Covalent Organic Frameworks: A Promising Materials Platform for Photocatalytic CO$_2$ Reductions

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Abstract: Covalent organic frameworks (COFs) are a kind of porous crystalline polymeric material. They are constructed by organic module units connected with strong covalent bonds extending in two or three dimensions. COFs possess the advantages of low-density, large specific surface area, high thermal stability, developed pore-structure, long-range order, good crystallinity, and the excellent tunability of the monomer units and the linking reticular chemistry. These features endowed COFs with the ability to be applied in a plethora of applications, ranging from adsorption and separation, sensing, catalysis, optoelectronics, energy storage, mass transport, etc. In this paper, we will review the recent progress of COFs materials applied in photocatalytic CO$_2$ reduction. The state-of-the-art paragon examples and the current challenges will be discussed in detail. The future direction in this research field will be finally outlooked.

Keywords: covalent organic frameworks; CO$_2$ reduction; photocatalysis; functional materials design

1. Introduction

In recent years, the anthropogenic consumption of fossil fuels increased with an exponential growth rate. This led to energy exhaustion, environmental pollution and the warming-up effect by CO$_2$ emission. All of these have severely threatened the sustainable and harmonious development of human society. The environment and energy crises have become the most lethal enemies for human-beings. Developing efficient approaches to reduce the enormous amount of carbon dioxide being discharged into the environment has become an urgent need. In nature, plants use sunlight to convert carbon dioxide into glucose by photosynthesis [1]. Inspired by this, chemists began to focus on the effective methods to use solar energy to convert CO$_2$ into valuable chemicals about a century ago [2,3]. Due to the more and more serious situations facing to the human society, converting CO$_2$ into fuels, other useful stock and fine chemicals has become one of the best options and been considered as “the holy grail” to reduce green-house gas emission, counter the energy shortage and ameliorate the environment pollution. It can be seen as the solution to “kill three birds with one stone” [4–7].

CO$_2$ reduction reaction is a considerably endothermic process, which makes it unfavorable in thermodynamics. Compared with water-splitting reaction, the CO$_2$ reduction reaction requires more in-put energy (see Equations (1)–(6) below). Different from common thermocatalytic CO$_2$
reduction, photocatalytic materials not only need to convert solar energy into electron/hole pair possessing oxidizing and reducing ability, but also should be capable of reducing the activation energy for CO2 reduction. Photocatalytic CO2 reduction is a multi-electron transfer process which produces different reduction products at varied redox potentials as shown in Table 1 below. The key to obtain a certain reduction product is to choose the photocatalytic materials with matched energy band position and E0. However, this is considered to be the most challenging point in CO2 reduction. The following processes shown in Table 1 often occur simultaneously, leading to the formation of multiple reduction products. Moreover, the proton can also act as the electron-acceptor to generate H2 by-product, worsening the CO2 reduction selectivity.

\[
\begin{align*}
\text{H}_2\text{O}(l) & \rightarrow \text{H}_2(g) + \frac{1}{2}\text{O}_2(g), \quad \Delta G^0 = 237 \text{ kJ/mol}, \; \Delta E^0 = 1.23 \text{ V.} \\
\text{CO}_2(g) & \rightarrow \text{CO}(g) + \frac{1}{2}\text{O}_2(g), \quad \Delta G^0 = 257 \text{ kJ/mol}, \; \Delta E^0 = 1.33 \text{ V.}
\end{align*}
\]

\[
\begin{align*}
\text{CO}_2(g) + \text{H}_2\text{O}(l) & \rightarrow \text{HCOOH}(l) + \frac{1}{2}\text{O}_2(g), \quad \Delta G^0 = 286 \text{ kJ/mol}, \; \Delta E^0 = 1.48 \text{ V.} \\
\text{CO}_2(g) + 2\text{H}_2\text{O}(l) & \rightarrow \text{H}_2\text{O}(l) + \frac{3}{2}\text{O}_2(g), \quad \Delta G^0 = 703 \text{ kJ/mol}, \; \Delta E^0 = 1.21 \text{ V.}
\end{align*}
\]

**Table 1. Reduction potentials of CO2.**

| Reaction | E0 (V) vs NHE at pH = 7 |
|----------|------------------------|
| 2H+ + 2e- → H2          | -0.41                  |
| CO2 + e- → CO2-         | -1.9                   |
| CO2 + 2H+ + 2e- → HCOOH | -0.61                  |
| CO2 + 2H+ + 2e- → CO + H2O | -0.53               |
| CO2 + 4H+ + 4e- → C + 2H2O | -0.2                 |
| CO2 + 4H+ + 4e- → HCHO + H2O | -0.48                |
| CO2 + 6H+ + 6e- → CH3OH + H2O | -0.38               |
| CO2 + 8H+ + 8e- → CH4 + 2H2O | -0.24                |
| 2CO2 + 8H2O + 12e- → C2H4 + 12OH- | -0.34               |
| 2CO2 + 9H2O + 12e- → C2H5OH + 12OH- | -0.33              |
| 3CO2 + 13H2O + 18e- → CH3OH + 18OH- | -0.32               |

In order to effectively utilize solar energy to convert CO2 to useful chemicals, a key issue should be addressed first; that is, to design and prepare highly-efficient “artificial leaf” materials...
which act as the antenna to absorb solar energy, transform the solar energy to chemical energy and drive the kinetically sluggish and thermodynamic unfavored CO₂ reduction reaction. Since the first report by Inoue and co-workers in 1979 of successfully transforming CO₂ into various hydrocarbons in aqueous solution using semiconductors [8] under sunlight illumination, a renaissance of photocatalytic CO₂ reduction surged. Various inorganic semiconductors were extensively investigated to screen efficient CO₂ photocatalytic reduction catalysts [9–13]. Besides photocatalysts themselves, the choice of electron-donor, photosensitizer and sacrificial reagents can also largely affect the overall efficiency of CO₂ photocatalytic reduction. Using TEOA, ascorbic acid or alcohol as electron-donor and sacrificial reagent would usually provide a much higher CO₂ reduction yield than H₂O because of their lower oxidizing potential and rapid electron-transfer kinetics, but exploiting H₂O as electron-donor and avoiding sacrificial reagent is more ideal from the consideration of sustainability. Although applying photosensitizer such as Ru(bpy)₃Cl₂ can remarkably increase CO₂ photocatalytic reduction efficiency by more effective light-harvesting, combining the photocatalysis and photosensitization function in semiconductor photocatalyst itself without the requirement of additional photosensitizer would be preferable due to the avoidance of the use of expensive and toxic transition-metal complex. If more delicate design and structural optimization of photocatalysts materials can be achieved, the ultimate goal of applying H₂O as the only electron-donor without extra sacrificial reagent and using semiconductor as the sole photocatalyst without noble-metal photosensitizer is not difficult to realize.

