Voltage-dependent Gating Rearrangements in the Intracellular T1–T1 Interface of a K⁺ Channel

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The intracellular tetramerization domain (T1) of most eukaryotic voltage-gated potassium channels (Kv channels) exists as a “hanging gondola” below the transmembrane regions that directly control activation gating via the electromechanical coupling between the S4 voltage sensor and the main S6 gate. However, much less is known about the putative contribution of the T1 domain to Kv channel gating. This possibility is mechanistically intriguing because the T1–S1 linker connects the T1 domain to the voltage-sensing domain. Previously, we demonstrated that thiol-specific reagents inhibit Kv4.1 channels by reacting in a state-dependent manner with native Zn²⁺ site thiolate groups in the T1–T1 interface; therefore, we concluded that the T1–T1 interface is functionally active and not protected by Zn²⁺ (Wang, G., M. Shahidullah, C.A. Rocha, C. Strang, P.J. Pfaffinger, and M. Covarrubias. 2005. J. Gen. Physiol. 126:55–69). Here, we co-expressed Kv4.1 channels and auxiliary subunits (KChIP-1 and DPPX-S) to investigate the state and voltage dependence of the accessibility of MTSET to the three interfacial cysteines in the T1 domain. The results showed that the average MTSET modification rate constant (kMTSET) is dramatically enhanced in the activated state relative to the resting and inactivated states (~260- and ~47-fold, respectively). Crucially, under three separate conditions that produce distinct activation profiles, kMTSET is steeply voltage dependent in a manner that is precisely correlated with the peak conductance-voltage relations. These observations strongly suggest that Kv4 channel gating is tightly coupled to voltage-dependent accessibility changes of native T1 cysteines in the intersubunit Zn²⁺ site. Furthermore, cross-linking of cysteine pairs across the T1–T1 interface induced substantial inhibition of the channel, which supports the functionally dynamic role of T1 in channel gating. Therefore, we conclude that the complex voltage-dependent gating rearrangements of eukaryotic Kv channels are not limited to the membrane-spanning core but must include the intracellular T1–T1 interface. Oxidative stress in excitable tissues may perturb this interface to modulate Kv4 channel function.

INTRODUCTION

Vital electrophysiological processes in the brain and heart depend on the precise orchestration of intramolecular motions in voltage-dependent K⁺ channels (Kv channels). Current models of Kv channel activation gating propose that the opening of the main gate namely depends on the electromechanical coupling between segments S4 (voltage sensor) and S6 (activation gate), which are membrane-spanning regions of the Kv subunit (Yellen, 1998; Horn, 2000; Lu et al., 2002; Tristani-Firouzi et al., 2002; Bezanilla and Perozo, 2003; Long et al., 2005b). However, recent work has suggested the contribution of the intracellular NH₂-terminal tetramerization domain (T1) to activation gating (Cushman et al., 2000; Minor et al., 2000; Robinson and Deutsch, 2005). Namely, these studies demonstrated that the mutations cause dramatic shifts in the voltage dependence of channel activation. For instance, the T46V mutation in the rat Kv1.2 channel stabilizes the closed state by destroying a buried hydrogen bond network between T46 and D79 in the T1–T1 interface without significantly changing the tertiary structure of the protein (Minor et al., 2000). In contrast, another T1 mutation (N136A) in the Aplysia Kv1.1 destabilizes the closed state and changes the tertiary structure near the central axis of the T1 tetramer (Cushman et al., 2000).

