Pillared clays from natural resources as catalysts for catalytic wet peroxide oxidation: Characterization and kinetic insights

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ABSTRACT
Pillared clays with Zr and Fe/Cu/Zr polycations have been prepared from natural clays found in large deposits of Kazakhstan and assessed as catalysts for the catalytic wet peroxide oxidation (CWPO), using 4-nitrophenol (4-NP) as model compound. The performance of the catalysts was followed by measuring the concentration of 4-NP, H₂O₂ and the total organic carbon (TOC), considering C_{4-NP} = 5 g L⁻¹, C_{H₂O₂} = 17.8 g L⁻¹, C_{cat} = 2.5 g L⁻¹, initial pH = 3.0 and T = 50°C. At those selected conditions, the pillared clays showed higher activity than natural clays in the CWPO of 4-NP. The conversion of the model pollutant was complete when Fe/Cu/Zr-PILCs were used, with the TOC removal reaching 78.4% after 24 h with the best Fe/Cu/Zr-PILC. The H₂O₂, 4-NP and TOC time-evolution was well described by a kinetic model based on TOC lumps in three blocks, considering the initial TOC (corresponding to 4-NP), the production of oxidizable intermediates and the formation of refractory products.

Keywords: Advanced oxidation process, Clay-based catalyst, CWPO, Kinetic modeling, Wastewater treatment, 4-nitrophenol

1. Introduction
In modern life, especially in productive activities, water plays an important role. There are about 39 thousand rivers and temporary streams in Kazakhstan. Among them, over 7 thousand are more than 10 km long. Most of the rivers in Kazakhstan belong to the internal closed drainage basins of the Caspian and Aral Seas and Balkhash, Alakol and Tengiz Lakes [1]. The majority of industries are situated around these rivers and lakes in order to use the water and to enable the discharge of some effluents for the aquatics systems. However, some of those streams can be harmful, making nowadays the contamination one of the primary issues of environmental authorities, requiring solutions in order to reduce its impact by using effective and inexpensive wastewater treatment methods, such as advanced oxidation processes (AOPs).

Nowadays, the scientific community shows an increasing interest in the development of AOPs (e.g., Fenton process, photo catalytic oxidation and electro catalytic oxidation) for the removal of recalcitrant and non-biodegradable organic compounds from aqueous streams. AOP are particularly useful technologies to treat recalcitrant or non-degradable compounds present in wastewaters in a wide range of concentrations (0.1-10 g/L) [2,3], difficult to remove by the conventional biological processes. The Fenton process consists on the use of iron and hydrogen peroxide, commonly known as Fenton’s reagent (H₂O₂ + Fe²⁺), for the generation of hydroxyl radicals (E°ₐₒ = 2.80 V), highly oxidizing species capable of degrading the organic matter present in wastewater effluents. The loss of the homogeneous catalyst is a major drawback that limits in practice the application of the Fenton process [4], since there is the necessity of a continuous feeding of iron and the corresponding additional separation processes and sludge disposal. In addition, although the Fenton process can completely destroy organic pollutants to harmless products, it requires a tight range of pH to operate (typically between 2 and 3, depending on the effluent [5]), and the need...
The catalytic activity of the prepared PILCs is compared to the per sulphate and ferrous sulphate as pillaring cation precursors. Karatau (Kazakhstan) deposits, using zirconium tetrachloride, cop-
PILCs prepared from natural clays extracted from Akzhar and water) considering monometallic (Zr) and trimetallic (Fe/Cu/Zr) 4-NP in aqueous reaction media (used as model synthetic wast-
tion mechanism and kinetics are of highly importance. as model compound in AOP studies and the knowledge of its degrada-
to carry oxygen to tissues and organs [19,20]. Thus, the use of 4-NP can develop a blood disorder which reduces the ability of the blood
previous work shows that the intercalation of different pairs of cations on natural clays, as zirconium (Zr), copper (Cu), iron (Fe), leads to improved catalytic activity for the oxidation of organic compounds, mainly tested with phenol, extensively used as model pollutant [10]. In CWPO, PILCs have been used as catalysts, taking advantage of its porous structure to adsorb organic compounds on its surface, which actively participates in the oxidation process in the presence of hydrogen peroxide. The use of 4-nitrophenol (4-NP) as model compound to assess the performance of catalytic systems in AOPs has been less explored than phenol, the mecha-
nistic insights lacking consolidated data. However, few reports in CWPO [11-14], photo oxidation by UV/H2O2 [15] and, in previous works on our group with CWPO reduced graphene oxide materials [16], magnetic carbon xerogels [17] and carbon nanotubes [2], reveal that the main reaction intermediates are hydroquinone, benzoguai-
none, catechol and several low molecular weight carboxylic acids (e.g., malonic, malic, maleic and acetic acids). To the best of our knowledge, there are no works related to the development of kinetic models for the CWPO of 4-NP and nowadays most of the studies regarding CWPO processes present only pseudo-first order rate equations describing the disappearance of a target pollutant [18].
4-NP is a toxic and bio-refractory compound that can damage the central nervous system, liver, kidney and blood of humans and other living beings. It has been shown in the literature that 4-NP can develop a blood disorder which reduces the ability of the blood to carry oxygen to tissues and organs [19,20]. Thus, the use of 4-NP as model compound in AOP studies and the knowledge of its degrada-
tion mechanism and kinetics are of highly importance.

