Research Article

Photocatalytic Degradation of a Water Soluble Herbicide by Pure and Noble Metal Deposited TiO$_2$ Nanocrystalline Films

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We present the photocatalytic degradation of a water soluble sulfonylurea herbicide: azimsulfuron in the presence of titania nanocrystalline films. Efficient photodegradation of herbicide was achieved by using low-intensity black light tubes emitting in the Near-UV. The degradation of the herbicide follows first-order kinetics according to the Langmuir-Hinshelwood model. Intermediate products were identified by the LC-MS-MS technique during photocatalytic degradation. In order to increase photodegradation rate of the herbicide, we examined the effect of titania modification by depositing noble metals at various quantities and valence states. The presence of platinum at neutral valence state and optimum concentration induced higher photodegradation rates while silver-modified titania exhibited similar photocatalytic rates with those obtained with pure nanocrystalline TiO$_2$ films. Finally, the effect of initial pH value was also examined. Acidic or alkaline media were unfavorable for azimsulfuron photodegradation.

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1. INTRODUCTION

Photodegradation of various organic pollutants by photocatalysis, using wide bandgap semiconductors, has been extensively studied [1–3]. Among them, TiO$_2$ a relatively inexpensive semiconductor exhibits high photocatalytic activity, stability in aqueous solution, nontoxicity [4, 5], and so forth. However, TiO$_2$ usage has a few drawbacks; for example, it absorbs only in the UVA part of the light spectrum where solar radiation is only 2-3% of the total reaching the surface of the Earth [6]. Moreover, the application of TiO$_2$ for photocatalytic oxidation of organic molecules is limited by high charge carrier recombination rates that results in low quantum efficiency. In recent years, surface metallization of TiO$_2$ has received considerable attention as an option to overcome the high degree of charge carrier recombination [7–10]. Platinum, and some other noble metals, may be used for this purpose thus providing an electron sink. In addition, they may extend TiO$_2$ absorbance in the Visible [11]. The presence of a metal at the surface of TiO$_2$ results in the formation of a Schottky barrier at the metal-semiconductor interface, which facilitates the interfacial electron transfer and subsequently encourages the charge carrier separation [12].

Among the various organic substances, which are known as water pollutants, herbicides are a major pollution source for both underground and surface waters. Advanced oxidation processes are used, among others, also for the degradation of herbicides [13, 14]. Azimsulfuron (AZS, see Scheme 1 for chemical structure) belongs to the class of sulfonylurea herbicides, which have a broad spectrum of weed control, low application rate, and low animal toxicity [15, 16]. Sulfonylurea herbicides, in addition to playing an important role in modern agriculture, are also degradable by heterogeneous photocatalysis, as it has been proven in the past [17, 18]. In the present work, sol-gel prepared TiO$_2$ films, which were further modified with noble metal ions, were examined for the photodegradation of AZS in water. The effect of various parameters, such as the amount of metal deposits and pH value of herbicide aqueous solution, were studied in order to evaluate the optimum conditions for the photocatalytic oxidation of AZS.
Scheme 1: Chemical structure of AZS.

2. EXPERIMENTAL

Azimsulfuron, N-[[4,6-dimethoxy pyrimidin-2-yl]-amino] carbonyl]-1-methyl-4-(2-methyl-2H-tetrazole-5-yl)-1H-pyrazole-5-sulfonamide (AZS, 99.3% purity, Scheme 1), was donated by DuPont de Nemours, Puteaux, France. All the rest of reagents were obtained from Sigma-Aldrich, Mo, USA. Millipore water was used in all experiments.

2.1. Description of the photocatalytic reactor

The cylindrical reactor schematically shown in Figure 1 was used in all experiments [19]. Air was pumped through the gas inlet using a small pump to ensure continuous oxygen supply to the reaction solution while simultaneously agitating it. In cases where experiments were carried out in the absence of oxygen, the solution was deoxygenated by nitrogen flow and the openings were sealed. Four black light fluorescent tubes of 4 W nominal power were placed around the reactor. The whole construction was covered with a cylindrical aluminum reflector. Cooling was achieved by air flow from below the reactor using a ventilator. The catalyst was in the form of four-glass rings, covered on both sides with nanocrystalline TiO₂ film. Film deposition is described below. The glass rings were of 38 mm of diameter and 15 mm height, stacked and coaxially placed inside the reactor. Thus, the total surface of the photocatalyst film was approximately $2 \times 71.6 = 143 \text{ cm}^2$. The intensity of radiation reaching the surface of the film on the side facing lamps was measured with an Oriel radiant power meter and found equal to 0.79 mW/cm².

