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Advances in the synthesis of Fe-based bimetallic electrocatalysts for CO₂ reduction

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Achieving carbon neutrality and slowing down global warming requires research into the electrochemical CO_2 reduction reaction (CO_2RR), which produces useful compounds. Utilizing renewable energy to meet carbon-neutral energy goals produces single-carbon (C_1) and multi-carbon (C_{2+}) goods. Efficient and selective electrocatalysts are essential to advancing this revolutionary technology; bimetallic Fe-based catalysts work better than their monometallic counterparts because multiple metals work synergistically to reduce CO_2 levels. A thorough summary of recent developments in the synthesis of Fe-X bimetallic catalysts will be provided in this review, with an emphasis on key performance indicators like stability, faradaic efficiency, potential, current density, and primary product production. In addition, this analysis will look at representative instances of Fe bimetallic catalysts that are well-known for their selectivity in generating particular alcohols and hydrocarbons, clarifying the mechanics behind CO_2 reduction, pointing out existing difficulties, and examining the potential of electrosynthesis processes in the future.

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1. Introduction

Over the past 170 years, human activity has caused a sharp rise in CO₂ emissions, which has resulted in ocean acidification and global climate change. Terrestrial ecosystems only absorb around 30% of CO₂ produced by humans annually, which is insufficient to offset anthropogenic emissions, and the amount



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of CO_2 in the atmosphere has dramatically increased in recent decades, reaching 400 ppm for the first time in human history. Since the late 1950s, the Mauna Loa Observatory in Hawaii has been continually monitoring atmospheric CO_2 , and as of 2024,



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CO₂ concentrations are approximately 417 ppm. One of the main causes of climate change, this represents a sharp rise from pre-industrial levels of about 280 ppm.³⁻⁵ According to predictions, if present emission trends continue, CO₂ levels might surpass 450 ppm by 2030. If immediate mitigation measures are not taken, some models forecast significantly higher concentrations.⁶ Over 450 ppm could dramatically raise the probability of catastrophic climate consequences, including more severe and frequent heatwaves, rising sea levels, and disturbances to

ecosystems and food security.7 Although CO₂ is a very stable molecule that is usually inert, it can undergo electrochemical activation and be transformed into reduced products through the CO2 reduction reaction (CO₂RR) with the help of protons in solution and appropriate cathodic reduction potentials.8,9 Several methods have been suggested to change CO2 into value-added products, such as chemical transformation, 10,11 reduction by photocatalysis, 12-14 reduction by electrocatalysis, 15,16 thermal catalysis, 17,18 photothermal catalysis,19 as well as biological conversion.20,21 The simplicity, mild reaction conditions, environmental compatibility, and possible integration with energy from renewable sources of electrocatalytic CO2 reduction (eCO2RR) make it stand out among these applications. Since CO2 has a linear, symmetrical structure with a zero-dipole moment, it is stable and challenging to activate in electrocatalytic reduction. Because of the molecule's symmetry, the opposing dipoles of the C=O bonds cancel each other out, resulting in a low electron density around the molecule's center, especially close to the carbon atom. The molecule's general lack of a dipole moment is a result of the decreased electron density surrounding the carbon center. It is more difficult to activate CO2 due to its low electron density. Effective electrocatalytic reduction requires the catalyst to capture and hold CO₂ molecules long enough for the reaction. However, CO2 lacks a strong dipole and other reactive properties, making it more difficult to activate for the reduction process. This demands extremely effective catalysts that can change CO2's electron density or aid in breaking its strong C=O bonds.22,23 A significant overpotential is needed for CO2 activation due to the dissociation energy needed to break the C=O bond, which is more than 750 kJ mol⁻¹, leading to poor energy efficiency and high operational costs.24 The shale gas revolution of the last twenty years, especially the large amounts of ethane (3-12% fraction) from shale gas, has changed the dynamics of the global energy market and produced an excess of ethane despite its relatively low market price, which is especially noticeable in the United States.25,26 Studies on CO2 reduction for CO and carbon-based energy sources²⁷⁻²⁹ have gained increasing attention owing to rising atmospheric CO2 levels and expanding energy demands, with a focus on finding inexpensive, efficient, and selective catalysts.30-32 These catalysts include homo-bimetallic sites (Fe-Fe, Co-Co, Ni-Ni, Cu-Cu) that indicate enhanced reactivity in comparison to monometallic counterparts, while heterobimetallic catalysts remain relatively overlooked.33-41

Efficient CO_2 reduction catalyst design is hard due to stability, huge potentials, ^{29,42} restricted solubility, competing HER, and slow kinetics. ^{43–46} Reduced activation barriers are

critical for improving electrocatalyst efficiency and selectivity. Despite the beneficial features of iron-group metallic alloys and compounds, their efficacy as catalysts in CO2 reduction remains insufficient due to low activity and stability. 47,48 Because Fe is so readily available, it is essential to build highly efficient Fe-N-C catalysts. With the abundance of iron, there is a need to produce these catalysts. Fe-porphyrins treated with phenolic groups showed remarkable CO faradaic yields exceeding 90% without degradation, emphasizing Fe-N4 sites in macrocycles as active centers.49 The incorporation of Fe atomically into nitrogendoped carbon substrates, such as Fe-N-C catalysts, has demonstrated remarkable catalytic reactivity towards CO2 reduction to CO, with Fe-N₄ sites largely studied as active sites in several investigations. 50-53 Fe species provide dynamic surface manipulation, which is critical for understanding structure dynamics and rational catalyst designing in CO2 electroreduction reaction (CO2ERR).54-56 Recent studies reveal that heteroatom inclusion in carbon support alters the electrical environment, allowing tailored Fe sites to lower the energy of activation limitations in electrocatalysis.57 Fe or Cu-based metals/alloys are widely used as catalysts in CO2 reduction; Fe has significant catalytic activity and a minimal energy barrier, whereas Cu has excellent CO2 adsorption traits and resistance to coking.58-61 In addition to heteroatom inclusion, the use of a second metal atom in Fe-based materials also termed dualatomic catalysts (DACs) improves catalytic activity synergistically. CO2 reduction relies on DACs, while Ru, Fe, Mn-based homogeneous, and Cu-based heterogeneous catalysis provide viable alternatives. 62-72 The adsorbate-metal surface interaction in Fe-N-C single-atom catalysts (also known as SACs) is influenced by the shift of the d-band center.73 Determining the intensity and kind of these interactions between molecules is largely dependent on this change. Consequently, it has a major effect on the catalytic activity. Integrating heteronuclear metal atoms such as Ni, Co, and Zn permits electronic structure adjustment, which facilitates adsorbate absorption as well as desorption on the surface of the catalyst.74-76 Recently, multiple reviews have investigated the CO2ERR, spanning diverse catalysts like as copper-palladium nanoalloys,9 Cu-based nanocrystals,77 bimetallic chalcogenides,78 bimetallic catalysts with atomic sites,79 Bi-based,80 Ni-based,81 Sn-based,82 and carbidebased bimetallic catalysts.83 Fe-based bimetallic electrocatalysts are superior to other metals because they are more affordable, widely available, and have better selectivity for CO2 reduction products. When it comes to stability, efficiency, and scalability, they can perform better than single-metal catalysts like Cu. There is currently no review that provided in-depth analysis of the research on Fe-based bimetallic electrocatalysts for CO2 reduction. Given the growing importance of electrolytic CO₂ reduction, the performance of Fe-based bimetallic catalysts merits a thorough examination. This research focuses on the production, implementation, and mechanistic understanding of these catalysts in CO₂ electrocatalysis, covering a wide spectrum of product forms. Furthermore, the study highlights the problems and opportunities in developing and comprehending Fe-based bimetallic electrocatalysts, which offer useful insights for future research paths in this sector.

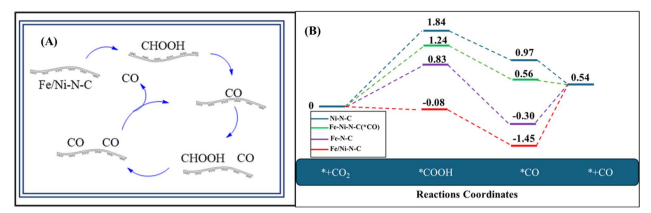


Fig. 1 (A) Proposed paths for reduction of CO_2 into CO over Fe/Ni-N-C. (B) Using DFT, the electrochemical reduction of CO_2 to CO on Fe-N-C, Ni-N-C, and Fe/Ni-N-C with and without adsorbed *CO was represented by a free energy diagram.

2. CO₂ reduction pathways over Febased bimetallic electrocatalysts

Fe/Ni-N-C materials were used as electrocatalytic reduction (ECR) catalysts by Huiying Tian and coworkers. These substances were utilized to speed up the electrochemical processes that produce CO.84 Fig. 1A illustrates the CO2 reduction into CO over the Fe/Ni-N-C catalyst. For the catalytic reduction of CO₂ into CO, the Fe/Ni-N-C catalyst provides sites for CO₂ to bind, and Fe/Ni acts as active centers for the reduction. For binding, CO2 accepts electrons and protons from the electrolyte solution and converts them into intermediate CO precursors such as CHOOH. This reaction is typically completed in two steps CO₂ → COOH → CO* as mentioned in Fig. 1. The Fe/Ni-N-C electrocatalyst achieved an impressive (faradaic efficiency of CO) FE_{CO} of 92.9% at -0.677 V vs. (reversible hydrogen electrode) RHE, indicating great efficiency. The system retained an elevated current density and faradaic efficiency for the generation of CO (FE_{CO}) when applied in a continuous flow cell at scale, holding onto over 89% shortly after 40 hours of electrolysis. Because of the binary metals combined effect, charge transfer rates were increased, resulting in favorable kinetics and long-term, effective electrochemical performance.⁵⁷

