

# Ion Transit Pathways and Gating in ClC Chloride Channels

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**ABSTRACT** ClC chloride channels possess a homodimeric structure in which each monomer contains an independent chloride ion pathway. ClC channel gating is regulated by chloride ion concentration, pH and voltage. Based on structural and physiological evidence, it has been proposed that a glutamate residue on the extracellular end of the selectivity filter acts as a fast gate. We utilized a new search algorithm that incorporates electrostatic information to explore the ion transit pathways through wild-type and mutant bacterial ClC channels. Examination of the chloride ion permeation pathways supports the importance of the glutamate residue in gating. An external chloride binding site previously postulated in physiological experiments is located near a conserved basic residue adjacent to the gate. In addition, access pathways are found for proton migration to the gate, enabling pH control at hyperpolarized membrane potentials. A chloride ion in the selectivity filter is required for the pH-dependent gating mechanism. *Proteins* 2004; 57:414–421. © 2004 Wiley-Liss, Inc.

## INTRODUCTION

ClC chloride channels are conserved from prokaryotes to eukaryotes and are involved in many biological functions, including acidification of the stomach<sup>1</sup> and intracellular vesicles, excitability of skeletal muscle, and salt and water transport across epithelia.<sup>2</sup> Mutations in genes coding for ClC channels are associated with several diseases.<sup>3</sup> Recently, it has been determined that ClC homologues in *E. coli* perform as anion-selective electrical shunts coupled with proton pumps that respond to extreme acid conditions;<sup>4–6</sup> the experiments have additionally revealed that the bacterial structure, while closely related to eukaryotic ClC channels, is a proton/chloride exchange transporter<sup>7</sup> (see discussion section below). Early physiological evidence from the Torpedo ClC-0 channel suggested a double-barreled structure that conducts with two equally-spaced open current levels during intermittent burst periods, which are separated by periods of the inactivated state.<sup>8,9</sup> The channels show evidence of multi-ion permeation.<sup>10</sup>

The recent determination of the crystal structures of wild-type and mutant bacterial ClC channels has given dramatic proof of the double-barreled geometry of the channels.<sup>11,13</sup> Sequence alignment exhibits a substantial degree of conservation between bacterial and eukaryotic ClC channels; the similarity is especially strong in the selectivity filter region. Mutational studies on eukaryotic channels correlate well with the locations of key residues in the bacterial

structures.<sup>2</sup> Thus the bacterial structures can be expected to yield insights into the functioning of eukaryotic channels.<sup>12</sup>

Each monomer of the *E. coli* channel contains 18  $\alpha$ -helices, which are arranged to create an hourglass pore for chloride ion passage.<sup>11</sup> The distribution of  $\alpha$ -helices is quite complex, creating an antiparallel architecture within each monomer. Intracellular and extracellular vestibules are separated by a narrow selectivity filter roughly 15 Å in length. Four highly conserved regions stabilize a chloride ion in the filter. The conserved sequences occur at the N-termini of the  $\alpha$ -helices; the neighboring partial charges create a favorable electrostatic environment for binding of the chloride ion. The pore exhibits substantial curvature. A glutamate residue near the extracellular end of the filter blocks the pore, and Dutzler and coworkers<sup>11</sup> have proposed that this residue gates the channel. Since the open probability and the external chloride concentration are closely coupled, it was suggested that a chloride ion replaces the glutamate gate to initiate permeation.

Following the discovery of the bacterial structure, Dutzler and coworkers<sup>13</sup> examined the gating mechanism in ClC channels by determining the crystal structures of two *E. coli* mutants (E148A and E148Q), in which the proposed glutamate gate is replaced by alanine or glutamine, respectively. In both mutants, the side chain is moved from the glutamate binding site, which blocks the pore, suggesting open conformations. Three chloride ions are bound in the mutant structures, one at the previously observed filter binding site ( $S_{cen}$ ), one near the intracellular entrance to the filter ( $S_{int}$ ) and one near the glutamate binding site close to the extracellular end of the filter ( $S_{ext}$ ). In conjunction with the bacterial structures, ClC-0 channels were mutated to study altered physiological behavior. The open probability increased significantly for the mutants, providing further evidence linking the glutamate to gating function. In addition, lowering the external pH for the wild-type channel increased the open probability, making it similar to the two mutants. A gating charge value close to 1 best fit the open probability versus voltage curve; it has been proposed that the chloride ion itself is the gating charge for the ClC channels as it moves from an external binding site into a filter binding site.<sup>10,14</sup>

Previous work showed the important role of external chloride ion concentration<sup>10,14,15</sup> and pH<sup>15</sup> in the voltage-dependent gating mechanism. Increasing external chlo-

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ride concentration at a fixed pH shifts the open probability versus voltage curves to the left (increased open probability at higher chloride ion concentration); depolarization leads to near unity values of open probabilities. Decreasing pH increases the open probability under hyperpolarization conditions, but it does not shift the curve horizontally. Chen and Chen<sup>15</sup> developed a gating model that involves two separate voltage-dependent gating mechanisms: one that emphasizes depolarization potentials and one that describes pH-dependent gating at hyperpolarization. This model accurately fits the gating data.

