

Review

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Posted Date: 9 April 2025

doi: 10.20944/preprints202504.0806.v1

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Review

Recent Developments in the Use of Covalent Organic Frameworks for Photocatalytic Water Decontamination

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Abstract: Water pollution by persistent organic and inorganic contaminants implies a significant trouble for ecosystems and public health. Organic substances such as dyes, pharmaceutical residues, pesticides, and phenolic compounds are increasingly detected in water due to industrial and agricultural activities. Alongside these, toxic heavy metals contribute to the complexity of water treatment challenges. Conventional remediation methods often fall short due to high operational costs or limited efficiency. In this context, photocatalysis has emerged as a promising approach for pollutant degradation in water under light irradiation. In this sense, covalent organic frameworks (COFs), a class of porous, crystalline materials formed by covalent linkage of organic units, offer great advantages as photocatalysts. Their tunable electronic properties, structural diversity, and high stability under aqueous conditions make them ideal for visible-light-driven processes. This review explores the structural features that govern the photocatalytic activity of COFs, including conjugation, bandgap modulation, and donor-acceptor structures. Mechanistic insights into photocatalytic degradation are also discussed. Finally, examples of pre-designed COFs are presented with their application in photodegradation of water pollutants and their main reactive oxygen species (ROS) involved in the photodegradation mechanism. Overall, this review aims to provide a foundation for the rational design of COFs in advanced water treatment technologies.

Keywords: covalent organic framework; photocatalysis; water decontamination

1. Introduction

The limited availability of safe drinking water is a growing problem worldwide.[1] In fact, water is related to many aspects of economic and social challenges and is itself one of the main objectives of sustainable development. In recent decades, several new organic compounds have found their way into water supplies because of intense human activities. Among others, synthetic dyes, pesticides, pharmaceutical and phenolic compounds have increasingly contaminated water sources.[2] Synthetic dyes, extensively used in the textile, food, and cosmetic industries, are highly stable and their strong coloration can reduce light penetration, affecting aquatic photosynthesis.[3,4] Pharmaceutical compounds, including antibiotics, or analgesics, enter water bodies and, even at trace levels, they can contribute to the development of antibiotic resistance.[5] Additionally, pesticides, widely used in agriculture, can leach into groundwater and surface water, leading to long-term environmental persistence.[6] Phenolic compounds, commonly found in petrochemical plants, and paper production, are known for their high toxicity and resistance to degradation with a significant threat to aquatic ecosystems and human health.[7] The adverse effects of these pollutants on flora, fauna and humans can be catastrophic. In addition to their potential toxicity, organic compounds can cause taste and odor problems in water. Unfortunately, once introduced into the environment, some

of these compounds are extremely persistent due to their high chemical stability. Apart from organic molecules, a high concentration of heavy metals and other inorganic pollutants contaminate water in industrial environments.[8] In general, these pollutants are not biodegradable, so they can persist largely in the environment. The accumulation of heavy metals is a serious public health problem. For example, Cr(VI) is easy to be absorbed by the human body acting as a mutagen, carcinogen, and teratogen.[9] The sources of these metals are diverse and are largely related to several industrial applications in chrome plating, dye manufacturing, textile industry, aircraft industry, leather tanning, wood preservation, and mud drilling.

Considering the consequences of water pollution, it is obvious that they constitute a global problem in today's interconnected world. Therefore, it is highly desirable to look for integrative approaches that simultaneously address several types of contaminants without the formation of significant toxic by-products. A classic way to decontaminate water is the use of chlorine or ClO₂ for oxidative degradation of organic and biological matter.[10] In addition, among the conventional processes to remove heavy metals from wastewater, the most used is chemical precipitation, which involves the formation of their insoluble hydroxides, carbonates or sulfides, followed by sedimentation and filtration.[11] Although effective, these procedures involve costly infrastructures, the use of toxic reagents and the extensive formation of residues. It is therefore necessary to explore alternatives that avoid these drawbacks. For this reason, new strategies have been explored in recent years that take advantage of light energy to trigger photocatalytic processes in the degradation or detoxification of pollutants.[12] Thus, many photooxidative processes are based on the light-induced formation of reactive oxygen species mediated by photosensitizers. These ROS can be radical species, such as hydroxyl anion and/or superoxide, or singlet oxygen, which are effective in the oxidative degradation of organic and biological pollutants. On the other hand, photoreductive transformations, which are usually concomitant to oxidation of water or additional organic molecules, can also be applied to detoxify heavy metal species.

Although the ability of separating the photosensitizer from solution is crucial in water treatment, heterogeneous systems have been scarcely explored in pollutant photodegradation.[13] A typical approach relayed on the study of TiO2-based materials, which usually require ultraviolet irradiation.[14] However, reticular chemistry, based on linking molecular building blocks by strong bonds to make open frameworks, offers many new possibilities on the discovery of new photoactive systems able to use visible-light to trigger predetermined processes. Particularly relevant are covalent organic frameworks, which are based in the atomically precise integration of pure organic units through covalent bonds to create predesigned skeletons and pores, constituting a main family of reticular porous materials.[15] COFs possesses a highly tunable and predictable pore size and structure, easily tailored functionality and versatile covalent combination of building blocks. Because they are composed of light-weight elements linked by strong covalent bonds, they have low mass densities and possess high thermal stabilities.[16] Furthermore, a wide variety of organic building blocks is available with a range of different organic groups that result on well stablished condensation reactions. Therefore, outstanding properties of COFs make them very attractive and prompted us to investigate their application into light-mediated decontamination of water.

In recent years, the application of covalent organic structures to photocatalytic processes has experienced a great blossoming.[17–20] As a specific application of this field, the degradation of pollutants has become one of the main focuses of interest, due to its social relevance. Although this area is rapidly reaching considerable maturity, important open questions remain. For example, structure-reactivity relationships in photoactive COFs are currently the subject of debate in current literature. In this review we present a critical description of the structural principles affecting the photocatalytic activity of COFs. In particular, conjugation and bandgap tuning of materials, together with donor-acceptor architectures, are reviewed as main strategies to modulate photocatalytic activities. In addition, understanding photocatalytic mechanisms is valuable information needed to rationally design improved photocatalytic systems. Thus, this review presents the current knowledge of the photocatalytic pathways determined for COFs. Finally, numerous examples of application of

COFs to the decontamination of pharmaceutical compounds, organic dyes, pesticides, phenolic compounds and heavy metal species are summarized in the final part of this manuscript. Considering the large amount of available literature, this review article is not intended to be an exhaustive compilation of published studies. Instead, we have selected the cited literature according to two main criteria: 1) We have given priority to the most recent reports and 2) We have chosen the most representative examples to illustrate the concepts presented in this manuscript. A systematic literature review, such as the one presented in this manuscript, should serve as a guide for future studies devoted to the design of new materials with photocatalytic properties useful for water decontamination.

2. Results

2.1. Structure-Activity Relationships

The photocatalytic efficiency of covalent organic frameworks (COFs) for water decontamination is strongly linked to their structural features. Crystal alignment, defect presence, and overall morphology determine their band structure, which, in turn, impacts their photocatalytic performance. These factors influence the modulation of COF optoelectronic properties, ensuring efficient charge and energy transfer processes, both essential for the photodegradation of water pollutants.

