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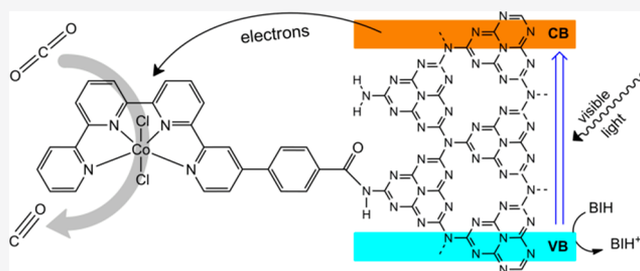
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Efficient Visible-Light-Driven CO₂ Reduction by a Cobalt Molecular Catalyst Covalently Linked to Mesoporous Carbon Nitride

Bing Ma, Gui Chen,* Claire Fave, Lingjing Chen, Ryo Kuriki, Kazuhiko Maeda, Osamu Ishitani,* Tai-Chu Lau,* Julien Bonin, and Marc Robert*

ABSTRACT: Achieving visible-light-driven carbon dioxide reduction with high selectivity control and durability while using only earth abundant elements requires new strategies. Hybrid catalytic material was prepared upon covalent grafting a Co–quaterpyridine molecular complex to semiconductive mesoporous graphitic carbon nitride (mpg-C₃N₄) through an amide linkage. The molecular material was characterized by various spectroscopic techniques, including XPS, IR, and impedance spectroscopy. It proved to be a selective catalyst for CO production in acetonitrile using a solar simulator with a high 98% selectivity, while being remarkably robust since no degradation was observed after 4 days of irradiation (ca. 500 catalytic cycles). This unique combination of a selective molecular catalyst with a simple and robust semiconductive material opens new pathways for CO₂ catalytic light-driven reduction.



INTRODUCTION

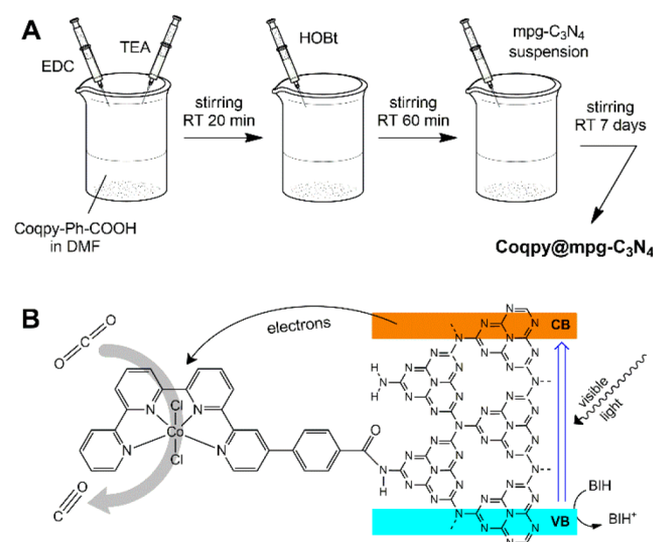
CO₂ may be used as a renewable feedstock for making fuels or commodity chemicals directly from sunlight energy. But achieving photochemically driven conversion of CO₂ remains a grand challenge, especially if one considers that only abundant materials should be used in view of future large scale applications. One approach consists of associating a molecular catalyst, typically a metal complex, to a robust semiconductive material that will efficiently absorb visible-light photons and transfer energy under the form of electrons to the catalyst. In this connection, carbon nitride (C₃N₄) is a synthetic polymer which received considerable attention in the past 15 years, not only as an earth-abundant visible-light photocatalyst^{1–3} for water splitting,^{4–6} CO₂ reduction,⁷ and inclusion in fuel cells^{8,9} but also for applications in organic synthesis,¹⁰ water depollution,¹¹ or sensors.¹² This medium band gap (ca. 2.8 eV for the mesoporous phase)^{13,14} semiconductor possesses several allotropic phases, including α -, β -, cubic, quasi-cubic, and graphitic phases, the latter (g-C₃N₄) being the most stable.¹⁵ g-C₃N₄ is a graphite-like layered structure composed of conical nitrogen bridges and triazine or heptazine rings. It has high chemical stability, a good electron-withdrawing structure, and visible light response activity.

Since the pioneering work of Lehn on Re and Ru complexes,^{16–18} the photochemical reduction of CO₂ with a molecular catalyst has proven to be a powerful way to target a specific product,¹⁹ from two-electron-reduced products (CO, formate) to fuels such as methane (eight-electron reduc-

tion)^{20–22} and to reach a high selectivity thanks to the control of the chelating environment and the steric and electronic effects of the ligands. Most of these molecular catalysts, including those based on earth-abundant transition metals such as Fe or Mn, mainly lead to the formation of carbon monoxide or formate.^{23,24} However, molecular systems could suffer from progressive structural degradation due to photochemical instability or secondary reactions from/with byproducts. Bimolecular reactions (between the catalyst, the sensitizer, and the electron donor/acceptor) may additionally severely limit catalytic performances of homogeneous systems. To date, only a few semiconductor solid catalysts (e.g., metal oxides such as TiO₂) have been reported for visible-light-driven catalysis,^{23,25,26} and most of them suffer from poor catalytic selectivity, mainly because of the competition with hydrogen evolution. For all of these reasons, an attractive option is to combine an efficient and selective molecular catalyst with a solid-state sensitizer material such as carbon nitride. It may be achieved by adsorption or by covalent linking, so as to integrate the main components for the photochemically induced CO₂ conversion.^{27,28} Several recent studies have reported the combination of various types of C₃N₄ materials

