We rationally designed an epitaxial Au–Cu heterostructure as a model system to accurately correlate the atomic reconstruction with catalytic performance and revealed a brand-new insight into tandem CO$_2$-to-C$_2^+$ alcohols conversion on bimetallic interface. We illuminated that the in situ formed AuCu alloy and Cu shell served as the active phases for $^*$CO buildup and $^*$CO-to-C$_2^+$ alcohols conversion, respectively, and enriching $^*$CO was a crucial step to keep the durable production of C$_2^+$ alcohols.
Dynamic restructuring of epitaxial Au–Cu biphasic interface for tandem CO2-to-C2+ alcohols conversion

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SUMMARY
Interfacing Cu with a secondary metal is an effective strategy to enhance the production of value-added C2+ fuels in CO2 electroreduction. However, such a biphasic interface generally suffers unclear dynamic reconstruction/phase transformation, rendering the structure-function correlation elusive. Here, we studied a model system of epitaxial Au–Cu heterostructure, which exhibits a ~150 mV more positive onset potential for C2+ alcohols and a 400-fold improved alcohols production over hydrocarbons, relative to primitive Cu. We unambiguously revealed a dynamic restructuring of such heterostructure, from phase-separated bimetals to alloy-supported core-shell nanoclusters, driven by the oxidation/reduction of Cu(0) at the interface. A distinct tandem mechanism was proposed, and the buildup of *CO was identified as crucial to keeping the production durability of C2+ alcohols. This work fills in the voids of direct observation on the dynamic restructuring of the bimetallic interface and establishes a paradigm to understand the tandem CO2-to-C2+ alcohols conversion from an atomic view.

INTRODUCTION

The over-reliance on fossil fuels gave rise to the global-warming crisis, and thus technologies that are capable of converting greenhouse gas CO2 to value-added chemicals are therefore of interest.¹⁵ Electrochemical CO2 reduction (CO2R) holds a great promise to transform CO2 to valuable fuels and potentially serves as a supplement to the petrochemical feedstock.³,⁴ Among various products from CO2R, multi-carbon (C2+) fuels, such as ethylene, ethanol, and n-propanol, etc., are highly demanded because of their high energy densities and ease of storage/transportation.⁵–⁷ Despite tremendous efforts since the pioneering work of Hori et al.,⁸–¹⁰ the main hurdle for CO2R remains the development of highly active, selective, and durable electrocatalysts. Owing to the optimal binding energy with the intermediate *CO, Cu is the only recognized catalyst that can catalyze carbon dimerization and produce value-added C2+ fuels.¹¹–¹³ However, CO2R on monometallic Cu is processed at very high overpotentials and, in most cases, suffers from a poor selectivity and yield rate toward C2+, heavily impeding their practical applications.

Interfacing Cu with a secondary metal to form a bimetallic catalyst has been shown effective to enhance the selectivity toward C2+ products from tandem electrocatalysis,¹⁴–¹⁶ benefiting from the tunable binding energy between the bimetallic catalyst and key intermediates, such as *CO. For instance, Pd–Cu and Ag–Cu bimetallic catalysts have been proven to effectively improve the selectivity toward ethylene,¹⁷–¹⁹
and Au–Cu will boost the production of C$_2$+ alcohols.\textsuperscript{20,21} Most of these studies to date established good understandings toward the improved performances from a theoretical perspective without an in-depth empirical atomic-scale structure characterization. However, the structural dynamics of catalysts in electrocatalysis complicate such theoretical understanding, in particular when phase transformation and surface restructuring occur during CO$_2$R. The lack of accurate atomic-scale structural characterization would particularly impede the fundamental understanding of tandem CO$_2$-to-C$_2$+ conversion on a bimetallic interface.

