

# Enhancement of ciprofloxacin degradation in aqueous system by heterogeneous catalytic ozonation

Katia González-Labrada, Romain Richard, Caroline Andriantsiferana, Héctor Valdés, Ulises J. Jáuregui-Haza, Marie-Hélène Manero

## ▶ To cite this version:

Katia González-Labrada, Romain Richard, Caroline Andriantsiferana, Héctor Valdés, Ulises J. Jáuregui-Haza, et al.. Enhancement of ciprofloxacin degradation in aqueous system by heterogeneous catalytic ozonation. Environmental Science and Pollution Research, 2018, pp.1-10. 10.1007/s11356-018-3559-9. hal-01980447

## HAL Id: hal-01980447 https://hal.science/hal-01980447

Submitted on 14 Jan 2019  $\,$ 

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



### **Open Archive Toulouse Archive Ouverte**

OATAO is an open access repository that collects the work of Toulouse researchers and makes it freely available over the web where possible

This is an author's version published in: http://oatao.univ-toulouse.fr/21527

Official URL: <u>https://doi.org/10.1007/s11356-018-3559-9</u>

#### To cite this version:

González-Labrada, Katia and Richard, Romain<sup>®</sup> and Andriantsiferana, Caroline<sup>®</sup> and Valdés, Héctor and Jáuregui-Haza, Ulises J. and Manero, Marie-Hélène<sup>®</sup> Enhancement of ciprofloxacin degradation in aqueous system by heterogeneous catalytic ozonation. (2018) Environmental Science and Pollution Research. 1-10. ISSN 0944-1344

Any correspondence concerning this service should be sent to the repository administrator: <u>tech-oatao@listes-diff.inp-toulouse.fr</u>

## Enhancement of ciprofloxacin degradation in aqueous system by heterogeneous catalytic ozonation

Katia González-Labrada<sup>1</sup> • Romain Richard<sup>2</sup> • Caroline Andriantsiferana<sup>2</sup> • Héctor Valdés<sup>3</sup> • Ulises J. Jáuregui-Haza<sup>4</sup> • Marie-Hélène Manero<sup>2</sup>

#### Abstract

Fluoroquinolones are extensively used in medicine due to their antimicrobial activity. Their presence in water inhibits microor-ganism activity in conventional wastewater treatment plants. This study aims to evaluate the technical feasibility of applying heterogeneous catalytic ozonation to eliminate ciprofloxacin (CIP) as a representative of fluoroquinolone antibiotics normally present in municipal wastewater discharges. Experiments were conducted in a semi-batch stirred slurry reactor, using 0.7 L of 100 mg L<sup>-1</sup> CIP aqueous solution, at pH 3 and 30 °C. Experimental results show that single ozonation can easily oxidise CIP molecules (68%) within the first 5 min, leading to the generation of refractory oxidation by-products. However, when hetero-geneous catalytic ozonation is applied using iron oxide supported on MFI synthetic zeolite, total degradation of CIP is observed at 5 min and a higher mineralisation rate is obtained. A novel sequential process is developed for CIP mineralisation. In a first step, a flash single ozonation is applied and CIP molecules are broken down. Then, a catalytic ozonation step is conducted by adding the Fe/MFI catalyst into the reactor. As a result of catalyst addition, 44% of Total Organic Carbon (TOC) is eliminated within the first 15 min, compared to single ozonation where only 13% of TOC removal is reached in the same time. The application of this sequential process to a real wastewater effluent spiked with CIP leads to 52% of TOC removal.

Keywords Advanced oxidation process · Antibiotic · Catalyst · Ciprofloxacin · Ozonation · Wastewater treatment

#### Introduction

Over the last 15 years, pharmaceuticals have been receiving increasing environmental attention due to their negative effects. Thousands of tons of pharmaceuticals are used every year not only to treat human and animal diseases, but also in farming and aquaculture and these compounds have been becoming a serious threat to human health and environment (Gomes et al. 2017). Most of the pharmaceuticals designed to target specific metabolic pathways may become detrimental even at very low concentration. Such chemicals are very persistent and can often be found in wastewater treatment plants (Rivera Utrilla et al. 2013; Padhye et al. 2014; Andreu et al. 2016; Monteiro et al. 2016).

In particular, antibiotics have raised environmental concern because their presence even at low concentration can be harmful and toxic after prolonged exposure (Kümmerer 2003; Martínez 2008; Graham et al. 2010; Knapp et al. 2012; Larsson 2014; Li et al. 2016). Fluoroquinolones group antibiotics are extensively used in medicine due to their antimicrobial activity. Their presence in water inhibits microorganism activity in conventional wastewater treatment plants. After administration, most of the fluoroquinolones are incompletely metabolised and excreted in urine and faeces as a parent compound. Fluoroquinolones are strongly sorbing antibiotics tightly bound to organic matter in faeces, soil and clay minerals (Ötker and Akmehmet-Balcıoğlu 2005). As conventional wastewater treatment systems are unable to reliably remove a large amount of antibiotics (Kümmerer 2003; Rivera Utrilla et al. 2013), new ways of treatment have been developed for some decades.

In recent years, a considerable interest has been focused in the development and application of Advanced Oxidation Processes (AOPs) for the elimination of pharmaceuticals from contaminated waters (Oller et al. 2011: Lastre-Acosta et al. 2015; Sayed et al. 2016; Gomes et al. 2017; Adityosulindro et al. 2017; Velichkova et al. 2017). AOPs involve the generation of hydroxyl radicals that are able to degrade organic pollutants which are highly stable and very difficult to mineralise (Andreozzi et al. 1999; Legube and Leitner 1999; Kasprzyk et al. 2003). AOPs, such as H<sub>2</sub>O<sub>2</sub>/UV, photo-assisted Fenton, and ozonation at high pH values, have been used to oxidise micro-contaminants (Tay and Madehi 2015; Chávez et al. 2016; Ou et al. 2016). However, the efficiency of these homogeneous processes can be reduced by the presence of radical scavengers such as carbonates, bicarbonates, and natural organic matter due to the inhibition of radical chain reactions (von Sonntag and von Gunten, 2012). Therefore, it is necessary to find new process alternatives to overcome such drawbacks.

