The electroreduction of C1 feedgas to high-energy-density fuels provides an attractive avenue to the storage of renewable electricity. Much progress has been made to improve selectivity to C1 and C2 products, however, the selectivity to desirable high-energy-density C3 products remains relatively low. We reason that C3 electrosynthesis relies on a higher-order reaction pathway that requires the formation of multiple carbon-carbon (C-C) bonds, and thus pursue a strategy explicitly designed to couple C2 with C1 intermediates. We develop an approach wherein neighboring copper atoms having distinct electronic structures interact with two adsorbates to catalyze an asymmetric reaction. We achieve a record n-propanol Faradaic efficiency (FE) of (33 ± 1)% with a conversion rate of (4.5 ± 0.1) mA cm$^{-2}$, and a record n-propanol cathodic energy conversion efficiency ($EE_{\text{cathodic half-cell}}$) of 21%. The FE and $EE_{\text{cathodic half-cell}}$ represent a 1.3× improvement relative to previously-published CO-to-n-propanol electroreduction reports.
The CO₂ electroreduction reaction (CO₂RR) to high-energy-density liquid products is an attractive avenue to achieving the storage of renewable energy. In recent years, much progress has been made in CO₂RR, but the main products reported have been C₁ (CO, CH₄, methanol, and formate) and C₂ (ethylene, acetate, and ethanol) products. n-propanol, a high-value and high-energy-density C₃ product that can be generated via either CO₂ or CO electroreduction, has been produced with low-to-moderate Faradaic efficiencies (FEs) in prior reports.

In CO₂RR, the formation of multi-carbon products starts with the formation of the CO intermediate, followed by the CO electroreduction process. Recently, significant progress has been made in CO₂RR to CO with a FE nearly 100%. The ultimate goal of high selectivity to high-value-added C₃ products from CO₂RR, it is of interest to improve significantly the FE for CO electroreduction (CORR) to C₃ products.

The formation of C₃ products from CORR relies on the sequential formation of two carbon–carbon (C–C) bonds, the main reaction mechanism for C₃ formation reported previously. Cu provides excellent C–C coupling and produces multi-carbon chemicals in the electroreduction of CO; however, the selectivity towards C₃ products on Cu has remained low. The generation of C₂ products from CO requires multiple product/intermediate formation steps, and it is prone to the competing production of a wide variety of chemical products.

Increasing selectivity in the electroreduction of CO to C₃ products is thus an important challenge to address in the field of electrocatalysis. Until now, catalysts for CORR have focused on Cu and oxide-derived Cu catalysts, and a number of factors have been found to increase performance: these include engineering the oxidation state of the atoms making up the metal catalyst, as well as grain-boundary effects and the selective formation of the desired facets. However, the main products of these Cu and oxide-derived Cu catalysts have been C₂ chemicals (ethanol, acetate, and ethanol), and the selectivity to C₃ products has saturated in recent manuscripts based on Cu catalysts.

