

# Probing Hydronium Ion Histidine NH Exchange Rate Constants in the M2 Channel via Indirect Observation of Dipolar-Dephased <sup>15</sup>N Signals in Magic-Angle-Spinning NMR

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**Supporting Information** 

ABSTRACT: Water-protein chemical exchange in membrane-bound proteins is an important parameter for understanding how proteins interact with their aqueous environment, but has been difficult to observe in membrane-bound biological systems. Here, we demonstrate the feasibility of probing specific water-protein chemical exchange in membrane-bound proteins in solidstate MAS NMR. By spin-locking the <sup>1</sup>H magnetization along the magic angle, the <sup>1</sup>H spin diffusion is suppressed such that a water-protein chemical exchange process can be monitored indirectly by dipolar-dephased <sup>15</sup>N signals through polarization transfer from <sup>1</sup>H. In the example of the Influenza A full length M2 protein, the buildup of dipolar-dephased <sup>15</sup>N signals from the tetrad of His37 side chains have been observed as a function of spin-lock time. This confirms that hydronium ions are in exchange with protons in the His37 NH bonds at the heart of the M2 proton conduction mechanism, with an exchange rate constant of ~1750 s<sup>-1</sup> for pH 6.2 at -10 °C.

D iological membranes are composed of crowded membrane-bound proteins/peptides in a lipid environment. They exhibit diverse shapes and conduct many essential biological processes, such as inter- and intracellular signal transduction, protein localization, and trafficking required synergistic effects between the proteins and their surrounding complex lipid environs.<sup>2-4</sup> It has become clear that the membrane lipids have profound impacts on the structure and function of membrane-bound proteins and peptides.<sup>5-9</sup> The implications of the lipid-protein interactions have been increasingly recognized in past years and many solid-state NMR techniques have been developed to study such lipidprotein interactions. 10-14 For instance, the influence of proteins on membrane lipid dynamics and vice versa could be investigated through deuterium line-shapes<sup>15,16</sup> or by relaxation parameters.<sup>17,18</sup> Spin diffusion from lipids into membrane proteins allows for the detection of nuclear spin magnetization in proteins for the direct investigation of lipid-protein interactions.<sup>19–22</sup> Magic-angle-spinning (MAS) recoupling techniques<sup>23</sup> are also commonly used for measuring distances from the lipid head groups to specific sites on proteins so as to study the insertion and alignment of the proteins/peptides into the lipids.<sup>24-26</sup> Water-protein chemical exchange<sup>27-31</sup> is

another important parameter in the dynamic relationship between proteins and their surrounding environment that has been investigated in the past primarily through <sup>1</sup>H-<sup>15</sup>N heteronuclear correlation (HETCOR) experiments. <sup>32–34</sup> Direct observation of water—protein exchange is challenging in the presence of strong <sup>1</sup>H spin diffusion associated with the relatively rigid membrane-bound protein environments. Here, we propose one-dimensional (1D) chemical exchange measurements for probing the specific water—protein chemical exchange kinetics of membrane-bound proteins using the Influenza A full length M2 protein (M2FL).

The M2 protein is a 97-residue membrane protein that assembles as a tetrameric bundle that conducts protons at a slow rate  $(10^2 - 10^3/s)$  when activated at low pH.<sup>35,36</sup> There are two debated proton transport mechanisms: (1) the proton is transferred through the breaking and reforming of H-bonds between two pairs of His37 dimers<sup>27,37</sup> or (2) individual His37 residue shuttles protons through imidazole ring reorientations and exchanging protons with water without the process of forming intermonomer His37 H-bonds.<sup>29,30</sup> M2 spectra dramatically vary depending on the M2 constructs and lipids used in sample preparation. For instance, a set of two resonances for His37 was observed in the conductance domain M2 (22-62) in DOPC/DOPE lipids<sup>38</sup> and M2(18-60) in POPC and DPhPC,<sup>39-42</sup> as well as the M2FL protein in Escherichia coli membranes,43 suggesting that the histidine tetrad exhibits a dimer of dimer conformation. However, the M2 conductance domain M2(22-62) in viral-envelope-mimetic lipid membranes shows a single set of His37 resonances and did not bind amantadine.<sup>29</sup> Nevertheless, the <sup>1</sup>H-<sup>15</sup>N HETCOR spectra from either the M2FL in DOPC/DOPE<sup>31</sup> or the truncated M2 protein in viral-envelope-mimetic lipid membranes<sup>30</sup> show correlations between water and His37 imidazolium nitrogen. These results indicate that water molecules are involved in proton conductance, although their interpretations yielded different conductance mechanisms. A direct measurement of the water-protein chemical exchange may shed light onto how the water with hydronium ions interacts with the protons in the His37 tetrad.