To thoroughly investigate the combined effect of Ni-N-C, Fe-N-C, and Fe/Ni-N-C on ECR, DFT studies were performed utilizing the computational hydrogen electrode technique.85 In electrocatalytic CO2 reduction, the Fe atom's electrical properties and catalytic action are largely determined by its spin orientation.86,87 Fe can exist in both high and low spin states in bimetallic Fe-based catalysts, which affects the electron distribution in the d-orbitals and changes the CO2 and intermediate adsorption strength.88 While a low-spin level may produce more stable, less reactive configurations, a high-spin state can increase the activation of CO2 by offering more accessible electron states.89 The impact of these spin arrangements on the CO₂ reduction reaction mechanism and efficiency is clarified using DFT simulations. The major catalytic sites, Me-N₄ motifs, were used as single-site models based on prior investigations. 51,90 The electrocatalytic reduction (ECR) process involved typical two-electron and two-proton transfer reactions,

culminating in the creation of *COOH and *CO intermediates. The symbol asterisk (*) represents the active site. Ni-N-C, Fe-N-C, and Fe/Ni-N-C optimized geometries served as computational models for the investigation.91,92 On Fe-N-C sites, the rate-determining step was $*CO \rightarrow CO_{(g)}$, while on Ni-N-C sites, it was $CO_{2(g)} \rightarrow *COOH$. Fe/Ni-N-C generated *COOH intermediates easily, while *CO desorption was difficult. Fe/Ni-N-C adsorbed with *CO intermediates had a much smaller free energy shift for the rate-determining phase $*CO \rightarrow CO_{(g)}$, indicating easier desorption. This shows that Fe/Ni-N-C provides more active sites by efficiently combining the benefits of Ni-N-C and Fe-N-C sites, increasing CO generation catalytic activity. CO2 adsorption on Fe-Ni bimetallic sites, electron and proton transfer pathways to form *COOH within *CO intermediate, and subsequent CO(g) desorption to regenerate Fe-Ni-N-C (*CO) are the suggested ECR reaction routes on Fe/Ni-N-C. This highlights the increased catalytic activity seen in the studies. The Fe, Ni bimetallic nitrogen-doped carbon successfully lowered the energy barriers of *COOH intermediate production and *CO-to-CO, improving favorable kinetics and increasing ECR activity, verified by the DFT calculations. According to the findings of the calculations above, the suggested ECR reaction pathways of CO2 to CO on Fe/Ni-N-C and their energy diagram using DFT calculation are shown in Fig. 1.84

3. Bimetallic graphene catalysts: mechanistic pathways for CO₂ reduction and CH₄ production

Previous studies have shown that single-atom-doped graphene is exceptionally efficient for catalyzing CO₂RR.^{93–96} Researchers have studied diverse doping techniques for adding transition metals to the graphene, demonstrating that bimetal single-atom-doped catalysts have greater catalytic performance than standard single-atom-doped catalysts.^{97–100} Run Zhang *et al.*, employed DFT calculations for examining CO₂RR on three bimetal-doped graphene catalysts, Cu–Ni/DG, Cu–Fe/DG, and Fe–Ni/DG. Different reduction pathways yield various products

Table 1 Summary of adsorption energy of different reaction intermediates for the production of CH_4 on Cu-Fe/DG and Fe-Ni/DG catalysts

	Potential energy	(eV)
Reaction intermediates	Cu-Fe/DG	Fe-Ni/DG
*CO	-1.7	-2.81
*HCOOH	0.78	-0.27
*CH ₂ O	-1.46	-2.05
*CH ₃ OH	0.97	-0.44
*CH ₄	0.70	-0.76

such as CH_4 , CH_3OH , HCOOH, and CO. In the initial stages of CO_2RR on doped graphene, CO_2 adsorption occurs, characterized by analysis of E_{ads} , electron density difference, density of state (DOS), as well as Integrated Crystal Orbital Hamiltonian Population (ICOHP). Compared to Cu-Fe/DG, CO_2 reacts more strongly with iron-based Fe-Ni/DG. The catalytic performance of the material is improved by this greater contact. Electron density difference, DOS, and ICOHP studies reveal more robust interactions between certain dopants (Fe and Ni) and CO_2 . Salve Table 1 summarizes the adsorption energies of process intermediates on bimetal-doped Fe catalysts. When CO_2 is first protonated, it produces *COOH or *OCHO, which can then be hydrogenated again to generate CO or HCOOH. These changes proceed in many ways:

$$COOH \rightarrow *CO + H_2O \rightarrow * + CO$$

and

On Cu–Fe/DG, Cu–Ni/DG, and Fe–Ni/DG catalysts, the high adsorption of CO and HCOOH encourages continued reduction as intermediates. Nevertheless, significant free energy barriers prevent CO and HCOOH from being desorbed from the catalyst's surface, which presents problems for product release. In the CO $_2$ reduction reaction, CH $_3$ OH is a potential product. Four pathways for CO $_2$ \rightarrow CH $_3$ OH involve *CO or *HCOOH as intermediates. *CO undergoes the following reactions:

*CO
$$\rightarrow$$
 *CHO \rightarrow *CH₂O (exothermic)

or

*CHO is exothermically converted to *CH₂O on Cu-Ni/DG and Cu-Fe/DG. On the other hand, *CHO into *CHOH conversions on Fe-Ni/DG are endothermic.

This distinction draws attention to the different energetics of different catalysts for these reactions.

The thermodynamic favorability of *CH $_2$ O formation is highlighted by its reduced variance in free energy, which is why it is preferred over *CHOH formation. This preference highlights the role that energetics play in identifying the paths of reactions. Six routes for CO $_2 \rightarrow$ CH $_4$ reduction were investigated, using *CO or *HCOOH as intermediaries. *CO undergoes the following conversions:

or

It resulted in *CHO or *COH and finally CH₄ by additional hydrogenation. *CO prefers *CHO production because of the lower free energy fluctuation. On Cu-Ni/DG and Cu-Fe/DG,

While, Fe-Ni/DG prefers,

Path 3 is the best route from Path 1 to Path 5, demonstrating its effectiveness and fit for the intended change. Nevertheless, route 6 (*OCHO \rightarrow *HCOOH) doesn't work as the best route for Fe–Ni/DG due to significant free energy fluctuation. The optimized pathway is Path 6, which is exothermic on Cu–Fe/DG and Cu–Ni/DG.¹⁰¹

*CHO
$$\rightarrow$$
 *CH₂OH \rightarrow *OCH₃ \rightarrow *CH₄ (exothermic)

Because CO₂ interacts with Fe or Ni atoms more strongly than its interaction with Cu, Fe–Ni/DG is more stable than Cu–Ni/DG and Cu–Fe/DG. The catalytic potential of Fe–Ni/DG for CO₂ conversion reactions is highlighted by its improved stability. Fe–Ni/DG becomes more prominent in bimetal-doped graphene systems because of its increased stability, which implies that it can support effective and long-lasting catalytic activity. Graphene doped with Cu, Fe, and Ni shows significant selectivity for CO₂ reduction over hydrogen evolution (HER), suggesting that these materials are effective catalysts for CO₂ conversion processes, with various product outcomes seen for the initial protonation step of CO₂ on these catalysts.

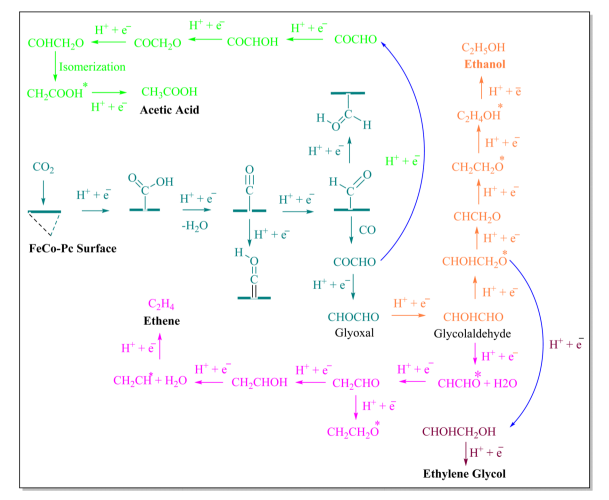
4. FeCo-Pc catalysts for multi-carbon (C₂) product formation

FeCo-Pc catalyst with dual metal-nitrogen active sites for efficient CO₂RR. FeCo-Pc overcomes the challenge of C-C coupling seen in single-atom catalysts, ^{102,103} enabling the production of

Table 2 Summary of reaction intermediates generates during the formation of C_{2+} products with their possible potential energy over FeCo-Pc catalyst

Compound	Reaction intermediate	Potential energy (eV)	
C_2H_4	COOH*	0.13	
2 1	CO*	0.23	
	CHO*	0.68	
	COCHO*	0.70	
	CHOCHO*	0.38	
	СНОНСНО*	0.55	
	CHCHO*	0.27	
	CH ₂ CHO*	0.17	
	CH ₂ CHOH*	0.06	
	CH ₂ CH*	0.51	
C_2H_5OH	CHOHCH ₂ O*	0.41	
	CHCH ₂ O*	0.21	
	CH ₂ CH ₂ O*	0.40	
	$C_2H_2OH^*$	0.71	
CH ₃ COOH	COCHOH*	0.57	
3	COCH ₂ O*	1.59	
	COHCH ₂ O*	0.26	
	CH ₂ COOH*	0.60	
CH ₂ OHCH ₂ OH	CHOHCH ₂ OH*	0.16	
	CH ₂ OHCH ₂ OH*	0.20	

C2 products, as shown in Table 2. These C2 products include C₂H₄, CH₂OHCH₂OH, C₂H₅OH, and CH₃COOH with enhanced selectivity, due to the cumulative effects of Fe and Co dual active sites anchored within phthalocyanine (FeCo-Pc).104 The computations are performed using the Gaussian 09 program, PBE exchange-correlation functional, and 6-31G* basis sets. 105,106 C-C coupling processes in CO2 reduction are critical for comprehending multi-carbon product generation. 107,108 Water plays an important role in the reduction of CO₂ because it influences intermediate hydration, provides protons for product production, and functions as a solvent for ion transport. Additionally, through the oxygen evolution reaction (OER), it competes with CO2 reduction and affects catalyst behavior. 109,110 CO was identified as a crucial intermediate, discovered by in situ spectroscopy. 111,112 PCETs generate C₁ intermediates like CHO* and COH*, with thermodynamics and kinetics assessed on FeCo-Pc surfaces. CO to CHO* (formyl group) is produced by further reducing *CO and adding an extra proton and electron. Usually, this process produces more valuable chemicals such as alcohols and aldehydes. In the case of COH* protonation step is required for the formation of COH*, but not the complete reduction required to generate *CHO. In the synthesis of other C₁ products, such as ethanol or



Scheme 1 Possible electrochemical reaction pathways of CO_2 over Fe-based bimetallic catalyst into C_{2+} products.

