Cobalt-Porphyrin Catalyzed Electrochemical Reduction of Carbon Dioxide in Water II: Mechanism from First Principles

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Abstract

We apply first principles computational techniques to analyze the two-electron, multi-step, electrochemical reduction of CO₂ to CO in water using cobalt porphyrin as a catalyst. Density Functional Theory calculations with hybrid functionals and dielectric continuum solvation are used to determine the steps at which electrons are added. This information is corroborated with ab initio molecular dynamics simulations in an explicit aqueous environment which reveal the critical role of water in stabilizing a key intermediate formed by CO₂ bound to cobalt. Using potential of mean force calculations, the intermediate is found to spontaneously accept a proton to form a carboxylate acid group at pH<9.0, and the subsequent cleavage of a C-OH bond to form CO is exothermic and associated with a small free energy barrier. These predictions suggest that the proposed reaction mechanism is viable if electron transfer to the catalyst is sufficiently fast. The variation in cobalt ion charge and spin states during bond breaking, DFT+U treatment of cobalt 3d orbitals, and the need for computing electrochemical potentials are emphasized.
I. INTRODUCTION

CO$_2$ capture from flue gas and its conversion to useful products, including fuel molecules, has emerged as an important paradigm for a carbon-neutral economy. At discussed in the preceding paper of this series (henceforth “paper I”), high (∼70%) yield of carbon monoxide (CO) has been demonstrated in cobalt macrocycle-catalyzed electrochemical reduction of carbon dioxide (CO$_2$) in water at applied voltage of about -1.0 volt. The mechanisms of CO$_2$ reduction in non-aqueous solvents, for which much more negative potentials are needed, have been examined using a variety of methods.

Co(I)P-catalyzed CO$_2$ reduction in water and protic solvents, which requires a much less negative voltage for the onset of reaction than in organic solvent has received less fundamental studies. The present theoretical work focuses on the mechanism of this electrochemical reaction in aqueous media. As discussed in the preceding paper in this series (henceforth “Paper I”) which examines the structures, energetics, and charge states of reaction intermediates in detail, the reaction likely takes place in the following logical sequence of steps:

\[
\begin{align*}
\text{CoP} + \text{CO}_2 & \rightarrow \text{CoPCO}_2; \\
\text{CoPCO}_2 + \text{H}^+ & \rightarrow \text{CoPCOOH}; \\
\text{CoPCOOH} & \rightarrow \text{CoPCO} + \text{OH}^-; \\
\text{CoPCO} & \rightarrow \text{CoP} + \text{CO}.
\end{align*}
\]

“CoP” will henceforth denote cobalt porphine, the simplest porphyrin species, adopted in this work for ease of calculations. Key questions to be addressed include: (1) why the reaction, which involves protonation to form carboxylate acid motifs (COOH) with pK$_a$ typically on the order of 4.5, readily proceeds despite the fact that pH > 7 in experiments; (2) whether all steps are thermodynamically downhill; (3) whether the free energy barriers of the intermediate steps are low enough to be consistent with the observed reaction rate; and (4) at what stages the two electrons are added.

In Eqs. above, we have intentionally left out the charge states of the intermediates as yet unassigned in experiments. Since the two electrons can be added at any step(s), numerous mechanisms are consistent with these equations. The sequence of electron injection is governed by the redox potentials ($\Phi_{\text{redox}}$) of the pertinent reaction intermediates relative
to the applied voltage. The half-cell potentials of various charge states of cobalt porphyrins are known, but not those for the CO- and CO$_2$- bound complexes.

In this work, we use a combination of *ab initio* molecular dynamics (AIMD) techniques and Density Functional Theory (DFT) calculations with various exchange-correlation functionals and the polarizable continuum model (hereafter referred to as “DFT+pcm”) to study Eqs. [1][4]. These methods inform and support each other. The more economical DFT+pcm calculations approximate the aqueous environment as a dielectric continuum, allowing a global overview of the entire electrochemical reaction and a survey of the numerous reaction intermediates. Thus, DFT+pcm results reported in Paper I are used to extract $\Phi$$_{\text{redox}}$ for all possible intermediates. Redox potentials consist of hydration free energy ($\Delta$G$_{\text{hyd}}$) and ionization potential/electron affinity contributions. While $\Phi$$_{\text{redox}}$ and $\Delta$G$_{\text{hyd}}$ have also been calculated with the AIMD method and explicit treatment of the aqueous environment, these have so far been limited to monoatomic ions and molecules much smaller than porphyrins comprising a minimum of 37 atoms. Our B3LYP DFT+pcm redox potentials predict that CoPCOOH$^-$ is the key intermediate. To corroborate this $\Phi$$_{\text{redox}}$ conclusion, we also conduct more costly AIMD simulations with explicit water molecules. The hydration structures of the reaction intermediates predicted therein help explain the redox potential trends.

Next, we perform AIMD calculations on the individual reactions identified as the key steps in DFT+pcm calculations. Although it is possible that the electron addition steps of the two-electron reduction of CO$_2$ are rate-limiting, modeling the electron transfer rate via Marcus theory approach may not answer the most interesting scientific questions. This is because the electron injection rate almost certainly depends on engineering aspects such as the electrical contact between the gas diffusion electrode and the polymerized catalyst. Besides, it is necessary to demonstrate the viability of other steps in which electron transfer is not involved. We omit the electrode, focus on cobalt porphine molecule dispersed in water, and use AIMD to study two reactions involving [Co(II)PCOOH]$^-$: deprotonation (the reverse of Eq. [2]) and the cleavage of the C-OH bond (Eq. [3]). The protonation reaction proves critical to the efficient removal of one of the CO$_2$ oxygen atoms and reduction of the carbon atom from the +4 to the +2 formal charge state. The $p$K$_a$ of [Co(II)PCOOH]$^-$ will be determined using AIMD potential of mean force (PMF) in a manner previously applied to silanol groups on silica surfaces. As for the free energy change and barrier associated
with the breaking of the C-OH bond (Eq. 3) prior to releasing CO (Eq. 4), there is as yet no experimental reaction rate for direct comparison. The turnover rate per catalytic active site is obscured by the undetermined proportion of active CoP molecules actually participating in the reactions. But our PMF with an approximate reaction coordinate yields a barrier which suggests that CO gas evolution should proceed readily if electron transfer from the electrode is fast. It is known experimentally that Co(II)P binds weakly to CO;\textsuperscript{40,41} our gas phase calculations also shows that the Co(I)P-CO complex is weakly bound. Therefore once the C-OH bond is severed to form a OH\textsuperscript{−}, the subsequent steps (Eq. 4) should be exothermic, fast, and non-rate-determining, and they do not require further theoretical studies.

This work provides fundamental understanding specific to the CO\textsubscript{2} reduction mechanism in water using macromolecule catalysts, highlights the critical role played by the protic solvent water in lowering the voltage needed for the reaction, and may shed light on ways to further improve and modify the cobalt porphyrin catalyst. It is also of general interest to the fledging field of computational electrochemistry. The multistep nature of the reaction emphasizes the importance of accurate calculation of redox potentials.\textsuperscript{42–45} The protonation reaction (Eq. 2) is accompanied with a change in the Co ion charge state, which can be considered a form of coupled proton/electron process.\textsuperscript{42,46} We show that this can induce hysteresis in AIMD simulations. Finally, this study in explicit water, comprising AIMD trajectories exceeding 200 ps in duration, is greatly facilitated by the use of an empirical DFT+U method\textsuperscript{47} that can treat the localized 3\textit{d}-orbitals of the Co ion accurately without resorting to more costly theoretical methods such as hybrid functionals.

