Photo-Fenton oxidation of phenol with magnetite as iron source

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Abstract

In this work, magnetite-catalyzed Fenton reaction was investigated under UVA irradiation for the degradation of phenol as model compound. Four kinds of magnetite were used having different particle size, surface area and FeII content. Different kinetic behaviors were observed, thereby underscoring the strong implications of surface and chemical properties of magnetite. The size and surface area of the particles seemed to be less important, while the FeII/FeIII ratio played some role. Despite the link between magnetite reactivity and its structural FeII content, light-induced reduction of FeIII to FeII was found necessary to promote the Fenton-based reactions. As surface FeII may be oxidised or otherwise unavailable, initial photoactivation may be needed to trigger the Fenton reactivity. Two major driving forces were highlighted that account for the photoactivity of magnetite at pH 3: (i) the formation of intermediates such as hydroquinone that are able to reduce FeIII to FeII, and (ii) the accumulation of dissolved Fe due to magnetite dissolution, both in dark and under irradiation. Very interestingly, the photo-Fenton degradation of phenol was also observed under neutral conditions. In this case, for two out of four samples, the degradation rates were quite near those found at pH 3, which is usually reported as the optimum pH value of the process. The magnetite ability to promote photo-Fenton reactions even under circumneutral pH conditions, the limited iron leaching and its easy magnetic separation makes magnetite a promising catalyst in wastewater treatment applications.

Keywords: photo-Fenton; magnetite; advanced oxidations process; heterogeneous photo-Fenton oxidation.
1. Introduction

The classic Fenton process involves aqueous ferrous ions and $\text{H}_2\text{O}_2$ that react together to form $\cdot\text{OH}$, in a reaction that can be expressed as follows (in acidic solution, Fe$^{\text{II}}$ is usually present as Fe$^{2+}$ and Fe$^{\text{III}}$ may be present as FeOH$^{2+}$) [1]:

$$\text{Fe}^{\text{II}} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{\text{III}} + \text{OH}^- + \cdot\text{OH} \quad (1)$$

Reaction (1) is a stoichiometric process, but Fe is often used in catalytic amount because of the subsequent pathways of $\text{H}_2\text{O}_2$ decomposition [2], which regenerate Fe$^{\text{II}}$ that reacts again in (1). The classical Haber-Weiss view of the process reads as follows [3]:

$$\text{FeOH}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{\text{II}} + \text{HO}_2\cdot + \text{H}_2\text{O} \quad (2)$$

$$\text{FeOH}^{2+} + \text{HO}_2\cdot \rightarrow \text{Fe}^{\text{II}} + \text{O}_2 + \text{H}_2\text{O} \quad (3)$$

The production of $\cdot\text{OH}$, a powerful reactant that is able to degrade a wide variety of xenobiotics at diffusion-limited reaction rates, accounts for the widespread use of the Fenton reaction among Advanced Oxidation Processes (AOPs) for the abatement of recalcitrant compounds in water and wastewater. Reaction (1) is the fast step of the Fenton process, while reactions (2,3) are considerably slower. Therefore, when Fe$^{\text{II}}$ and $\text{H}_2\text{O}_2$ are mixed in the presence of a pollutant, one often observes a very fast first step followed by a considerable slowing down of the reaction. This issue complicates the use of Fe$^{\text{III}}$, which is much cheaper than Fe$^{\text{II}}$ in the classic Fenton process. It also explains why several variants of the Fenton reaction have been proposed. An example is the photo-Fenton process, where the formation of Fe$^{\text{II}}$ from Fe$^{\text{III}}$ takes place photochemically (L in reaction (5) is an organic ligand such as oxalate) [4, 5]:

$$\text{FeOH}^{2+} + \text{hv} \rightarrow \text{Fe}^{\text{II}} + \cdot\text{OH} \quad (4)$$

$$\text{Fe}^{\text{III}}\cdot\text{L} + \text{hv} \rightarrow \text{Fe}^{\text{II}} + \text{L}^{\cdot\cdot} \quad (5)$$

