

Common Mechanism of Pore Opening Shared by Five Different Potassium Channels

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ABSTRACT A fundamental question associated with the function of ion channels is the conformational changes that allow for reversibly opening/occluding the pore through which the cations permeate. The recently elucidated crystal structures of potassium channels reveal similar structural motifs at their pore-forming regions, suggesting that they share a common gating mechanism. The validity of this hypothesis is explored by analyzing the collective dynamics of five known K^+ channel structures. Normal-mode analysis using the Gaussian network model strikingly reveals that all five structures display the same intrinsic motions at their pore-forming region despite the differences in their sequences, structures, and activation mechanisms. Superposition of the most cooperative mode profiles shows that the identified common mechanism is a global corkscrew-like counterrotation of the extracellular and cytoplasmic (CP) regions, leading to the opening of the CP end of the pore. A second cooperative mode shared by all five K^+ channels is the extension of the extracellular and/or CP ends via alternating anticorrelated fluctuations of pairs of diagonally opposite monomers. Residues acting as hinges/anchors in both modes are highly conserved across the members of the family of K^+ channel proteins, consistent with their presently disclosed critical mechanical role in pore gating.

INTRODUCTION

The functions of ion channels include establishing a resting membrane potential, controlling cell volume, and regulating flow of ions across epithelial cells (1). Diseases caused by mutations in ion channels, termed as channelopathies, potentially impair cell-cell communication and lead to neurodegenerative disorders or muscular diseases such as multiple sclerosis (2). The molecular conformational mechanisms underlying the (dys)function of ion channels remain, however, elusive.

Membrane proteins are typically composed of three domains: the extracellular (EC) domain exposed to the periplasm, the intracellular/cytoplasmic (CP) domain buried in the cytoplasm, and the transmembrane (TM) domain embedded in the lipid bilayer. The TM domain is composed of a bundle of helices, contributed by four identical monomers in potassium (K^+) channels, which encloses a pore, or a channel, through which ions are conducted. The pore regions of most K^+ channels are considered to have similar structural characteristics: a narrow selectivity filter at the EC side, followed by a large cavity in the middle, and ending in a long gating region that connects to the CP region.

Potassium channels are $\sim 10^4$ times more selective to K^+ ions than to Na^+ ions (3). This propensity for K^+ is attributed to the selectivity filter residues (GYG) also known as the “signature sequence” conserved in both eukaryotic and prokaryotic K^+ channels (3). One of the fundamental questions associated with the gating of K^+ channels is the location of the activation gate and the molecular conformational

changes that ensure a reversible opening/occlusion of the pore.

The bacterial K^+ channel from *Streptomyces lividans* (KcsA) is the first K^+ channel crystallized and structurally determined (3). Since then, several more K^+ channel structures have been determined, all tetramers like KcsA, albeit in different conformations and with different structural topologies, including MthK (4), KirBac 1.1 (5), KvAP (6), and Shaker (7). The crystal structures of MthK and KvAP are considered to be in the “open” form, whereas those of KcsA and KirBac are “closed”. The organisms, resolutions, and Protein Data Bank (PDB) (8) identifiers associated with these five structures are listed in Table 1, along with the size of their monomers and the residue numbers corresponding to the signature sequence GYG in each case.

Sequence alignment of these five structurally known K^+ channels reveals 18–28% sequence similarity with KcsA in the pore-forming domain (Fig. 1 *a*). The pore regions also share a common structural topology (Fig. 1 *b*), mainly four pairs of transmembrane (TM) helices contributed by each monomer, denoted as TM1 (*outer, yellow*) and TM2 (*inner, green*) helices, connected by the P-loop region (*red*) on each monomer. The P-loop regions each consist of a short P-helix, the selectivity filter, and the turret (Fig. 2). Gating is activated by different factors in these K^+ channels: KcsA is gated by pH change (3), MthK by intracellular Ca^{2+} binding (4), KirBac by phosphatidylinositol 4,5-bisphosphate (PIP₂) modulation (9) and ATP binding at the cytoplasmic β -domains (5), KvAP (6) and Shaker (7) by voltage difference across the membrane. KirBac and MthK have an additional CP domain attached to their TM domain. Their monomers contain two TM helices each, like KcsA. KvAP (6) and Shaker (7), on the other hand, contain six TM helices: they

Submitted January 4, 2006, and accepted for publication February 14, 2006.

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0006-3495/06/06/3929/12 \$2.00

doi: 10.1529/biophysj.105.080093

TABLE 1 Potassium channels structurally characterized to date

Protein	(reference)	PDB code*	Organism	Resolution (Å)	N†	GYG‡
KcsA§	(54)	1k4c	<i>Streptomyces lividans</i>	2.0	103	77–79
MthK	(4)	1lnq	<i>Methanobacterium thermoautotrophicum</i>	3.3	80	61–63
KirBac	(5)	1p7b	<i>Burkholderia pseudomallei</i>	3.5	94	112–114
KvAP	(6)	1orq	<i>Aeropyrum pernix</i>	3.2	106	198–200
Shaker	(7)	2A79	<i>Rattus norvegicus</i>	2.9	109	376–378

*For a discussion of the Protein Data Bank, see Berman et al. (8).
†Number of residues (N) refers to those in the pore-forming regions of the monomers, which have been included in the computations.
‡Residue numbers corresponding to the signature sequence GYG in each PDB file.
§The coordinates of KcsA in the high-resolution K⁺ channel/Fab complex determined by Zhou et al (54) are considered in this study.

are known as 6TM structures, although their pore-forming region is structurally similar to that of KcsA (Fig. 1 *b*). The structural similarity at the pore region suggests a similar gating mechanism shared by all K⁺ channels, although the validity of this hypothesis needs to be tested. More importantly, the conformational motions that underlie the gating of the pore for K⁺ conduction remain to be elucidated. This study aims at answering these fundamental questions.

Several experimental studies have been carried out to date, to elucidate the conformational changes associated with pore opening in K⁺ channels (9–14). Prominent among these are the studies by Perozo and co-workers (13,14), where the molecular events associated with the activation of KcsA were investigated by site-directed spin-labeling and electron paramagnetic resonance spectroscopy. Based on their results, Perozo and co-workers suggested that when viewed from the EC side, the M2 helices rotate in a counterclockwise direction while swinging away from the permeation pathway, thus increasing the diameter of the pore cavity (13). This motion was pointed out to be accompanied by that (counterclockwise rotation) of the M1 helices. These experiments also indicated the intrinsic rigidity of the selectivity filter as compared to the flexibility of the M2 helices, particularly toward the CP side.

Although these structural and spectroscopic studies provide insightful details on the channel-pore interactions, the precise conformational changes that accompany the opening/closing of the channel during gating and their molecular basis remain elusive. The increasing experimental data on K⁺ channel structures now provides a foundation for structure-based computational studies. The availability of the atomic coordinates of KcsA has indeed generated a plethora of computational studies (15–27), and with the additional K⁺ channel structures elucidated in the last two years, a comparative analysis of cooperative motions near the pore region can potentially provide new insights, which is the focus of this study.

