

Modeling Operando Electrochemical CO2 Reduction

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# Modeling Operando electrochemical CO<sub>2</sub> reduction

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## Abstract

Since the seminal works on the application of density functional theory (DFT) and the computational hydrogen electrode to electrochemical CO<sub>2</sub> reduction (eCO<sub>2</sub>R) and hydrogen evolution (HER), the modeling of both reactions has quickly evolved for the last two decades. Formulation of thermodynamic and kinetic Linear Scaling Relationships for key intermediates on crystalline materials have led to the definition of activity volcano plots, overpotential diagrams, and full exploitation of these theoretical outcomes at laboratory scale. However, recent studies hint at the role of morphological changes and short-lived intermediates in ruling the catalytic performance under operation conditions, further raising the bar for the modeling of electrocatalytic systems. Here, we highlight some novel methodological approaches employed to address eCO<sub>2</sub>R and HER reactions. Moving from the atomic scale to the bulk electrolyte, we first show how *ab initio* and machine learning (ML) methodologies can partially reproduce surface reconstruction under operation, thus identifying active sites and reaction mechanisms if coupled with microkinetic modeling. Later, we introduce the potential of DFT and ML to interpret data from *Operando* spectroelectrochemical techniques, such as Raman spectroscopy and Extended X-ray Absorption Fine Structure characterization. Next, we review the role of electrolyte and mass transport effects. Finally, we suggest further challenges for computational modeling in the near future as well as our perspective on the directions to follow.

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# 1 Introduction

Electrocatalysis is a crucial process which “facilitates conversion between electrical and chemical energy in fuel cells and electrolysis devices”.<sup>1</sup> Conversion of chemical energy into electrical energy takes place in a “galvanic” cell through spontaneous reactions occurring at the two electrodes once connected through a conductor.<sup>2</sup> Instead, exploitation of electrical energy to carry out chemical reactions takes place in an “electrolytic” cell by applying an external voltage across the electrodes greater than the open circuit potential.<sup>2</sup> In this review, we focus on electrocatalytic processes applied to electrolytic cells.

Due to the technological advances and consequent cost reduction enabled by the economy of scale of renewable energy within the last decades,<sup>3</sup> sustainable storage technologies as conversion of water to hydrogen (Hydrogen Evolution Reaction, HER) or electrochemical reduction of carbon dioxide (eCO<sub>2</sub>R) will likely compete soon with the exploitation of fossil fuels.<sup>4,5</sup> The uptake of both processes has quickly evolved from pioneering outlooks at the beginning of the last century,<sup>6</sup> mainly as a result of key investigations,<sup>7–11</sup> as well as practical breakthroughs.<sup>3,4,12</sup> Over the last decade, most of the computational studies have been centered in going beyond Linear Scaling Relationships (LSR) and Computational Hydrogen Electrode (CHE),<sup>8</sup> typically applied to estimate the thermodynamics and kinetics of well-defined catalytic systems. In the present review we focus on the material gap between ideal structural models and the *in situ* catalytic phases under reaction conditions.

Very recently, new challenges are arising from processes occurring across the electrocatalytic cell during operation,<sup>13–15</sup> such as electric field, electrolyte, and mass transport effects, thus requiring more complex methodological approaches to rationalize the overall performance.<sup>16</sup> Many heavy methodological developments have been introduced to consider solvents either implicitly or explicitly and the effect of electric field (*i.e* the jDFT<sub>x</sub> scheme).<sup>17–19</sup> These approaches aim at incorporating these terms into the Kohn-Sham DFT formalism in an increasing self-consistent manner.<sup>18</sup> In contrast, much less attention has been paid to other modeling aspects, like the surface modifications induced at the electrode-electrolyte

interface or the impact of the current in restructuring the materials.

After a short preliminary discussion on the state-of-the-art research works, this review aims at giving a subjective, yet accurate overview about the advances in modeling electrochemical CO<sub>2</sub> reduction which have occurred for the last five years. Across the article, mentions to hydrogen evolution (competing reaction) and Oxygen Evolution Reaction (OER, anodic process) will instead be limited to specific cases. We start by discussing the main methodologies developed to rationalize eCO<sub>2</sub>R on crystalline materials under well-defined reaction conditions (Section 2). Here we review approaches to reproduce the effect of electrochemical potential,<sup>20</sup> solvation,<sup>21–26</sup> and the concepts of network analysis and dimensionality reduction inherent to Linear Scaling Relationships.<sup>27,28</sup> Section 3 highlights new insights and corresponding challenges from *Operando* studies of electrocatalytic processes. Moving from catalyst to diffusion layer, we consider (1) reconstruction of active sites, (2) detection and identification of key reaction intermediates, and (3) electrolyte effects. In the core of the review, we discuss novel theoretical frameworks to tackle these experimental challenges. Initially, we focus on *ab initio* and machine learning techniques to model surface reconstruction under eCO<sub>2</sub>R conditions (Section 4). Secondly, we underline the role of DFT vibrational analysis and microkinetic studies to complement spectroscopy and kinetic data to define reaction pathways for eCO<sub>2</sub> up to *n*-propanol (Sections 5-6). After, we introduce the relevance of multiphysics modeling and *ab initio* simulations to assess potential influences of pH and electrolyte species in CO<sub>2</sub> reduction activity and selectivity (Section 7). Section 8 offers a short discussion over applications of machine learning to (1) the development of atomistic potentials toward high computational efficiency and (2) the automatic extraction of structural properties from experimental data. Finally, we conclude the review by offering our own perspective on future developments needed in the field of computational electrocatalysis, as well as an outlook on relevant remaining research questions.

## 2 CO<sub>2</sub> reduction on crystalline materials

### 2.1 Hori's Milestones

The pioneering concept introduced by Giacomo Luigi Ciamician in 1912 – *By using suitable catalyzers, it should be possible to transform the mixture of water and carbon dioxide into oxygen and methane*<sup>6</sup> – is now a reality. In the field of eCO<sub>2</sub>R, research has advanced from preliminary reports of polarization signals due to CO<sub>2</sub> reduction on mercury cathodes,<sup>29</sup> to production of added value C<sub>2+</sub> products at high selectivities and current densities.<sup>4,12</sup> Catalytic search and mechanistic studies carried out by Hori and collaborators during the last forty years have elucidated the complexity of this reaction, thus providing key insights and experimental data for posterior theoretical rationalization.<sup>9</sup>

Since many of the milestones in the field naturally derive from Hori's work, we find it adequate to recall the main findings below.<sup>7,30-37</sup>

1. **Product distribution.** eCO<sub>2</sub>R on Cd, Sn, Pb, In mainly accounts for formate. Zn is selective to formate or CO, whilst Cu is the only material which allows formation of C<sub>2+</sub> compounds. Ag and Au catalyze CO<sub>2</sub> reduction to CO, whereas Ni and Fe are only active for HER.<sup>7</sup>
2. **Cation effect.** On a copper catalyst, cations with larger ionic radius promote CO<sub>2</sub> (CO) reduction to ethylene to the detriment of CH<sub>4</sub> production and eCO<sub>2</sub>R activity correlates with cation ionic radius.<sup>32</sup>
3. **CO as key eCO<sub>2</sub>R intermediate. pH effect (vs SHE reference).** High CO coverage ( $\sim 90\%$ ) is observed during CO<sub>2</sub> reduction to hydrocarbons,<sup>33</sup> suggesting the relevance of adsorbed CO species. Ethylene and ethanol selectivity increases for bulk alkaline pH, whilst acidic pH favors methane production.<sup>37</sup> Thus, the rate-determining step (RDS) to form methane is expected to depend on pH, whilst ethylene production is pH-independent.<sup>37</sup> Spectroscopic signals for adsorbed CO are observed between  $-0.8$

and  $-1.0$  V vs NHE,<sup>34</sup> and are promoted by higher temperatures.<sup>30,31</sup>

4. **Copper facet sensitivity.** Cu(111) favors eCO<sub>2</sub>R to methane. Cu(100) enables ethylene production, which is further promoted on (111) steps on (100). Conversely, (100) planes with (110) steps are mainly selective to ethanol.<sup>35,36</sup>
5. **Surface poisoning.** Metals contaminants, such as Fe<sup>2+</sup> and Zn<sup>2+</sup>, can poison copper electrodes during operation, causing its deactivation.<sup>38</sup>

## 2.2 Computational modeling

Many of the observations of Hori *et al.* for eCO<sub>2</sub>R on well-defined catalysts (Section 2.1) in addition to evidence on other electrocatalytic reactions have been elucidated during the last decade through the introduction of the Computational Hydrogen Electrode<sup>9</sup> and the observation of Linear Scaling Relationships between CO<sub>2</sub> reduction intermediates (Sections 2.2.4).<sup>11,39</sup>

### 2.2.1 The “Anderson” Electrode

Albeit typically overlooked, the contribution of Anderson and his research group has been fundamental to the application of computational chemistry to electrocatalysis.<sup>20</sup> Already in 1982, Anderson and Ray studied hydrogen evolution on a Fe<sub>5</sub> cluster from H<sub>3</sub>O<sup>+</sup> and H<sub>2</sub>O reduction over a 4 V potential range simulated by shifting the Fe ionization potential.<sup>40</sup> This elegant approach was one of the first strategies to explicitly include potential bias in estimation of activation energies. Besides, the validity of this work is further highlighted by the assessment of different proton sources for HER, topic which has recently become very relevant in the field.<sup>41,42</sup> Few years later, Anderson and Kang extended their previous model by introducing an electron donor in the simulation cell to assess hydrogen evolution on diamond.<sup>43</sup> In this study, they employed H<sub>3</sub>O<sup>+</sup> as proton source and CH<sub>4</sub> as a model molecule for the C-H bond (H on the diamond system). Besides, they placed a Li atom

coordinated to 1-2 water molecules 10 Å far away from the CH<sub>4</sub> molecule. Since equilibrium is assumed at the electron transfer, the electrochemical potential  $U$  of Li is given by Equation 1, where  $IP$  stands for the ionization potential of the alkali cation and 4.6 V represents the potential vs SHE of the electron. Different applied potentials were then estimated by simply tuning Li ionization potential. Remarkably, the use of alkali metals as electron donors has been reintroduced in 2016 by Nørskov and co-workers to assess explicit field effects in electrochemical CO<sub>2</sub> reduction on Ag,<sup>44</sup> and recently an analogous approach has allowed to rationalize the key role of cations in eCO<sub>2</sub>R and HER on Au, Ag, and Cu.<sup>42,45</sup>

$$U = (IP/|e|^- - 4.6) \text{ V} \tag{1}$$

One year later (1999), Anderson and Albu extended this framework to a general electron donor entity,  $D$ , representing the working electrode.<sup>46,47</sup> The chemical potential  $\mu(D)$  of electrons at the electrode is defined as their Fermi energy,  $E_F$ , or equivalently as the negative of the electrode thermodynamic workfunction  $\phi$ . On the standard hydrogen electrode scale, the electrode potential is then given by Equation 2, where the formula can be further simplified recalling the average value of 4.6 eV for  $\phi_{\text{H}^+/\text{H}_2}$  and that  $\phi = -\mu(D)$ . The electron transfer to a specific reduction entity  $R$  is assumed to take place when the electron affinity of  $R$ ,  $E_{\text{ea}}$ , matches the chemical potential of the donor,  $\mu(D)$ . The electron affinity of  $R$  can be estimated computationally as the difference between the energy of  $R$  and  $R^-$  with constant structure. Thus, different structures can be sampled and the ones which fulfill the relation  $E_{\text{ea}} \sim \mu(D)$  identified. Finally,  $U$  can be derived from  $E_{\text{ea}}$  from the lowest energy configuration, see Equation 2.<sup>46,47</sup>

$$U = \phi - \phi_{\text{H}^+/\text{H}_2} = E_{\text{ea}} - 4.6 \text{ eV} \tag{2}$$

The last generalization of the “Anderson” electrode was developed to assess Oxygen Reduction Reaction (ORR) on a Pt<sub>3</sub>Cr(111) catalyst.<sup>48,49</sup> First, Roques and Anderson con-

sidered the well-known relationship between the variation of Gibbs free energy  $\Delta G^0$  for a given reduction reaction and the reversible potential  $U^0$ , Equation 3, where  $n$  is the number of electrons involved in the reduction and  $F$  is the Faradaic constant.<sup>49</sup>  $\Delta G^0$  can be estimated through the the reaction energy  $E_r$  plus a constant  $k$  which includes enthalpic, entropic, and solvation contributions. Equation 3 can then be further expanded as Equation 4, where the 4.6 V factor is due to the electron energy at 0 V vs SHE. In the computational study, Equation 4 allowed to determine the reversible potential for the reduction of OH radical (*i.e.* the reactant) to water (*i.e.* the product) with significant accuracy:  $U_{\text{theo}}^0 = 2.99$  V vs  $U_{\text{exp}}^0 = 2.81$  V.

Generalizing this approach, the reversible potential for a specific reduction reaction on a given catalyst  $M$  can be estimated by computing reactant  $R$  and product  $P$  binding strengths on the catalytic surface and correcting for the reversible potential given in aqueous solution,  $U_{\text{aq}}^0$ , see Equation 5.<sup>48,49</sup> As pointed out in Ref. 20 the only real difference between this methodology and the successive Computational Hydrogen Electrode formalism<sup>8</sup> is the choice of the OH radical as reference compound instead of  $\text{H}_2$ . Otherwise, both approaches are conceptually identical, since they include applied potential effects as linear corrections on the intermediates adsorption energies.

$$U^0 = -\frac{\Delta G^0}{nF} \quad (3)$$

$$U^0 = -\left(\frac{E_r}{nF}\right) - 4.6 \text{ V} + k \quad (4)$$

$$U^0(M) = U_{\text{aq}}^0 + \frac{[E_{\text{ads}}(P)_M - E_{\text{ads}}(R)_M]}{nF} \quad (5)$$

### 2.2.2 The Computational Hydrogen Electrode

Few months after the introduction of the ‘‘Anderson’’ electrode in February 2004,<sup>48,49</sup> Nørskov *et al.* introduced the Computational Hydrogen Electrode formalism to assess the effect of proton-coupled electron transfers (PCETs) on the thermodynamics of the oxygen reduc-

tion reaction on Pt(111) (Figure 1a).<sup>8</sup> This procedure is based on six steps and it is only applicable for reaction routes which occur *via* PCETs:

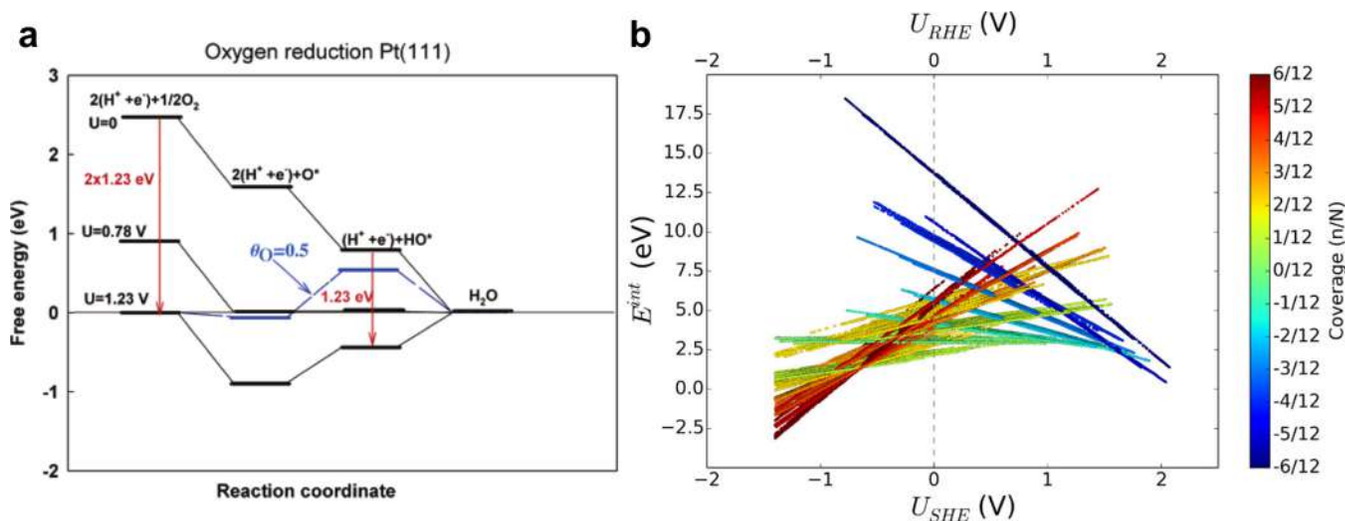
1. Following the Standard Hydrogen Electrode (SHE) scale, at standard conditions (pH = 0, 1 H<sub>2</sub> bar,  $T = 298.15$  K,  $U = 0$  V vs SHE), we can relate the chemical potential of a proton/electron couple to one half of H<sub>2</sub> Gibbs free energy.
2. Solvation contributions must be included either through explicit consideration of water molecules in the supercell,<sup>8,22,25</sup> or through implicit continuum models based on solvent dielectric permittivity.<sup>23,24,26,50–52</sup> Mixed schemes with explicit and implicit terms are less widely used than in homogeneous catalysis.
3. The Gibbs free energy of all the intermediates which involve an electron transfer (ET) must be shifted by  $-n|e^-|U$ , where  $n$  is the overall number of electron transferred (with electric charge in elementary unit  $|e^-|$ ) and  $U$  the applied bias. Equation 6 reports a practical example to calculate the Gibbs free energy variation for CO<sub>2</sub> reduction to COOH *via* 1 PCET.

$$\Delta G^*_{\text{COOH}} = G_{\text{products}} - G_{\text{reactants}} = G^*_{\text{COOH}} - (G_{\text{CO}_2} + \frac{1}{2}G_{\text{H}_2} - |e^-|U) \quad (6)$$

4. Electric field effects are estimated as  $\vec{p} \cdot \vec{E}_{\text{EDL}}$ , where  $\vec{p}$  is the dipole moment of the adsorbate and  $\vec{E}$  is the electric field within the Electrical Double Layer (EDL).
5. At pH  $\neq 0$ , the Gibbs free energy of the proton is linearly corrected *via*  $k_{\text{B}} \cdot T \cdot \ln(10) \cdot \text{pH}$ , in line with the Reversible Hydrogen Electrode (RHE) standard.
6. Gibbs free energy of adsorption must account for zero point energies and entropic contributions, including vibrational terms for adsorbed molecules).

The CHE formalism permitted the first mapping of experimental evidence to first principles calculations, such as cyclic voltammograms for H on Pt surfaces<sup>53</sup> and Pourbaix diagrams,<sup>54</sup> thus allowing significant advances in the field.<sup>55</sup> In the first application of the CHE

to CO<sub>2</sub> reduction,<sup>9</sup> overpotentials to form H<sub>2</sub>, HCOOH, CO, and CH<sub>4</sub> obtained by DFT showed excellent agreement with experimental data from Hori's eCO<sub>2</sub>R studies on copper (Section 2.1).<sup>30,31</sup> After more than a decade of success, the CHE is considered the fundamental framework to model the role of potential in the thermodynamics of electrochemical reactions under the assumption that activation energies can be initially disregarded.<sup>56</sup> Nevertheless, alternative approaches have been developed to include potential effects on reaction kinetics, as nicely summarized in this recent Review.<sup>56</sup>



**Figure 1:** **a**, Determination of the Gibbs free energy diagram for oxygen reduction on Pt(111) at  $U=0$  V vs SHE,  $U = +0.78$  V vs SHE, and  $U = +1.23$  V vs SHE through the CHE formalism. The energy profile at  $+1.23$  V vs SHE for 0.5 ML oxygen coverage is highlighted in blue.<sup>8</sup> **b**, Interface energy vs  $U_{SHE}$  and  $U_{RHE}$  at pH = 0 calculated for different snapshots of *ab initio* molecular dynamics simulations carried out on a water/Au(111) interface with different number  $n$  of H<sup>+</sup> added to the system.  $N$  denotes the number of surface Au atoms (12).<sup>57</sup> Figures respectively adapted with permission from Ref. 8 (Copyright 2004 American Chemical Society) and Ref. 57 (Copyright 2016 American Chemical Society).

### 2.2.3 Beyond the Computational Hydrogen Electrode

Since the CHE formalism accounts for electric field effects only *a posteriori* from the dipole moment of the adsorbate (step 4 in Section 2.2.2),<sup>8</sup> recently Rossmeisler *et al.* developed a generalized formalism (GCHE) to include explicit interactions between electric field, intermediates, and solvent molecules.<sup>58</sup> Along with postulating that the bulk electrolyte is in

equilibrium and that the interface is in equilibrium with bulk electrodes and electrolyte, the GCHE assumes that the interface region is charge neutral and large enough to screen all charges.<sup>57</sup> Thus, a direct correlation exists between electron wave function, obtainable for a metallic surface through DFT, and electrode potential vs SHE at the electrochemical equilibrium. Additionally, the electrode potential can be referred to the RHE scale by correcting for the different pH.

According to the GCHE formalism,<sup>58</sup> the electrochemical potential of a proton/electron couple,  $\mu_{\text{H}^+ + \text{e}^-}$ , can be calculated from Equation 7, where  $\Delta G_{\text{d}}$  is the dissociation energy for  $\frac{1}{2}$  H<sub>2</sub>,  $\Delta G_{\text{i}}$  is the H ionization energy,  $\phi_{\text{e}^-}$  and  $\phi_{\text{H}^+}^0$  are respectively the work functions of electron and proton in solution (at pH = 0), measured with respect to the vacuum. Since for  $U = 0$  V vs SHE and pH = 0 the electrochemical potential of the proton/electron couple is zero, Equation 7 can be simplified to Equation 8, where  $\phi_{\text{e}^-}(\text{SHE}) = 4.44$  eV is the work function of an electron in a metal at the standard hydrogen electrode potential. Thus,  $\mu_{\text{H}^+ + \text{e}^-}$ ,  $\phi_{\text{e}^-}$ , and pH are related *via* Equation 8 for a given metal/solution interface.  $\phi_{\text{e}^-}$  can be calculated from the electrode potential  $U$ , obtained from the interface dipole within the simulated cell, Equation 9.

$$\mu_{\text{H}^+ + \text{e}^-} = \Delta G_{\text{d}} + \Delta G_{\text{i}} - \phi_{\text{H}^+}^0 - 2.3 \cdot k_{\text{B}} \cdot T \cdot \text{pH} - \phi_{\text{e}^-} \quad (7)$$

$$\mu_{\text{H}^+ + \text{e}^-} = \phi_{\text{e}^-}(\text{SHE}) - 2.3 \cdot k_{\text{B}} \cdot T \cdot \text{pH} - \phi_{\text{e}^-} \quad (8)$$

$$U_{\text{SHE}} = \frac{\phi_{\text{e}^-} - \phi_{\text{e}^-}(\text{SHE})}{e^-} \quad (9)$$

Since interface dipole, electrode potential, and electron work function are constant for a given structure, the Gibbs free energy of a metal/solution interface normalized per surface atom is defined at the electrochemical equilibrium (pH = 0) by Equation 10.  $n$  and  $N$  are respectively the number of hydrogens and surface metal atoms within the systems, while  $G_{N,n}$ ,  $G_{N,0}$ ,  $G_{\text{H}_2}$  are the Gibbs free energies of metal/solution interface, clean surface, and gas phase H<sub>2</sub>. These energies can be either calculated *via* DFT simulations or derived from

experimental data. Equation 10 can be extrapolated to account for  $\text{pH} \neq 0$  as in Equation 11, where the proton/electron couple chemical potential can be obtained from Equations 8 and 9, knowing  $\text{pH}$  and estimating electron work function from the interface dipole.

$$G^{\text{interface}}(\mu_{\text{H}^+ + \text{e}^-} = 0, \phi_{\text{e}^-}) = \frac{G_{N,n} - G_{N,0}}{N} - \frac{1}{2} \frac{n}{N} G_{\text{H}_2} \quad (10)$$

$$G^{\text{interface}}(\mu_{\text{H}^+ + \text{e}^-}, \phi_{\text{e}^-}) = \frac{G_{N,n} - G_{N,0}}{N} - \frac{1}{2} \frac{n}{N} G_{\text{H}_2} - \frac{n}{N} \mu_{\text{H}^+ + \text{e}^-} \quad (11)$$

Following the GCHE principles relevant electrochemical properties, such as potentials of zero charge, Gibbs isotherms, and differential capacities as a function of  $\text{pH}$ , were estimated from DFT simulations on a Au/water interface.<sup>57</sup> Besides, the same methodology was applied to a Pt/water interface to estimate OH coverage vs bulk  $\text{pH}$ .<sup>59</sup> In Ref. 57, hydrogen atoms were either added or removed from the solvation layer and the resulting systems underwent *ab initio* molecular dynamics at 300 K with time step 0.5 fs to account for 90000 states with different number of protons in the system. Thus, the authors estimated the Gibbs free energy of the metal/solution interface as a function of applied potential and  $\text{pH}$  from Equation 11, see Figure 1b. They calculated the Gibbs free energies of Au/water system, the clean surface, and electron work function for all the AIMD snapshots with DFT. Finally, static properties of the interface were derived from grand canonical Metropolis Monte Carlo averages on the snapshots to define an ensemble with physical values of surface coverage and proton excess at any given electrode potential and  $\text{pH}$ .

Further approaches to relate thermodynamics/kinetics and electrochemical potentials involve the explicit insertion of electric field in slabs through an artificial dipole moments,<sup>44,60–63</sup> or the self-consistent solution of the Poisson-Boltzmann (PB) equation and the Kohn-Sham equations.<sup>64–66</sup> We refer to Ref. 20 for an accurate review of these methodologies.

### 2.2.4 Linear Scaling Relationships

In agreement with the Brønsted-Evans-Polanyi relation, first derived from experimental evidences,<sup>67,68</sup> kinetic Linear Scaling Relationships (kLSR) were demonstrated between activation barrier and reaction energy for different reactions.<sup>69–73</sup> Few years later Abild-Pedersen *et al.* developed a set of scaling properties between adsorption energies for hydrogen-containing molecules, the so-called thermodynamic Linear Scaling Relationships (tLSR), see Figure 2a.<sup>10</sup> The rationale behind these dependencies can be traced back to bond-order conservation theory and valence (octet) rules for light elements. As suggested by the *d*-band model,<sup>74,75</sup> the reactivity of a transition metal is mainly given by its *d*-band and this property affects an adsorbate depending on the bond order with the surface. When comparing the same metal in two environments, the system characterized by higher *d*-band center is more reactive, due to enhanced hybridization between the bonding and anti-bonding adsorbate states and the metal electronic states.<sup>74</sup>

According to the tLSR,<sup>10</sup> the adsorption energy for a given intermediate containing a hydrogen atom correlates with the bond order and  $\Delta E_{sp}$ , parameter which depends on *sp* contribution to binding energy. In the case of adsorption of  $\text{CH}_x$  molecules on surfaces, tLSR are mathematically described by Equation 12, where  $\gamma(x) = \frac{x_{\max} - x}{x_{\max}}$ ,  $\xi = \Delta E_{sp}^{\text{CH}_x} - \gamma(x)\Delta E_{sp}^{\text{C}}$ , and  $x_{\max}$  is the maximum number of H atoms which can bond to C.

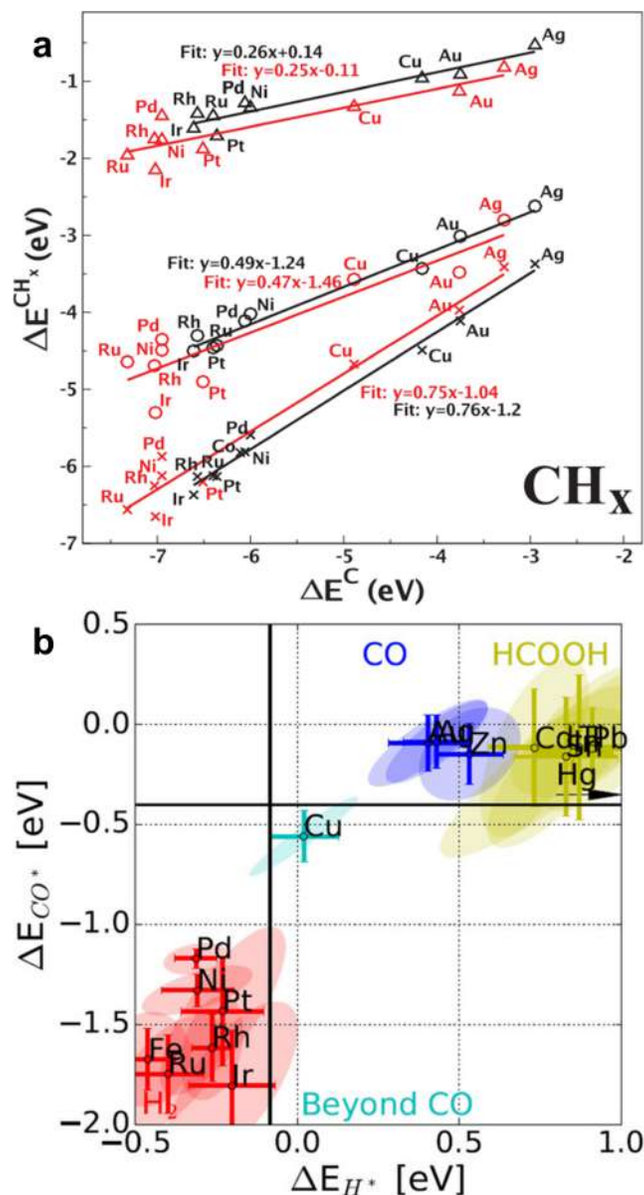
$$\Delta E^{\text{CH}_x} = \gamma(x)\Delta E^{\text{C}} + \xi \tag{12}$$

Analogous tLSR were demonstrated for  $\text{NH}_x$ ,  $\text{OH}_x$  and  $\text{SH}_x$  intermediates,<sup>10</sup> thus suggesting their validity for other electrocatalytic reactions as well. In oxygen evolution reaction, binding energies of the reaction intermediates (\*OH, \*OOH, and \*O) fulfill universal tLSR on both metallic<sup>76</sup> and oxidic systems,<sup>77</sup> thus leading to limitation of catalytic performance. Such limiting effect has been nicely exemplified through volcano plots correlating activity (or overpotential) for OER and energetic descriptors, as \*O binding strength<sup>76–79</sup> or energy

differences (e.g.  $\Delta E_{*O} - \Delta E_{*OH}$ ).<sup>80</sup> Similar volcano plots were defined for hydrogen evolution reaction, correlating  $*H$  binding and HER activity as first proposed by Trasatti.<sup>41,81</sup> Three decades later, Nørskov *et al.* employed this concept to rationalize the outstanding performance of Pt for HER, which is due to its thermoneutral  $*H$  chemisorption energy.<sup>82</sup>

In addition to tLSR based on electronic effects, analogous relationships were demonstrated between adsorption energies and coordination numbers of the active species,<sup>83-85</sup> predicting that surface defects exhibit stronger binding of cathodic reaction intermediates due to their lower coordination and consequently higher localized electronic density. Whilst the valence of the adsorbate (Equation 12) rules the slope of tLSR, both valence and local coordination of the catalyst center provides the offset. Hence, proper synthetic guidelines can tune the interplay of these effects to design systems with high performance by combining both electronic and structural dependencies of surface reactivity.<sup>85</sup> The application of this theoretical framework enabled the prediction of catalytic reactivity of defective surfaces for ORR and the identification of product-specific active sites motifs for  $eCO_2R$  on Cu.<sup>86,87</sup> Apart from coordination environment, local strain affects binding properties on metals as well, since lattice expansion leads to up-shift of the metal  $d$ -band and consequent stronger adsorption of reaction intermediates. Such phenomenon was first introduced by Mavrikakis and co-workers for  $*O$  and  $*CO$  on Ru and Au.<sup>88,89</sup> We anticipate here that this strain effect may be the physical foundation of the previously discussed coordination number-based linear scaling relationships.<sup>83-85</sup>

Since both thermodynamic and morphological linear scaling relationships are based on the  $d$ -band model, we here briefly mention a recent study from our group, highlighting the shortcomings of this framework to rationalize the adsorption energies of group 12 metals, such as Zn and Cd.<sup>90</sup> The reactivity of these two metals cannot be entirely described by their  $d$ -band center, suggesting the need for a second parameter. Hence, the authors introduced a second physical descriptor, the reduction potential,<sup>91</sup> to account for ionic effects on top of solely covalent interactions ruled by the  $d$ -band center. A statistical Principal Component



**Figure 2:** a, Adsorption energies of  $\text{CH}_x$  ( $x = 1, 2, 3$  as crosses, circles, and triangles, respectively) vs adsorption energy of C on transition metals. Molecules in vacuum and clean surfaces were taken as energy references. Data points for close-packed and stepped surfaces are colored in black and red respectively.<sup>10</sup> b,  $\Delta E_{\text{CO}^*}$  and  $\Delta E_{\text{H}^*}$  as descriptors to classify transition metals into  $\text{H}_2$  evolving catalysts (strong H binding), systems selective to  $\text{C}_{2+}$  (weak H binding, mild CO binding), and metals evolving CO (no H binding, weak CO binding) or  $\text{HCOO}^-$  (no H neither CO binding).<sup>11</sup> Black lines highlight boundaries for H ( $x$ -axis) and CO adsorption ( $y$ -axis). Figures adapted with permission from Ref. 10 (Copyright 2007 American Physical Society) and Ref. 11 (Copyright 2017 Wiley-VCH) respectively.

