



Cite this: *Energy Environ. Sci.*, 2016, 9, 3161

A CO₂ adsorption-enhanced semiconductor/metal-complex hybrid photoelectrocatalytic interface for efficient formate production[†]

Xiaofeng Huang,^a Qi Shen,^a Jibo Liu,^a Nianjun Yang^{*b} and Guohua Zhao^{*a}

In photoelectrochemical CO₂ conversion, the concentration of fixed CO₂ on the photocathode surface is of primary concern. Herein, a CO₂ adsorption-enhanced semiconductor/metal-complex hybrid photoelectrocatalytic interface was established by utilizing a carbon aerogel as the CO₂ fixation substrate. In CO₂ reduction photoelectrocatalysis, Co₃O₄ was employed as the light harvester, and Ru(bpy)₂dppz was utilized as the electron transfer mediator and CO₂ activator. The CO₂ surface concentration exhibited a 380-fold increase on this hybrid interface than that on Co₃O₄/FTO. The CO₂ conversion to formate occurred at an onset potential of -0.45 V (vs. normal hydrogen electrode, NHE) under photoelectrochemical conditions, 160 mV more positive than its thermodynamic redox potential. At an applied potential of -0.60 V (vs. NHE), the selectivity of the formate yield reached 99.95%, with a production rate of approximately $110 \mu\text{mol cm}^{-2} \text{ h}^{-1}$ and a Faradaic efficiency of 86%. Such a conversion has an electron transfer rate of $2.94 \times 10^{-3} \text{ cm s}^{-1}$. The CO₂ conversion to formate was confirmed to be an instantaneous proton-coupled electron transfer process, originating from the rapid photoelectrochemical activation of bpy and dppz in Ru(bpy)₂dppz as well as the synergic effect of the promoted CO₂ adsorption and the applied molecular catalysis.

Received 1st April 2016,
Accepted 25th July 2016

DOI: 10.1039/c6ee00968a

www.rsc.org/ees

Broader context

In natural photosynthesis, CO₂ is fixed by ribulose-1,5-biphosphate carboxylase/oxygenase (RuBisCo) and then reduced by a regulated proton-coupled electron transfer process. Thus, there are three aspects to achieve a highly effective process of CO₂ conversion to fuels by mimicking natural photosynthesis: enhanced CO₂ adsorption, rapid electron transfer, and decreased energy input. For a high CO₂ surface concentration, molecular catalysts, such as Ru(ii) bipyridyl complexes, are promising candidates due to their unique properties in CO₂ binding affinity. The application of photoelectrochemical methods that integrate photocatalysis with electrocatalysis provides a promising strategy for CO₂ reduction at a low overpotential. This study demonstrates the use of highly porous and adsorptive carbon aerogels (CA) to promote CO₂ fixation as well as the employment of a robust visible-light harvester Co₃O₄ photoelectrocatalyst and Ru(bpy)₂dppz as a molecular catalyst to accelerate electron transfer for CO₂ reduction. This interface enables high CO₂ adsorption, a rapid electron transfer rate and a high yield of formate, an important industrial C1 stock. This reduction occurred at a relatively low onset potential under photoelectrochemical conditions and with a moderate Faradaic efficiency, originating from the multiple synergies among the enhanced CO₂ adsorption from CA, the distinguished photoelectrocatalytic activity of Co₃O₄ and the rapid electrochemical kinetics of Ru(bpy)₂dppz.

Introduction

Natural photosynthesis converts CO₂ into a series of intermediate products (e.g., malate and pyruvate) and finally to glucose with the assistance of solar light.¹ In such a carbon recycling process,

CO₂ is fixed by ribulose-1,5-biphosphate carboxylase/oxygenase (RuBisCo). Solar light is harvested, and the electrons are generated by cytochrome. A proton-coupled electron transfer is then regulated and mediated by nicotinamide adenine dinucleotide phosphate hydride, leading to the photochemical reduction of CO₂.¹ Unfortunately, such a process has a low conversion efficiency. Processes mimicking natural photosynthesis, so-called artificial photosynthesis, have been developed to increase this conversion efficiency and provide more pathways to produce valuable chemicals using CO₂ as the carbon stock.²⁻⁸

CO₂ adsorption is of the greatest concern during the CO₂ reduction process because the reduction kinetics are highly correlated with the CO₂ concentration. However, the free CO₂

^a School of Chemical Science and Engineering, Tongji University, 1239 Siping Road, 200092, Shanghai, China. E-mail: g.zhao@tongji.edu.cn

^b Institute of Materials Engineering, University of Siegen, Paul-Bonatz Str. 9-11, Siegen 57076, Germany. E-mail: nianjun.yang@uni-siegen.de

[†] Electronic supplementary information (ESI) available: Detailed experimental methods, SEM and TEM characterization, chronocoulometry, EQCM results, BET and BJH results, XRD patterns, CVs, UV-DRS, Tauc plots, Mott-Schottky plot, and RDE results. See DOI: 10.1039/c6ee00968a

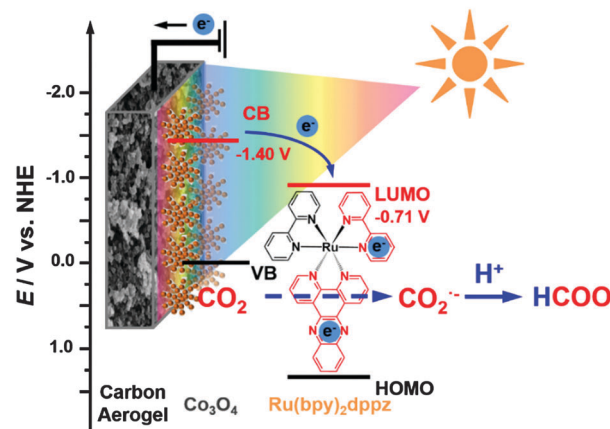


concentration in aqueous solution is only approximately 0.034 M, which severely deteriorates the aqueous heterogeneous reduction. Although various CO_2 absorptive materials have been proposed,^{9–11} their CO_2 adsorption performance and mechanism during catalytic processes have not been investigated in-depth. Moreover, the obtained efficiency for the CO_2 reduction still remains to be further increased.

Semiconductor-based photoelectrochemical methods are currently popular and have been successful for CO_2 reduction in that the synergistic conjunction of photoelectrocatalysis (namely, electrocatalysis and photocatalysis) facilitates the separation of photo-induced electrons and holes under applied electric fields.^{6,7} Light irradiation-induced band bending compensates the required overpotential for CO_2 reduction, causing the CO_2 conversion to occur at relatively positive potentials. Surprisingly, the majority of semiconductors exhibited non-specific CO_2 surficial binding. In other words, the CO_2 concentration on the surface of the semiconductors was rather low. Therefore, only slow reduction kinetics were reported.^{6,7} In addition, the majority of semiconductors have poor conductivities, which deteriorated the electron transfer rate as well. Thus, photoelectrochemical CO_2 reduction still occurred at relatively negative potentials in these systems, for example, at -1.0 V for $\text{MoS}_2/\text{TiO}_2$ and InP/TiO_2 (vs. normal hydrogen electrode, NHE, the same as below),^{12,13} at -1.1 V for ZnTe ,⁵ at -1.2 V for p-GaP and GaAs,¹⁴ and at -0.9 V for $\text{Mg}-\text{CuFeO}_2$.³ Furthermore, inevitable hydrogen evolution was observed, leading to relatively low Faradaic efficiency.

As an alternative to those semiconductors, molecular catalysts, such as Ru(II) or Re(II) bipyridyl complexes, have been employed because they present unique properties in CO_2 binding affinity, and high CO_2 surface concentrations are expected.^{15,16} Furthermore, the formation of a chemical bond between CO_2 and the metal centre will lead to a high turnover rate and high selectivity of CO_2 conversion during a catalytic cycle.^{15,16} The main drawback of these catalysts is that additional pathways are required to activate these molecular catalysts. In most cases, an extremely negative potential is required. For example, potentials of -1.52 , -1.6 , and -1.73 were applied for $\text{Ru}(\text{bpy})(\text{tpy})(\text{NCCH}_3)$,¹⁷ $\text{Re}(\text{tBu}-\text{bpy})(\text{CO})_3\text{Cl}$,¹⁸ and $\text{Re}(\text{pbn})(\text{CO})_3\text{Cl}$,¹⁹ respectively, to achieve the catalytic reduction of CO_2 . These molecular catalysts encountered destabilization (e.g., dimerization or dissolution) during a long-term electrolysis process, resulting in rapidly reduced photoelectrochemical activity for CO_2 conversion.^{20,21}

The complementary combination of a semiconductor and a molecular catalyst was then developed for photoelectrochemical CO_2 conversion.^{14,22–24} These combinations accelerated electron transfer generation and lowered the activation barriers for both molecular catalysts and CO_2 binding. The highly efficient and highly selective generation of hydrocarbon fuels, such as methanol and formic acid, has been reported on these systems.^{14,22–24} However, these systems have shown significant flaws, such as high overpotentials (e.g., -1.0 V on H-Si-Re(tBu-bpy)(CO)₃Cl),¹⁴ poor selectivity of reduction products (e.g., the simultaneous formation of HCOOH , H_2 and CO on Ag/TaON-Ru(bpy)₃-(CH₂)₂-Ru(bpy)(CO)₂Cl₂, while HCOOH



