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Physical basis for distinct basal and mechanically-gated activity of the human K⁺ channel TRAAK

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Summary

TRAAK is a mechanosensitive two-pore domain K⁺ (K2P) channel localized to nodes of Ranvier in myelinated neurons. TRAAK deletion in mice results in mechanical and thermal allodynia and gain-of-function mutations cause the human neurodevelopmental disorder FHEIG. TRAAK displays basal and stimulus-gated activities typical of K2Ps, but the mechanistic and structural differences between these modes are unknown. Here, we demonstrate that basal and mechanicallygated openings are distinguished by their conductance, kinetics, and structure. Basal openings are low conductance, short duration, and due to a conductive channel conformation with the interior cavity exposed to the surrounding membrane. Mechanically-gated openings are high conductance, long duration, and due to a channel conformation in which the interior cavity is sealed to the surrounding membrane. Our results explain how dual modes of activity are produced by a single ion channel and provide a basis for the development of state-selective pharmacology with the potential to treat disease.

Graphical Abstract

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Declaration of Interests

The authors declare no competing interests.

Inclusion and Diversity

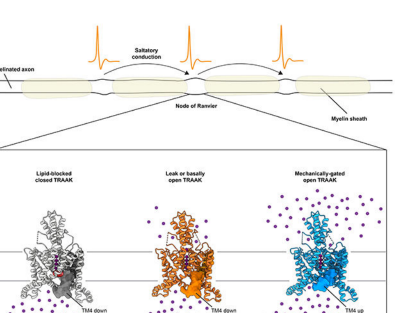
While citing references scientifically relevant for this work, we also actively worked to promote gender balance in our reference list.

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Author Contributions

S.G.B., R.A.R., and B.S. conceived of the project, analyzed data, and wrote the manuscript. R.A.R. performed all aspects of the biochemistry and structural biology. B.S. performed all aspects of the electrophysiology. R.A.R. and B.S. generated constructs for recording. B.L. cultured the hybridoma for antibody purification. S.G.B. supervised the project and secured funding.



EHEIG es

eTOC blurb

The K2P-type ion channel TRAAK involved in action potential propagation generates leak currents and is further activated by mechanical force. Rietmeijer, Sorum et al. study TRAAK gain-of-function mutations including those that cause the human neurodevelopmental FHEIG syndrome and show leak and mechanically-gated openings have distinct structures and electrical properties.

Introduction

Two-pore domain K⁺ (K2P) channels display low basal activity (also referred to as leak, background, fluxgated, or resting activity) that is important for setting and maintaining the resting membrane potential in many cells (Enyedi and Czirják, 2010; Renigunta et al., 2015). In addition, K2Ps are further activated by diverse physical and chemical stimuli including membrane tension, temperature, voltage, pH, signaling lipids, and calcium to regulate cellular electrical excitability. K2Ps are physiologically relevant targets of volatile anesthetics and antidepressants and their dysregulation has been implicated in pathologies including migraine, depression, pulmonary hypertension, atrial fibrillation, diabetes mellitus, Birk-Barel syndrome, and FHEIG syndrome (Facial Dysmorphism, Hypertrichosis, Epilepsy, Intellectual disability, and Gingival outgrowth) (Enyedi and Czirják, 2010; Barel et al., 2008; Bauer et al., 2018; Heurteaux et al., 2006; Lafrenière et al., 2010; Ma et al., 2013; Royal et al., 2019; Schmidt et al., 2015; Vierra et al., 2015).

TRAAK is a mechanosensitive K2P expressed in the nervous system of jawed vertebrates and localized to nodes of Ranvier, the gaps in myelinated axons where the action potential is regenerated during saltatory conduction(Brohawn et al., 2019; Kanda et al.,

2019; Maingret et al., 1999). TRAAK accounts for ~25% of nodal background K⁺ conductance and maintains the resting membrane potential and voltage gated-Na⁺ channel availability required for high frequency spiking (Brohawn et al., 2019; Kanda et al., 2019)). Altering TRAAK activity in animals has physiological consequences; TRAAK knockout mice display mechanical and thermal allodynia and mechanical hyperalgesia while gain-of-function mutations in humans (TRAAK_{A198E} and TRAAK_{A270P}) underlie the neurodevelopmental disorder FHEIG (Bauer et al., 2018; Noël et al., 2009).

TRAAK basal activity can be activated ~100-fold by increased membrane tension (Brohawn et al., 2014a, 2014b; Sorum et al., 2021). A model for mechanical gating primarily involves conformational changes in transmembrane helix 4 (TM4) (Brohawn et al., 2014b). In this model, nonconductive (closed) TRAAK adopts a TM4 "down" conformation and conductive (open) TRAAK adopts a TM4 "up" conformation. When TM4 is down, a lipid acyl chain can enter the channel cavity below the selectivity filter through membrane facing openings and sterically block the conduction path. Movement of TM4 up seals the membrane facing openings, prevents lipid access, and permits ion conduction. Structures of these two conformations provide a biophysical explanation for mechanosensitivity: shape changes upon opening, including an expansion in cross sectional area and increase in cylindricity, are energetically favored by membrane tension.

Notably, single channel recordings of TRAAK, and the closely related mechanosensitive K2P channels TREK1 and TREK2, suggest the presence of multiple distinct open and closed states (Bang et al., 2000; Kang et al., 2005; Maingret et al., 1999; Patel et al., 1998; Sorum et al., 2021; Xian Tao Li et al., 2006). In addition, alternative models for gating K2Ps posit closures involving C-type gating at the selectivity filter (Bagriantsev et al., 2011; Chatelain et al., 2012; Cohen et al., 2008; Li et al., 2020; Lolicato et al., 2020; Niemeyer et al., 2010; Piechotta et al., 2011; Sandoz et al., 2009; Schewe et al., 2016), closures involving dewetting of the channel cavity (Aryal et al., 2014, 2015), and openings that do not involve upward movement of TM4 (Lolicato et al., 2014; McClenaghan et al., 2016). For TRAAK, and other K2Ps, basally open and stimulus-activated currents display different voltage dependence (Schewe et al., 2016), consistent with structurally distinct open states. Here, we show basal and mechanically-gated TRAAK open states are distinct in their conductance, kinetics, and structure and present a model for channel gating that associates conformational and functional states.

Results

Basal and mechanically-gated TRAAK openings are physically distinct

A macroscopic recording from a patch containing wild-type TRAAK (TRAAK_{WT}) illustrates basal activity and robust activation by membrane tension (generated by negative pressure applied through the patch pipette) that are characteristic of the channel (Fig. 1A,B). We performed single channel recordings to determine whether basally open and mechanically activated TRAAK have different biophysical characteristics consistent with distinct channel open states. By varying applied pressure over approximately an hour of single channel records, we sampled a wide range of TRAAK_{WT} activity (0.01 open probability (P_O) 0.98) (Fig. 1C–F, S1A,B, Table S1). We analyzed records in three

bins according to P_O (Fig. 1C). We found that in addition to a simple increase in P_O , mechanically activated TRAAK openings are distinct from basal openings in two ways.

First, mechanical force favors higher conductance openings (Figs. 1G–I, S2). Two conductance states are observed in TRAAK_{WT} a low conductance opening with a unitary current of ~1 pA and a high conductance opening with a unitary current of ~2 pA. In unstretched patches at low P_O (0.01 P_O 0.07), the two conductance states are observed at roughly similar frequency (Fig. 1G,M). At intermediate P_O (mid P_O , 0.38 P_O 0.69), more high conductance openings are observed (Fig. 1H,M). In records mechanically activated to high P_O (0.93 P_O 0.98), almost all openings are high conductance (Fig. 1I,M). Notably, unitary currents of these openings do not change significantly as a function of P_O , but the relative proportion of each conductance state does (Fig. 1G–J,M).