In this study, we report the results obtained in the CWPO of 4-NP in aqueous reaction media (used as model synthetic wast-
water) considering monometallic (Zr) and trinodallic (Fe/Cu/Zr) PILCs prepared from natural clays extracted from Akzhar and Karatau (Kazakhstan) deposits, using zirconium tetrachloride, copper sulphate and ferrous sulphate as pillaring cation precursors. The catalytic activity of the prepared PILCs is compared to the extracted natural clays and a kinetic model is developed for the CWPO of 4-NP.

2. Materials and Methods

2.1. Reagents and Chemicals

Clays were obtained from the Karatau and Akzhar deposits (located in the Zhambyl region of south of Kazakhstan). Zirconium tetra-
chloride, reactor grade (99.5 %), was supplied from Alfa Aesar, and ferrous sulphate heptahydrate (99.5%) and copper sulphate pentahydrate (99.9%) were obtained from Skat reagents. Hydrogen peroxide (30% w/v), used as oxidant in the treatment of the synthetic wastewater, was purchased from Fluka. Titanium (IV) oxysulphate (TiOSO4, 15 wt.% in dilute sulphuric acid, 99.99%), hydrochloric acid (HCl, 37 wt.%), and sodium sulphite (Na2SO3, 98 wt.%) were purchased from Sigma-Aldrich. Sodium hydroxide (NaOH, 98 wt.%) was obtained from Panreac. 4-NP (98 wt.%) and 4-nitroace-
tochol (98 wt.%) were acquired from Acros Organics and Fluka, respectively. Methanol (HPLC grade), glacial acetic acid (analytical reagent grade) and acetoniitriile (HPLC grade) were obtained from Fisher Chemical. All chemicals were used as received without further purification. Distilled water was used throughout the work.

2.2. Preparation of Pillared Clays

Two natural clays extracted from Karatau (KNC) and Akzhar (ANC) deposits were used as raw materials to prepare the pillared clays. First, the extracted natural clays were extensively washed with HCl (2 M) at 50°C, in order to eliminate non-honed content of metal and other contaminants present inside the clays. Pillared clays were then prepared from the washed natural clays, using zirconium tetrachloride as a source of zirconium polycations, Ferrous sulphate and copper sulphate were used as source of iron and copper polycations. The pillaring solution was prepared by slow addition of NaOH (0.2 M) to the solution containing the polycation precursors at room temperature until pH = 2.8 were obtained. The resultant solution was aged for 24 h at room temperature. The clay pillaring process kept a ratio of 10 mmol of total metal per gram of washed clay, irrespective Zr-PILCs or Fe/Cu/Zr-PILCs were prepared. The final materials were dried at 350 K for 24 h and calcined during 2 h at 823 K considering a heating rate of 275 K min⁻¹.

2.3. Characterization Methods

The physico-chemical characteristics of the natural and pillared clays were determined by X-ray spectral analysis performed with a spectrometer Inca Energy from Oxford instruments, using an electron microprobe (EMP) of the brand Superprobe 733 from JEOL. X-ray diffraction analysis was performed on an automated diffrac-
tometer DRON-3 with CuKa - radiation, b-filter. A semi-quantitative basis was carried out on the diffraction patterns of powder samples using the method of equal weights and artificial mixtures. The dif-
fraction patterns were interpreted using ICDD data: a base of powder -filter. A semiquantitative
energy dispersive spectrometer (EDS). Transmission electron microscopy (TEM) was performed in a LEO 906E instrument operating at 120 kV, equipped with a 4Mpixel 28 × 28 mm CCD camera from TRS.

