

Structure and Function of the Influenza A M2 Proton Channel

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Received May 25, 2009; Revised Manuscript Received June 17, 2009

ABSTRACT: The M2 protein of influenza A viruses forms a tetrameric pH-activated proton-selective channel that is targeted by the amantadine class of antiviral drugs. Its ion channel function has been extensively studied by electrophysiology and mutagenesis; however, the molecular mechanism of proton transport is still elusive, and the mechanism of inhibition by amantadine is controversial. We review the functional data on proton channel activity, molecular dynamics simulations of the proton conduction mechanism, and high-resolution structural and dynamical information of this membrane protein in lipid bilayers and lipid-mimetic detergents. These studies indicate that elucidation of the structural basis of M2 channel activity and inhibition requires thorough examination of the complex dynamics and conformational plasticity of the protein in different lipid bilayers and lipid-mimetic environments.

FUNCTION OF THE M2 PROTON CHANNEL OF INFLUENZA A VIRUSES

The M2 protein of influenza A and B viruses forms tetrameric proton channels that are important for the viral life cycle. After the virus enters the infected cell by endocytosis, the M2 proton channel opens in response to the low pH of the endosome, allowing proton flux into the virus, which triggers the dissociation of the viral RNA from the matrix proteins and the fusion of the viral and endosomal membranes. These events release the viral RNA to the cytoplasm for replication by the host cell (1). In a later stage of virus replication, the M2 protein maintains the high pH of the trans-Golgi network and prevents premature conformational changes of hemagglutinin in viruses with a high pH optimum of hemagglutinin-induced fusion (2).

The influenza A M2 (AM2) protein contains a short N-terminal periplasmic domain, a transmembrane (TM) domain, and a C-terminal cytoplasmic tail (Figure 1). It is one of the smallest ion channel proteins and thus an excellent system for elucidating the structure–function relation of ion channels. Extensive mutagenesis, electrophysiology (3, 4), and sedimentation equilibrium experiments (5) have been conducted to characterize the function and stability of AM2 (for recent reviews, see refs 6 and 7). The AM2 proton channel is also the target of the amantadine class of drugs, one of only two anti-influenza drugs currently available. However, the efficacy of amantadine dropped by two orders of magnitude between 2002 and 2007, although the 2008 seasonal flu strains were largely sensitive to amantadine. The resistance mainly resulted from the S31N mutation in the M2 TM domain (8). Thus, elucidating the mechanism of amantadine inhibition of AM2 has great public health relevance.

Recently, several high-resolution structural studies were reported that shed light on the structural basis of AM2 proton conductance and inhibition. In this work, we summarize the main functional data of AM2 and high-resolution structural information available about the TM domain to promote future investigations of this intriguing and far from understood membrane protein.

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Proton Transport, Selectivity, and Activation. The proton conductance of AM2 has been well studied both in vivo and in vitro since its discovery as an ion channel (3). The gold standard of M2 functional assays is the replication of live viruses in infected cells, followed by the “silver standard” of whole-cell conductance data in *Xenopus laevis* oocytes. The wild-type protein is activated below pH 6.2, and its current is sensitive to amantadine. The single-channel activity under a -130 mV voltage is ~ 0.16 $\mu\text{A}/\text{ng}$ of protein at pH 6.2 for the A/Udorn/72 strain of M2 (4), which corresponds to 12 aA per channel, a conductance of ~ 100 aS, or ~ 100 protons per second. Extraction of single-channel currents requires quantification of the amount of protein and the assumption that all proteins form open channels or the knowledge of the fraction of open channels.

For in vitro liposome assays, several additional factors such as the orientation of the protein in the membrane, the size and buffering capacity of the liposomes, and the fluxes of other monovalent ions (Na^+ and K^+) can affect the measured conductance. One of the most reliable liposome assays employed an intravesicular pH-sensitive dye for measuring proton fluxes and yielded a single-channel current of 2.7 aA at pH 5.7 and 18 °C (conductance of 29 aS) for full-length M2 in DMPC/DMPS vesicles (9). These values agree well with the oocyte result (4). Several other liposome assays gave more divergent values because of different experimental designs and protein constructs (10–12). However, the general consensus is that the single-channel conductance of M2 ranges from tens of attosiemens to a few hundred attosiemens, indicating low transport rates of ten to several hundred protons per second at mildly acidic pH. It is thought that the low currents are a result of the low physiological proton concentration (9), although the idea that M2 may have some characteristics of transporters has also been proposed (13).

There is general consensus that AM2 is selective for protons over Na^+ and K^+ by a factor of $1.5\text{--}3.0 \times 10^6$ (9, 14, 15), and that the increased proton current at low exterior pH (pH_{out}) is due to true activation of the channel rather than mere mass action, since an increased pH_{out} stops the outward current of acidified cells (15, 16). The site of activation is His₃₇, the only residue in the

MSLLTEVETP¹⁰ IRNEWGRCRN²⁰ DSSDPLVVA³⁰A SIIGILHLIL⁴⁰ WLDRLFFKC⁵⁰
 IYRFFEHLGK⁶⁰ RGPSTEGVPE⁷⁰ SMREEYRKEQ⁸⁰ QSAVDADDSH⁹⁰ FVSIELE

TM domain

FIGURE 1: Amino acid sequence of influenza A/Udorn/72 M2 protein. The TM domain containing the crucial His₃₇ and Trp₄₁ residues (red) is underlined.

TM domain that can be protonated within the relevant pH range. Mutation of His₃₇ to either Gly or Glu results in channels that are no longer modulated by pH_{out} (17). Solid-state NMR (SSNMR) studies of His₃₇ in TM peptide M2(22–46) suggest that it is the third protonation at pH 6.3 that opens the channel (18).

