Hydrophobic interaction between contiguous residues in the S6 transmembrane segment acts as a stimuli integration node in the BK channel

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Large-conductance Ca\(^{2+}\)- and voltage-activated K\(^{+}\) channel (BK) open probability is enhanced by depolarization, increasing Ca\(^{2+}\) concentration, or both. These stimuli activate modular voltage and Ca\(^{2+}\) sensors that are allosterically coupled to channel gating. Here, we report a point mutation of a phenylalanine (F380A) in the S6 transmembrane helix that, in the absence of internal Ca\(^{2+}\), profoundly hinders channel opening while showing only minor effects on the voltage sensor active–resting equilibrium. Interpretation of these results using an allosteric model suggests that the F380A mutation greatly increases the free energy difference between open and closed states and uncouples Ca\(^{2+}\) binding from voltage sensor activation and voltage sensor activation from channel opening. However, the presence of a bulky and more hydrophobic amino acid in the F380 position (F380W) increases the intrinsic open–closed equilibrium, weakening the coupling between both sensors with the pore domain. Based on these functional experiments and molecular dynamics simulations, we propose that F380 interacts with another S6 hydrophobic residue (L377) in contiguous subunits. This pair forms a hydrophobic ring important in determining the open–closed equilibrium and, like an integration node, participates in the communication between sensors and between the sensors and pore. Moreover, because of its effects on open probabilities, the F380A mutant can be used for detailed voltage sensor experiments in the presence of permeant cations.

INTRODUCTION

Large-conductance Ca\(^{2+}\)- and voltage-activated K\(^{+}\) channels (BK, Slo1) increase their open probability in the presence of membrane depolarization and/or during an increase in the intracellular Ca\(^{2+}\) concentration (Marty, 1981; Pallotta et al., 1981; Latorre et al., 1982). The BK channel is one of the most broadly expressed channels in mammals and plays important roles in both excitable and nonexcitable cells. For example, in vascular smooth muscle cells it regulates the contractile tone, in neurons and nonexcitable cells. For example, in vascular smooth muscle cells it regulates the contractile tone, in neurons and nonexcitable cells it colocalizes with voltage-dependent calcium channels (B Crest, 1993; Edgerton and Reinhart, 2003; Gessner et al., 2005) and is involved in the control of neurosecretion, and in hair cells it is involved in frequency tuning (Lancaster and Nicoll, 1987; Brayden and Nelson, 1992; Gola and Crest, 1993; Edgerton and Reinhart, 2003; Gessner et al., 2005; Miranda-Rottmann et al., 2010). BK channels are tetramers in which the pore-forming α subunit is coded by a single gene (Slo1; KCNMA1) ubiquitously expressed across mammalian tissues (Toro et al., 1998). However, BK channels display a variety of phenotypes in different cells and tissues as a result of alternative splicing, metabolic regulation and modulation by β (Orio et al., 2002) and γ (Yan and Aldrich, 2012) subunits. This diversity of phenotypes is fundamental for their adequate function in each tissue.

The voltage-sensing domain (VSD) and pore-gating domain (PGD) of BK channels share similarity with voltage-dependent K\(^{+}\) channels (Kv). However, an evolutionary relationship between them seems distant (Yu and Catterall, 2004). Evidence suggests that pore domains in both channels are different. For instance, the inner pore in BK channels is larger (Brelidze et al., 2003; Li and Aldrich, 2004; Zhou et al., 2011) than in Kv channels. Also, the ion permeation gate of BK channels resides at the selectivity filter (Piskorowski and Aldrich, 2006), whereas that of Kv channels is controlled by an intracellular gate (Liu et al., 1997). Moreover, Zhou et al. (2011) found that BK S6 transmembrane helix residues are rotated relative to Shaker, which is a Kv channel.
Mutations in the S6 affect BK channel gating. For example, mutations on human BK F380 residue (Lippiat et al., 2000) or its corresponding murine BK-F315 (Wang and Brenner, 2006) have profound effects on gating and unitary conductance. There is evidence that this residue participates in a state-dependent interaction with L312 (L377 in human BK) of a neighboring subunit to hold the channel in the closed configuration (Wu et al., 2009), whereas the neighboring residue M314 (M377 in hSlo) changes its degree of pore exposure during opening (Chen and Aldrich, 2011). Interestingly, Wu et al. (2009) found that mBK F315 mutants show changes in Ca$^{2+}$ and voltage sensitivity, suggesting that this residue participates in the allosteric communication between voltage sensors and the pore gate domain. However, it is difficult to decide which parameters are being affected by the mutations on the basis of macroscopic currents alone. In particular, the mBK mutant (F315A) is able to reach the membrane, but the very small currents detected suggest that either function or expression is impaired in the mutant channel (Wu et al., 2009). Here, we found that the corresponding human mutation (hSlo-F380A), given the large gating currents we detected, is robustly expressed in the oocyte plasma membrane, indicating that expression and voltage sensor movements were not greatly affected. However, ionic currents noticeably changed. In the presence of 100 µM internal Ca$^{2+}$, a large rightward shift of the voltage activation curve was observed, with a reduction in the apparent voltage sensitivity. In the absence of Ca$^{2+}$, small ionic currents of BK F380A were detected only at voltages >300 mV. Therefore, F380A mutant gating currents can be observed in solutions containing 110 mM K$^+$ without interference of ionic currents in a broad range of voltages. Based on the Horrigan and Aldrich (HA) allosteric model for BK channel gating (Horrigan and Aldrich, 2002), these results can be explained by a large increase in the free energy difference between open and closed states and a decrease of the allosteric factors that couple the Ca$^{2+}$ binding with the voltage sensor and the voltage sensor with the pore opening. The same behavior was observed when the 380 residue was mutated to other small hydrophobic amino acids. The behavior of BK F380A makes it an excellent candidate for further understanding BK voltage-dependent activation as gating currents can be measured in the presence of K$^+$. To test the effects of increasing the hydrophobicity of the 380 residue, we studied the effects of the F380W mutation on BK channel gating. The open–closed equilibrium was increased and the coupling between Ca$^{2+}$ and voltage sensors with the pore domain was severely weakened in the F380W mutant. Our results indicate that the 380 residue participates in a hydrophobic interaction with L377 that stabilizes the open state while functioning as a link for the communication between Ca$^{2+}$ and voltage sensors and the pore domain.

