Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2021, 12, 5865

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 19th January 2021 Accepted 16th March 2021

DOI: 10.1039/d1sc00354b

rsc.li/chemical-science

Effects of applied voltage on water at a gold electrode interface from *ab initio* molecular dynamics†

Zachary K. Goldsmith, D* Marcos F. Calegari Andrade D and Annabella Selloni **D**

Electrode—water interfaces under voltage bias demonstrate anomalous electrostatic and structural properties that are influential in their catalytic and technological applications. Mean-field and empirical models of the electrical double layer (EDL) that forms in response to an applied potential do not capture the heterogeneity that polarizable, liquid-phase water molecules engender. To illustrate the inhomogeneous nature of the electrochemical interface, Born—Oppenheimer *ab initio* molecular dynamics calculations of electrified Au(111) slabs interfaced with liquid water were performed using a combined explicit—implicit solvent approach. The excess charges localized on the model electrode were held constant and the electrode potentials were computed at frequent simulation times. The electrode potential in each trajectory fluctuated with changes in the atomic structure, and the trajectory-averaged potentials converged and yielded a physically reasonable differential capacitance for the system. The effects of the average applied voltages, both positive and negative, on the structural, hydrogen bonding, dynamical, and vibrational properties of water were characterized and compared to literature where applicable. Controlled-potential simulations of the interfacial solvent dynamics provide a framework for further investigation of more complex or reactive species in the EDL and broadly for understanding electrochemical interfaces *in situ*.

Introduction

Aqueous electrochemical interfaces are central to heterogeneous catalysis^{1,2} and emerging energy conversion technologies.^{3–5} Electrical double layers (EDL) of solvated ions notably form at such interfaces, however, water molecules will predominate the immediate interface even for concentrated electrolyte solutions⁶ and are known to exhibit starkly different properties and reactivity than water in the bulk liquid.⁷ It is therefore imperative to both characterize water at electrochemical interfaces and develop the computational tools for doing so as a step towards modeling more sophisticated EDL environments.⁸

Experimental studies of the structure of the electrode-water interface, often by X-ray absorption spectroscopy (XAS), have underscored its anomalous structure and hydrogen bonding patterns. 9,10 Specifically, water forms at least one structural layer against common electrodes such as Ag and Au with densities higher than bulk and with dangling OH bonds at both positive and negative voltage biases. 11 Directly studying electrochemical interfaces *in situ* has been largely accomplished *via* surface-

Department of Chemistry, Princeton University, Princeton, NJ 08544, USA. E-mail: zkg@princeton.edu; aselloni@princeton.edu

† Electronic supplementary information (ESI) available. See DOI: 10.1039/d1sc00354b

specific vibrational spectroscopies such as surface-enhanced infrared absorption (SEIRAS),^{12,13} plasmon-enhanced Raman spectroscopy,¹⁴ and sum-frequency generation (SFG).^{15,16} These techniques have elucidated dependencies of the vibrational spectra of water at the interface on the voltage applied, consequences of the strong electric fields in the EDL.¹⁷ Nonetheless, experimental probes of the structure and composition of the EDL are challenging to perform, incomplete in scope, and would benefit from improvements in atomistic modeling.

Realistic simulation of the EDL and processes therein is an outstanding challenge for theory and computation as well. Classical MD and Monte Carlo simulations of water at electrified surfaces¹⁸⁻²⁰ have been instructive but are limited in their microscopic resolution. Notably, empirical approaches lack instantaneous polarization of species in the interfacial electric field that may give way to novel properties and/or reactivity. Quantum chemical simulations, most practically density functional theory (DFT), are capable of describing molecular polarization and bond-breaking, albeit at greater computational cost. DFT with *a posteriori* corrections for the thermodynamic effects of the electrode potential^{21,22} have been useful in elaborating catalytic mechanisms, but neglect specific effects of the strong, induced electric fields at the interface.

Recent advances in periodic DFT with continuum solvation, ²³⁻²⁶ self-consistent analytical EDL models, ^{27,28} as well as formally grand-canonical approaches^{29,30} have enabled studies of more sophisticated systems that include a controlled potential. Applications of such implicit and/or hybrid solvation methods have included potentiostatic simulations along key nuclear coordinates associated with heterogeneous catalysis31-34 and surface-enhanced spectroscopy.35,36 Dynamical ab initio studies including voltage bias have mostly considered just the electrode37 or limited numbers of molecules38-40 and/or monolayers.14,41-44

In this work, we performed ab initio molecular dynamics of water at a Au electrode interface with fixed amounts of charge on the Au electrode as a means of modulating the electrode potential. The electrode potentials were measured at frequent times for each simulation and their converged average values yielded a physically reasonable differential capacitance for the system. The recovery of the system's electronic properties enabled analyses of the structure and dynamics of interfacial water under quantifiable potentials. In particular, we elucidated the effect of voltage on the density of water with respect to surface normal as well as the characteristic orientations of water molecules near the electrode as a function of the voltage bias. Furthermore, our computed vibrational densities of states of the interfacial water under bias corroborated phenomena previously reported by surface-enhanced spectroscopies. Moreover, we realized ab initio-quality simulations of an electrode under quantifiable voltage biases in contact with several layers of explicit solvent, a long outstanding benchmark for computation. We anticipate that atomistic modeling of electrochemical interfaces will be a powerful tool for understanding how chemical mechanisms are shaped by the polarization and statistical dynamics of the EDL.

2. Methods

Computational system

We devised a model system that was small enough to enable several AIMD trajectories while maintaining the essential physical chemistry of both phases at the interface. In addition, Au(111)/water was chosen so that no faradaic charge transfer would occur on the timescale of the simulations, maintaining our focus on the interfacial system near equilibrium. The lack of faradaic charge transfer furthermore allowed for the constant charge ensemble employed to best approximate that of a constant potential,45 which is the more practical experimental condition. The 3D-periodic computational unit cell was therefore comprised of a 4 × 4 × 3 Au(111) slab (48 Au atoms) interfaced with 56 molecules of water. The surface-parallel cell dimensions as determined by the Au(111) slab's size were approximately 10.2 Å \times 11.8 Å and the total height of the orthorhombic cell was 36 Å. The computational cell is depicted in Fig. 1.

