Structural Models for the KCNQ1 Voltage-Gated Potassium Channel†

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ABSTRACT: Mutations in the human voltage-gated potassium channel KCNQ1 are associated with predisposition to deafness and various cardiac arrhythmia syndromes including congenital long QT syndrome, familial atrial fibrillation, and sudden infant death syndrome. In this work 3-D structural models were developed for both the open and closed states of human KCNQ1 to facilitate structurally based hypotheses regarding mutation—phenotype relationships. The KCNQ1 open state was modeled using Rosetta in conjunction with Molecular Operating Environment software, and is based primarily on the recently determined open state structure of rat K,1.2 (Long, S. B., et al. (2005) Science 309, 897–903). The closed state model for KCNQ1 was developed based on the crystal structures of bacterial potassium channels and the closed state model for K,1.2 of Yarov-Yarovoy et al. ((2006) Proc. Natl. Acad. Sci. U.S.A. 103, 7292–7207). Using the new models for KCNQ1, we generated a database for the location and predicted residue—residue interactions for more than 85 disease-linked sites in both open and closed states. These data can be used to generate structure-based hypotheses for disease phenotypes associated with each mutation. The potential utility of these models and the database is exemplified by the surprising observation that four of the five known mutations in KCNQ1 that are associated with gain-of-function KCNQ1 defects are predicted to share a common interface in the open state structure between the S1 segment of the voltage sensor in one subunit and both the S5 segment and top of the pore helix from another subunit.

This interface evidently plays an important role in channel gating.

KCNQ1 is a voltage-gated potassium channel that participates critically in human physiology and is subject to several heritable disease-linked mutations (2, 3). The most common splice variant of KCNQ1, also known as K,LQT1 or K,7.1, contains 676 residues and consists of a cytosolic N-terminal domain followed by the voltage sensor (S1−S4), a canonical pore domain (S5-P-S6) and a long cytosolic C-terminus (see Figure 1). The best-characterized physiological function of KCNQ1 relates to its expression in cardiac tissue, where it coassembles with KCNE1 (minK), a single transmembrane-span protein to form a channel complex that mediates the slow delayed rectifier current, I,Ks, a critical electrical component of the cardiac action potential necessary for myocardial repolarization (2–5).

Disruption of KCNQ1 function by dominant mutations causes more than 50% of genotype-known cases of congenital long QT syndrome type 1 (LQTS1), a potentially fatal arrhythmic disorder (6–10). In the autosomal dominant form of the disease, Romano–Ward syndrome, mutant KCNQ1 exerts dominant-negative effects that cause loss-of-function. Recessive loss-of-function mutations in KCNQ1 cause Jervell and Lange–Nielsen syndrome (JLNS), characterized by both long-QT syndrome and congenital deafness (8, 9, 11). Deafness in this syndrome highlights the important role that KCNQ1 plays in tissues outside the cardiovascular system including the kidney, stomach, and ear. The association with congenital hearing loss, for example, reflects an important role for KCNQ1 channels in recycling potassium ions to reestablish transmembrane potentials following auditory stimulation in the cochlea and vestibule of the inner ear (2, 11, 12). Of the spectrum of disease-causing mutations in the KCNQ1 gene, by far the most common are missense mutations that result in single amino acid residue replacements (6–9).

While LQTS1 and JLNS are the consequence of KCNQ1 mutations that induce varying degrees of loss-of-function, there are also rare mutations that result in gain-of-function (1, 13–17, 55). Such mutations have been associated with familial atrial fibrillation, the short-QT syndrome, and sudden infant death syndrome (SIDS).

While several mutagenesis and electrophysiology studies have probed structure–function relationships for KCNQ1 (2, 3, 5) and while a high-resolution structure for one of its extramembrane domains was recently determined (18), this

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1 Abbreviations: AF, atrial fibrillation; DN, dominant negative; E1, the KCNE1 protein; JLNS, Jervell and Lange–Nielsen syndrome; LQTS1, long-QT syndrome, type 1; P-helix, pore helix; Q1, the KCNQ1 protein; rmsd, root-mean-squared deviation; RWS, Romano–Ward syndrome (autosomal dominant LQTS1); SIDS, sudden infant death syndrome; WT, wild type.
The gap opening and gap extension penalties were set to 3.0 and 0.3, respectively. These parameters were previously optimized using a small database of manually refined membrane protein alignments (Dong, Smith, and Meiler, unpublished).

