Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2024, 15, 8451

d All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 14th March 2024 Accepted 30th April 2024

DOI: 10.1039/d4sc01735h

rsc.li/chemical-science

Enhancing C₂₊ product selectivity in CO₂ electroreduction by enriching intermediates over carbon-based nanoreactors†

Min Wang,^{ab} Chunjun Chen, ^{ab} Shuaiqiang Jia,^{ab} Shitao Han,^{ab} Xue Dong,^{ab} Dawei Zhou,^{ab} Ting Yao,^{ab} Minghui Fang,^{ab} Mingyuan He,^{ab} Wei Xia, ^b** * Haihong Wu * * and Buxing Han * b*** * and Buxing Han * b

Electrochemical CO_2 reduction reaction (CO_2RR) to multicarbon (C_{2+}) products faces challenges of unsatisfactory selectivity and stability. Guided by finite element method (FEM) simulation, a nanoreactor with cavity structure can facilitate C–C coupling by enriching *CO intermediates, thus enhancing the selectivity of C_{2+} products. We designed a stable carbon-based nanoreactor with cavity structure and Cu active sites. The unique geometric structure endows the carbon-based nanoreactor with a remarkable C_{2+} product faradaic efficiency (80.5%) and C_{2+} -to- C_1 selectivity (8.1) during the CO_2 electroreduction. Furthermore, it shows that the carbon shell could efficiently stabilize and highly disperse the Cu active sites for above 20 hours of testing. A remarkable C_{2+} partial current density of–323 mA cm⁻² was also achieved in a flow cell device. *In situ* Raman spectra and density functional theory (DFT) calculation studies validated that the * CO_{atop} intermediates are concentrated in the nanoreactor, which reduces the free energy of C–C coupling. This work unveiled a simple catalyst design strategy that would be applied to improve C_{2+} product selectivity and stability by rationalizing the geometric structures and components of catalysts.

Introduction

The escalating emission of greenhouse gases exacerbates climate change, posing a greater threat to the future of humanity. Carbon dioxide (CO_2), one of the primary greenhouse gases, plays a crucial role in this process. Consequently, the electrocatalytic reduction of CO_2 into valuable chemicals and fuels holds immense significance in mitigating climate change, promoting a closed carbon cycle, and facilitating the storage of renewable electricity. The primary outputs of the electrochemical CO_2 reduction reaction ($\mathrm{CO}_2\mathrm{RR}$) consist of C_1 products and multicarbon (C_{2+}) products. Within these products, C_{2+} products hold greater economic significance within the chemical industry due to their higher commercial value. Currently,

obtain high-productivity of C_{2+} products.⁷⁻¹⁰ Among them, copper (Cu) is recognized as the unique metal to efficaciously break the stable CO_2 bonds and form C–C bonds, a key step towards C_{2+} products.¹¹⁻¹⁴ However, the challenge of C_{2+} selectivity still limits its economic competitiveness, and more advanced strategies are needed to design high-performance catalysts.

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Based on previous studies, C-C coupling is considered a crucial route for the formation of C₂₊ products in the CO₂RR, whereas the C-C coupling reaction relies heavily on the *CO intermediate dimerization.¹⁵ Achieving a local high concentration of *CO intermediates at the active sites is crucial for initiating the dimerization process between neighbouring *CO intermediates.¹⁶ But *CO species tended to diffuse from the catalyst surface to the electrolyte, and it may result in their premature discharge from the active sites.¹⁷ This premature release severely hampers the efficiency of C-C coupling, thus reducing the overall performance of the CO₂RR process.¹⁸ A simple way to increase the local concentration of *CO species is to increase the diffusion resistance and extend the diffusion path length.

