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Role of water states on water uptake and proton transport in Nafion using molecular simulations and bimodal network

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1. Introduction

Optimal water management is critical to the operation of proton-exchange-membrane fuel cells (PEMFCs) [1–3]. Key in importance is keeping the electrolyte membrane, e.g., Nafion[®] hydrated to ensure good proton conductivity [4], while minimizing liquid water in other PEMFC layers for ease of gaseous fuel transport [5,6]. Hydration of Nafion and its transport properties are related to hydration-induced nanophase morphology, causing separation into hydrophobic-polymer-backbone and water-filled domains surrounded by hydrophilic sulfonic-acid side chains [4].

The water-filled domain size has a distribution of 1–1000 nm, with a mean size of $\langle L_d \rangle = 4$ nm [7–9], and this nanosize influences the water state and its transport properties. The surface energy from the domains influences the thermodynamic state of water, i.e., its chemical potential, and this directs the water adhesion toward the domain surface (adsorption). In particular, when the domain size is smaller than the intermolecular force-field distance (~1 nm), the surface force fields overlap each other, and this promotes water filling [10]. In addition, the domain size is much smaller than an average collision distance of the water molecules at

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ABSTRACT

Using molecular simulations and a bimodal-domain network, the role of water state on Nafion water uptake and water and proton transport is investigated. Although the smaller domains provide moderate transport pathways, their effectiveness remains low due to strong, resistive water molecules/domain surface interactions. The water occupancy of the larger domains yields bulk-like water, and causes the observed transition in the water uptake and significant increases in transport properties.

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near ambient pressure and temperature, and this significantly reduces their traveling distances (diffusion). The surface energy also impedes their motions, which in turn limits the transport, and this becomes more pronounced in the smaller domain size due to the overlapped surface force fields. So, these domain-sizedependent water states and their roles on transport properties are essential to the fundamental understanding of hydrated Nafion and optimal fuel cell design.

Nafion hydration and transport properties have been experimentally studied for water uptake [11-17], water self-diffusion [18-20], and proton conductivity [20,21]. These observed behaviors have been explained by both continuum and atomistic modeling [22-24] with the former being focused on macroscopic, thermodynamic descriptions [25-27], and the latter on hydrationassociated nanoscopic polymer dynamics [7,8,23,28-35]. Although both approaches have shown that the number of water molecules per hydrophilic, sulfonic-acid site, i.e., water content $\langle \lambda_{H_2O} \rangle$, correlate with the transport properties, their domain-size dependence has not been well explored, nor have these two approaches been correlated to each other. This lack of understanding has been in part due to limited methodologies. Macroscopic approaches have by nature relied on homogeneous treatments, and have been challenging to provide atomic-level insights on hydration and transport. Molecular approaches have also been limited to small spatial coverage for the domain-size distribution, and not extended



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to explain the domain-size-dependent macroscopic water uptake and its relation to transport properties. Thus, there is a critical need to bridge the two spectra. A previous study has attempted to bridge this gap using a macroscopic domain-network model, relating the adsorption-capillary water states in the Nafion domains to water uptake and proton conductivity [27]. However, this study utilized an empirical relation for the atomistic level water behavior, which is articulated here by rigorous molecular simulations. Along this line, we elucidate the roles of the domain-size-dependent water states on the water uptake and water/proton transport properties both in the individual domains and in the transport pathways throughout the domain network. In particular, we examine the nanoscopic water states and transport in presence of the surface force fields, as continuation of our previous, domain-sizedependent water distribution and its relation to the transport properties [6]. For this combined nano- and macroscopic water descriptions, we propose molecular simulations combined with a bimodal domain-size network model.

2. Proposed bimodal network model

2.1. Bimodal domain size

The microstructure of hydrated Nafion including its connectivity and hydrophilic-site distribution, is crucial to the water states, transport properties, and mechanical properties. Much effort has been made over last few decades to characterize the structure underlying its unique, high proton conductivity, however, due to its low crystallinity and complex chemical composition, an exact structure is still contentious [4], which will return to at the end of this section. Since the primary focus here is the water state and the transport properties, an idealized bimodal-domain distribution network for hydrated Nafion is proposed using uniform density of the hydrophilic sites. Such a construct is guided by our small-angle X-ray scattering (SAXS) measurement, the reported domain-size distribution measurement [9], and the available bimodal clusternetwork model [7,8]. As shown in Fig. 1(a), the large domains are



Fig. 1. (a) The proposed bimodal network model, showing hydration-dependent morphological evolution as a function of water activity (or hydration). (b) The cluster-network model [7,8]. (c) The structural-inversion network model [41]. (d) The sandwich-like model [30]. (e) The percolation network model [31]. (f) The parallel, long-channel network model [34].

