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Perspective

Outer-coordination sphere in multi-H⁺/multi-e⁻ molecular electrocatalysis

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SUMMARY

Electrocatalysis is an indispensable technique for small-molecule transformations, which are essential for the sustainability of society. Electrocatalysis utilizes electricity as an energy source for chemical reactions. Hydrogen is considered the "fuel for the future," and designing electrocatalysts for hydrogen production has thus become critical. Furthermore, fuel cells are promising energy solutions that require robust electrocatalysts for key fuel cell reactions such as the interconversion of oxygen to water. Concerns regarding the rising concentration of atmospheric carbon dioxide have prompted the search for CO₂ conversion methods. One promising approach is the electrochemical conversion of CO₂ into commodity chemicals and/or liquid fuels, but such chemistry is highly energy demanding because of the thermodynamic stability of CO₂. All of the above-mentioned electrocatalytic processes rely on the selective input of multiple protons (H⁺) and electrons (e⁻) to yield the desired products. Biological enzymes evolved in nature to perform such redox catalysis and have inspired the design of catalysts at the molecular and atomic levels. While it is synthetically challenging to mimic the exact biological environment, incorporating functional outer coordination spheres into molecular catalysts has shown promise for advancing multi-H⁺ and multi-e⁻ electrocatalysis. From this Perspective, herein, catalysts with outer coordination sphere(s) are selected as the inspiration for developing new catalysts, particularly for the reductive conversion of H^+ , O_2 , and CO_2 , which are highly relevant to sustainability. The recent progress in electrocatalysis and opportunities to explore beyond the second coordination sphere are also emphasized.

INTRODUCTION

Increasing energy and environmental concerns have prompted the search for alternative energy resources. Despite the remarkable progress achieved in the past decades, the commercialization of clean and sustainable energy technologies is still limited (Lewis and Nocera, 2006). Hydrogen gas is an attractive fuel owing to its high enthalpy of combustion (–143 kJ/mol), and is a primary feedstock for fuel cells (Lewis, 2007; Turner, 2004). However, the industrial-scale production of H₂ still depends on metallic Pt or its alloys, highlighting the need for efficient and inexpensive catalysts (Winter and Brodd, 2004). Thus, exploring robust catalysts comprising earth-abundant elements has become immensely important for producing H₂ or performing the key fuel cell reactions.

The rapidly increasing atmospheric CO_2 level (at present >410 ppm) due to the combustion of carbonbased fuels is alarming and requires urgent attention(Ballantyne et al., 2012). Although CO_2 reduction chemistry began in the 1940s, steady accomplishments in this research area began in the late 1970s, followed by the confluence of climate concerns early in this century(Keith et al., 2018; Robert, 2016). The fundamental research goal is to capture atmospheric CO_2 and convert it into commodity chemicals such as methanol, formic acid, ethylene, and carbon monoxide(Boot-Handford et al., 2014; Centi and Perathoner, 2009). These transformations require selective delivery of both protons (H⁺) and electrons (e⁻) to generate specific products(Robert, 2016).

Progress in the development of electrocatalysts to promote multi-H⁺ and multi-e⁻ transfer reactions has been accomplished during the last 40 years, yet the challenges in the organization of proton and electron

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Table 1. Thermodynamic potentials of the electrochemical reactions of interest					
Reactions	Standard potential ^a (vs. NHE)	Standard potential ^b (vs. Cp ₂ Fe ^{+/0})			
$2H^+ + 2e^- \rightarrow H_2$	0.00 V	-0.28			
$\mathrm{O_2} + 4\mathrm{e^-} + 4\mathrm{H^+} \rightarrow 2\mathrm{H_2O}$	+1.23 V	+1.21			
$\mathrm{CO}_2 + 2\mathrm{e}^- + 2\mathrm{H}^+ \rightarrow \mathrm{CO} + \mathrm{H}_2\mathrm{O}$	-0.10 V	-0.12			
$CO_2 + 2e^- + 2H^+ \rightarrow HCOOH$	-0.20 V	-0.84°			
^a Reported at pH 0.(Machan, 2020; Pegis et al., 2015; Zhao et al., 2020).					

^bReported in MeCN.(Pegis et al., 2015).

^cCalculated based on the conversion factor of -0.64 V vs Cp₂Fe^{+/0}.(Pegis et al., 2015).

