Titanium dioxide immobilization in recycled aluminum net for degradation of textile dye by Heterogeneous Photocatalysis through neural networks

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Abstract

In this work, recycled aluminum nets were used as TiO$_2$ support for the treatment of the textile dye Remazol golden yellow (RGY 150). The immobilization of 100 mg of TiO$_2$ was accomplished after aluminum net calcination. The following experiments were carried out in a photocatalytic bench reactor: preliminary tests, a $2^3$ factorial experimental design with the best system and the degradation kinetic of TiO$_2$ supported systems. Characteristic peaks of aluminum and TiO$_2$ (anatase and rutile) were observed by XRD and the catalyst film formation was verified by SEM. The system UV/H$_2$O$_2$/TiO$_2$ performed higher efficiency, with 99% of dye degradation in 50 minutes, followed by the system UV/H$_2$O$_2$/TiO$_2$sp with 93% of dye degradation. Whereas the supported catalyst presents execution advantages and high degradation rates, the factorial design was realized for this system, achieving best degradation (100% degradation in 35 minutes). It was obtained k = 0.114 min$^{-1}$ and R$^2$ = 0.994 in a pseudo-first order model adjustment. The use of artificial neural network was proven to be efficient to predict the degradation of a textile dye with an absolute error of 0.0181. After treatment, there was a decrease of dye toxicity.

Keywords: Aluminum net. Advanced Oxidation Process. Remazol golden yellow. Photocatalysis. Phytotoxicity.

Resumo

Neste trabalho, redes de alumínio recicladas foram utilizadas como suporte de TiO$_2$ para o tratamento do corante têxtil Remazol amarelo ouro (RGY 150). A imobilização de 100 mg de TiO$_2$ foi realizada após calcinação do material de alumínio. Foram conduzidos em reator de bancada fotocatalítico testes preliminares, um experimento fatorial $2^3$ com o melhor sistema e a cinética de degradação em sistemas com TiO$_2$ suportado em rede de alumínio. Foram observados picos característicos de alumínio e TiO$_2$ (anatase e rutile) por DRX e a formação do filme catalítico foi verificada por MEV. O sistema UV/H$_2$O$_2$/TiO$_2$ apresentou maior eficiência, com 99% de degradação do corante em 50 minutos, seguido do sistema UV/H$_2$O$_2$/TiO$_2$sp com 93% de degradação do corante. Enquanto o catalisador suportado apresenta vantagens de execução e altas taxas de degradação, o planejamento fatorial foi realizado para este sistema, apresentando melhor degradação (100% de degradação em 35 minutos). Foi obtido k = 0,114 min$^{-1}$ e R$^2$ = 0,994 em um sistema de ajuste de modelos de pseudo-primeira ordem. O uso da rede neural artificial mostrou-se eficiente para predizer a degradação de um corante têxtil, com erro absoluto de 0,0181. Após o tratamento, houve uma diminuição da toxicidade do corante.


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1 Introduction

The textile industry is one of the major producers of waste water containing dyes and chemical products (ASGHAR; RAMAN; DAUD, 2015; HASSAAN; NEMR, 2017). The environmental impact by the textile sector provides a representative pollutant potential, once the industrial wastes are very diverse and their chemical composition vary according to the industry typology and the involved processes. This effluent class is considered toxic and contains non-biodegradable compounds and the dyes derived from industrial processes are largely responsible for this characteristic (ALISANFI et al., 2007). Therefore, it is necessary to develop alternative methods for waste treatment that ensure a contaminant level decrease after treatment.

Moreover, through these processes, the organic molecules complete or partial oxidation can happen from the formation of very reactive free radicals species, such as hydroxyl radicals (OH•), promoted by a radiation source (M-BRA et al., 2019; LI et al., 2017; POURAKBAK et al., 2016). They are mainly characterized by the non-selectivity and the use of potent oxidant species, as hydrogen peroxide (H₂O₂), or semiconductors, as titanium dioxide (TiO₂), in the radical generation (ROZAS et al., 2016; MOREIRA et al., 2017).

