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Solid-State NMR Investigation of the Conformation, Proton Conduction, and Hydration of the Influenza B Virus M2 Transmembrane Proton Channel

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Abstract

Together with the influenza A virus, influenza B virus causes seasonal flu epidemics. The M2 protein of influenza B (BM2) forms a tetrameric proton-conducting channel that is important for the virus lifecycle. BM2 shares little sequence homology with AM2, except for a conserved HxxxW motif in the transmembrane (TM) domain. Unlike AM2, no antiviral drugs have been developed to block the BM2 channel. To elucidate the proton-conduction mechanism of BM2 and to facilitate the development of BM2 inhibitors, we have employed solid-state NMR spectroscopy to investigate the conformation, dynamics and hydration of the BM2 TM domain in lipid bilayers. BM2 adopts an α-helical conformation in lipid membranes. At physiological temperature and low pH, the proton-selective residue, His19, shows relatively narrow ¹⁵N chemical exchange peaks for the imidazole nitrogens, indicating fast proton shuttling that interconverts cationic and neutral histidines. Importantly, pH-dependent ¹⁵N chemical shifts indicate that His19 retains the neutral population to much lower pH than His37 in AM2, indicating larger acid-dissociation constants or lower pKₐ’s. We attribute these dynamical and equilibrium differences to the presence of a second titratable histidine, His27, which may increase the proton-dissociation rate of His19. 2D ¹H-¹³C correlation spectra probing water ¹H polarization transfer to the peptide indicates that the BM2 channel becomes much more hydrated at low pH than at high pH, particularly at Ser12, indicating that the pore-facing serine residues in BM2 mediate proton relay to the proton-selective histidine.

Graphical abstract
Introduction

While influenza viruses are best known for causing devastating flu pandemics in history, they exert just as much economic and social burden by causing seasonal flu epidemics. Two types of influenza viruses, A and B, are responsible for the seasonal flu. On average influenza A infections are more common in the seasonal flu\(^1\), but the influenza B virus becomes dominant in the spring months, and in some seasons can account for up to 50% of infections (data available from CDC FluView, [www.cdc.gov/flu/weekly/\(^2\)].

The currently approved adamantane class of antiviral drugs works only against the influenza A virus by binding to its M2 protein (AM2) and blocking the conduction of protons into the virus, a process that is important for the virus lifecycle\(^3\)–\(^7\). However, these drugs have no effect against the influenza B M2 protein (BM2), which also has proton-channel activity\(^8\). The lack of inhibition is not surprising: BM2 and AM2 share little sequence homology, with only a HxxxW motif conserved in their respective TM domains (Fig. 1a)\(^9\). The histidine is responsible for the acid activation and proton selectivity of the channel, whereas the tryptophan is responsible for channel gating\(^8\),\(^10\). Sequence alignment of the two TM domains show that the BM2 channel pore is significantly more polar than the AM2 channel, with multiple Ser residues lining the pore, suggesting the reason for the inability of the hydrophobic adamantane to inhibit BM2\(^11\). A high-resolution solution NMR structure of the BM2 TM domain has been determined using BM2(1–33) bound to DHPC micelles\(^12\). The result shows a coiled-coil tetramer with three Ser residues (S9, S12, S16) lining the pore and a bulky Phe residue (F5) lying at the N-terminal entrance. The structure of the cytoplasmic domain was also determined in the same study using a BM2(26–109) construct. Liposome assays indicate that BM2(1–33) exhibits rimantadine-insensitive proton conductance, and in *Escherichia coli* lipid extracts, the single-channel conductance of BM2(1–33) is 2-fold higher than the conductance of AM2(18–60)\(^12\). Since AM2(18–60) has the same specific activity as full-length AM2\(^13\), and whole-cell voltage-clamp assays of full-length AM2 and BM2 indicate similar inward proton currents\(^8\), these data together indicate that BM2(1–33) encompasses the full channel activity of the intact protein. Similar to AM2, BM2’s channel activity is important for virus uncoating\(^8\),\(^14\) and for preventing hemagglutinin from prematurely adopting a low-pH conformation during transport to the cell surface\(^8\),\(^15\),\(^16\). The
cytoplasmic domain of BM2 carries out the separate function of incorporation of M1 and viral ribonucleoprotein complexes at the virion budding site during virus assembly\textsuperscript{17,18}. While AM2 and BM2 channels are functional analogs, various differences exist in their pH- and voltage-dependent proton conduction. The BM2 TM domain contains a second histidine (H27), C-terminal to the gating Trp23, whose mutation to Ala reduces the inward H\textsuperscript{+} current by \textasciitilde26\%. Whole-cell electrophysiological assays showed that BM2 has significant outward current under positive voltages, when the external pH (pH\textsubscript{out}) is high and the internal pH (pH\textsubscript{in}) is low, conditions under which AM2 shows negligible outward current\textsuperscript{8}. To understand the basis for these functional differences, and to facilitate the design of antiviral drugs against influenza B infection, it is important to determine the structure of the BM2 TM domain in lipid bilayers and to investigate the mechanisms of action of the functional residues, His19 and Trp23.

A large amount of structural and dynamical information about the HxxxW motif has been obtained on the AM2 protein. The His37 residue in AM2 is now known to shuttle protons into the virion by proton exchange with water.\textsuperscript{15}N chemical exchange peaks have been observed in both TM peptide and cytoplasmic-containing constructs of AM2 and in the S31N mutant of AM2\textsuperscript{19–22}. The linewidths of the exchange peaks differ, indicating that proton transfer rates are sensitive to protein structural differences and electrostatic effects. This proton shuttling involves ring reorientations\textsuperscript{23} and $\tau$ - $\pi$ tautomerization\textsuperscript{22}. Measured imidazole $^1$H chemical shifts at low and high temperatures suggest hydrogen bonding between histidine and water\textsuperscript{7}; in addition His-His hydrogen bonding has also been proposed\textsuperscript{21,24}. His37-Trp41 cation-$\pi$ interaction at low pH has been suggested by $^{13}$C-$^{19}$F\textsuperscript{25} and $^{13}$C-$^{15}$N\textsuperscript{26} distance experiments and UV resonance Raman experiments\textsuperscript{27}, and has been proposed to regulate proton release from His to the C-terminal water molecules\textsuperscript{28,29}, in agreement with the gating function of Trp\textsuperscript{10}. Computed free-energy profiles of proton transport and 2D IR data indicate that proton conduction at low pH is facilitated by an expanded channel width and increased hydration and solvent dynamics\textsuperscript{30,31}. The relation between channel hydration and His-mediated proton shuttling is of special interest, because the BM2 TM domain has a more polar pore-lining surface than AM2, suggesting that the histidine-mediated proton transfer dynamics might differ from that of AM2.

