Molecular Properties and Chemical Transformations Near Interfaces

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1. INTRODUCTION

Chemistry rarely occurs in a homogeneous aqueous phase but, instead, occurs inside niches, within crevices, between surfaces, and at impurity sites involving interfaces between two or more phases of gases, liquids, or solids. Interfacially induced changes of molecular properties feed up at an operational level to a diverse range of chemical, biophysical, and applied environmental and energy systems including the following: catalysis for targeted energy reactions,7 intergranular corrosion in metals,8 aerosol and atmospheric particles,9,10 and reaction rate acceleration in nanodroplets11 and microdroplets.12 However, at present, we lack the ability to accurately predict most molecular processes in such systems because interfacial and confinement effects violate our understanding of these same processes gained from bulk phase studies. In particular, when the correlation lengths in space and/or time of the interfacial system become comparable to that of a given molecular property, how that property is altered will now depend sensitively on the specifics of the interfacial geometry and chemistry. In this Feature Article, our team seeks to unify the aspects of interfacial features that are predictive for functional outcomes by a systematic strategy of changing the nature of the interface from a prototypical air–water interface,13,14 and then expanding into more complex condensed phase environments.

The structural organization and thermodynamics of ion solvation as well as ice nucleation and growth15 under complex environments are demonstrably different from those in the bulk phase. In many real-world applications, the active role that water ubiquitously plays in interfacial chemistry16 also is an important theme as we contrast air–water vs solid–water interfaces to better reveal how key structural motifs change the thermodynamic behavior of ion solvation at the gas vs solid interface.17,18 With careful studies of ice nucleation in the bulk,19 we contrast nucleation and growth properties of water under confinement between graphene sheets20 or at grain boundaries21 which has important implications for chemical reactions on ice22−25 and gaseous SO2 uptake in ice.26

We also discuss the elucidation of the principles that control reactivity in heterogeneous environments where interfacial processes are inextricably connected to chemical transformations such as “on-water” chemistry and other types of fundamental organic reactions that have been found to be catalyzed by microdroplet interfaces.17,28 Classic organic reactions such as retro-Diels−Alder help to elucidate the effects of nanoconfinement on solvent properties and their effects on reactivity,29 as well as the diffusive and mesoscale confinement in droplets and aerosols that can alter overall reaction rates and...
mechanisms. Heterogeneous catalysis and electrocatalysis enable important chemical reactions that are thermodynamically uphill, such as the splitting of water to produce H₂ and the reduction of CO₂ to fuels, to be carried out at ambient conditions. These interfaces under in situ conditions structure32 and drive chemical transformations through multiple intermediates33 and pose a remarkable series of challenges for theory and computation to elucidate the sequence of events at the molecular scale.34−36

We highlight the theoretical and experimental advances that have enabled our team to tackle systems that are heterogeneous and profoundly complex in composition. There is a substantial experimental gap in methods that can achieve microsecond and submicrosecond mixing times, have critical time scales for observing reactive intermediates (such as radicals) in condensed phase reactions, and require an integrated spectroscopy platform with a colliding droplet reactor to probe fast events. In addition to seminal work using soft X-rays coupled to liquid jets,38 the recently developed velocity map imaging (VMI)39 method uses X-ray photoelectron spectroscopy to probe aerosols and nanoparticles.40,41 Ambient-pressure XPS (APXPS) with soft (<2 keV) and tender X-rays (2−6 keV)33,42−46 has enabled the ability to probe solid−liquid interfaces and is ideal for buried interfaces between tens of nanometers thick liquid electrolytes. We have developed new theoretical approaches for understanding the nonequilibrium dynamics in clays47 and confined aqueous systems,48 the transport properties and phase behavior of glasses,49 and the self-assembly of amphiphilic structures.50 Finally, we summarize our Feature Article with new experimental and theoretical efforts that will be needed to more seamlessly connect structure and dynamics that cross disparate scales in order to better understand emergent phenomena for interfacial systems in the future.

