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X-ray Crystal Structure of the Influenza A M2 Proton Channel S31N Mutant in Two Conformational States: An Open and Shut Case

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Abstract

The amantadine-resistant S31N mutant of the influenza A M2 proton channel has become prevalent in currently circulating viruses. Here, we have solved an X-ray crystal structure of M2(22–46) S31N that contains two distinct conformational states within its asymmetric unit. This structure reveals the mechanism of adamantane resistance in both conformational states of the M2 channel. In the Inward_{open} conformation, the mutant Asn31 side chain faces the channel pore and sterically blocks the adamantane binding site. In the Inward_{closed} conformation, Asn31 forms hydrogen bonds with carbonyls at the monomer–monomer interface, which twists the monomer helices and constricts the channel pore at the drug binding site. We also examine M2(19–49) WT and S31Nusing solution NMR spectroscopy and show that distribution of the two conformational states is dependent on both detergent choice and experimental pH.

Graphical Abstract

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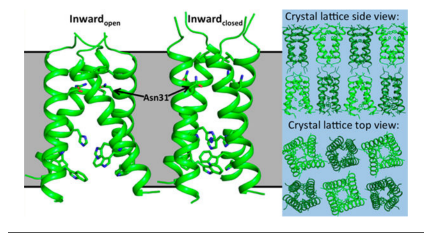
ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.9b02196. Supplemental figures and tables and detailed experimental methods (PDF)

Crystallographic data for M2 S31N mutant in two distinct conformational states (PDB code 6MJH) (CIF)

The authors declare no competing financial interest.



INTRODUCTION

Resistance to influenza antivirals is an ongoing concern. Most currently circulating strains of the virus are resistant to the adamantanes,^{1,2} which target the influenza A matrix 2 (M2) proton channel.^{3,4} Some strains are resistant to both neuraminidase (NA) inhibitors and M2 channel blockers.⁵ The most prevalent mutation in currently circulating influenza viruses is S31N,^{6–9} which is found in over 98% of adamantine-resistant strains of influenza.¹

The M2 channel is homotetrameric and proton selective.^{10–14} The minimum construct needed for proton conductance is the transmembrane (TM) domain,^{15,16} which contains gating residues His37 and Trp41.^{17,18} The TM domain of M2 is well-conserved relative to the rest of the protein,¹⁹ with only a small number of mutations resulting in fit viruses. Mutating one residue of the M2 monomer introduces four perturbations into the channel, so pore-lining mutations can have a large effect on conductance. The rate of proton conductance in M2 is finely tuned to allow for the acidification of the viral interior and uncoating of viral RNA before hemagglutinin-mediated membrane fusion occurs.^{20,21} As a result, only a few pore-lining mutations are observed in transmissible viruses, with the most notable being S31N, V27A, and L26F.^{2,22,23} These mutations have proton conductance rates and pH activation curves that resemble the wild type (WT) channel.²⁴ Because the S31N mutation is so prevalent, it is an important target for structural characterization and drug discovery.

Two conformational states of the M2 channel have been experimentally characterized: the Inward_{closed} and Inward_{open} states. Near neutral pH, M2 primarily adopts the Inward_{closed} state, in which the C-terminus of the channel is tightly packed and each monomer helix has a kink at residue Gly34. The Inward_{closed} state of M2 WT has been characterized using solid-state NMR,^{25,26} solution NMR,^{27,28} and X-ray crystallography,^{29,30} with good agreement between the structures in the TM domain. In the Inward_{open} conformation, the C-terminus of the channel opens and the pore-lining residues become exposed to bulk solvent. The Inward_{open} conformational state has been characterized primarily using X-ray crystallography.^{30–32} Bending or straightening of the monomer helices at Gly34 interconverts between these two conformations.^{33,34}In lower pH ranges, conformational

equilibrium between the Inward_{closed} and Inward_{open} states results in peak broadening in NMR spectra.^{28,33,34}

In addition to this conformational equilibrium, simultaneous equilibria exist between the possible charge states of the gatingHis37 tetrad (Figure 1). A number of NMR titration experiments have been carried out to determine the pK_a 'sassociated with successive protonation of the four His37 gating residues. There exists some variation between the reported values resulting from differences in the hydrophobic environment used for reconstitution.^{35–39} Nevertheless, in all of these studies, protonation to the +2 charge state occurs slightly above or near neutral pH, with protonation to the +3 state occurring within physiologically relevant low pH ranges (4–5).

