

Carboxy-Terminal Truncations Modify the Outer Pore Vestibule of Muscle Chloride Channels

Simon Hebeisen* and Christoph Fahlke*[†]

*Department of Physiology, RWTH Aachen, Aachen, Germany; and [†]Centro de Estudios Científicos, Valdivia, Chile

ABSTRACT Mammalian CIC-type chloride channels have large cytoplasmic carboxy-terminal domains whose function is still insufficiently understood. We investigated the role of the distal part of the carboxy-terminus of the muscle isoform CIC-1 by constructing and functionally evaluating two truncation mutants, R894X and K875X. Truncated channels exhibit normal unitary conductances and anion selectivities but altered apparent anion binding affinities in the open and in the closed state. Since voltage-dependent gating is strictly coupled to ion permeation in CIC-1 channels, the changed pore properties result in different fast and slow gating. Full length and truncated channels also differed in methanethiosulphonate (MTS) modification rate constants of an engineered cysteine at position 231 near the selectivity filter. Our data demonstrate that the carboxy-terminus of CIC channels modifies the conformation of the outer pore vestibule.

INTRODUCTION

CIC-type anion channels and transporters are expressed in prokaryotic and eukaryotic cells and fulfill diverse physiological functions. Nine human CIC channels (hCIC-1 to hCIC-7, hCIC-Ka, and hCIC-Kb) play roles in the regulation of cell excitability (1–3), transepithelial chloride transport (4,5), and pH adjustment in certain cell compartments (6–8). Bacterial homologs function as secondary active, proton-coupled chloride transporters (9) and were proposed to provide an electrical shunt for an outwardly directed proton pump linked to amino acid decarboxylation (10).

Prokaryotic and eukaryotic CIC isoforms exhibit a unique, evolutionarily conserved double-barreled architecture with two ion conduction pathways, each formed by a single subunit (11–13). In contrast to prokaryotic isoforms, all eukaryotic channels have large carboxy-termini that reside in the cytoplasm and contain two cystathionine- β -synthase (CBS) domains (14,15). For CIC-0 and CIC-1, complete removal of the carboxy-terminus has been demonstrated to render the corresponding protopore nonfunctional (16–20). On the other hand, point mutations or deletions within the region distal to the second CBS result in functional channels with altered voltage-dependent gating (18,21). Since CIC channels are thought to be gated by the permeant anion and to lack an endogenous voltage sensor (22–24), modification of gating properties by a cytoplasmic domain is intriguing. Mutations within the distal carboxy-terminus might alter CIC channel gating either by affecting intrinsic pore properties or, alternatively, by modifying inherently voltage-dependent conformational changes of the channel protein that were not identified before.

We generated and functionally characterized two truncation mutants of hCIC-1 (R894X and K875X) (Fig. 1 A). R894X represents a naturally occurring mutation identified in patients with recessive myotonia congenita (25), and K875X hCIC-1 was engineered to remove the whole region distal to the second CBS domain. Both mutations affect the interaction between the permeating anion and its binding sites within the ion conduction pathway, and these alterations result in changed channel gating. Our results demonstrate that the carboxy-terminus modifies the outer pore vestibule of CIC-type chloride channels and further support the concept of CIC channel gating by permeant anions.

MATERIAL AND METHODS

Expression of WT and mutant hCIC-1

Truncations were introduced into the plasmid pRc/CMV-hCIC-1 containing the full length wild-type (WT) hCIC-1 cDNA (26) by polymerase chain reaction-based strategies. For all constructs, polymerase chain reaction-amplified sequences were verified by direct sequencing, and two independent mutant clones were used for expression studies. Some of the experiments were performed with WT and mutant pSVL-hCIC-1 constructs obtained by subcloning the respective cDNA into the pSVL vector (Amersham Biosciences, Piscataway, NJ). Mutant and WT hCIC-1 channels were transiently expressed in tsA201 cells with passage numbers between 3 and 20 and examined typically 2 days after transfection with a calcium phosphate precipitation method (27). To identify cells with a high probability of expressing recombinant ion channels, cells were cotransfected with a plasmid encoding the CD8 antigen and incubated 5 min before use with polystyrene microbeads precoated with anti-CD8 antibodies (Dynabeads M-450 CD 8, Dynal, Great Neck, NY) (28). For some experiments examining WT hCIC-1, stably transfected human embryonic kidney (HEK) 293 cells (29) were used. Functional properties of channels stably expressed in HEK 293 cells and those transiently expressed in HEK 293 or tsA201 cells were indistinguishable from one another.

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Address reprint requests to Christoph Fahlke, Abteilung Neurophysiologie, Medizinische Hochschule Hannover, Carl-Neuberg-Str. 1, 30625 Hannover, Germany. Tel.: 49-511-532-2774; Fax: 49-511-532-2776; E-mail: fahlke.christoph@mh-hannover.de.

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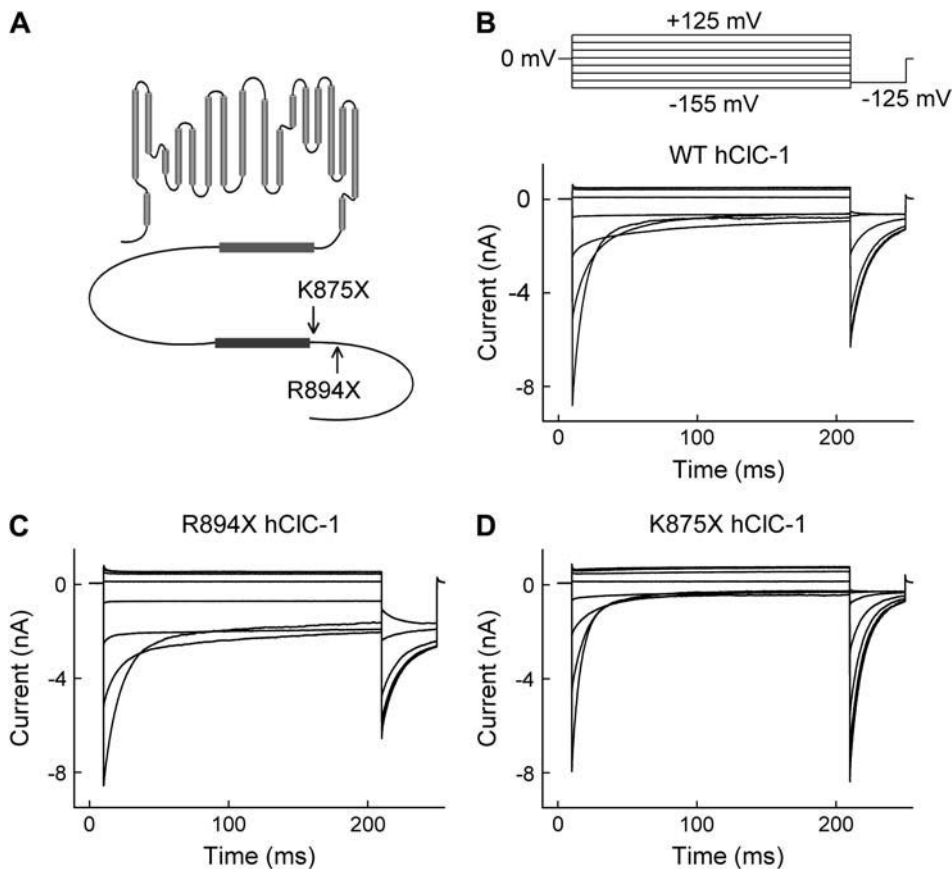


