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Cu-Ag tandem catalysts for high-rate CO₂ electrolysis towards multicarbons

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Summary: Electrochemically upgrading CO₂ to carbon-neutral multicarbons (C_{2+}) is a promising technology for CO₂ recycling and utilization. Since such transformations involve multiple elementary steps, a tandem strategy becomes attractive as catalysts can be optimized for specific reaction steps independently. Related strategies have been demonstrated under low working current densities; however, the applicability of a tandem strategy towards high-rate CO₂ electrolysis to C_{2+} is unknown. Here, we demonstrate that a Cu-Ag tandem catalyst can enhance the multicarbon production rate in CO₂RR by decoupling high-rate CO₂ reduction to CO on Ag and subsequent CO coupling on Cu. With the addition of Ag, the partial current towards C_{2+} over a Cu surface increased from 37 mA/cm² to 160 mA/cm² at -0.70 V vs RHE in 1M KOH while no mutual interference between two metals was observed. Moreover, the normalized intrinsic activity of C_2H_4 and C_2H_5OH in the tandem platform under CO₂ reduction conditions is significantly higher than Cu alone under either pure CO₂ or CO atmosphere. Our results indicate that the CO-enriched local environment generated by Ag can enhance C_{2+} formation on Cu beyond CO₂ or CO feeding, suggesting possible new mechanisms in a tandem three-phase environment.

1 1. Introduction

2 CO₂ electrolysis is a promising technique for CO₂ recycling and utilization¹⁻², offering carbon-neutral chemical feedstocks³ for 3 downstream applications. Electrochemical CO₂RR towards single carbon products have achieved enormous progress⁴, especially 4 for carbon monoxide⁵⁻⁹. However, in the periodic table, copper (Cu) is the only metal that can efficiently catalyze CO₂ to multicar-5 bon products (C₂H₄, C₂H₅OH, CH₃COOH, n-C₃H₇OH, etc.), which was firstly reported by Hori and co-workers¹⁰. Since multicarbon 6 products are higher-value and energy concentrated^{1, 3}, widespread attention is paid to the unique catalytic property of the Cu 7 surface. Both theoretical¹¹ and experimental¹² works demonstrate that sluggish C-C coupling kinetics over a pure Cu surface im-8 pedes industrial level production of multicarbon products. Thus, various multicomponent catalyst design principles have been 9 utilized to improve the catalytic performance of Cu-containing catalysts, such as alloy formation¹³⁻¹⁵, surface doping¹⁶⁻¹⁷, ligand 10 modification¹⁸⁻²⁰, interface engineering²¹⁻²⁴.

11 One attractive multicomponent strategy is to break down the multiple reaction steps of CO₂-to-C₂₊ to have optimized compo-12 nents perform each step. In recent years, this "tandem catalysis" strategy has been successfully demonstrated in thermal heterogeneous catalysis²⁵⁻²⁷. Since it has been widely accepted that *CO (* denotes surface adsorbed species) and CO related intermedi-13 ates are critical for CO₂RR towards multicarbons^{4, 28-29}, the tandem strategy is also suitable for CO₂ electrolysis. Due to excellent 14 CO formation ability over Ag and Au, Cu-based bimetallic tandem platforms have been invoked for CO₂RR³⁰⁻³⁴. However, the prac-15 16 ticality of such a strategy at high production rates (*i.e.* > 200 mA/cm²) is still in question, due to CO₂ mass transport limitations in 17 previous works. The gas diffusion electrode (GDE) can eliminate sluggish feed gas transportation kinetics by constructing an efficient three phase interface³⁵⁻³⁶. Nevertheless, the micro-environment around the catalyst may be considerably different during 18 19 high-rate electrolysis compared with aqueous conditions³⁷. Thus, to establish decoupled electrocatalytic CO₂-to-C₂₊ paradigms in 20 high-rate electrochemical environments, the efficacy of a tandem platform should be evaluated in a gas diffusion electrode flow 21 cell.

22 Here, in order to evaluate whether a tandem strategy can be utilized in high-rate CO₂ reduction, we designed a model two-23 component catalytic architecture in alkaline electrolyte with commercial Cu and Ag powder (Fig. 1a). Ag is selected for its high 24 activity for CO formation during CO₂RR³⁸ and its thermodynamically immiscible nature with Cu at room temperature, thereby de-25 convolving enhancement due to bulk elemental alloying³⁹. Although 1M KOH electrolyte is ultimately unsustainable for long term 26 practical electrolysis due to electrolyte instability from reaction with CO₂, it is useful as a model environment to evaluate catalytic 27 concepts at the early stage³⁷. Compared against Cu alone, a significant enhancement of the partial current towards C_{2+} products 28 (C₂H₄, C₂H₅OH, CH₃COOH) is observed over the tandem catalyst during CO₂ electrolysis, with a concurrent selectivity shift towards oxygenates over C₂H₄. Both post-electrolysis characterization and electrochemical CORR results confirm that the active sites over 29 30 Ag and Cu surface work independently during CO₂RR, supporting the importance of a CO-rich local environment generated by the 31 Ag component for enhancing C-C coupling on the Cu component in accordance with the tandem strategy. Furthermore, the nor-32 malized intrinsic activity towards C2+ products indicate that the Ag-generated microenvironment in tandem CO2RR is superior to 33 both pure CO₂ and CO atmosphere for Cu-catalyzed multicarbon production. This phenomenon suggests the existence of mecha-34 nisms for CO2 reduction enhancement specifically stemming from tandem strategies and their resulting chemical microenviron-35 ments.

36 2. Results

37 2.1 Electrode characterization. A Cu-Ag tandem architecture catalyst was prepared by physical mixture of Cu and Ag na-38 nopowders on carbon paper support and utilized in gas diffusion flow cells (Fig. S1) for high current CO₂ electrolysis. The 39 morphology of commercial Cu and Ag powders is revealed with TEM (Fig. S2), showing that the crystalline feature size of both Cu 40 and Ag is tens of nanometers, but heavily agglomerated with no distinct morphology. The electrode fabricated with these metal 41 powders was name as Cu_xAg_y, representing a Cu-Ag tandem catalyst with x µg/cm² Cu and y µg/cm² Ag respectively (see experi-42 mental procedures). Then, these electrodes were imaged under SEM. The thickness of the catalyst layer in the as-prepared 43 Cu₅₀₀Ag₁₀₀₀ catalyst on carbon paper is several microns (Fig. 1b, 1c). SEM after electrolysis shows no significant structural change 44 on the micron scale from the initial structure (Fig. S3).

45 Elemental characterization through XRD, XPS and EDS mapping (Fig. 1d, 1e, S4) provide further insight into the nature of the 46 tandem catalyst. Both XRD patterns and XPS spectra of the as-prepared tandem Cu₅₀₀Ag₁₀₀₀ catalyst show characteristic Cu and 47 Ag related peaks without any observable peak shifting. Although the Cu LMM Auger peak indicates that the surface of nano pow-48 der has been oxidized, numerous studies⁴⁰⁻⁴⁴ indicate that such a naturally formed oxide shell can be quickly reduced under work-49 ing condition. Despite the ongoing debate in the community towards the existence and function of oxygen residue during CO₂RR⁴⁵⁻⁴⁷, we note that such a phenomenon will not influence the conclusions related to tandem interactions between Ag and Cu 50 51 of the later discussion in this work. Meanwhile, it is widely reported that Cu can undergo surface reconstruction and structural evolution during CO₂ electrolysis,⁴⁸⁻⁵¹ and previous reports have also invoked Cu-Ag alloy¹³⁻¹⁵ or surface alloy¹⁶ strategies to im-52

53 prove CO₂RR catalytic performance. Thus, we conducted post-electrolysis characterization of the tandem Cu₅₀₀Ag₁₀₀₀ catalyst to 54 determine whether the structure is maintained. Previous works have shown structural and electronic differences owing to strong 55 Ag interactions with Cu: for example, up to 0.8° Cu(111) peak shift in XRD could be found for a Cu-Ag alloy system¹⁴ whereas up to 0.3 eV Cu 2p_{3/2} peak shift in XPS was reported for a Cu-Ag dimer.²³ In contrast, no peak shift of Cu or Ag could be observed for the 56 57 tandem Cu₅₀₀Ag₁₀₀₀ catalyst in either XRD, XPS or Cu LMM Auger peak after electrolysis, indicating the structural maintenance of 58 this tandem catalyst and absence of electronic interactions between Ag and Cu throughout electrolysis. This absence is likely due 59 to the bulk-like nature of Ag and Cu used, in addition to the mild conditions in which the electrode is fabricated, resulting in ther-60 modynamically favored separation⁵². Importantly, this does not preclude the Ag-Cu surface and interfacial alloying observed in 61 other reports which use more energetic fabrication conditions.

62 2.2 Enhanced CO₂RR catalytic performances toward C₂₊ products over tandem Cu-Ag catalysts. The polarization response 63 curve of Cu₅₀₀Ag₁₀₀₀ in Fig.2a shows higher geometric current density than Cu₅₀₀ or Ag₁₀₀₀ alone under the same potentials. Inter-64 estingly, partial current densities toward C_{2+} products over different catalysts are also observed to be substantially higher for 65 Cu₅₀₀Ag₁₀₀₀, which cannot be explained simply through the individual contributions of Cu₅₀₀ and Ag₁₀₀₀ (Fig. 2b). Explicitly, Ag₁₀₀₀ 66 does not contribute to C-C coupling reactions in the potential range from -0.5 V to -0.8 V vs RHE. Thus, all partial current toward 67 C2+ products should come from the Cu surface. Based on linear interpolation, the partial current toward C2+ products over Cu500 68 surface is only about 37 mA/cm² at -0.70V, while the corresponding C₂₊ partial current density over Cu₅₀₀Ag₁₀₀₀ is 160 mA/cm². As 69 the theoretical Cu loading density is constant between Cu_{500} and $Cu_{500}Ag_{1000}$, the significant enhancement of the C_{2+} partial current 70 density over the tandem catalyst beginning at around -0.55 V vs RHE should be attributed to the additionally loaded Ag. The en-71 hancement of C₂₊ products is shown to be dependent on the extent of Ag loading (Fig. S5), further reinforcing that Ag is responsi-72 ble for the increase in C_{2+} production.

