Posttreatment of Olive Mill Wastewater by Immobilized TiO₂ Photocatalysis

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ABSTRACT

A photocatalytic reactor with UV/TiO2 was used for the posttreatment of olive mill wastewater after anaerobic digestion. A factorial experimental design was adopted to determine the statistical significance of each parameter tested, namely, initial chemical oxygen demand (COD), pH, treatment time and recirculation flow and possible interactions in three response variables: phenols, color and COD removals. Removal efficiencies of 90.8 \pm 2.7%, 79.3 \pm 1.9% $50.3 \pm 6.3\%$ were obtained for total phenols (TPh), color and COD respectively. TPh and color were almost completely removed after 24 h of treatment, while the COD removal was partial. Because increasing the treatment time is economically unfeasible a recirculation to the anaerobic reactor should be considered. Regarding the most significant variables, the TPh removal efficiency is dependent of the initial COD concentration; the color removal efficiency decreased with increasing COD concentration and pH; and, the COD removal efficiency is directly linked with the treatment time. The interaction between the initial COD and treatment time affect negatively the response variables tested because of the inactivation of some active sites of the TiO₂ paper.

INTRODUCTION

Olive oil has been considered a fundamental ingredient of a healthy diet, especially in the Mediterranean area. A significant increase in production during the last years was observed. Therefore, one of the major concerns of olive oil industry is the treatment of high amounts of generated olive oil mill wastewater (OMW). Its annual production is in the range of some million cubic meters throughout the European Union (1). Oil mill wastewater (OMW) is heavy polluted and hence it cannot be disposed in the environment without any treatment (2). However, a common way of dealing with OMW is to discharging it directly into the sewer, confine it in a central lagoon or store it in a small pond nearby the mill, where it is left to evaporate until the next season. These ponds are often leaking causing ground water pollution and malodor. The setting up of more stringent regulations

(Table 1) concerning public waste disposal, imposed higher interests in development of new technologies and procedures for the purification of OMW (3). Among those processes, biological methods, whether aerobic (4) or anaerobic (5), have been recognized as viable options. Due to high contents of chemical oxygen demand (COD), up to 220 g L⁻¹ (6), OMW can be an excellent substrate for renewable energy production, such as in anaerobic digestion (AD) processes, which produce biogas (7) and biohydrogen.

The main problems of OMW are the high lipidic and phenolic compounds concentration. Phenols can go from 1 to 24 g L⁻¹ (8). These characteristics may cause an inhibitory effect when OMW is directly used in an anaerobic treatment. Phenolic compounds are considered one of the major problems associated with OMW not only due to their recalcitrance but also because of their strong brown to dark color. The combination of environmental and economic advantages makes the use of biological treatments and advanced oxidation processes (AOP) a suitable solution for the removal of toxic and colored compounds from water (2). AOP are based on the use of the hydroxyl radical (OH[•]) as primary oxidant of organic pollutants. Photocatalysis is an emerging AOP process (9). The main mechanism of photocatalysis is the formation of highly reactive and nonselective hydroxyl radicals upon irradiation of semiconductor particles with near UV radiation (wavelength $[\lambda]$ <400 nm). The photogenerated electron/hole pairs (e⁻/h⁺) are used as reducing/ oxidizing agents as they can migrate to the surface of the semiconductor particle to form oxidizing species in the presence of oxygen. The positively charged holes oxidize the organic compounds adsorbed on the catalyst surface. In addition, hydroxyl radicals are produced in aqueous system. The hydroxyl radicals serve as a strong oxidizing agent for the conversion of organic molecules to carbon dioxide (10). Due to its stability TiO2 is accepted to be one of the most suitable semiconductors for photocatalysis. It is the most studied, however, a large band gap (3.2 eV) of anatase TiO₂ restricts its use to the narrow lightresponse range of UV light ($\lambda < 400$ nm, only about 3–5% of total sunlight). Nevertheless, efforts are reported in the literature to decrease the band gap of TiO₂ and improve its activity under visible light (11,12). This can significantly improve the energy demand of advanced oxidation technologies and overcome its main drawback, i.e. high energetic costs. The fundamentals of photocatalysis, employing the semiconductor TiO₂ as catalyst, have been extensively reported (9,10,13). The process is suitable for wastewater remediation because it occurs under ambient conditions, it avoids the formation of photocatalyzed intermediate

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Table 1. Independent and response variables tested with respective initial high and low values used in the factorial experimental design and Portuguese legislation limits.

