Journal of Materials Chemistry A



PAPER

View Article Online
View Journal | View Issue



Cite this: J. Mater. Chem. A, 2022, 10, 20059

In situ regeneration of copper catalysts for longterm electrochemical CO₂ reduction to multiple carbon products†

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The valorization of carbon dioxide (CO₂) via electrochemical CO₂ reduction (ECR) has attracted great interest as a pragmatic approach to tackle greenhouse gas emissions. Multiple carbon (C_{2+}) products, such as ethylene (C_2H_4) , ethanol (C_2H_5OH) , and propanol (C_3H_7OH) , are highly valuable chemicals and of great demand. Copper (Cu)-based catalysts are so far the only electrocatalytic materials that allow CO2 reduction to C₂₊ products at industrially relevant current densities (≥100 mA cm⁻²). However, most Cubased catalysts are unstable in long-term reactions (>100 hours), with the main reasons being the potential-induced surface reconstruction, deposition of impurities, and catalyst aggregation and leaching, among others. Herein, we report an in situ catalyst regeneration strategy that can extend the operation time of Cu-based catalysts. By periodically adding segments of anodic currents to electrolysis, a Cu catalyst is partially oxidized to CuO_x in each cycle, as confirmed by in situ Raman studies, leading to the restoration of the catalytically active sites for C_{2+} products. We found that the oxidation current density and time significantly affect the selectivity and stability of Cu catalysts. Applying this strategy to a Cu catalyst – which is stable for ~ 5 h towards C_{2+} products during a continuous electroreduction under neutral-pH conditions, we were able to extend the operating time to \sim 120 h in a flow cell system. The catalyst maintained a high faradaic efficiency (FE) for C_2H_4 of $\geq 50\%$ at a fixed cathodic current density of 150 mA cm⁻² for over 60 h and continued to operate with a C_2H_4 FE \geq 40% for the entire length of the reaction time. This work opens up an avenue to enhance the stability of Cu electrocatalysts, via controlling the operating procedure during electrolysis.

Received 4th April 2022 Accepted 19th July 2022

DOI: 10.1039/d2ta02709g

rsc.li/materials-a

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† Electronic supplementary information (ESI) available. See https://doi.org/10.1039/d2ta02709g



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Introduction

The imminent threat of rising global temperatures has resulted in shifting weather patterns and disruption of nature's normal balance. Reducing carbon dioxide (CO₂) emissions is critical and urgent; otherwise, humanity will be impacted by more and more severe weather events, together with food insecurity and global health issues. Carbon capture, utilization, and sequestration play an essential role in mitigating CO₂ emissions.^{1,2} The development of CO₂ utilization technologies such as electrochemical CO₂ reduction (ECR) offers a pragmatic strategy to tackle emissions,² as CO₂ is converted into numerous valuable products without concerns of its immediate re-emission to the atmosphere.³

A wide range of electrocatalytic materials for ECR have been investigated and reported in the literature. Electrocatalysts with high selectivity (FE > 80%) and long-term stability (>100 h) when performing at industrially relevant current densities (>100 mA cm⁻²) have been reported for single carbon (C₁) products such as carbon monoxide4-7 and formic acid/formate.8-10 Cu and Cubased catalysts have drawn great interest for ECR studies, not only because of their natural abundance and low cost, but also due to their unique feature as the only catalysts capable of converting CO2 into valuable C2+ products at high current density.11-17 Recent improvements in Cu-based catalysts and cell design have promoted ECR at elevated current densities even beyond 1 A cm⁻².14 In some cases, high selectivity under alkaline and neutral-pH conditions was achieved, with FE values in the range of 40-72% and 41-52% for $C_2H_4^{13,14}$ and $C_2H_5OH_5^{12,15}$ respectively. Interestingly some high-performing ECR Cu-based electrocatalysts with $C_2H_4 \ge 75\%$ 18,19 have also been demonstrated with stability \le 100 h and current density < 100 mA cm⁻². Nevertheless, the reported stability of ECR has been limited to a few operating hours 13,20,21 for these C_{2+} products. In most cases, the Cu-based catalysts are rapidly deactivated, particularly when ECR reactions are conducted in neutral-pH or acidic media, as the competing hydrogen evolution reaction (HER) prevails after a short period.²² Notably, performing ECR under alkaline conditions has previously been shown to extend the catalyst operating time while displaying good selectivity for C₂₊ products. However, under these conditions, CO₂ directly reacts with KOH and forms bi/carbonate salts, necessitating additional energy expenses for electrolyte and CO₂ regeneration.13 Thus, there is a need to design suitable electrocatalysts and/or devise electrolysis procedures that can stably produce C₂₊ products in neutral or acidic environments to make the ECR technology more economically competitive.

