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Effects of Surface Roughness on the Electrochemical Reduction of CO₂ over Cu

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Abstract

We have investigated the role of surface roughening on the CO$_2$ reduction reaction (CO$_2$RR) over Cu. The activity and product selectivity of Cu surfaces roughened by plasma pretreatment in Ar, O$_2$, or N$_2$ were compared with that of electrochemically polished Cu those. Differences in total and product current densities, the ratio of current densities for HER (the hydrogen evolution reaction) to CO$_2$RR, and the ratio of current densities for C$_2$ to C$_1$ products depend on the electrochemically active surface and are nearly independent of plasma composition. Theoretical analysis of an electropolished and roughened Cu surface reveals a higher fraction of undercoordinated Cu sites on the roughened surface, sites that bind CO preferentially. Roughened surfaces also contain square sites similar to those on a Cu(100) surface but with neighboring step sites, which adsorb OC-COH, a precursor to C$_2$ products. These findings explain the increases in the formation of oxygenates and hydrocarbons relative to CO and the ratio of oxygenates to hydrocarbons observed with increasing surface roughness.
Electrochemical CO$_2$ reduction reaction (CO$_2$RR) offers a promising route for the production of chemicals and fuels using renewable electricity generated from wind and solar energy.$^{1-4}$ Among metallic electrocatalysts evaluated for this purpose, copper is the only metal that produces hydrocarbons and oxygenated products with high Faradaic efficiency (FE).$^{5-10}$ Previous studies have shown that the activity and selectivity of Cu are strongly dependent on the surface morphology of the metal, as well as its local reaction environment (electrolyte composition and pH).$^{11-16}$ For example, Cu(100) and Cu(211) surfaces are more active than Cu(111) surfaces and more effective in promoting C-C bond formation on both single crystal Cu electrodes$^{17, 18}$ and Cu nanoparticles.$^{19, 20}$ Recent theoretical calculations have shown that square sites of Cu(100) bind *OCCO and *OCCHO more strongly than do sites on Cu(111), and that the step sites of Cu(211) facilitate the kinetics of CO dimerization relative to those on Cu(111), resulting in a higher C$_2^+/C_1$ selectivity on square and stepped facets.$^{21, 22}$ The role of other low-coordination Cu sites beyond those present on low-index facets is the subject of continuing discussion. An investigation of the CO$_2$RR over size-controlled Cu nanoparticles has also shown that the population of low-coordination Cu surface sites increases with decreasing Cu particle size (from 15 nm to 2 nm), and that high coordination site result in higher H$_2$ and CO selectivities and lower hydrocarbon (CH$_4$ and C$_2$H$_4$) selectivities.$^{23}$ By contrast, a comparative investigation of CO$_2$RR on (100)-, (111)-, and (751)-preferentially orientated Cu thin film electrodes revealed that both Cu(100) and (751) surfaces with Cu coordination numbers from 6 to 8 promote C-C bond formation relative to more highly coordinated sites on Cu(111) surfaces.$^{24}$ More recently, a study of the electrochemical reduction of CO has suggested that highly porous Cu electrode could favor C$_2^+$ oxygenates selectivity with a large suppression of competitive HER; however, the underlying causes for these observations remains unclear.$^{25}$

Oxide-derived Cu with enriched surface grain boundaries have also been reported to be efficient for reducing CO$_2$ to multi-carbon products.$^{26-29}$ It has been hypothesized that this enhanced C$_2^+$ products selectivity on oxide-derived Cu could arise from surface Cu$^{+}$ retained under reaction conditions and stabilized by subsurface oxygen$^{28, 30}$ or adsorbed halide anions$^{31, 32}$ e.g., I$^-$, based on quasi in situ spectroscopic experiments. However, this hypothesis has been challenged by recent DFT calculations,$^{33, 34}$ $^{^{18}}$O labeling,$^{35}$ electrochemical Raman spectroscopy$^{36, 37}$ and in situ synchrotron XAS and XRD experiments,$^{38, 39}$ suggesting the near-surface oxygen and/or Cu(I) species are not sufficiently long-lived to be present under reaction conditions at highly reducing
cathode potentials. A more plausible interpretation for the enhanced \( C_{2+} \) selectivity of oxide-derived Cu is offered by a recent theoretical analysis of sites present on a rough Cu nanoparticle. This work concludes that surface twin boundaries in the oxide-derived Cu associated with concave defects with respect to Cu(100) planes serve as active sites for C-C bond formation by stabilizing OC-COH species, one of the precursors to \( C_{2+} \) products.

