Probing the Pore of ClC-0 by Substituted Cysteine Accessibility Method Using Methane Thiosulfonate Reagents

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ABSTRACT ClC channels are a family of protein molecules containing two ion-permeation pores. Although these transmembrane proteins are important for a variety of physiological functions, their molecular operations are only superficially understood. High-resolution X-ray crystallography techniques have recently revealed the structures of two bacterial ClC channels, but whether vertebrate ClC channel pores are similar to those of bacterial homologues is not clear. To study the pore architecture of the *Torpedo* ClC-0 channel, we employed the substituted-cysteine-accessibility method (SCAM) and used charged methane thiosulfonate (MTS) compounds to modify the introduced cysteine. Several conclusions were derived from this approach. First, the MTS modification pattern from Y512C to E526C in ClC-0, which corresponds to residues forming helix R in bacterial ClC channels, is indeed consistent with the suggested helical structure. Second, the ClC-0 pore is more accessible to the negatively charged than to the positively charged MTS compound, a pore property that is regulated by the intrinsic electrostatic potential in the pore. Finally, attempts to modify the introduced cysteine at positions intracellular to the selectivity filter did not result in larger MTS modification rates for the open-state channel, suggesting that the fast gate of ClC-0 cannot be located at a position intracellular to the Cl⁻ selectivity filter. Thus, the proposal that the glutamate side chain is the fast gate of the channel is applicable to ClC-0, revealing a structural and functional conservation of ClC channels between bacterial and vertebrate species.

KEY WORDS: MTS modification • SCAM • charge selection • state dependence

INTRODUCTION

The ClC channel family comprises members widely distributed in different living species from bacteria to vertebrate animals (Jentsch et al., 1999, 2002; Maduke et al. 2000; Hille, 2001). These channels are unique in that two identical ion-conducting pores are present in a channel molecule. Recently solved high-resolution X-ray structures of the bacterial ClC channels have provided a basis for understanding the molecular operations of these anion channels (Dutzler et al., 2002, 2003). The structures show that ClC channels are homodimers containing two pores, and each pore is formed entirely by a subunit containing 18 α-helices named helix A to R. These helices wrap around a common center, and the amino acids at the NH₂-terminal end of the α-helices are brought together to coordinate the bound chloride (Cl⁻) ion at a site (the central site, Scen) likely to serve as the ion selectivity filter (Dutzler et al., 2002). Besides the Scen site, there are two other Cl⁻ binding sites found in the ion permeation pathway: one internal to the Scen site, and the other located external to Scen at a position of the negative charge on a glutamate residue (Dutzler et al., 2003). This glutamate residue, which corresponds to glutamate 166 (E166) of ClC-0, projects its negatively charged side chain into the ion permeation pathway to occlude the ion flux. Competition of Cl⁻ with this glutamate side chain has been proposed to be responsible for the coupling of gating to the ion permeation (Dutzler et al., 2002, 2003). Although not as clear as the pore in the KcsA K⁺ channel (Doyle et al., 1998), the bacterial ClC channel structures also suggest the Cl⁻ permeation pathway. In particular, basic amino acid residues in the internal and external pore entrances were thought to be important in funneling the permeant ions to the pore (Dutzler et al., 2002).

Although these interesting structural features in the bacterial ClC channels are very informative, the difficulty in recording the bacterial ClC channels with conventional electrophysiological techniques has impeded the studies of the structure-function relation in bacterial ClC channels. ClC-0 from the electric organ of *Torpedo* rays (Jentsch et al., 1990; O’Neill et al., 1991) provides a model system to understand the channel functions. Significant characterization of ClC-0 properties has been obtained from functional recordings of this...
Therefore, whether the 20% sequence-identity employed substituted cysteine–accessibility method (SCAM)* between ClC-0 and bacterial ClC channels, we employed bacterial ClC channel structures to define the pore residues of the ClC-0 channel.

A direct exploration of the ClC-0 pore will thus complement bacterial ClC channel structures to provide a structural basis for identifying crucial charged residues in the pore (Dutzler et al., 2000). The crystal structures of bacterial ClC channels, however, have suggested only one gate, the side-chain of a glutamate residue. One might be curious which charged side chain of the glutamate to open the pore is used to explain the previous functional observation that the gating of CIC-0 is tightly coupled to ion permeation (Pusch et al., 1995; Chen and Miller, 1996). Finally, a positively charged lysine residue at the inner pore mouth of ClC-0 has been demonstrated to control the pore conductance (Middleton et al., 1996; Chen and Chen, 2003). These channel properties derived from functional studies of CIC-0 appear to fit the structures of the bacterial CIC channels quite nicely.