The inactive back side of the Au(111) slab was in contact with vacuum ($\varepsilon = 1$) and the surface atoms on this side were held fixed in the direction perpendicular to the surface (z). A region of dielectric continuum representative of bulk water ($\varepsilon = 78.3$) was imposed beyond the region of explicit water molecules. This hybrid explicit/implicit approach enabled us to use a smaller system in expensive electronic structure calculations

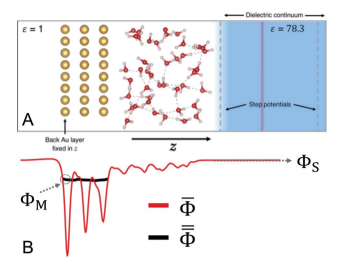


Fig. 1 (A) Schematic illustration of the Au(111)/water computational unit cell. Au atoms are gold in color, O atoms are red, and H atoms are white. The light blue color indicates the region of dielectric continuum water ($\varepsilon = 78.3$). The inactive side of the Au slab is in contact with vacuum ($\varepsilon = 1$). The transparent red plane represents the neutralizing counter-charge and the black dashed vertical lines are indicative of the step potentials used to separate the explicit and implicit water regions. (B) Representative xy plane-averaged electrostatic potential $\bar{\Phi}$ (red) and its corresponding macroscopically-averaged potential within the Au slab $ar{\Phi}$ as functions of z for the system in (A). The values Φ_{M} and Φ_{S} are illustrated as the value of $ar{\Phi}$ in the back of the Au slab and the value of $ar{\Phi}$ deep in the continuum region, respectively.

than a fully explicit system. More critically, this hybrid approach made it possible to establish both a region of "bulk solution" and a vacuum on the inactive side of the Au slab. The presence of the vacuum ensured the selective localization of excess charge at the interface with water. 33 The continuum solvent was included using the Environ^{24,46} plug-in to Quantum-ESPRESSO, 47,48 which solves the Poisson equation for the dielectric continuum in contact with the atomistic system selfconsistently with the Kohn-Sham equation. Specifically, we employed a "solvent-aware" implementation of the selfconsistent continuum solvation method24 and solved the relevant Poisson equation using periodic boundary conditions only in the surface-parallel dimensions to avoid solvation interactions between unit cells. A plane of charge equal and opposite that on the electrode was positioned within the region of implicit water to maintain the overall neutrality of the periodic cell. In addition, a dipole correction was employed in the vacant region near the boundary of the cell50 to negate spurious electrostatic interactions via periodicity.

Additionally, a planar step function potential⁵¹ was positioned at a distance from the Au surface that ensured a total density of 1 g mL⁻¹ of the 56 water molecules. The step function, which in practice was error function-shaped, had a height of 0.25 Ry. To maintain 3D periodicity, this region was terminated near the top of the unit cell and beyond the position of the counter-charge plane by a step function of equal size and opposite directionality. These features are illustrated in Fig. 1.

2.2 Ab initio molecular dynamics calculations

Edge Article

We performed DFT-based Born-Oppenheimer molecular dynamics (BOMD) trajectories using the PWscf code of the Quantum-ESPRESSO package.47,48 We employed optimized norm-conserving Vanderbilt pseudopotentials⁵² and a plane wave cutoff energy of 60 Ry,53 which is greater than the suggested cutoff for these pseudopotentials. Only the gamma point was used to sample the surface Brillouin zone. We used the PBE functional54 for all DFT calculations with the exception of the BOMD trajectory in which there was no excess charge on the electrode (i.e. at the potential of zero charge (PZC)). For this we used PBE with a Tkatchenko-Scheffler⁵⁵ correction for van der Waals (vdW) interactions (PBE + TS), which are known to be especially important for unbiased metal-water interfaces. 56,57 Contributions from vdW interactions were not expected to be as important for describing water at electrode surfaces carrying excess charge, at which configurations with water oriented parallel to the surface were expected to be disfavored. 40 Moreover, we demonstrate herein that the qualitative results of this study were not impacted by the inclusion of vdW correction only at PZC.

We carried out five BOMD trajectories of our model electrochemical half-cell, each with a different amount of charge localized at the Au surface. The total electrode charges per unit cell σ used were -0.4e, -0.2e, 0e, +0.2e, and +0.4e, where e is the elementary charge. These amounted to at most $\pm 0.025e$ per surface Au atom, if distributed evenly. We demonstrate below that these σ represented modest applied potentials smaller in magnitude than ± 0.5 V vs. PZC. The BOMD runs were performed with a timestep of ~0.48 fs, which was made possible by using the mass of D for all H atoms, and at 330 K, as is typical for AIMD simulations of bulk liquid water.58 The results below regarding hydrogen bonding and diffusivity are indicative that water was sufficiently liquid in our simulations. The physical Au and O masses were used otherwise. We employed an NVT ensemble with a velocity-rescaling thermostat with a tolerance of ± 100 K. Each trajectory spanned over 10 ps of integration time, which we will later show was sufficient to recover a physical differential capacitance for the system. The initial coordinates were generated via a geometry minimization in a classical force field57 with no excess charges followed by partial DFT-level geometry optimization at each trajectory's respective electrode charge. The 3D electrostatic potential Φ , as obtained by solving the Kohn-Sham equation self-consistently, was recorded every 100 time steps (\sim 48 fs) of each trajectory.

2.3 Calculation of electrode potential

Here we briefly introduce the absolute electrode potential, its constituent quantities, and demonstrate the simple utility of measuring the computed electrode potentials relative to the system's potential of zero charge (PZC). The absolute electrode potential $\Phi^{\rm abs}$ can be expressed as, ^{59,60}

$$\Phi^{\rm abs} = \Phi_{\rm e} - \frac{\mu_{\rm M}}{F} \tag{1}$$

where $\Phi_{\rm e}$ is the difference between the electrostatic potentials in the metal $\Phi_{
m M}$ and in bulk solution $\Phi_{
m S}$, also known as the contact or Volta potential difference. 59,60 In eqn (1), $\mu_{\rm M}$ is the Fermi energy of the bulk metal (the negative of its work function), and F is Faraday's constant. The quantities $\Phi_{\rm S}$ and $\Phi_{\rm M}$ were computed at each MD step at which Φ was recorded. This was achieved by averaging the computed electrostatic potential Φ in the xy plane to obtain $\bar{\Phi}$. Then, we took as Φ_S the value of $\bar{\Phi}$ deep in the continuum solvent region of the cell, 25,36,61 where $\bar{\Phi}$ is unchanging as a function of z (Fig. 1B). To measure the metal potential, we macroscopically-averaged $\bar{\Phi}$ in the Au slab using the optimized layer spacing of 2.4 Å and took as μ_{M} the average of this quantity over the back 1 Å of the slab. Utilizing the partially-frozen back layer provided a more stable metal potential (Fig. 1B), although alternative definitions for Φ_{M} did not qualitatively change our results (see Fig. S3†). The resulting values of $\Phi_{\rm e}$ presented below represent the difference between the quantities $\Phi_{\rm M}$ and $\Phi_{\rm S}$ for each trajectory.