In this study, we investigate the effects of plasma pretreatment of polished Cu foils in different gas atmospheres on their \( CO_2 \)RR activity and selectivity. Since \( O_2 \)-plasma treatment will cause both chemical and physical modifications to Cu surfaces, we pretreated Cu by \( Ar^+ \) ion bombardment in an Ar plasma in order to isolate the effects of surface roughening. These studies show that the changes in the distribution of \( CO_2 \)RR products is attributable to changes Cu surface topography created by the plasma pre-treatments. Roughened Cu surfaces containing a high proportion of under-coordinated Cu sites that exhibit stronger CO adsorption energies than more highly coordinated site present of planar surfaces. Consistent with this finding, the fraction of CO formed from \( CO_2 \) released as CO decreases and the fraction converted to hydrocarbons and alcohols increases with increasing roughness. Our experimental efforts are supported by an analysis of the distribution of sites on a simulated roughened surface of Cu. This work shows that the roughened surface contains a much higher proportion of under-coordinated sites, and in particular sites that adsorb OCCOH, a suggested precursor to \( C_{2+} \) products, more strongly than do the more highly coordinated sites present on a Cu(100) surface.

**Surface Characterization Before and After Plasma Pre-treatment**

Figures 1A and 1B show atomic force microscopy (AFM) and scanning electron microscopy (SEM) images, respectively of the electrochemically polished Cu foil. Fig. 1C presents the AFM topography image of Cu foil after 10-min Ar plasma treatment. In contrast to the polished foil, the Ar plasma-treated foil is much rougher, exhibiting extensive ridges and valleys. The arithmetic surface roughness factor \( (S_a) \) increases from 1.28 nm for the electropolished foil to 7.68 nm after \( Ar^+ \) sputtering based on an AFM scan of a 500 nm x 500 nm area (Fig. S1). SEM images of Ar plasma treated Cu foil before and after \( CO_2 \)RR electrolysis are shown in Fig. 1D. In agreement with the topography determined by AFM, the 2D projection exhibits a surface structure...
covered with pits and islands generated by Ar⁺ bombarding and re-deposition of Cu atoms. Similar surface roughening effects were also observed for N₂ and O₂ plasma treated Cu foils (Fig. S2).

**Figure 1.** Cu electrodes characterization. AFM and SEM images of electrochemically polished Cu foil (A, B) acquired before and after a 10-min plasma pre-treatment in Ar. Images (A) and (C) show the reconstructed 3D topography of the surface obtained from AFM scans, images (B) and (D) show typical SEM images of Cu foils taken before and after 1-h CO₂RR electrolysis at -1.0 V in CO₂-saturated 0.1 M CsHCO₃.

To further quantify the surface roughness of Cu foils electrodes, we determined the electrochemically active surface area (ECSA) by measuring the double-layer capacitances and then calculating the relative roughness of plasma pre-treated Cu compared to electropolished Cu (Table 1 and Fig. S3). In general, all of the plasma treatments increased the roughness of Cu, and prolonged pretreatment time led to a more roughened surface. For the same pretreatment duration, Cu foils treated in either an Ar or O₂ plasma were rougher than that exposed to an N₂ plasma. These differences are likely due to the larger ion size of Ar⁺, the more aggressive etching of O²⁻,⁴³,⁴⁴ as well as the pronounced surface structural rearrangement caused by removal of oxygen from copper oxide during electrochemical reduction.²⁶,³⁴ We also note that plasma pretreatment in an
N$_2$ plasma for more than 10 min did not increase the surface roughness of Cu further, in contrast to what was observed for pretreatment in an O$_2$ plasma.

