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Effect of electrochemically-driven technologies on the treatment of endocrine disruptors in synthetic and real urban wastewater

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Abstract

While it remains a challenging task, having a fundamental understanding of the influence of water matrix composition is of central importance for the removal of pollutants in electrochemical treatment processes. This work studied the effect of the complexity of different water matrices on the removal of endocrine disruptors using different electrochemical advanced oxidation processes (EAOPs) based on H₂O₂ electrogeneration. The study helped improve the understanding of synthetic and real urban wastewater, and the effective mechanisms for the treatment of pollutants in these effluents. The results obtained showed that, regardless of the medium applied, there was an increase in pollutants mineralization in the following order, based on the EAOPs method employed: electrochemical oxidation with H_2O_2 electrogeneration (EO- H_2O_2) < electro-Fenton (EF) < photoelectro-Fenton (PEF). Despite the slower degradation kinetics of the EO-H₂O₂ process, wider it suitable implementation. its pН range makes for Aside the degradation/mineralization performance of the EO-H₂O₂ process, a time-course analysis was conducted on the nitrogen and chloride-based ions in the solution investigated since their presence is vital for determining the final use of the treated effluents based on the parameters of water quality. These parameters include the established maximum concentration levels of the compounds in the water matrix allowed for human consumption or the legal limits of discharge for wastewater treatment facilities. The findings help shed light on the role of coexisting species in water matrix, since this affects the efficiency and competitiveness of EAOPs. Clearly, having detailed knowledge of the water matrix composition will help one to have a better understanding of EAOPs, and this will enable to obtain higher degradation levels in water treatment processes.

Keywords: Bisphenol A; methyl paraben; paracetamol; inorganic species; real wastewater; water matrix

1 Introduction

Currently, water scarcity and pollution are among the major challenges faced by humanity. This situation has been dramatically intensified by the increase in world population and the incessant quest for the improvement of life quality [1]. In fact, some of the so-called "new helpful daily life products", as is the case of plastic bottles and containers, linings of metal food cans, detergents, flame retardants, food, toys, drugs and cosmetics, and pesticides, have been shown to have adverse effects on both the environment and humans over time [2– 7]. The natural and man-made chemicals that are used in manufacturing these products end up being disposed of into the environment spreading across the ecosystem; these chemicals contain endocrine disruptors, which have been found to be extremely hazardous to humans and the ecosystem as a whole [8]. The endocrine disruptors may mimic or interfere with the body's hormones causing developmental, reproductive, brain, and immune-related problems. Many studies published in the literature have reported the presence of endocrine disruptors in wastewater disposed of into the environment and in water bodies [9,10]. Conventional water treatment processes have been demonstrated to be inefficient when it comes to the degradation of these recalcitrant organic pollutants [11,12]. Electrochemical advanced oxidation processes (EAOPs) have emerged as a sustainable alternative for the removal of endocrine disruptors where the electron is used as green reagent for the *in-situ* generation of oxidants [13]. The degradation of organic pollutants by EAOPs occurs via strong reactive oxygen species such as hydroxyl radical ([•]OH); this radical is a non-selective oxidant with a high standard reduction potential of $E^{\circ} = 2.80$ V vs SHE. In the case of the electrochemical oxidation (EO) process, [•]OH radical is obtained from water electrolysis occurring on the

surface of the anode (Eq. 1) [14–16]. The •OH electrogenerated during anodic oxidation remains adsorbed on the electrode surface and the degradation of organic compounds is limited by mass transfer.

$$M + H_2O \rightarrow M(^{\bullet}OH) + H^+ + e^-$$
(1)

One will note, however, that other indirect electrochemical processes can promote the formation of $^{\circ}$ OH in the bulk. This is the case of electrochemical processes based on Fenton reaction (2). The electro-Fenton (EF) process relies on the use of carbonaceous cathodes to promote the selective bi-electronic reduction of oxygen to hydrogen peroxide (E⁰=0.695 V vs SHE) via reaction (2) [17]. Electrogenerated H₂O₂ can then react with Fe²⁺ in the solution to yield $^{\circ}$ OH through reaction (3) [18]. The EF system not only allows the continuous dosing of H₂O₂ without the need of transport, storage, and handling but also enables the fast regeneration of the Fe²⁺ catalyst from the electrochemical reduction of Fe³⁺ through reaction (4) [19–21].

$$O_{2(g)} + 2H^+ + 2e^- \rightarrow H_2O_2 \tag{2}$$

$$H_2O_2 + Fe^{2+} \rightarrow Fe^{3+} + {}^{\bullet}OH + OH^-$$
(3)

$$\mathrm{Fe}^{3+} + \mathrm{e}^{-} \to \mathrm{Fe}^{2+} \tag{4}$$

The degradation performance of the EF process can be markedly enhanced when the treated solution is subjected to simultaneous irradiation with UV light through the photoelectro-Fenton (PEF) process. The light conveyed into the solution promotes photon-driven reactions, such as the regeneration of iron catalyst based on reaction (5), and mineralization of iron-carboxylate complexes through charge transfer reactions from the ligand to the metal based on the general reaction (6) [22,23].

$$Fe(OH)^{2+} + hv \rightarrow Fe^{2+} + {}^{\bullet}OH$$
(5)

$$Fe(OOCR) + hv \rightarrow Fe^{2+} + CO_2 + R^{\bullet}$$
(6)

Despite the promising results obtained from the degradation of organic persistent pollutants, there are a few questions that remain unanswered regarding the competitiveness of electrochemically-driven Fenton processes. Previous studies have shown the high competitiveness of electrochemical treatment processes during the treatment of ultrapure water containing only one target pollutant. However, the treatment of endocrine disruptors can be affected by the coexistence of different species in real water matrices. To the best of our knowledge, this problem has neither been thoroughly explored nor has it been widely established in the literature [24]. This work aims to investigate the effect of the complexity of the constituents of natural water matrices on the removal of endocrine disruptors. To this end, the present study sought to evaluate the efficiency of EAOPs in the treatment of synthetic and real urban wastewater contaminated with endocrine disruptors with the aim of investigating the effect of the water matrix. The study also quantified the accumulated amount of H₂O₂, pollutant degradation rates, total organic carbon (TOC) removal, and the amount of carboxylic acids generated. Engineering figures of merit were used to evaluate the operational requirements for electrical energy per order (EE/O) and the efficient use of electron delivered in terms of Faradaic efficiency. The understanding of these key aspects can help pave the way toward the large-scale industrial application of this technology for commercialization purposes.

2 Experimental

2.1 Reagents

The endocrine disruptors used for the experiments included bisphenol A (BPA, Sigma Aldrich), methyl paraben (MetP, Supelco), and paracetamol (PAC, Sigma Aldrich) with > 99% purity. Analytical grade sodium sulfate (Na₂SO₄) and iron (II) sulfate heptahydrate (FeSO₄·7H₂O), acquired from Sigma-Aldrich, were used in the experiments. Carboxylic acids with > 99% purity, acquired from Sigma-Aldrich, were used as analytical standards (*i.e.*, acetic, formic, malic and oxalic acids). The ionic species present in the water matrices were quantified by ion chromatography using standard solutions prepared from inorganic salts containing K⁺, Mg²⁺, Ca²⁺, NH₄⁺, NO₂⁻, NO₃⁻, Cl⁻, ClO₃⁻, ClO₄⁻ and SO₄²⁻ (acquired from Specsol). All solutions were prepared using ultrapure water from a Millipore Milli-Q system with resistivity >18.2 MΩ cm at 25 °C.

