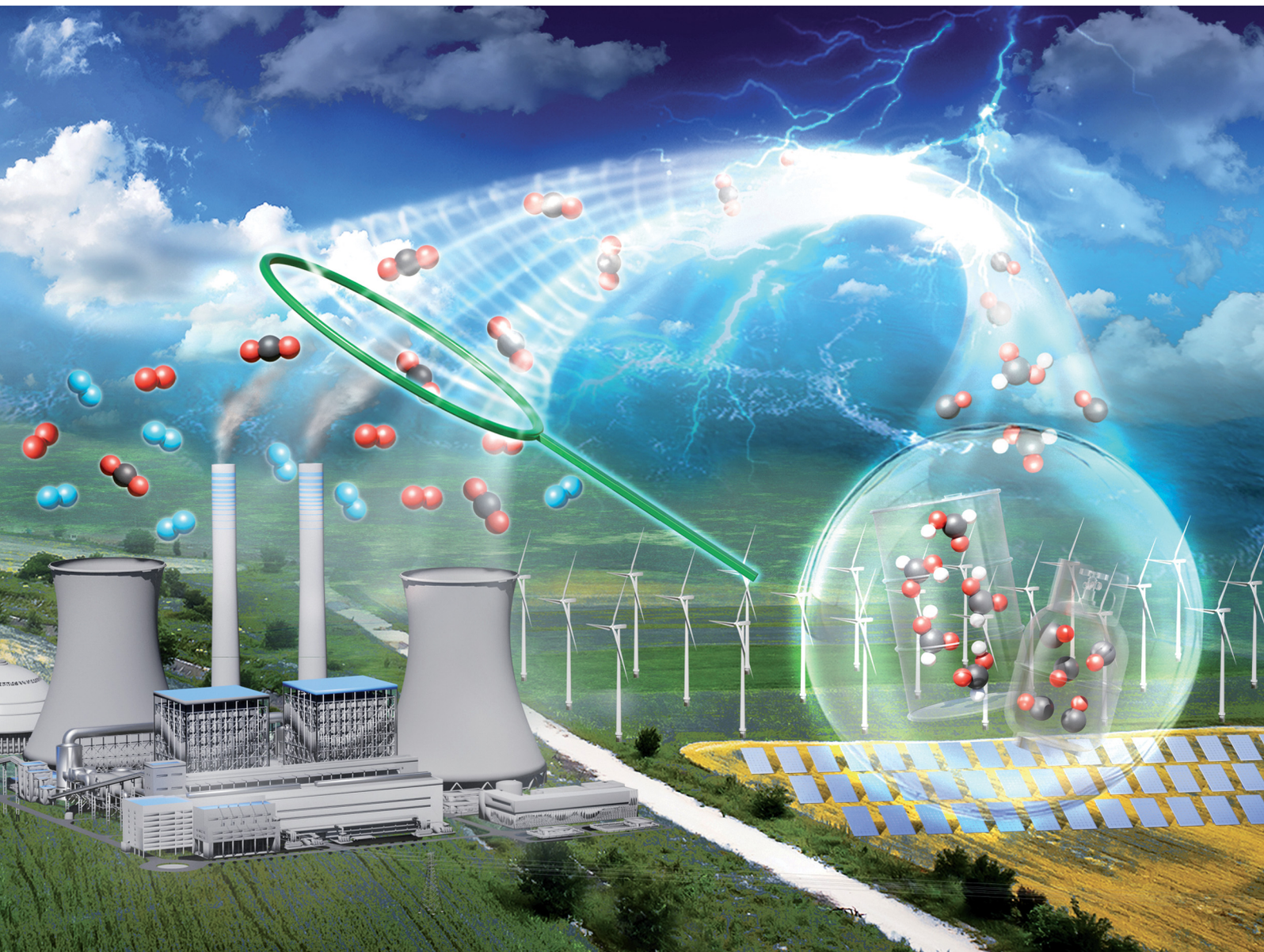


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REVIEW ARTICLE

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Electrochemical valorization of captured CO₂: recent
advances and future perspectives



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Electrochemical valorization of captured CO₂: recent advances and future perspectives†

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The excessive emission of CO₂ has led to severe climate change, prompting global concern. Capturing CO₂ and converting it through electrochemistry into value-added products represent promising approaches to mitigating CO₂ emissions and closing the carbon cycle. Traditionally, these two processes have been performed independently, involving multiple steps, high energy consumption, and low efficiency. Recently, the electrochemical conversion of captured CO₂, which integrates the capture and conversion processes (also referred to as electrochemically reactive CO₂ capture), has garnered increasing attention. This integrated approach bypasses the energy-intensive steps involved in the traditional independent process, including CO₂ release, purification, compression, transportation, and storage. In this review, we discuss recent advances in the electrochemical conversion of captured CO₂, focusing on four key aspects. First, we introduce various capture media, emphasizing the thermodynamic aspects of carbon capture and their implications for integration with electrochemical conversion. Second, we discuss product control mediated by the selection of different catalysts, highlighting the connections between the conversion of captured CO₂ and gas-fed CO₂. Third, we examine the effect of reactor systems and operational conditions on the electrochemical conversion of captured CO₂, shedding light on performance optimization. Finally, we explore real integration systems for CO₂ capture and electrochemical conversion, revealing the potential of this new technology for practical applications. Overall, we provide insights into the existing challenges, potential solutions, and thoughts on opportunities and future directions in the emerging field of electrochemical conversion of captured CO₂.

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1. Introduction

The excessive emission of CO₂, the primary anthropogenic greenhouse gas, has caused severe climate change, presenting one of the most urgent global challenges.^{1,2} The amount of CO₂ released into the atmosphere due to human-related activities is massive, with emissions of ~35 Gt CO₂ just in 2021.³ For this

reason, atmospheric CO₂ levels have surged from 280 ppm before the Industrial Revolution to 420 ppm.⁴ Consequently, global temperatures have risen by ~1 °C compared to pre-industrial times, with projections indicating a potential increase exceeding 2 °C by the end of this century if CO₂ emissions remain unchecked.^{4,5} Climate change has already resulted in severe events on earth, including glacier retreats, sea level elevation, extreme weather events, and ecosystem degradation.

Carbon capture, utilization, and storage (CCUS) technologies have been developed rapidly over the past decades to mitigate CO₂ emissions and reduce atmospheric CO₂ concentration.^{6,7} CCUS typically involves multiple steps, including CO₂ capture from flue gas by amine or other alkaline solutions/materials, release of the captured CO₂ by temperature or pressure swings, followed by CO₂ purification, compression, and transportation before it is finally stored or utilized.⁸ The long-term storage of CO₂ is mainly implemented by mineral carbonation, along with oceanic and underground geological storage.⁹ However, CO₂ can also be utilized directly in food preservation and production, as well as in industrial

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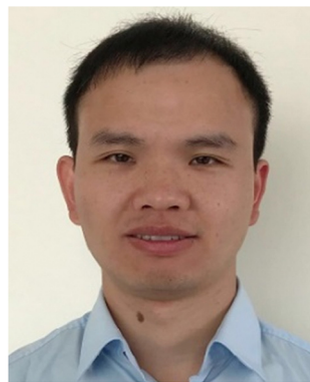
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processes,¹⁰ or indirectly by conversion into various value-added products and fuels through thermochemical,^{11,12} photochemical,^{13–15} biochemical,¹⁶ and electrochemical techniques.¹⁷ Among these techniques, electrochemical conversion of CO₂ stands out due to several advantages such as mild reaction conditions, environmental friendliness, use of renewable electricity, and controllable reaction rate and product selectivity.¹⁸ In particular, gas diffusion electrodes and flow cell configurations have been effectively utilized to boost the productivity of electrochemical CO₂ reduction at industrial-relevant current densities (>200 mA cm⁻²).^{19,20}

To date, the CO₂ capture from diluted sources (0.04–15%) and the follow-up electrochemical conversion have been operated in three distinct routes, *i.e.*, (1) independent, (2) coupled, and (3) integrated (Fig. 1).^{21,22} Route 1 involves multiple decoupled processes, where CO₂ capture occurs independently of electrochemical CO₂ conversion. Route 2 entails CO₂ capture and release stages, immediately followed by the electrochemical valorization of the released CO₂. Route 3 exhibits complete integration between the capture and electrochemical conversion of CO₂, bypassing the CO₂ stripping step required in Routes 1 and 2. For all three routes, producing value-added



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Alonso Rosas-Hernández

Alonso Rosas-Hernández pursued his education at the National Autonomous University of Mexico (UNAM), where he obtained his BS and MS degrees in Chemistry. He then completed his doctorate in 2017 at the Leibniz Institute for Catalysis (LIKAT) under the supervision of Prof. Matthias Beller. Following this, he joined Prof. Jonas Peters' group at the California Institute of Technology (Caltech) as a postdoctoral scholar. In 2021, he

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Kim Daasbjerg

Kim Daasbjerg obtained his PhD in 1993, conducting research in organic electrochemistry under the supervision of Prof. Henning Lund at Aarhus University. After completing his doctoral studies, he undertook postdoctoral research at the Royal Institute of Technology in Stockholm in 1994, working with Prof. Gábor Mérenyi. He then began his academic career at Aarhus University, where he earned a Doctor of Science degree in 2006

and was appointed Professor in 2010. His research expertise spans electrochemistry, polymer brushes, responsive polymers, smart hybrid materials, and composites incorporating carbon substrates such as graphene. In recent years, he has leveraged his combined expertise in these fields to tackle the scientifically complex and socially important challenge of converting carbon dioxide, a key greenhouse gas, into valuable building blocks for the chemical industry and energy sector.



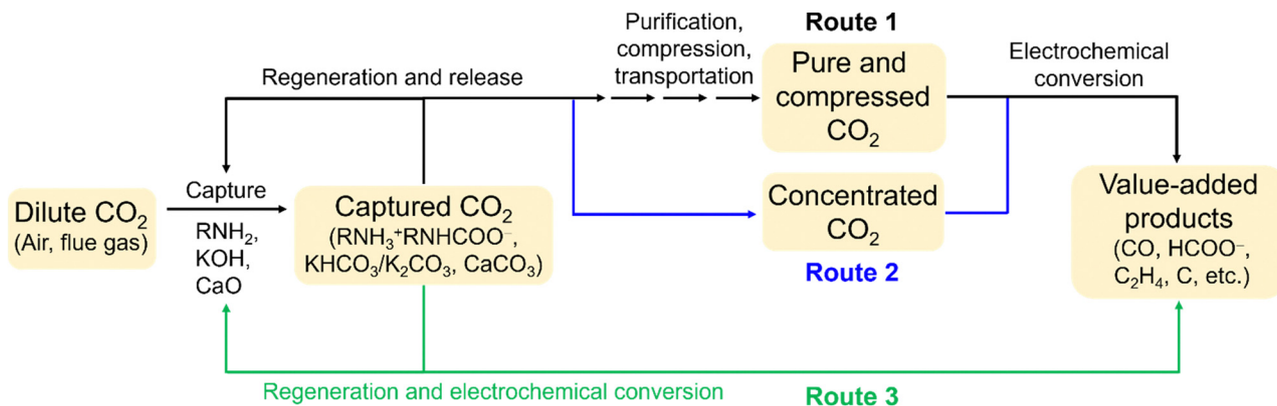


Fig. 1 Three routes for CO₂ capture and electrochemical conversion: (1) independent, (2) coupled, and (3) integrated.

chemicals and fuels helps offset capture expenses, rendering commercial deployment economically more feasible.

1.1. Route 1 (independent)

The independent systems inherently incur energy penalties and costs associated with CO₂ release, purification, compression, and transportation. In particular, the CO₂ desorption step can be energy-intensive if thermal regeneration at high temperatures is required. Nevertheless, Route 1 configuration can emerge as a viable scenario, considering the potential incompatibility in operating conditions between capture technologies that utilize temperature or pressure swings and the electrochemical conversion of CO₂ under ambient conditions. This configuration offers flexibility for independently optimizing the capture and conversion stages. However, it is important to note that standalone electrochemical systems using gas-fed CO₂ often face a significant challenge due to the typically low single-pass CO₂ conversion efficiency (<20%).^{23–25} Consequently, this leads to low product concentration and significantly escalates downstream separation/purification costs.^{26,27}

1.2. Route 2 (coupled)

An advantage of the coupled CO₂ capture and conversion system is that it bypasses the CO₂ purification, compression, and transportation steps, thereby reducing the complexity and energy consumption of the overall process. Here, the CO₂ for the downstream conversion is directly supplied by the upstream CO₂ capture unit, which usually releases CO₂ by electrochemically driven redox couples or pH swings.^{28,29} Note that the coupled cascade system still involves independent CO₂ capture and conversion, as well as the energy-intensive CO₂ release step. Accordingly, the problem of low single-pass conversion efficiency for CO₂ exists. In addition, by enhancing the overlap between the capture and conversion steps, fewer degrees of freedom are available for optimizing the electrocatalysts toward higher activity and selectivity for a targeted CO₂ reduction product.

1.3. Route 3 (integrated/reactive capture)

Compared with Route 2, these systems offer the advantage of reducing the total energy requirements for capture and

conversion by eliminating CO₂ release and many other follow-up steps, which typically incur a significant energy penalty. By directly converting the captured CO₂ at the electrode surface, the regeneration of the capture media and the production of valuable products occur simultaneously. In this sense, Route 3 integrates the two processes of CO₂ capture and follow-up electrochemical conversion and has also been coined electrochemically reactive capture of CO₂.^{30–32}

As a result, Route 3 represents a more straightforward and sustainable approach for CCUS than the other two routes, at least at first glance. This unique advantage has stimulated increasing activities in the study of direct electrochemical valorization of captured CO₂ in the past several years. For instance, several reaction media, such as amine/carbamate solution, hydroxide/carbonate/bicarbonate solution, and molten oxide/carbonate, have proven effective in mediating CO₂ capture and subsequent electrochemical conversion. This offers the opportunity to directly transform CO₂ from flue gas or the air into valuable products such as carbon monoxide, formate, methane, and ethylene. Despite the significant advances and great prospects, several critical challenges still need to be addressed to advance such an emerging technology for large-scale applications.

These challenges lie in the unclear mechanism and low efficiency of the conversion of captured CO₂, poor integration of CO₂ capture and conversion processes, and others. In this context, a timely and thorough review article summarizing the progress and discussing the challenges in this emerging field would benefit the community.

In this review, we summarize the recent advances in the strategy of reactive CO₂ capture, with a specific focus on the electrochemical conversion of captured CO₂. Techno-economic analyses comparing the three different routes for CO₂ capture and conversion can be found in recent reviews and are thus not covered in this paper.^{30,33} Moreover, readers with interests in related topics such as electrochemical CO₂ capture^{34,35} and electrochemical CO₂ conversion³⁶ are referred to the respective reviews.

This review is structured around four aspects of the technology: capture media, electrocatalysts, system optimization, and practical considerations for continuous operation (Fig. 2). We



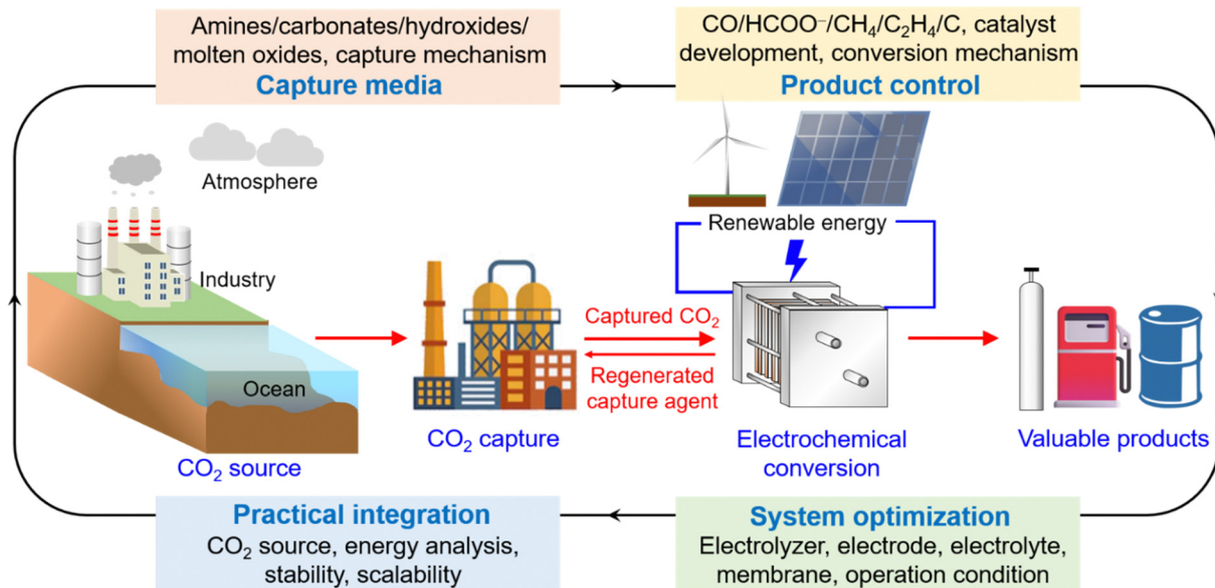


Fig. 2 Themes covered in this review on the electrochemical conversion of captured CO₂.

start by introducing various types of capture media, emphasizing the thermodynamic aspects of carbon capture and their implications for integrated Route 3 processes. We move on to examine different electrocatalyst compositions and morphologies, discussing their influence on product distribution from the conversion of captured CO₂ and its relationship with the conversion of gas-fed CO₂. Next, we explore the effects of reactor systems and operational conditions on the electrochemical conversion of captured CO₂, aiming to dissect the key factors determining the conversion performance. Finally, we present examples of integrated CO₂ capture and conversion processes. Furthermore, we offer insights into the challenges and potential solutions and present our perspectives on the opportunities and future directions for integrating CO₂ capture and electrochemical conversion. This exploration uncovers promising avenues for practical applications of this emerging field in achieving more sustainable CO₂ capture and conversion, potentially contributing to mitigating global climate change.

2. Properties of capture media

The electrochemical conversion of captured CO₂ from diluted sources marks a significant advancement in the field of CCUS. Historically, CO₂ capture and subsequent electrochemical valorization were treated as separate research domains. The electrochemical valorization has primarily focused on novel electrocatalyst development, electrolyte selection, cell configuration engineering, and using gas-fed concentrated CO₂ without considering the carbon source (Routes 1 and 2 in Fig. 1). However, the emergence of highly efficient electrochemical systems for CO₂ reduction over the past decade has reshaped the field.^{37,38} Consequently, the CO₂ capture step and the

conversion of resulting captured CO₂ have begun to be integrated to optimize the system (Route 3 in Fig. 1). This section explores the various capture media wherein the captured CO₂ can be utilized for electrochemical conversion, encompassing aqueous amines, alkaline solutions, and molten oxides. In this section, our focus is directed toward the underlying mechanisms of carbon capture and the thermodynamic aspects of chemical CO₂ sorption.

2.1. Amine solutions

The fundamental process of CO₂ removal from a gas stream leveraging its reversible binding with amines was discovered in 1930.³⁹ Initially utilized to separate CO₂ from natural gas and hydrogen, such a chemical reactivity has since been employed in amine scrubbing for post-combustion capture of CO₂ on small-scale gas and coal-fired plants.⁴⁰ Currently, amine scrubbing is the most cost-effective and widely used technology for CO₂ capture from industrial flue gases in power plants, oil refineries, cement production, and other applications. This is due to the high absorption rate and large CO₂ capacity of amines. For example, NRG Energy Inc. and JX Nippon Oil & Gas Exploration Corp. have launched a post-combustion carbon capture project called Petra Nova, which aims to remove over 90% of the CO₂ from the flue gas of a coal-fired power plant in Texas, USA using an amine solution.⁴¹ However, it is important to note that the high temperatures in flue gas environments can lead to amine losses due to thermal degradation and evaporation. Furthermore, the presence of poisoning gases such as SO_x and NO_x can decrease the yield of captured CO₂. While amine scrubbing is often hailed as the most cost-effective post-combustion CO₂ capture technology, it remains relatively expensive in terms of scalability. Therefore, ongoing research endeavors aim to enhance amine scrubbing processes, address their limitations, and optimize efficiency.



The capture of CO₂ using amine-based technologies involves two primary steps: absorption of CO₂ and CO₂-stripping for amine regeneration. The process is initiated by directing a CO₂-diluted gas stream at or near ambient temperature through the amine solution. During this phase, CO₂ is captured *via* a thermodynamically favorable reaction with the amine, yielding carbamate and ammonium ions (eqn (1)). Amines generally display high values of absorption enthalpies (ΔH_{abs}), ranging from -90 to -60 kJ mol⁻¹ when flue gas (12–15% CO₂ concentration) is employed.⁴² Subsequently, the amine undergoes regeneration through stripping with water vapor at temperatures ranging from 100–120 °C, which shifts the equilibrium in eqn (1) toward the reactants. Following this, water is condensed from the stripper vapor, leaving behind pure CO₂. The main energy demand in the process comes from stripping CO₂ from the CO₂-rich amine stream, which requires 115–140 kJ mol-CO₂⁻¹. This step consumes more energy than is released during the absorption phase because it produces a higher-purity CO₂ stream than the original flue gas. Thus, the critical thermodynamic parameters governing the amine scrubbing process revolve around the chemical interaction between the amine and CO₂.