Table 1. Determined surface roughness of post electrolysis Cu foils with different plasma pretreatments.

<table>
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<th>Ar Plasma 5-min</th>
<th>Ar Plasma 10-min</th>
<th>Ar Plasma 20-min</th>
<th>O$_2$ Plasma 5-min</th>
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Figure 2. Chemical state characterization of Cu electrodes. Ex situ (A) XPS and (B) Raman spectra recorded on Cu foils after 10-min plasma treatments under different gaseous atmosphere.

Figure 2A shows the ex-situ core level X-ray photoelectron spectra (XPS) of the Cu 2p region after plasma treatments in different atmospheres. All four samples show the predominant peak at 932.4 (Cu 2p$_{3/2}$) and 952.2 eV (Cu 2p$_{1/2}$), corresponding to Cu(0) or Cu(I). For the O$_2$ plasma treated Cu, two other satellite peaks show up at ~935 eV and ~944 eV, which are assigned to Cu(II) species. Evidence for Cu(II) cation was also obtained from the Cu LMM region of the Auger spectrum (see Fig. S4). Raman spectra of these four Cu samples are shown in Fig. 2B. No obvious Raman scattering feature is observed for electropolished Cu or Cu exposed to a N$_2$ or an Ar plasma. By contrast, features for both Cu$_2$O (~218, 402 and 526 cm$^{-1}$) and CuO (~290, 332 and 617 cm$^{-1}$) are observed after O$_2$ plasma pretreatment,$^{45}$ in agreement with XPS spectra and Raman studies reported previously.$^{28,32}$
Electrochemical Activity and Selectivity

Figure 3. Electrochemical reduction of CO$_2$ over Cu foil electrodes after 10-min plasma pretreatments under different gaseous atmosphere. (A) Geometric area normalized, (B) ECSA-normalized partial current densities and (C) selectivity of major products during CO$_2$RR over Cu foil electrodes after 10-min plasma pre-treatments under different atmosphere.

The electrochemical CO$_2$RR performance of plasma pre-treated Cu foils was evaluated by 1-h chronoamperometric electrolysis at -1.0 V vs. RHE, for which the CO$_2$ consumption rate is below 17.0 nmol s$^{-1}$ cm$^{-2}$ (Fig. S5).$^{46}$ 0.1 M CsHCO$_3$ was employed as the supporting electrolyte, based upon our previous studies showing that Cs$^+$ cations enhance the field stabilization of the intermediates critical to the formation C$_2$ products$^{13, 14, 47}$ (further evidence for influence of cation identity is given in Fig. S6). The superficial current densities shown in Fig. 3A all increase with plasma pretreatment, in the order of the increasing ECSA. To account for this effect, Fig. 3B shows the current densities for all four samples normalized by the ECSA. The ECSA-normalized current densities for H$_2$ and CH$_4$ are not strongly changed by plasma pretreatment, whereas those for CO and HCOO$^-$ decreases and those for all C$_2^+$ product increases in the order no pretreatment < N$_2$ plasma pretreatment < Ar plasma pretreatment < O$_2$ plasma pretreatment. We note in particular that the rate of CO evolution decreases by more than an order of magnitude upon Ar or O$_2$ plasma pretreated Cu compared to that for electropolished Cu, and the rate of C$_3$ products generation - allyl alcohol and n-propanol - increases by a factor of 3 to 5. The FEs of the principal products of the CO$_2$RR generated on electropolished and plasma pretreated Cu are illustrated in Fig. 3C. After 10-min plasma pretreatment, the selectivity toward hydrogen evolution (HER) and methane generation does not change very much, while the FEs for CO and HCOO$^-$ decrease in the order of
polished Cu > N₂ plasma pretreated Cu > Ar plasma pretreated Cu ≈ O₂ plasma pretreated Cu. By contrast, the FEs for C₂+ products - C₂H₄, C₂H₅OH and n-propanol - follow the reverse trend.

