Single-atom catalysts: stimulating electrochemical CO₂ reduction reaction in the industrial era

Zedong Zhang and Dingsheng Wang

Carbon monoxide and formic acid play a significant role in industrial processes and are exceedingly economical C₁ products in electrochemical CO₂ reduction reactions (ECR). To improve the yield and catalytic activity in ECR processes, numerous researchers have optimized catalysts and upgraded the electrolyzer. An increase in the electrocatalytic current density and production yield can significantly enable industrial electrochemical conversion from CO₂ to CO/HCOOH. However, the crucial factor restricting the industrialization of ECR is the lack of high-efficiency catalysts. Thus, the development of efficient catalysts for ECR has been ongoing for decades. Traditional nanostructure catalysts suffer from poor product selectivity, unstable structure, and inhomogeneous catalytic active centers, seriously affecting the industrial process. In contrast, single-atom catalysts (SACs) process a uniform coordination environment of active centers, maximum atom-utilization efficiency, and impressive stability, which are suitable as catalysts for industrial ECR. This perspective highlights and summarizes the designed coordination of SACs to obtain a high yield of CO/HCOOH. Moreover, we discuss the fabrication of the electrolyzer and the method of techno-economic assessment (TEA). Finally, we summarize the opportunities and challenges for ECR in industrial processes.

1. Introduction

The continued growth in carbon dioxide emissions has caused ongoing climate change and severe disasters globally.¹ Thus, individual countries plan to peak the CO₂ emission and implement carbon neutrality before the middle of the 21st century, which urgently require the development of carbon capture, storage, and transfer technologies. CO₂ transfer with high efficiencies and low emissions is the holy grail of catalytic chemistry. Traditional thermocatalytic CO₂ reduction is widely used in the industry, which benefits from mature technology and high yield of reduction.²–⁴ However, thermocatalysis typically uses hydrogen as the reductant, and the catalytic temperature is high (~573 K). Photocatalytic CO₂ reduction utilizes solar energy exclusively to reduce CO₂ to CO or hydrocarbon fuels with water and photoelectrons.⁵ However, its low yield and poor reduction selectivity restricts wide research on CO₂ reduction in the industry.
Fe, Co, Ni, Zn, Pd, Ag, Cd, and Ir SACs prefer reducing CO2 to also strengthens the electrons transfer on the catalyst surface. These properties restrict the application of nanoparticles in industry. The strong metal-support interaction (SMSI) of SACs result in their aggregation and inactivation. Furthermore, they presented an in-depth understanding of the structure–activity relationship of SACs and their optimizations in the ECR process. The Faraday efficiency (FE), current density, energy efficiency (EE), cell voltage (CV), and catalytic stability have gradually become satisfactory to meet the demand of industrial ECR for C1 products. Among the studies on ECR systems, the electrolysers have developed from the H-cell to the flow cell and membrane electrode assembly (MEA) electrolyzer. The current density and single-pass conversion have significant improvements, which make the large-scale reduction of CO2 to C1 products possible.15,26

Techno-economic assessment (TEA) is widely applied to evaluate the economic viability of the ECR process for C1 products.27–29 There are many benchmarks to assess, including the revenue from the products, electricity cost, electrolyzer cost, separation consumption, and catalyst cost. The critical factors for catalysts are the catalyst cost, stability, and selectivity in the ECR, which determine where the catalysts can be scaled up in industry or not.29 SAC catalysis of ECR usually exhibit excellent C1 product (CO/FA) selectivity, low cell potential with high EE value, and outstanding catalytic stability as a result of the strong metal–support interaction.31,32 In this perspective, we mainly focus on the SACs in ECR to produce C1 products and the challenges in designing and synthesizing effective SACs. Further, we discussed developing an ECR electrolyzer and combining it with traditional TEA methods to predict the possibilities and challenges of SACs in industrial processes (Fig. 1).

2. SACs for simple C1 products

In general, the catalytic process for 2e− products (from CO2 to C1 products, such as CO and FA) involves the following steps: (1) the concentration of CO2 feedstock; (2) diffusion of CO2 in electrolyte, and then to the catalytic interface; (3) adsorption and catalytic conversion of CO2 into *OCHO or *COOH.
intermediates on the catalyst surface; (4) desorption of over-saturated CO\(_2\) and the C\(_1\) products; and (5) transport of products and collection of redundant CO\(_2\) feedstock. Understanding the structure–activity relationship focuses on the adsorption/desorption steps and the reaction activation energy during the catalytic process. As one of the critical 2e\(^-\) C\(_1\) products in ECR, CO is commonly associated with carbon-bound intermediates, as shown in Fig. 2, where protons or electrons in the electrolyte induce the metal-catalytic active sites to generate metal\(^*\) (M\(^*\)) and adsorb CO\(_2\) to form a carbon-bond (M–C). The intermediates on the catalytic interface are M–\(\text{COOH}\), which further undergo dehydroxylation to CO. The oxygen-bound intermediate theory is used to explain the catalytic process, where FA: M–\(\text{OCOH}\) is the key intermediate, which is further converted to FA products through the dihydroxylation mechanism. Surface catalysis of these 2e\(^-\) C\(_1\) products occurs at single dispersed atomic active centers. Different species of metal elements and the coordination environment of the single dispersed metal atoms are the crucial factors affecting the activity and selectivity in ECR.\(^{33}\)

2.1. Carbon-supported SACs

Carbon-supported SACs is an essential branch of SACs, which are effective catalysts for ECR. Previous work showed that most carbon-supported SACs with abundant nitrogen-doped are stable activated single-metal ions to form the homogeneous microcosmic structure of M–N\(_2\)-C.\(^{\text{40-48}}\) The universal methods for synthesizing carbon-supported SACs include molecular skeleton encapsulation,\(^{\text{44-48}}\) impregnation and pyrolysis,\(^{\text{49-51}}\) coordination regulation,\(^{\text{52-54}}\) and thermal dispersion.

