

# Correlation between degradation pathway and toxicity of acetaminophen and its by-products by using the electro-Fenton process in aqueous media

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| 1  | Degradation pathway and toxicity of acetaminophen by electro-Fenton processes in  |
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# 14 ABSTRACT

The evolution of the degradation by-products of an acetaminophen (ACE) solution was monitored by HPLC-UV/MS and IC in parallel with its ecotoxicity (*Vibrio fischeri* 81.9%, Microtox® screening tests) during electro-Fenton (EF) oxidation performed on carbon felt. The aromatic compounds 2-hydroxy-4-(N-acetyl) aminophenol, 1,4-benzoquinone, benzaldehyde and benzoic acid were identified as toxic sub-products during the first stage of the electrochemical treatment, whereas aliphatic short-chain carboxylic acids (oxalic, maleic, oxamic, formic, acetic and fumaric acids) and inorganic ions (ammonium and nitrate) were well identified as non-toxic terminal sub-products. Electrogenerated hydroxyl radicals then converted the eco-toxic and biorefractory property of initial ACE molecule (500 mL, 1 mM) and subsequent aromatic subproducts into non-toxic compounds after 2 h of EF treatment. The toxicity of every intermediate produced during the mineralization of ACE was quantified, and a relationship was established between the degradation pathway of ACE and the global toxicity evolution of the solution. After 8 h of treatment, a total organic carbon removal of 86.9% could be reached for 0.1 mM ACE at applied current of 500 mA with 0.2 mM of  $Fe^{2+}$  used as catalyst.

Key words: Carbon felt, Advanced Oxidation Process, TOC, Mineralization, By-products, Toxicity

#### 1 **1. Introduction**

Over the past few years, pharmaceutical drugs have become a severe problem for the domestic 2 wastewater treatment (Schwarzenbach et al., 2006). Pharmaceuticals such as anti-inflammatories, 3 analgesics, lipid regulators, antibiotics, anti-epileptics, betablockers and oestrogens are used 4 extensively for both humans and livestock. When consumed, 10-90% of these drugs are 5 metabolized. A significant quantity of the active ingredients is therefore excreted, mainly through 6 7 urine and faeces, and collected in urban sewer systems or discharged directly into the environment by livestock. Many studies have confirmed the presence of a large number of medicinal substances 8 9 in all environmental compartments: ground and surface water (Andreozzi et al., 2003; Leónidas A. Pérez-Estrada et al., 2005), drinking water (Ternes et al., 2002), tap water (Doll and Frimmel, 10 2003), ocean water, sediment and soil (Halling-Sørensen et al., 1998); this indicates the 11 12 inefficiency of conventional water treatment processes to remove these kinds of micro-pollutants. Even if the amount of these pharmaceuticals in aquatic media is low, usually with concentrations 13 lower than 10 µg L<sup>-1</sup> (Ternes, 1998; Huerta-Fontela et al., 2011; Huerta et al., 2016), its 14 continuous input constitutes at long term a real risk for aquatic and terrestrial organisms (Zhou et 15 16 al., 2013; García-Mateos et al., 2015). Some pharmaceuticals are indeed suspected to affect the endocrine system of fishes, and available data on antibiotics indicate that they can exert toxic 17 effects on algae and invertebrates and may favour the development of multi-drug resistant 18 microbial strains (Zuccato et al., 2000; Balcıoğlu and Ötker, 2003). Among pharmaceuticals, 19

acetaminophen (ACE) is an analgesic molecule consumed worldwide, and is usually selected by researchers as an emerging contaminant model (García-Mateos et al., 2015). ACE is also called paracetamol, and its chemical name is N-acetyl-p-aminophenol. This drug has been detected in European wastewater treatment plant (WWTP) with a concentration of 6  $\mu$ g L<sup>-1</sup> (Ternes, 1998), up to 10  $\mu$ g L<sup>-1</sup> in natural water resource in the USA (Kolpin et al., 2002) and more than 65  $\mu$ g L<sup>-1</sup> in the Tyne River, UK (Kabdaşli et al., 1999; Roberts and Thomas, 2006).