This paper is organized as follows: The theoretical methods used are discussed in Sec. 2. Section 3 describes the redox potential predictions which determine the electron addition steps, and then focuses on the deprotonation and C-O bond cleavage reactions of the key intermediate [Co(II)PCOOH]\textsuperscript{−}. Section 4 compares the method used in the present work to our previous CO\textsubscript{2}-related simulations;\textsuperscript{48} it further looks to the future and briefly discusses new computational techniques that may facilitate the modeling of demanding electrochemical processes. The accuracy of electronic structure calculations which underlie our mechanistic predictions will also be addressed.\textsuperscript{49} Section 5 concludes and summarizes the paper. An appendix discusses minor hysteresis issues encountered in one part of the calculations.
II. METHOD

All B3LYP or PBE plus dielectric continuum calculations apply the Gaussian suite of programs version g03,50 All DFT+U calculations use the VASP code.51,52 The Supporting Information document (SI) Sec. S1 provides a brief comparison between these packages.

DFT plus dielectric continuum (DFT+pcm) calculations apply the Becke-3-parameter-Lee-Yang-Parr (B3LYP)53,54 or the Perdew-Burke-Ernzerhof (PBE)55 functional, and the PCM dielectric continuum model.32 Other details are described in Paper I.29 Zero-point energy (ZPE) contributions to Eqs. 2 and 3 are estimated at T=0 K, also using Gaussian50 and the 6-31+G* basis set. The H2O-OH− complex is assumed to be the H+ accepting or OH− containing species in ZPE calculations.

The redox potential of the A(n+1)−/An− couple is the electron affinity (EA) of An− plus the difference in hydration free energies (∆Ghyd difference between A(n+1)− and An−). Both EA and ∆Ghyd are readily obtained from DFT+pcm total free energies that include zero point energy corrections and finite temperature contributions. Where available, DFT+U Φredox are estimated by substituting ground state DFT+U energies for B3LYP ones but retaining B3LYP dielectric hydration and ZPE information.71 All reported Φredox are referenced to the accepted value of 4.44 volt for the standard hydrogen electrode half-cell potential. The effect of the choice of DFT functionals on Φredox will be addressed in Sec. IV.

Spin-polarized AIMD simulations apply Vienna ab-initio simulation program (VASP),51,52 the PBE functional, Γ-point Brillouin zone sampling, 400 eV planewave energy cutoff, deuterium masses for all protons to allow Born-Oppenheimer dynamics time steps of 0.25 fs, a 10−6 eV energy convergence criterion, and T=425 K NVT conditions using a Nose thermostat. At T=400 K, the PBE functional yields average pure water structure consistent with experimentally observed water g(r) at T=300 K.56 Since the porphine ring exhibits significant ruffling fluctuations that explore a large configurational space, we raise the temperature by an extra 25 K to promote better sampling statistics. AIMD simulations apply 13.64 Å×13.64 Å×13.64 Å simulation cells which contain a CoPCOOH complex and 71 H2O molecules.57

Semi-local functionals such as PBE are generally inadequate for treating first row transition metal complexes like cobalt,58,59 although they are more successful with transition metal dimers.60 In general, transition metal complexes have been a challenge to DFT methods.49
We apply the following reaction to benchmark the prediction energetics:

\[ \text{[Co(III)P]}^+ + \text{CO} \rightarrow \text{[Co(III)P} - \text{CO}]^+. \] (5)

While Co(II)P itself seems well represented by DFT functionals,\textsuperscript{61,62} Paper I has shown that the widely used PBE and B3LYP functionals predict Eq. 5 binding energies which differ from the experimental value\textsuperscript{41} by about 0.5 eV in opposite directions. To deal with this problem, we augment the PBE functional with DFT+U\textsuperscript{47} applied to the partially occupied 3d-orbitals of the Co ion. With a judicious choice of the U parameter, this approach has been shown to give accurate predictions for organometallic compounds, although agreement between theory and experiments has not been universal.\textsuperscript{63–70} Using VASP PAW pseudopotentials,\textsuperscript{52} setting \( U=2.5 \) eV yields a binding energy for Eq. 5 that agrees with experimental data reported for the Co(III)TPP-CO complex.\textsuperscript{29,41,71} At \( T=0 \) K, these two complexes exhibit very similar VASP/PBE binding energies of 1.319 and 1.351 eV respectively; they differ by only 0.74 kcal/mol, suggesting that the experimental Co(III)TPP-CO binding free energy is a good metric for benchmarking the theoretical Co(III)P-CO predictions. As neither \( \text{CO}_2 \) nor CO strongly binds to cobalt porphyrins at most accessible Co charge states, Eq. 5 is the only binding constant available in the experimental literature as a benchmark. While this value of \( U \) may not be optimal for all Co charge states, referencing to Eq. 5 appears the most justifiable empirical route. The accuracy of AIMD simulations depend on the DFT+U method used, as will be discussed in Sec. IV.

As reported in Paper I,\textsuperscript{29} the optimal spin states of all gas phase B3LYP and PBE CoP complexes are low spin except \([\text{Co(I)PCOOH}]^{2-}\) and the unligated \([\text{Co(III)P}]^+\), both of which are triplets. DFT+U calculations at \( T=0 \) K yield the same optimal spin states.

Equation 2 involves calculating the \( pK_a \) of \([\text{Co(II)PCOOH}]^-\). \( pK_a = -\log_{10} \exp(-\beta \Delta G^{(0)}) \) has been successfully computed for molecules and surfaces in liquid water using the AIMD technique.\textsuperscript{39,72–75} Here \( \beta = 1/k_B T \), and \( \Delta G^{(0)} \) is the standard state deprotonation free energy,

\[
\Delta G^{(0)} = -k_B T \ln \left\{ C_0 \int_0^{R_{\text{cut}}} dR A(R) \exp[-\beta W(R)] \right\}. \tag{6}
\]

\( C_0 \) denotes 1.0 M concentration, \( R \) is the reaction coordinate, \( A(R) \) is a phase space factor to be discussed in Sec. III, \( R_{\text{cut}} \) is the cutoff distance delimiting the reaction and product valleys in the free energy landscape, and \( W(R) \) is the potential of mean force (PMF), referenced
such that \( W(R) = 0 \) as \( R \rightarrow \infty \). \( R_{\text{cut}} \) is taken as the onset of the plateau where \( W(R) \rightarrow 0 \).

The umbrella sampling technique\(^7\) and a 4-atom reaction coordinate

\[
R = R_1 - R_2 - R_3
\]

are applied to compute \( W(R) \). Here \( R_1 \) is the distance between the COOH acid proton and the oxygen atom on the designated proton-accepting H\(_2\)O, while \( R_2 \) and \( R_3 \) are the distances between this O atom and the two protons originally on the designated H\(_2\)O molecule, respectively.\(^39\) As \( R_2 \) and \( R_3 \) are about 1.0 Å for intact O-H covalent bonds, it can be readily inferred that \( R \sim -1.0 \) Å is consistent with deprotonation and CO\(^-\)/H\(_3\)O\(^+\) contact ion pair formation while \( R > -0.4 \) Å indicates an intact CO-H bond.\(^39\) Designating a special H\(_2\)O molecule can be done without loss of generality because all water molecules are interchangeable and only one is at any time is close enough to the acid proton to be considered a potential proton acceptor. Harmonic potentials of the form

\[
U(R) = B(R - R_o)^2,
\]

with \( B \) values between 2 to 4 eV/Å\(^2\) are applied to 7 umbrella windows with \( R_o \) spanning the range of \( R \) to be sampled. We reference the p\(K_a\) of [CoPCOOH]\(^-\) relative to the free energy of water-autoionization\(^23\) computed using the same reaction coordinate and elevated temperature and assumed to exhibit p\(K_w\)=14. As an validation test, our AIMD p\(K_a\) methodology has been applied to formic acid in water, yielding an acidity constant within a fraction of a pH unit of the experimental value (SI Sec. S2).