Recently, heterogeneous AOPs based on the combination of ozone with activated carbon or zeolites have been shown to improve the generation of free radicals. So the impact of inhibition effect produced by radical scavengers is less critical in such a process (Valdés and Zaror, 2006; Valdés et al., 2009 Reungoat et al. 2010; Valdés et al. 2010, 2012a, b; Sui et al. 2012; Aboussaoud et al. 2014; Jalali 2016). In such heterogeneous processes, chemical surface properties of microporous materials have been claimed to play an important role in the reaction mechanisms (Valdés and Zaror, 2006; Valdés et al., 2009; Merle et al. 2010; Valdés et al. 2012a, b). Consequently, heterogeneous catalytic ozonation with natural or synthetic zeolites has been receiving increasing attention in recent years (Martins et al. 2014; Ikhlaq et al. 2015; Vittenet et al. 2015; Gomes et al. 2018).

Lately, the use of zeolites containing iron species as catalysts has gained increasing interest because of the combination of their adsorptive and catalytic properties. Among catalysts, iron type catalysts can be recommended because of the abundance of this metal on earth. Iron type catalysts have already been used in ozonation processes to remove different compounds such as phenol, aniline, carboxylic acids, chlorobenzene, chlorophenols, dyes or natural organic matter (Beltran et al. 2005). Fe-containing zeolites are interesting candidates as heterogeneous catalyst because of their high catalytic activity in the oxidation of organic compounds, with minimal iron leaching (Chen et al. 2008; Gonzalez Olmos et al. 2011; Velichkova et al. 2017).

This work aims to study heterogeneous catalytic ozonation of antibiotic-contaminated waters, using ZSM-5 zeolite (Zeolite Framework Type ZSM-FIve; abbreviation: MFI.) doped with iron oxide nanoparticles. CIP was used here as a representative of a group of fluoroquinolone antibiotics widely used in human and animal therapy. CIP is a non-biodegradable antibiotic that has been detected in wastewater treatment plant effluents in concentrations from 5.6  $\mu$ g L <sup>1</sup> to 6.5 mg L <sup>1</sup> (Kümmerer et al. 2000; Hughes et al. 2012). As an antibiotic, CIP is of particular concern because its presence in natural waters could contribute to the rise of antibiotic resistance in microorganisms (Kümmerer 2003; Graham et al. 2010; Knapp et al. 2012; Martínez 2008; Larsson 2014). Very few studies have been found on CIP removal. Sorption on activated sludge and biodegradation catalysed by magnetite nanoparticles seem to give interesting results (Yang et al. 2017). More recently, membranebased processes have been focused on (Sadhwani et al. 2018). However, information that deals with the elimination of CIP and its oxidisation by-products using processes based on the combination of ozone and zeolites is not abundant. In particular in this study, the effect of single ozonation, single adsorption, single heterogeneous catalytic ozonation, and the application of a sequential process comprising of two step (single flash ozonation followed by catalytic ozonation) are evaluated in terms of CIP mineralisation. Finally, all processes are assessed using real effluent spiked with CIP.

#### **Materials and methods**

#### Materials

Ciprofloxacin (1-cyclopropyl-6-fluoro-4-oxo-7-piperazin-1ylquinoline-3-carboxylic acid, named here as CIP) was supplied by Sigma-Aldrich (St. Louis, MO, United States) with 99% purity. In a first step of this study, a synthetic wastewater: was used. It was prepared by dissolving CIP in deionised water. A complete geometric representation of the molecular structure of CIP, its solubility in water and its  $pK_a$  value are given in Table 1.

Table 1 Physical chemical properties of CIP



The adjustment of pH adjustment was carried out using 1 mol L<sup>1</sup> solution of HCl from Chem-Lab. Acetonitrile (HPLC quality) and o-phosphoric acid (85%) were provided by Merck (Darmstadt, Germany) and Fisher (Pittsburgh, United States), respectively, and were used in high-performance liquid chromatography (HPLC) analyses.

ZSM-5 zeolite with a framework type MFI doped with iron oxide (Fe<sub>2</sub>O<sub>3</sub>) nanoparticles (referred as Fe/MFI) was used here as a catalyst and was bought from Süd-Chemie AG (Munich, Germany) with a silicon/aluminium ratio of 27. This kind of ZSM-5 zeolite exhibits a three-dimensional pore network with 10-membered straight and sinusoidal ring channels and apertures of  $5.3 \times 5.6$  Å and  $5.1 \times 5.5$  Å, respectively (Pearce 1975). Physicochemical properties of Fe/MFI zeolite used in this study are listed in Table 2.

The physical-chemical characteristics of filtered real wastewater plant effluent sample, taken from Nailloux wastewater treatment plant (Occitanie, France), are shown in Table 3. Five-day biological oxygen demand (BOD<sub>5</sub>) analyses were assessed at the Departmental Laboratory, LD 31, Toulouse, France. In the experiments where real wastewater was applied; it was spiked with 100 mg L<sup>-1</sup> of CIP and used within 24 h after collection for assays of single ozonation and heterogeneous catalytic ozonation.

#### **Experimental procedures**

All experiments were performed at pH 3 and  $30 \pm 1$  °C in a semi-batch stirred slurry reactor (Ø100 mm x 100 mm) containing 0.7 L of a CIP aqueous solution (100 mg L<sup>1</sup>). Figure 1 displays a schematic representation of the experimental ozonation setup. Ozone was produced from dry pure oxygen supplied by Air Liquide (Paris, France), using a 4-HC, WEDECO ozone generator rated at 1.44 gO<sub>3</sub> h<sup>-1</sup>. A continuous flow of ozone (30 L h<sup>-1</sup>, 48 gO<sub>3</sub> m<sup>-3</sup>) was introduced into the reactor by means of a stainless steel diffuser. During single ozonation experiments at pH 3, no hydroxyl radical formation is expected (Hoigné 1998; Von Sonntag and von Gunten 2012). Under such conditions, radical oxidation route is almost inhibited and oxidation reactions are mainly due to direct action of ozone molecules. Samples were regularly collected in order to follow kinetic evolutions. Samples were filtrated and taken to determine the residual CIP concentration and Total Organic Carbon concentration (TOC).

