UC Berkeley

UC Berkeley Previously Published Works

Title

Bipolar membrane electrolyzers enable high single-pass CO2 electroreduction to multicarbon products.

Permalink

https://escholarship.org/uc/item/260210xz

Journal

Nature Communications, 13(1)

Authors

Xie, Ke Miao, Rui Ozden, Adnan et al.

Publication Date

2022-06-24

DOI

10.1038/s41467-022-31295-3

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at https://creativecommons.org/licenses/by/4.0/

Peer reviewed



ARTICLE



1

https://doi.org/10.1038/s41467-022-31295-3

OPFN

Bipolar membrane electrolyzers enable high singlepass CO₂ electroreduction to multicarbon products

Ke Xie^{1,5}, Rui Kai Miao ^{2,5}, Adnan Ozden², Shijie Liu ², Zhu Chen¹, Cao-Thang Dinh³, Jianan Erick Huang¹, Qiucheng Xu⁴, Christine M. Gabardo ², Geonhui Lee ¹, Jonathan P. Edwards ², Colin P. O'Brien ², Shannon W. Boettcher ⁴, David Sinton ² & Edward H. Sargent ¹

In alkaline and neutral MEA CO_2 electrolyzers, CO_2 rapidly converts to (bi)carbonate, imposing a significant energy penalty arising from separating CO_2 from the anode gas outlets. Here we report a CO_2 electrolyzer uses a bipolar membrane (BPM) to convert (bi)carbonate back to CO_2 , preventing crossover; and that surpasses the single-pass utilization (SPU) limit (25% for multi-carbon products, C_{2+}) suffered by previous neutral-media electrolyzers. We employ a stationary unbuffered catholyte layer between BPM and cathode to promote C_{2+} products while ensuring that (bi)carbonate is converted back, in situ, to CO_2 near the cathode. We develop a model that enables the design of the catholyte layer, finding that limiting the diffusion path length of reverted CO_2 to ~10 μ m balances the CO_2 diffusion flux with the regeneration rate. We report a single-pass CO_2 utilization of 78%, which lowers the energy associated with downstream separation of CO_2 by CO_2 to compared with past systems.

¹ Department of Electrical and Computer Engineering, University of Toronto, 10 King's College Road, Toronto, ON M5S 3G4, Canada. ² Department of Mechanical and Industrial Engineering, University of Toronto, 5 King's College Road, Toronto, ON M5S 3G8, Canada. ³ Department of Chemical Engineering, Queen's University, 19 Division Street, Kingston, Kingston, ON K7L 3N6, Canada. ⁴ Department of Chemistry and Biochemistry, University of Oregon, Eugene, OR 97403, USA. ⁵These authors contributed equally: Ke Xie, Rui Kai Miao. [©]email: sinton@mie.utoronto.ca; ted.sargent@utoronto.ca

 O_2RR for C_{2+} production requires the simultaneous achievement of high production rate and high energy efficiency^{1,2}. The current densities in flow cells (A in Table 1) and membrane-electrode assemblies (MEAs, B in Table 1) have reached industrially relevant levels (ethylene partial current density > 100 mA cm^{-2})³; however, the energy penalty associated with low single-pass CO_2 utilization (SPU:

the fraction of the CO_2 converted to the total input CO_2) has yet to be reduced to practical levels (SPU > 40%)⁴. Carbonate formation and crossover in typical CO_2RR electrolyzers limit the SPU to $\leq 25\%$ for C_{2+} (details in SI1), imposing energy penalties of 280-480 GJ in alkaline-media, and of 80-130 GJ in neutral-media, for the production of each ton of ethylene^{5,6}.

	Configuration diagrams	Cathode products	CO ₂ -originated carbonate mass balance	Catholyte	Cathode micro- environment	Max. total SPU for Cu cathode	Ref.
A	CO, Flow-cell Co. Crossover CO. H. etc. Co	C ₁ and C ₂₊	Loss in catholyte; migrate through AEM and reverted to ${\rm CO_2}$ at anode	OH [−] , HCO ₃ [−] , SO ₄ ^{2−}	Locally strong alkaline, with bulk alkaline or neutral catholyte	24%	21,41
В	AEM-based MEA CO,	C_1 and C_{2+}	Migrate through AEM and revert to CO_2 at anode	Solid-state polymer (AEM)	Locally strong alkaline	30%	21
С	Weak-acid modified BPMEA H,O CO, H, Hadden and the control of the	C ₁	Revert to CO ₂ at the surface of CEL	Solid-state polymer (PAA-PAH bilayers)	Locally weak acidic	N/A	42
D	High concentration analyte BPMEA HO OH HO	C ₁	Revert to CO ₂ at the surface of CEL	Solid-state polymer (CEL of BPM)	Locally strong acidic, cation effect	N/A	10
E	Buffer stationary catholyte BPMEA H,O O, HO, HO, HO H	C ₁ and C ₂₊ (C ₂₊ shown in SI3)	Revert to CO ₂ at the surface of CEL	>600 μm thick NaHCO ₃	Locally strong alkaline, with bulk neutral catholyte	15%	12
F	Non-buffer sub-100 jam-bhiok stationary catholyte BPMEA (this work) O, CO, IT OLD THE CO, THE	C_1 and C_{2+}	Reverted to CO_2 near (~12 μ m) the cathode	65 μm thick $K_2 SO_4$	Locally strong alkaline, with bulk neutral catholyte	78%	This work

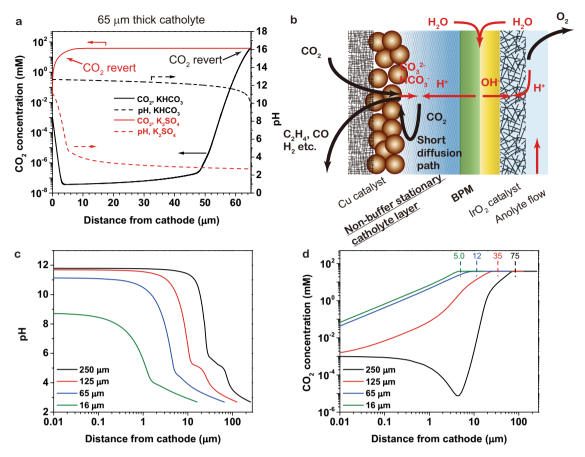


Fig. 1 The configuration and simulation of the electrolyzer design. a The CO_2 (solid lines) and pH distributions (dashed lines) in the 65 μ m-thick SC-layer. The positions where the (bi)carbonates revert to CO_2 are marked (red for non-buffering and black for buffering electrolyte). **b** The schemes and the mass transfer in the SC-BPMEA. **c** The pH distribution inside the SC-layer. **d** The dissolved CO_2 concentration profile inside the SC-layer. The virtual boundaries marked by dash lines are defined as the position where the CO_2 concentration becomes 1% lower than the bulk concentration. The numbers marked above are the distances between the virtual boundaries and the cathode surface.

Analysis of the CO_2 and carbonate mass balance in neutral-media electrolyzers (SI1) indicates that achieving high SPU requires that (bi)carbonate not cross the membrane to the anode, and that (bi) carbonate formed at the cathode must revert to CO_2 and remain available to participate in CO_2RR by returning to the cathode.

Bipolar membranes (BPMs) have been used to block $\rm CO_2$ crossover and convert (bi)carbonate back to $\rm CO_2^{7-9}$. In $\rm CO_2RR$ electrolyzers, commercial BPMs inhibit $\rm CO_2$ loss 10 , but the acidic cation-exchange layer (CEL) degrades the cathode's $\rm CO_2RR$ selectivity 10 .

To address cathode acidification, one may use a buffering catholyte (e.g., KHCO₃) between CEL and cathode^{9,11,12}; this approach (Fig. 1a) provides a pH at the CEL surface of ~3, and keeps the cathodic local pH >12. However, in a buffering catholyte, (bi)carbonate reverts to CO_2 near the surface of the CEL (solid black line in Fig. 1a), slowing mass transfer of the reverted CO_2 and reducing reactant availability for CO_2RR . This leads to the best SPUs reported of ~15% in prior BPM-based electrolyzers in C_{2+} electroproduction (SI3 and SI5). When the catholyte is flowing, the SPU is even lower, ~6% according to previous reports, because flowing catholyte removes reverted CO_2^{11} . Prior BPM-based electrolyzers (Table 1) have, as a result, not exceeded the 25% SPU limitation for C_{2+} electroproduction (SI3).