Second, mechanical force favors longer duration openings. Two kinetically distinct open states are observed in TRAAK_{WT} records: a short duration (~1 ms) opening and a long duration (~3 ms or longer) opening (Figs. 1K, S3). In unstretched patches at low P_O , almost all openings are short duration (Fig. 1M). In contrast, when mechanically activated to high P_O , almost all openings are long duration. At intermediate P_O , both short and long duration openings are observed. Open duration and unitary current are positively correlated: low conductance openings have a narrow duration distribution around 1 ms while high conductance openings have a wider duration distribution 2 ms, with longer openings observed at higher P_O (Fig. 1M).

In addition, mechanical force favors shorter duration closures. TRAAK_{WT} accesses two kinetically distinct closed states: a short duration (~0.5 ms) closure and a long duration (~3 ms or longer) closure (Figs. 1L, S4). Short duration closures of comparable length are observed in all records. However, long duration closures are only observed in low and intermediate P_O records and their prevalence and duration decrease as P_O increases (Figs. 1L, S3).

Single channel TRAAK_{WT} records can be well fit with a simple linear kinetic model relating the two closed states (long duration (C1) and short duration (C2)) and two open states (short duration/low conductance (O1) and long duration/high conductance (O2)), though additional transitions between states are likely possible (Fig. 1N, Table S1)(Sorum et al., 2021). The distribution of states and the duration of C1 and O2 are dependent on membrane tension. When mechanically activated to high P_O, long duration closures and short duration/low conductance openings are almost never observed, so data can be fit with a two-state equilibrium model between C2 and O2 states (Figs. 1I,K-M, S2-4, Table S1). At intermediate P_0 , the duration of long C1 closures is shorter than those at low P_0 and the duration of O2 openings is shorter than those at high P_O (Figs. 1H, K–M, S2–4, Table S1). We conclude that mechanically activated openings correspond to the O2 state while basal openings correspond to the O1 state. We speculate weak mechanical activation by resting membrane tension in the patch results in low Po O2 openings with a conductance distinct from O1 openings (Brohawn et al., 2014b; Opsahl and Webb, 1994). However, the duration of O1 and O2 openings at low P_{O} is similarly short and therefore indistinguishable kinetically (Fig. 1D,K).

We then asked whether we could identify the structural basis for these two distinct TRAAK open states. We reasoned that gain-of-function mutations could differentially stabilize particular open states and investigated three point mutants reported to dramatically increase channel activity: TRAAK_{A198E}, TRAAK_{A270P}, and TRAAK_{G158D}. TRAAK_{A198E} and TRAAK_{A270P} were recently identified in human patients with the neurological and developmental disorder FHEIG while TRAAK_{G158D} was identified as a pan-K2P gain-of-function mutation (Bauer et al., 2018; Ben Soussia et al., 2019).

Mechanical force and FHEIG mutations promote a TM4 up open state

We first consider the FHEIG mutations TRAAK_{A198E} and TRAAK_{A270P} (Figs. 2A, 3A). Macroscopic recordings of TRAAK_{A198E} or TRAAK_{A270P} channels display currents that are only subtly activated by mechanical force, consistent with the initial characterization of these mutations (Figs. 2B,C, 3B,C) (Bauer et al., 2018). These recordings are representative of the maximum level of channel activation observed before patch rupture (Figure S5A–C). Single channel records reveal a simple explanation for reduced mechanical activation of TRAAK_{A198E} and TRAAK_{A270P} relative to TRAAK_{WT}. TRAAK_{A198E} and TRAAK_{A270P} open probability is nearly one at rest and mechanical force does not change channel conductance, so stimulation can only slightly increase channel activity (Figs. 2D–G, 3D–G, S1C,D).

TRAAK_{A198E} and TRAAK_{A270P} single channel behavior closely matches TRAAK_{WT} when mechanically activated to high P_O. TRAAK_{A198E} and TRAAK_{A270P} openings are nearly all high conductance (with a unitary current ~2 pA) and long duration (~3 ms or longer), while closures are short duration (~0.5 ms) (Figs. 2F–J, 3F–J). The modest mechanical activation of TRAAK_{A198E} and TRAAK_{A270P} is due to an increase in long duration openings without significantly changing closed duration.

We determined crystal structures of TRAAKA198E and TRAAKA270P in the presence of K⁺ and in complex with a mouse monoclonal antibody Fab fragment (Figs. 2A, 3A, S6A-F, Table S2). Data were anisotropic and extended along the three crystallographic axes to 2.3, 2.4, and 3.3 Å resolution for TRAAK_{A198E} and 2.7, 2.9, and 3.9 Å resolution for TRAAK_{A270P}. As previously reported for this crystal form (Brohawn et al., 2013, 2014b), the conformation of one side of the TRAAK dimer is restricted because it is involved in forming contacts that propagate the lattice (Fig. S6G,H). We therefore focus our discussion on the opposing side of the TRAAK dimer (including TM4 from protomer B), which is conformationally unrestricted. In the same crystallization conditions, TRAAK_{WT} adopts a TM4 down nonconductive conformation; the membrane-facing lateral opening above TM4 permits hydrophobic acyl chains to access the channel cavity and block ion passage (Fig. 2K) (Brohawn et al., 2014b). TRAAKA198E and TRAAKA270P, however, adopt TM4 up conformations (Figs. 2L, 3K). Movement of TM4 up seals the membrane-facing lateral openings, preventing acyl chain access to the cavity and creating an unobstructed path for ion movement through the channel. This TM4 up conformation is similar to a TRAAK_{WT} TM4 up conductive conformation captured in the presence of a small molecule activator trichloroethanol (overall r.m.s.d = 0.2 Å between TRAAK_{A198E} or TRAAK_{A270P} and TRAAK_{WT} (PDB 4WFE), Fig. S7A,B,E) (Brohawn et al., 2014b). As observed in the

TRAAK_{WT} TM4 up conductive structure, density consistent with a K^+ ion is present in the cavities of TRAAK_{A198E} and TRAAK_{A270P} (Figs. 3L, S6I).

To confirm that the TRAAK_{A198E} TM4 up structure represents an open state, we determined a second structure in the presence of Tl⁺ (to 2.8, 2.8, and 3.8 Å resolution along the three crystallographic axes, Table S2) and used anomalous diffraction of Tl⁺ ions to unambiguously identify ion binding sites. Indeed, a Tl⁺ ion is identified in the channel cavity in addition to four sites in the selectivity filter (S1-S4) and one on the extracellular mouth of the pore (S0), consistent with a conductive conformation (Fig. 2M).

How do TRAAKA198E and TRAAKA270P promote TM4 up open states? Comparing TRAAK_{A198E} and TRAAK_{WT} suggests two mechanisms in this mutant (Fig. 2N). The first is a steric relay from A198E through F201 and Y271 that favors TM4 up. In TRAAK_{WT}, the TM4 down conformation involves movement of Y271 1 ~ towards the cytoplasm relative to TM4 up. This requires the intracellular TM2-TM3 linkage to rotate 15° away from TM4 to prevent a clash between Y271 on TM4 and F201 on TM3. In TRAAKA198E, TM2-TM3 is similarly rotated out, but A198E pushes F201 closer to TM4 where it would clash with Y271 in a TM4 down conformation. A second possibility is that A198E causes local thinning of the lipid bilayer to favor TM4 up. While A198 is predicted to be embedded in the membrane inner leaflet facing hydrophobic residues on TM4 down, the negatively charged carboxylic acid of A198E likely favors hydration of this pocket, promoting TM4 up to maintain nonpolar interactions between its hydrophobic residues and lipid acyl chains (Brooks, B. R. and Brooks, III, C. L. and Mackerell, Jr. et al., 2009; Jo et al., 2008). Analysis of two additional mutants at this position are consistent with both mechanisms contributing to channel activation, with charge playing a larger role. A smaller negatively charged residue in TRAAKA198D activates the channel to a similar degree as TRAAKA198E based on maximum observed mechanical stimulation (Fig. S5E-G). In contrast, a bulky uncharged residue in TRAAKA198W still activates the channel relative to wild-type, but approximately two-fold less than TRAAK_{A198E} (Fig. S5A,E–G).

