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## Metal oxides-based photocatalyst for the efficient degradation of organic pollutants for a sustainable environment: a review

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Photocatalytic degradation is a highly efficient technique for eliminating organic pollutants such as antibiotics, organic dyes, toluene, nitrobenzene, cyclohexene, and refinery oil from the environment. The effects of operating conditions, concentrations of contaminants and catalysts, and their impact on the rate of deterioration are the key focuses of this review. This method utilizes light-activated semiconductor catalysts to generate reactive oxygen species that break down contaminants. Modified photocatalysts, such as metal oxides, doped metal oxides, and composite materials, enhance the effectiveness of photocatalytic degradation by improving light absorption and charge separation. Furthermore, operational conditions such as pH, temperature, and light intensity also played a crucial role in enhancing the degradation process. The results indicated that both high pollutant and catalyst concentrations improve the degradation rate up to a threshold, beyond which no significant benefits are observed. Optimal operational conditions were found to significantly enhance photocatalytic efficiency, with a marked increase in degradation rates under ideal settings. Antibiotics and organic dyes generally follow intricate degradation pathways, resulting in the breakdown of these substances into smaller, less detrimental compounds. On the other hand, hydrocarbons such as toluene and cyclohexene, along with nitrobenzene, may necessitate many stages to achieve complete mineralization. Several factors that affect the efficiency of degradation are the characteristics of the photocatalyst, pollutant concentration, light intensity, and the existence of co-catalysts. This approach offers a sustainable alternative for minimizing organic pollutants present in the environment, contributing to cleaner air and water. Photocatalytic degradation hence holds tremendous potential for remediation of the environment.

#### Introduction

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Urbanization and industrialization are cornerstones of modern civilization, underpinning significant advances in economic growth, technological innovation, and improved standards of living <sup>1</sup>. These processes have facilitated the development of cities, expanded infrastructure, and increased industrial productivity, creating myriad opportunities for societal progress 2,3 However, the rapid pace of urbanization and industrialization has also ushered in substantial environmental challenges, particularly through the generation of wastewater that contains a diverse array of organic pollutants <sup>4-6</sup>. These pollutants are frequently hazardous and contrary to conventional treatment procedures, presenting significant hazards to the both environment and public health 7-9. Industrial operations are major contributors to wastewater pollution, as they produce effluents laden with complex organic chemicals <sup>10-12</sup>. These chemicals are often by-products of various industrial processes and include a wide variety of substances such as antibiotics, organic dyes, nitrobenzene, cyclohexane, phenols, toluene, biphenyls, pesticides, fertilizers, hydrocarbons, plasticizing agents, detergents, oils, greases, proteins, and carbohydrates 13-15. The environmental impact of these pollutants is profound, as they can persist in the environment, bioaccumulate in wildlife, and enter human food chains, leading to chronic health issues and ecological damage <sup>16,17</sup>. The complexity and resilience of these organic pollutants necessitate the development of advanced treatment technologies <sup>18,19</sup>. Traditional biological treatment methods are often inadequate for fully degrading these pollutants due to their toxicity and chemical stability. In response to this challenge, Advanced Oxidation Processes (AOPs) have been developed and are increasingly being employed for the effective degradation of hazardous organic contaminants present in wastewater <sup>20–22</sup>. AOPs are distinguished by the production of extremely reactive species, such as hydroxyl radicals, that can indiscriminately oxidize a broad spectrum of organic pollutants. This process converts the pollutants into less dangerous chemicals or fully mineralizes them into carbon dioxide (CO<sub>2</sub>) and water (H<sub>2</sub>O) <sup>23</sup>. Among the various AOPs, photocatalytic degradation stands out as a particularly effective method <sup>20</sup>. Photocatalysis involves the use of semiconductor materials as catalysts to accelerate chemical reactions upon exposure to light. When semiconductor materials such as zinc oxide (ZnO), iron oxide (Fe<sub>2</sub>O<sub>3</sub>), titanium dioxide (TiO<sub>2</sub>), gallium

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phosphide (GaP), cadmium sulfide (CdS), and zinc sulfide (ZnS) are exposed to light, they generate electron-hole pairs that can generate reactive oxygen species <sup>24-26</sup>. These reactive species contain the very capability of breaking down complex organic pollutants into less harmful, simpler molecules and fully mineralizing them <sup>27,28</sup>. The advantages of photocatalysis are numerous and include low operational costs, the ability to accomplish full mineralization of contaminants without generating secondary pollution, and the capability to operate at ambient temperatures and pressures <sup>29</sup>. Among the various photocatalysts, titanium dioxide (TiO<sub>2</sub>) is the most extensively studied and broadly applied because of its exceptional chemical and photochemical stability, cost-effectiveness, low toxicity, and high activity under ultraviolet (UV) light. TiO<sub>2</sub>, with its wide band gap of approximately 3.2 eV, can mineralize a broad spectrum of organic contaminants, including herbicides, dyes, pesticides, phenolic compounds, and pharmaceuticals like tetracycline and sulfamethazine <sup>30,31</sup>. Nevertheless, the actual utilization of TiO<sub>2</sub> is somewhat restricted due to its dependence on UV light, which comprises just a minor portion of the solar spectral region <sup>32</sup>. To overcome this limitation, other semiconductor materials with broader light absorption properties are being explored. Tungsten trioxide (WO<sub>3</sub>) has emerged as a promising alternative due to its capability of absorbing visible light, making it more competent for photocatalytic oxidation of volatile organic pollutants under natural sunlight <sup>33,34</sup>. Additionally, silver nanoparticles (AgNPs) have gained significant attention as photocatalysts due to their high photostability, environmental friendliness, and catalytic properties that are dependent on their shape and size <sup>35</sup>. The effectiveness of photocatalytic systems in degrading organic pollutants is dependent on numerous operational parameters. These factors encompass the substrate concentration, photocatalyst quantity, pH of the solution, reaction medium temperature, light irradiation duration and intensity, photocatalyst surface area, dissolved oxygen content in the reaction medium, and the characteristics of both the photocatalyst and substrate <sup>29,36,37</sup>. Furthermore, the doping of photocatalysts with metal and non-metal ions can enhance their photocatalytic activity by modifying their electronic properties and extending their light absorption range <sup>38</sup>. It is important to optimize these parameters to maximize the degradation kinetics and overall efficiency of photocatalytic processes <sup>39</sup>. For instance, the proportion of the substrate to the photocatalyst must be carefully balanced to ensure that there are enough reactive sites for pollutant molecules to adsorb and react <sup>37</sup>. The pH of the solution can affect the charge and surface properties of the photocatalyst, influencing its interaction with pollutants. Temperature and light intensity also play significant roles in determining the rate of photocatalytic reactions, with higher temperatures and light intensities generally leading to increased reaction rates <sup>40–42</sup>. In this review, we focused on the degradation of six specific types of organic pollutants: antibiotics, organic dyes, nitrobenzene, toluene, oil, and cyclohexane. These pollutants represent a broad spectrum of chemical structures and environmental impacts, making them ideal candidates for studying the effectiveness of various photocatalysts under different operational conditions. We will delve into the various reaction parameters that are contrained achieving maximum degradation of these pollutants using different photocatalysts. This comprehensive analysis aims to provide insights into the optimal conditions and catalyst selections for effective wastewater treatment, contributing to the mitigation of environmental pollution and the protection of aquatic ecosystems.

# Photocatalytic degradation of chemical pollutants (organic dyes and antibiotics)

Chemical pollutants refer to a large group of contaminants that arise from different sources, including pharmaceuticals 43, personal care items <sup>44</sup>, pesticides <sup>45</sup>, and other synthetic chemicals <sup>46</sup>. Chemical pollutants, such as antibiotics and organic dyes, have significant adverse effects on the environment 47. Antibiotics, encompassing classes such as betalactams (e.g., penicillins, cephalosporins), macrolides (e.g., tetracyclines erythromycin), (e.g., doxycycline), aminoglycosides (e.g., gentamicin), quinolones (e.g., sulfonamides sulfamethoxazole), ciprofloxacin), (e.g., glycopeptides (e.g., vancomycin), and oxazolidinones (e.g., linezolid), are significant pharmaceutical pollutants <sup>48</sup>. Organic dyes, including azo dyes (e.g., methyl orange), anthraquinone dyes (e.g., alizarin), phthalocyanine dyes (e.g., copper phthalocyanine), triphenylmethane dyes (e.g., malachite green), xanthene dyes (e.g., fluorescein), and indigoid dyes (e.g., indigo carmine), are prevalent industrial pollutants <sup>49</sup>. Both types of pollutants are persistent in water bodies, posing substantial dangers to aquatic ecosystems and human health due to their toxicity, bioaccumulation potential, and the propagation of antibiotic-resistant bacteria <sup>50,51</sup>. The persistence and toxicity of these chemical pollutants effective remediation strategies, necessitate such as photocatalytic degradation, which utilizes light-activated catalysts to break down these harmful substances into harmless by-products, ensuring cleaner water and healthier ecosystems 52

#### **Organic Dyes**

A significant group of synthetic organic molecules produced by a variety of industries, including the leather, plastic, food, paper, textile, and medicinal sectors are known as dyes <sup>35</sup>,<sup>53</sup>. Due to their frequent application in various manufacturing sectors, dyes are inevitably accidentally released into the surroundings, particularly into either surface water or groundwater, where they may pose serious dangers to environmental and biological systems <sup>54</sup>,<sup>55</sup>,<sup>56,57</sup>. Over 700,000 tons of dyes are generated globally each year; 20% of these lost dyes reach the atmosphere and create pollution throughout processing or manufacturing, accounting for about 12% of the global total of dye generation. So the degradation is necessary of these organic dyes for maintaining the ecological balance <sup>58</sup>. Organic dyes are very detrimental to aquatic ecosystems, even This article is licensed under a Creative Commons Attribution 3.0 Unported Licence.

