Abstract: The presence of pharmaceutical compounds in the environment is a reality that calls for more efficient water treatment technologies. Photocatalysis is a powerful technology available but the high energy costs associated with the use of UV irradiation hinder its large scale implementation. More sustainable and cheaper photocatalytic processes can be achieved by improving the sunlight harvesting and the synthesis of semiconductor/carbon composites has proved to be a promising strategy. Carbamazepine, diclofenac, and sulfamethoxazole were selected as target pharmaceuticals due to their recalcitrant behavior during conventional wastewater treatment and persistence in the environment, as properly reviewed. The literature data on the photocatalytic removal of carbamazepine, diclofenac, and sulfamethoxazole by semiconductor/carbon materials was critically revised to highlight the role of the carbon in the enhanced semiconductor performance under solar irradiation. Generally it was demonstrated that carbon materials induce red-shift absorption and they contribute to more effective charge separation, thus improving the composite photoactivity. Carbon was added as a dopant (C-doping) or as support or doping materials (i.e., nanoporous carbons, carbon nanotubes (CNTs), graphene, and derived materials, carbon quantum dots (CQDs), and biochars) and in the large majority of the cases, TiO$_2$ was the semiconductor tested. The specific role of carbon materials is dependent on their properties but even the more amorphous forms, like nanoporous carbons or biochars, allow to prepare composites with improved properties compared to the bare semiconductor. The self-photocatalytic activity of the carbon materials was also reported and should be further explored. The removal and mineralization rates, as well as degradation pathways and toxicity of the treated solutions were also critically analyzed.

Keywords: carbamazepine; diclofenac; sulfamethoxazole; semiconductor composites; C-doping; activated carbon; CNT; graphene; CQD; biochar; sunlight-driven photocatalysis

1. Introduction

World’s population increase along with the expected economic development, are major challenges water and energy consumption. However, nowadays 800 million people do not have access to drinking water and 1000 million live without electricity [1]. Thus, all strategies that enable to maximize water reuse, and contribute to the preservation of water resources, as well as those promoting more efficient use of renewable energies (e.g., solar energy), are essential to help in minimizing water scarcity and energy demand.

Water treatment technologies upgrade is the key to face current water quality challenges. The upmost challenge for drinking water and wastewater treatment, as well as water reuse, is linked to the presence of natural organic matter (NOM) and micropollutants in raw water, the latter often
related to pharmaceutical compounds (PhCs). Several PhCs partially or totally resist to conventional treatments. So, it is crucial to upgrade the design and operation conditions of wastewater treatment plants (WWTPs), and water treatment plants (WTPs) with innovative, cost- and resource-effective solutions to achieve high-quality water standards.

Some of PhCs resist conventional treatments and are also highly resistant to photolysis and/or adsorption onto solid matter leading to their high persistence in the aquatic environment. Consequently, tertiary treatments are fundamental to assure the removal of these recalcitrant PhCs. These end-of-line treatments cover a large range of technologies: UV disinfection, ozone or chloride, and advanced water treatment technologies. This last class gathers activated carbons adsorption, membranes filtration, advanced oxidation processes (AOPs), and also combinations of two or more technologies. The use of technologies based on adsorption or membranes transfers the pollutant from the water to the pore structure of the adsorbent material or to the reject of the membrane filtration, originating a residue that still requests further treatment or disposal in a landfill. On the contrary, ideal AOPs allow total mineralization of the pollutant to CO₂, water and inorganic compounds, or at least degrade the pollutant into more innocuous compounds.

AOPs can be broadly defined as aqueous phase oxidation methods based on the formation of highly reactive species, such as (mainly but not exclusively) hydroxyl radicals (OH•), leading the target pollutants decomposition [2]. Extensive research has been carried out on the use of AOPs to treat industrial and urban effluents. AOPs are divided into homogeneous and heterogeneous processes when considering their occurrence in a single phase or not, respectively. Depending on the methodology of OH• generation, AOPs can also be classified as chemical, electro-chemical, sono-chemical, and photo-chemical processes [3].

PhCs are a very large group of organic micropollutants of emergent concern. Thus remediation studies usually focus on a small group of PhCs classified as recalcitrant to conventional water treatment technologies and representative of distinct therapeutic classes and chemical functionalities. Following this rational three PhCs were selected to be addressed in this review: carbamazepine (CBZ) an antiepileptic, diclofenac (DCF) an acidic anti-inflammatory and analgesic, and sulfamethoxazole (SMX) an antibiotic (Table 1).

| Pharmaceutical Molecular Structure | Therapeutic Class | Properties |
|-----------------------------------|------------------|------------|
| Carbamazepine (CBZ)               | Antiepileptic    | MW = 236.3 g/mol  
|                                   |                  | pKₐ = 13.9 [4]   
|                                   |                  | Log Kₐow = 2.45 [4] |
|                                   |                  | Water solubility (25 °C) = 18 mg/dm³ [4] |
|                                   |                  | kₜ = 1.1 × 10⁻¹⁰ atm m³/mol [4] |
|                                   |                  | K₉₉₉₉₉ prim < 0.020 L/gss [5] |
|                                   |                  | K₉₉₉₉₉ sec = 0.0012 ± 0.0005 L/gss [5] |
|                                   |                  | kₐbiol ≤ 0.01 L/gss·d [6] |
|                                   |                  | K₉₉₉₉₉ OH = 8.8 × 10¹⁰ ± 1.2 × 10⁹ 1/M s [7] |
| Diclofenac (DCF)                  | Analgesic        | MW (acid) = 296.1 g/mol  
|                                   |                  | MW (sodium salt) = 318.1 g/mol  
|                                   |                  | pKₐ = 4.15 (acid) [8] |
|                                   |                  | Log Kₐow = 4.51 [8] |
|                                   |                  | Water solubility (25 °C) = 2.37 mg/dm³ [8] |
|                                   |                  | kₜ = 4.73 × 10⁻¹⁵ atm m³/mol [8] |
|                                   |                  | K₉₉₉₉₉ prim = 0.495 ± 0.032 L/gss [5] |
|                                   |                  | K₉₉₉₉₉ sec = 0.016 ± 0.003 L/gss [5] |
|                                   |                  | kₐbiol ≤ 0.1 L/gss·d [6] |
|                                   |                  | K₉₉₉₉₉ OH = 7.5 × 10¹⁰ ± 1.5 × 10⁹ 1/M s [7] |
Table 1. Cont.

| Pharmaceutical Molecular Structure | Therapeutic Class | Properties |
|------------------------------------|-------------------|------------|
| Sulfamethoxazole (SMX)             | Antibiotic        | MW = 253.3 g/mol  
|                                    |                   | pK_a1 = 1.6, pK_a2 = 5.7 [9]  
|                                    |                   | Log K_ow = 0.89 [9]  
|                                    |                   | Water solubility (37 °C) = 610 mg/dm³ [9]  
|                                    |                   | K_H = 6.4 × 10^{-13} atm m³/mol [9]  
|                                    |                   | K_d = 0.256 ± 0.169 L/g_d [10]  
|                                    |                   | k_rad ≤ 0.2 L/g_d·d [6]  
|                                    |                   | K_OH = 5.5 × 10^9 + 7 × 10^9 L/M s [7]  

CBZ is an antiepileptic drug used to control seizures, is also used to relief neuralgia, and as an antidepressant in bipolar depression. DCF is a popular non-steroidal anti-inflammatory drug, having also antiarthritic and antirheumatic properties. In 2008, Zhang et al. estimated 1014 ton CBZ and 940 ton DCF consumption per year in accordance with IMS health data reporting 942 tons of CBZ and 877 tons of DCF sold in 2007 [11]. In 2013, the European Commission established the first watch list with 10 substances that may pose a significant risk to, or via, the aquatic environment (Directive 2013/39/EU). DCF was included in this list and maintained in next revision (Directive 2015/495/EU). The most recent revision removed DCF considering that sufficient high-quality monitoring data was already available (Directive 2018/840/EU). SMX is an extensively used antibiotic from the sulfonamide class of antibacterial compounds. It is used in clinic for the treatment of respiratory tract infections, severe urinary tract infections, and enteric infections [12], is also used in veterinary (to treat infections or for preventive care in aquaculture) and as herbicide [13].

In this context, this work starts reviewing the literature supporting the relevance of CBZ, DCF, and SMX as representatives of PhCs with recalcitrant behavior in water treatment and in natural photodegradation processes. Secondly, the use of semiconductor and carbon materials for the photocatalytic degradation of these PhCs will be addressed, highlighting the role of carbon materials, in particular for the improved sunlight harvesting.

2. Pharmaceutical Compounds as Pollutants of Emergent Concern

PhCs are essential in assuring high human health standards and well-being and are also largely employed in veterinary, being ubiquitous chemicals in modern societies. However, it is proven that the high consumption patterns of PhCs associated with their high chemical stability and low biodegradability, contributed to their widespread release in the aquatic environment (mainly for those highly soluble). Given the biological activity and recalcitrant behavior of several PhCs, they constitute a major treat firstly to aquatic life and consequently to the whole food chain. Besides the public health concerns regarding short and long-term toxicological effects related to the exposure to trace amounts of PhCs trough potable water and food chain, the environmental perspective of this problem is also a relevant topic.

The first report on PhCs occurrence in the environment dates from 1965 when Stumm-Zollinger and Fair addressed the presence of steroid hormones in the effluents of the U.S.A. WWTPs to understand their biodegradation in WWTP, and in receiving water bodies that could eventually supply drinking water to households [14]. In the 1970s clofibric acid, salicylic acid, and chlorophenoxyisobutyrate (PhCs and metabolites) were detected in the U.S.A. wastewater effluents [15,16]. Nowadays, PhCs are detected in all sorts of environmental waters worldwide, and some of them are part of a watch list for EU-wide monitoring to access their risk for aquatic environment and support future prioritization (2018/840/EU).
2.1. CBZ, DCF, and SMX Detections in Wastewater and Environmental Water

Since the early monitoring studies in 1980s and 1990s CBZ, DCF, and SMX have been detected in wastewater samples and in environmental waters in the ng/L–µg/L range (Table 2). Between 1996 and 2000 Heberer [17] monitored 24 PhCs in sewage treatment plants (STPs) and environmental waters from the urban area of Berlin and CBZ, DCF, and SMX were the compounds detected at the higher concentrations. The results allowed to conclude that only 8% CBZ and 17% DCF were removed during wastewater treatment. Consequently, not surprisingly, concentrations up to 1.1 µg/L CBZ and 0.6 µg/L DCF were detected in surface water of the Berlin urban area [17]. In 2001, Ternes reported negligible CBZ removal in a municipal STP, the treatment of DCF was more effective (69%) [18]. CBZ and DCF were also detected in the same concentration range in STPs effluents and environmental waters from other European countries and Brazil [19–24]. Frist monitoring studies also report the detection of SMX in STPs effluents at ≈0.9 µg/L but reaching a maximum of 2.0 µg/L [21,25].

Table 2. Early detections of carbamazepine (CBZ), diclofenac (DCF), and sulfamethoxazole (SMX) in wastewater and environmental water.

| PhC  | Type of Water                      | Concentration (µg/L) | Publication Year | Ref. |
|------|------------------------------------|----------------------|------------------|------|
| CBZ  | STP effluents                      | 3.7 (max 6.3)        | 1998             | [19] |
|      | Ground water                       | 0.82 (max 1.1)       |                  |      |
|      | STP influent (Frankfurt/Mains)     | 2.2                  | 2001             | [18] |
|      | STP effluent (Frankfurt/Mains)     | 2.0                  |                  |      |
|      | Ground water                       | Up to 1.1            |                  |      |
|      | Drinking water                     | Up to 0.030          |                  |      |
|      | STP influent (Berlin)              | 1.78 (max 3.80)      | 2002             | [17] |
|      | STP effluent (Berlin)              | 1.63 (max 5.00)      |                  |      |
|      | Surface water (Berlin)             | 0.4–1.1              |                  |      |
|      | Surface water (Berlin)             | 0.025–1.075          | 2002             | [24] |
|      | STP effluent (France)              | 0.98–1.2             |                  |      |
|      | STP effluent (Greece)              | 1.03                 | 2003             | [23] |
|      | STP effluent (Italy)               | 0.3–0.5              |                  |      |
|      | STP effluent (Sweden)              | 0.87                 |                  |      |
| DCF  | Sedimentation tank effluent        | ≤2                   | 1996             | [27] |
|      | River Rhine                        | 0.015–0.304          |                  | (in [21]) |
|      | Different rivers                   | 0.038–0.489          |                  |      |
|      | STP effluents                      | 1.6 (max 2.1)        | 1998             | [19] |
|      | Rivers and streams                 | 0.80 (max 1.2)       |                  |      |
|      | STP effluent (Brazil)              | 0.47–1.92            | 1998             | [22] |
|      | Effluent Swiss STP                 | 0.31–0.93            |                  |      |
|      | Swiss lakes/rivers                 | 0.001–0.370          |                  |      |
|      | Brazilian STP                      | 0.4 (max = 1.4)      | 1999             | [20] |
|      | River water                        | 0.02–0.06            |                  |      |
|      | STP influent (Frankfurt/Mains)     | 1.9                  | 2001             | [18] |
|      | STP effluent (Frankfurt/Mains)     | 0.58                 |                  |      |
|      | Ground water                       | 0.93                 |                  |      |
|      | Drinking water                     | Up to 0.006          |                  |      |
|      | STP effluent (Berlin)              | 3.02 (max 7.10)      | 2002             | [17] |
|      | STP effluent (Berlin)              | 2.51 (max 4.70)      |                  |      |
|      | Surface water (Berlin)             | < 0.1–0.6            |                  |      |
|      | Ground water                       | Up to 0.38           |                  |      |
|      | Surface water (Berlin)             | Up to 1.030          | 2002             | [24] |
|      | STP effluent (France)              | 0.25–0.41            |                  |      |
|      | STP effluent (Greece)              | 0.89                 | 2003             | [23] |
|      | STP effluent (Italy)               | 0.47–5.45            |                  |      |
Table 2. Cont.

| PhC   | Type of Water      | Concentration (µg/L) | Publication Year | Ref.     |
|-------|--------------------|----------------------|------------------|---------|
| SMX   | River water        | ≈1                   | 1983             | [28] (in [21]) |
|       | STP effluents      | 0.9 (max 2.0)        | 1999             | [25]    |
|       | Surface water      | 0.14 (max 0.48)      | 1999             |         |
|       | Ground water       | Up to 0.47           | 2001             | [26]    |
|       | Ground water       | Up to 0.410          | 2001             | [26]    |
|       | STPs effluent (Berlin) | 0.9                | 2002             | [17]    |
|       | STP effluent (France)  | 0.07–0.09            |                  |         |
|       | STP effluent (Greece) | 0.09                 | 2003             | [23]    |
|       | STP effluent (Italy)  | 0.01–0.03            |                  |         |
|       | STP effluent (Sweden) | 0.02                 |                  |         |

In 2010, Loos and co-workers analyzed 35 compounds (pharmaceuticals, pesticides, perfluorinated acids, benzotriazoles, hormones, and endocrine disrupters) in more than 100 river water samples from 27 European countries. Among all PhCs CBZ was the most frequently detected compound (95%) and at the highest concentration level (0.308 µg/dm³ for percentile 90%, and a maximum of 11.561 µg/dm³) [29]. Also DCF and SMX were detected with a frequency above the average frequency for all the compounds under study (61%), i.e., DCF was detected in 83% of the samples at concentrations of 0.043 µg/dm³ in 90% percentile, and maximum values of 0.247 µg/dm³, while SMX was detected in 75% of the analysis at concentrations of 0.104 µg/dm³ in 90% percentile, and maximum values of 4.072 µg/dm³ [29]. A survey on the occurrence of 59 polar organic persistent pollutants in 164 locations of European ground water (23 countries) identified 21 relevant compounds, among which CBZ (detection frequency 42%, max. 0.39 µg/dm³) and SMX (detection frequency 24%, max. 0.038 µg/dm³) [30]. The study also showed that CBZ was frequently detected above the European ground water quality standard for pesticides (0.1 µg/dm³) [30].