2. ION SOLVATION AT INTERFACES

Water’s ability to favorably solvate ions and polar molecules, and its inability to accommodate large nonpolar species, underlies many of its essential roles in shaping reactivities central to energy sciences and biophysics. A century of quantitative measurements, theoretical development, and modeling efforts have established a basic mechanistic view of aqueous solvation and many tools to explore its molecular underpinnings and chemical implications. Molecular dynamics simulations can, for instance, describe bulk ion solvation free energies with good to excellent accuracy,51 given a careful treatment of long-range electrostatic forces via Ewald summation conducted under periodic boundary conditions. This approach has been extremely reliable for structural, thermodynamic, and transport properties, once finite size corrections are applied.52

In many ways, however, our understanding of ion solvation remains markedly incomplete. Dielectric continuum theory (DCT), which embodies the classic physical perspective on charged solutes in polar solvents, fails to capture large thermodynamic distinctions between the solvation of positively and negatively charged ions that are otherwise identical. Spatially heterogeneous environments compound these shortcomings significantly, bringing into play modes of solvent response that can be neglected in the symmetric environment of a bulk liquid. Key interfacial behaviors, such as the surface enhancement of some anions and the exclusion of corresponding cations,53 lack a thorough understanding as a result, and a general theory for ion solvation at surfaces is still wanting. Surface-specific spectroscopies that might clarify the situation are both challenging to perform precisely and notoriously difficult to interpret reliably in microscopic terms.

To help address this gap, Geissler and Saykally have studied the adsorption of prototypical anions to the air−water interface by using the surface-selective method of second harmonic generation spectroscopy (SHG) along with computer simulations. By measuring SHG intensities as a function of bulk ion concentrations, simple Langmuir adsorption models yield the Gibbs free energy of adsorption, whose temperature dependence permits separation into enthalpic and entropic contributions. For the thiocyanate anion (SCN⁻), a prototypical chaotrope, surface adsorption is unexpectedly favorable enthalpically, which can be ascribed to a repartitioning of solvent molecules among bulk, interfacial, and ion hydration shell environments. The measured entropy change is negative (unfavorable), which we have ascribed to a suppression of

Figure 1. Solvation of a model ion at the interface between liquid water and graphene. (A) Snapshot from a molecular dynamics simulation, with the air/water (upper) and graphene/water (lower) interfaces rendered as smooth surfaces identified from a coarse-graining of the microscopic density field. Note the substantial suppression of capillary fluctuations at the lower interface. (B) Profiles of key thermodynamic quantities as functions of the solute’s distance z relative to the graphene sheet, computed by umbrella sampling. Features at small and large z highlight differences in solvation at the two interfaces. The entropy profile is nearly constant at small z (aside from a drop at the left-most point, where the solute begins to clash sterically with the graphene sheet), consistent with a reduced significance of interfacial shape fluctuations. Reproduced with permission from ref 18. Copyright 2017 National Academy of Sciences.
fluctuations in interfacial shape. As a route to further clarifying the mechanism that selectively drives ions to and away from the air/water interface, the Saykally group has developed a new experiment (deep UV sum frequency generation) for measuring the complete charge transfer to solvent (CTTS) spectrum of interfacial anions and has applied it to the prototypical cases of SCN− and the iodide anion. The spectra show significant differences from the bulk CTTS spectra and provide a new variable for constructing general theoretical models to explain interfacial ion behavior.

Geissler and Saykally have tested the generality of their mechanistic picture for ion adsorption at the air−water interface by examining solution in a different interfacial context, namely liquid water in contact with graphene. Deep UV second harmonic generation (SHG) measurements of the SCN− ion determined a free energy of adsorption at graphene within experimental error of that determined for air/water, in accord with the findings from molecular simulations that model anions absorb almost equally strongly at these two interfaces. Interestingly, however, computer simulations indicate that the enthalpic and entropic components of the adsorption free energy differ significantly in the two cases (Figure 1). Whereas the adsorption energy at the air/water interface is dictated by repartitioning of solvent molecules, direct interactions between the ion and graphene are dominant in the case of graphene/water. Moreover, interface shape fluctuations are suppressed by graphene even in the ion’s absence, so that the entropic consequences of capillary wave pinning by the ion are attenuated considerably.

Because long-range electrostatics figure prominently in aqueous solvation, quantifying ion adsorption thermodynamics in computer simulations requires careful consideration of finite size effects. To extrapolate from the nanometer simulation scale £ιm, one can presume that a macroscopic description like DCT applies on all larger scales. In the case of solvation in bulk liquid water, this strategy yields a simple yet accurate finite size correction. Geissler and co-workers derived an analogous correction for the interfacial case, obtaining similarly close agreement with observed system size dependence. In addition to providing a practical computational tool, successful extrapolation from small simulations (£ιm ∼ 1 nm) highlights the realism of DCT on scales larger than 1−2 molecular diameters, even for heterogeneous systems. Failures of DCT for predicting an aqueous response on smaller scales indicate the inappropriate omission of short-range physical features of water. Geissler and Mandadapu have recently shown that molecular quadrupole contributions can indeed largely account for errors in predicted ion solvation free energies and have suggested an elaboration of DCT that accounts implicitly for the geometric relationship between water’s dipole and quadrupole moments.