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at low concentrations (less than 1 ppm). Thus, it is essential and required to remove organic dyes from effluent <sup>59</sup>. The degradation mechanism of methylene blue dye is as follows <sup>60</sup>. Photocatalyst + hv(photon)  $\rightarrow$  Photocatalyst ( $e_{cb}$ <sup>-</sup>+ $h_{vb}$ <sup>+</sup>) (1)

$e_{cb}^{-} + O_2 \rightarrow \cdot O_2^{-}$	(2)	
$h_{vb}^{+} + H_2O \rightarrow \cdot OH + H^+$	(3)	
$h_{vb}^+ + OH^- \rightarrow \cdot OH$	(4)	
Methylene blue $+ \cdot OH \rightarrow Degra$	ded products	(5)
Methylene blue + $\cdot O_2^- \rightarrow Degr$	aded products (6)	
Degraded products +·OH/O <sub>2</sub> $\rightarrow$	CO <sub>2</sub> + H <sub>2</sub> O + Inorganic	ions (7)

Several metal oxides, such as ZnO, MgO, AgO, TiO<sub>2</sub>, Fe<sub>2</sub>O<sub>3</sub>, Mn<sub>2</sub>O<sub>3</sub>, CuO, and V<sub>2</sub>O<sub>5</sub> are frequently employed as photocatalysts in wastewater treatment processes to degrade dyes <sup>61</sup>. Zinc oxide (ZnO) is an oxidizing substance found in nature as the unusual mineral zincite. There have been attempts to use ZnO alongside other semiconductors for the photocatalytic destruction of an extensive variety of biological pollutants <sup>62</sup>. ZnO-based photocatalysts work according to the various parameter conditions. These parameters are mainly Ph, the initial concentration of dye or catalyst, the wavelength of the light & so on. The amount of photocatalytic reaction rate at the outermost layer of the catalyst can be influenced by the initial concentration of the substrate. To prevent the dispersion of light and the concentration impact of the exposed photocatalyst surface, the ideal photocatalyst concentration ought to be unique for heterogeneous photocatalysis processes <sup>63</sup>. Velmurugan et al. stated that the rate of degradation k dropped from 0.173 to 0.012 min<sup>-1</sup> when the dye concentration was increased from  $1 \times 10^{-4}$  to  $4 \times 10^{-4}$  M <sup>64</sup>. This is because many layers of adsorbed dye molecules have formed on the outermost layer of the catalyst, which prevents the photoreaction from occurring because there was not enough direct light interaction to produce hydroxyl radicals <sup>65</sup>. The first amount of dye has a significant influence on the degrading efficiency of MB 66. Sobana et al. used ZnO that was manually combined with activated carbon (AC-ZnO) and solar irradiation to study the impact of initial Direct Blue 53 (DB53) concentration over the concentration that ranged from 1×10<sup>-4</sup> to 9×10<sup>-4</sup> M <sup>67</sup>. Its numerous responsibilities make it extremely difficult to determine how the pH of a solution affects the efficacy of the dye photocatalytic degradation activity <sup>68</sup>. Velmurugan et al. stated the impact of pH in the range of 3-11 upon the photocatalytic breakdown of Reactive Red 120 (RR 120) over ZnO during solar light irradiation <sup>64</sup>. Photocatalytic breakdown of reactive orange 4 (RO4) and black 5 (RB5) dyes at various solution pH levels between 3 and 11<sup>69</sup>. The pH, which regulates the adsorption of organic compounds on the outermost layer of the photocatalyst, serves as one of the most crucial factors influencing photocatalysis effectiveness 70. Electromagnetic relationships between the outermost layer of the photocatalyst and the substrate of interest can be employed to clarify how pH affects photocatalysis outcomes <sup>27</sup>. Singh et al. stated that after exposing ZnO nanorods to UV radiation for 120 minutes, photodegradation activity levels of 7.169% and 47.63% for pH values of 4.5 and 10.5, correspondingly <sup>71</sup>.

Scientists' interest has been drawn to supported TiOarcatalyst utilization more and more over the past few years due of the prospective uses in the photocatalytic breakdown of organic contaminants such as organic dyes in air and water. Additionally, reports have it that when adsorbents are used to support TiO<sub>2</sub>, an ideal condition is created for the elimination or degradation of the compounds of interest <sup>72,73</sup>. To enhance TiO<sub>2</sub>-based photocatalysts on organic dye in wastewater, several conditions were adjusted. These crucial elements, which included light intensity, TiO<sub>2</sub> form, and structure, target type, pH level as well as doping type, all had an impact on the photocatalysis method's effectiveness <sup>58</sup>. If we want to discuss the parameters it is found that it is rather tough to comprehend how pH impacts the photodegradation process's efficacy <sup>29</sup>. TiO<sub>2</sub> exhibits amphoteric properties behavior, allowing for the development of either a positive or negative charge on its outermost layer <sup>74</sup>. Due to this, the adsorption of dye molecules over TiO<sub>2</sub> surfaces may be affected by changes in pH <sup>75</sup>. Bubacz et al. found that when pH is increased, so did the rate at which methylene blue was broken down photo-catalytically <sup>76</sup>. On the other hand, Neppolian et al. studied that acidic condition does not affect the degradation rate of the Reactive Blue 4 significantly enough 77. It has been found that organic dyes like Reactive Black 5 and Reactive Orange 4 degradation were enhanced in an acidic solution containing TiO<sub>2</sub><sup>69</sup>. Tanaka et al. discovered that at less acidic values, the positively charged TiO<sub>2</sub> layer absorbed more Acid Orange 7, and greater breakdown was accomplished <sup>78</sup>. A study has been conducted on the effects of pH on the adsorption as well as decolorization of Procion Red MX-5B (MX-5B) and Cationic Blue X-GRL (CBX). It was discovered that when the pH increased, MX-5B's adsorption was reduced <sup>62</sup>. Another key parameter for dye degradation using a TiO<sub>2</sub> catalyst is the dye amount or dye concentration. It has been found that the increased initial concentration of the dyes increase the degradation rate <sup>79</sup>, <sup>36</sup>. This is because when the dye's initial concentrations rise, the dye molecules become deposited on the outermost layer of the catalyst and consume a sizable proportion of UV light instead of the TiO<sub>2</sub> nanoparticles <sup>80,81</sup>. Neppolian et al. investigated how the original dye concentration affected the percentage of degradation. With the best possible catalyst loading, they changed the starting concentrations of Reactive Yellow 17 (8.9×10<sup>-4</sup> to 1.29×10<sup>-3</sup> M), Reactive Red 2 (4.169×10<sup>-4</sup> to 1.259×10<sup>-3</sup> M), and Reactive Blue 4 ( $1.9 \times 10^{-4}$  to  $5.9 \times 10^{-4}$  M) <sup>77</sup>. The dye degradation in a waterbased solution utilizing catalyst powder of  $TiO_2$  within a photocatalytic reactor is influenced by two additional parameters: the wavelength and intensity of the UV light irradiation source 82. Lower radiation wavelengths are thought to encourage the creation of electron holes, which would increase the catalyst's effectiveness 83. Ollis et al. said that at minimal light levels (0-20 mW/cm<sup>2</sup>), the rate would rise in an orderly manner as the intensity of light increased. The rate would rely on the square root of the light intensity at moderate light intensities (about 25 mW/cm<sup>2</sup>) but at intense light levels, the rate is independent of the light intensity <sup>84</sup> <sup>29</sup>. The degradation of Orange G was shown to be affected by light intensity in a range of 215 to 586 W/cm<sup>2</sup>. With a rise in light

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magnitude, Orange G's photolysis reaction rates climbed <sup>85</sup>. Rao et al. stated that acid orange 7 (AO7) photocatalytically breaks down at a pace that is roughly 1.5 times faster in direct sunlight compared to it is under synthetic UV radiation <sup>86</sup>. Another significant operational parameter for the organic dye degradation is temperature range <sup>36</sup>. The range of 40–50 C was determined to be the ideal operating temperature range. Since desorption of the produced products happens more slowly at

low temperatures than interface degradation<sub>Vi</sub>as<sub>Ar</sub>well<sub>ni</sub>as reactant adsorption, it restricts the reaction conversely the limiting step becomes the dye's adsorption on TiO<sub>2</sub> at an elevated temperature <sup>87</sup>. The rate constant is lowered at elevated temperatures due to the organics' and hydrated oxygen's reduced adsorptive ability. Consequently, the ideal temperature often falls between 293 and 353 K <sup>88 89</sup>.

