Fenton’s oxidation of Orange II solutions using heterogeneous catalysts based on saponite clay

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Abstract Fe-clay catalysts have been prepared and tested for Orange II oxidation with H2O2 in aqueous solution. The reaction is carried out in a batch reactor, using different hydrogen peroxide concentrations, and in a wide range of temperature and pH values. Twelve samples were prepared, with three different iron loads (7.5, 13.0 and 17.0 %, w/w), and using four iron salts as precursors, namely Fe(II) acetate, Fe(II) oxalate, Fe(II) acetylacetonate and Fe(III) acetylacetone. The samples were characterized using X-ray diffraction, thermal analysis, infrared spectroscopy and adsorption of nitrogen at 77K. The catalytic results show that these solids present good catalytic properties for the degradation and mineralization of Orange II solutions, allowing to reach, in the best conditions and after 4h of oxidation, 99% of dye degradation with 91% of TOC (Total Organic Carbon) reduction (at 70ºC), using only ca. 90 mg of clay catalyst per litre of solution. Nevertheless, 96% of dye removal with 82% of mineralization were also reached at 30ºC. Besides, the amount of iron released into the final solution is lower than 1 ppm, in the worst of the cases, and 0.09 ppm in the best case.

Keywords Clay-Fe catalysts, Fenton’s reagent, Heterogeneous catalysis, Orange II, Oxidation.

Introduction

The reaction of hydrogen peroxide with ferrous ion in acidic aqueous solutions (Fenton’s reagent) can be used as source of hydroxyl radicals for the degradation of organic pollutants in wastewater treatment (Gallard and Laat, 2001). The HO• radicals generated through reaction (1) are highly oxidative, non-selective, and able to decompose many organic compounds, including dyes (Swaminathan et al., 2003).

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Fe^{2+} + H_2O_2 → Fe^{3+} + OH^- + HO^-
\] (1)

However, the homogeneous Fenton process has a significant disadvantage: the removal of the sludge-containing Fe ions at the end of wastewater treatment is expensive and needs large amount of chemicals and manpower. To overcome this disadvantage some attempts have been made to develop heterogeneous Fenton process by means of heterogeneous catalysts, prepared by incorporating Fe ions or Fe oxides onto porous supports (Gemeay et al., 2003; Letaief et al., 2003). Transition metal complexes supported on different surfaces such as metal oxides, resins, and mixed (Al-Cu) pillared clay have also been used as potentially active catalysts for the decomposition of H2O2 and the oxidative degradation of organic contaminants. Among the porous solids used as supports for the iron phases, some are based on silica, alumina, silica-alumina and cation-exchanged resins, and all these catalysts have been used for degradation and mineralization of dyes (Gemeay et al., 2003). It is also important to remark the use of catalysts prepared by co-intercalation of two natural smectites (Wyoming SWy-l and Tunisia-Gafsa VI) with Fe-Al polycations, obtained by polymerisation of a mixture of FeCl3 and chlorhydrdrol (Letaief et al. 2003).

Pillared clays (PILCs) are one of the families of microporous solids developed by Molecular Engineering more studied in recent years. The PILCs are prepared by intercalation of appropriate polyoxocations into the clay, followed by calcination at relatively high temperatures (Belver et al., 2004), showing a bidimensional microporous network of molecular dimensions, with the pillars occupying the interlayer space defined by the clay layers. Alternatively, it can be said that the clay layers avoid the aggregation of the metallic oxide clusters, giving rise to much dispersed particles. The number and size of the pillars in the interlayer region condition the pore parameters of the pillared clay structure (Vicente et al., 2000).

In this work, several heterogeneous catalysts based on saponite impregnated with iron salts were prepared. A saponite has been intercalated with Al polycations, and the pillared solid obtained after calcination at 500ºC has been used as support for impregnation with iron. Four Fe-precursors have been used and for each one three different loads of iron (7.5, 13.0 and 17.0 Fe wt. % in the final catalysts) were considered. The obtained heterogeneous
catalysts are used in the oxidation of the non-biodegradable dye Orange II (O-II) in water solution, using a batch reactor, and carrying out experiments under a wide range of experimental conditions. The effectiveness of these catalysts in the oxidation of the dye, as well as the influence of the variables of the synthesis and of the conditions of reaction in the catalytic activity, are discussed.