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Because of the toxicity and persistency of pharmaceuticals, it is necessary to develop 27 powerful methods to ensure complete degradation of pharmaceutical pollutants as well as their 28 potentially toxic degradation by-products. Conventional technologies that are currently used, such 29 as activated carbon adsorption, flocculation, biological degradation, and chemical process like 30 chlorination, have the disadvantages of being poorly efficient for non-biodegradable compounds 31 or to transfer the pollutants into the solid phase (sludge or sorbent). Pollution is, in this case, only 32 displaced and not solved. Advanced oxidation processes (AOPs) (Ozonation, UV/H<sub>2</sub>O<sub>2</sub>, electro-33 and photo-Fenton) can be easily incorporated in pre-existing treatment plants and were found to 34 achieve efficient reduction of micropollutant discharge in the aquatic environment (Reungoat et 35 al., 2010; Reungoat et al., 2012; Eggen et al., 2014; Sirés et al., 2014; Moreira et al., 2017). 36 Because non-selective radicals such as hydroxyl radicals (OH) are generated, AOPs are considered 37 a competitive water treatment technology for the degradation of these refractory organic 38 micropollutants and the improvement of effluent quality from treatment plants (Gerrity et al., 39 2011; Giannakis et al., 2015). Several publications have highlighted the potentialities of AOPs 40 such as UV/H<sub>2</sub>O<sub>2</sub> (Frontistis et al., 2012; Lee et al., 2016); ozonation (Huber et al., 2005; Gerrity 41 et al., 2011; Zimmermann et al., 2011; Reungoat et al., 2012; Margot et al., 2013), Fenton and 42 photo-Fenton (Tekin et al., 2006; Kulik et al., 2008), semiconductor photocatalysis (Molinari et 43 al., 2006; Abellán et al., 2007), and electrochemical separation and degradation technologies 44 (Sirés and Brillas, 2012) to eliminate biorefractory pharmaceuticals. Recently, textural 45

modification of carbon felt (Le et al., 2016) and graphene deposition on graphite substrates (Le et 46 al., 2015a&b) have shown a great potential to enhance mineralization of the azo dye acid Orange 7 47 and ACE by the electro-Fenton (EF) process. Derived Fenton processes have also already proved 48 to be efficient methods to degrade ACE from water: combined EF and photo-Fenton using a 49 double cathode electrochemical cell (de Luna et al., 2012), coupling adsorption and photo-Fenton 50 using nano-zeolites and cobalt ferrite nanoparticles (Irani et al., 2015), bio-electrochemical 51 degradation in a microbial fuel cell-Fenton system (Zhang et al., 2015), or other methods relevant 52 to the photo-Fenton process (Trovó et al., 2008; Trovo et al., 2012; Cabrera Reina et al., 2015; 53 Rad et al., 2015). As a more detailed example, Sirés et al. could totally mineralize ACE in CO<sub>2</sub> 54 after 6 h of treatment by using the catalytic action of  $Fe^{2+}$ ,  $Cu^{2+}$  and UVA light to generate 55 hydroxyl radicals. During the electrolysis, several by-products were detected, such as 56 hydroquinone, p-benzoquinone, or short chain carboxylic acids (Sirés et al., 2006). Using the solar 57 58 photoelectro-Fenton (SPEF) method, ACE was efficiently mineralized in the study of Lucio Cesar Almeida et al. (Almeida et al., 2011). In particular, after 120 min of electrolysis, total organic 59 carbon (TOC) removal attained was 75% with an energy cost of 93 kWhkg<sup>-1</sup> TOC (7.0 kWhm<sup>-3</sup>) 60 and a mineralization current efficiency of 71%. In addition, many by-products including aromatic 61 compounds as well as carboxylic acids were adequately detected, such as hydroquinone, p-62 benzoquinone, 1,2,4-trihydroxybenzene, 2,5-dihydroxy-p-benzoquinone and tetrahydroxy-p-63 benzoquinone, maleic, fumaric, succinic, lactic, oxalic, formic and oxamic acids. In addition, the 64 combination of adsorption and photo-Fenton method by Irani et al. could lead to 99.80% of 65 paracetamol removal in 30 min (Irani et al., 2015). 66

Depending on the AOP technique used, several degradation pathways of ACE have been identified and already published (Andreozzi et al., 2003; Brillas et al., 2005; Skoumal et al., 2006; Yang et al., 2008a). All authors observed the formation of reactive by-products due to partial oxidation of the initial molecule and reaction with matrix components (Von Gunten, 2003). It is important to track the intermediates generated through degradation reactions and to know their

eco-toxicity, with the view to verify the safety of the water treatment process. Unfortunately, very few works were aimed at combining analytical chemistry to individual and global toxicity measurement of both products and solution during AOP treatment (Hamdi El Najjar et al., 2014) as presented in this study.