We show that a stationary, non-buffering catholyte with well-designed catholyte layer thickness (Fig. 1b) addresses these mass-transfer limitations while simultaneously providing the needed high local cathode pH (Fig. 1a). BPM-generated protons migrate farther in non-buffering catholyte than in buffering catholyte, since the latter

consumes protons at a higher rate. Here we show further quantitative analysis of the design requirements to achieve this target.

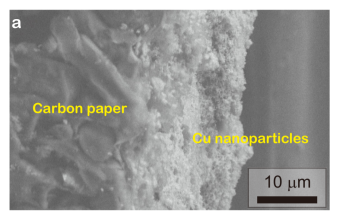
Results

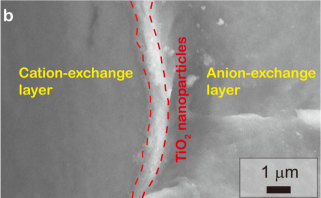
Finite-element numerical simulations of the stationary catholyte (SC)-layer. The composition and thickness of the catholyte layer influence the local pH, the efficiency of CO₂ regeneration and, thereby, the overall cell performance. We applied a one-dimensional multiphysics model in COMSOL to investigate the catholyte layer in BPM-based CO₂RR electrolyzers.

The CO_2 reactant is provided by two sources: the inlet CO_2 flow (gas) and the regenerated CO_2 (dissolved form, aq.) in the catholyte. To achieve high SPU, it is necessary to restrict the gaseous CO_2 feed^{13,14}. Under a restricted gaseous CO_2 availability, the cathode CO_2 supply relies more on regeneration (SI1): in an ideal case with 100% SPU and 100% C_{2+} selectivity, regeneration contributes 75% of the consumed CO_2 . Thus, the mass transport of regenerated CO_2 is most critical, and that transport is governed by catholyte composition and thickness.

At steady-state, electrolysis creates a pH gradient through the catholyte layer: the pH is high near the cathode and low near the CEL. The protons and (bi)carbonate ions recombine in the catholyte, forming CO₂ (aq.) that diffuses, in response to a concentration gradient, to the Cu catalyst.

Simulations resolve the local cathode environment as a function of dimensions, electrolyte, and running conditions (Figs. 1a, c, d, SI4





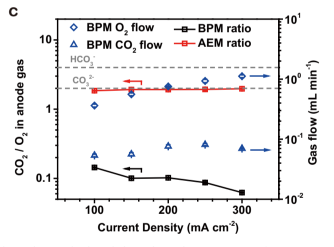


Fig. 2 Characterization of electrodes and BPM. Scanning electron microscopy (SEM) images of the cathode electrode (**a**) and the CEL/AEL interface of the custom BPM used for SC-BPMEA in neutral 0.1 M KHCO₃ anolyte (**b**). The custom BPM consists of a NafionTM 212 as CEL, Piperion (Versogen) as AEL, and a TiO₂ nanoparticle layer sandwiched in between as the water dissociation catalyst. **c** The CO₂/O₂ ratio in the anode gas stream for the conventional electrolyzers (red squares) and our SC-BPMEA (black squares) at various current densities. O₂ and CO₂ flow rates in the SC-BPMEA are also indicated. The plots show data obtained after 1h of continuous electrolysis at each current density. The data are from an SC-BPMEA with a 125 μm-thick SC-layer. Similar data were collected for 65 and 250 μm layers (difference below 5%). The error bars represent the standard deviation of three measurements.

and SI5). We selected 250, 125, 65, and 16 μm as the modelled thicknesses to correspond to commercially available materials.

Use of a buffering catholyte (e.g., KHCO₃) leads to a thick CO₂ (aq.) diffusion layer close to the catholyte thickness, since the CO₂

(aq.) is generated near the CEL surface, as shown in Fig. 1a (details in SI5). This effect reduces the CO_2 (aq.) mass-transfer efficiency. The experimental results presented and discussed in SI6 show similar trends to the simulation results.

In contrast, Fig. 1a, c illustrate that with a non-buffering catholyte layer (e.g., $0.5 \, \text{M} \, \text{K}_2 \text{SO}_4$) with thicknesses of 250, 125, and 65 μm , the local pH values near the cathode are greater than 11, which is sufficient to promote selectivity towards $\text{CO}_2 \text{RR}$ over HER¹³. Reducing the SC-layer thickness to 16 μm results in a cathode pH of 8.7, implying a lower selectivity toward $\text{CO}_2 \text{RR}$.

Figure 1c, d shows the simulated concentration profiles of CO₂ (aq.) in the non-buffering SC-layer. At steady-state, the CO₂ (aq.) is continuously supplied to the cathode to participate in CO₂RR, forming a concentration gradient (the boundary was defined here as the position where CO₂ concentration is 1% lower than the saturated concentration) to the cathode surface. Prior studies have termed the zone between the cathode and this boundary the diffusion layer 15. The thickness of the diffusion layer controls the efficiency of CO₂ (aq.) mass transport¹⁵. According to the simulations, the thicknesses of the diffusion layers are 75, 35, 12, and 5 µm for the catholyte layers with the thicknesses of 250, 125, 65, and 16 µm, respectively (marked in Fig. 1d). For reference, the CO₂ (aq.) diffusion layer thickness in H-cells (all CO₂ supplied in dissolved form) is typically 40-100 µm, and this does not support current densities exceeding 100 mA cm⁻².16 We expect that diffusion layers <40 µm, and a corresponding catholyte thickness <150 µm, are required for sufficient mass transport in a nonbuffering catholyte. To achieve similar mass transport in a buffering catholyte, the total thickness could not exceed 12 um, and the cathodic pH would not be sufficiently alkaline for selective CO₂RR (SI5).

The simulation results suggest the following design principles for the catholyte layer in a BPM-based electrolyzer: the local cathode pH and the diffusion layer thickness of the regenerated CO₂ increase as the catholyte thickness increases; the buffering capacity of the catholyte increases the diffusion layer thickness and reduces transport. Precise control of the thickness of a non-buffering catholyte should thus offer a route to high SPU, CO₂RR selectivity, and reaction rate.

System design for high SPU of CO₂ feedstock. Guided by the above analysis, we focused on a *stationary catholyte bipolar membrane electrode assembly* (SC-BPMEA) electrolyzer and incorporated a judiciously-designed catholyte layer and BPM (Fig. 1a).

The cathode was prepared by spraying Cu nanoparticles onto a hydrophobic carbon gas-diffusion layer for CO₂RR (Fig. 2a). The anode was IrO₂ supported on Ti felt for the oxygen evolution reaction (OER). A BPM (SEM in Fig. 2b) under reverse bias was employed with the anion exchange layer (AEL) contacting the anode and the CEL contacting the SC-layer (porous support saturated with electrolyte). The cathode was compressed onto the porous layer, and the anode and cathode flow-field plates sandwiched the system.

The BPM employed in this work sandwiched TiO_2 nanoparticles as the water dissociation catalyst¹⁷. This custom BPM can lower the cell voltage by ~1 V compared with commercial BPMs (e.g., Fumasep, details in SI2). The full cell voltage of such custom BPM-based electrolyzers is close to that of anion exchange membrane (AEM)-based systems.