represented by the mean domain size, 4 nm, while the small ones are determined by the Debye screening length δ_D , (the distance over which charge carriers screen out the surface electric field) [36],

$$\delta_{\rm D} = \left(\frac{\epsilon_0 \epsilon_{\rm r} k_{\rm B} T}{4\pi e^2 \sum_i n_i z_{e,i}^2} \right)^{1/2},\tag{1}$$

where ϵ_0 and ϵ_r are the free-space electric and relative permittivities, respectively, k_B is the Boltzmann constant, *T* is the temperature, and n_i and $z_{e,i}$ are the number density and the number of conduction electrons per contributing ions, respectively. $\delta_D = 0.23$ nm is predicted using an equivalent weight (EW) of 1100 g/mol, a density of 2 g/cm³ [37], and the number densities of two types of ions, i.e., the sulfonic-acid and hydronium. The calculated Debye length is the hard sphere diameter, $d_{H_2O} = 0.3$ nm, meaning that the effective interaction length is one layer of adsorbed water. This length is also consistent with the predictions using the Pekar–Marcus relation [38] and the mean field Poisson–Boltzmann theory [39]. Using 1.5 adsorbed-water layer and the two confining domain surfaces, a 1 nm nanogap (domain size) is set as the small domain size.

The bimodal-domain sizes lead to bimodal water filling (complete and partial water occupation), which results in selective transport pathways through the water-filled domains in a network, as shown in Fig. 1(a). During the water uptake, thermodynamic equilibria among neighboring domains in the network require that the connected domains have the same chemical potential. This condition results in first filling of the smaller domains due to their overlapping surface force fields, and then filling of the larger domains because of weaker surface interactions. The weak surface interactions mean more of a bulk-like water phase since the subsequent adsorbed-water layers are formed outside the Debye screening length. This finding has also been reported in previous studies [23,40], and this study further articulates the nanoscale-water behavior using molecular simulations.

The microstructure of hydrated Nafion has been primarily examined to understand the unique structure that yields its high proton conductivity. The primary models are the bimodal clusternetwork [7,8], the structural-inversion [41], the sandwich-like [30], the percolation-channel [31], and the parallel-channels models [34]. It has been known that the mean characteristic size of the microstructure is 4 nm and its size increases with water activity (hydration). The graphical descriptions of these models are shown in Fig. 1(b)–(f), with water activity as a variable.

The bimodal, cluster-network model, Fig. 1(b), proposes two characteristic cluster-domain sizes, such that the primary clusters are connected by the secondary transport channels, which evolve with water activity and become the larger percolated transport channels. Although this model has been the most popular and is similar to ours, it does not contain the viable molecular descriptions and cannot support the SAXS measurement [34]. In the structuralinversion model, Fig. 1(c), the swelling and dissolution processes have been phenomenologically described, showing a possible structural-inversion between the water and the polymer structure at high water activity, but no quantitative study has been done to date. In the sandwich-like model, Fig. 1(d), a simple nanogap has been proposed to predict the SAXS experimental results by quantifying the hydration-dependent structural change (swelling), but the water states and the resulting water/proton transport have not been addressed. In the percolation model, Fig. 1(e), the phenomenological descriptions on water state and its transport have highlighted the Schröder paradox which has been an inexplainable, larger water uptake increase ($\langle \lambda_{H_2O} \rangle = 14$ and 22) between fully vapor-equilibrated and liquid-equilibrated membranes at the same

chemical potential, and attempts have been made to relate it to the capillary percolation. However, capillary percolation occurs at lower hydration $\langle \lambda_{H_2O} \rangle = 5$ or 6, due to the strong surface interactions [6]. In the long parallel-channel model, Fig. 1(f), the proposed, long, cylindrical-domain network including semicrystal structures has best fitted the SAXS results, but the model has not been related to the nanoscopic descriptions on water states and the transport properties. It has also been found that the microstructure changes with hydration, temperature, and pretreatment, meaning that various morphologies may exist under different conditions [42]. However, the structure-related, nanoscale-water behaviors have not been addressed in any of the existing Nafion morphology models.

The key outcomes of this study rely on the nanoscale-water states and dynamics in the domains, and the resulting transport pathways in the network. These phenomena are related to the strength of the water/surface interactions which are affected by the microstructure. The spherical [7,8] and cylindrical [34] structures increase the specific surface area (total surface area per unit volume) compared to our nanogap. The increased surface area reduces the site density per unit surface area, and in turn weakens the surface force fields. This is expected to result in decreased water adsorption, reluctant water filling, and increased free-water motion. On the other hand, the spherical and/or cylindrical structures increase the surface curvature where the surface force fields overlap, which in turn strengthens them. The stronger fields lead to increased adsorption, favorable water filling (by reducing the capillary meniscus in the Kelvin–Cohan relation [43]), and hindered water motion. The two geometric effects above result in an expected, negligible net impact. It is implied that the nanoscalewater descriptions in our nanogap structure represent the water states and transport properties of hydrated Nafion without a significant loss of generality.

