J. Chil. Chem. Soc., 62, Nº 2 (2017)

ELECTROCHEMICAL AND PHOTO-ELECTROCHEMICAL PROCESSES OF METHYLENE BLUE OXIDATION BY Ti/TiO, ELECTRODES MODIFIED WITH FE-ALLOPHANE

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ABSTRACT

This work reports the degradation of methylene blue (MB) on Ti/TiO₂ and Ti/TiO₂/Fe-allophane electrodes in a pH 3 using 0.1 M Na₂SO₄ as support electrolyte. SEM micrographs show a homogeneous distribution of TiO₂ over the whole electrode surface forming nanotubes and nanopores. Fe-allophane modified electrode shows the formation of large-grains agglomerate on the electrode surface due to allophane, which provides a greater surface area to the electrode due to meso and micropore structures. Preliminary cyclic voltammetry show that Ti/TiO₂ has the typical voltammetric response due to Ti(III)/Ti(IV) pair. Diffusional problems were observed through of the film when the electrode is modified with Fe-allophane modifying the quasi-reversible process Ti(III)/Ti(IV). Different kind of methodologies in the degradation process were used: Electrochemistry (EC), Photochemistry (PC), Photoelectrochemistry (PEC) and Adsorption (Ads). These methods were developing to discard any reaction or interaction that is not of interest. On Ti/TiO₂ with PC and Ads methodologies was not observed any activity to MB degradation showing that is not photosensitive and that the interaction between this and surface electrode is low. But with EC and PEC degradation to 55% is reached after 3 hours of electrolysis. With Ti/TiO₂-Fe-allophane electrodes are observed a higher activity for all methodologies. The PC and Ads methods show that MB degradation reaches to ~20 % of the initial concentration. As mentioned above, the PC and Ads processes no show degradation on Ti/TiO₂, therefore the degradation it only due to the adsorption of MB in/on allophane coat behaving as concentrator matrix. A lower improvement is observed with EC process %, showing an improvement of the degradation with the presence of Fe-allophane. The results indicate that the main role of Fe-allophane on the electrode surface by oxidation products. With PEC is reached the higher degradation value of ~88 %, showing an improvement of the degradation with the presen

Keywords: Ti/TiO₂; Fe-Allophane; methylene blue; electrochemical and photoelectrochemical oxidation.

1. INTRODUCTION

The presence in the environmental of a wide range and variety of recalcitrant organic pollutants from agricultural, industrial and domestic activities made necessary to remove it through an efficient way using an environmental friendly technology. In this sense, advanced oxidation processes (AOPs) based on the combination of strong oxidants such as ozone, oxygen or hydrogen peroxide and the use of catalysts as transitions metals or semiconductors solid together with sources of radiation or ultrasound [1] have been widely proposed as a promissory way to degrade organic pollutant. The AOPs are based on the generation of a highly reactive species, which react with the most organic compounds by two principal ways: the first one is by direct electro-oxidation [2]. The nature, kind and the features of the electrode material is one the most important variables to determine which of these two ways follow the degradation process.

Among the solid catalysts, the use of semiconductor such as titanium dioxide (TiO_2) has been widely studied in photochemical processes due to their potential for mineralizing a wide variety of recalcitrant organic pollutants [3,4]. The titanium dioxide (TiO_2) is an excellent catalyst due its photostability, high photocatalytic activity, reusable characteristics, low cost and non-toxicity [3,5]. For this reasons in one of the most used support material for different electrochemical (EC) and photoelectochemical (PEC) degradation processes.

The photocatalytic degradation mechanism has been extensively discussed, but there are agreements that this starts with the photogeneration of an electronhole pair (Eq. 1) due to the UV illumination on TiO₂. These charged pairs might suffer a recombination or react with one water or oxygen molecules to produce reactive species (Eq. 2 and 3). The formation of hydroperoxyl radical is possible by electron transfer from TiO₂ particles to molecular oxygen and consecutive protonation (Eq. 4) [6].