The conventional computational approach for exploring the dynamics of KcsA has been to perform molecular dynamics (MD) simulations at atomic scale with explicit solvent and lipid molecules. Due to the large size of the system (typically >50,000 atoms), the generated trajectories are typically restricted to short times, of the order of nanoseconds. As such, MD simulations, including our previous work (15,16,20,26), have been unable to elucidate the collective structural changes involved in channel gating, which are at least one to two orders of magnitude slower. Many targeted or steered MD simulations have been resorted to, to accelerate the simulations and explore the functional motions of

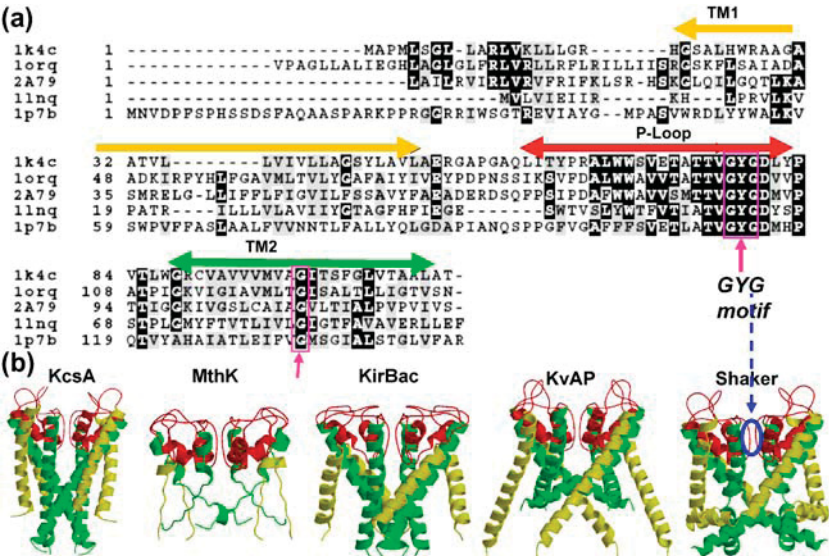
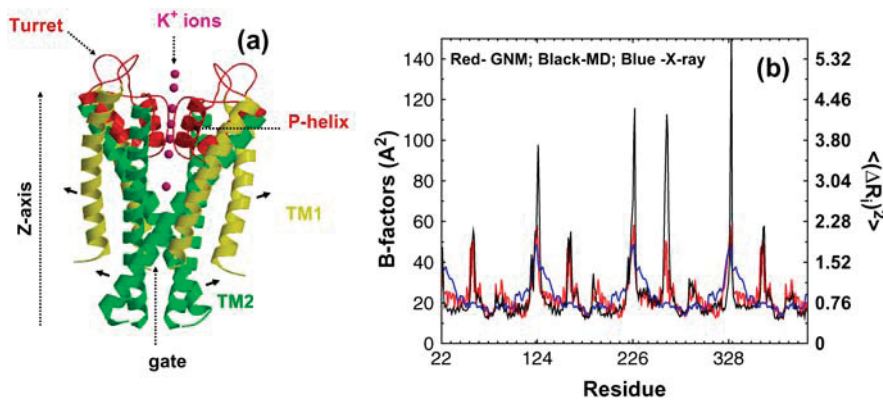


FIGURE 1 Sequence and structure of the pore region of five structurally known potassium channels. (*a*) Alignment of the pore-region sequences. See Table 1 for the nomenclature and source. The regions corresponding to the helices TM1 and TM2 and the P-loop are indicated by yellow, green, and red arrows, respectively. The alignment was performed using ClustalW (63). Fully or highly conserved regions are shown in black, moderately conserved in gray. Two regions of interest are the signature motif GYG at the selectivity filter, and the conserved G on TM2, both enclosed in magenta boxes (see also Fig. 2 *a*). (*b*) Structural comparison of the pore-forming regions of the K⁺ channels aligned in *a*. These are all tetrameric structures, the monomers of which contain either two TM helices (KcsA, MthK, and KirBac), colored yellow (TM1) and green (TM2), or six TM helices (KvAP and Shaker), denoted as S1–S6. Only the pore-forming helices, S5 and S6, equivalent to TM1 and TM2, are displayed here, along with the P-helix/loops region, colored red.



vidual residues, which are related as $B_i = 8\pi^2 \langle \Delta R_i^2 \rangle / 3$, where i is the residue index (abscissa). The three curves describe the results obtained from GNM analysis (red), MD simulations (black), and x-ray crystallography (blue). The correlation coefficients between experimental data and theoretical results are 0.94 (GNM) and 0.83 (MD), and that between the two theoretical results is 0.93.

K⁺ channels. In particular, the closed and open structures of KcsA and MthK, respectively, have been used as end points to model the gating mechanism (23,24). Biggin and Sansom simulated the opening of KcsA by gradually increasing the volume of a particle at the putative CP gate region (23). Tikhonov and Zhorov applied lateral forces on the inner ends of the M2 helices (see below) to open up the pore (24). There have been examples of notable successes in predicting the KcsA ion-binding sites (25,26) and in estimating the free-energy profile that facilitates ion permeation through the selectivity filter (27). Yet, the large-scale motions that cooperatively engage the entire molecule, and in particular those instrumental in gating, remained beyond the reach of MD simulations in general.

Another computational approach that has found wide applications in recent years for exploring long-time/large-scale dynamics is normal-mode analysis (NMA) with coarse-grained models. NMA has been applied to proteins since the early 1980s (28–30), but its utility as an efficient approach for a first assessment of functional motions has been fully recognized only recently, after verifying in many applications the strong correlation between the most cooperative (usually the lowest-frequency, also called global) modes of motions predicted by NMA and the functional motions inferred from experiments (31–33). A recent NMA by Ma and co-workers already demonstrated its utility for providing insights into the gating mechanism of KcsA (34). Given the insensitivity of global modes to detailed atomic interactions (35), reduced models (e.g., elastic network models) have been adopted for identifying such intrinsic dynamics, starting from the Gaussian network model (GNM) (36,37), built on statistical mechanical theories introduced for polymer networks (38), which led the way to a wealth of applications and extensions (32,33,39–52).

In this work, the equilibrium dynamics of KcsA, KirBac, MthK, KvAP, and Shaker (see Fig. 1) are analyzed using elastic network models. We focus on the slowest nonzero

FIGURE 2 Structure and dynamics of KcsA. (a) Ribbon diagram of the KcsA, based on the PDB structure deposited by Doyle et al. (3). Each monomer contains two TM helices, TM1 (outer; yellow) and TM2 (inner; green). The flow of K⁺ ions across the selectivity filter based on MD simulations (15) is schematically shown. Other regions of functional importance are the gate near the intracellular region and the turret at the extracellular surface. The short black arrows indicate the movements suggested by spin labeling experiments (13) for gate opening. The figure was made using Molscript (64). (b) Equilibrium fluctuations as a function of residue index. Left and right ordinates refer to the B-factors and mean-square fluctuations, respectively, of indi-

modes predicted by the GNM and the anisotropic network model (ANM) (41), and demonstrate that all five K⁺ channels possess the same type of intrinsic global dynamics for regulating the pore opening/closing at the putative intracellular gate. The most cooperative mechanism of conformational change invariably predicted for all five structures is a global torsion, manifested by the counterrotations of the EC and CP regions around the cylindrical axis, similar to a corkscrew mechanism, and simultaneously inducing an enlargement of the pore at the CP end. The second most cooperative mode is an alternating expansion/contraction of EC and/or CP ends via anticorrelated fluctuations of oppositely located pairs of monomers. This type of motion is likely to facilitate the binding and access of toxins to the central constriction zone. The results are shown to explain and complement the experimental observations of Perozo and co-workers (13,14) and permit us to identify the key sites/residues that control the functional mechanics of the potassium channels.

