The KCNQ1 (Kv7.1) COOH Terminus, a Multitiered Scaffold for Subunit Assembly and Protein Interaction

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The Kv7 subfamily of voltage-dependent potassium channels, distinct from other subfamilies by dint of its large intracellular COOH terminus, acts to regulate excitability in cardiac and neuronal tissues. KCNQ1 (Kv7.1), the founding subfamily member, encodes a channel subunit directly implicated in genetic disorders, such as the long QT syndrome, a cardiac pathology responsible for arrhythmias. We have used a recombinant protein preparation of the COOH terminus to probe the structure and function of this domain and its individual modules. The COOH-terminal proximal half associates with one calmodulin constitutively bound to each subunit where calmodulin is critical for proper folding of the whole intracellular domain. The distal half directs tetramerization, employing tandem coiled-coils. The first coiled-coil complex is dimeric and undergoes concentration-dependent self-association to form a dimer of dimers. The outer coiled-coil is parallel tetrameric, the details of which have been elucidated based on 2.0 Å crystallographic data. Both coiled-coils act in a coordinate fashion to mediate the formation and stabilization of the tetrameric distal half. Functional studies, including characterization of structure-based and long QT mutants, prove the requirement for both modules and point to complex roles for these modules, including folding, assembly, trafficking, and regulation.

The KCNQ channels represent a subfamily of voltage-gated K+ (Kv) channels, whose members (Kv7.1-5) are expressed in a wide variety of tissues. These channels play a major role in brain and cardiac excitability through the modulation of the cardiac potential waveform, the regulation of action potential generation and propagation, the tuning of neuronal firing patterns, and the modulation of neurotransmitter release (1, 2). Mutations in human KCNQ genes lead to major cardiovascular and neurological disorders, such as the cardiac long QT syndrome or neonatal epilepsy (3).

Like all Kv channels, the KCNQ α subunits share a common core structure of six transmembrane segments with a voltage-sensing domain (S1–S4) and a pore domain (S5–S6) (see Fig. 1A), likely to approximate the mammalian Kv1.2 channel α structure described by MacKinnon and co-workers (4). Often, KCNQ α is in complex with the integral membrane auxiliary subunit, known as KCNE1, IsK, or MinK. This protein alters channel properties, such as single channel conductance and activation kinetics while increasing channel density in the plasma membrane (5).

Several structural features of the Kv7 α family members distinguish them from the larger Kv channel superfamily. In particular, their primary sequence encodes a large (300–500 residues), intracellular COOH terminus (Fig. 1A). Sequence analysis predicts four helical regions (dubbed A–D) present in all family members (6). Helices C and D are thought to form coiled-coil assemblies. Yeast two-hybrid screens for proteins interacting with the COOH terminus revealed CaM, the ubiquitous Ca2+-sensitive protein, as a binding partner (6–8). Moreover, CaM constitutively associates with the channel (6–10). Helices A and B encode the binding site for CaM, and CaM association is required for proper channel assembly and function (6, 7, 9). LQT mutations that disrupt or significantly weaken the CaM interaction result in little or no complex and little channel expression in live cells (8, 11). Thus, CaM acts as an additional auxiliary subunit of the KCNQ channel complex. Additionally, Ca2+-CaM is a Ca2+-sensitive sensor for KCNQ1 function, transducing Ca2+ signals to stimulate IKS channels and producing a Ca2+-dependent left shift in the voltage dependence of channel activation. This Ca2+-sensitive IKS-current stimulation could increase the cardiac repolarization reserve, preventing the risk of ventricular arrhythmias.

Biochemical and functional studies have identified Kv7 COOH-terminal regions important for channel tetramerization and trafficking (12–16) based on deletion, truncation, and mutagenesis. Little work has directly examined on a protein level the postulated structural modules and their functional correlates. What precisely is the tertiary and quaternary organization of the COOH terminus, and what does it do that dis-
tistinguishes it from the other Kv channel subfamilies? Using a recombinant bacterial co-expression system (8), we have dissected the KCNQ1 COOH terminus protein complex. Our findings suggest that the KCNQ1 COOH terminus may be divided into two parts, where the membrane proximal half is important for functional expression, folding, and gating of the channel but not oligomerization, whereas the membrane distal half directs folding, oligomerization, partner specificity, and trafficking to the plasma membrane. This COOH terminus is a multifunctional platform, employing relatively simple structural modules to assemble a channel with high specificity in an apparent hierarchical manner.

EXPERIMENTAL PROCEDURES

General Subcloning and Mutagenesis

For bacterial expression vectors, PCR was used to engineer BamHI and NotI restriction sites onto the specific human KCNQ1 gene fragments. PCR product was ligated into doubly digested (BamHI and NotI) modified pETDuet vector. This pETDuet vector contains a His tag sequence and TEV protease recognition sequence upstream to multiple cloning site I with CaM inserted into multiple cloning site II. Positive clones were identified by restriction analysis and subsequently sequenced in all DNA constructs described below. In addition, all site-directed mutagenesis was performed by the QuikChange (Stratagene) method. Altered sequences were confirmed by DNA sequencing.

Subcloning of Δ Helix C—The KCNQ1 sequence in a pcDNA3 vector was used as a template for QuikChange, where primers were designed to delete helix C (residues 548–565). This modified vector then served as the template for PCR of an insert for the pETDuet vector.

Subcloning of Δ Loop—Construction was in two steps. First, PCR was used to engineer BamHI and EcoRI restriction sites onto a KCNQ1 gene fragment (residues 352–386). PCR product was ligated into a doubly digested (BamHI and EcoRI), modified pETDuet vector. Then PCR was used to engineer EcoRI and NotI restriction sites onto an additional gene fragment (residues 504–622). This PCR product was ligated into doubly digested (EcoRI and NotI) modified pETDuet vector already contained the fragment encoding residues 352–386.

Subcloning of COOH Terminus-GCN4-LI Chimera—Subcloning was in two steps. First, PCR was used to engineer BamHI and EcoRI restriction sites onto the KCNQ1 gene fragment (residues 352–594). PCR product was ligated into doubly digested (BamHI and EcoRI) modified pETDuet vector. Then the GCN4-LI sequence (encoding IEDKLEILSFLHYHENE-LARIKULLLG) was amplified by primers containing EcoRI and NotI sites in the 5’- and the 3’-flanking regions, respectively. GCN4-LI PCR product was ligated into doubly digested (EcoRI and NotI) modified pETDuet vector that already contained DNA encoding residues 352–594. Subsequently, the EcoRI restriction site was deleted by mutagenesis.

Expression and Purification of the Various KCNQ1 COOH Terminus-CaM Complexes

Transformed Escherichia coli BL-21 Tuner (Novagen), containing the “RIL” Codon Plus™ plasmid (Stratagene) cells were grown at 37 °C in LB medium, containing 100 μg/ml ampicillin, and 34 μg/ml chloramphenicol. Upon reaching an A₆₀₀ of 0.3, the temperature was lowered to 16 °C, and growth continued until the culture reached an A₆₀₀ of 0.6. Protein expression was induced with 135 μM isopropyl-1-thio-β-D-galactopyranoside. Cells were harvested after 14 h by centrifugation, frozen, and suspended in 120 ml of lysis buffer, buffer L (300 mM NaCl, 50 mM sodium phosphate, pH 8, 1 mM phenylmethylsulfonil fluoride). After lysis by a microfluidizer (Microfluidics), cell debris was removed by centrifugation at 20,000 × g. The soluble fraction was loaded onto a pre-equilibrated metal chelate Ni²⁺-CAM (Sigma) column (buffer A: 300 mM NaCl, 50 mM sodium phosphate, pH 8) at a flow rate of 1.0 ml/min. The column was washed with buffer A, containing 10 mM imidazole, until a stable base line was achieved. After elution with buffer A, supplemented with 125 mM imidazole, the protein eluate was then subjected to TEV protease in a ratio of 1:150. The proteolysis continued for 12 h. Subsequently, the sample was diluted 4-fold with 10% glycerol and loaded onto a pre-equilibrated SP-Sepharose (Amersham Biosciences) column (buffer S: 50 mM NaCl, 20 mM Tris, pH 7.5). The column was then washed with buffer S, and fractions were eluted with a shallow linear gradient of buffer S, containing 50–600 mM NaCl. Fractions were pooled and applied to a pre-equilibrated Superose 6 gel filtration column (Amersham Biosciences) with buffer F (150 mM NaCl, 20 mM Tris, pH 7.5, 1 mM dithiothreitol). The elution peak was concentrated to 3 mg ml⁻¹ using spin concentrators (Vivascience), divided into aliquots, and flash-frozen in liquid N₂. For purification of Δ helix D, Δ helices C-D, and Δ loop-CaM complexes, a Superdex-200 gel filtration column was employed instead of the Superose-6 gel filtration column.

