Structure- and Electrolyte-Sensitivity in CO₂ Electroreduction

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CONSPECTUS

The utilization of fossil fuels (i.e., coal, petroleum, and natural gas) as the main energy source gives rise to serious environmental issues, including global warming caused by the continuously increasing level of atmospheric CO₂. To deal with this challenge, fossil fuels are being partially replaced by renewable energy such as solar and wind. However, such energy sources are usually intermittent and currently constitute a very low portion of the overall energy consumption. Recently, the electrochemical conversion of CO₂ to chemicals and fuels with improved energy density driven by electricity derived from renewable energy has been recognized as a promising strategy towards sustainable energy.
The activation and reduction of CO$_2$, which is a thermodynamically stable and kinetically inert molecule, is extremely challenging. Although the participation of protons in the CO$_2$ electroreduction reaction (CO$_2$RR) helps lower the energy barrier, high overpotentials are still needed to efficiently drive the process. On the other hand, the concurrent hydrogen evolution reaction (HER) under CO$_2$RR conditions leads to lower selectivity toward CO$_2$RR products. Electrocatalysts that are highly active and selective for multicarbon products are urgently needed to improve the energy efficiency of CO$_2$RR. The reduction of CO$_2$ involves multiple proton-electron transfers and has many complex intermediates. Recent reports have shown that the relative stability of the intermediates on the surface of catalysts determines final reaction pathways as well as the product selectivity. Furthermore, this reaction displays a strong structure-sensitivity. The atomic arrangement, electronic structure, chemical composition, and oxidation state of the catalysts significantly influence catalyst performance. Fundamental understanding of the dependence of the reaction mechanisms on the catalyst structure would guide the rational design of new nanostructured CO$_2$RR catalysts. As a reaction proceeding in a complex environment containing gas/liquid/solid interfaces, CO$_2$RR is also intensively affected by the electrolyte. The electrolyte composition in the near surface region of the electrode where the reaction takes place plays a vital role in the reactivity. However, the former might also be indirectly determined by the bulk electrolyte composition via diffusion. Adding to the complexity, the structure, chemical state, and surface composition of the catalysts under reaction conditions usually undergo dynamic changes, especially when adsorbed ions are considered. Therefore, in addition to tuning the structure of the electrocatalysts, being able to also modify the electrolyte provides an alternative method to tune the activity and selectivity of CO$_2$RR. In situ and operando characterization methods must be employed in order to gain in depth understanding on the structure- and electrolyte-sensitivity of real CO$_2$RR catalysts under working conditions.

This Account provides examples of recent advances in the development of nanostructured catalysts and mechanistic understanding of CO$_2$RR. It discusses how the structure of a catalyst (crystal orientation, oxidation state, atomic arrangement, defects, size, surface composition, segregation, etc.) influences the activity and selectivity, and how the electrolyte also plays a determining role in the reaction activity and selectivity. Finally, the importance of in situ and operando characterization methods to understand the structure- and electrolyte-sensitivity of the CO$_2$RR is discussed.
1. Introduction

The electrochemical conversion of CO₂ provides a potential solution to address some of the future energy needs by converting greenhouse gases into valuable fuels or feedstocks. The use of renewable energy sources to supply the electrons needed to electroreduce CO₂ makes this process attractive, so that the intermittent electricity coming from solar, wind, or hydro-power technologies could be complemented by the energy stored in the chemical bonds of high-energy-density products coming from captured CO₂. However, this molecule is extremely stable, being the CO₂ electroreduction reaction (CO₂RR) an uphill process that requires high overpotentials and presents numerous other challenges. The multiple electron transfer steps required to reduce CO₂ into fuels, accompanied by the competing hydrogen evolution reaction (HER), leads to poor selectivity towards multicarbon compounds (e.g. ethylene and ethanol). The later species and other so-called C₂+ products are highly sought due to their higher value, greater energy density, and broader applicability than C₁ products. Therefore, the development of highly active and selective catalysts, as well as an optimum technology that improves the effective CO₂ conversion, are key challenges for practical application of the CO₂RR.