2.2. Nanocrystalline Titania films and metal deposition

Titania films were deposited by following the previously reported procedure [20, 21]. Briefly, for 25 mL solution, 3.6 g of the nonionic surfactant Triton X-100 (polyoxyethylene-10-isooctylphenyl ether) was mixed with 20 mL of ethanol, followed by addition of 1.6 mL of glacial acetic acid and 1.8 mL of titanium isopropoxide under vigorous stirring. Self organization of the surfactant in this original sol creates organized assemblies that act as templates defining nanoparticle size. The surfactant is burned out during calcination. After a few minutes stirring, the glass rings described above, which were previously thoroughly washed, sonicated in ethanol and dried in a N₂ stream, were dipped in the above sol and withdrawn slowly by hand. After the film was dried in air for a few minutes, it was calcined in an oven. The temperature was increased in a ramp rate of 20°C/min up to 550°C and left at that temperature for about 10 minutes. When the titania-covered rings were taken out of the oven, they were transparent and optically uniform. The above procedure was repeated several times in order to reach the quantity of catalyst necessary for the purposes of the present work. The final mass of titania on the four glass rings was 80 mg (20 mg on each glass ring). Noble metal ions were deposited on titania films by adsorption from aqueous solutions containing one of the following metal salts: Na₂PtCl₄·xH₂O or AgNO₃ at various concentrations (from $10^{-4}$ to $10^{-3}$ M for the platinum salt and from $5 \times 10^{-4}$ to $10^{-2}$ M for the silver salt). After the last layer of TiO₂ was deposited and immediately after the film was taken out from the oven, the rings were submerged in the salt aqueous solution and were left for half an hour in the dark. Then, the rings were washed, dried, and subjected to UV radiation for 30 minutes; or they were additionally heated at 500°C for 15 minutes. UV and heat treatment were performed to reduce cationic species to neutral metallic particles.

2.3. Photodegradation procedure

The reactor was filled with 100 mL water in which 6 mg of AZS were dissolved. Standard solutions were employed in all experiments. AZS is not adsorbed on titania films, however, about an hour of equilibrium was allowed before illumination. The concentration of AZS in solution was monitored by absorption spectrophotometry at 240 nm using quartz cuvettes. In order to assure linear relationship between absorbance and concentration, preliminary measurements have been made. The linear relationship was established in the range of $0 \rightarrow 60 \text{ mg L}^{-1}$. Thus, the presently chosen
concentration 60 mg L\(^{-1}\) was the maximum concentration of the established linear concentration versus absorbance range. It was found that this concentration is also close to the solubility limit of AZS in water. A 3 mL aliquot was collected for performing absorption spectrophotometry, and it was poured back into the reactor after measurement. The first concentration measurement was done in the dark. It was then verified that the herbicide was not adsorbed on the titania film. Finally, the products of photodegradation have been monitored by LC-MS-MS analysis.

2.4. Apparatus and methods

Absorption measurements were made with a Cary 1E spectrophotometer. LC-MS-MS analysis was carried out on a Varian 1200L mass spectrometer equipped with electrospray ionization interface (ESI). The ion trap was connected to a Varian 1200 LC system consisting of two individual pumps, a membrane degasser, an autosampler with temperature control, and a thermostatic column control. The chromatographic separation was performed on a reversed phase Varian Polaris C-18 column (150 mm × 2.1 mm, 5 μm particle size) at 30°C. The concentration of AZS at each degradation stage was calculated by using an external calibration curve from LC-MS-MS measurements of AZS at five different concentrations. Stock solution was prepared by diluting the appropriate amount of the analytical standard with water/methanol (70/30) at concentration 60 ppm. In order to determine the degradation products formed during AZS photocatalysis, ESI ionization in both negative- and positive-ion modes was investigated. The results indicated that the ESI source at positive-ion mode with collision energy at 30 V was most appropriate for the analysis of the degradation products.