Measurements of the $\rm CO_2/O_2$ ratio in the anode gas stream show that the SC-BPMEA effectively prevents $\rm CO_2$ crossover, as required for high SPU (SI1)^{8,11}. In agreement with the previous studies^{8,11}, the AEM-based MEA (AEMEA) showed an anode $\rm CO_2/O_2$ ratio of ~2 for current densities ranging from 100 to

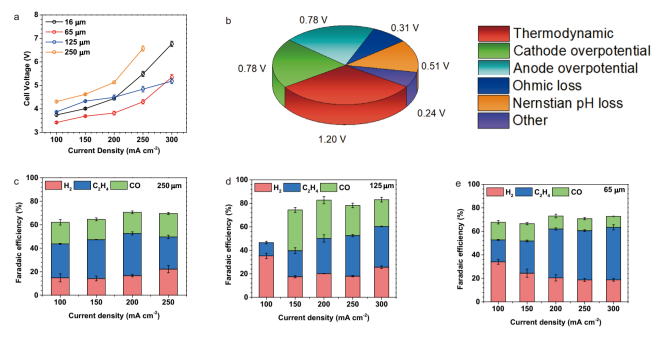


Fig. 3 Investigations on the CO₂RR performance of the SC-BPMEA using catholyte thicknesses of 250, 125, 65, and 16 μm. All the results are collected at 35 °C with a CO₂ flow rate of 10 sccm cm⁻² (normalized by the geometric area of the cathode), a catholyte of 0.5 M K₂SO₄, and an anolyte of 0.1 M KHCO₃. **a** The dependence of cell voltages on current density. **b** Distribution of voltage losses measured in the SC-BPMEA with 65 μm thick catholyte operating at 200 mA cm⁻² (cell voltage = 3.82 V). The breakdown of the voltage is explained in SI7 of the Supplementary Information. The dependence of the CO₂RR gas products FE on the current density for the SC-BPMEAs with the catholyte thickness of 250 (**c**), 125 (**d**), and 65 μm (**e**). The FE result for 16 μm is shown in Supplementary Fig. 12. The error bars represent the standard deviation of three measurements.

 $300~\text{mA}~\text{cm}^{-2}$ (Fig. 2c). In conventional AEMEAs, the anionic charge carriers are $\text{CO}_3{}^2-$, and thus suffer the loss of one molecule of CO_2 for every two electrons transferred. The anode CO_2/O_2 ratio in the SC-BPMEA (0.06 at $200~\text{mA}~\text{cm}^{-2}$) is one order of magnitude lower. Control experiments confirm that the CO_2 detected in the anode is not due to acidification of anolyte (using 0.1 M K_2SO_4 instead of 0.1 M KHCO $_3$ resulted in a similar CO_2/O_2 ratio, Supplementary Fig. 3). The anode CO_2/O_2 ratio decreases as the operating current density increases, an effect we ascribe to an increased flux of protons toward the cathode. This flux decreases the pH at the CEL surface and reduces the diffusion of CO_2 and $\text{HCO}_3{}^-/\text{CO}_3{}^-$ in the $\text{CEL}^{18,19}$.

Impact of the thickness of the SC-layer on CO₂RR. As predicted from simulations, the thickness of the stationary catholyte has a major impact on cell voltage. The cell voltage of the SC-BPMEA decreases as the thickness of the SC-layer decreases (Fig. 3a) from 250 μm (5.1 V, 200 mA cm⁻²) to a minimum at 65 μm (3.8 V, 200 mA cm⁻²). Further thinning the catholyte to 16 μm resulted in higher voltage (4.4 V, 200 mA cm⁻²)—an effect of the lowerporosity support layer used in the 16 µm case (<20% vs. >70% for the thicker layers, see Supplementary Fig. 13a and SI7). A longer ion migration path and higher ohmic resistance partially explain the 0.67 V cell voltage increase as the stationary catholyte thickness increases from 65 to 125 µm. Based on the independently measured ohmic resistance (Supplementary Fig. 13a), increasing the SC-layer thickness from 65 to 125 µm imposes an ohmic voltage increase of merely $0.07~\mathrm{V}$ at $200~\mathrm{mA~cm^{-2}}$. Similarly, compared to $65~\mu m$, the 250 µm SC-layer increases the ohmic voltage loss by 0.24 V at 200 mA cm^{-2} , while the cell voltage increases by 1.3 V.

The simulations (Fig. 1d) indicate that the thicker SC-layer results in longer transport distances for dissolved CO_2 . The CO_2 regeneration rate inside the SC-layer also depends on the current density, and for thicker SC-layers (e.g., >125 μ m), CO_2 bubbles are more prone to form near the CEL. These bubbles obstruct ion

migration, increasing the ohmic resistance of the SC-BPMEA. Electrochemical impedance spectroscopy measurements (Supplementary Fig. 13d, e) also support this finding. An applied current of 200 mA cm $^{-2}$ resulted in an insignificant change to the high-frequency resistance (HFR) of the SC-BPMEA with a 65 μm -thick SC-layer; while, in contrast, the HFR of the SC-BPMEA with a 125 μm -thick SC-layer increased by 120% after applying 200 mA cm $^{-2}$ for 20 min, leading to a cell voltage 0.6 V higher than for the 65 μm SC-layer.

The cell voltage of the SC-BPMEA with a 65 μ m SC-layer operating at 200 mA cm⁻² is 3.8 V, comparable to the AEM-based neutral-media MEAs operating at similar conditions (difference <±0.05 V)^{20–22}. This result demonstrates that the cell voltage of a BPM-based CO₂RR electrolyzer can be as low as that of an AEM-based electrolyzer with a current density of up to 200 mA cm⁻², while suppressing unwanted crossover and providing high SPU.

Figure 3b shows the breakdown of the 3.8 V cell voltage, determined using methods reported previously^{13,21} (SI7). The factors making up the cell voltage include the thermodynamic potential, cathode overpotential, anode overpotential, ohmic loss, and Nernstian/concentration overpotential (i.e., due to pH gradient)²¹. The sum of these factors accounts for most of the cell voltage, suggesting that the water dissociation overpotential at the AEL/CEL interface of the BPM is small in the SC-BPMEA at 200 mA cm⁻², in agreement with previous reports employing BPMs fabricated in this way (SI2 and ref. ¹⁷).

The thickness of the SC-layer also affects selectivity towards CO_2RR . With thicknesses of 65, 125, and 250 μm , the H_2 Faradaic efficiencies (FEs) are consistent (~20% at 200 mA cm⁻², Fig. 3c-e), confirming that high local pH conditions are maintained the cathode in these cases (Fig. 1c). However, reducing the thickness to 16 μm increases the H_2 FE to 88% at 200 mA cm⁻² (Supplementary Fig. 14), consistent with a cathodic pH that is reduced due to fast proton transport through a thin

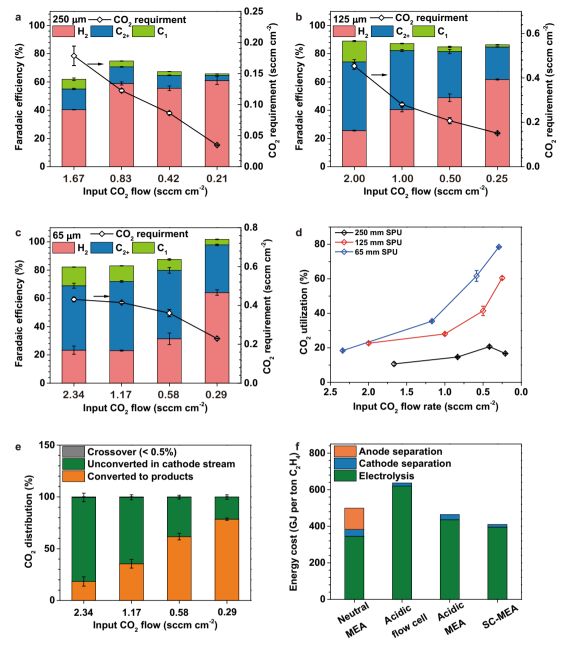


Fig. 4 The exploration of the CO_2RR performance and energy intensity of SC-BPMEA with restricted reactant availability. All the measurements were conducted at 35 °C and 200 mA cm⁻², and the data were collected after 2 h of continuous operation. **a-c** The FE distributions and the CO_2 requirements (total CO_2 converted to products) of the SC-BPMEAs with different catholyte thickness and input CO_2 flow rates (sccm normalized by electrode area). C_1 refers to CO_2 , formate, and methane. C_{2+} refers to ethylene, ethanol, acetate, and n-propanol. **d** The total CO_2 single-pass utilization (the CO_2 -to-ethylene single-pass conversion see Supplementary Fig. 13a) for the SC-BPMEAs with different catholyte thickness and input CO_2 flow rates. **e** Carbon balance in SC-BPMEA with 65 μm CO_2 at different input CO_2 flow rates. See Supplementary Fig. 13b for plots on a logarithmic scale. **f** The comparison of the energy cost distributions among state-of-art CO_2 -to-ethylene electrolyzers. The SC-BPMEA case refers to the input CO_2 flow rate of 1.17 sccm cm⁻². The error bars represent the standard deviation of three measurements.