2.2. SAXS Measurement

The bimodal-domain size are further informed by our analysis of SAXS measurements. Nafion 117 specimens were tested using SAXS at the Advanced Light Source at the Lawrence Berkeley National Laboratory, with a X-ray wavelength of 1.239 Å. In preparation for the tests, the samples were boiled for 1 h in a 1 M H₂SO₄ solution to remove any ionic impurities and to ensure a fully protonated state. Samples were mounted in a sealed sample holder such that they were suspended above a pool of pure water, saturated LiCl solution, or desiccant, and allowed to equilibrate over 4 h with data taken every 5 min. The sample holder is shown in Fig. 2(a). Kapton-covered holes through the holder allowed unimpeded X-ray access to the hanging sample while maintaining a sealed environment. Background spectra were collected for each test and assumed to apply for the duration of the drying data gathering. The measured scattering X-ray intensities were subtracted from the background scattering, and then were normalized by the transmitted intensity. Fig. 2(b) shows the scattered intensities (arbitrary unit) with respect to the scattering wave vector (q = 0.05 to 0.3 Å⁻¹) at water activities of $a_{H_2O} = 0$, 0.11, and 1.0. One (ionomer) peak is observed for dry Nafion $(a_{H_2O} = 0)$, and this shifts to lower q with an increase in water activity (q = 0.185 to 0.15 Å⁻¹). This is primarily related to the water-filled, domain size growth by swelling (the domain size change of 3.3-4.2 nm) [4,7]. Indeed, the intensity at the low q (matrix knee) is related to the semicrystalline structures from long polymer matrix chains, which is important to its mechanical properties, while the intensity at the large q (Prod's law) represents the domain surface area [34,44]. In addition, we note that the water morphologies in the domains **a** Nafion Holder for SAXS Measurement



b SAXS Measurements and Predictions





Fig. 2. (a) Holder for SAXS measurement. (b) Measured, normalized SAXS intensity $I^*(\mathbf{q})$ as a function of the scattering wave vector \mathbf{q} for Nafion 117 at $a_{H_{2O}} = 0, 0.11, \text{ and } 1, T = 298 \text{ K}, \text{ and } p = 1 \text{ atm. Variations of the predicted intensities at the peaks using Eq. (2) are also shown. (c) The proposed cubical domains for the polymer matrix, the adsorbed-water layer, and the water-fillable layer.$

(partial/complete water occupation) can be another source of the peak shift, although the thin, adsorbed-water layers are the secondary X-ray scatters.

To model this, cubical domains are constructed in a network, as shown in Fig. 2(c). Each domain includes polymer matrix, an adsorbed-water layer, and a water-fillable layer as sources of the X-ray scattering. The last layer represents the potential, waterfilling layer at high water activity and is not expected to influence significantly the water transport due to the absence of surface interaction with water. Practically, this water-fillable layer is occupied by the polymer matrix, where it swells/shrinks with hydration. However, we use idealized, rigid domains, along with the water-filling layer, for feasible molecular simulations. The rigorous descriptions associated with the adsorbed/water-fillable requires modifications of the existing sandwich-like model [30], and after adding these the layers, we have

$$I^{*}(q) = \alpha \int_{0}^{1} \frac{2}{\pi} \int_{0}^{\pi/2} S^{2}(L_{X}, \phi, a) d\phi \Big[\Delta n_{e,p} \big(S(L_{Z}, a) - S(L_{Z,d}, a) \big) \\ + \Delta n_{e,ad, H_{2}O} \Big(S(L_{Z,d}, a) - S(L_{Z,wf}, a) \Big) \\ + \Delta n_{e,g, H_{2}O} S(L_{Z,wf}, a) \Big]^{2} da,$$
(2)

where

$$S^{2}(L_{X},\phi,a) = \frac{\sin\left[0.5qL_{X}\cos(\phi)(1-a^{2})^{1/2}\right]}{0.5qL_{X}\cos(\phi)(1-a^{2})^{1/2}}$$
$$\frac{\sin\left[0.5qL_{X}\sin(\phi)(1-a^{2})^{1/2}\right]}{0.5qL_{X}\sin(\phi)(1-a^{2})^{1/2}}L_{X}^{2},$$
(3)

$$S(L_Z, \phi, a) = \frac{\sin(0.5qL_Z a)}{0.5qL_Z a} L_Z, \quad a = \cos(\phi),$$
(4)