In the heterogeneous photocatalysis, the TiO₂ has been used in aqueous medium dispersed or immobilized on some support surface. Its use in dispersion arrangement in aqueous medium is energetically more efficient because of the higher contact area available for the catalyst. However, the immobilized form presents the advantage that it is not necessary to separate dispersed TiO₂ particles in the end of treatment (BARROS et al., 2014).

In order to avoid this disadvantage, recent studies utilized diverse materials for TiO₂ immobilization, such as glass (CUNHA et al., 2018; OBLAK et al., 2018), polymeric membranes (BARROS et al., 2014; MUKHERJEE et al., 2014), concrete (DELNAVAZ et al., 2015) and metallic supports (MOZIA et al., 2012). Those materials must present rugosity for catalyst particles fixation (BARROS et al., 2016) and good adherence to prevent leaching after immobilization (CUNHA et al., 2018).

During the studies of advanced oxidation processes is common the lack of knowledge about every existent reaction in the degradation. Many times the degradation kinetic is approximated by a pseudo-first order model, as observed in several works (AJMAL et al., 2016; KUMAR et al., 2018; SOLEYMANI; CHAHARDOLI; KAYKHAIL, 2016). Nonetheless this approximation does not consider the actual degradation process complexity. One modeling technique utilized when knowledge is not available about a complex process is the Artificial Neural Network (ANN) (ROCHA et al., 2017; DAS, MAITY AND BASU, 2014). The interesting feature in neural networks is the ability to learn from experimental data exposed to them, and thus to generate a model that can approximate any curve (CSÁJI, 2001).

The ANN are able to fit in a phenomenon behavior when submitted to the experimental conditions of the process to be modeled, without the phenomenon knowledge involved in the process. The ANN has been applied to solve hard problems that would not be solved by other methods. The neural networks can model one- and multidimensional cases, what make them a practical tool to analyze kinetic patterns of complex reactions, as in this work, as well as in other academic works, for example Rocha et al. (2017) and Das, Maity and Basu (2014).

This work objective was to evaluate the efficiency of aluminum soda can seals (post-consumer recycled material) as support materials (sp) for TiO₂ films in the degradation through heterogeneous photocatalysis of the dye Remazol golden yellow RNL 150 (RGY 150), a synthetic dye commonly used by the textile industry. The usage of aluminum seals as TiO₂ support presents economic and environmental perspectives, once the material is abundant and has low cost and aimed mechanical properties. The degradation evaluation was realized by kinetic modeling for different reaction orders and by Artificial Neural Networks.

2 Materials and Methods

In the degradation assays, the textile dye Remazol golden yellow RNL 150 (RGY 150), obtained from F. Trajano Aromas e Ingredientes LTDA was used. The titanium dioxide was obtained from Evonik Degussa LTDA, comprising 75% of anatase and 25% of rutile, with superficial area of 50 m².g⁻¹ and particle size of 20 nm. The hydrogen peroxide 50% (v.v⁻¹) was obtained from Coremal Pochteca°.

2.1 TiO₂ calcination and immobilization in aluminum nets

Supports crafted from aluminum seals were prepared from post-consumer soda cans. The can seals were cleaned and washed with distilled water. The nets were acidified with HCl 37% for 20 seconds, succeeding to calcination at 490°C in muffle (SPLABOR, SP-1200DRP7) for 26 hours before TiO₂ immobilization.

In the next, TiO₂ immobilization on the calcined aluminum nets was carried out by the methodology described by Barros et al. (2014). The total immobilized quantity was 100 mg of catalyst in each support. After TiO₂ immobilization, the samples were taken to muffle at 490°C for 4 hours for best catalyst adherence on support.