For the purposes of analyzing the effects of electrode potential on various properties of water we referred potentials to the system's PZC, *i.e.* $\sigma=0$ e. The referenced potentials $\Delta \Phi_{\rm e}$ are expressed as,

$$\Delta \Phi_{\rm e} = \Phi^{\rm abs}[\sigma] - \Phi^{\rm abs}[\sigma = 0] = \Phi_{\rm e}[\sigma] - \Phi_{\rm e}[\sigma = 0] \tag{2}$$

In this expression it is evident that we do not need the bulk metal's Fermi energy to compute the referenced electrode potentials. While certain implementations of *e.g.* Poisson–Boltzmann boundary conditions may make it possible to obtain the absolute potentials from *ab initio* calculations of small systems, 27,41,62 the PZC-referenced potentials will suffice to quantify the effects of electrode polarization on water. In the present application, the referenced electrode potentials associated with each σ will emerge as the trajectory averages of the $\Phi_{\rm e}$, referred to hereafter as φ and computed as,

$$\varphi = \Delta \langle \Phi_{\rm e} \rangle \tag{3}$$

In the following section we will first present the dynamics and convergence of each trajectory's $\Phi_{\rm e}$ before subsequently analyzing the structure, dynamics, and vibrational spectra of interfacial water as a function of φ , the trajectory-averaged potentials *versus* PZC.

Results and discussion

3.1 Electrical properties of Au(111)/water

3.1.1 Time evolution of electrode potentials. The calculated plane-averaged electrostatic potentials $\bar{\Phi}$ for each electrode charge as well as the full trajectory averages are shown in Fig. 2. The electrostatic potential wells observed include one for each of the layers of the Au slab and at least one characteristic of a structural layer of water at the interface. In the region of dielectric continuum, to the right of the potential energy step in Fig. 2, $\bar{\Phi}$ is unchanging in z, making the calculation of $\Phi_{\rm S}$ straightforward. The metal potentials as measured at the back of the slab were relatively stable as a function of time, as shown

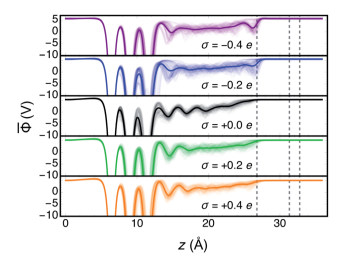


Fig. 2 Plane-averaged electrostatic potentials $\bar{\Phi}$ measured every $\sim\!48$ fs (transparent) and the trajectory-averaged $\langle\bar{\Phi}\rangle$ (opaque) for all five electrode charges. The deeper potential wells associated with the three Au slab layers have been cut off for clarity in the water region. The dashed vertical lines in each represent the locations of (from left to right) the potential step up, the counter charge plane, and the potential step down.

in Fig. S2 and S3.† In the two trajectories with negative electrode charges, and therefore positive counter charges, we observed electrostatic potential wells near the boundary of the region of explicit water. While these are indicative of a spurious interaction between some water molecules and the counter charge plane, the below analyses demonstrate that the system's electrical properties and more immediate interfacial water layers were nonetheless physically realistic.

At each electrode charge, $\bar{\Phi}$ fluctuated as a consequence of the changing atomic structure during the molecular dynamics, as depicted in Fig. 2. Moreover, for these fixed σ trajectories, motion of both the Au and water atoms yielded changes to the electrode potentials $\Phi_{\rm e}$ over time, plotted in Fig. 3A. The values of $\Phi_{\rm e}$ for the five trajectories largely preserve the intuitive ordering according to the electrode charges throughout the

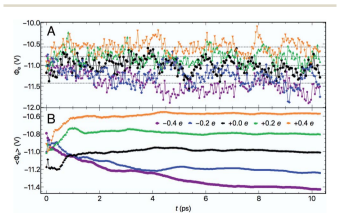


Fig. 3 (A) Electrode potentials $\Phi_{\rm e}$ versus trajectory time for five electrode charge and (B) the cumulative time-averaged electrode potentials $\langle \Phi_{\rm e} \rangle$ as a function of time.

dynamics. After at most 5–6 ps of dynamics, the cumulative time averages $\langle \Phi_e \rangle$ (shown in Fig. 3B) were well converged. Although the 10 ps trajectories were short relative to molecular dynamics studies of interfacial water, ⁶³ the converged electrical properties of the system allowed for a qualitative investigation into water's structure and dynamics under quantifiable voltage bias.

3.1.2 Referenced potentials and differential capacitance. The averaged potentials *versus* PZC φ in the five trajectories were (from most to least negative) -0.42 V, -0.24 V, 0 V, 0.21 V, and 0.44 V and are presented as such in Fig. 4. The error bars shown in Fig. 4 reflect those obtained via a block-averaging of each trajectory and are all smaller than \sim 0.07 V. On the contrary, the full-trajectory standard deviations of φ for each trajectory were \sim 0.2 V. In other words, the BOMD trajectories with fixed electrode charges σ exhibited large fluctuations of the potentials around a convergent average (Fig. 3). The large fluctuations of the potential in each trajectory can be attributed predominantly to the fluctuations in $\Phi_{\rm M}$, as $\Phi_{\rm S}$ were comparatively more stable (Fig. S1†). Conversely, fixed-potential ab initio approaches³⁰ have engendered large fluctuations of the electrode charge. We emphasize that the fixed-charge approach was appropriate because the Au(111)/water system was electrochemically inert on the timescale of the simulations. In general, a fixed-potential approach is more comparable to experiment.