Although the above properties of CIC-0 appear to find support from the bacterial CIC channel structures, other aspects of the channel remain uncertain. First, the Torpedo ClC-0 channel is known to have two distinct gating mechanisms, a fast and a slow (inactivation) gating, each operating at very different time scales (for review see Miller and Richard, 1990; Maduke et al., 2000). The crystal structures of bacterial CIC channels, however, have suggested only one gate, the side-chain of a glutamate residue. One might be curious which gating mechanism, the fast or the slow gating of CIC-0, corresponds to the glutamate gate observed in bacterial CIC channel structures. Even though bacterial CIC channel structures show a glutamate side chain in each pore, suggesting that this side chain is more likely to be the fast gate, the absence of the structure for the COOH-terminal half of the CIC-0 molecule makes this assertion less certain. Second, the bacterial CIC channel structures provide a structural basis for identifying the pore-lining residues in the ion-permeation pathways. In comparison with the structure of KcsA K+ channel, however, the two ion permeation pathways in bacterial CIC channels are more obscure. For example, the bound Cl− ions at the selectivity filter are nearly invisible from the pore entrances. The curved ion permeation pathway, as defined by the bound ions and the crucial charged residues in the pore (Dutzler et al., 2002, 2003), is void of water filled space as the one observed in the KcsA K+ channel (Doyle et al., 1998). Therefore, whether the 20% sequence-identity Torpedo homologue adopts a similar pore structure is unknown. A direct exploration of the ClC-0 pore will thus complement bacterial CIC channel structures to define the pore residues of the ClC-0 channel.

To understand if the pore architecture is conserved between ClC-0 and bacterial CIC channels, we employed substituted cysteine–accessibility method (SCAM)* and used methane thiosulfonate (MTS) reagents to probe the putative inner pore region of CIC-0. SCAM has been widely employed in a variety of cation and anion channels (Akabas et al., 1992; Lu and Miller, 1995; Cheung and Akabas, 1997; Fahlke et al., 1997; Liu et al., 1997; Karlin and Akabas, 1998; Lu et al., 1999; Reeves et al., 2001). By comparing the reaction rates of the negatively and positively charged MTS reagents with the cysteine in the pore, it is possible to measure the extent of anion to cation selectivity in the pore (Cheung and Akabas, 1997; Pascual and Karlin, 1998; Wilson et al., 2000). Studying the accessibility of the introduced cysteine to MTS reagents at open and closed states would also reveal potential state-dependent modifications, which may be argued to produce the presence of a gate (Liu et al., 1997). In the present study, we introduce cysteine at the positions Y512 to E526 (see also Ludewig et al., 1997), residues that potentially form an α helix (helix R, according to the bacterial CIC structures) extending from the S_{cen} site to the proposed inner pore mouth. We also place a cysteine in place of S123. The side chain of the corresponding serine of the bacterial CIC channel coordinates the bound Cl− at S_{cen}. Comparisons of the modification rates of MTS reagents with the introduced cysteine at these positions reveal that a more positive intrinsic electrostatic potential is encountered by the MTS reagents as the introduced cysteine is located deeper in the pore. Altering the side-chain charge of the amino acid residues in the pore also significantly changes the MTS modification rate, suggesting that the charges of the pore residues participate in selecting anionic over cationic MTS reagents. We have also conducted experiments to examine if there is a state-dependent modification by MTS reagents of a cysteine placed at various positions in the pore.

**Materials and Methods**

**Mutagenesis, Channel Expression, and Transfection**

Site-directed mutagenesis was performed using PCR-based techniques, and the mutations were confirmed by DNA sequencing. All the cysteine mutants in the present study are created on the background of the C212S mutation (Lin et al., 1999), and therefore this inactivation-suppressed mutant will be referred to as the wild-type (WT) channel. For channel expressions in *Xenopus* oocyte, all the mutants were constructed in the pBluescript vector (Stratagene). The method for RNA synthesis and the injection of RNA into oocytes have been described in previous studies (Chen, 1998; Lin et al., 1999; Lin and Chen, 2000; Chen and Chen, 2003). For channel expression in human embryonic kidney (HEK) 293 cells, mutant plasmids in pBluescript vector were digested with two restriction enzymes, KpnI and AgeI (New England Biolabs, Inc.). The cDNA fragments containing the mutations were sub-cloned into a WT ClC-0 constructed in the pcDNA3 vector.

HEK 293 cells were plated into a 35-mm culture dish (Corning, Inc.) at a density of $3 \times 10^5$ per well one day before transfection.
tion. The medium was the antibiotics-free Dulbecco’s Modified Eagle Medium supplemented with 20 mM L-glutamine and 10% fetal bovine serum. Upon transfection, cells usually grew to >90% confluence, and 1–3 μg of the channel cDNA and 0.4–0.7 μg of green fluorescence protein (GFP) cDNA were co-transfected into the cells with Lipofectamine™ 2000 (GIBCO BRL) according to the procedures from the manufacturer. On the day when the recordings were performed (usually 24–48 h after transfection) the cells were dissociated into individual cells with 1 × EDTA/Trypsin (GIBCO BRL) and plated onto coverslips coated with 0.01% poly-L-lysine (Sigma Aldrich). The cells were allowed to settle down over the coverslip for two hours, and identification of the transfected cells was facilitated through GFP protein using a fluorescence microscope (Leica DM IRB).

