Sonocatalytic oxidation of EDTA in aqueous solutions over noble metal-free Co$_3$O$_4$/TiO$_2$ catalyst

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- Sonochemistry
- Ultrasound
- Advanced oxidation processes
- EDTA
- Heterogeneous catalysis

Abstract

The sonocatalytic degradation of EDTA in aqueous solution was studied under ultrasound irradiation (345 kHz, 73 W, acoustic power 0.20 W·mL$^{-1}$, Ar and Ar/O$_2$ saturating gases, $T = 20–50$ °C) in the presence of Co$_3$O$_4$/TiO$_2$ and Pt/TiO$_2$ nanocatalysts. About 90% of EDTA ($C_0 = 5 \times 10^{-3}$ M) was oxidized during ultrasonic treatment at 40 °C in the presence of the Co$_3$O$_4$/TiO$_2$ catalyst and Ar/O$_2$ gas mixture. By contrast, Pt/TiO$_2$ catalyst exhibited much lower sonocatalytic activity in this system. Suggested mechanism of EDTA oxidation in the presence of Co$_3$O$_4$/TiO$_2$ catalyst involved the generation of oxidizing radicals by acoustic cavitation and Co(II)-Co(III) redox process. Quite low apparent activation energy of the sonocatalytic process ($E_a = 19$ kJ mol$^{-1}$) was attributed to diffusion of reagents in the vicinity of the active sites of catalyst. Sonocatalytic degradation of EDTA is accompanied by formation of iminodiacetic acid, formic acid, oxalic acid, glycolic acid and acetic acid as intermediate products in an agreement with radical-driven mechanism.

1. Introduction

Ethylenediaminetetraacetic acid (EDTA) is widely used in food, cosmetic and pharmaceutical industries [1]. While EDTA is nontoxic itself, it is a metal-mobilizing pollutant that requires its removal from wastewater. In nuclear industry, EDTA is an obvious decontaminating agent due to its ability to form strong chelating complexes with a large number of radionuclides [2]. However, the formation of such complexes may impede further radionuclide removal from the liquid waste, i.e. through chemical precipitation or ion exchange [3]. Therefore, EDTA must be decomposed or at least defunctionalized prior to radionuclide specific treatments, whereas the methods of its decomposition should use minimal amounts of reagents, in order to minimize the volume of final waste. Advanced Oxidation Processes (AOP), based on the generation of highly reactive radical species, can potentially enable the effective oxidation of chelating agents in waste solutions [4]. Indeed, different AOPs have been lately considered for EDTA oxidation, such as catalytic Fenton-like processes [5,6], ozonolysis [7], hydrothermal oxidation [8], photolysis [9] and photocatalysis [10]. All of these techniques have some limitations, for instance, large amount of catalyst and H$_2$O$_2$ are required for Fenton processes. In the same way, ozone is not an eco-friendly and healthy product and the others treatments need high energy input. For all these reasons, researches have to be carried out to find more ecological and cost effective approaches for the treatment of wastewaters.

Among the different AOPs, sonochemistry stands as a promising technology that can provide an answer to this EDTA degradation issue. Indeed, this technique enables the in situ production of radical species. The chemical effects of power ultrasound come from acoustic cavitation, which is a set of consequent events: nucleation, growth and violent collapse of microbubbles in liquids submitted to ultrasonic vibrations. In aqueous solutions, cavitation causes splitting of water molecules leading to formation of OH$^-$ radicals and H atoms, and to the products of their mutual recombination: H$_2$O$_2$ and H$_2$, respectively [11]. Recent spectroscopic studies of light emitted by collapsing bubbles, called sonoluminescence, revealed formation of nonequilibrium plasma inside the bubble [12]. Therefore, each cavitation bubble can be considered as microscopic plasma reactor producing highly reactive species in liquids at almost room bulk temperature. Indeed, the sonochemical degradation of EDTA in homogeneous aqueous solutions has been reported in the presence of oxygen [13] and hydrogen peroxide [14]. However, despite the extreme local conditions inside the cavitation bubbles, the overall reaction rate was found to be quite low. Sonochemical oxidation of EDTA can be significantly accelerated in the presence of zero-valent...
iron particles via Fenton-like mechanism [15]. However, iron dissolution during this process would require additional purification stage after EDTA degradation.