Solid-state NMR spectroscopy is a powerful and versatile tool for studying the structure and dynamics of membrane proteins in lipid bilayers. Increasing evidence from studies of many membrane peptides and proteins indicates that the structures of small oligomeric membrane proteins are more susceptible to changes by the membrane environment compared to large membrane protein complexes and machineries\textsuperscript{32–37}, and some of these structural changes are functionally relevant. Solid-state NMR based structural and dynamical studies of membrane proteins in phospholipid bilayers instead of detergents are therefore essential for obtaining biologically authentic mechanistic information. In this study, we investigate the backbone conformation of BM2(1–33) using chemical shifts of site-specifically labeled peptides, then focus on the structures and dynamics of the proton-selective His19. We also investigate channel hydration using water-protein $^1$H-$^{13}$C 2D correlation experiments. We show that in cholesterol-containing eukaryotic-mimetic membranes, BM2(1–33) is stably $\alpha$-helical at high and low pH and in two different membranes examined, and His19 exhibits a

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number of cationic and tautomeric states that in totality resemble those seen for His37 in AM2 but that have a shifted pH dependence. $^{15}$N NMR spectra of His19 yielded four pK$_a$'s, which are about one pH unit lower than the pK$_a$'s of His37 in AM2. The BM2 channel hydration is both pH- and membrane-dependent: negatively charged membranes increased the pore hydration compared to neutral membranes, and the pH dependence of BM2 hydration is larger than that of AM2. These results suggest the structural reasons for some of the functional differences between these two viral proton channels, and how the amino acid sequence affects the histidine-mediated proton conduction.

**Materials and Methods**

**Synthesis of isotopically labeled BM2(1–33)**

A peptide corresponding to residues 1–33 (MLEPFQILS ICSFILSALH FMAWTIGHLN QIKR) of the B/Maryland/1/2001 strain of influenza virus was synthesized using Fmoc solid-phase peptide synthesis protocols as described previously. Briefly, ChemMatrix Rink Amide resin was used for the peptide synthesis. For the coupling step, 5 equivalents of amino acid, 5 equivalents of HCTU, and 10 equivalents of diisopropylamine were used and the reaction was carried out in DMF for 5 minutes at 75°C. For the Fmoc deprotection step, 5% piperazine in DMF plus 0.1 M HOBt were used, and the reaction was similarly carried out in DMF for 5 minutes at 75°C. Final peptide cleavage from the resin was carried out in a cocktail that consists of 95% trifluoroacetic acid, 2.5% triisopropylsilane, and 2.5% H$_2$O for 2.5 hours at room temperature. Cleaved peptide was precipitated in cold ether and purified by preparative HPLC (Vydac C4 Column). Uniformly $^{13}$C, $^{15}$N-labeled amino acids (Sigma-Aldrich and Cambridge Isotope Laboratories) were introduced at positions S12, A17, His19 and G26.

**Membrane Sample Preparation**

The BM2(1–33) peptide was reconstituted into two lipid membranes, POPC : POPG : cholesterol (PC/PG/Chol) at molar ratios of 60% : 20% : 20%, and POPC : POPE : sphingomyelin : cholesterol (VM+) at equimolar concentrations. Both lipid mixtures mimic eukaryotic membranes by including cholesterol, but the former contains the anionic lipid POPG, since this mixture is commonly used in M2 proton conductance assays and membrane scission assays. Anionic lipids are also present in the plasma membrane and the virus lipid envelope at ~15 mol%, but usually in the form of phosphatidylserine instead of phosphatidylglycerol. The neutral VM+ lipid mixture allows us to compare the structure of BM2 with that of AM2 in the same membrane. The lipids were mixed together in chloroform, then the solvent was removed by nitrogen gas. The peptide was dissolved in 1 ml of 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP) or 2,2,2-trifluoroethanol (TFE), then mixed with the lipids. The bulk solvent was removed again by nitrogen gas, then the powder was placed under vacuum at room temperature overnight followed by lyophilization for 2–3 hours to fully remove fluorinated solvents. The resulting lipid-peptide mixture was suspended in buffer, vortexed and freeze-thawed 8–10 times to create uniform vesicles. The proteoliposome solution was spun at 50,000 rpm at 4°C for 4 hours to obtain homogenous membrane pellets. The pellets were allowed to dry slowly in a desiccator to 40–50%
hydration by mass, then transferred into 4 mm or 3.2 mm magic-angle-spinning (MAS) rotors for solid-state NMR experiments.

Seven BM2 samples were prepared at different pH in the two membranes. For the PC/PG/Chol membrane, three samples were prepared at pH 7.5 (10 mM Tris, 1 mM EDTA, 0.1 mM Na$_3$N), pH 5.5 and pH 4.5 (10 mM citric acid/citrate, 1 mM EDTA, 0.1 mM Na$_3$N) and the peptide : lipid molar ratio was 1 : 15. For the VM+ membrane, four samples were prepared at pH 6.5 (20 mM Bis-Tris, 2 mM EDTA, 0.2 mM Na$_3$N), pH 5.5 (10 mM citric acid/citrate, 1 mM EDTA, 0.1 mM Na$_3$N), pH 4.5 (10 mM citric acid/citrate, 1 mM EDTA, 0.1 mM Na$_3$N) and pH 4.0 (20 mM citric acid/citrate, 2 mM EDTA, 0.2 mM Na$_3$N). The peptide : lipid molar ratio was 1 : 13.3. The pH of the membrane samples was measured at two time points during sample preparation: after freeze-thawing to create homogeneous vesicle solutions, and after ultracentrifugation, where the supernatant pH was measured. The measured pH was within 0.1 pH units of the desired value.

Membrane samples for dynamic nuclear polarization (DNP) experiments

BM2(1–33) was reconstituted into a deuterated lipid mixture denoted d$_{31}$-VM$^+$, which contains d$_{31}$-POPC, d$_{31}$-POPE and cholesterol at a molar ratio of 40% : 40% : 20%.$^{41}$ The partially deuterated phospholipids were used to increase the efficiency of polarization transfer from the radical to the peptide. About 1 mg of BM2 was dissolved in TFE, then mixed with the d$_{31}$-VM$^+$ lipid mixture in chloroform at a peptide/lipid molar ratio of 1 : 22.5. The organic solvents were removed under nitrogen gas, then the mixture was lyophilized overnight. The peptide-lipid mixture was suspended in a 10 mM pH 5.5 citrate buffer and centrifuged at 40,000 rpm at 4°C overnight to obtain a homogeneous pellet, which was equilibrated to a hydration level of ~40 wt%. A small amount of a stock DNP solution, d$_8$-glycerol : D$_2$O : H$_2$O (60 : 30 : 10 by volume) containing 10 mM of the biradical AMUPol was titrated into the membrane pellet$^{41}$. The pellet was vortexed to distribute the radical uniformly, and D$_2$O was added to reach a D$_2$O : H$_2$O volume ratio of 3 : 1. The sample was then allowed to equilibrate again to ~40% hydration, then packed into a 3.2 mm sapphire rotor and sealed with a silica plug for DNP experiments at 117 K.

Solid-state NMR experiments

Solid-state NMR experiments were carried out on Bruker 400 MHz (9.4 Tesla) and 800 MHz (18.8 Tesla) NMR spectrometers using 4 mm or 3.2 mm $^1$H/$^{13}$C/$^{15}$N MAS probes. Typical radiofrequency (rf) field strengths were 71–83 kHz for $^1$H, 62.5–71 kHz for $^{13}$C and 45–50 kHz for $^{15}$N. Chemical shifts were referenced to the CH$_2$ signal of adamantane at 38.48 ppm on the tetramethylsilane (TMS) scale for $^{13}$C, the amide signal of N-acetylvaline at 122.0 ppm on the liquid ammonia scale for $^{15}$N, and the H$_{\gamma}$ signal of the lipid phosphocholine head group at 3.26 ppm on the TMS scale for $^1$H.