Subcloning, Expression, and Purification of GST-Helix C and GST-Helices C-D

Both KCNQ1 gene fragments (residues 535–572 or 535–622, respectively) were amplified by PCR with primers containing BamHI and NotI sites in the 5’- and 3’-flanking regions, respectively. PCR product was ligated into doubly digested (BamHI and NotI) pGEX-4T-1 vector (Amersham Biosciences). Expression was as above. Cells were suspended in phosphate-buffered saline (PBS) (pH 7.4) with 1 mM phenylmethylsulfonil fluoride. After lysis, cell debris was removed by centrifugation. The soluble fraction was loaded onto a pre-equilibrated glutathione-Sepharose column (buffer A: PBS, pH 7.4), at a flow rate of 1.0 ml/min. The column was washed with buffer A and eluted with buffer B (100 mM Tris pH 8, 100 mM NaCl, 20 mM glutathione), and the protein was applied to a pre-equilibrated Superdex 200 gel filtration column with buffer F. The elution peak was concentrated and stored as above.

Expression and Purification of Selenomethionine Protein

The selenomethionine Δ loop-CaM complex was prepared by inhibition of the methionine pathway. Inoculum was grown from a single transformed colony in 10% LB medium. The cells were pelleted, and media supernatant was removed, prior to the addition to 2 liters of New Minimal Medium, containing Kao and Michayluk vitamin solution (Sigma), 100 μg/ml ampicillin, and 34 μg/ml chloramphenicol. Upon reaching an A₆₀₀ of 0.3,
the temperature was lowered to 16 °C. Lysine, phenylalanine, and threonine (100 mg/liter), isoleucine, leucine, and valine (50 mg/liter), and DL-selenomethionine (50 mg/liter) were added 45 min before induction, when the culture reached an A600 of 0.6. Protein expression was induced with 135 μM isopropyl 1-thio-β-D-galactopyranoside over a 16-h period. Purification of the selenomethionine protein was similar to that of the native protein, except that 5 mM β-mercaptoethanol was added to all solutions to prevent oxidation of the selenomethionine derivative protein.

**Peptide Purification**

The helix D synthetic peptide (residues 585–621, acetylated and amidated at the NH2 and COOH termini, respectively) was purified by reverse-phase high pressure liquid chromatography using a C18 column (Vydac) with a shallow acetonitrile gradient of 35–60% (both solvents were supplemented with 0.05% trifluoroacetic acid).

**Analytical Ultracentrifugation**

Sedimentation velocity analysis of the different protein samples were carried out at 40,000 or 50,000 rpm and 5 °C (except Δ helices C-D and point mutants that were centrifuged at 10 °C) in an XL-A analytical ultracentrifuge (Beckman-Coulter Inc.) with a UV-visible optics detection system, using an An60Ti rotor and 12-mm double sector centerpieces. All of the proteins were equilibrated in 20 mM Tris-HCl, 150 mM NaCl, 0.1 mM dithiothreitol, pH 7.5, buffer except for CT-GCN4LI, which was in PBS. The sedimentation velocity runs were done at different protein concentrations ranging from 0.1 to 1.5 mg/ml. Sedimentation profiles were registered every 5 min at the appropriate wavelength (ranging from 230 to 280 nm). The sedimentation coefficient distributions were calculated by least-squares boundary modeling of sedimentation velocity data using the c(s) method (17) as implemented in the SEDFIT program. The corresponding standard s-values (s20,w) were obtained from the experimental s-values upon correction for density, viscosity, and protein concentration (18), using the SEDNTERP program (19).

Sedimentation equilibrium experiments were carried out at multiple speeds (9000, 13,000, 22,000, 32,000, 40,000, and 43,000 rpm) and wavelengths (230, 238, 250, and 280 nm) with a UV-visible optics detection system, using an An60Ti rotor and 12-mm double sector centerpieces. All of the proteins were equilibrated in 20 mM Tris-HCl, 150 mM NaCl, 0.1 mM dithiothreitol, pH 7.5, buffer except for CT-GCN4LI, which was in PBS. The sedimentation velocity runs were done at different protein concentrations ranging from 0.1 to 1.5 mg/ml. Sedimentation profiles were registered every 5 min at the appropriate wavelength (ranging from 230 to 280 nm). The sedimentation coefficient distributions were calculated by least-squares boundary modeling of sedimentation velocity data using the c(s) method (17) as implemented in the SEDFIT program. The corresponding standard s-values (s20,w) were obtained from the experimental s-values upon correction for density, viscosity, and protein concentration (18), using the SEDNTERP program (19).

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Analytical Gel Filtration Chromatography

Proteins were injected onto a Superose 6 10/300 GL or Superdex 200 10/300 GL or Superdex 75 10/300 GL (Amersham Biosciences) column. Proteins were eluted with buffer F at a flow rate of 0.5, 0.7, or 0.8 ml/min for Superose 6, Superdex 200, or Superdex 75 columns, respectively.

**Crystallization**

Crystals were grown in 12–16% polyethylene glycol 8000, 0.1 M Tris, pH 7–8, at 19 °C by sitting drop vapor diffusion. Equal volumes (1 μl) of frozen stock protein (15 mg/ml) were mixed with reservoir solution. Native crystals appeared after 2 weeks. Heavy atom soaks were prepared by adding 1 mM K2ReCl6 directly to drops containing crystals for 24 h. Four selenomethionine protein crystals appeared after several months. Crystals were cryoprotected by sequential dilution with reservoir solution, including 24% glycerol, and then loop mounted and flash frozen in liquid N2.

**Structure Determination**

Diffraction data were obtained under standard cryogenic conditions and processed with HKL (23). We executed a single wavelength anomalous diffraction experiment on the selenomethionine crystals, using the anomalous selenium peak. Heavy atom site location and experimental phases calculation was performed with CNS (24) and SHARP (25, 26), respectively. Multiple isomorphous replacement with anomalous scattering from the selenium and rhenium derivative data sets against the native data set were used for experimental phase calculation. Subsequent density modification with SOLOMON (26) gave electron density maps of excellent quality. An initial model was built with ARP/wARP (27) and subsequently refined with Refmac5 (28) with rounds of model building.
Structural and Functional Studies of the KCNQ1 COOH Terminus

Electrophysiology

Recordings were performed using the whole cell configuration of the patch clamp technique. Signals were amplified using an Axopatch 200B patch clamp amplifier (Axon Instruments), sampled at 2 kHz and filtered at 800 Hz via a 4-pole Bessel low pass filter. Data were acquired using pClamp 8.2 software in conjunction with a DigiData 1322A interface. The patch pipettes were pulled from borosilicate glass (Warner Instrument Corp.) with a resistance of 4–7 megaohms. The intracellular pipette solution contained 130 mM KCl, 1 mM MgCl₂, 5 mM K₂ATP, 5 mM EGTA, 10 mM HEPES, adjusted with KOH to pH 7.4 (290 mOsM). The external solution contained 140 mM NaCl, 4 mM CaCl₂, 1.2 mM MgCl₂, 11 mM glucose, 5.5 mM HEPES, adjusted with NaOH to pH 7.4 (310 mOsM). Series resistances (8–15 megaohms) were compensated (75–90%) and periodically monitored.