**Table 1:** Data for the photocatalytic degradation of organic dyes using various catalysts.

Pollutants	Pollutant Conc. (mg/L)	Catalyst	Catalyst Conc. (mg/L)	Operational Conditions	Lamp, Power (W)	Degradation Percentage (%), Time (min)	Ref.
MB	63.97	Mn-doped ZnO		Visible Light; Light intensity: 18.6 lux	Tungstane lamp, 500	50%, 10	90
IC	10	CA–CNT/TiO <sub>2</sub> –NH <sub>2</sub>		UV Light (315- 400 nm), pH =2, temp = 80 °C	40	100%, 180	91
MB	30	CA–CNT/TiO <sub>2</sub> –NH <sub>2</sub>		UV Light (315- 40 nm), pH =2, temp = 80 °C	40	80%, 300	91
RhB	6	nanostructured TiO <sub>2</sub>	0.0001- 0.0005	UV Light, pH = around 7		93.8%, 190	92
AR57	30	TiO <sub>2</sub>	0.0005	UV Light, pH = 7.18, Temp = 400 °C		90.7%, 190	92
CR	75	ZnO	0.00016	Solar Light, pH = 6		97%, 120	93
MB	50	Cu-doped ZnO (NPs)		Visible Light	300	85%, 60	94
RhB	10	WO <sub>3</sub> /Ag <sub>2</sub> CO <sub>3</sub>		Visible Light	Metal halide lamp, 70	99.7%, 8	95
MB	64	Undoped ZnO		Visible Light; Light intensity: 18.6 lux	Tungsten lamp, 500	50%, 30	67
MO	10	2%Al- 2%Ni-ZnO	500	Visible Light	Halogen lamp, 100	99%, 30	96
MB		Nano ZnO		UV light	Mercury vapor lamp, 8	97.64%, 120	97
MO	25	ZnO/Cu <sub>2</sub> O		UV Light	Tungsten lamp, 200	73%, 180	98
AR27		Ce-ZnO	0.004	Solar Light, Degradation steadily increased till pH=12		90%, 60	79
MO	15	ZnO NFs		UV Light		99.46%, 50	99
MO		Natural zeolites Supported TiO <sub>2</sub>	0.0006 - 0.004	UV light, pH = 4		96.58 %, 100	100

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AB25	100	MgAl <sub>2</sub> O <sub>4</sub> nanoparticles		UV , pH = 3		99.86%, 35	<b>101</b> View Article C
MB	50	ZnO NPs		UV/Visible,		100% <sup>D</sup> under <sup>10</sup>	39 <b>.02</b> 4NA00
						UV irridation in	
						20 min, 91%	
						under visible	
						light in 60 min	
RhB	10	$TiO_2/g-C_3N_4$		Solar Light,	Compact	95%, 120	103
				Temp: 350 <sup>0</sup> C	xenon		
					lamp, 300		
MO	10	[Zn(L)(H₂O)]·H₂O		UV Light,	High	83,8%, 120	104
					pressure Hg		
					lamp, 300		
MB	5	ZnO nanowires		UV Light,	High	96%, 120	105
					pressure Hg		
					lamp, 50		
IC	10	TiO <sub>2</sub> –NH <sub>2</sub> NPs		UV light, pH = 2	UV lamp,	100%, 180	91
				Temp: 80 <sup>0</sup> C	40		
RhB	0.001	The floral-like LaFeO <sub>3</sub>		Visible Light,	High	, 720	106
					pressure Hg		
					lamp, 150		
RhB	23.4	ZnO	400	pH = 7		, 180	107
RO16	20-60	TiO <sub>2</sub>	90	UV light,	Xenon	87% after 20	108
				pH=7.0, Temp:	lamp,	min, 70% after	
				25 °C		20 min, nearly	
						100% after	
						120 min	
MO		PbBiO <sub>2</sub> Br		Visible light	Xe arc	95%, 60	109
					lamp, 300		

#### \*MB = Methylene Blue, \*IC = Indigo Carmine, \*RhB = RhodamineB, \*AR57 = Acid Red 57, \*CR = Congo Red, \*MO = Methyl Orange, \*AR27 = Acid Red 27, \*AB25 = Acid Blue 25, \*RO16 = Reactive Orange 16



Fig. 1: Working procedure of the photocatalyst for the dye degradation

### Antibiotics

Due to their extremely stable and non-biodegradable nature, antibiotics accumulate in the ecosystem as a result of overuse and uncontrolled environmental discharge <sup>110,111</sup>. The release of diverse antimicrobial pollutants and their varied toxicity provide

a significant challenge for researchers trying to find a solution <sup>112,113</sup>. The excessive accumulation of antibiotics in natural environments has presented a significant peril to ecological systems <sup>114,115</sup>. Unfortunately, water treatment traditional methods such as adsorption, filtration, and biodegradation are ineffective in effectively removing antibiotics due to their significant durability and limited biodegradability. Hence, the

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development of novel technologies is vital to ensure the efficient elimination of antibiotics <sup>116–119</sup>. Due to its advantageous characteristics cost-effectiveness, of environmental sustainability, and high efficacy, heterogeneous photocatalysis has become a process of great promise for wastewater treatment, which relies on the direct utilization of sunlight to effectively degrade and subsequently mineralize organic pollutants, has emerged as a promising approach to tackle diverse environmental challenges <sup>120–122</sup>. Furthermore, it is crucial to provide an overview of frequently utilized photocatalytic nanomaterials and their specific use in breaking down popular antibiotics. This is necessary to validate their practical superiority and efficacy as catalysts for the process of photodegradation <sup>123–125</sup>. The degradation mechanism of Ciprofloxacin antibiotic in the presence of different photocatalysts is provided <sup>126</sup>.

Photocatalyst + hv(photon)  $\rightarrow$  Photocatalyst ( $e_{cb}^{-}$  +  $h_{vb}^{+}$ ) (8)

 $e_{cb}^{-} + O_2 \rightarrow O_2^{-} \qquad (9)$  $h_{vb}^{+} + H_2O \rightarrow OH + H^+ (10)$ 

 $h_{vb}^{+}$  + OH<sup>-</sup> → ·OH (11)

Ciprofloxacin +  $\cdot$ OH  $\rightarrow$  Degraded products (12)

Ciprofloxacin +  $\cdot O_2^- \rightarrow$  Degraded products (13)

Degradation products +  $\cdot$ OH/O<sub>2</sub>  $\rightarrow$  CO<sub>2</sub> + H<sub>2</sub>O + Inorganic ions (14)

Yang et al. researched the degradation of Ciprofloxacin using g- $C_3N_4/TiO_2$  nanocomposites by the help of visible light irradiation utilizing a 300 W Xe visible lamp where the authors observed 88% CIP degraded in 180 minutes 127. Verma explored the degradation of amoxicillin (AMX) by the utilization of TiO<sub>2</sub> photocatalysis and sono-photocatalysis and achieved the highest degradation rate (80%) of AMX at pH of 7.0 under UV irradiation at a power density of 672 W/m<sup>2</sup> <sup>128</sup>. Zhang examined the mechanism and kinetics of photocatalytic degradation of tetracycline (TC) utilizing a supramolecular organic photocatalyst called three-dimensional network structure perylene diimide (3D-PDI) <sup>129</sup>. Fan et al. synthesized three different structures of Bi-modified titanate nanomaterials (Bi-TNM) utilizing the hydrothermal technique and carefully adjusted variables to break down paracetamol (ACT). The study revealed that Bi-Titanate nanoribbons, when used at a concentration of 1 gL<sup>-1</sup>, had the most effective photocatalytic degradation capability, achieving a rate of 88% <sup>130</sup>. The catalytic efficiency of NiS and NiS immobilized within the magnetite polypyrrole core/shell matrix (Fe<sub>3</sub>O<sub>4</sub>@PPY) was examined for the degradation of cephalexin. The study also examined the photocatalytic breakdown of cefalexin using the NiS-PPY-Fe<sub>3</sub>O<sub>4</sub> photocatalyst, which was exposed to sunshine. The photocatalyst demonstrated a removal efficiency of over 80% over a 30-minute timeframe <sup>131</sup>. Payan studied the creation of photocatalysts using Cu-TiO2@functionalized single-walled carbon nanotubes and found that sulfamethazine can be fully destroyed under solar irradiation within 300 minutes <sup>132</sup>. Rajeev et al. synthesized BN/CdAl<sub>2</sub>O<sub>4</sub> composites and evaluated their photocatalytic ability to degrade cefoxitin sodium (CFT) antibiotic in an aqueous solution. The findings demonstrated that a nearly complete degradation of CFT, reaching approximately 100%, was seen within 240 minutesie at line (BiOBr) photocatalyst using the PVP-assisted was produced by solvothermal technique. The PVP-capped BiOBr exhibits a removal efficiency of 94% and 99.8% for the antibiotics ofloxacin (OFL) and norfloxacin (NOR) respectively, when exposed to visible light <sup>134</sup>. Yinyan prepared Z-scheme CdTe/TiO2 heterostructure photocatalysts decomposing 78% tetracycline hydrochloride (TC-H) within 30 min of irradiation under visible light <sup>135</sup>. Wei examined the photocatalytic efficiency of four tetracycline antibiotics using BiVO<sub>4</sub>/TiO<sub>2</sub>/RGO composites. The BiVO<sub>4</sub>/TiO<sub>2</sub>/RGO photocatalyst demonstrated significant photocatalytic activity and compatibility, providing efficient separation of photo-generated carriers with oxidation capabilities and high reduction <sup>136</sup>. Najmeh synthesized a novel heterojunction Z-scheme MnWO<sub>4</sub>/Bi<sub>2</sub>S<sub>3</sub> using a hydrothermal technique to study the photocatalytic behavior of catalysts in the decomposition of metronidazole (MTZ) and cephalexin (CFX) under LED light exposure where CFX achieved a maximum degradation efficiency of 78.8%, whereas MTZ earned a maximum degradation efficiency of 83.3% <sup>137</sup>. Akbar et al. studied manufactured nanostructured photocatalysts composed of tin oxide (SnO<sub>2</sub>) and cerium oxide (CeO<sub>2</sub>). These photocatalysts were employed to degrade the antibiotic tetracycline hydrochloride (TC) under visible light. The most optimal outcome seen among the examined photocatalysts had a TC removal effectiveness of approximately 97% within a 120minute timeframe under visible-light exposure <sup>138</sup>. An investigation was conducted on the photocatalytic degradation of pharmaceutical micropollutants of Penicillin G (PG) in a photoreactor. The proficiency of the photocatalytic process was increased by the inclusion of persulfate sodium (PPS). The inclusion of PPS greatly enhanced the efficiency of the photolysis process, resulting in a considerable improvement of 72.72% compared to the traditional photocatalysis system, which achieved 56.71% efficiency <sup>139</sup>. Bouyarmane synthesized TiO<sub>2</sub>-hydroxyapatite nanocomposites precipitating a redissolved natural phosphate mineral in ammonia using the concurrent gelation of titanium alkoxide. These nanocomposites were then subjected to degradation of the drug testing in a solution under ultraviolet light. When utilizing 40TiHAp as a photocatalyst, ciprofloxacin and ofloxacin were destroyed through photodegradation in 15 minutes and 120 minutes, respectively <sup>140</sup>. A simple solvothermal technique was employed to synthesize a novel Cu<sub>3</sub>P-ZSO-CN p-n-n heterojunction photocatalyst for the degradation of the antibiotic tetracycline (TC) under exposure to visible light. The degradation efficiency for TC was found to be 98.45% 141. Muneeb et al. synthesized ACT-X nanocomposites using activated carbon and TiO2 to enhance the inherent