Up to now, homogeneous transition-metal complex based catalysts [14,15], heterogeneous inorganic semiconductor catalysts [16,17], and recently developed inorganic-organic hybrid metal-organic frameworks (MOFs) materials [18–20] have found applications in CO₂ photocatalytic transformations. MOFs materials combined the long-comings of homogeneous organometallic-based molecular catalysts and inorganic semiconductor photocatalysts. On one hand, the metal centre and organic ligand can facilitate efficient light harvesting. On the other hand, the transition metal plays a pivotal role in the activation and transformation of CO₂ due to its high affinity towards CO₂ through metal-carbonyl coordination interaction. Moreover, the metal centre can experience fruitful oxidative addition and reductive elimination steps, and stabilize the intermediate state during CO₂ reduction. Further, MOFs are highly porous materials with a huge amount of CO₂ docking sites. The well-defined structure with high crystallinity confirmed the establishment of accurate structure-activity relationships (SARS) as crystalline inorganic semiconductor photocatalysts do. However, since most MOFs materials are composed by weak coordination bonds, only few are stable enough to endure the aqueous condition and achieve the ideal goal of reducing CO₂ with H₂O as the electron donor. Since, in 2005, Yaghi and co-workers first synthesized COF-1 material by the self-condensing of phenyl diboronic acid forming boroxine bonds into a 2D porous polymer with a long-range ordered structure, COFs materials have seen a burgeoning development [21]. Compared with their MOFs counterparts, COFs materials, which are connected via strong covalent bonds, are more difficult to obtain a good macroscopic crystal [22]. By thermodynamically controlled reversible chemistry based on “error-checking” and “proof-reading” mechanisms, a number of COFs materials were synthesized by various methods such as solvothermal, ultrasonic treatment, ionothermal, mechanochemical treatment, room-temperature Lewis-acid catalysis conditions and the interfacial synthesis for COFs films. In comparison with other organic semiconductor-based photocatalyst materials such as carbon nitride, COFs materials possess the tunable skeleton and pore structures to offer plenty of docking sites for catalytically active species and provide a large surface area to fix carbon dioxide [23]. Compared with other porous polymeric materials such as hyper-crosslinked polymer, conjugated microporous polymer, COFs materials own long-range ordered crystalline structure and its structures are clearer than other amorphous porous polymeric materials, which facilitates the demonstration of structure-activity-relationship (SARS) and further optimizing of its performance. Moreover, the pore engineering provides specific interaction with CO₂ by non-covalent bonding, which facilitates its adsorption and reaction in the nano-channels. The pore structure can act as a nano-reactor to
shape a confined micro-environment to facilitate the CO₂ adsorption and reduction products desorption. This selective mass transport flow will highly benefit the catalyst performance. Besides, COFs ordered π-array structure can provide a pre-organized channel for rapid and efficient charge-carrier transport [24], which will further improve the activity of the photocatalytic reduction of CO₂. By condensing electron-rich and electron-deficient moiety into a bicontinuous alternative structure, the 1D column array of donor-donor and acceptor-acceptor can be thus formed [25]. This super-heterojunction structure possesses better charge-transfer and collection behavior in comparison with inorganic semiconductors and common organic 1D polymeric material. Besides these two advantages, the tunability of COFs structure can accommodate different π-conjugated module, electron donor and acceptor groups, which will form excellent antennae and achieve the effective harvesting of the whole sunlight spectrum extending from ultra-violet (UV), visible-light (Vis) to near-infrared light (NIR). These crucial factors, the superb charge carrier transport and collection, the light harvesting, and the adsorptive performance ensure COFs to be promising material platforms for photocatalytic CO₂ reduction [26].

There have been several excellent reviews on COFs materials application in photocatalysis [27], mainly focused on photocatalytic hydrogen evolution [28] or organic transformations [29]. This review will focus on the applications of COFs-based photocatalysts for the CO₂ reduction field. The discussion will be divided into three parts according to the photocatalyst system. Firstly, we will focus on metalated COF photocatalysts and hybrid photocatalysts of COFs materials with homogeneous molecular CO₂ reduction catalyst. The second part will discuss pure COFs photocatalysts for CO₂ reduction without transition-metal moiety. Besides, the covalent triazine frameworks (CTFs) which are capable of photocatalytic CO₂ reduction will also be incorporated. Finally, the current challenge and the future direction of this field will be presented in the conclusion.

2. COFs Application for Photocatalytic CO₂ Reduction

2.1. Metalated COFs and Hybrid COF Photocatalyst Systems with Homogeneous CO₂ Reduction Catalysts

Compared with pure COFs photocatalytic CO₂ reduction materials, metalated COFs and hybrid COF catalyst systems conjugated with homogeneous metal-complex molecular CO₂ reduction catalysts often exhibited better activity. This activity enhancement was mainly attributed to the presence of transition-metal elements, which usually possess outstanding CO₂ adsorption and catalytic capability or have the ability to considerably strengthen the photo-induced charge generation and accelerate charge-transfer process.

In 2019, Lan et al., developed the first metalated COF material for heterogeneous photocatalytic CO₂ reduction. The authors reticulated 2,6-diaminoanthraquinone (DQ) with 2,4,6-triformylphloroglucinol (TP) in DMA (N,N-dimethylacetamide)/1,3,5-trimethylbenzene/6M acetic acid mixed solvent heated at 120 °C for 72 h [30]. The schematic diagram of the reaction mechanism is demonstrated in Figure 1. The as-prepared 2D-anthraquinone-based DQTP-COF possessed plentiful metal coordinating sites via anthraquinone O atoms. Upon the sonication and heating in the aqueous solution with cobalt salt, Co(II) ion was incorporated into the COF structure. The as-prepared DQTP-COF-Co material showed good CO₂ reduction ability under visible-light irradiation (λ ≥ 420 nm) sensitized by Ru(bpy)₃Cl₂ with triethanolamine (TEOA) as electron donor. The formation rate of CO was as high as 1.02 × 10⁸ μmol g⁻¹ h⁻¹, which was much higher than the non-metalated DQTP-COF, DQTP-Ni-COF and DQTP-Zn-COF samples. The selectivity for CO over H₂ was moderate (59.4%). However, when Zn(II) ion was loaded into the DQTP-COF structure, HCOOH was generated with an excellent selectivity up to 90% and a formation rate of 152.5 μmol g⁻¹ h⁻¹. The authors proposed a “two-pathway” mechanism for this distinguishing selectivity. Two pathways forming either HCOOH or CO share the same intermediate, i.e., M-COOH (see Figure 1). If the coordination environment for -COOH was electron-rich, the C-O bond was prone to be cleaved forming CO product, while the electron-deficiency environment would facilitate the
HCOOH formation via proton-coupled-electron-transfer process (PCET), which by-passed the C-O cleavage pathway. Since Co is a good π-electron donor, CO was the main product; while for Zn-based COF, the π-electron acceptor metal centre would promote the formation of HCOOH. While for Ni-COF, which was neither electron-rich nor deficient, the generation rate of CO and HCOOH was almost identical. Lan’s report was the first example using metalated-COF material as heterogeneous CO₂ photocatalyst.

