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The sixth transmembrane segment is a major gating component of the TMEM16A calcium-activated chloride channel

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Summary

Calcium-activated chloride channels (CaCC) formed by TMEM16A or TMEM16B are broadly expressed in the nervous system, smooth muscles, exocrine glands, and other tissues. With two calcium-binding sites and a pore within each monomer, the dimeric CaCC exhibits voltage dependent calcium sensitivity. Channel activity also depends on the identity of permeant anions. To understand how CaCC regulates neuronal signaling and how CaCC is, in turn, modulated by neuronal activity, we examined the molecular basis of CaCC gating. Here we report voltage modulation of TMEM16A-CaCC involves voltage-dependent occupancy of calcium and anion binding site(s) within the membrane electric field as well as a voltage-dependent conformational change intrinsic to the channel protein. These gating modalities all critically depend on the sixth transmembrane segment.

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TMEM16A is an ion channel protein with functions in a multitude of cell types. Peters *et al.* identified a specific domain that integrates the responses of this channel to three separable physiological stimuli.

Author Contributions

Declaration of Interests

The authors report no competing interests.

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Introduction

While cation channel gating and its contribution to the physiology of excitable cells have been studied extensively, gating mechanisms of anion channels are far less well understood. TMEM16A (TransMEMbrane protein of unknown function 16A, also known as Anoctamin-1 or ANO1) is a calcium-activated chloride channel (CaCC) (Caputo et al., 2008; Schroeder et al., 2008; Yang et al., 2008) in many epithelial cell types (Huang et al., 2009), nociceptive neurons (Cho et al., 2012), smooth muscles (Davis et al., 2010; Huang et al., 2012), and tumor cells (Ardeleanu et al., 2009; Duvvuri et al., 2012). A hallmark feature of TMEM16A-CaCCs is the apparent voltage dependence of its Ca²⁺ sensitivity. To understand how TMEM16A-CaCCs might contribute to cellular physiology, we examined how this channel is controlled by membrane potential, Ca²⁺ ions, and permeant anions.

TMEM16A-CaCC channels are homodimers of subunits each containing ten membrane spanning segments (Paulino et al., 2017; Dang et al., 2017). Intracellular Ca²⁺ ions are coordinated by the side chains of six amino acid residues (Yu et al., 2012; Tien et al., 2014; Dang et al., 2017; Paulino et al., 2017), and Ca²⁺ binding results in the activation of a channel that prefers permeation of Cl⁻ anions over small monovalent cations, and larger I⁻ anions over Cl⁻ anions (Caputo et al., 2008; Schroeder et al., 2008; Yang et al., 2008). Structural information suggests that anions traverse through a passage formed by portions of transmembrane helices 3–8 (Paulino et al., 2017; Dang et al., 2017). Mutations of residues surrounding this passage can modify anion selectivity (Peters et al., 2015; Dang et al., 2017) and anion/cation preference (Yang et al., 2012). The two pores of the dimeric channel likely function independently (Jeng et al., 2016; Lim et al., 2016).

TMEM16A open probability depends on not only intracellular Ca²⁺ but also membrane voltage and external anions (Perez-Cornejo et al., 2004; Contreras-Vite et al., 2016). While nanomolar intracellular concentrations of Ca²⁺ produce an outwardly rectifying current, micromolar and higher concentrations of Ca²⁺ elicit an Ohmic current The transition between rectifying and Ohmic currents can be achieved at lower concentrations of intracellular Ca²⁺ if extracellular Cl⁻ is replaced with more permeant anions such as I⁻ (Cruz-Rangel et al., 2017). A kinetic model of gating has been proposed to account for contributions from these different stimuli (Contreras-Vite et al., 2016), but the molecular mechanisms for these gating processes are unclear. We propose that CaCC gating stems from the following voltage dependent processes: i) Ca²⁺ binding occurs at a site within the membrane voltage field, causing an increase in the apparent binding affinity of the channel for Ca²⁺ at depolarizing voltages; ii) extracellular anions potentiate anion flux by binding to one or more modulatory sites inside the electric field of a multi-ion pore; and/or iii) TMEM16A channels undergo voltage dependent conformational changes leading to a nonlinear conductance/voltage relationship.

Here, we performed molecular dynamics (MD) simulations of the fungal homolog nhTMEM16 and incorporated TMEM16A structural information to dissect gating behavior in TMEM16A channel mutants. We found that transmembrane segment 6 (TM6), which is α-helical in both nhTMEM16 (Brunner et al., 2014) and TMEM16A (Paulino et al., 2017; Dang et al., 2017), is flexible in MD simulations of nhTMEM16 (Figures S1A and S1B),

and appears capable of adopting multiple stable conformations in TMEM16A. Furthermore, TM6 contributes residues to both the putative pore and the Ca^{2+} binding site (Brunner et al., 2014; Tien et al., 2014; Paulino et al., 2017; Dang et al., 2017). An alanine mutagenesis scan identified mutations that shifted the EC_{50} for Ca^{2+} gating at +60 mV. Subsequent analysis revealed that several of these mutations decoupled the Ca^{2+} , anion, and voltage-dependent gating processes, allowing us to study these processes in isolation. Our fast solution exchange pharmacology and patch clamp analyses established that depolarization, intracellular Ca^{2+} , and external anions all act upon TM6 to permit the flow of anions through the channel. Furthermore, the transfer of a mildly voltage-dependent intrinsic gating element involves at least two conformational changes of or about TM6, which influence and are influenced by the binding of intracellular Ca^{2+} ions and the identity of the permeating anion.

Results

Internal calcium and external anions modify voltage-dependence of TMEM16A channel activation

Gating of TMEM16A-CaCCs is sensitive to intracellular Ca^{2+} level and the identity of permeant anions. To characterize these gating processes, we transiently expressed in HEK293 cells TMEM16A from *mus musculus* with pEGFP fused to its C-terminus and channel properties identical to those wild-type channels (Tien et al., 2014; Peters et al., 2015), henceforth referred to as "wild-type", and employed whole cell patch clamp to record currents in voltage clamp mode. Exposing cells expressing wild-type TMEM16A to 400 nM Ca^{2+} allowed the channel to act as an outward rectifier, with the primary flux being inward flowing Cl^- ions (Figures 1A–1C). Replacement of Cl^- ions with I^- ions caused shifts in the reversal potential (Peters et al., 2015) as well as a sensitizing (left) shift in voltage dependence (Figures 1A–1C). Increasing intracellular Ca^{2+} to 1 mM caused the channel to a nearly voltage-independent conductance-voltage relationship (Figures 1A–1C). The physical bases for how membrane voltage, intracellular Ca^{2+} ions and extracellular permeant anions interact to contribute to TMEM16A gating remain intriguing open questions (Figure 1D).

Molecular dynamics simulation of nhTMEM16 and cryo-EM structures of TMEM16A-CaCC channels suggest the 6th transmembrane segment contributes to both anion permeation and calcium binding

To analyze the molecular basis of channel gating, we examined an atomic resolution structure of nhTMEM16 from *Nectria haematococca*, Protein Data Bank ID: 4WIS (Brunner et al., 2014), and several recently published structures of the TMEM16A channel, PDB ID: 5OYB, 5OYG, 6BGI and 6BGJ (Paulino et al., 2017; Dang et al., 2017), beginning with molecular dynamic simulations from the nhTMEM16 crystal structure (Bethel and Grabe, 2016). Briefly, nhTMEM16 was embedded in a phosphatidylcholine bilayer, and we ran a number of atomistic simulations consisting of 3.2 µs total aggregate time. Reasoning that components of the channel involved in gating would likely be dynamic and simulation may result in new protein conformations indicative of the gating motions, we aligned snapshots of the protein taken every 16 ns and calculated the root mean squared fluctuation (RMSF) of TM6 as well as the other two helices harboring residues involved in Ca²⁺-binding (TM7 and

TM8). Figure S1A shows the TM6 helix from our 200 snapshots overlaid onto the structure of the first snapshot. TM6 has a relatively high degree of flexibility, consistent with a poorly ordered or highly mobile region. For instance, the RMSF of the cytoplasmic end of TM6 is 1.5 Å (Figure S1B). Consistent with the notion that TM6 represents a region of flexibility, two recent studies of TMEM16A structures revealed that the conformation of TM6 depends on the cryo-EM conditions (Paulino et al., 2017; Dang et al., 2017). TM6 of TMEM16A reconstituted in nanodiscs (Dang et al., 2017) is largely α -helical, whereas the C-terminal half of TM6 in TMEM16A solubilized with Lauryl Maltose Neopentyl Glycol is unresolved, indicative of flexibility (Dang et al., 2017).