$TiO_2 + hv \rightarrow e^2 + h^+$	(1)
$e^{-} + O_{2} \rightarrow O_{2}^{+}$	(2)
$h^+ + H_2O \rightarrow H^+ + HO^{\bullet}$	(3)
$e^{-} + O_{2}^{-} + H^{+} \rightarrow HO_{2}^{\bullet}$	(4)

The photocatalyts TiO_2 have been used to degrade a large variety of recalcitrant organic pollutants [2, 3, 7, 8, 9]. It has been reported that the photocatalytic performance of TiO, is primarily determined by its

physicochemical properties such as crystal structure, particle size, surface area and microporosity. The most used TiO_2 is the anatase structure due to photocatalytic properties to generate the hole-electron pair when it is irradiated with UV light by process mentioned above.

It has been widely reported that the large band gap of TiO_2 limits its photoresponse to ultraviolet region (< 400 nm) [10]. As is known, only a small proportion (approximately 4%) of the solar spectrum corresponding to UV region [11, 12]. Thus, the development of new efficient photo-catalysts must be developed thinking in a device that uses a higher spectra region as visible light, could involve the visible light absorption, which is 42% approximately of total solar energy. Also, many studies have demonstrating that the use of nanostructures could improve the catalytic efficiency of photo-catalysts due to large surface area and micro and mesoporosity, which could increase the adsorption of target molecules and reach higher reaction ratios [10,11,13].

In this sense, there is evidence that the modification of TiO, with other narrow band gap materials, such as iron oxide could be made possible to absorb the photons in the visible range improve the adsorption properties of the surface [12, 14, 15, 16]. In the case of iron oxide studies reported that the Fe³⁺ ions increase the photocatalytic efficiency of TiO₂ since Fe³⁺ can serve as shallow trapping sites and Fe loading on TiO,-supporting montmorillonite catalyst (Fe-TiO2-MMT) for β-naphtol degradation under UV-A irradiation exhibited higher surface areas, pore volumes and extended absorption profiles through longer wavelength [5]. The mesoporous structures of the supported catalysts improved their dark adsorption capacities and degradation abilities. The existence of Fe enhances the electron transfer reaction in the catalysis. Studies photo-catalytic oxidation of 4-chlorophenol under visible ultraviolet radiation using mesoporous structures of Fe_2O_3 /TiO₂ [12], showing that the Fe₂O₂ incorporation allows the light absorption in the visible solar irradiation improvement the 4-chlorophenol oxidation. Through synthesized α -Fe₂O₂/ TiO, nanotubes [9] used for phenol degradation by heterogeneous electro-Fenton which is compared with α -Fe₂O₃ nanoparticles and TiO₂ nanotubes. They reported that the composite Fe₂O₃/TiO₂ nanotubes electrodes were more effective, obtaining a 100% of removal efficiency after 60 minutes of treatment.

Therefore, the use of highly active photocatalyts able to absorb the solar radiation in a wide region (UV and visible range), with a porous structure and large surface area that allow a higher adsorption of the organic could be there a promissory alternative in the degradation of recalcitrant organic pollutants by electro-chemical methods.

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The use of clays minerals and TiO, as support is a promissory alternative due that the clays exhibit large surface area and micro and mesoporosity, besides some clay are in natural association with iron species, as is the case of allophane nanoclays. Allophane is a nano-clay size mineral of a widespread occurrence in volcanic ash soils (mostly Andisols). Their unitary structure consists in hollow spherules, highly porous with a diameter in the nanoscale range (3-5 nm) [17]. Due allophane properties, such as, large surface area, unique structure and highly porosity, iron oxide supported allophane nanoclays with a hydrous feldspathoids structure were reported as highly active and stable in the phenol oxidation by heterogeneous Fenton reaction [18] and for the atrazine oxidation in the heterogeneous electro-Fenton process [19]. Recently, has been carried out a study with allophane-TiO, nanocomposite photocatalyst to degrade trichloroethylene (TCE) under UV irradiation. The catalytic efficiency was compared with the TiO, and the allophane-TiO, mixture [20]. TCE was transformed into intermediate products, on the TiO₂, during the UV irradiation further this compound was rapidly absorbed on the allophane and gradually degraded after diffusing to the TiO2. The diffusion rate on the nanocomposite was faster than on the physical mixture due to high dispersion of allophane into the TiO₂. In the same way, it was also used dyedispersing allophane-titania composite electrodes to study its photoelectric conversion properties [19] and other studies the photocatalytic decomposition rate of gaseous acetaldehyde using TiO,/allophane mixed powders with acid treatment was performance [21]. Despite of this studies that using a mixture of allophane nanoparticles and TiO2, Fe-allophane/TiO2 as efficient and stable photocatalyts for organic pollutant oxidation by electrochemical and photoelectrochemical methods have not studied yet.