Among all metals, Cu is the only metal capable of efficiently reducing CO₂ to hydrocarbons and alcohols, while hydrogen, CO, and formate are the major products produced by other metals.¹ The CO₂RR is a structure-sensitive process, meaning that the atomic arrangement at the catalyst surface affects the adsorption and/or bond cleavage of the reactants, thus determining the catalytic performance. Recent reports have opened new perspectives in exploring both compositional and morphological effects towards the rational design of more selective catalysts. Moreover, optimizing the reaction environment has also been demonstrated to be critical to address some or the challenges of this reaction, such as the low solubility of CO₂, changes in pH, and stabilization of certain active species.

In this Account, we provide an overview of the current understanding of the CO₂RR, focusing on the strategies followed so far for the development of more efficient and selective electrocatalysts. Recurrent topics throughout our report are the need of having access to morphologically and chemically well-defined material systems in order to establish structure/chemical state/composition-reactivity correlations, and to able to study them in situ and under operando reaction conditions in order to gain mechanistic insight into CO₂RR.
2. From Molecular Catalysts to Single-Site Heterogeneous Catalysts

Due to the intrinsically high metal atom efficiency, molecular catalysts usually show high turnover frequency (TOF) for many chemical reactions, including CO\textsubscript{2}RR\textsuperscript{2,3}. Cobalt porphyrins incorporated in covalent organic frameworks exhibited 90 % Faradaic efficiency (FE) and TOF of 9400 h\textsuperscript{−1} with an overpotential of −0.55 V for CO\textsubscript{2}RR to CO\textsuperscript{2}. Single site heterogeneous catalysts, with isolated metal atoms embedded in a nitrogen-doped carbon matrix, have been explored for CO\textsubscript{2}RR\textsuperscript{4,−6}. Ju et al.\textsuperscript{4} studied the CO\textsubscript{2}RR performance of a family of single site, metal- and nitrogen-doped nanoporous carbons (Figure 1) containing active M-N\textsubscript{x} sites (M = Mn, Fe, Co, Ni, Cu). Fe-N\textsubscript{x} and Ni-N\textsubscript{x} single-site catalysts showed a unique reactivity and FE for reducing CO\textsubscript{2} to CO, comparable to the mass-based activity of state-of-the-art Au catalysts. Wang et al.\textsuperscript{5} further reported that single-site Co catalysts exhibit a remarkable activity and selectivity towards CO\textsubscript{2}RR if the Co-N coordination number decreases from 4 to 2, due to the facilitated activation of CO\textsubscript{2} to the CO\textsuperscript{2−} intermediate over low-coordinated sites.

![Figure 1](image)

**Figure 1.** (a) CO production rate of M-N-C catalysts vs. potential. Inset: schematic structure of the M-N-C catalysts. (b) Free energy diagrams of HER and CO\textsubscript{2}RR at −0.8 V\textsubscript{RHE} for M-N-C catalyst. Reproduced with permission from ref. 4. Copyright 2017 Nature Publishing Group.
3. Metal Cluster and Nanoparticle (NP) Size Effects