MATERIAL AND METHODS

GNM

Detailed description of the GNM can be found in the literature (33,36). We discuss here only the salient features of the model.

The structure is represented by a network of N nodes identified by the α -carbons. The pairs of nodes within a cutoff distance R_c of 7.0 Å are assumed to be connected by uniform springs of force constant γ , representative of the interactions that stabilize the native fold. The interresidue contact topology is fully defined by the $N \times N$ Kirchhoff matrix, the elements of which are defined as

$$\Gamma_{ij} = \begin{cases} -1 & i \neq j, \text{ and } R_{ij} \leq R_c \\ 0 & i \neq j \text{ and } R_{ij} > R_c \\ -\sum_j \Gamma_{ij} & i = j \end{cases} \quad (1)$$

The inverse of Γ provides information on the mean-square (MS) fluctuations $\langle \Delta R_i^2 \rangle$ of residues (diagonal elements) and their cross-correlations

(off-diagonal elements), in a manner similar to the inverse of the Hessian in NMA, i.e.

$$\langle (\Delta R_i)^2 \rangle = (3k_B T / \gamma) [\Gamma^{-1}]_{ii}, \quad (2)$$

where T is the temperature and k_B the Boltzmann constant.

Global modes of motion

A major utility of the GNM is the decomposition of the fluctuation spectrum into a set of normal modes, or the extraction of the slow (or global) modes with minimal computational cost. The contribution of the k^{th} nonzero mode ($1 \leq k \leq N - 1$) to $\langle (\Delta R_i)^2 \rangle$ is given by (36)

$$[(\Delta R_i)^2]_k = (3k_B T / \gamma) \lambda_k^{-1} [\mathbf{u}_k]_i^2, \quad (3)$$

where \mathbf{u}_k and λ_k are the respective k^{th} eigenvector and eigenvalue of Γ . λ_k scales with the frequency of mode k , and the i^{th} element of \mathbf{u}_k , $[\mathbf{u}_k]_i$, provides a measure of the displacement of residue i along the k^{th} mode coordinate. In particular, $[\mathbf{u}_1]_i$ reflects the mobility of residue i in the first (slowest) mode (note that $\lambda_0 = 0$). Its distribution as a function of residue index i is termed the global mode shape.

The slow modes usually have high collectivity (51). Several studies have shown that a small subset of slow modes are usually implicated in molecular motions relevant to biological function (31–33, 39–49, 51–53). The most constrained (or least mobile) residues in these modes play a key mechanical role, such as acting as hinge centers or regulating the concerted movements of entire domains. A major utility of the GNM is indeed the identification of such mechanically critical sites.

Anisotropic network model (ANM)

The GNM provides information on the magnitude of fluctuations, not on their directions, and these fluctuations are isotropic by definition, i.e., $\langle (\Delta x_i)^2 \rangle = \langle (\Delta y_i)^2 \rangle = \langle (\Delta z_i)^2 \rangle = \langle (\Delta R_i)^2 \rangle / 3$. To characterize the directions of motions, we adopt the ANM (41). Γ is replaced therein by the $3N \times 3N$ Hessian matrix \mathbf{H} , the elements of which are found from the second derivatives of the ANM potential

$$V_{\text{ANM}} = (\gamma/2) \sum_i \sum_{j>i} \Gamma_{ij} (R_{ij}^0 - R_{ij})^2 \quad (4)$$

with respect to α -carbon positions. Note that the ANM potential differs from the one implicitly assumed in the GNM,

$$V_{\text{GNM}} = (\gamma/2) \sum_i \sum_{j>i} \Gamma_{ij} (\mathbf{R}_{ij} - \mathbf{R}_{ij}^0) \cdot (\mathbf{R}_{ij} - \mathbf{R}_{ij}^0), \quad (5)$$

in that V_{ANM} depends on the magnitudes of the initial (R_{ij}^0) and final (R_{ij}) interresidue distances, not penalizing the changes in orientation that maintain the interresidue distances, whereas V_{GNM} depends on the vectorial difference. That is, GNM takes account of internal orientational deformations in addition to distance changes. In this respect, the ANM is less realistic than the GNM, but it has the advantage of yielding information, albeit approximate, on the components of $\Delta \mathbf{R}_i$.

Generation of accessible and potentially functional reconfigurations

Conformational changes favored by a given mode k can be assessed using the ANM-predicted eigenvector $\mathbf{u}_k^{\text{ANM}}$ in

$$[\mathbf{R}(\pm s)]_k = \mathbf{R}^0 \pm s_k \lambda_k^{(-1/2)} \mathbf{u}_k^{\text{ANM}}. \quad (6)$$

Here, \mathbf{R}^0 and \mathbf{R} represent the $3N$ -dimensional vectors of the original and instantaneous positions, respectively, of all amino acids; the subscript k refers to the mode that induces the conformational change; and s_k is a parameter uniformly scaling the size of the deformation induced by mode k (52).

Structural details on the 2TM pore

Most K^+ channels are homotetrameric and fourfold symmetric around the pore (cylindrical, z) axis. The pore is surrounded by four pairs of TM helices, called TM1 and TM2 in KcsA. As illustrated for KcsA (Fig. 2 *a*), the outer helices (TM1) are exposed to the hydrophobic lipid environment, whereas the inner helices (TM2) face the pore. The entire length of the pore is ~ 34 Å. The P-loop is comprised of the selectivity filter, the P-helix, which spans only the upper half of the bilayer, and the exposed loops, known as the turret (3). The four P-loops together form the EC vestibule, enclosing a narrow selectivity-filter region ~ 10 Å in length, which opens up into a large central cavity of ~ 10 Å in diameter. This central cavity is connected to the cytoplasm by a hydrophobic pore ~ 18 Å in length, containing the putative gate region. The pore-forming regions of MthK, KirBac, KvAP, and *Shaker* have topology similar to that of KcsA (Fig. 1 *b*). No ions were included in our model, since the global modes are insensitive to the inclusion or exclusion of single nodes.

MD simulations

MD simulations of KcsA (1k4c) were carried out in a lipid bilayer solvated with explicit water molecules and three potassium ions in the selectivity filter and the cavity using GROMACS (53). An equilibration run of 300 ps was performed first, during which the protein backbone atoms were restrained by a harmonic potential and the lipid (phosphatidylethanolamine) and water molecules were allowed to relax. This was followed by a production run of 7 ns, during which the position constraints on the protein were removed.

RESULTS AND DISCUSSION

Equilibrium fluctuations

Before proceeding to the characterization of the global modes of motions, it is of interest to assess the applicability of the GNM insofar as the equilibrium dynamics of the ion channels is concerned. To this aim, we compared the distributions of residue fluctuations predicted for KcsA by the GNM with 1), experimental data (x-ray crystallographic B-factors), and 2), results from full atomic MD simulations. The experimental B-factors $B_i = 8\pi^2 \langle (\Delta R_i)^2 \rangle / 3$ are taken from the PDB file 1k4c (54) (Table 1); and the fluctuations from MD simulations are computed from trajectories of 7 ns with explicit solvent and lipid molecules, using the averages over 140 snapshots taken at 50-ps intervals.