In this article, we examine the wild-type and mutant *E. coli* CIC channels via molecular modeling techniques to explore permeation pathways, selectivity and chloride concentration as well as the pH dependences of gating (see ref. 16 for a concise summary of the current experimental work). We employ a new algorithm that exhaustively searches for favorable pathways for ion transit. The results provide a detailed picture of the course of ion transit through the complex bacterial CIC channels.

## METHODS

We utilized two algorithms to search for ion permeation pathways. The first method was developed by Smart and coworkers.<sup>17</sup> The algorithm requires knowledge of an initial location within the pore. Simulated annealing is employed to locate the geometric pore center (point at which the largest hard sphere can be inserted without overlap with the protein) in a horizontal plane at a given location  $z$ . The ‘energy’ for the statistical weight is  $-R$ , where  $R$  is the distance to the edge of the nearest protein atom. The algorithm then moves up or down in the  $z$  direction (perpendicular to the membrane plane) and initiates a new simulated annealing step. The method generates unambiguous results for well-defined pathways without too much curvature (for example, the KcsA channel). However, due to the soft nature of the annealing potential and the reliance on only geometric features, many irrelevant pathways are located for a more complex structure like the CIC channel. These must be sorted based on geometric and/or electrostatic features to determine which ones are viable for ion transit. The problem is especially severe for the wild-type *E. coli* structure, in which the E148 side chain blocks the permeation pathway. As an initial step, we modified the HOLE algorithm to allow for movement along pores with strong curvature. The problem of locating many candidate pores was accentuated, however, with the increased flexibility of the modified HOLE geometry-based search.

To overcome these difficulties, we have developed a new search algorithm (TransPath) that is more robust than the HOLE algorithm. Here we briefly summarize the essential features of the new algorithm; details are presented in ref. 18. We first generated a continuum dielectric model of the protein by discretization on a three-dimensional grid.\* We

numerically solved the variable dielectric Poisson equation either with the CHARMM<sup>20</sup> Poisson–Boltzmann code or with our own efficient multigrid solver.<sup>21†</sup> Cuts in horizontal planes (parallel to the membrane plane) at several  $z$  locations were examined in order to locate high dielectric spots; any viable transmembrane pathway must pass through a high dielectric spot in the horizontal planes. Configurational Bias Monte Carlo<sup>22</sup> trajectories were initiated from each of these spots. The energy for the statistical weight includes four contributions: a geometric factor  $1/R$  (a harsher potential than in the HOLE algorithm), a hard sphere potential between the individual chain segments and between the chain segments and protein atoms, the electrostatic potential from the variable dielectric protein model and an external potential that nudges the biased random walk either up (extracellular) or down (intracellular). Each continuous term includes a scaling factor to yield comparable magnitudes for the various contributions during the biased random walks.

We generated swarms of Monte Carlo trajectories and ranked their importance based on the Rosenbluth weight.<sup>22</sup> Failed trajectories that did not make their way out of the protein were discarded. In this way, we collected an ensemble of trajectories that pass from one side of the channel to the other. Once the trajectories were generated and a representative collection was chosen based on the Rosenbluth weight, simulated annealing steps were conducted in 85–95 horizontal planes (with a plane separation of 0.5 Å) to locate the geometric pore center. The advantages of this algorithm are that no initial knowledge of the pore is required and that the annealing step is conducted after generation of an electrostatically driven statistical ensemble of viable pathways.

