Proton Release from the Histidine-Tetrad in the M2 Channel of the Influenza A Virus

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ABSTRACT: The activity of the M2 proton channel of the influenza A virus is controlled by pH. The tautomeric state and conformation of His37, a key residue in the M2 transmembrane four-helix bundle, controls the gating of the channel. Previously, we compared the energetics and dynamics of two alternative conformations of the doubly protonated state at neutral pH, namely, a 4-fold symmetric “histidine-box” and a 2-fold symmetric “dimer-of-dimers”, and proposed a multiconformation model for this charge state. Here, we elaborate this model by further studying configurations of the His37 tetrad in the triply protonated state and its subsequent deprotonation via quantum mechanics/molecular mechanics (QM/MM) molecular dynamics (MD) simulations, starting with the aforementioned configurations, to gain information about proton release in a viral membrane environment. Interestingly, the two configurations converge under acidic pH conditions. Protons can be transferred from one charged His37 to a neighboring water cluster at the C-terminal side of the channel when the Trp41 gate is open transiently. With limited backbone expansion, the free energy barrier for proton release to the viral interior at low pH is ∼6.5 kcal/mol in both models, which is much lower than at either neutral pH or for an isolated His37 cluster without a membrane environment. Our calculations also suggest that the M2 protein would seem to exclude the entrance of anions into the central channel through a special mechanism, due to the latter’s potential inhibitory effect on proton conduction.

INTRODUCTION

Proton transfer is a ubiquitous phenomenon in chemical and biological systems. Interestingly, the trajectory of a proton in the lumen of the so-called M2 channel of the influenza A virus is an example from Biology. The M2 integral membrane protein is a tetramer assembled from four identical subunits, each with 97 residues.1,2 As a prerequisite for release of genetic material to the cytoplasm, the interior of a viral particle must become acidiﬁed to reestablish in full the original conformation for the next cycle. Among the pore-lining residues, His37 and Trp41 were identiﬁed as the most important residues in the proton transfer through the M2 channel. The proton selective conductance in M2 is achieved by the presence of the His37 cluster: substitution of His37 with residues such as Gly or Ala induces a nonselective channel, indicating a mechanistic role of His37 in proton relay.6,7 Each of the four His37 may have two tautomeric forms at neutral conditions. A transfer cycle was proposed where a tautomerization of the His37 side chain occurs between the acceptance of a proton from the viral exterior and the release of a proton to the viral interior, to

reestablish in full the original conformation for the next cycle. On replacing the imidazole ring of the His37 side chain with a 4-thiazolyl group, which has the Nε1 as the only proton acceptor with pKa ∼2.5, this mutant shows inward proton flux resembling the wild type, thus demonstrating that tautomerization is not necessarily required for proton transport.8 However, rotation of the imidazole ring during proton translocation cannot be ruled out. Besides the functional role in proton selectivity, His37 is associated with channel activation: binding of a third proton to the His37 cluster under acidic conditions triggers a detectable proton ﬂux, with a peak conductance rate of 10⁹ protons per second.3,9 Trp41, on the other hand, serves as the channel gate.10,11 Substitution of Trp41 with smaller residues, like Gly, Ala and Phe, induces higher proton conductivity compared to the wild-type channel.10

Despite the relatively small size of the M2 protein, and several high-resolution structures being available, the mechanism of proton conduction through the channel is not completely understood (Figure 1A).12 Previously, we studied the addition of a proton to the His37 cluster from the extraviral

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104 protons per second.3,9 Trp41, on the other hand, serves as the channel gate.10,11 Substitution of Trp41 with smaller residues, like Gly, Ala and Phe, induces higher proton conductivity compared to the wild-type channel.10

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side, and compared the energetics and dynamics of two configurations of the doubly protonated state at neutral pH—a 4-fold symmetric "histidine-box" and a 2-fold symmetric "dimer-of-dimers"—and proposed a multiconfiguration model where these configurations are in moderately fast equilibrium. Here, we focus our analysis on the configuration of the His37···Trp41 quartet in the triply protonated state, and its subsequent deprotonation.

