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Article UV Light-Irradiated Photocatalytic Degradation of Coffee Processing Wastewater Using TiO₂ as a Catalyst

Gurudev Sujatha¹, Subramaniam Shanthakumar¹ and Fulvia Chiampo^{2,*}

- ¹ Department of Environmental and Water Resources Engineering, School of Civil Engineering, Vellore Institute of Technology (VIT), Vellore 632014, India; sujathagurudev@gmail.com (G.S.); shanthakumar.s@vit.ac.in (S.S.)
- ² Department of Applied Science and Technology, Politecnico di Torino, Corso Duca degli Abruzzi 24, 10129 Torino, Italy
- * Correspondence: fulvia.chiampo@polito.it; Tel.: +39-011-090-4685

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Abstract: The coffee industry generates a significant amount of wastewater that is rich in organic loads and is highly acidic. The present study investigates the potential of the heterogeneous photocatalytic oxidation process to reduce the pollutant load in coffee processing wastewater. The experimental runs were conducted to evaluate the effect of operative parameters such as pH, catalyst dosage, intensity of UV light irradiation, and addition of oxidant on Chemical Oxygen Demand (COD) and colour reduction. Significant results for COD and colour removal, 67%, and 70% respectively, were achieved at a pH of 4 with titanium dioxide (TiO₂), and a catalyst dosage of 500 mg/L, using four ultraviolet-C (UV-C) lamps of 16 W each. With the addition of hydrogen peroxide (H₂O₂) as an oxidant, the removal efficiency increased to 84% and 75% for COD and colour, respectively. Finally, the best results obtained by photocatalytic degradation using UV light were compared to those using solar light. Based on the investigation, it was inferred that the pollutant removal efficiency in coffee pulping wastewater was also considerably high under sunlight. These findings may have relevance in terms of application in countries where coffee processing is carried out and where sunlight irradiance is usually strong: the technique could be exploited to decrease the pollutant content of this wastewater sustainably.

Keywords: coffee pulping wastewater; photocatalytic degradation; titanium dioxide; Chemical Oxygen Demand; colour

1. Introduction

Coffee belongs to the genus *Coffea* of the *Rubiaceae* family, and the two well-known species of coffee grown are the Arabica and Robusta [1]. There has been a steady growth in world coffee production, reaching about 170 million 60 kg bags in the crop year 2019–2020, with India accounting for 3% of it [2]. Coffee is processed by dry and wet methods in which the wet method yields superior quality coffee compared to the dry one. In the conventional wet processing method, the coffee industry uses a massive quantity of water during the various stages of the process. Wet processing of coffee involves the removal of outer parts of the fruit, including the skin, pulp, mucilage layer, and the parchment. Thus, this is an essential step in coffee production that decides the quantity of water usage for coffee processing varies from 2.25 to 23 m³ per ton of processed fruit, suggesting the potential of wastewater generation [3,4].

The wastewater from wet coffee processing can be divided into two parts: (i) wastewater generated during the removal of mesocarp, that is, the pulp of the cherry which contains fermenting sugars,

and (ii) wastewater generated during fermentation and washing, which contains hydrolysed pectins. The resultant wastewater is acidic and rich in total suspended solids, which are biodegradable. Additionally, it contains proteins, carbohydrates, fibres, fats, caffeine, polyphenols, pectins, nitrates, ammonium, tannic acids, high levels of soluble organic matter, and deficient dissolved oxygen levels. It is characterized by unacceptable odour, colour, high Biological Oxygen Demand (BOD) and high Chemical Oxygen Demand (COD) [5–9]. The concentrations of pollutants vary with the quantity of water used when processing the fruits. Wastewater released from this operation without treatment has the potential to pollute the land and receiving water body, causing harmful effects on domestic users, aquatic life, livestock and the watercourse down the stream [10].

In developing countries, the disposal of wastewater from coffee processing is a severe environmental issue wherein the effluents are discharged unwisely into nearby natural water bodies which flow into rivers and/or infiltrate into groundwater and thus deteriorating the water qualities. There are several approaches, such as anaerobic–aerobic settling ponds, biogas reactors, land application by irrigation, and wetlands that are useful to control this pollution. In general, biological treatment systems could be suitable for this wastewater as it contains elevated BOD and COD [7]. However, the conventional biological treatment plants, usually adopted for municipal wastewater, should be modified to obtain a high pollution removal efficiency: at least, the neutralization (to increase the acidic pH) and the coagulation–flocculation steps (to remove colloids and recalcitrant suspended solids) should be added.