Alanine point mutants along TM6 produce significant shifts in calcium-dependent activation

Alignment of TM6 of TMEM16A with that of other family members (Figure 2A) demonstrates that it is particularly well conserved in TMEM16B, a highly similar CaCC. A model based on an atomic resolution cryo-EM structure of TMEM16A in nanodiscs was used to map the pore lining surface in dark grey (Figure 2B). Reasoning that TM6 at the interface between the permeation path and the Ca²⁺ binding site is ideally situated to contribute to channel gating, we performed alanine scanning mutagenesis of TM6 (Figure 2A). Of the 28 mutant constructs tested in HEK293 cells, 25 produced observable currents in inside out patches exposed to Ca^{2+} from the cytoplasmic side and held at +60 mV. Of the remaining three constructs with negligible currents, two (M638A and P654A) appeared to be poorly trafficked to the membrane, while L643A reached the membrane but was electrically non-functional (Figure S2A, S2B). To characterize channel mutants, we exposed inside out membrane patches from HEK293 cells to a series of solutions containing fixed concentrations of Cl^{-} ions but increasing concentrations of free Ca^{2+} ions (see sample traces in Figure 2C). We normalized the currents at various Ca²⁺ concentrations to current amplitude at a saturating concentration of Ca^{2+} (1 mM for most constructs; 10 mM for E650A) and fitted with sigmoidal concentration-response curves (equation 1 in STAR Methods), to derive the EC_{50} s and Hill coefficients (Figures 2D and 2E). Mutation at a known Ca²⁺ binding residue, E650 (Tien et al., 2014; Dang et al., 2017), and two asparagine residues, one of which may represent a 6th Ca²⁺-binding residue (Brunner et al., 2014; Dang et al., 2017), produced significant reductions in Ca^{2+} sensitivity (red bars with black hashmarks). Numerous other residues N-terminal to the Ca²⁺ binding residues of TM6 also produced statistically significant reductions in apparent Ca^{2+} sensitivity (Figure 2D). However, we also noticed that several residues within TM6 increased apparent Ca²⁺ sensitivity, including some that significantly reduced the Hill coefficient (Figure 2E). Because these mutated residues are not implicated in Ca²⁺ binding, we hypothesized that they may increase apparent Ca²⁺ sensitivity by producing conformational changes that facilitate either Ca²⁺ ion binding or channel activation after Ca²⁺ binding.

Alanine substitution of I637 or Q645 in TM6 enhances calcium sensitivity and enables channel activation in the absence of internal Ca²⁺

Next, we characterized mutants resulting in significant increases in apparent Ca^{2+} sensitivity, reasoning that they were likely involved in channel gating rather than directly affecting Ca^{2+} -binding. We tested all such alanine mutant constructs in Ca^{2+} -free internal solutions

using whole-cell patch clamp. Remarkably, I637A and Q645A, the two mutations producing the largest sensitizing shifts, resulted in an outward rectifying current voltage relationship within the physiological range in the absence of Ca^{2+} (Figures 3A–3C). Channels with mutations at G640, K641, L648, and I653 remained incapable of opening in the absence of Ca²⁺ up to +200 mV (Figure S3A, S3B). We confirmed the *bona fide* "Ca²⁺-independent" activation in Q645A by showing that channel activity remained the same with 5 mM BAPTA and 1 mM EDTA added to our normal "Ca²⁺ free solution" (which contained 5 mM EGTA) and with omission of Mg²⁺ ions, which do not normally activate TMEM16A (Ni et al., 2014), out of concern that the mutation may have sensitized the channel to Mg^{2+} . In addition, equivalent mutations in the TMEM16B channel also exhibited Ca²⁺-independent activation, albeit requiring more depolarizing voltages and producing smaller apparent whole cell currents (Figures S3C and S3D). To confirm that these mutations changed TMEM16A gating properties while preserving its channel permeation characteristics, we performed non-stationary fluctuation analysis of Q645A (Figures S4A and S4B; Heinemann and Conti, 1992) and showed that the single channel conductance of Q645A was 3.2 ± 0.4 pS, which is comparable to that of WT at 2.8 ± 0.5 pS (Figure S4C) and consistent with previous reports (Manoury et al., 2010; Lim et al., 2016; Ta et al., 2016). We also exposed this channel to a series of bi-ionic anion replacement solutions with internal solutions containing Cl⁻ and external solutions containing another anion species or 14 mM NaCl supplemented with D-mannitol, and found a selectivity sequence among the anions tested of P_I>P_{Cl}>P_F>P_{MES}>P_{Aspartate}>P_{Gluconate}, and a P_{Na+}/P_{Cl} of 0.10 (Figures S4D–S4F), consistent with previous findings for WT TMEM16A. Finally, we compared the conductance-voltage relationships of Q645A exposed to NaCl and NMDG-Cl and found no change in the waveform, suggesting that the monovalent cationic species did not influence the conductance-voltage relationship. Taken together, these results suggest that TM6 mutations are capable of shifting Ca²⁺ sensitivity and allowing the mutant channel to activate in the absence of Ca²⁺ at depolarizing voltages without affecting ion selectivity or the single channel conductance.

Permeant anions potentiate conductance via an externally situated binding site

Notably, the two mutants with sensitizing shifts in both voltage-dependent and Ca²⁺dependent gating remained sensitive to the concentration and identity of external anionic species, similar to an effect seen previously for heterologously expressed TMEM16A and endogenous CaCC in *Xenopus* oocytes (Perez-Cornejo et al., 2004; Contreras-Vite et al., 2016). Furthermore, plotting I637A and Q645A whole cell conductances for Cl⁻ and I⁻ against voltage revealed that the channel open probability (P_o) did not saturate even at strongly depolarizing (e.g., +150 mV) potentials (Figure 3C). When plotted with a semi-Log ordinate, normalized G(V) increased linearly with voltage even at strongly depolarizing potentials (Figure 3D), unlike outwardly rectifying potassium channels (Zagotta et al., 1994) with P_o approaching 1 under depolarizing conditions due to full activation of all voltage sensors. Replacing external Cl⁻ with I⁻ caused the G(V) curve to shift leftward while retaining a positive slope (Figure 3D). These results implicate the involvement of anions with a binding site accessible to the external medium in a rate-limiting process for the apparent voltage-dependence of channel conductance. To parse the effects of external anion binding, we recorded whole cell currents of Q645A with internal Cl⁻ replaced with aspartate ions that cannot easily pass through CaCC channels, to yield near unidirectional anionic fluxes with amplitudes proportional to the k_{ON} of anions from the extracellular milieu to an external binding site. As k_{ON} is also dependent on the concentration of anions in the bulk external solution, we applied varying concentrations of Cl⁻ or I⁻ and quantified an EC₅₀ for each anion at a series of strongly depolarizing voltages to eliminate the trace influence of internal anions (Figures S5AS5C). Subsequently, these EC₅₀s were Log transformed and fit with equation 2 (in STAR Methods) to establish the depth in the voltage field of this anion sensitive site for channel potentiation (Figure S5D). The linear fits to these points yielded an apparent z8 of 0.058 ± 0.013 for Cl⁻ and 0.057 ± 0.012 for I⁻. We reasoned that the outermost site of channel interaction with anions likely exists at that depth, consistent with a previous finding suggesting a shallow anion binding site and pore blocking site accessible from the extracellular space (Peters et al., 2015). Moreover, we reasoned that I⁻ binds to the same site with higher affinity than Cl⁻ because both ions had the same voltage dependence (z8).