![Figure 1. Proposed mechanism for the photocatalytic reduction of CO₂ with DQTP-COF-M ((M = Zn, Ni and Co). Copied with permission from Elsevier 2019 [30].](image)

Recently, Jiang et al., discovered that via a bottom-up preparation approach, COF-367-Co nanosheet (NS) exhibited considerably high activity for photocatalytic CO₂ reduction to CO under visible-light irradiation conditions [31], and the preparation procedures are shown in Figure 2. The authors discovered that during the condensation of 5,10,15,20-tetrakis(4-aminophenyl)porphinato)cobalt (Co-TAP) and biphenyl-4,4′-dicarboxaldehyde (BPDA) to fabricate COF-367-Co, the addition of excess amount of bulky BPDA analogue 2,4,6-trimethylbenzaldehyde (TBA) would induce the anisotropical growth of COF crystallite. The bulky additive can inhibit growth in the perpendicular direction of the nanocrystallite. Thus, the ultra-thin COF-367-Co nanosheet (NS) could be prepared even in gram scale with more exposed catalytic active sites. The COF-367-Co NS exhibited superior photocatalytic performance in comparison with the bulk COF-367-Co powder with 10,672 μmol g⁻¹ h⁻¹ CO yield and a selectivity of ca. 78% in aqueous solution sensitized by Ru(bpy)₃Cl₂ under λ ≥ 420 nm irradiation with ascorbic acid as electron donor. When bulk COF-367-Co powder material was used under other identical conditions, in sharp contrast, only 65 μmol g⁻¹ h⁻¹ of CO and 514 μmol g⁻¹ h⁻¹ of H₂ were evolved. This great improvement of activity was attributed to the ultra-thin 2D NS structure which facilitated the photo-induced electron transfer evidenced by the transient photoluminescence spectrometry and ultra-fast transient absorption experiments results. Further, the DFT theoretical calculation results demonstrated that cobalt atom plays the role as both CO₂ adsorption and activation sites, generates the key intermediate COOH⁺ with a 0.47 eV low energy barrier, while a much higher 0.74 eV energy barrier is required for proton reduction. This sharp difference accounted for the good selectivity of CO₂ reduction over proton reduction. This example still remains as the record-high activity for photocatalytic CO₂ reduction up till now in spite of the use of noble-metal sensitizer Ru(bpy)₃Cl₂ and ascorbic acid electron-donor.
Moreover, and activity could be realized with the same catalyst. The authors demonstrated that the condensation of TP and 5,5′-diamino-2,2′-bipyridine (BP) under solvothermal conditions generated TpBp-COF material. Moreover, the Ni single site catalyst could be incorporated into the COF structure only through simple impregnation treatment of TpBp-COF with Ni(ClO4)2. The state of single Ni sites on COF supporter was characterized by various techniques including aberration-corrected high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM). The as-synthesized TpBp-COF-Ni single site material exhibited considerably good performance for photocatalytic CO2 reduction under visible-light irradiation sensitized by Ru(bpy)2Cl2 with TEOA as electron donor. The CO formation rate was as high as 966 μmol g−1 h−1. An almost perfect 96% CO over H2 and other reduction products selectivity was achieved. Although TpBp-COF-Co provided higher reduction activity compared with TpBp-COF-Ni, the CO selectivity was poorer in comparison with Ni-based catalyst. The authors applied DFT calculation to investigate the origin of this extraordinary activity and selectivity. The calculation results indicated that the addition of H+ to L2Ni (L means ligand) could not occur easily due to the large 141.4 kcal mol−1 energy barrier, while the formation of L2Ni-CO2 adduct only required activation energy of 3.2 kcal mol−1. The following steps—including the sequential proton and electron addition to L2Ni-CO2, the transformation of Ni-COOH to Ni-CO2, and the release of CO—only needed to overcome considerably smaller energy barriers. Moreover, since the ΔG of COF-Ni adduct formation was less than Ni(bpy)2, the kinetics of CO2 reduction at the Ni single site supported on TpBp-COF were more favorable than H2 reduction. Zou’s example realized the first single metal site catalyst supported on COF to fulfill photocatalytic CO2 reduction with high reduction efficiency and perfect product selectivity.

Figure 2. Schematic illustration of synthesis of the COF-367 NSs for CO2 photoreduction to CO and its STM image. Copied with permission from ACS 2020 [31].

Figure 3. Schematic diagram of photocatalytic selective reduction of CO2 over Ni-TpBpy. Copied with permission from ACS 2019 [32].
Besides metalated-COF photocatalytic CO$_2$ reduction materials, the conjunction of COFs materials with other homogeneous CO$_2$ reduction catalysts is another strategy to achieve highly efficient and selective CO$_2$ reduction under visible-light irradiation. Noble-metal Re-complex molecular catalyst showed outstanding efficiency for CO$_2$ photocatalytic reduction when combined with COF materials. In 2018, Huang et al. reported the first rhenium-COF-based hybrid photocatalyst system, which exhibited superior selectivity up to 98% towards CO$_2$ reduction to CO and high activity, with 15 mmol g$^{-1}$ CO steady generation for more than 20 h, which accounted for a TON 48 and 22 times better than its homogeneous counterpart [33]. Their proposed catalysis mechanism is shown in Figure 4. This rhenium-COF hybrid photocatalytic system was constructed via the postsynthetic modification of the BPDA-TTA-COF which was generated by the Schiff base condensation reaction of 2,2-bipyridyl-5,5-dialdehyde (BPDA) and 4,4',4''-(1,3,5-triazine-2,4,6-triyl) trianiline (TTA) under solvothermal condition. Re-moiety was introduced by the ligand exchange procedure of bipyridine ligand in COF with chloride in the Re precursor Re(CO)$_5$Cl. The as-synthesized Re-COF was characterized by various spectrometric methods including FT-IR, XRD, BET and UV-Vis. To demonstrate the photophysical and photochemical property of this Re-COF hybrid photocatalyst, the TA (ultra-fast transient absorption) and XTA (X-ray transient absorption) including in situ diffuse reflectance UV-Vis-NIR absorption spectrometry techniques were utilized. TA and XTA results indicated that the materials undergo facile intramolecular charge transfer (ICT) through ET (electron transfer) route from photo-excited COF to rhenium moiety, while Re part acted as the CO$_2$ reduction catalytic active sites. From the in situ UV-Vis-NIR spectrum results, the authors discovered three key intermediates—TEOA$^-$-COF$^+$, TEOA$^-$-(COF-Re)$^-$ and TEOA$^-$-COF-Re(CO$_2$)$_2$ adduct—which are responsible for charge separation, induction period and rate-limiting step in the CO$_2$ photocatalytic reduction route. Huang’s example was the first COF-homogeneous metal-complex-based hybrid CO$_2$ photoreduction catalyst with high activity and selectivity.