2.2 Characterization of aluminum supports

The samples were analyzed in an XRD-6000 diffractometer from SHIMADZU, with radiation Cu Ka (λ = 1.5418 Å) using Nickel filter as a monochromator, power of 2 kVA, voltage of 30 kV and electric current of 30 mA. The scanning analyses were realized in the 20 range from 20° to 80°, step of 0.02° and velocity of 2°.s⁻¹. The characteristic peaks of the materials were compared with the crystallographic charts 21-1272 and 21-1276 (International Center for Diffraction Data) and the aluminum...
rings crystallography prior to the TiO\textsubscript{2} immobilization. The morphological evaluation of the aluminium support nets before and after acidification and calcination, and after TiO\textsubscript{2} immobilization was performed by Scanning Electron Microscopy (SEM, Tescan MIRA3).

### 2.3 Preliminary tests

Before each experiment in which TiO\textsubscript{2} was involved, the solution was kept in catalyst contact without radiation, under stirring, in order to reach the adsorption equilibrium. The systems UV, UV/TiO\textsubscript{2}, UV/H\textsubscript{2}O\textsubscript{2}, UV/H\textsubscript{2}O\textsubscript{2}/TiO\textsubscript{2}, UV/TiO\textsubscript{2}sp, UV/H\textsubscript{2}O\textsubscript{2}/TiO\textsubscript{2}sp, where sp means supported catalyst systems were evaluated in the degradation processes. The amount of TiO\textsubscript{2} and H\textsubscript{2}O\textsubscript{2} utilized was 100 mg and 1.59 mmol.L\textsuperscript{-1}, respectively. The TiO\textsubscript{2} mass used was based on the methodology described by Nascimento Júnior et al. (2018) and the H\textsubscript{2}O\textsubscript{2} concentration in stoichiometric calculus.

The tests were carried out in a photocatalytic batch reactor, that contains a UV lamp (245 nm) with power of 0.830 mW.cm\textsuperscript{-2}. The lamps were turned on 30 minutes before the experiments beginning for radiation stabilization. In cylindrical glass containers (height of 8.5 cm and diameter of 9,0 cm), 300 mL of the dye RGY 150 in concentration of 20 mg.L\textsuperscript{-1} were treated for 50 minutes.

### 2.4 Factorial Experimental Design (2\textsuperscript{3})

A 2\textsuperscript{3} factorial experimental design was conducted for the best system obtained in the preliminary batch tests. The studied parameters were H\textsubscript{2}O\textsubscript{2} concentration, TiO\textsubscript{2} mass and degradation time. The 2\textsuperscript{3} factorial design center point was done in triplicate as a means to determine the error, totaling 11 experiments. The studied levels are described in Table 1.

| Table 1 - Variables levels of the 2\textsuperscript{3} factorial design |
|-------------------|-----------------|------------------|
| Levels | TiO\textsubscript{2} (mg) | H\textsubscript{2}O\textsubscript{2} (mmol.L\textsuperscript{-1}) | Time (min) |
| -1    | 100             | 1.59             | 15           |
| 0     | 150             | 5.21             | 25           |
| +1    | 200             | 8.04             | 35           |

The Pareto diagram and the response surface methodology were afterwards to evaluate the statistical significance of the variables, based on the work of Barros Neto et al. (2010), through the program Statistic 6.0 (SANTOS et al., 2018).

### 2.5 Kinetic Study

A kinetic study was performed with the best conditions found in the factorial experimental design. The degradation kinetic was adjusted to the pseudo-first order, pseudo-second order and pseudo-third order models (Equations 1 to 4), by the software Excel 2013\textsuperscript{®}. In the equations, C is the concentration for time t and C\textsubscript{0} is the initial concentration. For the kinetic study, the correlation coefficient (R\textsuperscript{2}) for each studied order was determined and the velocity constant (k\textsubscript{app}) and the time of half-life (t\textsubscript{1/2}) of the photocatalytic reaction were obtained. The time of half-life was found by the Equation 6.