Though the equilibrium between the two conformational states is influenced by pH (and thus the protonation state of the gating His37 tetrad), the energetic difference between the two conformations is relatively small, and so each conformational state can be isolated in a range of pH conditions. The Inward_{open} state is observed in conformational equilibrium with the Inward_{closed} state at low pH, as evidenced by NMR and SSNMR studies showing peak broadening^{27,33,34,40} and increased hydration of the pore at low pH.^{41,42} However, previous structural studies have also isolated the Inward_{open} state at intermediate and high pH.^{31,32,43} The Inward_{closed} state is primarily observed in neutral to high pH ranges but has also been structurally characterized bound to drugs and inhibitors at low pH.³⁰ At very low pH (<5), where all four gating histidines are protonated, the Inward_{open} state is the predominant species. Experimental conditions can also influence the conformational landscape of M2, with certain membrane mimetics favoring one conformational state over the other.

Interconversion between the Inward_{closed} and Inward_{open} conformations is necessary for proton conduction in the M2 channel. In a study using inside-out macro patches of *Xenopuslaevis* oocytes, IV curves were fitted to a transporter-like model for proton conduction.⁴⁴ The proton conduction mechanism begins with the channel in the Inward_{closed} conformation with the His37 tetrad in the +2 charge state. Protonation to the +3 charge state triggers the channel to convert to the Inward_{open} conformation, which opens the Trp41 gate and allows a proton to be released into the viral interior. This is followed by a "recycling" step where the deprotonated (+2) Inward_{open} channel converts back to the Inward_{closed} conformation. This mechanism is consistent with the channel's low rate of proton conductance relative to the rate at which protons come on and off the gating His37 residues. ⁴⁵ This is best viewed as a linked equilibrium in which multiple states are explicitly considered as a function of pH.

Here, we have solved a crystal structure of both the Inward_{closed} and Inward_{open} conformations of M2(22–46) S31N in the same unit cell at low pH (5.0). This motivated us to further examine the pH dependence of the two conformational states using solution NMR. In physiologically relevant pH ranges, we find that the stability of the Inward_{closed} conformation reaches a maximum near pH 6 and becomes slightly less favored at higher pH and dramatically less favored at lower pH.

RESULTS AND DISCUSSION

X-ray Crystal Structure of the S31N Mutant in Two Conformational States.

Here, we present an X-ray crystal structure of the drug-resistant M2(22-46) S31N mutant in two distinct conformational states at 2.06 Å resolution (PDB: 6MJH) (Figure 2). The asymmetric unit of these crystals contains two M2 tetramers, with one tetramer adopting the Inward_{open} conformation and the second tetramer adopting the Inward_{closed} conformation. These crystals have P_{2_1} symmetry and lattice dimensions *a*, *b*, *c* (Å) = 36.29, 36.15, 76.45; α , β , γ (deg) = 90, 103.6, 90 (Table S1). The P2₁ symmetry results in a lattice where the Nterminus of the Inward_{open} tetramer packs against the N-terminus of the Inward_{closed} tetramer. The two conformations of the channel observed here are consistent with previously solved crystal structures. The Inwardonen tetramer is well-ordered and tightly packed at its N-terminus but more open near the C-terminus.^{31,32} The N-terminus of the Inward_{closed} tetramer has a slightly wider opening relative to that of the Inward_{open} conformation, though this region of the pore is narrowed in comparison to previously solved structures of M2 WT. The C-terminus of the Inward_{closed} conformation is well packed and consistent with previously solved structures.^{29,30} We have previously observed a number of crystal structures in a wide range of unit cells and crystal packing environments for both the wild type M2 channel and now also the S31N mutant (Figure S1 and Table S2). The uniformity of the structures in the Inward_{open} and Inward_{closed} states provides strong evidence that these conformational states are intrinsic to the tetrameric structure rather than being a product of crystal packing.

In previous studies, only the Inward_{open} conformation of the S31N mutant could be isolated through crystallization. The Inward_{open} tetramer seen here is very similar to the previously solved crystal structure of the S31N mutant in this conformational state,⁴⁶ with an all-atom RMSD of 0.509 Å. Solution NMR screening of M2 in various detergents identified a maltose neopentyl glycol detergent MNG-3-C8 that stabilizes the Inward_{closed} conformation (Figure 3). Using this detergentas an additive in crystallization trials allowed us to characterize the M2 S31N mutant in the Inward_{closed} conformation for the first time at atomic resolution. The backbone of the Inward_{closed} tetramer of M2(22–46) S31N is in good agreement with our previously solved solution NMR structure of M2(19–49) S31N in complex with a small molecule inhibitor of this mutant (2LY0),²⁸ with a CaRMSD of 1.4 Å.

