Vascular $K_{ATP}$ channel structural dynamics reveal regulatory mechanism by Mg-nucleotides

Min Woo Sung, Zhongying Yang, Camden M. Driggers, Bruce L. Patton, Barmak Mostofian, John D. Russo, Daniel M. Zuckerman, and Show-Ling Shyng

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Vascular tone is dependent on smooth muscle $K_{ATP}$ channels comprising pore-forming Kir6.1 and regulatory SUR2B subunits, in which mutations cause Cantu syndrome. Unique among $K_{ATP}$ isoforms, they lack spontaneous activity and require Mg-nucleotides for activation. Structural mechanisms underlying these properties are unknown. Here, we determined cryogenic electron microscopy structures of vascular $K_{ATP}$ channels bound to inhibitory ATP and glibenclamide, which differ informatively from similarly determined pancreatic $K_{ATP}$ channel isoform (Kir6.2/SUR1). Unlike SUR1, SUR2B subunits adopt distinct rotational “propeller” and “quatrefoil” geometries surrounding their Kir6.1 core. The glutamate/aspartate-rich linker connecting the two halves of the SUR-ABC core is observed in a quatrefoil-like conformation. Molecular dynamics simulations reveal MgADP-dependent dynamic tripartite interactions between this linker, SUR2B, and Kir6.1. The structures captured implicate a progression of intermediate states between MgADP-free inactivated, and MgADP-bound activated conformations wherein the glutamate/aspartate-rich linker participates as mobile autoinhibitory domain, suggesting a conformational pathway toward $K_{ATP}$ channel activation.

Significance

Vascular $K_{ATP}$ channels formed by the potassium channel Kir6.1 and its regulatory protein SUR2B maintain blood pressure in the physiological range. Overactivity of the channel due to genetic mutations in either Kir6.1 or SUR2B causes severe cardiovascular pathologies known as Cantu syndrome. The cryogenic electron microscopy structures of the vascular $K_{ATP}$ channel reported here show multiple, dynamically related conformations of the regulatory subunit SUR2B. Molecular dynamics simulations reveal the negatively charged ED-domain in SUR2B, a stretch of 15 glutamate (E) and aspartate (D) residues not previously resolved, play a key MgADP-dependent role in mediating interactions at the interface between the SUR2B and Kir6.1 subunits. Our findings provide a mechanistic understanding of how channel activity is regulated by intracellular MgADP.

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same condition (22–24). First, unlike in Kir6.2, Kir6.1 cytoplasmic domains (CDs) were displaced from the membrane too far to interact with PIP2 for channel opening. Second, unlike pancreatic channels, which have a predominant propeller-shaped conformation when bound to ATP and Glib (22, 24), vascular KATP channels held four distinct conformations, two resembling propellers and two quatrefoils, marked by varying degrees of rotation of SUR2B toward the core Kir6.1 tetramer. Importantly, a long segment of SUR not previously resolved in any KATP structures, linking NBD1 and transmembrane domain 2 (TMD2), was revealed within vascular KATP structures to mediate the cytosolic interface between SUR2B and Kir6.1. In particular, the linker’s unique 15 glutamate/aspartate residues termed the ED-domain (25) established a nexus of interactions engaging SUR2B-NBD2 with Kir6.1 C-terminal domain (CTD). Molecular dynamics (MD) simulations showed MgADP binding to NBD2 was accompanied by substantial reconfiguration at this nexus, revealing the ED-domain provides a mobile autoinhibitory interaction that guards the transition of SUR2B from MgADP-free inactivated state to MgADP-bound activated state. Together, our findings point to a structural pathway through which SUR regulates Kir6 channel gating.

Results and Discussion

Structure Determination of Kir6.1/SUR2B KATP Channels with ATP and Glib. Vascular KATP channels were purified from COSm6 cells coexpressing rat Kir6.1 and SUR2B (97.6 and 97.2% sequence identity to human Kir6.1 and SUR2B, respectively). COSm6 cells lack endogenous KATP channels and have been used extensively as a heterologous expression system for KATP channel structure–function studies (16, 26). Channels were solubilized in digitonin, purified via an SUR2B epitope tag, and imaged in the presence of 1 mM ATP (no Mg2+) and 10 μM Glib on graphene oxide–coated grids, as described in Materials and Methods.

In vascular KATP channel structures as in pancreatic channels, we found SUR2B anchored to Kir6.1 via interactions mediated by transmembrane helix 1 (TM1) of SUR2B-TMD0 and Kir6.1-TM1 (Fig. 1). However, conformational deviations from fourfold symmetry of the SUR2B were noted in two-dimensional class averages (SI Appendix, Fig. S1). To obtain clear SUR2B maps, we implemented symmetry expansion and extensive focused three-dimensional (3D) classification of Kir6.1 tetramer with individual SUR2B (Materials and Methods) (SI Appendix, Fig. S2), which isolated four 3D classes having identical Kir6.1 tetramer structures but different SUR2B orientations (Fig. 1 and SI Appendix, Fig. S2). When symmetrized, two of the 3D classes, designated P1 and P2, resembled the pancreatic channel propeller conformations previously reported (22, 24). The other two, designated Q1 and Q2, resembled the “quatrefoil conformation” reported for human pancreatic KATP in which the SUR1 NBDs are dimerized (27). Further refinement yielded cryoEM maps with overall resolutions of 3.4, 4.2, 4.0, and 4.2 Å for the P1, P2, Q1, and Q2 conformations, respectively (SI Appendix, Fig. S3). The maps were sufficient to build a full atomic model for all of Kir6.1 minus the disordered C terminus (365 to 424) for all