2.1.1. Molecular skeleton encapsulation strategy. The molecular skeleton encapsulation strategy involves the use of porous nanomaterials to encapsulate the metal ions or coordination compounds. Also, through pyrolysis treatment, the metal species can be reduced in the porous nanomaterial substrate. This strategy effectively restricts the agglomeration during heat treatment, where the traditional porous nanomaterials include MOFs, COFs, zeolites, and CNTs. Jaouen et al. sacrificed zeolitic imidazolate framework (ZIF-8) as a precursor of N-doped carbon materials.\(^{\text{55}}\) The imidazolate framework was the molecular skeleton to encapsulate different transition metal salts and prepare M–N\(_x\) (M = Mn, Fe, Co, Ni, and Cu) SACs through pyrolysis at 1050 °C. They found that all the carbon-supported SACs that exhibited preferable ECR gave particular CO products at \(-0.5\) V and \(-0.6\) V vs. RHE (Fig. 3a and b). This research also considered the volcano curve of current density versus the atomic number of the transition metal elements. In combination with DFT calculation, the computational reaction activity of the M–N\(_x\) sites shown in Fig. 3c follows the order of FeN\(_x\)-\(\text{H}_2\text{O}\) < MnN\(_x\) < Ni\(^{3+}\)N\(_x\) ≈ CuN\(_x\) < CoN\(_x\)-\(\text{H}_2\text{O}\) < Ni\(^{2+}\)N\(_x\). The tendency of the carbon-supported SACs in ECR can be attributed to the too high \(\text{COOH}\) adsorption energy over traditional Mn, Fe, Co-N\(_4\) structures, which increased the side-reaction of the HER (hydrogen evolution reaction). In contrast, Ni, Cu, and Zn-N\(_4\) structures have lower CO\(_2\) adsorption energy, which is conducive to the desorption of C\(_1\) products but at the expense of the active current density of the reaction. Jiang et al. used PCN-222 (M-TCPP (M = Fe, Co, Ni, and Cu; TCPP = tetrakis(4-carboxyphenyl)porphyrin)) to encapsulate transition metal atoms with TCPP coordination (Fig. 3d).\(^{\text{56}}\) Carbon-supported SACs were obtained through pyrolysis and removal of the metal Zr. They had a similar conclusion as the previous study, where Ni-N\(_4\) showed a perfect ECR performance compared to the other transition metals (Fig. 3e), which could reach 96.8% FECO at \(-0.8\) V vs. RHE and maintain its stability for 10 h at a current density of \(-27\) mA cm\(^{-2}\). The use of the host–guest interaction of metal–organic framework to encapsulate single atomic sites is a universal method to synthesize various element carbon-supported SACs.\(^{\text{57-58}}\) Zhang et al. explored the main group element indium, and the carbon-supported SACs with an In-N\(_4\) moiety exhibited a high probability to produce FA during the ECR.\(^{\text{59}}\) The In-N\(_4\) structure had good stability in the ECR with a reduction in the activity of less than 10% after 60 h and 96% FECO at 0.65 V vs. RHE. However, the strategy of molecular skeleton encapsulation with MOFs suffers from many problems as follows: (1) the metal nodes of the framework are challenging to treat cleanly after pyrolysis or acid etching post-treatment, where single zinc or zirconium atomic sites would affect the
activity of the reaction.\textsuperscript{20,21} Traditional methods of evaporating zinc at high temperatures result in poor reproducibility, which restricts the wide application of carbon-supported SACs in industry. (2) Tube furnaces have a significant deviation at high temperatures, and their energy consumption is too high to produce carbon-supported SACs. Wu et al. attempted to clarify the effect of different pyrolyzation temperatures (400 °C, 700 °C, 900 °C, and 1100 °C) on the Fe–N bond states for the ECR process.\textsuperscript{22} Compared to the Fe-N\textsubscript{4} carbon-supported SACs synthesized at 900 °C, the Fe-N\textsubscript{4} carbon-supported SACs synthesized at 1100 °C possessed a higher Fe\textsuperscript{CO} but lower current density, which was attributed to the higher temperature reducing the concentration of the iron element. This research also discussed the effect of different temperatures on zinc volatilization and different temperature-induced specific surface areas of carbon-supported SACs. Chen et al. regulated the nitrogen species of Fe-N\textsubscript{4} structures by regulating the atmosphere,\textsuperscript{23} pyrolysis temperature, and time, wherein the Fe-N\textsubscript{4} system with graphite N had the best Fe\textsuperscript{CO} of up to 97% at −0.6 V vs. RHE and good stability during the 24 h test under −0.6 V in an H-cell. Regulating the micromosic structure of single atoms and their coordination environment to optimize the conductivity and catalytic activity of carbon-supported SACs can truly guide more theoretical breakthroughs. Alternatively, precise regulation of the coordination environments of carbon-supported SACs lacks a universal synthesis method and enough characterization proof.

2.1.2. Impregnation and coordination regulation strategy. The ‘impregnation and coordination regulation strategy’ involves the pre-synthetic heteroatom doping of carbon, and then capturing metal atoms through the impregnating and pyrolysis process.\textsuperscript{24} This synthetic strategy can solve the problem of miscellaneous metals in the molecular skeleton encapsulation strategy. Zhang et al. primarily synthesized N-doped multiwall CNTs, and further mixed them with dicyanamide and manganese acetate.\textsuperscript{25} After low-temperature pyrolysis, they obtained Mn–C\textsubscript{3}N\textsubscript{4}/CNT catalysts with Mn-N\textsubscript{2} moieties. The D-band center models implied that the Mn-N\textsubscript{2} moieties possess a lower density of states (DOS) than the Mn-N\textsubscript{4} moieties (Fig. 4b and c, respectively). Calculation also solved the problem that the Mn-N\textsubscript{2} SACs have a too high *COOH adsorption energy. Single atomic Mn with low coordination numbers shows lower activity energy for the intermediate *COOH, which is beneficial to obtain C\textsubscript{1} products in the ECR. It also could keep the Fe\textsuperscript{CO} above 90% for 20 h under −0.55 V vs. RHE in an H-cell. Furthermore, the current density reached 15 mA cm\textsuperscript{−2} (Fig. 4a). Carbon-supported SACs with a hierarchical structure, enormous surface area, and hole capacity can effectively induce the contact between the active centers and the reactant.\textsuperscript{26,27} Li et al. used the hard-template method, as shown in Fig. 4d, to polymerize a nitrogen-containing polymer (melamine-resorcino-formaldehyde polymer) over SiO\textsubscript{2} nanospheres.\textsuperscript{28} After pyrolysis and etching treatments, the hierarchical N-doped carbon shell was further loaded with cobalt peptides to obtain SACs with Co-N\textsubscript{2} moieties. The comparative tests in the H-cell showed that this supported molecular catalyst had a higher current density and Fe\textsuperscript{CO} than conventional homogeneous molecular catalysts (Fig. 4e). Supported molecular catalysts are different from the common carbon-supported SACs obtained through pyrolysis, which have homogenous active center sites and can effectively avoid the differences in selectivity and reproducibility caused by the unevenness of the atomic structures. Nitrogen atoms stabilize the molecular catalyst on the carbon support to form a carbon-supported single-atom catalyst with M-N\textsubscript{2} moieties. The coordination numbers of the metallic centers are always more prominent than the original molecular catalysts. This is a typical type of coordination regulation strategy. Conversely, the stability of carbon-supported SACs should be considered without heat treatment, i.e., whether the molecular catalysts would change during the catalytic process. Liang et al. used the coordination regulation method to load peptide nitride (NiPc) structures regulated by different functional groups (−H, −CN, and −OMe) on the sidewalls of CNTs.\textsuperscript{30} The CNTs modified with NiPc−OMe exhibited outstanding Fe\textsuperscript{CO} and stability in the ECR test. Other functional groups (−H and −CN) were verified to increase the electron density of the nickel centers, which weakened the desorption of CO and affected the subsequent catalytic reaction.