His₃₇ Selectivity Filter and Trp₄₁ Gate in the TM Domain. His₃₇ and Trp₄₁ are the two most conserved residues in the M2 protein of influenza A strains (19) and are essential for channel function. His₃₇ protonation is responsible for the activation and selectivity of the channel. The wild-type (WT) A/Udorn/72 M2 protein exhibits a 50-fold conductance difference between pH_{out} 4.5 and 8.2. In contrast, many His₃₇ mutants, while still forming channels, are toxic to oocytes and have pH-independent currents, indicating nonspecific conductance of other cations (17). These mutants can be neither activated effectively by low pH_{out} (H37G, H37S, and H37T) (20) nor inactivated by high pH_{out} (H37G, H37E, and H37A) (17). The mutant channels are not proton-selective and are only partially and reversibly blocked by amantadine (20), in contrast to the WT protein (16). Although the H37 mutants are still tetrameric in oocytes (20), analytical ultracentrifugation (AUC) experiments indicate that H37A and H37F mutants are ~3.1 and ~1.8 kcal/mol less stable, respectively, than the WT peptide in DPC micelles (21).

When a high concentration of imidazole buffer is added to the bathing solution, proton-selective conductance and gating are partially restored to the His₃₇ mutants (20). This chemical rescue is specific to imidazole and does not require interaction with Trp₄₁. Thus, His₃₇ was proposed to be an intermediate proton acceptor and donor, consistent with SSNMR titration data of the TM peptide (18).

The M2 proton channel responds exclusively to external pH: low pH_{out} activates the channel, and high pH_{out} closes the channel, regardless of the interior pH. Mutagenesis data indicate that Trp₄₁ acts as the gate in concert with His₃₇ (22). At high pH_{out} and low pH_{in}, the WT protein does not exhibit outward current while W41A, -C, and -F mutants do, indicating loss of gating by the mutants (22). Further, intracellular Cu²⁺ inhibited the open W41A channel but not the WT channel, indicating that Trp₄₁ shields His₃₇ from coordination by Cu²⁺ (22, 23). These data suggest that the indole rings adopt distinct conformations at different pH_{out} values to obstruct or open the pore. Resonance Raman data indicate that the Trp₄₁ conformational change must be small to keep its environmental hydrophobicity unchanged (24). Concerted conformational changes seem necessary for His₃₇ to act as the selectivity filter and Trp₄₁ as the gate. Compared to His₃₇, the two Trp₄₁ mutants that abolish gating (W41A and W41F) (22) actually have enhanced tetramer stability in DPC micelles (21), indicating that this residue has evolved for function rather than stability.

Mechanism of Proton Transport. Molecular dynamics (MD) simulations have been used extensively to study the M2 proton transport mechanism. The basic premise is that protons hop along a water wire via the Grotthuss mechanism, but the water wire is broken at the narrow His₃₇ constriction (and possibly other locations along the channel). Thus, the mechanism

of passage of a proton through the His₃₇ tetrad is the central question. Two main models have been proposed: the gating model and the shuttle model. In the gating model, protonation of the His₃₇ tetrad at low pH causes pore opening by electrostatic repulsion, so that a continuous water wire is formed (25). Conformational changes of His₃₇ and possibly Trp₄₁ are necessary in this model. MD simulations suggest that a 60° change of the His₃₇ χ_2 angle may be sufficient to open the pore (26), and the excess proton may form a Zundel complex with two water molecules on the extra- and intracellular side of the histidine rings (27, 28). Early MD simulations postulated various charged states of His₃₇ for channel activation (29). However, with SSNMR data indicating that the third protonation of the His₃₇ tetrad is responsible for channel opening (18), recent simulations confirmed that the +3 state has the lowest proton permeation free energy (30). The channel water molecules that transport the protons were calculated to have 3-fold lower diffusion coefficients than bulk water (27, 31).

The second proton transport model is the shuttle model, in which the imidazole nitrogen facing the extracellular side binds one proton to form a biprotonated intermediate and then the other nitrogen releases the proton on the intracellular side and returns the imidazole to the neutral form (32). The initial state is then recovered by tautomerization or ring flip of the imidazole. This model explains the low proton transport rate and low-pH saturation of the channel, since both would be associated with the limiting rate of His₃₇ side chain torsional motion or tautomerization.

Simulations of the protein conformation at different pH values also shed light on the mechanism of proton transport. One proposal is that in addition to His₃₇, Val₂₇ may act as a secondary gate that breaks the water wire, especially in the presence of amantadine (33). The two gates may not be static but instead may move synchronously in response to His₃₇ protonation: at high pH, the Val₂₇ region opens while the His/Trp region narrows, and the opposite is found at low pH (13). Thus, instead of equating high pH with the closed state and vice versa, these simulations suggest a more complex view of an open_{out}–closed_{in} conformation at high pH and closed_{out}–open_{in} conformation at low pH. This model reconciles the different helix orientations and packing in different high-resolution structures (34–37).

Inhibition of the AM2 Channel. Three classes of M2 inhibitors have been identified so far: the adamantane analogues (38, 39), the spirene-containing compounds (40), and Cu²⁺ ions. The FDA-approved amantadine achieves its antiviral effect by inhibiting the M2 proton channel activity (3): 100 μ M amantadine nearly irreversibly blocks >90% of the channel activity of A/Udorn/72 (H3N2) M2 in several minutes at both acidic and neutral pH (16, 41, 42). For some other influenza strains, the amantadine block is moderately stronger (50%–2-fold) at neutral pH than at low pH. The Hill coefficient of the amantadine block is 1, consistent with the scenario that a single drug inhibits each functional channel (16).