Fig. 2 *b* compares the B-factors (*left ordinate*) or corresponding ms fluctuations (*right ordinate*) obtained for KcsA from GNM analysis (*red curve*), MD simulations (*black*) and x-ray crystallographic measurements (*blue*). The GNM curve is obtained using Eq. 2, with $\gamma = 4.82$ kcal/mol Å². Note that the absolute value of γ does not affect the profile, but uniformly rescales all fluctuations.

The correlation coefficients between the results for the respective pairs GNM/MD, GNM/x-ray, and MD/x-ray are 0.93, 0.94, and 0.83. The GNM results indeed exhibit an even better agreement with experimental data than that exhibited by MD simulations. The higher performance of GNM deserves attention given that this occurs despite the facts that 1), the GNM is a coarse-grained model that does not include any specific interaction at the atomic scale, whereas these are

included in MD force fields and simulations, and 2), the experimental data may also contain biases/errors due to static disorder or intermolecular contacts. One explanation is that the GNM results, being a unique analytical solution for the given structure, are devoid of any sampling inaccuracy that MD simulations may incur. One may indeed notice that the GNM yields identical results across all four monomers in accord with the fourfold symmetry of the structure, whereas the MD simulations yield different fluctuation behavior for the monomers due to random/incomplete sampling. A second explanation is that the GNM contains contributions, albeit approximate, from the complete spectrum of equilibrium modes, including, in particular, the slowest modes that involve the entire structure, which cannot be sampled within the 7-ns MD simulations.

The above analysis supports the utility and accuracy of the GNM in so far as the equilibrium fluctuations are concerned, but does not provide any information different from that already observed by experiments. We now proceed to the spectral decomposition of GNM dynamics to gain insights into the underlying dominant/cooperative modes that are potentially relevant to biological function.

Cooperative/global motions

Two mechanisms of global motions have been identified for all five examined structures, referred to hereafter as motions of types I and II, respectively. The motion of type I is doubly degenerate, i.e., it results from the combination of two symmetrically related modes with identical frequency; type II motion is nondegenerate. Although both modes are proposed to be functional, the second (type II) directly ensures pore opening, as described below.

Distribution of residue displacements driven by global modes

Fig. 3 presents an overview of the shapes of the global modes of type I (*left panels*) and both types (*right panels*) for the tetramer (*left*) or a monomer (*right*). The curves are the normalized distributions (of profiles) of residue mobilities (square displacements) induced by the particular modes (see Eq. 3). The left panels exhibit the fourfold symmetry of the motions consistent with the homotetrameric structures.

Two important observations are made with regard to the shape of the modes in Fig. 3. The minima in global mode of type I precisely coincide with the signature motif GYG (Table 1) of the ion channels in all five cases. The minima in global mode profiles have been shown in several studies to point to the mechanically critical sites of the examined structures (32,33). Strikingly, here the selectivity filter and, in particular, the signature sequence GYG, emerges as the most severely constrained element of the structure in all five structures. The position of the GYG motif is indicated by the “x” signs along the abscissa.

The selectivity filter residues are distinguished by their low mobilities not only in the motion of type I, but also in

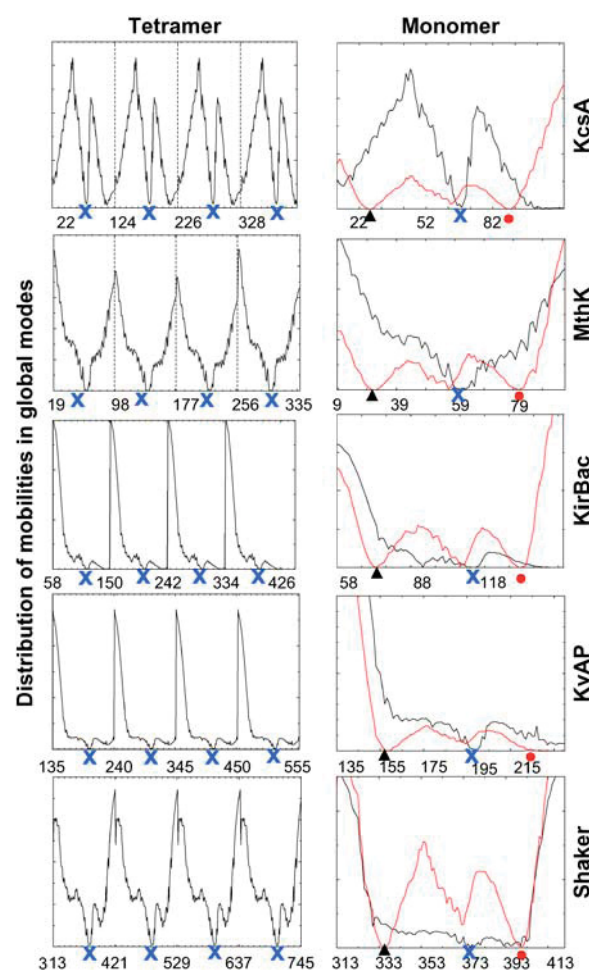


FIGURE 3 Residue mobilities induced by two dominant mechanisms of global motion. The panels describe the square displacements of residues driven by the two most cooperative (lowest-frequency) motions predicted by the GNM, referred to as global motions of type I and II, for all five structures examined (see Fig. 1 *b*). The left panels display the results for motion I, for the entire tetramer. This motion results from the combination of two degenerate modes that each activate the diagonally positioned pairs of monomers (see Fig. 3). The right panels display the mobilities induced by both modes I (black) and II (red). Results are displayed for monomers only, as they are repeated across all four monomers. Minima indicated by blue “x” and red dot symbols on the abscissa coincide with the signature sequence GYG, and the conserved Gly residues on the TM2 helices, respectively (Fig. 1 *a*). Those indicated by solid triangles on the right refer to small hydrophobic residues at central positions on TM1 helices, which closely interact with the conserved Gly on TM2 helices. These three sets define the key sites that act as hinges/anchors in coordinating the two global mechanisms.

type II motions, as can be inferred from the minima in the red curves in the right-hand panels. It is noteworthy to mention that our MD simulations of KcsA in an explicit solvent-bilayer environment (15) also revealed negligibly small fluctuations at the selectivity filter region. The high stability of the selectivity filter backbone is presumably a biological requirement for precise functioning (selective conduction) of K⁺ channels.

The second observation of interest relates to the minima in the modes of type II. These positions indicated by the red dots in the right-hand panels in Fig. 3 refer to centrally located residues in the TM1 and TM2 helices. The minima on TM2 helices are located at, or in the close vicinity of, the conserved Gly mentioned above, mainly at G⁹⁹ (KcsA), G⁸³ (MthK), G¹³⁴ (KirBac), A²²³ (KvAP), and L⁴⁰⁰ (*Shaker*). The occurrence of a glycine at a slow mode minimum suggests that this residue may serve as a “hinge” site for a global motion. Furthermore, the conservation of these glycines across all five ion channels (Fig. 1 *a*) suggests a conserved mechanism of motion (4). The minima on the TM1 helices (shown by the solid triangles along the abscissa in Fig. 3), on the other hand, occur at small hydrophobic residues—V³⁷ (KcsA), I³² (MthK), V⁷³ (KirBac), A¹⁵⁶ (KvAP), and L³³⁵ (*Shaker*)—that are spatially close to the conserved Gly on TM2. The small size of these residues allows for close van der Waals interactions and rotational flexibilities. These results suggest that these TM1 residues are possibly implicated, along with the conserved Gly on TM2 helices, in a pivotal role in the global motion of type II.