Improving the stability of Cu electrocatalysts in low pH media still remains a great challenge. ^{20,23-26} A wide array of reasons for Cu catalyst electrode failure have been reported, including poisoning by reaction products, ²⁷ deposition of metal impurities from the electrolyte, ²⁸ potential-induced surface reconstruction exposing inactive facets, ²⁹⁻³⁷ and dissolution and redistribution of the catalyst outer surface. ^{38,39} For gas diffusion electrodes (GDEs), which allow ECR to operate at industrially relevant current densities, catalyst deactivation is also due to the flooding of the

porous gas diffusion layer and the precipitation of bi/carbonate salts that block the transport of CO₂ gas to catalytic sites. To address the instability of Cu-based catalysts, strategies such as pulse electrolysis^{25,26,40-42} or employing electrolytically-cleaned electrolytes have been reported.43 It is also worth knowing that pulsed ECR has also been extensively studied towards enhancing C₂₊ product selectivity. 44-47 Pulsed potential electrolysis has been exploited towards enhancing the stability of Cu-based electrocatalysts. Jännsch Y. et al. 26 used 25 s at -1.38 V and 5 s at -1.0 V (vs. Ag/AgCl) to increase the stability of C2H4 production from less than 8 h to at least 16 h. Engelbrecht A. et al. 25 extended a 16 h potentiostatic electrolysis to about 85 h and 95 h via variation of cathodic pulse potential ($t_c = 25 \text{ s}, t_a = 5 \text{ s}, U_c = -1.6 \text{ V}, U_a =$ -0.18 V, vs. Ag/AgCl) in combination with two different modes of electrolyte management, respectively. They both achieved C₂H₄ FEs below 40% which were demonstrated at current densities below 100 mA cm⁻². However, to the best of our knowledge, none of these approaches have been able to demonstrate long-term stability towards C2+ products with high FE and at a high reaction rate in a flow cell configuration.

Herein, we report an *in situ* surface regeneration strategy to extend the operating time of a Cu electrocatalyst in neutral-pH media. The strategy is based on previous observation that oxide-derived Cu (OD-Cu) catalysts are effective for selective CO_2 reduction to C_{2+} products. ^{48,49} We exploited the copper oxide formed during the regeneration step of a thoroughly optimized oxidation and reduction cycle to target an extended ECR to C_{2+} products. With this *in situ* regeneration strategy, we achieved an order of magnitude improvement in operating time, with the Cu electrode operating at a cathodic current density of 150 mA cm⁻² for \sim 120 h. It maintained FE values for $C_2H_4 \geq 50\%$ for over 60 h and remained operating with C_2H_4 FE \geq 40% for the entire reaction time.

Results and discussion

Performance of Cu/PTFE catalysts at high current densities in a neutral-pH electrolyte

To fabricate Cu gas diffusion electrodes, we deposited metallic Cu on a porous polytetrafluoroethylene (PTFE) membrane with an average pore size of 0.45 microns, using the sputtering technique. The PTFE membrane was chosen because it has been demonstrated to be a durable gas diffusion layer, significantly suppressing flooding, allowing stable ECR to C2+ products in flow-cell configurations under highly alkaline conditions.13 The nominal thickness of the Cu layer was controlled, varying from 200 to 1000 nm. The morphologies of Cu/PTFE samples were characterized by scanning electron microscopy (SEM) (Fig. 1a-c). Cu particles were conformally deposited on PTFE fibers, forming a continuous Cu layer shell on the PTFE fiber core. The size of the Cu/PTFE fiber increases with increasing the Cu nominal thickness, leading to smaller pore size in the PTFE membrane (Fig. 1a-c and S1†). High magnification images show a similar Cu surface texture of small Cu particles with multiple grain boundaries for all Cu thicknesses (Fig. 1a-c and S1†).

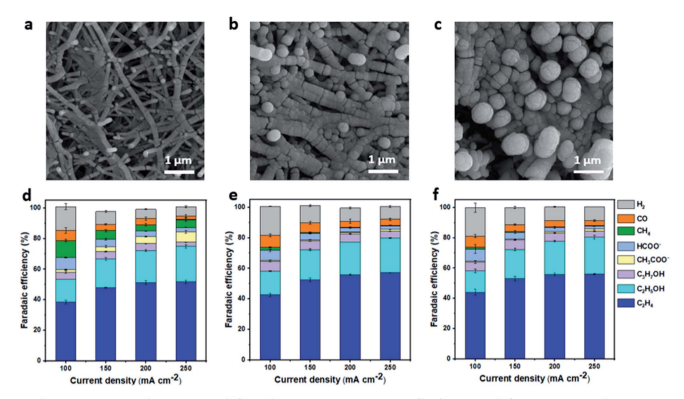


Fig. 1 Characterization and ECR selectivity of Cu/PTFE. Scanning electron microscopy (SEM) images of Cu/PTFE with nominal Cu thicknesses of 200 nm (a), 700 nm (b) and 1000 nm (c). CO_2 reduction product distribution at different cathodic current densities in 1 M KHCO₃ electrolyte using Cu/PTFE with nominal thicknesses of 200 nm (d), 700 nm (e) and 1000 nm (f). The FE error bar was calculated based on data collected from different experimental trials (n = 3). Potential data for the determination of ECR selectivity of Cu/PTFE for all samples are provided in the ESI (Fig. S3).†