**Table 3** Physical chemical characteristics of real and spiked wastewater matrix with  $100 \text{ mg L}^{-1}$  CIP and adjusted to pH 3

| Parameter        | Unit                             | Wastewater plant effluent | Spiked wastewater |
|------------------|----------------------------------|---------------------------|-------------------|
| рН<br>TOC        | mgO <sub>2</sub> L <sup>1</sup>  | 6.5<br>15.28              | 3.0<br>76.03      |
| BOD <sub>5</sub> | mgO <sub>2</sub> L <sup>-1</sup> | <2                        | 110               |

Four different processes were studied. First, single ozonation (i) and heterogeneous catalytic ozonation using 1 g L<sup>-1</sup> of Fe/MFI zeolite (ii) were conducted. After that, an original process was evaluated, with two different sequences: a quick ozonation step in order to break the CIP molecule (run during 5 minutes) followed by either an adsorption onto Fe/MFI zeolite (iii) or by a catalytic ozonation using Fe/MFI zeolite (iv). The influence of the catalyst dosage was assessed, applying different amounts of Fe/ZSM5 from 0 to 10 g L<sup>-1</sup> (v). Experiments were run in triplicate.

#### **Analytical methods**

CIP concentration was determined by HPLC (Varian ProStar 310 model). A RP-18 column (Super Sphere 100 model, 250 mm × 4.6 mm; 5  $\mu$ m) at 40 °C connected to an UV/VIS diode array detector and set at 278 nm was used for CIP detection. A mix of 15% of acetonitrile/acidified water (*v*/*v*) was applied as a mobile phase rated at 1.5 mL min <sup>1</sup>. A volume of 20  $\mu$ L was injected and CIP-eluted at around 9 min. A calibration curve ( $R^2 = 0.99$ ) was obtained using external standard, prepared with known concentrations of CIP.

Mineralisation of CIP was followed by TOC analysis, using a TOC-LCPN Shimadzu analyser equipped with a non-dispersive infrared detector. The injection volume was 50  $\mu$ L. Combustion reaction was achieved in a quartz pipe at 680 °C with platinum as a catalyst. Oxygen was used as a carrier gas with a flow rate of 150 mL min <sup>1</sup>.

#### **Results and discussion**

#### CIP adsorption onto the Fe/MFI zeolite

Preliminary experimental studies of kinetic adsorption show that the CIP adsorption onto the Fe/MFI zeolite is very slow.

| Table 2   | Physical chemical    |
|-----------|----------------------|
| propertie | es of Fe/MFI zeolite |
| (Velichk  | ova 2014)            |

| Commercial name | SiO <sub>2</sub> /<br>Al <sub>2</sub> O <sub>3</sub> | Fe<br>(%) | $S_{BET}$<br>$(m^2 g^{-1})$ | $V_{meso} (cm^3 g^{-1})$ | $V_{micro} (cm^3 g^{-1})$ | pH <sub>PZC</sub> <sup>a</sup> |
|-----------------|--|-----------|-----------------------------|--------------------------|---------------------------|--------------------------------|
| Fe SH 27        | 27   | 3.4       | 329                         | 0.05                     | 0.13                      | 2.6                            |

<sup>a</sup> pH of the point of zero charge ( $pH_{PZC}$ ) determined using the acidimetric alkalimetric titration method proposed by Stumm and Morgan (1996) Fig. 1 Schematic diagram of the experimental ozonation setup: (1) oxygen cylinder; (2) ozone generator; (3) gas flow meter; (4) ozone gas analyser/reactor bypass valves; (5) semi batch stirred slurry reactor; (6) sampling valve; (7) water collection trap; (8) ozone gas analyser; (9) ozone destruction system



Less than 5% of CIP removal is obtained in 1 h. This result could be explained by a size exclusion process because molecular dimensions of CIP are larger (13.2 Å; 7.8 Å; 3.7 Å) (Stewart 2012) than the dimensions of channels and pores of the zeolite  $5.3 \times 5.6$  Å and  $5.1 \times 5.5$  Å (Pearce 1975). At pH 3, the main part of CIP molecules carry on an overall positive charge (CIP<sup>3+</sup>) with a pKa<sub>1</sub> value of 3.64. Taking into account the low pH<sub>PZC</sub> (2.6), zeolite is able to exchange only a few cations in these conditions.

#### CIP homogeneous and heterogeneous ozonation

Figure 2 shows the variation of the dimensionless concentration of CIP and TOC as a function of time, during the application of single ozonation and heterogeneous catalytic ozonation promoted by Fe/MFI zeolite. A very fast degradation of the pharmaceutical compound is observed. A nearly total transformation (98%) of CIP is achieved in less than 15 min by single ozonation. However, an important amount of TOC (87%) still remains in the aqueous solution. The addition of 1 g L<sup>1</sup> of Fe/MFI zeolite enhances the oxidation kinetics during the first 30 min. As it can be seen in Fig. 2, a total degradation of CIP is obtained after 5 min of reaction time. A 42% of the TOC is removed after 15 min in comparison to 13% obtained when single ozonation is applied. These results could be mainly due to two effects: (1) the formation of radicals due to the adsorption of ozone onto the catalyst surface and its further decomposition leading to the formation of surface-bound O-radicals and hydroxyl radicals, which are active oxidative species (Kasprzyk-Hordern et al. 2003), (2) and the enhancement in the adsorption of some of ozonation by-products with smaller size, being able to diffuse and being adsorbed onto zeolite active sites. Hence an increase in the removal rate of CIP takes place by catalytic ozonation at the solid-liquid interface, and by CIP homogenous oxidation reactions in the liquid phase closed to the zeolite surface.