Visualization of mobilities

A better assessment of mobilities is possible by mapping the results in Fig. 3 into color-coded ribbon diagrams. Fig. 4

illustrates the results for KcsA, KirBac, and *Shaker*. The colors range from red (most mobile) to blue (static). The panels in the left two columns illustrate the results for each of the doubly degenerate modes of type I, and those in the right-most column refer to mode type II.

The pairs of diagonally opposite monomers exhibit identical fluctuation behavior in mode type I (Fig. 4, *left column*), and the roles of the pairs are inverted in the accompanying second degenerate mode (*middle column*). As will be shown below, in type I motion, a given pair of oppositely positioned TM1 helices move away from the pore region, giving rise to a distortion from circular symmetry into an ellipse when viewing the cross-sectional area while the other pair is stable, and vice versa. The motions are enhanced toward the EC region and peak at the turrets in KcsA, whereas a higher mobility toward the CP region is observed in KirBac and MthK (the latter not shown). *Shaker* and KvAP (the latter not shown) exhibit high mobilities at both ends. The cross-correlations and ANM results below show that the diagonally positioned pairs of monomers are subject to anticorrelated (i.e., concerted, but opposite-sense) fluctuations, allowing for the opening of the EC and/or CP ends via concerted, alternating movements of the pairs of monomers.

Motion of type II, on the other hand, emerges as a very robust mode, identically recruited by all five structures. The

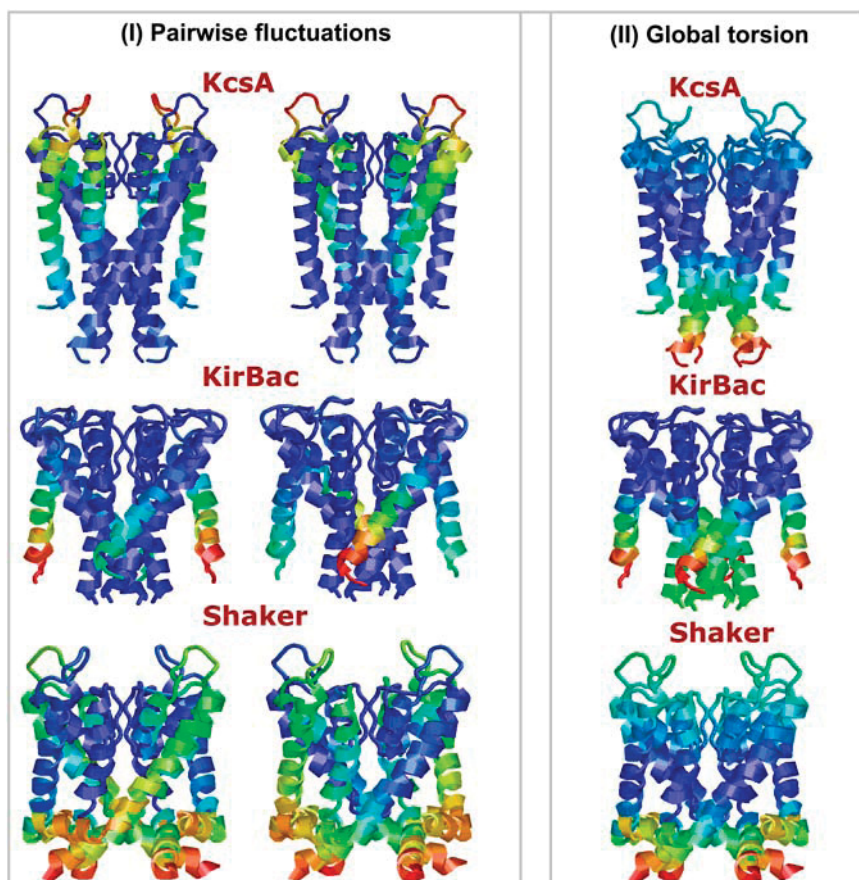


FIGURE 4 Mobilities in global modes shown by color-coded diagrams for KcsA, KirBac, and *Shaker*. The slowest global mode (mode type I) is twofold degenerate. The corresponding ribbon diagrams are shown on the left and middle columns. The second-lowest-frequency mode (mode type II) is illustrated in the right column. Color code is red, orange, green, cyan, and blue in order of decreasing mobility. Note that mode type I involves the pairwise motions of oppositely positioned monomers, whereas mode type II is cylindrically symmetric.

four monomers undergo the same fluctuation behavior in this mode, i.e., the motion is fourfold (or cylindrically) symmetric. This motion, amplified toward the CP end of the ion channels, will be shown below to be a global torsion that controls the gating of the pore.

The most cooperative (lowest-frequency) modes computed here therefore reveal that despite their sequence, structure, and activation mechanism dissimilarities, all five potassium channels bear a close resemblance to each other insofar as their global motions are concerned, and it will be shown below that the global mode of type II is directly involved in pore gating.

Master curves for all five K⁺ channels global dynamics

Although the mode shapes in Fig. 3 appear to be somewhat different, their superimposition after correcting for the residue insertions/deletions indicated by their sequence alignment (Fig. 1 *a*) leads to the master behavior of all ion channels displayed in Fig. 5. Fig. 5, *a* and *b*, refers to the global modes of types I and II, respectively. The residue numbers of MthK are used as reference along the abscissa.

Major features in these master curves are summarized as follows.

Insofar as the collective motions of type I (Fig. 5 *a*) are concerned:

All five structures exhibit a definite preference for severely constraining the signature motif GYG as a global minimum, irrespective of the differences in the mobilities of the other regions.

KcsA is distinguished by the peaks near the turrets and lower mobility toward both N- and C-termini, whereas all others exhibit increased mobilities toward their N-termini (at the CP region).

The C-terminus (i.e., CP ends of the TM2 helices), on the other hand, shows a mixed behavior. KcsA, KirBac, and KvAP are stable at this region, whereas MthK and Shaker show increased mobilities.

Insofar as the collective motions of type II (Fig. 5 *b*) are concerned:

All five chains again exhibit a high stability at the selectivity filter region.

A minimum is observed at a central position in both the TM1 and TM2 helices in all five structures, revealing a change in the direction of collective motion at a central plane. This inversion will be shown below to refer to a cross-sectional plane almost bisecting the TM helices. The minima on the TM2 helices exactly coincide with, or are close neighbors to, the conserved Gly residues pointed out in previous experiments to be functionally important in KcsA (G⁹⁹) (13) or to form a kink (e.g., G⁸³ in MthK (4)). Those on TM1 helices, on the other hand, occupy a position spatially close to these kink sites,

suggesting a possible bending of all helices, in addition to global inversion of the rotation direction, at the same central plane.

As will be shown below, the global motion of type II is mainly responsible for gating the pore. Indeed, this combination of global torsion with outward bending of the TM1 and TM2 helices' cytoplasmic halves, which will ensure the cooperative pore opening, is consistent with experimental observations.

Mechanisms of global motions and relevance to gating

The mechanisms of the global motions of type I and II are elucidated by ANM analysis.

