The chemical transformation of carbon dioxide (CO₂) has been considered as a promising strategy to utilize and further upgrade it to value-added chemicals, aiming at alleviating global warming. In this regard, sustainable driving forces (i.e., electricity and sunlight) have been introduced to convert CO₂ into various chemical feedstocks. Electrocatalytic CO₂ reduction reaction (CO₂RR) can generate carbonaceous molecules (e.g., formate, CO, hydrocarbons, and alcohols) via multiple-electron transfer. With the assistance of extra light energy, photoelectrocatalysis effectively improve the kinetics of CO₂ conversion, which not only decreases the overpotentials for CO₂RR but also enhances the lifespan of photo-induced carriers for the consecutive catalytic process. Recently, rational-designed catalysts and advanced characterization techniques have emerged in these fields, which make CO₂-to-chemicals conversion in a clean and highly-efficient manner.

Herein, this review timely and thoroughly discusses the recent advancements in the practical conversion of CO₂ through electro- and photoelectrocatalytic technologies in the past 5 years. Furthermore, the recent studies of operando analysis and theoretical calculations are highlighted to gain systematic insights into CO₂RR. Finally, the challenges and perspectives in the fields of CO₂ (photo)electrocatalysis are outlined for their further development.

1. Introduction

The increasing CO₂ emission in the ambient air has caused deteriorative global warming and ocean acidification problems. Accordingly, CO₂ conversion has been regarded as an effective approach to alleviate atmospheric CO₂ concentration in the past decades, aimed at maintaining the ecosystem carbon cycle. Inspired by natural photosynthesis, CO₂ capture and utilization via chemical reaction path have become a hot research topic in recent years, especially advanced technologies and rational-designed catalysts have attracted wide attention to maximize the efficiency of CO₂ conversion.

Traditional thermocatalytic CO₂ transformation as a popular approach has been developed to reduce CO₂ and further upgrade to important chemicals and fuels in an industrial scale. However, the application of thermocatalytic process is significantly affected by the excess combustion of fossil fuels, which can severely interrupt the carbon cycle and sustainable energy production. Hence, as clean and renewable energy techniques, electro- and photoelectrocatalysis have emerged for CO₂ reduction reaction (CO₂RR), which can be processed in a mild reaction condition (i.e., room temperature and ambient pressure) to realize CO₂ reduction to value-added chemicals production. Notably, CO₂ as an inert molecule requires excessive energy input to activate CO₂, and thus highly-efficient catalysts are required to provide active
conductive interfaces can accelerate the C = O activation process, which reduces CO₂ to hydrocarbons by the photogenerated carriers. Specifically, a semiconductor photocatalyst with a suitable band structure is of great significance to break the C = O bond. The bottom of the conduction band of a semiconductor should be more negative than the CO₂ reduction potential, while the top of the valence band should be more positive than the water oxidation potential, which could thereby simultaneously realize CO₂-to-fuels conversion and oxygen evolution reaction (OER). Notably, photoelectrocatalysis extends the material choices in comparison to photocatalysis, which results in a higher potential to meet the requirements of suitable band position and redox potential level for CO₂ conversion with the assistance of an external bias. Meanwhile, photoelectrocatalysis provides extra light energy to decrease the overpotentials for electrocatalytic CO₂ RR, which can also enhance the lifespan of photo-induced carriers in the whole catalytic process. Accordingly, some approaches (e.g., structure engineering, cocatalyst doping, and heterojunction design) have been employed to optimize the activity of catalysts, and thereby generate different carbonaceous compounds like CO, formate, alcohols, and hydrocarbons. In Figure 2, we summarize the major characteristics and working principle of the full-cell type of electro- and photoelectrocatalysis toward CO₂ conversion.[11,12a,14]

Herein, we present a timely and comprehensive review of the recent advances in electro- and photoelectrocatalytic CO₂ conversion during the past 5 years. First, the motivation and fundamentals of the catalytic technologies are mentioned. Then, since the developments of catalysts play a significant role in CO₂ electro- and photoelectrocatalysis, the important advances in design principles of catalysts have been emphasized in this review. There are some key points mentioned on synthetic strategies for preparing the (photo)electrodes and the main factors for improving the structure-activity relationships in these catalytic systems. Thirdly, the deep understanding of catalytic mechanism and reaction pathways of CO₂ reduction are discussed, mainly based on advanced techniques of in situ/operando characterizations and theoretical calculations. Last but not least, the challenges and future perspectives are addressed to promote the highly-efficient CO₂ utilization and upgradation, aimed at realizing the feasibility for industrial production. We hope this review can shed light on the chemical transformation of CO₂ molecules with the help of clean and renewable energy, and provide sufficient inspiration for researchers in this exciting field.

2. Electrocatalysis

2.1. Motivation and Principles

Although CO₂ thermocatalysis is the most popular catalytic route for large-scale industrial manufacture, whereas the enormous energy demand is often supplied via the excessive calcination of non-renewable fossil fuels. Therefore, converting CO₂ into fuels and chemicals in a more sustainable manner that is driven by renewable energy technologies, has gained extensive attention. Since the cost of electricity generated from solar, wind, and other clean energy sources continuously declines, electrocatalysis emerges as a greener strategy for CO₂ reduction and gains enormous attention in recent years. CO₂ is a thermodynamic stable molecule that requires excess energy of 806 kJ mol⁻¹ to ac-

Figure 1. Chronology of the electro- and photoelectrochemical CO₂ conversion. The research progresses in the recent 5 years (2016–2021) are highlighted in bold.[6]
tivate the C = O bond. Moreover, the first proton-coupled electron transfer process needs to overcome high energy barriers to form the surface-adsorbed species (i.e., *COOH and *OCHO intermediates). Accordingly, the external potentials should be applied to realize CO₂ activation and conversion at reasonable rates. However, different types of carbonaceous compounds could be simultaneously generated through multi-electrons (2, 4, 6, 8, 12, or even more) transfer pathways in CO₂ reduction due to the close thermodynamic redox potentials of these possible products. Table 1 summarizes the possible half-reactions of CO₂ reduction in aqueous solutions for the different hydrocarbons at 25 °C, 1 atm, and pH 7; Reproduced with permission. [15] Copyright 2019, Wiley-VCH.

### Table 1. The standard potentials (E°) of possible half-reactions of electrochemical CO₂ reduction in aqueous solutions for the different hydrocarbons at 25 °C, 1 atm, and pH 7.

| Possible half-reactions of ECR | E° (V vs SHE) |
|--------------------------------|---------------|
| CO₂ (g) + e⁻ → CO₂⁺⁻ | −1.90 |
| CO₂ (g) + 2H⁺ + 2e⁻ → HCOOH (l) | −0.55 |
| CO₂ (g) + 2H⁺ + 2e⁻ → CO (g) + H₂O (l) | −0.52 |
| CO₂ (g) + 4H⁺ + 2e⁻ → HCHO (l) + H₂O (l) | −0.48 |
| CO₂ (g) + 6H⁺ (l) + 6e⁻ → CH₃OH (l) + H₂O (l) | −0.38 |
| CO₂ (g) + 8H⁺ + 8e⁻ → CH₄ (g) + 2H₂O (l) | −0.24 |
| 2CO₂ (g) + 12H⁺ + 12e⁻ → C₂H₆ (g) + 4H₂O (l) | −0.38 |
| 2CO₂ (g) + 12H⁺ + 12e⁻ → C₂H₅OH (l) + 3H₂O (l) | −0.35 |
| 2CO₂ (g) + 14H⁺ + 14e⁻ → C₂H₆ (l) + 4H₂O (l) | −0.28 |
| 3CO₂ (g) + 18H⁺ + 18e⁻ → C₃H₇OH (l) + 5H₂O (l) | −0.30 |

The economic benefits of electrochemical CO₂ reduction (ECR) to different products depend on not only the market demand of the higher-value-added chemicals (C₂-C₄) but also the product separation costs, the further decrease of electricity costs, and the improvement of the electrocatalytic activity. Accordingly, different reaction pathways toward ECR were studied experimentally and theoretically in virtue of the applied electrocatalysts and reaction conditions, resulting in significantly different product distributions, as illustrated in Figure 4. [7,16] First, CO₂ molecules are adsorbed on the surface of catalysts and then activated via the proton-coupled electron transfer process to generate active intermediates (i.e., *OCHO and *COOH). Specifically, *OCHO species will form once the O atom of the activated *CO₂ species bonds to the electrocatalyst surface, and meanwhile, the C atom is protonated.

**Figure 2.** a) Characteristics of the technologies for electro- and photoelectrocatalytic CO₂ conversion. b) Schematic illustration of the working principles of the full-cell type (photo-)electrochemical CO₂ conversion systems. Reproduced with permission. [14] Copyright 2020, American Chemical Society.
2.2. Plausible Routes for CO₂ Conversion

Depending on the ECR products, the applied catalysts can be categorized into three classes to realize CO₂ activation and conversion. The following contents mainly focus on these pathways, and the special attention lies on the developed heterogeneous catalysts (e.g., metals, carbon materials) since the homogeneous ones always realize ECR in aprotic solution (for example, expensive ionic liquid) and are unsuitable for practical applications.

2.2.1. CO₂ to Formate/Formic Acid

The conversion of CO₂ to formate/formic acid has been regarded as an ideal choice for hydrogen carriers and HCOOH fuel cells, due to the advantages of easy storage and high safety. Realizing CO₂ electroreduction into formate/formic acid involves two elementary steps. First, CO₂ molecules go through a proton-electron pair transfer process to obtain an *OCHO intermediate. Then, the *OCHO intermediate further converts to *HCOOH via the subsequent proton-electron transfer at pH < 3.75 and thus generating formate/formic acid that can easily escape from the catalyst surface. However, the products could exist in the deprotonated form (HCOO⁻) at pH > 3.75. In recent five years, nanostructured metals like Sn, Bi, In, Pb, Sb, Pd, Co have been reported to be promising candidates for formate/formic acid production in ECR (Figure 5). The special focuses lie in the studies of their nanostructures, index planes, vacancies/defects, grain boundaries, and supports to achieve a high activity, satisfying selectivity, and long-term stability.

I) Sn-based materials have attracted wide attention owing to their abundance and environment-friendly properties. Many studies have concentrated on the optimal design of Sn, Sn oxides, and Sn sulfides experimentally and theoretically. For example, mesoporous SnO₂ nanosheets were fabricated by a self-templated synthetic method,[17] which possesses a large surface area and 3D hierarchical nanostructure, resulting in numerous undercoordinated sites or structural defects. Owing to the above merits, mesoporous SnO₂ nanosheets exhibited small overpotential (710 mV), high faradaic efficiency of formate (FE_{\text{formate}} = 83.0%), and long-term stability in ECR. SnS₂ monolayers were synthesized by He et al. via Li-intercalation and exfoliation process, which exhibited an extraordinary FE_{\text{formate}} up to 94% and excellent durability over 80 h in ECR.[18] The atomic-scale thickness accelerated the proton-electron transfer efficiency and the formation of *OCHO and *HCOOH intermediates, which were subsequently transformed into formate via a two-electron transfer pathway. In addition, the introduction of grain boundary could break the local spatial symmetry of materials, which significantly optimizes the binding energies of reaction intermediates, and thus realizes a high selectivity and energy efficiency for CO₂-to-formate conversion. Inspired by this, sub-2 nm ultrathin 1D SnO₂ quantum wires constructed by numerous quantum dots were fabricated,[19] presenting abundant grain boundaries on the catalyst surface (Figure 6a). The as-prepared SnO₂ quantum wires exhibited a...
peak value of \( \text{FE}_{\text{formate}} \) (87.3%) and energy efficiency (EE = 52.7%). Moreover, the \( \text{FE}_{\text{formate}} \) and \( \text{EE}_{\text{formate}} \) could maintain above 80% and 50% in a wide potential window, respectively (Figure 6b,c). Furthermore, the creation of structural defects such as oxygen vacancies in the catalysts was found to effectively improve \( \text{CO}_2 \) activation, electron mobility, the interaction between reaction intermediates and active sites, as well as desorption ability of \( \text{HCOOH} \) intermediates.\(^{20}\)

Daiyan et al. employ the industrially adopted flame spray pyrolysis technique to synthesize \( \text{SnO}_2 \) nanoparticles with the active oxygen hole center,\(^{21}\) realizing a high \( \text{FE}_{\text{formate}} \) of 85% and a current density of \(-23.7\ \text{mA cm}^{-2}\) at a potential of \(-1.1\ \text{V} \) versus RHE. Similarly, Chen’s group reported an “all in one” wavy \( \text{SnO}_2 \) network catalyst,\(^{22}\) which simultaneously possessed grain boundaries, oxygen vacancies, and low-coordinated active edge/corner sites. With
II) The potential of metallic Bi has recently been explored for highly selective formate production from ECR in an aqueous solution, which used to be commonly applied to CO₂-to-CO conversion in aprotic electrolytes before 2016. Element Bi locates close to the Sn in the periodic table and possibly provides similar electronic properties in ECR. Moreover, metallic Bi could suppress the competitive H₂ evolution since it exhibits high free energy of hydrogen adsorption (ΔGΘ). Bi-based catalysts always contain oxygen species, which are inevitably introduced during the synthetic process. Accordingly, Deng et al. reported a Bi oxides catalyst with a high concentration of Bi-O species by applying a time-dependent oxidation treatment.[23] The roles of the Bi-O species toward highly selective formate production in ECR were analyzed experimentally and theoretically (Figure 6d). The Bi-O structure was found to facilitate CO₂ adsorption and accelerate the rate-determining step (namely, *CO₂* + *H* → *OCHO*) by lowering the free energy for intermediate formation (Figure 6e). Therefore, the resulting Bi₂O₃ catalyst showed a maximum FE formate of 91% and a partial current density of 8 mA cm⁻² at −0.9 V versus RHE (Figure 6f). To further improve the durability of metastable Bi oxides, the impact of the morphology of Bi₂O₃ was investigated, demonstrating that the optimized Bi₂O₃ nanoparticles could achieve improved FE formate of 91% at the applied potential of −1.2 V versus RHE and stable performance over 23 hours.[24] To design a high-performance Bi-based catalyst for ECR, the coordinately unsaturated sites such as high-indexed planes and edge/corner sites can be incorporated onto the surface of the Bi electrodes, which enhances the stabilization of reaction intermediates onto the catalyst surface, and thus promotes CO₂ conversion. Inspired by this, some nanostructured Bi catalysts with different morphology have been explored, for example, nanodendrites, nanosheets, and monolayers to complement the optimization of surface structures, wavy SnO₂ catalyst showed maximum FE formate of 87.4% and EE formate of 57.5% at −1.0 V versus RHE.