Naturally occurring mutations in the TM domain that confer amantadine resistance include those at L26, V27, A30, S31, and G34, all N-terminal to G34 (8, 38, 43, 44) (Figure 1). Extensive whole-cell conductance studies have been conducted on these mutants. S31N, the most common resistant mutant (8), has an IC₅₀ of 199.9 μ M, more than 10 times higher than that of the WT (IC₅₀ = 16 μ M) (41). The resistance is not due to altered channel activity, as S31N has specific activity and acid activation very similar to those of the WT (4, 42). L26F has a ~35%

higher channel activity than S31N but is more sensitive to amantadine (42). G34E has a 3-fold higher specific activity than S31N and is ~5-fold less sensitive to amantadine than S31N (4). V27 mutants have mixed phenotypes: V27A and V27S are amantadine-resistant, V27T is amantadine-sensitive, and all V27 mutants have significantly increased (~3-fold) channel activities versus that of WT (4). In contrast, the A30T mutant has a greatly reduced but still detectable activity, while the A30P mutant is inactive (4). Thus, residue 30 appears to be important for maintaining a functional structure of the protein. Among the resistant mutants, S31N was found by thiol–disulfide exchange experiments to be stable in both bilayers and micelles (42), suggesting a reason for its dominance in current flu strains that is independent of the selective pressure of amantadine. The stability and similar channel activity of S31N compared to that of the WT protein argue against the model in which the S31N variant is resistant to amantadine by destabilizing the closed state and stabilizing the open state (45). The latter hypothesis was developed on the basis of a comparison of the NMR line widths of S31N and WT proteins in the presence of rimantadine. However, since S31N does not bind rimantadine, the more relevant comparison should be between the S31N spectra and the apo WT protein spectra.

In the C-terminal region of the TM domain, L38F, W41A, and D44A mutants incorporated into influenza viruses exhibited growth similar to that of the WT protein in infected cells, and the growth is sensitive to amantadine (41) despite the fact that the W41A mutant does not gate properly (22). Electric recordings in oocytes indicate that the IC_{50} of D44A is 15.8 μ M, indistinguishable from that of the WT protein. Thus, no amantadine-resistant mutations have been found C-terminal to G34, implicating the segment between L26 and G34 to be the amantadine-binding region (41). Further, when residues 6–18 of the amantadine-insensitive BM2 were replaced with AM2 residues 24–36, the chimeric protein showed amantadine-sensitive currents in oocytes (41), indicating that the C-terminus is not the pharmacologically relevant region associated with amantadine's antiviral effect. In contrast, a recent liposome proton flux assay of M2(18–60) found the D44A mutant to be resistant to rimantadine while the S31A mutant to be sensitive, supporting a C-terminal binding model (34, 45). This liposome assay result is in serious contradiction both with the chimeric protein result and with the finding from the “gold standard” assay that replication of live virus bearing the D44A mutation is inhibited by rimantadine (41).

The second class of M2 inhibitors consists of spirene-containing compounds, initially discovered from a growth impairment assay in which the M2 toxicity to yeast cells due to its proton channel activity is reversed by inhibitors (40). The first compound, BL-1743 {2-[3-azaspiro(5,5)undecanol]-2-imidazole}, has an inhibition rate, a Hill coefficient, and spectra of resistance similar to those of amantadine (46). However, BL-1743 inhibition is reversible (~12 min), while amantadine block is irreversible on the experimental time scale (~30 min). BL-1743 also differs from amantadine in its interaction with I35: the I35T mutant is resistant to BL-1743 but is reversibly inhibited by amantadine (46). The cause of this difference is suggested by a recent SSNMR study of the analogous spirene-piperidine compound, 3-azaspiro(5,5)undecane hydrochloride (IC_{50} = 0.92 μ M) (47). This compound reduces the level of dynamic disorder in the G34-I35 backbone of AM2, suggesting that spirene-amine inhibitors bind

more extensively and strongly with the TM helices than the adamantane inhibitors.

HIGH-RESOLUTION STRUCTURE OF THE AM2 TM DOMAIN

Orientation of the TM Helices in Lipid Bilayers. The TM peptide of AM2 has been a prime target for high-resolution structure investigation by a number of biophysical techniques. Both global structural information such as helix orientation and site-specific information of key residues have been obtained. The orientation of the TM helices in lipid bilayers has been extensively studied by SSNMR, EPR, and IR techniques. The majority of SSNMR studies measured orientation-dependent 15 N tensors on glass plate-aligned M2(22–46) (37, 48–51) and to a small extent on aligned full-length M2 (52, 53). From one-dimensional 15 N chemical shift anisotropies (CSA), Cross and co-workers obtained a helix tilt angle of 33–37° for M2(22–46) in DMPC bilayers (49, 50) and 33° in DOPC bilayers. Two-dimensional (2D) NMR spectra correlating 15 N– 1 H dipolar couplings and 15 N CSAs gave more precise orientational constraints, including both the tilt angle (τ) and the rotation angles (ρ) of individual residues. From these 2D spectra, the tilt angle of apo M2(22–46) in DMPC bilayers was refined to 38° (51), while the rotation angles around the helix axis are consistent with the pore- or lipid-facing positions of residues inferred from functional data [Protein Data Bank (PDB) entry 1NYJ].