![Diagram](image)

**Figure 4.** CO₂RR products distribution as a function of plasma treatment time. Both (A, B) O₂ and (C, D) Ar atmosphere plasma pretreatments were plotted. The sum of as-determined overall FEs are varied from 92.2% to 98.9% and all normalized to 100% in panels (A) and (C) for comparison.

The similarity of both C₂+ product selectivities and specific activities of Cu after Ar or O₂ pretreatment is particularly interesting. Therefore, we investigated the effects of the duration of plasma pretreatment in Ar and O₂ on the distribution of CO₂RR products. Fig. 4 shows these results for durations of 0, 5, 10, and 20 min, with the electropolished sample referenced as 0-min. The normalized FEs (referenced to 100%) are given in Figs. 4A and 4C. The FE for HER is cyan column, the FE for C₁ products (HCOO⁻, CO and CH₄) is orange column, the FE for C₂ products (C₂H₄ and C₂H₅OH) is purple column, and the FE for C₃ products (allyl alcohol and n-propanol) is magenta column. The selectivity ratio of HER/CO₂RR (black line) and the C₂+/C₁ ratio (red line) are illustrated in Figs. 4B and 4D. We observe similar product distributions throughout the time-
course for both O₂ and Ar plasma pre-treatments, which could largely rule out the (sub)surface oxygen effect. HER decreases to below 30% after 5-min plasma exposure in both cases and levels off with prolonged exposure. The FE for C₁ products also decreases from 0 to 10 min, then increases again due to enhanced methane evolution after a 20-min pretreatment. Over the same pretreatment time, the FE for C₃ alcohols reaches ~10%, whereas the FE for C₂⁺ products rises to ~60%. The ratio of C₂⁺/C₁ reaches a maximum value of ~9 after 10-min of pretreatment in either an Ar or O₂ plasma.

Since the Cu surface roughness increases with prolonged plasma-treatment time, we assessed whether the changes shown in Fig. 4 correlate with surface roughness. The choice of which product ratios to plot is guided by recent theoretical studies of the CO₂RR mechanism, which suggest that 2e⁻ products (HCOO⁻ and CO₂) are produced via the adsorbed intermediates *HCOO and *COOH, respectively, whereas both CH₄ and C₂⁺ products are produced via the reduction of *CO.⁵, 6, 48-5¹ Therefore, as illustrated in Fig. 5A, *CO serves as the key intermediate to CH₄ and C₂⁺ products. Figures 5B and 5C plot the selectivity ratios for C₂⁺ products versus C₁ products (HCOO⁻, CO, and CH₄) and versus CH₄ alone, respectively. The C₂⁺/C₁ ratio reaches a value of ~9 for a roughness of ~3 and then decreases thereafter, regardless of whether Cu is pretreated in an Ar or O₂ plasma. By contrast, the C₂⁺/CH₄ ratio increases monotonically to a value of 30 in the case of Ar plasma pretreatment but reaches a maximum value of ~20 and then decreases in the case of O₂ pretreatment.
Figure 5. CO$_2$RR products distribution as a function of Cu surface roughness. (A) Simplified flowchart of CO$_2$RR mechanism leading to C$_1$ and C$_2$+ products generation, * corresponds to adsorbed species, >2e reduction products of CH$_4$ and C$_2$+ that go through the reduction of *CO intermediate (*COR) are marked in light blue. (B) C$_2$/C$_1$ ratio, (C) C$_2$/CH$_4$ ratio, (D) *COR/(CO$_{(g)}$ + *COR) ratio, and (E) oxygenate/hydrocarbon (only >2e reduction products) ratio.