Variable	Units	Minimum (-1)	Maximum (+1)	Legislation limits*		
COD	${\rm mg~L}^{-1}$	1051	7370	150		
pН		4.8	7.5	6.0-9.0		
$t_{ m treatment}$	hour	8	24	n.a.		
$t_{ m contact}$	minute	1	2	n.a.		
TPh	$mg L^{-1}$	129	400	0.5		
Color	cm^{-1}	2.58	5.12	Undetectable at a dilution of 1:20		

^{*}Portuguese "Decreto-Lei no. 236/98, from 1/08/1998"— emission limit value that should not be exceeded for wastewater discharge. n.a.—not applicable.

stable products unlike direct photolysis techniques, and the oxidation of the substrates to CO₂ is complete. Because it is an inexpensive technology with a high turnover, it offers great potential as an industrial technology to detoxify wastewaters (10). Although TiO₂ is widely used in suspension with excellent performance, it requires expensive filtration to separate the catalyst from the treated water. To avoid this drawback, TiO₂ has been used immobilized on different types of supports (14,15).

Optimization of a photocatalytic process is difficult due to several interactions between the main variables controlling the process. To address the above concerns, and meet the demands of a structured research, proper experimental design techniques considering all factors and their possible interactions must be performed. Factorial experiments design allows to study the process behavior under conditions in which independent variables, called in this context factors, are varied simultaneously. Thus, it is possible to investigate the joint effect of two or more factors on a dependent variable. The factorial design also facilitates the study of interactions, giving insights on the effects of different conditions tested, on the identifiable subgroups of subjects participating in the experiment. The use of factorial designs leads to optimized parameters with a minimum set of experiments and to the possibility of obtaining a polynomial expression that describes the process yield (16).

The aim of this work is to study the photocatalytic oxidation of anaerobically pretreated OMW, regarding the effect of various operating conditions (initial COD, pH, treatment time and recirculation flow) on the removal of color, phenolic and organic compounds. A factorial design methodology was adopted to determine in a systematic way the statistical significance of each parameter and possible interactions.

MATERIALS AND METHODS

Olive mill wastewater. Olive oil mill wastewater used in these experiments was originated in a lab-scale up-flow anaerobic reactor of 2.5 dm³ (useful volume). The reactor consisted in an anaerobic chamber and an external jacket, which kept the temperature at 37°C. A solid–gas–liquid separator was connected at the middle of the reactor to avoid washout and to promote the degradation of the accumulated substrate onto the biomass (17). The effluent was immediately frozen until further use.

Photocatalytic reactor. The photocatalytic reactor (Fig. 1) consisted in a 37° slanted aluminum plate with a working area of 30 × 30 cm² (18,19). Coating it with a thin layer of polytetrafluoroethylene (PTFE) inactivated the aluminum surface. The solution to be treated flowed as a thin film from the top of the chamber over a nonwoven fabric made of cellulose fibers on which Tiona PC500 TiO₂ (18 g m²), UOP 2000 zeolite (2 g m²) and Snowtex 50 SiO₂ (20 g m²) have been fixed by compression (gift from Ahlstrom, Pont-Evêque, France). PC500 titania by Millennium Inorganic Chemicals was used (anatase: >99%; surface area >250 m² g¹¹, crystallites mean size = 5–10 nm). Titania PC500