The Fenton reaction is usually carried out in acidic conditions (optimum pH is 3), not only to keep Fe$^{\text{III}}$ dissolved but mostly because the $\cdot\text{OH}$ generation in reaction (1) has maximum effectiveness in an acidic environment. Furthermore, among the Fe$^{\text{III}}$ species, the FeOH$^{2+}$ hydroxo-complex that prevails at pH 3 has the highest reactivity towards reduction by $\text{H}_2\text{O}_2$ and $\text{HO}_2\cdot$ in reactions (2,3) [2, 3, 6]. The pH effect is a major issue where the still unresolved complexity of the Fenton mechanism comes to the surface, behind the apparent straightforwardness of reactions (1-5). Indeed, reaction (1) actually involves the formation of a highly oxidized Fe adduct, often indicated as ferryl (e.g. ferryl ion FeO$^{2+}$), which does not necessarily evolve into $\cdot\text{OH}$ [1, 7, 8]. The $\cdot\text{OH}$ formation is most effective (but by no means quantitative) at pH 3 and it usually decreases at higher or lower pH
values [2, 3]. Therefore, it is not surprising to find differences between the Fenton process and the expected reactivity of \( \bullet \text{OH} \) [1, 7, 9].

Recently, magnetite (Fe\( \text{II} \)Fe\( \text{III} \)O\(_4\), Fe\( \text{II} \)-Fe\( \text{III} \) mixed valence oxide) was successfully used as iron source in heterogeneous Fenton reactions, because Fe\( \text{II} \) plays an important role in the initiation of the Fenton process according to the classical Haber–Weiss mechanism. Magnetite, pristine, doped or coupled with other oxides (e.g. CeO\(_2\)) was shown to effectively catalyze the oxidative degradation of target compounds at circumneutral pH. It also exhibited good structural stability and excellent reusability [10-19]. Magnetite can be synthesized in the laboratory by various biotic and abiotic pathways. Abiotic procedures to form magnetite include co-precipitation of soluble Fe\( \text{II} \) and Fe\( \text{III} \) species, oxidation of hydroxylated Fe\( \text{II} \) species and ferric oxides transformation [20-24]. The morphology, crystallography and specific surface area of natural or synthetic magnetite can vary widely [25, 26]. Similarly to other iron oxides, magnetite exists as micrometric and nanometric particles in many natural and engineered environments. Because the specific surface area of nanosized particles is very large, their surface reactivity is exalted and they can play a potentially pre-eminent role in sorption and/or redox processes.

The heterogeneous photo-Fenton reaction can solve the problem of eliminating and re-using Fe from the reaction system at the end of the process, but the separation of the solid phase is still an open issue. The separation problem is even more important in the case of oxide nanoparticles, which are potentially more reactive because of the favorable surface-to-volume ratio. From this point of view, the fact that magnetite undergoes very easy magnetic separation from aqueous systems makes it a very interesting material to be tested for photo-Fenton reactivity. Moreover, it is very interesting to check whether, as for dark reactivity, the photo-activity of magnetite is maintained under circumneutral pH conditions, which would overcome the need of pH adjustment. At circumneutral pH the operational mechanism of the photo-Fenton process is still controversial, in particular in the presence of organic ligands for iron (EDTA, humic acids…) [27, 28]. Quite surprisingly, very few data are available about the use of magnetite in photo-Fenton chemistry. To fill in this knowledge gap, the present work has the goal of studying the photo-Fenton reactivity of magnetite toward the degradation of phenol. The latter was chosen because it is a substrate of well known behavior and it can be very helpful in the elucidation of reaction pathways [29].

The photo-Fenton reactivity of magnetite could depend on characteristics and surface properties such as crystallinity, surface area, Fe\( \text{II} \) content or Fe\( \text{II} \)/Fe\( \text{III} \) structural ratio. For this reason, dark and irradiation experiments were carried out with four different kinds of magnetite (two synthetic and two commercial) having different morphologies and structural properties. The aim was to point out the effects of particle size, surface area and Fe speciation on the magnetite ability to promote heterogeneous Fenton or photo-Fenton reactions, a topic that has attracted very little attention in the literature to date. Phenol was chosen as model compound in this study because of its well-known
photo-Fenton degradation, which considerably aids in the understanding of the reaction pathways [30, 31].

2. Experimental

2.1. Reagents and materials.

Phenol (purity grade 99%), 1,10-phenanthroline (99%), hydroquinone (98%), H₃PO₄ (85%), HClO₄ (70%), magnetite (97%) and methanol (gradient grade) were purchased from Aldrich, magnetite (98%) from Prolabo, H₂O₂ (35%) from VWR International. All reagents were used as received, without further purification. The aqueous solutions were prepared by using water of Milli-Q purity (TOC < 2 ppb, resistivity ≥18.2 mΩ cm).