Research at laboratory and pilot scale clearly demonstrated that AOPs can be applied to 76 improve the quality of effluents from municipal and industrial WWTPs by attenuating trace 77 organic contaminants (micropollutants) (Gerrity et al., 2011; Giannakis et al., 2015). However, a 78 potential disadvantage of these processes is the formation of unknown reactive by-products due to 79 partial oxidation of the targeted compounds and reaction with matrix components (von Gunten, 80 2003). Thus, the final toxicity of treated effluent should be determined together with the quantity 81 of targeted compounds in terms of the detection limits of the analytical protocols. In this study, the 82 EF reaction was applied to ACE solution. ACE and its by-products generated during the 83 84 degradation were monitored by HPLC-UV/MS, IC and TOC analysis. In addition, acute toxicity of the ACE solution under EF treatment as well as individual and mixed intermediate standards 85 86 were assessed through Microtox® toxicity test with Vibrio Fischeri marine bacteria. Through this study, the relationship between the ACE degradation pathway, the individual by-product toxicity 87 and the global toxicity of the solution could be newly and clearly established, thus providing a 88 more comprehensive understanding and safety control of the mineralization process of ACE by the 89 EF process. 90

91

#### 92 **2. Experimental**

93 2.1. Materials

ACE (acetaminophen: paracetamol), sodium sulphate (anhydrous, 99.0–100.5%), sodium hydroxide (99%), sulphuric acid (95–97%), iron (II) sulphate hepta-hydrate (99%), benzoquinone, benzaldehyde, benzoic acid, oxalic acid, maleic acid, oxamic acid, formic acid, fumaric acid and acetic acid were obtained from Sigma-Aldrich, USA. Bacterial strain of *Vibrio fischeri* NRRL B- 98 11177 involved in toxicity tests was obtained from Hach Lange GmbH, Germany. Osmotic 99 adjusting solution (MilliQ water with 22% NaCl) and diluent (MilliQ water with 2% NaCl) were 100 used for the preparation of the bacterial solution. The carbon felt was purchased from A Johnson 101 Matthey Co., Germany. TOC standard of 1000 mg L<sup>-1</sup> (Sigma-Aldrich, USA) and sodium 102 hydrogen carbonate ( $\geq$ . 99.5%, ACS, Karlsruhe, Germany) were used for TOC and inorganic 103 carbon (IC) calibration curves assessments, respectively.

104

# 105 2.2. Electrochemical system for ACE degradation

EF experiments were performed at room temperature in a 500-mL single cylindrical glass 106 cell with a two-electrode system. The applied current was controlled using a power supply 107 (Lambda Electronic, USA) in a galvanostatic mode. The carbon felt cathode ( $60 \text{ cm}^2$ ) was used as 108 a working electrode and a platinum cylindrical mesh as a counter electrode. The anode was 109 centred in the electrolytic cell, surrounded by the cathode, which covered the inner wall of the cell. 110 The distance between the electrodes was 1.5 cm. The aqueous solution of ACE, Na<sub>2</sub>SO<sub>4</sub> (50 mM) 111 as the supporting electrolyte and FeSO<sub>4</sub>.7H<sub>2</sub>O as the catalyst were prepared by adjusting the pH at 112 3.0 with sulphuric acid (H<sub>2</sub>SO<sub>4</sub>). The intermediate compound monitoring experiment was 113 performed in 500 mL solution of ACE 1 mM and/or 200 mL solution of ACE 0.1 mM with 114 FeSO<sub>4</sub>.7H<sub>2</sub>O (0.2 mM) at an applied current of 500 mA. Before the electrolysis, oxygen bubbling 115 was performed for 5 min to saturate the solutions, and the solutions were kept under continuous 116 magnetic stirring at a rate of 800 rpm during the experiment. 117

118 The specific energy consumption per unit TOC mass ( $EC_{TOC}$ ) was calculated by Eq. (1) (Ruiz et 119 al., 2011):

120  $EC_{TOC} (kWh g^{-1} TOC) = \frac{VIt}{(\Delta TOC)_{exp} V_s} (1)$ 

where V is the average cell voltage (V), V<sub>s</sub> is the solution volume (L),  $\Delta$ (TOC)<sub>exp</sub> is the experimental TOC decay (mg L<sup>-1</sup>), I is the applied current (A) and t is the electrolysis time (h).

