Removal of recalcitrant organic compounds from an industrial complex effluent by heterogeneous Fenton-type treatment†
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Graphical abstract
Industrial chemi-mechanical pulping effluents are not affected by conventional treatment. The main parameters involved in Fenton-type treatment were determined. Factorial experiments were applied to evaluate the most favorable range of conditions for CuO/γ-Al2O3 application. The highest reduction of all parameters was obtained at the superior level of all variables, achieving reductions of COD and TOC between 40 and 50%.

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Abstract
Because of their chemical complexity, industrial chemi-mechanical pulping effluents are evaporated and burned, in spite of the high associated cost involved in these processes. The aim of this study was to remove recalcitrant compounds from this kind of wastewater using a Fenton-type treatment. The main parameters involved in the process and their influence on the results were determined. Homemade catalysts based on CuO, Fe₂O₃, NiO and ZnO, supported on γ-Al₂O₃ have been tested for catalytic oxidation, and the CuO/γ-Al₂O₃ catalysts showed the greatest effect on total organic carbon (TOC) reduction (52.7%). A series of two-level factorial experiments was subsequently applied to evaluate the most favorable range of conditions for CuO/γ-Al₂O₃ application. The studied variables were hydrogen peroxide concentration ([H₂O₂], g/L), active phase content (metal oxide supported on alumina, %), mass of catalyst (metal oxide/alumina system, g), and reaction temperature (°C). The highest reduction of all parameters was obtained at the superior level of all variables with CuO/γ-Al₂O₃, achieving reductions of chemical oxygen demand (COD) and TOC between 40 and 50%. Increasing catalyst mass did not produce additional benefit. This variable has a significant effect only on the reduction of aromatic compounds. At its low level, reduction in aromatic content exceeded 80%. Color reduction was influenced only by temperature (maximum reduction of 90%).

Abbreviations: AOP, advanced oxidation process; BAT, best available technology; BET, Brunauer, Emmet, and Teller, BOD, biochemical oxygen demand; CMP, chemi-mechanical pulping process; COD, chemical oxygen demand; EDS, energy dispersive X-ray; HYP, high yield pulping process; HPLC, high performance liquid chromatography; SD, standard deviation; SEM, scanning electron microscopy; tc, treatment combination; TOC, total organic carbon; XRD, X-ray diffraction

Keywords: Advanced oxidation, Chemi-mechanical pulping effluent, Factorial experimental design, Heterogeneous type Fenton reactions, Recalcitrant compounds

1. Introduction
In Fenton reactions of model compounds, the evolution of oxidation mechanism of the system and its intermediates is widely understood [1–6]. On the contrary, in the case of real industrial streams, the catalytic parameters and catalytic conditions of the treatment are barely known.

Generally, the untreated liquid streams that result from processing of lignocellulosic materials have high biochemical oxygen demand (BOD), chemical oxygen demand (COD), suspended solids (mainly fibers), fatty acids, tannins, resin acids, lignin and its derivatives. Some contaminants are natural and other are xenobiotic, i.e. formed during the pulp and paper process, and recalcitrant to degradation [7, 8]. In alkaline sulfite chemi-mechanical pulping processes (CMP), wastewater contains low molecular weight hemicelluloses, sugars, organic acids and several compounds that are recalcitrant to microbiological treatment, such as aromatic derivatives from wood extractives and labile lignin fractions. As CMP processes do not have a chemical recovery system, diluted spent liquors are generally treated in conventional effluent treatment systems, but
recalcitrant compounds are not affected and are discharged as they are [9--11]. A proper effluent treatment is therefore mandatory.

In these processes (as in many other cases), the chemical structure of wastewater is scarcely established, which makes the design of an effluent treatment system very difficult. Since this effluent is recalcitrant to biological degradation and it might be toxic to aquatic species [12, 13], conventional treatment systems are not enough and pulp and paper mills are forced to adopt technologically advanced treatments [14, 15].

Advanced oxidation processes (AOP) exploit the high oxidation potential of hydroxyl radicals, which attack most of organic molecules [16]. The high reactivity of these radicals and their low selectivity during the oxidation process are useful attributes that make these technologies a promising option for the treatment of effluents containing refractory compounds [14, 17].

In many cases, the objective of the oxidation process is not complete mineralization, with the subsequent conversion from organic carbon to carbon dioxide, but the conversion of pollutants into biodegradable substances that do not cause inhibition problems in the biomass of conventional biological treatment, nor cause ecotoxicity problems in the discharge. The application of one method over the other mainly depends on the contaminant concentration and flow rate of the effluent [18, 19].

