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A review on electrochemical  $CO_2$ -to- $CH_4$  conversion for 1003854E sustainable energy future: From electrocatalysts to electrolyzers

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#### **Abstract**

The electrochemical reduction of carbon dioxide (CO<sub>2</sub>) to methane (CH<sub>4</sub>) offers a promising route to renewable fuels and carbon circularity, addressing urgent climate and energy challenges. However, key bottlenecks such as limited selectivity, sluggish reaction kinetics, and insufficient long-term stability still hinder the practical deployment of this reaction and technology. Fundamental research has uncovered promising electrocatalysts and mechanistic insights to overcome these limitations, yet translating these advances into scalable industrial solution remains a major challenge. This review addresses this critical gap by providing a comprehensive and focused overview of the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion, from fundamental reaction mechanisms to system-level implementation. This work systematically analyzes the most selective and active electrocatalysts developed to date, elucidating key design principle that govern CH<sub>4</sub> production. In addition, we assess the evolution of CO<sub>2</sub> electrolyzers tailored for CH<sub>4</sub>, comparing device configurations, operational strategies, and levels of technological maturity. Techno-economic evaluations are also integrated to identify bottlenecks and realistic near-term implementation scenarios. As demand for green CH<sub>4</sub> rises at a pace that outstrips conventional CH<sub>4</sub> growth, this technology emerges as a timely solution to decarbonize hard-to-abate sectors using renewable electricity.

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1. Introduction

electrochemical reduction<sup>22–24</sup>.

## The average atmospheric carbon dioxide (CO<sub>2</sub>) concentration in 2024 has once again reached an unprecedented level of 422 ppm, surpassing any historical human records. Projections indicate that the 2025 annual average CO<sub>2</sub> concentration is expected to exceed a peak value of 427 ppm<sup>1</sup> for the first time in the last 800,000 years<sup>2</sup>. This trend is anticipated to continue in an upward trajectory in the coming years, and this high concentration of CO<sub>2</sub> in the atmosphere is already contributing to a substantial rise in global temperature<sup>3,4</sup>, ocean acidification<sup>5</sup> and the disruption of the carbon cycle<sup>6</sup>. This forces scientists to intensify their efforts to explore and develop environmentally-friendly ways of energy production 7, with a focus on technologies that should aim to either reduce or (ideally) fully replace the use of conventional fossil fuel-based methods 8,9. Beyond this urgent need to foster the broad implementation of such renewable energy sources, sort- to mid-term solutions require a decrease of non-abatable CO<sub>2</sub>-emissions or even active strategies to capture atmospheric CO<sub>2</sub>. To strengthen the economic viability of such approaches, the latter CO<sub>2</sub>-capture<sup>10,11</sup> would ideally be followed by the conversion of the carbon dioxide into value-added chemicals useful for industrial applications<sup>12,13</sup>. The latter CO<sub>2</sub>-conversion encompasses different methods, including chemical conversion<sup>14–16</sup>, biological transformation <sup>17,18</sup>, photocatalytic reduction<sup>19–21</sup> and

These CO<sub>2</sub>-conversion methods are plagued by well-known drawbacks, such as high energy requirements, low conversion rates, etc. By comparison, the electrochemical CO<sub>2</sub> reduction reaction (ECO<sub>2</sub>RR) has recently emerged as a highly promising pathway due to the possibility to couple it with renewable electricity, as pictured in Figure 1<sup>25,26</sup>. Furthermore, some of the products of the ECO<sub>2</sub>RR (e.g., CO, C<sub>2</sub>H<sub>4</sub>) can serve as valuable fuels or chemicals that integrate perfectly into existing industrial processes. Moreover, this electrochemical reaction is well-known for its controllable and mild reaction conditions (i.e., close-to-atmospheric temperatures

and pressures). Nonetheless, there are still notable challenges associated with the ECO and a straight including their limited, operative current density (often << 1 A·cm<sup>-2</sup>), its poor selectivity towards producing a single desired product, and its lack of stability<sup>27–29</sup>.

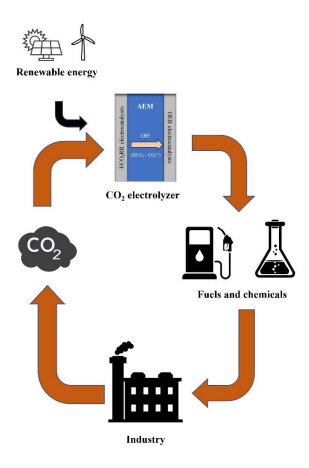


Figure 1: Illustration of the closed carbon cycle enabled by a  $CO_2$ -electrolyzer (in this example, relying on the use of an anion exchange membrane, AEM, and with the  $O_2$ -evolution reaction (OER) as the anodic counter-reaction) powered by renewable energy sources and that converts captured or emitted  $CO_2$  into chemicals or fuels for their direct usage in the industrial sector.

In an industrial context, two different application scenarios are envisaged for the ECO<sub>2</sub>RR. Firstly, the reaction can be conducted within a standalone electrochemical system in which the resulting products are stored for subsequent use in other industries. Alternatively, the CO<sub>2</sub> can be electrochemically converted into a value-added product that is directly coupled to a second reactor, thus avoiding (or at least mitigating) additional costs associated with product storage and transportation. This second approach is more demanding in terms of performance because

the resulting products must be of extremely high purity. Since they are fed directly into an off  $\frac{1}{2}$  industrial process, they need to meet strict quality standards to ensure efficiency and prevent any negative impact on the subsequent production state. This represents an important bottleneck in ECO<sub>2</sub>RR, as many materials electrochemically reduce CO<sub>2</sub> into a wide variety of carbon-containing chemicals and/or  $H_2$  with a poor selectivity<sup>30–32</sup>.

One of these possible ECO<sub>2</sub>RR-products is methane (CH<sub>4</sub>), which has otherwise found extensive use as an energy source in the fields of low-pollution power generation, liquidnatural-gas vehicles and so on<sup>30</sup>. CH<sub>4</sub> is particularly valuable where existing infrastructure for natural gas storage, distribution and consumption can be leveraged, 23 since 70-90% of this natural gas (which is mainly obtained from oil wells and coal beds<sup>33</sup>) is composed of CH<sub>4</sub><sup>34</sup>. As an extensive storage and distribution network already exists for natural gas, the development of electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion would enable a rapid and widespread distribution of net-zero-carbon energy services<sup>35,36</sup>. The global high purity methane gas market was valued at USD 101 billion in 2020 and is projected to reach USD 153 billion by 2025<sup>37</sup>, which is by far superior to other ECO<sub>2</sub>RR-derived products, such as CO (USD 3.3 billion in 2022 and projected to USD 4.5 by 2030)<sup>38</sup> or HCOOH (USD 2.1 billion in 2023 and projected to USD 3.8 billion by 2030)<sup>39</sup>. Nevertheless, several challenges impede the widespread implementation of electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion, including the low faradic efficiencies (FEs), sluggish kinetics and poor durability exhibited by most catalysts and reactor configurations. Moreover, the electrochemical production of CH<sub>4</sub> faces economic hurdles, as this gas possesses the lowest average market price among all CO<sub>2</sub>-derived products<sup>40</sup>. Specifically, the cost of CH<sub>4</sub> from the electrochemical reduction of CO<sub>2</sub> is expected to stand at approximately 2.5 - 10 \$\cdot \kg^{-1}\$, which is still higher than traditional CH<sub>4</sub>-synthesis methods (0.2) - 0.5 \$ kg<sup>-1</sup>)<sup>40</sup>. In this regards, technoeconomic analyses have suggested that the price of ECO<sub>2</sub>RR-derived CH<sub>4</sub> must be reduced to at least ≈1.0 \$ kg<sup>-1</sup> to effectively compete with

traditional CH<sub>4</sub>-production procedures<sup>40</sup>. Today, these elevated costs represent a major battlet color and consolidating the past achievements while addressing the remaining technoeconomic challenges.

With this motivation, in this review we comprehensively examine the advancements in electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion, focusing on the nature of electrocatalysts and the design of electrochemical cells and devices. We first discuss the fundamental principles of the ECO<sub>2</sub>RR, including reduction mechanisms and electric double layer (EDL) effects. Subsequently we delve deeply into the recent developments concerning electrocatalysts, operating conditions and CO<sub>2</sub> electrolyzers. Finally, we offer insights into the future prospects of ECO<sub>2</sub>RR and highlight the challenges that must be addressed to enhance the efficiency and effectiveness of this electrochemical reaction to accomplish technoeconomic requirements.

#### 2. Fundamentals of ECO<sub>2</sub>RR

#### 2.1 Thermodynamics and reaction mechanism

 $CO_2$  is a thermodynamically stable molecule with linear geometry and a dissociation energy of the C=O bond of  $\approx$ 750 kJ·mol<sup>-130,41</sup>. This high energy requirement suggests a substantial activation energy for direct C=O bond dissociation. In the  $CO_2$  reduction mechanism, this activation must be considered as the first step, where the linear geometry of the pristine molecule is transformed into a bent configuration in which the C=O bond is weakened through the formation of chemical bonds between the  $CO_2$  molecules and the active sites<sup>42</sup>. After  $CO_2$ -adsorption, the  $ECO_2RR$  goes through a series of reaction steps at the surface of the active

phase involving the cleavage of C-O bonds, C-C coupling, or C-H formation, and multiple colling electron transfer processes (e.g., 2 electrons for CO, 6 electrons for CH<sub>3</sub>OH) and eventually leading to different products<sup>43,44</sup>.

In theory, this variety of ECO<sub>2</sub>R-reactions generally exhibit their thermodynamic equilibrium potentials at values close to 0.0 V vs. the reversible hydrogen electrode (RHE), as shown in Table 1 which also includes the competing (and undesired) H<sub>2</sub>-evolution reaction (HER). All potentials were calculated based on the Gibbs free energy of reaction, using gas-phase thermochemistry data and Henry's law data for aqueous products<sup>24</sup>. Beyond these thermodynamic considerations, one must bear in mind that large overpotential are often required for achieving sufficiently high current densities due to the sluggish kinetics of these reactions<sup>45–47</sup>. Furthermore, the ECO<sub>2</sub>RR is highly sensitive to the applied potential, as the selectivity towards a given product usually follows the tendency of increasing until a maximum value, to then decrease when applying more negative potentials.

Table 1: Electrochemical CO<sub>2</sub> reduction potentials versus standard hydrogen electrode (SHE) at pH 7. Hydrogen evolution reaction is included for comparison purposes.

	Thermodynamic
Electrochemical Reaction	potential / V vs
	RHE
$CO_2 + 2H^+ + 2e^- \rightarrow HCOOH$	-0.12
$CO_2 + 2H^+ + 2e^- \rightarrow CO + H_2O$	-0.10
$CO_2 + 6H^+ + 6e^- \rightarrow CH_3OH + H_2O$	0.03
$CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O$	0.17
$2CO_2 + 12H^+ + 12e^- \rightarrow C_2H_4 + 4H_2O$	0.08
$2H^+ + 2e^- \rightarrow H_2$	0.00

**Figure 2:** Proposed mechanism for the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion and competitive reactions towards other CO<sub>2</sub>-derived products.

Concerning the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> reduction reaction, it has been proposed that CH<sub>4</sub> formation undergoes two stages schematized in Figure 2: the first one is the initial formation of a CO intermediate adsorbed to the catalyst surface (\*CO) and one water molecule, and the second stage is the subsequent hydrogenation of the \*CO into CH<sub>4</sub> <sup>42,48</sup>. The key point in the selectivity towards CH<sub>4</sub> formation appears to be related to the binding energy of the \*CO. Namely, if the binding energy of this intermediate is too weak, most of the \*CO will desorb as a carbon monoxide (CO), whereas a moderate binding energy between the active site and the C atom of \*CO intermediate is mandatory to reach high selectivity towards CH<sub>4</sub> formation. Computational studies have demonstrated that this hydrogenation can take place through (*a*) the hydrogenation of the C atom forming a so-called \*CHO intermediate, or (*b*) the weakening the C=O bond through the hydrogenation of the O atom, to form a \*C-OH intermediate (see Figure 2). The rate determining step in both mechanisms is the formation of the first

hydrogenation intermediate (i.e., \*CHO or \*COH). Here is worth pointing out that the \*COHICLE Online of the CH<sub>4</sub> or CH<sub>3</sub>OH, while \*CHO can also proceed to form C<sub>2+</sub> compounds. In pathway (*b*), water may be desorbed and lead to the ECO<sub>2</sub>RR to proceed through a \*C intermediate. After electron-proton pair additions, this intermediate undergoes reduction reactions towards \*CH, \*CH<sub>2</sub> and so on<sup>49</sup>.