characteristics of TiO<sub>2</sub>, resulting in improved light absorption in

the visible area. The ACT-4 photocatalyst has demonstrated the

maximum level of photocatalytic degradation (99.6%) for the

ceftriaxone (CEF) antibiotic <sup>142</sup>. The very first 3D hierarchical

ZnO/Bi<sub>2</sub>MoO<sub>6</sub> heterojunctions were synthesized using an in-situ

solvothermal technique. These heterojunctions exhibited a

remarkable efficiency of 100% in the photodegradation of the

ofloxacin (OFL) antibiotic. This exceptional performance can be ascribed to their reduced electron-hole recombination rate and large surface area <sup>143</sup>. A novel heterojunction photocatalyst (MoO<sub>3</sub>/g-C<sub>3</sub>N<sub>4</sub>) was synthesized using a straightforward hydrothermal calcination technique. The catalytic efficiency of this photocatalyst was assessed by measuring its ability to degrade tetracycline. The findings demonstrated that the 0D-2D MoO<sub>3</sub>/g-C<sub>3</sub>N<sub>4</sub> Z-scheme heterojunction outperformed the original g-C\_3N\_4 and achieved an impressive 85.9% removal efficiency within 100 minutes when exposed to visible light <sup>144</sup>. Elvira fabricated highly efficient photocatalysts by using electrochemical deposition and thermal treatment. These called photocatalysts. nanostructured homoiunction Bi<sub>2</sub>MoO<sub>6</sub>@Bi<sub>2</sub>MoO<sub>6-x</sub>, were able to effectively degrade and mineralize solutions containing various antibiotics (such as tetracycline, ciprofloxacin, and levofloxacin). After 180 minutes of radiation exposure, the photocatalysts achieved exceptionally high mineralization values (>95%) and nearcomplete degradation <sup>145</sup>. Peyman et al. examined the photocatalytic efficacy of Zn-Co-layered double hydroxide (LDH) nanostructures containing charcoal (BC) in the breakdown of gemifloxacin (GMF), a representative pharmaceutical contaminant. The results indicate that 92.7% of GMF underwent degradation through photocatalysis in the presence of the Zn-Co-LDH catalyst. The effectiveness of BC-incorporated

#### Zn-Co-LDH as a photocatalyst was greatly influenced by the concentration of the solute and the amount of photocatalyst used <sup>146</sup>. Elegant Z-scheme Composite hollow microspheres (CHMs) were made by sequentially controlling in situ hydrolysis and polymerization of WO<sub>3</sub>/g-C<sub>3</sub>N<sub>4</sub>. WO<sub>3</sub>/g-C<sub>3</sub>N<sub>4</sub> CHMs are the most effective photocatalytic degradation of CFS, with an 82% degradation efficiency after 2 hours of visible-light irradiation <sup>147</sup>. Yadav conducted research on the properties of photocatalyst magnesium titanate (MgTiO<sub>3</sub>) in the presence of visible light, specifically focusing on its interaction with lomefloxacin. The study found that a concentration of 30 mg/L of catalyst was the most effective in breaking down 10 mg/L of lomefloxacin using 30-W LED irradiation for a duration of 150 minutes <sup>148</sup>. The interaction between various surface facets of a semiconductor with suitable ratios can lead to improved performance in the degradation of photocatalytic processes. Juan et al. studied a material composed of bismuth called Bi<sub>4</sub>Ti<sub>3</sub>O<sub>12</sub> and examined that it showed improved degradation activity of tetracycline hydrochloride (TC-HCl) when exposed to irradiation 149. Mohammad investigated the degradation of cefazolin through exposure to immobilized and suspended TiO<sub>2</sub> on a glass plate. The findings indicate that the breakdown percentage of TiO<sub>2</sub> suspension at favorable pH conditions (pH 5) is 96.47% after 60 minutes of irradiation <sup>150</sup>.

Table 2: Data for the photocatalytic degradation of antibiotics using various catalysts.

Antibiotics*	Antibiot	Catalysts	Catalys	Operational	Lamn.	Degradation	Ref.
	ics		ts	Condition	Power	Percentage	
	Conc.		Conc.		(W)	(%). Time	
	(mg/L)		(mg/L)			(min)	
CIP	10	G-C <sub>3</sub> N <sub>4</sub> /TiO <sub>2</sub>	375	Visible- light	Xe, 300	88, 180	127
AMX	30	TiO <sub>2</sub>	450	UV light, pH 7	672	80, 270	128
тс	50	3D-PDI	25	Visible-light, pH 5	-	80, 150	129
АРАР	0.7	BiTNMs	1000	Visible light, pH 7	500	88, 180	130
CFX	50	NiS-PPY-Fe <sub>3</sub> O <sub>4</sub>	4000	UV- light, pH 5.5	Hg, 75	80, 30	131
SMZ	30	Cu-TiO <sub>2</sub> @functionalized SWCNT	900	UV-Vis light, pH 7	-	100, 300	132
CFT	15	BN/CdAl <sub>2</sub> O <sub>4</sub>	330	UV light, pH 7	108	100, 240	133
OFL	5	PVP capped BiOBr	10	Visible light	15	94, 240	134
NOR	5	PVP capped BiOBr	10	Visible light, pH 7.54	15	99.8, 240	134
ТС-Н	20	CdTe/TiO <sub>2</sub>	600	Visible light	300	78, 30	135
тс	0.01	BiVO <sub>4</sub> /TiO <sub>2</sub> /RGO	-	Visible light, pH 3	Xe, 1000	96.2, 120	136
СТС	0.01	BiVO <sub>4</sub> /TiO <sub>2</sub> /RGO	-	Visible light, pH 3	Xe, 1000	97.5, 120	136
отс	0.01	BiVO <sub>4</sub> /TiO <sub>2</sub> /RGO	-	Visible light, pH 3	Xe, 1000	98.7, 120	136
DXC	0.01	BiVO <sub>4</sub> /TiO <sub>2</sub> /RGO	-	Visible light, pH 3	Xe, 1000	99.6, 120	136
CFX	20	MnWO <sub>4</sub> /Bi <sub>2</sub> S <sub>3</sub>	1200	Visible light	Xe, 1000	78.8, 180	137
MTZ	20	MnWO <sub>4</sub> /Bi <sub>2</sub> S <sub>3</sub>	1200	Visible light	Xe, 150	83.3, 180	137
тс	10	SnO <sub>2</sub> /CeO <sub>2</sub>	200	Visible light, pH 9- 10	500	97, 120	138
PG	5	ZnO	800	UV light, pH 6.8	24	72.72, 150	139
CIP	20	40TiHAp	2000	UV light, pH 6.1	125	100, 15	140
OFL	20	40TiHAp	2000	UV light, pH 6.1	125	100, 120	140