The results of a systematic study regarding the fate of CBZ in eleven Austrian STPs conducted by Clara et al. in 2004, revealed that this antiepileptic medicine consistently resisted to degradation and adsorption during conventional wastewater treatment, and also during soil passage and subsurface flow, thus allowing to propose CBZ as an anthropogenic marker for detecting wastewater in the aquatic environment [31].

In 2008, Zhang et al. reviewed literature data on CBZ and DCF removal during wastewater treatment as well as their occurrence in water bodies [11]. According to monitoring data in WWTPs (1998–2007), CBZ is considered persistent since 70% of the studies reported only 0 to 10% removal of this PhC. For DCF the removal efficiencies reported varied between 0 and 80%. The justification for the distinct behavior can be related to the high resistance of CBZ to biodegradation at low concentration while DCF can be biodegraded at specific conditions (i.e., anoxic biofilm reactor, acidic conditions) [11]. Phototransformation during sunlight irradiation in WWTPs cannot be disregarded as a removal route. However, in deionized water CBZ presents very high stability with half-life of approximately 100 days while DCF and SMX present values two orders of magnitude lower (5.0 and 2.4 days, respectively) [11,23]. In real wastewater matrices the presence of organic matter that can behave as photosensitizers or inner filters must also be considered. In fact, humic acids act as inner filters for CBZ and DCF lowering their degradation, but in the case of SMX behave as photosensitizers promoting its phototransformation [11,23].

The recalcitrant nature of some PhCs during conventional wastewater treatment can be justified in light of their properties. According to Joss et al. [6], the biological degradation constants ($k_{biol}$, Table 1) of CBZ and DCF are indicative of less than 20% removal during wastewater treatment with conventional activated sludge (CAS) or membrane bioreactor (MBR) technologies, supporting the systematic low biodegradability during secondary treatment. Regarding CBZ, DCF, and SMX also, no sorption onto activated sludge is expected to occur due to distribution coefficient water/sludge values ($k_d$, Table 1).
much lower than reference value for significant sorption (0.500 L/gso) [11]. Volatilization could be another route to the removal of these PhCs during wastewater treatment but their low Henry constant values ($k_{H}$, Table 1) indicate that they are nonvolatile compounds.

A very recent study classified 13 PhCs, including DCF and SMX, as priority PhCs in the aquatic environment of China, according to the proposed ranking system that is supported on three criteria (occurrence, exposure potential, and ecological effects). [32]. DCF and erythromycin presented the highest predicted environmental risk. This prioritization is in line with other ranking studies since DCF is considered a priority PhC in eight reports, and SMX in six of the works [32]. In fact, Gou et al. identified 13 PhCs (including CBZ and DCF) and three metabolites as presenting high risk score since their estimated exposure is higher than predicted no-effect concentration (PNEC) [33]. Besse et al. calculated the risk quotient ratios with available ecotoxicity data of 76 PhCs and their metabolites and, according to the conditions required by the European Medicine Agency (EMEA) guidelines of 2006, CBZ, DCF, and SMX are three of the six PhCs identified as presenting environmental risk [34]. The Global Water Research Coalition (GWRC) consolidated data from 25 literature studies on 153 PhCs and, according with the seven more relevant criteria, CBZ, DCF, and SMX were identified as the top three PhCs of the group of 10 high priority PhCs [35]. In fact, these three PhCs present the highest number of occurrences in the literature (15, 12, 13, respectively) being the only ones that fulfil all the relevant criteria (i.e, regulation, consumption/sales, physicochemical properties, toxicity, occurrence—wastewater, river water, ground water, drinking water—, degradability/persistence and resistance to treatment) [35].

2.2. Photodegradation

As mentioned previously, the occurrence of CBZ, DCF, and SMX in environmental waters worldwide results from the low effectiveness of conventional water treatments as barriers for these PhCs, due to their high chemical stability and low biodegradability. Once in the environment, the fate of these PhCs in the water-cycle depends on their properties namely their hydrophobic/hydrophilic character that will determine their interactions with water, dissolved species, and solids, as well as, their stability under sunlight irradiation.

Photodegradation at the top level of surface waters may be a natural elimination pathway for recalcitrant PhCs in environmental waters. Direct and indirect photochemical processes can occur depending on the PhC nature, water matrix, and solar irradiation spectrum. Direct photolysis of an organic compound only occurs when the absorption spectrum of the compound overlaps the intensity spectrum of the incoming irradiation, resulting in the electronic excitation of the molecule that promotes its phototransformation. In indirect processes, photoactive compounds—photosensitizers—absorb sunlight producing reactive species such as hydroxyl radicals (OH) that can induce PhCs degradation. According to the UV spectra (Figure 1) it is possible to conclude that only a very small fraction of sunlight (i.e., UV-A and UV-B) is expected to promote direct photolysis of CBZ, DCF, and SMX.

![Figure 1. UV absorption spectra of CBZ, DCF, and SMX (5 mg/L solutions in inorganic matrix at pH 7.1) versus sunlight spectrum at sea level (ASTM G173-03 Global, [36]).](image-url)
Doll and Frimmel [37] evaluated the fate of CBZ in surface waters under natural and simulated solar light, concluding that although CBZ absorbs most of the incoming radiation compared with the other PhCs, it presents the lowest degradation rate constants. In real waters with low concentrations of NOM photodegradation of CBZ can be favored (indirect pathway by NOM-induced reactions), but in high concentration, NOM has a detrimental effect acting as the inner filter (radical scavenger and/or as precursor of reactive species). The presence of other PhCs also decreases the CBZ degradation rate (competitive inhibition) [37].

Andreozzi et al. studied the photodegradation of CBZ (8.0 × 10⁻⁶ mol/L) in bi-distilled water at pH 5.5 during solar light exposure determining a pseudo-first order kinetic constant of 5.7 × 10⁻³ h⁻¹ and a half-life time of 121.6 h, while in spiked natural water CBZ half-life time increased to 907 h [38]. Assays in the presence of nitrates reveal that these anions act as photosensitizers reducing CBZ half-life time to 69 h–11.2 h (5.0 × 10⁻⁴–1.5 × 10⁻² g/L). On the other hand, dissolved humic acid increased photodegradation half-life time of CBZ to 233.7 h possibly related to the reduction of direct photodegradation due to strong UV absorption of humic acid that acts as inner filters [38]. The laboratorial study developed by Yamamoto et al. also pointed out relatively high resistance of CBZ (20–100 µg/L) to solar light photodegradation with half-life times of 84 h, in August 2006, and 2100 h, in May 2007 [39].

Lam et al. studied the aquatic persistence of eight PhCs in sunlight-exposed pond water identifying CBZ and SMX as the most persistent with mean half-life times of 82 days and 19 days, respectively, due to their resistance to both biodegradation and photolysis [40].

The low photodegradability of CBZ is certainly related to its low quantum yield. In fact, according with the experimental quantum yields for the direct photolysis of PhCs in bi-distilled water at pH 5.5, CBZ has one of the lowest values (4.77 × 10⁻⁵ under sunlight) [23]. For SMX the value obtained under lamplight was 4.29 × 10⁻³ mol/Einstein and for DCF the results reported are 3.13 × 10⁻² mol/Einstein under lamplight, and 3.37 × 10⁻² mol/Einstein under sunlight [23]. Chiron et al. determined an average quantum yield of 1.5 × 10⁻⁴ mol/Einstein for CBZ direct photolysis in Milli-Q water under medium-pressure mercury lamp (highest contribution from 313, 366, 406 nm) [41]. In surface water at pH 7 quantum yields of CBZ irradiated with medium-pressure (205–500 nm) and low-pressure (253.7 nm) mercury lamps are slightly higher (0.6 × 10⁻³ and 2.3 × 10⁻³ mol/Einstein, respectively) [42].

The photodegradation of CBZ (2.1 × 10⁻⁴ mol/L) in artificial river and estuarine water (with humic acids, Fe(III), NO₃⁻, and Cl⁻) was studied by Chiron et al. who found a decrease in photodegradation rate with increasing pH, which was attributed to the distinct iron species in solution [41]. At acidic pH values CBZ photodegradation occurs via dissolved Fe(III) that originates OH• while at higher pH values Fe(III) colloids are predominant and produce lower amounts of OH• but are able to induce charge-transfer reactions, this will not be pure photodegradation processes since from the Fenton process it is known that iron can catalyse the degradation. However, under neutral to basic conditions the presence of chlorine enhance CBZ photolysis most probably due to the interaction between Fe(III) colloids and Cl⁻ leading to the formation of Cl₂• that albeit presenting slower degradation than OH• would be faster than charge-transfer processes occurring on the surface of Fe(III) colloids [41]. After 8 h irradiation, the toxic, mutagenic and carcinogenic acridine (major intermediate of CBZ direct photolysis) accounted for 10% of the initial carbamazepine species [41]. Since this transformation product has greater health and environmental risk than the pristine compound more effective degradation routes are needed.

In 2008, Matamoros et al. evaluated the photodegradation of four PhCs, including CBZ (10–40 mg/L), under simulated solar light and sunlight, in river water and sea water, as well as, distilled water for getting information regarding photodegradation pathways [43]. To assure irradiation close to real conditions, the authors used quartz vessels transparent to UV light. The results pointed out CBZ as the compound most resistant to photodegradation since in Milli-Q water it presented 38.5 h of half-life time, and in two river water or sea water, the value decreased to 8.25–14.4 h, revealing the role of DOC in indirect photodegradation. In the assays performed under sunlight the half-life time in river
water was eight times higher (67.4 h) compared with the value obtained under simulated conditions, being this decreased explained by the daily light variation.

Early studies by Buser et al. [22] reported a great decrease in DCF concentrations from a major tributary to a lake, to the outflow of the same lake (inflow from 0.005 to 0.370 µg/dm³ and outflow from <0.001 to 0.012 µg/dm³) estimating more than 90% DCF removal most likely due to photodegradation (not detected in sediments and laboratory experiments corroborate negligible absorption onto sediment particles). Lake water spiked with DCF presented minimal chemical and biological degradation under dark, however once exposed to sunlight a fast photodegradation occurred attaining the half-life time in less than 1 h of irradiation, most likely via direct photolysis [22]. In a further study the same authors were able to identify photodegradation products, namely 8-chlorocarbazole-1-acetic acid that was detected in all the experiments and photodegrades faster than DCF, and concludes that photochemical degradation is the predominant elimination process for DCF in the lake [44].

Literature studies report decomposition quantum yields for DCF from 0.031 to 0.22 mol/Einstein in the laboratory, in agreement with those determined in environmental conditions (0.094 and 0.13 mol/Einstein) [45]. According with simulated profiles and field measurements in a lake (three-month period) Tixier et al. estimated an average elimination rate of ≈2 h⁻¹ (0.082 days⁻¹) for direct DCF photolysis which is in accordance with the model simulation based on field data (overall elimination rate constant ≈2.1 h⁻¹ (0.088 days⁻¹), and eight days of half-life time), proposing this process as the main elimination route for DCF in environmental water [46]. The same study classified CBZ as fairly persistent with an overall elimination rate constant of ≈0.26 h⁻¹ (0.011 days⁻¹), and 63 days of half-life time [46].

Photodegradation profiles of DCF and SMX under artificial sunlight irradiation in dechlorinated tap water at pH 8 (mimicking surface water without bacteria or suspended solids) reveal that DCF, with a half-life time of 0.40 days, was by far much more sensitive to photolysis than SMX, which needed 54 days to decrease half of the initial concentration [47]. Although SMX has relatively low absorption in the UV-B region above 290 nm, according to the studies of Moore and Zhou in the mid-1990s, the susceptibility of this sulfonamide antibiotic to photodegradation is highly dependent on the solution pH [12,48]. In fact, under UV light the pseudo-first-order kinetic constant is higher than 0.15 min⁻¹ at pH < 4, and lower than 0.025 min⁻¹ at pH ≥ 7, with the inflection point matching the pKₐ value [48]. Also, the quantum yield of SMX photodegradation determined under UV irradiation is 0.47 mol/Einstein at pH 3 and only 0.084 mol/Einstein at pH 9 [48].