Scheme 1 Schematic plots of the CO_2 adsorption-enhanced $\text{Ru}(\text{bpy})_2\text{dppz}$ - $\text{Co}_3\text{O}_4/\text{CA}$ interface together with its energy level diagram and the possible reaction pathways for CO_2 conversion on this photocathode. Such a photoelectrocatalytic interface is composed of CA as the CO_2 -adsorption substrate, $\text{Ru}(\text{bpy})_2\text{dppz}$ as the molecular catalyst, and Co_3O_4 as the photoelectrocatalyst.

occupies only 56.5%),²³ and low Faradaic efficiency (e.g., only 62.3% on $[\text{Ru}(\text{dcbpy})_2(\text{CO})_2]_n\text{-p-InP-Zn}$).²⁴

Thus, the aim of this work is to construct an adsorptive photoelectrochemical interface for CO_2 reduction with better performance. The constructed interface is schematically shown in Scheme 1. In our concept, CO_2 fixation is promoted by utilizing an adsorptive substrate, a conductive, micropore-dominated three-dimensional carbon aerogel (CA) with a high surface area (up to $1815 \text{ m}^2 \text{ g}^{-1}$); solar light is efficiently harvested by Co_3O_4 micro-flowers, a visible-light driven and robust photoelectrocatalyst. These highly index-faceted Co(III)-enriched {121} structures are grown epitaxially inside CA networks. An enzyme-mimicking molecular catalyst, $\text{Ru}(\text{bpy})_2\text{dppz}$, is immobilized on $\text{Co}_3\text{O}_4/\text{CA}$ to accelerate and regulate the electron transfer for CO_2 reduction. Using such an interface, the CO_2 conversion to formate has been achieved at an onset reduction potential under photoelectrochemical conditions as low as -0.45 V, with a yield of approximately $110 \mu\text{mol cm}^{-2} \text{ h}^{-1}$, a selectivity of 99.95%, and a Faradaic efficiency of 86% at -0.60 V.

Experimental section

Fabrication of the $\text{Ru}(\text{bpy})_2\text{dppz}$ - $\text{Co}_3\text{O}_4/\text{CA}$ photocathode

Monolith bulky CA was synthesized via an ambient pressure resorcinol-formaldehyde (RF) drying method. The detailed synthesis process of CA is shown as Scheme S1 in the ESI.† A solvothermal reaction was utilized to synthesize $\text{Co}_3\text{O}_4/\text{CA}$. In a 50 mL acetone–water mixture ($V_{\text{acetone}} : V_{\text{water}} = 5 : 45$), 1.4552 g of $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, 0.2593 g of NH_4F , and 1.4019 g of hexamethylenetetramine were added. Magnetic stirring of this mixture for 10 min led to the formation of a pink transparent solution. The solution was then transferred to a Teflon-lined stainless steel autoclave, where one slide of CA ($4 \times 1.5 \text{ cm}^2$) was located. The solvothermal reaction was conducted at 95°C for 24 h. After cooling down to room temperature, deposition (pink to violet colour) occurred on the CA. The as-prepared

sample was carefully rinsed with acetone and then dried under vacuum at 60 °C for 2 h, followed by a calcination process in a N₂ atmosphere at 450 °C for 2 h. The ramping rate was 10 °C min⁻¹. After such a calcination process, the Co₃O₄/CA sample was synthesized. The amount of Co₃O₄/CA loading, averaged over the geometric area of the CA, was typically 8 mg cm⁻². A similar procedure was applied for the preparation of the control sample (Co₃O₄/FTO). The loading density of Co₃O₄ on FTO was 5 mg cm⁻². In this case, a piece of fluorine-doped tin oxide glass electrode (FTO, 4 × 1 cm²) was put into the autoclave instead of CA.

Chemical polymerization was used to decorate the Co₃O₄/CA with Ru(bpy)₂dppz (the detailed synthesis procedure for Ru(bpy)₂dppz is provided in the ESI†). Briefly, 0.0040 g of Ru(bpy)₂dppz (5.75 μmol) was dissolved in 1 mL of acetonitrile solution, denoted as solution A. Next, 0.0650 g of Fe(NO₃)₃·9H₂O and 8 μL of pyrrole were dissolved in 1 mL of ethanol, and the mixture was denoted as solution B. The stock solution for the electrode coating was obtained by mixing solutions A and B (with equal volumes) and shaking. One slide of Co₃O₄/CA was then coated with 200 μL of the stock solution in a dropwise manner, and the composite was dried at 60 °C for 5 min. After repeating the coating procedure 10 times, Ru(bpy)₂dppz-Co₃O₄/CA was obtained.

Electrochemical quartz crystal microbalance

To calculate the amount of CO₂ adsorbed onto the photocathodes, electrochemical quartz crystal microbalance (EQCM) experiments were conducted on a CHI440A (CH Instruments Inc., USA) with a Au-coated AT-cut quartz crystal (a fundamental frequency of 8 MHz) as the working electrode.

Approximately 10 mg of ground CA powder was ultrasonically dispersed in 1 mL of H₂O along with 20 μL of 2% Nafion-117 film solution (Alfa Aesar). After intense ultrasonication for 5 min, 10 μL of the dispersed solution was dip-coated onto the Au electrode and dried in air. The CA-modified electrode was further used in the EQCM studies.

In situ IR spectroelectrochemical experiments

For the *in situ* IR spectroelectrochemical experiments, the mercury–cadmium–telluride detector was cooled down to 77 K by liquid nitrogen. A self-made four-bottleneck cell with a CaF₂ window (2 mm thickness, 25 mm in diameter) was used as the electrochemical cell. A Pt disk electrode (5 mm in diameter), a Ag/AgCl (filled with saturated KCl) electrode, and a Pt foil (1 × 1.5 cm²) were used as the working, reference and counter electrodes, respectively. A mixture of 0.010 g mL⁻¹ Ru(bpy)₂dppz and 2% Nafion-117 was stirred in a sonication bath for 1 min. This mixture (10 μL) was dip-coated onto the Pt disk electrode and dried under ambient conditions. One hundred IR spectra were collected with a spectrum resolution of 8 cm⁻¹ and subsequently averaged at each potential. For these experiments, the solution was purged with CO₂ for at least 30 min to completely remove the dissolved oxygen and saturate the solution with CO₂.

Photoelectrochemical characterization and reduction of CO₂

To evaluate the performance of the Ru(bpy)₂dppz-Co₃O₄/CA catalyst, constant potential photoelectrolysis of CO₂ was conducted

in order to evaluate the performance of Ru(bpy)₂dppz-Co₃O₄/CA, which was conducted in a home-made H-type cell with a maximum volume of 100 mL. The as-prepared Ru(bpy)₂dppz-Co₃O₄/CA working electrode and the Ag/AgCl (filled with saturated KCl) reference electrode were placed in the cathodic chamber, while the counter electrode, a graphite plate (4 × 1 cm²), was placed in the anodic chamber. The two chambers were connected with 0.1 M NaHCO₃ but separated with a Nafion-117 proton exchange membrane (Dupont). Prior to the experiments, the electrolyte in the cathodic chamber was purged with high-purity CO₂ (99.99%) gas for more than 30 min at a flow rate of 20 mL min⁻¹. Negative potentials (0.0, -0.2, -0.4, -0.6, -0.8, -1.0 V) were applied to the photocathode through the electrochemical workstation. An APLS-SXE300 xenon lamp with a UV cutoff ($\lambda > 420$ nm, light intensity at 9 mW cm⁻²) was used as the light source and illuminated on the Ru(bpy)₂dppz-Co₃O₄/CA photocathode upon the addition of negative potential.

After such constant potential photoelectrolysis for 8 h, the reduction products were collected and quantitatively determined by HPLC and GC using the same procedure as described previously.²⁵ For the products in the aqueous phase, 0.2 mL of the liquid sample was collected and transferred into a 10 mL test-tube. The pH of the sample was adjusted to neutral by adding 0.2 mL of pH 7.6 phosphate buffer solution. Subsequently, 2,3,4,5,6-pentafluorobenzyl bromide (20 g L⁻¹, 1.0 mL) was added. The mixture was shaken for 1 min and then kept at 60 °C for another 1 h. The esterification product was extracted with 2.0 mL of *n*-hexane and centrifuged at 3000 rpm for 5 min. The upper layer was the organic phase, which was filtered through a 0.45 μm membrane. A C18 column was used with a mobile phase consisting of 65% methanol and 35% H₂O at a flow rate of 1.0 mL min⁻¹. The detection wavelength was 225 nm. For gaseous products, 1.0 mL of the gas sample was collected through a syringe. The detection conditions were an injection inlet temperature of 130 °C, an oven temperature of 80 °C, a detector temperature of 150 °C, N₂ carrier gas, and a gas flow rate of 0.2 L min⁻¹.

Prior to an isotopic ¹³C experiment, ¹³CO₂ (¹³C enrichment 98%) was purged into 0.1 M NaH¹³CO₃ (¹³C enrichment 98%) electrolyte solution for at least 30 min in order to fully expel oxygen and other impurity gases. The photoelectrochemical reduction of ¹³CO₂ saturated NaH¹³CO₃ (0.1 M) was identical to the procedure described in the Experimental section, holding the constant potential at -1.2 V vs. Ag/AgCl. Blank experiments using nitrogen purged Na₂SO₄ (0.1 M) were also conducted using an identical procedure. After reduction, 0.5 mL of catholyte solution was mixed with 0.1 mL of D₂O (Sigma Aldrich) containing 0.5 μL of DMSO as the internal standard. A one-dimensional ¹H nuclear magnetic resonance (NMR) spectrum was recorded with water suppression using a pre-saturation method.