1D $^{13}$C and $^{15}$N cross-polarization (CP) MAS spectra were measured at 313-243 K. To extract His19 pK$_{a}$'s, we measured $^{15}$N CP-MAS spectra of the four VM+ bound BM2 samples at 243 K on the 400 MHz spectrometer. About 100,000 scans were averaged for each sample to obtain sufficient sensitivity. The $^1$H-$^{15}$N CP contact time was 3 ms, and the $^1$H and $^{15}$N spin-lock matching condition was optimized by maximizing the 250-ppm...
unprotonated $^{15}$N peak in a pH 8.5 histidine model compound. The ratio of the integrated intensities of the unprotonated to the protonated $^{15}$N peaks in amino acid histidine ranged from 1.35 to 1.45, reflecting incomplete polarization transfer to the unprotonated nitrogen. This factor is taken into account in extracting the ratio of neutral to cationic His19 in BM2.

Two-dimensional (2D) $^{13}$C-$^{13}$C correlation spectra were measured using a proton-driven spin-diffusion experiment with dipolar-assisted rotational resonance (DARR) mixing$^{42}$. The 400 MHz data were acquired at 243 K with a spin diffusion mixing time of 50 ms, while the 800 MHz data were obtained at 263 K using a mixing time of 100 ms. 2D $^{15}$N-$^{13}$C heteronuclear correlation (HETCOR) spectra were measured at 263 K using a REDOR-based pulse sequence$^{43}$ with dipolar mixing times of 0.5–0.6 ms.

A 2D $^1$H-$^{13}$C HETCOR experiment with spin diffusion$^{44–46}$ was used to measure the hydration of BM2 residues. A $^1$H T$_2$ filter of 1–6 ms was used before spin diffusion to suppress the peptide $^1$H magnetization while retaining most of the water and lipid $^1$H magnetization. Two $^1$H spin diffusion mixing times were measured for each sample, a short mixing time of 4 ms and a long mixing time, which was 50 ms for the pH 7.5 PC/PG/Chol sample and 100 ms for the pH 5.5 PC/PG/Chol sample and the VM+ samples. Due to the different phase transition temperatures of the two membranes, the PC/PG/Chol samples were measured at 263 K while the VM+ samples were measured at 293 K. Water $^1$H T$_1$ relaxation times were measured using an inversion recovery sequence and range from ~230 ms to ~850 ms. The pH 7.5 PC/PG/Chol membrane showed the shortest water $^1$H T$_1$ relaxation time (~230 ms), thus we chose a long mixing time of 50 ms instead of 100 ms for this sample to obtain sufficient sensitivity. Based on our studies of water interactions with a number of membrane peptides and proteins such as AM2$^{46,47}$, antimicrobial peptides$^{34,48}$, and bacterial toxins$^{45,49}$, the 50 ms mixing is sufficient to equilibrate the water polarization with BM2. Water $^1$H cross-peak intensities with peptide $^{13}$C signals were extracted from the cross section of each 2D spectrum, and the intensity ratios between the short and long mixing-time spectra were corrected for $^1$H T$_1$ relaxation according to $S/S_0 = S(4$ ms$)·e^{4$ ms$/T_1}/S(100$ ms$)·e^{100$ ms$/T_1}$ or $S/S_0 = S(4$ ms$)·e^{4$ ms$/T_1}/S(50$ ms$)·e^{50$ ms$/T_1}$ to give the initial spin diffusion buildup intensity from water to the protein residue. Error bars for $S/S_0$ values were propagated from the signal-to-noise ratios of the peaks.

DNP experiment conditions

DNP spectra were measured on the pH 5.5 BM2 sample bound to the d$_{31}$-VM+ membrane on a 400 MHz/263 GHz DNP spectrometer at Bruker Biospin (Billerica, MA). The sample temperatures were ~108 K with the microwave (MW) off and ~117 K with the MW on at 130 mA. Double-quantum-filtered (DQF) 1D $^{13}$C spectra were measured using 2.2 ms of SPC5 $^{13}$C-$^{13}$C dipolar recoupling$^{50}$. 1D and 2D $^{15}$N-$^{13}$C dipolar filtered spectra were measured using a double CP pulse sequence with a $^1$H-$^{15}$N CP contact time of 1.25 ms and a $^{15}$N-$^{13}$C CP contact time of 9.0 ms.
Extraction of His19 pKₐ’s

His19 acid-dissociation constants (Kₐ’s) were extracted from the ¹⁵N CP-MAS spectra based on the fact that the neutral-to-cationic histidine concentration ratios are reflected by the relative intensities of the protonated NH peaks and the unprotonated nitrogen peak according to \[ \frac{[\text{His}]}{[\text{HisH}^+]} = 2\left(\frac{I_{\text{NH}}}{I_N}\right)^{-1}. \] ¹⁵N peak intensities were integrated from 258-240 ppm for the unprotonated imidazole nitrogen, which results exclusively from neutral histidine, and 200-150 ppm for the protonated nitrogen, which results from both neutral and cationic histidines. The \( \kappa \) in the equation is a correction factor for the reduced efficiency of \(^1\text{H}\)-¹⁵N CP for the unprotonated nitrogen compared to protonated nitrogen. The \( \kappa \) value is measured from the optimized CP spectrum of pH 8.5 amino acid histidine. The model-compound spectrum was measured with the same CP contact time, \(^1\text{H}\) and ¹⁵N radiofrequency matching conditions, and MAS frequency as for the BM2 samples. For these ¹⁵N CP experiments, the samples were spun at 7 kHz and a CP contact time of 3 ms was used. Uncertainties in the neutral-to-cationic histidine concentration ratios were propagated from the signal to noise ratios of the spectra.

The neutral-to-cationic histidine concentration ratios at different pH were fit using the equation

\[
\frac{[\text{His}]}{[\text{HisH}^+]} = 1 \cdot \frac{K_{a1} \cdot K_{a2} \cdot K_{a3} \cdot K_{a4}}{K_{a1} \cdot K_{a2} \cdot K_{a3} \cdot K_{a4}} \cdot \frac{K_{a1} \cdot K_{a2} \cdot K_{a3}}{K_{a1} \cdot K_{a2} \cdot K_{a3}} \cdot \frac{K_{a1} \cdot K_{a2} \cdot K_{a3}}{K_{a1} \cdot K_{a2} \cdot K_{a3}} \cdot \frac{K_{a1} \cdot K_{a2} \cdot K_{a3}}{K_{a1} \cdot K_{a2} \cdot K_{a3}}
\]

in MATLAB’s Curve Fitting Toolbox. Among the four pKₐ’s, pK₁ has the highest value and is for the first protonation event, while pK₆ corresponds to the last protonation event and has the lowest value. Uncertainties in the determined pKₐ’s were estimated from the uncertainty ranges of the histidine concentration ratios.

