282:8044–8051. doi:10.1074/jbc.M611334200
- Shafir, Y., S.R. Durell, and H.R. Guy. 2008. Models of the structure and gating mechanisms of the pore domain of the NaChBac ion channel. *Biophys. J.* 95:3650–3662. doi:10.1529/biophysj.108.135327
- Stary, A., Y. Shafir, S. Hering, P. Wolschann, and H.R. Guy. 2008. Structural model of the Ca(V)1.2 pore. *Channels (Austin)*. 2:210–215.
- Sunami, A., A. Tracey, I.W. Glaaser, G.M. Lipkind, D.A. Hanck, and H.A. Fozzard. 2004. Accessibility of mid-segment domain IV S6 residues of the voltage-gated Na⁺ channel to methanethiosulfonate reagents. *J. Physiol.* 561:403–413. doi:10.1113/jphysiol.2004.067579

- Tikhonov, D.B. 2007. Ion channels of glutamate receptors: structural modeling. *Mol. Membr. Biol.* 24:135–147. doi:10.1080/09687860601008806
- Tikhonov, D.B., and B.S. Zhorov. 2005. Modeling P-loops domain of sodium channel: homology with potassium channels and interaction with ligands. *Biophys. J.* 88:184–197. doi:10.1529/biophysj.104.048173
- Tikhonov, D.B., and B.S. Zhorov. 2007. Sodium channels: ionic model of slow inactivation and state-dependent drug binding. *Biophys. J.* 93:1557–1570.
- Tikhonov, D.B., and B.S. Zhorov. 2008. Molecular modeling of benzothiazepine binding in the L-type calcium channel. *J. Biol. Chem.* 283:17594–17604. doi:10.1074/jbc.M800141200
- Tikhonov, D.B., and B.S. Zhorov. 2009. Structural model for dihydropyridine binding to L-type calcium channels. *J. Biol. Chem.* 284:19006–19017. doi:10.1074/jbc.M109.011296
- Tikhonov, D.B., I. Bruhova, and B.S. Zhorov. 2006. Atomic determinants of state-dependent block of sodium channels by charged local anesthetics and benzocaine. *FEBS Lett.* 580:6027–6032. doi:10.1016/j.febslet.2006.10.035
- Wang, S.Y., J. Mitchell, D.B. Tikhonov, B.S. Zhorov, and G.K. Wang. 2006. How batrachotoxin modifies the sodium channel permeation pathway: computer modeling and site-directed mutagenesis. *Mol. Pharmacol.* 69:788–795. doi:10.1124/mol.106.022368
- Weiner, S.J., P.A. Kollman, D.A. Case, U.C. Singh, C. Ghio, G. Alagona, S. Profeta, and P. Weiner. 1984. A new force-field for molecular mechanical simulation of nucleic-acids and proteins. *J. Am. Chem. Soc.* 106:765–784. doi:10.1021/ja00315a051
- Weiner, S.J., P.A. Kollman, D.T. Nguyen, and D.A. Case. 1986. An all atom force field for simulations of proteins and nucleic acids. *J. Comput. Chem.* 7:230–252. doi:10.1002/jcc.540070216
- Xiu, X., N.L. Puskar, J.A. Shanata, H.A. Lester, and D.A. Dougherty. 2009. Nicotine binding to brain receptors requires a strong cation-pi interaction. *Nature.* 458:534–537. doi:10.1038/nature07768
- Zhen, X.G., C. Xie, A. Fitzmaurice, C.E. Schoonover, E.T. Orenstein, and J. Yang. 2005. Functional architecture of the inner pore of a voltage-gated Ca^{2+} channel. *J. Gen. Physiol.* 126:193–204. doi:10.1085/jgp.200509292
- Zhorov, B.S., and D.B. Tikhonov. 2004. Potassium, sodium, calcium and glutamate-gated channels: pore architecture and ligand action. *J. Neurochem.* 88:782–799.
- Zhorov, B.S., E.V. Folkman, and V.S. Ananthanarayanan. 2001. Homology model of dihydropyridine receptor: implications for L-type $\text{Ca}(2+)$ channel modulation by agonists and antagonists. *Arch. Biochem. Biophys.* 393:22–41. doi:10.1006/abbi.2001.2484
- Zong, Y., S.K. Mazmanian, O. Schneewind, and S.V. Narayana. 2004. The structure of sortase B, a cysteine transpeptidase that tethers surface protein to the *Staphylococcus aureus* cell wall. *Structure.* 12:105–112. doi:10.1016/j.str.2003.11.021