Comparing TRAAK_{A270P} and TRAAK_{WT} provides a structural explanation for channel activation: a proline at position 270 produces a kink in TM4 that closely approximates the TRAAK_{WT} TM4 up conformation (Fig 3M). In TRAAK_{WT}, movement of TM4 up depends on a hinge at a conserved glycine G268. The A270P mutation instead disrupts backbone hydrogen bonding in TM4 to create a ~27° bend and ~9.5 Å upward movement of TM4 relative to TRAAK_{WT} TM4 down.

We conclude that the TRAAK_{A198E}, TRAAK_{A270P} and TRAAK_{WT} TM4 up structures correspond to the high conductance/long duration mechanically activated open state O2. The following lines of evidence support this assignment: (i) TRAAK_{A198E} and TRAAK_{A270P} have an open probability 0.9 (Figs. 2F,H; 3F,H), (ii) the structures are similar to a TRAAK_{WT} mechanically activated conductive conformation (Figs. 2K,L; 3K), (iii) the structures show unobstructed conduction paths with ions in the channel cavities (Figs. 2M, 3L, S6I), (iv) the predominant TRAAK_{A198E} and TRAAK_{A270P} unitary current is indistinguishable from TRAAK_{WT} O2 (Figs. 1I, 2G, 3G), and (v) the mean duration of

the TRAAK_{A198E} and TRAAK_{A270P} open states are indistinguishable from TRAAK_{WT} O2 (Figs. 1K, 2I, 3I, S3).

A TM4 down open state promoted by a pan-K2P activating mutation underlies basal activity

We next consider TRAAK_{G158D} (Fig. 4A). In macroscopic recordings, mechanical force activates TRAAK_{G158D} a maximum of ~2-fold; more than TRAAK_{A198E} and TRAAK_{A270P}, but less than TRAAK_{WT}(Fig. 4B–D, S5A–D). We note that recordings and quantification of mechanical activation in macroscopic patches (Fig. 1A,B; 2B,C; 3B,C; 4B–D) correspond to maximum mechanical activation observed prior to membrane rupture. Pressure values required for maximal activation vary between patches and are not directly comparable (Fig. S5A–D) due to variation in patch geometry that results in different relationships between applied pressure and membrane tension generated that opens the channel (Moe and Blount, 2005). Single channel analysis shows that, similar to TRAAK_{WT}, mechanical activation of TRAAK_{G158D} involves an increase in both open probability and conductance (Fig. 4E–H, S1E–I). However, the resting open probability of TRAAK_{G158D} (P_O ~0.7) is much higher than TRAAK_{WT} (P_O ~0.04) so mechanical stimulation of TRAAK_{G158D} are low conductance with a unitary current ~1 pA while mechanically activated openings are higher conductance with a unitary current ~2 pA (Figs. 4G–I, S2).

Kinetic analysis shows that TRAAK_{G158D} closely resembles TRAAK_{WT} except that the long duration closed state C1 is absent in the mutant (Figs. 4J, S3, S4). TRAAK_{G158D} accesses only a single short closed state (Fig. 4K). Like TRAAK_{WT}, TRAAK_{G158D} has two open states, one short duration/low conductance (~1 ms / 1 pA) and one long duration/high conductance (~3 ms or longer / 2 pA) (Fig. 4I,J). Mechanical stimulation favors longer duration/high conductance openings. (Figs. 4L, S2, S3).

We determined the crystal structure of TRAAK_{G158D} (to 2.9, 3.1, and 3.7 Å resolution along the three crystallographic axes) in the presence of K⁺ (Fig. 4A, Table S2). TRAAK_{G158D} adopts a TM4 down conformation like TRAAK_{WT} crystallized under the same conditions (overall r.m.s.d = 0.2 Å between TRAAK_{G158D} and TRAAK_{WT} (PDB 4WFE))(Fig. 4M). However, spherical shaped density in the TRAAK_{G158D} cavity is consistent with a K⁺ ion rather than the hydrophobic acyl chain present in the TRAAK_{WT} TM4 down structure (Fig. 4N). Attempts to use Tl⁺ anomalous diffraction to unambiguously determine the identity of the spherical shaped density in the TRAAK_{G158D} cavity were inconclusive due to limited size of TRAAK_{G158D} crystals grown in Tl⁺ and correspondingly weak overall anomalous signal. However, the shape of the cavity density and high resting open probability of the mutant channel (P_O ~0.7) are most consistent with the TRAAK_{G158D} structure representing a TM4 down open state.

TRAAK_{G158D} likely promotes a TM4 down open state through a simple electrostatic consequence of the mutation (Fig. 4O, a comparison to TRAAK_{WT} is shown in Fig. S7M–P). G158D projects into the channel cavity ~5 Å underneath the selectivity filter and increases hydrophilicity of the ion conduction path close to where acyl chains are observed in nonconductive TM4 down TRAAK_{WT} structure. The electronegative cavity

of TRAAK_{G158D} is expected to disfavor acyl chain access and channel block, instead promoting ion occupancy in the cavity and a conductive open state. The increased cavity polarity may contribute to the observed difference in voltage-dependance between TRAAK_{WT} and TRAAK_{G158D} (Fig. 1A,B; 4B,C).

We conclude that the TM4 down open state captured in the TRAAKG158D structure corresponds to the low conductance/short duration basal open state O1 in TRAAK_{WT}. The following data support this assignment: (i) TRAAK_{G158D} has a high resting open probability (Fig. 4E), (ii) a K⁺ ion rather than an acyl chain is observed in the TRAAK_{G158D} cavity (Fig. 4N), (iii) unitary current of the TRAAK_{G158D} basal open state is indistinguishable from TRAAK_{WT} O1 (Fig. 4I), (iv) mean dwell time of the TRAAK_{G158D} short duration opening is statistically indistinguishable from TRAAK_{WT} O1 (Fig. 4J), and (v) like TRAAK_{WT}, the basal low conductance/short duration TRAAK_{G158D} O1-like open state can be mechanically activated to a high conductance/long duration O2-like open state (Fig. 4F–J). Consistent with (v), TRAAK_{G158D} can adopt a TM4 up-like conformation (observed on the side of the channel involved in forming crystal contacts) and TRAAK_{G158D} TM4 up is expected to be energetically favored in the presence of membrane tension relative to $TRAAK_{G158D}$ TM4 down because it has an increased cross-sectional area (Brohawn et al., 2014b) (Fig. S7C). The energetic difference between TM4 down and TM4 up states may be different in the mutant; trichloroethanol has no effect on TRAAK_{G158D} current in whole cell recordings under conditions that activate TRAAK_{WT} (Fig. S5H–J).

We further conclude that the TM4 down lipid-blocked closed state observed in the TRAAK_{WT} structures corresponds to the long duration closed state C1. Analogous long duration closures are not observed in in TRAAK_{G158D}, TRAAKA_{A198E}, or TRAAK_{A270P} (Figs. 1L, 2J, 3J, 4K). In TRAAK_{A198E} and TRAAK_{A270P}, this is likely because TM4s are favored in an up conformation that seals lateral membrane openings, preventing lipid access to the cavity (Figs. 2L, 3K). In TRAAK_{G158D}, this is likely because the mutation increases polarity of the channel cavity to disfavor occupancy of hydrophobic acyl chains (Fig. 4O).

Discussion

Together, these data support a model for TRAAK gating shown in Figure 5 that maps structural conformations to a linear four-state kinetic model derived from single-channel records (Fig. 1N). Long duration closures (C1) correspond to the TM4 down, lipid blocked conformation captured in TRAAK_{WT} structures (Brohawn et al., 2014b). Low conductance/short duration basal openings (O1) correspond to the TM4 down, conductive conformation captured in the TRAAK_{G158D} structure (Fig. 4). High conductance/long duration mechanically activated openings (O2) correspond to the TM4 up, conductive conformation captured in structures of TRAAK_{A198E}, TRAAK_{A270P}, and TRAAK_{WT} with activating small molecules (Fig. 2,3). Structures of two other gain-of-function mutations, TRAAK_{G124I} and TRAAK_{W262S}, have been previously reported (Lolicato et al., 2014), although their open probability, single channel behavior, and cavity ion occupancy is not known. However, both adopt TM4 down-like conformations and may promote O1 like openings similarly to TRAAK_{G158D}.