Here we explore instead a doping strategy involving different metal-doped Cu (M-doped Cu) catalysts in an attempt to increase C₃ production in CORR. The low C₃ selectivity on Cu catalysts is associated with the low rate of C–C bond formation, including C₁–C₁ and C₁–C₂ coupling. Mechanisms underpinning C₁–C₁ coupling to C₂ products have been explored extensively in prior studies, while C₁–C₂ coupling to C₃ products is less explored. Here we screen the propensity to catalyze C₁–C₁ and C₁–C₂ coupling using density functional theory (DFT) based on M-doped Cu catalysts. We find that, among different M-doped Cu candidates explored, Ag-doped Cu is expected to offer the highest activity for both C₁–C₁ and C₁–C₂ coupling, and we pinpoint a role for the asymmetric C–C coupling active site in this high activity. Specifically, this active site consists of two neighboring Cu atoms that exhibit different electronic structures: this asymmetry among the neighbors’ energetics is determined by the combination of strain and ligand effects arising upon Ag doping. We then fabricate Ag-doped Cu nanocatalysts via a galvanic replacement approach. We demonstrate that the synthesized Ag-doped Cu catalyst exhibits higher FEs for n-propanol compared to all previous CORR reports. This leads to superior energy conversion efficiency in the cathodic half-cell (EEcath = feed concentration vs. Cu). Several M-doped Cu systems (M = Ag, Au, Ru, Rh, and Pd) were considered because bimetallic catalysts have been shown to tune catalyst performance in other catalytic reactions. CO dimerization is one reaction pathway for C₁–C₁ coupling, so we used the barrier of CO dimerization to describe the readiness of C₁–C₁ coupling. Due to the abundance of CO species in CORR, we used the barrier of COCO and CO coupling as the indicator for the C₁–C₂ coupling (Supplementary Figs. 1–14 and Supplementary Tables 1–3). As shown in Fig. 1a, among the M-doped Cu systems studied, calculation results show that Ag-doped Cu possesses the lowest activation energies for both C₁–C₁ and C₁–C₂ coupling, suggesting that Ag-doped Cu is a promising catalyst for the formation of C₃ products from CO.

We carried out further theoretical investigations to uncover the physical origins of enhancement in C₂–C₁ and C₂–C₂ coupling on Ag-doped Cu. We used a model with one Ag atom doped in a Cu (111) slab with a 3 × 3 unit cell, which corresponds to about 3% doping concentration for four layers of Cu atoms. By a margin of 0.63 eV, Ag at the surface of Cu(111) is more favorable than in...
subsurface of Cu(111), inducing us to focus on surface-localized Ag in our ensuing studies.

As the radius of the Ag atom is larger than that of Cu atom, Ag doping produces surface strain. The bond length of Cu–Cu changed from 2.57 Å on the Cu(111) surface to 2.55 and 2.48 Å on the Ag-doped Cu surface, resulting in asymmetric compressive strain (Supplementary Fig. 15). The ligand effect caused by Ag doping in Cu also has the potential to affect C1–C1 and C1–C2 coupling. To evaluate the effect of the strain and ligand, we built a model with the same strain of Ag-doped Cu but without Ag substitution by fixing the bond length of Cu surface (denoted Cu with strain). We calculated the activation energies for C1–C1 and C1–C2 coupling steps on Cu, Cu with strain, and Ag-doped Cu models (Supplementary Table 4). Calculation results show that both compressive strain and ligand effects contribute to enhanced activity for C1–C1 and C1–C2 coupling (Supplementary Fig. 16a).

Ag doping in Cu leads to two classes of neighboring Cu atoms (denoted Cu-a and Cu-b atoms, Supplementary Fig. 16b). The two classes exhibit distinct electronic structures, as a result of strain and ligand effects. As to the strain effect, the bond length between Cu-a and Cu-b atoms is 2.55 Å, while the bond length between two Cu-b atoms is 2.48 Å (Supplementary Fig. 15), suggesting that Cu-a is more isolated than Cu-b. As to ligand effects, Cu-a coordinates with nine Cu atoms, while Cu-b coordinates with eight Cu atoms and one Ag atom (Supplementary Fig. 16). We term a pair of adjacent Cu atoms with different electronic structures an asymmetric C–C coupling active site. In this reaction, the asymmetric site interacts with two CO to yield asymmetric reactants—two adsorbed CO on Cu-a and Cu-b atoms with different electronic structures—enhancing C1–C1 coupling. In light of the similarity with C–C coupling, the same site can further promote C1–C2 coupling between asymmetric C1 and C2 intermediates (Supplementary Fig. 17). These findings are in ways analogous to the enhanced coupling effect of Cu6 and Cu + proposed by Goddard and co-workers. Taken together, these DFT calculation results suggest that Ag-doped Cu with asymmetric C–C coupling active sites is a good candidate for C3 production in CORR which appear to support both C1–C1 dimerization and C1–C2 coupling. It is worth mentioning that there exist other C1–C2 coupling possibilities: we also examined another C2–C2 coupling mechanism, OC-OCCOH (refs. 33,39), and found that Ag-doped Cu had a barrier of 0.76 eV, lower than 0.88 eV on Cu (Supplementary Figs. 18, 19, and Supplementary Table 5).