2.2 Aqueous matrices

The electrochemical treatments were performed in five different water matrices numbered (i)-(v) aiming at evaluating the effect of the specific components in the water matrices. First, the endocrine disruptors were treated under conventional conditions using ultrapure water containing Na₂SO₄ as single supporting electrolyte. These experiments were conducted in (i) 50 mmol L⁻¹ Na₂SO₄ solutions prepared in Milli-Q water with conductivity of 7.20 mS cm⁻¹. Second, similar conditions were employed using NaCl as alternative single supporting electrolyte – this is representative of one of the most evaluated electrolytes in the literature. Solutions of (ii) 50 mmol L⁻¹ NaCl were prepared in Milli-Q water with conductivity of 6.50 mS cm⁻¹. Third, a mixture of both electrolytes, which consisted of (iii)

25 mmol L^{-1} Na₂SO₄ + 25 mmol L^{-1} NaCl solutions in Milli-Q water with conductivity of 6.90 mS cm⁻¹, was also evaluated.

Considering that real water matrices contain complex mixtures of inorganic species, artificial water matrices were prepared using a synthetic wastewater effluent composition reported elsewhere [25]. The synthetic wastewater effluent (iv) was prepared in Milli-Q water with conductivity of 2.25 mS cm⁻¹ using the following composition: 2.10 mmol L⁻¹ KNO₃, 10.3 mmol L⁻¹ NaCl, 7.04 mmol L⁻¹ Na₂SO₄, and 1.5 mmol L⁻¹ NH₄Cl. Finally, real urban wastewater (v) was also subjected to electrochemical treatment; this effluent was collected after aerobic / anaerobic bacterial treatment from a wastewater treatment plant (WWTP) in the city of Bariri, São Paulo State, Brazil – the effluent was stored at 4 °C until its characterization (Table 1) and use.

Physical Characteristics	Values
pН	7.70
Conductivity (mS cm ⁻¹)	2.40
Total organic carbon (mg C L ⁻¹)	13.48
Chemical Characteristics	Values (mg L ⁻¹)
Ammonia (NH4 ⁺)	5.81
Calcium (Ca ²⁺)	30.94
Magnesium (Mg ²⁺)	6.07
Potassium (K ⁺)	15.54
Sodium (Na ⁺)	90.68
Chloride (Cl ⁻)	53.62
Nitrate (NO ₃ ⁻)	8.67
Nitrite (NO ₂ ⁻)	0.98
Sulfate (SO4 ²⁻)	20.43

Table 1. Physico-chemical characteristics of the real urban wastewater.

2.3 Treatment of endocrine disruptors by EAOPs

The bench-scale electrochemical assays were performed in a 150 mL electrochemical cell under controlled temperature of 25 °C. To perform the assays, commercial electrodes were used. A 2500 ppm boron-doped diamond (BDD) thin film doped on a Si substrate (acquired from NeoCoat) was used as anode and a PTFE gas diffusion electrode (supplied by Sainergy Fuel Cell) connected to a pure O₂ line with a gas flow of 500 mL min⁻¹ was used as cathode. Both electrodes had a geometric area of 3.0 cm² and were kept at an interelectrode gap distance of 1.0 cm. Experiments were conducted galvanostatically under a fixed applied current density (*j*) using a Minipa MPS-3005B DC power supply coupled to Instrutemp ITMDB ITM 100 amperemeter and Instrutemp MDB-450 voltmeter. Before the experiments, the anode and cathode were activated for 60 minutes at *j* of 100 mA cm⁻² in NaSO₄ 50 mmol L⁻¹ solution.

The EF and PEF treatments were performed using 0.5 mmol L⁻¹ Fe²⁺ as catalyst for the Fenton process. For the PEF treatment process, the solution was simultaneously illuminated using a 5W UVC lamp (Pen-Ray model 11SC-2.12) with irradiance of 19 W m⁻². Different operating conditions were tested: pH = 3, 7 and 10 (adjusted with dilute solution of 1 M H₂SO₄ or 1M NaOH); applied current density (*j*) = 16.7, 33.3 and 66.7 mA cm⁻²; initial concentration of BPA = 0.07, 0.14 and 0.28 mmol L⁻¹, with the following TOC values, 12.5, 25 and 50 ppm, respectively.

The optimized operating conditions ($j = 33.3 \text{ mA cm}^{-2}$; pH = 3, and [BPA] = 0.14 mmol L⁻¹) were used for the removal of different endocrine disruptors in urban wastewater effluent from a wastewater treatment plant. The urban wastewater effluent was contaminated with 0.14 mmol L⁻¹ BPA, MetP and PAC, which were to be removed by three electrochemical

advanced oxidation processes (EAOPs): Electrogenerated H₂O₂ coupled to electrochemical oxidation (EO-H₂O₂), EF, and PEF processes. The electrical energy per order (EE/O) for the processes was calculated based on Eq. 7 [24,26].

$$EE/O(kWh m^{-3} order^{-1}) = \frac{6.39 \times 10^{-4} P}{V_s k_1}$$
(7)

where 6.39×10^{-4} is the conversion factor (1 h/3600 s/0.4343); *P*, which, depending on the process, can be the average of the cell potential (*P*_{cell}) or the sum of *P*_{cell} with the potential of the UV lamp (*P*_{lamp}) in V; *V*_s is the volume of the solution in L; and *k*₁ is the pseudo-first order kinetic constant in s⁻¹.

2.4 Analytical determinations

The pH and conductivity were measured using ION pHB500 meter and digimed DM3, respectively.

The accumulated concentration of H₂O₂ was determined by the molybdate complex method using a Shimadzu UV-1900 spectrophotometer. The absorbance of the colored complex formed with peroxide and (NH₄)₆Mo₇O₂₄ was measured at wavelength (λ) of 350 nm [27]. The current efficiency (*CE*_{H₂O₂) associated with the generation of the accumulated H₂O₂ follows the Faraday's law described by Eq. 8 below.}

$$CE_{\rm H_2O_2}(\%) = \frac{2 \,\mathrm{F} \,\mathrm{C}_{\rm H_2O_2} V_{\rm s}}{I \,t} \times 100$$
 (8)

where 2 corresponds to the number of electrons required for O₂ reduction toward H₂O₂ production; *F* stands for the Faraday constant (96,487 C mol⁻¹); $C_{H_2O_2}$ represents the H₂O₂

concentration in mol L⁻¹; V_s is the volume of the solution in L; *I* stands for applied current intensity in A; and *t* is the reaction time in seconds.

