Myotonia-related mutations in the distal C-terminus of ClC-1 and ClC-0 chloride channels affect the structure of a poly-proline helix

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Myotonia is a state of hyperexcitability of skeletal-muscle fibres. Mutations in the CIC-1 Cl− channel cause recessive and dominant forms of this disease. Mutations have been described throughout the protein-coding region, including three sequence variations (A885P, R894X and P932L) in a distal C-terminal stretch of residues [CTD (C-terminal domain) region] that are not conserved between CLC proteins. We show that surface expression of these mutants is reduced in Xenopus oocytes compared with wild-type CIC-1. Functional, biochemical and NMR spectroscopy studies revealed that the CTD region encompasses a segment conserved in most voltage-dependent CLC channels that folds with a secondary structure containing a short type II poly-proline helix. We found that the myotonia-causing mutation A885P disturbs this structure by extending the poly-proline helix. We hypothesize that this structural modification results in the observed alteration of the common gate that acts on both pores of the channel. We provide the first experimental investigation of structural changes resulting from myotonia-causing mutations.

Key words: CIC-1 chloride channel, common gating, myotonia, NMR, poly-proline helix, skeletal muscle.

INTRODUCTION

Mutations in the human skeletal-muscle voltage-dependent chloride channel CIC-1 cause recessive (Becker) [1] and dominant (Thomsen) myotonia congenita [1,2]. In both cases, mutations impair Cl− channel function [3,4]. In skeletal muscle, Cl− conductance performs a role similar to that of K+ conductance in neurons, i.e. stabilization of resting membrane voltage. Thus, if this conductance is reduced, the muscle becomes hyperexcitable [5].

Recessive myotonia is caused by mutations that lead to a loss or strong reduction of Cl− channel function. In contrast, most dominant mutations affect the common gate that acts on the two pores of the homodimeric Cl− channel, shifting the voltage dependence of activation to positive voltages [6,7].

The conformational changes that occur during common gating are still unclear [8]. Mutations in many parts of the coding region of CIC-1 modify the properties of the common gate. These include transmembrane regions, for instance helices at the contact sites between the two channel subunits [9,10]. Mutations impairing common gate function also occur in the large cytoplasmic C-terminus which contains two CBS (cystathionine-β-synthase) domains and which is present in all eukaryotic CLC proteins [11–15]. The recent crystal structure of the cytoplasmic C-terminus of the CIC-0 Cl− channel [16] provides a framework for further research. The structure revealed that the CBS1 and CBS2 domains associate as in other CBS-containing proteins [11,17]. Analytical ultracentrifugation experiments indicated that the CBS1–CBS2 complexes form homodimers, thereby extending the 2-fold transmembrane topology of the transmembrane region [16]. A model of the proposed quaternary structure has been proposed [16] in analogy with the crystal structure of a tandem CBS domain protein from Thermotoga maritima [18]. Interestingly, mutations in the residues from the predicted interface between CBS domains of distinct CLC subunits dramatically affect the common gate of CLC channels [11,19].

In eukaryotic CLC proteins, the sequence following the second CBS domain is of variable length, from seven (CIC-K) to 117 (CIC-1) amino acids. Unfortunately, due to proteolysis, this region is absent from crystals of the CIC-0 C-terminus [16]. In this region, a stretch of proline-rich residues of unknown significance for CIC-1 channel function has been identified [15]. The mutations in this region found in myotonia alter channel function by diverse mechanisms: some reduce macroscopic Cl− conductance with nearly normal channel properties of the remaining currents (e.g. R894X) [14,20–22]; others alter the voltage dependence of channel activation (e.g. A885P, present in the myotonic goat model) [23].

Here, we used a combination of structural, biochemical and functional methods to examine the molecular defect caused by these mutations. Our results provide the first structural information of the region after the second CBS domain of CLC proteins and reveal the likely structural alterations that are caused by several mutations found in myotonia.