Recent crystallographic studies of the isolated Kv3 and Kv4 T1 domains revealed that the tetrameric...
four-layer scaffold includes four C3H1 high-affinity Zn$^{2+}$ sites in the T1–T1 intersubunit interface (Bixby et al., 1999; Nanao et al., 2003). Surprisingly, however, our recent studies showed that the Zn$^{2+}$ site thiolate groups in Kv4 channels are not protected by Zn$^{2+}$ against chemical modification and that the T1–T1 intersubunit interface may play a role in channel gating (Wang et al., 2005). We have hypothesized that the L4 layer at the membrane side of the T1 domain and the S6 gate may undergo conformational changes associated with voltage-dependent activation. To test the coupling between a putative T1 conformational change and voltage-dependent activation gating, we probed here the state-dependent accessibility changes of the unprotected Zn$^{2+}$ site thiolate groups. Using thiol-specific reagents and patch-clamp electrophysiology combined with a concentration-clamp method, our experiments demonstrated a tight functional coupling between voltage-dependent gating and an apparent conformational change in the T1–T1 interface of the Kv4.1 channel. Moreover, cross-linking experiments supported the idea of a functionally critical and dynamic T1–T1 interface that contributes to channel gating. Altogether, these observations suggest strongly that the complex voltage-dependent gating rearrangements in eukaryotic Kv channels include coupled displacements involving the intracellular T1 domain. Therefore, the activation gating mechanism extends beyond the membrane-spanning core of the pore-forming subunits.

**MATERIALS AND METHODS**

**Molecular Biology**

Kv4.1 (mouse), DPPX-S (rat), and KChIP1 (rat) were maintained in pBluescript II KS, pSG5 (Stratagene), and a modified pBlue-script vector, pBlJS/KSM, respectively. DPPX-S and KChIP1 are gifts from B. Rudy (New York University, New York, NY) and Mark Bowly (Wyeth-Ayerst Research, Princeton, NJ), respectively. Eight Kv4.1 mutants were used in this study (Table I and Fig. 3). All mutations were created using the QuickChange site-directed mutagenesis (Stratagene) and confirmed by automated sequencing (Kimmel Cancer Institute of Thomas Jefferson University). The capped cRNAs for expression in Xenopus laevis oocytes were synthesized using the mMessage mMACHINE kit for in vitro transcription (Ambion).

**Heterologous Expression and Electrophysiology**

Kv4.1 wild-type and mutant channels were coexpressed along with two auxiliary subunits (KChIP1 and DPPX-S) as described previously (Wang et al., 2005). The expression of the Kv4.1 ternary complex was necessary because mutations in the putative Zn$^{2+}$ site yielded nonfunctional channels or inhibited expression profoundly. Our previous paper and an earlier study from the Pfafflin-gar laboratory showed that the apparently lethal phenotype of Zn$^{2+}$ site mutants can be corrected by coexpression of the channels with KChIPs (Kunjilwar et al., 2004; Wang et al., 2005); and we have found that DPPX-S boosts the expression of the channels even further (Wang et al., 2005), which made possible the recordings from inside-out macropatches.

All currents were recorded using an Axopatch 200A amplifier (Axon Instruments). To probe the gating state-dependent accessibility of the thiolate groups of the T1 domain, the membrane-impermeant thiol-specific reagent MTSET (2-trimethylammonium-ethyl-methane-thiosulfonate bromide; Toronto Chemicals) (200–400 μM) was applied to the intracellular side of inside-out macropatches at various membrane potentials (see online supplemental material, available at http://www.jgp.org/cgi/content/full/jgp.200509442/DC1). The composition of the solution in the patch electrodes was (in mM) 96 NaCl, 2 KCl, 1.8 CaCl$_2$, 1 MgCl$_2$, and 5 HEPES (pH 7.4, adjusted with NaOH), and that of the bath solution was (in mM) 98 KCl, 0.5 MgCl$_2$, 1 EGTA, 10 HEPES (pH 7.2, adjusted with KOH). The tip resistance of the borosilicate patch pipette was typically 1–2 MΩ. An online P/4 procedure was applied to subtract the passive leak current and capacitive transients. The currents were filtered at 1–5 kHz and digitized at 5–100 kHz. All recordings were obtained at room temperature (23 ± 1°C).

Data Acquisition and Analysis

Voltage-clamp protocols and data acquisition were controlled by a Pentium-III class desktop computer interfaced to a 12 bit A/D converter (Digidata 1200 using Clampex 8.0; Axon Instruments). Clampfit 8.0 (Axon Instruments) and Origin 7.0 (Origin Lab Inc.) were used for data reduction and analysis. To determine the MTSET modification rate constant, the peak currents were plotted as a function of the cumulative time of exposure to MTSET. The time constant ($\tau$) was computed from the best-fit exponential describing the time course of inhibition by MTSET, and the second-order rate constant $k_{b\text{MTSET}}$ was determined from this relationship: $k_{b\text{MTSET}} = (1/\tau \text{ [MTSET]})$. Data from at least three patches for each measurement are presented as mean ± SEM. The Student’s t-test (unpaired) was used to evaluate statistically significant differences between two groups of data.