III) Metallic In has a relatively higher cost than that of Sn and Bi, therefore attracting limited attention for large-scale applications. However, as an environmental-friendly metal, indium is one of the earliest investigated main group metals for formate formation toward ECR in a similar way as Sn and Bi. Bulk In reached a FE formate value of ≈95% at −1.55 V versus RHE in the 1990s,[28] unfortunately the required overpotential (1.36 V) and current density (5 mA cm⁻²) were unsatisfactory. To optimize the activity of In-based catalysts, dendritic In foams have been recently prepared by an electrodeposition method in an aqueous solution with the existence of Cl⁻ ions when using the hydrogen bubble dynamic templates.[29] The as-obtained In electrode possesses a large electrochemical surface area (ECSA) and needle-like dendrite nanostructures, showing an improved FE formate value of 86% at a relatively low potential of −0.86 V versus RHE. Since an oxide layer is usually inevitable on the surface of In under ambient conditions, In₂O₃ has also been regarded as the active species that benefit the CO₂ adsorption and intermediate binding capacity in ECR. Meanwhile, the close interactions between different components in a hybrid material offer synergistic effects for enhanced CO₂ electroreduction activity. Based on these proposed strategies, Zeng’s group synthesized a hybrid catalyst that is composed of porous In₂O₃ nanobelts and reduced graphene oxide (In₂O₃-rGO) by a facile two-step process.[30] The theoretical calculations and electrochemical microkinetic analysis demonstrated that the presence of chemical coupling within the In₂O₃-rGO hybrid caused the changes in electronic density, contributing to the higher extent of electron transfer on the catalyst surface. Accordingly, the electron-rich structure of In₂O₃-rGO significantly affected the adsorption ability of reaction intermediates, therefore facilitating the formation of key intermediate *HCOO⁻* during CO₂-to-formate conversion. The aforementioned results clarified the experimental observations that In₂O₃-rGO hybrid exhibited enhanced FE formate (1.4-fold) and current density (3.6-fold) than that of bare In₂O₃ nanobelts (Figure 7a–c). Similarly, Mou and co-workers reported an In₂O₃@C catalyst,[10] which showed the peak FE formate value of 87.6% and satisfactory durability (12 h) at
Figure 7. a) Gibbs free energy diagrams for CO$_2$ reduction to formate on In$_2$O$_3$-rGO hybrid, In$_2$O$_3$/rGO, and In$_2$O$_3$ catalysts. b) FE$_{CO}$, FE$_{formate}$, and c) FE$_{formate}$ for In$_2$O$_3$-rGO hybrid, In$_2$O$_3$/rGO, and In$_2$O$_3$ catalysts. The color codes in (c) apply to (b). Reproduced with permission. Copyright 2019, American Chemical Society. d) TEM image of Sb nanosheets. e) FE$_{formate}$ values of bulk Sb, Sb nanosheets, and Sb nanosheets-graphene composite catalysts at different potentials. Reproduced with permission. Copyright 2017, Wiley-VCH. f) TEM image and the size distribution histogram of Pd-B/C catalyst. g) Potential dependent FE$_{formate}$ of Pd-B/C, Pd/C (home), and Pd/C (com) catalysts. Reproduced with permission. Copyright 2019, John Wiley & Sons, Inc. h) TEM image of partially oxidized Co 4-atom-thick layers. i) FE$_{formate}$ at each applied potential and j) ECSA-corrected Tafel plots for formate production of partially oxidized Co 4-atom-thick layers (red), Co 4-atom-thick layers (blue), partially oxidized bulk Co (violet), and bulk Co (black). Reproduced with permission. Copyright 2016, Springer Nature Limited.

an overpotential of only 710 mV. These performances were significantly superior to that of In$_2$O$_3$ nanoclusters due to the increased ECSA and the positive effects of carbon black.

IV) Pb-based catalysts perform promising activity and selectivity for formate formation in ECR, whereas its high toxicity brings adverse impacts on human health and the environment, and thus hindering their practical applications. Kanan’s group prepared PbO$_2$-derived nanocrystalline Pb films by in situ electrochemical reductions, which significantly suppressed HER and reached FE$_{formate}$ values above 90% from −0.7 to −1.0 V versus RHE. Importantly, the metastable Pb oxide (hydroxide) species covered on Pb surface favor the ECR process and act as passivation layers for HER. To simultaneously realize a higher current density with an optimal FE$_{formate}$, arylaliphatic amines, aminophenyl, and nitrophenyl derived Pb catalysts were synthesized. The enhanced CO$_2$-to-formate performance was due to the effectively improved CO$_2$ capture, suppressed H$_2$ generation, and lowered overpotential. Accordingly, the amine-derived Pb electrocatalysts showed FE$_{formate}$ of 94% and a partial current density for formate production ($j_{formate}$) of 9.5 mA·cm$^{-2}$ at an applied potential of −1.09 V versus RHE. To further fabricate an efficient catalyst with enhanced electrical conductivity, and larger specific surface area, a multi-walled carbon nanotube aerogel supported Pb catalyst was synthesized, which could provide a rapid mass transfer pathway for reactants and lead to sufficient contact between CO$_2$ molecules and active sites. Finally, the as-obtained electrocatalyst performed high FE$_{formate}$ (84.6%) and large current density (28 mA cm$^{-2}$) with long-term stability (10 h).

V) Sb is closed to metallic Sn and Bi in the periodic table, while it remains unexploited for ECR. This is probably due to the limited active sites on the surface of bulk Sb, which have a strong impact on its practical application in the catalytic field. Interestingly, the pristine Sb material possesses a rhombohedral layered structure, which can be easily exfoliated into 2D form (namely, antimonene). Due to its satisfactory stability and specific features, researchers have preliminarily tried to study its physiochemical properties and explored the catalytic performance. Li and co-workers successfully synthesized 2D Sb nanosheets with few layers by cathodic exfoliation, which exposed a higher density of active edge sites for CO$_2$ reduction (Figure 7d). The Sb nanosheets realized a peak value of FE$_{formate}$ (≈84%) at a moderate overpotential of 0.97 V. To further enhance...
FE\textsubscript{formate}, and current density of Sb nanosheets, anodically exfoliated graphene nanosheets were added to form an Sb nanosheets-graphene composite, which exhibited a higher FE\textsubscript{formate} value of 88.5% and a \textit{I\textsubscript{formate}} above 8 mA cm\textsuperscript{-2} at a decreased overpotential of 0.87 V for a long-term testing (>12 h). The improved CO\textsubscript{2}-to-formate behavior mainly resulted from the abundant active sites on 2D Sb nanosheets and strong interactions between Sb and graphene (Figure 7e).

VI) Pd-based catalysts have also been applied in electrocatalytic fields, performing satisfactory selectivity of C\textsubscript{1} compounds production in ECR.\cite{36} Detailed mechanisms have been studied to analyze the keys that dominate the activity and selectivity of catalysts. The optimized morphology and composition of nanostructured Pd catalysts, as well as, the in situ formed active phases (e.g., Pd-hydride) caused by applied potentials resulted in various selectivities toward reduction products via distinct reaction pathways and intermediates. Moreover, the introduction of heteroatoms (e.g., N, B) and supports (e.g., metal oxides, carbons) can improve the activity and selectivity of Pd-based catalysts and further maintain good durability for long-term ECR. Sargent’s group prepared different morphologies of Pd nanoparticles to expose high-density high-index surfaces,\cite{37} which optimized the binding capability of intermediates and thus realized preferable activity and selectivity of CO\textsubscript{2}-to-liquid fuels (mainly, formate) conversion. The as-obtained Pd nanoparticles exhibited a high FE\textsubscript{formate} value of 97% with a record current density of 22 mA cm\textsuperscript{-2} at a relatively low overpotential of −0.2 V. The abundant higher-index facets exposed on the Pd surface provided more step sites and undercoordinated atoms, which effectively decreased the energy barrier of reaction intermediates (HCOO\textsuperscript{−}) formation favoring formate production. By altering active phases of Pd nanoparticles under different applied potentials with the existence of \textsuperscript{1}H species, Gao et al. found that different products would be generated via a specific reaction pathway and preferable intermediates through advanced analysis of in situ X-ray absorption spectroscopy, in situ attenuated total reflection-infrared spectroscopy, and DFT results.\cite{38} The Pd catalyst contains a core–shell structure (α+β PdH\textsubscript{x}@PdH\textsubscript{y}) above −0.2 V versus RHE, which mainly generate formate via the HCOO\textsuperscript{−} intermediate, whereas the in situ formed β-PdH\textsubscript{x}@Pd below −0.5 V versus RHE prefers CO production via the COOH\textsuperscript{−} intermediate. Jiang and co-workers reported a boron-doped Pd catalyst (Pd-B/C),\cite{39} which effectively suppressed the CO poisoning phenomenon and selectively facilitated CO\textsubscript{2}-to-formate conversion. During ECR, Pd-B alloy would form once B atoms are inserted in the Pd lattice, and favors the formation of HCOO\textsuperscript{−} intermediates. Accordingly, the FE\textsubscript{formate} value achieves ≈70% over 2 h electrolysis at a low voltage of −0.5 V versus RHE on the Pd-B/C catalyst (Figure 7f,g). To broaden the potential window of Pd catalyst toward ECR, N, B-codoped TiO\textsubscript{2} nanotubes were fabricated and employed as a support to optimize the activity and selectivity of metallic Pd in CO\textsubscript{2} reduction.\cite{40} The dopant of N and B elements accompanied with the interactions between Pd and TiO\textsubscript{2} support synergistically improve the electronic properties of the composite catalyst, which effectively stabilizes the active Pd species (Pd hydride) and suppresses the CO formation pathway, and thus favoring the formate production in the larger potential window than the undoped electrodes. The concurrent CO poisoning phenomenon severely affects the CO\textsubscript{2}-to-formate conversion of Pd-based catalyst, which subsequently results in dehydration, lower formate selectivity, and decreased stability to a great extent. In this context, Lee and co-workers proposed a system-level strategy to effectively resolve the poisoning issue via a cyclic two-step electrolytic process.\cite{41} The researchers alternately applied the reduction and oxidation potentials during the ECR process to selectively remove CO from the surface of Pd during the anodic step, based on the different reversibility (redox potential) of formate and CO production reactions. Therefore, the cyclic two-step electrolysis maintains 100% current density durability and 97.8% selectivity toward formate production over 45 h testing, significantly superior to other conventional potentiostatic electrolysis.

VII) Other metallic catalysts like Co- and Mo-based catalysts are also regarded as the ideal alternatives for ECR due to their earth abundance and low overpotential/free energy for \textsuperscript{1}H species adsorption. Surface engineering has been explored to design highly efficient catalysts with low dimensions and large specific surface areas, which can modify the \textit{d}-band electronic structure of metal atoms, and thus construct suitable active sites for CO\textsubscript{2} reduction. Xie’s group successfully prepared atomic Co layers with high selectivity and long-term durability under relatively low overpotential for formate production.\cite{42} The optimal ultrathin morphology and oxidation state turned an inferior catalyst into an active one for CO\textsubscript{2} electroreduction, which facilitates the rate-determining proton transfer step via the strong stabilization of HCOO\textsuperscript{−} intermediate (Figure 7h–j). The Mo catalyst was optimized into a single-atomic scale to improve the atom utilization and charge transferability, which significantly enhances the catalytic activity and formate product selectivity with abundant active sites exposed.\cite{43}

2.2.2. CO\textsubscript{2} to CO

As important feedback of syngas, CO can be used in FT synthesis to further obtain a series of organic chemicals and intermediate products, such as, MeOH, gasoline, and diesel, etc. However, CO as a traditional CO\textsubscript{2} hydrogenation product has been commonly generated from the RWGS reaction, which is an endothermic reaction activated at relatively high temperatures (>300 °C). Accordingly, converting CO\textsubscript{2} to CO in an electrochemical manner is a promising alternative under ambient conditions, and CO can be easily extracted from the aqueous electrolytes for further applications, including chemicals, medicine, and the metallurgical industry.\cite{44} The electrocatalytic CO\textsubscript{2}-to-CO conversion undergoes a 2-proton/electron transfer process which involves two elementary steps: CO\textsubscript{2} molecule is first reduced and then hydrogenated to generate the surface-adsorbed \textsuperscript{0}COOH intermediate during a proton-coupled electron transfer process.\cite{45} Afterward, the \textsuperscript{0}COOH intermediate goes through the second proton/electron transfer process to convert into \textsuperscript{0}CO intermediate.
and then is desorbed from the catalyst surface to obtain the final product, gaseous CO. Over the past 5 years, a great deal of work has sprung up to uncover highly active and selective electrocatalysts for CO₂-to-CO conversion, such as metallic nanoparticles (Au, Ag, Pd, Cu, and Zn), single atoms (Fe, Co, and Ni), and carbon-based materials.