Fig. 1. Structures of the vascular KATP channel in the presence of ATP and Glib. (A) Schematics of SUR2B and Kir6.1 domain organization. (B) CryoEM density map of (Kir6.1)4SUR2B P1, side view. (C) Structural model of (Kir6.1)4SUR2B P1, side view. (D) Fourfold symmetrized structure model of P1 viewed from the top (i.e., extracellular side). (E) CryoEM density map of (Kir6.1)4SUR2B Q1, side view. (F) Structural model of (Kir6.1)4SUR2B Q1, side view. (G) Fourfold symmetrized structure model of Q1 viewed from the top.
conformations, with clear sidechain densities for most residues (SI Appendix, Fig. S3d) and also models for most of SUR2B (see Materials and Methods for details). Densities for ATP, Glib, and some lipids were well resolved (SI Appendix, Fig. S3d). Significantly, the Q1 conformation included definitive densities in SUR2B for L0, which is the linker connecting TMD0 and the ABC core, and also the N1-T2 linker, which connects NBD1 to TMD2; neither is resolved in the human pancreatic K\textsubscript{ATP} quatrofoil structure previously determined (27). The P- and Q-like conformations differ by a major rotation of the SUR2B-ABC core toward the Kir6.1 tetramer, clockwise when viewed from the extracellular side (Fig. 1D and G). P1 and Q1 were the dominant particle populations within the P- and Q-like forms, respectively, differing from P2 and Q2 by degree of rotation and specific features. We first focus on structural differences between P1 and Q1, which provided the highest resolutions.

**Structural Correlates of Kir6.1 Functional Divergence.** Although the Kir6.1 tetramer was similarly configured in all P and Q conformations for SUR2B, it included several features distinct from Kir6.2 in our published pancreatic channel structure determined under similar conditions with ATP and Glib (Protein Data Bank [PDB]: 6BAA). The Kir6.1 channel CD was extended intracellularly away from the membrane by \( \sim 5.8 \) Å, and simultaneously counterclockwise rotated (viewed from the extracellular face) by 12.4° (Fig. 2A). The Kir6.x CD is thus corkscrewed away from the membrane in Kir6.1/SUR2B, compared to Kir6.2/SUR1. Constrictions in the two cytoplasmic gates, namely the helix bundle crossing (F178) and the G-loop (G304, I305), indicate a closed Kir6.1 channel pore, similar to Kir6.2 under the same condition (SI Appendix, Fig. S4). However, the distance between the helix bundle crossing gate and the G-loop gate is significantly larger in Kir6.1 due to the un tethered CD.

In K\textsuperscript{+} channels, variations in the turret region surrounding the pore entryway have been shown to affect selectivity filter stability and ion conduction (28). Compared to Kir6.2, the turret of Kir6.1 contains an extra 11 aa (102-YAYMEKGITEK112). We found this sequence formed a helix and loop structure that extends the turret further out into the extracellular space (Fig. 2B and C). Functional studies using Kir6.1–Kir6.2 chimeras previously identified residues in Kir6.1 thought to impart its smaller unitary conductance, specifically M148 and N123-V124-R125 (29). M148 in Kir6.1 (replacing Kir6.2-V138) was proposed to reduce pore entrance diameter, while N123 in Kir6.1 (replacing Kir6.2-S113) was hypothesized to impact an intersubunit salt bridge between R146 and E150, which in other Kir channels is formed by corresponding residues and critical for channel conduction (29, 30). However, our structure found M148 facing the pore helix (Fig. 2C) rather than the entrance and that no significant difference exists in the adjacent pore diameters between Kir6.1 and Kir6.2, nor in their intersubunit salt bridges. Interestingly, N123-V124-R125 of Kir6.1 is located
between the turret extension and the pore loop and interacts with Y104 in the turret extension (Fig. 2C). Future mutagenesis studies will determine the contributions of these interactions to Kir6.1 channel conductance.

We next assessed structural differences between Kir6.1 and Kir6.2 in two elements interacted with associated activity at ATP PIP2 binding sites: the N-terminal amphiphatic helix known as the slide helix (SH), and the connecting strand between TM2 and the CTD called the C-linker (Fig. 2 B, D, and E). In our Kir6.2 structure (22), SH is bent halfway at the D58 position resembling a 3α helix (31). In contrast, SH in Kir6.1 formed a continuous helix extending toward the neighboring Kir6.1, thus compressing the PIP2 binding pocket. In Kir6.2, the C-linker forms a helix that tethers the CTD close to the membrane, which positions critical PIP2-binding residues such as R176 for PIP2 interaction. Rather different, the C-linker in Kir6.1 unwound into an unstructured loop stretching toward the cytoplasm, which deflected R186 (corresponding to Kir6.2-R176) away from the PIP2 binding site (Fig. 2B).