The networks of ordered waters within the M2 pore resemble those previously observed in crystal structures. In the Inward_{open} state, a continuous hydrogen-bonding network connects the waters below Asn31 to the gating His37 residues. The water network within the Inward_{closed} state closely resembles that of the X-ray crystal structure of M2 WT (PDB: 3LBW),²⁹ with four ordered waters forming hydrogen bonds to the four His37 δ -nitrogens below and two bridging waters positioned above.

Conformation of the Mutant Asn31 Side Chain.

In the crystal structure of M2(22–46) S31N reported here, the conformation of the resistance-conferring Asn31 side chain differs when the M2 tetramer is in the Inward_{open} state vs the Inward_{closed} state. In the Inward_{open} tetramer, the four Asn31 side chains face

toward the aqueous channel pore and form hydrogen bonds with symmetrically repeated Asn31 side chains from neighboring monomer units. The same side chain configuration was observed in our previously solved crystal structure of M2(22–46) S31N in the Inward_{open} state (PDB: 5C02).⁴⁶ However, when M2(22–46) S31N adopts the Inward_{closed} conformation in the absence of a drug, these Asn31 side chains face away from the channel pore and instead form hydrogen bonds with carbonyls at the tetramer's monomer–monomer interface. Interestingly, in a previous solution NMR structure of the M2 S31N mutant in complex with a small molecule inhibitor, the Asn31 side chain carboxamide was stabilized in the inward-facing conformation by interactions with a drug molecule.²⁸ In addition, Chou et al.⁴⁷ previously reported a solution NMR structure of M2(18–60) S31N in which the Asn31 side chain faces away from the channel's aqueous pore. However, Chou et al. model the Asn31 side chain as facing into the lipid bilayer. With the greater resolution now available using X-ray crystallography, we can see the Asn31 side chain carboxamide forming a bindentate interaction with a neighboring helix. In summary, it appears that the

conformation of Asn31 side chain can be modulated by either the conformational switch between Inward_{open} and Inward_{closed} state or drug binding. Consequently, understanding this conformational switch is critical for structure-based drug design.

Inward_{closed} Asn31 H-Bonding Motif in Other Structures.

In the M2(22–46) S31N Inward_{closed} tetramer, the Asn31 side chain engages in three hydrogen-bonding interactions. The nitrogen of the Asn31 carboxamide forms one intrahelix H-bond with the carbonyl from Val27 and a second H-bond with the Leu26 carbonyl from a neighboring monomer unit. The Asn31 side chain carbonyl oxygen also closely approaches the *a*-carbon of Ala30 from a neighboring monomer unit, with a distance of 3.3-3.5 Å between the two atoms, which is generally described as a C–H…O bond. The C–H…O hydrogen bonds are commonly observed in *a*-helical proteins between buried polar side chains and *a*-carbons, particularly in membrane proteins.^{48,49}

We interrogated the structural abundance of the Asn-to-backbone motif found in the Inward_{closed}M2 structure (Figure 4). We queried a nonredundant protein structural database of 20,383 structures for the motif, first starting with helical backbone superpositions (C, CA, N, O atoms) with RMSD 1.5 Å. Backbone residues 26–30 of chain A and 27–31 of chain B were queried with the program MASTER⁵⁰ for structural matches. All structural matches with an ASN at the position equivalent to 31B of M2 were then filtered by RMSD of the Asn side chain (CB, CG, OD1, ND2 atoms, RMSD 1 Å), using the program ProDy.⁵¹ A total of 114 close matches to the Asn motif were discovered, originating from a diverse set of membrane and soluble proteins (Table S3 contains a list of PDB accession codes). We conclude that this Asn hydrogen bonding motif is commonly observed to link *a*helices in a variety of proteins.

Examination of M2 Conformational States Using Solution NMR.