73 C₂₊ partial currents are further broken down to their main constituents to analyze how the tandem strategy influences product 74 distribution (Fig.2c). Here we define the enhancement factor (EF) as the tandem $Cu_{500}Ag_{1000}$ partial current divided by that of the 75 Cu₅₀₀ catalyst alone. Beginning from -0.55 V vs RHE, the EFs of C₂H₄, C₂H₅OH and CH₃COO⁻ all increase significantly when the ap-76 plied potential is more negative (Fig. 2c). Thus, relatively negative potential is essential to trigger the Ag-derived C2+ enhance-77 ment, achieving up to 6.43, 11.27, and 16.22 µg/h·cm²geom production rates toward C₂H₄, C₂H₅OH and CH₃COO⁻ respectively over 78 the Cu surface. Furthermore, oxygenate products are favored over C_2H_4 on the tandem catalyst, consistent with previous reports 79 that invoke enhancement by local CO concentration ^{31, 51}. At -0.70 V vs RHE, the EFs are mostly at the same range as -0.65 V vs 80 RHE, indicating that the tandem strategy induced enhancement is limited by factors beyond potential, such as the concentration 81 of CO local to the catalyst. Similar trends can also be observed from tandem catalyst with other Cu-Ag ratios (Fig. S6).

82 To further probe the role of local CO towards the C_{2+} enhancement, the net CO producing abilities over these catalysts are also 83 compared (Fig.2d). At potentials negative of -0.5 V vs. RHE, the C-C coupling reaction dominates such that net CO production rate 84 over Cu₅₀₀ remains at a relatively low level within the potential range. Meanwhile, Ag exhibits an exponential increase of CO pro-85 duction with respect to overpotential. Notably, the onset potential for C_{2+} enhancement is around -0.55 V vs RHE, where CO par-86 tial current on the pure Ag₁₀₀₀ catalyst is observed to take off. This indicates a direct correlation between C₂₊ enhancement on the 87 Cu₅₀₀Ag₁₀₀₀ catalyst and net CO production rate. Hence, the partial C₂₊ current enhancement is attributed to the high local CO cre-88 ated on the Ag surface within the tandem catalyst layer. Interestingly, Clark et al. invoked that the enhanced oxygenates selectivi-89 ty is not attributed to localized CO produced by Ag, but by strain effects induced by surface Ag doping which modulates C-C coupling activity and suppresses HER.¹⁶ Similarly, such HER suppression and oxygenates favoring phenomenon were also observed in 90 91 other Cu-Ag alloy catalysts¹⁴⁻¹⁵. The differing contributions of tandem effects on catalysis (*i.e.* whether localized CO may influence 92 further C_{2+} activity) between this work and previous works mentioned above may be due to the difference in electrolyte. It has 93 been reported that high pH electrolyte can significantly accelerate C-C coupling reaction over Cu surface³⁶, which would corre-94 spond to more efficient utilization of CO in our system. Notably, the correlated suppression of HER typically observed from Ag-95 alloying or doping is not found in our system (Fig. S7), suggesting that the Ag doping effect is mostly negligible here and that the 96 intrinsic catalytic activity of Cu is not influenced by Ag atomic interactions. More contextualization and comparison of relevant 97 works can be found in Fig. S8 and Table S1.

98 2.3 Active site analysis and performance comparison based on CORR. Considering the importance of localized CO produced by 99 the Cu-Ag tandem platform toward high rate C2+ chemicals production, we next investigated how such platforms behave under 100 CO reduction reaction (CORR) conditions. If the addition of Ag electronically influences Cu active sites, we expect that CO reduc-101 tion activity should differ between Cu and Cu-Ag catalysts⁵³. However, in stark contrast to our previous observations in CO₂RR, the 102 Cu₅₀₀Ag₁₀₀₀ catalyst exhibits neither an enhancement in total current nor an enhancement in C₂₊ formation under CO-reducing conditions (Fig. 3a, 3b, S9). Thus, in both Cu₅₀₀Ag₁₀₀₀ and Cu₅₀₀ catalysts, only the Cu component conducts CORR. Meanwhile, no 103 104 substantial enhancement is observed owing to the bimetallic platform in CORR across all multicarbon products, as shown explicit-105 ly in Fig. 3c, suggesting that the Cu₅₀₀Ag₁₀₀₀ catalyst has no intrinsic enhancement in CO coupling compared with Cu₅₀₀ alone.

106 In general, our CORR experiments strongly support the conclusion that Cu and Ag active sites operate independently. Ag pro-107 duces CO from CO₂ and this generated CO in addition to the CO generated on Cu itself, is further utilized for multicarbon produc-108 tion over the Cu surface. In other words, we posit that in accordance with our structural characterization, Ag does not modify the 109 Cu surface during tandem CO₂RR. We note that although electrocatalytic activity changes over longer periods of electrolysis, this 110 is due to commonly reported stability issues with flooding in carbon paper based gas diffusion electrodes^{36, 54-56}, rather than to 111 structural changes of the catalyst (Fig. S10). Moreover, given the similar activity between Cu and Cu-Ag catalysts in CORR when 112 catalyst preparation is simply normalized by Cu mass loading, we further hypothesize that intrinsic catalytic activity towards mul-113 ticarbons can be internally compared using Cu mass normalization.

114 2.4 Intrinsic catalytic performances of Cu catalyst under different local environments. Considering the production rate nor-115 malized by the mass of Cu as a measure of intrinsic catalyst activity for internal comparison, we sought to determine how the 116 local gas environments would influence C₂₊ partial current over Cu surface (Fig. S11). Unsurprisingly, for CORR the activity towards 117 all three main C₂₊ products over Cu₅₀₀ are larger than for CO₂RR. However, we also observed unexpectedly that the normalized C₂₊ 118 partial current over tandem Cu₅₀₀Ag₁₀₀₀ during CO₂RR is beyond that of Cu₅₀₀ during CORR. To exclude the influence of electron 119 transfer differences in CO₂RR and CORR (e.g. for C₂H₄, 12e⁻ during CO2RR and 8e⁻ during CORR), we used the normalized intrinsic 120 activity (in moles product per mass catalyst per time) to better represent the intrinsic performance over the Cu surface (Fig.4a-c). 121 The normalized intrinsic activities of tandem Cu₅₀₀Ag₁₀₀₀ towards C₂H₄, C₂H₅OH, CH3COO⁻ in CO₂RR are 3~6-fold larger than that of 122 Cu₅₀₀ in CO₂RR at -0.7 V vs RHE. Moreover, the normalized intrinsic activities of tandem CO₂RR towards C₂H₄ and C₂H₅OH are still 123 substantially larger than that of Cu₅₀₀ in CORR(Fig.4d), such that the local environment created by the tandem catalyst favors a 124 greater overall formation for C_{2+} than CORR on Cu alone. Given that the role of Ag is to supply Cu with a CO-rich local environment, 125 we considered how a similar amount of Cu in place of Ag would behave. The normalized catalytic activity over Cu₅₀₀Cu₁₀₀₀ is con-126 siderably lower than that of $Cu_{500}Ag_{1000}$, likely due to the deficient CO_2 -to-CO activity of a Cu surface per surface site. (Fig. S12). 127 Such results highlight the unexpected superiority of the tandem strategy for CO₂RR in a flow cell, which offers the opportunity to 128 enhance the efficiency of C-C bonding on a Cu surface beyond conventional CORR during high-rate electrolysis.

129 The underlying mechanism accounting for this change in product distribution is also of interest. We find that the tandem-130 derived C2+ intrinsic activity enhancements over a Cu500 surface can also be imitated with a blended gas of 2% CO and 98% CO2 (Fig. 131 S13). Therefore, a mechanistic difference in mixed CO₂-CO atmosphere in the tandem catalyst layer may contribute to the en-132 hancement of intrinsic activity towards a series of multicarbon products, similar to Strasser and co-workers' results in bicarbonate 133 at lower electrolysis rates⁵⁷. Other factors in the double layer may also be involved. For example, a relatively low CO coverage 134 over Cu surface (2.5%~10% atm) can promote C₂H₄ formation rate during CORR⁵⁸ while *H coverage would influence C₂H₅OH for-135 mation during CO₂RR⁵⁹ by adsorbate-adsorbate interaction. Furthermore, the surface pH may also be an important influencing 136 factor, due to subtle differences in the double layer given the presence of carbonate formation during CO₂RR³⁶. With many possi-137 ble mechanisms for the enhancement observed under tandem Cu-Ag CO₂RR conditions, Cu catalysts with well-defined surfaces as 138 well as advanced in-situ spectroscopy techniques for flow cells will be valuable future mechanistic tools.

139 **3.** Discussion

140 In this work, we established and evaluated a Cu-Ag tandem platform for achieving high current CO₂ electrolysis towards C₂₊ prod-141 ucts in a flow cell. In contrast with previous Cu-Ag works, Cu and Ag were confirmed to work independently in this system 142 through post-electrolysis characterization and control experiments with CO reduction, such that the localized CO produced by Ag, 143 with subsequent C-C bonding on Cu, is key towards high rate C_{2+} production. Furthermore, we found over the same Cu surface, 144 the turnover of C_2H_4 and C_2H_5OH in a tandem CO₂-reducing platform is substantially higher than the activity for Cu in either pure 145 CO_2 or pure CO-reducing conditions. Thus, we posit that the local environment created in high-rate tandem CO_2 electrolysis offers 146 new opportunities for expanding the catalytic space of CO₂ electroreduction. Deep investigation of local CO atmosphere in situ is 147 required to further understand high-rate tandem CO₂ electrolysis.

