A sequence in the carboxy-terminus of the α1C subunit important for targeting, conductance and open probability of L-type Ca^{2+} channels

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Abstract The role of the 80-amino acid motif 1572–1651 in the C-terminal tail of α1C Ca^{2+} channel subunits was studied by comparing properties of the conventional α1C\text{77} channel expressed in HEK-tsA201 cells to three isoforms carrying alterations in this motif. Replacement of amino acids 1572–1651 in α1C\text{77} with 81 non-identical residues leading to α1C\text{86} impaired membrane targeting and cluster formation of the channel. Similar to α1C\text{86}, substitution of its 1572–1598 (α1C\text{77K}) or 1595–1652 (α1C\text{77L}) segments into the α1C\text{77} channel yielded single-channel Ba^{2+} currents with increased inactivation, reduced open probability and unitary conductance, when compared to the α1C\text{77} channel. Thus, the C-terminal sequence 1572–1651 of the α1C subunit is important for membrane targeting, permeation and open probability of L-type Ca^{2+} channels.

Key words: Class C-type Ca^{2+} channel; Carboxyl tail; Targeting; Conductance; Inactivation; Fluorescence microscopy; GFP-labeled α1C subunit

1. Introduction

The voltage-gated L-type Ca^{2+} channel is an essential part of signal transduction systems in many cell types, triggering essential processes, such as muscle contraction [1,2] or neuronal gene expression [3–6]. It is composed of the pore-forming α1C subunit and auxiliary β- and δ-subunits [7,8]. The carboxy-terminus of the α1C subunit has attracted much attention [9–11] because of its potential involvement in channel gating. Two human splice variants of the principal 2138-amino acids pore-forming α1C subunit, α1C\text{77} and α1C\text{86}, differ in their carboxy-terminal tail sequence by 80 amino acid residues in positions 1572–1651 (Table 1; [12,13]). When expressed in oocytes, these two channel splice variants exhibit strong differences in inactivation properties [10]. Whole-cell Ba^{2+} currents of the α1C\text{86} channel inactivate significantly faster than those through α1C\text{77}. Furthermore, with Ca^{2+} as charge carrier, inactivation of the current through α1C\text{77} is greatly accelerated in contrast to the α1C\text{86} inactivation which is essentially Ca^{2+}-independent. Two independent determinants for Ca^{2+}-induced inactivation were identified within this sequence by segment exchange analysis [11], one located in the L-motif (1572–1598) and the other in the K-motif (1595–1651) (Table 1). A calmodulin binding IQ region (1624–1635) within the K-motif has been shown to be important for Ca^{2+}-dependent inactivation [14–16].

To further analyze the role of amino acids 1572–1651 in the regulation of Ca^{2+} channel activity, we studied here expression, subcellular localization as well as electrophysiological properties of the α1C\text{77} and α1C\text{86} channels and of two derived segmental mutants α1C\text{77K} and α1C\text{77L} (Table 1; [11]) in the HEK-tsA201 mammalian expression system. The results of our study indicate that the sequence 1572–1651 in the carboxy-terminal tail of the α1C subunit determines not only channel inactivation, but also membrane targeting, single-channel conductance and open probability of L-type Ca^{2+} channels.

2. Materials and methods

2.1. Materials

EGFP was purchased from Clontech (Heidelberg, Germany). The cDNA of the CD8 receptor (EBO pCD Leu2) was kindly provided by Richard Horn (Thomas Jefferson University Medical School, Philadelphia, PA, USA). The cDNA of the fusion construct of EGFP and the pleckstrin homology domain of phospholipase C β1 (EGFP-PH) was a gift from Tobias Meyer (Durham, North Carolina, USA). Tissue culture media and reagents were purchased from Life Technology, Vienna, Austria. (–)BAYK 8644 was from Research Biochemical International, Vienna, Austria and all other chemicals from Sigma, Vienna, Austria. Magnetic beads carrying antibodies against the CD8 receptor were purchased from Dynal, Hamburg, Germany.