Permeant anions and intracellular calcium affect an innate voltage dependent gating process of TMEM16A

We next sought to separate the effects of the modulatory site from other intrinsic components of the overall conductance-voltage relationship. We reasoned that eliminating Ca²⁺-dependent gating should allow us to probe whether additional voltage sensitivity exists, so we examined I637A and Q645A mutant channel activation in the absence of intracellular Ca²⁺. By fitting the whole cell G(V) curves between +150 and +180 mV in the presence of external Cl⁻ or I⁻ with equation 2 and using $z\delta = 0.06$ to establish an upper limit for the slope of the semi-Log plot, we could separate values for the apparent aniondependent effect producing non-saturating conductance at extreme depolarizations from a residual voltage dependence of the channel (Figures 3E–3G). Fitting this voltage dependence with a two-state Boltzmann relationship (equation 3 in STAR Methods) suggests that the mutant channel itself undergoes a discrete voltage-dependent rearrangement, henceforth referred to as "PA" for the voltage dependent probability of the channel being in the activated conformation. TMEM16A I637A has a $V_{1/2}$ of 72.2 ± 2.7 mV (n = 12) while Q645A has a $V_{1/2}$ of 52.6 \pm 5.3 mV (n = 14) in the presence of Cl⁻. Replacement of Cl⁻ with I⁻ led to a significant left shift in $V_{1/2}$ in I637A to -3.6 ± 3.4 mV (n =12) and in Q645A to -1.1 ± 6.2 mV (n = 13) (Figures 3F and 3G). Thus, extracellular anions permeating through the pore influence its intrinsic voltage-dependent gating, independent of Ca²⁺-occupancy of the mutant channel. We then examined the double mutant I637A Q645A, and found a negative shift in the voltage dependence compared to that of either single mutant (Figures 3H and 3N). This result suggests that these two mutations likely introduce separate, distinguishable changes to gating stemming from movements of or about TM6.

It was notable that alanine substitution for the G640 residue between I637 and Q645 also increased Ca^{2+} sensitivity (Figure 2D). Glycine residues found along the pore lining helices of ion channels such as KcsA and the *Shaker* family potassium channels are implicated as hinge bending elements crucial for channel gating (Jiang et al., 2002; Ding et al., 2005). Typically glycine to alanine mutations would reduce hinge bending. Remarkably, G640A

facilitated activation of the TMEM16A by Ca^{2+} . This raises the possibility that if G640 acts as a flexible hinge or inflection point, it may favor a closed channel conformation or hamper Ca^{2+} binding. We tested whether the reintroduction of glycine nearby could restore normal Ca^{2+} -dependence and found that two double mutations of G640A, with I636G or M638G, both decreased Ca^{2+} -sensitivity to near wild-type levels (Figures 3I–3K). Thus, the presence of a glycine at or near position 640 is important for Ca^{2+} -dependent gating at physiologically relevant Ca^{2+} concentrations. We next asked whether this glycine might also contribute to the equilibrium of voltage-dependent gating of I637A or Q645A. Recording of double mutants I637A G640A or Q645A G640A in Ca^{2+} -free solutions in whole cell patch clamp conditions revealed that G640A imparted an additional left-shift in voltagedependence to Q645A but not I637A (Figures 3L–3N). This result suggests that the flexible hinge afforded by G640 plays a role in Q645A but not I637A channel gating, further reinforcing the notion that TM6 likely undergoes at least two sequential movements during voltage-dependent gating.

A lysine in TM6 contributes voltage-dependence to TMEM16A activation

The Ca²⁺ sensitivity was increased by alanine substitution for the K641 basic residue within the pore region (Figures 2D and 4A), which has been implicated in both anion conductance and gating of the channel (Peters et al., 2015). Because the double mutant of K641A with Q645A did not show expression in HEK293 cells, we examined the charge neutralization and charge reversal double mutant constructs Q645A K641Q and Q645A K641E using whole cell patch clamp in Ca²⁺ free internal solution (Figure 4B), and found significant right-shifts to the voltage-dependent P_A (Figures 4C, 4D and 4F). These shifts persisted even when extracellular Cl⁻ was replaced by I⁻, albeit with smaller magnitudes. Furthermore, Q645A K641E exhibited a left shift in P_A in the presence of 1 mM intracellular Ca²⁺ similar to that in Ca²⁺-free solutions (Figures 4E and 4F). However, Ca²⁺ ions were no longer able to push the channel to its "Ohmic" conformation, nor were they able to relieve the dependence of P_A on external anionic species as observed in the wild-type channel with high Ca²⁺. Taken together, these results suggest that K641 at the lower end of TM6 is critical for regulating the dependence of channel gating on voltage, Ca²⁺ and anions (Figure 4G).

Mutation of calcium-binding residues shifts TMEM16A voltage-dependence

TM6 harbors 2 of the 6 amino acids critical for coordinating Ca²⁺, including 5 acidic residues and an asparagine. Having found that both N646 and N647 produced right-shifts in the EC₅₀ for Ca²⁺ (Figure 2D), we examined aspartate substitution for either asparagine using inside-out patch clamp electrophysiology to test whether introduction of an additional negative charge to the Ca²⁺ binding sites causes a sensitizing shift in the EC₅₀ for Ca²⁺. N646D but not N647D caused a potent increase in Ca²⁺-sensitivity [EC₅₀: 216 ± 15 nM (n = 9); Figures S6A and S6B], consistent with the observation that N646 participates in Ca²⁺ binding (Dang et al., 2017) (Figure 5A). As I637A and Q645A are mutations within TM6 capable of facilitating activation of the channel in the absence of Ca²⁺ ions, they also provided the opportunity to parse the role of Ca²⁺ from the voltage-dependence of this domain. P_A of I637A N646A and I637A E650A double mutants in Ca²⁺-free solutions was the same as that of I637A single mutant (Figure 5B). Conversely, both N646A and E650A produced strong sensitizing shifts in P_A when placed in the Q645A background (Figure 5C).

We considered two possible explanations for this phenomenon: either these Ca^{2+} binding residues represent negative "gating charges," or the absence of Ca^{2+} in its binding site results in destabilization of the configuration of the Ca^{2+} binding sites due to electrostatic or steric effects. As N646 is not charged, we favored the second explanation, and to test it, we made additional double mutants of Q645A with the remaining Ca^{2+} -binding residues. Q645A E698A appeared similar to wild-type, but Q645 E701A, Q645A E730A and Q645A D734A all resulted in strong sensitizing shifts in P_A (Figures 5D–5F), supporting the notion that the negative charges on these residues may repel each other in the absence of Ca^{2+} as a stabilizing ligand. Indeed, E698A is spatially removed from the other 5 residues and is likely to only interact with one of the two Ca^{2+} ions, while E650, E701 and E730 form a ring (Figure 5A, lower panel) and are likely to interact with both Ca^{2+} ions.

Crosstalk between voltage and calcium-dependent gating in TMEM16A

To investigate the interaction between voltage and Ca^{2+} in the wild-type channel, we used an internal solution containing 400 nM Ca^{2+} to activate the channel and estimated the voltage-dependent P_A as before, and found that the wild-type TMEM16A had a residual voltage dependence (Figures 6A and 6C). As in Q645A, WT TMEM16A showed differential gating effects for Cl⁻ and I⁻, as external I⁻ caused a desensitizing shift in P_A (Figures 6A and 6C). We then examined how P_A varied with Ca^{2+} . Reducing intracellular Ca^{2+} to 200 nM caused a right-shift whereas increasing intracellular Ca^{2+} to 1500 nM led to a left-shift in P_A (Figures 6B and 6C), suggesting that voltage and Ca^{2+} likely combine to determine the conformation of TM6.