inventories remain the same, which is a fundamental requirement for achieving product selectivity and catalytic stability. To facilitate ideal electron and mass transport, biological systems, such as metalloproteins, employ highly optimized chemical cascades to drive multi-H⁺ and multi-e⁻ transfer processes that include secondary sphere effects. A few examples of such enzymes are: (1) hydrogenases, which catalyze reversible H₂ evolution reactions in aqueous media at or near thermodynamic potential (Madden et al., 2012; Shafaat, 2021), (2) cytochrome c oxidase (CcO), which selectively reduces O_2 with $4H^+/4e^-$ (Kaila et al., 2010; Wikström et al., 2015), and (3) carbon monoxide dehydrogenases (CODH), which reversibly catalyze the reduction of CO₂ to CO(Svetlitchnyi et al., 2001). Thus, such enzymatic mechanisms have inspired the study of redox reactions in molecular electrocatalysis where the selective coupling of H⁺ and e⁻ is the central focus(Amanullah et al., 2021). Although establishing a bioidentical environment in synthetic models is challenging, research efforts to incorporate nature's features into molecular electrocatalysts have afforded promising results(Amanullah et al., 2021; Dey et al., 2017). In this context, the concept of a second coordination sphere integrated with the catalyst's active site is the state-of-the-art approach. This Perspective emphasizes the application of the outer coordination sphere in electrocatalysis in which synthetic catalysts mimic the feature(s) at the active site of enzymes. The present discussion is limited to three examples of energy-related electrocatalysis, the H_2 evolution reaction (HER), O_2 reduction reaction (ORR), and CO_2 reduction reaction (CO₂RR), with a primary focus on how second coordination spheres direct multi-e^{-/} multi-H⁺ transfer and promote chemical transformations. Although there are many reviews on the HER(Amanullah et al., 2021; McKone et al., 2014), ORR(Machan, 2020; Pegis et al., 2018), and CO₂RR(Francke et al., 2018; Robert et al., 2020), herein, electrocatalysts with second and higher coordination spheres that mimic the function of biological motifs are highlighted, thus inspiring novel molecular designs for energy-related electrocatalysis. We also extend the discussion to include the effects of the higher coordination sphere as a promising strategy for enhancing the catalytic efficiency that includes kinetic and thermodynamic aspects. A list of the standard thermodynamic potentials for some selective multi-e⁻/multi-H⁺ reactions of interest is presented in Table 1 that will guide to estimate the overpotential for the electrocatalysts discussed later in this Perspective.

H₂ EVOLUTION REACTIONS

The prime objective of the hydrogen economy is achieving large-scale H₂ production from water as an energy source. However, the challenge for H₂ production by electrocatalysis is to achieve a high turnover frequency (TOF) at low overpotential using earth-abundant catalysts. In nature, hydrogenases in many microorganisms such as algae and bacteria couple $2H^+$ and $2e^-$ to produce H₂, or oxidize H₂ molecules with a TOF of up to 10^4 mol/s per mole of the enzyme in the aqueous medium(Thauer et al., 2010). Although immobilizing enzymes as thin films on electrodes has proven to be a viable strategy, the catalytic activity is not satisfactory (2–4 mA per 1 mM hydrogenase)(Vincent et al., 2007). Therefore, designing molecular HER catalysts that mimic the structural features found in nature remains an ongoing active research focus.

Enzymatic strategies

Three types of hydrogenases catalyze the HER: [NiFe]-hydrogenases, [FeFe]-hydrogenases, and [Fe]-hydrogenases (Fontecilla-Camps et al., 2007; Lubitz et al., 2014). Herein, we highlight the mechanisms by which the first two types, ([NiFe]-hydrogenases and [FeFe]-hydrogenases), which are most common in nature, catalyze the HER. First-row transition metals (Ni and Fe) form the active sites of [NiFe]-hydrogenases. The metals are bridged by two cysteine (Cys) thiolates (Figure 1A). Additionally, the Ni center is ligated with two other terminal Cys groups that direct H⁺ transfer while promoting the interconversion of H⁺

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A Enzymatic HER



^B Bioinspired electrocatalysts bearing second-coordination sphere (SCS)



c Examples of SCS effects in HER for synthetic HER electrocatalysts



Figure 1. Enzymatic and synthetic HER models

(A) [NiFe]-hydrogenases and [FeFe]-hydrogenases known for enzymatic HER(Shafaat, 2021). (B) Selected bio-inspired molecular HER electrocatalysts bearing second-coordination sphere (SCS). (C) Two proposed intermediates ($Ni-N_2P_2$ -Int1/Int2) for $Ni-N_2P_2$, one intermediate ($Ni-NPNH_2$ -Int) for $Ni-NPNH_2$, and two intermediates (FeFe-R-Int1/Int2; R groups are shown in (B)), showing different intermediate stabilization effects of SCS in H⁺ transfer events during HER(Helm et al., 2011; Tatematsu et al., 2016).

and H₂(Lubitz et al., 2014; Shafaat, 2021). It is proposed that four intermediate species, Ni-SI_a, Ni-L, Ni-C, and Ni-R (Figure 1A), are generated by [NiFe]-hydrogenases during the HER (identical intermediates are generated during H₂ oxidation)(Shafaat, 2021). The coordinated Cys thiolate groups at the Ni center systematically control H⁺ delivery, with consequent H₂ evolution at nearly zero overpotential. Notably, the three FeS clusters connect the active site of the [NiFe]-hydrogenases to the surface of the enzyme surface and facilitate multi-electron transfer events for the HER.