EXPERIMENTAL

Solid-phase synthesis of the peptides

All peptides were synthesized using the Fmoc (fluoren-9-ylmethoxy carbonyl)/tBu (t-buty1) strategy and 10 mol of TBTU [O-(benzotriazol-1-yl)-N,N,N′,N′-tetramethyluronium tetrafluoroborate]/HOBT (1-hydroxybenzotriazole)/DIEA (N,N-di-isopropylethylamine) as the coupling mixture. Amino acids were...
purchased from Novabiochem. The final peptide was cleaved with 94% TFA (trifluoroacetic acid)/2.5% water/2.5% EDT (ethanedithiol)/1% TIS (tri-isopropylsilane) and precipitated in cold diethyl ether. The crude material of each reaction was purified by preparative HPLC until a sample with a purity of at least 96% (characterized by HPLC–MS) was obtained.

NMR

Peptide assignments were obtained from two sets of two-dimensional experiments run either in 90% water/10% D2O (NOESY–TOCSY pair) or in 100% D2O [ROESY (rotating-frame Overhauser enhancement spectroscopy)–TOCSY pair], acquired at 800 MHz and at 285 K. Resonance assignment was done manually using the program CARA[24] (see Supplementary Figure 1 at http://www.BiochemJ.org/bj/403/bj4030079add.htm). Peptide models were generated with CNS using the restraints from the NOESY and the ROESY experiments. In each case a set of ten structures was calculated and submitted to a water refinement protocol.

Functional and biochemical methods in *Xenopus* oocytes

Capped complementary RNA of CLC channels (CIC-1: 10 ng; CIC-0: 1 ng) was expressed in *Xenopus* oocytes as described in [11]. Measurements were done in ND96 medium (96 mM NaCl, 2 mM KCl, 1.8 mM CaCl2, 1 mM MgCl2 and 5 mM Hepes buffer at pH 7.4). Pulse protocols for tail current analysis of CIC-1 were performed as described in [11]. Fast and slow gates of CIC-0 were studied using protocols described previously [8,11]. The Clampfit program (Axon Instruments) was used for fitting. Two batches of oocytes, each batch containing *n* = 6 oocytes, were used. Inside-out patch–clamp experiments were performed using an EPC-7 amplifier (List) and the custom GePulse acquisition program. The solutions had the following composition: the standard intracellular solution contained (in mM) 100 NMDG (N-methyl-d-glucamine)-Cl, 2 MgCl2, 10 Hepes and 2 EGTA at pH 7.3; the standard extracellular (pipette) solution contained (in mM) 100 NMDG-Cl, 5 MgCl2 and 10 Hepes. PBS/1% Triton X-100-solubilized extracts from oocytes were obtained and the protein was quantified with a BCA (bicinchoninic acid) kit (Pierce). Equal amounts of proteins were separated on SDS/7.5% polyacrylamide gels and electrotransferred (Bio-Rad Laboratories) to PVDF membranes. A ponceau staining was performed on PVDF membranes to control protein loading. After incubation with primary and secondary antibodies, antigen–antibody complexes were visualized by enhanced chemiluminescence (ECL®; Amersham) and exposed on X-ray films. Quantification of protein levels was done using ImageJ. Surface expression using antibodies against HA (haemagglutinin) epitope and chemiluminescence was performed as previously described [11]. Results shown were obtained with *n* = 10 oocytes in at least two batches of oocytes.

Molecular biology

Constructs were made using recombinant PCR with two mutagenesis primers and two external primers flanking SacI and EcoRI restriction sites. The final PCR fragment was digested with SacI and EcoRI and cloned in the vector pTLN-N-HA-CIC-1-HAloop already digested with the same restriction sites and were then sequenced. In CIC-1, an HA epitope was introduced between helices L and M after the glycosylation site (HAloop), which resulted in the sequence VKHAGYPYDVPDYADEP (HA epitope in boldface). In the N-terminus of CIC-1, an HA epitope was added after the second amino acid to improve detection by Western blot. These tags did not significantly affect conductance or the voltage dependence of gating [11].