The complex SMX speciation profile allows concluding that while the nonionized form is extremely susceptible to photodegradation under UV or sunlight irradiation, the anion form, predominant at pH 5.6 or higher (pH values of wastewater effluent and environmental water), is more stable. The high photocatalytic activity of the SMX anionic form offers and explanation to the slower and less effective photolysis of SMX in wastewater compared with deionized water, both at natural pH (4.8 and 8.1, respectively) [49] or in deionized water with pH values increasing from 3 to 10 [50]. This behavior is a strong indication that SMX resistance in the environment is most likely due to its resistance to photodegradation in slightly basic conditions. The presence of natural water constituents (i.e., fluvic acids and suspended solids) were reported to have a negative effect on the photodegradation of SMX (1–10 mg/L), in the case of fluvic acid at 50 mg/L the half-life is five times higher compared to deionized water [50]. The type of organic matter is also relevant for the photolysis of SMX, Ryan et al. verified that effluent organic matter (EfOM) enhanced SMX photodegradation, while NOM had no effect, probably due to the higher concentration of nitrates in EfOM, which act as photosensitizers for the production of hydroxyl radicals, and also due to the presence of triplet excited state organic matter in EfOM [51]. Although acidic pH favors SMX photodegradation, the mineralization degree is negligible due to the formation of abundant transformation products [49,52]. 4-amino-N-(5-methyl-2-oxazolyl)benzenesulfonylamide is the main photodegradation product identified, and although presenting the same empirical formula of SMX suffered a photoisomerization reaction of the isoxazole ring [12,40,49]. Trovó et al. evaluated the acute toxicity of SMX photodegradation products for
Daphina magna and Vibrio fischeri concluding that the first was the most sensitive organism (from 60% to 100% immobilization) [49]. Since SMX was totally removed after 30 min irradiation toxicity was attributed to the degradation products. Gmurek et al. also studied the toxicity of the long-term SMX phototransformation products under simulated solar light towards Vibrio fischeri growth and luminescence [52]. Regardless of the irradiation time, the bioluminescence inhibition of the mixture of transformation products retained the toxicity of the parent SMX solution. Regarding growth inhibition, the increase of irradiation times shifted the curve to higher concentrations but the growth inhibition decrease was not proportional to the degradation of the SMX entities [52].

3. Photocatalysis

In 1972, Fujishima and Honda discovered the photoelectrocatalytic water splitting by using TiO$_2$ electrodes [53] and paved the way for more efficient use of light in chemical processes. Since then photocatalysis has rapidly expanded for a large range of environmental and energy applications [54–61], is considered a potential solution to the worldwide energy shortage and for counteracting environmental degradation [60]. As stated by Fujishima and co-workers, in a time-scale of ten years the basic scientific knowledge on TiO$_2$ photocatalysis allowed the development of a technological field reaching the real large-scale industrial exploitation [62], is thus a great example of fundamental science solving real world challenges.

More efficient use of the solar spectrum for an effective photodegradation of persistent PhCs as is the case of CBZ, DCF, and SMX can be achieved by applying materials with photocatalytic properties, as is the case of semiconductors. As previously mentioned TiO$_2$ is by far the most popular photocatalyst due to its unique properties: inertness, high photostability and photoactivity, cost-efficiency and also non-toxicity to humans and environment [61]. Other metal oxides (e.g., ZnO, WO$_3$), and their derivatives have also been explored for the degradation of organic compounds [63]. Besides the classical metal oxide semiconductors, the pioneering work by Wang et al. in 2009 reported the H$_2$ evolution by metal-free polymeric graphitic carbon nitride (g-C$_3$N$_4$) under visible light [64], thus proposing the first metal-free photocatalyst. Since then these organic solids—g-C$_3$N$_4$ and derivatives, as well as other conducting polymers like polyaniline (PANI), polyacetylene, polythiophene, polypyrrole, poly-p-(phenylenevinylene)—proved to be promising visible-light-responsive organic photocatalysts for several key reactions (e.g., photochemical water splitting, oxidation reactions, CO$_2$ mitigation, pollutant degradation and bacterial disinfection) [65–68], as well as, playing important roles in biomass conversion, i.e., sustainable chemistry processes [66].

Photocatalysis is initiated when a semiconductor absorbs a photon ($h\nu$) with energy equal to, or greater than, the band gap ($E_g$) of the semiconductor, causing the excitation of an electron ($e^-$) from the valence band (VB) to the conduction band (CB), which leads to the formation of an electron-hole ($e^-/h^+$) pair (reaction (1)). These photogenerated charge carriers ($e^-/h^+$) can recombine or migrate to the surface of the catalyst. Recombination can occur (i) in the bulk of the semiconductor, (ii) and/or mediated by surface species, (iii) or adsorbed molecules. When charge carriers migrate to the surface of the photoactive material and are scavenged by the species adsorbed onto the surface of the semiconductor, they induce the redox reactions of the adsorbed species. The competition between the fast kinetic of recombination and slow kinetic of charge separation and transfer (i.e., migration) to the surface of the material determines the efficiency of the photocatalytic process. Consequently, the first requisite to boost the photo-oxidation of the target pollutant is to prevent the $e^-/h^+$ recombination. The photo-generated hole is a strong oxidizing agent that can directly react with the target pollutant or with water to produce hydroxyl radicals (OH$^\bullet$) (reaction (2)), the former occurs when the redox potential of the pollutant is less negative than that of semiconductor valence band while in the latter the oxidation of the pollutant occurs through radical chain reactions. On the opposite the electron is a strong reducing agent able to react with electron acceptor species, as is the case of dissolved oxygen to
create superoxide radicals ($O_2^{•−}$) (reaction (3)), or other species with a redox potential more positive than that of the photocatalyst conduction band.

$$\text{Photocatalysis initiation: } \text{semiconductor} + \nu \rightarrow e^{−} + h^{+} \quad (1)$$

$$\text{photo-oxidation: } h^{+} + OH^{−} \rightarrow OH^{•} \text{ or } h^{+} + H_2O \rightarrow OH^{•} + H^{+} \quad (2)$$

$$\text{photo-reduction: } e^{−} + O_2 \rightarrow O_2^{•−} \quad (3)$$

The efforts to optimize photocatalysts have been mainly focused on the increase of light efficiency, mainly by searching for solar-light-responsive photocatalysts and minimizing electron-hole recombination [60,63].

Among photocatalysts, nanosized semiconductor particles have met superior photocatalytic properties due to their large surface area, distinctly good dispersion capability and quantum size effects [63,69]. Yet, the large scale application of semiconductor nanoparticles has two major constraints: aggregation in suspension leads to fast loss of active sites, and consequent decreased photocatalytic efficiency, and the challenging separation, recovery, and reuse [63].

The synthesis of semiconductor hybrids and/or composites by coupling a semiconductor with organic/inorganic materials is a promising option to overcome the above mentioned drawbacks and challenges [70,71]. Usually, a hybrid material consists of a mixture of two or more phases at the nanometer scale or molecular level, and the hybrid material features do not correspond to the sum of the properties of the single phases but to synergistic interactions among them [70]. Typically, a composite consists of the dispersion of a micron-level phase or nanometer-size structures in a matrix, with the properties of the composite resulting from the addition of each phase contribution. To the best of our knowledge in photocatalysis most of times the terms hybrid and (nano)composite are indistinctively used to denote a material composed by two or more compounds and/or materials, regardless the establishment or not of specific chemical interactions between both phases, and the relative proportion of each one in the final mixture.

Due to their extensive surface area, black body properties, and high charge carrier mobility, carbon materials have been largely explored as dopants, hybridizing materials or supports of semiconductors. They are, undoubtedly, a very effective route to improve photocatalytic activity, as proved by the number of literature studies reviewing TiO$_2$/carbon performance in the last decade [72–85]. For example Khalid et al. compared the performance of TiO$_2$/carbon nanomaterials against bare TiO$_2$ concluding that carbon doping and composites with activated carbons, fullerene, carbon nanotubes (CNTs), or graphene lead to materials with enhanced photocatalytic activity, adsorption capacity, electron scavenging and sensitization ability, extended visible light absorption, and also easier separation [73].

The enhanced photocatalytic activity due to the addition of carbon to a semiconductor can be reasoned in light of two mechanisms (i) improved charge separation in the presence of carbon, (ii) carbon acting as photosensitizer (Figure 2) [86]. When porous carbons are added to semiconductors, several synergistic effects due to adsorption must be considered. The charge separation mechanism by semiconductor/carbon composites was proposed by Woan et al. considering that carbon materials with metallic conductivity are able to trap the electron generated in the semiconductor, therefore hindering the recombination (Figure 2a) [76]. Wang et al. proposed the mechanism where the carbon material acts as photosensitized by transferring electrons to the conduction band of the semiconductor (Figure 2b) [87].
Doping semiconductors is an option to improve their optical properties since the replacement of metal or nonmetal atoms by other elements, or the addition of atoms in interstitial positions, ideally maintaining the crystalline structure of the semiconductor, can shift the onset of the response from the UV to the visible region [88,89]. Several metals (e.g., W, V, Fe or Cu) and nonmetals (B, C, N, F, P, S, Cl, and Br) have been used to successfully dope TiO₂ [88]. Among the nonmetals, N- and C-doping have similarities since both favor the formation of oxygen vacancies, enhancing the photocatalytic activity of TiO₂ in the visible region [89]. The theoretical study by Valentin et al. reveals that at low C concentration the characteristics of the C-doped TiO₂ depends on the O availability during...
the synthesis: substitution of O by C atoms and O vacancies are favored under O₂-poor conditions, while interstitial C and substitution of Ti by C atoms is preferred at O₂-rich conditions [89].

Irie et al. attributed the visible light responsivity of C-doped TiO₂ (anatase), prepared by mild oxidation of TiC and further annealing under O₂, to the 0.32% of C atoms located at O sites [62]. Albeit the very small carbon content in the doped TiO₂, it was enough to shift the absorbance edge and shoulder to the visible light region (bandgap narrowing). C-doped TiO₂ (anatase) catalysts can be prepared by hydrolysis of TiCl₄ with nitrogen bases (e.g., tetrabutylammonium hydroxide) followed by calcination at 400 °C, being proved that optical properties are dependent on the doping percentage [90]. Materials with 2.98% and 0.43% of C absorb light at 400–700 nm and were five times more active than an N-doped for the artificial solar light degradation of 4-chlorophenol, while the unmodified TiO₂ was almost inactive [90]. TiO₂-C-0.43% was also effective for the gas-phase photodegradation of acetaldehyde, benzene, and carbon monoxide in air under diffuse indoor light [90].

Recently C-doped semiconductors have also been successfully tested for the photocatalytic degradation of CBZ and DCF under UV and visible light (Table 3) and, as expected, TiO₂ is by far the most explored semiconductor. The majority of the studies use initial DCF or CBZ concentrations in the mg/dm³ range, deionized water, and single-solute conditions.

In 2017, Surenjan and co-authors reported the effectiveness of C-doped spherical TiO₂ in the anatase form for the photodegradation of DCF and CBZ after 4 h under visible light since after this time none of the pharmaceuticals or respective degradation intermediates and products were identified (mineralization degrees higher than 98%) [91]. The characterization of the doped photocatalyst reveals that C atoms may substitute Ti sites or occupy interstitial positions. The authors found that in the presence of glucose the degradation rate of CBZ decreases (from 0.0348 min⁻¹ in single-solute to 0.0249 min⁻¹) indicating that glucose acts as a sacrificial agent. In dual-solute conditions (i.e., DCF and CBZ) the rate of CBZ degradation has always a more pronounced decrease than that of DCF but even thought a complete removal of both PhCs was achieved in 210 min [91].

In a distinct approach, Sambandam et al. evaluated the effect of the morphology of C-doped TiO₂ (anatase form) catalytic activity for the degradation of CBZ under visible light [92]. The authors proposed that in comparison with other crystal morphologies the rice grain shaped catalysts enhanced visible light degradation of CBZ (k₁ of 0.094 min⁻¹ vs. 0.059–0.02 min⁻¹ for the other morphologies) due to the conjugation of co-exposed low energy/high energy [101]/[001] facets thermodynamically stable or not, and high surface area (229 m²/g vs. 33–62 m²/g for the other morphologies) [92].