Having found that both voltage and Ca²⁺ play a role in TM6-dependent gating, we varied holding potentials and pulse durations to probe the time-dependence of gating effects in wild-type TMEM16A to ascertain how these two stimuli interact to produce the complex gating behavior. Notably, the wild-type channel exhibits a markedly biphasic activation upon depolarization, with an apparent "instantaneous" (I_{inst}) component followed by a second, more slowly activating component (Figure 6D). First, TMEM16A in whole cell clamp exposed to 400 nM intracellular Ca²⁺ was held at a series of prepulse voltages. We then stepped the membrane potential to a supraphysiological range in 10 mV intervals and compared the size of Iinst immediately following the voltage jumps. Only a small fraction of the current was activated very quickly from -120 mV (Figure 6D, left panel arrows): mean I_{inst} activated upon pulse to +120 mV was 0.09 ± 0.01 of the size of the steady state current at +120 mV, where the driving force for Cl^{-} is the same size. As this I_{inst} activates upon depolarization, we propose that it represents a fraction of channels with a single Ca²⁺ ion bound that are closed but "primed" for activation upon depolarization. Following the step to +120 mV, activation to another open state with a slower time course can then proceed (see Figure 6D, inset). When these same cells were held at +120 mV prior to the voltage steps and then stepped from +120 mV back to -120 mV, the inward I_{inst} was 0.86 ± 0.02 the magnitude of the steady state current at +120 mV, demonstrating that prolonged activation at +120 mV allows a much larger inward Iinst current to be briefly observed following steps to negative potentials. Instantaneous current amplitudes from different prepulse holding potentials normalized against the baseline activation at +120 mV are shown in Figure 6E.

In contrast, when the same experiment is performed with the outwardly rectifying Q645A mutant in the absence of internal Ca^{2+} , I_{inst} at +120 mV following a -120 mV prepulse was 0.65 ± 0.04 of the steady state at +120 mV (Figure 6F), while the inward I_{inst} amplitude at -120 mV following a +120 mV prepulse was 0.29 ± 0.05 of the magnitude at +120 mV. Thus, while Q645A channels can be more quickly activated from a -120 mV holding potential by depolarization, prolonged depolarization does not stabilize a conformation that allows for prolonged inward current (Figure 6G). Reasoning that this likely resulted from the absence of Ca²⁺ binding, we performed a similar analysis of I_{inst} using the Ca²⁺-binding mutant E650A, in the presence of 1 mM internal Ca²⁺. Under these conditions, E650A demonstrated I_{inst} of 0.80 \pm 0.09 of the +120 mV steady state magnitude when pulsed from -120 mV, but only 0.21 \pm 0.03 in the inward direction when pulsed to -120 mV from +120mV, and indeed, the Iinst magnitude appeared identical at every pulse voltage independent of prepulse level (Figures S7A and S7B). We propose that E650A eliminates 1 of the 2 Ca²⁺binding sites, and for the WT channel, initial Ca²⁺ binding influences TM6 to adopt a conformation that permits binding of the second Ca²⁺. Moreover, TM6 movement is voltage-dependent, and all three steps (first Ca²⁺ binding, TM6 movement, and second Ca²⁺ binding) are required for full activation.

Next, we probed the kinetics of activation and deactivation after prolonged periods of activation (at +120 mV) or deactivation (at -120 mV) (Figure 6H). We fit time constants to these time courses to estimate activation and deactivation rates (using equations 4.1 and 4.2 in STAR Methods), the onset of the slow component of deactivation (Figure 6H, top panels, arrows), and the loss of instantaneous current after activation (Figure 6H, bottom panels, arrows). We found that for Q645A with no internal Ca²⁺, I_{inst} remained constant following activation for different durations, and the time course of deactivation exhibits a single, "fast" exponential decay even after prolonged depolarizing pulses. In contrast, wild-type TMEM16A with 400 nM intracellular Ca^{2+} showed pronounced biphasic activation at +120 mV (Figure 6H, see also Figure 6D), and increasing the duration of this pulse before hyperpolarization to -120 mV increased the relative contribution of a more slowly deactivating component at -120 mV. Quantification of the fast and slow components of the deactivating current following a 300 ms pulse to +120 mV revealed that substituting Cl⁻ with I⁻ further increased the relative contribution of slow deactivation (Figure 6I). Conversely, reducing intracellular Ca²⁺ from 400 nM to 200 nM decreased the relative contribution of slow deactivation with either Cl⁻ or I⁻ as the external anion (Figure 6I). Thus, the rate of onset of this slow phase is dependent on both intracellular Ca²⁺ and extracellular anions. The non-Iinst components of current onset during depolarization were fit with double exponential time constants (τ_{a1} , fast; τ_{a2} ; slow) for WT channels with 200 and 400 nM internal Ca²⁺ and with external Cl⁻ and I⁻. The amplitude of the slowly deactivating component at -120 mV as a portion of the total initial current at -120 mV $(\tau_{ON} \text{deact}_{s/t})$ increased with the same time course as τ_{a2} of current activation, which was insensitive to the change in internal Ca²⁺ or external anionic species (Figure 6J). Instead, the balance between the instantaneous and slow activation rates were affected by anions and Ca²⁺ (Figure 6I), suggesting that conformational changes in the protein, rather than the ion binding events, are likely rate limiting for full activation of the channel.

We also modeled channel deactivation from +120 mV to -120 mV with double exponentials $(\tau_{d1}, \text{fast}; \tau_{d2}; \text{slow})$ and compared them to the decay in the amplitude of the instantaneous current of a subsequent pulse to +120 mV following pulses with variable length to -120 mV (Figure 6H, lower panel, arrows; Figure 6K). We found that the time course of the "loss of instantaneous" ("L.O.I.") current occurred with a τ similar to τ_{d2} . Slow deactivation (τ_{d2}), unlike for activation, appeared to be slowed by exchange of Cl⁻ for I⁻ (Figure 6K), suggesting hysteresis in the sequence of events leading to activation and anion permeation. We also noticed that the fast τ_{d1} varied with both intracellular Ca^{2+} and the species of permeant anion; however, the contribution of this τ became negligible with longer activating pulses in wild-type (but not in Q645A with 0 Ca²⁺ or E650A with 1 mM Ca²⁺ where both τ of deactivation and its amplitude were unchanged by increasing prepulse duration) suggesting that it may represent deactivation from a conformation with only a single bound Ca^{2+} ion. Altogether, these results are consistent with a model where consecutive conformational changes in TM6 occur: occupancy of the first Ca²⁺ binding site leads to the outwardly rectifying conductance with fast kinetics while occupancy of the second Ca²⁺ binding site leads to the voltage-independent conductance with slow kinetics.

Discussion

Channel potentiation by Cl⁻ or I⁻ ions represents a positive feedback regulation of channel activity by permeant anions. Anion discrimination is especially relevant to TMEM16A function in the thyroid (Twyffels et al., 2014), where it allows for I⁻ influx in the apex of the thyrocyte. Newly discovered roles for TMEM16A and the closely related TMEM16B in neurons also benefit from the complex gating apparatus of these channels. In thalamocortical neurons, which act as integrators of sensory input and relay information to the cortex, TMEM16B mediates spike frequency adaptation to allow neuronal self-inhibition (Ha et al., 2016), and the multi-phasic Ca^{2+} and voltage-dependent gating is ideally suited to allow it to do so. At low [Ca²⁺], CaCC channels do not conduct outward currents, so they do not contribute to the resting potential in most neurons. As $[Ca^{2+}]$ rises in the cell following action potential firing or excitatory synaptic inputs, Ca²⁺ binds to its first binding site "priming" the channel to be activated upon strong depolarization. This priming will not change the action potential threshold of the neuron. With depolarizing spikes, a small population of channels activate quickly to allow an inward flux of Cl⁻ (outward current), and when the spike frequency increases, so does the influx of Ca^{2+} , increasing the number of CaCC channels able to activate on the time scale of neuronal spiking and providing negative feedback against high frequency activation of mature central neurons. The identification of TM6 as a voltage and Ca²⁺-dependent gating element further improves the outlook for designing and understanding the pharmacological profiles of channel modulators. Several TMEM16A inhibitors have been identified, including ostensible pore blockers (Peters et al., 2015) as well as several with still unknown mechanisms of action (Namkung et al., 2011; Oh et al., 2013; Seo et al., 2016). However, binding sites for these drugs have not been mapped. Identifying a transmembrane segment linking permeation and gating would facilitate future development and improvement of compounds to influence TMEM16A activity for both in vitro studies and in vivo applications.