Fig. 2 Comparison of the variation of TOC and CIP concentrations during homogeneous and heterogeneous ozonation. Open markers represent single ozonation; filled markers mean heterogeneous catalytic ozonation promoted by Fe/MFI zeolite (1 g L<sup>1</sup>): ( $\circ$ / $\bullet$ ) dimensionless

concentration of CIP and  $(\Delta/\blacktriangle)$  dimensionless concentration of TOC. Experimental conditions: 0.7 L of an aqueous solution of CIP with 100 mg L<sup>1</sup> of initial concentration, continuous inlet flow of ozone (30 L h<sup>1</sup>, 48 gO<sub>3</sub> m<sup>3</sup>) at pH 3 and 30 °C



Based on previous studies that reveal the formation of oxidation by-products during the ozonation of CIP molecule such as dealkylated and hydroxylated compounds, with smaller size and molecular mass than CIP molecule (De Witte et al. 2008; Liu et al. 2012), a sequential process was tested: a flash ozonation of the CIP solution during 5 min to transform large CIP molecules, followed by an adsorption process to remove small size oxidation by-products. Unfortunately, these ozonation by-products are hardly adsorbed, as it can be seen in Fig. 3. However, a better result is obtained when a catalytic ozonation process is applied instead of single adsorption. Under such conditions, a noticeable enhancement on the efficiency is found; up to 50% of TOC is removed after 40 min of catalytic ozonation. Similar results were obtained by Sui and co-workers (Sui et al. 2012). A 38% reduction of TOC was achieved using catalytic ozonation promoted by MnO<sub>2</sub>/MWCN, where a 30% of CIP remained adsorbed on the surface of MnO<sub>2</sub>/MWCN. Therefore, the 25% of TOC removed from single ozonation compared to the 39% obtained after 25 min indicates that the presence of Fe/MFI zeolite increases TOC removal. Fe/MFI zeolite seems to provide a surface where CIP oxidation by-products are adsorbed and react with radicals generated from ozone interaction with active sites of zeolite surface.

In order to evaluate this behaviour, different Fe/MFI zeolite concentrations were used. Figure 4 shows the effects of the Fe/MFI zeolite concentration  $(0-10 \text{ g L}^{-1})$  in the TOC removal of



Fig. 4 Effect of the dosage of Fe/MFI zeolite during two different sequential processes (× stands for single flash ozonation during 5 min). Five minute single flash ozonation followed by 40 min of adsorption onto Fe/MFI zeolite using ( $\blacklozenge$ ) 1 g L<sup>1</sup>, ( $\diamond$ ) (5 g L<sup>1</sup>) and ( $\Box$ ) 10 g L<sup>1</sup>. Five minute single flash ozonation followed by 45 min of heterogeneous

catalytic ozonation promoted by Fe/MFI zeolite using ( $\blacktriangle$ ) 1 g L<sup>1</sup>, ( $\Delta$ ) 5 g L<sup>1</sup> and ( $\blacksquare$ ) 10 g L<sup>1</sup>. Experimental conditions: 0.7 L of an aqueous solution of CIP with 100 mg L<sup>1</sup> of initial concentration, continuous inlet flow of ozone (30 L h<sup>-1</sup>, 48 gO<sub>3</sub> m<sup>-3</sup>) at pH 3 and 30 °C

the solution after 5 min of single flash ozonation. As it can be observed, TOC removal rate is improved as catalyst concentration is raised. TOC removal is enhanced for Fe/MFI zeolite load of 10 g L<sup>1</sup> (44%) within the first 15 min if compared with zeolite load of 1 g L<sup>1</sup> (27%).

During heterogeneous catalytic ozonation, TOC removal rate is increased to higher values than those obtained when a single ozonation process is followed by a single adsorption step, as it is shown in the Fig. 4. In this process, zeolite not only provides a contact surface where the CIP oxidation by-products and ozone are adsorbed, but also participates in ozone decomposition, yielding a higher quantity of hydroxyl radicals that increases TOC removal rate. Ozone could react with acid surface groups of metal oxide. Thanks to its resonance structure, ozone features high electron density on one of its oxygen atoms resulting in strong affinity to Lewis acid sites with the subsequent decomposition (Kasprzyk-Hordern et al. 2003). Adsorbed ozone on Lewis acid sites could convert to radicals ( $O_2 \cdot$  or HO·), acting as initiators and/or promoters in free radical chain reactions (Valdés et al. 2010).

A kinetic study of CIP direct and heterogeneous catalytic ozonation with multi-walled carbon nano-tubes supported manganese oxides (MnO<sub>2</sub>/MWCNT) as a catalyst has shown similar results, using Monte Carlo simulation, with respect to the effect of catalyst load (Jalali 2016). This method was applied with the purpose of modelling the experimental data for the ozonation of CIP over MnO<sub>2</sub> and MnO<sub>2</sub>/MWCNT and (Sui et al. 2012). Under the experimental conditions investigated (pH 7, 25 °C, CIP concentration of 10 mg L<sup>1</sup>, and different initial amount of MnO<sub>2</sub> and MnO<sub>2</sub>/MWCNT) a positive effect on CIP degradation is attained when the highest dose of MnO<sub>2</sub>/ MWCNT is applied. A lower pH value shows to be more favourable for CIP degradation in the catalytic ozonation. The efficiency of ozone treatment in the decontamination and regeneration of contaminated natural zeolite has also been evaluated by other research groups (Ötker and Akmehmet-Balcıoğlu 2005). Kinetic experiments for zeolite were performed using 200 mg L<sup>1</sup> of enrofloxacin solutions at pH values of 5, 7 and 10. It is claimed that a sufficient ozone concentration is able to decompose enrofloxacin adsorbed on natural zeolite, where enrofloxacin sorption interactions with natural zeolite are controlled by ionic functional groups resulting in ionic binding of the antibiotic and the zeolite (Ötker and Akmehmet-Balcioğlu 2005). Catalytic ozonation in the removal of ibuprofen from water using ZSM-5 zeolites has been also investigated by Ikhlaq and collaborators (Ikhlaq et al. 2014). Different SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> ratios were used. under different pH conditions in the range between 3 and 13 (490 mL of water spiked with 20 mg L  $^{1}$  of ibuprofen, 5 g of catalyst, applying 0.5 mg min<sup>1</sup> of ozone flow rate, at 20°C). Results indicate that ibuprofen shows higher adsorption on zeolites at pH 3 than at pH 7.2 and 13. Furthermore, adsorption capacity of zeolite resulted to be dependant of the  $SiO_2/Al_2O_3$  ratio. Catalytic ozonation using ZSM-5 zeolites is effective at pH 3 and 7.2, with the highest activity observed in the case of Z25 zeolites. However, zeolites did not show any activity at a basic pH of 13. This may be due to the high concentration of OH ions that are responsible for the high rates of aqueous ozone decomposition at pH 13 (Ikhlaq et al. 2014). Moreover, some other results show that heterogeneous catalytic ozonation using natural zeolite increases methylene blue removal rate with respect to the single ozonation process (Valdés et al. 2010, 2012a, b).