Sources of Error

To test whether or not a mutant was inhibited by MTSET, we used continuous intracellular application of the reagent while the current was evoked by a 250-ms step depolarization to +80 mV from a holding potential of −100 mV (3 s, start-to-start) (Wang et al., 2005). The rate constants determined from these experiments were slow and similar to those obtained from measurements in the resting state (Figs. 1 and 2). This similarity is expected because even during continuous application, the channels spend most of the time in the resting state at −100 mV. In general, however, the measurement of very slow MTSET modification rate constants (<50 M$^{-1}$ s$^{-1}$) in the resting state was less accurate. This difficulty resulted from the limited survival of some inside-out patches, which did not always allow the modification to reach steady state. Under continuous application, steady state was reached typically in ~4–5 min. To minimize the error, the constant term of the exponential function (see above) was fixed by assuming the value obtained from those experiments that were long and stable enough to reach steady-state. For wild-type and the mutants C11xA, C12xA, and C13xA, the fractional steady-state level of the inhibition by MTSET ranged approximately between 0.1 and 0.3 (Wang et al., 2005). The slow
rate constants in the resting state were not corrected for the possible loss of some reagent due to hydrolysis during the application period. At neutral pH, the half-life of MTSET is 10–11 min (Karlin and Akabas, 1998).

**Online Supplemental Material**

Rapid reagent application and verification of the switching and exchange processes were performed as described in the online supplemental materials (available at http://www.jgp.org/cgi/content/full/jgp.200509442/DC1).

**RESULTS**

**Kv4.1 Cysteines in the T1 Interfacial Zn$^{2+}$ Site Undergo State-dependent Modification by MTSET in the Presence of Auxiliary Subunits**

To investigate the possible functional coupling between the transmembrane activation machinery and the T1 domain, all but three intracellular Zn$^{2+}$ site cysteines in the T1–T1 interface of the Kv4.1-α subunit were mutated to alanines (Kv4.1-C11xA) (Table I); and the wild-type or mutant Kv4.1-α subunit was coexpressed with the Kv4 auxiliary subunits KChIP1 and DPPX-S (ternary complex) as shown previously (Wang et al., 2005). The functional impact of the remaining cysteines was probed upon chemical modification with a thiol-specific reagent. Fig. 1 shows that internal application of the membrane-impermeant MTSET irreversibly inhibits ternary complexes of wild-type or C11xA channels in the resting, activated, or inactivated state. In contrast, and regardless of the gating state of the channel, MTSET had no effect on the C14xA mutant, which has no remaining intracellular cysteines. More significantly, the rate of the inhibition of ternary C11xA was strongly gating state dependent (Fig. 2), which is in agreement with previously published results obtained in the absence of auxiliary subunits (Wang et al., 2005). When MTSET was applied to channels in the resting state (240 ms, at −120 mV), the time course of the inhibition was very slow ($k_{MTSET} = 0.038 \text{mM}^{-1}\text{s}^{-1}$). In sharp contrast, when a 7-ms pulse of MTSET was applied immediately following current activation by a strong step depolarization to +80 mV, the rate constant of inhibition was $\sim 475$-fold faster ($k_{MTSET} = 19 \text{mM}^{-1}\text{s}^{-1}$). When a 120-ms pulse of MTSET was applied at the end of a long step depolarization to +80 mV to test the inactivated state (Fig. 1), the rate constant of inhibition by MTSET was intermediate ($k_{MTSET} = 0.33 \text{mM}^{-1}\text{s}^{-1}$) between those for resting and activated channels. These observations confirm that the chemical modification of at least one thiolate group in the T1 interfacial Zn$^{2+}$ site causes inhibition of the Kv4.1 channel, implying that these intersubunit interfaces play a critical functional role (Wang et al., 2005). Importantly, the state-dependent cysteine accessibility rate constants are correlated with the main functional states of the Kv4.1 channel. Thus, a dynamic T1–T1 interface could adopt three distinct conformations in resting, activated, and inactivated channels because the targeted intracellular cysteines in the C11xA mutant are all located in the interfacial Zn$^{2+}$ site. These conformational correlatives are not induced by the eleven Cys→Ala mutations or the presence of auxiliary subunits because we observed similar state dependence and rate constants with wild-type and C11xA in the absence or presence of KChIP1 and DPPX-S (Table I).