COFs are semiconducting materials formed by the valence band (VB), composed of filled atomic orbitals, and the conduction band (CB), formed by empty orbitals. These bands are separated by an energy gap (Eg), containing forbidden energy states. When irradiated with photons of higher energy than the COF band gap, electrons are excited from the VB to the CB. However, fast electron-hole recombination occurs in most semiconductors, leading to poorer optoelectronic properties and lower photocatalytic activity. Modifying semiconductor structure to trap or extract charge carriers is an established strategy for improving charge separation, and thus photocatalytic efficiency. This review focuses on three key strategies for optimizing COF structures and fine-tuning the COF optoelectronic properties, namely: 1) Incorporation of donor-acceptor units into the COF framework to improve charge separation; 2) Enhancement of π -conjugation across the COF framework; and 3) Modulation of the bandgap energy and COF band positions for improved photocatalysis.

2.1.1. Donor-Acceptor

Incorporating donor-acceptor (D-A) units into the backbone of photoactive COF is a powerful approach to enhance photocatalytic degradation of water pollutants. Pre-designing COFs containing alternating electron-rich donor and electron-deficient acceptor fragments broadens their optical absorption, while also promoting efficient separation and transfer of photogenerated electron-hole pairs.[21] This structure significantly reduces recombination, extending electron lifetimes and enabling the generation of reactive oxygen species (ROS), such as hydroxyl radicals and superoxide radical anions, which are critical for oxidizing and degrading organic pollutants in water.

The spatial arrangement of donor and acceptor units within COFs, often incorporating heteroatoms, is a critical factor for efficient charge transfer. For instance, the introduction of benzothiadiazol (acceptor) with halogen substituents and pyrene (donor) units within an imine-based COF has shown to suppress charge recombination and enhance water splitting (**Figure 1a**).[22] Li et al. synthesized an ethylene-linked COF (Py-BSZ-COF) also using pyrene and benzothiadiazole as donor and acceptor fragments (**Figure 1b**).[23] This combination resulted in a lower bandgap and faster charge separation, producing superoxide radicals under visible light for catalytic reactions, including oxidative amine coupling and thioamide cyclization to 1,2,4-thidiazole. Furthermore, incorporating benzothiadiazole as a building block extended the light absorption, leading to a redshift in the absorption edge when compared to equivalent COFs but without this unit.

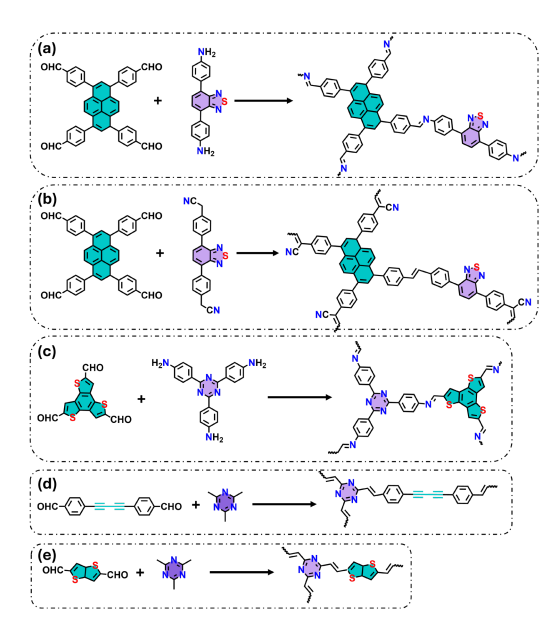


Figure 1. Examples of donor-acceptor precursors used to form new photoactive COFs.

Additionally, triazine-based COFs offers many structural advantages. The triazine moiety with a large number of nitrogen atoms which act as electron acceptor improves charge separation and ROS generation. For example, Qing et al. designed imine-linked COFs incorporating triazine (acceptor) and benzotrithiophene (donor), achieving spatially separated donor-acceptor centers (**Figure 1c**).[24] This D-A arrangement enhanced electron distribution within the COF framework and codirectional charge transfer, resulting in high H₂O₂ production. Similarly, Chen et al. designed vinylene-linked COFs combining triazine (acceptor) and diacetylene (donor) derivatives (**Figure 1d**), and demonstrated impressive photocatalytic degradation of phenol and norfloxacin by more than 96% in 15 minutes.[25] Deng et al. reported vinylene-linked COFs (TMT-TT-COF) with triazine and heteroatom-rich donor molecules (thieno [3,2-b]thiophene-2,5dicarboxaldehyde), creating narrow bandgaps and efficient charge transport (low recombination rates), resulting in high performance for photocatalytic H₂O₂ production and bisphenol A (BPA) degradation (**Figure 1e**).[26]

In summary, the donor-acceptor design in COF promotes long-range charge transport to targeted pollutants or oxygen molecules for generating ROS, key agents for the degradation of organic pollutants.

2.1.2. Conjugation Tunning

Increasing the degree of π -conjugation within COFs is another strategy to improve photocatalytic performance. Extensive π -conjugation in COFs allows better light absorption and faster charge transport, thereby enhancing charge transfer (from light-absorbing regions to reactive sites) while reducing recombination. Higher conjugation facilitates the delocalization of electrons across the COF framework, optimizing its electronic properties for photocatalysis.

The ability to fine-tune the degree of conjugation in COFs enables optimization for specific light sources and pollutant targets. Lin et al. synthesized two hydrazone-linked COFs with biphenyl (BPh–COF) or acetylene (AC–COF) to evaluate the effect of conjugation on photocatalytic efficiency (**Figure 2**).[27] The biphenyl ligands in the BPh–COF offer rigidity and strong conjugation. However, the ethynyl skeleton in the AC-COF enhances planarity and improves the π -conjugation. This significantly improved the optoelectronic properties of AC-COF compared with BPh-COF by narrowing the bandgap energy and improving charge transfer, ultimately enhancing photocatalytic efficiency for oxidative coupling amines. Furthermore, fully conjugated carbon-carbon double bonds in vinylene-linked COFs' prevent spatial distortions, leading to a more efficient conjugated structure.

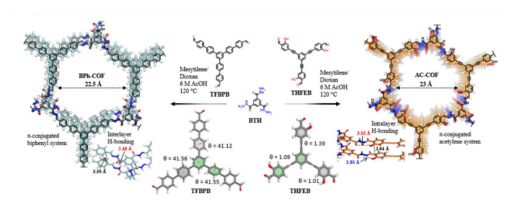


Figure 2. Example of extended π -conjugated COFs (adapted from [27]).

Additionally, Deng et al. presented a post-synthetic modification of imine COFs by sulfurization, leading to thiazole-linked frameworks with enhanced π - conjugation.[28] The resulting sulfurized COFs exhibited superior charge transport, increasing the photocatalytic H₂O₂ production when applied in oxygen reduction reactions (ORR) by nearly six times higher than pristine COF. This work highlights the importance of conjugation tunning and details a post-synthetic sulfurization for improving the local conjugation of imine-linked COFs and therefore, their photocatalytic activity.

2.1.3. Band-Gap Modulation and Band Positioning for Optimal Photocatalysis

Modulating the COF bandgap and positioning the VB and CB are key strategies for optimizing electron transfer and maximizing photocatalytic efficiency. A narrower bandgap allows COFs to absorb more visible light, making them ideal for sunlight-driven photocatalysis in water remediation. Additionally, aligning VB and CB of the COFs with the redox potentials of pollutants enables more effective electron-hole, increasing photocatalytic degradation rates.