The alignment is most confident in the S5-P-S6 region, where the algorithm can draw from significant sequence identity (~38%) in addition to secondary structure and transmembrane prediction data. Within the voltage sensor domain the sequence identity is reduced to ~17%. Here unambiguous alignment of each transmembrane helix (S1 through S4) relies heavily on secondary structure and transmembrane prediction data. The resulting alignment between the KCNQ1 target and K_1.2 template sequences was used for model building and is shown in Figure 1.

**Construction of Open and Closed State Models.** Construction of the open state model of KCNQ1 using the X-ray structure for K_1.2 as the template and the sequence alignment shown in Figure 1 is complicated by two issues. First, in the original K_1.2 open state crystal structure there are a number of residues that are not resolved in the connecting segments of S1 with S2, S2 with S3, and S3 with S4. Second, side chains were not resolved for S1 and S3 transmembrane segments and were reported in the PDB file as polyalanine segments. This missing information makes it impossible to build the KCNQ1 sequence/structure directly onto the K_1.2 scaffold for S1 or S3. However, using a hybrid method of comparative modeling and Rosetta-Membrane techniques Yarov-Yarovoy et al. addressed these issues by building the missing pieces of K_1.2 into the remainder of the structure (34). This completed K_1.2 crystal structure of the full-length open state of K_1.2 was used as the structural template for the open channel state of KCNQ1.

Because there are no experimentally determined closed state structures of mammalian potassium channels, we used the K_1.2 closed state model of Yarov-Yarovoy et al. (34) as the template for generating the KCNQ1 closed state model. This template was built from the closed state structure of KcsA in the pore domain. For the voltage sensor domain, a hybrid approach using domain assembly and *de novo* protein construction was employed (34).

MOE software (*Molecular Operating Environment*, version 2006.08, Chemical Computing Group) was used to create initial monomer models of the open and closed states of KCNQ1. For each state, twenty candidate models were generated and scored with the MOE-Homology functionality. MOE-Homology utilizes a segment-matching procedure (35) in combination with modeling of indels based on the methods of Fechteler et al. (36) The resulting open and closed state
monomer ensembles are presented in Figure 2. Four copies of the optimally scoring candidate from each ensemble in Figure 2 were then transformed into the corresponding tetramer structure using the BIOMT records reported in the K,1.2 crystal structure using UCSF Chimeras software (37). The intermonomer interfaces of the resulting tetramer models were then optimized using Rosetta (38) to repack the side chains, followed by all-atom energy minimization with AMBER9 (39) to allow for small backbone adjustments. This Rosetta/AMBER repack/minimize cycle was repeated until the backbone conformation and AMBER energy had converged. These refined open and closed state tetramer models are illustrated in Figure 3.

RESULTS

Development and Validation of Open and Closed State Models for KCNQ1. The development of the KCNQ1 models was highly dependent on the use of Rosetta algorithms. The Rosetta software suite is one of the best comparative modeling packages (40) as assessed in recent CASP experiments (41), particularly for cases where sequence identity is low and when flexible segments are present in the molecule of interest. Indeed, physically realistic homology models built with Rosetta can be more accurate than their experimental structural templates (42). Rosetta has recently been used to develop models for membrane protein structures (34, 43).

Rosetta homology and de novo methods were employed to construct KCNQ1 models in the open and closed states using the open and closed state hybrid experimental/model structures (34) of K,1.2 as templates (see Methods). This approach assumes that both KCNQ1 states are similar to the corresponding states of K,1.2.

The open and closed channel models were analyzed with Procheck (44) and Maxisoftware (http://sw-tools.pdb.org/apps/MAXIT/index.html) to assess their quality. All chiral centers were found to be correct, and no close contacts (<2.2 Å) were observed. All bond lengths fell within acceptable ranges, with just 2 (open) and 4 (closed) bond angles falling outside of 6 × the rmsd of dictionary values. All main chain parameter assessments (Ramachandran plot, nonbonded interactions, peptide planarity, H-bond energies) were well within expected ranges for high-resolution structural models (2.0 Å or better). In the open state model, all residues fell within favored (90%), allowed (9%), or generously allowed (1%) regions of the Ramachandran plot. In the closed model, only 5 residues (0.6%) fell into disallowed regions, with the remaining 99.4% being in favored (87%), allowed (11%), or generously allowed (1.5%) regions. Analysis of side chain conformations further confirmed the quality of these models with 99.7% (open) and 99.2% (closed) of χ1 and χ2 rotamers inhabiting favorable conformations. This is well within the expected bandwidth of precision for high-resolution structural models and represents a positive indicator of the reliability of these models.

