Characterizing the hydrophobic-to-hydrophilic transition of electrolyte structuring in proton exchange membrane mimicking surfaces†

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The surface density of charged sulfonic acid head groups in a perfluorosulfonic acid (PFSA) proton exchange membrane determines the hydrophilicity of the ionic channels and is thus critical for the structuring and transport of water and protons. The mechanism through which the head group density affects the structuring of water and ions is unknown, largely due to experimental challenges in systematically varying the density in an appropriate model system resembling the ionic channels. Here, we present a model system for PFSA membrane ionic channels using self-assembled monolayers with a tunable surface density of sulfonic acid and methyl groups to tune surface hydrophilicity. Atomic force microscopy force–distance measurements were used to quantify the hydration forces and deduce the interfacial electrolyte structure. The measured force profiles indicate a pronounced change of the electrolyte layering density at the surface with an unexpectedly sharp hydrophobic-to-hydrophilic transition when the surface shows a contact angle of ~37°. Using an extended Derjaguin–Landau–Verwey–Overbeek model including the Hydra force, we quantify diffuse double layer charges and characteristic hydration lengths as a function of sulfonic acid group density on the surface. Translating our results to PFSA membranes, these findings have two implications: (1) the density of sulfonic acid head groups can have a dramatic effect on the local solvent structuring of water inside the ionic channels and (2) they support a view where two types of water (solution) exist in PFSA ionic channels – a structured (shell/surface) and a non-structured (bulk) water. This offers an interesting perspective on how different head group densities lead to changes in water and proton transport and macroscopic membrane conductivity properties based on hydration layer characteristics.

Introduction

Perfluorosulfonic acid (PFSA) proton exchange membranes (PEMs) are one of the key components for PEM fuel cells (PEMFCs) and largely determine the performance of PEMFCs.† Nafion membranes are the most widely used PFSA membranes in PEMFCs because of their outstanding chemical stability for long-term use and reasonable proton conductivity. Improving the proton conductivity of Nafion to optimize PEMFC performance is an active area of research. In a Nafion membrane, hydrophilic sulfonic acid head groups (–SO3H) aggregate to form ionic channels of about 3–6 nm in diameter that are distributed in a hydrophobic Teflon matrix. In different versions of Nafion, the degree of hydrophilicity of the channels is tuned by a variation of the ~SO3H density in the polymer. While water and proton transport are known to occur through these ionic channels, the fluid structure in the channels and the exact transport mechanism are not well understood at the molecular scale, which hinders bottom-up design of PEMs with improved transport properties.

Previous work by our group and others has reported that the chemical constitution of the ionic domains – particularly the surface density of sulfonic acid groups – in Nafion membranes strongly affects water structuring, water transport, and proton transport under fully hydrated conditions. However, these works were based on Nafion systems with given ~SO3H densities, and predictions of the effect of sulfonic acid group density on structural and transport membrane properties has emerged almost exclusively from theory. To experimentally elucidate the influence...
of sulfonic acid group surface density on the structural and transport properties of Nafion membranes, an appropriate model system for the ionic surfaces that allows systematic variation of the surface hydrophilicity/hydrophobicity\(^5,\!^12\) combined with a method to measure liquid structures and surface interactions on the molecular scale is required.

To emulate the pore environment inside Nafion ionic channels, we functionalized gold surfaces with short aliphatic thiols terminated either by –SO\(_3\)H (hydrophilic head group) or by –CH\(_3\) (hydrophobic head group) with a systematic variation in hydrophilicity (–SO\(_3\)H/–CH\(_3\) ratio) to span degrees of sulfonation that can be found in conventional PEMs. We analyzed the effect of surface hydrophilicity on the hydration forces between two opposing, similarly functionalized surfaces in an aqueous acidic electrolyte with atomic force microscopy (AFM) force–distance measurements, which have proven to be effective in characterizing fluid structuring between two surfaces.\(^13\)

In particular, surface interaction forces were measured, and hydrophilic and hydrophobic hydration were modeled according to the empirical Hydra model.\(^14\) With this AFM-Hydra model approach, we characterize the electrolyte structure between two opposing, functionalized Au surfaces of varying degree of sulfonation (hydrophilicity), which approximates the channel environment in PFSAs membranes.