It has been recently shown that power ultrasound accelerates the catalytic oxidation of carboxylic acids in the presence of powdered Pt/TiO₂ and Ar/O₂ gas mixtures [16]. The sonocatalytic effect was attributed to the strong dispersion of the Pt/TiO₂ catalyst together with the generation of chemically active species during water sonolysis. Though active, noble metals are quite expensive and, therefore, alternative catalytic systems need to be found. In the present paper we report, for the first time, quantitative sonochemical defunctionalization of EDTA in the presence of a noble metal-free catalyst Co₃O₄/TiO₂. Moreover, we found that Co₃O₄/TiO₂ exhibits much higher sonocatalytic activity than Pt/TiO₂ in studied reaction. Ultrasonic frequency and gas composition were chosen on the basis of previous researches [12,16].

2. Materials and methods

2.1. Reagents

Solids Na₂H₂EDTA·2H₂O (98%, Fluka AG), Co(NO₃)₂·6H₂O (98%, Alfa Aesar), TiO₂ anatase (99%, Sigma Aldrich, 95% anatase, 5% rutile), TiO₂ P25 (Aeroxide®, Evonik) and H₂PtCl₆ (999%, Strem) were used as received without further purification. All solutions were prepared with deionized water (Milli-Q 18.2 MΩ cm at 25°C). Ar/O₂ gas mixture with 20 vol.% of O₂, Ar of 99.999% purity and O₂ 99.999% purity were supplied by Air Liquide.

2.2. Synthesis of catalysts

Co₃O₄/TiO₂ catalysts with 6.4%wt., 3.3%wt. and 1.3%wt. of Co₃O₄ were prepared by excess solution impregnation using Co(NO₃)₂·6H₂O solution as cobalt precursor. The solution with suspended TiO₂ anatase powder was stirred one hour, and then dried at 65°C for 30 min in a rotary evaporator. The resulting material was further dried in an oven at 80°C for 20 min and subsequently calcinated at 550°C for 4 h in order to ensure a maximal yield of crystallized Co₃O₄ [17].

Pt/TiO₂ catalyst was prepared by incipient wetness impregnation technique, using H₂PtCl₆ solution and P25 TiO₂ as the support for a Pt nominal content of 1%. After drying Pt(IV) was reduced with hydrazine hydrate solution, washed until the absence of chloride ions (control with Ag⁺) and dried overnight under vacuum at room temperature. This technique enables a complete reduction of Pt(IV) at room temperature without modification of TiO₂ support [18].

2.3. Techniques of catalyst characterization

High-resolution transmission electron microscopy (TEM) measurements were carried out in a Jeol JEM 2200 FS device, equipped with a numeric camcorder. An ultrasonic bath was used to disperse the samples in deionized water. Then, one drop was deposited on carbon-coated copper grid and dried in atmospheric air prior to analysis. Powder XRD measurements were performed on the readily-prepared, fresh and used oxide catalysts in a Bruker D8 Advance Diffractometer, equipped with a Cu Kα (λ=1.54056Å) radiation source and a Lynx Eye detector. Powder XRD patterns were collected in the 10–90 2-theta degree range with a step size of 0.01° and a step time of 1 s. N₂ adsorption study at −196°C was achieved with a BELSORB-Mini II device (BEL-Japan). Specific surface area was calculated with BET method. Temperature programmed reduction (H₂-TPR) was performed in a BELCAT-M apparatus from BEL Japan equipped with a thermal conductivity detector. In each experiment 60 mg of catalyst were degassed at 100°C for 2 h and then heated to 900°C at 7.5 °C min⁻¹ heating rate in the presence of 5% vol. H₂/Ar gas mixture.