The relationship between the surface charges σ and the full trajectory-averaged electrode potentials versus PZC φ was found to be nearly perfectly linear, as depicted in Fig. 4. We ascribe the observed symmetry of the potentials around PZC to the lack of electrolyte ions in our system. The inverse slope of the linear fit to these five data points represents the differential capacitance $C_{\rm d}$ of the Au(111)/water system, $14.3 \pm 2.2~\mu {\rm F~cm}^{-2}$. This $C_{\rm d}$ is within the rather large range of experimentally measured values, 11–40 $\mu {\rm F~cm}^{-2}$, 64 -66 which depend significantly on experimental conditions including but not limited to the concentration and identity of the electrolyte ions. In the absence of ions, we are satisfied to recover a reasonable

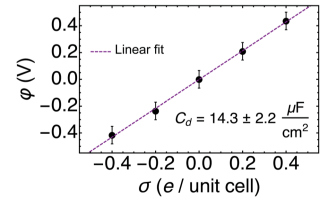


Fig. 4 Full trajectory-averaged, referenced electrode potentials φ as a function of the electrode surface charge per unit cell σ . The dashed purple line represents a linear regression to the five data points whose inverse slope corresponds to a differential capacitance, $C_{\rm d}$, of 14.3 \pm 2.2 $\mu {\rm F}$ cm⁻².

Edge Article Chemical Science

differential capacitance for Au(111)/water towards the bottom of the measured range of $C_{\rm d}$. Similar values of $C_{\rm d}$ have been achieved for Au(111)/water with a fully implicit solvent with specifically tuned parameters²⁵ and with a monolayer of water. Our results recover this macroscopic electrical property from short AIMD trajectories and an effectively fully explicit solvent.

3.2 Structural and dynamic response of water to applied voltages

3.2.1 Water density versus surface-perpendicular z. The density profile of water (Fig. 5) versus z was calculated vis-à-vis the O atom number density to determine the effects of the metal interface and the applied potential φ on water's structure. At each φ an interfacial layer of water formed, centered 3–3.5 Å from the surface with a density maximum of at least 1.5 and up to about 3 times that of bulk water. At PZC ($\varphi = 0$ V) and at positive φ water demonstrated larger initial density peaks than at negative φ . The first density peaks at negative φ were found to be further from the Au(111) surface and had smaller maximum densities of between 1.5 and 2 times that of bulk. It is intuitive that the more concentrated negative charges on O are selectively repelled from the surface when it is charged negatively and conversely attracted when the surface is charged positively.9 The non-negative φ systems exhibited second water layers with densities greater than bulk, and a third such layer can even be observed for the system at PZC. The negative φ systems only demonstrated one such structural layer each. In the negative φ systems, we observed a structural layer of water near the imposed boundary of the region, likely due to interactions with the neutralizing counter charge. Given that the overall electrical properties were recovered despite this spurious behavior, the structural and dynamical properties of the more immediately interfacial water were expected to be physical regardless.

The structural similarities of the first water layers at PZC and at $\varphi > 0$ V have been previously observed by XAS.¹¹ Furthermore, the position and maximum of the first density peak at PZC is in agreement with recent AIMD studies of Au(111)/water.^{68,69} An earlier AIMD study of this system at PZC predicted a narrowly resolved first water layer with a density maximum of \sim 4 times that of bulk water,¹¹ although this work was performed with PBE

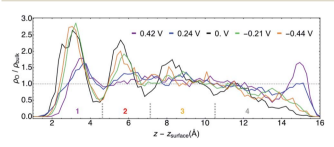


Fig. 5 Calculated O atom number densities $\rho_{\rm O}$ relative that for experimental bulk water $\rho_{\rm bulk}$ versus z corrected for the position of the Au(111) surface $z_{\rm surface}$ for all potentials φ . The horizontal dashed line represents the experimental bulk water density. The three short, dashed, gray vertical lines approximate the boundaries of the water layers 1–4 for which subsequent analyses are performed.

water at 300 K, which is now known to likely be over-structured.⁷⁰

For the purposes of further analyses, we designated four characteristic layers of water for each trajectory, the approximate positions of which are delineated in Fig. 5. The layer boundaries were chosen at local density minima and differed slightly for each φ . Hereafter we will refer to water in these spatial regions by the numbers shown in Fig. 5, with layer 1 being the most immediate interfacial layer, followed by layer 2. Layer 3 is yet further from the surface and appears in terms of the density to be most similar to bulk water. We will neglect layer 4 in the subsequent analyses given its aforementioned unphysical behavior.

3.2.2 Orientations of interfacial water under voltage bias. Beyond the formation of structural layers, we investigated how φ influenced the orientation of water's geometric dipoles. Fig. 6A is the histogram of the cosine of the angles formed between the water bisectors and a unit vector parallel to the z axis, only for layer 1 at each φ . The maximum $P(\cos \theta)$ at nonzero φ in Fig. 6A demonstrate that when the Au surface was charged positively the bisectors were selectively oriented away from the surface (O atoms oriented towards the surface) and when the surface was negatively charged the water dipoles were selectively oriented towards the surface (O atoms oriented away from the surface). These results follow intuitively from

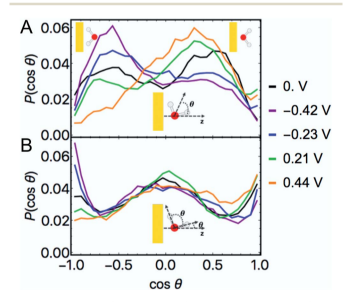


Fig. 6 Histograms of the cosines of the respective angles θ formed between a unit vector in the direction of the z axis and the bisectors of water (above) and the unit vectors corresponding to all OD bonds (below), in which $P(\cos\theta)$ is the normalized frequency. Inset illustrations indicate the relevant angles θ for each plot in which the yellow rectangle represents the Au slab, the O atom is red, and the D atoms are white. These data correspond only to water in layer 1, the water molecules nearest the surface. In each case, a $\cos\theta=1$ refers to the associated vector pointing away from the Au surface whereas $\cos\theta=-1$ indicates that vector being oriented towards the surface. These extrema as they pertain to the water bisectors are depicted by the cartoons in the top corners of the upper panel. $\cos\theta=0$ indicates that the bisector/OD bond is parallel to the surface.