Figure 5D show that the fraction of CO produced by the CO$_2$RR converted to CH$_4$ and C$_2$+ products increases from 0.68 for a roughness of 1.0 to 0.96 for a roughness of 1.5. Further increasing the roughness to ~4.0 increases this ratio to 0.98. The trend observed in Fig. 5D suggests
that the binding energy for CO increases monotonically with surface roughness, consistent with temperature-programmed desorption experiments\textsuperscript{26} and recent theoretical calculations on simulated Cu particle surface.\textsuperscript{41, 52} The theoretical work also predicts that the stability of \*OC-COH, a critical precursor to C\textsubscript{2} products, increases with surface roughness and in particular with the formation of concave sites at grain boundaries between Cu(100) and Cu(111) surfaces. The downturn in the ratio of FEs for \textsuperscript{C\textsubscript{2+}} product formation and CH\textsubscript{4} formation seen in Fig. 5C when the roughness exceeds ~3.0 for O\textsubscript{2} plasma pretreated Cu might be attributable to the formation of sub-nanometric Cu clusters (i.e., surface dimer and trimer adatoms) that could serve as active sites for selective CO\textsubscript{2}-to-CH\textsubscript{4} conversion.\textsuperscript{53} Finally, Fig. 5E shows the surface roughness dependence of the ratio of FEs for \textsuperscript{C\textsubscript{2+}} oxygenated products to that for \textsuperscript{C\textsubscript{2+}} hydrocarbons. The selectivity to \textsuperscript{C\textsubscript{2+}} oxygenated products (ethanol, allyl alcohol and n-propanol) increases by a factor of two with increasing roughness, most likely due to the lower surface coverage by \*H, as reflected in the HER trend of Fig. 4, and the reduced likelihood of hydrogenating C–C intermediates on the roughened Cu surfaces. Similarly, the selectivity to HCOO\textsuperscript{−} decreases as the roughness increases from 1.0 to 2.6, and levels off thereafter (Fig. S7). We suggest that this trend is due to the lower availability of terrace sites on roughened Cu surfaces, which are needed to bond bidentate \*HCOO, the precursor to HCOOH and hence HCOO\textsuperscript{−} upon desorption of HCOOH into the alkaline electrolyte.\textsuperscript{6}

A further factor that can contribute to enhancing the ratio of \textsuperscript{C\textsubscript{2+}}/CH\textsubscript{4} on roughened vs smooth Cu surfaces is the higher local pH near the surface of the roughened Cu; therefore, we considered the possible effect of pH on the observed results. The higher pH near the roughened surface is ascribable to the higher rate of OH\textsuperscript{−} generation rate per geometric electrode area. We note that this proposal is consistent with the control experiment presented in Fig. S8, illustrating the effects of increased bicarbonate concentration, and the earlier findings of Hori et al., who reported a 4-fold enhancement in the \textsuperscript{C\textsubscript{2+}}/CH\textsubscript{4} ratio upon increasing the surface pH from 8.5 to 9.5.\textsuperscript{54} In our studies, we observed only a ~2.6-fold increase in the \textsuperscript{C\textsubscript{2+}}/CH\textsubscript{4} ratio due to an estimated increase in the surface pH change from 9.76 to 9.88 for Cu treated for 5-min vs. 10-min in an Ar plasma. The pH for these experiments was estimated assuming a mass-transfer boundary layer thickness of ~40 µm and a bicarbonate concentration of 0.1 M.\textsuperscript{46, 55} We also note that it has been reported that the increase in surface pH as a consequence of electrolyte polarization should suppress CH\textsubscript{4} production, but keep \textsuperscript{C\textsubscript{2+}} activity constant at a given potential.\textsuperscript{56, 57} Figure 3B shows that while the intrinsic activity (ECSA-normalized partial current density) for forming CH\textsubscript{4} decreases following plasma
pretreatment, the intrinsic activities for producing \( \text{C}_2^+ \) products, especially oxygenated products, increases. These observations suggest that the enhanced \( \text{C}_2^+ \) selectivity observed after plasma treatment is not attributable to the small change in the local pH.