2.2. Molecular biology

2.2.1. Preparation of eukaryotic expression plasmids encoding the α1C channel isoforms. 77pcDNA3 coding for α1C\text{77} [17] used for eukaryotic transfection was inserted into plasmid pcDNA3 (Invitrogen) and contained the Kozak consensus sequence (5’-cctgga-3’) upstream of the initiation codon. 86pcDNA3 was prepared by replacing the SfiI (3341)-AatII (5494) fragment of plHCC77 with the corresponding fragment [10]. To prepare 77KpcDNA3 and 77LpcDNA3, the PHLCC77K and PHLCC77L plasmids [11], respectively, were digested with BamHI, blunt-ended using the Klenow DNA polymerase, digested with PvuII (2760) and the obtained 3’-terminal 3.9-kb fragments were ligated into 77pcDNA3 to replace the corresponding fragment in the PvuII (2760)NotI (blunt-ended) cassette.

2.2.2. Construction of the 5’-GFP-labeled α1C\text{77} and α1C\text{86}. A 5’-terminal HindIII linker upstream Kozak sequence and 3’-terminal BglII linker were incorporated into the flanking regions of the open reading frame of the pGreen Lantern-1 DNA (Life Technology, Gai-
thersburg, MD, USA) ORF by PCR using sense 5'-aagatgcggccac-
catgag-3' and antisense 5'-agatgttgtatagctgctc-3' primers, respectively. The HindIII/BglII fragment (green Lantern-1 cassette) was ligated into the 77pcDNA3 vector at HindIII/BamHI sites so that the 5'-GFP-77pcDNA3 construct encoded the enhanced GFP fused to a C177 via RSAT tetrapeptide. 5'-GFP-56pcDNA3 was prepared by replacing the 3.9-kb PvuII/IolI fragment of 5'-GFP-77pcDNA3 with the respective fragment of pHLC86B.

Nucleotide sequences of all PCR products, as well as ligation sites were verified using the ABI Prisms™ Dye Terminator Cycle Sequencing Kit with AmpliTag DNA Polymerase (Perkin Elmer, Norwalk, USA).

2.3. Cell culture and transfection of tsA201 cells
tsa201 cells were cultured in DMEM medium supplemented with streptomycin (100 μg/ml) and 10% FCS in a humidified atmosphere (95%) at 5% CO2 and 37°C. Cells were used for 12–14 passages and were passed every 4 days. Transfection was performed using SuperFect (Qiagen, Hilden, Germany). In brief, cells exhibiting confluence of about 30–50% were transfected with 2.5 μg of total cDNA (molar ratio of αC:β2:δ:EGF-CD3 = 1:1:6:14:2.5:0.7). Transfection efficiency was estimated by the fraction of cells showing EGFP fluorescence, and was in the range of 20–60%. Expression of CD8 receptor and binding of CD8 antibody-coated beads (Dynal) was used as visual marker to identify cells in electrophysiological experiments. Binding of the beads (range 3–15 per cell) was estimated to occur in 1–5% of the total number of cells.

2.4. Microscopy

Cells transfected with αC constructs carrying green fluorescent protein (GFP) at their amino-terminus were imaged either by conventional fluorescence microscopy or by two-photon confocal fluorescence microscopy [18]. Conventional fluorescence microscopy was performed on an inverted microscope (Axiovert 100TV, Zeiss, Oberkochen, Germany) equipped with a 100× NA 1.3 objective (Zeiss) and a Seniscam slow scan CCD camera (PCO CCD Imaging, Kellheim, Germany). In confocal microscopy, cover slips carrying the cells were mounted onto a piezo-driven scanning stage (PI, Waldbronn, Germany) mounted on an inverted microscope (AxiovertS 100TV, Zeiss) equipped with a 100× NA 1.3 objective (Zeiss). The size of the point-spread function was measured to be 0.25 μm in lateral and 0.8 μm in axial direction. The fluorescence was excited by the light from a fs-Ti:sapphire laser (Tsunami, Spectra Physics, Mountain View, CA, USA) tuned to 910 nm. The average power at the objective was kept to 5 mW equivalent to an average intensity of ~6 MW/cm². Appropriate filters in the detection path (XLP800+LP750, Newport, Irvine, USA; BG39-3, Schott, Mainz, Germany) allowed to reliably discriminate the GFP fluorescence. The fluorescence was detected by a photon-counting avalanche diode (SPCM-AQR-141, EG&G, Dummer,erry, Canada) in combination with an interface card (PMS300, Becker and Hickl, Berlin, Germany). The dwell time of the counter was set to 10 ms.