Table 3. Photocatalytic degradation of CBZ and DCF by C-doped semiconductors.

| PhC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|-------------------------|--------------------|-----|
| CBZ | TiO₂C    | Microwave synthesis: spherical catalyst (anatase) of 50 nm | 250 mg/dm³ of catalyst allow higher and faster degradation | [91] |
| DCF |          | [TiO₂/C] = 50 - 350 mg/dm³ | High [PhC] lowers degradation efficiency, likely due to the higher competition of PhCs and intermediates for active sites | |
|     |          | [PhC] = 50 - 250 µg/dm³ (each PhC) | Optimum light intensity: 8000 lx | |
|     |          | Continuous O₂ | pH 7.5 allows higher degradation rate of CBZ and DCF | |
|     |          | pH 6.5 – 8.5 | At pH > 7.5 repulsion between the photocatalyst, negative surface charge and excess of OH⁻ may prevent the formation of OH⁻ | |
|     |          | Visible (intensity 4000–10,000 lx) | In the presence of glucose CBZ degradation rate decreases | |
|     |          | 60 min dark (ads) + 4 h light (deg) | Dual-solute (DCF and CBZ) degradation profiles: DCF removal was faster (0.0334 min⁻¹) while CBZ removal rate slightly decreases compared with single-solute assay (0.0348 min⁻¹ vs. 0.0247 min⁻¹) | |
|     |          | Single-solute and multi-solute | After 4 h irradiation no CBZ or DCF is detected, > 98% TOC removal at higher [PhC]₃ (5 mg/dm³) and catalyst loading (500 mg/dm³) | |
The contribution of C-doping for enhanced semiconductor photodegradation of DCF under simulated solar light is clearly illustrated in the manuscript published by Cordero-Garcia et al. [94]. C-doping at percentages as low as 0.1–1.8 wt.% has a positive effect on: (i) the narrowing of the WO3/TiO2 band gap, (ii) the decrease of bandgap values to the visible light region (i.e., Eg values (in eV) of 3.15 for TiO2−0.04%C, 3.10 for WO3/TiO2−0.05%C, 3.05 for WO3/TiO2−0.1%C, and 2.98 for WO3/TiO2−0.18%C), and (iii) lower recombination e−/h+ [94]. The incremental slight displacement of the adsorption edge of the catalysts to the visible region proved to increase their mineralization efficiency for DCF, from 65% in the bare TiO2 to 76.2% in the WO3/TiO2, since WO3 acts as acceptor center for the photogenerated electrons. Values around 80% were attained for the C-doped mixed oxide (78.1% and 82.4% for, respectively WO3/TiO2−0.1%C and WO3/TiO2−0.18%C). Regardless the catalyst, the DCF degradation kinetic is always one order of magnitude faster than the TOC removal profile pointing out the formation of degradation products (possibly quinine imide compounds) more recalcitrant than parent DCF. The synergetic effect of C and WO3 on the properties of anatase under solar irradiation was explained by the authors according to the mechanism presented in Figure 4. The incorporation of C atoms created a narrow C2p band just above TiO2 O2p valence band allowing

| PbC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|-------------------------|-------------------|------|
| CBZ | TiO2:C   | • Microwave synthesis: anatase TiO2/C catalyst (50–70 nm) with controlled shape | • Mainly graphite carbon is deposited on the surface of TiO2 | [92] |
|     |          | • [TiO2/C] = 230 mg/dm³ | • From the four TiO2/C morphologies the rice grain shaped nanocrystal catalysts presented enhanced visible light degradation of CBZ | |
|     |          | • [CBZ]0 = 50 µg/dm³ | • CBZ photodegradation obeys the pseudo-first order kinetic model for all morphologies with kₜ (in min⁻¹) following the sequence of 0.094 (rice grain) >> 0.059 (spherical) > 0.044 (distorted spherical) > 0.020 (hexagonal) | |
|     |          | • Continuous O₂ flow 300 mL/min | | |
|     |          | • Visible (150 W, ~400 nm, 7700 lux) | | |
|     |          | • 45 min dark (ads) + 2 h light (deg) | | |
|     |          | • Single-solute & deionized water | | |
| DCF | TiO2:C   | • Sol-gel synthesis: mesoporous TiO2 (with Amberlite and calcination at 300 and 550 °C) | • TiO2:C (pure anatase) present the best photocatalytic performance attainment ≈ 80% DCF degradation, ≈80% mineralization, and pseudo-first order rate constants of 1.0964 × 10⁻³ min⁻¹ and 1.3242 × 10⁻³ min⁻¹ | [93] |
|     | TiO2:C,N | | • High photoactivity of TiO2/C attributed to increased rates of e− transfer with increasing specific surface area. | |
|     |          | • [Catalyst] = 1 g/dm³ | | |
|     |          | • [DCF]₀ = 10–50 mg/dm³ | | |
|     |          | • pH = 7 | | |
|     |          | • Simulated solar light (Xe lamp 1500 W) > 290 nm | | |
|     |          | • Single-solute | | |
|     |          | | | |
|     |          | • Continuous flow reactor | | |
| DCF | WO3/TiO2-C | • Sol-gel method: under acidic conditions (<2 wt.% WO3, 0.1 or 0.18 wt.% C) | • C-doping narrows WO3/TiO2 band gap and decreases Eₕ | [94] |
|     | WO3/TiO2 | • [Catalyst] = 1 g/dm³ | • All catalysts attain total DCF degradation after 250 kJ m⁻² of accumulated energy but mineralization is favored at higher C-doping | |
|     | TiO2 | • [DCF]₀ = 10 mg/dm³ | • During degradation, pH decreases due to the formation of carboxylic acids and Cl⁻ release | |
|     |          | • pH = 7 | • DCF degradation kinetic faster than mineralization process (10⁻³ vs. 10⁻⁵ min⁻¹): intermediate products more recalcitrant than parent DCF | |
|     |          | • Simulated solar light (Xe lamp 1500 W) > 290 nm | • Synergetic effects among semiconductors and C allows high catalytic activity of WO3/TiO2-C under solar light (changes in electronic structure) | |
|     |          | • Single-solute | | |
|     |          | | | |
| DCF | ZnO/TiO2 | • Composites obtained by high-thermal processing (900 °C) | • ZnO/C has the highest efficiency (60% DCF degradation in 20 min) | [95] |
|     | ZnO/C | • [Catalyst] = 0.5 g/dm³ (TiO2 – Aerioxide P25) | • TOC removal profile similar to that of DCF indicating that TOC decrease is connected with direct DCF decomposition, 35% TOC remaining corresponds to DCF and transformation products | |
|     | TiO2 | • [DCF]₀ = 10 mg/dm³ | • DCF photodegradation follows pseudo-first order kinetic model: kₜ (ZnO/C) = 8 kₜ (P25) and kₜ (ZnO/C) = 2.5 kₜ (ZnO) | |
|     | | • pH = 7 | • Defects in ZnO structure of ZnO/C contribute to better DCF removal | |
|     | | | • Ads of ZnO/C have the highest efficiency (60%) in C-doped TiO2 | |
| | | • Single-solute | | |
| DCF | TiO2:C | • C-doped anatase TiO2 coated on glass plate (12.5% C) | • TiO2/C adsorbed UV-A light and also a range of visible light due to its smaller bandgap compared with bare TiO2 | [96] |
|     | | • [DCF]₀ = 0.5 mg/dm³ | • OH• were generated and DCF was degraded (< quantification limit) | |
|     | | • pH = 6.2 – 7.2 | | |
|     | | • UV-A (3 lamps 15 W, max 365 nm) | | |
|     | | • Single-solute & deionized water | | |

ads-adsorption, deg-degradation
the visible light absorption and further transference of these photogenerated electrons to the Ti 3d conduction band. The dual-oxide W/Ti system allows the trapping of the photogenerated electrons improving the charge carrier separation. In both oxides, the holes can react with water to produce OH\(^{•−}\) while the e\(^{−}\) can react with adsorbed O\(_2\) that is reduced to superoxide radicals and further reacts with protons to form HO\(_2\)^{•−} [94].

![Figure 4. Mechanism of charge carrier separation proposed by Cordero-Garcia et al. [94] for photoexcited WO\(_3\)/TiO\(_2\)-C (Reprinted with permission [94], Copyright 2019, Elsevier).](image)

3.2. Semiconductor/Activated Carbon

As mentioned, the use of semiconductors has several drawbacks related namely with aggregation, high recombination of photogenerated e\(^{−}/h^{+}\) pairs, limited activity under solar light and post-separation. To overcome these issues the scientific community explores different strategies. Back in early 1989, Tanguay et al. reported that anatase TiO\(_2\) supported on carbon felt was photoactive for the degradation of dicholomethane, and allowed easy separation of the catalyst from the reaction mixture [97]. Between 1998 and 2001, Matos et al. reported that the addition of an amorphous carbon phase (i.e., activated carbon) to semiconductor in mixed suspensions enhance the photoactivity for the degradation of organic pollutants [98–100]. The immobilization of semiconductor photocatalysts onto solid supports is an option to improve their performance, and among the several materials explored, activated carbons meet the criteria requested to be considered valuable supports since they (i) allow strong adherence between catalyst and support, (ii) the catalyst reactivity is not affected by the attachment process, (iii) have high specific surface area, and (iv) have a high adsorption affinity towards the pollutants [71]. In fact, several studies reveal that the use of activated carbon as support material can increase the photodegradation rate by progressively allowing an increasing quantity of substrate to come in contact with the immobilized metal oxide by means of adsorption [71], by minimizing the recombination e\(^{−}/h^{+}\) of the metal oxide [101], and due to the black character of activated carbons harvesting the solar spectrum is favored. The studies aiming to evaluate the synergetic effects of semiconductor/carbon nanocomposites revealed that an increased adsorption of the contaminants onto the activated carbon phase, followed closely by a transfer through an interphase to the TiO\(_2\) phase, gives a complete photodegradation process [71,101]. In 2011, Lim et al. reviewed the synergistic adsorption-photocatalytic processes of TiO\(_2\)/activated carbon composites addressing the challenges and further developments for water treatment and reclamation. Authors propose the coupling of [70].
these composites with membrane separation technologies to recover and regenerate the composites and also to enhance solar harvesting by using N-doped activated carbons [74]. The improvement of solar spectrum harvesting and catalysts immobilization strategies are also identified by Chong et al. as urgent needs in order to provide cost-effective photocatalytic technologies for water treatment sector [57].

Furthermore, as recently highlighted in the review on the photochemical activity of nanoporous carbons by Bandosz and Ania [102], activated carbons can play more than a secondary role in photochemical processes. Since 2010, works by Ania and co-workers have proven photoactivity of solar light irradiation is used.

The works reporting the use of semiconductor/AC composites for the photodegradation of CBZ, DCF, and SMX are listed in Table 4 and, again, TiO$_2$ is the semiconductor tested and in only one study solar light irradiation is used.

| PhC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|-------------------------|--------------------|------|
| SMX | TiO$_2$/PAC | • Composites: physical, mechanical and chemical mixtures  
• [TiO$_2$] = 0.5 g/dm$^3$ (Aeroxide P25)  
• [PAC] = 2.5–35 mg/dm$^3$ (commercial)  
• [PAC] = 0.5 mg/dm$^3$ (each PhC)  
• UV-A (1.0 mW/cm$^2$), UV-B (2.5 mW/cm$^2$), UV-C (3.65 mW/cm$^2$)  
• 60 min dark (ads)  
• Multi-solute (5 PhCs)  
• Deionized water, synthetic matrix and spiked real water (tap water, river water, sea water, wastewater) | • In deionized and river water UV-C allows higher overall removal  
• Photocatalyst deactivation in synthetic and spiked real waters (mainly with HCO$_3^-$ and in wastewater effluent)  
• In spiked sea water, ads-des yields comparable to deionized water  
• In deionized water physical mixture attains higher overall efficiency (~90%) than mechanical or chemical mixture composites (60–80%)  
• In deionized water, degradation by TiO$_2$ and TiO$_2$/PAC mixture fits the first-order kinetic model: fastest for DCF and slowest for SMX  
• Regardless of the type of water, PAC enhances overall efficiency and individual removal yield  
• DCF 100% removal independent of PAC addition  
• Cumulated kinetic constants (after dark) are worse than single PhC but still fit first-order model: 0.069 min$^{-1}$ in deionized water, 0.070 min$^{-1}$ in seawater, but only 0.029 min$^{-1}$ in wastewater | [107] |
| DCF | TiO$_2$ | | | |
| CBZ | TiO$_2$/PAC | • 1.5 wt. ratio TiO$_2$/PAC (agglomerates in suspension)  
• pH 7.5 (phosphate buffer)  
• [TiO$_2$] = 0.1 g/dm$^3$ (Aeroxide P25)  
• [PAC] = 0 - 20 mg/dm$^3$ (commercial - Norit)  
• [PAC] = 8 - 9 mg/dm$^3$ (each PhC)  
• UV light (Hg lamp)  
• Single-solute & deionized water | • Steam activated PAC (steam-PAC) originated higher amount of mixed agglomerates with TiO$_2$ in suspension: tested in a hybrid process  
• Steam-PAC did not allow CBZ degradation, neither did the other 3 PhCs  
• Steam-PAC addition seemed to inhibit the rate of CBZ degradation: initial adsorption (17 to 59%) led to lower degradation rate (i.e., 0.009-0.012 min$^{-1}$ (TiO$_2$+PAC) vs. 0.022 min$^{-1}$ (TiO$_2$))  
• High CBZ affinity for PAC may decrease mobility, preventing the contact between CBZ and TiO$_2$ resulting in a lower degradation rate. Turbidity increase due to PAC addition can also justify the low performance of the TiO$_2$/PAC mixture.  
• For a PhC with lower adsorption affinity (iomeprol), the detrimental effect of turbidity was compensated by the synergistic effect TiO$_2$/PAC | [108] |
| CBZ | TiO$_2$ | | | |
| CBZ | PAC | | | |
was counterbalanced by a slower photodegradation rate and the values of dissolved organic carbon in the water matrix. DCF and SMX present almost constant overall removal efficiencies regardless of the presence of the carbon particles, justify the low performance of the composite TiO$_2$/PAC when compared with the bare TiO$_2$. The initial removal of CBZ by adsorption was counterbalanced by a slower photodegradation rate and the values of dissolved organic carbon indicate that the rate of mineralization was much lower than the rate of degradation.

The enhancement of photoactivity through the immobilization of TiO$_2$ onto AC was proved by Alalm et al. that studied the degradation of DCF, among other PhCs, using a solar reactor in a competitive scenario. The removal efficiency for DCF was improved from 68% with bare TiO$_2$ to 85% using TiO$_2$/AC. Moreover, the composite also allowed total removals of amoxicillin and...
ampicillin. The photocatalytic degradation of the four PhCs in multi-solute conditions follows the Langmuir-Hinshelwood kinetic model. According to this mechanism, before degradation species must be adsorbed and this last process is faster for the TiO$_2$/AC composite regardless the PhC. Higher removal rates with TiO$_2$/AC were obtained in a large pH range (3–10), being the highest value observed for pH $\geq$ 7, which is, close to the values usually found in secondary effluent (wastewater). This is an advantage of this process in comparison with photo-Fenton (faster and complete degradation in 1–2 h) that is favored at pH 3, causing corrosion of mechanical facilities and the need for an extra neutralization step. Finally, according to the authors’ estimation, operating and total costs of using TiO$_2$/AC solar photocatalysis were lower than those for bare TiO$_2$ [109], thus highlighting the potential of these systems to be used in real water treatment facilities.

3.3. Semiconductor/CNT

Carbon nanotubes (CNTs) whose synthesis was reported by Iijima in 1991 [111] are another interesting class of carbon materials due to their outstanding structural, thermal and electronic properties associated with their one dimensional (1D) structure [73]. CNTs are classified as single-walled CNT (SWCNTs) and multi-walled CNT (MWCNTs) [111], presenting specific surface areas $> 150$ m$^2$/g, excellent mechanical properties and their high electron mobility can be compared to those of metals, i.e., they may exhibit metallic conductivity [76]. As highlighted in several reviews published over the last decade, the combination of all these attributes makes CNT superior supports to enhance semiconductors photoactivity [72,73,76,78,85,112].