Immunocytochemistry

COS7 cells were grown on 13-mm diameter coated glass coverslips in 24-well plates. Cells were rinsed for 5 min in PBS and subsequently fixed for 20 min in 4% paraformaldehyde in PBS. Following extensive washes in PBS, the cells were permeabilized by incubation with 10% normal goat serum in PBS containing 0.2% Triton X-100 for 20 min. Cells were then washed twice for 10 min each in PBS containing 1% normal goat serum. Cells were incubated at 4 °C overnight with a rabbit anti-KCNQ1 (Alomone Laboratories) channel or mouse anti-FLAG (M2; Sigma) antibodies. After a 3 × 5-min wash in PBS, cells were incubated for 1 h at room temperature with secondary antibodies, CY2-conjugated anti-rabbit IgG (1:200; Jackson Immunoresearch) and RRX-conjugated anti-mouse IgG (1:1000; Jackson Immunoresearch), respectively. Cells were viewed, and digital images were taken using a Zeiss 510 META confocal microscope using the 488-nm argon or 543-nm HeNe excitation laser lines.

Cell Surface Biotinylation Assays

Biotinylation of surface proteins was carried out by incubating cells with 1 mg/ml EZ-Link Sulfo-NHS-SS-Biotin (Pierce) in PBS containing 1 mM phenylmethylsulfonyl fluoride and 5 mM EDTA (solution A) for 30 min at room temperature. The reaction was terminated by incubating cells for 5 min in solution A containing 20 mM glycine, followed by three washes in solution A. Cells were lysed in buffer P (50 mM Tris-Cl, pH 7.5, 20 mM HEPES, pH 7.5, 150 mM NaCl, 5 mM EDTA, 1.5 mM MgCl₂, 10% glycerol (w/v), 1% Triton X-100 (v/v)), containing 1 mM phenylmethylsulfonyl fluoride and 10 μl/ml protease inhibitor mixture (Sigma), for 1 h at 4 °C under rotation. Cell lysate was cleared by centrifugation, and biotinylated proteins were precipitated by incubation with streptavidin-agarose beads (Pierce) overnight at 4 °C. The beads were washed six times with buffer P. Proteins were eluted by incubation with sample buffer at room temperature and then resolved by SDS-PAGE and Western blotted.

Inductively Coupled Plasma (ICP) Spectrometry

Peptide samples were suspended at room temperature in a mixture of metal chloride salts containing Tris (10 mM), pH 7.5, 100 mM NaCl, and a 20 μM concentration of the indicated metals for 2 h at 0.35 mg/ml. Subsequently, peptide was dialyzed for 16 h against the same buffer without metals. Samples were prepared for analysis by incubation in a hot water bath at 80 °C for 2 h with the following addition. One-mL volumes of concentrated nitric acid were added to 1–2-mL sample batches in polypropylene 50-ml tubes. Liquid residues were taken up in deionized water and made up to a 10-ml volume with the same solvent. Metal concentration in the acid extract was determined by ICP atomic emission spectrometry. An ICP atomic emission spectrometer with cross-flow nebulizer was used (Spectroflame Modula E from Spectro, Kleve, Germany).

RESULTS

Oligomeric State of the KCNQ1 COOH Terminus-CaM Complex—Work by Pitt and co-workers (11) suggested previously that a soluble KCNQ1 COOH terminus-CaM complex is tetrameric using chemical cross-linking, consistent with the universal tetrameric structure of Kv channels. Using a similar molecular complex, comprising soluble COOH terminus and constitutively bound CaM (Fig. 1, B and C), we probed its oligomerization state (i.e. its molecular weight) by sedimentation equilibrium analysis. The results indicate that the complex is best modeled as a tetramer with four subunits of KCNQ1 COOH terminus and four bound molecules of CaM (Fig. 1D). Data analysis, assuming a single sedimenting solute species, yields a buoyant molecular weight of 54,000 ± 3000 (solid line). This value is very close to that calculated from the addition of the buoyant molecular weights of four molecules of the COOH terminus (4 × 8400) and four of CaM (4 × 4500). It is not compatible with a 4:2 complex (dotted line).

Sedimentation velocity studies that evaluate homogeneity/dispersity of the same COOH terminus-CaM complex provide convincing evidence that the primary species (89–95% of the total loading concentration, varying by experiment) is this very same tetrameric complex (8.2 S) with a minor species that represents a higher molecular weight aggregate (11.4 S) (Fig. 1E). Importantly, there is no evidence of a free CaM species, indicating that CaM is stably bound, since measurements of CaM alone exhibit a symmetrical sharp distribution with a standard s value of 2.8 S, compatible with the expected value for monomeric CaM (Fig. 1E). No such peak is detected in the profile of the COOH terminus-CaM complex. The hydrodynamic behavior of the 4:4 COOH terminus-CaM complex (fractional ratio f₁₀₀ = 1.8; see Ref. 29) is compatible with the solute being an elongated, nonglobular protein. Moreover, calcium is not required for this assembly, since size exclusion chromatography of a COOH terminus-CaM₁₂₃₄ complex, utilizing a CaM mutant that does not bind calcium, behaves like a COOH terminus-CaMᵢₓᵢₓ complex (Fig. 1F).

CaM Organization—To explore the possibility that CaM may induce oligomerization by its association with the COOH-terminal proximal half, we coexpressed CaM with a protein segment spanning helix A and B but truncated afterward (Δ helices C-D plus CaM) (Fig. 1, B and C). Sedimentation velocity and size exclusion chromatography indicates that it behaves as a 1:1 complex of CaM to Δ helices C-D; i.e. CaM does not cross-link COOH terminus subunits, inducing their oligomerization.
neither in the presence of calcium (1 mM) nor in its absence (5 mM EDTA). Since the intervening loop between helices A and B is long, binding by CaM of these two helices may be in either parallel or anti-parallel helical arrangements. Nevertheless, the loop is not necessary for CaM binding, since its removal does not prevent formation of the loop-CaM complex (Fig. 1, B and C), consistent with an anti-parallel helix bundle as the target of CaM.

Role of the Distal COOH Terminus (Helices C and D)—Earlier studies on KCNQ have assigned regions from the distal COOH terminus, encompassing sequences predicted to form the two helices C and D with what has been called an assembly or tetramerization domain. Both of these predicted helices are amphipathic and have varying probabilities to form coiled-coils. We tested the basic prediction of helicity for helix C. Using CD spectroscopy, we measured the spectrum of helix C peptide (Fig. 2, A and B). The results indicate clearly that helix C takes an α-helical conformation (Fig. 2C). Having established that this module is helical, we then asked if it self-associates. Sedimentation equilibrium studies demonstrate that helix C dimerizes, presumably by a coiled-coil interaction, as supported by reversible thermal denaturation, a property characteristic of coiled-coils (Fig. 2C, inset, and supplemental Fig. 2 for the heptad repeat assignment), while further associating as a dimer of dimers.

We then asked what helix C will do in the context of the COOH terminus-CaM but without the confounding action of helix D. Sedimentation studies, both equilibrium and velocity, of COOH terminus Δ helix D-CaM (Fig. 2, A and B) unambiguously demonstrate that the complex is best modeled as a dimer (i.e. two COOH terminus subunits bound to two CaMs with a buoyant molecular weight half that of the full-length COOH terminus-CaM complex) (solid line gradient in Fig. 2E). The data cannot be accommodated by a gradient expected for a 4:4 complex (Fig. 2E, dashed line). The hydrodynamic behavior of Δ helix D-CaM complex (s value of the main sedimenting species was 5 s, open circles in Fig. 2F) also deviates from that expected for a rigid globular solute (frictional ratio f/f₀ = 1.7). We conclude that, like the isolated helix C module, helix C in its COOH-terminal context directs dimerization. It seems likely that during biosynthesis of the intact channel, this dimeric module forms a dimer of dimers due to high local concentration.