2.5. Electrophysiology

Single-channel patch-clamp recordings [19] were obtained with a List L/E PC 7 amplifier from successfully transfected tsA201 cells. Ba2+ currents through single Ca2+ channels were recorded in the cell-attached configuration according to [20]. The dihydropyridine Ca2+ channel activator (–) BAYK 8644 (2.5 μM) was included in the pipette solution to facilitate channel activity in the cell-attached configuration. This allowed for correct estimation of the number of channels in the patch. Pipettes (GC150F-7.5F) were fabricated from borosilicate glass (Clark Medical Instruments, Pangbourne, UK) and had resistances of 4–6 MΩ. Sigmaocto was used to reduce pipette capacitance. Single-channel currents of the channels were evoked by repetitive depolarizations (0.66 Hz) applied for 0.480 s from a holding potential of −80 to 0 mV (αC(77) and 10 mV (αC(86), αC(77), and αC(86)) and filtered at 4 kHz. All experiments were performed at room temperature.

2.6. Analysis of electrophysiological data

Single channel current amplitudes were determined by analyzing all point amplitude histograms fitted by Gaussian curves employing pClamp 6.0.3 software. Estimation of Ca2+ channel activity was primarily based on determination of overall channel open probability (P). P was calculated from the time course of mean channel activity Np (N is the total number of channels; [21]) determined for each depolarizing voltage pulse defined as a sweep, and obtained by averaging all Np values divided by the number of channels in the patch. The number of channels N was estimated by the method described in [22]. Channel availability (Pp) and open probability (Po) according to P = PpPo were calculated in multi-channel experiments employing a recently developed analysis method [20,22,23]. Pp was defined by the ratio (×100 in %) of depolarizing sweeps with channel activity to the total number of sweeps, and Po represents a channel’s open probability in non-blank sweeps. Open times were determined from non-overlapping channel openings [20]. Since Pp was usually <10% and N ≤ 3 in experiments used for open times analysis, open time distributions are expected to be biased. Distributions were corrected for missed events by excluding events <1 ms [20]. Open times histograms were fitted with multieponential functions employing a Chebyshev fitting routine from pClamp 8 software.

2.7. Statistics

Averaged results are presented as mean ± S.E.M. for the number of experiments usually given in parentheses. Two-tailed t-test was used for statistical comparison considering differences statistically significant at P < 0.05.

3. Results

The human Ca2+ channel isoforms αC(77) and αC(86) (Table 1) were transiently co-expressed with β2s and δ subunits in HEK-tsA201 cells. The expression pattern of these channels and electrophysiological properties of their single-channel currents were analyzed to characterize the functional role of the sequence 1572–1651. To narrow structures within the sequence 1572–1651 critical for functional properties, two segmental mutants, i.e. αC(77L) and αC(77K), were studied in which

Table 1 Structure of the variable parts in the carboxy-terminal tail of the investigated αC subunits

<table>
<thead>
<tr>
<th>αC(77)</th>
<th>αC(77L)</th>
<th>αC(77K)</th>
<th>αC(86)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IKTENHGQNAELRA1IK1W1RTKMTKLLQVQFPAGDDEV1GKYATFL1QTEYRFRKLKQEGGQLGKPSQRNALS</td>
<td>ETESGQWQYQKAEASLRRKSSHSF</td>
<td>SSSHKSTPKNKLSSGGSSTGWEDARALEQVLARGGCGWLSLREERGHPHPLGLF</td>
<td>ETESGQWQYQKAEASLRRKSSHSF</td>
</tr>
</tbody>
</table>

Amino acid sequences of αC(77) (1572–1651) and αC(86) (1572–1652) are shown on the top and bottom, respectively. Indicated amino acids of αC(86) replace the respective residues in the amino acid sequence of αC(77). In αC(77L) and αC(77K) subunits, indicated segments of αC(86) replace the respective motifs L (1572–1598) and K (1595–1651) of the αC(77) subunit. Note that the overlapping 4-amino acid segment SSSH has been proven not to contribute to the kinetics, voltage- or Ca2+-dependence of inactivation ([11]). Bold residues are located in identical positions between αC subunits.
spected by eye using an inverted fluorescence microscope with 100× magnification for the GFP-labeled αIC,77 and αIC,86 did not significantly affect Ca2+ channel function in both whole-cell and single-channel experiments (data not shown). Conventional fluorescence microscopy revealed a different pattern of subcellular distribution for the GFP-labeled αIC,77 and αIC,86 subunits, when co-expressed with α2δ and β2a subunits (Table 2). The αIC,77 subunit was mainly located in the plasma membrane with the formation of punctate clusters visible in 16% of these cells. In contrast, the αIC,86 subunit showed mainly cytoplasmic localization besides some plasma membrane staining without any indication of cluster formation. Fig. 2 shows two-photon confocal fluorescence images of equatorial slices through the αIC-expressing tsA201 cells which illustrate the typical patterns of subcellular distribution as observed also in the conventional fluorescence microscopy. The fluorescence is attributed to GFP-labeled proteins as confirmed by images of cells transfected with non-fluorescent subunits (Fig. 2A). As control for plasma membrane localization, expression of the GFP-labeled pleckstrin homology (PH) domain that has been found to be enriched in the plasma membrane [26] is shown (Fig. 2B). The difference in cellular localization between αIC,77 and αIC,86 as given in Table 2 is clearly evident from comparison of Fig. 2C,D. Expression of each αIC subunit alone resulted in similar expression patterns showing mainly intracellular localization (data not shown).