Results and discussion

Enhanced CO₂ adsorption

For photoelectrochemical CO₂ conversion, the surface concentration of CO₂ on the photocathode, namely, CO₂ fixation, is a

chief concern. A high surface concentration of CO_2 on the photocathode accelerates the photoelectrochemical kinetics. Herein, the surface concentration of CO_2 (Γ_{ads}), the normalized amount of adsorbed CO_2 with the electrochemical active surface area (S_{EASA}), was adopted as the parameter to evaluate the efficiency of CO_2 fixation on the photocathodes.

The results in Table S1 (ESI†) demonstrate the advantage of using a CA substrate for CO_2 fixation. This statement is further supported by the EQCM results of CA in a CO_2 -saturated electrolyte under negative potentials, as shown in Fig. S1 (ESI†). The mass addition is 6–10 times heavier on activated CA than that on an Au electrode. For example, at -0.4 V, a mass addition of 174 ng cm^{-2} was obtained on a CA substrate, whereas it was barely observed on an Au quartz substrate. At -0.6 V, the mass addition on the CA substrate was 8.2-fold larger than that on the Au quartz substrate. At -0.9 V, the mass addition reached 143.5 ng cm^{-2} on the CA substrate. When the applied potentials were more negative than -0.9 V, the mass addition on the CA substrate was even larger than that on the Au quartz substrate. CA is not capable of reducing CO_2 electrochemically; thus, the mass addition on the CA substrate is mainly ascribed to the promoted electro-sorption of carbonaceous species (e.g., CO_2 and HCO_3^-) from the electrolyte. The carbonaceous species are significantly promoted because the micropores contributed a high Brunauer–Emmett–Teller surface area (S_{BET}) of up to $1815 \text{ m}^2 \text{ g}^{-1}$ (Fig. S2, ESI†).

This high S_{BET} offers numerous sites for CO_2 adsorption, as indicated by Fujishima *et al.* and Yaghi *et al.*,^{21,26} as CO_2 molecules tend to be adsorbed in the micropores of a material. This value is also higher than that of common porous carbon materials, such as ordered mesoporous carbon ($812.3 \text{ m}^2 \text{ g}^{-1}$), commercially available Vulcan ($237.9 \text{ m}^2 \text{ g}^{-1}$) and carbon nanocoil ($233 \text{ m}^2 \text{ g}^{-1}$).²⁷ The superior performance of CA supports the notion that CO_2 adsorption on other high-surface-area carbon materials (e.g., activated carbon, carbon nanotubes) might result from the following aspects. First, the microporous feature of CA (Fig. S2, ESI†) offers numerous sites for CO_2 adsorption. Second, the activated CA possesses a relatively high surface area ($1815 \text{ m}^2 \text{ g}^{-1}$), which provides many sites for Co_3O_4 loading. According to our measurements, the Co_3O_4 loading amount was increased to 8 mg cm^{-2} compared to 5 mg cm^{-2} on FTO. The S_{BET} of $\text{Co}_3\text{O}_4/\text{CA}$ of $713 \text{ m}^2 \text{ g}^{-1}$ (Fig. S2a, ESI†) is considerably higher than that of commonly designed porous inorganic semiconductor electrodes.^{28,29} This result is further supported by the well-maintained micropore domination of the electrode (Fig. S2b, ESI†). Based on the voltammetry (Fig. S3, ESI†) and chronocoulometry (Fig. S4, ESI†) for $\text{Co}_3\text{O}_4/\text{CA}$, Γ_{ads} exhibits a 20-fold increment ($0.25 \text{ pmol cm}^{-2}$) compared to that ($0.01 \text{ pmol cm}^{-2}$) of $\text{Co}_3\text{O}_4/\text{FTO}$. The electrochemically active surface area (S_{EASA}) was determined to be 8015 cm^2 (Table S1, calculated from Fig. S5, ESI†) with respect to that of $\text{Co}_3\text{O}_4/\text{FTO}$ (316 cm^2). Experimental tests on the effects on CO_2 fixation of other porous carbon-based substrates are currently in progress.

As shown in the scanning electron microscopic (SEM) images of $\text{Co}_3\text{O}_4/\text{CA}$ (Fig. S6, ESI†), the CA backbone is clearly

visible even after the solvothermal growth of Co_3O_4 microflowers. The XRD patterns of both CA and Co_3O_4 in $\text{Co}_3\text{O}_4/\text{CA}$ can also be clearly observed (Fig. S7, ESI†). In other words, such an epitaxial growth pattern fully exposes the adsorption sites to CO_2 , which results in an increase in the Γ_{ads} for CO_2 . Moreover, the 3D structure of CA allows for the penetration of electrolytes into the pores of the electrode, leading to an increase in the interfacial area between the electrode and the electrolyte. Finally, CA possesses a high conductivity (electrical resistivity $< 40 \text{ m}\Omega$), similar to those of carbon nanotubes and graphene.

Γ_{ads} further increases to $3.79 \text{ pmol cm}^{-2}$ when $\text{Co}_3\text{O}_4/\text{CA}$ is coated with a molecular catalyst, $\text{Ru}(\text{bpy})_2\text{dppz}$, which is 15.2 and 300 times larger compared to those on $\text{Co}_3\text{O}_4/\text{CA}$ and $\text{Co}_3\text{O}_4/\text{FTO}$, respectively. However, the S_{EASA} of $\text{Co}_3\text{O}_4/\text{CA}$ is approximately 4 times larger than that of $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, likely due to the blockage of $\text{Co}(\text{III})$ active sites by immobilized $\text{Ru}(\text{bpy})_2\text{dppz}$ during the chemical polymerization. Nonetheless, the active area of this electrode is maintained at $1.64 \times 10^{-2} \text{ cm}^2$ (Table S1 and Fig. S3, ESI†). The existence of the $\text{Ru}(\text{bpy})_2\text{dppz}$ molecular catalyst promotes CO_2 fixation on the photocathode with an enhancement factor of 300. Γ_{ads} was also normalized by the catalyst weight (Table S1, ESI†) and exhibited the same trends as that normalized by S_{EASA} . Hence, the enhancement of CO_2 adsorption and photoelectrochemical density is in fact the synergic effect of multiple factors: the electrosorption from the CA substrate, the $\text{Co}(\text{III})$ sites from the high index faceted Co_3O_4 on CA and the electrochemical activity of $\text{Ru}(\text{bpy})_2\text{dppz}$. These results confirm that CA, Co_3O_4 and $\text{Ru}(\text{bpy})_2\text{dppz}$ co-promote CO_2 fixation and lead to a high CO_2 surface concentration on such an interface. Thus, accelerated kinetics are expected for photoelectrochemical CO_2 conversion.

Photoelectrochemical CO_2 conversion

The reactivity of $\text{Ru}(\text{bpy})_2\text{dppz}$ as a homogenous catalyst toward electrochemical CO_2 conversion was first studied in CO_2 -saturated NaHCO_3 solution. As shown in Fig. S8 (ESI†), the CO_2 reduction potential remains at -0.40 V in the current aqueous solution, which is at least 950 mV more positive than those reported in organic solvents.³⁰ This is the first time that a molecular catalyst, polybipyridyl $\text{Ru}(\text{II})$, was observed to convert/reduce CO_2 at such a low potential, indicating the high electron transfer ability of $\text{Ru}(\text{bpy})_2\text{dppz}$ toward CO_2 conversion.

CO_2 conversion on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ was investigated in detail using a potentiodynamic mode. As seen in Fig. 1a, CO_2 conversion/reduction occurs at an onset potential of -0.45 V under photoelectrochemical conditions (red line). This potential is an “underpotential” of approximately 160 mV with respect to the thermodynamic redox potential for CO_2 to formic acid.³¹ The peak potential is approximately -0.61 V, considerably lower than other reported values, e.g., -1.33 V on $\text{p-Cu}_2\text{O}$ immobilized by $\text{Re}(\text{tBu-bipy})(\text{CO})_3\text{Cl}$ ³² and -0.6 V on p-InP-Zn decorated by $[\text{Ru}(\text{dcbpy})_2(\text{CO})_2]_n$.²⁴ On $\text{Co}_3\text{O}_4/\text{CA}$, the reduction potential is approximately -0.62 V with an onset potential of -0.52 V under photoelectrochemical conditions (dashed lines). Moreover, a second peak appears at -0.73 V. On $\text{Co}_3\text{O}_4/\text{FTO}$, the onset potential is -0.67 V (dotted lines), indicating an entirely