![Figure 4. Proposed catalytic mechanism of Re-COF.](image)

In 2019, Zou et al. studied a highly efficient, stable and recyclable photocatalyst by embedding photoactive rhenium complex (Re(CO)$_5$Cl) into bipyridine-based Tp-Bpy COFs [34], as shown in Figure 5. The experimental results of FT-IR, N$_2$ isotherm sorption and X-ray diffraction demonstrated that the Re-TpBpy-COF material was constructed as the designed structure. Upon the analysis of the N 1s deconvoluted X-ray photoelectron spectrometry results, it could be confirmed that the pyridinic nitrogen acted as the anchor group for rhenium complex. The as-prepared Re-TpBpy-COF hybrid material showed excellent photocatalytic CO$_2$ reduction activity and
The CO formation rate was as high as 270.8 μmol g⁻¹ h⁻¹ for the first cycle under λ ≥ 390 nm irradiation with TEOA as electron donor. Moreover, the Re-TpBpy-COF photocatalyst possessed good recyclability and reusability providing almost negligibly reduced CO evolution activity after 24 h continuous irradiation, while their homogeneous molecular complex photocatalyst counterpart Re(bpy)(CO)₃Cl showed half the CO formation rate of Re-BpTpy after 4 h irradiation. This example exhibited that the Bpy-based COF material could act as an efficient supporter to anchor photo-active Re-based CO₂ reduction catalyst. Moreover, the Tp moiety of COF provided extra stabilizing effects due to the keto-enamine tautomerism-induced conjugation stabilization effect. From the above two examples (Huang and Zou), we can rationalize that Bpy-based COF can be excellent supporter acting similar as non-innocent ligands to render stability for the homogeneous Re-complex molecular photocatalyst, achieving a recyclable and reusable hybrid CO₂ photocatalytic reduction system. The Re-Bpy-COF based hybrid photocatalyst can be a promising hybrid material system, due to its high activity and stability towards CO₂ reduction to CO in the presence of an electron donor.

**Figure 5.** Top: Illustration for the construction steps of TpBpy and Re-TpBpy-COF. Down: Time course of CO production during photocatalytic CO₂ reduction on (a) TpBpy-COF, (b) Re-Bpy, and (c) Re-TpBpy-COF photocatalysts under visible light irradiation. Insert: evolution of the CO vibration band during the CO₂ reduction on Re-TpBpy-COF (time resolution 1 h/spectrum). Copied with permission from Elsevier 2019 [34].

Apart from the above-mentioned BPDA-TTA-COF and Tp-Bpy-COF, Cooper et al., further improved the CO₂ photocatalytic reduction performance of Re-COF-based materials by introducing pyrene moiety and applying Knoevenagel condensation reaction to fabricate sp²-c-COF structure [35], the synthesis process is demonstrated in Figure 6. The as-prepared Bpy-sp²-c-COF material
could accommodate CO₂ reduction catalyst Re(CO)₅Cl by its metal-coordinating Bpy unit. The sp²-c conjugation structure led to much better visible-light absorption and charge-separation/transfer ability. Furthermore, the authors utilized steady-state and time-resolved emission spectrometry, time-resolved femto-second transient absorption experimental results to detect the formation of a key long-lived (non-emissive) charge-separation state. Moreover, the DFT calculation results indicated that upon visible-light irradiation, the photo-induced electron dwelled in pyrene fragment would transfer to Bpy fragment and further relay to the complexed Re-based CO₂ reduction catalyst. This Re-Bpy-sp²-c-COF exhibited excellent CO₂ reduction performance—yielding 1040 μmol g⁻¹ h⁻¹ and CO over H₂ selectivity up to 81% using TEOA as sacrificial reagents in acetonitrile solution. This example was, up to now, the record activity of photocatalytic CO₂ reduction using hybrid COF-homogeneous CO₂ reduction catalyst hybrid materials.

![Figure 6. Synthesis of Bpy-sp²-c-COF and Re-Bpy-sp²-c-COF. Conditions for Bpy-sp²-c-COF: KOH (4 M) 1,2-dichlorobenzene and 1-butanol (1: 1 mixture), 120 °C, 72 h. Copied with permission form Royal Society of Chemistry 2020 [35].](image)

Besides the Re-based COF/homogenous molecular CO₂ photocatalytic reduction system, other earth-abundant transition-metal complexes could also act as efficient components with COF to construct highly effective visible-light induced photocatalytic CO₂ reduction hybrid material. Very recently, Islam et al. reported that a Co(dmg): co-catalyst can play pivotal role in assisting TFPG(1,3,5-Triformylphloroglucinol)-DAAQ(2,6-diaminoanthraquinone)-COF for CO₂ reduction under blue light irradiation (λ = 445 nm) [36], as demonstrated in Figure 7. It was interesting that the CO₂ photoreduction product was not CO but formic acid. The catalyst system exhibited good activity providing TON up to 125, excellent stability and recyclability without apparent loss of activity for five cycles. The authors proposed a mechanism for this catalytic cycle. Initially, under 455 nm blue light irradiation, only TFPG-DAAQ-COF material was excited because Co(dmg), TEOA and the product HCOOH absorption spectrum all fall out of this region. After the excitation of COF material to its higher energy state, the electron/hole formation process occurs. The highly-reducing conduction band electron reduces Co(III) to Co(II) complex. The Co(II) complex possessed the ability to coordinate CO₂ molecules and transformed them into HCOOH by the redox ability of Co(II) metal centre. The oxidized Co(III) species was sequentially reduced by the excited COF molecule. The COF radical cation species was quenched by TEOA to regenerate ground state COF molecule. After irradiation, the newly excited COF molecular re-entered into the next catalytic cycle. This example indicated the possibility to use an earth-abundant CO₂ reduction co-catalyst to facilitate COF-based photocatalysis.
Figure 7. **Top:** Proposed reaction cycle for the photocatalytic CO$_2$ reduction using COF and Co(dmg) as co-catalysts. **Down:** Proposed reaction cycle for the photo-catalytic reduction of CO$_2$ into HCOOH. Copied with permission from Elsevier 2019 [36].