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methane, this is frequently a transitional stage. CO dimerization, proposed as the initial step to C₂₊ products, and carbene (CH₂) coupling with CO* to form COCH₂ or CH₂CO*, are investigated.42,113 However, the generation of "dead-end" intermediates, such as COCO* and other C-C linked intermediates, tightly bound to FeCo-Pc, incurs high energy consumption, creating kinetic obstacles and surface contamination. Electrolytes C-C coupling pathways are influenced by pH,114 with high pH favoring CO* to COH* conversion and CO* coupling with CHO* to generate COCHO*, boosting C2 synthesis. 115,116 COCHO* formation has a lower activation barrier compared to CHO*-CO* precursors, demonstrating its thermodynamic favorability and potential to provide more C₂ products in CO₂RR on FeCo-Pc surfaces.117 In an aqueous solution, the relative stability of COOH* as well as OCHO* in CO2RR against H* in HER affects the competing processes of the H2 evolution reaction (HER).118 FeCo-Pc prefers CO2RR over HER due to larger free energy changes for COOH* and OCHO* production against adsorbed H*. COOH* has a smaller overpotential than OCHO*, indicating FeCo-Pc prefers CO₂RR. 119,120 The electro-conversion of CO2 to C2H4 is important for the C2H4 industry, however, it faces challenges with an elevated overpotential and multielectron transfer processes. 121,122

FeCo-Pc catalysts increase CO₂RR by favoring COOH* over H*, resulting in C₂H₄ generation. During CO reduction, CHO* formation takes precedence over COH* formation. The ratelimiting step for C₂H₄ generation is coupling CHO* with CO* to generate COCHO*.22 FeCo-Pc surfaces aid in producing C2H4 by reducing COCHO* to glycolaldehyde and hydrogenating further. FeCo-Pc catalysts provide a viable avenue for the electrochemical process to transform CO₂ into ethanol (C₂H₅OH), a critical commodity chemical, via C-C coupling reactions. The procedure is optimized by hydrogenating typical intermediates with ethylene (C2H4). CHOHCHO* is found as a selectivitydetermining molecule. Thermodynamically, C2H5OH is produced through the optimum process of CHOHCHO* hydrogenation to CHOHCH2O*.117,123,124

The rate-limiting step (RLS) for C₂H₅OH creation involves the hydrogenation process of CH2CH2O* to C2H4OH*, which has a greater barrier than the formation of C₂H₄. The preference for CHCHO* or CHOHCH2O* production during CHOHCHO* reduction determines the selectivity of C₂H₄ and C₂H₅OH. Increasing potential increases the feasibility of producing C₂H₄ and C₂H₅OH on FeCo-Pc, with all fundamental stages downward energetically at -0.66 V-RHE. Previous investigations have discovered ethylene glycol to be a negligible product in CO₂RR utilizing catalysts such as Au, Ru, and Cu.125-127 Calvinho et al., recently proved that CO₂RR may be converted to ethylene glycol (CH₂OHCH₂OH) using a transition-metal phosphide catalyst. CHOHCHO*, like C₂H₄ and C₂H₅OH, determines the selectivity of CH₂OHCH₂OH production. Protonation of CHOHCH₂O* results in the formation of CHOHCH2OH*, which is preferred over CHCH₂O*. This is then transformed into ethylene glycol. Geometry optimization demonstrates that CH2OH is not chemisorbed, indicating that it prefers the formation pathway over C₂H₅OH. Both CH₂OHCH₂OH and C₂H₅OH have an identical kinetic barrier for CHOHCH2O* production, resulting in

C₂H₄ selectivity. 104,128-130 Possible electrochemical reduction pathways for CO₂ into C₂ products are shown in Scheme 1.

Fe-based bimetallic electrocatalysts: advanced pathways for CO2 reduction

As previously reported, nitrogen-doped carbon nanotubes with Fe/Fe₃N nanoparticles improve the catalytic performance of the oxygen reduction process (ORR) by exposing active areas and enabling electron transport.131-134 Before pyrolysis, Fe-doped zinc-imidazole frameworks (ZIF-8) were changed with phosphomolybdic acid hydrate (PMo), resulting in the formation of Fe nanoparticles contained within molybdenum and nitrogenco-doped carbon scaffolds (Fe-NP/MNCF). In CO2 electrolysis powered by a Zn-air battery (ZAB), Fe-NP/MNCF served as a dual-functional catalyst during ORR and CO₂RR. To synthesize, Fe(NO₃)₃·9H₂O, Zn(NO₃)₂·6H₂O, and PMo were dispersed in 2.5 mL of water that was deionized by ultrasound. The precursor was calcined at 900 °C in an argon environment for 2 hours to produce Fe-NP/MNCF, as illustrated in Fig. 2A. 135,136

Nitrogen-doped carbon was obtained by annealing ZIF-8 at 950 °C in an argon environment. The final product, called ZIF-NC, has a three-dimensional porous structure. 137 FeNi@N-CNTs catalysts were developed by wet impregnation and thermal processing. The heat was applied at 1100 °C to produce FeNi@N-CNTs-X, where X is the temperature at which they were annealed (Fig. 2B). By encasing the FeNi alloy in N-CNTs, this synthesis method enhances the catalytic properties for potential CO₂ reduction uses. Better CO₂ER activity and stability were demonstrated by FeNi@N-CNTs-1100, which showed over 90% CO faradaic efficiency spanning a wide potential range (-0.47 to -0.97 V vs. RHE). Optimized *COOH adsorption and *CO desorption were credited with improving catalytic activity and CO selectivity. 138,139

Xiao Han et al. employed a solution approach to create FeNi precursors, which were subsequently transformed into a variety of FeNi-NC catalysts via one-step pyrolysis. Dissolved $Ni(NO_3)_2 \cdot 6H_2O$, $Zn(NO_3)_2 \cdot 6H_2O$, and $Fe(NO_3)_3 \cdot 9H_2O$ in 80 mL of methanol and stirred thoroughly. Separately, another 80 mL of methanol was used to dissolve 2-methylimidazole and added to the metal nitrate solution. The resultant mixture was agitated constantly for 8 hours to produce the catalyst precipitate. The resulting precipitate washed away with the solvent methanol centrifuged, evaporated at 60 °C, and powdered to produce Fe₃Ni₇-ZIF samples with different Fe/Ni ratios, as illustrated in Fig. 2C. These catalysts attain about 100% overall Faraday efficiency by promoting CO2 electroreduction into CO and H2. Furthermore, we discovered that altering the applied potential across a large range throughout the procedure makes it simple to change the syngas ratio from 1:1 to 6:1 (CO/H₂). Because of its versatility, syngas can be utilized to manufacture fuels and raw materials for chemicals.140

The unique features of doped Cu in Fe-N-C catalysts, including its numerous oxidation states, which facilitate fast electron transfer,87,141 ability to particularly manufacture C2+

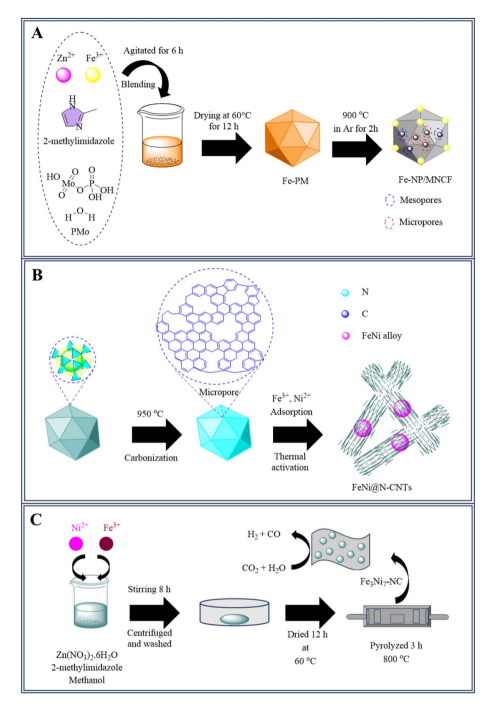


Fig. 2 (A) Synthesis of Fe-NP/MNCF (B) synthesis of FeNi@N-CNTs and (C) synthesis of Fe $_3$ Ni $_7$ -N-C.

products, ¹⁴²⁻¹⁴⁴ and improved interaction with CO_2 to limit hydrogen development, have sparked great interest in CO_2RR . The Fe/Cu–N–C catalyst, which was produced by adding a copper promoter to a mixture of iron and carbon sources and then pyrolyzing it, has outstanding CO_2 reduction efficiency with more than 90% CO faradaic productivity (FE_{CO}) in a broad potential range (-0.5 to -0.7 V) and remarkable stability, with FE_{CO} maintained after 10 hours of electrolysis. To make the Fe/Cu–N–C catalyst, Shulin Zhao *et al.*, mixed tris(2,4-pentanedionato)iron(III), Cu-acetylacetonate, along with meso-tetra(4-

methoxyphenyl) porphin in CHCl $_3$ and stirred it at 60 °C for the period of 3 h. Rotational evaporation was used to extract the solvent from the mixture after 30 minutes of sonication following the addition of zinc oxide. After the powder was produced, it was heated to 900 °C in an argon atmosphere for two hours, then it was leached for six hours at 80 °C in 0.5 M $\rm H_2SO_4$ and allowed to dry overnight. Table 3 shows the comparative analysis of heteronuclear Fe-based catalysts for $\rm CO_2$ electroreduction.

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Comparative analysis of heteronuclear Fe-based catalysts for CO₂ electroreduction

Catalyst	Synthesis method	Key features	CO ₂ reduction efficiency	Stability	Reference
Fe-NP/MNCF	Fe-doped ZIF-8 modified with PMo, pyrolysis at 900 °C	Molybdenum and nitrogen co-doped carbon scaffold	The dual-functional catalyst for ORR and CO ₂ RR	Used in Zn-air battery-powered CO ₂ electrolysis	135 and 136
ZIF-NC	Annealing ZIF-8 at 950 °C	Three-dimensional porous structure	1	1	137
FeNi@N-CNTs-X	Wet impregnation and thermal processing at 1100 °C	Encapsulated FeNi alloy in N-CNTs, enhanced CO ₂ reduction	>90% CO faradaic efficiency (-0.47 to High stability -0.97 V vs. RHE)	High stability	138 and 139
$\mathrm{Fe_3Ni_7} ext{-ZIF}$	Solution approach, pyrolysis	Various Fe/Ni ratios to adjust performance	\sim 100% faraday efficiency, tunable syngas ratio (1:1 to 6:1 CO/H ₂)	High stability, broad potential range	140
Fe/Cu-N-C	Cu promoter added to Fe/carbon mixture, pyrolysis at 900 °C, acid leaching	Enhanced electron transfer, improved ${\rm CO}_2$ adsorption	>90% CO faradaic efficiency (-0.5 to -0.7 V)	Stable after 10 hours of electrolysis	145
C-Fe-Co-ZIF	Impregnation of ZIF-8 with Fe and Co, $$ Bimetallic Co–Fe catalyst for CO $_2$ pyrolysis	Bimetallic Co–Fe catalyst for CO_2 electroreduction	+10% CO faradaic efficiency $vs.$ pure Co-ZIF	+10% CO faradaic efficiency $vs.$ pure Co- $\rm H_2/CO$ ratios tunable (0.8 to 4.2), 93% ZIF	146
Fe/Mn-N-C	Potassium citrate calcination, Fe and Mn doping, pyrolysis at 800 °C	Atomic dispersion of Fe and Mn for CO 94% CO faradaic efficiency at $-0.5~\mathrm{V}$ selectivity (RHE)) 94% CO faradaic efficiency at -0.5 V (RHE)	$>$ 80% FE $_{ m CO}$ after 12 hours	146 and 147