Herein, an epitaxial Au–Cu heterostructure was employed as a model system to understand the dynamic reconstruction/phase transformation of the bimetallic interface under CO$_2$R. Comparing with monometallic Cu, the epitaxial Au–Cu exhibits a \textasciitilde 150 mV more positive onset potential toward C$_2$+ alcohols and a 400-fold improvement on the production of alcohols over hydrocarbons at a low overpotential. We probed the bimetallic interface at an atomic scale and observed an unambiguous interface restructuring, from a phase-separated Au–Cu interface to unique AuCu alloy-supported epitaxial Au@Cu core-shell nanoclusters, driven by the oxidation-reduction process of Cu(0) on electrode-electrolyte interface. Combining with \textit{in situ} Fourier-transform infrared (FTIR) spectroscopy, finite-element simulation, and density functional theory (DFT) calculation, we illuminated that the \textit{in situ} formed AuCu alloy and Cu shell serves as the active phase for *CO buildup and *CO-to-C$_2$+, alcohols transition, respectively, and enriching *CO is a key step to keep the durable production of C$_2$+, alcohols. This work reveals the empirical demonstration to track the atomic restructuring of a model bimetallic system, which not only sets a paradigm to correlate the surface structure and catalytic performance but also brings a brand-new insight to tandem CO$_2$R on bimetallic electrocatalysts.

RESULTS AND DISCUSSION

Structural characterization of epitaxial Au–Cu heterostructure

A bimetallic interface with a well-defined atomic configuration is fairly important for the subsequent study on dynamic reconstruction. Here, we chose an epitaxial Au–Cu heterostructure as a model system. Typically, Au nanoclusters were electrochemically deposited on a monocrystalline Cu substrate, which was fabricated via our recently developed “seeded growth” technique.\textsuperscript{22} The single-crystal nature of Cu support was corroborated by X-ray diffraction (XRD) and selected area electron diffraction (SAED) taken over a large area (Figures S1 and S2). The atomic force microscopy (AFM) and scanning electron microscopy (SEM) images demonstrated an even dispersion of uniform Au nanoclusters after electrodeposition, with a typical size of \textasciitilde 10 nm (Figures 1A and S3). X-ray photoelectron spectroscopy (XPS) measurements showed that the atomic content of Au can be well controlled in the range of 2\%–10\% depending on the deposition time (Figure S4). As shown in Figure 1B, the Au 4f XPS spectra can be fitted into two symmetric peaks with binding energies of 87.9 (4f$_{5/2}$) and 84.2 (4f$_{7/2}$) eV, respectively, indicative of the existence of Au(0). The Cu 2p$_{3/2}$ peak was deconvoluted into three-subpeaks at 932.5, 933.5, and 935.0 eV, respectively, confirming the mixed valence states of Cu(0) and Cu(II). The presence of Cu(II) can be attributed to the formation of an ultrathin CuO and CuCO$_3$ layer after exposing electrode in air.\textsuperscript{20,23}

Next, electron microscopy was employed to illuminate the as-prepared Au–Cu interface at an atomic scale. The cross-sectional transmission electron microscopy (TEM) sample was prepared using focused ion beam (FIB), with detailed methods provided in experimental procedures and Figure S5. It was observed that the Cu surface

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displayed a rough geometry, having series of peaks with vertex angles distributed in the range of 52°–63° (see Figure S2 and Table S1). The scanning TEM (STEM) image illuminated uniform Au clusters coated on the highly uneven Cu surface after electro-deposition (Figure S6). Elaborate TEM imaging and electron diffraction analysis revealed that quantities of Cu peaks exposed (100) terraces (Figures S7 and S8), which is highly likely due to the lowest required formation energy under such Au deposition condition. Furthermore, an epitaxial Au–Cu interface was disclosed by the aberration-corrected high-angle annular dark-field (HAADF)-STEM image based on Z-contrast (Figure 1C). The lattice spacing of {200} increased from 1.82 Å in the bulk of Cu to 1.99 Å at the Au–Cu interface and further increased to 2.04 Å on the top surface of Au (to protect the Au–Cu interface during the FIB cutting, a thin Pt layer was deposited on top of Au).

**Figure 1.** Characterization of the as-prepared epitaxial Au–Cu heterostructure
(A) An AFM image (left) and corresponding height profile (right) along the white line.
(B) Au 4f and Cu 2p XPS analysis.
(C) A typical HAADF-STEM image of Au–Cu biphasic interface (top) and intensity line profiles extracted from the solid box (bottom), showing the lattice spacing of (200) increased from 1.82 Å in the bulk of Cu to 1.99 Å at the Au–Cu interface and further increased to 2.04 Å on the top-surface of Au (to protect the Au–Cu interface during the FIB cutting, a thin Pt layer was deposited on top of Au).
(D) FFT patterns taken from Au and Cu regions in (C) at the zone axis of <001>.
(E) EDS elemental maps at Au–Cu interface, showing the separate phases of Au and Cu.