Methods

Preparation of the catalysts

Saponite clay from Yunclillos (Toledo, Spain) was kindly supplied by TOLSA (Madrid, Spain). The fraction with particle size smaller than 2 µm was obtained by dispersion in water and controlled decantation of the natural clay, and was then used for intercalation/pillaring experiment. It is a well ordered smectite with basal spacing of 14.4 Å, BET specific area of 152 m²/g and cation exchange capacity (CEC) of 0.9 meq/g.

Saponite was intercalated with \([\text{Al}_{13}\text{O}_{4}(\text{OH})_{24}(\text{H}_2\text{O})_{12}]^{7+}\) polycations (in short, \(\text{Al}_{13}\)) by using a standard procedure (Bottero, et al., 1980; Lahav, et al., 1978). First, intercalating \(\text{Al}_{13}\) solution was obtained by careful hydrolysis of an \(\text{Al}^{3+}\) solution, derived from \(\text{AlC}_{3}\cdot6\text{H}_2\text{O}\) (Panreac, purissimum), with 1M NaOH (Panreac, purissimum), using a ratio \(\text{OH}^-/\text{Al}^{3+}=2.2\), stirring vigorously and aging 24 h (final pH=4.1). Once the solution was aged, it was added to a previously (12 hours) prepared suspension of saponite in water, employing an Al/clay ratio of 5 mmol/g. The new suspension was stirred for 24 h, and then washed by centrifugation and dialysis until absence of chloride (Ag⁺ test). The solid obtained was dried at 70°C for 16 h, and then heated to 500°C at a heating rate of 1°C/min and maintained at this temperature for four hours. The solid obtained was used as support for impregnation. As this is the support used for the preparation of all supported catalysts, it may always be referred to as “the support”.

Impregnation was carried out by means of the incipient wet impregnation method. As indicated, \(\text{Al}_{13}\)-pillared saponite was always used as support, and four different Fe-salts as precursors (Fe(II) acetate, Fe(II) oxalate, Fe(II) acetylacetonate and Fe(III) acetylacetonate). For each precursor, the amounts needed for obtaining 7.5, 13.0 and 17.0 Fe wt. % in the final catalysts were calculated. These amounts were dissolved in the minimum amount of the appropriate solvent (water for the acetate and the oxalate, and acetone for both acetylacetonates), the obtained solutions being used for the impregnation. The first two precursors were very soluble and a single impregnation step was needed, while five cycles were needed for both acetylacetonates because of their low solubility. After completing the impregnation, the solids were calcined at 500°C following the calcination procedure described above, thus obtaining the final catalysts. These catalysts are designated by a precursor-amount notation, for example Fe(II) acetate-7.5 designates the solid prepared using as precursor Fe(II) acetate and containing 7.5 wt. % of Fe in the final catalysts.

Characterization of the catalysts

- Powder X-ray diffraction patterns (XRD) of the solids were recorded using a Siemens D-500 diffractometer at 40 kV and 30 mA using filtered Cu Kα radiation (\(\lambda=1.5418\) Å) over a 20 range from 2 to 65 degrees.
- FT-Infrared spectra were recorded in the 4000-350 cm⁻¹ region with a Perkin-Elmer 1730 FT-IR spectrometer, using a He-Ne laser source (\(\lambda=632.8\) nm), in KBr pellet (0.001 g sample with 0.3 g KBr), and 15 scan per minute to improve the signal-to-noise ratio.
- BET specific surface areas were determined by adsorption of nitrogen at 77K, measuring five adsorption points by using a Micromeritics Gemini apparatus.
- Simultaneous thermal gravimetric (TG) and differential scanning calorimetric (DSC) analyses of samples were carried out using a TA-SDT Q600 Instrument. Samples of about 40 mg were heated in air (flow rate = 100 mL/min) from room temperature up to 900 °C, with a heating rate of 10 °C min⁻¹.

