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Solvent-Adsorbate Interactions and Adsorbate-Specific Solvent Structure in Carbon Dioxide Reduction on a Stepped Cu Surface

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Abstract

In this work, the structure of water and its interactions with various carbon dioxide reduction intermediates adsorbed on a Cu(211) surface is investigated using density functional theory. We find that the presence of adsorbates has a significant and adsorbate-specific effect on the local water structure, and that solvation can stabilize adsorbate conformations different than those found in vacuum. We describe relationships between the hydrogen bonding capability of an adsorbate, the dipole moment of the adsorbate, the energetic strength of water-adsorbate interactions, and the change induced in the local water orientation by the adsorbate. Mechanistic implications are discussed. We investigate and quantify the error associated with using arbitrary locally optimized solvent structures in calculations of relevant physical quantities, such as solvated binding energies and work functions. Possible effects of thermal motion on calculations of the work function are investigated using ab initio molecular dynamics.

Introduction

The electrolyte is a critical part of electrochemical systems. It facilitates the separation of the oxidation and reduction half-reactions within the electrochemical cell, conducts ions, and directly interacts with the reaction intermediates at the electrochemical interface. Experimentally, it has been demonstrated that many properties of the electrolyte, such as pH, solvent concentration, transport properties and specific ion concentrations significantly affect selectivity and activity of various electrocatalytic processes. A systematic understanding of electrolyte effects would facilitate the rational design of the electrolyte as an integral part of the optimization of electrochemical processes. Theoretical and computational approaches play an important role towards this end, given the experimental challenges associated with investigating reaction processes in situ and operando.

The structure of water on metal surfaces has been studied for many years. Experiments on close-packed transition metal surfaces such as Pt(111) using a variety of x-ray
spectroscopic methods\textsuperscript{14} as well as Pd(111) and Ru(0001) using scanning tunneling microscopy (STM)\textsuperscript{15} reveal a predominantly hexagonal, ice-like structure under vacuum conditions and low temperatures. Theoretical investigations of these hexagonal structures using density functional theory (DFT) revealed that on more reactive surfaces such as Ru(0001), the surface water layer may be partially dissociated,\textsuperscript{16} while an intact structure without dissociation was predicted on Pt(111).\textsuperscript{14} The simple ice-like descriptions have since been called into question\textsuperscript{17} and a diverse collection of water structures on flat metal surfaces has been described.\textsuperscript{18–22} It has also been observed experimentally using STM that water binds more strongly to low coordination sites such as step edges on Pt\textsuperscript{23} and Cu,\textsuperscript{24} leading to additional features compared to flat surfaces. Experimental investigations of oxide\textsuperscript{25} and bimetallic\textsuperscript{26} surfaces have characterized the effects of various surface heterogeneities on water structure. There is also experimental evidence that adsorbed species can have specific effects on the water structure.\textsuperscript{27}

The many experimental observations have prompted theoretical investigations on water structure on elemental metals,\textsuperscript{17,18,28–35} oxides,\textsuperscript{25,36–40} bimetallic alloys,\textsuperscript{26,41} and stepped transition metal facets.\textsuperscript{32,42–44} Recent theoretical work has provided a systematic understanding of how various surface properties affect surface binding of water.\textsuperscript{45} Several studies have also investigated the effects of explicitly including pH.\textsuperscript{40,46,47} Although the reaction thermochemistry is often investigated without the explicit consideration of the electrolyte,\textsuperscript{48} it has been demonstrated that ions can significantly affect the thermodynamics due to field effects.\textsuperscript{49–51} Calculations of electrochemical reaction barriers generally must also explicitly consider at least the reacting ions, which usually requires some amount of explicit solvation.\textsuperscript{52–54} For this reason, many studies have investigated the effect of including explicit solvent structures and electric field effects in calculations on reaction mechanisms and kinetics.\textsuperscript{8,9,49–51,53–66} Of the studies including explicit electrolyte species, some use structures which are locally optimized, representing local minima in the potential energy surface, while others employ \textit{ab initio} molecular dynamics simulations to sample thermodynamic ensembles at realistic tem-
temperatures. The drawback of using only local minima is that the possible effects of thermal fluctuations are not explicitly included. However, using *ab initio* molecular dynamics simulations can be orders of magnitude more computationally expensive than studying locally optimal structures. In both cases, it is difficult to determine whether relevant configurations have been sufficiently sampled. Even with molecular dynamics, the simulation lengths used are often on the order of approximately 10-100 ps, which may not be long enough to sufficiently sample statistically independent configurations due to the correlation time scales for water relaxation near metal interfaces at room temperature, which may be on the order of tens of picoseconds or longer.67,68