Though these approaches reduce the pollution level, the total cost makes it expensive for companies. Hence, there is a need to curb this problem through innovative and eco-friendly techniques. The heterogeneous photocatalysis has proved to be efficient for the treatment of wastewater [11–13]. It can degrade organic compounds such as alcohols, carboxylic acids, phenols into harmless products such as carbon dioxide and water [14,15].

Photocatalysis is considered an effective system for the mineralization of many organics through the generation of radicals such as HO• and $O_2^{\bullet-}$, reducing the organic load of effluents considerably using procedures with relatively low costs [16,17]. The principle involves the initial absorption of photons with energy equal to or greater than the bandgap energy of the semiconductor, leading to the formation of electrons and holes. The semiconductors are mainly titanium dioxide (TiO₂), zinc oxide (ZnO), cadmium sulphide (CdS), zinc sulphide (ZnS). Among these, TiO₂ in anatase form has been most commonly studied due to its ability to break down organic pollutants and achieve complete mineralization. Moreover, TiO₂ has many advantages over the other catalysts as it is easily available, relatively inexpensive, and chemically highly stable [18–20].

A chemical transformation due to acceleration by the catalyst with light is called photocatalysis. The reaction mechanisms are widely known and can be resumed by Equations (1)–(5) [21,22]:

$$TiO_2 + h\upsilon \rightarrow e_{cb}^{-} + h_{vb}^{+}$$
⁽¹⁾

$$h_{vb}^{+} + H_2 O \rightarrow H^+ + HO \bullet$$
⁽²⁾

$$h_{vb}^{+} + R \rightarrow \text{intermediates} \rightarrow CO_2 + H_2O$$
 (3)

$$\mathbf{e_{cb}}^- + \mathbf{O}_2 \to \mathbf{O}_2 \bullet^- \tag{4}$$

$$HO \bullet + R \rightarrow intermediates \rightarrow CO_2 + H_2O$$
 (5)

The hydroxyl radical has been pointed out as the main responsible species for the oxidative degradation of organic pollutants. Figure 1 shows the mechanism of photocatalysis on an irradiated TiO₂ particle.

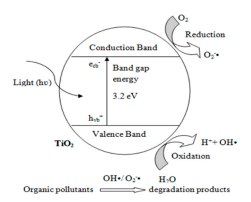


Figure 1. Photocatalytic mechanism for TiO₂ photocatalyst.

Besides, the presence of oxidants, for example, H_2O_2 , can improve the pollution removal, due to the generation of the hydroxyl radical:

$$e_{cb}^{-} + H_2O_2 \rightarrow HO\bullet + OH^{-}$$
(6)

The major factors affecting the TiO₂/UV light process are initial organic load, catalyst loading, reactor's design, temperature, solution pH, UV irradiation time, light intensity, and presence of ionic species. Previous studies on photocatalysis for the degradation of pollutants in industrial wastewater generated by the textile [23–25], refinery [26], pharmaceutics, and pesticides [27–29], cork [30] and olive mill [31] sectors reveal the effectiveness of the treatment method. However, the detailed literature review indicates that not many studies have been conducted on coffee pulping wastewater treatment using photocatalysis [32,33]. Satori and Kawase [32] studied the colour removal from coffee processing wastewater using ZnO particles. Their findings showed that the dissolved oxygen plays a role in colour abatement, achieving efficient removal only when its concentration was high. When the dissolved oxygen had small concentration or it was absent, the colour removal kinetics increasingly slowed down. The oxygen sparging would seem mandatory to obtain efficient removal, adding an operative cost. Zayas et al. [33] found that the combined process of chemical coagulation–flocculation advances oxidation processes, resulting in 87% abatement of COD and colour in the examined wastewater, with process time equal to 120 min. As a whole, the high efficiency is achieved with a complex and expensive sequence of operation, and from the application point of view, this could be an evident drawback.

The present study aimed to evaluate the technical feasibility of photocatalysis for the treatment of coffee pulping wastewater. The study was conducted by irradiating the samples with UV light in a photoreactor and with the sunlight in the presence of a catalyst. The effect of operative parameters such as pH, catalyst dosage, type of catalyst (TiO₂ and ZnO were tested), light power on the removal of COD, and colour were highlighted as preliminary results for the future process study on a pilot scale.