Lan and co-workers designed and synthesized a series of crystalline porphyrin-tetrathiafulvalene covalent organic frameworks for artificial photosynthesis [37] to combine water oxidation and CO$_2$ reduction in a single COF photocatalyst system. The proposed catalysis mechanism is shown in Figure 8. Electron-deficient metalloporphyrin (TAPP) complexes have excellent light harvesting ability and possess potential CO$_2$ binding ability, while electron-rich tetrathiafulvalene (TTF) is a superb π-electron donor with rapid electron-transfer property. In this 2D COF material, the strong covalent-coupling effect between TAPP and TTF allows the photo-induced-charges (excitons) to be effectively transferred between TTF and TAPP moieties, resulting that the photo-excited electrons resided on porphyrin and holes on TTF can be effectively exploited for separate reduction and oxidation reaction, respectively. Similar with photosynthesis in natural plant chlorophyll, H$_2$O was used as the sole electron-pool for CO$_2$ reduction. Especially, for this COF photocatalyst system, there was no need to add extra photosensitizer, sacrificial reagent and noble-metal co-catalyst. By adjusting and optimizing the monomer chemical structure and metal centre of TTCOFs, TTCOF-Zn achieves the highest photocatalytic CO$_2$ reduction performance yielding 12.33 μmol CO in 6 h irradiation with about 100% selectivity, while H$_2$O acted as final electron-donor and hole-trapping reagent evidenced by the detection of O$_2$ evolution. $^{18}$O and $^{18}$O isotope-labeling experiments unambiguously indicated that CO was produced from CO$_2$ source while O$_2$ was from H$_2$O. Photoluminescence quenching and time-resolved photoluminescence experiments demonstrated that TTCOF-Zn has more efficient charge-transfer property and longer excited-state life-time for electron-transfer events. From DFT calculation, more direct and clearer evidence for understanding the structure-function relationship of multi-site and multi-electron photocatalytic processes were provided. This discovery is especially significant since only H$_2$O and COF are required for photocatalytic CO$_2$ reduction without any additive. Moreover, this example provides new perspectives for the design of the next generation crystalline photocatalysts applied for CO$_2$ reduction.
In recent years, accompanying the tremendous advance in the field of the synthesis and application of covalent organic frameworks, more and more pure COFs CO$_2$ photocatalytic reduction catalysts—without the aid of metal centre or metal-complex CO$_2$ reduction co-catalyst, photosensitizer and even sacrificial reagent—have appeared in the research field. The investigation of these pure COFs CO$_2$ photocatalytic systems is meaningful, since the application of these pure organic systems represents the future direction for large-scale CO$_2$ utilization due to these catalysts being more sustainable and green, without having the secondary environmental effects as the metalated and metal-complex co-catalyzed COFs system do. Although the efficiency of these pure COFs systems is still much lower in comparison with its metalated counterparts, they possess more advantages and long-comings in consideration of sustainability and recyclability. Even though pure COFs CO$_2$ photocatalysts are still in their infancy stage, they represent the future direction for COF-based CO$_2$ photoreduction catalysts.

CO$_2$ photocatalytic reduction can form various products including CO, HCOOH, methanol, formaldehyde, methane and a small amount of C$_2$- products. By the elaborate design and optimization of COFs monomer and topological alignment, differentiated products other than CO could be yielded in the aqueous photocatalytic CO$_2$ reduction by COFs materials. Zhu and co-workers realized metal-free and additive-free photocatalytic CO$_2$ reduction by azine-based ACOF and N$_2$-COF [38], which is shown in Figure 9. Different from Lan’s TTCOF-Zn, liquid fuel compound methanol was isolated as photo-reduction product by ACOF and N$_2$-COF. H$_2$O was applied as electron donor although the O$_2$ detection experiments were not conducted. The amount of generated CH$_3$OH over N$_2$-COF catalyst is 13.7 μmol g$^{-1}$ during 24 h irradiation by visible-light, which is more superior to the performance of ACOF and g-C$_3$N$_4$. The N$_2$-COF was revealed to be a robust catalyst possessing high stability in a five-run cycling test of photocatalytic CO$_2$ reduction and only exhibited slightly lower yield after first two cycling runs. Moreover, the PXRD peaks, FT-IR and N$_2$ adsorption–desorption curves of the used catalyst did not show obvious change after the five-run cycling test compared with the fresh one. Based on the DFT calculation results, N$_2$-COF possesses a suitable HOMO and LUMO energy level (~4.95 eV and ~2.71 eV) which is both large enough to trigger CO$_2$ reduction and small enough to harvest a major portion of visible-light photons. The intramolecular charge transfer event can be induced by photons with wavelengths less than 480 nm. Moreover, the photo-induced hole can be efficiently quenched by H$_2$O via hydrogen bond interaction on polar azine moiety, while the photo-induced electron on LUMO can
sufficiently reduce CO2 on the catalyst surface to methanol due to the matched energy level. Since H2O oxidation is a kinetically sluggish process, the COF material is negatively charged. The N3-COF demonstrated higher activity due to its electron-withdrawing triazine building block compared with phenyl in ACOF, which can more effectively stabilize the negatively charged intermediate species on COF material. Although the activity of the current example of ACOF and N3-COF is not high, the tunability and metal-free property of these COFs-based materials rendered them excellent candidates to supplement current prevalently investigated organic and inorganic semiconductor-based CO2 photo-reduction catalysts.

Almost at the same time with Lan’s TTCOF photocatalyst system, Kong and co-workers discovered highly effective metal-free donor-acceptor type carbazole-triazine COF photocatalyst, which could photo-reduce CO2 to CO using H2O as electron source and hole scavenger without any sacrificial reagent or additives [39], as shown in Figure 10. Experimental results showed that CO and O2 are the sole products from the visible-light induced photocatalytic reduction of gaseous CO2. The generation rate of CO could be optimized to 102.7 μmol g⁻¹ h⁻¹ with the concomitant formation of stoichiometric amount of O2 (51.3 μmol g⁻¹ h⁻¹). 13C and 18O isotope tracing experiments exhibited that CO and O2 products were actually evolved from the photocatalytic reduction of CO2 and oxidation of H2O. Moreover, the authors used in situ FT-IR and DFT calculation to expound on the photo-reaction mechanism. This paper possessed important status in COF photocatalytic reduction of CO2 since the structure-functionality relationship was well demonstrated in this report. The authors indicated that electron donor-acceptor structure was essential for effective CO2 reduction in additive-free COF photocatalysis, which corroborated with Lan’s work using TTCOF to reduce CO2.

**Figure 9.** Schematic diagram for the photocatalytic reduction of CO2 over azine-based COFs upon visible light irradiation. Copied with permission from Elsevier 2018 [38].