**Materials and Methods**

**Site-directed mutagenesis and channel expression**

Human BK α subunit (GenBank accession no. U11058) was provided by L. Toro (University of California, Los Angeles, Los Angeles, CA). Full-length cDNA for hSlo1 was subcloned into pBusta. Mutations were obtained by the QuikChange site-directed mutagenesis kit (Agilent Technologies). The constructs were checked by sequencing. To prepare mRNA, the plasmids were linearized with Enzyme NotI and then in vitro transcribed by mMESSAGE mMACHINE (Ambion). *Xenopus laevis* oocytes were injected with ~50 ng of a subunit mRNA 4–8 d before recording.

**Electrophysiology**

Using the patch-clamp technique, ionic currents ($I_\alpha$) were recorded in inside-out and outside-out configurations, and gating currents ($I_g$) were recorded in the inside-out configuration only. All of the internal solutions contained (mM) 110 X-MeSO$_4$ and 10 HEPES. 5 mM EGTA was used as Ca$^{2+}$ chelator for “zero Ca$^{2+}$” solutions. These solutions contained an estimated 0.8 nM, based on the presence of ~10 µM contaminant [Ca$^{2+}$] (Cox et al., 1997), 1 mM HEDTA was used as Ca$^{2+}$ buffer for solutions containing 0.1–3 µM free Ca$^{2+}$. No Ca$^{2+}$ chelator was used in 100 µM free Ca$^{2+}$ solutions. For 0.1–3 µM free Ca$^{2+}$ solutions, CaCl$_2$ was added to reach the desired [Ca$^{2+}$] from a stock solution containing X mM CaCl$_2$, and HCl was added until reaching 4 mM [Cl$^{-}$]. The external (pipette) solution contained (mM) 110 X-MeSO$_4$, 10 HEPES, and 2 MgCl$_2$. $I_\alpha$ currents and unitary channel recordings were recorded using internal and external K$^+$ instead of X.

**Nonstationary noise analysis**

To determine the unitary conductance of the channels at 100 and −100 mV, we used a nonstationary fluctuation analysis (Sigworth, 1980; Alvarez et al., 2002). The mean current and variance of isochrones were calculated from 200 current relaxation traces in response to a 100-mV pulse and a −100-mV tail pulse. The unitary current was calculated by fitting the variance versus mean current data to a straight line.

**Limiting slope analysis**

In the case of the F380A mutant, it was possible to obtain the limiting slope of the $P/O(V)$ curve at 0 internal Ca$^{2+}$. In this case the number of channels, $N$, was obtained by dividing the maximum ON gating charge by the charge per channel, which was supposed to be 2.4 $e_0$ (4 $z_0$ see below). Because in the case of the F380A mutant we can measure gating currents and macroscopic ionic currents ($I$) simultaneously, the probability of opening $P_O$ can be calculated as $P_O = I/N$, where $i$ is the single-channel current. From the noise analysis we have $i$ at −100 mV ($I$=100mV); however,
the BK \( i(V) \) curve is nonlinear. Because we cannot determine the \( I(V) \) relation for the F380A single channel, this nonlinearity was taken into account by multiplying \( I - 100 \text{mV} \) by the ratio of the macroscopic conductances obtained at the end of the activation pulse and at the beginning of the tail current for the mutant channel at 100 \( \mu \text{M Ca}^{2+} \) (see Fig. 1 B) for the same voltages at which the \( P_0 \) was calculated in Fig. 3 B.

Data analysis

Data analysis was performed using Clampfit 10 (Axon Instruments) and Excel 2007 (Microsoft). OFF gating currents were integrated between 0 and 400 \( \mu \text{s} \) after the OFF voltage step to obtain the OFF net charge movement (\( Q_{\text{OFF}} \)). To determine the charge displaced between closed states, \( Q \), the first 50 \( \mu \text{s} \) of the ON gating currents were fitted to a single exponential function and the area under the curve was integrated, as described in Fig. S1. In all cases, leakage current was subtracted by shifting the current by its mean value during 1 ms before the voltage step. \( Q_{\text{OFF}}(V) \) and \( Q(V) \) data were fitted using a Boltzmann function:

\[
Q(V) = \frac{Q_{\text{MAX}}}{1 + e^{-zQ(\frac{V - V_0}{RT})}},
\]

where \( Q_{\text{MAX}} \) is the maximum charge, \( z_Q \) is the voltage dependency of activation, \( V_0 \) is the half-activation voltage of gating current, \( T \) is the absolute temperature (typically 295 K), \( R \) is the universal gas constant. \( Q_{\text{MAX}}, \ V_0, \) and \( z_Q \) were determined by using the Solver complement of Microsoft Excel to find the minimum of the sum of the squares of the differences between experimental and calculated charges. Instantaneous tail currents were fitted to a Boltzmann function of the form:

\[
G(V) = \frac{G_{\text{MAX}}}{1 + e^{-zG(\frac{V - V_0}{RT})}},
\]

where \( G_{\text{MAX}} \) is the maximum tail current, \( z_G \) is the voltage dependency of activation, and \( V_0 \) is the half-activation voltage of the ionic current. All the analyses that follow were performed using \( G/G_{\text{MAX}} \) or \( Q/Q_{\text{MAX}} \) values.

To analyze \( G(V) \) and \( Q(V) \) curves, a mean \( V \) and \( V_a \) (\( <V_a> \)) were obtained at each \( \text{Ca}^{2+} \) concentration. All of the corresponding \( G(V) \) and \( Q(V) \) curves were aligned by shifting them along the voltage axis by the mean \( \Delta V_0 \) (\( <V_a> - V_0 \)) or \( \Delta V_0 \) (\( <V_a> - V_0 \)) to get a mean curve that did not alter the voltage dependence. For each mean curve, the data were binned in 25-mV intervals.

