Interaction between residues in the Mg\(^{2+}\)-binding site regulates BK channel activation

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As a unique member of the voltage-gated potassium channel family, a large conductance, voltage- and Ca\(^{2+}\)-activated K\(^+\) (BK) channel has a large cytosolic domain that serves as the Ca\(^{2+}\) sensor, in addition to a membrane-spanning domain that contains the voltage-sensing (VSD) and pore-gate domains. The conformational changes of the cytosolic domain induced by Ca\(^{2+}\) binding and the conformational changes of the VSD induced by membrane voltage changes trigger the opening of the pore-gate domain. Although some structural information of these individual functional domains is available, how the interactions among these domains, especially the noncovalent interactions, control the dynamic gating process of BK channels is still not clear. Previous studies discovered that intracellular Mg\(^{2+}\) binds to an interdomain binding site consisting of D99 and N172 from the membrane-spanning domain and E374 and E399 from the cytosolic domain. The bound Mg\(^{2+}\) at this narrow interdomain interface activates the BK channel through an electrostatic interaction with a positively charged residue in the VSD. In this study, we investigated the potential interdomain interactions between the Mg\(^{2+}\)-coordination residues and their effects on channel gating. By introducing different charges to these residues, we discovered a native interdomain interaction between D99 and E374 that can affect BK channel activation. To understand the underlying mechanism of the interdomain interactions between the Mg\(^{2+}\)-coordination residues, we introduced artificial electrostatic interactions between residues 172 and 399 from two different domains. We found that the interdomain interactions between these two positions not only alter the local conformations near the Mg\(^{2+}\)-binding site but also change distant conformations including the pore-gate domain, thereby affecting the voltage- and Ca\(^{2+}\)-dependent activation of the BK channel. These results illustrate the importance of interdomain interactions to the allosteric gating mechanisms of BK channels.

INTRODUCTION

The large conductance, voltage- and Ca\(^{2+}\)-activated K\(^+\) (BK) channel plays important roles in many physiological processes, such as muscle contraction, neural transmission, and cardiac pacemaker output (Brayden and Nelson, 1992; Robitaille et al., 1993; Meredith et al., 2006). A functional BK channel is composed of four Slo1 subunits, each of which comprises multiple structural modules (Fig. 1, A and B): a membrane-spanning domain including the pore-gate (S5 and S6) and voltage-sensing domain (VSD; S1 to S4), and a large cytosolic domain that contains two Ca\(^{2+}\)-binding sites to serve as the Ca\(^{2+}\) sensor (Adelman et al., 1992; Schreiber and Salkoff, 1997; Xia et al., 2002; Zhang et al., 2010). Conformational changes of the VSD upon membrane depolarization and conformational changes of the cytosolic domain upon Ca\(^{2+}\)-binding control BK channel activation.

Intracellular Mg\(^{2+}\) activates BK channels (Golowasch et al., 1986; Oberhauser et al., 1988; Shi and Cui, 2001; Zhang et al., 2001) by binding to the interface between the cytosolic and membrane-spanning domains (Shi et al., 2002; Xia et al., 2002; Yang et al., 2008). The Mg\(^{2+}\)-binding site is composed of residues E374 and E399 that are located in the cytosolic domain and D99 and N172 in the membrane-spanning domain (Fig. 1, A and B). The bound Mg\(^{2+}\) activates the channel by repulsing gating charge R213 in the voltage sensor S4 through an electrostatic interaction (Hu et al., 2003; Yang et al., 2007, 2008) (Fig. 1 A). On the other hand, the oxygen atoms from the four Mg\(^{2+}\)-coordination residues contribute to the formation of an octahedral sphere for Mg\(^{2+}\) binding with Mg-O internuclear distance ranging from 2 to 2.2 Å (Dudev and Lim, 2003). Nevertheless, it is not known whether these physically proximate coordination residues, three of which are...
acids with negative charges, interact with one another in the absence of Mg$^{2+}$, and if such interactions indeed exist, how they contribute to the allosteric mechanism of BK channel gating in general.

In this study, we discovered that a native interdomain interaction between Mg$^{2+}$-coordination residues D99 and E374, located in the membrane-spanning domain and the cytosolic domains, respectively, inhibits the activation of the WT BK channel. Further analysis indicates that both D99 and E374 also interact with other residues to affect BK channel activation, making the function of this native interaction difficult to resolve. Therefore, to further study the effects of the interactions between Mg$^{2+}$-coordination residues on the structure and activation of BK channels, we engineered different charges at the position of neutral residue N172 in the Mg$^{2+}$-binding site. We found that the charges at position 172 sense the charges at residue 399, and this electrostatic interaction regulates fundamental aspects of BK channel activation, including voltage- and Ca$^{2+}$-dependent activation and the intrinsic open probability of the activation gate. Our current work thus suggests that interaction between Mg$^{2+}$-binding coordination residues participates in BK channel activation.

**MATERIALS AND METHODS**

**Mutagenesis and expression**

The mutations were made from the *mbr5* splice variant of mSlo1 (provided by L. Salkoff, Washington University, St. Louis, MO) (Butler et al., 1993) (available from GenBank accession no. GI:347143) using overlap-extension PCR. The PCR-amplified regions for all the mutations were verified by sequencing. RNA was transcribed in vitro with T3 polymerase (Ambion) and injected into *Xenopus laevis* oocytes (stage IV–V), each with an amount of 0.05–50 ng. Experiments were performed after 2–5 d of incubation at 18°C. The procedures for harvesting oocytes and housing *Xenopus* were approved by the Washington University Animal Studies Committee and performed according to the protocols.