Generally, the paths we found collapsed into one or a few prototype paths upon annealing. The method has been tested by comparison with previous results on the potassium channel<sup>23</sup> and with several HOLE results on the CIC mutant channels. We have found this method to be faster and more thorough than the HOLE method, and the incorporation of electrostatic information from the start aids in locating important paths for ion transit. We note that we do not include ion self-energy contributions in the potential appearing in the TransPath algorithm (which would require separate Poisson solutions for each chloride ion location). These energies are always positive and will have significant values in narrow pore regions (for example, roughly 0.5 V for the gramicidin pore, ref. 25). It is shown in ref. 25 that small protein fluctuations stabilize the ions and to a large extent cancel the self-energies. For example, the fluctuations lead to a reduction in the potential of mean force (which includes the dielectric self-energy) of nearly 0.6 V for a potassium ion in the gramicidin channel relative to results for a rigid channel geometry. There is no guarantee of a similar reduction in the chloride channel, but it is clear that protein fluctuations are important during ion passage (both for geometric and

\*Dielectric constants of 4 and 80 were used for the protein and water, respectively. The calculations were performed without a surrounding membrane; some test calculations were completed with a membrane that had no noticeable effect on the pore potential. The dielectric profile for the electrostatic calculations utilized the Connolly

surface. Default charges were assumed. The pore radii for the hole search calculations used hard core parameters.<sup>19</sup>

†CHARMM v. 28 was used in our calculations.

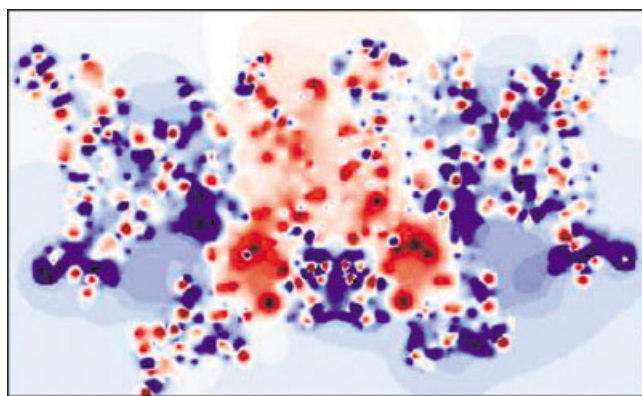


Fig. 1. Electrostatic map of the *E. coli* wild-type structure. Blue shows positive potential, and red shows negative potential.

electrostatic reasons). The self-energy contribution is shown for the chloride path through the rigid E148Q mutant structure below to illustrate its magnitude. The rest of the computational results present the potential due to the variable dielectric protein model but do not include the ion self energies. The  $pK_a$  calculations for the E148 protonation state presented at the end of the results section include self-energy contributions for the charged residue in solution and in the protein environment.

The paths located are single representative paths, which include geometrically and electrostatically favorable features; of course the diffusional trajectories of individual ions can follow an infinite number of paths around these idealizations. The purpose of this work is to explore candidate ion transit pathways and gain insights into the relative energetics during ion passage.

## RESULTS

We first present an electrostatic map of the wild-type C1C structure from ref. 13 (Fig. 1). The potential is shown in a vertical cut through the channel in a plane slightly off from center. The map shows the positive potential (blue) profile along the chloride ion path in each of the monomers. The positive profile is even more distinct in the mutant E148Q, which is proposed to correspond to an open structure. Also, notice a large domain of negative potential (red) at the top of the channel centered in the region around the dimer interface. This negative potential arises from several acidic residues (six on each monomer). This domain plays a crucial dual role: it directs chloride ions outward toward the entrance to the two anion transit pathways, and it creates a favorable electrostatic environment for penetration of protons to the glutamate gate (see below).

Next we examine the chloride ion transit path through the mutant structures (E148A and E148Q; we label the wild-type structure E148). Illustrations of the paths appear in Figures 2(a,b), and the pore radii and potentials are shown in Figure 3. The radius profile illustrates the selectivity filter region (roughly 15 Å) separating the broad intracellular and extracellular vestibules. The filter region is quite narrow ( $R_{\min} = 1.36$  Å for E148A and 1.05 Å, near the glutamine, for E148Q compared to the chloride ion radius of 1.8 Å); clearly