Recent MD simulations of Kir2.2 in membranes containing mixed phospholipids showed that PS can also occupy the PIP2 binding pocket (33). Because no exogenous PIP2 was added to our protein sample, the simplest interpretation of the structural data is that the more abundant PS resides in the Kir6.1 PIP2 binding pocket. However, a possibility remains that the density includes endogenous PIP2 or other phospholipids copurified with the channel. CryoEM densities matching ATP were clearly resolved in Kir6.1 tetramers, at sites located between the N-terminal domains and CTDs of adjacent Kir6.1 subunits, matching sites in Kir6.2/SUR1 channels. However, unlike for Kir6.2, ATP had fewer close residue interactions in Kir6.1 due to displacement of the Kir6.1-CD. In particular in pancreatic channels, SUR1-K205 (in L0) directly participates in binding ATP at its inhibitory site (23, 27, 34), while the corresponding vascular channel residue SUR2B-K203 was displaced from potential ATP binding (Fig. 2D). Thus, the constellation of ATP interactions was sparser and hence likely weaker when Kir6.1-CD was displaced from the membrane.

Taken together, the translocation of the Kir6.1-CD away from the membrane compromised binding of both ATP and PIP2. This correlates well with the basal inactivity and reduced ATP sensitivity of the vascular KATP channel compared to Kir6.2 channels (35, 36). Rotation and downward movement of the Kir6.2-CD have been detected in minor subclasses of ATP- and Glib-bound pancreatic Kir6.2/SUR1 structures (23, 24), indicating similar dynamics occur but less stably persist. Moreover, translation and/or rotation of the CD is observed in Kir2, Kir3, and bacterial Kir channels (37–40), and recent cryoEM studies of Kir3 channels found that increased PIP2 concentra-
tions shift particle distributions toward those having CD teth-
ered close to the PIP2 membrane sites (41). Thus, a common model of KATP channel activity involves channel opening dependent on PIP2 binding, which in turn depends on engagement by the Kir6.x-CD modulated by its vertical translocation/rotation. Accordingly, in vascular Kir6.1 channels, a greater energy barrier is involved in rotating the CD upward to interact with PIP2 than in pancreatic channels whose Kir6.2-CD is more stably tethered to the membrane. This explains why Kir6.2-containing pancreatic channels are spontaneous, while Kir6.1-containing vascular channels are not. By extension, vascular channel activation by Mg-nucleotides likely involves SUR2B-controlled upward movement of the Kir6.1-CD. It has been shown that Kir6.1 binds PIP2 with higher affinity than Kir6.2 in biochemical assays and that once activated by Mg-nucleotides, vascular KATP channels are highly stable and more resistant to PIP2 depletion by polylysine than pancreatic channels (32). In our Kir6.1 structure, the Kir6.1-R70 sidechain is directed toward the lipid density in the PIP2 binding pocket (Fig. 2E). Interestingly, in Kir6.2 the corresponding residue is a proline (P69). It is possible that this sequence variation may contribute to the higher PIP2 affinity and stability of open vascular channels, but future studies are necessary to investigate this. Higher-PIP2 affinity also accounts for long-standing results showing activated vascular KATP channels are much less sensitive to ATP inhibition, as increased PIP2 interaction reduces ATP inhibition in KATP channels (16).

**SUR2B Dynamics.** Focused 3D classification resolved four distinct conformations, P1, P2, Q1, and Q2, showing variable SUR2B orientations (SI Appendix, Fig. S5 and Movie S1). P conformations differed from Q conformations by a large rotation of the ABC core of SUR2B relative to the Kir6.1 tetramer (~41° between P1 and Q1, about the axis defined by N447 in TMD1 and N69 in TMD0, respectively, compared to 63° rotational difference between the propeller and quatrefoil conformations that SUR2B occupies in human pancreatic NBD-dimerized channels measured from the equivalent residues). Within P and Q, P1 and Q1 particles were located over P2 and Q2. Transitions from P1 to P2 and Q1 to Q2 involved alternative rotation steps: P1’s ABC core was 10° further away from Kir6.1 than P2’s, while Q1’s ABC core was 8° closer to Kir6.1 than in Q2. In short, Q1 was the tightest quatrefoil and P1 the most extended propeller. 3D variability analysis in CryoSPARC (SI Appendix, Fig. S6a) indicated SUR2B subunits moved independently between P- and Q-like positions (Movie S2). Further multibody refinement in RELION3 revealed greater heterogeneity within Q1 conformations than in P1, indicating wider dynamic range (SI Appendix, Fig. S6b and Movie S3).

Accompanying rotation, the SUR2B-ABC core also tilts away from Kir6.1. Tilting elevated the ABC core TMD in the Q conformations relative to P conformations (by 2.6 A from P1 to Q1, measured at SUR2B-Y370; SI Appendix, Fig. S5b). Between the pancreatic KATP propeller and quatrefoil forms (NBDs dimerized), the entire ABC-TMDs elevate ~3 A without tilting (27). Tilt in our Q conformations may represent a less defined conformational position, as in the propeller-only conformation. In the NBDs-dimerized pancreatic KATP quatrefoil is the dominant class. Here, Q conformations were less common than P conformations among vascular KATP channel structures in which the NBDs remain separated (SI Appendix, Fig. S2 and Table S1). Probabilities of SUR adopting P- or Q-like conformations therefore correlate with NBD dimerization state, although both occur regardless.