The pH of point of zero charge ($pH_{PZC}$) was determined by pH drift tests, following the procedure described elsewhere [21,22]. Briefly, five NaCl (0.01 M) solutions were prepared as electrolyte and initial pH was adjusted to 2, 4, 6, 8 and 10 using HCl and NaOH 0.1 M solutions. Then, samples of 0.05 g of pillared clays were loaded in each 20 mL of the previous solutions. The equilibrium pH of each suspension was considered after 48 h under stirring (320 rpm) at room temperature and, hence, the pH was measured (final pH). The $pH_{PZC}$ value was determined by intercepting the curve ‘final pH vs initial pH’ with the straight line ‘initial pH = initial pH’.

Fourier Transform Infrared Spectroscopy (FTIR) spectra of natural and PILCs were obtained with a FTIR instrument (Infraspek, Model FSM 1201, Russia, St-Petersburg) with a resolution of 1 cm$^{-1}$ and scan range of 7,800 to 400 cm$^{-1}$ using a sample based on 1% of clay with KBr.

2.4. Catalytic Runs

The CWPO of 4-NP in aqueous medium was carried out in a 250 mL well-stirred glass reactor maintained at 323 K. The reactor was loaded with 100 mL of a concentrated (5.0 g/L) 4-NP aqueous solution, considered as a model system to simulate high-loaded wastewaters [2,23–25]. The initial pH of the solution was adjusted to 3 by adding H$_2$SO$_4$ and NaOH solutions (not buffered). The stoichiometric quantity of hydrogen peroxide needed for complete mineralization of 4-NP was then added. The catalyst was loaded (2.5 g/L) after homogenization of the resulting solution, that moment being considered as $t_0 = 0$ h. All experiments were carried out for 24 h. Several samples were withdrawn from the medium of reaction at previously selected time points to follow the course of the 4-NP conversion and the appearance of intermediate compounds, measured by high-performance liquid chromatography (HPLC). For that purpose, a Jasco HPLC system equipped with a UV-Vis detector (UV-2075 Plus), a quaternary gradient pump (PU-2089 Plus) for solvent delivery (1 mL min$^{-1}$) and a Kromasil 100-5-C18 column (15 cm × 4.6 mm; 5 μm particle size; reversed-phase) was employed. Total Organic Carbon (TOC) and H$_2$O$_2$ concentrations were also measured during each run, using respectively a Shimadzu TOC-L CSN analyzer and a colorimetric method based on TiOSO$_4$ [16].

2.5. Kinetic Modelling

The kinetic modelling of the CWPO results obtained with the natural clays Karatau and Akzhar and with the PILCs Fe-Cu-Zr-Karatau and Fe-Cu-Zr-Akzh has been performed to describe the time-course evolution of 4-NP (used as target pollutant), H$_2$O$_2$ and TOC concentrations, following the procedures described elsewhere [18,22]. The rate of disappearance of each given compound (i) in the reaction medium, $r_i$ (mol g$^{-1}$ s$^{-1}$), is given by Eq. (1):

$$-r_i = \frac{1}{C_{cat}} \frac{dC_i}{dt}$$

where $C_{cat}$ refers to the catalyst concentration (g L$^{-1}$), $C_i$ to the concentration of compound i, such as H$_2$O$_2$, 4-NP or TOC (mol L$^{-1}$) and $t$ representing the time of reaction (h). The rate of disappearance $r_i$ can be expressed as a function of the concentration of the species involved in the reaction.

The numerical integration of the rate equations was solved by using the Microsoft Excel Solver (Microsoft Office 2010, Microsoft Corp.) for least-squares minimization. The models were also evaluated by the sum of square residual (SSR) and determination factor ($R^2$), calculated by applying Eq. (2) and Eq. (3), respectively:

$$SSR = \sum_{n=1}^{N} (C_{exp,i,n} - C_{model,i,n})^2$$

$$R^2 = 1 - \frac{\sum_{n=1}^{N} (C_{model,i,n} - C_{model,i})^2}{\sum_{n=1}^{N} (C_{exp,i,n} - C_{model,i,n})^2}$$