was coated on nonwoven paper (natural and synthetic fibers, 2 mm thick) using an inorganic binder. The binder was an aqueous dispersion of colloidal SiO₂ (EP1069950B1 European patent) (19). A transparent glass sheet covered the reacting chamber to avoid evaporation of the solution. The treated sample (300 mL) was stored in a reservoir and continuously circulated in the system by a peristaltic pump, thereby permitting optimal distribution of the liquid over the catalytic support. PTFE tubing was used. The reservoir was open to air and the sample constantly agitated to ensure sufficient oxygenation. Artificial irradiation was provided by two UV lamps emitting light with a wavelength around 365 nm (F15T8, BLB 15W: Duke, Essen, Germany). The lights were positioned in parallel to the reactor. Light was turned on at the beginning of each experiment. After each experimental run, the photocatalytic reactor was rinsed with de-ionized water under UV irradiation. The photocatalytic reactor using an immobilized catalyst avoids the need of a final step of filtration for the extraction of the catalyst from the effluent.

Analytical methods. Total COD was determined using test kits (Hach Lange, Düsseldorf, Germany). Total and volatile suspended solids (TSS and VSS) were determined according to Standard Methods (20). Total phenolic (TPh) content was determined colorimetrically at 765 nm on a spectrophotometer (Hach Lange). The Folin-Ciocalteau reagent was used according to the procedures described in (21). Gallic acid monohydrate was used as standard to quantify the concentration of total phenols in OMW. UV–Visible spectrophotometry using a JASCO V-560 device in the range 200–800 nm, measured sample absorbance. Changes in sample absorbance at the wavelength of 390 nm were monitored to assess the extent of decolorization occurring during photocatalytic posttreatment.

Factorial experimental design. A statistical approach was chosen based on a factorial experimental design to infer about the effects of four variables on the treatment performance and possible interaction effects. Four independent variables (X_1-X_4) , which typically affect the efficiency of photocatalysis, in general, and the photocatalytic reactor used, in particular, were taken into account. The variables (Table 1) were the initial COD concentration, initial pH, treatment time ($t_{\rm treatment}$) and cycle duration expressed as the time that takes to the sample volume pass through the reactor ($t_{\rm contact} = {\rm Volume/Flow}$). In this study, the initial TPh concentration and color are dependent of the initial COD and therefore were not considered as independent variables. The pH limits imposed correspond to the pH of the samples coming from the anaerobic digestion step (+1) and the pH of the raw OMW (-1). It was adjusted with HCl 8 ${\rm m}$. The photocatalysis efficiency was assessed in terms of response factors, namely, TPh removal (Y_1), Color removal at $\lambda = 390$ nm (Y_2) and COD removal (Y_3).

The experimental design consisted in a full factorial experimental set with 16 (2^n , with n=4 variables) runs. The experiments were randomly performed. The software package Design-Expert [®] (Stat-Ease, Inc., Minneapolis) was used to perform the design matrix of the experiments and their statistical analysis. Estimation of the average effect, the main effects of each individual variable, as well as their interaction effects were determined according to Eq. (1)

$$Y_{i} = \beta_{0} + \sum \beta_{i} \chi_{i} + \sum \beta_{ij} \chi_{i} \chi_{j} + \sum \beta_{ijk} \chi_{i} \chi_{j} \chi_{k} + \beta_{ijkl} \chi_{i} \chi_{j} \chi_{k} \chi_{l}$$

$$\tag{1}$$

Where Y_i is the predicted response variable removal efficiency (%); β_0 is the constant coefficient; β_i is the coefficient of the variable X_i ; β_{ij} , β_{ijk} and β_{ijkl} are the interactions coefficients; and X_i , X_j , X_k and X_l are the independent variables. Removal efficiencies data were processed for

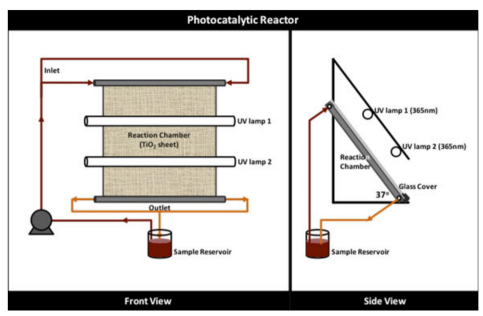


Figure 1. Schematic presentation of the photocatalytic reactor.