Previous studies have demonstrated that the confinement effect can alter the diffusion kinetics and effectively improve the local concentration of key intermediates. Sargent *et al.* applied finite element method (FEM) simulations and experimental

[&]quot;Shanghai Key Laboratory of Green Chemistry and Chemical Processes, State Key Laboratory of Petroleum Molecular & Process Engineering, School of Chemistry and Molecular Engineering, East China Normal University, Shanghai 200062, China. E-mail: wxia@chem.ecnu.edu.cn; hhwu@chem.ecnu.edu.cn; hanbx@iccas.ac.cn

^bInstitute of Eco-Chongming, 20 Cuiniao Road, ChenjiaTown, Chongming District, Shanghai 202162, China

Beijing National Laboratory for Molecular Sciences, CAS Key Laboratory of Colloid and Interface and Thermodynamics, CAS Research/Education Center for Excellence in Molecular Sciences, Institute of Chemistry, Chinese Academy of Sciences, Beijing 100190, China

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d4sc01735h

analyses to explore the C-C coupling reactions by the confinement effect.20 They demonstrated that Cu2O with the cavity structure led to a high surface coverage of intermediates and improved the electrocatalytic conversion of CO-to-propanol. Yu et al. conducted a comprehensive investigation on the multicavities of Cu₂O via intermediate confinement to enhance the selectivity of CO₂ electroreduction to C₂₊ fuels.²¹ These findings revealed that this unique cavity structure effectively suppressed the loss of key intermediates, and enhanced their local concentration at active sites, thus increasing the chance of C-C coupling reactions. Studies on the steric confinement effect in the CO2RR have substantiated its capability to enhance the selectivity of multicarbon products. 22,23 However, a persistent challenge in current studies about Cu-based nanoreactors is Cu surface reconstruction and compositional changes during electrolysis, which makes product selectivity and activity decline significantly.24 Also, the designability of pure Cu-based catalysts is very limited.

It is known that supported catalysts, in which active sites are anchored on supports, have some obvious advantages, such as the size and dispersion of the active sites can be tuned, and the synergy of the active sites and supports. Specifically, carbon materials exhibit advantages in electrocatalysis, including strong stability, ease of morphological adjustments and optimized reaction intermediate adsorption that make it an excellent candidate for supporting active sites. Kia *et al.* proposed a carbon protected indium oxide electrocatalyst, where the carbon layer not only prevents the reductive corrosion of indium oxide during electrolysis, but also optimizes the intermediate adsorption, thus improving the stability and activity of CO₂ reduction. Therefore, it is important to introduce a carbon layer into catalyst design to achieve electrochemical stability while maintaining high activity.

Combining steric confined carbon supported nanoreactors with highly active Cu sites would establish a distinctive class of catalysts with both exceptional C2+ product selectivity and stability. Thus, we began using FEM simulations to explore the influence of cavity structures on the accumulation of *CO intermediates and C-C coupling. Guided by FEM results, we synthesized a Cu/C-cavity nanoreactor in which Cu species are supported in the carbon shell. The unique structure endows the Cu/C-cavity catalyst with high CO₂ electrocatalytic performance. As a result, the nanoreactor achieved a remarkable multicarbon (C_{2+}) product faradaic efficiency (FE) of 80.5% and a higher C_{2+} to- C_1 ratio of 8.1 at -1.2 V vs. RHE. Based on the highly dispersed Cu active sites on the carbon support to prevent agglomeration, the nanoreactor performed continuous electrolysis for more than 20 hours. In situ Raman spectra and density functional theory (DFT) calculation results revealed that the cavity of the nanoreactor could concentrate *CO_{atop} intermediates and reduce their dimerization barriers. This work introduces a convenient and efficient synthesis strategy to enhance the selectivity of multicarbon products and stability for the electrochemical CO₂RR. This strategy could applicable to other reactions, rationalizing the morphology and active components of the catalyst, and concentrating key intermediates through nanoconfinement to improve the selectivity of ideal products.

Results and discussion

It is assumed that the cavity structure nanoreactor could retard the diffusion kinetics through the confinement effect and concentrate *CO species, thus increasing the possibility for C-C coupling. To verify the hypothesis, we applied FEM simulations to explore the prospects of cavity structure in enhancing C2+ product selectivity. A hollow spherical shell model with circular openings (with an outer diameter of 39 nm and an inner diameter of 6.5 nm) was used to represent the nanoreactor immersed in an electrolyte (see details in the ESI†). The out flux of *CO and C2 products on the inner and outer surfaces of the nanoreactor was monitored. Results of the simulation indicated that CO₂ molecules first diffused to the surface (Fig. 1a). Then, CO2 was adsorbed and reduced into *CO at the interior and outer surfaces of the spherical shell (Fig. 1b). Finally, the *CO species could desorb from the surface as C1 products (CO assumed) or be coupled with other *CO to form C2 products (C2H4 assumed). The surface C2 species could diffuse into the electrolyte or continue to react with another *CO to produce C₃ products. Consistent with our hypothesis, the cavity structure could restrict the diffusion of internally generated *CO intermediates to the outside of the cavity (Fig. 1b, arrows), leading to the accumulation of *CO species inside the cavity and the concentration of the *CO species was significantly increased (Fig. 1b, colour map). The results lead to a high concentration of *CO intermediates required for the formation of C2 products, and facilitated the conversion to C_2 products (Fig. 1c).