To probe the function of helix C in a cellular context, we recorded currents from Chinese hamster ovary cells transfected with a helix C deletion channel construct. This
deletion obviates functional expression (i.e. production of a detectable K^+ current) (Fig. 2G). The expressed protein appears to be retained intracellularly, as shown by immunocytochemical labeling and cell surface biotinylation assays (Fig. 2, H and I, respectively).

What function does helix D perform? To address this question, we measured CD spectra of helix D peptide (residues 585–621), confirming its helical conformation (data not shown). We examined the peptide’s oligomeric state using both sedimentation equilibrium and velocity. The results show a tetrameric species (single species with a molecular weight of 18,400 ± 1500; see Fig. 3A and supplemental Fig. 1B) over a wide range of concentrations (10–100 μM) versus a monomeric one (molecular weight of 4200), proving that helix D forms a stable tetrameric coiled-coil with a globular shape (frictional ratio f/f0 = 1.3), in contrast to helix C. Next, we purified a KCNQ1 COOH terminus Δ helix C-CaM complex (Fig. 3, B and C), allowing us to directly examine the action of helix D without the potential confounding effects of helix C, shown above to also self-associate. Sedimentation velocity establishes that the Δ helix C-CaM complex tetramerizes much like the WT COOH terminus-CaM complex (Fig. 3D). Therefore, helix D induces a tetramerization state upon the KCNQ1 COOH terminus, independent of the helix C segment.

Deletion of helix D abrogates KCNQ1 current (12), underlining its requirement for proper function. If the helix is perturbed by substitution of proline at position 609 (Ile → Pro), Roden and co-workers (15) report that no functional current is detectable, and the channel is found intracellularly. Presumably, this mutation disturbs the helix sufficiently to disrupt the predicted coiled-coil. These investigators ascribed the elimination of functional current to the essential role of helix D in trafficking the channel to the plasma membrane. We there-
fore examined the consequences of this mutation in our expression system. The mutated protein was insoluble (data not shown), similar to that seen when CaM is not coexpressed or its binding sites are perturbed (8, 11). This result hints that viable channel folding and assembly requires properly folded helix D. If helix D directs tetramerization, then perhaps we could replace it with an analogous coiled-coil that performs the same task. Such coiled-coils have been used in similar contexts (30). We utilized GCN4-LI, a modified GCN4 coiled-coil that forms a parallel, tetrameric coiled-coil (31). The GCN4-LI module was introduced into KCNQ1 COOH terminus at residue 594, preserving completely the first helical turn of helix D (Fig. 3B). This chimeric protein was soluble (Fig. 3C), and a fraction of it behaved like WT in sedimentation velocity experiments (Fig. 3D), corroborated by size exclusion chromatography (supplemental Fig. 1C), but when transfected into cells it produced no current and was retained intracellularly without evidence of cell surface expression (Fig. 3, E–G).

**Crystallographic Studies of the Helix D Tetramer**—What is the molecular mechanism by which helix D acts? We obtained the atomic details of this molecular complex using x-ray crystallography. Crystallization trials with COOH terminus Δ loop-CaM complex (Fig. 1, B and C) yielded small crystals after several weeks. NH2-terminal sequential of the crystallized protein identified the sample as KCNQ1 starting with residue 574, just upstream of helix D. We conclude that a contaminating protease cleaved the Δ loop protein in the predicted random coil region between helices C and D, allowing the helix D assembly to crystallize. Single wavelength anomalous diffraction and multiple isomorphous replacement with anomalous scattering diffraction experiments using a selenomethionine protein crystal and a rhenium-soaked crystal, respectively, produced excellent electron density maps to 2.0-Å Bragg spacings (Table 1). The current refined model includes KCNQ1 residues 585–621 with good stereochemistry, 52 solvent molecules, and an Rfactor of 25.5%. Residues 574–584 are not observed and must be disordered.

The crystallographic studies reveal a tetrameric parallel-oriented coiled-coil quaternary structure (Fig. 4A). The crystallo-graphic asymmetric unit contains a dimeric coiled-coil, where-upon a 2-fold crystallographic dyad generates the presumed biological tetrameric unit. Thus, formally the helix D assembly is a dimer of dimers, with pseudo-4-fold symmetry that is close to perfect C4 symmetry. The coiled-coil cylinder is 50 Å long with a diameter of about 27 Å, composed of helices of nine turns or about four and half heptads. Coiled-coil parameterization analysis by the program TWISTER (32) shows the assembly to have the canonical heptad (7/2) repeat, left-handed supercoiling, and other Crick parameters (Table 2) with the signature of a classical tetrameric parallel coiled-coil. The buried residues of the a and d heptad positions maintain alternating perpendicular and parallel knobs-in-hole packing, respectively, as computed by Socket (33). The perpendicular and parallel terms refer to packing angles for the side chains, as defined by Harbury et al. (31). This arrangement was observed for GCN4-LI. The tetramer is capped at its carboxyl-terminal end with a histidine ring coordinating a metal ion (Fig. 4B).

The coiled-coil interactions constituting the hydrophobic core and the solvent-exposed residues involved in interhelical interactions are summarized schematically in Fig. 4, C and D. The a and d positions of the heptad are taken by branched, apolar residues generally conserved in both the KCNQ1 orthologs and the Kv7 subfamily. Notably, helix D of KCNQ1, as seen in the structure and sequence alignment (supplemental Fig. 2), has a longer coiled-coil, by one heptad repeat or two helical turns longer than the other KCNQ members. The one exception to the branched hydrophobic residues in positions a and d is the capping histidine found in position a. Kim and co-workers (31), in studies using the dimeric GCN4 as a scaffold, proposed a set of rules for predicting the oligomeric state of a coiled-coil. They posited that unbranched Cβ (Leu) residues at the a position with branched Cβ residues (Ile, Val) at the d position will favor tetrameric assemblies. Contrary to this proposal, the tetrameric KCNQ1 helix D d positions have almost as many unbranched residues as branched.

The helix D assembly buries a total of 6540 Å2, an area typical of coiled-coil architectures, and reflects a character more akin to globular protein core than standard protein-protein interfaces. We measured the stability of this coiled-coil using thermal denaturation, assayed by CD spectroscopy (Fig. 4E). The measurements indicate a Tm of ~35 °C with reversibility of the process. We note this seemingly low value for tetrameric coiled-coils, despite efficient interior packing (34, 35).

Having established the coiled-coil architecture of both helix C and D modules separately and determined their thermal stability, we examined the consequences of having them in the tandem arrangement found in the intact protein. Specifically, would the relatively low Tm values and reversible folding/un-
Structural and Functional Studies of the KCNQ1 COOH Terminus

A. helix D

B. C-terminus

C. Δ helix C

D. Sedimentation Coefficient

E. WT KCNQ1

F. F

G. normalized ratio

WT KCNQ1        KCNQ1-GCN4LI

0.00 0.02 0.04 0.06 0.08 0.10

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folding seen for the individual modules be the same in this type of tandem organization? To this end, we constructed a GST fusion protein, beginning at helix C and extending to just beyond the end of helix D, which was expressed, cleaved from the GST, and purified (Fig. 4, F and G). This protein was characterized by analytical size exclusion chromatography, which suggests that it is most likely a tetrameric complex, as expected (supplemental Fig. 1D). We further analyzed the complex with CD spectroscopy, where it shows the anticipated helical structure (Fig. 4H). Thermal denaturation experiments demonstrate a more complex denaturation curve than that seen for both of the individual modules with a marked increase in the Tm (about 45 °C) and two different phases in the unfolding, one in the 30–40 °C range with a later phase occurring between 55 and 65 °C (Fig. 4H, inset). Moreover, the folding was not reversible. We conclude that the tandem coiled-coil architecture confers appreciable stability, including a possible kinetic deterrent to unfolding/dissociation of the distal COOH terminus.