Analysis (PCA) applied to DFT thermochemical data for 71  $\text{*H-}$ ,  $\text{*O-}$ , and  $\text{*C-}$ terminated  $\text{C}_1$ - $\text{C}_2$  adsorbed species on 12 close-packed metal surfaces confirmed validity of both covalent and

ionic descriptors to determine surface reactivity. By enabling prediction of surface reactivity, the developed framework enables quick screening of binding energy of intermediates and consequently it is transferable to the assessment of any reaction of choice.<sup>92</sup>

Following the discussion on application of tLSR to eCO<sub>2</sub>R, Bagger *et al.* applied these concepts to classify the selectivity toward C<sub>1</sub>-C<sub>2</sub> for *d*- and *p*-block metals,<sup>11,93</sup> observed experimentally by Hori *et al.* (Section 2.1).<sup>7</sup> A  $\Delta E_{\text{*HCOO}}$  vs  $\Delta E_{\text{*COOH}}$  LSR failed to differentiate between catalysts selective to CO (Ag, Au) and HCOOH (Zn),<sup>11</sup> thus they proposed three other descriptors:  $\Delta E_{\text{*COOH}}$ ,  $\Delta E_{\text{*H}}$ , and  $\Delta E_{\text{*CO}}$ , see Figure 2b.<sup>11</sup> The authors noted that metals which strongly bind \*H are mainly selective to H<sub>2</sub>, whilst CO-producing catalysts can adsorb both \*COOH and \*H. Besides, ideal systems for catalyzing eCO<sub>2</sub>R to HCOOH must have endothermic \*H adsorption. As for eCO<sub>2</sub>R to C<sub>2</sub> products, a selective catalyst must exhibit an intermediate CO binding to hinder its release to the electrolyte, yet enabling C-C coupling.<sup>93</sup> Thus, Cu sits at the sweet spot, since it exhibits strong enough CO binding, but thermoneutral H adsorption. Finally, oxygen binding energy was proposed as a descriptor for alcohol vs hydrocarbon competition and this prediction was later confirmed by experimental reports on eCO<sub>2</sub>R to methanol or methane on Au, Cu, and Ag electrodes.<sup>11,94,95</sup> As a crucial remark, we highlight that the above-mentioned descriptors are specifically tailored to single metals, thus alloys or oxide-derived materials may have completely different reactivity, see for instance eCO<sub>2</sub>R to C<sub>2</sub> products (specifically ethylene glycol) on Fe<sub>2</sub>P<sup>96</sup> and nickel oxygenates.<sup>97</sup>

In a following study, Bagger *et al.* included considerations on both thermodynamics and surface morphology to rationalize Hori's results on copper single crystals (Section 2.1)<sup>35,36</sup> Since CO and H binding energies did not vary adequately to account for the different eCO<sub>2</sub>R selectivity, the authors considered surface coordination number ( $N_{\text{coord}}$ ) and adsorption energy ( $\Delta E$ ) of an extended list of intermediates (\*OH, \*OCCOH, \*O, \*H, \*COOH, \*CO, \*C) as potential descriptors. By applying Principal Component Analysis on experimental Faradaic efficiencies, the authors demonstrated that structural properties can describe

C<sub>2+</sub> compounds selectivity on copper better than binding energies:<sup>39</sup> (111) facet promotes methane, (110) acetaldehyde, (100) ethylene, whilst  $n(100) \times (110)$  steps are responsible for ethanol production.

### 3 Challenges from electrocatalytic processes under working conditions

As we have seen, computational simulations on purely crystalline models were very effective in reproducing and rationalizing experimental observations on single crystals and under well-defined conditions (Sections 2.2.2-2.2.4). Particularly, all the terms contributing to the Gibbs free energies have been benchmarked extensively, demonstrating that the equations to obtain the energies are robust and contain all the key elements to constitute an accurate first order approach. However, limitations have been identified as the thermodynamic descriptor inherent to the CHE approach might not be sufficient. For instance, in the HER process the typically assumed  $\Delta G_H = 0$  eV target of the volcano plot has been challenged due to coverage, kinetic, and the explicit applied overpotential effects.<sup>98,99</sup> In addition, new challenges are arising from *Operando* characterization when electrocatalytic systems are ruled by dynamic processes. This particularly affects the material gap between the prepared catalyst and its state under reaction conditions along with mass transport and concentrations at the interfaces between the electrode, electrolyte, and bulk solution.

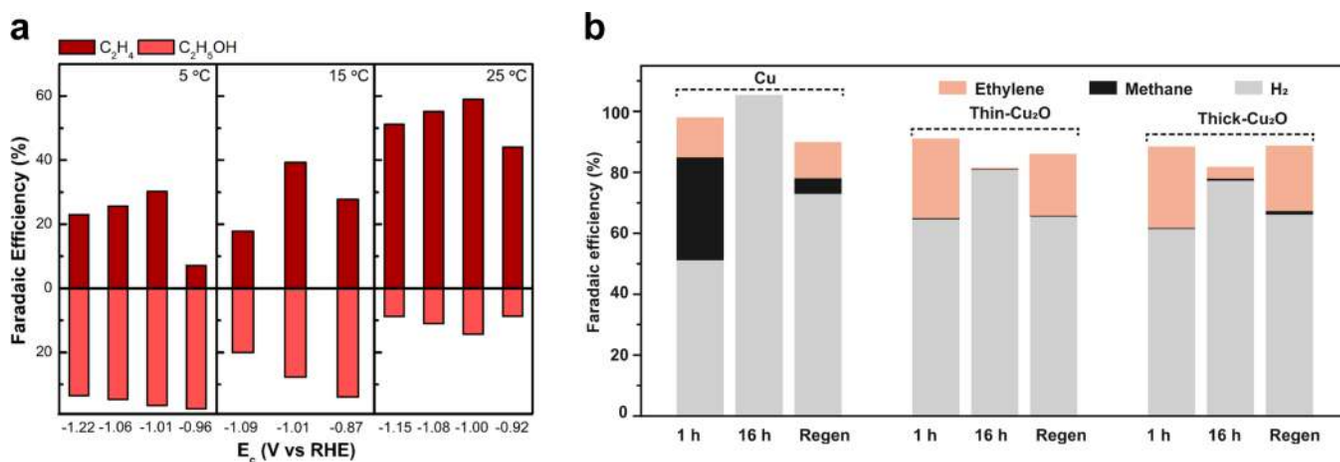
#### 3.1 Reconstruction of active sites during operation

Recently, many different observations of surface reconstruction towards optimized electrocatalytic performance were put forward. In the field of electrochemical CO<sub>2</sub> reduction, copper surface reconstruction has been discussed to a large extent during the last decades.<sup>14,63,100–103</sup> *Operando* characterization *via* Electrochemical Scanning Tunneling Microscopy (EC-STM) and Raman spectroscopy highlighted the reconstruction of polycrystalline copper (pc-Cu)

to (100) facets,<sup>102</sup> and nanocuboids.<sup>63</sup> Air-oxidized copper naturally reconstructed toward Cu(100) facets at cathodic bias,<sup>104</sup> as well as Cu(110) and copper materials including both polycrystalline and (111) facets.<sup>105</sup> Generally, product distribution and activity of copper catalysts strongly depends on the synthetic procedures, thus suggesting a completely different morphology on reconstructed surfaces than single crystals.<sup>35,36,106</sup> Specifically, oxide-derived copper mainly produce ethylene,<sup>107-114</sup> ethanol,<sup>109</sup> *n*-propanol,<sup>109,115</sup> and traces of acetate and ethane,<sup>109,116</sup> in line with the eCO<sub>2</sub>R performance of copper nanoparticles at higher overpotentials<sup>117-120</sup>

Recent studies further confirm the role of metastable or kinetically trapped phases, surface reconstruction, and solvent molecules in tuning catalytic performance.<sup>14,107,112,114,121</sup> Carbon isotopic labelling (<sup>12</sup>CO<sub>2</sub> vs <sup>13</sup>CO<sub>2</sub>) differentiated between three specific active sites on oxide-derived copper (OD-Cu) catalyst, selective to (1) ethanol and acetate; (2) ethylene, and (3) 1-propanol, whilst no difference was found for crystalline Cu.<sup>122</sup> Furthermore, a Cu<sub>2</sub>O-derived material showed an outstanding CO reduction selectivity toward ethanol and acetate at very low cathodic bias ( $\sim 40\%$  and  $> 10\%$  Faradaic efficiency at  $-0.3$  V vs RHE for 0.1 M KOH respectively),<sup>109</sup> suggesting residual oxygen or solvent water to take part into CH<sub>3</sub>COO<sup>-</sup> formation.<sup>123,124</sup> A successive study on C<sup>16</sup>O reduction in a H<sub>2</sub><sup>18</sup>O electrolyte on crystalline copper facets confirmed that around 65%, 80%, and 70% of ethanol, acetate, and 1-propanol products had at least one oxygen from solvent molecules.<sup>125</sup> The presence of low amount of residual oxygen during CO<sub>2</sub> reduction conditions has been demonstrated by oxygen isotopic labelling ( $< 1\%$  of the original content within 100 nm).<sup>119</sup> Recently, an independent study has detected the presence of residual oxygen under reductive conditions through grazing incident hard X-ray Photoelectron Spectroscopy.<sup>126</sup> The authors identified those species as oxygen present at defects and/or vacancies on the surface and oxygen intercalated within metal layers. Arán-Ais *et al.* showed that C<sub>2+</sub> products formation correlates with the amount of surface defects and percentage of Cu(I) site during pulsed CO<sub>2</sub> electroreduction.<sup>14</sup> In particular, the extent of Cu(100) terraces promotes ethylene selectivity,

whilst Cu(I)-Cu<sup>0</sup> sites, together with defects, lead to higher ethanol production. An elegant assessment of Cu(I) reducibility through pulsed eCO<sub>2</sub>R at different temperatures further indicated that, when the kinetics of Cu(I) reduction is minimized ( $T = 278.14$  K), ethanol is the main C<sub>2</sub> reduction product to the detriment of ethylene formation (Figure 3a).<sup>127</sup> Finally, DiDomenico *et al.* proposed that pulsed CO<sub>2</sub> electrolysis may have two-fold impact on eCO<sub>2</sub>R: (1) “self-cleaning” of the catalysts from surface poisons (from  $E_{\text{anodic}} > -0.5$  V) and (2) enhanced C<sub>2+</sub> selectivity due to electrolyte/electric field effects (from  $E_{\text{anodic}} > -0.1$  V).<sup>128</sup>



**Figure 3:** **a**, Faradaic efficiencies of ethylene and ethanol for eCO<sub>2</sub>R in 0.1 M KHCO<sub>3</sub> on Cu at different cathodic bias and applied temperatures. Anodic pulse: +0.6 V vs RHE, timesteps for anodic and cathodic pulses: 1 s.<sup>127</sup> **b**, Product distribution on Cu, thin Cu<sub>2</sub>O, and thick Cu<sub>2</sub>O for CO<sub>2</sub> reduction in a 0.1 M K<sub>2</sub>CO<sub>3</sub> electrolyte at  $-1.0$  V vs RHE after 1 hour, 16 hours of continuous operation, and regeneration of the oxide layer by exposing the catalyst to air for two weeks after operation.<sup>129</sup> Figures adapted with permission from Ref. 127 (Copyright 2021 American Chemical Society) and Ref. 129 (Copyright 2021 National Academy of Sciences) respectively.

Recently, it was highlighted that almost no CO<sub>2</sub> reduction products are formed in absence of oxygen or surface defects.<sup>129,130</sup> After 16 hours of operation at  $-1.0$  V vs RHE in CO<sub>2</sub>-saturated 0.1 M K<sub>2</sub>CO<sub>3</sub> electrolyte, hydrogen evolution was the only electrocatalytic reaction observed on both pc-Cu and a Cu/Cu<sub>2</sub>O catalyst, unless oxygen was reintroduced in the system (Figure 3b). Since the authors demonstrated through *in situ* Raman spectroscopy that Cu<sub>2</sub>O was fully reduced after only 1 hour at  $-1.0$  V vs RHE, they highlighted the rel-

evance of disordered  $\text{Cu}^0$  for  $\text{C}_2\text{H}_4$  production. Additionally,  $\text{CO}_2$  reduction performance differs significantly depending on the pre-treatment of the catalyst, as shown in a recent work on well-defined  $\text{Cu}(100)$  and  $\text{Cu}(111)$  facets. These catalysts were prepared by pristine atomically flat Ultra-High Vacuum (UHV) specifically to reproduce Hori's results.<sup>35,36</sup> UHV-prepared, sputtered, electrochemically polished, and  $\text{O}_2$  plasma treated copper surfaces presented similar  $\text{eCO}_2\text{R}$  reduction current densities normalized by their electrochemically active surface area,<sup>130</sup> in line with a recent outlook.<sup>131</sup> Higher Faradaic efficiencies toward  $\text{eCO}_2\text{R}$  products on electrochemically polished and plasma treated samples were entirely due to suppression of  $\text{H}_2$  production,<sup>130</sup> expected to happen due to variations of surface pH.<sup>131</sup> Hydrocarbons were observed only for electrochemically polished or plasma-treated samples, thus surface defects were deemed responsible for tuning CO binding energy, crucial property to enable the C-C pathway. These features were observed by CO Temperature Programmed Desorption (TPD) analysis.

Briefly extending the discussion to OER catalysts, a pristine  $\text{SrIrO}_3$  catalyst performed better than  $\text{IrO}_x$  and  $\text{RuO}_x$  for OER due to strontium leaching during operation.<sup>13</sup> The resulting non-stoichiometric catalyst enables only 0.27-0.29 mV overpotential for 30 hours of continuous operation. X-ray Photoelectron Spectroscopy (XPS) on Ir 4*f* and Sr 3*d* observed a decrease of Sr signal to 25% of its original value after only 30 minutes and 20% after 24 hours. Thus, a novel structure was entirely formed *in situ* and became extremely active after 2 hours of electrochemical testing. Besides, the typically employed Ni materials reorganize extensively to form oxyhydroxy-compounds<sup>132-135</sup> which may be further doped by iron coming from nonpurified electrolytes.<sup>54,136-138</sup> Under operation, nickel catalysts evolve from  $\alpha\text{-Ni}(\text{OH})_2$  at low potentials to  $\gamma\text{-NiOOH}$  at high potentials as indicated by Ni K-edge X-Ray Absorption Spectroscopy (XAS) measurements.<sup>139</sup> In addition to the morphological evolution, the layer-layer stacking is typically enlarged by water insertion and all these structural changes lead to improvements in the reactivity of the active sites as well as the mass transport of the reaction intermediates. The complexity of the system increases when

Ni electrocatalysts are doped with Ir ions, since these dopants may coordinate with water molecules in solution, adsorb on the surface, and/or intercalate across the NiOOH lattice.<sup>54</sup>

### 3.2 Short-lived reaction intermediates

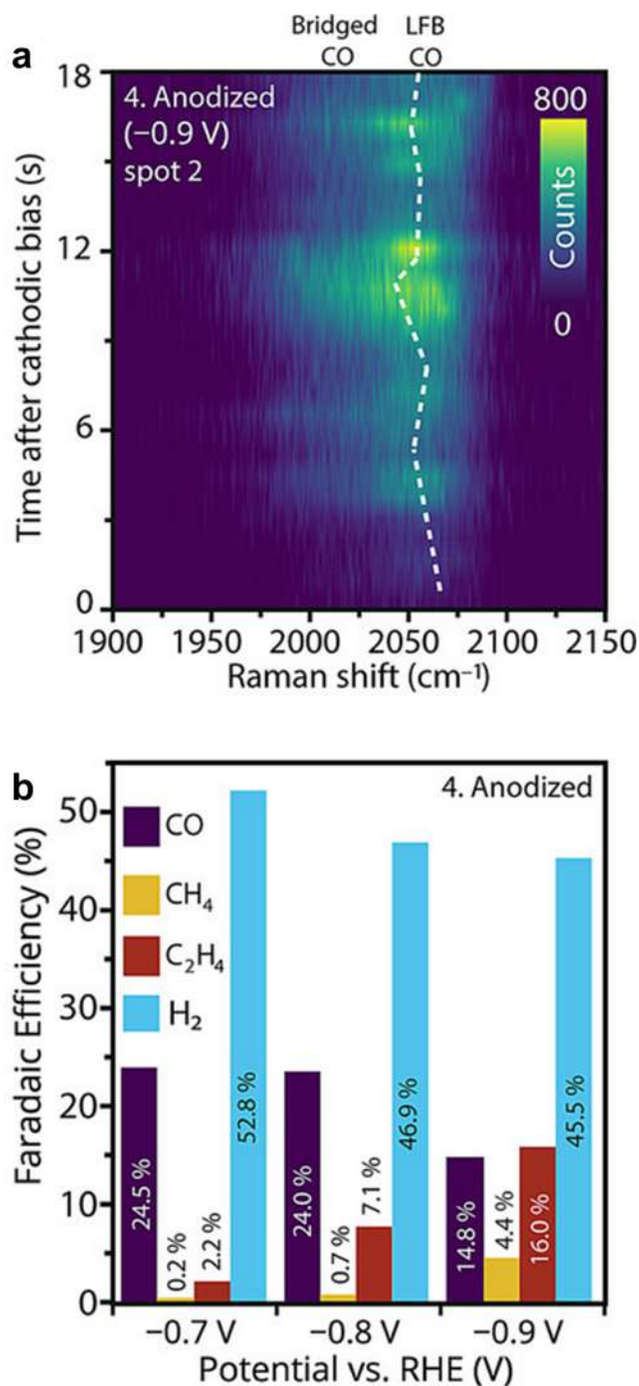
Many computational and experimental mechanistic studies have been dedicated to disentangle the key intermediates and elementary steps of catalytic reactions,<sup>9,37,140–142</sup> toward an accurate description of the reaction pathways for eCO<sub>2</sub>R reduction on transition metals.<sup>143,144</sup> Steady adsorbates have been successfully identified through spectroscopic techniques,<sup>145</sup> thus confirming theoretical predictions on adsorbed species.<sup>9</sup> However, short-lived intermediates formed during reaction pose a new challenge to both spectroscopy,<sup>146</sup> and computational modeling. These species are typically crucial for differentiating mechanistic routes and ruling activity, as in the case of OCCO<sup>-</sup> along the C<sub>2+</sub> route,<sup>16</sup> and their unequivocal identification needs an integrated approach involving experimental and theoretical characterization.<sup>16,147</sup>

As a clear proof of the relevance of short-lived intermediates, lately new features have been reported through *Operando* techniques.<sup>148–150</sup> Surface-selective infrared spectroscopy has been employed to sample CO adsorption configurations on copper.<sup>151</sup> By integrating the C=O vibrational bands associated to bridge adsorption site (1800–1900 cm<sup>-1</sup>) and top adsorption site (~ 2100 cm<sup>-1</sup>) at different cathodic bias, top-bound CO was indicated as the only active species for both eCO<sub>2</sub>R and CO reduction (COR). Besides, the authors observed that this specific adsorption configuration partially converts to unreactive bridge-bound CO below CO saturation regime.<sup>151</sup> An *et al.* further confirmed the presence of a reactive CO configuration *via* Time-Resolved Surface Enhanced Raman Spectroscopy (TR-SERS) for CO<sub>2</sub> reduction on copper nanoparticles. They revealed a highly dynamic, short-lived CO intermediate with low frequency C=O vibration (~ 2060 cm<sup>-1</sup>, Figure 4a), appearing at -1.0 V vs RHE, the sweet spot for ethylene formation (Figure 4b).<sup>152</sup> Concurrently, an independent work observed the increase in Cu-CO stretching band at the expense of Cu-

CO rotation band for high CO partial pressure and high negative potential and related this feature to C-C coupling.<sup>153</sup> Lately, two vibrational signals at  $\sim 490\text{ cm}^{-1}$  and  $\sim 530\text{ cm}^{-1}$  have been detected on Ag-decorated  $\text{Cu}_2\text{O}$  catalysts using *Operando* SERS for potentials more negative than  $-1.0\text{ V}$  vs RHE, with a concurrent increase in selectivity toward ethanol, 1-propanol, and acetaldehyde.<sup>154</sup> A proper assignment of these bands may shed light on the preferential selectivity of Ag-based catalysts toward oxygenates<sup>155,156</sup>

### 3.3 Electrolyte effects

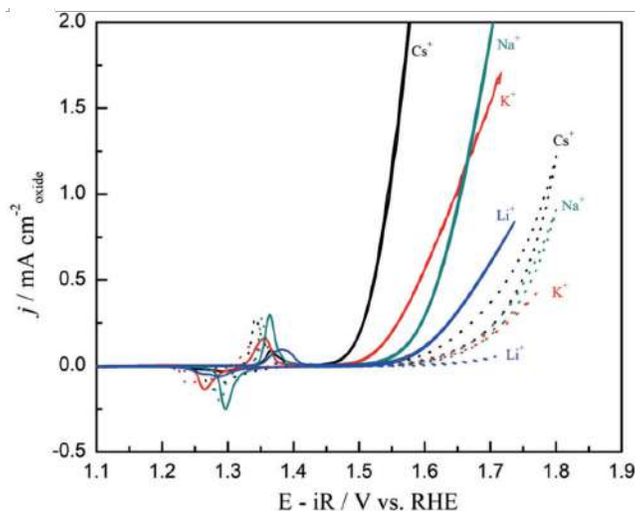
Electrolyte, as well as mass transfer of reaction intermediates, are crucial parameters for electrocatalytic reactions, as  $\text{CO}_2$  reduction,<sup>158,159</sup> hydrogen evolution,<sup>42,160,161</sup> and oxygen evolution,<sup>162</sup> since concentration profiles at the interface might severely depart from the bulk electrolyte composition.<sup>2,16,162–164</sup> For instance, alkaline pH at the surface is expected to promote  $\text{CO}_2$  reduction to  $\text{C}_{2+}$  to the detriment of  $\text{C}_1$  products.<sup>165–167</sup> Laser-microstructured cavities on copper allowed the local tuning of surface pH and  $\text{CO}_2$  concentrations, confirming high Faradaic efficiency (F.E.) toward ethanol, ethylene, and propanol for alkaline pH and intermediate  $\text{CO}_2$  concentration.<sup>167</sup> Further investigations with pulsed  $\text{CO}_2$  reduction proved a two-fold effect of anodic bias in lowering the surface pH and stabilizing and/or generating polarized sites.<sup>168</sup> Whilst for a anodic bias of  $+0.9\text{ V}$  vs RHE the authors observed a 10% enhancement of  $\text{C}_2$  selectivity with respect to the lack of anodic pulses, a more positive bias ( $E_{\text{anodic}} = +1.2\text{ V}$  vs RHE) resulted in increased  $\text{CH}_4$  production due to consumption of  $\text{OH}^-$  at the surface and a consequently more acidic pH.<sup>141</sup> Finally, ions in the electrolyte are also crucial for cathodic reactions as e $\text{CO}_2$ R and HER. Cations are expected to interact with reaction intermediates through local electric field effects,<sup>32,169–171</sup> or inducing water electrolysis from their solvation shell,<sup>172</sup> whilst anions are mainly responsible for surface poisoning due to specific adsorption.<sup>2,173–176</sup> Besides, the impact of ions is not only limited to transition metals, as demonstrated by observed cation effect for  $\text{CO}_2$  reduction on perovskite oxides.<sup>177</sup>



**Figure 4:** **a**, Time-Resolved Surface Enhanced Raman Spectroscopy (TR-SERS) heat map on anodized mechanically polished Cu at  $-0.9$  V vs RHE, showing a low-frequency band due to an highly dynamic CO intermediate. Raman spectra interval: 717 ms. **b**, Catalyst performance at different applied potential in a  $\text{CO}_2$ -saturated  $0.1$  M  $\text{KHCO}_3$ . Figures adapted with permission from Ref. 152 (Copyright 2021 Wiley-VCH under Creative Commons Attribution 4.0 International License<sup>157</sup>).

Whilst simulations on crystalline models successfully accounts for some of these phenomena, the assessment of the interplay of cation, surface pH, and mass transfer effects is so far limited. Recently, specific anion and cation contributions were employed in an empirical way to promote long-term stability for CO<sub>2</sub> reduction on continuous-flow electrolyzers.<sup>15</sup> Since flow cells undergo deactivation due to formation of salt at the cathode,<sup>176</sup> Endrődi *et al.* designed a novel *Operando* procedure, where fresh electrolyte solution (1.0 M CsOH) was introduced into the CO<sub>2</sub> feed every 12 hours to regenerate and activate the silver cathode. Through this process, they achieved partial current density for CO up to 420 mA cm<sup>-2</sup> for more than 200 hours. They attributed the enhanced performance of this proof-of-concept setup to physical interactions, such as removal of salt at the cathode from the fresh solution and promoting effects from the cations. Besides, benchmark studies of the reaction conditions indicated that CO Faradaic efficiency depended on bulk pH, peaking at pH = 13.6 ( $j_{\text{CO}} > 200 \text{ mA cm}^{-2}$ ), thus suggesting the concurrent influence of surface pH effects.<sup>178</sup> Very recently, a fundamental study demonstrated the triggering role of alkali metals in enabling CO<sub>2</sub> reduction on polycrystalline Au, Ag, and Cu in acidic media due to stabilization of a short-lived \*CO<sub>2</sub><sup>-</sup> intermediate.<sup>45</sup> Such evidence was reported independently for eCO<sub>2</sub> on a copper Gas Diffusion Electrode (GDE) configuration, even though the authors observed such effect only for nonacidic bulk pH.<sup>179</sup> In fact, different pH regimes lead to variations in cation concentration at the surface,<sup>180</sup> as rationalized by Goyal *et al* through fundamental experiments and theoretical insights.<sup>161</sup>

Cation effects extend beyond eCO<sub>2</sub>, since they have been identified as well for HER, ORR, and OER on different electrodes. Along the alkali group, as the ionic radius increases, higher reduction activities are typically expected (Figure 5), however the mechanism underlying this promotional phenomenon is widely debated. Gao *et al.* reported that the alkali-metal ions significantly influence the oxidation state of Mn in MnO<sub>x</sub> OER catalysts.<sup>182</sup> Specifically, Li<sup>+</sup> ion stabilizes the Mn<sup>2+</sup> species, which is OER-inactive, thus leading to low OER performance for small radius cations.<sup>182</sup> Instead, Garcia *et al.* postulated that the superior OER catalytic



**Figure 5:** Cyclic voltammograms of NiOOH at pH 13 during OER in different electrolytes. Solid lines: unpurified electrolytes; dotted lines: purified electrolytes. Measurements were performed under rotation of the working electrode (1600 rpm) and collected at  $0.010 \text{ V s}^{-1}$ .<sup>181</sup> Adapted with permission from Ref. 181 (Copyright 2019 Wiley-VCH).

performance of larger cations  $M^+$  is due to specific interactions with Ni-OO<sup>-</sup> species to form NiOO<sup>-</sup> ···  $M^+$  complexes, whose formation is increasingly stabilized as the cation radius becomes larger.<sup>181</sup>

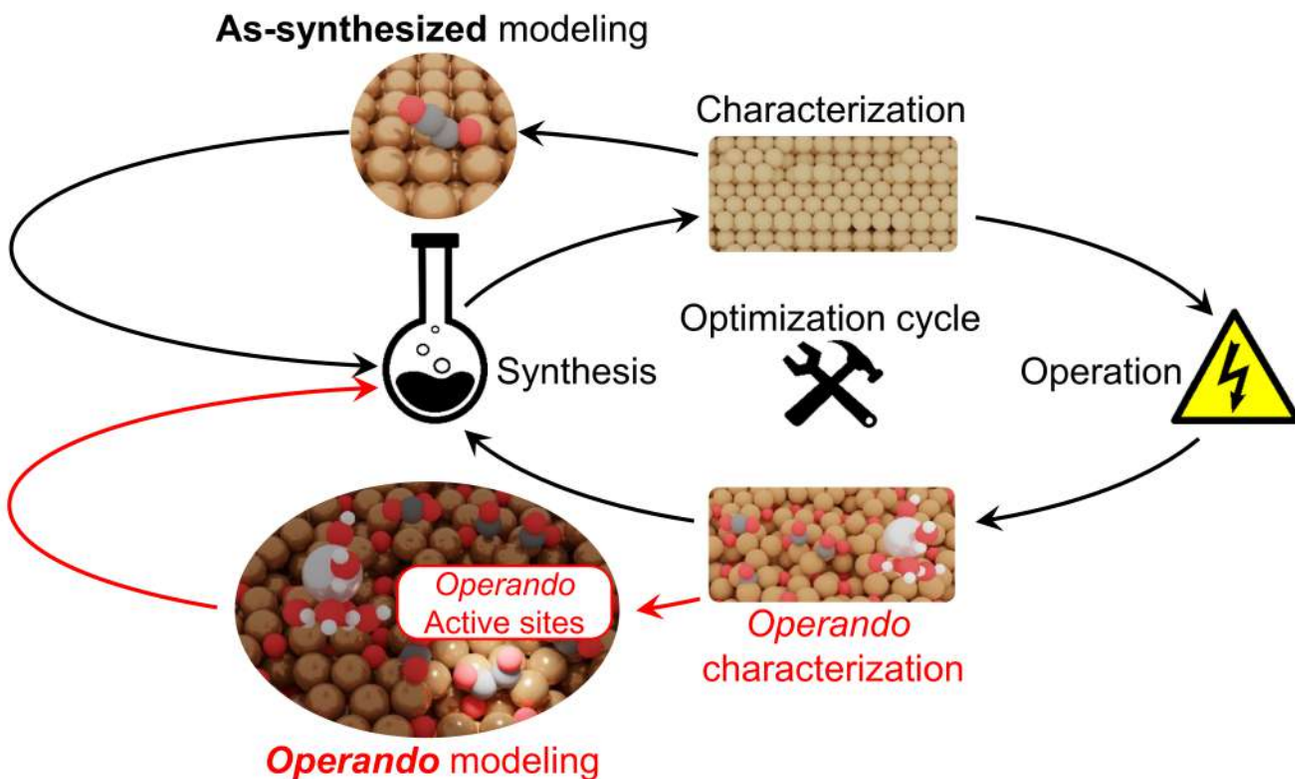
On top of cations, anions are relevant for OER as well, since they may accumulate at the anode and form complexes on the surface. Eventually, these species may lead to deactivation and/or degradation of the employed catalysts,<sup>162</sup> thus considerable effort has been devoted recently to tuning the chemical environment at the anode.<sup>183</sup> Typically, these anionic complexes generated on the surface are hard to identify since they evolve during operation, converting to other chemicals depending on surface pH and applied potential.<sup>184</sup> For instance, for a Pt(111) catalyst in sulfuric acid solution different anionic adsorbates have been proposed such as sulfate,<sup>185</sup> bisulfate,<sup>186</sup>  $\text{SO}_4^{2-} \cdots \text{H}_3\text{O}^+$  ion pairs,<sup>187</sup> or  $\text{OH}^-$ .<sup>188</sup> To further increase the complexity of these processes, mutual dependencies between cations and anions are also expected, since cations may act as counterions and stabilize anions species. García *et al.* proposed a similar mechanism for anions on Pt, suggesting a two layer morphology with chemisorbed (bi)sulfate on the Pt catalyst and adsorbed cations on the

(bi)sulfate layers.<sup>189</sup>

### 3.4 Perspective for *Operando modeling*

Recent experimental challenges summarized in Sections 3.1-3.3 cannot be fully rationalized through simulations on as-synthesized crystalline models,<sup>190</sup> which in brief we here define as “*as-synthesized*” modeling (Section 2.2). New approaches must be developed to reproduce electrocatalytic conditions, addressing the material gap and mass transport phenomena within the diffusion layer<sup>2</sup> toward the definition of an “*Operando*” modeling framework.<sup>123</sup> Predictions achieved through this new approach must then be validated by *ad hoc* experimental studies under well-defined conditions.<sup>190</sup>

This framework should aim at realistically assessing (1) electric field/adsorbate driven surface reconstruction, (2) evolution of active sites and consequent changes in catalyst reactivity, (3) electrolyte effects as cation/anion/solvation contributions to electrocatalytic reactions, (4) mass transfer, and (5) surface/bulk pH effects. These five challenging phenomena are certainly crucial in tuning the overall performance of electrocatalytic cells, thus in our opinion the field of computational modeling should tackle these challenges, possibly by integrating these processes into generalized, multiphysics models. While insights from computational studies on as-synthesized catalysts (in brief, *as-synthesized modeling*, Figure 6) are only partially valid under operation, “*Operando*” modeling can properly tackle challenges from *Operando* characterization. Thus, by assessing more “realistic” conditions, this novel framework could enable the identification of potential-, adsorbate-, and pH-dependent active sites and trigger a second step in the optimization cycle, Figure 6.



**Figure 6:** Catalyst optimization cycle and contribution from “*as-synthesized*” modeling and “*Operando*” modeling. Characterization of pristine systems and *Operando* tracking of structural changes and/or intermediates provide fundamental understanding of the role of the active sites. After two decades of modeling of as-synthesized catalysts, theoretical methods must now move toward the accurate modeling of the overall reaction environment during operation.