FIGURE 1 Carboxy-terminal truncations alter gating properties of hClC-1 channels. (A) Topology and (B–D) representative current recordings of WT and mutant channels.

Electrophysiology

Standard whole-cell, excised inside-out or outside-out patch clamp recordings were performed as previously described (19,30). The standard extracellular solution contained (in mM): NaCl (140), KCl (4), CaCl_2 (2), MgCl_2 (1), HEPES (5), pH 7.4; the standard intracellular (in mM): NaCl (120), MgCl_2 (2), EGTA (5), HEPES (10), pH 7.4. Cells were clamped to 0 mV between two test sweeps. To exchange the external or the internal solution, cells/excised patches were moved into the stream of solutions containing external or internal solutions whose composition was altered by substituting NaCl equimolarly with sodium glutamate, NaI, or NaNO_3 . Fast opening rate constants of K875X hClC-1 channels (see Figs. 3 C and 4 C) were determined using an intracellular solution containing 230 mM NaCl and an external solution with 250 mM NaCl that was partly substituted with Na glutamate to change $[\text{Cl}^-]_o$. To study block of chloride currents by I^- or NO_3^- , instantaneous current amplitudes were determined after a 200 ms prepulse to +75 mV before and after application of a given anion concentration. Agar bridges were used to connect the amplifier and the bath solution. Junction potentials calculated using the JPCalc software (Dr. P. Barry, University of South Wales, Sydney, Australia (31)) and offset potentials measured at the end of each experiment were used to correct results. Modification by 2-(trimethylammonium)ethyl-methanethiosulphonate (MTSET) and 2-aminoethyl-methanethiosulphonate (MTSEA) was performed as described (32). Modification rates were determined at holding potentials of −80 mV, 0 mV, and +75 mV.

Data analysis

Data were analyzed by a combination of PulseFit and PulseTools (HEKA Electronics, Lambrecht, Germany), Clampfit (Axon Instruments, Foster

City, CA), and SigmaPlot (Jandel Scientific, San Rafael, CA) programs. Instantaneous current amplitudes were calculated by extrapolating deactivation time courses with biexponential functions to the moment of the voltage step. To determine the unitary current amplitude and the open probabilities of mutant hClC-1 channels, nonstationary noise analysis was performed as described (19,30). To determine the time course of current deactivation, a sum of two exponentials and a time-independent value ($I(t) = a_1 \exp(-t/\tau_{\text{fast}}) + a_2 \exp(-t/\tau_{\text{slow}}) + c$) were fit to data recorded during a series of hyperpolarizing voltage steps from a holding potential of 0 mV. To obtain the voltage dependence of the relative open probability, the instantaneous current amplitudes after a voltage step to −125 mV were normalized by their maximum value and plotted versus the preceding potential. The absolute open probability at 0 mV determined by noise analysis was then used to calculate the voltage dependence of the open probability (19) (Fig. 2 A). The voltage dependences of the opening of the fast and the slow gate were distinguished as described (33) (Fig. 2, C and D). The relative open probability of the slow gate is determined by measuring normalized tail current amplitudes after 200-ms prepulses to various potentials and a short fixed pulse to positive voltages inserted before the test step (Fig. 2 C, inset). For a separation of fast and slow gating, the short pulse has to be long enough to fully activate the fast gate without significantly affecting the slow gate. Time constants of fast and slow activation were determined at +180 mV for $[\text{Cl}^-]_o = 10$ mM or 150 mM using envelope protocols (33) (data not shown), demonstrating that a pulse duration of 500 μs satisfies these requirements for all constructs. Activation curves for the fast and for the slow gate were fit with the sum of a voltage-independent minimum open probability (P_{min}) and a voltage-dependent term: $I(V) = \text{Amp}/(1 + e^{(V - V_{0.5})/kV}) + P_{\text{min}}$.

The voltage dependences of the fast deactivation time constants (Fig. 2 B) were fitted in two steps. First, the τ -V curve of R894X hClC-1 was fitted with