**Electrophysiology**

Inside-out patches were formed from oocyte membrane by borosilicate pipettes of 0.8–1.5-MΩ resistance. Macroscopic currents were recorded using a patch-clamp amplifier (Axopatch 200-B; Molecular Devices) and PULSE acquisition software (HEKA). The current signals were low-pass filtered at 10 kHz by the amplifier’s four-pole Bessel filter and digitized with 20-µs intervals. The pipette (extracellular) solution comprised (mM): 140 potassium methanesulphonic acid, 20 HEPES, 2 KCl, and 2 MgCl$_2$, pH 7.2. Intracellular solutions contained: 140 mM potassium methanesulfonic acid, 20 mM HEPES, 2 mM KCl, 1 mM EGTA, and 22 mg/L 18GluP, pH 7.2. CaCl$_2$ standard solution was added to obtain the desired free [Ca$^{2+}$] from 1.0 to 112 µM, which was verified by a Ca$^{2+}$-sensitive electrode (Thermo Electron). The nominal 0-µM [Ca$^{2+}$] solution contained an additional 4 mM EGTA with no added Ca$^{2+}$, and the free [Ca$^{2+}$] was calculated to be ~0.5 nM. EGTA was excluded from the intracellular solution containing 200 µM [Ca$^{2+}$]. The treatment of MTS reagents or DTT was performed by perfusing the intracellular side of the excised patch for 5 min with the corresponding solution. Before each experiment, the nominal 0-µM [Ca$^{2+}$], solution was used to dilute the stock MTS reagents (500x) or DTT (10x) to the final concentration of 1 mM (MTSES), 0.2 mM (MTSET), 1 mM (MTSACE), or 10 mM (DTT). The MTSET solution was freshly prepared right before each perfusion because its lifetime is ~10 min, whereas the MTSES and MTSACE solutions were prepared every 30 min. The DTT solution was freshly prepared and used for the day. G-V relations of the same mutation before and after chemical modifications were obtained using different groups of patches. All the experiments were performed at room temperature (22–24°C).

**Analysis**

The tail current amplitudes at ~80 mV were measured to determine the relative conductance. Each G-V relation was normalized and fitted with the Boltzmann equation:

$$
G / G_{\text{max}} = \frac{1}{1 + \exp \left( \frac{-z(V - V_{1/2})}{kT} \right)}
$$

where $G / G_{\text{max}}$ is the ratio of conductance to maximal conductance, $z$ is the number of equivalent charges, $e$ is the elementary charge, $V$ is membrane potential, $V_{1/2}$ is the voltage where $G / G_{\text{max}}$ reaches 0.5, $k$ is Boltzmann’s constant, and $T$ is absolute temperature. Error bars in this paper represent SEM unless stated otherwise. Unpaired Student’s t test or two-way ANOVA were performed to detect a significant difference among the results, and a p-value smaller than 5% is considered significant. A p-value of multiple comparison is adjusted using the Bonferroni correction. The sample size in statistical analysis is between 3 and 16 in this paper.

**Open probability at negative voltages**

The open probability at negative voltages was measured by single-channel recordings using patches containing hundreds of channels (Horrigan et al., 1999; Cui and Aldrich, 2000). The open probability measured at negative voltages is combined with the corresponding G-V relation to construct a $P_e V$ relation, which is fitted to the following model, based on the conceptual framework provided by Horrigan, Cui, and Aldrich (HCA; Horrigan et al., 1999):

$$
P_e = \frac{1}{1 + \exp \left( \frac{-z_e F (V - V_0)}{RT} \right)} \times \frac{1 + \exp \left( \frac{-z_F (V - V_e)}{RT} \right)}{1 + \exp \left( \frac{-z_F (V - V_{e0})}{RT} \right)}
$$

where $z_e$ is the charge associated with gate opening when all the voltage sensors are at their resting state, $z_f$ is the charge associated with voltage-sensor movements, and $L_o$ is the intrinsic open probability at $V = 0$ while all the voltage sensors are at their resting state. $V_0$ and $V_e$ are the voltages for half of the voltage sensors to be at their activation state at the closed and the open conformations of the gate, respectively (Horrigan and Aldrich, 2002).

**RESULTS**

A native interdomain interaction within the Mg$^{2+}$-binding site participates in channel activation

To probe the native interaction that might exist between negatively charged D99 and E374/E399, we introduced positively charged Arg mutation to each of these residues and tested their perturbation on channel
To examine if the interaction between D99 and E374 depends on electric charges, we measured the effect of E374R on the G-V relation in the background of mutation D99A; E374R has a smaller effect on the G-V relation (Fig. 1 E), consistent with an electrostatic interaction between D99 and E374.

During further study of this interdomain interaction, we found that residues D99 and E374 are involved in interactions with other sites such as R213 in the voltage sensor S4 (unpublished data). To further understand the role of the interdomain interaction within the Mg2+-binding site in channel activation without the interference of other types of interactions, we engineered artificial electrostatic interactions between residues at positions 172 and 399 and characterized the effects of these interactions in the remaining part of this work.