Based on results from several approaches presented in this study, we propose a model for how the molecular machinery involved produces synergistic channel gating interactions between internal Ca²⁺, external anions, and intrinsic components of the channel (Figure 7). In the absence of internal Ca^{2+} , the wild-type channel is closed, but can bind Ca^{2+} ions. As $[Ca^{2+}]$ increases, association of one Ca^{2+} ion allows anions to enter the outer pore, and "primes" the channel to undergo a fast voltage-dependent conformational change involving TM6 to mediate anion conductance upon depolarization. Replacement of external Cl⁻ with I ⁻ likely increases the dwell time and/or interaction of the anion in the outer pore to stabilize this conformation of TM6, thus favoring the subsequent voltage-dependent conformational changes at less depolarizing voltages. Depolarization allows the channel to enter its first open state with fast kinetics, and this state favors binding of a second Ca²⁺ ion. Association of a second Ca²⁺ ion results in the stabilization of the fully activated conformation of TM6, resulting in much slower deactivation kinetics and ostensibly Ohmic anion conductance. When I637 or Q645 was replaced with alanine, the conformational change normally associated with the first Ca²⁺ binding event could take place in the absence of divalent cations, allowing the channel to be opened to its first conductive conformation by depolarization alone (see Figures 3E-3G, 7B); however, these two mutations produce nonequivalent effects on gating and likely point to the existence of separable, sequential conformational changes in TM6 (Figure 7E), which are also strongly dependent on the properties of several other amino acids on TM6. Without Ca^{2+} binding, channels cannot enter the second open state, which requires bound Ca^{2+} to be stable, thus explaining the absence of the slower activating current and failure of these mutant channels to adopt an "Ohmic" conductance.

The idea that TM6 is a dynamic segment sensitive to external stimuli is consistent with previous results examining this region of the protein. TM6 had been predicted to form an external reentrant loop based on topology prediction algorithms (Yang et al., 2008), before studies using cysteine accessibility and cross-linking analysis refined the topology to show that TM6 indeed traversed the membrane (Yu et al., 2012; Tien et al., 2014). While the key residues highlighted in this study are well conserved with the closely related TMEM16B-CaCC, the conservation is poor for family members that are not Cl⁻ channels (Figure 2A). However, the Ca²⁺-binding residues in TM6-8 are highly conserved, and our MD simulations of nhTMEM16, and the TMEM16A structures (Paulino et al., 2017; Dang et al., 2017) demonstrate a high likelihood that this portion of the structure adopts several stable conformations involving different packing of TM6 with the surrounding helices. Thus, the role of TM6 in transducing Ca²⁺-binding into protein conformational change is likely conserved. G640, a proposed hinge examined in this study, influences the interaction between Ca²⁺-binding and conformational changes in TM6 associated with voltage-dependent channel gating.

While we could identify an innate voltage-dependence to the channel by using site-directed mutagenesis to eliminate the requirement of Ca²⁺ for activation, the underlying source of the voltage-dependence is unclear. The slope of voltage dependence was shallow: for example the G(V) slope (k in equation 3 in STAR Methods) of Q645A was 39 ± 4 mV, corresponding to an apparent transferred gating charge of $0.65 \pm 0.06 q_e$ per subunit – much smaller than the 3–4 q_e per subunit for voltage-gated ion channels (Zagotta et al., 1994). The charge may

be carried by TM6 itself – K641, when mutated to alanine, did shift the "Boltzmann" component of gating both with and without internal Ca^{2+} – however, the mutant channel did retain underlying sensitivity to voltage, suggesting that other residues contribute to the gating charge. Of the side-chains predicted to coordinate Ca^{2+} , five are acidic and may also be sensitive to voltage (Yu et al., 2012; Tien et al., 2014; Paulino et al., 2017; Dang et al., 2017). Moreover, 5 of the 6 Ca^{2+} coordinating side chains caused shifts in voltage-sensitivity when mutated in the Q645A background. However, as with K641 mutants, removal of one of these negative charges shifted, but did not abolish, voltage-dependence. Furthermore, the persistent ability of the more permeant I⁻ to modify voltage-dependence suggests that anion presence in the permeation pathway may have a significant role in the conformational changes associated with gating.

In summary, we have determined that the 6th transmembrane segment, which harbors both pore lining residues and Ca²⁺ binding sites (Dang et al., 2017), represents a critical component of the TMEM16A gating mechanism. As other regions of the protein have also been implicated in regulating channel gating, including the N-terminus, the C-terminus, and several of the intracellular loops (Ferrera et al., 2009; Ferrera et al., 2011; Xiao et al., 2011), future studies will be necessary to elucidate how these regions may interact either directly or allosterically with the machinery described here. Since some members of the TMEM16 family show vastly different channel function and/or scramblase behavior dependent on structural motifs in the same region of the protein (Yang et al., 2012; Yu et al., 2015; Bethel and Grabe, 2016), testing for homologous regions in those proteins may help in developing a theory to explain the functional disparities between family members and to identify commonality in regulatory mechanisms.

STAR Methods text

CONTACT FOR REAGANT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Lily Jan (lily.jan@ucsf.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell lines—HEK293 cells (ATCC, RRID:CVCL_0045) were maintained at 37 °C and 5% CO2 in Dulbecco's Modified Eagle's Medium supplemented with 10% Fetal Bovine Serum and 1% penicillin/streptomycin, and were passaged upon reaching confluency (every 2–4 days) using 0.05% Trypsin-EDTA. These are a commercially available cell line and are authenticated by the manufacturer. Sex: female.

METHOD DETAILS

Molecular dynamics—Molecular dynamics simulations of nhTMEM16 were performed to predict regions of structural mobility in the protein. Modeller 9.15 (Sali and Blundell, 1993) was used to build in loops missing from the crystal structure (PDB: 4WIS). Using CHARMM-GUI (Lee et al., 2016), nhTMEM16 was embedded in a POPC bilayer. The entire system consisted of 356,426 atoms with 710 lipids. Simulations were run at constant pressure (1 bar) and temperature (303.15 K) using the CHARMM36 forcefield (Klauda et

al., 2010). Using the CUDA enabled version of Amber (Case et al., 2005), eight parallel simulations were run for 400 ns each, yielding an aggregate simulation time of $3.2 \,\mu s$.

Generation of cDNA reagents—TMEM16A (splice variant 'a') and TMEM16B clones from *mus musculus* (NCBI: NP_001229278.1,) in a pEGFP-N1 expression vector were used for all experiments. Single point mutations were introduced using Phusion polymerase for PCR, as instructed by the manufacturer, and mutagenic oligonucleotides were purchased from IDT-DNA Technologies (Iowa). Mutagenic primers consisted of reverse complimentary 33-mers centered upon the codon representing the targeted amino acid, whereupon nonsynonymous base pairs were introduced to the sequence by PCR. Mutant clones were confirmed by Sanger sequencing at Quintara Biosciences (South San Francisco, CA).

Transfection—24 hours before recording, cells were transiently transfected with TMEM16A constructs using Lipofectamine 2000 (Invitrogen), in antibiotic free media, as recommended by the manufacturer. On the morning before recording, cells were re-plated onto poly-L-lysine coated coverslips by dispersion with 0.05% trypsin followed by neutralization and dilution with bath medium. Cells were allowed to settle for at least 1 hour prior to recording.

