



Universiteit
Leiden
The Netherlands

A spongy nickel-organic CO₂ reduction photocatalyst for nearly 100% selective CO production

Niu, K.; Xu, Y.; Wang, H.; Ye, R.; Xin, H.L.; Lin, F.; ... ; Zheng, H.

Citation

Niu, K., Xu, Y., Wang, H., Ye, R., Xin, H. L., Lin, F., ... Zheng, H. (2017). A spongy nickel-organic CO₂ reduction photocatalyst for nearly 100% selective CO production. *Science Advances*, 3(7), e1700921. doi:10.1126/sciadv.1700921

Version: Not Applicable (or Unknown)
License: [Leiden University Non-exclusive license](#)
Downloaded from: <https://hdl.handle.net/1887/57813>

Note: To cite this publication please use the final published version (if applicable).

CHEMICAL PHYSICS

A spongy nickel-organic CO₂ reduction photocatalyst for nearly 100% selective CO production

Kaiyang Niu,^{1,2*} You Xu,^{3,4*} Haicheng Wang,^{2,5*} Rong Ye,^{2,6} Huolin L. Xin,⁷ Feng Lin,⁸ Chixia Tian,⁹ Yanwei Lum,^{1,2} Karen C. Bustillo,¹⁰ Marca M. Doeff,⁹ Marc T. M. Koper,¹¹ Joel Ager,^{1,2} Rong Xu,^{3,4†} Haimei Zheng^{1,2†}

Solar-driven photocatalytic conversion of CO₂ into fuels has attracted a lot of interest; however, developing active catalysts that can selectively convert CO₂ to fuels with desirable reaction products remains a grand challenge. For instance, complete suppression of the competing H₂ evolution during photocatalytic CO₂-to-CO conversion has not been achieved before. We design and synthesize a spongy nickel-organic heterogeneous photocatalyst via a photochemical route. The catalyst has a crystalline network architecture with a high concentration of defects. It is highly active in converting CO₂ to CO, with a production rate of $\sim 1.6 \times 10^4 \mu\text{mol hour}^{-1} \text{g}^{-1}$. No measurable H₂ is generated during the reaction, leading to nearly 100% selective CO production over H₂ evolution. When the spongy Ni-organic catalyst is enriched with Rh or Ag nanocrystals, the controlled photocatalytic CO₂ reduction reactions generate formic acid and acetic acid. Achieving such a spongy nickel-organic photocatalyst is a critical step toward practical production of high-value multicarbon fuels using solar energy.

INTRODUCTION

Rapid fossil fuel consumption induces environmental burden and energy crisis (1–3). Excessive anthropogenic CO₂ emission is a significant concern because of its hastening impact on climate change (4–6), acidification of ocean (7), crop yield reduction (8), extinction of animal species (9), and damage to human health (10, 11). Removal of excessive CO₂ from the atmosphere (12), particularly converting CO₂ to fuels using solar energy, is currently a global research endeavor (13–15). Discovering novel catalysts that can reduce the stable CO₂ molecules and convert them to liquid fuels with high activity and selectivity is essential (13, 14). To date, despite the progress that has been made in investigating the photocatalytic reduction of CO₂ (15–19), controlling the reaction to yield a specific product among many possible reaction species, including CO, H₂, CH₄, and formic acid, remains a great challenge (16, 20, 21). Finding photocatalysts that can efficiently convert CO₂ to CO and largely suppress other competing photocatalytic reactions, such as H₂ evolution, would be a critical step forward toward practical solar-to-fuels conversion for the production of high-value multicarbon fuels (15, 17, 22).

We recently developed a laser-chemical method and synthesized active transition metal hydroxide catalysts with a high concentration of defects for water oxidation (23). Specifically, we used an unfocused infrared laser to initiate the reactions between transition metal ions and triethylene glycol (TEG) and obtained a series of metal hydroxide-TEG

composites with a distorted layered structure (23). This disordered structure enhances the accessibility of water molecules to the active sites and enables efficient electrocatalysis of alkaline water oxidation (23). Such a laser-chemical strategy may be applied to the discovery of many other catalysts, for instance, novel nanostructured metal-organic heterogeneous catalysts for CO₂ reduction reaction.

When designing catalysts for CO₂ reduction, the material's ability to capture the CO₂ molecules is another significant consideration (24). Metal-organic frameworks (MOFs) with high surface area and tunable pores have been used for gas capture and heterogeneous catalysis (25, 26). Typically, MOFs have highly ordered crystalline structures constructed by coordinating metal ions or clusters with rigid organic linkers, most often the aromatic carboxylic acid molecules (27), such as terephthalic acid (TPA). In light of MOF structure design, we replace part of the rigid linkers (for example, TPA) in traditional MOFs with soft molecules (for example, TEG) by laser, considering the comparable molecular length of TEG to TPA (fig. S1). When the TEG molecules, which lack essential carboxylic groups for the perfect framework construction, are weaved into the metal-TPA framework, their substitution of TPA linkers may frustrate the growth of highly ordered MOF crystals, resulting in disordered and defective metal-organic hybrids for effective CO₂ fixation.

Here, we design a model metal-organic CO₂ reduction catalyst, with Ni²⁺ ions as active metal centers, TPA as a rigid linker, TEG as a soft linker, and dimethylformamide (DMF) as a solvent, via laser-induced solution reactions. The as-synthesized catalyst, labeled as Ni(TPA/TEG), has a crystalline network architecture with considerable defects and performs nearly 100% selective gas production (CO over H₂ evolution) with a high CO production rate of $\sim 1.6 \times 10^4 \mu\text{mol hour}^{-1} \text{g}^{-1}$. Further metal decorations (that is, Rh and Ag) of the Ni(TPA/TEG) catalyst lead to controlled photocatalytic CO₂ reduction reactions that generate formic acid and acetic acid.

RESULTS AND DISCUSSION

Structure determination of the Ni(TPA/TEG) catalyst

As shown in Fig. 1A, the Ni(TPA/TEG) composite forms a disordered spongy network structure, in which Ni, O, and C are uniformly distributed (fig. S2). In comparison, the solution without TEG, Ni(TPA) only,

¹Department of Materials Science and Engineering, University of California, Berkeley, Berkeley, CA 94720, USA. ²Materials Sciences Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, USA. ³School of Chemical and Biomedical Engineering, Nanyang Technological University, 62 Nanyang Drive, Singapore 637459, Singapore. ⁴SinBeRISE (Singapore-Berkeley Research Initiative for Sustainable Energy) CREATE, 1 Create Way, Singapore 138602, Singapore. ⁵National Center for Materials Service Safety, University of Science and Technology Beijing, Beijing 100083, P. R. China. ⁶Department of Chemistry, University of California, Berkeley, Berkeley, CA 94720, USA. ⁷Center for Functional Nanomaterials, Brookhaven National Laboratory, Upton, NY 11973, USA. ⁸Department of Chemistry, Virginia Tech, Blacksburg, VA 24061, USA. ⁹Energy Storage and Distributed Resources Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, USA. ¹⁰National Center for Electron Microscopy, Molecular Foundry, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, USA. ¹¹Leiden Institute of Chemistry, Leiden University, 2300 RA Leiden, Netherlands. *These authors contributed equally to this work.

†Corresponding author. Email: rxu@ntu.edu.sg (R.X.); hmzheng@lbl.gov (H.Z.)