2.1.3. Thermal dispersion strategy. The thermal dispersion method, which involves reducing nanoparticles at high temperature to release metal atoms on N-doped carbon materials, is a type of synthetic method with significant industrial prospect. A milestone was achieved in the study that showed that a 2 nm Pd nanoparticle could disperse into Pd single atoms during the carbonization of ZIF-8.\textsuperscript{31} Initially, Ostwald ripening occurs the early stages of medium-temperature carbonization, causing the size of the Pd particle to increase to about 5 nm. With an increase in temperature, zinc is evaporated from the previous position, which prompts the Pd single atoms to be released. Metallic zinc is convenient to form Pd–Zn bonds\textsuperscript{32} and Pt–Zn bonds,\textsuperscript{33} and thus the vaporization of zinc could drive the dispersion of Pd or Pt nanoparticles at high temperature. Another universal method to convert nanoparticles into single
atoms is the use of a reducing atmosphere (NH₃ and H₂) to induce the evaporation of nanoparticles and transfer single atoms to the N-sites of the carbon support.⁶¹–⁶⁴ Li et al. utilized dicyanamide to produce ammonia and promote the conversion of bismuth nanoparticles to Bi-N₄ moieties.⁶⁴ This Bi-N₄ catalyst exhibited 97% FECO under −0.5 V vs. RHE in the H-cell test.

In summary, the strategies for the synthesis of carbon-supported SACs can be classified into the four above-mentioned categories. The molecular skeleton encapsulation method is commonly used in specific mesoporous carbon materials [MOFs,⁶⁵ COFs,⁶⁶ HOFs, etc.]. The advantage of this method is that it can be universally employed to synthesize carbon-supported SACs with different elements (Mn, Fe, Co, Ni, Cu, Zn, Ag,⁶⁷ Cd,⁶⁸ In, Sn,⁶⁹–⁷⁰ Sb,⁷¹ and Bi), which show superior performances to produce C₂ products in ECR. However, carbon-supported SACs obtained with this method often suffer from inhomogeneous active centers and poor reproducibility. These disadvantages restrict their large-scale application in industry. Impregnation and pyrolysis is a method to load single-metal atoms following pre-synthetic heteroatom doping of the carbon substrate. The metal sites do not require high-temperature treatment for this method. The coordination regulation method can be employed to design the structure of substrates and precisely synthesize homogenous carbon-supported SACs. Both the impregnation and pyrolysis methods and coordination regulation methods can be employed to fabricate SACs with a more obvious micromolecular structure and produce carbon-supported SACs on the gram-scale, which is beneficial for industrial ECR. The thermal dispersion method is a novel notion of recovering the disabled nanoparticle catalysts as efficient carbon-supported SACs.⁷² However, there is still a lack of research on the dispersion mechanism and the size and capacity limits of nanoparticles.

2.2. Nanocrystal/cluster-supported SACs

2.2.1. The synthetic methods. Nanocrystal catalysts have specific lattices and defect effects. Single-metal atoms anchored on different lattices or defects can regulate the selectivity of the products. Different from the characteristics of inert carbon substrates, nanocrystals typically have high catalytic activity, which modify the single metallic atoms at the interface and can regulate the local electron density at the active centers.⁷³–⁷⁸ The impregnation method is a general strategy to synthesize nanocrystal/cluster-supported SACs, which is different from carbon-supported SACs, where galvanic replacement (GR)⁷⁹,⁸⁰ and cation/anion exchange reaction⁸¹,⁸² can occur on nanocrystals/clusters. Zheng et al. developed a method to deposit single Pd atoms on the (100) and (110) facets of Cu nanocrystals through a GR reaction.⁸³ Their different catalytic performances provide advanced perception for synthesizing nanocrystal-supported SACs. Sargent et al. synthesized Cu nanoclusters with single-dispersed gold atoms on their surface, which could effectively convert CO₂ to C₂ products in a flow cell.⁸⁴ Cu nanomaterials are widely used to produce C₂ in ECR. Single metallic atom-doped Cu nanoparticles can regulate the selectivity of C₂ products. Cation exchange reaction (CER) methods were previously applied for the synthesis of intermetallic compounds with specific proportions. Li’s group use the CER method to synthesize Pt single atomic site Cu₃S catalysts.⁸⁵ However, there are only a few studies on the synthesis of nanocrystal-supported SACs and their application in the ECR reaction. Under the guidance of theory, Li et al. found that Au nanoparticles loaded on defective CeO₂ spontaneously dispersed into Au atoms at high temperature.⁸⁶ Inspired by this, Wang et al. synthesized Ag nanoparticles/MnO₂ nanorods and converted them to Ag single-atomic sites/MnO₂ (Fig. 5a).⁸⁷ This Ag SAs/MnO₂ possessed high CO product selectivity in the ECR (FECO of up to 95.7% at −0.85 V vs. RHE, and the current density was nearly 6 mA cm⁻² in the H-cell) and good catalytic stability for 9 h at −0.85 V. A more common method for the preparation of nanocrystal/cluster-supported SACs is the “one-pot” strategy. Yao’s group mixed metal salts precursors with an Fe/Au atomic ratio of 1 : 19 and reduced them to Fe₁/Au single-atomic alloy catalysts via a “one-pot” method.⁸⁸ This material was shown to possess high activity and selectivity in the ECR for CO products. Single-atom Fe doping also amplified the mass-specific activity to 399 mA mg⁻¹, which is much higher than that of pure Au nanoparticles (Fig. 5b–d). Jiao et al. added a Cd salt precursor during the fabrication of thiolate-protected Au nanoclusters (Auₓ(SR)ₓ) and achieved single-atomic Cd dispersed gold nanoclusters (Fig. 5e).⁸⁹ This regulation improved the FECO and current density to yield C₁ products (Fig. 5f). Unsatisfactorily, the modification of inexpensive metal elements on noble metal nanomaterials has not reduced catalyst cost, which restricts

![Fig. 5](https://example.com/f5.png)