### Materials and methods

All reactants were used as received unless otherwise stated. Solvents were MilliQ water (Millipore, resistivity of \(\geq 18 \text{ MΩ cm, TOC below 2 ppb}\), ethanol (\(\geq 99.8\%\), Sigma-Aldrich, Germany), and hexane (\(\geq 95\%\), Sigma Aldrich, Germany). Functionalized surfaces were prepared by immersing gold substrates and tips in ethanol solutions of sodium 3-mercaptopropanesulfonate (HS(CH\(_2\))\(_3\)SO\(_3\)Na, 90\%, Sigma-Aldrich, Germany) and 1-butanethiol (HS(CH\(_2\))\(_3\)-CH\(_3\), 98%, Alfa Aesar, Germany) and allowing the SAM to form overnight. 0.1 M HCl (32% w/w, Merck, Germany) electrolyte solutions were prepared and the pH was adjusted to 1 using diluted NaOH or HCl as required. The molar fraction (\(X_{\text{SO}_3\text{H}}\)) of HS(CH\(_2\))\(_3\)SO\(_3\)H in HS(CH\(_2\))\(_3\)-CH\(_3\) was varied between 0 and 1. The total molar concentration of thiols in each thiol–ethanol solution was 1 mM. Different Au-substrates were prepared by immersion into solutions with molar fractions given in Table 1. Additional details for preparing different Au-substrates and the measurements using these Au-substrates are shown in the ESI\(^\dagger\) (Table S1) in detail.

#### Atomic force microscopy (AFM)

AFM force curves were recorded on a JPK Nanowizard (JPK Instruments, Germany) at an approach speed of 0.5 \(\mu\)m s\(^{-1}\) in solution. The spring constant of the used cantilevers was determined using the thermal noise method to be between 250 and 500 pN nm\(^{-1}\). Two sets of 100 deflection–displacement curves were recorded at random positions on a given surface and converted into force–distance profiles consisting of one extend and one retract trace. This method was repeated using 3–4 sets of surfaces and tips for each molar fraction. The experimental setup is schematically shown in Fig. 1. \(R_{\text{eff}}\) is the effective tip radius, \(T_{\text{SAM}}\) is the thickness of the monolayer, \(D_t\) stands for the relative distance between the tip and the surface, and \(F\) is the force that is quantified by the AFM. Data fitting to the force–distance curves is explained in detail in the Results section and in the ESL\(^\dagger\).

#### Surface-enhanced Raman spectroscopy (SERS)

SERS-active Au-substrates were prepared by electrochemically roughening an Au-coated glass slide (see Experimental details in ESI\(^\dagger\)). The thickness of the pristine Au layer on the glass slide was 200 nm. Surfaces of varying degree of hydrophilicity were prepared by immersing the roughened Au-substrates in the respective thiol–ethanol solutions (Table 1) overnight. SERS maps were taken on a Bruker Senterra Raman microscope with an excitation wavelength of 785 nm. The laser power used in all experiments was 1 mW. A 50\(\times\), NA 0.75 objective (Olympus) was used to focus the laser onto the sample and to collect the SERS signal. The \(x \times y\) (11 \(\times\) 11) pixels spatial images were captured from three different spots of three independent samples for each X. The step size between two neighboring pixels was 0.2 \(\mu\)m in both \(x\)- and \(y\)-axes. All spectra were obtained with 10 s acquisition time. The spectral background was subtracted using an iterative-polynomial algorithm with a polynomial order of 2.\(^15\)