Electrochemical reduction of CO₂
The CO₂R catalytic activity of Au–Cu heterostructure was examined in CO₂-saturated 0.1 M KHCO₃ with a gas-tight H cell. Considering the facet influence, we prepared Cu(100) and Au(100) as control samples, respectively (see details in experimental procedures). To quantitatively determine and compare the catalytic activity, gas products were quantified through the sampling system of an online
gas chromatography (GC), whereas the liquid products were collected and analyzed by 1H nuclear magnetic resonance (NMR) spectroscopy. 1 h chronoamperometry (CA) experiments were used to examine the activities of electrocatalyst across a broad range of potentials. The average experimental data, collected for at least three independent samples tested at each potential, were summarized in Figures S10–S12, including J_total-V curves, Faradaic efficiencies (FEs), and production rates for each product.

One can see that Cu(100) exhibited a comparable product distribution with that reported in the literature, generating more C_2+ hydrocarbon products, such as ethylene, than C_2+ alcohols. In sharp contrast, only hydrogen (H_2), carbon monoxide (CO), and formic acid were detected on Au, in particular, at −0.65 V versus reversible hydrogen electrode (RHE), and CO with a maximum FE of ~60% was the dominant CO_2R product (Figure S11). As shown in Figure 2A, an increasing FE and yield rate of CO on Cu, Au–Cu heterostructure, and Au were observed, consistent with a lower activation energy of Au for CO production, as unveiled by DFT calculations (Figure S13). The distinguished CO production can be also ascribed to the different binding energies between metal and *CO intermediate, e.g., the weak binding energy between Au and *CO promotes CO production, whereas the moderate binding energy between Cu and *CO is beneficial for further *CO reduction and >2e^− fuels (>2 mol of e^− are consumed for the generation of product per mol in CO_2R) production. Au–Cu heterostructure exhibited a fairly modest CO producing capability, with a maximum yield rate of ~18 μmol cm^−2 h^−1 at ~0.85 V versus RHE, which was ~10-fold higher than Cu, but 3 times inferior to Au.

Figure 2. Electrochemical CO_2R measurements
(A) Production rate and FE of CO on Au, Au–Cu, and Cu electrodes. (B) Production rate of C_2+ alcohols on Au–Cu and Cu electrodes. (C and D) Potential dependence of the molar ratio of C_2+ alcohols to hydrocarbons (C) and alcohols to (CO + >2e^− products) (D) on Au–Cu and Cu. (E) The stability of production rates of H_2, CO, and C_2+ alcohols on Au–Cu at −0.75 V versus RHE. All measurements were performed in a CO_2-saturated 0.1 M KHCO_3 electrolyte (pH 6.8), and the error bars represent one standard deviation of triplicate measurements.
For >2e⁻ fuels, Au–Cu heterostructure demonstrated a distinct production of C₂⁺ alcohols, with a ~150 mV more positive onset potential than Cu and a ~2.5-fold higher CO₂-to-C₂⁺ alcohols conversion rate at ~1.0 V versus RHE (Figure 2B). We further analyzed the production rates of hydrocarbons (e.g., methane and ethylene) (Figure S12), which followed competing production pathways with C₂⁺ alcohols.²⁶,²⁷ The mole ratio of C₂⁺ alcohols/hydrocarbons was used to demonstrate this competition. As shown in Figure 2C, this ratio increased dramatically with an overpotential dropping, indicating C₂⁺ alcohols are more preferable at less negative potentials. In particular, Au–Cu heterostructure outperformed Cu to produce alcohols and suppress hydrocarbons, as demonstrated by the positively shifted onset potential of alcohols (~0.7 V versus RHE on Au–Cu and ~0.85 V versus RHE on pure Cu), and more importantly, a 400-fold higher ratio of C₂⁺ alcohols/hydrocarbons at the onset potential of C₂⁺ alcohols. As *CO is regarded as the intermediate for >2e⁻ products, we next calculated the mole ratio of C₂⁺ alcohols/(CO + >2e⁻ products) to elucidate the electrochemical efficiency of *CO-to-C₂⁺ alcohols conversion (Figure 2D). The higher ratio can be clearly identified on Au–Cu heterostructure, which was almost 6-fold higher than that on Cu at ~0.85 V versus RHE, further demonstrating a preferred C₂⁺ alcohol production. For the durability test, the time-dependent production rate and current density of each product on Au–Cu were plotted in Figures 2E and S14, respectively. It was observed that such Au–Cu heterostructure exhibited a slightly declined CO₂R performance with time, e.g., the yield of CO and C₂⁺ alcohols dropped by ~40% (from 9.2 to 5.6 μmol cm⁻² h⁻¹) and ~49% (from 0.38 to 0.20 μmol cm⁻² h⁻¹) after 2 h electrocatalysis at ~0.75 V versus RHE, respectively. Instead, the production of H₂, as a major competing reaction, increased by ~16%. Such a performance evolution motivated us to probe the structural dynamics on Au–Cu interface down to an atomic scale, in order to elucidate the structure-function correlation.