Furthermore, the microenvironments over the interface of the fully closed and fully open structures were also monitored by FEM. The solid and fragment models were used to represent the fully closed and fully open structure catalyst immersed in an electrolyte. As *CO species tended to escape from the catalyst surface to the electrolyte, both the solid and fragment structures cannot prevent the diffusion of *CO from its surface to the electrolyte (Fig. S1a-f†). Therefore, these two structures were ineffective in enhancing the coverage of *CO species at active sites, resulting in no enhancement in the yield of C_{2+} products. The solid and fragment structures exhibited limitations in promoting C–C coupling. Fig. 1d shows coincident simulating results of the C_{2+}/C_1 ratios of the three structures. The C_{2+} -to- C_1 simulated ratios of cavity, solid and fragment structures were 7.81, 2.44, 1.73, respectively.

We also simulated the time-dependent variation of C_{2+} concentration on the cavity, solid and fragment structure models, and the results are shown in Fig. 1e. The C_{2+} concentration on the three models kept increasing with time, and the cavity model was obviously larger than that of the solid and fragment structure models at the same time. This distinction arose from the cavity structure, which significantly slowed down the diffusion kinetics, and improved the local coverage of *CO, thus facilitating the formation of C_{2+} products. In contrast, the diffusion of the *CO on solid and fragment structures cannot be retarded. These simulation results showed that cavity structures could promote the formation of C_{2+} products through steric nanoconfinement effects (Fig. 1f).

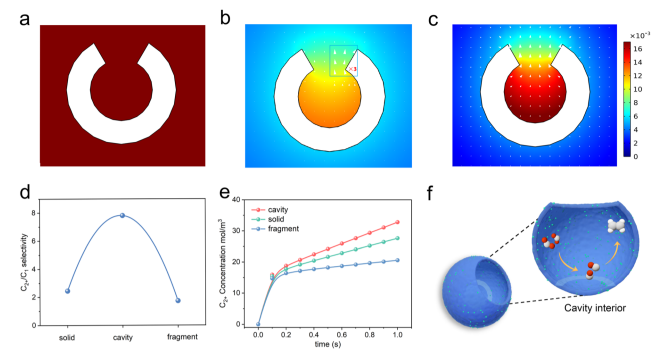


Fig. 1 Computed concentration and flux distribution of species. (a) CO_2 , (b) * CO_3 , and (c) CO_2 + concentrations (color scale, in mol CO_2 + concentrations) distributions (arrows) on the cavity structure. (d) Simulation results of the C_{2+}/C_1 product selectivity on the cavity, solid and fragment structure. (e) Simulation results of the time-dependent variation of C_{2+} concentration on cavity, solid and fragment structure. (f) The diagram displays how the cavity confinement effect promotes *CO intermediate dimerization and transformation to C₂H₄. Red, oxygen; grey, carbon; white, hydrogen.

Under the guidance of FEM simulation results, based on the synthesis mechanism of the self-template method,28 the carbonbased nanoreactor (Cu/C-cavity) was synthesized. The synthetic protocol of the Cu/C-cavity catalyst is illustrated in Fig. 2a. First, the template MET-5 was synthesized by a solvothermal method²⁹ (Fig. S2†), and then coated with a polydopamine (PDA) outer layer to obtain MET-5@PDA (Fig. S3†). Thermogravimetric analysis of MET-5 and MET-5@PDA was carried out (Fig. S4†). The removal rate of template MET-5 could be regulated by controlling the calcination time and heating rate, which results in different structures,30 and the Cu/C-cavity and Cu/C-fragment (fully open structures) catalysts could be obtained. The Cu/C-solid (fully closed structures) was obtained by carbonizing MET-5.