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ТС	10	Cu <sub>3</sub> P/ZnSnO <sub>3</sub> /g-C <sub>3</sub> N <sub>4</sub>	500	Visible light	Xe, 500	98.45, 60ew Article	141 Online
CEF	100	ACT-4	844	Visible light	LED bulb,	P9916,1260/D4NAC	051 <b>42</b> A
					50		
OFL	10	ZnO/Bi <sub>2</sub> MoO <sub>6</sub>	250	Visible light, pH	Daylight	100, 240	143
				7.54	lamp, 15		
ТС	10	MoO <sub>3</sub> /g-C <sub>3</sub> N <sub>4</sub>	500	Visible light	-	85, 9100	144
TC, CPX, and/or LFC solution	60	Bi <sub>2</sub> MoO <sub>6</sub> @Bi <sub>2</sub> MoO <sub>6-x</sub>	300	Visible light, pH 7	LEDs, 6.2	>95, 180	145
GMF	15–35	Zn-Co-LDH@BC	750	UV light, pH 5.5	10	92.7, 130	146
CFS	500	WO <sub>3</sub> /g-C <sub>3</sub> N <sub>4</sub>	25	UV light	Xe, 300	82, 120	147
Lomefloxacin	10	MgTiO <sub>3</sub>	30	UV light, pH 7	LED light,	83, 150	148
					30		
TC-HCI	20	Bi <sub>4</sub> Ti <sub>3</sub> O <sub>12</sub>	10	UV light	Xe, 300	75.5, 150	149
Cefazolin	20	TiO <sub>2</sub>	400	UV light, pH 5	15	96.47, 60	150
SMX	-	TiO <sub>2</sub>	0.002	UV light	Xenon,	88, 360	151
					1500		
SMX	10	TiO <sub>2</sub>	250	UV light, pH 4.1 –	UV, 9	Close to 100,	152
				5.4		120	
OA	20	Titanium Degussa P-25	0.001	pH 7.5	Black	90, 30	153
					light		
					lamp		
Norfloxacin	-	C-TiO <sub>2</sub>	0.0002	Visible light	-	78, 70	154
ТС	40	metal ion@TiO <sub>2</sub> /HNTs		Visible light	Xenon,	76.54, -	155
					500		
Chloramphenicol	50	TiO <sub>2</sub>	0.001	UV light	Osram	90,90	156
					Dulux, 9		
SMX	100	TiO <sub>2</sub>	0.0005	UV light	Xenon	80.360	157
					lamp		

\*CIP = Ciprofloxacine, AMX = Amoxicillin, TC = Tetracycline, APAP = Acetaminophen, CFX = Cephalexin, SMZ = Sulfamethoxazole, CFT = Cefoxitin sodium, OFL = Ofloxacine, NOR = Norfloxacinw, CTC = Chlorotetracycline, OTC = Oxytetracycline, DXC = Doxycycline, MTZ = Metronidazole, PG = Penicillin G, CEF = Cefixime, GMF = Gemifloxacin, CFS = Ceftazidime, SMX = sulfamethoxazole, OA = oxolinic acid



Fig. 2: Working procedure of the photocatalysts for the antibiotics degradation

# Other industrial pollutants (toluene, nitrobenzene, cyclohexene, and refinery oil)

Industrial chemical pollutants are a subgroup of chemical pollutants specifically connected with industrial operations <sup>158</sup>. They encompass a wide spectrum of chemicals used or produced in manufacturing, refining, and other industrial processes <sup>159</sup>. Industrial chemical pollutants, including toluene, cyclohexene, nitrobenzene, and refinery oil, pose significant

environmental threats due to their widespread use and high toxicity <sup>160,161</sup>. Toluene, an industrial solvent, pollutes air, water, and soil, causing harm to aquatic organisms and long-term environmental damage <sup>162</sup>. Cyclohexene, used in chemical production, contributes to air and water pollution, affecting aquatic life <sup>163</sup>. Nitrobenzene, a dye and pharmaceutical precursor, contaminates soil and water, posing toxic and carcinogenic risks <sup>164</sup>. Refinery oil, a byproduct of petroleum refining, causes extensive damage through spills and leaks, affecting marine and terrestrial ecosystems <sup>165</sup>. Photocatalytic degradation is crucial for mitigating these pollutants, as it offers

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an efficient, eco-friendly method to break down these toxic substances, preventing their persistence in the environment and safeguarding both ecosystems and human health  $^{\rm 166}.$ 

#### Toluene

As one of the pollutants that pose a risk to human health and the ecosystem, toluene has been classified as a priority pollution; for this reason, emission management is required <sup>167,168</sup>. Owing to the serious issues that toluene causes, various methods for toluene abatement have been developed <sup>169</sup>. The rapid growth in industrialization and urbanization has played a notable role in the emergence of severe environmental issues <sup>170,171</sup>. Toluene, a volatile organic molecule, can induce skin inflammation, respiratory ailments, chronic and acute intoxication, neurotoxicity, and reproductive toxicity 172-176. Therefore, it is necessary to enhance the efficacy of eliminating indoor toluene vapors. Methods to counteract atmospheric pollution can be classified as either chemical or physical approaches <sup>177,178</sup>. Physical approaches include adsorption, the process of condensation, and separating membranes. Chemical approaches encompass combustion, low-temperature plasma, biological, and photocatalytic treatments <sup>179,180</sup>. Photocatalysis is regarded as a very promising option for environmental cleaning among these techniques. Photocatalytic technologies provide the benefits of being non-toxic and cost-effective, requiring gentle reaction conditions, and producing no secondary pollutants <sup>136,181</sup>. Almost all the hydrocarbon degrades as the following mechanism <sup>182,183</sup>.

Photocatalyst + hv(photon)  $\rightarrow$  Photocatalyst ( $e_{cb}^{-}$  +  $h_{vb}^{+}$ ) (15)

 $e_{cb}^{-} + O_2 \rightarrow \cdot O_2^{-} \quad (16)$ 

 $h_{vb}^{+} + H_2O \rightarrow \cdot OH + H^+$  (17)

 $h_{vb}^{+} + OH^{-} \rightarrow OH$  (18)

Toluene +  $\cdot$ OH  $\rightarrow$  Hydroxylated intermediates (19)

Hydroxylated intermediates +  $\cdot$ OH  $\rightarrow$  Degradation products (20)

Degradation products +  $\cdot OH/O_2 \rightarrow CO_2 + H_2O$  + Inorganic ions (21)

Ming et al. utilized a hydrothermal technique to synthesize In<sub>2</sub>S<sub>3</sub> in a nanoscale form. This nanomaterial was then employed to fabricate a composite photocatalyst consisting of  $In_2S_3$  and g-C<sub>3</sub>N<sub>4</sub>. The process of toluene photocatalytic decomposition was investigated, and a feasible mechanism was proposed. The heterojunctions exhibited the  $\ln_2 S_3/g - C_3 N_4$ highest photocatalytic degradation when a 40% loading of  $In_2S_3$  was used <sup>184</sup>. Birgitta enhanced the TiO<sub>2</sub> catalyst by introducing sulfur, and nitrogen (S, N) components and reduced graphene oxide (rGO) through doping. The most efficient photocatalytic degradation of Toluene was achieved using a combination of 1wt% reduced graphene oxide (rGO) and 0.05wt% nitrogendoped titanium dioxide  $(N_{0.1}TiO_2)$  <sup>185</sup>. Van stated that the nanostructured Ir-doped TiO<sub>2</sub> is a highly effective photocatalyst that produces a superb material for reducing the risk of gaseous toluene. The material had a large surface area and had a consistently spherical shape of 10-15 nm in diameter <sup>186</sup>. The composite of PIL (poly ionic liquid) @TiO2 was formed using two different concentrations of polymerized ionic liquid (low and high). The composite was then assessed for its ability to degrade toluene. The findings indicated that 印印印印印印印 composite exhibited higher activity compared to the PIL (high) @TiO<sub>2</sub> composites <sup>187</sup>. Zhiguo synthesized a novel hierarchical heterostructured photocatalyst consisting of TiO<sub>2</sub>/Bi/Bi<sub>2</sub>MoO<sub>6</sub> using a solvothermal technique. On the outermost layer of flower-like Bi<sub>2</sub>MoO<sub>6</sub> nanospheres, the TiO<sub>2</sub> nanoparticles were evenly dispersed. The results suggest that the combination of TiO<sub>2</sub> can greatly improve the effectiveness of Toluene photocatalytic oxidation of toluene using the hierarchical heterostructure TiO<sub>2</sub>/Bi/Bi<sub>2</sub>MoO<sub>6</sub> <sup>188</sup>. Yuxi used zinc chloride  $(ZnCl_2)$ , zinc nitrate  $(Zn (NO_3)_2)$ , and zinc acetate  $(Zn (CH3COO)_2)$ to modify activated carbon fiber (ACF). Subsequently, titanium dioxide (TiO<sub>2</sub>) was loaded onto the modified ACFs. The study investigated the photocatalytic performance and adsorption of  $TiO_2/ACF-Ac$  modified by  $Zn(CH_3COO)_2$ ) highest for the removal of toluene <sup>189</sup>. The presence of a three-dimensional (3D) and directed structure enables efficient absorption of photons and rapid diffusion of volatile organic compounds (VOCs), surpassing the capabilities of catalysts in powder form. The researchers successfully created uniform and free-standing nanowire (NW) arrays of p-type Cu<sub>2</sub>O by subjecting Cu (OH)<sub>2</sub> NWs to heat treatment. The Cu<sub>2</sub>O NWs, as they are created, exhibit exceptional performance in degrading 30 ppm toluene, with a degradation rate of 99.9% achieved within 120 minutes <sup>190</sup>. Pejman et al. used a hydrothermal technique to deposit synthesized SrTiO<sub>3</sub> onto graphene oxide (GO). Photocatalysts that were artificially created were utilized for the process of gaseous toluene breaking down dynamically using photocatalysis while being exposed to UV radiation <sup>191</sup>. Rostami synthesized the TiO<sub>2</sub> and Bentonite photocatalyst by a method called co-precipitation and evaluated its catalytic efficiency in degrading Para Nitrotoluene (PNT) <sup>192</sup>. Oxygen vacancies (OVs) can regulate photocatalytic activity by altering their electrical and/or band structures. A wide bandgap p-block metal combination containing OVs, indium oxyhydroxide (InOOH), was produced using a one-pot hydrothermal approach was used to investigate the effect of OVs in photocatalytic decomposition and toluene ring breakage. Validated modified InOOH improves photocatalytic potency by decreasing the energy limitation of critical intermediates reaction during toluene degradation <sup>183</sup>. Xiaolong enhanced the performance of the C-USTiO<sub>2</sub> photocatalyst by applying it to carbon cloth and conducted a study on its ability to continuously degrade toluene under LED light exposure. The results demonstrated that the removal of the degraded toluene can exceed 80% when a large concentration of CO<sub>2</sub> is produced, and it exhibits exceptional cycle stability lasting for over 180 minutes <sup>193</sup>. Muyan et al. researched the use of CeO2 nanorods for the degradation of toluene using vacuum ultraviolet (VUV) catalytic oxidation. CeO<sub>2</sub> nanorods were utilized in a system that involved VUVphotolysis, UV-PCO, OZCO, and UVOZCO processes. Utilizing VUV light instead of ozone catalytic oxidation can significantly enhance the efficiencies, increasing them from 12.9% to 83.2% when combined with the suggested catalyst <sup>194</sup>. An efficient electrochemical method consisting of two steps was devised to produce a nanotube array of atomically dispersed Au-loaded