The lowest energy representatives of the open and closed state ensembles were used to construct models for the channel tetramers (Figure 3). PDB files for these models are provided in the Supporting Information. Following a short energy minimization AMBER force field energies were calculated and compared to the calculated energies for the corresponding K,1.2 template structures. It was found that the open state KCNQ1 tetramer yielded a calculated energy (−10.031 kcal) that is slightly more favorable than K,1.2 (−9425 kcal), with a similar result being observed for the closed state KCNQ1 (−9364 kcal) relative to closed K,1.2 (−9210 kcal).

The results of model validation described above lend confidence to these models. Nevertheless, the limitations of the models need to be recognized and are summarized as follows.
Figure 3: Open and closed state models for the S1–S6 domain of KCNQ1. (A) Left: Extracellular view. Right: View from membrane plane. (B) Intracellular stereoview, with the atoms of Ser349 being highlighted in van der Waals representation.
In both open and closed state KCNQ1 structures the pore domains are modeled with a higher certainty than the voltage sensors because of the larger number, greater completeness, and higher resolution of the available crystal structures for the pore domain, as well as because of the higher sequence identity between the Kv 1.2 and KCNQ1 pore domains relative to the voltage sensors. On the other hand, the extracellular loop regions of these models, which occur primarily at deletions/insertions in the sequence alignment (Figure 1), are modeled at a lower precision (Figure 2) since the conformation of these loops is based entirely on de novo methods and not on experimentally determined regions of the templates.

The open state model is expected to be generally reliable because, except for a few short regions in the voltage sensor domain (see Methods), the entire backbone of the open state is based directly on the experimentally determined structure of Kv 1.2. The closed state model, however, is based on a Kv 1.2 template that was itself created with a hybrid of comparative and de novo modeling techniques. Therefore, the closed state model of KCNQ1 is probably less reliable than the open state model, particularly in the voltage sensor domain. The open and closed state models in Figure 2 have been color-coded to indicate segments that are modeled directly from high-resolution X-ray crystal structures, from low-resolution crystal structures (with details filled in using Rosetta), or based solely on Rosetta prediction.

Description of the Open and Closed State Models of KCNQ1. The open and closed state models of KCNQ1 are illustrated in Figure 3. The topology of the models, the sequential/topological locations of the disease mutations, and key differences between KCNQ1 and Kv 1.2 are summarized in Figure 4 and in Supplementary Figure 1 (Supporting Information). The opening and closing of the KCNQ1 pore is defined by differences in the conformation of the cytosolic-proximal segment of S6 between the open and closed states. In the closed state S6 is a nearly ideal R-helix over its full transmembrane span, with an intrinsic tendency to kink at Pro343 being suppressed by the voltage sensor, as mediated by the downward leverage of the helical S4-S5 linker on the cytosol-proximal segment of S6 from the same subunit. The closest S6-S6 crossing point in the tetrameric closed state occurs near the cytosol-membrane interface and constricts the pore such that ions are unable to enter the inner vestibule of the channel. The channel opens because of changes in the orientation and conformation of the S1-S4 voltage sensor domain that pulls the S4-S5 linker away from the pore to release tension on S6. This permits S6 to kink at Pro343 so that its C-terminus reorients, cantilever-like, away from the central axis of the channel, opening the pore to the cytosol.