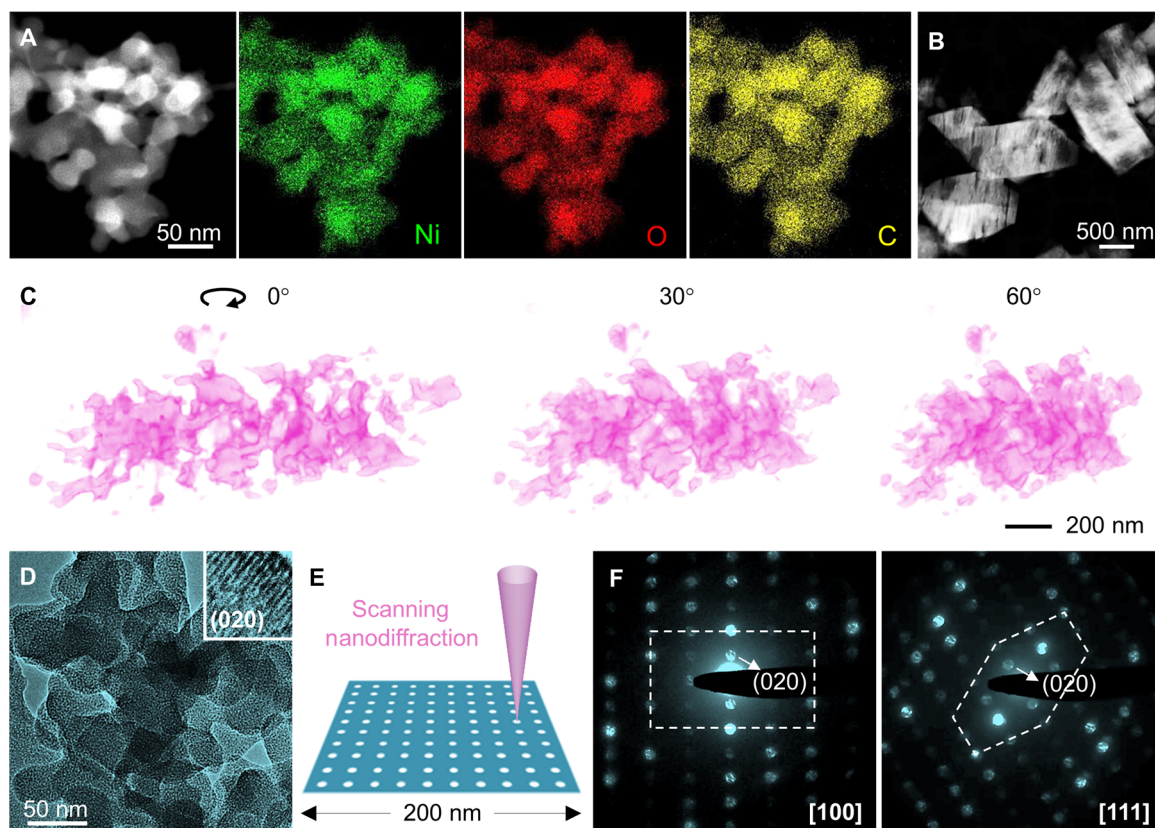


Fig. 1. Structure of the laser-chemical tailored spongy Ni(TPA/TEG) catalyst. (A) Scanning TEM (STEM) images and energy-dispersive x-ray spectroscopy (EDX) mapping of the spongy Ni(TPA/TEG) nanostructure. (B) STEM image of the Ni(TPA/TEG) particles. (C) Three-dimensional tomographic reconstruction of a fraction of spongy Ni(TPA/TEG) composite (movie S1). (D) TEM image of the spongy Ni(TPA/TEG) nanostructure. The inset high-resolution TEM image displays the defective (020) lattices [$d_{(020)} = 1.02$ nm] of an orthorhombic crystal. (E) Scanning electron nanodiffraction series taken from the Ni(TPA/TEG) particle by a scanning nanoprobe with an electron beam size of ~ 3 nm. The probe step size is 10 nm with an exposure time of 0.5 s at each step and a total beam current of ~ 5 pA. (F) Diffraction patterns showing the [100] and [111] orientations of the orthorhombic Ni(TPA/TEG) composite (movies S2 and S3). The dimensions of the diffraction patterns are $11.9 \text{ nm}^{-1} \times 11.9 \text{ nm}^{-1}$.

forms large particles (Fig. 1B). A three-dimensional electron tomographic reconstruction of the spongy Ni(TPA/TEG) architecture reveals various mesopores in the structure (Fig. 1C and movie S1), which closely resembles the pore features identified from the N_2 physisorption measurements (fig. S3). Figure 1D shows a typical transmission electron microscopy (TEM) image of the spongy Ni(TPA/TEG) composite, where defective lattices with a d -spacing of 1.02 nm are captured. To further interpret the structure of Ni(TPA/TEG) composite, we acquire a scanning nanobeam diffraction data set using an electron beam with a size of ~ 3 nm, a total beam current of ~ 5 pA, and an exposure time of 0.5 s, where the electron beam damage to the metal-organic material has been evidently minimized (Fig. 1E). Single-crystalline diffraction patterns along the [100] and [111] orientations of the Ni(TPA/TEG) composite are captured from two different regions of the spongy network (Fig. 1F and movies S2 and S3), showing an orthorhombic structure similar to that of the Ni(TPA) particles (fig. S4). Changes of the diffraction patterns are observed from movies S2 and S3, indicating defects (that is, grain boundaries) in the spongy Ni(TPA/TEG) catalyst (fig. S5).

To verify that the soft TEG molecules have been incorporated into the Ni(TPA) framework (Fig. 2A), we compare the structure of laser-synthesized Ni(TPA) and Ni(TPA/TEG) composites in detail. The x-ray diffraction (XRD) pattern (Fig. 2B) shows that the Ni(TPA) composite has an orthorhombic structure where the Ni-TPA units construct the

framework (fig. S4). The spongy Ni(TPA/TEG) has a crystal structure similar to that of Ni(TPA), but slight differences exist in the peak positions and widths of the x-ray lines, which may result from the exchange of linkers and solvent molecules in the structure (28). In the Fourier transform infrared (FTIR) spectra (Fig. 2C), we can see that both samples have clear bands of $\nu(\text{COO}^-)$ (1375 and 1575 cm^{-1}) and ring breathing (815 cm^{-1}) from the TPA linkers. Characteristic bands of δNCO (690 cm^{-1}) and $\nu(\text{CO})$ (1685 cm^{-1}) from DMF molecules are found in Ni(TPA), indicating that DMF molecules may reside in the MOF cavities and coordinate with Ni^{2+} through carbonyl groups (28). Meanwhile, distinct bands of $\nu(\text{OH})$ (1065 and 3374 cm^{-1}) related to TEG are found exclusively in Ni(TPA/TEG), and no DMF bands are detected, indicating that TEG molecules exist in Ni(TPA/TEG); DMF molecules that originally occupied the Ni(TPA) framework cavities prefer to leave. The EDX spectrum (Fig. 2D) also shows that no nitrogen (from DMF) can be detected in Ni(TPA/TEG). Thermogravimetric analysis (TGA) curves in Fig. 2E, both displaying three stages of mass losses, indicate differences between the two samples. For Ni(TPA), the mass loss measurements of 6, 18, and 50% are from H_2O , DMF, and TPA, respectively (28); for Ni(TPA/TEG), the mass losses of 16, 26, and 26% are from H_2O , TEG, and TPA, respectively. The large differences in the mass loss of TPA in the two samples (50% versus 26%) indicate that soft TEG molecules have replaced part of the rigid TPA linkers, causing the formation of a spongy Ni(TPA/TEG)

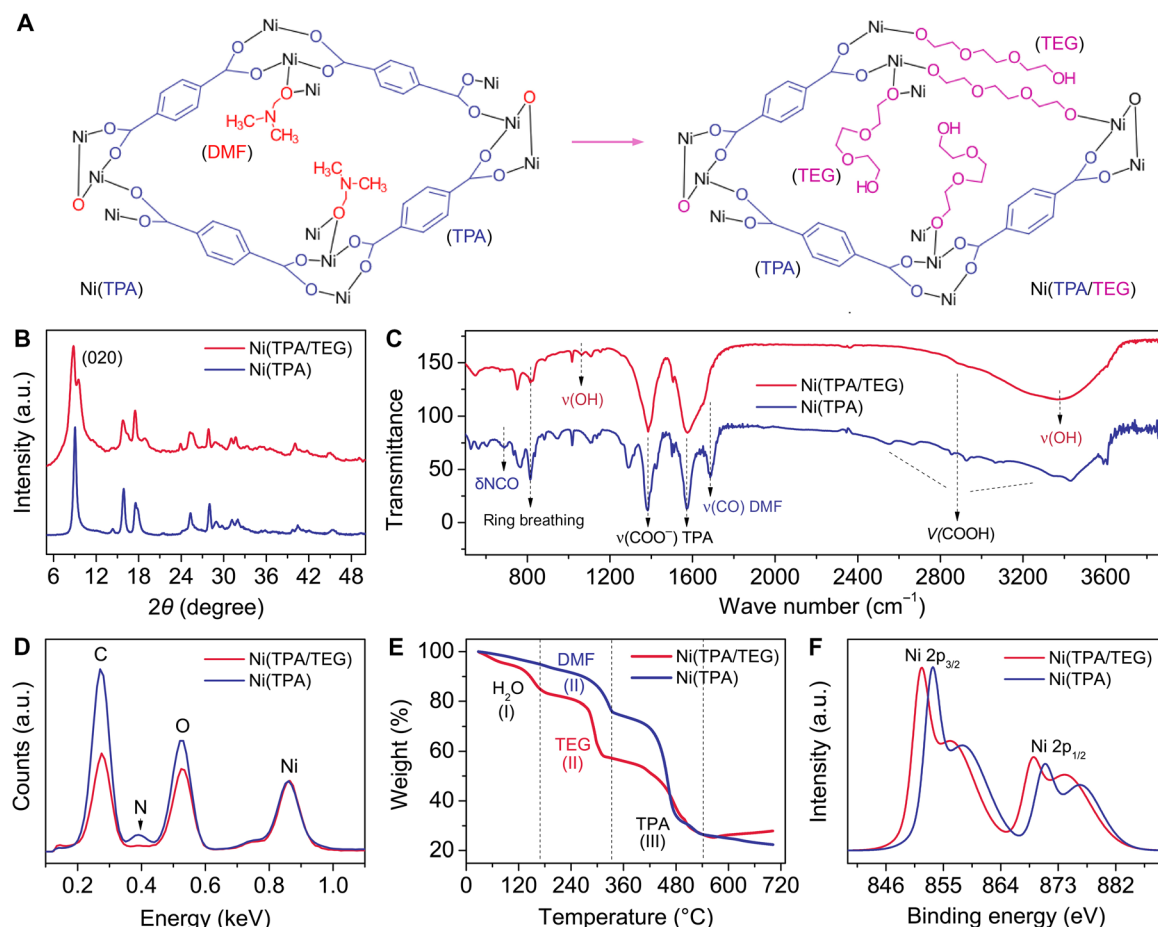


Fig. 2. Comparison of laser-chemical tailored Ni(TPA/TEG) and Ni(TPA) composites. (A) Proposed design strategy of the disordered spongy Ni(TPA/TEG) composite by introducing soft Ni-TEG building units into a Ni(TPA) framework through laser-chemical reaction. XRD patterns (B), FTIR spectra (C), EDX spectra (D), TGA curves (E), and XPS spectra (F) of the laser-chemical tailored Ni(TPA/TEG) and Ni(TPA) composites. a.u., arbitrary units.

network. Because of the varying chemical environment of Ni^{2+} , the $\text{Ni}2p$ peaks in the x-ray photoelectron spectroscopy (XPS) spectra (Fig. 2F) shifted to the right in Ni(TPA/TEG) compared to Ni(TPA).