Electrophysiological Recordings

The excised inside-out patch configuration (Hamill et al., 1981) was used throughout all recordings. Borosilicate glass electrodes were pulled by PP-830 puller (Narashige Co.), and when filled with the pipette solution, have resistance of ~1–2 MΩ. For recordings of HEK 293 cells, the standard bath (intracellular) solution consisted of (in mM): 130 NaCl, 5 HEPES, 5 MgCl₂, 1 EGTA, pH = 7.4. The pipette (extracellular) solution contained (in mM): 130 NMDG-Cl, 5 HEPES, 5 MgCl₂, 1 CaCl₂, pH = 7.4. In the experiments using an extracellular solution containing only 4 mM Cl⁻, the 130 mM NMDG-Cl in the pipette solution was replaced with 130 mM NMDG-glutamate and 5 mM MgCl₂ was changed to 1 mM. For single-channel recordings, the bath and the pipette solutions were described previously (Lin and Chen, 2000; Chen and Chen, 2003). The pH of the bath and the pipette solution was titrated with NaOH and NMDG, respectively.

Macroscopic currents recorded from HEK 293 cells were low-pass filtered at 1kHz (digital filter associated with the acquisition software) and were digitized by Digidata 1320 acquisition board (Axon Instruments, Inc.) at 5 kHz with pClamp8 software. Out of the 16 mutants, only L521C did not produce large enough current for functional studies. To evaluate the mutation effects of introducing a cysteine residue to various positions, a series of voltage steps was delivered to the membrane patch to elicit a family of macroscopic currents that allows a calculation of the P_o of the channel. Examples from the WT channel and several cysteine mutants are displayed in Fig. 1. Such a voltage protocol contains a prepulse voltage step to 60 or 80 mV for 100 ms, followed by a 0-mV voltage step. The initial current was measured at the steady-state. However, the modifications of K519C and S125C with MTS were monitored at −40 mV because the current increases much more at the negative potential than at the positive potential. In experiments using 4 mM Cl⁻, the current was monitored by a voltage step to 80 mV followed by a 0-mV voltage step. The initial current at 0 mV reflects the amount of the current flowing through the fully open channels at the preceding voltage step of 80 mV. These measured currents were plotted against the accumulative MTS exposure time, or the accumulative MTS exposure (concentration × time) in Figs. 8 C and 9 A.

Single-channel Recordings

To compare the MTS modification effects of the cysteine mutants between macroscopic and single-channel current levels, MTS modifications at single-channel level were also performed in some of the mutants expressed in Xenopus oocytes (see Fig. 3). Detailed procedures of single-channel recordings and analyses were described in the previous papers (Lin et al., 1999; Lin and Chen, 2000; Chen and Chen, 2001, 2003).

Data Analysis

Except where indicated, all MTS modification processes were fitted to a single-exponential equation,

\[
I_t = I_\infty + (I_0 - I_\infty) \exp(-t/\tau),
\]

where \(I_t\) is the measured current at the accumulative exposure time \(t\), \(I_0\) is the initial current, \(I_\infty\) is the current when the modification is reached to a steady-state, and \(\tau\) is the time constant of the modification process. The curve fitting was performed with an unweighted least-squares method using Clampfit 8 software (Axon Instruments, Inc.). The time constant \(\tau\) and the concentration of MTS reagents were used to calculate the second order rate constant \(k\) according to the following equation:

\[
k_{MTS} = 1/(\tau[MTS]).
\]

All data points in this study were the average of at least three independent measurements, and the numbers are presented as mean ± SEM.

RESULTS

Mutating the amino acid to cysteine sequentially from residue 512 to 526 of ClC-0 that align to helix R of the bacterial ClC channel gave rise to functional mutants in most of the cases. Except K519C and S125C, most of these cysteine mutants have a similar pattern in the macroscopic current as that of the WT channel (Fig. 1). For the K519C mutant, the slight outward rectification of the instantaneous current is likely due to the rectification of the single-channel current (Middleton et al., 1996), whereas the slower current deactivation is
due to a reduction in the fast-gate closing rate. However, the fast-gate $P_o$ of this mutant, like that of the WT channel, is close to unity at the most depolarized potential, allowing the construction of the steady-state $P_o$-$V$ curve. The only mutant that prevents the evaluation of the fast-gate $P_o$ from macroscopic current is S123C. The $P_o$ of this mutant does not reach to unity even at 80 mV, as judged from the prominent noise from the recording trace (also see Fig. 8 A). Direct observations of the channel behaviors at the single-channel level also indicate that the fast-gate closing rate of S123C is very fast (see below), resulting in a very fast macroscopic current deactivation that cannot be easily separated from the capacitance current.