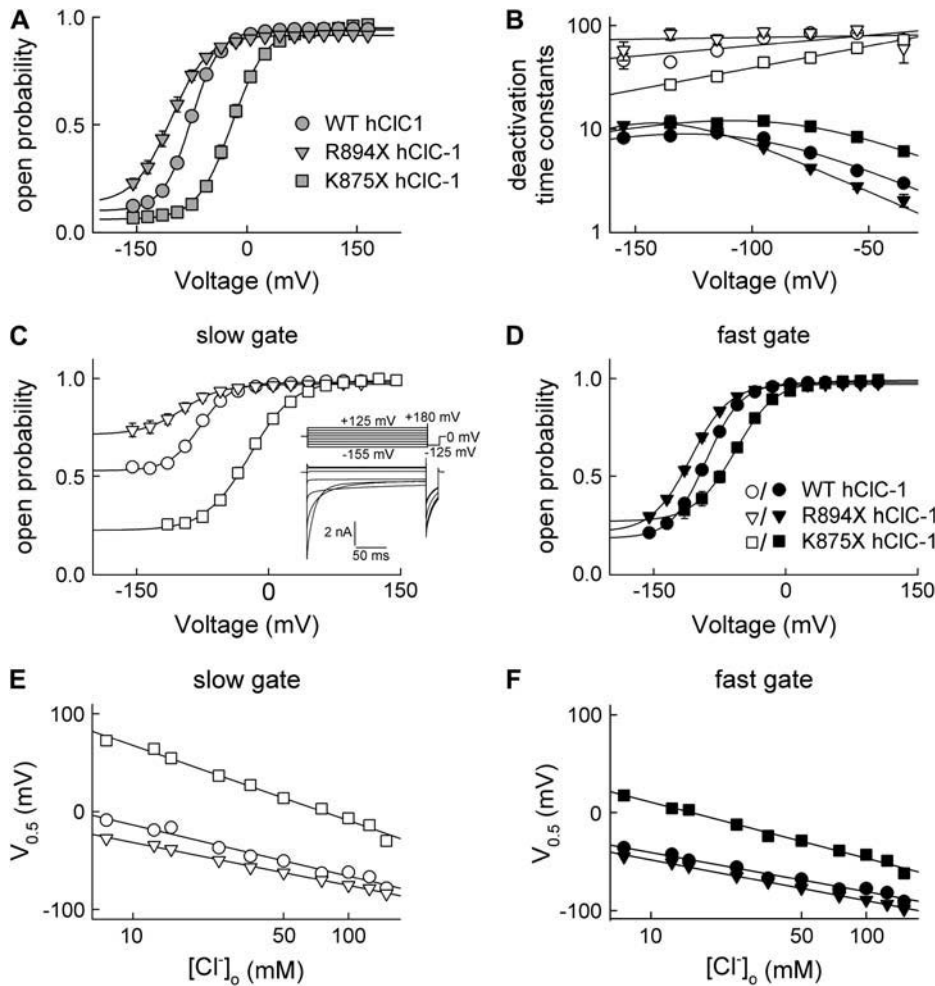


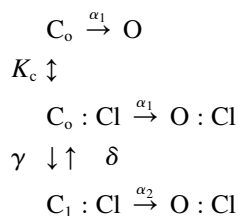
FIGURE 2 Carboxy-terminal truncations modify the voltage dependence of fast and slow gating. (A) Voltage dependence of the open probability. (B) Voltage dependence of the fast (solid symbols) and slow (open symbols) deactivation time constants for WT (●), R894X (▼), and K875X (■) hCIC-1 channels. Mean \pm SE ($n > 58$). The voltage dependence of the fast deactivation time constants was fitted with $\tau = 1/(\alpha_o e^{a_{\text{aco}}V/kT} + \beta_o e^{-b_{\text{zo}}V/kT})$, giving an equivalent gating charge of 1 for the opening rate constant and 0.75 for the closing time constants. Slow time constants were fitted with single exponentials. (C and D) Voltage dependence of the slow (C) and the fast (D) gate of WT (●) and mutant (R894X, ▼, and K875X, ■) channels. (E and F) Plot of the midpoint of the voltage dependence of slow (E) and fast (F) gating versus $[\text{Cl}^-]_o$. Mean \pm SE ($n > 8$).

$\tau = 1/(\alpha_o e^{a_{\text{aco}}V/kT} + \beta_o e^{-b_{\text{zo}}V/kT})$, with α_o and β_o as opening/closing rate constants at 0 mV, and a and b as equivalent gating charges defining the voltage dependences of these rate constants. The voltage dependence of the deactivation time constants of WT and K875X hCIC-1 was then fitted to the same equation with fixed equivalent gating charges. Slow time constants were fitted with single exponentials (Fig. 2 B). Because of the limited range of values, the voltage dependence of opening and closing rate constants cannot be deduced from these values.

We calculated opening rate constants of the fast and the slow gate for various voltages by dividing the open probability of the particular gate by the fast or the slow deactivation time constant, respectively (34):

$$\alpha(V) = \frac{P_{\text{fast/slow}}(V)}{\tau_{\text{fast/slow}}(V)}. \quad (1)$$

Fast and slow gating was then further analyzed using the gating scheme developed by Chen and Miller for the fast gate of CIC-0 (24):



Opening rate constants measured at a given potential were plotted versus $[\text{Cl}^-]_o$ (Fig. 4) and fit with

$$\alpha(V, [\text{Cl}^-]_{\text{ex}}) = \frac{\alpha_1(V) + [\alpha_1(V) + \gamma(V)][\text{Cl}^-]_{\text{ex}}/K_c}{1 + [\text{Cl}^-]_{\text{ex}}/K_c}. \quad (2)$$

These fits provide the opening rate constants in the absence of external Cl^- (α_1), the rate constants describing a conformational change of the Cl^- -liganded channel that permits rapid channel opening (γ), and the Cl^- dissociation constant of the closed channel (K_c) (24). α_1 , γ , and K_c were determined at various voltages, and the voltage dependences of these parameters were fit with the following: $\alpha_1(V) = \alpha_1(0) e^{z_{\alpha_1}V/kT}$, $\gamma(V) = \gamma(0) e^{z_{\gamma}V/kT}$, and $K_c(V) = K_c(0) e^{z_{K_c}V/kT}$, respectively. The so-determined amplitudes and the voltage dependences of α_1 and γ were then optimized by fitting Eq. 2 to the voltage dependence of opening rate constants at different $[\text{Cl}^-]_o$ (Table 1). For these fits, the K_c s obtained from the concentration dependences of opening rate constants were used as fixed values. We tested different voltage dependences of K_c and obtained best results with voltage-independent K_c s.

To determine rate constants of MTS modification, the time course of modification was fit with a single exponential giving the time constant of modification. Pseudo-first order rate constants were calculated as the inverse of the modification time constant, and dividing by the concentration of the MTS reagents provided the second order rate constants (see Fig. 7).