Concentrations of ACE and their aromatic oxidation by-products were analysed by a HPLC 125 system (Waters 717 Autosampler and Waters 616 Pump) with a Photodiode Array Detector 126 (Waters 2996 Photodiode Array Detector) fitted with a reverse phase Thermo Scientific C18 127 column (L=250 mm, I.D=4.6 mm, and 5-µm particle size). The column temperature was 128 thermostated at 30°C. The injection volumes were 20 µL. The mobile phase was constituted of 129 buffer A (HPLC grade water + 0.1 % (v/v) trifluoroacetic acid) and buffer B (HPLC grade 130 acetonitrile + 0.1 % (v/v) trifluoroacetic acid). The flow rate was 1 mL.min<sup>-1</sup>. The eluent gradient 131 started with 10% of eluent B, gradually increasing to 70% in 35 min. Eluent B was elevated to 132 90% in 5 min, and the system was kept for 5 min, returning to 10% in 1 min and equilibrating for 133 9 min (returning to initial condition and re-equilibrating the column). UV detection was performed 134 at  $\lambda$ =254 nm for substituted benzene structure monitoring. 135

The degradation products of ACE were identified by an LC-MS system. LC-MS was 136 performed with an Alliance e2695 (Waters HPLC pump and autosampler system), and a Quattro-137 138 Micro mass spectrometer was equipped with an Electrospray probe (Waters Micromass, Wythenshawe, Manchester, UK). The detection conditions were as follows: capillary potential of 139 3.5 kV, cone voltage of 30 V, source temperature of 120°C, desolvation temperature of 450°C, 140 cone gas flow of 50 L h<sup>-1</sup> and desolvation gas flow of 450 L.h<sup>-1</sup>. Nitrogen was used as the 141 nebulizer gas. A Waters column HSST3 (L=100 mm, D.I= 2.1 mm, and 5-µm particle size) was 142 used to separate intermediates. The column was set at room temperature (22°C). The injection 143 volumes were 20  $\mu$ L. The mobile phase comprised buffer A (HPLC grade water + 0.1% (v/v) 144 formic acid) and buffer B (HPLC grade acetonitrile + 0.1% (v/v) formic acid). The flow rate was 145 146 0.25 mL min<sup>-1</sup>. The eluent gradient started with 10% of eluent B and was kept for 1 min, gradually increasing to 90% in 2 min, and the system was then kept for 1 min, returning to 10% in 1 min and 147 148 equilibrating for 2 min (returning to initial condition and re-equilibrating the column). The eluent from the chromatographic column successively entered the UV-Vis diode array detector (Waters 149

PDA996) and the electrospray ionization (ESI) interface of the mass spectrometer. Mass
spectrometry (MS) analysis was performed in a positive mode. The mass range was 90–400 (m/z).

The eluted components having specific retention time were characterized by their UVspectra and/or their mass. The quantification using the standard addition method (HPLC-UV) or external calibration against standard (LC-MS) was performed. Short-chain acids were identified and quantified using the HPLC-UV system already described, but fitted with a SHODEX KC 811 (D.I=8 mm, L=300 mm and 7-µm particles) column, which was thermostated at 30°C. UV detection was performed at  $\lambda$ =210 nm. The mobile phase was buffer A (HPLC grade water + 0.1% (v/v) Phosphoric acid 85%). The flow rate was 0.50 mL min<sup>-1</sup>.

Inorganic ions (NO<sub>3</sub><sup>-</sup>, NH<sub>4</sub><sup>+</sup>) generated during decomposition of ACE were identified by an ion chromatography equipment. For anion, Dionex ICS-1000 system was connected to an AS19 column (4 mm  $\times$  250 mm), and the eluent KOH was applied according to the elution gradient: 10 mM in 10 min, then gradient in 20 min to 45 mM. Dionex ICS-900 was used to measure the cation concentration in the treated solution with the column CS12A and 20-mM methane sulphonic acid as eluent. Both columns were thermally controlled at 30°C, and DS6 conductivity detector was used.

The TOC and the IC of the initial and treated samples were determined using a TOC-L CSH/CSN Shimadzu (Japan) analyser. Calibration curves for TOC and IC analysis were constructed by automatic dilution on the same machine from solutions of TOC standard of 1000 mg/L and sodium hydrogen carbonate.