3. Results and Discussion

3.1. Characterization of the Natural Clays and PILCs

3.1.1. X-Ray diffraction

The powder X-ray diffraction patterns of the natural clays are shown in Fig. 1. In the diffractogram of the natural clays, the peaks at $2\theta = 6.2^\circ$ (d$_{001}$=14.19 Å) and $2\theta = 19.9^\circ$ (d = 4.5 Å) represents the characteristic reflection of montmorillonite (represented as M in Fig. 1), as reported by Is Fatimah [26]. The diffraction peaks at (20) 20.85°, 26.6°, 36.5º, 39.5º and 50.1º revealed the presence of quartz (as Q in Fig. 1), an impurity in the natural clays [13,26]. The reflections at (20) 23.01°, 29.5° and 36.0° was due to calcite (represented as C in Fig. 1) [27].
A semi-quantitative analysis was carried out from the XRD spectra in order to determine the composition of the natural clays in quartz, muscovite, feldspar, calcite, hydrated aluminum silicate and Kaolinite was determined (Table S1). As can be observed, these natural clays contain a high quantity of impurities in form of quartz and calcite (56.7 and 45.9% for ANC and KNC, respectively). The content of quartz on raw natural clays can be different depending on the deposit of the clay. Djeffal et al. [28] reported the use of natural clay with 53.86% in quartz in the CWPO of phenol. The calcite presented in the raw clays is normally replaced during cation exchange in the pillaring process, as done in this work.

3.1.2. EMP

The chemical composition of the natural and pillared clays is shown in Table 1, as determined by Inca Energy with a dispersive spectrometer from Oxford Instruments. As observed, the natural clays used in this work are rich in iron (7.9-9.9%), which can play an important role in the decomposition of hydrogen peroxide to produce hydroxyl radicals for the oxidation of pollutants in CWPO. The analysis on the composition of the pillared clays indicates the successful enrichment of pillaring Zr cations in Zr-KPILC and Zr-APILC, by exchange with Ca cations. In Akzhar pillared clays the quantity of Zr is 36.3% and in Karatau pillared clay it is 35.1%. As expected, the content of Fe on the Fe/Cu/Zr-PILCs increases in the modified clays in comparison with the natural clays, suggesting the exchange and fixation of the intercalating metals in the interlayer space. This incorporation is particularly noticed on Fe/Cu/Zr-APILC, with the presence of 22.8% of Fe in the Fe/Cu/Zr-APILC, against 7.9% in the corresponding natural clay (ANC), whereas 11.5% of Fe was reached in the Fe/Cu/Zr-KPILC, against 9.9%. However, in the Fe/Cu/Zr-KPILC, the increment of Cu and Zr content was higher, reaching 9.6 and 5.1%, respectively; whereas 6.1 and 0.9% were achieved in the Fe/Cu/Zr-APILC. The total weight percentage in the three metals for Fe/Cu/Zr-APILC and Fe/Cu/Zr-KPILC resulted in 26.2 and 29.8%, respectively. It is also evidenced that the solids modified with Fe/Cu/Zr have lower Si/Al ratios (Si/Al < 3.7) than in the Zr-PILCs (Si/Al > 4.1), meaning that oxides get preferentially stabilized in the interlayer space of the clay, following the targeted cationic exchange mechanism [29]. Several works [13,14,30-32] dealing with different natural clays, modified with Al, Fe and Cu metal cations to produce pillared interlayered catalysts (Al/Fe, Al/Cu and Al/Cu/Fe), show to incre-

3.1.3. Fourier transforms infrared spectroscopy

The FTIR spectra of the ANC and of the APILCS are shown in Fig. 2. The general shape of all spectra is similar, indicating that the layered crystal structure of the natural clay is not strongly affected by the pillaring process. Peaks between the FTIR spectra of the pillared samples and of the natural clay suggest its successful modification.

The broad band at 3,444 cm$^{-1}$ is due to bending and stretching vibrations of H$_2$O and the band at 1,631 cm$^{-1}$ is due to an overtone of the bending vibration of water [33]. The band at 1,046 cm$^{-1}$ is attributed to the stretching vibration of the Si-O bond, while the band corresponding to the bending vibration was observed at 875 cm$^{-1}$. The bending vibration bands of the Si-O-Si and Si-O bonds indicate that the natural clay holds two types of layered structures, one related to plane layer structure, and the other related to the interlayer structure with the appearance of the Si-O band at 536 cm$^{-1}$.