SC-layer. Without restricting CO_2 availability (the performance in Fig. 3 was recorded at a CO_2 flow rate of 10 sccm cm⁻²), the SC-BPMEAs with the SC-layer thickness of 65, 125, and 250 μ m show similar ethylene FE of 35–43%.

Assessment of SPU in SC-BPMEA. By suppressing the crossover of CO_2 (e.g., <0.5% of total CO_2 input at 200 mA cm⁻², Figs. 2c and 4e), the SC-BPMEA surpasses the SPU of conventional CO_2 -to- C_{2+} electrolyzers, in which carbonate is the dominant charge

carrier. Measuring the CO_2 SPUs with a restricted CO_2 flow rate is a direct approach to determining the upper bound of SPU in the CO_2RR electrolyzers¹³.

As the inlet CO_2 flow rate decreased, the C_{2+} FE of the SC-BPMEA at 200 mA cm⁻² decreased, accompanied by an increase in the H_2 FE (Fig. 4a-c). With SC-layer thicknesses of 65 μ m (Fig. 4c), as the input CO_2 flow rate decreases from 1.17 to 0.58 and 0.29 sccm cm⁻², the C_{2+} FE decreases from 49 to 48% and 34%, while the H_2 FE increases from 23 to 31% and 64%. This shift is consistent with a CO_2 mass transport limitation CO_2 13, 20,21.

The stationary catholyte thickness affects the SPU of the SC-BPMEA. The SPU gradually increases up to 21, 61, and 78% for the SC-BPMEAs with SC-layer thicknesses of 250, 125, and 65 μ m, respectively (Fig. 4d). These results demonstrate that high CO₂ conversion efficiencies are possible using SC-BPMEAs with SC-layer thicknesses of 125 and 65 μ m.

For a given CO_2 flow rate, a thicker SC-layer produces a lower SPU (Fig. 4d). In the SC-BPMEA, reactant CO_2 is available from the inlet gas stream and regeneration in the SC-layer. With unrestricted CO_2 supply (Fig. 3c–e), the H_2 FEs are similar for different stationary cathode layer thicknesses, indicating that both the CO_2 availability and local pH are unaffected by catholyte thickness under excess supply conditions. The simulations suggest that the thicker SC-layer results in a lower dissolved CO_2 flux to the cathode due to the smaller concentration gradient (Fig. 1d). Compared to the SC-BPMEAs with thinner SC-layers, CO_2 availability with thicker SC-layers decreases more significantly with reducing CO_2 flow rate, leading to a more dramatic increase in H_2 FE (Fig. 4a–c).

The experimental trends are generally consistent with those of the simulations. The SC-BPMEA with a dissolved CO₂ diffusion layer thicker than 75 μm (representing a 250 μm SC-layer) fails to surpass the SPU limit because of insufficient mass transfer. In contrast, a 65 μm SC-layer facilitates efficient mass transport of the regenerated CO₂ (diffusion layer thickness of 12 μm) and simultaneously promotes high local cathode pH.

As demonstrated in SI10, SC-BPMEAs using acidic and alkaline electrolytes achieve carbon efficiencies comparable to those using neutral electrolytes. The compatibility of SC-BPMEAs with a range of electrolytes offers flexibility in the selection of cathode and anode catalysts. In contrast, acidic $\rm CO_2$ -to- $\rm C_{2+}$ electrolyzers have only been demonstrated with precious metal anodes 13,14 .

As shown in Supplementary Fig. 18, the SC-BPMEA shows > 50-h stability operating at 200 mA cm $^{-2}$ with limited $\rm CO_2$ availability (CO $_2$ input flow rate of 1.42 sccm cm $^{-2}$). This operating stability is competitive with that of the neutral-electrolyte-based $\rm CO_2$ -to-C $_2$ + electrolyzers 23,24 .

Can a cation-exchange membrane replace the BPM in SC-BPMEA? We attempted to extend the SC-layer strategy in a CEM-based MEA cell (i.e., SC-CEMEA, Fig. 5a) using an acidic anolyte with pH < 2.4, expecting a lower cell voltage than the SC-BPMEA while maintaining high SPU. We found that in the SC-CEMEA, the CO₂ crossover was essentially eliminated. This observation is ascribed to the lower pH near the stationary catholyte layer/CEM interface, as shown in Fig. 5a.

SC-CEMEA shows a lower full cell voltage (Fig. 5b) compared to the SC-BPMEA presented, partly due to the lower resistance of the CEM and the absence of water dissociation overpotential. Meanwhile, it has a reasonable CO_2RR selectivity over HER (Fig. 5c) due to the cation effect and high local pH induced by the presence K⁺ in the SC-layer (Fig. 5a). However, this design is not amenable to steady-state operation without continuous addition of acid and salt to the anolyte, as the initial pH gradient will be eliminated due to co-ion transport and neutralization. We found the CO_2RR selectivity decreases over time and approaches 100% H_2 after ~ 3 h.

We also observed that the SC-CEMEA design periodically ejects electrolyte from the cathode flow channel, likely due to poor water balance. On the anode, the OER generates one proton per one electron transfer. The charge carriers across the CEM are primarily $\mathrm{H^+}$, although neutral ion pairs will diffuse as well. At the cathode $\mathrm{K^+}$ makes up the electrochemical double layer at the Cu surface, and the steady-state $\mathrm{K^+}$ profiles are governed by the

electric and chemical-potential gradients that develop under operation, which usually takes tens of seconds¹⁹. H⁺ migrates to the cathode and combines with OH⁻ (or CO₃²⁻/HCO₃⁻), producing water at the cathode. The protons also drag water molecules (~1 per proton)²⁵ by electro-osmosis. We accordingly calculate the water balance for different cathode products as listed in Table 2. The water generated and transported to the cathode appears to dilute and push out the electrolyte in the stationary catholyte layer, of which the volume is small (ca. 10 µL per cm² electrode area). This phenomenon results in flooding of the cathode (as confirmed experimentally) and loss of supporting electrolyte, thus degrading performance. In the BPMEA design, it is likely that the BPM slows co-ion transit across the membrane, compared to the CEM, by the large outward flux of OH⁻ and H⁺ from the water dissociating junction.

Energy assessment of the SC-BPMEA with optimal SC-layer.

The energy costs (measured in gigajoules per tonne of the target product, GJ/t) for a CO_2 -to- C_{2+} electrolyzer include the electrolysis electrical energy, cathodic stream separation, and anodic stream separation 13 . CO_2RR performance metrics of importance include cell voltage, target product FE, SPU, and CO_2 crossover 5 . High SPU and high energy efficiency have not been accomplished simultaneously in C_{2+} electroproduction. In SC-BPMEAs, a higher SPU reduces the energy required for cathode separation, but the accompanying decrease in the ethylene selectivity (Fig. 4c) elevates the specific energy requirement 5 . We carried out a total energy assessment of the SC-BPMEA and other state-of-art CO_2 -to-ethylene electrolyzers and summarized the results in Table 3 , Fig. 4 f, and Supplementary Table 2 .

The energy consumption of an alkaline CO₂RR electrolyzer² is listed in Supplementary Table 2. In such systems, CO₂ and OH⁻ react to form carbonate continuously. This carbonate has to be recovered to maintain the CO₂RR performance of such a system, consuming 5.5 GJ per tonne CO₂⁵. In the alkaline CO₂RR electrolyzers, ca. 63 tonne of CO₂ transforms to carbonate to produce 1 tonne of ethylene, representing an energy penalty of 350 GJ⁵. This costs at least \$1900 per tonne of ethylene, while its market price is \$800–1000 per tonne²⁶. The alkaline electrolyzers thus do not allow for ethylene electrochemical production to be yet profitable.

In neutral-media CO₂RR electrolyzers, recovering the CO₂ from the anodic gas stream results in significant energy costs. In the context of highly selective conversion (i.e., CO₂-to-ethylene with unity selectivity), the recovery process requires an energy input of 52 GJ to produce every tonne of product. In practice, due to non-unity product selectivity, the process is even more prohibitive, i.e., requiring an energy penalty of 80–130 GJ for producing one tonne of ethylene⁵.