Herein, quantum mechanics/molecular mechanics (QM/MM) molecular dynamics (MD) simulations reveal insights into the function and mechanism of this protein. Specifically, molecular simulations complemented by crystallographic and NMR (nuclear magnetic resonance) determined structural data, as well as physiological measurements provide mechanistic clues in understanding this rate-limiting step of proton release toward the intraviral side. Then, we calculated potentials of mean force (PMFs) of the deprotonation by constrained QM/MM MD simulations: the PMFs of both models have barriers ~6.5 kcal/mol. Compared to previous simulation studies, the local electrostatic environment of the hydrated membrane significantly lowers the energy cost. Finally, we explored the possible role of a Cl− anion in the pore at low pH, which was found to stabilize the highly charged 3+ histidine cluster, but also to impede proton release by stabilizing the hydronium species within the channel pore.

**COMPUTATIONAL DETAILS**

For clarity, we named the “dimer-of-dimers” configuration at the 2+ state (with 2 of 4 His37 side chains charged) at neutral pH as the “D2-model”, and the subsequent 3+ state obtained by adding a third proton as the “D3-model" (Figure 1B). Correspondingly, the “histidine-box” at 2+ state and the subsequent 3+ state structures are designated as the “H2-” and “H3-models" (Figure 1C), respectively.

**Classical MD Simulations.** The 1.65 Å resolution X-ray structure (pdb entry: 3LBW, residues 25–46) was used as the initial structure. Two and three of the four His37 residues were initialized to be charged in the 2+ and 3+ states, respectively. The protein was then embedded in a previously equilibrated 80 × 80 Å2 lipid bilayer containing 171 1-palmytoyl-2-oleoyl-sn-glycero-3-phosphatidylcholine (POPC) molecules, hydrated by 10,031 water molecules. Potassium and chloride ions were added to neutralize the system, and to provide a solution buffer with ionic concentration of 150 mM.

Periodic boundary conditions were applied, and long-range electrostatic interactions were treated with the particle mesh Ewald method. A time step of 2 fs was used, and all of the bonds involving hydrogen atoms were constrained with the SHAKE method. After an equilibration period with gradually released harmonic restraints, production simulations were performed at 310 K and 1 atm using Langevin temperature and Langevin piston pressure coupling schemes. The TIP3P force field was used for water molecules, and CHARMM force field was used for the protein and lipid atoms. Simulations were performed with NAMD, and continued for 200 ns for the 2+ state, and 300 ns for the 3+ state.

**Quantum-Mechanical Optimization of the His37···Trp41 Quartet Structure at 3+ State.** To prepare the His37···Trp41 quartet structures for the subsequent H3- and D3-models at the acidic condition with QM/MM MD simulations, we followed the protocol used in our previous work to optimize the His37···Trp41 quartet structure at the 2+ state with the ONIOM method implemented in Gaussian 03. On the basis of the optimized His37···Trp41 quartet structure in the H2- and D2-models, a third proton was added to a neutral His37 side chain, and ONIOM was used to optimize the system after this addition. Then, the H3 and D3-models were constructed with the original quartet in the corresponding H2(D2)-model replaced with the newly optimized quartet at the 3+ state. Therefore, the initial backbone of the H3(D3)-model is similar to the preceding H2(D2)-model. With the quartet structure held fixed, we then relaxed the remaining atoms of the H3(D3)-model with classical MD simulations for 20 ns. We took the last snapshot from the MD simulations, and minimized the system with the H37···Trp41 quartet fixed. The minimized structures were used for the following QM/MM MD simulations.
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Figure 2. Dihedral angle distributions of the His37 and Trp41 side chains from the D- and H-models are plotted at neutral and acidic pH conditions. The top panel shows the plots at neutral conditions (2+ state), which are labeled as a−f; the corresponding plots at acidic conditions (3+ state) are labeled as a′−f′, shown in the bottom panel. The experimental structures at either condition are used for comparison. Panels a and b show the correlation between $\chi_1$ and $\chi_2$ on His37 side chain from the QM/MM and classical MD simulations, respectively. Panels c and d show the correlation between $\chi_1$ and $\chi_2$ on Trp41 side chains. Panels e and f show the correlation between $\chi_2$ on His37 and $\chi_2$ on Trp41.