Therefore, the aim of this work is study of methylene blue degradation by electrochemical (EC) and photoelectrochemical (PEC) methods using Ti/ TiO₂ electrodes modified with allophane nanoclays. Methylene blue has been widely used in the dye and pigment industries and has been selected as model pollutant due to its environmental significance, ease of analysis, and relative solubility in water [22].

2. EXPERIMENTAL

Reagents

All chemicals were used as received. Titanium foil (99.7%), Nafion® perfluorinated ion-exchange resin and methylene blue were purchased from Sigma-Aldrich. H,SO₄ (95-97%) and HF (48-51%) were supplied from Merck.

Preparation of Fe-allophane

Allophane nanoclays with a theoretical Si/Al ratio of 3.0 and similar structure to hydrous feldspathoids were synthesized by co-precipitation method [23]. The allophane nanoclays were impregnated with 6% wt. iron oxide (Fe-allophane) with a wet impregnation method previously reported [17].

Preparation and characterization of Ti/TiO_2 and Ti/TiO_2 modified with Feallophane electrodes.

The Ti/TiO₂ electrode was prepared by electrochemical anodized in presence of $0.5 \text{ M} \text{ H}_2\text{SO}_4$ and 0.04 M HF. The titanium plate (6 x 2.5 cm, 0.25 mm thickness) were cleaned with acetone-ethanol (50:50) and washed with

distilled water under ultrasonic bath. The anodized process was performed at controlled potential (25 V) with a Power Source MCP Lab Electronics M-10 QD305, under continuous stirring during 4 hours.

Ti/TiO₂ electrodes were modified painting the electrode surface with a dispersion prepared according to method reported previously [24]. In brief, 0.5 mL of Nafion® perfluorinated ion-exchange resin and 4 mL of an aqueous dispersion containing 30 mg of Fe-allophane were mixed and ultrasonically homogenized. The solvent was evaporated under air, in a stove at controlled temperature (60°C) for 2 h. The Ti/TiO₂ with Fe-allophan modified electrode (Ti/TiO₂/Fe-allophane) was calcined at 460°C for 4 hours to activate the catalytic properties of TiO₂.

The surface morphology of the electrodes was examined by scanning electron microscopy (SEM) using a Tescan Scanning Electron Microscopy (SEM).

The electrodes were also characterized by cyclic voltammetry (CV). CVs were obtained with CH-Instruments potentiostat electrochemical workstation model CHI760D. A platinum wire and saturated calomel electrodes (SCE) were used as counter and reference electrodes, respectively. The CVs were registered at 0.1 V s⁻¹ and a solution at pH 3 under nitrogen-saturated, using 0.1 M Na_sSO_s as support electrolyte.

Methylene blue oxidation by electrochemical and photo-electrochemical methods

Methylene blue (MB) oxidation by electrochemical method was studied using the Ti/TiO₂ and Ti/TiO₂ modified with Fe-allophane (geometric area of 20 cm²) as working electrode. A platinum wire and saturated calomel electrode (SCE) were used as counter electrode and reference electrode, respectively. A single compartment electrochemical quartz cell was employed using a typical three-electrode arrangement. The electrochemical oxidation (EC) was performed using a CH-Instrument potentiostat at controlled potential (-1.0 V) in a mechanically stirred solution during 3 hours. The initial methylene blue concentration was 15 mg/L (100 mL), prepared in 0.1 M Na₂SO₄ adjusted at pH 3 and 0.1 mL of the samples were withdrawn at predetermined time intervals.