TABLE 1 Gating properties of WT and truncated CIC-1 channels

	P_{open} under standard conditions $V_{0.5}$ (mV)	Fast gate						$K_{\text{C fast}}(\text{Cl}^-)$ at -35 mV (mM)
		$V_{0.5 \text{ fast}}$ under standard conditions (mV)	$P_{\text{min fast}}$ under standard conditions	$\alpha_{1 \text{ fast}}(0)$ (s^{-1})	$z (\alpha_{1 \text{ fast}})$	$\gamma_{\text{fast}}(0)$ (s^{-1})	$I (\gamma_{\text{fast}})$	
hCIC-1	-75.9 ± 1.7 $n = 79$	-90.5 ± 0.8 $n = 13$	0.18 ± 0.03 $n = 13$	0.69	-0.61	677.9	0.42	54.1 ± 3.1 $n = 9$
R894X hCIC-1	-96.2 ± 0.8 $n = 58$ $p < 0.01$	-107.8 ± 1.8 $n = 8$ $p < 0.01$	0.23 ± 0.02 $n = 8$ $p = 0.51$	0.42	-0.55	1088.7	0.50	33.0 ± 1.5 $n = 10$ $p < 0.01$
K875X hCIC-1	-21.4 ± 1.1 $n = 71$ $p < 0.01$	-59.0 ± 1.5 $n = 13$ $p < 0.01$	0.27 ± 0.05 $n = 13$ $p = 0.25$	0.52	-0.47	258.4	0.33	155.0 ± 20.5 $n = 5$ $p < 0.01$
		Slow gate						$K_{\text{C slow}}(\text{Cl}^-)$ at -35 mV (mM)
		$V_{0.5 \text{ slow}}$ under standard conditions (mV)	$P_{\text{min slow}}$ under standard conditions	$\alpha_{1 \text{ slow}}(0)$ (s^{-1})	$z (\alpha_{1 \text{ slow}})$	$\gamma_{\text{slow}}(0)$ (s^{-1})	$z (\gamma_{\text{slow}})$	
hCIC-1		-74.7 ± 2.2 $n = 14$	0.54 ± 0.02 $n = 14$	0.46	-0.61	16.43	0.54	16.7 ± 0.8 $n = 6$
R894X hCIC-1		-98.4 ± 2.0 $n = 8$ $p < 0.01$	0.77 ± 0.08 $n = 8$ $p < 0.01$	n.d.	n.d.	n.d.	n.d.	n.d.
K875X hCIC-1		-25.5 ± 1.8 $n = 13$ $p < 0.01$	0.23 ± 0.01 $n = 13$ $p < 0.01$	0.55	-0.60	13.08	0.51	45.3 ± 6.7 $n = 7$ $P < 0.01$

n.d.: not determined.

All data are shown as mean \pm SE from at least three cells and at least two distinct transfections. In general, mean \pm SE are small and, in the figures, often smaller than the symbols. For statistical evaluation, Student's *t*-test was applied.

RESULTS

Functional expression of WT and C-terminally truncated hCIC-1 channels

Fig. 1 shows representative current recordings from tsA201 cells expressing WT (Fig. 1 *B*), R894X (Fig. 1 *C*), and K875X (Fig. 1 *D*) hCIC-1 channels. Cells were held at 0 mV, and voltage steps between -155 mV and $+125$ mV were applied, each followed by a fixed test step to -125 mV. WT and mutant currents increased instantaneously upon voltage steps in the positive as well as in the negative direction. At positive potentials, the current amplitudes were time independent; upon membrane hyperpolarization the instantaneous rise was followed by a slower decrease of the current amplitude due to channel deactivation. WT and mutant channels displayed comparable macroscopic current amplitudes but differed in the time course and degree of deactivation (Fig. 1).

To determine the voltage dependence of the open probability for WT and mutant channels, instantaneous current amplitudes were measured at a voltage step to -125 mV

after prepulses to different voltages, plotted versus the preceding potential and normalized to the open probability at 0 mV obtained by nonstationary noise analysis (Fig. 2 *A*). Both C-terminal truncations change the voltage dependence of activation but leave the maximum absolute open probability unchanged (Fig. 2 *A*). The activation curve of R894X hCIC-1 channels was shifted to more negative potentials and the voltage dependence of K875X hCIC-1 to more positive potentials compared to WT hCIC-1 (Table 1).

CIC-1 channels are known to exhibit two different gating processes. Fast gating opens and closes individual protopores, whereas slow cooperative gating steps act on both protopores together (35). Time constants of fast and slow deactivation were determined through fitting a sum of two exponentials and a time-independent value to current relaxations at negative potentials. Both deactivation time constants are voltage dependent, and carboxy-terminal truncations cause a shift of these voltage dependences (Fig. 2 *B*). To separate the open probabilities of the fast and the slow gate, we used the method developed by Accardi and Pusch (33). A short pulse to $+180$ mV, inserted before the test step into the pulse protocol given in Fig. 1 (Fig. 2 *C*, *inset*), fully activates the fast gate and permits acquisition of the relative open probability of the slow gate (Fig. 2 *C*). Dividing the channel open probability by the relative slow gate activation curve yields the open probability of the fast gate (Fig. 2 *D*). Since

maximum absolute open probabilities are close to 1 for all tested conditions, absolute open probabilities are equal to relative open probabilities in all cases. WT and mutant channels differ in the voltage dependence of the fast gate and in the voltage dependence and minimum open probability of the slow gate (Fig. 2, *C* and *D*, Table 1).

Gating was investigated for various external chloride concentrations, and under all conditions tested, a linear relationship between midpoint of activation and the logarithm of the $[Cl^-]_o$ was observed (Fig. 2, *E* and *F*). This is in agreement with the notion that the voltage dependence of fast and slow gating is conferred entirely by the permeant anion and further supports the notion that the stimulus for CIC-1 activation is a change of the electrochemical gradient for Cl^- , rather than one of the transmembrane voltage (23).

We next determined opening rate constants for both gates from open probabilities and relaxation time constants under various $[Cl^-]_o$ and voltages (Fig. 3). Such an analysis is restricted to voltages, where relaxation time constants can be reliably determined, i.e., only in a voltage range between -155 mV and -25 mV for the fast gate and -115 mV and -25 mV for the slow gate. For the slow gate of R894X hCIC-1 channels, relative current components and voltage-dependent changes of absolute open probabilities are very small, precluding accurate determination of the open proba-

bility and the deactivation time constants for several $[Cl^-]_o$. We therefore did not calculate slow opening rate constants for this mutant.

Augmenting the $[Cl^-]_o$ increases fast and slow opening rate constants at more positive voltages, leaving rate constants at very negative voltages unaffected. Opening rate constants at different $[Cl^-]_o$ converge at negative voltages but do not cross over. All these results demonstrate that fast and slow gating of CIC-1 can be adequately described by the gating scheme developed by Chen and Miller (24) for the *Torpedo* CIC isoform, CIC-0.

Within this gating scheme, opening rate constants are a function of three different parameters: the dissociation constant of anion binding sites accessible in the closed state (K_C), the opening rate constant in the absence of external Cl^- (α_1), and the rate constant (γ) describing a conformational change of the Cl^- -liganded channel that permits rapid channel opening (24). These parameters were determined for fast and slow gating of WT and truncated CIC-1 channels in two steps. We first plotted opening rate constants of the fast and the slow gate versus $[Cl^-]_o$ and fitted the dependences with Eq. 2 (Fig. 4). These fits provide values for K_C in a voltage range of 30 mV (Fig. 5, *A* and *B*). α_1 and γ exhibit a significant voltage dependence that cannot be reliably determined from these fits because of the limited voltage range. We therefore computed these values by fitting the voltage dependence of fast and slow opening rate constants for different $[Cl^-]_o$ (Figs. 3 and 5, Table 1).