SSNMR studies show that the equilibrium between conformational forms of M2 depends markedly on the phospholipid composition of bilayers in which the protein is reconstituted. ^{33,34,52–54} Similarly, in solution NMR experiments, the conformation of the M2 channel is highly dependent on the micelle or bicelle composition.⁵⁵ Figure 3 shows NMR spectra for

M2(19-49) WT in a variety of detergents before and after addition of an inhibitor, which is known to stabilize the Inward_{closed} conformation of M2.33,56 We observe an equilibrium between both conformational states through examination of backbone peaks and also a single peak corresponding to Trp41 in the TM domain. The Inward_{closed} conformation has well-dispersed and sharp chemical shifts, while the peaks of the Inward_{open} conformation are broader and show less dispersion in the proton dimension. In our previous solution NMR studies,⁵⁷ we developed a simple, fast way to identify the two conformations through examination of the chemical shift of the Trp41 indole proton. The peak at ~11.0 ppm corresponds to the Inward_{closed} conformation, and the peak at ~10.2 ppm corresponds to the Inward_{open} conformation. For comparison, the average chemical shift of this Trp indole proton is 10.1 ppm in the BMRB (Biological Magnetic Resonance Data Bank). When M2(19-49) WT is solubilized in LPPG, only the Inward_{open} conformation is observed in both the absence and presence of amantadine. When M2(19-49) WT is solubilized in MNG-3-C8 detergent, only the Inward_{closed} conformation is observed both with and without amantadine. Using this detergent as an additive in the present study aided crystallization trials of M2(22-46) S31N in the Inward_{closed} state. In previous crystallization trials using octyl glucoside as a detergent additive, only the Inwardopen conformation of M2 S31N could be isolated.⁴⁶ We note that nonionic detergents appear to favor crystallization of M2 and speculate that MNG-3-C8 stabilizes the Inward_{closed} conformation because of its additional length relative to OG.

The Inward_{open} and Inward_{closed} conformational equilibrium of M2 is also highly dependent on pH. After screening a variety of detergents (Figure 3), we observed that C_{14} -betaine allows the two conformations to interexchange within the physiological pH range. In Figures 5 and 6, we selected the aromatic $C\delta$ –H δ peak of F48 as a probe to perform quantitative analysis of the conformational equilibrium because the F48 peaks from the two conformations clearly separate from each other and also do not overlap with any other peaks during the course of titration. In addition, F48 is near the end of TM, so it is sensitive to opening and closing of M2's C-terminus. The data clearly show that the stability of the Inward_{closed} state is relatively low at the highest pH, reaches a maximum as the pH is decreased to near pH 6, and then decreases as the pH is decreased further. The Inward_{closed} conformation becomes undetectable at pH 4.3 or lower. Similar behavior is seen for S31N, although the overall stability is somewhat lower in this particular micelle, C_{14} -betaine (Figure 6).

Figure 5 shows that the Inward_{closed} state of the WT peptide is somewhat less stable than the Inward_{open} state at high pH. Interestingly, the Inward_{closed} conformation becomes more stable as the pH is lowered to near 6.0, after which it drops off more precipitously with further decreasing pH, and the Inward_{closed} state is not detectable at pH 4.3 or lower. Similar results were seen for the S31N mutant (Figure 6). To explain this behavior, it is important to consider the protonation constants for both states. Thus, the data in the scheme in Figure 1 were used to determine the p K_a values for protonation of the Inward_{closed} and Inward_{closed} states (Figure 7, Scheme S1, and Table S4). This analysis indicated that the peak in stability near pH 6 is a direct result of energetic coupling between protonation and conformational change. The computed p K_a for the first two protonations of the Inward_{closed}state (6.5 ± 1.1 and 7.4 ± 1.2) are higher than the corresponding p K_a for the Inward_{open} state (approximately

 5.9 ± 0.1), resulting in stabilization of the Inward_{closed}state up to about pH 6 (because here the protons have a higher affinity for the Inward_{open} state and, hence, drive the equilibrium toward this state relative to high pH where both His37 residues are neutral). The p K_a for the final two protonations of the His37 tetrad are 4.4 ± 0.8 and <4. As a result, decreasing the pH below 6 leads to progressive destabilization, and the Inward_{closed} state is entirely destabilized at low pH.