**Preparation and characterization of nanocatalysts.** We sought to prepare experimentally Ag-doped Cu catalysts. We employed a galvanic replacement reaction driven by the difference in the reduction potential of Ag vs. Cu (ref. 46). Firstly, we deposited a thin layer of commercial Cu nanoparticles with average size of 100 nm on a carbon-based gas diffusion layer (GDL) via spray-coating (Supplementary Fig. 20a). The Cu gas diffusion electrode (GDE) was then immersed in N2-saturated 5 μmol L−1 AgNO3 aqueous solution at 65°C for 1 h to obtain the Ag-doped Cu GDE. The Ag-doped Cu catalyst retains the particle size and the morphology of the pristine Cu nanoparticles (Fig. 2a, b, and Supplementary Figs. 20 and 21). Electron energy loss spectroscopy (EELS) elemental mapping showed that Ag and Cu elements were uniformly distributed in the particle (Fig. 2c and Supplementary Fig. 22). In transmission wide angle X-ray scattering (WAXS) data and powder X-ray diffraction (XRD) patterns, we observed peaks for Cu2O in both Cu and Ag-doped Cu GDE (Fig. 2d, e, and Supplementary Fig. 23), which were attributed to oxidation of Cu in air during the preparation of GDE. Due to the low concentration of Ag (4.0% atomic fraction determined by XPS), a peak shift relative to Cu is not observed in WAXS data of the Ag-doped Cu GDE (Fig. 2e and Supplementary Table 6). X-ray photoelectron spectroscopy (XPS) of the Ag-doped Cu GDE also showed the existence of CuO on the surface and confirmed the presence of Ag0 (Fig. 2f, g).

**CORR performance and Operando X-ray absorption spectroscopy.** We then investigated the performance of the Ag-doped Cu GDE in CORR flow cell reactors (Supplementary Fig. 24). Flow cells overcome the mass transfer limitation of CO and produce a triple-phase interface that allows the gas reactant to contact the catalyst-electrolyte interface during the reaction. Figure 3a shows FE for C2+ products in the applied potential range of −0.36 to −0.56 V with reference to the reversible hydrogen electrode (RHE) in 1 M KOH electrolyte. The liquid products (n-propanol, ethanol, and acetate) and gas products (ethylene and H2) were quantified using nuclear magnetic resonance (NMR) and gas chromatography, respectively (Supplementary Fig. 25 and Supplementary Table 7). In the applied potential range of −0.36 to −0.56 VRHE, the total C2 FS on Ag-doped Cu GDE are higher than that on Cu GDE: indeed the total FE of C2+ products on Ag-doped Cu GDE reaches about 80% at −0.56 VRHE. In particular, at a low potential of −0.46 VRHE, the Ag-doped Cu GDE records a high n-propanol FE of (33 ± 1%) with the partial n-propanol current density of (4.5 ± 0.1) mA cm−2, whereas n-propanol FE on pristine Cu is (22 ± 1%) (Fig. 3b). This impressive n-propanol FE represents the highest value reported for n-propanol production via CO2RR and CORR (Supplementary Table 8). The higher FE for C2+ and C3 products on Ag-doped Cu relative to Cu is consistent with predictions from DFT. The intrinsic activities for n-propanol production on Ag-doped Cu and Cu are reported via the partial current density for n-propanol formation normalized to the electrochemical surface area (ECSA) (Supplementary Figs. 26, 27, and Supplementary Table 9). The ECSA-normalized partial n-propanol current density on Ag-doped Cu is 0.124 mA cm−2, which is 3 times that on Cu. The n-propanol EEcathodic half-cell reaches 20% at a low potential of −0.46 VRHE when the overpotential of oxygen evolution in anode side is assumed to be 0. After correcting for ohmic loss (Supplementary Table 10), the n-propanol EEcathodic half-cell reaches 21% under a low overpotential of 0.616 V. This EEcathodic half-cell is higher than the best prior reports by a margin of 1.3× (Supplementary Sections 3, 4, and Supplementary Table 8).