Prior to 2001, the oriented NMR studies were conducted on peptides that were first mixed with lipids in organic solvents before hydration. These samples were not pH controlled and may be acidic due to the TFA salt associated with the peptide. After 2001, the oriented membranes were prepared by an aqueous protocol in which M2 was reconstituted into lipid bilayers by detergent dialysis at a controlled pH before alignment on glass plates. The amantadine-bound M2(22–46) and the full-length M2 were studied in this way. In DMPC/DMPG lipid bilayers, the 2D spectra of uniformly 15 N-labeled full-length M2 at pH 8.0 showed a TM helix with a 25° tilt, which is smaller than the tilt of the TM peptide alone, and an 80° tilted helix, which is attributed to an amphipathic helix C-terminal to the TM domain (52). Hydrogen–deuterium (H–D) exchange of full-length M2 showed that the TM helix underwent fast H–D exchange, consistent with its participation in the formation of an aqueous pore, while the 80°-tilted helix resisted H–D exchange, indicating that it is shielded from water. Full resonance assignment is necessary to elucidate the orientation and depth of this C-terminal amphipathic helix.

When amantadine binds, M2(22–46) exhibits a pronounced kink at G34 in DMPC bilayers, with a tilt angle of 31° for the N-terminal segment and 20° for the C-terminal segment (PDB entry 2H95) (Figure 5a) (48). While it is tempting to conclude that amantadine caused this kink, the bound peptide spectra were obtained from aqueous prepared samples at pH 8.8, under which condition there are no comparable apo peptide data. It is thus possible that the kink may already be present in the apo state, especially given the known propensity of Gly to perturb α -helices. Indeed, other high-resolution structures of AM2 suggest a G34 kink already in the apo state. For example, MAS NMR data of DLPC-bound M2 suggest a kink of 5° for the bound peptide and 8° for the apo peptide (36), while the crystal structure of apo M2 at neutral pH shows a distribution of helix kink angles (35).

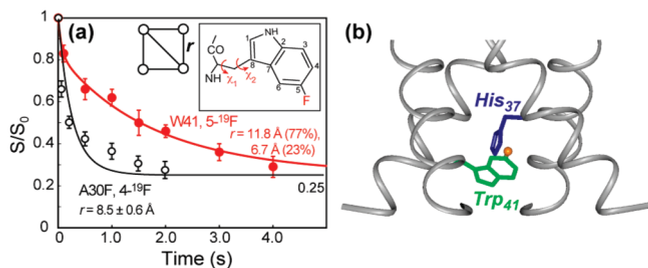


FIGURE 2: (a) Interhelical ¹⁹F–¹⁹F spin diffusion data of 5-¹⁹F-Trp₄₁ WT M2 (red) and 4-¹⁹F A30F M2 (black) in DMPC bilayers (59, 67). The best-fit distances confirm the tetrameric state of the peptide and constrain the Trp₄₁ side chain (χ_1 and χ_2) torsion angles (inset). (b) Trp₄₁ t90 and His₃₇ t-160 rotamers proposed from the distance data.

Direct comparison of the orientation of amantadine-free and amantadine-bound M2(22–46) prepared under the same conditions was reported recently (54). Using unoriented DLPC vesicles and dialysis-reconstituted M2(22–46), Hong and co-workers determined the helix orientations by exploiting the uniaxial diffusion of the helical bundles around the bilayer normal. The N-terminal half of the TM segment was found to have a tilt angle of 35° in the apo state and a slightly larger tilt angle of 38° in the amantadine-bound state (55) (Figure 5d), indicating that the drug slightly enlarges the pore vestibule.

The TM helix orientation is also sensitive to membrane thickness. EPR spectra of N-terminally nitroxide-labeled M2(22–46) reconstituted into DLPC, DMPC, DOPC, and POPC bilayers showed the least dipolar broadening in the thinnest DLPC membrane and the strongest broadening in the thickest POPC membrane (56). The broadening is attributed to shorter interhelical distances as the peptide decreases its tilt angle in the thicker membrane. The orientational change was detected directly in SSNMR spectra of unoriented M2(22–46). The TM helices exhibited a tilt angle of 35° in DLPC bilayers but a much smaller angle of 26° in the thicker POPC bilayers (54). These results are consistent with AM2 adjusting its TM helix orientation to minimize the hydrophobic mismatch with the lipid bilayer, as also observed for other membrane peptides (57, 58). These results revise an earlier conclusion that the M2 helix orientation is intrinsic to the peptide and unaffected by the membrane (50), made on the basis of the fortuitously similar orientations of the peptide in DMPC and DOPC bilayers because of the similar thickness of these two bilayers.

The SSNMR and EPR orientational studies do not detect evidence of rotational angle change of M2 by pH: both the organic solvent-prepared samples, which likely have acidic pH, and the aqueous samples at high pH gave the same rotation angles. These results are consistent with the functional requirement for a hydrophilic pore. Distances between the Trp₄₁ indole 5-¹⁹F of neighboring helices were also found to be the same (11.8 Å) at pH 7.5 and 4.5 (Figure 2), consistent with the lack of ρ change (59). Thus, it is surprising that a recent IR study of M2(22–46) in DMPC bilayers reported a one-residue (100°) rotation of the helices between low and high pH (60). The ρ angle of the low-pH samples was found to agree well with early SSNMR data (37, 51), while that of the high-pH samples differs by ~100°. When the ρ angles were combined with the +3 protonated His₃₇ tetrad, MD simulations found that the rotated helices reduce the level of exposure of His₃₇ to water at high pH, thus rationalizing channel closure (61). The ρ angles were

extracted from diagonal line widths of 2D IR spectra, which reflect the electrostatic environment of the ¹³C=¹⁸O groups, the main contributor of which is thought to be the water contact. However, the 2D line shape analysis requires the distinction of homogeneous and inhomogeneous line widths, consideration of population relaxation times, and consideration of water-independent electrostatic effects such as the hydrogen bond network of the peptide. Thus, the origin of the diagonal line width change is complex. In addition, linear Fourier transform infrared (FT-IR) dichroism data for the high-pH samples reported in this work do not show a clear sinusoidal dependence on residue number, casting further doubt on the high-pH ρ angle result (60).

