Basis for allosteric open-state stabilization of voltage-gated potassium channels by intracellular cations

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The open state of voltage-gated potassium (Kv) channels is associated with an increased stability relative to the pre-open closed states and is reflected by a slowing of OFF gating currents after channel opening. The basis for this stabilization is usually assigned to intrinsic structural features of the open pore. We have studied the gating currents of Kv1.2 channels and found that the stabilization of the open state is instead conferred largely by the presence of cations occupying the inner cavity of the channel. Large impermeant intracellular cations such as N-methyl-d-glucamine (NMG+) and tetraethylammonium cause severe slowing of channel closure and gating currents, whereas the smaller cation, Cs+, displays a more moderate effect on voltage sensor return. A nonconducting mutant also displays significant open state stabilization in the presence of intracellular K+, suggesting that K+ ions in the intracellular cavity also slow pore closure. A mutation in the S6 segment used previously to enlarge the inner cavity (Kv1.2-I402C) relieves the slowing of OFF gating currents in the presence of the large NMG+ ion, suggesting that the interaction site for stabilizing ions resides within the inner cavity and creates an energetic barrier to pore closure. The physiological significance of ionic occupation of the inner cavity is underscored by the threefold slowing of ionic current deactivation in the wild-type channel compared with Kv1.2-I402C. The data suggest that internal ions, including physiological concentrations of K+, allosterically regulate the deactivation kinetics of the Kv1.2 channel by impairing pore closure and limiting the return of voltage sensors. This may represent a primary mechanism by which Kv channel deactivation kinetics is linked to ion permeation and reveals a novel role for channel inner cavity residues to indirectly regulate voltage sensor dynamics.

INTRODUCTION

Voltage-gated potassium (Kv) channels are responsible for repolarizing the membrane and regulating firing rates in electrically excitable cells (Hille, 2001). Activation of Kv channels relies on the coupling of a voltage-sensing domain (VSD; transmembrane segments S1–S4) to a highly selective potassium (K⁺)-conducting pore domain (S5–S6; Long et al., 2005b). The pore creates an energetically permissible route for charged K⁺ ions across the hydrophobic membrane bilayer. Selectivity filter binding sites stabilize dehydrated K⁺ ions to confer K⁺ selectivity, and an aqueous inner cavity funnels hydrated K⁺ ions to the selectivity filter (Y. Zhou et al., 2001; Long et al., 2005b). The conduction pathway becomes accessible to intracellular ions from the cytosol only when the intracellular gate opens and allows the passage of ions into the cavity (Liu et al., 1997).

Voltage-dependent activity is generated through VSD movements that couple to the pore and displace charge before the channel opens. These movements comprise several kinetically distinct and nonconductive closed states that are occupied before the channel pore opens at the intracellular gate (Zagotta et al., 1994b; Schopka and Sigworth, 1998; Ledwell and Aldrich, 1999; del Camino et al., 2005). The coupling of the VSD to the pore gate is mediated largely by a strong interaction between the S4–S5 linker and the cytosolic end of S6, where the activation gate resides (Lu et al., 2002; Long et al., 2005a,b). Gating charge, which reflects the movements of the VSD, has been shown to be slower to return from voltages at which the channel pore opens than from potentials where the open state has not been populated (Perozo et al., 1993; Zagotta et al., 1994b; Chen et al., 1997; Olcense et al., 1997; Batulan et al., 2010; Lacroix et al., 2011). It appears that the channel pore must close before any of the gating charges can return, as the open pore prevents the return of the tightly coupled voltage sensor to its resting conformation, a process referred to as voltage sensor immobilization (Armstrong, 1971; Bezanilla et al., 1991). The kinetics of the return of the gating charge are therefore controlled by two factors: the force exerted by negative voltage, pulling the positively charged residues in the voltage sensor down, and...
the opposing force conferred by the stability of the open channel pore, which has to close before the voltage sensor can return.

The factors that affect the stability of the open pore fall broadly into two categories: extrinsic (or allosteric), in which elements disparate from the pore impact pore stability, and intrinsic, in which the stability is determined by the local peptide conformation. Extrinsic mechanisms include the inhibition of activation gate closure by the N-terminal peptide in wild-type (WT) Shaker channels, which immobilizes gating charge while it prevents pore closure (Bezanilla et al., 1991). Intracellular applied quaternary ammonium ions greatly slow the recovery of ionic and gating currents and stabilize the open pore in a similar way (Armstrong, 1971; Choi et al., 1993; Melischuk and Armstrong, 2001), and both of these examples highlight the rate-limiting nature of the first transition away from the open state. Alternatively, if an open state has an intrinsic conformation that is more stable than the closed state, the rate of leaving that state will be slowed. Processes such as slow inactivation have been linked to the slow return of gating charge, suggesting that conformational rearrangements, associated with C-type inactivation processes, might physically stabilize the voltage sensor in the activated state (Bezanilla et al., 1982; Fedida et al., 1996; Olcese et al., 1997; Wang et al., 1999). More recently, stabilization of the VSD after depolarization has been observed in nonactivating HCN (hyperpolarization activated cyclic nucleotide gated) channels (Bruening-Wright and Larsson, 2007) and a Ciona intestinalis voltage-sensitive phosphatase, which lacks a pore (Villalba-Galea et al., 2009). These studies suggest that the voltage sensor might “relax” by itself and adopt an intrinsically distinct, stable conformational state (Villalba-Galea et al., 2008). Even in the absence of inactivation, or quaternary ammonium ions, or prolonged depolarizations that induce VSD relaxation, a slowing of the return of charge is still observed after depolarization. Gating currents recorded from nonconducting Shaker IR W434F channels in K⁺ internal solutions still display a slowing of I\text{Goff}\_\text{IR} after brief depolarizations to potentials at which the channels open (Perozo et al., 1993), and varying the major monovalent cation in intracellular and extracellular solutions has been shown to influence the rate of return of gating charge (Chen et al., 1997; Wang et al., 1999; Melischuk and Armstrong, 2001; Varga et al., 2002; Goodchild and Fedida, 2012). It is clear that both extrinsic and intrinsic factors can impact directly on the stability of the open state and the subsequent rate of channel deactivation, but their relative contributions, although currently unknown, are often dependent on specific conditions and the channel type.