where *S* is the form factor, Δn_e is the electron density variation, $L_{Z,d}$ is the domain size, $L_{Z,wf}$ is the water-fillable layer thickness, L_X is the lateral domain size, and α is the scaling factor. In the network, the individual domains are randomly rotated, and hydrated Nafion has a water-filled domain volume ratio, $\epsilon = 0.4$ [9,27,45]. Using $L_X = L_Z = 6$ nm guided by our experimental result at $a_{H_2O} = 1$ (Table 1), the domain population ratios are 0.53 and 0.47 and the domain volume ratios are 0.22 and 0.78, for $L_{Z,d} = 1$ and 4 nm, respectively, as summarized in Table 2. Using the bimodal-domain size, the predicted $L_{Z,wf}$ (which will be predicted using molecular simulations in the next section), and the domain population ratio, the predicted peaks for $a_{H_2O} = 0, 0.11$, and 1 show reasonable agreements with the measured data, particularly near q > 0.125 Å as shown in Fig. 2(b). The peak in q (characteristic length) is related to an average distance associated with the L_X or L_Z , $L_{Z,d}$ and $L_{Z,wf}$, and the domain-size distribution. The predictions using the parameters given in Table 1 agree with the measured peak at $q \sim 0.15$ to 0.185 Å⁻¹.

2.3. Bimodal water filling

The bimodal water filing is related to the bimodal strength of the water/surface interactions, and it is analyzed by the isosteric heat of adsorption [46]. Using Eq. (A.1), the heat of adsorption is calculated as a function of the number of adsorbed-water layers, N_{ad,H_2O} , for the two domain sizes, as shown in Fig. 3. The total adsorbed-water-layer thickness, $\delta_{H_2O,ad}$ (summation of two layers from top and bottom) using $d_{H_2O} = 0.3$ nm is also shown at the top *x*-axis. The Debye screening length is also marked to show water-behavior changes by the surface force fields. As the adsorbed-water layer grows beyond the Debye length, the heat of adsorption decreases toward the bulk value, i.e., the heat of evaporation (46 kJ/mol at T = 300 K for MSPC/E water model [47]), since the surface interactions are significantly screened. In the small domains, the heat of

 Table 1

 Summary of the SAXS model parameters for Eqs. (2)–(4).

<i>a</i> _{H₂O}	L _Z and L _X , nm	$L_{Z,wf}$, nm in $L_{Z,d} = 1$ nm	$L_{Z,wf}$, nm in $L_{Z, d} = 4$ nm	$\Delta n_{e,p}$, Å $^{-3}$	$\Delta n_{e,ad,\mathrm{H_2O}},$ Å ⁻³	$\Delta n_{e,g,\mathrm{H_2O}},$ Å ⁻³
0	4.8	1	4	-0.0875	0.02	-0.0045
0.11	5.0	0	3.7	-0.0875	0.02	-0.0045
1	6.0	0	0	-0.18	0.08	-0.1

 Table 2

 Ratio of domain populations and volumes

	$L_{Z,d} = 1 \text{ nm}$	$L_{Z,d} = 4 \text{ nm}$
$N_{L_{z,d},i}/N_d$	0.53	0.47
$V_{L_{Z,d},i}/V_{L_Z}$	0.22	0.78

adsorption is nearly doubled for small adsorption, and is still higher compared to the bulk heat of evaporation at large adsorption. In contrast, in the large domain it is only 15% higher for the small adsorption and then becomes nearly the same as the bulk. This indicates that the small domains significantly change water behavior due to strong interactions, whereas the large domains lead to bulklike water due to no significant surface interactions. In addition, an increase in sulfonic-acid-site density, $n(SO_3^-) = 1.75$ to 2.5 nm⁻², strengthens the water interactions with the surfaces. This change is more pronounced in the larger domains because the smaller domains already have strong surface forces (overlapping force fields). This result implies that the water-state changes with respect to the hydrophilic-site concentration (or equivalent weight) are more significant in the larger domains than those in the smaller domains. The strength of the surface forces is also related to the water filling, which will be discussed further below.

3. Water uptake

The bimodal-size and water-filling model succeeds in explaining water uptake and its significant increase near high water activities. The water states of each domain size are explored using the grand canonical molecular dynamics-Monte Carlo (GCMD-MC) simulation, as described in Appendix A, and the results are integrated over the entire network through the prescribed chemical potential (water activity). One of the important factors for the water states in the domains is the surface sulfonic-acid density $n(SO_3^-)$. The reported values have a range of $0.51-3.3 \text{ nm}^{-2}$ as summarized in Table 3 [8,9,35,41]. Here, using $n(SO_3^-) = 1.5$, 1.75, and 2.5 nm⁻², the number of water molecules per volume of each domain, n_{HaO} , are



Fig. 3. Variations of the MD predicted isosteric heat of adsorption $\Delta h_{lg,ad}$ as a function of the number of adsorbed-water layers at the sulfonic-acid surface densities, $n(SO_3^-) = 1.75$ and 2.5 nm⁻², and the domain sizes, $L_{Z,d} = 1$ and 4 nm. The adsorbed-water thickness is also shown at the top *x*-axis. The bulk heat of evaporation for MSPC/ E water model at T = 300 K [47] and the Debye screening length δ_D are shown.