Electrolyte effects can be modeled using a wide variety of atomistic and continuum approaches. In general, classical force fields and continuum implicit solvation models allow for the exploration of significantly larger system sizes at much reduced computational cost. There are several challenges involved in accurately modeling a metal-water interface using classical molecular dynamics, including the implementation of an ideally polarizable metal model which includes the detailed electrostatic induction effects in the metal, specific adsorption, and constant electrode potential.67–69 Although these challenges have been addressed in several applications of classical force fields to metal-water interfaces,67–71 the use of such classical force field methods involves the careful implementation of sophisticated models which may be less readily transferable to new systems than *ab initio* models. Implicit solvation has also been successfully applied to the study of water-metal interfaces. Although methods such as joint density functional theory72 and reference interaction site model73 are capable of calculating the equilibrium structure of solvent in combination with DFT calculations, most implicit solvent methods used in combination with DFT74–77 by design do not provide finely detailed structural information about the solvent at atomic resolution. Understanding atomic-level detail in the solvation structure can provide important understanding of how the specific solvation structure may affect energetics and mechanisms.

In this study we use DFT and an explicit solvent model of the metal-water interface cou-
pled with a global optimization algorithm to study the structure of water and its interactions with adsorbates on a stepped Cu(211) surface. We focus on a set of key adsorbed intermediates relevant to the CO\textsubscript{2} and CO reduction reactions\textsuperscript{50,57,78}. We apply a constrained minima hopping algorithm\textsuperscript{79,80} to thoroughly sample the local minima inherent to the potential energy surface of the solvated systems. We characterize the water structure near Cu(211) and how it is changed by the presence of adsorbates, and we describe systematic relationships between these structural changes and the energetics associated with the adsorbate-water interactions. We quantify errors associated with the use of explicit solvent in DFT calculations by quantitatively comparing physical quantities of the global minima with those associated with statistics from the distributions of local minima results. We also consider the possible effect of thermal motion on calculations of the work function.

**Computational Details**

Density functional calculations were performed using the Quantum ESPRESSO electronic structure suite\textsuperscript{81}, using the Atomic Simulation Environment (ASE) as a user interface\textsuperscript{82}. The dispersion-corrected Bayesian Error Estimation Functional BEEF-vdW\textsuperscript{83–86} was used as the exchange-correlation functional. The Vanderbilt ultra-soft pseudopotentials\textsuperscript{87–90} were used to represent the core electrons, and the valence electrons were expanded in a plane wave basis, and the Brillouin zone was sampled using a Monkhorst Pack (MP) k-point grid\textsuperscript{91}. Details are provided in the Supporting Information (SI).

Global optimization of the water structures was carried out using a modified version of the constrained minima hopping algorithm\textsuperscript{79,80}. It should be emphasized that application of minima hopping does not guarantee that the global minimum configuration will be found. Moreover, the very large number of potential local minima in systems of this size\textsuperscript{79,92} combined with the computational expense of DFT, complicates determining convergence of the results relative to number of iterations. This is further discussed in the SI. *Ab initio* molec-
ular dynamics calculations were performed at constant temperature, number of particles, and volume (NVT) as implemented in ASE. The hydrogen bonding interactions were investigated using an analysis based on the bond orders as calculated using properties of the charge density with Chargemol and GPAW. The hydrogen bonds drawn in the atomistic visualization figures were produced using simple geometric criteria validated by comparison with the bond order analysis. Details are provided in the SI.