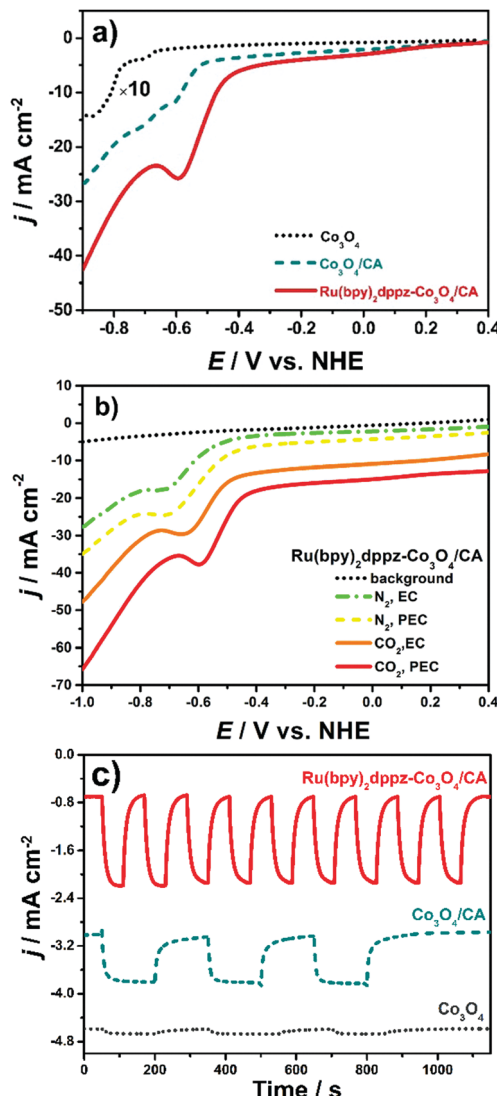


Fig. 1 (a) Linear sweep voltammograms (LSVs) of $\text{Co}_3\text{O}_4/\text{FTO}$ (dotted line), with a magnification factor of 10), $\text{Co}_3\text{O}_4/\text{CA}$ (dashed line) and $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ (solid line) in CO_2 -saturated 0.1 M NaHCO_3 at a scan rate of 50 mV s^{-1} under illumination conditions; (b) LSVs of $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ in N_2 -purged $0.1 \text{ M Na}_2\text{SO}_4$ (dotted line), N_2 -purged 0.1 M NaHCO_3 with (dashed line)/without (dash-dotted line) light irradiation, and CO_2 -saturated 0.1 M NaHCO_3 without (solid line) light irradiation at a scan rate of 50 mV s^{-1} ; (c) amperometric $i-t$ curves on $\text{Co}_3\text{O}_4/\text{FTO}$ (dotted line), $\text{Co}_3\text{O}_4/\text{CA}$ (dashed line) and $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ (solid line) in CO_2 -saturated $0.1 \text{ M Na}_2\text{SO}_4$ at a potential of -0.40 V . The lines were shifted in the Y-axis direction for comparison.

different photoelectrochemical reduction pathway. The positive shift of the CO_2 reduction potential on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ is likely due to the increased surface concentration of CO_2 and the catalytic role of $\text{Ru}(\text{bpy})_2\text{dppz}$. Based on these peak potentials, the heterogeneous electron transfer rate constant (k_s) for CO_2 reduction was calculated using a method provided in the literature.³³ k_s is $2.94 \times 10^{-3} \text{ cm s}^{-1}$ for $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, which is 26% larger than that ($2.19 \times 10^{-3} \text{ cm s}^{-1}$) for $\text{Co}_3\text{O}_4/\text{CA}$.

This higher k_s value suggests an accelerated and enhanced electron transfer on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$. Such a method to

evaluate the efficiency of CO_2 conversion is entirely new. According to our knowledge, this method has not been presented in the literature to date. Considering the CO_2 reduction potential and k_s on both electrodes, the catalytically active centre for CO_2 reduction is $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, not $\text{Co}(\text{m})$. This statement is further demonstrated from the Tafel plots described below.

The current density reaches approximately 8.1 mA cm^{-2} on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, which is considerably more intense than sole $\text{Ru}(\text{bpy})_2\text{dppz}$ and those reported using similar molecular catalyst–semiconductor composites. For instance, on $\text{Ru}(\text{dcbpy})_2(\text{CO})_2/\text{p-InP}$ and CO -dehydrogenase/p-NiO, the photocurrent density reached only the level of $\mu\text{A cm}^{-2}$ at similar potentials. A photocurrent density larger than -2.0 mA cm^{-2} was obtained on a $\text{Re}(\text{tBu-bpy})(\text{CO})_3\text{Cl}/\text{Cu}_2\text{O}$ photocathode, but a potential of -2.0 V was applied.³² So, photoelectrochemical CO_2 conversion on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ occurs at a low overpotential but with a high photocurrent density.

The current densities shown in this paper were obtained using the geometric areas of the electrodes. Due to their porous structures, CA-based photoelectrodes will have a higher S_{EASA} value. Their S_{EASA} values were determined using surface-sensitive redox probes of $\text{Fe}(\text{CN})_6^{3-/-4-}$ (Table S1 and Fig. S3, ESI[†]). A 2.7-fold enhancement was observed after loading the same amount of Co_3O_4 onto FTO. The current densities were then re-calculated using their S_{EASA} values. The magnitude of the current densities followed the same trend, namely, in the order of $\text{Co}_3\text{O}_4/\text{FTO} < \text{Co}_3\text{O}_4/\text{CA} < \text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, the same as that obtained from the current densities normalized by the geometric area. To compare our results with those obtained using other porous materials presented in the literature, the current densities shown throughout the paper were then calculated using the geometric areas of the photoelectrodes, the most frequently applied approach for electrochemical and photoelectrochemical CO_2 reduction.

Fig. 1b shows the LSVs on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ in 0.1 M NaHCO_3 solution purged with N_2 or CO_2 . The pH value of 0.1 M NaHCO_3 is in the range of 8.3–8.5 after being purged with N_2 . Once it was saturated with CO_2 , the pH value decreased to the range of 6.5–7.0.³⁴ Such a weak acidic environment provides a favourable protic environment for CO_2 reduction using a $\text{Ru}(\text{bpy})_2\text{dppz}$ catalyst.^{2–5} Notably, the peak potential for CO_2 reduction is 50 mV more positive than that of its N_2 -purged counterpart, with the peak current density doubled. These results clearly confirm the involvement of protons in CO_2 reduction. Fig. 1b shows the effect of light irradiation on CO_2 conversion as well. On $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ without light irradiation, the peak current density of the CO_2 reduction decreases by a ratio of 35% along with a 50 mV negative shift in the peak potential compared to the case when light irradiation is applied. Similar tendencies were observed on $\text{Co}_3\text{O}_4/\text{CA}$ and $\text{Co}_3\text{O}_4/\text{FTO}$ (Fig. S9, ESI[†]). This phenomenon could be due to the photoelectrocatalytic properties of the $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ electrode.

An amperometric photocurrent response was further investigated at a fixed potential of -0.40 V . As shown in Fig. 1c, the steady photocurrent density reaches approximately 1.5 mA cm^{-2}

on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, with a stabilizing time of only 40 s, whereas on $\text{Co}_3\text{O}_4/\text{CA}$, the peak photocurrent intensity decreases by approximately 7 times compared to $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$. However, the peak photocurrent intensity is 8–10 times higher than that on $\text{Co}_3\text{O}_4/\text{FTO}$. The stabilization times for $\text{Co}_3\text{O}_4/\text{CA}$ and $\text{Co}_3\text{O}_4/\text{FTO}$ are 150 and 200 s, respectively. The potentiodynamic voltammograms in 0.1 M Na_2SO_4 also reflected that $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ showed a distinguished photoelectrochemical activity. The difference between the photocurrent (j_{PEC}) and dark current (j_{EC}), denoted as $j_{\text{PEC}} - j_{\text{EC}}$, is summarized in Fig. S10 (ESI†). $j_{\text{PEC}} - j_{\text{EC}}$ is approximately 10 times higher on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ than on FTO. This result could be explained by several reasons. First, the $\{12\bar{1}\}$ crystal facet of Co_3O_4 (with an inter-planar spacing of 0.286 nm) being finely exposed (Fig. S11, ESI†), which enriches triply uncoordinated Co(III) sites. Such a high-index-facet Co_3O_4 facilitates the photoelectrochemical reduction of CO_2 , as our previous work has indicated.⁴ Second, the epitaxial growth manner of high-index-facet Co_3O_4 on CA provided a direct electron transfer channel. Photo-induced holes could rapidly transfer to the highly conductive CA network. Electrons and holes are efficiently separated. Finally, the ingenious merging of photocatalysis and electrocatalysis on a single surface has been found to efficiently separate the photo-induced carriers, resulting in a reduction of the overpotential for CO_2 conversion.^{2,4} The applied negative potential creates a more upward bending for Co_3O_4 , a p-type semiconductor. Then, the driving force for photoelectrons to cross the semiconductor–electrolyte junction is enlarged, resulting in an enhanced photocurrent density. In the meantime, the light-induced upward band-bending of $\text{Co}_3\text{O}_4/\text{CA}$ lifts the Fermi level, which compensates part of the required applied negative potential under dark conditions, yielding a reduced overpotential for CO_2 reduction. All of these effects reveal the essential role of Co_3O_4 as the photoelectrocatalyst in a photoelectrochemical strategy for CO_2 conversion.

Comparing the cyclic voltammograms of different photoelectrodes, the lowest overpotential and the highest photocurrent density for CO_2 conversion were obtained on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$. Apart from the respective contributions from the enhanced adsorption on CA and distinguished photoelectrochemical properties of $\text{Co}_3\text{O}_4/\text{CA}$, the molecular catalyst $\text{Ru}(\text{bpy})_2\text{dppz}$ also exhibits excellent electrochemical reductive activity toward CO_2 reduction, as suggested by the above-mentioned k_s . The synergistic effect among those effects exerted on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ has superiority in magnifying the photocurrent density, reducing the overpotential, and accelerating and regulating the electron transfer pathway for the photoelectrochemical CO_2 reduction process.