2.2. Synthesis of magnetite samples

Experiments were conducted with four different kinds of magnetite. Among them, two (S1 and S2) were prepared in the lab from two different Fe(III)-oxyhydroxides, the third one (S3) was purchased from Prolabo and the fourth (S4) from Aldrich. S1 and S2 were prepared by starting from 2-line ferrihydride and lepidocrocite (γ-FeOOH), respectively. The ferrihydrite and lepidocrocite were synthesized as explained in previous work [32], according to the methods proposed by Schwertmann and Cornell [33]. All the Fe³⁺ precipitates were washed several times to remove electrolytes, centrifuged and then dried. Starting from these materials, S1 and S2 were then prepared by Fe²⁺-induced mineralogical transformations of synthetic ferric oxyhydroxides, as explained in detail in previous works [32, 34, 35]. The suspensions were vigorously stirred for two days, they were then centrifuged and the solid was dried in a glove box.

2.3. Sample characterization

To identify the crystal structure of minerals, the solid samples were analyzed by X-ray powder diffraction (XRD). The XRD data were collected with a D8 Bruker diffractometer, equipped with a monochromator and a position-sensitive detector. The X-ray source was a Co anode (λ = 0.17902 nm). The diffractogram was recorded in the 3-64° 2θ range, with a 0.0359° step size and a collection time of 3 s per point.

Transmission Electron Microscopy (TEM) analysis was also performed to obtain information regarding morphology, size, shape and arrangement of particles. TEM observations were carried out with a Philips CS20 TEM (200 kV) coupled with an EDAX energy dispersive X-ray spectrometer. The solid powder was re-suspended in 2 mL ethanol under ultrasonication and a drop
of the suspension was evaporated on a carbon-coated copper grid, which was placed on filter paper for analysis.

The specific surface area of the iron oxides was determined by multipoint N₂–BET analysis, using a Coulter (SA 3100) surface area analyzer. The point of zero charge (PZC) of the tested magnetites was determined by potentiometric titration of the oxides in a thermostatic double-walled Pyrex cell at 293 K in 0.001, 0.01 and 0.1 M NaCl solutions, according to the method of Parks and de Dyne [36]. The N₂ gas was constantly passed through the suspensions to bubble out CO₂. The pH value of the suspensions was adjusted with titrant solutions (HCl or NaOH) and recorded with an Orion pH meter model 710A, connected to a combined glass electrode. Blank titrations were also performed with similar solutions in the absence of solid.

The Fe²⁺ content in the oxide structure was determined by chemical analysis after acid dissolution with 6 M HCl. Ferrous and total iron concentrations were determined using a modified 1,10-phenanthroline method [37]. The Fe²⁺/Fe³⁺ ratios of oxides are reported in Table 1. All chemical analyses were performed in triplicate.

The degree of aggregation of the magnetite primary particles in circumneutral and acidic (10⁻³ M HClO₄) aqueous suspensions was assessed with an ALV-NIBS (Langen, Germany) Dynamic Light Scattering (DLS) instrument, equipped with a Ne–He laser and an ALV-5000 multiple tau digital correlator. The scattered light intensity for each sample was recorded for at least 30 s at 298 K, at an angle of 173° with respect to the incident beam. For each sample the measurement was repeated for at least three times on 30, 20 and 10 mg dm⁻³ suspensions. The hydrodynamic radius was calculated using the cumulant method and it was expressed as the average value for all the tested concentrations.

### 2.4. Irradiation experiments.

Magnetite stock suspensions at 1 g L⁻¹ loading were prepared by ultrasonication (Branson 2200 ultrasonic bath, 40 kHz). The suspensions for irradiation (50 mL total volume in a beaker) were prepared by adding magnetite for a final loading of 0.2 g L⁻¹ (unless otherwise reported), as well as H₂O₂ (where relevant) and phenol from separate stock solutions. The natural pH value of the system was ~6, measured with a combined glass electrode connected to a Metrohm 713 pH meter. When required, the pH of the system was adjusted to 3 upon addition of HClO₄. Irradiation took place under a Philips TL 09N UVA lamp, with emission maximum at 365 nm. The lamp irradiance on top of the irradiated suspensions was 18 W m⁻² in the 295-400 nm range, measured with a CO.FO.ME.GRA. (Milan, Italy) power meter. The TL 09N lamp was chosen because UVA radiation is present in the sunlight spectrum, while preliminary experiments showed that visible light alone was ineffective in inducing degradation processes. Samples were mechanically stirred.
during irradiation. A picture showing the adopted experimental set-up is provided in the Supplementary Material (hereafter SM), in Figure SM1.