**Atomic reconstruction of Au–Cu interface**

Atomic-scale HAADF-STEM imaging was carried out to reveal the Au–Cu interface after 1 h electrocatalysis at ~0.75 V versus RHE, and a unique core-shell nanostructure with a typical size of 10–15 nm was observed (Figure 3A). The interplanar d-spacing in the case of the shell can be assigned to Cu(100) planes. As shown in Figure 3B, the lattice fringe spacing of {200} increased from 1.82 Å on the top surface of Cu to 1.85 Å at the nanocluster/substrate interface, highly likely due to the tensile strain because of Au incorporation. Likewise, we characterized the core of this unique structure and identified that Au centered at the core, with a {200} interlayer spacing of 1.96 Å (Figure S15). The valence state of Au remained the same as pristine Au–Cu structure, as demonstrated by XPS analysis (Figure S16). The EDS elemental maps and FFT patterns further verified the location of Au and Cu (Figures 3C, 3D, and S17) and confirmed the Au@Cu core-shell structure. Therefore, the discrete Au nanocluster before CO₂R has been encapsulated in a Cu nanoshell. It is notable that the atomic migration of Cu onto Au was epitaxially, leading to a curved Cu surface together with a nano-space between two neighbored Cu shells.

In addition, the atomic-scale HAADF-STEM image and EDS elemental map at the planar interface corroborated an atomic interdiffusion between Au and Cu atoms (Figures 3E and 3F). The brighter spots showing a higher Z-contrast are corresponding to the Au atoms, proving their diffusion into the Cu matrix. Corresponding FFT patterns at the interface showed superlattice (100) diffraction points (Figure 3G), which is a strong evidence for an intermetallic alloy.²⁸ Such an evolution happened across the whole surface, and more HAADF-STEM and
EDS results on other positions can be found in Figures S17 and S18. This unique finding validates the statement that Au–Cu biphasic interface undergoes a severe reconstruction during CO$_2$R, leading to a structural transformation from phase-separated bimetals to AuCu alloy-supported epitaxial Au@Cu core-shell nanoclusters.

The dynamic restructuring of Au–Cu interface can be ascribed to the oxidation/reduction of Cu(0) during electrocatalysis, as illustrated by the schematic illustration shown in Figure 3H. At an open circuit potential, the Cu(0) exposed to electrolyte without Au protection is prone to be oxidized to Cu(I) and Cu(II),$^{29,30}$ and such chemical oxidation damages the well-defined atomic structure because of oxygen incorporation. Some Cu atoms are even released to electrolyte and generate a certain concentration of Cu ions, as verified by the inductively coupled plasma optical
emission spectrometry (ICP-OES) (Table S2). Under a cathodic bias, the oxygen removal will lead to the reduction of Cu(I) and Cu(II) and spontaneously leave abundant of atomic vacancies, which will greatly facilitate the atomic interdiffusion between Au and Cu, leading to AuCu alloy after applying a cathodic potential. At the same time, the electrochemical reduction will result in a redeposition of Cu ions from electrolyte, and such redeposition is prone to occur locally on the top surface of Au nanoclusters owing to their lower adsorption energies, followed by the formation of epitaxially Au@Cu core-shell nanoclusters. As far as we know, this is the first time to report the dynamic restructuring of Cu-based bimetallic heterostructure down to an atomic scale during tandem CO2R electrocatalysis.