Eq. (1), including ANOVA to obtain the interaction between the process variables and the responses. Statistical significance was established at the P < 0.05 level.

RESULTS AND DISCUSSION

Anaerobic digestion step

Olive oil mill wastewater samples were collected from an up-flow anaerobic reactor operating with an organic loading rate of $1.0\pm0.1~\rm kg~\rm COD~m^{-3}~d^{-1}$ and a COD removal efficiency of $79\pm5\%$. TPh removal was around 40% ($\pm10\%$) and the dark color was not removed during the AD step. Therefore, a subsequent step for TPh and color removal is necessary. A constrain to the photocatalytic process is the high TSS concentration present in raw OMW. The anaerobic reactor was operating with removal efficiencies of $70\pm5\%$ of VSS and $40\pm5\%$ of TSS when the samples were collected. A large part of the OMW solids are being retained/degraded in the AD step representing a major advantage for the use of photocatalysis as a posttreatment. TSS at the beginning of the experiments were $215\pm20~\rm mg~L^{-1}$. Details about the anaerobic reactor operation and results are described elsewhere (17).

Photocatalysis

A statistical approach based on a factorial experimental design that would allow us to infer about the effect of four variables with a relatively small number of experiments was chosen. The classical univariate approach is suitable for factors that are independent but is time consuming because the response is investigated independently for each factor while all other factors are held at a constant level. Moreover, univariate methods do not take interactive effects between factors into account. The experimental matrix and the corresponding response variables obtained in each run are presented in Table 2.

Total phenols removal. Total phenols are one of the major problems in OMW treatment. Initial concentration of TPh varied between 129 and 400 mg L $^{-1}$, which stay far away from the legislation limits imposed by the Portuguese law (Table 1). TPh are the main compounds targeted by the photocatalysis technique, therefore its removal was the first response variable studied. Initial COD decrease from 7370 to 1051 mg L $^{-1}$ (and concomitant decrease in initial concentration of TPh) caused an increase in TPh removal efficiency from $33 \pm 7\%$ to $74 \pm 15\%$ (Table 2). Initial COD was identified as the most significant variable for the model (78.9% contribution). However, there are other significant variables, such as $t_{\text{treatment}}$ (13.2%), pH (2.8%) and an interaction between COD and $t_{\text{treatment}}$ (4.3%). This can be described by the following model equation:

$$Y_1 = 53.1 - 20.6Z_1 - 3.9Z_2 + 8.4Z_3 - 4.9Z_1Z_3 \tag{2}$$

where Y_I is the total phenols removal efficiency (%), Z_i is the transformed forms of the independent variables according to:

$$Z_i = \frac{\chi_i - \frac{\chi^+ + \chi^-}{2}}{\frac{\chi^+ + \chi^-}{2}} \tag{3}$$

where X_i is the original (untransformed) values of the variables, and X^+ and X^- are the upper and lower limit value of the variables.

Equation (2) and Table 2 show that at lower concentration of TPh the photodegradation efficiency is high, but it decreases as the concentration increases. When the pollutant concentration increases, the radicals OH* needed to achieve a similar degradation increases. However, the formation of hydroxyl radicals remained constant in time because the catalyst concentration and the light intensity did not change. A dependency between the photocatalysis efficiency and the initial concentration of phenol was also detected in (22). According to the authors, the competitive adsorption on the surface of TiO₂ particles between the hydroxyl radicals generated by the active site OH⁻ and phenol molecules adsorbed on the surface catalyst, which react with radicals, can affect the overall degradation rate. Slow diffusion of

Table 2. Design matrix of the 2^4 factorial experimental design and observed response factors as well as removal efficiencies of COD, TPh and color (at $\lambda = 390$ nm) and specific electric energy consumption per unit of olive oil mill wastewater mass ($E_{\rm EM}$).