**QM/MM MD Simulations.** In the QM/MM scheme, as in our previous study, we included in the QM region the His37-...Trp41 quartet and thirteen water molecules nearby (corresponding to the three water clusters identified in the 3LBW.pdb structure). Six waters just above the His37 tetrad, two in the His37-...Trp41 cavity, and five below the Trp41 gate. These atoms were treated at the density functional theory (DFT) level using a mixed Gaussian and plane-waves treatment, while the rest of the system, including the protein, the lipids, and waters, was treated with the MM method. The generalized gradient approximation (GGA) functional BLYP with the D2 level dispersion-correction (BLYP-D2), the triple-$\zeta$ basis set with two polarization functions, and the Goedecker, Teter, and Hutter type pseudopotentials were used. A wavelet-based Poisson solver was used to remove the spurious interactions of the QM region atoms with the periodic images. A box size of $32 \times 32 \times 32$ Å$^3$ was used for the QM region so that the buffer between QM atoms and the box edge is ~8 Å. We saturated the broken bonds with hydrogen atoms, and combined the QM and MM subsystems with the IMOMM methodology, where the scaling factors of 1.355, 1.384, and 1.416 were used to relate the QM C–H bond distance to the MM $C\alpha–C$, $C\alpha–N$, and $C\alpha–C\beta$ distances, respectively.

For the QM/MM MD simulations, after 2 ps equilibration for each model, we accumulated 9 ps trajectories for the D3- and Hutter type pseudopotentials were used. A box size of $32 \times 32 \times 32$ Å$^3$ was used for the QM region so that the buffer between QM atoms and the box edge is ~8 Å. We saturated the broken bonds with hydrogen atoms, and combined the QM and MM subsystems with the IMOMM methodology, where the scaling factors of 1.355, 1.384, and 1.416 were used to relate the QM C–H bond distance to the MM $C\alpha–C$, $C\alpha–N$, and $C\alpha–C\beta$ distances, respectively.

For the QM/MM MD simulations, after 2 ps equilibration for each model, we accumulated 9 ps trajectories for the D3-model with and without Cl, and 4 and 7 ps for H3-model with or without Cl, respectively.

To characterize the free energy profile for proton release from the His37 cluster in M2 protein, we followed the well-tested protocol used in the previous work from our group to run constrained QM/MM MD simulations. The starting configuration was adopted from the preceding 3+ state QM/MM MD simulations. The reaction coordinate $\zeta$ was defined as the distance between the Ne on one His37 side chain and its neighboring He2, in the range of 1.0 Å to 1.6 Å. With the reaction coordinate $\zeta$ fixed in each window, constrained QM/MM MD simulations were carried out for times varying between 5 and 8 ps to obtain a converged PMF. Forces after the initial 1 ps of each constrained simulation were collected to calculate the mean constraint force. Free energy profiles for the proton release were estimated by thermodynamic integration of the mean force along the reaction coordinate.

All of the QM/MM MD simulations were carried out with the CP2K program with a time step of 0.4 fs. The temperature was 310 K, using the Nose–Hoover chains thermostat with a time constant of 1 fs.

**Natural Bond Orbital (NBO) Analysis.** NBO was used to calculate the partial charge at the MP2/6-311++G(d,p) level, performed with Gaussian 03. In those calculations, only the side chains of the His37 and Trp41 residues were included, with the broken $C\alpha–C\beta$ bonds saturated by hydrogen atoms.

### RESULTS AND DISCUSSION

**Structural Integrity.** The conformation of the His37 cluster in both the D3-model and the H3-model was quantified by a single parameter, $\lambda$, defined as the projection of the atomic coordinates of His37 along the displacement vector connecting the two models, where $\lambda = 0$ represents the “dimer-of-dimers” (D2-model) and $\lambda = 1$ the “histidine-box” (H2-model). Upon binding the third proton, as shown in Figure 1D, the $\lambda$-value in the H3-model remains at ~1, indicating that the protonation does not change the conformation of the His37 cluster significantly. In the D3-model, His37 shifts from $\lambda = 0$ to $\lambda = 0.4$, showing that the D2-model and the corresponding D3-model are connected by a pH-dependent transformation. Furthermore, this transformation appears to form an intermediate state of the one leading from the “dimer-of-dimers” ($\lambda = 0$) to the “histidine-box” ($\lambda = 1$). This conformational change has been previously proposed for the “dimer-of-dimers” at acidic conditions, where one dimer is broken upon taking a third proton, to avoid electrostatic repulsion. The structure of the His37 cluster after this
conformational change is closer to the “histidine-box” conformation.