TRAAK_{WT} under basal conditions transitions primarily between C1, O1, and C2 states. The channel predominantly adopts a lipid-blocked TM4 down closed state C1, but spontaneous delipidation in the absence of protein conformational changes results in short O1 openings until lipid rebinding. These O1 openings underlie basal activity. Mechanical force favors the TM4 up open state O2 because it involves shape changes (relative to TM4 down) that are energetically favored in the presence of membrane tension, including an increase in cross-sectional area and cylindricity(Brohawn et al., 2014b). Mechanical force disfavors the TM4 down states C1 and O1 because the relatively smaller cross-sectional area and more wedge-shaped structure is disfavored in the presence of membrane tension. Higher membrane tension increasingly favors promotes and stabilizes TM4 up, resulting in longer duration O2 openings and correspondingly shorter duration C1 closures.

The gain of function mutations TRAAK_{A198E} and TRAAK_{A270P} mimic mechanically activated TRAAK_{WT} by promoting a TM4 up conductive conformation. As a consequence, TRAAK_{A198E} and TRAAK_{A270P}, like mechanically activated TRAAK_{WT}, primarily transition between O2 and C2 states. Gain of function mutant TRAAK_{G158D} mimics basally open TRAAK_{WT}, promoting a conductive TM4 down conformation by disfavoring lipid entry and channel block at low membrane tension. As a consequence, TRAAK_{G158D} primarily transitions between O1, C2, and O2 states. Like TRAAK_{WT}, mechanical stimulation of TRAAK_{G158D} disfavors O1 to favor transitions between O2 and C2 states.

Differences in the chemical and geometric properties of the conduction path likely explain differences in properties of basal O1 and mechanically-activated O2 openings. The cavity diameter is larger and more polar near the cytoplasmic entrance to the channel in the TM4 up O2 open state compared to a TM4 down O1 open state (Fig. S7M–P). These differences are expected to result in lower access resistance to the cavity, which would explain the higher conductance of O2 compared to O1 openings.

The short duration closure (C2) does not yet have a defined structural correlate. We observe C2-like closures of similar duration in all TRAAK variants analyzed (Figs. 1L, 2J, 3J, 4K), suggesting they may occur independently of TM4 position or cavity environment. One possibility is that C2 closures represent gating at the selectivity filter. While gating conformational changes at the selectivity filter have not been observed in TRAAK, they have been hypothesized from functional data and visualized in other K2Ps including the mechanosensitive TREK1 in low [K⁺] and pH-gated TASK2 in physiological [K⁺] and low pH (Li et al., 2020; Lolicato et al., 2020). Alternatively, C2 closures could correspond to cavity dewetting that has been proposed to occur based on molecular dynamics simulations (Aryal et al., 2014, 2015) or other, yet to be visualized, conformational rearrangements.

Notably, some evidence suggests that the physical basis for distinct modes of TRAAK activity shown here could be shared among K2P channels. Differences between basal and stimulus-activated openings, including their kinetics and voltage dependence (Schewe et al., 2016), have been reported for other K2Ps, suggesting that physically distinct open states could similarly underlie different modes of channel activity (Bang et al., 2000; Gnatenco et al., 2002; Kang et al., 2005; Maingret et al., 1999; Sorum et al., 2021). Mutations analogous to TRAAK_{G158D} were found to activate all K2P channels, with the extent of

activation correlated with the polarity of the introduced amino acid change (Ben Soussia et al., 2019). Membrane-facing openings have been observed in TM4-down-like structures of TREK2 (Dong et al., 2015), TWIK1 (Miller and Long, 2012), TASK1 (Rödström et al., 2020), and TASK2 (Li et al., 2020) in addition to TRAAK (Brohawn et al., 2012) (Fig. S7D–L). Acyl chains that could block conduction were modeled in TWIK1, TREK2, and TRAAK structures (Brohawn et al., 2012; Dong et al., 2015; Miller and Long, 2012), though their absence in other K2Ps may be due to limited resolution of, or partial occupancy in, structures determined to date. This raises the possibility that lipid-mediated block could be more common among K2Ps than currently appreciated. Like TRAAK, basal activity of other K2Ps could result from transient unbinding of lipid from channel cavities and ion passage through an unobstructed conduction path, while stimulus gating could involve conformational changes that prevent lipid entry into the channel entirely (Brohawn et al., 2012, 2014b; Dong et al., 2015; Li et al., 2020; Miller and Long, 2012; Rödström et al., 2020).

STAR Methods

Resource availability

Lead contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Stephen Brohawn (sbrohawn@berkeley.edu).

Materials availability—Plasmids used in this study are available upon request.

Data and code availability—Single channel analysis code written in Python for this paper is available on Github: (https://github.com/BrohawnLab/ElectrophysiologyScripts/tree/singles0).

The X-ray crystallographic coordinates and structure factors for TRAAK_{A198E} in K⁺ (7LJ5), TRAAK_{A198E} in TI⁺ (7LJA), TRAAK_{A270P} in K⁺ (7LJ4), and TRAAK_{G158D} (7LJB) are available at the Protein Data Bank.

Experimental model and subject details

Cell lines—Oocytes were harvested from female *Xenopus laevis* frogs obtained from Nasco and housed at the University of California, Berkeley in the Office for Animal Care and Use (OLAC) facilities. The use of oocytes was approved by the Animal Use and Care Committee at the University of California, Berkeley under AUP-2019-11-12743.

Method details

Electrophysiology—Full length hsTRAAK (Uniprot Q9NYG8-2) was codon optimized for eukaryotic expression, synthesized (Genewiz), and cloned into a modified pGEMHE vector using Xhol and EcoRI restriction sites such that the mRNA transcript encodes hsTRAAK 1-419 with an additional "SNS" at the C-terminus. Point mutants G158D, A198E, and A270P were generated by inverse PCR of the parent construct and verified by Sanger sequencing. cRNA was transcribed from linearized plasmids *in vitro* using T7

polymerase, and 0.1–10 ng cRNA was injected into *Xenopus laevis* oocytes extracted from anaesthetized frogs.

Currents were recorded at 25°C from inside-out patches excised from oocytes 1–5 days after mRNA injection. Pipette solution contained (in mM) 15 KCl, 135NaCl, $2MgCl_2$, 10 HEPES, pH = 7.4 with NaOH. The solution in the bath contained 150 KCl, 2 MgCl₂, 10 HEPES, 1 EGTA, pH = 7.1 with KOH. All single channel records were made at a holding voltage of 0 mV and presented in physiological convention.

Currents were recorded using an Axopatch 200B Patch Clamp amplifier at a bandwidth of 1 kHz and digitized with an Axon Digidata 1550B at 50 to 500 kHz. To maintain the integrity of brief TRAAK dwell times, records were not further filtered unless explicitly stated. Baseline correction was performed manually based on the observable closed state for each recording; since there was no solution exchange in our experiments, no associated large drifts in baseline were observed. Portions of TRAAKWT records were assigned as low, mid, or high Po for analysis. Low Po was observed prior to pressure application, while mid and high P_O behavior was observed after pressure application. Portions of mutant records during pressure application were analyzed separately from mutant records prior to pressure application. High POTRAAK_{WT} and mutant records with pressure were made with the highest achievable pressure prior to patch rupture. Single channel open-close transitions were idealized by half-amplitude threshold crossing and dwell time event lists were generated for each baseline corrected record. Dwell time histograms were generated from the dwell time event lists using custom built software (Sorum et al., 2015), while the fits for these histograms were generated in GraphPad Prism version 9.0 using single, double, or triple Gaussian fits.