**Figure 10.** Proposed photoactive electron transfer and reaction pathway in the photoreduction of CO2 (PET = photoinduced electron transfer). Copied with permission from Wiley 2019 [39].
Very recently, Zhou and co-workers achieved highly efficient photocatalytic reduction of CO2 using TAPPB-COF with gaseous water vapor as the sole reductant without any sacrificial reagents and additives in a gas-solid reaction [40]. The authors modified the COF-366 structure by introducing electron-withdrawing bromine group into this porphyrin COF photocatalyst. The introduction of bromine group enhanced the electron density on LUMO according to the DFT calculation results, which contributed to the change of the valence band potential from 0.80 V to 1.10 V (vs. NHE at pH = 7). The up-lifting of the valence band energy greatly facilitated the oxidation of water since the driving force was much larger in comparison with the pristine COF-366 because the redox potential of the pristine COF-366 valence band potential was too close with O2/H2O (0.82 V vs NHE at pH = 7). The average yield of CO was 24.6 μmol g⁻¹ h⁻¹ under simulated sunlight irradiation (200 ≤ λ ≤ 1000 nm) for TAPPB-COF, which was about 3 times that of the pristine COF-366 materials and 1.5 times that of the g-CN yield under other identical conditions. Moreover, not only was the activity of COF photocatalyst enhanced, the selectivity of CO over H2 was considerably high (95.6% selectivity). This example indicated that the design and tuning of the electronic property of the monomer unit and material modification via a bottom-up approach could have considerably large influences on the COFs semiconductor valence band (VB) position, which is a key factor for the efficiency of water oxidation semi-reaction. If an appropriate VB potential was generated, the facilitated water oxidation semi-reaction would accelerate the other photo-reduction of CO2 semi-reaction.

Besides pure COF-based CO2 photocatalyst, the metal-oxide water oxidation semiconductor could be hybridized to construct Z-scheme heterojunction structure with COF to fulfill efficient overall CO2 reduction only with water in the absence of additive, photosensitizer or sacrificial reagent. CO formation yield could be optimized to 69.67 μmol g⁻¹ h⁻¹ [41], which is the record-high CO2 photoreduction performance without photosensitizer, sacrificial reagent and co-catalysts with only water as electron donor. The authors developed COF-316/COF-318 materials by Sn/Ar reaction between aromatic phenol 2,3,6,7,10,11-hexahydroxynaphthalene (HHTP) and two kinds of o-difluoroarenes, i.e., tetrafluorophthalonitrile (TFPN) or 2,3,5,6-tetrafluoro-4-pyridinecarbonitrile (TFPC) as precursor to synthesize two robust PAE-COFs, and the schematic diagram is shown in Figure 11. The as-prepared COF-316/COF-318 was further covalently linked to inorganic semiconductors such as TiO2, Bi2WO6 and α-Fe2O3 to form COF/inorganic semiconductor hybrid structure. The pyridinecarbonitrile-based COF-318/TiO2 provided the best photocatalytic CO2 reduction activity creating new record for pure COF-based CO2 photocatalytic system. The heterojunction structure was confirmed from transient photocurrent response, photoluminescence and electrochemical impedance spectrometry characterization techniques, these experimental results indicated that COF-318/TiO2 catalyst garnered more effective photo-induced charge carrier separation capability and less recombination probability, and much improved electric conductivity in comparison with the semiconductors displaying better charge-transfer property. Moreover, the DFT calculation and in situ XPS experimental results pointed out that upon 365 nm UV irradiation, both COF-318 and TiO2 components in COF-318/TiO2 were excited. The photo-electrons in TiO2 conduction band transferred to COF-318 pyridinic and cyano nitrogen group, which was evidenced from the positive shift binding energy of Ti 2p orbital meaning that the electron density on Ti sites was reduced. Due to the strong adsorption and interaction with CO2 molecules at pyridinic and cyano nitrogen sites, the electron was conveniently transferred to CO2 coupling with adsorbed proton to yield the key intermediate COOH-. The second proton-coupled-electron-transfer (PCET) was calculated to be the potential-determining-step (PDS). From the DFT results, although the PCET process occurring at cyano nitrogen had a lower over-potential (0.93 eV vs 1.59 eV for pyridinic nitrogen), the first-step activation of CO2 to form COF-CO2 adduct was more favorable at pyridinic nitrogen sites, which made it possible for the PCET process to occur at both pyridinic and cyano nitrogen sites. The authors demonstrated that the coexistence of pyridinic and cyano group in COF-318 facilitated both CO2 activation and reduction processes, which was superior than COF-316 possessing only cyano group. The water oxidation occurred at the TiO2 photo-induced VB
holes naturally. Lan’s report was the only example to hybrid inorganic semiconductors with COF-based material to fulfill highly effective CO₂ photoreduction. The delicate Z-scheme type II heterojunction design made it the record-high performance for CO₂ photoreduction in the absence of photosensitizer, sacrificial reagents and CO₂ reduction catalyst with only H₂O as electron donor up to now.

![Schematic representation of the preparation of COF-318-SCs via the condensation of COF-318 and semiconductor materials.](image)

**Figure 11.** Schematic representation of the preparation of COF-318-SCs via the condensation of COF-318 and semiconductor materials. Copied with permission from Wiley 2020 [41].

### 2.3. CTF-Organometallic Complex-based CO₂ Photocatalytic Systems

As relatively similar porous organic polymers compared to covalent organic frameworks, covalent triazine frameworks (CTFs) composed by the triazine units have demonstrated their potential to be utilized as photo-responsive materials and organic semiconductors for a wide range of photochemical applications including water-splitting, CO₂ reduction and organic transformations due to its nitrogen-rich structure, excellent photo- and chemical stability [42,43].

As early as 2016, Baeg et al. developed a 2D triazine-based CTF material for photocatalytic CO₂ reduction application in conjunction with a Rh-complex electrocatalyst, β-NAD⁺ and formate hydrogenase enzyme and ascorbic acid as electron donor [44], as demonstrated in Figure 12. The authors delicately designed and prepared the perylene-based 2D-CTF materials through a co-condensation reaction between cyanuric chloride and perylene diimide, which maneuvered the challenging synthesis of perylene diimide-based carbonitrile monomers. Furthermore, the authors constructed the CTF-Rh-NAD⁺-formate hydrogenase artificial photosynthetic system for CO₂ reduction with ascorbic acid as sacrificial reagent. The CO₂ reduction product was HCOOH, with a formation rate as high as 881.3 × 10⁹ μmol g-cat⁻¹ h⁻¹ for the CTF photocatalyst under visible-light irradiation λ ≥ 420 nm at 100 mW cm⁻² light density. This CO₂ reduction rate was extremely high. Although the formate hydrogenase enzyme mainly accounted for this extraordinary CO₂ reduction performance, it could be confirmed that the perylene diimide-based 2D CTF really contributed to the much enhanced visible-light harvesting ability, since the HCOOH formation rate was only 261.85 × 10⁹ μmol g-cat⁻¹ h⁻¹ for the PDI monomer under other identical conditions. According to the cyclic voltammetry experimental results, an electron relay mechanism was proposed. Upon the absorption of visible-light, CTF material was excited. The electron dwelling on HOMO was activated to jump to the LUMO energy level in CTF material. The highly-reducing CTF LUMO electron would transfer to Rh(III) complex. The reduced Rh(I) complex further reduced NAD⁺ to NADPH. The NADPH acted as the reducing reagent for CO₂ transformation to HCOOH under the formate hydrogenase enzyme catalysis conditions. The oxidized NAD⁺ re-entered into the catalytic cycle and was further reduced by Rh(I) to regenerate Rh(III) and NADPH. NADPH continuously...
Reduced CO₂ to HCOOH and was recycled by Rh(I) species. The oxidized Rh(III) was reduced by CTF-LUMO electron, while CTF⁺ was quenched by ascorbic acid to regenerate the catalytic active CTF species. This hybrid catalysis assembly unified photocatalysis, transition-metal catalysis, organocatalysis and biocatalysis power to render extremely high CO₂ reduction efficiency. This example set a benchmark for the further development of CTF- and COF-based photocatalytic CO₂ reduction applications.