### Results

#### Macro- and microscopic surface chemistry characterization

The variation of the surface density of sulfonic acid groups was macroscopically characterized by static contact angle measurements (Fig. 2a). The contact angle measured on modified Au samples decreases approximately linearly with the increasing \(X_{\text{SO}_3\text{H}}\), indicating an increasing hydrophilicity of the surface. The monotonic trend suggests a strong correlation of solution \(X_{\text{SO}_3\text{H}}\) and the corresponding surface concentration. To confirm that our sample preparation protocol results in a homogeneous distribution of sulfonic acid groups (as opposed to microscale hydrophilic/hydrophobic phase separation and domain formation), Au-coated surfaces were characterized with SERS. Two example SERS spectra are shown in Fig. 2b for \(X_{\text{SO}_3\text{H}} = 0\) and \(X_{\text{SO}_3\text{H}} = 1\) that highlight spectral differences between the –CH\(_3\) and –SO\(_3\)H terminated thiols. The peak at 1060 cm\(^{-1}\) is assigned to the symmetric stretching vibration of sulfonic acid groups \(\nu(\text{S–O})\)^16 and the peak at 1095 cm\(^{-1}\) to the carbon chain skeletal vibration \(\nu(\text{CCCC})\).\(^17\) Thus, the relative amount of –CH\(_3\)/–SO\(_3\)H can be quantified as the band intensity ratio \(Q = I(\text{CCCC})/I(\text{S–O})\). The intensities of the two peaks (\(I(\text{CCCC})\) and \(I(\text{S–O})\)) were obtained by integrating the wavenumber ranges marked by grey shading in Fig. 2b.

| Table 1 Molar fraction \((X_{\text{SO}_3\text{H}})\) of HS(CH\(_2\))\(_3\)-SO\(_3\)H/HS(CH\(_2\))\(_3\)-CH\(_3\) in ethanol |
|---|---|---|---|---|---|---|---|
| X | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| X | 0 | 0.2 | 0.4 | 0.5 | 0.6 | 0.8 | 1 |
Symmetric functionalization by self-assembled monolayers on both the Au surface and the AFM tip (right) to mimic Nafion ionic channels (left). The aliphatic chains are attached to the gold via thiol bonds: \(-\text{CH}_3\), \(-\text{SO}_3\text{H}\) head groups. 

Images of \(Q\) for the surface prepared from the thiol–ethanol solution with \(X_{\text{SO}_3\text{H}} = 0, 0.5, \text{and 1}\) are shown in Fig. 2b (inset). Three different locations on these samples show that \(Q\) decreases with increasing \(X_{\text{SO}_3\text{H}}\) as expected, and none of the surfaces shows any microscale heterogeneity or phase separation within our optical spatial resolution of \(\sim 400\) nm. Fig. 2a (black) shows that the average \(Q\) values for each surface coating – computed by averaging \(Q\) from all pixels in three images per surface (\(N = 121\) spectra per image) – decreases almost linearly with increasing \(X_{\text{SO}_3\text{H}}\), consistent with the contact angle results. Characterization of the surfaces using SERS quantitatively shows that the surface density of sulfonic acid groups can be controlled by tuning the molar fraction \(X_{\text{SO}_3\text{H}}\) in the thiol solutions and that the surface distribution of the hydrophilic groups is homogeneous within our optical resolution.

**Nanoscale surface and solvent characteristics**

For each \(X_{\text{SO}_3\text{H}}\), we measured force–distance (F–D) characteristics between thiol-functionalized Au-coated AFM tips and Au surfaces with the same \(X_{\text{SO}_3\text{H}}\) on both the tip and the surface. As all recorded curves for a given molar fraction gave nearly the same response over the entire surface, we concluded that the prepared surfaces were homogeneously covered with hydrophilic and hydrophobic thiol molecules without nanoscale domain formation above the typical AFM contact area of \(<5–10\) nm\(^2\). From each group of 100 measured force curves taken at different locations, 15 approach curves were randomly selected and averaged into a mean curve. Averaging more than 15 curves did not improve the accuracy or information content. 100 curves were usually recorded to ensure that more curves than necessary were available for post-experimental analysis. Fig. 3 shows experimental average F–D characteristics and the fit curves from the extended Derjaguin–Landau–Verwey–Overbeek (DLVO) Hydra model. The experimental fluctuation between the 15 curves, presumably due to thermal noise, is by the grey dots.