2. Materials and Methods

2.1. Characterization of Wastewater

Coffee wastewater was collected from the pulping process from a coffee processing industry located in Karnataka, India. The samples were collected in pre-cleaned bottles, stored at 4 °C in the icebox, and transferred to the laboratory for characterization. The physicochemical parameters such as colour, COD, BOD, pH, turbidity, total suspended solids (TSS), total dissolved solids (TDS), suspended solids (SS), nitrate-nitrogen and phosphate were analyzed as per the standard procedure prescribed by American Public Health Association (APHA) [34].

The samples were taken from the same huge tank at the company site; therefore, each experimental run was completed with wastewater having the same chemical and physical properties.

2.2. Catalyst and Reagents

The catalysts used in the photocatalytic degradation process were commercial titanium dioxide (TiO₂) and zinc oxide (ZnO). The chemicals used were analytical grade. Different concentrations (0.1 M and 1 M) of HCl and NaOH were used to adjust the initial pH. All the experiments were carried out using double distilled water.

2.3. Experimental Set-Up

The experimental set up consisted of a cubical container (L = 45 cm) with 4 UV-C lamps, with 16 W each and a peak wavelength of 254 nm. The container walls were covered with Al foil for light reflection. Two exhaust fans were fixed on each side of the reactor walls to maintain the temperature inside the reactor. A 1000 mL beaker was placed on a magnetic stirrer inside the container.

2.4. Experimental Methodology

UV photocatalytic degradation of coffee processing wastewater was performed in batch mode for a constant test duration of 3 h. This time was chosen as precautionary, considering that several pollutant compounds contained into coffee wastewater are complex molecules (for example, proteins, phenols, pectins). The working volume of 500 mL was kept in the beaker placed on the magnetic stirrer inside the photoreactor. The various influencing parameters such as pH (2–10), dosage of the catalyst (125–2000 mg/L), UV light irradiation (16 W–64 W), and oxidant dose (600–1200 mg/L of H_2O_2) were studied by varying the range of each parameter and keeping constant the others. At the end of experimentation, samples of the reaction mixture were withdrawn and filtered using Whatman paper filter (grade no. 42, pore size = $2.5 \mu m$) and analyzed for COD and colour to compare it with the initially measured values of these parameters to evaluate the UV photocatalytic removal efficiency. At the optimized conditions of UV photocatalytic degradation of coffee processing wastewater, the experiments were carried out for the solar photocatalytic process by placing the photoreactor in any light condition to compare its performance to the UV photocatalytic one. The sunlight irradiance was measured by a laboratory lux meter at Vellore Institute of Technology (12.97" N, 79.16" E) when the tests were done, that is, in February. The tests were carried out when the sun had the best irradiance during the day, even if the constant value of irradiance could not be achieved. The coffee effluent COD was measured following the closed reflux titrimetric method using a COD mineralizer according to APHA standards [34], while decolourization was measured as a decrease in absorbance at 465 nm by a UV-VIS spectrophotometer. The percentage reduction in COD and colour of the samples were calculated using the Equations (7) and (8), respectively:

$$\text{COD removal } \% = \frac{C_i - C_f}{C_i} \times 100 \tag{7}$$

Colour removal % =
$$\frac{Cl_i - Cl_f}{Cl_i} \times 100$$
 (8)

where: C_i = initial COD (mg/L), C_f = final COD (mg/L), Cl_i = initial colour (colour unit), Cl_f = final colour (colour unit).

The experimental runs had no replicas, while the wastewater analyses were replicated twice. The results, as the average of the doubled replicated analyses, will be shown without error bars and standard deviations, since two replicates cannot support a reliable statistical analysis.

2.5. Kinetic Model for COD Removal

In some experimental runs, the COD concentration was monitored along the run, and this allowed us to model the process kinetics.

The pseudo-first-order model was applied, namely:

$$r = dC/dt = -kC \tag{9}$$

where r is the reaction rate, C is the residual COD concentration, t is time, k is the reaction rate constant. The integration of Equation (9) gives the well-known expression:

$$C(t) = C_0 \exp\left(-kt\right) \tag{10}$$

where C(t) is the COD concentration (mg/L) at time t, C₀ is the initial COD concentration (mg/L), k is the reaction rate constant (min⁻¹), t is time (min).