3. RESULTS AND DISCUSSION

The films made by the sol-gel procedure described in the Experimental section consisted of anatase nanocrystals of about 15 nm average size. Size polydispersity is limited and the active surface area is around 110 m\(^2\)/g, as measured by nitrogen adsorption-desorption isotherms according to the BET analysis [21].

3.1. Photodegradation studies of AZS with pure TiO\(_2\) films

It has been proposed that titanium dioxide mediated photodegradation involves the generation of electron-hole pairs [2], which migrate to the photocatalyst surface forming surface bound hydroxyl and superoxide radicals according to following reactions:

\[
\begin{align*}
    \text{TiO}_2 + \text{hv} & \rightarrow e_{cb}^+ + h_{vb}^+; \\
    \text{O}_2 + e_{cb}^+ & \rightarrow \text{O}_2^-; \\
    \text{H}_2\text{O} + h_{vb}^+ & \rightarrow \cdot \text{OH} + \text{H}^+.
\end{align*}
\]

(1)

It is commonly known that the hydroxyl and superoxide radicals are the primary oxidizing species in the photodegradation process. These oxidative reactions result in the photodegradation of AZS. The efficiency of this process depends on oxygen concentration. The absorption spectra of the AZS solution at various stages of photodegradation are presented in Figure 2, where it is obvious that the concentration of intact AZS decreased during illumination in the presence of titania. The characteristic peak at 240 nm completely disappears after 15 hours of illumination, using low-power black light source. Practically, no photolysis of AZS has been observed at the present illumination level, as seen by curve number 2 of Figure 3. The presence of oxygen during photocatalytic experiments is absolutely necessary and this is proved by the data of curve number 3 in Figure 3. Absence of oxygen leads to negligible photodegradation. Photodegradation followed Langmuir-Hinshelwood kinetics:

\[
\ln \left( \frac{A_0}{A} \right) = k_{\text{app}} t.
\]

(2)
The corresponding first-order rate constant was 0.0020 min^{-1}. By taking into account the fact that no AZS was adsorbed on the titania films, as already said, it is obvious that photodegradation is carried out in the liquid phase and it is also obvious that O_2•^- species, which are only formed in the presence of oxygen, play a major role. Furthermore, the fact that photodegradation is carried out in the liquid phase explains why it is relatively slow. As a matter of fact, it is known that photodegradation rates are higher when the target substance is adsorbed on the photocatalyst.

### 3.2. Photodegradation of AZS in presence of noble metals deposited on TiO_2 films

Silver or platinum metal ions were deposited on the TiO_2 surface by submerging titania for several minutes in aqueous solutions of metal salts. Metal cations could be adsorbed in TiO_2 films at substantial quantities because of the relatively high specific surface area of the films. The platinum complex is in the form of divalent tetrachloride ion. By accepting two photo-generated electrons, it is converted into zero valence metal according to the following reaction:

\[
\text{PtCl}_4^{2-} + 2e^- \rightarrow \text{Pt}^0 + 4\text{Cl}^-.
\]  

(3)

However, XPS measurements for UV illuminated samples gave peaks with binding energies of 72.85 and 76.13 eV, consistent with published values [22] for Pt^{10}. In addition to UV exposure, thermal heating (500 °C) of the metal-deposited films gave a variation of the binding energies of platinum, which were then located at 70.5 and 73.86 eV, consistent with Pt^{0} [22]. It is obvious that weak UV light intensity from black light irradiation was not enough for the complete transformation of tetrachloroplatinate ion into zero valence platinum. However, it is possible that Pt^{11} is finally transformed into Pt^{0}, due to the prolonged UV illumination during photodegradation. In the case of silver ions, the results were more predictable as only one electron is enough to reduce silver ions and create zero valence noble metal on the surface of TiO_2, according to the following reaction:

\[
\text{Ag}^{+1} + e^- \rightarrow \text{Ag}^0.
\]  

(4)