The curves shown in Fig. 3 indicate a pronounced change of the F–D characteristics from an attractive force profile at \(X_{\text{SO}_3\text{H}} = 0\) (hydrophobic surface) to a fully repulsive profile at \(X_{\text{SO}_3\text{H}} = 1\) (hydrophilic surface), with a transition from attractive to repulsive forces at \(X_{\text{SO}_3\text{H}} \approx 0.4\). This degree of surface hydrophilicity seems to mark the threshold between the two regions where water is attracted to or depleted from the interface, which is confirmed by deduction of the Hydra parameters from the extended DLVO Hydra model fits as explained below.

The F–D characteristics indicate a strong attractive force at \(X_{\text{SO}_3\text{H}} = 0\) that cannot be explained solely by attractive van der Waals (vdW) forces; an additional well-known “hydrophobic attraction”\(^{11}\) acts due to a purported pronounced depletion of water at the hydrophobic interface.\(^{18,19}\) With increasing surface...
charge and hydrophilicity, water as well as ions are attracted toward the surface. As a result, the F–D characteristics switch from attractive to repulsive due to steric hydration repulsion originating from strongly surface-bound molecules, which can be described by an extended DLVO Hydra model (eqn (1)).\textsuperscript{20–23} This approach includes van der Waals (vdW) attraction, electrostatic double layer repulsion, and an additional hydration interaction parameter ($H_Y$):

$$ F_{\text{tot}} = F_{\text{vdW}}(\lambda_{\text{Hy}}) + F_{\text{EDL}}(\lambda_{\text{D}}, \sigma) + F_{\text{Hy}}(\gamma_{\text{Hy}}, H_Y, \lambda_{\text{Hy}}) \quad (1) $$

In eqn (1), the total measured force, which is commonly normalized by the effective radius $F_{\text{tot}}/R_{\text{eff}}$, is the sum of three independent force terms: (1) the attractive vdW force $F_{\text{vdW}}$, characterized by a distance-dependent Hamaker constant $A_{\text{Hy}}$, (2) the electrostatic double layer repulsion $F_{\text{EDL}}$, of the interacting surfaces with Debye length $\lambda_{\text{D}}$, and charge density $\sigma$, and (3) a term describing an additional contribution related to the hydrophobic properties of the surface: the Hydra force, $F_{\text{Hy}}$. $F_{\text{Hy}}$ is described by a recently suggested hydration parameter Hydra, $H_Y$, which corresponds essentially to the fraction of hydrophobic area at an interface and references the magnitude of the hydration interaction to a purely hydrophobic interaction ($H_Y = 1$) with a hydrophobic interface tension $\gamma_{\text{Hy}} = 45 \text{ mN m}^{-1}$.\textsuperscript{14} It becomes negative for repulsive hydration interactions. The Hydra decay length $\lambda_{\text{Hy}}$ characterizes the effective range of the hydration interaction and hence the hydration layering at the interacting surfaces. Below roughly $D < 2\lambda_{\text{D}}$ (i.e. 2 nm for the used electrolyte concentration), the electrostatic double layer force deviates from the Poisson–Boltzmann equation and the fitted hydration interaction $F_{\text{Hy}}$ overpowers the electrostatic contribution.