Woan et al. summarized the three mechanisms proposed in the literature to explain the enhancement of photoactivity in TiO$_2$/CNT composites: (i) CNTs inhibit recombination, (ii) photosensitization due to e$^-$/h$^+$ pair generation in the CNTs, and (iii) CNTs act as impurities through Ti-O-C bonds [76]. In mixtures or nanocomposites semiconductor/CNT there are strong interfacial electronic effects between both players and the large electron-storage capacity of CNTs allows them to accept photon-excited electrons. As a consequence of the inherent electron density delocalization of the carbon matrix, the e$^-$/h$^+$ recombination is retarded or hindered [76,102]. MWCNT were reported to have self-photoactivity under visible light due to the presence of structural defects and vacancies [113], justifying a large number of studies using MWCNT instead of SWCNT [85]. The sensitizing role of MWCNT for visible light photoactivity was proposed by Wang et al. [87]: the electron photogenerated in the CNT is injected into the conduction band of the TiO$_2$ and the positively charged CNT remove an e$^-$ from the TiO$_2$ valence band creating a h$^+$, allowing the reaction with O$_2$ (e$^-$) and H$_2$O (h$^+$) to generate the oxygen reactive species (O$_2^-$ and OH$, $ respectively). The third mechanism is more complex since two distinct contributions of TiO$_2$/CNT are reported [76]. In the first one MWCNT act as carbon-doped TiO$_2$ since the Ti-O-C bond extends the light absorption to longer wavelengths with potential to improve photoactivity and solar-light harvesting. The second is related to the electronic configuration of the CNTs that justifies the highest performance of CNTs prepared by arc-discharge CNT over CVD-grown ones, due to higher electrical conductivity and less defects of the former [76].

Owing to their great potential the use of semiconductor/CNT composites for the photodegradation of organic contaminants has been of considerable scientific interest and the works focused on the photodegradation of CBZ, DCF and SMX are listed in Table 5. As in the previous sections, TiO$_2$ is the most tested semiconductor. Half of the works evaluated the performance of the composite under solar irradiation, and the majority use single-solute conditions and deionized water, not allowing the comparison of the behavior of the material, nor accounting for competition of other PhCs or organic matter.
**Table 5.** Photocatalytic degradation of CBZ, DCF, and SMX by semiconductor (single-walled carbon nanotubes, MWCNT—multi-walled carbon nanotubes).

| PhC          | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|--------------|----------|--------------------------|--------------------|------|
| SMX          | TiO<sub>2</sub>SWCNT | 0.58 mg TiO<sub>2</sub>/mg SWCNT [TiO<sub>2</sub>] = [TiO<sub>2</sub>/SWCNT] = 0.1 mg/dm<sup>3</sup> [TiO<sub>2</sub>—Aeroxide P25 and synthetic red-like nanocrystals] | PhC mixture photocatalysis follows the Langmuir-Hinselwood mechanism showing first-order kinetic model | [114] |
| DCF          | TiO<sub>2</sub> | Same conditions as CBZ | In ultrapure water under UV light TiO<sub>2</sub>/SWCNT is more effective than TiO<sub>2</sub> for DCF, SMX, and more PheCs; for the remaining 16 PheCs, (including CBZ) TiO<sub>2</sub>/MWCNT is comparable or slightly better | |
| CRZ          | TiO<sub>2</sub>/SWCNT | 30 min in dark (ads) Multi-solute (22 PhCs) Deionized water and raw water (secondary wastewater effluent) | In real wastewater effluent under UV light, lower degradation rates compared with values in ultrapure water in similar conditions | |
| SMX          | WO<sub>3</sub>/MWCNT | Hydrothermal synthesis: 400 mg Na<sub>2</sub>WO<sub>4</sub>·2H<sub>2</sub>O for 2, 4 or 8 mg MWCNT [WO<sub>3</sub> = [WO<sub>3</sub>/MWCNT] = 0.5 g/dm<sup>3</sup> (Commercial MWCNT) [SMX] = 10 mg/dm<sup>3</sup> Simulated solar light 420 – 630 nm 1 h dark (ads) + 1 h light (deg) Single-solute & deionized water | CNT content increase in the WO<sub>3</sub>/MWCNT composites enhances visible light absorption (red-shift) Band gaps: 2.80 eV (W0.1:2), 2.65 eV (WO<sub>3</sub>/MWCNT-2), 2.52 eV (WO<sub>3</sub>/MWCNT-4) and 2.32 eV (WO<sub>3</sub>/MWCNT-8) Similar photoluminescence emission spectra for bare WO<sub>3</sub> and composites, but composites have lower e<sup>−</sup>/h<sup>+</sup> recombination and charge separation is improved as the CNT content increases Removal under solar irradiation (3 h): WO<sub>3</sub> (25%) < composites (42-75%) WO<sub>3</sub>/MWCNT-4 composite is the smarter choice The higher the WO<sub>3</sub>/MWCNT-4 dose (0.25 - 2.00 g/dm<sup>3</sup>) the better the SMX removal efficiency (40-88%) After 4 reuse cycles, WO<sub>3</sub>/MWCNT-4 only lost ~5% removal efficiency All radicals contribute to the SMX degradation with WO<sub>3</sub>/MWCNT-4 under solar light, being OH<sup>−</sup> and O<sup>2</sup>− the most important ones Intermediates identified and 4 processes proposed for SMX photocatalytic degradation by WO<sub>3</sub>/MWCNT-4 | [115] |
| CRZ          | TiO<sub>2</sub>MWCNT<sub>ox</sub> | Mixture and sol-gel synthesis: 10.1 wt% titania to MWCNT<sub>ox</sub> [TiO<sub>2</sub>] = [TiO<sub>2</sub>/MWCNT<sub>ox</sub>] = 0.1-2 g/dm<sup>3</sup>, optimum 0.5 g/dm<sup>3</sup> (TiO<sub>2</sub>—Aeroxide P25 and lab-made (anatase and rutile), commercial MWCNT<sub>ox</sub>) [ZnO] = 0.5 g/dm<sup>3</sup> (com. Eronikov) [CBZ] = 8 mg/dm<sup>3</sup> pH = 6 (natural CBZ solution pH) 200 cm<sup>3</sup>/min O<sub>2</sub>/Ar (0-100 vol.% of O<sub>2</sub>, optimum 50 vol.% of O<sub>2</sub>) [H<sub>2</sub>O<sub>2</sub>] = 0-10 mM, optimum 5 mM UV-C and Near UV-Vis (NUV-Vis) 30 min dark (ads) + 1 h light (deg) Single-solute & deionized water | Composite TiO<sub>2</sub>/MWCNT<sub>ox</sub> absorbs at higher wavelengths than TiO<sub>2</sub> advantage considering sun-light harnessing Photolysis and photocatalytic degradation of CBZ upon UV-C irradiation, under NUV-vis light CBZ photolysis is negligible compared with photocatalysis Addition of O<sub>3</sub> favors faster CBZ photodegradation UV irradiation: (1) addition of MWCNT does not promote CRZ photocatalytic degradation (pure photochemical process) (2) TiO<sub>2</sub> efficiency: anatase > P25 > rutile NUV-vis irradiation (photocatalytic regime): (1) positive synergy for P25 and synthetic TiO<sub>2</sub> with 70% anatase and 30% rutile (2) CRZ photocatalytic degradation trend TiO<sub>2</sub>-P25 > mixture TiO<sub>2</sub> + MWCNT > TiO<sub>2</sub>/MWCNT<sub>ox</sub> > H<sub>2</sub>O<sub>2</sub> 5 mM > TiO<sub>2</sub>/MWCNT<sub>ox</sub> (3) ZnO degradation of CRZ 17% higher than with TiO<sub>2</sub>-P25 2 pathways proposed for CRZ photodegradation | [116] |
| CRZ          | TiO<sub>2</sub>SiO<sub>2</sub>/MWCNT<sub>ox</sub> | Sol-gel synthesis: composites with 0.15 – 17.8 wt.% TiO<sub>2</sub> [Catalyst] = 0.5 g/dm<sup>3</sup> [P:HC] = 10 mg/dm<sup>3</sup> (each PhC) TiO<sub>2</sub>—Aeroxide P25 for comparison UV (1.0 mW/cm<sup>2</sup>) 30 min dark (ads) + 2 h light (deg) Single-solute | Composites with anatase TiO<sub>2</sub> (7-8 nm): T<sub>g</sub> reduces from 3.2 eV to 2.2 eV as CNT content increases and visible light harvest improved Up to 3.5 wt.% CNT act as a dopant in TiO<sub>2</sub>SiO<sub>2</sub>, for higher CNT content TiO<sub>2</sub>SiO<sub>2</sub> crystals are supported on the outer CNT surface Faster degradation with the composite presenting 17.8 wt.% of CNT (Pseudo-first order kinetic constant 0.0131 – 0.0743 min<sup>−1</sup>) Distinct decomposition pathway with TiO<sub>2</sub>-P25 and composites Addition of CNT during TiO<sub>2</sub>SiO<sub>2</sub> synthesis: enhances TiO<sub>2</sub> activity, changes CBZ degradation mechanism, and transformation products in model wastewater have low toxicity to D. magna and V. fischeri | [117] |
| DCF          | TiO<sub>2</sub>MWCNT<sub>ox</sub> | Same catalyst ref. [116] [TiO<sub>2</sub>] = [TiO<sub>2</sub>/MWCNT<sub>ox</sub>] = 0.1-2 g/dm<sup>3</sup>, optimum 1 g/dm<sup>3</sup> [DCF] = 8 mg/dm<sup>3</sup> pH = 6 (natural DCF pH) 200 cm<sup>3</sup>/min O<sub>2</sub>/Ar [H<sub>2</sub>O<sub>2</sub>] = 0-5 mM, optimum 5 mM UV-C and Near UV-Vis (NUV-Vis) 30 min dark (ads) + 1 h light (deg) Single-solute & deionized water | Addition of O<sub>3</sub> favors faster DCF photodegradation Complete photolytic and photocatalytic degradation of DCF upon UV-C and NUV-vis light after 30 min irradiation UV irradiation: DCF degradation trend anatase > rutile > TiO<sub>2</sub>/MWCNT<sub>ox</sub> > no catalyst > TiO<sub>2</sub>-P25 NUV-vis irradiation (photolytic + photocatalytic regime): DCF photocatalytic degradation trend TiO<sub>2</sub>-P25 > anatase > no catalyst > TiO<sub>2</sub>/MWCNT<sub>ox</sub> > rutile Considering the 8 identified intermediates the photocatalytic degradation of DCF is proposed | [118] |
In 2015 Murgolo et al. tested the multi-solute photocatalytic degradation of 22 PhCs, in the ppm concentration range, under UV-C and simulated solar light in deionized water and secondary wastewater effluent. The first-order constant rates for DCF, SMX, and CBZ are summarized in Figure 5 [114]. In the most favorable experimental conditions—UV irradiation and deionized water—for DCF and SMX the first-order constant rates obtained with composite TiO\textsubscript{2}/SWCNT are much higher than with TiO\textsubscript{2} P25, while for CBZ TiO\textsubscript{2} similar values were observed. For secondary wastewater effluent irradiated with UV-C light the photodegradation rates of DCF and SMX are also favored with the composite and a detrimental effect of the real wastewater is clearly shown. Regarding simulated solar light irradiation, the only interesting results were those obtained with TiO\textsubscript{2} in ultrapure water. The photodegradation rate of the three PhCs addressed in this review follows the order DCF > SMX > CBZ [114]. So, in line with the high persistence of CBZ in the environment, due to the low photolysis profiles in water streams, according to these results CBZ is also more resistant to photodegradation regardless the irradiation source and water matrix. The reuse of TiO\textsubscript{2}/SWCNT was tested for five cycles with five PhCs, including SMX and CBZ, under UV light. No variation of rate constants were observed for all but ibuprofen. Lastly, it is important to stress that this work clearly demonstrates the importance of performing studies in real water matrices since the results obtained in synthetic solutions do not allow a direct translation to water treatment facilities.

![Figure 5](image_url)

**Figure 5.** Effect of irradiation light and water matrix in first order kinetic constants (k) for DCF, SMX, and CBZ multi-solute photodegradation by TiO\textsubscript{2} and TiO\textsubscript{2}/SWCNT (data from [114]).

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**Table 5.**

| PhC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|-------------------------|-------------------|------|
| DCF | SiO\textsubscript{2}-TiO\textsubscript{2}/MWCNT | Sol-gel method: (0.01 wt.% MWCNT) basic and acid conditions, calcined in air (400°C) | All prepared catalysts were more effective than TiO\textsubscript{2} P25 for the adsorption-degradation of DCF under UV-A and solar light | [119] |
| DCF | SiO\textsubscript{2}-TiO\textsubscript{2} | [Catalyst] = 0.5 g/dm\textsuperscript{3} | Regardless of the light source, MWCNT alone removes \( \approx 50\% \) DCF by adsorption and \( \approx 30\% \) more by photocatalytic degradation | |
| DCF | TiO\textsubscript{2} lab-made and P25 (Evonik) | [DCF] \textsubscript{0} = 10 mg/dm\textsuperscript{3} | In composites, MWCNT mainly contributes to degradation | |
| DCF | UV-A and visible light | | SiO\textsubscript{2}-TiO\textsubscript{2} presented higher photocatalytic activity than catalysts doped with MWCNT and are more active under visible light than under UV-A | |
| DCF | 30 min dark (ads) + 1 h light (deg) | | After treatment with SiO\textsubscript{2}-TiO\textsubscript{2} and regardless of the light source the bioluminescent inhibition of *V. fischeri* decreased to values \( \approx 20\% \) while the toxicity of the solution before treatment was \( \approx 90\% \) | |

ads-adsorption, deg-degradation
Other studies listed in Table 5 systematically report enhanced visible light harvesting for the composites’ semiconductor/CNT when compared with the bare semiconductor [115–119].