RESULTS

Effect of mutations associated with myotonia on the C-terminal distal region of CIC-1

Among mammalian CLC proteins, the skeletal-muscle Cl-channel CIC-1 has the longest C-terminal region after the second CBS domain (Figure 1A). Here we refer to this stretch as the CTD (C-terminal domain) region. Three myotonia-causing mutations
have been described in the CTD region of CLC-1: A885P (identified in a myotonic goat model [23]), P932L [25,26] and the truncation R894X [14,20–22]. The A885P mutation is located in a segment downstream of the CBS2 domain that is conserved among CLC-0, CLC-1 and CLC-2 (Figure 1A). The R894X is localized a few residues downstream from this segment, whereas the P932L mutation is more C-terminal (Figure 1A). Each of these mutations was introduced in human CLC-1 and studied by voltage-clamping and biochemical analyses after heterologous expression in Xenopus oocytes. Monitoring with an HA antibody (Figure 1B) revealed that all three mutations reduced surface expression, with the most drastic effect observed for the truncation R894X. Compared with wild-type CLC-1, protein expression levels as monitored by Western blotting were similar for mutants A885P and P932L and dramatically reduced for mutant R894X (Figure 1B, inset).

Similar to what has been described previously [23], mutation A885P gave functional Cl− channels with reduced macroscopic current amplitude and altered gating behaviour. The voltage of half-maximal activation was $-10.6 \pm 1.1 \text{ mV} (n = 7)$ [$V_{0.5}$ (wild-type) $= -68.5 \pm 1.2 \text{ mV} (n = 68)$], without any change in the slope of $P_{\text{rev}}$ as a function of voltage. Analogous to what has been reported [14,21,26], mutations R894X and P932L did not change the voltage dependence of activation significantly, although they reduced current amplitudes in parallel with their surface expression levels (results not shown).

**Functional analysis of the CTD region**

To study the molecular basis of the defect caused by these mutations, we investigated which structures within the CTD region were crucial for channel function. First, we performed C-terminal deletion mutagenesis combined with functional and biochemical measurements. Consistent with the poor conservation between CLC proteins, the whole CTD region of CLC-1 could be deleted (in truncation E871X), without abolishing CLC-1 function (Figure 2A) and surface expression (Figure 2B). However, truncation of the protein before residue 858 (truncations 858X, 854X, 845X and 826X, which all occur within CBS2) abolished function (Figure 2A). In the truncation construct M854X, loss of function was due to a lack of surface expression (Figure 2B) and a reduced total expression (Figure 2B, inset).

It was puzzling that the myotonia-related mutant R894X severely reduced Cl− channel function (Figure 2A) and plasma membrane expression (Figures 1B and 2B), whereas deletion of further amino acids, i.e. in construct 871X, increased currents compared with R894X. We studied this effect by performing a fine mapping to find out at which position truncations lead to larger currents compared with R894X. Starting from the end of the protein until amino acid 891, C-terminal truncations progressively reduced channel function (Figure 2A). A further deletion (F887X) considerably increased CLC-1 currents, thereby defining the point of increased function around residues 891 and 887. Additional deletions reduced channel function mildly, until the truncations described above that abolished currents.

To further characterize these deletions, we examined the voltage of half-maximal activation of several of the truncation constructs (Figure 2C). As a rule, the C-terminal truncation constructs that yielded large conductance levels displayed shifts of the voltages of half-maximal activation ($V_{0.5}$) towards positive potentials. Starting with the truncation at amino acid 887, $V_{0.5}$ shifted to positive voltages, being maximally shifted by truncation 871X. However, $V_{0.5}$ was closer to wild-type values in the truncation 864X, corresponding to the last $\alpha$-helix of the second CBS domain.
Addition of consecutive alanines after amino acid 871 of ClC-1 dependent of the neighbouring amino acids. indicate that the function of this conserved segment may be in-