### 3.2. Single-channel characteristics of αIC,77 and αIC,86 channels

Single-channel currents were measured in the cell-attached configuration with Ba2+ as charge carrier in the presence of 2.5 μM (-)-BAYK 8644. Fig. 3A and B depict consecutive single-channel traces with corresponding ensemble average currents of αIC,77 and αIC,86, respectively. Consistent with whole-cell characteristics (data not shown; [10]), the ensemble average currents of αIC,86 showed a markedly faster inactivation rate as compared to the αIC,77 channel. Surprisingly, current–voltage relationships revealed a significantly lower single-channel Ba2+ conductance for the αIC,86 channel (Fig. 3C). A unitary conductance of 25.5 ± 0.8 pS (n = 14) was determined for αIC,86, whereas 29.3 ± 1.1 pS (n = 5) was calculated for the αIC,77 channel. Reversal potentials, however, did not significantly differ between these channels (Fig. 3C, inset) yielding 51.5 ± 4.1 and 57.8 ± 2.4 mV for αIC,77 and αIC,86, respectively.

To further evaluate the impact of the sequence 1572–1651 on Ca2+ channel gating characteristics, overall open probability (P0) which is the product of open probability (P0) times...
availability \( (P_s) \) were determined separately, and mean open times were estimated to reveal their contribution to the inactivation of channels (Fig. 4). It was found that \( P \) of \( K_{1C,77} \) was about 4-fold higher than that of the \( K_{1C,86} \) channel (Fig. 4A). This significant difference in \( P \) was mainly due to divergent \( P_o \), whereas \( P_s \) was not changed between the channels (Fig. 4B). To evaluate whether inactivation visible in ensemble average currents is mediated by a reduction in open times with progressive depolarization time, we performed separate analysis of open time distributions (Fig. 4C,D) restricted to 0–240 and 241–480 ms of depolarization. Open time distributions of both channels were fitted by two exponentials suggesting the existence of at least two distinct open states which apparently do not significantly change with time. The short open time constant and the proportion between short and long open time constants were rather similar for both channels. However, the long open time constant was almost 1.5-fold greater for \( K_{1C,77} \) as compared to the \( K_{1C,86} \) channel. The average mean open times of both channels (see Table 3) within the first and second time interval of depolarization did not significantly change suggesting an increase in average mean closed times to account for the increase in inactivation rate of \( K_{1C,86} \).

To further narrow critical regions within the sequence 1572–1651 that appears to be important for the functional properties characterized above, single-channel characteristics of the two segmental mutants \( K_{1C,77L} \) and \( K_{1C,77K} \) were studied.

3.3. Single-channel characteristics of the segmental mutants \( K_{1C,77L} \) and \( K_{1C,77K} \)

Single-channel characteristics of \( K_{1C,77K} \) and \( K_{1C,77L} \) channels are shown in Figs. 5 and 6, and summarized in Table 3. Consecutive current traces and their corresponding ensemble average currents (Fig. 5A,B) reflect the rapid inactivation observed in whole-cell currents (data not shown; [11]). Analysis of single-channel current–voltage relationships for \( K_{1C,77K} \) and \( K_{1C,77L} \) channels showed no significant difference either in their reversal potentials or in unitary conductances (Fig. 5C). Their conductances were also not significantly different to the \( K_{1C,86} \) channel, but were clearly distinct to that of the \( K_{1C,77} \) channel. Analysis of single-channel gating kinetics (Fig. 6) revealed that \( P \) of \( K_{1C,77K} \) and \( K_{1C,77L} \) (Fig. 6A) amounted on the average to 1.3 ± 0.4% \((n = 10)\) and 2.3 ± 1.1% \((n = 8)\) and were estimated to be not significantly different. Consistently, both \( P_s \) and \( P_o \) of the \( K_{1C,77K} \) and \( K_{1C,77L} \) channel were not significantly different (Fig. 6B). However, the time-restricted open time analysis (Fig. 6C) revealed a substantial reduction in mean open times of \( K_{1C,77K} \) with increasing depolarization time, which was not found with the \( K_{1C,77L} \) channel.
4. Discussion