For unitary current analysis, event lists for TRAAK_{WT} low P_O, TRAAK_{WT} mid P_O, TRAAK_{G158D}, and TRAAK_{G158D} with pressure were first generated in Clampfit 10.7, applying a 0.8 pA event detection cutoff. Currents were filtered to 10 kHz and were analyzed with a custom script in Python 3.7 using the pyABF module (Harden, 2019). Each point in the filtered record was annotated as closed (C1/C2), open low conductance (O1), or open high conductance (O2) using the events list. To account for the sharp detection cutoff, openings were padded by an additional point on either side of the cutoff (these points corresponded to 100 us of ~0.5-1 pA current during openings and closings). Open only event lists were then generated by removing all closed points. Open event only lists were binned by 0.2 pA in current and 0.2 ms in time to generate bubble plots relating unitary current and mean open duration for TRAAK_{WT} and TRAAK_{G158D}. Open only event lists were also used to generate open event current histograms to distinguish relatively rare low conductance openings from prevalent closed events. Mean unitary currents for TRAAK_{WT} low Po and $TRAAK_{WT}$ mid P_O were derived from Gaussian fits to these open event current histograms. Mean unitary currents for TRAAK_{WT} high Po and all mutants were derived from Gaussian fits to square root total event current histograms in order to distinguish conductance states when one state is present at higher frequency.

For TEVC recordings, the bath was ND96 with or without 5 mM trichloroethanol. Pipettes were filled 3M KCl and had resistances of 0.5-2 M Ω . Currents induced by voltage steps

were first recorded in ND96 bath solution. Next, half the bath solution was removed and replaced by the same volume of ND96 with trichloroethanol to ensure the bath volume remained constant between recordings.

Protein expression and purification—Human TRAAK (UniProt Q9NYG8-2) was cloned for expression in *Pichia pastoris* cells as previously described (Brohawn et al., 2012) with modifications described here. The construct used for purification included an additional 26 amino acid N-terminal sequence compared to Q9NYG8-1 that improved heterologous expression. The final construct is C-terminally truncated by 119 amino acids, incorporates two mutations to remove N-linked glycosylation sites (N104Q/N108Q), and is expressed as a C-terminal PreScission protease-cleavable EGFP-10xHis fusion protein. As a result, there is an additional amino acid sequence of "SNSLEVLFQ" at the C-terminus of the final purified protein after protease cleavage. Recombinant *Pichia* were grown in a 4 liter fermenter and harvested 48-72h after induction with methanol.

30-120g of frozen Pichia cells expressing TRAAK were disrupted by milling (Retsch model MM301) 5 times for 3 minutes at 25 Hz. All subsequent purification steps were carried out at 4 °C. Milled cells were resuspended in buffer A (in mM) 50 Tris pH 8.0, 150 KCl, 1 EDTA 0.1 mg/mL DNase1, 1 mg/mL pepstatin, 1 mg/mL leupeptin, 1 mg/mL aprotinin, 10 mg/mL soy trypsin inhibitor, 1 mM benzamidine, 100 µM AEBSF, 1 µM E-64, and 1 mM phenylmethysulfonyl fluoride added immediately before use) at a ratio of 1 g cell pellet per 4 mL lysis buffer and sonicated for 16 minutes with a duty cycle of 15 seconds of sonication per minute. The solution was ultracentrifuged at 150,000 xg for 1 hour at $4 \,^{\circ}$ C. Pellets were transferred to a Dounce homogenizer in buffer B (buffer A + 60 mM DM). Following homogenization, solutions were stirred for 3 hours at 4 °C followed by centrifugation at 33,000g for 45 minutes. Anti-GFP nanobody resin (1 mg purified anti-GFP nanobody conjugated to 1 mL resin) was washed in Buffer B and added to the supernatant at a ratio of 1 mL resin / 15 g Pichia cells. The solution was gently stirred for 3 hours. Resin was collected on a column and washed in 15 column volumes (CV) of Buffer C (buffer A +6 mM DM+ 150 mM KCl), followed by 2 CV Buffer D (buffer A +6 mM DM). The resin was resuspended in ~3 CV of Buffer D with 1 mg purified Precission protease and gently rocked in column overnight. Cleaved TRAAK was eluted in ~4 CV of Buffer D, concentrated (50 kDa MWCO), and applied to a Superdex 200 SEC column (GE Healthcare) equilibrated in Buffer E (20 mM Tris pH 8.0, 150 mM KCl, 1 mM EDTA, 4 mM DM). Peak fractions were pooled and concentrated to 200-300 µL for incubation with Fab 13E9 and subsequently applied to a Superdex 200 column (GE Healthcare) equilibrated in Buffer E. Fab 13E9 was prepared as described previously(Brohawn et al., 2014b). TRAAK-Fab complexes were pooled and concentrated to 25-33 mg/mL for hanging drop crystallization. For crystallization in Tl⁺, all purification steps were conduction as with KCl, with two exceptions: KCl is substituted for KNO3 for all buffers except Buffer E, where KCl is substituted with TINO₃.

Protein Crystallization—Crystals were grown in drops of $0.125-0.250 \mu$ L protein added to an equal volume of reservoir, in hanging drops over a 100 μ L reservoir at 4 °C. Reservoir for each mutant in KCl was 50mM Tris pH 8.8, 64-200 mM CaCl₂, 27–33%

(vol/vol) PEG400. Reservoir for each mutant in TINO₃ was 50 mM Tris pH 8.8, 64-200 mM Ca(NO3)₂, 27–33%(vol/vol)PEG400. Lower Calcium concentrations were observed to correspond to increased nucleation and rapid growth of large, well-ordered crystals. Crystals grew to ~100 μ m x 100 μ m x 200 μ m in 1–5 weeks. TRAAK A270P crystals rarely achieved sizes larger than 70 μ m x 70 μ m x 100 μ m after 5 weeks.

For cryoprotection, an approximately equal volume of mother liquor supplemented to be 30%(vol/vol) PEG400 was added to one side of the drop and crystals were moved through this solution with a cryoloop before being plunged into liquid nitrogen.

X-ray Data Collection, Model Building & Refinement—Data were collected at beamline 8.3.1 at the ALS, or 24-IDC & 24-IDE at the APS. Thallium-containing crystals were collected at 12680 eV, and Potassium-containing crystals were collected at 12663 eV. Were rotated 360 degrees, and data were collected in 0.20 degree wedges with a 0.25 second exposure time per wedge. Datasets were processed as individual wedges in XDS (Kabsch, 2010; Kabsch et al., 2010). Some datasets were truncated to eliminate redundant data displaying beam-induced damage prior to scaling with XSCALE, and merging with XDSCONV (Kabsch et al., 2010). Due to the anisotropy of the data, elliptically truncated resolution cutoffs generated by STARANISO (Tickle et al., 2018) were used to increased overall resolution of the maps. Structures were solved by molecular replacement using Phaser (McCoy et al., 2007) with an input model of nonconductive hsTRAAK for G158D (PDB ID 4WFF) and conductive hsTRAAK for A198E and A270P mutants (PDB ID 4WFE). Structure refinement was carried out in Coot (Emsley and Cowtan, 2004) and Refmac5 (Murshudov et al., 2011). Jelly body and automatically generated local NCS constraints were used throughout refinement. For the final round of refinement, three TLS groups per protein chain were incorporated. Molprobity was used to assess model geometry in later stages of refinement (Chen et al., 2010).

Ion occupancy in the conduction axis was determined by calculating either a model phased anomalous difference map for A198E crystals grown in Thallium, or a Polder map for G158D and A270P crystals grown Potassium. In all maps, density in the filter fits well to ions occupying the S1, 2, 3, and 4 positions. Similarly, densities above the filter (S0) are well fit with one or two ions. Density in the cavity below the filter is also fit with an ion. All ion occupancies are set to 1.

The angle and orientation of transmembrane helix 4 in each model was compared using UCSF Chimera (Pettersen et al., 2004). Cross-sectional area calculations were performed with CHARMM (Brooks, B. R. and Brooks, III, C. L. and Mackerell, Jr. et al., 2009; Jo et al., 2007, 2008). The area calculation used a surface calculated with 1.4 Å added to the van der Waals radii of protein atoms, and a probe radius of 3.5 Å to approximate lipid-accessible surface area. A water cylinder with a 6 Å radius was used to fill the cavity of TRAAK channels to exclude its contribution from the calculation. All calculations are made with channels containing symmetric protomers of the B chain, which does not make a contact in the crystal. Symmetric molecules were generated by rotation of the B protomer about the conduction axis, using pore helix and selectivity filter residues 119-133 and 228-242. Pore radii and pore-lining residue hydropathy were calculated with CHAP (Klesse et al., 2019)

using a pathway lining residue margin value of 0.55 and a hydrophobicity kernel bandwidth of 0.45.