CONCLUSIONS

Proteins in their native environment often have a number of possible conformational states. X-ray crystallography is a powerful technique for structural characterization because of its ability to isolate specific conformations and determine positions of individual atoms within particularly well-ordered crystals. However, conformational heterogeneity exists even within the ordered environment of the crystal lattice, both on the scale of individual side chain conformations and also on the scale of secondary and tertiary structure. Crystallization sometimes results in the isolation of multiple conformational states within a single crystal lattice. Examples of crystal structures containing more than one conformational state include structures of the *C. jejuni* CmeB multidrug efflux pump,⁵⁸ an RNA-dependent RNA polymerase from the rabbit hemorrhagic disease virus,¹⁷ and an mRNA capping enzyme.⁵⁹ These conformational states are often important for the function of the protein. In the case of the M2 channel, interconversion between the Inward_{closed} and Inward_{open} conformational states allows the channel to conduct protons through its narrow transmembrane pore.

Implications for Drug Resistance.

The conformation of the Asn31 side chain holds the key to adamantane resistance in the M2 channel. In the wild-type channel, the Ser31 side chain faces toward the center of the channel pore in both the Inward_{open} and Inward_{closed} conformational states, with the Ser31 hydroxyl engaging in a single intrahelix hydrogen bond with the carbonyl of Val27.^{29–32} As a result of this hydrogen bond, Ser31 engages in primarily hydrophobic interactions with the adamantyl cage of amantadine and rimantadine.^{30,31}

Based on the previously solved structure of S31N in the Inward_{open}conformation,⁴⁶the mechanism of drug resistance in the S31N mutant channel was hypothesized to be a steric block of the binding site as a result of replacing the Ser31 hydroxyl with the larger and more hydrophilic Asn31 carboxamide.⁴⁶ However, in this newly solved structure of M2(22–46) S31N in the Inward_{closed} state, Asn31 faces away from the center of the pore and forms hydrogen bonds at the monomer–monomer interface. The hydrogen bond between the Asn31 side chain and the Leu26 backbone carbonyl from a neighboring monomer twists each monomer helix at the channel's N-terminus. As a result of this twist, the Ala30 side chain that interacts with the monomer–monomer interface in the WT channel rotates toward the center of the pore in the S31N mutant channel. This constricts the channel at the adamantane binding site and elongates the pore at the N-terminus below the Val27 gate (Figure 8).

The shape of the pore in both conformational states is important for inhibition of the M2 channel. Both amantadine and rimantadine form stable complexes with both conformational

states of the M2 WT channel in X-ray crystal structures.^{30,31} This is consistent with inhibition of proton conduction upon addition of adamantanes at both high and low pH ranges;^{4,60} although the channel exists in conformational equilibrium at low pH,^{27,28,34} the adamantanes can bind to both states. The structures presented here will aid the design of new inhibitors to target both conformational states of the adamantane-resistant S31N mutant M2 channel.

Finally, this work provides an in-depth evaluation of the thermodynamically linked protonation and conformational equilibria in M2. The increased stability of the Inward_{closed} conformation as the pH is lowered from 8 to 6 directly reflects the well-known stability of partially protonated forms of M2, which have high pK_a 's for the first two protonation events on the gating His37 tetrad. In more stabilizing environments (e.g., the detergent MNG-3-C8 or certain lipid compositions),^{33,34,52–54} the small change in stability between pH 8and 6 does not lead to a large enough change in population of the states to be easily observed. However, as the pH is decreased below pH 6 the decrease in stability is quite pronounced and can be readily observed in many environments. The overall behavior reflects M2's ability to stabilize protons in the transmembrane pore, with special stabilization of the +2 state relative to either neutral or to +3 and +4 states. This can be understood in a mechanism whereby proton flux depends on the population of channel in the +2 state multiplied by the proton concentration.

METHODS

The M2(22–46) S31N construct was chemically synthesized as previously described⁶¹ and crystallized in the lipid cubic phase (LCP) with some modifications to the protocol described by Caffrey and Cherezov.⁶² Solution NMR was used to screen detergents for the stabilization of the Inward_{closed} conformational state; this identified an MNG detergent that was then used as an additive in crystallization trials. Crystals were diffracted at ALS 8.3.1 at a temperature of 100 K. Molecular replacement was carried out using structure 3LBW²⁹ as the starting model for the Inward_{closed} tetramer and 5JOO⁴³ as the starting model for the Inward_{closed} tetramer.

For solution NMR experiments, M2(19–49) WT and M2(19–49) S31N were doubly labeled with ¹⁵N and ¹³C and were prepared as previously described.²⁸ Spectra were recorded at 313 K on a Varian 600 MHz, a Bruker 600 MHz, or a Bruker 900 MHz spectrometer, as described in the Supporting Information.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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NOTE ADDED IN PROOF

In the course of reviewing the proofs we noticed that we had failed to reference two key papers concerning solid-state NMR characterization of M2: Mandala et al,⁶³ and Mandala et al.⁶⁴ These papers provide detailed structural and dynamic characterization of the two states of M2. Moreover they show that interconversion between the states occurs on approximately the same time scale as conductance.