Results and Discussion

Water Structure on Clean Cu(211) Surface

We investigated the structure of water on clean Cu(211) surfaces with 5 and 10-H$_2$O per supercell, containing 9 surface atoms per simulation supercell. Figure 1 shows the structures corresponding to the global minima. A single water molecule per supercell adsorbs oxygen-down on the step edge. Since the supercell contains 3 step edge sites, the coverage on the step edge sites is 1/3. This low coverage of the step edge results from the competition between water-water and water-surface interactions. Although the interaction of the water monomer with the step edge is favorable, the hydrogen bonding network favors a structure which promotes binding to sites other than step edges. An implication of this relevant to catalysis is that adsorption of a reaction intermediate on a step edge does not require the displacement or desorption of any step-adsorbed water, since 2/3 of the step sites are vacant. Higher energy, less stable structures with 2/3 step edge water coverages were also found, and the most stable of these structures and energies are provided in the SI, but no stable structures with full water coverage on the step edge were found.
Figure 1: Global optimal structures of water on clean Cu(211). a: 5-H$_2$O per supercell. b: 10-H$_2$O per supercell). The three panels show the projection of the structure in three orthogonal directions. The white rectangle indicates the periodic boundaries of the supercell.
Figure 1 shows that H$_2$O is oriented hydrogen-down at terraces and hydrogen-up at step-edge sites. On the solvated Cu(211) surface, the step edges carry a small partial positive charge where H$_2$O is adsorbed hydrogen-up, while the terrace sites carry a small partial negative charge. Results of Bader charge analysis are shown in the SI. The hydrogen bonding network is dominated by repeating pentagonal motifs. The oxygen-down adsorption at step edges, hydrogen-down interactions at terrace sites, and presence of pentagonal rings resemble results reported in a recent joint experimental-DFT study on Cu(511). This suggests that step features on facets other than 211 can have a similar effect on the interfacial water structure, and that the results reported in this work can be expected to extend to various types of Cu step sites.

In the most stable 10-H$_2$O configuration, also displayed in Figure 1, a similar structure near the surface is observed. The coverage adjacent to the surface remains similar to that used in the 5-H$_2$O calculations. This optimal structure illustrates that the water beyond the interfacial layer participates readily in hydrogen bonding with the interfacial water layer, in contrast to the hydrophobic interfacial layer described near some more strongly binding metal surfaces. This connectivity of the hydrogen bond network may be of relevance to proton transport mechanisms near the surface. A five-membered or pentagonal hydrogen-bonded ring structure is present in the 10-H$_2$O per supercell configuration, as in the 5-H$_2$O per supercell configuration. The preferential oxygen-down orientation at approximately 1/3 of the step edges and hydrogen-down orientation at terrace sites described for the 5-H$_2$O per supercell results is retained with the additional water molecules present.

**Solvation Effects on Binding Energies**

The solvated binding energy $\Delta G_{\text{bind}}^{\text{solv}}$ of an adsorbate can be expressed as the sum of two contributions:

1. energy of cavity formation $\Delta G_{\text{cav}}^{\text{solv}}$: the energy associated with reorganization of solvent from its relaxed equilibrium configuration with no adsorbate present into the solvation
shell configuration which it assumes in the presence of a given adsorbate, and

2. energy of solvated adsorbate insertion $\Delta G_{\text{ins}}$: the energy associated with insertion of the adsorbate from the gas phase into its minimum energy configuration, bound to the surface, within the already-configured equilibrium solvent shell.

The solvation energy $\Delta G_{\text{solv}}$ is the difference between the binding energies in solvent and vacuum, as written in equation 1.

$$\Delta G_{\text{solv}} = \Delta G_{\text{bind}} - \Delta G_{\text{bind}}^{\text{vac}} = \Delta G_{\text{cav}} + [\Delta G_{\text{ins}} - \Delta G_{\text{bind}}^{\text{vac}}]$$ (1)

Defining a “pre-organized solvation energy” $\Delta G_{\text{solv}}^{\text{pre}}$ as the term enclosed by brackets in equation 1, we decompose the solvation energy into two convenient terms,

$$\Delta G_{\text{solv}} = \Delta G_{\text{cav}} + \Delta G_{\text{solv}}^{\text{pre}}$$ (2)

where $\Delta G_{\text{solv}}^{\text{pre}}$ quantifies the strength of the interaction between the adsorbate and the already-organized optimal solvation shell. Quantifying the energies associated with solvent reorganization $\Delta G_{\text{cav}}$ and solvent-adsorbate interaction $\Delta G_{\text{solv}}^{\text{pre}}$ can be useful in more thoroughly understanding the solvation of adsorbed intermediates.