The scanning electron microscopy (SEM) images in Fig. 2b and S5† show the overall morphology of the Cu/C-cavity catalyst. Spherical particles with cavities were observed. In addition, transmission electron microscopy (TEM) images in Fig. 2c, S6 and S7† revealed the hollow morphology and open structures. The high-resolution TEM (HRTEM) image of the Cu/C-cavity catalyst displayed the lattice fringe of the crystal plane of Cu (111) (Fig. 2d). In addition, the SEM and TEM images confirmed the synthesis of Cu/C-solid and Cu/C-fragment catalysts (Fig. S8-S11†). The BET surface areas of Cu/C-cavity, Cu/C-solid, and Cu/ C-fragment are shown in Fig. S12.† The high-angle annular dark field (HAADF) image in Fig. 2e further confirms the hollow spherical structure formation of the Cu/C-cavity catalyst. The energy-dispersive X-ray spectroscopy (EDX) elemental mapping images showed uniform distribution of C, N, and Cu in the Cu/Ccavity catalyst.

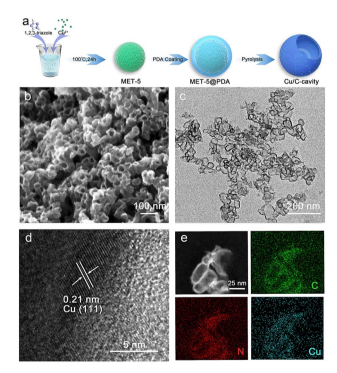


Fig. 2 Morphology characterization of the Cu/C-cavity catalyst. (a) The synthesis process of the Cu/C-cavity catalyst. Cu/C-cavity catalyst imaged by (b) SEM, (c) TEM, and (d) HRTEM. (e) High-angle annular dark field (HAADF) and mapping images of the Cu/C-cavity catalyst, showing the homogeneous distribution of C (green), N (red), and Cu (blue).

The chemical states and composition of the Cu/C-cavity, Cu/ C-solid and Cu/C-fragment catalysts were confirmed by a series of techniques. The powder X-ray diffraction (XRD) patterns results are presented in Fig. 3a, and the diffraction patterns at 43.2, 50.4, and 74.1 could be indexed to the Cu (111), Cu (200), and Cu (220) planes, respectively (PDF #04-0836). The XPS survey spectra illustrated the existence of Cu, O, N, and C elements in the three materials (Fig. S13a†). The Cu 2p_{3/2} peak and Cu 2p_{1/2} peak in the XPS spectra were observed at 932.3 eV and 952.2 eV (Fig. 3b), respectively. The Cu LMM Auger verified that the Cu species in the Cu/C-cavity, Cu/C-solid, and Cu/Cfragment catalysts existed in the form of Cu⁰ (918.3 eV, predominant) and Cu⁺ (914.7 eV). The minor amount of Cu⁺ may result from the partial oxidation of Cu during characterization. In addition, the N 1s peak could be deconvolved into pyridinic-N (398.48 eV), graphitic-N (401.4 eV), and oxidic-N (402.35 eV) peaks (Fig. S13b†).31 The C 1s spectrum could be fitted into three peaks, corresponding to C-C (284.6 eV), C-N (285.9 eV), and C-O (287.2 eV) (Fig. S13c†).32 The O 1s spectrum could be deconvolved into O-C (530.8 eV), O=C (532.1 eV), and O=C-O (533.5 eV) (Fig. S13d†).33

The structures of Cu/C-cavity, Cu/C-solid, and Cu/C-fragment catalysts were further evaluated by Raman analysis (Fig. 3c). The Raman spectra could be deconvoluted into four peaks by Gaussian–Lorentzian numerical simulation,³⁴ which were the graphene edges for the D1-band (*ca.* 1360 cm⁻¹), topological defects for the D3-band (*ca.* 1500 cm⁻¹), polyenes for the D4-band (*ca.* 1180 cm⁻¹) and graphitic lattice for the G-band (*ca.* 1580 cm⁻¹).³⁵ The graphitization degree of carbon was inferred by the ratio of integrated areas of the D1 and G.³⁶ The

calculated $I_{\rm D1}/I_{\rm G}$ ratios of the Cu/C-cavity, Cu/C-solid, and Cu/C-fragment catalysts were 1.65, 2.42, and 1.78, respectively, indicating that the Cu/C-cavity had a higher graphitization degree than Cu/C-solid and Cu/C-fragment catalysts.