Rotation of SUR2B between P to Q conformations incorporated significant local structural changes. Extracellular contacts between transmembrane bundle 1 (TMB1) and TMD0 restructured both protein–protein and protein–lipid interactions (SI Appendix, Fig. S7). Hydrophobic and electrostatic interactions in P1 are lost in Q1, including T338, L339, and F344 in the TM6-TM7 loop of TMB1, with L165 and R166 in TMS of TMD0. Moreover, a phosphatidylethanolamine molecule moved from between TM2 and TM7 in P1 to between TM5 and TM16 in Q1, likely stabilizing TMD0 and TMB1 interactions. Also noteworthy, in the pancreatic channel structure, SUR1 has an additional hydrophobic sequence (G237TVF240) which anchors the TM6-TM7 loop to TMD0 (SI Appendix, Fig. S7d) (23, 34). Absence of this sequence in SUR2B may impart flexibility that enables SUR2B to swing into Q conformations not observed in SUR1 when ATP and Glib are bound.

The SUR2B-L0 Linker and the Glib Binding Pocket. Transition between P and Q conformations repositioned cytoplasmic structural elements
including L0, the N1-T2 linker, and Kir6.1 N-terminus (Kir6.1(N)), unexpectedly affecting interactions between SUR2B and Kir6.1. In SURx, L0 connects TMD0 to the ABC core and is crucial to \( K_{ATP} \) gating (42–45). In SUR2B, we obtained two distinct L0 conformations, corresponding to P and Q conformers. In SUR2B-P1, we observed continuous cryoEM density of L0 (Fig. 3A and Q conformers. In SUR2B-P1, we observed continuous cryoEM density of L0 (Fig. 3A and Q conformations). Lipid density seen in P1 but absent in Q1 is shown in cyan. (B) Structure of (Kir6.1)\(_2\)SUR2B in P1 conformation showing L0 (red) viewed from the side (left) and from the cytoplasmic side near the membrane (right). The N1-T2 linker visible in these views is shown in green. (C) Structure of (Kir6.1)\(_2\)SUR2B in Q1 conformation viewed from the side and the cytoplasm. (D) Structure of (Kir6.2)\(_2\)SUR1 (PDB: 6BAA) bound to Glib and ATP for comparison.

Vascular \( K_{ATP} \) channels are inhibited by Glib but are ~10- to 50-fold less sensitive than pancreatic \( K_{ATP} \) channels (17, 47). Glib cryoEM density was well resolved in both P1 and Q1 conformations of the vascular \( K_{ATP} \) structure, where it bound within the same pocket of SUR2B (Fig. 4) as in SUR1 (24, 26). Also similar to pancreatic Kir6.2/SUR1 channels, cryoEM density of the distal KNt of Kir6.1 lay within the cleft between the two halves of the ABC core and immediately adjacent the Glib binding pocket (26). The structure model of the Glib binding site in P1 shows key interactions are largely conserved between SUR1 and SUR2B (Fig. 4C). However, the binding pose of Glib in the Q1 conformation is compressed compared to that in P1, with the Y1205 sidechain moved upward, which requires the 1-chloro-4-methoxy-benzene group to also move to avoid W423 in a neighboring helix (Fig. 4C). Also, an electrostatic interaction between chloride in Glib and nitrogen of R304 is eliminated in Q1. Worth noting, the SUR2B-Y1205 equivalent residue in SUR1 is S1238, and substitution of serine by tyrosine at this position has been shown to partly underlie Glib’s lower-affinity inhibition of SUR2 channels (48). Of particular interest, substitution of S1238 to Y in SUR1 converts the Glib inhibition of pancreatic Kir6.2/SUR1 channels from nearly irreversible to readily reversible similar to SUR2-containing channels (49, 50), suggesting the S1238Y mutation may affect Glib off rate. This may arise through steric hindrance from the flexible tyrosine sidechain, as observed in the Q1 conformation.