2.4. EDTA degradation experiments

A home-made 400 mL glass reactor enabled the control of temperature and gas atmosphere was used in the experiments (Fig. 1). This reactor was fixed on the top of a 345 kHz transducer (ELAC Nautik, 25 cm²), connected to a generator with a maximum electrical power of 125 W (T&C Power Conversion, Inc.). The electric load power was regulated at 73 W and the corresponding absorbed acoustic power was evaluated by thermal probe method at 0.20 W·mL⁻¹. The volume of sonicated solution was 200 mL, while additional mechanical stirring at 250 rpm was used in order to warrant a homogeneous suspension of catalyst particles [16]. The temperature during sonolysis was controlled between 20 and 50°C with a Huber Unistat Tango thermo-cryostat and measured by a thermocouple immersed into ultrasound-irradiated solution. Ar/O₂ gas mixture was bubbled through solution for 20 min at 204 mL·min⁻¹ flowrate before the experiment and kept during sonication. 400 mg of catalyst were dispersed in a minimal volume of water using ultrasonic bath prior to its injection to the EDTA solution. Sample aliquots were taken each hour to follow the evolution of EDTA, H₂O₂, intermediate degradation products and total organic carbon concentrations after their filtration using 0.2μm PTFE membrane. No EDTA adsorption in sampling filters was observed, since the EDTA concentration measured in the first filtered aliquot extracted from the reactor during the silent catalytic experiment was identical to the nominal initial concentration of EDTA used in the experiment. All experiments were reproduced twice to verify the suitability of our uncertainties. Formed CO₂ was continuously monitored in the outlet gas using a Prima BT mass spectrometer (Thermo Scientific).

A Thermo Scientific Evolution 220 UV–vis spectrophotometer was used for the analysis either of EDTA, titrated using a (Fe-TPTZ)₂⁺ complex, where TPTZ stands for Bis(2,4,6-tripyridyl-s-triazine) [19], or of H₂O₂, titrated with TiOSO₄ [20]. The total carbon in solution was followed using a Shimadzu TOC-VCSH analyzer calibrated with potassium phthalate solution. Various intermediate degradation products of EDTA were detected by ion chromatography in a Dionex AS-AP device equipped with a capillary column AS11. The used eluent for the
particles dispersed on the surface of TiO2 support. The average size of the product. In the case of 1.3%wt. Co3O4/TiO2, the cobalt oxide phase was not detected in the diffractograms, most probably due to the low cobalt oxide Co3O4 as the principal Co-phase, as well as of anatase, together with traces of rutile TiO2 (Fig. S1 of the Supplementary data). The presence of these two TiO2 phases is typical for commercial anatase catalysts (JCPDS file no. 00-042-1467) confirmed the presence of cobalt oxide Co3O4 in the 3.3%wt. Co3O4/TiO2, the cobalt oxide phase was not detected in the diffractograms, most probably due to the low cobalt concentration. The powder XRD patterns acquired for the Co-containing catalysts (JCPDS file no. 00-042-1467) confirmed the presence of cobalt oxide Co3O4 as the principal Co-phase, as well as of anatase, together with traces of rutile TiO2 (Fig. S1 of the Supplementary data). The presence of these two TiO2 phases is typical for commercial anatase product. In the case of 1.3%wt. Co3O4/TiO2, the cobalt oxide phase was not detected in the diffractograms, most probably due to the low cobalt concentration in this catalyst.

The HRTEM/EDX images shown in Figs. 2 and 3 for 1.3%wt. and 6.4%wt. Co3O4/TiO2 respectively and in Fig. S2 (Supplementary data) for 3.3%wt. Co3O4/TiO2 evidence presence of cobalt oxide nanoparticles dispersed on the surface of TiO2 support. The average size of Co3O4 nanoparticles calculated from HRTEM for each catalyst is shown in Table 1. Cobalt oxide nanoparticles around 10–15 nm are present in the 1.3%wt. Co3O4/TiO2 catalyst. Higher Co-loading results in an increase of the average Co3O4 particle size, i.e. in the 3.3%wt. Co3O4/TiO2 and 6.4%wt. Co3O4/TiO2 catalysts the size of the Co3O4 particles reaches 15–20 nm. Also certain cobalt oxide agglomeration can be observed at HRTEM images at high Co-loading. Whereas, crystallite size calculated by Scherrer equation (Table 1) were found to be equal to 39.7 ± 7.5 nm for 3.3%wt. Co3O4/TiO2 and 37.5 nm for 6.4%wt. Co3O4/TiO2. These values are consistent with HRTEM measurements taking into account quite broad particle size distribution and uncertainties of both techniques. No XRD information could be obtained for the 1.3%wt. Co3O4 sample because of low cobalt concentration. The specific surface area (Table 1) was measured quite low compared to TiO2 anatase specific surface area (45–55 m2 g−1) used in this work as well as other studies with TiO2 support [21]. This could be attributed to the calcination at 550°C which favors aggregation of cobalt oxide and titanium oxide particles [22]. The HRTEM micrographs acquired for the Pt/TiO2 catalyst are shown in Fig. 4 and reveal well dispersed 1–3 nm Pt nanoparticles on the TiO2 support as it was already reported in the literature [16].