The utilization of H\textsubscript{2}O\textsubscript{2} has emerged as a viable alternative among other oxidants, because it improves the efficiency of the oxidation, it is non-toxic and does not form any harmful by-products, and the reactions are carried out under non-critical conditions [20]. The process is performed at atmospheric pressure and low temperatures. Initial studies on AOP with H\textsubscript{2}O\textsubscript{2} were carried out using iron salts as homogeneous catalysts in the reactions known as “Fenton reactions”. The main disadvantage of homogeneous catalysts is the difficulty of the post-reaction separation of the catalysts from the product stream. It is then necessary to implement recovery operations to retrieve the used catalyst with the consequent increase of operational costs. An alternative solution to this problem is the deposition of the active phase on a porous solid, which characteristics are comparable to those of homogeneous catalysts, but that can be recovered by simple separation steps for reuse in other processes. Different variants of catalytic systems that work with active ions on a solid support are in study. In the field of heterogeneous catalysis, such reactions are known as “Fenton type reactions” [21--23]. The Fenton reagent’s mechanism has not been fully explained because of the variety of complex compounds and intermediates that are formed, and their subsequent reactions. During the reaction with the solution of H\textsubscript{2}O\textsubscript{2} and salts of Fe(II), organic compounds are oxidized with a radical chain mechanism [24], where hydroxyl radicals are capable of rapidly attacking organic substrates [25]. The main parameters that govern this system are: the initial concentration of H\textsubscript{2}O\textsubscript{2}, the concentration of Fe(II) and the temperature of the reaction [26].

High yield pulping processes (HYP) which combine chemical action prior to mechanical action are CMP. The unknown chemical structure and complexity of some contaminants in high yield pulping effluents represent a challenge. The aim of this study was to remove recalcitrant compounds from a CMP industrial effluent using a Fenton type treatment. The main parameters involved in the process and their influence on the results were
determined.
The first part of the work consisted of selecting efficient and stable metal oxides to eliminate recalcitrant substances. To select a particular catalytic system, the behavior of different catalysts consisting of metal oxides supported on alumina was evaluated, analyzing the oxidation of recalcitrant compounds in a Fenton type system. No reports on the treatment of HYP effluents treated by catalytic processes such as Fenton-type reactions have been found. Because of their chemical complexity, “best available technologies” (BAT) practice consists of evaporation and burning, in spite of the high associated cost involved in these processes. An experimental design consisting in a series of two-level factorial experiments was applied to evaluate the most favorable range of conditions for the oxidation of recalcitrant compounds.

**Materials and methods**

**2.1 Black liquor**
The spent liquor came from Papel Prensa S.A. integrated mill, located in San Pedro, Argentina, which produces 132,000 tons/year of soda-sulfite chemi-mechanical pulps from a mixture of willows, poplar and eucalyptus, and newspaper. In alkaline sulfite CMP wood chips are impregnated with sodium hydroxide and sodium sulfite (pH 9–10), and they are subsequently treated mechanically in a refiner. Chemical treatment is slight and lignin is not removed, but only softened. This type of pulping allows the use of low-density hardwoods as a fiber raw material. The actual effluent comprises several streams: 1) liquid waste from deinking plant; 2) water from wood treatment; 3) black liquor from the chemical treatment of wood (pH 7–8, deep red color, mainly composed by extractive and lignin derived from the product of the chemical reaction between wood and pulping liquor); 4) effluent from chemi-mechanical pulp washing; 5) effluent from the bleaching plant; and 6) white water from the paper making section of the mill. As effluent 3) contains the highest polluting load, dilutions of liquor 3) were performed with a liquor/water ratio of 1:50 to simulate the final mill effluent. The liquor was stored in plastic containers and was used without filtration. Oxidative processes were applied to this simulated effluent. The main characteristics of the industrial spent liquor are shown in Table 1.

**2.2 Catalyst preparation and characterization**

Samples of several metal oxides supported on alumina were tested at 70°C. Pellets of γ-Al₂O₃ (Alfa SASOL) were used as support and were impregnated using aqueous solutions of Cu(NO₃) · 2.5 H₂O (Riedel-de Haën, pa), Ni(NO₃) · 2.6 H₂O (Baker, pa), Zn(SO₄) · 7 H₂O (Riedel-de Haën, pa), citrate Fe(III) (BioPack, pa) as precursors. The incipient wetness impregnation method comprises mixing the support with an aqueous solution containing an appropriate amount of salt so that, after calcination, the catalyst contains the required metal content. The volume of the solution that is prepared is equal or slightly smaller than the pore volume of the support. The maximum load is limited by the solubility of the precursor in the solution. Solids were air dried during 24 h, then oven dried at 120 °C, and finally calcined at 900 °C in air atmosphere.