Location and Interactions of Disease-Linked Mutation Sites in KCNQ1. Disease-linked amino acid substitutions for the >80 channel domain sites indicated in the sequence/topology diagram of Figure 4 are listed in Supplementary Table 1 (Supporting Information), with a subset of 10 mutants being highlighted in Table 1. For the wild type amino acid side chains of each disease mutant site, we tabulated all amino acids that were within van der Waals contact of the channel opening and closing of the KCNQ1 pore.
<table>
<thead>
<tr>
<th>Amino Acid Mutation</th>
<th>Disease Phenotype</th>
<th>Site Location: Open State</th>
<th>Site Location: Closed State</th>
<th>Surface-Exposed Site? (Open/Closed)</th>
<th>VDW Contacts&lt;sup&gt;b&lt;/sup&gt; in Open State (amino acid, subunit letter, residue number)</th>
<th>VDW Contacts&lt;sup&gt;b&lt;/sup&gt; in Closed State</th>
<th>Known Properties of this Mutant</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ser140Gly</td>
<td>AF</td>
<td>S1, interacts both with middle of S4 and S5 from another subunit</td>
<td>S1, interface with S4</td>
<td>no/no</td>
<td>LEU A 137&lt;br&gt; SER A 143&lt;br&gt; THR A 144&lt;br&gt; GLN A 234&lt;br&gt; ARG A 237&lt;br&gt; TYR B 278</td>
<td>CYS A 136&lt;br&gt; LEU A 137&lt;br&gt; VAL A 141&lt;br&gt; MET A 159&lt;br&gt; ARG A 231&lt;br&gt; PHE A 232</td>
<td>When expressed alone, exhibits loss of function. When co-expressed with E1, shows dramatically enhanced current and rapid activation at all voltages. When co-expressed with WT and E1, channel properties that are intermediate between mutant/E1 and WT/E1 are observed (16). When expressed alone, similar to wild type in absence of KCNE1 (George, A.F., et al., unpublished). Favor open state in strictly KCNE1-dependent manner. See Discussion.</td>
<td>(16)</td>
</tr>
<tr>
<td>Val141Met</td>
<td>AF</td>
<td>S1, interacts with upper part of S5 and beginning of the pore helix, both from adjacent subunit</td>
<td>S1, oriented towards solvent</td>
<td>mostly/yes</td>
<td>ILE A 138&lt;br&gt; LEU A 142&lt;br&gt; TYR B 278&lt;br&gt; TYR B 299</td>
<td>LEU A 137&lt;br&gt; SER A 140&lt;br&gt; LEU A 142&lt;br&gt; PHE A 232</td>
<td>When expressed alone (no E1 or WT), very similar to WT but significantly higher current. When co-expressed with E1, loses much of its voltage dependence of activation—open at all potentials. Retains K&lt;sup&gt;C&lt;/sup&gt; selectivity. When co-expressed with E1 and WT, exhibits properties that are intermediate between V141ME1-only and WT/E1-only (14). Favor open state in a partially KCNE1-dependent manner. See Discussion.</td>
<td>(14;55)</td>
</tr>
<tr>
<td>Ser225Leu</td>
<td>LQTS1</td>
<td>S4, located near beginning of S4, oriented towards S3-S4 loop</td>
<td>S4, interacting with end of S1 from same subunit and at interface with S4 from another subunit</td>
<td>yes/slightly</td>
<td>VAL A 221&lt;br&gt; THR A 224</td>
<td>VAL A 221&lt;br&gt; ARG A 228&lt;br&gt; LEU B 282&lt;br&gt; TYR B 299</td>
<td>When co-expressed only with E1, current is reduced and V&lt;sub&gt;i&lt;/sub&gt; is shifted towards more positive potential. Trafficks normally. When co-expressed with WT, exerts DN effect to reduce current (17). Favor closed state (or disfavors open state), at least in presence of KCNE1.</td>
<td>(56;17)</td>
</tr>
<tr>
<td>Val254Leu</td>
<td>LQTS1</td>
<td>in S4-S5, interacting with S6 from some subunit and S6 from adjacent subunit</td>
<td>S4-S5, at interface with S6 from both same and different subunit</td>
<td>slightly/slightly</td>
<td>LEU A 250&lt;br&gt; LEU A 251&lt;br&gt; SER A 253&lt;br&gt; VAL A 255&lt;br&gt; HIS A 258&lt;br&gt; LEU A 347&lt;br&gt; GLY A 350&lt;br&gt; PHE A 351&lt;br&gt; LYS A 354&lt;br&gt; LEU B 342</td>
<td>LEU A 250&lt;br&gt; LEU A 251&lt;br&gt; SER A 253&lt;br&gt; VAL A 255&lt;br&gt; HIS A 258&lt;br&gt; ARG A 259&lt;br&gt; LEU A 353</td>
<td>Loss of function when expressed alone. When co-expressed with WT, DN effect on channel current. When co-expressed with WT and E1, DN effect on channel current and increase in the V&lt;sub&gt;i&lt;/sub&gt; for channel activation (37). Favor closed state (or disfavors open state) in KCNE1-independent manner.</td>
<td>(7;57)</td>
</tr>
<tr>
<td>Glu261Asp</td>
<td>JLNS</td>
<td>in S5, interacting with S6 from same subunit and close to S6 from adjacent subunit</td>