Chen et al. synthesized a family of hydrazide-based COFs modulating its band structure by incorporating different linkers.[29] They demonstrated improved bandgap tunning and charge transfer through intersystem crossing (ISC) from the S₁ excited state to the triplet state (T₁) under visible light. This boosted remarkable photocatalytic performance in applications like uranium extraction from water sources. However, it is well known that the photocatalytic efficiency of single-component catalysts is limited by the photocarrier recombination.[30] COF-based heterojunction has been reported as an alternative for improving band-bending. When COFs are combined with other semiconductors and form a heterojunction, band-bending occurs depending on the type of majority charge carriers (electrons or holes). In the literature, X-ray and Ultraviolet Photoelectron Spectroscopy (XPS and UPS) has been employed to determine Fermi level (*E_F*, energy level with the

50% probability of being occupied by an electron) position of different photocatalysts, which enables to predict their band bending. [31,32] Furthermore, band bending also occurs when a semiconductor is in contact with a wastewater solution. Electrons will flow until the Fermi level of the solid is equalized with the redox potential of the pollutant (E_{redox}), and the energy barrier formed on the solid-liquid interface is denoted as flat band potential (V_{fb}). This parameter can be experimentally estimated from electrochemical characterization by measuring the capacity of the semiconductor-electrolyte interface as a function of applied potential, known as Mott-Schottky plot.

Many examples of engineering band-bending in COFs have been reported to enhance charge carrier lifetime and interaction with pollutants, ensuring more efficient photocatalytic degradation. For example, covalent bonding between COFs and MOF has appeared as a promising research direction to enhance the degradation of organic pollutants. For instance, He et al. prepared a COF/MOF composite (NH₂-MIL-125 (Ti)/TTB-TTA) and, based on the valence band XPS spectra, they estimated the energy position of the VB and CB of the composite materials and the individual counterparts.[33] Yang et al. also adopts this strategy and combines Ti-MOF and COF with triazine moieties (**Figure 3a**).[34] Furthermore, they used photocurrent and electrochemical impedance spectroscopy (EIS) measurements and demonstrated an effective charge transfer in the heterojunction. This difficulted recombination and enabled an efficient photocatalytic degradation of bisphenol A.

Other examples of COF-based heterojunctions are their combination with well-known inorganic photocatalysts, solving the disadvantages of wide band gap and high electron-hole recombination rate of the inorganic semiconductors. For example, Liu et al. prepared Fe–TiO2@COF with exceptional photodegradation efficiency for methylene blue.[35] Furthermore, Liu et al. synthesized a COF-PD/AgI composite which showed excellent degradation of Rhodamine B and acetaminophen (Figure 3b).[36] Sun et al. constructed CdS/COF composite for the photodegradation of bisphenol A (BPA).[37] More sophisticated systems were reported by immobilization of electron mediator ([Cp*Rh(bpy)H₂O]²⁺) on COFs supported on silica matrix (Figure 3c).[38] In addition, the combination of COFs with visible light-responsive semiconductor photocatalyst graphite carbonitride (g-C₃N₄) has recently attracted great attention in the photocatalytic degradation of pollutants (Figure 3d). For instance, the heterojunction formed in composites g-C₃N₄@COF has been proven effective in enhancing charge mobility and reducing electron-hole recombination rates.[39–41] These composites exhibit high photocatalytic efficiency toward the degradation of organic pollutants due to the optimal balance between nitrogen content and graphitization degree.

Overall, COFs present a highly tunable structural platform for enhancing photocatalytic water decontamination. Structural modifications, particularly donor-acceptor integration, π -conjugation tuning, and modulation of bandgap and band position, are key strategies for improving photocatalytic performance. While challenges remain, such as minimizing charge recombination and maximizing light absorption, advanced spectroscopic techniques aid in determining the bandgap energy, positions of VB and CB, and charge mobility. This knowledge will be critical to establishing the best COF structures for degrading specific water pollutants with ongoing advancements in COF synthesis and functionalization, their potential for environmental remediation is expected to expand, offering green solutions for degrading persistent water pollutants.

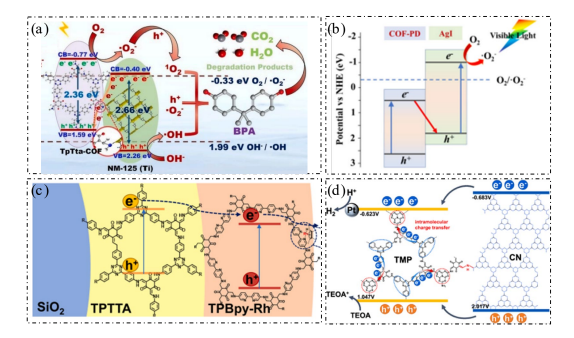


Figure 3. Examples of COF-based heterojunctions used as photocatalytic systems (adapted from [34,36,38,41]).

2.2. General Photocatalytic Mechanisms

Photocatalysis generally implies the absorption of a photon by a photocatalyst, leading to its excited electronic state. In this process, the photon energy (hv) must be equal to or greater than the energy gap between the valence and conduction bands (or the HOMO-LUMO gap in the case of molecular photocatalysts). Upon absorption, an electron is excited from the lowest occupied energy level to a higher energy state in the conduction band, increasing its reactivity.[42] The excited photocatalyst can return to its ground state through two primary mechanisms: Single Electron Transfer (SET) and Energy Transfer (EnT). In SET, an electron exchange occurs between the photocatalyst in its excited state and a substrate (such as a pollutant) or a cocatalyst within the reaction system.[43] In contrast, EnT involves the transfer the absorbed photon energy from the photoexcited catalyst to a substrate or cocatalyst via radiative or non-radiative way, inducing a chemical transformation.[44] Once this process concludes, the excited electron returns to the ground state (Figure 4).

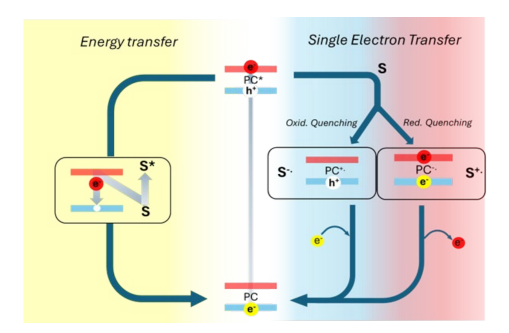


Figure 4. Schematic representation of energy transfer processes and singlet electron transfer process.

2.2.1. Electron Transfer: Reductive Quenching vs Oxidative Quenching

Electron transfer between the photocatalyst and a substrate (pollutant) or cocatalyst can proceed via two distinct pathways, depending on the redox potentials and the relative energy levels of the conduction and valence bands of the photocatalyst compared to the HOMO-LUMO levels of the substrate.[45]

One a one hand, if the reduction potential of the photocatalyst is higher than that of the substrate ($E^{S}_{red} < E^{PC}_{red}$), oxidative quenching occurs. This process is favored when the conduction band of the photocatalyst is energetically higher than the LUMO of the substrate. Oxidative quenching results in substrate reduction, generating a radical anion(S^{-}) and a radical cation of the photocatalyst (PC^{+}). The radical cation is subsequently reduced by a regenerator (R), closing the catalytic cycle. This regenerator may be a Sacrificial Electron Donnor (SED), or another compound involved in the mechanism (**Figure 5**). On the other hand, if the reduction potential of the substrate is higher than that of the photocatalyst ($E^{PC}_{red} < E^{S}_{red}$), reductive quenching is favored. In this case, electron transfer occurs from the substrate to the photocatalyst, generating an oxidized substrate intermediate (S^{+}) and a radical anion of the photocatalyst (PC^{-}). The radical anion then transfers its extra electron to the regenerator, completing the catalytic cycle (**Figure 5**).