Fig. 5 shows the H2-TPR profiles acquired for the different Co-containing catalysts. Table 2 summarizes the amounts of H2 consumed during TPR experiments for each catalyst. As expected, increasing Co-loading leads to higher H2-consumption during TPR, but this correlation is not directly proportional. TPR analysis reveals two reduction peaks common for all studied samples centered at 350°C and 480–500°C. According to the literature, the reduction of Co3O4 species on a TiO2 surface is a 2-step process involving the consecutive reduction of Co(III) to Co(II) and Co(II) to metallic Co [21,23,24]. Therefore, a first peak at about 350°C is expected to occur due to the reduction of Co3O4 to CoO, followed by a second peak at 450–500°C attributed to the reduction of CoO to metallic Co. Surprisingly, an intermediate reduction peak appeared at 500°C–520°C for the samples with higher Co3O4 content. As previously commented, HRTEM pointed out some agglomeration of the Co-phase with increasing of Co-loading. The coexistence of Co3O4 particles of different size may account for the splitting of the low temperature H2 consumption peak, i.e. larger Co3O4 particles will be reduced to CoO at higher temperatures [25]. The slight increase of the CoO reduction temperature from 500°C to 520°C with increasing of Co3O4 concentration from 3.3% to 6.4% (Fig. 5) can be attributed to the similar phenomenon. In the case of 2-step process, the area of the second peak should be three time more important than of the first one. However, in our study no direct relation could be found between each peak area. Then maybe, interaction with the support affects Co(III) and Co(II) reduction [21]. Another assumption could be the formation of intermediate nonstoichiometric cobalt oxides CoOx during H2-TPR reduction [26].

3. Results and discussion


All the catalysts prepared using cobalt salts were characterized upon their calcination, by means of both powder XRD and HRTEM techniques. In contrast, only HRTEM analysis was performed on the Pt/TiO2 catalyst. The powder XRD patterns acquired for the Co-containing catalysts (JCPDS file no. 00-042-1467) confirmed the presence of cobalt oxide Co3O4 as the principal Co-phase, as well as of anatase, together with traces of rutile TiO2 (Fig. S1 of the Supplementary data). The presence of these two TiO2 phases is typical for commercial anatase catalysts. The powder XRD patterns acquired for the Co-containing catalysts (JCPDS file no. 00-042-1467) confirmed the presence of cobalt oxide Co3O4 as the principal Co-phase, as well as of anatase, together with traces of rutile TiO2 (Fig. S1 of the Supplementary data). The presence of these two TiO2 phases is typical for commercial anatase catalysts. The powder XRD patterns acquired for the Co-containing catalysts (JCPDS file no. 00-042-1467) confirmed the presence of cobalt oxide Co3O4 as the principal Co-phase, as well as of anatase, together with traces of rutile TiO2 (Fig. S1 of the Supplementary data). The presence of these two TiO2 phases is typical for commercial anatase catalysts. The powder XRD patterns acquired for the Co-containing catalysts (JCPDS file no. 00-042-1467) confirmed the presence of cobalt oxide Co3O4 as the principal Co-phase, as well as of anatase, together with traces of rutile TiO2 (Fig. S1 of the Supplementary data). The presence of these two TiO2 phases is typical for commercial anatase catalysts.
almost independent of the solution temperature in the range of 20–40°C. In general, whatever the experimental conditions used, EDTA oxidation is quite slow in the absence of catalyst, as has been previously reported elsewhere [13].