Figure 1. D99 and E374 interact with each other at the domain interface of BK channels. (A) Cartoon showing the molecular mechanism of Mg2+ activation. S0–S6 are membrane-spanning segments. Mg2+ is coordinated by four residues including D99 and N172 from the membrane-spanning domain and E374 and E399 from the cytosolic domain. Bound Mg2+ interacts with R213 of the membrane-spanning domain to activate channel. Located in the vicinity of the Mg2+-binding site is residue Q397, which may interact with R213 when charged. (B) Cartoon showing the spontaneous formation of a disulfide bond between C99 and C397 located in the membrane-spanning and cytosolic domains, respectively. (C) Macroscopic current traces of inside-out patches expressing WT BK channels and D99R, E374R, D99R/E374R, and E399R mutant channels. Currents were elicited by voltages ranging from 0 to 200 mV, with 50-mV increments. The voltage was −50 mV before and −80 mV after the pulses. [Ca2+]: 0 µM. Bars, 1 nA. (D) G-V relations of WT, D99R, E374R, D99R/E374R, and E399R mutant channels. [Ca2+]: 0 µM. Solid lines are fittings to the Boltzmann equation. The fitting parameters are: WT (n = 4): V1/2 = 187 ± 4 and z = 1.29 ± 0.12; D99R (n = 14): V1/2 = 170 ± 3 and z = 1.31 ± 0.03; E374R (n = 6): V1/2 = 136 ± 2 and z = 1.29 ± 0.04; D99R/E374R (n = 5): V1/2 = 195 ± 2 and z = 1.21 ± 0.05; E399R (n = 3): V1/2 = 183 ± 5 and z = 1.34 ± 0.10. (E) V1/2 of the G-V relations. The fitting parameters for D99A and D99A/E374R are: D99A (n = 6): V1/2 = 198 ± 5 and z = 1.20 ± 0.05; D99A/E374R (n = 7): V1/2 = 186 ± 2 and z = 1.24 ± 0.06.
Electrostatic interactions between residues 172 and 399 after channel activation

As is the case with the residue pair of D99 and E374, N172 and E399 are also part of the Mg$^{2+}$-binding site (Shi et al., 2002; Yang et al., 2008) (Fig. 1, A and B). Shifts of the G-V curves to more positive voltages were observed when we introduced an artificial electrostatic interaction between positions 172 and 399 by modifying the double cysteine mutation (N172C/E399C) with charged MTS reagents, MTSET(+) and MTSES(−) (Fig. 2, A and B). Neither MTSET nor MTSES affects the control channel (C430A; Fig. 2 C), indicating that these MTS reagents react with the cysteines on residues 172 and 399. In contrast to the effect of MTSET(+) and MTSES(−), modification with neutral MTSACE did not affect the G-V relation of N172C/E399C (Fig. 2 D). Moreover, a subsequent MTSET(+) treatment failed to shift the G-V relation, indicating that both C172 and C399 were covalently modified by neutral MTSACE so that no free thiol groups at these two positions were available to react with MTSET(+). These results suggest that the engineered two charges with the same sign at 172 and 399 hinder channel activation through electrostatic repulsion with each other. Consistent with this mechanism, the effects of the charged MTS reagents on channel activation in 200 µM Ca$^{2+}$ were attenuated in 1 M NaCl intracellular solution (Fig. 3, C and D), indicating that increasing ionic strength indeed weakens the electrostatic repulsion between these two sites. The osmolarity change brought by 1 M NaCl has no effect on BK channel activation, as shown previously (Yang et al., 2010). Although ionic strength also affects WT channel activation in other [Ca$^{2+}$], the increase of ionic strength in 200 µM Ca$^{2+}$ had no effect on control channel activation (Fig. 3, A and B). Therefore, the effect of the charged MTS reagents on channel activation is caused by electrostatic interactions that are sensitive to the change of ionic strength of the solution.