Metal clusters and NPs have been explored for CO₂RR due to their high surface-to-volume-ratio and large content of low-coordinated surface sites. For example, Au₂₅ clusters experience a charge redistribution induced by a reversible Au₂₅-CO₂ interaction⁷ that results in a 200-300 mV lower overpotential and a CO production rate 200-700 times higher as compared to 5 nm Au NPs and bulk Au. On the other hand, metal NPs usually show a size effect for CO₂RR.⁸⁻¹³ Enhanced catalytic activity, unfortunately accompanied by a drastic increase in the selectivity for H₂ and CO in detriment of hydrocarbon production, was observed with decreasing Cu NP size (2-15 nm), Figure 2a.⁸ Although size-dependent changes in the electronic structure of clusters are important (a few tens of atoms to about 100 atoms), for larger NPs as those in the previous example, size-dependent changes in surface atomic coordination are held responsible for the distinct catalytic reactivity. Thus, the increased population of low-coordinated surface sites on the smaller Cu NPs (Figure 2b) was linked to the increased activity and surging H₂ and CO production rates. Au NPs showed a similar size effect, but with a more drastic increase in activity with decreasing size,⁹ along with a decrease in CO selectivity which was explained by the weaker binding of the COOH reaction intermediate in the presence of large H coverages on the small Au NPs (Figure 2c).¹⁰ Interestingly, a simultaneous increase in current density and CO FE was observed for Pd NPs with decreasing size (Figure 2d).¹¹ The different behavior between Au and Pd NPs was attributed to the similar activity of the competitive HER on the terrace, edge and corner sites of the Pd NPs, instead of the more favored HER on the undercoordinated sites of the Au NPs.¹⁰
Figure 2. (a) Faradic selectivity and (b) population of specifically coordinated surface atoms as a function of Cu NP size. (c) Molar ratio of $\text{H}_2$ and CO produced at $-1.2 \text{ V}_{\text{RHE}}$ as a function of NP size. (d) CO FE of Pd NPs with different sizes. Reproduced with permission from refs. 8, 10. Copyright 2014 American Chemical Society. Reproduced with permission from ref. 11. Copyright 2015 American Chemical Society.

4. Interparticle Distance/Loading Effect

Apart from the nanoscale structure and composition, mesoscale phenomena such as interparticle reactant diffusion and readsorption of intermediates also play a key role in reactivity. Cu NPs with narrow NP size distributions and uniform arrangements were used as model material system. For the same average NP size, a trend of increasing current density with decreasing interparticle (IP) distance was found. Furthermore, on samples with small IP spacing the diffusive transfer between neighboring NPs as well as the re-adsorption and subsequent reduction of the CO
intermediate to hydrocarbons was favored, Figure 3. However, the important role of changes in the local pH resulting from the distinct depletion of reactants and intermediates on samples with different spacings cannot be overlooked. Subsequently, Wang et al. explored indirectly the IP distance effect on the selectivity of CO$_2$RR on CuO$_x$ NPs by changing the areal particle density. They found that high areal particle density (small IP distance) improved ethylene production and attributed this to dynamically favored CO re-adsorption and reactive *CO dimerization at elevated local interfacial pH. The IP distance effect was also observed for the oxygen reduction reaction over Pt nanoclusters. A distinct potential distribution in the electric double layer (EDL) between the nanoclusters was suggested to lower the adsorption strength of oxygenated species and hence improve the activity at small IP distances. However, this effect cannot explain IP-distance dependent CO$_2$RR data since the smallest distances studied ($\approx$ 20 nm) are significantly larger than the ones deemed relevant for the EDL overlap in Ref$^{16}$ ($\leq$ 3 nm). The role of the Cu loading in the dynamic evolution from densely packed Cu NPs to cube-like particles was also observed by Kim et al. and Manthiram et al. during CO$_2$RR.
Figure 3. (a) Simulation results of the CO$_2$ concentration distribution based on diffusion equations. (b) Potential-dependent current density on 4.7 nm Cu NPs with different spacings. (c) Faradaic selectivity of 4.7 nm Cu NPs as a function of the interparticle distance. Reproduced with permission from ref. 14. Copyright 2016 American Chemical Society. (d) Potential-dependent ethylene/methane FE ratio over CuO$_x$ NPs. Reproduced with permission from ref. 15. Copyright 2017 Wiley.