As the SPU increases from 4 to 35%, we found a dramatic decrease in energy associated with cathode separation—from 85 to 15 GJ/t ethylene (Table 3), with the ethylene FE reduced by only 2%. Further increasing the SPU beyond 35% does not substantially reduce the energy cost associated with cathodic separation (Supplementary Table 2). This finding agrees with a recent energy analysis that in a (bi)carbonate-free $\rm CO_2$ -to- $\rm C_2$ + electrolyzer, improving SPU over 40% offers an insignificant benefit to the downstream separation cost⁴. Pursuing an SPU > 35% decreases ethylene FE by more than 4% when using the SC-BPMEA, and thus the increased input electricity cost exceeds the savings in the cathodic separation (Table 3 and Supplementary Table 2). Therefore, 35% SPU is the most favourable condition for the present SC-BPMEA.

The energy intensity of producing ethylene in SC-BPMEA is ~30% lower than that in conventional neutral-electrolyte-based

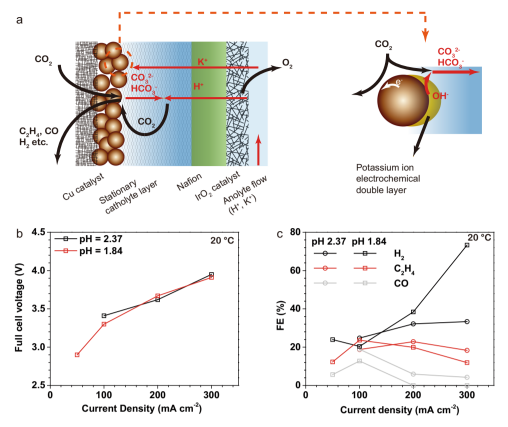


Fig. 5 Investigations on the CO₂RR performance of the SC-CEMEA. The stationary catholyte is $0.5 \, \text{M} \, \text{K}_2 \text{SO}_4$ and the anolyte is $0.5 \, \text{M} \, \text{K}_2 \text{SO}_4$ plus $0.1 \, (\text{pH} = 2.37)$ or $0.5 \, \text{M} \, (\text{pH} = 1.84) \, \text{H}_3 \text{PO}_4$. All the measurements were performed at $20 \, ^{\circ}\text{C}$ and a CO₂ flow rate of 10 sccm cm⁻². **a** The scheme of the SC-CEMEA. **b** Full cell voltages at various current densities. **c** The gas product FEs at various current densities.

Table 2 The cathode-anode water balance in an SC-CEMEA.								
Product (1 mol)	Water balance i	n the cathode (mol)	Water balance in the anode (mol)					
	Consumed	Generated	Dragged in	Net	Consumed and dragged out			
СО	1	2	2	+3	3			
HCOO-	1	1	1	+1	2			
CH₃COO−	5	7	7	+9	11			
C ₂ H ₄	8	12	12	+16	18			
C ₂ H ₅ OH	9	12	12	+15	18			
CH ₄	6	8	8	+10	12			
H ₂	2	2	2	+2	3			

 CO_2 electrolyzers (Fig. 4f and Table 3). In conventional neutral-electrolyte CO_2 -to-ethylene electrolyzers, the CO_2 crossover (at least 70%)²¹ costs 60–90 GJ per ton of ethylene to recover CO_2 from the anodic O_2 stream⁵. Notably, this energy penalty cannot readily be reduced, independent of optimizing catalysts and operating conditions (e.g., input CO_2 flow rates, reaction rates, operating temperature, and pressure)⁶. In contrast, crossover CO_2 in SC-BPMEA is < 0.5% of the total CO_2 input, minimizing the energy cost of anodic separation.

Recently, CO₂-to-ethylene conversion has been achieved in acidic electrolytes in both flow cell¹³ and MEA configurations¹⁴. These systems enabled CO₂ SPUs exceeding 75% and also mitigated the energy cost associated with anodic separation (Table 3). Owing to the strongly acidic environment, the flow cell enables an ethylene FE of 28% at a full-cell potential of 4.2 V. The acidic MEA used an anion-exchange ionomer coating on the catalyst layer to promote CO₂RR over HER. The modification of

the surface with the anion exchange ionomer resulted in a higher ohmic loss, and thus the cell required potentials of 3.8 and 4.4 V at 100 and 200 mA cm $^{-2}$, respectively. These devices thus eliminated the anodic $\rm CO_2/O_2$ separation energy but at the penalty of larger cell voltages and/or lower ethylene FEs. In contrast, SC-BPMEA shows a cell voltage of 3.8 V at 200 mA cm $^{-2}$ with an ethylene FE of 42%—voltages and selectivities comparable to the best conventional neutral-electrolyte $\rm CO_2$ -to-ethylene MEAs 21 . Compared to acidic systems, the energy intensity of the SC-BPMEA is 36% and 12% lower than acidic flow cell and acidic MEA, respectively (Fig. 4f and Table 3).

Discussion

We demonstrate a BPM-based CO_2 -to- C_{2+} MEA, with a judiciously-designed SC-layer between catalyst and BPM, that overcomes the (bi)carbonate-formation reactant loss issue without compromising performance. The composition and thickness

Table 3 Comparison of the energy intensity between various CO ₂ -to-ethylene electrolyzers and this work.							
Metrics	Neutral- MEA ^{a21}	Acidic flow cell ^{a13}	Acidic MEA ^{a14}	This work (10 sccm cm $^{-2}$)	This work (1.17 sccm cm $^{-2}$)		
Cell type	MEA	Flow cell	MEA	SC-BPMEA	SC-BPMEA		
Electrolyte	Neutral	Acidic	Acidic	Neutral	Neutral		
Full cell voltage (V)	3.75	4.20	3.80	3.82	3.82		
Ethylene FE (%)	45	28	36	42	40		
Current density (mA cm $^{-2}$)	150	1200	100	200	200		
Input CO ₂ flow rate (sccm cm ⁻²)	8	3	0.8	10	1.17		
Total CO ₂ SPU (%)	3	78	34	4.1	35		
CO ₂ -to-ethylene (%)	1.2	28	7.6	2.0	17		
Demonstrated stability (hours)	100	14	12	52			
Energy intensity (GJ per ton ethylene)							
Electrolyzer electricity	345	620	436	385	395		
Cathode separation	38	17	28	85	15		

^aThe energy intensities of reference CO₂-to-ethylene devices operating under the reported conditions are calculated, and those that provide the lowest energy intensity are presented in this table. ^bCrossover of CO₂ in the acidic flow cell, acidic MEA, and SC-BPMEA are each lower than 0.5% of input CO₂. Therefore, we assume the anodic separation energy to be 0. All the energy costs are normalized per ton of ethylene produced. The energy intensities of the SC-BPMEA operating at other CO₂ input flow rates are listed in Table S3 of Supplementary Information.

465

Op

470

of the SC-layer determine the CO₂RR performance and SPU via a strong influence on the local pH and the chemistry and transport of CO₂. The buffering capacity and the thickness of the SC-layer determine the efficiency of the regeneration, the transport, and the availability of reactant CO₂. These effects were predicted in simulations and supported by experiments. The SC-BPMEA design largely eliminates the energy penalty associated with the CO₂ loss in electrochemical CO₂ reduction.

116

499

Op

637

The performance of the SC-BPMEA might be further improved using, for example, ionic liquid or other organic salts as the catholyte, and by optimizing the porosity, structure, and hydrophobicity of the porous support layers. The CO₂RR performance of the SC-BPMEA might be improved with new cathodic catalysts, optimizing the loading and processing of the catalyst layer, and by implementing BPMs with further-lowered water-dissociation voltage loss. Broadly, the SC-BPMEA is a useful platform for evaluating CO₂RR catalysts operating with high CO₂ utilization. The strategy and findings presented here are also relevant to the electrochemical systems such as nitrate reduction and (bi)carbonate reduction, where controlling dissimilar microenvironments near each electrode is useful, and the exchange/transport of species (other than OH⁻ or H⁺) between cathode and anode is problematic.