Table S1 compares our results on the effects of surface roughness with those previously reported. Various methods have been used to roughen the surface of Cu. These include reduction of copper oxides produced by thermal and/or plasma oxidation, potential cycling in halogen-containing solutions, dendrite growth, etc. Regardless of the method used to achieve roughening, increased surface roughness generally leads to an enhanced \( \text{C}_2^+ \) FE and \( \text{C}_2^+/\text{C}_1 \) ratio. Most authors have attributed these effects enhanced surface roughness to a greater abundance of under-coordinated surface sites and defective sites that bind \( \text{^*CO} \) strongly and promote its further reduction to C-C bond formation. Unfortunately, these results cannot be compared with those reported here because of large difference in the modes of sample preparation and conditions for their investigation.

Theoretical Simulation and Analysis

Insights into why surface roughening causes an increase in the formation of \( \text{C}_2^+ \) products can be gained by atomic-level analysis of the Cu surface created by computational simulation of Ar\(^+\) bombardment roughening of a Cu(111) surface followed by thermal relaxation of the resulting surface. The simulation of roughening begins with a 17.6 nm \( \times \) 17.4 nm surface that 10.1 nm thick, which contains \( \sim \) 5,422 total surface atoms (see Fig. 6A). Figure 6B and the Supporting Movie 1 show that after bombardment of this surface with 1300 Ar\(^+\) cations, the Cu surface exposes 10,433 atoms, corresponding to an increase in surface roughness of 1.93. The simulated untreated Cu surface and the plasma-treated surface resemble the topography and roughness of the experimental observations, as seen in Figure 1.
Figure 6. Active site visualization by ReaxQM-Machine Learning. Images of the computationally produced Cu surface of (A) electro-polished and (B) after Ar plasma bombardment, and (C) predicted distribution of CO adsorption energies, $\Delta E_{\text{CO}}$. The three dashed lines indicate the CO adsorption energies on Cu(111), (100) and (211).

The ReaxQM-Machine Learning approach was used to predict the distribution of CO binding energies, $\Delta E_{\text{CO}}$, for both simulated electrochemically polished Cu and the Ar plasma roughened surfaces.\textsuperscript{41, 58} The electropolished Cu, shown in Fig. 6A, is dominated by close-packed low index sites, and, therefore, the mean values of $\Delta E_{\text{CO}}$ are close to those of Cu(111), (100) and (211) surfaces (Fig. 6C). By contrast, a much higher population of stronger CO binding sites, ranging from -1.10 to -1.54 eV (red sites along the ridges in Fig. 6B and the columns left of the Cu(211) line in Fig. 6C), appear after plasma pretreatment. The increased number of strong CO binding sites is consistent with the experimental observation of a reduced selectivity to CO formation following plasma pretreatment to roughen the catalyst surface. Since the adsorption of *CO and *H are expected to compete on the surface of Cu, the stronger adsorption of CO on roughened Cu is expected to result in a reduction of the *H/*CO. This reasoning would explain the increase in
the formation of oxygenated relative to hydrocarbon products observed with increasing roughening seen in Fig. 5E.

**Figure 7.** Visualization of square sites on computationally simulated Cu surfaces. (A) Electropolished Cu and (B) Ar plasma pretreated Cu, the surface square sites are marked in blue color. (C) Zoomed-in atomic structure of randomly chosen square sites on the plasma-treated Cu surface, both the square sites and the neighboring Cu step sites (if any) are highlighted in red.