In both models, the presence of a Cl\(^{-}\) ion in the His37…Trp41 cavity does not have a major impact on the protein’s conformation (Figure 1B–D), but has more profound effects on proton conduction, which is further described below.

Rotamer Structures. His37. We monitored the specific structural motifs of either model by exploring the rotamer structures of His37 and Trp41. As disclosed by experimental determined structures at different conditions, these two key residues at both high and low pHs adopt rotameric angle \(\chi_1 = 180^\circ\), whereas the angle \(\chi_2\) is relatively flexible. In particular, the side chain dynamics of the two residues are sampled in the classical MD simulations (Figure 2b and 2b’), largely because of the inability of the classical force field to reproduce the strongly hydrogen bonded “dimer-of-dimers” of the D2-model. In the D3-model (Figure 2a’), one of the charged His37 side chain in the broken dimer remains at the [180°, 60°] configuration with the other one swings at [180°, 150° ~ 220°], and the intact histidine-pair resembles those in the D2-model. Interestingly, this [180°, 60°] configuration is the characteristic of the low pH structure, which was occasionally sampled in the classical MD simulations at the 2\(^{+}\) state, but never reached in the corresponding QM/MM MD simulations. In both the H2- and H3-models, all of the four His37 side chains have \(\chi_{1,2}\) values of [180°, 180°].

Trp41. The Trp41 side chains feature a “flip” variant in the H-models (the Nε1-Hε1 bond of the indole group points toward the central pore) and a “flop” variant in the D-models (the same bond points away from the central pore). These variants correspond to \(\chi_2\) angles located at \(\sim 80^\circ\) (tp rotamer, the “flip” variant) and at \(\sim 280^\circ\) (tm rotamer, the “flop” variant), which are retained at both high and low pH conditions in each model (Figure 2c and 2c’). However, the \(\chi_2\) angle of Trp41 at the low pH conditions in the D3-model is more diffuse than the congener in the D2-model, indicating a more dynamic Trp41 gate in the D-model under acidic conditions. In contrast, this gate seems to be more rigid in the H-models, as evidenced by the similar distributions of \(\chi_2\) angle at neutral or acidic pH conditions. Due to the inward orientation of Trp41 in the H3-model, the excess proton is less easy to be released toward the intraviral end of the channel, as further described below. However, this orientation enables the Trp41 side chains to form water-mediated hydrogen bonding interactions with Asp44 (Figure 3), which was proposed to stabilize the Trp41 gate, a critical factor for the asymmetric proton flux. The outward orientation of the Trp41 in the D3-model, on the other hand, points the amide group toward the hydrophobic wall, which seems to be a less favorable factor for its gating.

It should be noted that, in the classical and QM/MM MD simulations of both D-models, only the tp rotamer of Trp41 was observed (Figure 2d and 2d’), as the “histidine-box” X-ray structure was used as the initial structure and no indole ring flipping happened in the submicrosecond time scale simulations.

The average Hη2…Hη2 distance of neighboring Trp41 residues in the D2- and D3-models are 4.7 and 7.2 Å, respectively, which are consistent with the data derived from \(^{19}\)F NMR measurement on \(^{19}\)F(HNM) M2TM sample (\(\sim 3.2\) Å at pH 8.0 and \(\sim 8.0\) Å at pH 5.3). In contrast, the average Hζ3…Hζ3 distance is \(\sim 11\) Å in both the H2- and H3-models. This is in line with a NMR experiment under different conditions, in which the \(^{19}\)F(Hζ3) M2TM sample was found to have the closest distance of \(\sim 11\) Å at both high and low pH conditions.

His37…Trp41 Coupling. As the M2 protein’s structure is sensitive to the change in the solubilizing environment, the diversity of the His37…Trp41 contact depends on the balance between different rotamers. The nature of the interaction between His37 and Trp41 differs between the D- and H-models, as the former is more sensitive to the protonation state (Figure 2c,e,f,f’). This difference in the His37…Trp41 packing and its response to the environment can be mainly attributed to the higher flexibility of the His37 side chain compared to Trp41. In the H-model, the orientations of the two residues and their coupling are less dependent on pH condition, showing only small thermal motions for both the imidazole ring of His37 and indole ring of Trp41.