These electron transfer pathways are very common in various photocatalytic transformations, where radical species act as key intermediates. The reactivity of these species depends on their lifetimes, while their selectivity is influenced by the reactive oxygen species (ROS) involved. Herein we describe some of the most frequent radical species involved in photocatalytic transformations.

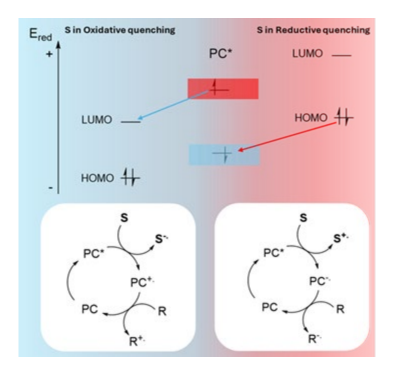


Figure 5. Schematic representation of oxidative and reductive quenching.

2.2.2. Generation of Radical Species: Superoxide, Hydroxyl, Organic Radicals

Radical species are compounds characterized by an unpaired electron. They commonly arise from homolytic cleavage of a covalent bond or, through SET-mediated oxidative or reductive quenching mechanisms involving the photocatalyst. These species generally exhibit high reactivity due to their inherent instability. In this section, we will review some of the most frequent radicals involved in photocatalytic oxidation processes related to organic photoactive materials.

Superoxide radical anion ($\cdot O_2$) is formed through oxidative quenching of molecular oxygen with the excited state of a photocatalyst. This radical results from electron transfer from the CB of the photocatalyst to the antibonding π orbital of triplet oxygen. Superoxide radicals are relatively stable due to their extended lifetime. The nature of this radical anion is predominantly oxidative (E_{red} ($\cdot O_2$ -/ H_2O_2) = 0.93 V) but can also act as a reductant, presenting in a protic media and E_{ox} (O_2 / $\cdot O_2$)= 0.33 V.[46] In fact, these radicals plays a crucial role in photocatalytic oxidative couplings and pollutant degradation. For example, the oxidative role of superoxide radical anion is commonly observed in photocatalytic oxidation of benzylic amines to form amine radical cations in oxidative couplings of amines[47] or in photocatalytic aerobic degradation of pollutants such as of methylene blue or Rhodamine B dyes.[48] Additionally, superoxide radical anion can be further oxidized to form singlet oxygen, or protonated (see **Scheme 1**) to form peroxyl radical ($\cdot OOH$).[49,50]

Hydroxyl radical(·OH) is a highly oxidant radical specie generated by the reduction of hydrogen peroxide via *Fenton*-type reactions or by the oxidation of hydroxyl anions. These species are commonly observed in metal oxide-based photocatalysis such as TiO₂.[51–53] However, only a few examples exist of COF materials with metal centers able to produce these hydroxyl radicals. In this sense, some composites with metal like Pd or Ni have been reported in oxidations promoted by hydroxyl radicals.[54,55]

Probably, the scarcity of examples of pure organic materials able to generate hydroxyl radical could be attributed to their lower charge mobility compared with metal-based photocatalysts, which restrict the diffusion of charge carriers to active adsorption sites on the material surface. Consequently, the homolytic cleavage of the O-H bond cleavage from water, hydrogen peroxide and hydroxyl anion species are difficulted.

Scheme 1. Scheme of the evolution of superoxide radical anion to form other oxidant radical species.

2.2.3. Energy Transfer: Singlet Oxygen and Direct Energy Transfer

Chemical transformations can also be promoted through direct energy transfer from the photocatalyst to the substrate, primarily via two different mechanisms, depending the type of excited state reached (**Figure 6**). In general, after the absorption of a photon, the photocatalyst reaches a primary excited state from which a slight energy decay attributed to vibrations and thermic loss of energy takes place. This process drives the PC* to a singlet excited state, which can furtherly evolve to a triplet excited state through Inter-System Crossing (ISC). Depending on the population of either singlet or triplet excited states, we can consider Förster resonance energy transfer (FRET) and Dexter energy transfer mechanisms.

Förster's mechanism implies long-range dipole-dipole (Coulombic) interactions from the excited photocatalyst to the substrate. The efficiency of this energy transfer depends on the relative geometric orientation and distance between interacting species. Since heterogeneous photocatalysis occurs at the catalyst's surface, proximity between the substrate and the catalyst active sites is crucial. Different parameters altering the substrate interaction with the photocatalyst, like non-covalent interactions, would have significant influences on the substrate adsorption and would impact energy transfer efficiency.

The Dexter mechanism relies on a short-range electron exchange between the excited photocatalyst and the substrate. Unlike SET, Dexter energy transfer mechanism does not produce charged intermediates and maintains a charge neutrality. This phenomenon requires orbital overlap between the donor (photocatalyst) and the acceptor (substrate), involving collisional interactions. For the viability of this mechanism, short distances (order of Angstroms) are required. In the case of a heterogeneous photocatalyst, an overlap between the "acceptor" substrate orbital and the "donor" band of the material would be required for an efficient energy transfer.

Finally, a well-known example of Energy Transfer activations is the conversion of triplet oxygen (3O_2) into singlet oxygen (1O_2), a highly reactive oxidant. This transformation has been widely used in photocatalyzed oxidation reactions involving heterogeneous photocatalysts.[56–58] However, the promotion of oxygen from its triplet to singlet excited state is not always favored. Efficient singlet oxygen generation depends on the photocatalyst's ability to undergo intersystem crossing (ISC) to a triplet state. If rapid fluorescence deactivation from the photocatalyst's singlet excited state occurs, the energy transfer process for singlet oxygen formation may be inefficient. Thus, photocatalysts with extended exciton lifetimes are more suitable candidates for singlet oxygen generation via energy transfer.[59]

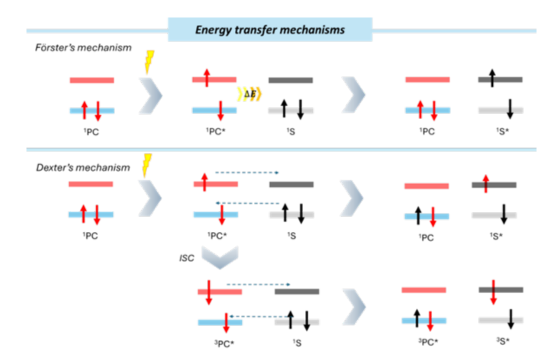


Figure 6. Schematic representation of energy transfer mechanisms.