For all constructs, we obtained best fits with voltage-independent K_C s, similar to the voltage-independent anion binding observed in CIC-0 (24). K_C s associated with fast gating were larger than the corresponding values of slow gating, whereas values of γ were larger for fast than for slow gating (Fig. 5, *C* and *D*, Table 1). The α_1 rate constants are similar for the fast and the slow gate. They increase with membrane hyperpolarization, whereas the voltage dependence of γ is opposite (Table 1). The voltage dependence of α_{1fast} and γ_{fast} differs from fast gating of the *Torpedo* isoform (Table 1). γ_{fast} is less voltage dependent, whereas α_{1fast} exhibits a steeper voltage dependence than the corresponding values of CIC-0.

Our results demonstrate that fast and slow activation gating of CIC-1 fits to a model in which anions bind voltage independently and their subsequent translocation to the electric field fully confers the voltage dependence of channel opening. The truncation-induced differences in hCIC-1 gating (Fig. 1) are caused by altered anion binding to the closed states. K875X hCIC-1 channels exhibit the largest K_C s, whereas those for WT and R894X hCIC-1 channels are significantly smaller ($K_C(Cl^-)$, Table 1). Both mutant channels also exhibit altered γ_{fast} rate constants. These results demonstrate that carboxy-terminal truncations modify the apparent affinity for anions and, in the case of fast gating, anion translocation. These variations result in altered voltage dependences of fast and slow gating.

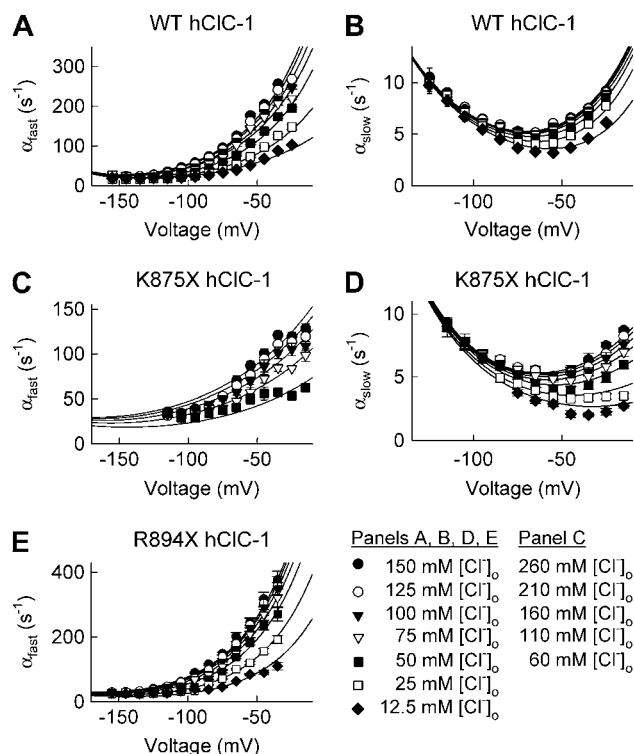


FIGURE 3 Voltage and chloride dependence of opening rate constants for the fast (*A*, *C*, and *E*) and the slow (*B* and *D*) gate of WT and truncated hCIC-1. Solid lines represent fits to Eq. 2. Mean \pm SE ($n > 5$).

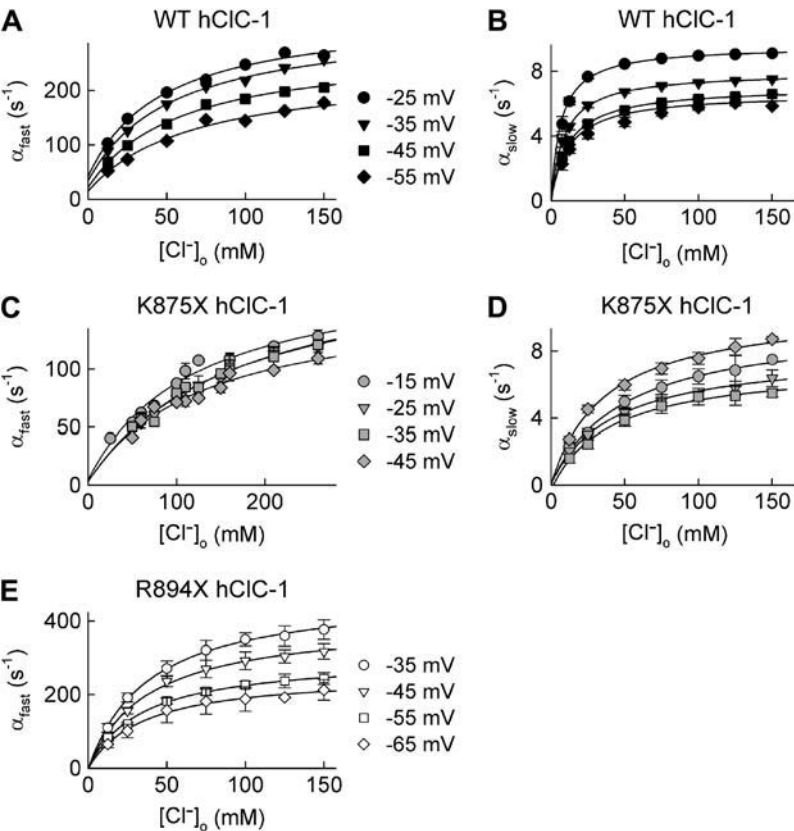


FIGURE 4 Opening rate constants for the fast and the slow gate of WT and truncated hClC-1. (A, C, and E) Dependence of the opening rate constants of the fast gate of WT (A), K875X (C), and R894X (E) hClC-1 channels on $[Cl^-]_o$. (B and D) Dependence of the opening rate constants of the slow gate of WT (B) and K875X (D) channels on $[Cl^-]_o$. Solid lines represent fits to Eq. 2. Mean \pm SE ($n > 5$).