The absorption bands at 3,622, 2,922 and 2,852 cm$^{-1}$ can be ascribed to –OH groups, typically found in other clays [33,34]. The presence of calcite in the raw material can be observed by the band found at 1,431 cm$^{-1}$ [27]. This corroborates the presence of calcite in the natural clays observed by XRD analysis. As can be observed, the carbonate band significantly decreases after the pillaring process, likely due to exchange of the ions.

Table 1. Chemical Composition of the Natural and Pillared Clays, Determined by Elemental Analysis

| Clay          | Na  | Mg  | Al  | Si  | K   | Ca  | Ti  | Mn  | Fe  | Cu  | Zr  |
|--------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| KNC          | 1.7 | 4.8 | 13.9| 44.8| 4.8 | 16.2| 3.4 | 0.5 | 9.9 | n.i.*| n.i.*|
| ANC          | 1.8 | 4.9 | 13.2| 48.4| 4.8 | 18.3| 0.7 | n.i.*| 7.9 | n.i.*| n.i.*|
| Zr-APILC     | 2.0 | 2.5 | 9.8 | 40.5| 4.8 | 0.8 | 0.5 | n.i.*| 2.8 | n.i.*| 36.3|
| Zr-KPILC     | 2.5 | 2.5 | 10.0| 41.8| 4.3 | 0.9 | 0.3 | n.i.*| 2.7 | n.i.*| 35.0|
| Fe/Cu/Zr-KPILC | 3.7 | 3.3 | 13.4| 47.5| 3.9 | 1.3 | 0.7 | n.i.*| 11.5| 9.6 | 5.1 |
| Fe/Cu/Zr-APILC | 1.9 | 2.6 | 11.9| 46.4| 5.6 | 1.2 | 0.6 | n.i.*| 22.8| 6.1 | 0.9 |

*n.i. = not identified

Fig. 2. FTIR spectra of ANC and APILCs.
3.1.4. Microscopy

The surface morphologies of the natural clays and of the PILCs prepared were observed by SEM and by TEM. In this work, SEM proves to be ideally suited for studying the configuration, texture of clay samples [35]. The corresponding SEM micrographs are shown in Fig. S1. Generally, the images of natural clays show layered and smooth surfaces. However, ANC and KNC (Fig. S1(A)-(B)) are not so smoother when compared to other non-pillared clays, such as the bentonite [33,36-40], whose surface is modified after pillarization showing a rougher surface and more porosity. The samples prepared by pillarization of ANC and KNC show a slight modification regarding the surface (slightly darker and rougher) and it is possible to observe higher particles when compared to the natural precursors.

The micrographs obtained by TEM analysis, in dark field mode, with natural and pillared clays are shown in Fig. 3. On the TEM images it is possible to observe characteristics of the materials, such as the stress, its crystallization, morphology and even its holography. On the TEM images of pillared clays shown in dark field mode the defects are evidenced, as well as fine particles present in the material, visible as dark-colored particles. These particles correspond to external aggregates of Zr which were used to modify the natural clays. This observation means that the impregnation of Zr on the natural clays also takes place along with its pillarization, likely due to an excess of the Zr precursor used. The phenomena has been also observed by other authors [41].

Fig. 3. TEM micrographs, in dark field mode, for (a-b) natural clays, (c-d) Zr-PILCs.
3. the Zirconia particles were highly dispersed on the natural clays, with some being anchored inside the natural materials. The Zr impregnation of the clays with high dispersion confers more active sites for the CWPO of 4-NP on the developed materials.

3.1.5. pHpzc determination
The pH at which the net total particle charge is zero is called the point of zero charge (pHpzc, or pHpzc), which is one of the most important parameters used to describe variable charge surfaces. The pHpzc of the different natural clays and PILCs used in this work are shown in Table S2. The results reveal that ANC and KNC have clearly basic character, since the values of pHpzc obtained for those clays were 9.3 and 9.1, respectively. However, the intercalation procedures used leads to a decrease of the basic character of the clays, resulting in pHpzc values lower than 6.3 for all PILC materials. Concretely, the intercalation of Zr species produced a shift of pHpzc from 9.3 (ANC) to 6.3 (Zr-APILC), and from 9.1 (KNC) to 6.1 (Zr-KPILC), for Akzhar and Karatau clays, respectively. Regarding the trimetallic Fe/Cu/Zr-PILCs, similar pHpzc decreases were observed, shifts to 6.2 and 6.0 being obtained for Fe/Cu/Zr-Akzhar and Fe/Cu/Zr-Karatau pillared clays, respectively. These values are in good agreement with the pHpzc values of Zr based pillared clays reported elsewhere [42]. The surface chemistry of the prepared PILC materials may thus be characterized as having a nearly neutral character which, when in contact with acidic aqueous solutions, will make the surface become with a positive charge, thus favorable for the attraction of negative anions present in solution.