It should be noted that the highest n-propanol FE on both Ag-doped Cu and Cu GDEs were achieved at relatively low potential (−0.46 VRHE), and n-propanol FE decreased when further increasing the potential to −0.56 VRHE (Fig. 3a and Supplementary Table 7). The total C2 product and ethylene FE on both Ag-doped Cu and Cu GDEs exhibited an increasing trend with increased applied potentials. This result can be explained by noting that the C–C coupling step for n-propanol formation becomes slow at high potential, and thus C2 intermediate protonation reaction is more favored compared with C–C coupling step for n-propanol formation26,34.

To study the chemical state of the catalyst during CORR, we performed operando X-ray absorption spectroscopy (XAS) of Ag-doped Cu GDE under operando CORR conditions at a constant applied potential of −0.46 VRHE. Operando Cu K-edge X-ray absorption near edge structure (XANES) spectra of Ag-doped Cu GDE show that Cu atoms were reduced to Cu0 in the first 10 s during CORR (Fig. 3c and Supplementary Fig. 28). Thereafter, the valence state of Cu is maintained at zero throughout CORR. Consistent results are shown in the operando extended X-ray adsorption fine structure (EXAFS). An EXAFS fitting analysis at the Cu K-edge showed the presence of Ag and its strong
interaction with Cu by having Cu–Ag distance value between pure Cu and pure Ag (Fig. 3d and Supplementary Table 11). Collectively, operando XAS results demonstrated that Ag atoms are doped in the lattice of Cu nanoparticles, and that each element remains in its metallic state during CORR. As a control experiment, we carried out operando Cu K-edge XANES on Cu GDE, and it also showed that Cu oxides were reduced to Cu0 in the first 10 s during CORR. These results further demonstrated that, rather than benefiting from oxidation states, the high FE of n-propanol on Ag-doped Cu GDE is associated with metallic states of Cu and might be ascribed to the different structures of Ag-doped Cu relative to Cu.

To explore further the role of Ag doping in Cu in facilitating C1–C2 and C1–C2 coupling, we investigated the CORR performance of Ag-doped Cu catalysts with different Ag concentrations. To vary the Ag concentration, we changed the immersion time of Cu with an atomic percentage of 2.4% Ag showed the highest n-propanol FE at -0.46 V_RHE (Supplementary Fig. 34). The analysis of operando Cu K-edge XANES of Cu–Ag-20 min and Cu–Ag-2 h GDE also demonstrates that Cu oxides are reduced to Cu0 in the first 10 s during CORR (Fig. 4c) and it is Ag-doped Cu in metallic state that is associated with the enhanced selectivity to n-propanol during CORR.

