Removal of organic pollutants from water by electro-Fenton and electro-Fenton like processes

Heng Lin

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Joint PhD degree in Environmental Science & Technology

Docteur de l’Université Paris-Est
Spécialité : Science et Technique de l’Environnement

PhD degree of Wuhan University
Speciality: Environmental Engineering

Thèse de doctorat d’université – PhD thesis

Heng LIN

Removal of organic pollutants from water by indirect electro-oxidation using hydroxyl (·OH) and sulfate (SO₄²⁻) radical species

Elimination des polluants organiques de l'eau par électrochimie indirecte basée sur les radicaux hydroxyles (·OH) and sulfate (SO₄²⁻)

To be defended on May 29, 2015

In front of the PhD committee

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Acknowledgements

I would like to thank gratefully to my supervisors Prof. Mehmet A. Oturan and Prof. Hui Zhang for giving me the chance to conduct the thesis work. It is my pleasure to work with Prof. Mehmet A. Oturan and Prof. Hui Zhang and I learned a lot of things from them.

I would also like to thank Dr. Nihal Oturan for giving me precious advice and technical supports.

I wish to express my gratitude to Prof. Ignacio Sirés (Barcelona University, Spain) and Prof. Zhihui Ai (Central China Normal University, China) for reading and evaluating my thesis. I also wish to thank Dr Nihal Oturan (Université Paris-Est, France) and Professor Feng Wu (Wuhan University, China) for their acceptance to be part of thesis jury.

I also wish to thank all my colleagues in Advanced Oxidation Lab of Wuhan University and Laboratoire Géomatériaux et Environmental of Université Paris-Est for their kindness help.

Thank for the fund support of China Scholarship Council (CSC) affiliated with the Ministry of Education of P.R. China.

Finally, I would especially like to thank my parents for always been there with encouragement.
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<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>2,4-DP</td>
<td>2-(2,4-dichlorophenoxy)-propionic acid</td>
</tr>
<tr>
<td>2,4,5-T</td>
<td>2,4,5-trichlorophenoxyacetic acid</td>
</tr>
<tr>
<td>2,4,5-TCP</td>
<td>2,4,5-trichlorophenol</td>
</tr>
<tr>
<td>4-NP</td>
<td>4-nitrophenol</td>
</tr>
<tr>
<td>AAS</td>
<td>Atomic absorption spectroscopy</td>
</tr>
<tr>
<td>ACF</td>
<td>Activated carbon fiber</td>
</tr>
<tr>
<td>AOPs</td>
<td>Advance oxidation processes</td>
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<tr>
<td>ANOVA</td>
<td>Analysis of Variance</td>
</tr>
<tr>
<td>ASP</td>
<td>Aspartame</td>
</tr>
<tr>
<td>BA</td>
<td>Benzoic acid</td>
</tr>
<tr>
<td>BBD</td>
<td>Box-Behnken design</td>
</tr>
<tr>
<td>BDD</td>
<td>Boron-doped diamond</td>
</tr>
<tr>
<td>BPA</td>
<td>Bisphenol A</td>
</tr>
<tr>
<td>CA</td>
<td>Clofibric acid</td>
</tr>
<tr>
<td>CV</td>
<td>Coefficient of variation</td>
</tr>
<tr>
<td>DSA</td>
<td>Dimensionally stable anode</td>
</tr>
<tr>
<td>DC</td>
<td>Direct current</td>
</tr>
<tr>
<td>DO</td>
<td>Dissolved oxygen</td>
</tr>
<tr>
<td>EAOPs</td>
<td>Electrochemical advanced oxidation processes</td>
</tr>
<tr>
<td>EC</td>
<td>Electrochemical</td>
</tr>
<tr>
<td>EE/O</td>
<td>Electrical energy consumption per order of magnitude</td>
</tr>
<tr>
<td>EFO</td>
<td>Electro-Fe(II)/Oxone</td>
</tr>
<tr>
<td>EI</td>
<td>Electronimpact</td>
</tr>
<tr>
<td>E.U.</td>
<td>European Union</td>
</tr>
<tr>
<td>GC–MS</td>
<td>Gas chromatography–mass spectrometry</td>
</tr>
<tr>
<td>GDEs</td>
<td>Gas diffusion electrodes</td>
</tr>
<tr>
<td>HPLC</td>
<td>High performance liquid chromatography</td>
</tr>
</tbody>
</table>
IC  Ion chromatography

$k_{\text{app}}$  Apparent rate constant values

MCE  Mineralization current efficiency

MLSS  Mixed liquor suspended solids

NDIR  Non dispersive infra-red absorption detector

OUR  Oxygen uptake rate

PDS  Peroxydisulfate

PMS  Peroxymonosulfate

PPCPs  Pharmaceuticals and personal care products

PS  Persulfate

RhB  Rhodamine B

RSM  Response surface methodology

RVC  Reticulated vitreous carbon

SAC  Saccharin

SCP  Sulfachloropyridazine

SMM  Sulfamonomethoxine

SOUR  Specific oxygen uptake rate

SUC  Sucralose

TCE  Trichloroethylene

TOC  Total organic carbon

$t_{\text{R}}$  Retention time

UV  Ultraviolet

XPS  X-ray photoelectron spectroscopy

XRD  X-ray diffraction
Abstract

In recent years, more and more refractory and toxic organic contaminants are detected in wastewater, surface water and ground water. Many of these organic pollutants can hardly be degraded by conventional water treatments. Advanced oxidation processes (AOPs), which based on in situ generation of strong oxidants, mainly the generation of hydroxyl radicals (·OH), have been applied to treat various non-biodegradable organic compounds in water. Conventional Fenton technology is a promising AOP for the treatment of organic contaminants in water. In Fenton process, hydroxyl radical, which is the second strongest oxidant (after fluorine), is formed through Fenton’s reaction and then degrade organic pollutants. Similar to conventional Fenton process, transition metal ions (Fe$^{2+}$, Co$^{2+}$, Ag$^+$, etc.) can also activate persulfate (PS) and generate sulfate radicals (SO$_4$$^-$). Sulfate radical is a powerful oxidant and can oxidize most of organic pollutants. This process is named Fenton-like process. There are some disadvantages existed in conventional Fenton and Fenton-like process. For example, a high concentration of Fe$^{2+}$ is required and a large amount of iron sludge is generated. In order to solve these problems, electro-Fenton and sulfate radical-based electro-Fenton-like processes are developed. Fe$^{2+}$ can be regenerated via cathodic reduction in electro-Fenton and electro-Fenton-like processes. Therefore, the Fe$^{2+}$ concentration used in these processes is much lower than that in Fenton and Fenton-like processes. In this paper, electro-Fenton and sulfate radical-based electro-Fenton-like processes were used to degrade artificial sweeteners and azo dyes. The removal efficiency, the oxidation mechanism, degradation pathway and toxicity evolution of target pollutants were investigated.

(1) A detailed discussion on the oxidative degradation of artificial sweetener aspartame (ASP) in acidic aqueous solution containing catalytic amount of Fe$^{2+}$ by using electro-Fenton process is reported. In electro-Fenton process, ASP could be completely removed in a 30 min reaction and the removal of ASP followed pseudo-first-order kinetics. The increase of Fe$^{2+}$ concentration and applied current to certain extent could increase the removal efficiency of ASP, while further increasing the Fe$^{2+}$ concentration and applied current could lead to the decrease of removal efficiency. Absolute rate constant of
hydroxylation reaction of ASP was determined as \((5.23 \pm 0.02) \times 10^9 \text{ M}^{-1} \text{ s}^{-1}\). When boron-doped diamond (BDD) electrode was used as anode, ASP could be completely mineralized in a 360 min treatment. Short-chain aliphatic acids such as oxalic, oxamic and maleic acid were identified as aliphatic intermediates in the electro-Fenton process. The bacteria luminescence inhibition showed the toxicity of ASP solution increased at the beginning of electrolysis, and then it declined until lower than the untreated ASP solution at the end of the reaction.

(2) The removal of artificial sweetener saccharin (SAC) in aqueous solution by electro-Fenton processes was performed. Experiments were carried out in an undivided cylindrical glass cell with a carbon-felt cathode and a DSA, Pt or boron-doped diamond (BDD) anode. The removal of SAC by electrochemically generated hydroxyl radicals followed pseudo-first order kinetics with all the anodes. The absolute rate constant of the SAC hydroxylation reaction was found as \((1.85 \pm 0.01) \times 10^9 \text{ M}^{-1} \text{ s}^{-1}\), which was determined by using the competition kinetic method. The comparative study of TOC removal efficiency during electro-Fenton treatment indicated a higher mineralization rate with BDD than Pt anode. Formation of oxalic, formic, and maleic acids were observed during electro-Fenton process. The evolution of toxicity of SAC and/or its reaction by-products based on the \(V. fischeri\) bacteria luminescence inhibition was studied.

(3) The removal of artificial sweetener sucralose (SUC) in aqueous solution by electro-Fenton processes was also performed in this study. Hydroxyl radical (’OH), a highly powerful oxidizing agent, was generated catalytically via the electrochemically assisted Fenton’s reaction in the bulk and water oxidation on the surface of anode. The effect of \(\text{Fe}^{2+}\) concentration and applied current on the mineralization of SUC was evaluated. A higher mineralization rate was obtained with BDD than that of Pt anode. Mineralization current efficiency (MCE) was calculated and better mineralization current efficiency was achieved at relatively short electrolysis time and low applied current. The concentration of the formed carboxylic acids and released inorganic ion was monitored by ion-exclusion chromatography and ion chromatography (IC). The toxicity of SUC and/or its reaction by-products was investigated based on the \(V. fischeri\) bacteria luminescence inhibition.
(4) The removal of Orange II by a novel electro/α-FeOOH/peroxydisulfate process is reported in this study. In electro/α-FeOOH/peroxydisulfate process, sulfate radicals were generated by activating peroxydisulfate (PDS) with goethite (α-FeOOH). When combined with electrochemical process, the Fe(III) on the surface of α-FeOOH converts to Fe(II) by cathodic reduction. The effect of initial pH on the decolorization of Orange II was investigated. Response surface methodology (RSM) based on Box-Behnken statistical experiment design (BBD) was applied to analyze the experimental variables. The positive and negative effects on the decolorization of Orange II were determined. The response surface methodology models were derived based on the results of the pseudo-first-order decolorization rate constant and the response surface plots were developed accordingly. The results indicated that the applied current showed a positive effect on the decolorization rate constant of Orange II. The interaction of α-FeOOH dosage and PDS concentration was significant. The ANOVA results confirmed that the proposed models were accurate and reliable for the analysis of the variables of EC/α-FeOOH/PDS process.