We also find that it is more effective to use lasers to cross-link the soft TEG and rigid TPA molecules together with the Ni^{2+} centers than to use a traditional heating process (see the Supplementary Materials). Laser irradiation appears to produce Ni-TEG building units that are indispensable for the spongy Ni(TPA/TEG) network construction (Fig. 2A). Because of the soft character of TEG molecules, various inhomogeneous configurations of Ni-TEG units can be generated. Mismatches between the soft Ni-TEG units and the rigid Ni-TPA units may introduce considerable defects, leading to the formation of disordered spongy Ni(TPA/TEG).

Evaluation of the photocatalytic activity for CO production

We apply the as-synthesized Ni-organic composites for visible light-driven photocatalytic CO_2 reduction in a solvent mixture of acetonitrile/water [considering the high solubility of CO_2 in acetonitrile (29)] under mild reaction conditions (20°C and 400 torr of CO_2), with triethanolamine (TEOA) as a sacrificial reducing agent and $\text{Ru}(\text{bpy})_3\text{Cl}_2 \cdot \text{H}_2\text{O}$ as a photosensitizer (18, 30). Five samples (figs. S6 to S8), that is, Ni(TPA) (L), Ni(TPA) (H), Ni(TPA/TEG) (L), Ni(TPA/TEG) (H), and Ni(TEG) (L), synthesized by both laser irra-

diation (L) and traditional heating (H), are examined. Figure 3A shows the CO evolutions from these five Ni-organic catalysts in a 6-hour photocatalytic reaction. The spongy Ni(TPA/TEG) (L) composite shows the highest activity, and the amount of CO is $95.2 \mu\text{mol}$ after a 2-hour reaction, giving a CO production rate of $15,866 \mu\text{mol hour}^{-1} \text{g}^{-1}$, which is several times higher than that from other samples. The total amount of CO produced on the spongy Ni(TPA/TEG) catalyst in 6 hours reaches $136.9 \mu\text{mol}$ (Fig. 3A), giving a turnover number of 11.5 for the 6-hour reaction (table S1). The CO production rate is also superior compared with many other reported heterogeneous CO evolution photocatalysts to the best of our knowledge (17), such as the Co_3O_4 platelets with $[\text{Ru}(\text{bpy})_3]\text{Cl}_2$ as a photosensitizer ($3523 \mu\text{mol hour}^{-1} \text{g}^{-1}$) (18), the sensitized TiO_2 particles with enzyme as a cocatalyst ($300 \mu\text{mol hour}^{-1} \text{g}^{-1}$) (31), and the sensitized $\text{BaLa}_4\text{Ti}_4\text{O}_{15}$ particles with Ag as a cocatalyst ($22 \mu\text{mol hour}^{-1} \text{g}^{-1}$) (16). Note that the soluble homogeneous metal complex catalysts, which have also been investigated for controlled CO_2 reduction (32–34), are not categorized here for comparison.

By testing the 2-hour yield of CO on different amounts of the Ni(TPA/TEG) catalyst, we obtain a roughly linear relationship between the amount of evolved CO and the amount of the catalyst (Fig. 3B). However, kinetically, we found that the CO production rate actually decreases with the increase in the amount of the catalyst

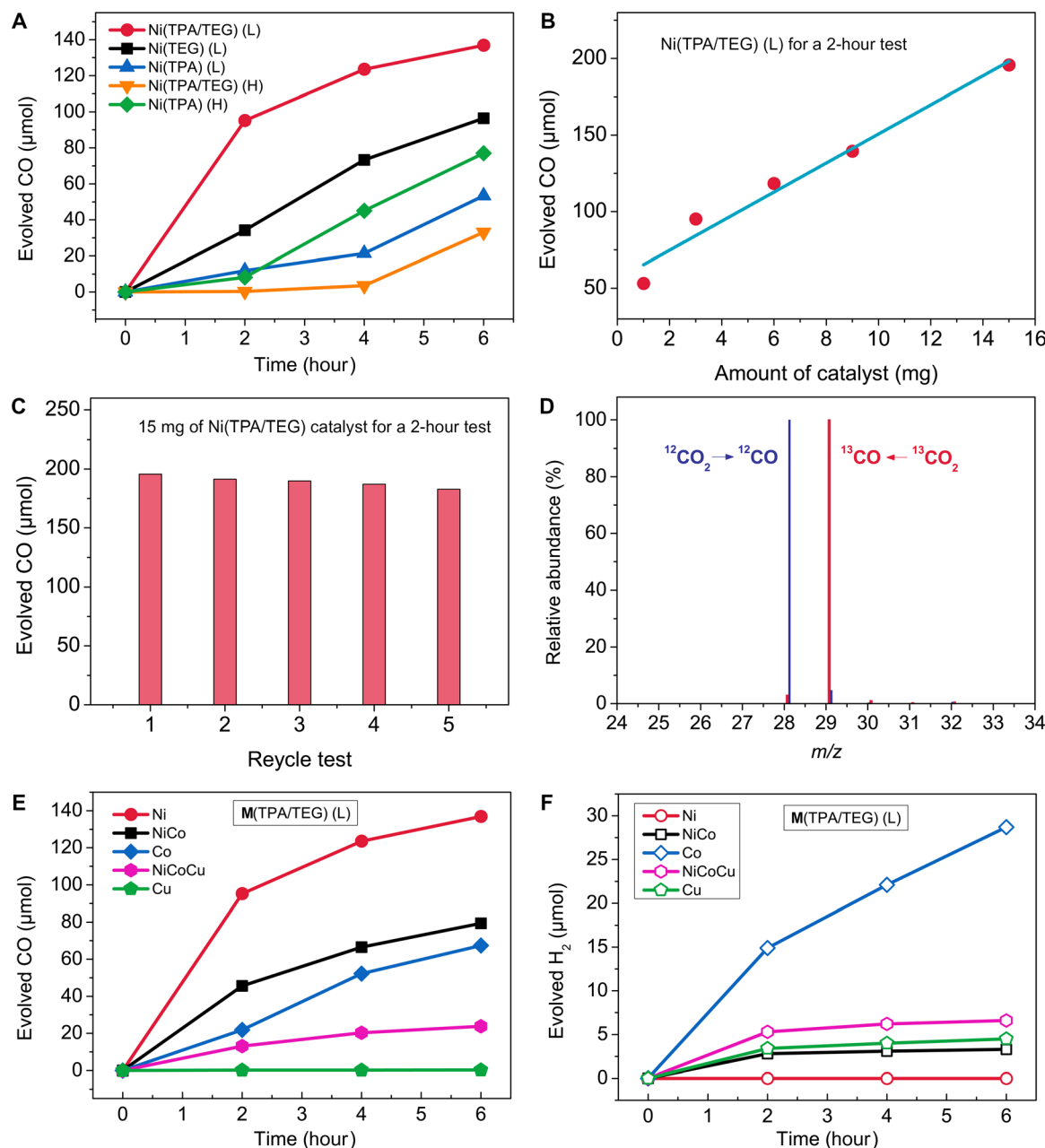


Fig. 3. Conversion of CO_2 to CO by photocatalysis. (A) CO evolution on five Ni-based catalysts with different combinations of TPA, TEG, and DMF. The composites synthesized by laser-chemical approach are labeled with "L"; the ones synthesized by traditional heating method are marked with "H." (B) CO production on different amounts of the Ni(TPA/TEG) catalyst. (C) Average yield of CO in the first 2 hours for five recycling tests. (D) MS of ^{12}CO (blue lines) and ^{13}CO (red lines) produced on the spongy Ni(TPA/TEG) catalyst by using $^{12}\text{CO}_2$ and $^{13}\text{CO}_2$ as gas sources, respectively. m/z , mass/charge ratio. (E) Comparison of CO evolution on five laser-synthesized $M(\text{TPA/TEG})$ ($M = \text{Ni}, \text{Co}, \text{Cu}$) catalysts. (F) Comparison of H_2 evolution on the five $M(\text{TPA/TEG})$ catalysts.

(fig. S9), where 1.0 mg of the Ni(TPA/TEG) catalyst gives a CO production rate of $\sim 26,620 \mu\text{mol hour}^{-1} \text{g}^{-1}$ in the same solution, indicating that more electrons generated from the photosensitizer molecules could have been transferred to the catalytic active sites. We have also tested the reusability of the spongy Ni(TPA/TEG) catalyst upon each 2-hour photocatalysis, where the catalyst has kept its activity and selectivity after recycling (Fig. 3C). It also exhibits excellent structural stability, and no obvious structural change is found after 24 hours of photocatalysis (fig. S10). To confirm the origin of the as-produced CO, we use isotopic $^{13}\text{CO}_2$ as feedstock gas for the photocatalytic re-

duction and examine the products by gas chromatography-mass spectrometry (GC-MS). A major signal at a mass/charge ratio of 29 on the mass spectrum corresponding to ^{13}CO (Fig. 3D, red lines) appears, which confirms that the as-detected CO originates from the CO_2 gas source (fig. S11).