The electronic information of the Cu/C-cavity, Cu/C-solid and Cu/C-fragment was further investigated by X-ray absorption spectroscopy (XAS) measurements. The Cu K-edge XANES spectra of the Cu/C-cavity, Cu/C-solid and Cu/C-fragment together with the references of Cu2O, CuO, and Cu foil are given in Fig. 3d. The X-ray absorption near edge structure (XANES) spectra (Fig. S14 and S15†) of Cu/C-cavity, Cu/C-solid and Cu/Cfragment were between the Cu foil and Cu2O, which proved that the Cu species was in the intermediate valence state between 0 and +1.37 The extended X-ray absorption fine structure (EXAFS) spectra (Fig. 3e) of Cu/C-cavity, Cu/C-solid and Cu/C-fragment displayed a main peak at 2.24 Å, corresponding to the metal Cu-Cu bond.³⁸ As shown in Fig. 3f, the Morlet Wavelet Transform (WT) of the k3-weighted extended X-ray absorption fine structure (EXAFS) further proved the existence of the Cu-Cu bond in the Cu/C-cavity, Cu/C-solid and Cu/C-fragment catalysts.39

The CO_2 electroreduction activity over the Cu/C-cavity, Cu/C-solid, and Cu/C-fragment catalysts was first investigated in 0.1 M CsI aqueous solution using a typical H type cell (Fig. S16†). As shown in Fig. 4a, the linear sweep voltammetry (LSV) curves over the Cu/C-cavity, Cu/C-solid, and Cu/C-fragment catalysts exhibited a higher reduction current density in the CO_2 -saturated electrolyte than the N_2 -saturated electrolyte, demonstrating their CO_2RR activity. Comparing these three catalysts, it was found that the Cu/C-cavity exhibited the highest reduction current density and the most positive onset potential

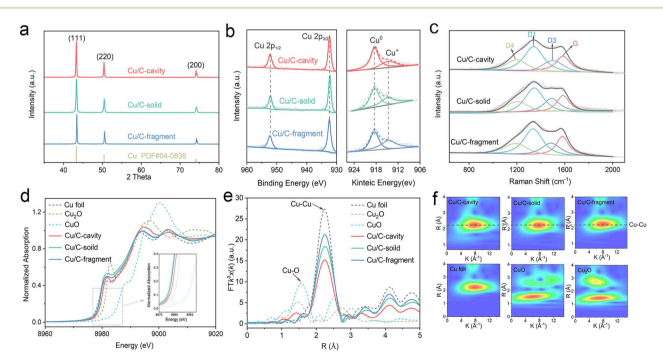


Fig. 3 Chemical structural characterization of different catalysts. (a) XRD, (b) XPS spectra of Cu 2p and Cu LMM Auger, (c) Raman spectra, (d) Cu K-edge XANES, (e) Fourier transformed Cu K-edge EXAFS spectra and (f) Morlet WT of the k3-weighted EXAFS of Cu/C-cavity, Cu/C-solid and Cu/C-fragment with the references.

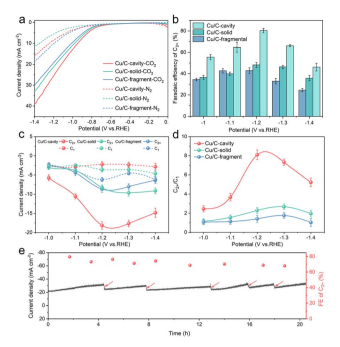


Fig. 4 CO_2 electrochemical reduction performance. (a) LSV curves on Cu/C-cavity, Cu/C-solid and Cu/C-fragment in CO_2 -saturated and N_2 -saturated 0.1 M CsI aqueous electrolyte. (b) C_{2+} FE from -1.0 V to -1.4 V vs. RHE. (c) C_{2+} and C_1 partial current densities at -1.0 V to -1.4 V vs. RHE in CO_2 saturated 0.1 M CsI aqueous electrolyte. (d) C_{2+} C_1 products selectivity ratio from -1.0 V to -1.4 V vs. RHE in CO_2 saturated 0.1 M CsI aqueous electrolyte. (e) Stability test of Cu/C-cavity at -1.2 V vs. RHE in CO_2 saturated 0.1 M CsI aqueous electrolyte, the arrows indicate the renewal of the electrolyte.