The production of Fe/Mn-N-C, a unique bimetallic catalyst consisting of iron and manganese atomic dispersion, involved the elevated temperatures calcination of an organic carbonbased porous precursor. The solution of potassium citrate monohydrate was initially calcined for an hour at 800 °C in a nitrogen atmosphere to create porous black carbon compounds. The resulting solid was dried in the oven for 12 hours at 80 °C after being rinsed with deionized water and a 1 M H₂SO₄ solution until it attained a neutral pH. A mixture consisting of carbon material, Fe(NO₃)₃·9H₂O, and MnCl₂·4H₂O in deionized H₂O was ultrasonically treated for an hour, centrifuged, and dried afterward. The resultant solid was combined with melamine in a particular mass ratio and then calcined at 800 °C, over a nitrogen environment for two hours to generate the Fe/Mn-N-C catalyst, 146 Fig. 3A. At a -0.5 V overpotential (RHE), the Fe/Mn-N-C catalyst produced a 94% Faraday efficiency (FE) for CO in the 0.1 M KHCO₃ electrolyte. This shows that, in these electrochemical circumstances, the catalyst has a high selectivity for CO synthesis. The catalyst's performance is notable when compared to previously published iron-based and manganese-based electrocatalysts, which include FeMn-N-C (FE_{CO} 80% at -0.5 V RHE), NFe-CNT/CNS (FE_{CO} 69% at -0.6 V RHE), and Mn-N-C (FE $_{CO}$ 70% at -0.6 V RHE). 53,148,149 Following just 12 hours of uninterrupted catalysis, the FE_{CO} was above 80%, suggesting good stability. Density functional theory (DFT) calculations show that the interaction of neighboring Fe-Mn centers lowers the potential for COOH* production and CO desorption.146

6. Development of atomically distributed Co–Fe catalysts for CO₂ reduction

Bimetallic Co–Fe catalysts that are atomically distributed were developed in two steps. Using this method, the catalysts were synthesized with accurate atomic-level dispersion of iron and cobalt through a series of synthesis steps. To ensure a successful yield without interference in the crystallization of Co-ZIF, Fe–Co-ZIF precursors were generated by an impregnation process that modified ZIF-8 into Co-ZIF and absorbed Fe source. Pyrolysis was then used to manufacture the final catalysts (C–Fe–Co-ZIF) for CO_2 electro-reduction, ¹⁵⁰ shown in Fig. 3B. The bimetallic catalysts produced more CO, with an additional 10% in CO Faradaic efficiency (FE) when compared to pure C–Co-ZIF. Adjustable H_2 /CO ratios (0.8 to 4.2) reached across a wide potential range, with a high overall FE CO + H_2 of 93% over 10 hours, showing the catalyst's capacity for efficient syngas production from CO_2 . ¹⁴⁷

7. Graphene oxide-based catalysts for CO₂ reduction

Graphene oxide (GO) was produced with graphite using the modified Hummers' method.¹⁵¹ GO suspension (2 mg mL⁻¹) was made by sonicating it in deionized water for 5 hours. Iron and nickel nitrates were introduced to the GO solution, which

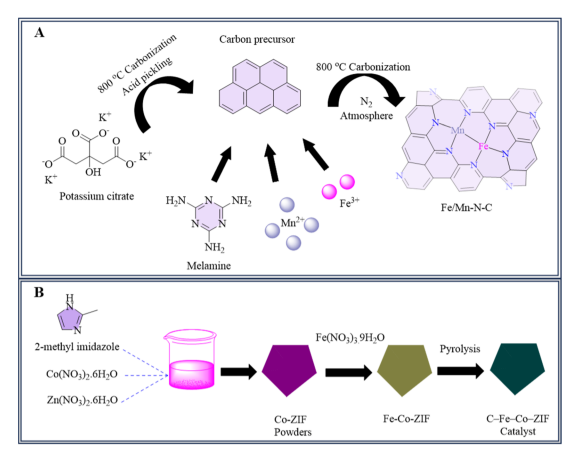


Fig. 3 Methodology for the synthesis of (A) Fe/Mn-N-C and (B) C-Fe-Co-ZIF catalysts

was sonicated for three more hours. The resulting slurry was heated to 180 °C in an autoclave lined with Teflon for 12 h before being freeze-dried to generate a columnar product. Subsequently, a chemical vapor deposition (CVD) process at 1000 °C with Ar and NH₃ was used to synthesize the H-NiFe/NG composite, followed by annealing with hot steam. A novel method involving steam-assisted chemical vapor deposition introduces surface oxygen vacancies (VO) into Ni-Fe BM NPs, creating electron-rich centers that activate CO2 molecules. 152,153 This method reduces the energy barrier for creating COOH* intermediates, increasing the reduction of carbon dioxide to CO while maintaining a faradaic efficiency of as high as 94% at -0.80 V (vs. RHE) along with excellent stability. Surface V_0 modified atoms of nickel have a vital role in increasing the electrocatalytic efficacy of reduction of CO₂ to CO, according to density functional theory simulations.154

8. Molecular catalyst-based heterostructures for CO₂ reduction

The design and synthesis of a molecular catalyst-based heterostructure for the reduction of CO_2 is still a serious issue. Molecular catalysts with transition-metal elements (Co, Ru, Fe, Ni, Cu) and ligands made of organic compounds (phthalocyanine, polypyridine, porphyrin) provide precise active sites and structural tunability for researching $\mathrm{CO}_2\mathrm{ER}$ processes. $^{155-159}$

These catalysts facilitate detailed investigations into CO₂ reduction catalysis. A crystalline bimetallic phthalocyanine heterostructure electrocatalyst (CoPc/FePc HS) was developed for CO2 reduction, achieving a remarkable CO2 to CO conversion efficiency of 99% at the potential of -0.87 vs. RHE and demonstrating outstanding stability over 10 h of electrocatalysis. Different Co/Fe molar ratios (3:1, 1:1, 1:3)160 of CoPc/FePc heterostructures, along with CoPc and FePc controls were synthesized by dispersing a mixture of CoPc and FePc in DMF and subjecting it to solvothermal treatment at 180 °C for 24 hours. Precipitates in the shape of purple microrods were gathered and cleaned with ethanol. They were then calcined for three hours at 450 °C in an Ar environment. CoPc/FePc heterostructures were formed as a consequence of this technique. This method provides a controlled approach to tailor the composition of bimetallic phthalocyanine heterostructures for CO₂ reduction applications. 161

9. $Cu-Fe-N_6-C$: a highperformance diatomic site catalyst for CO_2 reduction

Metal-nitrogen-carbon (M-N-C) catalysts have great potential for CO₂ electrocatalytic reduction because of their abundance of active sites and low-cost raw ingredients.^{52,162-165} Cu-Fe-N₆-C,

a new diatomic site catalyst coordinated with nitrogen and embedded into a carbon matrix, was developed, and synthesized. Cu-Fe-N₆-C was synthesized in two primary stages. First, PcCu-Fe-ZIF-8 is created by combining PcCu, zinc nitrate, iron nitrate, and 2-Me-imidazole, resulting in a blue precipitate that indicates uniform dispersion of Cu and Fe species inside the framework. PcCu-Fe-ZIF-8 becomes Cu-Fe bimetallic sites distributed on a nitrogen-doped carbon framework upon annealing at 1000 °C under Ar. The necessity for extra acid leaching treatment is eliminated by this technique. For a variety of processes, the resulting catalyst structure improves catalytic performance (Fig. 4). This catalyst outperformed individual Cu-N-C and Fe-N-C catalysts thanks to synergistic effects at bimetallic sites. Cu-Fe-N6-C demonstrated outstanding CO selectivity, with an exceptional faradaic efficiency of 98% at

-0.7 V, and maintained selectivity after 10 hours of electrolysis.

Experimental and theoretical investigations revealed that the

combined catalysis of several metallic sites increased CO₂ adsorption enthalpy, and lowered activation energy, resulting in

enhanced selectivity, activity, and stability, as well as decreased

impedance in CO₂ hydrogenation.¹⁶⁶

For CO₂ conversion, some Na-promoted Co-Fe bimetallic catalysts ranging in proximity and compositions were investigated. These catalysts are designed to use the strong selectivity of iron for olefins during CO2 hydrogenation, along with the high activity and reducibility of cobalt. The goal of this combination is to improve CO2 conversion operations' overall efficiency and selectivity. 167-170 Co-precipitation was used to produce Co-Fe bimetallic catalysts, which were then hydrothermally treated. The manufacture of uniform catalysts with regulated compositions and architectures is made easier by this technique. 171,172 Co(NO₃)₂·6H₂O and Fe(NO₃)₃·9H₂O were

dissolved sequentially in deionized water to achieve a [Co]²⁺ + [Fe]³⁺ concentration of 0.09 M, followed by the addition of $5 \text{ mol } L^{-1}$ NaOH solution until pH 11 was reached. The resultant hydroxide precipitates were hydrothermally treated at 150 ° C for 24 hours before being centrifuged, washed, and dried at 80 °C. The dried products were calcined at 400 °C for 3 hours to produce Co-Fe catalysts with various Co/Fe molar ratios (1/4, 1/ 2, 1/1, 2/1, and 4/1), designated as Co1Fe4, Co1Fe2, Co1Fe1, Co2Fe1, and Co4Fe1, respectively. The Co1Fe2 catalyst, having a Co/Fe molar proportion of 1/2 and proximity, permitted the quick reduction of CoFe₂O₄ to Co_xFe_y alloy and subsequently carbonization to χ - $(Co_xFe_{1-x})_5C_2$ alloy carbide. It demonstrated improved stability and performance in olefin production without deactivation over 500 h on-stream.173

Fe/Ni-N-C catalysts with 3D 10. carbon-based structures for CO2 reduction

A 3D carbon-based material was produced, featuring bimetallic centers174 that include NiNC and FeNC, which demonstrated synergistic effects advantageous to the CO2RR. The synthesis procedure involved numerous steps to produce various catalyst materials. Tripotassium citrate monohydrate was cooked at 800 °C under nitrogen, and then treated with sulfuric acid and water to create a porous carbon material. Next, a mixture containing carbon, nickel nitrate, iron nitrate, and glucose in water was processed using ultrasound and then combined with melamine. This mixture was heated at 800 °C under nitrogen to produce the NiNC/FeNC catalyst. 175 Further, specific catalysts like FePc@NiNC and NiPc@FeNC were prepared by treating

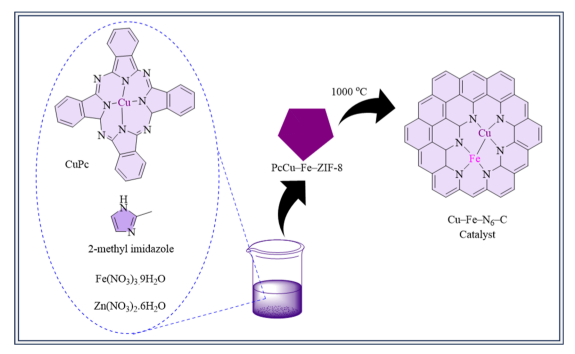


Fig. 4 Synthesis of Cu-Fe-N₆-C.

 K^{\dagger} Calcined at 800 °C 1M H₂SO₄ H_2O N_2 80 °C 12 h Tripotassium citrate monohydrate Carbon НО $Ni(NO_3)_2 \cdot 6H_2O$ Fe(NO₃)₃ HO α-d-glucose Calcined at 800 °C 60 °C for 12 h N₂, 2h FePc@NiNC NiNC/FeNC Catalyst Catalyst N,N-dimethylformamide Carbon porous material ŃΗ Melamine

Fig. 5 Methodology for the synthesis of FePc@NiNC catalyst.