The BPA concentration was monitored using a Shimadzu High Performance Liquid Chromatography (HPLC) model LC-20 AT, equipped with a Phenomenex Luna C18 (250 x 4.6 mm, 5 μ m) column and a Supelcosil C18 (4 x 3.0 mm i.d.) pre-column at 35 °C. The mobile phase employed consisted of a mixture of acetonitrile/water in the ratio 60:40 (v/v). The detection of the compound was performed using SPD-20A UV detector at wavelength of 280 nm, and well-defined peaks were observed in the chromatogram at retention time (t_r) of 4.7 min. Total organic carbon (TOC) was evaluated using Shimadzu TOC-VCPN equipment. TOC removal percentage was calculated based on Eq. 9.

$$TOC \ removal \ (\%) = \frac{TOC_0 - TOC_f}{TOC_0} \times 100 \tag{9}$$

where TOC_0 and TOC_f represent the total organic carbon at the initial and final time of the process (mg C L⁻¹), respectively. The current efficiency related to mineralization (MCE) was calculated using Eq. 10:

$$MCE (\%) = \frac{nFV_{s}\Delta(TOC)_{exp}}{4.32 \times 10^{7} m I t} \times 100$$
(10)

where *Vs* represents the volume of the solution (L), Δ (TOC)_{exp} is the difference between TOC₀ and TOC at time *t* (mg C L⁻¹), 4.32×10⁷ is the conversion factor (3,600 s h⁻¹ × 12,000 mg C mol⁻¹), *m* is the number of carbon atoms that belong to the BPA molecule (*m* = 15),

and *t* is specific time (h). When one considers the theoretical and total mineralization of BPA, one will find that the number of electrons (*n*) is 72 [13] (Eq. 11):

$$C_{15}H_{16}O_2 + 28 H_2O \rightarrow 15 CO_2 + 72 H^+ + 72 e^-$$
 (11)

The concentration of inorganic ions was determined by ion-chromatography using 850 Profissional IC ion chromatograph coupled to a 940 Professional IC module (Metrohm). The amount of each sample injection employed was 20 µL. The cationic species were analyzed using a Metrosep C4 column (150 mm/4.0 mm) and a Metrosep C4 Guardian/4.0 pre-column at 30 °C. The following was used as eluent: 1.7 mmol L⁻¹ nitric acid/0.7 mmol L⁻¹ dipicolinic acid at 0.9 mL min⁻¹. Analytical curves with concentrations ranging from 0 to 25 mg L^{-1} (R^2 > 0.99) were constructed for the following cationic species (retention times): Na⁺ ($t_r = 5.20$ min), NH₄⁺ ($t_r = 5.80$ min), K⁺ ($t_r = 7.70$ min), Ca²⁺ ($t_r = 16.00$ min), and Mg²⁺ ($t_r = 18.95$ min). The anionic species were analyzed in a Metrosep A Supp 5 column (150 mm/4.0 mm) and a Metrosep A Supp 5 Guardian/4.0 pre-column at 30 °C, using an eluent composed of 3.2 mmol L⁻¹ Na₂CO₃/1.0 mmol L⁻¹ NaHCO₃ at 0.7 mL min⁻¹. An analytical curve with concentrations ranging from 0 to 100 mg L⁻¹ was constructed for each of the following anionic species: Cl⁻ ($t_r = 5.45 \text{ min}$), NO₂⁻ ($t_r = 6.60 \text{ min}$), ClO₂⁻ ($t_r = 4.30 \text{ min}$), ClO₃⁻ ($t_r = 4.30 \text{ min}$)), ClO₃⁻ ($t_r = 4.30 \text{$ 8.00 min), NO₃⁻ ($t_r = 9.75$ min), PO₄³⁻ ($t_r = 13.50$ min), and SO₄²⁻ (15.70 min). The carboxylic acids generated were analyzed using the ion chromatograph mentioned previously. The following was used for the conduct of the analysis - separation column: Metrosep Organic Acids (250 mm/7.8 mm) from Metrohm; solvent: a diluted H₂SO₄ solution (0.75 mmol L⁻¹); and suppressor: NaCl 50 mmol L⁻¹. Analytical curves of different carboxylic acids were constructed for concentrations ranging between 0-100 mg L^{-1} ($R^2 > 0.99$) using the following:

oxalic acid ($t_r = 7.60 \text{ min}$), malic acid ($t_r = 9.85 \text{ min}$), acetic acid ($t_r = 13.60 \text{ min}$), and formic acid ($t_r = 15.25 \text{ min}$).

3 Results and discussion

3.1 Generation and accumulation of H₂O₂ in media of different complexity

Sulfate and chloride are ionic species which are commonly present in wastewater. These inorganic ions provide the essential conductivity required for the conduct of electrochemical treatments. Apart from sulfate and chloride, real wastewater also contains other ions and compounds, and this makes the matrix more complex to handle. Fig. 1a illustrates the effect of the electrolyte on the accumulation of electrogenerated H₂O₂ during electrolysis conducted for a period of 360 min in an undivided cell at pH 3.0, with current density of 33.3 mA cm^{-2} . As can be observed, regardless of the medium applied, there is a progressive accumulation of H_2O_2 caused by its continuous electrogeneration from oxygen reduction via reaction (2), which is followed by the occurrence of a plateau after 240 min of treatment. This behavior is attributed to the simultaneous degradation of accumulated H₂O₂ through its oxidation at the anode or further reduction to water at the cathode, which leads to the generation of a *pseudo*steady concentration. Despite the similarities observed for sulfate and chloride media, there is a marked difference in the maximum concentration obtained at the steady state. When sulfate is used as medium, a relatively higher H_2O_2 (26.6 mmol L⁻¹) concentration is obtained compared to that obtained when chloride is used as medium (21.2 mmol L⁻¹). The relatively lower H₂O₂ accumulation observed in chloride media can be attributed to the role played by electrogenerated active chlorine species in the process. The oxidation of chloride ion on the surface of the anode leads to the generation of chlorine ($Cl_{2(aq)}$, E°= 1.36 V vs SHE) based on Eq. (12). The hydrolysis of chlorine yields hypochlorous acid (HClO, E°= 1.49 V vs SHE) and Cl⁻ (Eq. 13) [28]. The pH plays an important role in the distribution of active chlorine species by defining their speciation. For instance, HClO has an acid-base equilibrium described in Eq. (14), with pKa = 7.55, which may lead to the preferential accumulation of the weaker oxidant hypochlorite (OCl⁻, E° = 0.89 V vs SHE) [29]. Electrogenerated active chlorine species are strong oxidants that can scavenge H₂O₂ [30]. The experiment conducted using mixed electrolyte composition of 25 mmol L⁻¹ Na₂SO₄ and 25 mmol L⁻¹ NaCl generated a maximum accumulated H₂O₂ of 23.9 mmol L⁻¹ between the individual electrolytes. This result is in agreement with the scavenging role of active chlorine species which are accumulated at a lower rate for lower concentrations of chloride.

$$2Cl^{-} \rightarrow Cl_{2(aq)} + 2e^{-} \qquad pH < 3 \qquad (12)$$

$$Cl_{2(aq)} + H_2O \rightarrow HClO + Cl^- + H^+ \qquad 3 < pH < 8$$
(13)

$$HClO \rightleftharpoons OCl^- + H^+ \qquad pH > 8 \qquad (14)$$



Fig. 1. a) Accumulation and b) current efficiency of H_2O_2 over time. Operating conditions: EO-H₂O₂ at pH 3.0 and temperature of 25 °C using a stirred undivided cell with BDD anode and carbon-PTFE cathode, at current density (*j*) = 33.3 mA cm⁻², in different media: (•) 50 mmol L⁻¹ Na₂SO₄, (•) 25 mmol L⁻¹ Na₂SO₄+ 25 mmol L⁻¹ NaCl, (•) 50 mmol L⁻¹ NaCl, (•) synthetic effluent, and (•) real wastewater.