In this study, we sought to understand the mechanism of open-state stabilization in Kv1.2 channels by varying the intracellular ionic composition and mutating pore residues. Although Kv1.2 gating currents have not previously been studied, we used Kv1.2 rather than Shaker channels for two reasons: (1) Kv1.2 is resistant to inactivation as a result of a valine residue at position V381 (T449 in Shaker), which allowed us to investigate gating independently from inactivation processes; and (2) the availability of a high resolution crystal structure of the open-state channel allowed us to interpret our results from a structural perspective. Our results show that the stabilization of voltage sensors associated with the open state is critically regulated by the occupancy of a specific site in the inner pore. This suggests that the open pore is not intrinsically more stable than the closed pore, but that it is controlled extrinsically by the occupancy of the inner pore by both permeant and nonpermeant ions, which create an energy barrier to pore closure.

**MATERIALS AND METHODS**

**Cell culture and molecular biology**

WT Kv1.2 channel cDNA were mutated within the mammalian expression vector pGW1. Point mutations were generated using the QuikChange II Site-Directed Mutagenesis kit (Agilent Technologies) and confirmed by sequencing. Channels were expressed in mammalian tsA201 cells grown and maintained in MEM (Invitrogen) at 37°C in an air/5% CO₂ incubator. Media contained 10% bovine serum (Gibco) with 100 U/ml penicillin and 100 µg/ml streptomycin (Invitrogen). On the day before transfection, cells were washed with MEM, treated with trypsin/EGTA for 1 min, and plated on 25-mm² glass coverslips. WT Kv1.2-pGW1 and green fluorescent protein–pcDNA3 cDNA were then transiently cotransfected using Lipofectamine 2000 transfection reagent according to the manufacturer’s instructions (Invitrogen). 24–48 h after transfection, coverslips with adherent cells plated on the surface were washed in external solution and then placed in a recording chamber containing the external bath solution. All recordings were performed at room temperature (~20°C).

**Patch-clamp electrophysiology**

Whole-cell current recording and data acquisition were performed using an amplifier (Axopatch 200B) controlled with pClamp10 software (Molecular Devices). Patch electrodes were pulled from thin-walled borosilicate glass (World Precision Instruments) on a horizontal micropipette puller (Sutter Instrument) and fire polished before use. Electrodes had resistances of 1–2 MΩ when filled with internal solutions. Liquid junction potentials were calculated using the junction potential calculator function in pClamp10. Membrane potentials were corrected for junction potentials >3 mV that arose between the pipette and bath solution by applying the appropriate offset potential before recording, unless otherwise stated. Leak subtraction was routinely applied using a −P/8 or −P/6 protocol from the holding potential unless otherwise stated. Currents were filtered at 10 kHz and sampled at 100 kHz for gating currents and deactivation tail current recordings or filtered at 2–5 kHz and sampled at 25 kHz for other ionic current recordings. Only isolated, unconnected cells were used for recording, and ≥80% series resistance compensation was applied in all whole-cell recordings to minimize any error in membrane voltage and low-pass filtering effects resulting from series resistance and cell capacitance.
Solutions

Gating current recordings. The extracellular bath solution contained the following (mM): 140 TEACl, 10 HEPES, and 1 CaCl₂, pH adjusted to 7.4 with HCl. Intracellular pipette solution contained the following (mM): 140 Na⁺Cl, 10 HEPES, and 5 EGTA, pH adjusted to 7.4 with NaOH. Intracellular pipette solution was made from WT Kv1.2 channels was that the OFF gating currents were sensitive to the presence of intracellular Na⁺.

Whole-cell ionic current recordings. The extracellular solution contained the following (mM): 135 NaCl, 5 KCl, 1 MgCl₂, 2.8 Na-acetate, 10 HEPES, and 1 CaCl₂, pH adjusted to 7.4 with NaOH. Intracellular pipette solution contained the following (mM): 130 KCl, 5 EGTA, 1 MgCl₂, 10 HEPES, 4 Na₇-ATP, and 0.1 GTP, pH adjusted to 7.2 with KOH.

Data analysis

Charge-voltage (Q-V) measurements were obtained by integrating the ON gating (IgON) or OFF gating currents (IgOFF) over an 11-ms period, unless otherwise stated. Conductance-voltage (G-V) relationships were constructed from isochronal tail current measurements at −40 mV after families of 50-ms depolarizing prepulses were applied to activate the channels, from a holding potential of −100 mV. Q-V and G-V relationships were normalized to the maximum value and fit with a single Boltzmann function of the form $A/A_{\text{max}} = (1 + \exp(zF(V - V_{\text{0.5}})/RT))^{-1}$, where $A/A_{\text{max}}$ is the normalized conductance (G/Qmax) or charge (Q/Qmax), $z$ is the equivalent charge, $V_{\text{0.5}}$ is the half-activation voltage, $F$ is Faraday’s constant, $R$ is the gas constant, and $T$ is the temperature in kelvin. Fitting was performed on data from individual experiments using Microcal Origin software (version 8), and the fit parameters quoted in the text are the means ± SEM of the individual fits. In figures, curves are fit to mean data for display purposes. Statistical significance was assessed using unpaired Student’s t tests and considered significant for $P < 0.05$.