Tetramer Stability and Packing. In mammalian cells and oocytes, the functional state of M2 is tetrameric, as shown by sucrose gradient, chemical cross-linking (62), and conductance experiments (5). In detergents and lipid bilayers, the TM peptide retains the tetrameric state under a wide range of conditions, and the monomer–tetramer equilibrium is readily shifted by membrane thickness, pH, amantadine binding, cholesterol, and amino acid sequence. The environmental effects on the free energy of tetramer association have been the focus of study using AUC in micelles (21, 63) and thiol disulfide exchange in lipid vesicles (64). It was shown that M2(19–46) tetramers are increasingly stabilized in the direction of detergents \ll DLPC < DMPC < POPC. Among lipid bilayers, the hydrophobic thickness-matched bilayers produce the most stable tetramers (64), but even the thinnest bilayer (DLPC) induces 100-fold tighter tetramer association than detergents. Both amantadine and cholesterol shift the equilibrium to tetramers (64, 65). Compared to external environmental factors, the sequence requirement for tetramer formation is much more lenient: Ala and Phe mutations of a number of pore residues retain stable M2(22–46) tetramers in DPC micelles, with the exception of the functionally essential His₃₇ (21). However, more stable tetramers are often less functional tetramers, as manifested by their weakened ability to bind amantadine (63). For example, the most amantadine-resistant S31N mutant forms 0.4 kcal/mol more stable tetramers than the WT protein (42). This inverse relationship between stability and function suggests that the multiple M2 functions (proton conduction, gating, and inhibitor binding) may require conformational flexibility, which cannot be provided by highly stable mutants locked in the wrong conformational minima.

Direct distance constraints for M2 tetramers in lipid bilayers were recently reported on the basis of ¹⁹F spin diffusion SSNMR experiments that probe ¹⁹F–¹⁹F distances in the ~10 Å range (66, 67). So far, four interhelical ¹⁹F–¹⁹F distances have been reported for M2(22–46) in DMPC bilayers. On 4-¹⁹F-labeled A30F M2, the ¹⁹F spin diffusion equilibrium value of 1/4 directly proved the tetrameric state, and the decay trajectory yielded a distance of 8.5 ± 0.6 Å between nearest neighbor helices (Figure 2a). This distance rules out helix tilt angles smaller than 20° and constrains the Phe χ_1 angle. The ¹⁹F NMR data of V27F suggest incomplete tetramer formation (partial dimers), which is consistent with the reduced stability of this mutant seen in AUC experiments (21). The L38F NMR data also indicated distance heterogeneities (59).

His₃₇ and Trp₄₁ Side Chain Conformation. A semiquantitative ¹³C–¹⁵N distance between His₃₇ N π and Trp₄₁ C γ was measured on DMPC-bound M2(22–46) at pH 7 and 38 °C using SSNMR REDOR experiments (37). Significant dipolar dephasing was observed that corresponded to a minimal coupling of

63 ± 12 Hz (37) or a maximum distance of 3.9 Å. The limiting nature of this result stems from the fact that the measurements were taken at a physiological temperature where M2(22–46) undergoes uniaxial diffusion around the bilayer normal, which averages the dipolar coupling (50, 54). Scanning the possible His and Trp rotameric states indicates that this C–N distance must be between helices i and $i + 1$ and suggested that His₃₇ adopts the t-160 ($\chi_1 = 180^\circ$, and $\chi_2 = -160^\circ$) rotamer while Trp₄₁ adopts the t-105 rotamer (PDB entry 1NYJ) (37). Subsequent reexamination of the conformational space of the two residues indicates that other possible solutions also exist: His₃₇ can adopt the t-160 or t60 rotamer, while Trp₄₁ can adopt the t-105 or t90 rotamer. Density functional theory calculations and MD simulations suggest that the most stable combination is (t60, t90) (26).

¹⁹F spin diffusion NMR experiments provided a direct distance constraint on the Trp₄₁ side chain conformation. The i to $i + 1$ distance between 5-¹⁹F-Trp₄₁ of M2(22–46) was measured to be mainly 11.8 Å at pH 7.5 (59) (Figure 2a), and the value does not change significantly with amantadine binding and pH. When the helix orientation is fixed to 35° and the channel diameter is fixed to 10.2 Å between G34 C α , the only Trp₄₁ rotamer that is consistent with the ¹⁹F–¹⁹F distance is t90, which is consistent with the MD prediction (26). Combined with the measured His₃₇–Trp₄₁ C–N distance, the (t-160, t90) rotamer pair was proposed for the closed state of His₃₇ and Trp₄₁ (59). This conformation suggests that it may not be Trp₄₁ alone but rather the cation– π interaction between His₃₇ and Trp₄₁ that is responsible for the gating and shielding of His₃₇ from intracellular ions (Figure 2b).

A third Trp₄₁ rotamer ($\chi_1 = -100^\circ$, $\chi_2 = 100^\circ$) was proposed for the high-pH state of M2 on the basis of analysis of ¹⁹F NMR line shapes under the combined effects of ¹⁹F CSA and ¹⁹F–¹⁹F interhelical dipolar couplings (68). However, the presence of intermediate time scale broadening of the spectra and the lack of information about the orientational angles between the ¹⁹F chemical shift tensor and the F–F dipolar tensors make this analysis a severely underdetermined problem. The resulting rotamers are also inconsistent with any of the computational predictions (26).