With regard to the electrogeneration and accumulation of H_2O_2 for the simulated and real effluents, lower H_2O_2 concentrations were obtained – approximately 17.6 mmol L⁻¹ and 16.1 mmol L⁻¹, respectively, compared to the previous experiments using Na₂SO₄, Na₂SO₄ + NaCl, and NaCl. The increase in complexity of the medium contributed toward a slight decrease in concentration of the oxidant generated due to the increased probability of the number of parasitic reactions that were expected to occur as a result of the additional inorganic ions and the natural organic matter that were present in the medium.

The evaluation of the Faradaic efficiency provides a quantitative understanding of the impact of scavenging oxidants in the solution investigated. Fig. 1b illustrates the gradual decrease of overall efficiency over time of the electrogeneration of H₂O₂ due to the coexistence of parasitic reactions that consume this oxidant (Eqs. 15-19) [30–32]. The most noticeable difference associated with the electrolyte composition can be observed at the beginning of the experiments where the $CE_{H_2O_2}$ values were 74.1% for Na₂SO₄, 60.6% for Na₂SO₄+NaCl, 44.5 % for NaCl, 34.2% for the synthetic effluent, and 29.3% for the real wastewater. Note that this implies a noticeable 2.5-fold decrease in H₂O₂ electrogeneration efficiency in the real water samples, which is attributed to the constituents of this effluent, compared to the solution containing only Na₂SO₄. This dramatic decrease in $CE_{H_2O_2}$ (loss of *about* 45%) may undermine the performance of H₂O₂-based AOPs and should be considered during the implementation of the technology in industrial scale production. Fortunately, the experimental results show that the amount of electrogenerated H₂O₂ required for the supporting electrolyte employed.

$$BDD + H_2O_2 \rightarrow BDD(HO_2^{\bullet}) + H^+ + e^-$$
(15)

$$H_2O_2 + 2H^+ + 2e^- \rightarrow 2H_2O$$
 (16)

$$H_2O_2 + Cl_2 \rightarrow O_2 + 2H^+ + 2Cl^-$$
 (17)

$$H_2O_2 + HOCI \rightarrow O_2 + H_2O + H^+ + Cl^-$$
 (18)

$$H_2O_2 + OCl^- \rightarrow O_2 + H_2O + Cl^-$$
⁽¹⁹⁾

3.2 Impacts of water matrix on the efficiency of EAOPs in the removal of endocrine disruptors

Bisphenol A (BPA) is a common endocrine disruptor found in natural water bodies. The present work investigated the effect of water matrix composition on the degradation of 0.14 mmol L⁻¹ BPA through the application of different EAOPs under comparable conditions of pH 3.0 and applied density of 33.3 mA cm⁻².

The nature of the EAOPs employed has a direct impact on the kinetics of BPA degradation (decay) as can be noted in Fig. 2. Slower degradation rates were observed for the mass transfer limited EO-H₂O₂ process, while the Fenton-based processes recorded 4.0-25 times faster BPA removal (Fig. 2 - b, d, f). Here, one will notice the remarkably faster removal rate observed between the treatment processes where the electrolysis time of 60 min for the EO-H₂O₂ process was reduced down to roughly 10 min for electrochemically driven Fenton processes (Fig. 2a, c, e). These results point to the key role played by two factors in accelerating the degradation of BPA: i) the *in situ* generation of homogeneous **•**OH radicals, derived from the Fenton reaction (3); and ii) the role of UVC light in the photolytic process described in reaction (20). The results obtained here are in agreement with the findings published previously in the literature which showed that the application of electro-Fenton techniques led to higher removal of organic persistent pollutants from water matrices [33,34].

$$H_2O_2 + hv \rightarrow 2^{\bullet}OH$$

Even though similar patterns of behavior were observed for different water matrices when the EAOPs were compared under certain conditions, a closer look at the data revealed that the water matrix played a major role in the kinetics of BPA removal for each individual electrochemically-driven technique. These differences are more noticeable for the EO-H₂O₂ process where different electrolytes greatly influenced the degradation kinetics. The degradation of BPA via the EO-H₂O₂ treatment process in the presence of Na₂SO₄ led to a gradual decrease of BPA concentration, where 39.6% BPA removal was obtained after 60 min of treatment. For purposes of comparison, the presence of chloride anion enhanced BPA removal with a sharp increase in the contaminant decay. For the treatments conducted using Na₂SO₄+NaCl and NaCl, the complete removal of BPA was achieved after 30 and 60 min of treatment. The corresponding degradation rate constants (k_1) obtained under the different media employed were 2.12x10⁻⁴ s⁻¹ ($R^2 = 0.998$), 9.22x10⁻⁴ s⁻¹ ($R^2 = 0.993$), and 1.98x10⁻³ s⁻¹ (R²=0.996), for Na₂SO₄, NaCl, and Na₂SO₄+NaCl, respectively. This result can be attributed to the presence of sulfate and chloride species in the water matrix and their possible coelectrogeneration of additional oxidants apart from $^{\circ}$ OH: peroxydisulfate (S₂O₈²⁻, E⁰ = 2.01 V vs SHE, Eq. 21) and active chlorine species (Eqs. 12-14). A comparison between peroxydisulfate and active chlorine species shows that the latter has relatively greater oxidizing power [9]. It is worth noting that, compared to •OH radicals, these extra oxidants - peroxydisulfate and active chlorine species, have weaker oxidation potentials, and have mainly distinct homogeneous character. These electrogenerated oxidants are yielded in the bulk of the solution.

$$SO_4^{2-} + SO_4^{2-} \rightarrow S_2O_8^{2-} + 2e^-$$
 (21)

Although the presence of sulfate and chloride improves the pollutant degradation, depending on the concentration of these species, they may also act as scavengers of •OH radicals (Eqs. 22 - 26) [35]. This deleterious effect is observed for the experiment with 50 mmol L⁻¹ NaCl. The Cl⁻ concentration evolution was followed over time and its conversion occurred as follows: 4.5%, 9.3% and 22.1% for the NaCl, Na₂SO₄+NaCl, and synthetic effluent water matrices, respectively. Based on the results obtained, the conversion of higher amount of Cl⁻ implied the generation of greater amount of active chlorine species.

$$\mathrm{SO4}^{2-} + \mathrm{^{\bullet}OH} \to \mathrm{SO4}^{-\mathrm{^{\bullet}}} + \mathrm{OH}^{-}$$
 (22)

$$Cl^{-} + {}^{\bullet}OH \rightleftharpoons ClO^{-} + H^{+} + e^{-}$$
(23)

$$ClO^{-} + {}^{\bullet}OH \rightleftharpoons ClO_{2}^{-} + H^{+} + e^{-}$$
(24)

$$ClO_2^- + {}^{\bullet}OH \rightleftharpoons ClO_3^- + H^+ + e^-$$
(25)

$$ClO_3^- + {}^{\bullet}OH \rightleftharpoons ClO_4^- + H^+ + e^-$$
(26)

The fundamental understanding derived from these experiments on simple electrolytes helped interpret the results obtained from the treatment of the simulated and real wastewater matrices (Fig. 2a). Apart from the effect of inorganic electrolytes, one also needs to point out the effect of organic matter present in real water samples (see Table 1). Organic matter that naturally exists in water effluents compete with the target pollutant (*i.e.*, BPA) for the electrogenerated oxidants (e.g., •OH radicals, active chlorine species, etc.), and this contributes to the decline in the removal efficiency. This undesired effect was found to be evident when the rate of BPA degradation in the real wastewater was compared with that observed for the simulated wastewater effluent in the absence of the organic species.