![Figure 8.](image)

**Figure 8.** a) CO product selectivity versus applied potential of bare Au, CN-Au, and Cl-Au. b) The schematic illustration of van der Waals interactions and electronic effects on modified Au. c) Correlation between binding energies of reaction intermediates and d-band centers up to 3/9 ML Cl and CN. Reproduced with permission. Copyright 2018, American Chemical Society. d) The three-phase interface of the Au/H₂O/CO₂. e) SEM image of Au/nanoPE membrane. f) FE CO (red curve) and FE H₂ production (black curve) by bilayer Au/nanoPE. Reproduced with permission. Copyright 2018, Springer Nature Limited. g) Top-down SEM image of the Ag foam with the primary macroporosity. h) Schematic illustration of the proposed reaction pathways toward CO₂-to-CO conversion on Au foam. Reproduced with permission. Copyright 2018, American Chemical Society.

I) Au-based materials are commonly regarded as the pioneer for CO₂ electroreduction with high selectivity of CO production (FE CO). However, the bulk morphology suppresses the activity of Au electrodes. Accordingly, the enhanced kinetics of the ECR in terms of nano-Au was studied by Chen’s group based on the in situ FTIR analysis, which directly detected the *COO⁻ and *COOH intermediates on Au surfaces. Based on the collected results, the possible reaction process of the CO₂ reduction was described in detail. Meanwhile, some effective strategies have further been employed to focus on the optimization of nanostructural Au materials, such as surface engineering, alloying, and supported Au composites, aiming at improving the catalytic CO₂-to-CO performance. The surface modification of catalysts is an effective strategy to enhance their catalytic activity due to the introduction of abundant active sites and the increased specific areas. Cho et al. modified the Au surface with additives by electroplating Au electrodes in an aqueous electrolyte with CN⁻ and Cl⁻ anions. Evidenced by the experimental and theoretical results, the addition of CN⁻ and Cl⁻ anions brings about van der Waals interactions and electronic effects, which significantly enhances the CO selectivity (FE CO = 80% at −0.39 V vs RHE) when compared to pristine Au (FE CO < 20%) (Figure 8a). The van der Waals and electronic contributions are responsible for physical interactions (mainly stabilization effect) among adsorbates and chemical absorption abilities onto the metal surfaces, respectively, which promotes the stabilization of *COOH intermediate and weakens the absorption of *H species (Figure 8b,c). In addition to the anion additives, amines and anchoring agent (thiol-tethered ligands) have also been reported as effective modifiers to optimize product selectivities in ECR, which play similar roles in the interactions between absorbates and Au electrodes with the existence of abundant low-coordinated sites. The construction of nanostructured surface could also bring about different coordination environments for surface atoms, which effectively adjusts their ECR performance. Narayanaru and coworkers modified the electrode roughness by changing the interfacial pH gradients, which converted the flat surface into nano and hierarchical structures. Such microheterogeneity of surfaces contributes to tunable product selectivity and high catalytic activity in virtue of the enriched active sites of Au. Alloying Au with other metals shifts the d-band centers of Au, which changes their electronic structures and alters the interactions between intermediates and substrates. Meanwhile, alloying can also bring about ge-
omeric effects, which leads to the atomic rearrangement around the active sites and further influences the absorption capabilities of reaction intermediates onto the alloy surface. Accordingly, Au-Pd alloys synthesized by Valenti et al. [51] and Au-Pt alloys fabricated by Ma et al. [52] showed distinct catalytic activity and selectivity for CO$_{2}$-to-CO conversion by varied binding strength of °COOH and °CO intermediates, which adequately demonstrated the synergistic effects of the changed electronic and geometric structures. This study would inspire more works on the synthesis of highly active and selective electrocatalysts at reduced overpotentials. Lately, some works focused on the support effect of Au catalysts, which showed great influences on the stabilization and dispersion of nanostructured Au, and provided abundant active sites and effective synergetic interactions between Au and support for enhanced ECR performance. Cui’s group reported a bilayer Au/nanoporous polyethylene (nanoPE) membrane via a sputtering method, [53] which exhibits an optimal FE$_{CO}$ of ≈92% with a high geometric current density for CO production (j$_{CO}$) of 25.5 mA cm$^{-2}$ at −0.6 V versus RHE. The nanoPE membrane not only promotes the CO$_{2}$ mass transport toward the reaction sites but also provides a high density of active sites at the three-phase (Au-H$_{2}$O-CO$_{2}$) interface due to its hydrophobic property (Figure 8d–f). Jin et al. reported an Au nanocatalyst supported on the N-doped carbon (AuNCs@CN), [54] where N heteroatom enriched the surface charge density and realized a high degree of dispersion of Au. When compared to pristine Au, N-doped carbon significantly increases the localized concentration of CO$_{2}$ around Au active sites, which improves its selectivity toward ECR by 50% and further decreases the total cost of the catalytic system.

II) Ag-based materials have been regarded as an ideal alternative for ECR owing to their abundant storage, low cost, and satisfying FE$_{CO}$ compared to that of Au. Recently, numerous works have been reported to concentrate on the structure optimization of Ag with low overpotential and high FE$_{CO}$ over 90%. Dutta et al. developed Ag-foam catalysts via a metal deposition approach with masses of low-coordination reaction sites, [55] which showed superb FE$_{CO}$ above 90% over a broad potential window from −0.3 to −1.2 V versus RHE (Figure 8g,h). Meanwhile, it presented long-term durability with FE$_{CO}$ retaining above 90% during 70 h ECR at −0.8 V versus RHE. Similarly, the sponge-like AgCu alloy with 3D nanoporosity possessed numerous holes and interior void space, [56] which led to larger ECSA and highly active local sites, thus resulting in the improved CO$_{2}$ adsorption, electron-proton transfer, °COOH binding capability, and °CO desorption. Other strategies have also been adopted to effectively stabilize reactants and reaction intermediates except for the structural design. For example, with the introduction of the amine functional group, Ag nanoparticles exhibited an improved selectivity for CO production (FE$_{CO}$ = 94.2%) and effectively suppressed HER owing to its destabilization of the bonded hydrogen species. [57] In addition, oxide-derived Ag catalysts were reported to be promising candidates for high-performance ECR. Ma et al. demonstrated an Ag catalyst derived from Ag oxides, which exhibited a decreased overpotential by more than 400 mV accompanied with a higher FE$_{CO}$ (≈80%) than that of untreated Ag (FE$_{CO}$ ≈ 4%) at a moderate overpotential of 0.49 V versus RHE. [58] The idea was applied to another work on Ag nanoparticles, [59] which incorporated stable oxygen species onto the surfaces and led to an improved CO selectivity and long-term stability. O$_{2}$ plasma-treated Ag foil was also employed to construct rougher surfaces and new active sites for Ag-based catalysts, [60] which performed optimized overpotentials for CO$_{2}$-to-CO conversion due to the stronger binding ability of reaction intermediates, mainly °COOH and °CO species. To further enhance the catalyst stability, SnO$_{2}$ species were electrodeposited on the O$_{2}$ plasma-treated Ag foil to construct a bimetallic system, which effectively maintained the chemical state, surface morphology, and composition of the catalysts. Similar phenomena were further evidenced by a recent study on a highly active Ag-alloyed Zn dendritic electrocatalyst, [61] which exhibited a high CO$_{2}$-to-CO selectivity of 91% and stability during 40 h testing without any obvious loss.

III) Pd-based materials are promising catalysts for CO and formate production in ECR, which are greatly influenced by the applied potentials, [62] morphologies, active sites, crystal facets, and heteroatoms doping. Recently, Chen and coworkers reported nanosized Pd octahedra with dominant Pd(111) facet, [62] which showed a high CO selectivity of 95% since the binding energies of reaction intermediates (°CO and °HOCO species) were optimized on the surface of in situ formed PdH(111) under reaction conditions (Figure 9a–c). Gong’s group synthesized 5-atomic-layer hexagonal Pd nanosheets with numerous exposed edge sites, [63] which reached a FE$_{CO}$ value of 94% at −0.5 V versus RHE without any activity loss in 8 h electrolysis. Meanwhile, the DFT calculations revealed that the increased amount of edge sites exposed more active centers with a generalized coordination number of 5, which certainly improved the catalytic performance due to the easier °COOH formation and °CO desorption. Another work was reported by the same group, [64] in which the Pd concave cubes enclosed with high-index (310) facets showed the peak FE$_{CO}$ value of 90.6%. Sun et al. engineered the surface of Pd nanoparticles by the introduction of CeO$_{2}$ interfaces and Pd heteroatoms, [65] which could reach a FE$_{CO}$ value over 84% and a high mass activity for CO formation of 92 mA m$^{-2}$.

IV) Cu-based materials are well-known catalysts for high-value-added hydrocarbons production instead of CO in ECR due to their relatively high CO binding capacity. However, many efforts have also been made to adjust the absorption ability of reaction intermediates, which altered the main product from hydrocarbons to CO. To enlarge the number of active sites and atomic utilization, ultrathin Cu nanosheets [66] and Cu atom-pair catalysts [67] were synthesized, which promoted the mass and electron transfer, the interaction between reactants/intermediates and Cu substrates, and intrinsic activity, thus generating optimized efficiencies, selectivities, and stability with relatively low overpotentials for CO production. In addition, some works focused on combining Cu with other metals or metal oxides to break the scaling relations and contribute to highly selective CO for-
Figure 9. a) TEM image for Pd octahedra. b) Faradaic efficiencies of reduction products for Pd octahedra in ECR. c) DFT calculated free energy diagrams of ECR on Pd(111), Pd(100), PdH(111), and PdH(100). Reproduced with permission.[62] Copyright 2019, Wiley-VCH. d) FE-SEM image of hexagonal Zn. e) FE CO of Zn foil and hexagonal Zn at various constant potentials ranging from $-0.6$ to $-1.1$ V versus RHE. f) CO$_2$ reduction pathway on Zn (002) (black solid line) or Zn (101) (red solid line) at $-0.71$ V versus RHE. Reproduced with permission.[75] Copyright 2016, Wiley-VCH.

formation at a decreased onset potential, which was caused by the balance between *COOH and *CO binding energies on the catalyst surface. Furthermore, CuAu,[68] CuPd,[69] CuZn,[70] CuCu,[71] CuFe,[72] In-doped Cu@Cu$_2$O,[73] and Cu/SnO$_x$[74] had been studied to shift the main products to CO.

V) Zn-based materials have also been popular alternatives for ECR due to their abundant storage, low toxicity, and high CO selectivity. Woo’s group reported a hierarchical hexagonal Zn catalyst,[75] which realized a highly efficient (FE$_{\text{CO, max}} = 95\%$) and stable performance (30 h) for CO production. The optimized (101)/(002) facet ratio effectively stabilized *COOH intermediate and suppressed HER process (Figure 9d–f). However, Zn is regarded as an active metal that can easily form surface oxide layers, which on the other hand minimizes the density of active sites and increases HER activity. Inspired by a promising “Trash to Treasure” approach, a facile anodic oxidizing method was applied to regulate ZnO species with lattice dislocation,[76] which successfully converted CO$_2$ to syngas (CO + H$_2$) with satisfactory ratios in a wide potential window. Besides, strategies of converting ZnO to porous Zn catalysts via a pre-treatment process[77] or in situ reduction[78] during the ECR process, led to an optimal stabilization capacity of the key intermediates toward CO$_2$-to-CO conversion and therefore contributing to high FE$_{\text{CO}}$ and $J_{\text{CO}}$ values.