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173 *2.4.Toxicity tests* 

174 The Microtox® bacterial toxicity test is based on the general principles described by ISO
175 (2007) (Romero et al., 2008). The toxicity of ACE and its intermediate by-products formed during

its degradation was determined by measuring the effect on the luminescence of marine bacteria.
Bacterial luminescence was measured using a Microtox® Model 500 Analyzer (Modern Water
Inc., UK). The bacterial strain used in this method was *Vibrio fischeri* NRRL B-11177. During
growth, this bacterium emitted luminescence that was related to its cellular respiration and cell
activity. As the activity of the bacterial cell could be reduced by the presence of toxic elements,
bioluminescence is therefore a very good indicator of state of the bacterium and thus of the global
toxicity of the sample.

This device allowed for performing the acute toxicity tests by using the software 183 MicrotoxOmni<sup>®</sup>. To identify the relative toxicity of each samples solution, an 81.9% screening 184 test was used to characterize the inter-sample toxicity variability: the influence of treatment 185 processes on the effluent toxicity was monitored with a sampling method performed as a function 186 of time. The test performed in this study is called 81.9% screening test because all samples were 187 188 diluted at 81.9% of the initial sample concentration by adding a 22% NaCl solution to allow V. fischeri normal activity and thus luminescence emission. Before measuring the bacterial 189 190 luminescence, pH of the samples were adjusted between 6.5 and 7.5 by sodium hydroxide or sulphuric acid, and filtration was then performed with a 0.2-µm syringe filter to eliminate any 191 precipitate or solid matter in the solution. 192

In the absence of toxicity, the luminescence of bacteria decreases over time under the effect of environmental conditions. Thus, it is necessary to compensate the errors due to these factors by considering the variability of the luminescence R(t) of the bacteria in a control solution (MilliQ water and NaCl), which gives the LU0 values. Equation (2) gives the corrected term R(t).

197 
$$R(t) = \frac{LU0(t)}{LU0(0)}$$
 (2)

where LU0(t) is the intensity of luminescence emitted by bacteria after a t=5 min or
t=15min of contact with the control solution (MilliQ water and NaCl) (Arbitrary Units (A.U.));
and LU0(0) is the initial intensity of luminescence emitted by bacteria before the addition of the
control solution (MilliQ water and NaCl) (A.U.).

The corrected inhibition rate at time t:I(t) intrinsically attributable to the sample toxicity is then calculated using Eq. (3). To simplify, I(t) is called "Inhibition" hereafter (Figs. 5, 6 and 7) and is expressed as percentage.:

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I(t) (%) = 
$$\left(1 - \frac{LU(t)}{R(t) \times LU(0)}\right) \times 100$$
 (3)

206

where LU(t) is the intensity of luminescence emitted by bacteria after a t=5 min or t=15min of contact with the sample (A.U.) and LU(0) is the initial intensity of luminescence emitted by bacteria before the addition of the sample (A.U.).

Furthermore, eco-toxicity tests were performed on the commercially available by-products to confirm the toxicity of the compounds formed during the degradation of the main molecule, ACE. These by-products and their mixtures were prepared with the maximum concentrations detected by HPLC analysis during the EF experiment.

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#### 215 **3. Results and discussion**

### 216 *3.1. Degradation pathway of acetaminophen during the EF process*

#### 217 *3.1.1.* Evolution of aromatic compounds

The degradation of ACE by the EF process formed some aromatic compounds such as 1,4-218 benzoquinone, benzaldehyde and benzoic acid identified by HPLC (Fig. 1) and 2-hydroxy-4-(N-219 220 acetyl) aminophenol identified by r LC/MS (Table S1). It was noted that at the beginning of the ACE degradation, 1,4-benzoquinone was the main aromatic by-product identified. Its highest 221 concentration reached 0.23 mM after 10 min of the degradation process; its concentration then 222 223 decreased, and this product disappeared totally after 120 min. The other identified aromatic intermediates are also detected at a very short time of decomposition process with a concentration 224 peak of 0.09 and 0.039 mM at 14 and 10 min for benzoic acid and benzaldehyde, respectively. 225 226 These aromatic compounds were totally decomposed after 120 min of electrolysis and were

227 converted into other aliphatic carboxylic acids and inorganic ions with the appropriate mass228 balance.

#### 229



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Fig. 1. Evolution of the concentration of ACE and the aromatic degradation by-products (RSD 2% n=3) during EF treatment of 500 mL ACE solution at  $C_0 = 1.0$  mM; pH =3;  $[Na_2SO_4]=50$  mM, I = 500 mA and  $[Fe^{2+}] = 0.2$  mM.