Due to the different rotameric conformations, distances between His37 and Trp41 are also distinct in the D- and H-models, which can be determined by different techniques. As shown in Table 1, the distances measured in our MD simulations are quantitatively consistent with experimental structures or NMR measurements. Taking the Cα(His37)…Hζ3(Trp41) distance as an example, at neutral pH condition, the average distances (Figure 4a) in the D2- and H2-models are 6.8 and 4.7 Å, respectively, corresponding to 6.5 Å in the 2L0j.pdb and 5.0 Å in the 3LBW.pdb.

Free Energy Cost of Deprotonation. To explore the free energy cost for the His37 cluster to release a proton at low pH conditions, we carried out constrained QM/MM MD simulations on the D3- and H3-models imbedded in the fully hydrated lipid bilayers, in which the geometrical parameter of...
Table 1. Geometric Parameters (in Å) of the His37···Trp41 Cluster in Different Computational Models and Experimental Structures

| Model | D2 | D3 | H2 | H3 | 2LOJ<sup>39</sup> | 3LBW<sup>15</sup> | 3BKD<sup>40</sup> | SSNMR<sup>54</sup> |
|-------|----|----|----|----|-------------------|-------------------|-------------------|-------------------|
| H(Nδ)···W(Cγ) | 4.7 | 4.0 | 5.4 | 5.7 | 5.3<sup>b</sup> | 5.5 | 4.2 | / |
| H(Cγ)···W(Hδ<sub>3</sub>) | 7.5 | 7.9 | 5.7 | 5.8 | 6.9 | 5.7 | 6.0 | 8.4<sup>c</sup> (6.2) |
| H(Cδ)···W(Hδ<sub>3</sub>) | 6.8 | 7.3 | 4.7 | 4.8 | 6.5 | 5.0 | 7.0 | / (6.0) |
| H(Cε)···W(Hδ<sub>3</sub>) | 5.5 | 5.6 | 6.3 | 6.5 | 4.9 | 6.0 | 5.0 | 8.4 (7.3) |
| W(Hγ<sub>3</sub>)···W(Hδ<sub>3</sub>) | 4.7 | 7.2 | 10.3 | 10.0 | 4.5 | ~11 | ~11 | ~11 |
| W(Hδ<sub>3</sub>)···W(Hε<sub>3</sub>) | 5.1 | 4 | 11.2 | 10.8 | 3.9 | ~11 | ~11 | ~11 |

<sup>a</sup>Ensemble average of the eight snapshots of the NMR structure. <sup>b</sup>Among the eight snapshots of the NMR structures, the shortest is 4.4 Å. <sup>c</sup>Data at pH = 8.5 and pH = 4.5 (in parentheses) are shown. SSNMR: solid-state NMR.

Figure 4. (a) Shown is the cation–π interaction between His37 and Trp41. The dashed line connecting His1 on His37 and Cy on Trp41 indicates the reaction coordinate used to scan the change of partial charge, as shown in panel b. NBO analysis was used to calculate the partial charge at the MP2/6-311+G(d,p) level.