ECR performance of Cu/PTFE GDEs was evaluated with a flow-cell reactor using 1 M KHCO3 catholyte. A bipolar membrane was used to separate the anode and cathode compartments. Nickel (Ni) foam and 1 M KOH solution were used as the anode and anolyte, respectively. The ECR of Cu/ PTFE catalysts was studied in the cathodic current density range of 100-250 mA cm⁻² using chronopotentiometry. Gaseous products were analyzed with in-line gas chromatography while liquid products were collected and analyzed using nuclear magnetic resonance (NMR) (Fig. S2†). All Cu/PTFE samples showed FE values for C₂H₄ between 38 and 55% in the studied current range, with a total C₂₊ (C₂H₄, C₂H₅OH, C₃H₇OH, and C₂H₃O₂⁻) FE of >70% achieved at current densities of 150-250 mA cm⁻² (Fig. 1d-f). This performance is consistent with previous reports on Cu/PTFE catalysts using similar testing conditions.21,50 We observed that Cu thickness showed little impact on the overall product selectivity within the explored current density range. While the total C2+ was lower at 100 mA cm⁻² for all Cu/PTFE samples, the value increased about 15-20% at current densities between 150 and 250 mA cm⁻². At these higher current densities, the Cu/PTFE GDEs maintained a good C₂₊ FE of >70%. However, at current densities beyond 300 mA cm $^{-2}$ we observed a decrease in the total C_{2+} FE with a corresponding increase in H₂ and CH₄. The FE value of CH₄ was found to be the highest on the 200 nm Cu/PTFE sample and was approximately 10 times higher at all current

densities explored. These results suggest that the thickness of Cu/PTFE can be optimized to either achieve a high FE for C2+ products (thick Cu/PTFE) or a high FE for CH₄ (thin Cu/PTFE).

To study the stability of the GDEs, we performed the ECR reaction at a fixed current density of 150 mA cm⁻² and analyzed the gaseous products over time. The thickness of the Cu layer had a significant effect on the stability of Cu/PTFE. For the 200 nm Cu/PTFE, the C2H4 FE was stable at around 50% for about 1 h before it started decreasing rapidly, reaching 15% after 1.5 h of continuous reaction (Fig. 2a). After 1 h of reaction, the FE values of both CH₄ and H₂ started increasing, with H₂ being the dominant product after 1.5 h (Fig. 2a-c). The stability of Cu/PTFE catalysts was significantly improved with an increase in Cu nominal thickness. The 1000 nm Cu/PTFE maintained a C₂H₄ FE of 50% for 5 hours (Fig. 2a). For all Cu/ PTFE samples, CH₄ and H₂ were the two main products when the C₂H₄ FE started decreasing (Fig. 2b and c), while CO selectivity slowly decreased over time (Fig. S4†).

Previous studies on the stability of Cu catalysts for ECR in aqueous electrolytes have identified three main deactivation mechanisms among others. First, the Cu surface undergoes a reconstruction process, exposing the Cu sites (e.g. (111) facet) that are more selective towards CH₄ formation and H₂ production.51 This surface reconstruction usually occurs on the sub-10 nm scale. Second, metal impurities such as Fe and/or Ni present in the electrolyte, which can be deposited on the surface

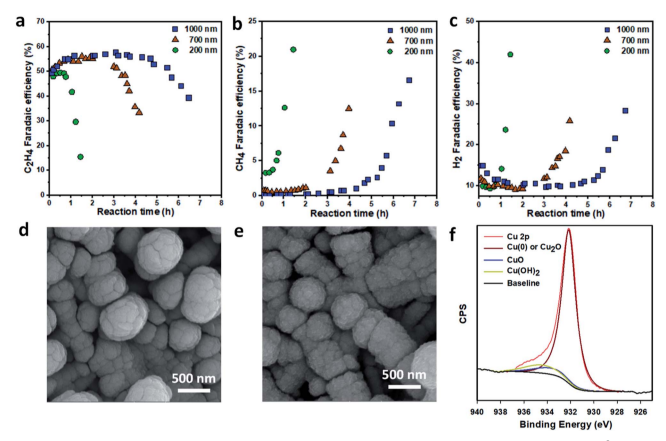


Fig. 2 Stability of Cu/PTFE in ECR. Variation of gas products over reaction time at a constant current density of 150 mA cm $^{-2}$ in 1 M KHCO $_3$ electrolyte: (a) C₂H₄, (b) CH₄ and (c) H₂. SEM images of Cu/PTFE catalysts with a nominal thickness of 1000 nm before (d) and after continuous electrolysis for 7 hours (e). High resolution Cu 2p XPS spectrum of Cu/PTFE catalysts after continuous ECR tests showing the presence of metallic Cu, copper oxide and copper hydroxide (f). Chronopotentiometric plots for the three samples are provided in the ESI (Fig. S5-S7).†

of Cu catalysts during electrolysis, form highly active sites for H₂ production.52 The last mechanism involves the reduction of oxidized Cu species (Cu⁺, Cu²⁺) in Cu-based catalysts during the ECR process. Because oxidized Cu species and/or sub-surface oxygen are critical for CO2 reduction to C2+ products, the disappearance of these species makes the catalyst less selective toward C₂₊ products over time.⁵³⁻⁵⁵ Furthermore, for gas-phase ECR in flow systems using gas diffusion electrodes, the flooding of the gas diffusion layer and the precipitation of bi/ carbonate salts can block the diffusion of the CO₂ reactant, leading to the suppression of CO₂ reduction.³