The effect of water on a few key elementary reaction energies relevant to CO$_2$ reduction catalysis was evaluated by comparing the reaction energies obtained including the explicit solvent in the model to the energies obtained in vacuum. The decomposition of solvation energy according to equation 2 is obtained by ‘single point’ energy calculations on structures derived from the full local minima structures by deleting specific atoms while holding all other atoms fixed, as described in the SI. The results for all adsorbates considered are summarized in Figure 2. The effects on the reaction free energy diagram are presented later in Figure 4. A noticeable yet small difference between 5 and 10-H$_2$O models is seen only in the larger C$_2$ adsorbates OCCO and OCCHO, which due to their size may extend further into the solvent.
phase than the smaller adsorbates do.

![Figure 2: Contributions to solvation energy of adsorbed intermediates. a: 5-H\(_2\)O / supercell. b: 10-H\(_2\)O / supercell). All quantities are Boltzann-weighted averages.](image)

The energetic cost of organizing the solvent into the optimal solvation shell can be quite large, on the order of 0.5 to 1 eV. The large cavity formation energy is evidence that adsorbates can induce significant changes in the local structure of water. The energetic cost of these structural changes scales closely with the energetic favorability of the adsorbate interactions with the new adsorbate-induced solvation structure. There is thus a significant cancellation between the cavity formation and insertion energies. More energetically favorable interactions between the adsorbate and the water (more negative \(\Delta G^{\text{pre}}_{\text{solv}}\)) are correlated with more extensive restructuring of the water near the adsorbate (more positive \(\Delta G_{\text{cav}}\)). The trends in \(\Delta G_{\text{cav}}\), and \(\Delta G^{\text{pre}}_{\text{solv}}\) for the 5-H\(_2\)O and 10-H\(_2\)O configurations are similar. Trends in \(\Delta G_{\text{solv}}\) are more subtle and difficult to confidently assess due to their smaller size, especially since the magnitudes of errors in DFT calculations for nonbonded interactions are on the order of 0.1 eV. The magnitude of the solvation energy roughly increases with adsorbate size in the 5-H\(_2\)O calculations, but this trend is not as clear in the 10-H\(_2\)O case, and OCCHO appears to be an exception to this trend in both cases. The physical significance of the energetic trends is more thoroughly explored in Figure 3 and the subsequent discussion.
Adsorbate-Specific Solvation Structure

The presence of adsorbates induces changes in the local solvent structure relative to that on the clean stepped Cu surface, as illustrated in Figure 3(a). More detailed images of the global minimal structures are available in the SI, as are direct comparisons of the 5-H$_2$O and 10-H$_2$O results.

Figure 3: Analysis of solvation structure. a: Illustration of global minima structures. b: Global minimum, Boltzmann-weighted average, and unweighted local minima distribution of water dipole potential $\Delta \Phi_{\text{orient}}$. c: Global minimum, Boltzmann-weighted average, and unweighted local minima distribution of hydrogen bonding characteristics. d: cavity formation energy $\Delta G_{\text{cav}}$ versus total hydrogen bonds for all adsorbates considered. e: Water dipole potential $\Delta \Phi_{\text{orient}}$ versus adsorbate vacuum dipole moment $\mu_{\text{ads}}$ in Debye for all adsorbates considered. f: Absolute value of the water dipole potential versus the cavity formation energy for all adsorbates considered. All quantities in d., e., and f. are Boltzmann-weighted averages over all local minima. Only the 10-H$_2$O data is shown in b. and c., and comparisons to 5-H$_2$O data are in the SI.

The global minimal structures displayed in Figure 3(a) suggest that hydrogen bonding can be a significant contributor to adsorbate-water interactions. In particular, COH donates a hydrogen bond to a nearby water, while CHO, OCCO, and OCCHO all accept at least one hydrogen bond and do not appear to donate any. The CO adsorbate does not participate in
hydrogen bonding to any appreciable extent. This is consistent with the observation from Figure 2 that CO interacts quite weakly with the solvent.