Table 3

Summary of various sulfonic-acid surface site densities in the literature.

$n(\mathrm{SO}_3^-)~\mathrm{nm}^{-2}$	Comments
1.62	110 sites per 4.65 nm diameter cluster
	for EW = 1100 g/mol [8]
3.3	$d_{\rm SO_3^SO_3^-} = 0.55$ nm for $T = 20-80$ °C [9]
1.82	the specific surface area of 0.55 nm ² per site [41]
0.51 to 1.23	$d_{\mathrm{SO}_3^\mathrm{SO}_3^-}=0.9$ to 1.4 nm for $\langle\lambda_{\mathrm{H}_2\mathrm{O}} angle=4$ to 15 [35]

shown in Fig. 4(a) and (b) at T = 300 K for $L_{Z,d} = 1$ and 4 nm, respectively. The GCMD-MC snapshots of completely/partially water-filled domains at the activity of 0.3 are also shown. Note that $n_{\rm H_2O} = 37 \text{ nm}^{-3}$ represents complete water filling. At low activities, there is only adsorbed-water layer without water filling, and its thickness grows as the activity increases. At the threshold activities, the domain begins to fill completely. The threshold activity increase in the large domains due to the weaker overlapping surface forces. However, no significant variation in the threshold is found in the small domains because of the strongly overlapped surface force fields. These predicted, nanoscale-water states are validated by the available modified BET theory, the Kelvin-Cohan (K-C) relation [27,48], and the Saam-Cole (S-C) transition theory (using the homogeneous water-water effective potential for the water-surface wall interactions beyond the Debye length [6,10]). Although these theories reasonably predict the water filling of the large domains, the prediction using Kelvin-Cohan relation is close to the GCMD-MC results with a large sulfonic-acid-site density i.e., $n(SO_3^-) = 2.5 \text{ nm}^{-2}$. The Saam–Cole theory predicts a delayed threshold. This delay is due to the large (bulk) surface tension, while the effective water-water intermolecular potential tends to be weaker than the surface-water interaction.

The sulfonic-acid-site density influences the adsorption and water filling. Higher surface energy or hydrophilic-site density (strong surface force, $n(SO_3^-) = 2.5 \text{ nm}^{-2}$ increases adsorption, and the water filling occurs at lower activity. However, the lower density (weak surface force, 1.5 nm^{-2}) decreases adsorption, and the filling does not occur at $a_{H_2O} = 1$, indicating that the surface is extremely hydrophobic and the expected water uptake with the fully vapor-equilibrated reservoir is significantly lower than the experimental results. Thus, we choose $n(SO_3^-) = 1.75 \text{ nm}^{-2}$ as the characteristic Nafion sulfonic-acid-site density for a good agreement with the available experimental water uptake data as well as the available water diffusivity and proton conductivity as shown below.

In general, the adsorbed water in Nafion is presented as water content defined by the number of water molecules per sulfonicacid site [21]. To predict the water content, $\langle \lambda_{H_2O} \rangle$, the total adsorbed water amount in the network is calculated by integrating the water states in the individual domains over the two domain sizes, as given by

$$\langle \lambda_{\rm H_2O} \rangle = \frac{\sum_{i} n_{\rm H_2O,ad} V_{L_{Z,d,i}}}{N(\rm SO_3^-)}, \tag{5}$$

where $n_{\text{H}_2\text{O},\text{ad}}$ is the number density of the adsorbed water in the individual domain, $V_{L_{z,d,i}}$ is the domain volume for each, and $N(\text{SO}_3^-)$ is the total number of sulfonic-acid sites. The predicted water content is shown in Fig. 4(c). It is small at low water activity, and increases with increasing water activity. At the threshold water activities, $a_{\text{H}_2\text{O}} = 0.1$ and 0.95, the water uptake significantly increases, which is related to the filling of the large domains. The first transition is caused by filling the smaller domains, $L_{Z,d} = 1$ nm, and the second is more pronounced by fully hydrating the larger



Fig. 4. (a) Variations of GCMD-MC results of the adsorbed water at $L_{Z,d} = 1$ nm with respect to the water activity, a_{H_2O} at T = 300 K and $n(SO_3^-) = 1.75$ nm⁻². The predicted results using available adsorption and water filling theories are also shown [6,10,27]. The GCMD-MC snapshot of the completely water-filled domain at the activity of 0.3 is also shown. (b) Variations of the adsorbed water at $L_{Z,d} = 4$ nm, T = 300 K, and $n(SO_3^-) = 1.5$, 1.75, and 2.5 nm⁻². The GCMD-MC snapshot of the patially water-filled domain at the activity of 0.3 is also shown. (c) Variations of the predicted water content $\langle \lambda_{H_2O} \rangle$, Eq. (5), as a function of the activity at T = 300 K. Available experimental result at T = 303 K is also shown [21].

domains, $L_{Z,d} = 4$ nm, due to a significant domain volume ratio. This agrees with the experimental result [21], especially at the large water content, while overpredicting at low water activities. The assumption of a rigid domain structure based on the fully hydrated Nafion causes the discrepancy at the low water activities. In addition, our model does not include hydrodynamic resistance among domains, or interfacial rearrangement and/or resistance [17,49,50], which also help to explain this overprediction.