Modifications of the WT and the cysteine mutants with MTS compounds have different functional consequences depending on the positions of the introduced cysteine and the charge of the modifying reagent. Even though there are 11 native cysteine residues in the WT channel (C212 has been removed in the "WT" channel referred in this study), applications of MTSES or MTSET to the intracellular side of the WT channel have little effect in altering the current (Fig. 2 A). On the other hand, MTSES decreases and MTSET increases the current of the K519C mutant (Fig. 2 B); the results directly reflect the modification effects on the single-channel conductance of this mutant (Middleton et al., 1996; Chen and Chen, 2003). For the other cysteine
mutants, MTS modifications can completely inhibit the current, partially alter the current, or have very little effect on the current (Fig. 3).

The overall results of MTSES and MTSET modifications of the 16 mutants examined in this study are shown in Fig. 4 A. 1 out of these 16 mutants, L521C, as indicated by the symbol “#” in Fig. 4 A, did not generate a large enough current for macroscopic current recordings (three runs of expression attempts). 2 out of those 15 functional mutants (K520C and E526C, indicated by the symbol “+” in Fig. 4 A) did not show an alteration of the current upon high concentrations (up to 1.5 mM) of MTSES and MTSET applications. At the present time, we could not differentiate if the absence of MTS modification effect in these two mutants is due to a lack of modification or because of no functional consequence after cysteine modification. For the rest of the mutants, both MTSES and MTSET can modify the introduced cysteine. In Q517C and I518C mutants, MTSET modification inhibits the current, while MTSES modification has little effect in altering the current. However, MTSES did modify both mutants because if the patch was first exposed to MTSES, subsequent MTSET modification was abolished (unpublished data). By the same argument, MTSET can modify P525C, but the modification has little functional effect. MTS modification rates in these three mutants (indicated by the symbol “★” in Fig. 4 A) therefore can be precisely measured only from one MTS reagent.

For the 13 cysteine mutants whose currents are susceptible to MTSES and/or MTSET modifications, the second order modification rate constants (VMO = −40 mV) are compared in Fig. 4 A. The modification rates of these mutants may seem to have no particular interesting pattern at a first glance. However, a careful comparison reveals that four positions stand out and show faster modifications than the rest of the mutants, P522C, K519C, I516C, and D513C. In every third position along the sequence, the MTS reagents give a faster modification rate. This might suggest that these four positions are more exposed to the aqueous solution. For the other positions, however, the introduced cysteine still reacts with both MTSES and MTSET significantly. Even for the two mutants S123C and Y512C, the modification rates at −40 mV for MTSES and MTSET are ~200–600 M−1s−1. Because these two residues are at the deepest locations in the pore, the other truly exposed pore residues should be more accessible to MTS reagents than these two residues. Thus, we set a criterion for picking up the exposed residues: their MTSES and MTSET modification rates are both larger than those of S123C and Y512C. Applying this criterion eliminates L524C and S514C from the 10 mutants that are functionally susceptible to both MTSES and MTSET modifications. In Fig. 4 B, the positions of the more exposed positions are shown in red, whereas the other positions in helix R are in yellow. It is interesting to observe that the “hot” spots are all positioned on one side of the suggested R helix, and this more exposed side appears to face the two permeant ions (see Fig. 4 B). Thus, this modification pattern is consistent with the idea that residues 512–526 of ClC-0 indeed form a helix, and suggests that these exposed residues likely line the wall of the pore.