Indeed, the slowest formation rate of H₂O₂ in the presence of EDTA than in water (Fig. 6b) suggests that the initial stage of EDTA sonochemical oxidation involves reactions with OH% and HO₂% radicals formed via water molecule splitting inside the cavitation bubbles:

\[ \text{H}_2\text{O} \rightarrow \text{H}^+ + \text{OH}^- \]  
\[ 2 \text{OH}^- \rightarrow \text{H}_2\text{O}_2 \]  
\[ \text{H}^+ + \text{O}_2 \rightarrow \text{HO}_2^- \]  
\[ 2 \text{HO}_2^- \rightarrow \text{H}_2\text{O}_2 \]  
\[ \text{OH}^- + \text{EDTA} \rightarrow \text{Oxidation Products} \]  
\[ \text{HO}_2^- + \text{EDTA} \rightarrow \text{Oxidation Products} \]

Actually, the Reactions (5) and (6) are competing with the reactions leading to H₂O₂ formation shown in the Eqs. (2) and (4). The plots in

Table 1
Co₃O₄ mean particle size obtained from HRTEM pictures and calculated from XRD patterns at 2-Thêta = 65.2° of Co₃O₄ using Scherrer equation, and specific surface area calculated by BET analysis. Particle size obtained by TEM was average from at least 20 parallel measurements.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>TEM measurement</th>
<th>XRD measurement</th>
<th>Specific surface area</th>
</tr>
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<tbody>
<tr>
<td>1.3%wt. Co₃O₄/TiO₂</td>
<td>13.5 ± 3.6 nm</td>
<td>39.7 ± 7.5 nm</td>
<td>7.7 ± 1.5 m²·g⁻¹</td>
</tr>
<tr>
<td>3.3%wt. Co₃O₄/TiO₂</td>
<td>18.0 ± 5.2 nm</td>
<td>39.7 ± 7.5 nm</td>
<td>7.9 ± 1.5 m²·g⁻¹</td>
</tr>
<tr>
<td>6.4%wt. Co₃O₄/TiO₂</td>
<td>19.5 ± 3.8 nm</td>
<td>37.5 ± 7.5 nm</td>
<td>7.2 ± 1.5 m²·g⁻¹</td>
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![Fig. 3. TEM picture (a) and EDX map (b) for the 1.3% wt. Co₃O₄/TiO₂ catalyst.](image1)

![Fig. 4. HRTEM images of the 1% wt. Pt/TiO₂ catalyst.](image2)

![Fig. 5. H₂-TPR profiles acquired for the 1.3% wt. Co₃O₄/TiO₂, 3.3% wt. Co₃O₄/TiO₂ and 6.4% wt. Co₃O₄/TiO₂ catalysts.](image3)

![Fig. 6b show the H₂O₂ formation during water sonolysis and in presence of EDTA. The difference between these plots evidences that only about 40% of oxidizing radicals reacts with EDTA and intermediate](image4)
products, while the remaining recombine into H$_2$O$_2$. The H$_2$O$_2$ evolution under argon atmosphere at 40°C and under Ar/20% O$_2$ atmosphere at 20°C is represented in Fig. S3 of the Supplementary data. Similar H$_2$O$_2$ evolution plot profile at 20°C and 40°C confirms the temperature independence on the H$_2$O$_2$ production yield and the EDTA degradation yield in homogeneous solutions.

3.2.2. EDTA sonocatalytic oxidation

The sonochemical oxidation of EDTA is largely enhanced in the presence of Co$_3$O$_4$/TiO$_2$ catalyst. Fig. 7a shows that under the studied experimental conditions about 90% of EDTA can be oxidized upon 7h ultrasonic treatment. Similar to homogeneous solutions, sonocatalytic reaction exhibits zero-order kinetics on EDTA. By contrast, the presence of Pt/TiO$_2$ enhances EDTA degradation of only 10% compared to sonochemical process without catalyst (Figs. 6a and 7b). This is a surprising result since noble metal-based catalysts usually show high catalytic performance in AOPs [16].