NiNC or FeNC with *N,N*-dimethylformamide and adding iron phthalocyanine (FePc) or nickel phthalocyanine (NiPc), respectively (Fig. 5). Each stage required precise chemical reactions and thermal treatments to generate the correct catalyst compositions. DFT models and observations show¹⁷⁶ Fe atoms are reactive and adsorption sites for CO_2RR , while substantial CO^* adsorption reduces stability. By adding Ni atoms, CO^* adsorption on Fe is decreased, changing the energy barriers and improving stability. The Fe–N₄ and Ni–N₄ sites work in concert to facilitate the rate-limiting processes ($CO_{2(g)} \rightarrow COOH^*, +0.95$ eV) in FePc@NiNC. Flexible syngas composition is made

possible by this synergy while high catalytic activity is maintained.¹⁷⁷

11. Ni/Fe-N-C: diatomic metalnitrogen catalysts for CO₂ reduction

A ZIF-8 was used to create a catalyst consisting of isolated diatomic metal-nitrogen species. Initially, Fe-doped ZIF-8 was made by combining zinc nitrate, iron nitrate, and 2-methylimidazole, maintaining that Fe ions were chemically bound to the organic ligand rather than being physically absorbed.¹⁷⁸ Fe-

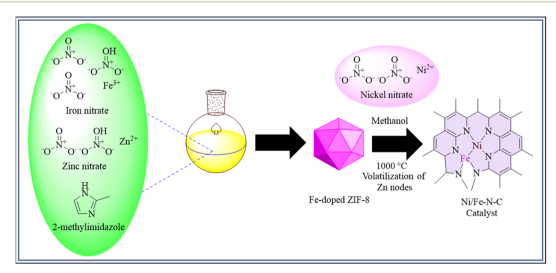


Fig. 6 Methodology for the synthesis of Ni/Fe-N-C catalyst.

Toluene CuCl₂ x 2 H₂O Stirred Stirred overnight 57 FeCl₂ x 2 H₂O AgNO₃

Fig. 7 Synthesis of ⁵⁷FeCu and ⁵⁷FeAg NPs for CO₂ERR.

Toluene

doped ZIF-8 was dissolved in n-hexane, and nickel nitrate methanol solution was added gradually. Nickel was contained within ZIF-8's tiny cavities using this method. Nickel was well incorporated into the framework owing to the steady infusion. 90,179 After thermal treatment at 1000 °C, the resulting catalyst, Ni/Fe-N-C, containing nitrogen-coordinated diatomic Ni-Fe species, was obtained (Fig. 6). For comparison, crystalline Ni-N-C and Fe-N-C catalysts were synthesized similarly. After 30 hours, the Ni/Fe-N-C catalyst retains 99% selectivity and over 90% CO faradaic efficiency from -0.5 to -0.9 V, which ended at 98% at -0.7 V. Synergistic Ni-Fe interactions lower CO₂ reduction reaction barriers and cause structural changes upon CO₂ adsorption, according to DFT research, improving the catalyst's performance.180

12. Micelle-encapsulated Fe-based nanoparticles for CO2 reduction

Inverse micelle encapsulation was used to produce size-selected nanoparticles (NPs) of Fe, FeCu, FeAg, Ag, and Cu. The poly(styrene)-block-poly(2-vinylpyridine) (PS-P2VP) diblock copolymer, which was obtained from Polymer Source Inc., was used in this procedure. Metallic salts (FeCl2, AgNO3, CuCl2, FeCl3) and the copolymer were dissolved in toluene. By taking advantage of the micelles encapsulating attributes, this technique made controlled nanoparticle manufacturing easier. 181,182 Specifically, isotopically enriched ⁵⁷FeCl₂ salt was employed for NRIXS measurements, prepared from iron foil with 95% ⁵⁷Fe isotopic enrichment using adapted literature procedures. 183 Following NP synthesis, the samples were soaked with carbon black powder and then treated using N2-plasma to eliminate the polymer, resulting in clean NP surfaces. The NPs were subsequently distributed into an ethanol/Nafion solution enabling electrode deposition, accompanied by further N2-plasma treatment to remove any remaining polymer before electrochemical evaluation. Fig. 7 shows that the production of 57Fe NPs involves mixing PS-P2VP in toluene to create reverse micelles, which were then added to 57FeCl2 salt and stirred for 72 h. Similar methods were utilized to create 57FeCu and 57FeAg NPs by changing the ratios of 57FeCl2 to CuCl2 or AgNO3 in the micellar solution. FeAg NPs had 36% CO faradaic selectivity at -1.1 V vs.RHE in 0.1 M KHCO₃, similar to pure Ag NPs, but FeCu NPs prefer H₂ evolution, similar to pure Fe NPs. 184 Table 4 summary of recently reported Fe-based bimetallic electrocatalysts for CO2 reduction.

13. Summary and outlook

Possibilities for the advancement of sustainable energy technology look promising for future studies on electrochemical CO₂ reduction with bimetallic catalysts. Optimizing catalyst compositions and structures to increase selectivity and efficiency in the production of CO, syngas, and other multi-carbon

Table 4 Summary of recently reported Fe-based bimetallic electrocatalysts for CO2 reduction

Catalyst	Electrolyte	Major product	FE (%)	Potential (V)	Current density (mA cm ⁻²)	Stability
Fe-NP/MNCF	0.5 M KHCO ₃	CO, H_2	87.50%	-0.7	10	36 h
FeNi@N-CNTs	0.5 M KHCO ₃	CO	90%	-0.47 to -0.97	20.18	35 h
Fe ₃ Ni ₇ -ZIF	0.5 M KHCO ₃	CO, H_2	81.30%	-0.9	-22.5	Good
Fe/Cu-N-C	0.5 M KHCO ₃	CO	97%	-0.6	74	10 h
Fe/Mn-N-C	0.1 M KHCO ₃	CO	94%	-0.5	-83.5	12 h
Fe/Ni-ZIF-8	0.5 M KHCO ₃	CO	89%	-0.677	26.92	40 h
FePc@NiNC	0.5 M KHCO ₃	CO, H_2	100%	-0.8	260	18 h
Ni/Fe-N-C	0.5 M KHCO_3	CO	98%	-0.7	7.4	30 h
H-NiFe/NG	0.1 M KHCO ₃	CO	94%	-0.8	18.2	20 h
PcCu-Fe-ZIF-8	0.1 M KHCO_3	CO	98%	-0.7	7	10 h
⁵⁷ FeAg NPs	0.1 M KHCO ₃	CO	36%	-1.1	0.35	2.5 h
Fe-Co-ZIF	0.5 M KHCO ₃	CO, H_2	93%	-0.55	8	10 h
CuFe/OG	_	CH_4	_	0.97	_	_
FeNi/DG	_	CH_4	_	-0.44	_	_
FeCo-Pc	_	C_{2+}	_	-0.66	_	_

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products is a crucial field of research. Fe-Ni, Fe-Ag, Fe-Mo, Fe-Co, Cu-Fe, and Fe/Mn-N are examples of novel metal combinations that present the potential for enhanced catalytic performance. The main goals of the research will be to comprehend the fundamental structure-activity correlations and stability of these bimetallic catalysts in practical working environments. The scalability of bimetallic catalysts for largescale commercial applications is limited by their typical synthesis, which involves intricate deposition, pyrolysis, and reduction methods. The development of more affordable, optimized synthesis techniques with improved loading capacities is necessary to meet this challenge and permit Fe-based bimetallic catalysts to be widely used in renewable energy systems. Moreover, a major challenge presented by the chemical instability of these catalysts is the reduction of active sites and changed performance caused by corrosion of the carbon substrate. Under practical circumstances, Fe-based bimetallic electrocatalysts for CO₂ reduction encounter difficulties such as low catalytic activity, poor selectivity for target products, and restricted stability. Controlling the chemical intermediates, improving the electrical and geometric properties, and interpreting the synergistic effects between metals are still major challenges. Precise control of Fe-based bimetallic catalysts' shape, structure, and atomic coordination is required to strike a compromise between stability and catalytic activity. The aim is to design specialized bimetallic catalysts with enhanced stability features and active sites outperforming existing catalysts. This will facilitate the development of scalable CO₂ conversion technologies for use in sustainable energy applications, assisting in the shift to a world without carbon emissions. The development of effective CO₂ electroreduction catalysts will be speed up by collaborative, multidisciplinary research that combines theoretical and experimental methods. Future research should concentrate on investigating novel bimetallic combinations that improve performance and customizing catalyst structures by nano-structuring. Enhancing these catalysts' scalability for industrial applications is also essential. Developments in reaction mechanism research, computational modeling, and in situ characterization methods will improve catalyst design and propel more effective CO2 conversion systems.

Data availability

No primary research results, software, or code have been included, and no new data were generated or analyzed as part of this review.

Author contributions

Ayesha Zafar: writing – original draft. Adnan Majeed: writing – review & editing and software. Abdul Ahad: formal analysis. Muhammad Adnan Iqbal: conceptualization, resources, supervision. Tanveer Hussain Bokhari: validation. Zanira Mushtaq: data curation, validation. Shahzaib Ali: visualization.

Conflicts of interest

The authors declare no conflict of interest.