To test whether the integrity of the T1 Zn$^{2+}$ site and the number of cysteines in this site are important in
the activated state and 400 μM for the resting and inactivated states. The inhibition by MTSET during the test pulse is not apparent because the chemical modification and the resulting inhibition were slow relative to channel gating.

determining the magnitude of $k_{\text{MTSET}}$ or its state dependence, we examined the ternary complexes of C12xA and C13xA. The coexpression of these mutants with KChIP1 and DPPX-S is necessary to rescue the lethal phenotype induced by the mutations of the Zn$^{2+}$ cysteines (Wang et al., 2005; Kunjilwar et al., 2004) (MATERIALS AND METHODS). C12xA has two intracellular cysteines (C131 and C132) in the Zn$^{2+}$ site, which cannot form an intersubunit metal bridge but an intrasubunit metal bridge with H104 (Nanao et al., 2003). By contrast, C13xA with only one intracellular cysteine in the Zn$^{2+}$ site (C110) could form a relatively weak metal bridge between C110 and H104 (Nanao et al., 2003). Inhibition of both mutant channels by MTSET was state dependent in a manner qualitatively similar to that observed with C11xA (Table I). From the three mutants, C11xA, C12xA, and C13xA, the average difference between the $k_{\text{MTSET}}$ of the activated and resting states was $\sim$200-fold, and between activated and inactivated states was $\sim$47-fold. Therefore, the integrity of the Zn$^{2+}$ site is not necessary to preserve the state dependence of $k_{\text{MTSET}}$. Moreover, $k_{\text{MTSET}}$ of the activated state increased proportionally with the number of cysteines in the T1 Zn$^{2+}$ site (Fig. 3 and Fig. 6, A and B). Our previous studies showed that Kv4.1 mutants with their Zn$^{2+}$ site either intact (C11xA) or disrupted (C12xA, C13xA, and C14xA) exhibit similar gating properties under the same condition (Wang et al., 2005). Thus, the relationship between $k_{\text{MTSET}}$ and the number of Zn$^{2+}$ site cysteines is unlikely to result from global channel distortions caused by the mutations. Instead, $k_{\text{MTSET}}$ is simply a function of the number of potential targets/subunit, with the most significant increase occurring when this number increases from 1/subunit to 2 or 3/subunit (Table I, Fig. 3, and Fig. 6, A and B).

Tight Correlation between the Chemical Modification of the T1–T1 Interface and Voltage-dependent Activation

