Unique gating properties of C. elegans ClC anion channel splice variants are determined by altered CBS domain conformation and the R-helix linker

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All eukaryotic and some prokaryotic ClC anion transport proteins have extensive cytoplasmic C-termini containing two cystathionine-β-synthase (CBS) domains. CBS domain secondary structure is highly conserved and consists of two α-helices and three β-strands arranged as β1-α1-β2-β3-α2. CIC CBS domain mutations cause muscle and bone disease and alter ClC gating. However, the precise functional roles of CBS domains and the structural bases by which they regulate ClC function are poorly understood. CLH-3a and CLH-3b are C. elegans ClC anion channel splice variants with strikingly different biophysical properties. Splice variation occurs at cytoplasmic N- and C-termini and includes several amino acids that form α2 of the second CBS domain (CBS2). We demonstrate that interchanging α2 between CLH-3a and CLH-3b interchanges their gating properties. The “R-helix” of ClC proteins forms part of the ion-conducting pore and selectivity filter and is connected to the cytoplasmic C-terminus via a short stretch of cytoplasmic amino acids termed the “R-helix linker”. C-terminus conformation changes could cause R-helix structural rearrangements via this linker. X-ray structures of three ClC protein cytoplasmic C-termini suggest that α2 of CBS2 and the R-helix linker could be closely apposed and may therefore interact. We found that mutating apposing amino acids in α2 and the R-helix linker of CLH-3b was sufficient to give rise to CLH-3a-like gating. We postulate that the R-helix linker interacts with CBS2 α2, and that this putative interaction provides a pathway by which cytoplasmic C-terminus conformational changes induce conformational changes in membrane domains that in turn modulate ClC function.

Introduction

Members of the CIC superfamily of anion transport proteins have been identified in plants, yeast, eubacteria, archaeabacteria and various invertebrate and vertebrate animals. CIC genes encode anion channels and Cl-/H+ exchangers and are expressed in plasma and intracellular organelle membranes where they perform essential physiological functions including transepithelial Cl- transport, systemic ion homeostasis, regulation of membrane potential and regulation of cytoplasmic and intra-organelle Cl- and H+ levels. Mutations in three of the nine human CIC encoding genes give rise to muscle, bone and kidney diseases.

Extensive electrophysiological and molecular biological studies have shown that CIC proteins are homodimers and that each monomer forms an independently gated protopore that functions as an anion channel or a Cl-/H+exchanger. X-ray crystallography studies of bacterial CICs have confirmed this model and demonstrate that each monomer comprises 18 α-helical domains (designated “A-R”), 17 of which are membrane embedded. The overall three-dimensional membrane structure of the bacterial CIC proteins is likely conserved throughout the CIC superfamily.

All eukaryotic and some bacterial CIC proteins have extensive cytoplasmic C-terminal domains following the R-helix. These intracellular C-termini contain two cystathionine-β-synthase (CBS) domains. The CBS domain is a 50–60 amino acid motif found in numerous diverse proteins. Mutations in CBS domains give rise to several diseases and alter ClC gating, indicating that they play a critical role in protein structure/function. Structural and biochemical studies have demonstrated that the cytoplasmic C-termini of eukaryotic CICs interact to form dimers.

Six CIC genes, termed clh (Cl channel homolog)-1-6, are present in the C. elegans genome. C. elegans CLH proteins are representative of the three mammalian CIC subfamilies. clh-3 encodes two splice variants, CLH-3a and CLH-3b, that exhibit striking differences in sensitivity to depolarizing voltages, activating voltages and extracellular pH and Cl-. The structural bases by which the cytoplasmic C-terminus controls CIC gating and activity are poorly understood. We