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 $WO_3/TiO_2$  for the oxidation of volatile organic compounds (VOCs). The presence of vacancies (OVs) on the surface of  $WO_3$ greatly improved the separation and movement of photogenerated carriers, as well as the adsorption of toluene. This resulted in an 85.5% mineralization and 95.4% degradation rate for the removal of toluene <sup>195</sup>. Jinze fabricated a hollow heterophase junction by applying a layer of amorphous TiO<sub>2</sub> onto anatase TiO<sub>2</sub> hollow spheres. The findings demonstrated that the application of the amorphous TiO<sub>2</sub> coating resulted in an augmentation of fine pores and intermediatevporesting the photocatalyst, leading to an improved capacity?for the photocatalyst, leading nanodiamonds to ZnO, the photo corrosion problem can be solved for photocatalytic degradation of gaseous toluene. Nanodiamond decorating resulted in lowered photoluminescence intensity and electrochemical impedance, enhancing ZnO light absorption, charge transfer, and toluene photocatalytic oxidation efficiency <sup>197</sup>.

**Table 3:** Data for the photocatalytic degradation of Toluene using various catalysts

Toluene	Catalyst	Catalyst	Operational	Lamp, Power	Degradation	Ref.
Conc.		Conc.	Condition	(W)	Percentage (%),	
(ppm)		(mg/L)			Time (min)	
60	$In_2S_3/g-C_3N_4$	50	Visible light, RH	Xe	89.7, 180	184
			50%–60%			
2	1wt%rGO/S <sub>0.05</sub> N <sub>0.1</sub> TiO <sub>2</sub>	500	Visible light, RH	Fluorescent,	72, 480	185
			60%	10		
1900	Ir Doped-TiO <sub>2</sub>	100	UV light, RH 70%	UV, 25	97, 8.5	186
50	PIL@TiO₂/m-GO	1000	UV, RH 40%	UV, 8	97, 24	187
-	TiO <sub>2</sub> /Bi/Bi <sub>2</sub> MoO <sub>6</sub>	2000	UV	Xe, 320	26.08, 120	188
843	$TiO_2$ / ACF-Ac fabricated by Zn	151.2	UV, RH 40%	Xe, 300	70, 2400	189
	(CH <sub>3</sub> COO) <sub>2</sub> )					
30	Cu <sub>2</sub> O NWs	172.26	Visible, RH 74%	Xe, 300	99.9, 120	190
60	SrTiO₃/rGO	400	UV light, RH 50%	UV, 8	98.65	191
50	TiO <sub>2</sub> /Bentonite	200	UV light	UV lamp	64, 120	192
50	InOOH	400	UV light	Xe, 300	75.8, 60	183
30	C-USTIO <sub>2</sub>	100	visible light, RH	LED, 1	80, 180	193
			50%			
30	CeO <sub>2</sub>	1000	VUV light, RH	VUV, 4	83.2, 144	194
			50%			
300	WO <sub>3</sub> /TiO <sub>2</sub>	-	LED light	-	95.4, 30	195
23.6	THS@Amorphous-TiO <sub>2</sub>	10	UV light	UV, 8	98.2, 240	196
50 ppm	TiO <sub>2</sub> /ND	100	UV	Xe, 50	100, 120	197
750	ZnAl <sub>2</sub> O <sub>4</sub>	-	UV	Black-light	90.25, 300	198
				fluorescent		
				lamp		
-	TiO <sub>2</sub>	-	UV, RH 35%	Germicidal	61.9, 180	199
				lamp, 15		
160	TiO <sub>2</sub>	-	UV, RH 25% -	Black lamp,	50-60, 5	200
			50%	10		
400	TiO <sub>2</sub>		UV	Iron	52,360	201
				halogenide		
				lamp, 500		
	TiO <sub>2</sub>		UV	Mercury	90, 120	202
				lamp, 300		

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Fig 3: Working procedure of the photocatalysts for the toluene degradation

#### Nitrobenzene

Since aromatic nitro compounds are frequently employed in industrial processes (such as the production of explosives, dyes as well as insecticides), they are present as contaminants in a variety of liquid sources, particularly surface water, and wastewater from industries <sup>203</sup>. Since nitrobenzene (NB) is identified as a significant contaminant, it is selected as a model pollutant. It is an extremely hazardous material and the highest permitted level of NB is 1 mg/L in wastewater <sup>204 205</sup>. Numerous factors, including the presence of anions, pH, light wavelength, and others, have an impact on nitrobenzene photocatalytic degradation utilizing UV radiation <sup>206</sup>.The degradation working mechanism of nitrobenzene in the presence of several photocatalysts is described <sup>207,208</sup>.

Photocatalyst + hv (photon)  $\rightarrow$  Photocatalyst ( $e_{cb}^- + h_{vb}^+$ ) (22)  $e_{cb}^- + O_2 \rightarrow \cdot O_2^-$  (23)

Nitrobenzene  $\rightarrow$  Catalyst surface (24)

Nitrobenzene +  $\cdot$ OH  $\rightarrow$  Activated nitrobenzene (25)

Activated nitrobenzene +  $\cdot$ OH  $\rightarrow$  Nitrophenol + Intermediate products (26)

Nitrophenol +  $\cdot$ OH  $\rightarrow$  Degraded products (27)

Degraded products  $+ \cdot OH \rightarrow CO_2 + H_2O +$  Inorganic ions (28) The study of the impacts of several factors, such as pH, anions, starting concentration, etc. has been introduced because the rate of breakdown of nitrobenzene utilizing controlled UV

radiation is guite significant when compared to solar radiation, a small amount of TiO<sub>2</sub> (0.05%, w/v) was used  $^{209,210}$ . Degussa P-25 TiO<sub>2</sub> was utilized as the photocatalyst in the majority of the nitrobenzene photocatalytic tests. Aldrich-TiO<sub>2</sub> (pure anatase with a BET surface area of roughly 250 m<sup>2</sup> g<sup>-1</sup>) was used in a few tests <sup>206</sup>. Matthews et al. used immobilized TiO<sub>2</sub> in a spiralshaped reactor for the photocatalytic degradation of NB and other chemicals & accomplished the degradation around 95% -100% at the initial concentration between 1.75 to 4.25 mg/L <sup>211</sup>. Degussa P-25 was applied as the catalyst in photocatalytic degradation tests, and UV lamps with lights radiating at  $\lambda$ max 253 and 365 nm, respectively, were used. The two bulbs produced nearly identical deterioration <sup>212</sup>. When it comes to 4chlorophenol degradation, it has been discovered that utilizing pulsed photocatalysis makes little distinction in terms of TOC elimination at shorter and longer wavelengths. It should be mentioned that 387 nm is the  $\lambda_{min}$  for anatase TiO<sub>2</sub> <sup>213</sup>. The pH has an impact on the ionizable organic molecules' photocatalytic breakdown. The significance of pH on the photocatalytic destruction of NB was assessed within a pH value range of 4–10, in a solution containing  $2.52 \times 10^{-4}$  M of pollutants. The ideal photocatalyst concentration was determined to be 0.5 wt.%  $Fe-TiO_2 = 250 mg/L$ , with an irradiation period of 60-240 minutes <sup>214</sup>. It has been discovered that, given the specified conditions, pH 7 is ideal for NB photocatalytic breakdown <sup>205</sup>.

able 4: Data for the catal	vtic degradation of nitrobenzene	using various catalysts

Nitrobenzene	Catalyst	Catalyst	Operationa	Lamp, Power (W)	Degradation	Ref.
Conc. (mg/L)		Conc.	l Conditions		Percentage (%),	
		(mg/L)			Time (min)	
50	$SrFeO_{3-\delta}$	0.001	UV	Mercury vapour, 125	99%, 360	215
50	P-25		UV	125	95%, 480	216
40	AuNPs/HPW/TiO <sub>2</sub> -NTs		Visible Light	Low-pressure mercury	90%, 30	217
				vapor lamp, 15		
	Ag/ZnO nanoflowers		UV	Tungsten lamp of 60	98%, 100	218
61.5	TiO <sub>2</sub>		Visible Light	125	58.46%, 210	219
	TiO <sub>2</sub> /g-C <sub>3</sub> N <sub>4</sub> /G	7.5	UV	Xenon lamp, 300	97%, 240	220
25	H <sub>3</sub> PW <sub>12</sub> O <sub>40</sub> supported on	10	Visible Light	Tungsten light, 500	88%, 390	221
	TiO <sub>2</sub>					
50			UV	Mercury vapor lamp, 200	96%, 240	222
50	TiO <sub>2</sub> -POMs		UV		86.4%, 180	223
1900	TiO <sub>2</sub> -SA-Arg particles		UV	UV Lamp	93.7%, 120	224