If the state dependence of $k_{\text{MTSET}}$ originates from the voltage-dependent activation process of the Kv4.1 channel, we expect a close correlation between the peak conductance–voltage ($G_{\text{PEAK}}$–$V_m$) and the $k_{\text{MTSET}}$–$V_m$ relations. To test this hypothesis, we examined the inhibition of the ternary C11xA complex by MTSET at various membrane potentials between $-120$ and $+120$ mV. We conducted these experiments as described above for the activated state (Fig. 1, middle) except for voltages between $-80$ and $-50$ mV, which are not sufficiently depolarized to induce significant current activation. In this voltage range, an 80-ms MTSET pulse was first applied during a 100-ms step depolarization to $-80$ or $-50$ mV; the membrane potential was then hyperpolarized for 240 ms at $-100$ mV to remove any inactivation induced by the first pulse, and lastly a 4-ms test pulse to $+80$ mV was applied to monitor the available current. Fig. 4 shows that inhibition by MTSET became more rapid with membrane depolarization and was well described as an exponential decay. Thus, $k_{\text{MTSET}}$ is voltage dependent. The best-fit fourth-order Boltzmann function estimated a maximal $k_{\text{MTSET}}$ on the order of 21 mM$^{-1}$s$^{-1}$ (Fig. 5 B). Furthermore, there is a close correlation between the $G_{\text{PEAK}}$–$V_m$ and $k_{\text{MTSET}}$–$V_m$ relations (Fig. 5 B), which suggests that the conformational change in the T1–T1 interface faithfully mirrors activation gating of the Kv4 channel. In that case, any factor that shifts the voltage dependence of activation gating should shift the voltage dependence of $k_{\text{MTSET}}$ as well. We tested this prediction in two ways: (1) coexpression of C11xA with DPPX-S only to induce a leftward shift of the $G_{\text{PEAK}}$–$V_m$ relation (unpublished data); and (2) exposure of the ternary complex of
C11xA to 400 μM Ni\(^{2+}\) (Cd\(^{2+}\) or Zn\(^{2+}\)) in the external pipette solution of the inside-out patch to induce a rightward shift of the \(G_{\text{PEAK}}-V_m\) relation (Song et al., 1998). We found indeed that the \(k_{\text{MTSET}}-V_m\) relation precisely followed the shifts induced by these two manipulations (Fig. 5). Therefore, the strong correlation between the voltage dependencies of \(G_{\text{PEAK}}\) and \(k_{\text{MTSET}}\) in three independent conditions is compelling evidence for a tight functional coupling between transmembrane voltage sensor and the intracellular T1–T1 intersubunit interface in a Kv channel during voltage-dependent gating.

**Disulfide Bond Cross-linking Across the T1–T1 Interface Inhibits Kv4.1 Activation**

If a movement of the T1–T1 interface is required for normal gating, locking the T1–T1 interface by forming a disulfide bond between two adjacent subunits should prevent the putative intersubunit displacement and thus suppress gating. Although the structure of the Zn\(^{2+}\)-free T1 site is not known, the available crystal structure of the T1 domain (Fig. 6, A and B) reveals that the distances between the β carbons of C110 and C131, C110 and C132, and C131 and C132 are 5.44 Å, 5.43 Å, and 3.97 Å, respectively. To form a disulfide bond, the β carbons of the cysteinyl groups in a rigid protein must be within 3.4–4.6 Å (Careaga and Falke, 1992); but in flexible proteins, the β carbons of the cysteinyl groups may be separated by as much as 15 Å (Falke and Koshland, 1987). Therefore, under proper conditions, the formation of intersubunit disulfide bonds in the channel tetramer is likely because our functional data suggest a dynamic T1–T1 interface. To test this hypothesis, we exposed the intracellular side of the channel to a mild oxidizing agent. Fig. 6 C shows that Cu/P (50 μM
Tightly Coupled Gating Motions in the T1 Domain

CuSO₄ and 200 μM phenanthroline; MATERIALS AND METHODS) inhibited the Kv4.1 current substantially when the intersubunit pair C110/C132 was available. Similarly, but to a lesser degree, inhibition was observed when the intersubunit pair C110/C131 was available. The inhibition was not reversible by washout of Cu/P, but the reducing agent DTT reversed it slowly (Fig. 7); and neither Cu²⁺ nor phenanthroline alone affected the Cu/P-sensitive mutants (unpublished data). In sharp contrast, when the intrasubunit pair C131/C132 or just C110 remained, there was no inhibition by Cu/P (Fig. 6 C). These observations suggest that the inhibition by Cu/P is not due to overoxidation of the thiolate to sulfinic or sulfonic acid but to the formation of inter-subunit disulfide bonds between C110 and C132 or between C110 and C131. Apparently, the latter pair formed the disulfide bond less efficiently than the former, suggesting that the spatial relationship between C110 and C132 is more favorable than between C110 and C131. To support this conclusion further, we investigated the inhibition by MTSET. If the cysteine pair forms a disulfide bond, the inhibition by MTSET would be reduced in a manner that reflects the efficiency of disulfide bond formation upon Cu/P treatment. Fig. 6 D shows that MTSET inhibited the currents by ~76–79% when all T1 thiolate groups remained free. However, when the putative T1–T1 disulfide bonds were formed after pretreating with Cu/P, the mutant channels harboring the pairs C110/C132 and C110/C131 were inhibited by only ~25% and 43%, respectively. These results showed that upon oxidation, fewer cysteines remained free and that disulfide bond formation was more efficient between C110 and C132, as hypothesized above. Altogether, the cross-linking results are consistent with channel inhibition resulting from strait-jacketing a functionally critical and dynamic T1–T1 interface during gating.