Detailed expressions for each force contribution are given in the ESI.\textsuperscript{†}

There are four fitting parameters (bolded in eqn (1)) in this equation: the tip radius $R_{\text{tip}}$, which varies between individual AFM tips typically in the range of 8 to 40 nm, Hydra decay length $\lambda_{\text{Hy}}$, and the charge density $\sigma$. All other parameters (see ESI†) can be fixed to known or measured values: the layer thickness $T_{\text{SAM}}$ (see XPS results in ESI†), Debye length $\lambda_{\text{D}}, \gamma_{\text{Hy}} = 45 \text{ mN m}^{-2}$, and the parameters that define the Hamaker constant for the studied systems. We assume constant charge boundary conditions for the electrostatic double layer force.

Fig. 3 shows the fitting of the data to the Hydra model for selected $X_{\text{SO}_3\text{H}}$ (red curves). All F–D curves can be fitted with the Hydra model except for the purely hydrophobic surface of $X_{\text{SO}_3\text{H}} = 0$ where the fitting deviates from the measured curves at $D \sim 1–2 \text{ nm}$ (marked in Fig. 3a), and the experimental curves show a very short-ranged strong repulsion that overpowers the hydrophobic attractions at $D \sim 5 \text{ Å}$. In our previous work with longer hydrocarbon chains, we recorded weak repulsive contributions within the attractive regime, albeit not as pronounced an effect as reported here.\textsuperscript{21} Similar short-range hydrophobic repulsion has been recently observed in experiments probing the hydrophobic force of diamond-like carbon.\textsuperscript{24}

Fig. 4 shows the extracted parameters charge density $\sigma$, hydration decay length $\lambda_{\text{Hy}}$, and the Hydra parameter $H_Y$, obtained from fitting the F–D curves to eqn (1) as a function of increasing hydrophilicity. The charge density magnitude (Fig. 4a) increases linearly with increasing $X_{\text{SO}_3\text{H}}$ to values of about $\sim 63 \text{ mC m}^{-2}$ for a fully hydrophilic surface, with an assumed $\sigma = 0$ for a completely hydrophobic (uncharged) surface (because $-\text{CH}_3$ groups are not charged). It is important to note that this fitted charge density is a measure for the charge at the plane of origin of the diffuse double layer. The plane of origin is typically $\sim 1 \text{ nm}$ from the actual surface plane, and the remaining (non-screened) charge is estimated to be 30% of that on the actual surface.\textsuperscript{25,26}

Fig. 4b shows the variation of the hydration decay length $\lambda_{\text{Hy}}$ as a function of $X_{\text{SO}_3\text{H}}$, $H_Y$ for hydrophobic surfaces is expected to be between 3 to 10 Å.\textsuperscript{27} Stock et al. showed that it is close to $7.2 \pm 1.2 \text{ Å}$.\textsuperscript{21} Our findings of $\lambda_{\text{Hy}}$ = 7 to 8 Å for the hydrophobic region of $X_{\text{SO}_3\text{H}} < 0.4$ compare well to the literature values. In the intermediate region, i.e. $0.4 < X_{\text{SO}_3\text{H}} < 0.6$, the $\lambda_{\text{Hy}}$ reaches a minimum of $\sim 2 \text{ Å}$, indicating considerably fewer strongly-surface-bound electrolyte constituents. At higher hydrophilic coverage, $\lambda_{\text{Hy}}$ reaches values of about 10 to 12 Å, indicating a surface-bound electrolyte that is interacting over longer distances as compared to hydrophobic surfaces, in good agreement with general observations of repulsive hydration forces.\textsuperscript{14}