Different strategies have been employed to balance the efficiency of the chemical absorption step and the total energy consumption for the amine regeneration. Predictably, enhancing the chemical interaction between CO₂ and the amine to improve absorption rates leads to increased energy input required to break the C–N bond of the carbamate group during amine regeneration, which is translated to the use of higher temperatures during the stripping step. For instance, primary and secondary amines display a very high heat of absorption since they can quickly generate chemically stable carbamates during their reaction with CO₂.⁴³ However, their regeneration energy requirement is high. In contrast, tertiary amines have a low heat of absorption and need less regeneration energy for solvent recovery, as bicarbonate, instead of carbamate, is formed for CO₂ capture with tertiary amines.

The most commonly used amine-based capture medium is a 30 wt% aqueous monoethanolamine solution, valued for its high CO₂ absorption rate, large absorption volume, and low cost.⁴⁴ Other frequently employed amines include diethanolamine and methyldiethanolamine, which are examples of secondary and tertiary amines, respectively. Although these aqueous amine solutions are widely used for CO₂ absorption, they have drawbacks, such as amine loss and thermal decomposition. A reported strategy to overcome such issues is mixing monoethanolamine with other secondary or tertiary amines with low energy consumption for regeneration.⁴⁵ This results in a mixture in the absorber unit that simultaneously displays high absorption capacity, high absorption rate, and low regeneration energy consumption. For instance, a 2:1 mixture of monoethanolamine and diethylenetriamine has been reported to increase CO₂ absorption capacity by 53% and removal

efficiency by 31% with respect to the commonly used 30 wt% monoethanolamine aqueous solution.⁴⁶

Since 2010, amine scrubbing has been successfully applied to capture CO₂ from natural gas, hydrogen, and other gases with low oxygen content.⁴⁰ For instance, aqueous monoethanolamine solution can capture CO₂ from flue gas of coal-fired plants.⁴⁷ Unfortunately, it has displayed oxidative and thermal degradation under these conditions. When the oxygen content in the gas stream is in the range of 15%, a more concentrated amine solution (30 wt%) is required to increase the rate of CO₂ capture, which is inhibited by the oxidation of the amine in the presence of oxygen. Additionally, thermal degradation has been minimized by operating at lower temperatures (100 °C).⁴⁸

Stern *et al.* proposed in 2013 an alternative process that uses an electrochemical swing based on Cu²⁺/Cu to strip CO₂ from the carbamate adduct and regenerate the amine.⁴⁹ The electrochemically mediated amine regeneration (EMAR) cycle leverages the high stability of the Cu²⁺-amine complexes compared to the carbamate, thus favoring the dissociation of CO₂ from the latter. Formation constants of cupric amine complexes exhibit values of 10¹⁸ for polyamines such as ethylenediamine, while binding constants between amines and CO₂ typically fall in the order of $\sim 10^3$.⁵⁰

Currently, the EMAR technology has potential for large-scale applications in systems following Route 2 in Fig. 1. One of its most notable advantages is its ability to achieve high amine utilization, reducing the capital costs for auxiliary equipment due to lower circulation rates.⁵¹ Additionally, the EMAR technology requires less energy input since the capture and strip of CO₂ are performed at room temperature. Since the process requires only electrical energy, decentralized devices can be developed at point sources to capture and convert CO₂ as needed.

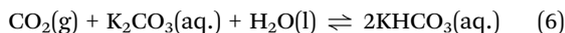
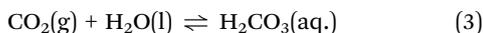
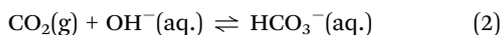
2.2. Aqueous alkaline solutions

CO₂ capture in many industrial processes has been successfully performed using inorganic salt solutions of various weak acids.⁵² The most commonly employed salts are sodium and potassium carbonate, which, due to their alkaline pH (typically 9–11), absorb acid gases such as CO₂. For instance, the traditional Benfield process, which has been in use since the 1950s, was developed as an alternative to amine scrubbing and uses hot K₂CO₃ solutions to absorb CO₂ from a synthesis gas under relatively high CO₂ partial pressures and high temperatures.⁵³

The capture of CO₂ using aqueous K₂CO₃ solutions depends on the established equilibria when CO₂ dissolves in water and how these equilibria are affected by pH changes. In open systems, increasing the pH raises the total concentration of dissolved carbon, while lowering the pH causes CO₂ to be released, thereby decreasing CO₂ absorption. When CO₂ passes through aqueous K₂CO₃, it can react *via* two pathways. First, bicarbonate ions can be directly formed from the reaction of CO₂ with hydroxide ions (eqn (2)). Alternatively, CO₂ can be hydrated by water to produce carbonic acid (eqn (3)), which then deprotonates to form bicarbonate (eqn (4)). In either case, the resulting decrease in pH is buffered by the equilibrium



between the carbonate and bicarbonate species (eqn (5)). Overall, K_2CO_3 in water captures CO_2 as potassium bicarbonate (eqn (6)), with a ΔH_{abs} value of approximately -40 kJ mol^{-1} .⁵⁵



Either eqn (2) or (3) represents the process's rate-limiting step, depending on pH, as the deprotonation/protonation reactions in aqueous systems (eqn (4) and (5)) are considered barrierless. Accordingly, in solutions with $pH > 9$, the generally faster reaction between CO_2 and hydroxide ions dominates. In contrast, in solutions with $pH < 7$, the slower CO_2 hydration reaction becomes more prominent. For K_2CO_3 solutions, where the pH is usually > 9 , the reaction with hydroxide ions predominates and serves as the rate-determining step.

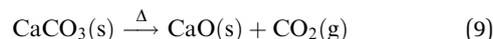
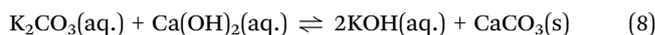
The release of CO_2 from $KHCO_3$, along with the regeneration of K_2CO_3 (reverse of eqn (6)), takes place through a pressure swing between the absorption and desorption units, eliminating the need for an additional temperature swing. This engineering approach enables absorption and regeneration at the same elevated temperature ($\sim 100 \text{ }^\circ\text{C}$), improving absorption kinetics and reducing energy consumption by nearly half.⁵³

Aqueous K_2CO_3 solutions offer several advantages over traditional amine-based absorption liquids, including lower toxicity and nonvolatility, which minimize absorbent losses and decrease solvent replacement costs. Most importantly, carbonates exhibit resistance to absorbent degradation and are relatively inexpensive compared to many amines. A practical challenge associated with the carbonate absorption medium is that the precipitation of bicarbonate salts limits the use of highly concentrated solutions. However, recent research has shown that purposeful precipitation of bicarbonate salts can facilitate high-pressure stripping processes, reducing the regeneration energy requirement and liquid circulation rates.⁵⁴ For example, an Australian company, KC8 Capture Technologies, uses a K_2CO_3 solution to capture 90–95% of CO_2 emissions from flue gas, forming $KHCO_3$, which can then precipitate and be subjected to CO_2 release and K_2CO_3 regeneration.⁵⁵ The regeneration energy requirement of the technology is 88–110 $\text{kJ mol}^{-1}CO_2$, which is lower than that typically needed for amine-based processes. Another challenge for the carbonate absorption medium is its slow kinetics for CO_2 absorption, which can be improved by adding activators or promoters such as amines^{53,56} and enzymes.^{57,58}

Analogous to carbonate solutions, aqueous hydroxide solutions can also be used to capture CO_2 , but in this case, carbonate salts are produced as described in eqn (7). The produced carbonate can continue to capture CO_2 to form bicarbonate.



Initially proposed in 1999 as a solution for direct air capture (DAC), the high absorption rate of this process results from the large thermodynamic driving force of the reaction between dissolved CO_2 and hydroxide ions ($-\Delta H_{abs} = 90\text{--}100 \text{ kJ mol}^{-1}$).⁵⁹ Although the original process was limited by reduced capture capability due to the low solubility of the used $Ca(OH)_2$, most recent implementations of this technology use KOH , which has a solubility in water over two orders of magnitude greater than that of $Ca(OH)_2$.⁶⁰ Once K_2CO_3 is formed in the absorption chamber (eqn (7)), the next step is the precipitation of calcium carbonate and the regeneration of reusable KOH in solution (eqn (8)). The calcium carbonate is then subjected to a calcination process at $\sim 900 \text{ }^\circ\text{C}$ to liberate CO_2 and generate CaO (eqn (9)), which generates the calcium hydroxide required in the precipitation chamber by reaction with water. Industrial applications of this technology for DAC have shown promising results by achieving a capture rate of $1 \text{ Mt } CO_2 \text{ year}^{-1}$ (Carbon Engineering ULC.).⁶¹ However, this plant needs high-grade heat ($230 \text{ kJ mol}^{-1}CO_2$), which is supplied by burning natural gas.



2.3. Molten metal oxides

Capture of CO_2 using alkali and alkaline-earth metal oxides has been employed in large-scale processes for over 150 years. Its first application involved purifying hydrogen by removing CO_2 with CaO in a gasification process. However, it was not until 1999 that Shimizu *et al.* introduced the carbonate looping process, which paved the way for CO_2 capture by metal oxides at medium to high temperatures ($> 200 \text{ }^\circ\text{C}$).⁶² In this process, CaO reacts with post-combustion CO_2 to form $CaCO_3$ (eqn (10)), followed by its thermal decomposition to CaO and the release of pure CO_2 . Compared to liquid amine scrubbing, the higher operating temperatures of metal oxides provide the advantage of incurring lower energy penalties when the carbon capture process is implemented into power plants or other high-temperature processes.⁶³

The most used metal oxide sorbents utilize alkaline-earth metals such as calcium and magnesium to capture CO_2 by forming the corresponding carbonates (eqn (10) and (11)). Although the carbonation reactions for CaO and MgO are exergonic, relatively high operating temperatures are required for the CO_2 capture step to achieve adequate CO_2 capture rates. This ensures that the carbonation reaction is primarily controlled by thermodynamics, leading to high process efficiencies. For example, continuous CO_2 capture from ambient air has been demonstrated using CaO particles at temperatures ranging from 365 to 400 $^\circ\text{C}$, achieving a CO_2 removal efficiency of 99.9% after five consecutive cycles.⁶⁴



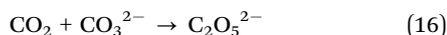
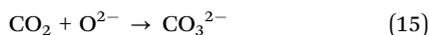
Using oxides of alkali metals such as Li , Na , and K as sorbents is not often considered for large-scale CO_2 capture



applications. The formed carbonates (eqn (12)–(14)) typically require temperatures exceeding 1200 °C to release the absorbed CO₂ and regenerate the metal oxides. However, these monovalent oxides have played an important role in altering the CO₂ sorption mechanisms of CaO and MgO.⁶⁵ When used as promoters, the alkali metal oxides readily transform into their carbonate forms in the presence of CO₂ with ΔH_{abs} of approximately -300 kJ mol^{-1} .⁶⁵ The melting points of Li₂CO₃, Na₂CO₃, and K₂CO₃ are relatively low, ranging from 730–900 °C, and eutectic mixtures of the three components possess melting points as low as ~ 400 °C. Taking advantage of these low melting points, a strategy has been devised to enhance the rate and extent of CO₂ uptake by CaO and MgO-based sorbents. This involves coating sorbents with binary or ternary molten alkali carbonates, which enhances CO₂ transport within the sorbent.



Huang *et al.* reported the remarkable effects of alkali carbonate molten salts on the performance of CaO particles for CO₂ capture.⁶⁶ The (Li–K)₂CO₃ molten salt coating was found not only to promote CO₂ uptake but also to facilitate CO₂ desorption from CaO. Particularly at low temperatures of 500 and 600 °C, the CO₂ capture capacity increased significantly from 1.19 and 3.26 mmol g⁻¹ to 6.93 and 10.38 mmol g⁻¹, respectively, using 7.5 mol % (Li–K)₂CO₃. The coating of alkali carbonate molten salts is believed to prevent the formation of a rigid CaCO₃ layer on the surface of the CaO particles. This facilitates the ongoing reaction of dissolved CO₂ with O²⁻, originating from CaO, in the liquid molten salts, resulting in the formation of CO₃²⁻ (eqn (15)). The (Li–K)₂CO₃ molten salt is particularly effective in enhancing CO₂ uptake due to the high O²⁻ migration, which ensures the continuous counter diffusion of CO₃²⁻ and O²⁻.⁶⁷ Furthermore, CO₂ can be captured by the reaction with the continuously delivered CO₃²⁻ to form dicarbonate ions (eqn (16)).



In 2015, Harada *et al.* disclosed that coating MgO particles with molten alkali metal nitrates could significantly improve the CO₂ uptake,⁶⁸ which increased from $<1 \text{ mmol g}^{-1}$ for uncoated MgO to 10.2 mmol g^{-1} at 300 °C with the coating. They also demonstrated the molten alkali metal nitrates prevented the formation of a rigid carbonate layer on the surface of MgO, which would otherwise be impermeable to CO₂. The high concentration of oxide ions in the molten salt restricted the generation of this rigid layer and facilitated the generation of carbonate ions (eqn (11)).⁶⁹ This resulted in the rapid formation of MgCO₃ and eased the regeneration of particles at moderate temperatures.

2.4 Electrochemical reactions of captured CO₂

The physicochemical properties of each capture medium will affect the thermodynamics of the electrochemical conversion of the corresponding CO₂-adduct to any of the possible products. As depicted in Fig. 3, the electrocatalytic conversion of the captured CO₂ is an energetically uphill pathway, with the initial energy level determined by the CO₂ absorption enthalpy of the capture medium employed. For instance, the electrochemical conversion of bicarbonate, derived from the CO₂ capture with aqueous carbonate solutions (eqn (6)), is less endothermic than the direct conversion of the carbamate obtained from the reaction between CO₂ and aqueous amines (eqn (1)). This is translated to a higher energy input for the electrochemical conversion of the carbamate as the result of the stronger interaction between CO₂ and the capture medium.

The electronic and steric properties of the CO₂-adduct are also crucial in determining the kinetics of the electrochemical conversion of captured CO₂. The energies of intermediates involved in the pathways leading to the reduced products are influenced by the chemical structure of the captured CO₂ species, thereby affecting the reaction kinetics. Moreover, the chemical composition and morphology of the catalyst can alter the binding energies of such reaction intermediates, consequently impacting reaction rates and favoring specific products. Thus, there is a complex interplay between the nature of the captured CO₂ species and the electrocatalyst, collectively determining the kinetics for generating the products.

For instance, carbamates can be directly converted at the cathode, producing CO or other carbon-based products while regenerating the amine for another capture cycle (eqn (17)). The efficiency of the electrochemical conversion of CO₂ captured as carbamate can be affected by several factors. First, the energy required to break the C–N bond in the carbamate group increases with stronger binding interactions between CO₂ and the amine, which raises the overpotential of the electrocatalytic reaction. Moreover, the selectivity toward carbonaceous

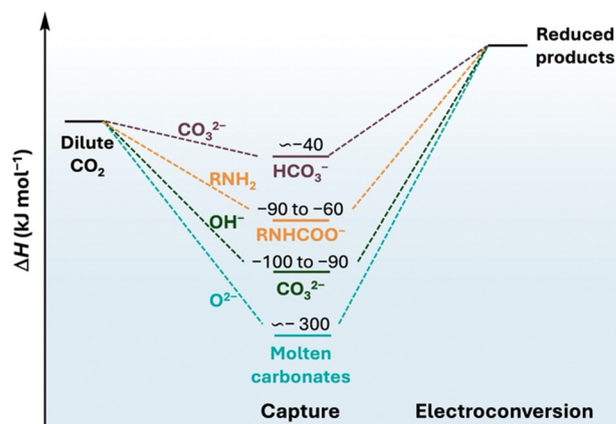
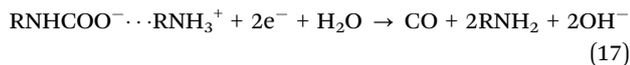


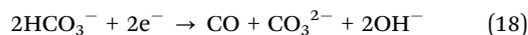
Fig. 3 Schematic representation of the energetics for the electrochemical conversion of captured CO₂. Enthalpy changes in CO₂ absorption are shown for each capture medium. For ease of representation, energy levels are not to scale.



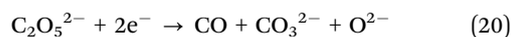
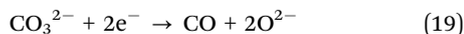
products can be adjusted by changing the concentration of the amine solutions to improve the mass transport of the amine-CO₂ adduct to the electrode, thereby suppressing the undesired hydrogen evolution reaction (HER). Also, considering that the electrochemical conversion of carbamate follows an inner-sphere electron transfer mechanism,⁷⁰ the distance between the carbamate and the electrode surface is crucial in determining the kinetics of the electrocatalytic reaction. Consequently, the structure of the amine will significantly influence the efficiency of carbamate conversion.



In the case of CO₂ capture using hydroxide or carbonate solutions, bicarbonate is the final product (eqn (2) and (6)). The bicarbonate solution can undergo electrochemical reduction to obtain CO or other products while regenerating the initial carbonate ions (eqn (18)). A challenging aspect of bicarbonate reduction systems is resolving the electrochemically active species, considering the dynamic acid-base equilibria that define the relative concentrations of CO₂, bicarbonate, and carbonate at the electrode surface (eqn (2)–(5)).



The CO₂ capture by molten oxide generates molten carbonate (eqn (15)), which can be reduced at elevated temperatures (400–900 °C) at the cathode to form CO or other products while regenerating the oxide (eqn (19)). Similarly, dicarbonate ions can be directly reduced at the cathode, regenerating the carbonate and oxide ions (eqn (20)). These electrochemical transformations are unique since they are carried out at much higher temperatures than the conversion of carbamate or bicarbonate solutions. It has been shown that adjusting the temperature can effectively control the selectivity of the electrocatalytic reactions.⁷¹



The electrocatalytic conversion of CO₂ captured by various capture media, as described above, facilitates the conversion of CO₂ from diluted sources into value-added products. Producing carbonaceous products other than carbon monoxide, such as formate, ethylene, and carbon materials, entails more complex reaction pathways. The next section will explore these topics in detail.

3. Catalyst-mediated product control for converting captured CO₂

The electrocatalyst is crucial in converting captured CO₂, affecting the type and distribution of products, and the conversion efficiency. Recent progress in the electrochemical valorization of captured CO₂ has made it possible to produce various products, including CO, HCOO⁻, CH₄, C₂H₄, C₂H₅OH, and carbon materials, as summarized in Table 1.

These products are important chemicals, fuels, or materials with many industrial applications. Carbon monoxide is a key component of syngas, which is extensively used to synthesize long-chain hydrocarbons through the Fischer-Tropsch process.¹¹⁴ In addition, CO is widely utilized in synthesizing various carbonyl compounds *via* carbonylation reactions.¹⁸ Formate and formic acid are used in leather and textile manufacturing, serve as preservatives in livestock feed, and act as efficient H₂ carriers in fuel cells.¹¹⁵ Methane, the primary component of natural gas, serves as a gas fuel for electricity generation, heating buildings, transportation, and industrial processes. Ethylene is an important building block for industrial chemical production, particularly in the manufacture of polyethylene, the most widely used plastic worldwide.¹¹⁶ Ethanol is commonly used as a solvent, disinfectant, precursor, liqueur, and liquid fuel. Carbon materials, with varying morphology and physicochemical properties, are extensively utilized in applications such as adsorption, separation, and catalysis.¹¹⁷

3.1. Mechanism of conversion of captured CO₂

There are two possible pathways for the electrochemical valorization of captured CO₂, each influencing catalyst selection. (a) An indirect pathway, where the captured CO₂ is first converted back to CO₂ at the electrode-electrolyte interface and then undergoes electrochemical reduction (Fig. 4a); (b) a direct pathway, where the captured CO₂ is directly reduced at the electrode (Fig. 4b).

In the indirect conversion, the captured CO₂ (such as bicarbonate or carbamate) dissociates, producing CO₂ *in situ* at the electrode-electrolyte interface (Fig. 4a). This dissociation process is typically triggered by either temperature or proton flux changes within the electrolyzer. Currently, most reported instances of electrochemical conversion of captured CO₂ in aqueous solutions follow this indirect pathway (Table 1).^{72–74} Because of the *in situ* CO₂ generation, the catalyst selection closely aligns with conventional electrochemical CO₂ conversion, *i.e.*, Ag and single-atom catalysts (SACs) facilitate CO production, Bi and Sn catalyze formate formation, while Cu-based catalysts are suitable for producing hydrocarbons and alcohols. The key aspect of the indirect pathway hinges on balancing the rate of *in situ* CO₂ release with the rate of CO₂ reduction at the electrode surface to achieve the conversion of the captured CO₂ with high activity and selectivity. This requires precise design of the catalyst structures and properties, as well as optimization of the electrolyzer, electrolyte, and other operating conditions.