Transition metal ions with switchable electronic states have long been considered promising active sites for diverse photocatalytic or electrocatalytic reactions, such as water splitting (35–37), CO oxidation (38), and CO_2 reduction (39). Consequently, we use the laser-chemical method to synthesize four additional samples and compare different

metal ions, that is, Ni^{2+} , Co^{2+} , and Cu^{2+} (fig. S12), as active sites for the photocatalytic CO_2 reduction. Results show that the spongy $\text{Ni}(\text{TPA}/\text{TEG})$ catalyst is still the most active catalyst for CO evolution, and the activity worsens with the incorporation of Co and Cu ions (Fig. 3E). The amount of CO generated on the $\text{Ni}(\text{TPA}/\text{TEG})$ catalyst is almost two times that of the CO generated from $\text{Co}(\text{TPA}/\text{TEG})$ in a 6-hour photocatalytic reaction. The $\text{Cu}(\text{TPA}/\text{TEG})$ catalyst barely generates any CO, which is distinctly different from the metal Cu catalyst that has superior activity for CO evolution from the electrocatalytic reduction of CO_2 (40).

Evaluation of the CO production selectivity

We find that CO ($\text{CO}_2 + \text{H}_2\text{O} + 2\text{e}^- \rightarrow \text{CO} + 2\text{OH}^-$) is the only detectable gas product from the photocatalytic CO_2 reduction on the TPA-containing Ni-organic catalysts. H_2 evolution ($2\text{H}_2\text{O} + 2\text{e}^- \rightarrow \text{H}_2 + 2\text{OH}^-$), usually acting as a major competing reaction in the CO_2 reduction system for many transition metal-based catalysts (18, 20), has been completely suppressed (table S1). Thus, a near 100% selectivity of CO production (over H_2 evolution) is achieved. Note that no other potential competing gas products, such as CH_4 and C_2H_4 (17), have been detected in our experiments either.

For comparison, we have detected considerable amounts of H_2 from other laser-synthesized $M(\text{TPA}/\text{TEG})$ ($M = \text{Ni}, \text{Co}, \text{Cu}$) catalysts (Fig. 3, E and F, and fig. S13), and CO selectivity values of 96, 70, 78.2, and 4.8% were measured for $\text{NiCo}(\text{TPA}/\text{TEG})$, $\text{Co}(\text{TPA}/\text{TEG})$, $\text{NiCoCu}(\text{TPA}/\text{TEG})$, and $\text{Cu}(\text{TPA}/\text{TEG})$, respectively. We have also detected a fair

amount of H_2 (25.7 μmol) from the hydroxylated TPA-free $\text{Ni}(\text{TEG})$ catalyst in addition to the CO evolution (96.5 μmol) after a 6-hour reaction (fig. S14), which is analogous to the Ni-based hydroxides for practical H_2 evolution from the electrocatalysis of water (36).

Tuning the selectivity for liquid fuels production from CO_2

Furthermore, considering the potential of noble metal electrodes for CO_2 reduction (41, 42), we decorate the spongy $\text{Ni}(\text{TPA}/\text{TEG})$ with noble metal nanocrystals, that is, Rh and Ag, in pursuit of tuning the selectivity of liquid fuels production from the photocatalytic CO_2 reduction (15, 21). Figure 4 (A to C) shows the Ag-decorated $\text{Ni}(\text{TPA}/\text{TEG})$ catalyst, where Ag nanocrystals with an average diameter of ~ 6 nm are well dispersed on the $\text{Ni}(\text{TPA}/\text{TEG})$. We examine the liquid products after a 6-hour reaction on three catalysts, that is, undecorated $\text{Ni}(\text{TPA}/\text{TEG})$, Rh-decorated $\text{Ni}(\text{TPA}/\text{TEG})$, and Ag-decorated $\text{Ni}(\text{TPA}/\text{TEG})$ (Fig. 4D). For the $\text{Ni}(\text{TPA}/\text{TEG})$ catalyst without any decoration, we measure formic acid (HCOOH) with a concentration of 29.2 μM and acetic acid (CH_3COOH) with a concentration of 72.5 μM in addition to CO. With the decoration of Rh and Ag nanocrystals, the amounts of CO decrease drastically, whereas the amounts of liquid products significantly increase. Formic acid (HCOOH) with a concentration of 313.5 μM is mainly obtained on the Rh-decorated $\text{Ni}(\text{TPA}/\text{TEG})$ catalyst, and CH_3COOH with a concentration of 195.6 μM is the major product for the Ag-decorated $\text{Ni}(\text{TPA}/\text{TEG})$ catalyst, where the origin of CH_3COOH has been confirmed from the flowing CO_2 source by MS (fig. S15). Note that CO,

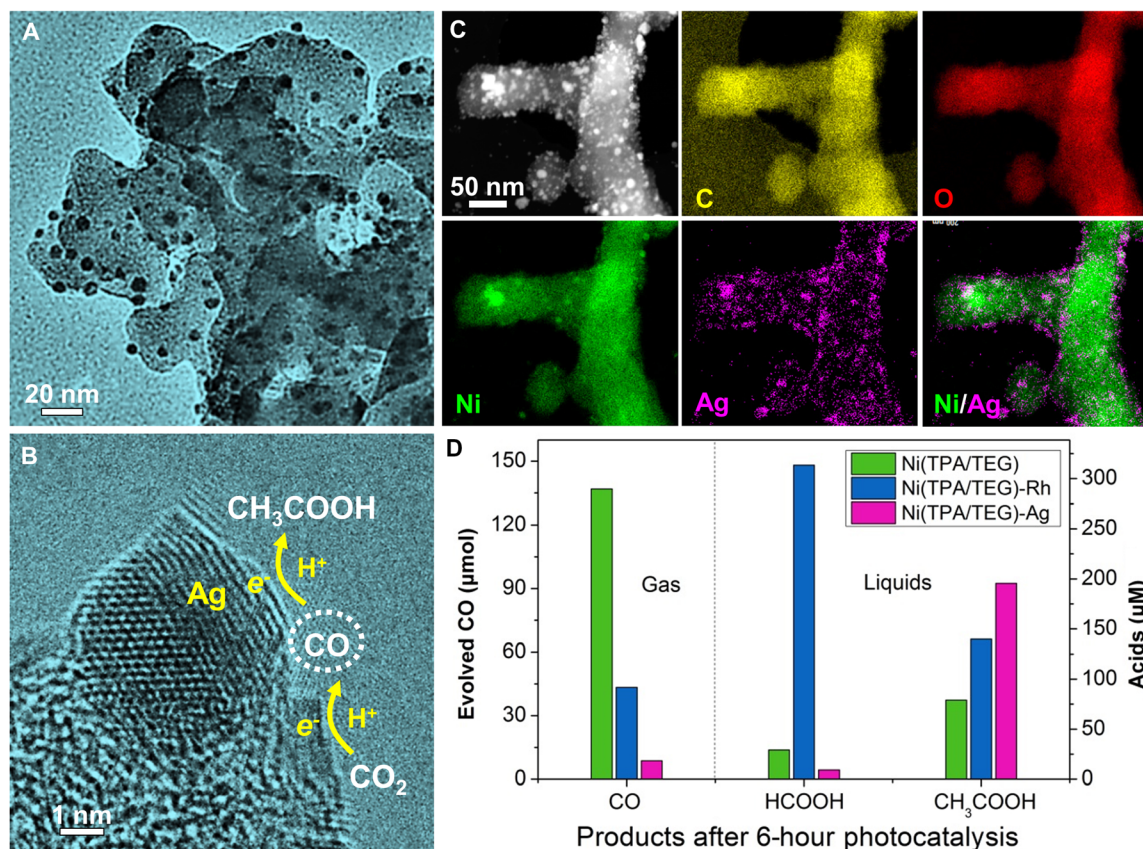


Fig. 4. Generation of liquid products on metal-decorated $\text{Ni}(\text{TPA}/\text{TEG})$ composites. Low-magnification (A) and high-resolution (B) TEM images of the $\text{Ni}(\text{TPA}/\text{TEG})$ composite decorated with Ag nanocrystals. (C) EDX mapping of the as-prepared $\text{Ni}(\text{TPA}/\text{TEG})$ -Ag composite. (D) Comparison of the amount of the products (CO, HCOOH, and CH_3COOH) generated from photocatalytic CO_2 reduction on $\text{Ni}(\text{TPA}/\text{TEG})$, $\text{Ni}(\text{TPA}/\text{TEG})$ -Rh, and $\text{Ni}(\text{TPA}/\text{TEG})$ -Ag catalysts.

formic acid, and acetic acid reflect the overall product distribution of the photocatalytic CO₂ reduction reaction; no other liquid products, such as methanol, ethanol, or propanol, are detected in this experiment.

Enhanced production of liquid fuels from photocatalytic reduction of CO

By assuming that the evolved CO may be further consumed for the production of acids, we also conduct control experiments using CO, instead of CO₂, as the gas feedstock for the photocatalytic reduction reaction. As a result, largely enhanced yields of acids are obtained. For instance, the amount of HCOOH evolved from the CO reduction is 24 times higher than that from the CO₂ reduction on the Ni(NPA/TEG) catalyst (Fig. 5A), and the amount of CH₃COOH produced from the CO reduction reaction for the Ag/Rh-decorated Ni(NPA/TEG) catalyst exhibits a sixfold increase over the results from the CO₂ reduction reaction (Fig. 5B and fig. S16). Besides the increase in acid production, another important C2 product, that is, ethanol [with a concentration of 270.6 μM for Ni(NPA/TEG)-Rh and 262.2 μM for Ni(NPA/TEG)-Ag], has also emerged from the 6-hour photocatalytic CO reduction reaction.