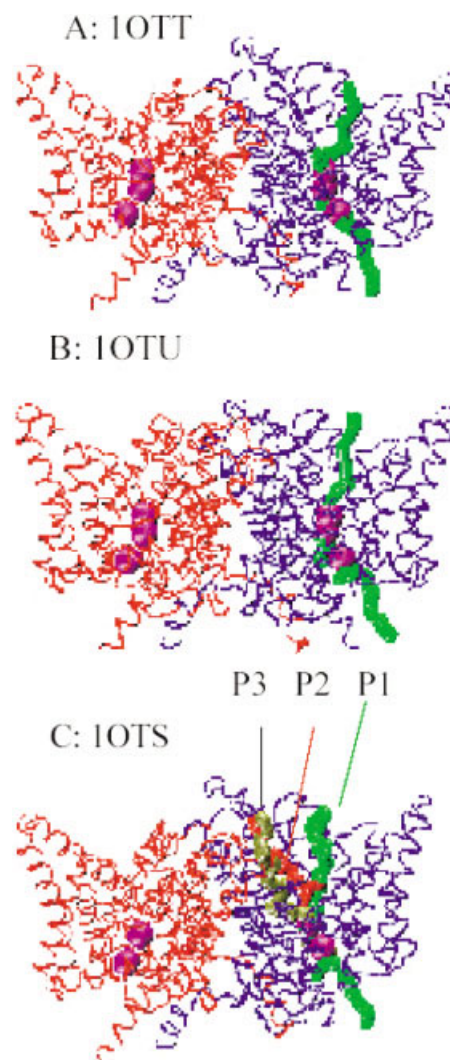


Fig. 2. Images of the various ion transit paths. The bacterial C1C channels are shown from the side with the z-direction vertical. The red and blue portions of the proteins are the two dimers composing the double-barreled channels. (a) The mutant structure E148A. The purple spheres locate the three negative ion binding sites  $S_{\text{int}}$ ,  $S_{\text{cen}}$  and  $S_{\text{ext}}$ . The green polymer follows the chloride ion pathway. (b) The mutant structure E148Q. Colors as in (a). (c) The wild-type structure. The three paths (P1, P2 and P3) are discussed in the text. The protein images were created with VMD software.<sup>48</sup>

protein fluctuations are required for the passage of anions. The same behavior is observed in the selectivity filter of the KcsA channel ( $R_{\min} = 0.78$  Å from TransPath, while the potassium ion radius is 1.4 Å). The potential profile displays a very large positive region through the filter (with a maximum of 1.7 V) and broad low-level positive domains in the two vestibules arising from exposed basic residues.<sup>26</sup> The large positive potential suggests that one or more chloride ions will be located in the filter.<sup>12,26</sup> The potential profile for the E148Q structure with a chloride ion placed at  $S_{\text{cen}}$  is presented in Figure 3(e). The potential exhibits a large positive lobe near  $S_{\text{ext}}$  of magnitude 1.1 V, which indicates that there is a strong tendency to attract a second anion into the filter even when the  $S_{\text{cen}}$  site is occupied. When the glutamine is mutated back to a charged glutamate at the