Moreover, we emphasize that although a highly alkaline electrolyte is beneficial for new catalytic architecture exploration and demonstration, serious concerns remain regarding its sustainability and stability issues due to the reaction between KOH and CO- $^{2^{37}}$. This may partially account for the lack of long-term stability observed in this work (Fig. S10). Translating the catalytic concept demonstrated here to truly practical CO₂ electrolysis requires systematic design that reduces CO₂ interaction with alkaline electrolyte while ideally still leveraging the catalytic benefits of local hydroxide. One avenue of interest may be to substitute bulk KOH electrolyte with local metal hydroxide at the cathode. Further exploration of catalyst design with cell engineering as a multi-scale strategy in sustainable electrolytes will undoubtedly be needed to enable practical tandem CO₂ electrolysis systems.

Experimental procedures

Resource Availability

Lead Contact

Further information and reasonable requests for resources and materials should be directed to and fulfilled by the Lead Contact, Peidong Yang (p_yang@berkeley.edu)

Materials Availability

All chemicals were purchased from commercial resources and used as received.

Data and Code Availability

There is no dataset or code associated with this work.

Materials

Commercial Cu powder (40-60 nm particle size (SAXS)), commercial Ag powder (<100 nm particle size), deuterium oxide (D₂O, 99.9 atom % D), dimethyl sulfoxide (DMSO, ACS reagent, \geq 99.9%) and Nafion 117 ionomer solution (~5% in water and alcohol mixture) was purchased from Sigma-Aldrich. KOH pellets (>85%) were purchased from Fisher Chemicals. Anion exchange membrane (Fumasep FAA-3-PK-130) and Sigracet carbon papers were ordered from Fuel Cell Store. Ag/AgCl (3M KCl) reference electrodes were purchased from CH Instrument. Platinum gauze (99.9% metal basis) was purchased from Alfa Aesar. Carbon dioxide (CO2, 4.5LS), carbon monoxide (4.0 research) and Argon gas were purchased from Praxair. Deionized water (DI water with 18.2 M Ω ·cm, <5 ppb TOC) was obtained from a Millipore water system.

Electrode preparations

The following procedures describe the fabrication of a $Cu_{500}Ag_{1000}$ catalyst, with other catalysts produced in a similar fashion and adjusting for mass. 12 mg commercial Cu or Ag nano-powder was firstly mixed with 4 ml ethanol, 40 µl Nafion ionomer solution and then sonicated for 1h in the water bath to produce a uniform ink. Cu and Ag inks were mixed together with 1:2 volume ratio and then spray-cast on carbon paper with a spray gun (SPEEDAIRE, Mod. 48PX90). After that, the fabricated electrode was stored in a vacuum desiccator overnight to remove residual ethanol. $Cu_{500}Ag_{1000}$ represents a Cu-Ag tandem catalyst with 500 µg/cm² Cu and 1000 µg/cm² Ag theoretically. The weight difference of carbon paper before and after spraying was measured by an analytical balance to confirm the loading amount.

Characterization

The image of Cu and Ag powder morphology was collected by a transmission electron microscopy (Tecnai G2 T20 S-TWIN) under 200 kV. Scanning Electron Microscopy (Ultra 55-FESEM) is operated at 5 keV with 5 mm working distance to characterize the fabricated electrode morphology and elemental distribution before and after electrolysis. XRD (Bruker D8) patterns were collected using a Cu target. X-ray photoelectron spectroscopy (XPS) (Thermo Scientific K-Alpha) measurement was conducted using an Al K α source. For all post-electrolysis samples, electrodes were taken after CO₂ electrolysis at 200 mA/cm² constant current density for 30 min, then rinsed with deionized water and isopropanol repeatedly in order to remove as much electrolyte residue as possible and dried in a vacuum desiccator before measurement.

Electrochemical experiments and products characterization

All CO_2 and CO involved electrolysis were conducted in a gas diffusion flow cell. Before electrolysis, electrolyte was firstly bubbled with argon for at least 30 min to remove dissolved oxygen. In this work, 1 M KOH was used as the electrolyte. The gas flow rate during electrolysis was controlled by a mass flow controller at 25 sccm while the electrolyte flow rates in cathode and anode channels were managed by two low-flow chemical metering pumps at 5 ml/min. The flow-out electrolyte would be directly collected for product analysis or transferred to waste bottle instead of feeding back to the cell in order to eliminate pH drop induced by neutralization reaction between KOH and CO_2 . During all electrolysis, the cathode and anode were separated by a Fumasep anion exchange membrane. Ag/AgCl (3M KCl) was used as reference electrode. Considering batch to batch difference and harsh working conditions (high current and high pH), the Ag/AgCl reference electrode was frequently calibrated and replaced if needed.

During chronopotentiometry (CP), both gas and liquid products were collected and analyzed. Sampling was conducted at least 300 s after the electrolysis began in order to let the system reach steady state. No automatic iR correction was applied during CP experiments. After product collection, the cathodic potential and resistance between working electrode and reference electrode

were precisely measured by a subsequent current interrupt (CI) experiment. The back-calculated potentials were based on the following equation:

 $E_{vs RHE} = E_{vs Ag/AgCl} + iR + E_{Ag/AgCl vs RHE}$

The RHE potential is calculated using a pH value of 14. After electrolysis, gas products, including CO, CH₄, C₂H₄ and H₂ were detected and quantified using gas chromatography (Agilent 7890B GC system). Three GC columns (1.5 ft HayeSep Q, 6 ft HayeSep Q and 6 ft Molsieve 5A) are used to separate different gases. CO and CO₂ were additionally transformed into CH₄ with a methanizer. H₂ was detected by TCD (Ar flow) while other gases were detected by FID. Liquid products were analyzed by 1H nuclear magnetic resonance (Bruker AV600 with a 5 mm Z-gradient triple resonance 1H/BB Prodigy cryo-probe) with water suppression by excitation sculpting. DMSO was used as an internal standard for quantitative NMR. Through the calibration curve, the concentration of the products can be determined (unit: mM for liquid products and ppm for gas products). Partial current density of the products can be calculated with the following equations:

 $j_{gas product} = c_{gas product} \times 10^{-5} \times e^{-1}$ transfer number $\times F \times v_{gas} / (V_m \times S)$

and

 $j_{\text{liquid product}} = c_{\text{liquid product}} \times e^{-transfer number} \times F \times v_{\text{electrolyte}} / S$

where:

j: partial current density (mA/cm²)

c: concentration of the product in gas phase (ppm) or electrolyte (M)

F: Faraday constant (96485 C/mol)

v: flow rate (mL/s)

V_m: molar volume at room temperature and pressure (293 K, 1 atm)

S: electrode geometric area (cm²)

Subsequently, faraday efficiency (FE) can be obtained with:

 $FE = j_{product} / j_{total} \times 100\%$

Specifically, in this work, enhancement factor (EF) at specific potentials was defined as:

EF = jproduct on tandem catalyst / jproduct on single Cu catalyst

And normalized intrinsic activity towards multicarbons was calculated with following equation: normalized intrinsic activity = $j_{product}$ / (F × e⁻ transfer number × d_{cu})

where:

d_{Cu}: Cu loading density (mg/cm²)

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Author Contributions

C.C. designed the systems, collected and analyzed data, and wrote the manuscript. Y.L., J.J and M.L. analyzed data and wrote the manuscript. S.Y. and S.L. collected data. M.B.R. contributed to system setup. P.Y. designed the systems, analyzed data, and wrote the manuscript.

Declaration of Interests

The authors declare no competing financial interest.

References

(1) Kibria, M. G.; Edwards, J. P.; Gabardo, C. M.; Dinh, C. T.; Seifitokaldani, A.; Sinton, D. and Sargent, E. H., (2019). Electrochemical CO2 Reduction into Chemical Feedstocks: From Mechanistic Electrocatalysis Models to System Design. Adv. Mater. *31*, 1807166.

(2) Seh, Z. W.; Kibsgaard, J.; Dickens, C. F.; Chorkendorff, I.; Norskov, J. K. and Jaramillo, T. F., (2017). Combining theory and experiment in electrocatalysis: Insights into materials design. Science 355, eaad4998.

(3) Bushuyev, O. S.; De Luna, P.; Dinh, C. T.; Tao, L.; Saur, G.; van de Lagemaat, J.; Kelley, S. O. and Sargent, E. H., (2018). What Should We Make with CO 2 and How Can We Make It? Joule 2, 825-832.

(4) Ross, M. B.; De Luna, P.; Li, Y.; Dinh, C.-T.; Kim, D.; Yang, P. and Sargent, E. H., (2019). Designing materials for electrochemical carbon dioxide recycling. Nat. Catal. 2, 648-658.

(5) Ross, M. B.; Li, Y.; De Luna, P.; Kim, D.; Sargent, E. H. and Yang, P., (2019). Electrocatalytic Rate Alignment Enhances Syngas Generation. Joule 3, 257-264.

(6) Li, T.; Lees, E. W.; Goldman, M.; Salvatore, D. A.; Weekes, D. M. and Berlinguette, C. P., (2019). Electrolytic Conversion of Bicarbonate into CO in a Flow Cell. Joule 3, 1487-1497.

(7) Ren, S.; Joulié, D.; Salvatore, D.; Torbensen, K.; Wang, M.; Robert, M. and Berlinguette, C. P., (2019). Molecular electrocatalysts can mediate fast, selective CO2 reduction in a flow cell. Science 365, 367-369.

(8) Gu, J.; Hsu, C.-S.; Bai, L.; Chen, H. M. and Hu, X., (2019). Atomically dispersed Fe3+ sites catalyze efficient CO2 electroreduction to CO. Science 364, 1091-1094.