Results

Backbone conformation and dynamics of BM2(1–33) in lipid membranes

We measured 1D and 2D ¹³C spectra as a function of pH, membrane composition and temperature (Figs. 2, 3) to determine the global conformation and dynamics of BM2(1–33). At low temperature (243 K), the labeled residues exhibit strong signals at \( \alpha \)-helical chemical shifts in both PC/PG/Chol and VM+ membranes. The S12 Ca and C\( \beta \) cross peaks are close to the diagonal, characteristic of the \( \alpha \)-helical conformation. Distinct \( \alpha \)-helical Ca and C\( \beta \) chemical shifts are also observed for A17 and G26. Interestingly, for S12, A17 and G26, Ca and CO chemical shifts decrease at low pH compared to high pH (Table 1), suggesting that the local conformations at these residues become less ideally \( \alpha \)-helical at acidic pH. The linewidths also increase at low pH, indicative of larger conformational heterogeneity in the open state. Increasing the temperature to 303 K weakened most of the intensities of the PC/PG/Chol bound peptides while retaining the intensities of VM+ bound peptides (Fig. 2a, b). This indicates that BM2(1–33) undergoes intermediate-timescale motion in PC/PG/Chol bilayers, which interferes with \(^1\text{H}\) decoupling and \(^1\text{H}\)-¹³C cross polarization, but is largely immobilized in the VM+ membrane. While the two membranes both contain POPC.
and cholesterol, POPE and sphingomyelin have significantly higher phase transition temperatures than POPG, thus the VM+ membrane is more viscous than the PC/PG/Chol membrane, slowing down BM2(1–33) motion.

DNP experiments at cryogenic temperature (~117 K) enhanced the sensitivity sufficiently to allow us to resolve the BM2 signals from the lipid background signals through $^{15}$N-$^{13}$C and $^{13}$C-$^{13}$C dipolar filters (Fig. 2c–e). A sensitivity enhancement factor of 21–27 was achieved using a recently optimized membrane preparation protocol for DNP NMR. With the removal of the natural abundance $^{13}$C signals of lipids and cryoprotectants, we resolved two His19 Cβ signals at 29 and 27 ppm and the S12 Ca and Cβ signals at ~60 ppm (Fig. 2d, e).

**pH- and membrane-dependent structures of His19**

1D $^{13}$C spectra (Fig. 2), 2D $^{13}$C-$^{13}$C correlation spectra (Fig. 4) and 2D $^{15}$N-$^{13}$C correlation spectra (Fig. 5) provide detailed information on the chemical structure of the proton-selective residue, His19, under different pH and membrane compositions. In the aromatic region of the 1D spectra (Fig. 2), both the pH 7.5 PC/PG/Chol sample and the pH 5.5 VM+ sample show Cγ and Cδ2 chemical shifts of ~135 ppm and ~114 ppm, which are diagnostic of neutral histidines. The DNP-enhanced spectra show additional intensities at ~118 ppm, which can be assigned to Cδ2 of cationic histidine. 2D $^{13}$C-$^{13}$C correlation spectra (Fig. 4) allowed unambiguous assignment of the $^{13}$C signals of the various tautomeric and cationic states of histidine. In the aliphatic region, the pH 7.5 PC/PG/Chol sample and the pH 5.5 VM+ sample both show two Ca-Cβ cross peaks, while the pH 5.5 PC/PG/Chol sample and the pH 4.5 VM+ sample exhibit four Ca-Cβ peaks. The nature of these Ca-Cβ peaks can be identified based on their correlations with the sidechain Ce1, Cδ2 and Cγ chemical shifts. For the higher-pH spectra, the (54.8, 29.7) ppm cross peak can be assigned to the τ tautomer due to its correlation with the 113.5 ppm Cδ2 chemical shift, while the (54.3, 27.1) ppm cross peak can be assigned to the π tautomer because of its connectivity to the 124 ppm Cδ2 peak. In the lower-pH spectra, the τ tautomer Ca-Cβ peak is still present, while two cross peaks with Cβ chemical shifts of ~26 ppm can be assigned to cationic histidines, cat2 and cat3, based on their correlations with Cγ and Cδ2 chemical shifts of ~127 ppm and ~117 ppm. The fourth cross peak at (58.0, 28.9) ppm is correlated with the same Ce1 and Cγ peaks as the τ tautomer, but also correlates with a 116.5-ppm Cδ2 peak, which thus reveals it to be a cationic histidine. This state is similar to the previously observed cat1 state of AM2 His37. Finally, in the pH 4.5 PC/PG/Chol sample, the spectrum simplifies to only cationic histidine, with two partially overlapped Ca-Cβ cross peaks (Fig. 4c). These chemical shifts are summarized in Table 1.

These 2D $^{13}$C-$^{13}$C correlation spectra show that the VM+ membrane produces the same set of His19 structures as the PC/PG/Chol membrane, but at 1–2 lower pH units. Thus, the negatively charged membrane facilitates proton association with His19. The τ and π tautomers that coexist in the pH 5.5 VM+ membrane have weak but clear cross peaks with each other, for example between the τ Cβ chemical shift and the π Cγ chemical shift at (30, 126) ppm and between π Cδ2 and τ Cδ2 at (125, 114) ppm (Fig. 4d). These inter-tautomer
cross peaks indicate that both τ and π tautomers exist in the same tetramer instead of being separated into different channels.  

2D $^{15}$N-$^{13}$C HETCOR spectra (Fig. 5) give complementary information about the pH-dependent structure distribution of His19. At pH 5.5, the backbone N-Cα cross peaks show relatively narrow linewidths of 1.5 ppm. Decreasing the pH to 4.5 increased the backbone linewidths, similar to the situation of the 2D $^{13}$C-$^{13}$C correlation spectra. The aromatic region of the 2D spectrum is simple for the pH 5.5 sample, with two τ tautomer peaks and one π tautomer peak (Fig. 5a). But at pH 4.5 the 2D spectrum is much more complex, showing multiple cationic species (Fig. 5b). Four Ne2-C62 cross peaks are resolved at $^{13}$C chemical shifts of 113 – 118 ppm, and can be assigned to the τ tautomer and three cationic histidines, while two Nδ1-Cγ cross peaks are observed and can be assigned to cat2 and cat3.

Fig. 6 compares the pH-dependent 2D $^{13}$C-$^{13}$C correlation spectra encoding the $^{13}$C chemical shifts of His19 in BM2 and His37 in AM2. With decreasing pH, His19 progresses from a dual τ–π tautomeric mixture to a four-state neutral and cationic mixture. This change occurs from pH 7.5 to pH 5.5 in the PC/PG/Chol membrane and from pH 5.5 to pH 4.5 in the VM+ membrane. At sufficiently low pH, which is 4.5 for the PC/PG/Chol membrane and pH 4.0 for the VM+ membrane, a single cationic state is observed. The cytoplasmic-containing AM2(21–97) also resolved four cationic His37 species, cat1 to cat4 (Fig. 6b), with similar chemical shifts as those of BM2 His19; however, the four states do not occur at the same pH. These four species have been assigned to histidines in variously charged tetrads, +1 to +4, based on the pH at which they become populated. The shorter AM2 TM peptide also showed similar tautomeric chemical shifts (Fig. 6c), but fewer cationic species were found compared to the cytoplasmic-containing AM2. Importantly, the pH at which the multiple histidine states appear differs qualitatively between BM2 and AM2: BM2 concentrates a larger number of histidine species into a narrower pH range, which is around pH 4.5 for the VM+ membrane and about pH 5.5 for the PC/PG/Chol membrane. AM2(21–97) shows three cationic species at pH 5.4 while AM2(22–46) shows only two cationic states at pH 5.2. Since these AM2 and BM2 spectra were measured in similar eukaryotic-mimetic lipid mixtures of VM+ and VM, the different pH dependences reflect real differences in the acid-dissociation equilibria of the histidines. Specifically, BM2 His19 retains neutral tautomers to lower pH than His37 in AM2, indicating lower pKₐ’s (vide infra).