Methylene blue oxidation by photo-electrochemical (PEC) process was studied in the above described conditions. UV medium pressure mercury lamp UVP Black Ray B-100 AP was positioned in front of the quartz electrochemical cells. The light source was positioned perpendicularly at 30 cm lighting of the working electrode.

Methylene blue concentration for both EC and PEC experiments was determined immediately by measuring the absorbance of the solution at 665 nm (λ_{max} for MB) using a Cary 1e UV-visible spectrophotometer.

3. RESULTS AND DISCUSSIONS

Characterization of Ti, Ti/TiO, and TiO/Fe-allophane electrodes by SEM.

Figure 1 shows the SEM micrographs for the Ti base, Ti/TiO₂ and Ti/TiO₂/Fe-allophane electrodes. Ti base electrode (Fig.1a) shows the typical characteristic of an amorphous and non-homogeneous surface with defects generated in the laminating process and/or mechanical polishing. In the Ti/TiO₂ electrode (Fig. 1b) it is possible to see the homogeneous size particle of TiO₂ formed on Ti sheet and distributed all over the surface.



Figure 1: SEM image of (a) Ti base and (b) Ti/TiO₂, (c) Ti/TiO₂/Fe-allophane electrodes used at MB degradation.

The possible reactions that occur throughout the formation at anodized process are described by (eq. 5 and 6)

$$\begin{aligned} &\Gamma i + 2H_2O \to TiO_2 + 4H^+ + 4e^- & (5) \\ &\Gamma iO_2 + 6F^- + 4H^+ \to [TiF_6]^{2-} + 2H_2O & (6) \end{aligned}$$

Fluoride ions present in an aqueous medium assist the formation of TiO₂ nanotubes and nanopores as shown in the insert in Fig. 1b. Whereas the electrode is formed in an aggressive medium the concept for the origin of the porosity it is validated [25] due to the acid medium especially hydrofluoric acid used in anodizing. The pore formation produces a greater surface area of the electrode of Ti/TiO₂ compared to the Ti base electrode.

Finally, the electrode surface with Ti/TiO₂/Fe-allophane (Fig. 1c) is observed rougher than the other electrodes. The Fe-allophane image shows that larger grains agglomerates are formed on the surface. This provides a greater surface area the electrode due to presence of meso and micropores in the structure. It has been noticed that the use of allophane with a higher amount of silicon in their composition produces a tetrahedral polymer with greater spaces in their structure improving the porosity of the material [26].

Characterization of Ti/TiO_2 and TiO_2/Fe -allophane electrodes by cyclic voltammetry

 Ti/TiO_2 and Ti/TiO_2 /Fe-allophane electrodes were characterized by cyclic voltammetry (Figure 2). This methodology allowed clearly distinguish the redox Ti(III)/Ti(IV) pair characteristic of TiO_2 formed on a Ti electrode (Fig. 2a peaks I and II)). The redox reaction involved is described by (eq. 7).

$$Ti^{III}OOH \rightarrow Ti^{IV}O_2 + H^+ + e^-$$
 (7)

On the other hand, in the Ti/TiO_2 electrode modified with Fe-allophane was not possible to observe the couple redox Ti(III)/Ti(IV) (Figure 2b) because the dispersion of Fe-allophane used to paint the electrode produce diffusional problems through of the Nafion added during the preparation of $Ti/TiO_2/Fe$ -allophan electrode.

Furthermore, studies of oxygen reduction reaction (ORR) was performance to determine the activity to this reaction. The ORR would be essential to produce reactive species for both Ti/TiO, and Ti/TiO_/Fe-allophane electrodes as indicated in equation 2 and 4 above. In the presence of O₂ (Figure 2 a and b solid line) the cyclic voltammetry of Ti/TiO₂ show an increase in current associated with the oxygen reduction reaction (ÔRR). On the other hand, Feallophane electrode the higher ORR current (10 mA) was associated to the presence of Fe centre, which it is known as reaction catalyst oxygen reduction.