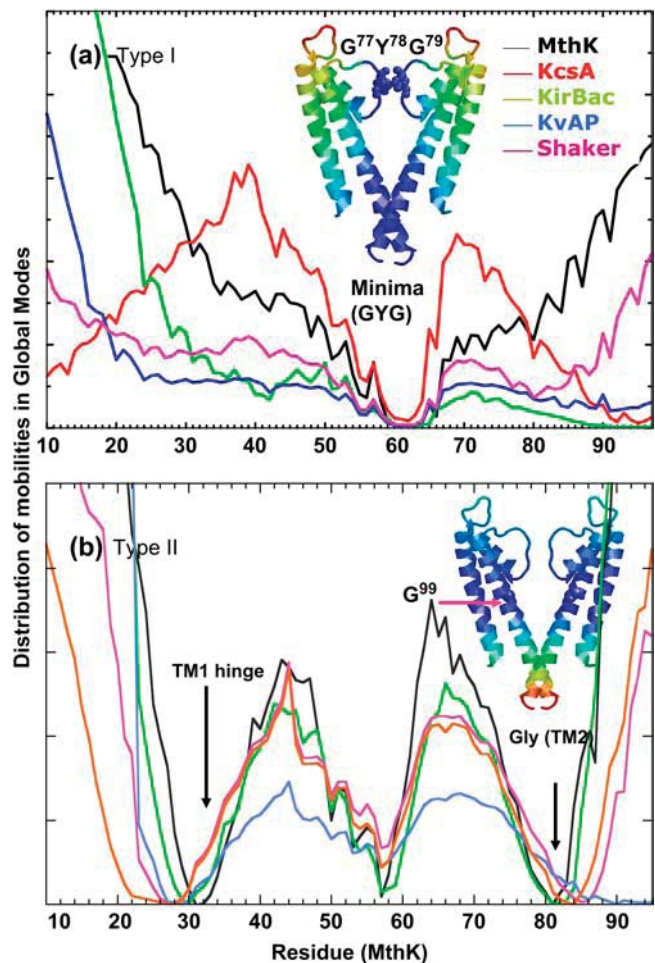


FIGURE 5 Master curves for global dynamics of potassium channels. (*a*) Superposition of the most cooperative global mode shape (type I) for KcsA (red), KirBac (green), KvAP (blue), Shaker (magenta), and MthK (black), plotted for one monomer. The abscissa refers to the residue indices of MthK. (Inset) Ribbon diagram of KcsA, with two monomers removed for clarity, highlighting the selectivity-filter residues GYG, in space-filling representation. (*b*) Superposition of the global mode shape (type II) for the same set of proteins. The abscissa refers to the residue indices of MthK. (Inset) Ribbon diagram of KcsA, with two subunits removed for clarity, highlighting the conserved glycine on TM2 in space-filling representation.

We first focus on KcsA. The strategy is as follows: we identify the ANM modes that are equivalent to the above-determined GNM modes of types I and II by comparing the eigenvectors derived from $\mathbf{\Gamma}$ and \mathbf{H} , and sorting pairs that yield the most correlated square displacements. ANM thus permits us to compute the x , y , and z components of the fluctuations driven by the global motions of type I and II, or the deformation vectors, which upon substitution in Eq. 6 permit us to generate the “deformed” states, or the alternative conformations sampled by the actions of modes I and II. These alternative states are consistent with the notion of the native (macro) state as an ensemble of microstates defined by the normal fluctuations in the neighborhood of the known structure.

Fig. 6 illustrates the alternative conformations sampled by KcsA near its original (PDB) structure. Two different pathways of reconfiguration are shown, referring to global modes of types I and II. Fig. 6 *a* displays the side views of the native state and two conformations favored by mechanisms I and II. Fig. 6 *b* displays the top views of the same structures. The central structure is the original structure, and the four alternative forms are again those sampled by mechanisms I and II, but this time we display both the positive and negative departures away from the original state by adopting the scaling parameters of $\pm s_I = 75$ or $\pm s_{II} = 100$ for visualizing the respective modes (see Eq. 6). This parameter scales the size of the motion without affecting its mechanism.

We note that positive and negative fluctuations are equally probable, by definition, in the immediate neighborhood of the global energy minimum in NMA. Larger-scale deformations may entail a preference in one direction or another due to the anharmonicity of the energy landscape. This study sheds light on the initial events only, in the close neighborhood of the native state.

The emerging mechanisms of motions are as follows:

Motion of type I involves the concerted anticorrelated opening/closing of the EC ends of the diagonally positioned pairs of monomers. As a given pair of monomers move away from each other, the opposite pair come close, such that the cross-sectional area viewed from the top oscillates between a circular (native) and elliptic (deformed) shape, the long axis of the ellipse alternating between the two perpendicular directions, depending on the “stretched” monomers. The pore region remains practically fixed during this motion, whereas the EC ends of the molecule, and in particular the turrets, undergo large swinging motions.

Motion of type II is a global torsion. During this cooperative motion, the EC regions of the molecule rotate counterclockwise, whereas the CP domains rotate clockwise, and vice versa. The amplitude of the motion becomes larger with increasing distance from a stationary plane that comprises the central parts of TM1 and TM2 helices. As pointed out above, this central

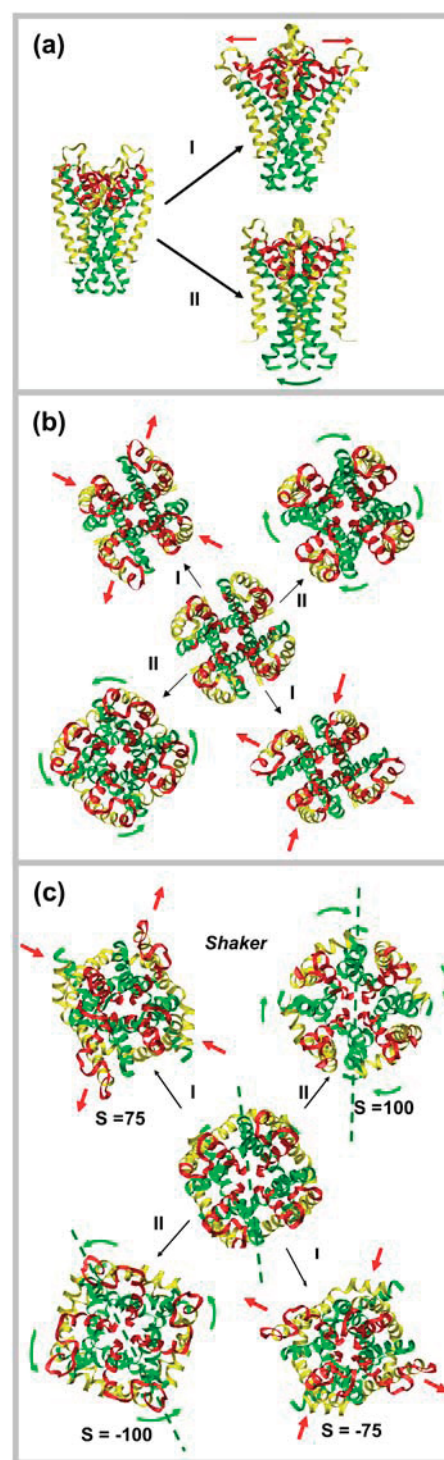


FIGURE 6 Dynamic equilibrium between fluctuating conformations. Ribbon diagrams represent different deformed models for KcsA (*a* and *b*) and Shaker (*c*), generated using Eq. 6 for mode types I and II. (*a*) Side view of deformed model for type I mode for $s_I = -75$ (top) and $s_I = -100$ (bottom). (*b*) Top view of deformed models for mode type I obtained using $s_I = -75$ (top right), $s_I = +75$ (bottom left), and for mode type II, $s_{II} = -100$ (top left), $s_{II} = 100$ (bottom right). The figure in the middle is that of the crystal structure. (*c*) Counterpart of *b* for Shaker.

plane includes the conserved glycines on TM2 and a set of small hydrophobic residues of TM1 helices that form a stable core. The amplification of the motion away from this plane is particularly pronounced toward the CP ends of TM2 helices, and confers a remarkable expansion and mobility at the intracellular gate region, whereas the selectivity filter remains fairly rigid. The pore enlargement is induced by the rotations in either direction, as can be discerned in Fig. 6 *b*. This structure-induced ability of KcsA to open the pore via concerted tilting and rotation of the TM1 and TM2 helices was also pointed out by Ma and co-workers (34) in their insightful NMA of KcsA intrinsic dynamics. Interestingly, the same mechanisms of motions are observed here not only for KcsA, but across all five potassium channels structurally characterized to date. The counterpart of Fig. 6 *b* for *Shaker* is given in Fig. 6 *c*.