with a metal complex,²⁹ for example, based on Ru,^{30–33} Ni,³⁴ or Co^{14,35,36} for photochemical catalysis. To date, the best catalytic performance with mesoporous mpg-C₃N₄ for CO₂ reduction to formate was obtained with a *trans*-(Cl)-[Ru[4,4'-(CH₂PO₃H₂)₂-2,2'-bipyridine](CO)₂Cl₂] (RuP)³⁷ immobilized complex in a mixed DMA:TEOA (4:1, v:v) solvent. The turnover number (TON_{formate}) reached 1061 after 20 h irradiation, with a selectivity of ca. 80%. With regard to CO₂ reduction to CO, the best performance for systems only containing abundant elements was recently obtained by [Fe(2,2':6',2'':6'',2''-quaterpyridine qpy)(H₂O)₂]²⁺ (Fe(qpy))³⁸ as a homogeneous catalyst in a mixed ACN:TEOA (4:1, v:v) solvent, TON_{CO} reaching 155 along with a very high selectivity (97%) and a high apparent quantum yield (ca. 4.2%). Poisoning of the iron catalyst with CO, however, hampers the long-term activity. The cobalt analogue, Coqpy, has also been shown to act as a selective and efficient homogeneous catalyst for the CO₂-to-CO photochemical reduction when sensitized by the tris(bipyridine)ruthenium(II) (Ru(bpy)₃)²⁺ complex, with longer stability than Fe(qpy).³⁹ To have a system incorporating only earth-abundant elements and to also enhance electronic interaction between the sensitizer and the catalytic sites, we have constructed a new molecular-material hybrid upon covalently attaching the metal complex Coqpy-Ph-COOH to the semiconductive polymer (mpg-C₃N₄) through an amide linkage (Scheme 1). This

Scheme 1. (A) Preparation of the Coqpy@mpg-C₃N₄ Hybrid Assembly and (B) Illustration of the Visible-Light-Driven CO₂ to CO Reduction Process^a



^aBIH is the sacrificial electron donor (see text). The Coqpy-Ph-COOH compound is abbreviated Coqpy in all hybrid names.

robust association led to excellent long-term stability upon several days of irradiation, while efficient electron transfer to the cobalt catalyst allowed for efficient and selective CO₂ reduction to CO.

RESULTS AND DISCUSSION

Molecular Materials Hybrid Assemblies Preparation.

Mesoporous graphitic carbon nitride (mpg-C₃N₄)⁴⁰ and graphitic nanosheets of graphitic carbon nitride (nsg-C₃N₄)³² were prepared according to previously reported methods. Coqpy@

mpg-C₃N₄ was prepared (Scheme 1A) by the addition of 1-ethyl-3-(3-(dimethylaminopropyl)carbodiimide (EDC, 3.05 mg, 16 μmol) and triethylamine (TEA, 2.2 μL, 16 μmol) to a stirring suspension of Coqpy-Ph-COOH (4.48 mg, 8 μmol; see Figures S1 and S2 for synthesis and characterization) in DMF (5 mL). The mixture was stirred for 20 min at room temperature, and then a 1-hydroxybenzotriazole (HOBt, 2.14 mg, 16 μmol) solution in 0.5 mL of DMF was added and kept stirring for 1 h. Next, mpg-C₃N₄ (80 mg) suspension in 2 mL of DMF was introduced into the mixture. After stirring for 7 days, the product was isolated by filtration on a nylon membrane (0.1 μm). Excess Coqpy-Ph-COOH and other impurities were removed through washing cycles, including sonication, filtration, and resuspension of the solid in DMF (200 mL). UV-vis spectroscopy was used to check the filtrate to ensure that no Coqpy-Ph-COOH remained in the final washing. After filtration, Coqpy@mpg-C₃N₄ was washed with ultrapure water several times, following the above procedure. Finally, the precipitate was dried under vacuum to yield the Coqpy@mpg-C₃N₄ hybrid. Coqpy@nsg-C₃N₄ was similarly synthesized by using nsg-C₃N₄ instead of mpg-C₃N₄. The adsorbed sample, noted Coqpy-mpg-C₃N₄, was prepared by dissolving Coqpy-Ph-COOH (1 mg) into a mpg-C₃N₄ (25 mg) suspension in 25 mL of DMF. After stirring overnight, the final solid was obtained by a similar post-treatment including filtration and washing cycles, but no sonication. In a final step, the precipitate was dried under vacuum to yield the Coqpy-mpg-C₃N₄, for which the Co complex is physically adsorbed onto the carbon nitride material. For the mixed sample, noted Coqpy-Ph-COOH/mpg-C₃N₄, mpg-C₃N₄ was dispersed into acetonitrile and sonicated to obtain a uniform suspension. Then a known amount of Coqpy-Ph-COOH solution was added into the suspension, and the photocatalysis experiments started immediately.