The second pathway is the direct conversion, which bypasses the formation of CO₂ before the reduction process (Fig. 4b). Here, the captured CO₂ (such as bicarbonate or carbamate) serves as the primary reactant, setting this approach apart from conventional electrochemical CO₂ conversion. One significant advantage of focusing on (bi)carbonate and carbamate instead of CO₂ is achieving much higher reactant concentrations. So far, only a few documented examples have utilized the direct pathway in aqueous bicarbonate and carbamate, primarily under high-temperature



Table 1 Recent progress of catalyst-mediated product control in the electrochemical conversion of captured CO₂ (since 2019)

Cathode materials	Captured CO ₂	Electrolyzer	Pathway	Products	FE (%)	$ j $ (mA cm ⁻²) ^a	T (°C) ^b	Ref.
Ag nanoparticles (NPs) on Ag film	K ₂ CO ₃	MEA with BPM	Indirect	CO	28	100	R.T.	72
Ag NPs	K ₂ CO ₃	MEA with cation exchange membrane (CEM)	Indirect	CO	46	200	R.T.	73
Ag NPs	K ₂ CO ₃	MEA with CEM	Indirect	CO	42	200	R.T.	74
Ag NPs-coated porous carbon	KHCO ₃	MEA with BPM	Indirect	CO	37	100	R.T.	75
Ag NPs	KHCO ₃	MEA with BPM	Indirect	CO	82	100	R.T.	76
Ag NPs	KHCO ₃	MEA with BPM	Indirect	CO	~65	200	R.T.	77
Ag NPs	KHCO ₃	MEA with CEM	Indirect	CO	99	51.5	R.T.	78
Ag (free-standing porous electrode)	KHCO ₃	MEA with BPM	Indirect	CO	59	100	R.T.	79
Ag foam	KHCO ₃	MEA with CEM	Indirect	CO	40	100	R.T.	80
Ag foam	KHCO ₃ + DTAB ^c	MEA with BPM	Indirect	CO	80	100	R.T.	81
Ag (electrodeposited)	KHCO ₃ + DTAB ^c	MEA with BPM	Indirect	CO	85	100	50	82
Ag (Br-modified)	NH ₄ HCO ₃	H cell	Indirect	CO	77.8	18 ^d	R.T.	83
Ag/C	Captured CO ₂ from the air using KOH solution	MEA with BPM	Indirect	CO	13	50	R.T.	84
Ag NPs on a Ag film	CO ₂ -captured monoethanolamine	Flow cell	Direct	CO	72	50	60	85
Ag (Cl-modified)	CO ₂ -captured 2-amino-2-methyl-1-propanol (AMP)	H cell	Indirect	CO	91	6	R.T.	86
Au (foil)	CO ₂ -captured AMP	Flow cell	Indirect	CO	45	10	75	87
Ni SAC	CO ₂ -captured monoethanolamine	MEA with BPM	Indirect	CO	64.9	50	R.T.	88
Ni SAC	KHCO ₃	MEA with BPM	Indirect	CO	67.2	100	R.T.	89
Ni SAC	KHCO ₃	MEA with BPM	Indirect	CO	93	200	R.T.	90
CoPc@CNT	KHCO ₃	MEA with CEM	Indirect	CO	96.2	300	R.T.	91
FeNi alloy (foam)	Na ₂ CO ₃ -K ₂ CO ₃	Undivided cell	Direct	HCOO ⁻	90.5	100	800	92
Bi-coated carbon	KHCO ₃	MEA with BPM	Indirect	HCOO ⁻	64	100	R.T.	93
Bi electrodeposited on carbon	NH ₄ HCO ₃	MEA with AEM	Indirect	HCOO ⁻	75	100	40	94
SnO ₂ /C	Captured CO ₂ from air using KOH solution	MEA with BPM	Indirect	HCOO ⁻	16	50	R.T.	84
Sn (rotating disc electrode)	KHCO ₃	H cell	Indirect	HCOO ⁻	66	~3 ^d	R.T.	95
Sn (foil)	KHCO ₃	H cell	Indirect	HCOO ⁻	18	6	R.T.	96
Sn (foil)	KHCO ₃	Undivided cell	Direct	HCOO ⁻	83	146 ^d	100	97
Pb (foil)	KHCO ₃	Flow cell	Indirect	HCOO ⁻	50	25	75	87
Cu (foam)	CO ₂ -captured AMP	MEA with BPM	Indirect	CH ₄	27	400	R.T.	98
Cu (mesh)	KHCO ₃	MEA with BPM	Indirect	CH ₄	27	500	R.T.	99
Fe	Li ₂ CO ₃ -Na ₂ CO ₃ -K ₂ CO ₃ -LiOH	Undivided cell	Direct	CH ₄	63.3	15	575	100
Fe sheet	Li _{1.427} Na _{0.359} K _{0.214} CO ₃	Undivided cell	Direct	CH ₄	42.7	400	650	101
Cu/Ag (bilayer)	KHCO ₃	MEA with BPM	Indirect	C ₂ H ₄ , C ₂ H ₅ OH, C ₃ H ₇ OH	41.6	100	R.T.	102
Cu/CoPc-carbon nanotube (CNT)	K ₂ CO ₃	MEA with CEM and interposer	Indirect	C ₂ H ₄	34	300	R.T.	103
Cu/Ag (NPs)	K ₂ CO ₃	MEA with BPM	Indirect	C ₂ H ₄	9	150	R.T.	104
Ag-Cu(OH) ₂ nanowire on Cu foam	K ₂ CO ₃ /KHCO ₃	MEA with BPM	Indirect	C ₂ H ₄	12	100	R.T.	105
Cu/Zn (Muntz brass sheet; 3:2 alloy)	Li ₂ CO ₃ -Na ₂ CO ₃	Undivided cell	Direct	carbon nano-scaffold	85	200	670	106
Sn (liquid cathode)	Li ₂ CO ₃ -Na ₂ CO ₃ -K ₂ CO ₃	Undivided cell	Direct	Sn-C spheres	84	N.A. ^e	550	107
SnO ₂	CaCO ₃ -CaCl ₂ -NaCl	Undivided cell	Direct	Sn@CNT	85.1	N.A.	800	108
Steel (galvanized)	CO ₂ -captured Li _{1.5} Na _{1.5} BO ₃	Undivided cell	Direct	CNT	55	120	600	109
Steel (galvanized)	Li ₂ CO ₃	Undivided cell	Direct	Carbon platelets	N.A.	200	730	110
Ni	Li ₂ CO ₃ -K ₂ CO ₃ -Na ₂ CO ₃	Undivided cell	Direct	Graphite	N.A.	5-100	650-750	111
Ni	LiCl-Li ₂ CO ₃ -LiBO ₂	Undivided cell	Direct	Carbon nanofibers	93.8	35	550	112
Ni	Li ₂ CO ₃	Undivided cell	Direct	Carbon nano-onion	N.A.	150	750	113

^a $|j|$: current density. ^b T: temperature. ^c DTAB: dodecyltrimethylammonium bromide. ^d Calculated based on data from references. ^e N.A.: not available.



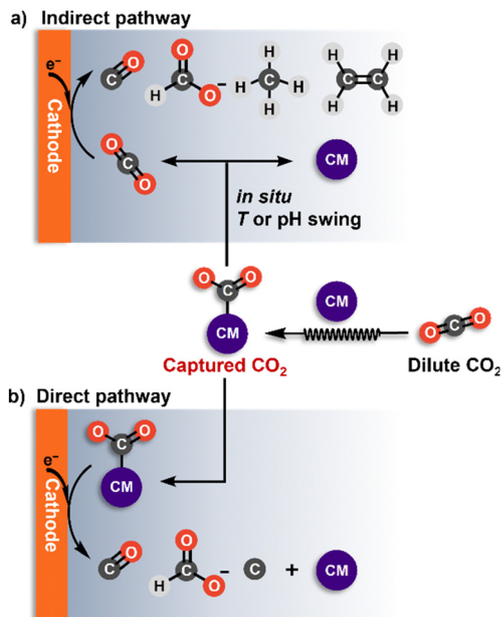


Fig. 4 Illustration of the two pathways of electrochemically reactive CO_2 capture: (a) Indirect with the captured CO_2 converted back to CO_2 at the electrode–electrolyte interface before its electrochemical reduction; (b) direct with the captured CO_2 as a reactant in the electrochemical step. CM: capture medium.

conditions.^{85,97,118} Similarly, in molten salts, the captured CO_2 (as CO_3^{2-}) can be directly converted, using the high temperature to overcome the energy barrier for reduction.

The direct conversion of bicarbonate or carbamate, particularly to C_{2+} products, presents a formidable challenge, with few studies reported. It has been observed that the direct pathway for carbonate conversion does not involve CO intermediates.¹¹⁸ As a result, this pathway cannot generate C_{2+} products, as these are primarily formed through CO–CO coupling. To effectively catalyze the conversion of bicarbonate or carbamate, the catalysts need to have a strong affinity for the reactant. Since heterogeneous electron transfer is an inner-sphere reaction, the proximity between the reactant and the electrode surface is crucial for the electrolysis reaction. Unlike charge-neutral CO_2 molecules, bicarbonates and carbamates are anions that tend to migrate toward the anode, resulting in low concentrations at the cathode interface. Therefore, a positively charged catalyst surface may aid in attracting the anionic reactant through electrostatic interactions. Thus, careful tuning of the electronic configuration and the morphology of electrocatalysts, along with surface modification, is essential to promote efficient reactant adsorption. Additionally, a more comprehensive understanding of the mechanistic pathway for the direct conversion of bicarbonate or carbamate is needed to investigate the potential for C–C coupling in this system.

3.2. Production of CO

In the electrochemical conversion of captured CO_2 into CO, silver nanomaterials are extensively utilized as catalysts, similar to their prominent role in the electrochemical reduction of

CO_2 .^{72,75,119} Li *et al.* employed Ag as the electrocatalysts for the electroreduction of a carbonate electrolyte, achieving a CO faradaic efficiency (FE_{CO}) that decreased from 28% to 12% as the current density increased from 100 to 300 mA cm^{-2} .⁷² In this process, the carbonate reacts with protons from the bipolar membrane (BPM) to generate CO_2 *in situ* near the electrode surface (eqn (21)), following the indirect pathway depicted in Fig. 4a. In another work, Li *et al.* discovered that pure Ag foil was ineffective in catalyzing the direct reduction of NH_4HCO_3 , while a Br-modified Ag electrode exhibited higher efficiency with a significant FE_{CO} of 77.8% and a current density of 13.8 mA cm^{-2} at -0.6 V vs. RHE.^{83,120} They hypothesized that the Br modification could increase the number of active sites on the Ag surface, promoting the release of CO_2 from NH_4HCO_3 and its subsequent reduction.

Mezza *et al.* conducted a systematic study on the effect of Ag mass loading on the performance of bicarbonate electrolysis to CO (eqn (18)) through the indirect pathway. Their findings showed that as the Ag loading rose from 116 to 565 $\mu\text{g cm}^{-2}$, FE_{CO} improved from 55% to 77%, while the partial current density of CO rose from 6 to 13 mA cm^{-2} . Further increases in the Ag loading did not lead to additional improvement because a balance was reached between the availability of active sites and the electrode permeability.¹²¹ It is important to note that there is a trade-off between selectivity and activity of CO production under these conditions. The decrease in FE_{CO} at higher current densities is due to an imbalance between the utilization rate of the *in situ* generated CO_2 and the local CO_2 concentration on the catalyst surfaces.



Lee *et al.* proposed a direct pathway for using carbamate electrolyte as the reactant in electrochemical conversion (Fig. 4b). They achieved an impressive FE_{CO} of 72% at 50 mA cm^{-2} in a flow cell using a Ag electrocatalyst as the cathode in an electrolyte consisting of 30 wt% monoethanolamine with 2 M KCl at 60 °C (eqn (17)).⁸⁵

SACs have emerged as promising candidates for the electrolytic conversion of captured CO_2 , due to their exceptional selectivity for CO, high activity, and maximum atom efficiency. Kim *et al.* demonstrated that a Ni–N/C SAC exhibited more selective production of CO (78.3%) compared to commercially available Ag (cAg) catalysts (38.3%) in CO_2 -captured monoethanolamine (Fig. 5a).⁸⁸ They ascribed this enhanced performance to the much higher potential of zero charge (PZC) of Ni–N/C compared to cAg, as a positively shifted PZC can increase the surface cation charge density at the same applied potential in the reduction reaction. This, in turn, stabilizes the CO_2^- intermediate and enhances the CO_2 reduction activity. This relationship was further confirmed by studying the cation sensitivity using additional catalysts (AuAg alloy, cPd, and cAu catalysts; Fig. 5b and c). Ni–N/C could maintain a FE_{CO} of up to 50% as the cation size increased from K^+ to diethanolammonium (Fig. 5d–i), showing selective CO production in different



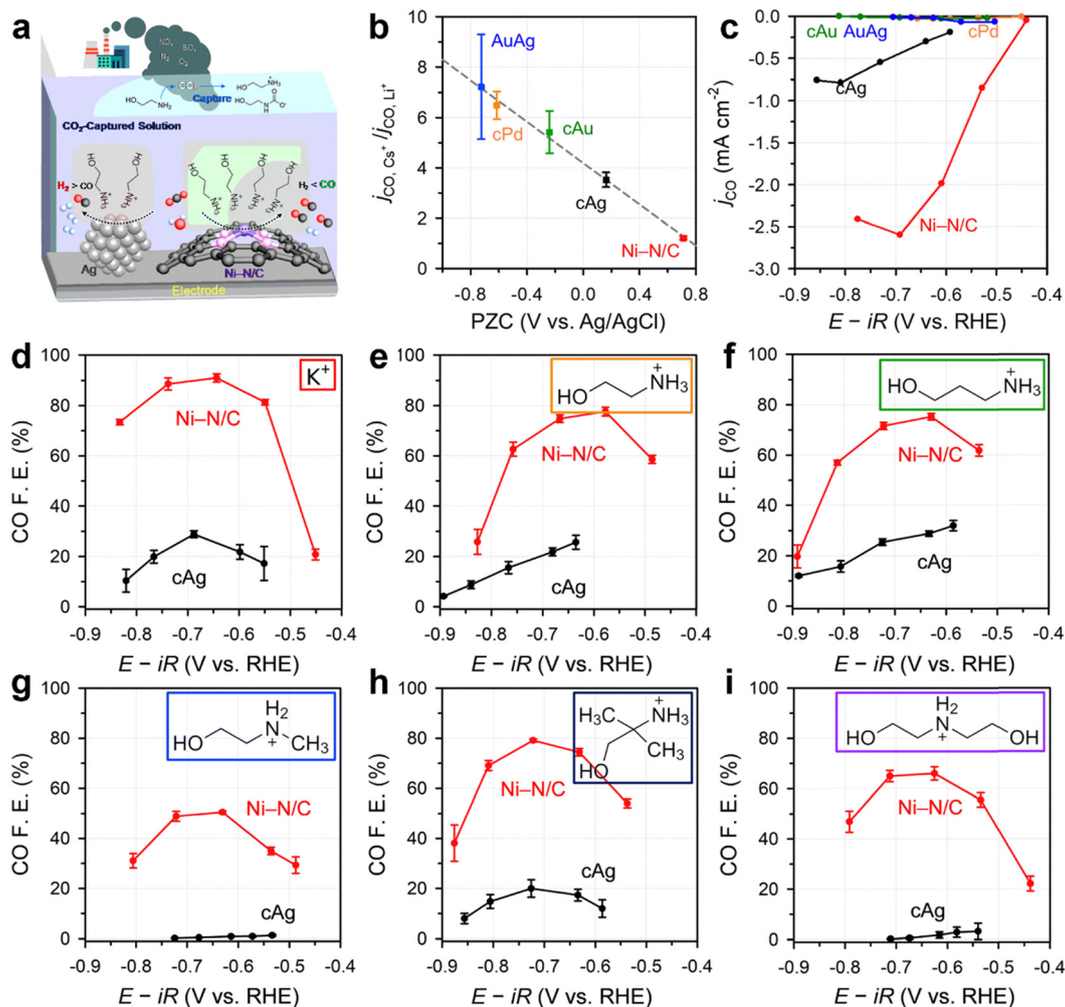


Fig. 5 (a) Schematic illustration of the superior performance of Ni-N/C catalysts compared with Ag in electrochemical conversion of captured CO₂ to CO. (b) and (c) Relationship between cation sensitivity of the catalysts and CO production performance: (b) normalized partial current density, j_{CO} , at -1.7 V vs. NHE with respect to the PZC of the catalysts in 0.05 M carbonate electrolytes; (c) j_{CO} of the catalysts for the electrolysis of CO₂-absorbing 5 M monoethanolamine solution. FE_{CO} of Ni-N/C and cAg for electrolysis in CO₂-absorbing (d) 1 M KHCO₃, (e) 1 M monoethanolamine, (f) 1 M 3-amino-1-propanol, (g) 1 M 2-(methylamino)ethanol, (h) 1 M 2-amino-2-methyl-1-propanol, and (i) 1 M diethanolamine Ar-saturated solutions. Reproduced with permission from ref. 88.

CO₂-captured media. In contrast, the FE_{CO} of cAg dropped to almost zero as the bulkiness of the amine increased.

In addition, Yue *et al.* used a Ni SAC derived from a metal-organic framework (MOF) to boost the electrolytic bicarbonate conversion to CO, achieving a FE_{CO} of 67.2% at 100 mA cm⁻².⁸⁹ The abundant mesopores in the Ni SAC promoted the transport of the *in situ* released CO₂ reactant. Cu electrodes have also demonstrated potential in the electrocatalytic conversion of carbamate into CO. Zhang and Kraatz showed that Cu electrodes significantly enhanced FE_{CO} to 58% compared to glassy carbon electrodes (2.3%) when using ethylenediamine as the capture media at -0.76 V vs. RHE.¹²² This differs substantially from numerous reports that typically show Cu-based materials catalyzing the conversion of gas-fed CO₂ to C₂₊ products.^{123,124} Furthermore, the current density rose notably from 0.63 to 18.4 mA cm⁻². While the morphology and crystalline properties

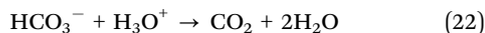
of the Cu electrodes used were not specified, these factors are likely crucial in CO production.

3.3. Production of HCOO⁻

Hori and Suzuki were the first to report the electrocatalytic reduction of bicarbonate into formate at a mercury electrode at relatively low current densities (<1 mA cm⁻²).¹²⁵ They suggested that the electrolytic reduction of HCO₃⁻ is a very slow process due to the slow dissociation of HCO₃⁻ to CO₂ (reverse of eqn (2)). Min and Kanan employed a Pd electrocatalyst that achieved a formate faradaic efficiency (FE_{formate}) of $\sim 54\%$ at 6.0 mA cm⁻² in aqueous bicarbonate solutions without requiring any CO₂ input.¹²⁶ They hypothesized that bicarbonate first decomposed into CO₂, followed by the subsequent reduction of CO₂ to formate (Fig. 4a, indirect pathway). In a flow cell equipped with a BPM, Bi catalysts demonstrated high efficiency



in converting HCO_3^- to formate, achieving a $\text{FE}_{\text{formate}}$ of 64% at 100 mA cm^{-2} in a 3 M KHCO_3 electrolyte.⁹³ In this process, the proton flux facilitated by the BPM converted HCO_3^- to CO_2 (eqn (22)), which was subsequently reduced to formate on the Bi surface. Similarly, electrodeposited Bi on a carbon paper substrate exhibited excellent performance in an ammonium bicarbonate (NH_4HCO_3)-fed electrolyzer, using *in situ* CO_2 as the primary reactant at slightly elevated temperature ($\sim 40 \text{ }^\circ\text{C}$).⁹⁴ Because NH_4HCO_3 ($36 \text{ }^\circ\text{C}$) has a much lower decomposition temperature than ethanolamine- CO_2 ($120 \text{ }^\circ\text{C}$) and KHCO_3 ($150 \text{ }^\circ\text{C}$), it can provide larger amounts of CO_2 to the Bi electrode, resulting in higher current densities at a smaller energy input.



Pei *et al.* demonstrated that a dynamic $\text{Sn}^{\delta+}/\text{Sn}$ interface significantly enhanced the direct reduction of bicarbonate to formate, achieving an optimal partial current density of 121 mA cm^{-2} for formate with a FE of 83% achieved in a 3 M KHCO_3 solution at $100 \text{ }^\circ\text{C}$.⁹⁷ Their study indicated that formate was generated directly from bicarbonate, rather than from CO_2 produced by the dissociation of bicarbonate at high temperatures (Fig. 4b, direct pathway).

Ma *et al.* revealed that carbonate intermediates adsorbed onto a Cu electrode during the electrochemical CO_2 reduction in a KHCO_3 electrolyte over a potential range from -1.0 to 0.2 V vs. RHE (Fig. 6a).¹¹⁸ At potentials of -0.4 V vs. RHE and more negative, these intermediates were reduced to formate. Although the $\text{FE}_{\text{formate}}$ was quite low (0.61%), this finding demonstrated the potential for directly reducing carbonates using a modified Cu electrode. Pulse electrolysis experiments conducted under a N_2 atmosphere with $0.05 \text{ M K}_2\text{CO}_3$ aqueous solution as the electrolyte corroborated the direct electroreduction of carbonate at the Cu surface (Fig. 6b).¹¹⁸ *In situ* Raman spectroscopy at the applied potential of -0.05 V vs. RHE , confirmed the formation of a $^*\text{CO}_3^{2-}$ intermediate (where * indicates it is bound to the surface). Upon switching the potential to -0.50 V vs. RHE , the surface-bound carbonate was reduced to formate, as detected by high-performance liquid chromatography. Isotope labeling confirmed that the carbonate was the carbon source of the produced formate. In contrast,

constant potential electrolysis at -0.50 V vs. RHE under the same reaction conditions yielded no formate.