(9) Liu, M.; Pang, Y.; Zhang, B.; De Luna, P.; Voznyy, O.; Xu, J.; Zheng, X.; Dinh, C. T.; Fan, F.; Cao, C.; de Arquer, F. P.; Safaei, T. S.; Mepham, A.; Klinkova, A.; Kumacheva, E.; Filleter, T.; Sinton, D.; Kelley, S. O. and Sargent, E. H., (2016). Enhanced electrocatalytic CO2 reduction via field-induced reagent concentration. Nature *537*, 382-386.

(10) Hori, Y.; Murata, A. and Takahashi, R., (1989). Formation of hydrocarbons in the electrochemical reduction of carbon dioxide at a copper electrode in aqueous solution. J. Chem. Soc., Faraday Trans. 1 85, 2309-2326.

(11) Garza, A. J.; Bell, A. T. and Head-Gordon, M., (2018). Mechanism of CO2 Reduction at Copper Surfaces: Pathways to C2 Products. ACS catal. 8, 1490-1499.

(12) Ma, M.; Djanashvili, K. and Smith, W. A., (2016). Controllable hydrocarbon formation from the electrochemical reduction of CO2 over Cu nanowire arrays. Angew. Chem. Int. Ed. 55, 6680-6684.

(13) Hoang, T. T. H.; Verma, S.; Ma, S.; Fister, T. T.; Timoshenko, J.; Frenkel, A. I.; Kenis, P. J. A. and Gewirth, A. A., (2018). Nanoporous Copper-Silver Alloys by Additive-Controlled Electrodeposition for the Selective Electroreduction of CO2 to Ethylene and Ethanol. J. Am. Chem. Soc. 140, 5791-5797.

(14) Li, Y. C.; Wang, Z.; Yuan, T.; Nam, D. H.; Luo, M.; Wicks, J.; Chen, B.; Li, J.; Li, F.; de Arquer, F. P. G.; Wang, Y.; Dinh, C. T.; Voznyy, O.; Sinton, D. and Sargent, E. H., (2019). Binding Site Diversity Promotes CO2 Electroreduction to Ethanol. J. Am. Chem. Soc. 141, 8584-8591.

(15) Lee, S.; Park, G. and Lee, J., (2017). Importance of Ag–Cu biphasic boundaries for selective electrochemical reduction of CO2 to ethanol. ACS catal. 7, 8594-8604.

(16) Clark, E. L.; Hahn, C.; Jaramillo, T. F. and Bell, A. T., (2017). Electrochemical CO2 Reduction over Compressively Strained CuAg Surface Alloys with Enhanced Multi-Carbon Oxygenate Selectivity. J. Am. Chem. Soc. 139, 15848-15857.

(17) Zhou, Y.; Che, F.; Liu, M.; Zou, C.; Liang, Z.; De Luna, P.; Yuan, H.; Li, J.; Wang, Z.; Xie, H.; Li, H.; Chen, P.; Bladt, E.; Quintero-Bermudez, R.; Sham, T. K.; Bals, S.; Hofkens, J.; Sinton, D.; Chen, G. and Sargent, E. H., (2018). Dopant-induced electron localization drives CO2 reduction to C2 hydrocarbons. Nat. Chem. *10*, 974-980.

(18) Buckley, A. K.; Lee, M.; Cheng, T.; Kazantsev, R. V.; Larson, D. M.; Goddard Iii, W. A.; Toste, F. D. and Toma, F. M., (2019). Electrocatalysis at Organic-Metal Interfaces: Identification of Structure-Reactivity Relationships for CO2 Reduction at Modified Cu Surfaces. J. Am. Chem. Soc. 141, 7355-7364.

(19) Hoang, T. T. H.; Ma, S.; Gold, J. I.; Kenis, P. J. A. and Gewirth, A. A., (2017). Nanoporous Copper Films by Additive-Controlled Electrodeposition: CO2 Reduction Catalysis. ACS catal. 7, 3313-3321.

(20) Han, Z.; Kortlever, R.; Chen, H. Y.; Peters, J. C. and Agapie, T., (2017). CO2 Reduction Selective for C>/=2 Products on Polycrystalline Copper with N-Substituted Pyridinium Additives. ACS Cent. Sci. *3*, 853-859.

(21) Chang, X.; Wang, T.; Zhao, Z. J.; Yang, P.; Greeley, J.; Mu, R.; Zhang, G.; Gong, Z.; Luo, Z.; Chen, J.; Cui, Y.; Ozin, G. A. and Gong, J., (2018). Tuning Cu/Cu2 O Interfaces for the Reduction of Carbon Dioxide to Methanol in Aqueous Solutions. Angew. Chem. Int. Ed. 57, 15415-15419.

(22) Bai, S.; Shao, Q.; Wang, P.; Dai, Q.; Wang, X. and Huang, X., (2017). Highly Active and Selective Hydrogenation of CO2 to Ethanol by Ordered Pd-Cu Nanoparticles. J. Am. Chem. Soc. 139, 6827-6830.

(23) Huang, J.; Mensi, M.; Oveisi, E.; Mantella, V. and Buonsanti, R., (2019). Structural Sensitivities in Bimetallic Catalysts for Electrochemical CO2 Reduction Revealed by Ag-Cu Nanodimers. J. Am. Chem. Soc. 141, 2490-2499.

(24) Varandili, S. B.; Huang, J.; Oveisi, E.; De Gregorio, G. L.; Mensi, M.; Strach, M.; Vavra, J.; Gadiyar, C.; Bhowmik, A. and Buonsanti, R., (2019). Synthesis of Cu/CeO2-x Nanocrystalline Heterodimers with Interfacial Active Sites To Promote CO2 Electroreduction. ACS catal. *9*, 5035-5046.

(25) Yamada, Y.; Tsung, C.-K.; Huang, W.; Huo, Z.; Habas, S. E.; Soejima, T.; Aliaga, C. E.; Somorjai, G. A. and Yang, P., (2011). Nanocrystal bilayer for tandem catalysis. Nat. Chem. 3, 372-376.

(26) Xie, C.; Chen, C.; Yu, Y.; Su, J.; Li, Y.; Somorjai, G. A. and Yang, P., (2017). Tandem Catalysis for CO2 Hydrogenation to C2-C4 Hydrocarbons. Nano Lett. 17, 3798-3802.

(27) Li, Z.; Qu, Y.; Wang, J.; Liu, H.; Li, M.; Miao, S. and Li, C., (2019). Highly selective conversion of carbon dioxide to aromatics over tandem catalysts. Joule 3, 570-583.

(28) Nitopi, S.; Bertheussen, E.; Scott, S. B.; Liu, X.; Engstfeld, A. K.; Horch, S.; Seger, B.; Stephens, I. E. L.; Chan, K.; Hahn, C.; Norskov, J. K.; Jaramillo, T. F. and Chorkendorff, I., (2019). Progress and Perspectives of Electrochemical CO2 Reduction on Copper in Aqueous Electrolyte. Chem. Rev. 119, 7610-7672.

(29) Li, F.; Li, Y. C.; Wang, Z.; Li, J.; Nam, D.-H.; Lum, Y.; Luo, M.; Wang, X.; Ozden, A.; Hung, S.-F.; Chen, B.; Wang, Y.; Wicks, J.; Xu, Y.; Li, Y.; Gabardo, C. M.; Dinh, C.-T.; Wang, Y.; Zhuang, T.-T.; Sinton, D. and Sargent, E. H., (2019). Cooperative CO2-to-ethanol conversion via enriched intermediates at molecule–metal catalyst interfaces. Nat. Catal. *3*, 75-82.

(30) Fu, J.; Zhu, W.; Chen, Y.; Yin, Z.; Li, Y.; Liu, J.; Zhang, H.; Zhu, J. and Sun, S., (2019). Bipyridine-assisted assembly of Au nanoparticles on Cu nanowires to enhance electrochemical reduction of CO2. Angew. Chem. Int. Ed. 58, 14100-14103.

(31) Lum, Y. and Ager, J. W., (2018). Sequential catalysis controls selectivity in electrochemical CO2 reduction on Cu. Energy Environ. Sci. 11, 2935-2944.

(32) Gurudayal, G.; Perone, D.; Malani, S.; Lum, Y.; Haussener, S. and Ager, J. W., (2019). Sequential Cascade Electrocatalytic Conversion of Carbon Dioxide to CC Coupled Products. ACS Appl. Energy Mater. 2, 4551-4559.

(33) Morales-Guio, C. G.; Cave, E. R.; Nitopi, S. A.; Feaster, J. T.; Wang, L.; Kuhl, K. P.; Jackson, A.; Johnson, N. C.; Abram, D. N.; Hatsukade, T.; Hahn, C. and Jaramillo, T. F., (2018). Improved CO2 reduction activity towards C2+ alcohols on a tandem gold on copper electrocatalyst. Nat. Catal. *1*, 764-771.

(34) Gao, J.; Zhang, H.; Guo, X.; Luo, J.; Zakeeruddin, S. M.; Ren, D. and Gratzel, M., (2019). Selective C–C Coupling in Carbon Dioxide Electroreduction via Efficient Spillover of Intermediates As Supported by Operando Raman Spectroscopy. J. Am. Chem. Soc. 141, 18704-18714.

(35) Li, J.; Chen, G.; Zhu, Y.; Liang, Z.; Pei, A.; Wu, C.-L.; Wang, H.; Lee, H. R.; Liu, K.; Chu, S. and Cui, Y., (2018). Efficient electrocatalytic CO2 reduction on a three-phase interface. Nat. Catal. 1, 592-600.

(36) Dinh, C.-T.; Burdyny, T.; Kibria, M. G.; Seifitokaldani, A.; Gabardo, C. M.; de Arquer, F. P. G.; Kiani, A.; Edwards, J. P.; De Luna, P. and Bushuyev, O. S., (2018). CO2 electroreduction to ethylene via hydroxide-mediated copper catalysis at an abrupt interface. Science *360*, 783-787.

(37) Burdyny, T. and Smith, W. A., (2019). CO2 reduction on gas-diffusion electrodes and why catalytic performance must be assessed at commerciallyrelevant conditions. Energy Environ. Sci. 12, 1442-1453.