From examination of the helix D complex architecture that is both elegantly simple and symmetric, we discern three pertinent molecular features. The first (i) is the interface by which the polypeptide chain folds with the others, forming the hydrophobic core of the complex, primarily but not exclusively mediated by the interior a and d heptad positions. The second feature (ii) is the intersubunit interface and surface that contains stereochemically specific interactions (e.g. hydrogen bond and van der Waal interactions with a special emphasis on salt bridge networks). This feature furnishes biological specificity in helix D complex formation, possibly specifying homotypic or heterotypic combinations in the Kv7 subfamily. The third feature (iii) is the solvent-accessible molecular surface available for protein-protein interactions with other proteins, which is likely to play an important role in trafficking and regulation of the channel (15, 36). We probed these three features with a functional survey of 17 different structure-based point mutations in the helix D region, designed to perturb relevant structural determinants (Table 3). Certain residues are likely to contribute to more than one molecular feature.

When we sought to disrupt the coiled-coil hydrophobic core (molecular feature i), only specific changes had an impact. We focused not just on residues of the a and d positions but also on buried residues from other heptad positions. For example, G589M, L602I, or L609P had no significant effect, as assessed by current recorded from transfected cells (Table 3). These results suggest a measure of plasticity in the helix D complex's folding determinants. In contrast, mutations previously performed by Roden and co-workers (15) (L606P/L613P, L606D/L613D, I609P, and L602D/I609D) that essentially abrogate cell surface

### TABLE 1

Crystallographic statistics

|                     | Native | SeMet | Re |
|---------------------|--------|-------|----|
| Data collection and phasing statistics |        |       |    |
| Wavelength (Å)      | 0.9792 | 0.9788| 0.9788 |
| Space group         | P3121  | P3121 | P3121 |
| Unit cell parameters (Å) | a = b = 51.3, c = 71.1 | a = b = 51.6, c = 72.1 | a = b = 51.4, c = 71.0 |
| Total reflections   | 67,168 | 131,874| 257,557 |
| Unique reflections  | 7656   | 1795  | 7214 |
| Completeness (%)    | 99.2 (99.5) | 96.3 (99.5) | 99.5 (100) |
| Rmerge (%)          | 6.5 (33.7) | 5.0 (12.6) | 5.7 (13.2) |
| Rfree (%)           | 19.6 (3.36) | 47.4 (21.0) | 52.9 (26.0) |
| Resolution range (Å)| 50–2.0 | 50–2.5 | 50–2.5 |
| Fc/〈Fc〉           | 76.4/9 | 71.0/12.1 |
| Phasing power       | 1.88   | 0.743 | 0.145 |
| Figure of merit     | 0.3    | 0.3   | 0.3   |
| Beamsline (ESRF)    | 0.3    | 0.3   | 0.3   |

Refinement statistics

|                     | Native | SeMet | Re |
|---------------------|--------|-------|----|
| No. of reflections (working/test) | 7,283/354 | 7,283/354 | 7,283/354 |
| a (Å)               | 2.0    | 2.0   | 2.0   |
| Rwork/Rfree (%)     | 22.3/25.5 | 22.3/25.5 | 22.3/25.5 |
| r.m.s. deviation from ideality |          |       |      |
| Bond lengths        | 0.008  | 0.008 | 0.008 |
| Bond angles         | 1.36   | 1.36  | 1.36  |
| B factors (Å2)      | 1.850  | 1.850 | 1.850 |
| Average B factor (Å2) | 32.3   | 32.3  | 32.3  |
| No. of protein atoms/solvent | 642/52 | 642/52 | 642/52 |

*Values of the highest resolution shell are given in parentheses.

**R**merge = ΣhklΣit[Iit] - (Iit)/ΣhklΣit[Iit], where Iit is the intensity of a reflection, and (Iit) is the average of all observations of this reflection.

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**FIGURE 3.** Oligomeric states and functional data of helix D module. A, sedimentation equilibrium gradient of helix D peptide. The solid line shows the best fit gradient corresponding to a tetramer, whereas the other lines denote trimer, dimer, and monomer gradients, as indicated. B, schematic depiction of recombinate constructs. All of the constructs were coexpressed with CaM C. SDS-PAGE of relevant proteins. Samples shown are composite from different gels. All samples were run in the presence of 1% n-dodecyl β-D-maltopyranoside (DDM) distribution. All proteins are in complex with CaM. Experiments were done at 50,000 rpm and 10 °C. The s values have been corrected to standard conditions. The estimated molecular masses for the 7.6, 12.4, and 16.3 s peaks are 170, 350, and 530 kDa, respectively. E, representative current traces of WT KCNQ1 and the KCNQ1-GCN4LI mutant. From a holding potential of −90 mV, Chinese hamster ovary cells were stepped for 3 s from −70 to +60 mV in 10-mV increments and repolarized for 1 s at −60 mV. F, immunocytochemical mapping of COS cells transfected with WT KCNQ1 and KCNQ1-GCN4LI mutant plasmids and detected with rabbit anti-KCNQ1 antibodies. Results clearly show that although the KCNQ1-GCN4LI mutant exhibits essentially intracellular localization with strong accumulation in the endoplasmic reticulum, the WT channel protein displays intracellular but also significant plasma membrane distribution, denoted by arrowheads. Images were taken using a Zeiss 510 metaconfocal microscope with a 458-nm excitation argon laser line. G, cell surface biotinylation of KCNQ1-GCN4LI. FLAG-tagged WT KCNQ1 and KCNQ1-GCN4LI fusion were expressed in HEK 293 cells, labeled with NHS-LC-Biotin, pulled down with streptavidin beads, and probed with anti-FLAG antibodies. Signals were corrected to channel protein input, normalized to WT, and expressed as ratios (n = 3; **, p < 0.001, two-tailed t test).
Structural and Functional Studies of the KCNQ1 COOH Terminus

A

B

C

D

E

helix D

F

C-terminus

GST-helices C-D

G

helices C-D

H

helices C-D

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of coiled-coils and may facilitate specificity of assembly in two ways; first, determination of the partner subunit, and second, number of subunits in the assembly. Charged residues confer specificity by requiring an oppositely charged residue to be in proximity and by selecting against a partner whose proximal residue is like charged. Nonetheless, obvious empirical rules are not easily derived, since many expected perturbations of this molecular feature like T600L, Q601A, or D603G appear to have little effect (Table 3).

To probe the importance of the complex’s solvent accessible residues vis-à-vis their potential role to generate a platform for interacting proteins (molecular feature iii), we mutated almost all of the helix D residues with accessibility greater than 40% (Table 3). The preponderance had little or no effect. However, three mutations, G589D, A590W, and N593G, located in the NH$_2$-terminal third of helix D reduced the expressed K$^+$ currents in the transfected cells.

KCNQ1 has been implicated genetically in long QT syndrome. Lesions responsible for this pathophysiological phenotype are found throughout the structural gene, and several localize to helix D (supplemental Table 1). A representation showing their location on the complex’s molecular surface is depicted in Fig. 5A. Strikingly, these mutations predominantly cluster in the NH$_2$-terminal third of the assembly and include the loop upstream of the assembly, forming a contiguous surface. Curiously, all of the mutations involve non-$\alpha$ or $\delta$ coiled-coil position residues. Thus, they either perturb partner specification in assembly (e.g. R594H) or formation of protein-protein interaction interfaces or both. Apparently, they do not perturb folding and tetrameric assembly. One mutation, G589D, has been shown to abrogate binding of the A kinase-anchoring protein protein yotiao, important for sympathetic regulation of the cardiac action potential as mediated by $\beta$-adrenergic receptor signaling (36). Other possibilities could involve proteins important for trafficking of the channel complex to the plasma membrane (15).