At selected time intervals, 1 mL aliquots (precisely measured) were withdrawn from the irradiated suspension, added with 0.5 mL methanol to quench the Fenton reaction, and filtered on Millipore HV syringe filters (Teflon, 0.45 µm pore diameter). The clear solutions underwent analysis by high-performance liquid chromatography coupled to diode array detection (HPLC-DAD). The instrument used was a VWR Hitachi Elite chromatograph, equipped with L-2200 Autosampler (injection volume 60 µL), L-2130 quaternary pump for low-pressure gradients, L-2300 column oven (set at 40ºC), and L-2455 DAD detector. The column used was a RP-C18 LichroCART (VWR Int., length 125 mm, diameter 4 mm), packed with LiChrospher 100 RP-18 (5 µm diameter). Elution was carried out with a 40:60 mixture of methanol: aqueous H₃PO₄ (pH 2.8) at 1.0 mL min⁻¹ flow rate, with detection at 220 nm. Under these conditions the retention times were 1.5 min for hydroquinone (HQ) and 4.9 min for phenol. The column dead time was 0.9 min.

In some runs the concentration of dissolved Fe was also checked. In this case 5 mL aliquots were withdrawn, magnetite was separated by a magnet and the supernatant was filtered. If needed, 50 µL of concentrated HClO₄ were added to acidify the solution. The clear sample was analyzed with a Varian Liberty 100 Inductively Coupled Plasma-Optical Emission Spectrometer (ICP-OES), provided with a Czerny-Turner monochromator, a Sturman-Masters spray chamber, a V-groove nebulizer and a radio frequency (RF) generator at 40.68 MHz.

3. Results and Discussion

3.1 Synthesis and characterization of the magnetite samples

Magnetite (Fe₂O₄) is the only pure oxide of mixed valence and it is usually represented by the formula (Fe³⁺)₄₋₈[Fe³⁺Fe²⁺]₄₋₈O₄ [20]. It has a cubic spinel structure with iron in both tetrahedral and octahedral sites. Tested magnetites were characterized by XRD (Fig. 1a). For the S3 and S4 samples, five diffraction peaks at 2θ = 21.2°, 35°, 41.2°, 50.4° and 62.8° could be assigned to Fe₂O₄, magnetite [33]. The d-space values of these main peaks were 2.53, 2.96, 2.09, 4.85 and 1.71 Å, which may respectively correspond to the more intense lines of magnetite 311, 220, 400, 111 and 422. It should be noted that the XRD pattern of S1 and S2 shows the same peaks that are less intense. The broad nature and low intensity of the peaks in the spectra of S1 or S2 can result from nanosized particles (Fig.1b), which may exhibit poor crystallinity [33, 38].

TEM images show that magnetite particles are highly aggregated and exhibit irregular shapes and non-uniform size (Fig. 1b). S1 particles were smaller with quasi-spherical shape. The shape of S2
particles was between hexagonal to octahedral, while S3 exhibited non-uniform size and shape. The TEM image of S4 shows more or less rhombohedral particles, with crystals varying between 100 and 300 nm in length. TEM combined with energy-dispersive X-ray spectrometry (EDXS) yields an elemental analysis of the sample, so that elemental ratios can be calculated by EDXS and compared with the known mineralogical composition. EDX microanalysis showed the characteristic Fe/O ratio of magnetite (~ 0.75) for all samples. The particle diameter range deduced from all images is reported in Table 1.

As the Fe\(^{II}\) content is a key parameter in the heterogeneous Fenton reaction, the Fe\(^{II}\)/Fe\(^{III}\) ratio was determined by chemical analysis for each magnetite sample and it is reported in Table 1. S1 and S4 have the lowest Fe\(^{II}\) content and they are quite far from stoichiometric magnetite [24]. While S4 is a commercial sample, S1 may still contain some impurities as residuals of ferrihydrite particles from the synthesis method. TEM images of S1 and its high surface area and PZC value (close to that of ferrihydrite, i.e. ~8.2 [13]) support this statement. Overall, the tested magnetites have different particle size (S1 < S2 < S4 < S3), while the reverse order was found for BET surface area and PZC (Table 1). The pH values of the point of zero charge (pH\(_{\text{PZC}}\)) estimated from potentiometric titration are close to those reported in the literature (Table 1) [39, and references therein].