Figure 2. Electrostatic repulsion between residues 172 and 399 inhibits channel activation. (A) Macroscopic current traces of inside-out patches expressing N172C/E399C before and after treatment of negative (ES) or positive (ET) reagent. Currents were elicited by voltages of −50, 0, 50, and 100 mV. The voltage was −50 mV before and −120 mV (top) or −80 mV (middle and bottom) after the pulses. [Ca$^{2+}$]: 200 µM. (B) G-V relation of N172C/E399C shifts after treatment of ES or ET reagent. Open symbols, [Ca$^{2+}$]: 0 µM; closed symbols, [Ca$^{2+}$]: 200 µM. Solid lines are fittings to the Boltzmann equation. The fitting parameters are: N172C/E399C untreated, 0 [Ca$^{2+}$], (n = 3): $V_{1/2} = 197 ± 5$ and $z = 1.19 ± 0.22$; 200 µM [Ca$^{2+}$], (n = 7): $V_{1/2} = 2 ± 7$ and $z = 1.20 ± 0.09$; after MTSES treatment, 0 [Ca$^{2+}$], (n = 3): $V_{1/2} = 250 ± 25$ and $z = 0.86 ± 0.60$; 200 µM [Ca$^{2+}$], (n = 7): $V_{1/2} = 47 ± 6$ and $z = 1.13 ± 0.27$; after MESET treatment, 0 [Ca$^{2+}$], (n = 3): $V_{1/2} = 271 ± 10$ and $z = 1.19 ± 0.44$; 200 µM [Ca$^{2+}$], (n = 4): $V_{1/2} = 61 ± 1$ and $z = 1.13 ± 0.03$. (C) G-V relation of the control C430A after treatment of ES or ET reagent. Open symbols: [Ca$^{2+}$]: 0 µM; closed symbols: [Ca$^{2+}$]: 200 µM. Solid lines are fittings to the Boltzmann equation. The fitting parameters are: C430A untreated, 0 [Ca$^{2+}$], (n = 7): $V_{1/2} = 198 ± 2$ and $z = 1.13 ± 0.03$; 200 µM [Ca$^{2+}$], (n = 6): $V_{1/2} = −19 ± 4$ and $z = 1.48 ± 0.08$; after MTSES treatment, 0 [Ca$^{2+}$], (n = 6): $V_{1/2} = 204 ± 6$ and $z = 1.11 ± 0.04$; 200 µM [Ca$^{2+}$], (n = 3): $V_{1/2} = −7 ± 4$ and $z = 1.26 ± 0.13$; after MESET treatment, 0 [Ca$^{2+}$], (n = 3): $V_{1/2} = 195 ± 2$ and $z = 1.00 ± 0.06$; 200 µM [Ca$^{2+}$], (n = 3): $V_{1/2} = −14 ± 6$ and $z = 1.54 ± 0.49$. (D) G-V relations of N172C/E399C after treatment of neutral MTSACE (ACE) and/or positive ET. [Ca$^{2+}$]: 0 µM. Solid lines are fittings to the Boltzmann equation. The fitting parameters are: N172C/E399C untreated (n = 3): $V_{1/2} = 197 ± 5$ and $z = 1.19 ± 0.22$; after MTSES treatment (n = 4): $V_{1/2} = 199 ± 5$ and $z = 1.11 ± 0.21$; after MTSET treatment (n = 3): $V_{1/2} = 271 ± 10$ and $z = 1.19 ± 0.44$; after MESET and MTSET treatments (n = 3): $V_{1/2} = 207 ± 4$ and $z = 1.19 ± 0.22$. N172C/E399C mutation is on the background of C430A to eliminate the effects of MTS reagents on the native C430 (Zhang and Horrigan, 2005).
residue 399 was neutral (Fig. 4 B). This suggests that the charges at residue 172 probably interact with other charges besides E399. By testing the voltage at half-maximum of G-V relation (V1/2) against the charge type at 172 and 399 for 26 mutations and chemical modifications, ANOVA detects a significant difference among the V1/2 values with different charge type at residue 172 (P = 0.049) but not with the charge type at residue 399 (P = 0.364) (Fig. 4 C and Table 1). However, although charges at 399 do not interact with other charges, the

Figure 3. Electrostatic interactions between residues 172 and 399 depend on ionic strength. (A–D) G-V relations in control solution (200 µM [Ca2+]i solution) or in control solution with an additional 1 M NaCl for the control (A), N172C/E399C (B), N172C/E399C after MTSES (C), or N172C/E399C after MTSET (D). All G-V curves are fitted to the Boltzmann equation (lines). The fitting parameters are: the control channel under control solution (n = 6): V1/2 = −19 ± 4 and z = 1.48 ± 0.08; under an additional 1 M NaCl (n = 3): V1/2 = −24 ± 3 and z = 1.36 ± 0.08; N172C/E399C under control solution (n = 7): V1/2 = 2 ± 7 and z = 1.20 ± 0.09; under an additional 1 M NaCl (n = 3): V1/2 = 7 ± 2 and z = 1.32 ± 0.11; under an additional 1 M NaCl (n = 3): V1/2 = 17 ± 2 and z = 1.29 ± 0.05; N172C/E399C after MTSES under control solution (n = 4): V1/2 = 60 ± 1 and z = 1.13 ± 0.03; under an additional 1 M NaCl (n = 4): V1/2 = 39 ± 2 and z = 1.14 ± 0.03. The control and N172C/E399C mutation are on the background of C430A.

Figure 4. Effects of charge type at residues 172 and 399 on channel activation. G-V relation is shifted by charge type at residue 172 (B) but not by residue 399 (A). [Ca2+]i: 0. Solid lines are fittings to the Boltzmann equation. The fitting parameters are: N172Q (n = 4): V1/2 = 181 ± 5 and z = 1.11 ± 0.20; N172Q/E399N (n = 5): V1/2 = 186 ± 4 and z = 1.15 ± 0.18; N172Q/E399R (n = 5): V1/2 = 184 ± 4 and z = 1.04 ± 0.17; E399C/N172D (n = 8): V1/2 = 188 ± 4 and z = 1.25 ± 0.23; E399C/N172R (n = 5): V1/2 = 132 ± 5 and z = 1.30 ± 0.29; E399C (n = 6): V1/2 = 201 ± 4 and z = 1.23 ± 0.22. (C) V1/2 of a variety of N172 and E399 mutations with or without chemical modifications listed in Table 1 versus charge type at residue 172 (left) or 399 (right). Solid lines are fittings to linear regression. The slope ± fitting SD of linear regression is −16.1 ± 7.5 for the charge type at 172 and −2.6 ± 7.4 for the charge type at 399. (D) V1/2’s of mutations in Table 1 are averaged based on the interaction type between 172 and 399.
data in Table 1 suggest that charges at 399 affect channel activation by interacting with the charges at 172 because ANOVA shows that the interaction between 172 and 399 affects channel activation with a p-value of 0.0006 (Fig. 4 D and Table 1).

Thus, channel activation is altered by the electrostatic interaction between the charges at residues 172 and 399. Different types of interaction—repulsion, attraction, or no interaction—can be introduced to cause different effects on channel activation. To avoid the complication of varying the charge at 172, which interacts with other charges in addition to that at 399, we changed the interaction type by varying the charge at 399 and keeping the same residue at 172 for each set of the following experiments.