## 4 Modeling reconstruction under electrochemical conditions

Surface sensitivity, *i.e.* the contribution of surface morphology to catalytic activity and selectivity, has been extensively assessed in the field of catalysis,<sup>191</sup> leading to the development of Linear Scaling Relationships correlating adsorption energies of reaction intermediates and coordination number of the active sites (Section 2.2.4). Application of these concepts to crystalline domains suggest that undercoordinated sites bind adsorbates more strongly,<sup>83–85</sup> motivating both the high reactivity of defects for eCO<sub>2</sub>R reduction,<sup>14,129,192</sup> and the raise of competing HER for surface atoms with very low coordination number.<sup>117,193</sup>

Even though defined synthetic procedures and catalyst treatments have proved effective to increase the density of surface defects,<sup>112,194–197</sup> reconstruction phenomena occurring during operation have prevented the reproducibility and scalability of these approaches.<sup>63,102,198–200</sup> Additionally, poor modeling of reconstruction processes hinders the identification of active sites during operation and thus prevents the definition of guidelines to maximize these reaction centers. The material gap, important in thermal catalysis, is even more acute in the context of electrochemistry, as the potential and electrolyte are thermodynamic sinks to which the simulations are typically not coupled. Furthermore, thermodynamic considerations alone cannot reproduce local morphologies originated from kinetic processes,<sup>201</sup> such as metastable and kinetically trapped ensembles which might be extremely active in electrocatalysis.<sup>202–207</sup> A comprehensive investigation of kinetic effects driving surface restructuring through purely static DFT simulations is limited by the large configuration space of defective surfaces, thus such studies can be performed only under well-defined mechanistic hypothesis<sup>208</sup> or through machine learning (ML).<sup>209</sup> The ML strategy is promising, yet not fully realizable to date. Besides, for suitable application to electrocatalysis it should be generalized to include electrolyte species and adsorbates. As an ultimate consequence, experimental studies are not yet able to fully exploit theoretical insights from models based

only on thermodynamic contributions and crystalline materials.

In this section, we will discuss new methodologies to (1) assess the thermodynamics and kinetics of adsorbate-driven surface reconstruction on crystalline facets,<sup>100,208</sup> and (2) to reproduce metastable phases formed during catalytic reactions.<sup>54,210,211</sup> This second strategy is particularly relevant, since application of molecular dynamics enables the assessment of surface patterns which originate from non thermodynamic processes. Nevertheless, the robustness of such DFT-MD approach must be properly validated through *ad hoc* experiments, as Korpelin *et al.* have recently highlighted.<sup>212</sup> All in all, detailed computational characterization of these reconstructed surfaces allows to identify active sites responsible for improved catalytic performance as well as local effects which tune activity and selectivity (Sections 4.3-4.4).<sup>166,210,211,213</sup>

## 4.1 Restructuring processes on “as-synthesized” models

Copper reconstruction has been observed through different characterization techniques and under distinct conditions. Low-Energy Electron Diffraction (LEED) and Scanning Tunneling Microscope (STM) have detected surface restructuring under ultrahigh vacuum conditions in presence of O and H adsorbates,<sup>214–219</sup> or graphene overlayers.<sup>220</sup> Besides, this continuous process occurs as well under electrocatalytic conditions, as reported through *Operando* EC-STM for HER<sup>221,222</sup> and with electrolytes typical of CO<sub>2</sub> reduction.<sup>63,100,102,103,105,198,223,224</sup> Overall, cathodic bias alone or surface modifications *via* adsorption of H or C species are expected to be the main promoters of such reconstruction,<sup>224</sup> nevertheless at very negative potential specific cation adsorption may also play an active role.<sup>225,226</sup>

Researchers from the groups of Prof. Buonsanti and Prof. Marzari assessed the extent of morphological evolution in copper under CO<sub>2</sub> reduction conditions and described a potential-driven degradation mechanism due to CO and H adsorption *via* Grand-Potential density functional theory simulations.<sup>100</sup> Copper nanocubes with different sizes (16, 41 and 65 nm) were subjected to CO<sub>2</sub> electrolysis in a 0.1 M KHCO<sub>3</sub> electrolyte at  $-1.1$  V vs RHE.

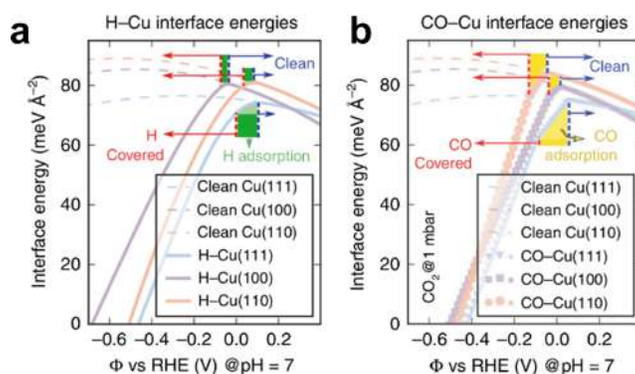
Although some cuboids were still observable after 6 hours, after 12 hours the nanoparticles underwent a full degradation of the crystalline facets, which caused a marked increase of HER. Nanoclustering was observed as well in a CO<sub>2</sub>-free electrolyte, thus suggesting the instrumental role of applied potential for enabling this phenomenon. To assess the influence of both adsorbates and applied potential, the authors employed DFT simulations on specific crystalline facets of copper, either clean or covered by H, CO, and H+CO with solvation included through a self-consistent polarizable continuum model.<sup>227</sup>

A grand canonical approach was applied to predict the potential-dependent equilibrium shape of nanoparticles under electrochemical conditions.<sup>228</sup> In case of adsorption of a proton, the dependence of the free energy of a catalytic system versus the potential  $U$  at the RHE scale is defined by Equation 13.  $\Delta G$  is the Gibbs free energy for hydrogen adsorption, corrected for entropic and zero-point energy contributions, obtained from DFT. The last term accounts for the variation of configurational entropy of the surface upon H adsorption. Contribution for the bulk pH is intrinsically included in the RHE dependence of the potential. The electrode potential  $U$  is obtained from the variation of electrostatic potential between vacuum and the Fermi energy of the metal, which was induced in the study by introducing opposite charges localized respectively on the slab and at 3 Å from the surface, and then correcting for the potential of zero charge. Hydrogen coverage  $\theta$  at equilibrium can be estimated as a function of applied potential  $U$  by solving Equation 13 for  $\Delta G_{\text{tot}} = 0$ . Consequently, the surface energy  $\gamma$  of a given system in presence of an adsorbed species is calculated from the electrocapillary equation knowing  $\theta$  and  $U$ . This method can be extrapolated to any species involving an electron transfer from the surface, i.e. CO.

$$\Delta G_{\text{tot}} = |e^-|U_{\text{RHE}} + \Delta G(\theta) + k_{\text{B}}T \ln \left( \frac{\theta}{1-\theta} \right) \quad (13)$$

This methodology reproduced the well-documented thermodynamic stability trend of copper facets without potential:  $\gamma_{110} > \gamma_{100} > \gamma_{111}$ , which implies that the formation of

(111) facet requires less energy than (100) and (110) cleaving. However, at cathodic bias, the adsorption of H and CO reduces the interface free energy, leading to negative surface energies for potential more negative than  $-0.5$  V vs RHE (Figure 7a-b). A negative surface energy implies instability of the given facet and its degradation or evolution toward a stable domain.<sup>229,230</sup> Thus, the authors proposed that surface dissolution starts at the edges of the nanocuboids and more negative potentials determine a stabilization of non cubic domains, *i.e.* (111), to the detriment of cubicity. The predicted triggering effect of negative enough applied potential to enable reconstruction was confirmed by experimental studies at more positive voltages ( $-0.7$  and  $-0.3$  V vs RHE), for which no significant degradation was observed.



**Figure 7: a-b**, Grand potential interface energies calculated for H-covered and CO-covered Cu surfaces in aqueous solution.<sup>100</sup> Adapted with permission from Ref. 100 (Copyright 2018 Springer Nature under Creative Commons Attribution 4.0 International License<sup>157</sup>).

In a following study in collaboration with our group,<sup>63</sup> the application of a graphene-layer on top of polycrystalline copper accounted for a different reconstruction process. After 4 hours, polarization at  $-1.0$  V vs SHE in a  $0.1$  M  $\text{KHCO}_3$  electrolyte, *Operando* EC-STM and Raman spectroscopy revealed the evolution of pc-Cu to copper nanocuboids.<sup>63</sup> Suggesting a minor, likely kinetic effect of reaction intermediates due to the protective graphene overlayer, the authors rationalized the nanostructuring process through sole electrostatic considerations. In the modeling section, we explicitly induced electric field on the chosen copper surfaces *via* a dipole correction,<sup>231,232</sup> as described in the DFT datasets generated and stored in the ioChem-BD database.<sup>233</sup> Thus, by modeling the electrical double layer as

a parallel plane capacitor under the assumptions of high polarization and large electrolyte concentration,<sup>2</sup> we derived a dependence of copper surface energy  $\gamma$  on applied electric field  $\vec{E}$  and solvent dielectric permittivity,  $\epsilon_r$ .<sup>63</sup> The intensity of  $\vec{E}$ , modulated by  $\epsilon_r$ , determines an increased electronic density at the surface, thus stabilizing the overall system. Open facets as Cu(100) and Cu(110) can better accommodate electronic density than (111) due to the lower coordination of their surface sites.<sup>232</sup> Thus, upon application of high cathodic bias, these domains are more stable than close-packed domains. Finally, surface morphologies with shorter radius of curvature experience a higher electric potential, thus boosting the reconstruction even further.

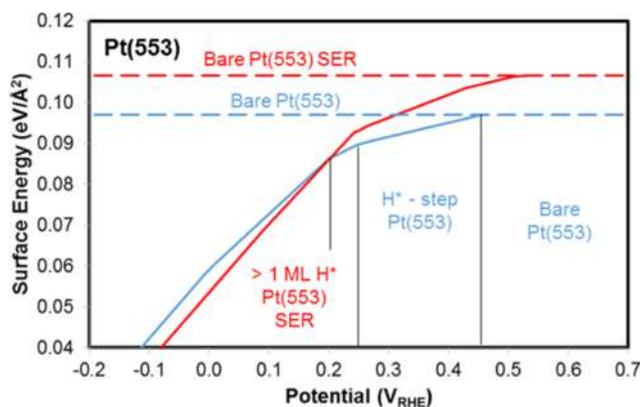
Analogous thermodynamic considerations can be applied as well to assess the stability of nanoparticles (NPs). According to the Wulff theorem, at standard conditions the most abundant facet on a nanoparticle is the one which presents the lowest surface energy.<sup>234</sup> Thus, by estimating surface energies of different crystalline domains, it is possible to represent nanoparticles through their Wulff construction.<sup>235</sup> However, under reaction conditions, Ostwald ripening and degradation mechanisms can occur along with the formation of complexes at the surface. The window of stability of nanoparticles can be estimated following the theoretical framework developed by Ouyang *et al.*<sup>236</sup> For instance, considering the case of Ni nanoparticles on N-doped supports under eCO<sub>2</sub>R conditions, the Gibbs free energy of disintegration  $\Delta G_{\text{NP}}^{\text{dis}}$  toward Ni(CO)<sub>*x*</sub> was computed depending on the nanoparticle radius, temperature, and pressure, Equation 14.<sup>237</sup>  $E_{\text{surf-}n\text{CO}}^{\text{f}}$  represents the energy of the Ni center on carbon support coordinated to *n* CO molecules,  $\Delta\mu$  is the excess chemical potential of CO,  $\Delta E'_{\text{NP}}(R)$  is the energy of the CO-covered nanoparticle, *R*, *T*, *p*, and *S* are respectively radius, temperature, pressure, and entropy. By comparing  $\Delta G_{\text{NP}}^{\text{dis}}$  with the energy of an isolated Ni adatom, Pršlja *et al.* concluded that at very high coverage ( $\theta_{\text{CO}} = 1.00$  ML) and close to standard conditions (*T* = 300 K, *p* = 0.1 mbar), Ni nanoparticles smaller than 4 nm naturally disintegrate into Ni(CO<sub>2</sub>) complexes.<sup>237,238</sup>

$$\Delta G_{\text{NP}}^{\text{dis}}(R, T, p) = E_{\text{surf-}n\text{CO}}^{\text{f}} - n \cdot \Delta \mu_{\text{CO}}(T, P) - \Delta E'_{\text{NP}}(R) - TS \quad (14)$$

Extending those concepts to HER, a similar thermodynamic approach was employed to elucidate the occurrence of the so-called “third hydrogen peak” in the cyclic voltammograms on Pt electrodes at +0.22 V vs RHE.<sup>208</sup> This feature has been linked to (110) steps sites, (111) terraces, or (110)-(2 × 1) domains through accurate studies on single crystal surfaces.<sup>239–241</sup> McCrum *et al.* calculated the surface energies of Pt(553) and Pt(533) characterized respectively by (111) terraces with (110) and (100) steps in presence of adsorbed hydrogen. The onset potentials for hydrogen adsorption on the (110) and (100) edges was calculated as +0.46 eV and +0.56 eV for Pt(553) and Pt(533) respectively. Overall, this process was more favorable on steps rather than (111) terraces due to the lower coordination number of these defects.<sup>83–85</sup> The authors observed that by applying more negative potential, hydrogen adsorption becomes thermodynamically favorable on the terraces as well and \*H coverage increases. Consequently, the surface energy of the overall system decreases and the gradient of such effect depends on the number of electrons transferred per unit area, *i.e.* on hydrogen coverage (Figure 8).<sup>208</sup> The surface energy  $\gamma$  of the system in presence of H adsorption was calculated through Equation 15.  $G_x$  are the Gibbs free energy of adsorbed hydrogen (\*H), clean surface (\*), and 1/2 hydrogen molecule ( $\frac{1}{2}\text{H}_2$ ),  $U$  is the applied potential vs RHE,  $U_{\text{pzc}}$  is the potential of zero charge,  $\vec{p}_x$  are the electric dipole moments for adsorbed hydrogen (\*H) and clean surface (\*),  $d$  is the thickness of the double layer, and  $A$  is the geometric area of the surface.

$$\gamma(U_{\text{RHE}}) = \gamma(U_{\text{RHE}} = 0 \text{ V}) + \frac{G_{*\text{H}} - G_* - G_{\frac{1}{2}\text{H}_2} + |e^-|U + |e^-|(U - U_{\text{pzc}}) \frac{\vec{p}_{*\text{H}} - \vec{p}_*}{d}}{A} \quad (15)$$

The surface energies of the roughened facets, labeled as Pt(553)-SER and Pt(533)-SER, decrease as well under more negative potential. These systems were modeled starting from



**Figure 8:** Surface energy of the pristine and roughened (denoted as SER) Pt(553) facet with (solid lines) and without (dashed lines)  $\text{H}^*$ . Changes in surface energies slope are due to variation in hydrogen coverage, as indicated as insets.<sup>208</sup> Adapted with permission from Ref. 208 (Copyright 2019 American Chemical Society).

a pristine surface, either Pt(553) or Pt(533), with 1/3 of the step atoms translated to the minimum energy adsorption site close by. Thus, this corner at the edges showed opposite crystalline facet than the pristine steps: (100) for Pt(553)-SER, whilst a (110) for Pt(533)-SER. Since the (100) corner has lower coordination than the pristine (110) step-edges sites, this site binds hydrogen stronger and thus Pt(553)-SER presents a lower surface energy than Pt(553) from +0.20 V vs RHE (Figure 8). Thus, below this critical bias, the pristine (533) facet naturally reconstructs to its roughened form, giving rise to the experimental peak observed at +0.22 V vs RHE. Instead, this process is not favorable on Pt(533), since the pristine (100) step-edges sites on Pt(533) adsorb H strongly than the reconstructed (110) corner, thus Pt(533) is more stable than Pt(533)-SER for any applied potential. Kinetic studies further confirmed the crucial role of hydrogen coverage to drive platinum reconstruction. The concerted translation of two neighboring Pt – the first to create the corner and the second to fill the vacancy created – is kinetically hindered by a high activation barrier of 1.75 eV in absence of hydrogen. Instead, at high  $\text{H}^*$  coverage such kinetic barrier decreases to 0.93 eV and the reaction is exergonic. Thus, H adsorption leads to a reduction of the kinetic barrier  $> 0.8$  eV, yielding rates observable experimentally.

## 4.2 Modeling *Operando* surface reconstruction

Metastable phases or kinetically trapped ensembles stand at the core of electrocatalytic reactivity.<sup>207</sup> Albeit their thermodynamic instability, configurational entropy or kinetic processes can locally stabilize these active sites,<sup>204,242,243</sup> leading to enhanced activity and selectivity.<sup>244,245</sup> For instance, stability and role of polarized sites is a crucial dilemma in the field of eCO<sub>2</sub>R, following the first observation by Kanan and Li of the enhanced activity of a Cu<sub>2</sub>O-derived catalyst.<sup>108</sup> Even though copper is expected to be fully metallic under cathodic bias,<sup>246–248</sup> many direct and indirect experimental evidences hint at the presence of either kinetically trapped or metastable active sites under reduction conditions.<sup>14,111,202,206,249–258</sup> Thus, further insights from theory are needed and DFT modeling on crystalline surfaces is unable to fully reproduce the experimental data.<sup>205,259,260</sup>

We report here two possible approaches to mimic the complexity of copper surfaces under eCO<sub>2</sub>R conditions, respectively (1) *ab initio* molecular dynamics (MD) modeling,<sup>261</sup> and (2) methodologies based on machine learning (ML) techniques (Section 8 ).<sup>262</sup> The outcomes from DFT-MD simulations must be properly validated by experimental evidences, since commonly employed thermostats may lead to non physical temperature variations and large temperature gradients within the simulation cells.<sup>212</sup> Instead, the ML strategy will likely play a determinant role in the future, as can be easily extended as well to the stability of other components of electrocatalytic devices, such as membranes for fuel cells for instance.<sup>263</sup> Besides, alternative methods can serve the scope, as effective medium theory<sup>264,265</sup> which was recently employed to mimic surface roughening on copper electrodes derived from cuprous oxide, phosphide, nitride, and sulfide precursors.<sup>266</sup> As a last remark to guide the reader, in this section we limit the discussion to methodology and structural properties, while we cover the insights on active sites and local polarization effects in Sections 4.3 and 4.4 respectively.

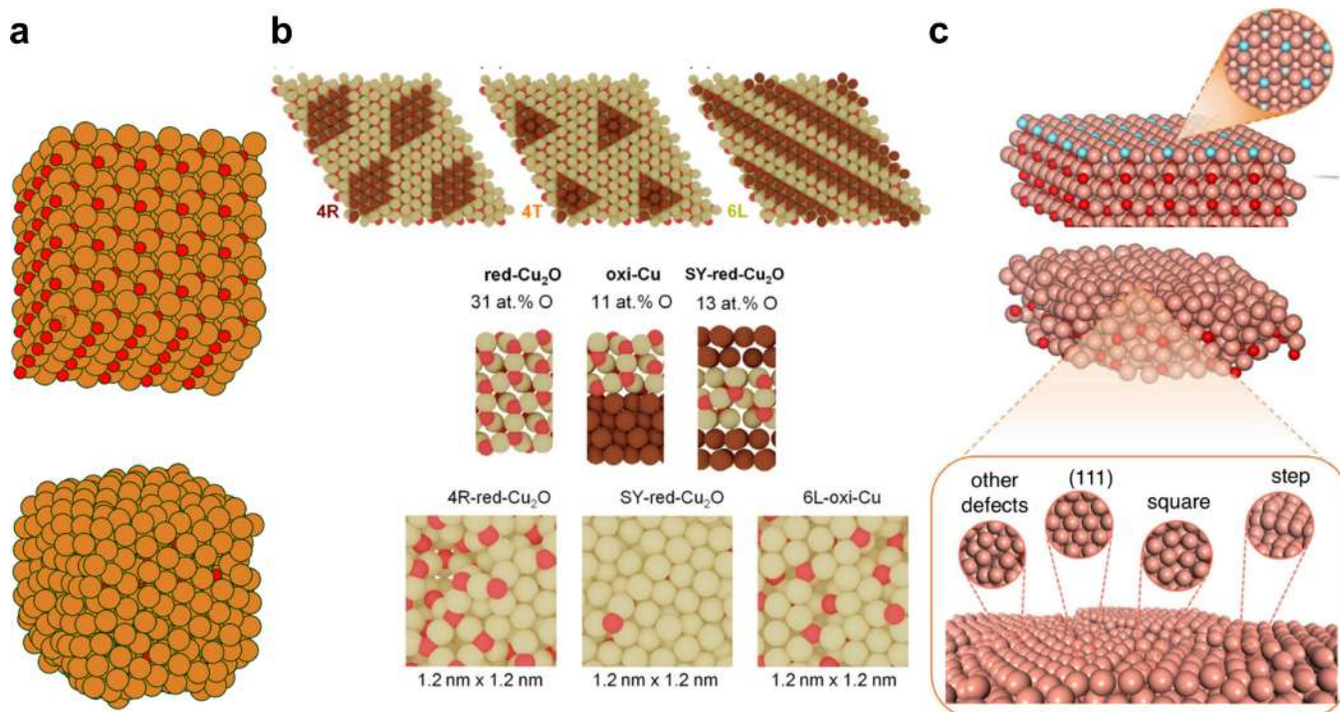
The first study which aimed at reproducing reconstruction on an oxide-derived model employed an *ab initio* molecular dynamics (AIMD) approach. Since subsurface oxygen was calculated to be unstable on a crystalline slab system in contrast with experimental obser-

vations,<sup>202,203</sup> Liu *et al.* assessed a disordered Cu<sub>2</sub>O-derived nanocube.<sup>205</sup> By employing a Self-Consistent Charge Density Functional Tight Binding scheme (SCC-DFTB),<sup>267</sup> they initially built a (5 × 5 × 5) Cu<sub>2</sub>O supercell accounting for 1.7 nm width and composed of 500 Cu atoms and 250 O atoms (Figure 9a). To simulate reduction processes occurring at cathodic bias, the surface oxygens were fully depleted. Besides, oxygen atoms within the two outermost layers were removed ten at a time, and after each removal a structural optimization was performed until no oxygen atoms were exposed. The resulting systems, which accounted for 500 Cu and 91 O, underwent a minima hopping algorithm with an initial NVE molecular dynamics temperature of 500 K for 50 cycles with 1 fs time step.<sup>268</sup> Oxygen diffusion from subsurface sites to the surface was tested within the SCC-DFTB framework, suggesting that such process is hindered on the distorted nanocube geometry. Migration of oxygens to the surface was possible only for edge and corner subsurface sites, however these configurations were endothermic, thus confirming the overall stability of subsurface oxygens.

Recently, we applied a similar strategy, albeit in a more systematic manner.<sup>210</sup> Since OD-Cu is usually synthesized either through oxidation of copper foils or electrochemical reduction of copper oxides,<sup>109,247</sup> we built two classes of model to mimic these regimes, labelled respectively red-Cu<sub>2</sub>O and oxi-Cu (Figure 9b). The red-Cu<sub>2</sub>O class was derived from a pristine (2√3 × 2√3)-R30° Cu<sub>2</sub>O(111) supercell with 2.1 nm lateral size. We fully depleted the outermost oxygen layer (12/144) and we partially removed subsurface oxygens (4-6/144). The elimination of subsurface oxygen was done following three geometric shapes, rhomboidal (R), triangular (T), and linear (L) to enhance clustering, pitting, and formation of grain boundaries. The three resulting *nS* systems were characterized by equivalent suboxide formation energy within 0.01 eV per Cu atom (*n* = number of subsurface oxygen removed; *S* = elimination motif). The simulated O content, 30-31 at.%, was higher than values reported experimentally (10-20%),<sup>111,112</sup> due to contribution from the bulk oxide. Besides, a 7 layers-thick symmetric slab (SY-red-Cu<sub>2</sub>O) was employed to model deep reduction conditions, after having depleted all the oxygens within the three outermost Cu<sub>2</sub>O layers

from each side (120/168, O content of 13 at.%, Figure 9b). Finally, systems mimicking copper oxidation were obtained by depositing three  $\text{Cu}_2\text{O}$  layer on a  $(5\sqrt{3} \times 5\sqrt{3})\text{-}R30^\circ$  Cu(111) matrix and removing surface and subsurface oxygens according to the protocol previously defined (11 at.% O, Figure 9b). Analogously, these three systems were isoenergetic within 0.01 eV. The seven simulated models underwent AIMD simulations with the PBE density functional (Ref. 269) for 10 ps at 700 K (3 fs time steps with a canonical ensemble, NVT, Nosé-Hoover thermostat).<sup>261,270</sup> The full AIMD trajectories are freely available on the open-source ioChem-BD database.<sup>271</sup> Benchmark tests with a Hubbard correction  $U_{\text{eff}} = U - J = 7 - 1$  eV on the  $d$  orbital of Cu,<sup>272-276</sup> and different AIMD temperatures (300 K, 500 K) confirmed the robustness of the computational setup. As a proof of agreement with experimental data, simulated STM images of the reconstructed surfaces resembled experimental observations on  $\text{Cu}_2\text{O}/\text{Cu}(111)$  materials under CO autocatalytic reduction.<sup>277</sup> Secondly, the overall atomic roughness for our models was significantly higher than crystalline Cu (0.8-1.4 Å vs 0.32 Å, respectively).<sup>210,278</sup> Finally, both simulated Cu  $2p$  and O  $1s$  features,<sup>279,280</sup> as well as Raman spectra,<sup>281</sup> agreed with the experimental reports.<sup>202,251,282-284</sup>

Previously, Huang *et al.* employed a different approach, which combined DFT with Grand Canonical techniques and an artificial Neural Network (NN) to investigate Cu nanoparticles.<sup>213</sup> As-synthesized nanoparticles cannot be explicitly model through DFT simulations due to computational limits, since 10 nm NP accounts for  $\sim 200000$  atoms and  $\sim 10000$  surface sites. Thus, the authors designed the copper nanoparticle through the reactive force field framework (ReaxFF) developed earlier,<sup>285-287</sup> and the semi-empirical embedded-atom model to model interactions between copper atoms.<sup>288</sup> To simulate chemical vapor deposition, copper atoms were subsequently introduced in the computational system at a rate of 3.2 atoms per ns for 30 ns. Later, 38 annealing cycles were simulated to optimize the structure. Each cycle consisted of a temperature ramp from 300 K till 1200 K (5 ps), 5 ps stabilization at 1200 K, cooling down to 300 K (5 ps). Finally, the resulting system was relaxed through ReaxFF for 20 ps at 300 K. Up to 400 random sites were chosen from the



**Figure 9:** Molecular dynamics-based approaches to reproduce surface reconstruction occurring for oxide-derived copper catalysts under operation. Depending on the oxygen content, different surface motifs are reproducible. **a**, OD-Cu nanocube containing 250 unit cells initially, then manually reduced to 500 Cu (brown) and 91 O (red) atoms prior to apply minima hopping.<sup>205</sup> **b**, Partially oxidic phases for O-lean surfaces.<sup>210</sup> Red and light/dark brown highlight oxygen and oxidic/metallic copper atoms, respectively. **c**, Defective Cu facets for O-free surfaces.<sup>211</sup> Red and brown atoms represent O and Cu, respectively, whilst light blue atoms highlight the surface oxygens initially removed. Adapted with permission from Ref. 205 (Copyright 2017 American Chemical Society), Ref. 210 (Copyright 2020 American Chemical Society), and Ref. 211 (Copyright 2021 Springer Nature under Creative Commons Attribution 4.0 International License<sup>157</sup>) respectively.

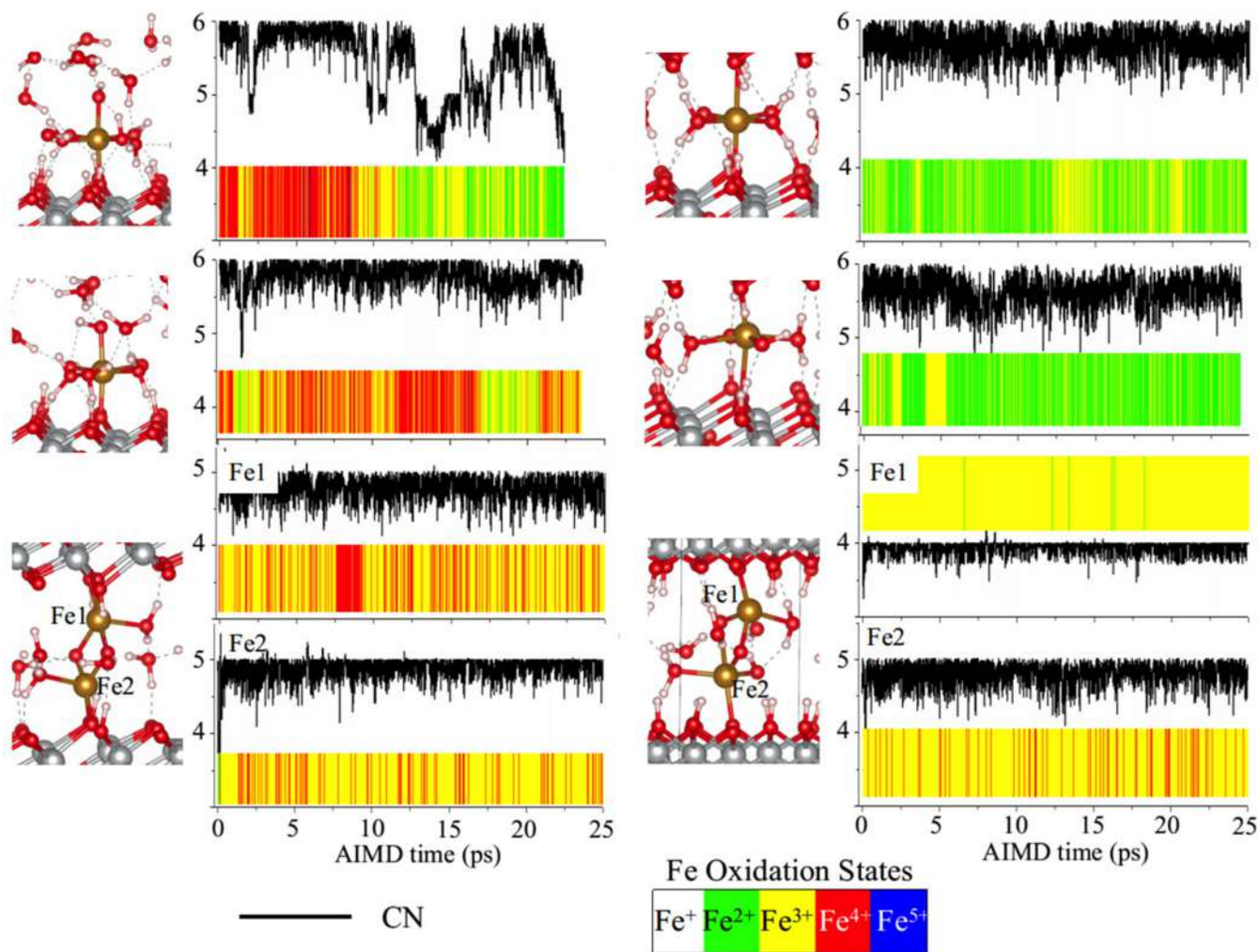
final structure and the local environment of each site up to 8 Å was integrated into a neural network. Each neighboring atom was transformed into a 12 two-body and 18 three-body molecular descriptors for a 2-layer NN with 50 nodes in each layer. The 400 sites were further partitioned into training, validation and test sets with an 8:1:1 ratio.

Later, Cheng *et al.* developed a framework integrating molecular dynamics and a neural network potential (NN-MD)<sup>289</sup> to assess the reconstruction occurring on a oxygen-lean copper material.<sup>211</sup> Initially, the whole surface oxygen layer was depleted from a pristine Cu<sub>2</sub>O(111), since previous studies suggested oxygens to be unstable at the electrode/electrolyte

interface under CO<sub>2</sub>R conditions (Figure 9c).<sup>247,290</sup> Then, the resulting system underwent NN-MD for 1 ns at 300 K to mimic the first stage of the reduction process. Since some of the subsurface oxygens migrated to the surface, O atoms with lower vacancy formation energy and coordination number 3 were removed, since they are expected to be the least stable during reduction. An additional 1 ns NN-MD was carried out and the previous step was repeated until no surface oxygen was present to derive six different OD-Cu models. The surface morphology largely evolved, accounting for (111) facets ( $\sim 53\%$ ), square domains ( $\sim 12\%$ ), steps ( $\sim 16\%$ ), and other defects ( $\sim 19\%$ ), see Figure 9c.<sup>211</sup>

The above mentioned methodologies are applicable as well to assess reconstruction processes for other electrocatalytic reactions, as the effect of potential dopants on catalytic performance for OER. For instance, in our group we have studied the formation of NiO<sub>x</sub>H<sub>y</sub> phases under OER conditions.<sup>54,291</sup> Based on computed Pourbaix diagrams, a NiO<sub>x</sub>H<sub>y</sub> catalyst is expected to evolve to the  $\gamma$ -Ni(OH)<sub>2</sub> phase when in contact with water solution. Besides, a successive transition to  $\gamma$ -NiOOH is envisioned when a positive potential is applied.<sup>54,292</sup> These phase transitions happen concurrently with water intercalation.<sup>293-295</sup> In fact, due to strong water adsorption and H-bonds, a NiOOH surface fully covered by water molecules results in a negative surface energy. This indicates that the water will intercalate between NiOOH layers, which agrees with the volume expansion observed in experimental studies.<sup>293-295</sup> When Fe is introduced in the system as dopant, the Fe<sup>2+</sup>/Fe<sup>3+</sup> ions may also penetrate into nickel layers further increasing layer-layer distance and active inner surface. Consequently, the surface of a Fe-doped NiO<sub>x</sub>H<sub>y</sub> electrode in water solution is expected to become gel-like under work conditions. Through AIMD simulations, we found that the adsorption of iron ions on the NiOOH surface is stable. However, we observed that the local chemical environment of Fe ions, such as the number of coordinated water molecules, is dynamic, as shown in Figure 10. Water molecules coordinated with iron ions frequently exchange their proton (coupled with electron) with oxygen at the catalytic surface. This behavior dynamically changes the oxidation state of Fe and further enables more flexible va-

lence state of Ni ions, from  $\text{Ni}^{3+}$  to  $\text{Ni}^{2+}$ , lowering the overall potential for oxygen evolution to 0.23 V.