Methods

Anode separation

Overall energy

Materials. Phosphoric acid (H₃PO₄, 85%), potassium sulfate (K₂SO₄, 99%), potassium bicarbonate (KHCO₃, 99.7%), potassium chloride (KCl, 99%), potassium hydroxide (KOH, 99.95%), copper nanoparticles (25 nm), NafionTM 1100 W (5 wt. % in a mixture of lower aliphatic alcohols and water) and isopropanol (IPA, 99%) were purchased from Sigma Aldrich and used as received. Titanium oxide as received. The porous supports were also purchased from Fisher Scientific and used as received. The porous supports were also purchased from Fisher Scientific: 125 μm PVDF (0.45 μm pore size), 65 μm PTFE (0.44 μm pore size) and 16 μm PC (0.4 μm pore size). NafionTM 212, NafionTM XL, Fumasep (FAS-PET-130) and titanium (Ti) felt were purchased from Fuel Cell Store. Iridium(IV) chloride hydrate (Premion*, 99.99%, metals basis, Ir 73% min) was purchased from Alfa Aesar. The water used in this study was 18 MΩ Milli-Q deionized- (DI-) water. Nafion membranes were activated through the following procedure: 1 h in 80 °C 1 M H₂SO₄—1 h in 80 °C H₂O₂—1 h in 1 M H₂SO₄—stored in DI-water. Fumasep was used as received and stored in 1 M KCl. Piperion (40 μm) was purchased from W7Energy and stored in 0.5 M KOH.

Fabrication of water dissociation catalyst layer of the custom bipolar membrane (BPM). The water dissociation catalyst layer was fabricated following a similar procedure in a previous report¹⁷. TiO₂ nanoparticles inks were prepared by sonicating the mixture of TiO₂, DI-water, and IPA with the weight ratio of 1: 833: 2833 for 30 min. TiO₂ nanoparticle ink was spray-coated onto a Nafion 212

membrane, of which the edges were sealed by Kapton tape. The exposed membrane dimension was 2.2 cm \times 2.2 cm. The nominal loading of $\rm TiO_2$ is 0.2 mg cm $^{-2}$. The $\rm TiO_2\text{-}coated\ Nafion^{TM}$ was immediately used for assembling electrolyzers once prepared.

Op

410

Electrode preparation. For the $\mathrm{CO_2RR}$, we prepared the gas diffusion electrodes (GDEs) by spray-depositing a catalyst ink dispersing 1 mg mL⁻¹ of Cu nanoparticles and 0.25 mg mL⁻¹ of NafionTM 1100 W in methanol onto a hydrophobic carbon paper. The mass loading of Cu NPs in the GDE was kept at 1.5 mg/cm². The GDEs were dried in the air overnight prior to experiments.

The OER electrode preparation procedure involves: etching the Ti felt in hydrochloric acid at 70 °C for 40 min; rinsing the etched Ti felt with DI water; immersing the Ti felt into an Ir(IV) chloride hydrate solution; drying and sintering the Ir-loaded Ti felt. The loading, drying, and sintering steps were repeated until a final Ir loading of 1.5 mg cm $^{-2}$ was achieved.

Assembly of the stationary catholyte membrane electrode assembly (SC-BPMEA). The MEA set (5 cm²) was purchased from Dioxide Materials. A cathode was cut into a 2.1 cm × 2.1 cm piece and placed onto the MEA cathode plate with a flow window with a dimension of 2.2 cm × 2.2 cm. The four edges of the cathode were sealed by Kapton tape, which also made the flow window fully covered. The exposed cathode area was measured every time before the electrochemical tests, in the range of 3.1 to 4.2 cm². Onto the cathode, a porous support layer (2 cm \times 2 cm with various thicknesses, 250 μm was stacking two 125 μm-thick PVDF) saturated with desirable electrolyte (sonicated in electrolyte for 15 min to degas) was carefully placed. This porous support layer serves as the 'stationary catholyte layer (SC layer).' The considerations of membrane selection can be found in SI2 and SI4 of the Supplementary Information. When using the custom BPM, a TiO2-coated Nafion membrane was placed onto the SC-layer with the TiO₂ layer facing up, then covered by a Piperion (5 cm \times 5 cm) membrane. When using Fumasep BPM, the membrane was placed with its cation-exchange layer (CEL) facing the cathode side. An IrO2 loaded Ti felt (2 cm × 2 cm) was placed onto the anion-exchange layer (AEL) of the BPM.

Scanning electron microscopy (SEM). Images of cathode and custom BPM were captured by an FEI Quanta FEG 250 environmental SEM.

Electrochemical measurements. Throughout all experiments, CO_2 flowed to the cathode side at 10 sccm cm $^{-2}$ unless otherwise specified, while the anode side was fed with neutral 0.1 M KHCO $_3$ at 10 mL/min by a peristaltic pump unless otherwise specified. The electrochemical measurements were performed with a potentiostat (Autolab PGSTAT204 with 10A booster). The cell voltages reported in this work are not iR corrected. The system was allowed to stabilize at the specific conditions for > 1000 s before recording the results. All the error bars represent standard deviations based on three measurements.

Product analysis. The CO₂RR gas products, oxygen, and CO₂ were analyzed by injecting the gas samples into a gas chromatograph (Perkin Elmer Clarus 590) coupled with a thermal conductivity detector and a flame ionization detector. The gas chromatograph was equipped with a Molecular Sieve 5A Capillary Column and a packed Carboxen-1000 Column with argon as the carrier gas. The volumetric gas

flow rates in and out of the cell were measured with a bubble column. The FE of a gas product is calculated as follows:

$$FE_i = x_i \times \frac{VP}{RT} \times \frac{n_i F}{I} \tag{1}$$

Where x_i is the volume fraction of the gas product i, V is the outlet gas flow rate in L s⁻¹, P is atmosphere pressure 101.325 kPa, R is the ideal gas constant 8.314 J mol⁻¹ K⁻¹, T is the room temperature in K, n_i is the number of electrons required to produce one molecule of product F is the Faraday Constant 96485 C mol⁻¹, and J is the total current in A.

The liquid products from the cathode side of the SC-BPMEA were collected using a cold trap cooled to 0 °C. The collected liquid was combined with anolyte (some crossover liquid product) for quantifying by the proton nuclear magnetic resonance spectroscopy ($^1\mathrm{H}$ NMR) on an Agilent DD2 500 spectrometer in $\mathrm{D}_2\mathrm{O}$ using water suppression mode and dimethyl sulfoxide (DMSO) as the internal standard. For each plot of liquid product quantification, fresh anolyte was used, and the duration of the collection was 30 min. The FE of a liquid product is calculated as follows:

$$FE_i = m_i \times \frac{n_i F}{Jt} \tag{2}$$

Where m_i is the quantity of the liquid product i in mole, t is the duration of product collection (1800 s).

The CO₂ SPU calculation is detailed in SI1 of Supplementary Information.

COMSOL one-dimensional modelling. The electrochemical reaction model was performed by COMSOL Multiphysics version 5.5. This simulation was built upon previous modelling work $^{27-30}$. The local pH and different species concentrations were simulated for different catholyte thicknesses (16, 65, 125, and 250 μm). Two different catholytes (K_2SO_4 and KHCO_3) were used in the simulation. All the chemical reactions between species were considered in this one-dimensional modelling. The simulation (Fig. 6) included a 50 μm thick gas diffusion layer (GDL), a 0.1 μm thick Cu cathode catalyst (CL), a catholyte region with various thicknesses indicated above, and a cation exchange layer (CEL) boundary.

Constant concentration (Dirichlet) boundary conditions were used. Specifically, a constant concentration 37.8 mM of CO_2 was assumed within the GDL layer, as this region is in direct contact with the input CO_2 flow and thus assumed to be at equilibrium with gas phase CO_2 over this region for the purposes of the simulation. The BPM was interpreted as a boundary with a constant species concentration (1 M H_3O^+ at the CEL surface)^{18,31}, because it was assumed to generate protons as the dominant ionic charge carrier at a constant rate under constant current density (200 mA cm⁻²).

A user-controlled mesh is employed in the COMSOL simulation. Edge type of mesh is used for GDL, CL, catholytes, respectively. Specifically, the mesh distribution is predefined with an interval of 500 nm for GDL and catholytes, and an interval of 5 nm for CL.