In this study, we characterized functional expression of the human L-type Ca^{2+} channel splice variants $\alpha_{1C;77}$ and $\alpha_{1C;86}$ as well as their segmental mutants $\alpha_{1C;77K}$ and $\alpha_{1C;77L}$. Our results strongly suggest that the sequence 1572–1651 in the carboxy-terminal tail of $\alpha_{1C}$ is critical for subcellular protein targeting and significantly affects single-channel conductance and gating of L-type Ca^{2+} channels (summarized in Table 3).

4.1. Expression and subcellular localization pattern of splice variants

Estimation of Ca^{2+} channel density in transfected tsA201 cells points to stronger expression of the $\alpha_{1C;77}$ channel as compared to $\alpha_{1C;86}$. Consistently, we observed a different pattern of subcellular distribution for these splice variants using fluorescent $\alpha_{1C}$ subunits. The GFP-$\alpha_{1C;77}$ subunit, when expressed together with auxiliary subunits, was mainly localized in the plasma membrane, and in 16% of cells formed clusters, whereas the $\alpha_{1C;86}$ subunit was detected predominantly in the cytoplasm. This distinct expression pattern did not change with time between the first and third day after transfection.

![Fig. 3. Single-channel characteristics of the $\alpha_{1C;77}$ and $\alpha_{1C;86}$ channels. Single-channel traces obtained from (A) $\alpha_{1C;77}$ and (B) $\alpha_{1C;86}$ channels by repetitive depolarizations (0.66 Hz) from a holding potential of −80 to 0 and 10 mV, respectively. Corresponding average currents are shown at the bottom of A and B. Dotted lines indicate zero current level. C: Corresponding current–voltage relationships of typical experiments with unitary conductance ($\sigma$) as indicated. Inset in C depicts respective mean values showing a significant difference ($P<0.02$) between unitary conductances of the $\alpha_{1C;77}$ and $\alpha_{1C;86}$ channels, but not in their reversal potentials.](image-url)
(data not shown). Hence, targeting of the $\alpha_{1C,86}$ subunit to the plasma membrane appears to be less efficient. Previously, the targeting of $\alpha_{1C}$ subunit to plasma membrane has been demonstrated to be critically dependent on the interaction with the $\beta$ subunit [25,27–29]. Accordingly, in the absence of $\beta_{2a}$, both $\alpha_{1C,77}$ and $\alpha_{1C,86}$ subunits were diffusely distributed in the cytoplasm (data not shown). The interaction with the $\beta$ subunit, however, is expected to be similar for $\alpha_{1C,77}$ and $\alpha_{1C,86}$, as both subunits retain invariant $\beta$-subunit binding domain in the cytoplasmic linker between repeats I and II of the core protein. Our results are consistent with the idea that the carboxyl tail of the conventional $\alpha_{1C,77}$ channel contains additional structures critical for membrane targeting that appear to be disrupted in $\alpha_{1C,86}$. Furthermore, the segmental mutants $\alpha_{1C,77K}$ and $\alpha_{1C,77L}$ exhibited an expression pattern which resembles that of $\alpha_{1C,86}$ (data not shown). Recent data supporting our findings suggest that the C-terminus of the $\alpha_{1C}$ subunit is required for proper plasma membrane targeting of L-type Ca$^{2+}$ channels ([30]; B. Flucher, personal communication). Our study allowed this additional critical structure to be narrowed to the 80-amino acid motif 1572–1651 at the $\alpha_{1C}$ tail.