To select the metal system, γ-Al₂O₃ (I) pellets of 2.5 mm in diameter and 190 m²/g of surface area (determined by the Brunauer, Emmet, and Teller method (BET) method) were used. The final charge of the
Four catalysts (CuO/Al₂O₃, NiO/Al₂O₃, ZnO/Al₂O₃ and Fe₂O₃/Al₂O₃) was 5% by weight. In addition, experiments were carried out using two commercial pellets catalyst: Haldor-Topsoe (52% CuO, 25% ZnO) and Engelhard (12.5% CuO, 0.3% NiO) for comparison.

To evaluate the factors that influence the process, γ-Al₂O₃ (II) pellets 1.8 mm in diameter and 200 m²/g of surface area (according to the manufacturer) were used. The final load of catalysts used in this stage was established according to the experimental design.

The support and the catalysts were characterized using the following techniques:

- Surface areas were calculated from nitrogen adsorption at −196 °C by using a Micromeritics FlowSorb II 2300.
- Oxide structures and cluster size of the synthesized metal oxides supported on alumina were determined by techniques of powder X-ray diffraction (powder XRD) using PANanalytical, X’Pert Pro equipped with Cu Kα radiation. The patterns were recorder over a range of 10° < 2θ < 70° and compared to the JCPDS files to confirm phase identities.
- The surface morphology of the catalysts was examined by means of a scanning electron microscopy (SEM).

The elemental composition of the catalysts was determined by energy dispersive (EDS) X-ray spectroscopy using an EDAX Genesis XM4-Sys 60 equipment, equipped with a EDAX multichannel analyzed model EDAM IV, sapphire detector Si(Li) and Be window, super ultra-thin, software EDAX Genesis version 5.11.

### 2.3 Experimental design

To determine the magnitude of the effect of each variable (factor) on the studied system, experiments were conducted according a 2⁴ factorial design. In this design, the number of combinations of treatments (tc) is given by 2ⁿ, where n corresponds to the number of the analyzed experimental variables. In this study, n = 4, therefore tc = 2⁴ = 16 treatments. The studied variables were hydrogen peroxide concentration [H₂O₂] in g/L, active phase content, that is, content of metal oxide supported on alumina (%), mass of catalyst, i.e. metal oxide/alumina system (g) and reaction temperature (°C). The analysis of each experiment was performed using coded levels (+1 and --1 values), which arise from the true values converted to independent unit values of the factors. The real and coded values of each level of the experimental factors are shown in Table 2.

The H₂O₂ dosage was based on the stoichiometric ratio with respect to COD. This was calculated assuming complete oxidation of COD to CO₂ and H₂O. The H₂O₂ stoichiometric concentration respect to COD value was 1.98 g/L. To set H₂O₂ levels, the feasibility of reducing the treatment cost was assessed, so the lowest (sub-stoichiometric I) and uppermost (upper-stoichiometric II) values of the experimental design correspond to H₂O₂/COD ratios of 0.9:1, and 1.2:1, respectively.

### 2.4 Catalytic activity

A 250 mL PYREX glass batch reactor with a glass stopper equipped with a condenser, a thermocouple, and pH meter were used. To minimize external mass transport effects, experiments were carried out with a high-
speed magnetic stirrer (1200 rpm). Testing was performed in contact with air at atmospheric pressure. The reaction volume was 100 mL (with a dilution of 1:50 such as initial values of COD and TOC of the diluted liquor were 931 and 433 mg/L, respectively). The experiments were conducted using $\gamma$-Al$_2$O$_3$. The H$_2$O$_2$/COD ratio for the selection of the metallic system analysis was fixed as 9.4:1. H$_2$O$_2$ in excess (18.6 g/L) was used to select the most suitable catalyst metal system, since the extent of recalcitrance of this effluent was not known. The oxidation reaction was carried out at 70°C and with 1 g catalyst in all assays. All catalytic reactions were developed without pH adjustment to record the natural tendency of each catalyst system, allowing to avoid pH control systems and to compare the activity of the different catalyst systems.

Once the catalyst was added to the liquid mixture in the reactor, and the system reached the reaction temperature, initial time was recorded ($t = 00$, corresponding to the sample M00). At this point the corresponding amount of H$_2$O$_2$ was added, initiating the oxidation reaction (time zero, $t = 0$, corresponding to the sample M0). 2.5 mL samples were taken at different time intervals such as 0, 15, 30, 60, 120, 180 and 240 min (each sample was labeled from M0 to M6 according to time intervals) and were kept in plastics vials and then analyzed. The total oxidation reaction time was 240 min (M6). The analyzed responses throughout the reaction were: percentage of TOC reduction (TOC % reduction), percentage of aromatic content reduction (aromatic % reduction), percentage of COD reduction (COD % reduction), percentage of color reduction (color % reduction) and H$_2$O$_2$ consumption.