Structure-property correlation
We employed operando FTIR spectroscopy to probe the adsorbate intermediates on both Au–Cu and Cu at different potentials. Four major peaks were explicitly discerned on Au–Cu surface (Figure 4A). The peak A appearing at 1,393–1,401 cm\(^{-1}\) was attributed to CO\(_3^{2-}\), whereas peak B at 1,620 cm\(^{-1}\) was assigned to the H–O–H bending mode of H\(_2\)O (\(\delta_{(\text{H–O–H})}\)). The band at 1,760 cm\(^{-1}\), peak C,
was in close proximity to the stretching of *CHO, which served as an important precursor for the subsequent C–C coupling.\(^3\)\(^4\) The band at \(\sim 2,100\ \text{cm}^{-1}\) (peak D), intensifying at more negative potentials, was attributed to the C\(=O\) stretching mode of atop-bonded *CO adlayers.\(^3\)\(^5\) In contrast to Au–Cu, Cu surface exhibited two strong bands assignable to CO\(_2^2\) and H\(_2\)O at 1,393–1,401 cm\(^{-1}\) (peak A) and 1,620 cm\(^{-1}\) (peak B), respectively. The peak D for *CO adsorbates at \(\sim 2,100\ \text{cm}^{-1}\) was much more inexplicit across the potential window. The negative band at \(\sim 2,345\ \text{cm}^{-1}\) on Cu was assigned to CO\(_2\) molecules in aqueous electrolyte.\(^3\)\(^2\) These spectroscopic results demonstrated that Au–Cu electrode exhibited a higher coverage of *CO, a critical intermediate in the producing pathway of C\(_2^+\) alcohols. The enriched *CO provided a microenvironment that can lower down the reaction barrier for C–C coupling and steered the selectivity toward C\(_2^+\) alcohols at a lower overpotential.\(^3\)\(^6\) Moreover, such in situ produced *CO cannot be replaced by CO supplying reagent, as the production rates of alcohols from the direct CO reduction on Cu and Au–Cu were both orders of magnitudes lower than that from CO\(_2\)R (Figure S19). Furthermore, Cu–Au exhibited scarcely any production preference toward C\(_2^+\) alcohols comparing with pure Cu. These observations experimentally proved the importance of locally enriched *CO for tandem catalysis on reconstructed Au–Cu surface.

The conventional view on tandem catalysis stated that CO\(_2\) would be first reduced to *CO on the secondary metal, such as Zn,\(^1\)\(^6\) Pd,\(^1\)\(^7\) Ag,\(^1\)\(^8,\)\(^1\)\(^9\) and Au,\(^2\)\(^0\) etc., building up a super-high *CO concentration in the vicinity, which would undergo dimerization and further be reduced to a higher-valued product on Cu. By contrast, our results proved that the biphasic interface will experience a significant dynamic reconstruction under CO\(_2\)R, which leads to evolution of active phases, suggesting that the classic tandem picture of electrocatalysis on biphasic interface may be insufficient. From our observations, the Au nanoclusters were fully encapsulated in a Cu shell and could not serve as efficient CO producing sites. Instead, the in situ generated AuCu alloy served as the active phase to buildup *CO at the interface, as inspired from Kim et al.\(^3\)\(^7\) Following, the in situ formed Cu shell would convert *CO to C\(_2^+\) alcohols. This mechanism was schematically shown in Figure 4B. Such a tandem picture can be also visualized via finite-element simulation (see details in experimental procedures). We set up a full micro-kinetic model that took into account all key elementary steps from CO\(_2\) to ethanol, including CO\(_2\)-to-*CO conversion on AuCu alloy support and Cu shell as well as *CO-to-ethanol conversion on Cu shell. As demonstrated in Figure 4C, *CO mainly generated on AuCu alloy surface and quickly diffused to the electrolyte (as CO molecules). Instead, ethanol preferred to generate at the Cu shell neighboring the AuCu alloy, where *CO was mostly enriched, unveiling a synergistic interaction of two phases for the tandem CO\(_2\)-to-alcohols production.