His₃₇ Protonation State. The charged state of His₃₇ gives crucial information about the activation of the M2 proton channel. The pK_a of His₃₇ in micelle-bound M2(22–46) was measured by ¹H solution NMR to be 6.8 for the monomer and 6.4 for the tetramer in the absence of drug (65). Amantadine decreases the tetramer pK_a to 5.8. UV resonance Raman experiments on M2(22–46) in POPE/POPS membranes found a pK_a of 5.7 based on the intensity of the 1407 cm⁻¹ band (24). This imidazole band has the same intensity in liposomes and in SDS micelles at pH 4.0, indicating that all four His residues are protonated at pH 4.0 (24).

By far the most definitive study of His₃₇ protonation states came from ¹⁵N SSNMR experiments on M2(22–46) in DMPC/DMPG bilayers by Cross and co-workers (18), who monitored the imidazole N δ 1 and N ϵ 2 ¹⁵N isotropic chemical shifts as a function of pH. Two neutral tautomers were found at pH 8.6, with the unprotonated nitrogen exhibiting a large isotropic shift of ~230 ppm. Between pH 8.0 and 7.0, two +1 dimer species, His–HisH⁺, were found. On the basis of the relatively large ¹⁵N line widths in this pH range, which suggest intermediate time scale motion, low-barrier hydrogen bonds between imidazole and imidazolium within each dimer were proposed. Below pH 6.5 and 5.0, the third and fourth His residues are protonated. Quantifica-

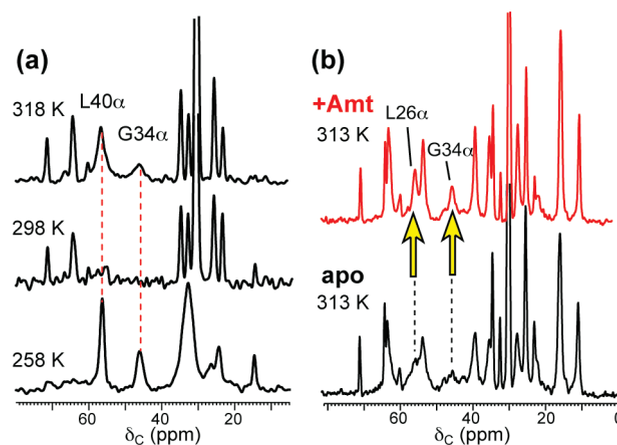


FIGURE 3: ¹³C MAS spectra of DLPC-bound M2(22–46). (a) Variable-temperature ¹³C MAS spectra indicate large-amplitude intermediate time scale motion at ambient temperature (54). (b) Amantadine narrows the line widths of most residues at 313 K (55).

tion of the spectral intensities yielded four pK_a values at 8.2, 8.2, 6.3, and <5.0. Thus, the first two histidines titrate cooperatively at higher pH than the aqueous pK_a of His (6.0). Since the +1 dimers already exist at neutral pH, they are proposed to form a histidine-locked state that occludes the pore (18). The binding of the third proton below pH 6.3 was proposed to break the 2-fold symmetry of the +1 dimers and thus activates the channel.

Amantadine binding changes the titration curve of the His₃₇ tetrad (69). The 230 ppm unprotonated ¹⁵N peak persisted to a much lower pH of 6.0 in the presence of the drug. Intensity analysis shows that the bound peptide has only one pK_a near 5.4. The nature of this apparently highly cooperative protonation event is still unknown. In addition, the ¹⁵N peak associated with the His–HisH⁺ dimer is no longer observed in the amantadine-bound peptide, suggesting that the neutral tautomers directly form biprotonated histidines in the presence of amantadine. Further studies are necessary to elucidate the amantadine-induced changes in His₃₇ protonation.

Extensive Conformational Dynamics of AM2. The M2 homotetramer exhibits extensive conformational dynamics in lipid bilayers and lipid-mimetic detergents, which are also well documented by SSNMR and solution NMR spectra. Membrane-bound apo M2(22–46) shows dynamically broadened lines in both unoriented vesicle spectra (36, 55) and oriented spectra (70) at ambient temperature in a range of model membranes: DLPC, DMPC/DMPG, and POPC. The fact that the line broadening is mainly dynamic in origin rather than due to static conformational distributions is proved by the temperature dependence of the line widths. The broadest lines are observed between 298 and 263 K (54), above and below which the lines narrow (Figure 3a). Acidic pH broadens the lines compared to neutral pH for both membrane-bound M2(22–46) (70) and detergent-bound M2(18–60) (34). In all solvent environments and for both M2(22–46) and M2(18–60), amantadine noticeably narrows the line widths (Figure 3b). The line broadening or narrowing occurs for all residues; thus, it reflects the dynamic property of the whole protein rather than segmental motion.