**DNP-enhanced 2D spectra at cryogenic temperature**

Sensitivity enhancement by DNP allowed the observation of additional features in 2D $^{13}$C-$^{13}$C and $^{15}$N-$^{13}$C correlation spectra at pH 5.5 (Fig. 4f, 5c). Where previously only neutral τ and π tautomers were detected at 263 K, we now also observe the signals of cat1, cat2 and cat3, while the π tautomer cross peaks are below the detection limit (Fig. 4f).

Spectral integration indicates that the cationic species represent 50–60% of the total intensities under the DNP experimental condition, while the higher-temperature 2D spectra at pH 5.5 do not show detectable cationic histidine intensities. Instead, the DNP spectrum at pH 5.5 is more similar to the high-temperature 2D spectrum at pH 4.5, which shows ~80% cationic intensities. This suggests that the effective pKₐ’s of the histidine are higher at cryogenic temperature than at high temperature. The 2D $^{15}$N-$^{13}$C correlation spectrum (Fig.
5c) shows similar results, with the intensity pattern approaching the pH 4.5 spectrum at high temperature (Fig. 5b).

**His19 proton exchange equilibria and dynamics**

To obtain quantitative information about the His19 protonation equilibria, we measured the 1D $^{15}$N CP-MAS spectra of VM+ bound BM2 from pH 6.5 to pH 4.0. Since there are in principle four acid dissociation constants for the tetrameric channel, the pK$_a$ extraction requires a minimum of four samples. The spectra were measured at 243 K where proton transfer dynamics are suppressed. Fig. 7a shows the expected progressive decrease of the 250-ppm unprotonated $^{15}$N peak intensity with decreasing pH. As shown before\textsuperscript{19}, the relative concentration of neutral and cationic histidines can be obtained from the intensity ratio $I_{\text{NH}}/I_N$. Fig. 8b and Table 2 show that [His]/[His$^+$] decreases rapidly from 10.8 to 0.14 over the pH range of 6.5 to 4.0 for the VM+ bound peptide. Fitting these values to equation (1) yielded pK$_{a}$'s of 6.1±0.1, 5.7±0.1, 4.5±0.3, and 4.2±0.4. The two lowest pK$_{a}$'s have the largest uncertainty, due to the fact that the unprotonated $^{15}$N peak intensity is weak at pH 4.0, thus the percentage of neutral histidines is difficult to determine with high precision. Alternative fits assuming only two distinct pK$_{a}$'s can also agree with the data, and extraordinarily high precision and accuracy in the concentration ratios as well as the sample pH would be required to distinguish these different models. Nevertheless, the BM2 data differs from the recent studies of His37 in AM2(18–60) and in the cytoplasmic-containing AM2(21–97), whose $^{15}$N spectra could not be adequately fit to four pK$_{a}$'s\textsuperscript{20,53}. Regardless whether there are two or four distinct pK$_{a}$'s, the observation that the two highest pK$_{a}$'s cluster and the two lowest pK$_{a}$'s cluster is unambiguous, and this translates to charged-tetrad population curves that have a high percentage of the +2 state over a relatively wide pH range, preceded by the persistence of the fully neutral (0 charge) and +1 tetrads above pH 6, and followed by quick onsets of the +3 and +4 channels below about pH 4.5 (Fig. 8c).

Incorporating BM2(1–33) into the anionic PC/PG/Chol membrane caused a shifted pH dependence of the $^{15}$N spectral intensities: the pH 7.5 PC/PG/Chol spectrum resembles the pH 5.5 VM+ spectrum while the pH 5.5 PC/PG/Chol spectrum approaches the pH 4.5 VM+ spectrum (Fig. 7b). This is consistent with the 2D spectra that indicate a shift of the His19 protonation equilibria to higher pH in the anionic membrane compared to the neutral membrane.

To investigate the proton transfer kinetics, we measured $^{15}$N CP spectra at high temperature (Fig. 7c). Since BM2 tetramers undergo intermediate-timescale motion in the PC/PG/Chol membrane, high-temperature intensities can only be observed for VM+ bound peptide. At 303 – 313 K, the pH 5.5 VM+ sample retained the neutral tautomer peaks at 251 ppm and 160–170 ppm, but in addition a broad band of intensities between 170 and 250 ppm emerged, suggesting chemical exchange of the imidazole nitrogens with a distribution of equilibrium constants. Most importantly, at pH 4.5, the spectrum exhibits an exchange peak at 213 ppm and additional intensities at ~185 ppm. These intermediate $^{15}$N chemical shifts are identical to the values seen for His37 in AM2, and can be assigned to Ne2 and N61 chemical exchange among the τ tautomer, π tautomer, and cationic histidine$^{19,22}$. Thus, the
chemical-exchange equilibria among the three histidine states are conserved between AM2 and BM2.

BM2 channel hydration from water-protein 2D correlation experiments

To investigate whether BM2 channel hydration differs from AM2, we measured 2D water-peptide $^1$H-$^{13}$C correlation spectra as a function of the $^1$H spin diffusion mixing time. Fig. 9a shows a representative 2D spectrum of BM2(1–33) in the PC/PG/Chol membrane at pH 7.5, measured with a $^1$H mixing time of 50 ms. Spectra measured at a shorter mixing time of 4 ms give information about the initial spin diffusion rate from water to the peptide. At pH 7.5 (Fig. 9b), the intensity of the short-mixing spectrum is 20–30% of the long-mixing spectrum for the $\text{C}_\alpha$ and $\text{C}_\beta$ peaks, while the His19 aromatic signals have higher intensities of >40%. At acidic pH, the 4-ms $\text{C}_\alpha$ and $\text{C}_\beta$ peak intensities increase compared to the higher pH sample, indicating faster water-protein polarization transfer. The His19 sidechain intensities are the highest, showing 80–90% of the equilibrium intensities. Thus, BM2 is much more hydrated at low pH than at high pH, consistent with an open channel.

Fig. 9d–f summarizes the S/S$_0$ ratios of the resolved $^{13}$C signals of BM2(1–33). For each membrane, the water cross peaks are higher at low pH than at high pH. Between the two lipid membranes, the PC/PG/Chol bound BM2 exhibits higher water cross peaks than the VM+ bound peptide at the same pH, indicating that the anionic membrane facilitates channel hydration. Thus, the viscous and neutral VM+ membrane contains less hydrated BM2 channels than the PC/PG/Chol membrane. Among the four labeled residues, the His19 sidechain has the highest water cross peaks, consistent with the polar and pore-facing nature of this residue. Interestingly, S12 exhibits a large pH-dependent change in the water cross peak intensity. In the PC/PG/Chol membrane, the S/S$_0$ values changed from 0.29±0.04 at pH 7.5 to 0.42±0.04 at pH 5.5, and decreasing the pH by 1 unit in the VM+ membrane caused the S/S$_0$ values to increase from 0.14±0.03 to 0.38±0.03. The latter change is larger than that of all other residues measured so far in AM2 and BM2, including S31 in AM2 (Fig. 9g), indicating that the pore-facing serine residues in BM2 play a prominent role in channel hydration, which should promote H$^+$ relay to the HxxxW motif.