A “reflecting boundary condition” potential \( V(R_{\text{OH}}) \) sets an approximate 1.2 Å distance of closest approach between water protons and the hydroxyl oxygen atom. It preserves the identity of the deprotonated -COOH (or H\(_2\)O in the case of water auto-ionization) by preventing proton transfer via the Grotthuss mechanism. \( V(R_{\text{OH}}) = B(R_{\text{OH}} - R_1)^4 \), where \( B=200 \) eV/Å\(^4\) and \( R_1=1.3 \) Å, is imposed whenever \( R_{\text{OH}} < R_1 \). \( R_{\text{OH}} \) is the distance between the hydroxyl (COH) oxygen and all H\(^+\) other than the original COOH proton. Related boundary potentials have been applied to AIMD simulations of other chemical reactons.\(^48,76\) \( V(R_{\text{OH}}) \) is only necessary in the deprotonation umbrella sampling window with the most negative \( R \). As this leftmost umbrella sampling window exhibits a \( W(R) \) variation of only \( \sim 0.6 \) kcal/mol (see Sec. III), the effect of \( V(R_{\text{OH}}) \) should be small — much less than 0.6 kcal/mol.
FIG. 1: Snapshot of (a) [Co(I)P]−; (b) [Co(I)P]2− (see footnote 9) for description of charge states; 
(c) [Co(I)PCO2]−; (d) [Co(I)PCO2]2−; (e) Co(II)PCOOH; (f) [Co(II)PCOOH]−; in water. Panels 
(a)-(b) show the absence of hydrogen bonding between the negatively charged CoP and water. 
In (c)-(f), most H2O molecules are omitted; only those forming hydrogen bonds with the CoP 
structure are shown. Panels (e) and (f) depict the C-OH group in the trans (exo) and cis (endo) 
configuration, respectively. All snapshots are taken after 1-2 ps of short AIMD runs. Pink: Co; 
grey: C; red: O; white: H.

For the C-OH cleavage reaction (Eq. 3), the reaction coordinate is taken to be $R_{C-O}$, the 
distance between the COOH carbon and the hydroxyl oxygen atom. Harmonic potentials of 
the form Eq. 8 are applied to 10 umbrella sampling windows, but with $R_{C-O}$ replacing the 
4-atom coordinate $R$ in this case. The phase space factor $A(R)$ in the free energy expression 
analogous to Eq. 6 becomes $4\pi R_{C-O}^2$.

Dealing with slow, diffusive degrees of freedom is a significant challenge in AIMD calcula-
tions of bond-breaking free energies. During the stretching/cleavage of the C-OH bond,
rotational phase factors of $k_B T \log[4\pi(R_{C-O})^2]$ should naturally emerge in well-converged evaluations of Eq. 3. However, rotation of the nascent product $\text{OH}^-$ ion about the carbonyl carbon atom in water is too slow on AIMD time scales to accurately reproduce this rotational entropy. To give a better converged $W(R)$, the $x$- and $y$-coordinates of the carbonyl C and hydroxyl O atoms are kept identical and fixed while their $z$ coordinates are allowed to vary. No other atom is frozen in AIMD simulations. The rotational entropy is restored by multiplying $4\pi R_{C-O}^2$ to the $R_{C-O}$ probability distribution function at each $R_{C-O}$.\textsuperscript{48,76,77} The SI, Sec. S3, shows that this constraint has modest effect on the sampling of the Co-C-O angular distribution despite the bulkiness of the CoP group.

The deprotonation free energy change in Eq. 2 is a state function which should not depend on the reaction coordinate chosen provided that equilibrium sampling is achieved. Using the one-dimensional coordinate $R$, however, we observe some hysteresis due to the picosecond time scale relaxation of the charge state of the cobalt ion as the extent of deprotonation varies. For Eq. 3, we are interested in the free energy barrier in addition to the free energy change. Umbrella sampling yields a free energy barrier estimate which depends on and is generally underestimated by any chosen reaction coordinate because some trajectories with forward velocity can recross the “transition state” point and do not proceed to product formation.\textsuperscript{78,79} To assess the validity of the computed barrier, we perform transmission coefficient ($\kappa$) calculations,\textsuperscript{78,79} to be discussed in more detail below. The transition path sampling method\textsuperscript{80} is the rigorous approach to compute free energy barriers; although more costly, it will be considered in the future. Multi-dimensional metadynamics,\textsuperscript{81–83} new deprotonation coordinates,\textsuperscript{75} the self-consistent DFT+U method,\textsuperscript{68} and new DFT functionals,\textsuperscript{84} may also benefit future AIMD-based electrochemical calculations, and will be mentioned in Sec. IV.

The amount of water in the simulation cell is determined using grand canonical Monte Carlo simulations at constant water chemical potential, the Towhee Monte Carlo code,\textsuperscript{85} and the SPC/E water model.\textsuperscript{86} The cell size is identical to that used in AIMD simulations (13.64 Å$^3$). The temperature is set at $T=300$ K because the SPC/E model, unlike DFT/PBE, yields reasonable water structure at room temperature. One charge-neutral Co(III)PCOOH is placed frozen in its DFT-optimized configuration in the simulation cell. The CoP and COOH Lennard-Jones force field parameters are approximated with those of Mn(II)P\textsuperscript{87} and the formate anion, respectively. The atomic partial charges are assigned
TABLE I: Redox potential of various species using three functionals/computational methods, in volts. ZPE are computed using the B3LYP functional; asterisks indicate that dielectric solvation contributions also come from B3LYP. Convergence cannot be achieved when applying DFT+U to some doubly negatively charged systems in the gas phase. The electron affinities of these species are listed in the SI, Sec. S4. DFT+U calculations are performed using VASP; B3LYP and PBE calculations apply the Gaussian suite of codes.

| Species              | Reduced   | Oxidized  | B3LYP | PBE  | DFT+U |
|----------------------|-----------|-----------|-------|------|-------|
| Co(I)P^−             | Co(II)P   | -1.71     | -0.89 | -1.46*|
| Co(I)P^2−            | Co(I)P^−  | -2.11     | -2.31 | NA   |
| Co(I)PCO_2^2−        | Co(I)PCO_2^− | -1.82  | -1.78*| -1.68*|
| Co(II)PCOOH^−        | Co(III)PCOOH | -1.17 | -2.14 | -1.07*|
| Co(II)PCOOH^2−       | Co(II)PCOOH^− | -2.17 | -2.08*| NA   |

using Mulliken charge analysis of a gas phase B3LYP/6-311+G(d,p) Gaussian calculation. 4×10^8 Monte Carlo moves are attempted, 30% of them being water insertion/deletions. The most probable number of water molecules in the simulation cell is determined to be 71, and this is the water content used in AIMD simulations. A net −|e| charge and a neutralizing background are then imposed on the final CoPCOOH/H_2O configuration from the Monte Carlo run, and AIMD simulations are initiated. We choose a few sampling windows along the reaction coordinates as seed windows. With the appropriate umbrella sampling potentials of the selected windows turned on, AIMD pre-equilibration is conducted for 2 ps at T=500 K. Maximally localized Wannier function analyses confirm that the extra electron resides on CoPCOOH. Then the system is further equilibrated at the target temperature T=425 K for another 2 ps before statistics are collected. The starting configurations for all other sampling windows are taken successively from snapshots of adjacent windows a few picosecond into their AIMD trajectories.