Electrostatic interaction between residues 172 and 399 alters voltage and Ca\textsuperscript{2+} dependence

BK channels are activated by voltage and Ca\textsuperscript{2+} via distinct allosteric mechanisms (Horrigan and Aldrich, 2002; Cui et al., 2009). Would interactions between the Mg\textsuperscript{2+}-coordination residues specifically alter voltage or Ca\textsuperscript{2+}-dependent activation, or do such interactions affect both mechanisms? To address this question, we examined the voltage-dependent activation (measured as $V_{1/2}$ of the G-V relation in 0 [Ca\textsuperscript{2+}]) and Ca\textsuperscript{2+}-dependent activation (measured as shift of the G-V, $\Delta V_{1/2}$, caused by an increase of [Ca\textsuperscript{2+}], from 0 to the saturating 112 µM) for several mutations and found that both are affected by the 172–399 electrostatic interactions (Fig. 5). Mutations of E399 to neutral (C or N) and positive charge (R) altered both voltage (Fig. 5 A) and Ca\textsuperscript{2+} (Fig. 5 B) -dependent activation on the background of N172R or N172D, but had no effect on channel activation on the background of WT or N172Q where residue 172 is neutral. Interestingly, the results show that regardless of the charge type at residues 172 or 399, a repulsive 172–399 interaction shifted the voltage dependence to higher voltage ranges (Fig. 5 A) and increased Ca\textsuperscript{2+} sensitivity (Fig. 5 B) as compared with that of an attractive 172–399 interaction in

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The charge type is listed for each mutation using symbols (+, positive; −, negative; 0, neutral), along with the amino acid and MTS reagent. As a result, the electrostatic interaction between 172 and 399 can be repulsion (Rep, between like charges), attraction (Attr, between opposite charges), or 0 (not both are charged). The p-values are calculated using ANOVA test (see Materials and methods). Mutations 1–18 are on the background of C430A.

\textsuperscript{a}A p-value of <0.05 indicates that a significant difference among the $V_{1/2}$ values was detected with different charge or interaction type.
but with larger intervals at low [Ca2+]i for N172R/E399R, such that the G-V curves cluster at high [Ca2+]i (Fig. 5, E and F). Consistently, the distribution of G-V curves of N172R/E399C shows intermediate distribution along the voltage axis (Fig. 5, D and F). On the other hand, the slope of G-V relations does not vary significantly at different [Ca2+]i or among different mutant channels (Fig. 5, C–F). These results indicate that the BK channel becomes more sensitive to both saturating and intermediate [Ca2+]i when the 172–399 interaction changes from attraction to repulsion. Based on the above experiments, we conclude that both voltage- and the same group. This result is similar to that of the native interaction between D99 and E374 (Fig. 1).

The effect of 172–399 interactions on Ca2+-dependent activation is not limited to the activation by saturating Ca2+ concentration; it also affects the activation by intermediate Ca2+ concentration. Fig. 5 (C–E) shows the G-V relations with varying [Ca2+]i, ranging from 0 to 112 µM for mutations of E399 to C or R on the background of N172R. The mutations change the 172–399 interaction from attraction to none or repulsion. With increasing [Ca2+]i, the G-V relations shift on the voltage axis with approximately equal intervals for N172R (Fig. 5, C and F).

Figure 5. The 172–399 electrostatic interaction alters voltage- and Ca2+-dependent activation. (A) V1/2 at 0 [Ca2+]i indicating changes in voltage-dependent activation. The mutations are grouped such that the negative residue E399 (open bars) is mutated to neutral (gray bars) or positive (closed bars) residues on the backgrounds where residue 172 contains a negative, positive, or no charge. (B) ΔV1/2 = V1/2 at 0 [Ca2+]i − V1/2 at 112 µM [Ca2+]i, indicating Ca2+ activation by saturating [Ca2+]i. Mutations are the same as in A. Asterisks in A and B indicate significant difference between the labeled pairs as identified in multiple comparisons (P < 0.017 as adjusted by the Bonferroni correction). The group of N172R mutations in A and B are made on the background of C430A. There is no significant difference in the number of equivalent gating charges among all the G-V relations. (C–E) G-V relation for N172R (C), N172R/E399C (D), and N172R/E399R (E) in varying [Ca2+]i. Solid lines are fittings to the Boltzmann equation. The fitting parameters of the mutations in A–E are:

- N172R: 0 Ca (n = 4): V1/2 = 139 ± 1 and z = 1.24 ± 0.05; 1.0 Ca (n = 4): V1/2 = 91 ± 2 and z = 1.43 ± 0.09; 4.6 Ca (n = 4): V1/2 = 48 ± 3 and z = 1.28 ± 0.06; 9.3 Ca (n = 4): V1/2 = 23 ± 4 and z = 1.37 ± 0.02; 112 Ca (n = 3): V1/2 = −12 ± 3 and z = 1.33 ± 0.07; 4.6 Ca (n = 4): V1/2 = 36 ± 2 and z = 1.41 ± 0.03; 9.3 Ca (n = 2): V1/2 = −10 ± 1 and z = 1.48 ± 0.02; 112 Ca (n = 3): V1/2 = −27 ± 5 and z = 1.48 ± 0.04; N172R/E399R 0 Ca (n = 4): V1/2 = 172 ± 4 and z = 1.29 ± 0.05; 1.0 Ca (n = 4): V1/2 = 118 ± 8 and z = 1.30 ± 0.14; 4.6 Ca (n = 4): V1/2 = 41 ± 4 and z = 1.25 ± 0.07; 9.3 Ca (n = 4): V1/2 = 14 ± 2 and z = 1.30 ± 0.09; 112 Ca (n = 3): V1/2 = −4 ± 6 and z = 1.34 ± 0.05; N172D 0 Ca (n = 4): V1/2 = 20 ± 4 and z = 1.18 ± 0.20; 112 Ca (n = 3): V1/2 = 20 ± 2 and z = 0.96 ± 0.15; N172D/E399N 0 Ca (n = 10): V1/2 = 200 ± 4 and z = 1.26 ± 0.07; 112 Ca (n = 2): V1/2 = 32 ± 10 and z = 1.34 ± 0.04; N172D/E399R 0 Ca (n = 4): V1/2 = 153 ± 5 and z = 1.28 ± 0.07; 112 Ca (n = 4): V1/2 = 16 ± 2 and z = 1.02 ± 0.02; WT 0 Ca (n = 7): V1/2 = 164 ± 2 and z = 1.18 ± 0.05; 112 Ca (n = 5): V1/2 = −7 ± 2 and z = 1.55 ± 0.07; E399N 0 Ca (n = 9): V1/2 = 170 ± 4 and z = 1.18 ± 0.03; 112 Ca (n = 5): V1/2 = −1 ± 3 and z = 1.59 ± 0.07; E399R 0 Ca (n = 7): V1/2 = 172 ± 3 and z = 1.30 ± 0.09; 112 Ca (n = 7): V1/2 = −7 ± 6 and z = 1.52 ± 0.05; N172Q 0 Ca (n = 4): V1/2 = 181 ± 2 and z = 1.11 ± 0.09; 112 Ca (n = 5): V1/2 = −22 ± 2 and z = 1.42 ± 0.05; N172Q/E399N 0 Ca (n = 5): V1/2 = 186 ± 3 and z = 1.11 ± 0.07; 112 Ca (n = 4): V1/2 = −14 ± 4 and z = 1.43 ± 0.05; N172Q/E399R 0 Ca (n = 5): V1/2 = 184 ± 3 and z = 1.10 ± 0.02; 112 Ca (n = 4): V1/2 = −13 ± 3 and z = 1.46 ± 0.07. All these mutations are on the C430A background. (F) ΔV1/2 (open symbols) and z (closed symbols) versus [Ca2+]i for the G-V relations in C–E.
Ca²⁺-dependent activation are altered by the electrostatic interaction between residues 172 and 399.

Electrostatic interaction between residues 172 and 399 alters local conformation
Because the Mg²⁺-binding site is not near the Ca²⁺-binding site according to the recent crystal structures of the cytosolic domain (Y. Wu et al., 2010; Yuan et al., 2010, 2012), the interaction between residues 172 and 399 has to alter the Ca²⁺-dependent activation through allosteric effects. To understand the underlying structural mechanism causing these allosteric effects, we first tested whether the 172–399 interaction could alter local conformation in the vicinity of the Mg²⁺-binding site.

The effect of 172–399 electrostatic interactions on the spontaneous formation of a disulfide bond between C99 and C397 (Yang et al., 2008) was monitored. Residues 99 and 397 are located in the vicinity of residues 172 and 399. Previously, we have shown that cysteines at these two residues spontaneously form a disulfide bond (Fig. 1 B) (Yang et al., 2008). The formation of disulfide bonds requires proper orientation of the two cysteine residues and the distance between the Cα atoms of the two cysteine residues to be within the range of 2.9 to 4.6 Å (Hazes and Dijkstra, 1988). If the 172–399 interaction alters the Cα–Cα distance to be outside of this range or changes their relative orientation, the formation of the disulfide bond would be prevented. Our previous studies

![Figure 6](image-url)

Figure 6. The 172–399 electrostatic interaction disrupts the formation of the interdomain disulfide bond. (A) G-V relations before (open circles) and after (closed symbols) treatment of MTSET (ET), MTSES (ES), or DTT when E399 is mutated to Asn (middle) or Arg (right) on the background of D99C/Q397C/N172R (left). “DTT, ET” in the middle panel represents the treatment of DTT followed by MTSET. [Ca²⁺]: 0. Solid lines are fittings to the Boltzmann equation. The fitting parameters are: D99C/Q397C/N172R untreated (n = 8): V1/2 = 153 ± 3 and z = 1.3 ± 0.02; after ET (n = 5): V1/2 = 140 ± 2 and z = 1.37 ± 0.04; after DTT (n = 6): V1/2 = 147 ± 3 and z = 1.28 ± 0.06; D99C/Q397C/N172R/E399N untreated (n = 6): V1/2 = 183 ± 3 and z = 1.14 ± 0.07; after ET (n = 8): V1/2 = 187 ± 5 and z = 1.07 ± 0.05; after DTT (n = 11): V1/2 = 161 ± 3 and z = 1.26 ± 0.04; after DTT and ET (n = 3): V1/2 = 218 ± 5 and z = 0.88 ± 0.06; D99C/Q397C/N172R/E399R untreated (n = 9): V1/2 = 203 ± 4 and z = 1.12 ± 0.05. (B) V1/2 of the G-V relations shown in A for the corresponding mutations and chemical modifications. Cartoons on the top show the type of the 172–399 electrostatic interactions (arrows) as a result of residue 399 charges and its consequence on the formation of the disulfide bond between D99C and D397C (line) based on the results (see Results). Asterisks indicate significant difference between labeled pairs as identified in multiple comparisons (P < 0.017 for the left and right panels and P < 0.0083 for the middle panel as adjusted by the Bonferroni correction). All the mutations are on the background of C430A.
showed that the formation of the C99–C397 disulfide bond restricts channel activation; thus, breaking the disulfide bond by DTT treatment would facilitate channel activation (Yang et al., 2008). In addition, when C397 was modified by MTSET(+) or MTSES(−), the charge attached to C397 would interact with R213 in S4 to change channel activation (Yang et al., 2008), whereas the C99–C397 disulfide bond protects C397 from MTS modification so that MTSET(+) or MTSES(−) treatment should not alter channel activation. If no spontaneous disulfide bond is formed between C99 and C397, the thiol group on C397 will be available for MTSET(+) or MTSES(−) modification, resulting in a shift of G-V relation, whereas DTT treatment should no longer have any effect on G-V relation.