The results of a more quantitative analysis of the hydrogen bonding characteristics is displayed in Figure 3(c). The bars in Figure 3(c) constitute histograms of the unweighted distributions of hydrogen bond numbers in all the local minima. The unfilled bars indicate the number of hydrogen bonds accepted by the adsorbate and the filled bars indicate number of hydrogen bonds donated by the adsorbate in each local minimum configuration. The vertical lines overlayed on these histograms indicate the calculated values of the Boltzmann-weighted averages over all local minima, as well as the values at the global minima, of the number of hydrogen bonds accepted by and donated by each adsorbate as indicated in the legend. The agreement between the results evaluated at the global minima and the Boltzmann-weighted averages over all local minima indicates that the global minima structures shown are representative of the Boltzmann-weighted average results. The correlation between $\Delta G_{\text{cav}}$ and the total number of adsorbate-water hydrogen bonds, plotted in Figure 3(d), indicates that the energetic cost of solvent restructuring in response to the presence of the adsorbate scales with the extent of hydrogen bonding between the solvent and the adsorbate.

To quantify how the presence of adsorbates affects the local orientation of water, we calculate the electrostatic potential energy change across the isolated water molecules in the absence of the metal surface and the adsorbate. We refer to this here as the ‘water dipole potential’ $\Delta \Phi_{\text{orient}}$ where the subscript emphasizes that it is the orientation contribution only, consistent with previous work involving this quantity. The quantifies the effect of water orientation on the work function of the surface, neglecting the polarization contribution. The dominant contribution to this potential drop is the dipole moment density of the water molecules. A positive water dipole potential as defined here corresponds to a net hydrogen-down structure, while a negative water dipole potential corresponds to a net hydrogen-up structure.
The structures associated with clean Cu(211) as well as adsorbed CO show nearly zero net water dipole orientation. The result for clean Cu(211) is consistent with previous studies of water near metal electrodes.\textsuperscript{29,34} The observation that CO induces only a very small change in the local water orientation is once again consistent with its very weak interaction with the solvent (Figure 2). In contrast, the other adsorbates induce significant changes in the local water orientation. COH induces a positive water dipole potential, while CHO, OCCO, and OCCHO induce negative water dipole potentials. This indicates that COH induces a net hydrogen-down structure, while CHO, OCCO, and OCCHO induce net hydrogen-up structures, at the coverages studied. A correlation between the water dipole potential and the adsorbate vacuum dipole moment is apparent in Figure 3 (e). This means that the average dipole orientation induced in the water is antiparallel to the direction of the adsorbate dipole. The most stable orientation of a dipole in response to the field created by another spatially fixed dipole depends on its position relative to the fixed dipole and its axis. Directly vertically above the fixed dipole along its axis, the most stable alignment is parallel, while in the plane bisecting the axis of the fixed dipole the most stable alignment is antiparallel. In this case, where the adsorbate can be thought of as having a fixed dipole and the water molecules are relatively free to reorient in response to its presence, there are more water molecules laterally beside the adsorbate on the surface than there are vertically directly above it, and thus on average the antiparallel alignment is stabilized. This suggests that the correlation described here is expected to be specific to this particular adsorbate coverage, and could change as adsorbate coverages increases.

The magnitude of the water dipole potential is correlated with the magnitude of the cavity formation energy, as shown in Figure 3 (f). Coupled with the correlation between the adsorbate dipole moment and the water dipole potential, this suggests that the electrostatic interaction between the dipole moment of the adsorbate and the local water structure has a role in the restructuring of the water in response to the presence of an adsorbate. Note that in these plots $\Delta G_{\text{solv}}^{\text{pre}}$ and $\Delta G_{\text{cav}}$ could be used essentially interchangeably except for a
sign change, since those quantities are also correlated (Figure 2). Together, the correlation plots in Figure 3(d-f) suggest that the local solvent structure restructures in response to the presence of an adsorbate by forming hydrogen bonds with the adsorbate and aligning with the field induced by the adsorbate dipole moment, and that the energy associated with this solvent restructuring $\Delta G_{cav}$ and with the resulting water-adsorbate interactions $\Delta G_{solv}^{\text{pre}}$ both scale with these structural descriptors.