(5) The decolorization of Orange II in aqueous solution by magnetite (Fe₃O₄) activated peroxydisulfate (PDS) oxidation in an electrochemical reactor (EC/Fe₃O₄/PDS process) was performed in this study. Various parameters were investigated to optimize the process, including initial pH, current density, PDS concentration and Fe₃O₄ dosage. The stability of Fe₃O₄ particles was observed by recycle experiments. The X-ray photoelectron spectroscopy (XPS) was applied to investigate the surface properties of Fe₃O₄ before and after reaction. GC–MS analysis was employed to identify the intermediate products and a plausible degradation pathway of Orange II was proposed. The change of acute toxicity during the treatment was investigated by activated sludge inhibition test. The TOC removal efficiency was 30.0% in a 90 min treatment.

**Keywords:** Electro-Fenton; Electro-Fenton-like process; Hydroxyl radicals; Sulfate radicals; Organic pollutants
Chapter 1 Introduction
1.1 Background

Water quality and availability is a challenging problem facing our society all over the world (Rivas et al. 2009; Bernal-Martínez et al. 2010). In recent years, more and more organic contaminants from contaminated soil, agricultural runoff, industrial wastewater and hazardous compounds storage leakage are detected in wastewater, surface water and ground water. The presence of these organic compounds in water poses serious threat to public health since most of them are toxic, endocrine disrupting, mutagenic or potentially carcinogenic to humans, animals and aquatic life in general (Babuponnusami and Muthukumar 2014). Many organic pollutants are considered as toxic and detrimental even when present at very low concentrations. Moreover, many organic contaminants cannot be eliminated by conventional physical separation methods or cannot be degraded by biological processes due to the recalcitrant nature of the pollutants present (Pérez et al. 2002). Therefore, more powerful wastewater treatment methods are required.

Advance oxidation processes (AOPs) have been proved to be effective for the degradation of many toxic/persistent organic contaminants from aqueous medium such as coloring matters, pesticides, artificial sweeteners and pharmaceuticals and personal care products (Kim and Tanaka 2009; Wang et al. 2012; Calza et al. 2013; Janin et al. 2013; El-Ghenemy et al. 2014; Rodrigo et al. 2014; Wang et al. 2014). Moreover, AOPs have been successfully used as pretreatment methods to reduce the concentration of toxic organic compounds that recalcitrant to biological wastewater treatments (Stasinakis 2008). AOPs are based on the in-situ generation of hydroxyl radical (•OH, \( E^0(\text{•OH}/\text{H}_2\text{O}) = 2.80 \text{ V/SHE} \)) (Evgenidou et al. 2007; Salazar et al. 2013; Oturan and Aaron, 2014), which is the second strongest oxidizing agent after fluorine (Mousset et al. 2014a). AOPs are especially efficient for degrading aromatic molecules due to the electrophilic aromatic substitution of hydroxyl radical which then lead to the opening of the aromatic ring (Brillas et al. 2009; Mousset et al. 2014b). The most commonly used AOPs for the removal of organic pollutants from water is based on the Fenton’s reagent (an aqueous mixture of \( \text{Fe}^{2+} \) and \( \text{H}_2\text{O}_2 \) that produces hydroxyl radicals)
(Babuponnuam and Muthukumar 2012; Basturk and Karatas 2014). It has been used as an attractive and effective technology for the degradation of various organic pollutants (Hermosilla et al. 2009a; Hermosilla et al. 2009b; Zazo et al. 2009; Hermosilla et al. 2012) because of the lack of toxicity of the reagents, eventually leaving no residues and the simplicity of the technology (Sun et al. 2007; Nidheesh and Gandhimathi 2012). However, the conventional Fenton process has the disadvantages of requirement of high Fe$^{2+}$ concentration and the formation of Fe sludge which limit its application (Nidheesh and Gandhimathi 2012).

Electro-Fenton process, in which H$_2$O$_2$ is produced electrochemically and Fe$^{2+}$ can be regenerated throughout the process, overcomes these disadvantages of conventional Fenton process. Compared to the conventional Fenton process, the electro-Fenton process has the advantage of regeneration of Fe$^{2+}$ and avoiding the storing and transport of the H$_2$O$_2$. Moreover, electricity as a clean energy source is used in the process, so the overall process does not create secondary pollutants (Jiang and Zhang 2007; Brillas et al. 2009). The electro-Fenton process has been successfully applied in wastewater treatment and the process efficiency has been experimentally confirmed by different authors for the treatment of media containing pesticides (Oturan et al. 2009; Yatmaz and Uzman 2009; Abdessalem et al. 2010), herbicides (Kaichouh et al. 2004; Da Pozzo et al. 2005; García et al. 2014), synthetic dyes (Cruz-González et al. 2010; Ghoneim et al. 2011; Kourdali et al. 2014), pharmaceuticals (El-Ghenemy et al. 2013; Feng et al. 2014; Yahya et al. 2014), and so on.

As a strong oxidant, persulfate (PS) is another popular reagent in the wastewater treatment because it is more stable than H$_2$O$_2$ and it can be transported over a long distance without alteration. Similar to the conventional Fenton process, PS can be activated by transition metals to generate sulfate radicals ($\text{SO}_4^{\cdot-}$), which have a high standard redox potential ($E^0 = 2.6$ V/SHE) (Shukla et al. 2010; Deng and Ezyske 2011) and can oxidize organic compounds into small molecules and carbon dioxide (Gayathri et al. 2010). This process is also named Fenton-like process (Fernandez et al. 2003; Fernandez et al. 2004).

Like conventional Fenton process, Fe$^{2+}$ is hard to be regenerated after conversion
to Fe$^{3+}$ in sulfate radical-based Fenton-like process, thus high concentration of Fe$^{2+}$ is required and a large amount of iron sludge is produced. In order to solve this problem, electrochemical (EC) technology was combined with Fenton-like process (electro-Fenton-like process) and used to treat herbicide 2,4,5-trichlorophenoxyacetic acid (2,4,5-T) (Wang and Chu 2011), azo dye Orange II (Wu et al. 2012), bisphenol A (BPA) (Lin et al. 2013), clofibric acid (CA) (Lin et al. 2014) and landfill leachate (Zhang et al. 2014). However, the electro-Fenton-like processes mentioned above are all homogeneous oxidation processes. The removal of metal ions from the effluent requires additional separation and disposal (Zhong et al. 2011). Therefore, various kinds of solid catalysts instead of soluble metal ions have received much attention. Compared with soluble metal ions, solid catalyst can be recovered and reused without being lost along with the effluent. As a result, different kinds of iron metal oxides (such as magnetite and goethite) are employed in the electro-Fenton-like process in this study.

In this work, electro-Fenton and electro-Fenton-like processes in the presence of different oxidants and catalysts were studied. As an emerging contaminant, artificial sweeteners are an essential part of food technology as a group of discrete chemical substances that possess intense sweetness. Synthetic dyes are one of the most common toxic pollutants in the natural environment around the world. These two types of organic compounds were selected as model pollutants in this thesis work.

1.2 Research goals and objective

The objective of this work was to study the removal of target pollutants from water using electro-Fenton and electro-Fenton-like processes. The optimal oxidation conditions were identified and the oxidation mechanisms were examined. To meet these goals, the following specific objectives were defined:

1. Conduct experiments to investigate the effect of the operation parameters on the degradation and mineralization of artificial sweetener aspartame (ASP) by electro-Fenton process. The concentration of inorganic ions and carboxylic acid released during the treatment was monitored.
2. Conduct experiments to investigate the degradation and mineralization of artificial sweetener saccharin (SAC) using electro-Fenton process. The comparison among different anode materials on the degradation and mineralization was investigated. Moreover, the toxicity of SAC and its byproducts during electro-Fenton process was determined.

3. Conduct experiment to investigate the mineralization of artificial sweetener sucralose (SUC) using electro-Fenton process. The influence of the operation parameters on the mineralization of SUC was investigated. The Mineralization current efficiency (MCE) was calculated to evaluate the mineralization capacity.

4. Fe$^{2+}$ was replaced by goethite to activate PS and degraded azo dye Orange II in sulfate radical-based electro-Fenton-like process. The response surface methodology models were derived based on the results of the pseudo-first-order decolorization rate constant and the response surface plots were developed accordingly.

5. Magnetite was used as heterogeneous catalyst to degrade Orange II in sulfate radical-based electro-Fenton-like process. The surface properties of Fe$_3$O$_4$ before and after reaction were investigated. The intermediates of Orange II during the treatment were identified and a plausible degradation pathway was proposed. The change of acute toxicity during the treatment was investigated.

1.3 Structure of the thesis

This dissertation is composed of eight chapters.

Chapter 1 is an introduction, including background information of electro-Fenton and electro-Fenton-like processes and the chosen organic pollutants, research objectives and organization of this thesis.

Chapter 2 presents a review of recent work related to this study. Four parts are described:

2.1 Artificial sweeteners
2.2 Synthetic dyes
2.3 Advanced oxidation processes (AOPs)
2.4 Iron metal oxides application in AOPs

The following three chapters are the presentation of experiments results related to electro-Fenton process.