The carbonate reduction mechanism was modeled using density functional theory (DFT) calculations according to the two pathways shown in Fig. 6. The results indicated for the direct pathway that carbonate adsorption at Cu sites is favorable at -0.05 V vs. RHE (Fig. 6b; first step of red lines). However, at this potential, the protonation of $^*\text{CO}_3^{2-}$ to form $^*\text{HCO}_3^-$ is unfavorable (Fig. 6b; red lines). Upon switching the potential to -0.50 V vs. RHE , the formation of $^*\text{HCO}_3^-$ becomes feasible, along with the subsequent reaction steps leading to formate production (Fig. 6b; black lines). Notably, carbonate with $\Delta G_{\text{ads}} = 0.68 \text{ eV}$ cannot be adsorbed at -0.50 V vs. RHE (Fig. 6b; gray lines) due to increased repulsion between CO_3^{2-} and the negatively charged electrode. This explains the lack of formate production during constant potential electrolysis experiments.

For the indirect pathway, which involves the release of CO_2 before electrochemical reduction (Fig. 6a), the calculations show that a $^*\text{CO}_3^{2-}$ intermediate forms through the reaction of CO_2 with a residual $^*\text{O}$ atom on the Cu surface. As expected, this reaction pathway toward formate production becomes energetically favorable at -0.50 V vs. RHE (Fig. 6a; black lines). Note that this pathway is stoichiometric in reactants and cannot continue once the $^*\text{O}$ atoms are consumed. Overall, the experimental and computational findings demonstrate that carbonate can be directly reduced at electrode surfaces, providing a possible method to activate the otherwise inert CO_2 -adduct.

3.4. Production of CH_4

Methane is a highly desirable product due to its superior heat value of 55.5 MJ kg^{-1} , significantly greater than that of CO (10.2 MJ kg^{-1}) and methanol (22.7 MJ kg^{-1}).¹²⁷ Lees *et al.* reported that a Cu catalyst modified with cetyltrimethylammonium bromide (CTAB) facilitated CH_4 formation at a partial current density of $120 \pm 10 \text{ mA cm}^{-2}$ and with a yield of $34 \pm 7\%$ during bicarbonate electrolysis, where the *in situ* generated CO_2 served as the actual reactant.⁹⁸ A one-dimensional continuum model of the cathode compartment suggested that H^+ from the BPM in the bicarbonate electrochemical reactor would

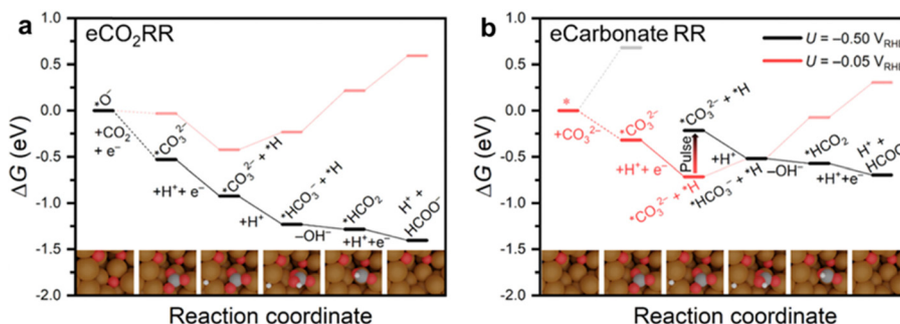


Fig. 6 Energy profiles of (a) CO_2 reduction by CO_2 adsorption on an $^*\text{O}$ site at Cu and further reduction to formate at an applied potential of -0.5 V vs. RHE , and (b) CO_3^{2-} reduction to formate at applied potentials of -0.05 V vs. RHE (red), -0.50 V vs. RHE (black), and with the potential pulsed from -0.05 to -0.50 V vs. RHE . Reproduced with permission from ref. 118.



neutralize hydroxide generated at the cathode surface, favoring CH₄ over multicarbon products.

In the electrochemical conversion of captured CO₂, CH₄ is predominantly generated at elevated temperatures, with molten salts as the capture medium.¹⁰⁰ When molten carbonate salts are used, the type and distribution of products are largely determined by the properties of these salts, as well as the electrolysis conditions, including temperature and applied potential. In this setup, cathode materials are essential as they act as electron-transfer agents, which can greatly influence the overpotential and efficiency of the reactions. For effective electrolysis in molten salts, the cathode material must fulfill two criteria: (1) resistant to corrosion by the highly alkaline molten salts during high-temperature electrolysis. (2) Unable to alloy with Li, Na, or K at high temperatures in molten salts, as this would compromise the electrode structure.¹⁰⁰

Wu *et al.* reported on the electrolysis of CO₂ and H₂O employing Fe and Ni electrodes as the cathode and anode, respectively, in an alkali carbonate/LiOH electrolyte at 600 °C.¹²⁸ The electrolysis products comprised 64.9% CH₄, 34.8% H₂, and 0.3% C₂₊. Bai *et al.* systematically compared the overpotential and CH₄ yield of different cathode materials, including Fe, stainless steel (SS304), Ti, and cupronickel sheets, in molten Li_{1.427}Na_{0.359}K_{0.214}CO₃ ~ 0.15LiOH at 650 °C.¹⁰¹ Their study found that Fe, cupronickel, and SS304 exhibited similar cathodic overpotentials, all higher than that of Ti. Notably, the highest CH₄ yield (33.3%) was achieved with the cupronickel cathode, exceeding the yield from the standard Fe cathode (25.4%) despite its higher activity. This enhanced CH₄ production from the cupronickel cathode is likely due to the synergistic electronic effects of Cu and Ni atoms, which improve the adsorption of reaction intermediates for CH₄ production during electrolysis.

3.5. Production of C₂₊

The electrochemical conversion of captured CO₂ into C₂₊ products offers greater potential in terms of market size and value compared to C₁ products. However, this area of research has not been as extensively explored. Lee *et al.* applied Cu/Ag bilayer electrodes with engineered bilayer cation- and anion-conducting ionomers to convert bicarbonate into C₂₊ products such as acetate, ethylene, ethanol, and propanol.¹⁰² The *in situ* generated CO₂ was first converted to CO on the Ag layer (Fig. 4a, indirect pathway), which was then efficiently transformed into C₂₊ products on the Cu layer, due to the locally concentrated CO. They achieved a maximum FE of 41.6 ± 0.4% for C₂₊ products at a current density of 100 mA cm⁻². Prajapati *et al.* developed a fully integrated system for CO₂ capture and reduction using Cu mesh catalysts, achieving a steady-state CO₂ reduction FE of 57% and a high FE of 40% for C₂H₄ at a current density of 200 mA cm⁻² (see Section 5.2).¹²⁹

Recently, Song *et al.* demonstrated the direct conversion of carbonate solutions to C₂H₄ using a customized Cu–Ag catalyst in a BPM-MEA (membrane electrode assembly) setup. Although the carbonate electrolysis resulted in a relatively low FE of ~10% for C₂H₄ production, the Cu–Ag electrode configuration

effectively achieved nearly zero CO₂ concentration in the outlet stream. Increasing the operating temperature to 50 °C further enhanced C₂H₄ production, with a higher partial current density of 18–20 mA cm⁻² compared with 10 mA cm⁻² at room temperature.¹⁰⁴

3.6. Production of carbon materials

In addition to the common products from CO₂ reduction, valuable carbon materials, such as graphene and nanotubes, can be obtained in the electrochemical conversion of molten carbonate salt. The cathode's function as a substrate for carbon deposition affects the nucleation and growth of these carbon materials.¹³⁰ For example, a Zn galvanized steel cathode surface was shown to yield a high proportion of carbon nanofibers (>80%), a result not achievable with a 316 stainless steel cathode.¹³¹

Hu *et al.* successfully produced graphene using stainless steel as the cathode material in molten salt electrolysis.¹³² They proposed that active Fe atoms on the cathode surface catalyze carbon growth, leading to the formation of Fe₃C as a key intermediate product. Subsequently, active Fe atoms are inserted into the interlayer of the graphene sheets, enabling continuous layer-by-layer growth through micro-explosion reactions in the interlayer space caused by the intense generation of CO. Moreover, different cathode materials were found to influence the structure of the carbon products, with Cu rods resulting in ball-structured graphene and Ni cathodes catalyzing the formation of flat graphene sheets.

Pint *et al.* demonstrated the electrochemical synthesis of carbon nanotubes from the capture and conversion of ambient CO₂ using Fe as the cathode and further revealed a correlation between the thickness of the Fe metal layer and the diameter of the carbon nanotubes.¹³³ This highlights the importance of active metal atoms on the cathode as nucleation sites that catalyze the growth of carbon films.

3.7. Designing catalysts with CO₂ capture capability

Recently, a growing body of research has focused on using catalysts capable of capturing CO₂. In these studies, the capture and conversion of CO₂ are integrated within the catalyst itself, eliminating the need to rely on the electrolyte. For example, Liu *et al.* demonstrated this by depositing amine-linked covalent organic frameworks (COFs) onto a flat Ag electrode.¹³⁴ They created a molecularly defined interface for the electrochemical reduction of CO₂ to CO, achieving significantly higher efficiency and selectivity than a bare Ag electrode. This improved performance was attributed to the synergistic interaction between the COFs and the Ag electrode interface. The porous nature of the COFs facilitated CO₂ diffusion to the electrode surface, while the amine functional groups near the electrode surface boosted the CO₂ conversion efficiency by promoting the formation of a carbamate intermediate. Similar effects were observed in another study, where Liu *et al.* coated Ag nanoparticles with a thin 2 nm layer of 4-aminobutylphosphonic acid confined in poly(vinylpyrrolidone).¹³⁵



Yan *et al.* used redox-active 2-amino-5-mercapto-1,3,4-thiadiazole (AMT) functionalized gold nanoparticles to achieve electrochemical carbon capture and conversion from flue gas.¹³⁶ The AMT ligand not only acts as a selective capture agent to concentrate CO₂ near the active site but also serves as a protective layer to inhibit O₂ reduction. As a result, a maximum FE_{CO} of 80.2% at -0.45 V vs. RHE in an H-type cell and 66.0% at a voltage of 2.7 V in a full cell was achieved in simulated flue gas (15% CO₂, 4% O₂, balanced with N₂). Notably, a microporous conductive Bi-based MOF (BiHHTP, HHTP = 2,3,6,7,10,11-hexahydroxytriphenylene) was recently reported to efficiently capture CO₂ from dilute sources under high humidity and catalyze the electroreduction of the adsorbed CO₂ into formic acid with a high current density of 80 mA cm⁻² and a FE of 90% at a low cell voltage of 2.6 V.¹³⁷ This high performance was attributed to the CO₂ capture capability of the micropores on BiHHTP and the lower Gibbs free energy of formation for the key intermediate *OCHO on the exposed Bi sites.

Generally, amine-based molecules^{135,138,139} and porous framework materials^{140–142} are employed as agents for capturing CO₂. The immobilization of amine groups onto the catalysts will enable the catalysts to concentrate CO₂ molecules near the active sites, thereby enhancing the electrochemical conversion of CO₂. Furthermore, functional groups such as amines help stabilize key intermediates during CO₂ reduction, improving selectivity for specific products. Using catalysts with CO₂ capture capabilities is particularly valuable when dealing with flue gas, as it significantly suppresses the competing reduction of O₂.

4. Reactor and system optimization

Besides the capture medium and the catalyst, several other factors influence the outcome of the electrochemical conversion of captured CO₂, including the electrolyzer design, electrode configuration, ion exchange membranes, electrolyte with additives, and reaction conditions. This section will explore these aspects in detail.

4.1. Batch cell versus flow reactor

The electrochemical conversion of captured CO₂ can be conducted in a batch or flow process.¹⁴³ The conversion of molten carbonate has typically been carried out in an undivided cell equipped with heating elements under batch conditions, as high temperature is required for this process (Fig. 7a). At elevated temperatures, the electrolysis of molten carbonate can typically achieve a current density of 100–200 mA cm⁻² and a FE of 60–90% for carbonaceous products.^{108,144} However, for the conversion of captured CO₂ in aqueous solution (such as bicarbonate and carbamate) at room temperature, both batch (*e.g.*, H-cell) and flow (*e.g.*, MEA) reactors have been used. Nonetheless, the reaction process and outcomes vary significantly depending on the type of reactor employed.

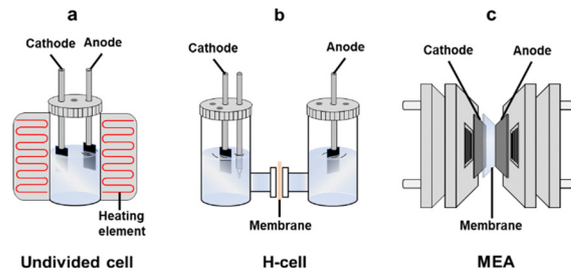


Fig. 7 Three types of reactors used for the electrochemical conversion of captured CO₂: (a) an undivided cell for molten carbonate electrolysis at high temperatures, (b) an H-cell, and (c) an MEA for electrolysis of aqueous bicarbonate and carbamate solutions at room temperature.

In an H-cell, the potential can be easily adjusted to control the product distribution, but this setup typically achieves low current density and FE (Fig. 7b). For instance, Hu *et al.* carried out electrolysis of a KHCO₃ solution in an H-type cell using a hybrid catalyst composed of cobalt porphyrin and carbon nanotubes, achieving a current density of a few mA cm⁻² and FE_{CO} of 18%.¹⁴⁵ This performance is much lower than that of gas-fed CO₂ conversion using the same H-cell with the same catalyst and is also significantly poorer than that achieved in an MEA setup. Similarly, Chen *et al.* conducted the electrochemical reduction of carbamate formed through CO₂ capture by 30% monoethanolamine solution in an H-cell using various smooth or porous metal electrodes, such as In, Sn, Bi, Pb, Pd, Ag, Cu, and Zn.¹⁴⁶ The highest FE_{CO} and FE_{HCOO-} reported were 39% and 45%, respectively, with current densities below 50 mA cm⁻². These studies indicate that the electrochemical conversion performance of aqueous bicarbonate and carbamate solutions in an H-cell reactor is relatively low. This is due to the very low concentration of *in situ* liberated CO₂, produced only from the dissociation of RNHCOO⁻ or HCO₃⁻ (reverse of eqn (1) and (2)), and the mass transport limitation of the liberated CO₂ in the H-cell.¹²⁵ The liberated CO₂ must diffuse through the reaction layer to be reduced at the electrode surface.¹⁴⁶

The current density and FE can be significantly enhanced when converting captured CO₂ in flow reactors, such as MEA electrolyzers (Fig. 7c). This enhancement is due to the specific configuration within the MEA electrolyzers, which includes the CO₂-captured electrolyte, catalyst layer, and membrane. Besides the dissociation of RNHCOO⁻ or HCO₃⁻, a much higher concentration of CO₂ can be generated near the catalyst layer due to the reaction of RNHCOO⁻ or HCO₃⁻ with H⁺ produced from the heterolytic dissociation of water at the BMP. As a result, most studies on captured CO₂ conversion have been conducted using MEA electrolyzers, achieving significantly higher FE up to 90% and larger current density up to 200 mA cm⁻² in some cases (Table 1).^{90,91,98} These performance metrics are comparable to those achieved with gas-fed CO₂ conversion and are suitable for meeting the requirements of industrial applications.^{147,148}

In summary, conducting the conversion of aqueous bicarbonate/carbamate solutions in an H-cell offers high flexibility and quick startup, but it often results in low overall efficiency.



In contrast, MEA electrolyzers operate with higher efficiency, leading to increased production rates, which makes them more advantageous for large-scale applications. The next section will further explore how the key components of MEA electrolyzers impact the performance of the electrochemical conversion of bicarbonate/carbamate solutions.

4.2. Electrode configuration

In the electrochemical conversion of gas-fed CO_2 , gas diffusion electrodes (GDEs) are required to achieve a large current density.¹⁴⁹ Increasing the hydrophobicity of GDEs can enhance performance, suppress the flooding of the electrodes, and thus maintain stability for CO_2 electrolysis over a longer period.^{150,151} However, the electrodes for the electrolysis of captured CO_2 differ significantly from those typically employed for gaseous CO_2 electrolysis because the source of CO_2 is different. In general, a certain level of hydrophilicity is needed for these electrodes to ensure adequate contact between the catalyst on the electrode and the CO_2 -captured solution, whether to facilitate the direct conversion of the CO_2 -captured solution or reduction of the *in situ* generated CO_2 .

Lees *et al.* systematically investigated the impact of electrode structure on the electrolysis of bicarbonate solutions using a Ag catalyst.⁷⁶ They found that adding hydrophobic polytetrafluoroethylene (PTFE), a microporous layer (MPL), or both, negatively affected bicarbonate electrolysis, significantly reducing FE_{CO} . The hydrophobic PTFE and MPL hinder the transport of solvated HCO_3^- ions through the GDE, thereby decreasing the rate of *in situ* CO_2 generation at the electrode *via* the reaction of HCO_3^- with H^+ from the bipolar membrane (eqn (22)). They also examined the effect of Nafion loading in the catalyst layer and observed that FE_{CO} initially increased to a plateau of $\sim 58\%$ and then decreased as the Nafion loading increased at a current density of 100 mA cm^{-2} . The optimal Nafion loading was found to be 4.0 wt%. This behavior is likely due to variations in the catalyst layer morphology; lower Nafion loading leads to poor adhesion of the Ag catalyst to the electrode, while higher loading leads to a densified catalyst layer that blocks the pores of the gas diffusion layer. Similar behavior has been observed in the electrochemical conversion of gas-fed CO_2 .¹⁵² By removing the hydrophobic components of the GDE and optimizing the catalyst coverage, optimal performance for bicarbonate electrolysis was achieved, with a FE_{CO} of 82% at 100 mA cm^{-2} .

Based on this understanding, the same group developed a free-standing porous Ag electrode, which demonstrated higher hydrophilicity than the commonly used GDE and could be more easily integrated into commercial electrolyzers.⁷⁹ As a result, the free-standing Ag electrode achieved a respectable FE_{CO} of 59% at 100 mA cm^{-2} under ambient pressure for converting 3 M KHCO_3 .

Later, Lee *et al.* successfully modeled the generation, diffusion, and consumption of chemical species, identifying an optimal electrode architecture for the electrochemical conversion of carbonate solution to C_{2+} products over a Cu catalyst.¹⁰³ Their modeling results showed that the spacing distance

between the cation exchange layer (CEL) and the catalyst layer (CL) influenced the concentrations of CO_3^{2-} , H^+ , *in situ* generated $\text{CO}_2(\text{g})$, and $\text{CO}_2(\text{aq.})$ in the carbonate solution (Fig. 8). When the CEL and the CL were closely spaced, the local pH at the CEL decreased to no more than 10, as the diffusion of CO_3^{2-} and OH^- could easily neutralize the H^+ coming from the CEL. This prevented the *in situ* generation of $\text{CO}_2(\text{g})$ or products during electrolysis. However, when the spacing distance between the CEL and the CL was in the range of 100–300 μm , optimal conditions for converting carbonate to C_{2+} products could be achieved, with a low pH (< 4) at the CEL, promoting the *in situ* generation of a high concentration of $\text{CO}_2(\text{g})$ ($> 4 \text{ vol}\%$) at the CL. Despite this, the pH remained > 13 at the CL, as OH^- was continuously produced by the conversion of the *in situ* generated $\text{CO}_2(\text{g})$ from carbonate. The high local pH favored the C–C coupling to form C_{2+} products over C_1 products. Finally, increasing the spacing distance to $> 540 \mu\text{m}$ decreased the concentration of $\text{CO}_2(\text{g})$ and hindered the carbonate conversion due to the increased likelihood of CO_2 being recaptured over the extended layer distance.

Based on this modeling, the authors used a porous hydrophilic mixed cellulose ester as an interposer to create a well-defined spacing between the CEL and the CL. The electrolysis of carbonate solutions at 250 mA cm^{-2} over a Cu catalyst using an interposer thickness of 130–270 μm increased $\text{FE}_{\text{C}_{2+}}$ to 40%. This was three times larger than with an interposer thickness of 60 μm ($\text{FE}_{\text{C}_{2+}} = 14\%$) and nearly twice as high as observed in the 540 μm case ($\text{FE}_{\text{C}_{2+}} = 25\%$). These results highlight the critical role of electrode configuration design in enhancing the selective electrochemical conversion of carbonate solution to C_{2+} products.

4.3. Ion exchange membrane

Ion exchange membranes serve as separators and regulate ion transport between the cathodic and anodic chambers of the electrolyzer. The choice of ion exchange membrane has a significant impact on the electrolysis process (Fig. 9). BPMs are widely used in the electrochemical conversion of bicarbonate solutions because HCO_3^- can react with the H^+ produced by the BPM to generate *in situ* CO_2 at the BPM–electrode interface (eqn (22)). The *in situ* generated CO_2 subsequently receives electrons from the electrode and undergoes reduction to produce various value-added products such as CO, formate, and C_2H_4 . Using BPMs for bicarbonate conversion offers several benefits, including rapid water dissociation, minimized crossover of generated products, and extended operational lifespan.¹⁵³ However, employing BPMs requires an additional potential of 0.83 V for H^+ generation *via* water dissociation under standard conditions, resulting in a large overpotential, increased energy consumption, and reduced energy efficiency.