It is interesting to note that in the presence of pristine TiO$_2$ the reaction rate is very similar to that measured in homogeneous solutions, i.e. in absence of any solid material. This indicates that the active sites of the sonocatalytic reaction are situated on the surface of Co$_3$O$_4$ or Pt and not on TiO$_2$ support. However, both Co$_3$O$_4$/TiO$_2$ and Pt/TiO$_2$ were found to be completely inactive under silent conditions, i.e. under mechanical stirring without ultrasound. In the light of these results, the following mechanism of EDTA sonocatalytic degradation with the Co$_3$O$_4$/TiO$_2$ catalyst can be suggested: i) in a first step EDTA molecules are adsorbed on the surface of the Co$_3$O$_4$ sites, ii) Co(II) of Co$_3$O$_4$ mixed oxide is then oxidized to Co(III) by highly reactive radicals produced by the cavitation bubbles, and iii) EDTA is finally oxidized by Co(III), being consequently reduced to Co(II). The proposed reaction mechanism depicted in Fig. 8 is confirmed by recently published data about strong interaction of EDTA molecules with Co$_3$O$_4$ surface [27]. The observed low catalytic performance of Pt/TiO$_2$ catalyst in studied process can be explained by the difference in the reaction mechanism which involves adsorption of O$_2$ molecules at Pt surface in a first step instead of EDTA molecule in the case of Co$_3$O$_4$/TiO$_2$ catalyst [16]. Therefore, the initial stage of EDTA sonocatalytic oxidation with Pt/
The TiO₂ catalyst would be limited by the reactions with radicals in homogeneous solution or cavitation bubble/solution interface rather than by redox process on Pt/TiO₂ surface.

For better understanding of the reaction mechanism, the influence of temperature on the sonochemical oxidation of EDTA was evaluated in the range of 20–50 °C. The derived Arrhenius linear plot is shown in Fig. S4 of the Supplementary data. The relatively low apparent activation energy calculated from these data (Eₐ = 19 ± 2 kJmol⁻¹) suggests that the thermal effect is governed by reagent diffusion rather than by the splitting of EDTA chemical bonds, since the influence of the temperature on the intrinsic chemical reaction rate is expected to be much stronger [8]. Indeed, this conclusion is in line with the proposed radical-induced mechanism of EDTA sonochemical oxidation.

In order to elucidate the role of H₂O₂ in the reaction mechanism, the experiment of EDTA degradation was performed at 40 °C under mechanical stirring and Ar/20%O₂ atmosphere without ultrasound in the presence of Co₃O₄/TiO₂ catalyst and under progressive addition of H₂O₂. No oxidation of EDTA was detected in this experiment indicating that H₂O₂ itself is not involved in the reaction mechanism. In contrast, quite rapid decomposition of H₂O₂ was observed with a rate of 2.4 μmol min⁻¹ in the presence of Co₃O₄/TiO₂ (Figs. 9 and S5 of the Supplementary data) in agreement with recently published results [28,29].

Table 3 summarizes the kinetics of EDTA degradation as a function of Co₃O₄ content. The normalized EDTA oxidation rate clearly increases with decreasing of cobalt oxide loading. This effect can be most probably attributed to the decrease of Co₃O₄ particle size with decreasing of Co₃O₄ content. According to literature, the enhanced catalytic activity of small Co₃O₄ nanoparticles is related to their higher surface density of Co(III) moieties [30,31]. It is noteworthy that Co(III) is known to form very strong complexes with EDTA [8], which subsequently leads to more efficient sonochemical oxidation.

Finally, the influence of oxygen concentration in Ar/O₂ gas mixture was investigated under the same reaction conditions (2 g L⁻¹ of 6.4% wt. Co₃O₄/TiO₂, 40 °C, 200 mL, 5 mM EDTA, 345 kHz, Pₑ = 0.20 W mL⁻¹). Pure argon, Ar/20%O₂ and Ar/40%O₂ gas mixtures were used. The results are shown in Fig. S6 of the Supplementary data. Without oxygen in the system, EDTA degradation rate is very similar to the one observed under ultrasonic irradiation without catalyst. In addition, no significant differences were observed between 20% and 40% of oxygen in the gas mixture. One can conclude that oxygen is essential to provide maximal yield of OH⁻ and HO₂⁻ radicals via H atom scavenging without being a limiting species in this system.