**Mechanistic Implications**

The specificity of the local water structure to each adsorbed intermediate suggests that elementary reaction steps, such as proton transfers and C-C coupling, are accompanied by rearrangements in the local water structure. This adsorbate-specific solvation structure may affect the energetic barriers associated with relevant elementary reaction steps in several ways. Even if solvation does not significantly change the relative stabilities of the initial and final states of an elementary step, it could have an effect on the kinetics if significant reorganization is required due to different solvation structures in the initial and final states, since the solvent re-orientation energy barrier contributes to the barrier associated with the elementary step.

As an example of the possible mechanistic significance of solvent re-organization, we consider two possible C-C coupling reactions: $2{\ast CO} \rightarrow {\ast OCCO}$ and ${\ast CO} + {\ast CHO} \rightarrow {\ast OCCHO}$. In Figure 3(b,c), it is clear that both ${\ast OCCO}$ and ${\ast OCCHO}$ induce hydrogen-up local water structures and accept approximately two hydrogen bonds. Examining the reactants, ${\ast CO}$ is associated with a neutral water structure and accepts no hydrogen bonds, while ${\ast CHO}$ induces a hydrogen-down local water structure and accepts one or two hydrogen bonds. Thus, the solvation structure of ${\ast CHO}$ is quantitatively similar to that of the corresponding final state, ${\ast OCCHO}$, while the solvation structure of ${\ast CO}$ is quite different than that of the corresponding final state, ${\ast OCCO}$, in hydrogen bonding, orientation, and cavity formation energy. This suggests that the water re-organization contribution to the barrier may
be greater in CO-CO coupling than in CO-CHO coupling, and this solvent re-organization contribution would promote C-C coupling of *CHO relative to *CO. Previous computational work has indicated a lower activation barrier for C-C coupling with increasing hydrogenation of the reactants even when explicit solvent is not considered, and the present results indicate that the effects of solvent re-organization could also contribute to this trend.

Thermal fluctuations in the solvation structure could likewise influence the subsequent elementary reaction steps by pre-solvation, which can play an important role in the rates of many aqueous processes, including proton transfer, in an analogous manner to the role of solvent reorganization in electron transfer reactions. For example, when instantaneous thermal fluctuation of the water structure near adsorbed CO cause a momentarily hydrogen-up structure, then this pre-organized solvation structure may promote the formation of CHO via proton transfer relative to COH, since CHO is stabilized by a locally hydrogen-up solvation structure while COH is stabilized by a locally hydrogen-down solvation structure. It is also plausible that proton transfers to the surface are more facile from predominantly hydrogen-down water structures, while a predominantly hydrogen-up water structure could increase this barrier since a water molecule must rotate before the proton can be transferred. A shortcoming of this study is that the explicit solvent effects on the transition state structures are not studied, and the dynamic effects of pre-organization and reorganization on reaction events are not directly studied. This motivates future work focused on understanding the kinetic effects of the local electrolyte structure.

**Comparison of Global Minima to Local Minima Distributions**

In addition to searching for a global minimal structure, the minima-hopping configuration sampling procedure finds a diverse collection of local minima. In this work the minima hopping procedure was continued until the distributions of energy from this sampling method were observed to relatively converge as described in the SI. Assuming that the collection of local minima found in this procedure is roughly representative of the collection of inherent
local minima in the potential energy surface of the system, we can compare results obtained using statistics of this collection to those evaluated at the global minima. This is of practical importance because many DFT calculations which include explicit solvent structures do not employ exhaustive global minimization or configuration sampling procedures, as was done in this work. A comparison of results obtained from statistics of the representative local minima distribution to those obtained from global minimization is a step towards quantifying the accuracy and precision of using solvent structures which have not been thoroughly globally minimized. Of primary interest are the free energies and the work functions, which are shown in Figures 4 and 5, respectively. We emphasize that this error analysis is associated with the computational explicit-solvent model itself, rather than a comparison to experimentally measured quantities.

Figure 4: Comparison of free energies of adsorbed intermediates calculated from the global minima structures to those computed from statistics of the local minima distributions. The error bars quantify one standard deviation of the results computed from the local minima distributions.