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Results

Large deletion mutations in CLH-3b CBS domains give rise to CLH-3a-like gating. Figure 1 shows the sequence of the predicted intracellular domains of CLH-3a and CLH-3b. The membrane-associated domains of the two channels are identical and have been omitted for clarity. Splice variations in CLH-3a and CLH-3b are highlighted in yellow and green, respectively. Location of CBS domains were predicted using the domain recognition programs PFAM, InterProScan and MyHits and are outlined with boxes.
Figure 2. CLH-3b CBS domain deletion mutations give rise to CLH-3a-like gating. Current-to-voltage relationships, activation voltages and activation by bath acidification are shown in (A–C), respectively. **p < 0.001 and *p < 0.01 compared to wild-type CLH-3b. Effects of depolarized holding potentials are shown in (D–F). (D) Representative current traces showing hyperpolarization induced current activation in cells held at depolarized holding potentials. Cells were held at voltages of -20 to 60 mV for 3 sec before currents were activated by stepping to -120 mV. (E) Current activation by depolarized holding potentials. Peak current amplitudes are normalized to that measured following a holding potential of -20 mV (I_{-20 mV holding potential}). (F) Effect of holding potential on current inactivation. Mean pseudo-steady-state current (I_{steady-state}) was measured over the last 20 msec of the -120 mV test pulse and normalized to peak current (I_{peak}) amplitude. Values in (A–C, E and F) are means ± S.E. (n = 5–12). CLH-3bΔ822-838 is a deletion mutant lacking the last 17 amino acids of CBS2.
CLH-3a and CLH-3b show strong inward rectification and activation at hyperpolarized voltages (Fig. 2A). However, CLH-3a activation requires much stronger hyperpolarization. Mean activation voltages were -54 mV for CLH-3a and -27 mV for CLH-3b (Fig. 2B). Deletion of the last seventeen amino acids of CBS2 in CLH-3b induced a hyperpolarizing shift in activation voltage (Fig. 2A and B). The mean activation voltage of CLH-3b∆822-838 was ∼-52 mV and was not significantly (p > 0.6) different from that of CLH-3a.

CIC channels are activated by extracellular acidification. Reduction of bath pH to 5.9 activates CLH-3a and CLH-3b ∼3.2- and ∼1.6-fold, respectively. Deletion of amino acids 822–838 significantly (p < 0.001) increased the pH sensitivity of CLH-3b resulting in a ∼4.4-fold activation by reducing bath pH to 5.9 (Fig. 2C).

CLH-3a shows unique sensitivity to depolarized holding voltages termed pre-depolarization induced potentiation. As shown in Figure 2D and E, increasing the degree of holding potential depolarization increases the extent of or “potentiates” hyperpolarization-induced current activation. Currents potentiated by pre-depolarization also undergo slow, partial inactivation (Fig. 2D and F). Wild-type CLH-3b is insensitive to depolarized holding voltages (Fig. 2D–F). However, the deletion mutation in CBS2 induced sensitivity to pre-depolarization that resembled that of CLH-3a (Fig. 2D–F). Taken together, data in Figure 2 and our previous studies demonstrate that disruption of either CBS1 or CBS2 structure in CLH-3b gives rise to channels with CLH-3a-like biophysical properties.

In contrast to CLH-3b, deletion of the last 11 amino acids of CBS1 or the last 17 amino acids of CBS2 in CLH-3a had no effect on voltage and pH sensitivity (Fig. 3A and B) or the response to depolarized holding potentials (Fig. 3C and D). These data demonstrate that the unique gating properties of CLH-3a are insensitive to disruption of CBS domain conformation.

Splice variation of CBS2 determines the gating characteristics of CLH-3a and CLH-3b. The primary sequence of CBS domains in diverse proteins is variable, but the motif has a highly conserved secondary structure consisting of an N-terminal β-strand (β1) followed by an α-helix (α1), two β-strands (β2 and β3) and an α-helix (α2). As shown recently by Dutzler and coworkers, CBS1 and CBS2 domains play an important role in determining
the overall homodimeric structure of the cytoplasmic regions of CIC proteins. Large deletions in either CBS1 or CBS2 are expected to dramatically disrupt C-terminus conformation and function. Our deletion mutation studies (Figs. 2 and 3) as well as previous work\textsuperscript{18,19} demonstrate that overall CBS domain architecture plays a critical role in determining CLH-3b but not CLH-3a functional properties.