Figure 2: Cyclic voltammetry of (a) Ti/TiO₂ and (b) Ti/TiO₂/Fe–allophane. O₂ (solid line) and N₂ (dot line). Electrolyte solution 0.1 M Na, SO₄ adjusted to pH 3. 0.1 V s⁻¹.

Electrochemical and photoelectrochemical oxidation of methylene blue

Degradation of MB with Ti/TiO, electrode

The MB degradation was performance with Ti/TiO_2 and Ti/TiO_2 /Feallophane electrodes by different methodologies to evaluate the occurrence of other kind of reaction, i.e., the MB degradation not occur by other way of reaction. The only paths of reaction of interest are those described by the equations 1 to 4. The methodologies used are Adsorption on the electrode surface without the external factors mainly may occur through electric potential and electromagnetic radiation; electrochemistry using with electrolysis at controlled potential; photochemistry with mercury lamp UV light irradiation and photoelectrochemistry that is a combination between photochemistry and electrochemistry. The different reactions ways electrochemistry and photoelectrochemistry consider the formation of species highly reactive to degrade an organic substrate present in solution [27]. The comparison of different methodologies used is shown in Figure 3. Photochemical degradation (Fig. 3 curve •) shows a decrease in the MB concentration of 2.9%, this is due to the low reactivity of MB to UV light and therefore no processes degradation were observed, which is reflected in its low value of degradation constant of 0.06 min⁻¹ (see Table 1). Similar result is obtained for MB adsorption method (Fig. 3 curve \mathbf{V}) with the Ti/TiO₂ electrode this is mainly because there is not interaction between the surface Ti/TiO₂ electrode and MB molecule. Both MB molecule and titanium dioxide have positive charge at this pH which produces an electrostatic repulsion and an increase of net positive charge on surface [8,4].

For electrochemical (Fig. 3 curve \blacktriangle) and photo-electrochemical methods (Fig. 3 curve \blacksquare), the percentages achieved are similar between them giving

45.8 and 43.7%, respectively. These lower differences are produce by its degradation constant being for EC and PEC of $0.78 \times 10^{-3} \text{ min}^{-1}$ and $0.61 \times 10^{-3} \text{ min}^{-1}$, respectively. These results were not expected for photo-electrochemical. The oxygen reduction reaction by electrochemical could produce hydrogen peroxide that has capacity to degrade the MB partially. However, regarding the photo-electrochemical reaction, it was desired to produce synergic effect between photochemical with hole-electron pair formation and electrochemical avoid the recombination.

Results obtained were similar those observed by electrochemical method since the UV irradiation has not effect on MB degradation. This might be attributed to the formation of residues during the degradation reaction which are deposited over surface as a barrier blocking the path of UV radiation to the electrode surface. Therefore, the active Ti/TiO₂ site where the formation of reactive species occurs is occupied by residue of reaction.

The low increased in the degradation of MB when is irradiated with UV light during electrolysis, it is due that the hole-electron pair which are charge carriers are being inactivated either by recombination among them or parallel reaction, promoted by the irradiation of UV light on the oxide film. Some authors have proposed the existence of localized zones where the holes and/ or electrons are deactivated by process of recombination. These sites would be associated with the morphology of the oxide film such as structural defects or impurities [29, 30].



Figure 3: (a) Concentration decay of a 15 ppm MB solution and (b) pseudo first order decay for MB degradation showed in (a). (\blacktriangle) EC, (\bullet) PC, (\blacksquare) PEC and (\blacktriangledown) adsorption methodologies using Ti/TiO₂ electrodes at 0.1 M Na₂SO₄ adjusted to pH 3.

Degradation of MB with Ti/TiO./Fe-allophane modified electrode

Figure 4 shows the results of absorbance decrease for MB degradation on Ti/TiO_2 modified with Fe-allophane electrodes. This figure shows in both EC and PEC cases that there is a great decrease in the absorbance of MB to the maximum wavelength.

From absorbance values are obtained the maximum wavelength obtaining the ratio A/A_0 or $[MB]/[MB]_0$ which shows for the concentration decay of MB by EC and PEC (Figure 5).