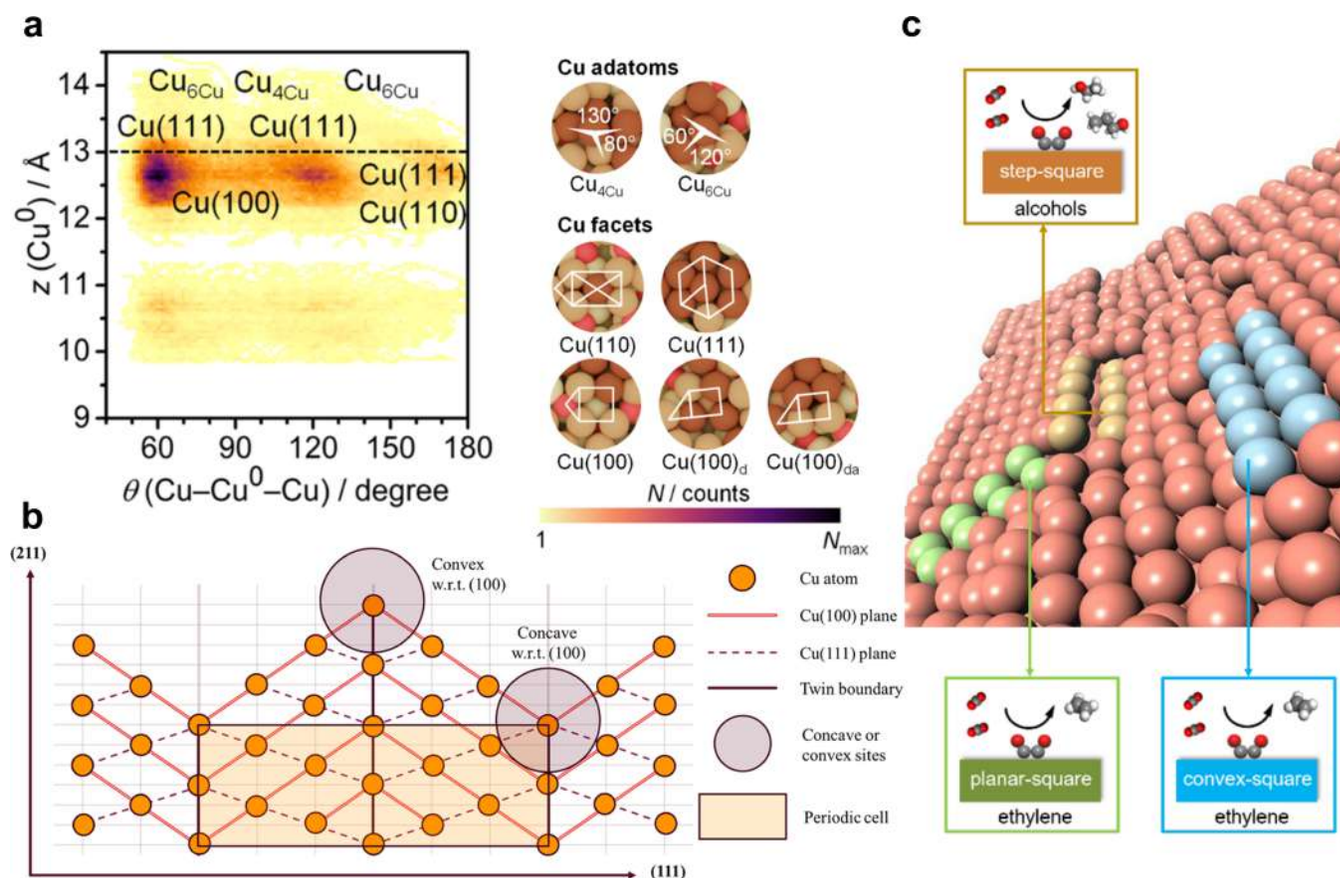
**Figure 10:** Evolution of coordination numbers (CNs) and oxidation states (OSs) of Fe ion adsorbed and/or intercalated for different Ni-based (oxy-)hydroxide models ( $\text{NiO}_x\text{H}_y$ ) during AIMD simulations.<sup>54,291</sup> Figures adapted with permission from Ref. 54 (Copyright 2020 American Chemical Society).

### 4.3 Identification of active sites

In addition to properly reproduce morphological changes at the surface, the methodologies described in Section 4.2 enable the identification of crucial surface ensembles and active sites responsible for the catalytic performance.<sup>205,210,211,213</sup>

The AIMD-based study on oxygen-lean  $\text{Cu}_2\text{O}$  catalysts (Figure 9a, Section 4.2) highlighted the presence of copper in three well-defined structural and electronic states: metallic  $\text{Cu}^0$ , suboxidic  $\text{Cu}^{\delta+}$ , and oxidic  $\text{Cu}^+$ , respectively coordinated with 0, 1, and 2 oxygen atoms.<sup>210</sup> Remarkably, suboxidic  $\text{Cu}_2\text{O}_{0.5}$  ensembles have been observed experimentally,<sup>206,255,256</sup> and reported with an average Cu-O coordination number of 1.1 estimated through EXAFS,<sup>256</sup> confirming theoretical predictions. While metallic and oxidic copper exhibited the typical oxidation states of bulk Cu and  $\text{Cu}_2\text{O}$ , suboxidic Cu accounted for a Bader charge between 0.1 and 0.4  $|e^-|$ , in line with experimental evidence of formation of polarized  $\text{Cu}^{\delta+}$  sites during thermal and electrochemical reactions.<sup>206,255,256</sup> Besides, all the three copper species were significantly undercoordinated if compared to surface atoms on crystalline facets and each oxygen coordination further lowered Cu-Cu coordination number by one unit. Thus, average  $\bar{N}_{\text{Cu-Cu}}$  of 4.9, 3.6, and 2.0 units for  $\text{Cu}^0$ ,  $\text{Cu}^{\delta+}$ , and  $\text{Cu}^+$  were estimated from these OD-Cu models, showing good agreement with experimental characterization of OD-Cu (respectively 6.6, 3.08, and 2.21).<sup>256,296</sup> The abundance of residual  $\text{Cu}^+$  species within the surface layers ranged between 10 and 40 at.% (for red- $\text{Cu}_2\text{O}$  and oxidic Cu classes respectively), in line with recent experimental observations on  $\text{Cu}_2\text{O}$  nanocubes at  $-0.95$  V vs RHE (20 at.%  $\text{Cu}^+$ )<sup>258</sup> and Cu(100) under pulsed  $\text{CO}_2$  electroreduction (7-11 at.% at  $-1$  V vs RHE).<sup>14</sup> Finally, sampling the occurrence of interatomic angles for a given central species ( $\text{Cu}^0$ ,  $\text{Cu}^{\delta+}$ ,  $\text{Cu}^+$ , and O), we detected 14 recurrent ensembles. Darker areas in the 2D histogram reported in Figure 11a correspond to higher density of similar structural morphology. By mapping recurrent angles with geometric patterns visible on the reconstructed surfaces, we characterized surface defects (4- and 6-coordinated copper adatoms) and crystalline reconstructions toward (100), (110), and (111) facets, either fully metallic or partially polarized. These observations closely agree with experimental reports on surface reconstruction toward open facets,<sup>63,100,102,223</sup> as well as high surface density of defects.<sup>14,112,195,199</sup> Finally, a remarkable  $\text{Cu}_3^{\delta+}\text{O}_3$  feature appeared at the surface, which was characterized in an independent experimental work.<sup>255</sup> As for oxygen species, subsurface

configurations adopted the bulk-like tetrahedral shapes,  $O_{4Cu,t}$  as well as possible distorted 3- and 5-fold coordinations,  $O_{3Cu,d}$  and  $O_{5Cu}$ . Remarkably, few oxygens reconstruct toward planar configurations near the surface,  $O_{3Cu,p}$  and  $O_{4Cu,p}$ , and may be the species found stable as a  $Cu_2O_{0.5}$  stoichiometry for a  $Cu(OH)_2$ -derived electrode under  $CO_2$  reduction conditions at  $-1.0$  V vs RHE.<sup>256</sup>



**Figure 11:** Relevant ensembles identified on OD-Cu and Cu models. **a**, Recurrent surface ensembles on O-lean OD-Cu structures (Figure 9a), defined by the local geometry of a given Cu cluster.<sup>210</sup> Oxygen atoms are shown in red, whilst copper is depicted in brown, from dark ( $Cu^0$ ) to light ( $Cu^+$ ) depending on its oxidation state. **b**, Theoretically-designed FCC copper containing (100) and (111) planes, as well as twin boundaries.<sup>213</sup> **c**, On a O-free OD-Cu surface (Figure 9c), step-square (dark yellow), planar-square (green), and convex-square (light blue) domains were characterized and related to alcohols and ethylene formation.<sup>211</sup> Adapted with permission from Ref. 210 (Copyright 2020 American Chemical Society), Ref. 213 (Copyright 2018 American Chemical Society), and Ref. 211 (Copyright 2021 Springer Nature under Creative Commons Attribution 4.0 International License<sup>157</sup>) respectively.

To assess a small sample of their computationally designed copper nanoparticle (Section

4.2),<sup>213</sup> scientists from Prof. Goddard III’s group selected 400 surface sites and calculated CO binding energy on these active species through their reactive force field framework, ReaxFF (Refs. 285–287). Further, they extrapolated the corresponding values for all the 10000 surface sites of their original NP *via* the NN trained during the study. As a result, they identified specific active sites with stronger binding than low crystalline facets, *i.e.* (100) and (211), in line with previous TPD experiments on copper.<sup>297</sup> Then, applying the formation energy of the protonated CO dimer, \*OCCOH, as a potential descriptor for C<sub>2+</sub> selectivity, they observed that square motifs similar to (100) accounted for favorable binding of this intermediate. Assessing the local geometry of the favorable square domains, a specific (100) terrace coupled to (111) was reported to further enhance this process, in agreement with experimental observations on the promoting effect of Cu(S)[*n*(100) × (111)].<sup>35,36,298</sup> As a final guideline to catalytic design synthesis, the authors proposed a periodic FCC copper structure with high surface density of this (111)-(100) motif as a promising active ensemble for increasing C<sub>2+</sub> formation (Figure 11b), since it showed the most favorable formation energy for OCCOH (0.35 eV and 0.41 eV either on convex or concave sites). This devised catalyst was predicted to grant 97% F.E. toward multi-carbon products, assuming a linear correlation between density of twin boundaries and C<sub>2+</sub> selectivity and no influence on HER.

Identification of the active sites on the O-lean OD-Cu developed with the NN-MD methodology (Figure 9c, Section 4.2)<sup>211</sup> and correlation with eCO<sub>2</sub>R selectivity was achieved following a reductionist approach alike the previous study.<sup>213</sup> The authors selected 155 surfaces sites out of the reconstructed slabs and calculated \*CO and \*COCO binding energy for these active centers *via* the NN potential.<sup>289,299</sup> The robustness of the NN potential was confirmed by benchmark tests through single-points DFT simulations, which returned low systematic offset of 0.08 eV and 0.13 eV for  $\Delta E^*_{\text{CO}}$  and  $\Delta E^*_{\text{COCO}}$  respectively. The CO dimer was chosen since it is generally assumed as the RDS toward formation of multicarbon products on copper.<sup>16</sup> In line with the results from Huang *et al.*,<sup>213</sup> square motifs exhibited the strongest \*COCO binding energy, accounting for an average energy cost for C-C

coupling of 1.27 eV. This conclusion agrees with an earlier theoretical study by Bagger *et al.*<sup>39</sup> and experimental insights from Hori *et al.* (Section 2.1).<sup>35,36</sup> By further differentiating between the squared domains, Cheng *et al.* characterized four surface motifs: planar-square (p-sq), step-square (s-sq), concave-square (cc-sq), and convex-square (cv-sq) facets, which were extracted as slab models to calculate the energetics of CO dimerization through accurate DFT simulations. All structures allowed for a less endothermic C-C coupling in comparison to (111) and (211) facets. Besides, p-sq, s-sq, and cv-sq sites permitted a facile kinetics, with an activation barrier lower than 0.7 eV vs 0.85/1.19 eV on the crystalline domains. C<sub>2+</sub> selectivity of these ensembles was mapped according to their \*CH<sub>2</sub>CHO adsorption energy, since this adsorbate is expected to be the intermediate ruling competition between ethanol and ethylene.<sup>16,143,300</sup> The authors observed that planar and convex squares promote the cleavage of the C-O toward ethylene, whilst step-square resembling Cu(S)-[*n*(111) × (100)] motifs inhibit the hydrocarbon pathway (Figure 11c), in good agreement with previous experimental and theoretical suggestions.<sup>35,36,39</sup> The different surface reactivity was rationalized through considerations on the local morphology. p-sq and cv-sq can stabilize \*O after C-O bond breaking on 4-fold sites, whilst the s-sq cleaving presents the lowest coordination number,<sup>83,83,84</sup> thus the strongest binding of the acetaldehyde intermediate which enables its further protonation. Additionally, C-O bond length was employed as descriptor for ethylene and ethanol selectivities. As expected, a shorter bond length is representative of a stronger bond, which consequently hinders C-O breaking, favoring formation of C<sub>2</sub>H<sub>5</sub>OH. Finally, the authors extended their theoretical framework by comparing performance of experimentally annealed OD-Cu with models obtained with different AIMD temperature ramps. Remarkably, C<sub>2+</sub> current density correlated with the atomic ratio of the three identified ensembles (p-sq, cv-sq, s-sq) estimated for the AIMD-annealed slabs. eCO<sub>2</sub>R activity toward ethylene and alcohols exhibited a linear dependence on (p-sq+cv-sq) and (s-sq) atomic contents respectively, confirming the selectivity of step-square domains toward ethanol and planar/concave-square morphology toward C<sub>2</sub>H<sub>4</sub>. However, according

to both theoretical and experimental observations, the highest density of active sites, and consequently highest performance of the catalyst, was achieved for the non-annealed Cu<sub>2</sub>O catalyst, whilst thermal treatments worsened both parameters.<sup>211</sup>

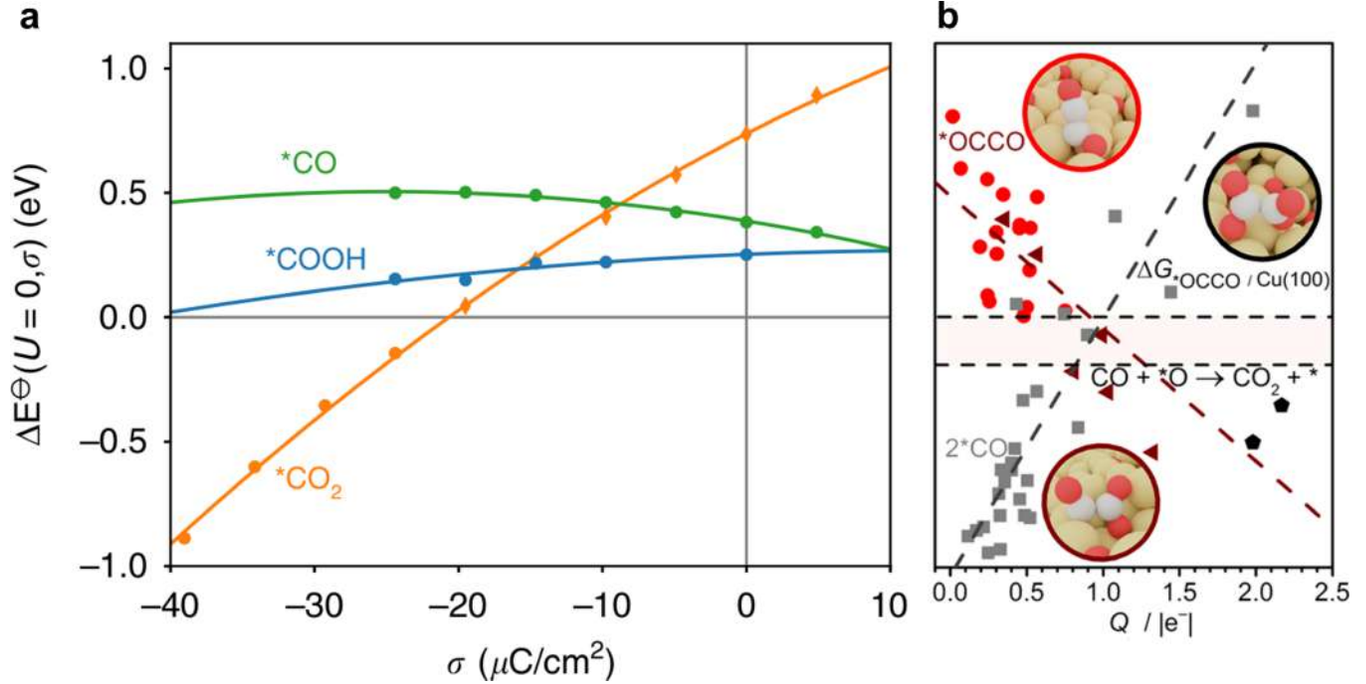
## 4.4 Surface polarization effects

As demonstrated through the *d*-band model,<sup>74,75</sup> surface reactivity of transition metals depends on their *d*-band center calculated with respect to their Fermi energy, *i.e.* a proxy of the degree of filling of the *d*-band (Section 2.2). In general, the capability of a catalytic center to bind intermediates is driven by its local electronic density. For cathodic reactions, low-coordinated surface defects exhibit stronger binding of the reaction intermediates due to their higher electronic density, resulting in tLSR based on average coordination numbers (Section 2.2.4).<sup>83-85</sup> In fact, the electronic density at the surface correlates with the effective potential, *i.e.* applied potential vs the potential at the point of zero charge (PZC).<sup>2,63,171,208</sup> Besides, undercoordinated sites are locally more charged due to their higher local curvature, as predicted by Green's theorem.<sup>63,301</sup> In addition to local variations of surface electronic density, the applied potential leads to the build-up of the electrochemical double layer, so that local electrostatic interactions can further stabilize or destabilize reaction intermediates.<sup>44</sup>

According to the Computational Hydrogen Electrode,<sup>8</sup> electric field/adsorbate electric dipole interaction can be estimated by a  $\vec{p} \cdot \vec{E}_{\text{EDL}}$  term which is added to the adsorption energy of the intermediate (Section 2.2.2). Following this concept, average electronic density at the surface, proxy of the effective electric field across the EDL, is a key factor to tune formation energies of specific compounds, as shown in Figure 12a for \*CO, \*COOH, and \*CO<sub>2</sub> adsorption during CO<sub>2</sub> reduction.<sup>166</sup> On top of these average fields effect, local electrostatic phenomena are expected to play a crucial role in triggering catalytic activity for eCO<sub>2</sub>R on otherwise inactive catalysts.<sup>42,45,97</sup> In fact, recent theoretical and experimental observations pointed out that dopants may further tune the local polarization of surface sites, so that limitations from tLSR can be overcome due to the formation of active sites with significantly

different electronic properties. For instance, we rationalized the enhanced eCO<sub>2</sub>R selectivity of S-doped copper toward formate through local electronic effects.<sup>302,303</sup> As an adatom, since S is more electronegative than Cu, it withdraws electronic density from the neighboring copper sites and thus acts as a tethering center to adsorb either CO<sub>2</sub> or a hydride. Then, adsorbed CO<sub>2</sub> selectively reduces to formate upon protonation of the C atom, whilst the \*H pathway undergoes an Heyrovský-like mechanism to the same product (Section 5.2, Figure 15). The negative charges localized on S is instrumental to enable both processes and to limit unselective eCO<sub>2</sub>R pathways on neighboring copper sites. Instead, positively charged surface sites are expected to influence reactivity in an opposite manner. Sn<sup>δ+</sup> sites in Cu-Sn alloys were predicted to weaken \*COOH and \*CO binding energy, thus leading to enhanced selectivity toward formate and CO depending on Sn/Cu ratio.<sup>304-306</sup> Cu<sup>+</sup>-Cu<sup>0</sup> pair have been deemed responsible for the enhanced C<sub>2+</sub> selectivity of OD-Cu materials.<sup>307,308</sup> Recently, a joint experimental and theoretical study employed TPD, *in situ* Attenuated Total Reflection Fourier Transform Infrared spectroscopy (ATR-FTIR), and DFT to suggest that strongly bound, bridge-adsorbed CO (CO<sub>B</sub>) is a precursor for methane formation, whilst a Cu<sup>+</sup> site accounts for weaker binding of linearly adsorbed CO, enabling CO dimerization to ethylene.<sup>257</sup> A similar weakening effect of polarized sites on binding energy was reported by Ismail *et al.*,<sup>309,310</sup> on a Au/Pb interface, where subsurface oxygen dopants hindered poisoning of the Pb surface, thus allowing formation of methane. An analogous electrostatic repulsion between positively charged surface atoms and adsorbates has been mentioned for a In-Pd solid solution, where In substituents weaken \*H and \*CO binding energy due to their partial positive charge, +0.64 |e<sup>-</sup>| estimated through Bader charge analysis.<sup>311,312</sup> Finally, similar tuning of eCO<sub>2</sub>R selectivity due to surface polarization were proposed for nickel oxygenates.<sup>97</sup>

Recently, we have systematically assessed the influence of polarized sites on CO<sub>2</sub> reduction activity and selectivity on the AIMD-derived OD-Cu models discussed previously (Figure 9a, Section 4.2).<sup>210,313</sup> We observed Linear Scaling Relationships of \*CO<sub>2</sub>, \*OCCO, and 2\*CO



**Figure 12:** **a**, Formation energies for  $^*\text{CO}_2$  (orange),  $^*\text{COOH}$  (blue), and  $^*\text{CO}$  (green) on Au versus the surface charge density, taking  $\text{CO}_2$ ,  $\text{H}_2\text{O}$ , and clean metallic surface as energy reference.<sup>166</sup> **b**,  $2^*\text{CO}$  and  $^*\text{OCCO}$  formation energies as a function of surface polarization  $Q$ . An asymmetric OCCO dimer on metallic copper, deprotonated glyoxylate, and oxalate intermediate are reported as red, dark red, and black insets respectively.  $Q$  is defined as the sum of the absolute Bader charges of the active sites, Equation 17.<sup>210</sup> Adapted with permission from Ref. 166 (Copyright 2020 Springer Nature under Creative Commons Attribution 4.0 International License<sup>157</sup>) and Ref. 210 (Copyright 2020 American Chemical Society).

formation Gibbs free energies vs surface polarization  $Q$  (Equation 16, Figure 12b).  $Q$  was obtained as the sum of the absolute values of the Bader charges  $q$  of the active sites ( $X$ ) binding O or C, Equation 17. Overall,  $\Delta G_{^*\text{CO}_2}$ ,  $\Delta G_{^*\text{OCCO}}$ , and  $\Delta G_{2^*\text{CO}}$  exhibited different dependencies on  $Q$ , with  $\beta < 0$  for intermediates adsorbed *via* a  $\eta_{\text{C},\text{O}}^2$  configuration ( $^*\text{CO}_2$ ,  $^*\text{OCCO}$ ) and  $\beta > 0$  for adsorbates solely bound through the terminal carbons ( $2^*\text{CO}$ ).

$$\Delta G = \alpha + \beta Q \quad (16)$$

$$Q = \frac{1}{N_{X(\text{C})}} \left| \sum_{i=1}^{N_{X(\text{C})}} q_{X_i(\text{C})} \right| + \frac{1}{N_{Y(\text{O})}} \left| \sum_{i=1}^{N_{Y(\text{O})}} q_{Y_i(\text{O})} \right| \quad (17)$$

Overall, surface polarization strengthens the binding of  $\eta_{\text{C},\text{O}}^2$  configurations, whilst it leads

to a weaker binding of C-bound compounds. Following these principles,  $^*\text{CO}_2$  adsorption is increasingly stabilized by surface polarization, thus suggesting a triggering role of electrode charging for the enhanced eCO<sub>2</sub>R activity of OD-Cu materials. Eventually, highly negatively charged sites such as near-surface oxygens ( $\text{O}_{\text{ns}}$ ) can combine with strongly bound  $^*\text{CO}_2$  adsorbates to form carbonate species, which limit the overall activity of the system through surface poisoning.<sup>121</sup> On the contrary, adsorption of CO molecules as separate entities on the surface is destabilized by increased surface polarization, whilst  $^*\text{OCCO}$  formation is increasingly promoted. Mildly polarized active sites, such as a  $\text{O}_{\text{ns}}$ -Cu couple, makes C-C coupling thermoneutral, thus enabling the possible formation of a deprotonated glyoxylate,<sup>314</sup> characterized by a low kinetic barrier (0.53 eV) (dark red inset in Figure 12b). This compound is a well-known intermediate for prebiotic CO<sub>2</sub> reduction,<sup>315</sup> although it has never been reported for the electrocatalytic counterpart.<sup>146</sup> Finally, high surface polarization characteristic of a  $\text{O}_{\text{ns}}$  pair can lead to formation of oxalates at the surface, which passivate the catalytic layer similarly as carbonate and may cause copper dissolution.

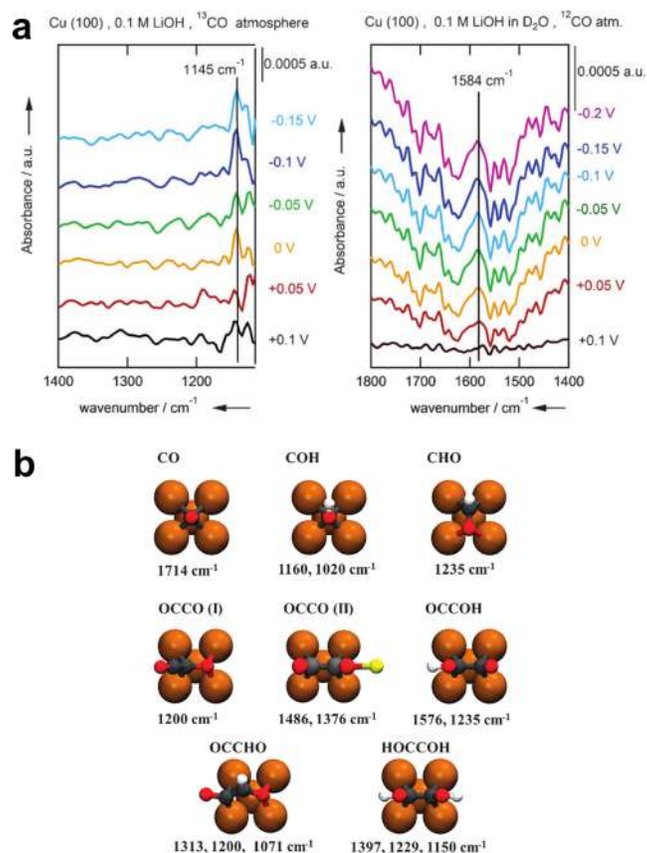
## 5 Reaction Pathways

The application of *Operando* techniques has evolved exponentially,<sup>150,316</sup> providing theoreticians with new challenges (Section 3.2), as only a direct mapping to computational structures allows for real insights into active sites and intermediates. Whilst initially spectroelectrochemical techniques could only shed light on catalytic properties under well-defined conditions or long-lived adsorbates on the surface, new advances have allowed to detect crucial, yet metastable reaction intermediates,<sup>147–149,151</sup> and to define complex reaction pathways and mechanisms.<sup>151–153,317</sup> In the incoming section, we review three recent works which highlight how systematic integration of spectroscopic and computational studies enabled the rationalization of the eCO<sub>2</sub> reaction mechanism (Section 5.1). Later, we describe the reaction routes for CO<sub>2</sub> reduction, hydrogen evolution, and oxygen evolution reactions (Sections 5.2-5.3.)

## 5.1 Spectroscopy-assisted identification of key reaction intermediates

A key step for deciphering the CO<sub>2</sub> reduction pathway puzzle was the identification of the RDS for C<sub>2+</sub> formation, the OCCO<sup>-</sup> intermediate.<sup>16</sup> As observed by Hori *et al.*,<sup>37</sup> CO<sub>2</sub> (CO) reduction to ethylene is pH-independent, thus the rate-determining step for C<sub>2</sub>H<sub>4</sub> formation must involve an electron transfer (Section 2.1). Since methane production is instead pH dependent,<sup>37,141</sup> the reaction pathways toward these two hydrocarbons must bifurcate at early stages. Whilst several studies suggested the formation of a negatively charged CO dimer as the RDS,<sup>140,300</sup> only a joint spectroscopic/DFT study could undeniably demonstrate the existence of this intermediate on the surface.<sup>147</sup> The authors observed two vibrational bands on Cu(100) in a <sup>13</sup>CO-saturated 0.1 M LiOH/D<sub>2</sub>O electrolyte from +0.1 to -0.15 V vs RHE, 1191 cm<sup>-1</sup> and 1584 cm<sup>-1</sup>, through *in situ* Fourier transform infrared (FTIR) spectroscopy, see Figure 13a. Transmission spectra of potential CO reduction intermediates as formaldehyde, formate, methanol, acetaldehyde, and acetic acid did not show compatible features, thus neglecting the assignment of these bands to other C-containing species. Finally, DFT-based vibrational analysis was crucial in identifying C-O-H and C=O stretching modes of a hydrogenated CO dimer (OCCOH) as responsible for the features observed (Figure 13b).

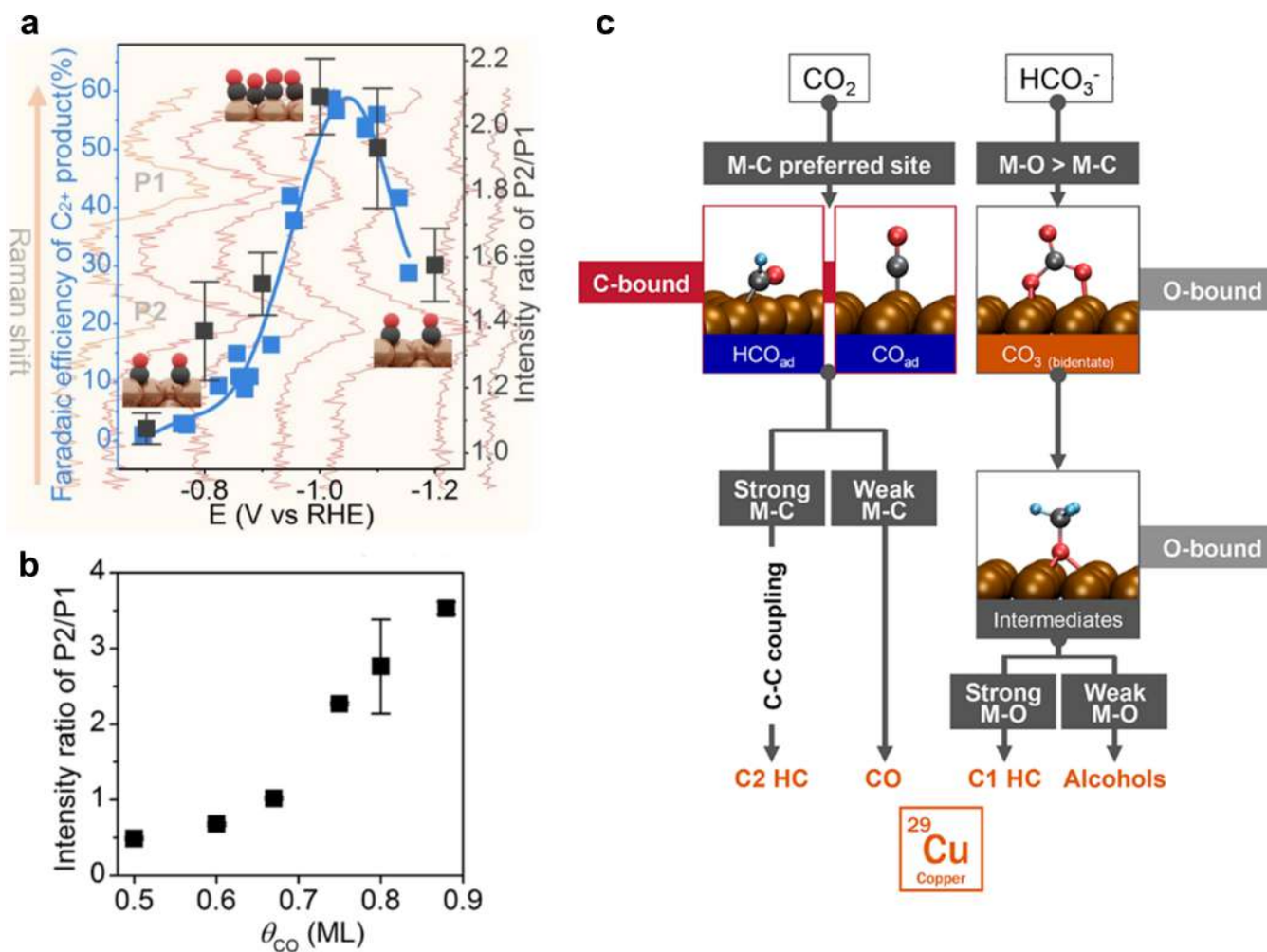
DFT vibrational analysis and *Operando* characterization can be further employed to provide mechanistic insights. In a recent collaboration with our group, computational modeling correctly reproduced experimental observations from *Operando* Surface Enhanced Raman Spectroscopy on CO<sub>2</sub> reduction on Cu<sub>2</sub>O nanocubes. As introduced in Section 3.2, scientists from Prof. Roldán-Cuenya’s research group observed a remarkable correlation between C<sub>2</sub>H<sub>4</sub> selectivity and intensity ratio between Cu-CO stretching and CO rotation bands (Figure 14a).<sup>153</sup> By assessing different CO surface coverages on Cu *via* DFT and estimating the related CO vibrational frequencies, we demonstrated a correlation between computationally-estimated intensity ratio and theoretical CO surface coverage (Figure 14b),<sup>318</sup> in line with



**Figure 13:** C-C coupling intermediate during CO reduction on Cu(100) observed and identified *via in situ* FTIR spectroscopy and DFT-based vibrational analysis. **a**, Spectra recorded on Cu(100) for a  $^{13}\text{CO}$ -saturated  $\text{D}_2\text{O}$  electrolyte in 0.1M LiOH exhibits a  $1145\text{ cm}^{-1}$  and  $1584\text{ cm}^{-1}$  assigned to  $^{13}\text{C-OH}$  and  $^{12}\text{C=O}$  stretching bands. **b**, Among  $\text{C}_1$  and  $\text{C}_2$  intermediates, a single(double)-hydrogenated CO-CO dimer presents analogous DFT vibrational frequencies as observed experimentally.<sup>147</sup> Adapted with permission from Ref. 147 (Copyright 2017 Wiley-VCH).

experimental observations for different CO concentrations in the electrolyte. Furthermore, the more intense Cu-CO stretching band was attributed to weakly-bond CO adsorbed on top sites at high coverages, which showed the lowest activation barrier for CO-CO dimerization.