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References

- 1 S. A. Rekker, K. R. O'Brien, J. E. Humphrey and A. C. Pascale, *Nat. Clim. Change*, 2018, **8**, 489–492.
- 2 J. Rogelj, P. M. Forster, E. Kriegler, C. J. Smith and R. Séférian, *Nature*, 2019, **571**, 335–342.
- 3 Z. Chen, G. Zhang, H. Chen, J. Prakash, Y. Zheng and S. Sun, *Renewable Sustainable Energy Rev.*, 2022, **155**, 111922.
- 4 K. Heine, in *The Quaternary in the Tropics: A Reconstruction of the Palaeoclimate*, Springer, 2024, pp. 11–84.
- 5 S. Munassar, Development of pre-operational mesoscale inverse modelling system to quantify CO2 sources and sinks over Europe, PhD dissertation, Friedrich-Schiller-Universität Jena, Jena2024.
- 6 V. P. N. Bharathi, K. Muthuswamy, B. Natarajan, B. Vasudevan, S. Appavu, R. Marimuthu, K. Shanmugam and D. Rajaram, *Fresenius Environ. Bull.*, 2024, 1581.
- 7 A. Lucas, Clim. Risk Manag., 2021, 31, 100257.
- 8 A. Vasileff, C. Xu, Y. Jiao, Y. Zheng and S.-Z. Qiao, *Chem*, 2018, 4, 1809–1831.
- 9 F. Zhou, J. Zhang, Y. Zhang, Y. Wu, Y. Wang and W. Luo, Coord. Chem. Rev., 2024, 509, 215802.
- 10 F. Juliá-Hernández, T. Moragas, J. Cornella and R. Martin, *Nature*, 2017, 545, 84–88.
- 11 K. Sekine and T. Yamada, *Chem. Soc. Rev.*, 2016, **45**, 4524–4532.
- 12 H. Rao, L. C. Schmidt, J. Bonin and M. Robert, *Nature*, 2017, 548, 74–77.
- 13 W. Kim, B. A. McClure, E. Edri and H. Frei, *Chem. Soc. Rev.*, 2016, 45, 3221–3243.
- 14 S. Fang, M. Rahaman, J. Bharti, E. Reisner, M. Robert, G. A. Ozin and Y. H. Hu, *Nat. Rev. Methods Primers*, 2023, 3, 61.
- 15 F. Li, A. Thevenon, A. Rosas-Hernández, Z. Wang, Y. Li, C. M. Gabardo, A. Ozden, C. T. Dinh, J. Li and Y. Wang, *Nature*, 2020, 577, 509–513.
- 16 Y. Chen, X.-Y. Li, Z. Chen, A. Ozden, J. E. Huang, P. Ou, J. Dong, J. Zhang, C. Tian and B.-H. Lee, *Nat. Nanotechnol.*, 2024, 19, 311–318.
- 17 D. M. Koshy, S. S. Nathan, A. S. Asundi, A. M. Abdellah, S. M. Dull, D. A. Cullen, D. Higgins, Z. Bao, S. F. Bent and T. F. Jaramillo, *Angew. Chem.*, 2021, 133, 17613–17621.
- 18 Q. Wang, H. Wang, X. Ren, R. Pang, X. Zhao, L. Zhang and S. Li, *J. Phys. Chem. Lett.*, 2023, **14**, 8421–8427.
- 19 Y. Dong, R. Song, Z. Zhang, X. Han, B. Wang, S. Tao, J. Zhao, A. N. Alodhayb, Z. Chen, X. Yi and N. Zhang, *Cell Rep. Phys. Sci.*, 2024, 5(10), 102227.

20 S. Gleizer, R. Ben-Nissan, Y. M. Bar-On, N. Antonovsky, E. Noor, Y. Zohar, G. Jona, E. Krieger, M. Shamshoum and A. Bar-Even, Cell, 2019, 179, 1255-1263.

- 21 T. E. Miller, T. Beneyton, T. Schwander, C. Diehl, M. Girault, R. McLean, T. Chotel, P. Claus, N. S. Cortina and J.-C. Baret, Science, 2020, 368, 649.
- 22 Y. Y. Birdja, E. Pérez-Gallent, M. C. Figueiredo, A. J. Göttle, F. Calle-Vallejo and M. T. Koper, Nat. Energy, 2019, 4, 732-745.
- 23 L. Zhang, Z.-J. Zhao, T. Wang and J. Gong, Chem. Soc. Rev., 2018, 47, 5423-5443.
- 24 S. Lu, H. Hu, H. Sun, F. Yang, H. Zhu, M. Du, Y. Jin and W. Zhang, Green Chem., 2024, 26(10), 5744-5769.
- 25 B. M. Tackett, E. Gomez and J. G. Chen, Nat. Catal., 2019, 2,
- 26 Z. Xie, L. R. Winter and J. G. Chen, Matter, 2021, 4, 408-440.
- 27 C. S. Diercks, Y. Liu, K. E. Cordova and O. M. Yaghi, Nat. Mater., 2018, 17, 301-307.
- 28 J. Qiao, Y. Liu, F. Hong and J. Zhang, Chem. Soc. Rev., 2014, 43, 631-675.
- 29 E. E. Benson, C. P. Kubiak, A. J. Sathrum and J. M. Smieja, Chem. Soc. Rev., 2009, 38, 89-99.
- 30 H. Takeda, C. Cometto, O. Ishitani and M. Robert, ACS Catal., 2017, 7, 70-88.
- 31 S. Zhang, Q. Fan, R. Xia and T. J. Meyer, Acc. Chem. Res., 2020, 53, 255-264.
- 32 H. Yang, Q. Lin, C. Zhang, X. Yu, Z. Cheng, G. Li, Q. Hu, X. Ren, Q. Zhang and J. Liu, Nat. Commun., 2020, 11, 593.
- 33 M. Cheng, Y. Yu, X. Zhou, Y. Luo and M. Wang, ACS Catal., 2018, 9, 768-774.
- 34 D. E. Polyansky, D. C. Grills, M. Z. Ertem, K. T. Ngo and E. Fujita, ACS Catal., 2022, 12, 1706-1717.
- 35 T. Ouyang, H. H. Huang, J. W. Wang, D. C. Zhong and T. B. Lu, Angew. Chem., 2017, 129, 756-761.
- 36 C. Zhang, P. Gotico, R. Guillot, D. Dragoe, W. Leibl, Z. Halime and A. Aukauloo, Angew. Chem., Int. Ed., 2023, 62, e202214665.
- 37 M. E. Ahmed, S. Adam, D. Saha, J. Fize, V. Artero, A. Dey and C. Duboc, ACS Energy Lett., 2020, 5, 3837-3842.
- 38 J. A. Intrator, D. A. Velazquez, S. Fan, E. Mastrobattista, C. Yu and S. C. Marinescu, Chem. Sci., 2024, 15(17), 6385-6396.
- 39 X. Jiang, X. Li, Y. Kong, C. Deng, X. Li, Q. Hu, H. Yang and C. He, J. Energy Chem., 2023, 76, 462-469.
- 40 Y. Yao, J.-H. Wu, G. Liu, R. Zhang, Z.-S. Yang, S. Gao, T.-C. Lau and J.-L. Zhang, ChemCatChem, 2024, 16(10), e202301705.
- 41 Y. Yao, J. H. Wu, G. Liu, R. Zhang, Z. S. Yang, S. Gao, T. C. Lau and J. L. Zhang, ChemCatChem, 2024, e202301705.
- 42 R. Kortlever, J. Shen, K. J. P. Schouten, F. Calle-Vallejo and M. T. Koper, J. Phys. Chem. Lett., 2015, 6, 4073-4082.
- 43 W. Zhang, Q. Qin, L. Dai, R. Qin, X. Zhao, X. Chen, D. Ou, J. Chen, T. T. Chuong and B. Wu, Angew. Chem., Int. Ed., 2018, 57, 9475-9479.
- 44 J. Zou, C. Y. Lee and G. G. Wallace, Advanced Science, 2021, 8, 2004521.

- 45 J. E. Huang, F. Li, A. Ozden, A. Sedighian Rasouli, F. P. García de Arquer, S. Liu, S. Zhang, M. Luo, X. Wang and Y. Lum, Science, 2021, 372, 1074-1078.
- 46 Y. Jia, F. Li, K. Fan and L. Sun, Adv. Powder Mater., 2022, 1,
- 47 S. Gao, Z. Sun, W. Liu, X. Jiao, X. Zu, Q. Hu, Y. Sun, T. Yao, W. Zhang and S. Wei, Nat. Commun., 2017, 8, 14503.
- 48 T. Li, G. Luo, K. Liu, X. Li, D. Sun, L. Xu, Y. Li and Y. Tang, Adv. Funct. Mater., 2018, 28, 1805828.
- 49 H.-J. Yang, J. Dong, Y.-H. Hong, W.-F. Lin, Z.-Y. Zhou and S.-G. Sun, Electrochem. Commun., 2018, 97, 82-86.
- 50 Q. Cheng, K. Mao, L. Ma, L. Yang, L. Zou, Z. Zou, Z. Hu and H. Yang, ACS Energy Lett., 2018, 3, 1205-1211.
- 51 A. S. Varela, W. Ju and P. Strasser, Adv. Energy Mater., 2018, 8, 1703614.
- 52 X. Wang, Y. Pan, H. Ning, H. Wang, D. Guo, W. Wang, Z. Yang, Q. Zhao, B. Zhang and L. Zheng, Appl. Catal., B, 2020, 266, 118630.
- 53 A. S. Varela, N. Ranjbar Sahraie, J. Steinberg, W. Ju, H. S. Oh and P. Strasser, Angew. Chem., Int. Ed., 2015, 54, 10758-10762.
- 54 Z.-F. Huang, J. Song, Y. Du, S. Xi, S. Dou, J. M. V. Nsanzimana, C. Wang, Z. J. Xu and X. Wang, Nat. Energy, 2019, 4, 329-338.
- 55 J. Wang, L. Gan, W. Zhang, Y. Peng, H. Yu, Q. Yan, X. Xia and X. Wang, Sci. Adv., 2018, 4, eaap7970.
- 56 X. Liang, J. Xiao, W. Weng and W. Xiao, Angew. Chem., 2021, 133, 2148-2152.
- 57 F. Pan, B. Li, E. Sarnello, S. Hwang, Y. Gang, X. Feng, X. Xiang, N. M. Adli, T. Li and D. Su, Nano Energy, 2020, 68, 104384.
- 58 H. Lv, L. Lin, X. Zhang, D. Gao, Y. Song, Y. Zhou, Q. Liu, G. Wang and X. Bao, J. Mater. Chem. A, 2019, 7, 11967-11975.
- 59 Y. Li, B. Hu, C. Xia, W. Q. Xu, J. P. Lemmon and F. Chen, J. Mater. Chem. A, 2017, 5, 20833-20842.
- 60 J. Lu, C. Zhu, C. Pan, W. Lin, J. P. Lemmon, F. Chen, C. Li and K. Xie, Sci. Adv., 2018, 4, eaar5100.
- 61 J. Ko, B.-K. Kim and J. W. Han, J. Mater. Chem. C, 2016, 120, 3438-3447.
- 62 M. Behrens, F. Studt, I. Kasatkin, S. Kühl, M. Hävecker, F. Abild-Pedersen, S. Zander, F. Girgsdies, P. Kurr and B.-L. Kniep, Science, 2012, 336, 893-897.
- 63 Z. Chen, P. Kang, M.-T. Zhang, B. R. Stoner and T. J. Meyer, Energy Environ. Sci., 2013, 6, 813-817.
- 64 S. C. Mandal, K. S. Rawat, S. Nandi and B. Pathak, Catal. Sci. Technol., 2019, 9, 1867-1878.
- 65 H. Peng, M. T. Tang, X. Liu, P. S. Lamoureux, M. Bajdich and F. Abild-Pedersen, Energy Environ. Sci., 2021, 14, 473-
- 66 Y. Hori, H. Wakebe, T. Tsukamoto and O. Koga, Surf. Sci., 1995, 335, 258-263.
- 67 S. C. Mandal and B. Pathak, ACS Appl. Nano Mater., 2021, 4, 11907-11919.
- 68 S. Kar, A. Goeppert and G. S. Prakash, J. Am. Chem. Soc., 2019, 141, 12518-12521.