To identify the deactivation mechanism in our system, we first checked if flooding and salt precipitation were the main reasons. ECR at 150 mA cm⁻² was performed continuously until the C_2H_4 FE decreased to around 25%. The current density was then reduced to 125, 100, 75 and 50 mA cm⁻² and the gaseous products were analyzed accordingly. The FEs for H2 and CH4 remained high with lowering the current density, implying that CO₂ diffusion limitation was not the reason for the drop in C₂H₄ FE (Fig. S8†). To further confirm that flooding was not the major mechanism for the deactivation of our Cu/PTFE sample, we investigated electrode flooding in our flow cell system using a capacitance change measurement.⁵⁶ We quantified the electrochemical double-layer capacitance in an attempt to track the ingress of the electrolyte into the GDE during electrolysis. This capacitance measurement obtained through the cyclic voltammetry technique, in the non-faradaic region with variable scan rates, can offer insights into the changes that occurred at the electrode-electrolyte interface. The cyclic voltammetry was carried out between -0.05 V and -0.20 V vs. Ag/AgCl with scan rates of 10, 50, 100, 150 and 200 mV s^{-1} . We observed no significant change in the electrochemical double-layer capacitance (Fig. S9-S11†). The capacitance was found to be around 0.5756 mF cm⁻² at the open circuit potential before the start of continuous electrolysis, 0.5309 mF cm⁻² after 3 h of electrolysis and 0.5143 mF cm⁻² after approximately 6-7 h of continuous operation (Fig. S11†). The FE of C₂H₄ was about 56% after 3 h and decreased to about 42% at the end of the continuous operation.

To check for a change in the catalyst surface morphology, we compared the SEM images of the Cu/PTFE sample before and after continuous ECR tests (Fig. 2d and e). The surface texture of Cu/PTFE appeared to change after the reaction as the particle size became smaller. While this could be due to the surface construction, it may also have originated from the oxidation of the Cu surface when the sample was exposed to air after the reaction. High resolution Cu 2p X-ray photoelectron spectroscopy (XPS) of the sample (1000 nm) shows the presence of CuO

and Cu(OH)₂ after the reaction (Fig. 2f). The survey XPS spectra show no peaks for Fe and Ni in all samples while small peaks of C, F and Cl were detected (Fig. S12†). These data suggest that impurity could not be the main cause of lower C₂H₄ selectivity. We therefore reason that surface reconstruction and changing of the Cu oxidation state and sub-surface oxygen could be the main contributors to the deactivation of Cu/PTFE catalysts.

Regeneration of Cu catalysts by in situ oxidation

The development of highly selective OD-Cu catalysts for CO2 reduction to C₂₊ products^{48,49,57-60} motivated us to pursue an in situ regeneration of the deactivated Cu catalyst, using an electrochemical oxidation process to transform Cu back to Cu oxide. We reason that this oxidation step may address the main deactivation mechanisms of Cu-based catalysts since it can reproduce the selective OD-Cu catalysts. In addition, it might also enable the dissolution of metal impurities such as Fe on the Cu surface back to the electrolyte.

To investigate the recovery of Cu catalyst selectivity toward C₂H₄ via in situ electrochemical oxidation, we first performed continuous ECR tests at a current density of 150 mA cm⁻² until

the C₂H₄ FE decreased to around 10-15% (Fig. 3a). Next, we performed repeated oxidation-reduction cycles until a stable C₂H₄ FE was achieved (Fig. 3a). The oxidation current density was varied between 0.4 and 1 mA cm⁻² while the reduction current density and time were fixed at 150 mA cm⁻² and 5 min, respectively. The FE of C₂H₄ was analyzed at the end of each reduction cycle. The oxidation time was studied to maximize C_2H_4 FE. In all cases, the oxidation charges $(Q = I \times t)$ explored were either 36, 24, or 12 mC. As shown in Fig. 3b-e, the C₂H₄ FE increases after each oxidation-reduction cycle while the FEs for H₂ and CH₄ slowly decrease (Fig. S13-S16†). The FE for CO remains relatively constant with the oxidation-reduction cycle (Fig. S13-S16†). C₂H₄ FE values >50% were achieved at all oxidation current densities, with higher applied charges allowing the catalysts to operate with higher C₂H₄ FE for each oxidation current density. This suggests that a higher oxidation current density only needs a shorter time to recover the high FE for C_2H_4 .