DISCUSSION

Others have established strong correlations between cysteine accessibility changes in the S4 or S6 transmembrane segments and gating charge movements or pore opening, respectively (Yang and Horn, 1995; Larsson et al., 1996; Yang et al., 1996; Liu et al., 1997; Baker et al., 1998; Mannikko et al., 2002). These correlations are the bases of the proposed mechanisms of activation gating in voltage-gated cation channels. Our results strongly suggest that the coupled conformational changes extend beyond the S4 and S6 segments into the interfacial Zn²⁺ site of the intracellular T1 domain in Kv4 channels.

Working Models and Mechanisms

From our earlier work and this study, we conclude that the Kv4 T1–T1 interface is functionally active because chemical modification of the Zn²⁺ site thiolate groups or intersubunit disulfide bridges involving these groups cause channel inhibition (Fig. 6) (Wang et al., 2005). This inhibition may result from a steric local perturbation or strait-jacketing of the T1 domain, respectively. In addition, we demonstrated that the cysteine accessibility is much faster in the activated state than in the resting or inactivated states (Fig. 3; Table I), and that \( k_{\text{MTSET}} \) is dependent on the membrane potential (Figs. 4 and 5). Importantly, this voltage dependence follows the \( G_p-V \) relation faithfully (Fig. 5). Thus, \( k_{\text{MTSET}} \) is tightly correlated with the channel’s conductance change. Fig. 8 B illustrates hypothetical working models that attempts to explain these observations.

![Figure 5](image-url)
To create this cartoon models, we used the 3D crystal structure of Kv1.2 in the open state as a template (Fig. 8 A). Kv1.2 and Kv4.1 are expected to share similar structural features. Note that the voltage-sensing domains (VS) are connected to the pore domain (P) via the S4–S5 linkers, and to the T1 domain via the S1–T1 linkers; and that the T1 domain sits just below the S6 segments (and the S6 tails; not depicted). The latter is critical because the S6 helix bundle at the internal mouth of the channels is the main gate that controls the opening of the pore. Given these general features, including a relatively restricted space between the membrane-spanning core of the channel and the T1 domain, we propose that the closed T1–T1 interface and the S6 tail (or post-S6 segment) bury the critical Zn$^{2+}$ site cysteines in the resting state. Therefore, the cysteine accessibility to MTSET is low. When membrane depolarization activates the channel, two alternative hypotheses may explain the dramatic increase in cysteine accessibility at the T1 Zn$^{2+}$ sites (Fig. 8 B).