4.2. Electrophysiological features of $\alpha_{1C}$ splice variants and segmental mutants in single-channel experiments

The $\alpha_{1C,77}$ channel exhibited a significantly slower inactivation than $\alpha_{1C,86}$ and the two segmental mutants $\alpha_{1C,77K}$ and $\alpha_{1C,77L}$, as judged from ensemble average currents. These data substantiated the role of the sequence 1572–1651 and of its L- and K-segments (see Table 1) as molecular determinants of voltage-dependent inactivation [11]. In addition, current–voltage relationships implied the sequence 1572–1651 as affecting the unitary conductance. The three $\alpha_{1C}$ channel isoforms which exhibited rapid inactivation of Ba$^{2+}$ currents showed

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**Fig. 4.** Gating characteristics of single $\alpha_{1C,77}$ and $\alpha_{1C,86}$ channels. (A) Overall open probability and (B) availability as well as open probability of the respective channel. Numbers of experiments are indicated in parentheses and significant differences are marked by an asterisk. Open time histograms of (C) $\alpha_{1C,77}$ and (D) $\alpha_{1C,86}$ channels constructed from the indicated number (n) of experiments for the first (0–240 ms) and second (241–480 ms) time interval of depolarizations. Open times (τ) and proportions (P) were calculated from the respective biexponential open time distributions fitted by a Chebyshev algorithm.
a significantly lower single-channel conductance compared to the \( \alpha_{1C,77} \) channel. We discovered that substitution of segment L (1572–1598) or K (1595–1651) in \( \alpha_{1C,77} \) by the corresponding sequences of the \( \alpha_{1C,86} \) subunit was sufficient to reduce unitary conductance of the resulting \( \alpha_{1C,77L} \) and \( \alpha_{1C,77K} \), respectively, channels to almost the same level as that seen in the \( \alpha_{1C,86} \) channel. This finding was unexpected, as the pore-structure remains identical in all channels studied which is consistent with no significant difference in their reversal potentials. The fact that each of the two adjacent cytoplasmic motifs K and L of the \( \alpha_{1C,86} \) tail reduced unitary conductance to the same level seen in the \( \alpha_{1C,86} \) channel suggests that there are two independent determinants for this effect. In view of recent results obtained with a Shaker channel [31], a link between cytoplasmic domains and permeation properties appears possible. However, the sequence 1572–1651 is not a critical determinant of permeation, as its deletion still yields functional channels [32], but rather has a modulatory role.

4.3. Molecular determinants of gating kinetics

The 4–7-fold lower \( P \) of the \( \alpha_{1C,86} \) channel and of its segmental mutants \( \alpha_{1C,77K} \) and \( \alpha_{1C,77L} \) compared to the \( \alpha_{1C,77} \) channel is mediated by a decrease in \( P_o \), whereas \( P_S \) is not significantly changed. As \((-)\)-BAYK 8644 was used in all

Fig. 5. Single-channel characteristics of single \( \alpha_{1C,77K} \) and \( \alpha_{1C,77L} \) channels. Single-channel traces of (A) \( \alpha_{1C,77K} \) and (B) \( \alpha_{1C,77L} \) channels recorded by repetitive depolarizations (0.66 Hz) from a holding potential of \(-80\) to 10 mV. Corresponding average currents are shown at the bottom of A and B. Dotted lines denote zero current level. C: Corresponding current–voltage relationships of typical experiments are shown with estimated unitary conductances (\( \sigma \)) and reversal potentials (\( E_{rev} \)) as indicated. Inset depicts respective mean values showing no significant difference between both unitary conductances and reversal potentials of \( \alpha_{1C,77K} \) and \( \alpha_{1C,77L} \) channels.
experiments, the difference in $P_o$ could be at least in part due to different affinities of the channels to dihydropyridines [33]. To largely compensate for these different affinities to (−)-BAYK 8644, supramaximal concentrations of 2.5 μM (−)-BAYK 8644 were used. Hence, the sequence 1572–1651 appears to determine $P_o$ of the Ca$^{2+}$ channel. In accordance, Ca$^{2+}$-induced inactivation of the channel, which critically depends on this sequence [10], is manifested in a reduction of $P_o$ while leaving $P_s$ unchanged [20].

To evaluate the impact of the sequence 1572–1651 on channel experiments, the difference in $P_o$ could be at least in part due to different affinities of the channels to dihydropyridines [33]. To largely compensate for these different affinities to (−)-BAYK 8644, supramaximal concentrations of 2.5 μM (−)-BAYK 8644 were used. Hence, the sequence 1572–1651 appears to determine $P_o$ of the Ca$^{2+}$ channel. In accordance, Ca$^{2+}$-induced inactivation of the channel, which critically depends on this sequence [10], is manifested in a reduction of $P_o$ while leaving $P_s$ unchanged [20].