By assigning spectroscopic signals to adsorbed species on different catalysts, DFT-based vibrational analysis can define complex reaction pathways and descriptors for catalyst performance. Vibrational bands for  $\text{CO}_2$  reduction on Cu, Au, and Pt were detected through *in situ* Surface Enhanced Infrared Absorption Spectroscopy (SEIRAS) at different applied potentials and later attributed to key reaction intermediates.<sup>317</sup> The authors observed a sin-



**Figure 14:** Unveiling reaction mechanisms and key intermediates with joint investigations involving density functional theory and *Operando* SERS. **a**, eCO<sub>2</sub> reduction to ethylene on Cu<sub>2</sub>O-derived catalysts exhibits a peak at around  $U = -1.0$  V vs RHE analogously to the ratio between Cu-CO stretching (P2) and CO rotation (P1) bands. **b**, Comparable P2/P1 ratio are reproduced *via* DFT for CO adsorption on Cu(100) at high coverage, suggesting a coverage driven C-C coupling mechanism.<sup>153</sup> **c**, eCO<sub>2</sub> reduction reaction mechanism on Cu defined through a joint *in situ* SERS/DFT-vibrational study.<sup>317</sup> Adapted with permission from Ref. 153 (Copyright 2021 American Chemical Society) and Ref. 317 (Copyright 2019 American Chemical Society).

gle signal for CO<sub>2</sub> reduction on Au,  $\sim 2100$  cm<sup>-1</sup>, which DFT ascribed to linearly-adsorbed CO. In fact only CO was produced at low cathodic bias (F.E. > 50% at  $-0.3$  V vs RHE). Four vibrational bands were instead reported on Pt: 2074 and 1831 cm<sup>-1</sup> for  $U < +0.2$  V vs RHE); 1620 and 1376 cm<sup>-1</sup> for anodic bias. DFT analysis attributed these signals to linearly-bond CO, CO on a bridge site, adsorbed COOH, and bidentate bicarbonate, respec-

tively. No eCO<sub>2</sub>R reaction products were detected, in line with CO poisoning expected for Pt.<sup>7</sup> Finally on Cu, adsorbed species varied depending on the applied potential. Bidentate carbonate was indicated responsible for the band at 1495 cm<sup>-1</sup> between +0.3 V vs RHE and -0.2 V vs. RHE, in good agreement with DFT calculations on Cu(100) and Cu(111) (1510 cm<sup>-1</sup> and 1533 cm<sup>-1</sup> respectively). O-bound intermediates such as \*OCH<sub>3</sub> (1390 cm<sup>-1</sup>), \*OCH<sub>2</sub>CH<sub>3</sub> (1340 cm<sup>-1</sup>), and bidentate \*HCOO (1240 cm<sup>-1</sup>) were observed from +0.1 V vs RHE until -0.9 V vs RHE. Besides, vibrational features for \*CHO (1477 cm<sup>-1</sup>) and \*CO (~ 1900 cm<sup>-1</sup>) were detectable from -0.3 V vs RHE and -0.6 V vs RHE, respectively. With regard to the catalytic performance, mainly CO was formed at -0.3 V vs RHE (F.E. = 30%), whilst CO<sub>2</sub> reduction to formate (F.E. ~ 15%) and ethylene (F.E. ~ 20%) become relevant from -0.9 V vs RHE at expenses of CO. Since only \*CHO and \*CO were detected within this potential windows, these intermediates were deemed responsible for C-C dimerization on Cu. Finally, the authors rationalized Pt, Au, and Cu reactivity assuming the strength of M-O and M-C bonds (*M* = metal) as descriptor (Figure 14c). In the case of copper, sites with stronger C affinity preferentially catalyze C-C coupling, whilst a weaker C binding allows for CO production. Conversely, sites which absorb eCO<sub>2</sub>R intermediates *via* a terminal oxygen are selective toward C<sub>1</sub> products or alcohols in case of strong or weak M-O bond, respectively.

## 5.2 Mechanism for CO<sub>2</sub> reduction and hydrogen evolution

From very early reports by Hori *et al.*, CO has been considered as the first key intermediate toward formation of C<sub>1</sub>-C<sub>2</sub> hydrocarbons such as methane and ethylene (Section 2.1).<sup>33,319</sup> Thus, electrochemical CO<sub>2</sub> reduction to CO is expected to occur either *via* (1) a proton-coupled electron transfer (PCET) to form a \*COOH species or (2) an electron-transfer (ET) toward a metastable \*CO<sub>2</sub><sup>-</sup> adsorbate, followed by a quick proton transfer (PT) toward \*COOH, see Figure 15a)<sup>9,143</sup> The preferential pathway depends on the given catalytic system.<sup>143,320</sup> Weak to mild \*CO binding elements as Cu, Ag, Au have been demonstrated to

favor the  $^*\text{CO}_2^-$  route,<sup>45</sup> while strong binding elements as Pt and Pd and single atom catalysts may prefer the  $^*\text{COOH}$  pathway.<sup>320</sup> As evidence of the  $^*\text{CO}_2^-$  route, experimentally  $\text{eCO}_2\text{R}$  activity was reported to be pH-independent (vs SHE) on mercury and gold.<sup>321–323</sup> High  $\text{CO}_2$  reduction activity was observed on Hg in media with low proton availability (dimethylformamide)<sup>321</sup>, while CO partial current density was reported to be independent from bicarbonate and proton concentration and first order on  $\text{CO}_2$  pressure on Au.<sup>323</sup>

Generalizing these evidences to other catalysts, the formation of adsorbed  $^*\text{CO}_2^-$  intermediate has been hypothesized as the rate-determining step for electrochemical  $\text{CO}_2$  reduction on transition metals,<sup>16</sup> while  $^*\text{COOH}$  formation has been proposed to limit  $\text{eCO}_2\text{R}$  activity on single atom catalysts.<sup>320</sup> Interestingly, recent works shows that alkali metal cations play a significant role along the  $\text{eCO}_2\text{RR}$  pathway.<sup>44,45,170,171</sup> Specifically, Koper and co-workers observed that no  $\text{CO}_2$  reduction products were catalyzed on Cu, Ag, and Au without an alkali cation in solution,<sup>45</sup> thus suggesting that explicit cation- $\text{CO}_2$  interactions are needed to stabilize adsorbed  $\text{CO}_2^-$  on these metals. If instead  $^*\text{COOH}$  formation *via* PCET from  $\text{CO}_2$  were to be the first step in  $\text{eCO}_2\text{R}$ , as generally assumed in studies employing the CHE framework,<sup>8,9</sup> then cation stabilization should be less relevant and  $\text{CO}_2$  reduction should take place at large overpotentials even without cation.<sup>171</sup>

In addition to  $^*\text{CO}_2^-$  and  $^*\text{COOH}$  formation, recently Chan and co-workers proposed  $^*\text{CO}_2$  adsorption prior to ET as the RDS for  $\text{eCO}_2\text{R}$  on transition metals,<sup>166,320</sup> since a previous study suggested quick electron transfer kinetics for  $\text{CO}_2$  activation on Au. In this work,<sup>324</sup> Nørskov, Chan, and coworkers described a reaction scheme involving an initial physisorption of the  $\text{CO}_2$  intermediate and a successive electron transfer only at the transition state for  $\text{CO}_2$  activation, confirmed by Bader charge for activated  $\text{CO}_2$  of  $-0.35 |e^-|$ . The authors then indicated average electric field/ $\text{CO}_2$  dipole interaction as crucial to facilitate  $\text{CO}_2$  adsorption.<sup>44</sup> Since no proton transfer is involved in this step, this reaction mechanism fulfills the experimental observation of no pH dependence of  $\text{CO}_2\text{R}$  rate vs SHE. As a short remark on this study, we observe that  $\text{CO}_2$  dipole depends on  $\text{CO}_2$  activation angles, thus

such dipole stabilization effect may become relevant only after CO<sub>2</sub> activation (bending) due to electron transfer. New studies should carefully assess this aspect, to verify whether average field/dipole interactions are sufficient to stabilize \*CO<sub>2</sub> before its activation<sup>171,324</sup> or alternative promoters, as cations, are keys.<sup>44,45</sup>

Following the reaction route toward C<sub>1</sub> (from center to left in Figure 15a), the \*COOH formed from CO<sub>2</sub> *via* PCET or ET + PT then undergoes a successive proton-coupled electron transfer to form adsorbed \*CO. Elements with weak \*CO binding, as Au, Ag, and Zn, can release this adsorbate in solution, thus leading to high selectivity toward CO. Instead, for mild to strong \*CO binding catalysts, the CO adsorbate can evolve toward other C<sub>1</sub> reduction products, such as methanol and methane, *via* successive proton-coupled electron transfers (Figure 15a).<sup>143</sup> The pH dependence (in SHE scale) of methane formation was first observed by Hori *et al.* for CO reduction on polycrystalline copper,<sup>37</sup> and then confirmed for both CO and CO<sub>2</sub> reduction on Cu(100) and Cu(111) by Schouten *et al.*<sup>141</sup> While methane selectivity is remarkable on copper for cathodic bias more negative than -1.0 V vs RHE, methanol formation on pure Cu is limited to Faradaic efficiencies lower than 1%.<sup>7,106</sup> Both products form either from a \*CHO or \*COH precursor via 5 and 3 proton-coupled electron transfers respectively (Figure 15a).<sup>143</sup> Since the pioneering work of Peterson *et al.*,<sup>9,325</sup> density functional theory simulations have reported the former intermediate (\*CHO) to be thermodynamically more stable, however recent kinetic analysis have indicated that methane formation occurs through the COH\* pathway.<sup>326</sup> Generally, Cu(111) has been regarded as selective toward methane<sup>141</sup> due to an effective stabilization of key intermediates on fcc sites.<sup>39,257</sup> In fact, \*COH/\*CHO and further reduced species accommodate nicely on hollow sites,<sup>39</sup> which can effectively donate electronic density to the adsorbate.<sup>257</sup>

Formate (or formic acid depending on bulk pH) is the last major C<sub>1</sub> product observed during eCO<sub>2</sub>R.<sup>106</sup> This chemical is expected to originate either from (1) coupling of a hydride with CO<sub>2</sub> *via* sequential PCET and ET,<sup>16,143,302</sup> or a (2) \*CO<sub>2</sub><sup>-</sup> intermediate adsorbed in a η<sub>O,O</sub><sup>2</sup> configuration *via* ET (see central mechanism in Figure 15a).<sup>141</sup> Alternately, the second

mechanism may proceed as well directly through a PCET to  $^*\text{OCHO}$ .<sup>327</sup> Strong  $^*\text{H}/^*\text{CO}$  binding metals as Pt and Pd favors the first route, while weak binding elements as Sn and In exhibit endergonic  $^*\text{H}$  binding, thus favoring the  $^*\text{OCO}^-/^*\text{OCHO}$  route.<sup>11</sup> Then, the hydride-mediated pathway continues with an Heyrovský-step, where the  $^*\text{H}^-$  adsorbate couple with a  $\text{CO}_2$  in solution leading to  $\text{HCOO}^-$  (Figure 15a). On the other hand, the  $^*\text{OCO}^-$  gets protonated to formate (Figure 15a).<sup>327</sup> Evolution of  $\text{HCOOH}$  depends on bulk pH and may occur as well as a chemical step in solution due to acid/base reactions.<sup>302</sup>

$\text{eCO}_2\text{R}$  to multicarbon products to date is limited to copper-based catalysts and is assumed to occur from CO-CO dimerization *via* ET (Figure 15b),<sup>16</sup> since the rate-determining step to form these products is pH-independent (Section 2.1).<sup>37,141</sup> The peculiar ability of copper to catalyze this reaction has been attributed to its mild CO binding, which prevents CO poisoning yet allowing the coupling reaction under fairly negative reduction potentials.<sup>11</sup> CO-CO dimerization takes place at high CO coverages and from activated CO adsorbed on atop sites, as demonstrated by suppression of CO rotation band<sup>153</sup> and dynamic C=O stretching band lower than  $2060\text{ cm}^{-1}$ .<sup>152</sup> The sluggish kinetics of C-C coupling ( $E_a > 1\text{ eV}$ )<sup>153</sup> should in principle involve poor  $\text{C}_2$  formation rates, while in contrast high rates are reported experimentally.<sup>12</sup> Under experimental conditions, cations may stabilize the  $^*\text{OCCO}^-$  dimer<sup>328</sup> and possibly facilitate the electron transfer, thus motivating this discrepancy between experiments and simulations.<sup>45</sup> In fact, short-lived adsorbates with vibrational frequencies attributable through DFT analysis to hydrogenated dimer ( $^*\text{OCCOH}$ ) or cation-coordinated dimer were spectroscopically observed under CO reduction on Cu(100), see Section 5.1.<sup>147</sup> In addition to the CO-CO dimerization mechanism, other C-C coupling precursors have been proposed, such as  $^*\text{CO-CH}$ ,<sup>155</sup>  $^*\text{CO-}^*\text{CH}_2$ ,<sup>37</sup>  $^*\text{CO-}^*\text{CHO}$ ,<sup>144,329</sup>  $^*\text{CO-}^*\text{CHO}$ ,<sup>65,329</sup> and  $^*\text{C-CO}$ .<sup>326</sup> However none of these species have been observed experimentally for  $\text{eCO}_2\text{R}$  on copper.<sup>5,16,330</sup> These alternative C-C steps may be relevant instead for different catalysts and CO coverage regimes.<sup>155,331</sup> Recently, Buonsanti, Calle-Vallejo and co-workers proposed a  $^*\text{CO-}^*\text{CH}$  coupling step on mixed Cu-Ag nanoparticles to motivate their enhanced selec-

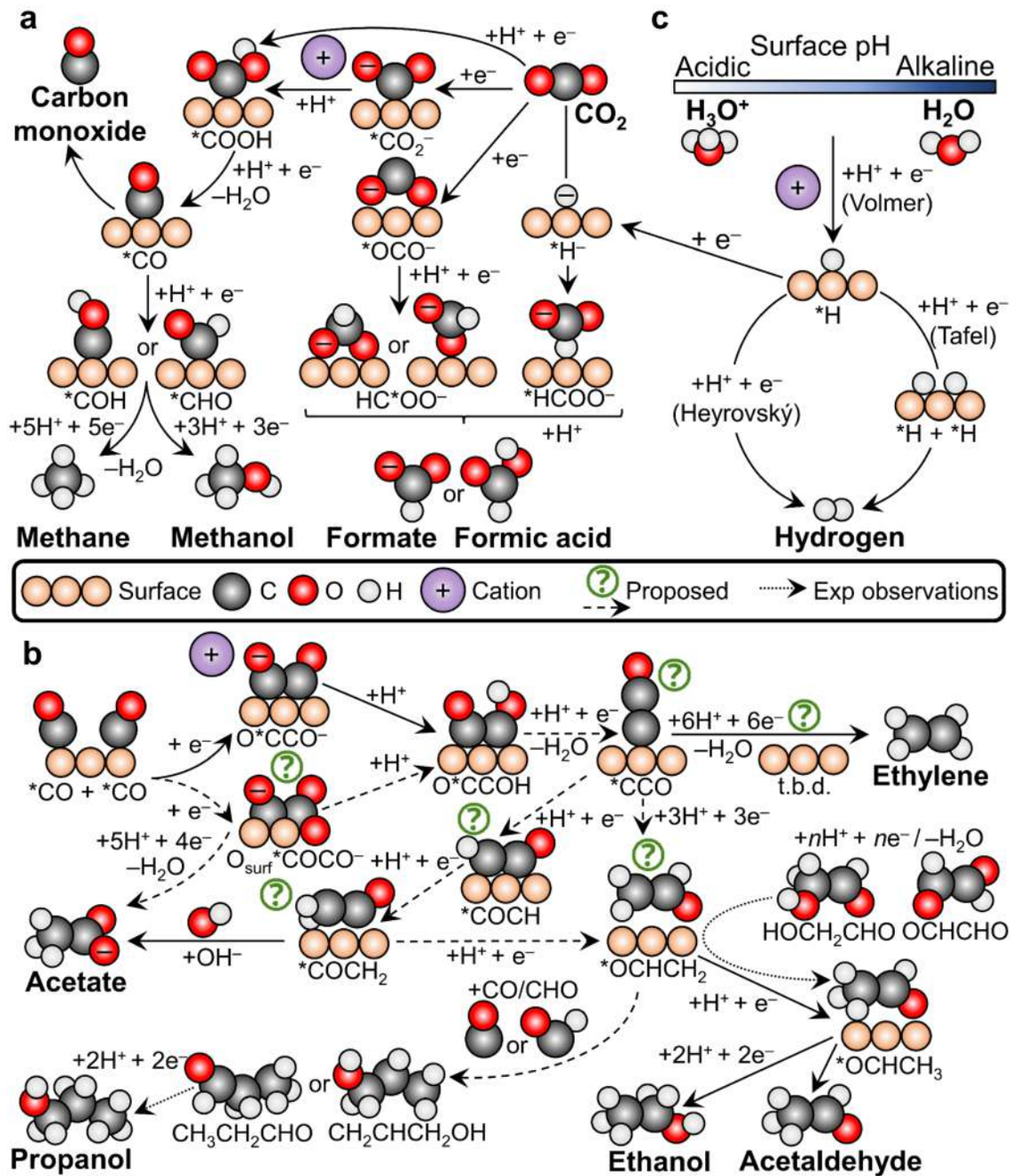
tivity to ethanol.<sup>155</sup> In line with this reaction route, Chang *et al.* employed CO adsorption isotherms and electrokinetic data to demonstrate that under low CO coverage typical of COR, CO-CO is unlikely to be the rate-determining step toward C<sub>2+</sub> formation.<sup>331</sup>

Following the pathways toward multicarbon species, the transient OCCO<sup>-</sup> gets quickly protonated to \*COCO<sub>H</sub>, which then undergoes successive proton-coupled electron transfers toward ethylene, ethanol, and *n*-propanol, the main C<sub>2+</sub> products (Figure 15b).<sup>106,143</sup> Spectroscopic techniques have not managed yet to detect the short-lived intermediates linking \*COCO<sub>H</sub> to reduction products.<sup>16,146,149</sup> Thus, even though many potential species have been proposed theoretically<sup>5,143,144,329</sup> we avoid here any detailed discussion on them and refer the reader to other works.<sup>5,332</sup> Instead, we address here reaction species and pathways which were verified experimentally through spectroscopic or specific reduction experiments. \*OCHCH<sub>2</sub>, formed from \*OCCO<sub>H</sub> *via* 5 proton-coupled electron transfers (Figure 15b), is generally assumed as the selectivity-determining intermediate between ethanol and ethylene.<sup>16</sup> Schouten, Koper *et al.* reported fully selective reduction of glyoxal (OCHOCH) and glycolaldehyde (HOCH<sub>2</sub>CHO) to acetaldehyde and ethanol on polycrystalline copper on phosphate buffer by Online Electrochemical Mass Spectrometry (OLEMS).<sup>140</sup> Since both glyoxal and glycolaldehyde can undergo 3 and 1 PCETs to \*OCHCH<sub>2</sub> (Figure 15b),<sup>144</sup> this intermediate is a precursor of ethanol. To further assess the relevance of this precursor, Calle-Vallejo and Koper calculated the thermodynamic cost for further protonation of \*OCHCH<sub>2</sub> either to (1) \*OCHCH<sub>3</sub> or (2) \*O + C<sub>2</sub>H<sub>4</sub>.<sup>300</sup> Both steps are exergonic, with the ethylene selective route, (2), more favorable than the ethanol/acetaldehyde one, (1), by 0.2 eV.<sup>300</sup> Since ethylene is the major C<sub>2</sub> product and it is typically formed at higher rates than ethanol,<sup>35,36,106</sup> the authors concluded that \*OCHCH<sub>2</sub> should be the key reaction intermediate to rule selectivity toward ethylene and ethanol, so-called selectivity-determining step (SDS).

Alternatively, other theoretical studies employed thermodynamic considerations to suggest earlier bifurcation in the ethylene/ethanol route, with among others \*CHCO<sub>H</sub>,<sup>329</sup> \*CO-

CHOH,<sup>144,329</sup> and other ketenes as potential ethylene precursors (Figure 15b). These precursors may then lead to formation of ethylene oxide adsorbed  $*(O)CH_2CH_2$ ,<sup>329</sup> which proved to be fully selective to ethylene on copper in a phosphate buffer.<sup>140</sup> While several sound theoretical evidences confirm a  $*OCHCH_2 \rightarrow *OCHCH_3 \rightarrow$  ethanol route on copper and OD-Cu,<sup>140,142,333</sup> a  $*OCHCH_2$  mediated ethylene pathway seems at odds with the full selectivity of glyoxal and glycolaldehyde to ethanol.<sup>140</sup> Besides, an early bifurcation between the ethylene and ethanol pathways would reconcile experimental and theoretical observations on the role of atomic oxygen to trigger  $C_2H_5OH$  formation.<sup>14,109,125,127,210</sup> Recently, we have proposed a subsurface oxygen-mediated deprotonated glyoxylate (Figure 15b) as the precursor leading to ethanol formation on oxide-derived copper,<sup>210,314</sup> where this alcohol is reported as main reaction product (F.E. > 40% at  $-0.3$  V vs RHE) without any trace of ethylene.<sup>109</sup> Otherwise, this intermediate could undergo one 1 PT and 5 successive PCETs to acetate (Figure 15b), observed concurrently with ethanol at low overpotential on OD-Cu (20% F.E. at  $-0.25$  V vs RHE)<sup>109</sup> Besides, competition between ethanol and ethylene during pulsed  $CO_2$  electroreduction was reported to depend on the cell temperature, with lower temperature ( $5$  °C) favoring the alcohol while higher temperature ( $25$  °C) promoting hydrocarbon formation.<sup>127</sup> This interesting result was attributed to Cu(I) (or equivalently residual oxygen) reducibility.

As a further proof of the role of atomic oxygens from OD-Cu or solvation, more than 60% of oxygen contained in ethanol formed on crystalline copper facets arises from water molecules, as demonstrated for  $C^{16}O$  reduction in  $H_2^{18}O$  solvent.<sup>125</sup> This evidence was confirmed also for acetate and 1-propanol, thus to rationalize this process the authors suggested a  $OH^-$  insertion mechanism on a  $*CCH$  backbone, which then reduce to the three products. Instead, successive theoretical and experimental studies attributed the observation to oxygen exchange between acetaldehyde reduced from CO and water, thus disregarding any role of solvation molecules in earlier reaction stages.<sup>123,124,335</sup> In addition, analogous insertion mechanisms of solvent or subsurface atomic oxygen were predicted. Heenen *et al.*



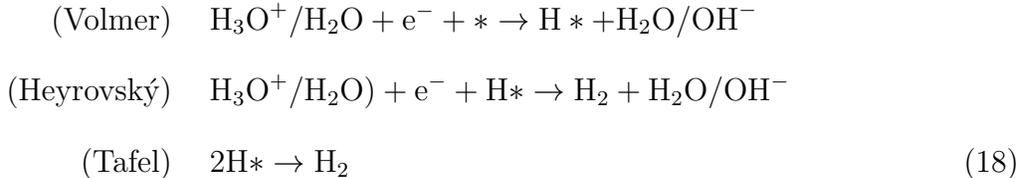
**Figure 15:** eCO<sub>2</sub>R pathway toward major **a**, C<sub>1</sub> and **b**, C<sub>2+</sub> products on metals. Experimental observations are taken from Refs. 140 and 334. The CO-CO dimer bound to a near-surface oxygen is the deprotonated glyoxylate geometry predicted on OD-Cu.<sup>210,314</sup>  $n$  is equal to 3 and 1 for reduction of glyoxal (OCHOCH) and glycolaldehyde (HOCH<sub>2</sub>CHO) to \*OCHCH<sub>2</sub>. **c**, Hydrogen evolution reaction mechanism on metals, as defined in Ref. 41. The key role of cations in enabling CO<sub>2</sub> adsorption, C-C coupling, and Volmer step for water dissociation has been proposed in Refs. 44,45, 328, 42,160,161. Full arrows represent confirmed routes.

employed an *ab initio* kinetic-transport model to describe the mechanism for acetate formation on copper, proposing  $\text{H}_2\text{CCO}$  coupling to  $\text{OH}^-$  in solution as the main precursor toward  $\text{CH}_3\text{CO}_2^-$  with ketene as common  $\text{C}_{2+}$  intermediate (Figure 15b, further details on the mechanism in Section 6).<sup>336</sup> Such analogy between acetate and ethanol reaction route is further supported by  $^{13}\text{CO}_2$ - $^{12}\text{CO}$  electroreduction experiments on OD-Cu copper. Different  $^{13}\text{C}/^{12}\text{C}$  ratios were detected in the reaction products, namely ethylene, ethanol, acetate, and *n*-propanol.<sup>122</sup> Ethylene and propanol showed a high  $^{13}\text{C}$  fraction (0.3-0.5 isotopic composition), thus suggesting a more favorable reduction from  $^{13}\text{CO}_2$  precursors, while acetate and ethanol present an equivalent low  $^{13}\text{C}$  fraction (0.25-0.4), thus preferential formation from paired  $^{13}\text{CO}_2$ - $^{12}\text{CO}$  or  $^{12}\text{CO}$ . This led to the conclusion that the active sites on OD-Cu involved in acetate/ethanol formation might be different from the ones enabling ethylene and propanol formations

Experimental and theoretical insights into  $\text{C}_{3+}$  products are limited by the complexity of the reaction mechanism and the low selectivity of these chemicals, which prevents spectroscopic observation of the potential intermediates and comprehensive screening of electrochemical reduction experiments for key reagents. Even though sequential CO trimerization has been proposed in the past,<sup>337</sup> currently the expected coupling mechanism toward  $\text{C}_3$  products involves a  $\text{OC}(\text{H})\text{CH}_2\text{-C}(\text{H})\text{O}$  precursor,<sup>156,334</sup> potentially involving allyl alcohol ( $\text{CH}_2\text{CHCH}_2\text{OH}$ ) and propionaldehyde ( $\text{CH}_3\text{CH}_2\text{CHO}$ ) at later reduction stage (Figure 15b).<sup>334</sup> In fact, F.E. toward propanol, allyl alcohol, and propionaldehyde for  $\text{eCO}_2\text{R}$  exhibits similar dependencies on applied potential for a polycrystalline copper catalysts<sup>106</sup> and were equally promoted on Ag-decorated  $\text{Cu}_2\text{O}$  nanocubes<sup>154</sup> As for  $\text{C}_4$  products, to date two main reaction pathways have been proposed for the formation of 1-butanol, the only species observed at significant rate.<sup>94,338</sup> Both reaction schemes highlight the key role of an acetaldehyde precursor. Ting *et al.* have indicated that  $\text{CO}_2$  first reduces to acetaldehyde on OD-Cu under alkaline bulk pH.<sup>94</sup> Acetaldehyde then undergoes a base-catalyzed aldol condensation to crotonaldehyde, which is then reduced to butanal, 1-hydroxybutyl,

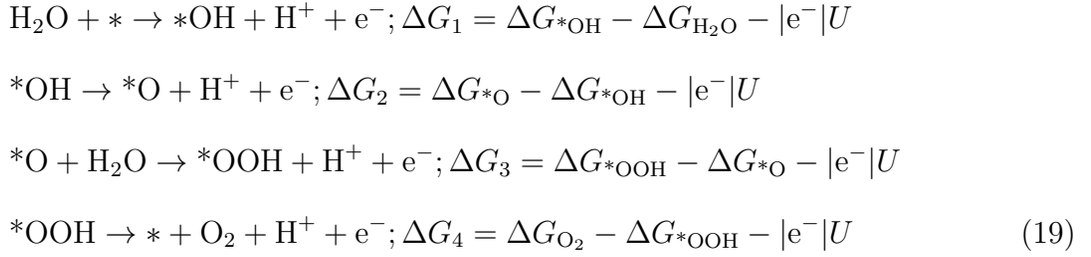
and finally 1-butanol. On the other hand, Choi *et al.* have highlighted a different reduction mechanism on phosphorus-rich copper, involving formation of acetaldehyde from aldehyde self-condensation of two formate precursors. Finally, acetaldehyde undergoes aldol condensation and sequential hydrogenation to 1-butanol.<sup>338</sup>

Moving to the reaction competing with eCO<sub>2</sub>R, hydrogen evolution (Figure 15c), we refer the reader to the nice perspective by Dubouis and Grimaud for a detailed discussion.<sup>41</sup> In short, HER first involves reduction of a proton *via* a proton-coupled electron transfer, the so called Volmer step, Scheme 18. Then, this \*H species can evolve molecular H<sub>2</sub> either through a second proton-coupled electron transfer from solution, Heyrovský step, or the coupling of two adsorbed protons, Tafel step, Scheme 18. H<sub>3</sub>O<sup>+</sup> and H<sub>2</sub>O in solution are typically assumed as proton sources at acidic and alkaline surface pH (Figure 15c)<sup>41</sup> Nevertheless, under electrochemical CO<sub>2</sub> reduction, HER is generally assumed to occur through water reduction (H<sub>2</sub>OR), thus releasing OH<sup>-</sup> upon reaction,<sup>16</sup> since proton reduction is diffusion limited even at acidic bulk pH.<sup>339,340</sup> Proton reduction does not typically exhibit cation effect, while water reduction strongly depends on cation concentration and species.<sup>42,160,161</sup> Such dependence is discussed in details in Section 7.1.



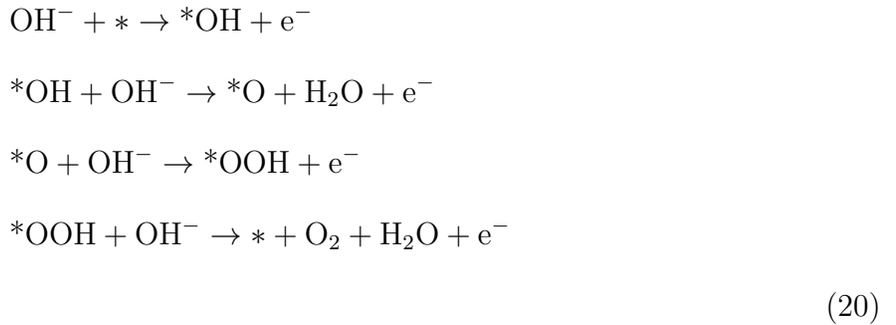
### 5.3 Mechanism for oxygen evolution

In acidic environments, the OER mechanism can be described as below, Scheme 19.<sup>77</sup> In the scheme, \* is adsorption site, \*OH, \*O, and \*OOH are adsorbed OH group, O atom, and OOH group respectively.  $\Delta G_i$  is variation of Gibbs free energy related for each step *i*.



The total energy increase in these four steps is 4.92 eV, which is the energy cost for water splitting. The overpotential is defined by:  $\eta^{\text{OER}} = \max[\Delta G_i]/|\text{e}^-| - 1.23 \text{ V}$ . The ideal catalyst should evenly divide the total energy cost, thus resulting in zero overpotential. Based on this mechanism, the adsorption energies ( $E_{\text{ads}}(X) = E(X*) - E(*) - E(X)$ ,  $X = \text{OH}$ ,  $\text{O}$ , and  $\text{OOH}$ ) of  $\text{OH}$ ,  $\text{O}$ , and  $\text{OOH}$  determines the OER energy profile. To some extent,  $E_{\text{ads}}(\text{OOH})$  correlate with  $E_{\text{ads}}(\text{OH})$  through tLSR, as discussed in detail in Section (2.2.4). On one hand, a strong  $\text{O}$  binding, *i.e.* large  $\Delta G(\text{O}_{\text{ads}})$ , leads to large  $\Delta G_3$  (Scheme 19), hence a large overpotential is expected. On the other hand, a weak  $\text{O}$  adsorption results in large  $\Delta G_2$  and  $\Delta G_4$  (Scheme 19), leading equally to large overpotential. An optimal catalyst is predicted to have  $\Delta G(\text{O}_{\text{ads}}) = 2.46 \text{ eV}$ .

In alkali environments,<sup>77</sup> OER process follows the Scheme 20. for which the considerations reported for alkaline pH applies as well.