Replacing the BPM with an anion exchange membrane (AEM) or a cation exchange membrane (CEM) can lower the cell voltage but may negatively impact the conversion efficiency. Li *et al.* found that an electrolysis cell with a BPM showed a higher $\text{FE}_{\text{formate}}$ (+35%) for KHCO_3 conversion than a corresponding AEM cell, which was attributed to the higher local CO_2



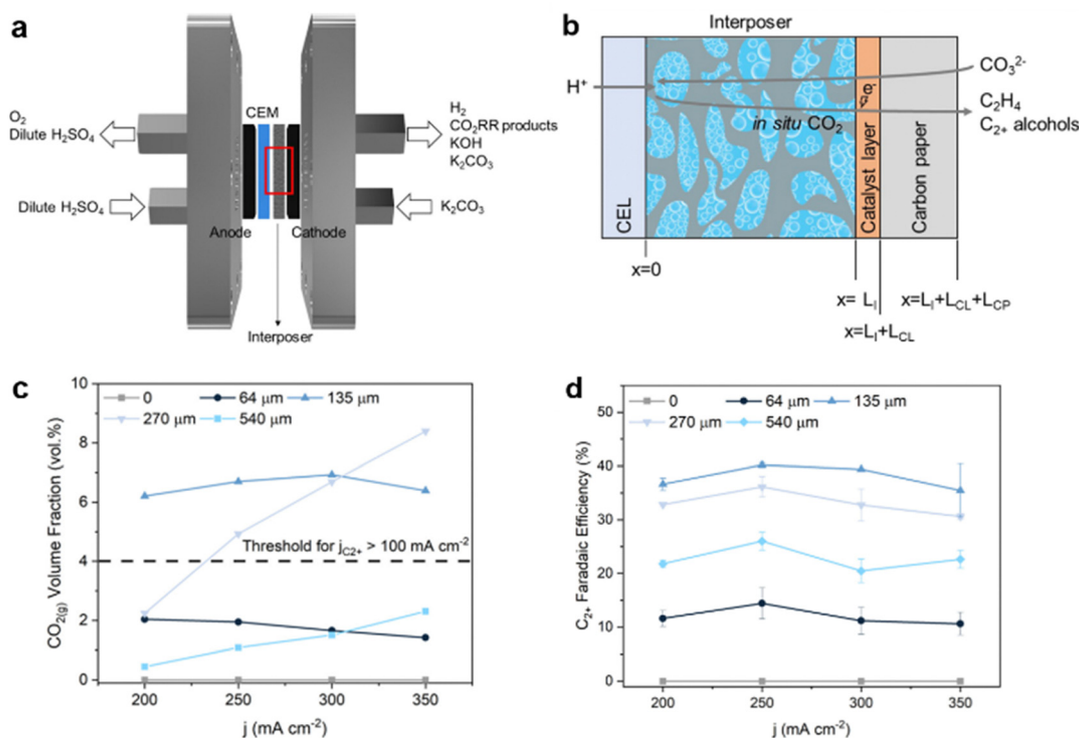


Fig. 8 (a) MEA electrolyzer for aqueous carbonate conversion, where the cathode and anode are separated by an interposer. (b) Schematic of the cathode configuration, including the cation-exchange layer (CEL), the interposer, catalyst layer, and carbon paper. (c) CO₂(g) volume fraction for different spacing (*L_I*) conditions, *i.e.*, 0, 64, 135, and 540 μm at current densities of 200, 250, 300, and 350 mA cm⁻² in 1.5 M K₂CO₃ electrolyte. (d) FE of C₂+ products in carbonate electrolysis using a Cu electrocatalyst in 1.5 M K₂CO₃ electrolyte with various interposer thicknesses from 0–540 μm. Reproduced with permission from ref. 103.

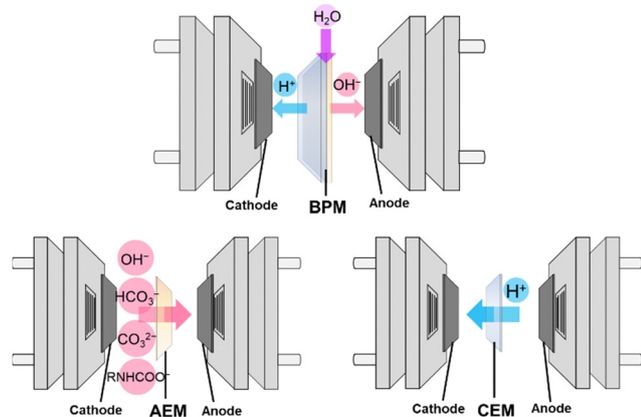


Fig. 9 Schematic illustration of three typical types of ion exchange membranes used in electrochemical conversion of captured CO₂.

concentration at the electrode in the BPM system.⁹³ In the AEM cell, CO₂ was formed only through the bicarbonate dissociation equilibrium (reverse of eqn (2)). However, the trend reversed when using a different bicarbonate (NH₄HCO₃) solution, where replacing the BPM with the AEM resulted in an increased FE_{formate}.⁹⁴ This was explained by the favorable microenvironment created by NH₄⁺, which would suppress the HER at the electrode-membrane interface. It is important to note that

formate crossover could be observed in the AEM cell, as AEMs are designed for anion exchange.¹⁵⁴

Utilizing a CEM can also create an acidic environment at the cathode-membrane interface, as protons are generated from water oxidation (*i.e.*, the oxygen evolution reaction (OER)) or the acidic electrolyte in the anodic chamber permeating through the membrane.¹⁰³ These protons can react with the captured CO₂ (in the form of bicarbonate or carbamate) to produce high local concentrations of free CO₂ near the catalyst layer, thereby enhancing the electrolysis efficiency. To ensure effective proton transport through the CEM, it is crucial to maintain a neutral or acidic electrolyte in the anodic chamber. However, this condition makes the OER more challenging, requiring a larger overpotential.¹⁵⁵ To address this issue, Zhang *et al.* conducted the hydrogen oxidation reaction at the anode instead of the conventional OER during electrolysis of 3 M KHCO₃ and used a CEM in place of the BPM. This approach allowed them to produce CO at a high partial current density of 220 mA cm⁻² at a cell voltage of only 2.3 V.⁸⁰

4.4. Electrolyte with additives

As highlighted in the preceding section, the distance between the reactant and the electrode has a crucial impact on the reaction outcomes. Consequently, using a CO₂-captured amine solution as the electrolyte presents efficiency challenges, unlike



converting the bicarbonate solution. This disparity arises because the bulky carbamate (RNHCOO^-) and ammonium (RNH_3^+) ions impede mass transport at the electrochemical double layer (EDL).

To address this issue, Khurram *et al.* investigated the introduction of various electrolyte salts, including LiPF_6 , LiClO_4 , NaClO_4 , KClO_4 , and TBAClO_4 , into CO_2 -captured amine solutions in dimethyl sulfoxide.¹⁵⁶ They observed that while the anions had a minor impact, the cations significantly influenced the speciation of amine- CO_2 adducts. Specifically, pairing the K^+ cation with RNHCOO^- could enhance the rates of electrochemical reactions due to improved K^+ transfer from the bulk solution to the reaction site. Therefore, the judicious selection of strong electrolyte salt that facilitates rapid cation transfer is essential for achieving high efficiency in converting amine-captured CO_2 .

Later, Lee *et al.* demonstrated that introducing alkali metal ions into the aqueous CO_2 -captured amine solution could significantly improve the electrochemical conversion of the amine- CO_2 adduct (carbamate).⁸⁵ They proposed that the alkali cations (*e.g.*, K^+) modified the EDL to facilitate the heterogeneous electron transfer from the catalyst to the carbamate (Fig. 10a). Using this approach, efficient conversion of the carbamate to CO was achieved with a Ag catalyst, resulting in a $\text{FE}_{\text{CO}} = 72\%$ at 50 mA cm^{-2} .

Surfactants exert a large influence on the efficiency of gaseous CO_2 conversion,^{157,158} as well as the electrochemical conversion of captured CO_2 . Chen *et al.* reported that adding the

cationic surfactant, CTAB, suppressed the competing HER, thereby greatly enhancing the conversion of carbamate derived from the CO_2 capture by monoethanolamine (Fig. 10b).¹⁴⁶ This enhancement was evident in the increase of $\text{FE}_{\text{formate}}$ from 2.4% to 45.4% over an In electrode. In contrast, other surfactants, such as sodium dodecylsulfate (anionic) and TritonX-100 (non-ionic), showed a negligible effect, indicating that the performance of the carbamate conversion depends on the type of surfactant added. Ahmad *et al.* found that adding CTAB to the CO_2 -captured 2-amino-2-methyl-1-propanol solution increased the FE_{CO} over a Cl-modified polycrystalline Ag electrode due to a more compact EDL.⁸⁶ Similarly, for the conversion of a bicarbonate solution, Lees *et al.* observed that the presence of just 3 mM CTAB increased the CO_2 reduction current density from 11 to 162 mA cm^{-2} and the FE_{CH_4} from 0 to 27% at an applied current density of 400 mA cm^{-2} .⁹⁸ This performance improvement was attributed to reduced oxide coverage on the Cu electrode during electrolysis when CTAB was added.

4.5. Operation parameters

The concentration of the captured CO_2 significantly impacts the electrochemical performance of the system. For bicarbonate conversion, higher concentrations lead to higher FE of carbonaceous products.⁹³ Increasing the bicarbonate concentration accelerates the rate at which HCO_3^- dissociates and reacts with H^+ from the membrane to liberate CO_2 . Thus, a high

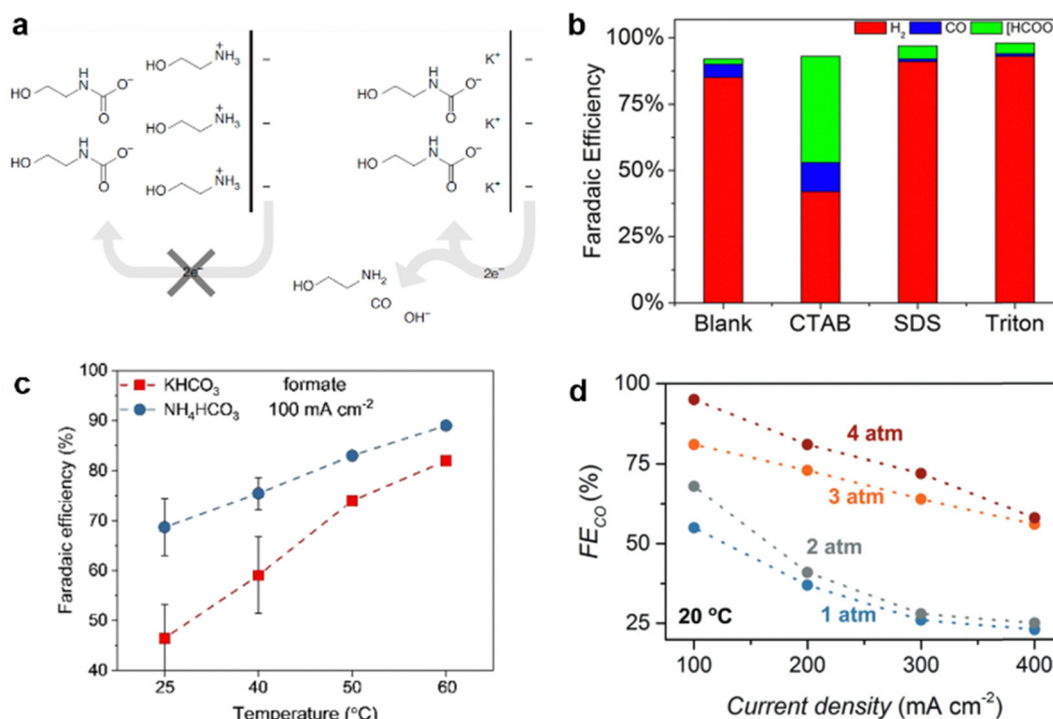


Fig. 10 Effect of (a) strong electrolyte KCl (reproduced with permission from ref. 85), (b) surfactant (reproduced with permission from ref. 146), (c) temperature (reproduced with permission from ref. 94), and (d) pressure (reproduced with permission from ref. 79) on the conversion of aqueous carbamate or bicarbonate solution.



concentration (3 M, near saturation) is typically used for bicarbonate conversion.

Temperature can influence the performance of bicarbonate conversion in several ways. First, increasing the temperature of the electrolyte is expected to promote CO_2 generation by shifting the bicarbonate equilibrium toward CO_2 (reverse of eqn (2)). Moreover, a temperature rise causes an increase in pH due to the formation of OH^- , which suppresses the HER. Raising the temperature also enhances mass transport, thereby accelerating the electrochemical process. This effect has been observed in numerous studies, resulting in improved FE for forming carbonaceous products (Fig. 10c).^{79,94} Similarly, the effect of the temperature was exploited in the conversion of CO_2 -captured amine (carbamate) solutions. Pérez-Gallent *et al.* reported that the reaction rate of the conversion of CO_2 -captured 2-amino-2-methyl-1-propanol in a propylene carbonate solution was significantly enhanced when the temperature was raised from 15 to 75 °C.⁸⁷ The increase in reaction rate was attributed to the much faster liberation of CO_2 at elevated temperatures, which was eight times higher at 75 °C than at 15 °C.

The effect of temperature becomes more complex in the electrochemical conversion of molten carbonate salts. Deng *et al.* reported that the electrolysis of CaCO_3 -containing molten LiCl-KCl over a Ni electrode produced carbon materials with various shapes and morphologies.¹⁵⁹ At 450 °C, micron-sized hollow carbon spheres and ultrathin carbon sheets were the major products, depending on the applied cell voltage. As the temperature increased to 550 and 650 °C, a range of other carbon materials, such as quasi-spherical carbon particles, coral-like carbon, and carbon nanofibers, were formed. In a subsequent study, it was demonstrated that raising the operational temperature led to an increase in current density or a reduction in cell voltage for the conversion of CaCO_3 in the LiCl-KCl melt.¹⁶⁰ These conditions facilitated the dissolution of both the reactant CaCO_3 and the product CaO in the melt, thereby preventing the accumulation of solid CaO on the electrode. As a result, the process exhibited greater durability and energy efficiency. The energy consumption for producing 1 kg of carbon was calculated to be 16.3 kW h kg^{-1} at 650 °C, compared with 31.4 kW h kg^{-1} at 450 °C.

Zhang *et al.* demonstrated that pressure, in addition to temperature, affects the performance of bicarbonate conversion (Fig. 10d).⁷⁹ When the inlet pressure of the electrolyzer was increased from 1 to 4 atm, FE_{CO} increased from 55% to 95%, and the cathodic energy efficiency from 22% to 34% at a current density of 100 mA cm^{-2} . At 4 atm pressure, an impressive current density of 400 mA cm^{-2} for bicarbonate conversion with $\text{FE}_{\text{CO}} > 55\%$ could be achieved. This performance enhancement was attributed to the kinetically improved supply of CO_2 to the catalyst at elevated pressures.

The authors also explored how the electrolyte flow rate influenced the performance of bicarbonate conversion. They increased the flow rate from 30 to 100 mL min^{-1} , which led to an increase in *in situ* generated CO_2 and, consequently, a higher FE_{CO} . This effect was attributed to the enhanced convective mass transport of HCO_3^- . Building on this finding, they

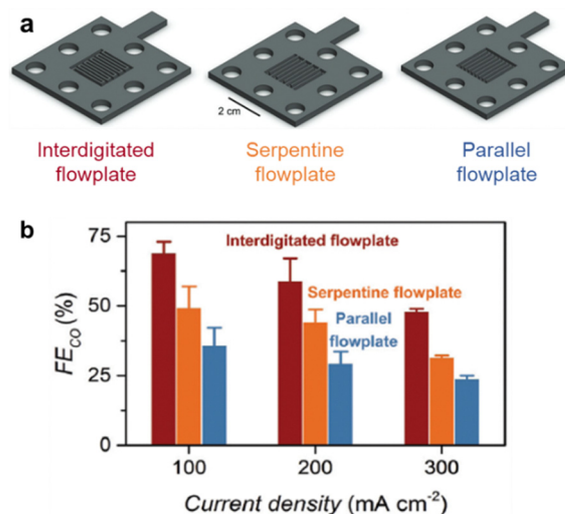


Fig. 11 (a) Interdigitated, serpentine, and parallel cathodic flow plate designs used in bicarbonate electrolysis. (b) FE_{CO} as a function of current density for interdigitated, serpentine, and parallel flow plates tested in a bicarbonate electrolyzer equipped with porous silver electrodes. Reproduced with permission from ref. 79.

modified the flow plate geometry to further enhance convective mass transport. Three types of flow plates were tested, *i.e.* interdigitated, serpentine, and parallel (Fig. 11). Convection was the primary mode of mass transport in the interdigitated flow plate, while diffusion was dominant in the parallel and serpentine flow plates. As a result, the electrolyzer with the interdigitated flow pattern exhibited higher FE_{CO} ($\approx 69\%$ at 100 mA cm^{-2}) compared to the other two. These results demonstrate that enhancing the convective mass transport can increase FE_{CO} significantly for converting captured CO_2 .

5. Practical integration of CO_2 capture and conversion

In this section, we present practical examples that fully integrate the two processes of CO_2 capture and electrochemical conversion (Fig. 1, Route 3). We begin by examining technologies based on different CO_2 sources, including air,^{161,162} flue gas,^{163,164} and ocean water,^{165,166} as these sources significantly influence the outcome of the integration due to differences in CO_2 concentration and purity.

5.1. Conversion of captured CO_2 from various sources

5.1.1. Air. CO_2 in the atmosphere can be captured and utilized through DAC technologies. In the earth's atmosphere, CO_2 constitutes ~ 400 ppm. Although this concentration has devastating climate impacts, it is relatively low in absolute terms, imposing an entropy penalty on DAC technologies. The minimum energy required to separate a stream of air with a CO_2 concentration of 400 ppm into one stream with 200 ppm CO_2 and another with 99% CO_2 , all at the same temperature and pressure, is about 20 kJ mol^{-1} .³⁵ However, the efficiency of real-world DAC technologies is likely to be limited to



~5%.¹⁶⁷ Therefore, the energy requirement to achieve a 99% concentration of captured CO₂ that can be effectively converted electrochemically is ~400 kJ mol-CO₂⁻¹.

If conventional energy sources such as coal were used to power a DAC system with 5% efficiency, there would be no net CO₂ capture, as producing the required 400 kJ to capture one mol of CO₂ would generate 2.5 moles of CO₂. This is also true for natural gas, the least carbon-intensive fossil fuel, which emits one mol of CO₂ while generating 396 kJ of energy. Therefore, CO₂-free power sources such as nuclear, wind, solar, and hydro must drive air capture systems. For example, the energy requirement of 400 kJ mol-CO₂⁻¹ is equivalent to ~2500 kW h t-CO₂⁻¹, which could be provided by a large 3 MW wind turbine operating for 1 h.

Integrating the CO₂ capture and electrochemical conversion processes is challenging because the rate of capture is usually much lower than the achievable rate of conversion. A study exemplifying this issue involved absorbing atmospheric CO₂ over 8.5 h using an alkaline solution (1 M KOH), resulting in the formation of a bicarbonate/carbonate mixture (Fig. 12).⁸⁴ This solution was then used as the catholyte in a zero-gap flow electrolyzer to produce industrially significant carbon products. At a current density of 50 mA cm⁻², a FE_{formate} of 16% (using SnO₂/C as the electrocatalyst) and FE_{CO} of 13% (using Ag/C) were achieved. To improve electrochemical performance, it would be necessary to capture higher concentrations of CO₂ by increasing the base concentration and/or extending the absorption period. From an economic perspective, this

would reduce the attractiveness of integrating the two technologies.

Recently, Almajed *et al.* evaluated the feasibility of directly integrating DAC with (bi)carbonate electrolysis using KOH and K₂CO₃ solutions as the capture media (Fig. 13a).¹⁶⁸ They found that the presence of CO₃²⁻ alongside HCO₃⁻ in the effluent from the air contactor significantly reduced electrolysis performance due to incomplete CO₂ capture, ultimately decreasing the CO₂ capture fraction from 78% to ≤1%. As a result, they estimated that air contactors would need to be 5–14 times larger than those typically required for DAC to produce suitable effluents for (bi)carbonate electrolysis, leading to unfavorable process economics. Furthermore, they demonstrated that regenerating the capture media after the electrolysis was insufficient for effective CO₂ recapture. This incompatibility between DAC and (bi)carbonate electrolysis was attributed to the mismatched pH values between the air contactor outlet/electrolyzer inlet and the air contactor inlet/electrolyzer outlet. To resolve this issue, the authors suggested acidifying the air contactor effluent and basifying the electrolyzer effluent by incorporating an electro dialysis unit or directly feeding acidic/basic streams to the respective effluents (Fig. 13b). However, this approach would inevitably increase both capital and operational costs.