The origin of the broad line widths and their narrowing by amantadine have been examined in detail by SSNMR (54, 71). The main motion accounting for the broad line width is the whole-body uniaxial diffusion of the tetramers around the bilayer

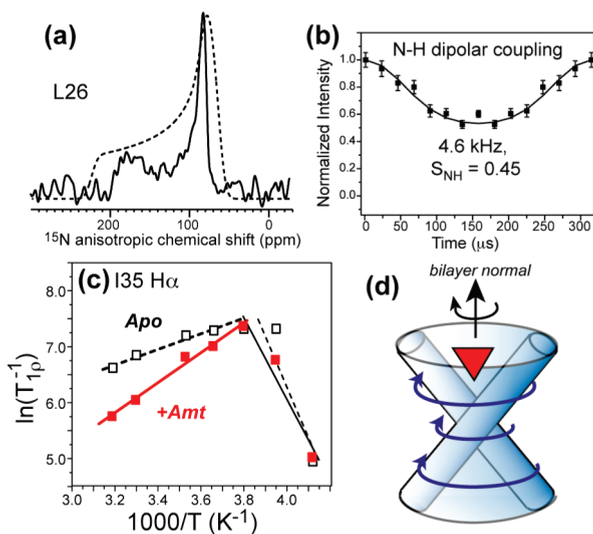


FIGURE 4: Uniaxial diffusion of M2(22–46) helical bundles around the bilayer normal. (a) ^{15}N CSA of L26 is uniaxially averaged from the rigid-limit pattern (dashed line) (54). (b) L26 N–H dipolar coupling is reduced from the rigid-limit coupling. (c) Amantadine decreases ^1H $T_{1\rho}$ relaxation rates at high temperatures, indicating it accelerates protein motion (71). (d) Schematic of M2 uniaxial diffusion. Amantadine binding creates better packed tetramers, thus accelerating rotational diffusion.

normal, as shown by motionally averaged ^2H , ^{15}N , and ^{13}C NMR line shapes (Figure 4a,b) (54). The correlation time of this uniaxial diffusion has been estimated from relaxation studies to be $\sim 3 \mu\text{s}$ (71) for DLPC-bound apo M2(22–46) at 313 K, which is very close to the time scale of the radio frequency pulses and nuclear spin interactions ($2\text{--}4 \mu\text{s}$), thus causing intermediate time scale broadening. Amantadine binding reduces the correlation time by 2–3-fold at 313 K (71) (Figure 4c), thus avoiding exchange broadening and narrowing the NMR lines. The higher diffusion rates suggest that amantadine promotes the formation of more homogeneous helical bundles (Figure 4d), which is consistent with the increased tetramer stability observed in thiol disulfide exchange experiments (64). The NMR relaxation data also indicate that excess amantadine in the bilayer increases the membrane viscosity. Since protein rotational diffusion is generally sensitive to membrane viscosity, cholesterol-rich virus-mimetic membranes immobilize this diffusion and were found to suppress exchange broadening and give rise to high-resolution NMR spectra over a wide temperature range (72).

Whether the whole-body uniaxial diffusion has functional importance is not known. If the viral envelope is in a liquid-ordered phase, then M2 is most likely immobilized, but if the virus envelope contains significant liquid-disordered domains, then the fact that M2 is poorly incorporated into raftlike microdomains (73) should promote uniaxial diffusion in the functional state. Most spectroscopic studies so far were conducted on M2 bound to non-cholesterol-containing membranes or bound to detergents, which favor the whole-body motion. Thus, site-specific conformational dynamics have not been separated from the whole-body motion. In this light, the larger line widths of M2 at low pH are at least partly due to global motional changes, and their interpretation in the context of pH activation needs to be considered carefully (34, 70).

Amantadine Binding in High-Resolution Structures. Two atomic-resolution structures of TM-containing portions of AM2 were recently determined using X-ray crystallography and solu-

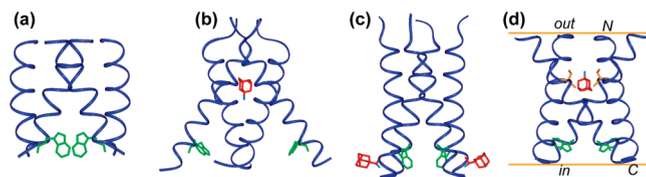


FIGURE 5: High-resolution structures of the TM domain of AM2. (a) Amantadine-bound orientational structure of M2(22–46) in DMPC bilayers at pH 8.8 from oriented-membrane SSNMR (PDB entry 2H95) (48). The drug binding site was not directly studied but was implicitly assumed to be in the pore at the N-terminal side. (b) Amantadine-bound crystal structure of M2(22–46) in OG at pH 5.3 (PDB entry 3C9J) (35). (c) Rimantadine-bound solution NMR structure of M2(18–60) in DHPC micelles at pH 7.5 (PDB entry 2RLF) (34). Only the TM part is shown. (d) Amantadine-bound structure of M2(22–46) in DLPC bilayers at pH 7.5 from MAS SSNMR (PDB entry 2KAD) (36). The amantadine position is a hypothesis based on the chemical shift perturbation of S31. The bilayer planes are drawn for reference. In each structure, two of the Trp₄₁ indole rings (green) are shown, and the drug molecules are colored red. In panel c, only two rimantadine molecules are shown for the sake of clarity.

tion NMR, which shed light on the pH activation of the channel but led to two different models of M2 inhibition (74).

The crystal structures were determined on M2(22–46) in octyl β -D-glucopyranoside (OG) at pH 7.3 in the absence of amantadine (PDB entry 2BKD) (2.0 Å resolution) and at pH 5.3 in the presence of amantadine (PDB entry 3C9J) (3.5 Å resolution) (35). In both structures, the N-terminal half of the TM domain has a tilt angle of $\sim 35^\circ$, consistent with SSNMR data (51, 54). The orientation of the C-terminal half of the TM helix depends on the pH: at low pH, the helices are uniformly straight and diverge toward the C-terminus, creating an open cavity, but at pH 7.3, two of the four helices bend and create a small pore in the His₃₇/Trp₄₁ region. The amantadine-bound structure, determined with a 1.3:1 protein:amantadine molar ratio, shows a single drug molecule in the pore, surrounded by residues that confer drug resistance (Figure 5b). Thus, the amantadine binding site and stoichiometry agree with electrophysiology data (16), and the crystal structure supports an occlusion model for channel inhibition.