The photoelectrochemical CO_2 conversion on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ was further studied in CO_2 -saturated 0.1 M NaHCO_3 by varying the applied potential. Its variation with the potentials applied is summarized in Fig. 2a. The amplitude of $j(\text{CO}_2) - j(\text{N}_2)$ increases when the potential is more negative, following the order of $\text{Co}_3\text{O}_4/\text{FTO} < \text{Co}_3\text{O}_4/\text{CA} < \text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$. A sharp increase in the amplitude of $j(\text{CO}_2) - j(\text{N}_2)$ occurs at -0.4 V. At a reduction potential of -0.6 V, the net

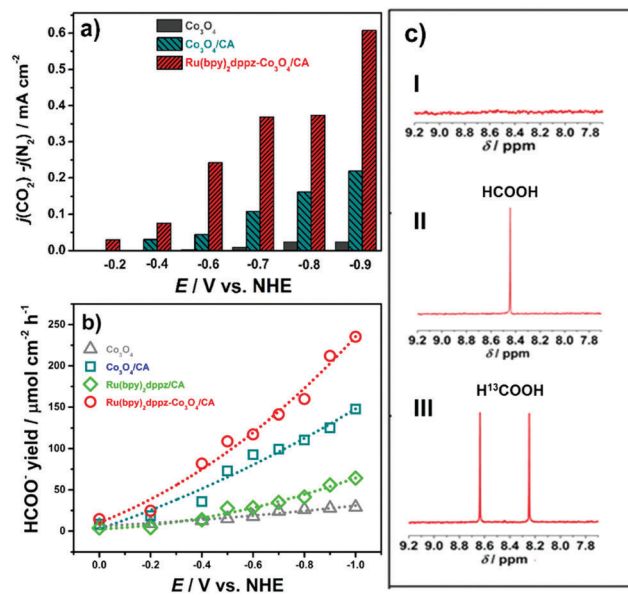


Fig. 2 (a) Variation of $j(\text{CO}_2) - j(\text{N}_2)$, on $\text{Co}_3\text{O}_4/\text{FTO}$ (grey), $\text{Co}_3\text{O}_4/\text{CA}$ (cyan) and $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ (red) with the applied potential; (b) variation of the yield rates of formate on the photocathode of $\text{Co}_3\text{O}_4/\text{FTO}$ (gray triangle), $\text{Ru}(\text{bpy})_2\text{dppz}/\text{CA}$ (green rhombus), $\text{Co}_3\text{O}_4/\text{CA}$ (cyan square) and $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ (red circle) with the applied potentials. The dashed lines are for guiding the eyes. (c) ^1H NMR spectra of the reduction product in N_2 -purged Na_2SO_4 (I), CO_2 -saturated NaHCO_3 (II) and $^{13}\text{CO}_2$ -saturated $\text{NaH}^{13}\text{CO}_3$ (III). The concentrations of these solutions were 0.1 M.

photocurrent reaches -0.35 mA cm^{-2} . This value is twice as high as that on $\text{Co}_3\text{O}_4/\text{CA}$ and nearly 10 times higher than that on $\text{Co}_3\text{O}_4/\text{FTO}$.

Under identical photoelectrolysis conditions, the yield rate of formate on these four photocathodes is in the order of $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA} > \text{Co}_3\text{O}_4/\text{CA} > \text{Ru}(\text{bpy})_2\text{dppz}/\text{CA} > \text{Co}_3\text{O}_4/\text{FTO}$ at the same applied potentials, as shown in Fig. 2b. For example, $869.8 \mu\text{mol}$ formate was produced at a potential of -0.6 V on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$. This value is considerably higher than that presented previously ($384 \mu\text{mol}$) on hierarchical Co_3O_4 , even at -0.7 V.⁵ If the electrode area is considered, the predicted yield rate of formate on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ is approximately $450 \mu\text{mol cm}^{-2}$ at the onset potential of -0.45 V under photoelectrochemical conditions. At -0.7 V, the yield rate of formate reaches approximately $900 \mu\text{mol cm}^{-2}$. Considering the reduction time span further, the yield rate of formate reaches nearly $110 \mu\text{mol cm}^{-2} \text{ h}^{-1}$, even at a reduction potential of -0.6 V. The yield rate of formate on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ is higher than most reported values, e.g., nearly 29-, 5.2-, and 2-fold higher than that on $\text{Ru}(\text{II})-\text{Re}(\text{I})$ multinuclear complexes,²² $\text{Ru}(\text{II})$ electro-polymerized p-InP-Zn,²⁴ and $\text{Cu}_2\text{ZnSnS}_4$,³⁵ respectively. The estimated Faradaic efficiency for the yield of formate is approximately 86%, which is comparable to that on semiconductor–molecular photocatalytic systems as well as on other semiconductor-based photocatalytic systems, such as Mg-doped CuFeO_2 ³ and $\text{Au}/\text{ZnO}/\text{ZnTe}$.⁵ Notably, formate yield rates on the $\text{Ru}(\text{bpy})_2\text{dppz}/\text{CA}$ were lower than that on the $\text{Co}_3\text{O}_4/\text{CA}$ at each investigated potential, the reason is described below.

To confirm the carbon source for the reduction product of formate, isotopic measurements were conducted using $^{13}\text{CO}_2$ -saturated $\text{NaH}^{13}\text{CO}_3$ (0.1 M) electrolyte solution.^{36,37} Control experiments were also performed in N_2 -purged Na_2SO_4 (0.1 M) solutions. These related $^1\text{H-NMR}$ spectra are shown as Fig. 2c. The spectrum obtained from the N_2 -purged Na_2SO_4 (0.1 M) solutions is featureless (Fig. 2c-I) in the chemical shift range of 9.2 to 7.7, indicating that no formate is produced when no carbonaceous species (e.g., CO_2 , HCO_3^-) are present. In Fig. 2c-II, a singlet peak at the chemical shift value of 8.42 is detected, likely resulting from the yield of HCOO^- from CO_2 -saturated NaHCO_3 . When $^{13}\text{CO}_2$ -saturated $\text{NaH}^{13}\text{CO}_3$ solution is used, a doublet peak is shown at the chemical shift values of 8.64 and 8.25 (Fig. 2c-III). This doublet peak indicates the production of $\text{H}^{13}\text{COO}^-$.³ In other words, formate is obtained from the ^{13}C species in the electrolyte rather than from the impurities in the electrolyte or on the surface of the photocathode. Photoelectrochemical CO_2 reduction thus occurs on the adsorption-enhanced molecular catalyst–semiconductor hybrid interface.

To investigate the energy conversion efficiency, the turnover number (TON) and turnover frequency (TOF) for photoelectrochemical CO_2 reduction were estimated at different potentials on this interface. These values are tabulated in Tables S2 and S3 (ESI†). $\text{Co}_3\text{O}_4/\text{CA}$ has a 3- to 5-fold increment in both TON and TOF compared to $\text{Co}_3\text{O}_4/\text{FTO}$, resulting from the effect of improved CO_2 adsorption. This result further demonstrates the importance of the primary CO_2 fixation process. On the $\text{Ru}(\text{bpy})_2\text{dppz}$ molecular catalyst immobilized interface, the magnitudes of both TON and TOF were increased by approximately 1 order. This trend was also proven by the results from $[\text{Zn}(\text{II})\text{TRP}]^{4+}/[\text{SiW}_{12}\text{O}_{40}]^{4-}$ ³⁸ and COF-367-Co,²¹ both of which possess a high surface area, thus resulting in a high TON value. The TON value was increased with the negative shift of the potential. For example, at -0.6 V vs. NHE, the TON value already reached 978.7, with a TOF value of 122. In Table 1, these results are further compared with those shown in the literature. The higher TON value (978.7) in our case compared to those reported (including those of similar semiconductor–molecular complex model systems (e.g., $\text{InP}/[\text{MCE2-A} + \text{MCE4}]\text{-TiO}_2/\text{Pt}$,³⁹ the NiO-RuRe complex⁴⁰ and $\text{Ru}(\text{dcbpy})/\text{N-Ta}_2\text{O}_5$ ⁴¹)) demonstrates the distinguished photoelectrochemical performance of the concurrent $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ system.

Hydrogen, as the only side product detected, only occupied 0.05% of the total products, which is indicative of the high

selectivity of $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ toward CO_2 reduction. On $\text{Co}_3\text{O}_4/\text{CA}$, the hydrogen production is 3 times (Fig. S12b, ESI†) higher, but the yield rate of formate is approximately 1.5 times lower (Fig. 2b). On $\text{Ru}(\text{bpy})_2\text{dppz}/\text{CA}$ (Fig. S12c, ESI†), lower amounts of H_2 were detected after applying a potential of -0.9 V. On $\text{Co}_3\text{O}_4/\text{FTO}$ (Fig. S12d, ESI†), the yield rate of formate is the lowest, although no hydrogen is detected. The selectivity for $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ is higher than those of $\text{Ru}(\text{n})$ -electro-polymerized p-InP-Zn^{24} and $\text{Cu}_2\text{ZnSnS}_4$,³⁵ in which non-negligible amounts of H_2 and CO were generated. Therefore, the selectivity of the CO_2 conversion to formate is extremely high on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ (99.95%).