We thus focused our characterization efforts on five point mutations of helix D that perturbed current expression, in an attempt to define their molecular mechanism. The residues’ locations on the helix D coiled-coil are graphically depicted in Fig. 5B. These mutations should impinge upon the molecular features described above. Specifically, G589D, a LQT mutation, along with A590W, N593G, and possibly H620S, may affect both intersubunit interfaces and the surface available for protein-protein interactions. Mutations L602M, H620S, and possibly A590W may affect the core of the complex. The current-voltage relationships (I-V curves) determined for the various mutants (Fig. 5, C and D) indicate that some (N593G, L602M, and H620S) give rise to moderately impaired current magni-

expression, are easily rationalized in light of the structure. All of these mutations change $\alpha$ or $\delta$ heptad position hydrophobic residues to either helix-kinking proline or a negative charge, almost certainly disrupting the coiled-coil assembly. On the other hand, when these same residues were replaced by alanine in that same study, surface expression approached that of WT, underscoring the complex’s plasticity. The alanine replacements are neutral; they do not negatively or positively select against heptad position partners, a principle clearly critical in coiled-coils (33, 37).

Hydrogen-bonded salt bridges play a prominent role in the intersubunit interface (molecular feature ii). For example, Arg$^{594}$ (c position in $\alpha$ subunit) hydrogen-bonds to Glu$^{596}$ (e position in $\beta$ subunit), which concomitantly engages Lys$^{598}$ (g position in $\alpha$ subunit) (Fig. 4, C and D). This network is typical of coiled-coils and may facilitate specificity of assembly in two ways; first, determination of the partner subunit, and second, number of subunits in the assembly. Charged residues confer specificity by requiring an oppositely charged residue to be in proximity and by selecting against a partner whose proximal residue is like charged. Nonetheless, obvious empirical rules are not easily derived, since many expected perturbations of this molecular feature like T600L, Q601A, or D603G appear to have little effect (Table 3).
tude, whereas the current magnitude of others (G589D and A590W) is severely diminished. Previously published reports for G589D conflict; Kass and co-workers (36) published data showing unimpaired K⁺ currents and normal plasma membrane location, whereas Roden and co-workers demonstrated complete K⁺ current inhibition, as we observe, and intracellular...
lar channel retention (15). In order to determine if these mutants’ current deficiency was due to problems with targeting, we performed cell surface biotinylation assays on four mutants. The results are shown in Fig. 5E. The mutants exhibit diminished but variable cell surface expression when compared with WT. To some degree, functional current expression correlates with cell surface expression.

We then asked if the mutations might be causing difficulties in assembly. We addressed this question by expressing and purifying the point mutants in complex with CaM in our recombinant system. These complexes were then studied by sedimentation velocity and/or size exclusion chromatography. The results are shown in Fig. 5, F and G. The WT COOH terminus-CaM complex provides a size and shape standard for the 4:4 tetrameric assembly with c(s) distribution centered around 8 s. We quantitated the fraction of mutant complex that sediments at the peak corresponding to a tetrameric assembly from the derived c(s) distribution of each individual sedimentation velocity experiment. The mutants G589D, A590W, L602M, form 76, 69, and 71% tetrameric assemblies, respectively, whereas the WT forms 88%. Generally, the mutants exhibit more polydispersity than the WT. Based on size exclusion chromatography, the mutants N593G and H620S behaved almost exactly like the WT. We conclude that all of the mutants appear to assemble into proper tetrameric complexes, albeit with slighter lower efficiency than WT. Therefore, their lower functional expression does not stem from flawed assembly but from either faulty trafficking or other defective regulation or both.

As noted earlier, the helix D complex is capped by a histidine ring that coordinates a metal ion. This appears to be a novel structural feature not yet observed in structures from natural sequences, although similar structures have been engineered for designer coiled-coils. First, we sought to identify the metal ion. To this end, we purified, desalted, and suspended a synthetic peptide of helix D in buffer containing appropriate cations, and/or size exclusion chromatography. The results are shown in Table 4. The WT COOH terminus-CaM complex provides a size and shape standard for the 4:4 tetrameric assembly with c(s) distribution centered around 8 s. We quantitated the fraction of mutant complex that sediments at the peak corresponding to a tetrameric assembly from the derived c(s) distribution of each individual sedimentation velocity experiment. The mutants G589D, A590W, L602M, form 76, 69, and 71% tetrameric assemblies, respectively, whereas the WT forms 88%. Generally, the mutants exhibit more polydispersity than the WT. Based on size exclusion chromatography, the mutants N593G and H620S behaved almost exactly like the WT. We conclude that all of the mutants appear to assemble into proper tetrameric complexes, albeit with slighter lower efficiency than WT. Therefore, their lower functional expression does not stem from flawed assembly but from either faulty trafficking or other defective regulation or both.

As noted earlier, the helix D complex is capped by a histidine ring that coordinates a metal ion. This appears to be a novel structural feature not yet observed in structures from natural sequences, although similar structures have been engineered for designer coiled-coils. First, we sought to identify the metal ion. To this end, we purified, desalted, and suspended a synthetic peptide of helix D in buffer containing appropriate candidate divalent metal salts (20 μM) or in buffer alone. Then samples were dialyzed extensively against buffer alone and analyzed by ICP spectrometry (Table 4). The data clearly indicate that the preferred metal ion was Cu2+. We expect that Zn2+ can bind as well (data not shown). Structurally, this identification makes good inorganic chemical sense, since Cu2+ is well suited for the observed square pyramid 5-coordination as a water molecule is bound above the metal ion. In addition, the coordination lengths are appropriate (2.17 Å). Other copper-binding proteins, such as laccase, an oxidoreductase from fungi (38), have similar stereochemistry, wherein four histidines chelate the copper.

![FIGURE 6. Quaternary structure model of the KCNQ1 COOH terminus-CaM complex.](link to image)

We ask if metal binding is necessary for assembly formation by using thermal denaturation as measured with CD. The addition of metal did not change the melting temperature, indicating no stabilizing effect. We then turned to mutagenesis and functional assays to evaluate the significance of metal binding. When we mutated the histidine 620 to alanine, no appreciable effect was seen on current recorded from transfected cells. When the histidine was changed to serine, although not significant, some inhibition (~50%) of the current was detected (Table 3 and Fig. 5, C and D).

**DISCUSSION**

We outline our findings with a working model for the KCNQ1 COOH terminus structure and its functional correlates. A structural model is drawn in Fig. 6. The COOH terminus structure may be best summarized by dividing it into two halves, proximal and distal to the membrane. The proximal half binds CaM. The configuration of CaM association that best agrees with our data has one CaM binding by sandwiching helices A and B of a single subunit. Accordingly, CaM does not play
a direct role in oligomerization but apparently participates in forming a module, pivotal for regulating channel gating (8, 11). Nevertheless, apoCaM does play a critical role in correct COOH-terminal folding, since its absence results in no functional channels and no soluble COOH terminus (8, 11), whereas Ca$^{2+}$-CaM is not required, as seen from experiments using CaM$_{1234}$ (Fig. 1F) (11). Also, calcium does not induce dimerization of the proximal COOH terminus as observed in our sedimentation velocity experiments with $\Delta$ helices C-D-CaM, unlike the SK channel structure (39). Helices A and B are connected by a long stretch of polypeptide, dubbed above the intervening loop, that has little sequence conservation, regions of low complexity, and little predicted secondary structure. This intervening loop is not required for formation of the tetrameric COOH terminus-CaM complex (Fig. 1C).