Discussion

The data shown here give the first detailed structural and dynamical information about the proton-conducting heart of BM2 in lipid bilayers. The chemical shifts of the four labeled residues indicate that the BM2 TM peptide is $\alpha$-helical in a range of pH and in two different membranes. Although the number of labeled residues is too small to permit more detailed structural conclusions, it is nevertheless interesting that the three non-histidine residues change their chemical shifts at low pH in a direction that corresponds to less ideal $\alpha$-helical conformations at low pH. This trend is opposite that of the AM2 TM peptide, which exhibits more ideal $\alpha$-helical chemical shifts, and straight helices, at acidic pH$^{55}$. The non-ideal helical chemical shifts seen in BM2 may be related to the coiled coil structure found for DHPC-bound BM2, even though the solution NMR structure was solved at pH 7.5$^{12}$. More complete structural measurements are required to elucidate the high-resolution backbone conformation and oligomeric assembly of BM2 in lipid bilayers.

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The pH-dependent $^{15}$N spectra and 2D correlation spectra indicate that His19 protonates at significantly lower pH than His37 in both the TM and cytoplasmic-containing constructs of AM2. The average pK$_a$ of His19 in BM2-TM is 5.1 (Table 3). In comparison, the average pK$_a$'s of His37 in AM2(22–46) and AM2(21–97) are 5.9 and 6.3, respectively, in similar virus-mimetic membranes$^{19,20}$. The earliest measurement of His37 pK$_a$ in AM2(22–46) was conducted in the significantly different model membrane DMPC/DMPG, and found a much higher average pK$_a$ of ~6.9$^{24}$. In another recent study, the intermediate construct AM2(18–60), which includes the TM helix and the amphipathic helix, was measured in the negative-curvature membrane formed by diphytanoylphosphocholine$^{56,57}$. This study resolved two pK$_a$'s with an average of 6.1$^{53}$. Therefore, despite the significant variations in the protein construct lengths and membrane compositions, the His37 pK$_a$'s in AM2 measured under all conditions so far are higher than the BM2 His19 pK$_a$'s found here. The His19 titration curve parallels the titration curve of His37 in the cytoplasmic-containing AM2(21–97) but is shifted by ~1.2 pH unit lower (Fig. 8b). Compared to the AM2 TM peptide, the BM2 titration curve drops more steeply at low pH. Overall, the His19 tetrads remain predominantly in low charged states of +2, +1 and 0 till about pH 5, below which the +3 and +4 tetrad populations surge (Fig. 8c). This result is found for the VM+ membrane, which is the membrane used in two of the previous four AM2 studies, thus the pK$_a$ decrease reflects real differences in the protonation equilibria of His19 in BM2 and His37 in AM2.

The lower pK$_a$'s of His19 was surprising, because the BM2 channel pore is lined with polar residues such as S12, S19 and S26 (Fig. 1), which would be expected to facilitate proton relay to His19 and thus stabilizing the cationic state of His19. In general, altered equilibrium constants imply altered structures, thus the BM2 TM helix structure and structural equilibrium must differ in crucial ways from the AM2 structure to account for this pK$_a$ decrease. The solution NMR structure of micelle-bound BM2(1–33) shows a coiled coil (Fig. 9i) while AM2 in the same micelles exhibits straighter helices. This secondary structure difference, if verified to persist in the lipid bilayer, may affect the proton-transfer equilibria at His19. Since there is yet no high-resolution bilayer-bound structure of BM2, we focus our consideration on the equality $K_a = k_{off}/k_{on}$ between the acid-dissociation constant and the ratio of the first-order proton dissociation rate constant $k_{off}$ and the second-order proton-association rate constant $k_{on}$. The lower pK$_a$'s or larger K$_a$'s indicate larger $k_{off}$ and/or smaller $k_{on}$. Information about the proton-exchange kinetics is partly contained in the high-temperature His19 $^{15}$N spectra (Fig. 7c), which show a 213-ppm exchange peak with a full width at half maximum of 400 ± 140 Hz. In comparison, the exchange peak of His37 in the AM2 TM peptide has a larger linewidth of 640 ± 70 Hz. Since proton dissociation is the rate-limiting step in proton transfer, the narrower His19 exchange peak suggests a larger $k_{off}$ than that of His37 in AM2. Although $k_{on}$ may be increased by the serine triplets in the N-terminal half of the TM helix, the $k_{off}$ may increase more, thus shifting the equilibrium towards the neutral state. This interpretation of a preferentially increased $k_{off}$ is consistent with liposome assays that indicate a larger H$^+$ flux of BM2(1–33) compared to AM2(18–60)$^{12}$.

What might be the structural basis for an increased proton-dissociation rate constant for His19? We propose that the second titratable histidine, His27, C-terminal to the HxxxW motif and one helical turn away from Trp23, may speed up proton release from the His19-
Trp23 pair. A UV resonance Raman study showed that His27 has a higher pKₐ than His19⁵⁸, indicating that His27 is more cationic than His19 at the same pH and thus consistent with this hypothesis. Mutation of His27 to Ala was found to decrease the H⁺ conductance by ~25% compared to the wild-type, indicating that His27 plays an active role in H⁺ conduction through the BM2 channel. Importantly, this second histidine is absent in the AM2 TM domain, whose equivalent position harbors an Arg45, which is constitutively charged under the full pH range of the channel. Thus, this single amino-acid difference may explain the observed destabilization of the cationic state of His19 or the lowering of the pKₐ's.

At high temperature, the exchange-averaged ¹⁵N chemical shifts (213 ppm and ~185 ppm) of BM2 are the same as for AM2. We have previously shown that this interconversion among different histidine structures is most observable in low charged tetradś²², because they contain a high concentration of neutral histidine, and correspond a concentration ratio of 2 : 1 : 1 for the interconverting τ tautomer, π tautomer, and cationic histidine. Thus, the resolved ¹⁵N chemical exchange peak does not reflect the full population distribution, and an altered histidine-water proton-transfer equilibrium at low temperature can coexist with a conserved histidine interconversion equilibrium at high temperature.

The above quantified pKₐ's are obtained from BM2 bound to the neutral and relatively viscous VM+ membrane. The membrane composition also affects the relative concentrations of neutral and cationic histidines. The ¹⁵N spectra of the peptide in the anionic and more fluid PC/PG/Chol membrane clearly indicate that His19 protonates at higher pH in this membrane. We attribute this change to increased local proton concentrations and enhanced protein conformational motions necessary for proton binding and release in the negatively charged and fluid PC/PG/Chol membrane. If the former accounts for the entire extent of change in the cationic/neutral histidine concentration ratios, then the true equilibrium constant would be unaffected because the higher local H⁺ density rather than the average sample pH should be used in considering the equilibrium constant. The importance of electrostatic charges on the H⁺ transfer equilibria is also observed in a recent study of the cytoplasmic-containing AM2, which shows higher pKₐ's for His37 than the TM peptide. This result was attributed to the anionic nature of the cytoplasmic domain²⁰, which should elevate the local proton concentrations.