VI) Carbon-based materials have been considered as highly active and stable electrocatalysts owing to their rapid ion and electron transfer capacity, as well as, chemical and thermal stability. Tremendous efforts have been made to study their applications in HER, oxygen reduction reaction, and OER, whereas they perform notorious activity toward ECR possibly due to the lack of active sites for CO$_2$ activation. Accordingly, introducing heteroatoms into the carbon framework has been demonstrated to be an effective approach to enhance CO$_2$ reduction activity by tuning the electronic properties of adjacent carbon atoms. Some carbon-based materials, such as, N-doped carbon nanotube, graphene, carbon fibers, carbon black, nanoporous carbon, and diamonds had been summarized in our previous review,[7] which presented high CO selectivities and low overpotentials in ECR. Besides, Guo and co-workers revealed the intrinsic activity of ECR on N-doped carbon materials including graphene and CNT, through DFT and ab initio molecular dynamic calculations.[79] Theoretical results calculated the barriers for CO$_2$ activation and intermediates formation on different structures of carbon materials to verify the active sites and the selectivities of different products, which revealed the vital role of N species in activating CO$_2$ reduction and restraining HER kinetics. Since the pyrolyzation of zeolitic imidazolate frameworks (ZIFs) has been known as a strategy to provide N-doped carbons, Lin’s group synthesized a composite material combining the pyrolyzed ZIFs with multi-walled CNTs, which reached an extraordinary FE$_{\text{CO}}$ value of approximately $100\%$ at a moderate overpotential of 740 mV.[80] Meanwhile, Lin’s group further introduced the Fe element into the pyrolyzed ZIFs/multi-walled CNTs hybrid and achieved a high FE$_{\text{CO}}$ value of $97\%$ at a decreased overpotential of only 440 mV. Similarly, Xu and co-workers prepared a 3D N-doped mesoporous carbon-supported Ni catalyst, which exhibited excellent FE$_{\text{CO}}$ of $\approx 98\%$ at $-0.7$ V versus RHE and long-term durability during the 25 h test.[81] The enhanced performance of carbon-based materials containing N species and transition metals (M-N-C) could be explained by DFT studies, which were reported in a recent review of Strasser’s group.[82] When com-
Figure 10. a) High magnification aberration-corrected HAADF-STEM image of atomic Fe/N-doped graphene catalyst. b) Potential-dependent $F_{\text{ECO}}$ in ECR on Fe/N-doped graphene single-atom catalysts prepared at different annealing temperatures. c) Free energy diagram for ECR to CO on Fe-N$_4$ moieties supported on graphene sheets. Reproduced with permission.[83] Copyright 2018, Wiley-VCH. d) AC-HAADF-STEM images of Co-N$_5$/HNPCs catalyst at different magnifications. e) The comparisons of $F_{\text{ECO}}$ values of Co-N$_5$/HNPCs and CoPc catalysts at various constant potentials. f) Calculated free energies of CO$_2$RR on Co-N$_5$/HNPCs and other metal phthalocyanines (FePc, CoPc, NiPc, and CuPc). Reproduced with permission.[86] Copyright 2018, American Chemical Society. g) $F_{\text{ECO}}$ values of NiPc-OMe MDE, NiPc MDE, NiPc-CN MDE, and P-NiNC at different applied potentials. h) Calculated free energy diagrams of NiPc-CN-H$_2$, NiPc-H$_2$, and NiPc-OMe-H$_2$ for CO$_2$RR at $-0.11$ V versus RHE. Reproduced with permission.[92] Copyright 2020, Springer Nature Limited.

Compared with N-doped carbon, the incorporation of transition metals could further enhance the reactant mass transport, electron-proton mobility, and electronic density of the carbon framework. The doping of transition metals also played an essential role in the interplay between key intermediates and metal sites, which significantly influences the ECR performance. In particular, to realize high CO selectivity, the binding energy of *H intermediate on the metal center should be reduced to suppress the HER side reaction. Meanwhile, to further decrease the overpotential of CO$_2$ to CO conversion, the absorption capability of *COOH and *CO intermediates should be optimized on metal sites.

VII) Currently, some inexpensive transition metals (e.g., Fe, Co, and Ni) with bulk morphology perform superior HER and easy CO-poisoning phenomenon, which suppress their CO$_2$ reduction to a great extent. The limited densities of active sites and poor intrinsic activity of bulk electrocatalysts hinder their ECR performance. Accordingly, atomically dispersed transition metal catalysts have recently been the popular options for CO$_2$ to CO conversion due to their high atomic utilization, tunable coordination structures, and suitable electronic effects. Many studies have sprung up to explore transition metal single atoms or porphyrin-like catalysts in ECR, with a special focus on the synthetic methods and their structure-performance relationship. N-doped carbon frameworks have been regarded as promising supports to achieve the atomic dispersion of non-noble metals. For example, graphene can be used as suitable support to stabilize these atomic active sites because of its specific surface area, high electronic conductivity, and controllable surface engineering. Meanwhile, heteroatom doping can further adjust the electronic structures of graphene, which promotes the strong interaction between heteroatoms and metals and thus realizes the high dispersion of metal atoms onto the graphene.

Tour et al. anchored atomic Fe sites on the N-doped graphene support, which exhibited an optimal $F_{\text{ECO}}$ value of 80% at a low overpotential (Figure 10a,b).[83] DFT calculations were further conducted to understand the mechanism of enhanced CO$_2$ to CO conversion with the existence of Fe-N$_4$ moieties. The results indicated that the Fe-N$_4$ centers effectively lowered the formation energy of *COOH intermediates and desorption ability of *CO intermediates, thus favoring the CO production.
Besides, N-doped graphene support could also stabilize atomically dispersed Fe species to form the Fe-N\textsubscript{s} active sites, which formed from an additional axial pyrrolic N ligand coordinated to Fe-N\textsubscript{b} moieties.[84] The DFT results revealed that Fe-N\textsubscript{s} sites could make full use of the electron density of Fe 3d orbitals, which suppressed the Fe-CO π back-donation effect and thus accelerated CO desorption. Herein, the Fe-N-C catalyst exhibited an extraordinary CO selectivity (FE\textsubscript{CO} = 97%) at a low overpotential of 350 mV. To further improve the ECR performance of the Fe-N-C catalyst, Li’s group incorporated S atoms in N-doped microporous carbon layers, which effectively tuned the electronic structure Fe-N active centers.[85] The N, S co-doping upshifted the Fermi energy of Fe 3d and increased charge density of Fe atoms on Fe-N\textsubscript{s} sites, which facilitated CO\textsubscript{2} activation and strengthened the interaction between active sites and key intermediates, and thereby achieved a high FE\textsubscript{CO} of 98% at a moderate overpotential of 490 mV and long-term stability during a 30 h test without obvious activity degradation.

Atomically dispersed Co has also been a promising active site for highly effective and selective ECR. Li et al. synthesized Co single atoms supported on hollow N-doped porous carbon spheres (HNPCSS), which were coordinated with the adjacent N atoms to form the Co-N\textsubscript{s} site (Figure 10d).[86] The Co-N\textsubscript{s}/HNPCSS catalyst exhibited high selectivity toward CO\textsubscript{2}-to-CO conversion with FE\textsubscript{CO} above 90% over a wide potential window of −0.57 to −0.88 V versus RHE (Figure 10e). DFT studies revealed that the extraordinary performance was mainly ascribed to the atomically dispersed Co-N\textsubscript{s} sites, which effectively facilitated CO\textsubscript{2} activation, *COOH intermediate formation, and *CO desorption (Figure 10f). Co-N\textsubscript{s} moieties within Co porphyrin or supported over carbon substrates have also been reported for selective CO\textsubscript{2}RR. The mechanism and reaction pathways of these catalysts in ECR were comprehensively investigated, aimed at gaining new insights into the efficient catalytic systems. Koper and co-workers reported a DFT study of Co porphyrins in ECR.[87] They found that the CO\textsubscript{2} molecule was first activated into CO\textsuperscript{−} anions and bound tightly onto a Co porphyrin. Then, CO\textsuperscript{−} acted as a Bronsted base and obtained a proton from a water molecule, and then formed the [Co(P)(COOH)] intermediate. Afterward, gaseous CO was generated as the main product through a decoupled proton-electron transfer process. Accordingly, Co porphyrins can realize high faradaic yield, turnover frequency, and low overpotential when catalyzing CO\textsubscript{2} to CO. For example, a metallloporphyrin-tetrathiafulvalene catalyst-based covalent organic framework was prepared with Co porphyrin as building units,[88] which exhibited a high FE\textsubscript{CO} value of 91.3% at −0.7 V versus RHE and long-term stability over 40 h in aqueous electrolyte. To further enhance the CO\textsubscript{2} reduction performance of Co porphyrin, the catalyst could be incorporated into various organic frameworks owing to its structural flexibility. Sun et al. synthesized Co porphyrin nanotubes with a large diameter to study the curvature effect on the CO\textsubscript{2}RR selectivity, which preferred CO formation at a low overpotential because of its weak adsorption of *CO intermediate.[89] To improve the conductivity and stability of the overall catalytic system, carbon-based supports have been applied to immobilize the Co porphyrin forming strong π–π interactions between the porphyrins and carbon supports. Daasbjerg’s group explored a facile method to directly immobilize unmodified cobalt meso-tetraphenylporphyrin (CoTPP) onto CNT,[90] which showed a high selectivity of CO production above 90% at a relatively low overpotential. In comparison, the pristine unsupported CoTPP catalyst exhibited poor product selectivity and slow reaction rate of CO\textsubscript{2} conversion and required an excess high overpotential. Similarly, Han et al. covalently grafted Co porphyrins onto the CNT surface by a substitution reaction with a quite high loading of 10 wt%,[91] which achieved an extraordinary FE\textsubscript{CO} of 98.3% and a high current density of 25.1 mA cm\textsuperscript{−}2 with long-term durability at a low overpotential of 490 mV. In addition, the turnover frequency of CO production was improved by more than three times higher when compared to the physically mixed Co porphyrin/CNTs composites. The optimized configuration strengthened the catalyst-substrate interaction, which synergistically improved the electron transfer to the intermediates and the long-term stability of the overall catalytic system.

Ni phthalocyanines were also incorporated onto CNTs as an efficient composite for the CO\textsubscript{2}RR application. Liang et al. synthesized Ni phthalocyanine with -OME/-H ligands embedded on carbon nanotubes as a molecularly dispersed electrocatalyst (NiPc-OME MDE), which achieved superior catalytic performance in terms of activity (high current density = 300 mA cm\textsuperscript{−}2), selectivity (FE\textsubscript{CO} > 99.5%), and stability (40 h) (Figure 10g).[92] Observed from DFT results (Figure 10h), the energy barrier for *COOH formation is smallest in the case of NiPc-OME, which is the rate-determining step toward CO\textsubscript{2}-to-CO conversion, thus facilitating the activity and selectivity of CO production. Moreover, analyzed from Ni–N bond orders in NiPc molecules, the results indicated that NiPc-OME possessed the strongest Ni–N bonds, and the existence of methoxy ligand preferred easier CO desorption, which synergistically avoided structure collapse of catalyst and made NiPc-OME MDE an ultra-stable CO\textsubscript{2}RR catalyst. Liu and co-workers synthesized atomically dispersed Ni-N\textsubscript{s} active sites,[93] which anchored the planar Ni-N\textsubscript{s} onto the N atom in the carbon matrix to form NiPc/NC catalyst. Owing to the synergistic effect of the coordinatively unsaturated Ni-N\textsubscript{s} sites and the surface pyridinic N species, NiPc/NC exhibited high FE\textsubscript{CO} over 93% in a wide potential window from −0.5 to −0.8 V versus RHE, and the optimal value can up to 98% at a relatively low potential of −0.5 V versus RHE. Similarly, Ni and N co-doped mesoporous carbon materials were fabricated by Qiu’s group,[94] which brought synergetic effects to enhanced CO\textsubscript{2} activation, accelerated reactant, and electron transfer, as well as, a higher density of active sites in the form of Ni–N moieties in the unique ordered mesoporous structure and therefore realizing high FE\textsubscript{CO} over 90% in a broad potential range from −0.75 to −1.20 V versus RHE. Lou et al. successfully decorated isolated Ni atoms onto hollow N-rich carbon plates (Ni-NC),[95] which greatly facilitated the electron transfer and boosted the redox capability for CO\textsubscript{2} reduction. The Ni atoms acted as the active sites that could adjust the energy configuration of Ni-NC catalyst, which achieved highly-efficient electrochemical CO\textsubscript{2}RR with enhanced current densities and high selectivity of nearly 100% toward CO formation in a wide potential range. Li et al. conducted a solid-state diffusion process between the N-doped carbon and bulk Ni metal to synthesize Ni single-atom catalysts on self-supported and hierarchical carbon substrate.[96] The resultant catalyst exhibited a high current density of 48.66 mA cm\textsuperscript{−}2 and a high FE\textsubscript{CO} of 97% at
–1.0 V versus RHE, reaching an industrial-scale level. The aforementioned works broaden the applications of non-noble materials into scalable, efficient, and stable catalysts.

2.2.3. CO₂ to Hydrocarbons/Oxygenates

The technologies of CO₂-to-formate and CO₂-to-CO conversion are comparatively mature in ECR, which can be directly used in the chemical industries. However, the synthesis of other hydrocarbons and oxygenates requires overcoming higher energy barriers for CO₂ adsorption, activation, and the subsequent step-wise transformations of *CO intermediates, which determine the yield of C₂₄ products of ECR, as illustrated in Figure 4. The *CO dimerization is the pre-requisite step for C₂H₄, CH₃CH₂OH, and n-C₃H₇OH production. After the successive proton-electron pair transfer process, the CH₃CHO intermediate is generated from the *CO-CO dimerization, which is a rate-determining step for C₂H₄ and C₂H₅OH production. Furthermore, n-C₃H₇OH can be produced by inserting CO into the stabilized *CH₃CHO intermediate. C₃H₄ and CH₃COO– share the same *CH₂ intermediate, which is generated from a series of hydrogenation reduction steps of *CO intermediate. Subsequently, the insertion of *CO into *CH₂ generates CH₃COO– product, or the protonation of *CH₃ gives *CH₂ intermediate, which produces C₂H₄ via *CH₂ dimerization. Accordingly, the formation of multi-carbon compounds has been regarded as a complex and uncontrollable process, which is mainly attributed to the involvement of various intermediates and non-electrochemical steps. Hence, the fabrication of highly efficient electrocatalysts for hydrocarbon and oxygenate formation deserves more research attention. Various strategies (e.g., morphology and structure design, alloying/dealloying, surface modification, and support modification) have been employed to optimize the structure of the catalysts to achieve higher formation rates and selectivities toward desired hydrocarbons and oxygenates.