Open Access Article. Published on 18 March 2021. Downloaded on 8/2/2025 9:18:22 AM

Chemical Science

electrostatic attraction of O to the positively charged surface

electrostatic attraction of O to the positively charged surface and of H/D to the negatively charged surface. Nonetheless, these maxima appear not at $\cos\theta=\pm 1$ but at intermediate positive and negative values, *i.e.* at these modest voltages water's geometric dipoles are not entirely oriented to the electric fields. We also observed non-negligible probabilities of water molecules being oriented counter to the prevailing electrostatics for both positive and negative φ . At PZC, we observed a bimodal distribution of water bisectors with more being oriented generally away from the surface by a small margin.⁷¹ This is in subtle contrast to literature characterizations of room temperature water at the Au(111) interface as being similar to a 2D ice, in which the bisectors would be predominantly parallel to the surface and $P(\cos\theta)$ would be maximal near $\cos\theta=0.^{11}$ In general, the bisector orientation statistics in layer 1 appear to smoothly change as a function of φ .

The histogram of the cosine of the angles formed between all water OD vectors and a unit vector parallel to the z axis in layer 1 is given in Fig. 6B. These plots demonstrate local maxima at -1 for the negative φ and at +1 for the positive φ , *i.e.* OD bonds pointed perpendicularly towards and away from the surface, respectively, in accordance with the bond's geometric dipole. At each non-zero φ we additionally observe significant frequencies near $\cos\theta=0$, indicative of a significant proportion of OD bonds being nearly parallel to the surface in layer 1.

The complimentary observations of water bisectors not being entirely oriented along the applied fields and the significant proportion of OD bonds being roughly parallel to the surface suggest a general characterization of water's structure in layer 1: water's geometric dipole being partially oriented according to the local field while sustaining hydrogen bonding interactions with neighboring water molecules. Nonetheless, interfacial water at room temperature still exhibits considerable disorder within this paradigm at the modest values of φ studied.

This characterization is further supported by an analysis of the hydrogen bonding properties of water in our simulations that found on average at least 1 and up to more than 2 intralayer hydrogen bonds among layer 1 water at every potential (Fig. S5†). In addition, our calculations suggest fewer hydrogen bonds formed by water in layer 1 at negative φ than at PZC and positive φ , on average (Fig. S5†). This is consistent with the observation of more dangling OH bonds at negative potentials than at positive potentials by XAS.11 The hydrogen bonding properties of water at an Au(111) interface at PZC are in accordance with a previous BOMD study.71 The average number of hydrogen bonds in layer 3, the most bulk-like water, ranged from 3.42-3.58 (Fig. S7†). This compares favorably to the experimental value for bulk water at 330 K, 3.58.72 Comparable BOMD simulations of bulk water estimate this value to be approximately 3.85, 3.77, and 3.60 for PBE, PBE + TS, and SCAN, respectively.58

The biasing voltage affects the orientation of water in sub-interfacial layers 2 and 3 as well, as shown in Fig. S8 and S9.† We observed little attenuation of the bisector orientation preferences of water in layers 2 and 3 at non-zero φ . In the case of the PZC trajectory, water no longer shows any significant orientation pattern beyond layer 1.

3.2.3 Diffusion coefficients of water. We furthermore utilized the controlled-potential AIMD trajectories to understand the effects of applied potential and distance from the surface on the diffusivity of water parallel to the surface. We computed the parallel diffusion coefficients D_{\parallel} , shown in Fig. 7, via Einstein's relation and the mean-square displacements (MSDs) of the O atoms (see Fig. S10–S12†). The error bars presented in Fig. 7, obtained via block-averaging the MSD for each trajectory, are relatively large as a consequence of the short trajectory lengths.

For water in layer 1, we observed an increase in the D_{\parallel} at negative φ relative to both PZC and to bulk D_2O . At PZC and at positive φ we observed little difference within error of D_{\parallel} relative to bulk D_2O , further evidence contrary to layer 1 water having ice-like properties. For layer 2, we found a minimum D_{\parallel} at PZC, with D_{\parallel} at PZC being below, but within the error, of bulk D_2O . Furthermore, D_{\parallel} at both positive and negative potentials in layer 2 was found to be higher than that of bulk D_2O . In layer 3 we observed a similar response of D_{\parallel} to φ as in layer 2. Note that in layer 3, the most bulk-like water in our small simulation, D_{\parallel} at PZC was nearly identical to that of bulk D_2O (Fig. 7).

The observation of increased parallel diffusivity of water in layer 1 at negative φ can be rationalized by the smaller average number of hydrogen bonds observed under these conditions (Fig. S5-S7†). Conversely, the total average numbers of hydrogen bonds were roughly the same for the trajectories of φ ≥ 0, for which the layer 1 parallel diffusion coefficients were similar. The increased D_{\parallel} in layers 2 and 3 at $\varphi \neq 0$ are more difficult to rationalize on the basis of hydrogen bonding, given that their total numbers and ratios of intra- and inter-layer hydrogen bonding are not very different from PZC in these layers (Fig. S5-S7†). However, there is literature precedent for water's diffusion being faster in small, oscillatory external electric fields,73 which may describe the sub-interfacial water layer. Recent AIMD simulations demonstrated that such electric fields induced frequent rearrangements of the hydrogen bonding network and consequentially increased water's diffusivity. 3 We also note that D_{\parallel} in each layer is generally lowest at

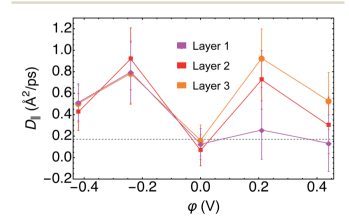


Fig. 7 Diffusion coefficients parallel to the surface D_{\parallel} as a function of applied potential φ for each of the first three layers of D₂O. The gray dashed line corresponds to the 2D diffusion coefficient for bulk D₂O at room temperature.

PZC, the only trajectory in which the vdW correction was included, despite the inclusion of vdW corrections in AIMD being known to increase the diffusivity of bulk water.⁵⁸

3.3 Water vibrational densities of states *versus* electrode potential

We computed the vibrational density of states (VDOS) for water in each layer and at each potential from the BOMD trajectories. The VDOS spectra calculated by Fourier transforming the velocity autocorrelation functions for each trajectory are shown in Fig. 8. For the purposes of qualitatively assessing the effects of distance from the electrode and φ on the VDOS spectra, we will focus on the peaks associated with the bending (δDOD , \sim 1200 cm⁻¹) and stretching (ν OD, \sim 2400–2600 cm⁻¹) modes of D₂O. The former yielded a single, relatively narrow peak, whereas the latter exhibited a much broader peak that in some cases was evidently bimodal. Bimodal vOH(D) signals at the Au(111) interface have been previously observed by SEIRAS12 and SFG15 and from BOMD of similar systems at PZC.74 We emphasize that VDOS spectra are related to but not exactly coincident with the spectra associated with any surfaceenhanced spectroscopy.