The statistical uncertainty in each sampling window is estimated by splitting the trajectory into four equal parts, calculating the standard deviation for W(R_2) − W(R_1), where R_1 and R_2 are the boundary values in the window, and then dividing by √4 to yield an approximate error bar for the entire trajectory. The overall uncertainty convolves the standard deviations in all windows. The statistics are generated with AIMD trajectories of at
least 10 ps in duration; in a few windows where the uncertainties are large, 15-20 ps AIMD runs are conducted.

The maximally localized Wannier function analysis is used to determine the charge/spin state of the cobalt ion in the reactants, productions, and reaction intermediates in liquid water. As discussed in the SI Sec. S1, the Mulliken charge decomposition technique, popular in the quantum chemistry community, has not been implemented in the planewave basis VASP code. Alternative charge decomposition methods such as simply integrating the charge/spin densities within some radius around the transition metal ion tend to yield ambiguous results. Hence the Wannier approach appears the most useful method in our condensed phase, periodically replicated simulation cell setting.

III. RESULTS

A. Redox Potentials, Hydration Structures, and Electronic Structures

The redox potential values are listed in Table I. To anticipate the conclusions, we find that the predicted B3LYP DFT+pcm absolute redox potentials are not in good agreement with experimental values. However, these potentials referenced to one redox couple yield qualitative results consistent with insights gained from AIMD simulations, and they allow us to assign the charge state of the key intermediate species.

Thus, at first glance, B3LYP appears to predict a Co(I)P/Co(II)P redox couple value which disagrees with the average experimental value of −0.67 volt (−0.5 to −0.84, depending on the porphyrin ring substituent and solvent). The redox potential also strongly depends on the DFT functional used, in contrast to the findings of Ref. 62 for the Co(II)P-NO/[Co(III)P-NO]+ couple. DFT+U and B3LYP redox potentials track each other while the PBE functional yields substantially different values. Systematic errors in hybrid DFT plus dielectric continuum redox potentials have been reported in the literature, particularly those associated with the B3LYP functional. While part of the discrepancy in Table I may be due to DFT inaccuracies, the DFT+U redox potential, fitted to Eq. 5 is also off by 0.8 volt. Thus uncertainties arising from the use of the dielectric approximation used to calculate \( \Delta G_{\text{hyd}} \) as well as the lack of ring substituents in our calculations may also be responsible. Although PBE appears to yield a \( \Phi_{\text{redox}} \) for the [Co(I)P]−/Co(II)P couple close
FIG. 2: Mechanism of CO₂ reduction with electron addition deduced from hybrid DFT plus dielectric continuum redox potential calculations. Red denotes key intermediates; green species should undergo fast reactions.

To the experimental value, it performs worse for the benchmark reaction Eq. 5 than DFT+U. The result for that equation is deemed more reliable because the pertinent experiments were performed in aprotic solvents which interact weakly with the reactants.

To proceed, we focus on relative B3LYP redox potential values and assume that all electrochemical measurements occur near the B3LYP [Co(I)P]⁻/Co(II)P voltage. As discussed in the Introduction, this corresponds to the experimental condition where the onset of CO gas evolution is within a few tenths of a volt of the Co(II)P reduction potential. The B3LYP redox potentials of other species relative to [Co(I)P]⁻/Co(II)P (−1.71 volt) then determine whether an additional electron has been incorporated at that step. A similar approach has been used in Ref. 43, where it has also been suggested that relative B3LYP redox potentials are much more reliable than absolute values. Thus the B3LYP functional predicts that Co(III)PCOOH is reduced to [Co(II)PCOOH]⁻, while [Co(I)P]⁻ and [Co(II)PCOOH]⁻ are not reduced to [Co(I)P]²⁻ and [Co(II)PCOOH]²⁻, respectively, because their required voltages are much more negative than −1.71 volt. The experimental reduction potentials for [Co(I)P]⁻ in aprotic solvents are indeed -0.51 to -1.2 volt more negative than that for Co(II)P. However, unlike [Co(I)P]⁻ itself, [Co(I)PCO₂]⁻ is already reduced to [Co(II)PCO₂]²⁻ at the [Co(I)P]⁻/Co(II)P voltage with a mere additional -0.11 volt. In other words, it is much easier to add an electron to [Co(I)PCO₂]⁻ in water than [Co(I)P]⁻.
These trends can be explained with DFT+U/AIMD simulations in explicit water. The snapshots in Fig. 1 are generated using short, 1-2 ps AIMD trajectories, which are sufficient to yield qualitative, well-equilibrated hydration structures if not highly precise average hydration numbers. Despite their net negative charges, both [Co(I)P]− and [Co(I)P]2− are effectively hydrophobic plates which do not form hydrogen bonds with water molecules. No water protons are observed within 2.5 Å, a typical hydrogen bonding cutoff distance, of the porphine N and Co atoms (Figs. 1a-b). This is in contrast to the charge neutral and positively charged Mn(II)P and Mn(III)P, where 1 or 2 H2O molecule strongly coordinates to the Mn site. CO2, a famously inert molecule, also fails to hydrogen-bond with water. However, when they combine to form [Co(I)PCO2]− and [Co(I)PCO2]2−, the resulting complexes form 4 to 5 hydrogen bonds with water through the partially negatively charged O atoms on the CO2, which now adopts a bent geometry like a carbonate or a carboxylate anion (Figs. 1c-d). The enhanced interaction with water evidently facilitates the accommodation of an extra electron. This finding can be significant not just for electrochemical reduction of CO2, but also for CO2 capture in general. The B3LYP DFT+pcm results reflect this CO2− polarization information despite the fact that water is treated implicitly there.

We also consider the hydration structures of Co(III)PCOOH and [Co(II)PCOOH]− (Figs. 1e-f). The C-OH trans (exo) and cis (endo) configurations are almost isoenerggetic in gas phase Co(III)PCOOH, within 1.06 kcal/mol of each other. In contrast, [CoP(II)PCOOH]− forms a strong intramolecular hydrogen bond between the COOH acid proton and one of the nitrogen atoms on the porphine ring when the proton is in the cis position (Fig. 1f). This COOH proton cannot form hydrogen bond with other water molecules. In the gas phase, this structure is 4.93 kcal/mol more stable than the trans configuration where the OH points outwards (Fig. 1e). This feature will play a prominent role in acid-base reactions (Eq. 2). Intramolecular hydrogen bonds have also been suggested to facilitate CO2 binding and chemical reduction in the literature.

The calculated redox potentials (Table I) suggest an overall mechanism shown in Fig. 2. The CO2 adsorption and the first electron insertion steps are likely simultaneous and cooperative. This is because the B3LYP functional predicts that the following equation,

\[ [\text{Co(I)}P]^2− + \text{CO}_2 \rightarrow [\text{CoP(I)}\text{PCO}_2]^2−, \]  

(9)
exhibits a very significant free energy gain of 27 kcal/mol in water (treated as a dielectric continuum).\textsuperscript{29} However, \([\text{Co(I)P}]^-\) should not be readily reduced to \([\text{Co(I)P}]^{2-}\) at or slightly below the \([\text{Co(I)P}]^-/\text{Co(II)P}\) half cell voltage, and \(\text{CO}_2\) is not strongly bound to \([\text{Co(I)P}]^-\). This suggests that the \(\text{CO}_2\) may be thought of as part of the solvent, and the electron transfer to the cobalt complex as a solvent/\(\text{CO}_2\) fluctuation-mediated process akin to the Marcus theory picture.\textsuperscript{38} As mentioned in the introduction, the electron transfer rate may depend on the electrical contact between the catalyst and the gas-diffusion electrode, and is not the main subject of this study. The protonation and C-OH cleavage steps in Fig. 2 are at the heart of the catalytic function, and must be fast and spontaneous; understanding of the scientific principles involved can lead to improved catalysts and reaction conditions. In the next subsections, we consider these steps in detail using the AIMD method. \(\text{Co(II)P}\) is known to be weakly bound to \(\text{CO}.\textsuperscript{40}\) Although experimental data is not available, B3LYP/6-31+G* calculations indicate that the gas phase \(\text{Co(I)P-CO}\) binding energy is only 1.76 kcal/mol at \(T=0\) K. Once C-OH cleavage is achieved, the rest of the reaction (Eq. 4) should proceed rapidly.