3.2. CWPO of 4-NP

3.2.1. Screening of catalysts
Our work aims to develop affordable and active catalysts based on natural and pillared clays modified with Zr and Fe/Cu/Zr for use in purification processes of wastewaters containing organic pollutants by CWPO. Fig. 4(a) shows the conversion of 4-NP as a function of reaction time, obtained with the two natural clays and the four pillared clays considered in this work. The pillared clays were found to have excellent catalytic properties in the oxidation of 4-NP. It should be noted that the initial natural clay is rather inactive when compared to the modified materials (pillared clays). While the natural clays achieved only 32% of 4-NP conversion after 8 h of reaction, the materials modified with metals revealed 100% of 4-NP conversion in just 4 h of reaction. The materials modified with Fe/Cu/Zr species show better results than the solids modified with Zr species. While the conversions observed with the clays modified only with Zr reaches 100% after 4 h of reaction, the Fe/Cu/Zr-KPILC and Fe/Cu/Zr-APILC show 100% conversion of 4-NP in only 2 h of reaction. In all the experiments carried out with the natural clays, almost complete oxidation of 4-NP could only be obtained after 24 h of reaction (98% conversion). The analysis of the results obtained clearly emphasizes that the pillared clays have much superior catalytic activity than the natural clays towards the removal of 4-NP by CWPO. Therefore, the importance on the modification of natural clays by Zr, Fe and Cu cations to increase its catalytic activity in the CWPO of 4-NP is placed in evidence.

Fig. 4. Evolution of (a) 4-NP and (b) H2O2 against time of reaction by CWPO with the natural and pillared clays (Operating conditions: C4-NP = 5 g L⁻¹, CH2O2 = 17.8 g L⁻¹, Ccat = 2.5 g L⁻¹, initial pH = 3.0 and T = 50°C).

Fig. 4(b) shows the results obtained for H2O2 decomposition. It is clearly seen that the inclusion of Fe and Cu metals in the pillared clays drastically increases the activity of the materials for 4-NP oxidation. Nevertheless, the rate of H2O2 decomposition appears fairly similar for both Zr-KPILC and Zr-APILC during the first 4 h of reaction.

The conversion of 4-NP reached with any of the prepared PILCs is higher when compared with other catalysts used in previous works at the same operating conditions, such as graphene oxide materials [16] and carbon nanotubes [2]. The CWPO of 4-NP using clays as catalyst has been scarcely explored. Ayodele et al. [12] achieved a 4-NP degradation of 100% after 10 min, but using higher concentration of the catalyst and hydrogen peroxide (4 g L⁻¹ of Cu-pillared synthetic bentonite with H2O2 in an excess of 20%
and light irradiation to assist the process) for much lower initial concentrations of 4-NP (75-100 ppm). Chirchi et al. [11] studied the degradation of similar initial concentrations of 4-NP (1 mM) in the presence of Fe- and Fe-Al-pillared synthetic bentonite (1 g L⁻¹), reaching complete conversions after 5 h. The prepared KPILC and APILC allow to completely remove high loads of 4-NP (5 g L⁻¹) in less time (2 h). The mineralization level obtained in the experiments was followed by measurements of the TOC conversion. The results obtained are represented in Fig. 5.

As expected, the removal of TOC is higher when pillared clays (conversion of TOC close to 70%) are used and compared to the natural clays. The highest TOC removal (65.1% after 8 h) was obtained with Fe/Cu/Zr-APILC. Under the same conditions, the natural clays presented only 7.0% of TOC conversion after 8 h. After 24 h (results not shown), the removals of TOC were 74.5 and 78.5% with Fe/Cu/Zr-KPILC and Fe/Cu/Zr-APILC, respectively.