Zhu et al. prepared composites WO₃/MWCNT with increasing CNT contents verifying that higher CNT amount lower both the band gap energy and e⁻/h⁺ recombination. The composite WO₃/MWCNT-4 (4 g of MWCNT) presented the best compromise between the band gap energy (2.52 eV), the dispersion in water and the amount of MWCNT [115]. SMX degradation efficiency under simulated solar light followed the order WO₃ (25%) < WO₃/MWCNT-2 (42%) < WO₃/MWCNT-4 (65%) < WO₃/MWCNT-8 (73%). The composite WO₃/MWCNT-4 only lost 5% efficiency after four reuse cycles [115]. The mechanism proposed for the SMX photocatalytic degradation under visible light irradiation involves the transference of the photo excited electron from the WO₃ conducting band to the CNT which acts as an electron trap to hinder e⁻/h⁺ recombination. The electron will react with O₂ to produce O₂•⁻, while the hole reacts with water and/or surface hydroxyl to generate OH•, these radicals will decompose the SMX [115].

Martinez et al. used TiO₂/MWCNTₓ and TiO₂ for the photocatalytic degradation of CBZ [116] and DCF [118] under UV-C and near UV-Vis irradiation (single-solute and deionized water). The TiO₂/MWCNTₓ with a 10:1 weight ratio absobes at higher wavelengths than TiO₂ (P25, anatase or rutile) allowing improved sunlight harnessing [116,118]. The addition of 21% of O₂ (V/V) contributes to a huge increase in the degradation rate of CBZ and DCF by P25 leading to half-life times lower than 5 min under near UV-vis light [116,118]. The faster degradation of the PhCs is attributed to the reaction between them and HO₂•/O₂•⁻, and/or to the strong electrophilic character of dissolved O₂ which may reduce unfavorable e⁻/h⁺ recombination. Albeit the improved sunlight harvesting, compared with anatase for UV irradiation or P25 for near UV-Vis irradiation, the composite has worst photoactivity for CBZ and DCF degradation (UV-C or near UV-vis), possibly due to OH• inhibition in the presence of MWCNTs. The fact that CBZ and DCF are electron-rich molecules that may transfer e⁻ to MWCNT conducting band may also justify the slower oxidation process [116,118].

Czech & Buda tested nanocomposites TiO₂-SiO₂/MWCNT with 0.01% to 17.8% of CNT against TiO₂-SiO₂ for the photocatalytic degradation of CBZ [117] and DCF [119]. The addition of SiO₂ promotes the dispersion of TiO₂ [119] while the addition of MWCNT steadily reduces the band-gap of the composite TiO₂-SiO₂/MWCNT from 3.2 eV to 2.2 eV when the CNT content increases from 0.15% to 17.8%, shifting absorption to visible [117]. CNT acts as a dopant for up to 3.5 wt.% CNT but as support for higher concentrations. Under UV-A irradiation TiO₂-SiO₂/MWCNT with 17.8 wt.% of CNT allows faster CBZ degradation, through a decomposition pathway distinct from that obtained by P25, and the formation of degradation products with low toxicity to D. magma and V. fisheri. Both composites with TiO₂-SiO₂/MWCNT with 0.01 wt.% of CNT was effective than bare P25 for the adsorption and degradation of the DCF under UV-A or solar light. Most probably due to their small weight percentage in composites, MWCNTs mainly contributes to the degradation step [119]. However SiO₂-TiO₂ presented even higher photocatalytic activity than those doped with MWCNT, being more active under visible light than under UV-A and allowing a decrease of bioluminescent inhibition of V. fisheri (from > 90% to values ≤ 20%) [119]. It must be also highlighted that MWCNTs are effective solar-responsive metal-free photocatalysts for DCF degradation regardless the light source (UV-A and visible). In fact MWCNTs removed ≥ 50% DCF by adsorption plus more 30–40% by photocatalytic degradation [119].

3.4. Semiconductor(r)GO

Graphene is a two-dimensional (2D) sheet of carbon atoms connected by sp² bonds with an aromatic π electron system [120], it has a high thermal conductivity (≈5000 W/(m × K)), allows excellent mobility of charge carriers at room temperature (2000 cm²/(V × s)), high optical transmittance (≈97.7%), and has an extremely high theoretical specific surface area (≈2600 m²/g) [121,122]. The first report of graphene synthesis and properties dates from 2004 by Novosevov et al. and since then graphene has...
been extensively explored by the scientific community in several applications, and it also caught the attention of industry.

Graphene is the precursor of graphene oxide (GO) and reduced graphene oxide (rGO), both showing high efficiency for water treatment due to their surface functionalities (e.g., hydroxyl, carboxyl and epoxy) [72]. Due to the exceptional features of graphene and derived nanomaterials, these solids are attractive for many applications mainly in environmental protection and energy production/storage. The use of graphene-based semiconductor photocatalysts has been a hot research topic since graphene joins the most promising properties of CNTs and activated carbons, i.e., the high electron mobility with high pore structure and adsorption capacity. The advances in graphene-based semiconductor photocatalysis have been addressed in several review papers [72,73,80–84,121–123].

The enhanced photoactivity of semiconductor/graphene composites can be understood in light of the mechanisms previously described for semiconductor and activated carbon or CNT composites. The advantages and shortcomings of the use of activated carbons, CNTs and graphene on the semiconductor composites were systematized by Awfa et al. [72] identifying that easy recovery from reaction media follows the trend activated carbon > CNT > graphene, while the overall process cost follows the opposite trend. High mechanical strength and improved thermal stability, large electron storage capacity, and superior metallic conductivity are some of the advantages of adding CNT and graphene to semiconductors. Due to graphene’s 2D structure, the interfacial contact between graphene and the semiconductor usually provides higher photoactivity [72]. The more difficult separation of the semiconductor/graphene composites can be overcome during the synthesis to obtain, for example, 3D aerogels [124], composites immobilized onto a solid supports [125] or adding magnetic features [126]. As reviewed by Dong et al. [127] the enhanced visible light photocatalytic activity of semiconductor/graphene composites is attributed to (i) the improved $e^-/h^+$ mobility and suppression of the photo-generated $e^-/h^+$ recombination, (ii) to their higher surface adsorption capacity for organic molecules through $\pi-\pi$ interactions, and (iii) full and intimate contact through chemical bonds of the metal semiconductor with carbon and oxygen atoms.

Certainly due to their potential graphene-based materials were the most representative group of composites studied for the photocatalytic degradation of CBZ, DCF, and SMX. It must be also noted that the majority of studies listed in Table 6, were tested under under visible irradiation, and only two works evaluate the performance of the photocatalysts in real water. In all but one study TiO$_2$ was the semiconductor selected.

| PhC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|-------------------------|--------------------|-----|
| SMX | TiO$_2$/GO (0.1–10 wt.% GO:TiO$_2$) coated on optical fibers | Without catalyst: CBZ removal is negligible while SMX decreases 30% | [125] |
| CBZ | TiO$_2$/rGO (0.1–10 wt.% GO:TiO$_2$) coated on optical fibers | Without catalyst: CBZ removal is negligible while SMX decreases 30% | [125] |
| | Catalyst: 30 fibers of 10 cm coated with composite and TiO$_2$ (synthesized and Aeroxide P25) | Photocatalytic activity: synthesized TiO$_2$ < TiO$_2$ P25 < TiO$_2$/rGO-2.7% (> 50% CBZ and > 90% SMX removal) | [125] |
| | [PhC]$_0$ = 5 mg/dm$^3$ (each PhC) | While SMX removal is more effective than of CBZ (> 90% vs. ~ 40%) both reach similar mineralization (54–59%) after 3 h irradiation: faster SMX degradation but intermediates need similar time as those of CBZ to achieve mineralization | [125] |
| | pH 6 | TiO$_2$/rGO-2.7% durability assessed for ibuprofen during 15 cycles (45 h total contact time) under UV-vis irradiation: >80% removal | [125] |
| | UV-vis (UV-B, UV-A and visible) | | |
| | 3 h dark (ads) + 3 h light (deg) | | |
| | Single-solute & desonized water | | |
Table 6. Cont.

| PhC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|--------------------------|-------------------|------|
| SMX | TiO$_2$/rGO | Hydrothermal synthesis | TiO$_2$/rGO is an efficient activator of persulfate for visible light SMX degradation: 52% SMX degradation (0.055 min$^{-1}$) and 26% mineralization | [129] |
| SMX | TiO$_2$/GO | Hydrothermal synthesis | TiO$_2$/GO and TiO$_2$/rGO improve light harvesting and e$^-$ for persulfate activation | [129] |
| SMX | TiO$_2$/GO | Microwave hydrothermal method: 1 – 10% GO and 0.2 mg TiO$_2$ P25. Aerioxide | Enhancing visible light harvesting and e$^-$ for persulfate activation | [130] |
| CBZ | TiO$_2$/GO | Microwave hydrothermal method: Bioactivity of CBZ degradation, 1% and 1.5% allow higher first-order kinetic rates than P25. Higher amounts of GO have a detrimental effect on photodegradation. | | [130] |
| CBZ | TiO$_2$/GO | Physical mixture or hydrothermal synthesis (1:1 to 4:1 wt.% ratio TiO$_2$/GO, with L-ascorbic acid) | The composite prepared ex-situ (direct use of TiO$_2$ nanoparticles) with 2:1 wt.% TiO$_2$/GO presented the best performance (=99% CBZ degradation, 0.0473 min$^{-1}$ pseudo-first-order rate constant) | [124] |
| CBZ | TiO$_2$/rGO | Physical mixture or hydrothermal synthesis (1:1 to 4:1 wt.% ratio TiO$_2$/GO, with L-ascorbic acid) | The composite prepared ex-situ (direct use of TiO$_2$ nanoparticles) with 2:1 wt.% TiO$_2$/GO presented the best performance (=99% CBZ degradation, 0.0473 min$^{-1}$ pseudo-first-order rate constant) | [124] |
| DCF | TiO$_2$ nanotubes/Pd-rGO | Photoelectrode prepared by electro-deposition | The TiO$_2$ nanotubes/Pd-rGO photoelectrode has enhanced photocurrent density and charge carrier concentration, attaining 58.4% for DCF degradation under visible light | [131] |
| DCF | Ag-BiO$\text{I}$/rGO | Hydrothermal synthesis | Ag-BiO$_2$/rGO is the best performing catalyst (0.026 min$^{-1}$), attaining 100% DCF removal and 55.8% mineralization in 80 min (visible light) | [132] |
Lin et al. synthesized TiO$_2$/rGO with rGO weight percentages 0.1–10% coated and tested the photoactivity of the composites for the photocatalytic degradation of three PhC, including SMX and CBZ, under UV and visible light [125]. The best performing photocatalyst was TiO$_2$/rGO-2.7% due to (i) lower band gap energy, (ii) formation of a heterojunction interface leading to improved charge separation [125]. Under UV-vis irradiation SMX removal is more effective than CBZ (92% vs. 54%) but both PhCs attain similar mineralization degrees (54–59%) indicating that although SMX presents faster degradation its oxidation intermediates need similar time as those of CBZ to achieve mineralization [125]. The immobilization of the composite onto optical fibers proved to be a promising strategy since after 15 cycles (i.e., corresponding to 45 h) under UV-vis irradiation the ibuprofen removal efficiency by TiO$_2$/rGO-2.7% was over 80% [125].

SMX photocatalytic degradation by TiO$_2$/rGO composites under visible light was also tested by Yang et al. demonstrating that TiO$_2$/rGO is an activator of persulfate for SMX and other PhCs degradation through SO$_4^{2-}$ and OH• [129]. Karaolia et al. studied SMX adsorption and photocatalytic degradation in spiked real urban wastewater effluent verifying that bare P25 is more efficient that TiO$_2$/rGO prepared photocatalitically or by hydrothermal synthesis [128].

Three studies report CBZ photochemical degradation using single-solute conditions and UV irradiation. The best performing TiO$_2$/graphene composite attained higher CBZ degradation than bare TiO$_2$ [124,126,130] and adding magnetic properties [126] or 3D graphene structure [124] allowed several reuse cycles. Appavoo et al. [130] and Nawaz et al. [124] concluded that there is an optimum percentage of GO or rGO to assure the best compromise between CBZ adsorption and photocatalytic degradation. Appavoo et al. report mineralization degrees of 80% when composites with 1% and 2% of GO are used, while P25 only reaches 64%.

Metals or peroxodisulfate were used as additive to improve photocatalytic degradation of DCF with semiconductor/graphene composites under visible light [131–133]. Cheng et al. conclude that both Pd and rGO in the TiO$_2$ nanotubes/Pd-rGO contribute to the improved light harvesting and effective separation of the photogenerated charge carriers, and identify OH• as the most relevant radicals for DCF degradation [131]. Li et al. reported 100% DCF removal and 55.8% mineralization when irradiating Ag-BiOI/rGO under visible light during 80 min, maintaining the performance after threereuse cycles [132]. The authors propose that Ag enhance the separation e–/h$^+$ while the rGO with extended π-electron conjugations has strong charge transfer capability and is the potential sink of e–. Moreover, rGO can function as e– acceptor, and transfer due to its excellent charge carrier conductivity [132]. The work by Chen et al. followed a more applied perspective by evaluating the effect of the water matrix and solution pH in the DCF degradation by TiO$_2$/rGO in the presence of peroxodisulfate under visible light [133]. The authors verified that the increase of solution pH has a detrimental effect on DCF degradation rate and environmental factors also interfere in the degradation.
The presence of Cl\(^{-}\) facilitates DCF degradation but the presence of HCO\(_3\)\(^{-}\) as well as tap water, lake water, river water matrices decrease DCF degradation. In this photodegradation system both O\(_2\)\(^{•-}\) and h\(^+\) play significant roles in DCF degradation, peroxodisulfate behaved like an electron acceptor to enhance e\(^-\)/h\(^+\) separation and generation of additional reactive oxygen species, and rGO served as electric conductor, so both additive increase photocatalytic efficiency of the composite for DCF degradation [133].