3.2.3. Analysis of intermediate products

The plots shown in Fig. 10 evidence less effective TOC removal than EDTA degradation (Fig. 6a) which implies the formation of intermediate products. TOC data are supported by CO₂ mass spectrometric analysis in the outlet gas. For instance, in the experiment with Co₃O₄/TiO₂ catalyst 1.98 mmol of CO₂ were measured corresponding to 2 mmol. TOC abatement (Fig. 10). Indeed, ion chromatography analysis allowed to identify the presence of iminodiacetic acid, formic acid, oxalic acid, glycolic acid and acetic acid in the solutions treated with ultrasound. Chemical structure of these products and typical chromatograms are shown in Figs. S7 and S8 of the Supplementary data respectively. The concentrations of these products measured after 7 h of sonolysis are summarized in Fig. 11. The fragments of EDTA degradation observed in our experiments are very similar to those reported for EDTA γ-radiolysis in oxygenated solutions [32]. Radiolytic degradation of EDTA occurs via the reactions with OH⁻ and HO₂⁻ radicals. Therefore, the similarity of the intermediate products for both processes would indicate the radical mechanism of EDTA sonochemical degradation. Iminodiacetic acid is formed at the earlier stage of EDTA oxidation. By contrast, formic acid is a product of deep EDTA degradation. Consequently, lower concentration of iminodiacetic acid and higher concentration of formic acid in solution treated with Co₃O₄/TiO₂ catalyst compared to sonicated homogeneous solution indicate more profound EDTA degradation in sonochemical process.

The ultrasonic irradiation in the presence of Pt/TiO₂ catalyst is less effective for TOC removal compared to Co₃O₄/TiO₂ catalyst (Fig. 10), which agrees with general lower catalytic activity of platinized catalyst for EDTA oxidation. On the other hand, the concentrations of iminodiacetic acid, formic acid and oxalic acid are clearly lower in the presence of Pt/TiO₂ particles than in homogeneous solutions (Fig. 11). Therefore, one can conclude that this catalyst improves the degradation of intermediates, which lead to somewhat better TOC abatement compared to homogeneous solution (Fig. 10). This is traced back to the difference in the reaction mechanisms with Co₃O₄/TiO₂ and Pt/TiO₂ catalysts. In the case of Pt/TiO₂, the principal mechanism related to the activation of O₂ molecules on Pt surface provides better efficiency for oxidation of small carboxylic acids [16] formed as intermediates during sonochemical degradation of EDTA.

Through chemical analysis allowed to establish a carbon balance in solution summarized in Fig. S8 of the Supplementary data. In sonochemical process with Co₃O₄/TiO₂, some unidentified species can reach 35% of initial carbon balance. On the other hand, the concentration of such products is negligible for the process in homogeneous solution which can be related to some difference in the reaction mechanism with and without catalyst.

3.2.4. Catalyst stability

The stability of both catalysts was assessed in the presence of 5 mM EDTA with mechanical stirring and Ar/O₂ atmosphere under ultrasonic irradiation and silent conditions during 7 h at 40 °C. ICP OES analysis showed that only 0.2% and 1.4% for Pt and Co respectively of an equivalent metal quantity were leached into the solution under ultrasound. Metal leaching under silent conditions was found to be equal to 0.2% and 2% for Pt and Co respectively. These results are within the ICP technique uncertainty and point out a good catalyst stability, and practically rules out the possibility of any secondary homogeneous oxidation reaction occurring within the liquid phase.