In one scenario, the voltage-dependent displacement of the S4 sensor in the VS domain moves the S4–S5 linker and allows the opening the S6 helix bundle (Lu et al., 2002; Tristani-Firouzi et al., 2002; Long et al., 2005b). The latter opens the pore and exposes the T1–T1 interface, which undergoes a quasi-simultaneous conformational change through a possible direct interaction between the post-S6 segment and the T1–T1 interface at the level of the L4 layer (Fig. 8 B, bottom pathway). Ultimately, these conformational changes expose the Zn$^{2+}$ site cysteines and increase the accessibility to MTSET. The interaction between post-S6 and
the T1–T1 interface may be critical for the opening of the pore because T1–T1 cross-linking appeared to be sufficient to inhibit the channel. Also, because the S6 helix bundle is the main activation gate that controls the opening of the pore, the contributions of post-S6 help to explain the tight correlation between the voltage dependencies of $k_{	ext{MTSET}}$ and peak conductance. In the alternative scenario, the voltage-dependent displacement of the S4 segment moves the S4–S5 segment to open the pore and induces a rearrangement of the VS domain (Chanda et al., 2005), which propagates into the T1 domain via the S1–T1 linker. This propagated conformational change could open the T1–T1 interface locally at the Zn$^{2+}$ site (L4 layer); however, to expose the cysteines and increase the accessibility to MTSET, the post-S6 segments would have to move too in a manner that is strictly coupled to the T1–T1 displacements. Therefore, if the T1–T1 interface cannot shift (e.g., upon intersubunit cross-linking), the S6 helix bundle cannot complete the pore opening. The VS domain-driven T1–T1 displacements alone cannot account for the observed voltage dependence of $k_{	ext{MTSET}}$ because this is tightly correlated with the Gp-V relation and the necessary movements of the voltage sensors (i.e., the Q-V relation) are expected to occur at more negative membrane potentials. Currently, our data cannot distinguish between these scenarios, and the mechanisms responsible for the inhibition are not known. Nevertheless, these working models provide concrete frameworks to investigate the striking voltage dependence of the putative conformational changes in the T1–T1 interface of Kv channels and their role in gating.

In contrast to the ~260-fold change in cysteine accessibility between resting and activated channels, the average change is much smaller between resting and inactivated channels (~3.7-fold) (Fig. 2; Table I). This observation may also be significant because it suggests that the closed and inactivated conformations of Kv4 channels are structurally alike, which is consistent with the presence of closed-state inactivation (Bahring et al., 2001; Beck and Covarrubias, 2001; Shahidullah and Covarrubias, 2003; Jerng et al., 2004). It is also in agreement with a model of closed-state inactivation induced by the decoupling between the S4 voltage sensor and the S6 gate (Shin et al., 2004). In this decoupled state, the resting and inactivated states of the channel may become functionally indistinguishable.

**Physiological Significance**

The proposed displacements could have important functional consequences. For instance, the T1–T1 interface may report the activation status of the channel. Consequently, the redox potential of the cell may modulate the functional activity of the T1–T1 interface in a state-dependent manner. A recent study has demonstrated acute state-dependent redox modulation of putative Kv4 channels in internally dialyzed cardiac myocytes and implicated intracellular sulfhydryl groups (Rozanski and Xu, 2002). Based on our results, we propose that the Kv4 Zn$^{2+}$ site cysteines are potential targets of physiological redox modulation in the heart. This modulation may exist in other excitable tissues and affect other Kv channels (Kv2 and Kv3), which also harbor the Zn$^{2+}$ site cysteines in the T1 domain (Bixby et al., 1999).