Truncation mutations exhibit altered anion binding to the open channel

We next studied anion binding to WT and mutant chloride channels in the open state. Various permeant anions, such as I^- and NO_3^- , block chloride current through hClC-1 by binding to sites within the ion conduction pathway with higher affinity than Cl^- (27,36,37). The block of chloride

currents by anions applied to the internal or the external membrane side allows one to distinguish between two anion binding sites of the open channel, one accessible from the inside and one from the outside. The two anion binding sites differ in their apparent binding affinity and in their effect on gating properties (27,36,37). Fig. 6 A shows the concentration dependence of external I^- block measured at -115 mV for WT and mutant channels. In all cases a Michaelis-

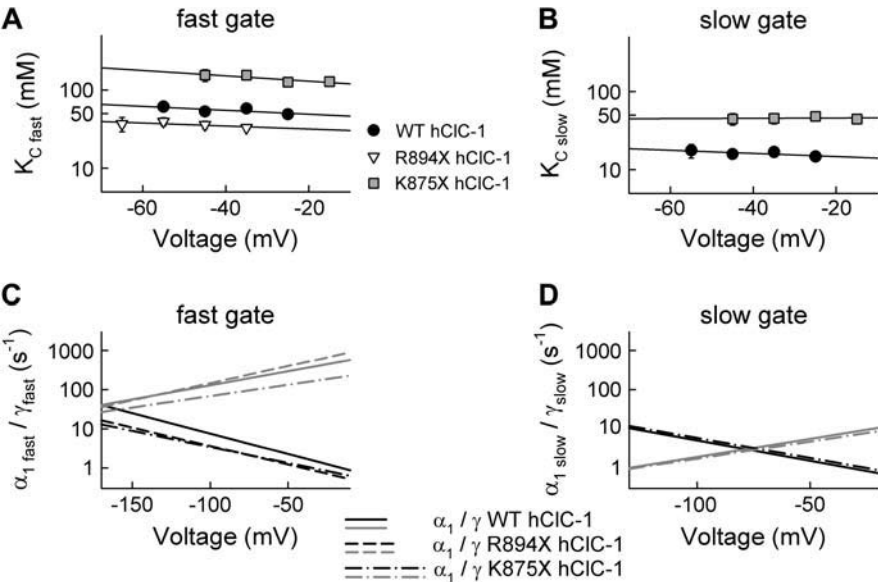


FIGURE 5 Alteration of gating parameters by carboxy-terminal truncations. (A and B) Voltage dependence for the chloride dissociation constants (K_{CS}) of the gating binding site for the fast (A) and the slow (B) gate. Mean \pm SE ($n > 5$). (C and D) Voltage dependence of α_1 and γ for the fast (C) and the slow (D) gate obtained from fitting the voltage dependence of the opening rate constants with Eq. 2.

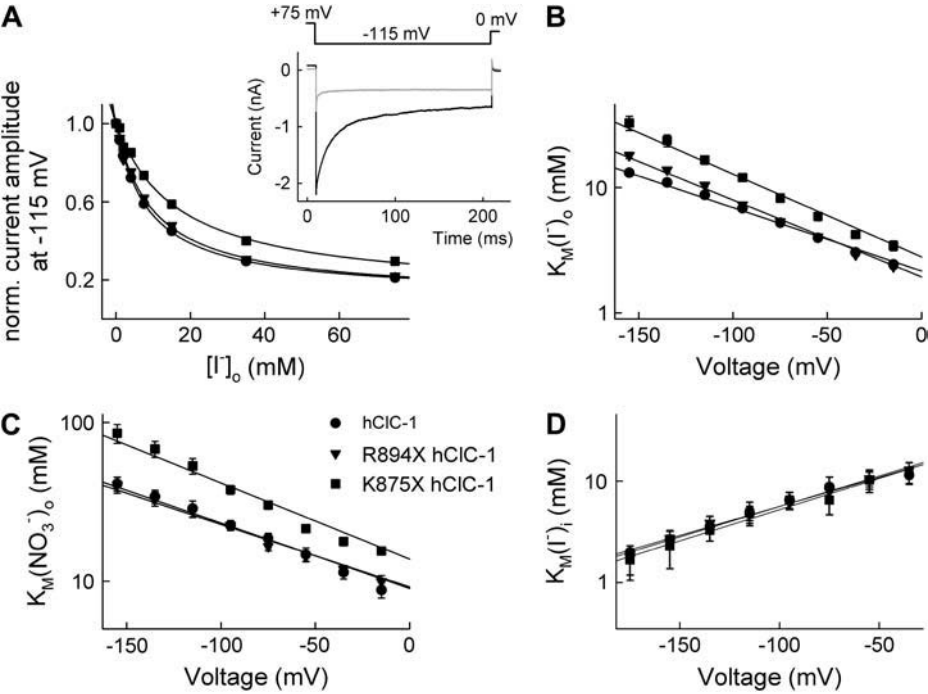


FIGURE 6 Anion current block by external and internal anions. (A) Concentration dependence of current block by external I^- at -115 mV for WT (●), R894X (▼), and K875X (■) hCIC-1 channels. The solid lines represent a fit to the Michaelis-Menten function. (Inset) Representative whole-cell recordings from a cell expressing WT hCIC-1 channels at 0 (black line) and 20 mM external I^- (shaded line). (B–D) Voltage dependence of the K_M s determined by fits shown in (A) for external I^- (B), external NO_3^- (C), and internal I^- (D).

Menten relationship with a Hill coefficient of 1 adequately described the data, providing the Michaelis constants (K_M s) for WT and mutant channels. WT, R894X, and K875X hCIC-1 channels display different K_M s for external I^- (Fig. 6 B, Table 2), and K875X differ from WT and R894X hCIC-1 in block by external NO_3^- (Fig. 6 C, Table 2). In contrast, no differences were observed for internal I^- (Fig. 6 D, Table 2).

The unitary current amplitude (single channel current at -160 mV determined by nonstationary noise analysis (Table 2)) as well as the selectivity among anions (anion permeability ratios determined from reversal potential measurements (32): $P_{SCN} \sim P_{Cl} > P_{Br} > P_{NO_3} > P_I > P_F$) remained unaffected in mutant channels. The observed functional alterations, i.e., the changed apparent affinities for anion binding to the closed state and to the external binding site of the open channel, together with the unaltered selectivity, unitary conductance, and block by internal anions, sug-

gest that the outer vestibule of the ion conduction pathway is altered in truncated hCIC-1 channels.