Quantification and statistical analysis

Data analysis was performed using Excel (Microsoft) Prism 9.1 (GraphPad). Statistical details are described in the Results, Figures, and Figure Legends. All data are mean \pm SEM. Exact values of n are reported where appropriate. Depending on the experiment, n represents number of patches, number of cells, or number of crystals.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- The K2P TRAAK has physically distinct leak and mechanically-activated open states
- FHEIG syndrome causing mutations promote mechanically-activated openings
- A pan-K2P activating mutation in the channel cavity promotes leak openings
- Leak openings occur with TM4 down and mechanically-activated openings with TM4 up

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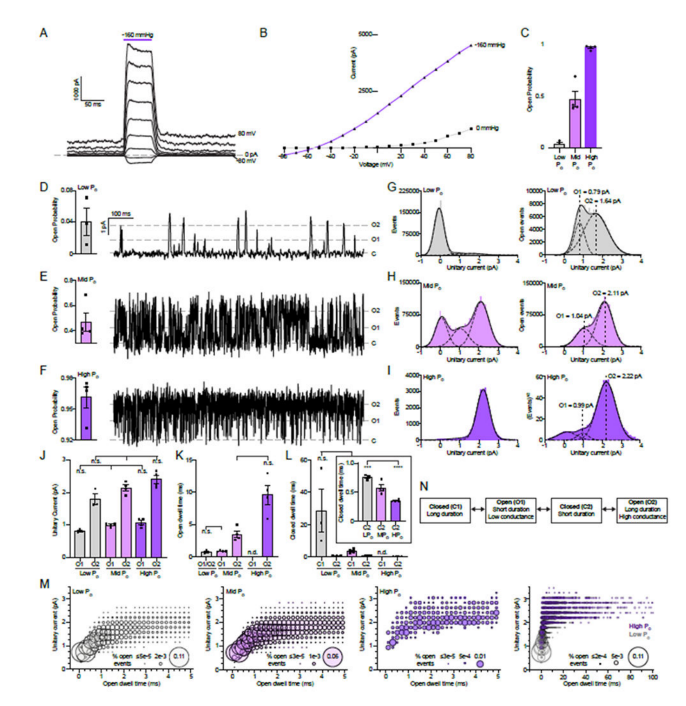


Figure 1 –. Distinct basal and mechanically-activated open states in TRAAK_{WT}. (A) Macroscopic currents recorded across an inside-out patch pulled from a TRAAK-expressing cell in response to a voltage step protocol ($V_{hold} = 0$, $V_{test} = -80$ to 80, V = 10 mV, 20 mV intervals displayed) with a pressure step applied at each voltage (purple bar). (B) Current-voltage relationship from (A). (C) Open probability calculated from single channel records (0.04 ± 0.02 , 0.47 ± 0.07 , and 0.96 ± 0.01 for low P_O , mid P_O , and high P_O , respectively (mean \pm sem, n = 3, 4, and 4 patches)). All single channel data in the paper were recorded at $V_{hold} = 0$ mV in a ten-fold gradient of [K⁺] and are presented in

physiological convention. (D-F) 1 s portion from representative low P_{O} , mid P_{O} , and high Po recordings, respectively. (G-I) All event (left) and open-only or square root all event (right) current histograms from representative low P_O, mid P_O, and high P_O recordings, respectively. (J) Unitary currents of O1 and O2 states (0.81 ± 0.03 pA and 1.80 ± 0.16 pA, 0.99 ± 0.03 pA and 2.14 \pm 0.09 pA, and 1.07 \pm 0.07 pA and 2.41 \pm 0.13 pA for low $P_O,$ mid P_O , and high P_O recordings, respectively (mean \pm sem, n= 3, 4, and 4 patches)). (K) Open dwell time of O1 and O2 states (0.80 ± 0.15 ms, 0.91 ± 0.05 ms and 3.52 ± 0.51 ms, and 9.60 ± 1.46 ms for low P_O, mid P_O, and high P_O recordings, respectively (mean \pm sem, n= 3, 4, and 4 patches)). (L) Closed dwell times of C1 and C2 states (28.70 \pm 13.50 and 0.77 \pm 0.03 ms, 3.59 \pm 0.67 and 0.58 \pm 0.05 ms, and 0.35 \pm 0.01 ms for low Po, mid Po, and high P_O recordings, respectively (mean \pm sem, n=3, 4, and 4 patches, n.d. not determined, n.s. not significant, ***p = 0.0006, ****p < 0.0001 (one-way ANOVA with Tukey correction)). (M) Unitary current-open dwell time relationships for low Po, mid Po, and high Po open events and an overlay of low Po and high Po relationships at expanded time scale. Bubble size is proportional to percentage of open events. (N) Four state TRAAK gating model. See also Figures S1–5, Table S1.

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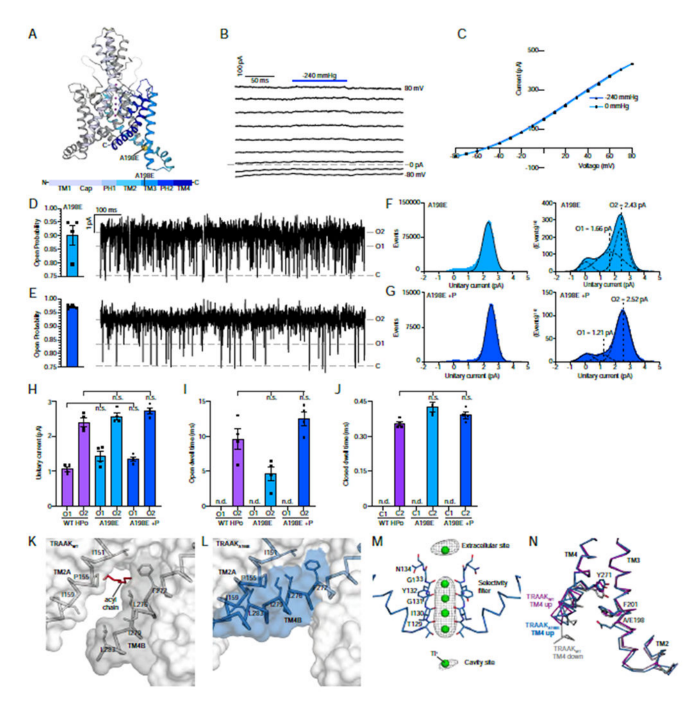
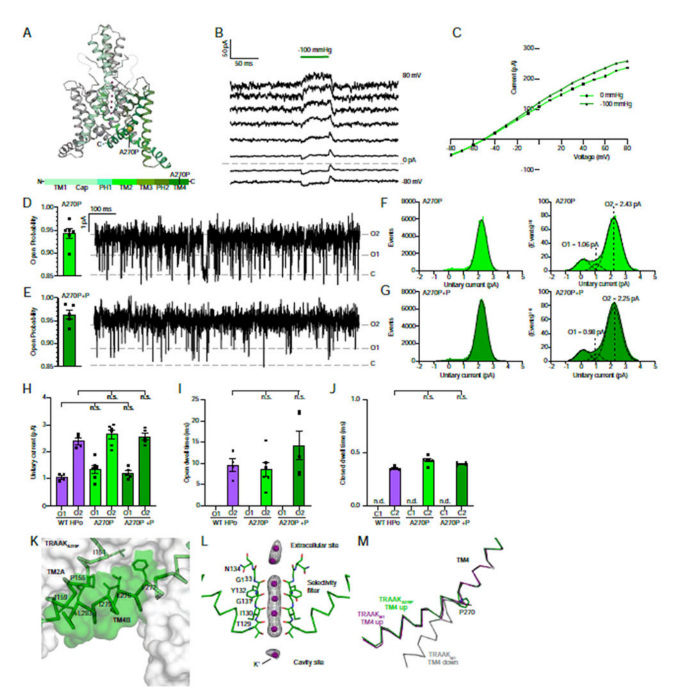


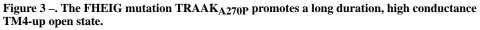
Figure 2 –. The FHEIG mutation \mbox{TRAAK}_{A198E} promotes a long duration, high conductance TM4-up open state.