Independent variables matrix					Removal efficiency				
Test	X_I COD mg L ⁻¹	<i>X</i> ₂ рН	X_3 $t_{ m contact}$ hour	X_4 $t_{ m cycle}$ minute	Y ₁ TPh %	Y_2 Color ($\lambda = 390$)	Y ₃ COD %	$E_{\rm EM}$ kWh kg $^{-1}$ COD _{removed}	
1	-1	+1	-1	+1	56.6	19.0	27.8	2735	
2	-1	+1	+1	+1	83.3	49.0	41.3	5536	
3	-1	+1	-1	-1	57.1	32.7	37.2	2049	
4	-1	+1	+1	-1	87.3	65.7	52.5	4352	
5	-1	-1	-1	+1	64.9	45.5	18.6	4092	
6	-1	-1	+1	+1	91.0	76.6	59.3	3855	
7	-1	-1	-1	-1	63.5	49.8	34.0	2238	
8	-1	-1	+1	-1	86.1	77.4	56.5	4044	
9	+1	+1	-1	+1	26.3	21.6	35.2	308	
10	+1	+1	+1	+1	29.9	23.5	39.5	823	
11	+1	+1	-1	-1	23.5	16.9	34.9	311	
12	+1	+1	+1	-1	29.9	18.0	35.8	910	
13	+1	-1	-1	+1	34.8	42.7	29.7	366	
14	+1	-1	+1	+1	43.9	46.2	33.4	975	
15	+1	-1	-1	-1	31.0	34.4	37.3	291	
16	+1	-1 -1	+1	-1 -1	40.9	36.4	42.5	767	
10	71	-1	r1	-1	40.9	50.4	42.3	707	

the intermediates from the catalyst surface can result in the block of active sites on the catalyst, resulting in a reduction of the degradation rate. When the initial concentration is high, the number of active sites available is decreased by the phenolic compounds, and because the intensity of irradiation light was constant, the OH $^{\bullet}$ and O $_2$ $^{-\bullet}$ radicals formed on the surface of the catalyst were also constant; then, the attack of radicals on the compound molecules decreased while, simultaneously, the photodegradation efficiency decreases. At low concentration, the number of catalytic sites will not be a limiting factor and the degradation rate will be proportional to the substrate concentration, according to the apparent first-order kinetics (23).

The only variable that has a positive effect on the TPh removal is the treatment time. Indeed, an increase in $t_{\rm treatment}$ from 8 to 24 h caused an increase in the removal efficiency from 45 \pm 17% to 62 \pm 28% (Table 2). However, a negative interaction effect between $t_{\rm treatment}$ and initial COD [Eq. (2)] was detected. Indeed, the increase in substrate concentration can lead to the generation of intermediates, which may adsorb on the surface of the catalyst and gradually block the catalyst action. Also, the accumulation of fouling may cause the decrease of incident light and decrease the phenolic compounds degradation and decolorization efficiency.

The pH has a key effect on the oxidation potential of OH. radicals because of the direct relation of the oxidation potential with the pH value (E0 = 2.8 V and E14 = 1.95 V) (24). In this study, pH was identified as a significant variable affecting negatively the TPh removal efficiency [Eq. (2)] and decolorization (see subsection Color removal), meaning that higher efficiencies will be obtained at lower pH values. TPh and Color removals efficiencies changed from 57 \pm 23% to 49 \pm 26% and from $51 \pm 17\%$ to $31 \pm 18\%$, when the initial pH was 4.8 and 7.5, respectively (Table 2). Because the color removal is caused by phenolic compounds removal, the observed pH dependency can be explained by the fact that at pH 7.5, the phenolic compounds (weak acids, with a pKa of 4.5) are fully dissociated (25), and the TiO2 surface is negatively charged (its point of zero charge "pzc" is 6.8) (26). Thus, at this pH, there are repulsive forces between the negatively charged catalyst surface and the phenolic

compounds, resulting in very low adsorption. At pH 4.5, where the ${\rm TiO_2}$ surface is positively charged and the phenolic compounds are protonated, higher adsorption capacity can be achieved (27).