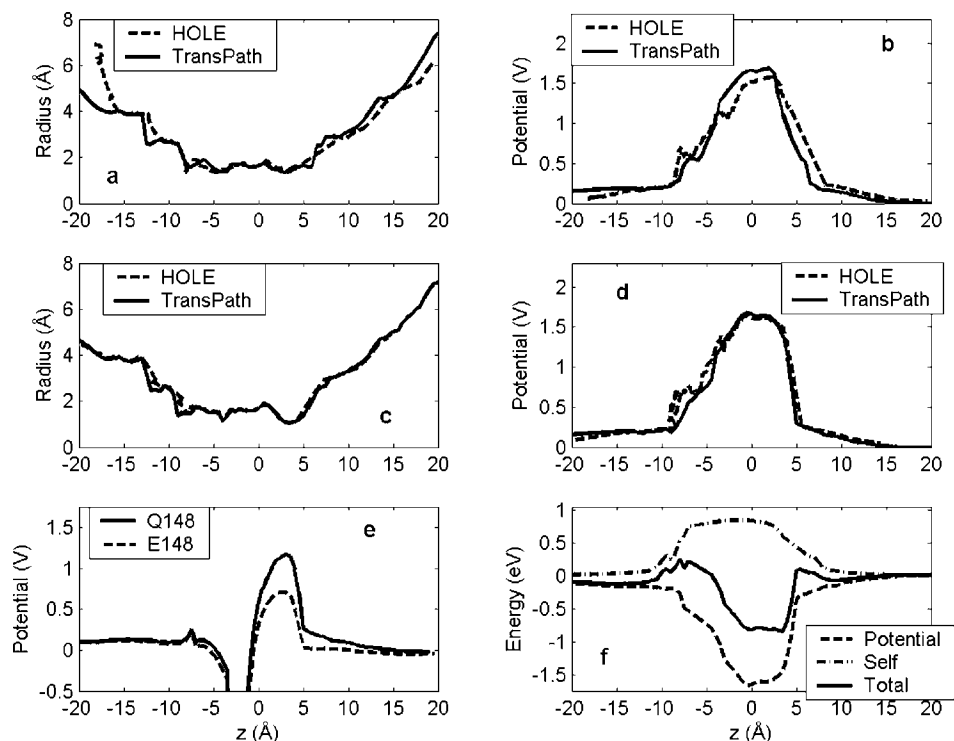


Fig. 3. Mutant radius (a,c) and potential (b,d) profiles for chloride ion pathways. Plots (a,b) are for the E148A mutant, while (c,d) are for the E148Q mutant. Results from HOLE and TransPath are compared. Dashed lines are the HOLE results, and the solid lines are the TransPath results. (e) The potential profile through the E148Q structure with a chloride ion located at  $S_{\text{ext}}$  (solid). The dashed curve is for the same structure with the glutamine mutated back to a charged glutamate residue. (f) Chloride ion self-energy calculation for the E148Q structure. The potential due to the protein is multiplied by the chloride ion charge to generate the attractive potential due to the protein (long dashed). The self energy (dot/dashed) and the total energy (solid) display the relative magnitude of the self energy in relation to the protein potential.

same location, the potential is still 0.7 V at  $S_{\text{ext}}$ . This result is consistent with the fact that the ClC-0 channel can conduct even at high pH values, where the E148 residue is likely charged in both closed and open conformations.<sup>15</sup> The large binding energy in the filter is partially counteracted by a self-energy contribution in a continuum-level treatment. The self-energy contribution for the fixed E148Q structure is displayed in Figure 3(f). It attains a magnitude of 0.9 V at the center of the narrow filter and rapidly drops toward zero in the two vestibules. Slight barriers exist in the total ion energy profile near the intracellular and extracellular entrances to the filter. As discussed above, however, the self-energy can be partially compensated by electrostatic effects due to small-amplitude protein fluctuations (ref. 25).

Consistent with the above results, both the X-ray structure<sup>13</sup> and results from physiological experiments<sup>10</sup> indicate that ClC channels display multi-ion conduction. The chloride ion path has substantial curvature; it is directed toward the portion of the vestibules distant from the dimer interface in order to access regions of positive potential. The path confirms the initial proposal of Dutzler and coworkers<sup>11</sup> and passes the three chloride binding sites (at  $z = -8.73$  Å,  $-2.71$  Å and  $1.01$  Å) determined from the crystal structures.<sup>13</sup> Both the HOLE and TransPath search algorithms located this path. Pathway searches on the wild-type structure, on the other hand (below), exhibited a narrower restriction near the glutamate ( $R_{\text{min}} = 0.71$  Å),

and the electrostatic potential profile is considerably reduced in magnitude compared to the mutant profiles. Therefore, the path search results support the proposal that the two mutant structures are indeed open, and the wild-type channel is closed.

We now focus on the wild-type structure to address the mechanism of gating starting from the closed state. The experiments of Chen and Chen<sup>15</sup> show that the channel opens at a pH of 9.6 under depolarization. This suggests that the glutamate gate can operate while charged. In Chen and Chen's two-mechanism model, this correlates with the depolarization-driven rate and carries a gating charge of approximately one. The electrostatic profile [Fig. 4(b)] along the chloride ion path [P1 of Fig. 2(c)] displays a positive shoulder of 0.2 V (7 kT) in the neighborhood of  $z = 8$  Å. This location is very near Arg147 (and just prior to a drastic narrowing of the pore radius). We label this location  $S_{\text{bs}}$  and propose that this is the external binding site suggested by Chen and Miller.<sup>14,16</sup> In their model, the chloride ion binds to the  $S_{\text{bs}}$  site with little voltage dependence, presumably because the site is located in the vestibule and is exposed to the external solution. The site attracts external chloride ions to a location near the gate.<sup>27</sup> The distance between the Glu148 and Arg147 side chains decreases substantially (from 6.72 to 3.56 Å) between E148 and E148Q. One possibility is that, during gate opening, a chloride ion at  $S_{\text{bs}}$  replaces the charged