Catalytic activity

Chemical oxidation of 0.1 mM Orange II (Fluka p.a.) solutions was carried out in a batch reactor of 1 L of capacity, with constant stirring by means of a FALC F30ST magnetic stirrer (230 rpm), and with a permanent control of the temperature by a thermostatic bath (Huber, Polystat CC1 unit). After stabilisation of temperature and pH, the catalyst was added, and the beginning of the reaction (t=0) was taken when \(\text{H}_2\text{O}_2\) was added. Absorbance, temperature and pH were permanently measured by means of a Philips PU8625 UV/VIS spectrophotometer, a
temperature sensor, and a pH EDT Instruments RE 357 TX electrode, respectively. For the on-line determination of the absorbance at λ=486 nm (characteristic wavelength of the Orange II molecule) a flow-through cell was used, and recirculation was obtained by a peristaltic Watson-Marlow 5055 pump, using a flow-rate of 100 mL/min. Acquisition of data was carried out by Labview 5.0 software, from National Instruments. The total duration of each oxidation experiment was 4 hours. In several cases, Total Organic Carbon (TOC) was measured using a Shimadzu 5000A spectrophotometer, model TOC-5000 CE, equipped with an automatic sample injector. After taking the sample and before measuring TOC, reaction was stopped by adding Na$_2$SO$_3$ (that instantaneously consumes H$_2$O$_2$), followed by centrifugation and filtration for the perfect separation of the catalyst from the liquid phase. The total Fe in the final solution was determined using a UNICAM 939/959 atomic absorption spectrophotometer.

**Results and discussion**

**Characterization of the Catalysts**

Intercalation/pillaring experiment was carried out under standard conditions, and goes in a successful way. The solid intercalated with Al$_{13}$ polycations and then calcined at 500°C, used as support for further experiments, shows the typical features of Al-PILCs. It has a layered structure with a basal spacing of 18.2 Å (Fig. 1A), and a specific surface area of 240 m$^2$/g. The impregnated solids maintain the layered structure, but with a remarkable loss of ordering compared to the support. These catalysts, once calcined at 500°C, show a weak diffraction effect, between 16.8 and 17.3 Å (Fig. 1A). However, as can be observed in these diffractograms, although the long-range ordering in the c-axis is low, it is not negligible, and all solids show a certain degree of layered-structure ordering. At the same time, the impregnation-drying process, mainly in the solids treated with acetylacetonate solutions, for which various impregnation cycles are needed, and the further calcination at 500°C, produce a certain collapse and delamination of the structure. Reflections independent of c-axis ordering do not show any variation with respect to the support, indicating that the impregnation does not affect the individual layers, but only their stacking. On the other hand, it may be underlined that no peaks due to iron phases are observed in the diffractograms. Considering the preparation method used and the results obtained from other techniques, it may be reasonably expected that phases close to Fe$_3$O$_4$ be formed after calcination of the impregnated solids at 500°C, by the removal of the organic moieties of the precursors, and oxidation, if needed, of iron. However, no oxides, neither oxi-hydroxides, peaks are found in the diffractograms, although the amounts of iron used in the impregnations are relatively high. So, it may be proposed that iron is as a very disperse phase on the surface of the support.

FT-IR spectra of the support and of various impregnated solids, both dried and calcined, are shown in Fig. 1B. The support has the bands characteristic of Al-PILCs. Thus, in the high wavenumber region, the spectrum is dominated by the stretching vibrations of the hydroxyl groups bonded to metallic cations and to water molecules. Bending of water molecules also produces an important effect close to 1630 cm$^{-1}$, while the bands characteristic of the tetrahedral layer of the clay dominate the region of low wavenumbers, being the Si-O-Si band, at 1007 cm$^{-1}$, the more intense of the spectrum, and appearing at lower wavenumbers the bands due to M-O bonds in the octahedral layer (mainly Mg-O and Fe-O, because of the chemical nature of saponite). Solids impregnated and dried show, in all cases, the bands due to the support, also showing bands assigned to the precursors. The bands due to the support do not show important variations with respect to the support before impregnation, only small differences in intensity. This is an expectable observation, because of the low influence of the impregnation on the structural bonds of the clay, where only surface hydroxyl groups may be affected, giving rise to interfacial coordination bonds with Fe$^{2+}$ and Fe$^{3+}$ cations from the precursors. On the other hand, the effects due to the precursors are strongly intense, as can be expected because precursors containing organic moieties have been employed. The characteristic bonds of each precursor are observed in each case, with characteristic vibrations of C-H bonds, carboxylate or carbonyl groups, etc. All these effects completely disappear when calcining the impregnated solids at 500°C, showing that organic moieties are completely removed at this temperature, thus transforming the precursor molecules into iron oxide-like phases. However, no peaks due to Fe-oxides, neither oxy-hydroxides, are observed in the spectra, suggesting once again a high dispersion of the iron phases.