Possible mechanisms for the photocatalytic reduction reactions of CO₂ or CO

On the basis of the abovementioned results, we propose the following mechanism for the photocatalytic CO₂ reduction reactions on the

spongy Ni(TPA/TEG) catalyst (Fig. 6). Upon visible light irradiation, the photosensitizer [Ru(bpy)₃]²⁺ is excited and then reductively quenched by the TEOA sacrificial electron donor (18, 30, 43), which gives rise to the reduced species of [Ru(bpy)₃]²⁺ (Fig. 6A). Subsequently, the reduced species of [Ru(bpy)₃]²⁺ could transfer an electron to the spongy Ni(TPA/TEG) catalyst, which then participates in reducing the CO₂ molecules fixed on the catalyst (Fig. 6B). In the 2-hour yield tests of CO production in the solution with different amounts of Ni(TPA/TEG), we have found that the CO production rate decreases with the increase in the amount of catalyst (fig. S9), indicating that the electron transfer from [Ru(bpy)₃]²⁺ to the catalyst could be a rate-determining step for the CO₂ reduction reaction, where a diffusion-limited event may have occurred in this heterogeneous catalytic system (44).

For the selective generation of CO from CO₂, we propose the formation of a basic intermediate (45) [that is, CO₂ radical anion (CO₂^{•-})] in the initial reaction step (CO₂ + e⁻ → CO₂^{•-}), which acts as a Brønsted base and reacts easily with H₂O to form CO (CO₂^{•-} + H₂O + e⁻ → CO + 2OH⁻). On the other hand, in our proton-deficient photocatalytic reaction medium (pH 8), water (the proton donor) could provide another nonbasic intermediate, H[•] (H₂O + e⁻ → H[•] + OH⁻), before the H₂ evolution (H₂O + H[•] + e⁻ → H₂ + OH⁻) (46). For the spongy Ni(TPA/TEG) catalyst, it is likely that the Ni-TPA coordination units are unfavorable for the binding of H[•] on the active sites, limiting the proton transfer and the formation of H₂. In addition, the flexible Ni-TEG units, with superior structural resistance to the aqueous environment (23), enable the disordered spongy network construction, where the open and defective structure provides more accessible Ni²⁺ active sites to capture and stabilize the CO₂^{•-} intermediates, leading to the efficient production of CO.

Subsequently, the evolved CO can be further reduced to liquid fuels through proton-coupled multielectron reaction processes (Fig. 6C and fig. S17). Figure 6D shows the proposed conversion pathways leading to the formation of HCOOH, CH₃COOH, and CH₃CH₂OH via proton-coupled one-, four-, and eight-electron steps, respectively, in the electrolyte with a pH value of ~8. For the formation of HCOOH, we propose the following reactions



where ^{*}CO is quickly protonated to ^{*}CHO and then hydroxylated to HCOOH. In the pathway of CH₃COOH formation, ^{*}CO is continuously hydrated to ^{*}CHO → ^{*}CHOH → ^{*}CH₂OH → ^{*}CH₃OH, which bonds with the adsorbed ^{*}CO to form CH₃COOH. As to the formation of CH₃CH₂OH, the dehydroxylation of the as-formed ^{*}CHO could be a critical rate-limiting step to produce ^{*}C that can be further protonated to ^{*}CH → ^{*}CH₂ → ^{*}CH₃ (47), and the C-C coupling between ^{*}CH₃ and multiprotonated ^{*}CO (that is, CH₂OH) could lead to the formation of CH₃CH₂OH (48).

To further tune the proposed reactions, we performed the CO reduction reaction on the Ni(TPA/TEG)-Ag in the electrolyte with a pH value of ~13 (fig. S18). We found that CH₃OH (184.09 μM), CH₃CH₂OH (149.39 μM), HCOOH (438.98 μM), and CH₃COOH (276.99 μM) are produced at pH 13 in 6 hours, which is distinct from the liquid products generated at pH 8, that is, CH₃OH (0 μM), CH₃CH₂OH

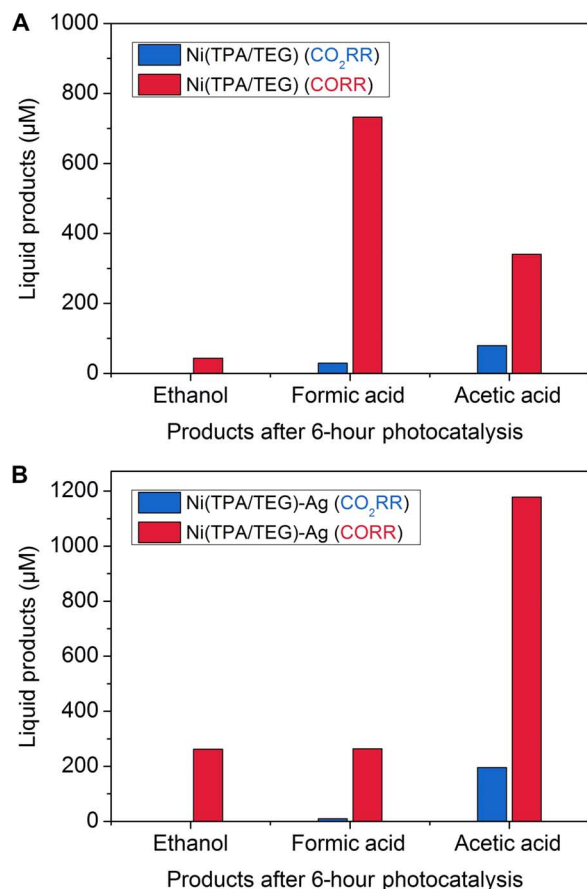


Fig. 5. Comparison of the liquid products generated from photocatalytic CO₂ reduction reactions (CO₂RR) and CO reduction reactions (CORR) on two catalysts. (A) Ni(TPA/TEG). (B) Ni(TPA/TEG)-Ag.

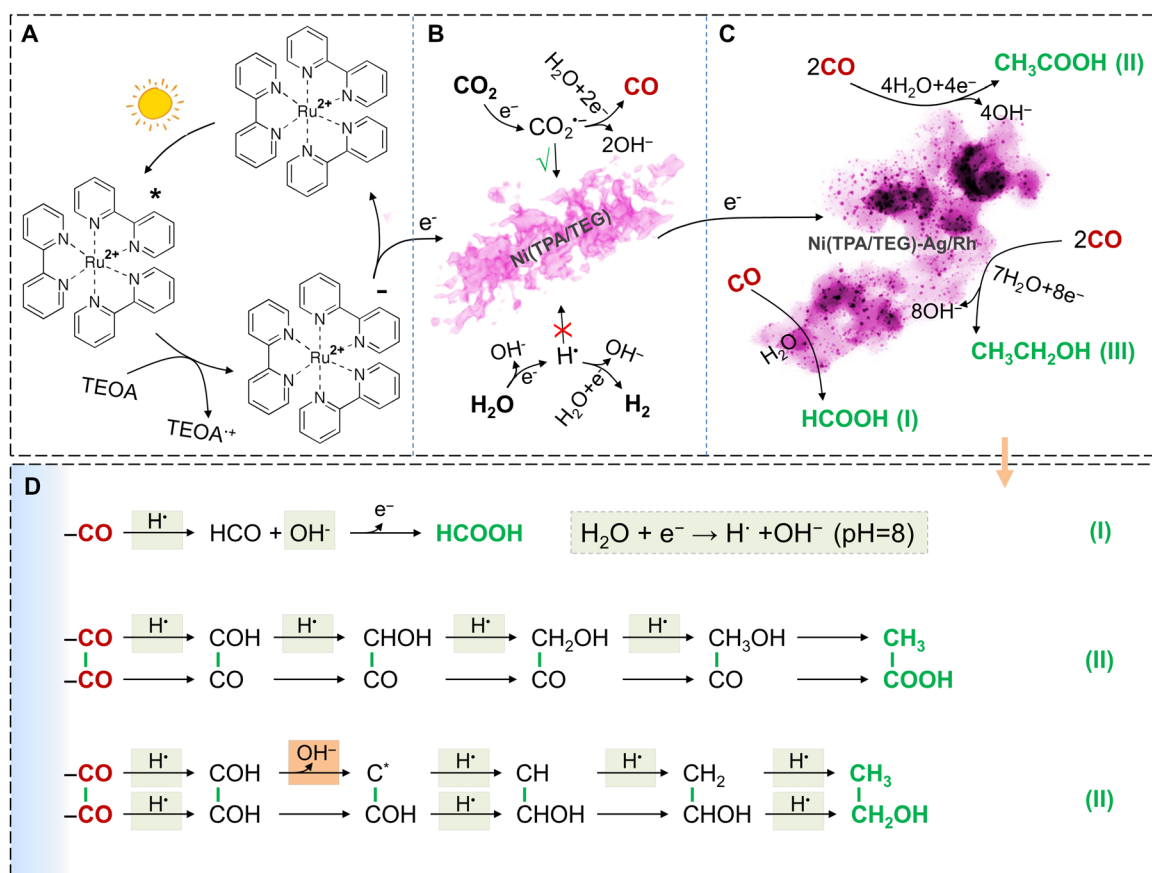


Fig. 6. Proposed mechanisms for the photocatalytic conversion of CO_2 to CO and of CO to other liquid products. (A) Visible light reduction of the photosensitizer $[\text{Ru}(\text{bpy})_3]^{2+}$, which transfers an electron to the $\text{Ni}(\text{TPA/TEG})$ catalyst to convert CO_2 to CO (B) and to $\text{Ni}(\text{TPA/TEG})\text{-Ag/Rh}$ catalysts for the generation of HCOOH , CH_3COOH , and $\text{CH}_3\text{CH}_2\text{OH}$ from further reduction of CO (C). The STEM image in (C) is the Ag-decorated $\text{Ni}(\text{TPA/TEG})$ catalyst (the original STEM image is shown in fig. S15). (D) Possible conversion pathways leading to the formation of HCOOH , CH_3COOH , and $\text{CH}_3\text{CH}_2\text{OH}$ via proton-coupled one-, four-, and eight-electron steps, respectively.