TOC conversion in blank trials (Al$_2$O$_3$ + H$_2$O$_2$ + effluent; H$_2$O$_2$ + effluent; Al$_2$O$_3$ + effluent) was negligible.

2.5 Analytical Techniques

Solids were determined according to Tappi T629 [27]. Inorganic content was assessed by determining the ashes at 525°C according to Tappi T211 [28]. The color of the effluent was measured by spectroscopy, measuring the absorbance at 450 nm (TECHCOMP spectrometer). COD was measured following the technique SM 5220-B (Standard Methods for the Examination of Water and Wastewater) [29]. To assess oxidation, total organic carbon (TOC) was measured using a TOC analyzer (Shimadzu, TOC-VCPN model). The evolution of aromatic compounds and the acids present in liquors, and peroxide consumption during the reaction, were quantified by high performance liquid chromatography (HPLC) (Waters, Massachusetts, USA) using an Aminex-HPX87H column under the following conditions: 4 mM H$_2$SO$_4$ as eluent, flow rate of 0.6 mL/min, 35°C, and UV diode array detector (organic acids at 210 nm and aromatic compounds at 254 nm).

2.6 Statistical evaluation

All results were statistically analyzed using the Statgraphics software at 95% of confidence level.

3 Results and discussion

3.1 Characterization of the catalysts

The characterization results of the $\gamma$-alumina and tested catalysts are summarized in Table 3. Both the impregnation process and the thermal treatment induced a decrease in the surface areas [30], but despite the
calcination temperatures, the results show acceptable surface areas after calcination, which provides the ability to achieve high oxidation values in further catalyst experiments. Therefore, the key point in producing high-performance catalysts is to increase or maintain the surface area [31]. BET values from Table 3 also indicate that the use of γ-Al₂O₃ is very important to provide high surface areas [32]. The performance of metal oxides as catalyst supports depends on their crystalline structure and textural properties. Although porosity of the catalyst was not measured, it can be assumed that porosity was slightly affected by calcination. Mesoporous materials such as γ-Al₂O₃ are preferred because they offer the advantage of avoiding pore plugging [33]. Mesopores are the paths for reactants and products, rendering high dispersion of active phase [34] and increase the amount of accessible active sites, improving catalytic reactions [35].

X-ray diffraction analysis revealed the presence of the characteristic peaks of the γ-Al₂O₃ phase (see Fig. 1). No peaks of other metals-containing phases were detected, only the main peaks corresponding to the γ-alumina phase, which are 2θ = 66.7° (100), 46.1° (80), 37.4° (60), and 39.7° (30). This result shows that the support is thermally stable and no structural changes occurred during the preparation of the catalyst, at least measurable by this technique [36]. From the absence of diffraction lines associated with the studied metal oxides, it can also be deduced that the concentration of impregnated active phase is quite low and the formed particles are well dispersed oxides [37], preventing its detection by XRD, as reported for Fe₂O₃/Al₂O₃ [30], CuO/Al₂O₃ [38], NiO/Al₂O₃ [39], and ZnO/Al₂O₃ [40].

The dispersion of metal species on the surface of the catalysts was analyzed by SEM-EDAX. Figure 2 shows SEM pictures, EDS spectra and the quantitative analysis (active phase content) obtained for CuO/Al₂O₃(I), NiO/Al₂O₃(I), and ZnO/Al₂O₃(I) catalysts. Fe₂O₃/Al₂O₃(I) SEM and EDAX spectra are not shown, but the quantification gave 7.37% of Fe₂O₃. This difference of 5% can be attributed to experimental errors, mostly of the equipment of measure. For all catalyst systems analyzed by SEM, the supported particles are well dispersed and distributed homogeneously over the substrate. The composition of each catalyst via spectrometry and EDX was determined as the average of four replicas. Catalysts based on Zn and Fe presented sulfur and chlorine impurities, respectively. The presence of the impurities was due to precursors used in both cases, which were not fully eliminated by thermal treatment during calcination.