<td>S5, interacts with S6 from both same and different subunit, also interacts with S4-S5 linker from different subunit</td>
<td>partial/slightly</td>
<td>HIS A 258&lt;br&gt; GLN A 260&lt;br&gt; THR A 264&lt;br&gt; THR A 265&lt;br&gt; PRO A 343&lt;br&gt; ILE A 346&lt;br&gt; LEU A 347</td>
<td>ARG A 259&lt;br&gt; GLN A 260&lt;br&gt; LEU A 262&lt;br&gt; THR A 265&lt;br&gt; LEU A 342&lt;br&gt; ILE A 346&lt;br&gt; PHE D 351&lt;br&gt; LYS D 354</td>
<td>When co-expressed with E1 exhibits minimal current. When co-expressed with both E1 and WT exhibits only minimal current—strong DN effect (58). Exhibits severe mistrafficking with or without E1 or WT co-expression. Also causes mistrafficking of co-expressed WT (59). Misfolds/mistrafficks in KCNE1 independent manner. Also causes mistrafficking of co-expressed WT.</td>
<td>(10;59;60)</td>
</tr>
</tbody>
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Smith et al.
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<tr>
<th>Amino Acid Mutation</th>
<th>Disease Phenotype</th>
<th>Site Location: Open State</th>
<th>Site Location: Closed State</th>
<th>Surface-Exposed Site? (Open/Closed)</th>
<th>VDW Contacts(^a) in Open State (amino acid, subunit letter, residue number)</th>
<th>VDW Contacts(^b) in Closed State</th>
<th>Known Properties of this Mutant</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ile274Val</td>
<td>SIDS</td>
<td>S5, located at interface between S1 and S4 of another subunit</td>
<td>S5, modest interaction with S6 from same subunit</td>
<td>partial/yes</td>
<td>PHE A 270 LEU A 273 TYR A 278 TYR A 299 LEU D 137 MET D 238</td>
<td>PHE A 270 LEU A 273 TYR A 278 LEU A 303 PHE D 232 LEU D 236</td>
<td>Similar to WT in the absence of E1. When co-expressed with E1 exhibits higher current than WT, faster activation kinetics, and slow deactivation, although no significant change in (V_{1/2}); (TE Rhodes, AF George et al., unpublished). Favors open state in KCNE1-specific manner. See Discussion.</td>
<td>(13)</td>
</tr>
<tr>
<td>Ala300Thr</td>
<td>reported as LQTS1 but may be AF</td>
<td>at beginning of P-helix, oriented towards S6 of another subunit</td>
<td>P-helix, interacts with top of S6 from different subunit</td>
<td>yes/yes</td>
<td>SER A 298 TRP A 304</td>
<td>SER A 298 TYR A 299 ASP A 301 LYS B 326</td>
<td>When co-expressed only with E1, current is reduced and (V_{1/2}); is shifted towards more negative potential and activation is accelerated. Trafficks normally. When co-expressed with WT, no loss of WT current. (17) Favors open state. See Discussion.</td>
<td>(17:61)</td>
</tr>
<tr>
<td>Trp305Ser</td>
<td>JLNS</td>
<td>P-helix, interacts with filter, filter-S6 connector, and S6, all from same subunit and also filter from adjacent subunit</td>
<td>P-helix, interacts with S6, P-helix and filter, all from same subunit and also filter from adjacent subunit</td>
<td>no/no</td>
<td>SER A 276 VAL A 280 GLY A 306 THR A 309 LYS A 318 VAL A 319 VAL A 319</td>
<td>SER A 276 VAL A 280 PHE A 296 ALA A 302 TRP A 304 GLY A 306 THR A 309 LYS A 318 VAL A 319</td>
<td>When expressed alone or with E1: complete loss of function. When co-expressed with E1 and WT, observed channel properties match that of WT(62). Probably misfolds/mistrafficks prior to co-assembly with WT.</td>
<td>(10:62:63)</td>
</tr>
<tr>
<td>Val307Leu</td>
<td>Short QT syndrome (gain-of-function)</td>
<td>P-helix, at interface with S6 from another subunit, also some contact with S5 from same subunit</td>
<td>P-helix, at interface with S6 from a different subunit, also some contact with S5 from same subunit</td>
<td>slightly/slightly</td>
<td>LEU A 273 LEU A 303 VAL A 308 VAL A 310 SER B 330</td>
<td>LEU A 273 LEU A 303 VAL A 308 THR A 311 LYS B 326 SER B 330</td>
<td>Co-expression with E1 and in presence or absence of WT leads to more rapid channel activation and a shift to more negative (V_{1/2}) (1). When expressed in absence of E1 or WT, similar current to WT, slightly delayed activation, and no inactivation (unlike WT-only) (64). Favors open state in a partially KCNE1-dependent manner (see Footnote 2).</td>
<td>(1) (64)</td>
</tr>
<tr>
<td>Ser349Trp</td>
<td>LQTS1</td>
<td>S6, oriented towards center of pore, exposed to solvent</td>
<td>S6, interacting with other S6 segments</td>
<td>yes/no</td>
<td>GLY A 345 ILE A 346</td>
<td>ARG A 259 ILE A 346 GLY D 348 PHE D 351 ALA D 352 VAL D 355</td>
<td>See Discussion.</td>
<td>(10:54)</td>
</tr>
</tbody>
</table>