The Co charge and spin states predicted for B3LYP DFT+pcm optimal structures\textsuperscript{29} and in DFT+U/AIMD aqueous phase snapshots generally agree with each other. For example, we consider an AIMD snapshot of \([\text{Co(I)PCO}_2]^{2-}:\text{H}_3\text{O}^+\) contact ion pair\textsuperscript{39} in water (Fig. 3a, obtained using Eq. 8 with \(B=3\) eV/Å\(^2\) and \(R_o = -1.12\) Å).\textsuperscript{37} A maximally localized Wannier function analysis reveals that 6 occupied \(d\)-spin-orbitals are centered within 0.1 Å of the Co atom; another 2 occupied spin-orbitals are 0.86 to 0.88 Å away from the Co, and 1.04 and 1.02 Å away from the \(\text{CO}_2\) carbon atom, respectively. (The Co-C distance is 1.90 Å in this snapshot.) This electronic configuration is consistent with a dative covalent bond donated by \([\text{Co(I)P}]^{2-}\) to \(\text{CO}_2\). The singly occupied highest occupied HOMO state is delocalized on the porphyrin ring, as has been discussed in Paper I and is confirmed in the spin density plot of Fig. 3a. We designate this species \([\text{Co(I)PCO}_2]^{2-}\).

A key exception to the agreement between AIMD and B3LYP DFT+pcm calculations is \([\text{Co(II)PCOOH}]^-\). Figure 3b depicts an AIMD snapshot of this species, obtained with harmonic constraint \(R_o=-0.4\) Å which yields a COOH (i.e., protonated \(\text{CO}_2\)) group and also forces a \(\text{H}_2\text{O}\) molecule to accept a hydrogen bond from the COOH proton.\textsuperscript{39} A Wannier analysis of this snapshot (Fig. 3b) reveals 7 occupied \(d\)-spin-orbitals centered around 0.2 Å of the Co atom. Another two occupied spin-orbitals are localized along the Co-C bond,
FIG. 3: Spin densities as protonation of $[\text{Co(I)PCO}_2]^{2-}$ proceeds. Gold/blue: regions with net positive/negative spin densities. The isosurface spin density values are different for all panels to facilitate visualization, and are $1.32 \times 10^{-4}$, $2.88 \times 10^{-4}$, $2.32 \times 10^{-4}$, and $2.72 \times 10^{-4} |e|/\text{Å}^3$ for panels (a), (b), (e), and (f), respectively. The color scheme of the stick figures for water and CoP is as in Fig. 1. (a) Well-equilibrated $[\text{Co(I)PCO}_2]^{2-}$:$\text{H}_3\text{O}^+$ contact ion pair, obtained using a harmonic umbrella potential (Eq. 8) with $B=3.0 \text{ eV/Å}$, $R_o = -1.12 \text{ Å}$. The singly occupied molecular orbital (HOMO) state is delocalized on the porphyrin ring. (b) $[\text{Co(II)PCOOH}]^-$, $B=3.0 \text{ eV/Å}$, $R_o=-0.4 \text{ Å}$; the harmonic potential forces a $\text{H}_2\text{O}$ molecule to accept a hydrogen bond from the COOH proton. (c) & (d) reprise panels (a) and (b), respectively, omitting the spin densities to reveal the atomic positions more clearly. The dark blue stick figure above CoP in panel (c) represents the transient $\text{H}_3\text{O}^+$. (e) & (f): Starting the trajectory from panel (a), $R_o$ is suddenly switched from $-1.12 \text{ Å}$ to $-0.4 \text{ Å}$. The panels are taken 1.188875 ps and 1.189125 ps into this trajectory and bracket the transition.

within 0.6 and 0.8 Å of the C-atom respectively. These Wannier orbital centers are closer to C than Co, and the electronic configuration is consistent with a $[\text{COOH}]^-$ group attached to a $[\text{Co(II)P}]^0$. The HOMO state is a Co $d$-orbital; the spin density of this species (Fig. 3b) is far more localized than that of $[\text{Co(I)PCO}_2]^{2-}$ (Fig 3a). In contrast, B3LYP calculations with dielectric approximation for water suggest a $[\text{Co(I)P}]^{2-}$-COOH$^+$ electronic configuration, and the HOMO is a $\pi$-orbital on the porphyrin ring there. This difference most likely
reflects the explicit treatment of molecular water in the DFT+U/AIMD simulation which helps to stabilize a COOH$^-$ group via hydrogen bonding. In the SI, Sec. S5, explicit treatment of water molecules in the first hydration shell of the COOH group is indeed shown to yield a Co(II) charge state.

The change of electronic structure coupled to protonation of [Co(I)PCO$_2$]$^2^-$ has consequences for AIMD simulations. Figures 3e-f depict the spin density transition in real time, occurring about 1.2 ps after the harmonic umbrella sampling potential (Eq. 8) is suddenly switched from $R_o = -1.12$ Å (consistent with that in Fig. 3b) to $R_o = -0.4$ Å (Fig. 3b). As alluded to above, this change in $R_o$ leads to reprotonation of [Co(I)PCO$_2$]$^2^-$ from its H$_3$O$^+$ neighbor and the motion of a water molecule towards the CO$_2^-$ group. Simultaneously, the electronic structure relaxes to [Co(II)PCOOH]$^-$ due to the large driving force arising from the nuclear motion. However, if the driving force is not sufficient, e.g., if $R_o$ is switched to an intermediate $-0.7$ Å, the system may take much longer than 1.2 ps to spontaneously sample the Co(II) charge state. As discussed below, this leads to a slight hysteresis in the umbrella sampling calculation.

B. Protonation of [CoPCO$_2$]$^2^-$

Figure 4a depicts the $W(R)$ associated with the deprotonation reaction

$$[\text{Co(I)PCO}_2]^2^- + \text{H}^+ \rightarrow [\text{Co(II)PCOOH}]^-.$$  (10)

Our 4-atom coordinate $R$ effectively interpolates between the large negative $R$ deprotonated plateau region where $W(R) \rightarrow 0$, related to water-separated ion pairs,$^{39}$ and the protonated $R > -0.4$ Å region where the shallow curvature of $W(R)$ is governed by hydrogen bonding between the acid proton and a water molecule serving as a hydrogen bond acceptor. The distribution of shortest distance between a water oxygen and the COOH acid proton, obtained in an unconstrained (i.e., $B = 0$ in the Eq. 8) AIMD simulation of [Co(II)PCOOH]$^-$ in water, is expressed as a free energy profile ($W(r_{O-H})$) in the inset of Fig. 4a. The optimal $r_{O-H}$ is 3.2 Å. Recalling the definition Eq. 7 and the fact that $R_2$ and $R_3$ are O-H covalent bonds of lengths $\sim 1$ Å, this optimal value implies that a true minimum in $W(R)$ should not emerge until $R \sim 1.2$ Å. The optimal $r_{O-H}$ distance is larger than the canonical hydrogen bond cutoff distance of 2.5 Å, and reflects the inability of the COOH proton, engaged
in a strong intramolecular hydrogen bond to one of the CoP nitrogen atoms, to donate a hydrogen bond to water molecules (Fig. 1f). Indeed, along the AIMD trajectory, there is only a 2% probability that the COOH proton and any water oxygen atoms are within 2.5 Å of each other. Fortunately, the reaction coordinate $R$ and the umbrella constraining potentials enforce hydrogen bond donation from the acid proton to water, a prerequisite to deprotonation.