(a) *In situ* AC-TEM monitoring of the Ag nanoparticle thermal dispersion phenomenon to fabricate Ag-SA/MnO₂. Reprinted with permission from ref. 87 Copyright 2021, Wiley-VCH. (b) FECO of the Fe₁/Au SAA and Au nanoparticle catalyst at different reduction potentials. (c) Mass activity of Fe₁/Au and Au nanoparticle catalyst at different reduction potentials in ECR. (d) Schematic diagram and EDS mapping figures of Fe₁/Au. Reprinted with permission from ref. 88 Copyright 2021, the American Chemical Society. (e) Schematic diagram of Cd₇Au₁₉ and Au₂₃ nanocluster catalysts. (f) FECO of the Cd₇Au₁₉ and Au₂₃ nanocluster catalysts. Reprinted with permission from ref. 89 Copyright 2021, Wiley-VCH.
their application in industry. Reducing the dosage of noble metals and anchoring noble metal atoms on transition metals is more beneficial for the practical application of SACs. Cao et al. used the molecular skeleton encapsulation method to synthesis Ni nanoclusters with single-dispersed Zn atoms. Although this method provides different types of active centers, it is also profitable to explore whether modifying the metal atoms on the nanocrystals/clusters could optimize the stability and activity performance in the ECR.

2.2.2. The perspective of nanocrystal/cluster-supported SACs. Currently, the synthesis of nanocrystal/cluster-supported SACs is still a novel field for effectively electrically reducing CO\textsubscript{2} to simple C\textsubscript{1} products. SACs suffer from a common problem, where the number of active sites is always too low to obtain high catalytic activity in many fields.\textsuperscript{90-95} Many studies indicate that SACs possess outstanding turnover frequency (TOF). However, their low yield (compared to commercial catalysts, at the same volume/quality) impedes the scale-up of the catalytic process. Nanocrystals and nanoclusters have high catalytic activities and specific nanosize effects. Transition metals are inexpensive with abundant reserves, and thus these elements can be effectively utilized to synthesize nanomaterials. Noble metals exhibit unique catalytic performances when downsized to the atomic level, promoting the catalytic selectivity of transition metals. Combining single noble metal atoms and transition metal nanomaterials can be employed to optimize the synthesis of efficient SACs. Inspired by the development of nanocrystals, the strategies for the synthesis of nanocrystal/cluster-supported SACs in the future mainly include “one-pot” and “multi-steps” methods. The “one-pot” method is easy for the preparation of SACs. However, more metal atoms are assembled in the interior, which causes the atomic utilization of noble metal to be lower than 100%. The “multi-step” method is normally employed to fabricate nanomaterials initially, and GR, CER and nanoparticle conversion reactions are employed to synthesize SACs. Noble metal atoms can be effectively anchored at the catalytic interface with the “multi-step” method, and thus the atomic utilization of noble metal elements is nearly 100% (Table 1).

3. SACs for deeply reduced C\textsubscript{1} products

Deeply reduced C\textsubscript{1} products initially include the 8e\textsuperscript{-}-transfer product methane and 6e\textsuperscript{-}-transfer product methanol. The catalytic mechanism is the same as that to obtain simple C\textsubscript{1} products. Metal–carbon or metal–oxygen intermediates are formed on the single atomic site, and there is no C–C coupling process. Alternatively, deeply reduced C\textsubscript{1} products need more electrons to reduce the 2e\textsuperscript{-} products (CO and FA). Currently, the consensus is that after the 2e\textsuperscript{-} reduction process, the *CO intermediate is preferentially reduced by protons/electrons to *CHO and *OCH intermediates rather than desorbed to produce CO (Fig. 6).\textsuperscript{13,34-36} Catalytically, the formation of the *CHO intermediate and *OCH is the rate-determination step in producing CH\textsubscript{4} and CH\textsubscript{3}OH. Therefore, regulation of the adsorption energy of the *CO intermediate is the key to catalyzing deeply reduced C\textsubscript{1} products.\textsuperscript{97}

Zheng et al. synthesized Cu single-dispersed CeO\textsubscript{2} structures via the impregnation and pyrolysis method,\textsuperscript{98} where Cu single atoms were trapped in the defects of CeO\textsubscript{2} through exchange of the valence of Ce\textsuperscript{4+} and Ce\textsuperscript{3+}, presenting the oxidation of +1 (Fig. 7a). In the ECR test with an H-cell, the Cu-SA/CeO\textsubscript{2} catalyst showed a notable FE\textsubscript{CH\textsubscript{4}} of 58% at −1.8 V vs. RHE, and the current density reached 70 mA cm\textsuperscript{-2} simultaneously (Fig. 7b). The synergistic effect of Cu single atoms and the substrate promotes the adsorption stability of the CO intermediate. Single-dispersed copper atoms suppress the occurrence of C–C coupling, and thus Cu-SA/CeO\textsubscript{2} catalysts can efficiently produce the deeply reduced product. Xin et al. used the coordination regulation and pyrolysis method for the synthesis of a carbon-supported SAC with a Zn-N\textsubscript{4}–C moiety.\textsuperscript{99} In 1 M KHCO\textsubscript{3}, this catalyst exhibited 83% FE\textsubscript{CH\textsubscript{4}} with a current density of 31.8 mA cm\textsuperscript{-2} (Fig. 7c and d) and stability for 35 h without an obviously decrease at a reduction potential of −1.8 V. In the 6e\textsuperscript{-}-transfer production of methanol, as the competition product of CH\textsubscript{4}, the rate-determination step (RDS) of the ECR process is the formation of the metal–oxygen intermediate, which requires the regulation the *CO adsorption energy. Han et al. synthesized a defected CuO structure with Sn single atoms using H\textsubscript{2} plasma-treated CuO to produce oxygen defects and trapped Sn atoms.\textsuperscript{100} This catalyst showed good selectivity for methanol in CO\textsubscript{2}-saturated ionic liquid (nearly 88.6% FE for methanol at −2.0 V). A peptide nitrile cobalt (CoPc) was also found to convert CO\textsubscript{2} to methanol after loading CNT. Liang et al. modified the peptide nitrile with amidogen (CoPc–NH\textsubscript{2}) and loaded CoPc–NH\textsubscript{2} on CNT the substrate.\textsuperscript{101} The FE for methanol could reach 44% at the beginning of ECR at −1.0 V vs. RHE and the largest partial current density of 10.6 mA cm\textsuperscript{-2} was observed for the CoPc–NH\textsubscript{2}/CNT catalyst. There is also a lack of research exploring the relation between SACs and the deeply reduced C\textsubscript{1} product. Most reports indicate that the single-dispersed atom sites block the transition state of C–C coupling. According to the Sabatier principle, regulating the adsorption energy of *CO on the active center can selectively produce 2e\textsuperscript{-}, 6e\textsuperscript{-} or 8e\textsuperscript{-} products. However, the deeply reduced C\textsubscript{1} products face the challenges of low catalytic activity, poor stability, and impaired conversion, which should be promoted in the future.