The distal COOH-terminal half contains two tandemly arranged coiled-coils. This half directs oligomerization due to two structural components, namely the helix C module, the first coiled-coil that is dimeric, and subsequently the helix D module, the second coiled-coil that is tetrameric. These two components are linked by a proteolytically susceptible linker, strongly suggesting that this 22-residue span is mobile and not well structured. Two independent observations support that conclusion: first, adventitious proteolysis in this region enabled crystallization of the helix D complex from the COOH terminus $\Delta$ loop-CaM complex, and second, residues 574–586, although present in the protein, are not discernible in the electron density map.

The two coiled-coils are intrinsically different, since the helix C module fundamental unit appears to be dimeric, probably parallel due to structural considerations. This unit undergoes concentration-dependent self-association into a dimer of dimers. This concentration-dependent self-association may explain findings by Ghosh et al. (11) that a fraction of COOH terminus (truncated at residue 555)-CaM complex tetramerizes, as found by chemical cross-linking performed at 1 mg/ml protein concentration. Indeed, this truncation leaves intact a significant portion of the helix C module. Similarly, Schwake et al. (14) found that mutants disrupting the helix D module in KCNQ2 or -3 still afforded functional current, albeit significantly diminished, suggesting that the helix C module is sufficient for tetramerization. In contrast, the helix D module fundamental unit is a stable tetramer, even at lower concentrations. Despite the difference in quaternary structure, their individual thermal stabilities are about the same. Thermodynamically, both of the specific modules are less stable than engineered coiled-coils, not unlike some natural coiled-coils that have quite low melting points (40). These $T_m$ results may have implications for the dynamics of KCNQ1 biosynthesis and assembly. However, when they fold/unfold in their native tandem arrangement, there is increased thermal stabilization caused either by structural or kinetic reasons. Thus, the distal COOH terminus assembles by a concerted action of its particular modules.

Inspection of sequence conservation (supplemental Fig. 2) in the distal COOH terminus reveals that the helix C module is strongly conserved within the Kv7 subfamily, at both buried and surface-accessible residues. The buried residue conservation may emphasize a general role in promoting oligomerization, whereas the conservation of surface residues may hint at its role as a surface for associating proteins involved in trafficking and regulation. The helix D module differs in its conservation characteristics, exhibiting markedly lower overall levels within the Kv7 subfamily with the amino-terminal third more conserved, including solvent-exposed residues (non-$a$ and $d$ positions). We propose three nonexclusive possibilities for the significance of this conservation. First, the lesser Kv7 subfamily conservation is in line with a role that specifies the type of oligomerization (i.e. homotypic and/or heterotypic). Second, coiled-coils undergo folding, which may be initiated by so-called “trigger” sequences. Examples of this phenomenon have been isolated in certain coiled-coils, and a consensus sequence has been formulated (41). Other studies have shown that trigger sequences are not found in all coiled-coils (42). The KCNQ1 sequence conforms to this trigger sequence except for the first two positions (Leu $\rightarrow$ Ile$^{588}$ and Glu $\rightarrow$ Gly$^{589}$). Thus, the conserved NH$_2$-terminal end may comprise a trigger sequence for helix D assembly. When we perturbed this sequence region (residues 588–594), removing the entire helix D sequence in our attempt to substitute helix D with GCN4-LI, as described above, the COOH terminus-CaM complex aggregated in a nonspecific manner (data not shown). However, when we preserved this region (residues 588–594), we were able to construct a viable chimeric COOH terminus-GCN4-LI (Fig. 3, B–D), emphasizing the sequence’s importance for formation of the helix D complex. Third, this region generates a surface for essential interacting proteins, conservation being a hallmark of such interfaces (36). Finally, we note the relative sequence conservation of the linker between the helix C and D modules, discerned also through the prism of LQT syndrome mutations in this region (supplemental Table 1). This feature may be due to specific requirements for folding, as just described for the KCNQ1-GCN4 chimeras. Alternatively, these conserved sequences are required for other proteins to bind.

Two salient structural features may explain the observation that human KCNQ1 homotetramerizes exclusively (i.e. it is partner-specific), whereas other members can heterotetramerize or homotetramerize (13, 16). First, the KCNQ1 helix D should be about two helical turns longer than the other family members. This additional heptad repeat will prefer assembly with a partner of equal length and sequence in order to realize most fully its folding potential. Indeed, the recent report of the KCNQ4 helix D complex structure supports this notion (43). Second, human KCNQ1 contains a metal coordination site capable of constructing a viable chimeric COOH terminus-GCN4-LI (Fig. 3, B–D), emphasizing the sequence’s importance for formation of the helix D complex. This, third, region generates a surface for essential interacting proteins, conservation being a hallmark of such interfaces (36). Finally, we note the relative sequence conservation of the linker between the helix C and D modules, discerned also through the prism of LQT syndrome mutations in this region (supplemental Table 1). This feature may be due to specific requirements for folding, as just described for the KCNQ1-GCN4 chimeras. Alternatively, these conserved sequences are required for other proteins to bind.

Regarding this latter feature, our findings lead us to posit that since the histidines are positioned at the $a$ heptad repeat position, pointing into the interior of the coiled-coil, and alanine is found frequently at the $a$ position, the assembly or folding of tetrameric helix D is maintained. However, when serine (a hydrophilic residue infrequently found at that position and incapable of coordinating a metal ion due to its short length) is introduced, assembly is affected. Therefore, the metal coordination found in helix D of human KCNQ1 may have physiological relevance by providing an additional mechanism for assembly partner specificity. A possible structural precedent for this type of mechanism is found in Kv channels that contain the
tetramerization domain, T1. In that case, certain subtypes encode a Zn$^{2+}$ binding motif that determines partner subtype specificity (44). Further experimentation will be required to test whether metal coordination may indeed be a KCNQ1 specificity determinant.

Both of the helix C and D modules in KCNQ1 are nonredundant at the functional level, although structurally there may be a degree of overlap. Our cellular studies show that the function of helix C is complex, since a construct like Δ helix C, despite its apparent biochemical viability, failed to reach the plasma membrane, implying that helix C plays a role in channel trafficking (Fig. 2, G–I). Hence, we assign it as a module for proper assembly and trafficking to the plasma membrane. Our experiments with helix D clearly demonstrate that it encodes a complex function also. Thus, the tetramerization functionality could be replaced in part on the biochemical level with GCN4-LI but is insufficient for functional expression, suggesting a role in trafficking, implicitly by association of proteins to this region (15).

Our analysis of helix D point mutants (e.g. A590W) that demonstrate an ability to assemble to levels approaching that of WT nonetheless shows impaired functional expression, attributable in part to faulty trafficking. We speculate that such mutations may perturb binding surfaces required for association of trafficking or regulatory proteins, particularly in the amino-termini of the helix D module. Such an interpretation may also explain the mechanism for LQT mutations (e.g. G589D) in this region.

What does the COOH terminus do? T1, the intracellular domain amino-terminal to the membrane domain in several K$^+$ channel subfamilies, is not essential for Kv1 and Shaker channel formation. Jan and co-workers (30) suggested that it enhances channel assembly by increasing the local concentration of subunits in a multistep process. Inherent to this hypothesis is the premise that another oligomerization element exists that resides in the membrane domain of the protein. T1 then improves quaternary structure formation efficiency. In contrast, KCNQ1 channel formation cannot be coaxed out of truncations lacking its COOH terminus. Once again, the membrane domain must contribute considerably to channel assembly but is not sufficient. The COOH terminus must provide the necessary energy deficit for assembly. We propose that the helix C module may contribute adequate free energy for this deficit at some level, thereby creating a dimer of dimers, although this may vary from Kv7 subfamily member to member (14).