\[
\frac{dC}{dt} = -kC \quad \rightarrow C = C_0 e^{-kt} \quad (1)
\]

\[
\frac{dC}{dt} = -kC^n \quad \rightarrow C = \frac{1}{t} \left(2\sqrt{(n-1)} \frac{kt}{C_0} + C_0\right) \quad (2)
\]

\[
n = 2, \quad \frac{C(t=0)}{C_0} = 1 \quad \rightarrow C = \frac{1}{kt + 1} \quad (3)
\]

\[
n = 3, \quad \frac{C(t=0)}{C_0} = 1 \quad \rightarrow C = \frac{1}{\sqrt{2kt + 1}} \quad (4)
\]

\[
n = 1, \quad \ln\left(\frac{[C]}{[C_0]}\right) = -kt \quad (5)
\]

\[
t_{1/2} = \frac{\ln 2}{k} \quad (6)
\]

The kinetic model using ANN was also established for the best factorial design conditions. The ANN was constructed using time as the input value and dimensionless concentration as the output, generating a curve of concentration depending on time, according to other authors such as Göb et al. (1999), Giroto et al. (2006), Rocha et al. (2017) and Das, Maity and Basu (2014). The software utilized for the neural adjustment was developed by the research group through Unity 3D\textsuperscript{®}. The ANN is composed by input, hidden layers and output with different structure and number of neurons. The chosen net structure was 1:2:1 (one input variable, time, one hidden layer with two neurons and one output variable).

The achieved software was able to adjust this neural network to real data in the learning stage. The learning method was based on the particles swarm optimization, random search and genetic algorithm, partial and initially presented by Kennedy and Eberhart (1995). The error criteria applied was the mean absolute error for the studied system. A feedforward neural network was used and the activation function adopted was sigmoid between 1 and -1. To realize the ANN analysis, the input and output values were normalized and stablished between -1 and 1 (t = 0 and t = 35 min). Furthermore, the output concentrations (C/C\textsubscript{0} = 0 e C/C\textsubscript{0} = 1) were renormalized, lest the training makes the network goes away to a saturation zone, as exposed by Haikin (1999). The training time utilized was 600 seconds to reach a limit residual error.

### 2.6 Toxicity assays

The toxicity assay consisted to expose lettuce seeds of the species Lactuca Sativa (obtained from Feltrin Semen-
tes, Brasil) to dye samples and the best treatment system observed in the studies, following Meyer et al. (1982). The negative control was done with distilled water and the positive with boric acid solution (3%). The assay was carried out five times. The seeds were exposed to the dye and its treatment for 120 hours in the dark and temperature at 22 °C. The radicles growth, the inhibition rate and the inhibition concentration of the radicle growth for 50% of organisms (IC_{50}) were calculated.

### 3 Results and Discussion

#### 3.1 Characterization of the aluminum support

The XRD spectrums of TiO$_2$ (100 mg) immobilized in aluminum support are showed in Figure 1. The qualitative interpretation of the compounds present in the catalyst was realized based on the Inorganic Crystal Structure Database (ICSD): 21-1272 e 21-1276.

Figure 1 - XRD diffractogram of TiO$_2$ immobilized in aluminum nets (anatase (A), rutile (R) e aluminum (Al))

Figure 2 - SEM analyses for a) aluminum support, b) acidified and calcined aluminum support and c) TiO$_2$ supported in aluminum net.
It is observed in Figure 1 the presence of the TiO$_2$ anatase and rutile phases, as well as the metallic substrate of aluminum (nets). The peaks 25.53° (101), 38.60° (112), 43.22° (200) and 62.6° (204) are referred to the anatase phase (JCPDS 00-021-1272) for different crystal plans. The rutile phase (JCPDS 00-021-1276) is represented by the peak 44.77° (210). The peak in 77.24° is probably related to the metallic substrate of aluminum.

The SEM scans displayed in Figure 2 demonstrate the aluminum support surface before calcination, after calcination and after calcination and immobilization of 100 mg of TiO$_2$.