5. Bimetallic Catalysts

Bimetallic catalysts, with various compositions and atomic arrangements, might be used to tune the binding strength of intermediates, to enhance the reaction kinetics and change the reaction pathways for CO$_2$RR. The relatively low selectivity towards hydrocarbons and oxygenates on Cu NPs may be improved by designing bimetallic CuPd catalysts (Figure 4a). Phase-separated CuPd catalyst were found to achieve high selectivity (>60 %) for C$_2$, while ordered CuPd catalyst with an alternating Cu-Pd arrangement exhibited the highest selectivity for C$_1$ (>80 %).

Geometric/structural effects, as well as electronic effects induced after mixing Cu with Pd, appear to play an important role in determining the subsequent reaction pathway of the *CO intermediate. Similar geometric effects were also observed for AuCu NPs. In contrast to AuCu alloy NPs which were more active for HER, ordered AuCu NPs selectively converted CO$_2$ to CO with a FE of 80 % (Figure 4b). However, the negative potentials being used for the CO$_2$RR process are expected to induce significant segregation in bimetallic NPs, such as the formation of an Au-rich surface.

Along these lines, core-shell AuFe@Au NPs with surface defects stabilized by subsurface Fe were found to exclusively produce CO with a 100-fold increase in mass activity compared to Au NPs, Figure 4c. AgSn@SnO$_x$ core-shell NPs showed a volcano-like relationship between formate production and Sn concentration, attributed to the compromise of favorable stabilization of OCHO* by lattice expansion and the electron conductivity loss due to the increased thickness of the SnO$_x$ shell (Figure 4d). A greater alcohol to ethylene ratio was achieved by including sulfur atoms in Cu NPs. According to DFT calculations, the resulting Cu vacancies in the NP shell were responsible for the higher alcohol production.
In addition to NPs, bimetallic films were also used for CO$_2$RR. By varying the amount of Zn in Cu$_x$Zn film catalysts, the selectivity for ethanol versus ethylene production could be tuned, Figure 5a. It was discussed that CO spillover from Zn to adjacent Cu sites increased the population of free CO, which participates in the formation of ethanol by inserting itself into the bond between the Cu surface and *CH$_2$. Ni$_x$Ga$_y$ (Figure 5b) constitutes an example of hydrocarbon production on non-Cu materials. The introduction of Ga into the Ni films weakened the Ni-CO interaction,
resulting in a comparable CO binding strength as CO-Cu. The formation of more reduced products beyond CO was recently described to depend on an optimum binding of both, CO and H.\textsuperscript{27} This strategy was applied to Pd-modified-Au foils,\textsuperscript{28} which also produce hydrocarbons (< 5 % FE).

**Figure 5.** (a) Ethanol FE and ethanol/ethylene ratio as a function of the Zn content in Cu\textsubscript{x}Zn films. The insert is a SEM image of a Cu\textsubscript{4}Zn film. (b) Potential dependent hydrocarbon FEs for Ni\textsubscript{5}Ga\textsubscript{3} films. Reproduced with permission from refs. 25, 26. Copyright 2016 American Chemical Society.

6. **Facet Effect: from Single Crystal Electrodes to Shaped NPs.**

Over the past decades, single crystal studies have provided in depth insight on the influence of the surface structure and order in electrocatalytic processes of technical interest.\textsuperscript{1} As aforementioned, Cu is unique at converting CO\textsubscript{2} into hydrocarbons and alcohols, mainly due to its moderate CO binding and the stabilization of certain intermediates.\textsuperscript{29} Moreover, the product selectivity is highly dependent on the crystalline orientation: Cu(111) yields mainly methane, while Cu(100) favors ethylene, and Cu(110) promotes the production of ethanol, acetate and acetaldehyde.\textsuperscript{1,30} Theoretical studies have shed light into the reaction mechanisms and intermediates responsible for the different selectivity on (111) and (100) surfaces\textsuperscript{31}, showing that the atomic configuration of the Cu(100) surface is the most favorable towards the pathway that yields ethylene. This mechanism was argued to take place via CO dimerization, Figure 6a. In this sense, high surface coverage of adsorbed CO* was found essential for the selective formation of
ethylene (Figure 6b).\textsuperscript{32} Moreover, Cu(751) surfaces show higher selectivity for C\textsubscript{2+} products at lower overpotentials than lower indexed surfaces, Figure 6c.\textsuperscript{33} However, the role played by the distinct facets in CO\textsubscript{2}RR is not fully understood, and further studies using model surfaces are still needed to understand the combined effect of surface structure and other parameters such as oxygen content.

