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2D Covalent Organic Frameworks as Intrinsic Photocatalysts for Visible Light-Driven CO₂ Reduction

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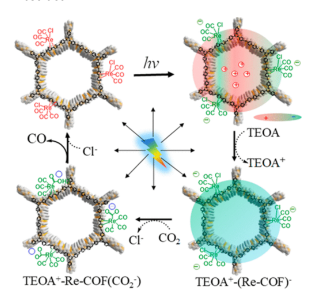
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Abstract



Covalent organic framework (COF) represents an emerging class of porous materials that have exhibited great potential in various applications, particularly in catalysis. In this work, we report a newly designed 2D COF with incorporated Re complex, which exhibits intrinsic light absorption and charge separation (CS) properties. We show that this hybrid catalyst can efficiently reduce CO_2 to form CO under visible light illumination with high electivity (98%) and better activity than its homogeneous Re counterpart. More importantly, using advanced transient optical and X-ray absorption spectroscopy and *in situ* diffuse reflectance spectroscopy, we unraveled three key intermediates that are responsible for CS, the induction period, and rate limiting step in catalysis. This work not only demonstrates the potential of COFs as next generation photocatalysts for solar fuel conversion but also provide unprecedented insight into the mechanistic origins for light-driven CO_2 reduction.

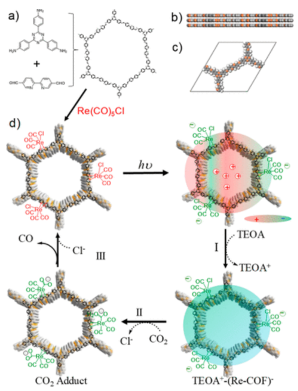
Efficiently capturing CO_2 and simultaneously converting it to chemical fuels driven by solar energy is a promising approach to address energy crisis and climate issues. (1–3) The essential challenge in reaching this elusive goal is to formulate a rationally designed photocatalytic system that can effectively couple a given photosensitizer (PS) with an appropriate molecular catalyst (MC), thereby enabling efficient photosensitization of multielectron

reduction catalysis. (4–6) While many molecular- or semiconductor-based photocatalytic systems have been designed, they all suffer difficulties, such as poor CO_2 adsorption, inappropriate architecture of active sites, rapid charge recombination (CR) or low selectivity etc. (1,7–9)

As an emerging class of crystalline porous materials, covalent organic frameworks (COFs) represent a versatile platform offering new promise for photocatalytic CO_2 reduction.(10–20) COFs are built from periodic organic building blocks via covalent bond formation, providing an innovative approach for the construction of robust photocatalytic materials with built-in PS (i.e., extended π -conjugation) and MC (e.g., incorporated via postsynthetic modification), thereby facilitating efficient charge separation (CS) and precise determination of the nature of the incorporated MC. Moreover, these structurally diverse materials, with large surface areas and readily tunable pore sizes are expected to provide an ideal scaffold for CO_2 adsorption, diffusion, and activation. However, this undeniable potential has yet to be realized, with some recently studied initial systems exhibiting moderate efficiencies.(21–24) Given the inherent advantages of COFs as photocatalysts and their potential impact on the global energy crisis,(25–29) no time should be wasted in undertaking a well-designed plan to further develop useful devices.

Addressing this pressing need, herein, we report a newly designed COF photocatalyst with a photoactive 2D triazine COF as PS and incorporated tricarbonylchloro(bipyridyl) Re complex (Re(bpy)(CO) $_3$ Cl) as CO $_2$ reduction MC (denoted Re-COF). We show that Re-COF can effectively reduce CO $_2$ to CO with high selectivity (98%) and durability upon visible light illumination. More importantly, the combination of *in situ* and time-resolved absorption spectroscopy uncovered the key intermediate species that are responsible for CO $_2$ reduction.