3.5. Semiconductor/Carbon Quantum Dots (CQDs)

Carbon quantum dots (CQDs) were firstly reported by Xu et al. [134] in 2004 just a few months earlier than graphene, but these fluorescent carbon nanoparticles only received their most common name – CQD – from Sun et al. in 2006 [135]. CQDs are approximately flat or quasi-spherical carbon nanoparticles (<10 nm) composed by amorphous to nanocrystalline cores with predominantly sp\(^2\) carbon (i.e., graphitic carbon and/or graphene and GO sheets) fused by diamond-like sp\(^3\) hybridized carbon intersections [136,137]. CQDs have excellent aqueous solubility due to the considerable amounts of carboxylic surface moieties, presenting oxygen content between 5 and 50% [135–137]. CQDs have demonstrated capability of harnessing long wavelength light and their ability to exchange energy with solution species offers excellent opportunities for their use as photocatalysts [136]. Furthermore, CQDs present up-conversion fluorescent emission, an optical phenomenon wherein the fluorescence emission wavelength is shorter than the used excitation wavelength. This property is particularly important for in vivo bioimaging since near IR (NIR) region is usually preferred [136] but is also a very valuable feature to improve semiconductor photoactivity under full sunlight spectrum [137]. In fact, CQDs proved to be excellent additives in solar photocatalytic systems since their ability to harness the full spectrum of sunlight and up-convert visible light into shorter wavelength light triggers the semiconductor charge separation (i.e., e\(^-\)/h\(^+\) generation and hinders recombination) which by reaction with H\(_2\)O/O\(_2\) will produce the active oxygen species needed to promote contaminants degradation [136,137]

Regarding the target PhCs, to the best of our knowledge, so far semiconductor/CQD composites were only explored for the photodegradation of DCF (Table 7). The three very recent studies explore the solar light photoactivity of the composites in single-solute conditions, and only one evaluates the influence of distinct water matrices.

Table 7. Photocatalytic degradation of DCF by semiconductor/CQD.

| PhC  | Catalyst Experimental Conditions | Concluding Remarks | Ref. |
|------|---------------------------------|--------------------|-----|
| DCF  | BiOCOOH/CQD TiO\(_2\)          | Composites obtained by ultrasonic dispersion (3h at 180 °C) | BiOCOOH adsorbs at >360 nm (UV) but the incorporation of 1–5% CQDs redshifts the adsorption spectra and allows superior visible light adsorption (\(E_g = 3.42–2.81\) eV) | [138] |
|      | [Catalyst] = 0.6 g/dm\(^3\)     | Under visible light BiOCOOH/CQD with 2.0 wt.% of CQD reaches 98% DCF degradation while the bare BiOCOOH only attains 51.5% degradation, the commercial P25 degrades less than 15% of the DCF |     |
|      | [TiO\(_2\)-Aeroxide P25]       | CQD on the surface of BiOCOOH operates as photosensitizers |     |
|      | pH 7                            | O\(_2\)\(^{•-}\) and holes contribute for the higher BiOCOOH/CQD photoactivity |     |
|      | Kinetic, by-product and toxicity: Visible light (350 W Xe lamp, 1.15 mW/cm\(^2\)) with 420 nm cut-off | In 2 h under visible light BiOCOOH/CQD mineralize, detoxify (V. fisheri, D. magna, and D. subspicatus tests) and dechlorinate DCF |     |
|      | Photocatalyst mechanism tests: UV-A (390–400 nm, 1.6 mW/cm\(^2\)) Blue (455–460 nm, 3.84 mW/cm\(^2\)) Green (515–530 nm, 4.5 mW/cm\(^2\)) Red (655–660 nm, 4.07 mW/cm\(^2\)) | 90% DCF degradation after 4 CQD/BiOCOOH reuses under visible light |     |
|      | Near-IV (750 nm, 3.1 mW/cm\(^2\)) | Intermediaries identified and degradation mechanisms proposed |     |
|      | 30 min dark (ads) + 1–2 h light (deg) | Single-solute |     |

*Note:* Table 7 includes the experimental conditions and concluding remarks for the photodegradation of DCF using semiconductor/CQD composites. The table highlights the effectiveness of CQDs in enhancing the photocatalytic activity of semiconductors, particularly in the visible light range, and their ability to mineralize and detoxify contaminants.
Chen et al. report the superior performance of BiOCOOH/CQD composites under visible-light for the degradation of DCF when compared with P25 [138]. The incorporation of CQDs allows the composite to absorb in the visible range presenting band gaps 3.42–2.84 eV, as the percentages of CQDs increase from 1 to 5%. The best photocatalytic activity is obtained for BiOCOOH/CQD-2% (98% DCF removal vs. 51.5% for BiOCOOH and only 15% for P25), allowing to attain 90% DCF even after four reuse cycles. This behavior was mainly attributed to the generation of O$_2$•• and holes [138]. It is also remarkable that in just 2 h this composite effectively mineralized, detoxify and dechlorinaited DCF, and it also presented high performance for the degradation of the antibiotic sulfadimidine (>90%) and rhodamine B (>75%), proving its potential for application in water remediation strategies. Authors propose CQDs act as photosensitizers of the BiOCOOH, with the electron transfer properties
of the CQDs being beneficial to constrain the $e^-/h^+$ recombination and their up-conversion properties contributing to enhancing the visible photoactivity of the semiconductor [138].

In a more sustainable approach, Liu et al. tested metal free composites $g$-$C_3N_4$/CQD for the visible light photodegradation of DCF [140]. Once again, the introduction of optimized amounts of CQDs improved visible light harvesting and increased $e^-/h^+$ separation. Composite $g$-$C_3N_4$/CQD-0.05 showed the fastest total DCF removal just 1 h irradiation and degrades 90% of the DCF after five reuse cycles [140]. Alkaline medium favors a faster DCF degradation. $O_2^•−$ plays the dominant role in the DCF degradation process which is not dependent on the holes, which allowed the proposal of a photosensitation-like mechanism. After 1.5 h of irradiation of $g$-$C_3N_4$/CQD-0.05, the DCF mineralization reached 54% and although, in general, the intermediates are less toxic than the parent compound they are still classified as very toxic, pointing out the need of longer irradiation times [140].

In 2017, Song et al. used $g$-$C_3N_4$ for the photodegradation of sulfonamides, including SMX, under visible light attaining 90% removal under distinct water quality parameters [141]. SMX was degraded by $O_2^•−$ and holes, and the degradation pathways were proposed.

Wang et al. compare the efficacy of composites of $TiO_2$ nanosheets with two distinct surface energy facets or P25 with N-doped CQD for DCF degradation under broad spectrum irradiation [139]. The results reveal that TiO$_2$ facets play a major role in the photodegradation process with the superior performance of TiO$_2$(001)/N-CQD. This behavior was attributed to the synergistic effect of the high oxidation activity of exposed (001) facets, up-conversion properties of the N-CQDs and efficient charge separation in the direct Z-scheme heterojunction [139]. Under visible light, composites with TiO$_2$(101) and TiO$_2$(001) photodegrade, respectively, around 80% and more than 90% of DCF after 1 h of irradiation, while composite with P25 and the three bare TiO$_2$ samples degraded less than 20% of DCF [139]. Despite the high DCF removal by TiO$_2$(001)/N-CQD, after 1 h of visible light irradiation, only 50% mineralization was achieved. After four reuse cycles, the composite effectively degrades 80% of DCF. Contrarily to what was reported by Liu et al. [140] for $g$-$C_3N_4$/CQD, with this composites the increase of solution pH from 3 to 11 has a detrimental effect on the DCF degradation rate (from 0.09 to 0.01 min$^{-1}$) [139]. The influence of the water matrix was studied showing that the presence of transition metals and humic acids restrain the photodegradation of DCF. On the contrary the assays in natural waters (river water, sea water, and wastewater effluent) show that photocatalytic degradation of DCF is only slightly inhibited (84.0–84.6% vs. 90.9%, in deionized water) [139].

3.6. Semiconductor/Char

Chars are typically obtained by the carbonization of biomass through the removal of other elements than carbon, and transformation of the precursor in an amorphous carbon-rich material usually named biochars [142]. Chars can also be designated as pyrochars, hydrochars or acid-chars depending on the carbonization process, i.e., pyrolysis, hydrothermal, or acid-mediated carbonization, respectively [142,143]. Chars have incipient pore networks and depending on the precursor and synthesis procedure (i.e., carbonization method, temperature, and time) it is possible to tailor their surface chemistry from highly acidic to basic surface chemistries rich in oxygen groups or even other electron rich atoms (e.g., nitrogen, sulfur, or phosphorus) in order to meet with the needs of a specific process. The oldest known use of char (carbonized wood, coal, or partially devolatilized coals), also known as charcoal, dates to the Stone Age when the material was used as black pigment for cave painting [144]. Greeks and Romans used charcoal in the treatment of various diseases, and Hindus for water treatment. In the nineteenth century, it played a crucial role in sugar refining, and as gas filters in London sewage ventilation systems [142]. Recently Chen et al. compared the photoactivity of a pyrochar vs. a hydrochar, obtained from the same precursor, verifying that due to the abundant photocative oxygenated surface groups, the hydrochar could generate more reactive species under sunlight thus enhancing the degradation of the target pollutant (sulfamidimine) [143].

Along with the carbon materials previously discussed, biochars have also been tested to enhance semiconductor activity. The works focused on SMX and CBZ photodegradation under UV or visible
light irradiation are gathered in Table 8. All the studies were performed in single-solute conditions and in one case the photocatalytic activity of the semiconductor/char composites in deionized water and spiked river water were compared.

Xie and co-workers reported 80% SMX removal after 3 h under simulated visible light using a Zn-TiO$_2$/biochar composite [145]. This composite outperformed TiO$_2$/biochar and bare TiO$_2$, which was explained considering that Zn-doping effectively reduced the agglomeration of TiO$_2$ and decrease the crystal size. On the other hand, the higher photocurrent of Zn-TiO$_2$/biochar justifies the higher charge transfer and separation [145]. The photoactivity of Zn-TiO$_2$/biochar slightly decreases after the first cycle but remains almost constant in the next four reuses. Water matrix has a crucial role in the composite performance: in spiked river water only 53.83% SMX removal, the presence of SO$_4^{2-}$, Cl$^-$, and NO$_3^-$ also has an inhibitory effect possibly due to trapping of OH$^.$ or capture of h$^+$ to hinder production of OH$^.$ [145].

CBZ photodegradation was studied by Li et al. with magnetic Fe$_3$O$_4$/BiOBr/biochar, Fe$_3$O$_4$/BiOBr, and bare BiOBr [146]. At pH 6 Fe$_3$O$_4$/BiOBr/biochar (10% biochar, 0.05 g Fe$_3$O$_4$) is the best performing photocatalyst: highest rate constant (0.01777 min$^{-1}$), CBZ removal similar to that of BiOBr (95%), 70% mineralization (only $\approx$ 30% for BiOBr, $\approx$ 55% for Fe$_3$O$_4$/BiOBr). The species OH$^.$, h$^+$ and O$_2^-$ seem to take part in the CBZ photodegradation, suggesting that oxygen radicals play the most important role. After five reuse cycles under visible irradiation, Fe$_3$O$_4$/BiOBr/biochar degraded 90% of the CBZ, and in the fourth cycle more than 60% mineralization is achieved. Like in the previous study the presence of Cl$^-$ and SO$_4^{2-}$ has detrimental effect on CBZ degradation. Humic acid has opposite effects since while they are present at low concentration photodegradation decreases but at higher concentrations they behaved as photosensitizers promoting CBZ photodegradation [146].

Table 8. Photocatalytic degradation of CBZ and SMX by semiconductor/char.

| PbC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|-------------------------|-------------------|-----|
| SMX | Zn-TiO$_2$/biochar | • Sol-gel method: Zn(NO$_3$)$_2$ (1%, 10% & 15%), 1 g of pretreated reed straw biochar and 20 cm$^3$ tetrabutyl titanate (calcined at 300 °C)  
• [Catalyst] = 0.625–2.5 g/dm$^3$  
• [SMX]$_0$ = 10 mg/dm$^3$  
• pH 2.0–10.97  
• Simulated visible light (50 W/Xe lamp, >420 nm)  
• 30 min dark (ads) + 3 h light (deg)  
• Single-solute  
• Deionized water & spiked river water | SMX removal: Zn-TiO$_2$/biochar $>$ TiO$_2$/biochar $>$ TiO$_2$, up to 80% SMX removal in 3 h and $\approx$55% COD  
Zn-TiO$_2$ biochar with Ti/Zn mass ration of 1:1 is the best performing material for degrading SMX (0.085 min$^{-1}$)  
pH 5 and 1.25 g/dm$^3$ catalyst allow faster and higher SMX removal  
Assays in spiked river water reveal decreased efficiency of Zn-TiO$_2$/biochar (53.83%) compared with deionized water ($>80\%$)  
Common anions (e.g., SO$_4^{2-}$, Cl$^-$ and NO$_3^-$) have an inhibition effect on SMX degradation possibly due to OH$^.$ trapping or capture of h$^+$ (hinder OH$^.$ production)  
Zn-TiO$_2$/biochar photoactivity slightly decreases after 1st cycle but remains almost constant in the 4 next ones  
SMX photodegradation mechanism by Zn-TiO$_2$/biochar under visible light proposed, degradation products identified and 4 possible degradation pathways proposed | [145] |
| CBZ | Fe$_3$O$_4$/BiOBr/Biochar  
Fe$_3$O$_4$/BiOBr  
BiOBr | • One-step hydrolysis method: 5%, 10%, 20% and 30% (in wt.%) reed straw biochar  
• [Catalyst] = 1 g/dm$^3$  
• [CBZ]$_0$ = 10 mg/dm$^3$  
• pH 3–10  
• Visible LED light (50 W, 475 nm)  
• 1 h dark (ads) + 3 h light (deg)  
• Single-solute | Amounts of Fe$_3$O$_4$ and biochar influence the photoactivity of the composite for CBZ degradation under visible light during 3 h: 95% removal attained for 10% biochar and 0.05 g of Fe$_3$O$_4$  
At pH 6 Fe$_3$O$_4$/BiOBr/biochar is the best performing photocatalyst: highest rate constant (0.01777 min$^{-1}$), CBZ removal similar to that of BiOBr (95%) associated to mineralization degree of 70% (only $<30\%$ for BiOBr and $\approx$55% for Fe$_3$O$_4$/BiOBr)  
Optimum catalyst dosage is 1.0 g/cm$^3$ and solution pH is 7.1: faster (0.02292 min$^{-1}$) and almost 100% CBZ removal  
Cl$^-$ and SO$_4^{2-}$ inhibit CBZ photodegradation  
Humic acids role depends on concentration: low conc. $-$ detrimental, high conc. $-$ photosensitizers promoting CBZ photodegradation  
After 5 reuse cycles Fe$_3$O$_4$/BiOBr/biochar degrades 90% CBZ under visible light, after 4 reuse cycles mineralization $>60\%$  
OH$^.$, h$^+$ and O$_2^.$ seem to take part in the CBZ photodegradation with a suggestion that oxygen radicals play the most important role  
15 main intermediates and 2 ring-rupturing products identified, 4 possible reaction pathways proposed | [146] |
Although in optimized conditions high and fast degradation of the target pollutants can be attained, pore structure, the 42% CBZ removal may result from the combined adsorption and photocatalytic photodegradation under UV light irradiation in the presence of \( O_3 \) degree or evaluated the toxicity of the solution after photocatalysis.

mainly attributed to adsorption (nanoporous solid), in the case of the biochar, that has an incipient surface area that allows adsorbing CBZ that is further photodegraded by the reactive oxygen species produced by the semiconductor [148].