#### **Kinetic modelling**

In this section, a kinetic model was developed taking into account a synergistic mechanism that combines adsorption and oxidation reactions at the zeolite–liquid interface and oxidation reactions in the bulk-liquid phase. Similar approach has been proposed for cationic dye removal from contaminated waters using heterogeneous catalytic ozonation in the presence of natural zeolite, acid-treated natural zeolite and volcanic sand (Valdés et al. 2010, 2012a, b).

Hence, catalytic ozonation of CIP oxidation by-products (represented here as TOC) in the presence of Fe/MFI zeolite might include both direct reaction of molecular ozone and indirect reactions involving non-selective free radicals coming from ozone decomposition taking place in the bulk-liquid phase, adsorption of low molecular size CIP oxidation by-products at zeolite active sites and surface reactions of adsorbed complexes with molecular ozone and/or with free radicals. Assuming that the process rate is controlled by chemical reactions, ozonation of CIP oxidation by-products in the presence of Fe/MFI zeolite could be represented by a simplified and unbalanced reaction mechanism, comprising the two phases, as follows:

Bulk liquid phase reactions:

$$TOC + O_3 \xrightarrow{k_2} CO_2 + H_2O - \frac{d[TOC]_1}{dt}$$
$$= k_1[O_3][TOC] \tag{1}$$

$$TOC + R \xrightarrow{k_2} CO_2 + H_2O - \frac{d[TOC]_2}{dt} = k_2[R][TOC]$$
(2)

where, [TOC], [O<sub>3</sub>] and [R] represent the concentration of CIP oxidation by-products analytically measured as Total Organic Carbon, dissolved ozone, and ozone decomposition reactive species in water (e.g. HO<sup>,</sup>, HO<sub>2</sub>·/O<sub>2</sub>·, O<sub>3</sub>·, H<sub>2</sub>O<sub>2</sub>/HO<sub>2</sub>), respectively.

*Surface reactions at zeolite active sites*  $(Z^*)$ :

$$TOC + Z^* \stackrel{k_3}{\underset{k_3}{\Rightarrow}} Z^* TOC \quad r_3 \quad k_3 \{Z^*\}[TOC] \ k_3 \{Z^* \ TOC\}$$
  
(3)

$$Z^* TOC + O_3 \xrightarrow{k_4} CO_2 + H_2O \quad r_4 = k_4[O_3] \{Z^* TOC\}$$

$$(4)$$

$$Z^* - TOC + R \xrightarrow{k_5} CO_2 + H_2O \quad -r_5 = k_5[R] \{Z^* - TOC\}$$
(5)

The overall rate of change of the surface complex concentration of CIP oxidation by-products  $\{Z^*-TOC\}$  could be expressed as follows:

$$\frac{d\{Z^* \ TOC\}}{dt} = k_3\{Z^*\}[TOC] \ (k_3 + k_4[O_3] + k_5[R])\{Z^* \ TOC\}$$
(6)

where {*Z*\*} and {*Z*\*–TOC} stand for surface concentration of zeolite active surface sites and surface complex of CIP oxidation by products, respectively.  $k_1$  denotes rate constant of homogeneous reaction between CIP oxidation by-products and ozone (M<sup>1</sup> s<sup>-1</sup>),  $k_2$  is the rate constant of the homogeneous reaction between CIP oxidation by-products and radicals generated by ozone decomposition (M<sup>-1</sup> s<sup>-1</sup>),  $k_3$  is the forward rate constant of surface complex formation of CIP oxidation byproducts (M<sup>-1</sup> s<sup>-1</sup>),  $k_3$  is the reverse rate constant of surface complex formation of CIP oxidation by-products (M<sup>-1</sup> s<sup>-1</sup>),  $k_4$ is the surface reaction rate constant between CIP oxidation byproducts and ozone (M<sup>-1</sup> s<sup>-1</sup>),  $k_5$  is the surface reaction rate constant between CIP oxidation by-products and radicals generated by ozone decomposition (M  $^{1}$  s  $^{1}$ ).

Applying the kinetic modelling recently proposed by Valdés and co-workers (Valdés et al. 2012a, b), the overall rate expression of catalytic ozonation of CIP oxidation by-products using ZSM-5 zeolite doped with iron oxide nano-particles could be represented as a sum of bulk-liquid and surface reactions at zeolite active sites:

$$-\mathbf{r}_{overall} = \mathbf{r}_{bulk} + \mathbf{r}_{surface} = \left(\mathbf{k}_{bulk} + \mathbf{k}_{surface}\left\{\mathbf{Z}^*\right\}\right)[TOC] (7)$$

Integrating, with initial conditions  $[TOC]_t = [TOC]_0$  leads to:

$$n\frac{[\text{TOC}]_{t}}{[\text{TOC}]_{0}} = -k_{\text{overall}}t$$
(8)

with,

$$\mathbf{k}_{overall} = \mathbf{k}_{bulk} + \mathbf{k}_{surface} \{ \mathbf{Z}^* \}$$
(9)

where,  $r_{bulk}$  and  $r_{surface}$  are removal rates of CIP oxidation byproducts in the bulk-liquid phase and at the active sites of zeolite surface, respectively.  $k_{bulk}$  and  $k_{surface}$  represent pseudo-first-order rate constant of ozonation reactions taking place in the bulk-liquid phase and pseudo-second-order rate constant of the degradation of CIP oxidation by-products due to surface ozonation reactions, respectively. [TOC]<sub>t</sub> and [TOC]<sub>0</sub> are the concentration of CIP oxidation by-products globally determined as Total Organic Carbon at any time t and time zero, respectively. Such kinetic model is limited in this study to describe catalytic ozonation of CIP oxidation byproducts using ZSM-5 zeolite doped with iron oxide nanoparticles without mass transfer limitations.