There is a distribution of possible calculated free energies for each adsorbate which arises from the collection of possible solvation structures. Figure 4 illustrates that the expectation values of these distributions are quite close to the physically relevant values given by the global minima. For CO, COH, and CHO, the results found using the median or arithmetic mean of the local minima distributions are nearly indistinguishable from those from the
global minima. For OCCO and OCCHO, using the median or mean tends to underpredict the effect of solvation, resulting in a slightly less stable energy than does the global minimum result. The largest error is in the prediction of the OCCO solvated binding energy, for which the error is approximately 0.15 eV in the 5-H$_2$O per supercell case and approximately 0.1 eV in the 10-H$_2$O per supercell case. In all other cases, the difference between the median and mean results and the global minima results is less than 0.1 eV. The bias for the clean solvated surface is zero here because the corresponding solvated surface is always used as the reference for the binding energies. The mean and median operations are also applied to these reference states. This means that the reference energies corresponding to the median and mean structures are greater than the reference energies corresponding to the global minima. This is why the binding energies calculated from the local minima distributions can be lower than those calculated from the global minima.

Convergence of the expectation values of the local minima results to the physically correct values is practically useful only if the variances of these distributions are also relatively small, such that a small number of samples drawn from this collection could yield a reasonable result. The error bars in Figure 4 indicate the positions of one standard deviation above and below the given statistic on the free energy diagram. This is equivalent to approximately a 68% prediction interval for a given free energy if only a single structure is sampled, at random, from the collection of possible local minima. This quantifies the expected deviation from the global minimum result caused by using an arbitrarily chosen solvation structure. For the 5-H$_2$O per supercell case, these standard deviations are on the order of 0.1 to 0.15 eV, while for the 10-H$_2$O per supercell case, the standard deviations are slightly higher, but always less than 0.2 eV. This suggests that as more water molecules are included, the variance introduced into the calculation by using an arbitrary local minimum solvent structure increases.

A similar analysis can be applied to the work function, which is often used in DFT calculations to quantify the absolute electrode potential.$^{52,105}$ Figure 5 displays the distribution of work functions from the collection of local minima structures for each adsorbed interme-
diate. Except in the case of CO with 5-\text{H}_2\text{O} per supercell, the expectation values of these distributions are quite close to the work functions of the global minimum structures.

![Graph showing work functions of solvated Cu(211) with adsorbed intermediates calculated from the global minima structures to those computed from the local minima distributions.]

Figure 5: Comparison of work functions of solvated Cu(211) with adsorbed intermediates calculated from the global minima structures to those computed from the local minima distributions.

The correspondence between the expectation values of the local minima distributions and the global minima results indicates that the work function at the global minimum structure may be approximated by sampling a number of local minima from a representative sample. In practice, since the variance is quite large, the work function of a single arbitrarily chosen local minimum structure is often quite different from that at the global minimum. This underscores the importance of sufficiently sampling solvent configuration when the physical quantity of interest depends very sensitively on the details of the solvent structure, as is the case with the work function.\(^9\)

Finally, we consider again the hydrogen bonding results shown in Figure 3(c). The unweighted distributions of hydrogen bonding characteristics from the entire collections of local minima sampled reveal that the higher-energy local minima also generally display
hydrogen bonding characteristics qualitatively similar to those of the global minima: the
adsorbates with global minima that accept hydrogen bonds have local minima collections
which predominantly accept hydrogen bonds, and those with global minima that donate
hydrogen bonds have local minima collections which predominantly donate hydrogen bonds.
This is of practical importance because it means that structural conclusions drawn from the
study of arbitrarily chosen local minima may often be consistent with those drawn from
rigorously optimized global minima structures.

The Effect of Thermal Motion on the Work Function

The results discussed in the previous sections have focused on the local minima in the po-
tential energy surface of the solvent structure. As the temperature is increased from 0 K to
more experimentally relevant temperatures, thermal excitations or fluctuations away from
these local minima become increasingly important. To evaluate the possible effects of ther-
mal motion on the structure, we performed short \textit{ab initio} molecular dynamics simulations
starting from three arbitrarily chosen distinct local minima of the clean surface with 5 H\textsubscript{2}O
per supercell and no adsorbate. For each of these molecular dynamics simulations, the work
function recorded at each time step. Each system was equilibrated with a thermostat set
at 300 K for 500 fs. After equilibration, the simulation was continued for another 1500 fs,
during which equilibrium statistics were recorded. Details are provided in the SI. The work
function results are summarized in Figure 6.
Figure 6: Comparison of work functions at local minima to ensemble averages and fluctuations from *ab initio* molecular dynamics simulations.