Chapter 3 is entitled “Treatment of aspartame in aqueous solution by electro-Fenton process”. The removal of ASP in electro-Fenton process was investigated. The effect of Fe$^{2+}$ concentration and applied current on the degradation and mineralization of ASP was evaluated. The absolute rate constant for the reaction between ASP and 'OH was determined by using the competition kinetic method. The concentration of the formed carboxylic acids and released inorganic ion was monitored by ion-exclusion chromatography and ion chromatography (IC). The toxicity of ASP and/or its reaction byproducts was studied based on the *V. fischeri* bacteria luminescence inhibition.

Chapter 4 is entitled “Treatment of saccharin (SAC) in aqueous solution by electro-Fenton process”. The removal of artificial sweeteners saccharin (SAC) in aqueous solution by electro-Fenton processes was performed in this study. Different anode materials on the removal and mineralization of SAC were tested. The absolute rate constant of the SAC hydroxylation reaction was determined. The evolution of carboxylic acids was determined by ion-exchange chromatography. The evolution of toxicity of SAC and/or its reaction byproducts based on the *V. fischeri* bacteria luminescence inhibition was studied.

Chapter 5 is entitled “Treatment of sucralose (SUC) in aqueous solution by electro-Fenton process”. The removal of artificial sweeteners SUC in aqueous solution by electro-Fenton processes was carried out in this study. The effect of Fe$^{2+}$ concentration and applied current on the mineralization of SUC was evaluated. The concentration of the formed carboxylic acids and released inorganic ion was monitored by ion-exchange chromatography and ion chromatography. The toxicity of SUC and/or its reaction byproducts was investigated based on the *V. fischeri* bacteria luminescence inhibition.

The following two chapters are the parts devoted to the experiments results of electro-Fenton-like process.

Chapter 6 is entitled “Decolorization of Orange II in water by
electro/$\alpha$-FeOOH/PDS process”. In this section, the removal of Orange II by electro/$\alpha$-FeOOH/peroxydisulfate process was investigated. Response surface methodology (RSM) based on Box-Behnken statistical experiment design (BBD) was applied to analyze the experimental variables. The positive and negative effects on the decolorization of Orange II were determined. The response surface methodology models were derived based on the results of the pseudo-first-order decolorization rate constant and the response surface plots were developed accordingly.

Chapter 7 is entitled “Decolorization of Orange II in water by electro/Fe$_3$O$_4$/PDS process”. In this chapter, the decolorization of Orange II was performed in aqueous solution by Fe$_3$O$_4$ activated per oxydisulfate (PDS) oxidation in an electrochemical reactor (EC/Fe$_3$O$_4$/PDS process) was performed. Various parameters were investigated to optimize the process. The surface properties of Fe$_3$O$_4$ before and after reaction was investigate. The intermediates of Orange II during the treatment were identified and the degradation pathway was proposed. The change of acute toxicity during the treatment was investigated.

Finally, the thesis manuscript ends with Chapter 8 entitled "General conclusions and perspectives”.

References


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Chapter 2 Literature review
2.1 Artificial sweeteners

Artificial sweeteners are one of the most important and interesting classes of emerging contaminants. Artificial sweeteners have been classified as nutritive and non-nutritive depending on whether they are a source of calories (Whitehouse et al. 2008). The nutritive sweeteners include the monosaccharide polyols (e.g. sorbitol, mannitol, and xylitol) and the disaccharide polyols (e.g. maltitol and lactitol). They are approximately equivalent to sucrose in sweetness (Dills Jr 1989). The non-nutritive sweeteners, better known as artificial sweeteners, include substances from several different chemical classes that interact with taste receptors and typically exceed the sweetness of sucrose by a factor of 30 to 13,000 times (Whitehouse et al. 2008). The most popular artificial sweeteners are aspartame (ASP), saccharin (SAC), sucralose (SUC) and acesulfame. The chemical structure and main characteristics of these commonly used artificial sweeteners were presented in Table 2-1.

Artificial sweeteners are water contaminants that are highly specific to wastewater. Different from other emerging trace contaminants, such as pharmaceuticals and personal care products (PPCPs), artificial sweeteners have been considered in environmental sciences only recently (Loos et al. 2009; Mead et al. 2009; Richardson 2010; Richardson and Ternes 2011; Lange et al. 2012). Excretion after human consumption is one of the major source of artificial sweeteners in the environment (Kokotou et al. 2012). The artificial sweeteners can also enter into wastewater treatment plants from households and industries and they eventually reside in the receiving environmental bodies from effluents (Houtman 2010).
Table 2-1 The chemical structure and main characteristics of some commonly used artificial sweeteners

<table>
<thead>
<tr>
<th>Name</th>
<th>Chemical structure</th>
<th>Molecular formula</th>
<th>Molecular weight (g/mol)</th>
<th>CAS number</th>
<th>Water solubility (g/L)</th>
<th>Number of times sweeter than sucrose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspartame</td>
<td><img src="image1" alt="Aspartame" /></td>
<td>C_{12}H_{18}N_{2}O_{5}</td>
<td>394.31</td>
<td>22839-47-0</td>
<td>~10 (25 °C)</td>
<td>200</td>
</tr>
<tr>
<td>Saccharin</td>
<td><img src="image2" alt="Saccharin" /></td>
<td>C_{7}H_{5}NO_{3}S</td>
<td>183.18</td>
<td>81-07-2</td>
<td>4</td>
<td>200-700</td>
</tr>
<tr>
<td>Sucralose</td>
<td><img src="image3" alt="Sucralose" /></td>
<td>C_{12}H_{19}Cl_{3}O_{8}</td>
<td>397.63</td>
<td>56038-13-2</td>
<td>283 (20 °C)</td>
<td>600</td>
</tr>
<tr>
<td>Acesulfame</td>
<td><img src="image4" alt="Acesulfame" /></td>
<td>C_{4}H_{5}NO_{4}S</td>
<td>163.15</td>
<td>33665-90-6</td>
<td>270 (20 °C)</td>
<td>200</td>
</tr>
</tbody>
</table>
Artificial sweeteners are extensively tested for potential adverse health effects on humans because they are used as food additives (Schiffman and Gatlin 1993; Kroger et al. 2006; En Humanos and Revisión 2009). Although the measured concentrations of some artificial sweeteners range up to microgram per liter levels in surface water, groundwater and drinking water, there is a huge safety margin regarding potential adverse health effects (Lange et al. 2012). Acceptable daily intake value of artificial sweeteners is 5 mg/kg of body weight per day and are thus three to four orders of magnitude above the maximum possible daily human intake by drinking water. Adverse human health effects for the application of artificial sweeteners have been reported in several studies (Lange et al. 2012; Toth et al. 2012). The long-term health effects resulting from the chronic exposure to low levels of these compounds are largely unknown (Mawhinney et al. 2011).

2.1.1 Aspartame

Aspartame, the best known of the various artificial sweeteners, is a dipeptide methyl ester (Table 1), namely N-L-α-aspartyl-L-phenylalanine methyl ester. It was discovered by J. Schlatter in 1965 and commercialized by Searle & Co. under the brand name Nutrasweet™ (Galletti et al. 1995). As the “first generation” sweeteners (Berset and Ochsenbein 2012), ASP is a white dipeptide, crystalline, low-caloric sweetener, which is 180-200 times sweeter than sucrose (Conceição et al. 2005). ASP is relatively stable in its dry form. When pH below 3.0, it is unstable and is hydrolyzed to aspartylphenylalanine. Above pH 6.0, it is transformed to 5-benzyl-3,6-dioxo-2-piperazine acetic acid (Sardesai and Waldshan 1991; Kokotou et al. 2012). ASP was found in all of the surface waters at a concentration up to 0.21 mg/L in Tianjin, China (Gan et al. 2013). The risks of ASP ingestion would be in the toxicity of its metabolism products. One of its metabolites is methanol which can cause acidoses and blindness (Conceição et al. 2005). ASP was rarely degraded by AOPs and it was treated by electro-Fenton process for the first time in this study.
2.1.2 Saccharin

Saccharin (SAC), one of the “first generation” artificial sweeteners, was discovered accidentally by Fahlberg and Remsen at Johns Hopkins University in 1879 when they were studying the oxidation of o-toluene-sulfonamides (Bruno et al. 2014). SAC is about 200-700 times sweeter than sucrose (Assumpção et al. 2008; Lange et al. 2012). Nowadays SAC is approved in more than 90 countries and is widely applied in many pharmaceutical and dietary products (Filho et al. 2003). For example, in the European Union (E.U.) SAC is used as an additive in animal feed for piglets, pigs, bovines and calves, and it is also the major degradation by-product of certain sulfonylurea herbicides (Buerge et al. 2010; Kokotou et al. 2012). SAC was slowly absorbed and not metabolized by the human organism, so it is consequently an appropriate artificial sweetener for diabetics (Filho et al. 2003). SAC has been detected in municipal wastewaters and in rivers in many countries, such as Germany, Switzerland and China (Gan et al. 2013; Van Stempvoort et al. 2011). For example, SAC was detected in surface waters and ranged from 50 ng/L to 0.21 mg/L in Tianjin, China (Gan et al. 2013). It is crucial to find efficient water treat technologies for the removal of SAC from aqueous media. In this work, electro-Fenton process was employed to degrade SAC in water.

2.1.3 Sucralose

Sucralose (SUC), which is a chlorinated disaccharide derived from sucrose, is one of the most popular artificial sweeteners and has shown increasing trend of consumption (Toth et al. 2012). SUC is about 600 times sweeter than the parent molecule, sucrose (Grotz and Munro 2009; Sharma et al. 2014). Nowadays, SUC is approved in more than 80 countries and is used widely in food products and pharmaceuticals (Sharma et al. 2014). SUC is not metabolized by the human body due to the orientation of the glycosidic linkage, and consequently, up to 92% of the consumed SUC is excreted unchanged in both urine and feces (Sharma et al. 2014; Toth et al. 2012). SUC assesses potential adverse effects on health, since it provokes symptoms, such as, increase in
blindness, mineralization of pelvic area and epithelial hyperplasia (Calza et al. 2013). Nowadays, SUC has been detected in wastewaters and rivers in many countries, such as America, France, Italy and China (Calza et al. 2013; Gan et al. 2013). Recently, SUC is considered as an emerging contaminant the by Environmental Protection Agency (EPA) due to its occurrence in environmental waters and persistence (half-life up to several years) (Richardson 2011; Richardson and Ternes 2011; Calza et al. 2013). Unfortunately, SUC is hard to degrade by conventional wastewater treatments processes (Labare and Alexander 1994; Torres et al. 2011; Sharma et al. 2014). It is crucial to find more powerful water treat technologies for the removal of SUC from polluted effluents.