We attempted to correlate the yields of *CO and ethanol with the size of reconstructed Au@Cu clusters (Figures S20 and S21). Notably, the size increasing of Au@Cu cluster will lead to a higher coverage of Cu shell, whereas a lower coverage of exposed planar AuCu alloy. Figure 4D evidenced that at a given potential, both *CO and ethanol exhibited size-dependent production rates. The generation of *CO strongly declined with the increasing size of Cu shell, which could be attributed to the drop of AuCu area exposed. In stark contrast, the production of ethanol first showed a weakly positive correlation with Au@Cu cluster size below 5.5 nm, which, however, turned to a modestly negative correlation as the size increasing, suggesting the optimal size of Cu shell would be 5.5 nm. It is worth noting that the yields of *CO and ethanol showed a similar correlation when the size was higher than 9 nm,
indicating that although a larger curving Cu shell provided more active sites for *CO-to-C2+ alcohols conversion, the buildup of *CO determined the production rates of C2+ alcohols.

To unveil the influence of surface reconstruction on *CO enrichment, time-dependent in situ FTIR and DFT calculation were further performed. The FTIR spectra in Figure 4E showed that the intensity of *CHO and atop-bonded *CO (peak C and D, at 1,760 cm⁻¹ and ~2,100 cm⁻¹, respectively) gradually weakened with time prolonged, suggesting the reconstruction was detrimental to buildup *CO intermediate. DFT calculation was explored to reveal how the *CO coverage influenced C2+ alcohols generation on Cu surface (see supplemental information for details). We calculated the energy diagrams associated with hydrogenation along the formation pathway of ethanol at different *CO coverages. Results demonstrated that the energy barrier required for *HCCOH-to-*HCCHOH (rate-determining step) increased at a lower *CO coverage (Figure S22). Therefore, it is reasonable to ascribe the gradually dropped yield of C2+ alcohols to the limited supply of *CO intermediate from surface reconstruction. These observations were consistent with the declined yields of CO and C2+ alcohols with time shown in Figure 2E.

Conclusions
In summary, we studied a model system of epitaxial Au–Cu heterostructure having a biphasic interface for CO₂R, which demonstrated a remarkable tandem CO₂-to-C2+ alcohols conversion. Utilizing aberration-corrected HAADF-STEM, a dynamic structural reconstruction/transformation from bimetallic separated-phase into alloy-supported core-shell nanoclusters was evidently unveiled during the reaction. We proposed a novel CO₂-to-C2+, alcohols conversion mechanism distinguished with the conventional view and correlated the evolution of CO₂R performance to the limited supply of *CO intermediates. This work establishes the empirical demonstration of dynamic structural reconstruction/transformation in the bimetallic model system and sets up a paradigm to understand the tandem CO₂-to-C2+ alcohols conversion from an atomic view.

EXPERIMENTAL PROCEDURES
Resource availability
Lead contact
Further information and requests for resources and reagents should be directed to the lead contact, Liming Zhang (zhanglm@fudan.edu.cn).

Materials availability
The materials generated in this study will be made available on request.

Data and code availability
All of the data are available from the corresponding author upon reasonable request.

Synthesis of epitaxial Au–Cu bimetallic heterostructure
The single-crystal Cu substrate was prepared via the “seeded abnormal grain growth” strategy following our previous report. Prior to deposit Au, the single-crystal Cu substrate was first smoothed by electrochemical polishing in an electrolyte composed of H₃PO₄ and H₂SO₄ with a voltage of 2 V for 2 s. After cleaning, the single-crystal Cu substrate was immediately moved to an electrolyte consisting of 0.05 mM HAuCl₄, 0.1 mM KCl, 0.1 mM K₂SO₄, and 1 mM H₂SO₄. The Au
nanoclusters were epitaxially grown on Cu via electrochemical deposition at −0.3 V versus Ag/AgCl for 30 s. After that, the Au–Cu heterostructure was cleaned by 0.1 M HClO₄ and milli-Q water to remove the residual acid.