The results obtained from the experiments also showed that other inorganic compounds, such as potassium, nitrate, and ammonia, did not exert any significant influence on the decay of the contaminant under the EO-H₂O₂ treatment technique.

Similar patterns of behavior were observed for the EO-H₂O₂ treatment mechanism with regard to the pollutant mineralization in terms of total organic carbon (TOC) removal (Table 2). Discrete TOC removal rates ranging between 14.8-31.5% were obtained after 60 min of treatment by EO-H₂O₂; this range of TOC removal rates is commonly observed for electrochemical oxidation processes. This outcome can be attributed to the mass transfer limitation and the quantity of intermediate compounds generated which can have a more recalcitrant character than the compounds initially present in the wastewater effluent. Interestingly, the highest TOC removal (31.5%) was obtained under the treatment using the simulated wastewater effluent. The higher degree of mineralization obtained under the treatment with simulated wastewater effluent may be attributed to the solution composition. The synthetic medium contains NO₃⁻ and NH₄⁺, in addition to sulfate and chloride ions. On the one hand, the NH4⁺ cation has negligible impact on the mineralization of pollutants (the aspects related to its reactivity with active chlorine species will be discussed in detail later). On the other hand, according to recent literature reports [36], the use of water matrices with NO₃⁻ can yield a higher TOC removal due to N-containing species produced from NO₃⁻ electro-reduction which may contribute toward the mineralization of by-products, leading to higher mineralization. Finally, the result obtained for mineralization current efficiency estimated based on Eq. 10 indicated the following in terms of process efficiency: Synthetic effluent > $NaCl + Na_2SO_4 > NaCl > Na_2SO_4$. This result is in agreement with the results obtained for TOC removal shown in Table 2.

The results obtained related to the effect of the composition of the water matrix showed comparable patterns of behavior for the electrochemically-driven Fenton processes. The contaminant degradation was found to occur faster in the simulated effluent (k_1 =7.23×10⁻³ s⁻¹ – EF, k_1 =9.05×10⁻³ s⁻¹ – PEF), and this was closely followed by Na₂SO₄+NaCl (k_1 =5.85×10⁻³ s⁻¹ – EF, k_1 =6.91×10⁻³ s⁻¹ – PEF), real wastewater (k_1 =5.11×10⁻³ s⁻¹ – EF, k_1 =5.93×10⁻³ s⁻¹ – PEF), and Na₂SO₄ (k_1 =4.45×10⁻³ s⁻¹ – EF, k_1 =5.34×10⁻³ s⁻¹ – PEF) (Figs. 2 - c, d, e, f, and Table 2). The slightly faster degradation kinetics of the endocrine disruptor observed under the PEF treatment technique can be attributed to the role played by irradiation in the process. The use of UVC irradiation can yield additional °OH radicals from H₂O₂ photolysis (Eq. 28). Furthermore, UVC irradiation can activate other electrogenerated oxidants, which can, in turn, lead to the formation of higher oxidizing species, such as sulfate radical SO4^{-•} based on reaction (27) or chlorine radical (Cl[•]) via reactions (28) and (29).

$$S_2 O_8^{2-} + h\nu \to 2 S O_4^{-\bullet} \tag{27}$$

$$HOCl + hv \to {}^{\bullet}OH + Cl^{\bullet}$$
⁽²⁸⁾

$$ClO^{-} + hv \to O^{-\bullet} + Cl^{\bullet}$$
⁽²⁹⁾

The TOC removal rates for the matrices investigated saw between 1.2-1.8 fold increase for EF and between 1.4-2.4 fold increase for PEF compared to EO-H₂O₂ (Table 2). The higher mineralization percentage obtained for the PEF process is related to photodecarboxylation reactions induced by light irradiation based on reaction (6). In terms of BPA mineralization, the superior mineralization efficiency observed for the Fenton-based processes over the EO-H₂O₂ process was also reflected in the slightly higher MCE values obtained for the former, which ranged between 10.9 - 17.7% (Table 2). The type of water matrix and the electrochemical process suitable for the treatment of endocrine disruptors (EDs) are key elements that need to be considered when one seeks to obtain high rates of degradation. With that in mind, the next step of this study aimed to elucidate the applicability of EAOPs in the treatment of a mixture of EDs in real matrices.



Fig. 2. Normalized BPA concentration decay over time (a, c, d) and corresponding pseudo-first-order kinetic analysis (b, d, f) for EO-H₂O₂ (a, b), EF (c, d) and PEF (e, f) processes, respectively. Operating conditions: a stirred undivided cell with BDD anode and carbon-PTFE cathode, with pH of 3.0, j = 33.3 mA cm⁻², and temperature of 25 °C, using

different media (●) 50 mmol L⁻¹ Na₂SO₄, (▼) 25 mmol L⁻¹ Na₂SO₄+ 25 mmol L⁻¹ NaCl,
(■) 50 mmol L⁻¹ NaCl, (▲) synthetic wastewater effluent and (◆) real wastewater.

Table 2. Key fitted and calculated parameters obtained from the experiments using EO-H₂O₂, EF and PEF for the oxidation of 0.14 mmol L^{-1} bisphenol A in different media at pH 3 and current density of 33.3 mA cm⁻².

Experiment	$k_1 imes 10^{-3} / \text{ s}^{-1}$	R ²	TOC removal / %*	MCE / %
EO-H ₂ O ₂				
Na ₂ SO ₄	0.212	0.998	14.8	5.95
NaCl	0.922	0.993	22.8	9.13
Na ₂ SO ₄ +NaCl	1.34	0.997	28.3	11.4
Simulated	1.82	0.995	31.5	12.7
Real**	0.391	0.992	18.3	-
EF				
Na ₂ SO ₄	4.45	0.993	27.2	10.9
Na ₂ SO ₄ +NaCl	5.85	0.994	35.5	12.9
Simulated	7.23	0.992	38.7	15.6
Real**	5.11	0.997	28.6	-
PEF				
Na ₂ SO ₄	5.34	0.995	35.1	14.1
Na ₂ SO ₄ +NaCl	6.91	0.993	40.1	16.4
Simulated	9.05	0.998	44.2	17.7
Real**	5.93	0.992	35.4	-

*After 60 min of treatment.

**TOC0=38 mg C L-1

3.3 Electrochemical treatment of real urban wastewater containing a mixture of endocrine disruptors

Polluted waters may contain more than one recalcitrant endocrine disruptor. A more complex matrix is certainly more complicated to handle during the treatment process. Figure 3 shows the impact of coexisting pollutants methyl paraben (MeP) and paracetamol (PAC) in the degradation of BPA. Looking at Figs. 3a and 3b, one will observe that the degradation kinetics of BPA decreased significantly (2.4 - 3.0x) in this real wastewater medium under the application of the EAOPs. The coexistence of persistent organic pollutants decreased the kinetic rate of BPA removal under the EO-H₂O₂ treatment process from $3.91 \times 10^{-4} \text{ s}^{-1}$ in the real effluent down to $1.64 \times 10^{-4} \text{ s}^{-1}$. Given that the operational conditions were the same, this decline in BPA removal can be explained by the non-selective nature of •OH radical and other oxidants. Thus, the coexisting pollutants compete for the same amount of oxidants electrogenerated, and this leads to a decrease in the overall removal rate.