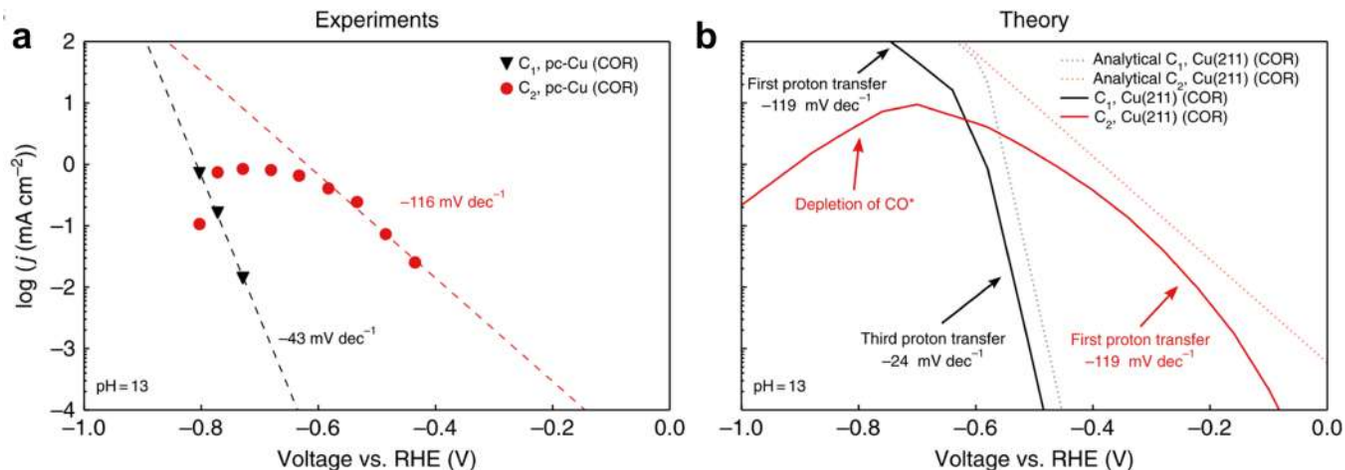
## 6 Microkinetic modeling

Mechanistic studies for the large reaction network involved in eCO<sub>2</sub>R couple well with multi-scale methods like microkinetic modeling (MKM), as this technique can help delineate which reaction pathways have the dominant rates for selective production of the target hydrocarbon or alcohol products. Even though microkinetic modeling is vastly used in heterogeneous catalysis, its application to electrocatalysis is limited by the complexity embedded in estimation of electrochemical activation barriers and potential contributions from electric field and solvent.<sup>341–343</sup> Nevertheless, in the past few years the use of microkinetic modeling has picked up pace and several studies have used MKM to simulate polarization curves for selective CO conversion from CO<sub>2</sub>.<sup>44,320,344–346</sup> A few recent studies have also used MKM to understand the complex reaction network for the formation of C<sub>1</sub> and C<sub>2</sub> hydrocarbon and alcohol products (see for instance Ref. 347), thus the review mainly focuses on these works which go beyond assessment of CO formation.

As discussed in Section 2.1, Hori’s work on Cu surfaces showed that the pH dependence is different for C<sub>1</sub> and C<sub>2</sub> product formations,<sup>37</sup> indicating that the RDS for methane and methanol involves a proton-coupled electron transfer while the RDS for C<sub>2</sub> products involves an electron transfer.<sup>16,141,143</sup> Further, the obtained Tafel slopes are usually smaller for C<sub>1</sub> products as compared to the C<sub>2</sub> products (see Figure 16a), even though Tafel slope analysis may be not always statistically relevant due to physical non idealities, *e.g.* charge transfer, electrostatic screening of adsorbates, etc.<sup>348</sup> To explain the differences in Tafel slope and pH dependence for the formation of C<sub>1</sub> vs C<sub>2</sub> products from CO, Chen and co-workers performed microkinetic modeling for CORR on Cu(211).<sup>349</sup> The insights on the rate determining steps were used to obtain analytical expressions for polarization curves, which aided the interpretation of the MKM results. Using free energy diagrams, the authors predicted that the first proton-electron transfer of OCCO\* intermediate is rate-determining for the formation of C<sub>2</sub> products at low potentials (−0.5 V vs RHE), while the third PCET along CO reduction, \*CHOH formation, is rate determining for C<sub>1</sub> products (Figure 16b). This led to a difference

in Tafel slopes for the two classes of products ( $C_1$  vs  $C_2$ ) based on Equation 21, where  $n$  here corresponds to the number of PCETs which occur before the RDS,  $U$  is the potential vs SHE, and  $k$  is the reaction rate based on Butler-Volmer kinetics. At high negative potentials ( $-1$  V vs RHE), the RDS step changes from PCET to  $^*OCCOH$  to CO dimerization toward  $C_2$  products, while the first proton-electron step is defined as the RDS for  $C_1$  product formation. This led to an increase in Tafel slope for  $C_1$  products, while for  $C_2$  products the current density decreases, as the rate has second order dependence on CO coverage, which depletes as the products are formed.

$$\frac{\partial U}{\partial \log(k)} = -\frac{2.3k_B T}{(n + \beta)e} \quad (21)$$



**Figure 16:** **a**, Experimental polarization curves on polycrystalline Cu for CO reduction toward  $C_1$  and  $C_2$  products at pH = 13. Data are taken from Ref. 350. **b**, Polarization curves predicted by the full microkinetic model and the associated analytical approximation on Cu(211) for CO reduction toward  $C_1$  and  $C_2$  products at pH = 13. Adapted with permission from Ref. 349 (Copyright 2020 Springer Nature under Creative Commons Attribution 4.0 International License<sup>157</sup>).

In terms of pH dependence, experiments showed that there is a drastic decrease in onset potential for formation of  $C_2$  products as we move towards alkaline conditions, while no relevant impact is observed for  $C_1$  products.<sup>350</sup> This was again rationalized by MKM-based analytical expressions derived for current densities, which showed that the earlier the RDS

along the pathway, the larger the decrease in onset potential with pH. Since the RDS for  $C_2$  products involves a PCET which occurs at an earlier stage than the  $C_1$  products, this motivates the stronger impact of pH on overpotentials for  $C_2$  products. Finally, the authors suggested as well depletion of  $C_2$  products at high potentials for alkaline conditions using the microkinetic model.

Peng, Tang, and coworkers also performed microkinetic modeling on Cu(100) for CORR to understand the dominant pathways that lead to the formation of methane and ethylene, used in the study as proxy for  $C_1$  and  $C_2$  product formations.<sup>326</sup> The major focus of the paper was to build mechanisms for  $C_1$  and  $C_2$  formations using surface carbon ( $C^*$ ) as a plausible intermediate. Through their MKM results, they found that for wide potential and pH windows methane is formed through the  $COH^*$  pathway, which followed through a  $*C$  intermediate. As for the ethylene pathway, at neutral pHs and potential windows of interest for  $C_{2+}$  products (from  $-0.4$  to  $-1$  V vs RHE), the  $OC-C^*$  route is the dominant pathway, which shifts to the traditional  $OCCOH$  pathway at alkaline pHs.

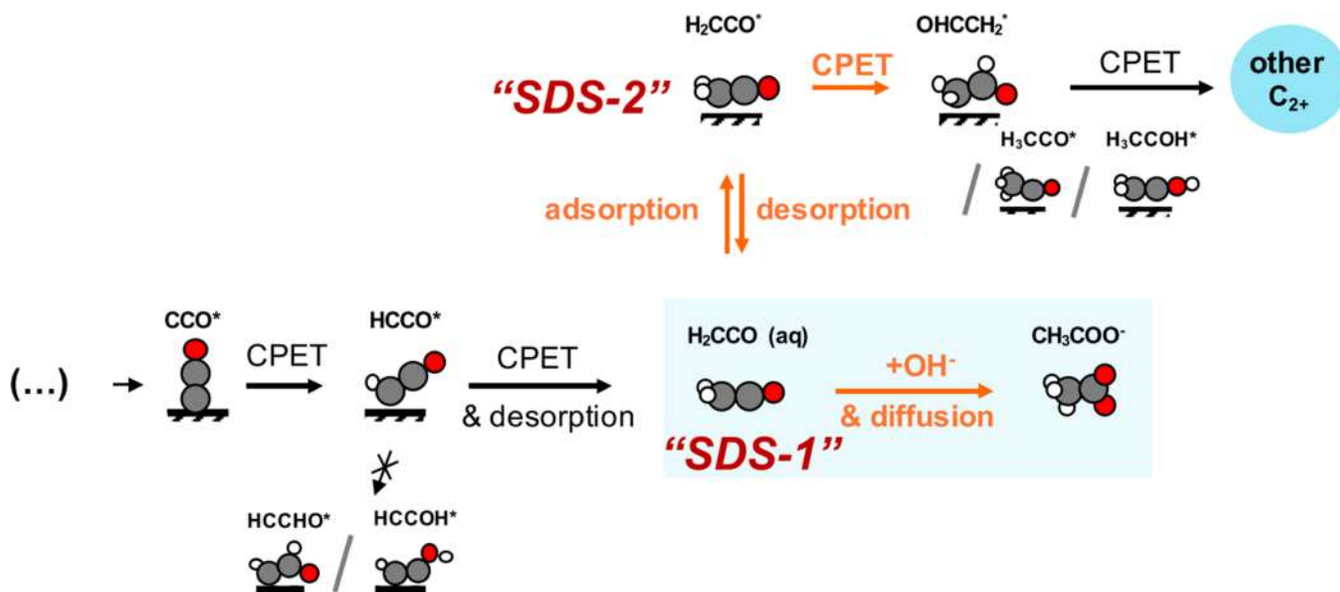
Furthermore, the MKM analysis at neutral pHs illustrated that the protonation of  $C^*$  and the  $C^*-CO$  coupling are the bifurcation points which ultimately determine the competition between  $C_1$  and  $C_2$  products. Using these insights, a simplified rate expression was derived for ratio of rates for  $C_1$  vs  $C_2$  formations in terms of the barriers for  $C^*$  protonation and  $C^*-CO$  coupling. This analytical expression was then used to delineate the trends across different Cu facets. The trend in predicted  $C_2/C_1$  products selectivities was calculated as follows:  $Cu(511) > Cu(100) > Cu(211) > Cu(111)$ , observation which is in good agreement with experiments by Hori *et al.* on single crystal surfaces (Section 2.1).<sup>35,36</sup>

While microkinetic modeling only considers processes which occur at the surface, transport equations need to be integrated to estimate the concentration of species near the electrode. For pathways involving reaction intermediates in solution, these mass transfer effects can ultimately determine which reaction pathways have the highest rates. In the following lines, we focus on two studies where mass transfer has been proved to play a pivotal role

towards defining which reaction mechanism is favored among various plausible alternatives.

Singh *et al.* used multi-scale modeling approaches to look at CO formation from CO<sub>2</sub> on Ag(110) surface by including mass transfer effects.<sup>351</sup> They considered three reaction mechanisms (RM) for the formation of COOH\* and CO\* in which the proton donor is either H\*, H<sub>2</sub>O\*, or H<sub>2</sub>O in solution (denoted as RM-1, RM-2 and RM-3). Remarkably, RM-3 was the only pathway which involved \*CO<sub>2</sub><sup>-</sup> as a plausible surface intermediate in the reaction mechanism. The DFT energetics were inputted into the microkinetic model and the appropriate rate equations for the surface reactions were solved. Next, the current densities from the microkinetic model and the diffusion coefficients of species in the electrolyte were inputted into the continuum transport model and the equations were solved iteratively till the bulk pH and CO<sub>2</sub> concentration converged. Through this systematic approach, the authors found that even though the RM-3 pathway had the highest activation energies for the CO formation, only this reaction scheme could reproduce the experimental correlation observed between current densities and CO<sub>2</sub> partial pressure.<sup>352</sup> Further, the most abundant adsorbed species were found to be \*COOH and \*CO<sub>2</sub><sup>-</sup> which is in agreement with the spectroscopic observations on silver catalysts. Finally, the authors concluded that explicit assessment of mass transfer effects enable better reproduction of the experimental trends of CO and H<sub>2</sub> partial current densities vs applied potential.

More recently, Chan and co-workers used a coupled microkinetic and transport model to study the mechanism for acetate formation on Cu surfaces.<sup>336</sup> The authors proposed a reaction scheme where the surface intermediate ketene (CH<sub>2</sub>CO\*) desorbs and reacts with OH<sup>-</sup> in solution. Overall, two possible selectivity-determining steps (SDS) for acetate formation were envisioned (Figure 17): (1) reaction of H<sub>2</sub>COO (aq) with OH<sup>-</sup> vs readsorption of this species (SDS-1) (2) desorption of H<sub>2</sub>CO\* into solution vs further proton electron steps to form other C<sub>2</sub> products (SDS-2). It is important to note that the SDS-1 only involves chemical steps, while the SDS-2 involves both a chemical and a PCET step. Energetics and initial concentrations of the reactants were then inputted in the integrated multi-scale model,



**Figure 17:** Schematic of the ketene pathway, including the two selectivity-determining steps towards acetate, either *via* a reaction in solution (SDS-1), or further reduction of the adsorbed ketene ( $\text{H}_2\text{CCO}^*$ ) to form other  $\text{C}_2$  products (SDS-2). The excluded alternative reduction steps are reported in the bottom left. Adapted with permission from Ref. 336 granted by the authors under Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License.<sup>353</sup>

which showed that acetate selectivity followed a U-shaped dependence on applied potential vs SHE, in agreement with experimental data. At low overpotentials, the PCET step towards the formation of  $\text{H}_2\text{COO}^*$  from the ketene intermediate is limiting. As the potential shifts to more negative values, this PCET step becomes more favorable, consequently leading to decrease of acetate selectivity (SDS-2). Finally, at very high overpotential,  $\text{H}_2\text{COO}^*$  formation is now facile and acetate selectivity is now ruled by the competition between  $\text{H}_2\text{COO}^*$  reduction or desorption and further reaction with  $\text{OH}^-$  in solution (SDS-2). Thus, the formation of acetate is now dependent on local  $\text{OH}^-$  concentration (SDS-1), which increases at more negative potentials due to water and  $\text{CO}_2$  reduction, leading again to a U-shaped dependence on applied potential. Moreover, the effects of pH and surface roughness are also confirmed by specific experiments. Intriguingly, this solution phase mechanism are at odds with other proposed mechanisms for acetate formation<sup>125</sup> (Section 5.2), consequently the authors proved the power of an integrated transport/kinetic framework for finding the

dominant reaction pathways which drive the formation of a specific product.

## 7 Modeling electrolyte and bulk mass transfer effects

In addition to Faradaic processes at the surface, phenomena occurring within the diffusion layer are crucial in electrocatalysis.<sup>2</sup> As shown in Section 3.3, a proper understanding of local electrolyte effects can be employed to enhance the performance of eCO<sub>2</sub>R and HER.<sup>15,42,161,354</sup> Explicit cation effects are well-known to promote the overall activity and selectivity of these reactions,<sup>32,161</sup> whilst surface poisoning from electrolyte anions have the opposite effect.<sup>174–176</sup> Besides, mass transfer within the diffusion layer is also instrumental for the industrial exploitation of eCO<sub>2</sub>, motivating the need for optimized configurations such as gas diffusion electrodes.<sup>355,356</sup> Lastly, chemical reactions within the diffusion layer must be properly accounted, since they can limit the overall performance of electrocatalytic devices<sup>357</sup> or tune product distributions through chemical instead of electrochemical steps.<sup>94,358</sup> For instance, the conversion of CO<sub>2</sub> and OH<sup>-</sup> to CO<sub>3</sub><sup>2-</sup> is the main source of energy loss in CO<sub>2</sub> reduction under neutral to basic bulk pH.<sup>357</sup> Consequently, industrially relevant single-pass CO<sub>2</sub> utilization and conversion/energy efficiency can be achieved only at acidic pH, as demonstrated independently by Huang *et al.*<sup>179</sup> and Monteiro *et al.*<sup>359</sup> As further examples of chemical steps within the diffusion layer, formation of carboxylic acids and alcohols during CO<sub>2</sub> reduction conditions was demonstrated to occur through disproportionation of aldehydes due to high alkaline surface pH instead of direct eCO<sub>2</sub>R.<sup>358</sup> Besides, eCO<sub>2</sub>R to 1-butanol (F.E. = 0.056%,  $j = -0.080 \text{ mA cm}^{-2}$  at  $-0.48 \text{ V vs RHE}$ ) on an oxide-derived copper catalyst was proposed to happen through a final chemical step, *i.e.* hydrogenation of 1-hydroxybutyl formed from acetaldehyde reduction (for further details on the mechanism, see Section 5.2).

## 7.1 Cation effects

Electrolyte effects are known in eCO<sub>2</sub>R since the initial observations by Hori *et al.* illustrating higher activity for larger ionic radius alkali cations (Section 2.1).<sup>33,170,171,360</sup> Following Frumkin’s theory,<sup>361</sup> the authors attributed this enhancement to modification of the local electrostatic field within the electrical double layer through specific cation adsorption. In particular, weakly hydrated cations can better accumulate at the Outer Helmholtz plane (OHP), thus leading to a more intense electrostatic field.<sup>33</sup> However, the occurrence of specifically adsorbed cations under CO<sub>2</sub> reduction conditions is a topic of vivid debate in the field.<sup>362</sup>

Mills *et al.* employed density functional theory to estimate the equilibrium potential  $U_0$  for cation adsorption on fcc(111) facets of Ag, Au, Ni, Pd, Pt, Equations 22 and 23.<sup>363</sup>  $G^*_{\text{cat}}$ ,  $G^*$ , and  $G_{\text{cat}^+}$  are the Gibbs free energies for the metal/water/cation system, metal/water, and solvated cation alone, respectively.  $G_{\text{cat}^+}$  was estimated from the free energy of cation in gas phase, corrected by experimental solvation energy at 1 M solution. The term which depends on potential at PZC ( $U_{\text{pzc}}$ ) and cation (system) electric dipole moment,  $|\vec{p}^*_{\text{cat}}|$  ( $|\vec{p}^*|$ ), is included to account for the partial positive charge retained by the cation upon solvation, which generates a dipole moment along the surface normal direction. Finally,  $d$  represents the thickness of the electrical double layer.



$$U_0 = \frac{-G^*_{\text{cat}} + G^* + G_{\text{cat}^+} + U_{\text{pzc}} \frac{|\vec{p}^*_{\text{cat}}| - |\vec{p}^*|}{d}}{|e^-| + \frac{|\vec{p}^*_{\text{cat}}| - |\vec{p}^*|}{d}} \quad (23)$$

By referring Equation 23 to the Normal Hydrogen Electrode reference and carrying out benchmarks with different numbers of water molecules within the solvation layer, the authors estimated equilibrium adsorption potentials for cations more negative than  $-0.80$  V vs NHE (Cs<sup>+</sup> adsorption on Pt) and far from CO<sub>2</sub> reduction conditions for Ag and Au ( $U_0 < -2.0$  V vs NHE) (Table 1).<sup>363</sup> Although these equilibrium adsorption potentials are significantly more negative than cathodic bias for eCO<sub>2</sub>R, we note that the role of near-surface solvent or

other electrolyte effects may further stabilize cation specific adsorption on metals. Thus, this process should not be disregarded. In a successive study from the same authors, iodine and potassium co-adsorption was suggested to occur on copper under CO<sub>2</sub> reduction conditions at low cation coverage ( $\theta = 0.11$  ML) and alkaline pH, leading to a preferential increase of binding energy of O-terminated species.<sup>225</sup> In line with such prediction, recently a Total Reflection Surface Enhanced Infrared Absorption Spectroscopy (ATR-SEIRAS) study of CO adsorption configurations on Pt and Au revealed that at low cathodic bias the band for linearly-bond CO disappears, likely due to displacement by K<sup>+</sup> at the OHP.<sup>364</sup>

**Table 1:** Estimated equilibrium potential  $U_0$  (V vs NHE) for cation adsorption on fcc(111) electrode surfaces. Data taken from Ref. 363

system	H <sup>+</sup>	Li <sup>+</sup>	Na <sup>+</sup>	K <sup>+</sup>	Cs <sup>+</sup>	Ba <sup>2+</sup>
Pt(111)	-0.07	-1.30	-0.90	-0.98	-0.80	-1.31
Pd(111)	+0.09	-1.63	-1.19	-1.35	-1.09	-1.43
Ni(111)	+0.05	-2.06	-1.61	-1.85	-1.68	-1.86
Au(111)	-1.29	-2.76	-2.40	-2.74	-2.64	-2.46
Ag(111)	-0.65	-2.63	-2.11	-2.52	-2.44	-2.62

Following our overview on cation effects, an increasing number of experimental reports attributed the trends on eCO<sub>2</sub>R activity and selectivity to non electric fields effects, such as cation coordination with intermediates at the surface,<sup>365</sup> or increased polarizability of adsorbates.<sup>366</sup> To unveil this phenomenon, Singh *et al.* employed a 1D mathematical model for an electrochemical cell for eCO<sub>2</sub>R with anolyte, catholyte, Pt anode, Cu (and Ag) cathode, a anion-exchange membrane to separate anolyte and catholyte, and different species within the electrolyte: CO<sub>2</sub>, HCO<sub>3</sub><sup>-</sup>, CO<sub>3</sub><sup>2-</sup>, H<sup>+</sup>, OH<sup>-</sup> and alkali metal cations (Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, Rb<sup>+</sup>, Cs<sup>+</sup>).<sup>172,367</sup> This model was solved using the COMSOL multiphysics software and including ion migration, diffusion, acid-base equilibrium, gas-liquid transport of CO<sub>2</sub>, hydrolysis of cations, and kinetics for both OER and CO<sub>2</sub>.<sup>172</sup> Since the  $pK_a$  for an hydrated cation represents the ease to undergo hydrolysis, the authors estimates this parameter in the bulk electrolyte and in the proximity of the Cu and Ag cathodes at -1.0 V vs RHE, respectively. Whilst cation  $pK_a$  at the bulk increases following the ionic radius, since larger

ionic radius cations exhibit lower electrostatic interaction with neighboring water molecules, this trend overturns near the cathode surface due to the higher electrostatic field generated by weakly hydrated cations (larger ionic radius), Table 2. The authors derived an analytical equation to estimate the  $pK_a$  for cation hydrolysis at the cathode, which is proportional to both cation/water and cation/cathode electrostatic interactions. These terms are shown in Equation 24, where  $z$  is the effective charge of the hydrated cation,  $\sigma$  the surface charge density,  $r_{M-O}$  the sum of cation (M) and oxygen (O, from water) radii, and  $r_{H-EI}$  the distance between hydration shell and cathode surface. Equation 24 was verified experimentally by measuring the  $pK_a$  of monovalent alkali metal cations through potentiometric titration, thus  $A$  and  $B$  were calculated by fitting the analytical equation to the experimental data.

$$pK_a = -A \left[ \frac{z^2}{r_{M-O}} + 2\pi\sigma z r_{H-EI} \left( \sqrt{1 + \frac{r_{M-O}^2}{r_{H-EI}^2}} - 1 \right) \right] + B \quad (24)$$

Since pH at the surface is rather alkaline under CO<sub>2</sub> reduction conditions in a neutral bulk electrolyte,<sup>16</sup> then K<sup>+</sup>, Rb<sup>+</sup>, Cs<sup>+</sup>, having low  $pK_a$ ,<sup>367</sup> can undergo hydrolysis, thus leading to pH buffering and consequent higher concentrations of CO<sub>2</sub> at the surface. In absence of alkali cations, an alkaline pH at the surface involves rapid neutralization of CO<sub>2</sub> by a neighboring OH<sup>-</sup> to form carbonate, so that CO<sub>2</sub> surface concentration decreases significantly with consequent mass transfer limitations on eCO<sub>2</sub>R activity. Hence, the well-known cation effect on eCO<sub>2</sub>R was solely rationalized through hydrolysis of electrolyte cations, suggesting that weakly hydrated cations have a lower  $pK_a$  at the surface, thus they allow for an increased pH buffering and higher CO<sub>2</sub> availability. The continuum model employed resulted in remarkable agreement with experimentally measured current densities on Cu and Ag at -1.0 V vs RHE for the different cations assessed. However, a recent study estimated CO<sub>2</sub> concentration at a Au electrode from ATR-SEIRAS measurements and demonstrated a negative correlation of this parameter with cation ionic radius,<sup>368</sup> since weakly hydrated cations promote CO<sub>2</sub> reduction to a larger extent, thus leading to higher CO<sub>2</sub> consumption

and lower local concentration. Instead, the prediction of higher pH buffering for cations with larger ionic radius was confirmed by ATR-SEIRAS characterization, where the authors probed pH at a gold/electrolyte interface from the  $\text{CO}_2/\text{HCO}_3^-$  bands ratio.<sup>369</sup>

**Table 2:**  $\text{p}K_a$  for cation hydrolysis in the bulk electrolyte (bulk) and at the surface layer of copper (Cu) and silver (Ag) cathodes at  $-1.0$  V vs RHE,<sup>172</sup> obtained by solving Equation 24. Data taken from Ref. 172

cation	bulk	Cu (Surf)	Ag (surf)
$\text{Li}^+$	13.6	11.64	13.16
$\text{Na}^+$	14.2	10.26	11.44
$\text{K}^+$	14.5	7.95	8.49
$\text{Rb}^+$	14.6	6.97	7.23
$\text{Cs}^+$	14.7	4.31	4.32

Some of the proponents of the hydrolysis hypothesis carried out a follow up study with lower applied potentials, so that mass transfer contributions to  $\text{eCO}_2\text{R}$  activity and selectivity could be neglected. Specifically, Resasco *et al.* demonstrated that Faradaic efficiencies for  $\text{eCO}_2\text{R}$  on Cu(100) vary by increasing the  $\text{CO}_2$  flow rate for potentials more negative than  $-1.1$  V vs RHE,<sup>170</sup> thus suggesting the critical role of  $\text{CO}_2$  diffusion under the regime of their previous study.<sup>172</sup> Overall, production rates of formate, ethylene, and ethanol showed a positive correlation with the ionic radius of alkali cations for Cu(100) and Cu(111), whilst hydrogen, carbon monoxide, and methane formations were not affected. Similarly, both CO and  $\text{HCOO}^-$  partial current densities on Ag and Sn increased following the trend of cation ionic radius. Small additions of larger size cations on a 0.1 M  $\text{LiHCO}_3$  electrolyte with constant ionic strength suddenly changed the partial current density toward ethylene, ethanol, and formate on Cu(100), suggesting that larger cations accumulate more at the surface/electrolyte interface. To rationalize the last evidence, the authors employed a simplified model for a Cu/electrolyte interface: a Cu(111) supercell with two ice-like water bilayers, one solvated cation, 12 Å of vacuum, and explicit electric potential.<sup>44,170</sup> Constrained minima hopping (CMH)<sup>268</sup> was carried out on the system to optimize cation solvation shell. Finally, the thermodynamic driving force for each cation to accumulate at a distance of 6.25

Å from Cu(111) was estimated from the lowest energy configurations of the solvated cations and the cation reduction potentials.<sup>91</sup> In fact, the energy of the cation/electron pair in the bulk electrolyte  $G_{\text{cat}^+ + \text{e}^-, \text{bulk}}$  can be derived by applying the CHE<sup>8</sup> formalism to cation reduction (Equation 25), where cat and solv stand for cation and explicit solvent molecules within its coordination shell. At the equilibrium potential for cation reduction,  $U_0$ , both processes are in equilibrium, thus the energies of products and reactants are equivalent. Thus,  $E_{\text{cat}^+ + \text{e}^-, \text{bulk}}$  at a given potential  $U$  can be calculated from Equation 26. Since the energy of the cation/electron pair at the OHP,  $E_{\text{Cat}^+ + \text{e}^-, \text{OHP}}$ , can be estimated from the CMH simulations, the thermodynamic driving force for cation accumulation is given by Equation 27.



$$E_{\text{cat}^+ + \text{e}^-, \text{bulk}} = E_{\text{cat}} - |\text{e}^-|(U - U_0) \quad (26)$$

$$\Delta E_{\text{cat}, \text{OHP}} = E_{\text{cat}^+ + \text{e}^-, \text{OHP}} - E_{\text{cat}^+ + \text{e}^-, \text{bulk}} \quad (27)$$

The dependence of  $\Delta E_{\text{cat}, \text{OHP}}$  on potential  $U$  (vs SHE) is derived from the work function  $\phi$  of the interface referred to the work function of the SHE electrode and it is scaled through the partial electron transfer to the cation upon migration from bulk electrolyte to the OHP.<sup>170</sup> Taking  $\Delta E_{\text{cat}, \text{OHP}}$  for  $\text{Li}^+$  as a reference, accumulation is more exothermic for larger size cations (Figure 18a), thus suggesting that cation trend on eCO<sub>2</sub>R is mainly due to larger concentrations at the OHP of  $\text{K}^+$  and  $\text{Cs}^+$  versus  $\text{Li}^+$  and  $\text{Na}^+$ , respectively. The hypothesis of the preminent role of accumulation to drive eCO<sub>2</sub>R activity was further confirmed by an almost equivalent cation-induced electric field near a CO molecule adsorbed on Cu(111), independent from the cation species in the vicinity.<sup>170</sup> Higher accumulation at the surface for cations with larger ionic radius was confirmed independently in a recent theoretical study solving generalized modified Poisson-Nernst-Planck equations for a CO<sub>2</sub> reduction system.<sup>164</sup> The mathematical details behind the model will be discussed in Section 7.3. For a cathodic bias of  $-0.32$  V vs PZC,  $\text{Li}^+$ ,  $\text{Na}^+$ ,  $\text{K}^+$ , and  $\text{Cs}^+$  concentration at the OHP accounted for

$\sim 4$  M,  $\sim 4.5$  M,  $\sim 6$  M, and  $\sim 6$  M respectively.