3. ICE NUCLEATION AT INTERFACES

While effective control of ice nucleation under different environmental conditions is highly desirable, in reality it has proven to be incredibly challenging. This is partly because water does not usually freeze at its equilibrium melting point but can exhibit significant supercooling and the resulting metastability makes it difficult to control the ice nucleation event at the desired temperature. Previous work has shown that static electric fields have been effective in controlling ice nucleation without the need for nucleating agents, but the highest freezing temperature (Tf) of water that can be achieved in an electric field (E) was still uncertain until recently when Ahmed and co-workers performed a systematic study of the effect of an electric field on water freezing (Figure 2). By varying the thickness of a dielectric layer and the voltage across it in an electrowetting system, they showed that Tf increases sharply with increasing E, but only to a point (Figure 2A). Using classical heterogeneous nucleation theory, they found that this behavior is due to saturation in the contact angle of the ice embryo with the underlying substrate, a saturation that occurs at −3.5 °C after a critical value E of 6 × 106 V/m is reached. They also showed that it is possible to overcome this freezing saturation by controlling the uniformity of the electric field with carbon nanotubes (Figure 2B), in which they achieved a Tf of −0.6 °C using a larger field strength of E of 3 × 107 V/m. In this case, a solid−liquid interface alters the mechanism due to changes in electric fields that are not uniform which they hypothesize leads to an enhancement of ice nucleation (Figure 2C).

While bulk water freezes at low temperatures, depending on pressure, into different ice phases, water at interfaces can exhibit a far richer phase diagram and phase coexistence depending on experimental parameters. Real-world experiments prepare and characterize nanoscale encapsulated water by employing interfaces that are inherently flexible and using irreversible kinetically controlled processes such as capillary compression or epoxy encapsulation so that the water content in the system is fixed. Most computational studies have focused on water confined within carbon nanotubes or between graphene sheets, in which the models reflect hydrophobic confining surfaces that are...

Figure 2. Temperature−time curve showing the onset of freezing. (A) The temperature of water was decreased linearly at the rate of 2 K min−1 until there was by a sudden temperature jump to 0 °C due to the release of latent heat. The inset shows pictures of the sample in liquid and solid phases during the experiment. The scale bar is 2 mm. (B) A scanning electron microscopy picture shows the carbon nanotube bundles on the alumina substrate, circled by a dashed line. The scale bar is 30 μm. (C) Also shown is a microscale mechanism on the surface of a carbon nanotube inserted in water. The electric field strength is not uniform at the solid−liquid interface, which may cause microflow near the surface and facilitate ice nucleation. Adapted with permission from ref 19. Copyright 2020 American Chemical Society.
almost always rigid, and the processes are assumed to be at equilibrium.

By lifting these two idealized conditions, the Head-Gordon lab showed that the nanoconfinement at fixed water densities was dramatically different for graphene that was treated as a rigid interface as compared to that of a highly flexible two-dimensional carbon allotrope. While the rigid slit nanocapillaries exhibit sharp differences in the confinement distances at the transition between an \((n)\)-layer and an \((n + 1)\)-layer water state, the flexible graphene walls allow the nanoconfined water to spatially split into an \((n)\)-layer and an \((n + 1)\)-layer coexisting state (Figure 3). Furthermore, the flexible walls introduce a very different sequence of ice phases and phase coexistence with vapor and liquid phases than that observed with rigid walls, with evident boundaries between square ice (as seen experimentally), vapor, and liquid states. The flexible confining sheets also induce small patches of hexagonal ice that nucleate and dissipate on the time scales of the simulation, suggesting that water is close to a solid–liquid phase transition. This leads to many new questions about crystallization kinetics and how one can control aqueous-based nucleation events at interfacial boundaries and the use of confinement.

Mandadapu et al. have been focusing on the dynamics of supercooled liquids and solids, particularly formation of polycrystalline solids, which consists of single crystals oriented in different directions and solid–solid interfaces called grain boundaries. To this end, most materials, including water, capable of forming polycrystals can also form glasses or amorphous solids. Whether a polycrystal or glassy structure is formed depends on the protocol or method by which the material is prepared such as the cooling rate or the annealing schedule. The resulting microstructure is a complex process involving competition between crystallization and vitrification, where the size of the crystals and resulting dynamics of the grain boundaries in polycrystals depend on the aforementioned formation protocols. Slower cooling rates produce larger grains while higher cooling rates produce smaller grains and glass formation.