The hydrogen evolution potentials were further estimated based on the potential-dependent hydrogen evolution profiles shown in Fig. 2b and Fig. S12a–d (ESI†). The potentials on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, $\text{Co}_3\text{O}_4/\text{CA}$, $\text{Ru}(\text{bpy})_2\text{dppz}/\text{CA}$ and $\text{Co}_3\text{O}_4/\text{FTO}$ are -0.6 V, -0.7 V and -0.9 V vs. NHE, respectively. However, there are no clear cathodic peaks shown in Fig. 1 for hydrogen evolution because the $\text{Ru}(\text{n})$ centre on the molecular catalyst $\text{Ru}(\text{bpy})_2\text{dppz}$ has fully coordinated to the nitrogen atoms from two 2,2'-bipyridine and dppz ligands. Then, both the bipyridine ligand and dppz are not able to dissociate from $\text{Ru}(\text{n})$ during the electrochemical processes. In other words, there will be no opportunity to generate any catalytic wave for hydrogen evolution. These results indicate that this unwanted reaction has not been involved in our system during CO_2 reduction. Thus, an improved conversion efficiency and high selectivity are expected.

The stability of the proposed photoelectrocatalytic interface was examined by recording the XRD patterns and cyclic voltammograms of $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ before and after a long-term photoelectrochemical CO_2 reduction. The XRD patterns of both CA and Co_3O_4 do not vary after the 8 h photoelectrochemical reduction of CO_2 (Fig. S13, ESI†). The peak current density of $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ is slightly enhanced after 100 cycles of cyclic voltammetry (Fig. S14, ESI†). These facts confirm the high stability of our molecular catalyst–semiconductor-assembled photocathode.

The underlying mechanism of such a synergistic effect, the high selectivity for formate and its relation to the photoelectrochemical CO_2 reduction is discussed. Prior to this discussion, the interfacial energetics between the two components were analysed. The band-gap (Fig. S15, ESI†) and the flat-band potential (Fig. S16, ESI†) of $\text{Co}_3\text{O}_4/\text{CA}$ were estimated to be 1.87 V and 0.37 V, respectively.

Table 1 TON and TOF compared to results presented in the literature

Catalyst	Condition	Product	TON (TOF)	Ref.
$\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$	-0.6 V vs. NHE, 9 mW cm^{-2} Xe lamp, 8 h	HCOO^-	978.7 (122)	This work
$\text{Ru}(\text{bpy}-\text{H}_2\text{PO}_3)_2(\text{CO})_2\text{Cl}_2-\text{C}_3\text{N}_4$	400 W Hg lamp ($\lambda > 400$ nm), 20 h	HCOOH	1100 (55)	42
$\text{Ru}(\text{bpy})_3-(\text{CH}_2)_2-\text{Re}(\text{CH}_3\text{-bpy})(\text{CO})_3\text{Cl}$	Hg lamp ($\lambda > 500$ nm), 24 h	HCOOH	25 (1.04)	43
$\text{Ru}(\text{H}_4\text{P}_2\text{O}_6-\text{C}_2\text{H}_4\text{-bpy})(\text{CO})_2\text{Cl}_2\text{-mpg C}_3\text{N}_4$	450 W Hg lamp ($\lambda > 400$ nm), 20 h	HCOOH	~ 210 (10.5)	44
$\text{InP}/[\text{MCE2-A} + \text{MCE4}]$	AM 1.5G, 24 h	HCOO^-	> 17 (0.7)	39
$\text{Ru}(\text{dcbpy})/\text{N-Ta}_2\text{O}_5$	Xe lamp ($410 < \lambda < 750$ nm), 20 h	HCOO^-	90 (4.5)	45
NiO-RuRe complex	-1.2 V vs. Ag/Ag^+ , 300 W Xe lamp, 5 h	CO	32 (6.4)	40
Zn-TPP-Re complex	200 W Hg lamp, $\lambda > 375$ nm, 50 h	CO	12.8 (0.26)	46
$\text{Re}(\text{bpy})(\text{CO})_3\text{Cl}$	-1.25 V vs. SHE, 14 h	CO	300 (22)	47
$[\text{Co}(\text{CR})\text{Cl}_2](\text{ClO}_4)$	LED light strip ($\lambda > 460$ nm), 22 h	CO	268 (12.18)	48



Provided that the valence band of Co_3O_4 is 0.1 V more positive than the flat band potential, the valence band of Co_3O_4 was estimated to be 0.47 V. So the conduction band of $\text{Co}_3\text{O}_4/\text{CA}$ was estimated to be -1.40 V vs. NHE ($E_g = |E_{\text{CB}} - E_{\text{VB}}|$). In contrast, the energy difference between the HOMO and the LUMO for $\text{Ru}(\text{bpy})_2\text{dppz}$ is approximately 2.29 eV, as calculated using the intersection wavelength at 540.8 nm from its normalized UV-vis and fluorescence spectra in acetonitrile (Fig. S17a, ESI†). According to the Ru(III)/Ru(II) redox potential (Fig. S17b, ESI†), the LUMO energy level of $\text{Ru}(\text{bpy})_2\text{dppz}$ was calculated to be -0.71 V. $\text{Ru}(\text{bpy})_2\text{dppz}$'s LUMO energy level is then 0.69 V more positive than the conduction band of $\text{Co}_3\text{O}_4/\text{CA}$. Therefore, the photo-induced electrons on Co_3O_4 transfer to the LUMO of the molecular catalyst, as confirmed by various reports.^{24,32,35,40,41} Although the value of the conduction band and the redox potential of the molecular complex vary with the pH value of the electrolyte, the shift of the redox potential of a similar molecular complex $[\text{Ru}(\text{phen})_2(\text{ptpb}\beta)]^{2+}$ by a rate of -62 mV pH⁻¹⁴⁹ is in line with the typical -59 mV pH⁻¹ for semiconductors.

Electrochemical characterization data can also support such an electron transfer process. The onset potential of electrochemical CO_2 reduction using $\text{Ru}(\text{bpy})_2\text{dppz}$ was at -0.4 V vs. NHE, which is similar to that for photoelectrochemical CO_2 reduction using $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4$ (-0.45 V vs. NHE). Moreover, the cathodic current density of electrochemical CO_2 reduction on $\text{Ru}(\text{bpy})_2\text{dppz}$ (*ca.* 0.63 mA cm⁻²) was far smaller than that on $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4$ (*ca.* 15.0 mA cm⁻²). From *in situ* IR spectroelectrochemical spectra (Fig. 3a), the upward IR peaks at 1419 and 1446 cm⁻¹, assigned to the A1 mode of the C–C–H deformation bending vibration on $\text{bpy}^{\cdot-}$ and $\text{dppz}^{\cdot-}$,^{50,51} validate such a statement. A relatively strong broad upward peak at 1716 cm⁻¹ displays the same trend. This peak likely arises from the stretching of C=O from the as-formed formate.^{52,53} These facts suggest that the electrons are transferred from the excited Co_3O_4 to $\text{Ru}(\text{bpy})_2\text{dppz}$, and then take part in photoelectrochemical CO_2 reduction. As a result of such a photo-induced electron transfer to the LUMO of $\text{Ru}(\text{bpy})_2\text{dppz}$, CO_2 conversion/reduction occurs in the adsorptive substrate CA, the photocatalyst Co_3O_4 and the molecular catalyst $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$, exactly as illustrated in Scheme 1. Such an electron transfer process is the core of such an interface. Note that $\text{Ru}(\text{bpy})_2\text{dppz}$ in its activated form is capable of reducing CO_2 into formate with a two-electron process. The potential of its singly reduced form, $\text{Ru}(\text{bpy})_2(\text{dppz}^{\cdot-})$, is -0.95 V vs. NHE in MeCN.³⁰ Its doubly reduced state, $\text{Ru}(\text{bpy})(\text{bpy}^{\cdot-})(\text{dppz}^{\cdot-})$, is possible to be obtained by merging $\text{Ru}(\text{bpy})_2\text{dppz}$ and Co_3O_4 . This is because the electrons at the conduction band of Co_3O_4 bear only an energy of -1.40 V. In the presence of Co_3O_4 , the formation of a band-alignment interface based on $\text{Ru}(\text{bpy})_2\text{dppz}$ and Co_3O_4 allows the direct injection of photo-induced electrons to the LUMO of $\text{Ru}(\text{bpy})_2\text{dppz}$, which greatly suppresses the quenching of the molecular catalyst in the pure water phase.⁴⁹ Eventually, the photocatalytic activity of $\text{Ru}(\text{bpy})_2\text{dppz}$ is remarkably improved. This is further supported by the fact that the rate yield of formate, along with the TON/TOF values of the