The FeS cluster in [FeFe]-hydrogenases plays similar roles in redox processes, but the proton transfer steps for proton relay are controlled by an additional secondary amine unit from the second coordination sphere (Figure 1A).(Amanullah et al., 2021) [FeFe]-hydrogenase catalyzes the HER in a more complicated manner





Table 2. Selective electrocatalysts ^a and their activity for HER						
Catalysts	TOF (s ⁻¹)	Overpotential (V)	Solvent	Proton source		
Ni-NP ₂	106, 000	0.72	MeCN	[(DMF)H] ⁺ + H ₂ O		
Ni-N ₂ P ₂	5670	~0.3 ^b	MeCN	CF ₃ CO ₂ H		
	590	$\sim 0.4^{b}$	MeCN	[(DMF)H] ⁺		
Ni-NP ₂ PhBr	3300	0.76	MeCN	$[(DMF)H]^+ + H_2O$		
Ni-NPNH ₂	8400	0.59	MeCN	CH ₃ CO ₂ H		

^aFor FeFe-x, x = 1–10, see (Amanullah et al., 2021).

^bCalculated based on the reported data. Overpotential = standard potential (corrected considering the pK_a of the acid used) – applied potential.

than [NiFe]-hydrogenase(Amanullah et al., 2021). The key intermediate for [FeFe]-hydrogenases is Fe(I)Fe(I) with a protonated bridgehead N atom (Figure 1A). The ensuing protonation step involves electron transfer from the FeS cluster to form a terminal Fe(II)Fe(II)-hydride (H) species (Figure 1A) that releases H₂ upon the addition of a proton. Thus, the [FeFe]-hydrogenases deliver H⁺ based on the presence of a bridgehead amine functionality in the second coordination sphere during the HER.

Bioinspired electrocatalysts for HER

Since the crystallographic data for [NiFe]-hydrogenase were initially reported in 1995, the synthesis of biomimetic models using synthetic inorganic and organometallic tools has been attempted(Ahmed and Dey, 2019; Simmons et al., 2014; Tard and Pickett, 2009). However, only a few synthetic models can mimic the structure-function relationship observed in the enzymatic cycle(Brazzolotto et al., 2016). Although modeling the synthesis of bimetallic complexes containing Ni and Fe or other metals (e.g., Co, Mn, Ru, etc.) remains challenging, a considerable number of bioinspired mononuclear Ni electrocatalysts bearing proton donor(s) in the second coordination sphere have been reported, and have demonstrated remarkable performance for homogeneous electrocatalysis in organic media(Bullock and Helm, 2015; Helm et al., 2011; Wiese et al., 2013). Herein, we highlight a few examples from the library of electrocatalysts for the HER that incorporate pendant Brønsted base(s) in the outer-coordination sphere. Dubois, Helm, and others reported diphosphine-chelating Ni electrocatalysts, Ni-NP₂, Ni-N₂P₂, and Ni-NP₂PhBr (Figure 1B), which utilize flexible amine functionalities in the second coordination spheres to shuttle H⁺, as well as stabilize the Ni-hydride species (Ni-N₂P₂-Int1 and Ni-N₂P₂-Int2) during the HER (Figure 1C).(Helm et al., 2011; Rountree and Dempsey, 2015; Wiese et al., 2013) Thus, the roles of these flexible arms in the outer coordination sphere resemble the function of Cys in [NiFe] hydrogenases. These molecular Ni electrocatalysts exhibit high turnover frequencies (TOF, $3,300-100,000 \text{ s}^{-1}$, Table 2) for the HER in strong acids such as protonated dimethylformamide ([(DMF)H]⁺ ($pK_a = 6.1$))(McCarthy et al., 2014), and anilinium ($pK_a = 10.6$)(McCarthy et al., 2014) in wet (>1 M H₂O) MeCN(Helm et al., 2011; Rountree and Dempsey, 2015). Furthermore, Masuda et al. reported a Ni^{II} electrocatalyst (Ni-NPNH₂, Figure 1B) with a comparatively high TOF of 8,400 s⁻¹ (Table 2) for H₂ evolution from a weak acid, acetic acid ($pK_a = 23.5$) (Tatematsu et al., 2016). This catalyst employs a phosphinopyridyl ligand with amine groups in the second coordination sphere to stabilize the Ni-hydride intermediate, Ni-NPNH2-Int (Figure 1C). Overall, these HER electrocatalysts showed stable catalytic currents over the duration of controlled potential electrolysis and the production of H_2 was confirmed using gas chromatographic analysis.

[FeFe]-hydrogenases also inspired the investigation of binuclear Fe complexes as molecular HER electrocatalysts. However, the commonly synthesized binuclear Fe complexes include six CO ligands (Figure 1B) and both Fe centers are expected to be reduced to Fe(0) before protonation. These binuclear synthetic hexacarbonyl Fe complexes (FeFe-x, x = 1-10, Figure 1B) contain an amine unit with a variety of R-substituents as a Brønsted base. The Brønsted base in the second coordination sphere is protonated in the presence of a strong acid, and thus resembles the function of the bridgehead amine in [FeFe]-hydrogenases (Amanullah et al., 2021).