2.3. Pollutants Tackled

Covalent organic frameworks offer a promising solution for water decontamination through the photocatalytic degradation of major pollutants, including pharmaceutical compounds, organic dyes, toxic metals, pesticides and phenolic compounds. **Table 1** summarizes various COF-based photocatalyst capable of degrading common pharmaceutical contaminants such as the analgesic acetaminophen and three antibiotics: tetracycline, sulfamethazine and sulfamethoxazole (**Figure 7**).

Table 1. COF-based photocatalysts for degradation of pharmaceutical compounds in water.

Pollutant	COF name (linkage type)	Design principle	ROS involved	Ref.
	COF-R (imine)	Donor-acceptor COFs	\cdot O ₂ - / \cdot OH	[60]
Tetracycline	sp ² c-COF (olefin)	Extended conjugation of COFs	·OH	[61]
	MoS ₂ /COF (imine)	MoS ₂ /COF heterojunction	·OH / ·O ₂ -	[62]
	COF-TD1 (imine)	Donor-acceptor COFs	·O ₂ -	[63]
Acetaminophe	COF-PD/AgI (imine)	COF/AgI heterojunction	·O ₂ -	[36]
n	Heptazine-COF@TiO ₂ (imine)	COF/TiO ₂ heterojunction	·O2 ⁻	[64]
Sulfamethazin e	SQ-COF-1 (imine)	Extended conjugation of COFs	·O ₂ -	[65]
	COF/CN-3 (C-N bond)	COF/ g-C ₃ N ₄ heterojunction	·O ₂ - / ¹O ₂ / ·OH	[66]
Sulfamethoxaz ole	COF-909(Cu) (imine)	COF(Cu)	·O ₂ -/·OH	[67]

Figure 7. Chemical structures of pharmaceutical compounds commonly found as water pollutants.

Tetracycline (TC) photodegradation has been explored through various synthetic strategies, including donor-acceptor COFs, extended π -conjugated COFs and inorganic/COF heterojunctions. One approach involves imine-linked COF (COF-R) functionalized with electron-withdrawing (-OH and -OMe) and electron-donating (-Cl, -Br) substituents (**Figure 8a**). Mechanistic studies indicated that TC degradation primarily occurs through direct oxidation by photogenerated holes, with ROS, particularly \cdot O₂- and \cdot OH, also contributing to the TC photodegradation process. The electron-withdrawing groups favor direct oxidation by lowering the valence band energy, whereas electron-donating groups enhances ROS generation. [60]

Another strategy involves the development of olefin-linked COFs with extended π -conjugation, such as sp²c-COF, which facilitates the delocalization of photogenerated electrons and promotes TC direct oxidation by holes (**Figure 8b**).[61] Electron paramagnetic resonance (EPR) analysis confirms the generation of both ·OH and ·O2⁻, although scavenger studies indicate that only ·OH plays a significant role in TC photodegradation. Additionally, semiconductor/COF heterojunctions, such as MoS2/COF, have been employed for TC photodegradation (**Figure 8c**). [62] In this system, photogenerated electrons in the COF conduction band (CB) transfer to the MoS2 CB, while holes oxidize TC directly. The electrons in MoS2 react with molecular O2 to generate ·O2⁻, which subsequently forms ·OH via interaction with H2O, enhancing photocatalytic performance.

For acetaminophen (ACTP) degradation, donor-acceptor COFs have been designed using thiadiazole (donor) and quinone (acceptor) fragments, as COF-TD1 (**Figure 8d**).[63] These fragments facilitate charge separation by localizing photogenerated electrons at the acceptor centers and holes at the donor centers, reducing recombination. Mechanistic studies reveal that while direct oxidation by holes is possible, ·O₂- is the primary oxidant species. COF-based heterojunctions have also been investigated for ACTP photodegradation. For example, Liu et al. developed a COF-PD/AgI heterojunction that primary generates ·O₂-, which oxidize the acetaminophen molecules (**Figure 8e**).[36] Through hydroxyl radicals were detected by EPR, they are inactive in ACTP degradation but contribute to other photocatalytic processes such as bacteria degradation. Direct oxidation by holes was negligible. Similarly, Ge et al. reported a Heptazine-COF@TiO₂ heterojunction for photocatalytic degradation of dyes and pharmaceuticals (**Figure 8f**).[64] Under visible light irradiation, photogenerated electrons in the COF generate ·O₂- or are transferred to the TiO₂ CB, further producing ·O₂-, which plays a crucial role in ACTP degradation.

Photodegradation of other antibiotics, such as sulfamethazine (SMT) and sulfametathoxazole (SMX), have been also included in **Table 1**. Ben et al. synthesized extended π -conjugated COFs via imine condensation of squaric acid (SQ) with 1,3,5-tris(4-aminophenyl)benzene (TAPB), forming SQ-COF-1 (**Figure 8g**).[65] The SQ center possesses a more negative charge density than the TAPB fragment, which is mainly positive. Upon illumination, photogenerated electrons localize on SQ, while holes remain on TAPB fragments, increasing their lifetime. Scavenger studies indicate that SMT photodegradation proceeds through \cdot O₂- oxidation. Furthermore, Qi et al. studied SMT degradation using a COF-based hybrid material, combining PC-COF with g-C₃N₄, forming the COF/CN-3 photocatalyst (**Figure 8h**).[66] Under irradiation, COF CB electrons react with molecular O₂ to generate \cdot O₂-, which further evolves into 1 O₂, both serving as primary reactive species in SMT photodegradation. Additionally, SO₄-· and ·OH radicals were identified as photocatalytic intermediates. Finally, for sulfamethoxazole (SMX) degradation, **Table 1** includes an N-rich COF coordinated with Cu, designated COF-909(Cu) (**Figure 8i**).[67] Cu coordination with COF nitrogen

atoms reduces the bandgap energy and shifts the energy band positions, enhancing photocatalytic performance. Mechanistic studies employing EPR detected \cdot OH and \cdot O₂ species. However, a scavenger study is lacking to determine the relative significance of these ROS species in SMX degradation.

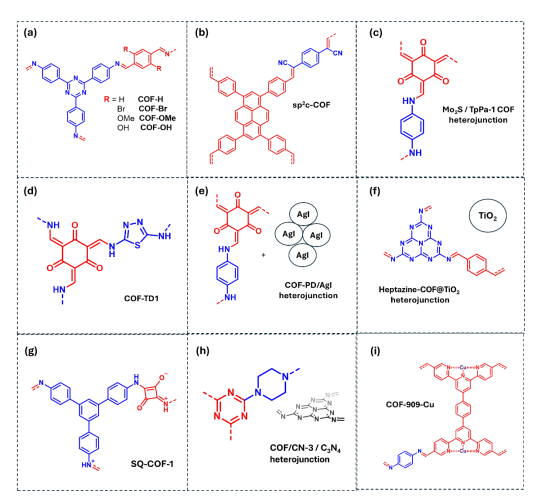


Figure 8. COF-based materials employed as photocatalysts for the degradation of pharmaceutics in water.

These studies highlight the versatility of COFs in photocatalytic water decontamination, demonstrating their efficiency in degrading pharmaceutical pollutants through various photocatalytic mechanisms, including direct oxidation and ROS-mediated degradation.