Figure 1a shows schematics for water—protein chemical exchange, where M represents water molecules, I is the specific proton in the protein that chemically exchanges with

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**Figure 1.** (a) Schematics for the water-protein chemical exchange system. (b) Pulse sequence used to probe specific water-protein exchange via indirect <sup>15</sup>N observation. (c) Simulated buildups of the dipolar-dephased <sup>1</sup>H magnetization I as a function of  $t_{SL}$  at various exchange rate constants  $k_{IM}$  using  $T_{I\rho}^{H} = 16.2$  ms and  $p = 10^{-6}$ .

hydronium ions at an exchange rate constant of  $k_{IM}$  and S is a spin such as <sup>15</sup>N that covalently bonds with the specific proton. With the hydronium ion concentration p in the pool of M, the exchange rate constant  $k_{MI}$  from M to I is  $pk_{IM}$ . Thus, the exchange process can be characterized by the classical Solomon equations<sup>44</sup> when the <sup>1</sup>H magnetization is spin-locked for a period of time  $t_{SL}$  along the magic angle (MA) by a Lee–Goldburg (LG) sequence:<sup>45</sup>

$$\frac{\mathrm{d}M(t_{SL})}{\mathrm{d}t_{SL}} = -\left(\frac{1}{T_{1\rho}^{M}} + pk_{IM}\right)M(t_{SL}) + k_{IM}I(t_{SL}) \tag{1}$$
$$\frac{\mathrm{d}I(t_{SL})}{\mathrm{d}t_{SL}} = pk_{IM}M(t_{SL}) - \left(\frac{1}{T_{1\rho}^{I}} + k_{IM}\right)I(t_{SL}) \tag{2}$$

Here, *M* and *I* represent the <sup>1</sup>H magnetizations along the MA for water and specific protein protons, respectively. The cross relaxation among protons is neglected because the <sup>1</sup>H spin diffusion is sufficiently suppressed along the MA<sup>45</sup> (Figures S1 and S2). For simplicity, we assume that the *M* and *I* protons have the same spin–lattice relaxation time in the LG spin-lock (LGSL) field, i.e.,  $T_{1\rho}^{M} = T_{1\rho}^{I} = T_{1\rho}^{H}$ .

Figure 1b shows the pulse sequence for probing the specific water—protein chemical exchange through indirect observation of <sup>15</sup>N signals. After <sup>1</sup>H excitation, a rotational-echo double-resonance (REDOR)<sup>23</sup> based dipolar dephasing scheme is used to prepare the initial <sup>1</sup>H magnetizations, M(0) and I(0). A 35.3° pulse flips the <sup>1</sup>H magnetizations to the MA, followed by a LG sequence<sup>45</sup> to spin-lock the <sup>1</sup>H magnetizations, during which the water—protein chemical exchange takes place. The *S* spin is then brought into a short contact with the *I* protons by applying the ramped rf field on the *S* spin such that the cross-polarization (i.e., LGCP) is established to transfer the *I* magnetization into the *S* spins.<sup>46,47</sup> Thus,  $I(t_{SL})$  can be monitored indirectly by its nearby *S* spins as a function of  $t_{SL}$ .

Without dephasing I(0), as documented in eq S9,  $I(t_{SL})$  contains the chemical exchange term that is scaled by the population difference between the hydronium ions and the specific proton involved in the chemical exchange, and thus is not sensitive to their exchange process. When I(0) is dephased by REDOR before LGSL,  $I(t_{SL})$  is derived as

$$I(t_{SL}) = pM(0)\{1 - \exp[-(p+1)k_{IM}t_{SL})]\}$$
  
$$\exp(-t_{SL}/T_{1\rho}^{H})/(p+1)$$
(3)

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Clearly,  $I(t_{SL})$  builds up through the chemical exchange process  $k_{IM}$  and is proportional to the population of the hydronium ions in water, as shown in Figure 1c.