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### 235 3.1.2. Evolution of short-chain carboxylic acids

The evolution of carboxylic acid concentration during the electrolysis treatment of a 1.0-236 mM ACE aqueous solution at pH 3.0 is reported in Fig. 2. These by-products are usually found in 237 ACE degradation by AOPs such as ozonation (Skoumal et al., 2006) or electrogenerated hydrogen 238 peroxide process (Reungoat et al., 2010). Oxalic, maleic, oxamic, formic, acetic and fumaric acids 239 were detected at 11.6, 13.59, 15.23, 20.59, 22.25 and 22.89 min of retention time, respectively, 240 241 during the degradation experiment. Figure 2 shows that fumaric and oxamic acids were generated after around 20 min of electrolysis, while the other by-products such as maleic, acetic, formic and 242 oxalic acids were detected only after 3 h. The accumulation of oxalic acid is well known in 243 Fenton-related processes because of the formation of iron-oxalate complexes that are slowly 244 removed by 'OH. At the end of the EF treatment (8 h), fumaric, acetic and formic acids were 245 completely removed from solution, while oxalic and maleic acid were still present in the solution. 246

Singularly, oxamic acid concentration remained high (around 0.52 mM) after 8 h of treatment,
with a TOC removal of 51%. This indicated that oxamic acid remained stable under our
experimental conditions.

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Fig. 2. Evolution of the concentration of carboxylic acids (RSD 2% n=3) during EF degradation of ACE: Oxalic acid ( $\bullet$ ), Acetic acid ( $\Box$ ), Formic acid ( $\Delta$ ), Oxamic acid ( $\bullet$ ), Fumaric acid (\*), Maleic x 10 ( $\bigcirc$ ). Conditions: 500 mL ACE solution at C<sub>0</sub> = 1.0 mM; pH =3; [Na<sub>2</sub>SO<sub>4</sub>]=50 mM, I = 500 mA and [Fe<sup>2+</sup>] = 0.2 mM.

256

#### 257 3.1.3. Evolution of inorganic ions

The mineralization of ACE and their intermediates also produced inorganic ions. 258 Ammonium and nitrate concentrations were monitored during the 8 h of treatment (Fig. 3). Nitrate 259 was measured at very low concentrations compared to ammonium. As observed by E. Brillas et al. 260 (Brillas et al., 2009), the first limiting step of ACE degradation is the hydroxylation reaction 261 leading to 2-hydroxy-4-(N-acetyl) aminophenol (Table 1), which is then converted into oxamic 262 acid. The presence of  $Fe^{2+}$  in the solution as catalyst could lead to the concomitant production of 263 persistent iron-oxamate complexes, which are very difficult to mineralize by •OH, thus limiting 264 the oxidation ability of the EF process. Therefore, nitrate was clearly detected and quantified only 265 after 2 h of treatment and reached 0.02 mM in 8 h, while ammonium concentration reached up to 266 0.38 mM at the same time (Fig. 3). At the end of the treatment process, the initial nitrogen-267

containing compound (1 mM) was converted into oxamic acid (0.52 mM), ammonium (0.38 mM)
and trace of nitrate (0.02 mM) (Figs. 2 and 3). Therefore, a total of 0.92 mM of N in by-products
was quantified, which means that the nitrogen mass balance is confirmed, considering that around
8% of nitrogen lost during the EF process could be explained by the fact that nitrogen could be
transferred to the volatile phase through N<sub>2</sub> and NH<sub>3</sub> apparition (Hammami et al., 2008).

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Fig. 3. Evolution of the inorganic ions concentration (RSD 5% n=3) during EF degradation of ACE. Conditions: 500 mL ACE solution at  $C_0 = 1.0$  mM; pH =3; [Na<sub>2</sub>SO<sub>4</sub>]=50 mM, I = 500 mA and [Fe<sup>2+</sup>] = 0.2 mM.

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# 279 3.1.4. Degradation pathway of ACE mineralization during the EF process

After 8 h of treatment, almost 90% of organic compounds in the initial ACE solution (200 mL at 0.1mM) were mineralized, proving the efficiency of the EF process for removing the pharmaceutical pollutants (Fig. S1). The specific energy consumption in 120 min was calculated at 3.6 kWh g<sup>-1</sup> TOC, which is a low value, proving the interest of the system, compared to the reported value of 93 kWh g<sup>-1</sup> TOC obtained by the solar SPEF method (Almeida et al., 2011).