(A) Crystal structure of TRAAK_{A198E}. Side view from the membrane plane with one protomer gray and the second protomer colored according to the key below. A198E is shown as a yellow sphere and K⁺ ions are colored purple. (B) Macroscopic currents from a TRAAK_{A198E}-containing patch in response to a voltage step protocol (V_{hold} = 0, V_{test} = -80 to 80, V = 10 mV, 20 mV intervals displayed) with a pressure step applied at each voltage (dark blue bar). (C) Current-voltage relationship from (B). (D,E) Open probability calculated from all TRAAK_{A198E} and TRAAK_{A198E} with pressure (+P) records

(left, $P_{O} = 0.90 \pm 0.04$ and $P_{O} = 0.97 \pm 0.003$ (mean \pm sem, n = 4 patches) and 1 s portion from representative recordings. (F,G) All event (left) and square root all event (right) current histograms from representative recordings. (H) Unitary currents of TRAAK_{WT} HP_O, TRAAK_{A198E}, and TRAAK_{A198E}+P O1 and O2 states (1.07 \pm 0.07 and 2.41 \pm 0.13 pA, 1.44 ± 0.14 and 2.57 ± 0.10 pA, and 1.35 ± 0.06 and 2.72 ± 0.09 pA, respectively). (I) Open dwell times of TRAAK_{WT} HPO, TRAAKA198E, and TRAAKA198E+P O2 states $(9.60 \pm 1.46 \text{ ms}, 4.63 \pm 1.03 \text{ ms}, 12.46 \pm 1.02 \text{ ms}, \text{respectively})$. (J) Closed dwell times of TRAAK_{WT} HP_O, TRAAK_{A198E}, and TRAAK_{A198E}+P C2 (0.35 \pm 0.01 ms, 0.42 \pm 0.02 ms, and 0.39 ± 0.01 ms, respectively). For H-J, data are mean \pm sem, n = 4 patches; n.d., not determined; n.s., not significant (one-way Anova with Dunnett correction). (K) View of the membrane-facing lateral opening in a TRAAK_{WT} TM4-down structure (PDB 4WFF). A cavity-bound lipid acyl chain blocks conduction. (L) TRAAKA198E in the same view as (K). A TM4-up conformation seals the membrane opening. (M) Ions in a TRAAKA198E-Tl⁺ structure. Anomalous density (grey) around Tl⁺ ions (green) displayed at 2.5 σ (extracellular / selectivity filter / cavity ions). (N) Overlay of the TM2-TM3-TM4 interaction from TRAAKA198E and TRAAKWT structures. A198E sterically promotes a TM4-up open state. See also Figures S1-7, Tables S1,S2.

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(A) Crystal structure of TRAAK_{A270P}. Side view from the membrane plane with one protomer gray and the second protomer colored according to the key below. A270P is shown as a yellow sphere and K⁺ ions are colored purple. (B) Macroscopic currents from a TRAAK_{A270P}-containing patch in response to a voltage step protocol (V_{hold} = 0, V_{test} = -80 to 80, V = 10 mV, 20 mV intervals displayed) with a pressure step applied at each voltage (dark green bar). (C) Current-voltage relationship from (B). (D,E) Open probability calculated from all TRAAK_{A270P} and TRAAK_{A270P} with pressure (+P) records

(left, $P_O = 0.94 \pm 0.01$ and $P_O = 0.96 \pm 0.01$ (mean \pm sem, n = 6 and 5 patches) and 1 s portion from representative recordings. (F,G) All event (left) and square root all event (right) current histograms from representative recordings. (H) Unitary currents of TRAAK_{WT} HP_O, TRAAK_{A270P}, and TRAAK_{A270P}+P O1 and O2 states (1.07 ± 0.07 and 2.41 ± 0.13 pA, 1.34 ± 0.15 and 2.65 ± 0.16 pA, and 1.21 ± 0.09 and 2.57 ± 0.12 pA, respectively). (I) Open dwell times of TRAAK_{WT} HP_O, TRAAK_{A270P}, and TRAAK_{A270P}+P O2 states (9.60 \pm 1.46 ms, 8.56 \pm 1.69 ms, and 14.47 \pm 3.30 ms, respectively). (J) Closed dwell times of TRAAK_{WT} HP_O, TRAAK_{A270P}, and TRAAK_{A270P}+P C2 (0.35 \pm 0.01 ms, 0.43 \pm 0.01 ms, and 0.39 ± 0.01 ms, respectively). For H-J, data are mean \pm sem, n = 4, 6, and 5 patches; n.d., not determined; n.s., not significant (one-way Anova with Dunnett correction). (K) View of the membrane-facing cytoplasmic half of TM4 in TRAAK_{A270P}. A TM4-up conformation seals the membrane opening. (L) Ions in the TRAAKA270P-K⁺ structure. Polder omit F_o - F_c density (grey) around K⁺ ions (purple) displayed at 5 and 5.5 σ for extracellular and selectivity filter and cavity ions, respectively. (M) Overlay of TM4 from TRAAKA270P and TRAAKWT structures. A270P kinks TM4 to promote a TM4-up open state. See also Figures S1–7, Tables S1,S2.

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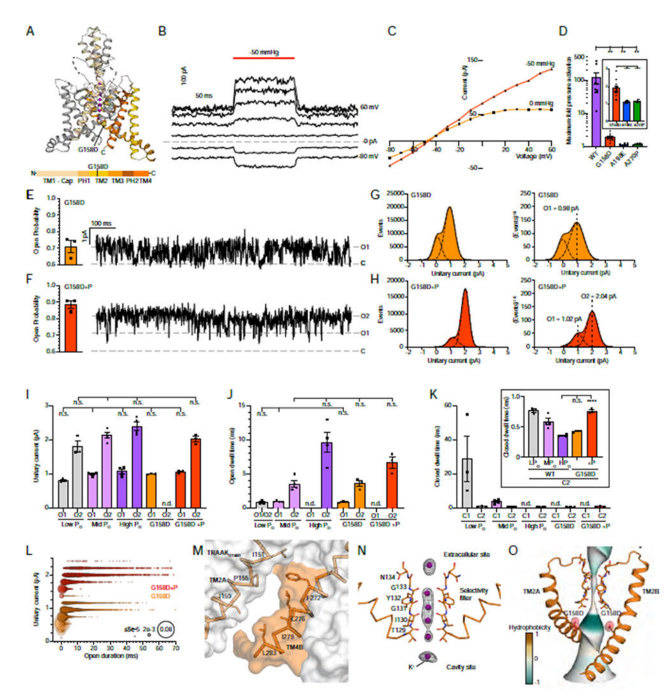


Figure 4 –. The gain-of-function mutation \mbox{TRAAK}_{G158D} promotes a short duration, low conductance TM4-down open state.