### 3.2 Selection of the metallic system

Catalytic activity tested on different systems is summarized in Fig. 3 (as TOC % reduction). The NiO/Al₂O₃ catalyst (reduction of 3.7%) showed a statistically similar behavior to that of the blank test (p < 0.05); i.e. which means that does not produce a significant effect on percentage of TOC reduction. These results differ from those observed by other authors [37, 41], who confirmed the high oxidizing power of supported nickel applied to several types of compounds and the strong interaction between the metal particles and the support. These differences could be explained by the low active phase content and low working temperature used in this work. The catalyst NiO/Al₂O₃, and Engelhard (39.6% of TOC reduction), showed significant differences in their behavior (p < 0.05), but it is presumed that the high conversion rate achieved by the bimetallic catalyst is not the result of nickel action. Haldor-Topsoe and ZnO/Al₂O₃ (47.1 and 38.7% of TOC reduction,
showed no significant differences in their behavior \((p < 0.05)\). Copper and zinc appear to have a synergistic effect. In addition, \(\text{CuO/Al}_2\text{O}_3\) and \(\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3\) catalysts presented a significant effect on TOC (52.7 and 30.10%, respectively). The oxidative action of \(\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3\) was not significantly different from \(\text{ZnO/Al}_2\text{O}_3\) \((p < 0.05)\). Therefore, \(\text{CuO/Al}_2\text{O}_3\), \(\text{ZnO/Al}_2\text{O}_3\) and \(\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3\) catalysts have the greatest effect on the oxidation of recalcitrant compounds, and consequently, the highest values on the percentages of TOC reduction. The maximum conversion values achieved were generally low. This may be due to the high value of the initial concentration of peroxide, since \(\text{H}_2\text{O}_2\) in excess can generate radical capturing compounds (scavenging effect), decreasing the oxidation efficiency [42].

Leaching of the active phase was not observed in any of the tests when working with an excess of \(\text{H}_2\text{O}_2\), possibly also due to the slight variation of pH of the solutions, which remained near neutrality [43, 44]. This trend can be explained by the formation of sodium carbonate, whereas the reaction proceeds because of the oxidation of liquor components. It should be noted that alumina used as support gives the catalyst certain chemical properties that influence pH, among other conditions of the reaction. The pH values of the oxidation reactions were close to 7.3, the value in the blank trials using only the support (alumina as catalyst). In cases when homemade catalyst was used, the pH was also close to this value (see Table 4), indicating that the pH of the reaction medium is also controlled by the support [45]. Acid and basic sites at the surface of \(\gamma-\text{Al}_2\text{O}_3\) behave exactly as acid and basic substances in solution, probably buffering the reactions with the alkaline liquor. Therefore, they respond reversibly to a change in the pH of the solution with the concomitant variation of the total surface charge [46]. Thus, it is possible that the higher catalytic activity of a catalyst based on copper is also based on the favorable pH of the reaction [47]. On the contrary, optimal pH for an iron based catalyst is between 3 and 4, which discourages their use for practical applications [48].

Copper oxide was selected as oxidation catalyst for the rest of the work, as it generated one of the highest percentages of mineralization. Moreover, the pH range of the reaction media (near neutral) is considered optimal for its use [49], which represents an additional advantage when working with alkaline liquors because pH control is not needed.

**3.3 Effect of temperature, hydrogen peroxide concentration, active phase content, and mass of catalyst**

The results of the analysis of the factorial design at 240 min of reaction (M6) are shown in Table 5.

The effect of factors is defined as the change in the studied response as a consequence of the change in factor level, from the lower to the upper level and vice versa. The effect of the studied factors (and their levels) on the percentage of TOC reduction, aromatic compounds reduction, and COD reduction can be seen in Fig. 4. Standard deviations (SD) for the means of TOC, aromatic compounds and COD were ±1.83, ±2.22, and ±1.28 respectively. The percentage of TOC reduction, aromatic compounds reduction, and COD reduction can be seen in Table 6.

Temperature was the most important factor for TOC reduction over the studied range \((p\text{-value}: 0.0002)\). The oxidation degree reached a maximum at the highest temperature, as shown in Fig. 4 (70°C). A potential solution to improve mineralization degree with the same dose of \(\text{H}_2\text{O}_2\) is to increase temperature, because it determines the radicals generation rate [50, 51], reducing reaction costs [52, 53]. The detrimental effect of
competitive reactions and scavenging, which appears when the ratio of oxidizing agent is not correct (for example, when hydrogen peroxide concentrations is above the optimum), is partly offset by the positive action of increasing temperature [26]. Temperature is also the most influential factor in COD and aromatic compounds reduction (p-value of 0), as shown in Fig. 4. The greatest percentages of reduction were achieved at the highest value of the variable (70°C). Discoloration of the effluent showed the same behavior (p-value: 0.0084).