To study the competition between crystallization and vitrification, Mandadapu and co-workers proposed a simple but physically rich coarse-grained lattice model, called the Arrow–Potts model that entails the essential physics of both glass-forming liquids and polycrystals. Specifically, the model supports a phase transition between a liquid phase with glassy dynamics and a crystal phase with multiple orientations. As the name suggests, the Arrow–Potts model combines the Arrow model, which is a kinetically constrained model used in prior studies of glassy dynamics, and the Potts model, a generalization of the Ising model. Using Monte Carlo simulations, the model reproduces the nonmonotonic behaviors on time scales of crystallization as a function of temperature typically encoded in the time–temperature transformation (TTT) diagrams. The model shows the emergence of two classes of polycrystalline structures limited by crystal nucleation or growth. At high temperatures, crystallization is dominated by the nucleation of compact and coarsening proceeds through hierarchical relaxation pathways within the supercooled liquid, resulting in fractal and ramified crystals that contain a large amount of interfaces.

To explain and unify these two regimes, Mandadapu et al. employed the framework of Kolmogorov–Johnson–Mehl–Avrami theory to combine the field theory of nucleation, and a random walk theory for crystal coarsening kinetics, both of which govern the high- and low- regimes, respectively. Figure 4 shows the nonmonotonic behaviors of the crystallization time scales as a function of temperature when quenched or cooled from the liquid state to the crystalline state, along with the characteristic polycrystalline microstructures at high and low temperatures. As previously mentioned, grain boundary interfaces result from the mismatch of orientation of two single crystal grains that meet during a cooling protocol. The grain
is the average shown. Adapted from ref 21.

boundaries contribute an unfavorable energy to the system, and there is a strong thermodynamic driving force to eliminate them. Furthermore, the evolution of the microstructure in crystalline systems toward its equilibrium structure proceeds through the slow migration of grain boundaries. The structural and dynamical aspects of the grain boundaries in this process are therefore important for understanding how polycrystalline materials evolve over time. It is well-known that boundaries through the small angle mismatched grains are known to consist of stacked dislocations with some order.71,72

However, the physical state and the dynamics of the high-angle grain boundaries are not completely understood. It is possible that, just as the bulk liquid exists in liquid crystal and supercooled states, the grain boundaries may have a life of their own and exist in different states such as liquid, crystalline-like, and disordered glassy states. These aspects provide interesting avenues for future studies of complex systems such as polycrystalline ice nucleation and growth, which has multiple crystalline polymorphs, both in the bulk and under confinement.

4. CHEMICAL REACTIVITY AT INTERFACES

Chemical transformations in realistic systems rarely occur in a single phase and often involve transport and reactions across interfaces. Ahmed, Head-Gordon, Limmer, and Wilson have examined the chemical coupling of bulk and interfacial processes to examine how reaction mechanisms and rates are modified in systems that have simple interfaces and dimensions that are small compared to the characteristic reaction diffusion lengths.

The Diels–Alder (DA) and retro-DA reactions in bulk water are well-understood and of great importance in organic chemistry. The Head-Gordon group recently studied the retro-Diels–Alder (retro-DA) reaction between cyclopentadiene (CPD) and methyl-vinyl-ketone (MVK) in bulk water at both low and high densities, as well as under two different manifestations of nanoconfinement (Figure 5).29 They simulated the entire system at the same level of QM theory, including the water molecules, with ab initio molecular dynamics (AIMD) based on density functional theory (DFT) using one of the best semilocal functionals available, B97M-rV,74 which they have shown previously reproduces structural and dynamical properties of water at a level of accuracy on par with hybrid functionals.74 To characterize the CPD-MVK retro-DA reaction in the different solvent environments, they mapped the free energy surface (FES) along two collective variables in the gas phase, in bulk water at two different densities, and in two different nanoconfined environments, using well-tempered metadynamics.

They found that low densities or nanoconfinement does not change the retro-DA activation free energies; these are comparable to those observed in bulk water within $k_B T$. These results are remarkable in that extremely different solvation environments lead to nearly identical acceleration rates. This is because the mechanism, which is the stabilization of the transition state through a highly local water hydrogen bonding to promote the elongation of the first bond-breaking event, remains the same for all environments, including that under strong nanoconfinement. For this reason, the implications for on-water and microdroplet chemistry for accelerated Diels–Alder reactions must be a complete mechanistic change and possibly reactants that are altered so completely as to destroy the original reaction altogether.