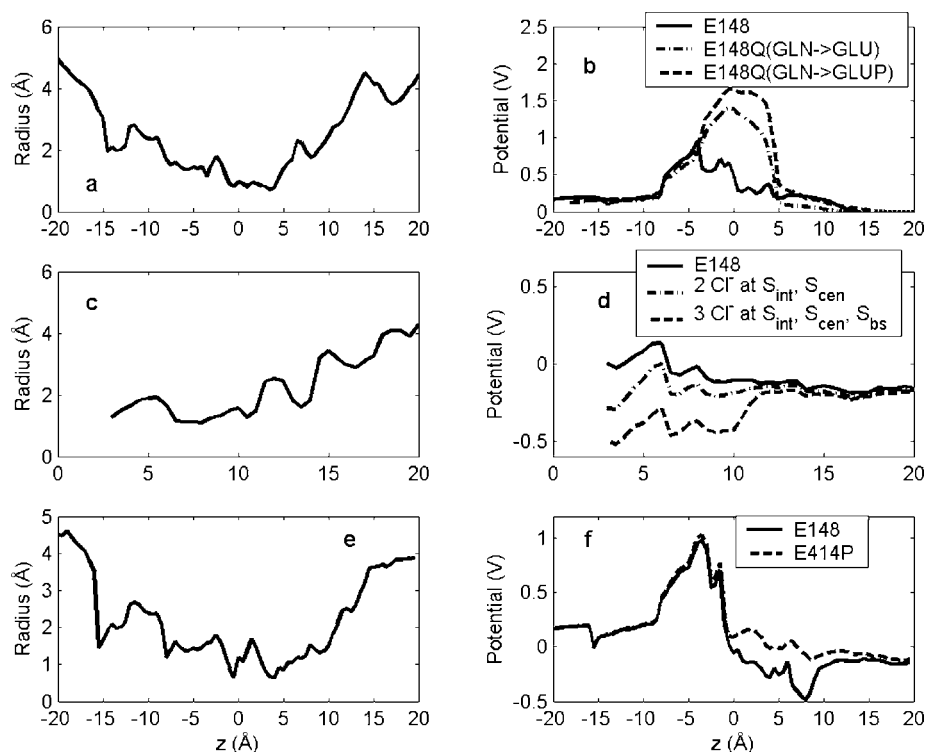


Fig. 4. E148 pathway radii and potentials: the pore radius and potential profile for paths P1 (a,b), P2 (c,d) and P3 (e,f). Plot (b) displays three potential profiles: the chloride path for E148 (solid), the chloride path for E148Q with the glutamine mutated to neutral Glu (dashed), the chloride path for E148Q with the glutamine mutated to a charged Glu (dot/dashed). For the P2 path potential profiles (d), the solid line represents E148 with no chloride ions, the dot/dashed is for two chloride ions, one at  $S_{int}$  and one at  $S_{cen}$ , and the dashed has an additional chloride at  $S_{bs}$ . The P3 potential profile (f) is solid, and the dashed line is the profile when Glu414 is neutralized.

glutamate in a concerted process whereby the ion fills the  $S_{ext}$  location while the charged Glu148 side chain moves into closer proximity to the Arg147 side chain. Notice that the potential profile for the chloride ion path with the E148Q glutamine mutated back to a charged Glu residue still maintains a strong positive potential entering the filter [Fig. 4(b)].

The pH-dependent hyperpolarization-driven mechanism corresponds to the second mechanism proposed by Chen and Chen,<sup>15</sup> with a gating charge of approximately  $-0.3$ . This mechanism leads to increased opening rates at low pH and negative potentials. Presumably, protonation of the Glu148 gate results in increased open probabilities as the neutral side chain is freed from the strong binding energy at  $S_{ext}$  when charged. In order for this second mechanism to work, protons must access the Glu148 side chain. We propose that the acidic residues at the top center of the channel provide the favorable electrostatic environment required for proton propagation to the gate. Recent simulation work on proton diffusion through channels indicates the important role of electrostatic driving forces in proton migration through channels.<sup>28,29</sup> To explore this proposal, we converted our test charge in the TransPath search algorithm to a positive charge to locate candidate proton access paths.

The first candidate proton path [P2, Figs. 2(c) and 4(c,d)] follows the side of the extracellular vestibule nearer the

dimer interface. The computed pore is quite wide in the vestibule, and the potential in regions away from the gate is negative ( $-0.2$  V minimum near the extracellular exit, and  $R_{min} = 1.09$  Å near Arg147). While protons will diffuse along this direction to near  $z = 7$  Å, there exists a potential barrier for penetration of the proton to Glu148 due to the Arg147 side chain. If we place chloride ions at  $S_{int}$ ,  $S_{cen}$  and  $S_{bs}$ , however, the potential drops considerably to large negative values [Fig. 4(d)]. If chloride ions are placed at  $S_{int}$  and  $S_{cen}$ , an intermediate profile is obtained. Therefore, instead of the original proposal that protonation of an external site may induce chloride permeation directly,<sup>30</sup> the opposite process is suggested by these results; chloride binding at  $S_{bs}$  alters the potential profile so a proton may access the gating region. Experimentally, the influence of external chloride ion concentration on the pH-dependent gating mechanism is not entirely clear (ref. 15, Fig. 7).