The use of the pendant base in the second coordination sphere to control H⁺ transfer in the HER is not limited to bioinspired Ni-based electrocatalysts. Iron-porphyrins bearing distal triazole residues in the outer sphere (Fe-trz-Fc, Fe-trz-^tBu, Figure 2A) have also shown promise for catalyzing the HER in organic and aqueous media(Rana et al., 2017). These iron-porphyrins catalyze the HER at the Fe(I) center, and the





Figure 2. Examples of iron-porphyrins for HER

(A) Iron-porphyrins bearing triazole units in the second coordination sphere. (B) Proposed overall HER mechanism for the complexes in (A), derived from the original work reported by Dey et al(Amanullah et al., 2021). The black circle around Fe represents the porphyrin rings.

catalytic currents increase upon the addition of acid. Notably, density functional theory (DFT) calculations supported the proposed mechanism for the HER using these iron-porphyrins, where the Fe(I)-center forms Fe^{III}-hydride species upon protonation and the protonated triazoles in the second coordination sphere stabilize such intermediates through dihydrogen bond formation (Figure 2B). The red dotted bond in Figure 2B indicates stabilization of the H-bond between the protonated triazole residues and Fe^{III}-hydride species. Thus, such second coordination spheres in iron-porphyrins provide thermodynamic advantages in the HER by increasing the affinity of H⁺ for the metal-hydride intermediate(Liu et al., 2012).

ORR

Electrochemical reduction of O_2 with 4H⁺/4e⁻ for H₂O production is an important reaction in fuel cells and secondary metal- O_2 batteries(Behling, 2013). Water must be selectively produced via the ORR in the fuel cell and in metal- O_2 batteries as other partially reduced oxygen species, such as the 2H⁺/2e⁻ reduction product, H₂O₂(Borup et al., 2007), could harm the membrane separators. Therefore, the development of ORR electrocatalysts capable of selective O_2 -to-H₂O conversion near the thermodynamic potential has gained significant attention in the last five decades(Machan, 2020). Although there are numerous review articles dealing with ORR electrocatalysis(Dey et al., 2017; Pegis et al., 2018), we herein focus on bioinspired electrocatalysts possessing second-coordination spheres for achieving enhanced catalytic activity and stability, and reduced overpotential.

Biological ORR strategies

Nature has optimized metalloproteins, such as CcO, to favor the 4H⁺/4e⁻ reduction of O₂(Kaila et al., 2010; Wikström et al., 2015), in which the active sites comprise an Fe-containing heme (heme a₃), a distal Cu, and a tyrosine amino acid (tyrosine 244, Figure 3A); these species control the delivery of electrons and protons to ensure the selective reduction of O₂. The heme a₃ is an iron porphyrin that putatively generates a (Fe^{IV} = O)porphyrin^{•+} complex during catalysis(Dey et al., 2017). The Cu site and tyrosine244 supply electrons, and proton delivery is managed by a transmembrane H-bonding network that also supports the essential proton-pumping activity of CcO(Yoshikawa and Shimada, 2015). Thus, the ORR selectivity is controlled by sequential H⁺/e⁻ transfer events. Multicopper oxidases (MCOs) are another example of O₂-reducing metalloenzymes that have multiple redox centers (one type 1, one type 2, and two type 3 Cu centers) at the active center (Figure 3B).(Solomon et al., 1996) The mechanism of the ORR in MCOs preferentially follows the concerted 4e⁻/4H⁺ reduction of O₂ to H₂O(Solomon et al., 2014). The oxidation states of copper alternate between Cu¹ and Cu¹¹ to provide the required 4e⁻. Impressively, some MCOs facilitate the ORR at nearly the thermodynamic potential of the O₂/H₂O couple(Cracknell and Blanford, 2012). Overall, such structure– function relationships at the active sites of CcO and MCO provide strong motivation for the design of molecular ORR electrocatalysts.

Bioinspired ORR electrocatalysts

Designing bioinspired metalloporphyrins for selective 4H⁺ and 4e⁻ transfer and evaluating their performance are contemporary research topics. In this context, Iron tetraphenylporphyrin (FeTPP), a simple







Figure 3. Enzymatic and synthetic ORR models

Active site of (A) cytochrome c oxidases and (B) multicopper oxidases. The figures are adapted from the work reported by Dey et al. (Dey et al., 2017). (C) Symmetric Fe-porphyrins, (D) Hangman porphyrins, and (E) asymmetric metalloporphyrins bearing proton relays(Dey et al., 2017).