Kinetic modeling

Our experiments suggested that the kinetics of Na⁺ (Na⁺) dissociation (kon) was voltage dependent (see Fig. 3 C) and was therefore modeled using rate constants of the form

$$k_{\text{on}} = k_{\text{on}(0 \text{mV})} \times \left[ B^+ \right] \times \exp \left( \frac{zFV}{RT} \right)$$

and

$$k_{\text{off}} = k_{\text{off}(0 \text{mV})} \times \exp \left( \frac{zFV}{RT} \right),$$

where $k_{\text{on}(0 \text{mV})}$ and $k_{\text{off}(0 \text{mV})}$ are the voltage-independent rate constants reflecting the intrinsic chemical binding energy of the interaction, $[B^+]$ is the concentration of Na⁺, $z$ is the Na⁺ charge, and $\delta$ is the fraction of the membrane field sensed by the ions. As the majority of charge is carried during transitions between closed states (Ledwell and Aldrich, 1999), occupancy of the O:B state will trap channels in the open state and voltage sensors in the activated state. The dissociation and association rate of Na⁺ ions entering and leaving the cavity will then determine the rate-limiting step in voltage sensor return and thus be reflected in the kinetics of IgOFF.

We could not measure IgOFF at 0 mV, as voltage sensors are in the deactivated state (see Fig. 3). The rate of relaxation to a new equilibrium between two states is defined by $1/(k_{\text{on}} + k_{\text{off}})$ and at 0 mV, therefore, simplifies to $1/(k_{\text{on}(0 \text{mV})} \times [\text{Na}^+] + k_{\text{off}(0 \text{mV})})$. Substituting our estimate of the rate of onset of the development of slow charge return (1/τ) at 0 mV of 270.27 s⁻¹ (see Fig. 2 C), an apparent $k_{\text{on}(0 \text{mV})}$ of 1,929.71 M⁻¹ s⁻¹ for Na⁺ association was calculated. This is likely to represent a lower limit, as the derivation of the rate of onset of the slow OFF gating would be dependent on channel opening and therefore be rate limited by the opening transition, which may be slower than Na⁺ association. These experimentally predicted $k_{\text{on}(0 \text{mV})}$ and $k_{\text{off}(0 \text{mV})}$ values were then used in the model. The voltage dependence of the dissociation rate of Na⁺ from the pore in the model was set at $\alpha = 1.6$, thus assuming an equivalent voltage dependence of the association and dissociation rates.

Simulations were performed using the program IonChannelLab (De Santiago-Castillo et al., 2010) and Berkeley Madonna with the integration period set at 10 µs (100 kHz). Simulated gating currents were then used in the model. The voltage dependence of the dissociation rate of Na⁺ from the pore in the model was set at $\alpha = 1.6$, thus assuming an equivalent voltage dependence of the association and dissociation rates.

Online supplemental material

Fig. S1 demonstrates the resistance of Kv1.2 to inactivation over the time course of the gating current experiments and the insensitivity of this inactivation to the presence of intracellular Na⁺. Supplemental materials and methods for this experiment and the docking simulation are also available. Online supplemental material is available at http://www.jgp.org/cgi/content/full/jgp.201210823/DC1.

RESULTS

Striking slow OFF gating current of the Kv1.2 channel voltage sensor

As stated in the Introduction, it is the stability of channel open states relative to the closed states that determines the rate of return of gating charge when channels close (Armstrong, 1971; Bezanilla et al., 1991; Melishchuk and Armstrong, 2001). The first observation that was made from WT Kv1.2 channels was that the OFF gating currents at −100 mV displayed fast kinetics when returning from pulses up to −50 mV (Fig. 1, A and B). However, at more positive pulse potentials, a slow component in the IgOFF kinetics emerged along with a decrease in the peak amplitude, illustrated by individual traces in

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whereas the OFF gating currents become slowed and blunted after larger depolarizations. The $Q_{\text{off}}$-$V$ relationship was normalized to the maximum $Q_{\text{on}}$ and illustrates that only around 70% of the $Q_{\text{on}}$ had recovered within 11 ms as a result of the slowing of $I_{\text{gOFF}}$ (Fig. 1 C). The normalized peak $I_{\text{gOFF}}$ was plotted against the voltage of the preceding depolarizing pulse. Error bars represent SEM.

Fig. 1 B. These gating currents are measured with the permeant K$^+$ and Na$^+$ ions in standard physiological solutions replaced by NMG$^+$ and TEA$^+$. What is most obvious is the lack of symmetry between the ON gating currents that represent the movement of the charged voltage sensor from resting to activated and the OFF gating currents. The ON gating currents rise rapidly and show typical voltage-dependent kinetics as increasing depolarization increases the rate of charge movement,
Figure 2. Rate of induction of the slowing of Kv1.2 charge return and the voltage dependence of return of stabilized voltage sensors. (A) Representative gating current demonstrating the slowing in the kinetics of $I_{g_{off}}$ currents at a repolarizing voltage of $-100$ mV as the duration of depolarization to 0 mV was extended in 0.5-ms increments ($n = 4$). (B) Isolated traces from A after depolarization of 0.5, 1.5, 3.5, 9.5, and 14.5 ms. (C) Plot of the $Q_{off}/Q_{on}$ as a function of pulse duration. $Q_{off}$ was calculated by integrating the first 11 ms of $I_{g_{off}}$. (D) Representative gating currents obtained using the two-pulse protocol shown. Return of charge was observed after a 50-ms depolarizing pulse to 10 mV by a series of hyperpolarizing pulses in 10-mV increments from $-40$ to $-190$ mV. Linear membrane leak was subtracted online using a $-P/8$ leak subtraction protocol from a subtraction holding potential of $-140$ mV ($n = 6$). (E) The first 11 ms of $I_{g_{off}}$ were integrated, and the values were plotted against voltage and compared with the $Q_{on}$-V relationship from Fig. 1 D. Error bars represent SEM.
the return of stabilized voltage sensors. A simple comparison of the G-V with the Q off-V and peak Ig OFF plot illustrates that slowing of the OFF gating current correlates with potentials at which the open probability (Po) of the channels rapidly increases, thus linking the process of the slowing of charge return to the open pore.