TOC conversions are considerably lower than 4-NP conversions, owing to the formation of more refractory products (intermediate compounds). Fig. 5(b) shows the evolution of TOC removal in the experiment carried out with Fe/Cu/Zr-APILC, detailing the experimental TOC of the reaction media ($C_{\text{TTOC,4-NP}}$), theoretical TOC of 4-NP (determined from the concentration of 4-NP by HPLC and taking into account its TOC contribution, $C_{\text{TOC,4-NP}}$) and, finally, the resultant TOC obtained by the subtraction of the $C_{\text{TTOC,4-NP}}$ to $C_{\text{Tinitial,TOC}}$, which can be considered as the TOC of the oxidizable intermediates and final refractory products. As can be observed, after 4-NP oxidation, oxidizable compounds are produced, since the $C_{\text{TTOC,4-NP}}-C_{\text{TOC,4-NP}}$ curve show a maximum concentration close to 1 h of reaction time. Then, when all 4-NP is almost removed, the $C_{\text{TTOC,4-NP}}-C_{\text{TOC,4-NP}}$ still decreases, evidencing the occurrence of oxidizable intermediates. According to previous works [16], the first stage leads to aromatic intermediates (4-nitrocatechol, phenol, hydroquinone, resorcinol, catechol and benzoquinone). Then these intermediates are further oxidized to low molecular weight carboxylic acids (malic, maleic, malonic, oxalic, acetic and formic acids), conferring to the aqueous media solution more acidity (final pH was found to be less than the initial pH of 3).

3.2.2. Kinetic modeling

Based on the TOC contributions described above (4-NP, oxidizable intermediates and refractory products), it is possible to describe the evolution of TOC by lumping into three blocks (TOCA, TOCB and TOCC) to adjust the results to a simplified kinetic model (Eq. (4)-(6)).

$C_{\text{Tinitial,TOC}} = C_{\text{TOC,4-NP}} + C_{\text{TOC,B}} + C_{\text{TOC,C}}$ (4)

$\text{TOCA} + H_{2}O_{2} \rightarrow CO_{2} + \text{TOC}_B + H_{2}O$ (5)

$\text{TOCA} + H_{2}O_{2} \rightarrow \text{TOC}_C + H_{2}O$ (6)

TOCA represents the initial TOC and, thus, TOCA was assumed as the theoretical TOC of the 4-NP, $C_{\text{Tinitial,4-NP}}$ which is converted to CO₂ (justifying the decrease of TOC since the beginning of the CWPO process) and TOCB, corresponding to the lump of the oxidizable intermediates. Once formed, these compounds also evolve to CO₂ and to the final refractory products, TOCC. According with this, the following equations can be considered (Eq. (7)-(8)):

$\text{TOCA} = C_{\text{Tinitial,4-NP}}$ (7)

$\text{TOCA} + \text{TOCB} = C_{\text{Tfinal,TOC}} - C_{\text{Tinitial,4-NP}}$ (8)

Simple power-law rate equations, of first reaction order with respect to each reactant ($H_{2}O_{2}$, TOCA, TOCB and TOCC) have been checked for the prediction of $H_{2}O_{2}$ consumption and TOC removal,
the following initial conditions:

constant value relating the production of the refractory products \((t\leq 0)=0\) mol L\(^{-1}\)

for TOC, a good determination coefficient is observed when using the proposed kinetic model, which was fitted for all experimental data, is able to represent the data with accuracy (Fig. S2). The solution of these equations was solved by using numerical methods (ODE 45) implemented in the MATLAB environment. The numerical solution of these equations was solved by using numerical methods (ODE 45) implemented in the MATLAB environment. The numerical solution was performed by using the ode45 function in MATLAB. As observed previously, the pillaring process increases substantially the catalytic activity of the clays in the CWPO of 4-NP, compared to the parent natural clays. In fact, the formation of oxidizable intermediates is harder to oxidize compared to 4-NP.

The determination coefficients \((R^2)\) show a good agreement between the experimental and the simulated data for \(H_2O_2\) and 4-NP concentrations. For TOC, a good determination coefficient is obtained in the fitting of the experimental data obtained from Fe/Cu/Zr-PILCs. However, the regression of TOC does not show a good accuracy for the modelling of the data obtained with the natural clays (KNC and ANC). The results were maintained in order to compare the kinetic constant values obtained using the Fe/Cu/Zr-PILCs with the corresponding natural clays. Nevertheless, the proposed kinetic model, which was fitted for all experimental data, is able to represent the data with accuracy (Fig. S2). The validation of this model is also illustrated by the parity plots shown in Fig. S3.