synthetic model of CcO, provides a remarkable electrocatalytic platform for the homogeneous ORR(Pegis et al., 2018). Modifying the **FeTPP** ring at the *meso*-positions using weak Brønsted bases or acids such as 4-pyridyl (**FeP-4Py**)(Matson et al., 2012) and 4-carboxyphenyl (**FeP-4CO₂H**, Figures 3C) and 3(Carver et al., 2012) respectively, afforded improved ORR activity compared to **FeTPP** under homogeneous conditions. The catalytic ORR activity was further improved by introducing 2-pyridyl (**FeP-2Py**)(Matson et al., 2012) or 2-carboxyphenyl (**FeP-2CO₂H**)(Carver et al., 2012) substituents at the *meso*-positions of the **FeTPP** ring. The effects of the position of pendant proton relays in the second-coordination sphere (Figure 3C) on the selectivity and kinetics of O₂ to H₂O conversion were compared (Table 3).(Dey et al., 2017) Hangman catalysts are another remarkable example of asymmetric porphyrins (Figure 3D), where one *meso*-position carries a dibenzofuran moiety that is further connected to a -CO₂H unit to enable proton relay(Chang et al., 2003). The presence of the single proton relay enhanced the ORR activity and selectivity toward H₂O formation(Rosenthal et al., 2007).

Asymmetric metalloporphyrins bearing second-coordination spheres have also been immobilized onto graphite electrode surfaces for heterogeneous ORR in aqueous electrolytes(Sinha et al., 2015, 2019). The objective is to study proton relay on solid electrode surfaces and how molecules with pendant proton relay behave on carbon surfaces during the ORR in an aqueous solution. For example, FeTPPy (Figure 3E) bearing a single pyridyl group showed stable catalytic currents over 5 h of controlled-potential electrolysis (CPE) in O_2 -saturated 1 M H₂SO₄ (pH 0), whereas the catalytic current of the parent FeTPP declined after 15 min of CPE under identical electrochemical conditions(Sinha et al., 2015). Additionally, both FeTPP and FeTPPy immobilized on edge plane graphite showed similar Faradaic efficiencies of ~90% toward H₂O production(Sinha et al., 2019). Notably, the inclusion of a single pyridyl group in FeTPPy at the *meso*-position did not affect the overpotential compared to that of FeTPP for O_2 -to-H₂O conversion at pH 0.

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Catalysts	TOF (s ⁻¹)	Overpotential (V)	Solvent	Proton source
FeTPP	40,000	0.8	DMF	Et ₃ NH ⁺
FeP-2CO ₂ H	200	0.4	MeCN	[(DMF)H] ⁺
FeP-2Py	600	$\sim 0.8^{b}$	aqueous (pH 0)	HOTf
FeTPPy	7840	$\sim 1.0^{b}$	aqueous (pH 0)	H_2SO_4
СоТРРу	15,960	~0.8 ^b	aqueous (pH 0)	H ₂ SO ₄
CoTPNMe ₃	1430	$\sim 1.0^{b}$	aqueous (pH 0)	H_2SO_4
	1183	~0.8 ^b	aqueous (pH 7)	H ₂ SO ₄

Furthermore, replacing the Fe(III) center in **FeTPPy** with Co(II) yielded another ORR electrocatalyst, **CoTPPy**, which catalyzed the ORR with selective H₂O production, without compromising the overpotential and catalytic stability. The effect of a single pyridyl group in changing the selectivity of Co-porphyrins is remarkable, as most Co-porphyrins, including **CoTPP**, promote the $2H^+/2e^-$ reduction, leading to H₂O₂ production, which is undesirable for fuel cells(Sinha et al., 2019). Furthermore, electrostatic stabilization of the Co-oxo intermediate during the ORR using the cationic ancillary group -NMe₃⁺ in the second coordination sphere of **CoTPNMe₃** also afforded pH-dependent selectivity toward H₂O formation (Figure 3E).(Zhang and Warren, 2020) The above-mentioned examples validate the high applicability of second coordination spheres in enhancing the catalyst stability and selectivity for the ORR.