**Figure 6.** (a) Deconvolution of the first electron–proton transfer on Cu(100). Reproduced with permission from ref. 31. Copyright 2013 Wiley. (b) Relationship between ethylene formation and CO evolution on Cu(100). Reproduced with permission from ref. 32. Copyright 2017 American Chemical Society. (c) Oxygenate/hydrocarbon ratios as a function of potential for Cu(111), Cu(751), and Cu(100). Reproduced with permission from ref. 33. Copyright 2017 Proceedings of the National Academy of Science.
Nanoparticles synthesized with a particular morphology generally have a well-defined surface structure. Therefore, tailoring the shape of the NPs can lead to materials with superior activity and selectivity. Facet effects on metals like Pd\textsuperscript{34}, Ag\textsuperscript{35} and Au\textsuperscript{36} have been explored for CO\textsubscript{2}RR, showing the importance of the structure for CO and formate production. For instance, Ag(100) facets lead to an ultralow overpotential for CO\textsubscript{2} reduction to CO on triangular Ag nanoplates (Figure 7a), while enhanced activity and selectivity for either formate or CO was found for Pd and Au NPs, respectively (Figures 7b-c). Regarding Cu, cubic-shaped NPs composed of (100) facets were found to display increased ethylene selectivity while suppressing methane production\textsuperscript{37,38}. The presence of small well-ordered (100) domains, combined with numerous steps and edges could be responsible for this selectivity trend as proposed for stepped surfaces\textsuperscript{1}. However, other facts might also determine the selectivity towards the formation of C\textsubscript{2+} products. Gao et al\textsuperscript{38} recently demonstrated the combined effect of morphology, defect density and oxygen content on the activity and selectivity of Cu nanocubes modified by plasma treatments. The presence and stabilization of subsurface oxygen and Cu\textsuperscript{+} species under reaction conditions were key parameters to favor C\textsubscript{2+} product selectivity (Figure 7d).
Figure 7. (a) Active adsorption site density on Tri-Ag-NPs. Reproduced with permission from ref. 35. Copyright 2017 American Chemical Society. (b) Free energy diagrams for CO₂RR to formic acid on Pd. Reproduced with permission from ref. 34. Copyright 2016 American Chemical Society. (c) CO FE of an Au film and shaped Au NPs at −1.0 V_RHE. Reproduced with permission from ref. 36. Copyright 2015 American Chemical Society. (d) Current density and ethylene FE at −1.0 V_RHE for different plasma-treated Cu nanocubes and an electropolished Cu foil. Reproduced with permission from ref. 38. Copyright 2017 American Chemical Society.

7. Stability and Support Effect.

Dynamic changes in the morphology of shaped electrocatalysts in the course of the CO₂RR cannot be neglected, since they might have ascribed concomitant changes in the resulting activity and selectivity. For example, surface roughening, defect and pore formation, and the disappearance of well-defined (100) facets on Cu nanocubes deposited on carbon, together with a reduction of...
CuO\textsubscript{x} species were found to be responsible of a significantly lower selectivity towards ethylene and ethanol in favor of methane.\textsuperscript{39} The effect of the support was also explored, showing an enhanced morphological durability and greater stability of Cu\textsuperscript{+} species under CO\textsubscript{2}RR when a Cu foil was used as support instead of carbon. Furthermore, the Cu cubes on Cu-foil\textsuperscript{38} displayed a lower overpotential and higher FE for C\textsubscript{2+} products than those on C (Figure 8a).\textsuperscript{39} Carbon supports with different morphologies have also been explored for Cu NPs in order to increase the CO surface concentration and thus facilitate its dimerization and ethylene production (Figure 8b).\textsuperscript{40} Strong interactions at metal/metal-oxide interfaces were found to be responsible for higher reactivity and stability. For instance, Au NPs supported on CeO\textsubscript{x} showed higher CO FE than those supported on C, which was ascribed to an enhanced CO\textsubscript{2} activation at the Au-CeO\textsubscript{x} interface (Figure 8c).\textsuperscript{41}