4. Water self diffusion

Water self-diffusion occurs through the completely water-filled transport pathways [Fig. 1(a)]. Hydration controls not only nanoscale-water motions (local diffusivity) in the individual domains, but also the transport channel connectivity (or tortuosity of transport pathways). In the individual domains, the diffusivities, D_{H_2O} are calculated using the molecular simulations and the Green–Kubo autocorrelation, Eq. (A.2), at T = 300 K as a function of a_{H_2O} , and the results are reported in Table 4. Next, the effective diffusivity $\langle D_{H_2O} \rangle$ is calculated using 30×30 domains in the 2-D bimodal network as depicted in Fig. 1(a), and the results are shown in Fig. 5(a) as a function of the predicted water content [Fig. 4(c)]. No calculation size effect is found up to 100×100 domains. Calculation details have been given previously [51].

At $\langle \lambda_{\rm H_2O} \rangle \sim 2(a_{\rm H_2O} \sim 0.05)$, the effective diffusivity is significantly low compared to that in bulk liquid water because the movement of water molecules are significantly hindered by the strong surface force fields (most of water molecules are within the Debye screening length, Table 4). The transport pathways are also significantly limited by the partially-filled domain network. At $\langle \lambda_{\rm H_2O} \rangle \sim 4 (a_{\rm H_2O} \sim 0.1)$, the effective diffusivity begins to increase primarily by approaching the bulk-like diffusivity in the waterfilled small domains and by increasing the transport pathways. At $\langle \lambda_{\rm H_2O} \rangle \sim 5(a_{\rm H_2O} \sim 0.3)$, the adsorbed layers in large domains grow beyond the screening length, and for $\langle \lambda_{H_2O} \rangle > 5$, further filling of the large domains increases the effective diffusivity. These results are in good agreement with experiments [18] and available MD results [52,53], and our multiscale approach better predicts compared to the full atomistic simulations [52,53], where the limited simulation size may not represent the domain-sizedependent water transport. With a consideration of hydrationdependent Nafion morphology (swelling), i.e., reduced domain sizes at low water activity, the water uptake is expected to decrease especially at the low water content, and this consideration results in better agreement with the experimental data. For validation, our MD-predicted bulk-liquid diffusivity at T = 298 K is in excellent agreement with existing data [54].

The role of the water-filled transport channels on the effective water diffusivity is elucidated by examining the pathway tortuosity. In permeable media, the effective self diffusivity $\langle D_{H_2O} \rangle$ is related to

Table 4

MD predicted D_{H_20} , as a function of a_{H_20} and $\langle \lambda_{H_20} \rangle$ at T = 300 K for $L_{Z,d} = 1$ and 4 nm.

$a_{\rm H_2O}(\langle\lambda_{\rm H_2O}\rangle)$	$D_{\mathrm{H}_2\mathrm{O}}$, m ² /s at $L_{Z,d}=1~\mathrm{nm}$	$D_{\mathrm{H_2O}}$, m ² /s at $L_{Z,d}=4~\mathrm{nm}$
0.05 (2.0)	$5.70 imes 10^{-10}$	5.82×10^{-10}
0.1 (3.9)	1.21×10^{-9}	$7.64 imes 10^{-10}$
0.3 (4.8)	1.25×10^{-9}	2.13×10^{-9}
0.5 (5.4)	1.27×10^{-9}	2.25×10^{-9}
0.7 (6.3)	$1.29 imes 10^{-9}$	2.45×10^{-9}
0.8 (6.9)	$1.29 imes 10^{-9}$	3.03×10^{-9}
0.9 (7.2)	1.30×10^{-9}	3.20×10^{-9}
1.0 (13.1)	$1.29 imes 10^{-9}$	2.80×10^{-9}



Fig. 5. (a) Variations of water self-diffusion coefficient in Nafion as a function of average water content at 303 K. The experiential results [18] and the available MD results [52,53], are also shown along with the experimental result for bulk liquid water at T = 298 K [54]. (b) Variations of the tortuosity with respect to the water content using the normalized effective diffusivity. The domain water-filling regimes are also marked.