There are also several competitive reactions that can tune the selectivity towards CH<sub>4</sub> formation from CO<sub>2</sub>. In mechanism (a), the selectivity for CH<sub>4</sub> generation over CH<sub>3</sub>OH is determined by the last reduction step, where \*CH<sub>2</sub>OH can undergo two different pathways leading to the hydrogenation of either the oxygen or the carbon atoms. At the same time, it has been demonstrated that C<sub>2</sub>H<sub>4</sub> and CH<sub>4</sub> generation share similar pathways until the \*CHO intermediate<sup>48</sup>. If C-C coupling reactions are favored at the electrocatalyst's surface, the formation of an OHC\* - \*CHO intermediate will predominate. Thus, in the search of highly selective CO<sub>2</sub>-to-CH<sub>4</sub> electrocatalysts, the active sites should favor the completion of the CH<sub>4</sub> pathway avoiding C-C coupling.

The HER is the main competing reaction in the cathodic reduction of CO<sub>2</sub> in aqueous electrolytes, since it features a similar equilibrium potential (Table 1) and is particularly facile to catalyze in acid solutions<sup>47</sup>. The first reaction step involves the reduction of protons or H<sub>2</sub>O molecules in the electrolyte to produce adsorbed H (H\*, known as the Volmer step) that is, subsequently further reduced to generate H<sub>2</sub> molecules. If the catalyst's active sites possess a strong H\* adsorption, the HER will predominate and ECO<sub>2</sub>RR will be suppressed, and thus a good ECO<sub>2</sub>RR electrocatalyst must feature a weak binding strength towards H\* and a moderate CO<sub>2</sub> activation.

#### 2.2 Electric double layer

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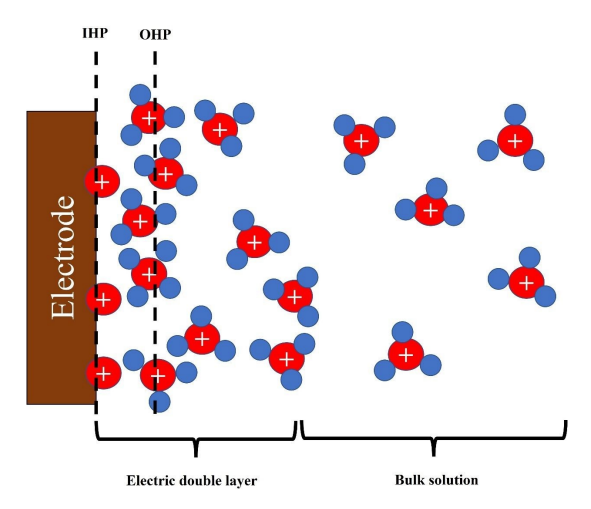
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Although the electrolyte is sometimes considered chemically inert in many electrochemical reactions, the identity of the ions present in it is known to affect the reaction rates and product selectivity of many electrochemical reactions in impactful ways<sup>50</sup>. This is in part due to the electric double layer (EDL) that forms when a solution containing ions is in contact with a solid surface that holds stationary charges<sup>51</sup>. The EDL is the result of the attractive and repulsive electrostatic interactions between the ions in the electrolyte and the charged surface, which creates a locally varied electric potential. The understanding of the EDL in electrochemical reactions has significantly advanced over the decades. The proposed theories include (i) the blocking of the active sites by ions of the electrolyte, (ii) the redistribution of the potential drop in the double layer, which heavily affects the driving force for electron transfer, (iii) the interaction of the interfacial electric field with the electric dipole moments and polarizabilities of adsorbed intermediates, (iv) chemical interaction between ions and reaction intermediates, (v) the buffering of the interfacial pH by hydrated ions, and (vi) the alteration of the interfacial water structure<sup>52</sup>. To fully understand the EDL, it is worth mentioning that all these theoretical concepts are not strictly separable but interrelated.

For cathodic reactions like the ECO<sub>2</sub>RR, the electrode is negatively polarized and its negative charge translates into an excess of cations and a depletion of anions in the vicinity of its surface (Figure 3)<sup>53</sup>. The outer Helmholtz plane (OHP) represents the closest approach of hydrated ions to the surface, while the inner Helmholtz plane (IHP) is formed by the ions that are chemically adsorbed onto the surface of the electrode.

Many studies have reported effects of the electrolyte's cation(s) on the ECO<sub>2</sub>RR in aqueous media. One illustrative example is what is observed by substituting Li<sup>+</sup> with Cs<sup>+</sup>, where the latter cationic species leads to a substantial enhancement of the ethylene selectivity during ECO<sub>2</sub>RR on Cu(100) and Cu(111)-oriented thin films <sup>54</sup>. Such cationic effect is mainly related to the tendency of cations to chemically or physically adsorb on the electrode surface, which

is governed by the reaction energetics and hydration shell of the cations<sup>55</sup>. The hydration hydration shell of the cations<sup>55</sup>. The hydration hyd



**Figure 3:** Illustration of the inner Helmholtz plane (IHP) and outer Helmholtz plane (OHP) in the electric double layer. Red spheres represent the cation, blue spheres represent water molecules acting as a solvation sphere. Anions in the solution were not incorporated for the sake of clarity.

#### 3 Electrocatalyst Design

Electrocatalysts are essential for the ECO<sub>2</sub>RR, since they speed up the rate of this high high had a complex and kinetically demanding electrochemical reaction. These electrocatalysts hold the potential to substantially improve and modulate the selectivity and kinetics of the electrochemical CO<sub>2</sub> reduction, all while preserving their intrinsic properties. The urgent need to address climate change and reduce anthropogenic CO<sub>2</sub> has pointed out the key role of electrocatalysts design for ECO<sub>2</sub>RR in the coming years.

In the 80s and 90s, ECO<sub>2</sub>RR-electrocatalysts were classified into different groups according to the metal phase and resulting product selectivity<sup>59</sup>. However, it is currently known that activity and selectivity do not only depend on the nature of the metal phase, but also on complementary properties such as the electronic configuration, catalysts support, surface chemistry, morphology or electric conductivity<sup>59–62</sup>. It is proven that tuning the electronic properties, composition and morphology of the electrocatalysts can significantly modify the density and turn-over frequency of the sites/phases active for the ECO<sub>2</sub>RR.

Thus, in this section we present and discuss recent developments on ECO<sub>2</sub>RR electrocatalysts based on their initial composition, to then delve into the influence of different electronic and structural properties that can tune the ECO<sub>2</sub>RR performance, with especial emphasis in the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion. To provide a quick overview for the readers, we summarize in Table 2 the most active catalyst reported to date, highlighting key electrochemical parameters.

#### 3.2 Metal electrocatalysts

A simple but vague classification can be made by considering the nature of the metallic phase for bulk materials<sup>47</sup>. Generally speaking, most of the transition metals electrocatalyze the HER more easily than the ECO<sub>2</sub>RR in aqueous media, rendering them useless for practical application when seeking CH<sub>4</sub> formation. This HER-selective materials group includes Ti and

Pt, among others. Complementarily, some transition metals exhibit higher ECO<sub>2</sub>RR<sub>O1</sub>vs. HER ricle Online performance with a preferred selectivity towards CO formation, as it is the case of Au, Ag and Zn, or preference towards HCOOH formation, as is the case of In, Sn and Cd. In a more particular scenario, Cu-based electrocatalysts have arisen as the only category capable of converting CO<sub>2</sub> into hydrocarbons, including our target product, CH<sub>4</sub><sup>59</sup>.

Bulk electrodes in the form of metal foils, normally require high overpotentials to catalyze CO<sub>2</sub> conversion, and thus many studies have recently focused their efforts on nanostructured materials with a higher ratio of surface-accessible active sites. Moreover, nanostructured materials can exhibit rough surfaces, combinations of oxidation states, small crystalline features and defects that can enhance their ECO<sub>2</sub>RR-performance with regards to the corresponding, bulk transition metal. In the following subsection we tackle the effect of structural, chemical and electric properties that result in a high selectivity towards CH<sub>4</sub> formation.

**Table 2:** Summary of representative electrocatalysts for electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion, including their reported active sites, maximum CH<sub>4</sub> Faradaic efficiency (FE), operating current density, applied potentials, stability metrics (if reported), and corresponding references. Current densities marked with \* refer to partial CH<sub>4</sub> current densities. \*\* refers to applied voltage instead of applied potential.

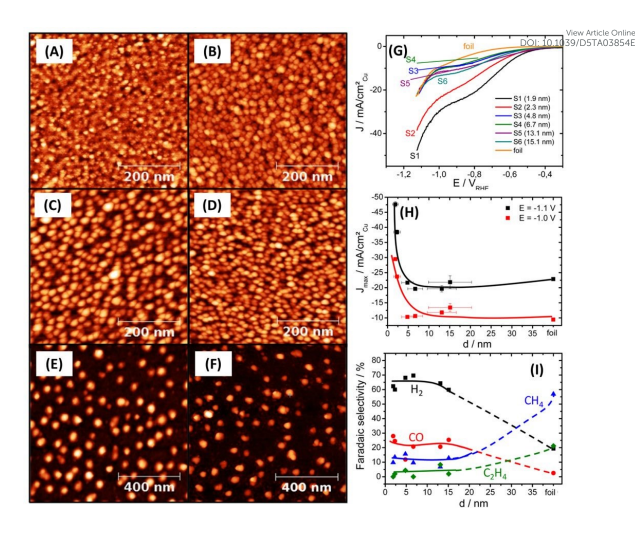
Active site	CH <sub>4</sub> FE / %	j/ mA·cm <sup>-2</sup>	E / V vs. RHE	Stability	Electrolyzer	Reference
Cu nanoparticles	76	11	-1.35	-	H-type cell	(71)
Cu nanoparticles	58	105	-1.00	-	Flow cell	(87)
Cu nanoparticles	73	234	-1.10	-	Flow cell	(88)
Cu nanoparticles	60	230	4 V**	50 h	Zero-gap cell	(88)
CuN <sub>4</sub>	55	35	-1.25	12 h	H-type cell	(93)
CuN <sub>2</sub> B <sub>2</sub>	73	292	-1.46	8 h	Flow cell	(95)
CuN <sub>2</sub> O <sub>2</sub>	78	40	-1.44	6 h	H-type cell	(97)
La <sub>5</sub> Cu <sub>95</sub>	65	300	-1.72	-	Flow cell	(99)
Pd single atoms on Cu	60	118*	-1.10	-	Flow cell	(103)
N <sub>x</sub> C-encapsulated Ag nanoparticles	44	7	-1.40	10 h	H-type cell	(108)
N-doped carbons	15	30	-0.90	-	H-type cell	(133)
F- and N-doped carbons	99	0.2	-0.80	-	H-type cell	(150)
Cu single-atoms	62	136	4 V**	110 h	Zero-gap cell	(171)
Cu single-atoms	82	400	-0.90	5 h	Flow cell	(201)

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#### 3.2.1 Cu-based electrocatalysts

As mentioned before, copper (Cu) outstands over other metals due to its capability for converting CO<sub>2</sub> into hydrocarbons, including CH<sub>4</sub>, C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>6</sub>, with low selectivity towards a single product<sup>63</sup>. However, understanding the factors that determine this catalytic activity remains a challenge. Modelling works grouped all Cu crystal facets in order of the ECO<sub>2</sub>RR activity, being Cu(211) the most active surface, followed by Cu(110), Cu(100), and finally Cu(111)<sup>64</sup>. The findings indicate that Cu(211) is the surface of pure Cu that exhibits the best electrocatalytic activity for both CO and CH<sub>4</sub> formation, which unfortunately make them non selective towards the formation of one unique product. Experimentally, initial studies with Cu foils 65 demonstrated that the preparation conditions of the Cu surface affect the selectivity and efficiency towards certain products. The experiments showed that methane is produced when the Cu surface is cleaned with HCl rather than HNO<sub>3</sub> or oxidized in air, suggesting that the oxidation state of Cu significantly influences the reaction mechanisms. Some years later, a few studies demonstrated that improved hydrocarbon selectivity was observed in oxidized Cu foils due to the presence of Cu<sub>2</sub>O sites, which act as a more active phase for certain hydrocarbons than metallic Cu<sup>66–68</sup>. More precisely, the work of Mistry et al<sup>66</sup> demonstrated that a H<sub>2</sub> plasma treatment of polycrystalline Cu led to a metallic Cu surface that yielded a large C<sub>1</sub> product selectivity, with a FE for CH<sub>4</sub> formation of 37 %. In contrast, an O<sub>2</sub> plasma treatment of polycrystalline Cu produced an oxidized surface with nearly 0 % C<sub>1</sub> products selectivity, but a high conversion to C<sub>2</sub> products, including a FE for C<sub>2</sub>H<sub>4</sub> of 60 %.