The P1 to Q1 translocation was accompanied by more substantial change to the opposite side of the Glib binding pocket at F215, T227, and Y228 of the L0 linker (Fig. 4C). Previous studies of L0 of SUR1 have shown that Glib binding indirectly involves Y230 and W232 (Y228 and W230 in SUR2B), which stabilize the TM helices lining the Glib binding pocket (22); mutation of these residues to alanine reduces sensitivity to Glib (49, 50). In SUR2B P1 conformation, we found the hydrophobic Y228 and W230 sidechains, as well as F215 in the lower part of L0, similarly stabilized the TM helices along the Glib binding pocket (Fig. 4C), as occurs in SUR1. Specifically, F215 lay buried in a hydrophobic cavity formed by W230 from L0 and Y371, F2107, and L1206 from TMB1. However, in Q1, L0 was significantly remodeled at the interface with TMB1. In particular, a loop segment including P218-Y228 seen in P1 is raised and transformed into a helix in Q1. This helical element newly filled the hydrophobic cavity between TMD0 and TMB1, otherwise occupied by lipids in P1 (Fig. 3A). As further consequence in Q1, Y228 and F215 in L0 are displaced from the cavity, and Y371 and T227 occupy the space vacated by the sidechain of Y228. The movement of Y228 out of the cavity eliminates Y228. The movement of Y228 out of the cavity eliminates the hydrophobic packing between L0 and the TM helices lining the Glib binding pocket, thus disrupting the integrity of the pocket.
in similar fashion to the Y230A mutational effect in SUR1 (51). Lastly, the density of Kir6.1Nt in the ABC core central cleft also differed between P1 and Q1 (Fig. 4D). In P1, a strong continuous density of KNt was present, braced by R804 and N1090 of SUR2B, a pair of residues guarding entry to the cleft. The KNt density in Q1 was considerably weaker and discontinuous, indicating a more-labile conformation that may contribute to weak Glib binding at its adjacent pocket (26, 52). In summary, as the SUR2B-ABC core changes from P conformation to Q, L0 and Kir6.1Nt undergo remodeling that affects the Glib binding pocket.

The N1-T2 Linker. In all published pancreatic K<sub>ATP</sub> channel structures, the critical N1-T2 linker of SUR1 has remained unresolved (22–24, 26, 27, 34, 53), suggesting dynamic instability. In the density map of our vascular Kir6.1/SUR2B channel from the P1 particle set, the C-terminal end of the N1-T2 linker was sufficiently resolved (Fig. 5A), and we were able to build a polyalanine helical structure into the density map (residues 961 to 976). Density for the rest of N1-T2 (residues 911 to 960) remained largely unresolved in P1. However, the density for the entire linker was apparent in the map for our vascular K<sub>ATP</sub> channel Q1 structure (Fig. 5B), although resolution of residues 911 to 960 was insufficient for modeling. Specifically, the linker extended from NBD1 through the space between the two NBDs, then continued through the gap between the outer surface of NBD2 and the adjacent CTD of Kir6.1, before connecting to TMD2 (Fig. 5B and SI Appendix, Fig. S8). The location of the SUR2B N1-T2 linker contrasts sharply with locations of corresponding linkers in other ABCC proteins, including the CI<sup>−</sup> channel CFTR and the yeast cadmium transporter Ycf1p. In CFTR, the N1-T2 linker equivalent is known as the R domain, which is phosphorylated by PKA to allow CFTR gating by Mg-nucleotides. In the unphosphorylated CFTR structure, the R domain is wedged in the cleft between the two halves of the ABC core, preventing NBD dimerization (54). In the phosphorylated CFTR structure, the R domain relocates to the outer surface of NBD1 (SI Appendix, Fig. S8), which allows NBD dimerization, hence CFTR gating by Mg-nucleotides (55, 56). In the Ycf1p structure, the N1-T2 linker is found at the outer surface of NBD1 similar to phosphorylated CFTR (57) even though the NBDs are separate. The peculiar location of the SUR2B N1-T2 linker suggests the linker has adopted a separate role in regulating functional coupling between SUR2B and Kir6.1.

In SUR2, the N1-T2 linker includes at its C-terminal end a stretch of 15 aa consisting exclusively of negative-charged glutamate and aspartate designated the ED-domain (947 to 961) (SI Appendix, Fig. S8), which is unique among all ABCC proteins. Previous mutational studies have implicated the ED-domain in transducing MgADP binding in SUR2A to opening of Kir6.2 (25). Disruption of the ED-domain prevented the normal activation response to MgADP and to pinacidil, a potassium channel opener. In the Q1 structure, the density corresponding to the ED-domain is sandwiched between NBD2 and Kir6.1-CTD (Fig. 5) and surrounded by positively charged residues from Kir6.1Nt, Kir6.1-CTD, and NBD2 of SUR2B (Fig. 5B), an array of partners for electrostatic interactions. To understand the potential molecular interactions and their functional relevance, we employed MD simulations of the (Kir6.1)4SUR2B Q1 structure.

**MD Simulations Reveal MgADP-Dependent Dynamic Interactions between the ED-Domain, NBD2, and Kir6.1-CTD.** To assess conformational dynamics of the ED-domain and its interacting...
partners and how they may be dependent on the nucleotide binding status at the two NBDs, we performed simulations under two conditions. In one, ATP is bound to Kir6.1 and NBD1 of SUR2B, as present in our cryoEM structure. In the second condition, Mg$^{2+}$ is included with ATP bound at NBD1, and MgADP is docked into NBD2 (Fig. 6A). In both conditions, Glib was omitted from the structure to allow the SUR2B TMDs to be free of constraint during simulations. To assess reliability, three independent 1-μs simulations for each condition were carried out (SI Appendix, Fig. S9a). As with many biomolecular simulations, ours do not exhibit true equilibrium-like repeated fluctuations about mean values (58), although the combined 6 μs permitted structural inferences (Fig. 6C and E). The root-mean-square fluctuation (RMSF) analyses showed high degrees of fluctuations of NBD1, the N1-T2 linker, and NBD2 (SI Appendix, Fig. S9b), consistent with overall lower resolutions of these domains in cryoEM maps (SI Appendix, Fig. S3b). However, particular interactions between the ED-domain and NBD2 depended on whether NBD2 was occupied by MgADP, and in turn those ED-domain–NBD2 interactions controlled direct interaction of NBD2 with Kir6.1-CTD.