To estimate $pK_a$ via Eq. 10, we find the most probable $O_{water}$-$H^+$ hydrogen bond distance $r_{O-H}$ at each $R$, thus locally converting $W(R)$ to $\tilde{W}(r_{O-H})$, and perform a spline fit to that probability distribution. The result is matched to $\tilde{W}(r_{O-H})$ in the small $r_{O-H}$ region obtained in the aforementioned AIMD run where the umbrella potential is absent. See the Fig. 4a inset. Integrating over $r_{O-H}$ with a $4\pi r_{O-H}^2$ volume element, which takes the place of the phase space factor $A(R)$ in Eq. 6, referencing to water auto-ionization computed at a similar elevated temperature, and adding a $-0.57$ kcal/mol zero point energy correction estimated from gas phase B3LYP calculations, $pK_a=9.0\pm0.4$ is predicted. Thus, $[Co(II)PCOOH]$ does not behave like an ordinary carboxylate acid with $pK_a\sim4.5$. The significant reduction of acidity indicates that protonation of $[Co(II)CO_2]^{2-}$ is exothermic at the experimental pH $>7$ conditions.

In contrast, a preliminary study of the deprotonation $W(R)$ of Co(III)PCOOH reveals that $W(R)$ is almost independent of $R$ for $R<-0.4$ Å (not shown). Our reaction coordinate has been used to calculate $pK_a$ down to 3.8 where a finite curvature persists in $W(R)$. This suggests that the $pK_a$ of Co(II)PCOOH is less than 3.8, much lower than that of $[Co(II)PCOOH]^-$. Thus, adding an electron to the CO$_2$-ligated catalyst evidently enhances its ability to hold on to excess protons. While this may appear obvious in retrospect, the strong intramolecular hydrogen bonding in $[Co(II)COOH]^-$(Fig. 1f) also likely contributes to its higher $pK_a$. This preliminary $pK_a$ estimate for Co(III)PCOOH suggests that protonation of CoPCOO$^-$ cannot occur spontaneously at the experimental pH $\sim7$, further confirming the B3LYP DFT+pcm $\Phi_{\text{redox}}$ prediction that $[Co(II)PCOOH]^-$, not $[Co(III)PCOOH]$, is the key intermediate (Fig. 2).

Figure 4b shows that the hydration numbers $N_w$ do not significantly vary with $R$. However, the hydration structure may determine whether hysteresis in the cobalt charge state occurs as $R$ increases. The full curve in Fig. 4 collates contributions from all windows, such that windows 1 and 3 are initiated from $[Co(I)PCO_2]^{2-}$ configurations generated in a
FIG. 4: (a) Red: Potential of mean force ($W(R)$) associated with $[\text{Co(II)PCOOH}]^-$ deprotonation. The most negative values of $R$ are associated with complete deprotonation while $R > -0.4$ Å refers to the protonated state. Inset: the violet line depicts the $\bar{W}(r_{\text{O-H}})$, computed using unbiased AIMD simulations; the red line depicts $\bar{W}(r_{\text{O-H}})$ transformed from the biased (umbrella sampling) $W(R)$ and is a cubic spline fit. (b) Hydration numbers ($N_w$) of the carbonyl (green) and hydroxyl (blue) oxygen atoms on the COOH group. $N_w$ is defined as the number of water protons within 2.5 Å of these oxygen sites. Cobalt is in the Co(I) state in windows 1-3 and Co(II) state in 4-7, except for the dashed curves (window 3) where Co(II) prevails. The dashed lines shows that window 3 is affected by the cobalt charge state hysteresis (see text).
hysteresis in umbrella sampling simulations. Fortunately, only the intermediate $R_0 = -0.7 \, \text{Å}$ window 3 suffers from this problem. The maximal underestimation of $[\text{Co(II)PCOOH}]^- \text{pK}_a$ this can introduce is the difference between the full and dashed curves in window 3, which is 0.76 kcal/mol or 0.55 pH unit. This hysteresis issue is discussed in more detail in the appendix.

Using a purely dielectric continuum treatment of water and the B3LYP/6-31+G* method, Paper I predicts deprotonation $\text{pK}_a$ of 13.8 and 7.6 for $[\text{Co(II)PCOOH}]^-$ and Co(III)PCOOH, respectively. They are several pH units higher than AIMD estimates. The discrepancies are partly due to the lack of explicit water molecules in the B3LYP DFT+pcm calculations, but they also reflect the substantial ($\sim 5$ kcal/mol) variation in deprotonation free energies when using different DFT functionals (Table 4 in Paper I).

C. C-OH bond Cleavage Reaction

Finally, we apply umbrella sampling to study

$$[\text{Co(II)PCOOH}]^- \rightarrow [\text{Co(II)PCO}] + \text{OH}^-.$$  (11)

We use the C-O distance of the C-OH bond as the reaction coordinate. Figure 5a shows that this reaction is exothermic. Wannier analysis reveals that the system remains in the Co(II) state as $R_{C-O}$ varies, and no hysteresis is observed. After accounting for standard state correction, the $4\pi R_{C-O}^2$ rotational contribution, a $-2.5$ kcal/mol ZPE correction, and integrating $\exp[-\beta W(R_{C-O})]$ in the reactant channel as in Eq. 6 we obtain a free energy of reaction $\Delta G^{(0)} = -8.5\pm1.1$ kcal/mol. The barrier height is a low $\Delta G^{(0)*} = 5.2\pm0.6$ kcal/mol confirms that the activation free energy is fairly low. We have not attempted to compute ZPE for $\Delta G^{(0)*}$ which requires a projecton operation that removes the reaction coordinate; however, in a study of a simple C-OH bond breaking reaction ZPE has been found to be small, reducing $\Delta G^{(0)*}$ by 0.8 kcal/mol.

The predicted activation barrier may depend on the $R_{CO}$ coordinate chosen. As in Ref. 48, we have computed the transmission coefficient. Thus, in the umbrella sampling window containing the $W(R)$ turning point, we randomly choose 10 configurations at the top of the barrier, half with positive velocities $dR_{C-O}/dt$ and half with negative ones, restart AIMD trajectories without umbrella sampling potentials, and determine the ratio $\kappa$ that
FIG. 5: (a) $W(R)$ for the C-OH cleavage reaction, Eq. 11. Inset: $W(R)$ for CO$_3$H$^-$ → CO$_2$ + OH$^-$ from Ref. 48. Snapshots at points A-D are depicted in Fig. 6. (b) Hydration numbers ($N_w$) for the carbonyl (green) and hydroxyl (blue) oxygen atoms as $R_{C-O}$ varies.

The C-OH cleavage activation barrier is almost a factor of 3 smaller than that previously found for the uncatalyzed CO$_3$H$^-$ → CO$_2$ + OH$^-$ reaction (Fig. 5a inset). Even though the comparison is not perfect, in that the carbon atom is not reduced to its +2 oxidation state in the previous work, the cobalt porphyrin has clearly and drastically reduced the C-OH cleavage barrier.