**Figure 4:** Tapping-mode atomic force microcospy images of Cu nanoparticles supported on SiO<sub>2</sub> (4 nm)/Si(111): (**A**) S1, (**B**) S2, (**C**) S3, (**D**) S4, (**E**) S5, and (**F**) S6. (**G**) Linear sweep voltammetries recorded on glassy carbon supports, S1-S6, in a CO<sub>2</sub>-saturated 0.1 M KHCO<sub>3</sub> solution acquired at room temperature and at a scan rate of 5 mV s<sup>-1</sup>. Current densities were normalized by the Cu particle surface area after subtraction of the glassy carbon background signal at -1.1 V vs RHE. A Cu foil is included as reference. (**H**) Particle size effect during catalytic CO<sub>2</sub> reduction. The faradaic current densities at -1.1 and -1.0 V vs RHE are plotted against the size of the Cu nanoparticles. The current densities have been normalized by the Cu particle surface area after subtraction of the glassy carbon background signal. (**I**) Particle size dependence of the faradic selectivity towards various reaction products during the CO<sub>2</sub> reduction reaction. Reproduced from reference <sup>69</sup> with permission from the American Chemical Society, copyright 2014.

Recently, it was demonstrated that the presence of an oxidizing agent in the electrolyte can also modulate the ECO<sub>2</sub>RR of Cu electrocatalysts. As shown in the literature<sup>70</sup>, oxygen-containing species not only enhance the rate of the ECO<sub>2</sub>RR, but also tune the selectivity for certain products. The presence of H<sub>2</sub>O<sub>2</sub> accelerates the rate of CH<sub>4</sub> production by a factor of 200 in Cu electrodes, whereas in O<sub>2</sub>-containing solutions a strong decrease of the CH<sub>4</sub> production is observed. The authors relate such a behavior to an increase in the surface concentration of oxygen-containing Cu species that, when stabilized, can significantly enhance the ECO<sub>2</sub>RR activity and selectivity of such electrodes, in a very similar way to what was discussed above

for pre-oxidized Cu samples. In addition, the same authors claimed that the selectivity of Stao3854E different products can be tuned by the chemical structure of the oxygen-containing species.

#### 3.2.1.1 Cu Nanoparticles electrocatalysts

With the advance in nanoparticle synthesis, controlling the size, composition, structure and morphology of Cu nanoparticles is nowadays possible. For such nanoparticulate materials, there are three main parameters that determine the ECO<sub>2</sub>RR performance: their composition, size and support. The size of Cu nanoparticles has proven to play a key role in ECO<sub>2</sub>RR catalysis. Cu nanoparticles showed high selectivity towards CH<sub>4</sub> formation (FE of 76 %), which is substantially superior to the CH<sub>4</sub> FE of polycrystalline Cu (44 %)<sup>71</sup>. Nevertheless, Cu nanoparticles with a diameter below 15 nm tend to enhance competitive reactions, such as H<sub>2</sub>-evolution and electrochemical CO<sub>2</sub>-to-CO formation, lowering significantly the FE towards CH<sub>4</sub><sup>69</sup>. A controlled synthesis of Cu nanoparticles of different sizes is illustrated in the AFM images in Figure 4A-F. As shown in Figures 4G and 4H, the smaller the nanoparticle diameter, the higher the current density in linear sweep voltammetry curves. This enhanced current density is not only associated with an improvement of the ECO<sub>2</sub>RR performance, but also to an increase in the HER activity. The smaller nanoparticles exhibit a substantial enhancement in selectivity towards H<sub>2</sub>, whereas larger nanoparticles tend to favor ECO<sub>2</sub>RR-products<sup>69</sup>. As a result, large nanoparticle sizes are recommendable for hydrocarbons production (Figure 4I).

Another essential aspect when designing Cu-based electrocatalysts is the catalytic support. The main goal of this support is to inhibit nanoparticle agglomeration during catalyst synthesis and under working conditions<sup>72</sup>, and to provide diffusion channels for the supply of the reactant and the evacuation of the products<sup>63</sup>. Moreover, catalytic supports are also essential to assure electrical conduction paths to the metal sites. The most common supports are based on carbon

materials<sup>73</sup>, although other materials can also act as effective supports, such as polymers<sup>74 $\omega$ 7/6ticle Online CeO<sub>2</sub>77,78 or molybdenum-based 2D materials<sup>79–81</sup>.</sup>

Carbonaceous materials are low-cost and abundant catalytic supports, which makes them promising for reducing electrocatalyst costs<sup>28</sup>. Anchoring metals on carbon materials has been an excellent approach for large-scale electrocatalyst production<sup>82</sup>. The metal phase provides the active sites for the ECO<sub>2</sub>RR, while the carbon provides a high surface area that reduces diffusion limitations and exposes a higher number of Cu active sites, while assuring that the metal-support assembly remains electrically conductive <sup>28</sup>. Carbon materials, especially those that are doped with heteroatoms, feature excellent nanoparticle-anchoring capabilities owing to their metal-heteroatom chemical bonds. Nitrogen (N) is by far the most studied heteroatom<sup>69,83–85</sup>, since its similar size to carbon and unique electronic configuration generates electron delocalization in the carbon layers, resulting in an n-type-like semiconductor, as the substitution of a carbon atom by nitrogen introduces an extra electron into the carbon structure, which can move through the electron cloud, enhancing electrical conductivity. Additionally, N atoms strongly interact with metal phases, avoiding nanoparticle leaching and agglomeration and thus enhancing the electrocatalyst's *operando* stability <sup>86</sup>.

One example of such N heteroatom effect was observed by Dai et al. <sup>87</sup>, who anchored Cu on a carbon support through strong metal-heteroatom interaction by pyridine-based N-functionalities (pyridine-substituted graphdiyne (Py-GDY) obtained by cross-coupling of 1,3,5-triethynyl-2,4,6-tris(4-pyridyl)benzene. The non-doped homologous material (GDY) exhibited larger and more heterogenous Cu-nanoparticle sizes, and while the catalyst prepared with Py-GDY showed excellent ECO<sub>2</sub>RR selectivity with a CH<sub>4</sub> FE of 58 % at a current density of 105 mA cm<sup>-2</sup>, the catalyst made with the non-doped GDY only had a methane Faradaic efficiency of 37 % at the same current. The authors attributed this enhanced selectivity for CH<sub>4</sub>-formation to the pyridyl groups, which led to strong metal-N interaction that resulted in

uniformly dispersed Cu nanoclusters of about 2 nm<sup>87</sup>. In a similar work<sup>88</sup>, the authors prepared action of about 2 nm<sup>87</sup>. In a similar work<sup>88</sup>, the authors prepared action of a similar work<sup>88</sup>, the authors prepared action of a similar work<sup>88</sup>, the authors prepared action of a similar work<sup>88</sup>, the authors prepared excellent properties towards CH<sub>4</sub> selectivity, with a maximum FE of 73 % and a CH<sub>4</sub>-specific current density of 230 mA cm<sup>-2</sup> at -1.1 V vs. RHE, which is again much higher than that of the analogous catalyst prepared on the non-doped support at the same potential (CH<sub>4</sub>-specific current density and FE below 50 mA cm<sup>-2</sup> and 20 %, respectively). In this case, the high methane selectivity was attributed to the pyrrolic N in the vicinity of the Cu nanoparticles, which accelerates the hydrogenation of reaction intermediates.

In summary, the surface functional groups in carbon supports have proven to be capable of modulating the selectivity and activity of electrochemical CO<sub>2</sub>-reduction towards CH<sub>4</sub> conversion, and thus such functionalities may play a decisive role in the future development of highly effective electrocatalysts for this reaction.

#### 3.2.1.2 Cu Single-Atom electrocatalysts

The maximized atomic utilization and well-defined coordination of single atom sites make them the most promising active centers among Cu-based ECO<sub>2</sub>RR-electrocatalysts<sup>89</sup>. The rational design of the coordination of the metal center, including the nature and number of the coordination atoms affects substantially the electronic structure of the metal site and should cause significant changes in the reaction pathway and product selectivity with regards to the extended metal surface. A Cu-N<sub>4</sub> coordination is the most desirable configuration for ECO<sub>2</sub>RR due to its high stability, stemming from its optimal thermodynamic interaction between Cu atoms and reaction intermediates<sup>89</sup>. It is widely acknowledged that Cu single atoms often produce hydrocarbons with only one carbon atom (i.e., C<sub>1</sub> products) because the lack of adjacent active sites restrain C-C coupling. Beyond this general observation, some works report high selectivity towards CH<sub>3</sub>OH or CH<sub>4</sub>, while a few other studies reported a preference for

CO formation<sup>90,91</sup>. The reasons for this discrepancy remain unanswered, with new study stilled online focusing more on electrocatalytic results rather than in the understanding of these disparities.

As an example, Cu single atoms displaying the above Cu-N<sub>4</sub> configuration and incorporated in carbon nanofibers electrochemically reduced CO<sub>2</sub> to CH<sub>3</sub>OH with a FE of 44 % and at a current density of 93 mA cm<sup>-2</sup> <sup>92</sup>. DFT modelling showed that these Cu-N<sub>4</sub> single atoms possess a high adsorption energy for the \*CO intermediate<sup>92</sup>, and that the free energy barrier of the subsequent \*CHO to \*CHOH step is much lower than that of the \*CHO to \*CH<sub>2</sub>O path, thus favoring methanol over CH<sub>4</sub>- or CO-production. However, in a different study CuN<sub>4</sub> single atoms obtained from Cu phthalocyanine exhibited a high FE towards CH<sub>4</sub> (55 % at a current density of –35 mA cm<sup>-2</sup>)<sup>93</sup>. At the same time, a high CO<sub>2</sub>-to-CO conversion was recently reported with CuN<sub>4</sub> single atoms anchored on carbon materials<sup>94</sup> that showed a FE towards CO of > 90 %. These are just a few examples of the large variety of selectivity trends reported for CuN<sub>4</sub> sites and for which the disparities remain elusive. All these previous works have employed DFT calculations to support the reported selectivity and mechanisms, in principle proving the thermodynamic viability of the individual ECO<sub>2</sub>RR pathway towards CH<sub>3</sub>OH, CH<sub>4</sub> or CO, but without modelling all other possible reaction mechanisms, which make these results inconclusive.

Another important factor that may account for these discrepancies is the nature of the supporting matrix. Beyond simply stabilizing the Cu–N<sub>4</sub> moieties, the substrate can strongly influence charge distribution, binding energies of intermediates, and even site stability. For instance, CuN<sub>4</sub> decorated on N-doped carbon dots have been reported to deliver FEs above 80% for ethanol<sup>95</sup>, while Cu single atoms embedded in a porphyrin-based MOF produced CH<sub>4</sub> with a FE of  $\approx 80\%^{96}$ . However, the substrate is clearly not the sole determinant: another study on Cu single atoms in a porphyrin-based MOFs instead showed a high selectivity towards acetate (>40%) with negligible CH<sub>4</sub> production<sup>97</sup>. These contrasting examples highlight that

the catalytic outcome arises from a complex interplay between the atomic site and physicochemical properties of the support, and that decoupling these contributions remains a major challenge. It should also be emphasized here that many metal-free catalysts, especially carbon-based materials, are catalytically active for the ECO<sub>2</sub>RR (see Section 3.3 Metal-free carbon-based electrocatalysts) which further complicates the interpretation of activity trends.

A potential strategy to address the discrepancies is the application of advanced in-situ/operando spectroscopic techniques capable of resolving reaction intermediates under realistic electrochemical conditions. We believe that by directly tracking adsorbed species and identifying the rate-determining steps the field can move beyond purely thermodynamic DFT predictions, which often neglect alternative pathways and dynamic effects under operating conditions. At the same time, the current understanding of the role of the supporting material remains limited. In this regard, in-situ/operando characterization will be essential not only to determine the intrinsic activity of CuN<sub>4</sub> sites, but also to understand why their catalytic properties vary so markedly depending on other factors such as the local coordination environment, the supporting material, and the surrounding reaction microenvironment.

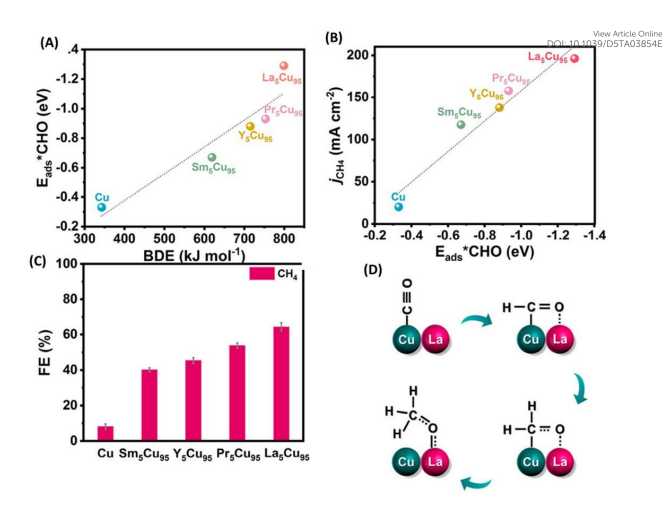
Nevertheless, given the current lack of mechanistic consensus, many studies have adopted an empirical approach by tuning the local environment of the SAC's metal centers to directly modulate the ECO<sub>2</sub>RR selectivity. The introduction of B atoms in the local coordination of the Cu single atoms, especially in the Cu-N<sub>2</sub>B<sub>2</sub> configuration, results in a significant increase of the catalyst's selectivity towards CH<sub>4</sub> formation, with a FE of 73 % at a partial current density of - 292 mA cm<sup>-2</sup> 98, as opposed to only 30 % CH<sub>4</sub> at barely 100 mA cm<sup>-2</sup> at the same potential (- 1.46 V vs RHE) for the analogous Cu-N<sub>4</sub>-based material. The authors attribute this improvement brought along by the introduction of B to a lower energy requirement for the \*CO to \*CHO thermodynamic barrier.