Entry of the voltage sensor into the stabilized state occurs over the time course of channel openings

In these first recordings of OFF gating currents from Kv1.2, not only is a slowing of charge return extensive, but the onset is extremely rapid, and the voltage dependence of charge return is displaced to very negative potentials. The time course of the onset of the slow OFF gating (Fig. 2 A) was comparable with that of channel opening rather than an inactivation process that is limited in scope and occurs over a much longer time scale in Kv1.2 (Fig. S1 A) and that is not perturbed by the presence of intracellular NMG+ ions replacing K+. The emergence of the slow Ig OFF is clearly visible in the isolated traces shown in Fig. 2 B and represents a stabilization of the voltage sensor in the activated state. Fig. 2 C shows a plot of Q off/Q on that was fit with a single exponential function, giving a time constant for the stabilization of the voltage sensor at 0 mV of 3.7 ± 1.1 ms (n = 4).

The rate of the return of stabilized charge displayed a marked hyperpolarizing shift in the voltage dependence of the charge return (V0.5 = −114.2 ± 4.5 mV, z = 1.8 ± 0.1; n = 6; Fig. 2 E) compared with the Q on-V (Fig. 1 C) and suggested that to return charge to the resting state from this state over a fixed period of time required significantly more energy than was required to move the charge into the activated state in the first place. To accurately resolve the kinetics of charge recovery, we used a double pulse recovery protocol as shown in Fig. 3 A. The plots of charge recovery over time were fit with a single exponential function at holding potentials between −80 and −100 mV, where the channels are fully closed (Fig. 3 B). Now,

$$1 / \tau = I_{\text{Goff}(0 \text{ mV})} \times \exp \left(-\frac{z \delta}{RT} \right),$$

where $I_{\text{Goff}(0 \text{ mV})}$ represents the theoretical rate of gating charge return at 0 mV and $\delta$ is the voltage dependence.
The effects of varying intracellular cations on rate of charge return. (A) Representative current traces recorded using an intracellular TEA\(^+\) solution were evoked by 12-ms depolarizing pulses applied in 10-mV increments from a holding potential of \(-100\) mV. (B) Gating currents recorded with Cs\(^+\). (C) Representative currents obtained by depolarizing the membrane to 0 mV for 50 ms before repolarization to a series of pulses from \(-40\) to \(-190\) mV with TEA\(^+\). Linear membrane leak was subtracted online using a \(\Delta P/8\) leak subtraction protocol from a subtraction holding potential of \(-140\) mV. (D) Return of charge in the presence of intracellular TEA\(^+\) and Cs\(^+\) was measured during an 11-ms charge integration period after the depolarizing pulse and plotted against voltage. The \(Q_{\text{off}}\)-V for Cs\(^+\) had \(V_{0.5} = -38 \pm 1.8\) mV, \(z = 2.8 \pm 0.2\) (\(n = 3\)). Using intracellular TEA\(^+\) did not significantly affect \(Q_{\text{on}}\) (\(Q_{\text{on}}\)-V: \(V_{0.5} = -30.3 \pm 1.4\) mV, \(z = 2.3 \pm 0.3\); \(n = 5\)). The \(Q_{\text{off}}\)-V for TEA\(^+\) gave fitted values of \(V_{0.5} = -125.9 \pm 3.9\) mV and \(z = 2 \pm 0.2\) (\(n = 3\)). Error bars represent SEM.

Figure 4.

Intracellular ions, including physiological concentrations of K\(^+\), alter voltage sensor dynamics

We next sought to investigate the basis for the stability of the open state in Kv1.2. Slow OFF gating in Shaker channels is caused by the N-terminal region of the channel entering the pore in the same way as intracellular TEA\(^+\), through blockade of pore closure (Bezanilla et al., 1991; Perozo et al., 1992). More recently, a slowing of OFF gating has been correlated with the ionic radii of group 1 cations used in the internal solution, with larger cations causing more severe slowing (Wang et al., 1999). Here, we have examined the effect of two cations with previously described disparate effects in Kv channels: a large blocking cation, TEA\(^+\), which causes severe slowing, and a small cation, Cs\(^+\), that previously caused an increase in the rate of charge return (Fig. 4).

TEA\(^+\) (Fig. 4, A and C) caused a severe slowing of \(Q_{\text{off}}\) at potentials that correlated with channel opening and a comparable \(Q_{\text{on}}\) shift to NMG\(^+\) (Figs. 1–3). Using Cs\(^+\) solution (Fig. 4 B), we observed a small outward ionic current, as a result of slight Cs\(^+\) permeability through Kv1.2 channels, that masked \(Q_{\text{on}}\). We were, however, able to resolve \(Q_{\text{off}}\) at \(-100\) mV in these cells by adding 3.1 mM Cs\(^+\) to the extracellular solution, setting the reversal potential for Cs\(^+\) to \(-100\) mV to eliminate any contamination from ionic tail currents at that voltage (Chen et al., 1997). The Cs\(^+\) OFF gating currents were
slowed after depolarizations more positive than −30 mV but recovered fully within the integration period (Fig. 4 B). In this situation, little shift of \( Q_{\text{off}} \) was seen with Cs\(^+\) (Fig. 4 D), suggesting that because TEA\(^+\) and NMG\(^+\) internal solutions required a greater energy to recover gating charge than Cs\(^+\), the voltage sensor was stabilized in the activated state by the larger cations more effectively than by Cs\(^+\).