As shown in Figure 1, CBS2 domains of CLH-3a and CLH-3b predicted using the domain recognition programs PFAM,\textsuperscript{30} InterProScan\textsuperscript{31} and MyHits\textsuperscript{32} are alternatively spliced at the last six amino acids. This region forms part of α2. The sequences of the six splice variant amino acids in CLH-3a and CLH-3b are VCFLIS and LRLAIY, respectively.

We postulated that splice variation of α2 may be a critical determinant of channel biophysical properties. To test this hypothesis, we interchanged the six alternatively spliced amino acids between the channels. CLH-3b + VCFLIS exhibited a hyperpolarized activation voltage (Fig. 4A), increased pH sensitivity (Fig. 4B) and sensitivity to depolarized holding potentials (Fig. 4C and D). Replacement of the last six amino acids in CBS2 of CLH-3a with the analogous amino acids in CLH-3b (CLH-3a + LRLAIIE) had little effect on activation voltage or pH sensitivity (data not shown).

The response of CLH-3a + LRLAIIE to pre-depolarization was variable between cells. Of the 15 cells that were patch-clamped from two separate transfections, we found that two cells showed a strong response to pre-depolarization resembling that of wild-type CLH-3a, seven cells showed no response and six cells exhibited a weak response (data not shown). This variability suggested to us that additional spliced amino acids may contribute to channel gating properties.

To examine CBS2 structure in greater detail than that predicted by domain recognition programs, we used Rosetta software\textsuperscript{33,34} to develop 20,000 candidate structures of amino acids 803–811 (i.e., VCFLISRKK) for CLH-3a and 833–841 (i.e., LRLAIYELQ) for CLH-3b. Figure 5 is a histogram showing the number of amino acids in this region that adopt a helical conformation and the number of Rosetta models that predict

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**Figure 4.** Interchanging alternatively spliced amino acids in CBS2 interchanges channel functional properties. (A) Activation voltage. **p < 0.0001 compared to wild-type CLH-3b (B) Activation by bath acidification. *p < 0.05 compared to wild-type CLH-3b (C) Current activation by depolarized holding potentials. (D) Effect of holding potential on current inactivation. Data in (C and D) were acquired as described in Figure 2 legend. Values are means ± S.E. (n = 5–11). CLH-3b + VCFLIS and CLH-3a + LRLAIYELQ are chimeras in which alternatively spliced amino acids in CBS2 have been interchanged between the two channels.
To determine whether the last three amino acids of CLH-3a (i.e., RKK) independently regulate channel properties, we exchanged them with the analogous amino acids of CLH-3b (i.e., YLQ). Replacing RKK with YLQ (i.e., CLH-3a + YLQ) had no significant effect on pH sensitivity or activation voltage (p > 0.7) or sensitivity to depolarized voltages (data not shown). Similarly, CLH-3b + RKK showed reduced pH sensitivity similar to that of wild-type CLH-3b and was insensitive to pre-depolarization (data not shown). However, the activation voltage for CLH-3b + RKK was significantly (p < 0.0001) hyperpolarized to a mean ± S.E. value of -51 ± 1 mV (n = 13), which is similar to that of wild-type CLH-3a (see Fig. 2B).