(38) Zheng, T.; Jiang, K. and Wang, H., (2018). Recent Advances in Electrochemical CO2 -to-CO Conversion on Heterogeneous Catalysts. Adv. Mater. 3, 1802066.

(39) Lee, C. W.; Yang, K. D.; Nam, D. H.; Jang, J. H.; Cho, N. H.; Im, S. W. and Nam, K. T., (2018). Defining a Materials Database for the Design of Copper Binary Alloy Catalysts for Electrochemical CO2 Conversion. Adv. Mater. 30, 1704717.

(40) Zhuang, T.-T.; Pang, Y.; Liang, Z.-Q.; Wang, Z.; Li, Y.; Tan, C.-S.; Li, J.; Dinh, C. T.; De Luna, P.; Hsieh, P.-L.; Burdyny, T.; Li, H.-H.; Liu, M.; Wang, Y.; Li, F.; Proppe, A.; Johnston, A.; Nam, D.-H.; Wu, Z.-Y.; Zheng, Y.-R.; Ip, A. H.; Tan, H.; Chen, L.-J.; Yu, S.-H.; Kelley, S. O.; Sinton, D. and Sargent, E. H., (2018). Copper nanocavities confine intermediates for efficient electrosynthesis of C3 alcohol fuels from carbon monoxide. Nat. Catal. *1*, 946-951.

(41) Li, Y.; Kim, D.; Louisia, S.; Xie, C.; Kong, Q.; Yu, S.; Lin, T.; Aloni, S.; Fakra, S. C. and Yang, P., (2020). Electrochemically scrambled nanocrystals are catalytically active for CO2-to-multicarbons. Proc. Natl. Acad. Sci. U.S.A. 117, 9194-9201.

(42) Farmand, M.; Landers, A. T.; Lin, J. C.; Feaster, J. T.; Beeman, J. W.; Ye, Y.; Clark, E. L.; Higgins, D.; Yano, J.; Davis, R. C.; Mehta, A.; Jaramillo, T. F.; Hahn, C. and Drisdell, W. S., (2019). Electrochemical flow cell enabling operando probing of electrocatalyst surfaces by X-ray spectroscopy and diffraction. Phys. Chem. Chem. Phys. 21, 5402-5408.

(43) Arán-Ais, R. M.; Scholten, F.; Kunze, S.; Rizo, R. and Cuenya, B. R., (2020). The role of in situ generated morphological motifs and Cu (i) species in C 2+ product selectivity during CO 2 pulsed electroreduction. Nature Energy 5, 317-325.

(44) Lum, Y. and Ager, J. W., (2018). Stability of Residual Oxides in Oxide-Derived Copper Catalysts for Electrochemical CO2 Reduction Investigated with (18) O Labeling. Angew. Chem. Int. Ed. 57, 551-554.

(45) Gao, D.; Zegkinoglou, I.; Divins, N. J.; Scholten, F.; Sinev, I.; Grosse, P. and Roldan Cuenya, B., (2017). Plasma-Activated Copper Nanocube Catalysts for Efficient Carbon Dioxide Electroreduction to Hydrocarbons and Alcohols. ACS Nano 11, 4825-4831.

(46) Jung, H.; Lee, S. Y.; Lee, C. W.; Cho, M. K.; Won, D. H.; Kim, C.; Oh, H. S.; Min, B. K. and Hwang, Y. J., (2019). Electrochemical fragmentation of Cu2O nanoparticles enhancing selective C-C coupling from CO2 reduction reaction. J. Am. Chem. Soc. 141, 4624-4633.

(47) Chou, T. C.; Chang, C. C.; Yu, H. L.; Yu, W. Y.; Dong, C. L.; Velasco-Velez, J. J.; Chuang, C. H.; Chen, L. C.; Lee, J. F.; Chen, J. M. and Wu, H. L., (2020). Controlling the Oxidation State of the Cu Electrode and Reaction Intermediates for Electrochemical CO2 Reduction to Ethylene. J. Am. Chem. Soc. 142, 2857-2867.

(48) Kim, D.; Kley, C. S.; Li, Y. and Yang, P., (2017). Copper nanoparticle ensembles for selective electroreduction of CO2 to C2–C3 products. Proc. Natl. Acad. Sci. U.S.A. 114, 10560-10565.

(49) Huang, J.; Hormann, N.; Oveisi, E.; Loiudice, A.; De Gregorio, G. L.; Andreussi, O.; Marzari, N. and Buonsanti, R., (2018). Potential-induced nanoclustering of metallic catalysts during electrochemical CO2 reduction. Nat. Commun. *9*, 3117.

(50) Kim, Y. G.; Baricuatro, J. H.; Javier, A.; Gregoire, J. M. and Soriaga, M. P., (2014). The evolution of the polycrystalline copper surface, first to Cu(111) and then to Cu(100), at a fixed CO(2)RR potential: a study by operando EC-STM. Langmuir *30*, 15053-15056.

(51) Osowiecki, W. T.; Nussbaum, J. J.; Kamat, G. A.; Katsoukis, G.; Ledendecker, M.; Frei, H.; Bell, A. T. and Alivisatos, A. P., (2019). Factors and Dynamics of Cu Nanocrystal Reconstruction under CO2 Reduction. ACS Appl. Energy Mater. 2, 7744-7749.

(52) Osowiecki, W. T.; Ye, X.; Satish, P.; Bustillo, K. C.; Clark, E. L. and Alivisatos, A. P., (2018). Tailoring Morphology of Cu-Ag Nanocrescents and Core-Shell Nanocrystals Guided by a Thermodynamic Model. J. Am. Chem. Soc.

(53) Wang, X.; Wang, Z.; Zhuang, T.-T.; Dinh, C.-T.; Li, J.; Nam, D.-H.; Li, F.; Huang, C.-W.; Tan, C.-S. and Chen, Z., (2019). Efficient upgrading of CO to C 3 fuel using asymmetric CC coupling active sites. Nat. Commun. 10, 5186.

(54) Wang, Y.; Shen, H.; Livi, K. J.; Raciti, D.; Zong, H.; Gregg, J.; Onadeko, M.; Wan, Y.; Watson, A. and Wang, C., (2019). Copper Nanocubes for CO2 Reduction in Gas Diffusion Electrodes. Nano Lett. 19, 8461-8468.

(55) Luc, W.; Fu, X.; Shi, J.; Lv, J.-J.; Jouny, M.; Ko, B. H.; Xu, Y.; Tu, Q.; Hu, X.; Wu, J.; Yue, Q.; Liu, Y.; Jiao, F. and Kang, Y., (2019). Two-dimensional copper nanosheets for electrochemical reduction of carbon monoxide to acetate. Nat. Catal. 2, 423-430.

(56) Jouny, M.; Luc, W. and Jiao, F., (2018). High-rate electroreduction of carbon monoxide to multi-carbon products. Nat. Catal. 1, 748-755.

(57) Wang, X.; de Araujo, J. F.; Ju, W.; Bagger, A.; Schmies, H.; Kuhl, S.; Rossmeisl, J. and Strasser, P., (2019). Mechanistic reaction pathways of enhanced ethylene yields during electroreduction of CO2-CO co-feeds on Cu and Cu-tandem electrocatalysts. Nat. Nanotechnol. 14, 1063-1070.

(58) Li, J.; Wang, Z.; McCallum, C.; Xu, Y.; Li, F.; Wang, Y.; Gabardo, C. M.; Dinh, C.-T.; Zhuang, T.-T.; Wang, L.; Howe, J. Y.; Ren, Y.; Sargent, E. H. and Sinton, D., (2019). Constraining CO coverage on copper promotes high-efficiency ethylene electroproduction. Nat. Catal. 2, 1124-1131.

(59) Luo, M.; Wang, Z.; Li, Y. C.; Li, J.; Li, F.; Lum, Y.; Nam, D. H.; Chen, B.; Wicks, J.; Xu, A.; Zhuang, T.; Leow, W. R.; Wang, X.; Dinh, C. T.; Wang, Y.; Wang, Y.; Sinton, D. and Sargent, E. H., (2019). Hydroxide promotes carbon dioxide electroreduction to ethanol on copper via tuning of adsorbed hydrogen. Nat. Commun. *10*, 5814.