J) Cu has been regarded to be the most popular alternative for ECR that can effectively produce hydrocarbons and oxygenates. Although the bulk Cu shows limited activity for ECR, the morphology and composition regulations have been demonstrated to be effective strategies to improve its catalytic performance toward ECR to hydrocarbons and oxygenates, including nanostructuring (e.g., shape, exposed facet, crystal phase, defective site, oxidation state, and particle size), alloying (Cu-M bimetal), surface modification, and support addition.

**Influence of Nanostructuring on Activity:** Chen et al. have revealed shape-dependent activity of Cu for ECR, which demonstrated that cube-like and hexahexagonal decahedron-like Cu single crystals preferred C₂H₄ and C₂H₅OH production while octahedron-like Cu produced C₁ products. The different morphologies and structures determine the atomic arrangement of the Cu surface, which could alter the intermediate species and their binding energies and thus result in different reaction pathways and product distributions. Similarly, Wang’s group investigated the effect of Cu shape on the ECR activity and selectivity. When compared to Cu nanospheres with a similar size, Cu nanocubes performed an enhanced faradaic efficiency for C₂H₄ formation (FE_{C₂H₄} = 60%) and a partial current density of 144 mA cm⁻² by gas diffusion electrodes. The improved CO₂-to-C₂H₄ performance was mainly attributed to the exposed facets on the surface of Cu nanocubes and the alkalinity effects of electrolytes. At high overpotentials (~0.6 to ~0.8 V vs RHE), C–C couplings preferred to occur at (100) planes rather than (110) planes, which resulted in the higher C₁ selectivity on Cu nanocubes than that on Cu nanospheres. Tan et al. reported a hierarchically nanoporous Cu skeleton with large specific surface area, which not only provided the higher density of active sites but also accelerated the transport of the electron and reactant. Moreover, the decoration of vanadium oxide further facilitates the water dissociation and adjusts the *H adsorption energy, which lowers the energy barrier for the intermediate formation and the subsequent C–C coupling process, as evidenced by the DFT calculations. Therefore, the resultant np-Cu@VO₂ catalyst achieved a FE value of 30.1% for C₃H₇OH production with an ethanol partial current density of 16 mA cm⁻² at ~0.62 V versus RHE, which is a fourfold higher compared to the pristine nanoporous Cu.

The crystal phase of a metallic catalyst can also affect its ECR activity to a certain extent. Most of the reported Cu catalysts for CO₂ reduction are of face-centered cubic (fcc) crystal phase, which is the thermodynamically most stable phase of Cu. However, Zhang and co-workers recently reported unconventional 4H-Cu and hexaphase 4H/fcc Cu catalyst by using the 4H and 4H/fcc Au as templates, which exhibited enhanced activity for C₂H₄ formation than that of fcc Cu (Figure 11a,b). As evidenced by theoretical calculations, C–C coupling reactions between *CO and *CHO intermediates are more energetically favored on the surface of 4H Cu and 4H/fcc Cu and thus realize higher C₂H₄ selectivity (Figure 11c).

Nano-defective structure has also been constructed to facilitate the adsorption and accumulation of reaction intermediates onto the catalyst surface. Zhang et al. reported that the Cu nanosheets with abundant structure defects (nano-defective Cu NSs) could achieve high selectivity for C₂H₄ production (FE_{C₂H₄} = 83.2%) in ECR (Figure 11d,e). The study demonstrated that the nano-defective structure contributed to the enrichment of reaction intermediates and OH⁻ on the surface of the electrocatalyst, and therefore enhancing C₂H₄ formation via the C–C coupling reaction (Figure 11f). Twin boundary is another well-defined and stable defective structure, which could expose truly 1D and atomic string structure, thus effectively improving the activity of the twin catalysts. Meanwhile, the exposed atoms in the twin boundary have identical chemical environments, therefore providing a simple model for theoretical exploration. Sun’s group synthesized Cu twin boundaries on polished Cu electrode using a pulsed electrochemical deposition method. As evidenced by the intermediate experiments and DFT calculations, twin boundaries provided advantages for the stabilization of *COOH and *CHO intermediates, facilitating the conversion of absorbed *CO into CH₂. Herein, the Cu catalyst with twin boundary atoms exhibited a local partial current density of 1294 mA cm⁻² and an intrinsic FE_{C₂H₄} value of 92%.

Since Cu is an active metal and can be easily oxidized to Cu⁺ species on the surface during the electrocatalytic reaction, some works attempted to understand the relationship between the ECR activity and Cu⁺ species. For example, Huang and co-workers synthesized Cu₂O nanoparticles enclosed with different crystal facets, which synergistically reached a good preference for C–C couplings. The authors ascribed the good performance to...
the presence of abundant low-coordinate Cu\(^{2+}\) ions on the surface and thus boosting C\(_2\)H\(_4\) generation. Especially, Cu\(_{10}\)O nanoparticles with (111) and (100) facets exhibited an optimal FE of 59\%. Similarly, Sun’s group designed a new perovskite-type Cu\(_3\)N nanocube with Cu(I) active species for selective CO\(_2\)RR.\(^{[106]}\) The 25 nm Cu\(_3\)N nanocubes showed an optimal C\(_2\)H\(_4\) selectivity of 60\% and stability with a 20 h testing at \(-1.6\) V versus RHE. The experimental results and DFT calculations suggested that the (100) Cu(I) species stabilized by the Cu\(_3\)N structure favored CO-CHO coupling, which resulted in high selective for C\(_2\)H\(_4\) formation.

To design of Cu nano-catalysts with ultrasmall dimension has been regarded as another effective method to achieve improved catalytic activity and selectivity. To construct a unique electronic structure and maximize atom utilization, Cu-based single-atom catalysts have been reported for improved ECR activity. Although CO was the most common product for Cu catalysts, alcohols were also reported to be the possible products. For example, He and co-workers synthesized isolated Cu atoms onto the carbon nanofibers in a large-scale production method.\(^{[105]}\) The self-supported catalyst was directly used in CO\(_2\)RR, which realized high Faradaic efficiency for methanol formation of 44\%. Recently,
Xu et al. reported a highly-dispersed Cu single-atom catalyst using amalgamated Cu-Li as the precursor,[106] which could optimize the electronic state and the stereochemistry of the active centers. The catalyst realized a single-product faradaic efficiency of 91% at −0.7 V versus RHE toward CO₂-to-ethanol conversion (Figure 11g,h). Moreover, it showed a relatively low onset potential of −0.4 V versus RHE and satisfying stability over a 16 h electrocatalytic test. The superior faradaic efficiency for ethanol formation originated from the high dispersion of Cu atoms, which benefited the reactant activation and intermediate formation as suggested by first-principles calculations (Figure 11i).

Influence of Cu Alloying on Activity: In addition to the structural optimization, the development of Cu-based bimetallic catalysts has attracted wide attention for ECR application, which can break the scaling relationship and tune the interactions between catalysts and key intermediates, and thus optimize the product distributions and reach faster kinetics. Chen’s group designed a model system that combined Cu nanowire with different atomic ratios of Ag as electrocatalysts for ECR.[107] The study demonstrated that Cu₆₈Ag₃₂ nanowire exhibited the peak selectivity for methane production with a FEₐCH₄ value of ≈60%, approximately threefold higher than that of Cu nanowires (Figure 12a,b). Meanwhile, analyzed by in situ XRD, XAS, and Raman techniques, a dynamic reconstruction was detected on the surface of the catalyst, which could bring about the reoxidation of Cu⁰ and effectively stabilize the chemical state of Cu, and thus generating active species under reaction process to produce desirable products. Ag was used as an assistant to activate Cu toward the generation of multi-carbon products according to the tandem mechanism. For example, Buonsanti et al. synthesized Ag-Cu nanodimers where Ag nanoparticles acted as nucleation seeds for the Cu domain. Due to the tandem catalytic mechanism and unique electronic effects,[108] bimetallic Ag-Cu showed an enhanced FEₐCH₄ value by three times higher than that of pure Cu (Figure 12c,d). The Ag-Cu nanodimers consisted of two segregated domains, which provided an optimal interface and Cu domain size for the improvement of *CO₂ dimerization, and finally achieved the maximum CₓHᵧ yield.[109] The experimental and theoretical results indicated that the synergistic effect of Langmuir-Hinshelwood *CO + *CH₄, and Cu-Ag boundaries could contribute to the abundant CO spillover and facilitate the subsequent CO dimerization, and therefore improved the selectivity of ethanol formation (Figure 12e–i). Au has been regarded as another effective promoter for Cu-based catalysts to realize satisfactory activity toward ECR. Accordingly, Sun et al. assembled Au nanoparticles on Cu nanowires with 4,4’-bipyridine as a linker to form an Au-bipy-Cu composite catalyst.[110] With the Au/Cu atomic ratio of 50%, the Au-bipy-Cu composite performed a high FE for total products of 90.6% at −0.9 V versus RHE, with a maximum FE value of 25% for methanol formation among the liquid carbonaceous product (formate, methanol, and acetate) distribution (75%). Au-catalyzed ECR could enrich CO near Cu sites, which accelerated CO dimerization. Moreover, the addition of bipy Lewis base centers further promoted the catalytic activity of Cu catalysts since they could effectively stabilize the *CO₂ intermediates and provide abundant protons on Cu sites, synergistically resulting in high selectivities for multi-carbon compounds production. Ternary Cu-Au/Ag nanoframe reported by Peng et al. was also an ideal catalyst to break the linear scaling relationship of the interactions between catalyst and intermediates, and overcome the kinetic barriers of CO₂ conversion.[111] Evidenced by operando and DFT studies, Ag/ Au structures played a synergistic effect for high content of CO formation (CO₂ → *CO), while Cu was responsible for the CO–CO coupling (2*CO → *OCCO), thereby realizing a satisfying FEₐCH₄ value of 69 ± 5% with extraordinary catalytic stability and material integrity. The work emphasized the cooperative roles of tandem effect, electronic effect, and defect engineering, which offered some insights for achieving highly efficient CO₂-to-C₂₄
compounds conversion. Owing to its low cost and non-toxicity properties, bimetallic CuZn is an ideal alternative for ECR to realize the high activity and selectivity of carbonaceous products. Cuena’s group employed Cu100−xZnx nanoparticles to explore the activity-structure relationship.[70] When the concentration of Zn in the range from 10% to 50%, CH4 was determined to be the main product with a peak value of FECH4 ≈70%, while HER side-reaction was effectively suppressed. However, when the content of Zn continuously increased, the selectivity of CH4 decreased and CO became the main product in ECR. In virtue of operando XAFS and XPS analysis, the alloying of Cu atoms with Zn played a vital effect in the product selectivity. The variations of Cu-Zn interaction during ECR could switch the selectivity from CH4 on Cu-ZnO species to CO over CuZn alloy species. Therefore, when Cu and Zn atoms homogeneously distribute in the CuZn catalysts, CO could spill from Zn to Cu sites, thus greatly improving the formation of high value-added products. Furthermore, Goddard’s group synthesized Cu-Bi nanoparticles, which exhibited a high FFCH4 value of 70.6% at −1.2 V versus RHE, ≈25 times higher than that of Cu nanoparticles.[112] The experimental results indicated that the partially oxidized Cu acted as the possible active sites due to the electron withdrawal ability of Bi, which significantly lowered the formation barrier for *COH formation, and thus realized high activity and selectivity in ECR.

Influence of Surface Modification on Activity: Surface modification of Cu catalysts with organic additives is another effective strategy to optimize the electronic structure and alter the interactions between catalysts and intermediates and thereby facilitating the kinetic rates and product selectivity toward CO2RR. Wang’s group designed an amino acid (zwitterionic glycin) modified Cu electrode, which stabilized the key intermediate *CHO and further enhanced the selectivity of hydrocarbons including C2H6, C3H6, and C4H8.[113] Similarly, Zhang et al. introduced −NH2 terminated methylcarbamates onto Cu surface, which accelerated CO conversion by 17% and achieved an extraordinary FFCH4 value of 81.6% due to the affinity interaction between Lewis acid (−NH2) and Lewis base (CO2).[114] DFT results emphasized that the −NH2 groups could stabilize *CO and *CHO intermediates onto Cu surface, which facilitated the subsequent hydrogenation reaction toward hydrocarbons production. Pyridine has also been selected as another additive to improve the catalytic activity and selectivity of Cu electrodes. For example, Agapie et al. synthesized a novel catalyst combining polycrystalline Cu electrode with N-substituted pyridinium additives, which achieved a high selectivity for C2 and C3 hydrocarbon production, especially with a quite high ratio of C2/C1 (>100).[115] Similar to the previous work, Agapie’s group employed a simple approach to integrate a polycrystalline Cu electrode with an N,N’-ethylene-phenanthrolinium dibromide additive to form a composite catalyst,[116] which realized the selectivities for C2 products up to 70% with stable catalytic activity as well as structural integrity upon 40 h test at relatively low overpotentials. The organic additives played essential roles in the improved ECR process. In detail, the organic additives could stabilize the nanostructure on the surface of Cu, which improved C–C dimerization and suppressed HER and CH4 formation, thus realizing the highly efficient and durable C2 chemical formation during the ECR process.