Although this role might explain its outstanding conservation, this functionality is not sufficient for full maturation of the channel. To fully specify the correct partners for the tetramer and also stabilize oligomerization, the tetrameric helix D module is required. A similar functional architecture has been shown for the related EAG family of potassium channels (45). Previous and present results intimate that it provides a protein–protein interaction platform for trafficking and regulation, in some measure analogous to T1. Consequently, the whole COOH terminus constitutes an elongated, nonglobular, multitiered scaffold for associating proteins like CaM and others central to the signaling process (46).

Finally, the layering, physically, of oligomerization predicts a staged channel assembly, whose temporal sequence at this point remains unknown. The use of coiled-coils in channel assembly perforce couples folding (i.e. tertiary structure formation) with oligomerization (i.e. quaternary structure formation), parallel to conclusions drawn by Deutsch and Robinson (47) regarding the biosynthesis of K$^+$ channels that employ T1.

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REFERENCES

1. Jentsch, T. J. (2000) Nat. Rev. Neurosci. 1, 21–30
2. Robbins, J. (2001) Pharmacol. Ther. 90, 1–19
3. Jentsch, T. I., Hubner, C. A., and Fuhrmann, J. C. (2004) Nat. Cell Biol. 6, 1039–1047
4. Long, S. B., Campbell, E. B., and Mackinnon, R. (2005) Science 309, 897–903
5. Abbott, G. W., and Goldstein, S. A. (2001) Mol. Interv. 1, 95–107
6. Yus-Najera, E., Santana-Castro, I., and Villarroel, A. (2002) J. Biol. Chem. 277, 28545–28553
7. Wen, H., and Levitan, I. B. (2002) J. Neurosci. 22, 7991–8001
8. Shamgar, L., Ma, L., Schmitt, N., Hattin, Y., Peretz, A., Wiener, R., Hirsch, I., Pongs, O., and Attali, B. (2006) Circ. Res. 98, 1055–1063
9. Gamper, N., Stockand, J. D., and Shapiro, M. S. (2003) J. Neurosci. 23, 84–95
10. Gamper, N., Li, Y., and Shapiro, M. S. (2005) Mol. Biol. Cell 16, 3538–3551
11. Ghosh, S., Nunziato, D. A., and Pitt, G. S. (2006) Circ. Res. 98, 1048–1054
12. Schmitt, N., Schwarz, M., Peretz, A., Abitbol, I., Attali, B., and Pongs, O. (2000) EMBO J. 19, 332–340
13. Schwarz, M., Jentsch, T. I., and Friedrich, T. (2003) EMBO Rep. 4, 76–81
14. Schwarz, M., Athanasiadu, D., Beimgraben, C., Blanz, J., Beck, C., Jentsch, T. I., Saftig, P., and Friedrich, T. (2006) J. Neurosci. 26, 3757–3766
15. Kanki, H., Kupershmidt, S., Yang, T., Wells, S., and Roden, D. M. (2004) J. Biol. Chem. 279, 33976–33983
16. Maljevic, S., Lerche, C., Seehom, G., Alekov, A. K., Busch, A. E., and Lerche, H. (2003) J. Physiol. 548, 353–360
17. Schuck, P. (2000) Biophys. J. 78, 1606–1619
18. van Holde, K. E. (1986) in Physical Biochemistry, 2nd Ed., pp. 110–136, Prentice-Hall, Englewood Cliffs, NJ
19. Laue, T. M., Shah, B. D., Ridgeway, T. M., and Pelletier, S. L. (1992) in Analytical Ultracentrifugation in Biochemistry and Polymer Science (Harding, S. E., Rowe, A. J., and Horton, J. C., eds) pp. 90–125, Royal Society of Chemistry, Cambridge, UK
20. Cole, J. L. (2004) Methods Enzymol. 384, 212–232
21. Rivas, G., Stafford, W., and Minton, A. P. (1999) Methods 19, 194–212
22. Minton, A. P. (1994) in Modern Analytical Ultracentrifugation (Schuster, T. M., and Laue, T. M., eds) pp. 81–93, Birkhauser, Boston, MA
23. Otwinowski, Z., and Minor, W. (1997) Methods Enzymol. 276, 307–326
24. Brunger, A. T., Adams, P. D., Clore, G. M., DeLano, W. L., Gros, P., Grosse-Kunstleve, R. W., Jiang, J. S., Kuszewski, J., Nilges, M., Pannu, N. S., Read, R. J., Rice, L. M., Simonson, T., and Warren, G. L. (1998) Acta Crystallogr. Sect. D Biol. Crystallogr. 54, 905–921
25. de La Fortelle, E., and Bricogne, G. (1997) Methods Enzymol. 276, 472–494
26. Abrahams, J. P., and Leslie, A. G. W. (1996) Acta Crystallogr. Sect. D Biol. Crystallogr. 52, 30–42
27. Morris, R. J., Perrakis, A., and Lamzin, V. S. (2003) Methods Enzymol. 374, 229–244
28. Murshudov, G. N., Vagin, A. A., and Dodson, E. J. (1997) Acta Crystallogr. Sect. D Biol. Crystallogr. 53, 240–255
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29. Waxman, E., Laws, W. R., Laue, T. M., Nemerson, Y., and Ross, J. B. (1993) Biochemistry 32, 3005–3012
30. Zerangue, N., Jan, Y. N., and Jan, L. Y. (2000) Proc. Natl. Acad. Sci. U. S. A. 97, 3591–3595
31. Harbury, P. B., Zhang, T., Kim, P. S., and Alber, T. (1993) Science 262, 1401–1407
32. Strelkov, S. V., and Burkhard, P. (2002) J. Struct. Biol. 137, 54–64
33. Walshaw, J., and Woolfson, D. N. (2001) J. Mol. Biol. 307, 1427–1450
34. Kleywegt, G. J., and Jones, T. A. (1994) Acta Crystallogr. Sect. D Biol. Crystallogr. 50, 178–185
35. Fleming, P. J., and Richards, F. M. (2000) J. Mol. Biol. 299, 487–498
36. Marx, S. O., Kurokawa, J., Reiken, S., Motoike, H., D’Armiento, J., Marks, A. R., and Kass, R. S. (2002) Science 295, 496–499
37. Woolfson, D. N. (2005) Adv. Protein Chem. 70, 79–112
38. Ducros, V., Brzozowski, A. M., Wilson, K. S., Brown, S. H., Ostergaard, P., Schneider, P., Yaver, D. S., Pedersen, A. H., and Davies, G. J. (1998) Nat. Struct. Biol. 5, 310–316
39. Schumacher, M. A., Rivard, A. F., Bachinger, H. P., and Adelman, J. P. (2001) Nature 410, 1120–1124
40. Mason, J. M., Schmitz, M. A., Muller, K. M., and Arndt, K. M. (2006) Proc. Natl. Acad. Sci. U. S. A. 103, 8989–8994
41. Kammerer, R. A., Schulthess, T., Landwehr, R., Lustig, A., Engel, J., Aebi, U., and Steinmetz, M. O. (1998) Proc. Natl. Acad. Sci. U. S. A. 95, 13419–13424
42. Lee, D. L., Lavigne, P., and Hodges, R. S. (2001) J. Mol. Biol. 306, 539–553
43. Howard, R. J., Clark, K. A., Holton, J. M., and Minor, D. L., Jr. (2007) Neuron 53, 663–675
44. Bixby, K. A., Nanao, M. H., Shen, N. V., Kreusch, A., Bellamy, H., Pfaffinger, P. J., and Choe, S. (1999) Nat. Struct. Biol. 6, 38–43
45. Jenke, M., Sanchez, A., Monje, F., Stuhmer, W., Weseloh, R. M., and Pardo, L. A. (2003) EMBO J. 22, 395–403
46. Kass, R. S., Kurokawa, J., Marx, S. O., and Marks, A. R. (2003) Trends Cardiovasc. Med. 13, 52–56
47. Robinson, J. M., and Deutsch, C. (2005) Neuron 45, 223–232