Interestingly, similar effects were found when introducing a low concentration (5%, 5%) and Agicle Online catalyst weight basis) of Cu single atoms in CeO<sub>2</sub> nanorods. Such an addition induced a substantial increase in the CH<sub>4</sub> selectivity up to a FE of 58% due to the creation of oxygen vacancies surrounding the Cu-N<sub>4</sub> sites<sup>99</sup>. Similarly, the introduction of oxygen atoms coordinated directly to the Cu single atoms (i.e. in the form of CuN<sub>2</sub>O<sub>2</sub>) can also efficiently modify the electronic structure of the metal center for reducing the thermodynamic barriers towards CH<sub>4</sub> formation<sup>100</sup>. As a result, the CuN<sub>2</sub>O<sub>2</sub>-containing electrocatalysts shows a superior CH<sub>4</sub> selectivity with a FE of 78 % at -1.44 V with a total current density of 40 mA·cm<sup>-2</sup>.

In summary, the modulation of the chemistry in the local coordination or vicinity of Cu-N<sub>4</sub> sites seems to be a promising strategy to reach high selectivities towards CH<sub>4</sub> formation. Further studies with different heteroatoms, defect engineering and pioneering synthesis strategies are mandatory to obtain even more selective materials and a better understanding about the mechanisms behind these phenomena.

#### 3.2.1.3 Cu-based bimetallic electrocatalysts

The electrocatalytic activity of CO<sub>2</sub>-to-CH<sub>4</sub> electrochemical conversion appears to be related to the linear relation between the adsorption energies of \*CO and \*CHO intermediates, which restricts the selectivity towards CH<sub>4</sub> formation. The inclusion of a second metal either in the form of a single atom (using the so-called dual-site configuration) or through the alloying of metallic Cu with a second element can provide a new degree of freedom to modulate this adsorption energy difference, allowing a better selectivity towards CH<sub>4</sub><sup>101</sup>.



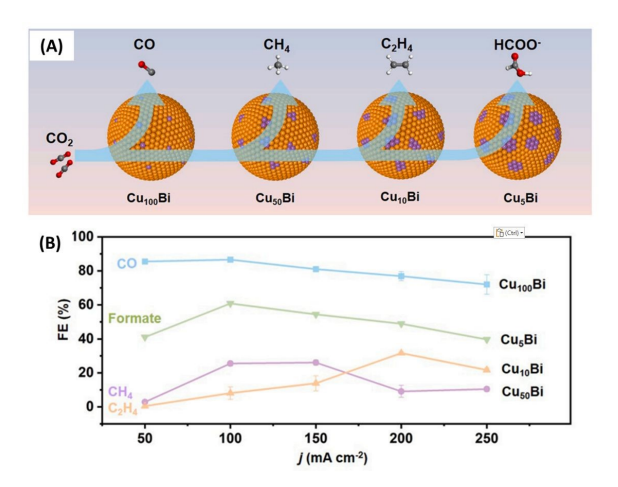
**Figure 5:** (**A**) Relation between calculated \*CHO adsorption energy (energy required to convert one molecule from the lowest adsorbed state to the lowest gaseous state) and the bond dissociation enthalpy (BDE) of the metal-oxygen bond. (**B**) Relation between the partial current density of CH<sub>4</sub> at total current density of 300 mA cm<sup>-2</sup> (in CO<sub>2</sub>-saturated 1 M KOH solution) and the adsorption energy of \*CHO for Cu and M<sub>5</sub>Cu<sub>95</sub> electrocatalysts. (**C**) FE of CH<sub>4</sub> on Cu and M<sub>5</sub>Cu<sub>95</sub> electrocatalysts at a total current of 300 mA cm<sup>-2</sup>. (**D**) Proposed reaction pathways of ECO<sub>2</sub>RR to CH<sub>4</sub> on the La<sub>5</sub>Cu<sub>95</sub> surface. Reproduced from reference <sup>102</sup> with permission from the American Chemical Society, copyright 2023.

Dual Cu adjacent sites have also been proposed as a promising pathway to increase the selectivity towards hydrocarbon products<sup>89,103</sup>. Although this system is not strictly a bimetallic material, we find it useful to include it in this section because the results of adding an additional Cu atom closely resemble those observed in bimetallic materials rather than in single-atom Cu catalysts. In dual-site Cu catalysts, two neighboring Cu atoms are anchored onto a support and work cooperatively to enhance CO<sub>2</sub> reduction activity. In these systems, Cu atoms serve as active sites for the \*CO<sub>2</sub> adsorption and when two \*CO<sub>2</sub> intermediates are adsorbed close to each other, the proximity significantly favors C-C coupling reactions, leading to the formation of hydrocarbons, especially C<sub>2+</sub> products<sup>103</sup>. Such a synergistic effect of dual Cu sites was

further demonstrated through the tuning of the neighboring distance between Cu sites of the products of the neighboring distance between Cu sites of the products. At modulated the selectivity to different hydrocarbons, allowing the formation of C<sub>2+</sub> products. At Cu concentrations above 4.9 mol %, the distance between adjacent CuN<sub>4</sub> species was close enough to enable C-C coupling of ECO<sub>2</sub>RR intermediates, resulting in high selectivity towards C<sub>2</sub>H<sub>4</sub><sup>104</sup>. In contrast, concentrations below 2.4 mol % demonstrated a high selectivity towards CH<sub>4</sub><sup>104</sup>. Thus, the metal content and corresponding distance between CuN<sub>4</sub> sites is of high relevance for the product selectivity.

In the case of bimetallic alloys, a large number of studies that have tried to overcome the low selectivity of Cu-based electrocatalysts for the formation of CH<sub>4</sub> through the synthesis of Cubased bimetallic alloys. More precisely, trace amounts (from 0 to 7 at.%) of oxophilic metals (La, Pr, Y, and Sm) were introduced into a pure Cu matrix for evaluating the ECO<sub>2</sub>RR performance<sup>102</sup>. As observed in Figure 5A, DFT calculations showed that the oxophilicity of the alloying metal correlates strongly with the adsorption energy of \*CHO, which is known to be a key parameter in ECO<sub>2</sub>RR selectivity. Experimentally, the introduction of La in the Cu structure in a La<sub>5</sub>Cu<sub>95</sub> ratio resulted in an excellent partial current density of 194 mA cm<sup>-2</sup> (Figure 5B) with a high FE towards CH<sub>4</sub> of 64.5 % at 300 mA·cm<sup>-2</sup> (Figure 5C). This is attributed to the alloying of Cu with La, which not only stabilizes the \*CHO intermediates, but also promotes the splitting of the C-O bond in the \*CH<sub>3</sub>O reaction intermediate by forming a stable La-O bond (Figure 5D). Similar conclusions were found through the introduction of Bi in a Cu aerogel<sup>105</sup>, which resulted in the variation of the Cu<sup>2+</sup>/Cu<sup>+</sup> ratio on the electrocatalysts surface and in turn led to different FEs towards ECO<sub>2</sub>RR products (Figure 6A). In particular, the highest FE towards CH<sub>4</sub> was obtained for a Cu<sub>50</sub>Bi material that displayed a FE towards CH<sub>4</sub> of 26% at a total current density of 150 mA cm<sup>-2</sup>. Additionally, as observed in Figure 6B, the tuning of the Cu<sub>x</sub>Bi composition leads to a significant modification of the ECO<sub>2</sub>RR selectivity, with FEs towards CO above 80 % in the case of Cu<sub>100</sub>Bi, or 60 % FE towards

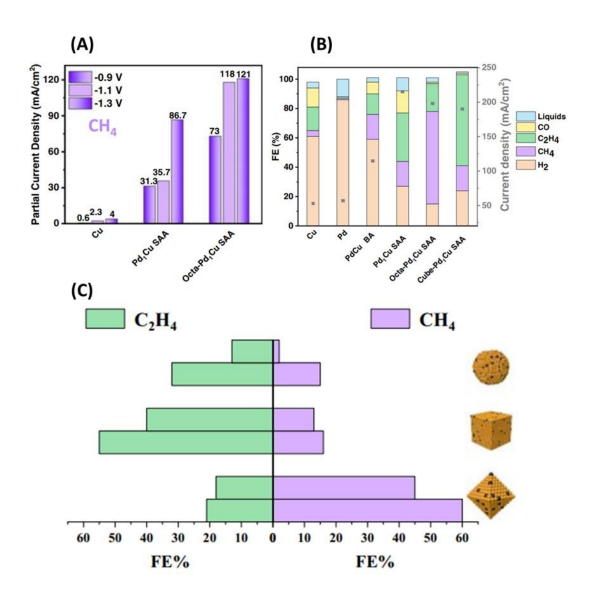
formate with  $Cu_5Bi$ . Interestingly, the higher  $Cu^{2+}/Cu^+$  ratio increased significantly  $\frac{1}{39}$   $\frac{1$ 



**Figure 6:** (A) Products distribution of the ECO<sub>2</sub>RR with different  $Cu_XBi$  compositions. (B) Electrocatalytic performance (in  $CO_2$ -saturated 1 M KOH solution) of the corresponding  $Cu_XBi$  aerogels, showcasing the FEs for ECO<sub>2</sub>RR products at different applied current densities for  $Cu_{100}Bi$ ,  $Cu_{50}Bi$ ,  $Cu_{10}Bi$  and  $Cu_5Bi$ . Reproduced from referece <sup>105</sup> with permission from Elsevier, copyright 2022.

Another interesting work tried to integrate H-affine elements within Cu. Although platinum-group elements are known for their high HER-selectivity, when atomically dispersed platinum atoms are integrated in a shaped-controlled Cu catalyst, they increase the selectivity towards hydrocarbon products. For instance, Pt-group single atoms act as a H\* supplier and facilitate the hydrogenation of \*CO intermediates in polycrystalline Cu surfaces<sup>106</sup>. The incorporation of Pd single atoms to Cu-based electrocatalysts led to an increase in the CH<sub>4</sub> partial current density from 2.3 to 118 mA cm<sup>-2</sup> (Figure 7A), with an improvement of the FE towards CH<sub>4</sub>

from 2 to 60 % at -1.1 V vs RHE (Figure 7B). Moreover, the alloy phase morphology is  $\frac{V_{ext}}{AO3854E}$  relevant, since spherical or cubic structures resulted in high  $C_{2+}$  yields, whereas octahedral structures promoted  $CH_4$  formation (Figure 7C). Chiefly, this strategy of adjusting the metal nature and metal doping to tune the \*CHO adsorption energy can be extended to increase the hydrocarbon selectivity of Cu electrocatalysts towards products other than  $CH_4^{107,108}$ .



**Figure 7:** (A) Partial current density for CH<sub>4</sub> at different voltages for polycrystalline Cu, polycrystalline Pd<sub>1</sub>Cu configuration and shape-controlled Octa-Pd<sub>1</sub>Cu configuration. SAA stands for single-atom alloys. (B) Comparison of CO<sub>2</sub> reduction FE (%) and current densities of Pd<sub>1</sub>Cu configuration and analogues. The FE was obtained from amperometric i·t curves at -1.1 V vs RHE (in a CO<sub>2</sub>-saturated 0.5 M KHCO<sub>3</sub> solution) and quantifying the products over 30 min. (C) FE distribution of CH<sub>4</sub> and C<sub>2</sub>H<sub>4</sub> obtained using Cu and shape controlled Pd<sub>1</sub>Cu materials. Reproduced from reference <sup>106</sup> with permission from Springer Nature, copyright 2023.

#### 3.2.2 Silver-based electrocatalysts

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Ag-based electrocatalysts are of the greatest interest for CO production because of this metal's relatively low cost (particularly when compared with noble metals) and high selectivity for this product<sup>109</sup>. This is why most of the works that focus on Ag electrocatalysts for ECO<sub>2</sub>RR do so with a strong focus on CO production. However, the size, crystal structure and morphology of silver (nano)catalysts can further tune their ECO<sub>2</sub>RR performance. As a result, Ag-based electrocatalysts in which the properties of the pristine metal have been tuned through the addition of a second metal phase have recently arisen as a promising approach for the electrochemical conversion of CO<sub>2</sub> into CH<sub>4</sub>.