Hydrogen peroxide concentration showed significant influence on TOC conversion rates (p-value: 0.0243), COD (p-value: 0.0336) and aromatic content (p-value: 0.0005). Even if the best oxidation rates were attained working with the highest H$_2$O$_2$ concentration, conversion rates were not as high as expected (Fig. 4). Peroxide did not have significant effect on discoloration of the effluent. This behavior is not in agreement with findings of other authors [54, 55], but H$_2$O$_2$ concentrations used in the present study were low.

Only the catalyst mass had a significant effect on the reduction of aromatic compounds (p-value: 0.0207). The highest reduction was achieved with 0.5 g of catalyst load (Fig. 4). Increasing catalyst mass within the studied range (0.5–1 g) did not result in an additional benefit in COD and TOC reduction, although active sites for H$_2$O$_2$ decomposition and for organic compounds adsorption are increased [56]. Significant reductions of TOC and COD were achieved by increasing active phase content (p-values: 0.0179 and 0.0002 for TOC and COD, respectively), suggesting that the active phase (catalyst after calcination) is not in the bulk of the support but on the surface (Fig. 4).

The interaction between active phase and temperature showed to be highly significant (p = 0.0183). A synergistic effect can be observed in TOC, COD, and aromatic reduction when both variables were at their uppermost level (2.5 % and 70°C). The interactions between [H$_2$O$_2$] and active phase (p =0.0194), and [H$_2$O$_2$] and catalyst mass (p = 0.0077) were only significant on percentage of COD reduction.

The significance of each factor over the response varied along the oxidation reaction, so this must be taken into account to select and optimize the catalytic system. The effect of each significant factor on the percentages of TOC and of aromatic compounds reductions is shown in Fig. 5 (non-significant factors and interactions are not shown). Figure 5 shows that maximum removals were produced at high levels of temperature, hydrogen peroxide, and active phase for TOC, whereas in the case of aromatic compounds, high levels of temperature and peroxide, and low level of catalyst mass were necessary.

According to Table 5, final pH remained almost constant despite the different tested conditions. Similar final pH values probably are an indication that there are similar distribution of intermediates and/or similar accumulation of carboxylic acids at the end of each oxidation reaction [57]. As the only identified species were acetic acid, oxalic acid, sulfur compounds, sodium carbonate and other salts [10, 11], probably these last ones buffered the oxidation reaction and avoid higher degree of mineralization. It is known that those compounds are recalcitrant and interfere the Fenton oxidation process [58, 59].

Since H$_2$O$_2$ consumption is critical in this catalytic process, the oxidation reactions were also evaluated using the “η parameter” (Table 5). This parameter, as defined by several authors [60], represents the reagent consumption considered as the amount of TOC (mg/L) and COD (mg/L) converted per unit of decomposed H$_2$O$_2$ (g/L). From the comparison of treatment conditions which differ only in the temperature of reaction, the
efficiency of \( \text{H}_2\text{O}_2 \) does not decrease when increasing temperature as should be expected if \( \text{H}_2\text{O}_2 \) is decomposed due to thermal instability. On the contrary, a higher temperature implies a faster, and more efficient cooper-catalyzed \( \text{H}_2\text{O}_2 \) conversion into radicals, enhancing mineralization. Other authors [54] also found that \( \text{H}_2\text{O}_2 \) conversion into hydroxyl radicals is accelerated as the temperature increases, and the amount of \( \text{H}_2\text{O}_2 \) available to scavenge these radicals is accordingly small, allowing a higher \( \text{H}_2\text{O}_2 \) consumption yield for a given conversion of TOC and COD.

The long-term stability of a catalyst is essential for industrial applications. A catalyst batch was retained and reused again at the same conditions (tc: 12 of the experimental design), without treatment between runs maintaining a similar alkaline liquor concentration. The solid was just recovered from the solution after the reaction, dried at room temperature and tested again. The TOC removal achieved and the \( \text{H}_2\text{O}_2 \) consumed were similar after five consecutive cycles, indicating that it is possible to reuse the catalyst without affecting its efficiency. The analysis of TOC reduction profiles in the repeated experiments showed a slight induction period which marginally lengthened the consecutive runs. Nevertheless, the results observed for the fifth reuse of the catalyst showed that it was still active and able to be used.