Model fitting

We fit the experimental data using the HA model (Horrigan and Aldrich, 2002). In this model the opening reaction is described by the voltage-dependent equilibrium constant \( L \). \( J \) is the equilibrium constant that defines the voltage sensor resting-active equilibrium (\( J_f \)), whereas the \( \text{Ca}^{2+} \) binding is described by the equilibrium constant \( K \). The model includes allosteric factors between any pair of modules: \( C, D, \) and \( E \) for the coupling between calcium sensors and pore opening, the coupling of the voltage sensors and pore opening, and the calcium sensor–voltage sensor allosteric coupling, respectively (see Scheme 1 below).

\[
P_0 = \frac{L(1+J+K+JKE)}{L(1+J+K+JKCDE)}
\]

where

\[
J = J_e^f \frac{e^{\gamma Jf(V - V_a)}}{RT},
\]

\[
K = \frac{[\text{Ca}^{2+}]}{k_d},
\]

and

\[
L = L_e^f \frac{e^{\gamma Lf(V - V_a)}}{RT}.
\]

\( f \) is the resting–active equilibrium for the voltage sensor at 0 mV with no calcium bound. \( L_0 \) is the equilibrium constant for channel opening with all voltage sensors in the resting state and not calcium bound at 0 mV. The parameter \( k_d \) is the dissociation constant of a single calcium-binding site with all voltage sensors at rest and the channel closed. The parameters \( z_L \) and \( z_J \) are the voltage dependence of the equilibria \( L \) and \( J \), respectively.

For WT BK channel, \( L_0 \) and \( z_L \) were obtained from Orio and Latorre (2005). The equilibrium constant that defines the voltage sensor active–resting equilibrium at zero voltage, \( f_0 \), and the apparent number of gating charges per voltage sensor, \( z_Q \), in the absence of \( \text{Ca}^{2+} \) were obtained from the data shown in Fig. 3 E because

\[
Q(V) = \frac{Q_{\text{MAX}}}{1 + e^{\gamma Q(V - V_0)/(RT)}},
\]

where

\[
J_e^f = \frac{e^{\gamma Jf(V - V_0)}}{RT},
\]

and \( \gamma \) is the half-activation voltage of the \( Q(V) \) curve. Next, we calculated the allosteric parameter \( E \) assuming that the fast ON gating currents, \( Q \), in the presence of saturating calcium occurs essentially between closed states; in this case \( f \) is multiplied by \( E \). So, \( E \) can be calculated using the equation:

\[
J_E = \frac{e^{\gamma Jf(V - V_0)/(RT)}}{RT},
\]

where \( z_Q \) and \( V_0 \) are the voltage dependence and the half-activation voltage of the \( Q(V) \) curve at 100 \( \mu \text{M Ca}^{2+} \). The values obtained for \( L_0, z_L, j_0, j_e, k_d \) and \( E \) were introduced into the model as fixed parameters, and we calculated the allosteric parameter \( D \) by minimizing the sum of square of \( G(V) \) data in the absence of internal \( \text{Ca}^{2+} \) with the Solver tool of Excel. Once \( D \) was obtained, we performed a global fit to the \( G(V) \) data obtained at seven different \( \text{Ca}^{2+} \) concentrations to estimate \( C \) and \( k_d \).

To fit the F380A data, we fixed \( z_L \) from Orio and Latorre (2005). \( E, j_0 \), and \( z_Q \) were calculated using the data for the mutant channel shown in Fig. 3 E in the same manner as we did for the WT channel. \( k_d \) was assumed to be identical to WT. Finally, to obtain \( L_0, D, \) and \( C \), the \( G(V) \) data obtained at different \( \text{Ca}^{2+} \) concentrations was globally fitted to the HA model. \( L_0 \) and \( D \) were fitted without any restriction, except the condition that \( L_0D' = 6.59 \times 10^{-3} \) (see Fig. 2 B).
The F380W data were fitted to the HA model using the same \( z_1 \) as before, and \( k_f \) and \( z_2 \) were obtained from the fit to the \( Q(V) \) data shown in Fig. 4 C in a similar way as was done for WT BK. \( k_0 \) was supposed as identical to WT. Previous parameters obtained were introduced into the model, and we calculated the allosteric parameters \( L_C \) and \( D_{by} \) minimizing sum of the square of \( G(V) \) data, in the absence of internal Ca\(^{2+}\). Finally, \( E \) and \( C \) were obtained by performing a global fit to the \( G(V) \) data obtained at different Ca\(^{2+}\) concentrations using the HA model.

To test whether a fit parameter was statistically different between two datasets, a global fit in which the parameter of interest was restricted to be equal among all datasets was compared against an independent fit of each dataset. A comparison between independent and global fits was done by an extra sum of squares \( F \) test, using the formula

\[
F = \frac{(SS_{gb} - SS_{ind}) / (DF_{gb} - DF_{ind})}{SS_{ind} / DF_{ind}},
\]

where \( SS \) and \( DF \) are the sum of squares and the degrees of freedom of the fit, respectively; \( gb \) and \( ind \) denote the global and the independent fit, respectively. All of the analyses were performed with Excel, and the conversion of the \( F \) values to \( p \) values was performed with Prism software (GraphPad Software). \( p \) values <0.01 were considered significant.

**Molecular modeling**

BK 3-D model was built using the sequence of human BK channel (UniProt ID: Q12791) stored in GenPept. Based on the sequence alignment of Carvacho et al. (2008), the segments SS–S6 were aligned against the sequence of MthK crystal in Protein Data Bank (accession no. 4HYO; Posson et al., 2013), achieving 35% of sequence identity. Several models were obtained using Modeller v9.10 (Sali and Blundell, 1994), from which the lowest energy structure based on DOPE potential was chosen to build a tetrameric model by structural aligning with MthK crystal structure. The resulting model was then evaluated using the ANOLEA software (Melo and Feytmans, 1998).