Mapping a key segment in voltage-dependent CLC channels

Comparison of the amino acid sequence between CLC proteins close to the identified border revealed an amino acid segment that is conserved in ClC-0, ClC-1 and ClC-2, but that is absent from CLC-K channels and CLC transporters (Figure 1A). Interestingly, the myotonia-related mutation A885P is located in this conserved segment. In ClC-1, the amino acid sequence of this segment is LRPPLASFR, corresponding to amino acids 880–888 (Figure 1A). We studied this conserved segment in more detail by mutagenesis experiments of human ClC-1 and analysed the voltage dependence of mutants in this segment (Table 1). Starting from the end of CBS2, consecutive amino acids were deleted from amino acids 872–880. Only after deleting nine amino acids [Del (872–875)] was the voltage dependence of activation significantly different from wild-type ClC-1. Secondly, changing the conserved segment, adding consecutive amino acids 880–888 from CLC-K channels and CLC transporters (Figure 1A). Two experiments suggested that ClC-0 can be used as a model to study the functional effect on gating of ClC-1 mutations in this segment. First, swapping the CTD domain of ClC-1 by the CTD domain of ClC-0 resulted in a channel with a voltage of half-maximal activation of 787 (D787X, the first amino acid after the conserved segment) resulted in a channel with wild-type voltage dependence (Figure 3). Proceeding further, with the ClC-0 truncation F785X (at the end of the conserved segment), produced a channel with altered gating (Figure 3). The mutant showed inverted voltage dependence, becoming activated by hyperpolarizing voltage pulses (Figure 3). A similar phenotype has been described for the mutant A783P, that reproduces in ClC-0 the above described myotonia related A885P mutation in ClC-1 [27]. We concluded that mutations in this conserved segment alter voltage dependence of gating in ClC-0 and ClC-1.

This segment is highly conserved between ClC-1 and ClC-0, with the exception of an arginine to proline change (Arg881 in ClC-1, Figure 1A). Two experiments suggested that ClC-0 can be used as a model to study the functional effect on gating of ClC-1 mutations in this segment. First, swapping the CTD domain of ClC-1 by the CTD domain of ClC-0 resulted in a channel with a voltage of half-maximal activation of −65.9 ± 3 mV (n = 6), not significantly different from wild-type ClC-1. Secondly, changing this arginine to a proline in ClC-1 or the opposite in the equivalent position in ClC-0, yielded channels with wild-type properties (results not shown). Hence, we concentrated our further analysis on the ClC-0 channel, for which single-channel analysis is possible.

Three-dimensional structure of the conserved segment

To gain insights into the structure of this segment, we performed two types of experiments: (i) we changed each single residue from 82

Table 1 Voltage dependence of currents from several mutants around a conserved segment (in italics in Figure 1) after CBS2 of the CTD of ClC-1

<table>
<thead>
<tr>
<th>Mutations</th>
<th>Voltage</th>
</tr>
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<tbody>
<tr>
<td>Consecutive deletion of amino acids after 871 of ClC-1</td>
<td></td>
</tr>
<tr>
<td>Del (872)</td>
<td>−56.2 ± 3.1 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Del (872–873)</td>
<td>−63.9 ± 3.1 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Del (872–874)</td>
<td>−73.1 ± 3.3 mV (n = 13, three batches)</td>
</tr>
<tr>
<td>Del (872–875)</td>
<td>−52 ± 2.5 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Del (872–876)</td>
<td>−54.8 ± 3.9 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Del (872–877)</td>
<td>−55.9 ± 2.4 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Del (872–878)</td>
<td>−44.2 ± 3.3 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Del (872–879)</td>
<td>−58.2 ± 3.5 mV (n = 6, one batch)</td>
</tr>
<tr>
<td>Del (872–880)</td>
<td>−9.1 ± 3.3 mV (n = 7, two batches)</td>
</tr>
<tr>
<td>Alanine scanning of three amino acids from 872–886 of ClC-1</td>
<td></td>
</tr>
<tr>
<td>Ala (872–874)</td>
<td>−78.9 ± 4.0 mV (n = 8, three batches)</td>
</tr>
<tr>
<td>Ala (875–877)</td>
<td>−55.8 ± 5.1 mV (n = 10, two batches)</td>
</tr>
<tr>
<td>Ala (878–880)</td>
<td>−35.9 ± 2.5 mV (n = 9, two batches)</td>
</tr>
<tr>
<td>Ala (881–883)</td>
<td>−30.4 ± 5.7 mV (n = 9, three batches)</td>
</tr>
<tr>
<td>Ala (884–886)</td>
<td>−13.2 ± 4.3 mV (n = 6, two batches)</td>
</tr>
<tr>
<td>Addition of consecutive alanines after amino acid 871 of ClC-1</td>
<td></td>
</tr>
<tr>
<td>Ala (1)</td>
<td>−64.8 ± 1.6 mV (n = 11, two batches)</td>
</tr>
<tr>
<td>Ala (2)</td>
<td>−73.6 ± 2.0 mV (n = 11, two batches)</td>
</tr>
<tr>
<td>Ala (3)</td>
<td>−78.8 ± 2.6 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Ala (4)</td>
<td>−74.2 ± 1.5 mV (n = 12, two batches)</td>
</tr>
<tr>
<td>Ala (5)</td>
<td>−73.6 ± 1.9 mV (n = 11, two batches)</td>
</tr>
<tr>
<td>Ala (6)</td>
<td>−70.2 ± 2.9 mV (n = 13, two batches)</td>
</tr>
</tbody>
</table>