The kinetic rate is often used to give the half-life time $t_{1/2}$, that is, the time by which the initial concentration is reduced to 50%: at $t = t_{1/2}$, $C = C_0/2$. For the pseudo-first-order model, the half-life time depends only on the reaction rate constant, and it is equal to

$$t_{1/2} = \ln 2/k = 0.693/k \tag{11}$$

3. Results

3.1. Characteristics of Wastewater

The physicochemical properties of raw coffee pulping wastewater were analyzed, and the results are presented in Table 1. It can be noted that the wastewater is acidic with a high concentration of COD and colour, which imposes a challenge for the selection of treatment technology. As per the Central Pollution Control Board (CPCB) of Indian Statute, the maximum permitted value of BOD concentration discharge into natural water bodies is 1000 mg/L, and for land application, it is 100 mg/L.

Parameter	Value
pH	4.44
Colour Unit (CU)	1023
Electrical Conductivity (mS/cm)	1718
Total Dissolved solids (mg/L)	2110
Total Suspended Solids (mg/L)	4223
Total Solids (mg/L)	6333
Chemical Oxygen Demand (mg/L)	28,800
Biochemical Oxygen Demand (mg/L)	14,000
Phosphate (mg/L)	10
Nitrate (mg/L)	23
Nephelometric Turbidity Unit (NTU)	40.6

Table 1. Characteristics of the raw coffee pulping wastewater.

3.2. Effect of Initial pH

The effect of initial pH was one of the most significant factors which influence the photocatalytic degradation of wastewater, as it affects the charge on the catalyst particles, size of aggregates and the position of conductance and valence bands, thus affecting the adsorption of pollutants that happens at the surface of photocatalysts [35,36].

Figure 2 shows the effect of pH in the range of 2–10. It can be noted from the diagram that the maximum COD reduction and colour removal was achieved at the acidic pH of 4, which is about the pH of the raw coffee wastewater. At a pH of 8 and after 3 h of photocatalytic treatment, the COD and colour removal efficiency was 33% and 45%, respectively, while at a pH of 10, the process provided unsuitable degradation for future application, even if higher than zero. The COD removal was increased by up to 67.4% under acidic conditions (pH of 4) and was then decreased to 60% as pH was further reduced to 2. A similar trend was observed for colour removal. This can be attributed to the fact

that TiO_2 shows an amphoteric character, that is, positive or negative charge can be developed on its surface. The surface of the TiO_2 is positively charged under acidic conditions and negatively charged under alkaline conditions. In acidic media, electrons tend to move into the surface of a catalyst for the electrostatic attraction between the positive charged TiO_2 and electrons. They react with the oxygen molecules absorbed around the TiO_2 surface to form oxidizing species such as $O_2\bullet^-$ and $\bullet OOH$ [26]. The maximum oxidizing capacity of TiO_2 is thus at a lower pH, notwithstanding the decrease in reaction rate at a low pH due to excess H⁺ [37].

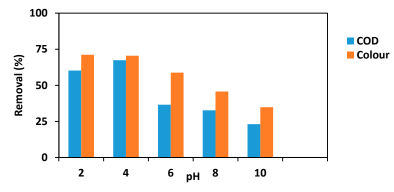
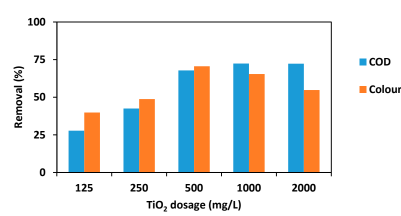


Figure 2. Effect of pH on photocatalytic degradation (process conditions: TiO₂ dosage—500 mg/L, reaction time—3 h, UV light irradiation—64 W).

Moreover, the catalyst has a pH of zero and a point charge of around 6.2, as given by the producer and confirmed by other scientific publications (it was not measured during the experimental runs). Considering the meaning of this parameter and concerning these results, at a pH < 6.2, the particle surface is covered with positive charges and can attract anionic compounds. The results are coherent with this, since the best results were achieved at an acidic pH, and at a pH of 6, the removal efficiency is lower for both the monitored parameters (COD and colour). Regarding coffee processing wastewater, many pollutants tend to be polar or in anionic form; for example, many polyphenols present negatively charged hydroxyphenyl groups that can react with the positively charged particles.

Therefore, the selection of pH needs to be appropriate to achieve maximum degradation efficiency.

3.3. Effect of Catalyst Dosage



At a pH of 4, the TiO₂ dosage was varied from 125 mg/L to 2000 mg/L, and the results are presented in Figure 3.