The third example concerns molten metal oxides, such as Li₂O, which have shown potential for directly sequestering CO₂ from the air at elevated temperatures, resulting in the formation of carbonate salts.¹³¹ These carbonates can then undergo electrochemical reduction, producing solid carbon

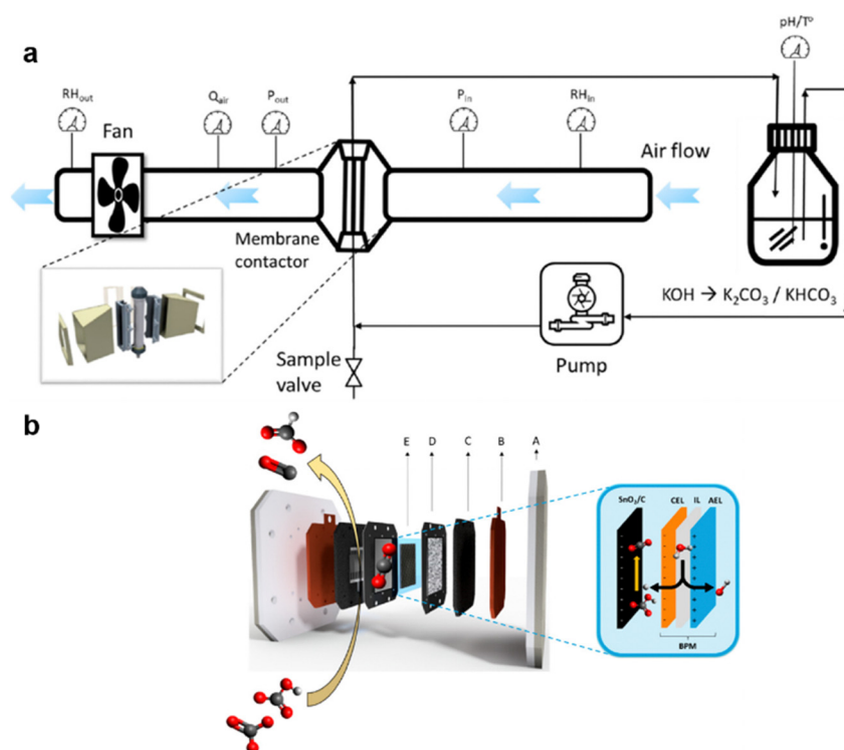


Fig. 12 Integration of atmospheric CO₂ capture and the subsequent electrochemical conversion of the captured CO₂: (a) diagram of the CO₂ capture setup and (b) the electrolytic cell for converting captured CO₂. Reproduced with permission from ref. 84.



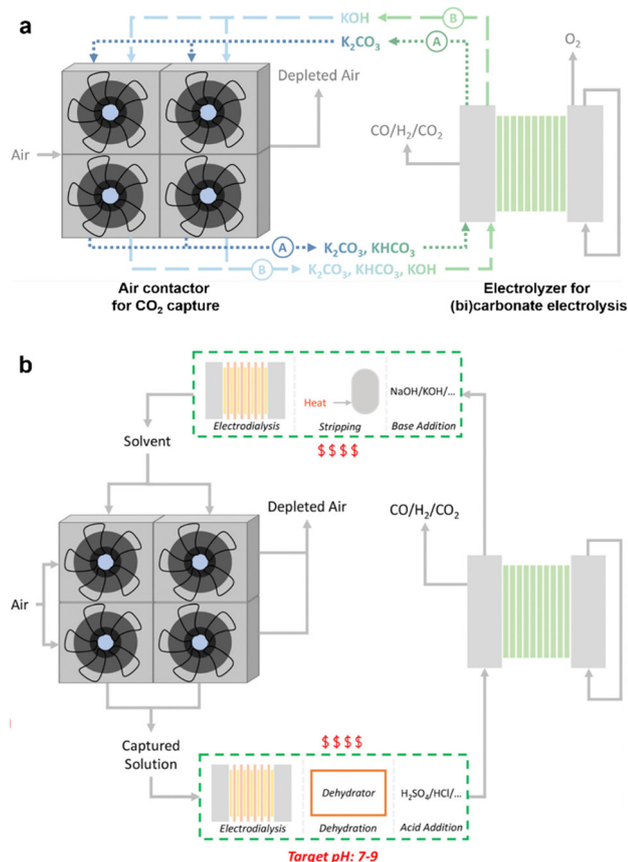


Fig. 13 Schematic of the integration route proposed in the literature, with the air contactors on the left and the electrolyzer stacks on the right. (a) The two pathways for CO₂ capture from the air are (A) K₂CO₃-based and (B) KOH-based. (b) Potential solutions to address the different pH requirements of the capture and conversion processes are shown inside the green dashed boxes at the top and bottom. Reproduced with permission from ref. 168.

on and attached to the cathode and oxygen at the anode. This process has been demonstrated on a laboratory scale to successfully convert ambient air into carbon and oxygen, producing, e.g., 10 g h⁻¹ of carbon nanofibers at 100 A.

These examples highlight the feasibility of integrating electrochemical conversion with CO₂ capture from the air. However, the efficiency of the integration is currently not satisfactory due to the slow CO₂ capture. One approach to overcome this limitation is to use a significantly larger amount of capture agent while vigorously feeding air into the solution under stirring. Alternatively, existing methods could be suitable for small, decentralized units with minimal time constraints. Integrating electrochemical conversion with DAC technology provides the flexibility to be deployed anywhere, making it ideal for reducing emissions from dispersed sources when combined with renewable energy for power generation. Nevertheless, the widespread implementation of DAC infrastructure presents challenges, including substantial investment and land use. Furthermore, DAC technology is relatively new and less mature, which may pose hurdles regarding scalability and cost-effectiveness, at least in the short term.

5.1.2. Flue gas. Flue gas refers to emissions from industrial operations, power generation, and combustion processes. The CO₂ content in flue gases is 2–3 orders of magnitude higher than in the atmosphere, depending on the fuel sources, such as coal (12–15% CO₂) and natural gas (3–4% CO₂). Consequently, the minimum energy input required to concentrate CO₂ in flue gas to a highly concentrated stream is reduced by a factor of three with respect to DAC systems. Therefore, extracting CO₂ from flue gas is more efficient than from the atmosphere, making it a promising approach for reducing CO₂ emissions.

In most cases, the capture agents and methodologies developed for DAC can also be applied to flue gas and *vice versa*. This is similarly true for the integrated CO₂ capture and conversion approach. However, significant amounts of gas impurities in flue gas, such as NO_x and SO_x, can cause considerable inefficiencies and increase cost, as they may interfere with both the capture and the conversion processes. For example, removing SO₂ from flue gas by converting it into CaSO₃ requires a substantial energy input of 380 kJ mol⁻¹.¹⁶⁹ Removing NO_x involves its selective reduction to N₂ over a supported vanadia catalyst using ammonia as a reductant. Although this chemical reaction is exergonic, ~500 kJ of energy is required per mol of NO_x removed in commercial systems.¹⁶⁹

Most studies on the electrochemical conversion of flue gas neglect these impurities by using simulated flue gas, which consists of a variable mixture of CO₂ and usually N₂. In general, the risk of interference from acidic impurities like SO₂ and NO₂ is minimal when CO₂ is captured using alkaline (bi)carbonate solutions, as these can neutralize such impurities.^{36,170,171} Among the NO_x gases, the relatively inert NO is usually the predominant component, requiring oxidation to NO₂ using, e.g., hydrogen peroxide for effective removal.¹⁷⁰ Unfortunately, even small amounts of impurities can negatively impact catalyst performance and, consequently, the conversion process. Therefore, several studies on gas-fed CO₂ have thoroughly investigated this issue, with findings highly relevant for integrated processes, especially those involving *in situ* generated CO₂.

Ko *et al.* examined the influence of NO_x on Cu, Ag, and Sn catalysts by introducing 8300 ppm NO into a CO₂ stream.¹⁷² For Ag and Sn catalysts, NO reduction reactions reduced FE by 35%. Similarly, introducing 10 000 ppm SO_x led to a decrease in FE by 25–40% when targeting CO or formate.¹⁷³ In both cases, the catalytic efficiency was restored upon reintroducing a pure CO₂ stream into the reactor. For the Cu catalyst, a significant shift in selectivity toward formate was observed, along with a decrease in the production of C₂₊ products, highlighting Cu's high sensitivity to SO₂ impurities.¹⁷³

Removing these impurities beforehand may be necessary, although it often requires a combination of specialized and expensive technologies for each impurity. A more effective approach would be to develop catalysts that can tolerate these impurities. This was demonstrated in a study,¹⁷⁴ where stability experiments were conducted with the presence of ~200 ppm SO₂ or NO in the feed gas stream using Bi₂O₃ (producing formate) and Ag (producing CO) as catalysts. Encouragingly,



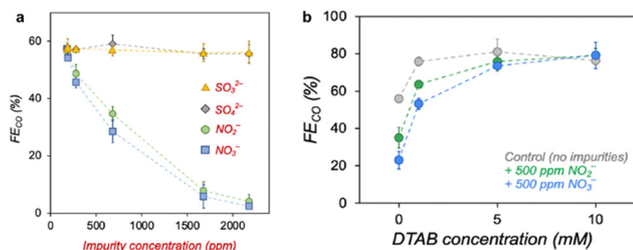


Fig. 14 FE_{CO} for electrolysis of 3 M KHCO₃ solution at 100 mA cm⁻² using a porous silver foam cathode with (a) 0–2000 ppm of flue gas impurities and (b) as a function of DTAB concentration. Reproduced with permission from ref. 81.

the results showed consistent performance and high FE (>90%) for both catalysts toward the target products over 20 h.

Zhang *et al.* evaluated the effects of nitrogen and sulfur contaminants (NH₄⁺, NO₃⁻, SO₄²⁻, and SO₃²⁻) on the catalytic performance of a porous silver electrode for the conversion of 3 M bicarbonate.⁷⁹ At concentrations of 100 ppm for these ions, there was no significant effect on FE_{CO} or FE_{H₂}, except for NO₃⁻, which reduced FE_{CO} from 56% to 28%. At 500 ppm NO₃⁻, no CO or H₂ was produced due to the competitive reduction of NO₃⁻. Later, Pimlott *et al.* conducted a detailed study on the impact of nitrogen- and sulfur-containing impurities at varying concentrations.⁸¹ They observed that adding anions such as SO₄²⁻ and SO₃²⁻ (originating from SO_x impurities) in 3 M KHCO₃, had no impact on product formation (Fig. 14a). In contrast, dissolved NO₂⁻ and NO₃⁻ ions (originating from NO_x impurities) at a concentration of 2000 ppm in the same electrolyte reduced FE_{CO} by up to 55% at a current density of 100 mA cm⁻². This decrease in FE_{CO} was attributed to the relatively more positive electrochemical potentials at which NO_x anions are reduced compared to CO₂, rather than catalyst degradation. Switching to a fresh KHCO₃ solution restored performance, indicating minimal long-term impact from NO₃⁻. This demonstrates that direct bicarbonate conversion is more tolerant to impurities than gaseous CO₂ conversion.

An alternative strategy to minimize the impact of NO_x impurities is to remove NO₂⁻ and NO₃⁻ from the bicarbonate solutions before they reach the electrolyzer, using methods such as reverse osmosis, ion exchange, distillation, or biodenitrification.¹⁷⁵ However, these processes can be cumbersome. To address this, Pimlott *et al.* demonstrated that adding surfactants like 10 mM dodecyltrimethylammonium bromide (DTAB) to the bicarbonate solution suppressed the reduction of aqueous NO_x impurities and increased the FE_{CO} from 56% to 80% at 100 mA cm⁻², even slightly higher than without impurities (Fig. 14b).⁸¹ This improvement occurs because the surfactant's alkyl chains suppress the HER and increase the local CO₂ concentration by displacing ions near the catalyst surface.^{157,176} From this perspective, adding surfactants to the electrolyte solution could be a cost-effective way to mitigate the poisonous effects of anionic contaminants during the conversion of captured CO₂.

Understanding the effect of oxygen on the electrochemical conversion process is crucial, given that it is often present in

flue gas streams at concentrations of several percent. A recent study examined an electrolyzer supplied with either CO₂ or bicarbonate.¹⁷⁷ When 10% O₂ was introduced into the CO₂ stream entering a gas-fed electrolyzer, CO selectivity decreased by over 90%. In contrast, the presence of O₂ in the bicarbonate solution did not affect the performance of a liquid-fed electrolyzer. Regardless of the O₂ concentration in the gas streams (ranging from 0–100%), the conversion of bicarbonate solutions consistently produced CO at a Ag/C electrode with a FE_{CO} of ~65% at a current density of 100 mA cm⁻². This indifference to O₂ is a very encouraging result for integrated CO₂ capture and electrochemical conversion. Nonetheless, a potential strategy to mitigate the adverse effects of O₂, even for the feed-stream approach, is to operate the electrolyzer at a high total current density to ensure all O₂ at the electrode is reduced. Although this approach entails higher electricity usage, the associated costs appear negligible for low O₂-containing feed streams (≤3% O₂) at 300 mA cm⁻².¹⁷⁴

Many industrial factories already have infrastructure for capturing and processing flue gas, which facilitates the integration of CO₂ capture with conversion. However, this does not change the fact that implementing CO₂ conversion technologies in industrial settings requires significant initial investments in infrastructure and equipment. Another limitation of flue gas capture is that it is confined to areas with substantial industrial activity, making it ineffective for addressing CO₂ emissions from dispersed sources such as transportation, buildings, or agriculture.

5.1.3. Ocean. The ocean acts as a major reservoir for carbon, absorbing CO₂ from the atmosphere through oceanic carbon sequestration. Dissolved CO₂ levels in surface ocean waters vary depending on factors such as temperature, pressure, and biological activity, with an average concentration of ~50 times greater than in the air. Ocean capture can be achieved using natural seawater or through alkalinity enhancement, where seawater is treated with alkaline substances to increase its capacity to absorb CO₂.^{165,178}

The potential for integrating ocean capture with electrochemical conversion is immense. Although studies to date have focused on the release/capture of CO₂ or the coupled capture/conversion approach (Fig. 1, Route 2), we will describe one such study as a source of inspiration for future integration efforts. Digdaya *et al.* reported a reactor operated using ocean water with a bipolar membrane electro dialysis (BPMED) cell followed by vapor-fed CO₂ reduction.¹⁶⁵ The BPMED cell utilized a one-electron reversible redox couple (Fe(CN)₆³⁻/Fe(CN)₆⁴⁻) at the electrodes instead of the conventional water-splitting reaction. This demonstrated efficient capture of CO₂ (released upon acidifying the bicarbonate-containing ocean water) at an electrochemical energy consumption of 155.4 kJ mol⁻¹ or 0.98 kWh kg⁻¹ of CO₂, with a CO₂ capture efficiency of 71%.

Once CO₂ was released and led to the vapor-fed electrochemical reduction, various fuels and chemicals, such as CO, CH₄, ethylene, and ethanol, could be produced with a total FE of up to 73% at current densities of 58 mA cm⁻² when using a Cu electrocatalyst. Similarly, CO could be selectively obtained with



a FE of up to 95% at a current density of 11.2 mA cm^{-2} using a Ag electrocatalyst. It is important to note that the energy required for CO_2 capture constitutes only a small fraction of the total capture and conversion energy. For example, converting CO_2 to CH_4 would require 13.9 kWh kg^{-1} of CO_2 . To date, this process has been executed coupled, with the CO_2 being released from the bicarbonate-containing ocean water and directed to the vapor-fed electrode. However, there is nothing preventing the integration of these steps, as bicarbonate can be electrochemically converted to products directly or indirectly (Table 1).

5.2. Continuous operation of CO_2 capture and conversion

With a comprehensive understanding of CO_2 capture processes and the conversion of captured CO_2 , the natural progression is to integrate these processes to streamline operations and enhance overall efficiency. Below, we highlight three studies that have carefully considered aligning the two processes for continuous operation. While the first two examples utilize simulated flue gas, the third employs pure CO_2 .

Wang and Luo discussed both a coupled and an integrated system designed for CO_2 capture, electrocatalytic reduction, and purification of gaseous products.²³ An alkaline solution was generated *via* a BPMED cell to capture CO_2 from the simulated flue gas ($\text{N}_2:\text{CO}_2 = 5:1$). The absorbed CO_2 was then released by acidifying the absorbent solution with the acidic solution generated from the BPMED cell. The liberated CO_2 was electrocatalytically reduced into CO at a Ag GDE using a flow-cell setup. However, due to low single-pass efficiency, the CO concentration reached only $\sim 10\%$ at 200 mA cm^{-2} and $\sim 15\%$ at 300 mA cm^{-2} . Nevertheless, any unreacted CO_2 could be absorbed by the alkaline solution produced by the BPMED cell, resulting in a pure stream of syngas. The authors also explored an integrated process, bypassing the CO_2 release process, by examining the performance of a Ag electrode directly in 1 M KHCO_3 (Fig. 1, Route 3). In this scenario, a FE_{CO} of 40% was achieved at a current density of 50 mA cm^{-2} . Unfortunately, the current density could not be further increased, as this led to the continuous decline in CO production, accompanied by a gradual increase in H_2 from the HER.^{88,94} Thus, the integrated approach will require significant optimization before it can be scaled up and operated stably.

Prajapati *et al.* integrated CO_2 capture and electrochemical conversion using a continuous flow system (Fig. 15).¹²⁹ To capture CO_2 , they employed a migration-assisted moisture-gradient (MAMG) process, where gaseous CO_2 was absorbed as HCO_3^- in a CO_2 -binding organic liquid. The captured CO_2 was then led through an anion exchange membrane under an electric field to an aqueous solution, where it was converted to dissolved CO_2 . For CO_2 reduction, they devised an electrochemical cell setup using a Cu mesh electrode, enabling the extraction of CO_2 -free products such as CO, CH_4 , and C_2H_4 . In practice, they operated the MAMG CO_2 capture unit at a current of 600 mA and the CO_2 reduction unit at a current density of 200 mA cm^{-2} , which achieved a 40% FE for C_2H_4 from a simulated flue gas feed and could be operated stably for

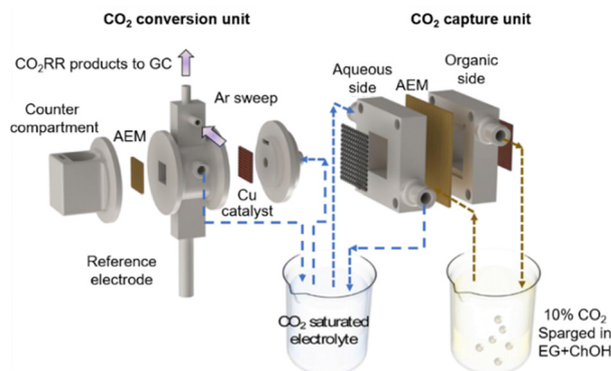


Fig. 15 Illustration of the integrated system featuring MAMG CO_2 capture and electrochemical conversion. Reproduced with permission from ref. 129.

24 h. To effectively integrate the CO_2 capture and CO_2 reduction processes without diminishing the overall efficiency or necessitating downstream purification or recycling of CO_2 , the rate of CO_2 capture must match the rate of CO_2 reduction.

Bai *et al.* developed a molten-salt-based electrolyzer capable of continuously capturing and converting CO_2 with 100% efficiency into CH_4 and CO, along with H_2 from the HER (Fig. 16).¹⁰¹ Notably, the CO_2 fed into the reactor was completely captured by the molten salt electrolyte by carefully optimizing the feeding rate. The highest CH_4 selectivity (33%) was achieved at 400 mA cm^{-2} using a commercial CuNi_{18} sheet as the cathode and a corrosion-resistant Ni sheet as the anode in a molten $\text{Li}_{1.427}\text{Na}_{0.359}\text{K}_{0.214}\text{CO}_3 \sim 0.15\text{LiOH}$ electrolyte at 650°C (reactor dimensions: diameter = 2 cm, length = 4 cm). Pure CO_2 was supplied to the reactor at a flow rate of 20 mL min^{-1} . Unlike previous molten-salt single cells based on the one-pot method, this reactor operated without the need for membranes and with a conversion efficiency of 100% for a single pass, eliminating the need to separate unreacted CO_2 from the gaseous products. Unfortunately, the operation of this system lasted for only 3.6 h, and a gradual increase in the electrolysis voltage was observed, showing its unsatisfactory stability. In general, molten salt systems require substantial energy input to maintain operations at high temperatures. In addition, for molten salt systems that generate carbonaceous

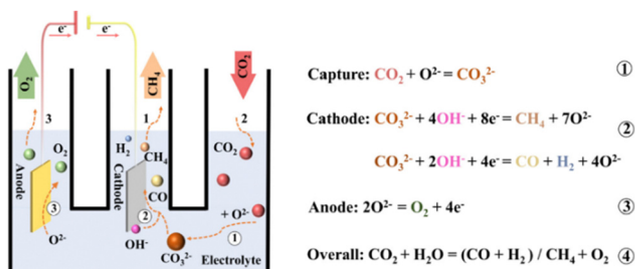


Fig. 16 Schematic illustration of a reactor for reactive CO_2 capture, featuring a commercial CuNi_{18} sheet as the cathode and a corrosion-resistant Ni sheet as the anode in molten $\text{Li}_{1.427}\text{Na}_{0.359}\text{K}_{0.214}\text{CO}_3 \sim 0.15\text{LiOH}$. Reproduced with permission from ref. 101.



products,^{131,132} regular pauses in production are necessary to remove carbon deposits accumulated on the cathode.

Although examples of the practical integration of CO₂ capture with subsequent conversion are limited, they have demonstrated the feasibility and potential of this technology. One approach involves running the two processes sequentially, with the conversion starting only after the capture is complete. The cycle can then restart after the conversion and regeneration of the capture agent. Alternatively, the conversion of captured CO₂ can be conducted concurrently with the capture process, which appears promising if the capture can be performed efficiently, as in the case of CO₂ from flue gas. However, if this is not the case, this approach may result in overall low conversion efficiency and selectivity issues due to the low steady-state concentration of the captured CO₂.