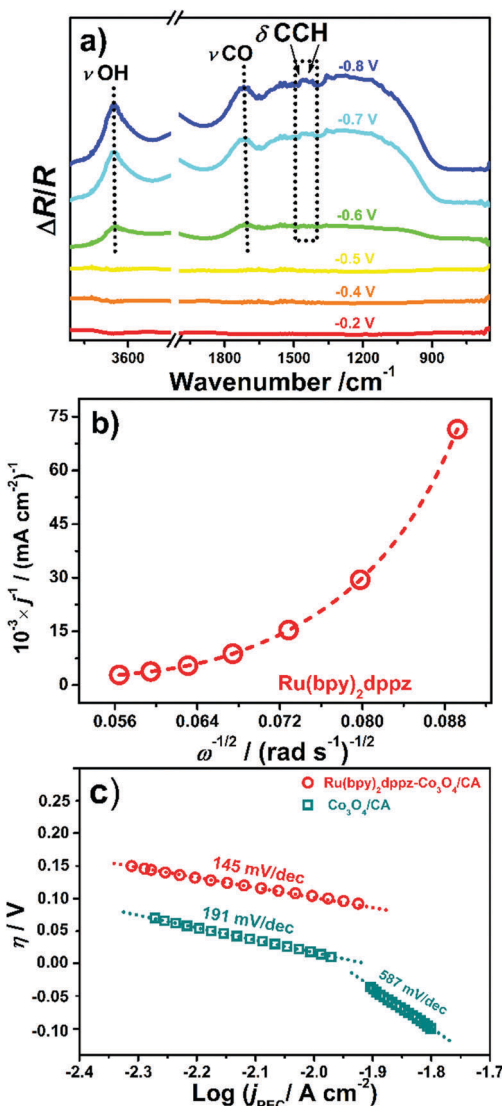


Fig. 3 (a) *In situ* photoelectrochemical IR spectra of $\text{Ru}(\text{bpy})_2\text{dppz}$ within the wavenumber ranges of 4000 – 3400 cm⁻¹ and 2000 – 650 cm⁻¹; (b) Koutecky–Levich fitting of the steady-state current on a $\text{Ru}(\text{bpy})_2\text{dppz}$ -coated glassy carbon electrode. (c) Tafel plots of $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ (circles) and $\text{Co}_3\text{O}_4/\text{CA}$ (squares) at a scan rate of 50 mV s⁻¹. All of these electrochemical measurements were conducted in CO_2 -saturated 0.1 M NaHCO_3 .

$\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4/\text{CA}$ after 8 h of reduction was almost 20 times higher than that on the $\text{Ru}(\text{bpy})_2\text{dppz}/\text{CA}$ (Fig. 2b). And on the basis of the Marcus–Gerischer model,^{54,55} the negative bias applied to the $\text{Ru}(\text{bpy})_2\text{dppz}-\text{Co}_3\text{O}_4$ system suppresses the back electrons transferred from $\text{Ru}(\text{bpy})_2\text{dppz}$ to Co_3O_4 . The utilization of the electrons is then greatly improved, and a high heterogeneous electron transfer rate of 2.94×10^{-3} cm s⁻¹ is reached.

It also noteworthy that the absence of the stretching vibrations of Ru–H, N–H, and C–H in the IR spectra indicates that the protons are not directly bonded to $\text{Ru}(\text{bpy})_2\text{dppz}$.^{56,57} A broad peak at approximately 3691 cm⁻¹ emerges from the featureless IR spectra when the applied potential is higher than -0.6 V vs. NHE (Fig. 3a).



This newly emerged upward peak is assigned to the stretching vibration of a non-hydrogen bonded hydroxyl. Water molecules are attracted and nearly dissociate from their bulky form due to the existence of localized electrons on the bpy ligand in Ru(bpy)₂dppz, thus causing the appearance of the peak at 3691 cm⁻¹. Hydrodynamic voltammograms using rotating glassy carbon disk electrodes were recorded using the electrode coated with Ru(bpy)₂dppz (Fig. S18a, ESI[†]) and Co₃O₄ (Fig. S18b, ESI[†]). Fig. 3b shows the related Koutecky–Levich plot for the electrode coated with Ru(bpy)₂dppz, indicating its non-linear features. This non-linearity is quite helpful for CO₂ conversion in that it indicates that CO₂ and the surrounding water fix into the outer sphere of Ru(bpy)₂dppz, apart from its diffusion and surface reaction with Ru(bpy)₂dppz under hydrodynamic conditions.⁵⁸ Taking a Langmuir–Hinshelwood bimolecular reaction mechanism as a model,⁵⁹ these results clearly confirm the enzyme-mimicking role of the molecular catalyst, Ru(bpy)₂dppz, for CO₂ conversion. In contrast, a linear reaction is observed on the electrode coated with Co₃O₄ (Fig. S18c, ESI[†]), indicating different electrokinetics between Co₃O₄ and Ru(bpy)₂dppz.

The rate-determining step of the CO₂ conversion was estimated through Tafel analysis (Fig. 3c). For Ru(bpy)₂dppz-Co₃O₄/CA, a Tafel slope of 145 mV dec⁻¹ is determined. Due to the porous feature of CA, it is slightly deviated from its classical value of 118 mV dec⁻¹.^{60,61} Therefore the rate-determining step of photoelectrochemical CO₂ conversion on Ru(bpy)₂dppz-Co₃O₄/CA is the single electron transfer step rather than the proton transfer step. This result is in agreement with our previous study regarding the photoelectrochemical CO₂ conversion on hierarchical Co₃O₄.⁴ This finding also supports the notion that the protonation of the reaction intermediate occurs rapidly in CO₂-saturated solution. Namely, faster electrode kinetics for CO₂ conversion were obtained with the help of Ru(bpy)₂dppz. In contrast, a Tafel slope of 191 mV dec⁻¹ is observed for Co₃O₄/CA, which originates from the porous structure of CA and is indicative of its slow photoelectrochemical kinetics in the absence of Ru(bpy)₂dppz. In addition, the activation energy of the CO₂ reduction in our system, defined as the energy required to activate CO₂ into its radicals,^{62–64} was calculated to be 1.047 kJ mol⁻¹ (see the ESI[†] for details). This value is presented for the first time in this study and therefore further density functional-based theoretical calculations are required to confirm this value.

Based on the above analysis, the activation of Ru(bpy)₂dppz is accomplished only by the photoelectrochemical process from the irradiated Co₃O₄/CA. Ru(bpy)₂dppz is actually “catalysed” by Co₃O₄/CA rather than merely an electron conductor. Therefore, the reduction performance is markedly enhanced with the assistance of the photoelectrocatalyst of Co₃O₄. Therefore, as schematically shown in Scheme 1, the possible pathways for CO₂ conversion on Ru(bpy)₂dppz-Co₃O₄/CA involve the transfer and mediation of electrons, protons, and the activated form of Ru(bpy)₂dppz.

Conclusions

In summary, an adsorption-enhanced Co₃O₄/Ru(bpy)₂dppz hybrid photoelectrocatalytic interface has been established.

The CO₂ adsorption is remarkably increased through the utilization of a high-surface-area and micropore-dominated carbon aerogel substrate. The epitaxial growth of high-index-facet Co₃O₄ along with the intrinsic highly conductive network of the carbon aerogel enables an elevated electron transfer rate with the assistance of a molecular catalyst of Ru(bpy)₂dppz. Eventually, such an interface bears a unique proton-coupled electron-transfer reactivity and rapid electron kinetics toward CO₂ reduction. Through the simultaneous activation of the bpy and dppz ligands of the Ru(II) molecular catalyst, the photoelectrochemical conversion of CO₂ to formate has been realized on such a photocathode with an onset potential as low as -0.45 V vs. NHE. At a potential of -0.6 V vs. NHE, the yield rate of formate reaches approximately 110 μmol cm⁻² h⁻¹, with a selectivity of 99.95% and a Faradaic efficiency of 86%. The mechanism of such a rapid photoelectrochemical electron transfer process is explained as the synergistic effect of the photoelectrochemical activation of bpy and dppz inside Ru(bpy)₂dppz as well as the remarkably promoted CO₂ adsorption on Co₃O₄/CA. Although future work on the effect of the morphology of Co₃O₄/CA (e.g., the size, facets, and shape of Co₃O₄ on CA) and the loading amount of Co₃O₄/CA (e.g., distribution and density) as well as the light density on the efficiency of the CO₂ conversion must be conducted, such an adsorption-enhanced molecular catalytic photoelectrocatalytic interface has potential for application in the production of useful chemicals from CO₂ carbon stock in the future.

Acknowledgements

The authors acknowledge Dr Sanjun Zhang (State Key Laboratory of Precision Spectroscopy, East China Normal University) for his assistance with spectral data analysis and discussion on the electron transfer mechanism. G. Z. acknowledges the financial support from the National Natural Science Foundation of China (NSFC) under projects No. 21537003 and 21477085. N. Y. acknowledges the financial support from the German Research Foundation (DFG) under project YA344/1-1.

Notes and references

- 1 N. J. Brown, C. A. Newell, S. Stanley, J. E. Chen, A. J. Perrin, K. Kajala and J. M. Hibberd, *Science*, 2011, **331**, 1436.
- 2 Q. Shen, Z. Chen, X. Huang, M. Liu and G. Zhao, *Environ. Sci. Technol.*, 2015, **49**, 5828.
- 3 J. Gu, A. Wuttig, J. W. Krizan, Y. A. Hu, Z. M. Detweiler, R. J. Cava and A. B. Bocarsly, *J. Phys. Chem. C*, 2013, **117**, 12415.
- 4 X. F. Huang, T. C. Cao, M. C. Liu and G. H. Zhao, *J. Phys. Chem. C*, 2013, **117**, 26432.
- 5 J.-W. Jang, S. Cho, G. Magesh, Y. J. Jang, J. Y. Kim, W. Y. Kim, J. K. Seo, S. Kim, K.-H. Lee and J. S. Lee, *Angew. Chem., Int. Ed.*, 2014, **53**, 5852.
- 6 J. L. White, M. F. Baruch, J. E. Pander, Y. Hu, I. C. Fortmeyer, J. E. Park, T. Zhang, K. Liao, J. Gu, Y. Yan,

T. W. Shaw, E. Abelev and A. B. Bocarsly, *Chem. Rev.*, 2015, **115**, 12888.

7 L. I. Bendavid and E. A. Carter, *J. Phys. Chem. C*, 2013, **117**, 26048.

8 Y. K. Chen, N. S. Lewis and C. X. Xiang, *Energy Environ. Sci.*, 2015, **8**, 3663.

9 N. Kornienko, Y. B. Zhao, C. S. Kley, C. H. Zhu, D. H. Kim, S. Lin, C. J. Chang, O. M. Yaghi and P. D. Yang, *J. Am. Chem. Soc.*, 2015, **137**, 14129.