Spectroscopic Characterizations. Nitrogen adsorption/desorption experiment: The pore volume of ca. 0.50 cm³ g⁻¹ and the specific surface area of 111 m² g⁻¹ of synthesized mpg-C₃N₄ were measured by a nitrogen adsorption/desorption isothermal test at 77 K. The morphology was maintained through the grafting procedure and after photoreaction, with a final pore volume of 0.522 cm³ g⁻¹ and a specific surface area of 119 m² g⁻¹ (see Figure S3 and Table S1).

Infrared Spectra. IR spectra of mpg-C₃N₄, Coqpy-Ph-COOH, and the hybrid assembly are presented in Figure S4. Typical features of mpg-C₃N₄ can be observed for both bare material and hybrid assembly, including N-H stretching in the 3200 cm⁻¹ region with a -NH₂ specific stretching around 3280 cm⁻¹, an interchain NH...O-C bonding around 3170 cm⁻¹, triazine features around 810 cm⁻¹, and a broad band spanning the 1100–1600 cm⁻¹ region that can be assigned to in-plane C-N stretching and bending vibrations of the graphitic layers.⁴¹ In the hybrid spectrum, despite weak signals due to low catalyst loading, we still observed characteristic peaks at 1682 and 1558 cm⁻¹ attributed to C=O stretching of the amide link and to in-plane N-H deformation coupled with C-N stretching.

X-ray Photoelectron Spectroscopy (XPS). As shown in Figures S5 and S6 as well as Table S2, typical XPS signatures originating from C 1s, N 1s, and O 1s atoms within the hybrid are observed at binding energies of about 287, 400, and 532 eV.⁴¹ The C 1s spectrum is composed of three components: a major one at 288.9 eV corresponding to NC=N (carbon nitride), a second one at 284.9 eV from C-C/C-H, and a

third one at 286.3 eV from C–N attributed to the coupling of C_3N_4 and Coqpy-Ph-COOH units. The O 1s peak centered at 532.4 eV is attributed to the amide C=O. The N 1s peaks centered at 398.8 and 399.9 eV belong to C=N and N(C_3)₃ of the triazine motif, respectively. A weak N 1s peak centered at 404.8 eV is also observed for the nitride substrate. The peak centered at 401.1 eV is consistent with C–N–H from covalent amide linkage. On the other hand, a weak Co 2p signature at ca. 782 eV can be observed, proving the presence of cobalt inside the hybrid material.

Inductively Coupled Plasma–Optical Emission Spectroscopy (ICP-OES). Amounts of cobalt in the different hybrid materials (covalently linked or adsorbed) are given in Table 1. The equivalent concentration of Coqpy was also

Table 1. Amount of Cobalt in Hybrid Materials Determined by ICP and Equivalent Coqpy Concentration in Solution

hybrid	Co (wt %)	Coqpy ($\mu\text{mol g}^{-1}$)	[Coqpy] ^a (μM)
Coqpy@mpg- C_3N_4	0.00876	1.48	3.0
Coqpy/mpg- C_3N_4	0.00744	1.26	2.5
Coqpy@nsg- C_3N_4	0.02670	4.53	9.0

^aCalculated for 6 mg of hybrid material suspended in a 3 mL solution.

calculated and systematically used for TON calculation. From these measurements, it appears that the Coqpy loading is higher when nsg- C_3N_4 is used, most probably thanks to more favorable structural properties. Interestingly, equivalent Coqpy concentration is very similar to typical conditions used in homogeneous catalysis.^{42,43}

Cyclic Voltammetry (CV). The CV of Coqpy-Ph-COOH is shown in Figure S7. Under argon and upon reductive scan, a reversible wave at -0.7 V vs SCE is observed and is attributed to the Co^{II} – Co^I . Further reduction first leads to quasi-reversible waves at -1.1 V vs SCE and -1.4 V vs SCE due to combined metal and ligand-based reduction, in line with previous reports.^{39,44} Under a CO_2 atmosphere, an increase is observed on the latter wave, showing that Coqpy-Ph-COOH possesses the ability to catalytically reduce CO_2 . In the presence of a weak Brønsted acid (phenol), the catalytic current further drastically increases. The onset potential of the catalytic wave is positive to the position of the conduction band (CB) of mpg- C_3N_4 , located at ca. -1.35 V vs SCE, as reported elsewhere,¹ indicating that the illuminated semiconductive material can reduce the cobalt molecular catalyst.

Photoinduced Electron Transfer. Irradiation of mpg- C_3N_4 with visible light ($\lambda > 400$ nm) allows electron–hole separation with the promotion of an electron in the conduction band of the semiconductor. The efficiency of the electron transfer (ET) to Coqpy-Ph-COOH is a key parameter for the catalytic process which can be evaluated by mpg- C_3N_4 emission quenching experiments in the presence of either BIH or Coqpy-Ph-COOH (Figures S8 and S9). The steady-state emission spectrum of suspended mpg- C_3N_4 in acetonitrile shows two peaks of similar intensity at ca. 490 and 540 nm, in accordance with previous studies.⁴⁵ In the presence of BIH or Coqpy-Ph-COOH, we observe a moderate but linear emission decay with quencher concentration. Electron transfer rate constant determination through Stern–Volmer (S–V) analysis requires the excited-state lifetime of mpg- C_3N_4 . Several studies reported a very short lifetime for C_3N_4 excited states.^{4,46,47} In another recent paper the excited states and electron transfer dynamics of g- and mpg- C_3N_4 hybridized with

a Ru mononuclear complex were investigated.⁴⁵ It was shown that the emissive excited states of mpg- C_3N_4 decayed with a three-component (from 0.7 to 12 ns) kinetics, but these excited states were not quenched neither by a hole scavenger nor by an electron acceptor. It was further concluded that emissive excited states are mainly located in the bulk region of the carbon nitride and thus insensitive to the presence of quencher, whereas less to nonemissive excited states were mainly surface states responsible for ultrafast electron transfer (a few picoseconds).