Figure 5 a and b shows that all methodologies have first-order kinetics, i.e. only it will depend on the MB concentration, showing higher efficiency of the degradation by photoelectrochemical method. The high adsorption of MB molecules close to 80% at electrode occurs by a charge interaction, since the Fe-allophane has a higher proportion of Si with respect to Al. This would give negative charge on surface at pH acid and MB is positively charged, causing a charge attraction facilitating adsorption process. Negative surface charges that present allophane is explained by a virtual isomorphous replacement of Si⁴ by Al⁺³ [31].

Adsorption is not an expected result, since the electrode could be unrecovered and certainly this adsorption affect the results observed in the other methods and the relative degradation would not be a result of MB degradation, rather of represent MB degradation.

Furthermore, the decrease of MB concentration by photoelectrochemical methods was higher compared to electrochemical method, this occurs because the degradation constant for PEC process $(1.95 \times 10^{-3} \text{ min}^{-1})$ is higher than EC process $(1.26 \times 10^{-3} \text{ min}^{-1})$. This can be explained if it is considered the activation of the electrode in the presence of UV light generating the electron-hole pair in TiO₂ at electrode or the effect produced by Fenton iron oxides present in the electrode.

Brief, all methodologies for the electrode modified with Fe-allophane gave results of similar magnitude (Table 1). However, it is complex to point out what the process is contributing in the decreasing of MB concentration in each methodology.



Figure 4: Absorbance decay of a 15 ppm MB solution by (a) electrochemical and (b) photoelectrochemical methods.



Figure 5: (a) Concentration decay of a 15 ppm MB solution and (b) Pseudo first order decay for MB degradation showed in (a). (\blacktriangle) EC, (\bullet) PC, (\blacksquare) PEC and (\blacktriangledown) adsorption methodologies using Ti/TiO₂/Fe-allophane electrodes at 0.1 M Na₂SO₄ adjusted to pH 3.

4. CONCLUSIONS

SEM and voltammetry studies have made possible to identify the properties and relevant characteristics in each electrode. Through SEM, the morphology of the electrodes was observed. For Ti electrode a smooth surface was seen; in the Ti/TiO₂ electrodes the formation of pores at surface was observed and in the Ti/TiO₂/Fe-allophane electrodes the formation agglomerations and pores is the principal feature, attributed to the own porosity of Ti/TiO₂ used as a base and the structural characteristics of Fe-allophane.

Degradation of MB on Ti/TiO₂ electrodes showed a decrease of the MB concentration by electrochemical and photochemical methods reaching values of 43.7% and 45.8%, respectively. With these Ti/TiO₂ modified Feallophane electrodes was reached the highest values of the reduction of the MB concentration.

Additionally, higher efficiency is obtained with the electrode Ti/TiO₂modified Fe-allophane compared to the electrode of Ti/TiO₂. However, the results show that the photoelectrochemical application produces a greater decrease in MB in solution, possibly associated with a combination of UV and effect the Fenton reaction due to iron oxide present in the modified electrode.

Table 1: Percentage degradation and pseudo first order constant to MB with Ti/TiO_2 and Ti/TiO_2 /Fe-allophan electrodes obtained by EC, PEC, PC and adsorption methods. Time experiment 180 min.

	Ti/TiO ₂		Ti/TiO ₂ /Fe-allophan	
Methods	% decrease of [AM]	Pseudo first order constant / min ⁻¹ x10 ³	% decrease of [AM]	Pseudo first order constant / min ⁻¹ x10 ³
EC	45.8	0.78 ± 0.09	80.1	1.26 ± 0.07
PEC	43.7	0.61 ± 0.03	86.9	1.95 ± 0.09
PC	2.9	0.06 ± 0.03	73.1	1.61 ± 0.01
Ads	0	N.D.*	79.7	1.61 ± 0.07

(*) Not determined

ACKNOWLEDGEMENTS

This work was supported by FONDECYT-CONICYT Grant 11090336 and Dicyt-USACH 21441BG.

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