The R-helix linker plays a critical role in determining CLH-3a and CLH-3b gating properties. Data in Figure 4 demonstrate that splice variation of α2 of CBS2 plays a major role in determining the gating characteristics of CLH-3a and CLH-3b. Differences in the primary and secondary structure of α2 could alter local interactions within the channel that give rise to unique gating properties. To explore this possibility, we generated homology models of the cytoplasmic C-termini of CLH-3a and CLH-3b based on the 1.6 angstrom crystal structure of ClC-Ka (Fig. 6). The model of the cytoplasmic C-terminal domain was manually docked to the membrane-associated region of CLC-ec1 (Protein Data Bank accession CODE_1OTS) to provide context for the model (Fig. 6B). The cytoplasmic C-termini of both channels were modeled as homodimers as has been shown experimentally for CIC-0, CIC-5, CIC-Ka and CIC-Kb. Each pair of CBS domains within a monomer interact to form a dimer and the monomer-monomer interface is located between the two CBS2 domains (Fig. 6B).

A stretch of 20 cytoplasmic amino acids termed the R-helix linker connects the C-terminus of the R-helix of CLH-3a and CLH-3b to the N-terminus of CBS1 (Fig. 6C). The R-helix forms part of the channel pore and selectivity filter. The conformation of the R-helix and thus channel properties may be modulated by cytoplasmic domains. Crystal structures of CIC-0, CIC-Ka and CIC-5 cytoplasmic C-termini indicate that part of the cytoplasmic R-helix linker lies close to α2 of CBS2 (Fig. 6B). This close apposition suggests that the linker could interact with CBS2 and that this putative interaction may in turn play a role in mediating the differences in CLH-3a and CLH-3b gating induced by α2 splice variation.

Examination of the homology models indicated that F559 in the R-helix linker of CLH-3b lies very close to A836 in α2 of CBS2 (Fig. 6D). The orientation of the homologous phenylalanine in CLH-3a, F630, could be altered due to the presence of a bulkier leucine residue, L806, at the corresponding position.
As we have shown previously, CLH-3a and CLH-3b mutants in which the extracellular glutamate residue that forms the fast gate is mutated to cysteine show 7–8-fold differences in reactivity to the sulfhydryl modifying reagent MTSET. Deletion mutations in the CLH-3b C-terminus give rise to MTSET reactivity very similar to that of CLH-3a suggesting that cytoplasmic domains modulate the conformation of channel extracellular regions. To determine whether the F559D mutation alters the conformation of CLH-3b extracellular domains, we mutated the fast gate glutamate residue to cysteine (i.e., E167C). Consistent with our previous observations, this mutation gave rise to constitutively active channels that are no longer sensitive to voltage and extracellular protons (data not shown).

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Exposure of the E167C mutant to 1 mM MTSET inhibited whole cell current amplitude as we have shown previously.\textsuperscript{18} The mean ± S.E. time constant for inhibition was 283 ± 34 sec (n = 6). Mutation of F559 to aspartate reduced the inhibition time constant nearly 7-fold to a mean ± S.E. value of 41 ± 6.6 sec (n = 5). The striking difference in rates of MTSET inhibition for E167C and the E167C/F559D double mutant is remarkably similar to the difference observed between CLH-3a and CLH-3b.\textsuperscript{18} These data together with the results shown in Figure 7 demonstrate that mutation of a single amino acid in the R-helix linker, F559, to aspartate converts CLH-3b gating properties and extracellular cysteine reactivity to those resembling CLH-3a.

As shown in Figure 6D, the homology model predicts that F559 lies close to A836 in α2 suggesting that the two amino acids could interact. To test this possibility, we mutated A836 to aspartate. The A836D mutant gave rise to a small hyperpolarizing shift in activation voltage and potentiation behavior that was considerably less than that observed with wild-type CLH-3a (data not shown). We therefore mutated A836 to a larger, positively charged lysine. The A836K mutant exhibited voltage and pH sensitivity nearly identical to those of CLH-3a (Fig. 8A–D).