Pore enlargement induced by mode II: a closer view of the constriction zone

The expansion of the pore region induced by mode II can be directly viewed by plotting the radius of the central constriction zone as a function of the position along the *z* axis. The black curve in Fig. 7 displays the profile for the original (KcsA) structure. As illustrated in Fig. 7 *b*, the selectivity-filter is the region with the lowest (~ 1 Å) radius, shown in red, succeeded by the pore region of 1.5–2 Å radius (green). The central cavity enjoys significant radial enlargement (up to 4.5 Å, blue region), as does the EC end of the molecule. The fluctuating conformations induced by the action of mode II, on the other hand, lead to the solid and dotted curves shown in red in Fig. 7 *a*, which refer to the respective global torsions in opposite directions (positive and negative deformations induced by using $\pm s_{II}$). The selectivity filter remains practically unchanged during this reconfiguration, whereas there is a significant increase in the internal radius at the position of the intracellular gate. The pore-region diameter widens up to 7 Å using $\pm s_{II} = 100$. This behavior is illustrated by the lower diagram, where the expansion in the pore region can be visualized. The inset in Fig. 7 *a* also displays the relative position of the CP ends of the TM2 helices (blue and red, respectively, before and after global torsion). For simplicity, two opposite monomers are displayed here. Interestingly, this diagram shows close resemblance to that obtained by the superposition of the crystal structures of KcsA and MthK (4), supporting the view that the deformed state presently predicted is the open form of KcsA.

Calculations repeated for the four other K⁺ channels showed that the global torsion cooperatively modulating the opening of the pore is commonly shared by all potassium channels. See the movies (Supplementary Material, data 1–4) for an animated visualization of the conformational fluctuations associated with mode II for KcsA and *Shaker*.

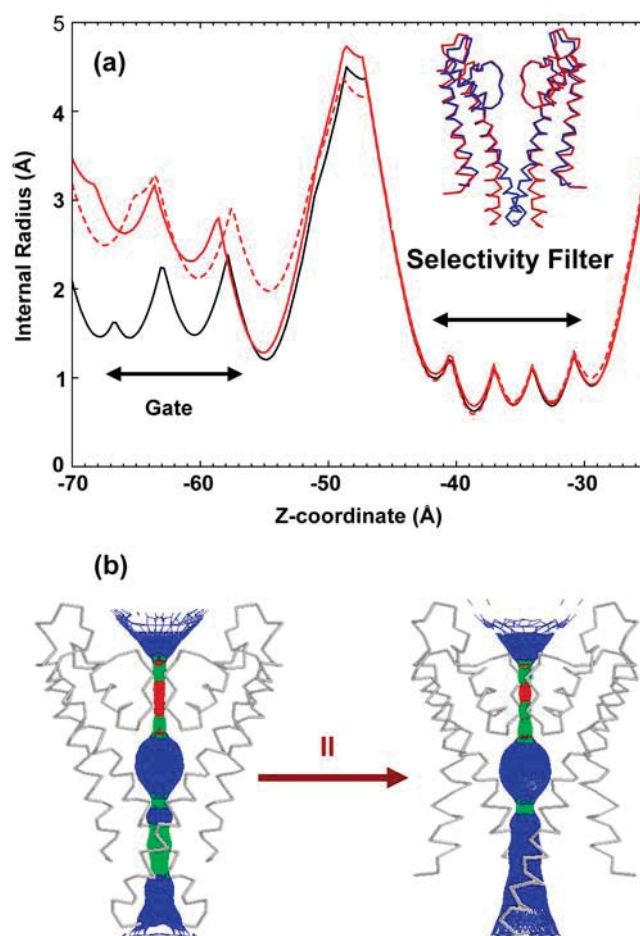


FIGURE 7 (a) Pore-radius profiles as a function of the position along the cylindrical (*z*) axis. The radii are computed for the crystal structure (black) and the deformed conformations at $s_{II} = -200$ (red solid line) and $s_{II} = +200$ (red dashed line). (b) Solid-sphere representation of the inner surface of the channel at the pore region for the crystal structure (left), and for the model of the open form (right) at $s_{II} = -200$, with the corresponding backbone structures superimposed onto the pore surface. The color code for the solid-sphere representation is: red, pore radius < 1.15 Å; green, 1.15 Å $<$ radius < 2.30 Å; and blue, radius > 2.30 Å. The pore-radius profiles were generated using HOLE (65). In the inset of *a* is the backbone of the crystal structure (blue) superimposed onto the model of the open form (red). Two monomers have been deleted for clarity.

Comparison with experimental data

Most K⁺ channels are known to be blocked by large organic cations such as tetrabutylammonium (TBA) from the CP side and tetraethylammonium (TEA) from both sides of the membrane (3). The size of these cations is ~ 7 – 12 Å. The fact that these molecules can enter the pore and bind at the CP end of the selectivity filter suggests that the gate must be opening up very wide to accommodate the cation. The crystal structure of MthK (4) also supports this argument, since in the crystal structure, the TM2 helices are bent away from the pore axis, increasing the diameter of the pore to > 12 Å. The top diagrams in Fig. 8, *a* and *b*, illustrate the experimentally proposed mechanisms for KcsA and MthK gating, respectively