### 3.4 Effect of the initial concentration of hydrogen peroxide and active phase in conditions out of the design

Fenton systems using \( \text{Cu}^{2+} \) catalysts require a larger stoichiometric excess of \( \text{H}_2\text{O}_2 \), compared to conventional Fenton systems, to overcompensate the competitive scavenging by \( \text{O}_2 \) (this treatment needs to work under aerobic conditions) [49]. To verify if this is the case in the current study, hydrogen peroxide concentrations much higher than stoichiometric values were tried (Fig. 6). Under these conditions (large excess of oxidizing agent) differences in mean reductions of TOC and COD were 34 and 40%, respectively, of those obtained with upper-stoichiometric II concentration, respectively (\( p \)-value < 0.05), as shown in Fig. 6. The three studied \( \text{H}_2\text{O}_2 \) concentrations resulted in significant differences on the reduction of aromatic compounds (\( p \)-value < 0.05). As \( \text{H}_2\text{O}_2 \) concentration increases, the amount of hydroxyl radicals available for the oxidation of pollutants also increases and the removal of recalcitrant compounds increases as well. Nevertheless, the efficiency of a dose of \( \text{H}_2\text{O}_2 \) 80% higher than that of the initial trials was very low (only 15–20% of TOC reduction). It is possible that the excess of \( \text{H}_2\text{O}_2 \) has promoted scavengers generation (radical capture compounds), decreasing the oxidation efficiency [61]. Final TOC conversions were 37, 40, and 62% for peroxide concentrations corresponding to sub-stoichiometric I, upper-stoichiometric II, and in excess, respectively.

Increasing the active phase induces the oxidant to generate more free radicals [62]. A correct balance between the mass of catalyst and the supported metal oxide is essential to maximize the amount of radicals generated in these systems [63]. Oxidation reactions using 5% of active phase were carried out (in addition to the design), but there were no statistically significant differences in COD and aromatic reductions between the three amounts of active phase tried (Fig. 6). TOC conversion decreased by 39.2% with respect to the other amounts of active phase when 5% was used (\( p \)-value < 0.05). Even if it is generally assumed that the concentration of the active phase has a strong influence on free radicals generation and it has also a great
influence on the kinetics reaction of TOC reduction [64, 65], it was not the case in this study. The relative difference between aromatic compounds and TOC removal was higher when using 5% of active phase. The results suggest that an important amount of the organic intermediates remain in the reaction mixture [66]. In conventional Fenton processes, the organic matter competes with the metallic substance for the hydroxyl radicals. It is then assumed that with 5% of active phase, the TOC at the end of the catalytic reaction is low, as the ratio metal content/ oxidizing agent decreased [42, 67].

**Concluding Remarks**

Catalytic oxidation in the presence of hydrogen peroxide could be an efficient alternative to treat effluents from chemi-mechanical pulp processes.

It was found that NiO/Al₂O₃ catalyst was not active, since it did not produce a significant effect on the TOC reduction.

Catalysts based on Cu, Fe, and Zn showed the best results in terms of TOC reduction and aromatic reduction along the oxidation reaction. Copper did not generate active phase leaching when working with an excess of H₂O₂.

When working with the Cu catalyst, temperature showed to be the most important factor for TOC, COD, and aromatic reduction of the simulated effluent in the studied range, being the only factor that significantly affected the effluent discoloration. The highest values of oxidation and discoloration were achieved at the uppermost temperature level (70°C).

Hydrogen peroxide concentration significantly reduced TOC and COD, although the achieved conversion (40–50%) was lower than expected in its uppermost level (2.5 g/L). However, reduction of the aromatic content was near 80%. The inability of the catalyst to achieve higher oxidation of refractory compounds was not a consequence of the availability of oxidizing agent.

Catalyst mass affected only the reduction of aromatic compounds, showing its greatest effect at its lowest level (0.5 g).

The highest level of active phase tested in the experimental design (2.5%) produced maximum reductions of TOC and COD. Working with high percentages of active phase (5%) did not produce any improvement.

The authors have declared no conflict of interest.

**References**


**Figure 1:** XRD pattern. A) Fresh catalyst used in the selection of metal system. B) Fresh catalyst used in the experimental design.

**Figure 2:** SEM images (2000×) and EDS spectra, a) CuO/Al2O3(I), b) ZnO/Al2O3(I), c) NiO/Al2O3(I).

**Figure 3:** TOC reduction (%) for different catalytic systems. Blank trial I: Al2O3/effluent; blank trial II: Al2O3/H2O2/effluent; blank trial III: H2O2/effluent (blank trials are batch reactions without active phase or
hydrogen peroxide.

**Figure 4:** Effect of significant factors on TOC reduction (%), aromatic compounds reduction, and COD reduction.

**Figure 5:** Reduction (%) of: a) TOC, b) aromatic compounds, for 240 min of reaction.

**Figure 6:** Influence of peroxide concentration and active phase content on the TOC reduction (%), aromatic compounds reduction (%), and COD reduction (%).