To evaluate the impact of the sequence 1572–1651 on channel

![Graph and Table]

**Fig. 6. Gating characteristics of single $\alpha_{1C,77K}$ and $\alpha_{1C,77L}$ channels.** (A) Overall open probability and (B) availability as well as open probability of the respective channel. Numbers of experiments are indicated in parentheses. Open time histograms of (C) $\alpha_{1C,77K}$ and (D) $\alpha_{1C,77L}$ channels constructed from the indicated number (n) of experiments for the first (0–240 ms) and second (241–480 ms) time interval of depolarizations. Open times ($\tau$) and proportions ($P$) were calculated from the respective biexponential open time distributions fitted by a Chebyshev algorithm.

### Table 3
Overview of the investigated properties of the $\alpha_{SC,77}$ and $\alpha_{SC,86}$ channels and their segmental mutants $\alpha_{1C,77K}$ and $\alpha_{1C,77L}$

<table>
<thead>
<tr>
<th>$\alpha_{IC}$</th>
<th>Subcellular localization</th>
<th>Ba$^{2+}$ current inactivation</th>
<th>$\sigma$ (pS)</th>
<th>$E_{rev}$ (mV)</th>
<th>$P_s$ (%)</th>
<th>$P_o$ (%)</th>
<th>$\tau_{mean}$ (ms)</th>
<th>$\tau_{mean}$ (ms)</th>
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<tbody>
<tr>
<td>77</td>
<td>mainly plasma membrane</td>
<td>slow</td>
<td>29.3 ± 1.1</td>
<td>51.5 ± 4.1</td>
<td>66.3 ± 6.0</td>
<td>10.7 ± 2.9</td>
<td>1.87</td>
<td>1.86</td>
</tr>
<tr>
<td>86</td>
<td>mainly cytoplasm</td>
<td>rapid</td>
<td>25.5 ± 0.8</td>
<td>57.8 ± 1.3</td>
<td>60.4 ± 5.3</td>
<td>3.9 ± 1.0</td>
<td>1.33</td>
<td>1.44</td>
</tr>
<tr>
<td>77K</td>
<td>mainly cytoplasm</td>
<td>rapid</td>
<td>24.2 ± 1.1</td>
<td>55.3 ± 2.8</td>
<td>70.8 ± 11.9</td>
<td>2.7 ± 1.8</td>
<td>0.84</td>
<td>0.79</td>
</tr>
<tr>
<td>77L</td>
<td>mainly cytoplasm</td>
<td>rapid</td>
<td>24.3 ± 1.0</td>
<td>58.9 ± 1.9</td>
<td>52.5 ± 4.9</td>
<td>3.7 ± 1.4</td>
<td>1.51</td>
<td>1.50</td>
</tr>
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</table>

Subcellular localization has been assessed using 5′-GFP-labeled $\alpha_{IC}$ subunits; $\sigma$ unitary conductance in 96 mM Ba$^{2+}$; $E_{rev}$, reversal potential; $P_s$, channel availability; $P_o$, channel open probability; $\tau_{mean}$, mean open times within 0–240 ms and 241–480 ms intervals of depolarization time.
nel inactivation, open time histograms were constructed for 0–240 and 241–480 ms time intervals of the depolarizing pulse. Both $\alpha_{1C.77}$ and $\alpha_{1C.56}$ as well as the segmental mutants $\alpha_{1C.77L}$ and $\alpha_{1C.77K}$ yielded open time histograms that implied two open states. Mean open times within the first and the second interval are rather similar (see Table 3) pointing to an increase in mean closed times to account for the faster second interval are rather similar (see Table 3) pointing to an increase in mean closed times to account for the faster inactivation rate of $\alpha_{1C.56}$ and its segmental mutants $\alpha_{1C.77K}$ and $\alpha_{1C.77L}$. Only the $\alpha_{1C.77K}$ channel showed a clear reduction in open time during the 241–480 ms time interval, which is, however, balanced by a concomitant change in the proportion between these open times.

In summary, the amino acid sequence 1572–1651 in the carboxy-terminal tail of the human $\alpha_{1C}$ subunit not only determines voltage- and $\text{Ca}^{2+}$-dependent inactivation, but, in addition, represents a key structure for L-type $\text{Ca}^{2+}$ channel membrane targeting, ion conductance and kinetics. The motifs K and L are independently critical for unitary conductance and open probability.

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