In Figure 2a), it is observed that the metallic material without acidification and calcination does not present porosity, while in Figure 2b) the aluminum support is presented porous and irregular after treatments. The immobilized TiO$_2$ layer is displayed as a uniform film (Figure 2c) after the TiO$_2$ distribution on the aluminum net and succeeding thermal fixation. According to Ramasundaram et al. (2015), in whose work steel wires were covered with TiO$_2$, the material porosity after thermal treatment allowed the formation of a catalyst layer, as well as in this work. Oblak et al. (2018) used stainless steel nets for the TiO$_2$ immobilization and the uniform distribution of material was noticed likewise.

3.2 Preliminary tests

In the preliminary tests, without radiation, it was observed that there was not degradation of the dye solution after 50 minutes in all studied conditions. This behavior was expected, since without radiation source, it is not possible to generate the free radicals (OH•) responsible for the organic dyes degradation. The same behavior was observed by Santos et al. (2018) and Nascimento Júnior et al. (2018) in their photocatalytic studies with dyes. In Figure 3, the degradation behavior of the RGY dye is represented for different AOP as a function of the experiment time.

The Figure 3 points out that the heterogeneous photocatalysis systems (UV/H$_2$O$_2$/TiO$_2$ and UV/H$_2$O$_2$/TiO$_2$sp) presented the best degradation rates among the tested systems, with efficiency of 99.93% and 93.41%, respectively, for 50 minutes. The systems with combined TiO$_2$ and H$_2$O$_2$ demonstrated the best results, what can be explained due to the synergic phenomenon that causes an increase in the global rate of radical hydroxyl generation (SALLEH et al., 2016).

The efficiency reduction for the immobilized TiO$_2$ system was expected related to the suspended system, once the use of TiO$_2$ dispersed in aqueous solution is energetically more efficient as a result of the larger catalyst contact area (BARROS et al., 2014; ROSTAMI-VARTOONI et al., 2016). The same behavior was noted by Alalm et al. (2016). Although the suspended system has presented best results related to degradation, the difference was only 6.52%. In addition, the immobilized catalyst choice offers advantages, for instance, it facilitates its recovery in the end of the process and allows good catalyst handling (LUDWICK et al., 2015; ERJAVEC et al., 2016; BARBOSA et al., 2019).

Figure 3 - Preliminary degradation tests for the dye RGY 150 in aqueous solution by oxidation processes under radiation UVC. [C$_0$] = 20 mg L$^{-1}$ (1:1); [H$_2$O$_2$] = 1.59 mmol.L$^{-1}$; m$_{TiO_2}$ = 100 mg
3.3 Factorial Experimental Design (2³)

A 2³ factorial experimental design with center point was conducted for the system selected in the preliminary tests (UV/H₂O₂/TiO₂sp). The utilized parameters were H₂O₂ concentration, TiO₂ mass and degradation time. The conditions applied to the design and the dye removal percentage are displayed in Table 2.

Table 2 – RGY degradation for levels of a 2³ Factorial Experimental Design

<table>
<thead>
<tr>
<th>Assay</th>
<th>H₂O₂</th>
<th>TiO₂</th>
<th>Time</th>
<th>Degradation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>27.13</td>
</tr>
<tr>
<td>2</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>29.73</td>
</tr>
<tr>
<td>3</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>35.25</td>
</tr>
<tr>
<td>4</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>59.93</td>
</tr>
<tr>
<td>5</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>71.50</td>
</tr>
<tr>
<td>6</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>93.57</td>
</tr>
<tr>
<td>7</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>76.80</td>
</tr>
<tr>
<td>8</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>100.00</td>
</tr>
<tr>
<td>9</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>70.06</td>
</tr>
<tr>
<td>10</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>74.15</td>
</tr>
<tr>
<td>11</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>72.23</td>
</tr>
</tbody>
</table>

The assay 8, that comprises the maximum values of every parameter in Table 2, presented the best degradation rates of RGY 150. Oancea and Meltzer (2013) perceived that the influence of the H₂O₂ amount applied for synthetic dyes removal obtained best degradation percentages for higher oxidant dosages. Also Pajootan et al. (2017), Santos et al. (2018) and Gupta et al. (2011) observed in their works that the TiO₂ increase caused higher efficiency on synthetic dyes removal in aqueous solution.