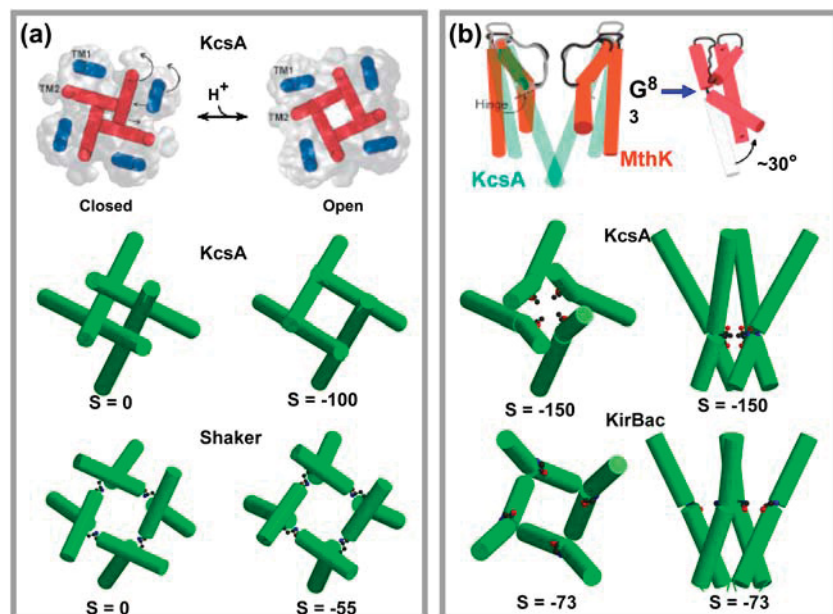


FIGURE 8 Cartoon representation of the TM2 helices of the deformed models of KcsA, KirBac, and Shaker, at different values of scaling parameter (Eq. 6). (a, top row) Cartoon representation of closed and open models for KcsA, proposed as a model for molecular mechanism of gating, based on experimental data (Fig. 7 in Perozo et al. (13)). (a, middle row) TM2 helices in the original structure ($s_{II} = 0$) and in the conformation induced by mode II ($s_{II} = -100$) for KcsA. (a, bottom row) TM2 helices of Shaker in the PDB structure ($s_{II} = 0$) and the predicted structure deformed along mode II ($s_{II} = -55$). (b, top row) Cartoon representation of KcsA (red) and MthK (green) TM structures, illustrating the model proposed by Perozo and co-workers for a “hinge-gating” mechanism at the intracellular gate. G^{83} acts as a hinge (Fig. 1 in Perozo (14)). (b, middle row) KcsA TM2 helices at $s_{II} = -150$, from the top (left) and side (right). Highlighted residue at the kink is T^{107} . (b, bottom row) KirBac TM2 helices at $s_{II} = -73$, from the top (left) and side (right). Conserved glycine (G^{134}) emerging as kink residue is highlighted.

(13,14). Such large conformational changes are also indicated in mutational studies done on Shaker (55). These studies, in fact, propose a similar opening mechanism for both the voltage-gated and voltage-independent K^+ channels. The torsional motion seen here in the most cooperative global mode, for all the proteins, irrespective of the gating ligand, supports the common-gating-mechanism view.

In Fig. 8 a, the opening of the KcsA pore from the closed form to the “open” form predicted by GNM (green diagrams) is clearly seen to be in accord with that proposed by models based on experimental data (13) and crystal structures (14). The diagrams illustrate the relative positions of the four TM2 helices, before deformation ($s_{II} = 0$), or as induced by mode II ($s_{II} \neq 0$). Similar results are presented for Shaker and Kirbac (Fig. 8, a and b, bottom row). In Shaker, since the pore is already open, this mode pushes the intracellular region of TM2 (beyond the Gly “kink”) outward and away from the pore axis.

A striking observation is the appearance of a kink, in addition to an overall enlargement, with increase in the extent of deformation. The kink occurs at T^{107} in KcsA, and at the conserved G^{134} in KirBac1.1 (Fig. 8 b), at $s_{II} = 150$ and 73, respectively. The residue T^{107} , which is a minimum in type I mode in KcsA, was also found to provide a binding site to the ion in the cavity in MD simulations (15); in the NMA of KcsA (34), the residues T^{107} – L^{110} were considered to be at a pivotal position in the gating mechanism. Notably, two mutants, A108S and A108T, have been observed (56) to dramatically increase the population of open KcsA conformers, consistent with the critical importance of this region for mediating pore opening. In addition, mutation of T^{107} in KcsA resulted in dramatic reduction in current (57). The “kink” formation presently predicted at this conserved

residue is in accord with the hypothesis (4) that the region at or around the conserved Gly serves as a “hinge” in the gating mechanism. It should be noted here that the kink in TM2 forms after the pore opens up, not before. Compare, for example, the results for KcsA for different extents of deformation (Fig. 8 a).

Finally, we note that in Shaker, the minima in the type II mode is at L^{400} , which is in close proximity to the “PVP” motif conserved in Kv channels. Cross-linking studies suggest that a distortion may occur at TM2 in Shaker (known as S6 in Kv channels) in the vicinity of the PVP motif (58,59), which lends further support to the physical realism of the predicted motions.

CONCLUSION

The process of channel opening requires structural rearrangement of the ion conduction pathway. The channels show narrow constrictions within the pore, which may be broadening and contracting as the channel gates. Such cooperative conformational changes, though anticipated by experimental results, have never been accessed so far, either by intensive MD simulations of KcsA equilibrium dynamics or by nonequilibrium simulations, except for the NMA conducted by Ma and co-workers (34). A simple analytical model purely based on contact topology/geometry has provided here a scaffold for the conformational changes involved not only at the putative intracellular gate but also at the EC turret region of KcsA. Most importantly, a common pore gating mechanism is revealed by the analysis in this study for all five structurally determined potassium channels. Although these K^+ channels differ in their sequences (Fig. 1 a), structures (Fig. 1 b), and activation

mechanisms, by focusing on their pore-forming region, and optimally aligning their sequence, master curves emerged for two dominant mode shapes shared by all five channels, as presented in Fig. 5.

Mode I is shown to be manifested by anticorrelated fluctuations of oppositely located monomers, to deform the cylindrical symmetry of the structure, and to expose the CP and/or EC regions to facilitate ligand/toxin binding from either end.

Mode II, on the other hand, which is proposed here to be the major mechanism involved in channel gating, is a global torsion, i.e., counterrotation of the two halves of the molecule, simultaneously inducing a substantial broadening in the pore region. This mode further gives rise to the formation of a kink at a conserved glycine in all K⁺ channels, after a certain level of enlargement of the pore region.

Although the TM helices and CP/EC domains undergo significant rearrangements in these global modes, the selectivity filter remains highly stable (see the minimum at the signature sequence GYG in modes I and II), consistent with the required precise operation (selective conduction) of the filter, which should not be disrupted by the global rearrangements that modulate the channel-gating. The atomic structure and interactions at the selectivity filter are indeed highly specific and selective, and cannot be described by the coarse-grained approach presently adopted. Several detailed simulations by other groups have indeed provided valuable insights on the dynamics of the selectivity filter at full atomic level in the presence of ions and water molecules (15,17–19,21). Our view is that the global motions characterized here do not interfere with the subtle operation of the selectivity filter, but regulate the collective rearrangements of the TM helices surrounding the pore region to ensure the predisposition of the structure to channel gating that may be triggered by various mechanisms in different K⁺ channels.

This study provides further evidence for the intrinsic ability of proteins to undergo cooperative changes required for their biological function (32,33,51,52,60–62). The current understanding is that the topology of interresidue contacts, or the overall architecture, defines their equilibrium dynamics, or the cooperative fluctuations that incur the lowest energy ascent in the neighborhood of the original equilibrium state; and these structure-induced fluctuations are those required for, or facilitating, biological functional mechanisms. The slowest modes unraveled by NMA are nothing other than these lowest-ascent pathways away from the global minimum, and due to their collective nature, these modes are relatively insensitive to atomic details, hence the utility of coarse-grained models such as GNM. This type of structure-encoded dynamics and allosteric effects are presumably further enhanced by structural symmetry in the case of multimeric enzymes or receptors (60), and the fourfold symmetric potassium channels are probably no exception to this generic behavior of biomolecular systems.

SUPPLEMENTARY MATERIAL

An online supplement to this article can be found by visiting BJ Online at <http://www.biophysj.org>.

We thank Dr. A. J. Rader and Dr. D. Tobi for insightful discussions. We thank Elsevier for permission to reproduce figures from works by Perozo and co-workers (13,14).

Partial support from National Institutes of Health grants 5R01-LM007994 and 1 P20 GM065805-01A1 is gratefully acknowledged.

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