The BK pore structure was embedded on a lipid membrane of phosphatidyl oleoyl phosphatidylcholine (POPC) hydrated with TIP3P water and neutralized by means of counteions at 0.110 M KCl. Also, potassium ions were located on sites S0, S2, and S4 of the selectivity filter as used by Moscoso et al. (2012). The final dimensions of the system were \( 12.5 \times 12.5 \times 12.0 \) nm\(^3\) and it was constituted by \( \approx \) 370,000 atoms. Next, the system was submitted to a molecular dynamics (MD) protocol. In brief, a first stage of 2,000 steps of energy minimization and 10 ns of equilibration was performed using an NPT ensemble at 298 K and 1 atm. Also, harmonic restraints of 0.5 kcal/mol\(\cdot\)Å\(^2\) on C\(\alpha\)s were applied to preserve the secondary structure of the model. F380A and F380W BK mutant systems were generated from the equilibrated BK WT system using the VMD mutator plugin (Humphrey et al., 1996). After equilibration, several production simulations were performed for each system, which altogether lasted \( \approx \) 40 ns. On the production phase, harmonic restraints of 0.2 kcal/mol\(\cdot\)Å\(^2\) were defined for C\(\alpha\)s of segment S5, allowing freedom of segment S6 for further analysis.

All MD simulations were performed with the software NAMD 2.9 (Phillips et al., 2005) using the CHARMM27 (Patel et al., 2004) and CHARMM36 (Klauda et al., 2010) force fields. Bonded and short- and long-range nonbonded interactions were integrated with a time step of 2, 2, and 4 fs, respectively. An 8-Å spherical cutoff was used for short-range nonbonded interactions, including a switching function from 7 Å for the van der Waals term and shifted electrostatics (Wells et al., 2012). The particle-mesh Ewald (PME) method was used with a grid spacing of \( \approx 1 \) Å.

**Online supplemental material**

This supplemental material contains one figure and its legend. Fig. S1 shows two examples of the way we determined the fast component of the ON gating current. Online supplemental material is available at http://www.jgp.org/cgi/content/full/jgp.201411194/DC1.

**RESULTS**

Mutations in the F380 residue produce large changes in conductance-voltage relationships

Macroscopic ionic currents for WT BK and the F380 mutants F380A and F380W in the nominal absence of internal Ca\(^{2+}\) and in the presence of several different Ca\(^{2+}\) concentrations were characterized in inside-out patches. Representative current records at nominally 0 and 100 µM internal Ca\(^{2+}\) concentrations are shown in Fig. 1 (A–C). At 100 µM internal Ca\(^{2+}\), the two BK high-affinity Ca\(^{2+}\)-binding sites were saturated, whereas at 0.8 nM (nominal “0” Ca\(^{2+}\)) the Ca\(^{2+}\)-binding sites were empty (Cox et al., 1997). Fig. 1 B shows that robust F380A currents can be recorded in the presence of several internal Ca\(^{2+}\) concentrations, but their behavior is quite different from the currents induced by the WT BK channel (Fig. 1 A). The faster kinetics and the lack of inward currents elicited by the test pulse both indicate that the voltages required to open the mutant channel are much larger compared with those required to activate WT BK channels. In fact, Fig. 1 E shows that although internal Ca\(^{2+}\) is able to promote a leftward shift in the F380A voltage-activation curve \( (G(V)) \), considerably larger voltages compared with the WT BK channel are needed to activate the F380A mutant channel (Fig. 1 D). This mutation also produces a significant decrease in apparent voltage dependence compared with the WT BK (Fig. 1, compare D and E). In the absence of intracellular calcium, the mutant F380A \( G(V) \) could not be measured, probably because it is right-shifted to positive voltages that were too large to be experimentally unreachable.

To understand the effects of substituting F380 with other hydrophobic amino acids, we tested mutants F380W, F380L, and F380I. Interestingly, introducing a larger and more hydrophobic amino acid in position 380 (F380W) produces a leftward shift of the \( G(V) \) in the absence of Ca\(^{2+}\) and at saturating Ca\(^{2+}\) concentrations (Fig. 1, C and F) compared with the WT BK. Surprisingly, at the two higher internal Ca\(^{2+}\) concentrations tested, a large fraction of channels remained open even at voltages so negative as -200 mV. In contrast, we found that at 100 µM Ca\(^{2+}\), the gating of mutants F380L and F380I was very similar to that of F380A (Figs. 1, G–I).
F380A decreases single-channel conductance and modifies gating kinetics without effects on the voltage dependence of the deactivation kinetics

We tested whether the F380A channel was a bona fide BK channel by analyzing its sensitivity to iberiotoxin (IbTx), known to be a potent and specific blocker of BK channels (Galvez et al., 1990; Candia et al., 1992; Giangiacomo et al., 1992; Kaczorowski and García, 1999). The effect of the toxin was checked in outside-out membrane patches. In the presence of 100 µM internal Ca²⁺, the current elicited by a 150-mV pulse was measured at different times in the presence of 100 nM IbTx in bath solution (Fig. 2A). In the presence of the toxin, currents decreased with time and were completely abolished after 5 min (t50). We also attempted to detect F380A mutant currents in the absence of internal Ca²⁺. In this case, the test pulse was 350 mV and robust gating currents and relative small ionic currents were observed (Fig. 2B). These ionic currents were fully blocked by IbTx after 5-min incubation. We concluded that these currents also are mutant BK channels instead of currents induced by oocyte endogenous channels.

Although it is possible to record unitary channels of WT BK and F380W in the absence of internal Ca²⁺, because of the very fast deactivation kinetics and the very low open probability at the voltages that unitary channels are typically recorded, we were unable to observe individual F380A channel openings (Fig. 2C). Therefore, we determined unitary conductance by using noise analysis (Sigworth, 1980; Álvarez et al., 2002). Unitary conductance of the mutants F380A and F380W (96 ± 6 pS and 106 ± 3 pS, respectively; mean ± SEM) was reduced compared with that of the WT channel (222 ± 20 pS; Fig. 2, D–F).