The triazine COF was synthesized from 2,2-bipyridyl-5,5-dialdehyde (BPDA) and 4,4',4"-(1,3,5-triazine-2,4,6-triyl) trianiline (TTA) by solvothermal reactions (Scheme 1a and SI). The formation of imine linkages between aldehyde and TTA in COF was confirmed by FT-IR spectrum (Figure 1a), where we observed the formation of C=N stretching modes at 1626 cm⁻¹ that is characteristic of imine in COF and the vanishing of amino band (3213–3435 cm⁻¹) and aldehyde band (1673–1692 cm⁻¹) that were present in TTA and BPDA.(15,16,30–32) The formation of the COF macrostructure was further supported by the additional absorption band (~440 nm) observed in its diffuse reflectance UV-visible spectrum (Figure 1b), which arises from the delocalized intramolecular charge transfer (ICT) band due to π -conjugation of TTA and BPDA.(33–35) Powder XRD patterns of COF show prominent diffraction peaks (Figure 1c), indicating its crystalline nature. The lattice model was simulated using Material Studio 8.0,(30) from which we obtained the most probable structure of COF with AA stacking mode (Scheme 1b,c). Pawley refinement of the simulated structure yields XRD patterns that agree well with the experimental data, as indicated by the negligible difference between the simulated and experimental data (middle panel of Figure 1c), suggesting the validity of the computational model.



Scheme 1. (a) Synthesis of COF and Re-COF; (b) Side View and (c) Unit Cell of AA Stacking COF; (d) Proposed Catalytic Mechanism for CO₂ Reduction

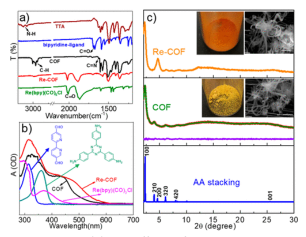


Figure 1. FT-IR (a) and diffuse reflectance UV–visible spectra (b) of COF, Re-COF and their starting materials. (c) Powder XRD patterns of Re-COF and COF obtained experimentally (green), through Pawley refinement (red), and via simulation using AA stacking mode (blue). The purple plot in the middle panel is the difference pattern between experimental and simulated data. The insets are the pictures and SEM images of Re-COF and COF.

Re moiety was then incorporated into COF via reaction between bipyridine ligand and Re(CO)₅Cl to form Re-COF (<u>Scheme 1</u>a and <u>SI</u>).(<u>36</u>) This is confirmed by the additional peaks at 1887, 1916, and 2022 cm⁻¹ in FT-IR spectrum of Re-COF, corresponding to –CO vibrational stretching of Re(bpy)(CO)₃ (<u>Figure 1</u>a).(<u>36–38</u>) The XRD patterns (<u>Figure 1</u>c) of Re-COF match well with the simulated data of COF, indicating the preservation of its crystal parameters after Re incorporation. This is further supported by the ¹³C NMR spectrum (<u>Figure S1</u>) and SEM images (inset of <u>Figure 1</u>c) of Re-COF and COF. In addition, the local coordination structure of Re in Re-COF measured using X-ray absorption spectroscopy retains that of Re(bpy)(CO)₃Cl (<u>Figure S2</u> and <u>Table S1</u>), suggesting that Re(bpy)(CO)₃Cl is well preserved in Re-COF. However, it is notable that the UV–visible spectrum

of Re-COF are extended to broader region (<u>Figure 1</u>b). This can be attributed to either the vibronic broadening of COFs, (39) or the increased delocalization due to the chelation of Re(bpy)(CO)₃Cl. The permanent porosity of COF and Re-COF was confirmed by N_2 sorption measurements at 77 K (<u>Figure S3</u>).