Another important group of toxic substances commonly found in water are heavy metals such as uranium U(VI) and chromium Cr(VI), which can be reduced to their non-toxic or less toxic forms, U(IV) and Cr(III), respectively, using COF-based materials (**Table 2**). In the case of uranium degradation, Zhang et al. reported a β -keto-enamine COFs (TpTt-COF) rich in heteroatoms and containing two electron-acceptor fragments (triazine and ketones) in close proximity (**Figure 9a**).[68] These structural features enhances planarity and extend π -conjugation of the COFs, improving the efficiency of photocarrier separation and boosting photocatalytic activity. This COF generates \cdot O₂ and \cdot O₂ radicals, where \cdot O₂ actively reduces U(VI) to insoluble U(IV) species such as UO₂, UO_{2.9}, UO_{2.87} and UO_{2.82}. However, \cdot O₂ can promote the reoxidation of U(IV) to U(VI), which should be minimized by adding scavengers of \cdot O₂.

Table 2. COF-based photocatalysts for reduction of U(VI) and Cr(VI) in water.

Pollutant	COF name (linkage type)	Design principle	ROS involved	Ref.
U(VI)	TpTt-COF (imine)	Donor-acceptor COFs	·O ₂ -	[68]
	DQTP-COF (imine)	Donor-acceptor COFs	·O ₂ -	[69]

	COF-AQ (imine)	Donor-acceptor COFs	-	[70]
	COF-4-Pd-AO (imine)	COF/Pd heterojunction	-	[71]
Cr(VI) —	COFs-OMe (imine)	Donor-acceptor COFs	·O2-	[72]
	TPB-BT-COF (imine)	Donor-acceptor COFs	-	[73]
	HDU-26-COF (imine)	Donor-acceptor COFs	·O ₂ -	[74]
	TAPP-2F-COF (imine)	Halogenated COFs	·O ₂ -	[75]

Furthermore, Ling et al. presented a COF containing anthraquinone units, which play a crucial role in charge separation and serve as active sites for U(VI) reduction (Figure 9b).[69] Experimental results identified ·O₂ as the dominant reactive species, while direct electron transfer also contributed to the U(VI) reduction process. The same COF material was further studied for various photocatalytic processes by Huang et al. who designated it as COF-AQ (Figure 9c).[70] In this study, EPR spectroscopy confirmed the generation of ROS species, and scavenger experiments using silver nitrate as an electron quencher demonstrated that the primary photocatalytic reduction mechanism for U(VI) using the COF-AQ involved direct electron transfer. Additionally, Hao et al. designed a COF by incorporating two amine precursors with bipyridine and diciano-biphenyl moieties and 4,4′,4″-(1,3,5-triazine-2,4,6-triyl)tribenzaldehyde (TATTA).[71] The resulting COF was subsequently functionalized with PdCl₂ to produce COF-4-Pd-AO, where Pd centers coordinated to bipyridine units act as active sites for adsorption and photocatalytic uranium reduction (Figure 9d). Mechanistic studies of this COF/Pd heterojunctions concluded that U(VI) species undergo direct electron transfer reduction to UO₂.

Regarding the photoreduction of Cr(VI) species , Zhang et al. investigated the effect of electron-withdrawing and electron-donating substituents in COFs, revealing that methoxy-functionalized COFs (COFs-OMe) exhibit the highest photocatalytic efficiency for Cr(VI) reduction (**Figure 9e**).[72] Mechanistic studies demonstrated that Cr(VI) photoreduction occurs through both direct electron transfer and ·O₂ generation, which actively participates in the reduction process. Furthermore, Chen et al. reported a benzodithiazole-containing COF (TPB-BT-COF) with exceptional Cr(VI) photoreduction to Cr(III) species, achieving over 99% efficiency without the need for sacrificial agents or pH adjustment (**Figure 9f**). Additionally, Lu et al. developed a COF containing thiophene and benzodithiazole units (HDU-6-COF) (**Figure 9g**).[74] Their mechanistic study confirmed that Cr(VI) reduction proceeds via direct electron transfer and ·O₂ involvement in the photocatalytic process.

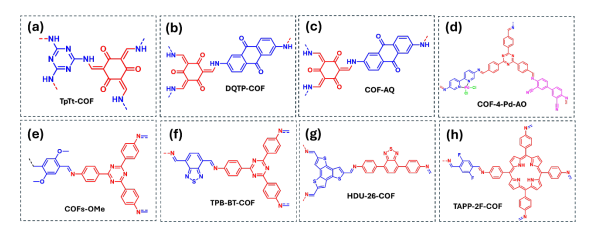


Figure 9. COF-based materials employed as photocatalysts for the reduction of U(VI) and Cr(VI) in water.

Organic dyes have become significant water pollutants due to their widespread use in industries such as food processing or textiles. **Table 3** presents various strategies of designing COF-based photocatalysts to remove these pollutants (**Figure 10**), particularly rhodamine B (RhB), methyl orange (MO), methylene blue (MB), mordant black 17 (MB17), Eriochrome black A and T (EBA and EBT), rose bengal (RB) and methyl green (MG).

Table 3. COF-based photocatalysts for degradation of organic dyes in water.

Pollutant	COF name (linkage type)	Design principle	ROS involved	Ref.
	COF-PD/AgI (imine)	COF/AgI heterojunction	·O ₂ -	[36]
Rhodamine B	MoS ₂ /COF (imine)	MoS ₂ /COF heterojunction	·O ₂ -/·OH	[62]
Kilouallille b	COF-HFeTBD (imine)	Donor-acceptor COFs	$^{1}O_{2}$	[76]
	HDU-105-COF (olefin)	Donor-acceptor COFs	·O ₂ - / ·OH	[48]
	HDU-105-COF (olefin)	Donor-acceptor COFs	·O ₂ - / ·OH	[48]
Methylene blue	C ₆ -TRZ-TPA COF (imine)	Donor-acceptor COFs	·O ₂ - / ·OH	[77]
	TTO-COF (olefin)	Extended conjugation of COFs	•O2 ⁻	[78]
Methyl Orange	TTO-COF (olefin)	Extended conjugation of COFs	·O ₂ -	[78]
	COF _{A+C} (imine)	N-rich COFs	·O ₂ - / ·OH	[79]
Rose Bengal	C ₆ -TRZ-TPA COF (imine)	Donor-acceptor COFs	·O ₂ -/·OH	[77]
Mordant black 17	TpBD-COF (imine)	Pure COF	$\cdot O_{2^{-}} / {}^{1}O_{2} / \cdot OH$	[80]
Eriochrome black	TpBD-COF (imine)	Pure COF	$\cdot O_2$ - / 1O_2 / $\cdot OH$	[80]
T	TFA-TTA-COF (imine)	Fluorinated COF	·O ₂ - / ¹O ₂	[81]
Eriochrome black A	TFA-TTA-COF (imine)	Fluorinated COF	·O ₂ - / ¹O ₂	[81]
Methyl green	CuO/COF (imine)	CuO/COF heterojunction	·O ₂ -	[82]

Figure 10. Chemical structures of common dyes pollutant in water.

Regarding rhodamine B, two COF-based heterojunctions (COF-PD/AgI and MoS₂/COF) previously discussed in **Table 1**, have also been explored for its photocatalytic degradation (**Figure 11a,b**).[36,62] In both cases, ·O₂ is the primary reactive species responsible for degradation, while in the MoS₂/COF system, ·OH radicals also participate in the RhB photodegradation. Additionally, Liu et al. reported a D-A COF incorporating porphyrin units coordinated with Fe centers (COF-HFeTBD) (**Figure 11c**).[76] Mechanistic studies reveals that the active species in this system is singlet oxygen species (¹O₂), generated via oxidation of ·O₂, although ·O₂ itself is not directly involved in the RhB oxidation.