Immunocytochemistry—To examine surface expression of TMEM16A-GFP mutants, transfected HEK293 cells were immunostained using a cell-surface marker and overlap of this marker with GFP. After a 24 hour transfection period, cells were fixed in 2% paraformaldehyde for 10 minutes on ice. Cells were washed two times in 1× Phosphate Buffered Saline (PBS), 10 minutes each. Cells were incubated in blocking solution with 1% heat inactivated goat serum and 1:150 wheat germ agglutinin (Vector Labs, Burlingame CA) for 1 hour at room temperature. Cells were then washed in 1× PBS three times, 10 minutes each. The coverslips with the cells were then mounted onto Fisher premium frosted microscope slides with DAPI Fluoromount-G (SouthernBiotech). Cells were visualized using a Leica SP8 confocal microscope at $63 \times .30$ µm thick Z-stacks were acquired.

Patch clamp electrophysiology

Patch pipettes: All electrophysiology experiments were performed using borosilicate glass purchased from Sutter Instrument Company (Novato, CA). Patch pipettes were pulled from 1.5/0.86 (OD/ID) glass and polished to 2–2.5 M Ω resistance (inside-out patch) or 3–5 M Ω (whole cell patch). Rapid solution exchange pipettes were pulled from Theta glass (Sutter) and manually broken to approximately 50 μ M in diameter by scoring with a ceramic tile.

Solutions: For patch clamp electrophysiology, coverslips were transferred into a recording bath solution containing 140 mM NaCl, 10 mM Hepes and 5 mM EGTA, pH7.2/NMDG. All recordings were made at room temperature. Internal solutions containing buffered Ca²⁺ were made at pH 7.2 by combining Ca(OH)₂ and EGTA to a ratio predicted by CaBuf software (KU Leuven) and then adding 140 mM NaCl or NMDG-Cl and 10 mM Hepes to the desired volume. Ca²⁺ concentrations were subsequently measured directly against calcium standards using a commercial Ca²⁺ buffer kit (Invitrogen) in a Fluo-8 or Fluo-8FF fluorescent assay in

a BioTek Synergy H4 plate reader. Solutions containing free Ca^{2+} greater than 100 µM were made by adding CaCl directly to an EGTA-free version of our 140 mM NaCl solution, to the desired concentration. Solutions for biionic experiments were made by substituting the desired anions or cations for Na⁺ or Cl⁻ in equimolar concentrations. Solution osmolality was adjusted to 305 ± 5 mOsm by addition of D-mannitol. All laboratory chemicals were purchased from Sigma-Aldrich, unless otherwise noted.

Data collection and equipment: Perfusion exchange was performed using a VM-8 perfusion apparatus with Octaflow software (ALA Scientific). Recordings were made using an Axon Instruments Multiclamp 700 with Digidata 1440, and were collected into pClamp10 software. All patch clamp seals were allowed to reach at least 3 G Ω resistance before patch rupture, but typical seal resistance usually exceeded 10 G Ω . All recordings were made using a 1M KCl agar bridge to prevent baseline fluctuation at the reference electrode.

Rapid solution exchange experiments used an Automate piezo driven exchanger with Thetaglass pipette applicators pulled from 1.5 mm OD Theta glass (Sutter Instrument). Data were collected at 10 kHz sampling rate and low pass filtered online at 2 kHz (biionic recordings), or at 100 kHz and filtered at 10 kHz, then reduced by a factor of 3, resulting in digitization at 33.3 kHz for non-stationary analysis recordings.

QUANTIFICATION AND STATISTICAL ANALYSIS

Patch clamp electrophysiology

Data analysis: All offline data analysis was performed using Graphpad Prism 6, Clampfit 10 and Microsoft Excel. Concentration-dependence curves for Ca2+ or permeant anions (X-) were generated by fitting data to an equation of the form:

$$\frac{I}{I_{MAX}} = \frac{1}{1 + \left(\frac{K_D}{[ion]}\right)^{n_H}}$$
Equation 1

where I/IMAX denotes current normalized to the maximum amplitude in the highest [Ca2+] or [X-] tested, KD denotes the dissociation constant for that ion and nH denotes the Hill coefficient.

Voltage-dependence of EC50 values for gating by extracellular anions were calculated by fitting data to an equation of the form:

$$LOG(EC_{50}) = LOG(EC_{50})_{0 mV} - \frac{z\delta FV}{RT}$$
 Equation 2

Normalized conductances were parsed by dividing the resulting $z\delta$ values from the overall G(V) relationship to reveal Boltzmann shaped residual curves, which were fit to voltagedependent gating data using an equation of the form:

$$G(V) = \frac{1}{1+e\left(\frac{V_{1/2}-V}{k}\right)}$$
 Equation 3

where $V_{1/2}$ represents the voltage where half of the channels are active, and k represents a slope factor for the sigmoidal curve.

To quantify the time courses of channel activation and deactivation, voltage clamp traces were fit with single or double exponential curves with equations of forms, respectively:

$$I_t = I_0 - A\left(1 - e^{(-t/\tau)}\right) \quad \text{Equation 4.1}$$

or

$$I_t = I_0 - \left[A_1 \left(1 - e^{-t/\tau_1} \right) + A_2 \left(1 - e^{-t/\tau_2} \right) \right]$$
 Equation 4.2

Single channel conductance was estimated using non-stationary fluctuation analysis of current traces by plotting signal variance against amplitude and fitting a polynomial function:

$$\sigma^2 = iI - \frac{I^2}{n}$$
 Equation 5

where σ^2 denotes signal variance, I denotes macroscopic current, i denotes single-channel current and n the number of channels in the patch. Single channel conductance (γ) was calculated by dividing i by (Vm-ERev) for Cl⁻ ions. Outlying values for signal variance (ie: system noise vs. channel noise) were identified and eliminated by iteratively calculating standard deviation across samples at each time point and eliminating values larger than 4 S.D. from the mean after each calculation until an unchanging value for variance was reached, as has been described (Heinemann and Conti, 1992). Permeability ratios were determined from biionic conditions using a reduced form of the Goldman-Hodgkin-Katz voltage equation:

$$E_{REV} = \frac{RT}{zF} Ln \frac{P_X [X^-]_o}{P_{Cl} [Cl^-]_I} \quad \text{Equation 6}$$

where z represents the valence of the anion, PX represents relative permeability of ion species "X" and F, R and T have their usual thermodynamic meanings.

Sequence alignments were performed using Clustal Omega and visually rendered in BoxShade via the ExPasy web portal.

Statistical analysis: Statistical analysis employed Student's T-tests for single comparisons, or one-way ANOVA followed by Bonferroni post-hoc tests for statistical significance for multiple comparisons. All "center" values from fit curves represent the means of those parameters fit to the individual data, and all data where curves could be unambiguously fit in Graphpad were included, except where sigmoidal fits estimated a halfway point that fell outside the minimal or maximal values of the independent variable used to generate the data. To increase statistical power and avoid error due to multiple comparisons, data for individual constructs under given recording conditions were pooled for analysis, and a single ANOVA statistical test with posthoc analysis performed for any comparisons involving those datasets. For individual constructs, comparison of conductance in Cl⁻ vs I⁻ containing solutions were made using Student's T-test, but no dataset was subjected to T-test more than once. P < 0.01was used as a threshold for significance, and all values in text, and all data presented in summary graphs, are cited or shown as mean \pm S.E.M. n represents the number of individual cells or patches used in recordings, and is displayed either in summary graphs where possible, or in the figure legends where summary graphs are not shown or where display of n in figure would be aesthetically impractical. **, *** and **** represent p < 0.01, p < 0.001 or p < 0.0001 levels of significance, respectively.

Immunocytochemistry—Image files were deconvolved using AutoQuantX3. Deconvolved images were analyzed for colocalization using ImarisColoc. Pearson's correlation coefficients were determined between TMEM16A and WGA, with the membrane of TMEM16A-transfected cells identified as the region of interest. Statistical significance was determined using 1-way ANOVA analysis followed by Bonferroni's posthoc test for significance. P < 0.01 was used as a threshold for significance, and all values in text, and data are shown as mean \pm S.E.M. n represents the number of individual cells used and is displayed in summary graph. **, *** and **** represent p < 0.01, p < 0.001 or p < 0.0001 levels of significance, respectively.