The comparison of MTSES and MTSET modification rates in these more exposed residues reveals another interesting observation: they are quite similar to each other for most of the mutants even though MTSET is known to have a higher intrinsic reactivity with the free thiol in the bulk solution (Stauffer and Karlin, 1994). To examine this issue more closely, we calculate the ratio of the modification rates between MTSES (kMTSES) and MTSET (kMTSET) for those residues that are likely
exposed to the aqueous solution. This ratio reflects the charge selection at the position where the introduced cysteine is placed, and may be used to imply the anion to cation (or vice versa) selectivity of the pore (Cheung and Akabas, 1997; Pascual and Karlin, 1998; Wilson et al., 2000). Fig. 4 C plots the ratios of $k_{\text{MTSES}}/k_{\text{MTSET}}$ for the six mutants shown in red in Fig. 4 B, those for the mutants S123C and Y512C, and the ratio of modifying $\beta$-mercaptoethanol ($\beta$-ME) taken from the study of Stauffer and Karlin (1994). It is interesting to observe that as the position is deeper in the pore, the ratio $k_{\text{MTSES}}/k_{\text{MTSET}}$ generally becomes larger, with two exceptions: K519C and D513C—both mutations convert a charged amino acid to a cysteine. The deviation from the general trend of increasing the ratio at positions 519 and 513 was therefore ascribed to the alteration in the intrinsic electrostatic potential of the pore due to the neutralization of charged residue by the mutation. Because the inner pore residues of ClC-0 control the conductance of ClC-0 (and thus the Cl$^{-}$/H$^{+}$ flux) electrostatically (see Chen and Chen, 2003), we suspected that the same electrostatic regulations also contribute to the selection of the charged MTS reagents. To examine this possibility, we made a double mutant E127Q/K519C, and compared its MTS modification rates with those of the K519C mutant. At the same membrane potential, removing the negative charge at position 127 increases...
The MTSES modification by \(11601\)fold (Fig. 5). Unfortunately, in the presence of E127Q mutation, modifying the cysteine at position 519 with MTSET did not have an effect on the macroscopic current (unpublished data). This is because in the E127Q background, the channels with a positive and a neutral charge at position 519 have similar conductance (see Chen and Chen, 2003). Without a prominent modification effect, the MTSET modification rate cannot be precisely determined in the E127Q/K519 double mutant.

To circumvent this problem, we introduce cysteine at position 515 and alter the charge at positions 127 and/or 519. Fig. 6 A shows the experiment for MTSES modification of I515C in various background charge combinations for positions 127 and 519. The second order rate constants for MTSES and MTSET modifications and the ratios of \(k_{\text{MTSES}}/k_{\text{MTSET}}\) for these mutants are plotted in Fig. 6 B. It can be seen clearly that as the overall charge from positions 127 and 519 is made more negative, the MTSES modification rate is reduced while the MTSET modification rate is increased. When these positions are made more positive, the MTSES modification rate increases as expected, but the MTSET modification rate is paradoxically increased. However, the ratio of \(k_{\text{MTSES}}/k_{\text{MTSET}}\) increases monotonically by \(~110\)-fold when comparing the one in Q127/K519 channel with that in the E127/E519 channel. Like the regulation of the single-channel conductance, position 127 has a stronger effect in altering the absolute value of MTS modification rate. For example, removing one positive charge at position 519 (Q127/K519 versus Q127/M519) reduces the MTSES modification rate by \(~10\)-fold, while adding a negative charge onto position 127 (Q127/K519 versus E127/K519) reduces the MTSES modification rate by \(~20\)-fold. However, in terms of \(k_{\text{MTSES}}/k_{\text{MTSET}}\) ratio, altering the charge at position 127 is roughly equivalent to changing the charge at position 519.

Thus, the above results suggest that the pore architecture of ClC-0 appears to be similar to those seen in the bacterial CiC channels—the modification pattern from Y512C to E526C is consistent with a helical structure, and the charges at these positions also help select anions over cations. Therefore, helix R of ClC-0 is likely to line the wall of the pore. Because mutations at the inner pore mouth change the fast gating property of the channel (for example, K519E mutation, see Pusch et al., 1995; Chen and Chen, 2003), we ask if there is a physical gate internal to the Scen site. Modifications of the introduced cysteine have been used to suggest the presence of a physical gate if the introduced cysteine reacts with MTS reagents significantly faster when the channel is open than when the pore is closed (Liu et al., 1997). At more superficial positions, positions 523 and 519, no significant difference in the MTS modification rate was observed between conditions in which the fast gate \(P_o\) is \(~0.25\) and \(~0.97\) (unpublished data). In another experiment, the cysteine at a deeper position, I515C, was examined (Fig. 7). In this experiment, two approaches were used to dissect out the state dependence from the voltage dependence of MTS modifications. First, we took advantage of the fact that the fast-gate \(P_o\) is significantly changed by varying \([\text{Cl}^-]_o\) (Fig. 7 A), and compared the MTSES modification rate at the same voltage but at different \([\text{Cl}^-]_o\) (Fig. 7 B). Second, we studied MTSES and MTSET modifications at different voltages (Fig. 7, C and D). In both approaches, the modification rates for MTSES and MTSET in various conditions were \(~300–700\) \(M^{-1}s^{-1}\), and we did not observe a consistent increase of MTS modification rate when the \(P_o\) of the fast gate is increased. In contrast, the MTSES modification rate appears to decrease as the \(P_o\) of the fast gate increases.
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The voltage-dependent change of the fast-gate \( P_o \) is most prominent in the mutant S123C. Fig. 8 A shows the macroscopic current, and Fig. 8 B shows the single-channel recording traces of this cysteine mutant. Because the closing rate of the channel is very fast, we use an internal Cl\(^{-}\) concentration of 2,400 mM to slow down the closing rate of the channel in single-channel recording experiments (Chen and Miller, 1996). However, even under such a high internal Cl\(^{-}\) concentration, the closing rate of the channel is so fast that the deactivation of the macroscopic current cannot be easily separated from that of the capacitance current. Even at a voltage of 80 mV (the most depolarized voltage in Fig. 8 A), the large noise from the recording trace suggests that the fast-gate \( P_o \) is not close to unity. Therefore, the severe outward rectification in the macroscopic current recording is most likely due to a significant increase of \( P_o \) when the voltage is depolarized. We therefore compare the MTSES modification rates at four voltages (Fig. 8, C and D). Again, the modifications do not show a consistent pattern of higher modification rate when the \( P_o \) of the channel is raised.