\(^a\) An exhaustive listing for all disease mutants is found in Supplementary Table 1 in the Supporting Information.

\(^b\) A residue was deemed to be roughly within van der Waals contact of the disease mutant site when at least one atom from the side chain of that residue was within 4 Å of at least one atom of the wild type side chain at the disease mutation site.
electrophysiological and/or trafficking data, when available, for each mutant. As exemplified in Table 1, disease mutations in KCNQ1 can be grouped into three classes: mutations that result in gain of function, mutations that result in full or partial loss of function as a consequence of KCNQ1 misfolding and/or mistrafficking, and mutations that result in full or partial loss of function due to altered conductance or gating.

DISCUSSION

This work has generated the first atomic-detail models for the open and closed states of KCNQ1 that include its voltage sensor domain. These structural models should, of course, be regarded as working models that are subject to revision as additional experimental structural data becomes available. Nevertheless, even with these caveats in mind, these models immediately provide intriguing and sometimes unforeseen insight into KCNQ1 channel function. This is here illustrated by three examples.

Implications for Interactions of KCNE1 with KCNQ1. While KCNQ1 functions as a voltage-gated potassium channel even in the absence of accessory subunits, its activity under physiological conditions is often profoundly modulated by KCNE1 or other members of the KCNE family of accessory proteins. Evidence indicates that KCNE1 and KCNQ1 coassemble early in the secretory pathway to form a complex that trafficks to the cell surface. In cardiac myocytes, this complex is responsible for $I_{Ks}$, a potassium current that contributes to membrane repolarization during the cardiac action potential (2–5). There is much evidence that the transmembrane and juxtamembrane cytosolic domains of KCNE1 are critical for binding to and regulating KCNQ1 (4, 45–47). Moreover, there is evidence that at least part of the KCNE1 transmembrane domain is in direct contact with the S6 segment of the channel (4, 20, 46–51), although there has been debate regarding whether the KCNE1 transmembrane domain directly participates in the ion conduction pathway.