Figure 8. (a) EC-AFM images of Cu-cubes on HOPG before and during CO\textsubscript{2}RR at -1.1V\textsubscript{RHE}, and effect of the support (C vs. Cu) in methane and ethylene FEs. Reproduced with permission from
8. Roughness and Defects

Nanostructured materials show better catalytic activity and selectivity than flat surfaces. The presence of low coordinated sites such as steps, edges and defects\textsuperscript{42}, grain boundaries\textsuperscript{43} or certain periodic spacing between the facets at the electrode surface\textsuperscript{1}, are thought to be responsible for the enhanced reactivity. The \textit{in situ} electrochemical reduction of metal oxides offers a way to prepare roughened and highly defective structures that exhibit higher current densities compared to flat surfaces due to their larger electrochemically active surface areas.\textsuperscript{44} However, the selectivity for CO\textsubscript{2}RR of these systems is also drastically modified. Roughened and highly defective Cu surfaces display preferential selectivity towards multicarbon products and oxygenates.\textsuperscript{38,42,45–47}

On the other hand, high CO yield at lower overpotentials has been found for highly porous Au, Ag and Zn catalysts. For example, the improved catalysis in oxide-derived (OD) Au electrodes, which showed $>96$ % FE for CO at -0.35 V vs RHE over the course of 8 h electrolysis, has been ascribed to a better stabilization of the CO\textsubscript{2}$^{\text{●―}}$ intermediate (Figure 9a).\textsuperscript{48} In the same way, O\textsubscript{2}-plasma pre-treated highly defective nanostructured Ag catalysts showed CO FE above 90 % at -0.6 V vs RHE, a notable enhancement over flat Ag foils, which reduces CO\textsubscript{2} at overpotentials below -0.9 V vs RHE\textsuperscript{49}. The \textit{operando} X-ray absorption fine-structure spectroscopy (XAFS) and \textit{quasi in situ} X-ray Photoelectron Spectroscopy (XPS) characterization of these samples revealed that the surface AgO\textsubscript{x} species are reduced within the first 3 min of the reaction, and that subsurface oxygen or AgO\textsubscript{x} is not detected either in the bulk of the foil after 35 min, ruling out the role of AgO\textsubscript{x} or O in their improved reactivity, Figure 9b. Instead, DFT calculations indicated that defects created by the plasma treatment lead to locally enhanced negative electric fields which resulted in a decrease in the overpotential needed to produce CO.

Oxide-derived Cu catalysts have been proven to be very selective for the formation of C$_2$+ products. Structural factors such as roughness effects, defects and grain boundaries left behind by the oxygen pre-treatment have been initially deemed responsible for the former unusual reactivity, although more recently the presence of Cu$^+$ and residual subsurface oxygen have been suggested to influence the formation of ethylene and alcohols. Still, significant reduction of the OD Cu catalyst to metallic state is bound to occur when electroreducing CO$_2$ \cite{Kundu2021}. In fact, a number of studies have reported metallic Cu as the only stable and active species during CO$_2$RR \cite{Yang2019, Kundu2021}. However, special sites can be created in the Cu structures to stabilize the Cu$^+$ species or sub-surface oxygen. For instance, the inclusion of boron as a modifier element can change the electronic structure of Cu and stabilize Cu$^{δ+}$. The latter was theoretically and experimentally linked to a higher selectivity towards C$_2$ products from CO$_2$RR \cite{Matsui2018}.