Fig. 4: Working procedure of the photocatalysts for the nitrobenzene degradation

#### Cyclohexane

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A common volatile organic compound (VOC) that presents significant dangers to both humans, as well as the surroundings, is cyclohexane <sup>225</sup>. An extremely significant industrial procedure is the breakdown of cyclohexane to produce cyclohexanol as well as cyclohexanone which is utilized globally as chemical precursors for the synthesis of caprolactam and adipic acid <sup>226</sup>, <sup>227</sup>. Photocatalytic techniques for the degradation of cyclohexane in both solid heterogeneous as well as homogenous stages have received a lot of research in recent years <sup>228</sup>. In heterogeneous environments, semiconductors along with oxides are being used as photocatalysts to oxidize cyclohexane. A number of semiconductors have been used, including CeO<sub>2</sub>, WO<sub>3</sub>, Sn/Sb, ZrO<sub>2</sub>, ZnO, V<sub>2</sub>O<sub>5</sub>, SnO<sub>2</sub>, Sb<sub>2</sub>O<sub>4</sub> and mixed oxides 229. In the presence of various types of photocatalysts, cyclohexene degrades as the following mechanism <sup>230,231</sup>.

$$\begin{split} & \text{Photocatalyst} + hv(\text{photon}) \rightarrow \text{Photocatalyst} \left( e_{cb}^{-} + h_{vb}^{+} \right) (29) \\ & h_{vb}^{+} + H_2O \rightarrow \cdot OH + H^+ (30) \\ & h_{vb}^{+} + OH^- \rightarrow \cdot OH (31) \end{split}$$

Cyclohexene + $\cdot$ OH $\rightarrow$ Intermediate products (32)
Intermediate products + $\cdot$ OH $\rightarrow$ Further degraded products
(33)

Intermediate products +  $\cdot OH/O_2 \rightarrow CO_2 + H_2O$  + Inorganic ions (34)

Xiao et al. discussed the photocatalytic characteristics of silver nanoparticles loaded on the nanocrystals of tungsten oxide when cyclohexane was being photo-catalytically degraded <sup>232</sup>. In standard manufacturing processes, cyclohexane is degraded at 150 °C using a homogenous cobalt-based catalyst <sup>228</sup>. Variations in the emitted photon flux and the irradiation wavelength during continuous irradiation result in notable variations in substance outputs and selectivity values during the photocatalytic degradation of cyclohexane by the help of TiO<sub>2</sub> in a pure liquid organic phase <sup>233</sup>. The photodegradation of cyclohexane with hydrogen peroxide at ambient temperature, assisted by a copper (II)-exchanged Y zeolite (CuY). Following 6 processing, hours of cyclohexanol and cyclohexyl hydroperoxide with 37% and 54% selectivities, respectively, resulted as the major products <sup>234</sup>.

**Table 5:** Data for the catalytic degradation of cyclohexane using various catalysts

Cyclohexene	Catalyst	Catalyst	<b>Operational Conditions</b>	Lamp, Power (W)	Degradation	Ref.
Conc. (mg/L)		Conc.			Percentage	
		(mg/L)			(%), Time (min)	
200 mg/L	Pt/TiO <sub>2</sub>		UV, Temp: 100 <sup>o</sup> C with	Osram Ultra-Vitalux	Close to 100%,	235
			0.5% Pt loading	lamp, 300		
	Ti <sup>3+</sup> self-doped TiO <sub>2</sub>		Visible Light, Temp: 40 °C	Xenon lamp	95%, 420	236
	WO <sub>3</sub> /Co-Pt	100	UV		93%, 720	237
	Degussa P-25	0.001	Visible Light	Metal halide lamp	Around 40%,	238
					180	
	WO <sub>3</sub> –TiO <sub>2</sub> mixed	50	Visible Light	Xenon lamp, 500	97%, 60	239
	catalysts					
	Au/TiO <sub>2</sub>	0.001	Visible Light, Temp: 25 °C	Mercury lamp of 50	50%,	240
	Degussa P25	0.001	UV, Temp: 650 <sup>0</sup> C	Xenon lamp, 450	, 60	241
523	TiO <sub>2</sub>		UV, Temp: 30 <sup>o</sup> C	Black light lamp, 20	63%, 5	242
	TiO <sub>2</sub>	0.001	UV,	Mercury lamp, 50	, 10	240
	Ag-substituted and	0.001	UV, Temp: Below 35 <sup>0</sup> C	A high-pressure	Around 10%, 6	243
	impregnated nano-			mercury vapor		
	TiO2			lamp, 80		

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 TiO <sub>2</sub>		UV, Temp: 140 ºC – 180 ºC	A high-pressure	<b>Over 90%,</b> <sub>View A</sub>	244 rticle Online
			mercury lamp, 100	DOI: 10.1039/D4	NA00517A
 TiO <sub>2</sub>	0.001	UV, Temp: 60 ºC	Medium pressure	Over 95%,	245
			mercury-vapor		
			lamp, 450		
 Na <sub>4</sub> W <sub>10</sub> O <sub>32</sub>	0.05	UV,	Medium pressure	, 3	246
			mercury-arc lamp,		
			125		
 Fe-modified C-		Visible Light, Temp: 25 <sup>0</sup> C	Xenon lamp, 300	, 5	247
doped Cr <sub>2</sub> O <sub>3</sub>					



Fig. 5: Working procedure of the photocatalysts for the cyclohexane degradation

#### **Refinery Oil**

Several methods may be used for the treatment of oil refinery effluents which include adsorption, Fenton oxidation, electro floatation-coagulation, photocatalytic degradation/oxidation, chemical flocculation-coagulation, and membrane filtration <sup>248-</sup> <sup>253</sup>. These procedures either produce insignificant impurities or need prolonged durations to eradicate the impurities <sup>254-256</sup>. Conventional methods like adsorption or membrane separation produce an inferior contaminant by moving the contamination from one phase to another, and the reusability of adsorbents is uncertain <sup>257,258</sup>. Bacterial degradation requires a significant amount of time to break down pollutants and is not suitable for the majority of organic compounds found in oil refinery wastewater <sup>259-261</sup>. Photocatalytic degradation techniques contain significant attention due to their ability to break down wide range of organic compounds utilizing suitable photocatalysts 52,262,263. The degradation of pollutant chemicals is caused by the hydroxyl radical (OH), which can react with organic compounds and break them down and degrade them <sup>264,265</sup>. The mechanism for refinery oil degradation in the presence of various photocatalysts is given <sup>266,267</sup>.

nanocomposites with varying amounts levels of BiOI deposited

via sequential ionic layer adsorption and reaction (SILAR) perform well in water under visible (>400nm) irradiation for crude oil degradation. The BiOI/TiO2 heterojunction separates photogenerated charges, improving degradation efficiency <sup>268</sup>. Actual wastewater from a refinery, containing a variety of aromatic and aliphatic organic compounds, was treated using nanoparticles (specifically TiO<sub>2</sub> and ZnO). The degradation ability of the organic contaminants was reduced from 98.57% to 89.482% when the photocatalysts changed from  $TiO_2$  to  $ZnO^{-267}$ . Dheeaa investigated the application of the ZnO/TiO<sub>2</sub>/H<sub>2</sub>O<sub>2</sub> using visible light (1000 W/m<sup>2</sup>), to decrease the total organic carbon (TOC) content in the actual petroleum wastewater obtained from Sohar Refinery Company (SRC). The treatment efficiency for total organic carbon (TOC) at pH 5.5 increased at significant percentage compared to the TiO<sub>2</sub> procedure <sup>269</sup>. Zahra et al. examined the photocatalytic oxidation of organic contaminants in petroleum refinery wastewater (PRWW) utilizing synthesized nano-TiO<sub>2</sub> incorporated into Fe-ZSM-5 zeolite and UV light. Results indicate optimal photodegradation efficiency at 3 g'l-1 photocatalyst concentration, pH 4, 45°C temperature, and 120 min UV irradiation <sup>270</sup>. Shahrezaei investigated TiO<sub>2</sub> photocatalysis for the primary degradation of phenol and phenolic compounds in refinery wastewater. At optimal conditions, 90% phenol removal was achieved in 2 hours <sup>271</sup>. The user created a composite membrane by combining polyvinylidene and titanium dioxide (PVDF/TiO<sub>2</sub>) and then treated it using the hot-pressed method. This treatment was done to increase the bonding between the TiO<sub>2</sub> and the membrane surfaces, to employ the membrane to degrade oil in wastewater.