Interestingly, the Fenton-based processes were also affected by the presence of species that competed for the electrogenerated oxidants. In the case of the EF process, which attained complete BPA removal in the real matrix after 10 min of treatment, only 65.9% of BPA was degraded in the same matrix when MeP and PCA were present at the same initial concentration. Instead, 40 min of EF treatment were required to degrade BPA in the solution containing the interfering species. The competition for hydroxyl radical between the different pollutants decreased the pseudo first-order rate constant of BPA removal in real water from $5.11 \times 10^{-3} \text{ s}^{-1}$ down to $1.72 \times 10^{-3} \text{ s}^{-1}$ (almost a 3-fold decrease). The PEF process exhibited identical pattern of behavior in which the kinetic constant of BPA removal declined from $5.93 \times 10^{-3} \text{ s}^{-1}$ to $2.13 \times 10^{-3} \text{ s}^{-1}$. Depending on the nature of the coexisting pollutants, the light employed in the process may be absorbed by the organic substances in solution, decreasing

its transport. However, the main cause of the decline in kinetic constant seemed to be associated with the competitive consumption of electrogenerated oxidants.

The study of the mineralization of the solution investigated gave us the results discussed previously. The rate of mineralization obtained was in the following order: $PEF >> EF > EO-H_2O_2$, where the use of only direct photolysis was insufficient to achieve the incineration of the organic substances (commonly observed for photostable compounds).

The experiment using EO-H₂O₂ presented a steady TOC decay, leading to 40.8% of TOC removal after 300 min of treatment. This gradual mineralization is attributed to the presence of heterogeneous BDD(•OH) and the control of organic substances by mass transport from/toward the anode surface. The test conducted under the EF process exhibited a noticeable total organic carbon removal of about 53.6% during the first 180 min, remaining practically constant thereafter. This quick initial mineralization may be linked to the oxidation of some intermediates by the homogeneous •OH radicals and the weaker oxidants electrogenerated in the bulk, which can lead to the formation of recalcitrant compounds that are sluggishly destroyed by these same oxidants or by BDD(•OH). These recalcitrant compounds have been reported to be mainly composed of carboxylic acids which form stable complexes with iron (III). The formation of these complexes leads to the trapping of the catalyst in the Fenton reaction process, slowing down the magnitude of the reaction. In addition, these stable organo-complexes have demonstrated to be more resistant to oxidation than non-complexed acids. These effects have been associated with the strong deceleration of TOC removal at longer electrolysis periods under the EF treatment process [37,38]. The use of the PEF treatment process enabled us to overcome this constraint through the application of UVC light which promoted the photodecarboxylation of the complexes, leading to almost complete TOC removal from the water matrix containing the mixture of

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endocrine disruptors at the end of the treatment time. To improve our understanding of the process involving the generation of intermediate compounds, the concentration of shortchain carboxylic acids – which are final by-products of the PEF process, was analyzed by HPLC. Fig. 4c shows the evolution of the concentration of acetic, formic, malic, and oxalic acids over time in the degradation process of real wastewater containing 0.14 mmol L⁻¹ BPA, 0.14 mmol L⁻¹ MetP, and 0.14 mmol L⁻¹ PAC. There was a gradual maximum accumulation of the carboxylic acids tested and all of them were practically removed after 300 min of treatment. The acetic acid, which was probably obtained mainly from the cleavage of the benzene rings of the endocrine disruptors, presented a maximum accumulation of 17.70 mg L⁻¹ during the first 60 min. Formic, malic, and oxalic acids persisted in the oxidation process of the carboxylic acids and presented maximum accumulations of 14.19, 3.96 and 22.32 mg L^{-1} at 120 min, 60 min, and 60-120 min, respectively. These acids were directly mineralized to CO₂. The main conclusion that can be drawn here is that the EAOPs are capable of completely mineralizing the organic load of polluted effluents irrespective of the nature of the persistent organic pollutants contained in the water matrix. However, the degradation kinetics of a specific target pollutant (such as BPA) may experience a significant decline in water matrices containing many interfering pollutants compared to pure water or real matrices containing only one target pollutant. In this context, complete TOC removal may be a straightforward indicator of water quality for researchers or analysts when the EAOP treatment technique is applied in centralized treatment plants.



Fig. 3. a) Normalized BPA concentration decay in a mixture of different endocrine disruptors (0.14 mmol L⁻¹ BPA, 0.14 mmol L⁻¹ methyl paraben, and 0.14 mmol L⁻¹ paracetamol) over time using EO-H₂O₂, EF and PEF processes and b) respective pseudo-first order kinetics. c) Normalized TOC decay in a mixture of different endocrine disruptors over time using photolysis, EO-H₂O₂, EF, and PEF processes. d) Evolution of the carboxylic acids concentration over time under the PEF process. Operating conditions: real wastewater, pH 3.0, j = 33.3 mA cm⁻², 0.5 mmol L⁻¹ Fe²⁺ used as catalyst for the Fenton reactions, and UVC lamp for the PEF.

3.4 Optimization of operating conditions for the electrochemical oxidation process

Although Fenton-based processes are regarded highly competitive, the techniques are found to be limited to a narrow acidic pH range of about 3.0. Indeed, this is a major limitation when it comes to the treatment of large volumes of matrices that require pH adjustment prior and after the treatment. In contrast, EO-H₂O₂ can be potentially competitive under extensive

range of operation conditions. The present study sought to investigate the influence of different variables on the degradation efficiency and competitiveness of EO-H₂O₂. Fig. 4 illustrates the impact of different operating conditions, which include pH, applied current density, and initial BPA concentration, on the performance of the EO-H₂O₂ treatment process.

The pH is one of the major constraints in the search for competitive treatment techniques for implementation. Fig. 4a shows a steady decrease in the contaminant degradation with the increase in pH values based on the application of the EO-H₂O₂ treatment process. A similar pattern of behavior was also observed for TOC removal. BPA removal of 71% was obtained for the contaminant concentration at pH 10 after 60 min of treatment ($k_1 = 3.17 \times 10^{-4} \text{ s}^{-1}$, R² = 0.995). Furthermore, an increase of 1.8-fold ($k_1 = 5.84 \times 10^{-4} \text{ s}^{-1}$, $R^2 = 0.993$) and 5.7-fold (k_1 1.82 x10⁻³ s⁻¹, $R^2 = 0.995$) in k_1 was observed for pH 7.0 and pH 3.0, respectively. Under the acidic medium, the use of the EO-H₂O₂ treatment process led to a sharp removal of BPA (89.3%) in the first 20 min of treatment, and the compound disappeared completely after 40 min. In both water matrices, the EO-H₂O₂ process was found to be mostly influenced by the BDD([•]OH) radicals and the electrogeneration of active chlorine species. As discussed previously, the presence of chloride enhances the degradation kinetics due to the action of homogeneous active chlorine species in the bulk. However, the oxidation capabilities of active chlorine species are intimately related to the pH, which also defines their speciation. Indeed, the oxidation of the pollutant in the presence of chlorine species is more extended in acidic than in alkaline media due to the higher standard potential of HClO ($E^0 = 1.49$ vs. SHE) compared to ClO⁻ ($E^0 = 0.89$ vs. SHE) [29]. The electrogeneration of hydroxyl radical and its oxidation capabilities are also affected by the pH [39] though not as strongly as the effect observed in chlorine species [40]. This observation can help explain the pattern of

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behavior similarly observed for the TOC removal in the inset of Fig. 4a. The energy required to remove one unit of BPA (EE/O) was determined for each pH in the simulated medium based on Eq. 7. The experiment at pH 3.0 yielded EE/O value of 2.80 kWh m⁻³ order⁻¹; this value is 3.3-fold and 6.3-fold inferior to the values obtained at pH 7.0 and 10, respectively (Fig. 4b). The results of these experiments showed that the best initial pH to work with is the acidic pH (~3.0), although the major strength of EO-H₂O₂ is its possible application under the natural pH of water effluents (6.0-8.0).