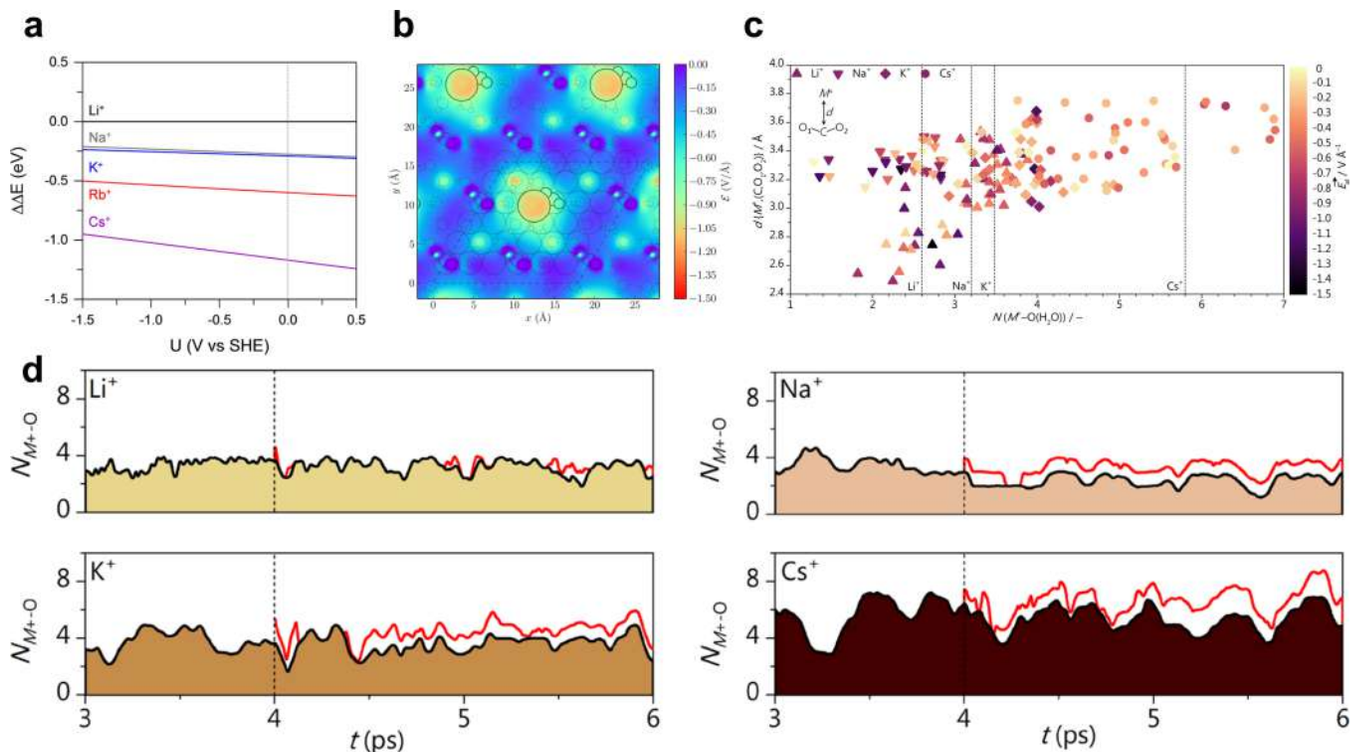
As for specific cation effects, an earlier study by Chen *et al.* rationalized the local stabilization of reaction intermediates through the dependence of adsorbate formation energy on its electric dipole moment  $\vec{p}$  and polarizability  $\alpha$ , as well as on the electric field  $\vec{E}$ , Equation 28.<sup>44</sup> By calculating CO<sub>2</sub> adsorption energy versus the applied electric field  $\vec{E}$ , induced either by an external dipole potential under vacuum,<sup>231</sup> or insertion of solvated cations close to the surface (K<sup>+</sup> and 1-ethyl-3-methylimidazolium, EMIM<sup>+</sup>), the authors proved the correctness of the former hypothesis.<sup>44</sup> The presence of a solvated cation close to the surface leads to very localized electric field in the order of  $-1.0$  V Å<sup>-1</sup>, which quickly decays beyond a radial distance of 5 Å (Figure 18b). The electric field at a given radial distance  $\vec{r}$  from the cation was derived from the variation of electrostatic potential due to charged species, Equation 29, where surf, solv, and cat stand for surface, solvent molecules, and cation respectively. Hence, the cation-induced electric field,  $\vec{E}_{\text{cat}}$ , is given by Equation 30, where  $\vec{r}_{\text{ads}}$  and  $\vec{r}_{\text{cat}}$  are the coordinates of adsorbate and alkali cation. The additional negative sign highlights that the electric field is estimated for a negative unit charge.

$$E(\vec{E}) = E(\vec{E} = 0 \text{ VÅ}^{-1}) + \vec{p} \cdot \vec{E} - \frac{\alpha |\vec{E}|^2}{2} \quad (28)$$

$$U(\vec{r}) = U_{\text{surf/solv/cat}} - U_{\text{surf}} - U_{\text{solv}} - U_{\text{cat}} \quad (29)$$

$$-\vec{E}_{\text{cat}} = - \left( - \frac{U(\vec{r}_{\text{ads}}) - U(\vec{r}_{\text{cat}})}{\|\vec{r}_{\text{ads}} - \vec{r}_{\text{cat}}\|} \right) \hat{r} \quad (30)$$

Recently, we applied the formalism mentioned above (Equations 29, 30) to demonstrate that cation-induced electric field strongly depends on cation coordination shell and cation-adsorbate distance (Figure 18c). In general, electric field is more intense for low cation-water coordination numbers and short distances, in excellent agreement with a recent experimental study of cation effect on eCO<sub>2</sub>R on Au.<sup>370</sup> The employment of a more accurate model of the catalyst/electrolyte interface revealed an additional chemical effect of the cation in addition



**Figure 18:** Role of metal cations in eCO<sub>2</sub>R. **a**, The thermodynamic driving force for cation accumulation at the OHP of a Cu(111) electrode is stronger for alkali cations with larger ionic radius.<sup>170</sup> **b**, Cation-induced electric field on the  $z$ -plane corresponding to a CO<sub>2</sub> adsorbed on a (6 × 6) Ag supercell. K<sup>+</sup> coverage account for  $\frac{1}{36}$  ML. Solvent atoms are indicated as dashed circles, whilst cation, CO<sub>2</sub>, and Ag atoms are highlighted as black circles.<sup>44</sup> **c**, Dependence of alkali cation-induced electric field on solvation shell ( $x$ -axis) and cation-adsorbate average distance ( $y$ -axis).<sup>45</sup> **d**, Cation-water (filled area) and cation-CO<sub>2</sub> (red line) coordination number during 2 ps of AIMD simulations on a 4-layer ( $3\sqrt{3} \times 3\sqrt{3}$ ) Au(111) model with 72 water molecules and 1 alkali cation.<sup>45</sup> Adapted with permission from Ref. 170 (Copyright 2017 American Chemical Society), Ref. 44 (Copyright 2016 American Chemical Society), and Ref. 45 (Copyright 2021 Springer Nature) respectively.

to the mean-field electrostatic stabilization previously discussed (Equation 28).<sup>44,171</sup> In fact, we applied AIMD simulations on a ( $3\sqrt{3} \times 3\sqrt{3}$ )-R30° Au(111) supercell with 72 explicit water molecules pre-optimized for a Pt(111)/water system (20 ps).<sup>22,45</sup> The Au/water system underwent an initialization of 2 ps AIMD simulation (1 fs time step) to optimize the solvation layer, then an additional 2 ps upon introduction of an alkali cation (Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, and Cs<sup>+</sup>) close to the surface, and a final optimization for 2 ps upon adsorption of a CO<sub>2</sub> molecule in the vicinity of the cation. The datasets related to this work are available on the ioChem-BD

database.<sup>371</sup>

By assessing the cation-oxygen radial distribution function and defining cation local coordination shell during the AIMD simulations on the Au/water/cation/CO<sub>2</sub> system, we observed a local coordination between cation and adsorbate (Figure 18d), which was more frequent for weakly hydrated cations, such as K<sup>+</sup> and Cs<sup>+</sup>.<sup>45</sup> Upon coordination, a short-range cat<sup>+</sup> ··· O(CO<sub>2</sub>) interaction occurs, which accounts for a stabilization of CO<sub>2</sub> adsorption by 0.5 eV. This promotional effect is rather independent of the cation species, thus this evidence confirms that CO<sub>2</sub> reduction activity trends must be attributed to accumulation processes rather than distinct cation stabilization.<sup>170</sup> While CO<sub>2</sub> adsorption is endergonic by  $\sim +0.8$  eV for the cation-free case, this process becomes almost exergonic if facilitated by a neighboring cation, thus supporting the experimental results from Prof. Koper’s group that metal cations are key to enable eCO<sub>2</sub>R on gold, silver, and copper. The promoting effect of cation is highlighted as well by (1) the decrease of the O-C-O angle,  $\alpha \sim 130^\circ$ , which indicates activation of the CO<sub>2</sub> molecules, (2) and the enhancement of the first electron transfer from the surface. As a consequence of the theoretical and experimental evidences reported in the study,<sup>45</sup> an updated mechanism for CO<sub>2</sub> reduction to CO was proposed, involving the instrumental role of cation in stabilizing the CO<sub>2</sub><sup>-</sup> intermediate at the surface prior to protonation to COOH. Besides, cation-CO<sub>2</sub> coordination number was proposed as a unique descriptor for eCO<sub>2</sub>R activity and very recently its validity was extended to bi- and trivalent cations as well.<sup>42</sup> In addition to CO<sub>2</sub> adsorption, cation stabilization effect may be crucial for C-C coupling as well.<sup>147</sup>

Overall, cation accumulation at the OHP,<sup>170</sup> mean-field effects,<sup>171</sup> and localized electrostatic stabilization<sup>44,45</sup> are the three main phenomena ruling cation effect on CO<sub>2</sub> reduction, and must be properly accounted in theoretical modeling. While accumulation and mean-field effects rule CO<sub>2</sub> reduction activity and any trends based on cation ionic radius, local electrostatic interactions are key for enabling the overall process by stabilizing crucial reaction intermediates. These considerations support the accuracy of semi-quantitative predictions

on eCO<sub>2</sub>R activity trends from theoretical models based solely on concentration and mean-field phenomena. For instance, Ringe *et al.* applied a 1-dimensional continuum electrolyte model,<sup>171</sup> the size-modified Poisson Boltzmann approach,<sup>372</sup> to describe the induced charge density at the density at a given cation concentration and applied potential, Equations 31, 32.  $\epsilon_{\text{bulk}}$  is the bulk dielectric permittivity,  $z_i$  and  $C_i$  are respectively ionic charges and concentrations (negative sign for anions, positive for cations),  $F$  and  $R$  the Faraday and Ideal Gas constant,  $T$  the temperature, and  $\Phi$  the electrostatic potential within the electrical double layer.  $\chi_0$  represents the ion-occupied volume fraction given by  $2C_{\text{bulk}}a^3N_A$  where  $N_A$  stands for the Avogadro constant and  $a$  is the lattice cell length in the statistical model, *i.e.* the ion diameter in this case. Equation 32 can be solved applying Dirichlet boundary conditions at the end of the diffusion layer ( $\Phi = 0$  for  $x = x^\infty = 80 \mu\text{m}$ ) and Robin boundary condition at the electrode surface, Equation 33, where  $\sigma$  is the cation-induced surface density,  $\Phi^{\text{M}}$  and  $\Phi^{\text{M,PZC}}$  are respectively the electrode potential of the metal relative to the bulk electrolyte and its potential at the PZC.  $C_{\text{gap}}$  stands for the interfacial Helmholtz capacitance. By solving Equations 31, 32, and 33 the authors derived the dependence of surface density  $\sigma$  on potential  $\Phi$  and ion size  $a$ . The variation of Gibbs free energy of CO<sub>2</sub> and OCCO adsorption driven by the increased electronic density at the surface (Equation 28) was obtained through density functional theory, so it was possible to estimate CO and C<sub>2</sub> partial current density as a function of  $\sigma$ . Finally, by coupling continuum model and DFT, the authors mapped the dependence of eCO<sub>2</sub>R activity and C<sub>2</sub> selectivity on the ion size  $a$  for silver and copper, showing excellent agreements with experimental reports.<sup>171</sup>

$$\epsilon_{\text{bulk}} \frac{d^2\Phi}{dx^2} = \sum_i z_i C_i[\Phi] \quad (31)$$

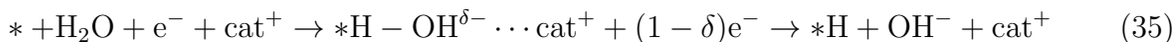
$$C_{\pm}[\Phi] = C_{\text{bulk}} \frac{e^{\pm \frac{z_i F \Phi}{RT}}}{1 - \chi_0 + \chi_0 \cosh\left(\frac{z_i F \Phi}{RT}\right)} \quad (32)$$

$$\sigma = \epsilon_{\text{bulk}} \frac{d\Phi}{dx} = C_{\text{gap}} [\Phi^{\text{M}} - \Phi^{\text{M,PZC}} - \Phi(x=0)] \quad (33)$$

In addition to CO<sub>2</sub> reduction, cation effects have been reported as well for hydrogen evolution at alkaline pH, albeit showing different trends depending on the catalysts. HER has been observed to decrease following the alkali group for Pt and Ir, whilst this trend overturns for Au and Ag.<sup>373</sup> Prof. Koper and his research group recently proposed a novel mechanism to rationalize such effect on transition metals, based on theoretical and fundamental insights.<sup>42,160,161,374,375</sup> By increasing cation concentration at a constant bulk pH, a reaction order  $\sim 0.6$  of HER on cation concentration was observed at bulk pH = 11 on a gold electrode. However, such dependence turned negligible (reaction order  $\sim 0$ ) for pH = 12 and lead to a negative reaction order of around  $-1$  at pH = 13. Besides, *in situ* SERS detected a red shift of the vibrational band related to H adsorption on gold when the bulk pH increases, thus suggesting changes in hydrogen binding energies. Thus, both evidences hint at the occurrence of a convoluted effect between electrolyte pH and cation concentration, where both parameters enhance water reduction to a certain extent.<sup>160</sup> Overall, Goyal *et al.* proposed cation to promote the rate-determining Volmer step for water reduction on gold, Equation 34, by stabilizing the transition state associated with the reaction as in Equation 35. Later, McCrum *et al.* extended the validity of this scheme as well to Pt, for which Na<sup>+</sup> enhances water dissociation as well.<sup>374</sup> In the same study, the authors unveiled the role of OH<sup>-</sup> binding energy to tune HER activity, predicting a 3-D volcano plot for HER activity using  $\Delta G_{*OH}$  and  $\Delta G_{*H}$  as suitable descriptors.

To rationalize the convolution between cation and pH effects, Goyal *et al.* assumed enhancement of HER at pH = 11, saturation of such effect at pH = 12 due to achievement of maximum concentration of cations at the surface, and a site blocking effect for pH = 13, motivating the negative reaction order of  $\sim -1$ .<sup>161</sup> Based on these hypothesis, the authors derived a general empirical rate ( $\nu$ ) law, Equation 36, where  $k$  is the rate constant (we employ here the standard notation, even though the rate depends on the temperature),  $\alpha$  the transfer coefficient,  $F$  and  $R$  the Faraday and Ideal gas constants,  $U$  the applied potential vs the standard potential of the reaction,  $T$  the temperature,  $\Gamma_{cat,surf}$  and  $\Gamma_{cat,max}$  respectively

cation surface concentration and its saturation value, and  $\gamma$  an empirical reaction order of HER on  $\Gamma_{\text{cat,surf}}$ . Hence, for low cation concentrations at the surface, *i.e.*  $\Gamma_{\text{cat,surf}} \ll \Gamma_{\text{cat,max}}$ , Equation 36 correctly reproduces an empirical reaction order of  $\gamma$ . Instead, the correlation between pH and cation surface coverage can be described through the dependence of cation coverage on the effective potential  $U_{\text{eff}}$  and the bulk cation concentration  $C_{\text{cat,bulk}}$ , assumed to follow a Frumkin isotherm in Equation 37. In this equation,  $K$  is the standard equilibrium constant for cation adsorption at the surface and  $g$  is the Frumkin interaction parameter, higher than 0 in case of repulsion between adsorbates. Since experimental results were obtained at constant potential  $U$  vs the RHE, the potential at the PZC for the Au electrode,  $U_{\text{pzc}}$ , shifts positively for increasing surface pH ( $\text{pH}_{\text{surf}}$ ), thus  $U_{\text{eff}}$  exhibits a negative pH-dependent offset, Equation 38. Recently, Monteiro *et al.* rationalized experimental evidences of cation effects on Pt and Au by defining a generalized cation promotion/inhibition scheme based on the interaction between specific transition metal and alkali cation.<sup>375</sup>

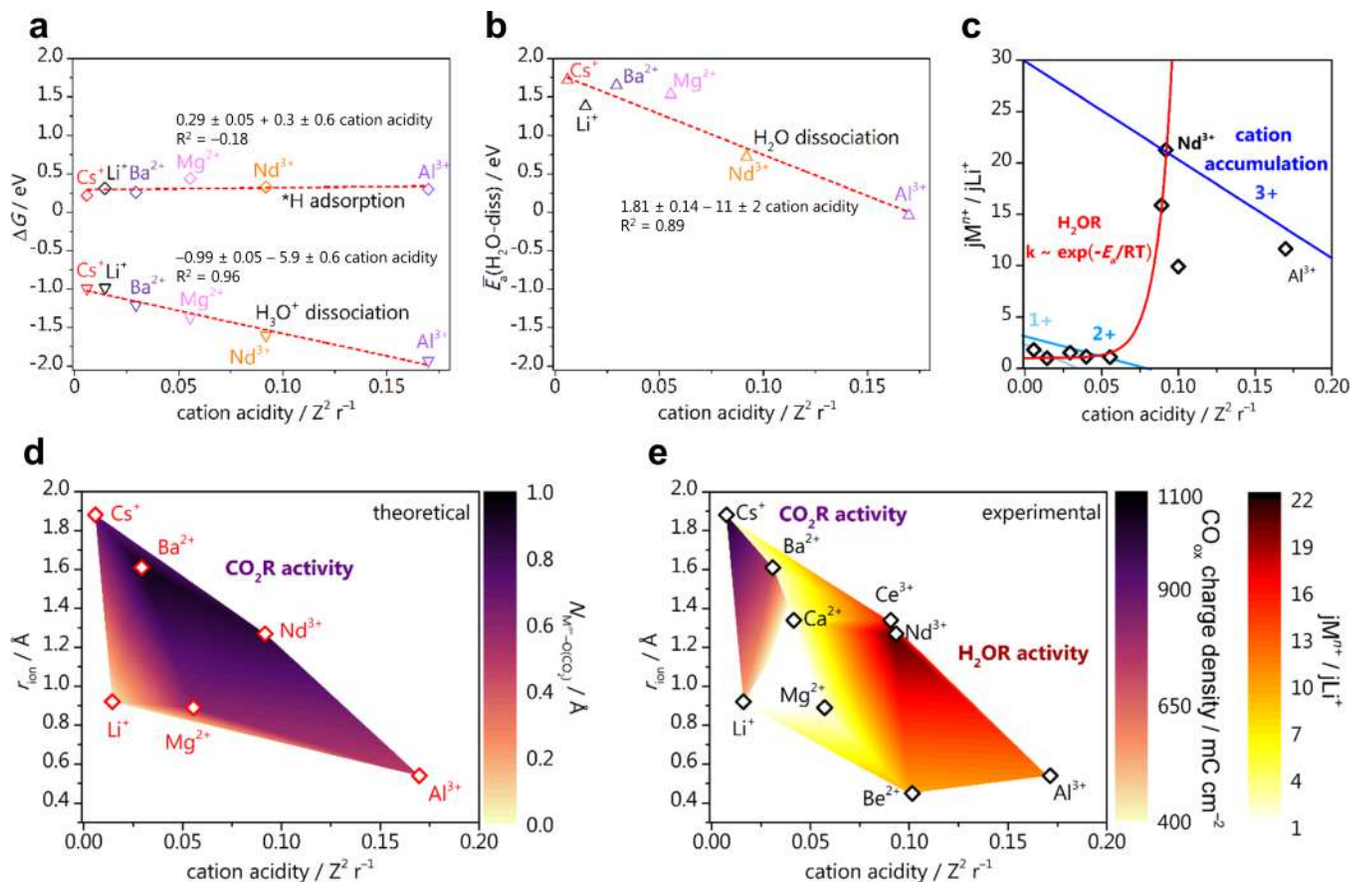


$$\nu = k \left( 1 - \frac{\Gamma_{\text{cat,surf}}}{\Gamma_{\text{cat,max}}} \right) e^{-\frac{\alpha F U}{RT}} \Gamma_{\text{cat,surf}}^\gamma \quad (36)$$

$$\frac{\Gamma_{\text{cat,surf}}}{\Gamma_{\text{cat,max}} - \Gamma_{\text{cat,surf}}} = K e^{\left(-\frac{F(U - U_{\text{pzc}})}{RT}\right)} e^{\left(-g \frac{\Gamma_{\text{cat,surf}}}{\Gamma_{\text{cat,max}}}\right)} C_{\text{cat,bulk}} \quad (37)$$

$$U_{\text{eff}} = U - U_{\text{pzc}} = U - U_{\text{pzc}}(\text{pH} = 0) - k_{\text{B}} T \ln 10 \text{pH}_{\text{surf}} \quad (38)$$

Finally, a collaborative experimental-computational study between our and Prof. Koper's research groups extended the previous insights on alkali cations to multivalent species, assessing cation effects on both  $\text{eCO}_2\text{R}$  and HER on polycrystalline gold at  $\text{pH} = 3$ . The DFT datasets related to this work are available on the ioChem-BD database.<sup>376</sup> Experimentally, activity and onset for water reduction were observed to correlate with cation acidity, defined



**Figure 19:** **a**, Gibbs free energy for \*H adsorption (diamonds) and  $\text{H}_3\text{O}^+$  dissociation (down-pointing triangles) vs cation acidity. **b**, Activation barrier for water dissociation (up-pointing triangles) vs cation acidity. **c**, Normalized activity for water reduction estimated from experimental data vs cation acidity. The qualitative fit decouples the contributions from water dissociation kinetics (red) and cation accumulation (shades of blue) for alkali (1+), divalent (2+), and trivalent (3+) species. **d**, Theoretical prediction of  $\text{eCO}_2\text{R}$  performance for different cationic species using cation- $\text{CO}_2$  coordination number as activity descriptor. **e**, Heat map of  $\text{CO}_2$  reduction (purple shades) and  $\text{H}_2\text{O}$  reduction (red shades) performances at high overpotential vs cation ionic radius and cation acidity.<sup>42</sup> Adapted with permission from Ref. 42 (Copyright 2022 American Chemical Society).

as the ratio of charge and ionic radius, while no effect was found on proton reduction. Thus, significant  $\text{eCO}_2\text{R}$  activity in presence of acidic cations was reported only at low applied potential, for negligible water reduction, while  $\text{Ba}^{2+}$  and  $\text{Cs}^+$  accounted for the highest  $\text{CO}$  formation rates at high overpotential. To rationalize these evidences, we applied a similar computational framework as for our previous study,<sup>45,371</sup> *i.e.* AIMD simulations at  $T = 300$  K on a gold supercell with explicit solvation previously optimized and different cations ( $\text{Li}^+$ ,

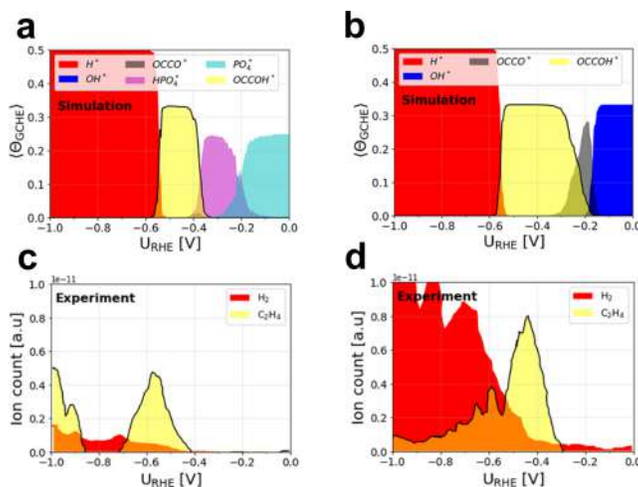
Cs<sup>+</sup>, Mg<sup>+</sup>, Ba<sup>2+</sup>, Al<sup>3+</sup>). Electric field effects ( $\vec{E} = -0.3 \text{ V \AA}^{-1}$ , roughly corresponding to  $-0.7 \text{ V vs SHE}$ ) and realistic cation concentration (2 cations, 0.07 ML coverage, 1.0-1.6 M) were assumed in the model, which was validated by comparing experimental and theoretical values for cation coordination shell and hydrated radius.<sup>362,377</sup>

The explicit insertion of electric field effects enables the observation of electrostatic repulsion among acidic cations, suggesting that nonacidic cations accumulate more at the interface. Specifically, we observe that the gradient of the thermodynamic driving force for cation accumulation correlates with cation acidity, which then acts as a proper descriptor of cation trends among different species. As for the rationale behind specific cation promotion of hydrogen evolution, we assessed the energetics related to the dissociation of H<sub>3</sub>O<sup>+</sup> and H<sub>2</sub>O, expected to be the proton sources at acidic and neutral/alkaline surface pH,<sup>41</sup> and \*H adsorption, typically assumed as the key descriptor for HER activity.<sup>11</sup> H<sub>3</sub>O<sup>+</sup> dissociation is kinetically barrierless and thermodynamically favored for any assessed cation, and the Gibbs free energy related to this step scales linearly with cation acidity, Figure 19a. Instead, \*H adsorption is not affected by the change of cation (Figure 19a), thus both these evidences motivate the absence of any specific cation effect on proton reduction. The activation energy for water dissociation strongly correlates with cation acidity, ranging from  $E_a$  larger than 1.5 eV for Cs<sup>+</sup> till negligible activation on acidic cation (Figure 19b). Thus, cation acidity rules water reduction due to its impact on cation accumulation and water dissociation kinetics, leading to the definition of a HER activity volcano plot (Figure 19c). Alkali and bivalent cations exhibit similar HER activity since they account for low water dissociation rate, thus cation accumulation at the interface has a negligible promoting role due sluggish kinetics. Instead, acidic cations enable high H<sub>2</sub>O dissociation rates, thus HER activity is mainly driven by the cation accumulation trend within the trivalent group, motivating the highest performance for Nd<sup>3+</sup>, present at higher surface concentrations than Al<sup>3+</sup>. In addition to defining a general mechanism for cation effect on water reduction, this study highlights the interplay between promotional effects on CO<sub>2</sub>R and HER and the need for an

integrated assessment of both contributions. In fact, even though the validity of cation-CO<sub>2</sub> coordination number was here generalized to multivalent cations (Figure 19d) it serves its scope only at low overpotential, where water reduction was negligible, Figure 19e.

## 7.2 Anion effects

Anion species in the electrolyte can poison the electrode surface under cathodic conditions,<sup>174,175</sup> limiting the performance of optimized devices such as Gas Diffusion Electrode due to salt formation at the catalyst layer.<sup>15,176</sup> Prof. Rossmeisl and his group have successfully employed the GCHE framework (Section 2.2)<sup>57,58</sup> to quantify poisoning effects in CO<sub>2</sub> reduction and identify the stability region of adsorbed species vs applied potential and bulk pH,<sup>174,175</sup> showing excellent agreement with experimental reports.<sup>378</sup>



**Figure 20:** Surface poisoning of anions during electrocatalytic reactions. **a-b**, Simulated coverage for adsorbed species on Cu(100) at pH = 7 (left) and pH = 13 (right) derived from interface phase diagrams estimated *via* the GCHE formalism.<sup>57,58</sup> **c-d**, CO reduction products on Cu(100) in a H<sub>3</sub>PO<sub>4</sub> (pH = 7, left) and NaOH (pH = 13, right) electrolyte, respectively measured by Online Electrochemical Mass Spectrometry (OLEMS).<sup>378</sup> Adapted with permission from Ref. 174 (Copyright 2019 American Chemical Society).

The first application of the GCHE to assess explicit electrolyte effects aimed at rationalizing the pH dependence of CO reduction on copper and its lower onset potential than HER, reported in a previous experimental work for phosphate (pH = 7) and NaOH (pH

= 13) aqueous electrolytes.<sup>378</sup> The authors carried out AIMD simulations on copper facets with explicit water molecules, electrolyte species, and reaction intermediates.<sup>174</sup> Different \*H and \*OH coverages were sampled from zero to half coverages, while \*H<sub>x</sub>PO<sub>4</sub>, \*OCCO, and \*OCCOH were introduced individually in the supercell. The resulting configurations generated by AIMD were then screened through a Boltzmann weighting of the energies, thus allowing to estimate key parameters. In general, \*OCCO was reported stable only on Cu(100), because of large stabilization by the solvent. Besides, \*OCCO protonation to \*OCCOH or \*OCCHO was reported more favorable than \*H adsorption, thus motivating the lower onset potential for ethylene formation rather than HER. As for the dependence of ethylene selectivity on bulk electrolyte pH, both pH and applied potential were included in the energy of all the AIMD configurations through the GCHE scheme to define interface phase diagrams of the adsorbed species (Figure 20a,b). Besides showing excellent agreements with experimental results at pH = 7 (Figure 20c) and pH = 13 (Figure 20d),<sup>378</sup> simulations suggested that ethylene formation is limited by adsorption of anions (\*OH, \*HPO<sub>4</sub>, \*PO<sub>4</sub>) in the low overpotential region. Instead, at high overpotential, \*OCCOH adsorption is hampered by an increased \*H coverage. In the same study, carbonate poisoning was suggested to have a similar role for CO<sub>2</sub> reduction, although carbonate blocked copper facets until larger overpotential than that required for \*OH adsorption, thus motivating the difference in activity for CO and CO<sub>2</sub> reduction.<sup>350</sup> As a last conclusion from their study, the authors proposed that the experimental difficulty in observing the OCCO<sup>-</sup> intermediates are due to the very tiny potential window where this intermediate is stable before his protonation to \*OCCOH. Recently, the same authors carried out a similar investigation of the CO reduction process on Cu(100) and Cu(111), combining voltammetric CO-displacement measurements and AIMD to characterize the Cu/electrolyte interface and the role of phosphate poisoning in tuning CO adsorption properties.<sup>175</sup> \*HPO<sub>4</sub> and \*PO<sub>4</sub> were suggested as the predominant adsorbed species on Cu(100) and Cu(111) respectively, limiting CO adsorption to more negative potentials.

### 7.3 Surface pH and bulk mass transfer

The Computational Hydrogen Electrode is a powerful tool to account for bulk pH effects, since this term is explicitly accounted through a linear correction to adsorption energies of intermediates (Section 2.2).<sup>8</sup> Hence, this framework has proven effective in rationalizing experimental evidences from Tafel plot analysis<sup>379</sup> and derive fundamental insights on reaction mechanisms.<sup>349</sup> However, recent technological and conceptual advances allow to measure experimentally the pH within the diffusion layer (the so-called “surface pH” or “local/localized pH”),<sup>380</sup> through the Scanning Electro-Chemical Microscopy (SECM) setup<sup>381,382</sup> (even time resolved),<sup>383</sup> spectroscopic strategies (e.g  $\text{HCO}_3^-/\text{CO}_3^{2-}$  vibrational bands),<sup>369,384</sup> and Rotating Ring-Disc Electrode (RRDE) voltammetry.<sup>385</sup> For cathodic reaction such as  $\text{CO}_2$  reduction, pH at the surface was reported to vary up to 5 units from the bulk value due to proton consumption or  $\text{OH}^-$  generation from  $\text{eCO}_2\text{R}$  and HER,<sup>16</sup> depending on the overall current density.<sup>384–386</sup> Variation of pH at the surface is only one of the relevant mass transfer effects occurring under electrocatalytic conditions,<sup>387,388</sup> thus models based on solving numerically the Nernst-Planck equations have been developed since the seminal work of Gupta *et al.*<sup>163</sup> Currently, two main theoretical approaches are employed to assess mass transfer within the diffusion layer and its influence on reaction activity and selectivity: (1) numerical solution of modified Poisson-Nernst-Planck (PNP) equations,<sup>164</sup> and (2) multiphysics modeling with finite element methods.<sup>167,389</sup>

As an example of the first framework, Bohra *et al.* investigated the  $\text{CO}_2$  reduction process on CO-producing catalysts, such as CO and Au, through a 1D model based on generalized solution of modified PNP equations.<sup>164</sup> Such generalization was needed to account for steric effects happening at realistic electrolyte concentrations. The authors only accounted for  $\text{CO}_2$  and  $\text{H}_2\text{O}$  reduction as Faradaic processes at the surface, while water dissociation and bicarbonate/carbonate/ $\text{CO}_2$  chemical equilibria reactions were considered within the diffusion layer. The bulk concentration of electrolyte species was assumed constant and estimated from experimental values. Three generalized modified Poisson-Nernst-Planck equations were

considered to model mass transfer within the diffusion layer, Equations 39-41.  $C_i$ ,  $\vec{J}_i$ ,  $z_i$ ,  $D_i$ , and  $a_i$  are respectively concentration, flux, charge, diffusion coefficient, and effective solvated diameter of the species  $i$ .  $R_i$  is the rate of production of species  $i$  from a  $p$  homogeneous reactions between water dissociation and bicarbonate/carbonate/ $\text{CO}_2$  equilibria.  $R$ ,  $F$ ,  $N_A$  are respectively Ideal Gas constant, Faraday constant and Avogadro number, while  $\Phi$  and  $T$  stand for potential vs the point of zero charge of the electrode and temperature.  $\epsilon_0$  and  $\epsilon_r$  are the vacuum and dielectric permittivity of the electrolyte, and  $\epsilon_r$  is approximated by a continuous function ranging from 80.1 (value for water at room temperature) to 6 (value for water at dielectric saturation).<sup>16</sup>

$$\frac{\partial C_i}{\partial t} = \nabla \cdot \vec{J}_i + \sum_p R_p \quad (39)$$

$$\vec{J}_i = -D_i \nabla C_i - \frac{D_i C_i z_i F}{RT} \nabla \phi - D_i C_i \left( \frac{N_A \sum_{i=1}^n a_i^3 \nabla C_i}{1 - N_A \sum_{i=1}^n a_i^3 C_i} \right) \quad (40)$$

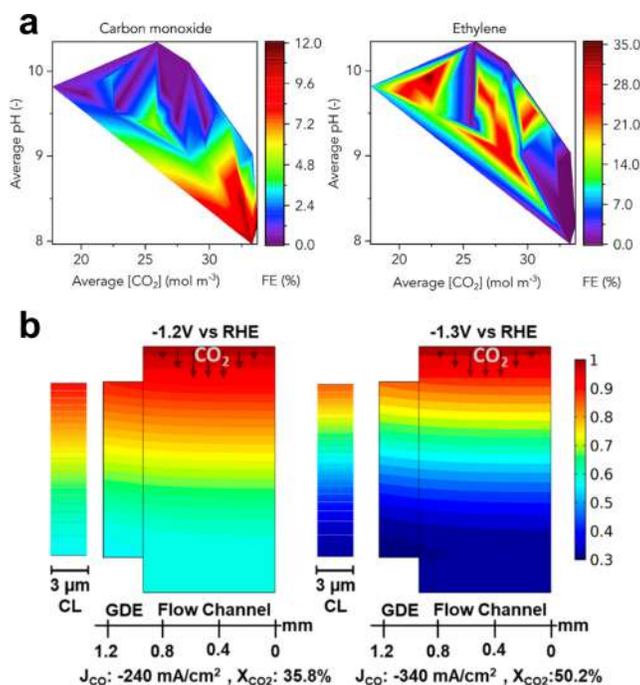
$$\nabla \cdot (\epsilon_0 \epsilon_r \nabla \phi) = -F \sum_{i=1}^n z_i C_i \quad (41)$$

Equations 39-41 were then numerically solved, assuming the electric field at the OHP as Neumann and Dirichlet boundaries conditions and the predictions from the GNPNP scheme were benchmarked with state-of-the-art models (reaction-diffusion and Poisson-Nernst-Planck). Several key parameters were calculated as continuous functions of applied potentials vs PZC and distances from the OHP: the effective potential, pH,  $\text{CO}_2$ , cation concentrations, etc.<sup>164</sup> Overall, significant variations of pH were reported already for low current densities ( $j = 10 \text{ mA cm}^{-2}$ ) together with very intense electric fields at the OHP ( $-0.7 \text{ V \AA}^{-1}$ ) and low water relative permittivity ( $\epsilon_r = 50$ ) even at mild applied potential ( $-0.32 \text{ V vs PZC}$ ).<sup>164</sup>

Veenstra *et al.* mapped  $\text{CO}_2$  product distribution with surface pH and applied potential through a joint experimental/computation study.<sup>167</sup> First, they synthesized well-defined microstructured CuO electrodes, generating surface cavities with tunable length through Laser ablation ( $L = 0, 20, 40, 60, 100, 130 \mu\text{m}$ ). Using these cavities as micro-probes for surface

pH, they characterized CO<sub>2</sub> selectivity in a 0.1 M KHCO<sub>3</sub> electrolyte. Then, they employed structural geometries and eCO<sub>2</sub>R performances as input data for multiphysics 3D modeling through the COMSOL platform to target concentrations and fluxes of key intermediates depending on the local morphology of the electrode. The computational scheme included eCO<sub>2</sub>R and HER, chemical equilibria for carbonate species, and diffusion, as in previous models.<sup>163,390</sup> Besides, the model explicitly considered the buffering role of cation hydrolysis suggested by Resasco *et al.*<sup>170</sup> The simulation accounted for three regions: (1) the volume of the surface cavity, (2) eight adjacent cavities to encompass potential contributions from surrounding environment, (3) a diffusion layer with tunable thickness depending on the mass transfer regime, initially set to 50  $\mu\text{m}$ . Bulk concentrations and zero flux for non-reactive species were imposed as boundaries conditions beyond the diffusion layer. Since CO<sub>2</sub> and pH did not show significant variation across the  $(x, y)$  planes, the initial 3D model was simplified to a 1D setup, only considering the  $z$ -direction. A pH increase up to 3 units was calculated at the end of the cavities, and this effect strongly depended on applied potential and length of the micro-probes. By relating computational estimations with experimental results for eCO<sub>2</sub>R selectivity, the authors generated selectivity maps for different reaction products, unveiling the role of surface pH and average CO<sub>2</sub> concentration at the electrode (Figure 21a highlight the case of carbon monoxide and ethylene). These separate analysis were later combined into a general selectivity maps, which lead to remarkable fundamental insights on reaction mechanisms toward C<sub>1</sub>, C<sub>2</sub> and C<sub>3</sub> on copper.<sup>167</sup>

Kas *et al.* employed an analogous multiphysics framework to model CO<sub>2</sub> reduction on a GDE with Ag catalyst layer.<sup>389</sup> The 2D model accounted for Butler-Volmer kinetics for both eCO<sub>2</sub>R and H<sub>2</sub>O,<sup>2</sup> as well as CO<sub>2</sub>/bicarbonate/carbonate equilibria reactions within the electrolyte. The input parameters for modeling the GDE were retrieved from experimental characterization.<sup>391</sup> Overall, significant variations of CO<sub>2</sub> concentrations and partial current densities were estimated within the GDE area, both across ( $x$ -direction) and along ( $y$ -direction) the flow channel (Figure 21b). Whilst the first variation is driven by concentra-



**Figure 21:** Surface pH and mass transfer effects in electrochemical CO<sub>2</sub> reduction. **a**, Heat maps for CO and C<sub>2</sub>H<sub>4</sub> F.E. vs surface pH and CO<sub>2</sub> concentration.<sup>167</sup> White areas correspond to pH/[CO<sub>2</sub>] pairs not allowed under the given reaction conditions. **b**, Heat plots of gaseous CO<sub>2</sub> mole fraction in the gas flow channel and gas diffusion electrode for different applied potentials, 5 mL min<sup>-1</sup> flow rate, and 1 cm<sup>2</sup> electrode area. Data employed in the model, (partial current density  $j_{CO}$  and single-pass conversion  $X_{CO_2}$ ) are defined below each panel, whilst gaseous mole fractions of CO<sub>2</sub> in the catalyst layer (CL) are reported in the inset.<sup>389</sup> Adapted with permission from Ref. 167 (Copyright 2020 Elsevier) and Ref. 389 (Copyright 2021 American Chemical Society).