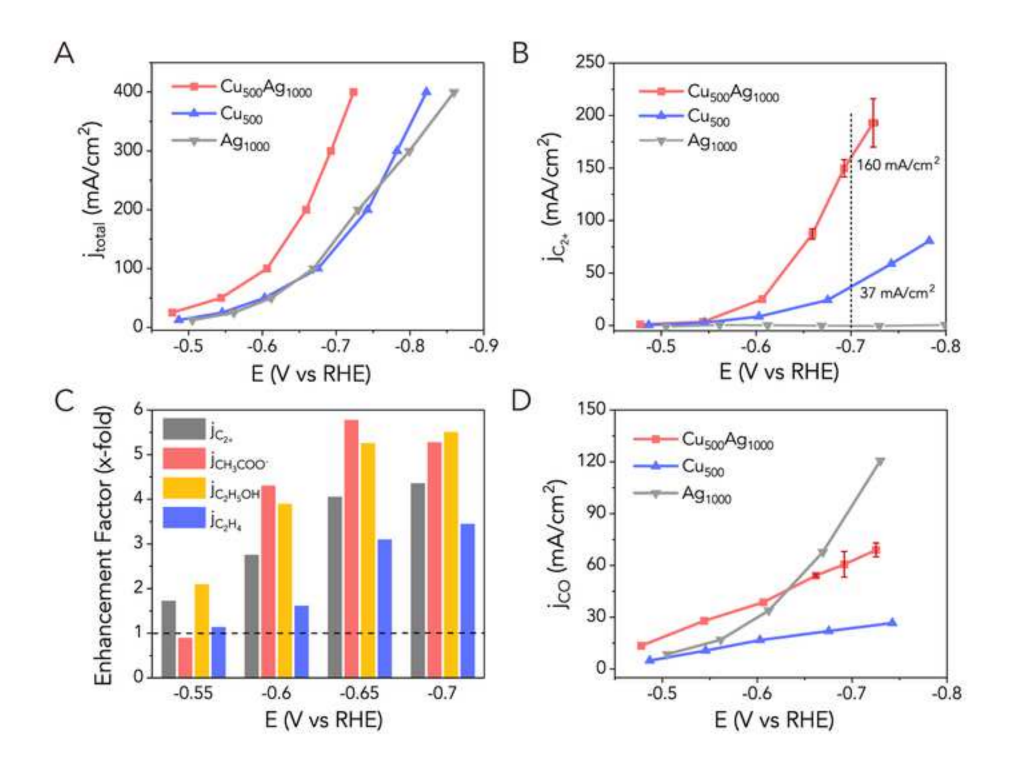
Figure 1. Cu-Ag tandem architecture design and characterizations. (a) Scheme of the Cu-Ag tandem platform for high rate CO₂ electrolysis and C₂₊ products formation. (b) Top view SEM image of Cu₅₀₀Ag₁₀₀₀ catalyst over carbon paper before electrolysis. (c) Cross-section SEM image of Cu₅₀₀Ag₁₀₀₀ catalyst over carbon paper before electrolysis. (d) XRD patterns of Cu₅₀₀Ag₁₀₀₀ catalyst over carbon paper before and after electrolysis. Inset: Cu(111) and Ag(200) range enlarged. Standard PDF index of Cu (ICDD PDF# 99-0034) and Ag (ICDD PDF# 99-0094) are plotted for comparison. (e) XPS spectrum of Cu₅₀₀Ag₁₀₀₀ catalyst over carbon paper before and after electrolysis, with Cu₅₀₀ and Ag₁₀₀₀ for comparison. Post-electrolysis samples were prepared after conducting CO₂RR at 200 mA/cm² constant cathodic current density for 30 min.

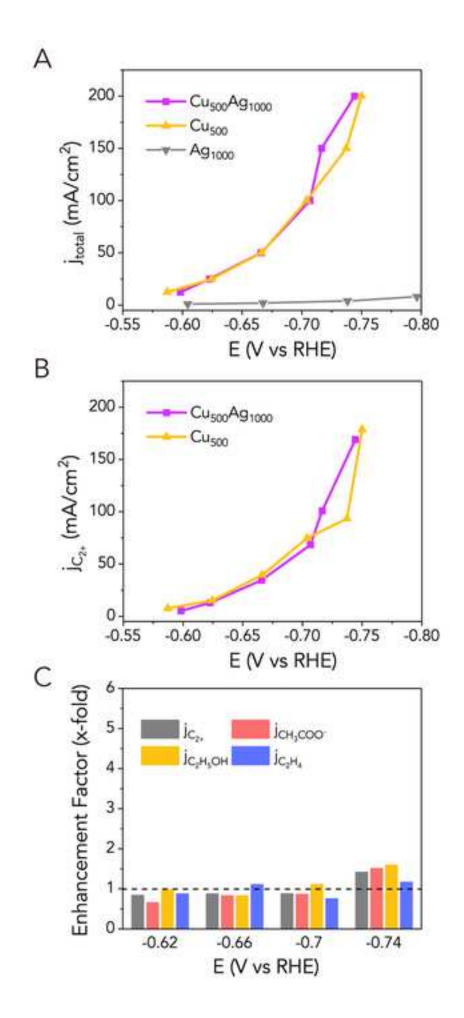
Figure 2. CO₂**RR performance of tandem catalysts comparing with pure metal catalysts.** (a) Total geometric current density curve of $Cu_{500}Ag_{1000}$, Cu_{500} and Ag_{1000} catalysts. (b) Partial current density towards C_{2+} products over $Cu_{500}Ag_{1000}$, Cu_{500} and Ag_{1000} catalysts. (c) Enhancement factors (EF) for total C_{2+} and individual products over four different potentials (-0.55V, -0.60V, -0.65V and -0.70V vs RHE) in CO_2RR , which is defined as the tandem $Cu_{500}Ag_{1000}$ partial current divided by that of Cu_{500} catalyst alone. Linear interpolation was used to obtain the estimated current values at these potentials. (d) CO production partial current density over $Cu_{500}Ag_{1000}$, Cu_{500} and Ag_{1000} catalysts. Data points with error bars (1 SD) in all figures based on average value of three separate experiments results.

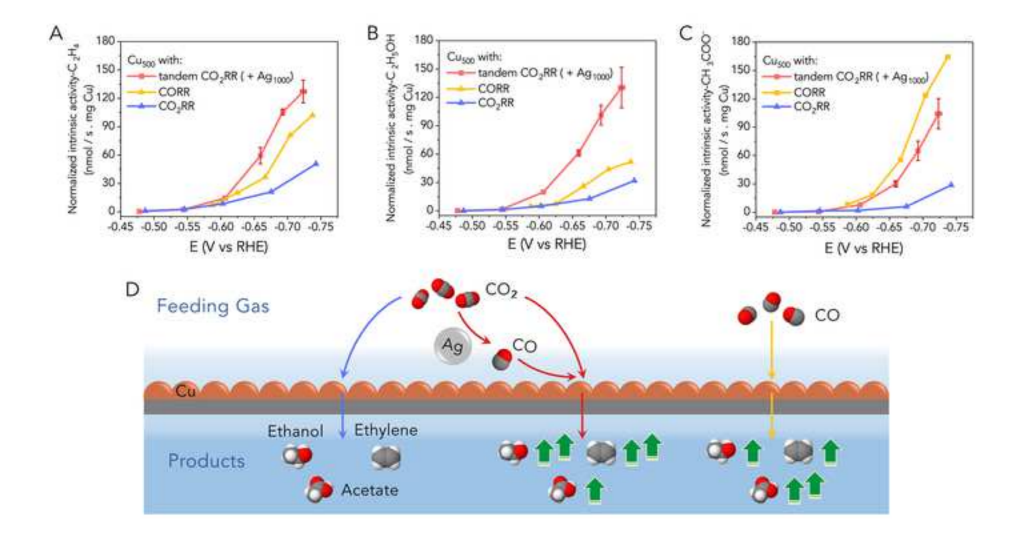
Figure 3. CORR performance of tandem catalysts comparing with pure metal catalysts. (a) Total currents of CO electroreduction reaction (CORR) over $Cu_{500}Ag_{1000}$, Cu_{500} and Ag_{1000} catalysts. (b) C_{2+} products partial currents of CORR over $Cu_{500}Ag_{1000}$ and Cu_{500} catalysts. (c) Enhancement factors (EF) for total C_{2+} and individual products over four different potentials (-0.62V, -0.66V, -0.70V and -0.74V vs RHE) in CORR, calculated as described in Figure 2.

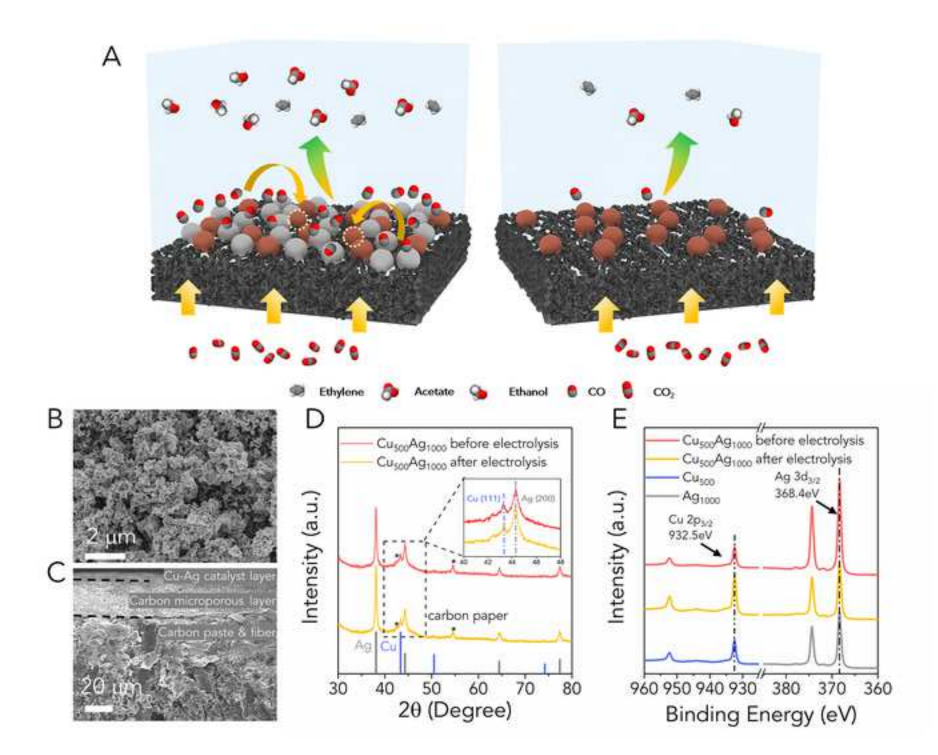
Figure 4. Normalized intrinsic activity analysis and corresponding schematic illustration. Normalized intrinsic activity towards (a) C_2H_4 (b) C_2H_5OH (c) CH_3COO^- over the Cu surface with different local environments (tandem CO₂ in red, fed CO in yellow, and fed

 CO_2 in blue). Data points with error bars (1 SD) in all figures based on average value of three separated experimental results. (d) Scheme representing the catalytic difference provided by the local environment produced by the tandem catalyst (red) vs. standard CORR (yellow) or CO_2RR (blue).









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- Figure S13. Imitating tandem architecture catalytic performance with a blended gas of 2% CO and 98% CO₂.