Finally, we studied the modification of the cysteine introduced at another position at S cen, Y512. Fig. 9 A shows the comparison of MTSES and MTSET modifications at two voltages where the values of fast-gate \( P_o \) are very different. It can be clearly seen that when the fast-gate \( P_o \) is raised from \(-80\) to \(+60\) mV, the MTSES modification rate is significantly reduced while the MTSET
modification rate remains roughly the same. Fig. 9 B shows the averaged results of MTSET modification at three voltages; the modification rates are all the same. On the other hand, the MTSES modification rate of the Y512C mutant varies according to voltages (Fig. 9 C), and the modification shows a consistent pattern—the more depolarized the membrane potential, the slower the MTSES modification rate. This appears to be a state-dependent modification because the change in the modification rate correlates more with the $P_o$ of the channel but not with the membrane voltage. This state-dependent modification is further supported by an experiment in which the MTSES modification was conducted in a different $[\text{Cl}^-]_o$. In the presence of a higher $[\text{Cl}^-]_o$, the fast-gate $P_o$ is larger, and the MTSES modification rate is consistently smaller than that at a low $\text{Cl}^-$, even though the modification voltage is the same (Fig. 9 D). Such a pattern of modification rate is not consistent with a gated access of MTS reagents to the position behind a gate. Instead, it may suggest a gating-associated change in the pore that alters the MTSES modification rate.

**DISCUSSION**

The present study employs SCAM with charged MTS reagents to examine the accessibility of the amino acid residues of ClC-0 that correspond to those on the helix R of the bacterial ClC channels. The modification pattern from residue 512 to 526 reveals that the modification rate is faster in every third position for both MTSES and MTSET modifications. The most rapid MTSES and MTSET modification rates at four positions (D513C, I516C, K519C, and P522C) are close to the rates of modifying $\beta$-ME in the bulk solution, suggesting that these residues are likely to be freely accessible to the applied MTS reagents. These four positions as well as two other positions (I515C and Y523C) that show higher modification rates than those of S123C and Y512C are located on one side of the R helix that faces the permeant ions (Fig. 4 B). These results argue that the structure of the helix R in ClC-0 is likely to be similar to that of the bacterial ClC channels. For the rest of the positions in helix R of ClC-0, MTSES and MTSET can still modify the introduced cysteine with a smaller rate. The fact that the modification occurs at most positions may be ascribed to the possibility that the packing of the helix to the channel is not very tight. Consequently, the crevice between helices may allow MTS reagents to gain access to most positions in helix R.

Besides the helical arrangement of the positions from 512 and 526, we have obtained two other pictures regarding the functions of the ClC-0 channel. First, the inner pore region of ClC-0 strongly selects anionic over cationic MTS reagents. Second, we find no evidence of a physical gate internal to the central Cl$^-$-binding site. We discuss below the implications of MTS modification rates in the intrinsic electrostatic potential of the pore and in the gating mechanism of the channel.

**Anion to Cation Selection by the Intrinsic Electrostatic Potential in the Pore**

The selection of an anionic over a cationic MTS reagent by the pore of ClC-0 is already present even in modifying the cysteine at the superficial positions such as Y523 and P522. The ratios of MTSES and MTSET modification rates are close to 1 at these positions in contrast to a 124-fold difference for these two compounds to react with the thiol group of $\beta$-ME in the
bulk solution (for β-ME, \(k'_{\text{MTSES}}/k'_{\text{MTSET}} = 0.08\); see Stauffer and Karlin, 1994). The selection for anionic over cationic MTS reagents thus may explain the deviations of the K519C and D513C modifications from the general trend of increasing the ratio of \(k_{\text{MTSES}}/k_{\text{MTSET}}\) (Fig. 4 C). This selection for anions is even more prominent as the position of the cysteine is deeper (Fig. 4 C). At the deepest positions, S123C and Y512C, MTS reagents react with these two cysteines two- to fourfold faster than MTSET—transforming into an anion to cation ratio (\(k_{\text{MTSES}}/k_{\text{MTSET}}/0.08\)) of 20–50 (Cheung and Aka-bas, 1997).