The distance between Ne2 on one His37 side chain and its neighboring He2 was selected as the reaction coordinate ζ. Among the three charged His37 residues, the one with the [γ1, γ2] of [180°, 60°] was selected to release a proton, as this His37 has an orientation similar to those found in the low pH condition X-ray structure.40 The free energy profiles for deprotonation and the representative snapshots along the reaction pathways are shown in Figure 5. The calculated PMFs was obtained by thermodynamic integration of the average force along the deprotonation pathway, where all free energy values are shown relative to the value of the lowest PMF calculated in each system (Figure Sα). The deprotonation curves are quite similar between the D3- and H3-models, but the presence of Cl− imposes significant perturbation on His37 deprotonation (describe later). In the D3-model, the minimum of the free energy profile is located at ζ = 1.024 Å, corresponding to the triply charged His37 cluster state (Figure Sα,b). Along with the increase of the Ne2−He2 bond until ζ = 1.30 Å, the PMF increases, showing that the proton moving from His37 to the neighboring water is energetically unfavorable (Figure Sα,b). At ζ = 1.30 Å, the proton is shared by His37 and the neighboring water, with the average Ne2(His37)···O(water) distance of 2.52 Å, where the proton is closer to the water (the averaged He2···O(water) distance is 1.24 Å). The newly formed hydronium is stabilized by the surrounding hydrogen-bond network, which extends to the waters at the C-terminal side (Figure Sb<sup>b</sup>). At high pH conditions, these interactions are interrupted as the Trp41 gate is closed. Meanwhile, one proton of the hydronium points to the neighboring water, and is ready to be released. Beyond this point, the PMF decreases. The barrier height for proton release is 6.35 kcal/mol. The additional proton on the hydronium is expected to release to the C-terminal side of the pore. At ζ = 1.5 Å, the proton transfer is complete (the averaged He2···O(water) distance is 1.06 Å), and the hydronium group is separated from His37 by three water molecules. The mechanism of proton release in the H3-model (Figure Sα,c−c′) is very similar to that in the D3-model, with the free energy barrier height of 6.63 kcal/mol, located at ζ = 1.36 Å: at this point, a hydronium is formed next to the deprotonated His37 (with the Ne2(His37)···O(water) and He2···O(water) distances of 2.53 and 1.22 Å, respectively).

Previously, we demonstrated that two alternative configurations (the D2- and H2-models) are capable of stabilizing the charges in the pore at neutral pH condition.34 Here we find that, under acidic conditions, the mechanism of proton release from the His37 cluster is much less dependent on its configuration (the D3- or H3-models). In other words, for a charged His37 side chain, which is ready to lose a proton, it is largely unaffected by the configuration of the remaining three His37 residues. Therefore, the interplay between His37 residues, the pH sensor of the channel, is pH dependent during the proton relay. The Trp41 gate and the membrane environment, on the other hand, are likely to play a role in modulating the rate of deprotonation and thus the conductance.

Different computational approaches have previously been used to explore the deprotonation from the charged His37 cluster in the M2 protein.34−36 In one recent paper from our group, the deprotonation from a single histidine and from the His···Trp motif in water was investigated with DFT-based MD simulations.34 For a single histidine, the free energy barriers were calculated to be 10−10.1 kcal/mol at the Nδ position, and 10.5−10.6 kcal/mol at the Ne position. A recent study from Voth and co-workers combined an empirical valence bond and a DFT-based QM/MM approach to study the proton release at different protonation states, reporting barrier heights of ~13 and ~10 kcal/mol for proton release from the triply and quadruply charged His37 cluster, respectively.36

When a single Trp residue is included in the calculation with a position resembling that in the protein, the barrier for the Nδ was found to be 9.1 kcal/mol, ~1 kcal/mol lower than that in the isolated histidine in water box,34 suggesting a role of the Trp41 gate in regulating the proton relay through His37, in addition to its known ability to generate an asymmetric flow.15 Specifically, charge transfer from His37 to the bound Trp41 through the cation−π interactions was witnessed, as assessed by the partial charge on each group (Figure 4b), consistently with the calculated electrostatic potential.23 Presumably, this charge delocalization decreases the energy cost to lose a proton.
In the present work, the energy cost for deprotonation in the membrane environment is ~6.5 kcal/mol, significantly lower than the one in the His···Trp motif or the isolated His. Apparently, the present work provides a more realistic model in studying proton transfer, where the backbone of the transmembrane domain of the protein, the Trp41 gate, and the explicit membrane environment have significant effects on modulating the dissociation of protons from His37. For the latter one, to be specific, the confined environment of the pore and the low dielectric constant of the membrane destabilize the charged cluster, and thus facilitate the deprotonation of the charged His37. By using the following equations and assuming $\Delta G = 0$ for simplicity between His37 and bulk water, the upper limit of the pKa of His37 in the membrane environment is estimated to be 4.7 in the D3-model and 5.0 in the H3-model, respectively, based on the major barrier heights for deprotonation in both models. Experimentally, the third pKa of the His37 cluster—when three of four residues are protonated—was determined as 6.3$^{37}$ and 4.9 ± 0.3$^{48}$ respectively, by using different protein constructs and lipid membrane compositions.