A general reaction sequence for the EF degradation of ACE in acidic medium is proposed in Fig. 4. The process was initiated by •OH radical attack either on the C(2) position of ACE (I), which led to the formation of 2-hydroxy-4-(N-acetyl)aminophenol (II), or on its C(4)-position

which led to 1,4-benzoquinone (III). The formation of these intermediate products could also be 288 proposed in other AOP processes during the degradation of ACE (Yang et al., 2008b). 2-Hydroxy-289 4-(N-acetyl) aminophenol (II) was then degraded into benzoic acid (V), benzaldehyde (IV) and 290 acetamide. Subsequently, all aromatic by-products were further oxidized to produce aliphatic 291 organic acids such as fumaric, oxalic, acetic, formic and maleic acids. Furthermore, oxamic acid 292 was generated from the oxidation of acetamide (Skoumal et al., 2006). At the end of the 293 294 degradation process, unlike oxamic and oxalic acids which persisted under our experimental conditions, all these last by-products were transformed into CO<sub>2</sub>, H<sub>2</sub>O and inorganic ions. 295



Fig. 4. General reaction sequence proposed for acetaminophen (ACE) mineralization in an acidicmedium by hydroxyl radicals during the EF process.

*3.2. Toxicity evolution of ACE solution during the EF process* 

*3.2.1. Global toxicity* 

The global toxicity evolution of 500 mL of 1.0-mM ACE solution during 8 h of EF 303 treatment is shown in Fig. 5. The acute toxicity of the treated solution after 5- and 15-min time 304 contact with V. fischeri bacterial strain increased rapidly at the early stage of the treatment and 305 reached 100% between 2 and 55 min. This result is relevant and consistent with regard to the 306 previously proved formation of toxic aromatic by-products such as 1,4-benzoquinone, benzoic 307 acid and benzaldehyde (Lee and Chen, 2009; Das et al., 2010; Dirany et al., 2012). After 60 min, 308 the toxicity markedly declined which well agreed with aromatic compounds found only at trace 309 levels at this stage of the EF reaction (see part 2.1.1 and Fig. 1). The predominance of very low 310 toxic short-chain carboxylic acids in this period did not significantly contribute to the inhibition of 311 the bacterial luminescence; consequently, the solution toxicity became zero at about 120 min. 312 Subsequently, the inhibition value was continuously maintained at around zero until 8 h where 313 short-chain carboxylic acids were continuously transformed under •OH attack during the EF 314 315 process. The two exposure times measured yielded identical inhibition (%) results.

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**Fig. 5.** Inhibition of luminescence of *V. fischeri* marine bacteria during EF treatment of 500 mL of 1 mM ACE at pH=3.0;  $[Na_2SO_4]=50$ mM, I=500 mA,  $[Fe^{2+}]=0.2$ mM. Two exposure time of the bacterial strain to the treated solution were used (5 and 15 min). RSD (n = 5): 12% for inhibition < 20%; 5% for 20% < inhibition < 70% and 0% for inhibition > 70%.

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# 323 *3.2.2. Toxicity of intermediate sub-products*

The toxicity of individual and mixed aromatic by-products taken as standard solutions was investigated by the ecotoxicity assays. Their concentration was selected with regard to those analysed during the ACE degradation. For this purpose and according to by-products formation shown in Fig. 2, specific times of EF treatment were considered: 10, 60, 80 and 120 min corresponding to different mixture solutions and concentrations of by-products (Table S2).

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As shown in Fig. 6a, b, c and d, ACE itself is not a hazardous pollutant for this strain of 330 bacteria because its toxicity was very low even at the high concentration (0.5 mM) found after 331 10 min of treatment. Similarly, benzoic acid was also a low toxic by-product because its 332 negative toxicity values towards the bacterial strain during the degradation process signifies a 333 positive effect on bacteria growth and activity (Velegraki et al., 2010). In contrast, 1,4-334 benzoquinone showed an important toxic effect as the inhibition ratio was around 100% even at 335 very low amount, 0.04 mM (60-min degradation), compared to other formed aromatic 336 intermediates. Although benzaldehyde was not as toxic as 1,4-benzoquinone, this compound 337 338 still largely affected bacterial luminescence (Paulraj et al., 2011). It caused an acute toxicity between about 50% and 20% at the concentration of 0.039 mM (10 min) and 0.002 mM (120 339 min), respectively. In addition, the mixtures of these aromatic compounds at four periods from 340 341 10, 60, 80-120 min were ecotoxically assessed (Fig. 6e). The inhibition ratio reached 100% at the early stage of the treatment (10 and 60 min), again proving the high toxicity of 1,4-342 benzoquinone. This agrees with previously published results (Rubio et al., 2011). Subsequently, 343 a remarkable detoxification was observed after 80 and 120 min due to the decline of the 344 corresponding toxic aromatics. 345