**Figure 12.** Schematic illustration of the CTF film photocatalyst-enzyme coupled system involved in the exclusive production of formic acid from CO₂. Rhox = [Cp⁺ Rh(bpy)H₂O]²⁺, Rh₄ = Cp⁺ Rh(bpy), Rh₄ = [Cp⁺ Rh(bpy)H⁺]; Cp⁺ = pentamethylcyclopentadieny1, bpy = 2,2'- bipyridine. Copied with permission from RSC 2016 [44].

Apart from the CTF-enzyme-based hybrid CO₂ reduction system, artificial CTF-organometallic composite materials could also act as effective CO₂ photocatalytic reduction catalysts. Cao et al. developed a pyridine-based CTF material and coordinated a single site Re(CO)₆Cl CO₂ reduction catalyst at py-CTF pyridine-triazine bi-dentate chelation sites [45]. The modified Re-py-CTF material showed enhanced CO₂ photoreduction activity up to a 353.05 μmol g⁻¹ h⁻¹ formation rate of CO within 10 h under (200 nm–1100 nm) full sunlight spectrum irradiation in a solid–gas system with TEOA vapor as electron donor—see Figure 13. Moreover, the Re-py-CTF-based photocatalyst exhibited much improved photo-stability upon Re leaching. The heterogeneous photocatalyst system provided good recyclability with negligible loss of activity after five consecutive cycles in the solid–gas system. The authors discovered that when soaked in an acetonitrile/TEOA solution, the solid/liquid photocatalytic reaction system showed a much inferior CO formation yield only after the first cycle due to two factors: firstly, the CO₂ diffusion and mass-transport in the solid-gas system is more efficient than the solid–liquid system because CO₂ can more effectively transport through the pores of CTF material in solid-gas system while in the solid-liquid system, this mass-transport is hindered partly because the solvent molecules can compete with CO₂ for the CTF pore adsorption site. Secondly, about half the Re catalyst was leached into the bulk solution in the solid–liquid system. This example demonstrated the important role for CTF-based material in maintaining the activity of homogeneous transition-metal complex CO₂ reduction catalysts by shielding them from inactivation caused by the leaching effect. This report proved that the solid/gas CO₂ photoreduction may be preferable over solid/liquid reaction in some cases, especially those involving metal-complex moieties, due to the reduced possibility for catalytic active metal component leaching.
Figure 13. Top: Synthesis of CTF-py and Re-CTF-py materials. The inset shows the two types of pores (unit A and unit B). Down: (a) CO evolution over CTF-py, Re-CTF-py and the physical mixture of CTF-py and Re(CO)\(_5\)Cl (Re and CTF-py) in the solid–gas system. (b) Photocatalytic activity of Re-bpy and Re-CTF-py in solid–gas and liquid–gas systems. (c) Production yield of CO over the Re-CTF-py photocatalyst to measure its reproducibility by cycling tests. Copied with permission from RSC 2018 [45].

Besides CTF-based materials with noble-metal cocatalysts for photocatalytic CO\(_2\) reduction, earth-abundant Co could also act as efficient catalytic center to synergistically reduce CO\(_2\) with CTF under visible-light irradiation. Bi et al. reported that by loading cobalt oxide on CTF-1 material, its performance on CO\(_2\) reduction to CO was increased by 44-fold in comparison with pristine CTF-1 and 12-fold with CoO\(_x\) in TEOA/H\(_2\)O/MeCN solution and Ru(bpy)\(_3\)Cl\(_2\) as photosensitizer [46]. The improvement on catalytic performance is mainly due to the enhanced CO\(_2\) capture capacity, extended visible-light absorption and efficient transfer of the charge originated from the modification of cobalt.

The performance of the state-of-the-art example of a COF-based photocatalysis system for CO\(_2\) reduction is summarised in Table 2. To compare the activity of the COF-based photocatalyst for CO\(_2\) reduction with other semiconductor photocatalyst, we summarized the CO\(_2\) reduction efficiency by exploiting H\(_2\)O as electron-donor in various systems in Table 3.
Table 2. The summary of photocatalytic CO$_2$ reduction performance for COFs with different structure.

| Type   | COFs                  | Light           | Additive                        | Conditions       | Product Yield          | Selectivity (%) | Ref |
|--------|-----------------------|------------------|---------------------------------|------------------|------------------------|-----------------|-----|
| Enamine| DQTP-COF-Co           | λ ≥ 420 nm       | Ru(bpy)$_2$Cl$_2$/TEOA         | Liquid (MeCN)    | CO: $1.02 \times 10^8$ µmol·g$^{-1}$·h$^{-1}$ | 59.4            | [31]|
| Imine  | COF-367-Co NS         | λ ≥ 420 nm       | Ru(bpy)$_2$Cl$_2$/ascorbic acid| Liquid (MeCN)    | CO: 10,672 µmol·g$^{-1}$·h$^{-1}$ | 78             | [32]|
| Enamine| TpBp-COF-Ni           | λ ≥ 420 nm       | Ru(bpy)$_2$Cl$_2$/TEOA         | Liquid (H$_2$O)  | CO: 966 µmol·g$^{-1}$·h$^{-1}$ | 96             | [33]|
| Imine  | Re-COF                | λ ≥ 390 nm       | TEOA                           | Liquid (MeCN)    | CO:780 µmol·g$^{-1}$·h$^{-1}$ | 98             | [34]|
| Examine| Re-TpBpy COFs         | λ ≥ 390 nm       | TEOA                           | Liquid (MeCN/H$_2$O) | CO: 270.8 µmol·g$^{-1}$·h$^{-1}$ | N/A            | [35]|
| Olefin | Bpy-sp2-c-COF         | λ ≥ 420 nm       | TEOA                           | Liquid (MeCN)    | CO: 1040 µmol·g$^{-1}$·h$^{-1}$ | 81             | [36]|
| Enamine| TFPG-DAAQ-COF         | λ = 445 nm       | TEOA                           | Liquid (MeCN)    | HCOOH: TOF = 6         | N/A            | [37]|
| Imine  | TTCOF-Zn              | 420–800 nm       | None                           | Liquid (H$_2$O)  | CO: 2.055 µmol·g$^{-1}$·h$^{-1}$ | 100            | [38]|
| Azine  | N$_2$-COF             | 420–800 nm       | None                           | Liquid (H$_2$O)  | CH$_3$OH:0.57 µmol·g$^{-1}$·h$^{-1}$ | N/A            | [39]|
| Imine  | TT-COF                | λ > 420 nm       | None                           | Solid-gaseous H$_2$O | CO: 102.7 µmol·g$^{-1}$·h$^{-1}$ | 98             | [40]|
| Imine  | TAPPB-COF             | 200–1000 nm      | None                           | Solid-gaseous H$_2$O | CO: 24.6 µmol·g$^{-1}$·h$^{-1}$ | 95.6           | [41]|
| Arylether| COF-318-SCs         | 380–800 nm       | None                           | Solid-gaseous H$_2$O | CO: 69.67 µmol·g$^{-1}$·h$^{-1}$ | N/A            | [42]|
Table 3. The summary of photocatalytic CO₂ reduction efficiency using H₂O as electron-donor for COFs and other semiconductor materials [47–61].