4. Development of ECR electrolyzers

In the development of ECR, the design of the electrolyzer has pushed the lab-scale electric reduction of CO\textsubscript{2} towards large-current and yield industrial-scale. The design of ECR electrolyzers mainly focuses on the following aspects: (1) effectively separating the products of the cathode and anode and preventing the reduction products of ECR oxidation at the anode. The crossover effect is deemed initially to decrease the reduction efficiency in the ECR. (2) Increasing the solubility and mass transfer rate of CO\textsubscript{2}. (3) Reducing the ohmic loss of the electrolyzer and the instantaneous volatility of local pH, which decrease the EE of electrolyzers. (4) Promoting the exchange and separation of products and continuous CO\textsubscript{2} conversion.
Table 1 Summary of the performance of different SACs in the ECR

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Mass loading (mg cm(^{-2}))</th>
<th>Electrolyte</th>
<th>Product</th>
<th>Reduction potential (V vs. RHE)</th>
<th>FE (%)</th>
<th>TOF (h(^{-1}))</th>
<th>Current density (mA cm(^{-2}))</th>
<th>Cell</th>
<th>Stability time [h]</th>
<th>Reference DOI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ag(<em>{1})-N(</em>{3})/PCNC</td>
<td>—</td>
<td>0.1 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.37</td>
<td>95</td>
<td>7.6</td>
<td>H cell 10</td>
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<td></td>
<td>10.1021/acsami.1c03638</td>
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<tr>
<td>Ni(<em>{1})-N(</em>{3})</td>
<td>0.53</td>
<td>0.5 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.80</td>
<td>96.8</td>
<td>27</td>
<td>H cell 10</td>
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<td></td>
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<tr>
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<td>0.1 M KHCO(_{3})</td>
<td>CO</td>
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<td>95.3</td>
<td>8.4</td>
<td>H cell 10</td>
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<td></td>
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<tr>
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<td>—</td>
<td>0.5 M KHCO(_{3})</td>
<td>HCOOH</td>
<td>−0.95</td>
<td>96</td>
<td>8.87</td>
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<td>Cd-N(_{3})</td>
<td>1.5</td>
<td>0.5 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.728</td>
<td>91.4</td>
<td>5</td>
<td>H cell 10</td>
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<tr>
<td>Fe(_{1})/graphitic N</td>
<td>—</td>
<td>0.1 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.60</td>
<td>97</td>
<td>6.87</td>
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<tr>
<td>Sb-N(_{2})</td>
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<td>0.5 M KHCO(_{3})</td>
<td>HCOOH</td>
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<td>4</td>
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<td>CoN(_{3})-C</td>
<td>—</td>
<td>0.2 M NaHCO(_{3})</td>
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<td>Bi-N(_{3})</td>
<td>—</td>
<td>0.1 M NaHCO(_{3})</td>
<td>CO</td>
<td>−0.5</td>
<td>97</td>
<td>3.9</td>
<td>H cell 4</td>
<td></td>
<td></td>
<td>10.1021/jacs.9b08259</td>
</tr>
<tr>
<td>Ni-SA/NC</td>
<td>3.00</td>
<td>1 M KOH</td>
<td>CO</td>
<td>1.06</td>
<td>96.9</td>
<td>213.2</td>
<td>Flow 20 (H cell)</td>
<td></td>
<td></td>
<td>10.1016/j.nanoen.2020.105689</td>
</tr>
<tr>
<td>Mn-N(_{1})-C</td>
<td>1.00</td>
<td>[Bmim]BF(<em>{4})/ CH(</em>{3})CN(<em>{2})(</em>{4})</td>
<td>HCOOH</td>
<td>1.06</td>
<td>98.8</td>
<td>22.7</td>
<td>H cell 20</td>
<td></td>
<td></td>
<td>10.1038/s14467-020-18143-y</td>
</tr>
<tr>
<td>Ni-Pc</td>
<td>0.05</td>
<td>1.0 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.61</td>
<td>99.5</td>
<td>400</td>
<td>MEO 40</td>
<td></td>
<td></td>
<td>10.1038/s14560-020-0667-9</td>
</tr>
<tr>
<td>Cu/Ni(OH)(_{2})</td>
<td>1</td>
<td>0.5 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.50</td>
<td>92</td>
<td>11 315~6</td>
<td>H cell 22</td>
<td></td>
<td></td>
<td>10.1126/sciadv.1701069</td>
</tr>
<tr>
<td>Cu(<em>{2})Sn(</em>{1})</td>
<td>0.57</td>
<td>0.5 M KHCO(_{3})</td>
<td>CO</td>
<td>−1.0</td>
<td>95.3</td>
<td>12.5</td>
<td>H cell 10</td>
<td></td>
<td></td>
<td>10.1021/acscatal.1c02556</td>
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<tr>
<td>Cu(<em>{3})Sn(</em>{1})</td>
<td>0.57</td>
<td>0.5 M KHCO(_{3})</td>
<td>HCOOH</td>
<td>−1.2</td>
<td>95.4</td>
<td>24.1</td>
<td>H cell 10</td>
<td></td>
<td></td>
<td>10.1100/adma.202005821</td>
</tr>
<tr>
<td>Pd(_{1})Ag</td>
<td>1</td>
<td>0.1 M KHCO(_{3})</td>
<td>HCOOH</td>
<td>−0.23</td>
<td>94</td>
<td>5</td>
<td>H cell 1h</td>
<td></td>
<td></td>
<td>10.1021/acscatal.1c02491</td>
</tr>
<tr>
<td>Au(<em>{1.5})Cd(</em>{2})(SR)(_{16})</td>
<td>0.02</td>
<td>0.5 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.60</td>
<td>90</td>
<td>45</td>
<td>H cell —</td>
<td></td>
<td></td>
<td>10.1021/acscatal.8b0104</td>
</tr>
<tr>
<td>Ag(<em>{1})/MnO(</em>{2})</td>
<td>—</td>
<td>0.5 M KHCO(_{3})</td>
<td>CO</td>
<td>−0.85</td>
<td>95.7</td>
<td>4</td>
<td>H cell 9</td>
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<td></td>
<td>10.1039/DEEO01486A</td>
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<tr>
<td>Fe(_{1})/Au</td>
<td>0.1</td>
<td>0.5M KHCO(_{3})</td>
<td>CO</td>
<td>−0.9</td>
<td>96.3</td>
<td>17.5</td>
<td>H cell 4</td>
<td></td>
<td></td>
<td>10.1021/acscatal.0c04291</td>
</tr>
<tr>
<td>Cu/CeO(_{2})</td>
<td>—</td>
<td>0.1 M KHCO(_{3})</td>
<td>CH(_{4})</td>
<td>−1.8 V</td>
<td>58</td>
<td>56</td>
<td>H cell 2</td>
<td></td>
<td></td>
<td>10.1021/acscatal.8b01014</td>
</tr>
<tr>
<td>Cu-N(_{1})</td>
<td>0.5</td>
<td>0.1 M KHCO(_{3})</td>
<td>CH(_{4})</td>
<td>−1.4</td>
<td>24.8</td>
<td>6.84</td>
<td>H cell 10</td>
<td></td>
<td>10 h</td>
<td>10.1021/acscenergylett.0c00018</td>
</tr>
<tr>
<td>Cu-N(_{1})</td>
<td>0.5</td>
<td>0.1 M KHCO(_{3})</td>
<td>CH(_{4})</td>
<td>−1.4</td>
<td>13.9</td>
<td>3.83</td>
<td>H cell 10</td>
<td></td>
<td>10 h</td>
<td>10.1021/acscenergylett.0c00018</td>
</tr>
<tr>
<td>Zn-N(_{4})</td>
<td>0.4</td>
<td>1 M KHCO(_{3})</td>
<td>CH(_{4})</td>
<td>−1.8 (V vs. SCE)</td>
<td>85%</td>
<td>39.9</td>
<td>H cell 35</td>
<td></td>
<td></td>
<td>10.1021/jacs.9b12111</td>
</tr>
</tbody>
</table>