5.3. Energy cost analysis

An important aspect of the practical integration of CO₂ capture and conversion is the analysis of the energy costs of the entire process, from capturing CO₂ from dilute sources to generating the desired product. This analysis is crucial for assessing the feasibility of the technology from both technical and economic perspectives. Since CO is the most reported product from the electrolysis of both gas-fed CO₂ and captured CO₂ systems, we first compare the energy consumption for producing CO between independent and integrated CO₂ capture and electrochemical conversion (Table 2). In the analysis, we considered three key steps that incur the main energy costs in the entire chain of CO₂ capture and conversion, *i.e.*, CO₂ capture/release, electrolysis, and product separation. The energy consumption analysis used experimentally reported data, including faradaic efficiencies, full-cell potentials, and CO₂ utilization efficiencies. As a result, the calculated energy values inherently carry some degree of uncertainty due to the variability and limitations of the experimental conditions and measurements (see the ESI† for details).

The independent CO₂ capture and electrochemical conversion strategy (Fig. 1, Route 1) requires a substantial energy investment to release CO₂ from the capture media. For DAC technologies using potassium hydroxide solutions, the energy required to release CO₂ from the subsequently formed CaCO₃ is 178.3 kJ mol⁻¹.¹⁸³ Furthermore, in alkaline CO₂-fed electrolyzers, CO₂ gas can be lost due to its reaction with OH⁻ ions generated during the reduction reaction. Therefore, the CO₂ utilization efficiency, defined as the percentage of input carbon converted to a targeted product, is typically no greater than 20% for electrolyzers that generate CO.^{179,180} In other words, to obtain one mol of CO product, 5 moles of CO₂ need to be captured, thus leading to an energy consumption of 892 kJ mol-CO⁻¹ for this CO₂ regeneration step. As expected, the energy required for the CO₂ capture and release steps with CO₂-fed electrolyzers increases as the CO₂ utilization efficiency decreases. For the electrolysis step, gas-fed CO₂ electrolyzers, whether in flow cell or MEA configurations, require relatively low energy costs, ranging from 480 to 633 kJ mol-CO⁻¹. This is due to the high level of optimization of the electrocatalysts for

Table 2 Comparison of the energy requirements between integrated (Route 3) and independent (Route 1) pathways for the production of CO, HCOO⁻, CH₄, and C₂H₄^a

Route	System	Product	CO ₂ utilization (%)	CO ₂ regeneration ^b (%)	FE (%)	Electrolysis EE (%) ^c	Electrolysis ^b (%)	Outlet CO ₂ (%)	Product purity (vol%)	Product separation ^{bd} (%)	Total energy ^b (kJ mol ⁻¹)	CCU EE (%)	Ref.
Route 1	CO ₂	CO	20	892	77	41	633	50	26	1250	2775	9	179
	CO ₂	CO	17	1049	99	54	480	36	32	1050	2579	10	180
Route 3	HCO ₃ ⁻	CO	60	0	70	25	1014	25	54	200	1214	21	90
	CO ₃ ²⁻	CO	100	0	25	35	734	0	25	0	734	35	72
	Carbamate	CO	90	0	72	40	634	0	72	0	634	41	85
Route 1	CO ₂	HCOO ⁻	9	1981	96	29	917	N.A.	N.A.	265	3163	9	181
Route 3	HCO ₃ ⁻	HCOO ⁻	70	0	85	40	499	0	0.8 M	265	764	26	94
Route 1	CO ₂	CH ₄	1	13715	62	16	5495	99	1	38000	57210	1	182
Route 3	HCO ₃ ⁻	CH ₄	25	0	27	4	20584	75	7	1500	22084	4	98
	Molten salt	CH ₄	100	0	43	21	36989 ^e	0	16	0	36989	3	101
Route 1	CO ₂	C ₂ H ₄	5	3566	70	34	3974	77	3	15400	22940	6	19
Route 3	CO ₃ ²⁻	C ₂ H ₄	100	0	34	10	13971	0	8	0	13971	10	103

^a See the ESI for details on the calculation procedure. ^b Unit: kJ mol-product⁻¹. ^c EE: Energy efficiency. ^d Separation energy of unreacted CO₂ from reaction products. ^e Electrolysis energy without thermal considerations is 4881 kJ mol-product⁻¹; an extra 32 108 kJ mol⁻¹ has been added due to the process being conducted at 650 °C (see the ESI).



the CO₂-to-CO conversion, resulting in systems displaying high FEs, large current densities, and low cell voltages.

After electrolysis, the outlet gas stream consists of CO, unreacted CO₂, and H₂ byproduct, which requires separation before obtaining pure CO. Greenblatt *et al.* outlined general approaches to separating product mixtures resulting from CO₂ valorization.¹⁸⁴ The unreacted CO₂ and unwanted H₂ gases can be separated through, *e.g.*, pressure/temperature-based swings, adsorption/desorption, and membrane separation. Since the CO₂ utilization efficiency is low, the percentage of unreacted CO₂ at the outlet gas stream can be as high as 50%, and the purity of CO is estimated to be 26%.¹⁷⁹ Such a low CO purity leads to high energy consumption for the separation step, estimated at 1250 kJ mol-CO⁻¹. In total, the energy consumption from cradle to grave is 2775 kJ mol-CO⁻¹. Given that the theoretical energy consumption for converting gas-fed CO₂ to CO is 257 kJ mol-CO⁻¹,³⁸ the global carbon capture and utilization (CCU) energy efficiency for the entire chain is 9%.

In the case of the electrochemically reactive capture of CO₂ (Fig. 1, Route 3), the absence of energy costs associated with the CO₂ capture/release step is a significant advantage, highlighting the potential of this technology in reducing energy consumption. The primary energy cost in the integrated pathway is the electrochemical conversion of the captured CO₂ solution. For instance, in the electrochemical conversion of a bicarbonate solution to CO,⁹⁰ the energy consumption for this step is 1014 kJ mol-CO⁻¹, which is higher than for the electrolysis of gas-fed CO₂. However, the CO₂ utilization efficiency for converting bicarbonate is 60%, significantly higher than that of gas-fed CO₂ electrolyzers. Since some *in situ* generated CO₂ exits the electrolyzer without being converted to CO, the concentration of CO in the outlet gas stream is 54%, thus requiring 200 kJ mol-CO⁻¹ to separate the small amount of unconverted CO₂. The total energy consumption for the entire process, from CO₂ capture in dilute sources to CO production *via* a bicarbonate system, amounts to 1214 kJ mol-CO⁻¹, which is significantly lower than for the independent CCU process. Moreover, the global CCU energy efficiency of the integrated pathway is 21%, twice as high as that of the independent CCU process.

In the case of carbonate and carbamate solutions, the *in situ* generated CO₂ can be quantitatively converted to CO, as almost no CO₂ is detected in the outlet gas stream.^{72,85} This suggests that the CO₂ utilization efficiency for both cases is nearly 100%. In this scenario, the total energy consumption for converting carbonate or carbamate solutions can be estimated at 734 and 634 kJ mol-CO⁻¹, respectively. These values are approximately four times lower than those for independent systems, highlighting their potential for practical applications (Table 2). The global CCU energy efficiencies are calculated to be 35% for the carbonate system and 41% for the carbamate system, significantly higher than those achieved with CO₂-fed electrolyzers. Notably, if the targeted product is syngas (a combination of CO and H₂), the separation energy can be considered zero since no unreacted CO₂ is present in the outlet gas stream.

The estimation of the separation energy for the CO product focused exclusively on the energy required to remove unreacted

CO₂, without accounting for the separation of the H₂ byproduct. This approach is based on several considerations. First, CO₂ removal is the most energy-intensive step,¹⁸⁵ while extracting H₂ from syngas after CO₂ removal requires significantly less energy. For example, the energy cost for CO₂ separation can vary widely, ranging from 100–900 kJ mol⁻¹ of CO₂, depending on the separation method used. For the energy calculations, a fixed value of 500 kJ mol⁻¹ of CO₂ was assumed for separating unreacted CO₂ from the gaseous output.⁷² In contrast, the energy required for H₂ removal from syngas (with a ratio of ~1:3 CO:H₂) is about 85 kJ mol-H₂⁻¹.¹⁸⁶ Secondly, the H₂ concentration in the gas stream is typically much lower than that of unreacted CO₂ or the primary product, CO. As a result, the energy cost for H₂ separation generally accounts for a small fraction of the overall energy consumption for various CO₂ capture and conversion methods.

For instance, in cases where FE_{CO} ≥ 70%, the estimated energy cost for H₂ separation is ≤ 34 kJ mol-CO⁻¹, representing < 2% of the total energy consumption for both Route 1 scenarios and under 5% for the two Route 3 scenarios (*e.g.*, bicarbonate and carbamate conversion). In the context of carbonate conversion, where FE_{CO} is only 25% (*i.e.*, CO/H₂ = 1:3), the estimated energy cost for H₂ separation rises to 255 kJ mol-CO⁻¹. Even with this increased cost, the total energy consumption for Route 3 (*e.g.*, carbonate conversion, 989 kJ mol-CO⁻¹) remains lower than that for the two Route 1 scenarios (2775 and 2559 kJ mol-CO⁻¹, respectively). It is important to note that the energy cost for H₂ separation heavily depends on the gas stream composition and the required purity levels of CO and H₂. Consequently, the cost will vary based on the specific components of the gas stream. These factors introduce complexity and result in uncertainties that are challenging to quantify. Given the relatively minor contribution of H₂ separation and its associated complexities, these energy costs are not included in Table 2.

Among the different types of captured CO₂ (*i.e.*, bicarbonate/carbonate/carbamate), the energy consumption for converting carbamate to CO/syngas is the lowest, which aligns with the weaker binding energy of amines with CO₂ compared to hydroxide solutions. Therefore, from an energy consumption standpoint, integrating CO₂ capture by amine with the electrolysis of the resulting carbamate to produce CO/syngas is the most advantageous option.

Similar conclusions can be drawn for the generation of HCOO⁻ as the product. In the case of CO₂-fed electrolyzers, due to the low CO₂ utilization efficiency (9%), 1981 kJ mol-formate⁻¹ of energy is required for the CO₂ capture and release steps.¹⁸¹ Additionally, considering the electrolysis and purification steps, the total energy consumption amounts to 3163 kJ mol-formate⁻¹, corresponding to a global CCU energy efficiency of 9%. In contrast, the direct electrochemical upgrading of a bicarbonate solution in the integrated process requires only 764 kJ mol-formate⁻¹, leading to a global CCU energy efficiency of 26%.⁹⁴ These efficiency values demonstrate the promising potential of the integrated strategy in producing an important chemical such as HCOO⁻.



A key difference between the systems producing CO and those generating HCOO^- lies in the purification methods to separate HCOO^- from the liquid electrolyte solution. For one thing, the formate after electrolysis usually has a low concentration (0.8 M) in the electrolyte, which brings difficulty for separation (Table 2). For another, since HCOO^- is a charged product that can combine with electrolyte cations to form salts, an additional acidification step is required to obtain the more valuable formic acid. The resulting formic acid in water can then be separated by azeotropic distillation or extraction, which requires 265 kJ mol^{-1} .¹⁸⁷

In producing highly reduced products such as CH_4 and C_2H_4 , CO_2 -fed electrolyzers typically exhibit low CO_2 utilization efficiency. For example, electrolyzers converting CO_2 into CH_4 have demonstrated efficiencies of $\sim 1\%$,¹⁸² while those producing C_2H_4 achieve $\sim 5\%$.¹⁹ These low efficiencies result in high energy costs associated with the CO_2 capture and release steps. Moreover, for both independent and integrated pathways, the energy costs of the electrolysis step are significantly higher compared to electrolyzers that generate CO or HCOO^- . This is due to the greater number of electrons required to convert CO_2 into CH_4 (8 electrons) or C_2H_4 (12 electrons), as opposed to the 2 electrons needed for producing CO or HCOO^- , substantially increasing the energy consumption of the electrolysis. For instance, in the direct electrochemical conversion of a bicarbonate solution to CH_4 ,⁹⁸ the energy required for the electrolysis step can be calculated to amount to $20\,584 \text{ kJ mol}^{-1}$. However, when considering the CO_2 capture/release and product purification steps, the direct conversion of bicarbonate into CH_4 has a lower total energy consumption ($22\,084 \text{ kJ mol}^{-1}$) compared to CO_2 -fed electrolyzers ($57\,210 \text{ kJ mol}^{-1}$). The higher concentration of CO_2 in the electrolyzer outlet makes the independent pathway more energetically expensive, as more energy is required to separate the unreacted CO_2 from CH_4 than in the integrated approach.

An energy analysis of CH_4 production using molten salts indicates that even in an electrolysis step carried out at 650°C , the integrated strategy is energetically more favorable.¹⁰¹ With molten salts, the CO_2 utilization efficiency is exceedingly high, and the energy consumption for the electrolysis step is relatively low (4881 kJ mol^{-1}). However, additional energy is required to heat the electrolyzer to 650°C , amounting to $32\,108 \text{ kJ mol}^{-1}$. This brings the total energy cost to $36\,989 \text{ kJ mol}^{-1}$, which is still lower than the total energy costs for producing CH_4 using a CO_2 -fed electrolyzer ($57\,210 \text{ kJ mol}^{-1}$).

Based on the energy consumption analysis shown in Table 2, we conclude that the integrated CO_2 capture and electrochemical conversion strategy (Fig. 1, Route 3) is more energy-efficient than independent processes (Fig. 1, Route 1), regardless of the products formed. This makes the integrated approach potentially more economical. The analysis also suggests that CO or syngas would be the most feasible initial targets, followed by HCOO^- , as the total energy consumption for producing CO and HCOO^- from dilute CO_2 sources *via* electrochemically reactive carbon capture is lower than for other products such as CH_4

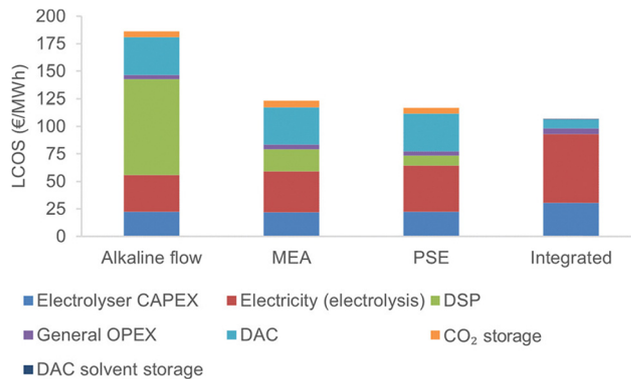


Fig. 17 Comparison of the LCOS of different routes of capture and electrochemical conversion of CO_2 from air. The terms "alkaline flow," "MEA," and "PSE" (porous solid electrolyte) refer to the independent route using these respective cell configurations. The "General OPEX" category accounts for operational and maintenance costs. Reproduced with permission from ref. 188.

and C_2H_4 . This conclusion is further supported by Debergh *et al.*,¹⁸⁸ who examined the economics of electrochemical syngas production *via* DAC. The authors compared the levelized cost of syngas (LCOS) for different CO_2 capture and electrochemical conversion routes from air. They found that the integrated route offers the lowest LCOS due to reduced downstream separation (DSP) and CO_2 capture costs from the air (*i.e.*, DAC), even though it incurs higher electrolysis costs (Fig. 17). Furthermore, their analysis revealed that CO is the most promising target for the integrated route among the various products.

Highly reduced products such as CH_4 and C_2H_4 have important applications as chemical feedstocks. To improve the economic viability of producing these valuable compounds *via* the integrated route, it is essential to reduce the overall energy consumption. A significant portion of the energy used in these processes is attributed to the direct electrolysis of captured CO_2 solutions. Currently, the electrocatalytic materials employed in these processes suffer from high overpotentials and low selectivity for CH_4 and C_2H_4 , which drives up energy costs for electrolysis. Addressing this challenge requires developing novel electrocatalysts optimized for directly upgrading captured CO_2 solutions to lower energy consumption and enhance the economic competitiveness of CH_4 and C_2H_4 production. Methanol and ethanol, both valuable as fuels and chemical intermediates, can be produced through the electrochemical conversion of gas-fed CO_2 . However, there are currently only a few reports of direct electrochemical conversion of captured CO_2 into these products with low selectivity (*e.g.*, $\text{FE}_{\text{ethanol}} < 20\%$).¹⁰² Therefore, developing efficient and selective catalysts for the integrated process in these cases is essential for accurately evaluating their energy requirements and economic viability.

It should be noted that the integrated route for CO_2 capture and conversion demonstrates a similarly high CO_2 utilization efficiency as thermochemical CO_2 conversion by hydrogenation, typically ranging between 70% and 90%.^{189–191} The CO_2



hydrogenation reactions are generally conducted at temperatures between 300 and 700 °C, enabling the production of valuable products such as CO, HCOOH, CH₃OH, and CH₄. While both CO₂ valorization methods achieve comparable CO₂ conversion efficiencies, thermochemical processes tend to produce higher CO₂ emissions due to their energy-intensive conditions, unless powered by carbon-neutral heat sources.

5.4. Stability evaluation

The energy costs mentioned above are based on electrolysis data obtained from short-term experiments, typically lasting several hours to tens of hours. For industrial-scale applications, it is crucial to comprehensively study the long-term stability of integrated CO₂ capture and electrochemical conversion systems operating at high current densities. Unfortunately, only relatively few studies have addressed this issue.

Table 3 summarizes studies investigating the stability of electrochemical conversion of captured CO₂ for at least 5 hours at current densities of ≥ 50 mA cm⁻². In an early report, Li *et al.* demonstrated the stable operation of an integrated system for 145 h using a Ag catalyst (Fig. 18a and b).⁷² Here, a KOH solution was used to capture CO₂, resulting in the formation of carbonate, which was then reduced to syngas while regenerating the KOH solution for another round of CO₂ capture *via* electrolysis. Throughout the operation, the current density remained stable at ~ 180 mA cm⁻², and the H₂/CO ratio consistently ranged between 2 and 3. However, slight fluctuations in the syngas ratio and a decrease in FE_{CO} were observed, attributed to metal contamination and deposition over time.

Later, Zhang *et al.* conducted electrolysis of a 3 M KHCO₃ solution over 80 h at an applied current density of 65 mA cm⁻² using a free-standing porous silver electrode.⁷⁹ Throughout the experiment, FE_{CO} decreased by only 3% and the cell voltage increased by just 100 mV, demonstrating the good stability of the system. However, this stability was achieved by manually refreshing the 3 M KHCO₃ electrolyte every 500 s. When the free-standing electrode was replaced with silver-carbon composite electrodes, a much larger decrease in FE_{CO} (16%) was observed over the 80 h electrolysis. Both Li *et al.* and Lees *et al.* also demonstrated that refreshing the captured CO₂ electrolyte was essential to maintaining good stability in their electrolysis systems.^{75,76} Without refreshing the electrolyte, a gradual decrease in FE_{CO} was observed due to the consumption of (bi)carbonate, resulting in an increase in pH over electrolysis time (Table 3).

Recently, Song *et al.* investigated the electrolysis of a 3 M KHCO₃ solution with continuous CO₂ purging for 29 h using a Ni SAC. They found that the system remained very stable for continuous CO production with FE_{CO} of $>90\%$ during the first 18 h due to the regeneration of bicarbonates by the CO₂ purging (Fig. 18c).⁹⁰ Under such conditions, electrolyte refreshing had no impact on the electrolysis performance, and therefore, it was not required. In contrast, the electrolysis of gas-fed CO₂ and 3 M KHCO₃ without CO₂ purging showed a rapid decline in FE_{CO} under the same conditions. In addition, a slight decrease of FE_{CO} to $<90\%$ was observed after 20 h of

Table 3 Stability assessment of electrochemical conversion of captured CO₂ (since 2019)^a

Captured CO ₂	$ j $ (mA cm ⁻²)	Cell voltage (V)	Product FE	Duration of electrolysis (h)	Reason for performance decrease	Ref.
Captured CO ₂ in 2 M KOH	~ 180	3.8	CO: 33–25%	145	Contamination by metal deposition	72
3 M KHCO ₃	100	~ 3.5	CO: 37–27% ^b	5	Consumption of KHCO ₃ and pH increase	75
3 M KHCO ₃	100	3.5–3.7	CO: 42–32% ^b	8	Consumption of KHCO ₃ and pH increase	76
3 M KHCO ₃	65	3.4–3.5	CO: 88–85% ^b	80	Consumption of KHCO ₃ and pH increase	79
Captured CO ₂ in 2 M KOH	100	3.3	CO: 40–36%	23	N.A.	73
1 M K ₂ CO ₃	200	~ 3.7	CO: 42–35%	25	N.A.	74
3 M KHCO ₃ with CO ₂ purging	200	~ 3.2	CO: 95–86%	29	Partial detachment of Ni SAC	90
Captured CO ₂ in 2 M KOH	200	3.5–3.8	C ₂ products: 42–32%	22.5	Degradation of the CEM interposer pore structure	103
Captured CO ₂ in 2 M KOH	100	3.4–3.5	C ₂ products: 22–18%	50	Reduction of exposed Cu(OH) ₂ nanowire catalyst	105
5 M K ₂ CO ₃	100	~ 3.7	C ₂ H ₄ : 10–9%	23	Consumption of K ₂ CO ₃ ; degradation of Cu–Ag catalyst	104

^a Data selected with $|j| \geq 50$ mA cm⁻² and electrolysis time of ≥ 5 h. ^b Electrolyte refreshed at regular intervals.