As an example of this, a 2017 study showed that Ag-Co bimetallic electrocatalysts with electronic properties different from those of the monometallic Ag counterpart featured a methane FE of 20 % at an applied voltage of 2.0 V (vs nearly 0 % for the monometallic catalyst) thanks to the atomic rearrangement and concomitant tuning of the binding strength of the \*CO intermediate, which favors CH<sub>4</sub><sup>110</sup>. Specifically, Co atoms provide free electrons to the vacant orbitals in Ag, forming Ag-Co bonds that induce large changes in the electron density of the metals and shift the ECO<sub>2</sub>RR selectivity towards CH<sub>4</sub> instead of CO. Similarly, the deposition of a N<sub>x</sub>C shell surrounding the Ag nanoparticles also serves to tune the selectivity products<sup>111</sup>, in this case not through a modification of silver's electronic properties, but by prolonging the residence time of the \*CO intermediate on the catalyst's surface and increasing the number of hydrogenation reactions. As a result, the FE for CH<sub>4</sub> formation was enhanced above 44 % after the creation of the N<sub>x</sub>C nanoshell, as compared to nearly 0 % in the absence of the latter (i.e., for bare Ag-nanoparticles)

In summary, although there are not many studies that focus on modifying the ECQ R RAO3854E selectivity of Ag electrocatalysts, the few works discussed above demonstrate that the tuning silver's properties can also be a promising strategy towards CH<sub>4</sub> formation.

#### 3.1.3 Cobalt-based electrocatalysts

DFT computations indicate that the chemisorption of the \*CO<sub>2</sub> intermediate is the rate-limiting step upon ECO<sub>2</sub>RR on CoN<sub>4</sub> sites<sup>112</sup>. While CO is the main product for such Co-based electrocatalysts<sup>113</sup>, CH<sub>4</sub> is eventually produced in minor amounts due to the hydrogenation of the \*CO intermediate. This is confirmed experimentally, since a FE towards CO of 97-99 % was obtained when investigating Co phtalocyanine as the ECO<sub>2</sub>RR active site, and negligible formation of CH<sub>4</sub> was observed<sup>76,114</sup>. Nevertheless, even if the majority of Co-based ECO<sub>2</sub>RR electrocatalysts are designed for CO production, strategies again exist to improve their selectivity towards CH<sub>4</sub>.

For instance, electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion has been achieved when employing photo-assisted electrochemical methods<sup>115</sup>. The addition of light illumination during electrocatalysis promotes the stabilization of the \*CO intermediate adsorbed on the Co-N<sub>4</sub> sites, allowing its subsequent hydrogenation to yield CH<sub>4</sub>. Along these lines, the incorporation of Co phthalocyanine to a Zn-N-C materials caused a 100-fold enhancement in the selectivity of the latter materials towards CH<sub>4</sub> formation<sup>116</sup>. DFT calculations revealed that the first reduction of CO<sub>2</sub> molecules occurs at the Co-N<sub>4</sub> sites, but the resulting \*CO intermediate is transferred onto the Zn-N<sub>4</sub> sites for further conversion into CH<sub>4</sub><sup>116</sup>.

Interestingly, a strong effect of the supporting electrolyte was also observed in Co porphyrins<sup>117</sup>. Cation size affects the reaction in two ways: large cations, such as K<sup>+</sup>, enhance the HER because smaller cations retain more water, creating steric hindrance that slows H<sup>+</sup>

diffusion. Conversely, small cations, such as Li<sup>+</sup>, stabilize the key intermediate of conversely, small cations, such as Li<sup>+</sup>, stabilize the key intermediate of conversion of CO<sub>2</sub> into CO and CH<sub>4</sub>. <sup>117</sup>.

#### 3.2.3 Fe-based electrocatalysts

Fe is among the most promising transition metals for catalyzing many electrochemical reactions due its high abundance and corresponding low-cost  $^{118-121}$ . This means that tuning the electronic properties of Fe electrocatalysts is not a new strategy in materials science, but how this approach affects these materials' ECO<sub>2</sub>RR performance is still unknown. Furthermore, the poor HER-activity of Fe-based electrocatalysts (typically requiring overpotentials > 400 mV  $^{82}$ ) underscores the capabilities of these electrocatalysts for promoting CO<sub>2</sub>-electroreduction over H<sub>2</sub>-production.

Iron, especially in the form of single-atoms with an FeN<sub>4</sub> coordination, is known to electrochemically reduce CO<sub>2</sub> into CO with a high selectivity (i.e., a FE > 90%)<sup>122–124</sup>. The correspondingly poor selectivity towards CH<sub>4</sub> formation is mainly associated to the isolated nature of the FeN<sub>4</sub> active sites. Computations indicate that ECO<sub>2</sub>RR in FeN<sub>4</sub> sites can indeed proceed beyond CO<sup>125</sup>. Specifically, since FeN<sub>4</sub> sites are excellent CO producers and therefore can form a large amount of \*CO intermediates, these reaction intermediates can be further reduced until yielding hydrocarbons. However, this work suggested that highly active CH<sub>4</sub> formation may require an extended surface or nearby proton source to lower the protonation barrier<sup>125</sup>.

We are only aware of one study in which Fe exhibited CH<sub>4</sub> formation<sup>126</sup>. This work explored structure-activity relations in FeN<sub>4</sub> catalysts, revealing that the catalytic activity is strongly influenced by the nitrogen functionalities. Specifically, XPS data suggested that pyridinic N and FeN<sub>4</sub> moieties serve as active sites, as they promote CO<sub>2</sub> adsorption and facilitate electron transfer. Operando EXAFS results indicated a change in the Fe oxidation state from Fe<sup>2+</sup> to

Fe<sup>1+</sup> at potentials of approximately -0.9 to -1.1 V vs. RHE, coinciding with the onset of 35/154 A03854E production. The authors attributed this CH<sub>4</sub> formation to the FeN<sub>4</sub> sites with the Fe atom in the +1 oxidation state. While CH<sub>4</sub> production remained very low (< 1 %), this redox transition offers a potential pathway to enhance the CO<sub>2</sub>-to-CH<sub>4</sub> activity of FeN<sub>4</sub> sites by tuning their electronic configuration.

#### 3.3 Metal-free carbon-based electrocatalysts

In the pursuit for low-cost ECO<sub>2</sub>RR electrocatalysts, metal-free carbon-based materials have arisen as an excellent alternative to those containing metals<sup>28,127,128</sup>. Nevertheless, even if these metal-free electrocatalysts have been extensively proven to be effective for other electrochemical reactions such as oxygen reduction<sup>129,130</sup> and hydrogen evolution <sup>130,131</sup>, their ECO<sub>2</sub>RR performance is still far from that of the metal-containing materials. Pristine, undoped carbon materials exhibit minimal catalytic activity towards the ECO<sub>2</sub>RR because the electron density across their carbon layers is homogenously distributed<sup>132</sup>. A common strategy to improve this poor catalytic performance involves altering and tuning their electronic properties through the introduction of heteroatoms, surface functionalities, as well as defect engineering and curvature. Incorporating functional groups or defects to the basal plane of the carbon layers usually induces a redistribution of the electron density through electron withdrawal-donation effects that localize the charge in carbon atoms adjacent to defects or heteroatoms. This electron delocalization near heteroatoms tends to attract the reagent molecule more easily, and when the molecule is chemically adsorbed to the active sites, the polarization of the carbon electrode induces its subsequent reduction.

Nitrogen is the most studied heteroatom when it comes to doping carbon materials due to its similar size to carbon, but also because of its larger electronegativity (3.04 for N and 2.55 for C). N-doped carbon materials have proven to be effective electrocatalysts for multiple

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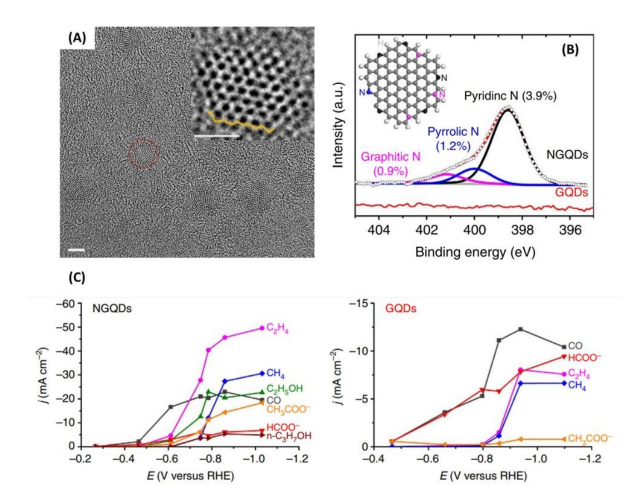
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electrochemical reactions, including ECO<sub>2</sub>RR. Nitrogen atoms can be featured in call wasticle Online materials in the form of different functionalities that can be classified according to their binding energies<sup>133–135</sup>: pyridinic-N (398.4 eV), pyrrolic/pyridonic-N (400.3 eV), graphitic-N (also called quaternary-N, 401.2 eV) and oxidized-N (402.9 eV). In general, the defects located at the edge of the carbon layers are assumed to be more active than those at the basal plane, with N-doped zigzag species being the most active site among these. 136 Figure 8a showcases HRTEM images of N-doped graphene quantum dots (NGQDs) 1-3 nm in size that were prepared through exfoliating and cutting graphene oxide with in situ N doping. The N 1s XPSspectrum confirmed the N doping in the form of pyridinic, pyrrolic and graphitic N species (Figure 8B). Pristine non-doped GODs showed indeed moderate ECO2RR-activity thanks to their large edge density, with a current density of around 10 mA cm<sup>-2</sup> at -0.85 V vs. RHE and high selectivity towards CO (FE = 10%) and HCOO (FE = 6 %) $^{137}$ . However, the high N content at the basal plane of NGQD resulted in a significant improvement of the catalytic activity with a current density of 100 mA cm<sup>-2</sup> at the same potential, and an enhanced selectivity for CH<sub>4</sub> and C<sub>2</sub>H<sub>4</sub> products (Figure 8C, 8D)<sup>137</sup>. Among such N species, pyridinic functional groups appear to be responsible for variations in the selectivity in metal-free carbon-based electrocatalysts, as this edge-type N species substantially decreases the energy barrier for the formation of the \*COOH intermediate, which mainly leads to CO production<sup>138,139</sup>. For high CH<sub>4</sub> or C<sub>2+</sub> production, N species located in the basal plane, such as graphitic N, appear to be the most promising in terms of selectivity. However, for graphitic N, the extra electron located in the  $\pi^*$  antibonding orbital is less accessible for CO<sub>2</sub> chemisorption, as proved by the 1 eV higher chemisorption energy of graphitic N when compared to pyridines<sup>140</sup>. Nevertheless, even with this thermodynamic barrier, graphitic N remains better than the N-free, pristine carbon, as demonstrated by its better ECO<sub>2</sub>RR-activity<sup>140</sup>.

Here it is worth noting that both graphitic and pyridinic N induce an electron withdrawal effection of the Article Online N induce an electron withdrawal effection of the Article Online N induce an electron withdrawal effection of the Article Online N induce an electron withdrawal effection of the Article Online N induce an electron withdrawal effection of the Article Online N induce an electron withdrawal effection of the Article Online N induce an electron withdrawal effection of the Article Online N induce and electron withdrawal effection of the Article Online N induce and electron withdrawal effection of the Article Online N induce and electron withdrawal effection of the Article Online N induce and electron withdrawal effection of the Article Online N induce and electron withdrawal effection of the Article Online N induce and electron withdrawal effection of the Article Online N induce and electron withdrawal effection of the Article Online N induce and electron with the Article Online N induce and electron with the Article Online N induce and electron of in the adjacent carbon atoms, which leads to a redistribution of the charge density in the carbon matrix<sup>139–141</sup>. This results in a negative electron density in the N atom and a corresponding positive charge density in the first neighbor carbon atoms. These negatively charged sites act as active centers onto which the positively charged C atoms from the CO<sub>2</sub> molecules chemisorb, forming a N-CO<sub>2</sub> intermediate. In the case of pyridines, the chemisorption occurs via sp<sup>2</sup>-to-sp<sup>3</sup> hybridization, facilitating CO<sub>2</sub> chemisorption. However, graphitic N already forms three chemical bonds with adjacent carbon atoms, making the creation of an additional chemical bond with the CO<sub>2</sub> molecule highly energy demanding. This is why the carbon atoms in the vicinity of the N functionality are considered the active sites in the case of graphitic N groups. However, in modelling graphitic N, it is common to propose C-CO<sub>2</sub> chemisortion on the carbon atom located in the *ortho* position<sup>142</sup>, which appears unlikely when considering these C-atoms' positive charge induced by the N functionality. Therefore, it is anticipated that the active sites involve the C atom in the meta position to the N through a C-CO2 interaction, or the C atom in *ortho* position through the interaction with the oxygen atoms of the CO<sub>2</sub> molecule through a C-OCO intermediate. As such, further studies are mandatory to unravel the mechanisms behind the ECO<sub>2</sub>RR in N-doped carbon materials.

Other heteroatom elements have also been studied as doping agents in carbon materials, such as B<sup>143–145</sup>, F<sup>146,147</sup>, P<sup>148–151</sup> or S<sup>152,153</sup>. Furthermore, heteroatom doping with these elements has also been carried out with the aim of tuning the electronic structure of the carbon layers and induce larger electron delocalization for further ECO<sub>2</sub>RR studies. One particularly interesting article claimed to reach a FE towards CH<sub>4</sub> formation of 99 % with an F- and N-codoped carbon material<sup>154</sup>. However, this high selectivity was only reached with a very low current density of 0.2 mA cm<sup>-2</sup>. Recently, adjacent N and B atoms in N- and B-codoped carbon materials have

also proven a high CH<sub>4</sub> selectivity (i.e., a FE of 68 %) and a current density of  $\approx 15$ <sub>DM. Access/DSFA03854E</sub> -0.5 V vs RHE<sup>155</sup>.