The maximal FE for C₂H₄ was found to be a function of the oxidation charge regardless of the oxidation current density applied in our studied range (Fig. 3f) with a FE of 53-56% being

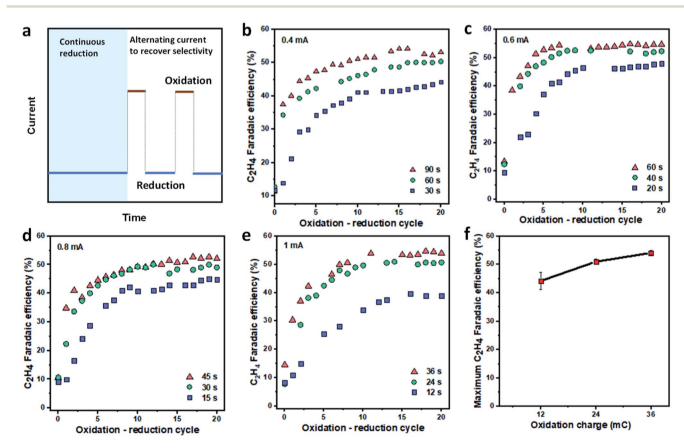


Fig. 3 Regeneration of Cu catalysts by in situ oxidation. (a) Oxidation-reduction program for the recovery of C_2H_4 FE. All samples were operated continuously at a fixed reduction current density of 150 mA cm $^{-2}$ until the C_2H_4 FE decreased to around 10–15%. The catalysts were then subjected to 20 repeated oxidation-reduction cycles. Each oxidation-reduction cycle involves an oxidation step at a designated current and time, followed by ECR at 150 mA cm $^{-2}$ for 5 minutes. The gas products were analyzed at the end of each reduction cycle. The recovery of FE for C_2H_4 is shown with oxidation current densities of 0.4 mA cm⁻² (b), 0.6 mA cm⁻² (c), 0.8 mA cm⁻² (d) and 1 mA cm⁻² (e) at different oxidation periods. The dependence of maximum C₂H₄ FE on oxidation charge (f). The deviation was calculated based on data at different oxidation current densities but with similar oxidation charges (n = 4). The FE values for CO, H₂ and CH₄ during the oxidation–reduction cycle are provided in the ESI (Fig. S13-S16†). The morphology of Cu/PTFE after in situ electrochemical oxidation is presented in the ESI (Fig. S17†).

recovered when the oxidation charge was \geq 24 mC. In addition, at each given oxidation current density and time condition, we found that the C_2H_4 FE quickly recovered in the first 10 cycles before reaching a plateau. The presence of such a plateau suggests that (i) the 5 min ECR reaction after each oxidation step is not long enough to cause catalyst deactivation and (ii) the ratio between the catalytic sites for C_2H_4 and other products is dependent on the applied charge of the oxidation current density, which governs the final C_2H_4 FE.

In situ Raman studies on oxidized Cu species

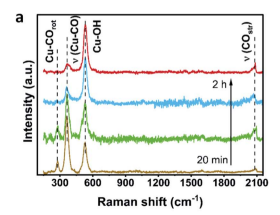
To shed light on how the oxidation current density and time affect product selectivity, we performed operando Raman analysis. This technique allowed us to investigate the catalyst structure and surface adsorbates during reduction and oxidation. We first studied the surface of Cu/PTFE catalysts during a continuous ECR process. The CO intermediate is usually observed by surface enhanced Raman scattering (SERS) at three Raman shift regions: 280-290 cm⁻¹ for the restricted rotation of adsorbed CO, 360-370 cm⁻¹ for Cu-CO stretching, and 1800-2100 cm⁻¹ for C≡O stretching.^{61,62} All in situ Raman measurements in this study were performed using a 785 nm laser which exhibits suitable SERS activity. Signal enhancements were performed and control experiments with two different laser wavelengths were carried out to study the effect of laser excitement wavelength (Fig. S18†). The Raman shift in the range of 520-530 cm⁻¹ can be assigned to the Cu-OH band. It has been shown that higher intensity of the Cu-CO stretching band can be assigned to a higher coverage of the CO intermediate and higher C₂₊ selectivity.⁶³ A significant decrease in the intensity of restricted rotation of adsorbed CO and Cu-CO stretching compared to the Cu-OH can be seen by increasing the reduction time (Fig. 4a). This is a sign of the deactivation of copper by less CO coverage resulting in a reduced amount of C₂H₄. These observations are in agreement with the drop in C₂H₄ FE over the reaction time.

Given the observed removal of adsorbed CO species and a significant drop in C_2H_4 formation, we performed in situ

Raman studies using the oxidation-reduction strategy as a remedy against this deactivation. As the oxidation current density was applied, the Cu-CO and Cu-OH species disappeared, and the CuO_x bands emerged in the range of 450-700 cm⁻¹. These oxide species were visible after 20 s of oxidation current (Fig. 4b). The CuO_r bands disappeared entirely while the Cu-CO and Cu-OH species reappeared after the reduction cycle. It has been shown that these CuO_r crystals reduced into small Cu grains developing OD catalysts which facilitated the C-C coupling and accounted for enhanced C2H4 selectivity. 63-65 Interestingly, by applying cathodic current the restricted rotation of adsorbed CO and Cu-CO stretching is recovered, confirming the re-coverage of CO. By repeating the oxidation-reduction cycles, more intense Cu-CO bonds were formed, which indicates enhanced CO coverage. Roldán Cuenya et al.63 showed that the intensity of ratio of the Cu-CO stretching band to the CO rotation band can be correlated to the CO coverage and a higher ratio leads to more C-C coupling and C2+ formation. Our SERS analysis illustrates an increasing trend of this ratio with increasing the number of oxidation-reduction cycles. This confirms a better CO coverage on the surface of the catalyst leading to the sustained C₂H₄ FE (Fig. S19†).