At a more mechanistic level, the emerging multitasking picture of the T1 domain reveals three separate but fundamentally important functions: (1) T1 determines specific subunit coassembly within Kv subfamilies (Xu et al., 1995; Li et al., 1992; Bixby et al., 1999); (2) T1 is the anchoring site for auxiliary subunits of Kv channels (Gulbis et al., 2000; Scannevin et al., 2004; Callsen et al., 2005; Long et al., 2005a); and (3) in conjunction with the membrane spanning core and possibly other intracellular regions of the Kv channel, T1 contributes to the molecular rearrangements that govern gating. Perhaps the objective of this contribution is to allow the expansion of the lateral intracellular portals of the channel, which could favorably influence the opening of the main S6 gate and the rapid access of K$^+$ and the intracellular inactivation gate to the internal mouth of the Kv pore. This hypothetical expansion is reasonable because the lateral intracellular portals are clearly apparent.
displacements as the S6 helix bundle expands to open the pore. In both pathways, the speculative conformational changes expose the T1–T1 interfaces and the post-S6 segments and the L4 layers of T1 may drive the T1–T1 mediated by the S1–T1 linker. Instead, a direct interaction between the interfaces does not depend on a propagated conformational change of the T1 domain. The blue cylinders represent the post-S6 segments (right) was used as a template to develop the schematic model of the Kv1.2 channel (Long et al., 2005a). The different colors depict distinct functional domains of the Kv channel: pore domain (P, blue), voltage sensing domain (VS, green; including the S4–S5 linker), and the T1 domain (T1, orange). The S1–T1 linkers (gray) connect the VS domain to the T1 domain. Dashed lines represent the approximate boundaries of the lipid bilayer. A section of the channel at the level of the internal boundary of the lipid bilayer is shown on the right-hand side of the model. This view illustrates the fourfold symmetry of the channel oligomer with the S6-tails (blue) sitting just above the T1 domains and the S1–T1 linkers extending from each corner of the oligomer. (B) Two hypothetical pathways that explain the voltage dependence of the cysteine accessibility changes in the T1–T1 interface. The section shown in A (right) was used as a template to develop the schematic model of the T1 domain. The blue cylinders represent the post-S6 segments sitting on the T1–T1 interface when the pore is closed. At rest, the T1–T1 interfaces are buried. The upper pathway assumes VS domain-driven displacements of the T1–T1 interfaces via the S1–T1 linkers, which allow the concerted movement of the S6 helix bundle and the resulting opening of the pore. The lower pathway assumes the quasi-simultaneous displacements of the T1–T1 interfaces and the opening of the main S6 gate. In this case, exposing the T1–T1 interface does not depend on a propagated conformational change mediated by the S1–T1 linker. Instead, a direct interaction between the post-S6 segments and the L4 layers of T1 may drive the T1–T1 displacements as the S6 helix bundle expands to open the pore. In both pathways, the speculative conformational changes expose the Zn$^{2+}$ site cysteines in a manner that reflects the increase in conductance resulting from the opening of the main activation gate.

Figure 8. Coupling between voltage-dependent activation and the putative movements of the T1–T1 interfaces and the post-S6 segments. (A) Structural model of a Kv channel based on the 3D crystal structure of the Kv1.2 channel (Long et al., 2005a). The different colors depict distinct functional domains of the Kv channel: pore domain (P, blue), voltage sensing domain (VS, green; including the S4–S5 linker), and the T1 domain (T1, orange). The S1–T1 linkers (gray) connect the VS domain to the T1 domain. Dashed lines represent the approximate boundaries of the lipid bilayer. A section of the channel at the level of the internal boundary of the lipid bilayer is shown on the right-hand side of the model. This view illustrates the fourfold symmetry of the channel oligomer with the S6-tails (blue) sitting just above the T1 domains and the S1–T1 linkers extending from each corner of the oligomer. (B) Two hypothetical pathways that explain the voltage dependence of the cysteine accessibility changes in the T1–T1 interface. The section shown in A (right) was used as a template to develop the schematic model of the T1 domain. The blue cylinders represent the post-S6 segments sitting on the T1–T1 interface when the pore is closed. At rest, the T1–T1 interfaces are buried. The upper pathway assumes VS domain-driven displacements of the T1–T1 interfaces via the S1–T1 linkers, which allow the concerted movement of the S6 helix bundle and the resulting opening of the pore. The lower pathway assumes the quasi-simultaneous displacements of the T1–T1 interfaces and the opening of the main S6 gate. In this case, exposing the T1–T1 interface does not depend on a propagated conformational change mediated by the S1–T1 linker. Instead, a direct interaction between the post-S6 segments and the L4 layers of T1 may drive the T1–T1 displacements as the S6 helix bundle expands to open the pore. In both pathways, the speculative conformational changes expose the Zn$^{2+}$ site cysteines in a manner that reflects the increase in conductance resulting from the opening of the main activation gate.

between the membrane-spanning core and the T1 domain in the recently published crystal structure of a mammalian Kv channel in the open state (Kobertz et al., 2000; Kim et al., 2004; Long et al., 2005a).

Conclusion

We have provided compelling evidence for tight coupling between voltage-dependent activation of a Kv4 channel and conformational changes involving the intracellular T1–T1 interface. We propose that the complex structural rearrangements that control fast and efficient activation gating of eukaryotic Kv channels include propagated movements in the conserved L4 layer of the T1 domain and a post-S6 COOH-terminal segment that may contact the T1–T1 interface. These findings suggest novel ways to regulate Kv channel gating in excitable tissues.

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