As light absorption and CS are the key initial step to determine whether Re-COF can be used as photocatalyst for CO_2 reduction, we first examined the CS dynamics in Re-COF using transient absorption (TA) spectroscopy following 530 nm excitation (where the TA spectra of Re(bpy)(CO)₃Cl show negligible signal). The TA spectra of COF (Figure 2a) show a positive feature at 600 nm, which can be assigned to the absorption of excited ICT,(41,42) and a negative band centered at 500 nm, which is likely due to stimulated emission (SE)(40,41) (Figure S4) although the contribution from ground state (GS) bleach cannot be completely excluded. The formation (rising component) of the excited ICT (COF@ICT, Figure 2b) is ultrafast with ~200 fs time constant and the CR time is 19.4 ps (Table S2). Compared to that of COF, the TA spectra of Re-COF show a similar SE band and the formation of the excited ICT state (Figure 2c). However, its excited ICT band is much broader than that in COF, which can be attributed to its broader GS spectrum. More interestingly, this excited ICT state of Re-COF (Re-COF@ICT, Figure 2b) exhibits much longer lifetime ($\tau = 171$ ps) than that of COF, suggesting that the incorporation of Re(bpy)(CO)₃Cl inhibits CR in Re-COF.

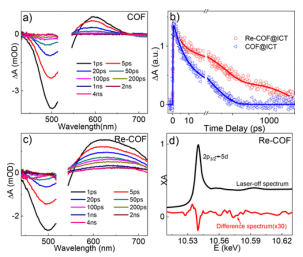


Figure 2. Transient optical spectra of COF (a) and Re-COF (c) following 530 nm excitation. (b) Comparison of kinetics for ICT of COF and Re-COF. (d) XANES spectrum of Re-COF at Re L_3 -edge. The bottom panel is the difference spectrum after subtracting the laser-off spectrum from laser-on spectrum collected at 150 ps delay time.

To gain insight on the role of Re(bpy)(CO)₃Cl play in elongating the excited ICT lifetime, we directly examined the electron density at the Re center in Re-COF using X-ray transient absorption (XTA) spectroscopy. Figure 2d shows the X-ray absorption near edge structure spectrum of Re-COF collected at Re L₃-edge, which is featured by a prominent white line transition corresponding to dipole allowed $2p_{3/2}$ –5d transition. (43,44) Also shown in Figure 2d is the difference spectrum obtained by subtracting the GS spectrum from the spectrum collected at 150 ps. The positive signal observed at 10.531 keV directly supports that Re edge shifts to lower energy, suggesting that photoexcitation of Re-COF leads to the reduction of Re center in Re-COF. This is further supported by the negative feature observed at 10.538 keV: the reduction of Re decreased the number of empty d orbitals, prohibiting $2p_{3/2}$ –5d transition and thus decreasing its absorption intensity. These results, together with elongated ICT lifetime observed in TA studies, suggest that the electrons in the excited ICT state of Re-COF are partially located in Re(bpy)(CO)₃Cl, i.e., electron transfer (ET) indeed occurs from COF to Re(bpy)(CO)₃Cl, which explains the retarded CR in Re-COF.

The demonstration of ET from COFs to Re(bpy)(CO)₃Cl suggests the feasibility of Re-COF as photocatalysts for solar fuel conversion. Accordingly, we proceeded to examine its photocatalytic activity for CO₂ reduction using TEOA as the sacrificial donor and Xe lamp (cutoff wavelength = 420 nm) as light source. Under the optimized conditions (SI and Figure S5). The system can generate ~15 mmol CO/g of Re-COF steadily for >20 h after ~15 min induction period (Figure 3a), accounting for a TON of 48 and 22 times better than its homogeneous counterpart (Figure S5). Because only 2% H₂ was produced in the gas phase (Figure S5), this system has high selectivity for CO₂ reduction to generate CO (98%). Isotopic experiment using 13 CO₂ was performed under the same catalytic conditions. The produced 13 CO (m/z = 29) shown by gas chromatography mass spectrometry (Figure S6) confirms that the generated CO comes from CO₂. The recycling experiments after every three hours of reaction show that the catalytic activity persists for at least 3 cycles (lower right inset of Figure 3a).

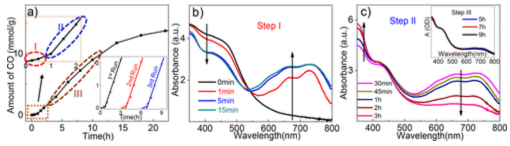


Figure 3. (a) Amount of CO produced as a function of time. The top left inset shows the zoomed in profile in the first 2 h of reaction, and the lower right inset shows the recyclability of the system after three 3-h experiments. The *in situ* diffuse reflectance UV–visible spectra of Re-COF under standard photocatalytic conditions within 15 min (b) and 3 h (c). The inset of panel c shows the *in situ* spectra collected from 5 to 9 h.