Our models for KCNQ1 suggest that the transmembrane domain of KCNE1 may be able to directly access the S5-P-S6 domain of the channel as part of its channel modulatory function without requiring a dramatic change in the conformation of KCNQ1. Looking down onto the open and closed state models of KCNQ1 in Figures 3 and 5, clefts are observed between the voltage sensor domains through which lipids and/or proteins might be able to laterally approach the

**Figure 5:** Surface representations of the KCNQ1 models compared to the K,1.2 models developed of Yarov-Yarov et al. (34) In all cases the view is from the cytosol.
pore domain to form direct contacts with S5 and S6 segments. We have carried out simple docking calculations using both open and closed state models of the channel that confirm that there is indeed sufficient space in KCNQ1 for the transmembrane helix of an accessory protein (such as KCNE1) to fit into the clefts between voltage sensors of adjacent subunits. This does not prove that this is the functionally relevant model of interaction between KCNE1 and KCNQ1, but does support the feasibility of such a model. Interestingly, the cleft between the voltage sensors of Kv 1.2 appears to be partially occluded relative to KCNQ1 (Figure 5), and this could explain why Kv 1.2 does not seem to be modulated by KCNE proteins (52, 53).

Structural Hypothesis for the Mechanism of Gain-of-Function Mutations. There are five disease-associated KCNQ1 mutations that are linked to rare gain-of-function $I_{Ks}$ phenotypes (Figure 4 and Table 1). The Ser140Gly mutation was associated with autosomal dominant atrial fibrillation. This mutant forms $I_{Ks}$ channels that activate much more rapidly and generate whole-cell currents ~3 times larger than wild type $I_{Ks}$ channels (16). A mutation at an adjacent site, Val141Met, was also associated with familial atrial fibrillation in the context of the short-QT syndrome and exhibits a biophysical phenotype that is very similar to that observed for the Ser140Gly mutant (14). KCNQ1 Ile274Val is a mutation recently identified in sudden infant death syndrome (13). Electrophysiological studies (Rhodes, et al., manuscript submitted) indicate that the Ile274Val mutation promotes gain-of-function channel behavior that is characterized by increased $I_{Ks}$ whole cell currents and altered deactivation.

The other two gain-of-function mutations, Ala300Thr and Val307Ile, do not result in increased current, but cause $I_{Ks}$ to activate more rapidly and at more negative potentials than wild type $I_{Ks}$ (1, 17).

When the positions of these 5 gain-of-function mutations are examined in the open state channel model, 4 of these sites, Ser140, Val141, and Ile274, and Ala300, seem to be located at the same interface between the voltage sensor from one subunit and the pore domain of another (Figure 6).22 This is very surprising given that these 4 sites span a wide range of sequence space, representing 3 distinct segments

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2 A fifth gain-of-function mutation that is linked to short-QT syndrome, Val307Leu, is not as easy to structurally rationalize from the models as the other four. Val307Leu involves a very conservative amino acid replacement at a residue located on the P-helix. However, unlike three of the other gain-of-function mutations, when Val307Leu is coexpressed with KCNE1, there is no augmentation in the whole-cell current when compared to the wild type (1). Rather, like Ala300Thr this mutation results in gain-of-function by inducing activation at more negative potentials and by accelerating activation kinetics. From the KCNQ1 models, the Val307Ile mutation is not obviously predicted to perturb the relative energies of the open and closed states. In both open and closed models Val307 is predicted to undergo similar interactions with the S5 segment of the same subunit and to participate in an interface with the S6 of a neighboring subunit. Site 307 may play an allosteric role in modulating the voltage dependence of channel opening and to affect the transition kinetics from the closed to open state. Interestingly, there are numerous loss-of-function disease-associated mutations at sites near Ile307 in both sequential and 3-D space (see Figure 4 and Supplementary Table 1). Whether this gain-of-function mutation is truly unusual or whether, in fact, many mutations in the pore region induce channel gain-of-function but are not detected in the human population is not known.
of the channel protein. Ser140 and Val141 are located on the extracellular end of S1 and make contacts across the subunit–subunit interface with residues located near the tops of both S5 and the P-helix from a different subunit. Ile274 is located in S5 and contacts Leu137 from S1 in another subunit. Ile274 is just below the S5 contact point with sites 140 and 141 from S1. The Ile274 side chain site is also highly proximal to Ile235 and Met238 in S4. Last, Ala300 is located on the P-helix at the interface between S6 from the same subunit (in proximity to Ile274) and is adjacent to Ser140 and Val141 in the voltage sensor from another subunit.