69 S. C. Mandal, A. Das, D. Roy, S. Das, A. S. Nair and B. Pathak, *Coord. Chem. Rev.*, 2022, 471, 214737.

RSC Advances

- 70 A. Zafar, T. H. Mawat, E. M. Atiyah, M. Adnan Iqbal, A. Majeed, G. Iram, R. Qureshi and S. Khalid, ChemistrySelect, 2024, 9, e202304566.
- 71 A. Zafar, C. Takeda, A. Manzoor, D. Tanaka, M. Kobayashi, Y. Wadayama, D. Nakane, A. Majeed, M. A. Iqbal and T. Akitsu, *Molecules*, 2024, 29, 398.
- 72 H. A. Petersen, T. H. Myren and O. R. Luca, *Inorganics*, 2020, 8, 62.
- 73 B. Hammer and J. K. Norskov, *Nature*, 1995, 376, 238-240.
- 74 H. X. Zhong, J. Wang, Q. Zhang, F. Meng, D. Bao, T. Liu, X. Y. Yang, Z. W. Chang, J. M. Yan and X. B. Zhang, *Adv. Sustainable Syst.*, 2017, 1, 1700020.
- 75 C. Zhu, H. Li, S. Fu, D. Du and Y. Lin, *Chem. Soc. Rev.*, 2016, 45, 517–531.
- 76 Y. Yan, H. Cheng, Z. Qu, R. Yu, F. Liu, Q. Ma, S. Zhao, H. Hu, Y. Cheng and C. Yang, *J. Mater. Chem. A*, 2021, 9, 19489– 19507.
- 77 B. Talukdar, S. Mendiratta, M. H. Huang and C. H. Kuo, Chem. - Asian J., 2021, 16, 2168–2184.
- 78 Q. Li, Y.-C. Wang, J. Zeng, X. Zhao, C. Chen, Q.-M. Wu, L.-M. Chen, Z.-Y. Chen and Y.-P. Lei, *Rare Met.*, 2021, 40, 3442–3453.
- 79 J. Fu, K. Liu, H. Li, J. Hu and M. Liu, *Environ. Chem. Lett.*, 2021, 1–20.
- 80 W. Chen, Y. Wang, Y. Li and C. Li, CCS Chem., 2023, 5, 544–567.
- 81 X.-H. Liu, X.-L. Jia, Y.-L. Zhao, R.-X. Zheng, Q.-L. Meng, C.-P. Liu, W. Xing and M.-L. Xiao, *Advanced Sensor and Energy Materials*, 2023, 100073.
- 82 F. Cheng, X. Zhang, K. Mu, X. Ma, M. Jiao, Z. Wang, P. Limpachanangkul, B. Chalermsinsuwan, Y. Gao and Y. Li, *Energy Technol.*, 2021, **9**, 2000799.
- 83 X. Bao, T. Wang and Y. Yang, *Mater. Chem. Front.*, 2024, 8(3), 627-651.
- 84 H. Tian, Z. Shui, M. A. Raza, L. Zhu and X. Chen, *J. Alloys Compd.*, 2023, **958**, 170544.
- 85 J. K. Nørskov, J. Rossmeisl, A. Logadottir, L. Lindqvist, J. R. Kitchin, T. Bligaard and H. Jonsson, *J. Phys. Chem. B*, 2004, **108**, 17886–17892.
- 86 Y. Li, D. Wang, Y. Guan, H. Liu, Y. Bao, N. Wu, X. Zhao, C. Sun, Z. Li and L. Lei, *Nano Energy*, 2024, **132**, 110370.
- 87 Y. Zhang, Q. Wu, J. Z. Y. Seow, Y. Jia, X. Ren and Z. J. Xu, *Chem. Soc. Rev.*, 2024, 53, 8123–8136.
- 88 A. Mahsud, M. Arif, W. U. Khan, T. Zhang, S. Hussain, M. Azam and Z. Lu, *Mol. Catal.*, 2023, 550, 113526.
- 89 A. Sen and G. Rajaraman, *Inorg. Chem.*, 2023, **62**, 2342–2358.
- 90 C. Zhao, X. Dai, T. Yao, W. Chen, X. Wang, J. Wang, J. Yang, S. Wei, Y. Wu and Y. Li, *J. Am. Chem. Soc.*, 2017, **139**, 8078– 8081.
- 91 X. Wang, S. Ding, T. Yue, Y. Zhu, M. Fang, X. Li, G. Xiao and L. Dai, *Nano Energy*, 2021, 82, 105689.
- 92 M. Duarte, N. Daems, J. Hereijgers, D. Arenas-Esteban, S. Bals and T. Breugelmans, *J. CO₂ Util.*, 2021, **50**, 101583.

- 93 C. Zhang, S. Yang, J. Wu, M. Liu, S. Yazdi, M. Ren, J. Sha, J. Zhong, K. Nie and A. S. Jalilov, Adv. Energy Mater., 2018, 8, 1703487.
- 94 Y. Cheng, S. Zhao, H. Li, S. He, J.-P. Veder, B. Johannessen, J. Xiao, S. Lu, J. Pan and M. F. Chisholm, *Appl. Catal.*, B, 2019, 243, 294–303.
- 95 L. R. L. Ting and B. S. Yeo, *Curr. Opin. Electrochem.*, 2018, **8**, 126–134.
- 96 N. Li, X. Chen, W.-J. Ong, D. R. MacFarlane, X. Zhao, A. K. Cheetham and C. Sun, ACS Nano, 2017, 11, 10825– 10833.
- 97 H. Zhang, J. Li, S. Xi, Y. Du, X. Hai, J. Wang, H. Xu, G. Wu, J. Zhang and J. Lu, *Angew. Chem.*, 2019, 131, 15013–15018.
- 98 L. J. Arachchige, Y. Xu, Z. Dai, X. L. Zhang, F. Wang and C. Sun, *J. Mater. Sci. Technol.*, 2021, 77, 244–251.
- 99 L. Jasin Arachchige, Y. Xu, Z. Dai, X. Zhang, F. Wang and C. Sun, J. Mater. Chem. C, 2020, 124, 15295–15301.
- 100 S. Dongare, N. Singh and H. Bhunia, Appl. Surf. Sci., 2021, 556, 149790.
- 101 R. Zhang, Y. Zhang, L. Liu, X. Li, Y. Tang, Y. Ni, C. Sun and H. Zhang, *Appl. Surf. Sci.*, 2022, **582**, 152472.
- 102 X. Cui, W. An, X. Liu, H. Wang, Y. Men and J. Wang, *Nanoscale*, 2018, **10**, 15262–15272.
- 103 W. Bi, C. Wu and Y. Xie, ACS Energy Lett., 2018, 3, 624-633.
- 104 L. Guo and S. Guo, *Int. J. Hydrogen Energy*, 2024, **51**, 1532–1544.
- 105 M. J. E. A. Frisch, gaussian 09, Revision d. 01, Gaussian. Inc, Wallingford CT 201, 2009.
- 106 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865.
- 107 G. Rostamikia, A. J. Mendoza, M. A. Hickner and M. J. Janik, J. Power Sources, 2011, 196, 9228–9237.
- 108 Y. Zhao, X. Chang, A. S. Malkani, X. Yang, L. Thompson, F. Jiao and B. Xu, *J. Am. Chem. Soc.*, 2020, **142**, 9735–9743.
- 109 X. Nie, M. R. Esopi, M. J. Janik and A. Asthagiri, *Angew. Chem., Int. Ed.*, 2013, 52(9), 2611.
- 110 Q. Zhao and E. A. Carter, J. Chem. Theory Comput., 2020, 16, 6528–6538.
- 111 A. D. Handoko, F. Wei, Jenndy, B. S. Yeo and Z. W. Seh, *Nat. Catal.*, 2018, **1**(12), 922–934.
- 112 R. Kas, O. Ayemoba, N. J. Firet, J. Middelkoop, W. A. Smith and A. Cuesta, *ChemPhysChem*, 2019, 20, 2904–2925.
- 113 Y. Zheng, A. Vasileff, X. Zhou, Y. Jiao, M. Jaroniec and S.-Z. Qiao, *J. Am. Chem. Soc.*, 2019, **141**, 7646–7659.
- 114 T. K. Todorova, M. W. Schreiber and M. Fontecave, *ACS Catal.*, 2019, **10**, 1754–1768.
- 115 H. Xiao, T. Cheng, W. A. Goddard III and R. Sundararaman, *J. Am. Chem. Soc.*, 2016, **138**, 483–486.
- 116 H. Xiao, T. Cheng and W. A. Goddard III, *J. Am. Chem. Soc.*, 2017, **139**, 130–136.
- 117 A. J. Garza, A. T. Bell and M. Head-Gordon, *ACS Catal.*, 2018, **8**, 1490–1499.
- 118 J. Zhao, J. Zhao, F. Li and Z. Chen, *J. Mater. Chem. C*, 2018, 122, 19712–19721.
- 119 Z. Zhao and G. Lu, ACS Catal., 2018, 8, 3885-3894.
- 120 Y. Feng, W. An, Z. Wang, Y. Wang, Y. Men and Y. Du, *ACS Sustainable Chem. Eng.*, 2019, **8**, 210–222.