To gain more mechanistic insight, we collected the *in situ* diffuse reflectance UV–visible spectra of the catalytic system under the catalytic conditions. Immediately following illumination, prominent absorption in 550–800 nm region that resembles the broad absorption of the ICT band in TA was observed (<u>Figure 3</u>b), and can thus be attributed to the formation of the excited ICT state. The intensity of this ICT band increases in the first 15 min, accompanied by the depletion of absorption at 400–500 nm with an isosbestic point at 539 nm, suggesting that such spectral evolution corresponds to the same process. While similar evolution was observed in the system without CO₂, significantly less evolution was observed in the absence of TEOA (<u>Figure S7</u>), suggesting that reduction quenching of Re-COF by TEOA with the formation of a formal TEOA*-(COF-Re)⁻ CS state contributes to the evolution. The time window for this spectral evolution agrees with the induction period and can thus be attributed to the accumulation of TEOA*-(COF-Re)⁻ CS state before catalysis initiates (Step I, <u>Figure 3</u>a).

After the induction period, the absorption at 550–800 nm decreases significantly within 3 h while the feature at <430 nm grows and a new isosbestic point is observed at 430 nm (Step II, Figure 3c), which results in a distinct feature from Step I and suggests the formation of a new intermediate species. Note that negligible or very slight evolution corresponding to Step II was observed in the system without CO_2 or by replacing Re-COF by COF (Figure S8), suggesting that the intermediate species formed during Step II is associated with CO_2 and Re moiety. After Step II, the spectral evolution stops (Step III, Figure 3c), consistent with the time window for steady generation of CO, suggesting that the system reaches an equilibrium state.

Together, these spectroscopic results point to a mechanistic pathway proposed in <u>Scheme 1</u>d. Upon illumination, the catalytic cycle initiates with the formation of ICT state which is quickly reduced by TEOA, forming a TEOA $^{+}$ -(COF-Re) $^{-}$ CS state (Step I). This formal CS state with reduced Re-COF is then able to capture CO $_{2}$ to form the next intermediate species (Step II). According to previous literature, (6,36–38,45,46) this

intermediate species is likely the CO_2 adducts such as $TEOA^+$ -(COF-Re[CO_2])- or/and $TEOA^+$ -(COF-Re[CO_2 H])-, which is formed after the dissociation of CI- from Re moiety. As the spectra persist during steady generation of CO (Step III), the consumption of the CO_2 adducts to eventually form CO represents the rate limiting step of the catalytic reaction.(34–36,43,44)

In conclusion, we report a newly designed COF hybrid catalyst by incorporating Re(bpy)(CO) $_3$ Cl into 2D triazine COF via postsynthetic modification. We show that this system can efficiently reduce CO $_2$ with better activity than its homogeneous counterpart and high selectivity and stability. Using TA and XTA spectroscopy, we show that this system can undergo facile ICT through ET from photoexcited COF to Re moiety. Using *in situ* diffuse reflectance UV–visible spectroscopy, we unraveled three key intermediate species that are responsible for CS, induction period, and rate limiting step in CO $_2$ reduction. These results not only demonstrated the great potential of COFs as effective solar fuel photocatalysts but also provided unprecedented new insight into the catalytic mechanism for CO $_2$ reduction.

The authors declare no competing financial interest.

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Supporting Information

The Supporting Information is available free of charge on the <u>ACS Publications website</u> at DOI: 10.1021/jacs.8b09705.

Synthesis and standard characterization of Re(bpy)(CO) $_3$ CI, COF, Re-COF, experimental details of transient and *in situ* spectroscopy, 13 C NMR of Re-COF, SEM images of COF, fitting parameters for TA and XAS, emission spectra, optimization of photocatalytic reactions, stability of COF in different solvents, and Re distribution (PDF)

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