Figure 1c shows the hydrodynamic radii of the tested magnetites, both at circumneutral pH and at pH 3 (1 mM HClO\(_4\)). Inspection of the particle size data reported in Table 1 (solid phase) and in Figure 1c (aqueous suspensions) suggests the following statements: (i) magnetite particles show a high degree of aggregation in aqueous suspension; (ii) the dimensions of the aggregates follow the same order as the primary particles (S1 < S2 < S4 < S3), while the reverse order was found for BET surface area and PZC (Table 1). The pH values of the point of zero charge (pH\(_{\text{PZC}}\)) estimated from potentiometric titration are close to those reported in the literature (Table 1) [39, and references therein].

3.2 Photo-Fenton degradation of phenol

Preliminary experiments showed that the photo-Fenton system with magnetite had an optimal loading of 0.2 g L\(^{-1}\) (higher loadings did not increase the rate of degradation considerably, see Figure SM2 in SM), and that 1 mM H\(_2\)O\(_2\) was quite effective in inducing degradation of 0.1 mM phenol. Therefore, such conditions were chosen as the test ones to compare the behavior of the different magnetite samples.

Figure 2 reports the time trend of phenol in the photo-Fenton system with the different magnetite samples at pH 3. Blank runs were also carried out, where one or more components of the system (H\(_2\)O\(_2\), magnetite and irradiation) were removed to highlight their effect toward phenol degradation. As far as blank runs are concerned, the following observations can be made: (i) phenol did not undergo significant photolysis under UVA at the adopted time scale (4 h), and also its degradation
in the presence of H\textsubscript{2}O\textsubscript{2} under irradiation (which could produce •OH [9]) was negligible; (ii) magnetite did not induce significant phenol degradation when irradiated without H\textsubscript{2}O\textsubscript{2} (suggesting that no photocatalytic processes were operational with irradiated magnetite) or when added with H\textsubscript{2}O\textsubscript{2} in the dark. The blank runs also showed that phenol adsorption on magnetite is negligible, which suggests that removal (when applicable) was due to actual transformation and not just to phase transfer. The results of most blank runs are not reported in Figure 2 for readability issues. Irradiation of magnetite with H\textsubscript{2}O\textsubscript{2} was required to trigger reactivity. The presence of Fe\textsuperscript{II} at the magnetite surface might suggest the possibility of a dark Fenton reaction, but Figure 2 shows that for a 4-h time scale (and in the absence of hydroquinone, vide infra) no significant phenol degradation was observed in the dark, in the presence of magnetite (S1 to S4) and H\textsubscript{2}O\textsubscript{2}.

As far as the photo-Fenton experiments are concerned (magnetite + H\textsubscript{2}O\textsubscript{2} under irradiation), Figure 2 shows that complete or almost complete phenol degradation could be achieved within 4 h or less. S2 and S3 were the most photoreactive samples, with less than 2 h irradiation required to halve the concentration of phenol. In the case of S1 and S4, 2-3 h irradiation was required to obtain the same result. The size of particles and particle aggregates did not seem to play a key role in reactivity, because S1 had the smallest particles and the particles of S4 were smaller compared to S3, but S2 and S3 were more reactive compared to S1 and S4. Interestingly, the most reactive samples (S2 and S3) were also those showing the highest structural Fe\textsuperscript{II} content (see Table 1). This observation may be consistent with a previous report on the magnetite-catalyzed Fenton reaction in the dark [15]. However, it is unlikely that the higher reactivity of S2 and S3 under photo-Fenton conditions may be explained by a simple mechanism in which Fe\textsuperscript{II} is directly released in solution where it takes part to the Fenton reaction. Indeed, if it were the case, one should expect significant phenol transformation with magnetite + H\textsubscript{2}O\textsubscript{2} in the dark, which was on the contrary negligible at the time scale of our experiments. The facts that light was needed to induce phenol transformation and that the role of irradiation was not linked to the photolysis of H\textsubscript{2}O\textsubscript{2} (no transformation took place with H\textsubscript{2}O\textsubscript{2} alone) might suggest that Fe\textsuperscript{III} photoreduction to Fe\textsuperscript{II} (see e.g. reaction 4 or its corresponding process on the oxide surface) was required to trigger the degradation process.