According to the feedstock and separation function of electrolyzers, we can divide the electrolyzers into the batch, semi-batch, and continuous electrolyzers.\(^{102}\)

4.1. Batch or continuous electrolyzers

Batch and semi-batch electrolyzers mainly refer to single-compartment and double-membrane electrolyzers (H-cell). The products need to be separated from the reaction system after the reaction each time. The primary issue associated with single-compartment electrolyzers is the crossover effect, which oxidizes the C\(_{1}\) products to CO\(_{2}\) and reduces the yield. The H-cell, which separates the cathode and anode with an ionexchange membrane, can efficiently decrease this crossover effect. However, the anion-exchange membrane (AEM) also causes crossover in alkali electrolytes after long-time ECR catalysis.\(^{183}\) The H-cell possesses a simple operation process, which is suitable for lab-scale research. The CO\(_{2}\) at the cathode normally encounters low saturation, which reduces the contact between the reactant and active catalytic centers. The inferior fluidity of electrolytes induces a poor mass transfer rate and restricts the increase in the current density of ECR. The OER product at the anode is mixed with CO\(_{2}\) or CO, which decreases the purity of the gaseous C\(_{1}\) product, and the waste O\(_{2}\) results in a decrease in the EE of the electrolyzer.\(^{184}\)

Zhang et al. proposed a solution to this problem. They used Cl\(^{-}\) to react at the anode and collected NaClO product.\(^{185}\)
CO2 reactant was electrically reduced on the Fe-Nx moiety SACs to produce CO and Cl− oxidized on RuO2 to produce ClO−, where the whole electrolyzer possessed 96% FE CO and 82% EE, significantly improving the EE compared to the OER wasted on the IrO2 anode. However, batch and semi-batch electrolysers are unsuitable for the efficient and continuous utilization of CO2 feedstock. Accordingly, the establishment of constant feeding and separation devices is the development trend of electrolysers. Continuous electrolysers should accelerate the fluidity of electrolytes, and consecutive feeding of CO2 can also facilitate the mass transfer and reduction in current density. The flow cell is a type of ordinary continuous electrolyzer, which can be divided into liquid-fed, vapor-fed, and microfluidic flow cells.106–108 The liquid-fed flow cell cathode and anode are separated by an electrolyte and often include a liquid cell and gas–liquid cell. Both humidified CO2 and dry CO2 can generate the reduction products on the cathode (Fig. 8a). The mass transfer is faster than batch electrolysers. However, the considerable distance between the cathode and anode causes ohmic loss, and higher partial pressure of CO2 produces a decrease in conductivity near the electrode. Thomas F. Jar­amillo’s group designed a liquid-fed flow cell with dry CO2 as the feedstock,109 which has a liquid volume of 8 mL and a gas volume of 3 mL and can obtain 25 reduction products in the ECR. This forms the prototype for the flow cell design (Fig. 8b). One of the potential solutions for reducing the ohmic loss is to decrease the distance between both electrodes. The vapor-fed flow cell is characterized by a minimum distance between electrode plates with lower ohmic loss, but the requirements for the separator membrane are much more demanding. Vapor-fed flow cells initially use humidified CO2 as the feedstock and have the desired mass-transfer rate and current density in the ECR. However, the ion-exchange membrane needs to withstand extreme current conditions. To date, the exploration of separator membranes with high intensity and ion selectivity is still a considerable challenge. The liquid-products in vapor-fed flow cells (Fig. 8c) diffuse slowly, which can cause the separator and SACs to be deactivated. Thus, third-generation microfluidic flow cells were designed (Fig. 8d), where there is a thin layer of flowing liquid between the cathode and the anode to transfer the liquid products. Dry or humidified CO2 rapidly contacts the SACs on the gas diffusion electrode (GDE), and the accelerated mass transfer also enhances the current density. The microfluidic flow cell is widely used in ECR tests, where dry CO2 contacts with superhydrophobic gas diffusion electrodes to produce C1 products. The advantage is that the gas and liquid can be partially separated by different electrolytic cells. However, achieving a uniform SAC coating on the GDE and the hydrophilic leakage of the GDE caused by the reduction voltage and alkaline environment are still challenges for further exploration.110