Five different electrode reactions were considered at the cathode catalyst layer in this simulation. Specifically, the hydrogen evolution reaction and CO_2 reduction reactions to CO_2 CP4, $\mathrm{C}_2\mathrm{H}_4$, and $\mathrm{C}_2\mathrm{H}_5\mathrm{OH}$ occurred at the cathode catalyst layer. In SC-BPMEA, the catalyst layer is immersed in a catholyte. Thus the simulation considers no gas-phase transport in the catalyst layer. The carbonate equilibrium reactions, corresponding catholyte buffer reactions, and a water dissociation reaction were considered in the catholyte region. The electrochemical reaction rates of the specific products were determined from experimental results. They are calculated based on the same manner as previous work 17 .

The electrochemical reactions at cathode catalyst layer:

$$2H_{2}O + 2e^{-} \rightarrow H_{2} + 2OH^{-}$$

$$CO_{2} + H_{2}O + 2e^{-} \rightarrow CO + 2OH^{-}$$

$$CO_{2} + H_{2}O + 8e^{-} \rightarrow CH_{4} + 8OH^{-}$$

$$2CO_{2} + 8H_{2}O + 12e^{-} \rightarrow C_{2}H_{4} + 12OH^{-}$$

$$2CO_{2} + 9H_{2}O + 12e^{-} \rightarrow C_{2}H_{5}OH + 12OH^{-}$$

The heterogenous electrochemical reaction rates are determined by the following equations:

$$r_i = \frac{I_i}{n_i F} * \frac{\varepsilon}{L_{\text{catalyst}}} \tag{3}$$

$$r_{\mathrm{CO_2}} = -\frac{I_{\mathrm{total}}}{F} \left(\frac{\mathrm{FE_{CO}}}{2} + \frac{\mathrm{FE_{CH4}}}{8} + \frac{\mathrm{FE_{C2H4}}}{12} + \frac{\mathrm{FE_{C2H5OH}}}{12} \right) * \frac{\varepsilon}{L_{\mathrm{catalyst}}} \tag{4}$$

$$r_{\rm OH^{-}} = \frac{I_{\rm total}}{F} * \frac{\varepsilon}{L_{\rm catalyst}}$$
 (5)

Where I_i represents the partial current density for CO, CH₄, C₂H₄, and C₂H₅OH occurred at the cathode catalyst layer, respectively. n_i represents the number of electrons transferred per mole reactant. F represents faraday's constant.

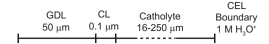


Fig. 6 The schematic of 1D COMSOL modelling. GDL and CL refer to gas diffusion layer and catalyst layer, respectively.

 I_{total} represents the total current density. The FEs for the specific product is determined by the experimental results shown in Fig. 3e. ε represents the catalyst porosity value. L_{catalyst} represents the cathode catalyst length.

The chemical reactions at the catholyte region and the corresponding forward k_f rate constants and reverse k_r rate constants taken from the literature³² (see Supplementary Table 3).

The Transport of Diluted Species physics model was used. The Nernst-Planck set of equations governed the species diffusion, and they were calculated in the same manner as previous work^{13,14}. Migration was ignored for simplicity as the experiments were performed in the concentrated electrolyte. The ion species transport is thus calculated by solving the two equations below.

$$\frac{\partial c_i}{\partial t} + \frac{\partial J_i}{\partial x} + r_i = R_i \tag{6}$$

$$J_i = -\frac{D_i \partial c_i}{\partial x} \tag{7}$$

$$D_i = \frac{\varepsilon_p}{\tau_{F,i}} * D_{F,i} \tag{8}$$

$$\tau_{F,i} = \varepsilon_p^{-1/3} \tag{9}$$

Where J_i is the molar flux, and r_i represents the heterogeneous electrode reactions for CO₂ reduction that were modelled at the cathode catalyst layer. R_i represents the rates of the homogeneous reactions indicated above. The Millington and Quirk model is used to determine the effective diffusivity, D_i . ε_p represents porosity coefficient. $\tau_{E,i}$ represents tortuosity coefficient.

The porosity value of 0.6 was used for the cathode catalyst and the porosity value of 1 for the catholyte region. The species diffusion coefficients are listed in Supplementary Table 4^{33-36} .

Henry's law and sets of Sechenov equation are applied to calculate the CO₂ concentration. The concentration of CO₂ in electrolytes depends on temperature and pressure. It is estimated in the same manner as previous work^{13,14}. The Sechenov coefficients are listed in Supplementary Table 5³⁷.

Energy assessment. We evaluated the energy consumptions for electrolyzer electricity, cathodic separation, and anodic separation in the context of ethylene. We consider the state-of-the-art CO₂RR systems from the literature, including alkaline flow-cell electrolyzers, neutral MEA electrolyzers, acidic flow-cells, and MEAs. This consideration is based on the performance metrics, including selectivity, productivity, and full-cell voltage—the combination reflects as energy intensity of producing multi-carbon products (i.e., ethylene). The proximity of these performance metrics will help refine the effect of anodic and cathodic separation on the energy requirement for producing ethylene. We summarize the input parameters to the model for all the systems. The energy assessment model, as well as the assumptions, are based on the previous work⁵. Ideally, it will be interesting to use experimental/modelling data corresponding to the exact gas composition from the CO2-to-C2+ device. However, at present, there is a gap in published literature. We, therefore, employed one of the most widely used models³⁸ (i.e., biogas upgrading) as the best approximation for evaluating the energy cost associated with cathode gas separation. The details of calculations for the carbon regeneration (for alkaline flow cell) and cathodic separation (for all the electrolyzers), can be found in previous work²⁶. The anodic separation (for neutral MEA electrolyzer) is modelled based on an alkaline capture solvent³⁹. The amount of CO2 crossover to the anode is calculated for one tonne of ethylene produced. The energy required to separate the CO2/O2 mixture is calculated based on a recent report by Carbon Engineering⁴⁰, in which 5.25 GJ/tonne CO₂ thermal energy and 77 kWh/tonne CO₂ are reported to be required to capture CO₂ and release at 1 bar. This energy consumption is a typical value for the alkaline capture process³⁹. For acidic flow-cell and MEA electrolyzers, we assume no energy cost associated with the anodic separation considering no CO₂ availability at the anodic gas stream¹³.

Data availability

All the data generated in this study are provided in the Supplementary Information and in the Source Data file. Source data are provided with this paper.