The presence of Cu$^+$ species and sub-surface oxygen during CO$_2$RR was experimentally demonstrated via operando XAFS (Figure 10a) \cite{Matsui2018}. Although the roughness of these plasma-treated...
OD-catalysts played a role in contributing to their increased activity, their outstanding ethylene selectivity (up to 60 %) was found to correlate with the presence of Cu⁺ and subsurface oxygen. The role of CuOₓ species and oxygen in Cu had been also previously discussed by others⁵⁵, but only based on ex situ measurements,⁵⁶ which had been questioned due to the facile oxidation of Cu upon air exposure for the characterization. A subsequent theoretical study described a synergistic effect between neighboring Cu⁺ and Cu⁰ in Cu surfaces leading to improved kinetics and thermodynamics of both CO₂ activation and CO dimerization, while hindering the C₁ pathways (Figure 10b).⁵⁷ Despite these promising results, the catalytic role of Cu⁺ and subsurface oxygen is a current topic of strong controversy.⁵⁸

![Figure 10](image_url)

Figure 10. (a) Hydrocarbon selectivity of plasma-treated Cu foils and EXAFS spectra with fits acquired under operando CO₂RR conditions. Reproduced with permission from ref. 45. Copyright 2016 Nature Publishing Group. (b) Free energy profiles of CO dimerization on a fully oxidized (red), metallic (blue) and Cu metal in oxidized (green) matrixes. Reproduced with permission from refs. 57. Copyright 2017 Proceedings of the National Academy of Science. (c) HCO₂H and CO FEs for Sn foil and in situ-deposited Sn/SnOₓ thin film electrodes. Reproduced with permission from ref. 60. Copyright 2012 American Chemical Society. (d) Potential-
dependent operando Raman spectra of a SnO\textsubscript{2} surface. Reproduced with permission from ref. 61. Copyright 2015 American Chemical Society.

The effect of the metal oxidation state and the presence of sub-surface oxygen is not only important for Cu. SnO\textsubscript{x} is essential for CO\textsubscript{2}R catalysis on Sn electrodes. DFT calculations point out that oxygen vacancies on the SnO surface are decisive for the selective reduction of CO\textsubscript{2} to formate.\textsuperscript{59} Moreover, SnO\textsubscript{x} appears to provide chemical functionality that stabilizes the CO\textsubscript{2}\textsuperscript{−}\textsuperscript{−} intermediate, mediating the direct electron transfer that yields CO and HCO\textsubscript{2}H (Figure 10c).\textsuperscript{60} Furthermore, based on operando Raman spectroscopy, a decrease in the formate FE was assigned to the reduction of SnO\textsubscript{x} to metallic Sn (Figure 10d).\textsuperscript{61} These results may open new perspectives in the preparation of metal/metal oxide composites with superior activity and selectivity.

10. Electrolyte Effect

CO\textsubscript{2}RR performance is generally measured in aqueous electrolytes, typically potassium bicarbonate solutions. Alkali metal cations have been reported to facilitate CO\textsubscript{2} adsorption and stabilize reaction intermediates.\textsuperscript{62} On Cu electrodes, increasing the cation size leads to a decrease in FE\textsubscript{s} of H\textsubscript{2} and C\textsubscript{1}, as well as an increase in C\textsubscript{2+} (Figure 11a).\textsuperscript{62–64} This selectivity change could be explained by the preferential hydrolysis of the cations, lowering the local pH near the electrode, leading to higher local CO\textsubscript{2} concentration.\textsuperscript{62} Bicarbonate is considered to contribute to the enhanced CO\textsubscript{2}RR activity by increasing the effective CO\textsubscript{2} concentration near the electrode surface.\textsuperscript{65–67} In addition, the bicarbonate solution can also suppress the increase of the local pH near the electrode surface, which significantly affects the reaction pathways for the production of methane and ethylene on Cu.\textsuperscript{68} CO\textsubscript{2}RR conducted in highly concentrated (10 M) KOH (high pH) over a Cu gas diffusion electrode revealed that the onset of ethylene evolution (−0.165 V vs RHE) occurs almost simultaneously with the CO production.\textsuperscript{69} Similarly, the major product on Ag switched from CO at pH 7 to formate at pH > 15 (Figure 11b).\textsuperscript{70}