We also mutated amino acids immediately adjacent to A836. Mutation of L835 to lysine (L835K) had no effect on the voltage or pH sensitivity of CLH-3b while mutating I837 to lysine (I837K) gave rise to CLH-3a-like behavior (Fig. 8A–D). Both amino acids are predicted to face away from the R-helix linker (Fig. 6D). Crystal structures of various tandem CBS domain containing proteins indicate that α1 and α2 lie close together and may physically interact.\textsuperscript{13,15,35,36} We examined the CLH-3b homology model to assess the relationship of α2 amino acids to α1 in CBS2. As shown in Figure 6D, L835 faces away from α1 while I837 faces towards it and could interact with amino acids.

**Figure 8.** Mutation of amino acids in α2 of CBS2 predicted to interact with F559 in the R-helix linker or α1 amino acids give rise to CLH-3a-like gating. (A) Activation voltage. (B) Activation by bath acidification. (C) Current activation by depolarized holding potentials. (D) Effect of holding potential on current inactivation. Data in (C and D) were acquired as described in Figure 2 legend. Values are means ± S.E. (n = 5–8). *p < 0.01 and **p < 0.0001 compared to wild-type CLH-3b. L833, L835, A836 and I837 are amino acids located in α2 of CBS2 of CLH-3b.
on this helix. This suggests that the L837K mutation may disrupt putative α1-α2 interactions. To test this possibility, we mutated L833, which also lies close to α1 (Fig. 6D), to glutamate (L833E). The L833E mutant exhibited CLH-3a-like behavior (Fig. 8A–D). Data shown in Figures 2–4, 7 and 8, and our previous findings indicate that mutations that disrupt CBS domain structure and/or putative interactions of CBS2 with the R-helix linker give rise to gating characteristics resembling those of CLH-3a.

Discussion

The CBS domain is a highly conserved motif found in diverse proteins including channels, transporters, kinases and metabolic enzymes of archaebacteria, euabacteria and eukaryotes. Mutations in CBS domains cause diverse inherited diseases such as cardiomyopathy, homocystinuria, retinitis pigmentosa, osteopetrosis and myotonia.

The precise function of CBS domains is poorly understood. In several diverse proteins, CBS domains have been shown to bind adenosyl compounds such as ATP, ADP and AMP that in turn regulate protein function. CBS domains in the MgE Mg<sup>2+</sup> transporter likely bind Mg<sup>2+</sup> and function as part of an intracellular Mg<sup>2+</sup> sensor that regulates transporter activity. Recent studies on the bacterial osmoregulatory organic osmolyte transporter OprA have shown that CBS motifs regulate the transporter on/off state by binding in an ionic strength dependent manner to charged membrane domains.

CBS motifs are typically present in proteins as two or four copies. Pairs of CBS motifs associate to form a dimeric structure known as a Bateman module or domain. The monomer-monomer interface between two CBS motifs occurs at β-sheets formed by β-strands 2 and 3. Meyer and Dutzler postulated a head-to-tail interaction between Bateman domains of CIC-0 cytoplasmic C-termini (i.e., CBS1 of one Bateman module interacts with CBS2 of the second domain). However, X-ray crystallography studies on dimers of CIC-5 and CIC-Ka cytoplasmic C-termini revealed a novel interface formed by the CBS2 motifs of each C-terminal monomer. This “tail-to-tail” arrangement of Bateman domains is shown in the CLH-3b homology model in Figure 6B.

Large deletion mutations in either CBS1 or CBS2 are expected to dramatically disrupt CBS domain tertiary structure and this in turn likely disrupts oligomerization. Our results demonstrate that disruption of CLH-3b CBS domain structure (Fig. 2) gives rise to channels with CLH-3a-like biophysical properties while disruption of CLH-3a CBS structure has little effect on channel gating (Fig. 3). These findings indicate that the overall CBS domain architecture of CLH-3b plays a critical role in determining the functional properties of the channel and that (2) that CLH-3a gating appears to be largely insensitive to changes in CBS domain conformation.