Figure 3. Effect of catalyst dosage on photocatalytic degradation (process conditions: pH—4, reaction time—3 h, UV light irradiation—64 W).

Regarding COD, it can be observed that as the dosage of the catalyst increases from 125 to 1000 mg/L, the percentage of its removal increases from 28% to 72%, with 72% also at a 2000 mg/L

dosage. The increased pollutant removal rate that follows the increase in the catalyst loading can be attributed to the fact that a more significant number of photons are adsorbed, thus accelerating the process.

About the reduction in colour, its trend shows the maximum at a 500 mg/L dosage, achieving 70%; with a further increase in the dosage, the colour removal efficiency decreases to 55% at a 2000 mg/L dosage. This is attributed to the rising of the light scattering due to the presence of TiO_2 particles excess [38]. Hence, the optimum dosage for the maximum degradation of coffee wastewater was considered as 500 mg/L.

3.4. Effect of UV Light Irradiation Power

The influence of light power on the removal efficiency was examined at a pH of 4 and a catalyst dosage of 500 mg/L for 3 h. The effect of UV light irradiation was analyzed by varying the output of UV lamp power from 16 W to 64 W (Figure 4). As expected, from the data, it can be observed that for both the parameters, the removal efficiency increases with an increase in the light irradiation, due to the generation of more electrons and holes [39]. The UV irradiation generates the photons required for the electron transfer from the valence band to the conduction one of a semiconductor photocatalyst. The energy of a photon is related to its wavelength, and the overall energy input to a photocatalytic process is dependent on light intensity. The degradation efficiency increases when more radiations fall on the catalyst surface, as more hydroxyl radicals are produced [40].

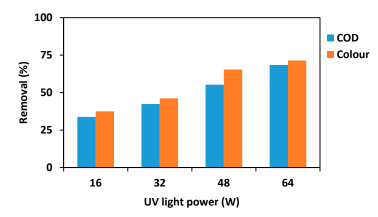


Figure 4. Effect of UV light irradiation power on photocatalytic degradation (process conditions: pH—4, reaction time—3 h, TiO₂ dosage—500 mg/L).

3.5. Effect of Catalyst Type

Although TiO_2 is the most commonly used active semiconductor photocatalyst, ZnO is a suitable alternative. ZnO has almost the same bandgap energy (3.2 eV) as TiO_2 , and its photocatalytic capacity is similar to that of TiO_2 [12]. Table 2 shows the results of degradation by the tested catalysts at optimized conditions. Figure 5 depicts the variation in the rate of degradation induced by TiO_2 and ZnO.

Table 2. Comparison of TiO₂ and ZnO on photocatalytic degradation (process conditions: pH—4, UV light irradiation—64 W, catalyst dosage—500 mg/L).

Description	% Removal		
Parameter	TiO ₂	ZnO	
COD	67.4	42.5	
Colour	70.2	48.7	

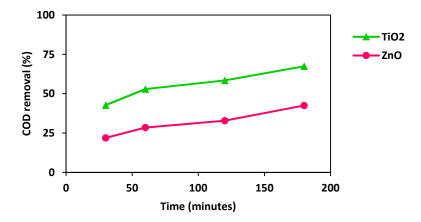


Figure 5. Effect of type of catalyst on photocatalytic degradation (process conditions: pH—4, reaction time—3 h, UV light irradiation—64 W, catalyst dosage—500 mg/L).

In the present study, zinc oxide showed lesser results compared to TiO_2 for the same conditions. Studies carried out by Sakthivel et al. [41] have revealed that the quantity of ZnO required to achieve the optimal photocatalytic activity is nearly double that for TiO_2 (anatase or rutile), and this supports the results obtained in the present study.