Color removal. Another problematic issue for the OMW treatment is its color. After the anaerobic treatment, the effluent did not accomplish the discharge limits (Table 1), presenting a strong dark color. UV/TiO₂ treatment can oxidize the colored phenolic compounds triggering the color removal. According to the factorial analysis, the main contributions for the color removal efficiency were the initial COD (33.4% contribution), pH (28.5%) and $t_{\rm treatment}$ (18.2%). Significant interactions must be considered namely, between COD and $t_{\rm treatment}$ (13.8%), COD and $t_{\rm contact}$ (4.4%), pH and $t_{\rm contact}$ (1.2%). This can be described in the following model equation:

$$Y_2 = 41.0 - 11.0Z_1 - 10.2Z_2 + 8.1Z_3 - 7.1Z_1Z_3 + 4.0Z_1Z_4 - 2.1Z_2Z_4$$
(4)

Where Y_2 is the color removal efficiency.

Effluent color decreased during the photo oxidation (Table 2). The main influence for the effluent decolorization efficiency is the initial COD concentration (and concomitant TPh concentration) because the effluent decolorization is coupled with the TPh degradation. The color removal efficiency represent around 68% of the TPh removal, in the experiments with 129 mg TPh $\rm L^{-1}$, increasing to around 90% during the runs with 400 mg TPh $\rm L^{-1}$. As expected, the OMW dark color is mainly due to the presence of phenolic compounds and their polymerized derivatives.

Increasing in the $t_{\rm treament}$ positively influenced the color removal efficiency. However, this assumption must be carefully analyzed because of the interaction effect between $t_{\rm treatment}$ and COD that negatively influences the process efficiency [Eq. (4)]. As explained in subsection total phenols removal, the increase in initial COD concentration may cause the gradual inactivation of active sites and consequently decrease the photoreactor efficiency. Confirmation of the previous interaction effect was

Table 3. Predicted response variables by the factorial experimental design model.

Variable response	Prediction	SD	95% CI low	95% CI high	95% PI low	95% PI high	95% TI low	95% TI high
TPh Color	90.8 79.3	2.7 1.9	87.5 76.4	94.1 82.1	84.0	97.6 84.4	77.4 69.0	104.3 89.5
COOD	50.3	6.3	44.4	56.2	74.1 35.5	65.1	21.4	79.1

SD—The standard deviation estimated on the ANOVA. 95% confidence interval (CI)—This is the low/high value of the 95% CI in which you could expect to find the average response of a group of runs. 95% PI—This is the low value of the prediction interval (PI) that will contain the true value of an individual observation with 95% CI. 95% TI-This is the lower bound of the tolerance interval (TI) containing a given proportion of all future outcomes with a given confidence level.

obtained by the decrease of the degradation rate in the longer tests, especially in the assays with 7370 mg COD L^{-1} . However, increasing the initial COD caused the removal efficiency decrease. This was expected because there are more molecules to be converted with the same incident light. Also, accumulation of wastes/fouling, especially with higher organic loadings, deteriorates the paper, causing the decrease in active sites and incident light intensity. This may explain why the color removal occurs mainly in the first hours of treatment.

Another important interaction occurs between initial COD and $t_{\rm contact}$. It seems that to counterbalance the increase in COD concentration it is necessary to increase the contact time between the effluent and the catalyst. This may be achieved by decreasing the circulation flow rate. When we increase the circulation flow rate, the thickness of the liquid film increases and less photons reaches the photocatalyst surface. This results in slower production of hydroxyl radicals, and consequently, lower photocatalytic reaction rates.