As shown in Fig. 6 (A and B, left), adding a positive charge to residue 172 (N172R) to the double cysteine mutation of D99C/Q397C made the resulting mutant channel no longer sensitive to DTT treatment but more activated after MTSET(+) treatment, suggesting that the electrostatic attraction between residues 172 and 399 prevented the spontaneous formation of the C99–C397 disulfide bond. Similar results were observed when converting the electrostatic interaction between residues 172 and 399 from attraction to repulsion by introducing two positively charged Arg residues at both sites (Fig. 6 A and B, right). In this experiment, we used MTSES(−) instead of MTSET(+) because a positive environment of N172R/E399R may potentially repulse the positively charged MTSET(+) to prevent it from modifying C397 (Elinder et al., 2001; D. Wu et al., 2010). These results indicate that both attractive and repulsive 172–399 interactions disrupt the spontaneous formation of the interdomain disulfide bond between C99 and C397. To further verify this, we abolished the 172–399 electrostatic interaction by removing the negative charge on residue 399 (E399N) of the D99C/Q397C/N172R triple mutant channel (Fig. 6, A and B, middle). The resulting D99C/Q397C/N172R/E399N channel can no longer be modified by MTSET(+), indicating the formation of the spontaneous disulfide bond between C99 and C397. Treating this mutant channel with DTT broke the disulfide bond and shifted the G-V relation to more negative voltages. The subsequent treatment with MTSET(+) shifted the G-V relation to even more positive voltages, further indicating the formation of the C99–C397 disulfide bond when the electrostatic interaction between residues 172 and 399 is absent. Collectively, the above chemical treatment experiments indicate that the electrostatic interaction between residues 172 and 399 can alter the distance or relative orientation between C99 and C397, thereby preventing the formation of the spontaneous disulfide bond. Therefore, the interaction between residues 172 and 399 induces conformational changes in the surrounding structures. It is worth noting that chemical modification resulted in a shallower slope of the G-V relation (Fig. 6). This is probably caused by incomplete modification such that the G-V relation is a mixture of the modified and intact channel species.

Electrostatic interaction between residues 172 and 399 alters intrinsic open probability of the pore

Besides its impact on channel activation by voltage and Ca2+, the 172–399 interaction also alters intrinsic properties of the channel, such as the open probability of the activation gate in the absence of voltage-sensor activation and Ca2+ binding. Previous studies have shown that the BK channel conducts K+ current with a small but finite Po at 0 [Ca2+]i and negative voltages (less than −80 mV) where the voltage sensors are at the resting state (Horrigan et al., 1999). Such an intrinsic Po without activation of the sensors is determined by the properties of the pore-gate and its interaction with other structure domains and thus can be used to detect conformational changes in the pore-gate domain caused by the 172–399 interaction. The intrinsic Po of BK channels is obtained by recording the unitary currents in a patch that contains hundreds of channels at negative voltages (less than −20 mV) (Horrigan et al., 1999; Cui and Aldrich, 2000) (Fig. 7 A). The Po-V relation was then combined with the G-V relation at high voltages to generate the complete curve of Po versus voltage (Fig. 7 B). Fig. 7 B (left) shows that on the WT background, where the native N172 is neutral, mutation E399R has no effects on BK channel Po at any voltage range, indicating that Po is not sensitive to the charge type at 399. However, on the N172D background, mutations E399N and E399R shift the voltage dependence of Po toward more negative voltage ranges (Fig. 7 B, right), indicating that the electrostatic interaction between residues 172 and 399 alters voltage-dependent gating. Moreover, at extreme negative voltages (less than −80 mV), where Po is independent of VSD activation, both repulsive (N172D) and attractive (N172D/E399R) 172–399 interactions lower Po, whereas Po is similar to that of the WT channel when there is no 172–399 electrostatic interaction (N172D/E399N) (Fig. 7, B, right, and C). By fitting the Po-V relation to the HCA model (Eq. 2) (Horrigan et al., 1999), we obtained parameter L0 (Fig. 7 D). L0 is the open probability at 0 voltage without activation of the voltage sensor or Ca2+-binding sites, which reflects the intrinsic property of the activation gate. We fitted the Po-V relations to the HCA model with fixed parameters zL = 0.1 and zJ = 0.57 to obtain L0 (Fig. 7 D). The result shows that when there is no interaction between residues 172 and 399, L0 is at the same level (WT, E399R, and N172D/E399N). However, both repulsive (N172D) and attractive (N172D/E399R) interactions lower L0 (the difference is more than five times of the standard deviation of fitting), indicating that the intrinsic pore gate property is altered by the 172–399 electrostatic interaction.
native 99–374 interactions based on the following observations: (a) they are all part of the Mg$^{2+}$-binding site (Shi et al., 2002; Yang et al., 2008); (b) one partner of the interactions, D99 or N172, is located in the membrane-spanning domain, while the other, E374 or E399, is located in the cytosolic domain; and (c) electric charges on these residues are important for the interactions; opposite charges in either pair favor channel activation as compared with the like charges (Figs. 1 and 5).