The Hydra parameter $H_Y$ (Fig. 4c) switches from positive to negative between $X_{\text{SO}_3\text{H}} = 0.3$ and 0.4, with values of about 0.7 for hydrophobic surfaces of $X_{\text{SO}_3\text{H}} \leq 0.2$ to about $-0.5$ at $X_{\text{SO}_3\text{H}} = 0.4$. The crossing from positive to negative reveals that the tip–surface forces during approach switch from attraction to repulsion between $0.3 < X_{\text{SO}_3\text{H}} < 0.4$. This $H_Y$ minimum is
For very hydrophobic surfaces \(X_{\text{SO3H}} < 0.4\), significant attraction forces in the F–D curves manifest themselves as vDW forces and hydrophobic forces,\(^{29}\) indicating a likely depletion of the solution (water) layer\(^{18}\) between the two functionalized Au surfaces. The water depletion is believed to occur because of the inability of the few hydrophilic R-SO\(_3\)H head groups to promote complete solvation of the surface, thereby creating only isolated hydrated regions surrounded by regions with depleted water. For \(X_{\text{SO3H}} > 0.4\), the F–D profile was reversed showing purely repulsive forces upon tip approach, indicating that repulsive hydration forces and EDL contributions become dominant and overpower attractive vDW forces. The molar fraction of \(X_{\text{SO3H}} = 0.4\) marks the threshold between the two surface compositions where water is attracted to or depleted from the interface as shown by the sign change in the Hy parameter and the related switch from attractive to repulsive interactions.

Our observation of a \(\lambda_{\text{Hy}}\) of less than 2 Å for \(X_{\text{SO3H}} = 0.4\) to 0.6 indicates a complete breakdown of any long-range hydration forces at distances larger than 3–5 Å from the surface into the solution. We speculate that the extended long-range structure of the solution between the surface and the cantilever is limited. Instead, a compact short-ranged structural order on the surface may occur by effective lateral bridging of water molecules across neighboring sulfonic acid head groups. As a result, only very short-ranged hydration forces are measured. This picture is comparable to the one for solvation of small hydrophobes (such as e.g. methane) in bulk water by caging of water around them.\(^{29}\) Fig. 5 summarizes the interpretation from our measurements in the different regimes of electrolyte structuring.

Looking at the charge densities derived from the F–D fits, it is interesting to compare these values with values expected for a fully saturated –SO\(_3\)H surface. A fully covered SO\(_3\)H surface is expected to have a charge of \(\sim -740\) mC m\(^{-2}\) on the surface under fully deprotonated conditions (see ESI† for calculation of charge densities of fully covered and deprotonated SO\(_3\)H surfaces). This estimate is \(\sim 10\)-fold larger than the maximal charge density of \(-63\) mC m\(^{-2}\) obtained for the pure SO\(_3\)H-terminated surface at the plane of origin of the diffuse double layer (not the surface) in the work at hand. This discrepancy brings up two plausible scenarios: either (i) most of the charge on the model surface decays within the compact inner layer or (ii) the sulfonic acid groups do not fully deprotonate in pH 1 solution in our experiments. While it is not possible to precisely pinpoint the magnitude of either effect, we can provide a reasonable estimate for the screening effect. Force spectroscopy results suggest that typically \(\sim 80 \pm 10\%\) of a surface charge is screened within the compact inner layer.\(^{25}\) Assuming an extreme 90% charge screening between the surface and the inner diffuse layer, the fitted diffuse layer charge density at \(X_{\text{SO3H}} = 1\) would suggest surface charge densities \((-63\) mC m\(^{-2}\) \(\times (1/0.1) = -630\) mC m\(^{-2}\)) which are close to the expected charge density value of a fully covered SO\(_3\)H surface (\(\sim 740\) mC m\(^{-2}\)). This numerical agreement demonstrates that charge screening between the surface and inner diffuse layer alone can potentially explain the lower charge densities extracted from fitting to our measurements. Moreover, with the SO\(_3\)H groups having a \(pK_a\) \(~ -2\) and our electrolyte solution having a pH = 1, it is reasonable to assume

**Discussion**

From the experimental results and those of the Hydra model fitting, we find clear trends in both the force profiles and model parameters as the functionalization of the surface progresses toward more hydrophilic coverage. The R-SO\(_3\)H molar fraction of \(X_{\text{SO3H}} = 0.4\) seems to be the critical value where the hydration properties change significantly as indicated by the Hy parameter. For very hydrophobic surfaces \(X_{\text{SO3H}} < 0.4\), significant attraction forces in the F–D curves manifest themselves as vDW forces and hydrophobic forces,\(^{29}\) indicating a likely depletion of