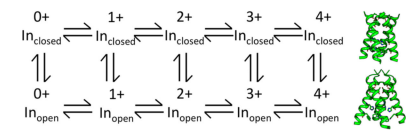


Figure 1.

Simultaneous equilibria between the His37 tetrad charge states and the two conformational states, $Inward_{closed}$ and $Inward_{open}$.

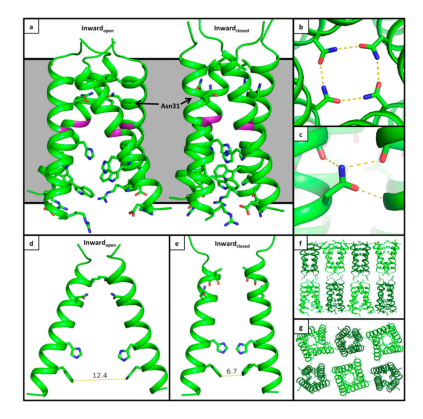


Figure 2.

X-ray crystal structure of M2(22–46) S31N in two conformational states. (a) The Inward_{open} conformation is at left, and the Inward_{closed} conformation is at right (PDB: 6MJH). The monomer helices in the Inward_{open} conformation are straight, and in the Inward_{closed} conformation they are kinked at Gly34 (shown in purple). As previously observed in structure 5C02, Asn31 faces the center of the pore and forms H-bonds with Asn31 side chains from neighboring monomers in the Inward_{open} conformation. In the Inward_{closed} conformation, Asn31 faces away from the pore and forms H-bonds with carbonyls at the monomer–monomer interface. (b) Top-down view showing hydrogen-bonding networks formed by Asn31 in the Inward_{open} conformation. (c) Side view of the hydrogen-bonding network formed by Asn31 in the Inward_{closed} conformation. (d) Side view of the Inward_{open} conformation, only chains F and H shown. (e) Side view of the Inward_{closed} conformation, in light green and the Inward_{closed} conformation in dark green. (g) Top-down view of crystal lattice.

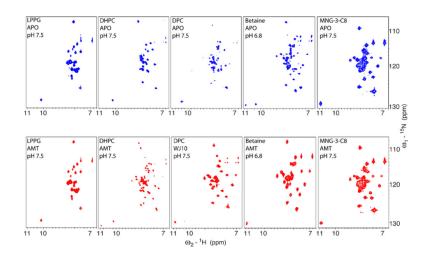


Figure 3.

Conformational equilibrium between the Inward_{open} and Inward_{closed} conformations for M2(19–49) WT is dependent on detergent choice in 2D NMR experiments. Solution NMR spectra of M2(19–49) WT in multiple detergents, without inhibitors (blue, top), and with inhibitors (red, bottom). The Trp41 indole proton peak (bottom left corner of each spectrum) can be used to distinguish the conformational states; a peak with chemical shift of ~11.0 ppm corresponds to the Inward_{closed} conformation. From left to right: In LPPG, M2 stays in the Inward_{open} conformation in absence and in the presence of amantadine. In DHPC, the presence of amantadine shifts a small portion of the protein in the Inward_{open} conformation. In DPC, the inhibitor WJ10 almost fully shifts the Inward_{open} to the Inward_{closed} conformation. In C₁₄-betaine, both conformations are observed in the absence of any drugs, and addition of amantadine fully shifts the protein to the Inward_{closed} conformation. In MNG-3-C8, the protein stays in the Inward_{closed} conformation.

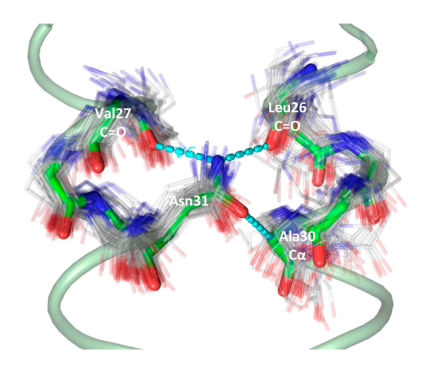


Figure 4.