Extending catalyst lifetime with the alternating current strategy

Having identified the oxidation conditions for recovering C_2H_4 FE, we set out to prolong the catalyst lifetime by performing an alternating oxidation–reduction current procedure (Fig. 5a). The reduction current density was fixed at 150 mA cm⁻² while the oxidation current density and time, as well as the reduction time, were varied to optimize the operation conditions. We first fixed the oxidation charge of 36 mC and varied the oxidation current density (from 0.4 to 1 mA cm⁻²) and oxidation time (36 to 90 s). The reduction current density and time were 150 mA cm⁻² and 5 min, respectively. We found that all operational procedures showed similar stability, maintaining C_2H_4 FE \geq 50% for at least 10 hours (Fig. 5b), doubling the time of the continuous reduction procedure. The FE for CO slowly



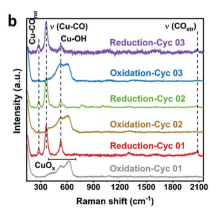


Fig. 4 In situ Raman studies on oxidized Cu species of Cu/PTFE. (a) In situ SERS of Cu/PTFE conducted at different times of reaction in $KHCO_3$ electrolyte at a current density of 30 mA cm⁻². (b) The structure and surface adsorbates during reduction and oxidation cycles at 0.6 mA cm⁻² oxidation current density for 20 s.

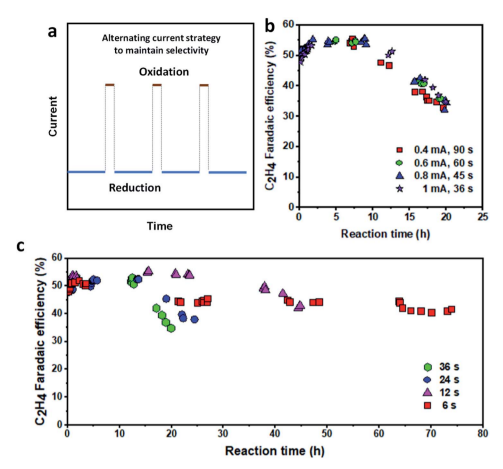


Fig. 5 Stability of the Cu/PTFE catalysts with the alternating current strategy. (a) Schematic illustration of the alternating current strategy. (b) Oxidation charge optimization at a fixed reduction current density of 150 mA cm⁻² for 5 min reduction time. The cathodic potential versus time plot is provided in Fig. S21.† (c) Oxidation time optimization at a fixed current density of 150 mA cm⁻² for 5 min reduction time and an oxidation current density of 1 mA cm $^{-2}$ (the oxidation time was varied between 6 s and 36 s). The cathodic potential versus time plot is provided in Fig. S23.†

decreased while H₂ and CH₄ FEs increased over time (Fig. S20†). Therefore, varying the oxidation current density and time plays a negligible role if the oxidation charge is fixed.

To further explore the effect of the oxidation charge, we then fixed the oxidation current density at 1 mA cm⁻² and varied the oxidation time from 6 to 36 s. The reduction current density and time are 150 mA cm⁻² and 5 min, respectively. While larger oxidation charge leads to higher C₂H₄ FE, the catalyst becomes less stable when the oxidation time (oxidation charge) is increased (Fig. 5c). With a 36 s oxidation time, the catalyst showed the highest C₂H₄ FE of 54% but was stable at above 50% for 14-16 h before dropping quickly to about 30% in 20 h of operation. In contrast, when the oxidation time was reduced to 6 s, the catalyst operated stably for up to 75 h, although with a lower C₂H₄ FE of 40% and above (Fig. 5c). Under all operating conditions, the FE of CO slowly decreased while those of H2 and CH₄ increased steadily with the reaction time (Fig. S22†). These results suggest that the oxidation time could not be tuned to achieve both high C₂H₄ FE and long stability simultaneously. While the catalyst can operate with an initial C2H4 FE of about 50% at all oxidation times, each operating condition tends to temporarily arrive at a constant FE as mentioned earlier, beyond

which the cumulative effect of the total oxidation time over the catalyst operating lifetime begins to significantly impact the catalyst performance. Therefore, it can be hypothesized that the total amount of charge (the product of a given oxidation charge and the number of times it was cycled) throughout the lifetime of the operating catalyst determines the overall stability of the Cu/PTFE catalyst electrode.

To extend the lifetime of the catalyst, we turned our attention to the reduction time. We reason that if the stability of the catalyst depends only on the oxidation charge, the overall operating time can be extended by increasing the reduction time. We fixed the reduction current density (150 mA cm⁻²), oxidation current density (1 mA cm^{-2}) and time (24 s) and varied the reduction time. As shown in Fig. 6a, increasing the reduction time extended the overall catalyst lifetime (time to maintain a C_2H_4 FE $\geq 40\%$). Based upon our optimized operating conditions, the longest catalyst lifetime of \sim 120 h was achieved at a reduction time of 45 min. At this condition, the C₂H₄ FE was maintained at over 50% for more than 60 h, which is around ten times longer than the continuous operation (5 h). The CO FE decreased rapidly in the first 20 h and is then relatively stable (Fig. S24†). The CH₄ FE slowly increased over time