This general conclusion from the Head-Gordon group that confinement effects on reactivity can be small for Diels–Alder reactions also resonates with the recent work of the Wilson group, who independently explored recent reports of dramatic reaction acceleration (up to $10^6$ times) in electrospray droplets.

There is emerging evidence that chemical reactions in droplets, films, and emulsions are accelerated by many orders of magnitude over the same reactions measured in macroscopic “beakers”.12,28,75−79 The molecular origin of these enhanced rates is not always known and in some cases may originate from the multiphase processes that occur in the electrospray droplets themselves.30 These include solvent evaporation, which enhances bulk reaction rates as well as the possibility of interfering gas phase reactions, both of which confound observing in-droplet chemistry. Nevertheless, the droplet or aerosol interfaces play a substantial role in chemical transformations due to the much larger fraction of interfacial molecules with reduced solvent coordination (vs bulk), and/or the charge state and surface acidity in micron-sized compartments.76

The reaction-diffusive length $L_{\text{rxn}} = \sqrt{D \tau}$ is the average distance a molecule travels prior to reaction. $L_{\text{rxn}}$ depends upon the diffusion constant ($D$) and $\tau$, the chemical lifetime (i.e., concentration or bimolecular rate constant). For highly reactive species, such as hydroxyl radicals in an organic liquid, $L_{\text{rxn}}$ is only a couple of nanometers.30 Less reactive molecules can also exhibit short $L_{\text{rxn}}$ in semisolid or glassy environments due to diffusive confinement.35 It is our hypothesis that, for reaction

![Figure 4. TTT diagram showing crystallization times for a chosen crystal fraction of 25% as a function of temperature for two idealized protocols. (i) Quenching a liquid state from above the melting temperature to a particular temperature $T$ (red circles) and (ii) crystallization of an equilibrated supercooled liquid (SL) at a particular temperature (green circles). The dashed vertical line indicates the melting temperature. The TTT diagram from the model exhibits nonmonotonic behaviors typical of many experimental TTD diagrams on different materials. Furthermore, the two protocols show significant deviations in crystallization time scales when crystals are grown at lower temperatures. The microstructures with compact crystalline structures at high temperatures and fractal structures at low temperatures are also shown. Adapted from ref 21.](https://doi.org/10.1021/acs.jpcb.1c03756)
containers that have substantial surface area and dimensions that are on the order of $L_{rxn}$, reactants increasingly sample the container walls (i.e., interfaces), where surface orientation and reduced solvent coordination can substantially alter transition states, rates, and mechanisms from those observed in the bulk liquid.

For example, Wilson and co-workers studied the heterogeneous reaction of hydroxyl radicals (OH) with aqueous citric acid nanodroplets.82,83 The diffusion constant of citric acid in aqueous solutions can be changed over many orders of magnitude by changing the relative humidity ($R_H$) around the droplet. This allows insight into how diffusive confinement (i.e., $L_{rxn}$) at the surfaces of glasses and semisolids (vs liquids) alters the fundamental coupling of acid–base and free radical reaction pathways (Figure 6). They found, using surface sensitive mass spectrometry,84,85 that the reaction depth was a sensitive function of the diffusion constant of citric acid in ∼100-nm-sized particles. At small diffusion coefficients, that reaction proceeded within 4−6 nm of a gas−particle surface, in contrast to larger diffusion coefficients where the entire particle contents were available for reaction. By simply changing the water content in the particle (by changing relative humidity), they were able to precisely change where the reaction occurs to explore more general aspects of how reaction pathways and intermediates depend upon nanoscale diffusively confined environments.

For aerosol particles that exist in highly viscous, diffusion-limited states, steep chemical gradients are expected to form during photochemical aging in the atmosphere. Under these conditions, species at the aerosol surface are confined and become more rapidly transformed than molecules that are shielded and reside in the particle interior. To examine the formation and evolution of chemical gradients at aerosol interfaces, Wilson and Ahmed considered the heterogeneous reaction of hydroxyl radicals (OH) on ∼200 nm particles of pure squalane (a branched, liquid hydrocarbon) and octacosane (a linear, solid hydrocarbon), and binary mixtures of the two to understand how diffusion limitations and phase separation impact particle reactivity.86 Aerosol mass spectrometry is used to measure the effective heterogeneous OH uptake coefficient and oxidation kinetics in the bulk, which are compared with the elemental composition of the surface obtained using X-ray photoemission. The differences in surface oxidation rates are analyzed using a previously published reaction diffusion model, which suggests that a 1−2 nm highly oxidized crust forms on octacosane particles, whereas in pure squalane the reaction products are homogeneously mixed within the aerosol. This