Inward_{closed} Asn31 hydrogen bonding motif is commonly observed in the PDB. The Inward_{closed} S31N structure backbone from chains A and B is shown with light green ribbon, with green sticks highlighting the search motif and hydrogen bonds shown as dashed cyan lines. Matching PDB structures are shown as gray lines.

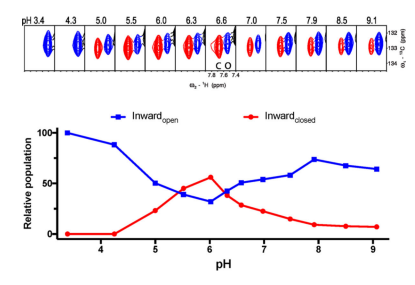


Figure 5.

F48 C δ -H δ peak shows conformational equilibration between the Inward_{open}(blue) and Inward_{closed} (red) states of M2(19–49) WT as a function of pH. Top: Only the Inward_{open} state is observed at pH 3.4 and 4.3. In the physiologically relevant pH range (5.0–7.5), both conformational states are observed. The peak of the Inward_{closed} conformation (red) gains intensity at pH 5.0, reaches a maximum at pH 6.0, and then decreases at higher pH. The data are normalized to the intensity observed at pH 3.4, and sum to slightly less than 100%, possibly due to a third state that is not detected due to chemical exchange or dynamics. Bottom: Fit of peak intensity vs pH for both conformational states. The intensity profile of the Inward_{open} conformation (blue) has a U-like shape with a minimum at pH ~6.0. The channel primarily adopts the Inward_{open} conformation at both very low and very high pH.

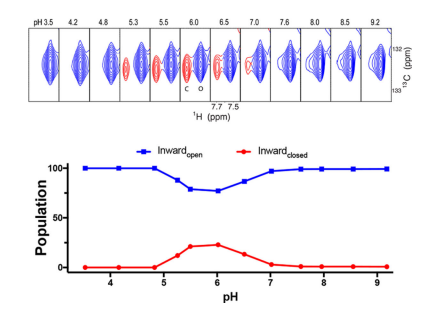


Figure 6.

F48 C δ -H δ peak of M2(19–49) S31N shows conformational interchange between the Inward_{open} and Inward_{closed} states of M2(19–49) S31N as a function of pH. Top: Similar to WT as seen in Figure 5, we show the F48 C δ -H δ peak from NMR spectra of the M2(19–49) S31N construct in a range of pH conditions. Bottom: Peak intensity vs pH for both conformational states. The intensity profile of the Inward_{open} conformation (blue) has a U-like shape with a minimum at pH ~6.0.

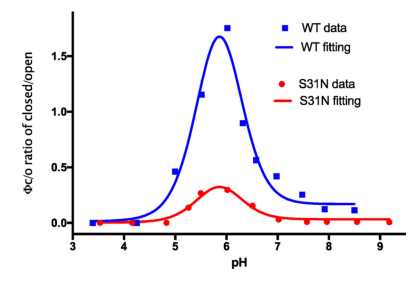


Figure 7.

Fitting of experimental data as a function of pH, giving R square values 0.96 and 0.95 for WT and S31N, respectively, and producing the three pKa's for the Inwardclosed state (pK1: 6.4 ± 1.1 , pK2: 7.4 ± 1.2 and pK3: 4.4 ± 0.8) and one pKa (5.9 ± 0.1) for the Inwardopen state. We set the pK4 as a constant 3, but the fit was insensitive to its value, so long as it was set less than 4.0.

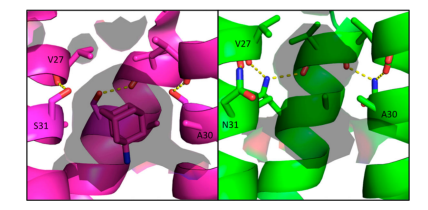


Figure 8.

Solvent-accessible surface area inside the M2 pore at the adamantane binding site. Amantadine-bound WT structure in the Inward_{closed} conformation 6BKK (left, magenta) vs S31N Inward_{closed} conformation (green, right); solvent-accessible surface area (calculated using a 1.4 Å probe) is shown as a transparent surface. In the WT channel, Ser31 forms an intrahelix hydrogen bond and partially faces the pore. In the S31N mutant, Asn31 faces the monomer–monomer interface, twisting the monomer helices such that Ala30 faces the center of the pore. This constricts and elongates the pore at the amantadine binding site for the S31N mutant M2 channel.