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Fig. 6. Toxicity of standard solutions of ACE and other aromatic compounds, taken alone (a, b, c,
d) or in mixture (e), under identical concentration, during the EF process on 500 mL (1-mM ACE)
and for different treatment times: (a) 10 min, (b) 60 min, (c) 80 min, (d) 120 min. (e) Mixture
solutions of standard ACE and aromatic compounds at different treatment times.

355 At the end of electrolysis, the oxidation by •OH generated during the EF process in solution led to the production of ring-opened products such as aliphatic carboxylic acids. These 356 acids such as formic acid or acetic acid are known to be low or non-toxic compounds (Liu et al., 357 2010). This suggestion was confirmed by their toxicity measurement presented in Fig. 7 in two 358 ways: individual test on each single standard compound in MilliQ water at its maximum 359 concentration measured during the EF process (Fig. 7a) and the combination (mixture) of all 360 carboxylic by-products at their maximal concentrations measured during three periods of the 361 electrolysis (Fig. 7b): 210–270 min (Mix 1); 270–330 min (Mix 2) and 330–480 min (Mix 3) 362 (Table S3). Acetic and oxamic acids toxicity was measured for similar maximum concentration of 363 about 0.5 mM, but oxamic acid seemed to promote higher toxicity than acetic acid with individual 364 toxicity values of 25.7% and 7%, respectively. The toxicity of maleic, oxalic and fumaric acids 365 366 was around 15%. In comparison with other intermediates, formic acid contributed negligibly to the 367 toxicity of the solution. Interestingly, a cocktail effect was observed: synergistic (Mix 1) as well as antagonist effects (Mix 3) were observed during the comparative study on single and mixture of 368 369 standard compounds. As observed in other studies with pesticides or antioxidants, the combinations of ACE carboxylic acid by-products did not induce an additive increase in 370 individual toxicity (Liu et al., 2010). The toxicity of three mixtures decreased gradually as a 371 function of treatment time, 18.7%; 18.1% and 0%, respectively, from 210 to 480 min. These 372 results confirmed that the EF process is an environmentally friendly technology, which can 373 374 efficiently degrade toxic pollutants into non-toxic compounds.



**Fig. 7.** Toxicity of (a) short-chain carboxylic acid by-products at their maximum concentrations (Table 3), and (b) Mixture of the short-chain carboxylic acid by-products corresponding to three periods: 210–270 min (Mix 1); 270–330 min (Mix 2) and 330–480 min (Mix 3) during EF treatment of 500 mL (1-mM ACE) at pH = 3;  $[Na_2SO_4] = 50$  mM, I = 500 mA,  $[Fe^{2+}] = 0.2$ mM.

380

#### 381 **4.** Conclusions

The evolution of TOC, intermediate compound concentrations and acute toxicity of ACE 382 solution in aqueous medium during the EF mineralization process was monitored through HPLC-383 UV/MS, IC, TOC analysis and Microtox® toxicity tests. During the first 60 min of treatment, the 384 acute toxicity of the treated solution reached 100%, which was identified on the basis of the 385 formation of toxic aromatic products such as 1,4-benzoquinone, benzoic acid and benzaldehyde. A 386 total detoxification of the solution was then accomplished after 120 min due to the decomposition 387 388 of ring-compounds into short-chain carboxylic acids. The mineralization of ACE by hydroxyl radicals (•OH) formed through the EF process also produced inorganic ions such as ammonium or 389 390 nitrate by bond breakage between nitrogen and carbon in the initial molecule. The mineralization of 200 mL of 0.1-mM ACE was 86.9% using an applied current of 500 mA and a catalyst 391 concentration of 0.2 mM. The relationship between the decomposition pathway and the evolution 392 of the ACE solution toxicity as a function of EF treatment time was clarified by the toxicity 393 measurement of both individual and mixture of standard compounds (ACE and sub-products). 394 This comprehensive degradation pathway of ACE can be further applied to other persistent 395 pharmaceuticals in an aquatic environment. 396

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