The most remarkable feature of phenol transformation with irradiated magnetite + H\textsubscript{2}O\textsubscript{2} is represented by the shape of its time evolution. In all the cases, although with different kinetics, phenol transformation was initially slow and then it gradually accelerated, before slowing down when almost all the substrate was degraded. Similar profiles have been observed in the phototransformation of nitrobenzene in the presence of soluble Fe\textsuperscript{III} and H\textsubscript{2}O\textsubscript{2} [40]. In AOPs, this behavior is less common than first-order degradation kinetics [29] and it deserves explanation. A first possibility (phenol adsorption on magnetite) can be excluded by blank experiments. An alternative explanation is that time is required for Fe dissolution to take place, and that the phenol reaction rate is highly dependent on the concentration of dissolved Fe. ICP-OES measurements indicated that total Fe was gradually released in solution from magnetites at pH 3 in the presence of
phenol and H$_2$O$_2$, both in the dark and under irradiation (see Fig. 3). The gradual increase of dissolved Fe concentration is consistent with the observed time trend of phenol, because phenol transformation rate is low when dissolved Fe is also low and it increases with increasing Fe in solution. However, a closer look at the experimental data indicates that the concentration of dissolved Fe is not the only or the main factor involved. The highest values of Fe concentration in solution were observed in the presence of magnetite S3 and H$_2$O$_2$ in the dark, which induced negligible transformation of phenol. In the irradiated systems, similar Fe time trends were observed with S1, S3 and S4, despite differences in reactivity. An important issue is that Fe$^{II}$ is considerably more reactive than Fe$^{III}$ towards H$_2$O$_2$, thus the speciation of Fe in the studied systems plays an even more important role than its total concentration in solution. Unfortunately it was not possible to quantify dissolved Fe$^{II}$ in the studied systems, because the colorimetric method for Fe$^{II}$ is less sensitive than the ICP-OES technique for total Fe, and because fast reaction with H$_2$O$_2$ [2, 3] would keep the Fe$^{II}$ concentration low.

In the dark Fenton reaction it has been shown that some aromatic compounds such as catechol and hydroquinone or even humic acids can enhance substrate degradation [40-42]. An important issue is that these compounds are involved in the reduction of Fe$^{III}$ to Fe$^{II}$, which considerably accelerates the slow second step of the Fenton process. In such a case, the shape of phenol time evolution would be due to the fact that the intermediates need first to be formed from phenol before they can accelerate the reaction. The pathway of Fe$^{III}$ reduction by hydroquinone (HQ) is shown below [41]:

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\text{Catechol and HQ are formed in the Fenton degradation of phenol [40], as typical intermediates of the reaction between phenol itself and } \cdot \text{OH or related/mimicking oxidants. In our system HQ could for instance be detected at concentration values up to 4-5 } \mu \text{M, as a result of the formation-transformation budget. To see if HQ can be responsible for the observed kinetics of phenol transformation, it was added to the system from the very start at an initial concentration of 50 } \mu \text{M. If the hypothesis depicted in reaction (6) is correct, the availability of HQ in much higher concentration should considerably enhance phenol transformation. Figure 2 shows that the presence}
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of HQ considerably accelerated phenol degradation, the most remarkable acceleration being observed with S1 and S4. This happens despite the fact that HQ is a scavenger of reactive species (including •OH [43]), thus it could compete with phenol for photo-Fenton transformation. Such a finding provides evidence that the reduction of Fe$^{III}$ to Fe$^{II}$ is very important in the photo-Fenton transformation of phenol in the presence of magnetite.

In the case of S2 (or S3) + H$_2$O$_2$, addition of HQ induced phenol degradation even in the dark (Figure 2), while no dark transformation could be observed without HQ. In contrast, HQ did not induce dark transformation of phenol in the presence of H$_2$O$_2$ with S1 and S4. This finding suggests that Fe$^{III}$ reduction would be easier for S2 and S3 compared to S1 and S4. At equal pH conditions, the speciation (and, therefore, the reactivity) of dissolved Fe$^{III}$ would be the same with all the studied magnetite samples. Therefore, differences in reactivity should involve Fe$^{III}$ at the surface of the solid. At first sight, S2 and S3 differ from S1 and S4 because of the higher content of structural Fe$^{II}$. This is not easily or directly linked to the photo-Fenton reactivity, because Fe$^{III}$ reduction is needed to start the reaction. However, a different stoichiometry of the solids could modify the reactivity or the accessibility of the surface sites, including the Fe$^{III}$ ones.