The carbon-based GDLS suffer from liquid penetration and gas diffusion blockage, termed flooding, over time. To overcome the flooding issue on carbon-based GDE after long operation time, we fabricated the Ag-doped Cu polytetrafluoroethylene (PTFE) electrode based on a configuration (graphite/carbon nanoparticle/Ag-doped Cu/PTFE electrode) developed by our group. The Ag-doped Cu layer was prepared by immersing a Cu layer in 5 μmol L\(^{-1}\) AgNO\(_3\) aqueous solution at 65 °C for 1 h. When a potential of -0.46 V_RHE was applied, the FE of n-propanol on the Ag-doped Cu PTFE electrode achieved 33% and operated stably over 200 min of CORR (Supplementary Figs. 35 and 36). We also compared the CO\(_2\)RR performance of Ag-doped Cu and pristine Cu to explore whether the asymmetric active sites enhance in this distinct context the C1–C2 and C1–C2 coupling (Supplementary Fig. 37 and Supplementary Table 12). In the
potential range from $-0.82$ to $-1.79$ V$_{\text{RHE}}$ (after $iR$ compensation), both C$_2^+$ and C$_3$ FEs on Ag-doped Cu catalysts are notably higher than those on pristine Cu at the potential of $-2.96$ V$_{\text{RHE}}$ ($-1.31$ V$_{\text{RHE}}$ after $iR$ compensation), the partial C$_2^+$ and n-propanol current densities of Ag-doped Cu are 308 ± 6 mA cm$^{-2}$ and 36 ± 2 mA cm$^{-2}$, and C$_2^+$ and n-propanol FEs on Ag-doped Cu are 62 and 7%, respectively, providing a doubling compared to pristine Cu.
Discussion
This work demonstrates Ag doping in Cu to facilitate C1–C3 and C1–C2 coupling and thus improve the selectivity to C3 products during CORR. DFT results show that the strain and ligand effects due to Ag doping jointly provide an asymmetric C–C coupling active site containing two neighboring Cu atoms with different electronic structures, and that these are capable of enhancing C1–C2 and C1–C2 coupling. Experimentally, we achieved a total C2+ FE of about 80% and a record n-propanol FE of (33 ± 1)% with a partial n-propanol current density (4.5 ± 0.1) mA cm−2 on Ag-doped Cu catalyst in CORR. The EEcathodic half-cell for n-propanol also reaches 21% at a low potential of 0.416 VRHE, with anode were circulated by two pumps at the rate of 10 mL min−1 between membrane and cathode, respectively. The electrolytes in cathode and anode chamber between anode and membrane, as well as the cathode chamber and the back side of cathode GDE at the rate of 50 mL min−1. Gas could diffuse into the interface between cathode and electrolyte, thus generating a triple-phase interface between gas, electrode, and electrolyte. The catalytic performance of cathodes was evaluated by performing potentiostatic electrolysis.

Methods
DFT calculations. In this work, all the DFT calculations were carried out with a periodic slab model using the Vienna ab initio simulation program (VASP) (https://www.vasp.at/). Detailed theoretical methods are found in the Supplementary Information.

Preparation of electrodes. A commercial Cu of 8.5 mg was dispersed in a mixture of 0.85 mL of methanol and 8.5 μL of 5% Nafion under ultrasonication for 30 min. The suspension was deposited on a carbon-based GDL using spray-coating with a catalyst loading of ~1 mg cm−2 to prepare the Cu GDE. We then immersed the prepared Cu GDE in 5 mol L−1 AgNO3 aqueous solution at 65 °C for a certain time period to prepare Ag-doped Cu GDE as cathodes. The main goal of the work was to focus on improving the efficiency of the cathodic side of CORR to propanol. Thus, we used Ni foam (1.6 mm thickness, MTI Corporation) as the oxygen electrode. The ORR performed in the flow cell reactor. The prepared GDEs, anion exchange membrane, and nickel foam were positioned and clamped together between silicone gaskets and PTFE fi-

Data availability
The data supporting this study are available in the paper and the Supplementary Information. All other relevant source data are available from the corresponding authors upon reasonable request.

Received: 15 February 2019; Accepted: 22 October 2019;
Published online: 29 November 2019