During simulations, the ED-domain exchanged interactions between surrounding positively charged residues from Kir6.1Nt, Kir6.1-CTD, and NBD2 (Fig. 5B and Movies S5 and S6). When MgADP was absent at NBD2, the first one-half of the ED-domain (947 to 953) was most frequently in contact with SUR2B-NBD2 Walker A K1348; this was infrequent with MgADP bound at NBD2. To quantify MgADP dependence of the ED-domain–Walker A interaction, we measured the minimum distances between the ED-domain residues 947 to 953 and K1348 throughout simulations, comparing results with and without MgADP at NBD2 (Fig. 6B and C). In the absence of MgADP, sidechain oxygens from ED residues were frequently within 4 Å of the sidechain nitrogen of K1348, supporting a salt bridge or strong electrostatic interaction (59). In contrast, in the presence of MgADP, sidechain oxygens from ED residues were frequently within 4 Å of the sidechain nitrogen of K1348, supporting a salt bridge or strong electrostatic interaction (59). In contrast, in the presence of MgADP, sidechain oxygens from ED residues were frequently within 4 Å of the sidechain nitrogen of K1348, supporting a salt bridge or strong electrostatic interaction (59). In contrast, in the presence of MgADP, sidechain oxygens from ED residues were frequently within 4 Å of the sidechain nitrogen of K1348, supporting a salt bridge or strong electrostatic interaction (59). In contrast, in the presence of MgADP, sidechain oxygens from ED residues were frequently within 4 Å of the sidechain nitrogen of K1348, supporting a salt bridge or strong electrostatic interaction (59). In contrast, in the presence of MgADP, sidechain oxygens from ED residues were frequently within 4 Å of the sidechain nitrogen of K1348, supporting a salt bridge or strong electrostatic interaction (59). In contrast, in the presence of MgADP, sidechain oxygens from ED residues were frequently within 4 Å of the sidechain nitrogen of K1348, supporting a salt bridge or strong electrostatic interaction (59).

NBD2 also frequently formed close contacts with Kir6.1-CTD in the absence of MgADP but not when NBD2 included MgADP (Movies S5 and S6). With no MgADP, a loop

Fig. 5. Comparison of cryoEM densities of Kir6.1 N terminus and SUR2B N1-T2 linker in P1 and Q1 conformations. (A) Overall cryoEM density of (Kir6.1)$_4$-SUR2B in gray with density of one Kir6.1 and its N terminus (KNt) highlighted in blue and density of the SUR2B N1-T2 linker highlighted in green. (B) Close-up view of the N1-T2 linker density in (Kir6.1)$_4$-SUR2B structure. Blue spheres are positively charged residues near the ED-domain. G1345 in the NBD2 Walker A motif and E1318 in the A-loop of NBD2 (\textsuperscript{1315}VRYEN\textsuperscript{1319}) are shown as reference points.
upstream of the Walker A motif in NBD2 (\textsuperscript{1315}VRYEN\textsubscript{1319}) named A-loop for aromatic residue interacting with the adenosine ring of ATP (60) frequently extended across the inter-subunit gap to interact with a cluster of positively charged residues in Kir6.1-CTD, including R323, K341, R347, and R352 (Movie S5). In direct contrast, when MgADP was bound to SUR2B-NBD2, the A-loop instead consistently interacted with MgADP at NBD2, far from the Kir6.1-CTD. The A-loop in SUR2B includes Y1317, which interacts with the adenine ring of bound MgADP at NBD2. Simultaneously, the dissociation of the ED-domain from Walker A K1348 that occurred with MgADP binding at NBD2 freed the ED-domain to move in between NBD2 and Kir6.1-CTD. There, the ED-domain interacted with positive-charged residues in Kir6.1-CTD (Movies S5 and S6). Effectively, the Kir6.1-CTD exchanged the A-loop for the ED-domain and stabilized each conformation. Quantitatively, minimum distances measured between E1318 in A-loop and the four positive residues in Kir6.1-CTD documented the closer relation of A-loop and Kir6.1-CTD throughout the simulations in the absence of MgADP than when MgADP was bound (Fig. 6E). Minimum distance below 4 Å sufficient for E1318 salt bridge formation was seen in all three runs lacking MgADP but only transiently (3.6 ns) in a single of...
The three runs with MgADP (Fig. 6E). The nucleotide-dependent dynamics between the NBD2 A-loop and Kir6.1-CTD was also shown by tracking distance between the C-α of E1318 and center of the mass of the C-α for the Kir6.1-CTD-positive residues (an example run for each condition shown in Fig. 6F).