The C-terminal end of CBS2 is alternatively spliced in CLH-3a and CLH-3b (Fig. 1). This spliced region forms the second α-helix domain, α2. Splice variation alters not only the primary sequence, but also is predicted to alter α2 secondary structure. Rosetta modeling suggested that α2 in CLH-3b is longer than that of CLH-3a (Fig. 5). Consistent with this, we found that insertion of the last six amino acids of the CLH-3a CBS2 domain (i.e., VCFLIS) into CLH-3b CBS2 gives rise to CLH-3a-like voltage- and pH-sensitivity (Fig. 4). Inserting the analogous six amino acids of CLH-3b (i.e., LRLAEY) into CLH-3a had little effect on activation voltage or pH sensitivity and had an inconsistent effect on sensitivity to depolarizing holding potentials. However, when the nine alternatively spliced amino acids comprising the longer α2 of CLH-3b (i.e., LRLAEYLYQ) were inserted into CLH-3a, the mutant channels were fully insensitive to depolarized voltages (Fig. 4C and D).

Given that large scale disruption of CBS structure by deletion mutations has no effect on CLH-3a gating, we suggest that splice variation of α2 in CBS2 of this channel disrupts critical interactions between CBS domains and other regions within the C-terminus. The absence of these interactions gives rise to the gating characteristics of CLH-3a. In contrast, both the primary sequence and predicted altered secondary structure of CLH-3b α2 mediate important functional interactions. Disruption of these interactions by mutation of either CBS1 or CBS2 gives rise to channels with CLH-3a-like properties.

It is important to note that the CLH-3a + LRLAEYLYQ chimeric channel did not fully recapitulate CLH-3b gating. Activation voltage and pH sensitivity of this chimera resembled those of wild-type CLH-3a (Fig. 4A and B). These results indicate that other cytoplasmic domains unique to CLH-3b play a role in determining these gating characteristics. Our previous studies have shown that deletion of the 101 amino acid linker domain between CBS1 and CBS2 in CLH-3b (Fig. 1) gives rise to channels with a strongly hyperpolarized activation voltage. In addition, deletion of the 160 amino acids immediately C-terminal to LRLAEYLYQ (Fig. 1) increases pH sensitivity and hyperpolarizes channel activation voltage (Dave and Strange, unpublished observations).

The molecular mechanisms by which cytoplasmic structures and intracellular signaling events regulate CIC channel/transporter function are poorly understood. Dutzler and colleagues have suggested that the membrane-associated R-helix may play an important role in this regulation. By virtue of its direct connection to the large cytoplasmic C-terminus, the R-helix may provide a pathway by which conformational changes in intracellular domains induce rearrangements of the pore that in turn alter channel/transporter activity.

A short, cytoplasmic linker connects the R-helix to CBS1. Interestingly, despite significant differences in primary sequence, part of the R-helix linkers of CIC-0, CIC-5 and CIC-Ka have similar and well ordered crystal structures. The conservation of this structure suggests that the linker likely plays an important role in channel function.

Crystal structures of CIC-0, CIC-5 and CIC-Ka indicate that α2 of CBS2 lies close to the cytoplasmic R-helix linker. Our homology models of CLH-3a and CLH-3b suggested putative interactions between α2 amino acids and an R-helix linker.
phenylalanine residue that could be altered by splice variation. Consistent with this, we found that mutation of CLH-3b F559 to aspartate or mutation of a closely apposed alanine residue in α2 gave rise to channels with gating properties and extracellular cysteine reactivity that fully recapitulated those of CLH-3a (Figs. 7 and 8, and Results). In contrast, mutation of the analogous R-helix linker phenylalanine residue in CLH-3a had no effect on channel activity.