![Figure 1](image-url)  
Figure 1. Characterization of BK F380 mutant ionic currents. (A) BK-WT currents (Iv) evoked by voltage steps (−100 to 300 mV; in 10-mV steps of 10 ms) at two different [Ca²⁺], (0 and 100 µM). (B) BK-F380A macroscopic currents recorded at 0 and 100 µM ([−100 to 350 mV; in 10-mV steps of 2 ms]). Note the difference in time scale because of the fast activation in the case of BK-F380A. (C) BK-F380W currents (Id) evoked by voltage steps (−220 to 300 mV; in 10-mV steps of 10 ms) in two different [Ca²⁺], (0 and 100 µM). (D–F) Relative conductance versus voltage (G/GMAX vs. V) curves were obtained from tail current measurements from experiments as those shown in A–C. Data were fitted using a Boltzmann function (solid lines). For the BK-WT channel, parameters were (mean ± SEM) V10 = 176 ± 8 mV, zG = 0.96 ± 0.08 at [Ca²⁺] = 0 µM, n = 5; V5 = 120 ± 10 mV, zG = 1.30 ± 0.10 at [Ca²⁺] = 0.1 µM, n = 5; V5 = 110 ± 8 mV, zG = 1.40 ± 0.10 at [Ca²⁺] = 0.2 µM, n = 6; V5 = 81 ± 5 mV, zG = 1.33 ± 0.12 at [Ca²⁺] = 0.5 µM, n = 5; V5 = 52 ± 8 mV, zG = 1.40 ± 0.06 at [Ca²⁺] = 1.5 µM, n = 5; V5 = 20 ± 6 mV, zG = 1.34 ± 0.11 at [Ca²⁺] = 3 µM, n = 6; V5 = −43.7 ± 11 mV, zG = 1.12 ± 0.04 at [Ca²⁺] = 100 µM, n = 5. BK-F380A: V5 = 620 ± 34 mV, zG = 0.32 ± 0.03 at [Ca²⁺] = 0.2 µM, n = 4; V5 = 492 ± 19 mV, zG = 0.43 ± 0.02 at [Ca²⁺] = 0.5 µM, n = 4; V5 = 349.8 ± 11.7 mV, zG = 0.42 ± 0.01 at [Ca²⁺] = 1.5 µM, n = 5; V5 = 321 ± 9 mV, zG = 0.45 ± 0.01 at [Ca²⁺] = 3 µM, n = 4; V5 = 292 ± 25 mV, zG = 0.48 ± 0.05 at [Ca²⁺] = 100 µM, n = 5; BK-F380W: V5 = 133 ± 3 mV, zG = 0.73 ± 0.03 at [Ca²⁺] = 0 µM, n = 17; V5 = 102 ± 6 mV, zG = 0.52 ± 0.02 at [Ca²⁺] = 0.1 µM, n = 6; V5 = 92 ± 7 mV, zG = 0.52 ± 0.01 at [Ca²⁺] = 0.2 µM, n = 7; V5 = 80 ± 5 mV, zG = 0.55 ± 0.03 at [Ca²⁺] = 0.5 µM, n = 7; V5 = 26 ± 5 mV, zG = 0.70 ± 0.01 at [Ca²⁺] = 3 µM, n = 7; V5 = −58 ± 3 mV, zG = 0.52 ± 0.03 at [Ca²⁺] = 100 µM, n = 14. (G and H) BK-F380I and BK-F380L currents (Id) evoked by voltage steps (−100 to 350 mV; in 10-mV steps of 2 ms) at [Ca²⁺] = 100 µM, respectively. (I) Relative conductance versus voltage (G/GMAX vs. V) curves were obtained from tail current measurements from the experiments shown in G and H. Data were fitted using a Boltzmann function (solid lines). For the BK-F380I channel, parameters were (mean ± SEM) V5 = −261 ± 6 mV, zG = 0.72 ± 0.08 at [Ca²⁺] = 0 µM, n = 100 µM, n = 4. BK-F380L: V5 = 267 ± 6 mV, zG = 0.58 ± 0.04 at [Ca²⁺] = 100 µM, n = 7. Black and red lines are the BK-F380I and BK-F380L curves and Boltzmann fitting of the BK channel WT and F380A mutant at [Ca²⁺] = 100 µM (as is seen in D and E), respectively.
A detailed study of the deactivation kinetics in a wide range of voltages indicates that, as expected from the $G(V)$ data, the whole $\tau(V)$ curve for the F380A mutant shifts to the right along the voltage axis (Fig. 2 G). At very negative voltages, only the backward closing rate constants contribute to $\tau(V)$, and the results in Fig. 2 G show that the closing rate of the F380A mutant becomes about fivefold faster with little effect on its voltage dependence compared with that of the WT channel. Note in Fig. 2 G that very fast deactivation kinetics of F380A would produce very brief openings at the single channel level, which could be difficult to record.

**Characterization of BK F380A mutant gating currents in the absence and presence of Ca$^{2+}$**

We were able to detect gating currents ($I_g$) of the F380A mutant in the presence (Fig. 3 A, inset) and absence (Fig. 3 D) of K$^+$ by using the inside-out patch-clamp configuration. Unlike WT BK, F380A gating currents can be measured in the presence of symmetrical $110 \text{ mM K}^+$ with minimal ionic current contamination (Fig. 3 A, inset). Note that the $Q(V)$ curves for F380A in the presence and absence of permeant cations are almost identical (Fig. 3 A).