the bulk diffusivity D_{H_2O} , permeable domain volume ratio ϵ , and tortuosity L_t^* , as given as [55]

$$\left\langle D_{\rm H_2O}\right\rangle = \frac{D_{\rm H_2O}\epsilon}{L_t^*}.$$
 (6)

The tortuosity is calculated using the homogeneous, bulk diffusivity for all the bimodal domains with a constant domain volume ratio of $\epsilon = 0.4$ as a function of water content. Note that $L_t^* = 1$ represents a full transport-network connectivity, which occurs at high water activity. Using the bulk diffusivity for all the domains, it predicts only the tortuosity without considering the surface-force-induced transport, as shown in Fig. 5(b). The tortuosity decreases with increasing water content, showing that the smaller domains form minor transport channels at low water content, and the larger domains provide major domain connectivity. Using the molecular-simulation-calculated diffusivity (surface-force-hindered diffusivity, Table 4), the tortuosity predicts 30–70% lower than the above predictions (without considering the surface-force hindrance). This resistance is caused by the transport impedance from the domain–surface interactions. Note that the

resistance difference is minimized where the water fillings occur, indicating that the complete water occupation yields the bulk-like water behavior.

5. Proton transport

In bulk liquid water, protons transfer via Grotthuss (or hopping) diffusion and translational diffusion (or vehicle motion). The former is the dominant mechanism, where a proton effectively hops over a hydrogen-bond network of water molecules using fast bond-breaking and making steps [56–60]. In the hydrated Nafion domain network, such transport is hindered by the strong surface force fields, and the effective proton conductivity decreases. Since the force fields are significant within the Debye length, no significant Grotthuss diffusion is considered, and the surface-force hindered translational hydronium diffusion is included (Table 5). Using the Nernst–Einstein relation [25,26], we have

$$\sigma_{\rm H^+} = \frac{e_{\rm H^+}^2}{k_{\rm B}T} \Big(n_{\rm H_3O^+} D_{\rm H_3O^+,D} + n_{\rm H^+} D_{\rm H^+,G} \Big), \tag{7}$$

where $e_{\rm H^+}$ is the proton charge, $n_{\rm H^+}$ is the number density of protons for Grotthuss diffusion, $n_{\rm H_3O^+}$ is the number density of hydronium for translational diffusion, $D_{\rm H^+,G}$ is the Grotthuss diffusivity, and $D_{\rm H_3O^+,D}$ is the translational diffusivity. $D_{\rm H_3O^+,D}$ is calculated using MD-simulated, 8 to 18 hydroniums, and the Green–Kubo autocorrelation relation Eq. (A.2), as presented in Table 5. The Grotthuss diffusivity is calculated using the available relaxation time for water-molecule rotations [25,26,61]. At T = 300 K, $D_{\rm H^+,G} = 7 \times 10^{-9} {\rm m}^2/{\rm s}$, using $D_{\rm H^+,G} = d_{\rm O} - O^2/6\tau_{\rm D}$, and relaxation time of $\tau_{\rm D} = 1.5$ ps, and O–O distance of 0.255 nm in H₉O₄⁺ [61]. For the total proton number density, equivalent weight of 1100 g/mol, and density of 2 g/cm³ are used, while for the bulk-like domains, $n_{\rm H^+}/(n_{\rm H^+} + n_{\rm H_3O^+}) = 0.5$ is assumed. We note that this population density ratio is so far not confirmed, even with the recent microscopic analyses and *ab initio*-molecular-dynamics simulations [56–60].

Similarly, using the 2-D, 30 \times 30 domain network and the Kirchhoff law at the domain junctions [27], the effective conductivity $\langle \sigma_{H^+} \rangle$ is calculated as a function of the water content $\langle \lambda_{H_2O} \rangle$, as shown in Fig. 6(a).

In Fig. 6(a), at $\langle \lambda_{H_2O} \rangle < 2$, proton conductivity is negligibly small, because the adsorbed water is mainly within the screening length. As the activity increases, the small domains readily fill, and at $\langle \lambda_{H_2O} \rangle \sim 4$ the proton conductivity significantly increases through bulk-like proton diffusion and hopping as marked by the onset of hopping at $L_{Z,d} = 1$ nm. However, in the large domains, the conductivity is still hindered, which in turn causes no significant proton conductivity increase. At $\langle \lambda_{H_2O} \rangle \sim 8$, the adsorbed-water layers in the large domains grow beyond the screening length,

Table 5
MD predicted $D_{H_3O^+,D}$, as a function of a_{H_2O} and $\langle \lambda_{H_2O} \rangle$ at $T = 300$ K for $L_{Z,d} = 1$ and
4 nm.