Fig. 5 Variation of pseudo first order rate constants of overall reactions of CIP oxidation by products ( $k_{overall}$ ) as a function of zeolite concentration. Experimental conditions: 0.7 L of an aqueous solution of CIP with 100 mg L<sup>-1</sup> of initial concentration, continuous inlet flow of ozone (30 L h<sup>-1</sup>, 48 gO<sub>3</sub> m<sup>-3</sup>) at pH 3 and 30 °C



| Treatment/time   | TOC $(mg O_2 L^{-1})$ | TOC<br>removal (%) | $\begin{array}{c} BOD_5 \\ (mg \ O_2 \ L^{-1}) \end{array}$ | BOD <sub>5</sub><br>removal (%) |
|--|-----------------------|--------------------|---|---------------------------------|
| Effluent   | 76                    |                    | 110   |                                 |
| Single ozonation of CIP (15 min)   | 41                    | 46                 | <2  | > 98                            |
| Single flash ozonation of CIP (5 min) followed by the adsorption of CIP oxidation by products onto Fe/MFI (1 g L $^{1}$ ) (15 min) | 48                    | 36                 | 6   | 94                              |
| Single flash ozonation (5 min) followed by heterogeneous catalytic ozonation with Fe/MFI (1 g L $^{1}$ ) (15 min)                  | 36                    | 52                 | <2  | 98                              |

The contribution of surface and bulk-liquid phase reactions to the catalytic ozonation of CIP oxidation by-products using Fe/MFI zeolite is estimated using Eq. (9), at pH 3. A series of experimental runs were performed with different zeolite concentration (see Fig. 5) and pseudo-first-order rate constants of overall oxidation reactions of CIP by-products,  $k_{overall}$ , were obtained from initial rate data.

Figure 5 illustrates the results of plotting the estimated koverall values as a function of zeolite concentration. Thus, the value of the pseudo-second-order rate constant of the degradation of CIP oxidation by-products due to surface ozonation reactions ( $k_{surface} = 0.0468 \times 10^{-3}$  g<sup>-1</sup> L s<sup>-1</sup>) and pseudo-firstorder rate constant of ozonation reactions taking place in the bulk-liquid phase ( $k_{bulk} = 0.4129 \times 10^{-3}$  s<sup>-1</sup>) are obtained from the line slope and the intercept of the line with "Y" axis, respectively.

Fe/MFI zeolite not only offers a contact surface for the adsorption and oxidation of CIP oxidation by-products but also takes part in aqueous ozone decomposition. Active surface sites of Fe/MFI zeolite could contribute to ozone decomposition, acting as initiators and/or promoters of radical chain reactions. Ozone could be adsorbed and decomposed at acidic sites of Fe/MFI zeolite such as acidic OH groups (Brønsted acid centres), "true" Lewis sites (aluminium-containing extraframework species), generating surface radicals (Ni and Chen 2001; Einaga and Futamura 2005; Kwong et al. 2008; Valdés et al. 2012a, b) that react with adsorbed CIP oxidation byproducts. A radical chain reaction continues in the bulkliquid phase leading to the formation of an ozonide anion,  $O_3$ , by the transfers of an electron from  $O_2$  to another ozone molecule, which is the chain reaction promoter and produces HO· radicals, contributing to organic oxidation in the liquid phase (Valdés et al. 2012a).

## Influence of treatment process on effluent's mineralisation and biodegradability

Table 4 shows experimental results of TOC and BOD<sub>5</sub> analyses after different sequential treatment processes using real wastewater spiked with CIP and adjusted to pH 3. As it can be observed in Table 4, BOD<sub>5</sub> value increases when CIP is added, being CIP the major organic compound in this matrix.

Each sequential treatment process reduces the value of BDO<sub>5</sub> from 110 to less than 2 mg  $O_2$  L<sup>-1</sup> after 15 min. When wastewater sample is ozonated during 5 min and then adsorption onto the zeolite takes place, the BOD<sub>5</sub> is slightly superior with 6 mg L<sup>-1</sup> compared to the other treatments. In the experiments, TOC is reduced from 41 to 36 mg  $O_2$  L <sup>1</sup> when single flash ozonation (5 min)-followed by heterogeneous catalytic ozonation (15 min) is applied compared to single ozonation (15 min) corresponding in both cases to the reduction of the BOD<sub>5</sub> compared to the initial BOD<sub>5</sub> of the effluent. These experiments display in a better way the results than those obtained without matrix effect. It is well known that the stability of ozone is mainly determined by its reaction with organic matter, carbonates and bicarbonates normally presented in water. They could act as inhibitors or radicals scavengers, prolonging the lifetime of ozone at typical pH values of wastewater (pH  $\leq$  8.5) (Von Sonntag and von Gunten 2012).

#### Conclusions

The results presented here demonstrate that single ozonation treatment can easily oxidise CIP molecules; however, low TOC removal is obtained due to the generation of refractory CIP oxidation by-products. An improvement in the mineralisation rate is achieved when heterogeneous catalytic ozonation is applied, using ZSM-5 zeolite doped with iron oxide. An increase on TOC removal rate occurs, from 13% for single oxidation to 42% for catalytic oxidation. A new sequential process is proposed in order to improve the efficiency of the mineralisation. In a first step, a single flash ozonation breaks down ciprofloxacin molecules. Then, a catalytic ozonation step promoted by Fe/ZSM-5 zeolite leads to a noticeable enhancement on the efficiency, achieving up to 50% of TOC elimination after 40 min of heterogeneous catalytic ozonation. In this heterogeneous system, the elimination of CIP oxidation by-products could be described by a synergistic mechanism that combines adsorption and oxidation reactions at the zeolite-liquid interface and oxidation reactions in the bulk-liquid phase. A sequential process of two-steps, conformed by flash ozonation followed by catalytic ozonation, could be a hopeful advanced oxidation process for the removal of non-biodegradable complex molecules with a high molecular mass and size.

Acknowledgements This research was possible thanks to the financial support of Laboratoire de Génie Chimique, Université de Toulouse and the project TATARCOP of Instituto Superior de Tecnologías y Ciencias Aplicadas (InSTEC) Universidad de La Habana. K. González Labrada expresses her gratitude to the Collaboration Services of the French Embassy in Cuba. H. Valdés gratefully acknowledges funding under CNRS Délégation Midi Pyrénées contract 618035.