Unfortunately, as discussed above, it was not possible to observe individual openings of F380A mutant, making it impossible to measure the probability of opening at very negative voltages (limiting slope determination) to obtain a direct estimate of the equilibrium constant that governs the closed–open equilibrium, $L$. Instead, thanks to the large rightward shift of the $G(V)$ promoted by the F380A mutant, we were able to calculate the limiting slope at voltages at which all voltages sensors are activated (Fig. 3 A). We used recordings as in Fig. 3 A (inset) to calculate the open probability at voltages from 300 mV (Fig. 3 B). To calculate the open probability, we estimated the total number of channels from the quotient between $Q_{\text{MAX}}$ in the patch and the charge per channel (estimated to be $4Z = 2.4e_0$ from the slope of the $Q(V)$). Once obtained, the $P_o(V)$ curve (Fig. 3 B), we can determine the voltage dependence of the intrinsic open–closed equilibrium, $\gamma$, and the equilibrium constant to 0 mV, $L_0^\ast$, by using the HA allosteric model, which predicts...
that under these experimental conditions the open probability is

\[ P_O = LD^4 / (1 + LD^4) \]  \hspace{1cm} (2)

Therefore, \( I_{O^*} = I_O D^4 \) obtained from extrapolation of the \( P_O(V) \) data of Fig. 3 B to 0 mV represents the product between the intrinsic equilibrium constant, \( I_O \), and the allosteric factor describing the interaction between channels opening and voltage sensor activation, \( D \), to the fourth power. The value of \( I_O D^4 = 6.6 \times 10^{-5} \) and \( z_L = 0.42 \) obtained from the fitting to the \( P_O(V) \) data show in Fig. 2 B to Eq. 2 was used as a further constraint when performing the global fit of the \( G(V) \) data of Fig. 6 A using the HA model. The value \( z_L \) found using this method agrees reasonably with the value obtained by Orio and Latorre (2005).

Robust gating currents for both WT and mutant channels can be recorded in the absence and presence of 100 µM internal Ca\(^{2+}\) (Fig. 3, C and D). In both cases, because of the very fast opening kinetics, to obtain only the gating current displaced between closed states, the gating charge was calculated by adjusting the first 50 µs of the ON gating currents to a single exponential and integrating the area under the curve. Compared with WT BK channels, F380A shifts the gating charge-voltage (\( Q_c(V) \)) curve along the voltage axis to the left by 20 mV in the absence of calcium with no appreciable changes in the voltage dependency of activation \( z_Q \) (Fig. 3 E). The values of the half-activation voltage of gating currents in the absence of internal Ca\(^{2+}\), \( V_0 \), and \( z_Q \) for the WT channel are in reasonable agreement with those previously reported (Horrigan and Aldrich, 1999, 2002; Bao and Cox, 2005; Contreras et al., 2012). Increasing intracellular Ca\(^{2+}\) promotes a leftward shift of the \( Q_c(V) \) curve for both the WT BK and the F380A mutant, accompanied by an increase in \( z_Q \). In the presence of 100 µM Ca\(^{2+}\), the half-activation voltage for the WT BK and the F380A mutant were 29 mV and 72 mV, respectively (Fig. 3 E). This result suggests that in the F380A mutant the interaction between voltage sensor activation and calcium binding has been weakened. Our results widely differ from that of Horrigan and Aldrich (2002) because they did not observe as large of changes in the half-activation voltage of the \( Q_c(V) \) in the presence of internal 70 µM Ca\(^{2+}\).

We note here that the half-activation voltage for the \( G(V) \) and \( Q_c(V) \) curves for the WT BK channel in 100 µM internal Ca\(^{2+}\) are \(-44 \) mV and 29 mV, respectively. These results indicate that when, on average, half of the channels are open, only 14% of the total gating charge has been displaced. Therefore, in the presence of saturating internal Ca\(^{2+}\), opening can proceed from deep closed states. Actually, when almost all BK channels are open, only \( \sim 50\% \) of the total voltage sensors have been activated.

Characterization of BK and F380 mutants gating currents in the absence of Ca\(^{2+}\).

To study the effects of other F380 mutations on voltage sensor activation, families of \( I_o \) as those shown in Fig. 4 (A–C, insets), evoked at different voltages (−90 to 350 mV), were determined for the fast ON gating current to
obtain the gating charge activation ($Q(V)$) relationships. $Q(V)$ data were fitted using Boltzmann functions, normalized to their maxima, and averaged to yield the results shown in Fig. 4 (A–C). Mutations F380l and F380L did not change voltage sensor activation significantly (Fig. 4 D). These results suggest that the F380 mutations to I or L do not affect the voltage sensor conformational changes that accompany the changes in applied membrane voltage but profoundly modify the channel gating uncoupling voltage sensor activation and pore and/or modifying the intrinsic closed–open equilibrium. In contrast, a large hydrophobic amino acid such as tryptophan slightly facilitates opening (Fig. 1, C and F) without greatly affecting the voltage sensor workings (Fig. 4 C). Thus, this suggests that the presence a strong hydrophobic amino acid in position 380 is needed for the adequate functioning of the channel.

F380A diminishes slow gating charge recovery in the absence of intracellular Ca$^{2+}$

A two-state model (i.e., resting–active) is adequate to describe the early movement of the voltage sensor in the BK (Horrigan and Aldrich, 1999). Late conformational changes of the voltage sensor produce a slow component of the gating charge that can be measured in the OFF gating current. To test the existence of a slow component, we searched for increases in the OFF net gating charge, $\Delta Q_{off}$, after voltage pulses of increasing duration were applied, as shown in Fig. 5 A. The origin of this slow component can be explained in terms of an allosteric voltage-gating scheme (Horrigan and Aldrich, 2002). This model predicts that the slow OFF gating charge develops with a time constant that is limited by the speed of channel opening (Horrigan and Aldrich, 1999). The slow component is quite reduced in the mutant compared with WT channel (Fig. 5, B and C). This result suggests that although the opening of WT occurs mainly when three or four voltage sensors are active, necessarily the opening of mutant channel occurs when all voltage sensors are active because of the weak voltage dependency of pore opening. We argue that the slow component is simply a consequence of the number of channels that were open at the end of a long duration voltage pulse (Horrigan and Aldrich, 2002). For example, in the case of the 200-mV pulse, most WT BK channel can reach the open configuration (Fig. 5 D). In this case, the OFF gating charge develops with a time constant close to the time constant that describes the activation of the ionic currents (Fig. 5 B). However, in the case of the F380A channel, the dramatic decrease in the slow component (Fig. 5 C) is simply a reflection of the very low probability of channel opening at 200 mV, and most of the gating charge is displaced between closed states (Fig. 5 E).