compared to Cu/C-solid, and Cu/C-fragment catalysts. This indicated that the Cu/C-cavity achieved a higher CO₂RR activity than the other two catalysts. The electrochemically active surface areas (ECSA) of the Cu/C-cavity, Cu/C-solid and Cu/Cfragment catalysts were obtained from the double-layer capacitance (C_{dl}) . In Fig. S17,† the Cu/C-cavity catalyst has the largest ECSA, which demonstrates that the Cu/C-cavity catalyst has the highest CO₂RR activity.⁴¹ Moreover, the Nyquist plots were recorded at the open-circuit potential to investigate the reaction kinetics of electrochemical processes. The electrochemical impedance spectroscopy (EIS) (Fig. S18†) shows that the Cu/Ccavity displayed the smallest Nyquist semicircle diameter compared to Cu/C-solid and Cu/C-fragment, suggesting a much faster interfacial charge-transfer kinetics.42 The local high ion concentration inside the cavity structure or the higher graphitization degree of Cu/C-cavity may improve the electrical conductivity, thus accelerating charge-transfer.43

According to the results in Fig. S19a,† the Cu/C-cavity catalyst demonstrated the production of C_{2+} products within the potential range of -1.0~V to -1.4~V νs . RHE. The results showed that the FE of C_{2+} products over the Cu/C-cavity catalyst could reach up to 80.5% (1446.17 ppm) at -1.2~V νs . RHE (Fig. 4b). This efficiency comprised 52.2% C_2H_4 , 18.8% C_2H_5OH , 5.4% CH₃COOH, and 4.2% n-PrOH (Fig. S20†). Fig. S21† shows the liquid products in D_2O (DMSO and phenol as an internal standard). Comparatively, the Cu/C-fragment and Cu/C-solid

catalysts tended to produce CO-dominated C1 products (Fig. 4b, S19b, c and S20†). The FE over C₂₊ products on Cu/C-cavity was much higher than that over the Cu/C-solid and Cu/C-fragment catalysts. The above experimental results proved that the steric confinement effect of the cavity structure could promote C2+ product selectivity. The partial current densities of C2+ and C1 products of Cu/C-cavity, Cu/C-solid, and Cu/C-fragment catalysts were compared at different applied potentials. As depicted in Fig. 4c, it was evident that the Cu/C-cavity catalyst exhibited the highest C_{2+} partial current density, reaching 18.14 mA cm⁻² at -1.2 V vs. RHE. In contrast, the Cu/C-solid and Cu/C-fragment showed lower maximum C2+ partial current densities of only 8.68 and 8.25 mA cm⁻², respectively. The $j_{C_{3}}$ normalised to ECSA and Cu content of Cu/C-cavity was larger than that of the Cu/C-solid and Cu/C-fragment catalysts (Tables S1 and S2†), indicating that the high C2+ products selectivity over the Cu/Ccavity resulted from its unique structure.

Fig. 4d discusses the ratios of C₂₊ to C₁ products over the Cu/ C-cavity, Cu/C-solid, and Cu/C-fragment catalysts. The Cu/Ccavity exhibited higher selectivity towards C2+ products across all tested potentials. Particularly at $-1.2 \text{ V} \nu s$. RHE, the Cu/Ccavity catalyst achieved maximum selectivity for C2+ products, with a $C_{2+}/C1$ ratio of approximately 8.1. The ratios over the Cu/ C-solid and Cu/C-fragment catalysts were 2.7 and 1.8, respectively. To further confirm the universality of our Cu/C-cavity structure, we investigated the electrochemical CO₂RR performance of Cu/C-cavity, Cu/C-solid, and Cu/C-fragment in 0.1 M KHCO₃ solution using a typical H type cell (Fig. S22†), and the FE of C₂₊ products over the Cu/C-cavity catalyst could reach up to 61.2% at -1.2 V vs. RHE, with a C_{2+}/C_1 selectivity ratio of approximately 6.14. The ratios of the Cu/C-solid and Cu/Cfragment catalysts were 2.06 and 1.31, respectively. The Cu/Ccavity also exhibited higher C₂₊ selectivity than Cu/C-solid and Cu/C-fragment in 0.1 M KHCO₃ solution. The experimental and simulated results showed excellent agreement, and it is also proved that the cavity structure is beneficial to improve the selectivity of C₂₊ products.