Our results show that the interactions between residues 172 and 399 affect both voltage- and Ca$^{2+}$-dependent activation through allosteric mechanisms. Such functional effects probably result from conformational changes that are proximal to the Mg$^{2+}$-binding site (Fig. 6) and distant from the pore-gate domain (Fig. 7). These conformational changes could interfere with the allosteric mechanisms of channel gating so that the voltage- and Ca$^{2+}$-dependent activation is altered as a result.

**DISCUSSION**

In this paper, a native interaction that participates in channel activation was identified between two of the Mg$^{2+}$-binding coordination residues, D99 and E374. To study the D99–E374 interaction while avoiding complications caused by interactions of these residues with those outside of the Mg$^{2+}$-binding site, we introduced artificial electrostatic interactions between the other two Mg$^{2+}$-binding coordination residues, N172 and E399, through mutagenesis and chemical modifications. The engineered 172–399 interactions mimic the native 99–374 interactions based on the following observations: (a) they are all part of the Mg$^{2+}$-binding site (Shi et al., 2002; Yang et al., 2008); (b) one partner of the interactions, D99 or N172, is located in the membrane-spanning domain, while the other, E374 or E399, is located in the cytosolic domain; and (c) electric charges on these residues are important for the interactions; opposite charges in either pair favor channel activation as compared with the like charges (Figs. 1 and 5).

Our results show that the interactions between residues 172 and 399 affect both voltage- and Ca$^{2+}$-dependent activation through allosteric mechanisms. Such functional effects probably result from conformational changes that are proximal to the Mg$^{2+}$-binding site (Fig. 6) and distant from the pore-gate domain (Fig. 7).

Our current discovery of the noncovalent interactions between the interdomain Mg$^{2+}$-coordination residues may add new insights to the understanding of the allosteric mechanisms of BK channel gating. Our previous studies showed that Mg$^{2+}$ bound to BK channels...
interacts with R213 in the S4 segment to favor voltage-sensor activation, and this interaction is stronger when the channel is at the open state (Yang et al., 2007). Both of these properties contribute to Mg$^{2+}$-dependent activation of the channel. However, a recent study demonstrated that Mg$^{2+}$ binding at the closed state also contributes to channel activation (Chen et al., 2011). Because the native interdomain interaction between D99 and E374 affects channel activation in the absence of Mg$^{2+}$, it is possible that Mg$^{2+}$ binding can disturb such an interaction at either the closed or open state to facilitate channel activation.

These results may also help our current understanding of the allosteric Ca$^{2+}$-dependent activation of BK channels. Recent structural studies suggest that the cytosolic domain undergoes large conformational rearrangements upon Ca$^{2+}$ binding (Yuan et al., 2012). Such a large conformational change in the cytosolic domain can open the gate in the membrane-spanning domain by pulling the C-linker that covalently connects these two domains (Niu et al., 2004; Yuan et al., 2012). Our current study illustrates that the noncovalent interactions between residues D99–E374 and charges at 172–399 in the Mg$^{2+}$-binding site can also affect Ca$^{2+}$-dependent activation (Fig. 5). Because the interacting partners, D99–E374 or charges at 172–399, are located on the membrane-spanning and cytosolic domains of the BK channel, respectively, the interactions are likely to change the interfacial conformations between the two domains. Such an interfacial conformational change is suggested by the results that the 172–399 interaction disrupts the spontaneously formed disulfide bond that cross-links the two structural domains between C99 and C397 (Fig. 6). Interfacial conformations between different structural domains have also been shown previously to be important for activation of various ion channels (Gordon et al., 1997; Jiang et al., 2002; Bouzat et al., 2004; Niu et al., 2004; Unwin, 2005; Long et al., 2007; Kawate et al., 2009; Taraska et al., 2009).

The noncovalent interactions at the interface between the membrane-spanning and cytosolic domains of BK channels might be much more complicated than the electrostatic interactions that we have shown in this work. For instance, charges at residue 172 still can affect channel activation when its cytosolic electrostatic partner E399 was neutralized (Fig. 4 B), suggesting that charges at residue 172 might be within an interaction network with other residues in vicinity. Considering the narrow distance between the membrane-spanning domain and the cytosolic domain, as well as the 71-residue-long S0–S1 linker that might squeeze into the interface, the interdomain noncovalent interactions, including electrostatic interactions, hydrophobic interactions, and hydrogen bonds, might tightly regulate the dynamic coupling of different structural domains to control BK channel gating. Our current study using electrostatic interaction as a tool to probe the interdomain interface will pave the way to further understand the domain–domain interactions and their effects on the allosteric gating mechanism of BK channels.

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Yang et al. 227