F380A mutation affects the closed–open equilibrium and coupling of voltage sensor activation to channel opening

To determine how the F380A and F380W mutations affect the phenotype of the BK channel, we fit our results to the HA allosteric model for WT BK and both mutants (Fig. 6, A–C). First, we fit the data of the WT BK using as initial values for the parameters of the model those reported by Orio and Latorre (2005), but using the $zJ$ values calculated from fitting the $Qc(V)$ curves to a Boltzmann function in the absence of intracellular calcium (Fig. 3 E). These results suggest that the F380 mutations to I or L do not affect the voltage sensor conformations that accompany the changes in applied membrane voltage but profoundly modify the channel gating uncoupling voltage sensor activation and pore and/or modifying the intrinsic closed–open equilibrium. In contrast, a large hydrophobic amino acid such as tryptophan slightly facilitates opening (Fig. 1, C and F) without greatly affecting the voltage sensor workings (Fig. 4 C). Thus, this suggests that the presence a strong hydrophobic amino acid in position 380 is needed for the adequate functioning of the channel.

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![Figure 4](https://example.com/figure4.png)

**Figure 4.** Gating currents of BK-F380 mutants in internal calcium absence. (A–C, insets) $I_g$ of mutants (F380I, F380L, and F380W) was recorded in inside-out macropatches at 0 µM [Ca$^{2+}$]. $I_g$ was evoked by 2-ms voltage steps of 10 mV going from −90 to 350 mV. (A–C) $Qc(V)$ relationships of BK and BK-F380 mutants were obtained by integrating each $I_g$ trace. Boltzmann fitting to the experimental data (mean ± SEM) is indicated by solid lines ($V_0 = 182 ± 9$ mV, $zQ = 0.56 ± 0.01 e_0$ in BK-F380I, n = 5; $V_0 = 173 ± 4$ mV, $zQ = 0.55 ± 0.01 e_0$ in BK-F380L, n = 7; $V_0 = 153 ± 6$ mV, $zQ = 0.56 ± 0.02 e_0$ in BK-F380W, n = 4). Dashed lines represent the Boltzmann function for the WT $Qc(V)$ obtained in the absence of intracellular Ca$^{2+}$ (Fig. 3 E). (D) $V_0$ of $Qc(V)$ of mutants (F380I, F380L, and F380W) compared with $Qc(V)$ of WT and F380A. Mean ± SEM.
described in Materials and methods using the results of Fig. 3 E, and $z_L$ and $L_0$ were obtained from Orio and Latorre (2005). The final values of the different parameters were obtained as described in Materials and methods. The parameters obtained for WT were in good agreement with those previously reported except for $C$ and $E$, which were smaller and greater, respectively. The $E$ value was surprisingly large compared with those reported in the literature (e.g., Horrigan and Aldrich, 2002; Orio and Latorre, 2005). However, we argue that there are no assumptions in the determination of this parameter because it was directly obtained from the gating charge experimental data.

The parameters of the model obtained for WT plus those obtained from the $Q_c(V)$ curves of F380A mutant were used as starting values to perform a global fit of the $G(V)$ for F380A mutant voltage activation at different $\text{Ca}^{2+}$ concentrations data (Fig. 6 B). The fit is reasonably described in Materials and methods using the results of Fig. 3 E, and $z_L$ and $L_0$ were obtained from Orio and Latorre (2005). The final values of the different parameters were obtained as described in Materials and methods. The parameters obtained for WT were in good agreement with those previously reported except for $C$ and $E$, which were smaller and greater, respectively. The $E$ value was surprisingly large compared with those reported in the literature (e.g., Horrigan and Aldrich, 2002; Orio and Latorre, 2005). However, we argue that there are no assumptions in the determination of this parameter because it was directly obtained from the gating charge experimental data.

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DISCUSSION

The intracellular mouth of Kv channels is a great deal narrower than the internal vestibule of BK channels (Li and Aldrich, 2004; Brelidze and Magleby, 2005; Ghatta et al., 2006; Geng et al., 2011; Zhou et al., 2011). Moreover, closed channels are blocked by both the Shaker “ball” peptide and large quaternary ammonium ions, indicating that the bundle crossing does not impede the passage of ions (Wilkens and Aldrich, 2006; Thompson and Begemisch, 2012). Although the evidence suggests that the BK activation gate is not intracellular as in Kv channels but may be located in the selectivity filter, there is evidence that certain key residues in the S6 transmembrane segment can play an important role in channel gating (Wang and Brenner, 2006; Wu et al., 2009; Chen and Aldrich, 2011; Zhou et al., 2011; Chen et al., 2014). Interestingly, some of the single point mutations introduced in S6 (e.g., L312Q and L312D, A313D, and A316D) produced permanently open channels at a wide range of voltages (Chen et al., 2014). Although it is reasonable to think that many amino acids could be involved in coupling, the results of other groups make us to believe that some residues are more important than others in determining BK channel gating. For example, Zhou et al. (2011) performed a cysteine scan throughout the murine BK S6 for each of the 28 amino acid residues. Consistent with our assumption that not all of the residues in S6 play the same role in gating, three residues, L312, F315, and P320, corresponding to L377, F380, and P385 in the human BK produced extremely shifted G(V) curves. In these cases Zhou et al. (2011) were forced to assume the half-activation voltages and the voltage dependence of the G(V) curves to perform an analysis of the periodicity of gating perturbations produced by cysteine mutations in the S6.