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$a_{\rm H_2O}(\langle\lambda_{\rm H_2O}\rangle)$	$D_{\mathrm{H_3O^+}}$, $\mathrm{m^2/s}$ at $L_{Z,d}=1~\mathrm{nm}$	$D_{\mathrm{H_3O^+}}$, $\mathrm{m^2/s}$ at $L_{Z,d}=4~\mathrm{nm}$
0.05 (2.0)	3.89×10^{-11}	3.65×10^{-11}
0.1 (3.9)	$2.25 imes 10^{-10}$	9.60×10^{-11}
0.3 (4.8)	$2.38 imes 10^{-10}$	2.62×10^{-10}
0.5 (5.4)	$3.07 imes 10^{-10}$	3.13×10^{-10}
0.7 (6.3)	$3.58 imes 10^{-10}$	3.58×10^{-10}
0.8 (6.9)	4.29×10^{-10}	3.69×10^{-10}
0.9 (7.2)	4.90×10^{-10}	1.22×10^{-9}
1.0 (13.1)	5.25×10^{-10}	1.51×10^{-9}



Fig. 6. (a) Variations of the predicted proton conductivity, as a function of water content, showing significant increases caused by water filling and bulk-like diffusivity, at T = 300 K. The predicted translational-diffusion proton conductivity, and the available experimental results at T = 303 K [21] and MD results T = 298 K [62] are shown. (b) Predicted domain connectivity with respect to the water content, and comparison with the normalized proton conductivity at T = 303 K. The domain water fillings are also marked.

and the bulk-like diffusion and hopping result in significant enhancement as indicated by the onset of hopping at $L_{Z,d} = 4$ nm. The detailed discussions on the water-filling related transport property transition are found in our previous work [6]. At high activities, water filling occurs and leads to the maximum conductivity. The diffusive proton conductivity is also shown, indicating that its contribution is only nearly one third. For comparison, a previously reported MD result is also shown, and it can be seen that it underestimates the proton conductivity without addressing the significant enhancement at $\langle \lambda_{\rm H_2O} \rangle \sim 5$ since it has not properly considered the domain-size-dependent transport properties [62]. Similar to Fig. 5(b), the normalized effective conductivity is shown in Fig. 6(b). At low water content, $\langle \lambda_{\rm H_2O} \rangle \sim 2$, the proton conductivity is a minimum due to the large hindrance from the ionic surface although there are water pathways through the smaller domains (the pathways are small as indicated by the small domain filling). At $\langle \lambda_{\rm H_2O} \rangle \sim 8$, the water molecules are outside the Debye length especially in the larger domains, and the conductivity increases significantly.

6. Conclusions

Using the proposed molecular simulations combined with the bimodal-domain-network model, we explained the role of water states on the water uptake and water and proton transport in Nafion, while addressing both nano- and macroscopic water behavior. The water states, i.e., adsorption and water filling, are domain-size dependent in the individual domains, and the water occupation forms selective transport pathways in the network. Although the smaller domains readily fill water and form network pathways, the strong overlapping surface forces retard the water and proton transport. However, in the larger domains, the water filling is delayed and their water occupation significantly increases both the water uptake and water and proton transport properties since the water molecules are beyond the Debye transport impeding length. Although idealized, this model provides nanoscopic insights on water states and the resulting transport, while addressing the macroscopic water description. The domain-sizedependent water descriptions contribute to complimentary understanding of the role of water states on water/proton transport in ion-conducting polymers. In addition, our grand canonical molecular dynamics - Monte Carlo (GCMD-MC) approach allows for predictions of the water content as a function of water activity, something that is challenging for conventional MD simulations. This methodology allows not only a direct comparison with wateruptake experiments, but also allows for extending the treatment to intrinsic, temperature-dependent water uptake with inclusion of the flexibility of the polymer matrix.

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Nomenclature

- A: cross-section area (m^2)
- a_{H_2O} : water activity
- D: diffusivity (m²/s)
- d: diameter (m)
- h: enthalpy (J/mol)
- k_B : Boltzmann constant (J/K)
- L: length or domain size
- *L_t*: tortuosity
- N: number of fluid particles, water layers, or domains
- *n*: number density (m^{-3})
- p: pressure (Pa)
- T: temperature (K)
- Greek symbols

 α : scaling factor

- €: water-filled domain volume ratio
- δ : thickness (m)

 $\begin{array}{l} \mu: \text{ chemical potential (J)} \\ \lambda_{H_2O}: \text{ water content} \\ \rho: \text{ density } (\text{kg/m}^3) \\ \sigma_l: \text{ surface tension } (\text{N/m}) \\ \sigma_{H}^+: \text{ proton conductivity } (\text{S/m}) \\ \tau: \text{ time } (\text{s}) \end{array}$

Subscripts ad: adsorption *d:* domain e: electron eff: effective f: fluid or water H_2O : water molecules H_3O^+ : hydronium lg: liquid—gas phase change, or saturation p: polymer so₃: sulfonic-acid wf: water-fillable X, Y, Z: coordinates of x, y, and z