Influence of Supports on Activity: Anchoring Cu on specific support brings about a synergistic effect for the catalytic system. First, the well-dispersed metal catalysts can expose abundant active sites, and thus accelerate the reaction rates toward ECR. Second, the strong interaction between Cu-based catalysts and supports can prevent the aggregation of Cu sites, which therefore significantly maintains the structure and composition of the catalysts and realizes long-term stability. For example, N-doped graphene,[117] N-doped carbon,[118] and MoS2 nanoflowers[119] have been selected as ideal substrates to strongly immobilize Cu nanoparticles via coordination structure or defect sites, which could stabilize the reaction intermediates and therefore improve the selectivities toward high value-added products such as CH2 and ethanol at moderate overpotentials in ECR. However, the metal utilization of nanoparticles is not satisfactory, for example, only 20% of atoms are exposed on a 2 nm particle. Thereafter, dispersing Cu as isolating single atoms emerges as a flourishing approach to maximum the atom economics and improve the catalytic performance in ECR, which can also increase the stability by preventing Cu atoms from oxidation or aggregation due to the strong metal-support interaction. However, Cu single-atom electrocatalysts commonly undergone the two-electron transfer process to generate CO and formate as the main products, while the further transformation into hydrocarbons or alcohols has been rarely reported so far. Accordingly, He et al. designed a facile method to realize the large-scale synthesis of atomically dispersed Cu on the surface of through-hole carbon nanofibers (CuSAs/TCNFs) (Figure 13a,b).[105] The as-obtained CuSAs/TCNFs catalyst reached a high faradic efficiency for methanol production (FEmethanol) value of 44% with an extraordinary partial current density for C1 chemicals of −93 mA cm−2 and great durability of 50 h (Figure 13c,d). DFT calculations indicated that the atomic Cu sites could stabilize *CO intermediate which could be further hydrogenated into methanol (Figure 13e,f). Noted that only C1 chemicals (i.e., methanol and CH2) could be produced using this catalyst in ECR. Accordingly, DFT calculations simulated the reaction pathways of CO2-to-methanol or CO2-to-CH4 on the surface of CuSAs/TCNFs, indicating that the *COH intermediate preferred to be hydrogenated into *CHOH (≈0.86 V) rather than *CO (≈1.88 eV), which was the key intermediate for CH4 formation. Accordingly, the atomic Cu sites on CuSAs/TCNFs preferred methanol formation rather than CH4. Other supports (e.g., CeO2,[120] TiC, and TiN[121]) have also been employed as ideal supports to prepare Cu-based single-atom catalysts by taking advantage of their oxygen vacancies or surface defect sites. The resulting electrocatalysts performed high selectivity for CH4 generation at quite low overpotentials, primarily because of the highly effective Cu atoms for CO2 activation, intermediate stabilization, and CO2 hydrogenation.

II) Most studies have shown that Au and Cu-based materials are regarded as the only choices to achieve highly selective and active performance for conversion of CO2 to C2 chemicals. Therefore, it is worthy to develop a novel efficient electrocatalyst from other metals for C2 hydrocarbons and oxygenates formation. Ag used to be reported preferring CO formation, while it was employed by Broekmann’s group for hydrocarbons production.[15] The researchers adopted an electrodeposition method to construct a novel foam-like Ag nanocatalyst, which created abun-
and OCH$_3$ intermediate contributed to high methanol selectivity. Meanwhile, the weaker interaction between alloys surface and OCH$_3$ intermediate binding capacity and thereby increase *CO surface coverage and suppressing the HER. Furthermore, the active sites effectively lowered the kinetic energy barrier for the subsequent *CO hydrogenation in *CHO intermediate. In addition, the surface sites enabled the stabilization of *CHO species for the final hydrocarbon production. Accordingly, Ag-foam catalyst achieved a maximum FECH$_4$ value of 51% at a higher potential of −1.5 V versus RHE, (Figure 14c), together with a maximum FE$_{CO}$ value of 8.6%. The Ag-foam catalyst designed in this work shared a common reaction mechanism with the traditional Cu-based catalysts, which provided new insights for CO$_2$-to-hydrocarbons/oxygenates conversion (Figure 14d). Roy et al. reported carbon-supported PtZn alloys in the ECR study for the first time, which realized FE$_{CO}$ value as high as 93.5% in an ionic liquid electrolyte, and the current density was sixfold higher than that of Cu electrocatalyst at the same reaction condition. Hexagonal BN sheet with a similar structure as graphene has also attracted increasing attention in the catalysis field. Recently, Luo et al. synthesized C-doped and line-defect embedded BN nanoribbons for ECR, which could switch selectivities of different products at various active sites (Figure 14g). For example, C-doped zigzag and armchair BN nanoribbons presented strong adsorption ability to the *CO intermediate, which promotes CH$_4$ formation after easy subsequent hydrogenation of *CO. When referred to the role of edge B atoms and C dopants, they would act as dual active sites to facilitate the couplings between *CH$_2$ and *CO intermediates and thereby contributing to a high selectivity for C$_n$H$_{2n}$ or ethanol formation. In conclusion, Cu-based materials, other transition metals, their alloys/carbides/chalcogenides, and metal-free catalysts have offered more choices and opportunities for satisfactory CO$_2$-to-hydrocarbon/oxygenate conversion with optimal activities, selectivities, and long-term stabilities.

Concluded from above-mentioned research work, different types of metal or non-metal based materials might exhibit various selectivities for a specific product. Therefore, we meticulously discussed the representative catalyst types and the corresponding catalytic mechanisms in each reaction pathway part, aiming at guiding the readers to rationally design the catalysts for highly-selective CO$_2$-to-specific product conversion. For example, due to the high intrinsic activity in converting CO$_2$ to produce C$_{1-3}$ oxygenates, including methanol and 1-propanol at quite low applied potential (−1.38 V vs Ag/AgCl,$^{123}$) it is the first time to report the metal alloy in generating C$_1$ product with the stability of the overall system more superior to that of Cu-based systems.

Furthermore, transition-metal carbides and chalcogenides have also attracted considerable attention for ECR due to their easy-available property, unique electronic and geometric structures. Peterson et al. found that Mo$_2$C could realize CO$_2$-to CH$_4$ conversion at low potentials and the onset potential is only −0.55 V versus RHE,$^{124}$ which is much lower than that of Cu electrocatalysts (onset potential of −0.80 V vs RHE) (Figure 14e). The main reason is that the carburizing transition metals could effectively tune the electronic structures of the catalysts and lower the free energy difference between *CO and *CHO intermediates, and thus decreasing the overpotential required for CH$_4$ production (Figure 14f). Because of the unique structure, the ultrathin MoTe$_2$ nanosheets were prepared by destroying the week van der Waals force between adjacent layers.$^{125}$ Due to the increased specific surface area and high density of low-coordinated surface sites, the ultrathin MoTe$_2$ nanosheets performed an optimal FE$_{CH4}$ value of 83% ± 3% and extraordinary durability of 45 h at −1.0 V versus RHE in an ionic liquid electrolyte, which could be comparable to the as-reported Cu-based catalysts toward CO$_2$RR. The aforementioned works provide new inspirations for the conversion of CO$_2$ to valuable chemicals by using other transition metals rather than metallic copper catalysts.

Recently, N-doped carbon materials, especially N-doped graphenes have been found to achieve the electrocatalytic reduction of CO$_2$ to CH$_4$ without metal sites, which process controllable structures, high stability, and easily accessible properties. Han et al. reported a graphene-like N-doped carbon material,$^{126}$ which realized FE$_{CH4}$ value as high as 93.5% in an ionic liquid electrolyte, and the current density was sixfold higher than that of Cu electrocatalyst at the same reaction condition. Hexagonal BN sheet with a similar structure as graphene has also attracted increasing attention in the catalysis field. Recently, Luo et al. synthesized C-doped and line-defect embedded BN nanoribbons for ECR,$^{127}$ which could switch selectivities of different products at various active sites (Figure 14g). For example, C-doped zigzag and armchair BN nanoribbons presented strong adsorption ability to the *CO intermediate, which promotes CH$_4$ formation after easy subsequent hydrogenation of *CO. When referred to the role of edge B atoms and C dopants, they would act as dual active sites to facilitate the couplings between *CH$_2$ and *CO intermediates and thereby contributing to a high selectivity for C$_n$H$_{2n}$ or ethanol formation. In conclusion, Cu-based materials, other transition metals, their alloys/carbides/chalcogenides, and metal-free catalysts have offered more choices and opportunities for satisfactory CO$_2$-to-hydrocarbon/oxygenate conversion with optimal activities, selectivities, and long-term stabilities.
CO, Au- and Ag-based materials have been intensively studied as the model catalysts for achieving high CO yield. Furthermore, systematic overviews have been unfolded on catalysts concerning design principles, covering the aspects of nanostructuring (e.g., shape, exposed facet, crystal phase, defective site, oxidation state, and size), alloying, surface modification, and support addition, to optimize the catalytic performance of the designed catalysts. These principles can act as the exemplary embodiments for other novel material design, so as to grasp the critical factors affecting the catalytic performance. Last but not the least, the above-mentioned contents emphasize the structure-activity relationships of electrocatalysts in ECR mainly from the aspects of the reactant diffusion/adsorption, intermediate types and the adsorption/desorption energy on the active sites of catalysts by in situ/operando characterizations and DFT calculations. Thus, the origin of selectivity toward a specific carbonaceous product on the basis of catalysis principles is more clear. Therefore, this section could provide a better mechanism understanding and guide for the development of the next-generation catalysts.

3. Photoelectrocatalysis

3.1. Motivation and Principles

When compared with photocatalysis, photoelectrocatalysis has more choices in semiconductors, which can better meet the requirement of suitable band position and redox potential level for CO₂ reduction with the assistant of an external bias. When compared with electrocatalysis, photoelectrocatalysis can decrease the overpotentials toward CO₂ reduction under light irradiation. In general, the photoelectrocatalytic CO₂ reduction processes can be classified into two systems, that is, half-cell and full-cell. How-

Figure 14. a) Side-view and b) top-down SEM images of the Ag foam. c) The plot of the Faradaic efficiencies as a function of applied potentials for Ag foam. d) Scheme illustration of the proposed reaction pathway of CO and hydrocarbon formations on Ag foam. Reproduced with permission.[55] Copyright 2018, American Chemical Society. e) CV of Cu sheet, Mo₂C sheet, and Mo₂C bulk powder for ECR in 0.1 m KHCO₃. f) Free energies of CO₂ conversion to CH₄ over Mo₂C(100) surface. Reproduced with permission.[124] Copyright 2016, American Chemical Society. g) Different active sites of BN for the switchable selectivities of reduction products. Reproduced with permission.[127] Copyright 2018, American Chemical Society.
ever, the latter has been regarded as an ideal photoelectrochemical (PEC) CO₂ conversion, which takes place in an aqueous electrolyte to produce carbonaceous products and oxygen in photocathode and photoanode via CO₂ reduction and H₂O oxidation reactions, respectively. Accordingly, highly efficient materials are required to be elaborately designed to simultaneously boost CO₂-to-fuels conversion and OER at a large photovoltage without any sacrificial reagents. Namely, the photoelectrode materials should possess a suitable band structure. In detail, the CB position of a photocathode semiconductor is required to be negative than the reduction potentials of CO₂-to-chemicals conversion aiming at achieving CO₂ activation and C=O bond cleavage, while the VB position of a photoanode semiconductor should be positive than the H₂O oxidation potential. Hitherto, various electrode materials have been developed for efficient photoelectrocatalytic CO₂ reduction, such as pristine semiconductors, heteroatom-doped semiconductors, metal/semiconductor heterojunctions, metal complex/semiconductor hybrids, and nanostructured semiconductors, etc. These adopted strategies aim at enhancing light absorption ability, photo-induced carrier mobility, and catalytic stability, finally realizing highly efficient photoelectrocatalytic CO₂ reduction with satisfying solar energy conversion efficiency.

3.2. Plausible Routes for CO₂ Conversion

Similar to the aforementioned techniques, the products generated from photoelectrocatalytic CO₂ conversion can be generally divided into four categories: Formate, CO, alcohols (mainly methanol and ethanol), and some other hydrocarbons. The different photoelectrodes, applied potential, and reaction conditions can lead to distinct CO₂ conversion pathways, therefore resulting in different products.