CO₂RR

Converting anthropogenic CO2 into sustainable liquid fuels (methanol, methane, formic acid, etc.), or into fuel precursors such as syngas (H₂ and CO), to yield hydrocarbons has gained immense attention in recent research trends(Robert, 2016). Electrochemical conversion is a very promising approach for reducing CO₂ via multi-H⁺/multi-e⁻ steps using electrocatalysts, including metal electrodes such as Cu, Sn, Au, or Pd (Cui et al., 2020; Godbold et al., 2021; Robert et al., 2020). Although heterogeneous CO₂RR models are highly suitable for industrial-scale CO₂ reduction, molecular CO₂RR electrocatalysts are desirable for elucidating the fundamental CO₂RR mechanism. In electrochemical CO₂RR catalysis, the electron-rich electrode surface supplies multiple electrons for the reaction, but multi-H⁺ transfer often compromises the product selectivity. Typically, the metal center of inorganic catalysts first gets reduced upon e⁻ transfer and follow the protonation-first pathway that forms a metal-hydride which reacts with another H⁺ to release H₂, or with CO₂ to yield formate. Alternatively, the reduced metal center can undergo CO₂-activation-first pathway and selectively produces CO upon sequential H⁺ transfer events(Barlow and Yang, 2019). The Sabatier principle states that such catalyst-substrate interaction is significant in catalysis and the related freeenergy should be favorable(Ooka et al., 2021). Furthermore, DuBois, Yang, Berben, and many others emphasized the hydricity of the metal hydride species that determine the free energy for the reaction between a metal hydride and H^+ or CO₂ and thus, controls the product selectivity (Barlow and Yang, 2019; Loewen and Berben, 2019; Rakowski Dubois and Dubois, 2009). However, similar thermodynamic landscape in CO₂RR also can be established by incorporating proton donors into the outer-coordination sphere of molecular electrocatalysts for supplying H⁺ during the CO₂RR(Costentin and Savéant, 2017). Additionally, chemical modification beyond the active site of molecular CO₂RR electrocatalysts offers advantages, such as (1) higher activity by stabilizing the CO_2 bound intermediate, (2) higher activity provided by proton-rich environments (i.e., proton relays and hydration spheres), and (3) optimization of the selectivity for the CO_2 reduction reaction.

Biological CO₂RR strategies

Enzymes such as [NiFe]CO dehydrogenases ([NiFe]CODHs) are natural substances that catalyze the CO₂ to CO interconversion reaction(Mondal et al., 2015). The mechanism of CO₂RR catalysis by these enzymes is still under investigation or complicated(Jeoung and Dobbek, 2007). It is generally proposed that CO₂ binds at the Ni center, and the Fe center coordinates with an O atom of CO₂. Interestingly, the same O atom forms an H-bond with a nearby lysine (Lys563, Figure 4A), and the other O atom of CO₂ forms an H-bond with a proximal protonated histidine (His 93, Figure 4A). Upon binding with the Ni center, the angle







Figure 4. Enzymatic and synthetic CO₂RR models

(A) Active site of [NiFe]CO dehydrogenases(Amanullah et al., 2021), (B) Fe-porphyrin bearing eight pendant proton relays (FeTPPOH₈)(Costentin et al., 2012), (C) Co-bipyridine with a bridging pendant amine group (CoBpy-amine)(Chapovetsky et al., 2018), (D) Zn-porphyrin containing 8 triazole units each with PEG proton relays(Williams et al., 2020), (E) Fe-porphyrin with a single PEG proton relay(Chaturvedi et al., 2021), (F) Mn-bipyridine with imidazole in the second coordination sphere(Sung et al., 2019).

of the C-O-C bond in CO₂ is proposed to be 132.5°, and the C–O bond is elongated to 1.26 Å due to charge transfer to the CO₂ antibonding orbitals via back-bonding(Amanullah et al., 2021). Overall, this is viewed as the enzymatic activation of CO₂ to CO₂•, where the proximal amino acids (Lys563 and His93) stabilize the intermediate through H-bonding. Thus, such biological CO₂RR catalysts motivate the design of artificial catalysts incorporating pendant outer-sphere residues. In this Perspective, we selectively chose some bioinspired electrocatalysts (Figure 4) that use the outer coordination spheres for the selective CO₂-to-CO conversion. If there are any such electrocatalysts that produce major products other than CO, we noted in our discussion.

Intermediate stabilization in molecular electrocatalysts

Stabilization of the bound substrate is critical for enhancing CO₂ reduction. Hydrogen bonding and electrostatic interactions effectively stabilize the [catalyst-CO₂•] adduct upon CO₂ binding at the reduced metal center, thereby minimizing the energy barrier ($\Delta G^{\#}$) or overpotential for CO₂RR electrocatalysis(-Costentin and Savéant, 2017). The FeTPPOH₈ electrocatalyst (Figure 4B) developed by Savéant et al. possesses a local proton source within the catalyst, serving as a remarkable example of enhanced catalytic activity due to sphere-assisted catalysis by functional groups in the second coordination sphere(Costentin et al., 2012). Such pendant phenolic groups in close proximity to the active site facilitate rapid protonation of the substrate, allowing for a lower overpotential and higher activity compared to iron porphyrins that do not have acidic groups in the second coordination sphere. In a study of cobalt complexes (Figure 4C), Marinescu et al. found that the pendant amine unit stabilizes the cobalt/carboxylate intermediate, and the number of pendant amines is directly proportional to the rate of catalytic CO₂ reduction(Chapovetsky et al., 2018). As a critical aspect, the study demonstrated a non-cooperative effect between the amine units in the higher coordination sphere, suggesting that each amine is individually involved in catalysis. DFT calculations reveal that the amine groups do not directly interact, but rather associate, with acid molecules to provide protons for stabilizing the intermediate(Chapovetsky et al., 2018).