The solution NMR structure was determined on rimantadine-bound M2(18–60) in DHPC micelles at pH 7.5 (PDB entry 2RLF) (34). The sample contains 0.75 mM protein, 40 mM rimantadine, and 300 mM DHPC; thus, the drug is in 50-fold excess over the protein or 200-fold excess over the channel. The structure showed the TM helix to be tilted by $\sim 15^\circ$ from the bilayer normal and the C-terminal amphipathic helix to be approximately perpendicular to the TM helix. Low pH broadens the line widths and increases the millisecond motion of Trp₄₁. Four protein-rimantadine NOEs were found to lipid-facing residues L40, I42, L43, and R45 near the C-terminus of the TM helix, with four rimantadine molecules per tetramer (Figure 5c). Since this external binding site is inconsistent with the large body of mutagenesis data that place drug-resistant mutations at pore-facing residues at the N-terminus, the authors proposed that rimantadine inhibits the channel by stabilizing its closed state and making the channel harder to open, whereas drug-resistant mutations destabilize the closed channel and make it easier to open. However, the allosteric inhibition model is inconsistent with the one-drug-per-channel stoichiometry of amantadine (16) and also contradicts virus replication assays and whole cell conductance data that showed D44A and R45A mutants to be amantadine-sensitive (41). The drug–protein

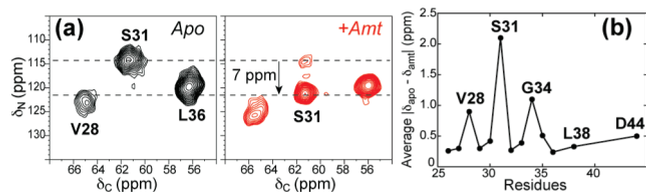


FIGURE 6: SSNMR evidence for amantadine binding near S31. (a) 2D ^{15}N - ^{13}C correlation spectra of DLPC-bound M2(22–46) in the absence (black) and presence (red) of amantadine (36). (b) Chemical shift perturbation by amantadine in bilayer-bound M2(22–46).

NOEs involve residues in the headgroup region of the micelle, exactly where the amphiphilic rimantadine is located based on paramagnetic relaxation NMR data (75, 76). At the high concentration of 40 mM, rimantadine constitutes 12 mol% of the micelle, making the surface binding site a likely secondary lipid binding site rather than the true inhibition site. Given the abundant reorientational dynamics of the peptide and amantadine discussed above (55, 71), the lack of NOEs between the N-terminal pore residues and rimantadine could arise from motion on unfavorable time scales that broadens the NMR signals and/or makes the NOE intensity vanish, or from micelle-induced helix orientation and packing changes that reject rimantadine in the pore. Thus, while this work shows the presence of rimantadine on the surface of the protein, they do not provide strong evidence against a pharmacologically relevant binding site in the pore.

It is unlikely for the different protein lengths to be the cause for the drug binding site difference, since the TM peptide is capable of binding amantadine, folding into appropriate tetramers, and having at least 50% of the proton conductance of full-length M2 (77).

A bilayer- and amantadine-bound structure of M2(22–46) has been reported from MAS SSNMR experiments (36, 55). The experiments were conducted at a peptide:amantadine:DLPC molar ratio of 1:8:15. The main conformational constraints came from ^{13}C and ^{15}N isotropic chemical shifts (CS) measured at 243 K where the protein dynamics is frozen. Comparison of the apo and bound peptide chemical shifts indicated that the M2(22–46) conformation is most perturbed by amantadine at S31, with a large ^{15}N CS change of 7 ppm, strongly suggesting drug binding near S31 (Figure 6a). The CS perturbation is also high for G34 and V28, while the C-terminal L38 and D44 are minimally affected (Figure 6b). The chemical shift-predicted ϕ and ψ angles for the apo and bound monomers also show that the apo peptide has a slightly larger kink than the amantadine-bound peptide. The monomer structure was combined with the helix orientation and interhelical ^{19}F - ^{19}F distances to lead to a tetramer structure for M2(22–46) at pH 7.5 (PDB entry 2KAD). The tetramer has a larger N-terminal vestibule than the other structures and differs in the Trp₄₁ conformation (36) (Figure 5d). SSNMR studies also found that amantadine slightly perturbs the side chain conformations of V27 and V28 (78).

FUTURE DIRECTIONS

While a large amount of biochemical and structural information has been obtained for the M2 proton channel, many open questions remain about this fascinating multifunctional protein. So far, the most complete three-dimensional structures were determined in detergents and micelles, which are only partial

mimics of lipid bilayers. Given the plasticity of the protein, high-resolution structural studies in lipid bilayers that mimic the virus envelope will be very valuable. Most structural studies so far used partial constructs of the full-length protein; thus, future studies of the full-length protein will clearly be desirable, especially for understanding the role of the C-terminal amphipathic helix, which was found to be able to affect ion channel function by interacting with the TM helix (79). Elucidating how M2 changes its conformation and dynamics between high pH and low pH is central to understanding its activation and will require atomic-resolution information of His₃₇ and Trp₄₁ conformation as a function of pH. Clearly, protein–drug distances in lipid bilayers will be essential for resolving the controversy of M2 inhibition. Finally, the ultimate public health benefit of this research is to develop new antiviral drugs to target the dominant mutants of influenza A viruses to prevent future influenza pandemics. Achieving this end will require a combination of pharmacological, biochemical, and structural studies.

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