The applied current density *j* is among the key parameters that exert influence over the electrokinetics of EAOPs. The current density defines the quantity of electrons that flow in the system and, consequently, the amount of oxidizing mediator species that can be electrogenerated [41]. The influence of current density was assessed in the range of 16.7 to 66.7 mA cm⁻². Fig. 4c shows that an increase in the applied *j* led to a decrease in the normalized concentration decay of BPA. These results help infer that the higher amounts of •OH radicals, peroxydisulfate, and active chlorine electrogenerated in the process contributed toward the increase in the degradation kinetics. In fact, the k_1 was doubled when the applied *j* was increased from 16.7 mA cm⁻² to 33.3 mA cm⁻². Nevertheless, almost no variation $(\sim 1.3x)$ was observed when the applied current density was increased further to 66.7 mA cm⁻ ² (Fig. 4d). These results allow one to infer that there is a transition from degradation kinetics limited by current to a degradation kinetics limited by the transport of pollutants toward the electrode surface. The excess of electrons in the process is consumed by parasitic reactions that do not lead to organic degradation reactions, but that consume/scavenge •OH radicals, as can be observed in reactions (22-26) and (31-33). Parasitic reactions affect the efficiency of the system and may unnecessarily increase the operational costs. The same pattern of behavior is also evident for the TOC removal, where the experiments conducted at current

densities of 16.7, 33.3, and 66.7 mA cm⁻² yielded TOC removal of 23.9%, 35.1%, and 39.8%, respectively (inset of Fig. 4c). Higher current densities resulted in higher degradation kinetics, which was, in turn, expected to lead to lower EE/O (Eq. 7); however, EE/O was found to be more affected by the increase of the cell potential (P_{cell}). Thus, based on the techno-economic aspects, one ought to find a middle-ground solution that minimizes retention time in the electrochemical cells while requiring an acceptable energy input. This parameter should be then defined for each specific case scenario.

$$BDD(^{\bullet}OH) \rightarrow BDD + \frac{1}{2}O_2 + H^+ + e^-$$
(31)

$$H_2O_2 + {}^{\bullet}OH \rightarrow HO_2 {}^{\bullet} + H_2O$$
(32)

$$2^{\bullet}\mathrm{OH} \to \mathrm{H}_{2}\mathrm{O}_{2} \tag{33}$$

An analysis of the impact of the initial concentration of BPA (0.07-0.28 mmol L⁻¹) provides valuable information regarding the range of concentration that can be efficiently treated within reasonable periods of time, as illustrated in Fig. 4e. As can be observed, a decrease in the initial concentration of the pollutant led to an increase in the degradation kinetics from $8.01 \times 10^{-4} \text{ s}^{-1}$ (R²= 0.982) at 0.28 mmol L⁻¹ to $1.53 \times 10^{-3} \text{ s}^{-1}$ (R²= 0.995) for 0.14 mmol L⁻¹ BPA. A further decrease of the concentration to 0.07 mmol L⁻¹ BPA also led to an increase of up to $2.78 \times 10^{-3} \text{ s}^{-1}$ (R²= 0.996) in k_1 (see Fig. 4f). Conducting experiments with higher concentrations of contaminants leads to the generation and accumulation of a larger number of intermediates and by-products, which tend to compete for the same amount of •OH radicals with the initial compounds. This ultimately leads to a decline in the pollutant degradation and an increase in EE/O (Fig. 5f). In the case of total organic carbon (TOC), the removal efficiency is found to be superior for the experiment with higher initial BPA concentration (inset Fig. 5e). It is worth noting that

mineralization depends on the effective transport of BPA to the electrode surface, which improves in the presence of larger organic load. Indeed, when there is a higher probability of provoking the formation of oxidants for organic degradation reactions, this tends to diminish the extent/magnitude of parasitic reactions. However, it has become clear that electrochemical processes can be applied in a wide range of concentrations, and these processes can be highly effective for the removal of trace pollutants when they are applied in combination with one another in wastewater treatment plants. Similar behavior was observed in real water matrices (see Fig. S1).



Fig. 4. Effect of a) initial pH, c) current density, and e) initial pollutant concentration on the decay of BPA over time. Insets: Corresponding TOC removal of the parameters after 60 min of treatment. Kinetic analysis based on a pseudo-first order degradation reaction (■) versus electrical energy per order (■) for b) initial pH, d) current density, and f) initial pollutant concentration.

3.5 Time-course analysis of coexisting inorganic ions during the treatment of real wastewater

Apart from the degradation performance, the time-course analysis of the ions in the solution under treatment is also a matter of great interest to this study. The inorganic ions were monitored during the treatment of [BPA]=0.14 mmol L⁻¹ at pH 3.0 under the EO-H₂O₂ treatment process at current density of 33.3 mA cm⁻². Looking at Fig. 5, one will observe that 71% of the initial chloride species were converted into active chlorine species. However, further oxidation of the inorganic chlorine species led to the formation of 1.23 mg L⁻¹ ClO₂⁻⁻ , 1.82 mg L⁻¹ ClO₃⁻, and 0.98 mg L⁻¹ ClO₄⁻. Chlorate and perchlorate are undesired species that may pose serious health risks to living organisms [42–44]. Under the framework of the World Health Organization (WHO), the maximum concentration level (MCL) of chlorate and perchlorate allowed in drinking water is 70 µg L⁻¹ [45]. The US Environmental Protection Agency (US-EPA) has established an even more stringent MCL of 15 µg L⁻¹ for chlorate and perchlorate in drinking water. Thus, the results obtained in the present study suggest that the treated water will not be suitable for drinking purposes without conducting additional treatment aimed at further reducing ClO3⁻ and ClO4⁻. However, one will note that the concentration level of these undesired species observed in the treated water may be in full compliance with the specified maximum effluent limit of up to 10 mg L⁻¹ of ClO₄⁻ allowed under the wastewater discharge permit in different wastewater treatment facilities [46].