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Table 6: Data for the photocatalytic degradation of refinery oil using various catalysts

Target compound	Compound Conc. (ppm)	Photocatalyst	Catalyst Conc. (mg/L)	Operation Condition	Lamp, Power (W)	Degradation Percentage (%), Time (min)	Ref.
Crude oil	200	BiOI/TiO <sub>2</sub>		isible light	LED, 13	85.62, 180	268
Refinery Oil	99.64	TiO <sub>2</sub>	100	UV light, pH	UV, 11	98.57, 120	267
Refinery Oil	99.64	ZnO	100	UV light, pH	UV, 11	89.48, 120	267
Oil in petroleum wastewater	15-22	TiO <sub>2</sub> /ZnO/ H <sub>2</sub> O <sub>2</sub>	H <sub>2</sub> O <sub>2</sub> =850 ZnO=590 TiO <sub>2</sub> =700	visible light, pH 5.5	LED, 1000	37	269
Petroleum refinery wastewater	500	TiO <sub>2</sub> /Fe-ZSM-5	3000	UV light, pH 4	UV, 8	66%, 120	270
Phenol	220	TiO <sub>2</sub>	100	UV light, pH 4	UV, 400	90%, 120	271
Phenol	70	Degussa P25	0.002-0.008	UV light, pH 5.5	Fluorescent T8 backlight- blue bulb, 18	76%, 90	272
Soap oil & grease	480	Degussa P25	0.002-0.008	UV light, pH 5.5	Fluorescent T8 backlight- blue bulb, 18	88%, 90	272
Refinery Oil		TiO <sub>2</sub>	100	UV light, pH 3	UV, 400	93.92%, 60	273
Refinery Oil		TiO <sub>2</sub>	0.0012	UV light, pH 4	UV, 11	40.68%, 120	274
Petroleum refinery wastewater		TiO <sub>2</sub>	100	UV light, pH 10	Mercury Vapor, 6	TOC = 32% & TN = 67 %, 90	275
Petroleum refinery wastewater		TiO₂/ZnO/ Degussa P25	0.0005- 0.005	UV light, pH 3.5 - 9	Mercury Vapor, 250	Phenols = 93% Dissolved organic carbon (DOC) = 63% Oil and grease (OG) = Over 50%, 60	276
Petroleum refinery wastewater	400-600	TiO <sub>2</sub>	100	UV light, pH 3, Temp: 45 °C	Mercury, 400	90%, 4	277
Petroleum refinery wastewater	Phenol = 0.002 COD = 1954 Oil & Grease = 298.8 Sulfide = 13.3	TiO <sub>2</sub>	0.001	O₃/UV, pH 7.16	Mercury, 100	Phenol = 99.9% COD = 31.6 % Oil & Grease = 5.2 % Sulfide = 53%, 5	278
Phenol	10	TiO <sub>2</sub>		UV, pH = 5		, 6	279
		•	•		•		

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Refinary Oil		TiO <sub>2</sub>		UV light, pH	UV, 60	83%, View Art	280 icle Online
				7-9		DOI: 10.1039/D4N	AUU517A
Petroleum	200-220	TiO <sub>2</sub>	100	UV light, pH	Mercury,	78%, 120	281
refinery				3, Temp: 45	400		
wastewater				°C			
Crude Oil	0.005	TiO <sub>2</sub> in Zeolite		UV light,	Mercury,	Linear alkanes =	282
				Temp: 400 <sup>o</sup> C	150	98.66%	
						Branched alkanes	
						= 97.31%	
						Cyclic alkanes =	
						96.04%	
						Aromatic	
						compounds =	
						99.54% Alkenes=	
						98.38% ,100	
Oil emulsion	25	Degussa P25	0.002	UV light, pH	Mercury,	In distilled water	283
in distilled/				4.5		92% & in artificial	
sea water						seawater 43%, 3	
Petroleum	COD = 1326	Green	0.00025	UV light,	UV,	99.9%, 90	284
refinery		Nanocatalyst					
wastewater		from the sepals					
		of waste tomato					
A synthetic	10,000	TiO <sub>2</sub>	0.002	UV light,	T8 black	68%, 30	285
oil-water					light blue		
emulsion					bulb, 18		
Refinery	100	4-Chlorophenol	178.5	UV light, pH	Mercury	, 80	286
wastewater				5	Lamp, 100		



Fig. 6: Working procedure of the photocatalysts for the refinery oil degradation

# Effects of crystal size & surface area in photocatalytic degradation

Organic chemicals and the photocatalyst's surface coverage are directly correlated, therefore surface morphology, such as crystal size and the surface area must be taken into account during the photocatalytic degradation procedure <sup>287,288</sup>. Since every chemical process occurs at the surface, the surface morphology of any photocatalyst is essential to its efficacy as a catalyst <sup>289</sup>. The anatase phase with a range of 2.59 to 12.00 nm in TiO<sub>2</sub> crystallite dimensions is visible in metal-doped TiO2 products. TiO<sub>2</sub> has a specific surface area of between 100 and 500 m<sup>2</sup>/g <sup>290,291</sup>. Sivalingam et al. used a

solution combustion process where 8–10 nm pure anatase phase TiO<sub>2</sub> with 156 m<sup>2</sup>/g BET surface area was created. This TiO<sub>2</sub> is commonly utilized for photocatalytic degradation of many dyes, including Orange G, Methylene Blue, Alizarin S, Methyl Red, and Congo Red. In this analysis, the crystal size of the photocatalyst was found as 8±2 nm <sup>292</sup>. The photoactivity of the photocatalysts increased due to the higher surface area. It has been found that the photoactivity of the TiO<sub>2</sub> while degrading the dye-like MB increased when the surface area of the catalyst increased from 63 m<sup>2</sup>/g to 156 m<sup>2</sup>/g <sup>293</sup>. For the maximum degradation of antibiotics like cefoxitin sodium, a novel BN/CdAl<sub>2</sub>O<sub>4</sub> composite with a surface area of 14.34 m<sup>2</sup> /g is used <sup>133</sup>. Mushtaq et al. found a decrease in the degradation rate of the ofloxacin antibiotic due to the increase in the particle size & decrease in the surface area of the photocatalysts <sup>294</sup>. The same

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scenario was also found during the advanced degradation of tetracycline antibiotics s by graphene-ordered mesoporous silica <sup>295</sup>. Zhou et al. used highly photoactive mesoporous anatase nanospheres that have a high specific surface area of 609 m<sup>2</sup>/g for the degradation of the toluene <sup>296</sup>. The highest specific surface area (130.3 m<sup>2</sup>/g) of nano-sized TiO<sub>2</sub> particles synthesized under ideal conditions is almost double that of Degussa P25 which is used for toluene degradation <sup>297</sup>. Rajesh et al. experimented with the degradation of nitrobenzene using nanocrystalline TiO<sub>2</sub> of different surface areas i.e., 259/199/166/124/91/2 m<sup>2</sup>/g <sup>216</sup>. Photocatalytic oxidation of cyclohexane over TiO<sub>2</sub> nanoparticles by molecular oxygen was carried out using different surface area photocatalysts ranging between 30-410 m<sup>2</sup>/g <sup>298</sup>. TiO<sub>2</sub> is made up of anatase and rutile with a mean particle size of 30 nm and a surface area of  $50 \text{ m}^2/\text{g}$ for the maximum degradation of refinery oil <sup>285</sup>.

#### Mechanism of photocatalytic degradation, Article Online

Photocatalytic degradation is a process where light energy, typically from UV or visible light, activates a photocatalyst, such as titanium dioxide (TiO<sub>2</sub>). When the photocatalyst absorbs light, it generates electron-hole pairs. These electron-hole pairs can initiate redox reactions that produce reactive oxygen species (ROS) like hydroxyl radicals and superoxide anions. These ROS are highly reactive and can break down organic pollutants, converting them into less harmful substances like water, carbon dioxide, and inorganic ions. The overall mechanism involves light absorption, generation of electronhole pairs, formation of ROS, and degradation of pollutants.



Fig. 7: Illustration of (a) Formation of free radicals, (b) Degradation of the organic pollutants by radicals, and (c) Overall photocatalytic degradation mechanism

#### Conclusion

Various photocatalysts are used depending on the variation in the organic pollutants. Titanium Dioxide (TiO<sub>2</sub>) is the most broadly applied photocatalyst, known for its maximum ability, stability, and non-toxicity. It is primarily activated by UV light. Zinc Oxide (ZnO) is another effective photocatalyst with properties similar to TiO<sub>2</sub> but with some advantages in certain conditions. Recent research includes materials like cadmium

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sulfide (CdS), tungsten oxide (WO<sub>3</sub>), and various metal-organic frameworks (MOFs) as effective photocatalysts. Scientists are working on photocatalysts that are triggered by visible light so that it is possible to improve the process's applicability and reduce energy consumption in the real world. This review is the scrutiny of variance in the degradation rate of the organic pollutants incorporating different conditions such as different pH levels, different concentration levels, various composite of the photocatalysts, different surface area & size of the photocatalysts, and so on. This review will help to identify the optimum parameters for the maximum amount of organic pollutant degradation. The goal of this field's ongoing research and development is to broaden the use of catalytic technologies and overcome current obstacles to ensure cleaner soil and water thus leading to a more sustainable environment. Greater prospects for the use of photocatalysis in the destruction of dangerous organic pollutants may arise from a greater understanding of the process and its operating parameters.

#### **Author Contributions**

Abdullah Al Miad and Shassatha Paul Saikat collected the data and wrote the draft and original manuscript. Md. Sahadat Hossain conceived and designed the review, analyzed the data, and assisted in writing the manuscript. Md. Kawcher Alam assisted in collecting data. Newaz Mohammed Bahadur and Samina Ahmed supervised the findings of this work. Samina Ahmed supervised the overall work and managed the required facilities.

#### **Conflicts of interest**

There are no conflicts to declare.

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