Accessibility of a substituted cysteine within the CIC channel pore

Using site-directed mutagenesis, a conserved fixed charge close to the CIC channel selectivity filter was identified as playing a role in defining the anion binding affinity for hCIC-1 (32). Neutralizing K231 (located in the N-terminus of the F helix in the atomic structure of a prokaryotic CIC isoform (13)) by substitution of a cysteine causes an inversion of the $Cl > I$ permeability of WT hCIC-1, and adding the positive charge back to C231 by reacting a MTSEA reagent restores WT anion selectivity (32). These results suggest that the positively charged side chain at position 231 is a major determinant of the electrostatic potential of the outer

TABLE 2 Pore properties of WT and truncated CIC-1 channels

	i at -160 mV (pA)	$K_M(I^-)_o$ at -115 mV (mM)	$K_M(NO_3^-)_o$ at -115 mV (mM)	$K_M(I^-)_i$ at -115 mV (mM)	MTSET modification of K231C ($s^{-1}M^{-1}$)
hCIC-1	0.28 ± 0.01 $n = 11$	8.7 ± 0.4 $n = 6$	28.6 ± 3.3 $n = 10$	4.8 ± 0.7 $n = 6$	1692 ± 126 $n = 42$
R894X hCIC-1	0.24 ± 0.01 $n = 8$ $p = 0.38$	10.4 ± 0.4 $n = 7$ $p < 0.01$	27.7 ± 2.5 $n = 8$ $p = 0.83$	4.8 ± 0.9 $n = 5$ $p = 0.91$	2227 ± 90 $n = 43$ $p < 0.01$
K875X hCIC-1	0.29 ± 0.02 $n = 7$ $p = 0.17$	16.5 ± 1.3 $n = 8$ $p < 0.01$	53.2 ± 6.0 $n = 8$ $p < 0.05$	4.9 ± 1.3 $n = 5$ $p = 0.62$	1088 ± 57 $n = 42$ $p < 0.01$

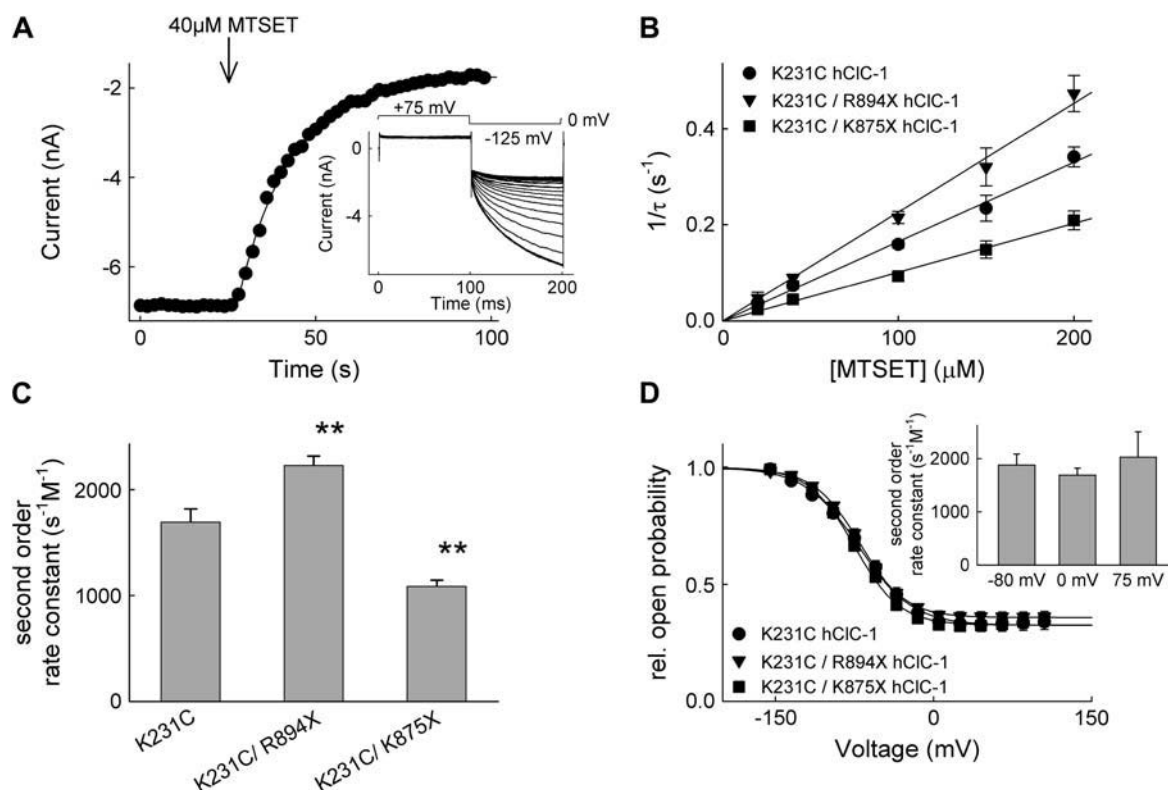


FIGURE 7 Truncation mutants alter cysteine accessibility of K231C. (A) Time-dependent reduction of K231C/K875X hCIC-1 channels after application of 40 μM MTSET. (Inset) Current recordings during modification by MTSET. The solid line represents the fit to a single exponential. (B) Concentration dependence of inverse modification time constants for various [MTSET] concentrations at 125 mV for K231C (●), K231C/R894X (▼), and K231C/K875X (■) hCIC-1 channels. Data were fit with linear regressions. (C) MTSET second order modification rates for K231C, K231C/R894X, and K231C/K875X hCIC-1 channels. Mean \pm SE ($n > 8$). (D) Voltage dependence of the relative open probability of K231C (●), K231C/R894X (▼), and K231C/K875X (■) channels. Solid lines represent fits to the Boltzmann equation (K231C: $V_{0.5} = -70.7 \pm 2.6$ mV ($n = 10$), K231C/R894X: $V_{0.5} = -69.8 \pm 0.9$ mV ($n = 14$, $p > 0.05$); K231C/K875X: $V_{0.5} = -75.4 \pm 2.8$ mV ($n = 9$, $p > 0.05$). (Inset) voltage dependence of MTSET second order modification rates for K231C hCIC-1. Mean \pm SE ($n > 7$).

vestibule. We studied whether an altered relative position of this positively charged side chain is responsible for the observed differences in anion binding. MTS modification rate constants were measured for K231C, K231C/R894X, and K231C/K875X hCIC-1 channels (Fig. 7). hCIC-1 channels without substituted cysteines exhibit no functional change upon exposure to MTS reagents (32). Fig. 7 A (inset) shows current recordings from a cell expressing K231C/K875X hCIC-1 during modification by 40 μM MTSET. Fig. 7 A gives the time course of isochronal current amplitudes determined at the end of a voltage step to -125 mV. Current amplitudes decayed in a monoexponential time course (Fig. 7 A). Time constants of current amplitude decay were determined for various [MTSET], and for all constructs a linear relationship between the inverse time constants and [MTSET] was observed (Fig. 7 B). This demonstrates that the obtained time course does indeed report modification of an engineered cysteine and not merely channel closure by a slow gating process. The different slopes were used to calculate second order rate constants of modification (Fig. 7 C). Constructs with truncated C-termini exhibit significantly different mod-

ification rate constants: K231C/R894X hCIC-1 is modified with a higher, K231C/K875X hCIC-1 with a lower second order rate constant (Fig. 7, A and C, Table 2).