Figure 3 Similar role of the conserved segment in the CIC-0 channel

Two-electrode voltage-clamp traces from oocytes expressing the indicated mutants. They were evoked by a pulse protocol consisting of a prepulse to 60 mV, followed by a series of test pulses ranging from 80 to −140 mV and a final pulse to −100 mV. Note the inverted voltage dependence of CIC-0 and mutant F785X, but not D787X.
the conserved segment in CIC-0 to alanine. Then, we functionally examined the gating of these mutants in order to find functional alterations that could follow periodic changes, as expected for a segment with a determinate structure [28] (Figure 4A), and (ii) we used NMR spectroscopy to resolve the structure of a peptide corresponding to the conserved segment from CIC-0 (Figure 4B).

Only three of these CIC-0 mutations produced an inwardly rectifying phenotype, as described for the truncation F785X. No functional changes were detected in the other alanine mutants (Figure 4A and results not shown). The three mutations that caused inward rectification affected the hydrophobic amino acids Leu778, Leu782 and Phe785. The spacing (778–782–785), i.e., a period of three to four residues, suggests that these three amino acids are located on the same face of a helix.

We used NMR to determine the structure of a peptide corresponding to the conserved segment. Similar approaches have been performed in other structural studies of proline-rich segments [29,30]. In addition, we reasoned that this approach was convenient for our segment, since the results obtained with the various mutants of CIC-1 and CIC-0 (Table 1 and Figure 3) were compatible with the notion that the conserved segment has a secondary structure that it is independent of the adjacent amino acids.

The interpretation of the NMR data (Supplementary Figure 1) indicated that the secondary structure of the peptide consisted of a short poly-proline helix type II [comprising the first leucine (L2) and the contiguous three prolines (P3 to P5)] followed by a disordered region (Figure 4B, left). In the lowest energy structure the three hydrophobic amino acids identified above were on the same face (Figure 4B, right).

The CIC-0 mutation A783P causes structural changes in the conserved segment

We hypothesized that the CIC-0 mutation A783P influences the structure of the conserved segment. To test this hypothesis, we synthesized a CIC-0 peptide variant in which this alanine residue was mutated to proline and studied its structure by NMR spectroscopy (Figure 5).

For the mutant peptide, we detected NOEs (nuclear Overhauser effects) consistent with a poly-proline helical conformation (all prolines were in the trans conformation) that spanned the LPPPLP region of the peptide (supplementary Figure 1). This finding indicated that the mutation extended the poly-proline helical segment (Figure 5, middle), which in the wild-type peptide ends after the first three prolines of the motif (Figure 5, top).

As a control, we studied the mutation S784P, one residue downstream of the alanine that is mutated in goat myotonia. In this second mutant peptide, a mixture of cis and trans conformations was observed for the new proline residue (instead of the only trans conformation detected in the peptide containing the mutation...
The left panels show two-electrode voltage-clamp traces from oocytes expressing the indicated mutants evoked by the pulse protocol described in Figure 4. Note the inverted voltage dependence of wild-type compared with that of the mutant A783P. The non-overlapping currents during the 60 mV prepulse observed for mutant S784P result from an accelerated closing kinetics of the slow gate, which is otherwise very similar to wild-type ClC-0. Right panels show schematic representations of each peptide studied by NMR. Peptide models are calculated on the basis of the NOEs observed in solution (see Supplementary Figure 1). The poly-proline conformation was clearly observed in solution, resulting in a pattern of NOEs that showed all prolines in the trans conformation. As a result of the observed restraints the structure calculation displays a high convergence in the region comprising the prolines, while the remaining residues are ill defined in solution. Six structures with less energy are represented, out of ten calculated for each peptide. Figures were generated with MOLMOL [33] (http://www.molmol.org).