10 Y. Fu, D. Sun, Y. Chen, R. Huang, Z. Ding, X. Fu and Z. Li, *Angew. Chem., Int. Ed.*, 2012, **51**, 3364.

11 H.-Q. Xu, J. H. Hu, D. K. Wang, Z. H. Li, Q. Zhang, Y. Luo, S.-H. Yu and H. L. Jiang, *J. Am. Chem. Soc.*, 2015, **137**, 13440.

12 P. Li, H. Hu, J. Xu, H. Jing, H. Peng, J. Lu, C. Wu and S. Ai, *Appl. Catal., B*, 2014, **147**, 912.

13 P. Li, X. Sui, J. Xu, H. Jing, C. Wu, H. Peng, J. Lu and H. Yin, *Chem. Eng. J.*, 2014, **247**, 25.

14 E. E. Barton, D. M. Rampulla and A. Bocarsly, *J. Am. Chem. Soc.*, 2008, **130**, 6342.

15 A. J. Morris, G. J. Meyer and E. Fujita, *Acc. Chem. Res.*, 2009, **42**, 1983.

16 J. Agarwal, E. Fujita, H. F. Schaefer and J. T. Muckerman, *J. Am. Chem. Soc.*, 2012, **134**, 5180.

17 Z. Chen, C. Chen, D. R. Weinberg, P. Kang, J. J. Concepcion, D. P. Harrison, M. S. Brookhart and T. J. Meyer, *Chem. Commun.*, 2011, **47**, 12607.

18 B. Kumar, J. M. Smieja and C. P. Kubiak, *J. Phys. Chem. C*, 2010, **114**, 14220.

19 Y. Matsubara, S. E. Hightower, J. Z. Chen, D. C. Grills, D. E. Polyansky, J. T. Muckerman, K. Tanaka and E. Fujita, *Chem. Commun.*, 2014, **50**, 728.

20 Y. Hayashi, S. Kita, B. S. Brunschwig and E. Fujita, *J. Am. Chem. Soc.*, 2003, **125**, 11976.

21 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 and C. J. Chang, *Science*, 2015, **349**, 1208.

22 Y. Tamaki, T. Morimoto, K. Koike and O. Ishitani, *Proc. Natl. Acad. Sci. U. S. A.*, 2012, **109**, 15673.

23 K. Sekizawa, K. Maeda, K. Domen, K. Koike and O. Ishitani, *J. Am. Chem. Soc.*, 2013, **135**, 4596.

24 T. Arai, S. Sato, K. Uemura, T. Morikawa, T. Kajino and T. Motohiro, *Chem. Commun.*, 2010, **46**, 6944.

25 J. D. Hong, W. Zhang, J. Ren and R. Xu, *Anal. Methods*, 2013, **5**, 1086.

26 T. Yamamoto, D. A. Tryk, K. Hashimoto, A. Fujishima and M. Okawa, *J. Electrochem. Soc.*, 2000, **147**, 3393.

27 S. Perez-Rodriguez, N. Rillo, M. J. Lazaro and E. Pastor, *Appl. Catal., B*, 2015, **163**, 83.

28 M. F. Wu, Y. N. Jin, G. H. Zhao, M. F. Li and D. M. Li, *Environ. Sci. Technol.*, 2010, **44**, 1780.

29 Y. N. Zhang, Y. F. Jin, X. F. Huang, H. J. Shi, G. H. Zhao and H. Y. Zhao, *Electrochim. Acta*, 2014, **130**, 194.

30 J. Fees, W. Kaim, M. Moschersch, W. Mattheis, J. Klima, M. Kerjelk and S. Zalls, *Inorg. Chem.*, 1993, **31**, 166.

31 J. L. Qiao, Y. Y. Liu, F. Hong and J. J. Zhang, *Chem. Soc. Rev.*, 2014, **43**, 631.

32 M. Schreier, P. Gao, M. T. Mayer, J. S. Luo, T. Moehl, M. K. Nazeeruddin, S. D. Tilley and M. Gratzel, *Energy Environ. Sci.*, 2015, **8**, 855.

33 J. G. Velasco, *Electroanalysis*, 1997, **9**, 880.

34 H. Zhong, K. Fuji, Y. Nakano and F. M. Jin, *J. Phys. Chem. C*, 2015, **119**, 55.

35 T. Arai, S. Tajima, S. Sato, K. Uemura, T. Morikawa and T. Kajino, *Chem. Commun.*, 2011, **47**, 12664.

36 X. Q. An, K. F. Li and J. W. Tang, *ChemSusChem*, 2014, **7**, 1086.

37 L. H. Zhang, D. Zhu, G. M. Nathanson and R. J. Hamers, *Angew. Chem., Int. Ed.*, 2014, **53**, 9746.

38 M. Garcia, M. J. Aguirre, G. Canzi, C. P. Kubiak, M. Ohlbaum and M. Issacs, *Electrochim. Acta*, 2014, **115**, 146.

39 S. Sato, T. Arai, T. Morikawa, K. Uemura, T. M. Suzuki, H. Tanaka and T. Kajino, *J. Am. Chem. Soc.*, 2011, **133**, 15240.

40 G. Sahara, R. Abe, M. Higashi, T. Morikawa, K. Maeda, Y. Ueda and O. Ishitani, *Chem. Commun.*, 2015, **51**, 10722.

41 S. Sato, T. Morikawa, S. Saeki, T. Kajino and T. Motohiro, *Angew. Chem., Int. Ed.*, 2010, **49**, 5101.

42 R. Kuriki, K. Sekizawa, O. Ishitani and K. Maeda, *Angew. Chem., Int. Ed.*, 2015, **54**, 2406.

43 A. Nakada, K. Koike, T. Nakashima, T. Morimoto and O. Ishitani, *Inorg. Chem.*, 2015, **54**, 1800.

44 K. Maeda, K. Sekizawa and O. Ishitani, *Chem. Commun.*, 2013, **49**, 10127.

45 S. Sato, T. Morikawa, S. Saeki, T. Kajino and T. Motohiro, *Angew. Chem., Int. Ed.*, 2010, **49**, 5101.

46 C. Matlachowski, B. Braun, S. Tschierlei and M. Schwalbe, *Inorg. Chem.*, 2015, **54**, 10351.

47 J. Hawecker, J. M. Lehn and R. Ziessel, *J. Chem. Soc., Chem. Commun.*, 1984, 328.

48 L. J. Chen, Z. G. Guo, X. G. Wei, C. Gallenkamp, J. Bonin, E. A. Mallart, K. C. Lau, T. C. Lau and M. Robert, *J. Am. Chem. Soc.*, 2015, **137**, 10918.

49 D. J. Boston, Y. M. Franco Pachon, R. O. Lezna, N. R. de Tacconi and F. M. MacDonnell, *Inorg. Chem.*, 2014, **53**, 6544.

50 P. K. Mallick, G. D. Danzer, D. P. Strommen and J. R. Kincaid, *J. Phys. Chem.*, 1988, **92**, 5628.

51 K. M. Omberg, J. R. Schoonover, J. A. Treadway, R. M. Leisure, R. B. Dyer and T. J. Meyer, *J. Am. Chem. Soc.*, 1997, **119**, 7013.

52 K. Jiang, K. Xu, S. Zou and W.-B. Cai, *J. Am. Chem. Soc.*, 2014, **136**, 4861.

53 J. Y. Wang, H. X. Zhang, K. Jiang and W. B. Cai, *J. Am. Chem. Soc.*, 2011, **133**, 14876.

54 B. H. Farnum, Z. A. Morseth, M. K. Brennaman, J. M. Papanikolas and T. J. Meyer, *J. Am. Chem. Soc.*, 2014, **136**, 15869.

55 N. S. Lewis, *J. Phys. Chem. B*, 1998, **102**, 4843.

56 Q. Wang, X. Wang and L. Andrews, *J. Phys. Chem. A*, 2011, **115**, 12194.

57 K. Toyohara, K. Tsuge and K. Tanaka, *Organometallics*, 1995, **14**, 5099.

58 R. Vargas, C. Borras, J. Mostany and B. R. Scharifker, *Electrochim. Acta*, 2012, **80**, 326.



59 Y. Kiya, O. Hatozaki, N. Oyama and H. D. Abruna, *J. Phys. Chem. C*, 2007, **111**, 13129.

60 Y. H. Chen and M. W. Kanan, *J. Am. Chem. Soc.*, 2012, **134**, 1986.

61 J. N. Soderberg, A. C. Co, A. H. C. Sirk and V. I. Birss, *J. Phys. Chem. B*, 2006, **110**, 10401.

62 S. Antonello, M. Musumeci, D. D. M. Wayner and F. Maran, *J. Am. Chem. Soc.*, 1997, **119**, 9541.

63 S. Antonello and F. Maran, *J. Am. Chem. Soc.*, 1999, **121**, 9668.

64 C. Ji, M. Ahmida, M. Chahma and A. Houmam, *J. Am. Chem. Soc.*, 2006, **128**, 15423.