3.6. Effect of Oxidant Addition

The addition of hydrogen peroxide significantly improved the efficiency of photocatalytic degradation of organic compounds. The oxidative photocatalytic degradation was investigated using raw coffee effluent, at a pH of 4, a TiO₂ dosage of 500 mg/L, and varying the concentration of H₂O₂ from 60 mg/L to 1200 mg/L (Figure 6). It can be noted that the removal efficiency first increased when hydrogen peroxide concentration was increased up to 1000 mg/L, reaching 84% of COD removal efficiency. Beyond this, the degradation was reduced to 73% after 3 h of exposure. Radical reaction mechanisms can explain this dual effect of H₂O₂. The added H₂O₂ could accelerate the reaction by producing additional hydroxyl radicals (Equation (6)). The generation of additional HO• and other oxidizing species increases the oxidation rate of intermediate compounds. However, by the further addition of H₂O₂, that is, at a high oxidant concentration, a considerable amount of hydroxyl radicals can be consumed by the oxidant itself [37,40]:

$$H_2O_2 + HO \bullet \to H_2O + HOO \bullet$$
(12)

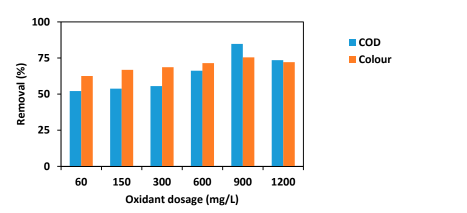


Figure 6. Effect of oxidant dose on photocatalytic degradation (process conditions: pH—4, reaction time—3 h, UV light irradiation—64 W, catalyst dose—500 mg/L).

The reaction described by Equation (12) is similar to a "parasite" reaction that consumes reactants.

3.7. Comparison of UV and Solar Photocatalytic Degradation

The utilization of sunlight for wastewater treatment is preferred over UV light due to its economics. Hence, in this study, the UV photocatalytic degradation was compared to the solar one (Figure 7). The average solar irradiance was found to be 716 W/m² during the daily period of experimentation. The pollutant removal efficiency was measured in terms of COD removal at different time intervals. Figure 7 shows that initially, the degradation is low in the solar photocatalytic process when compared to UV photocatalytic degradation efficiency, as the intensity of solar light was low. However, later, the pollutant removal increased up to 65% when the power of solar light reached its daily peak value (912 W/m²). Therefore, it can be inferred that solar photocatalytic degradation can be efficient in the regions receiving high radiation of solar light, reducing the energy cost, and making it sustainable.

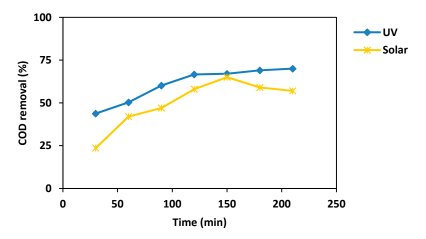


Figure 7. Influence of incident irradiance on COD removal over the 3.5-h duration test (process conditions: pH—4, UV light irradiation—64 W, catalyst dose—500 mg/L).

3.8. Kinetic Model for COD Removal

As shown in Section 2.5, the kinetics of the COD removal process can be described by the pseudo-first-order model. In this study, a couple of experimental runs were monitored along with the whole test duration, namely:

- Experimental run to check the effect of catalyst kind (green dots in Figure 5).
- Experimental run to check the influence of irradiance (blue dots in Figure 7).

By Equations (10) and (11), the kinetic rate constant and the half-life time were calculated, and Table 3 reports the results.

Table 3. Kinetic parameters for COD degradation by TiO_2 and ZnO (process conditions: pH—4, UV light irradiation—64 W, catalyst dosage—500 mg/L, C_0 —28,800 mg/L).

Catalyst	Run Duration (min)	k (min ⁻¹)	R ²	t _{1/2} (min)
TiO ₂	180	0.007	0.65	99
TiO ₂	210	0.008	0.71	87
ZnO	180	0.003	0.75	231
TiO ₂	180 and 210	0.007	0.49	99

For TiO₂, the experimental runs gave similar results, with the half-life time in the range of 87-99 min. In the bottom row, all the data for TiO₂ were modelled together.

As expected, ZnO had a much longer half-life time, showing the lower efficiency for COD removal of coffee processing wastewater.

4. Discussion

This study aimed to check the applicability of UV photocatalysis for the treatment of coffee processing wastewater with a TiO_2 as catalyst.

The experimental runs were carried out with industrial wastewater produced by an Indian company in Karnataka. In addition to the applicability of the photocatalysis, the effects of the operative parameters were tested, namely, initial wastewater pH, catalyst dosage, UV light power, kind of catalyst and oxidant dosage. Finally, the use of solar light instead of UV was tested, and the performances obtained in both cases were compared.

The application of photocatalysis to treat wastewater is not new. However, many laboratory studies were carried out usually with single pollutants characteristic of specific industrial sectors (textile, refinery, pharmaceutics, and personal care, paper, etc.), and very often focused on colour removal.