3.2.1. CO₂ to Formate/Formic Acid

Li et al. designed a CoOx nanosheet-like photocathode combined with the photoanode of TiO₂ nanotube arrays (Figure 15a),[128] which achieved a peak value of FEformic acid (60.9%) at a small applied potential (−0.85 V vs Ag/AgCl) in CO₂ electroreduction.
When compared with electrocatalysis, the PEC system exhibited an optimal formic acid production yield excess 80 μmol cm^{-2} h^{-1} with the assistant of solar energy and enhanced performance at the low potentials ranging from 0.5 to 0.9 V (vs Ag/AgCl), which realized efficient utilization of solar energy and high-performance CO2 conversion (Figure 15b,c). To boost the photocatalytic efficiency of pure semiconductors, heteroatoms-doping into the structure of catalysts has been demonstrated to be an effective approach to facilitate the CO2/intermediate activation. Herein, Yang et al. reported a Bi, S co-doped SnO2 catalyst for photoelectrocatalytic CO2 reduction,129 which provided abundant active sites and surface defects for formate production, and therefore achieved an ideal CO2 conversion process with a relatively low overpotential and high partial current density compared with pure SnO2 catalyst. The Bi, S co-doped SnO2 catalyst showed higher photocurrent density than that of SnO2, indicating that the heteroatoms doping could enhance the separation efficiency of photo-excited electron-hole pairs and improve the visible-light absorption ability due to the narrower bandgap. Besides, noble metals and nanostructuring have been adopted to improve the photoelectrochemical activity of semiconductors. For example, WO3 has been regarded as a popular semiconductor material with good optical and conductive properties for photoelectocatalytic reactions, while the wide bandgap and the rapid recombination of electron/hole pairs largely restrain the photoelectrochemical efficiency of WO3. Accordingly, Bal’s group employed a one-pot synthetic approach to prepare highly-dispersed Ag embedded on WO3 nanorods (Ag/WO3-NR).130 The synergistic interface of Ag/WO3-NR catalyst contributed to the highly efficient photoelectrocatalytic conversion of CO2 to formate. The SPR effect of Ag nanoparticles not only generated hot electron/hole pairs to activate CO2 hydrogenation (Figure 15d,e) but also selectively produced formate when applied external potentials. Therefore, Ag/WO3-NR exhibited a significantly enhanced current density and an optimal production rate of formate (31.7 mmol h^{-1}) at a given potential of −1 V versus Ag/AgCl in PEC CO2 reduction, which overcame the required overpotential for ECR system with the assistance of light irradiation. Chemically combining metal complexes with semiconductor photo-absorbers has been regarded as a promising way to attract more efficient photo-induced electrons for highly active CO2RR performance. Zhao and co-workers have designed a bimimetic photoelectrocatalyst composed of Co porphyrin and g-C3N4 nanosheet via π–π supramolecular interaction,131 which performed a satisfying formic acid production yield of 154.4 μmol during 8 h photoelectrocatalytic CO2 conversion with a TON of 137 and nearly 100% selectivity for liquid products at −0.6 V versus NHE with the existence of light source (Figure 15f,g). The Mott-Schottky plot and DFT calculations demonstrated the reaction mechanism of Co porphyrin/g-C3N4 nanosheet catalyst for PEC CO2 reduction as follows (Figure 15h). Under solar light irradiation, the photo-excited electrons could transfer from g-C3N4 nanosheet to the CB of Co porphyrin, while the holes flowed from Co porphyrin to the VB of g-C3N4 nanosheet, and thereby resulting in the formic acid product via the Co porphyrin-CO2 adduct with abundant protons. The aforementioned studies provide constructive insights into the synthesis of efficient PEC catalysts for formic acid/formate with a high production yield. On the other hand, these studies are conducive to enhancing the performance of photoelectrocatalytic CO2 reduction with improved reaction kinetics and clear

### 3.2.2. CO2 to CO

Photoelectrocatalytic CO2 reduction to syngas is an important platform to produce CO. So far, some semiconductors (e.g., TiO2, WO3, g-C3N4, and CdS) have realized efficient CO2 reduction half-reaction during photoelectrocatalysis, among which CdS is a promising candidate owing to its suitable bandgap for solar-light absorption and the appropriate CB position for CO2-to-fuels conversion. Accordingly, Yu et al. prepared a CdS catalyst with the presence of Cd vacancies via a facile hydrothermal approach by the addition of H2O2,132 which enhanced the separation efficiency of photo-excited electron/hole pairs and widened the light-response range (Figure 16a). Under visible light irradiation, the H2O2 treated CdS catalyst exhibited high CO selectivity and a satisfying CO formation rate of 316 μmol g^{-1} h^{-1}, which showed a 2.1-fold higher yield than that of the pristine CdS under a negative voltage (Figure 16b). Meanwhile, the photo-generated holes of CdS electrode would be consumed by the electrons from the external circuit during the PEC test and thereby preventing the CdS from being self-oxidized by the photo-induced holes and enhancing the stability of the catalytic system (Figure 16c). Additionally, other metals and metal complexes have also been adopted as cocatalysts for the selective CO2-to-CO conversion in photoelectrocatalysis. For example, Wang’s group reported an Ag nanoparticle decorated p-Si nanowire array catalyst,133 which could realize photoelectrocatalytic CO2 reduction to syngas with the adjustable H2/CO molar ratio ranging from 1 to 4. The Ag nanoparticles with the size of 8.2 nm provide a balance between *CO formation and CO adsorption, which selectively produced CO prod.
uct with a FE$_{\text{CO}}$ value of 53% with an optimal current density of 4.1 mA cm$^{-2}$ and high faradaic efficiency for syngas formation up to 97%. Si nanowires have been regarded as highly efficient photoanodes, which possess high activity, wide light-response range, rapid photo-induced carrier mobility, and long-term durability properties. Meanwhile, the CO adsorption ability could be regulated by modulating the sizes of Ag nanoparticles, which could influence the molar ratio of H$_2$/CO that is suitable for the production of different downstream products, such as, paraffin, olefins, and alcohols. Kang et al. proposed a hybrid photoelectrocatalyst combining Si wafer photoanode with Co complex photocathode ([Co(tpy)$_2$][PF$_6$]$_2$ or [Co(bpy)$_3$][PF$_6$]$_2$) toward CO$_2$-to-CO conversion,$^{134}$ realizing a photocurrent density of 1.4 or 1.0 mA cm$^{-2}$ with a photovoltage of 400 mV. Moreover, the FE$_{\text{CO}}$ value could reach up to 83 or 94% in a CH$_3$CN/MeOH electrolyte, which significantly improved the catalytic selectivity in photoelectrocatalytic CO$_2$ reduction system. The above studies shed light on the design of highly active photoelectrocatalysts for syngas production, which will inspire more research works on efficient photoelectrocatalytic CO$_2$ reduction by employing cost-effective, environmentally friendly, and homogeneous/heterogeneous systems.

### 3.2.3. CO$_2$ to Alcohols

Compared with formic acid/formate and syngas synthesis, alcohol synthesis from photoelectrocatalytic CO$_2$ reduction has more significance and commercial importance, which can be used as an environmental-friendly energy carrier that is compatible with the current industries. Plasmon-mediated electrocatalysis has been regarded as a promising approach to realize highly active and selective CO$_2$ conversion into higher value-added chemicals. For example, Lu et al. found that the plasmonic Au was an ideal catalyst to generate abundant electrons with high photon energy$^{135}$ which could be easily injected into CO$_2$ molecules and intermediates to activate the overall reaction system. Meanwhile, the generated photon flux would initiate C–C coupling toward high-order product formation (Figure 17a). Accordingly, plasmonic Au nanoparticles played a dual role, which could si-
multaneously act as a light absorber and an electrocatalyst to realize efficient photoelectrocatalytic CO₂ reduction, and thereby exhibiting a high faradaic efficiency of 52% for MeOH formation at an applied potential of −0.8 V versus RHE under 520 nm, 120 mW cm⁻² light irradiation (Figure 17b). By comparison, the Au catalyst could only generate CO product and no methanol was observed under the dark condition, which indicated that energetic charge carriers could form upon SPR excitation and further reduced the absorbed CO intermediate into MeOH via multi-electron transfer process. The study brought a new sight in designing plasmonic catalysts for photoelectrochemical CO₂ conversion systems. Cu as a promising candidate can transform CO₂ into alcohols through electrocatalytic hydrogenation process, whereas high energy barriers and inferior faradaic efficiency hinder its development. Herein, electrocatalysis combined with photo-excitation has been adopted in photoelectrocatalytic CO₂ reduction by using Cu-based catalysts. Navaee and Salimi reported a S-doped Cu₉O/CuO heterostructures,\(^{136}\) which showed enhanced activity and stability after S substitution toward photoelectrocatalytic CO₂-to-methanol/acetone conversion with an increased bandgap and an appropriate overpotential, in good agreement with the CB position of as-obtained S-Cu₉O/CuO nanoclusters. Besides, Zheng et al. designed a n-type Nb-doped TiO₂ nanotubes arrays (TNTs) with p-type CuFeO₂ as a heterostructural catalyst,\(^{137}\) which exhibited broadened light absorption and rapid photo-induced carrier mobility owing to its suitable bandgap and built-in field of a p-n junction. As a result, the CuFeO₂/Nb-TNTs photocathode realized a satisfying photoelectrochemical CO₂ reduction with high selectivity toward EtOH formation and high performance of photoresponse. The optimal current efficiency of EtOH could up to be 75% at −0.4 V with an EtOH yield of 0.66 μmol h⁻¹ cm⁻² (Figure 17c,d). Except for heteroatoms doping, metals and metal complexes have been introduced in photoelectrochemical reactions. Taking TiO₂ as an example, Luo et al. reported a Pd and rGO co-modified TNTs catalyst,\(^{138}\) which was supported by a Ti wire. With light illumination, the Pd nanoparticles not only enhanced the visible-light response region of the catalyst due to its SPR effect but also acted as the electron trapping centers. Subsequently, the photo-induced electrons flowed to the surface of rGO layers, which functioned as an electron accumulation “pool” to reduce the absorb CO₂ molecules. Therefore, Pd/rGO/TNTs/Ti exhibited improved production yields of MeOH (1624 nmol cm⁻² h⁻¹) and EtOH (536 nmol cm⁻² h⁻¹) under light irradiation assisted by an external potential (0.8 V bias potential) due to the synergistic effect of rGO and Pd nanoparticles. Notably, the ionization of CO₂ to HCO₃⁻ or CO₃²⁻ in an aqueous electrolyte was the critical step during photoelectrocatalytic CO₂ reduction, which was then reduced into MeOH or EtOH products. When referred to metal molecules cocatalyst, Zhao’s group embedded Ru-based complex (Ru-Py) onto the surface of TNTs by a covalently binding approach (Figure 17e),\(^{139}\) which could reach an enhanced cathodic photocurrent density of 1.79 mA cm⁻², 2.4 times higher than that of pure TNTs. Meanwhile, the system showed high selectivity for MeOH formation with a FE value of 63.9%, TON of 62.6, and production yield of 84.8 μmol at −0.9 V under 8 h light irradiation (Figure 17f). On the contrary, the non-covalent bonded system only achieved a MeOH yield of 41.3 μmol, which indicated the covalent linking played an essential role in carrier mobility and methanolization performance. The pyridyl that existed in Ru-Py provided abundant active sites for the pyridinium-formate immediate formation, which significantly contributed to the highly selective MeOH production (Figure 17g). Therefore, the aforementioned studies provided meaningful methodologies to optimize the design strategies of photocathodes for alcohol synthesis toward photoelectrochemical CO₂ reduction, which have made important significance in carbon resource recycling with the efficient utilization of solar energy and renewable electricity.

### 3.2.4. CO₂ to C₂, Hydrocarbons/Oxygenates

As an artificial photosynthesis process, the photoelectrocatalytic CO₂ reduction could transform CO₂ and H₂O into hydrocarbons and O₂. However, the product distribution is broad and has been usually C₁ and C₂ chemicals, such as formic acid/formate, methanol, ethanol, and CO. Accordingly, multi-carbon compounds have been regarded as higher value-added products, which provide promising pathways to complete a carbon-neutral cycle. However, there remain some challenges due to the inferior activity and poor selectivity for C–C coupling reactions. Therefore, 3D semiconductor heterostructures with abundant active sites have been designed as photocathodes to facilitate the C–C coupling pathway toward CO₂ reduction, aiming to improve the faradaic yields of multi-carbon hydrocarbons. Jing’s group fabricated a 3D C/N co-doped Zn₃:Co@Cu (Figure 18a) p-n heterojunction photocathode,\(^{140}\) which showed an optimal rate of 325μg h⁻¹ toward paraffin production at −0.4 V versus SCE and apparent moderate quantum efficiency (1.95%) of the photoelectrochemical cell (Figure 18b). The as-designed C/N-Zn₃:Co@Cu catalyst showed an improved separation efficiency of photoinduced electron-hole pairs due to the p-n heterojunctions. Moreover, heteroatom doping facilitated the absorption ability and activation capability of CO₂ molecules. Furthermore, Cu realized the significant FE for hydrocarbons and oxygenates due to its vital role in C–C coupling steps (Figure 18c). Except for metallic Cu, noble metals like Pd and Pt have also been employed as cocatalysts for efficient photoelectrocatalytic CO₂ reduction toward hydrocarbons production. For example, Pt/graphene deposited on Cu foam was used as a 3D binder-free photoelectrode,\(^{141}\) which could enhance the CO₂ reduction performance and promote C₁ products to higher value-added carboaceous compounds (e.g., acetic acid, propionic acid, and alcohols). Moreover, the carbon atom conversion rate of CO₂ reduction increased to 5040 nmol h⁻¹ cm⁻² at a voltage of 2 V (anode served as a reference electrode) together with an enhanced current density, which was 110 times higher than that in dark. Herein, the 3D Pt/graphene/Cu foam catalyst with high porosity provided abundant active sites for CO₂ activation and ¹⁸O dimerization for enhanced multi-carbon hydrocarbons synthesis. In another study, Pd/N-TiO₂/Co₂ photo cathode exhibited a peak apparent quantum efficiency of artificial photosynthesis cell of 1.78% at −1.0 V (anode served as a reference electrode) with a satisfying evolution rate (36.8 mm h⁻¹ g⁻¹) of total hydrocarbon (formate, methanol, and ethanol) formation (Figure 18d).\(^{142}\) Recently, poly pyrrole has been decorated on TiO₂ surface through an in situ oxidizing polymerization and temperature-controlled pyrolysis by Jing et al., which could
serve as a molecular cocatalyst for PEC CO₂ reduction. The resulting n-n heterojunctions (Figure 18e) achieved the highest selectivity (71.4%) for C₂⁺ hydrocarbons (alcohols, acetic acids, and acetone) production under light irradiation at −1.2 V (anode served as a reference electrode), since the pyridine-N sites and abundant defects prolonged the lifetime of photo-induced carriers and facilitated the selectivity toward C–C coupling pathway. Under the photoelectrochemical conditions, polypyrrole/TiO₂ catalyst could produce photo-excited electrons and holes through a built-in electric field. Due to the differences in the redox potentials of the two components, the electrons from the CB position of polypyrrole would flux to the CB of TiO₂, while holes were neutralized by the external electrons in the circuit. Finally, the absorbed CO₂ molecules were reduced to different C₂⁺ hydrocarbons via C–C coupling reactions. In conclusion, the deliberate design of heterojunctions, cocatalyst addition, and hierarchical nanostructuring can be regarded as promising approaches for selective hydrocarbons synthesis, which opened up new horizons for the fabrication of highly active photoelectrocatalysts toward artificial photosynthesis application.