(262.19 μM), HCOOH (263.58 μM), and CH_3COOH (1178.04 μM). At pH 13, the hydroxyl ions (OH^-) are commonly available for the hydroxylation of $^*\text{CHO}$, which favors the formation of HCOOH (fig. S18). On the contrary, the enhanced hydroxylation of $^*\text{CHO}$ may suppress the kinetics of the multiprotonation of $^*\text{CO}$ and the dehydroxylation of $^*\text{CHO}$ (Fig. 6D), resulting in a lower amount of CH_3COOH and $\text{CH}_3\text{CH}_2\text{OH}$ at pH 13. The appearance of CH_3OH may suggest a weak C–C coupling between $^*\text{CH}_3\text{OH}$ and $^*\text{CO}$ at pH 13 (leading to CH_3COOH at pH 8), which should be considered for the future CO_2/CO reduction catalyst design (49).

In summary, we have demonstrated a photochemical strategy for the design of novel nanostructured metal-organic materials, where the rigid TPA and soft TEG molecules are successfully cross-linked together with the Ni^{2+} centers. A spongy $\text{Ni}(\text{TPA/TEG})$ hybrid structure with a considerably high concentration of defects has been achieved. We found that the $\text{Ni}(\text{TPA/TEG})$ catalyst is remarkably active for CO production (with a production rate of $15,866 \mu\text{mol hour}^{-1} \text{g}^{-1}$) from the heterogeneous photocatalytic CO_2 reduction reaction, during which no other measurable competing gases such as H_2 or CH_4 are generated, thus giving a near 100% CO selectivity over other gases. When the spongy Ni-organic catalyst is enriched with Rh or Ag nanocrystals, formic acid and acetic acid can be produced selectively from the photocatalytic CO_2 reduction reactions. We propose to use the spongy $\text{Ni}(\text{TPA/TEG})$ catalyst in a “tandem catalyst” system to convert CO_2 into high-value liq-

uid fuels using visible light, where the selectively CO_2 -turned CO can be directly used as an intermediate gas reactant for the generation of liquid fuels, such as ethanol and acetic acid (fig. S19). More advanced metal-organic heterogeneous photocatalysts with improved CO_2 fixation and light-harvesting capabilities are expected to be fabricated using the photochemical strategy for efficient solar-to-fuels conversion.

MATERIALS AND METHODS

Laser-chemical synthesis of metal-organic photocatalysts

All chemicals including nickel nitrate (99.99%), cobalt nitrate (99%), copper nitrate trihydrate (99%), ethanol (99.5%), DMF, TPA, and TEG were purchased from Sigma-Aldrich and used as received. The deionized water was produced by the Milli-Q Integral water purification system. A Continuum Surelite III nanosecond pulsed laser was used as a power source, and the following were the typical parameters of operation: wavelength, 1064 nm; frequency, 10 Hz; pulse width, 7 to 8 ns; beam diameter, 0.9 cm; and 700 mJ per pulse. TEG solutions (1 ml) of 1.5 M transition metal nitrates were added into 5 ml of DMF solution of 0.5 M TPA; the mixed solutions were stirred for 30 min before laser irradiation. For the syntheses of NiCo- and NiCoCu-organic frameworks, molar ratios of Ni/Co (1:1) and Ni/Co/Cu (1:1:1) were used, respectively. Typically, 3-hour laser irradiation was required for a 6-ml mixed precursor solution to complete the reaction. For the

syntheses by heating method, the same precursor solutions in glass vials were heated at 110°C in an oven for 48 hours. Precipitates produced after laser irradiations or heating were rinsed with acetone/ethanol, centrifuged at 9000 rpm for three times, and then dried in air at 60°C to obtain powders.

Photocatalysis measurement of CO₂ reduction reactions

The visible light-driven photocatalytic CO₂ reduction was conducted in a closed gas circulation and evacuation system fitted with a top window Pyrex cell. A circulating cooling water system was used to maintain the reactor at around 20°C. In a typical reaction, 3 mg of catalyst (for each test), 2.5 mmol of Ru(bpy)₃Cl₂·6H₂O, and 2 ml of TEOA were added to 10 ml of acetonitrile/H₂O solvent mixture (CH₃CN/H₂O = 8:2). The pH value of the solution was ~8, which was tuned to 13 by 1 M NaOH aqueous solution for control experiment. The light source was a 300-W Xe lamp with a long-pass cutoff filter ($\lambda > 420$ nm). Before light irradiation, the reaction system was evacuated and refilled with high-purity CO₂ (99.995%; SOXAL) several times to remove air inside and finally filled with CO₂ gas to reach a pressure of 400 torr. The evolved gas was detected by online GC (Agilent 7890A) equipped with a thermal conductivity detector and a flame ionization detector (FID) at different times of the photoreaction. To evaluate catalyst reusability, 15 mg of the catalyst was applied for photocatalysis and recycled by centrifugation after a 2-hour reaction, and then mixed with 8 ml of CH₃CN, 2 ml of H₂O, 2 ml of TEOA, and 18 mg of Ru(bpy)₃Cl₂·6H₂O for the next run. The solution products in the liquid phase were analyzed separately at the end of the photoreaction. Alcohols in the liquid phase were analyzed by Agilent 7890A GC with an FID, a DB-WAX column, and helium as the carrier gas. Carboxylic acids in the liquid phase were analyzed using an Agilent 1260 high-performance liquid chromatograph (HPLC) with a PL Hi-Plex H column and variable wavelength detector (VWD) (210 nm). [Ru(bpy)₃]Cl₂·6H₂O (99.95%) and TEOA ($\geq 99.0\%$) were obtained from Sigma-Aldrich. The acetonitrile (HPLC-grade) was obtained from Merck.

The ¹³CO₂ isotope tracer experiment was performed under similar photocatalytic reaction conditions. The reactor containing 3 mg of the catalyst, 2.5 mmol of Ru(bpy)₃Cl₂·6H₂O, 2 ml of TEOA, and 10 ml of the CH₃CN-H₂O mixture solution (CH₃CN/H₂O = 8:2) was first evacuated to ensure air removal and then refilled with ¹³CO₂ gas (99 atomic % ¹³C; Aldrich) to reach a pressure of 400 torr. After 4 hours of irradiation, 200 μ l of gas products was withdrawn using a gas-tight syringe and then injected into a GC-MS (Agilent, GC Model 6890N/MS Model 5973) with a molecular sieve 5 Å column for analysis. The carboxylic acid analysis was carried out separately by using a Thermo Finnigan LCQ MS system.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at <http://advances.sciencemag.org/cgi/content/full/3/7/e1700921/DC1>

Materials characterizations

Three-dimensional tomographic reconstruction of the spongy Ni(TPA/TEG) network

Comparison of the samples synthesized by laser-chemical route and traditional heating method

Control experiments to confirm the origin of the evolved CO

fig. S1. Structure of the selected rigid TPA linkers and soft TEG linkers for the laser synthesis of the nickel-organic photocatalyst.

fig. S2. Morphology and composition of the spongy Ni(TPA/TEG) composite.

fig. S3. Nitrogen physisorption measurements of the spongy Ni(TPA/TEG) catalyst.

fig. S4. A comparison between the measured and simulated powder XRD patterns of the crystalline Ni(TPA) framework.

fig. S5. Electron diffraction of the Ni(TPA/TEG) catalyst.

fig. S6. STEM images show the morphologies of five nickel-organic composites synthesized either by the laser-chemical approach (L) or the conventional heating method (H).

fig. S7. Comparison of the structure of Ni(TPA/TEG) and Ni(TPA) composites synthesized by both laser-chemical (L) and heating (H) methods.

fig. S8. Brunauer-Emmett-Teller surface area of five as-synthesized nickel-organic composites, including Ni(TPA/TEG) (L), Ni(TEG) (L), Ni(TPA)-DMF (L), Ni(TPA)-TEG (H), and Ni(TPA)-DMF (H).

fig. S9. CO production rate on different amounts of the Ni(TPA/TEG) catalyst.

fig. S10. Powder XRD patterns of the spongy Ni(TPA/TEG) catalyst before and after 24-hour photocatalytic reactions.

fig. S11. Controlled photocatalysis experiments by applying the Ni(TPA/TEG) (L) catalyst in either CO₂ or He gas, and without adding catalyst in CO₂.

fig. S12. Comparison of the morphology and structure of the five laser-synthesized *M*(TPA/TEG) (*M* = Ni, Co, Cu) catalysts, including Ni(TPA/TEG), NiCo(TPA/TEG), Co(TPA/TEG), NiCoCu(TPA/TEG), and Cu(TPA/TEG).

fig. S13. Comparison of the yield of CO and H₂ on five laser-synthesized *M*(TPA/TEG) (*M* = Ni, Co, Cu) catalysts after a 6-hour photocatalytic CO₂ reduction reaction.

fig. S14. Comparison of the gas products evolved from the laser-synthesized Ni(TPA/TEG) and Ni(TEG) catalysts.

fig. S15. MS of acetic acid (CH₃COOH) produced from the photocatalytic reduction of CO₂ using the Ni(TPA/TEG)-Ag catalyst.

fig. S16. Comparison of the liquid products generated from photocatalytic CO₂RR and CORR on the Ni(TPA/TEG)-Rh catalyst.

fig. S17. STEM image of the Ag-decorated Ni(TPA/TEG) catalyst.

fig. S18. Comparison of the liquid products generated from photocatalytic CO reduction reactions on the Ni(TPA/TEG)-Ag catalyst at pH 8 and 13.

fig. S19. Proposed three-cell tandem photocatalytic CO₂ reduction system for liquid fuels production.

table S1. List of other heterogeneous catalysts used for photocatalytic CO₂-to-CO conversion.

movie S1. Three-dimensional reconstructed tomography of a fraction of the spongy Ni(TPA/TEG) composite.

movie S2. Scanning nanobeam diffraction pattern series showing the [100] orientation of the spongy Ni(TPA/TEG) composite.

movie S3. Scanning nanobeam diffraction pattern series taken along the [111] orientation of the spongy Ni(TPA/TEG) composite.