COD removal. COD removal efficiency is almost unaffected by the pH and $t_{contact}$, however, the $t_{treatment}$ has a large positive influence, with 40.3% contribution to the model described by Eq. (5). The interaction effect of $t_{\text{treatment}}$ and initial COD has a negative effect over the COD removal efficiency with 27.2% contribution to the following model:

$$Y_3 = 38.8 + 6.3Z_3 - 5.2Z_1Z_3 \tag{5}$$

Where Y_3 is the COD removal efficiency.

COD removal efficiency is directly linked with the treatment time. However, at high initial COD concentrations, due to the high energetic costs associated with this treatment, it is economically unfeasible to extend the treatment time until the legislation limits are achieved.

A substantial part of the initial COD is due to other compounds besides the phenolic compounds, which represent only 5-12% of the initial COD. These compounds, originally present in OMW or formed as secondary reaction by-products during the AD or photocatalysis steps, are more resistant to photocatalysis causing the difference between the COD and TPh removals efficiencies.

Once more is visible the negative influence of the $t_{\text{treatment}}$ and COD interaction [Eq. (5)]. This was detected for all the response variables Y, and was due to the deactivation of active sites and formation of intermediates that inhibit the photoreactor efficiency. Chatzisymeon and co-workers (28) used a similar approach but in a batch reactor with Degussa P25 TiO2 and obtained comparable results, although no significant interactions were detected. This main difference may be caused by the reactor characteristics. In a batch reactor with disperse catalyst and

constant agitation the deactivation of active sites with time is not as significant as in an immobilized catalyst.

Table 3 shows the predicted values for the three response variables tested. Removal efficiencies of 90.8 \pm 2.7%, 79.3 \pm 1.9% and 50.3 \pm 6.3% for TPh, color and COD, respectively, can be obtained with the experimental set up tested. In run 6 (Table 2), the maximum degradation of TPh (91%) and COD (59%) were achieved, whereas the color removal was 77%. The difference between degradation rates of TPh, color and COD could be explained by two factors: (1) the formation of small colorless organic molecules, which, at least temporarily, remain in the solution, for instance ethanol, glycerol and simple sugars, which can be considered as degradation intermediates; (2) the persistence, for varying periods of time, of only slightly colored or colorless macromolecules relatively resistant to the UV/TiO₂ attack, and hence likely to be carbohydrates (29). This is in good agreement with the interpretation of the TiO2 attack to OMW, i.e. preferential attack of the electron-rich aromatic and olefinic groups, and attack at a lower rate of aliphatic side chains and carbohydrates (30,31).

A possibility to increase the process efficiency is to increase the incident light intensity. Laoufi and co-workers (22) tested different light power sources. Increasing the light intensity from 15 to 400 W enhanced phenol degradation from 15% to 100%, after 4 h of irradiation. According to the authors, the photons present at high intensity were in excessive amounts leading to more reactive species generation and more destruction of phenol.

Energy consumption. One disadvantage of UV-photocatalysis is the need of energy that represents a bottleneck to the scale-up of this technique. As stated in subsection COD removal, the photocatalysis efficiency is linked with the UV lamp power (22,32). The specific electric energy consumption per unit of pollutant mass ($E_{\rm EM}$) [kWh kg $^{-1}$ COD_{removed}] can be calculated from the formula (33):

$$E_{\rm EM} = \frac{1000Pt}{V(C_i - C_f)} \tag{6}$$

Where P is the rated power of the UV lamp [kW], V is the volume [L] of OMW treated in the time t [h], C_i and C_f are the initial and final COD concentration [g L⁻¹] and the factor 1000 converts g to kg.