To answer the question of whether physiological concentrations of intracellular K\(^+\) ions could slow voltage sensor return in Kv1.2, we created a nonconducting mutant Kv1.2 W366F-V381T, analogous to the P-type inactivated Shaker W434F nonconducting channel. Fig. 5 A shows representative gating currents from this mutant recorded with K\(^+\) (Fig. 4 D), suggesting that because TEA\(^+\) and NMG\(^+\) internal solutions required a greater energy to recover gating charge than Cs\(^+\), the voltage sensor was stabilized in the activated state by the larger cations more effectively than by Cs\(^+\).

An inner cavity pore lining S6 residue I402 contributes to the interaction site for ionic stabilization of the voltage sensor

Based on the previous experiments, we considered the hypothesis that all intracellular ions that can occupy the Kv1.2 inner cavity will stabilize the open state of the channel and prevent voltage sensor return until they are expelled and, thus, that cations are a significant extrinsic regulator of open-state stability, channel pore closing, and charge return. In Shaker IR channels, a mutation of an inner pore cavity lining residue, I470, to the smaller C residue has previously been shown to allow TEA\(^+\) to reside in the closed inner cavity as a result of pore enlargement (Holmgren et al., 1997; Melishchuk and Armstrong, 2001).

In Kv1.2, a homologous mutation was made (Kv1.2-I402C). Gating currents recorded from Kv1.2-I402C with intracellular NMG\(^+\) no longer displayed slow OFF gating currents (Fig. 6 A). The OFF gating currents remained fast at all potentials, and the normalized \( Q_{\text{off}}-V \) showed saturating charge return. The \( Q_{\text{on}}-V \) \( V_{0.5} \) was not significantly different from Kv1.2 WT channels (Fig. 6 B), and the Kv1.2 transmembrane currents, the G-V relation, and the G\( V_{0.5} \) were not different from WT Kv1.2, suggesting that the activation and coupling of the voltage sensors to pore opening was unaltered by this mutation in S6. The absence of a discernable rising phase in Ig\( \text{OFF} \) after pulses that would open the pore suggested that the first open to closed transition was much faster in this mutant than in WT Kv1.2.

To compare Kv1.2-I402C Ig\( \text{OFF} \) kinetics with the WT channel, Ig\( \text{OFF} \) tracings at potentials between −80 and −100 mV were fit with a single exponential function to estimate the rate of charge return. Representative Ig\( \text{OFF} \) currents at negative potentials after a 50-ms depolarizing step to 0 mV to open channels are shown in the inset to Fig. 6 D. The mean Ig\( \text{OFF} \) rates were plotted against voltage and then fit with Eq. 1 to give the fitted values \( k_{\text{off}(0\, \text{mV})} = 103.1 \, \text{s}^{-1} \) and \( \delta = 0.4 \). The reduced \( \delta \) of Ig\( \text{OFF} \) rates observed in the I402C mutant result from the removal of a voltage-dependent rate-limiting transition, such as the exit of a charged intracellular ion. This observation is consistent with the mutant pore being able to close unimpeded around a blocking ion, or a significant reduction in the affinity of those ions for the mutant inner pore, increasing their dissociation rate such that they no longer limit the pore closing rate.

**Ionic deactivation kinetics is rate limited by K\(^+\) ion occupancy of the inner pore**

We have established that voltage sensor return is slowed in the presence of internal K\(^+\) ions (Fig. 5) and hypothesize that this is caused by the inability of the inner pore gate to shut when the cavity is occupied by an ion. To examine whether K\(^+\) ions in permeant ionic conditions could have a similar effect, we examined the deactivation kinetics of ionic currents in WT channels compared with Kv1.2-I402C channels. Deactivation tail currents recorded in 140 mM external K\(^+\) at −120 or −80 mV clearly show a faster rate of deactivation in Kv1.2-I402C (Fig. 6 E). The time for half the peak current to decay (\( t_{0.5, \text{decay}} \)) demonstrated that deactivation kinetics are threefold more rapid in Kv1.2-I402C at all voltages tested (\( n = 7–10 \)). These data confirm that there is a rate-limiting step in Kv1.2 channel deactivation that is reliant on a specific ionic coordination site within the inner cavity that must be vacated before the pore can shut.