$$\Delta G = \Delta G^0 + 2.303RT \log \left( \frac{[\text{His}][H^+]}{[\text{HisH}^+]} \right)$$

$$\Delta G^0 = 2.303RT pK_a$$

The backbone conformations in both D3- and H3-models are very similar to those of the relatively compact protein structure at neutral pH. Without significant expansion of the backbone, we demonstrate that both models are able to release a proton with a moderate energy cost. The presence of the C-terminal amphipathic helix anchoring at the lipid interface region is likely to limit the dynamic of the C-terminal segment$^{39}$ compared to what seen for the isolated transmembrane helices from electron paramagnetic resonance (EPR) experiments$^{39,50}$ and the X-ray structure determined at low pH conditions.$^{40}$

**Role of Anion in the Pore.** Finally, we investigated the possibility of a role for Cl$^-$ or other halide anions in the proton relay mechanism through the pore. In electrophysiology studies, the proton conduction rate was found to be unaffected by the concentration of Cl$^-$ in both oocytes$^{40,41}$ and liposomes. This finding is in apparent contrast with the attractive force that a triply charged His37 tetrad can exert on a Cl$^-$ ion: assuming for simplicity that the rate of proton transfer is only controlled by electrostatics, Cl$^-$ ions should increase the proton conduction rate. In a recent study of the calcium-release activated calcium channel (CRAC), a cation-conducting channel with a 6 Å-wide pore and a high density of basic side chains, we observed that Cl$^-$ anions in the pore significantly increase the rate of permeation of Na$^+$ cations.$^{53}$ Therefore, we extended the proton transfer calculations to a model including a Cl$^-$ ion bound to the His37 tetrad, to explore its impact on the proton conduction rate.

As shown in the PMF calculations, the presence of Cl$^-$ makes the free energy cost of deprotonation much higher. Following the increasing N–H distance, the PMF keeps rising: at $\zeta = 1.36$ Å, where the major barrier is located in the previous PMFs, the free energy value is 10.5 kcal/mol, 3.9 kcal/mol higher than that in the H3-model (Figure 5a). This demonstrates that the presence of Cl$^-$ creates an unfavorable pathway for a proton to be released. During the simulations, the Cl$^-$ ion is located in the highly charged His37 cluster, and has stable interactions with the water in the cavity below, which is formed by the His37 and Trp41 clusters (Figure 5d). The arrangement of the hydrogen bonding network of water molecules in the cavity could compensate in principle the energy cost for bond breaking of the proton release. Nevertheless, the electrostatic attraction between the anion and the proton attenuates the proton diffusion out of the cavity to the C-terminal even though the Trp41 gate is occasionally open. Therefore, the presence of anions in the His37···Trp41

![Figure 5](image-url)
cavity is an unfavorable factor for proton release, which the M2 channel can circumvent by maintaining its pore free of Cl\(^{-}\) ions through its Asp44 residues, two helical turns away from His37. This is confirmed by mutations of Asp44, which significantly impact the proton conduction profile in a salt solution with a [Cl\(^{-}\)]/([SO\(_4\)\(^{2-}\)]) ratio approximately 2:1.\(^{41}\)

**CONCLUSIONS**

In this study, we explored the different configurations of the His37-Trp41 cluster in the M2 protein at low pH conditions, and the subsequent depolarization via QM/MM MD simulations. With regard to the conformation of the key residues, two alternative models (D3 and H3) converge to a single depolarization mechanism, and feature similar free energy profiles to release a proton, with a barrier height of ~6.5 kcal/mol. Therefore, the multiconfiguration model proposed in our previous work,\(^{14}\) includes depolarization as a mechanism not only for proton conduction, but also for conformational exchange connecting the multiple configurations of this protein. The confined region between His37 and Trp41 also appears to significantly increase the proton affinity of a water molecule compared to the bulk solution. Calculations with a bound Cl\(^{-}\) ion in the pore show that halide anions may contribute to stabilizing the triply charged histidine cluster, but at the same time impede the release of protons. We suggest that only those viral strains capable of maintaining halide anions at the same time impede the release of protons. We suggest that only those viral strains capable of maintaining halide anions at the same time impede the release of protons. We suggest that this may be a general feature of proton conduction in viral channels.

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**Notes**

The authors declare no competing financial interest.

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