In the VDOS spectra for layer 1 water (top panel of Fig. 8) there is a clear dependence on φ . Both positive and negative φ have the effect of blue shifting the peaks associated with the δ DOD and ν OD vibrations. This phenomenon has been observed experimentally by SEIRAS. Moreover, the blue shifts of the two characteristic signatures with both positive and negative φ are consistent with the orientation of water's geometric dipoles with the induced fields ν is- \dot{a} - ν is the Stark effect. This qualitative conclusion holds despite the vdW correction (only employed at PZC) being known to blue shift both the δ DOD and ν OD peaks. Respectively.

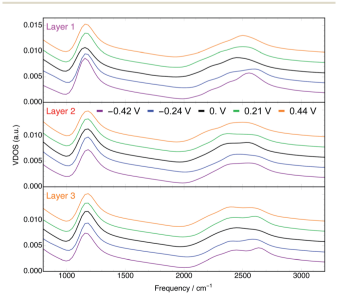


Fig. 8 Vibrational density of states of D_2O in layers 1 (top), 2 (middle), and 3 (bottom) at every φ . Spectra for each φ are vertically offset by arbitrary amounts in each plot for visual clarity.

Furthermore, the δ DOD peaks of layer 1 water appear at lower frequencies relative to those of water in layers 2 and 3 (Fig. S19†). This is suggestive of a weaker hydrogen bonding network⁷⁸ in layer 1, consistent with our observation of fewer overall hydrogen bonds formed in the interfacial layer (Fig. S5–S7†). In addition to the behavior of the peak positions, the integrated intensities of the δ DOD peaks demonstrate a minimum at $\varphi = 0$ V (Fig. S21†), consistent with SEIRAS data. ¹² Further analyses of the VDOS spectra, including the spectra that emerge from the velocities only in z, are presented in the ESI.†

The VDOS spectra for layer 2 water (middle panel of Fig. 8) are significantly less sensitive to φ than are those in layer 1. Nonetheless, we observe in layer 2 small potential dependencies of the ν OD peak (Fig. S17†). It is evident from the bottom panel of Fig. 8 and S18† that in layer 3, which is comprised of water even further from the electrode surface, the effects of φ on the VDOS spectra are mostly negligible. This observation is consistent with the electric field associated with the applied potential being nearly entirely screened by the first two layers of water.

4. Conclusions

In this work we studied the interface between electrified Au(111) slabs and heavy water using several BOMD trajectories and a mixed explicit–implicit solvation approach. The electrostatic potential differences between points in the interior of the metal and deep in the continuum solvent, $\Phi_{\rm e}$, were computed regularly throughout each BOMD simulation. The cumulative trajectory averages of $\Phi_{\rm e}$ all converged within 10 ps and the relationship between the total electrode charges and their corresponding average potentials yielded a differential capacitance of 14.3 $\mu {\rm F~cm}^{-2}$ for the system, in good agreement with experimental values for Au(111) electrodes with dilute electrolytes. Recovering the differential capacitance was validation for the methodology of utilizing several fixed electrode charge BOMD trajectories to study this inert electrochemical interface.

That the BOMD trajectories reliably quantified the potential φ enabled structural and dynamic analyses of water with respect to both φ and distance from the electrode surface. Our calculations demonstrated that water forms at least one structural layer with a density greater than that of bulk against the Au(111) electrode, and more than one such layer at PZC and at positive φ . In addition, an analysis of the orientations of water characterize a general structure of water under bias in layer 1 in which one OD bond is generally oriented towards or away from the electrode depending on the sign of the applied potential whereas the other OD bond is nearly parallel to the surface. We observed that the D_{\parallel} of layer 1 water is enhanced only at potentials negative of the PZC, and that D_{\parallel} increases for all nonzero values of φ in the second and third layers of water. Lastly, the VDOS were most significantly potential-dependent in layer 1 of water, in which the δDOD and νOD peaks exhibited blue shifts for all trajectories with $\varphi \neq 0$. The δ DOD peak positions supported the observation of a weakened hydrogen bonding network at the interface.

This work represents a step towards describing *in situ* aqueous electrochemistry with *ab initio* molecular simulation. Water comprises a significant portion of the EDL even for concentrated aqueous electrolyte solutions, necessitating that we intimately understand how its properties are altered by applied voltage at equilibrium and beyond. Looking ahead, the inclusion of ions that constitute the EDL as well as reliable descriptions of driven, faradaic charge transfer processes would vastly enhance the scope of these simulations and offer valuable microscopic insights into electrochemistry and its diverse applications.

Author contributions

Z. K. G. conceived of the project; Z. K. G., M. F. C. A., and A. S. designed the model system and simulations, analyzed the results, and wrote the paper. Z. K. G. and M. F. C. A. carried out BOMD simulations and corresponding post-processing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank Marko Melander for a helpful discussion of the electrode potential and Roberto Car for support and inspiration in designing the project. This work was supported by the Computational Chemical Science Center: Chemistry in Solution and at Interfaces under DOE Award DE-SC0019394.

Notes and references

- 1 Z. W. Seh, J. Kibsgaard, C. F. Dickens, I. Chorkendorff, J. K. Nørskov and T. F. Jaramillo, *Science*, 2017, 355, eaad4998.
- 2 K. Sakaushi, T. Kumeda, S. Hammes-Schiffer, M. M. Melander and O. Sugino, *Phys. Chem. Chem. Phys.*, 2020, 22, 19401–19442.
- 3 C. Zhao and W. Zheng, Front. Energy Res., 2015, 3, 23.
- 4 K. Sivula and R. van de Krol, Nat. Rev. Mater., 2016, 1, 15010.
- 5 V. R. Stamenkovic, D. Strmcnik, P. P. Lopes and N. M. Markovic, *Nat. Mater.*, 2017, **16**, 57–69.
- 6 A. J. Bard and L. R. Faulkner, *Electrochemical Methods Fundamentals and Applictions*, John Wiley & Sons, Inc., New York, 2nd edn, 2001.
- 7 J. O. Bockris and S. U. Khan, Surface electrochemistry: a molecular level approach, Springer Science & Business Media, 2013.
- 8 O. M. Magnussen and A. Groß, *J. Am. Chem. Soc.*, 2019, **141**, 4777–4790.
- 9 M. F. Toney, J. N. Howard, J. Richer, G. L. Borges, J. G. Gordon, O. R. Melroy, D. G. Wiesler, D. Yee and L. B. Sorensen, *Nature*, 1994, 368, 444–446.
- 10 S. V. Kalinin, O. Dyck, N. Balke, S. Neumayer, W.-Y. Tsai, R. Vasudevan, D. Lingerfelt, M. Ahmadi, M. Ziatdinov,