For all three trials, the time-averaged work function is comparable to the work function of the starting local minimum. This suggests that at time scales on the order of picoseconds, the time-averaged work function associated with a trajectory started from a given local minimum is similar to the work function evaluated at the corresponding local minimum. Therefore, there does not appear to be a significant systematic error in the work function calculation introduced by using its value at a local minimum structure rather than its ensemble average value at a nonzero temperature. On the other hand, this initial investigation indicates that thermal fluctuations can be substantial and motivates a more thorough investigation of thermal motion in future work.

The relative fluctuations are expected to be sensitive to the size of the simulated system. Here the relevant dimension is the geometric surface area of the electrochemical interface per supercell, which in this case is approximately 49 Å². These molecular dynamics simulations were carried out on clean Cu(211) with five rigid water molecules per supercell. We also emphasize here that understanding the accuracy and applicability of DFT for the structure and properties of liquid water is also an active research area. These considerations motivate further study of the effects of thermal energy on solvation structure near adsorbates.
Conclusions

In this work, the solvent structure near adsorbed carbon dioxide reduction intermediates on Cu(211) was investigated using DFT calculations coupled with a global minimization algorithm. We find that each adsorbate has a significant and unique effect on the local water structure. We quantify and systematically explain these adsorbate-specific structural changes by considering the relationships between the adsorbate-water hydrogen bonding, adsorbate dipole moment, the change in water orientation induced by the adsorbate, and the energetic strength of water-adsorbate interactions. Polar adsorbates can accept or donate hydrogen bonds and induce a net dipole moment orientation change locally in the solvent. The magnitude of this orientation change scales with the energetic cost associated with the organization of the solvent to the presence of an adsorbate, which in turn is closely correlated with the interaction strength between the water and the adsorbate. This analysis of adsorbate-specific solvation structure represents a step towards a systematic understanding of how changes in solvation structure interact with elementary reaction steps at electrochemical interfaces, providing initial mechanistic insight and directions for future work.

By comparing results obtained from global minima structures to results found by sampling the local minima distributions, we quantify the accuracy and precision associated with the common practice of using arbitrary locally optimal solvent structures without employing a thorough global minimization procedure. We find that the the solvation energies calculated from the global minima structures can be reproduced with reasonable accuracy (the largest error is approximately 0.15 eV, and most errors are less than 0.1 eV) and reasonably small variance using arbitrarily sampled local minima. The standard deviation of the local minima results is found to be on the order of 0.15 eV for 5 to 10 explicitly included water molecules. This roughly quantifies the error introduced in studies which use explicit solvent structures without thorough global minimization procedures. Comparisons to experimentally measured solvation energetics are not made here, and the error analysis is associated with the computational method itself.
We find that the work function corresponding to the global minima structures can also be reproduced by sampling arbitrary local minima, but in this case the variance is quite large due to the sensitive dependence of the work function on the detailed solvent structure. This indicates that for quantities which depend sensitively on the specific details of the solvent structure, thorough configurational sampling is crucial to ensure physically meaningful results. The variance in work function also motivates the future investigation of solvent dynamics using hybrid continuum-atomistic models or other controlled work function methods.

In our investigation of the possible effects of thermal motion on the water structure using \emph{ab initio} molecular dynamics, we find that at short time scales on the order of 2 ps the time-average of the work function is relatively close to the work function at the corresponding local minimum, while the substantial fluctuations observed provide motivation for future work to more comprehensively study these effects.

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**Supporting Information Available**

All data plotted in figures, computational details, convergence tests, hydrogen bonding analysis, Bader charge analysis, detailed images and descriptions of adsorbate structures, com-
parisons of 5 and 10-H$_2$O per supercell results, details of single point calculations, free energy corrections and other corrections and calculations used.

This material is available free of charge via the Internet at [http://pubs.acs.org/](http://pubs.acs.org/).

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