The use of mutagenesis, together with gating and macroscopic currents and MD simulations, allowed us in the present study to gain insight on the way that the S6 transmembrane segment participates as an integration node in the BK channel. Our results indicate that there are some key amino acid residues in the S6 that are central players in the coupling of electrical and calcium-binding energies with the pore opening. We have found an amino acid residue (F380) located in the S6 transmembrane segment that, once mutated to a small hydrophobic amino acid (alanine), profoundly modifies ion conduction without affecting the gating charge of the channel. Gating currents in the F380A mutant are found to be similar in their kinetic and steady properties to the WT BK channel. Moreover, the mutant channel allows the detection of gating currents in the presence of physiological K+ concentrations, a result which shows that the voltage sensor can undergo its normal gating transitions in the presence or in the absence of permeant ions. Furthermore, the fact that ionic currents...
can be detected when the internal Ca\(^{2+}\) concentration is high is a clear indication that this mutant channel can open upon membrane depolarization and is Ca\(^{2+}\) activated. In contrast, by using the allosteric HA model (Horrigan and Aldrich, 2002), it is possible to arrive at the conclusion that the F380A mutation uncouples voltage-dependent gating and calcium binding from voltage sensors and increases the energy difference between closed and open conformations of the channel by \(\sim 18.7\) kJ/mol.

The present results show that replacing the large and hydrophobic phenylalanine by the small and hydrophobic alanine stabilizes the closed conformation of the BK channel. These results are at odds with what was proposed by Wu et al. (2009), because it is expected that a disruption of the interaction between F380 and L377, which interact only in the closed state, would stabilize the open state. A more plausible possibility is that actually residues F380 and L377 interact in both the closed and open state and the proper interaction is required for the normal gating. In fact, it is interesting to note that the interaction between the corresponding residue pair in KcsA (I100-F103) is present in both states, closed and open, as has been observed in the crystal structures of the closed (Doyle et al., 1998), the open, and the open inactivated (Cuello et al., 2010) structures, but only the former was used by Wu et al. (2009) for their MD simulations. We built a new open BK pore 3-D model based on MthK open structure (Posson et al., 2013), which has a larger radius at the mutation position than the KcsA-based model reported by Wu et al. (2009). The side chains of Phe380 and Leu377 in the BK model are not confronted to the pore and form a stable hydrophobic ring (Fig. 7 C). The nonbonded interaction energy between residue 380 and L377 is decreased by 20 kJ/mol when F380 is replaced by alanine. We hypothesize that the main stabilizing factor of the interaction between F380 and L377 of a neighboring subunit is the van der Waals interaction linking these two amino acids. Replacement of F380 by smaller hydrophobic amino acid residues should have the effect of stabilizing the closed state; note that this mutation produces a deactivation rate only five times faster than WT BK channel, whereas changes in \(I_0\) are \(\sim 1,700\) times; this implies that the effect is mainly on the activation rate, which becomes

![Figure 7. Molecular BK model and MD analysis.](image)

(A) Model built for open BK WT channel pore region (green balls are K\(^+\) and blue balls are Cl\(^-\)). (B) Side view of the hydrophobic ring formed by the F380 and L377 residues in BK WT (one subunit is transparent to better view). (C–E) Top view of the hydrophobic ring formed by the 380 and 377 residues in BK WT, F380A, and F380W, respectively. (F) Interaction energy between L377 and 380 residues (WT is considered a reference).
340 times slower in the F380A mutant. In contrast, large hydrophobic amino acids such as tyrosine or tryptophan should strengthen the interaction between the L377 and 380 residue, increasing the intrinsic equilibrium constant $I_d$. As predicted, this mutation decreased the free energy difference between open and closed state, whereas the result of the in silico calculation of the energy between W380 and L377 was $-16.5$ kcal/mol when the F380 and L377 interaction energy is taken as reference (Fig. 7 F). Although it appears to be a clear negative correlation between the strength of the interaction and the free energy difference between open and closed states, the effects on the allosteric constants are not so clear and suggest that the hydrophobic ring is involved in the coupling between sensors and pore opening in a complex way. The coupling between voltage sensor and opening is decrease in both mutants, whereas E and C are decreased only in one of the mutants (F380A and F380W, respectively).

In the lack of a crystal structure of BK channel, we can only speculate about the possible interaction between the sensors and the structure formed by residues L377 and F380. Previous modeling (Carvacho et al., 2008) and the present molecular model (Fig. 7) show that the smallest diameter of the internal vestibule is attained at the level of F380. We hypothesized that the hydrophobic ring can be the fulcrum of a lever where the forces produced by the sensors converge. For example, the distances between the four S6 helices can increase or decrease in response to calcium binding, and these movements could produce a stretching or relaxing of the structure formed by the set of the four S6 helixes, whereas this structure is really pivoting at the level of the hydrophobic ring and the diameter to this lever does not change substantially. In the case of voltage sensor, if the S4–S5 linker plays a similar role in BK channels as it does in Kv channels where this linker interacts with the S6, it is possible that the hydrophobic ring plays the same role as it does when the channel is activated by Ca$^{2+}$.

The gating phenotype of the F380A in the high Ca$^{2+}$ mutant resembles that of the Shaker ILT mutant (Ledwell and Aldrich, 1999), where the voltage activation curve is shifted to the right along the voltage axis, whereas most of the gating charge moves at voltages at which channels are closed. In the ILT Shaker mutant, channel opening is associated with the charge movement linked to the last rate-limiting transition in the ILT activation pathway. Thus, the F380A mutant isolated the BK opening voltage-dependent transition from earlier gating transitions and may provide a valuable experiment tool for teasing out the molecular basis of structural transitions in the activation pathway of BK channels.

Additionally, our results also support the idea that the inner vestibule in BK channels plays an important role in determining unitary conductance (Lippiat et al., 2000). Our fluctuation analysis experiments show that the F380A and F380W mutations decrease the channel conductance by $\sim 60\%$ and 50\%, respectively.

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