Figure 4 reports the time evolution of phenol (initial concentration 0.1 mM) in the photo-Fenton reaction (1 mM H$_2$O$_2$ + 0.2 g L$^{-1}$ magnetite under UVA) at pH 6, the latter not varying significantly during irradiation. Blank runs are also reported, where no phenol degradation was observed. Differently from pH 3, pH 6 is definitely not the optimum for the Fenton or photo-Fenton reaction to take place [44-48]. This issue might account for the lack of reactivity at pH 6 of the magnetite samples S1 and S4, which were also the least reactive at pH 3. In the case of S2 and most notably S3, phenol transformation at pH 6 was not much modified compared to pH 3, when considering either the shape of the curve or the time required to reach a certain degree of phenol transformation. These data confirm the high reactivity of S2 and S3 under all the studied conditions. The time evolution of dissolved Fe at pH 6 is reported in Figure 3, where it is compared with the corresponding data at pH 3. The release of dissolved Fe in the dark was always lower at pH 6 than at pH 3, while under irradiation the difference between the two pH values was much less important. It is possible that the photoassisted reduction of Fe$^{III}$ would facilitate the release of iron in solution and that this process would become particularly important at pH 6. However, as already observed at pH 3, there was no straightforward correlation between dissolved Fe concentration and the ability of magnetite S1-S4 to degrade phenol. Furthermore, while the much lower dissolved Fe under irradiation at pH 6 compared to pH 3 could be consistent with the lack of photoreactivity of S4 at pH 6, in the case of S1 (unreactive at pH 6 as well) the amount of dissolved Fe under irradiation was comparable to (and even higher than) that at pH 3. In the cases of S2 and S3, which showed comparable photoreactivity toward phenol at both pH values, the Fe concentration was similar at pH 3 and 6 only for low irradiation times. At longer times, Fe accumulated in solution at pH 3 but not at pH 6. Such an overall scenario could be a consequence of the complexity of the processes
involved. In the presence of H\textsubscript{2}O\textsubscript{2}, dissolved Fe would mainly be in the form of Fe\textsuperscript{III} [2, 3], which could take part to the Fenton reaction only by undergoing reduction to Fe\textsuperscript{II}. The pH variation in the 3-6 range could have variable effects on the reduction process: on the one side, the concentration of FeOH\textsuperscript{2+} (the Fe\textsuperscript{III} form that undergoes the easiest reduction) would decrease with increasing pH, but on the other side HO\textsuperscript{2-} would be deprotonated to O\textsubscript{2-} that is more active towards Fe\textsuperscript{III} reduction [2, 3]. Furthermore, reduction could involve Fe\textsuperscript{III} on the oxide surface in addition to the dissolved one.

There are very different literature reports concerning the performance of the Fe\textsuperscript{III}-based heterogeneous photo-Fenton processes at –neutral vs. acidic pH. While in some cases considerable reactivity under neutral conditions is reported [49-51], in other cases a substantial decrease of the performance has been observed with increasing pH [52, 53]. It is possible that, as hypothesized in the present work, the effect of pH on reactivity depends on several factors such as Fe solubility and speciation and the availability and speciation of reducing agents, with different outcomes depending on the particular system under study.

Interestingly, 50 µM HQ at pH 6 slightly enhanced phenol transformation in the case of S2, but inhibited it with S3 (see Figure 4). A possible explanation is that HQ can both reduce Fe\textsuperscript{III} to Fe\textsuperscript{II}, thereby enhancing the Fenton process, and compete with phenol for the reaction with transient species including *OH [43]. The latter phenomenon would inhibit phenol transformation, and the HQ effect would be a balance between opposite trends. HQ could thus favor or inhibit phenol degradation depending on the magnetite sample and the operational conditions. There is also evidence that *OH and/or another oxidant with similar reactivity was involved in the transformation of phenol in the studied systems. Indeed, the addition of 2-propanol (0.01 M initial concentration) as *OH scavenger [43] to phenol + H\textsubscript{2}O\textsubscript{2} + magnetite (S2 or S3) under irradiation at both pH 3 and 6 was able to quench phenol degradation.