Received: 14 March 2022; Accepted: 10 June 2022; Published online: 24 June 2022

References

- De Luna, P. et al. What would it take for renewably powered electrosynthesis to displace petrochemical processes? Science 364, eaav3506 (2019).
- Dinh, C. T. et al. CO2 electroreduction to ethylene via hydroxide-mediated copper catalysis at an abrupt interface. Science 360, 783–787 (2018).
- Kibria, M. G. et al. Electrochemical CO₂ reduction into chemical feedstocks: From mechanistic electrocatalysis models to system design. Adv. Mater. 31, 1–24 (2019).
- Shin, H., Hansen, K. U. & Jiao, F. Techno-economic assessment of lowtemperature carbon dioxide electrolysis. Nat. Sustain 4, 911–919 (2021).
- Ozden, A. et al. Cascade CO₂ electroreduction enables efficient carbonate-free production of ethylene. *Joule* https://doi.org/10.1016/j.joule.2021.01.007 (2021).
- Rabinowitz, J. A. & Kanan, M. W. The future of low-temperature carbon dioxide electrolysis depends on solving one basic problem. *Nat. Commun.* 11, 10–12 (2020).
- Li, Y. C. et al. CO2 electroreduction from carbonate electrolyte. ACS Energy Lett. 4, 1427–1431 (2019).
- Ma, M., Kim, S., Chorkendorff, I. & Seger, B. Role of ion-selective membranes in the carbon balance for CO₂ electroreduction: Via gas diffusion electrode reactor designs. *Chem. Sci.* 11, 8854–8861 (2020).
- Vermaas, D. A. & Smith, W. A. Synergistic electrochemical CO₂ reduction and water oxidation with a bipolar membrane. ACS Energy Lett. 1, 1143–1148 (2016).
- Yang, K. et al. Cation-driven increases of CO₂ utilization in a bipolar membrane electrode assembly for CO₂ electrolysis. ACS Energy Lett. 6, 4291–4298 (2021).
- Ma, M., Clark, E. L., Chorkendorff, K. T. T. S. D. & Seger, B. Insights into the carbon balance for CO₂ electroreduction on Cu using gas diffusion electrode reactor designs. *Energy Environ. Sci.* 13, 977–985 (2020).
- Salvatore, D. A. et al. Electrolysis of gaseous CO₂ to CO in a flow cell with a bipolar membrane. ACS Energy Lett. 3, 149–154 (2018).
- Huang, J. E. et al. CO 2 electrolysis to multi-carbon products in strong acid. Science 372, 1074–1078 (2021).
- O'Brien, C. P. et al. Single pass CO₂ conversion exceeding 85% in the electrosynthesis of multicarbon products via local CO₂ regeneration. ACS Energy Lett. 6, 2952–2959 (2021).
- Xing, Z., Hu, L., Ripatti, D. S., Hu, X. & Feng, X. Enhancing carbon dioxide gas-diffusion electrolysis by creating a hydrophobic catalyst microenvironment. *Nat. Commun.* 12, 1–11 (2021).
- Clark, E. L. et al. Standards and protocols for data acquisition and reporting for studies of the electrochemical reduction of carbon dioxide. ACS Catal. 8, 6560–6570 (2018).
- Oener, S. Z., Foster, M. J. & Boettcher, S. W. Accelerating water dissociation in bipolar membranes and for electrocatalysis. *Science* 369, 1099–1103 (2020).
- Bui, J. C., Digdaya, I., Xiang, C., Bell, A. T. & Weber, A. Z. Understanding multi-ion transport mechanisms in bipolar membranes. ACS Appl. Mater. Interfaces 12, 52509–52526 (2020).
- Oener, S. Z., Ardo, S. & Boettcher, S. W. Ionic processes in water electrolysis: The role of ion-selective membranes. ACS Energy Lett. 2, 2625–2634 (2017).
- Ozden, A. et al. High-rate and efficient ethylene electrosynthesis using a catalyst/promoter/transport layer. ACS Energy Lett. 5, 2811–2818 (2020).
- Gabardo, C. M. et al. Continuous carbon dioxide electroreduction to concentrated multi-carbon products using a membrane electrode assembly. *Joule* 3, 2777–2791 (2019).
- Li, F. et al. Molecular tuning of CO₂-to-ethylene conversion. Nature 577, 509–513 (2020).
- Wang, Y. et al. Catalyst synthesis under CO₂ electroreduction favours faceting and promotes renewable fuels electrosynthesis. *Nat. Catal.* 3, 98–106 (2020).
- Wang, X. et al. Efficient electrically powered CO₂-to-ethanol via suppression of deoxygenation. Nat. Energy 5, 478–486 (2020).
- Xu, F. et al. Study of electro-osmotic drag coefficients in Nafion membrane in acid, sodium and potassium forms by electrophoresis NMR. J. Memb. Sci. 536, 116–122 (2017).
- Sisler, J., Shaihroz, K., Ip, A. H., Jaffer, M. W. S. S. A. & Sargent, E. H. Ethylene electrosynthesis: A comparative techno-economic analysis of alkaline vs membrane electrode assembly vs CO₂–CO– C₂H₄ Tandems. ACS Energy Lett. 6, 997–1002 (2021).
- Weng, L. C., Bell, A. T. & Weber, A. Z. Modeling gas-diffusion electrodes for CO2 reduction. *Phys. Chem. Chem. Phys.* 20, 16973–16984 (2018).
- Singh, M. R., Clark, E. L. & Bell, A. T. Effects of electrolyte, catalyst, and membrane composition and operating conditions on the performance of solar-driven electrochemical reduction of carbon dioxide. *Phys. Chem. Chem. Phys.* 17, 18924–18936 (2015).

- Singh, M. R., Goodpaster, J. D., Weber, A. Z., Head-Gordon, M. & Bell, A. T. Mechanistic insights into electrochemical reduction of CO₂ over Ag using density functional theory and transport models. *Proc. Natl Acad. Sci. USA* 114, E8812–E8821 (2017).
- Weng, L. C., Bell, A. T. & Weber, A. Z. Towards membrane-electrode assembly systems for CO₂ reduction: A modeling study. *Energy Environ. Sci.* 12, 1950–1968 (2019).
- Vermaas, D. A., Sassenburg, M. & Smith, W. A. Photo-assisted water splitting with bipolar membrane induced pH gradients for practical solar fuel devices. J. Mater. Chem. A 3, 19556–19562 (2015).
- Chen, Y., Lewis, N. S. & Xiang, C. Modeling the performance of a flowthrough gas diffusion electrode for electrochemical reduction of CO or CO₂. J. Electrochem. Soc. 167, 114503 (2020).
- Petr, V. Ionic conductiVity and diffusion at infinite dilution. CRC Handb. Chem. Phys. 96, 5–98 (1996).
- Aqion. Table of Diffusion Coefficients. https://www.aqion.de/site/diffusion-coefficients (2020).
- Applin, K. R. & Lasaga, A. C. The determination of SO42-, NaSO4-, and MgSO40 tracer diffusion coefficients and their application to diagenetic flux calculations. *Geochim. Cosmochim. Acta* 48, 2151–2162 (1984).
- Sugisaki, M. Soret coefficients and heat of transport of polyvalent electrolytes in an aqueous solution. Bull. Chem. Soc. Jpn. 48, 2751 (1975).
- Weisenberger, S. & Schumpe, A. Estimation of gas solubilities in salt solutions at temperatures from 273 K to 363 K. AIChE J. 42, 298–300 (1996).
- Ebbesen, S. D. & Mogensen, M. Techno-economic assessment of lowtemperature carbon dioxide electrolysis. *Nat. Sustain.* https://doi.org/10.1002/ 9780470456309.ch25 (2021).
- Renfrew, S. E., Starr, D. E. & Strasser, P. Electrochemical approaches toward CO₂ capture and concentration. ACS Catal. 10, 13058–13074 (2020).
- Keith, D. W., Holmes, G., St. Angelo, D. & Heidel, K. A process for capturing CO₂ from the atmosphere. *Joule* 2, 1573–1594 (2018).
- Ma, S. et al. One-step electrosynthesis of ethylene and ethanol from CO₂ in an alkaline electrolyzer. J. Power Sources 301, 219–228 (2016).
- Yan, Z., Hitt, J. L., Zeng, Z., Hickner, M. A. & Mallouk, T. E. Improving the efficiency of CO₂ electrolysis by using a bipolar membrane with a weak acid cation exchange layer. *Nat. Chem.* 13, 33–40 (2021)

Acknowledgements

The authors acknowledge Ontario Centre for the Characterization of Advanced Materials (OCCAM) for sample preparation and characterization facilities. This work received financial support from TOTAL S.E., Fund No. 508641. R.K.M. thanks NSERC, Hatch, and the Government of Ontario for their support through graduate scholarships. D.S. acknowledges the NSERC E.W.R Steacie Memorial Fellowship. S.W.B. acknowledges funding for the bipolar membrane work from the U.S. Office of Naval Research, grant number N00014-20-1-2517.

Author contributions

E.H.S. and D.S. supervised the project. K.X. introduced the concept, fabricated membrane with the advice from Q.X and S.W.B., performed electrochemical measurements and wrote the manuscript. R.K.M. performed $\rm CO_2$ utilization measurements, advised electrolyzer design, and contributed to the manuscript editing. A.O. prepared electrodes, advised electrolyzer design and contributed to the manuscript editing. S.L. performed the one-dimensional physics modelling. J.E.H, C.M.G., Q.S., C.T.D., Z.C., G.L., J.P.E. C.O. and S.W.B contributed to the discussions and manuscript editing.

Competing interests

K.X., R.K.M., D.S., and E.H.S. have filled a provisional patent application: US 63/201,675 on the SC-BPMEA structure described here. The remaining authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-022-31295-3.

Correspondence and requests for materials should be addressed to David Sinton or Edward H. Sargent.

Peer review information *Nature Communications* thanks Jose Solla-Gullón, and the other anonymous reviewers for their contribution to the peer review of this work.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit https://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2022