Different AOPs have been employed to remove SUC from water, including ozonation (Scheurer et al. 2010; Soh et al. 2011), ultraviolet (UV) irradiation (Soh et al. 2011), UV/H₂O₂ photooxidation (Keen and Linden 2013), photo-Fenton process (Calza et al. 2013) and TiO₂ photocatalysis process (Calza et al. 2013). When treated by 100 μM ozone, 6% SUC remained after 60 min (Soh et al. 2011). Since SUC does not have any evident sites for direct oxidation by ozone, the removal of SUC was mainly caused by hydroxyl radicals generated in the oxidative system (Soh et al. 2011; Sharma et al. 2014). In a photochemical reactor, SUC did not degrade after 5 h UV exposure (Soh et al. 2011) and it also did not decay significantly when UV light irradiation lasted for 24 h (Torres et al. 2011), because SUC has low molar extinction coefficient in the UV region (Sharma et al. 2014). When H₂O₂ was added to UV irradiation system, the removal of SUC was accelerated and 500 μg/L SUC could be almost totally removed when UV fluence was 4000 mJ/cm² (Keen and Linden 2013). When photo-Fenton and TiO₂ photocatalysis processes were introduced to treat SUC, the removal efficiency of 15 mg/L SUC were 97.1% and 88.9%, respectively (Calza et al. 2013). SUC was removed from aqueous media by electro-Fenton process for the first time in this study.

2.2 Synthetic dyes

Nowadays, there has been an increasing use of synthetic dyes in a large amount of industrial areas such as the textile, leather goods, pharmaceutical industry, food industry
and other chemical usages (Forgacs et al. 2004; Hai et al. 2007; Soon and Hameed 2011). In 1856, the world’s first commercially synthetic dye was discovered accidentally by William Henry Perkin (Saratale et al. 2011). Such dyes are defined as colored substances which can give fibers a permanent color. This color is able to resist fading upon exposure to sweat, light, water and many chemicals, including oxidizing agents and microbial attack (Rai et al. 2005). More than ten thousand commercially available dyes were developed and used in manufacturing by the end of the 19th century (Robinson et al. 2001). The growth of the worldwide textile industry has been accompanied by a rise in pollution due to wastewater contaminated with dyestuff (Parikh and Madamwar 2005). The water consumption and the wastewater generation from the textile industry (dry processing mill and woven fabric finishing mills) depend on the processing operations employed during the conversion of fiber to textile fabric (Dhanve et al. 2008). The textile industry is one of the greatest generators of liquid effluent pollutants which attributed to the high quantities of water used in the dyeing processes. Moreover, the processing stages and types of synthetic dyes applied during this conversion determine the variable wastewater characteristics in terms of pH, dissolved oxygen, organic and inorganic chemical content (Banat et al. 1996). It is estimated that over $7 \times 10^5$ tons of textile dyes are discharged in such industrial effluents every year (Robinson et al. 2001; Brillas and Martínez-Huitle 2015).

Dyes can be classified as Acid, Basic, Direct, Vat, Sulfur, Reactive, Disperse, metal complexes, etc., including antraquinone, indigoide, triphenylmethyl, xanthene and phthalocyanine derivatives (Martínez-Huitle and Brillas 2009; Brillas and Martínez-Huitle 2015). Among these dyes, azo dyes, which are characterized by one or two azo-bonds (–N=N–), account for over 70% of all dyestuffs used worldwide (Salazar et al. 2012, Saratale et al. 2011; Brillas and Martínez-Huitle 2015), making them the largest group of synthetic colorants and the most common synthetic dyes released into the environment (Chang et al. 2001; Zhao and Hardin 2007; Saratale et al. 2009). Inappropriate disposal of these dyes contaminated waters consistently would result in an even widely contaminated range of environmental matrixes, including surface, ground, drinking water, and soils. Unfortunately, due to the characteristic of biological
resistance and chemical stability, azo dyes can hardly be treated using classical wastewater treatments (Ganesh et al. 1994; Razo-Flores et al. 1997). For this reason, many studies have been focused mainly on the removal of azo dyes from waters (Shu and Chang 2005; Lucas and Peres 2006; Habibi and Talebian 2007; Migliorini et al. 2011; Yu et al. 2013).

### 2.3 Advanced oxidation processes (AOPs) in dye removal

Advanced oxidation processes (AOPs) have been precisely defined as “near ambient temperature and pressure water treatment processes which involve the generation of a powerful oxidizing agent such as hydroxyl radical in solution in sufficient quantity to effective water purification” by Glaze et al. at 1987. AOPs mainly use chemical, photochemical or electrochemical techniques to generate hydroxyl radicals (‘OH) which are a highly powerful oxidizing agents ($E^0 = 2.80$ V/SHE) and can bring about chemical degradation of organic pollutants (Zhang et al. 2007; Mollah et al. 2010). ‘OH is a non-selective and efficient oxidizing agent. It readily attacks a large number of organic chemicals and converts them to less complex and less harmful intermediate products. At sufficient contact time and proper operation conditions, it is practically possible to mineralize the target pollutant to CO$_2$ and H$_2$O, which are the most stable end-product of chemical oxidation (Ayoub et al. 2010). The advantage of AOPs overall chemical and biological processes is that they are “environmental-friendly” as they neither transfer pollutants from phase to the other (as in chemical precipitation, adsorption, and volatilization) nor produce massive amounts of hazardous sludge (Ince and Apikyan 2000). AOPs are considered as the promising processes for the wastewater treatment in 21$^{st}$ century (Munter 2001).

#### 2.3.1 Fenton process

The Fenton’s chemistry started as early as the end of the nineteenth century, when Henry J. Fenton reported that H$_2$O$_2$ could be activated by Fe(II) salts to oxidize tartaric acid (Fenton 1894). In a century since then, Fenton and related reactions have become
of great interest for their relevance to biological chemistry, synthesis and the chemistry of natural waters (Pignatello et al. 2006). In the 1930s, Haber and Weiss showed that the catalytic decomposition of H₂O₂ by iron salts obeyed to a complex radical and chain mechanism (Haber and Weiss 1934). Since then, interests in Fenton process has been definitely renewed, and Fenton’s reagent was applied to oxidize toxic organics originally appeared in the mid-1960s (Brillas et al. 2009).

The generally accepted mechanism of the Fenton process proposes that hydroxyl radicals are produced in accordance with Eq. (2-1) (Bautista et al. 2008; Oturan and Aaron 2014), while the catalyst is regenerated through Eq. (2-2), or from the reaction of Fe³⁺ with intermediate organic radicals (Eqs. (2-3)–(2-4)) (Walling 1975; Pignatello et al. 2006; Bautista et al. 2008). However, ferrous ions are consumed more rapidly than they are regenerated (Nidheesh and Gandhimathi 2012), and then high concentration of Fe²⁺ is required in Fenton process.

$$\text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{OH}^- + \cdot\text{OH} \quad k_1 = 76 \text{ M}^{-1} \text{s}^{-1} \quad (2-1)$$

$$\text{Fe}^{3+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{2+} + \text{HO}_2\cdot + \text{H}^+ \quad k_2 = 0.001-0.01 \text{ M}^{-1} \text{s}^{-1} \quad (2-2)$$

$$\text{RH} + \cdot\text{OH} \rightarrow \text{R}^- + \text{H}_2\text{O} \quad (2-3)$$

$$\text{R}^- + \text{Fe}^{3+} \rightarrow \text{R}^+ + \text{Fe}^{2+} \quad (2-4)$$

$$\text{R}^+ + \text{OH}^- \rightarrow \text{R}^- + \text{OH} \quad (2-5)$$

Nevertheless, a number of competitive reactions can also occur (Eqs (2-6)–(2-10)), which negatively affect the oxidation process (Bautista et al. 2008). The HO₂⁻ radical formed in Eq. (2-7) is characterized by a lower oxidization power and, therefore, is signifi cantly less reactive towards organic compounds (Rush and Bielski 1985; Trapido et al. 2009; Oturan and Aaron 2014).

$$\text{Fe}^{2+} + \cdot\text{OH} \rightarrow \text{Fe}^{3+} + \text{OH}^- \quad k_3 = 3.20 \times 10^8 \text{ M}^{-1} \text{s}^{-1} \quad (2-6)$$

$$\text{H}_2\text{O}_2 + \cdot\text{OH} \rightarrow \text{HO}_2\cdot + \text{H}_2\text{O} \quad k_4 = 2.70 \times 10^7 \text{ M}^{-1} \text{s}^{-1} \quad (2-7)$$

$$\text{Fe}^{2+} + \text{HO}_2\cdot \rightarrow \text{Fe}^{3+} + \text{OH}^- \quad k_5 = 1.30 \times 10^6 \text{ M}^{-1} \text{s}^{-1} \quad (2-8)$$

$$\text{Fe}^{3+} + \text{HO}_2\cdot \rightarrow \text{Fe}^{2+} + \text{O}_2 + \text{H}^+ \quad k_6 = 1.20 \times 10^6 \text{ M}^{-1} \text{s}^{-1} \quad (2-9)$$

$$\cdot\text{OH} + \cdot\text{OH} \rightarrow \text{H}_2\text{O}_2 \quad k_7 = 5.20 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \quad (2-10)$$

Basically, the Fenton process possesses some important advantages for wastewater
treatment, such as easy implementation in existing plants, easy-to-handle and relatively inexpensive chemicals, no need for energy input, and so on (Bautista et al. 2008; Oturan and Aaron 2014). It has been employed to degrade various kinds of organic pollutants in water, for example, dyes (Muruganandham and Swaminathan 2004; Sun et al. 2009), landfill leachate (Deng and Englehardt 2006; Hermosilla et al. 2009) and phenol (Kavitha and Palanivelu 2004; Zazo et al. 2005).