KEY RESOURCES TABLE

A Key Resources Table is attached separately to this submission.

REAGENT or RESOURCE	SOURCE	IDENTIFIER	
Antibodies			
Rhodamine-Wheat germ agglutinin	Vector Laboratories	Z0109	
Bacterial and Virus Strains			
Eschericia coli: DH5a strain, subcloning efficiency	Original: Invitrogen, maintained in house	Cat#18265017	
Biological Samples			
Chemicals, Peptides, and Recombinant Proteins			
Salts, acids, bases, chelators for recording solutions	Sigma-Aldrich	stock	
D-Mannitol	Sigma-Aldrich	stock	
Fluo 8-Potassium salt	AAT-Bioquest	Cat#20189	

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Fluo 8FF-Potassium salt	AAT-Bioquest	Cat#21102
Critical Commercial Assays		
Ca2+ Calibration Buffer Kit #2	Molecular Probes	Cat#3009
Deposited Data		
Experimental Models: Cell Lines		
Human: HEK293 cells	ATCC	RRID: CVCL_0045
Experimental Models: Organisms/Strains		
Oligonucleotides		
Mutagenic primers for PCR	IDT-DNA Technologies	Custom-made, described in STAR Methods
Recombinant DNA		
pEGFP-N1-TMEM16A from <i>Mus</i> <i>musculus</i> (previously generated in our lab)	Tien et al., 2013	N/A
pEGFP-N1-TMEM16B from <i>Mus</i> <i>musculus</i> (previously generated in our lab)	Tien et al., 2013	N/A
Software and Algorithms		•
WEBMAXC-chelation prediction software	Chris Patton, Stanford	http://maxchelator.stanford.edu/webmaxc/webmaxcE.htm
Clustal Omega sequence alignment software	European Bioinformatics Institute	https://www.ebi.ac.uk/Tools/msa/clustalo/
Boxshade multiple alignment software	K. Hofmann and M. Baron	https://www.ch.embnet.org/software/BOX_form.html
The PyMOL Molecular Graphics System, Version 2.0	Schrödinger, LLC	http://www.pymol.org
PClamp10 software suite	Molecular Devices	N/A
Graphpad PRISM 6	Graphpad Software, Inc.	N/A
AutoQuantX3	Bitplane	N/A
ImarisColoc	Bitplane	N/A
Amber 14	Case et al., 2005	http://ambermd.org/
MODELLER 9.15	Sali and Blundell, 1993	https://salilab.org/modeller/9.15/release.html
CHARMM-GUI	Lee et al., 2016	http://www.charmm-gui.org/
matplotlib	J. Hunter	https://matplotlib.org/
Visual molecular dynamics	NIH Center for Macromolecular Modeling & Bioinformatics	http://www.ks.uiuc.edu/Research/vmd/
Other		
Borosilicate thick wall glass 1.5 OD/0.86 ID	Sutter Instrument	BF150-86-10
Borosilicate theta glass 1.5 OD	Sutter Instrument	BT-150-10

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- TMEM16A is an ion channel gated by voltage, external anions and internal calcium
- Alanine scanning mutagenesis permits these gating modes to be analyzed individually
- The 6th transmembrane helix has a critical role in linking TMEM16A gating modes





A. Sample traces from whole cell patch clamped HEK293 cells expressing wild-type TMEM16A and stepped from -120 mV to +150 mV from a holding potential of 0 mV, in the presence of external 140 mM NaCl or NaI (anion indicated on the left) and internal 140 mM NaCl with 0 nM, 400 nM or 1 mM free calcium (indicated above). Red dashed lines indicate 0 nA. B. Current-voltage (IV) relationships for traces in A highlight the different factors affecting TMEM16A gating. TMEM16A is outwardly rectifying at moderate internal Ca²⁺ but loses rectification at high internal Ca²⁺, n = 5, 5 and 7 for 400 nM Ca²⁺ with Cl⁻,

400 nM Ca²⁺ with I⁻ and 1 mM Ca²⁺ with Cl⁻ traces, respectively. C. Steady-state whole cell conductances, derived from the data shown in B, are corrected against the reversal potential and normalized to +150 mV. D. A simplified schematic of TMEM16A gating illustrates a conceptual model of calcium, voltage and anion-dependent modulation of channel conductance. Error bars in B and C represent mean \pm SEM. See also Figures S1 and S2.





A. A representation of mouse TMEM16A TM6 placed above an unbiased sequence alignment of the TM5-6 linker and TM6 of mouse TMEM16A with the other nine mouse TMEM16 family members. Black shading indicates that 70% and grey shading indicates 50-70% of TMEM16 family members have the same amino acid as in TMEM16A, which is denoted in red. The alanine-scanned region, indicated by a green bar, includes residues of particular interest marked by red arrows. B. Cryo-EM structure of TMEM16A wuth TM6 highlighted in green, bound Ca²⁺ ions shown in blue, and the contours of an aqueous pore

shown in dark grey. C. Sample traces of inside-out patches pulled from HEK293 cells expressing the indicated construct and exposed to a series of increasing Ca²⁺ concentrations at +60 mV. Solutions a-f contained 150 nM, 300 nM, 400 nM, 600 nM, 5.5 μ M and 1 mM Ca²⁺, respectively. D. EC₅₀ values for Ca²⁺ concentration-response relationships from curves fit to traces as in panel C, with statistically significant right shifts highlighted in red, and significant left shifts highlighted in blue. Bars for residues that may directly coordinate Ca²⁺ ions are shown with diagonal hatches. "N.C." for constructs with protein expression but no current (see also Figure S2). n for WT, 19; L627A, 6; M628A, 5; E629A, 6; L630A, 6; C631A, 5; I632A, 8; Q633A, 4; L634A, 10; S635A, 9; I636A, 10; I637A, 8; L639A, 6; G640A, 9; K641A, 8; Q642A, 6; I644A, 8; Q645A, 7; N646A, 4; N647A, 5; L648A, 6; F649A, 7; E650A, 4; I651A, 10; G652A, 10; I653A, 11. E. Hill coefficients for fits as in panel D., with significant increased highlighted in red, and significant decrease highlighted in blue. EC₅₀ values and Hill coefficients were compared using one-way ANOVA followed by a Bonferroni post-hoc test for statistical significance. Data in panels D and E are expressed as mean ± SEM.

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A. Image of TMEM16A highlighting the position (in purple) of I637, G640, and Q645 in TM6 (green), with bound Ca²⁺ (blue). B. Current-voltage (IV) relationships for WT, I637A, and Q645A TMEM16A with 0 nM internal Ca²⁺ reveal that the I637A and Q645A mutations remove the requirement for Ca²⁺ for activation (see also Figure S3) as revealed by whole-cell patch clamp, using 10 mV voltage steps from -120 mV to +150 mV from a holding potential of -80 mV. C. Conductance-voltage (GV) relationships for I637A (red)

and Q645A (blue) derived from the IV curves shown in panel B, with filled circles depicting data gathered with chloride and open circles with iodide as the external anion. D. Log_{10} transformation of the data in C with grey dashed line indicating a non-saturating conductance at the highest positive voltages. G-V traces corrected for external aniondependence can then be fit with a two-state Boltzmann relationship (see also Figure S5). E. Sample traces from whole cell patch clamp recording of HEK293 cells expressing wild-type TMEM16A or channels bearing either or both Q645A and I637A mutations exposed to a series of voltage commands from a holding potential of -80 mV with external chloride and 0 nM internal Ca²⁺. Red dashed lines indicate 0 nA. F, G, and H. Graphs depicting normalized GV relationships after subtraction of non-saturating conductance ("Probability-activated" or PA) reveal a conductance with a Boltzmann profile. Solid lines in panel H represent PA with chloride and dashed lines with iodide as the permeant anion. I. Sample traces of inside-out patches exposed to a series of increasing Ca^{2+} concentrations at +60 mV. Solutions a-f contained 150 nM, 300 nM, 400 nM, 600 nM, 5.5 μ M and 1 mM Ca²⁺, respectively. J. Graph derived from data shown in I revealing the change in conductance with increasing Ca^{2+} . The G640A mutation increases sensitivity to Ca^{2+} , but mutating I636 or M638 to Gly restores the normal Ca²⁺ sensitivity. K. Bar graph showing that only the I636G and M638G mutations can rescue the G640A mutation. n was WT, 19; G640A, 9; GA I636G, 11; GA I637G, 4; GA M638G, 5; GA L639G, 4; GA A642G, 5; GA L643G, 6; GA I644G, 4. L and M. PA vs voltage relationships comparing the effect of I637A (L) and Q645A (M) mutations in the G640A background, with solid lines depicting the fits to I637A or Q645A mutations alone. N. A comparison of fitted $V_{1/2}$ s for combinations of I637A, Q645A, and G640A, representing information presented in panels F, G, H, L, and M. Data are expressed as mean \pm SEM and where error bars are invisible, they are contained within the points. See also Figures S3-S5.