**TEM samples preparation and characterization**

Cross-section TEM samples were prepared using FIB on a Thermofisher Helios G4 dual-beam system. Figure S5 schematically showed the procedure to prepare TEM samples by FIB. Pt was deposited on catalyst as a protection layer prior to the ion beam cutting. Typically, a thin Pt layer of 0.5 μm was first deposited using a mild electron beam (2 kV, 0.2 nA), which could eliminate any possible damages on the catalyst surface. Afterward, ion beam deposition (30 kV, 0.43 nA) was applied to deposit a thick Pt layer of 2.0 μm at the same region. Any possible damages from the ion beam deposition could be alleviated because the catalyst surface was already covered by Pt. To this end, the surface configuration of catalyst could be well preserved. Aberration-corrected STEM characterization was performed on a Thermofisher Themis Z microscope equipped with two aberration correctors under 300 kV. HAADF-STEM images were recorded using a convergence semi angle of 11 mrad, and inner and outer collection angles of 59 and 200 mrad, respectively. EDS was carried out using 4 in-column Super-X detectors.

**Other physical characterizations**

The XRD of the Cu foils was characterized by Bruker D2 PHASER. XPS (PHI 5000) measurements were carried out using a Mg Kα source, with the pressure inside the chamber maintained below 4 × 10⁻⁹ Torr and spectra were collected at a pass energy of 17.9 eV. The highest peak in C 1s spectra was shifted to 284.8 eV for charge correction. AFM (Cypher ES, AR) in tapping mode was used for geometry and morphology studies. Raman spectra were taken by a Horiba HR800 system with a laser excitation energy of 638 nm. The laser spot size was 1 μm and a laser power of 5 mW was used to avoid heating.

**Electrochemical CO₂R measurement**

The electrocatalytic characterization was carried out in a customized H cell, which has two compartments separated by an anion exchange membrane (Fumatech company, Fumasep FAA-3-PK130). Each compartment contains 6 mL of 0.1 M KHCO₃ aqueous solution (pH 6.8) and the compartment holding the working electrode is sealed to measure gaseous products. The electrolyte was purged with CO₂ at a constant flow rate of 20 sccm for 20 min before and during each measurement while stirring. A platinum wire and a Ag/AgCl (CHI, 3 M KCl) were used as counter and reference electrodes, respectively. The potentials measured were converted to RHE scale by \( V \text{(versus RHE)} = V \text{(versus Ag/AgCl, 3 M KCl)} + 0.210 \text{ V} + 0.0591 \times \text{pH} \). The solution resistance was compensated for 85% by the potentiostat and the rest 15% was post-corrected.

All electrochemical measurements were explored using a Biologic-SP300 potentiostat. Before electrolysis, linear sweep voltammetry (at a scan rate of 20 mV s⁻¹) was performed and then a set potential was applied for CA. During the 1-h constant potential electrolysis, the gaseous products were quantified using a GC (Agilent 7890B; column HP-PLOT and MS-5A). Sample for GC was collected at 20-min intervals and the separated gaseous products were analyzed by both thermal conductivity detector (for H₂) and flame ionization detector (for CO, methane, and ethylene). Liquid
products were analyzed afterward by 500 MHz NMR (AVANCE III HD) using dimethyl sulfoxide and phenol as internal standards. Solvent pre-saturation technique was implemented to suppress the water peak.

The FE and production rate can be calculated as follows:

**Calculation of the FE of gas products:**

$$\text{FE}(\text{gas}) = \frac{F_{\text{flow}} \times (C_{\text{gas}}/V_m) \times n \times F}{I_{\text{total}} \times 60} \times 100$$

**Calculation of the FE of liquid products:**

$$\text{FE}(\text{liquid}) = \frac{C_{\text{liquid}} \times V \times n \times F}{Q_{\text{total}}} \times 100$$

**Calculation of the production rate of products:**

$$\text{Production Rate} = \frac{\text{FE}_{\text{products}} \times I_{\text{total}}}{n \times F} \times 3,600 \times 1,000$$