Proton relays

Because the CO₂ reduction pathway favors proton-coupled electron transfer (PCET) steps, proton relays continuously furnish protons to the active site to enhance the catalytic activity. To mimic the complex enzymatic system found in nature, simpler proton relays have been designed for CO₂ reduction. Several hydrophilic groups have been utilized to achieve this shuttling effect, including phenol(Margarit et al., 2018; Nichols et al., 2020; Sinha and Warren, 2018), triazole(Lashgari et al., 2020; Sen et al., 2019), polyethylene glycol (PEG)(Chaturvedi et al., 2021; Lashgari et al., 2020), imidazole(Sung et al., 2019), ure-a(Gotico et al., 2020; Haviv et al., 2018), and amine(Jakobsen et al., 2021). The relay can be as short as a single OH group in the second coordination sphere, as achieved using an asymmetric Fe-porphyrin

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Figure 5. Selectivity in CO₂RR

(A) Schematic depicting selectivity differences of a Mn catalyst with different groups in the second coordination sphere(Rønne et al., 2020). (B) Two competing catalytic pathways for the catalytic reduction of CO_2 to CO or water to H_2 by non-heme iron complexes(Zee et al., 2020). Adapted with permission from (Zee et al., 2020). Copyright 2020 American Chemistry Society.

carrying a single *ortho*-phenolic group at the *meso*-position and three phenyl groups at the remaining *meso*-positions of the Fe-porphyrin ring(Sinha and Warren, 2018). Such asymmetric iron complex remained stable for approximately 2 h at -2.3 V vs. Fc^{+/0} after an initial drop in current. Our group observed that a triazole bundle attached to a zinc porphyrin complex (**ZnPEG3T8**, Figure 4D) afforded improved catalytic activity by enabling enhanced proton relays(Lashgari et al., 2020). Another example is an iron porphyrin with a single PEG chain in the second coordination sphere (Figure 4E).(Chaturvedi et al., 2021) It was demonstrated that proton relays do not have to completely create a "hydration sphere", as observed in many previous successful systems(Gotico et al., 2019; Roy et al., 2017). Imidazo-lium groups can also act as a proton shuttle. Nippe et al. designed and studied a manganese bipyridine complex (Figure 4F), where imidazolium groups created a hydration shell around the active site(Roy et al., 2017; Sung et al., 2019). The increased presence of protons caused by hydrogen bonding networks surrounding the active site can improve catalysis by furnishing the protons required for CO₂ reduction.

Selectivity enhancements

Achieving product selectivity in the CO₂ reduction reaction is a continuing challenge because of the large number of possible products (CO, formate, methanol, etc.), for which the thermodynamic potentials are very similar(Kuhl et al., 2012). Accomplishing the desired selectivity is one of the major obstacles hindering the progress of CO₂ electroreduction. The selectivity for CO₂ reduction is governed by the formation of different catalytic intermediates, which can be selectivity altered by the secondary coordination sphere's effect on the pK_a of the catalytic active site. A manganese bipyridine complex with different pendant groups is one example in which the selectivity is enhanced by modifying the second coordination ligand structure (Figure 5).(Rønne et al., 2020) When amine units were incorporated into the second coordination sphere, a Mn-hydride species was formed during catalysis, which is critical for formate production (Figure 5A). Alternatively, when a hydroxyl or methyl group was attached to the second coordination sphere, CO₂ was directly bound to the Mn center, and CO was the major product. The acidity of the environment surrounding the metal active site was modified by the ligand's second coordination sphere and therefore was able to be tuned toward a specific electrocatalytic product.

In addition to product selectivity within the realm of possible CO_2RR products, hydrogen evolution also competes with CO_2 reduction because of the favored thermodynamics of the former. Overcoming the thermodynamic preference of hydrogen evolution is of great interest in designing CO_2RR electrocatalysts. In another selectivity study by Long et al., the ligand choice in the second coordination sphere altered the catalytic pathway completely(Zee et al., 2020). With a strong protic unit such as OH, the iron non-heme catalyst favors the formation of the iron-hydride intermediate and therefore promotes water reduction. In comparison, when a weaker acid (–NHEt) is present in the second coordination sphere, the same iron active site preferentially forms the metallocarboxylate intermediate, leading to the reduction of CO_2 to CO (Figure 5B). Both cases suggest that the second coordination sphere can influence the preferred intermediate species, ultimately impacting the product selectivity. Finally, we noted the performance of the CO_2RR electrocatalysts discussed in this Perspective and





Table 4. Selective electrocatalysts and their activity for selective CO ₂ RR						
Catalysts	$TOF(s^{-1})$	Overpotential (V)	Solvent	Proton source		
FeTPPOH8	208,000ª	0.465	DMF	H ₂ O		
CoBpy-amine	16,900	-	DMF	TFE		
FePEGP	140,000	-	MeCN	H ₂ O		
Mn-Bpy-imidizole	_	-	MeCN	H ₂ O		
ZnPEGT8	1,843	0.63ª	DMF	H ₂ O		
MnBpy	5,500	0.60	MeCN	TFE		
Fe-pyd	2,067	-	MeCN	H ₂ O		
^a Estimated based on the reported data as described in Table 2.						

tabulated the kinetic and thermodynamic parameters that are associated with their CO_2RR reactivity in organic solvents (Table 4).