The results show that this is an expensive technique, but the $E_{\rm EM}$ substantially decrease as the initial COD increases (Table 2). Similar results were obtained in (26), where photocatalysis was used for the OMW pretreatment. Currently, coupling a photocatalytic oxidation as a pretreatment aiming the effluent detoxification and a subsequent biological step is gaining relevance. However, although the OMW detoxification may be important, this approach has the weakness of not take full advantage of the real valorization potential of this effluent. By using appropriate measures, such as biological community adaptation and combining feeding phases with batch degradation (17), is possible to prevent the biological step inhibition and achieve the effluent valorization through the production of an energy biocarrier (biogas). Afterward, the photocatalysis step may be used to polish the final effluent quality, removing phenols and color. The integrated process can achieve the OMW valorization and overcome the economical issues associated with the photocatalytic step. Moreover, the inconvenience of effluent dilution and/or solids removal should be kept in mind because photocatalysis alone is not capable of treating effluents with high organic and solid loadings, which can be removed in the AD step.

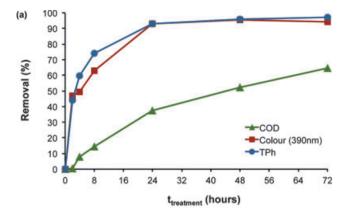
Treatment time effect. A second experiment, with initial conditions close to the optimal previously defined by the factorial analysis, was performed to evaluate the effect of extend the treatment time beyond 24 h. This parameter has been identified as having the highest positive effect in the models defined to maximize the removal efficiencies of phenols, color and COD. OMW initial COD was 1400 mg L⁻¹, TPh were 75 mg L⁻¹, t_{contact} was 2 min, and Color at 390 nm was 1.62 cm⁻¹. COD, TPh and color removal efficiencies were followed during 72 h (Fig. 2).

TPh and color removal were almost complete (>90%) after 24 h of operation. Fig. 3 shows the OMW samples at the beginning and after the treatment. The reactor efficiency to remove the OMW color is visible. However, the COD removal efficiency still increased from 38% (24 h) to 65% (72 h). If the fixed treatment time is too short, the intermediates remaining in solution could still be structurally similar to the initial biorecalcitrant compounds and therefore nonbiodegradable (24). However, the photocatalysis treatment time must be as short as possible to avoid high energy consumption, which represents about 60% of the total operational costs when using electric light sources (34). It was confirmed that a long treatment time is necessary to achieve an efficient COD removal. Therefore, a recirculation between the AD step and the photocatalytic reactor should be considered to degrade the residual COD.

CONCLUSIONS

A factorial experimental design was performed to optimize the performance of a photocatalytic reactor used for the posttreatment of olive mill wastewater. The Total phenols (TPh) removal efficiency was highly dependent of the initial COD concentration, being almost completely oxidized after 24 h. The color removal efficiency decreased with increasing COD concentration and pH, and the COD removal efficiency was directly linked with the treatment time. Regarding the interaction effects, it was identified a negative effect between the initial COD and treatment time. Increasing the treatment time of effluents with high COD may cause the inactivation of active sites of the TiO₂ paper, causing a drop in its efficiency. According to the factorial analysis, it is possible to obtain removal efficiencies of $90.8 \pm 2.7\%$, $79.3 \pm 1.9\%$ and $50.3 \pm 6.3\%$ for TPh, color and COD, respectively, in the best operational scenario tested, i.e. low levels of COD, pH and $t_{contact}$, and high $t_{treatment}$.

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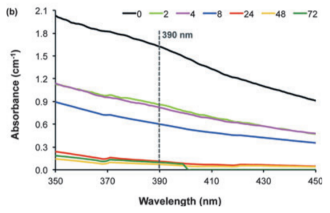


Figure 2. COD, TPh and Color (at 390 nm) removal efficiencies in the photocatalytic reactor (a). UV–Visible (350–450 nm) spectra of oil mill wastewater during the photocatalytic treatment (b).



Figure 3. Olive oil mill wastewater pictures at the beginning (a) and at the end (b) of the photocatalytic treatment.

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