**Fig. 4** Evolution of fitting parameters for Hydra model as function of \(X_{\text{SO3H}}\): (a) Charge density \(\sigma\), (b) hydration decay length \(\lambda_{\text{Hy}}\), (c) hydration parameter Hy.
that most SO$_3$H groups are deprotonated, even considering electrostatic repulsion effects (which have been shown to increase the solution pK$_a$ by 1–2 units in our experiments). Therefore, we surmise that reduced deprotonation likely plays a secondary effect.

For comparison of our model surfaces to Nafion, it is instructive to estimate the surface charge in prototypical Nafion ionic channels with an equivalent weight of 1100 g mol$^{-1}$ SO$_3$H, and an ionic channel diameter of ~5 nm.$^{31,32}$ Allowing for complete SO$_3$H deprotonation in a fully hydrated Nafion membrane at pH 7 water (more than 9 pH units above the solution pK$_a$ of SO$_3$H acid) and one charge for each SO$_3^-$ group, we calculate an approximate surface charge density of ~$370$ mC m$^{-2}$ in Nafion (see ESI† for Calculation). Again assuming a 90% charge screening between the surface and the inner diffuse layer (where we actually probe), we find that $X_{SO_3H} \sim 0.5$ provides a charge density comparable to Nafion ionic channels. In the context of our results, it would thus appear that Nafion membranes should exhibit similar values for $\lambda_{Hy}$ of $-0.45$ and for $\lambda_{Hy}$ of less than 2 Å as those found at $X_{SO_3H} \sim 0.5$ in our model system. Such a short hydration length implies local water structuring close to the channel surface, while the remaining solution toward the center of the ionic channel is essentially bulk-like and unstructured. Interestingly, we and others have proposed that two types of water exist in Nafion membranes in a core–shell structure$^{9,33}$ and that the shell water thickness near the sulfonic acid/Teflon backbone is estimated to be ~2 Å, which is quantitatively consistent with $\lambda_{Hy}$ at $X_{SO_3H} \sim 0.5$. Thus, we surmise that the moderately charged SAM surface of $X_{SO_3H} \sim 0.5$ appears to induce solution structuring very locally at the surface, comparable to the shell water in Nafion, that becomes bulk-like beyond this distance (i.e. core water in Nafion).

**Conclusions**

We have established a model system for PFSA membrane ionic channels using self-assembled monolayers with a tunable surface density of sulfonic acid and methyl groups to tune surface hydrophilicity. We quantified the hydration forces and deduced the interfacial electrolyte structure from AFM force–distance measurements. The measured force profiles indicate a pronounced change of the electrolyte layering density at the surface with an unexpectedly sharp hydrophobic-to-hydrophilic transition for $X_{SO_3H} \sim 0.4$. This inflection point can be attributed to a change from hydrophobic water depletion to strong water and ion adsorption at the surface for hydrophobic and hydrophilic surfaces, respectively, due to buildup of sufficient surface charge. Using an extended Derjaguin–Landau–Verwey–Overbeek model including the Hydra force, we quantify diffuse double layer charges and characteristic hydration lengths as a function of sulfonic acid group density on the surface. Interestingly, from $X_{SO_3H} \sim 0.4$–0.6 we find a significant breakdown of the long-ranged hydration effects beyond a few Å in force-versus-distance characteristics. The extracted surface charge density for surfaces with these molar fractions similar to charge densities estimated for fully hydrated Nafion, assuming a large decay of surface charge due to screening between the surface and inner diffuse layer. This suggests that the observed very short hydration layering and strong long-ranged repulsive hydration forces may also occur within Nafion ionic channels in support of the core–shell hypothesis of water in Nafion. The effect of electrolyte structuring on transport is yet unknown and will be the subject of follow-up studies.

**Conflicts of interest**

There are no conflicts to declare.

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