**Figure 8:** (A) TEM image of NGQDs (scale bar, 2 nm), with the inset showing a high magnification image of a single NGQD containing zigzag edges outlined by the yellow line (inset scale bar, 1 nm). (B) N 1s XPS-spectrum of the NGQD sample, deconvoluted into pyridinic, pyrrolic and graphitic N. The value in parentheses is the corresponding N atomic concentration. (C) Partial current densities as a function of the cathode potential for various electrochemical CO<sub>2</sub> reduction products when using NGQDs (left) and GQD (right) as the electrocatalysts. Reproduced from reference <sup>137</sup> with permission from Springer Nature, copyright 2016.

In summary, metal-free heteroatom-doped carbon materials are still at the earliest stage of development as ECO<sub>2</sub>RR-electrocatalysts. As observed in Table 2, their overall current density remains far below that of metal-containing electrocatalysts, with the best reported current density reaching 40 mA·cm<sup>-2</sup> so far. This large performance gap raises critical questions regarding their practical scalability. Nevertheless, an intriguing characteristic emerges when comparing both families of catalysts. While metal-containing electrocatalysts require very negative potentials to reach CH<sub>4</sub> formation, metal-free electrocatalysts can achieve high FE for

CH<sub>4</sub> already at moderate potentials between -0.5 and -0.9 V vs. RHE. This suggests that reduce online although less active in absolute terms, metal-free catalysts may intrinsically demand lower overpotentials to trigger efficient C-H bond formation. Importantly, long-term stability under operating conditions has not yet been demonstrated for metal-free catalysts, which remains a major barrier for their practical applications. Further advances in catalyst design and mechanistic understanding are therefore mandatory to assess whether such features can be leveraged for scalable and competitive implementation.

#### 4 Electrochemical CO<sub>2</sub> electrolyzers

The ECO<sub>2</sub>RR performance is not only influenced by the selection of the electrocatalyst, but also by the design of the electrochemical reactor (commonly referred to as a CO<sub>2</sub> electrolyzer) where CO<sub>2</sub> is converted into a value-added product. From an industrial perspective, one of the most crucial requirements for the practical deployment of CO<sub>2</sub> electrolyzers is their ability to operate at high current densities, typically at least 200 mA·cm<sup>-2</sup>. Achieving such high current densities is essential for ensuring economically viable reaction rates and maximizing CO<sub>2</sub> conversion efficiency at scale. However, maintaining high activity while preserving selectivity and stability poses significant engineering challenges, as it requires optimizing not only the catalyst, but also the overall reactor architecture (including the balance of plant), mass transport properties and ion management.

The geometry of the CO<sub>2</sub> electrolyzer, including the electrode and electrolyte configurations, plays a crucial role in determining the mass transfer efficiency of the gaseous reactant and the reaction products. These factors have a direct impact on key performance metrics, such as electrocatalytic selectivity, long-term stability and overall CO<sub>2</sub> conversion efficiency. While extensive research has been dedicated to the development and optimization of electrocatalysts to enhance CO<sub>2</sub> reduction activity and selectivity, less attention has been paid to the practical

implementation of these catalysts in application-relevant  $CO_2$ -electrolyzers, and comparative  $Y_{A03854E}$  fewer works can be found on this important topic.

Before delving into the different types of CO<sub>2</sub> electrolyzers, it is important to highlight the critical role of membranes, which are an essential yet often overlooked component in these systems. Membranes serve multiple key functions, such as enabling ion transport between the electrodes, preventing unwanted product crossover, and maintaining system stability, all of which directly impact the overall efficiency and selectivity of the electrolyzers. Since membranes are an essential component in CO<sub>2</sub> electrolyzers and exist in various types, it is important to first explain their nature and functions to understand their application in different electorlyzer types.

#### 4.2 Ion exchange membrane (IEM) CO<sub>2</sub>-electrolyzers

A fundamental component of any CO<sub>2</sub> electrolyzer is the membrane, which acts as a selective barriers that regulates the transport of ions between the electrode compartments while preventing the undesired crossover of reactants and products, which can lead to efficiency losses and reduced selectivity<sup>156</sup>. Depending on their ion transport properties, membranes can be classified into three main types: cation exchange membrane (CEM), anion exchange membrane (AEM) and bipolar membrane (BPM). Each of these membrane types has different characteristics that influence the reaction environment and can significantly influence electrolyte pH, ionic conductivity, and product separation, ultimately affecting the performance and feasibility of the system under industrially relevant conditions<sup>156</sup>.

role,

impact on the viability of CO<sub>2</sub> electrolyzers. The choice of IEM not only dictates performance metrics but also plays a crucial role determining operational costs, particularly large-scale applications. in Importantly, these costs are not limited to the intrinsic price of the membrane itself but extend to the entire system configuration that each membrane requires. Certain membranes impose operational constraints that necessitate additional infrastructure system or modifications to compensate their drawbacks. holistic approach was developed assess and optimize the economics factors involved the electrochemical conversion of CO<sub>2</sub> to CO based on different IEMs<sup>157</sup>. Although that study did not focus on CH<sub>4</sub> production, we

believe that it properly illustrates the costs

electrochemical

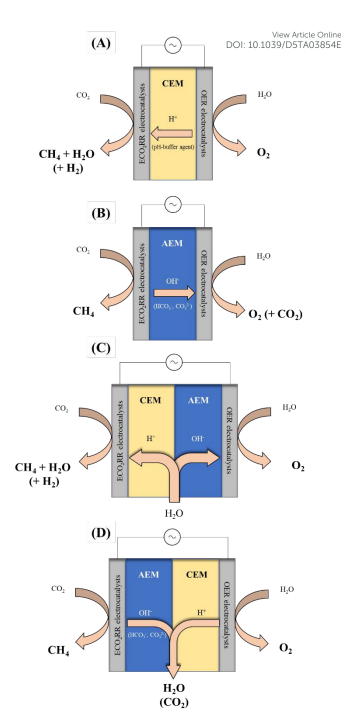
membranes also have a significant economic

Beyond

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their



**Figure 9:** Schematic representation of ion exchange membrane in CO<sub>2</sub> electrolyzers. (**A**) Cation exchange membrane (CEM)-type electrolyzer, (**B**) Anion exchange membrane (AEM)-type electrolyzer, (**C**) Reverse bias bipolar membrane (RB-BPM)-type electrolyzer and (**D**) Forward bias bipolar membrane (FB-BPM)-type electrolyzer.

associated to each IEM type for electrochemical CO<sub>2</sub> conversion. Mass and energy balances were computed using state-of-the-art literature data and revealed that 75 to 84 % of the total production costs can be attributed to the electrolyzer cost. According to the study<sup>157</sup>, AEMs are expected to be the most suitable membranes for ECO<sub>2</sub>RR-electrolysis due to the low electricity

and capital costs associated to their use, leading to an estimated CO-production cost of  $\approx 800^{\circ}_{A03854E}^{\circ}$   $\odot t_{CO}^{\circ}$ . This is closely followed by BPMs, which feature a competitive cost of  $\approx 840 \ \in t_{CO}^{-1}$ , and CEMs with a higher cost of  $\approx 1,100 \ \in t_{CO}^{-1}$ .

Given the importance of the membrane choice in both economic and performance terms, it is essential to delve into the specifics of each IEM type. In the following section, we will analyze the characteristics, advantages and challenges associated with each IEM category, providing a comprehensive understanding of how this influences CO<sub>2</sub> electrolysis systems.

# 4.1.1. Cation exchange membranes (CEMs)

Cation exchange membranes (CEMs) are a class of IEM that selectively allow the passage of cations, such as H+ or K+, while blocking the movement of anions. These membranes are widely used in electrochemical systems, including CO<sub>2</sub> reduction electrolyzers, due to their ability to maintain ionic conductivity and decrease the extent of unwanted crossover of reactants and products.

One of the major challenges of CEMs is the acidic pH at the cathode electrode, which promotes the competitive HER over  $CO_2$ -reduction. To mitigate this limitation, CEM-based  $CO_2$ -electrolyzers, in particular those with electrolyte fed at the anode compartment, can also be operated with a high concentration of  $K^+$  cations from the electrolyte solution, such as KHCO<sub>3</sub> or  $K_2CO_3$ . The high concentration of  $K^+$  in the electrolyte leads to the diffusion of  $K^+$  through the membrane, replacing  $H^+$  ions at the cathode, and consequently suppressing HER and improving the selectivity for  $CO_2$  reduction<sup>158</sup>. The resulting pH-buffer effect strongly correlates with the hydration shell of the electrolyte cation, which is in turn associated with its permeability and diffusion coefficients<sup>159</sup>. Considering an inverse relation between diffusion coefficient and cation hydration radii, the diffusion coefficient for alkali cations follows the order  $Cs^+ > K^+ > Na^+ > Li^{+159}$ . This implies that, from a device-operation perspective,

maintaining high selectivity towards CO<sub>2</sub> reduction requires precise control over the electrolyceroline composition and ion concentration. However, as we will explain later in the CO<sub>2</sub> electrolyzers section, this strategy is only useful for CO<sub>2</sub> electrolyzers that employ liquid electrolytes, which introduces several challenges, such as high electrical resistance, limited CO<sub>2</sub> solubility and concomitantly low current densities.

Another critical concern in CEM-based CO<sub>2</sub>-electrolysis relates to the crossover of CO<sub>2</sub>-products during operation, which leads to the need of including additional separation and regeneration processes and would increase these devices' operational complexity. Although CO<sub>2</sub> crossover is not a significant issue, CO<sub>2</sub>-derived alcohols, such as methanol and ethanol, are more problematic with regards to crossover due to their high diffusion coefficients in CEM<sup>160,161</sup>.

# 4.1.2. Anion exchange membranes (AEMs)

Anion exchange membranes (AEMs) serve as a key alternative to CEMs in  $CO_2$  electrolyzers, offering the advantage of selectively allowing the migration of anions, such as  $OH^+$ ,  $HCO_3^-$  and  $CO_3^{2-}$ , while blocking cations. AEMs are particularly advantageous in  $CO_2$  electrolysis because they create a neutral-to-alkaline environment at the cathode, which promotes the  $CO_2$  reduction reaction over the  $HER^{162}$ . As for the nature of the anionic species, while  $OH^-$  anions are produced from the reduction of  $H_2O$  and/or  $CO_2$  in the cathodic catalyst layer, they reaction with  $CO_2$  results in carbonate and bicarbonate anions (with the distribution of these species' concentrations being determined by the local pH), and all three species get transported through the AEM to the anode to yield  $H_2O$ ,  $O_2$  and  $CO_2$  molecules resulting from the oxidation of  $HCO_3^-$  and/or  $CO_3^{2-163-165}$ . This  $CO_2$ -regeneration at the anode decreases the electrolyzer's net carbon dioxide consumption, and makes it mandatory to put into place additional separation steps that substantially increases the electrolyzer  $cost^{166}$  and can be more energy-demanding

than that of the  $CO_2$ -electrolyzer itself<sup>163–165</sup>. Indeed, the separation of the  $CO_2$  that has crossed across the membrane to the anode compartment is estimated to be  $\approx 1.6$  times more energy demanding than the  $ECO_2RR$  step<sup>167</sup>.

Similar to what was discussed above for CEM-based CO<sub>2</sub>-electrolyzers, the crossover of products from cathode to anode is a major bottleneck for AEM CO<sub>2</sub>-electrolysis development. Indeed, it has been reported that more than 30 % of all cathodic liquid products cross the AEM to the anode compartment<sup>168</sup>. This crossover increases linearly with current density and CO<sub>2</sub> flow rate, and makes it significantly difficult to obtain high energy efficiencies. However, this product crossover is minimized when the targeted product is not a liquid but a gas, such as CH<sub>4</sub>, although crossover of volatile products through the GDE can also occur through evaporation<sup>168</sup>. Nevertheless, the high selectivity and efficiency of AEM-based electrolyzers compared to CEM-based ones nearly compensates such a substantial extra cost of recycling CO<sub>2</sub> from the anode<sup>162</sup>. This advantage stems from AEM electrolyzers' neutral-to-alkaline operative pHs in the cathode electrode, which mitigate the acidic conditions that facilitate the HER over the CO<sub>2</sub>RR in CEM-based devices (vide supra)<sup>169</sup>. This is also advantageous for the counter reaction in the anode, the oxygen evolution reaction (OER), since CEMs' acidic conditions limit the choice of OER electrocatalytic materials to scarce and expensive Ir-oxides, which would jeopardize such system's upscale potential due to their high costs<sup>170</sup>. Instead, such medium-to-high pHs at the anode allow the use of OER-electrocatalysts based on non-precious metals such as Co or Ni, which drastically reduces the electrolyzer price<sup>171–173</sup>.