The dynamic, tripartite interactions between the ED-domain, NBD2, and Kir6.1-CTD, and the dependence of these interactions on MgADP found in MD simulations significantly advances our understanding of the mechanism of SUR-mediated channel stimulation by Mg-nucleotides. In the absence of MgADP, the ED-domain has preferred interactions with NBD2 Walker A K1348, while the A-loop E1318 is engaged with Kir6.1-CTD. This hinders NBD2 from undergoing further conformational transition toward that of the NBDs-dimerized human pancreatic channel quatrefoil structure (27), which shows SUR1-NBD2 further rotated toward NBD1 and also away from the positively charged residues in Kir6-CTD (PDB: 6C30) (27). Upon MgADP binding to NBD2, the ED-domain is dissociated from K1348, while the NBD2 A-loop becomes stabilized by the bound MgADP, unable to extend toward Kir6.1-CTD. As a sequence of results, the ED-domain is free to move toward other surrounding positively charged residues including those in Kir6.1-CTD, which further prevents the interactions between NBD2 and Kir6.1-CTD, thus allowing NBD2 to undergo further rotation toward dimerization with NBD1. Supporting this understanding, an ion pair formed by R347 in the Kir6.1-CTD, with E1318 in the A-loop of SUR2B-NBD2, has previously been reported to play a role in channel activation by MgADP and the potassium channel opener pinacidil (61). Disruption of this ion pair by charge neutralization enhances MgADP/pinacidil gating, while charge swap restored wild-type–like sensitivity to MgADP/pinacidil (61). Our findings support the hypothesis that in order for NBDs to dimerize, interactions between SUR-NBD2 and Kir6-CTD must dissolve. Accordingly, disruption of the Kir6.1 R347-SUR2B E1318 salt bridge facilitates MgADP/pinacidil stimulation, as breaking the salt bridge promotes nucleotide binding at NBD2 and allows the further NBD2 movement needed for NBD dimerization and channel activation. The ED-domain in particular, by interacting with Walker A K1348, acts essentially as a mobile autoinhibitory motif, akin to autoinhibition mechanisms in many kinases (62), that occludes NBDs dimerization in the absence of MgADP and is deflected to permit dimerization.

Fig. 7. Residues mutated in Cantú patients mapped onto the Kir6.1/SUR2B channel structure. (A) Residues mutated are shown as blue (Kir6.1) or magenta (SUR2B) in P1 conformation as spheres (Left) or in stick model (Right). Rat SUR2B numbering is used. Corresponding human mutations with rat residue in parentheses are as follows: H60Y (H60), D207E (D207), G294E (G294), G380C (G377), P432 (P429), A478V (A475), D793V (D789), G815A (G811), Y985S (Y981), G989E (G985), H1005L (H1001), W1018G (W1014), T1019/K (T1015), S1020P (S1016), N1-T2 linker is colored green and labeled together with the second elbow helix leading to TM12 of TMD2 in SUR2B. Red numbers mark the TM helices shown.
when MgADP has bound to NBD2. In this way, the ED-domain functions as a gatekeeper to prevent unregulated channel activation in the absence of MgADP.

**Implications for Canůt Mutations.** Taken together, our structures and MD simulations capture conformations that appear intermediate between the NBD-separated inactive and NBD-dimerized active states. The structural knowledge sheds light on how Canůt mutations (Fig. 7A) may cause gain of function in vascular K<sub>ATP</sub> channels. In Kir6.1, V65M in the SH and C176S in the pore-lining helix likely enhance function by increasing channel Po, which has been demonstrated in equivalent mutations in Kir6.2 (63). Most Canůt mutations identified to date are in ABC<sub>C9</sub>. Significantly, many of them affect residues in TM12, including Y981 and G985 in the second elbow helix, and W1014, T1015, and S1016 at the top (Fig. 7). TM12 is connected to the N1-T2 linker (Fig. 7D). Many other Canůt mutations are in domains interacting with TM12, including a series throughout TM13 (F1035, C1039, C1046, S1050, and M1056), as well as H1001 in TM12 and R1112 in TM14, which interface TM13 (Fig. 7C). One most frequently mutated residue R1150 of TM15 is adjacent to the structured helix portion of the N1-T2 linker, C-terminal to the ED-domain (Fig. 7D). The interconnectivity of these residues and their association with the N1-T2 linker suggest they may in common govern a critical conformational change during channel gating by Mg-nucleotides at the NBDs. Consistent with this notion, Y981S, G985E, and M1056I have been shown to enhance channel response to MgADP stimulation (42). Of note, C1039Y in G985E, and M1056I have been shown to enhance channel nucleotides at the NBDs. Consistent with this notion, Y981S, which further corrupts the Mg-nucleotide binding site and static interaction between NBD2 A-loop and Kir6.1-CTD, that prevents unregulated activation. Specifically, ED-domain in the N1-T2 linker acts as an autoinhibitory motif of quatrefoil-like conformations, and SURx linkers and drug sensitivities. In contrast, the serendipitous appearance of quatrefoil-like conformations, and SURx linkers which have been missing in previous K<sub>ATP</sub> structures and are now seen at critical domain interfaces, affords insights into the long-sought structural dynamics shared by K<sub>ATP</sub> channels in regulating their activity. The Q conformations adopted by SUR2B are most simply interpreted as transitional states between the inactive NBD-separated and the active NBD-dimerized SUR conformations.