#### References

- Aboussaoud W, Manero MH, Pic JS, Debellefontaine H (2014) Combined ozonation using alumino silica materials for the removal of 2,4 dimethylphenol from water. Ozone Sci Eng 36:221 228
- Adityosulindro S, Barthe L, González Labrada K, Jáuregui Haza UJ, Delmas H, Julcour C (2017) Sonolysis and sono Fenton oxidation for removal of ibuprofen in (waste) water. Ultrason Sonochem 39: 889 896
- Andreozzi R, Caprio V, Insola A, Marotta R (1999) Advanced oxidation processes (AOP) for water purification and recovery. Catal Today 53:51 59
- Andreu V, Gimeno García E, Pascual JA, Vazquez Roig P, Picó Y (2016) Presence of pharmaceuticals and heavy metals in the waters of a Mediterranean coastal wetland: potential interactions and the influ ence of the environment. Sci Total Environ 540:278 286
- Beltran FJ, Rivas FJ, Montero de Espinosa R (2005) Iron type catalysts for the ozonation of oxalic acid in water. Water Res 46:3553 3564
- Chávez AM, Rey A, Beltrán FJ, Álvarez PM (2016) Solar photo ozona tion: a novel treatment method for the degradation of water pollut ants. J Hazard Mater 317:36 43
- Chen A, Ma X, Sun H (2008) Decolorization of KN R catalyzed by Fe containing Y and ZSM 5 zeolites. J Hazard Mater 156:568 575
- De Witte B, Dewulf J, Demeestere K, De Ruyck M, Van Langenhove H (2007) Critical points in the analysis of ciprofloxacin by high per formance liquid chromatography. J Chromatogr A 1140:126 130
- Einaga H, Futamura S (2005) Oxidation behavior of cyclohexane on alumina supported manganese oxides with ozone. Appl Catal B Environ 60:49 55
- Gomes JF, Costa R, Quinta Ferreira RM, Martins RC (2017) Application of ozonation for pharmaceuticals and personal care products remov al from water. Sci Total Environ 586:265 283
- Gomes JF, Quinta Ferreira ME, Costa R, Quinta Ferreira RM, Martins RC (2018) Paraben degradation using catalytic ozonation over vol canic rocks. Environ Sci Pol 25:7346 7357
- Gonzalez Olmos R, Holzer F, Kopinke FD, Georgi A (2011) Indications of the reactive species in a heterogeneous Fenton like reaction using Fe containing zeolites. Appl Catal A 398:44 53
- Graham DW, Olivares Rieumont S, Knapp CW, Lima L, Werner D, Bowen E (2010) Antibiotic resistance gene abundances associated with waste discharges to the Almendares River near Havana, Cuba. Environ Sci Technol 45:418 424
- Hoigné J (1998) Chemistry of aqueous ozone and transformation of pol lutants by ozonation and advanced oxidation processes. Quality and treatment of drinking water II. Springer, Berlin, pp 83 141
- Hughes SR, Kay P, Brown LE (2012) Global synthesis and critical eval uation of pharmaceutical data sets collected from river systems. Environ Sci Technol 47:661 677
- Ikhlaq A, Brown DR, Kasprzyk Hordern B (2014) Catalytic ozonation for the removal of organic contaminants in water on alumina. Appl. Catal B Environ 155:110 122

- Jalali HM (2016) Kinetic study of antibiotic ciprofloxacin ozonation by MWCNT/MnO<sub>2</sub> using Monte Carlo simulation. Mater Sci Eng C Mater Biol Appl 59:924 929
- Kasprzyk Hordern B, Ziólek M, Nawrocki J (2003) Catalytic ozonation and methods of enhancing molecular ozone reactions in water treat ment. Appl Catal B Environ 46:639–669
- Knapp CW, Lima L, Olivares Rieumont S, Bowen E, Werner D, Graham DW (2012) Seasonal variations in antibiotic resistance gene trans port in the Almendares River, Havana, Cuba. Front Microbiol 3:396
- Kümmerer K (2003) Significance of antibiotics in the environment. J Antimicrob Chemother 52:5 7
- Kümmerer K, Al Ahmad A, Mersch Sundermann V (2000) Biodegradability of some antibiotics, elimination of the genotoxicity and affection of wastewater bacteria in a simple test. Chemosphere 40:701 710
- Kwong C, Chao CY, Hui K, Wan M (2008) Removal of VOCs from indoor environment by ozonation over different porous materials. Atmos Environ 42:2300 2311
- Larsson DJ (2014) Antibiotics in the environment. Ups J Med Sci 119: 108 112
- Lastre Acosta AM, Cruz González G, Nuevas Paz L, Jáuregui Haza UJ, Teixeira AC (2015) Ultrasonic degradation of sulfadiazine in aque ous solutions. Environ Sci Pol 22:918 925
- Legube B, Leitner NKV (1999) Catalytic ozonation: a promising ad vanced oxidation technology for water treatment. Catal Today 53: 61 72
- Li X, Wang W, Dou J, Gao J, Chen S, Quan X, Zhao H (2016) Dynamic adsorption of ciprofloxacin on carbon nanofibers: quantitative mea surement by in situ fluorescence. J Wat Proc Eng 9:14 20
- Lin CE, Deng YJ, Liao WS, Sun SW, Lin WY, Chen CC (2004) Electrophoretic behavior and pKa determination of quinolones with a piperazinyl substituent by capillary zone electrophoresis. J Chromatogr A 1051:283 290
- Liu C, Nanaboina V, Korshin GV, Jiang W (2012) Spectroscopic study of degradation products of ciprofloxacin, norfloxacin and lomefloxacin formed in ozonated wastewater. Water Res 46:5235 5246
- Martínez JL (2008) Antibiotics and antibiotic resistance genes in natural environments. Sci 321:365 367
- Martins RC, Ramos CM, Quinta Ferreira RM (2014) Low cost catalysts to enhance ozone action on the depuration of olive mill wastewaters. Ind Eng Chem Res 53:15357 15368
- Merle T, Pic JS, Manero MH, Mathe S, Debellefontaine H (2010) Influence of activated carbons on the kinetics and mechanisms of aromatic molecules ozonation. Catal Today 151:166 172
- Monteiro MA, Spisso BF, dos Santos JRMP, da Costa RP, Ferreira RG, Pereira MU, da Silva Miranda T, de Andrade BRG, d'Avila LA (2016) Occurrence of antimicrobials in river water samples from rural region of the State of Rio de Janeiro, Brazil. J Environ Prot 7:230 241
- Ni C, Chen J (2001) Heterogeneous catalytic ozonation of 2 chlorophenol queous solution with alumina as a catalyst. Water Sci Technol 43:213 220
- Nowara A, Burhenne J, Spiteller M (1997) Binding of fluoroquinolone carboxylic acid derivatives to clay minerals. J Agric Food Chem 45: 1459–1463
- Oller I, Malato S, Sánchez Pérez JA (2011) Combination of advanced oxidation processes and biological treatments for wastewater decon tamination a review. Sci Total Environ 409:4141 4166
- Ötker HM, Akmehmet Balcıoğlu I (2005) Adsorption and degradation of enrofloxacin, a veterinary antibiotic on natural zeolite. J Hazard Mater 122:251 258
- Ou H, Ye J, Ma S, Wei C, Gao N, He J (2016) Degradation of ciproflox acin by UV and UV/H<sub>2</sub>O<sub>2</sub> via multiple wavelength ultraviolet light emitting diodes: effectiveness, intermediates and antibacterial activ ity. Chem Eng J 289:391 401