The reaction of an MTS reagent with a thiol side chain of a channel protein occurs in two consecutive steps. The MTS reagent first has to diffuse from the aqueous solution to the side chain and then reacts with it in a second step. The experimentally determined reaction rates give information about the rate-limiting step in the cascade. All measured rate constants were well below the values observed for the modification of thiols in aqueous solution (38). This observation, together with the linear relationship between [MTS] and pseudo first order reaction constants, indicates that reagent access from the aqueous solution is the rate-limiting step. The voltage dependences of the relative open probability of full length and truncated K231C channels virtually superimpose (Fig. 7 D), and modification rates of full length K231C are voltage independent between -80 mV and +75 mV (Fig. 7 D, inset). These data indicate that the different modification rates are not due to an altered gating but rather to different arrangements of the access pathway to the thiol side chain. We

conclude that the differences in rate constants between WT, R894X, and K875X channels demonstrate altered outer pore vestibules in the three constructs.

DISCUSSION

We investigated the functional effects of two carboxy-terminal truncations, R894X and K875X, of the human muscle chloride channel, hCIC-1. Truncated hCIC-1 channels exhibit unaltered unitary current amplitudes and relative anion permeabilities for SCN^- , NO_3^- , and I^- . They differ, however, in anion binding to open and closed channels demonstrating a role of the carboxy-terminus in defining pore properties.

It is well established that CIC channels are gated by the permeant anion (22–24), in marked contrast to voltage-gated cation channels. However, one report appeared to be incompatible with this concept. A point mutation in the distal carboxy-terminus (A885P) shifts the activation curve of muscle chloride channels to more positive values and causes muscle hyperexcitability and muscle stiffness in an animal model of myotonia, the myotonic goat (21). Since the distal carboxy-terminus does not contribute to the formation of the ion conduction pathway, the gating alteration by A885P could have been interpreted as indicative for inherently voltage-dependent gating transitions. Our results remove this remaining doubt. The distal carboxy-terminus modifies anion binding to the closed pore, thus explaining the changed voltage dependence by the A885P substitution within this region. Moreover, testing fast and slow gating of WT and mutant channels over a broad range of external anion concentrations, no deviation from the expectations of anion-dependent gating could be seen (Fig. 2).

Surprisingly, the truncations cause similar alterations of fast and slow gating of hCIC-1 channels. So far, these two gating transitions have been conceived as being two independent and structurally unrelated processes, with the slow gate as a structure closing both protopores together. Our experiments now demonstrate that both gating processes can be adequately described by the Chen-Miller model (24). In this model, the permeant anion binds voltage independently to the closed channel and subsequently translocates through the electric field, conferring the majority of the voltage dependence of channel opening. These findings suggest that opening of the fast as well as of the slow common gate requires anion translocation through an individual protopore. The K_{CS} associated with fast gating are larger than those obtained for slow gating. This implies that hCIC-1 anion conduction pathways exist in two closed states of differing apparent anion affinity, one with lower affinity corresponding to fast gating and one with higher affinity responsible for slow gating. Whereas the two gates dramatically differ in the translocation of the anion through the electric field, opening rate constants in the absence of

external Cl^- are quite similar (Table 1). A simple way of explaining our results is by proposing that fast and slow gating are resting and activated states of an allosteric gating process. A protopore might exist in two conformations, in one anion binding and translocation causes a slow co-operative opening of both of them, and in the other conformation fast opening of one protopore. Carboxy-terminal truncations modify anion binding in both conformations and thus alter fast and slow gating.

How does the distal carboxy-terminus modify anion binding to the pore of hCIC-1? The cytoplasmic channel regions might extend the ion conduction pathway by forming a cytoplasmic pore, as observed in inwardly rectifying potassium channels (39,40) or, alternatively, affect the conformation of the ion conduction pathway. At present, no structural information about the CIC carboxy-terminus is available, but several lines of evidence support the idea of a changed conformation of the pore of truncated hCIC-1 channels. Carboxy-terminal deletions have a pronounced effect on anion binding from the external side but no measurable effect on internal anion binding (Fig. 6). Moreover, anion binding is affected in the open (Fig. 6) as well as in the closed (Fig. 5) state. Lastly, MTS modification of an engineered cysteine demonstrates that there are discrete conformational differences between WT and truncated hCIC-1 channels (Fig. 7).

There are several earlier reports that carboxy-terminal modifications change gating of CIC channels. Fong et al. (41) reported that carboxy-terminal truncations and chimeras affect slow gating of CIC-0. A disease-causing point mutation (G715E) in human CIC-2 alters the anion dependence of gating (3), and splice variants of a *Caenorhabditis elegans* homolog, Clh-3, with modified N- and C-terminus exhibit altered voltage-dependent gating (42). Our findings provide a possible explanation for all these findings. The carboxy-terminus likely determines anion binding and thus anion-dependent gating of various CIC isoforms.

The carboxy-terminus does not only modify functional properties, but is also required for proper targeting of several CIC isoforms into the respective cell compartment (18–20,43,44). The functional effect of the carboxy-terminus is not restricted to the time period immediately after translation in the endoplasmic reticulum and therefore is not involved in correct folding of the hydrophobic core domain. Maduke et al. (18) demonstrated that injection of a purified protein containing the last 154 amino acids of CIC-0 fully restores the function of truncated CIC-0 channels when injected into *Xenopus* oocytes, even under conditions abolishing protein synthesis. This demonstrates that the carboxy-terminus interacts with other protein domains and that modification of the carboxy-terminal domain by intracellular signals permits adjustment of CIC channel pore conformations. The physiological relevance of such a regulation of pore properties arises from the unique gating mechanisms of CIC channels. In voltage-gated cation channels, an endogenous

voltage sensor moves across the membrane, and this movement can be modified by electrostatic or other interactions with regulatory pathways inside the cell, resulting in altered channel gating (45,46). Since CIC channels are gated by the permeant anion, changes of the voltage dependence of channel gating require altered anion binding to the closed pore. The functional interaction of the carboxy-terminus with the pore of hCIC-1 might make voltage-dependent gating processes accessible to intracellular signal cascades, permitting modification of the macroscopic CIC anion conductance by regulatory pathways.

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