A statistical analysis of the variables (parameters) significance was performed by the software Statistica Experimental Design 6.0, as displayed on the Pareto chart in Figure 4.

Figure 4 - Pareto chart for the results of the 2³ factorial experimental design for the system UV/H₂O₂/TiO₂sp

The interactions H₂O₂ vs TiO₂ and TiO₂ vs Time presented statistical significance. From the Pareto chart result, response surfaces were plotted and showed in Figure 5.

The analysis of the response surface in Figure 5b) points out that while the TiO₂ concentration and reaction time rise, the dye degradation percentage also increases. Overall, the photocatalytic processes efficiency is increased with larger amounts of titanium dioxide, since there are not excessive quantities of reagent. Its excess can cause solution opacity, which hampers the radiation absorption by the system (Alalm et al., 2016; Reza et al., 2017). The effect of combined TiO₂ and H₂O₂ cooperates for a higher formation of hydroxyl radicals and collaborates to avoid the recombination electron/hole (Salah et al., 2016). Hence, their amounts increase tend to raise the catalytic process efficiency.

According to experimental data, an empiric model was obtained as exposed by Equation 7 for the system UV/H₂O₂/TiO₂sp by the software Statistica 6.0. This model...
Figure 5 - Response surfaces of the dye degradation according to the variables: a) TiO$_2$ x H$_2$O$_2$ and b) TiO$_2$ x Time for the system UV/H$_2$O$_2$/TiO$_{2sp}$

Figure 6 - Residual analysis of the empiric model for the dye solution degradation (UV-C/H$_2$O$_2$/TiO$_{2sp}$ system)
describes the dye degradation efficiency (%), throughout the study range, for the significant variables.

\[ D(\%) = 64.67 + 5.50X_{TiO_2} + 7.95X_{H_2O_2} + 21.39X_{time} + 2.06X_{TiO_2}X_{H_2O_2} - 1.07X_{TiO_2}X_{time} \]  

(7)

The residual analysis by software Statistica 6.0 for the experimental data of the system UV-C/H_2O_2/TiO_2sp is presented in Figure 6.

Residual analysis accessed by the 2^3 factorial experimental design presented a good adjustment, with the experimental values significantly close to the predicted ones. The system pure error was 0.26, which confirms that the observed data were adjusted adequately.

### 3.4 Kinetic Study

The degradation of RGY 150 with TiO_2 supported in aluminum (UV/H_2O_2/TiO_2sp) was studied in the best conditions of the factorial design. Figure 7 shows the reaction kinetic for this system.

From Figure 7, it is possible to observe that more than 95% of the color was removed until 20 minutes of treatment. At 30 minutes, 98% of the dye was degraded. The index of 100% of degradation was reached in the end of 35 minutes of treatment.

Three kinetic models (pseudo-first order, pseudo-second order and pseudo-third order) were tested in order to obtain a general trend model. The pseudo-first order model reached 0.114 min^-1 for the kinetic velocity constant, k, and 0.994 for the correlation coefficient, R^2. The time of half-life estimated for the system was 7.93 minutes.

However, as seen in Figure 7 for the pseudo-second and pseudo-third order models, adjustments of superior order do not fit in experimental data as satisfactory as pseudo-first order curve. This can be proved by the parameters values obtained in the adjustments: in the pseudo-second order model, the R^2 was 0.8438 and in the pseudo-third order model, it was 0.7534, both lesser values that the one reached in Figure 6 by the pseudo-first order model. The superior order unfitness indicates that the pseudo-first order is the best adjustment for the tested set of models, what is also perceived in other works of AOP application for dye degradation (AJMAL et al., 2016; KUMAR et al., 2018; SOLEYMANI; CHAHARDOLI; KAYKHAI, 2016).