BEYOND SECOND-COORDINATION SPHERE

Coordination sphere effects can extend beyond those of the second coordination sphere. Higher coordination spheres enhance the catalytic efficiency by altering the surrounding chemical environment of the second coordination sphere. Shaw et. studied a nickel diphosphine HER catalyst with an amine-pendant phosphine ligand in the second coordination sphere and amino acid in the higher coordination sphere (Figure 6A).(Dutta et al., 2013) The higher coordination sphere acts as a proton shuttle for the proton relay to the second coordination sphere, providing an enzymatic effect. Our group studied the effects of PEG units in the higher coordination sphere, in conjunction with eight proton-donating triazole groups in the second coordination sphere (ZnPEG3T8 in Figure 6B).(Williams et al., 2020) The catalytic activity of ZnPEG3T8 was 30-fold higher than that of the control compound with alkyl chains in the higher coordination sphere (ZnC8T8). Another control experiment using a PEGylated, triazole-free complex (ZnPEG3T0)



Figure 6. Higher coordination spheres

(A) Schematic depicting the second and outer coordination spheres of a nickel diphosphine electrocatalyst for reversible hydrogen evolution(Dutta et al., 2013). (B) Schematic depicting the coordination sphere environments of ZnPEG3T8, ZnC8T8, and ZnPEG3T0 with CO Faradaic efficiencies (FE%) and current densities (J) at various potentials for all compounds(Williams et al., 2020).

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also showed inferior lysis efficiency, suggesting the cooperative effects of the second and higher coordination spheres. These examples demonstrate the long-distance effects of higher coordination spheres on catalytic performance, expanding the versatility of molecular designs.

OUTLOOK AND FUTURE DIRECTIONS

Designing ligand frameworks with outer-sphere coordination is an attractive strategy for developing bioinspired molecular electrocatalysts capable of multi-H⁺ and multi-e⁻ transfer. Such strategies also help establish the Sabatier principle(Ooka et al., 2021) by stabilizing the catalyst–substrate intermediates through substituent effects for homogeneous electrocatalysis. These effects include H-bonding, the acidity of the catalytic site, and through-space electrostatic effects. Stabilization of these intermediates lowers the energy barrier for product formation, thereby lowering the overpotential for electrocatalysis. However, there is generally a trade-off between lower overpotentials and the TOF. Therefore, incorporating multiple outer sphere effects in molecular electrocatalysts is promising for boosting kinetic factors with minimal compromise of the overpotential.

Although a library of impressive homogeneous electrocatalysts bearing outer coordination sphere(s) for multi-H⁺ and multi-e⁻ transfer catalysis has been documented, practical energy devices to address global energy challenges demand heterogeneous electrocatalysts. Therefore, the immobilization of molecular electrocatalysts on solid electrodes is an emerging trend in heterogeneous electrocatalysis(Brunner et al., 2020; Geer et al., 2021; Sinha et al., 2020; Zhang and Warren, 2020). The major challenge in developing these systems is optimizing the synthetic strategies for immobilizing molecules on the electrode surface. Conjugation of molecules on the electrode surface through covalent linkages(Jackson et al., 2019; Kaminsky et al., 2019) or non-covalent adsorption(Brunner et al., 2020; Maurin and Robert, 2016; Sinha et al., 2019) are ongoing strategies for developing heterogeneous electrocatalysts, where selectively controlling the number of H⁺ and e⁻ transfers is of central interest. Nevertheless, the molecular features, including the outer-sphere effects, are often complicated by the electrode properties upon immobilization of the molecules. This is primarily because the catalytic sites reside within the electric double-layer (EDL)(-Jackson et al., 2018). We propose that higher coordination spheres can increase the distance between the catalytic site and the electrode surface to facilitate electron tunneling across the EDL.

Despite significant current challenges in the practical development of electrocatalytic devices for multi-H⁺ and multi-e⁻ transfer catalysis on the industrial scale, the outer coordination sphere concept can facilitate the design and understanding of such catalysts to afford benchmark molecular electrocatalysts. Furthermore, the selectivity for multi-H⁺/multi-e⁻ redox reactions can also be enhanced through higher coordination sphere effects due to steric effects and changes in the pK_a. This in turn changes the major reduction product by limiting the reactivity of the intermediate species (i.e., metal hydrides). Thus, a holistic approach for catalyst design that extends beyond the second coordination sphere, and focuses on higher coordination sphere effects will push the field of electrocatalysis closer toward understanding enzymatic catalysis. Fine-tuning large ligand scaffolds for electrocatalysts is expected to further drive this progress.

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DECLARATION OF INTERESTS

The authors declare no competing interests.

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