(A) Crystal structure of TRAAK_{G158D}. Side view from the membrane plane with one protomer gray and the second protomer colored according to the key below. G158D is shown as a yellow sphere and K⁺ ions are colored purple. (B) Macroscopic currents from a TRAAK_{G158D}-containing patch in response to a voltage step protocol ($V_{hold} = 0$, $V_{test} = -80$ to 80, V = 10 mV, 20 mV intervals displayed) with a pressure step applied at each voltage (dark orange bar). (C) Current-voltage relationship from (B). (D) Maximum fold-activation by pressure of macroscopic TRAAK_{WT}, TRAAK_{G158D}, TRAAK_{A198E}, and TRAAK_{A270P}

currents with expanded scale in inset (119.8 \pm 45.38, 1.91 \pm 0.14, 1.12 \pm 0.02, and 1.16 \pm 0.02, respectively, mean \pm sem, n = 9, 11, 7, and 8, **p < 0.0021, ****p < 0.0001 (one-way ANOVA with Dunnett correction). (E,F) Open probability calculated from all TRAAK_{G158D} and TRAAK_{G158D} with pressure (+P) records (left, P_O = 0.71 \pm 0.04 and P_O = 0.89 \pm 0.02 (mean \pm sem, n=3 patches) and 1s portion from representative recordings. (G,H) All event (left) and square root all event (right) current histograms from representative recordings. (I) Unitary currents of TRAAK_{WT} LP_O, MP_O, HP_O, TRAAK_{G158D}, and TRAAK_{G158D}+P O1 and O2 states (0.81 \pm 0.03 pA and 1.80 \pm 0.16 pA, 0.99 \pm 0.03 pA and 2.14 \pm 0.09 pA, 1.07 ± 0.07 pA and 2.41 ± 0.13 pA, 0.99 ± 0.01 , and 1.06 ± 0.02 and 2.02 ± 0.08 pA, respectively). (J) Open dwell times of TRAAK_{WT} LPO, MPO, HPO, TRAAKG158D, and TRAAK_{G158D}+P O1and O2 states (0.80 \pm 0.15 ms, 0.91 \pm 0.05 ms and 3.52 \pm 0.51 ms, 9.60 ± 1.46 ms, 0.88 ± 0.07 and 3.58 ± 0.46 ms, and 6.63 ± 0.83 ms, respectively). (K) Closed dwell times of TRAAK_{WT} LPO, MPO, HPO, TRAAKG158D, and TRAAKG158D+P C1 and C2 states (28.70 ± 13.50 and 0.77 ± 0.03 ms, 3.59 ± 0.67 and 0.58 ± 0.05 ms, 0.35 \pm 0.01 ms, 0.42 \pm 0.01 ms, and 0.76 \pm 0.03 ms, respectively). For I-K, data are mean \pm sem, n = 3, 4, 4, 3 and 3 patches; n.d., not determined; n.s., not significant, ****p < 0.0001 (one-way Anova with Tukey correction). (L) Unitary current-open dwell time relationships for TRAAK_{G158D}, and TRAAK_{G158D}+P open events and an overlay at expanded time scale. Bubble size is proportional to percentage of open events. (M) View of the membrane-facing cytoplasmic half of TM4 in TRAAK_{G158D}. (N) Ions in the TRAAK_{G158D}-K⁺ structure. Polder omit F_0 - F_c density (grey) around K⁺ ions (purple) displayed at 6.5 and 6 σ for selectivity filter and extracellular and cavity ions, respectively. (O) The conduction path in TRAAK_{G158D} colored by hydrophobicity. G158D increases cavity electronegativity to promote a TM4-down open state. See also Figures S1-7, Tables S1,S2.

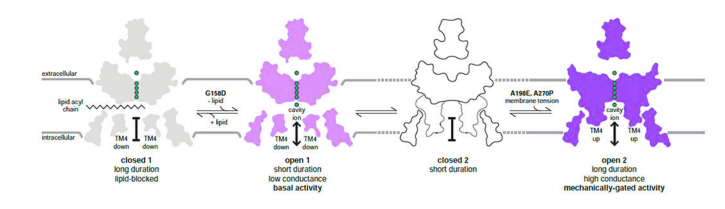


Figure 5 –. An integrated model for TRAAK gating with distinct basal and mechanically-gated open states.

Known structures are mapped to the linear-four state model for TRAAK gating. Basal activity corresponds to a TM4-down, low conductance, short duration open state O1. Mechanically-gated activity corresponds to a TM4-up, high conductance, long duration open state O2. Long duration closures correspond to a TM4-down lipid-blocked state C1. The unknown structure of the short duration closed state C2 is drawn without ions and the position of TM4 is indicated with dashed lines.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and virus strains		
STABL2	ThermoFisher	CAT#10268019
Chemicals, peptides, and recombinant proteins		
DM (Sol-Grade)	Anatrace	CAT#D322S
DM (Analytical Grade)	Anatrace	CAT#D322
pepstatin	Gold Biotechnology	CAT#P-020
leupeptin	Gold Biotechnology	CAT#L-010
aprotinin	Gold Biotechnology	CAT#A-655
soy trypsin inhibitor	Gold Biotechnology	CAT#T9128
benzamidine	Gold Biotechnology	CAT#B-6506
AEBSF	Gold Biotechnology	CAT#A-540
E-64	Gold Biotechnology	CAT#E-064
phenylmethysulfonyl fluoride	Gold Biotechnology	CAT#P-470
CNBr-conjugated sepharose	Cytiva	CAT#17043001
Critical commercial assays		
mMessage machine T7 transcription kit	ThermoFisher	CAT#AM1344
Deposited data		
TRAAK A198E-13E9 Fab complex in K+	This paper	7LJ5
TRAAK A198E-13E9 Fab complex in Tl+	This paper	7LJA
TRAAK A270P-13E9 Fab complex in K+	This paper	7LJ4
TRAAK G158D-13E9 Fab complex in K+	This paper	7LJB
Experimental models: Cell lines		
13E9 antibody hydriboma	Brohawn 2014	gift from R. Mackinnon, Rockefeller University
Xenopus laevis oocytes	This paper	N/A
Pichia pastoris SMD1163 (Alternative SMD1168)	ThermoFisher	CAT#C17500
Experimental models: Organisms/strains		
Xenopus laevis	Nasco	CAT#LM00531
Oligonucleotides	1	1
G158D_Fwd: GATATCTTGTTGGCTGGA	this paper	N/A
G158D_Rev: GAAGAGAGGGGATACCAAC	this paper	N/A
A198E_Fwd: GAAATGCTGTTCCTCCTCATC	this paper	N/A
A198E_Rev: GGACAGTACACGAACCAA	this paper	N/A
A270P_Fwd: CCTTACTTCGCTAGTGTGCTC	this paper	N/A
A270P_Rev: CAGTCCCAACAGAATCC	this paper	N/A

REAGENT or RESOURCE	SOURCE	IDENTIFIER
A198D_Fwd: CGTGTACTGTCCGATATGCTGTTC	this paper	N/A
A198D_Rev: GAGGAGGAACAGCATATCGGA	this paper	N/A
A198W_Fwd: CGTGTACTGTCCTGGATGCTGTTC	this paper	N/A
A198W_Rev: GAGGAGGAACAGCATCCAGGA	this paper	N/A
Recombinant DNA	. <u>I</u>	l
TRAAK pPICz vector	Brohawn et. al, 2012	N/A
TRAAK PGEM vector	Sorum et. al, 2021	N/A
Software and algorithms		1
Graphpad Prism Software	GraphPad Prism Software, Inc.	N/A
Excel	Microsoft, Inc.	https://www.microsoft.com/en-us/ microsoft-365/excel
XDS	Kabsch et al. 2010	https://xds.mr.mpg.de/
STARANISO	Tickle et al. 2018	http://staraniso.globalphasing.org/
Refmac5	Murshudov et al., 2011	N/A
UCSF Chimera	Pettersen et al., 2004	https://www.cgl.ucsf.edu/chimera/
Coot	Emsley et al. 2010	https://www2.mrc-lmb.cam.ac.uk/personal/ pemsley/coot
MolProbity	Chen et al. 2010	http://molprobity.biochem.duke.edu/
PyMol	Schrödinger, LLC	https://pymol.org/2/support.html#page-top
HOLE	Smart et al 1996	http://www.holeprogram.org/
CHAP	Kleese et al 2019	https://www.channotation.org/
CHARMM	Brooks et al., 2009; Jo et al., 2007, 2008	https://www.charmm-gui.org/
Python v3.6	Python	https://www.python.org
Python scripts for single channel event unitary current calculation, unitary current and dwell time 2D histogram calculation	This paper	https://github.com/sbrohawn/ ElectrophysiologyScripts/tree/singles
Antibodies	1	1
Anti-GFP nanobody	Addgene	CAT#61838