The evolution of TOC lumps by three defined blocks \((TOC_A, TOC_B\) and \(TOC_C\)) with the Fe/Cu/Zr-PILCs is depicted in Fig. 6. As can be observed, the sum of the three contributions of TOC is perceptible since the beginning of reaction. Interestingly, the model predicts significant differences between the pillared clays for \(TOC_A\) and \(TOC_C\) profiles, since \(TOC_C\) production is faster with the Fe/Cu/Zr-APIILC compared to Fe/Cu/Zr-KPILC.

This is consequence of a faster oxidation of \(TOC_B\) (higher value of \(k_7\) with Fe/Cu/Zr-APIILC). Accordingly, \(TOC_B\) evolution with Fe/Cu/Zr-APIILC describes the maximum concentration value \((24\) min before than Fe/Cu/Zr-KPILC \((36\) min). This means that the Fe/Cu/Zr-APIILC is more active in the CWPO of 4-NP (the degradation of the pollutant is faster). However the concentration of \(TOC_B\) with this catalyst at 8 h is close to 0 mmol L\(^{-1}\) and all TOC of the aqueous medium is consequence of the refractory products.

Table 2. Kinetic Parameters Obtained in Modeling of CWPO of 4-NP with Natural and Pillared Clays

| Catalyst          | \(k_{H_2O_2}\) (L g\(^{-1}\) h\(^{-1}\)) | \(k_1\) (L g\(^{-1}\) h\(^{-1}\)) | \(k_2\) (L g\(^{-1}\) h\(^{-1}\)) | \(k_3\) (L g\(^{-1}\) h\(^{-1}\)) | \(k_4\) (L g\(^{-1}\) h\(^{-1}\)) |
|-------------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|
| KNC               | 0.062                           | 0.037                           | 0.024                           | 0.128                           | 0.128                           |
| ANC               | 0.038                           | 0.041                           | 0.031                           | 0.080                           | 0.080                           |
| Fe/Cu/Zr-KPILC    | 2.216                           | 2.803                           | 2.365                           | 0.568                           | 0.215                           |
| Fe/Cu/Zr-APIILC   | 2.501                           | 2.752                           | 2.750                           | 1.328                           | 0.484                           |
Fig. 6. Evolution of the simulated lumped TOC removal (symbols as experimental data of TOC and theoretical TOC contribution of 4-NP) in the CWPO of 4-NP with Fe/Cu/Zr-PILCs from (a) Karatau and (b) Akzhar (Operating conditions: \[ C_{\text{4-NP}} = 5 \text{ g L}^{-1}, \ C_{\text{CH}_2\text{O}_2} = 17.8 \text{ g L}^{-1}, \ C_{\text{cat}} = 2.5 \text{ g L}^{-1}, \text{ initial pH} = 3.0 \text{ and } T = 50^\circ\text{C}. \) 

\[ \text{TOC} = \text{TOC}_a + \text{TOC}_b + \text{TOC}_c \]

\[ C_{\text{TOC}_a}/C_{\text{TOC}_0} \]

\[ C_{\text{TOC}_b}/C_{\text{TOC}_0} \]

\[ C_{\text{TOC}_c}/C_{\text{TOC}_0} \]

\[ \text{TOC}_b + \text{TOC}_c \]

\[ \text{TOC}_c \]

\[ \text{TOC}_b \]

\[ \text{TOC}_a \]

36 min

24 min

Time of reaction (h)

4. Conclusions

Zr and Fe/Cu/Zr-PILCs were prepared from natural clays of Karatau and Akzhar. The catalysts obtained by the simultaneous incorporation of Fe, Cu and Zr cations are highly efficient in the oxidation of 4-NP in aqueous medium at very mild conditions (50ºC and atmospheric pressure). Pillared clay materials showed higher catalytic activity in the oxidation of 4-NP when compared to the natural clays. A complete removal of the contaminant is achieved after 4 h when Zr pillared clays are used, whereas the removal is reached after 2 h when Fe/Cu/Zr pillared clays are considered. The highest mineralization was obtained with the Fe/Cu/Zr-PILCs. By the subtraction of the theoretical organic carbon contribution of 4-NP to the measured TOC it is possible to observe the production of the oxidizable intermediates, since a maximum peak of concentration is observed. According to that, it is possible to propose a kinetic lumped model for TOC removal based in the initial TOC (corresponding to 4-NP), oxidizable intermediates and the refractory products. The kinetic model can be used to reproduce suitable the experimental data obtained in the CWPO of 4-NP with pillared clays.

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