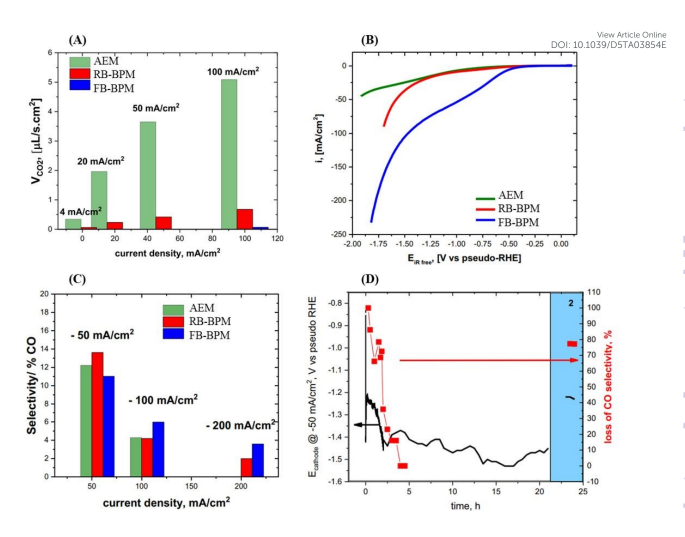
Palladium-based and Cu-based electrocatalysts were tested as cathode electrodes for the ECO<sub>2</sub>RR in AEM-based flow cell electrolyzers<sup>174</sup>. The Pd electrocatalysts showed a FE towards CO of 98 % with a current density of 200 mA cm<sup>-2</sup>. Similarly, Cu electrocatalysts were evaluated in the same conditions, with a FE of 82 % towards hydrocarbons with a current density of 350 mA cm<sup>-2</sup>. The electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> reduction reaction is, however, less

studied in AEM-type electrolyzers. Nevertheless, a few works have proven that the alkamaticle online pH of AEM-type electrolyzer can also produce high selectivity towards CH<sub>4</sub> formation, with FEs of 62 to 73 % <sup>88,175</sup>.

# 4.1.3. Bipolar membrane (BPM)

The above limitations regarding product and ion crossover have recently been reported to be reduced through the use of bipolar membranes (BPMs), which in their simplest version consist of an AEM and a CEM laminated against each other 156,171,176,177. Here is worth mentioning that two configurations can be applied when implementing a BPM in a CO<sub>2</sub>-electrolyzer. First, in the so-called reverse bias (RB) mode that has become the most common configuration in CO<sub>2</sub>-electrolysis, the anion- and cation-exchange layers (AEL, CEL) are disposed at the anode and cathode compartments, respectively (Figure 9). Moreover, water is splitted at the CEM-AEM interface, which therefore acts as H<sup>+</sup> and OH<sup>-</sup> supplier. The protons obtained from the water splitting are transported through the CEL to the ECO<sub>2</sub>RR electrocatalyst and react with the reduced species obtained from the ECO<sub>2</sub>RR. On the other side, OH<sup>-</sup> anions obtained from the water splitted at the BPM junction are transported by the AEL to the OER electrocatalysts, where they react with the oxidized species of the anode electrode.





**Figure 10**: (**A**) Volume flows of CO<sub>2</sub> produced at the anode side of a CO<sub>2</sub>-electrolyzer cell with an Au-based cathode and an IrO<sub>2</sub>-TiO<sub>2</sub> anode catalyst using different membrane configurations. The anode gas composition was analyzed using mass spectrometry at various cell current densities in galvanostatic mode. Cells were operated at 40°C under ambient pressure. The cathode was fed with pure, humified CO<sub>2</sub> and the anode with humidified Ar. (**B**) Cathode polarization curves of CO<sub>2</sub> - electrolysis cells with AEM (Fumasep AA30), RB-BPM (Fumatech FBM, 130μm thickness), and FB-BPM (Fumatech FBM, 130μm thickness) at 50 mV s<sup>-1</sup>, 40 °C and ambient pressure. (**C**) CO selectivities obtained in galvanostatic experiments at various fixed current densities. (**D**) Chronoamperometric measurements at – 50 mA cm<sup>-2</sup> (black line) and corresponding CO selectivity measured by *online* MS (red data) for the FB-BPM. The cathode was fed with 50/50 vol.% CO<sub>2</sub>/Ar, while the anode consisted of Pt/C catalyst fed with pure H<sub>2</sub>. Reproduced from reference <sup>165</sup> with permission from ECS, copyright 2019.

Large current densities have been obtained in the RB-BPM mode. Co phthalocyanine was reported as an excellent electrocatalysts for ECO<sub>2</sub>RR in a RB-BPM-type electrolyzer, reaching a CO FE of 62 % at a current density of 200 mA cm<sup>-2</sup> with aqueous CO<sub>2</sub>-saturated 1 M KOH anolyte<sup>178</sup>. The use of KOH as the anode electrolyte produces cation crossover that can damage the electrochemical devices. If KOH is substituted with pure-water, the current density is reduced to 100 mA cm<sup>-2</sup> for a similar FE at the same potential. Ni-based electrocatalysts were also tested in RB-BPM mode, with lower CO selectivity (FE of 30 %) with high current densities of 100 mA cm<sup>-2</sup> <sup>179</sup>. Interestingly, Cu-based electrocatalysts were also evaluated in

this configuration 180, with significant selectivity towards multicarbon products (C<sub>251</sub>.FE<sub>1</sub>. FE<sub>1</sub>. Fe<sup>2</sup> A03854E %) at a current density of 100 mA cm<sup>-2</sup>. At higher current densities, the HER predominates over CO or multicarbon selectivity.

The second configuration of the BPM electrolyzer is the so-called forward bias (FB) mode, in which the CEM is located in the anode and the AEM is in the cathode, thus contrary to the RB-BPM case (see Figure 9). This approach has gained attention in the last years due to the suppression of CO<sub>2</sub>-crossover drawbacks and the elimination of low pHs in the cathode. Regarding the former, the carbonate and bicarbonate anions generated at the cathode are transported by the BPM's AEM towards the CEM-AEM interface, where they recombine with H<sup>+</sup> obtained from the anodic water electrolysis and transported by the CEM to form water and CO<sub>2</sub>. Experiments have showed the absence of CO<sub>2</sub> gas in the anode electrode, implying that that the gas formed at the AEM-CEM interface diffuses back to the cathode compartment, thus successfully inhibiting net CO<sub>2</sub> crossover, as observed in Figure 10A <sup>165</sup>. However, it is important to note that this suppression of CO<sub>2</sub> crossover may not always be at play. The CO<sub>2</sub> generated at the BPM junction can also diffuse towards the anode, where it accumulates and transitions into the gas phase at the membrane-anode catalyst layer interface<sup>181</sup>. Over time, this accumulation can lead to structural damage, including catalyst layer perforation and BPM delamination, raising concerns about the stability of BPMs in CO<sub>2</sub> electrolyzers<sup>181,182</sup>. Nevertheless, larger current densities were obtained for the FB-BPM configuration (- 150 mA·cm<sup>-2</sup> at 1.7 V) compared to those of the RB-BPM (- 80 mA·cm<sup>-2</sup>) using the same cell components (Figure 10B). Additionally, at a fixed current density of - 100 mA·cm<sup>-2</sup>, a higher selectivity towards CO was obtained with the FB-BPM configuration ( $\approx 6$  %, compared to  $\approx 4$ % for AEM or RB-BPM — Figure 10C)<sup>165</sup>. The stability of this FB mode was also measured at 50 mA cm<sup>-2</sup>, revealing a substantial decrease in CO selectivity within the first 5 h due to water accumulation at the cathode, which progressively blocked CO<sub>2</sub> diffusion towards the electrocatalyst. However, after drying the cathode electrode for 24 h, the CO selectivity A03854E returned again to the initial values, confirming that excess water was responsible for the blockage (operation 2, blue region in Figure 10D) <sup>165</sup>.

In another study in which 1 M KOH was fed at the anode, it was shown that the dominant ion transport mechanism is water dissociation at the layer's interface in RB mode<sup>183</sup>, while the absence of water dissociation in FB mode reduces the applied voltage by around 3 V<sup>183</sup>. However, after 10 min of operation time, a drop in the cathodic potential is observed, which correlates with a decrease in CO selectivity. The authors hypothesize that this effect is due to the formation of potassium-based (bi)carbonate at the AEL-CEL interface<sup>183</sup>.

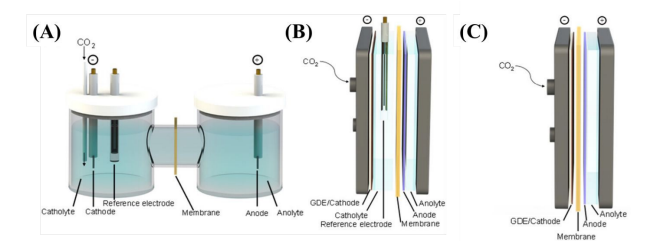
It is worth pointing that the FB-BPM configuration is significantly less studied than AEM or CEM  $\rm CO_2$  electrolysis, as it has only been under study for a few years. In particular, studies on the use of BPMs for  $\rm CO_2RR$  have only been published since  $2016^{184,185}$ , while FB-BPM research dates back to  $2021^{183,186}$ , highlighting the novelty of this approach. In general, only a few works have focused on the development of BPMs for  $\rm CO_2$  electrolyzers since then, but interesting results have already been obtained. BPMs have quickly achieved similar FEs and current densities to AEM and CEM. In such a short period, a few works have reported  $\rm FE_{\rm CO}$  values of 80 -100 % at 100 - 300 mA·cm<sup>-2187–189</sup>. We therefore believe that bipolar membranes possess the greatest potential for improvement due to their very early stage of development, but substantial advancements are still needed to render them fully applicable, especially with regards to their stability.

## 4.3 CO<sub>2</sub> electrolyzers

The design of the CO<sub>2</sub> electrolyzer plays a crucial role in determining the overall performance of the electrochemical reaction, as it directly affects mass transport, ion management, and the system's stability. Different electrolyzer configurations have been developed to address key

challenges such as achieving high current densities, optimizing gas-liquid interactions via high current densities via high curre

CO<sub>2</sub> electrolyzers can be broadly classified into three main categories (Figure 11)<sup>190</sup>: H-type and parallel plate cells, flow cells and zero-gap cells. In the following sections, each of these electrolyzer types will be discussed in detail, highlighting their operating principles, advantages, limitations and recent advancements.



**Figure 11:** Schematic illustration of different CO<sub>2</sub> electrolyzers: (**A**) H-type cell, (**B**) flow cell, and (**C**) zero-gap cell. Reproduced from reference <sup>190</sup> with permission from the American Chemical Society, copyright 2024

#### 4.2.1. H- and parallel-plate cell

H- and parallel-plate type electrolyzers are broadly used as lab-scale reactors for ECO<sub>2</sub>RR-studies due to their straightforward operation and simplicity. Both electrolyzers consists of a two-compartment electrochemical cell, where the cathode electrode is set in one compartment and the anode and reference electrodes are located in a separated one<sup>171,176,191,192</sup>. These cathodic and anodic compartments are separated by an IEM, which prevents the crossover and possible recombination of products during operation. During electrolysis, CO<sub>2</sub> gas is continuously fed into the cathodic reservoir solution, where it is reduced on the electrode surface. The gas output is subsequently directed to a gas chromatograph or a mass spectrometer

for accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate product analysis and quantification. Liquid products, such as CH<sub>3</sub>OH, are collected accurate products and analysis and quantification and analysis and quantification and analysis accurate product accurate product analysis and quantification and analysis accurate product accurate product analysis and quantification accurate products are considered accurate products and accurate product accurate product

This electrolyzer design mainly relies on the dissolution of  $CO_2$  into the cathode compartment electrolyte, whereby the use of a liquid solution limits the concentration of  $CO_2$  and its accessibility to the electrode surface. More precisely, aqueous electrolytes possess a  $CO_2$  solubility of  $\approx 30$  mM, which restricts the attainable current densities to values << 100 mA cm<sup>-2</sup> that cannot be considered as industrially relevant  $^{171,193}$ .

Instead, H- and parallel plate type CO<sub>2</sub> electrolyzers serve as effective tools for evaluating the kinetics of ECO<sub>2</sub>RR electrocatalysts, which is in turn extremely useful when comparing the catalytic performance of different materials. However, this cell design is not up-scalable due to its high electric resistance (typically around 5 Ω·cm<sup>2194</sup>), inefficient mass transfer and low CO<sub>2</sub> solubility, which as discussed above severely limit the currents attainable with such setups. As shown in Table 2, the highest current densities reported with the H-type cell design rarely surpass 40 mA·cm<sup>-2</sup>, and stability tests typically extend only for a few hours. Since this type of reactor is designed exclusively for the fundamental evaluation of catalyst potential, it is not meaningful to perform long-term durability studies, as H-type cells are not envisioned for industrial implementation. Therefore, degradation processes of membranes, catalysts, or other cell components are generally not investigated in these systems.