In our measurements, we observed that the two emission peaks were quenched at different rates both with BIH and Coqpy-Ph-COOH, which could be explained if these peaks correspond to slightly different excited states. However, the Stern–Volmer constant (Table S3) is much larger (by a factor >3900) in the case of the cobalt complex and largely counterbalances the smaller concentration of the catalyst as compared to the sacrificial donor (by a factor ca. 16) in the photocatalytic experiments. In the photocatalytic process, mpg- C_3N_4 is thus oxidized. Electron transfer is efficient and most probably proceeds from mpg- C_3N_4 to the cobalt(II) catalyst, with BIH sacrificially transferring electrons back to the material, even if we could not determine the exact rate constant. It must be noted that we cannot probe the second electron transfer to the as generated $Co(I)$ species, but we assume that the process is similar to the first one. We also measured the photocurrent response of mpg- C_3N_4 , Coqpy/mpg- C_3N_4 (obtained by simple mixing of Coqpy-Ph-COOH, and mpg- C_3N_4 as previously explained) and Coqpy@mpg- C_3N_4 (Figure 1). The materials were deposited at a FTO

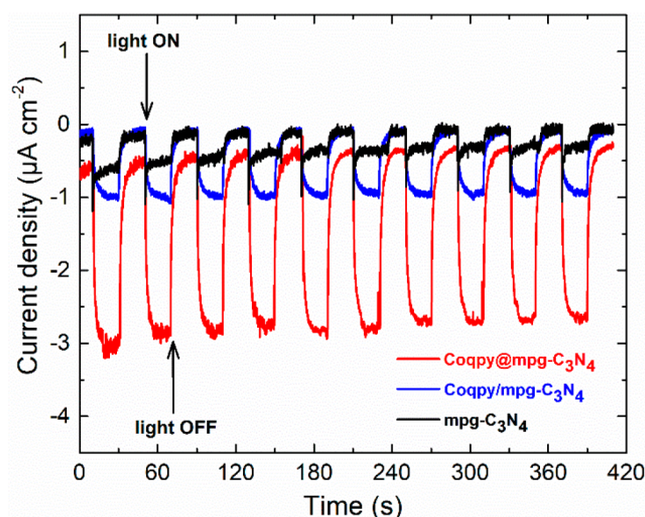


Figure 1. Transient (light on/off) photocurrent responses of mpg- C_3N_4 (black), Coqpy/mpg- C_3N_4 (blue), and Coqpy@mpg- C_3N_4 (red) hybrid ink dropped onto a FTO electrode connected to a platinum electrode (no bias, 0.1 M Na_2SO_4 solution).

electrode which was connected to a platinum electrode in a 0.1 M Na_2SO_4 solution without any bias (see the Experimental Section for details). The current responses upon light illumination of mpg- C_3N_4 and Coqpy/mpg- C_3N_4 were negligible, whereas the covalently linked Coqpy@mpg- C_3N_4 assembly showed a large, quickly rising, and stable current response, only with a slight attenuation being visible after ten cycles. This confirms that the covalent link between the two

moieties greatly enhanced electronic communication, as expected.

Upon running 10 consecutive illumination cycles, it was also observed that the current response was maintained with no significant attenuation or time delay. The electronic impedance spectroscopy (EIS) responses of mpg-C₃N₄, Coqpy/mpg-C₃N₄, and Coqpy@mpg-C₃N₄ were further measured (Figure 2), showing that the charge-transfer resistance of Coqpy@

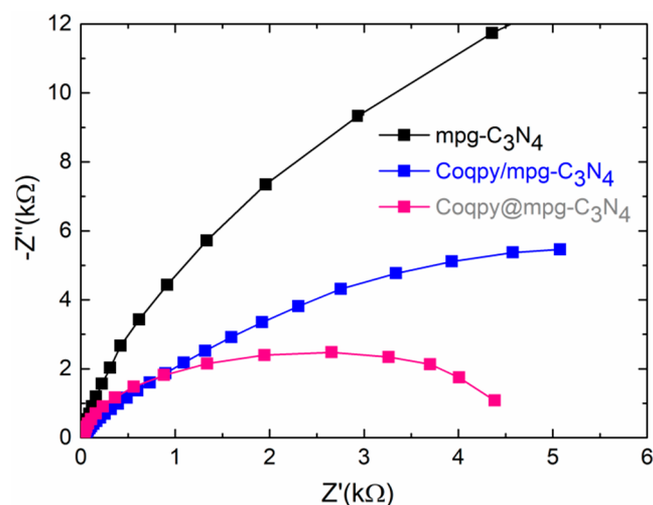


Figure 2. EIS (electrochemical impedance spectroscopy) responses of mpg-C₃N₄ (black), Coqpy/mpg-C₃N₄ (blue), and Coqpy@mpg-C₃N₄ (pink) hybrid ink dropped onto a FTO electrode in the dark (0.1 M Na₂SO₄ solution).

mpg-C₃N₄ is smaller than Coqpy/mpg-C₃N₄ and mpg-C₃N₄. The hybrid Coqpy@mpg-C₃N₄ thus possesses the lowest impedance, illustrating that the covalent linking significantly enhances conductivity of the material.