Our data are consistent with the hypothesis that amino acids in α2 of CBS2 and the R-helix linker interact, and suggest that this putative interaction could be important for determining the conformation of the R-helix and other associated regions that control channel gating. We postulate that splice variation of α2 in and lead to changes in channel function. This hypothesis is sup-

modulate the conformation of the R-helix and associated structures by which C-terminus conformational changes could regulate channel properties. We propose a mechanism in which the functional roles of ClC CBS domains and the R-helix linker in and structural studies are needed to directly test this hypothesis. It will be particularly important to determine the role played by the R-helix linker and CBS domain conformation in mediating the regulation of CIC channels and transporters by intracellular signaling events such as phosphorylation and ATP binding.

In summary, our studies have provided unique insights into the functional roles of CIC CBS domains and the R-helix linker in regulating channel properties. We propose a mechanism in which the R-helix linker interacts with CBS2 α2 and provides a struc-
tural pathway by which C-terminus conformational changes could modulate the conformation of the R-helix and associated structures and lead to changes in channel function. This hypothesis is sup-

ported by recent studies by Martinez and Maduke, which demonstrated that a point mutation in the R-helix linker of ClC-Kb abolishes extracellular Ca²⁺ activation of the channel and alters bromide permeability. Additional mutagenesis, modeling, biochemical and structural studies are needed to directly test this hypothesis. It will be particularly important to determine the role played by the R-helix linker and CBS domain conformation in mediating the reg-

ulation of CIC channels and transporters by intracellular signaling events such as phosphorylation and ATP binding.

**Methods**

**Transfection and whole cell patch clamp recording.** HEK293 (human embryonic kidney) cells were cultured in 35 mm diam-

eter tissue culture plates in Eagle’s minimal essential medium (MEM; Gibco, Gaithersburg, MD) containing 10% fetal bovine serum (Hyclone Laboratories, Inc., Logan, UT), non-

essential amino acids, sodium pyruvate, 50 U/ml penicillin and 50 µg/ml streptomycin. After reaching 50–70% confluency, cells were transfected using FuGENE 6 (Roche Diagnostics Corporation, Indianapolis, IN) containing 1 µg GFP and 1–8 µg of channel cDNAs ligated into pcDNA3.1. Point mutants were generated using a QuikChange Site-Directed Mutagenesis Kit (Stratagen, La Jolla, CA) and deletion mutants were generated by polymerase chain reaction (PCR)-based fusion strategies. For single amino acid substitutions, local structure was ana-

lyzed using a homology model (Fig. 6) to identify mutations that would be most disruptive to putative local interactions. All mutants were confirmed by DNA sequencing. Experimental protocols were performed on at least two independently transfected groups of cells.

Following transfection, cells were incubated at 37°C for 24–30 h. Approximately 2 h prior to patch clamp experiments, cells were detached from growth plates by exposure to Hank’s balanced salt solution containing 0.25% trypsin and 1 mM EDTA (Gibco) for 45 sec. Detached cells were suspended in MEM and then plated onto poly-L-lysine coated cover slips. Plated cover slips were placed in a bath chamber mounted onto the stage of an inverted microscope. Cells were visualized by fluorescence and differential interference contrast microscopy.

Transfected cells were identified by GFP fluorescence and patch clamped using a bath solution that contained 90 mM NMDG-Cl, 5 mM MgSO₄, 1 mM CaCl₂, 12 mM HEPES free acid, 8 mM Tris, 5 mM glucose, 80 mM sucrose and 2 mM glutamime and was titrated to pH 7.4 with HEPES free acid and Tris (297–303 mOsm), and a pipette solution containing 116 mM NMDG-Cl, 2 mM MgSO₄, 20 mM HEPES titrated to pH 7.0 with CsOH, 6 mM CsOH, 1 mM EGTA titrated to pH 8.3–8.5 with CsOH, 2 mM ATP, 0.5 mM GTP and 10 mM sucrose (pH 7.2, 274–280 mOsm). Low pH bath solution contained 90 mM NMDG-Cl, 5 mM MgSO₄, 1 mM CaCl₂, 5 mM glucose, 80 mM sucrose and 2 mM glutamine and was titrated to pH 5.9 using HEPES free acid, (297–303 mOsm).