### Table 1: Chemical characteristics of the spent liquor

| Total soluble solids (g/L) | 61.1 |
| pH | 7.4 |
| COD (mg/L) | 46 550 |
| TOC (mg/L) | 21 665 |
| Ashes at 525°C (% of total soluble solids) | 52.3 |
| Acetic acid (g/L) | 23.6 |
| Formic acid (g/L) | 0.3 |
| Propionic acid (g/L) | 0.6 |

### Table 2: Levels of each experimental factor in real and transformed values

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<thead>
<tr>
<th>Treatment (tc)</th>
<th>[H$_2$O$_2$] g/L</th>
<th>Active phase %</th>
<th>Mass g</th>
<th>Temperature ºC</th>
<th>[H$_2$O$_2$] Active phase</th>
<th>Mass g</th>
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### Table 3: Characterization Results of the Support and the Catalysts

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<tr>
<th>Catalyst</th>
<th>Metallic content (%)</th>
<th>Calcination temperature (ºC)</th>
<th>BET area (m²/g)</th>
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<tr>
<td>γ-Al₂O₃(I)</td>
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<tr>
<td>γ-Al₂O₃(II)</td>
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<td>--</td>
<td>200*</td>
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<tr>
<td>Haldor-Topsoe</td>
<td>52% CuO/25% ZnO</td>
<td>--</td>
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</tr>
<tr>
<td>Engelhard</td>
<td>12.5% CuO/0.3% NiO</td>
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<td>Fe₂O₃/Al₂O₃(I)</td>
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<td>CuO/Al₂O₃(II)</td>
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*According to the manufacturer

### Table 4: pH values and the end of catalytic reactions

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<td>ZnO/Al₂O₃(I)</td>
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*Remained almost constant during 240 min of reaction
Table 5: Variables values at the end of the reaction (240 min of reaction)

<table>
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<tr>
<th>Treatment (tc)</th>
<th>TOC (% reduction)</th>
<th>Aromatic (% reduction)</th>
<th>COD (% reduction)</th>
<th>Color</th>
<th>H$_2$O$_2$ (g/L)</th>
<th>pH*</th>
<th>TOC/H$_2$O$_2$</th>
<th>COD/H$_2$O$_2$</th>
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</table>

* Remained almost constant during 240 min of reaction

Table 6: Effect of significant factors and their levels on TOC reduction (%), aromatic compounds reduction (%), and COD reduction (%).

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<tr>
<th>Factor</th>
<th>Level</th>
<th>% of Reduction</th>
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<td></td>
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<tr>
<td>Temperature</td>
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<tr>
<td>H$_2$O$_2$</td>
<td>+</td>
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<tr>
<td>H$_2$O$_2$</td>
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<td>25</td>
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<tr>
<td>Active phase</td>
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<td>Active phase</td>
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<tr>
<td>Catalyst mass</td>
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<td>Catalyst mass</td>
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</tr>
</tbody>
</table>
Fig. 1
Fig. 2
Fig. 3

% of TOC Reduction

Catalysts tested:
- Engelhard
- Haldor-Topsoe
- Fe₂O₃/Al₂O₃
- CuO/Al₂O₃
- NiO/Al₂O₃
- zeo/Al₂O₃
- Fe₂O₃/Al₂O₃-Hom/Later
- Blank trial I
- Blank trial II
- Blank trial III
Fig. 4

![Graph showing % of Reduction for different factors: Temp (+), Temp (-), H2O2 (+), H2O2 (-), Active phase (+), Active phase (-), Catalyst mass (+), Catalyst mass (-).

The graph compares TOC, Aromatic compounds, and COD for each factor.

- Temp (+) with 70% reduction in TOC, 48% in Aromatic compounds, and 33% in COD.
- Temp (-) with 41% reduction in TOC, 37% in Aromatic compounds, and 33% in COD.
- H2O2 (+) with 52% reduction in TOC, 42% in Aromatic compounds, and 36% in COD.
- H2O2 (-) with 58% reduction in TOC, 57% in Aromatic compounds, and 32% in COD.
- Active phase (+) with 23% reduction in TOC, 24% in Aromatic compounds, and 23% in COD.
- Active phase (-) with 28% reduction in TOC, 29% in Aromatic compounds, and 33% in COD.
- Catalyst mass (+) with 42% reduction in TOC, 32% in Aromatic compounds, and 33% in COD.
- Catalyst mass (-) with 52% reduction in TOC, 23% in Aromatic compounds, and 23% in COD.
Fig. 5
Fig. 6