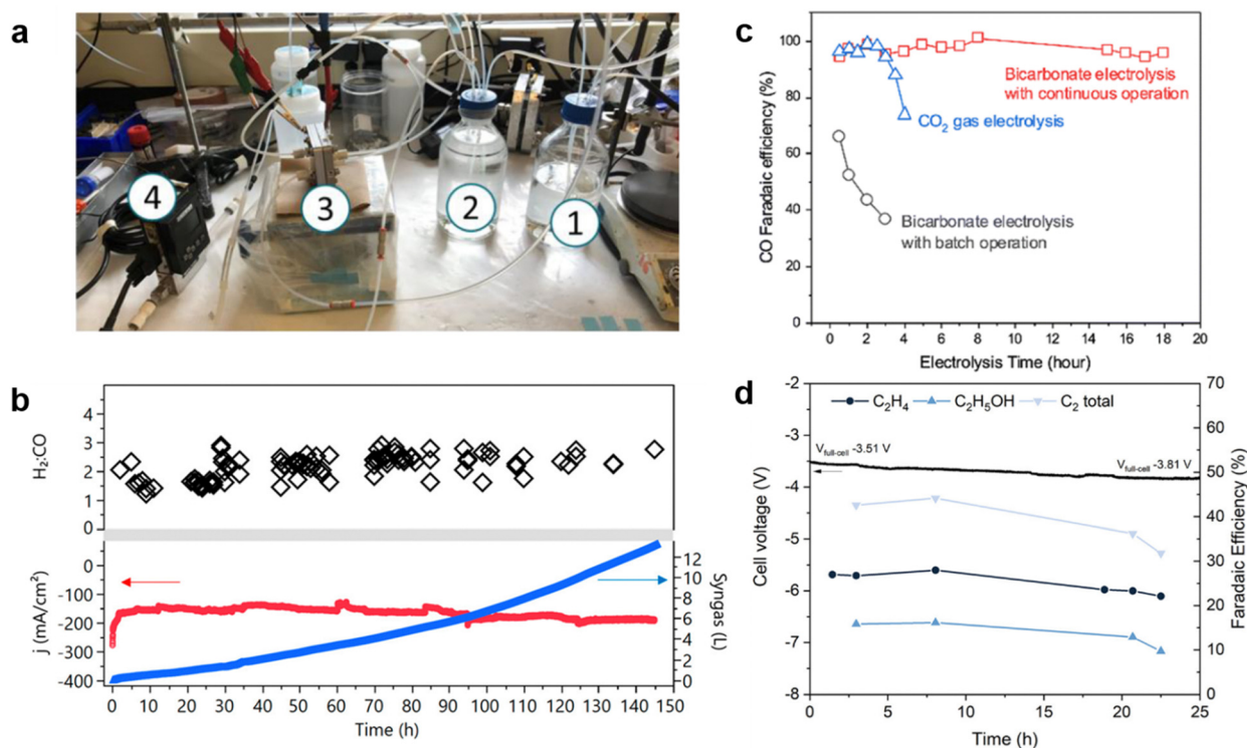


Fig. 18 Stability evaluation of the integrated CO₂ capture and electrochemical conversion. (a) Experimental setup, where CO₂ is captured into 2 M KOH (1) to generate carbonate, which is then pumped into a new bottle (2) and reduced in a direct carbonate cell (3). The gas products are measured by a mass flow meter (4) to determine the total volume. (b) Syngas production performance during electrolysis at a constant potential of 3.8 V using a Ag catalyst with the same experimental setup. Reproduced with permission from ref. 72. (c) Stability comparison of electrolysis between gas-fed CO₂ and 3 M KHCO₃ with/without CO₂ purging at a constant current density of 200 mA cm⁻² using a Ni SAC. Reproduced with permission from ref. 90. (d) Long-term operation for CO₂ capture and electrolysis in a 2 M KOH solution, performed with a Cu/CoPc-CNTs electrocatalyst and a 135 mm MEC interposer. Reproduced with permission from ref. 103.

electrolysis, which was attributed to the partial detachment of the catalysts.

Similar stability was observed in the electrochemical conversion of (bi)carbonate solution to C₂ products, such as C₂H₄. Ma *et al.* observed a decrease in FE for C₂ products and increased cell voltage during the first 10 h of electrolysis of captured CO₂ in a 2 M KOH solution using a Ag-Cu(OH)₂ nanowire catalyst.¹⁰⁵ The authors attributed this initial performance deterioration to the reduction of the exposed Cu(OH)₂ nanowires. After this phase, the system remained stable for 40 h of operation. Song *et al.* evaluated the stability of converting 5 M K₂CO₃ solution to C₂H₄ using a Cu-Ag catalyst at an applied current density of 100 mA cm⁻².¹⁰⁴ The FE for C₂H₄ was stable for the first 9 h but decreased after 18 h, attributed to various factors, including the consumption of K₂CO₃ and degradation of the Cu-Ag catalyst.

Lee *et al.* constructed a prototype system capable of operating CO₂ capture and electrochemical conversion to continuously produce C₂ products, similar to a previously reported system by the group (Fig. 18a).¹⁰³ The authors employed a Cu/CoPc-CNT catalyst (CoPc denotes cobalt phthalocyanine) and used a porous hydrophilic mixed cellulose ester (MCE) as the interposer to establish a well-defined spacing between the catalyst layer and the ion exchange membrane. They demonstrated continuous

operation of the setup for 20 h at a current density of 200 mA cm⁻². However, they observed a decline in performance after 8 h of operation (Fig. 18d), which was attributed to the degradation of the pore structure in the MCE membrane. This resulted in an increase in cell voltage and HER.

These results highlight the importance of optimizing the catalyst, electrolyte, interposer materials, and electrode configuration to enhance the stability of electrolysis of the captured CO₂ solution under relevant conditions. Note that during continuous operation, the electrolyte is refreshed through the reaction of CO₂ with the continuously basified electrolyte. One potential solution to address stability issues is the incorporation of additives or specifically engineered functionalities to prevent degradation of the catalyst and interposer. Until now, most stability studies have concentrated on the (bi)carbonate system, with limited research on the carbamate system. Furthermore, continuous CO₂ capture and electrochemical conversion over extended periods (≥1000 h) has not yet been demonstrated experimentally for either system, a crucial milestone for ensuring long-term durability in CO₂ capture and conversion.

5.5. Scalability consideration

Scalability is another important metric for integrated CO₂ capture and electrochemical conversion that has not yet been



Table 4 Summary of the three typical approaches for electrochemically reactive CO₂ capture

Systems	RNH ₂ (aq.)	OH ⁻ /CO ₃ ²⁻ (aq.)	Molten oxide
Captured CO ₂	RNHCOO ⁻ (aq.)	CO ₃ ²⁻ /HCO ₃ ⁻ (aq.)	Molten carbonate
Direct or indirect conversion	Both possible	Both possible	Direct
Main products	CO, HCOO ⁻	CO, HCOO ⁻ , CH ₄ , C ₂ H ₄	C, CO, CH ₄
Key benefits	Room-temperature operation; fast CO ₂ absorption kinetics	Room-temperature operation; improved stability and durability; larger j and FE for product formation	More compatible with flue gas systems; larger j and FE for product formation
Key challenges	Prone to thermal and oxidative degradation; lower j and FE for product formation	Inefficiency of CO ₃ ²⁻ (aq.) in capturing CO ₂ from the air	Operation at high temperature; batch production; interruptions due to buildup of solid carbon products on the cathode

thoroughly explored in the literature. The production rate, which reflects the scalability of a specific target product, is determined by the current density and FE achieved during electrolysis, as well as the geometric area of the electrode. Therefore, increasing these parameters is expected to enhance the production of target products. It should be noted, however, that there is a trade-off between current density and electrode area when aiming for a specific production rate. The current density determines the size of the electrolyzer. At lower current densities, the conversion of captured CO₂ is more energy efficient and incurs lower operating costs due to reduced energy losses from internal resistance, but this results in larger electrolyzers, increasing capital costs. In addition, it should be noted that scaling up production can be achieved either by increasing the size of the electrolyzer or by stacking multiple electrolyzers. An optimal current density balances operating and capital costs, estimated to be 200–400 mA cm⁻² for the electrochemical conversion of gas-fed CO₂.^{192,193} Such current densities are achievable for the electrolysis of captured CO₂ in many systems (Table 1).

To determine the electrolyzer size required to produce CO at a scale of 1 ton day⁻¹, we may use the optimal performance metrics reported by Song *et al.* (Table 3).⁹⁰ This calculation shows that operating electrolysis at a current density of 200 mA cm⁻² with FE_{CO} = 90% can achieve a CO production rate of 22.6 kg m⁻² day⁻¹. Accordingly, an electrolyzer with a total electrode area of 44.2 m² or an electrolyzer stack reaching this total area would be needed to reach the target production of CO. Doubling the current density is expected to reduce the electrolyzer size by half. Based on this calculation, scaling up the production of other products, such as C₂H₄, would require significantly larger electrolyzers due to the lower FE currently achieved for this product (<40%, Table 1).

6. Conclusions and perspective

6.1. Summary

The electrochemical conversion of captured CO₂ has garnered increasing attention in recent years due to its advantage of bypassing the traditional CO₂ release, purification, compression, transport, and storage steps. In this review, we outlined the four key factors that influence the success and applicability of this emerging technology. Table 4 summarizes the

three typical approaches for electrochemically reactive CO₂ capture.

(1) The choice of capture medium plays a crucial role in determining the thermodynamics and kinetics of the CO₂ absorption process and its subsequent electrochemical conversion. Each capture medium reviewed exhibits unique chemical properties that can be exploited to minimize the energy penalties associated with both capture and electrochemical conversion. Aqueous amines are the most widely studied medium for post-combustion CO₂ capture, although they are particularly vulnerable to thermal and oxidative degradation. Due to the strength of the C–N bond, the electrochemical reduction of the carbamate typically results in low current density values (<50 mA cm⁻²). Capturing CO₂ as a (bi)carbonate solution using aqueous carbonate and hydroxide solutions has emerged as a more economical and thermally stable alternative to amine scrubbing. The reduction of (bi)carbonate has shown higher reaction rates than carbamate reduction. However, several acid–base equilibria between CO₂ and (bi)carbonate complicate mechanistic investigations to identify the electrochemically active species at the electrode surface. Molten alkali carbonates are particularly beneficial for capturing CO₂ from flue gas, as this medium is compatible with the high temperature of such streams. Electrochemical conversion of captured CO₂ in molten carbonates typically occurs at temperatures ranging from 500–950 °C, with the added advantage of generating solid carbon in many cases, thereby eliminating the need for costly separation and purification processes.

(2) Various carbonaceous products can be produced through the electrochemical conversion of captured CO₂. The product types are determined mainly by the choice of catalysts, although the capture medium can also have an impact. For the electrochemical conversion of aqueous carbamate and (bi)carbonate solutions, CO is typically generated using Ag nanoparticles or SACs, formate is produced over Bi and Sn, while CH₄ and C₂H₄ are formed using Cu-based catalysts. This pattern mirrors that observed in the electrochemical conversion of gas-fed CO₂. In molten carbonate systems, the product outcome is more temperature dependent, with CO formed at ~900 °C and carbon materials at lower temperatures (~500 °C). Notably, the conversion of captured CO₂ can follow either an indirect pathway, where CO₂ is generated *in situ* and converted, or a direct pathway, where the captured CO₂ is directly converted at the electrode.



(3) Several other factors also significantly impact the electrochemical conversion of captured CO₂. Key elements such as the electrolyzer design, electrode materials, electrolyte composition, ion exchange membranes, and operating conditions (e.g., temperature, pressure, and flow plate design) can all affect conversion efficiency. These factors influence the availability of *in situ* generated CO₂ by shifting the dissociation equilibrium between carbonate, bicarbonate, or carbamate, and free CO₂, as well as impacting proton production and transport. Thus, optimizing these operational parameters is essential for improving the overall conversion efficiency.

(4) Based on these understandings, integrating CO₂ capture with subsequent conversion for practical applications becomes feasible. The outcomes of this integration will vary depending on the source of CO₂ (i.e., air or flue gas), as each source contains different levels of CO₂, O₂, and gas impurities. These factors impact, e.g., the availability of the CO₂ reactant, the charge transfer processes (e.g., CO₂ vs. oxygen reduction), and catalyst poisoning. For effective integration, the rates of capturing CO₂ and its subsequent conversion should be aligned. A comparison of energy costs between the conventional independent route (Fig. 1, Route 1) and the integrated route (Fig. 1, Route 3) shows that the integrated route consumes less energy and, as a result, achieves higher overall efficiency for the entire process, spanning from CO₂ capture from dilute sources to the production of the final products.

6.2. Challenges and potential solutions

Although previous studies have highlighted the significant potential for the electrochemical conversion of captured CO₂, several key challenges impede the rapid advancement of this emerging field and its large-scale implementation.

(1) The mechanism behind the conversion of captured CO₂ remains a topic of debate. It is not always clear whether the CO₂-bound capture agent itself or the CO₂ released from it is the actual species converted at the electrode. Most reports show that (bi)carbonate and carbamate are converted *via* the *in situ* generation of CO₂ at the electrode interface, although some reports indicate that they can also be directly converted. Due to the equilibrium between (bi)carbonate/carbamate and CO₂ during electrochemical conversion, determining the exact reaction pathway using only *ex-situ* techniques can be difficult. Kinetic studies are crucial for this clarification, and *in situ* *operando* techniques, such as infrared reflection-absorption spectroscopy and Raman spectroscopy, are valuable for identifying the adsorbed reactive species (bicarbonate/carbamate vs. CO₂) at the catalytic sites during the conversion process.^{194,195} For example, Lu *et al.* recently reported an electrolysis optical coherence tomography platform to visualize the chemical reactions occurring in a CO₂ electrolyzer, including reactants, intermediates, and products.⁷⁷ This platform, or a similar one, may serve as an effective tool for monitoring the dynamic movement of the various species during the electrolysis of captured CO₂, providing valuable insight into the mechanism. Furthermore, theoretical simulations based on DFT calculations can be employed to determine the adsorption energy of

reactive species and intermediates, as well as the free energy change of each elementary step in the conversion process.^{196,197}

(2) In general, the conversion efficiency for captured CO₂ is low. Over the past decades of intensive research, the electrochemical conversion of gas-fed CO₂ can now be achieved at a large current density (> 1 A cm⁻²) and with high FE (> 95%) for producing CO and formate.^{198,199} In comparison, the current density (usually < 200 mA cm⁻², although 400 mA cm⁻² can be achieved under certain conditions) and the product FE (usually < 80%, with 90% achievable under certain conditions) are much lower for the electrochemical conversion of captured CO₂ (Table 1), making the technology less efficient. The main reason for this is the low CO₂ concentration at the electrode/electrolyte interface, caused by the slow dissociation of carbonate/bicarbonate/carbamate to generate *in situ* CO₂ in the indirect conversion or the electrostatic repulsion between the negatively charged electrode and the carbonate/bicarbonate/carbamate ions in the direct conversion. Different strategies should be adopted to overcome these obstacles. For instance, modifying the electrode configuration, ion exchange membrane, and even the anolyte can enhance proton production and transport, thereby increasing the generation of *in situ* CO₂ and improving the conversion process. Drawing inspiration from electrochemical nitrate conversion, where negatively charged species are also involved, functionalizing the electrode surface with positively charged groups can reduce electrostatic repulsion and concentrate carbonate/bicarbonate/carbamate ions, promoting their direct conversion.^{200,201} Additionally, designing new catalysts specifically tailored for captured CO₂ conversion is crucial to enhance efficiency.

(3) Inefficient integration of CO₂ capture and conversion can hinder practical applications. To date, most studies have focused on improving the electrochemical conversion efficiency of captured CO₂ (i.e., carbonate, bicarbonate, or carbamate). A few examples demonstrate that integrating the two processes is feasible at the proof-of-concept stage. However, these reports rarely address the recyclability of the resulting electrolyte for CO₂ capture, nor do they explore the impact of different CO₂ sources or the long-term stability of the capture-conversion cycles. Currently, the electrolysis of captured CO₂ has been conducted for no longer than 150 h, with performance degradation arising mainly due to electrolyte basification and catalyst deactivation. Electrolyte basification can be mitigated by continuous and efficient CO₂ capture, along with a judicious selection of the capture medium, tailored to the specific operating conditions.

While air has a low concentration of CO₂, flue gas has a much higher CO₂ concentration but also includes gas impurities such as NO_x and SO_x. In an air capture scenario, a medium with selective and fast CO₂ uptake is necessary to concentrate the low levels of CO₂ for subsequent conversion. Molten oxides would be a suitable option in this case. For flue gas, aqueous amine and carbonate solutions can be effective for CO₂ capture, but careful attention must be given to selecting catalysts resistant to gas impurities. The decision to extract CO₂ from flue gas or ambient air depends on various factors,



including location, emission sources, available resources, and technological readiness. Additionally, new catalysts that demonstrate long-term stability for extended electrolysis periods are needed, or additives and specialized functionalities should be introduced to the electrode or electrolyte to prevent catalyst deactivation. Ideally, the electrochemical conversion process should regenerate the capture medium with minimal loss or degradation. Ultimately, the full integration of the capture and conversion processes must operate stably for hundreds or even thousands of hours without a significant decline in efficiency for both CO₂ capture and conversion.

One promising approach is integrating the capture and conversion sites at a molecular level. This approach closely mimics the working mechanism of enzymes, creating a precise microenvironment for molecular recognition and specific catalysis.²⁰² These studies highlight the strategy of tailoring microenvironments at a molecular level to enhance the performance of CO₂ capture and conversion.

(4) The scalability and economics of integrated systems have not yet been thoroughly explored. The few existing studies on such systems remain on the laboratory scale. In contrast, other large-scale CO₂ technologies have reached pre-commercial stages, including high-temperature electrochemical CO₂-to-CO production,²⁰³ thermal catalytic hydrogenation of CO₂ to methanol,²⁰⁴ and biological CO₂ upgrading to methane.²⁰⁵ These technologies can serve as valuable reference points for evaluating the scalability of integrated CO₂ capture and electrochemical conversion.

Any technology nearing scalability requires a comprehensive techno-economic analysis (TEA), which should extend beyond energy cost analysis. This means factoring in operational expenditures (OPEX) and capital expenditures (CAPEX) to fully understand the economic viability of the process.²⁰⁶ This assessment should encompass the entire value chain, including CO₂ capture methods (post-combustion or DAC), electrolysis using various capture media, and downstream processes for product purification and recycling of unconverted CO₂. While a detailed TEA is beyond the scope of this review, it would be highly beneficial for experts in the field to explore these aspects further. An important direction would be to conduct a comparative TEA of the integrated and independent routes over the entire life cycle of the CO₂ capture and conversion system.

To scale up the electrochemical conversion of captured CO₂ from a technological point of view, several parameters, such as current density, faradaic efficiency, and electrode area, must be optimized to increase product formation rates. However, once a target production rate is defined, a balance must be struck between current density and electrode area, as these factors heavily influence the economic feasibility of the electrolysis process. For instance, operating at lower current densities tends to be more energy efficient and reduces OPEX by minimizing energy losses from internal resistance. However, this approach requires larger electrolyzers, driving up CAPEX. Conversely, higher current densities might lower CAPEX but increase OPEX due to less efficient energy usage. In any case,

highly durable electrolyzer systems are essential to reduce overall costs, as frequent replacement of electrolyzer components incurs additional expenses and should, therefore, be minimized.

6.3. Future directions

In addition to addressing the challenges mentioned, there are numerous other opportunities to advance this emerging field. Two potential future directions are outlined below:

6.3.1. Exploring new capture agents for reactive CO₂ capture. So far, electrochemical conversion of captured CO₂ has primarily focused on carbamate, bicarbonate, and carbonate formed using conventional amine, carbonate, and molten oxide solutions as capture agents. Recently, various redox-active organic mediators, such as quinones and bipyridine-based types, have shown considerable promise for CO₂ capture through electrochemical swing.²⁰⁷ In this process, the redox-active mediator is first electrochemically reduced, triggering the CO₂-binding step. The resulting CO₂-adduct can then be further reduced at the electrode, leading to the production of various compounds and regeneration of the organic carrier. Notably, these redox-active mediators, as well as conventional amines, can be tethered to the active electrode surface, offering a novel approach for CO₂ capture and conversion.²⁰⁸ In this configuration, the captured CO₂ is located close to the electrode rather than in the electrolyte solution, reducing mass transport limitations and potentially increasing conversion efficiency. The use of ionic liquids also represents a promising pathway due to their strong affinity for CO₂ while still activating CO₂ conversion.²⁰⁹

6.3.2. Reactive CO₂ capture from the ocean. The ocean is a major carbon reservoir, with a CO₂ concentration ~50 times higher than in the air. As described in Section 5.1.3, a reported coupled system successfully captured and converted CO₂ from ocean water into valuable products such as CO and C₂H₄. Although not fully integrated, this highlights the immense potential of such an approach. The next step would be to fully integrate the ocean capture and conversion processes to reduce energy consumption by eliminating the CO₂ release step. However, several challenges must be addressed for successful implementation, such as optimizing electrochemical processes, developing corrosion-resistant catalysts, and addressing ecological impacts on marine life and ocean chemistry.

Author contributions

All authors contributed to the manuscript's writing, reviewing, and revising. K. Daasbjerg coordinated and supervised the overall project.

Data availability

All data supporting the findings in this review are available from the referenced sources listed in the manuscript. The data presented in the tables are derived from the corresponding



literature references. For the calculated values in Table 2, details of the calculation methodology are provided in the ESI.†

Conflicts of interest

There are no conflicts to declare.

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