| Photocatalyst | Light Source | Product Yield μmol·g⁻¹ h⁻¹ | Ref |
|---------------|--------------|-----------------------------|-----|
| TTCOF-Zn      | 420–800 nm 300 W Xe lamp | CO: 2.055                  | [38]|
| Ni-COF        | 420–800 nm 500 W Xe lamp | CH₃OH: 0.57               | [39]|
| TT-COF        | λ > 420 nm 300 W Xe lamp | CO: 102.7                 | [40]|
| TAPPB-COF     | 200–1000 nm Xe lamp | CO: 24.6                  | [41]|
| COF-318-SCs   | 380–800 nm 300 W Xe lamp | CO: 69.67                | [42]|
| CPO-27-Mg/TiO₂ | UV lamp 4 W 365 nm | CO: 4.09, CH₄: 2.35       | [47]|
| Well-crystallized ordered mesoporous TiO₂ | UV-Vis 300W Xe lamp | CO: 0.145, CH₄: 0.195 | [48]|
| TiO₂          | 300 W Xe lamp (λ ≥ 408 nm) | CO: 50.7, CH₄: 13.5       | [49]|
| Co-ZIF-9/TiO₂ | UV-Vis 300 W Xe lamp (200 < λ < 900) | CO: 17.58 | [50]|
| BiOBr         | 300 W Xe lamp (λ ≥ 400 nm) | CO: 87.4                | [51]|
| Defect-Rich Bi₉O₆Cl₃ Nanotubes | 300 W Xe lamp | CO: 48.6                 | [52]|
| ZIF-8/C₃N₄   | 300 W full-spectrum Xe lamp | CH₃OH: 0.75            | [53]|
| QS-COD (ZIF-67) | 200 W Xe lamp (AM 1.5) | CO: 46.3                | [54]|
| Bi₉O₆Cl₃/g-C₃N₄ | 300 W Xe lamp (λ ≥ 400 nm) | CO: 45.6, CH₄: 6       | [55]|
| Z-scheme CdS–WO₃ | 300 W Xe lamp (λ > 420 nm) | CH₄: 1.02              | [56]|
| HCP-TiO₂-FG   | 300 W Xe lamp (λ ≥ 420 nm) | CO: 27.62, CH₄: 21.63     | [57]|
| α-Fe₂O₃/g-C₃N₄ | 300 W Xe lamp (λ ≥ 420 nm) | CO: 27.2               | [58]|
| C-TiO₂-x@g-C₃N₄ | 300 W Xe lamp (λ ≥ 420 nm) | CO: 205                | [59]|
| TiO₂/N-doped-RGO | 400 W Xe lamp (λ = 250–400 nm) | CO: 44.56             | [60]|
| TiO₂/NH₂-UiO-66 | 1500 W Xe lamp (λ > 325 nm) | CO: 4.25              | [61]|

3. Conclusions

In conclusion, covalent organic frameworks are very promising materials for next-generation photocatalytic CO₂ reduction applications, since they possess multiple long-comings and advantages over traditional metal-oxide, metal-sulfide, noble-metal plasmonic inorganic semiconductor and inorganic-organic hybrid MOF-based photocatalysts. Firstly, due to COFs materials being composed of organic moieties and non-metal elements, the cost and availability of the COF’s monomer are superior to the metal-containing inorganic semiconductors and MOFs. Secondly, since the electronic and steric factor can be conveniently fine-tuned by the choice of differentiated monomer unit and the reticular chemistry, COFs materials provide us almost unlimited space in which to innovate and modify the structure to be more apt to CO₂ adsorption and activation, and to enlarge the absorption range of light from UV to Vis and NIR spectrum and to facilitate charge-carrier formation, separation and transfer process to achieve more efficient CO₂ photoreduction activity and selectivity. Thirdly, due to the property of the linking strong covalent bond, most COFs (except boron-based) are very inert to water, organic solvents, moisture and other conditions.
open-air conditions, with some robust keto-enamine-based Tp-COF being extremely stable and able to endure high-temperature boiling water and concentrated acid and base soaking for long time periods. This property is very different from their metal-containing MOF counterparts, which are woven by weak coordination bonds and most MOFs are unstable in water solution due to that the metal ion are prone to escape from the organic ligand confinement to become free ion in the bulk solution. The stability of COFs in multiple conditions made it promising for the activation of CO$_2$ in some extreme conditions. Fourthly, due to its huge surface-area and developed porous structure, a considerably large amount of catalytic active sites and light-harvesting antenna moieties are exposed on the surface in comparison with non-porous inorganic metal-oxide and metal-sulfide photocatalysts. The exposure of active sites promotes the photocatalytic reaction on the surface with the formation of more catalytic active species. Although COFs materials have many such advantages, there are still many gaps to overcome, and challenges to resolve in this research field. Firstly, the extreme difficulty in obtaining a large enough 2D COF single-crystal for common X-ray single crystal diffraction characterization still restricts the further development of 2D COFs multiple applications, including photocatalytic CO$_2$ reduction, since the current COF structure elucidation is mainly based on the calculation and the powder X-ray diffraction technique, which relies heavily on experiences, and much uncertainty accompanies the theoretical simulation results. The uncertainty of the COFs structure greatly restricted the elucidation of the SARS in its photocatalytic CO$_2$ reaction application. Besides, the CO$_2$ photoreduction rate is still too low. The catalyst activity should be improved from the current μmol g$^{-1}$ h$^{-1}$ scale to mmol g$^{-1}$ h$^{-1}$ and even mol g$^{-1}$ h$^{-1}$. The improvement of catalyst activity calls for deeper investigations into the mechanism of COFs-based CO$_2$ photocatalysts. Only if the black-box of the mechanism been deciphered could we design more effective COFs for CO$_2$ photoreduction. We are confident that if the above-mentioned issues are soundly addressed, COFs-based materials will have a prosperous and booming future, and become pivotal in settling the CO$_2$ reduction problem.

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