4. Conclusions

In this work, effective phenol degradation was obtained upon irradiation of magnetite in the presence of H\textsubscript{2}O\textsubscript{2} (heterogeneous photo-Fenton conditions). The initially low reaction rate considerably increased as the reaction progressed, most likely because of: (i) the formation of intermediates such as HQ that are able to reduce Fe\textsuperscript{III} to Fe\textsuperscript{II}, which takes part to the Fenton reaction, and (ii) the presence of dissolved Fe due to magnetite dissolution, both in dark and under irradiation. The need of Fe\textsuperscript{III} reduction, which is also supported by the important role of irradiation, might look surprising when considering that magnetite contains structural Fe\textsuperscript{II}. It is thus possible that the latter is not readily available for the Fenton reaction and that some form of initial activation (e.g. partial iron photodissolution) is needed. On the other hand, phenol degradation was most effectively achieved by the samples having the highest content of structural Fe\textsuperscript{II}. The size of the
primary particles seems to be less important, because it was not correlated with photo-Fenton activity in the studied samples. However, the size effect could be largely offset by important aggregation phenomena in the aqueous suspensions.

From an applicative point of view, the most interesting features of magnetite as shown in this study are:

- Reactivity upon absorption of UVA radiation, which makes cheap sunlight potentially applicable to carry out the reaction.
- Ability of some samples (most notably S2 and S3) to maintain reactivity also under circumneutral pH conditions, which could save the need to adjust and back-adjust pH.
- Limited iron leaching (up to or below the ppm range), which keeps dissolved Fe safely below the limits for wastewater discharge.
- Magnetic behavior, which greatly helps in the separation of magnetite from treated wastewater.

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References

Table 1. Some properties of the magnetite samples used in this study.

<table>
<thead>
<tr>
<th>Magnetite samples</th>
<th>Particle size range</th>
<th>SSA BET (m² g⁻¹)</th>
<th>PZC</th>
<th>Fe⁺⁺/Fe⁺⁺⁺ ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>30-50 nm</td>
<td>75 ± 2</td>
<td>8.1</td>
<td>0.34 ± 0.2</td>
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<tr>
<td>S2</td>
<td>60-80 nm</td>
<td>26 ± 1</td>
<td>7.8</td>
<td>0.42 ± 0.2</td>
</tr>
<tr>
<td>S3</td>
<td>1-2 µm</td>
<td>1.7 ± 0.2</td>
<td>7.4</td>
<td>0.43 ± 0.2</td>
</tr>
<tr>
<td>S4</td>
<td>100-300 nm</td>
<td>8.5 ± 0.5</td>
<td>7.6</td>
<td>0.30 ± 0.2</td>
</tr>
</tbody>
</table>
Figure 1: XRD (a) and TEM (b) images of the four used magnetite samples; (c) Hydrodynamic radii ($R_{\text{hyd}}$) of the magnetite particles/aggregates, measured by Dynamic Light Scattering in pure water and in 1 mM HClO$_4$. Error bars represent 95% confidence intervals.
Figure 2. Time trend of phenol (initial concentration 0.1 mM, pH 3 by HClO₄) under photo-Fenton conditions (phenol + H₂O₂ + magnetite S1-S4 + UVA) and in blank runs. When relevant, other conditions were as follows: 1 mM H₂O₂, 0.2 g L⁻¹ magnetite loading, 50 µM added HQ.
Figure 3. Total iron released in solution, both in the dark (open symbols) and under irradiation (solid symbols), at pH 3 by HClO$_4$ (circles) and at pH 6 (natural pH, triangles), in the presence of the studied magnetite samples. Conditions were as follows: initial phenol concentration 0.1 mM, 1 mM H$_2$O$_2$, 0.2 g L$^{-1}$ magnetite loading.
Figure 4. Time trend of phenol (initial concentration 0.1 mM, pH 6) under photo-Fenton conditions (phenol + H_2O_2 + magnetite S1-S4 + UVA) and in blank runs. When relevant, other conditions were as follows: 1 mM H_2O_2, 0.2 g L^{-1} magnetite loading, 50 μM added HQ.