Figure 5  NMR structure of a ClC-0 peptide containing a mutation identified in myotonia reveals structural differences caused by the mutation

The conserved segment is involved in the common gate

Finally, to study which gating process was altered by mutations in this conserved segment, we analysed the functional effects of a ClC-0 mutation of one hydrophobic amino acid from the conserved segment, L782A, using patch–clamp recordings.

 Following conventional pulse protocols, it was impossible to distinguish fast and slow gating in mutant L782A (Figure 7A). Currents activated rapidly at negative voltages, with a time constant of approx. 16 ms at $-140$ mV (Figure 7A) and deactivated at positive voltages with similarly fast kinetics. The steady-state
Distal C-terminal myotonia-related mutations

Figure 7 Mutations in the conserved segment affect the common gate

(A) Patch-clamp traces evoked from a patch containing several ClC-0 L782A channels (average of eight recordings). After a prepulse to −140 mV the voltage was stepped up to various values ranging from +80 to −140 mV. Finally, a ‘tail’ pulse was applied to +80 mV. In (B), the normalized initial current measured during the tail pulse is plotted towards the prepulse voltage and fitted by a Boltzmann function resulting in the parameters $V_{0.5} = -75$ mV, $z = 1.7$, $p_{min} = 0.0$. Comparable values for wild-type ClC-0 obtained with a longer pulse protocol to drive the slow gate into steady state are $V_{0.5} = -100$ mV, $z = 1.8$, $p_{min} \approx 0.1$ (results not shown). (C) The Figure shows recordings from a patch containing several L782A channels (−60 mV) and from a single-channel wild-type ClC-0 patch (−100 mV). Traces are at identical scales. The presence of more than one channel for the L782A patch was verified by pulses to more negative voltages (results not shown) at which the open probability increases markedly (see B). The short ‘burst’ duration of the mutant compared with wild-type is evident. Similar short burst durations were seen in all patches of mutant L782A ($n = 4$). The comparably long burst of wild-type ClC-0 is well documented (e.g. [34]). The dotted line indicates the closed channel current level.

The voltage dependence of the mutant (Figure 7B) was very similar to that of the slow gate of wild-type CIC-0 (see the legend of Figure 7). Single-channel recording showed that the macroscopic behaviour of mutant L782A mainly reflected a drastic alteration of the slow gate: bursts of openings at negative voltages were much briefer than for wild-type CIC-0 (Figure 7C). This finding precluded a quantitative analysis of the fast gate. Thus whether mutant L782A, in addition to this drastic effect on the slow gate, also alters the fast gate remains to be elucidated. However, our electrophysiological analysis demonstrates that mutations in this conserved segment alter the common gate that acts on both pores of the channel.

DISCUSSION

Using a combination of structural and functional methods, we have analysed three distinct mutations that cause myotonia.

The CIC-1 mutation A885P identified in a dominant myotonic goat model [23], is located in a protein segment that is conserved among the plasma membrane CLC channels CIC-0, CIC-1 and CIC-2. By studying wild-type and mutant peptides encompassing this segment in CIC-0 and CIC-1, we elucidated how this mutation changes the local protein structure. The most relevant difference between the wild-type and mutant CIC-0 and CIC-1 structures may be the reduced flexibility of the mutant. The core of the poly-proline helix in the wild-type peptide is formed by the first leucine (L2) and the following three contiguous prolines (P3, P4, P5 in Figure 4). Although this poly-proline helix contains a leucine or an arginine residue, this is not surprising, since bulky amino acids are found often to intercalate in proline-rich sequences, because they maintain the poly-proline structure [31].