However, Fenton process has some disadvantages which limit its application. Firstly, high concentration of Fe$^{2+}$ is required and large amount of Fe sludge is formed (Chou et al. 1999). The treatment of the sludge-containing Fe ions at the end of the wastewater treatment is expensive and needs large amount of chemicals and manpower (Nidheesh and Gandhimathi 2012). Secondly, Fenton’s reaction is limited by a narrow pH range (pH 2.5–3.5) because iron ions will be precipitated at higher pH values (Nidheesh and Gandhimathi 2012). Thirdly, as a liquid solution, the storing and transport of H$_2$O$_2$ are difficult. In order to solve these problems, Electrochemical advanced oxidation processes (EAOPs) based on Fenton’s reaction was developed.

2.3.2 Electrochemical Advanced Oxidation Processes (EAOPs) based on Fenton’s reaction

2.3.2.1 Fundamentals and categories

The electrochemical (EC) technology which based on the transfer of electrons, have received great attention for the prevention of pollution problems (Brillas et al. 2009; Oturan and Aaron 2014). The main advantage of EC technology is its environmental compatibility since the main reagent, electron, is a clean reagent. A large variety of EAOPs were developed by combining EC technology with AOPs in the last decade due to their environmental safety and compatibility (operating at mild conditions), versatility, high efficiency, and possibility of automation (Oturan and Aaron 2014). Among all the EAOPs, EAOPs based on Fenton’s reaction chemistry are
eco-friendly methods and avoid some disadvantages in conventional Fenton process that have recently attracted great attention for water remediation (Nidheesh and Gandhimathi 2012). EAOPs based on Fenton’s reaction chemistry can be generally divided into three categories.

The first one is electro-Fenton process. \( \text{H}_2\text{O}_2 \) is formed by the reduction of the dissolved oxygen on the cathode surface (Eq. (2-11)) in an electrolytic cell. \( \text{H}_2\text{O}_2 \) can then react with the externally added catalyst (\( \text{Fe}^{2+} \) or \( \text{Fe}^{3+} \)) to produce ‘OH (Eq. (2-1)). Moreover, \( \text{Fe}^{2+} \) which consumed in Fenton’s reaction can be regenerated by cathodic reduction (Eq. (2-12)) in electro-Fenton process which reduces the required concentration of initial \( \text{Fe}^{2+} \) to an catalytic amount (Figure 2-1a).

\[
\begin{align*}
\text{O}_2 + 2\text{H}^+ + 2\text{e}^- & \rightarrow \text{H}_2\text{O}_2 & (2-11) \\
\text{Fe}^{3+} + \text{e}^- & \rightarrow \text{Fe}^{2+} & (2-12)
\end{align*}
\]

Electro-Fenton process solves some problems existed in Fenton process. For example, higher degradation rate of organic pollutants is obtained because of the continuous regeneration of \( \text{Fe}^{2+} \) at the cathode and the on-site production of \( \text{H}_2\text{O}_2 \) avoids its dangerous transport and storage (Martínez-Huitle and Brillas 2009) and avoiding also the competition of wasting reactions (2-6), (2-7) and (2-10) (Oturan and Aaron 2014, Sirés et al. 2014).

The reporting cathode materials favored the electrogeneration of \( \text{H}_2\text{O}_2 \) were gas diffusion electrodes (GDEs) (Brillas et al. 1998; Panizza and Cerisola 2009; Barros et al. 2014; Yu et al. 2015), graphite (Yuan and Lu 2005; Yuan et al. 2006) and three-dimensional electrodes such as carbon-felt (Pimentel et al. 2008; Diagne et al. 2014; Olvera-Vargas et al. 2014), activated carbon fiber (ACF) (Wang et al. 2005; Lei et al. 2010; Wang et al. 2010a), reticulated vitreous carbon (RVC) (Xie and Li 2006; Martínez and Bahena 2009) and carbon sponge (ÖZcan et al. 2008). Among these cathode materials, carbon-felt has a high specific surface that favors the fast generation of Fenton’s reagent (\( \text{H}_2\text{O}_2 \) and \( \text{Fe}^{2+} \)) and has applied widely (Brillas et al. 2009). The commonly used anode materials in electro-Fenton process were high oxygen overvoltage anodes (M), such as \( \text{PbO}_2 \) dimensionally stable anodes (DSA), Pt and boron-doped diamond (BDD) anode. Organic pollutants can also be destroyed by
heterogeneous hydroxyl radicals M(’OH) electro-generated on high oxygen overvoltage anodes (Eq. (2-13)).

\[
M (\text{H}_2\text{O}) \rightarrow M (\cdot \text{OH}) + \text{H}^+ + e^- \quad (2-13)
\]

Dirany et al. 2012 treated antibiotics sulfachloropyridazine (SCP) in aqueous solution using electro-Fenton process. When the applied current was 300 mA, the initial pH of 0.21 mM SCP solution was fixed at 3.0, SCP could be totally removed in 10 min. The mineralization efficiency of SCP was nearly 100% in a 360 min reaction (Table 2-2).

The second category is Fered-Fenton process or EF-Fere process. Both components of Fenton’s reagent (H\textsubscript{2}O\textsubscript{2} and Fe\textsuperscript{2+}) are added externally to the reactor. Fe\textsuperscript{3+} which formed in the Fenton’s reaction can convert to Fe\textsuperscript{2+} via cathodic reduction (Eq. (2-12)) (Figure 2-1b). Zhang et al. 2007 degraded 4-nitrophenol (4-NP) in water by using Fered-Fenton process. The COD removal efficiency for 200 mg/L 4-NP was 70% in a 120 min reaction, while the COD removal efficiency was only 50% in Fenton process at the same concentration of Fenton’s reagent (H\textsubscript{2}O\textsubscript{2} and Fe\textsuperscript{2+}) (Table 2-2).

The third category is electrochemical peroxidation process. A sacrificial iron anode was used for Fe\textsuperscript{2+} electrogeneration from anodic dissolution via Eq. (2-14). H\textsubscript{2}O\textsubscript{2} is externally added to the treated solution to degrade the organic pollutants with ’OH from Fenton’s reaction (Figure 2-1c). In electrochemical peroxidation process, the coagulation of Fe(OH)\textsubscript{3} precipitate formed depending on the pH and the applied current can be an significant alternative route for the degradation of organic pollutants (Brillas et al. 2009).

\[
\text{Fe} \rightarrow \text{Fe}^{2+} + 2e^- \quad (2-14)
\]

Electrochemical peroxidation process was employed to treat hexachlorobenzene (HCB) in aqueous solution by Xie et al (Xie et al. 2005). 96.96% of the initial HCB could be removed in 3 h (Table 2-2).
Figure 2-1 Schematic representation of different categories of EAOPs based on Fenton’s reagent ((a) electro-Fenton process; (b) Fered Fenton process; (c) electrochemical peroxidation process;)

<table>
<thead>
<tr>
<th>Categories</th>
<th>Compound</th>
<th>Typical condition</th>
<th>Corresponding results</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electro-Fenton</td>
<td>SCP</td>
<td>BDD anode, CF cathode, ([\text{SCP}]_0 = 0.21 \text{mM})</td>
<td>100% removal in 10 min and near 100% TOC removal in 360 min</td>
<td>Dirany et al. 2012</td>
</tr>
<tr>
<td></td>
<td></td>
<td>([\text{Fe}^{2+}] = 0.2 \text{mM}), (I = 300 \text{mA}), pH 3.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fered-Fenton</td>
<td>4-NP</td>
<td>Stainless steel cylinder cathode, ([4-\text{NP}]_0 = 200 \text{mg/L})</td>
<td>About 70% COD removal in a 120 min reaction</td>
<td>Zhang et al. 2007</td>
</tr>
<tr>
<td></td>
<td></td>
<td>([\text{RuO}_2/\text{IrO}_2) anode, ([\text{H}_2\text{O}_2]) = 9.12 \text{mM}, \text{H}_2\text{O}_2) feeding time 60 min, (\text{Fe}^{2+}/\text{H}_2\text{O}_2 = 0.05), (I = 1.0 \text{A}), pH 5.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>electrochemical</td>
<td>HCB</td>
<td>Stainless steel electrode, gap 30% between electrodes 6 cm, (\text{H}_2\text{O}_2) concentration 500 mg/L, pH 2.5</td>
<td>96.96% HCB removal in a 3 h reaction</td>
<td>Xie et al. 2005</td>
</tr>
</tbody>
</table>

2.3.2.2 Reactors for EAOPs based Fenton’s reaction

Generally, the reactors in which EAOPs based Fenton’s reaction were carried out
can be divided into two categories, i.e., the undivided cells and divided cells. Majority of researches were carried out in undivided electrochemical cells (Brillas et al. 1998; Mansour et al. 2015; Thirugnanasambandham et al. 2015; Zhang et al. 2015a). The major advantage of using undivided cells is that the electrolysis requires lower cell voltage because the voltage penalty of the separator of divided cells is avoided (Brillas et al. 2009). Nidheesh et al. 2014 degraded Rhodamine B (RhB) in a 1000 mL undivided cylindrical glass cell of 10.4 cm diameter containing 750 mL RhB solution (10 mg/L) by using electro-Fenton process. Two graphite plate with the same dimensions (5 × 5 cm) were used as anode and cathode. The results indicated RhB can be totally removed in a 180 min reaction.