Visible-Light-Driven CO₂ Reduction Catalysis. As shown in Figure 3, visible light illumination of the catalyst suspension furnishes CO with very high selectivity (97%) and long-term durability since the catalytic material can evolve the

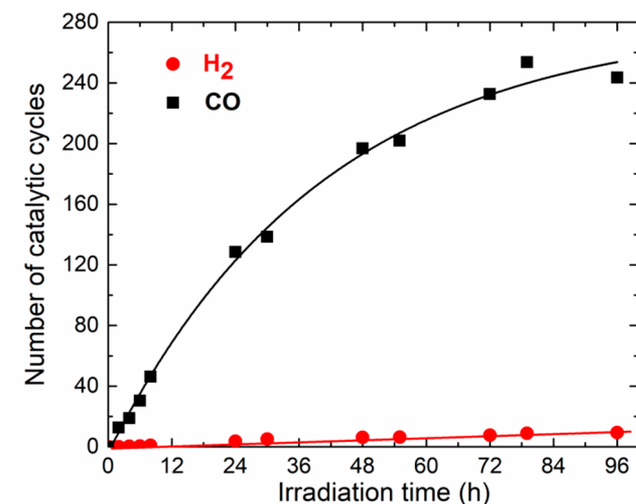


Figure 3. Generation of CO (black squares) and H₂ (red circles) over 4 days upon visible-light irradiation ($\lambda > 400$ nm) of a CO₂-saturated ACN solution containing 6 mg of Coqpy@mpg-C₃N₄, 0.05 M BIH (sacrificial donor), and 0.03 M PhOH (proton source).

gas product over a period of about 4 days, reaching a TON for CO of 254. The origin of carbon was asserted by running isotope-labeled experiments with ¹³CO₂ giving ¹³CO as the product (see MS spectrum, Figure S10). Table 2 reports data

Table 2. Visible-Light-Driven CO₂ Reduction with Coqpy Catalyst Covalently Attached to Mpg-C₃N₄

entry	catalyst ^a	product (μmol)			CO selectivity (%)
		H ₂	CO	TON _{CO} ^b	
1	Coqpy@mpg-C ₃ N ₄	0.06	1.15	128	98
2	Coqpy-Ph-COOH/mpg-C ₃ N ₄	0.04	0.31	37	88
3	Coqpy-mpg-C ₃ N ₄	0.03	0.15	21	83
4	Coqpy@mpg-C ₃ N ₄ ^c	0.013	0	0	0
5	Coqpy@mpg-C ₃ N ₄ ^d	0.001	0	0	0
6	mpg-C ₃ N ₄	0.012	0.017		59
7	Coqpy/Al ₂ O ₃	0.004	0.062	8	94
8	CoCl ₂ /mpg-C ₃ N ₄	0.013	0.018	1.7	58
9	Coqpy@nsg-C ₃ N ₄	0.018	0.81	27	98
10	Coqpy/mpg-C ₃ N ₄	0.035	0.22	26	86
11	Coqpy-mpg-C ₃ N ₄	0.09	0.52	58	85

^aReaction conditions: 6 mg of catalyst (3 μM Coqpy) in a CO₂-saturated ACN solution containing 0.05 M BIH and 0.03 M PhOH. A 6 mL quartz cell with a septum was used as the reaction vessel, and an AM1.5G solar simulator equipped with a 400 nm long-pass filter was used as irradiation source. Reaction time: 24 h. Amount of Coqpy in catalyst: covalently linked at mpg-C₃N₄ (Coqpy@mpg-C₃N₄), 3 μM ; mixed with carbon nitride (Coqpy/mpg-C₃N₄), 3 μM ; adsorbed at the carbon nitride surface (Coqpy-mpg-C₃N₄), 2.5 μM ; mixed with Al₂O₃ (Coqpy/Al₂O₃), 3 μM ; covalently linked at nsg-C₃N₄ (Coqpy@nsg-C₃N₄), 9 μM . ^bRelative to the amount of Coqpy or CoCl₂. ^cUnder an argon atmosphere. ^dIn the dark (results obtained with a Hg lamp as irradiation source are given in Table S4).

obtained after 24 h irradiation. TON_{CO} of 128 can be converted to 8 $\mu\text{mol g}^{-1} \text{h}^{-1}$ to facilitate comparison with other catalytic systems. No formate was identified from the liquid phase. Complete blank experiments are also given in Table 2, showing that all components of the catalytic system are needed. Experiments performed with cobalt chloride salt mixed with carbon nitride or Coqpy mixed Al₂O₃ suspension only furnishes a very low amount of reduction products (entries 7 and 8, respectively, Table 2). Moreover, simple mixing of the cobalt molecular complex with mpg-C₃N₄ considerably decreases the amount of CO produced (entries 2 and 3, Table 2), and concomitantly the selectivity for CO₂ reduction significantly diminishes.