The several conformations isolated from the cryoEM data set, together with the dynamics revealed by 3D variability analyses and captured by MD simulations, suggest a model hypothesis for how Mg-nucleotide interactions with SUR2B activates Kir6.1 (Fig. 8). In this model, individual SUR2B subunits transition between P and Q conformations. In the Q conformations and without Mg-nucleotides at NBD2, the ED-domain in the N1-T2 linker acts as an autoinhibitory motif that prevents unregulated activation. Specifically, ED-domain interaction with Walker A K1348 at NBD2 promotes electrostatic interaction between NBD2 A-loop and Kir6.1-CTD, which further corrupts the Mg-nucleotide binding site and promotes conformational change to the NBD-dimerized quatrefoil state for channel activation.

**Summary.** Insights into how a particular complex operates is often gained by comparing related complexes, anticipating that similarities and differences in structure and function will correlate. In this study, we sought to determine the cryoEM structure of vascular K<sub>ATP</sub> channels, composed of Kir6.1 and SUR2B, in the presence of ATP and Glib, for comparison to pancreatic K<sub>ATP</sub> channel (Kir6.2/SUR1) structures determined with the same conditions. The structures we obtained reveal multiple elements showing distinct configurations that may account for channel-specific conductance, ATP inhibition, and drug sensitivities. In contrast, the serendipitous appearance of quatrefoil-like conformations, and SURx linkers which have been missing in previous K<sub>ATP</sub> structures and are now seen at critical domain interfaces, affords insights into the long-sought structural dynamics shared by K<sub>ATP</sub> channels in regulating their activity. The Q conformations adopted by SUR2B are most simply interpreted as transitional states between the inactive NBD-separated and the active NBD-dimerized SUR conformations.
also withholds NBD2 from dimerization with NBD1. Addition of Mg-nucleotides relieves autoinhibition imposed by the ED-domain, coupling organization of the Mg-nucleotide binding site to liberation of NBD2 to rotate toward NBD1 for dimerization. Yet-to-be-determined mechanisms are required to explain how dimerization of NBDs in SUR2B leads Kir6.1–CTD to move up to the membrane to interact with PIP2 for channel opening. The model would predict that inhibitory ligands such as Gly or stimulatory ligands such as Mg-nucleotides or the potassium channel opener pinacidil, are able to shift the equilibrium of SUR2B toward P or Q conformations to drive channel closure or opening, respectively. It is important to note that dimerization of the NBDs was not observed during the 1-μs simulation in the presence of MgADP/MgATP; moreover, only one SUR2B is present in the simulation, which prevents consideration of potential structural impact of neighboring SUR2B subunits. Future structures with NBDs dimerized and MD simulations of the full channel are required to confirm and extend our understanding of KATP channel activation. This notwithstanding, we speculate the general scheme of the model applies to other KATP channels with variations to explain isoform-specific sensitivities for Mg-nucleotides and drugs. The structures presented here serve as a framework for understanding channel regulation and dysregulation and will aid development of isoform-specific pharmacological modulators to correct channel defects in Cantú and other diseases involving vascular KATP dysfunction.

Materials and Methods
Expression and purification of Kir6.1/SUR2B channels, cryo-EM imaging, data processing, and modeling were performed using published protocols (22, 23, 26, 64) and are described in detail in SI Appendix. Briefly, recombinant adenoviruses containing the coding sequences of Kir6.1 and FLAG-tagged SUR2B were used to infect COS7 cells and expressed channels purified via the FLAG tag. Purified channel complexes were spotted on grids coated with graphene-oxide, vitriﬁed, and imaged on a Titan Krios 300 kV cryoelectron microscope. Image processing and analysis were carried out in RELION-3.0 and CryoSPARC. Models were built by fitting previously published Kir6.2/SUR1 structures and in SWISS-MODEL and reﬁned in Coot and Phenix.

MD simulations were performed at all-atom resolution using AMBER 16 (65) with graphics processing unit (GPU) acceleration. The starting structure was developed from the Q1 model (four Kir6.1 and one SUR2B) with ﬂexible linkers built in SWISS-MODEL. Glyc was removed to allow the TMDs to relax during simulations. The structures were protonated at pH 7 and inserted in a bilayer membrane composed of 1-palmitoyl-2-oleoyl-phosphatidylcholine lipids and surrounded by an aqueous solution of 0.15 M KCl. Pairwise distances were analyzed from the simulated trajectories using the gmx pairdist tool in Gromacs 2019.4 (66). Detailed methods for MD simulations and data analysis are provided in SI Appendix.

Data Availability. CryoEM density maps have been deposited to the Electron Microscopy Data Bank (P1: EMD-23864, P2: EMD-23881, Q1: EMD-23880, and Q2: EMD-23882). Coordinates for (Kir6.1)4/SUR2B atomic models have been deposited to the Protein Data Bank (P1: 7MIT, P2: 7MJP, Q1: 7MIO, and Q2: 7MQ). MD simulation data have been deposited to the open-access repository Zenodo (5346127). All other study data are included in the article and/or supporting information.

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