- Padhye LP, Yao H, Kung'u FT, Huang CH (2014) Year long evaluation on the occurrence and fate of pharmaceuticals, personal care prod ucts, and endocrine disrupting chemicals in an urban drinking water treatment plant. Water Res 51:266 276
- Pearce H (1975) Zeolite molecular sieves structure, chemistry and use: by DA Breck, Wiley Interscience, Elsevier
- Reungoat J, Pic JS, Manero MH, Debellefontaine H (2010) Oxidation of nitrobenzene by ozone in the presence of faujasite zeolite in a con tinuous flow gas liquid solid reactor. Water Sci Technol 62:1076 1083
- Rivera Utrilla JM, Sánchez Polo MÁ, Ferro García G, Prados J, Ocampo Pérez R (2013) Pharmaceuticals as emerging contaminants and their removal from water. A review. Chemosphere 93:1268 1287
- Sadhwani JJ, El Kori AN, Melian Martel N, Del Rio Gamero B (2018) Removal of ciprofloxacin from seawater by reverse osmosis. J Environ Manag 217:337 345
- Sayed M, Ismail M, Khan S, Tabassum S, Khan HM (2016) Degradation of ciprofloxacin in water by advanced oxidation process: kinetics study, influencing parameters and degradation pathways. Environ Technol 37:590 602
- Stewart JJ (2012) Mopac 2012. Stewart Computational Chemistry, Colorado Springs
- Stumm W, Morgan JJ (1996) Aquatic chemistry: chemical equilibria and rates in natural waters. Wiley, Hoboke
- Sui MS, Xing L, Sheng SH, Guo H (2012) Heterogeneous catalytic ozon ation of ciprofloxacin in water with carbon nanotube supported manganese oxides as catalyst. J Hazard Mater 227:227 236
- Tay KS, Madehi N (2015) Ozonation of ofloxacin in water: by products, degradation pathway and ecotoxicity assessment. Sci Total Environ 520:23 31
- Valdés H, Zaror CA (2006) Heterogeneous and homogeneous catalytic ozonation of benzothiazole promoted by activated carbon: kinetic approach. Chemosphere 65: 1131 1136
- Valdés H, Tardón RF, Zaror CA (2009) Methylene blue removal from contaminated waters using O3, natural zeolite, and O3/zeolite. Water Sci. Technol. 60: 1419 1424

- Valdés H, Tardón RF, Zaror CA (2010) Effect of zeolite chemical surface properties on catalytic ozonation of methylene blue contaminated waters. Ozone Sci Eng 32:344 348
- Valdés H, Tardón RF, Zaror CA (2012a) Methylene blue removal from contaminated waters using heterogeneous catalytic ozonation pro moted by natural zeolite: mechanism and kinetic approach. Environ Technol 33:1895–1903
- Valdés H, Tardón RF, Zaror CA (2012b) Role of surface hydroxyl groups of acid treated natural zeolite on the heterogeneous catalytic ozona tion of methylene blue contaminated waters. Chem Eng J 211:388 395
- Velichkova FA (2014) Vers un procédé Fenton hétérogène pour le traitement en continu d'eau polluée par des polluants pharmaceutiques, École Doctorale Mécanique, Énergétique, Génie civil et Procédés (Toulouse)
- Velichkova FA, Delmas H, Julcour C, Koumanova B (2017) Heterogeneous Fenton and photo Fenton oxidation for paracetamol removal using iron containing ZSM 5 zeolite as catalyst. AICHE J 63:669 679
- Vittenet J, Rodriguez J, Petit E, Cot D, Mendret J, Galarneau A, Brosillon S (2014) Removal of 2, 4 dimethylphenol pollutant in water by ozonation catalyzed by SOD, LTA, FAU X zeolites particles obtain ed by pseudomorphic transformation (binderless). Microporous Mesoporous Mater 189:200 209
- Vittenet J, Aboussaoud W, Mendret J, Pic JS, Debellefontaine H, Lesage N, Faucher K, Manero MH, Thibault Starzyk F, Leclerc H, Galarneau A, Brosillon S (2015) Catalytic ozonation with  $\gamma$  Al<sub>2</sub>O<sub>3</sub> to enhance the degradation of refractory organics in water. Appl Catal A 504:519 532
- Von Sonntag C, von Gunten U (2012) Chemistry of ozone in water and wastewater treatment. IWA Publishing, London
- Yang Z, Xu X, Dai M, Wang L, Shi X, Guo R (2017) Accelerated cipro floxacin biodegradation in the presence of magnetite nanoparticles. Chemosphere 188:168 173