These observations clearly indicate that covalent grafting of the molecular catalyst enhances catalysis. Recently, polymerized cobalt phthalocyanine mixed with mesoporous carbon nitride was shown to evolve CO with ca. 90 TON and a selectivity in the range of 80%.¹⁴ In another approach, Co²⁺ single sites embedded into C₃N₄ material led to CO production with a selectivity inferior to 80% and ca. 200 TON.⁴⁸ In the former case, large loading will block incoming light to reach the semiconductive material while in the latter case it will drive to the formation of CoO, leading to decrease photocatalytic activity in both cases. The covalent attachment of an efficient and selective molecular catalyst to the polymeric material thus appears as a promising approach to catalyze CO₂ reduction, allowing for better control of the catalyst structure

on one hand and better selectivity and activity on the other hand.

To further investigate the stability of the hybrid assembly, we re-employed the same hybrid for four successive cycles. Each cycle corresponded to 24 h visible-light irradiation of a standard Coqpy@mpg-C₃N₄/BIH/PhOH ACN solution. After each cycle, the hybrid was washed with ACN with sonication and then centrifuged. After drying, the hybrid was dispersed again into a fresh ACN containing BIH and PhOH and then saturated with CO₂ for 30 min to start a new photocatalytic cycle. Results (Figure 4) show a remarkable stability of CO

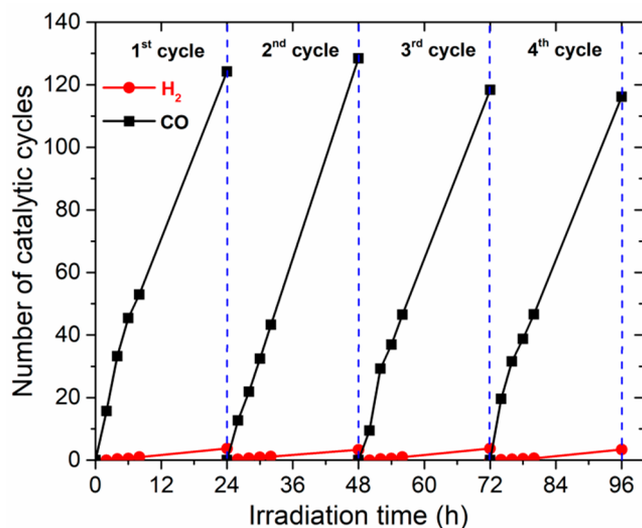


Figure 4. Generation of CO (black squares) and H₂ (red circles) during four consecutive 24 h irradiation cycles using the same hybrid material (see text).

production in each cycle, asserting the absence of hybrid degradation during irradiation. It also illustrates the ability to easily recycle the hybrid material for further use. A total number of ca. 500 catalytic cycles was reached after 96 h. Taking into account that CO₂-to-CO conversion necessitates two electrons, the apparent quantum yield for CO formation is $\Phi = 0.25\%$ (see the Supporting Information).

X-ray diffraction spectra of mpg-C₃N₄, nsg-C₃N₄, Coqpy-Ph-COOH, and Coqpy@mpg-C₃N₄ before and after irradiation are shown in Figure S12. Coqpy@mpg-C₃N₄ spectra before and after irradiation show mixed diffraction peaks including characteristic ones of the complex at 7.1°, 10.1°, 12.5°, 16.1°, 21.6°, 23.9°, 27.3°, and 29.9° as well as the peak of C₃N₄ at 27.5°. The strong diffraction peak due to the stacking of conjugated aromatic rings that can be assigned to the (002) plane of the heptazine-based g-C₃N₄. In addition, the comparison of hybrids patterns with one typical porous material, namely an aluminum silicate zeolite (JCPDS card 00-023-0214), shows common features. In addition, no characteristic diffraction peaks corresponding to cobalt oxide were observed. XPS data in Figure S6 also confirmed the presence of cobalt in Coqpy@mpg-C₃N₄ before and after photocatalysis as well as in the adsorbed Coqpy-mpg-C₃N₄. Absorption spectra of PhOH, BIH, qpy-Ph-COOH ligand, and Coqpy-Ph-COOH complex as well as photochemical mixture before and after irradiation are shown in Figure S13. The UV-vis absorption spectrum conserves the same spectral trends compared to before irradiation, albeit with a lower intensity.

No characteristic absorption peak of the ligand could be observed. All these results further support stability of the catalytic material upon long time irradiation.

CONCLUSION

The hybrid Coqpy@mpg-C₃N₄ material is among the first examples comprising a molecular catalyst covalently anchored to carbon nitride for CO₂RR and only containing abundant elements. Among such rare examples, Coqpy@mpg-C₃N₄ is highly selective and active toward CO production, showing in addition long-term stability. It thus combines the selectivity of molecular catalysts to the excellent stability of solid materials. The possibility to independently tune the structure of the metal complex and the structure of the semiconductive material (band gap and conduction band energy modulation) along with the ability to modulate the electronic interaction between the two components opens a new pathway for developing and optimizing highly active catalytic materials for the visible-light-driven reduction of CO₂.