121 H. Mei, H. Cheng, Z. Wang and J. Li, *Chem. Eng. Sci.*, 2017, **164**, 81–89.

- 122 X.-F. Qiu, H.-L. Zhu, J.-R. Huang, P.-Q. Liao and X.-M. Chen, *J. Am. Chem. Soc.*, 2021, **143**, 7242–7246.
- 123 T. Cheng, H. Xiao and W. A. Goddard III, *Proc. Natl. Acad. Sci. U. S. A.*, 2017, **114**, 1795–1800.
- 124 K. P. Kuhl, E. R. Cave, D. N. Abram and T. F. Jaramillo, *Energy Environ. Sci.*, 2012, 5, 7050–7059.
- 125 J. Tamura, A. Ono, Y. Sugano, C. Huang, H. Nishizawa and S. Mikoshiba, *Phys. Chem. Chem. Phys.*, 2015, **17**, 26072–26078.
- 126 K.-i. Tominaga, Y. Sasaki, T. Watanabe and M. Saito, *J. Chem. Soc. Chem. Commun.*, 1995, 1489–1490.
- 127 K. Schouten, Y. Kwon, C. Van Der Ham, Z. Qin and M. Koper, *Chem. Sci.*, 2011, 2, 1902–1909.
- 128 K. U. Calvinho, A. W. Alherz, K. M. Yap, A. B. Laursen, S. Hwang, Z. J. Bare, Z. Clifford, C. B. Musgrave and G. C. Dismukes, *J. Am. Chem. Soc.*, 2021, **143**, 21275–21285.
- 129 X. Xu, Z. Xia, X. Zhang, H. Li, S. Wang and G. Sun, *Nanoscale*, 2020, **12**, 3418–3423.
- 130 M. Qiao, Y. Wang, Q. Wang, G. Hu, X. Mamat, S. Zhang and S. Wang, *Angew. Chem., Int. Ed.*, 2020, **59**, 2688–2694.
- 131 Y. Zheng, F. He, J. Wu, D. Ma, H. Fan, S. Zhu, X. Li, Y. Lu, Q. Liu and X. Hu, ACS Appl. Nano Mater., 2019, 2, 3538– 3547.
- 132 H. Tan, J. Tang, J. Henzie, Y. Li, X. Xu, T. Chen, Z. Wang, J. Wang, Y. Ide and Y. Bando, ACS Nano, 2018, 12, 5674– 5683.
- 133 M. Lefèvre, E. Proietti, F. Jaouen and J.-P. Dodelet, *science*, 2009, **324**, 71–74.
- 134 B.-C. Hu, Z.-Y. Wu, S.-Q. Chu, H.-W. Zhu, H.-W. Liang, J. Zhang and S.-H. Yu, *Energy Environ. Sci.*, 2018, 11, 2208–2215.
- 135 T. Wang, Q. Zhang, K. Lian, G. Qi, Q. Liu, L. Feng, G. Hu, J. Luo and X. Liu, *J. Colloid Interface Sci.*, 2024, 655, 176– 186.
- 136 G. Li, X. Wan, Q. Zheng, M. Yang, Y. Xia, X. Qi, T. Wang and Z. Wu, *Colloids Surf.*, *A*, 2024, **699**, 134713.
- 137 E. Luo, H. Zhang, X. Wang, L. Gao, L. Gong, T. Zhao, Z. Jin, J. Ge, Z. Jiang and C. Liu, *Angew. Chem.*, 2019, 131, 12599– 12605.
- 138 L. Zhang, B. Geng, Y. Gao, H. Kang, P. Wang, C. Liu, H. Xiao, M. Zhao, J. Jia and H. Wu, *Chem. Eng. J.*, 2024, 481, 148086.
- 139 G. Li, X. Qi, G. Zhang, S. Wang, K. Li, J. Wu, X. Wan, Y. Liu and Q. Li, *Microchem. J.*, 2022, **179**, 107515.
- 140 X. Han, Y. Chang, T. Yue, M. Jia and J. Jia, *J. Alloys Compd.*, 2024, **971**, 172772.
- 141 W. Fan, Z. Li, C. You, X. Zong, X. Tian, S. Miao, T. Shu, C. Li and S. Liao, *Nano energy*, 2017, 37, 187–194.
- 142 Z.-Z. Wu, X.-L. Zhang, Z.-Z. Niu, F.-Y. Gao, P.-P. Yang, L.-P. Chi, L. Shi, W.-S. Wei, R. Liu and Z. Chen, *J. Am. Chem. Soc.*, 2021, 144, 259–269.
- 143 S. Zhang, S. Zhao, D. Qu, X. Liu, Y. Wu, Y. Chen and W. Huang, *Small*, 2021, **17**, 2102293.
- 144 C. E. Creissen and M. Fontecave, *Nat. Commun.*, 2022, **13**, 2280.

- 145 S. Zhao, Y. Sun, K. Lu, J. Wang, M. Qiao, Y. Wang, Y. Huang, Y. Wu and Y. Chen, *J. CO₂ Util.*, 2023, **70**, 102420.
- 146 C. Ma, H. Zhang, W. Kong, B. Shen and H. Lyu, *Environ. Funct. Mater.*, 2022, **1**, 284–297.
- 147 Z. Chen, G. Zhang, Y. Wen, N. Chen, W. Chen, T. Regier, J. Dynes, Y. Zheng and S. Sun, *Nano-Micro Lett.*, 2022, 14, 25.
- 148 F. Pan, H. Zhao, W. Deng, X. Feng and Y. Li, *Electrochim. Acta*, 2018, 273, 154–161.
- 149 F. Pan, W. Deng, C. Justiniano and Y. Li, *Appl. Catal., B*, 2018, **226**, 463–472.
- 150 X. Wang, Z. Chen, X. Zhao, T. Yao, W. Chen, R. You, C. Zhao, G. Wu, J. Wang and W. Huang, *Angew. Chem.*, 2018, 130, 1962–1966.
- 151 D. C. Marcano, D. V. Kosynkin, J. M. Berlin, A. Sinitskii, Z. Sun, A. Slesarev, L. B. Alemany, W. Lu and J. M. Tour, ACS Nano, 2010, 4, 4806–4814.
- 152 Z. Geng, X. Kong, W. Chen, H. Su, Y. Liu, F. Cai, G. Wang and J. Zeng, *Angew. Chem.*, 2018, 130, 6162–6167.
- 153 S. Huygh, A. Bogaerts and E. C. Neyts, *J. Mater. Chem. C*, 2016, **120**, 21659–21669.
- 154 Q. He, Y. Zhang, H. Li, Y. Yang, S. Chen, W. Yan, J. Dong, X. M. Zhang and X. Fan, Small, 2022, 18, 2108034.
- 155 C. Costentin, S. Drouet, M. Robert and J.-M. Savéant, *Science*, 2012, 338, 90-94.
- 156 S. Lin, C. S. Diercks, Y.-B. Zhang, N. Kornienko, E. M. Nichols, Y. Zhao, A. R. Paris, D. Kim, P. Yang and O. M. Yaghi, *Science*, 2015, 349, 1208–1213.
- 157 C. Yang, S. Li, Z. Zhang, H. Wang, H. Liu, F. Jiao, Z. Guo, X. Zhang and W. Hu, *Small*, 2020, 16, 2001847.
- 158 H.-x. Zhong, Q. Zhang, J. Wang, X.-b. Zhang, X.-l. Wei, Z.-j. Wu, K. Li, F.-l. Meng, D. Bao and J.-m. Yan, ACS Catal., 2018, 8, 3965–3970.
- 159 K.-H. Liu, H.-X. Zhong, S.-J. Li, Y.-X. Duan, M.-M. Shi, X.-B. Zhang, J.-M. Yan and Q. Jiang, *Prog. Mater. Sci.*, 2018, **92**, 64–111.
- 160 Y. Ma, J. Li, X. Liao, W. Luo, W. Huang, J. Meng, Q. Chen, S. Xi, R. Yu and Y. Zhao, Adv. Funct. Mater., 2020, 30, 2005000.
- 161 C. Yang, Z. Gao, D. Wang, S. Li, J. Li, Y. Zhu, H. Wang, W. Yang, X. J. Gao and Z. Zhang, Sci. China Mater., 2022, 65, 155–162.
- 162 J. Wang, X. Huang, S. Xi, H. Xu and X. Wang, *Angew. Chem., Int. Ed.*, 2020, **59**, 19162–19167.
- 163 Q. Wang, Y. Lei, D. Wang and Y. Li, *Energy Environ. Sci.*, 2019, **12**, 1730–1750.
- 164 J. Li, Y. Kuang, Y. Meng, X. Tian, W.-H. Hung, X. Zhang, A. Li, M. Xu, W. Zhou and C.-S. Ku, *J. Am. Chem. Soc.*, 2020, 142, 7276–7282.
- 165 L. Chen, C. He, R. Wang, Q. Li, J. Zeng, W. Liu, Y. Wang, Q. Wang, T. Ye and Y. Tang, *Chin. Chem. Lett.*, 2021, 32, 53–56.
- 166 R. Yun, F. Zhan, X. Wang, B. Zhang, T. Sheng, Z. Xin, J. Mao, S. Liu and B. Zheng, *Small*, 2021, 17, 2006951.
- 167 H. Yang, Y. Dang, X. Cui, X. Bu, J. Li, S. Li, Y. Sun and P. Gao, *Appl. Catal., B*, 2023, **321**, 122050.

168 F. Song, X. Yong, X. Wu, W. Zhang, Q. Ma, T. Zhao, M. Tan, Z. Guo, H. Zhao and G. Yang, *Appl. Catal.*, B, 2022, 300, 120713.

- 169 A. S. Ismail, M. Casavola, B. Liu, A. Gloter, T. W. van Deelen, M. Versluijs, J. D. Meeldijk, O. Stéphan, K. P. de Jong and F. M. de Groot, ACS Catal., 2019, 9, 7998–8011.
- 170 R. Satthawong, N. Koizumi, C. Song and P. Prasassarakich, *Catal. Today*, 2015, **251**, 34–40.
- 171 G. Wu, J. Wang, W. Ding, Y. Nie, L. Li, X. Qi, S. Chen and Z. Wei, *Angew. Chem.*, *Int. Ed.*, 2016, **55**, 1340–1344.
- 172 M. Ding, Asian J. Chem., 2014, 26, 1808.

RSC Advances

- 173 N. Liu, J. Wei, J. Xu, Y. Yu, J. Yu, Y. Han, K. Wang, J. I. Orege, Q. Ge and J. Sun, *Appl. Catal.*, *B*, 2023, 328, 122476.
- 174 H. Zhong, M. Ghorbani-Asl, K. H. Ly, J. Zhang, J. Ge, M. Wang, Z. Liao, D. Makarov, E. Zschech and E. Brunner, *Nat. Commun.*, 2020, **11**, 1409.
- 175 W. Cheng, P. Yuan, Z. Lv, Y. Guo, Y. Qiao, X. Xue, X. Liu, W. Bai, K. Wang and Q. Xu, *Appl. Catal.*, B, 2020, 260, 118198.
- 176 W. Ju, A. Bagger, G.-P. Hao, A. S. Varela, I. Sinev, V. Bon, B. Roldan Cuenya, S. Kaskel, J. Rossmeisl and P. Strasser, *Nat. Commun.*, 2017, **8**, 1–9.

- 177 X. Yang, T. Tat, A. Libanori, J. Cheng, X. Xuan, N. Liu, X. Yang, J. Zhou, A. Nashalian and J. Chen, *Mater. Today*, 2021, 45, 54–61.
- 178 F. Pan, H. Zhang, K. Liu, D. Cullen, K. More, M. Wang, Z. Feng, G. Wang, G. Wu and Y. Li, *ACS Catal.*, 2018, **8**, 3116–3122.
- 179 X. C. Huang, Y. Y. Lin, J. P. Zhang and X. M. Chen, *Angew. Chem., Int. Ed.*, 2006, **45**, 1557–1559.
- 180 W. Ren, X. Tan, W. Yang, C. Jia, S. Xu, K. Wang, S. C. Smith and C. Zhao, *Angew. Chem., Int. Ed.*, 2019, **58**, 6972–6976.
- 181 J. R. Croy, S. Mostafa, J. Liu, Y.-h. Sohn and B. Roldan Cuenya, *Catal. Lett.*, 2007, **118**, 1–7.
- 182 B. Roldan Cuenya, J. R. Croy, S. Mostafa, F. Behafarid, L. Li, Z. Zhang, J. C. Yang, Q. Wang and A. I. Frenkel, *J. Am. Chem. Soc.*, 2010, **132**, 8747–8756.
- 183 G. Winter, D. Thompson and J. Loehe, *Inorganic syntheses*, 1973, **14**, 99–104.
- 184 S. Kunze, P. Grosse, M. Bernal Lopez, I. Sinev, I. Zegkinoglou, H. Mistry, J. Timoshenko, M. Y. Hu, J. Zhao and E. E. Alp, Angew. Chem., 2020, 132, 22856– 22863.