In another approach, Wang et al. investigated a D-A olefin-linked COF (HDU-105-COF) for the photocatalytic degradation of RhB and MB (**Figure 11d**).[48] Mechanistic studies indicated that both \cdot O₂- and \cdot OH species contribute to the process, with \cdot O₂- playing a more significant role in RhB

degradation than in MB degradation. Similarly, Ruidas et al. reported a D-A imine-linked COF (C6-TRZ-TPA COF) capable of degrading MB and RB (**Figure 11e**).[77] This COF generates both ·O2 and ·OH radicals, which actively participate in the photodegradation of both dyes. Furthermore, Yang et al. designed an extended π -conjugated olefine-COF containing triazine units (TTO-COF) for the photocatalytic degradation of MB and MO (**Figure 11f**).[78] In this case, the degradation of both dyes is mainly driven by the generation of ·O2 radicals. Another example of MO degradation was reported by He et al., who developed an imine-linked, N-rich COF (COF_{A+C}) incorporating 4,4′,4″-(1,3,5-triazine-2,4,6-triyl) units (**Figure 11g**).[79] This material generates both ·O2 and ·OH radicals, which are responsible of the MO degradation (**Figure 16**).

Table 3 also highlights the use of COFs as photocatalyst for the degradation of other emerging dye pollutants. For example, Xue et al. synthesized an imine-linked COF (TpBD-COF) for the photodegradation of MB17 and EBT (Figure 11h).[80] Scavenger studies on MB17 photodegradation revealed that ·O₂ is the primary reactive species under visible light, while ¹O₂ and ·OH radicals also contribute but to a lower extent. Additionally, Qi. et al. developed an imine-linked, fluorinated 2D COF (TFA-TTA-COF) for the photocatalytic degradation of EBT and EBA (Figure 11i).[81] The dominant reactive species in the degradation of both dyes is ·O₂⁻, although ¹O₂ also plays a role in the process. Finally, Wang et al. investigated the photocatalytic degradation of MG using a CuO/COF heterojunction (Figure 11j).[82] Mechanistic studies demonstrated that ·O₂⁻ is the main ROS, while direct oxidation by photogenerated holes is also favored in this heterojunction system.

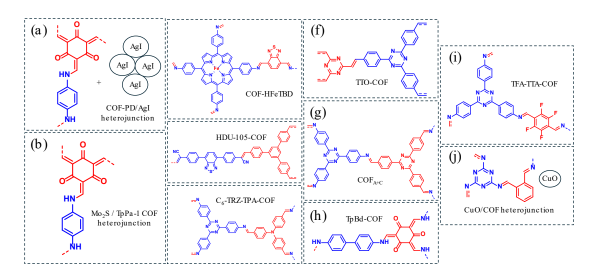


Figure 11. COF-based materials employed as photocatalysts for the degradation of dyes in water.

Pesticides are widely used in agriculture to protect plants from pesticides, however, their persistence and toxicity make them a significant water pollutant, leading to serious environmental concerns. **Table 4** presents examples of COF-based materials employed for the photocatalytic degradation of pesticides such as diazinon, parathion, imidacloprid and pymetrozine (**Figure 12**).

Table 4. COF-based photocatalysts for degradation of pesticides in water.

Pollutant	COF name (linkage type)	Design principle	ROS involved	Ref.
Diazinon and parathion	PS@COF-366 (imine)	Sulphured COF	$\cdot O_{2^{-}} / {}^{1}O_{2} / \cdot OH$	[83]
Imidacloprid	Fe ₃ O ₄ @HMN-COF	Fe ₃ O ₄ /COF		[84]
midacioprid	(imine)	heterojunction	-	[04]
Drymotrozino	Ag ₃ PO ₄ /TpPa-1-COF	Ag ₃ PO ₄ /COF	·O ₂ - / ·OH	[05]
Pymetrozine	(imine)	heterojunction	·O2 / ·OH	[85]

Figure 12. Chemical structures of common pesticides water pollutants.

Among them, pesticides such as diazinon and parathion have been effectively degraded using a sulfur-functionalized porphyrin-based COF, denoted as PS@COF-366 (**Figure 13a**).[83] in this system, the porphyrin units act as photoantennas, facilitating photocarrier separation and enhancing photocatalytic activity. The degradation mechanism involves multiple ROS species, including ·O₂, ¹O₂ and ·OH, with ·O₂⁻, playing the dominant role in the photodegradation process. Additionally, the polysulfide chains present in this COF assist in pesticide photodegradation by cleaving P-O bonds. Another example highlighted in **Table 4**-is the photocatalytic degradation of imidacloprid using an Fe₃O₄/COF heterojunction (Fe₃O₄@HMN-COF) (**Figure 13b**).[84] This system achieved high photocatalytic efficiency (> 98%) under optimized conditions. However, this study did not include a mechanistic investigation to identify the reactive species involved in the degradation process. Finally, a Ag₃PO₄/COF heterojunction (Ag₃PO₄/TpPa-1-COF) has been reported for the photocatalytic degradation of pymetrozine (**Figure 13c**).[85] Mechanistic studies revealed that the degradation process proceeds via the generation of ROS, specifically ·O₂⁻ and ·OH, in addition to a direct electron transfer to the substrate, further enhancing the photocatalytic performance.

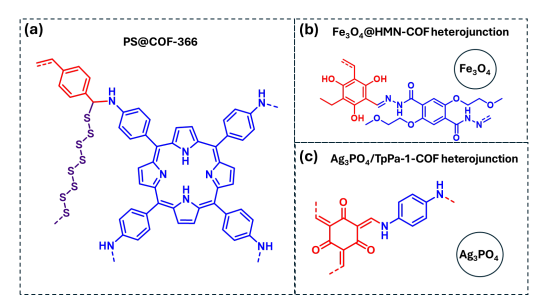


Figure 13. COF-based materials employed as photocatalysts for the degradation of pesticides in water.

Phenolic compounds are another major class of water pollutants, primarily originating from plastic industry, and pose a significant environmental threat due to their toxicity and persistence. **Table 5** presents recent studies utilizing COF-based materials as photocatalysts for the degradation of these pollutants, particularly phenol, bisphenol A and 4-chlorophenol (**Figure 14**).

Table 5. COF-based photocatalysts for degradation of phenolic compounds in water.

Pollutant	COF name (linkage type)	Design principle	ROS involved	Ref.
Phenol	COF _{A+C} (imine)	N-rich COFs	$\cdot O_{2^{-}} / \cdot OH$	[79]

	GCN/TD-COF (imine)	GCN/COF heterojunction	·O2 ⁻ / ·OH	[86]
	CdS/TFp-Azo-COF	CdS/COF	·O ₂ - / ·OH	[87]
	(imine)	heterojunction	02 / 011	
Bisphenol-A	NM-125(Ti) _{0.4} @TpTta-	MOF/COF	·O ₂ - / ¹O ₂ / ·OH	[34]
	COF (imine)	heterojunction	.02 / .02/ .011	
	MIL-101-NH ₂ @COF-	MOF/COF	ЮН	[00]
	TpMA (imine)	heterojunction	.011	[88]
4-chlorophenol	D:/TD DDDA COE (imina)	Bi/COF	O / OH	[89]
	bi/1r-bDDA-COF (imine)	heterojunction	·O ₂ - / ·OH	

Figure 14. Chemical structures of common phenolic compounds water pollutants.