References
1. Liu, M. et al. Enhanced electrocatalytic CO2 reduction via field-induced reagent concentration. Nature 537, 382–386 (2016).
2. Gao, S. et al. Partially oxidized atomic cobalt layers for carbon dioxide electroreduction to liquid fuel. Nature 529, 68–71 (2016).
3. Seifotokaldani, A. et al. Hydronium-induced switching between CO2 electroreduction pathways. J. Am. Chem. Soc. 140, 3833–3837 (2018).
4. Le, M. et al. Electrochemical reduction of CO2 to CH2OH at copper oxide surfaces. J. Electrochem. Soc. 158, E45–E49 (2011).
5. Zhang, W. et al. Electrochemical reduction of carbon dioxide to methanol on hierarchical Pd/SnO2 nanosheets with abundant Pd–O–Sn interfaces. Angew. Chem. Int. Ed. 57, 9475–9479 (2018).
6. Braeken, A. et al. Copper nano-catalysts for CO2 electroreduction to fuel. Science 363, 651–675 (2014).
7. Dinh, C.-T. et al. CO2 electroreduction to ethylene via hydroxy-mediated copper catalysis at an oxide-supported copper interface. Science 360, 783–787 (2018).
8. Mistry, H. et al. Highly selective plasma-activated copper catalysts for carbon dioxide reduction to ethylene. Nat. Commun. 7, 12123 (2016).
9. Liu, Y., Chen, S., Quan, X. & Yu, H. Efficient electrochemical reduction of carbon dioxide to acetate on nitrogen-doped nanodiamond. J. Am. Chem. Soc. 137, 11631–11636 (2015).
10. Zhuang, T.-T. et al. Steering post-C–C coupling selectivity enables high efficiency electroreduction of carbon dioxide to multi-carbon alcohols. Nat. Commun. 1, 421–428 (2018).
11. Ren, D., Ang, B. S.-H. & Yeo, B. S. Tuning the selectivity of carbon dioxide electroreduction toward ethanol on oxide-derived Cu2Zn catalysts. ACS Catal. 8, 8239–8247 (2014).
12. Kim, D., Kley, C. S., Li, Y. & Yang, P. Copper nanoparticle ensembles for selective electroreduction of CO2 to C2–C3 products. Proc. Natl Acad. Sci. USA 114, 10560–10565 (2017).
13. Hoang, T. T. H. et al. 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 (2018).
14. Jouny, M., Lue, W. & Jiao, F. General techno-economic analysis of CO2 electrolysis systems. Ind. Eng. Chem. Res. 57, 2165–2177 (2018).
15. Clark, E. L., Hahn, C., Jaramillo, T. F. & Bell, A. T. Electrochemical CO2 reduction over compressively strained CuAg surface alloys with enhanced multi-carbon oxygenate selectivity. J. Am. Chem. Soc. 139, 15848–15857 (2017).
16. Verdaguer-Casadevall, A. et al. Probing the active surface sites for CO2 reduction on oxide-derived copper electrocatalysts. J. Am. Chem. Soc. 137, 9808–9811 (2015).
17. Handoko, A. D. et al. Mechanistic insights into the selective electroreduction of carbon dioxide to ethylene on CuOx-Derived copper catalysts. J. Phys. Chem. C 120, 20038–20047 (2016).
18. Ren, D. et al. Mechanistic insights into the enhanced activity and stability of agglomerated Cu nanocrystals for the electrochemical reduction of carbon dioxide to n-propanol. J. Phys. Chem. Lett. 7, 20–24 (2016).
19. Rahaman, M., Dutta, A., Zanetti, A. & Broekmann, P. Electrochemical reduction of carbon dioxide at activated Cu mesh catalysts: an identical location (IL) study. ACS Catal. 7, 7946–7956 (2017).
20. Jiang, K. et al. Metal ion cycling of Cu foil for selective C–C coupling in electrochemical CO2 reduction. Nat. Catal. 1, 111–119 (2018).
21. Li, C. W., Ciston, J. & Kanan, M. W. Electroreduction of carbon monoxide to liquid fuel on oxide-derived nanocrystalline copper. Nature 508, 504–507 (2014).
22. Feng, X., Jiang, K., S. & Kanan, M. W. A direct grain-boundary-activity correlation for CO electroreduction on Cu nanoparticles. ACS Cent. Sci. 2, 169–174 (2016).
23. Raciti, D. et al. Low-overpotential electroreduction of carbon monoxide using copper nanowires. ACS Catal. 7, 4467–4472 (2017).