As the C-OH cleavage reaction proceeds, the carbonyl oxygen in the initially partially negatively charged COOH$^-$ functional group becomes part of an uncharged carbon monoxide
molecule (CO) weakly bound to Co(II)P, and the oxygen atom of the nascent CO exhibits a hydration number $N_w$ which steadily decreases (Fig. 5b). In contrast, the hydroxyl oxygen transitions towards a hydroxide anion (OH$^-$), and its $N_w$ increases to about 3.5. The $N_w$ value for the emerging OH$^-$ oxygen in the present heterogeneous environment is therefore similar to that predicted in bulk liquid water using the PBE functional. Figure 6 further depicts snapshots of the instantaneous hydration structure of the COOH group at different values of the reaction coordinate. Panel (b) represents a configuration where the COOH proton is intramolecularly hydrogen bonded to a porphine ring N atom (Fig. 1f). In panel (a), which corresponds to a kink in Fig. 5a, the COOH proton has instantaneously been donated to a N atom, forming a covalent bond with it. In the gas phase, this N-H bonded structure is 6.68 kcal/mol higher in energy than that of Fig. 6b. Nevertheless, in the aqueous phase, this configuration is occasionally observed.

IV. DISCUSSION

A. Comparison with Previous CO$_2$ Theoretical Work

In this subsection, we make comparisons with some computational aspects of Ref. 48 and with Paper I. Unlike Ref. 48, we have not constrained the OH bond rotation around the C-O axis in the cleaved COOH group and then corrected for the entropic contributions there. This is because the PBE functional we use is consistent with much faster OH$^-$ dynamics in water than the RPBE functional previously applied, and it is reasonable to assume that the OH rotation around the C-O axis is better-sampled than in RPBE simulations within 10 ps AIMD trajectories. This should only affect $\Delta G^{(0)}$, not $\Delta G^{(0)*}$, because the C-OH bond is not completely broken at the transition state at $R_{C-O} = 1.9 \text{ Å}$ and free OH rotation around the C-O axis does not occur there. The important qualitative conclusion of this work is that $\Delta G^{(0)}$ is exothermic and that C-OH bond breaking is thermodynamically downhill; the precise free energy change associated with Eq. 11 is less important.

We have not attempted to correct the AIMD/DFT+U $W(R)$ with single point MP2 calculations as was done in Ref. 48. The CoP systems examined in this work are too large for large-basis MP2. Furthermore, in Ref. 48, the DFT functional used was RPBE which
FIG. 6: (a)-(d): snapshots in the four umbrella sampling windows indicated in Fig. 5. AIMD simulations are conducted in an explicit liquid water environment; only a few water molecules are shown in these snapshots. In panel (a), the proton on the COOH group has migrated to one of the nitrogen atoms on the porphyrin ring. The color scheme is as in Fig. 1.

is arguably less accurate for heterogeneous C-O bond breaking than the PBE functional used herein.

Unlike Paper I, we have focused on Eq. 3 and not the proton-assisted variation

$$\text{[Co(II)PCOOH]}^- + H^+ \rightarrow \text{Co(II)PCO(OH}_2);$$

$$\text{Co(II)PCO(OH}_2) \rightarrow \text{[Co(II)PCO]} + H_2O.$$  \hspace{1cm} (12)

Thermodynamically the two are equivalent.$^{30}$ In terms of kinetics, which Paper I does not deal with, they will have different activation barriers. To estimate $\Delta G^{(0)*}$ for Eq. (12) we note that all AIMD $W(R)$ reported for protonation/deprotonation reactions in the literature have been monotonic, i.e., the free energy changes and activation barriers are the same. Assuming the second half of Eq. (12) is also fast to the point of being barrierless, which is a lower limit on the overall Eq. (12) barrier, $\Delta G^{(0)*}$ for Eq. 4 would be entirely due to the $\Delta G^{(0)}$ of the first half of this equation and related to the $pK_a$ of Co(II)PCO(OH$_2$). At the pH=7 experimental conditions, the $pK_a$ of the C-OH$_2$ group in Co(II)PCO-OH$_2$ will have to be above 3.2 in order to have a lower barrier than the $\Delta G^* = 5.2$ kcal/mol we find for Eq. 3. This appears unlikely; C-OH$_2$ groups tend to be very acidic and lose protons readily. In any case, if this
alternate, proton assisted route of C-O cleavage were faster, the barrier estimated in our PMF calculation (Fig. 5) would be an upper bound to the reaction activation free energy; the C-O bond breaking step would still exhibit fast dynamics, and the qualitative conclusion of this paper would be unchanged. We plan to revisit Eq. 12 in the future.

B. Alternative Computational Methods

A two-dimensional PMF calculation should remove the hysteresis behavior in the protonation reaction (Eq. 11, Fig. 4). A convenient second variable may be the hydration number of the O atom in the OH group. This is because the black dashed curve in Fig. 4a exhibits \( N_w = 1.49 \) for that oxygen, considerably higher than the \( N_w = 0.75 \) for the red, well-behaved \( W(R) \) segment, suggesting that the electronic structure is correlated with the average hydration number. 2-D PMF simulations would be substantially accelerated using the metadynamics method\(^{81-83} \) over traditional umbrella sampling.

Our \( pK_a \) calculations suggest that deprotonation of weak acid groups which exhibit intramolecular hydrogen bonding and do not donate hydrogen bond to water may require the use of a large number of sampling windows. This difficulty may be circumvented by reversibly annihilating the proton using an artificial reaction pathway\(^{75} \) in a what might be called a “molecular grand canonical Monte Carlo” approach\(^{101} \). This method does introduce the disadvantage of changing the net charge in the finite-sized simulation cell, and may require a number of new conformational constraints.

Finally, the self-consistent DFT+U approach\(^{68} \) has been tested for Eq. 5. This method has the potential to establish a \( U \) value without resorting to parameterization with experimental results. Our preliminary studies suggest that this approach yields a Co(III)P-CO binding energy that is too small, but further development of this promising approach is under way\(^{102} \).

The above theoretical considerations, touching on many newly developed techniques, emphasize the complexity and challenges associated with modeling electrochemical reactions in explicit-water aqueous phase simulations.
C. Accuracy of DFT functionals and redox potentials

Accurate DFT functionals and dielectric continuum approximation of the aqueous solvent are critical for DFT+pcm determination of redox potentials, which in turn govern the viable reaction intermediates and the overall reaction mechanism. We have so far considered PBE, B3LYP, and DFT+U electronic structure methods with the $U$ parameter in DFT+U fitted to an experimental binding constant (Table I). B3LYP and DFT+U redox potentials track each other and should predict the same reaction mechanism at voltages slightly more negative than the Co(I)P/Co(II)P couple. In contrast, the PBE functional predicts that $[\text{Co(II)COOH}]^-$ is extremely unstable with respect to the far more acidic $[\text{Co(III)COOH}]$ complex near the PBE Co(I)P/Co(II)P redox potential. The B3LYP $\Phi_{\text{redox}}$ predictions are more consistent with AIMD hydration structure considerations and the fact that the CO$_2$-reduction reaction readily occurs near neutral pH in a carbonate buffer (Sec. III).

More accurate DFT functionals may be used in the future to further examine the mechanism proposed herein. Candidates include the M06 class of functionals designed to yield better thermochemistry accuracy for transition metal complexes, the B4(XQ3)LYP functional which has been found to yield improved redox potentials for a suite of test cases, and Gutzwiller wavefunction based methods. The quality of the PCM dielectric continuum model should also be further examined.