However, reactive oxygen species and other weaker oxidants can also be generated at the anode in the undivided cell, which can complicate the electro-Fenton degradation process (Brillas et al. 2009). Therefore, it is harder to discuss the reaction mechanism in an undivided cell. Divided cells, in which cathode and anode are usually separated by membrane and salt bridge (Brillas et al. 2009) (Figure 2-2), are conducive to investigate the mechanism of cathodic Fenton process. Yuan et al. 2014 degraded 4-nitrophenol in a divided cell to investigate the mechanism of electro-Fenton process. Pozzo et al. 2005 employed an electrochemical cell divided by a cationic membrane to explore the production of hydrogen peroxide through the cathodic reduction of oxygen in acidic medium. Saltmiras and Lemley 2002 removed atrazine in a divided cell using anodic Fenton process in which ferrous ions were delivered via a sacrificial iron anode.
2.3.2.3 Improvements in electro-Fenton process

All the reactions mentioned above are homogeneous reactions, which have the disadvantages that the removal of metal ions from the effluent requires additional separation and disposal (Zhong et al. 2011). Therefore, various catalysts have been employed for the heterogeneous activation of $\text{H}_2\text{O}_2$. Garrido-Ramírez et al. 2013 used nanostructured allophane clays supported with iron oxide ($\text{AlSi}_2\text{Fe}_6$) as iron source of Fenton’s reaction to degrade atrazine in water by electro-Fenton process. 46 mM Atrazine could be totally removed in 8 h when the initial pH was 3.0. Iglesias et al. 2014 treated pesticide imidacloprid using a heterogeneous electro-Fenton system with ironalginate gel beads (EF-FeAB). The removal efficiency of 100 mg/L imidacloprid was 95% in a 120 min reaction.

2.3.3 Sulfate radical-based Fenton-like process

In recent years, activated persulfate (PS) oxidation is an emerging AOP for the treatment of organic pollutants (Rastogi et al. 2009; Tsitonaki et al. 2010). PS, which is discovered by M. Berthelot in 1878, is a stable oxidants with high aqueous solubility at ambient temperature (Cao et al. 2008). The commonly used PS in AOPs is
peroxydisulfate (PDS, $S_2O_8^{2-}$, $E^0 = 2.01$ V/SHE) and peroxymonosulfate (PMS, $HSO_5^-$, $E^0 = 1.82$ V/SHE). PS can be activated by transition metals (Fe$^{2+}$, Co$^{2+}$, Ag$^+$, etc.) (Romero et al. 2010; Zhao et al. 2010; Pagano et al. 2012; Wang and Chu 2012; Long et al. 2013) to generate sulfate radicals (SO$_4^{•−}$), which are promising in the treatment of organic contaminants in water ($E^0 = 2.6$ V/SHE) (Shukla et al. 2010; Cai et al. 2014; Wang et al. 2014). Transition metals activated PS process is similar to conventional Fenton process, so it also be named as Fenton-like process by Fernandez et al (Fernandez et al. 2003; Fernandez et al. 2004). Compared to other transition metals, Fe$^{2+}$ is inexpensive, nontoxic and effective, which has been widely used in catalytic oxidation process (Eq. (2-15) and (2-16)) (Wang and Chu 2011b; Li et al. 2015; Zhang et al. 2015b). Moreover, due to its unsymmetrical character, PMS might be more easily activated than PDS (Olmez-Hanci et al 2011).

$$S_2O_8^{2−} + Fe^{2+} → Fe^{3+} + SO_4^{•−} + SO_4^{2−} \quad (2-15)$$
$$HSO_5^- + Fe^{2+} → Fe^{3+} + SO_4^{•−} + OH^- \quad (2-16)$$

Similar to conventional Fenton process, there are some drawbacks encountered in sulfate radical-based Fenton-like process. For example, high ferrous ion dosage is required to activate PS due to the hard regeneration of Fe$^{2+}$. This results in large amount of iron sludge (Vicente et al. 2011). Like EAOPs based Fenton’s reaction, electrochemical process was combined with Fenton-like process and solved these problems (electro-Fenton-like process) (Wang and Chu 2011a; Wu et al. 2012).

In sulfate radical-based electro-Fenton-like process, Fe$^{2+}$ can be regenerated via cathodic reduction (Eq. (2-12)) of ferric iron ions. In addition, sulfate radicals could be produced via an electron transfer reaction (Eq. (2-17) and (2-18)) (Zhao et al. 2010; Zhou et al. 2011). Therefore, the degradation efficiency of organic pollutants was enhanced by coupling EC process and Fe$^{2+}$ activated PS process.

$$S_2O_8^{2−} + e^- → SO_4^{•−} + SO_4^{2−} \quad (2-17)$$
$$HSO_5^- + e^- → SO_4^{•−} + OH^- \quad (2-18)$$

According to different producing way of the PS and Fe$^{2+}$, electro-Fenton-like process can be divided into three categories.

In the first one, an iron electrode was used as anode and provided soluble ferrous
ions continuously through anodic oxidation (Eq. (2-14)). Once PS was added into the system, sulfate radicals were generated via the catalysis of ferrous ions (Eq. (2-17) or (2-18)) (Figure 2-3a).

Wang and Chu 2011a proposed an “electro-Fe(II)/Oxone (EFO)” process to degrade 2,4,5-trichlorophenoxyacetic acid (2,4,5-T). The EFO process demonstrated high 2,4,5-T removal efficiency and nearly completed the herbicide degradation within 10 min. Yuan et al. 2013 employed a similar system to degrade trichloroethylene (TCE). Cast gray iron was used as anode and provided Fe$^{2+}$, which could activate persulfate to generate sulfate radicals. 99% of 0.4 mM TCE could be transformed in a 20 min reaction time.

In the second category, both PS and Fe$^{2+}$ are added externally to the reactor. Fe$^{3+}$ which formed in Eq. (2-15) and (2-16) can transform to Fe$^{2+}$ via cathodic reduction (Eq. (2-12)) (Figure 2-3b). Wu et al. 2012 chose azo dye Orange II as the target pollutant and treated it using an “electro/Fe$^{2+}$/peroxydisulfate” process. 95% of 0.1 mM Orange II can be removed in a 60 min reaction with electrolyte concentration 0.1 M, Fe$^{2+}$ concentration 1 mM, PDS concentration 12 mM, and current density ($j$) 16.8 mA/cm$^2$. “Electro/Fe$^{2+}$/peroxydisulfate” process was also applied to the treatment of landfill leachate by Zhang et al. (Zhang et al. 2014). When PS concentration was 62.5 mM, Fe$^{2+}$ concentration was 15.6 mM, $j$ was 13.89 mA/cm$^2$ and initial pH 3.0, the COD removed by oxidation and coagulation was about 40% and 22%, respectively.

In the third category, Fe$^{2+}$ is replaced by Fe$^{3+}$ and added to the treated solution because Fe$^{2+}$ has the disadvantage of easily oxidized to Fe$^{3+}$ by air and ferrous solution should be stored under an acidic condition. Fe$^{3+}$ was first reduced to Fe$^{2+}$ (Eq. (2-12)) at the cathode and then activated PS to generate sulfate radicals. The regeneration of ferrous ions was enhanced by cathodic reduction (Eq. (2-12)) simultaneously (Figure 2-3c). “Electro/Fe$^{3+}$/PDS” and “EC/Fe$^{3+}$/PMS” processes were used to treat organic pollutants in water by our team (Lin et al. 2013, 2014). “Electro/Fe$^{3+}$/PDS” was used to degrade bisphenol A (BPA) which is a known endocrine disrupter and industrial chemical (Staples et al. 1998; Cleveland et al. 2014; Olmez-Hanci et al. 2015). BPA could be nearly completely removed in a 60 min treatment time with initial BPA
concentration of 0.22 mM, Fe$^{3+}$ concentration of 4 mM, PDS concentration of 20 mM, Na$_2$SO$_4$ concentration of 50 mM, initial pH value of 3.0 and the current density of 33.6 mA/cm$^2$. The mineralization efficiency reached 94.3% in a 120 min reaction at the same condition (Lin et al. 2013). Clofibric acid (CA), which is the active metabolite of several widely used blood lipid regulators, was treated by “EC/Fe$^{3+}$/PMS” process (Lin et al. 2014). When initial CA concentration was 50 mg/L, PMS concentration fixed at 20 mM, Fe$^{3+}$ concentration was 2.00 mM, Na$_2$SO$_4$ concentration was 50 mM, initial pH was 4.0 and the current density was 33.6 mA/cm$^2$, CA was completely removed after 60 min reaction.

![Figure 2-3](image)

Figure 2-3 Schematic representation of different types of electro-Fenton-like processes: (a) The first category; (b) The second category; (c) The third category)

All the electro-Fenton-like processes mentioned above are homogeneous processes. They have the same drawbacks as homogeneous electro-Fenton processes. In this study, iron metal oxides were used to replace Fe$^{2+}$ and activated PS.

### 2.4 Iron oxides applied in AOPs based on Fenton and Fenton-like process

Iron oxides are abundantly available minerals, presenting in the natural aqueous environment as suspended solid particles, and they also suspend in aerosol, clouds and fogs as fine particles (Hou et al. 2014; Rahim Pouran et al. 2014). Recently, iron oxides
are widely used in heterogeneous catalysis processes and constitute an attractive alternatives for the decontamination of soils, underground waters, sediments, and industrial effluents because they are natural, abundant, inexpensive, and environmentally friendly (Aredes et al. 2012; Xu et al. 2012; Rahim Pouran et al. 2014). Nowadays, several iron oxides and modified iron oxides are used to replace Fe$^{2+}$ in AOPs based Fenton and Fenton-like process (He et al. 2005; Guo et al. 2010; Ji et al. 2013; He et al. 2014; Sun et al. 2014).