24. Li, J. et al. Copper adparticle enabled selective electrosynthesis of n-propanol. Nat. Commun. 9, 4614 (2018).
25. Zhuang, T.-T. et al. Copper nanocavities confine intermediates for efficient electrosynthesis of C3 alcohol fuels from carbon monoxide. Nat. Catal. 1, 946–951 (2018).
26. Joump, M., Luc, W. & Jiao, F. High-rate electroreduction of carbon monoxide to methane/oil on carbon nanotubes. Nat. Catal. 1, 748–755 (2018).
27. Kortlever, R., Shen, J., Schouten, K. J. P., Calle-Vallejo, F. & Koper, M. T. M. Catalysts and reaction pathways for the electrochemical reduction of carbon dioxide. J. Phys. Chem. Lett. 6, 4073–4082 (2015).
28. Cheng, T., Xiao, H. & Goddard, W. A. Full atomistic reaction mechanism with kinetics for CO reduction on Cu(100) from ab initio molecular dynamics free-energy simulations at 298 K. Proc. Natl Acad. Sci. USA 114, 1795–1800 (2017).
29. Montoya, J. H., Shi, C., Chan, K. & Nørskov, J. K. Theoretical insights into a CO dimerization mechanism in CO2. J. Phys. Chem. Lett. 6, 2032–2037 (2015).
30. Liu, S. et al. Shape-dependent electrocatalytic reduction of CO2 to CO on triangular silver nanoplates. J. Am. Chem. Soc. 139, 2160–2163 (2017).
31. Jhong, H.-R. M. et al. A nitrogen-doped carbon catalyst for electrochemical CO2 to C2H4 reduction. ACS Catal. 7, 324–330 (2017).
32. Kortlever, R., Shen, J., Schouten, K. J. P., Calle-Vallejo, F. & Koper, M. T. M. Identical Location (IL) study. J. Phys. Chem. Lett. 7, 1471–1477 (2016).
33. Xiao, H., Cheng, T. & Goddard, W. A. Atomicistic mechanisms underlying selectivities in C1 and C2 products from electrochemical reduction of CO on Cu (111). J. Am. Chem. Soc. 139, 130–136 (2016).
34. Kuhl, K. P., Cave, E. R., Abram, D. N. & Jaramillo, T. F. New insights into the electrochemical reduction of carbon dioxide on metallic copper surfaces. Energy Environ. Sci. 5, 7050–7059 (2012).
35. Vayenas, C. G., White, R. E. & Gamboa-Aldeco, M. E. Modern Aspects of Electrochemistry. Electrochemical CO2 Reduction. Vol. 42 (Springer, New York, 2008).
36. Horii, Y., Takahashi, R., Yoshinami, Y. & Murata, A. Electrochemical reduction of CO at a copper electrode. J. Phys. Chem. B 101, 7075–7081 (1997).
37. Loiduice, A. et al. Tailoring copper nanocrystals towards C2 products in electrochemical CO2 reduction. Angew. Chem. Int. Ed. 55, 5789–5792 (2016).
38. Lum, Y., Cheng, T., Goddard, W. A. & Ager, J. W. Electrochemical CO reduction builds solvent water into oxygencate products. J. Am. Chem. Soc. 140, 9337–9340 (2018).
39. Pérez-Gallent, E., Figueiredo, M. C., Calle-Vallejo, F. & Koper, M. T. M. Spectroscopic observation of a hydrogenated CO dimer intermediate during CO reduction on Cu(100) electrodes. Angew. Chem. Int. Ed. 56, 3621–3624 (2017).
40. Goodpaster, J. D., Bell, A. T. & Head-Gordon, M. Identification of possible pathways for C–C bond formation during electrochemical reduction of CO2: new theoretical insights from an improved electrochemical model. J. Phys. Chem. Lett. 7, 1471–1477 (2016).
41. Xiao, H., Goddard, W. A., Cheng, T. & Liu, Y. Cu metal embedded in oxidized matrix catalyst to promote CO2 activation and CO dimerization for electrochemical reduction of CO2 to C2 hydrocarbons. Nat. Chem. 19, 974–980 (2018).
42. Zhu, W., Zhang, R., Qu, F., Asiri, A. M. & Sun, X. Design and application of foams for electrocatalysis. ChemSusChem 9, 1721–1732 (2016).
43. Ravel, B. & Nevill, M. ATHENA, ARTÉMIN, HEPHAESTUS: data analysis for X-ray absorption spectroscopy using IFEFFIT. J. Synchrotron Radiat. 12, 537–541 (2005).
44. Concorde, T. & Nika, software for two-dimensional data reduction. J. Appl. Crystallogr. 45, 324–328 (2012).