## 4.2.2. Flow cells CO<sub>2</sub> electrolyzers

In light of the limitations of H-type electrolyzers, flow cells have emerged as a promising solution to enhance CO<sub>2</sub> mass transfer <sup>195</sup>. These cells are structured into three compartments: a gas chamber, a cathodic compartment and an anodic compartment. The cathode electrode is typically constructed with a porous carbon gas diffusion layer (GDL) coated with the ECO<sub>2</sub>RR

electrocatalyst, located between the gas chamber and the cathodic compartment <sup>196</sup> Morrows <sup>196</sup> Noneth <sup>196</sup> In contrast to the conventional H-type CO<sub>2</sub> electrolyze, the gas diffusion medium provides pathways for gaseous CO<sub>2</sub> to directly access the cathode catalyst layer, thus avoiding the above discussed limitations in CO<sub>2</sub> availability intrinsic to its provision as a dissolved species in the electrolyte. As such, the electrolyte in the cathode compartment is also in contact with the ECO<sub>2</sub>RR electrocatalysts, so that the electrochemical reduction of CO<sub>2</sub> proceeds in a solid-liquid-gas interface. This design substantially improves mass transport efficiency and enables higher currents, particularly for the production of CO<sup>197</sup>–<sup>200</sup>, for which current densities of 500 – 1000 mA·cm<sup>-2</sup> with catalysts of different chemical compositions (Ni-, Co-, Cu- or Fe-based, among others) and electrolytes (such as CO<sub>2</sub>-saturated KHCO<sub>3</sub> or KOH solutions) have been demonstrated <sup>197</sup>–<sup>200</sup>. Furthermore, additional works have showcased the potential for hydrocarbon production when selecting appropriate Cu-based electrocatalysts, with current densities reaching more than 1000 mA·cm<sup>-2</sup> <sup>201</sup>–<sup>205</sup>.

Concerning electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion, flow cells have demonstrated large current densities with high CH<sub>4</sub> FE. As previously described, Cu nanoparticles supported on N-doped carbon materials were found to exhibit excellent catalytic performance for ECO<sub>2</sub>RR into CH<sub>4</sub><sup>88</sup>. The electrolyzer configuration consisted of an anion exchange membrane separating anodic and cathodic chambers. For electrochemical measurements, 1 M KOH was used as the electrolyte. At - 1.1 V vs RHE, the FE towards CH<sub>4</sub> formation was found around 70 % with a current density of 230 mA·cm<sup>-2</sup>.<sup>88</sup> As demonstrated in the following section, the authors also conducted electrochemical measurements in a zero-gap cells, with lower FEs but long stability (50 h). Additionally, a Cu(I)-based coordination polymer (NNU-33(S)) underwent a substitution of hydroxyl radicals for sulfate radicals during ECO<sub>2</sub>RR that resulted in an *in situ* dynamic crystal structure transformation to NNU-33(H), which reached a current

density of 391 mA·cm<sup>-2</sup> with a FE towards CH<sub>4</sub> of 82 % at - 0.9 V vs RHE<sup>206</sup>. These resultation continuous electroconversion.

Despite these encouraging results, there are still big challenges for the widespread implementation of large-scale CO<sub>2</sub> flow cells. One notable issue involves the flooding of the electrolyte in the GDL at the cathode gas chamber, causing channel blockages that limit CO<sub>2</sub> transfer from the gas chamber to the catalyst layer. This limitation significantly impacts the ECO<sub>2</sub>RR performance and leads to an increased HER selectivity after only a few hours of operation <sup>176,207</sup>. Furthermore, many of the used electrolytes, especially KOH, tend to produce carbonates and bicarbonates that precipitate during operation, hindering again the exposure to CO<sub>2</sub> through the blockage of the diffusion channels<sup>208</sup>. To address the flooding issue, three main strategies have been proposed<sup>209</sup>. The first is the control of the interface at the GDL-catalyst by using polymeric or hydrophobic substrates that reduce wettability and limit water penetration. The second is to design GDLs with adapted structures that allows excess electrolyte to drain through the electrode and be carried out of the cell with the gas flow. A third approach involves tuning the membrane thickness and CO<sub>2</sub> feed conditions (i.e. humidity) to control water transport and minimize salt precipitation<sup>210,211</sup>.

## **4.2.3.** Zero-gap cells

The high resistance caused by the liquid electrolytes circulated between the cathode and the anode in flow cells lowers the overall energy efficiency of these electrochemical systems<sup>208</sup>. Alternatively, zero-gap cells, also known as catholyte-free or membrane electrode assembly (MEA) cells, have been developed to eliminates the liquid electrolyte in the cathode compartment, thus addressing issues such as electrode flooding and ohmic resistance. The ohmic resistance is significantly reduced, reaching values as low as  $0.3~\Omega \cdot \text{cm}^2$  <sup>212</sup>. The zero-

by an IEM. One of the most promising qualities of zero-gap cells is the scalability of MEA technology, as multiple individual MEA cells can be assembled into a CO<sub>2</sub> electrolyzer stack, making it extremely relevant for industrial-scale processes<sup>208</sup>. In zero-gap cells, gaseous CO<sub>2</sub> (often humidified to enhance the membrane's ionic conductivity) is fed into the cathode

gap cell consists of a compact, sandwich-like design with the anode and cathode separated View Article Online Onli

through a GDE, and the anode electrode is exposed to the electrolyte. It is worth mentioning

that zero-gap cells equipped with AEM are the most commonly used configuration 116,175.

Few works were found about CO<sub>2</sub> conversion to CH<sub>4</sub> in zero-gap cells. The previously mentioned work about Cu nanoparticles supported on N-doped carbon materials showed a CH<sub>4</sub> selectivity above 70 % at -230 mA·cm<sup>-2</sup> in a flow cell reactor<sup>88</sup>. However, this methane FE dropped below 60 % after conducting measurements at the same current density in a CO<sub>2</sub> zero-gap electrolyzer. Both CO<sub>2</sub> electrolyzers employed an anion exchange membrane. Nevertheless, it is worth pointing out that the CH<sub>4</sub> selectivity and the current density in a zero-gap cell did not change substantially for 50 h at a voltage of 4 V, indicating an excellent stability.

Another exciting work can be found about electrochemical CO<sub>2</sub> conversion into CH<sub>4</sub> in a zero-gap cells, using again Cu nanoparticles in a MEA configuration with an anion-exchange membrane<sup>175</sup>. The authors demonstrated that a low coordination number in Cu nanoparticles are beneficial for reducing the hydrogenation energy requirement towards CH<sub>4</sub> production in alkaline conditions, counteracting the traditionally low energy requirements for C-C coupling. This MEA-catalyst combination operated for 110 h at a current density of 190 mA·cm<sup>-2</sup> with an average FE for CH<sub>4</sub> of 56%, which is the largest operating time for CO<sub>2</sub>-to-CH<sub>4</sub> conversion so far.

Despite these advantages, one of the most critical challenges of the zero-gap systems is advantages. precipitation of carbonates and bicarbonates within the catalyst layer and the GDL. Salt precipitation originates from the interplay of CO<sub>2</sub>, hydroxide ions and alkali cations (from the anolyte). Carbonates and bicarbonates accumulate within the catalyst layer, GDL and membrane, progressively blocking gas transport, increasing cell resistance and leading to higher HER selectivity. While dilute electrolytes greatly improve stability, as demonstrated by thousands of hours of continuous operation at moderate current densities<sup>213,214</sup>, they also increase cell resistance and limit performance at higher currents. This means that operating without cations prevents precipitation but drastically lowers activity<sup>213,214</sup>. Beyond avoiding precipitation, other strategies aim to remove salts once formed, typically through regeneration protocols. In these approaches, solvents are periodically injected into the cathode to dissolve the precipitated carbonates and restore CO<sub>2</sub> transport, although this inevitably complicates system operation and questions the viability for continuous industrial processes<sup>215</sup>. In the same line, some authors have proposed electrochemical pulse strategies, where the applied potential is periodically altered to redistribute ions and reverse salt precipitation<sup>215</sup>. Alternating between cell potentials of -3.8 V during operation and -2.0 V during regeneration enabled a 15-fold increase in stability compared to continuous operation without regeneration<sup>215</sup>.

#### 5 Future perspectives

A comprehensive overview of the historical and current landscape of the electrochemical CO<sub>2</sub> reduction reaction has been provided in this review, with particular attention to the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion. ECO<sub>2</sub>RR emerges as a straightforward solution for mitigating anthropogenic CO<sub>2</sub> in the atmosphere while producing value-added products. CH<sub>4</sub>, among all CO<sub>2</sub>-derived products, has arisen as a promising chemical due to the already existing

industrial infrastructure for its rapid implementation and utilization. Technoeconomic analyses active online underscore the near-term viability of the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion, although significant improvements are imperative to compete effectively with traditional CH<sub>4</sub> synthetic routes. The design of electrocatalysts and advancements in CO<sub>2</sub> electrolyzers stand as crucial milestones in the competitiveness of ECO<sub>2</sub>RR. Future perspectives and challenging targets for the successful implementation of electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion are outlined below.

- Metal electrocatalysts, in our assessment, hold promise for the near-term advancement
  in the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> conversion due to their competitive performance.
  Using Cu as active site have demonstrated the most straightforward strategy to obtain
  high selectivity towards CH<sub>4</sub> production. However, special emphasis should be given
  towards tuning the chemical environment of Cu with other metal electrocatalysts to
  enhance the selectivity in CH<sub>4</sub> production.
- Understanding why active sites, such as CuN<sub>4</sub>, exhibit such varied selectivity in ECO<sub>2</sub>RR remains a challenge. Cu single atoms have demonstrated remarkable selectivity towards CH<sub>4</sub>, CH<sub>3</sub>OH and CO, with FE above 70 %, in similar carbon materials. Nonetheless, comprehending the disparities in ECO<sub>2</sub>RR results among studies using comparable materials remains elusive. We understand most of the studies in electrocatalysts design prioritize outcomes over delving into the chemistry involved, however, fine chemistry is clearly playing a definitive role in the ECO<sub>2</sub>RR activity and selectivity. To design better electrocatalysts, the understanding of the electrochemistry that is behind these results is imperative.
- Metal-free carbon-based electrocatalysts have been proposed as promising materials for the ECO<sub>2</sub>RR. However, their ECO<sub>2</sub>RR performance is still far from competitive results. Despite its promising perspective, further efforts are still mandatory for carbon materials to compete with the state-of-the-art metal-containing electrocatalysts.

- Faradaic efficiency (FE) denotes the selectivity towards a particular product in ECO<sub>2</sub>RR. Despite the emphasis on FE in numerous studies, there must exist a crucial need to place greater scrutiny on energy efficiency (EE). EE, expressing the FE per voltage efficiency, provides a more meticulous indicator, especially for technoeconomic aspects. In the case of pathways with high number of electrons transferred, such as CH<sub>4</sub> (8-electron pathway), high voltages are usually required to obtain high selectivity. This is not reflected in FE but becomes essential in EE.
- We noticed the significant gap between fundamental and cell design researchers, which needs to be overcome to fulfill the cost and energy requirements for the electrochemical conversion of CO<sub>2</sub> into not only CH<sub>4</sub>, but all CO<sub>2</sub>-derived products. Fundamental research is imperative to understand kinetics and thermodynamics while producing the most active ECO<sub>2</sub>RR electrocatalysts but lacks practical applications. On the other hand, design of highly efficient electrochemical cells is useless if state-of-the-art electrocatalysts cannot be used in them.
- Advancements in CO<sub>2</sub> electrolyzer are crucial for raising the electrochemical CO<sub>2</sub>-to-CH<sub>4</sub> to competitive levels. H-type electrolyzers are useful electrochemical devices that provides valuable insights into mechanisms and fundamentals when comparing

hampers the industrial applications of these devices. More attention should be directed to flow or zero-gap electrolyzers, capable of providing current densities above 100 mA cm<sup>-2</sup>.

- Progress in ion exchange membranes is essential for the future electrochemical production of CH<sub>4</sub> from CO<sub>2</sub>. Both CEM and AEM have been widely studied in CO<sub>2</sub> electrolysis, showing significant drawbacks that hinder their worldwide application. CEM exhibits high HER in the cathode electrode, which reduces the ECO<sub>2</sub>RR selectivity. At the same time, AEM displays a large crossover issue, allowing CO<sub>2</sub>-products to cross the membrane and recombines at the anode electrode, reducing the energy efficiency.
- Recently, BPM has emerged as a promising alternative to mitigate crossover and pH challenges. However, the low stability under working conditions remains a key factor for the feasibility of these membranes. Particular relevance should be placed to the FB-BPM, as despite its novelty, it has reached comparable current densities to AEM, and RB-BPM. Moreover, in alignment with technoeconomic assessments, BPM are postulated as the most feasible membranes for the next generation of flow electrolyzers.

### **Conflicts of interest**

The authors declare no competing financial interest.

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# Data availability statements

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