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**Figure 5.** Retro-Diels−Alder in bulk vs near confined interfaces. (A) Collective variables $CV_1$ (green) and $CV_2$ (yellow) are the reactive bonds between diene and dienophile. (B, C) Free energy landscape, $\Delta G$, from AIMD using B97M-rV projected in the CV space and the minimum energy path between the cycloadduct reactant and dissociated diene-dienophile products. (D−G) The different simulated solvent environments: bulk mass density $\rho = 1$ g/cm$^3$ and $\rho = 0.66$ g/cm$^3$ (D, E) and in water nanoconfined between bent (F) and flat (G) graphene sheets. Reprinted with permission from ref 29. Copyright 2020 American Chemical Society.

**Figure 6.** Measurements for the heterogeneous reaction of OH with citric acid nanoparticles. Reaction depth ($L$) vs citric acid diffusion coefficient. Inset graphic showing $L$ relative to the nanoparticle size.
work illustrates how diffusion limitations can form particles with highly oxidized surfaces even at relatively low oxidant exposures, which is in turn expected to influence their microphysics in the atmosphere.

The Wilson group has also explored recent reports on dramatic acceleration (up to $10^6$ times) in droplets. For the reported phosphorylation reaction of sugars and the formation of ribonucleosides in droplets, careful control experiments have revealed in all likelihood that the observed reaction products and acceleration factors assigned to in-droplet chemistry have significant contributions from gas phase reactions. Using a newly developed quadrupole electrodynamic trap, they have further investigated confinement effects on a synthesis reaction using a fluorescent product. This approach allows precise control over droplet size ($10–80 \mu m$ in diameter) and avoids some of the ambiguities of electrospray mass spectrometry. The rate coefficient for the reaction of o-phthalaldehyde (OPA) with alanine in the presence of diethanolamine was found to be 25% larger than the equivalent reaction in the bulk, suggesting moderate effects of confinement on the bimolecular reaction.

Using stochastic reaction diffusion simulations, Wilson and coworkers have also examined accelerated imine synthesis in microemulsions reported by Fallah-Arraghi et al. Imine formation from the reaction of an aldehyde and an amine is unfavorable in aqueous solutions. However, Fallah-Arraghi et al. observed that imine synthesis was enhanced in micron-sized oil/water emulsions, with a 29-fold increase in the apparent rate of synthesis. They found that the reaction rate was accelerated in small compartments under dilute reactant concentrations (i.e., $[\text{aldehyde}]$ and $[\text{amine}]$). For example, the rate coefficient for imine synthesis became most pronounced in small compartments and dilute concentrations of reactions that could dominate the chemistry. This is illustrated in Figure 7 where deviations from $K_{eq}^B$ for imine synthesis become most pronounced in small compartments under dilute reactant concentrations (i.e., small compartments and dilute concentrations of reactants) can dominate the chemistry.

A key feature illustrated in eq 1 is that it is the fraction of molecules ($f_i$) at the interface that plays a large role in governing reactivity in microdroplets. For macroscale systems, $f_i$ is small so $K_{eq}^B \approx K_{eq}^o$. However, in micron-sized droplets or compartments, $f_i$ becomes sizable and in some cases (i.e., small compartments and dilute concentrations of reactions) can dominate the chemistry. This is illustrated in Figure 7 where deviations from $K_{eq}^o$ for imine synthesis become most pronounced in small compartments under dilute reactant concentrations (i.e., [aldehyde] and [amine]). For example, $K_{eq}^o$ is predicted to be nearly 5000X larger in a dilute, $r = 8.4 \mu m$ emulsion, compared to a macroscale reaction vessel.

The Limmer group has married path sampling tools with advanced force-field representations to describe simple chemistries in complex environments relevant to electrochemistry and atmospheric chemistry. With the isomorphism between rate constants and path ensemble partition functions leveraged, an algorithm to efficiently evaluate the spatial dependence of a rate constant, $\text{TPS+U}$, has been developed. Standard umbrella sampling is added on top of transition path sampling to bias the likelihood of a reactive event occurring at particular locations in an inhomogeneous system, such as near to or away from an extended interface. This algorithm has been used to trace out the dramatic slowing down of ion pair dissociation in the proximity of a platinum electrode. The algorithmic advances clarified that the nearly 2 orders of magnitude decrease in the rate was due to slow solvation fluctuations of constrained water molecules strongly confined to the first adlayer of the electrode. The dependence of the rate of ion pair dissociation through the premelting layer of ice has also been evaluated. As with the electrode interface, highly ordered water molecules at the liquid–solid interface were found to slow down dissociation, with a significant contribution to the rate afforded from the reduction of the effective diffusion controlling the reactive flux.