In our previous work we have employed the formation energy of *OC-COH (ΔE_{OC-COH})* as a descriptor for the selectivity of C\(_{2+}\) products formed during the CO\(_2\)RR, and have shown that the mean value of ΔE_{OC-COH} decreased from ~1.35 eV for random surface sites to ~0.50 eV for surface square sites.\(^{39}\) We also noted that concave defects located at Cu(100) plane next to a (111) step exhibited the most favorable values of ΔE_{OC-COH} for C\(_{2+}\) selectivity. In the present study, we identified the surface square sites on the simulated Cu surfaces for comparison as shown in Fig. 7A and 7B. The percentage of square sites to overall surface atoms increased from 20.9% on the
polished Cu surface to 38.5% on the surface produced by Ar plasma pretreatment. Fig. 7C shows the atomic structures of randomly chosen square sites occurring on the simulated surface produced by plasma treatment. The sites are similar to the Cu(100) configuration but have abundant neighboring step sites. As noted, the significantly increased percentage of surface square sites favors C-C bond formation and provides a rational for the enhanced C$_2^+$ selectivity that correlates with increased surface roughening caused by plasma pretreatment.

In conclusion, the present study investigated the effects of surface topography on the activity and product selectivity of electrochemical CO$_2$ reaction, demonstrating the distribution of products produced by CO$_2$RR on metallic Cu changes with Cu surface topography created by plasma pretreatment. The activity and product selectivity of electrochemically polished Cu was compared with those of roughened Cu surfaces prepared by different plasma pretreatments. The differences in overall CO$_2$RR activity from different plasma pretreatments are attributable changes in the electrochemically active surface area. Of particular note, we observe that with increasing surface roughness, the ratio of current densities for CO$_2$RR to HER, the ratio of the current densities for COR to the sum of the current densities for CO formation and COR, and the ratio of current densities for formation oxygenates to hydrocarbons all increase. These trends are interpreted based on an atomic-level analysis of the topography of roughened Cu surface. We find that CO formed by the electrochemical reduction of CO$_2$ binds more strongly on the roughened surface than on the electrochemically polished surface, suggesting that the ratio of adsorbed H to CO decreases with increasing surface roughness. These trends explain why the fraction of CO produced by CO$_2$ reduction converted to reduced products and the fraction of these products appearing as oxygenates rather than hydrocarbons increase with increasing surface roughness. A further effect of increasing surface roughness is an increase in the fraction of final products (oxygenates and hydrocarbons) containing two or more C atoms. Our analysis shows that surface roughening increases the fraction of square sites similar to those on a Cu(100) surface but having abundant neighboring step sites. Prior work has shown that such sites enhance the formation of C-C bonds required to form C$_2^+$ products during the electrochemical reduction of CO$_2$. Finally, we find that increased roughening of the Cu surface increases the ratio of current densities for C$_2^+$ to C$_1$ (CO, HCOO$,^-$, and CH$_4$) products up to a maximum value of ~ 9 for a surface roughness of ~ 3. Most of this downturn is ascribed to the enhanced formation of CH$_4$ on surfaces with roughness in excess of 3. The enhanced formation of methane on highly roughened surfaces is attributed to the formation of
surface dimer and trimer clusters on the Cu surface that have been shown by theoretical calculation
to serve as active sites for selective conversion of CO$_2$ to CH$_4$. In summary, the results of the
present work highlight the importance of surface topography and defect sites on the observed CO$_2$
reduction activity and selectivity, and suggest that rational surface structure engineering could
contribute to the development of Cu electrocatalysts exhibiting a high selectivity to multi-carbon
products.

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: xxx

Experimental and computational details; additional Cu electrodes characterization by AFM, SEM,
Auger, XPS and electrochemical double-layer capacitance measurements; Cu surface roughness
dependence of CO$_2$ consumption rate and FE$_{\text{formate}}$/FE$_{\text{(CO$_2$+COR)}}$ ratio during chronoamperometric
electrolysis; online differential electrochemical mass spectrometry study on the cation effect;
control experiments study on anion buffering effect and extra surface roughening at the presence
of iodide anion

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Table of Contents Graphic