Furthermore, the electrochemical CO2RR performances of Cu/C-cavity, Cu/C-solid, and Cu/C-fragment were studied in 1 M KOH using a flow cell within the potential range of -0.8 V to −1.2 V vs. RHE (Fig. S23†). The Cu/C-cavity catalyst exhibited much higher selectivity for C₂₊ products (Fig. S24†), and the FE of C_{2+} could reach up to 75.2% (6990.76 ppm) with a C_{2+}/C_1 ratio of 3.57 at -1.0 V vs. RHE. The $j_{C_{2+}}$ could reach -323 mA cm⁻², which is higher than those of most electrodes and reached industrial levels. In addition, we also conducted the stability test in a flow cell, which indicated that the stability of the electrode was satisfactory (Fig. S24e†). Fig. S25† shows the liquid products over Cu/C-cavity in the flow cell. However, the Cu/C-solid and Cu/C-fragment tended to produce CH₄-dominated C_1 products. The ratios of C_{2+}/C_1 over the Cu/C-solid and Cu/C-fragment catalysts were 1.15 and 1.34, respectively. The Cu/C-cavity also exhibited much higher C₂₊ selectivity than Cu/ C-solid and Cu/C-fragment in the flow cell. Consequently, the above results proved that the confined geometric structure facilitates C-C coupling and the formation of multi-carbon compounds, which aligned with our previous hypothesis.

The stability of catalysts is a crucial parameter for the CO₂RR. To assess the stability of the Cu/C-cavity catalyst, CO₂ electrolysis was carried out at -1.2 V vs. RHE. As in Fig. 4e, the current density and FE did not change obviously over a period of 20 hours. The Cu/C-cavity nanoreactor was characterized by XRD and XPS techniques (Fig. S26†) after electrocatalysis showed no apparent change in the chemical state. The Cu Kedge spectra indicated that the Cu/C-cavity nanoreactor after electrocatalysis retained the characteristic feature of Cu(0), as can be further confirmed by the Cu-Cu coordination at 2.23 Å (Fig. S27†). From the TEM images (Fig. S28†) of the Cu/C-cavity after the CO₂RR, the nanocavity structure basically remained intact, and no obvious agglomeration of particles was found. We conducted a comparative analysis of our carbon-based nanoreactor materials with other reported nanoconfinement reactors in the literature for CO2 electroreduction. The result proved that carbon-based nanoreactors have superior stability over other reported pure Cu-based nanoreactors (Table S3†). The superior stability is mainly from the Cu and C structure of the catalyst, and the carbon carrier effectively stabilizes the active site of Cu and protects it from the reaction environment.

To further explore the effect of nanoconfinement on the CO₂RR, *in situ* Raman spectroscopy was conducted to monitor the key intermediates *CO of surface adsorption. Fig. 5a shows the *in situ* Raman spectra using the Cu/C-cavity catalyst during the CO₂RR under different applied potentials in 0.5 M KHCO₃ solution. The bands at 307 and 394 cm⁻¹ were attributed to the CO frustrated rotation and Cu–CO vibration stretching, respectively, indicating the adsorption of *CO.⁴⁴ The band at 524 cm⁻¹ was ascribed to the adsorption of preliminary intermediates (such as CO₂ ad) on the active sites.⁴⁵ The band at 984 cm⁻¹ was assigned to the *COO. Meanwhile, a much stronger band at 1067 cm⁻¹, corresponding to carbonate could be observed. The adsorption bands in the range of 2007–2058 cm⁻¹ were attributed to the atop-bound *CO (*CO_{atop}), which is a key intermediate of the CO–CO coupling.⁴⁶ The

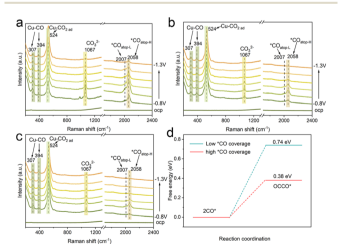


Fig. 5 Mechanistic studies. *In situ* Raman spectra of the (a) Cu/C-cavity, (b) Cu/C-solid, and (c) Cu/C-fragment catalysts during the CO2RR under different applied potentials. (d) The free energy of the *CO dimerization step at low (blue) and high *CO coverage (red) on Cu(111).