The subsequent leucine (Leu') may prolong this helix, depending on the amino acids that follow. When Leu' is followed by an alanine (wild-type), a residue allowing a certain flexibility, the
rigid part of the peptide is limited to the first leucine and the contiguous three prolines. In the peptide containing an alanine to proline mutation, in which Leu6 is followed by a proline, the poly-proline helix is extended by two extra residues (Leu6 and Pro7). In contrast, in the peptide where the Ser6 is changed to proline, Leu6 is followed by an alanine (Ala7), retaining some of the flexibility shown by the wild-type. It has been found that bulky side chains hinder the isomerization of prolines, whereas alanine does not contribute to maintain the poly-proline helical conformation, in agreement with our results [32]. Consistent with these data, mutant S784P shows nearly wild-type gating (Figure 5). In contrast, individually mutating the proline residues from this conserved segment to alanine did not result in gating alterations (Figure 4). From these experiments, we can conclude that it is critical for the function of the channel that the last part of the conserved segment is flexible and that a certain degree of flexibility in the first part of the segment does not impair a proper assembly of the motif.

Our interpretation of these results is based on the assumption that the structure of these peptides is a good model of the structure of this segment in the full-length channel. We believe that this is likely, at least for the part forming the stable poly-proline helix secondary structure, because our mutagenesis studies around this segment indicated that this region is independent from the neighbouring amino acids (Table 1) and the comparison with other poly-proline helices found in other proteins [29,30]. Proline-rich sequences are frequently found in inter-domain regions of proteins, and also on the surface of globular proteins of eukaryotic organisms, where they act as a binding site for interacting proteins [29]. When the number of consecutive prolines in a protein is equal or longer than four, this segment adopts a poly-proline type II helical conformation [29]. This conformation is very stable (prolines are sterically forced to adopt a trans-conformation), and this probably explains why poly-proline segments are found so often in otherwise unstructured regions of proteins [30]. However, ultimate proof of our hypothesis would be a full-length structure of the channel or at least of the complete cytosolic C-terminus. Unfortunately, the CTD region is absent from the recently solved crystal structure of the C-terminus of ClC-0, which lacks all amino acids after CBS2 [16]. To expand our knowledge of the function of this region, we expressed and purified a protein containing the CTD region and analysed it with NMR spectroscopy. Although the presence of secondary structural elements (some helical content) was ascertained, the lack of tertiary structure precluded a complete interpretation of the NMR data (R. Estévez and M. J. Macías, unpublished work).

At present, it is difficult to explain how extending the poly-proline helix type II alters the common gating of the channel. We can only speculate that this conserved segment of the channel may be binding to another part of the channel, and that changing the structure of this segment may alter this binding, causing gating alterations.

We have shown (Figure 1) that all myotonia-causing mutations identified in the CTD region of ClC-1 reduced plasma membrane expression, with the truncation R894X showing the strongest effect. Interestingly, truncating additional amino acids upstream of Arg894 partially rescued surface expression. Importantly, the relative increase in surface expression started in the conserved segment. In particular, three large hydrophobic amino acids (Leu774, Leu784 and Phe785 in ClC-0) were identified in this segment, mutations in which led to alterations in common gating kinetics. Although the last part of the conserved segment was found to be disordered in the peptide, we propose that these three residues lie on the same face and form a hydrophobic patch. This result leads us to speculate that all mutations identified in the CTD region will cause a distortion of the structure of this segment.

In addition to the conserved segment, the ClC-1 CTD region presents other proline-rich segments. Interestingly, another mutation found in myotonia (P932L) is present in one of these proline-rich segments, and patients carrying this mutation have a dystrophic phenotype in addition to myotonia [25]. Recent electrophysiological analysis of this mutant in HEK-293 cells (human embryonic kidney cells) did not reveal any functional defect [26]. Here, we show that mutating this residue reduces slightly the surface expression of the channel without changing its voltage dependence of gating. Probably, the identification of interacting proteins in this region is critical to understand in detail the severe phenotypes observed in patients carrying this particular mutation.

In summary, the present study is the first analysis of structural alterations caused by several myotonia-causing mutations. Certainly, whether these structural alterations have implications in ligand interactions is a question that still remains elusive and that will require additional experiments.
Distal C-terminal myotonia-related mutations


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