Figure 2 shows the  ${}^{15}N$  spectra of the His37-labeled M2FL in lipid bilayers at -10 °C using Figure 1b. Clearly, the  ${}^{15}N$ 



**Figure 2.** Expanded <sup>15</sup>N spectra of the His37-labeled M2FL (pH 6.2) in lipid bilayers at -10 °C without (black) and with (red) <sup>15</sup>N-dipolar dephasing at  $t_{SL}$  of 100 and 2000  $\mu$ s.

spectra (black) without <sup>15</sup>N dephasing show similar line-shapes and intensities at different  $t_{SL}$  values. With a short LGCP contact time (i.e., 200  $\mu$ s), the <sup>15</sup>N signals were only crosspolarized from the protonated <sup>15</sup>N sites of His37 side chains (i.e., the  $\tau$  state N $\varepsilon 2\tau$  and the charged state N $\delta 1$ + and N $\varepsilon 2$ +). The nonprotonated  $\tau$  state N $\delta 1\tau$  could hardly be polarized (Figure S3). These observed <sup>15</sup>N resonances are spread from 165 to ~200 ppm, whereas their correlated <sup>1</sup>H frequencies extend up to 19 ppm.<sup>31</sup> Such high <sup>15</sup>N and <sup>1</sup>H frequencies indicate the formation of short imidazole-imidazolium Hbonds.<sup>48</sup> When the <sup>15</sup>N selective dephasing was applied, those protons that directly bond with <sup>15</sup>N become null before the LGSL so that their bonded <sup>15</sup>Ns will not be polarized. As shown in the red spectra of Figure 2, the <sup>15</sup>N signals were largely reduced at a short  $t_{SL}$ . Clearly from Figure 2, much of the spectral intensity that appears at ~165 ppm in black were not observed in red even when  $t_{SL}$  was long, indicating that these signals at ~165 ppm belong to N $\epsilon 2\tau$  that is less accessible by the hydronium ions. This assignment was confirmed by the <sup>15</sup>N-<sup>15</sup>N correlation spectrum (Figure S4). It is clear from the red spectra in Figure 2 that the dipolar-dephased <sup>15</sup>N signals from the charged His37 N $\delta$ 1+ or N $\epsilon$ 2+ gained more intensity as  $t_{SL}$  increased, implying that their bonded protons gain magnetization during the LGSL. As <sup>1</sup>H spin diffusion is suppressed during the LGSL and any relayed transfer is largely eliminated, the observed gain can only be facilitated by chemical exchange between this particular proton and the hydronium ions. It is worth noting that, since the protons of the charged His37 H-N $\delta$ 1+ and H-N $\epsilon$ 2+ sites were dephased at the beginning of the LGSL, no signals from the charged His37 N $\delta$ 1+ or N $\epsilon$ 2+ protons were expected. However, in the presence of the water-protein exchange, the proton from the charged His37 H-N $\delta$ 1+ or H-N $\epsilon$ 2+ (presumably H-N $\epsilon$ 2+) was re-energized during the LGCP, such that the signals from the charged His37 N $\delta$ 1+ or N $\epsilon$ 2+ are still observed. For a given LGCP contact time, these signals strongly depend on the exchange rate constant between the water and the specific proton involved. In addition, any incomplete HN dephasing by REDOR, due to various NH bond lengths, also attribute to these initial signals. Therefore, for the indirect observation via LGCP, eq 3 should be modified to include an additional constant, whose value depends on the exchange rate constant and the LGCP contact time. When the exchange is faster than the LGCP contact time, the dipolar-dephased I(0) reaches its exchange equilibrium, such that  $I(t_{SL})$  no longer depends on  $t_{SL}$ . Shown in Figure S5, the dipolar-dephased <sup>15</sup>N signals for the His37-labeled M2FL (pH 6.2) at +23 °C were almost identical at  $t_{SL}$  = 100 and 1000  $\mu s$ , except for the N $\epsilon 2\tau$  resonance at ~165 ppm that gains in intensity as  $t_{SL}$  increases. This implies that the N $\epsilon 2\tau$  site in the His37 tetrad becomes accessible to water at +23 °C, whereas the protons in the His37 NH bonds exchange rapidly with hydronium ions, whose exchange rate constant could be estimated by the LGCP contact time used in the experiments, i.e., 1/200  $\mu s = 5000 \text{ s}^{-1}$ .