 $^*\mathrm{CO}_{\mathrm{atop}}$ peak could split into two bands. The low frequency band (LFB) at 2007 cm $^{-1\mathrm{is}}$ ascribed to $^*\mathrm{CO}_{\mathrm{atop}}$ on the terrace ($^*\mathrm{CO}_{\mathrm{atop}}$ –L), and the high frequency band (HFB) at 2058 cm $^{-1}$ is associated with $^*\mathrm{CO}_{\mathrm{atop}}$ on the low coordinated sites ($^*\mathrm{CO}_{\mathrm{atop}}$ –H). The $^*\mathrm{CO}_{\mathrm{atop}}$ peak intensity increased and then decreased with the potential scanned, indicating that $^*\mathrm{CO}$ intermediates were accumulated and then consumed.

Furthermore, the *in situ* Raman spectra of Cu/C-solid and Cu/C-fragment were also recorded (Fig. 5b and c). The intensities of the peaks for linearly adsorbed *CO_{atop} on the Cu/C-solid and Cu/C-fragment were weak, implying less accumulation of *CO. In comparison, the peak intensity of *CO_{atop} (2058 cm⁻¹) on the Cu/C-cavity was stronger than that over Cu/C-solid and Cu/C-fragment. The results suggested that *CO_{atop} intermediates were accumulated on the Cu/C-cavity, indicating that the cavity structure could enrich the local concentration of *CO intermediates.⁴⁷ Overall, the *in situ* Raman results revealed that the cavity structure exhibited a higher coverage of *CO_{atop} compared to the solid and fragment structure, thus accelerating the process of *CO dimerization, resulting in superior CO₂RR selectivity of C₂₊ products.

To further verify that the local high concentration of *CO could promote the rate of C–C coupling, the Gibbs free energy of the *CO dimerization step⁴⁸ was studied using DFT (Fig. 5d, S29 and S30†). The Gibbs free energy value for the C–C coupling of *CO was found to be 0.74 eV at low *CO coverage, and decreased to 0.38 eV at high *CO coverage. The results suggested that a lower energy barrier for the C–C coupling reaction appeared at high *CO coverage. Thus, the probability of C–C coupling could be promoted by a high *CO coverage, which is consistent with our experimental results.

Conclusions

In summary, we discussed the correlation between the geometric structures of catalysts and the selectivity of C2+ products. We have demonstrated that cavity nanoreactors showed a significantly improved electrochemical CO2RR performance. The Cu/C-cavity exhibited a high C2+ FE of 80.5% and C₂₊/C₁ selectivity ratio of 8.1, which is much higher than that over the Cu/C-solid and Cu/C-fragment. In addition, due to the highly dispersed Cu on the carbon support, the stability of Cu/C-cavity was better than that of most reported pure copperbased nanoreactor catalysts. The results of FEM simulation, in situ Raman and DFT supported that the cavity of nanoreactors enriched the local concentration of *CO intermediates, thus promoting the C-C coupling reactions. This research underscored the potential application of functionalized nanoreactors in the highly selective electrosynthesis of valuable fuels from CO₂, while shedding light on the morphology and composition of catalysts having a significant effect on the performance of the CO₂RR.

Data availability

The authors declare that all data supporting the findings of this study are available within the paper [and its ESI†].

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Author contributions

M. W., W. X., H. H. W. and B. X. H. proposed the project, designed the experiments, and wrote the manuscript. M. W. performed the whole experiment. S. Q. J., T. Y., X. D., D. W. Z., and M. H. F. performed the analysis of experimental data. W. X., H. H. W., M. Y. H. and B. X. H. co-supervised the whole project. All authors discussed the results and commented on the manuscript.

Conflicts of interest

The authors declare no competing interests.

Acknowledgements

The work was supported by the National Key Research and Development Program of China (2020YFA0710201; 2023YFA1507901), the China Postdoctoral Science Foundation Funded Project (2021M701211), and the National Natural Science Foundation of China (22293015, 21890761, and 22121002), and Fundamental Research Funds for the Central Universities. The X-ray absorption fine structure (XAFS) measurements were carried out at the 4B9A beamline of the Beijing Synchrotron Radiation Facility (BSRF).

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