The sensor/pore domain–domain interface at which these four mutations are located may help to stabilize the open channel state. Consistent with this possibility is the conservative nature of all four of these mutations. This is in sharp contrast to many loss-of-function mutations found at this interface, which typically are nonconservative substitutions that are predicted to destabilize this interface (see Supplementary Table 1), causing loss of function by disfavoring the open channel state or misfolding. These include Leu137Pro (S1), Arg231His (S4), Ile235Asn (S4), Leu239Pro (S4), Phe275Ser (S5), Tyr278His (S5), Tyr281Cys (S5), and Gly306Arg (P-helix).

While the verity of the model for the voltage sensor domain and voltage sensor/pore interface of the closed state channel model is deemed to be less reliable than for the open state model, it is interesting to observe that in the closed state model (Figure 6) the side chains for Ser140, Val141, and Ile274 are no longer predicted to be involved in intersubunit interactions, but rather appear to be lipid-exposed, which suggests that the fairly conservative gain-of-function mutations may have less of an impact on the energy of the closed channel state.

Importantly, the gain-of-function consequences of the Ser140Gly, Val141Met, and Ile274Val mutants are observed only when the channel is complexed with KCNE1 (13, 14, 16). In the absence of KCNE1, these mutant KCNQ1 channels exhibit near wild type channel properties. Therefore, any model explaining how these mutations induce gain-of-function must ultimately be elaborated to account for this KCNE1 dependence. While experimental data will ultimately be required to fully address this question, it should be noted that the voltage sensor/pore domain interface where these gain-of-function mutations occur sits near the base of the cleft between voltage sensor domains of adjacent subunits (see Figures 3 and 5). As noted in the above section, it seems probable that this cleft could accommodate the KCNE1 transmembrane domain, and we speculate that KCNE1 participates directly in the interface between pore and voltage sensor domains.

Structural Insight into Loss-of-Function KCNQ1 Mutations. Inferring the mechanisms by which loss-of-function mutations cause reduction or complete inhibition of whole-cell I_K, currents can be difficult, even with high quality functional, trafficking, and experimental 3-D structural data in hand. Nevertheless, the cataloging of structural predictions presented in Table 1 and in Supplementary Table 1 for the >80 KCNQ1 mutations associated with LQTS1 and JNLS can be employed to provide constraints that can disfavor possible structural explanations of mutant phenotypes. For example, even in the absence of detailed electrophysiological data loss-of-function due to the Ser349Trp mutation (see Table 1) seems very unlikely to be the consequence of enhanced stabilization of the closed state because the side chain of Ser349 sits at the intersubunit interface between S6 segments in the tightly constricted region of the pore (Figures 3B), where a tryptophan side chain cannot be accommodated without a major structural rearrangement in the closed state. Instead, channel loss-of-function resulting from the Ser349Trp mutation is more likely to be caused either by protein misassembly/mistracking or by occlusion of the open pore by the bulky tryptophan side chains clustered around the mouth of the conduction pathway. Such model-derived insight does not unambiguously resolve underlying disease mechanisms, but can be used to sharpen hypotheses for the molecular biophysical basis of disease—insight that should prove very helpful in designing decisive functional or biochemical experiments.

CONCLUSIONS

Despite the critical importance of KCNQ1 for normal human physiology, we are aware of no progress in expressing and purifying full length KCNQ1 or its channel domain at a level that would enable the initiation of high-resolution structural studies. In the absence of an experimentally determined high-resolution structure, models developed with the best available templates and computational methods can be valuable as the basis for testable hypotheses regarding mechanisms of channel function and modulation, as well as for the structural basis of disease mutation/phenotype relationships. In the case of the KCNQ1 models presented here, we are heartened by the fact that they provide an unanticipated but compelling hypothesis that a majority of known gain-of-function mutation sites are localized at the interface between S1 in the voltage-sensor domain of one subunit and the S5 and P-helix segments of the pore domain of another subunit.

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SUPPORTING INFORMATION AVAILABLE

Supplementary Table 1 providing descriptions of disease-associated mutations within the open and closed state models of KCNQ1. Supplementary Figure 1 providing a topology plot for KCNQ1 with annotations that summarize the differences between the open and closed state KCNQ1 models with each other and with the corresponding K_1.2 structures. Protein Data Bank format atomic coordinate files for KCNQ1’s open and closed state models. This material is available free of charge via the Internet at http://pubs.acs.org.

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