4. Design of Flow Cell Configurations

Nowadays, numerous attentions has concentrated on the overall system to extend the laboratory-scale operation of (photo)electrochemical CO₂ reduction to industrialized application, which comprehensively considers the performance of current densities, selectivities, and energy efficiencies. Except for the development of rationally-designed catalysts, promising strategies in electrode/reactor design have remained bottlenecks that require more effort. Accordingly, rationally-designed flow cell configurations have been employed to convert CO₂ molecules to high value-added chemicals and fuels. Schematic illustrations of H-type and flow cells are illustrated in Figure 19a,b to compare their differences. In an H-type cell, only CO₂ molecules near the working electrode can diffuse and participate in the reduction reaction, and thereby the current density is restricted by the limited solubility and diffusion rate of the reactant. Hence, the sluggish reaction kinetics require large overpotentials to drive CO₂RR, and the current density is usually unsatisfactory (<30 mA cm⁻²). The relatively small current density is far from the requirements for the practical industrial-scale implementation, of which the current density should achieve up to hundreds of mA cm⁻² and even 1 A cm⁻². In comparison, the flow cell with gas-diffusion electrodes (GDEs) has been regarded as an effective configuration, which circulates the electrolyte in the overall system. This not only facilitates the reaction kinetic but also stabilizes the local reaction environment as follows. First, the GDEs provide sufficient interaction between the solid catalyst, electrolyte, and CO₂, which effectively improves the CO₂ solubility and diffusion, and thereby enhancing the current density. Second, strong alkaline electrolytes (e.g., 1–10 m KOH) can be used in the flow cells, which significantly increases the pH value of electrolytes, thus enhancing the selectivities of carbonaceous products for CO₂RR and suppressing the competing HER. Moreover, the use of KOH electrolyte can decrease the solution resistance, thus further improving the energy conversion efficiency of the overall system.

Recently, many representative works have sprung up associated with the applications of flow cell configurations for EC and PEC CO₂ conversion. For example, Strasser’s group loaded Ni-N-C catalyst onto GDEs to create a three-phase interface in...
a flow cell toward electrochemical CO2-to-CO conversion.\textsuperscript{[146]} Compared to the H-type cell, this large-scale CO2 electrolyzer cell could be conducted at industrial current densities \( \approx \) 700 mA cm\(^{-2}\). A stable FE\textsubscript{CO} value of 85% was achieved for a 20-h long-term test with a large \( j_{\text{CO}} \) up to 200 mA cm\(^{-2}\). Such outstanding activity provides tremendous potential for the commercialization of the electrochemical CO\textsubscript{2}RR technique toward highly-selective C\textsubscript{1} chemical production. Besides CO\textsubscript{2}-to-C\textsubscript{1} conversion, Cu-based electrocatalysts have been employed in microfluidic CO\textsubscript{2} flow cells to realize highly-efficient C\textsubscript{2}+ chemicals production at industrially relevant rates. Jiao et al. reported the design of a highly porous Cu catalyst, which could realize an optimized selectivity of \( \approx \) 62% toward C\textsubscript{2}+ chemicals with a high current density of 653 mA cm\(^{-2}\) at a moderate applied potential of \(-0.67\) V versus RHE (Figure 19c–e).\textsuperscript{[147]} For PEC CO\textsubscript{2} conversion, the flow cell configuration also contributes significantly to improving the photocurrent density and product selectivity. For example, Lu et al. designed a membrane cathode assembly with Ag nanocubes as the electrocatalyst, which exhibited higher CO\textsubscript{2} solubility, better reactant diffusion, and inferior HER selectivity than that of the traditional CO\textsubscript{2} conversion process conducted in an H-type cell.\textsuperscript{[148]} Accordingly, the overall system achieved high electricity-to-chemicals efficiency of 92.1%. The photon-to-chemical energy efficiency reached 0.16% at the voltage of 1.2 V in a CO\textsubscript{2}-fed gas system.

Besides flow cells, the membrane electrode assembly (MEA) electrolyzer has gradually developed as another intelligent device for CO\textsubscript{2}RR, which is composed of a cathode, anode, and polymer electrolyte membrane. The electrocatalysts are directly coated on the two sides of the membrane or loaded on the GDEs, and then pressed to sandwich the membrane. Note that the membrane plays a significant role in MEA cells, exhibiting high ionic conductivity and high chemical stability. Accordingly, the MEA electrolyzers have been successfully employed in ECR applications to...
produce high value-added carbonaceous products (e.g., CO, formate, and ethylene).  

5. Conclusions and Perspectives

The gradual increase of atmospheric CO₂ concentration has caused serious global warming, energy, and environmental crises. Accordingly, researchers have made tremendous efforts in CO₂ emission reduction during past decades. Especially, CO₂ conversion through chemical reaction path has emerged as a promising approach for the highly-efficient CO₂ utilization and upgradation. Thereinto, numerous technologies have been adopted to improve the CO₂-to-chemicals efficiency, containing chemical reforming, thermocatalysis, photocatalysis, electrocatalysis, etc. Among which, electrocatalysis has been regarded as an advisable technology that can realize efficient CO₂ reduction in a clean and renewable manner driven by electricity. In comparison, photoelectrocatalysis as an intelligent technology can combine solar and electronic energy as an integral whole, which not only modulates product selectivity in CO₂ reduction by semiconductors with suitable band structure but also decreases the overpotentials with the assistance of solar energy compared to pure ECR. Meanwhile, the applied potentials indeed accelerate the separation of photo-generated hole/electron pairs to achieve long-term durability. Since multitudinous significant work has been continuously published regarding these fields, this review provides a timely and comprehensive overview of the state-of-the-art of electro- and photoelectrocatalytic CO₂ conversion into fuels and value-added chemical feedstocks, such as formate, CO, CH₄, alcohols, and C₂ hydrocarbons/oxygenates in the past 5 years. On account of the well-designed catalysts playing a significant role in the promoted development of electro- and photoelectroreduction of CO₂, this review pays special attention to the recent development of rational designing strategies for heterogeneous catalysts toward highly-efficient CO₂ conversion. The reaction pathways and catalytic mechanism toward different carbonaceous products via these technologies have been briefly discussed based on in situ/operando analysis and DFT calculations, which can provide a thorough understanding of key factors about rate-determining steps and the adsorption or formation of reactants/intermediates, and thus realizing optimal CO₂ reduction activity and high selectivity. Accordingly, when referred to the similarities between ECR and PEC CO₂ conversion, both techniques can effectively control the reduction products by modulating the reaction conditions (e.g., given potentials, band structures, reaction temperature). Meanwhile, both systems can be conducted in a clean and green manner driven by renewable energy sources (i.e., sunlight and electricity), and of which chemical consumption can be minimized to H₂O. Furthermore, the (photo-)electrochemical reaction setup is modular and compact for scale-up application. However, when referring to the differences that exist between ECR and PEC CO₂ conversion, the product selectivities in ECR mainly depend on the energy of electrons (namely, applied potentials) by tuning the linear-scaling relationship of CO₂-reduced intermediates binding energies onto the surface of catalysts. Concerning PEC CO₂ conversion, the electrode materials must be comprised of semiconductors. The product selectivities are not only affected by applied potentials, but also the energy of photoexcited electrons, which is determined by the CB edge of the semiconductor photoelectrodes.

However, the advancements in electro- and photoelectrochemical CO₂ transformations are still not satisfactory in terms of energy efficiency and technical and economic viability for practical application. There are still some challenges and perspectives are referred to as the following points. 1) As for catalysts, cost-effective materials and facile synthetic methodologies should be explored for large-scale production. Meanwhile, the rational design of nanostructured catalysts for highly selective toward one specific product is of great importance. In addition, the long-term durability of catalysts is regarded as another essential parameter that determines the practicability of catalytic systems in a practical device. In terms of photoelectrochemical CO₂ conversion, more efforts are required to be paid into novel catalysts (e.g., semiconductor photo-absorbers/catalysts, surface cocatalysts) to realize high solar-to-fuel efficiency, which should not only provide a narrow bandgap to absorb a wide spectral range of solar light, but also perform efficient CO₂ reduction with high selectivity and activity when given the external potentials. 2) As for reactor configurations, an H-type electrolytic cell with an ion-exchange membrane has been widely used in CO₂ electro- and photoelectroreduction. However, to meet the requirements of industrial-scale, ultra-high current density, and long-term stability should be achieved to make great progress in the high yields of valuable fuels and chemicals, providing more possibilities for future practical application. Therefore, the ration design of continuous flow cells and microfluidic have been made much effort to reach high activity in recent years. The GDEs, flow cell, and MEA electrolyzers can effectively adjust the local environment (e.g., pH value of electrolyte) and reaction conditions (e.g., CO₂ concentration) of CO₂RR system, which can not only improve the reactant flow rates, but also achieve higher partial current densities, and thereby promoting CO₂ conversion rates, efficiencies and selectivities. For example, GDEs provide a porous catalyst layer along with diffusion media to accelerate the reactant transport and distribution, aiming at achieving high current density and low transport losses. Moreover, the pH value of the employed electrolyte is an important factor in influencing CO₂RR activity and selectivity. Especially, enhanced activity toward high value-added C₂ products is favored at high pH values in the flow cell, which can hardly be achieved in a conventional H-type system. Furthermore, flow cells and MEA provide convenience for the separation of different reduction products. The above-mentioned advantages facilitate the large-scale employment of CO₂RR at industrial level. In addition, in the PEC CO₂ reduction field, full cell and tandem cell (photovoltaic cell-PEC reactor) has been designed with a suitable semiconductor as a photoanode for water oxidation, and simultaneously accelerate electron mobility for CO₂-to-chemicals reduction, facilitating the overall solar energy conversion efficiency, and thus realizes the long-term goal of 10% solar-to-fuel efficiency. 3) As for the measurement of FE and solar-to-fuel efficiency of EC and PEC CO₂ reduction, a standardized metrology method should be conducted to better compare the results from different research groups, mainly concerning illumination condition, selected electrolyte and reactors, CO₂ feeding, and product analysis methods, etc. 4) As for mechanism investigation, CO₂ conversion processes can be explored with the assistance of advanced analysis techniques, such as, operando...
spectrum (e.g., in situ XPS, IR, and EXAFS, etc.), and computational analysis. For example, theoretical calculations have recently been regarded as an essential method for catalyst screening, d-band center prediction, and Gibbs free energy trends. The DFT calculations and molecular dynamics simulations provide valuable information with existing data to conduct more sophisticated methods for holding great promise in fundamental studies of catalysis, and thereby, further pushing the development and prosperity of (photo-)electrochemical CO$_2$RR areas. Accordingly, the rate-determining steps, the adsorption behavior of intermediates, kinetic energy barriers, and thermodynamics limits would be dealt with, which reveals the key factors for improving the activity and selectivity of catalysts. Accordingly, the combination of experimental results and theoretical studies can figure out the close relationship between catalysts surface and complicated reaction networks.

In conclusion, the comprehensive summary of fundamentals, well-designed catalyst strategies, CO$_2$RR testing, and in-depth mechanism exploration of electro- and photoelectrochemical CO$_2$ conversion in this review will attract wide interest from both the academic and industrial communities concentrating on material science, heterogeneous catalysis, green chemistry, as well as energy and environmental catalysis.

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Conflict of Interest

The authors declare no conflict of interest.

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CO$_2$ conversion, electrocatalysis, photoelectrocatalysis, value-added chemicals
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