To evaluate this charge selection more closely, we adopt the analysis method introduced by Karlin and his colleagues (Pascual and Karlin, 1998; Wilson et al., 2000). The analysis reports the intrinsic electrostatic potential in the pore region under the assumption that the two MTS reagents are similar in all respects except charge. Therefore, the ratio of \(k_{\text{MTSES}}/k_{\text{MTSET}}\) should equal the ratio of the reaction rates of these two reagents with \(\beta\)-ME (\(k'_{\text{MTSES}}/k'_{\text{MTSET}}\) times a factor \(\rho\) that is a function of the intrinsic electrostatic potential in the pore (Pascual and Karlin, 1998; Wilson et al., 2000). Thus,

\[
\rho = \frac{k_{\text{MTSES}}/k_{\text{MTSET}}}{(k'_{\text{MTSES}}/k'_{\text{MTSET}})}
\]

(1)

and

\[
\rho = \frac{-z_{\text{MTSES}} - z_{\text{MTSET}}}{(F/RT)\psi},
\]

(2)

where \(z_{\text{MTSES}}\) and \(z_{\text{MTSET}}\) are the charge on MTSES and MTSET, respectively, F, R, and T have their usual meanings, and \(\psi\) is the intrinsic electrostatic potential. From Eqs. 1 and 2, we calculate the values of \(\rho\) and \(\psi\) for the mutants that show higher MTSES and MTSET modification rates than those of S123C and Y512C mutants (Table I). It can be seen that as the cysteine position becomes deeper in the pore, the value of the electrostatic potential \(\psi\) becomes more positive, a design of the pore that helps select anions to enter the pore. These intrinsic electrostatic potentials, however, probably do not reflect the potential at the position where the cysteine is introduced, but report the potential several Å away since MTSES and MTSET molecules are probably 6–7 Å in length (Pascual and Karlin, 1998; Wilson et al., 2000). If residues 512–526 of CIC-0 form an α helix, the linear distance between Y512 and K519 should be \(\sim 10–11\) Å. Taking into account of the side-chain length of the target cysteine, the charge on the MTS molecules is likely at a position close to the side chain of K519 when MTS reagents efficiently modify the cysteine at the selectivity filter. This may explain the largest anion to cation selection for modifying the Y512C mutant than modifying the cysteine at other positions. On the other hand, when MTS reagents are modifying I515C, the charge on the MTS molecules probably are 5 Å away in the intracellular side of K519.

The 5-Å distance, however, is short enough for the charged residues at the inner pore mouth to exert their electrostatic influences. Again, we use Eqs. 1 and 2 to calculate the intrinsic electrostatic potential of a location reported by MTS modification of I515C (Table II). In this case, an increase in the total positive charge (or a decrease in the negative charge) on E127 and K519 increases the ratio of \(k_{\text{MTSES}}/k_{\text{MTSET}}\) (Fig. 6 B). The calculated intrinsic electrostatic potential indicates that for an increase of a positive charge at E127 and K519, there is an \(\sim 20\)-mV increase in \(\psi\) (Table II). Altering the charge at the positions 127 and 519, however, probably also affects the local pH surrounding I515C because these three residues are close to each other. When the overall charge at positions 127 and 519 is made more positive, the percentage of thiolate species on the I515C side chain is likely to increase. This may explain the U-shape pattern in the MTSET modification rate (Fig. 6 B).

Although the thiolate formation on the thiol group of the introduced cysteine could alter the absolute MTS reaction rate, the changes in the modification rate cancel out by taking the ratio of \(k_{\text{MTSES}}/k_{\text{MTSET}}\). Therefore, the ratio \(k_{\text{MTSES}}/k_{\text{MTSET}}\) is monotonically increased when the sum of the charges at these two positions is more positive (Fig. 6 B). The roughly staircase jump in the intrinsic electrostatic potential upon altering one total charge at these two positions (\(\sim 20\) mV per charge) suggests that the contributions from these two charges in selecting charged MTS reagents are

| Cysteine mutant | \(\rho_{-40\text{mV}}\) | \(\psi\) |
|----------------|-----------------|-----------------|
| Y523C         | 9.28            | 28.42           |
| P522C         | 10.69           | 30.22           |
| K519C*        | 1.01            | 0.14            |
| I516C*        | 15.29           | 34.78           |
| I515C         | 23.32           | 40.16           |
| D513C*        | 8.09            | 26.67           |
| S123C         | 19.41           | 37.82           |
| V512C         | 48.02           | 49.38           |