The novelty of this study was the use of original industrial wastewater, and the removal of the pollution load (expressed as COD) and colour were the targets.

The experimental runs demonstrated the effectiveness of photocatalysis for the treatment of coffee pulping wastewater. Moreover, the experimental findings allowed us to affirm the following:

- An acidic initial pH gave better removal results than neutral or basic ones for both the targeted parameters (COD and colour). For COD, the highest removal efficiency was achieved at a pH of 4, with the values showing a maximum in the acidic field, while for colour this was not evident. The parabolic trend is supported by other studies [42,43] that found a similar effect of pH on pollution removal, even if in different ranges of pH. Panchangam et al. [44] found that the removal of perfluoroalkyl substances (PFAS) by photocatalysis with several catalysts was enhanced when the pH was acidic. Coffee pulping wastewater has an acidic pH; therefore, considering large-scale applications, this finding is positive since chemical concentration in the process reactions, and this should be exploited to apply treatment processes to wastewater with different industrial origins. However, they also showed the need for further investigations on the physical mechanisms involved in the whole process;
- The tests on catalyst dosage showed that this operative parameter must be optimized. At values higher than the one that gave the best result, the findings showed that COD removal did not increase and colour removal was reduced, as found by other authors with different operative conditions and types of wastewater [41,45,46]. The reason for this can be poor light use due to scattering and reduced paths of irradiation. This occurs for all the catalysts, albeit at different extents.
- When UV light irradiation power increased, the removal efficiency also increased, as expected, due to higher hydroxyl radical generation.
- In coffee processing wastewater, TiO₂ as a photocatalyst showed better pollution removal efficiency than ZnO at the same dosage; this is in line with other studies with the aim of removing textile dyes [47].
- The oxidant addition increased the removal efficiency of the concentration by up to 900–1000 mg/L; however, further additions caused a slight decrease in the process performance. The effect was similar to that of the pH, and the results impose a more detailed investigation to optimize the use of oxidants and acid.
- Sunlight demonstrated to be able to substitute UV light, notwithstanding the limits for its use, namely, a narrow-spectrum with a short wavelength, daily variable irradiance, and irradiation intensity. This was also reported in other studies [48] that showed the rate of degradation for the pollutants was faster using the UV/TiO₂ process than using solar/TiO₂. The wavelength of the visible light is higher than that of UV light, and it is proved that there are higher possibilities of trapping the electron-hole pairs with shorter wavelengths. Moreover, the solar spectrum consists of only 5% UV light, limiting the opportunity to obtain very good results for TiO₂

photocatalysis [49]. In any case, the exploitation of sunlight must be considered for equatorial and tropical countries where solar irradiance is high.

To summarize, the present study demonstrated the applicability of UV photocatalytic degradation to coffee pulping wastewater, achieving relevant COD and colour removal efficiency using TiO₂ as a catalyst at a pH of 4, and catalyst dosage of 500 mg/L over a 3-h period. The addition of an oxidant (H₂O₂) enhanced the degradation to 84% and 75% for COD and colour, respectively. Moreover, for this application, TiO₂ was more effective than ZnO. Finally, but not with minor relevance, sunlight was used, and the results compared to those achieved with the UV catalysis, confirming a lower removal efficiency. However, most coffee-processing work is completed locally where it grows, that is, in equatorial/tropical countries where the sunlight is strong and its irradiance can be efficiently exploited for photocatalysis, making the process sustainable.

The light source substitution should not be disregarded due to environmental and economic features; optimization of the process with sunlight may be of interest in coffee-processing countries with high solar irradiance, as this would increase its sustainability.

The study gave preliminary results that are fundamental to continue with challenging investigations and the development of the process in view of a future scale-up. However, there is a need to obtain more statistical data, that is, additional experimental findings, for a robust transfer of scale.

Finally, this study can be a very small contribution to photocatalysis for the treatment of wastewater produced by different industrial sectors, as summarized in recent reviews [50–53].

Author Contributions: Conceptualization, S.S.; methodology, S.S.; validation, S.S.; formal analysis, S.S. and F.C.; investigation, G.S.; resources, S.S.; data curation, S.S.; writing—original draft preparation, S.S.; writing—review and editing, S.S. and F.C.; visualization, G.S.; supervision, S.S.; funding acquisition, S.S. All authors have read and agreed to the published version of the manuscript.

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