**FE (gas):** FE of gas product, %
**FE (liquid):** FE of liquid product, %
**$F_{\text{flow}}$:** flow rate of CO$_2$, mL min$^{-1}$
**$C_{\text{gas}}$:** volume ratio of gas product, determined by on-line GC
**$V_m$:** the molar volume of an ideal gas at 1 atmosphere of pressure (molar volume at normal temperature and pressure is 0.0245 mL), mL mol$^{-1}$
**$C_{\text{liquid}}$:** concentration of liquid product after 1 h of electrolysis, determined by NMR, mol L$^{-1}$
**$V$:** volume of the electrolyte in the working cell, L
**$I_{\text{total}}$:** steady-state cell current
**$Q_{\text{total}}$:** total charge in 1 h of bulk electrolysis, C
**$n$:** number of transferred electrons for certain product
**$F$:** Faradaic constant, 96,485 C mol$^{-1}$

**DFT calculation**

All the calculations were performed using DFT as implemented in the Vienna ab initio simulation package (VASP). The ion-electron interactions were treated with the projected augmented wave (PAW) pseudopotentials, and the plane-wave basis set was cut off at 400 eV. The general gradient approximation (GGA) parameterized by Perdew, Burke, and Ernzerhof (PBE) was adopted to describe the exchange-correlation functional in structural relaxations. All structures were fully relaxed by a conjugate gradient method until the residual force component was less than 0.08 eVÅ. The centered (4 × 4 × 1) k-point grids were used. The constrained Broyden dimer method was used to locate the transition state. The solvent effect was simulated by using the continuum solvation model based on Poisson-Boltzmann equation, as implemented in the VASPsol code. The pressure of CO is set as 1 atm, the electrode potential U at −0.5 V versus RHE. The computational hydrogen electrode (CHE) is utilized to determine the coverage of H atoms at this electrode potential. The Cu (110) facet is covered by 1 ML *H, whereas the Au (110) face are covered by 0.5 ML *H, respectively. The Gibbs free energies ($\Delta G$) of different intermediates were calculated to characterize the CO$_2$R performance on Cu facets.
Finite-element simulation

Finite-element simulations were performed using the COMSOL Multiphysics software package. Three modules were used to establish a comprehensive chemistry-mass transport model of the Au–Cu heterostructure. First, the “Chemistry” module was used to define the tandem CO$_2$R steps. The formation of key CO intermediate from CO$_2$ on Cu and Au surface were both calculated, defined as the feedstock to ethanol generated from Cu substrate. Second, the “Transport of Diluted Species” module was used to solve the mass transport of three species. We determined the diffusion coefficients and rate constants of chemical reactions by sweeping these parameters over a large range (2 orders of magnitude) and fitting to those electrochemical CO$_2$R data. There were six parameters that we swept: three diffusion coefficients, $D_{\text{eq CO}_2}$, $D_{\text{eq CO}}$, and $D_{\text{eq ethanol}}$, as well as the rate constants, $K_{\text{f1}}$ and $K_{\text{f2}}$ of the CO$_2$ to CO on AuCu and Cu shell, and $K_{\text{f3}}$ of the CO-to-ethanol steps.

In situ spectroelectrochemical experiments

In situ FTIR experiments were carried out on a Nicolet 6700 spectrometer with an external mode, equipped with a mercury cadmium telluride detector and $\pi$-polarized light. The customized spectroelectrochemical cell was integrated with a prismatic CaF$_2$ window. The cell was filled with CO$_2$-saturated 0.1 M KHCO$_3$ electrolyte. In the thin layer configuration, each spectrum represented the average of 64 interferograms with a resolution of 4 cm$^{-1}$. The spectra presented with positive bands corresponded to the formation of species, whereas negative bands indicated the consumption of species. All in situ FTIR experiments were performed at an ambient environment, and Ag/AgCl and carbon rod were engaged as the reference and counter electrodes, respectively.

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.chempr.2022.08.016.

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AUTHOR CONTRIBUTIONS

L.Zhang designed and conceived the experiment. C.Z., S.Z., G.S., H.G., J.W., and X.G. fabricated the electrodes and performed the electrochemical characterization and data analysis. Z.Z. and K.L. carried out the synthesis of single-crystal Cu foils. L.Zhou and S.D. carried out the TEM characterization. Y.L. worked on the DFT calculations. C.Y. helped on the FEM modeling and data fitting. C.Z., S.D., and L.Zhang wrote the manuscript. All authors discussed the results and commented on the paper.
DECLARATION OF INTERESTS
The authors declare no competing interests.

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