N-species are also susceptible to chemical and electrochemical transformations. Chromatographic analyses show that the initial ammonia $(NH4^+)$ concentration of 5.81 mg L⁻¹ (4.5 mg L⁻¹ as N) was completely removed at the end of the treatment. The complete

disappearance of NH₄⁺ from the solution can be explained by its chemical reaction with the active chlorine species electrogenerated in the process. The chemical conversion of ammonia to innocuous nitrogen gas (N₂) occurs when a certain concentration of active chlorine species is obtained. This process occurs based on reaction (34), and it is referred to as the chlorine breaking point.

$$3HOCl + 2 NH_3 \rightarrow N_2 + 3H_2O + 3HCl$$

$$(34)$$

Under the EO-H₂O₂ treatment process, the initial nitrate (NO₃⁻) concentration was found to decrease from 8.67 mg L⁻¹ to 1.08 mg L⁻¹ (0.24 mg L⁻¹ as N) after 60 min of electrolysis. This behavior can be attributed to the gradual cathodic reduction of nitrate via direct charge transfer process, which is not highly favored in carbonaceous electrodes [47]. Nitrate reduction can yield nitrite (NO₂⁻) and NH₄⁺, based on reactions (35) and (36), respectively. Furthermore, NH₄⁺ evolves into N₂ based on reaction (37), as described previously; thus, this ion is not accumulated in the solution.

$$NO_3^- + 2 H^+ + 2 e^- \rightarrow NO_2^- + H_2O$$
 (35)

$$NO_{3}^{-} + 10 H^{+} + 8 e^{-} \rightarrow NH_{4}^{+} + 3 H_{2}O$$
(36)

$$2 \text{ NH}_4^+ + 8 \text{ H}^+ + 10 \text{ e}^- \rightarrow \text{N}_2 \tag{37}$$

The concentration of NO₂⁻ changed gradually from 0.98 mg L⁻¹ (0.30 mg L⁻¹ as N) to 0.93 mg L⁻¹ (0.28 mg L⁻¹ as N), and NO₂⁻ was further reduced to NH₄⁺, but the sustained yield of NO₂⁻ as stable by-product of NO₃⁻ electrochemical reduction maintained the level of NO₂⁻ during the electrolysis time under analysis. It is worth pointing out that the remaining concentrations of NO₂⁻ and NO₃⁻ were found to be below the MCLs for drinking water, which have been established to be 1.0 mg L⁻¹ and 10.0 mg L⁻¹ for NO₂⁻ and NO₃⁻, respectively.

These results show that there are some key water quality parameters that must be accounted for when it comes to assessing the efficiency of electrochemical treatments of organic pollutants. One will be able to thoroughly evaluate the full picture of the competitiveness of the treatment process once these quality parameters are accounted for. Although the Nspecies have met the quality criteria for drinking water, further treatment needs to be conducted in order to decrease the concentration of chlorine oxyanions.



Fig. 5. Nitrogen and chloride-based species identified at the end of 60 min of treatment. Operating conditions – pollutant concentration: 0.14 mmol L⁻¹ BPA; treatment process: EO- H_2O_2 ; pH 3.0; current density (j) of 33.3 mA cm⁻²; and temperature of 25 °C, using stirred undivided cell with BDD anode and carbon-PTFE cathode in real urban wastewater.

4 Conclusions

Having a comprehensive understanding of the effect of the complexity of different water matrices on the removal of endocrine disruptors is key to identifying specific areas for the

application of electrochemical techniques based on H₂O₂ electrogeneration. The results of the present study show that H₂O₂ oxidant can be generated and accumulated irrespective of the medium employed in the treatment of both simple and complex matrices. The application of three different EAOPs for the treatment of different media containing BPA showed that the process involving the contaminant removal occurred in different ways under the techniques investigated and with different levels of efficiency. The Fenton reactions and the presence of UVC light promoted higher degradation rates (about 4.0-25 times higher than the EO-H₂O₂ process) due to the greater amount of homogeneous [•]OH radicals generated in the system. The constituents of the water matrix, such as sulfate and chloride, and their concentration were found to play a key role in the generation of additional oxidants, such as peroxydisulfate and active chlorine species, which are regarded essential for the removal of endocrine disruptors. The undesired organic compounds present in the real wastewater investigated were found to affect the performance of the systems because they competed naturally with the target pollutant for the oxidants electrogenerated in the treatment process. Inorganic species, such as potassium, nitrate, and ammonia, showed no significant interference in the degradation of the contaminant. The coexistence of persistent organic pollutants (POPs) with the target pollutant (BPA) in the real wastewater investigated in this study increased the interference of these POPs in the degradation of BPA. Using the EO-H₂O₂ treatment process and under the application of the same operating conditions adopted in the other treatment processes, the kinetic rate of BPA decreased from $3.91 \times 10^{-4} \text{ s}^{-1}$ to $1.64 \times 10^{-4} \text{ s}^{-1}$ due to the higher competition for the same amount of oxidants from the intermediate species and new by-products generated in the process. Regarding the analysis of TOC removal, the formation of a plateau was clearly observed during the EF process; this was probably due to the generation of sub-products such as iron complexes which are

difficult to remove. This problem was overcome by the application of UVC light in the PEF process. Based on the time-course analysis of carboxylic acids, the results obtained show that the application of the PEF treatment mechanism can lead to a complete removal of the target compound from the system regardless of the nature of the competing persistent organic pollutants present in the wastewater matrix. Although the Fenton-based processes are highly competitive techniques, their acidic pH range requirement of approximately 3.0 hinders their broader applicability and implementation; this paves the way toward the use of the EO-H₂O₂ process since it provides one with the possibility of conducting the treatment under a wide range of pH. The optimal operating conditions obtained under the application of the EO-H2O2 process for the treatment of synthetic wastewater effluent were as follows: current density = 33.3 mA cm⁻²; pH = 3; and bisphenol A concentration = 0.14 mmol L^{-1} ; these optimal conditions yielded EE/O of 2.80 kWh m⁻³ order⁻¹. However, competitive levels of BPA degradation were obtained at pH 7.0 and 10.0. The evolution of inorganic species was monitored during the treatment of real wastewater under the optimized conditions mentioned above; the results obtained showed that the initial chloride species were converted to active chlorine species, and further oxidation of these species led to the formation of 1.23 mg L^{-1} ClO₂⁻, 1.82 mg L⁻¹ ClO₃⁻, and 0.98 mg L⁻¹ ClO₄⁻. The last two species are known to pose serious risks to human health and their concentration level observed in the treated effluent was far higher than the strictly low concentration levels established by the World Health Organization (70 µg L⁻¹) and the United States Environmental Protection Agency (15 µg L⁻¹) ¹). Indeed, the stringent legally established low concentration levels of these pollutants in water for human consumption make the use of the water treated here unsuitable for drinking purposes without further treatment to degrade or minimize its oxyanions content. It is worth noting, however, that the results obtained under the treatment process employed here show

that the effluents treated meet the requirements to be discharged in wastewater facilities since the concentration levels of the pollutants are far below the limit of $10 \text{ mg L}^{-1} \text{ CIO}_{4^-}$. Ammonia (NH4⁺) was not detected at the end of the treatment process. This outcome was attributed to the reaction of ammonia with the chlorinated species generated in the system and its conversion to innocuous nitrogen gas (N₂). With regard to NO₃⁻, its concentration recorded a decrease of approximately 87.5% at the end of the treatment. In fact, the reduction kinetics of the NO₃⁻ species is found to be widely affected by the type of electrodes material used. In addition, NO₃⁻ can further yield NO₂⁻ and NH4⁺. Nitrite concentration was found to be stable during the treatment process with almost no variation observed at the end of the treatment. Both NO₃⁻ and NO₂⁻ concentration levels were found to be within the maximum contamination levels allowed in drinking water.

This work highlights the challenges that different water matrices pose when it comes to the electrochemical treatment of organic pollutants aiming at obtaining potable water that is free from harmful pollutants or where the concentration levels of the compounds present in the water matrix are within the legal discharge limits allowed. Further studies need to be performed to improve the effectiveness of the EAOPs and to enhance their technology readiness level (TRL) to increase the competitiveness of these treatment processes.

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