It is noteworthy that the BET specific surface areas of all the samples (not shown for brevity) are comprised between 128 and 192 m$^2$/g, which represents a percentage of 53 - 80% of the value of the support. These values are relatively high considering the subsequent steps of the preparation of the catalysts, including impregnation, drying, and calcination at 500°C.
The thermogravimetric analyses curves of some of the impregnated samples (with Fe(II) oxalate) are displayed in Fig. 1C. Several weight losses can be observed. The removal of adsorbed water and physisorbed solvent occurs until ca. 150 °C, while the decomposition of the organic precursors occurs at higher temperatures (150-425 °C). The final weight loss observed in all cases in the 415-825 °C range is due to the dehydroxilation of the clay. For all samples, it is clear that the temperature of calcination guarantees the removal of the organic fragments and the obtaining of iron-like phases. The DSC curves for the samples containing 13 %wt. of Fe are shown in Fig. 1D. An endothermic peak is observed in the 30-150 °C for the sample containing acetate, as consequence of the highest weight loss of adsorbed water and solvent detected in the TG curve of this sample, with respect to those prepared with other precursors. The DSC peak corresponding to the decomposition of the most part of the organic material is centred at around 315 °C for acetate, 275 °C for Fe(II) acetylacetonate and 350 °C for oxalate and Fe(III) acetylacetonate. For these last three samples, other exothermic peaks of lower intensity can be observed in the 150-275 °C with associated weight losses in their corresponding TG curves. The removal of the last hydroxyl groups in the clay is observed as a clear weight loss close to 800°C, associated to an endothermal effect, and once this dehydroxilation is completed, it is followed by an exothermal effect corresponding to the phase transformation from saponite to enstatite, not associated with any weight loss in the TG curves. No peaks that could be associated to transformation of iron phases are observed.

Figure 1. Characterization of the catalysts: (A) XRD of the support and catalysts with 7.5 wt.% of iron calcined at 500°C, (B) FTIR of the support and impregnated solid with Fe(II) oxalate 17.0, before and after the calcination, (C) Thermogravimetric analysis of different dried samples impregnated with Fe(II) oxalate and (D) DSC curves of the samples impregnated with 13.0 wt. % of Fe.

Catalytic behaviour

Although the natural clay has already some iron (ca. 1 wt.%), no degradation of the OH solution was noticed when using it as catalyst under the typical conditions adopted in this work (\( C_{H_2O_2} = 6 \) mM, \( T = 30^\circ C \), \( pH = 3 \), \( w_{clay} = 91.5 \) mg/L, \( t = 4h \)), which may be related to the inaccessibility of such iron, located in the octahedral positions of the clay structure. Besides, there is any dye removal by adsorption (blank run in the same conditions but without hydrogen peroxide), what is due to the remarkably low concentrations of clay used in our work. Actually, the use of \( w_{clay} = 1 \)
g/L, an amount commonly found in the literature, provided about 36% removal by adsorption after 4h (but no equilibration was yet reached). Based in our previous work (Ramirez et al., 2006), among the twelve samples prepared two catalysts were chosen to study into more detail the effect of the temperature, pH and initial H₂O₂ concentration. They were those prepared with oxalate with Fe content of 7.5 and 17.0 wt. %. In this concern, it is especially remarkable the Fe(II) oxalate 17.0 catalyst, which showed a very good behaviour in terms of mineralization (81.6% of TOC reduction) and discoloration (95.9% of OII removal), after 4h of oxidation.