Galib and Limmer have studied the heterogeneous hydrolysis of $\text{N}_2\text{O}_5$, the primary sink of NOx compounds in the atmosphere, on the surface of aqueous aerosol (Figure 8). Such a study would be traditionally beyond the reach of standard computational tools due to the large time and length scales required to simulate interfacial chemical reactions with a realistic description of the environment. However, by employing an artificial neural network-based force field trained with ab initio reference data and employing importance sampling methods, they were able to study the steps whereby a gaseous $\text{N}_2\text{O}_5$ molecule is reactively taken up in an aqueous aerosol. Contrary to traditional wisdom, and the prevailing assumptions of a generation of atmospheric chemists, they found this process was dominated by interfacial rather than bulk hydrolysis due to the fast kinetics and slow solvation. Beyond describing the thermodynamics and molecular dynamics that lead to reactive uptake, they proposed and validated a simple model of the reactive uptake that can serve as a foundation for future field studies and rationalize a broad range of existing observations.

5. WHERE THEORY AND EXPERIMENT MEET AT CATALYTIC INTERFACES

Synchrotron-based soft X-ray spectroscopies, namely, soft X-ray absorption spectroscopy (XAS) and X-ray emission spectroscopy (XES), resonant inelastic soft X-ray scattering (RIXS), and ambient-pressure X-ray photoelectron spectroscopy (APXPS) with soft (<2 keV) and tender X-rays (2–6 keV) are necessary...
reveal that carbonate, oxalate, C−H/C−C, and LiO2 evolve at the surface, whereas LiO2 lies between Li° and the outermost carbonaceous component. Due to the peak evolution over time and increasing pressure, the study clarifies the reaction pathway in which oxalate will be formed and react as a reaction intermediate to create carbonate and CO. With the continued CO evolution, this pathway is self-recycling and can be proven by adding O2 to the CO2 gas environment. By adding O2 to the gas composition, the surface is completely oxidized, and no oxalate intermediate formation is evolving. Thus, the reaction pathway is bypassed, and carbonate is dominating the sample surface.

The Crumlin lab has shown that the utilization of APXPS revealed how CO2 interacts with a copper vs silver metal surface and how the water changed the adsorbates’ surface bonding properties. On the Cu(111) surface, the initial physisorption of linear CO2 and its transformation to a chemisorbed bent CO2 when water was introduced is a transition that occurs on the time scale of minutes to reach equilibrium. When both water and CO2 are subsequently removed, they observed the stable bent CO2 remaining on the surface confirming its strong bond to the surface. However, one of the challenges for characterizing metastable states, intermediates, and chemical reactions previously unobserved is the identification of what is measured using APXPS and other surface sensitive techniques. To address this challenge, a density functional theory (DFT) prediction approach that simulates adsorbate configurations and their respective free energies helps interpret experimental observables such as the measured binding energies. This process is depicted in Figure 10, showing how CO2 interacts this time with a silver surface and its transformation when exposed to water both through QM predictions and their agreement with experimentally collected APXPS data.

However, DFT at the lower rungs of Jacob’s ladder has yielded mixed success for modeling metal surfaces that are important to heterogeneous catalysis. Just recently, a benchmark study for metal surface relaxations and adsorption energies and site preferences for CO on the M(111) where M = Pt, Cu, Ag, Au metal surfaces found that while RPBE was performed, more recent meta-GGA DFT functionals such as SCAN102,103 and B97M-rV73,104 exhibited under-relaxed surfaces and strong overbinding of CO and did not predict the atop site for all metals observed at low CO adsorbate coverage. However, rarely are DFT functionals evaluated at the finite temperature conditions that are manifest in all catalytic experiments. To address this important aspect of electrocatalysis modeling, the Head-Gordon lab used AIMD in the NVT and NPT ensembles to better and more directly simulate the natural statistical fluctuations of an electrocatalytic surface under ambient conditions. They found that the good agreement for RPBE at 0 K worsened with experiment at finite temperature, whereas the B97M-rV73,104 meta-GGA functional better reproduced experimental bulk and surface relaxation properties. In turn, the CO adsorption energies and site preferences on the M(111) surfaces (where M = Pt, Cu, Ag, Au) using the meta-GGA functional also improved, predicting the atop site preference and correct trend for the remaining multicoordinated sites. Thermalization exposes the importance of interaction interactions, for which the B97M-rV functionals accurately predicts the weak binding energetics for the Ag(111) and Au(111) metals; these were found to be a mixture of chemisorbed and physisorbed CO species. Finally, because the B97M-rV73,104 functional also provides an excellent description of bulk
6. DISCUSSION AND FUTURE DIRECTIONS