Acknowledgements
This work was supported by the Ontario Research Fund Research-Excellence Program, the Natural Sciences and Engineering Research Council (NSERC) of Canada, the CIFAR Brain-Inspired Solar Energy program, and University of Toronto Connaught grant. The STEM lab was supported by NSERC. We thank F. Studt and L. Sun for technical support, Z. Larabai for assistance with the STEM measurements. C.-W.H., C.-S.T., Z.C., M.C. and S.-C.L. conducted the aberration-corrected STEM imaging part of this research was completed at the Center for Nanophase Materials Sciences, which is a DOE Office of Science User Facility. Synchrotron measurements were carried out at APS, an Office of Science User Facility operated for U.S. DOE Office of Science by Argonne National Laboratory, and was supported by the U.S. DOE under Contract No. DE-AC02-06CH11357; the Canadian Light Source and its funding partners, and CMS beamline of the National Synchrotron Light Source II, a U.S. DOE Office of the Science User Facility operated for the DOE Office of Science by Brookhaven National Laboratory. All DFT computations were performed on the IBM BlueGene/Q supercomputer with support from the Southern Ontario Smart Computing Innovation Platform (SOSCI) and Niagara supercomputer at the SciNet HPC Consortium. SOSCI is funded by the Federal Economic Development Agency of Southern Ontario, the Province of Ontario, IBM Canada Ltd., Ontario Centres of Excellence, Mitacs, and 15 Ontario academic member institutions. SciNet is funded by the Canada Foundation for Innovation, the Government of Ontario, Ontario Research Fund - Research Excellence, and the University of Toronto. J.L. acknowledges the Banting Postdoctoral Fellowships Program and the receipt of support from the CLSI Graduate and Post-Doctoral Student Travel Support Program. The authors thank T.P.W., Y.Z.F., L.M., and G.S. for technical support at the 9BM beamline of APS. C. M.G. acknowledges support from NSERC in the form of a postdoctoral fellowship. A.R.K. and L.I.R. would like to thank R.L. and M.F. for their assistance in WAXS measurements.

Author contributions
E.H.S. supervised the project. X.W. designed and carried out the experiments, analyzed the data, and wrote the paper. Z.W. carried out DFT simulations and wrote the corresponding section. J.L., D.H.N. and F.I. performed synchrotron X-ray spectroscopy measurements. C.-W.H., C.-S.T., C.G. and S.-C.L. conducted the aberration-corrected TEM characterizations. C.M.G. performed SEM characterization. P.T. and A.P. carried out XPS measurements. A.R.K. and L.I.R. carried out the WAXS measurement and analysed the WAXS data. T.-T.Z., C.-T.D., A.S., Y.P.W., A.H.I., B.S., A.O., S.O.K. and D.S. assisted with the discussions. All authors discussed the results and contributed to the preparation of manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41467-019-13190-6.

Correspondence and requests for materials should be addressed to E.H.S.

Peer review information Nature Communications thanks William Goddard, Feng Jiao and other, anonymous, reviewers for their contributions to the peer review of this work.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2019