A second possible proton pathway [P3, Figs. 2(c) and 4(e,f)] was found, which coincides with the vestibular path discussed above near the extracellular entrance but deviates approaching Glu148. An acidic residue (Glu414) is located near this path, which leads to a negative potential with a minimum value of  $-0.5$  V at  $z = 8$  Å [Fig. 4(f)]. The minimum radius of this path is  $0.63$  Å, so it is questionable whether protons associated with water can penetrate through such a narrow region unless there are substantial protein fluctuations. In addition, if the Glu414 residue is

protonated, the potential becomes unfavorable. It is interesting to note, however, that the Glu414 residue is conserved throughout the CIC family. Both paths (P2 and P3) terminate in close proximity to Glu148.

In addition, one path was found following the dimer interface with a minimum potential value of  $-0.8$  V at  $z = -3$  Å, but it reaches positive values at more negative  $z$ -locations. Therefore, this pathway is likely blocked to motion of ions of either charge. No clear access from this path to the filter region was apparent. Finally, we observed a pathway that follows the negative potential domain (red) in the lower portion of the channel (Fig. 1); this path may be involved in proton leakage through the channel (below). The path passes near three acidic residues (E113, E117, and E203).

Based on these results for the static structure, the vestibular path (P2) appears to be the most likely candidate for extracellular proton access to the proposed gating glutamate side chain due to its favorable electrostatic and geometric features. On the other hand, the conservation of the Glu414 residue through the CIC family suggests that it may play an important role in proton access. Protein fluctuations may allow for proton penetration through the narrower portions of the P3 path near Glu148. The (6 + 6) acidic residues near the channel top center are crucial for providing a negative electrostatic potential driving proton diffusion; when all of the acidic residues in this domain are protonated, the negative potential region is destroyed. Hyperpolarization provides an additional driving force for proton penetration to the gate. Mutations near P2 and/or involving the residues corresponding to Glu414 in eukaryotic channels should be able to test the relative importance of the proton access paths in gating.

Single-channel measurements have suggested that the  $pK_a$  of the CIC gate is roughly 5.3,<sup>15</sup> which implies a positive shift of about one unit. Electrostatic effects can substantially alter  $pK_a$  values in a protein environment.<sup>32–39</sup> To examine the protonation state of Glu148, we performed continuum dielectric calculations on the wild-type structure.\* The  $pK_a$  shift results are presented in Figure 5. With no chloride ions in the vicinity of the glutamate, the  $pK_a$  shift is large and negative (the  $pK_a$  is close to zero at 0.0 V transmembrane potential); the strong positive potential at the  $S_{ext}$  binding site makes the deprotonated state stable. If two chloride ions are placed at  $S_{int}$  and  $S_{cen}$ , the  $pK_a$  value of the glutamate shifts substantially upward, to a value of around 5 (the ion at  $S_{int}$  has little effect on the  $pK_a$ ). If a third chloride ion is placed at the  $S_{bs}$  location, the  $pK_a$  shifts to an even higher value (7.5). This suggests that protonation of Glu148 requires a chloride ion at  $S_{cen}$  in order to create a favorable electrostatic environment for protonation. An additional chloride

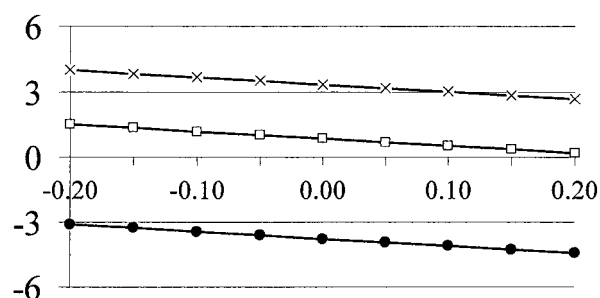


Fig. 5.  $pK_a$  shift versus applied voltage (in volts). Filled dots signify the absence of chlorides, open squares signify two chlorides at  $S_{int}$  and  $S_{cen}$ , and crosses signify the placement of an additional chloride at  $S_{bs}$ .

ion at  $S_{bs}$  further enhances the probability of protonation. The important role of internal chloride concentration on the pH-dependent gating has been observed experimentally.<sup>14,16</sup> This appears to occur through penetration of an internal chloride ion to  $S_{cen}$ .