- M. T. McDowell and E. Strelcov, *ACS Nano*, 2019, **13**, 9735–9780.
- 11 J.-J. Velasco-Velez, C. H. Wu, T. A. Pascal, L. F. Wan, J. Guo, D. Prendergast and M. Salmeron, *Science*, 2014, 346, 831– 834
- 12 K.-i. Ataka, T. Yotsuyanagi and M. Osawa, *J. Phys. Chem.*, 1996, **100**, 10664–10672.
- 13 N. Garcia-Araez, P. Rodriguez, V. Navarro, H. J. Bakker and M. T. Koper, *J. Phys. Chem. C*, 2011, **115**, 21249–21257.
- 14 C.-Y. Li, J.-B. Le, Y.-H. Wang, S. Chen, Z.-L. Yang, J.-F. Li, J. Cheng and Z.-Q. Tian, *Nat. Mater.*, 2019, **18**, 697–701.
- 15 S. Nihonyanagi, S. Ye, K. Uosaki, L. Dreesen, C. Humbert, P. Thiry and A. Peremans, *Surf. Sci.*, 2004, 573, 11–16.
- 16 W.-T. Liu and Y. R. Shen, Proc. Natl. Acad. Sci. U. S. A., 2014, 111, 1293–1297.
- 17 M. Osawa, M. Tsushima, H. Mogami, G. Samjeske and A. Yamakata, J. Phys. Chem. C, 2008, 112, 4248–4256.
- 18 J. I. Siepmann and M. Sprik, *J. Chem. Phys.*, 1995, **102**, 511–524
- 19 A. P. Willard, S. K. Reed, P. A. Madden and D. Chandler, Faraday Discuss., 2009, 141, 423–441.
- 20 Y. Zhang, H. B. de Aguiar, J. T. Hynes and D. Laage, J. Phys. Chem. Lett., 2020, 11, 624-631.
- 21 J. K. Nørskov, J. Rossmeisl, A. Logadottir, L. Lindqvist, J. R. Kitchin, T. Bligaard and H. Jónsson, *J. Phys. Chem. B*, 2004, **108**, 17886–17892.
- 22 J. K. Nørskov, T. Bligaard, A. Logadottir, J. R. Kitchin, J. G. Chen, S. Pandelov and U. Stimming, *J. Electrochem. Soc.*, 2005, **152**, J23–J26.
- 23 K. Letchworth-Weaver and T. A. Arias, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2012, **86**, 075140.
- 24 O. Andreussi, I. Dabo and N. Marzari, J. Chem. Phys., 2012, 136, 064102.
- 25 N. G. Hörmann, O. Andreussi and N. Marzari, J. Chem. Phys., 2019, 150, 041730.
- 26 J. A. Gauthier, C. F. Dickens, H. H. Heenen, S. Vijay, S. Ringe and K. Chan, J. Chem. Theory Comput., 2019, 15, 6895–6906.
- 27 G. Fisicaro, L. Genovese, O. Andreussi, N. Marzari and S. Goedecker, *J. Chem. Phys.*, 2016, 144, 014103.
- 28 F. Nattino, M. Truscott, N. Marzari and O. Andreussi, *J. Chem. Phys.*, 2019, **150**, 041722.
- 29 A. Bouzid and A. Pasquarello, J. Phys. Chem. Lett., 2018, 9, 1880–1884.
- 30 M. M. Melander, M. J. Kuisma, T. E. K. Christensen and K. Honkala, *J. Chem. Phys.*, 2019, **150**, 041706.
- 31 C. D. Taylor, S. A. Wasileski, J.-S. Filhol and M. Neurock, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2006, 73, 165402.
- 32 Y. Ping, R. J. Nielsen and W. A. Goddard, *J. Am. Chem. Soc.*, 2017, **139**, 149–155.
- 33 G. Kastlunger, P. Lindgren and A. A. Peterson, *J. Phys. Chem. C*, 2018, **122**, 12771–12781.
- 34 H. Nong, L. Falling, A. Bergmann, M. Klingenhof, H. Tran, C. Spöri, R. Mom, J. Timoshenko, G. Zichittella, A. Knop-Gericke, et al., Nature, 2020, 587, 408–413.
- 35 M. L. Clark, A. Ge, P. E. Videla, B. Rudshteyn, C. J. Miller, J. Song, V. S. Batista, T. Lian and C. P. Kubiak, *J. Am. Chem. Soc.*, 2018, **140**, 17643–17655.