The NMR observed reduction of BM2 His19 pKₐ's compared to AM2 His37 differs from UV resonance Raman data, which found that AM2 His37 has an average pKₐ of 5.7²⁷ while BM2(3–33) exhibits a midpoint of transition at pH ~6.5 for the pH-dependent quenching of W23 fluorescence emission by His19⁵⁸. These resonance Raman data were obtained from POPE/POPS (1:1) bound peptides. Two possible reasons could explain this discrepancy. First, the negatively charged POPE/POPS membrane may affect the BM2 and AM2 protonation equilibria differently compared to the neutral VM+ membranes used in the solid-state NMR experiments. Second, fluorescence quenching can be caused by changes in the environmental polarity and mobility of the Trp sidechain, in addition to His-Trp cation-π interaction, thus the interpretation of the resonance Raman spectra has inherent uncertainty. In comparison, the ¹⁵N NMR chemical shifts are unambiguous indicators of the chemical structure of histidine.
The water-peptide 2D correlation spectra show increased hydration at low pH than at high pH for BM2, similar to the AM2 behavior\textsuperscript{46}. The negatively charged PC/PG/Chol membrane at pH 7.5 gives higher water cross peaks than the VM+ membrane at pH 5.5, indicating that the VM+ membrane causes a tighter and less hydrated channel. The higher pore hydration at more acidic pH should facilitate His19 protonation by delivering protons through the water molecules rapidly to histidine. Interestingly, the VM+ bound S12 exhibits the largest pH-dependent increase of the water cross peak intensity among all residues measured so far in both BM2 and AM2. This acid-induced hydration increase is not solely due to the pH dependence of chemical exchange between the serine hydroxyl group and water, because S31 in AM2 does not show a particularly high S/S\textsubscript{0} value at low pH, and S12 at pH 5.5 has similar water cross peak intensities as the non-exchangeable A17 and G26. It is known that the chemical exchange rate of serine OH is ~900 s\textsuperscript{−1} at pH 7 and 35°C, which is two-fold smaller than the proton exchange rate of imidazole NH (1700 s\textsuperscript{−1})\textsuperscript{59}. Yet here S12 Ca has higher S/S\textsubscript{0} than H19 Ca at pH 4.5 (Fig. 9e). Thus, S12 hydration is truly increased at low pH, suggesting that this residue is important for mediating proton relay to the HxxxW motif.

The DNP spectra at cryogenic temperature show moderate differences in the pH equilibria of His19 from the high-temperature data. For the pH 5.5 sample, the concentration of cationic histidine increased to 40–50%, while at 263 K mainly neutral histidines are observed at the same prepared pH. We attribute this increased cationic concentration at low temperature to the different temperature dependences of the buffer and the imidazole pK\textsubscript{a}’s. The citrate buffer pK\textsubscript{a} has a linearly extracted temperature coefficient d(pK\textsubscript{a}/dT of -0.01\textsuperscript{60,61}, while the imidazole pK\textsubscript{a} has a larger temperature coefficient of -0.02. Thus, as the sample temperature decreases, the citrate pK\textsubscript{a} should increase but to a smaller extent than the imidazole pK\textsubscript{a}. This means that at low temperature, the imidazole sidechain will be more readily protonated by the buffer. The thermodynamic parameters for the ionization of buffer ions and imidazoles are not known to very low temperatures, and the exact freezing temperature of the channel water is also unknown. But the relative trend that the imidazole pK\textsubscript{a} has a more negative temperature coefficient than the citrate pK\textsubscript{a}’s is reliable, and favors protonation of His19 at low temperature. A potential acid-base equilibrium change between cryogenic temperature and ambient temperatures should thus be taken into account in DNP NMR experiments of pH-sensitive proteins and other biomolecules in buffered solution.

Conclusions

In its global conformational and dynamical behavior, BM2(1–33) is similar to the AM2 TM peptide: the tetrameric channel is \(\alpha\)-helical and undergoes intermediate-timescale motion in low-viscosity membranes at physiological temperature. However, important differences are found at the proton-selective histidine in the HxxxW motif, and in the pore-lining residues leading to this motif. His19 in BM2 remains neutral down to lower pH than His37 AM2, and cationic His19 appears in a narrower pH range than His37 in AM2. Thus, His19 in the tetrameric BM2 channel protonates with lower and more clustered pK\textsubscript{a}’s. Quantitative analysis of the \(^{15}\text{N}\) spectra yielded pK\textsubscript{a} values of 6.1, 5.7, 4.5 and 4.2, which are about 1 pH unit lower than His37 in AM2. We propose that the altered pK\textsubscript{a}’s result from the presence of a second titratable histidine, C-terminal to His19, in BM2. This second histidine is absent in AM2 and likely increases the proton dissociation rate constant of His19. This model is
consistent with the narrower His19 chemical exchange peak compared to AM2 His37 and with H+ flux and mutagenesis data. Similar to AM2, BM2 channels are more hydrated at acidic pH than at high pH, but compared to AM2, S12 in BM2 exhibits the largest acid-induced increase in the water cross peak intensity, consistent with the model that the Ser triplet lining the BM2 pore facilitates proton relay to His19 by increasing pore hydration. We found that the lipid membrane also plays a significant role in proton-transfer dynamics and channel hydration: anionic membranes facilitate His19 protonation at high pH and increase the channel hydration compared to neutral membranes. These results give rich insights into how the amino acid sequence and the membrane environment affect the functional structure and dynamics of this family of viral proton channels.

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References


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Figure 1.
Comparison of the amino acid sequences (a) of the BM2 and AM2 transmembrane domains. Lowercase letters indicate heptad repeat, with a and d indicating pore-facing positions. The conserved HxxxW motif is underlined and the $^{13}$C, $^{15}$N-labeled residues are shown in red. (b) Four-helix bundle organization of the BM2 and AM2 transmembrane domains, with the pore-facing a and d residues indicated.
Figure 2.
1D $^{13}$C CP-MAS spectra of BM2(1–33) as a function of pH, temperature, and membrane composition. (a) BM2 in the PC/PG/Chol membrane at pH 7.5. (b) BM2 in the VM+ membrane at pH 5.5. The VM+ membrane immobilizes the peptide while the PC/PG/Chol membrane promotes peptide motion at high temperature. (c–e) DNP-enhanced $^{13}$C spectra of BM2 at pH 5.5 in a partially deuterated VM+ membrane. (c) $^{13}$C CP spectrum with MW on. The sensitivity is 21–27 times that of the MW-off spectrum (not shown). (d) $^{15}$N-$^{13}$C dipolar filtered $^{13}$C spectrum. (e) $^{13}$C DQF spectrum.
Figure 3.
2D $^{12}$C-$^{13}$C correlation spectra of site-specifically labeled BM2(1–33) in different membranes and at different pH. (a) PC/PG/Chol at pH 7.5. (b) PC/PG/Chol at pH 5.5. (c) PC/PG/Chol at pH 4.5. (d) VM+ membrane at pH 5.5. (e) VM+’ membrane at pH 5.5 with DNP. (f) VM+ membrane at pH 4.5.
His19 regions of the 2D $^{13}$C-$^{13}$C correlation spectra of BM2(1–33) at different pH and in different lipid membranes. The spectra in (a–c) were measured at 243 K on PC/PG/Chol bound peptide. (a) pH 7.5, (b) pH 5.5, (c) pH 4.5. The spectra in (d–e) were measured at 263 K on VM+ bound peptide at (d) pH 5.5 and (e) pH 4.5. (f) DNP-enhanced spectrum of BM2(1–33) in the $d_{31}$-VM+’ membrane at pH 5.5. Spectra (a–c) were measured on a 400 MHz spectrometer under 7 kHz MAS, while spectra (d–e) were measured on an 800 MHz spectrometer with 14.5 kHz MAS. The DNP spectrum (f) was measured on a 400 MHz/263 GHz spectrometer at 117 K under 9 kHz MAS. The DARR mixing times were 50 or 100 ms in these spectra.
Figure 5.
2D $^{15}$N-$^{13}$C correlation spectra of His19 in BM2(1–33). (a, b) BM2 in the VM+ membrane, measured on an 800 MHz spectrometer with 14.5 kHz MAS at 263 K. (a) pH 5.5. (b) pH 4.5. (c) BM2 at pH 5.5 in the d$_{31}$-VM+’ membrane, measured on a 400 MHz/263 GHz DNP spectrometer under 9 kHz MAS at 117 K. Note the different neutral and cationic histidine distribution between high and low temperatures at pH 5.5, indicating temperature-induced pK$_{a}$ shifts.
Figure 6.
Comparison of His19 and His37 conformational distribution in BM2 and AM2 from 2D $^{13}$C-$^{13}$C correlation spectra. All peptides and proteins were bound to eukaryotic-mimetic lipid membranes, except for the pH 4.5 spectrum of BM2, which was bound to the PC/PG/Chol membrane. Spectra were measured at moderate low temperatures (243 – 273 K) where the peptides and proteins are immobilized. (a) BM2(1–33) in VM+ membranes at pH 5.5 and pH 4.5, and BM2(1–33) in the PC/PG/Chol membrane at pH 4.5. (b) AM2(21–97) in the VM+ membrane at pH 7.5 and pH 5.4. (c) AM2 TM peptide (residues 22–46) in the VM membrane at pH 7.0, 5.2 and 4.5. BM2 His19 exhibits the largest number of coexisting species, at pH 4.5. The fact that neutral tautomers persist to lower pH in BM2 indicates that
His19 protonates with lower and more clustered pKₐ’s. The spectrum of the pH 4.5 VM+ bound BM2 resembles the spectrum of the PC/PG/Chol bound BM2 at pH 5.5 in Fig. 4b, indicating the influence of the negatively charged lipid on the protonation equilibria.
Figure 7.
1D $^{15}$N CP spectra of BM2(1–33) as a function of pH, membrane composition and temperature. (a) Spectra of the VM+ bound BM2 from pH 4.0 to pH 6.5 at 243 K. (b) Spectra of PC/PG/Chol-bound BM2 at pH 5.5 and pH 7.5 at 243 K. (c) Comparison of high-temperature ($303 – 308$ K) $^{15}$N spectra of BM2 and AM2 at acidic pH. An $^{15}$N exchange peak is detected at ~213 ppm at pH 4.5 for BM2, at pH 5.2 for AM2-TM, and at pH 5.4 in the S31N mutant of AM2-TM. The exchange-averaged chemical shifts are the same in all
M2 samples, but the exchange linewidths differ. The BM2 exchange peak is fit (gray line) to give a linewidth of 400±140 Hz.
Figure 8.