The neural network resultant of the training with the degradation experimental data and its adjustment to them are displayed in Figure 8.

In Figure 8a), the input layer is related to normalized time values that generates concentration values, also normalized, as a result. During the calculus, the network multiply the input value times the weights (values above the lines that connect the neurons) and sum up to the bias (values above neurons). The value is then computed by the activation sigmoid function to get a result that is newly multiplied by a weight and succeeds to the next layer. Consequently the result is generated. This model is able to predict the dye degradation development in the given conditions.

The concordance between experimental and ANN predicted data is noted in Figure 8b). The mean absolute error was 0.0181, leading to a determination coefficient R^2 of 0.9967. This value is higher than the one observed in the pseudo-first order adjustment, what demonstrates that the neural network model can better describe the degradation behavior.
3.5 Toxicity assays

From the phytotoxicity bioassay, it was calculated the inhibition concentration for 50% of organisms exposed to the dye solution and the conditions with best results of the degradation studies. The radicle extent growth of *Lactuca sativa* in the dye solution and in its respective treatments was compared to the positive and negative control. The parameters germination, radicle length and IC$_{50}$ are displayed in Table 3.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Germination (%)</th>
<th>Radicle length (cm)</th>
<th>IC$_{50}$ (mg.L$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without treatment*a</td>
<td>93</td>
<td>2.1 ± 0.1</td>
<td>17.8 ± 0.2</td>
</tr>
<tr>
<td>Post-treatment</td>
<td>98</td>
<td>2.7 ± 0.2</td>
<td>22.5 ± 0.1</td>
</tr>
<tr>
<td>Positive Control*b</td>
<td>100</td>
<td>3.6 ± 0.2</td>
<td>-</td>
</tr>
<tr>
<td>Negative Control*c</td>
<td>0</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

*a*RGY 150 Solution 20 mg.L$^{-1}$  
*b*Distilled water  
*c*Boric acid solution 3%

It was observed, by Table 3, that the treatments applied to the dye solution decreased the acute phytotoxicity, once the IC$_{50}$ is inversely proportional to the toxicity. The negative control did not present germination and there was a growth of 0.6 cm for the sample after photocatalytic treatment comparing to the not treated sample. In the work of Santos et al. (2018), that studied the removal of the synthetic dyes bright blue and erythrosine (60 mg.L$^{-1}$) with titanium dioxide, a length variation of 0.43 cm between the treated and not treated samples was observed.

4 Conclusion

In the preliminary tests, a best behavior was observed for systems with combined TiO$_2$ and H$_2$O$_2$. In comparison with the suspended TiO$_2$ system, the catalyst supported on aluminum nets presents close dye degradation rates and does not need recovery procedures afterwards, what is an operational advantage. In the 2$^3$ factorial design for the supported catalyst system, the parameters amounts increase made the system more efficient, with good statistical significance and 100% of degradation in 35 minutes. With the best conditions ($m_{TiO2} = 200$ mg, [H$_2$O$_2$] = 8.04 mmol.L$^{-1}$, 35 minutes), a kinetic study was carried out and the system was better adjusted to the pseudo-first order model. The following kinetic coefficients were obtained: $k = 0.114$ min$^{-1}$, $R^2 = 0.994$ e $t_{1/2} = 7.93$ minutes. The neural network model presented still better results with the determination coefficient of $R^2 = 0.9967$. The degradation also diminishes the toxicity, what was proved by the phytotoxicity bioassay with lettuce seeds. The immobilized TiO$_2$ use is positive because is an ecological alternative due to the minimization of TiO$_2$ removal cost and the utilization of recycled material.
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Titanium dioxide immobilization in recycled aluminum-based zeolite catalysts: mechanisms, pathways and treatment