Some future directions for interfacial chemistry include increasing the level of integration of spectroscopic tools with scattering methods to be able to decipher the complex behavior on interfaces in order to deepen connections to theory. The IR and THz region of the water spectrum is highly affected by the hydrogen-bonding environment and is thus an excellent probe for the effect of electric field on its properties and will be applied to problems of ice nucleation under confinement and for electric fields on chemical reactivity as shown by the Head-Gordon lab. Attenuated total reflection (ATR), which allows direct insight into the aqueous interface and the double layer on electrocatalytic surfaces, could also be readily applied and coupled into the experimental platform. The beamlines at the Advanced Light Source (ALS), which is over 25 years old, will benefit from a new initiative to upgrade the ALS to a diffraction-limited storage ring (DLSR) that has full transverse coherence in its light output. The new capabilities will allow us to bridge the fundamental electronic, structural, and chemical dynamics of collective phenomena of extended systems in the condensed phase. Emerging exceptionally powerful X-ray spectroscopy methods such as APXPS and RIXS with a subnatural line width, as demonstrated recently to elucidate the structure of water, will enable the acceleration for future investigations to understand absorbate and solvent interactions on arbitrary metal surfaces and/or provide greater time- and space-resolution for understanding interfacial reactions. However, figuring out a seamless way of coupling theory and experiment to elucidate mechanisms poses a major challenge. Crumlin and co-workers have proposed creating a “Digital Twin”, the direct computational simulation of a device or system, which would eventually augment decision making and execution of optimal experimental strategies to drive physical knowledge acquisition for reactive systems of interests. This poses a huge challenge to a theoretical framework that is robust enough to “reply” to continuous updates from real experiments, and that predictive capabilities are strong enough to effect correct experimental adjustments. We envision that electronic structure calculations, and statistical mechanical sampling to capture grand canonical statistics and reaction path finding, can be coupled with work flows that can create massively parallel physical simulations that leverage leadership class computational resources. One could imagine that the simulation data will be further elaborated with data analytics or simulated with deep learning in order to construct massive chemical reaction networks etc. that will yield fundamental mechanistic understanding about catalytic and electrochemical reaction cascades to connect to experiments on a different water, it reveals its potential feasibility for better theoretical predictions at the solid–liquid interface in the future.

![Figure 10. QM predictions and experimental observations of Ag surface with CO₂ adsorption alone and with H₂O at 298 K. (a) Predicted structures for O=CO₂− on Ag surface. (b) C 1s APXPS spectra for Ag surfaces in the presence of 0.3 Torr CO₂ at 298 K. (c) b-CO₂ becomes stabilized by a pair of H₂O₄ species forming hydrogen bonds with ΔG of −0.18 eV with respect to desorbing H₂O and CO₂. (d, e) Adsorbed O=CO₂− stabilizes one or two H₂O₄ species via hydrogen bonds and two more waters with hydrogen bonds to the O₂⁻. O=CO₂− stabilizes the first, second, third, and fourth H₂O on this site with ΔG of −0.41, −0.45, −0.37, and −0.19 eV, respectively. (f) C 1s APXPS spectra and the theoretical peak deconvolution results for Ag surfaces in the presence of 0.3 Torr CO₂ and 0.15 Torr H₂O at 298 K. Figure reproduced from ref 46.](https://doi.org/10.1021/acs.jpcb.1c03756)
range of materials and solutions, and across experimental conditions.

Many of the efforts discussed above occur near enough to equilibrium that standard concepts like free energies and transition states can be used to rationalize observations. An exciting future direction concerns the development of formalisms and numerical and experimental tools to probe molecular properties and chemical transformations in instances far from equilibrium. When reactions are driven through photoexcitation and large energies must be dissipated in complex environments, nonequilibrium and nonadiabatic effects must be described on the same footing. Advances in the molecular theory of open quantum systems can provide guidance for what quantities encode molecular dynamics most transparently. Further, far from equilibrium conditions occur ubiquitously in confined systems where driving forces and relaxation times can be large. Advances in driven confined liquids would benefit disparate fields like separations, energy storage and generation, and atmospheric chemistry. Generalizations of transition state theory valid away from equilibrium, and notions of variational stability, recently developed can aid in guiding experimental design.

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Notes

The authors declare no competing financial interest.

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