4.2. MEA electrolysers and multi-assembly electrolysers

Accompanying the development of electrolysers, the current density of ECR has a noticeable improvement, gradually increasing the production scale of ECR from lab-scale to pilot-plants. However, it is difficult to achieve an ideal single-pass conversion rate of CO2 with a traditional single electrolyzer. C. Janaky et al. designed a multi-assembly flow cell based on the microfluidic flow cell, which improved the single-pass conversion by extending the transfer distance of CO2 molecules.111 As shown in Fig. 9a, dry CO2 was primarily treated to obtain humidity and pressure CO2 feedstock. The feedstock was bubbled into the multi-assembly flow cell. The current density for the production of CO products reached 250 mA cm−2 under normal pressure. In the 10 bar-pressurized ECR test, the current density reached up to 300 mA cm−2 with 95% FE CO and the CO2 single-pass conversion of this multi-assembly flow cell was nearly 40%. The CO product demonstrates its potential for the industrialization of the ECR. A further development of the electrolyzer is membrane electrode assembly (MEA) devices.
rubber membrane (APEM, PEM, AEM, or BPM), and anode catalysts for the OER (IrO₂, RuO₂, etc.). Pre-treated humidified CO₂ will produce and flow the products out on the same side of the GDE. The single-pass conversion of CO₂ can also be increased through the parallel multilayer MEA device (Fig. 9c). Compared to the traditional flow cell systems, the MEA electrolyzer is a complete battery system, which can achieve a higher ECR current density at a lower voltage. The battery assembled using MEA can obtain separated C₁ and O₂ products, improving the EE and FE of the process. This allows for a higher mass content of hydrogen results in a higher heat of combustion. H₂ is ideal energy but suffers from storage and transportation issues. The U.S. Department of Energy Laboratory proposes that the most valuable reduction product from an energy perspective is methane.¹¹³ Tao’s group reported that the ECR products from carbon utilization follow the order of CH₄, formic acid (FA), CO, C₂H₄, EtOH, and MeOH.¹¹⁴ However, combined with the development of ECR technology, the order of the catalytic performance for the C₁–C₃ products is CO, MeOH, CH₄, FA, ethylene, and acetic acid. The deeply reduced C₁ products methane and methanol are not ideal for scale-up application in industrial ECR, which have lower utilization of electrons (8e⁻ or 6e⁻ for one molecular product), lower conversion, and the mass yield of CH₄ is only 36%.

In contrast, the mass yield of CH₄ is 105% for FA in the ECR process. Assessment of the separation cost of the product and the sensitivity to market fluctuations shows that the most promising C₁ product in the industrial ECR is CO, followed by FA. Sargent’s group summarized the factors that TEA needs to consider from a laboratory perspective,¹¹⁵ including market size, price of products, battery EE (mainly for C₁ products), chemical efficiency (mainly for C₂ products), electrolyzer cost, operational cost, and separation cost of the whole process. Alternatively, as ECR technology becomes more mature, electricity prices, transportation costs, and CO₂ capture prices have become crucial factors affecting the economic benefits of the ECR industry.¹¹⁶ Establishing an accurate TEA method can predict the possibility of carbon economic development.

5. Techno-economic assessment of C₁ and key indicators for SAC performance

Techno-economic assessment (TEA) is typically used to evaluate the factors that affect the scale of the ECR through systematically analyzing the complete life cycle or half life cycle process of ECR industrialization. There are many considerations for the value of ECR products. From the perspective of energy storage, a higher mass content of hydrogen results in a higher heat of combustion. H₂ is ideal energy but suffers from storage and transportation issues. The U.S. Department of Energy Laboratory proposes that the most valuable reduction product from an energy perspective is methane.¹¹³ Tao’s group reported that the ECR products from carbon utilization follow the order of CH₄, formic acid (FA), CO, C₂H₄, EtOH, and MeOH.¹¹⁴ However, combined with the development of ECR technology, the order of the catalytic performance for the C₁–C₃ products is CO, MeOH, CH₄, FA, ethylene, and acetic acid. The deeply reduced C₁ products methane and methanol are not ideal for scale-up application in industrial ECR, which have lower utilization of electrons (8e⁻ or 6e⁻ for one molecular product), lower conversion, and the mass yield of CH₄ is only 36%.

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5.1. Assessment methods for industrial ECR

Currently, the feasibility methods for evaluating the prospects of the ECR industry include gross-margin analysis and single-variable sensitivity analysis. Paul J. A. Kenis et al. used the gross-margin model to analyze the economic feasibility of the ECR. Gross-margin analyzes the ratio of the net benefit generated by the ECR process to this revenue.²⁹ This value ordinarily should be higher than a fixed value (as shown in Fig. 10, eqn (1), (3), and (4), where the average of the commodity-chemicals
The investment of the whole life cycle includes the cost of the electrolysis cell, balance of the plant, catalyst and electrolyte, and electricity ($\lambda$ in Fig. 10 eqn (2), all values are state-of-the-art data). The advantage of this assessment method is that the calculation process is relatively simple. Gross-margin analysis is suitable for the rational analysis of a specific catalyst or technology process. However, in many ECR reactions, the current density, EE, FE, and peak and valley electricity prices are all fluctuating parameters, which can hardly be confirmed with specific values. The solution for gross-margin analysis is estimated using average values. Many research groups use single-variable sensitivity analysis to evaluate whether a C1 product can be profitable within specific parameters. Single-variable sensitivity analysis is more appropriate, given that it does not need to consider all the factors in the whole life cycle. Sargent et al. employed this method to analyze the possibility of the one-step ECR industrial production of ethylene, which is hopeless using current technology. When they estimated reducing the CO feedstocks to produce C2H4, they were surprised to find this catalytic process has high potential for the industrial production of C2. Jiao’s group performed a sensitivity analysis of the cost to produce C1 and C2 in industry (CO, FA, C2H4, and EtOH). Through this TEA research, they found the most susceptible cost for the industrial ECR production of CO is the price of electricity, followed by the single-pass conversion rate, which should be higher than 40%. Combined with the increased ohmic loss caused by high voltage, the current density also has a specific range (500–700 mA cm$^{-2}$). After, the service life cycle of MEA, CO2 capture price, product price, and FEO will affect the profits for the CO product in industrial ECR. The most susceptible cost in the industrial ECR production of FA is the current density (200–300 mA cm$^{-2}$) followed by the price of electricity and the service life cycle of MEA, etc. In contrast, the cost of the industrial ECR production of the C2 products of ethylene and ethanol can still achieve better market values with the state-of-the-art parameters tested with the advanced parallel electrolytic cells and high-efficient catalysts. Sensitivity analysis requires a sizeable statistical database and more calculation models. This TEA method is usually used to estimate the acceptance range of a specific factor affected by other factors (such as product cost and the origin of CO2) (Fig. 11).

5.2. Technical benchmarks related to ECR industrialization

5.2.1. The benchmarks of plant operating cost and electrolyzer. (1) Peak and valley electricity prices and CO2 capture techniques affect the cost of industrial ECR. The product value depends on the current floating market price. When estimating the possibility of applying SACs for industrial ECR, we can generally use the peak electricity price and optimized CO2 capture price (such as $40 per ton CO2 and $0.068 kW h$^{-1}$ electricity in 2017). For reduction products with less electrons (CO) in the ECR, the minimum selling price will be susceptible to the cost of CO2 capture. Currently, the CO2 capture cost is low enough to profit from the CO products. (2) Evaluating the economic parameters related to the electrolytic cell is an essential part of the ECR process. The cost of the electrolyzers has gradually decreased after the emergence of series electrolyzers. The MEA electrolyzer is designed to continuously feed CO2 and efficiently convert it to C1 with a high current density. However, MEA systems are generally affected by the surface hydrophilization caused by GDE leakage, which results in the side-reaction of the HER occurring. (3) Stability is still an important issue in industrial ECR. Paul J. A. Kenis and others used the gross-margin model to prove that the battery stability time is positively correlated with the ultimate benefit. There is a large coefficient factor. The conventionally used gross-margin model usually requires an ECR conversion of 4000 h to have a 30% net profit in the full lifecycle. However, the typical stability time for lab-scale catalysts is still less than 100 h. (4) Another influence factor consumable is the separator membrane, which is divided into PEM, AEM, and BPM. Different membranes greatly influence the suppression of
crossover problems and the diffusion of anions and organic products. According to the laboratory data statistics, the use of the AEM membrane can decrease the cost of industrial ECR for CO products, and the BPM membrane is more suitable for FA products.