**Kinetic model of intracellular cation modulation of pore closure**

To test the possibility that NMG\(^+\) acts as a limiting regulator of pore closure, we used a Markov state model to...
Figure 5. The nonconducting mutant Kv1.2 W366F,V381T shows slow OFF gating currents with intracellular K⁺ and Cs⁺. (A) Representative current traces recorded using K⁺ solution were evoked by 12-ms depolarizing pulses applied in 10-mV increments from a holding potential of −100 mV. (B) Individual representative gating current traces isolated from the same recording to illustrate the onset of slowing of the IgOFF current recorded at −100 mV after depolarizing steps from −50 to 10 mV in 20-mV increments. (C) Q-V relationships obtained from integrating IgON (Qon-V; Vₚ₀ = −39.4 ± 1.7 mV, z = 4.4 ± 1.3; n = 3) and IgOFF (Qoff-V) for 11 ms and fit with a Boltzmann function (n = 3). (D) Plot of the normalized peak IgOFF amplitude against voltage with a WT G-V curve in gray for comparison. (E) Representative current traces recorded using Cs⁺ solution. (F) Individual representative gating current traces isolated from the same recording from −50 and −10 mV. (G) Plot of the normalized peak IgOFF amplitude with internal Cs⁺ solution. Error bars represent SEM.
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Figure 6. Mutation of an S6 residue to enlarge the cavity relieved slowing of charge return and increased the rate of ionic deactivation kinetics. (A) Representative gating currents recorded from Kv1.2-I402C in the same conditions as Kv1.2 WT (Fig. 1 A) with NMG\textsuperscript{+}. (B) \( Q_{\text{on}} \)-\( V \) and \( Q_{\text{off}} \)-\( V \) relationships with values for the \( Q_{\text{on}} \)-\( V \) of \( V_{0.5} = -24.4 \pm 2.3 \text{ mV}, z = 2.9 \pm 0.8 \) (\( n = 7 \)) and for the \( Q_{\text{on}} \)-\( V \) of \( V_{0.5} = -25.6 \pm 2.3 \text{ mV}, z = 1.7 \pm 0.1 \) (\( n = 6 \)). (C) Representative ionic currents recorded from Kv1.2-I402C. The G-\( V \) relationship for Kv1.2-I402C was plotted with the WT (gray dashes) for comparison. Kv1.2-I402C, \( V_{0.5} = -1.1 \pm 2.2 \text{ mV}, z = 2.4 \pm 0.1 \) (\( n = 5 \)); \( P > 0.05 \) compared with WT. (D) To quantify the \( I_{\text{off}} \) rate, channels were depolarized to 0 mV for 50 ms to open channels, and the decay of \( I_{\text{off}} \) upon repolarization was fit with single exponential functions at potentials between -100 and -80 mV. Off rates (1/\( \tau \)) were as follows: -80 mV, 1/\( \tau \) = 369.8 ± 35.7 s\textsuperscript{-1};
simulated \( \text{IgOFF} \). The model was based on an established Shaker IR gating model in which the channel opens only once all four subunits have moved through two closed transitions into an active state (ZHA; Zagotta et al., 1994a). This model includes a modifying factor \( \theta \), which was set at 9.4, to slow the exit from the open state and reflect the cooperativity of the final concerted opening step and/or the combination of extrinsic and intrinsic open-state stability. The fast closed state present in the ZHA model to account for flickering single-channel openings was omitted in our model. The model, depicted in Fig. 7, was modified to incorporate an NMG\(^+\)-bound open channel state, O:B\(^+\). In this scheme, if the rate of NMG\(^+\) unbinding is significantly slower than the first open to closed transition (\( \delta \theta \)), it will become the dominant rate-limiting transition for pore closure and the subsequent return of gating charge. This is similar to the “foot in the door” mechanism known to operate in Shaker \( K^+ \) channels (Armstrong, 1971; Choi et al., 1993; Perozo et al., 1993).

Simulated OFF gating currents evoked by sequential depolarizations increasing by 20 mV from a holding potential of \(-100 \text{ mV}\) generated from the original ZHA model with \( \theta = 9.4 \) are shown in Fig. 7 B. The \( Q_{\text{on}} \) and \( Q_{\text{off}} \) relationships are symmetrical, with a \( V_{0.5} \) of \(-47 \text{ mV}\) indicating that all charge that had moved during the activation of the voltage sensors was returned within the integration period. However, when the \( Q_{\text{on}} \) was measured after depolarization to 0 mV for 50 ms to open channels, the \( V_{0.5} \) was somewhat hyperpolarized to \(-60 \text{ mV}\). This left shift reflected the slowing of \( \text{IgOFF} \) as a result of stabilization of the voltage sensor in the activated state by the slowed transition away from the open state. When compared with our data displayed in Fig. 2 E, it is apparent that Kv1.2 gating currents recorded with NMG\(^+\) displayed a greater left shift in \( Q_{\text{off}}(0 \text{ mV})-V \) than the ZHA Shaker model. To simulate the gating currents in the absence of any intrinsic stabilization, the open-state stabilizing factor \( \theta \) was set to 1 as shown in Fig. 7 C. In this case, the \( V_{0.5} \) for each \( Q_{\text{off}}-V \) curve was very similar to the \( Q_{\text{on}} \) (\(-46 \text{ mV}\)). This model recapitulated the data we gathered for the mutant channel Kv1.2-I402C, which showed a relief of open-state stabilization, suggesting that we might consider this channel’s kinetics consistent with an NMG\(^+\)-independent gating scheme (Fig. 6).

To assess the effect of internal NMG\(^+\) ions binding to the open state on the \( Q_{\text{off}}-V \) curves, we simulated the addition of 140 mM NMG\(^+\) ions into the model depicted in Fig. 7 using \( \theta = 1 \); that is, a channel that has no intrinsic open-state stabilization. Addition of internal NMG\(^+\) slowed \( \text{IgOFF} \) after depolarization to potentials at which the channel pore opens, recapitulating our observations for Kv1.2 gating currents (Fig. 1, A and B). The \( Q_{\text{off}}-V \) relationship shows that \( \sim 60\% \) of the \( Q_{\text{on}} \) is returned (Fig. 7 D), similar to the \( \sim 70\% \) we observed in Kv1.2 gating current recordings (Fig. 1 C). In addition, the \( Q_{\text{off}}(0 \text{ mV})-V \) curve was left shifted to \(-101 \text{ mV}\) as a result of the stabilization of the open state and consequent slowing of \( \text{IgOFF} \) analogous to the Kv1.2 gating currents recorded in the presence of NMG\(^+\) (Fig. 2 E). This simulation demonstrates that the effect of an intracellular cation trapping channels in the open state and rate-limiting voltage sensor return can quantitatively account for the slow OFF gating and resultant left shift in \( Q_{\text{off}}-V \) that we have observed in Kv1.2 gating currents.

**DISCUSSION**

Deactivation kinetics of \( K^+ \) channels are critically important in controlling membrane repolarization and influencing the frequency of action potential firing in excitable cells, yet the underlying mechanisms have remained unresolved. Here, we have demonstrated that the kinetics of deactivation and the associated voltage sensor return are controlled allosterically by ions occupying the inner cavity of open Kv1.2 channels, rather than by the intrinsic structural stability of the open pore domain alone.