EXPERIMENTAL SECTION

Chemicals. Cobalt(II) chloride hexahydrate (Acros Organics, 98%), 4-methoxycarbonylphenylboronic acid (J&K, 98%), Pd(PPh₃)₄ (J&K, 98%), 1-hydroxybenzotriazole hydrate (HOBt, Sigma-Aldrich, 97%), *N*-(3-(dimethylaminopropyl)-*N'*-ethylcarbodiimide hydrochloride (EDC, Sigma-Aldrich, >99%), *N,N*-dimethylformamide (DMF, Acros Organics, 99.8%), triethylamine (TEA, Acros Organics, 99%), phenol (PhOH, Fluka, Ultra >99.5%), and acetonitrile (ACN, Acros Organics, 99.8%) were used as received. Ultrapure water was obtained from a TKA MicroPure system. Argon (>99.998%) and ¹²CO₂ (>99.7%) were from Air Liquide, whereas ¹³CO₂ (99% content in atom ¹³C) was from Aldrich. The filter membrane (0.1 μm pore size, 47 mm diameter, hydrophilic membrane) was purchased from Millipore.

Synthesis. 1,3-Dimethyl-2-phenyl-2,3-dihydro-1*H*-benzo[*d*]-imidazole (BIH)⁴⁹ and 4-bromo-2,2':6',2'':6''-quaterpyridine⁵⁰ were prepared according to the literature. For the synthesis of 4-([2,2':6',2'':6''-quaterpyridin]-4-yl)benzoic acid (qpy-Ph-COOH), a mixture of 4-bromo-2,2':6',2'':6''-quaterpyridine (117 mg, 0.30 mmol), 4-methoxycarbonylphenylboronic acid (63 mg, 0.35 mmol), Pd(PPh₃)₄ (17.3 mg, 0.015 mmol), and Na₂CO₃ (159 mg, 1.5 mmol) in MeOH/H₂O (10 mL, 10:1) was refluxed for 24 h under argon. After evaporation of the solvent, the residue was washed with water and isopropanol to give crude methyl 4-([2,2':6',2'':6''-quaterpyridin]-4-yl)benzoate, which was used without further purification. To the crude methyl 4-([2,2':6',2'':6''-quaterpyridin]-4-yl)benzoate in THF/MeOH/H₂O (6 mL, 4:1:1) was added LiOH (3.0 mmol, 127 mg), and then the mixture was refluxed for 12 h. After cooling, the mixture was filtered and acidified with 1 M HCl. Upon removing organic solvent, a white product (qpy-Ph-COOH) was precipitated out. The solid was filtered and washed with water. After drying under vacuum for 12 h, qpy-Ph-COOH was obtained as a white solid (Figure S1). Yield: 87 mg (67%). ¹H NMR (400 MHz, DMSO-*d*₆): δ 8.98 (s, 1H), 8.92–8.88 (m, 3H), 8.85 (d, ³*J*_(H,H) = 4.0 Hz, 1H), 8.76 (d, ³*J*_(H,H) = 8.0 Hz, 1H), 8.63 (d, ³*J*_(H,H) = 8.0 Hz, 1H), 8.55 (d, ³*J*_(H,H) = 8.0 Hz, 1H), 8.28–8.22 (m, 3H), 8.17–8.12 (m, 4H), 8.03 (d, ³*J*_(H,H) = 4.0 Hz, 1H), 7.70 (t, ³*J*_(H,H) = 8.0 Hz, 1H).

For the synthesis of Coqpy-Ph-COOH, a mixture of qpy-Ph-COOH (43 mg, 0.1 mmol) and CoCl₂·6H₂O (36 mg, 0.15 mmol) in CH₃OH (20 mL) was stirred at room temperature for 24 h. After evaporation of solvent, the residue was washed with water and isopropanol to give Coqpy-Ph-COOHCl₂·2.5H₂O as a pale yellow solid (Figure S1). Yield: 50 mg (83%). Coqpy-Ph-COOHCl₂·2.5H₂O: Anal. Calcd (Found) for C₂₇H₁₈Cl₂CoN₄O₂·2.5H₂O: Co, 9.37 (9.74), C, 53.57 (53.68), H, 3.83 (3.66), N, 9.26 (9.15). ESI-MS in MeOH (Figure S2): *m/z* 244.7, [Coqpy-Ph-COOH]²⁺; 524.0, [Coqpy-Ph-COOH Cl]⁺.