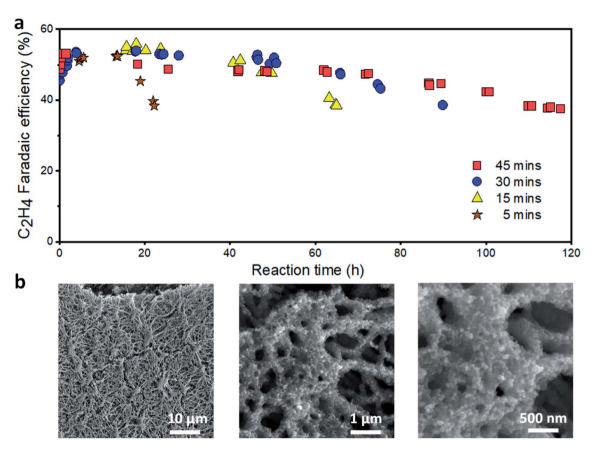


Fig. 6 Stability of the Cu/PTFE catalysts with the optimized alternating current strategy in 1 M KHCO₃. (a) Reduction time optimization (varying from 5 to 45 min) at 150 mA cm⁻² and a fixed oxidation current density and time of 1 mA cm⁻² and 24 s, respectively. (b) SEM of Cu/PTFE after the oxidation–reduction strategy using 45 min reduction at a current density of 150 mA cm⁻² and a fixed oxidation current density and time of 1 mA cm⁻² and 24 s, respectively. The cathodic potential *versus* time plot is provided in Fig. S27.†

while the H_2 FE varied in the range of 10–15% (Fig. S25 and S26†). Furthermore, under these operating conditions, we found that a further increase of the reduction time and/or oxidation charge does not improve the catalyst lifetime, at least for when we aim to maintain a C_2H_4 FE \geq 40%. An operation of the catalyst under the same conditions, but with a higher oxidation charge (60 mC), was carried out for \sim 58 h (Fig. S28†).

A limited Cu catalyst lifetime with the optimized oxidation and reduction conditions implies a major deactivation mechanism different from the potential induced surface reconstruction and surface impurities discussed above. To identify the major cause of the gradual decrease in C2H4 FE over the operating time, we characterized the Cu catalyst after the reaction using SEM. The catalyst layer (Fig. 6b) was found to be much thinner than that for the catalysts before (Fig. 2d) or after continuous ECR tests (Fig. 2e). We reason that the Cu at the Cu/ PTFE interface and at the top of the catalyst layer can migrate during the oxidation cycle *i.e.*, when Cu is oxidized to Cu oxide. This gradual migration of Cu species over the reaction time could lead to the redistribution of Cu on the PTFE substrate, forming Cu/PTFE locations with different Cu thicknesses. In a flow cell configuration, phenomena such as catalyst fragmentation⁶⁶ and agglomeration,^{37,67} Ostwald ripening,^{36,68}

particle dissolution and (electro-) redeposition³⁹ of dissolved copper species in the form of new smaller Cu fragments, as we observed in Fig. 6b, cannot be neglected. Over time, these effects could result in the formation of some areas with very thick Cu while the Cu layer may disappear in other areas, leading to the degradation of the Cu/PTFE catalyst layer when using the alternating current approach. In addition, the impact of other associated failure mechanisms resulting from the type of electrolyzer system architecture being used can lead to a systemic failure of the overall cell system. For a flow cell architecture, as we employed in this study, the additional impact of flooding and salt formation within the PTFE pores cannot be ignored for an extended testing time. These failure mechanisms, through the interdependent nature of the cell system architecture, directly impact and combine with the major failure resulting from our strategy to suppress the stability performance of the electrolyzer system and thereby limit its overall operating time.

Conclusions

In this work, we have introduced an *in situ* regeneration of Cu catalysts for stable CO_2 reduction to multiple carbon products. Our strategy involves an alternating current operating

procedure in which the catalysts are periodically regenerated by the oxidizing Cu surface. By optimizing the oxidation and reduction current density and time, a ten-fold increase in Cu catalyst lifetime has been achieved. Our optimal oxidationreduction strategy enables the Cu catalyst to operate for an extended time (~120 h) and maintain a significant FE of ≥50% at a fixed current density of 150 mA cm⁻² for the first 62 h, while the reaction continued with a selectivity of \geq 40% for the entire length of the operation. In situ Raman characterization confirmed the presence of copper oxide species during the oxidation cycle. This approach could potentially open up an opportunity for the long-term use of Cu and Cu-based catalysts for effective ECR to C2+ products and the realization of the electrochemical CO2 conversion technology.

Experimental

Electrode preparation

The copper cathode gas diffusion electrodes were prepared by magnetron sputtering using a Kurt J. Lesker PVD 75 vacuum deposition system. Pure Cu sputtering (99.99%) metal targets (3" diameter) were used with a DC source in pure argon plasma in a sputter-down configuration. The samples were affixed to a 150 mm diameter rotating platen with Kapton tape, with a target-to-platen distance of ~165 mm. Cu was sputtered onto a porous hydrophobic polytetrafluoroethylene (PTFE) gas diffusion layer with a 450 nm mean pore size using 120 W DC power and 4 mTorr pressure to achieve a deposition rate in the range of 0.76-1.06 angstroms per second. Approximately 200 nm, 700 nm, and 1000 nm nominally thick Cu films were sputtered onto the PTFE substrate.