**Temperature effect**

The results obtained for the OII degradation at four different temperatures (10, 30, 50 and 70°C), using catalysts prepared with Fe(II) oxalate with 7.5 and 17.0 wt. %, are shown in Fig. 2. The results show clearly that the reaction rate increases when increasing the temperature, which was expected due to the exponential dependency of the kinetic constants with the reaction temperature (Arrhenius law). Nevertheless, the final OII concentrations, after 4 hours of oxidation, are very similar at 50 and 70°C. In fact, the eliminations obtained at these temperatures are already quite similar after two hours of reaction, with values around 98%, whereas at 10°C the elimination is practically negligible (ca. 8% after 4 h of reaction for both samples). Other authors have also found similar results during catalytic wet peroxide oxidation of phenol over Fe-exchanged clays (Catrinescu, et al., 2003; Guo, et al., 2003), the similar performances achieved at higher temperatures being due to the accelerated decomposition of H₂O₂ into oxygen and water.

The overall TOC removal for both catalysts at different temperatures showed that, as expected, the mineralization increases with the temperature, the performances reached being better for the sample with 17.0% of Fe (Table 1). For this sample, it is remarkable that the TOC removal is near 91% at T = 70°C and around 82% at 30°C. Although lower than those obtained at higher temperatures, the values of OII and TOC removal achieved at 30 ºC might be considered satisfactory. Taking into account that a lower temperature might reduce the process costs, 30 ºC was the temperature chosen to carry out the following runs, where other parameters are changed. In addition, iron leaching is smaller at 30 ºC than at 70 ºC and is not very significant after 4h (<0.25 mg/L, thus being below the value of the EU directives).

**pH effect**

The results obtained for the OII degradation using the Fe(II) oxalate catalysts at three different pH values are displayed in Fig. 3. The best results of the OII degradation were obtained at pH 3.0, for both iron loads. At the lowest value of pH tested, pH = 2.0, the reaction is very slow and an important increase of decolorisation activity was only noticed after ca. 2.5 hours of reaction. However, at pH 3.5 performances achieved are even worst, with color removals after 4h of only 43 and 51% for 7.5 or 17.0 wt.% of Fe, respectively (Figs. 3A and 3B). It must be stressed that additional experiments have also been performed at pH 4.0 (not shown for brevity) but practically no dye degradation was produced, even for much longer reaction times. Regarding TOC removal, conclusions are similar, with better performances for both samples at pH 3.0 (Table 1). Curiously, this was exactly the best pH value found by Feng et al. (2006) during OII degradation using clay-based Fe nanocomposites with photo-Fenton process, with a reaction rate decrease similar to that reported by us when one deviates from such pH value. Other authors
also found, using pillared clays, that the reaction rate was much higher for the pH value corresponding to the optimum pH determined when homogeneous Fe species were used as catalyst (Barrault, et al., 2000).

Table 1. TOC removal (%) after 4h of oxidation.

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<th>Experimental conditions</th>
<th>Catalyst</th>
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<td>Temperature (ºC)</td>
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Figure 3. pH effect on the degradation of OII solution using different catalysts: (A) Fe(II) oxalate 7.5 wt. % and (B) Fe(II) oxalate 17 wt. % (C\textsubscript{H2O2} = 6 mM, T = 30ºC).

Figure 4 shows the effect of the reaction pH on the iron leaching. It is clear that iron lost is more significant at pH 2.0, especially for the 7.5 wt.% catalyst. Therefore, for long-term stability it would be preferable to work at higher pH values. Feng et al. (2006) also found that iron leaching was much more significant at pH around 2. The Fe leaching at this low initial solution pH can be attributed to the dissolution of iron oxide at very acidic conditions. At pH 2.0 the amount of iron leached into solution is much higher for the 7.5 wt.% catalyst than for the 17.0 wt.% one (Fig. 4A vs. 4B), even though the catalytic performance is not better (Fig. 3A vs. 3B), thus supporting the importance of the heterogeneous catalytic process. On the other hand, it is noteworthy that in the pH range studied (2.0-3.5), the sample with more iron (17.0 wt.%) shows similar or even lower percentages of iron lost, and thus can be in practice used for more operation cycles.