References (50–55)

REFERENCES AND NOTES

1. A. Cho, Energy's tricky tradeoffs. *Science* **329**, 786–787 (2010).
2. S. J. Davis, K. Caldeira, H. D. Matthews, Future CO₂ emissions and climate change from existing energy infrastructure. *Science* **329**, 1330–1333 (2010).
3. N. S. Lewis, D. G. Nocera, Powering the planet: Chemical challenges in solar energy utilization. *Proc. Natl. Acad. Sci. U.S.A.* **103**, 15729–15735 (2006).
4. S. R. Loarie, P. B. Duffy, H. Hamilton, G. P. Asner, C. B. Field, D. D. Ackerly, The velocity of climate change. *Nature* **462**, 1052–1055 (2009).
5. M. R. Allen, D. J. Frame, C. Huntingford, C. D. Jones, J. A. Lowe, M. Meinshausen, N. Meinshausen, Warming caused by cumulative carbon emissions towards the trillionth tonne. *Nature* **458**, 1163–1166 (2009).
6. K. B. Tokarska, N. P. Gillett, A. J. Weaver, V. K. Arora, M. Eby, The climate response to five trillion tonnes of carbon. *Nat. Clim. Change* **6**, 851–855 (2016).
7. R. A. Feely, C. L. Sabine, K. Lee, W. Berelson, J. Kleypas, V. J. Fabry, F. J. Millero, Impact of anthropogenic CO₂ on the CaCO₃ system in the oceans. *Science* **305**, 362–366 (2004).
8. T. Wheeler, J. von Braun, Climate change impacts on global food security. *Science* **341**, 508–513 (2013).
9. M. C. Urban, Accelerating extinction risk from climate change. *Science* **348**, 571–573 (2015).
10. J. S. Pal, E. A. B. Eltahir, Future temperature in southwest Asia projected to exceed a threshold for human adaptability. *Nat. Clim. Change* **6**, 197–200 (2016).
11. R. James, F. Otto, H. Parker, E. Boyd, R. Cornforth, D. Mitchell, M. Allen, Characterizing loss and damage from climate change. *Nat. Clim. Change* **4**, 938–939 (2014).
12. D. W. Keith, Why capture CO₂ from the atmosphere? *Science* **325**, 1654–1655 (2009).
13. N. S. Lewis, Research opportunities to advance solar energy utilization. *Science* **351**, aad1920 (2016).
14. M. Mikkelsen, M. Jorgensen, F. C. Krebs, The teraton challenge. A review of fixation and transformation of carbon dioxide. *Energy Environ. Sci.* **3**, 43–81 (2010).
15. J. L. White, M. F. Baruch, J. E. Pander III, Y. Hu, I. C. Fortmeyer, J. E. Park, T. Zhang, K. Liao, J. Gu, Y. Yan, T. W. Shaw, E. Abelev, A. B. Bocarsly, Light-driven heterogeneous reduction of carbon dioxide: Photocatalysts and photoelectrodes. *Chem. Rev.* **115**, 12888–12935 (2015).
16. K. Iizuka, T. Wato, Y. Miseki, K. Saito, A. Kudo, Photocatalytic reduction of carbon dioxide over Ag cocatalyst-loaded Al_{0.4}Ti_{0.6}O_{1.5} (A = Ca, Sr, and Ba) using water as a reducing reagent. *J. Am. Chem. Soc.* **133**, 20863–20868 (2011).