## DISCUSSION

The computational modeling described here yields chloride ion pathways in agreement with the initial proposal of Dutzler and coworkers.<sup>11</sup> The radius and potential profiles for the two mutant structures support their interpretation as open structures.<sup>13</sup> In addition, the results are consistent with the two-mechanism model put forward by Chen and Chen<sup>15</sup> for channel gating. The binding site at  $S_{bs}$  attracts chloride ions, which are optimally placed to enter the pore when the Glu148 side chain moves out of the way, likely in a concerted mechanism. This mechanism operates at positive voltages. At negative voltages, the gating is influenced by extracellular pH. We have located candidate proton access paths and shown that a chloride ion located in the filter is required to shift the  $pK_a$  value of the glutamate to the appropriate range. Strategically placed acid residues at the top center of the channel both direct chloride ions outward toward the entrance to the pore and create a favorable electrostatic environment for proton access. Since the gating mechanism appears to be relatively localized, further studies should be directed at molecular dynamics simulations to investigate the glutamate motions in charged and neutral forms, and in the presence of neighboring chloride ions.

Pusch and coworkers<sup>41–43</sup> have recently questioned the local glutamate gating hypothesis of MacKinnon and coworkers<sup>13</sup> based on blocking studies on the intracellular side of CIC-0 channels. They found substantially different affinities of the blocker for the open and closed states and suggested that this implies significant conformational changes (in the channel pore and filter) during gating. Part of the evidence for larger conformational changes came from studies of the mutant Y512F, which removes a hydroxyl group from the chloride binding site  $S_{cen}$ . While we cannot preclude conformational changes in our calculations based on fixed structures, we addressed the issue of this particular mutation via electrostatic modeling. First, we performed the mutation Y445F in the E148Q mutant structure and found virtually no change in the electrostatic profile for the chloride ion path through the filter region. This result shows that the hydroxyl

\*The  $pK_a$  calculations were performed with CHARMM v. 28, ref. 20. A protein dielectric constant of 10 was assumed; see ref. 37. The linearized Poisson-Boltzmann equation was solved with successive over-relaxation (SOR) relaxation. The grid size used was 0.5 Å. A membrane dielectric constant of 2 was used. A bathing solution of 150 mM was included. The water probe radius was taken to be 1.4 Å, with an ion exclusion radius of 2 Å. The membrane thickness was 34 Å. Default charges were assumed. We also utilized the UHBD code, ref. 40, with comparable parameters and results.

group is not crucial for determining the electrostatic profile through the filter. Second, the potential profile through the wild-type E148 filter region drops significantly from the E148Q profile due to the presence of the charged glutamate; the potential at  $S_{cen}$  decreases from 1.409 V in E148Q to 0.545 V in E148. Therefore, it can be expected that the chloride ion occupancy at  $S_{cen}$  in E148Q is substantially larger than it is in E148. Locating a chloride ion at  $S_{cen}$  in E148Q leads to a potential of 0.099 V at  $S_{int}$ , compared to 0.246 V for E148. Hence, the decreased affinity for the blocker in the open state could be equally explained by electrostatic effects due to increased occupancy of the open channel filter by chloride ions.

In this article, we have focused on chloride ion transport mechanisms through the bacterial CIC channels and the associated proton access to the proposed gating region. A recent study<sup>7</sup> has suggested possible proton/chloride antiport behavior in the bacterial structures. Since secondary active transport requires conformational transitions in the protein in addition to the fast chloride gate, this issue has not been addressed in the present study. A possible mechanism for moving protons completely across the protein is unknown. It is interesting to note, however, the two negative potential lobes in the lower part of Figure 1, which may provide zones through which protons diffuse, perhaps associated with protein rearrangements. In preliminary computational results for a homology model of the CIC-0 channel, the negative potential at the top channel center is maintained, but the two negative potential lobes in the lower domains of the protein disappear; the lack of a negative potential domain there would prevent proton transport across the channel. This change in electrostatic distribution is likely due to the replacement of the three acidic residues in the bacterial channel with basic or neutral residues in CIC-0 (E113 → K, E117 → R, E203 → V from the bacterial/CIC-0 alignment). Mutation studies involving these three residues, in addition to the observed disruption of proton coupling for the E148A mutation, could test their involvement in proton motion across the structure.<sup>7</sup> There does exist a close correspondence between the bacterial CIC structures and observed physiological behavior of CIC-0 channels through a range of mutational studies.<sup>2,12,13,16,44</sup> It is likely that, while the bacterial/eukaryotic similarity may not be complete, it does capture key features of chloride ion motion through the CIC channel family.\*

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\*Following submission of this paper, three papers appeared concerning molecular modeling of CIC chloride channels. They are listed here as refs. 45–47.

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