A series of 1D dipolar-dephased <sup>15</sup>N signals obtained here allows us to monitor the transient water-protein chemical exchange processes. Figure 3 shows the plot of the dipolar-



**Figure 3.** Normalized dipolar-dephased <sup>15</sup>N integral intensities as a function of  $t_{SL}$  for the His37-labeled M2FL (pH 6.2) at -10 °C. The red line represents the fitting curve with  $k_{IM} = 1750$  s<sup>-1</sup>. The dashed green and blue lines are the curves for  $k_{IM} = 2250$  and 1250 s<sup>-1</sup>, respectively.

dephased <sup>15</sup>N signals versus  $t_{SL}$ . We obtained  $T_{1\rho}^{H} = 16.2$  ms in separate experiments and then fitted eq 3 to yield  $(1 + p)k_{IM} =$  $1750 \pm 552 \text{ s}^{-1}$ . The concentration p of the hydronium ions in the M2 channel pore is about  $10^{-6}{}_{M}$  with the assumption that the pH in the pore is the same as in the bulk environment. Thus, the exchange rate constant between hydronium ions and the protons in the His37 NH bonds for the M2FL is on the order of  $1750 \pm 552 \text{ s}^{-1}$  at -10 °C. This represents an average value over a number of different His37 states with various exposures to hydronium ions.

Figure 4 shows a model for the water-protein exchange process for the His37 NH bonds at the heart of the His37 tetrad. The "initial" NH protons in the His tetrad are colored green and blue for the His C-D pair. The hydronium ion is attracted by the nonprotonated N $\delta$ 1 site resulting in both His37 residues in a dimer becoming charged. The hydronium ion based proton is colored red. These two imidazolium residues conformationally rearrange, due to charge repulsion with the newly protonated N $\delta$ 1+H site oriented toward the pool of externally exposed waters, whereas the original N $\varepsilon$ 2+H and the newly formed N $\varepsilon$ 2+H are both exposed to waters of the viral interior (Figure S6). The return of the His N $\delta$ 1+H proton to the waters of the viral exterior results in a futile cycle, while the absorbance of either His N $\varepsilon$ 2+H protons by waters of the viral interior results in a successful transport of a proton across the membrane. If the proton in the original imidazoleimidazolium H-bond is reabsorbed by water (right-hand path), the imidazolium donates its H-N $\delta$ 1 proton to reform the same His37 C-D pair with an imidazolium-imidazole H-bond utilizing a  $\pi$  state. This proton rapidly rearranges crossing the H-bond barrier to form a  $\tau$ -charge H-bonded pair, back to the



Figure 4. Chemical exchange model between water and the NH protons in the His37 tetrad of the M2 proton channel.

original state. When an interior water reabsorbs the proton of the newly formed N $\varepsilon$ 2+H, D becomes a  $\pi$  state and forms a new imidazolium-imidazole H-bond with H-N $\delta$ 1 of His A (lefthand path) with the proton rapidly crossing the H-bond barrier to form a  $\tau$ -charge hydrogen bonded pair, leading to a rotation of the imidazolium-imidazole bonding pairs<sup>37</sup> (Figure 7S). This continual process for bringing the hydronium protons into the His37 NH is essential for the buildup of the dipolar-dephased <sup>15</sup>N signals as a function of  $t_{SL}$ . Thus, the buildup of the dipolardephased <sup>15</sup>N signals, as observed here, discriminates the proton shuttling mechanism<sup>29,30</sup> from the hydronium pore mechanism in the M2FL proton channel in lipid bilayers.

In summary, we have demonstrated the feasibility of probing specific water-protein chemical exchange in membrane-bound proteins via indirect observation of dipolar-dephased <sup>15</sup>N signals in solid-state MAS NMR by spin-locking the <sup>1</sup>H magnetization along the MA. Although hydrogen-deuterium exchange<sup>49</sup> characterizes the accumulation of a slow exchange process, here the dynamics on a sub-microsecond scale are documented for this proton transport mechanism. By suppressing the <sup>1</sup>H spin diffusion, the pure chemical exchange during the LGSL is monitored. To the best of our knowledge, this is the first direct observation of kinetic water-protein chemical exchange processes on the submsec time scale in membrane-bound proteins/peptides. Thus, this new technique provides an opportunity to characterize structure-function relationships of membrane-bound species at the water-bilayer interface, and in particular to understand the nature of how Hbonds are formed and broken in biological systems during proton transport in the M2 proton channel or other transporters.

## ASSOCIATED CONTENT

### **S** Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.6b08376.

Materials and experimental details; derivation of Solomon equations; control experiments; brief discussion on HETCOR; <sup>15</sup>N MAS NMR spectra; modeled His37 orientations upon breaking of the H-bond (PDF)

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#### Notes

The authors declare no competing financial interest.

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