Patch electrodes with resistances of 3–7 MΩ were pulled from 1.5 mm outer diameter slanized borosilicate microhematocrit tubes. Whole cell currents were measured with an Axopatch 200B (Axon Instruments, Foster City, CA) patch clamp amplifier. Electrical connections to the patch clamp amplifier were made using Ag/AgCl wires and 3 M KCl/agar bridges. Data acquisition and analysis were performed using pClamp 10 software (Axon Instruments).

**Voltage clamp protocols and data analysis.** Whole cell currents were evoked by stepping membrane voltage for 1 sec between -120 mV and +100 mV in 20 mV increments from a holding potential of 0 mV. Test pulses were followed by a 1 sec interval at 0 mV. Current-to-voltage relationships were con-

structed from mean current values recorded over the last 25 msec of each test pulse. For pre-depolarization studies, cells were held at holding voltages of -20 mV to 60 mV for 3 sec and then stepped to -120 mV for 3 sec to activate channels. Peak hyperpolarization-induced current amplitude was measured over a 100 msec interval. Pseudo-steady-state current was measured over the last 20 msec of the -120 mV test pulse.

Channel activation voltages were estimated from current-to-voltage relationships. A line was drawn by linear regression analy-

sis of currents measured between 20 mV and 100 mV. A second line was drawn by linear regression analysis of currents measured between the first voltage at which inward current was detected and a second voltage 20 mV more negative. The intercept of these two lines is defined as the activation voltage.

**Homology modeling.** A homology model for the CLH-3b ytoplasmic C-terminus was generated based on the crystal structure of the CLC-Ka C-terminus (Protein Data Bank accession code 2PFI). We chose CLC-Ka as a template because biochemi-

cal data indicate that this crystal structure represents the native
dimeric state of the channel. 13 BCL::Align software 33 with manual adjustment of the linker region between CBS1 and CBS2 was used to align amino acids 549–615 and 783–841 of CLH-3b with the corresponding amino acids 542–681 of CLC-Ka. Modeller software 34 version 9.3 was used to develop 120 candidate structures for CLH-3b. The best-scoring structure by Modeller’s molpdf objective function was chosen and side chain orientations optimized from rotamer libraries were determined using SCWRL 3.35. The model was refined with AMBER 9 using 500 steps of energy minimization. 56,57

As shown in Figure 1, CBS1 and CBS2 of CLH-3a and CLH-3b have identical sequences except for the last six amino acids of CBS2, which encompasses part of the second α helix (α2). To investigate how splice variation affects α2 helical structure, we used Rosetta software 33,34 and the lowest energy structure generated by Modeller. The modeled domain includes amino acids 620–686 and 753–811 for CLH-3a and amino acids 549–615 and 783–841 for CLH-3b. Rosetta was used to predict de novo 20,000 candidate structures of amino acids 803–811 for CLH-3a and amino acids 620–686 and 753–811 for CLH-3b. These nine amino acids include the C-terminus of CBS2 and the following three amino acids, which are also alternatively spliced in CLH-3a and CLH-3b (Fig. 1). The remainder of the modeled region surrounding these nine amino acids was included for structural context and held in a fixed conformation.

**MTSET inhibition experiments.** [2-(Trimethylammonium) ethyl] methanethiosulfonate bromide (MTSET; Toronto Research Chemicals, Toronto, Ontario, Canada) was dissolved in water as a 400 mM stock and stored in 40 μl aliquots at -80°C until use. Working solutions with an MTSET concentration of 1 mM were made immediately before each experiment. Whole cell current amplitude in MTSET experiments was recorded by stepping membrane voltage to -100 mV for 500 msec every 1 sec from a holding potential of 0 mV.

**Statistical analysis.** Data are presented as means ± S.E. Statistical significance was determined using Student’s two-tailed t test for unpaired means. p values of ≤0.05 were taken to indicate statistical significance.

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