This electron is easy to be generated, either by UV illumination under black light irradiation or by thermal heating of the films [23]. XPS measurements gave similar results for both procedures of silver reduction. The photodegradation of AZS herbicide in the presence of platinum modified titania films is presented in Figure 4. The films subjected to both UV and thermal treatment gave better photocatalytic efficiencies as compared with the films, which were only UV treated. Furthermore, the original salt concentration, which obviously affects the quantity of deposited metal, affected photodegradation efficiency. When the platinum salt concentration was 5 × 10^{-4} mol L^{-1}, we obtained the highest photocatalytic efficiency. At higher metal load, efficiency dropped, possibly due to screening effects. The photocatalytic degradation of AZS in presence of silver loaded TiO_2 is presented in Figure 5(a). As in the case of platinum, metal load affected photocatalysis, 10^{-2} mol L^{-1} silver salt concentration giving the best results. Silver deposited on TiO_2 films gave the plasmon resonance absorption [23] spectra of Figure 5(b). Absorbance values verified that there is a direct relation between metal load and salt concentration in the aqueous solution. Comparative studies of AZS photodegradation with pure or metal-loaded TiO_2 showed that the presence of platinum is effective for the destruction of the herbicide contrary to silver, which gave similar results as pure TiO_2 films (see Figure 6). The role of platinum in photocatalysis is not fully understood. The degree of enhancement of the activity of TiO_2 by platinisation seems to depend highly on the substance to be photodegraded [24], the characteristics and the amount of deposited Pt [25], and the structural and morphological properties of original TiO_2 [26]. In general, it is accepted, as already said, that in platinized TiO_2, a better separation of charge carriers is observed.

### 3.3. AZS photodegradation intermediates

Intermediate products of AZS photodegradation were studied by LC-MS-MS. Two major ions were detected in all photodegradation stages, one with m/z: 199 and the other with m/z: 244. These two fragments exactly correspond to the amino-dimethoxyprymidinome and 1-methyl-4-(2-methyl-2H-tetrazole-5-y)-1H-pyrazole-5-sulfonamide moieties, respectively, which AZS consists of. Ion 244 yielded a major further fragment ion at m/z: 83.3, which could correspond to the sulfonamide SO_2NH_2 group. The amino-dimethoxyprymidin-month-containing ion yielded a major fragment ion at m/z: 74.5, which could correspond to the break down of the carbon link between the two nitrogen
3.4. pH effect on the photodegradation of AZS

The photodegradation of AZS was also studied at various pH values. The results are presented in Figure 7. pH values were modified by adding either H₂SO₄ or NaOH, both at a concentration of 0.1 mol L⁻¹. The fastest photodegradation was obtained at natural pH (4.7), that is, in a pure solution of AZS in water. Photodegradation rates in basic pH were low and even lower in acidic pH. This behavior could be explained by the fact that AZS possesses chargeable chemical groups. It is highly probable that at low pH values, AZS is positively charged, and at high pH values, it is negatively charged. However, as it is well known that the surface of titania is also charged with the same charge at the same pH [27]. Therefore, at low or high pH, there is a repulsion that separates the catalyst surface from the target molecule and makes it more difficult for the reactive radical species, which are generated on the surface of the catalyst, to reach the target molecules; hence the lower photodegradation rates in these cases. The right axis scale of all graphs of Figure 7 represents the change of pH in the solution during illumination. In strongly alkaline and acidic solutions, pH decreased during photodegradation, but in the case of natural solutions, a pH increase was observed. The interpretation of the pH effect on the photocatalytic process is a rather difficult task because of the multiple roles that various factors could play, such as electrostatic interactions with the semiconductor surface, chemical structure of the original substance and of its fragments, and formation of active radicals.

4. CONCLUSIONS

The herbicide azimsulfuron can be effectively photodegraded by employing pure or noble metal-modified titania nanocrystalline films as photocatalysts with black light tubes as low-intensity UV illumination source. The catalyst was deposited by the sol-gel method on glass rings; it could be
talline TiO₂ films. The pH of AZS aqueous solution affected photodegradation procedures. The best photocatalytic rates were achieved in the case of platinum modified nanocrystalline TiO₂ films. The pH of AZS aqueous solution affected photodegradation rates. The fastest rate was obtained in the case of natural pH of the solution.

Figure 7: Effect of pH variation in the photodegradation rates with pure nanocrystalline titania films (a) in acidic pH; (b) in natural pH of AZS in water; (c) in alkaline pH value.

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