5.2.2. The benchmarks of catalyst performance. Besides adjusting the factors in the ECR electrolyzer, to determine the feasibility of SACs in the ECR, it is necessary to consider the current density, single-pass conversion rate, FE, stabilization time (retaining 90% FE C1 stability), EE, and other factors. (1) Although an increase in cell potential is beneficial for the current density and favorable for the yield of C1, a too high cell potential will increase the cost. Ohmic loss aggravated at high current density reduces the stability of the battery. (2) The single-pass conversion ordinarily needs to be higher than 40% for CO and 80% for FA, which can be solved through parallel electrolyzers and an increase in electrode area for SACs with high FE CO or FE FA close to 100%. (3) In addition to improving the stability in the ECR, reducing the cost of the electrode (IrO2) and the GDE (catalyst) is also essential to achieving profits. High metal atom utilization of SACs can significantly reduce the cost of catalysts during the whole ECR process. (4) The EE can be promoted by regulating the FE and cell potential. The assessment of the ECR catalytic process shows there are still many challenges for SACs, such as the current density and FE, which are unsuitable for the standard of industrial ECR. The stability should be optimized to decrease the cost of the full lifecycle. Alternatively, industrial ECR is profitable with current SAC research progress in this field.121–123

6. Conclusion and outlook

This perspective summarized the methods for the synthesis of SACs with outstanding ECR performances for C1 products and briefly described the development of electrolyzers. Combining the ECR technical situation and evaluation standard of catalytic performance, we presented the benchmarks in the complete lifecycle of the industrial ECR and summarized the economic feasibility of applying SACs in the industrial electric conversion of CO2 to C1 with the guidance of previous techno-economic assessment. Efficient SACs should be designed with appropriate types of metal elements to achieve a higher current density, reduce the concentration of noble metals, facilitate stability during the ECR process, and maintain super-hydrophobicity in the alkaline electrolyte. As the conclusion in this perspective, Mn, Fe, Co, Ni, Cd, Ag, and Bi as the active center will produce CO during the ECR process. Also, the single-metal atoms in the main group prefer to produce FA in the ECR. The difference in the electronic structure of the active center caused by the metal species and the atomic coordination environment are the intrinsic factors that affect the ECR reaction. The adsorption energy of CO2 and various ECR products of the above-mentioned SACs directly affect the selectivity of the ECR process. Carbon-supported SACs have been researched in depth, but there are still many problems encountered, as follows: (1) the cost of catalyst synthesis; (2) the activity of the catalyst; (3) the uniformity and reproducibility of the catalyst; and (4) the difference in the microscopic atomic structure affect the selectivity of C1 products.

The main problem of the cost of SACs can be solved by reducing the cost of the substrate. ZIF-8 is a well-known precursor of carbon-supported SACs, which will consume the valuable ligands during the carbonization process. Research on the nitrogen species, the electronic structure of M1–N–C from ZIF-8, and the effectiveness of residual zinc will help substitute the carbon substrate with low-cost carbon materials. Compared with traditional carbon-supported SACs, nanocrystal/cluster-supported SACs have higher catalytic activity, more uniform microscopic atomic structure, and increased catalytic current density. However, their cost is higher than traditional carbon-supported SACs, which urge researchers to simplify the synthetic process for nanocrystals/clusters. Another crucial problem of SACs is the number of active centers. Densely active centers will improve the conversion and accelerate the catalytic process. The primary method to get dense metal SACs is the “bottom-up” strategy, which assembles small coordination compounds with high concentrations and carbonizes polymers to synthesize dense M–N–C SACs. Correspondingly, the “top-down” strategy may be used more frequently in the future. This indicates the conversion of nanoparticles or high-ordered single-atomic site dispersed metals to coordination-disordered SACs. The research on SACs for the ECR is mainly focused on the H-cell, which is not conducive to scaling-up in industry. Inspired by the development history of electrolyzers, evaluating the ECR performance (current density, FE, cycling stability, and single-pass conversion) of SACs with flow cell or multi-parallel MEA electrolyzers is a crucial step to facilitate the C1 synthesis scale from lab-scale to pilot-plant.

Under the target of carbon neutrality formulated by each country, the aim is to convert the originally economically unfavorable ECR process to a financially profitable industry due to the decrease in electricity prices and CO2 capture costs. Without considering the benefits of the policy carbon economy, both the gross-margin model and single-variable sensitivity analysis can be used to evaluate whether the industrial ECR process is profitable. Gross-margin model analysis is a quick and convenient judge for the profitability of a specific catalyst, and this can help analyze the availability of synthesized catalysts. Single-variable sensitivity analysis requires enormous statistical analysis. The deeply reduced C1 products (CH4 and CH3OH) are unfavorable in the entire lifecycle of industrial ECR at the current technicity. Conversely, simple C1 products (CO and FA) have more potential to yield a profit and realize a net-negative carbon economy. For a specific process, the factors and conditions that generate benefits can be analyzed to help understand the performance indicators that need to be regulated in the catalytic process (Fig. 11). Herein, we also discussed the feasibility of the electrochemical conversion of CO2 to CO and FA by SACs with guidance from the latest TEA progress. Consequently, some catalytic performance indicators (current density, FE, and EE) need to be optimized in the future.

In conclusion, SACs have improved the relevant catalytic indicators in the industrial ECR economic analysis, which are beneficial due to their high atomic utilization and specific
selectivity of products. Increasingly optimized technology and efficient SACs have facilitated the simple C_1 products (CO, FA) from the laboratory to scaled-up production in the industry. Although relevant factories at the domestic and foreign levels have already developed appropriate industrialized ECR devices, the use of SACs can still further help in process optimization. The three crucial issues are membranes, catalysts, and electrolyzers, which need to be optimized to provide a substantial basis for the industrial conversion of carbon. The era when SACs are used as critical catalysts in the industrial ECR is upcoming.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Notes and references


12 A. Han, Z. Zhang, X. Li, D. Wang and Y. Li, Small Methods, 2020, 4, 2000248.