V. CONCLUSIONS

In this work, we have applied first principles calculations to examine the mechanism of the multi-step, two-electron electrochemical reduction of CO$_2$ to CO in water using cobalt porphyrin (CoP) as catalyst. First we have extracted redox potentials from DFT plus dielectric continuum solvation calculations using the B3LYP functional and a dielectric continuum treatment of water. Even though the absolute value for the $[\text{Co(I)P}]^-/\text{Co(II)P}$ couple is not in good agreement with experiments, the relative values of various redox potentials allow us to determine where the electron transfers occur among the four intermediate steps. Due to the enhanced interaction of CO$_2$ with water when bound to cobalt porphine, $[\text{Co(I)PCO}_2]^2-$ and $[\text{Co(II)PCOOH}]^-$ are the key intermediates. This finding may be useful not just for electrochemical reduction of CO$_2$, but for CO$_2$ capture from flue gas as well.
AIMD umbrella sampling calculations show that the pKₐ associated with [Co(II)PCOOH]⁻ deprotonation is about 9.0. This indicates that the protonation of [Co(I)PCO₂]²⁻ is downhill at the bicarbonate buffer experimental conditions (pH ~7). The subsequent cleavage of the C-OH bond is also exothermic, and the activation free energy involved is estimated to be only 5.2 kcal/mol. If we assume a vibrational pre-factor of $k=0.1$ ps⁻¹, C-OH cleavage should occur in nanosecond timescale at T=300 K. Hence two key steps in the multistep reaction should proceed readily, and it is likely that the electron transfer between the gas diffusion electrode and the polymerized porphyrin catalyst is the rate limiting step of the CO₂ to CO reduction reaction in water.⁵⁻¹³

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Supporting Information Available

Further information are provided regarding differences between the VASP and Gaussian packages; formic acid pKₐ calculation as a benchmark of the reaction coordinate used in this paper; the distribution Co-C-H angles in AIMD simulations; gas phase electron affinities of reaction intermediates; and cobalt charge state in CoPCOOH⁻ in implicit versus explicit solvent environment. This information is available free of charge via the Internet at [http://pubs.acs.org/](http://pubs.acs.org/).
Appendix: Hysteresis in the Deprotonation AIMD Simulation

This appendix discusses in more detail the hysteresis behavior during pKₐ PMF calculations (window 3 in Fig. II). As mentioned in Sec. II, all AIMD trajectories ultimately originate from a Monte Carlo simulation-equilibrated classical force field configuration where a CoPCOOH is fixed in a gas phase optimized, intramolecularly hydrogen-bonded geometry (Fig. II). The [Co(II)PCOOH]⁻ is immersed in water and equilibrated using AIMD at several stretched values of the reaction coordinates associated with Eq. 10 and Eq. 11 using harmonic potentials U(R) (Eq. 8).

During the equilibration run for the deprotonated window 2 of Fig. I we impose Rₒ = −1.12 Å on the initial [Co(II)PCOOH]⁻ system, which turns into a [Co(I)PCO₂]²⁻:H₃O⁺ contact ion pair within 1 ps. A maximally localized Wannier function analysis of an AIMD snapshot confirms that, in this ion pair, Co has spontaneously switched to the +I charge state which should be favored for large negative R. Therefore the AIMD trajectory in window 2 (and that in window 1, spawned from window 2) yields an unambiguous Co(I) charge state. Likewise, windows 4-7 are spawned successively from a window 4 trajectory equilibrated using Rₒ = −0.4 Å, and they reflect a Co(II) charge state which is favorable in this R range. An additional test further confirms that the Co(II) charge state spontaneously occurs at less negative R. We start with a [Co(I)CO₂]²⁻, Rₒ = −1.12 Å configuration in a equilibrated window 2 trajectory and abruptly switch to Rₒ = −0.4 Å (i.e., effectively jumping from window 2 to window 4 in Fig. III). [Co(II)PCOOH]⁻ is recovered within 1.2 ps, consistent with the Co(II) charge state observed in the window 4 trajectory which originally started out as [Co(II)PCOOH]⁻. The change in electronic structure in real time, represented by the changes in the spatial distribution of the spin density, is depicted in Fig. III. Thus, after an approximately 1 ps equilibration run, the final Co charge state becomes independent of initial conditions.

Only window 3, located in the Co(I)/Co(II) transition region, exhibits a strong dependence on initial conditions. We have initially started from a snapshot in window 4 and then switched Rₒ = −0.4 Å to Rₒ = −0.7 Å. The latter value of Rₒ does not apparently contain sufficient driving force to rapidly alter the Co charge state, and the system remains [Co(II)PCOOH]⁻ throughout the 10 ps sampling trajectory. This yields the dashed
$W(R)$ segment in Fig. 4a which exhibits a slope at the window edge that matches poorly to the window 2, $[\text{Co(I)PCO}_2]^2^-:\text{H}_3\text{O}^+$ contribution. To make progress, we restart the window 3 simulation from a snapshot of window 2, abruptly switch the umbrella potential from $R_o = -1.12$ Å there to $R_o = -0.7$ Å, equilibrate for 1 ps, and collect statistics for 10 ps. The system remains in the Co(I) charge state throughout the trajectory. The corresponding $W(R)$ segment in window 3 (full curve in Fig. 4a) matches reasonably well with those in windows 2 and 4, and is taken to be the final result.

An ergodic AIMD simulation should in principle spontaneously sample both Co(I) and Co(II) charge states. Thus, the correct $W(R)$ in this intermediate $R$ region should be a weighted average of the full and dashed curves. Apparently the statistical weight for Co(I) is much larger, so that only including Co(I) $W(R)$ information already yields a smooth $W(R)$ curve in Fig. 4.

A secondary, less significant type of hysteresis associated the position of the COOH proton also becomes apparent in Fig. 7. When the COOH group is intact, and no H$_2$O accepts a hydrogen bond from the COOH proton, the hydroxyl group intramolecularly hydrogen-bonds to a N atom on the porphine ring in a cis configuration (Fig. 1f). Enforcing $U(R)$ with $R < 0.5$ Å imposes COOH-H$_2$O hydrogen bonding that breaks this intramolecular coupling (Fig. 7a). When $R$ is further reduced to $R \sim -1$ Å, the [COOH]$^-$ proton is detached from the hydroxyl oxygen, landing on the hydrogen-bonding accepting H$_2$O molecule. This newly formed H$_3$O$^+$ spontaneously migrates away from the hydrophobic porphine ring and coordinates to what is now a CO$_2$$^-$ group in the axial (trans) position. Now the excess proton sticks out of the porphine plane. When we increase $R_o$ to reprotonate the COOH group, the system remains in this isomeric form (Fig. 7b). Isomerization between the two may entail a free energy barrier of several kcal/mol even for a stretched O-H, and does not occur on AIMD timescales in window 3. The lack of isomerization should have no significant effect on the $W(R)$ of window 3, while the gas phase Fig. 11 $[\text{Co(II)PCOOH}]^-$ intramolecular hydrogen bonded configuration is stabilized over the Fig. 11 isomer by about 4.93 kcal/mol, that hydrogen bond is already broken when the carboxylate proton donates a hydrogen bond to a water molecular (Fig. 7a). This secondary hysteresis may be avoided altogether using an artificial deprotonation coordinate, although it is not obvious the electronic hysteresis will be avoided as well.
FIG. 7: Panels (a) & (b) correspond to the solid and dashed line in window 3, respectively. The color scheme is as in Fig. [1]

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