$H_2O_2$ Concentration effect

The initial hydrogen peroxide concentration was varied between 3 and 24 mM, using the same catalysts as in previous sections. The obtained results (Fig. 5) show, for both samples, a similar behavior in terms of dye degradation for the intermediate $H_2O_2$ concentrations (6 and 12 mM), whereas the reaction goes by more slowly when the concentration is smaller (3 mM) or higher (24 mM). The increase of the oxidant concentration from 3 to 12 mM leads to an increase in the reaction rate, as expected, because more radicals will be formed. Nevertheless, for a very high hydrogen peroxide concentration (24 mM) the performance decreases. The existence of an optimum hydrogen peroxide concentration is typical and well-known in Fenton’s oxidation. This optimum value was previously found by us to be 10 mM for OII degradation in homogenous Fenton reaction, using catalytic iron sulphate (Ramirez, et al., 2005). At higher $H_2O_2$ concentrations the scavenging of HO$^-$ radicals will occur, which can be expressed by the following reaction:
The influence of the H₂O₂ concentration on the mineralization (Table 1) is similar as for decolourisation, with an optimum oxidant concentration of 6 mM for both catalysts. In spite of the final performances attained seem to be only slightly affected by the peroxide dose, it is clear that for H₂O₂ concentrations above that value the final TOC removal decreases a little bit, this effect being similar to those reported by other researchers (Dutta, et al., 2001; Guedes, et al., 2003). Regarding iron leaching, it is noteworthy that, once again, the concentrations reached are always small (below 0.4 mg/L for both samples).

Finally, it is important to remark that we have performed some tests to evaluate the stability of the most promising catalyst (i.e., with low iron leaching and good catalytic performance): the Fe (II) oxalate 17.0 sample. OII degradation and TOC reduction were evaluated in 4 consecutive runs, being the catalyst recovered by filtration of the final effluent. Even though a slight activity decay was observed, which might be due to the iron loss (ca. 1.5% per cycle that represents a final concentration smaller than 0.3 mg/L, data not shown), OII conversion decreases only from 95.8 to 90.3% in 4 cycles, i.e., 16 h of operation. Regarding TOC reduction, in the 4 cycles final values were: 81.6, 81.4, 78.5 and 77.1%. In practice, this small deactivation could be compensated, if required, by adapting the reaction conditions (for instance slightly increasing the temperature along time).

**Conclusions**

- Twelve supported Fe-saponite catalysts have been prepared, by means of the incipient wet impregnation method, using a pillared clay support and four salts of Fe precursors at different Fe loads. The characterization of the
catalysts shows that the decomposition of the precursors gives rise to solids that present laminar structure, with active phases of Fe highly dispersed on the support, and high specific surfaces (in most of the cases with values comprised between 130 and 170 m$^2$/g), characteristics that make them potentially good catalysts for oxidation in the Fenton-like process.

- All the catalysts revealed to be quite active in the Fenton-like oxidation of Orange II, once clay concentrations used are much below than those usually found in the literature (typically around 1 g/L).
- The effects of hydrogen peroxide concentration, temperature and pH of the reaction medium were analysed in the present work. The obtained results show a high degradation of OII and of the intermediary oxidised compounds. At optimal conditions, 99% discoloration and 91% of mineralization were reached (after 4 h of reaction), using the catalyst prepared from Fe(II) oxalate with 17.0 wt% of Fe and in the following reaction conditions: T = 70°C, pH = 3.0 and $C_{H_2O_2} = 6$ mM. However, good performances with high selectivities to CO$_2$ and H$_2$O were also reached at significantly lower temperatures (30 °C).
- All the catalysts exhibit not only good catalytic activity but also a reasonable small iron leaching (below the EU directives values), indicating that the active phases are strongly fixed to the support (possibly iron strongly bonded to the aluminium pillars or engaged in small oxide clusters dispersed in the solid, inside or outside the porosity). This characteristic makes possible the Fe-saponite catalysts to have long-term stability, without generating iron hydroxide sludges.

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