Zhu et al. propose that continuous radical attack to the benzene and isoxazole rings leads to bond cleavage [115]. The degradation products identified present m/z down to 99, being mainly substituted aromatic compounds. However, none of the studies report the mineralization degree or evaluated the toxicity of the solution after photocatalysis.

### Table 8. Cont.

| PhC | Catalyst | Experimental Conditions | Concluding Remarks | Ref. |
|-----|----------|-------------------------|--------------------|------|
| CBZ | TiO\(_2\)/biochar | • Sol-gel method: TiO\(_2\)/coconut shell powder char (34.04 cm\(^3\))
• titanium-n-butoxide and 60 – 120 g biomass
• Pellets: mixing composites with 10% wt. of wheat flour, calcined at 500–800 °C for 2 h
• [Catalyst] = 60–120 g/dm\(^3\)
• [CBZ] = 10 mg/dm\(^3\)
• 24 g/dm\(^3\) \( O_2 \) flow (bottom-to-top)
• pH 3–11
• UV-C (10.5 mW/cm\(^2\))
• 60 min light (ads+deg)
• Single-solute & deionized water | Composites have higher adsorption and photoactivity than TiO\(_2\) or char
• Composite with 100 g biomass calcined at 700 °C has attains 98% CBZ removal possibly due to higher surface area and smaller crystallite size | [147] |

| CBZ | TiO\(_2\)/biochar | • Similar to ref. [147]
• [Catalyst] = 60–140 g/dm\(^3\)
• [CBZ] = 10 - 50 mg/dm\(^3\)
• 6–24 g/dm\(^3\) \( O_2 \) flow
• pH 3–11
• UV-A (4.2 mW/cm\(^2\)) and UV-C (10.5 mW/cm\(^2\))
• 60 min light (ads+deg)
• Single-solute & deionized water | Almost constant (0.05 min\(^{-1}\)) CBZ degradation rate for pH 3 to 11, attaining 89.2–94.4% CBZ removal
• \( O_2 \) flow increase improves the CBZ photodegradation rate constant more than 1.6 times (0.025 to 0.042 min\(^{-1}\))
• After 1 h of UV-C irradiation GAC and composite remove, respectively, >90% and >99% CBZ while TiO\(_2\) and coconut shell powder only remove 35–42% of CBZ
• Enhanced adsorption+degradation with the composite attributed to the surface area that allows adsorbing CBZ that is further photodegraded by the reactive oxygen species produced by the semiconductor
• Composite has superior recycling performance over 11 reuse cycles | [148] |

Khrasheh et al. assayed TiO\(_2\)/biochar, with coconut shell powder char, for the CBZ photodegradation under UV light irradiation in the presence of \( O_2 \) flow [147,148]. Regardless of the biochar amount, all composites presented higher adsorption and photocatalytic activity than bare TiO\(_2\) biochar and granular activated carbon (GAC). For pH 3 to 11 the degradation rate of CBZ is almost constant (0.05 min\(^{-1}\)) attaining 89.2–94.4% CBZ removal [148]. Interestingly after 1h of UV-C irradiation GAC and composite attain, respectively, >90% and >99% CBZ removal while bare TiO\(_2\) and biochar only attain 42% of CBZ removal [148]. While the 90% removal with GAC is mainly attributed to adsorption (nanoporous solid), in the case of the biochar, that has an incipient pore structure, the 42% CBZ removal may result from the combined adsorption and photocatalytic degradation. Regarding recyclability TiO\(_2\)/biochar also has better performance since after 11 reuse cycles it removes 60% CBZ while for the other solids the removal is <25% [148]. The enhanced removal of CBZ with the TiO\(_2\)/biochar composite is attributed to the surface area that allows adsorbing CBZ that is further photodegraded by the reactive oxygen species produced by the semiconductor [148].

#### 3.7. Overview of the Degradation Pathways and Intermediates/Products

One of the major challenges of photocatalytic degradation of PhCs is to achieve total mineralization. Although in optimized conditions high and fast degradation of the target pollutants can be attained, the mineralization rate is usually slower than the removal rate leading to the presence of degradation intermediates and/or products whose toxicity can be higher than that of the parent compound.

Among all the works focused on SMX photocatalytic degradation by semiconductor/carbon materials only two address the mechanism and propose decomposition pathways [115,145], in both cases under solar light irradiation. Despite the use of distinct composites—WO\(_3\)/MWCNT and Zn-TiO\(_2\)/biochar—four common transformation intermediates/products were identified and similar degradation pathways are proposed (Figure 6): (i) hydroxylation (OH\(^{\circ}\) attack) on the benzene ring or isoxazole ring of SMX generating mono- or di-hydroxyl derivatives, and (ii) then OH\(^{\circ}\) cleavages the S-N bond. [115,145]. Zhu et al. propose that continuous radical attack to the benzene and isoxazole rings leads to bond cleavage [115]. The degradation products identified present m/z down to 99, being mainly substituted aromatic compounds. However, none of the studies report the mineralization degree or evaluated the toxicity of the solution after photocatalysis.
while DCF is “mutagenic positive” only one of the degradation intermediates (obtained by chlorine cleavage are the transformation pathways generally proposed (Figure 7). The smallest degradation products identified are single–ring aromatic compounds with chlorine, hydroxyl, carboxyl, and amine substituents (m/z from 127 to 194). By using g-C3N4/CQD Liu et al. attained DCF mineralization degrees of 54% after 1.5 h of visible light irradiation and estimated the toxicity of DCF and its degradation intermediates/products by quantitative structure-activity relationship (QSAR) prediction [140]. The authors conclude that, according to acute toxicity values, although in general the degradation intermediates are less toxic than parent DFC they are still classified as very toxic (LD50 for rat is 244 mg/kg). However the photocatalytic process with g-C3N4/CQD significantly decreased the bioaccumulation factor and development toxicity, compared to parent DCF. Moreover while DCF is “mutagenic positive” only one of the degradation intermediates (obtained by chlorine removal and ring closure) has the same classification, and the remaining are “mutagenic negative” [140]. Czech et al. verified that after photodegradation treatment of DCF solution with SiO2-TiO2/MWCNT, and regardless the use of UV-A or visible light, the V. fisheri bioluminescence inhibition decreased to values around 20% (i.e., in the border between toxic and non-toxic classification) while the toxicity of the initial DCF solution was higher than 90% [119]. The photodegradation of DCF with BiOCCOH/CQD composites under visible light irradiation demonstrates that treatment time is a critical factor [138]. After 40–50 min of treatment DCF was totally removed, but the mineralization degree was only 47.9%, certainly due to the abundant degradation intermediates. The assays with V. fisheri, D. magna, and D. subspicatus systematically reveal that the toxicity of the degradation intermediates presents a maximum at 40–50 min of treatment (more than double of the parent DCF) [138]. However, after 2 h of treatment DCF was effectively mineralized, detoxified and dechlorinated, clearly demonstrating

![Figure 6. SMX degradation pathways and intermediates by WO3/CNT and/or Zn-TiO2/biochar under simulated solar light irradiation (based on [115,145]).](image-url)
the need of prolonged treatment times to assure higher mineralization degrees and consequently lower toxicity.

![Figure 7. DCF degradation pathways and intermediates by Ag-BiOI/rGO, TiO_2/rGO with peroxodisulfate, TiO_2[001]/N-CQD, g-C_3N_4/CQDs, and/or BiOCOOH/CQDs under visible light irradiation. The OH substituents which can appear in different positions in the rings cross the parenthesis mark (based on [132,133,138–140]).]

Regarding CBZ three of the four studies reporting degradation pathways and products used TiO_2 derived composites with MWCNT, rGO, and biochar [116,124,146]. Figure 8 resumes the CBZ degradation pathways and intermediates by TiO_2/rGO, TiO_2/MWCTox and/or Fe_3O_4/BiOBr/biochar under distinct irradiation conditions (UV, near UV-vis, and Visible LED light). All the proposed degradation pathways start with the hydroxylation of the aromatic system or the olefinic double bond, and two of the studies reported that the hydroxylated intermediates suffer ring cleavage to originate substituted single-ring aromatic compounds (m/z from 93 to 137) or even an aliphatic dicarboxylic acid (m/z of 115). The study reporting the use of Fe_2O_3/BiOBr/biochar for the degradation of CBZ under visible light attained 70% mineralization indicating that most of the degradation intermediates could be eliminated to small organic molecules [146]. The toxicity of model wastewater containing CBZ before and after photodegradation treatment with TiO_2/SiO_2/MWCNTox under UV light was evaluated by Czech et al. that reported lower toxicity of the treated solution to *V. fisheri* and *D. magna* when compared with the non-treated wastewater [117].
Figure 8. CBZ degradation pathways and intermediates by TiO$_2$/rGO, TiO$_2$/MWCNTox and/or Fe$_3$O$_4$/BiOBr/biochar under distinct irradiation conditions (UV, near UV-vis, and Visible LED light). Short-lived intermediates presented between [brackets] while those between (parenthesis) correspond to stable intermediates, the OH substituents which can appear in different positions in the rings cross the parenthesis mark (based on [116,124,146]).

Overall, regardless of the target PhC—SMX, DCF, and CBZ—and the composite used for photocatalytic degradation, the main products are mainly substituted single-ring aromatic compounds. The degradations pathways of SMZ and CBZ start with the hydroxylation of the aromatic moieties. DCF has the most complex degradation pathway since it can start by hydroxylation, decarboxylation and/or dechlorination. Besides OH\(^\bullet\) attack to the aromatic systems, O$_2$\(^\bullet\) plays a prominent role in the decomposition of the DCF carboxylic acid group. Despite the high number of degradation pathways, DCF is the PhCs with a larger amount of intermediates and products identified.

4. Summary and Outlook

The versatility of carbon element is well expressed in this review, the combination of C atoms with semiconductors (C-doping) or the use of carbon materials—activated carbons, CNT, graphene, and derived materials, CQD and biochars—as supports or doping materials in composites with semiconductors allows to enhance photocatalytic activity and sunlight harvesting. The black body character of the carbon materials associated with properties as distinct as texture, conductivity, or quantum effects are certain parameters responsible for the improved performance of the composites.

Regarding the three target pharmaceuticals of this review, DCF was by the most explored in photocatalytic degradation studies with semiconductor/carbon material (19 papers vs. 8 papers for SMX and 14 papers for CBZ), most likely due to its presence between 2013 and 2018 in the Watch.
Lists published by European Commission. The interest of the academy towards CBZ probably results from its high recalcitrant behavior in conventional wastewater treatment and high persistence in the environment that also supports the proposal of this pharmaceutical as a possible anthropogenic marker. In fact, also the few studies that evaluate the photocatalytic degradation of more than one of the three targets PhCs systematically identify CBZ as the PhC most resistant to photocatalytic degradation followed by DCF and SMX, in line with photolysis data and persistence after conventional wastewater treatment (biodegradation). Therefore, it is important to continue developing enhanced technologies to effectively restrain the dissemination of these recalcitrant PhCs, with special focus on CBZ, in the environment.

Regarding photocatalytic degradation of these PhCs, only few studies are focused on real water matrices, with the catalytic performance being mainly evaluated using deionized water solution at pH not close to the values of real water. However, the studies reporting data obtained in real water matrices reveal that the high performance of the catalyst in deionized water is not a guaranty for good results in spiked wastewater.

The majority of studies reviewed are focused on evaluating the removal efficiency of novel catalysts and composites for a specific contaminant, in most cases not assessing the mineralization rate. However the data available clearly demonstrates that the mineralization process demands longer operation times than the degradation/removal of the target PhC. Moreover, when toxicological assays are reported the data reveal that the transformation intermediates formed during the initial degradation process may be as toxic as, or even more toxic than, the pristine PhC.

As a final remark it must be stressed that regardless of the type of carbon material present in the composite, the literature data is almost unanimous to demonstrate the beneficial effect of the carbon doping or addition for the sunlight harvesting and better overall performance of the semiconductor/carbon catalysts. The enhanced photocatalytic activity of the semiconductor/carbon composites is mainly attributed to the electron transfer properties of the carbon material and better charge separation. However, there are also reports where the carbon material (CQDs) act as photosensitizers. The bibliometric analysis shows the great interest of the scientific community in the more conducting carbon materials—CNT and graphene-derived materials—and also the recent growing research in CQD. Nevertheless, more traditional carbon forms, as is the case of nanoporous carbons or biochars, may be more sustainable and feasible for large scale applications not only due to the production cost but mainly due to the toxicity of the crystalline carbon forms. In fact, in vivo studies reveal the health issues associated with CNT and graphene-derived materials due to their rigid structures that damage the cell walls [149,150]. Another advantage of nanoporous carbons is related to their well-known pore structure that, besides contributing to the overall removal of the target pollutant, may also allow the adsorption of the degradation intermediates and products thus contributing to overall decrease of the toxicity level.

Lastly, it must also be mentioned that self-photoactivity of some carbon materials for these recalcitrant PhCs, in both UV and solar irradiation, was observed in more than one work, point out the need to further explore the potential of metal-free carbon materials for the photocatalytic degradation of priority contaminants, ideally under sunlight irradiation and in real water matrices.

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