2.4.1 Goethite ($\alpha$-FeOOH)

Goethite, an iron oxyhydroxide mineral with chemical formula of $\alpha$-Fe$^{III}$O(OH), is a commonly used iron oxide among Fe(III) bearing minerals for heterogeneous Fenton and Fenton-like based AOPs due to its ability to operate in a wide range of pH, higher stability thermodynamically and being relatively low priced and environmentally friendly (Ortiz de la Plata et al. 2008; Wang et al. 2010b; Rahim Pouran et al. 2014; Sable et al. 2015). Goethite is also used in electro-Fenton process as a substitute of Fe$^{2+}$ to react with H$_2$O$_2$ and form •OH through Fenton’s reaction, avoiding the additional water pollution caused by the homogeneous catalyst. Expósito et al. 2007 applied goethite as active heterogeneous catalyst undergo Fenton’s reaction with electrogenerated H$_2$O$_2$ in electro-Fenton process using aniline as the model pollutant. When initial aniline concentration was 100 mg/L, $\alpha$-FeOOH dosage fixed at 1 g/L, Na$_2$SO$_4$ concentration was kept at 20 mM, initial pH was 3.0 and the applied current was 250 mA, the TOC removal efficiency of aniline was about 90% in a 25 h reaction.

2.4.2 Magnetite (Fe$_3$O$_4$)

Magnetite is a spinel iron oxide with chemical formula of (Fe$^{3+}$)$_{tet}$[Fe$^{2+}$Fe$^{3+}$]$_{oct}$O$_4$ where Fe$^{3+}$ cations occupy equally both octahedral and tetrahedral sites and Fe$^{2+}$ cations are placed only in octahedral sites (Rahim Pouran et al. 2014). Magnetite has gained considerable attention than other iron oxides in AOPs based Fenton and Fenton-like process due to its unique characteristics: (i) The Fe(II) in its structure may play an
significant role as an electron donor to initiate the Fenton’s reaction (Kwan and Voelker 2003; Moura et al. 2005; Hou et al. 2014); (ii) The octahedral site in the magnetite structure can easily accommodate both Fe(II) and Fe(III), which means that Fe(II) can be reversibly oxidized and reduced in the same structure (Moura et al. 2005); (iii) The production of more active systems by modification in the physico-chemical properties of the magnetite through isostructural substitution of iron by different transition metals (Moura et al. 2005; Hou et al. 2014); (iv) The magnetically easy separation of magnetite catalysts from the reaction system as a result of its magnetic property (Ai et al. 2011; Chun et al. 2012) and (v) Higher dissolution rate of magnetite compared to other iron oxides which lead to higher electron mobility in its spinel structure (Litter and Blesa 1992; Matta et al. 2008).

Chen et al. 2014 synthesized Fe₃O₄ nanoparticles (NPs) by the oxidation-precipitation method and investigated their catalytic properties by the peroxide oxidation of Orange II solution. After 60 min reaction, the decolorization efficiency of Orange II was 99.89% with initial Orange II concentration of 100 mg/L, Fe₃O₄ NPs dosage of 1.5 g/L, H₂O₂ concentration of 22 mM and initial pH value of 2.7. Yan et al. 2011 applied iron oxide magnetic nanoparticles (Fe₃O₄ MNPs) to activate PDS and degrade antibiotics sulfamonomethoxine (SMM) in aqueous solution. When initial SMM concentration was 0.06 mM, Fe₃O₄ MNPs dosage fixed at 2.40 mM, PDS concentration was 1.2 mM and initial pH was 6.8, SMM could be completely removed in a 15 min reaction.

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Chapter 3 Treatment of aspartame (ASP) in aqueous solution by electro-Fenton process
3.1 Introduction

In this study, a detailed discussion on the oxidative degradation of artificial sweetener aspartame (ASP) in acidic aqueous solution containing catalytic amount of Fe$^{2+}$ by using electro-Fenton process was reported. The kinetics of ASP degradation by •OH generated during electro-Fenton process has been investigated. The absolute rate constant of the reaction between ASP and •OH was determined by the competition kinetic method using benzoic acid as a reference competitor (Özcan et al. 2009a, Özcan et al. 2013). The effect of the applied current and Fe$^{2+}$ concentration on the degradation and mineralization of ASP was examined. The aliphatic short-chain carboxylic acids released during electro-Fenton process were monitored. The concentration of inorganic ions released into the solution was investigated. The variation of toxicity of ASP solution and its intermediates was assessed by Microtox method.

3.2 Materials and methods

3.2.1 Chemicals

ASP, N-L-α-aspartyl-L-phenylalanine methyl ester (C$_{14}$H$_{18}$N$_{2}$O$_{5}$, ≥98%), was obtained from Alfa Aesar and was used in the electrolytic experiments without further purification. Analytical grade anhydrous sodium sulfate from Sigma-Aldrich was used as supporting electrolyte. Ferrous sulfate heptahydrated of analytical grade was purchased from Acros Organics and used as catalyst source. Analytical grade benzoic acid from Prolabo (R.P. Normapur grade) was used as the competition substrate in several kinetic experiments. Analytical grade carboxylic acids and or other chemicals used for chromatographic analysis were purchased from Acros, Sigma, Riedel-de Haën and Fluka. All solutions were prepared with ultrapure water obtained from a Millipore Mill-Q system with resistivity > 18 MΩ cm at room temperature.
3.2.2 Procedures and equipment

Bulk experiments were carried out at room temperature in a 250 mL undivided cylindrical glass cell of 6 cm diameter containing 220 mL ASP solution. The cathode (working) electrode was a 87.5 cm$^2$ piece of carbon felt (17.5 cm × 4.5 cm, Carbon-Lorraine, France) which placed on the inner wall of the cell covering the totality of the internal perimeter. The anode (counter) electrode was a cylindrical Pt mesh (4.5 cm height, i.d. = 3.1 cm, Platecxis, France) or a 24 cm$^2$ thin-film BDD electrode (CONDIAS GmbH, Germany). The anode was centered in the cell, surrounded by the carbon felt. Electrolysis was conducted under constant current conditions using a Hameg HM8040-3 triple power supply (Germany).

Compressed air was bubbled starting 5 min before electrolysis at about 0.5 L/min through the aqueous solutions to saturate aqueous solution before starting electrolysis and it is maintained during electrolysis to ensure the continuous saturation of oxygen. H$_2$O$_2$ was supplied from the electro-reduction of O$_2$ dissolved in the solution in all the electrolyses. The degradation experiments were performed using 0.2 mM ASP solution with 50 mM Na$_2$SO$_4$ as supporting electrolyte. A catalytic quantity of ferrous ion was added into the solution before the beginning of electrolysis. The initial pH ($\text{pH}_0$) of ASP solutions was measured with a CyberScan pH 1500 pH-meter (Eutech Instrument, USA) and set at 3.0 ($\pm$ 0.1), adjusting by the addition of 1 M sulfuric acid.

3.2.3 High performance liquid chromatography (HPLC) analysis

The initial and residual ASP concentration was monitored by HPLC, which consists of a Merck Lachrom liquid chromatograph, equipped with a L-2130 pump and was fitted with a Purospher RP-18, 5 $\mu$m, 25 cm × 4.6 mm (i.d.) column at 40 °C, and coupled with a L-2400 UV detector at maximum absorption wavelength of 215 nm for ASP. The analysis was performed using a 70:30 (v/v) water (0.1% acetic acid)/methanol (0.1% acetic acid) isocratic solvent mixture as mobile phase at a flow rate of 0.5 mL/min. The injection volume was 20 $\mu$L.

The short-chain carboxylic acids were identified and quantified by ion-exclusion
HPLC consisting of a Merck Lachrom liquid chromatograph equipped with a L-7100 pump, a Supelcogel H column (250 mm × 4.6 mm, 9 μm (i.d.) particle size) and a L-7455 photodiode array detector at a selected wavelength of 220 nm. 0.1% H₂SO₄ solution was used as mobile phase with a flow rate of 0.2 mL/min. Calibration curves were achieved using standard solutions of related carboxylic acids. The identification of the carboxylic acids was performed by the comparison between the retention time (t_R) and internal standard addition method using standard solutions.

3.2.4 Ion chromatography (IC) analysis

Inorganic ions (NO₃⁻ and NH₄⁺) released during the treatment were monitored by ion chromatography with a Dionex ICS-1000 Basic Ion Chromatography System equipped with an IonPac AS4A-SC (anion exchange) and CS12 A (cation exchange) 250 mm × 4 mm column and fitted with a DS6 conductivity detector containing a cell heated at 35 °C under control through a Chromelon SE software. The mobile phase was a mixture of 3.6 mM Na₂CO₃ and 3.4 mM NaHCO₃ solution for anion-exchange column with a flow rate of 2.0 mL/min and 9 mM H₂SO₄ solution for cation-exchange with a flow rate of 1.0 mL/min. The volume of injections was 25 μL.

3.2.5 Total organic carbon (TOC) analysis

The TOC of the initial and electrolyzed samples were determined by Shimatsu TOC-V_CSH analyser according to the combustion catalytic oxidation method at 680 °C. The carrier gas was oxygen with a flow rate of 150 mL/min. Platinum was used as catalyst to carry out the combustion at 650 °C instead of 900 °C. The injection volume was 50 μL. Calibration of the analyser was obtained with potassium hydrogen phthalate standards.

3.2.6 Toxicity measurements

The evolution of the toxicity of treated solutions during the treatment was assessed by Microtox method according to the international standard process (OIN 11348-3)
using a Microtox® model 500. This measurements performed with the bio-luminescence
marine bacteria *V. fischeri* (Buerge *et al.* 2009; Buerge *et al.* 2010; Bokare and Choi
2014), provided by Hach Lange France SAS. Two values of the inhibition of the
luminescence (%) were measured after 5 min and 15 min of exposure to samples at 15
°C.

### 3.3 Results and discussion

#### 3.3.1 Effect of catalyst concentration on ASP degradation

A set of electrolysis was carried out with 0.2 mM (58.9 mg/L) ASP in aqueous
solutions to determine the influence of the main operating parameters on the ASP
degradation in electro-Fenton process. The results obtained show that the oxidative
degradation of ASP by electro-Fenton process fits well with the first-order kinetic model
and the apparent rate constant values (*k*<sub>app</sub>) calculated accordingly under different
operating conditions were given in table 3-1.