pKₐ extraction of His19 in BM2(1–33). (a) NH to N intensity ratios as a function of pH. (b) Neutral-to-cationic histidine concentration ratios as a function of pH. The BM2 data (black) are compared with previously measured AM2 TM peptide data (orange) and AM2(21–97) data (blue). The neutral histidine concentration in BM2 is higher than that of AM2 at similar pH. The extracted pKₐ’s for BM2 are 6.1, 5.7, 4.5 and 4.2, whose average is lower than that of AM2 constructs. The average pKₐ’s of the three M2 samples are indicated as solid lines at the bottom. (c) Populations of charged tetrads of His19 in BM2 as a function of pH. The intercepts of adjacent population curves correspond to the pKₐ’s.
Figure 9.
2D $^1\text{H}$-$^{13}\text{C}$ HETCOR spectra of membrane-bound BM2(1–33) to probe channel hydration.
(a) Representative 2D spectrum of PC/PG/Chol bound peptide at pH 7.5, measured with
a $^1\text{H}$ spin diffusion mixing time of 50 ms. (b) Water cross sections of BM2(1–33) between 4
ms and 50 or 100 ms for the pH 7.5 and pH 5.5 PC/PG/Chol samples measured at 263 K.
The S/S$_0$ values are indicated for H19 and S12. (c) Water cross sections of the aliphatic
region of the VM+ bound BM2 at pH 5.5 and pH 4.5, showing the large increase of the
initial buildup of the S12-water cross peak intensity at lower pH. (d) Water cross section of
the aliphatic region of VM-bound AM2 at pH 4.5, showing that the S31 S/S0 value is lower than that of S12 in BM2. (e) Initial buildup (S/S0 values) at high and low pH for PC/PG/Chol-bound BM2. (f) S/S0 values of VM+ bound BM2 at pH 5.5 and pH 4.5. (g) S/S0 values of the His19 sidechain in the two membranes. (h) S/S0 values of the Ca sites of AM2 residues and of the BM2 residues S12, A17 and G26 (shaded bars). S12 in BM2 shows the largest low-pH induced increase of the S/S0 value among all labeled residues of the two peptides. (i) Comparison of the solution NMR structure of BM2 (top, PDB: 2KIX) and AM2 (bottom, PDB: 2KQT) TM domains.
Table 1

$^{13}$C and $^{15}$N chemical shifts (ppm) of BM2(1–33)$^a$.

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$^a$The chemical shifts of S12, A17, and G26 are obtained from the PC/PG/Chol sample data, while the H19 chemical shifts are average values of the PC/PG/Chol and VM+ data. All $^{13}$C chemical shifts were taken from 2D $^{13}$C-$^{13}$C correlation experiments at low temperature, while $^{15}$N chemical shifts were taken from 1D $^{15}$N and 2D $^{15}$N-$^{13}$C correlation spectra at low temperature.
Table 2

BM2(1–33) His19 imidazole $^{15}$N intensities and neutral to cationic histidine concentration ratios.

<table>
<thead>
<tr>
<th>pH</th>
<th>$I_{syr/N}$</th>
<th>$\kappa$</th>
<th>[His]/[HisH$^+$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.0</td>
<td>21.7±10.9</td>
<td>1.44±0.04</td>
<td>0.14±0.07</td>
</tr>
<tr>
<td>4.5</td>
<td>7.0±0.9</td>
<td>1.33±0.04</td>
<td>0.47±0.06</td>
</tr>
<tr>
<td>5.5</td>
<td>3.1±0.3</td>
<td>1.35±0.03</td>
<td>1.54±0.15</td>
</tr>
<tr>
<td>6.5</td>
<td>1.6±0.2</td>
<td>1.35±0.04</td>
<td>10.8±1.4</td>
</tr>
</tbody>
</table>
Table 3

Comparison of histidine $pK_a$’s in influenza A and B M2 proteins in various lipid membranes.

<table>
<thead>
<tr>
<th>M2 constructs</th>
<th>Phospholipid membrane</th>
<th>His19 and His37 $pK_a$’s</th>
<th>Average $pK_a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>BM2(1–33)</td>
<td>POPC, POPE, SM, cholesterol</td>
<td>6.1, 5.7, 4.5, 4.2</td>
<td>5.1</td>
</tr>
<tr>
<td>AM2(22–46)</td>
<td>DPPC, DPPE, SM, cholesterol</td>
<td>7.6, 6.8, 4.9, 4.2</td>
<td>5.9</td>
</tr>
<tr>
<td>AM2(21–97)</td>
<td>POPC, POPE, SM, cholesterol</td>
<td>7.1, 5.4</td>
<td>6.3</td>
</tr>
<tr>
<td>AM2(22–46)</td>
<td>DMPC, DMPG</td>
<td>8.2, 8.2, 6.3, &lt; 5.0</td>
<td>&lt; 6.9</td>
</tr>
<tr>
<td>AM2(18–60)</td>
<td>DPhPC</td>
<td>7.6, 4.5</td>
<td>6.1</td>
</tr>
</tbody>
</table>