Edge Article Chemical Science

- 36 Z. K. Goldsmith, M. Secor and S. Hammes-Schiffer, ACS Cent. Sci., 2020, 6, 304-311.
- 37 N. Bonnet, T. Morishita, O. Sugino and M. Otani, Phys. Rev. Lett., 2012, 109, 266101.
- 38 A. Huzayyin, J. H. Chang, K. Lian and F. Dawson, J. Phys. Chem. C, 2014, 118, 3459-3470.
- 39 L. S. Pedroza, A. Poissier and M. V. Fernández-Serra, J. Chem. Phys., 2015, 142, 034706.
- 40 L. S. Pedroza, P. Brandimarte, A. R. Rocha and M. V. Fernández-Serra, Chem. Sci., 2018, 9, 62-69.
- 41 M. Otani and O. Sugino, Phys. Rev. B: Condens. Matter Mater. Phys., 2006, 73, 115407.
- 42 O. Sugino, I. Hamada, M. Otani, Y. Morikawa, T. Ikeshoji and Y. Okamoto, Surf. Sci., 2007, 601, 5237-5240.
- 43 M. Otani, I. Hamada, O. Sugino, Y. Morikawa, Y. Okamoto and T. Ikeshoji, Phys. Chem. Chem. Phys., 2008, 10, 3609-
- 44 S. Schnur and A. Groß, New J. Phys., 2009, 11, 125003.
- 45 C. Merlet, C. Péan, B. Rotenberg, P. A. Madden, P. Simon and M. Salanne, J. Phys. Chem. Lett., 2013, 4, 264-268.
- 46 O. Andreussi and N. Marzari, Phys. Rev. B: Condens. Matter Mater. Phys., 2014, 90, 245101.
- 47 P. Giannozzi, S. Baroni, N. Bonini, M. Calandra, R. Car, C. Cavazzoni, D. Ceresoli, G. L. Chiarotti, M. Cococcioni, I. Dabo, A. Dal Corso, S. de Gironcoli, S. Fabris, G. Fratesi, R. Gebauer, U. Gerstmann, C. Gougoussis, A. Kokali, M. Lazzeri, L. Martin-Samos, N. Marzari, F. Mauri, R. Mazzarello, S. Paolini, A. Pasquarello, L. Paulatto, C. Sbraccia, S. Scandolo, G. Sclauzero, A. P. Seitsonen, A. Smogunov, P. Umari and R. M. Wentzcovitch, J. Phys.: Condens. Matter, 2009, 21, 395502.
- 48 P. Giannozzi, O. Andreussi, T. Brumme, O. Bunau, Buongiorno Nardelli, M. Calandra, C. Cavazzoni, D. Ceresoli, M. Cococcioni, N. Colonna, I. Carnimeo, A. Dal Corso, S. de Gironcoli, P. Delugas, R. A. DiStasio, A. Ferretti, A. Floris, G. Fratesi, G. Fugallo, R. Gebauer, U. Gerstmann, F. Giustino, T. Gorni, J. Jia, M. Kawamura, H. Y. Ko, A. Kokalj, E. Küçükbenli, M. Lazzeri, M. Marsili, N. Marzari, F. Mauri, N. L. Nguyen, H. V. Nguyen, A. Otero-de-la Roza, L. Paulatto, S. Poncé, Rocca, R. Sabatini, B. Santra, M. Schlipf, A. P. Seitsonen, A. Smogunov, I. Timrov, T. Thonhauser, P. Umari, N. Vast, X. Wu and S. Baroni, J. Phys.: Condens. Matter, 2017, 29, 465901.
- 49 O. Andreussi, N. G. Hörmann, F. Nattino, G. Fisicaro, S. Goedecker and N. Marzari, J. Chem. Theory Comput., 2019, 15, 1996-2009.
- 50 L. Bengtsson, Phys. Rev. B: Condens. Matter Mater. Phys., 1999, 59, 12301.
- 51 T. Brumme, M. Calandra and F. Mauri, Phys. Rev. B: Condens. Matter Mater. Phys., 2014, 89, 245406.

- 52 D. R. Hamann, Phys. Rev. B: Condens. Matter Mater. Phys., 2013, 88, 085117.
- 53 Y.-C. Lam, A. V. Soudackov, Z. K. Goldsmith and S. Hammes-Schiffer, J. Phys. Chem. C, 2019, 123, 12335-12345.
- 54 J. P. Perdew, K. Burke and M. Ernzerhof, Phys. Rev. Lett., 1996, 77, 3865-3868.
- 55 A. Tkatchenko, R. A. DiStasio, R. Car and M. Scheffler, Phys. Rev. Lett., 2012, 108, 236402.
- 56 J. Carrasco, J. Klimeš and A. Michaelides, J. Chem. Phys., 2013, 138, 024708.
- 57 A. Berg, C. Peter and K. Johnston, J. Chem. Theory Comput., 2017, 13, 5610-5623.
- 58 L. Zheng, M. Chen, Z. Sun, H.-Y. Ko, B. Santra, P. Dhuvad and X. Wu, J. Chem. Phys., 2018, 148, 164505.
- 59 S. Trasatti, Pure Appl. Chem., 1986, 58, 955-966.
- 60 S. Trasatti, Electrochim. Acta, 1990, 35, 269-271.
- 61 S. Sakong and A. Groß, J. Chem. Phys., 2018, 149, 084705.
- 62 R. Jinnouchi and A. B. Anderson, Phys. Rev. B: Condens. Matter Mater. Phys., 2008, 77, 245417.
- 63 M. F. Calegari Andrade, H.-Y. Ko, R. Car and A. Selloni, J. Phys. Chem. Lett., 2018, 9, 6716-6721.
- 64 D. M. Kolb and J. Schneider, Electrochim. Acta, 1986, 31, 929-936.
- 65 N. Vasiljevic, T. Trimble, N. Dimitrov and K. Sieradzki, Langmuir, 2004, 20, 6639-6643.
- 66 B. Garlyyev, S. Xue, S. Watzele, D. Scieszka and A. S. Bandarenka, J. Phys. Chem. Lett., 2018, 9, 1927-1930.
- 67 E. Lust, in Electrical Double Layers. Double Layers at Singlecrystal and Polycrystalline Electrodes, American Cancer Society, 2007.
- 68 H. H. Heenen, J. A. Gauthier, H. H. Kristoffersen, T. Ludwig and K. Chan, J. Chem. Phys., 2020, 152, 144703.
- 69 J.-B. Le and J. Cheng, Curr. Opin. Electrochem., 2020, 19, 129-
- 70 R. A. DiStasio Jr, B. Santra, Z. Li, X. Wu and R. Car, J. Chem. Phys., 2014, 141, 084502.
- 71 J. Le, A. Cuesta and J. Cheng, J. Electroanal. Chem., 2018, 819,
- 72 A. Soper, F. Bruni and M. Ricci, J. Chem. Phys., 1997, 106, 247 - 254
- 73 Z. Futera and N. J. English, J. Chem. Phys., 2017, 147, 031102.
- 74 J. Le, Q. Fan, L. Perez-Martinez, A. Cuesta and J. Cheng, Phys. Chem. Chem. Phys., 2018, 20, 11554-11558.
- 75 S. D. Fried and S. G. Boxer, Acc. Chem. Res., 2015, 48, 998-
- 76 J. Xu, M. Chen, C. Zhang and X. Wu, Phys. Rev. B, 2019, 99, 205123.
- 77 K. Zhong, C.-C. Yu, M. Dodia, M. Bonn, Y. Nagata and T. Ohto, Phys. Chem. Chem. Phys., 2020, 22, 12785-12793.
- 78 T. Seki, K.-Y. Chiang, C.-C. Yu, X. Yu, M. Okuno, J. Hunger, Y. Nagata and M. Bonn, J. Phys. Chem. Lett., 2020, 11, 8459-8469.