Figure 4. Neutralization and charge reversal of K641 in TM6 shifts the voltage-dependence of TMEM16A and is sensitive to the presence of Ca^{2+}

A. Image of TMEM16A highlighting the positions of K641 (cyan) in TM6, with bound Ca²⁺ indicated in blue. B. Sample traces from whole cell patch clamp recording of HEK293 cells expressing TMEM16A with the Q645A mutation (left) or Q645A and K641E mutations (right), stepped from -120 mV to +150 mV in 10 mV steps from a holding potential of -80 mV. Chloride was the permeant anion in each case, with 0 nM internal Ca²⁺ used in the top row and 1 mM Ca²⁺ used in the bottom row. Red dashed lines indicate 0 nA. C and D. Graphs showing the P_A-V relationship for TMEM16A Q645A K641Q (C) and Q645A

K641E (D), with blue lines indicating fits to the Q645A single mutant. K641Q shifts the curve to the right and K641E causes a greater shift. E. P_A -V relationship of Q645A K641E mutant with 1 mM Ca²⁺ resembles that of the Q645A single mutant. Solid and dashed lines indicate the fit to Q645A K641E with 0 mM Ca²⁺ and external chloride (solid) or iodide (dashed). G. Schematic depicting possible changes in TMEM16A conformation after depolarization or Ca²⁺ binding. Position of residues and side chains do not reflect actual structure and are meant to be illustrative. Data are expressed as mean ± SEM and where error bars are invisible, they are contained within the points.

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Figure 6. Depolarization of TMEM16A channels causes a Ca²⁺-dependent stabilization of open conformation and slowing of deactivation kinetics

A. Representative traces of whole-cell patch clamp recordings of WT TMEM16A with 400 nM internal Ca²⁺ with external chloride (top) or iodide (bottom). B. Representative traces of whole-cell patch clamp recordings of WT TMEM16A with external chloride and 200 nM (top) or 1500 nM (bottom) internal Ca²⁺. In both A and B cells were stepped from –120 mV to +150 mV in 10 mV increments from a holding potential of –80 mV. C. Graph of P_A-V for data represented in A and B, showing a progressive left shift in the voltage-dependence upon increasing internal concentration of Ca²⁺. n was 200 Ca²⁺ Cl⁻_I, 5; 400 Ca²⁺ Cl⁻_I, 5; 400 Ca²⁺ Vlage-clamped from a pre-pulse potential of –120 mV (left) or +120 mV (right). Inset shows current immediately after voltage steps. E. "Instantaneous" current amplitudes from wild-type at voltages from –120 to +120 after prepulses to the indicated voltages, normalized to

amplitude at +120 mV, n = 5. F. Q645A with 0 internal Ca^{2+} stepped under voltage clamp from a pre-pulse potential of -120 mV (left) or +120 mV (right). Inset shows current immediately after voltage steps. G. "Instantaneous" current amplitudes from Q645A at voltages from -120 mV to +120 mV after prepulses to the indicated voltages, normalized to amplitude at +120 mV, n = 5 (see also Figure S7). H. Deactivation (top) and reactivation (bottom) time courses following activating and deactivating pulses of increasing duration to +120 mV and -120 mV, respectively, for Q645A with 0 Ca²⁺ (left) and WT with 400 nM internal Ca²⁺ (right). For WT, time constants of activation (cyan) and deactivation (orange) are overlaid with the onset of slow deactivation (top) and the decay of instantaneous current amplitude (bottom), respectively. I. Normalized amplitudes of the slow component of current deactivation compared to total for wild-type TMEM16A following 300 ms activating pulses to +120 mV, varying the concentration of intracellular Ca^{2+} and the species of extracellular permeant anion, as shown J. Slow and fast time constants of current activations for wild-type TMEM16A under the indicated conditions. Also shown is the time constant of onset of a slow component/total amplitude of deactivation during activations of increasing duration $(\tau_{ON} \text{deact}_{s/t})$ as indicated by arrows in panel E (top, right). K. Slow and fast time constants of current deactivations for wild-type TMEM16A under the indicated conditions. Also shown is the time constant of "loss of instantaneous" (L.O.I.) current, representing the recovery of the slowly activating current at +120 mV following deactivations of increasing duration at -120 mV as indicated by arrows in panel H (bottom, right), and the single exponential "fast" τ of deactivation for Q645A with 0 Ca²⁺ and E650A with 1 mM Ca²⁺ (see also Figure S7). n for data depicted in panels J and K was τ_{ON} deact_{s/t}, 7; 200 Ca²⁺ Cl⁻_L 200 Ca²⁺ I⁻_I, 9; 400 Ca²⁺ Cl⁻_I, 13; 400 Ca²⁺ I⁻_I, 8; τ_{OFF} "L.O.I.", 8; Q645A, 6; E650A, 6. Red dashed lines in panels A, B, D and F indicate 0 nA. All pooled data are expressed as mean \pm SEM. See also Figure S7.

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Figure 7. A dynamic interplay of Ca²⁺ and voltage controls TMEM16A gating A. Left, a simplified model depicting changes to TMEM16A. Two circles within the box represent the two Ca²⁺ binding sites, with filled circles indicating occupancy by Ca²⁺. The larger circle represents the pore, with a line indicating a closed pore, half-filled circle for the outward-rectifying open state, and fully filled circle for the 'Ohmic' open state. Right, A sample steady-state I-V distinguishing the two open states for channels with single or double Ca²⁺ occupancy. B. A basic gating scheme for the WT and Q645A mutant TMEM16A under conditions of 0 Ca²⁺_I. Left, the WT channel does not activate in response to voltage changes (up to +200 mV) without Ca²⁺ while the Q645A mutant (right), activates in response to membrane voltage potential changes as though with single Ca²⁺ occupancy, indicated by a gray filled circle. C. The WT channel, under conditions of moderate Ca²⁺ concentration, exists in equilibrium (gray dotted box) with the majority of channels not bound to Ca²⁺ and a small number bound to a single Ca²⁺. After depolarization, the few

channels with single Ca²⁺ occupancy open quickly and allow anion flux. As time passes an increasing number of channels enter the single-Ca²⁺-bound state whereupon they can be opened by depolarization, which also favors the binding of the second Ca²⁺ ion, stabilizing the channel and leading to slowly activating current upon depolarization. With elevated Ca²⁺ levels, the channel can readily bind two Ca²⁺ ions and enter a non-rectifying 'Ohmic' open state. D. A sample trace illustrates different phases of TMEM16A conductance after depolarization under moderate Ca²⁺ levels. A small, rapid outward current (1) results from opening of channels with single Ca²⁺ occupancy while the slower increase (2 and 3) reflect the need for channels to first bind Ca²⁺ before entering an open state. E. A model of TM6 gating in response to Ca²⁺ and voltage changes. Green residues are those in TM6 critical for the Ca²⁺ requirement for activation, while magenta indicates K641. Red circles and red residues indicate the Ca²⁺-binding residues of TM6, 7, and 8. Blue circles are Ca²⁺ ions.