Since the CO\textsubscript{2}RR performance is very sensitive to the composition and nature of the electrolyte, a number of studies exist trying to improve C\textsubscript{2+} production via electrolyte design.\textsuperscript{62,63,71,72} For example, by adding halide ions into the bicarbonate electrolyte, the CO\textsubscript{2}RR
activity could be significantly increased without sacrificing the intrinsically high C$_2$+ selectivity of plasma-oxidized Cu catalysts (Figure 11c). Halide ions such as I$^-$ can also induce significant nanostructuring of the oxidized Cu surface, even at open circuit potential. The stabilization of Cu$^+$ species made the halide effect on oxidized Cu different from that on metallic Cu, where an increased methane production is observed in contrast to C$_2$H$_4$ production over CuO$_x$.\textsuperscript{72}

In aqueous electrolytes, the reaction rate of CO$_2$RR is usually mass transport-limited due to the low CO$_2$ solubility in water (33 mM at 25 °C, 1 atm). To overcome this disadvantage, high CO$_2$ pressure\textsuperscript{73} and supercritical CO$_2$\textsuperscript{74} have been explored to increase the CO$_2$ concentration in aqueous electrolytes. Additionally, organic solvents such as acetonitrile, methanol and N,N-dimethylformamide are also used due to their improved CO$_2$ solubility compared to water. Ionic liquids can also be employed as co-catalysts, either as electrolytes dissolved in a solvent, or as combined solvent/electrolyte. Rosen et al.\textsuperscript{75} reported an electrocatalytic system with 1-ethyl-3-methylimidazolium tetrafluoroborate (EMIM-BF$_4$) as an electrolyte in water which was able to reduce CO$_2$ to CO at overpotentials below 0.2 V with >96 % FE. The ionic liquid lowered the energy of the (CO$_2$)$^-$ intermediate and the initial reduction barrier (Figure 11d).
11. Conclusion

In summary, structure- and electrolyte-reactivity relationships in CO2RR have been exemplified over a number of molecular, nanostructured, and single crystal catalysts. Although the dynamic
changes of the morphology, atomic arrangement, coordination number, and oxidation state of the catalysts under reaction conditions can already be monitored by advanced *operando* microscopy and spectroscopic techniques, improved surface sensitivity of some of this methods (e.g XAFS) is still needed when surface-modified bulk systems are studied. Furthermore, challenges still exist when trying to conduct truly surface-sensitivity measurements (e.g. XPS) not only *quasi in situ*, but also in a flow-cell configuration under potential control.

Moreover, the greater complexity of real catalysts compared to model single crystals further motivates researchers to try to synthesize well-defined nanoscale systems where one could obtain in depth understanding of the different parameters affecting CO₂RR by isolating, when possible, different contributions, as for example the role of the NP size, shape, structure, oxidation state, composition, interparticle distance, etc. Adding to this challenging task is the need of understanding the complexity of electrified solid/liquid and solid/liquid/gas interfaces. For instance, the adsorption of anions and cations on the electrode surface is usually different in the presence of applied potentials, and the adsorbed species also interact with CO₂ and reaction intermediates differently when electric fields are considered.

In the coming years, being able to gain in depth atomistic/molecular insight into the complexity of electrode/electrolyte interfaces, including their interaction under different environmental conditions will be key for understanding CO₂RR mechanisms and further designing more active and selective catalysts.

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