17. S. N. Habisreutinger, L. Schmidt-Mende, J. K. Stolarczyk, Photocatalytic reduction of CO₂ on TiO₂ and other semiconductors. *Angew. Chem. Int. Ed.* **52**, 7372–7408 (2013).
18. C. Gao, Q. Meng, K. Zhao, H. Yin, D. Wang, J. Guo, S. Zhao, L. Chang, M. He, Q. Li, H. Zhao, X. Huang, Y. Gao, Z. Tang, Co₃O₄ hexagonal platelets with controllable facets enabling highly efficient visible-light photocatalytic reduction of CO₂. *Adv. Mater.* **28**, 6485–6490 (2016).
19. S. Wang, W. Yao, J. Lin, Z. Ding, X. Wang, Cobalt imidazolate metal–organic frameworks photosplit CO₂ under mild reaction conditions. *Angew. Chem. Int. Ed.* **53**, 1034–1038 (2014).
20. A. Loidice, P. Lobaccaro, E. A. Kamali, T. Thao, B. H. Huang, J. W. Ager, R. Buonsanti, Tailoring copper nanocrystals towards C₂ products in electrochemical CO₂ reduction. *Angew. Chem. Int. Ed.* **55**, 5789–5792 (2016).
21. K. P. Kuhl, E. R. Cave, D. N. Abram, T. F. Jaramillo, New insights into the electrochemical reduction of carbon dioxide on metallic copper surfaces. *Energy Environ. Sci.* **5**, 7050–7059 (2012).
22. M. R. Singh, E. L. Clark, A. T. Bell, Thermodynamic and achievable efficiencies for solar-driven electrochemical reduction of carbon dioxide to transportation fuels. *Proc. Natl. Acad. Sci. U.S.A.* **112**, E6111–E6118 (2015).
23. K.-Y. Niu, F. Lin, S. Jung, L. Fang, D. Nordlund, C. C. L. McCrory, T.-C. Weng, P. Ercius, M. M. Doeff, H. Zheng, Tuning complex transition metal hydroxide nanostructures as active catalysts for water oxidation by a laser–chemical route. *Nano Lett.* **15**, 2498–2503 (2015).
24. J. Kothandaraman, A. Goepfert, M. Czaun, G. A. Olah, G. K. S. Prakash, Conversion of CO₂ from air into methanol using a polyamine and a homogeneous ruthenium catalyst. *J. Am. Chem. Soc.* **138**, 778–781 (2016).
25. H. Furukawa, K. E. Cordova, M. O’Keeffe, O. M. Yaghi, The chemistry and applications of metal–organic frameworks. *Science* **341**, 1230444 (2013).
26. K. Sumida, D. L. Rogow, J. A. Mason, T. M. McDonald, E. D. Bloch, Z. R. Herm, T.-H. Bae, J. R. Long, Carbon dioxide capture in metal–organic frameworks. *Chem. Rev.* **112**, 724–781 (2012).
27. W. Lu, Z. Wei, Z.-Y. Gu, T.-F. Liu, J. Park, J. Tian, M. Zhang, Q. Zhang, T. Gentle III, M. Bosch, H.-C. Zhou, Tuning the structure and function of metal–organic frameworks via linker design. *Chem. Soc. Rev.* **43**, 5561–5593 (2014).
28. C. G. Carson, K. Hardcastle, J. Schwartz, X. Liu, C. Hoffmann, R. A. Gerhardt, R. Tannenbaum, Synthesis and structure characterization of copper terephthalate metal–organic frameworks. *Eur. J. Inorg. Chem.* **2009**, 2338–2343 (2009).
29. Y. Tomita, S. Teruya, O. Koga, Y. Hori, Electrochemical reduction of carbon dioxide at a platinum electrode in acetonitrile–water mixtures. *J. Electrochem. Soc.* **147**, 4164–4167 (2000).
30. M. Kirch, J.-M. Lehn, J.-P. Sauvage, Hydrogen generation by visible light irradiation of aqueous solutions of metal complexes. An approach to the photochemical conversion and storage of solar energy. *Helv. Chim. Acta* **62**, 1345–1384 (1979).
31. T. W. Woolerton, S. Sheard, E. Pierce, S. W. Ragsdale, F. A. Armstrong, CO₂ photoreduction at enzyme-modified metal oxide nanoparticles. *Energy Environ. Sci.* **4**, 2393–2399 (2011).
32. H. Takeda, K. Ohashi, A. Sekine, O. Ishitani, Photocatalytic CO₂ reduction using Cu(I) photosensitizers with a Fe(II) catalyst. *J. Am. Chem. Soc.* **138**, 4354–4357 (2016).
33. S. Sato, T. Morikawa, S. Saeki, T. Kajino, T. Motohiro, Visible-light-induced selective CO₂ reduction utilizing a ruthenium complex electrocatalyst linked to a p-Type nitrogen-doped Ta₂O₅ semiconductor. *Angew. Chem. Int. Ed.* **49**, 5101–5105 (2010).
34. R. Kuriki, H. Matsunaga, T. Nakashima, K. Wada, A. Yamakata, O. Ishitani, K. Maeda, Nature-inspired, highly durable CO₂ reduction system consisting of a binuclear ruthenium(II) complex and an organic semiconductor using visible light. *J. Am. Chem. Soc.* **138**, 5159–5170 (2016).
35. U. Maitra, B. S. Naidu, A. Govindaraj, C. N. R. Rao, Importance of trivalency and the e_g¹ configuration in the photocatalytic oxidation of water by Mn and Co oxides. *Proc. Natl. Acad. Sci. U.S.A.* **110**, 11704–11707 (2013).
36. J. Luo, J.-H. Im, M. T. Mayer, M. Schreier, M. K. Nazeeruddin, N.-G. Park, S. D. Tilley, H. J. Fan, M. Grätzel, Water photolysis at 12.3% efficiency via perovskite photovoltaics and earth-abundant catalysts. *Science* **345**, 1593–1596 (2014).
37. M. W. Kanan, D. G. Nocera, In situ formation of an oxygen-evolving catalyst in neutral water containing phosphate and Co²⁺. *Science* **321**, 1072–1075 (2008).
38. X. Xie, Y. Li, Z.-Q. Liu, M. Haruta, W. Shen, Low-temperature oxidation of CO catalysed by Co₃O₄ nanorods. *Nature* **458**, 746–749 (2009).
39. S. Lin, C. S. Diercks, Y.-B. Zhang, N. Kornienko, E. M. Nichols, Y. Zhao, A. R. Paris, D. Kim, P. Yang, O. M. Yaghi, C. J. Chang, Covalent organic frameworks comprising cobalt porphyrins for catalytic CO₂ reduction in water. *Science* **349**, 1208–1213 (2015).
40. C. W. Li, M. W. Kanan, CO₂ reduction at low overpotential on Cu electrodes resulting from the reduction of thick Cu₂O films. *J. Am. Chem. Soc.* **134**, 7231–7234 (2012).
41. R. Kortlever, I. Peters, S. Koper, M. T. M. Koper, Electrochemical CO₂ reduction to formic acid at low overpotential and with high faradaic efficiency on carbon-supported bimetallic Pd–Pt nanoparticles. *ACS Catal.* **5**, 3916–3923 (2015).
42. K. P. Kuhl, T. Hatsukade, E. R. Cave, D. N. Abram, J. Kibsgaard, T. F. Jaramillo, Electrocatalytic conversion of carbon dioxide to methane and methanol on transition metal surfaces. *J. Am. Chem. Soc.* **136**, 14107–14113 (2014).
43. F. Gärtner, B. Sundararaju, A.-E. Surkus, A. Boddien, B. Loges, H. Junge, P. H. Dixneuf, M. Beller, Light-driven hydrogen generation: Efficient iron-based water reduction catalysts. *Angew. Chem. Int. Ed.* **48**, 9962–9965 (2009).
44. T. Maschmeyer, M. Che, Catalytic aspects of light-induced hydrogen generation in water with TiO₂ and other photocatalysts: A simple and practical way towards a normalization? *Angew. Chem. Int. Ed.* **49**, 1536–1539 (2010).
45. J. Shen, R. Kortlever, R. Kas, Y. Y. Birdja, O. Diaz-Morales, Y. Kwon, I. Ledezma-Yanez, K. J. P. Schouten, G. Mul, M. T. M. Koper, Electrocatalytic reduction of carbon dioxide to carbon monoxide and methane at an immobilized cobalt protoporphyrin. *Nat. Commun.* **6**, 8177 (2015).
46. A. Wuttig, M. Yaguchi, K. Motobayashi, M. Osawa, Y. Surendranath, Inhibited proton transfer enhances Au-catalyzed CO₂-to-fuels selectivity. *Proc. Natl. Acad. Sci. U.S.A.* **113**, E4585–E4593 (2016).
47. T. Cheng, H. Xiao, W. A. Goddard III, Full atomistic reaction mechanism with kinetics for CO reduction on Cu(100) from ab initio molecular dynamics free-energy calculations at 298 K. *Proc. Natl. Acad. Sci. U.S.A.* **114**, 1795–1800 (2017).
48. C. W. Li, J. Ciston, M. W. Kanan, Electroreduction of carbon monoxide to liquid fuel on oxide-derived nanocrystalline copper. *Nature* **508**, 504–507 (2014).
49. R. E. London, T. E. Walker, V. H. Kollman, N. A. Matwiyoff, Studies of the pH dependence of ¹³C shifts and carbon-carbon coupling constants of [U-13]aspartic and α-glutamic acids. *J. Am. Chem. Soc.* **100**, 3723–3729 (1978).
50. H. Jiang, Y. Cheng, L. Yuan, A Langmuir-like desorption model for reflecting the inhomogeneous pore structure of coal and its experimental verification. *RSC Adv.* **5**, 2434–2440 (2015).
51. S. Wang, X. Wang, Photocatalytic CO₂ reduction by CdS promoted with a zeolitic imidazolate framework. *Appl. Catal. B* **162**, 494–500 (2015).
52. S. Wang, Z. Ding, X. Wang, A stable ZnCo₂O₄ cocatalyst for photocatalytic CO₂ reduction. *Chem. Commun.* **51**, 1517–1519 (2015).
53. S. Wang, Y. Hou, X. Wang, Development of a stable MnCo₂O₄ cocatalyst for photocatalytic CO₂ reduction with visible light. *ACS Appl. Mater. Interfaces* **7**, 4327–4335 (2015).
54. R. Kuriki, K. Sekizawa, O. Ishitani, K. Maeda, Visible-light-driven CO₂ reduction with carbon nitride: Enhancing the activity of ruthenium catalysts. *Angew. Chem. Int. Ed.* **54**, 2406–2409 (2015).
55. H. Takeda, H. Koizumi, K. Okamoto, O. Ishitani, Photocatalytic CO₂ reduction using a Mn complex as a catalyst. *Chem. Commun.* **50**, 1491–1493 (2014).

Acknowledgments: We acknowledge C. Gammer for developing the Foundry users’ code for electron scanning diffraction experiments and analysis and Y. Liu at Lawrence Berkeley National Laboratory (LBNL) for help with FTIR spectroscopy measurements. **Funding:** We used TEM facilities in the Molecular Foundry at LBNL, which was supported by the U.S. Department of Energy (DOE), Office of Science, Office of Basic Energy Sciences, under contract #DE-AC02-05CH11231. Electron tomography studies used resources of the Center for Functional Nanomaterials, which is a DOE Office of Science facility, at Brookhaven National Laboratory under contract no. DE-SC0012704. This work was also supported by the Assistant Secretary for Energy Efficiency and Renewable Energy, Office of Vehicle Technologies of the DOE, under contract no. DE-AC02-05CH11231. All materials synthesis was carried out at LBNL and was supported by the DOE, Office of Science, Office of Basic Energy Sciences Materials Sciences and Engineering Division, under contract no. DE-AC02-05-CH11231 within the Early Career program MSE08 (to H.Z.). Y.X. and R.X. acknowledge financial support from Nanyang Technological University. K.N. and Y.L. were partially supported by the SinBeRise program of BEARS (Berkeley Education Alliance for Research in Singapore) at University of California, Berkeley. **Author contributions:** K.N. designed and performed the experiments, analyzed the experimental data, and wrote the manuscript. Y.X. and R.X. performed the photocatalysis experiments. H.W. performed the laser-chemical synthesis. H.L.X. performed the three-dimensional tomographic reconstruction. R.Y., F.L., C.T., Y.L., and K.C.B. helped on materials structure analyses. M.M.D., J.A., and M.T.M.K. contributed to the manuscript writing and catalytic mechanism discussions. All work was carried out under the supervision of H.Z. **Competing interests:** The authors declare that they have no competing interests. **Data and materials availability:** All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

Submitted 24 March 2017

Accepted 27 June 2017

Published 28 July 2017

10.1126/sciadv.1700921

Citation: K. Niu, Y. Xu, H. Wang, R. Ye, H. L. Xin, F. Lin, C. Tian, Y. Lum, K. C. Bustillo, M. M. Doeff, M. T. M. Koper, J. Ager, R. Xu, H. Zheng, A spongy nickel-organic CO₂ reduction photocatalyst for nearly 100% selective CO production. *Sci. Adv.* **3**, e1700921 (2017).

A spongy nickel-organic CO₂ reduction photocatalyst for nearly 100% selective CO production

Kaiyang Niu, You Xu, Haicheng Wang, Rong Ye, Huolin L. Xin, Feng Lin, Chixia Tian, Yanwei Lum, Karen C. Bustillo, Marca M. Doeff, Marc T. M. Koper, Joel Ager, Rong Xu and Haimei Zheng

Sci Adv **3** (7), e1700921.

DOI: 10.1126/sciadv.1700921

ARTICLE TOOLS

<http://advances.sciencemag.org/content/3/7/e1700921>

SUPPLEMENTARY MATERIALS

<http://advances.sciencemag.org/content/suppl/2017/07/24/3.7.e1700921.DC1>

PERMISSIONS

<http://www.sciencemag.org/help/reprints-and-permissions>

Use of this article is subject to the [Terms of Service](#)

Science Advances (ISSN 2375-2548) is published by the American Association for the Advancement of Science, 1200 New York Avenue NW, Washington, DC 20005. 2017 © The Authors, some rights reserved; exclusive licensee American Association for the Advancement of Science. No claim to original U.S. Government Works. The title *Science Advances* is a registered trademark of AAAS.