The imine-linked, N-rich COF_{A+C} previously discussed in **Table 3** and reported by He et al. was also investigated for the photodegradation of phenolic compounds (**Figure 15a**).[79] Mechanistic studies suggest that the photodegradation process is mainly driven by superoxide radicals (·O₂·), with hydroxyl radicals (·OH) playing a supporting role. Similarly, a graphitic carbon nitride/COF heterojunction system (GCN/TD-COF) demonstrated effective phenol degradation via mechanism involving ·O₂· and ·OH as the main reactive species (**Figure 15b**).[86] Beyond phenol degradation, different COF-based heterojunctions have been explored for the removal of bisphenol A, a widely used industrial compounds of high environmental concern (**Figure 26**). For instance, Sun et al. reported a CdS/COF heterojunction (CdS/TFp-Azo-COF) that achieved high photocatalytic efficiency for bisphenol A degradation (**Figure 15c**).[87] Mechanistic investigations identified ·O₂· as the dominant ROS, while ·OH played a secondary role in the photodegradation process.

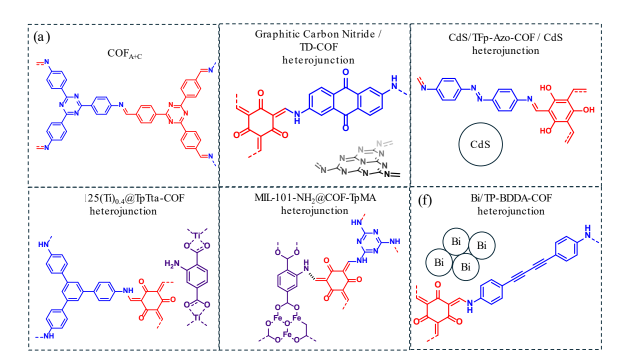


Figure 15. COF-based materials employed as photocatalysts for the degradation of phenolic compounds in water.

In addition, MOF/COF heterojunctions have been developed to enhance photocatalytic degradation. Yang et al. combined a Ti-based MOF with the TpTta-COF, forming the hybrid NM-125(Ti)_{0.4}@TpTta-COF (**Figure 15d**) .[34] Scavenger experiments revealed that the degradation of bisphenol A occurs through the generation of multiple ROS, including ·O₂-, ·IO₂ and ·OH. Lv et al. also explored a COF/MOF hybrid formed by combination of MIL-101-NH₂ and COF-TpMA and forming MIL-101-NH₂@COF-TpMA (**Figure 15e**).[88] This hybrid system demonstrated that bisphenol A degradation was mainly mediated by ·OH, with SO₄-· also participating in the oxidation process. Finally, the last example of **Table 5** consists in a Bi/COF heterojunction (Bi/TP-BDDA-COF) reported as an efficient photocatalyst for the degradation of 4-chlorophenol, with ·O₂- and ·OH identified as the key reactive species driving the process (**Figure 15f**).[89]

3. Concluding Remarks

In general, the review of the available literature presented in this manuscript highlights the enormous potential of COFs for photocatalytic water decontamination applications. The growing plethora of available building blocks, coupled with constant improvements in synthetic processes and in the design of covalent organic structures, make these materials a valuable target for environmental applications. The stablished mechanistic landscape and the dependence of the pathways followed on the structural features of materials resulted in stabilizing several structurereactivity relationships. In this review we highlighted the role of extended conjugation, the presence of donor-acceptor moieties and the strategies to modulate the optical bandgap as main tools to direct the photocatalytic activity of COFs. Such state of the art makes it foreseeable that new COF-type materials with improved activities for a wide variety of pollutants will emerge. Furthermore, dual photocatalytic systems are appearing in the literature, in which one pollutant is simultaneously oxidized and another is reduced, which promise to be an effective alternative for decontaminating water with pollutants of various kinds. Despite the wide range of pollutants tackled by photocatalytic systems based on COFs, some aspects related to water decontamination remain elusive. Specialty relevant challenge consists on micro/nano-plastic particles, which could be an important subject of future research.

The processes achieved by photocatalytic COFs constitute a very promising field of research of undoubted environmental interest. However, some aspects inherent to the nature of COF materials constitute major drawbacks for translating the basic research reported here into practical applications. First, water decontamination requires large-scale production of active materials, which should involve low-cost synthesis. At present, most photoactive COFs are constituted by high added-value organic building blocks. Therefore, the use of widely available organic structures, such as natural products, should be explored. In addition, COFs are often obtained as powdered samples. However, powders are difficult to incorporate into practical devices due to their difficult immobilization or inconvenient clogging phenomena. Therefore, COF processing strategies should be applied to incorporate these organic materials into parts or devices, which would involve generating structures of at least several millimeters in size.

Overall, the state of the art shown in this summary confirms that photocatalytic applications of COFs for water remediation have great growth potential. However, further progress is needed on new challenges such as deactivation of biological pathogens, removal of microplastics, large-scale production of COFs or their incorporation into practical devices. This review provides a starting point to encourage researchers to move towards new developments that expand the applicability of covalent organic frameworks to overcome common drawbacks of traditional water decontamination processes.

Author Contributions: All the authors have contributed to the search of literature, conceptualization and writing of this review.

Funding: Ruben Mas-Baalleste acknowledges the financial support provided by the Spanish Ministry of Science and Innovation MCIN/AEI/FEDER (10.13039/501100011033) through the project PID2022-141016OB-I00 via Proyectos de Generación de Conocimiento 2022. Alicia Moya acknowledges the Spanish Government and the Funds Next Generation of the European Union through the grant Maria Zambrano-UAM (CA3/RSUE/2021-00648). Miguel Sanchez-Fuente thanks Ministerio de Ciencia e Innovación for a FPI contract (PRE2020-092295).

Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

The following abbreviations are used in this manuscript:

COF Covalent Organic Frameworks
ROS Reactive Oxygen Species
VB Valence band
CB Conduction band
ISC InterSystem Crossing

EIS Electrochemical Impedance Spectroscopy

XPS X-ray Photoelectron Spectroscopy
UPS Ultraviolet Photoelectron Spectroscopy

D-A Donor-Acceptor BPA Bisphenol A

MOF Metal Organic Frameworks

V_{fb} Flat band potential

E_f Fermi level

HOMO Highest Occupied Molecular Orbital LUMO Lowest Unoccupied Molecular Orbital

SET Single Electron Transfer

EnT Energy Transfer PC Photocatalyst

SED Sacrificial Electron Donor

FRET Förster Resonance Energy Transfer

E_{red} Reduction Potential E_{ox} Oxidation Potential

EPR Electron Paramagnetic Resonance

TC Tetracycline ACTP Acetaminophen SMT Sulfamethazine SMX Sulfamethoxazole RhB Rhodamine B Methylene blue MB MO Methyl Orange MB17 Mordant Black 17 **EBA** Eriochrome Black A **EBT** Eriochrome Black T RB Rose bengal MG Methyl Green

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