This table lists the values of the anion-cationic selection ratio \(\rho\) and the intrinsic electrostatic potential \(\psi\) reported by the MTS modifications of the indicated cysteine mutants. \(V_{\text{SO}} = -40\) mV. The values of \(\rho\) and \(\psi\) were calculated from Eqs. 1 and 2, respectively. As discussed in the text, the value of \(\psi\) probably reports the potential sensed by the charge on the MTS reagent, which is \(\sim 10\) Å away from the listed cysteine position. The symbol “*” indicates a replacement of a charged residue by cysteine, and therefore the intrinsic electrostatic potential of these mutants show a deviation from the general trend.
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The apparent state dependence of Y512C is interesting in that the phenomenon is observed only when MTSES is used as the modifying reagent. Several possibilities might explain this phenomenon, yet each one having its own weakness. First, it might be possible that the side chain of Y512 is part of the fast gate, and when the gate is closed, this side chain directly points to the pore, and thus is more accessible to the MTES reagents in the pore. This proposal faces a severe problem because a similar state dependence would have occurred for MTSET modification. A second possibility is that perhaps in the open state, Cl\(^{-}\) ions coming from the extracellular side would physically knock off the MTS molecule in the pore, thus reduce the probability for the molecule to react with the cysteine at the selectivity filter. This argument is also weak because only Y512C mutant has such a state-dependent effect while the cysteine at other pore positions, such as S123C, does not show this phenomenon. In addition, if this were the mechanism, one should have observed at least part of the knock-off effect for the MTSET modification even though the interaction of Cl\(^{-}\) with MTSES and MTSET might not be exactly the same. The third possibility is that perhaps the fast gate is right behind Y512C (for example, the side chain of E166), and when the gate is open, the MTS reagent is pushed through the pore. Thus, MTS reagents can stay at the selectivity filter longer to react with Y512C when the gate is closed than when the gate is open. This argument again is not strong. Besides the same problem that a similar state-dependent modification does not occur for the S123C mutant, it is not known if a large molecule like MTSES can punch through the pore of CIC-0. The above proposals thus do not reasonably explain this peculiar state-dependent MTSES modification.

In the literature of MTS modification studies, a similar MTSES-only, state-dependent modification was observed when MTS modifications were used to evaluate the proximity of the outer end of the S4 segment to an external pore residue in Shaker K\(^{+}\) channel (Elinder et al., 2001). It was proposed that an increase in the local electrostatic potential due to the appearance of a positive charge can have two effects: the attraction or repulsion of MTS reagents to the reporter cysteine and an increase of the deprotonated state (the thiolate) of the thiol group of the reporter cysteine. These two effects which arise from an approaching positive charge (increase MTSES concentration and thiolate formation) both lead to a higher rate of MTSES modification, but they cancel out each other in the MTSET modification. In CIC-0, the electrostatic potential in the pore is prominent as judged from its controls in the ion flux (Chen...
and Chen, 2003) and in the selection of different charged MTS reagents (the above results). If during the fast-gate opening process a negative charge comes into the pore (or a positive charge moves away from the pore), the reduction in the electrostatic potential will reduce the MTSES modification rate. What are the potential sources for the change of the electrostatic potential in the pore? There are charged residues in the pore of ClC-0, including the side chain of E166 that may serve as the gate. The protein conformational change in the fast gating process surely involves a rearrangement of the positions of these charged residues, and this could result in a different intrinsic electrostatic potential between the closed and the open channels. Another possibility that could result in the changes in the intrinsic electrostatic potential comes from the binding of Cl\(^-\) to the pore when the channel is open. This possibility is directly suggested by the result in Fig. 9 D, in which the MTSES modification rates of the fully open Y512C pore are not the same when Cl\(^-\) concentrations are different. For example, at 60 mV, the fast-gate \(I_p\) of Y512C mutant is close to unity at both 4 and 142 mM [Cl\(^-\)],. However, different MTSES modification rates are observed under these two conditions—the higher the [Cl\(^-\)],, the smaller the MTSES modification rate.

In summary, three conclusions emerge from the present study. First, the MTS modification pattern suggests that the overall alignments of the residues on helix R in the inner pore region are quite similar between ClC-0 and the bacterial ClC channels—the hot residues face the permeant ions as if these residues line the wall of the ion permeation pathway (Fig. 4 B). Second, there is no moving structure on the intracellular side of the selectivity filter, which can physically gate the access of intracellular MTS reagents in response to the opening and closing of the fast gate. Thus, the hypothesis of the side chain of E166 being the fast gate of the channel is applicable to ClC-0. Finally, the electrostatic potential of the ClC-0 pore is prominent. This intrinsic potential in the pore is not only important in selecting charged MTS reagents, but is also likely responsible for the apparent state-dependent modification of Y512C. As the electrostatic potentials of ClC-0 pore is also critical for the functional properties of the channel, understanding the state-dependent MTSES modification on Y512C mutant may help further explore the gating and permeation of ClC-0.

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