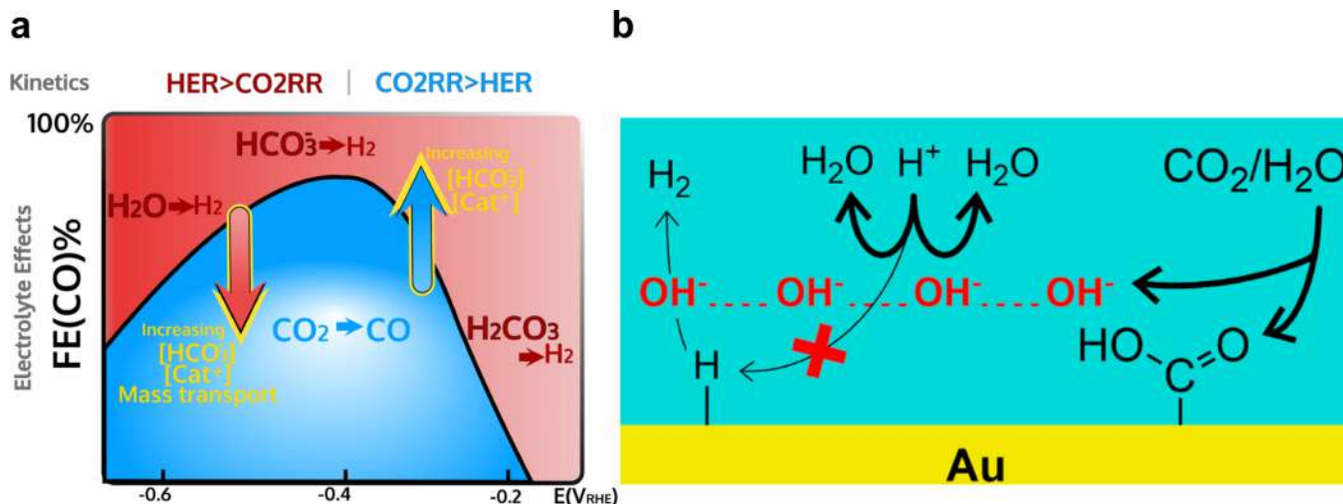
tion gradients due to CO<sub>2</sub> consumption at the catalyst layer, the second is due to the forced convective flow from the inlet. Instead, CO<sub>2</sub> concentrations and partial current densities were almost constant within the catalyst layer due to its small thickness (3 μm). Nevertheless, dramatic pH variations were calculated within the catalyst layer, leading to pH ~ 13 for current densities higher than 200 mA cm<sup>-2</sup> and reduction of anions diffusion thickness from 200 μm to few μm. Finally, the COMSOL model highlighted the existence of an interplay between applied potential, current density, and outlet concentrations. While high CO<sub>2</sub> flow rate leads to increased production rate due to enhanced mass transfer, it also determines larger ohmic loss and operational costs and lower single-pass conversions, thus limiting the industrial exploitation of such operating conditions.<sup>199</sup>

As a final remark in this section, we highlight that mass transfer effects can be crucial under well-defined conditions and may determine selectivity shifts due to interplay between different reaction precursors. Recently, Marcandalli *et al.* rationalized the Bell-shaped Faradaic efficiency of eCO<sub>2</sub>R to CO by identifying three different HER regimes (Figure 22a).<sup>339</sup> Through mechanistic investigations by a RRDE voltammetry, HER was suggested to occur through carbonic acid reduction at very low overpotentials. At more negative potential H<sub>2</sub>CO<sub>3</sub> reduction is presumably diffusion-limited, hence bicarbonate becomes the relevant proton donor. Until  $-0.5$  V vs RHE, CO<sub>2</sub> reduction to CO is increasingly promoted, due to the higher kinetics for this process than for HER. Lastly, water reduction initiates at more negative potentials and HER increases to the detriment of eCO<sub>2</sub>R, due to the higher kinetics for HER in this regime. Thus, ideal catalysts for eCO<sub>2</sub>R should be optimized within the low overpotential region ( $\leq -0.5$  V vs RHE), where this process is significantly more favorable than HER from bicarbonate and carbonic acid reductions.

In addition to this elegant work, the same research group demonstrated by Differential Electrochemical Mass Spectroscopy (DEMS) that HER can be entirely suppressed on gold under mildly acidic pH due to proton consumption by OH<sup>-</sup> generated by eCO<sub>2</sub>R (Figure 22b).<sup>340</sup> Hence, Faradaic efficiencies close to 100% can be achieved by carefully tuning CO<sub>2</sub> and H<sup>+</sup> surface concentrations so that excess OH<sup>-</sup> from eCO<sub>2</sub>R is fully neutralized by protons and does not undergo CO<sub>2</sub>/CO<sub>3</sub><sup>2-</sup> homogeneous reaction. Both fundamental approaches propose mass transfer optimization strategies which are scalable to industrial devices, as demonstrated in Refs. 179,359, thus highlighting even further that these are crucial parameters in electrocatalytic reactions and must be properly accounted for.

## 8 Application of Machine Learning

Since the advent of machine learning (ML) in the field of catalysis, over the past few decades this technique has been used to address shortcomings of atomistic simulations, such as



**Figure 22:** **a**, Bell-shaped Faradaic efficiency of  $\text{CO}_2$  reduction to CO on a gold electrolyte versus applied potential on bicarbonate electrolyte. Depending on the cathodic bias, carbonic acid, bicarbonate, and water are the main protons donors for HER.<sup>339</sup> **b**,  $\text{CO}_2$  partial pressure suppresses hydrogen evolution by consuming protons at the surface to form water, leading to F.E close to 100% for  $\text{CO}_2$  reduction on gold in acidic pH.<sup>340</sup> Adapted with permission from Ref. 339 (Copyright 2021 American Chemical Society) and Ref. 340 (Copyright 2021 American Chemical Society).

quantum mechanical (QM) and molecular mechanics (MM) methods.<sup>392</sup> On one hand, QM-based simulations are accurate and transferable, yet they are applicable only to small system sizes and short time scales. On the other hand, empirical MM-simulations are developed *ad hoc* for specific systems and thus are typically not transferable. As defined by Morawietz and Artrith in a nice recent review,<sup>392</sup> QM-based machine learning aims at increasing the computational efficiency of QM-based methods, without losing their transferability. Applications of machine learning to electrocatalysis usually falls into one of these four scopes:

1. Prediction of catalytic properties from datasets obtained through QM-based methods. This strategy consists of the automatic generation and analysis of QM data through ML or training of ML methods over QM data to predict relevant parameters and trends.
2. Extension of the range of applicability of QM-based methods to reach larger simulation cell size and time scale. This approach is based on the development of machine learning atomistic potentials to achieve analogous accuracy as QM-strategies.

3. Inverse Design, inversion of atomistic simulations (or experimental data) to extract atomic structures from a given set of properties.

To date, application of ML in the field of eCO<sub>2</sub>R has been mainly devoted to pursuing strategy (1), *i.e.* material discovery and data analysis, as we will briefly mention below. As a successful example of integration of both approaches, we cite the work by Ulissi *et al.*,<sup>393</sup> where thermodynamic/kinetic Linear Scaling Relationships and automatic identification of rate-determining steps were applied to study reaction of syngas to CO<sub>2</sub>, water, methane, methanol, acetaldehyde, and ethanol. For additional examples and a more in-depth review over these topics, we refer the readers to two recent reviews by Zhang *et al.*<sup>394</sup> and Mistry *et al.*<sup>262</sup> Instead, we here focus on the development of ML potentials to enhance the space and time scales of QM-simulations, strategy (2) discussed in Section 8.2), and data-driven structural analysis, strategy (3) discussed in Section 8.2. Finally, in Section 8.4 we give a short outlook on the future challenges in the field.

## 8.1 Material discovery and data analysis

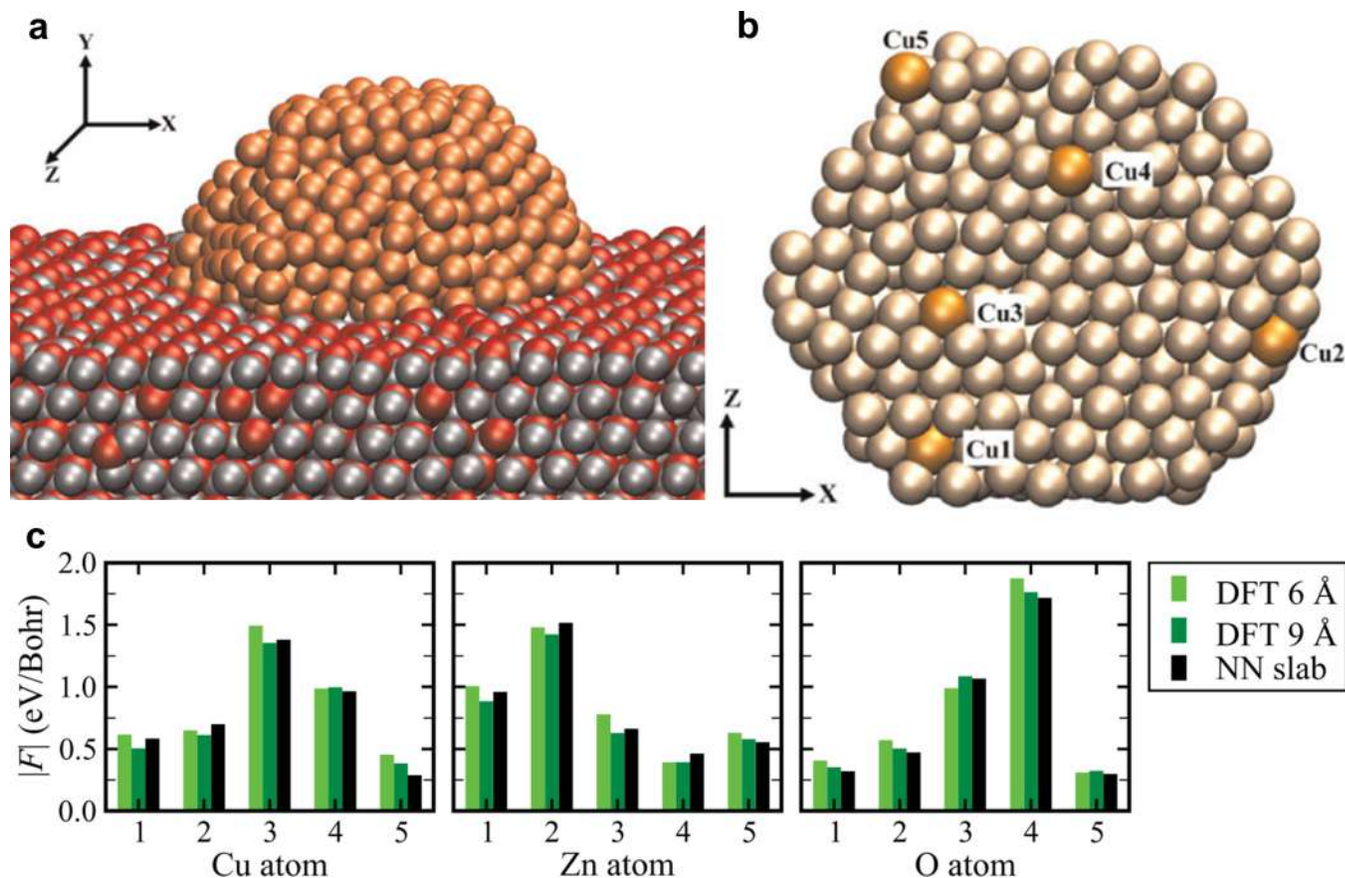
In one of the first application of ML to eCO<sub>2</sub>R, Li *et al.* designed a framework to predict limiting potential for CO<sub>2</sub> reduction to C<sub>1</sub>-C<sub>2</sub> products from  $\sim 250$  *ab initio* CO adsorption energies on (100)-terminated multimetallic Cu catalysts.<sup>395</sup> The properties of the adsorbing sites were parameterized based on their electronegativity and generalized coordination numbers and these parameters plus *ab initio* data were employed as input for a neural network with 75% of the dataset as training set and 25% for testing the model. Finally, linear scaling relationships were used to estimate binding energies of other C<sub>1</sub>-C<sub>2</sub> intermediates from CO adsorption energy and consequently to define a list of promising alloys with mild CO adsorption, as Cu-Ni-Cu<sub>3</sub>Y, Cu-Ni-Cu<sub>3</sub>Sc, Cu-Rh<sub>3</sub>Ti, Cu-Rh-Cu<sub>3</sub>V, and Cu-Rh-Cu<sub>3</sub>Mo.<sup>395</sup> The same research group applied a similar approach to screen over 1000 model alloys and calculate \*CO and \*OH binding energies on these structures, which were later used to predict reactivity towards methanol oxidation.<sup>396</sup> Independently, Ulissi, Sargent, and co-workers ap-

plied ML methods to screen through large number of intermetallic alloy compositions using CO binding energy as a descriptor for catalytic activity,<sup>397</sup> and eventually predicted Cu-Al alloys to contain multiple surface orientations with almost optimal CO binding energies.<sup>398</sup> The proposed Cu-Al compositions were then tested under eCO<sub>2</sub>R conditions and showed very high Faradaic efficiencies for ethylene, close to 85%.

## 8.2 Development of machine learning atomistic potentials

Since the pioneering works of Behler, Parrinello, and co-workers,<sup>399,400</sup> atomistic potentials generated through neural network have been successfully employed in electrocatalysis, as nicely reviewed in Ref. 392. Representation through Neural network of potential-energy surfaces as a function of the atomic positions has enabled faster assessment of different electrocatalysts, such as supported metal nanoparticles,<sup>401</sup> metal clusters on oxides,<sup>400</sup> Cu/Au nanoparticles.<sup>402,403</sup> This approach has been extended to fully ML-based<sup>211</sup> or hybrid molecular dynamics simulations,<sup>403</sup> in which ML is employed on limited numbers of MD snapshots to predict relevant properties of the system.<sup>404</sup> In this section, we highlight one of the first applications of neural network atomistic potentials to large size ternary system as Cu/ZnO,<sup>400</sup> while we refer the reader to two reviews by Behler for detailed insight into the mathematical formalism.<sup>405,406</sup>

Cu/ZnO materials have been extensively studied due to their reactivity for thermal CO<sub>2</sub> hydrogenation to methanol<sup>407</sup> and selectivity for eCO<sub>2</sub>R toward ethanol.<sup>408,409</sup> Nevertheless, an adequate computational assessment of defects, atomic rearrangements, and local stoichiometry is hindered by the large configuration space required and the low computational efficiency of DFT simulations. Thus, Artrith, Hiller, and Behler developed a framework to obtain neural network atomistic potentials from DFT energies of about 100000 well-defined Cu and ZnO structures, such as clusters, bulk, and surfaces. Their approach was validated first individually on Cu and ZnO. Surface energies estimated through NN-potentials for low index Cu facets laid within less than 5 meV Å<sup>-2</sup> from DFT values and bulk properties of



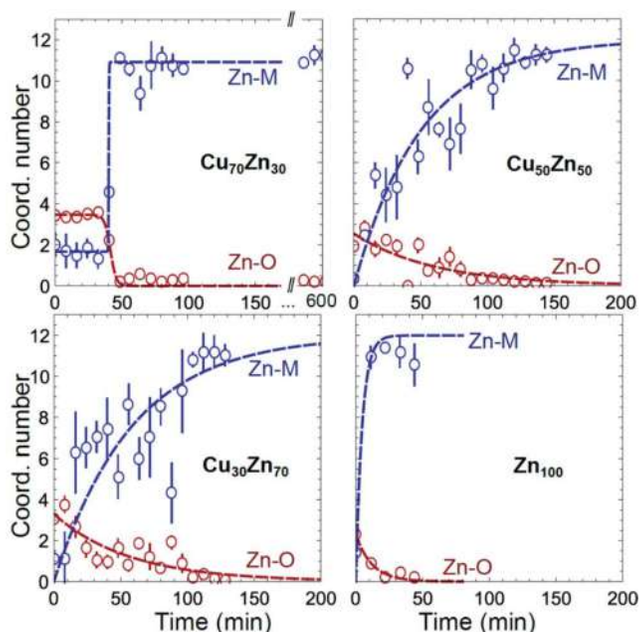
**Figure 23:** **a**, Snapshot of molecular dynamics simulation at  $T = 1000$  K of Cu<sub>612</sub> cluster on ZnO(10 $\bar{1}$ 0) surface. **b**, bottom-view of Cu cluster and selected sites at the Cu/ZnO interface. **c**, Neural network forces acting on selected atoms at the Cu(111)/ZnO(10 $\bar{1}$ 0) vs DFT forces obtained for clusters centered at these atoms with radius of 6 Å and 9 Å respectively.<sup>400</sup> Adapted with permission from Ref. 400 (Copyright 2013 Wiley-VCH).

ZnO were reproduced in excellent agreement as well. Besides, NN- and DFT-forces were equivalent for both systems. After benchmarking the neural network model, the potentials developed along the study were employed to simulate the interface between Cu and ZnO, modeled as a Cu<sub>612</sub> adsorbed on ZnO(10 $\bar{1}$ 0), Figure23a. Since the overall system contained 7524 atoms, its assessment through standard DFT would have been prohibitive instead. Local distortion was obtained through a NVT molecular dynamics simulation at 1000 K and five atoms at the interface were selected for each species (Cu, Zn, O), as shown in Figure23b for Cu. Comparison between NN-potential forces and DFT-forces for these sites confirmed once more the robustness of the framework (Figure 23c), suggesting a potential generalization

beyond ternary systems toward the inclusion of adsorbates.<sup>400</sup>

### 8.3 Inverse Design to extract structural data

As introduced before, machine learning can assist the assignment of fingerprints from *in situ Operando* characterization techniques and the consequent derivation of key structural properties for disordered systems,<sup>410</sup> thus is increasingly employed in materials science.<sup>411</sup> To mention a meaningful example of this *inverse design* strategy, Timoshenko *et al.* applied



**Figure 24:** Time-dependent Zn-O and Zn-M ( $M=\text{Zn-Cu}$ ) coordination numbers as extracted by neural network from time-dependent Zn K-edge EXAFS data for CuZn and Zn<sub>100</sub> nanoparticles.<sup>412</sup> Adapted with permission from Ref. 412 (Copyright 2020 The Royal Society of Chemistry with Creative Commons Attribution 3.0 Unported Licence<sup>413</sup>).

neural network to a set of Extended X-ray Absorption Fine Structure (EXAFS) data to extract structural information for CuZn nanoparticles under CO<sub>2</sub> reduction.<sup>412</sup> By training the NN with theoretical spectra for a copper foil, Cu<sub>2</sub>O, CuO, and Cu(OH)<sub>2</sub>, the authors could reproduce experimental radial distribution function with excellent accuracy. After having validated the methodology, Cu-O, Cu-Cu, and Cu-Zn radial distribution functions were extracted from experimental Cu K-edge EXAFS spectra recorded for as-synthesized

CuZn nanoparticles and catalyst after 1-7 hours of continuous CO<sub>2</sub> reduction. The authors tracked the variation of Cu-Cu/Zn distances over time as well as the coordination number for Zn-Cu/Zn and Zn-O (Figure 24). These structural properties were finally employed to rationalize the overall performance of the catalyst.<sup>414</sup>

## 8.4 Outlook on machine learning

Even though ML techniques are certainly promising in the field of computational modeling of eCO<sub>2</sub>R reaction, its application is still limited. Thus, further efforts are needed to employ these novel techniques within a consistent and benchmarked framework. Since the development of a general set of protocols facilitates future potential advances, we endorse here the list of best practices proposed in a recent perspective article by Artrith *et al.*<sup>415</sup> For simplicity, we report the main points below.

1. **Data sources.** The available data sources should provide high quality, quantity, and diversity. Datasets should be stored on FAIR databases (see for instance ioChem-BD<sup>416-418</sup> the Materials Project<sup>419</sup> and CatApp<sup>420</sup>).
2. **Data cleaning.** Data upload on accessible databases should involve cleaning steps to ensure that all the stored data are reusable. The cleaning steps should be properly described and preferentially applied through a semi/full automatic workflow.
3. **Data representation.** Different data representations should be considered to select a model and the choice should be properly described, motivated, and validated vs standard features sets.
4. **Model selection.** Since different approaches exist in ML, they should be properly accounted for and the choice of the model should be validated through baseline comparisons with simpler schemes.
5. **Model training and validation.** Parameters for training, validation, and test of the

model should be clearly stated. Besides, data splitting into the three categories should prevent data leakage.

6. **Code and Reproducibility.** The workflow should be reproducible, thus codes and related materials should be freely available in adequate long-term archiving.

Further, in terms of the four scopes previously introduced, material discovery for selective reduction of CO<sub>2</sub> to C<sub>2+</sub> hydrocarbons and alcohols should proceed hand in hand with determination of reaction mechanisms through multi-scale methods as microkinetic modeling. Successive high throughput screening studies should follow this preliminary assessment of the key reactions steps and intermediates. In our opinion such multi-scale approaches can greatly help the identification of generalized descriptors and increase the success rates towards finding better catalysts and improving the Faradaic efficiencies of various hydrocarbon products. As for the development of atomistic potentials, long-term MD simulations (of the order of nanoseconds) play a crucial role in determining active sites evolving under reaction conditions due to *Operando* restructuring of the catalyst (see Section 3.1). Since assessment of this process through *ab initio* MD simulations is limited to simplified models,<sup>205,210</sup> and may be affected by intrinsic inaccuracies<sup>212</sup> then accelerated ML-MD techniques will likely be determinant. The work from Behler, Artrith, and co-workers towards using artificial neural networks to simulate the potential energy surfaces is a major step in this direction.<sup>392,405,406</sup> Finally, novel techniques from data extraction and inverse design will significantly lower the impact of human biases in analysis of experimental data, thus increasing the overall accuracy of the resulting insights.

## 9 Conclusion and future challenges

The increasing accuracy of experimental methodologies to characterize electrochemical processes require significant progresses in computational modeling to mimic the full stack or device operation. In line with a terminology previously introduced,<sup>123</sup> *the eCO<sub>2</sub>R field needs*

*the development of an Operando modeling framework.*

On one side, the equations to obtain energies and chemical potentials from first principles simulations need to be more general, including promotional effects as external magnetic fields,<sup>421,422</sup> local surface polarization,<sup>97,210</sup> and specific stabilization from the electrolyte.<sup>45</sup> On the other hand, several issues need to be addressed regarding the materials and operation gap. In many cases, new techniques based on machine learning<sup>423</sup> can help integrate the different scales of the electrochemical processes and can provide strategies for dimensionality reduction, allowing unambiguous determination of descriptors for activity.<sup>424</sup>

Reconstruction, phase transitions, dissolution, and re-precipitation of species evolving from the as-synthesized compound requires the development of new computational models that are able to describe the effects related to stability. Novel methods will certainly benefit from the fast development of ionic potentials and artificial neural networks which will be crucial towards large scale molecular dynamics. The initial results in this field are very promising but the introduction of solvent and electrolyte has not been successfully implemented yet. Moreover, the characterization of pattern distribution and ensembles through graph theory and other ML approaches will be fundamental to identify the most active and selective sites.

Detailed modeling of complex reaction mechanisms, as CO<sub>2</sub> reduction to C<sub>3+</sub> products, requires significant computational advances toward the automatic generation of reaction networks, through graphs and autonomous energy evaluations with robust workflows which include human analysis of ambiguous calculations (from our own experience these ill-defined datasets account for about 15% of the total). Such frameworks should be preferentially integrated into multi-scale modeling techniques as microkinetic modeling to evaluate the most abundant surface intermediates and selectivity determining steps. Rationalization of pathways and fast evaluation of barriers linking key intermediates are crucial for further analysis of complex reaction networks. ML-based dimensionality reduction schemes can be employed to simplify the overall complexity and go beyond the standard definition of

catalytic descriptors.

Under electrocatalytic conditions, once the potential is applied, the material reacts and the applied bias also affects the solvent/electrolyte. Apart from cathodic/anodic potential, surface polarization is further influenced by the explicit interactions between cations and anions in the electrolyte or by their crossover through the membrane between cathodic and anodic compartments. Mass transfer phenomena both at the electrode/electrolyte interface and at the bulk electrolyte need to be considered for an accurate modeling of the operational device. In this area, microkinetic codes require a better standardization along with the fluid dynamic approaches. However these two approaches currently are difficult to couple, since the engineering programs lack a successful integration of first principles results.

Mapping experiments to computational results and establishing robust structure-activity relationships can only be done through a much closer interplay between experiments and theory, moving from qualitative or semi-quantitative analysis (e.g DFT vibrations for spectroscopy) to more adequate fingerprint identifications. Bridging technologies, like data processing of both experimental and computational datasets through neural network, will be instrumental to merge all the structural information in a seamless framework, thus enabling a successful identification of the active sites.

All these developments can be further steered by storing theoretical and experimental output on FAIR databases to promote reproducibility and reuse of the data. This is much more advanced in the area of first principle modeling where our approach ioChem-BD<sup>416–418</sup> and other initiatives such as the Materials Project,<sup>419</sup> CatApp,<sup>420</sup> etc. are getting momentum, while FAIR storage of experimental data is currently limited to few platforms (e.g. Zenodo<sup>?</sup> Generated datasets can be reused for heavy calculations or fast estimates, as we did for modeling the metal/electrolyte interface,<sup>22,42,45</sup> and then further rationalized through machine learning.<sup>28</sup>

Overall, the developments in the field during the last five years have been enormous and yet the main core questions remain open. A multidisciplinary approach, supported by new

researchers educated in the converging areas of Physics, Chemistry, Data, and Materials Sciences, will improve our understanding of the complexity of *Operando* electrochemical processes, hence enabling a full exploitation of fundamental insights in industrial devices.

## Notes

The authors declare no competing financial interests.

## Biographies

**Federico Dattila** Dr. Federico Dattila is a postdoctoral researcher in the Catalytic Reaction Engineering for Sustainable Technologies (CREST) group at the Polytechnic of Turin since March 2022. Previously, he worked in the group of Prof. Núria López at the Institute of Chemical Research of Catalonia (ICIQ, Barcelona Institute of Science and Technology, Tarragona, Spain) for FlowPhotoChem (grant-ID: 862453). His research focuses on modeling electrochemical CO<sub>2</sub> and CO reduction on transition metal catalysts. In 2020 he got his Ph.D. in Chemical Science and Technology at the Rovira i Virgili University under the co-supervision of Prof. Núria López and Dr. Rodrigo García-Muelas. During his Ph.D. he carried out two short stayings at the University of Leiden (Prof. Marc Koper's group) and at Avantium Chemicals BV (Dr. Klaas Jan P. Schouten's group). Prior to his Ph.D., he got a M.Sc. in Physics at the University of Turin in 2017, with a thesis on photoelectrochemical cells developed at Chalmers University of Technology (Gothenburg, Sweden).

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**Yecheng Zhou** Prof. Yecheng Zhou received the Ph.D. degree in Chemistry under the supervision of Dr. Angus Gray-Weale from The University of Melbourne, Australia, in 2017. After graduation, he has worked in the Department of Physics in Southern University of Science and Technology (Guangdong, China) as a research assistant professor and worked as a postdoctoral researcher in the group of Prof. Núria López at the Institute of Chemical Research of Catalonia (ICIQ), within the Barcelona Institute of Science and Technology (BIST) in Tarragona, Spain). At the end of 2019, he was appointed as an associate professor at Sun Yat-Sen University in China. His research interest is theoretical modeling of defect and surface chemistry in renewable energies, including perovskite and organic solar cells, and electrocatalysis.

**Núria López** Prof. Núria López has been active in the field of theoretical heterogeneous catalysis for the last 25 years. She obtained her Ph.D. at the University of Barcelona, Spain in 1999. Her post-doctoral studies were carried out in the group of Prof. Jens Nørskov at the Technical University of Denmark. Since 2005 she is group leader of the theoretical heterogeneous catalysis group at ICIQ. She has participated in about 250 publications in the areas of (Photo-Electro) Catalysis and collaborates with experimental groups worldwide.

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# Abbreviations

DFT = Density Functional Theory

LSR = Linear Scaling Relationships

CHE = Computational Hydrogen Electrode

EXAFS = Extended X-ray Absorption Fine Structure

eCO<sub>2</sub>R = Electrochemical CO<sub>2</sub> reduction

C<sub>n+</sub> = Chemical compounds from CO<sub>2</sub> reduction with carbon number  $\geq n$ .

HER = Hydrogen Evolution Reaction

OER = Oxygen Evolution Reaction

RDS = Rate-Determining Step

ORR = Oxygen Reduction Reaction

SHE = Standard Hydrogen Electrode

RHE = Reversible Hydrogen Electrode

PCET = Proton-Coupled Electron Transfer

ET = Electron Transfer

EDL = Electrical Double Layer

GCHE = Generalized Computational Hydrogen Electrode

ORR = Oxygen Reduction Reaction

PB = Poisson-Boltzmann

kLSR = kinetic Linear Scaling Relationships

tLSR = thermodynamic Linear Scaling Relationships

\* = adsorption site

\*molecule = adsorbed molecule

PCA = Principal Component Analysis

XPS = X-ray Photoelectron Spectroscopy

EC-ESTM = ElectroChemical Scanning Tunneling Microscopy

pc-Cu = Polycrystalline Copper

OD-Cu = Oxide-Derived Copper  
UHV = Ultra High Vacuum  
TPD = Temperature Programmed Desorption  
XAS = X-Ray Absorption Spectroscopy  
COR = CO reduction  
TR-SERS = Time-Resolved Surface Enhanced Raman Spectroscopy  
F.E. = Faradaic Efficiency  
GDE = Gas Diffusion Electrode  
LEED = Low-Energy Electron Diffraction  
NP = Nanoparticle  
ML = Machine Learning  
NN = Neural Network  
MD = Molecular Dynamics  
AIMD = Ab Initio Molecular Dynamics  
SCC-DFTB = Self-Consistent Charge Density Functional Tight Binding  
NVE = Microcanonical ensemble with no changes in moles, volume, and energy  
NVT = Microcanonical ensemble with no changes in moles, volume, and temperature  
ReaxFF = Reactive Force Field framework  
PZC = Point of Zero Charge  
ATR-FTIR = Attenuated Total Reflection Fourier Transform Infrared  
FTIR = Fourier Transform Infrared  
SEIRAS = Surface Enhanced Infrared Absorption Spectroscopy  
ATR-SEIRAS = Total Reflection Surface Enhanced Infrared Absorption  
PT = Proton Transfer  
SDS = Selectivity-Determining Step  
H<sub>2</sub>OR = Water reduction  
MMK = Microkinetic Modeling

CM = Quantum Mechanical

MM = Molecular Mechanics

CMH = Constrained Minima Hopping

OHP = Outer Helmholtz Plane

OLEMS = Online Electrochemical Mass Spectrometry

PNP = Poisson-Nernst-Planck

DEMS = Differential Electrochemical Mass Spectroscopy

FAIR = Findability, Accessibility, Interoperability, Reusability

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## Graphical TOC entry