Physical Characterization. UV–vis absorption spectra were measured with a Cary 60 spectrophotometer (Agilent Tech.). Emission quenching measurements were conducted with a Cary Eclipse fluorescence spectrophotometer (Agilent Tech.), with the excitation wavelength set at 400 nm. Emission intensities used for the Stern–Volmer analysis were taken at 488 and 543 nm, i.e., the emission maximum of mesoporous graphitic carbon nitride. Infrared spectra of Coqpy, C₃N₄, and Coqpy@C₃N₄ hybrid were measured in reflection mode by using a Spectrum BX FTIR spectrometer (PerkinElmer) equipped with a microscope with an MIR light source and a LiTaO₃ detector. Spectra were averaged over eight scans with a resolution of 4 cm⁻¹ and were normalized to the background signal. Samples were carefully dried under vacuum (overnight) prior to measurement. The composition (contents in oxygen, nitrogen, carbon, and cobalt elements) of Coqpy@mpg-C₃N₄ samples were determined by X-ray photoelectron spectroscopy (XPS) recorded on an ESCALAB 250 spectrometer (Thermo Sci.) accompanied by a microfocalized X-ray source (K α Al 1486.6 eV) and a double monochromator. The BET surface area and pore size distribution measurement were determined by BELSORP-Max (MicrotracBEL Corp., Japan). Inductively coupled plasma–optical emission spectroscopy (ICP-OES) was obtained with an iCAP 6300 ICP-OES CID spectrometer (Thermo Sci.) to determine the concentration in Co in the different hybrid materials. A test was performed with a RACHID detector including a diode array, a peristaltic pump, and argon plasma (analyzes present at 50 rpm, 1150 W).

Electrochemical and Photoelectrochemical Measurements.

For photocurrent measurements, 2 mg of each solid material was suspended into 250 μ L of DMF. The obtained catalytic ink was then dropped on one face of a fluorine-doped tin oxide (FTO) glass electrode (150 μ L for a 1 cm²) and allowed to dry under ambient conditions prior to use it as the working electrode. The Coqpy/mpg-C₃N₄ sample was prepared by grinding mpg-C₃N₄ with Coqpy-Ph-COOH. All electro- and photoelectrochemical experiments were performed in a glass cell with a quartz window in 0.1 M Na₂SO₄ electrolyte by using a conventional three-electrode system with a platinum wire as a counter electrode and a saturated calomel electrode (SCE) as reference (–0.241 V vs NHE). Light illumination was provided by a 300 W xenon lamp (Oriol Inst.). The amperometric photocurrent was measured for each switch on/off event by using an Autolab PGSTAT 128N potentiostat (Metrohm) with no bias voltage under the UV–vis light irradiation with light chopping every 20 s. Electrochemical impedance spectroscopy (EIS) plots were measured by using a PARSTAT 4000 potentiostat (Princeton Applied Res.) at the corresponding open circuit potential over the frequency ranging from 0.1 to 10⁶ Hz in the dark.

Photocatalytic Experiments. Photocatalysis, durability, and recycling tests were conducted in a quartz cell with an inner volume of 6 mL. Coqpy@mpg-C₃N₄ hybrid material (6 mg) was dispersed into an ACN solution (3 mL) containing BIH (0.05 M) as an electron donor and PhOH (0.03 M) as a proton source. The suspension was poured to the test cell and then sealed with a rubber septum. After bubbling CO₂ for ca. 30 min, the suspension was irradiated with an AM1.5G solar simulator equipped with a 100 W xenon lamp and a 400 nm long-pass optical filter. The temperature was controlled during each experiment by a water bath at 298 K. Control experiments (with no catalyst, no CO₂, or no light) and comparison experiments (using either Coqpy/mpg-C₃N₄, Coqpy-mpg-C₃N₄, Coqpy/Al₂O₃, Coqpy@nsg-C₃N₄, or CoCl₂) were conducted in the same conditions as the full system.

Sample Preparation for ICP-OES Analysis. Heterogeneous samples were preprocessed by acid digestion with concentrated nitric acid. In the present case, 2.0 mg of solid sample was introduced into 1.0 mL of concentrated nitric acid and heated on a hot plate. After cooling to room temperature, the reaction solution was filtered. Then the filtrate was diluted into a 10.0 mL volumetric flask of ultrapure water for testing.

Reduction Products Analysis. Gaseous reduction products in the sample headspace were analyzed by gas chromatography (GC) equipped with an activated carbon column and a thermal conductivity

detector (TCD, GL Sciences, GC323), with argon as the carrier gas. Calibration curves for H₂ and CO were established individually by filling pure gases to a tube with a graduated gastight syringe (Hamilton). ¹³C-labeled experiments were conducted following the same procedure but were also analyzed by a gas chromatograph mass spectrometer (GCMS-QP 2020, Shimadzu). Ionic chromatography (Dionex ICS-1100 ion chromatography system, Thermo Scientific) was employed to check the liquid reduction products in the solution.

Turnover Number (TON) Calculation. TON is defined as the maximum number of catalytic cycles reached until the system stopped reducing CO₂. Mole numbers of H₂ and CO were obtained from the conversion of GC peak areas into moles in the sample headspace thanks to calibration curves. Data points are the result of at least two individual experiments, and the relative error on TONs is ca. 5%, corresponding to the size of the data points. TON was based on the mole amount of Coqpy in hybrid Coqpy@C₃N₄.

ASSOCIATED CONTENT

UV–vis and infrared spectra, absorption, XPS, XRD, cyclic voltammetry, GC, GC-MS, emission quenching, SEM images and supplemental references (Tables S1–S3, Figures S1–S11) (PDF)

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Notes

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ABBREVIATIONS

g-C₃N₄, graphitic carbon nitride; mpg-C₃N₄, mesoporous graphitic carbon nitride; nsg-C₃N₄, nanosheet of graphitic carbon nitride; ACN, acetonitrile; DMF, *N,N*-dimethylformamide.

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