**Evidence for an occupancy theory of open pore stability**

We found that intracellular ions affect \( \text{IgOFF} \) without substantially altering \( \text{IgON} \), suggesting that an activated state of the voltage sensor was stabilized. Slowing of \( \text{IgOFF} \) correlated with depolarization to voltages at which the channel \( P_o \) rapidly increases (greater than \(-30 \text{ mV}\); Fig. 1 C), strongly favoring the interpretation that slowing was mediated through the open pore. In addition, the time course of the onset of the slow OFF gating current was fast and approximated the time course of channel activation (Fig. 2 A), indicating that pore opening was the critical step, rather than inactivation, which was minimal and much slower in this channel (Fig. S1 A). We assessed the voltage dependence of charge return and found that the \( Q_{\text{off}}-V \) was markedly left shifted with respect to the \( Q_{\text{on}}-V \) (Fig. 2). This left shift is the result of stabilization of the activated state of the voltage sensor, as more negative voltages are required to return the sensors from activated positions than was originally needed to move them into the activated states (Lacroix et al., 2011). These data support the conclusion that ions within the open pore stabilized the voltage sensor.

\(-90 \text{ mV}, 1/\tau = 441 \pm 25.7 \text{ s}^{-1}\); and at \(-100 \text{ mV}, 1/\tau = 511.2 \pm 62.2 \text{ s}^{-1}\). The voltage dependence of the rate of charge was assessed by plotting the \( \text{IgOFF} \) rate against membrane voltage and fit with an exponential function of voltage (Eq. 1). (E) Representative normalized ionic tail currents recorded in 140 mM external \( K^+ \) from Kv1.2 WT (black traces) and Kv1.2-I402C (gray traces) channels at \(-120 \text{ mV} \) and \(-80 \text{ mV}. \) Currents were recorded between \(-50 \) and \(-120 \text{ mV} \) after a 50-ms pulse to \( 30 \text{ mV} \) to open channels fully. The \( t_{0.5 \text{ decay}} \) was plotted against the voltage to quantify the increased rate of Kv1.2-I402C deactivation kinetics. Error bars represent SEM.

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Figure 7. Kinetic modeling of intracellular NMG\(^+\) binding can account for the observed slowing of IgOFF kinetics. (A) Schematic of the Markov model used to simulate gating currents. (B, left) Simulated gating currents with the depicted voltage protocols and a slowing factor \(\theta\) of 9.4 as in the original ZHA model. (right) Normalized Q-V relationships calculated by integrating gating current for 11 ms after the depolarizing pulse \((Q_{on}/Q_{on(max)})\), repolarization \((Q_{off}/Q_{on(max)})\), or hyperpolarization after 50 ms at 0 mV \((Q_{off}/Q_{on(0 mV)})\). (C) Repeat of simulations in B, but with the removal of the slowing factor \((\theta = 1)\). (D) Repeat of simulations in B, but with the addition of 140 mM NMG\(^+\). Error bars represent SEM.
in the activated state. A clue to the underlying mechanism emerged upon observing that the species of intracellular cations could critically regulate the presence or absence of slow OFF gating current (Fig. 4).

The mechanism of open pore stabilization

The inner pore cavity of Kv channels contains a binding site for permeating ions located at approximately the same position as the quaternary ammonium binding site, just below the selectivity filter (Choi et al., 1993; M. Zhou et al., 2001; Y. Zhou et al., 2001; Lenaeus et al., 2005). Occupancy of this site by inactivation peptides or intracellular blockers causes slow return of gating charge (Bezanilla et al., 1991; Perozo et al., 1992; Melishchuk and Armstrong, 2001). In Kv channels, the pore of the channel must close before the voltage sensors can deactivate; thus, intracellular pore blockers have the effect of slowing the return of voltage sensors, which is supported by kinetic studies of Kv channels that place an open state accessed from the last closed state that imposes pore closure as a requirement before voltage sensors can transition back to resting states (Bezanilla et al., 1994; Zagotta et al., 1994a; Schoppa and Sigworth, 1998). For Kv1.2 specifically, voltage clamp fluorometry studies of gating support this model because return movement of S4 follows the kinetics of ionic current deactivation (Horne et al., 2010) and is slowed by the Kv1.2 subunit, which confers N-type inactivation (Peters et al., 2009).

We used the Kv1.2 open-state crystal structure to investigate the possibility of TEA' or NMG' binding in the inner cavity through a docking simulation (Fig. 8, A and B). The docking demonstrated that both NMG' and TEA' can interact in a binding pocket within the inner cavity. We tested this model with a kinetic model (Fig. 7) that recapitulated the emergence of a slow component in IgOFF that became dominant after greater depolarizations. This reflected the stabilization of the voltage sensor and also resulted in a left shift of the $Q_{off}$-$V$ relation, confirming that the stabilization of the open channel pore by NMG' or another intracellular ion could account for the slowing of IgOFF we observed.

The affinity of ions for the inner cavity binding site is the basis of open pore stabilization

We expected the affinity of NMG' and TEA' to be comparable because of the similar left shift in $Q_{off}$-$V$ we observed...
Ions within the cavity stabilize open channels

Repulsive electrostatic interactions between the positively charged ions within the conduction pathway critically regulate the affinity of individual binding sites to allow high flux rates in a conducting channel (Hille and Schwarz, 1978). Considering these repulsive forces, we predict that the removal of positively charged permeant ions stabilized the TEA+/NMG+ binding site in Kv1.2 and enabled a higher affinity interaction than would normally be present with permeant ions in the pore. This mechanism could explain the apparent paradox that the nonconducting P-type inactivated mutant channel in internal K+ has significantly slower Ig OFF (Fig. 5, A and B) than ionic deactivation (Fig. 6 E). The removal of K+ ions from the selectivity filter in the P-type inactivated channel could explain the significantly slower Ig OFF (Q_{OFF(0 mV)} - V_{0.5} NMG+ = -114.2 mV vs. TEA+, -126 mV; Figs. 2 E and 4 D). The K_d of NMG+ binding (k_{off(0 mV)}/k_{on(0 mV)}) derived from our kinetic model was 57 µM (Fig. 7), a higher affinity than the reported K_d for internal TEA+ in Shaker channels of 390 µM (Choi et al., 1993). The increased affinity of our interactions could be explained by the lack of any competing permeant ions in our experiments, which would lead to a unique high affinity binding environment at the inner cavity site (Fig. 9 B).