Table 3

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>6.4%wt.</th>
<th>3.3%wt.</th>
<th>1.3%wt.</th>
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<tbody>
<tr>
<td>W (mmol h⁻¹ g₃Co₄⁻¹)</td>
<td>6.4</td>
<td>12.7</td>
<td>32.9</td>
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Fig. 9. Evolution of H₂O₂ quantity under 345kHz ultrasound (Pₑ = 0.20 W mL⁻¹, T = 40 °C, Ar/20%O₂).
3.2.5. Characterization of the spent catalysts

The Co-containing catalysts were characterized by XRD, BET and H2-TPR upon their exposure to ultrasound irradiation during the sonochemical experiments. XRD measurements of spent 6.4%wt. Co3O4/TiO2 catalyst show the same patterns as initial catalyst. The crystallite size of Co3O4 and the specific surface area of spent Co 3O4/TiO2 catalyst were also found practically equal to that before the sonocatalytic treatment (37.8 ± 7.5 nm and 6.9 ± 1.5 m2/g respectively). So, this catalyst was apparently stable during 7 h of ultrasonic treatment.

The H2-TPR profiles acquired for the fresh and spent 6.4%wt. Co3O4/TiO2 catalysts are shown in Fig. 12. The profile obtained for the spent catalyst differs from that acquired for the fresh catalyst indicating a redistribution of the Co-active sites upon the sonocatalytic reaction. The main reduction peaks are still present, however, their intensity and appearance clearly change. In addition, lower H2-consumption was obtained for the spent catalysts (25.5 μmol/g catalysts) than for the fresh one (64.5 μmol/g catalysts, Table 2), indicating the presence of already reduced Co-species upon the sonocatalytic reaction. This observation further confirms the sonocatalytic mechanism proposed in Fig. 8 involving Co(II)-Co(III) redox process in the reaction mechanism. Absence of Co(II) moieties in the XRD patterns of spent catalyst most likely is related to their low concentrations.

Fig. 10. Kinetics of TOC removal expressed in % (a) and mmol (b) during sonolysis of EDTA homogeneous solution and heterogeneous system with 6.4%wt. Co3O4/TiO2 catalyst and 1%wt. Pt/TiO2 (2 g L⁻¹ of catalyst, 40 °C, 200 mL, 5 mM EDTA, f = 345 kHz, P0,2 = 0.20 W mL⁻¹, 20%O2/Ar). TOC measurement error was about 5%.

Fig. 11. Intermediate products of EDTA solutions (2 g L⁻¹ of 6.4% wt. Co3O4/TiO2 or 1% wt. Pt/TiO2, 40 °C, 200 mL, 5 mM EDTA, f = 345 kHz, P0,2 = 0.20 W mL⁻¹, 20%O2/Ar). Concentration error was about 10%.

Fig. 12. H2-TPR profiles acquired for the 6.4% wt. Co3O4/TiO2 catalyst, before and after the sonocatalytic test.
4. Conclusions

The present study reveals advanced sonochemical degradation of CoOx/TiO2 noble metal-free catalyst in the reaction of EDTA oxidation. Quantitative EDTA degradation can be reached using high-frequency 345 kHz ultrasound and Ar/Ar gas atmosphere at near-room temperature. Surprisingly, the sonochemical activity of CoOx/TiO2 was found to be significantly higher than that of traditional Pt/TiO2 catalyst. This striking phenomenon can be attributed to the stronger interaction of EDTA with CoOx compared to metallic platinum. It is interesting to note that both catalysts are inactive under silent conditions. We suggest that the mechanism of sonochemical process involves the adsorption of EDTA at the cobalt oxide sites, the production of oxidizing radicals during cavitation bubble collapse and redox reaction at CoOx surface intermediated by Co(II)-Co(III) electron transfer. Relative low activation energy of the overall process equal to $E_a = 19$ kJ mol$^{-1}$ pointed out that the thermal effect is related to reagent diffusion rather than to excitation of chemical bonds. Sonochemical and sonochemical degradation of EDTA is accompanied by formation of several acidic intermediates, such as iminodiacetic acid, formic acid, oxalic acid, glycolic acid, and acetic acid. Similar intermediates have been observed during radical degradation of EDTA induced by $\gamma$-irradiation. Despite low catalytic activity toward EDTA oxidation, Pt/TiO2 catalyst improves sonochemical degradation of intermediate products compared to homogeneous solutions. To conclude, these results revealed a new possibility to decompose persistent organic pollutants, such as EDTA, with noble metal-free catalyst.

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Appendix A. Supplementary data

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References