Electrochemical CO₂ reduction

Electrochemical CO₂ reduction experiments were performed in a customized gas-fed flow cell configuration. The flow cell is made up of three compartments which are for CO2 gas, catholyte, and anolyte. Aqueous solutions of 1 M KHCO3 (Sigma-Aldrich) and 1 M KOH (Sigma-Aldrich) were used as the catholyte and anolyte, respectively. A bipolar membrane (Fumasep) was used for the experiment. The sputtered Cu/PTFE gas diffusion electrode (GDE), which is the working electrode, was carefully positioned between the catholyte chamber and a continuous flow of CO2 gas stream through the back of the cathode compartment. The catalyst side of the cathode compartment of the GDE was made to face the electrolyte. The cathode chamber was equipped with a leak-free Ag/AgCl in 3 M KCl reference electrode. The orientation of the bipolar membrane used was reverse biased with respect to the electrodes, where the cation-exchange layer (CEL) was made to face the cathode and the anion-exchange layer (AEL) was made to face the anode, thus maintaining separate pH in both sides of the electrolyzer compartment. Nickel foam (MTI Corp.) was used as the counter electrode in the anode compartment. Gaseous CO2 was allowed to flow past the back of the GDL at a constant flow of 30 sccm by means of a mass flow controller. Since only a fraction of the CO₂ gas is used in the reaction

process towards CO₂RR products, the electrolyzer outlet flow rate from the CO2 gas chamber was measured using a digital volumetric flow meter and was used in the calculation of the faradaic efficiency of the reaction products. The electrolytes were circulated in both compartments using peristaltic pumps. The electrochemical measurement experiments were performed using a potentiostat (Metrohm). The electrochemical potentials were compared against Ag/AgCl (3 M KCl) reference.

The gaseous products were quantified using online gas chromatography (GC, PerkinElmer Clarus 590). The GC was equipped with a thermal conductivity detector (TCD) and a flame ionization detector (FID). The chromatography instrument was directly connected to the outlet of the CO2 gas chamber of the customized gas-fed flow cell for continuous online analysis. The liquid products were analyzed using nuclear magnetic resonance spectroscopy (NMR). The ¹H NMR spectra of freshly collected liquid products were acquired on an Auto-400 ultrashield Bruker instrument operating at a denoted spectrometer frequency given in megahertz (MHz) at 25 °C in D₂O using water suppression mode, with dimethyl sulfoxide (DMSO) as a reference.

All potential data were presented versus Ag/AgCl. However, iR-compensation was performed, and the details are presented in the supplementary section (Fig. S29 and S30†).

Capacitance change measurement

The double layer capacitance was evaluated by regressing the measured working electrode charging current as a function of potential scan rate. The scan rates of 10, 50, 100, 150 and 200 mV s⁻¹ were applied. The potential range of -0.20 V to -0.05 V vs. Ag/AgCl was chosen only within limits where the HER and OER would be avoided and where capacitive current could be adequately evaluated. Charging current was determined as an average of the absolute values of the anodic and cathodic currents. The capacitance was determined as the value of the slope of the charging current versus the potential scan

Sample characterization

The surface morphology and composition of the catalysts were investigated using scanning electron microscopy (TESCAN-SEM system) operated at 5-10 kV acceleration voltage. X-ray photoelectron spectroscopy (XPS) measurements were carried out with a VG Escalab 220i-XL spectrometer using polychromatic Al Kα radiation (1486.6 eV).

In situ Raman

Raman spectroscopy was executed with a Renishaw Raman spectrometer equipped with a $\lambda = 785$ nm laser as the excitation source and a 63× immersion objective lens. All in situ experiments were carried out with a laser power of 0.1 mW over 10 acquisitions with an exposure time of 10 seconds. The customdesigned flow cell had an electrolyte reservoir and a gas channel separated by a gas diffusion electrode. The immersion objective was dipped in the electrolyte reservoir to collect the spectra. A Pt wire as the anode, sputtered copper on PTFE as the cathode,

and Ag/AgCl as the reference electrode were dipped in the electrolyte reservoir filled with 1.0 M KHCO $_3$ electrolyte. The area of the cathode in this configuration was 1 cm 2 . During the experiments, a continuous 30 sccm CO $_2$ gas flow was delivered to the cathode through the gas channel. The sputtered copper samples were reduced at 30 mA cm $^{-2}$ for at least 2 h before the oxidation/reduction cycles.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

C. A. O. acknowledges the financial support from Queen's University. T. N. N. acknowledges the support from Helen Co., Ltd. M. G. K. acknowledges the support from Canada First Research Excellence Fund at the University of Calgary. C. T. D. acknowledges the financial support from the Natural Sciences and Engineering Research Council of Canada (NSERC) and Queen's University. The authors thank Dr Graham Gibson at Nanofabrication Kingston for his assistance in sample preparation.

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