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Figure 9. Schematic diagram illustrating the proposed mechanism of open pore stabilization. (A) Channels in the absence of permeant ions or nonconductive mutant Kv1.2 W366F,V381T have vacant or collapsed selectivity filters. Upon depolarization, the channel activation gate opens, and intracellular ions can enter the inner cavity. Upon hyperpolarization, the pore cannot shut until the inner cavity is vacated, which rate limits the voltage sensor return. (B) In the presence of permeant ions K+ or Cs+, the selectivity filter is occupied. As the channel pore opens, the hydrated ions can enter the inner cavity, but the binding site is less stable because of strong ion-ion repulsive forces from the other ions in the selectivity filter. Upon hyperpolarization, the exit rate of the hydrated ions is faster as a result of repulsive forces, and the pore can shut faster than in the absence of permeant ions in the conduction pathway. SF, selectivity filter.
channel prevents the repulsive ion–ion interactions normally created within the highly occupied selectivity filter from displacing the intracellular K’ ion from its binding site, thus creating an atypical high affinity K’ binding site in the intracellular cavity (Fig. 9 A). This effect may also contribute in general to the increased slowing of IgOFF observed in channels that have undergone a pore-collapsing mutation or a P-type/slow inactivation process. Interestingly, internal Cs+ ions caused a similar rate of slowing in the conducting and nonconducting channels, which would suggest that the binding interactions of Cs+ ions within the inner cavity are not as strong as that of K’ but are still sufficient to be rate limiting to pore closure.

Although the ionic deactivation kinetics for K’ conductance are several-fold faster than the return of IgOFF in gating current experiments, the requirement for ions to vacate the pore in K’-containing solutions before closure is verified by our observation that ionic deactivation kinetics in the Kv1.2 1402C channel are threefold faster than the WT channel (Fig. 6 E). Therefore, the I402C channel also decreases any stabilizing effect that K’, and NMG+, ions occupying the pore may have produced.

Considering these data leads to a position in support of the occupancy hypothesis (Matteson and Swenson, 1986; Melishchuk and Armstrong, 2001) and provides a mechanistic basis for this effect. It follows that the critical determinant of open pore stability is not intrinsic intraprotein structural interactions but the allosteric effects of ionic occupancy of the inner cavity binding site creating an energy barrier to pore closure that has to be overcome.

The nature of the ion binding site in the inner cavity

It would be extremely informative in understanding our data to be able to examine a closed-state structure of a channel pore, but no crystal structures are currently available for closed Kv channels. Unfortunately, the KcsA closed-state structures are not a good model for the closed state of the Kv channel. KcsA has a considerably larger inner cavity than Shaker, as indicated by the fact that TBA, with a diameter of ~10 Å, has been crystallized and trapped in the inner cavity of the closed-state KcsA (Y. Zhou et al., 2001), but TEA (with a diameter of ~5 Å; Fig. 8 A) cannot be trapped in the closed cavity of the Shaker channel (Holmgren et al., 1997). In addition, cysteine modification experiments in the presence of intracellular TEA and TBA have demonstrated a pattern of reactivity in Shaker channels inconsistent with the linear pore lining helices of the KcsA channel, postulated to be related to the lack of a P-X-P helix breaking motif in the S6 of KcsA (del Camino et al., 2000). The KcsA structure does show that a fully hydrated K’ ion with eight water molecules resides in this cavity, indicating the possibility of a relatively stable site within a large cavity (Y. Zhou et al., 2001). The diameter of a hydrated K’ ion has been estimated at around 5.4 Å, a comparable size to the TEA+ ion, with a hydration energy of ±85 kcal/mol (Hille, 2001). The free energy difference between the open and closed states of the Shaker channel is estimated to be ~2 kcal/mol (Yifrach and MacKinnon, 2002). Energetically, a hydrated K’ ion residing in an open channel cavity could offer a significant barrier to channel pore closing because of the energetic costs associated with reorganizing the hydration shell.

The mechanism of slowing of Kv1.2 IgOFF kinetics is distinct from a voltage sensor conformational rearrangement

Slowing of IgOFF and the consequent left shifts in the Qoff-V relationship after prolonged depolarization have previously been linked to inactivation processes and a phenomenon known as voltage sensor relaxation, where the slowing of voltage sensor return is suggested to indicate a different pathway of voltage sensor transit from the relaxed state (Villalba-Galea et al., 2008; Lacroix et al., 2011). In our Kv1.2 experiments, however, the left shift of Qoff-V (Fig. 2) is unlikely to be associated with a relaxation process, as the durations of our depolarizing pulses are at most 50 ms, which is much slower than the time course over which relaxation is reported to occur (τ = ~1.8 s; Lacroix et al., 2011), and, indeed, we found no evidence of the Kv1.2 channel significantly inactivating over a greater time course than of our gating current recordings (Fig. S1 B). This leads to the conclusion that relaxation and/or inactivation cannot be the sole cause of slow return of OFF gating currents in Kv channels after depolarization.

Ion binding in the inner cavity offers an extrinsic mechanism by which channel deactivation kinetics are modulated

In summary, our results suggest that ion occupancy of the inner cavity of Kv1.2 channels impairs pore closure by slowing the open to closed transition and is therefore the cause of slowed IgOFF after depolarizations that open channels. The affinity of the ion for the cavity then becomes a primary determinant of pore closure, as depicted in Fig. 9 (A and B). Our data highlight the role that the channel inner cavity may play in fine-tuning deactivation kinetics of different channels, as variation in ion binding affinities and/or dissociation rates significantly affect deactivation kinetics. In essence, this represents an allosteric mechanism, separate from the VSD, through which it is possible for Kv channels to display varied deactivation kinetics.

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