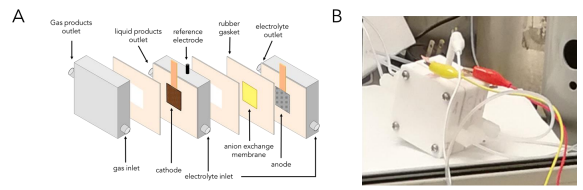


Figure S1. Gas diffusion cell configuration used in this work: (a) schematic illustration and (b) photograph.

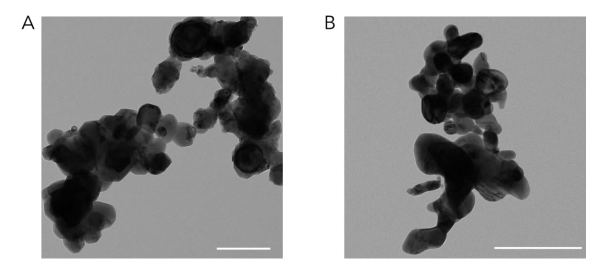


Figure S2. Additional morphology characterization of commercial Cu and Ag powders. (a) TEM image of commercial Cu (b) TEM image of commercial Ag. Scale bar: 200 nm.

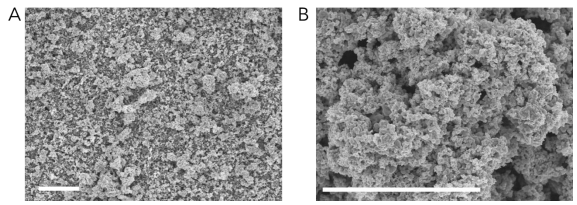


Figure S3. **SEM image of tandem Cu**₅₀₀**Ag**₁₀₀₀**/carbon paper after electrolysis.** (a) low magnification and (b) high magnification SEM image of tandem Cu₅₀₀Ag₁₀₀₀ catalyst after CO₂ electrolysis (30 min, 200 mA/cm², 1M KOH electrolyte). Scale bar: 2 μ m.

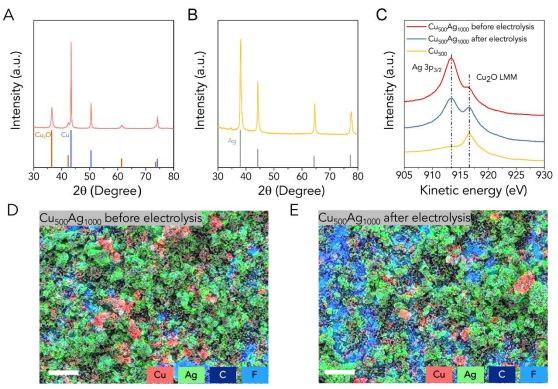


Figure S4. **Supplementary elemental characterization data.** (a) commercial Cu and (b) commercial Ag. Standard PDF index of Cu (ICDD PDF# 99-0034), Cu₂O (ICDD PDF# 99-0041), Ag (ICDD PDF# 99-0094) were also plotted for comparison. (c) LMM auger spectrum of tandem catalyst before and after electrolysis. Pure Cu catalyst was also characterized for reference. The auger peak of Cu₂O is partially overlapped with Ag 3p_{3/2}. Both XRD and XPS data confirm the existence of Cu₂O at particle surface, likely due to surface oxidation. Elemental mapping of tandem Cu₅₀₀Ag₁₀₀₀ catalyst (c) before and (d) after CO₂ electrolysis. Scale bar: 5 μm. Based on EDS elemental mapping, Cu and Ag powder were mixed at micron level, which is consistent with the TEM image in Fig S2. For Cu₅₀₀Ag₁₀₀₀ catalyst, it can be observed that Cu and Ag are segregated both before and after CO₂ electrolysis. This offers additional supporting evidence for confirming the lack of bulky alloy formation.

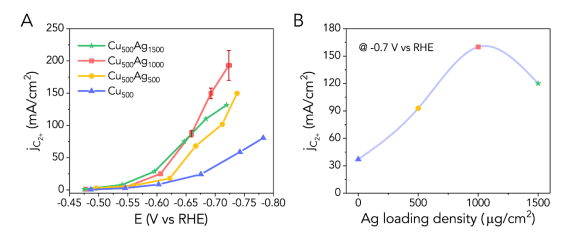


Figure S5. **Ag-loading dependent catalytic performance towards multicarbons over tandem catalysts.** (a) Partial current density towards C_{2+} products over tandem $Cu_{500}Ag_{1500}$, $Cu_{500}Ag_{1000}$, $Cu_{500}Ag_{500}$ and Cu_{500} catalysts in CO₂RR. (b) The relationship between Ag loading density in tandem catalyst and partial current density towards C_{2+} products at -0.70 V vs RHE. Partial current density values at -0.70 V vs RHE were obtained by linear interpolation. While Ag loading density below 1000 µg/cm², higher loading will result in larger C₂₊ partial current. But when Ag loading density exceed 1000 µg/cm², no further promotion could be observed, which may due to blockage of available Cu sites due to overcrowding of Ag.

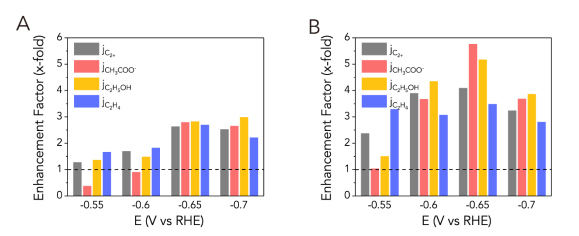


Figure S6. Enhancement factors (EF) between (a) $Cu_{500}Ag_{500}$ and Cu_{500} (b) $Cu_{500}Ag_{1500}$ and Cu_{500} during CO_2RR , which is defined as the quotient of the partial currents between two catalysts. Both total and specific C_{2+} products (C_2H_4 , C_2H_5OH , $CH3COO^-$) EFs were calculated over four different potentials (-0.55V, - 0.60V, -0.65V and -0.70V vs RHE). Linear interpolation was used to obtain the estimated current values at these potentials.

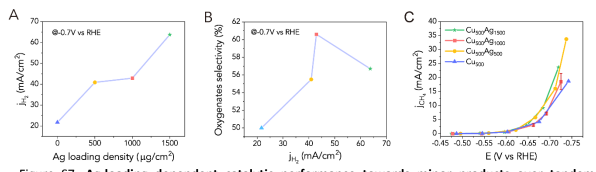


Figure S7. Ag-loading dependent catalytic performance towards minor products over tandem catalysts. (a) Partial current density towards H₂ over tandem $Cu_{500}Ag_{1500}$, $Cu_{500}Ag_{1000}$, $Cu_{500}Ag_{500}$ and Cu_{500} catalysts at -0.70 V vs RHE in CO₂RR. (b) The relationship between HER partial current density in tandem catalyst and oxygenates selectivity in multicarbon products at -0.70 V vs RHE. Partial current density values at -0.70 V vs RHE were obtained by linear interpolation. Generally, these results do not support a negative correlation between oxygenates selectivity and HER partial current density, which has been previously observed in Ag doped Cu catalyst¹. Instead, a positive correlation can be found between Ag loading density and HER partial current density, suggesting a different mechanism for C₂₊ enhancement in this work relative to Ag-doped Cu works. (c) Partial current density towards CH₄ over tandem $Cu_{500}Ag_{1500}$, $Cu_{500}Ag_{1000}$, $Cu_{500}Ag_{500}$ and Cu_{500} catalysts in CO₂RR. The methane production rate seems to be independent with Ag loading density, which may suggest that methane production is not CO-limited.

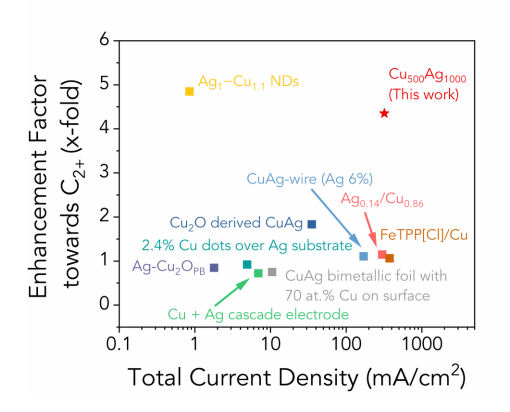


Figure S8. **Contextualization and comparison with relevant literature**. Total current density under most representative potential and the enhancement factors towards multicarbons are used as two indicators for comparison. More detailed information, including electrolyte, potential, total current density value and enhancement factors towards specific multicarbon products, can be found in table S1.

Although the alloying strategy (CuAg bimetallic foil; Ag0.14/Cu0.86; Ag-Cu2OPB) is a promising technique for tuning the selectivity of multicarbons, due to blockage of the surface, the value of EF towards total multicarbons are all around 1. Instead, tandem catalysts have a better potential to further improve EF (e.g. Cu2O derived CuAg). We attribute a higher EF in our work to the efficient CO-generating component in the catalyst (as compared with FeTPP[Cl] as an example) and high surface pH derived from electrolyte species and high electrolysis rate.

Note: (1) Enhancement factor (EF) here is defined as the quotient of partial current density between composed catalyst with reference pure Cu catalyst in literatures. Due to the varied morphology, ECSA and loading density of reference catalyst comparing with composed catalyst in some literatures, we remind readers of the potential imprecision contained in the calculated EF. (2) For the catalyst of Ag_{0.14}/Cu_{0.86}, offered comparison data are based on the same current density instead of same potential. (3) Due to the lack of same potential data, the product distribution under -0.84 V instead of -0.82 V vs RHE is used for the reference catalyst of FeTPP[CI]/Cu.

Table S1. **Catalytic performance and enhancement factor comparison of relevant literatures**. The values of current densities were either offered in literature or extracted from the corresponding figures. Bottom half of table represents GDE works whereas top half represents H-cell.

Catalyst	Electrolyte	Potential (V vs RHE)	Total current – density (mA/cm²)	Enhancement factor			
				C ₂ H ₄	C₂H₅OH	CH₃COOH & CH₃CHO	Total C ₂₊
CuAg bimetallic foil with 70 at.% Cu on surface ¹	0.1 M CsHCO₃	-1.05	10.1	0.6	1.2	4.5	0.8
Cu + Ag cascade electrode ²	0.1 M CsHCO ₃	-1.0	6.9	0.4	1.4	1.2	0.7
Ag-Cu ₂ O _{PB³}	0.2 M KCl	-1.2	1.8	0.3	1.9	0.9	0.9
2.4% Cu dots on Ag substrate ⁴	0.1 M KHCO3	-1.0	4.9	0.4	0.9	11.5	0.9
Cu ₂ O derived CuAg ⁵	0.1 M KHCO₃	-1.05	35.1	2.1	1.3	1.5	1.8
Ag ₁ -Cu _{1.1} NDs ⁶	0.1 M KHCO3	-1.1	0.85	4.9	N.A.	N.A.	4.9
Ag _{0.14} /Cu _{0.86} ⁷	1 M KOH	N.A.	300	0.7	1.4	3.7	1.2
CuAg-wire (Ag 6%) ⁸	1 М КОН	-0.6	170	1.3	0.9	0.8	1.1
FeTPP[Cl]/Cu ⁹	1 M KHCO3	-0.82	375	0.8	1.4	1.0	1.1
Cu ₅₀₀ Ag ₁₀₀₀ (This work)	1 М КОН	-0.7	320	3.4	5.5	5.3	4.4

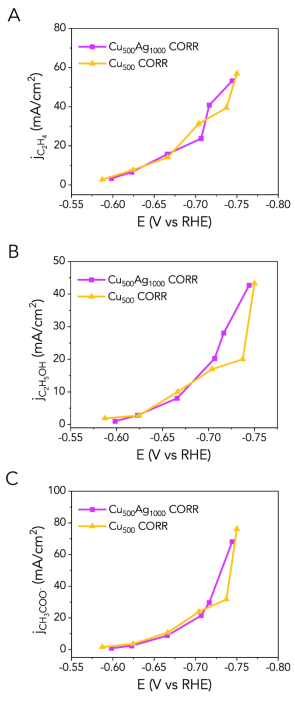


Figure S9. Partial current density of (a) C_2H_4 , (b) C_2H_5OH and (c) CH_3COO^- over both tandem $Cu_{500}Ag_{1000}$ and Cu_{500} catalyst in CORR. No significant difference is observed.

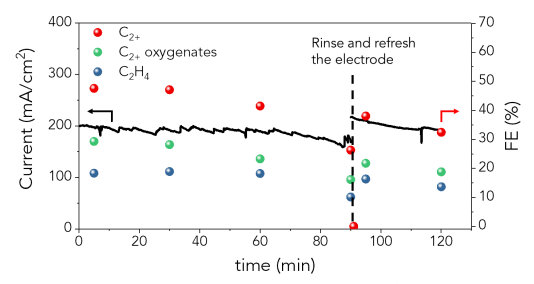


Figure S10. **Stability test under constant applied potential with 200 mA/cm² as starting current density.** The rapid catalytic deactivation happened after 1h electrolysis, which mainly due to the severe flood in gas diffusion layer. After the electrode fully deactivated, it was rinsed with DI water to remove electrolyte and precipitated out salts (*e.g.* K₂CO₃) thoroughly and then dried in vacuum chamber. The recovered catalytic activity during resumed CO₂ electrolysis indicates the deactivation mostly due to the instability of the carbon paper and the accumulation of carbonate salts from KOH-CO₂ reactivity.

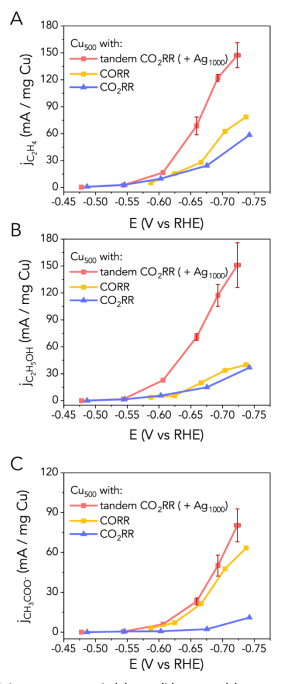


Figure S11. Normalized partial currents towards (a) C_2H_4 (b) C_2H_5OH (c) CH_3COO^- over Cu_{500} surface with different local environments.

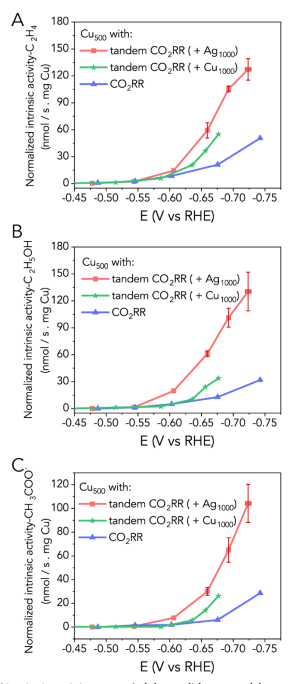


Figure S12. Normalized intrinsic activity towards (a) C_2H_4 (b) C_2H_5OH (c) CH_3COO^- over $Cu_{500}Cu_{1000}$ in CO_2RR . Cu_{500} and $Cu_{500}Ag_{1000}$ were also included for comparison.

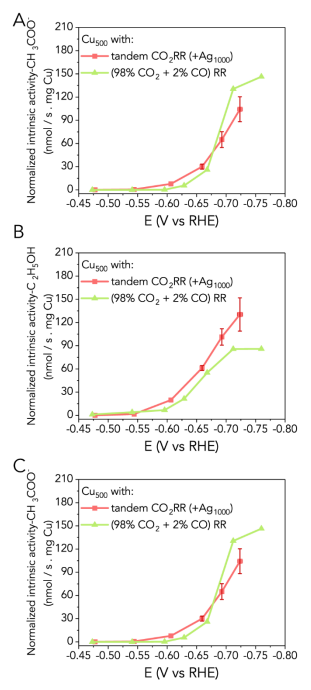


Figure S13. Imitating tandem catalytic performance with a mixed gas of 2% CO and 98% CO₂. Normalized intrinsic activity towards (a) C_2H_4 (b) C_2H_5OH (c) CH_3COO^- over Cu_{500} in a mixed gas atmosphere with 98% CO₂ and 2% CO. $Cu_{500}Ag_{1000}$ CO₂RR data were also included for comparison. Under comparable CO partial pressure, a similar intrinsic activity enhancement could also be observed on Cu_{500} , indicating the correlation between increased performance and the co-existence of CO and CO₂.

References:

(1) Clark, E. L.; Hahn, C.; Jaramillo, T. F. and Bell, A. T., (2017). Electrochemical CO2 Reduction over

Compressively Strained CuAg Surface Alloys with Enhanced Multi-Carbon Oxygenate Selectivity. J. Am. Chem. Soc. *139*, 15848-15857.

(2) Gurudayal, G.; Perone, D.; Malani, S.; Lum, Y.; Haussener, S. and Ager, J. W., (2019). Sequential Cascade Electrocatalytic Conversion of Carbon Dioxide to CC Coupled Products. ACS Appl. Energy Mater. 2, 4551-4559.

(3) Lee, S.; Park, G. and Lee, J., (2017). Importance of Ag–Cu biphasic boundaries for selective electrochemical reduction of CO2 to ethanol. ACS catal. *7*, 8594-8604.

(4) Lum, Y. and Ager, J. W., (2018). Sequential catalysis controls selectivity in electrochemical CO2 reduction on Cu. Energy Environ. Sci. *11*, 2935-2944.

(5) Gao, J.; Zhang, H.; Guo, X.; Luo, J.; Zakeeruddin, S. M.; Ren, D. and Grätzel, M., (2019). Selective C–C Coupling in Carbon Dioxide Electroreduction via Efficient Spillover of Intermediates As Supported by Operando Raman Spectroscopy. J. Am. Chem. Soc. *141*, 18704-18714.

(6) Huang, J.; Mensi, M.; Oveisi, E.; Mantella, V. and Buonsanti, R., (2019). Structural Sensitivities in Bimetallic Catalysts for Electrochemical CO2 Reduction Revealed by Ag-Cu Nanodimers. J. Am. Chem. Soc. *141*, 2490-2499.

(7) Li, Y. C.; Wang, Z.; Yuan, T.; Nam, D. H.; Luo, M.; Wicks, J.; Chen, B.; Li, J.; Li, F.; de Arquer, F. P. G.; Wang, Y.; Dinh, C. T.; Voznyy, O.; Sinton, D. and Sargent, E. H., (2019). Binding Site Diversity Promotes CO2 Electroreduction to Ethanol. J. Am. Chem. Soc. *141*, 8584-8591.

(8) Hoang, T. T. H.; Verma, S.; Ma, S.; Fister, T. T.; Timoshenko, J.; Frenkel, A. I.; Kenis, P. J. A. and Gewirth, A. A., (2018). Nanoporous Copper-Silver Alloys by Additive-Controlled Electrodeposition for the Selective Electroreduction of CO2 to Ethylene and Ethanol. J. Am. Chem. Soc. *140*, 5791-5797.

(9) Li, F.; Li, Y. C.; Wang, Z.; Li, J.; Nam, D.-H.; Lum, Y.; Luo, M.; Wang, X.; Ozden, A.; Hung, S.-F.; Chen, B.; Wang, Y.; Wicks, J.; Xu, Y.; Li, Y.; Gabardo, C. M.; Dinh, C.-T.; Wang, Y.; Zhuang, T.-T.; Sinton, D. and Sargent, E. H., (2019). Cooperative CO2-to-ethanol conversion via enriched intermediates at molecule–metal catalyst interfaces. Nat. Catal. *3*, 75-82.