**Table 3-1** Apparent rate constants (*k*<sub>app</sub>) obtained in electro-Fenton processes for ASP degradation,
assuming pseudo-first order kinetic model under different operating conditions. Operating conditions:

<table>
<thead>
<tr>
<th>Anode</th>
<th>[Fe&lt;sup&gt;2+&lt;/sup&gt;]/mM</th>
<th>I/mA</th>
<th><em>k</em>&lt;sub&gt;app&lt;/sub&gt; (min⁻¹)</th>
<th><em>R</em>²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt</td>
<td>0.05</td>
<td>200</td>
<td>0.15±0.01</td>
<td>0.994</td>
</tr>
<tr>
<td>Pt</td>
<td>0.1</td>
<td>200</td>
<td>0.24±0.01</td>
<td>0.992</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>200</td>
<td>0.34±0.01</td>
<td>0.995</td>
</tr>
<tr>
<td>Pt</td>
<td>0.3</td>
<td>200</td>
<td>0.26±0.01</td>
<td>0.996</td>
</tr>
<tr>
<td>Pt</td>
<td>0.5</td>
<td>200</td>
<td>0.22±0.02</td>
<td>0.999</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>50</td>
<td>0.14±0.01</td>
<td>0.997</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>100</td>
<td>0.15±0.01</td>
<td>0.998</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>300</td>
<td>0.34±0.03</td>
<td>0.968</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>500</td>
<td>0.17±0.01</td>
<td>0.997</td>
</tr>
</tbody>
</table>
The effect of catalyst (Fe$^{2+}$) concentration in ASP oxidation by electro-Fenton process was performed at current-controlled conditions and room temperature with Pt and BDD anode, respectively (Fig. 3-1). The Fe$^{2+}$ concentration was varied in the range of 0.05–0.5 mM in 50 mM Na$_2$SO$_4$ solution, at applied current 200 mA and initial pH 3.0. As can be seen in Fig. 3-1, ASP can be totally removed in a 20 min reaction using Pt anode, while it only needs 15 min to degrade ASP completely with BDD anode. This indicated that the use of BDD did lead to the acceleration of the ASP degradation. On the one hand, due to the low adsorption ability of •OH on BDD, the loosely bound BDD(•OH) formed at the anode surface (Eq. (3-1)) can readily react with organic pollutant, in contrast to the chemisorbed radicals typically formed at Pt surface which limited the oxidation ability of Pt(•OH) (Özcan et al. 2009a; Dirany et al. 2012). On the other hand, reactive BDD(•OH) is generated in much higher quantities than Pt(•OH) when electrolysis is operated at the current within the water discharge region (Brillas et al. 2005; Panizza and Cerisola 2005; Brillas et al. 2009) because of the large O$_2$ evolution overvoltage of the former.

$$\text{BDD} + \text{H}_2\text{O} \rightarrow \text{BDD}(\cdot\text{OH}) + \text{H}^+ + e^- \quad (3-1)$$

With both Pt and BDD anode, the removal efficiency and degradation rate of ASP increased significantly when Fe$^{2+}$ concentration increased from 0.05 to 0.2 mM (Fig. 3-1 and Table 3-1). At higher Fe$^{2+}$ concentration, a considerable amount of hydroxyl
radicals would be produced via Fenton’s reaction. These hydroxyl radicals react with ASP immediately, resulting in the increase of ASP degradation rate. However, further increasing the Fe\(^{2+}\) concentration to 0.5 mM, ASP removal efficiency and rate constant decreased evidently (Fig. 3-1 and Table 3-1). The negative influence of higher Fe\(^{2+}\) concentration might be attributed to the role of Fe\(^{2+}\) as scavenger of hydroxyl radicals (Eq. (3-2)) which takes place with a large rate constant \(k = 3.20 \times 10^8 \text{ M}^{-1} \text{ S}^{-1}\) (Sirés et al. 2007b; Oturan et al. 2010).

\[
\text{Fe}^{2+} + \cdot \text{OH} \rightarrow \text{Fe}^{3+} + \text{OH}^- \tag{3-2}
\]

Therefore, the reaction (3-2) becomes competitive for OH consumption at higher Fe\(^{2+}\) and consequently harms the oxidation of ASP.
Fig. 3-1 Effect of Fe$^{2+}$ (as catalyst) concentration on electro-Fenton degradation of 0.2 mM ASP with Pt (a) and BDD (b) anode versus carbon-felt cathode. Experimental conditions: $I = 200$ mA, pH$_0 = 3.0$, [Na$_2$SO$_4$] = 50 mM

3.3.2 Effect of applied current on ASP degradation

In electro-Fenton process, the applied current is a crucial parameter for the operational cost and process efficiency (Özcan et al. 2009a), because the formation rate of H$_2$O$_2$ (Eq. (3-4)), the regeneration rate of Fe$^{2+}$ (Eq. (3-3)) and consequently the generation rate of *OH through Fenton's reaction (Eq. (2-1)) are governed by this parameter. The applied current governs also the formation rate (Eq. (3-1)) and the amount of heterogeneous hydroxyl radicals BDD(*OH) or Pt(*OH). Therefore the effect of applied current for the treatment of 0.2 mM ASP in electro-Fenton processes using Pt/carbon-felt (Fig. 3-2a) and BDD/carbon-felt (Fig. 3-2b) cells was examined by using the applied current of 50, 100, 200, 300 and 500 mA.

As depicted in Fig. 3-2a, the total disappearance of ASP became faster at a higher current in Pt/carbon-felt cell. The $k_{app}$ increased from 0.14 to 0.34 min$^{-1}$ when the applied current increased from 50 to 300 mA (Table 3-1). A higher current can promote both the Fe$^{2+}$ regeneration (Eq. (3-3)) and the production of H$_2$O$_2$ (Eq. (3-4)) (Lin et al.
2014). However, further increasing the current to 500 mA resulted in the decrease of $k_{app}$ to 0.17 min$^{-1}$. The decrease in ASP oxidation efficiency at higher current can be related to the increase of parasitic reactions, such as H$_2$ evolution reaction (Eq.(3-5)) (Dirany et al. 2012).

$$Fe^{3+} + e^- \rightarrow Fe^{2+} \quad (3-3)$$

$$O_2 + 2H^+ + 2e^- \rightarrow H_2O_2 \quad (3-4)$$

$$2H_2O + 2e^- \rightarrow H_2 + 2OH^- \quad (3-5)$$

When it comes to the BDD/carbon-felt cell, the optimal current for ASP degradation was 200 mA (Fig. 3-2b). Moreover, it can be seen from Fig. 2 that the comparatively better performance of BDD was more pronounced at lower current intensities. When applied current is higher than 200 mA, the positive role of BDD favoring generation of \(^{\cdot}OH\) was insignificant (Table 3-1). This means the application of BDD anode can improve removal efficiency of ASP and reduce energy consumption by using lower current intensity. A similar behavior has been observed for the degradation of sulfachloropyridazine (Dirany et al. 2012). At current intensity above 200 mA, the removal efficiency decreased. This could due to the increase of side reactions: (i) H$_2$ evolution from H$_2$O reduction (Eq. (3-5)). (ii) The 4-e$^-$ reduction of O$_2$ leading to the formation of H$_2$O (Eq. (3-6)) which inhibited the H$_2$O$_2$ formation reaction (Eq. (3-4)) occurring on carbon-felt cathode (Özcan et al. 2009b). Moreover, the O$_2$ evolution reaction was more dominate than \(^{\cdot}OH\) on the BDD anode at high applied current (Özcan et al. 2009b). (iii) The formation of weak oxidant species like HO$_2^{\cdot}$ (Eq. (3-7)) which could react with \(^{\cdot}OH\) resulting in the formation of H$_2$O and O$_2$ (Eq.(3-8)) (Wu et al. 2012).

$$O_2 + 4H^+ + 4e^- \rightarrow H_2O \quad (3-6)$$

$$H_2O_2 + \cdot OH \rightarrow HO_2^{\cdot} + H_2O \quad (3-7)$$

$$HO_2^{\cdot} + \cdot OH \rightarrow H_2O + O_2 \quad (3-8)$$
Fig. 3-2 Effect of applied current on electro-Fenton degradation of 0.2 mM ASP by Pt (a) and BDD (b) anode versus carbon-felt cathode. Experimental conditions: [Fe^{2+}] = 0.2 mM, pH₀ = 3.0, [Na₂SO₄] = 50 mM
3.3.3 Determination of absolute constants for oxidation of ASP by hydroxyl radicals

The absolute rate constant of ASP treatment by hydroxyl radicals during electro-Fenton process was determined by competition kinetics method. Benzoic acid (BA) was selected as standard competitor because the absolute rate constant \( k_{abs,BA} \) of the reaction between BA and hydroxyl radicals is well known \( k_{abs,BA} = 4.30 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \) (Özcan et al. 2008; Sirés et al. 2007c; Oturan et al. 2010). Experiments were performed using Pt anode with 0.1 mM ASP solution, \( \text{Fe}^{2+} \) concentration 0.2 mM, applied current 50 mA and initial pH 3.0. The absolute rate constant of the oxidation of ASP was then calculated according to the Eq. \((3-9)\).

\[
\ln \left( \frac{[\text{ASP}]_0}{[\text{ASP}]_t} \right) = \left( \frac{k_{abs,\text{ASP}}}{k_{abs,\text{BA}}} \right) \ln \left( \frac{[\text{BA}]_0}{[\text{BA}]_t} \right)
\]

\((3-9)\)

Based on Fig. 3-3, the absolute rate constant for the oxidation reaction of ASP and \(^{\prime}\text{OH} \) \( k_{abs,\text{ASP}} \) was then determined as \((5.23 \pm 0.02) \times 10^9 \text{ M}^{-1} \text{s}^{-1} \). This value is closed to that reported by Toth et al. 2012 \((6.06 \pm 0.05) \times 10^9 \text{ M}^{-1} \text{s}^{-1} \) who determined the absolute rate constant by direct observation of the formation of the cyclohexadienyl radical adduct.

\[y = 1.1913x + 0.01624\]
\[R^2 = 0.9973\]

**Fig. 3-3** Determination of \( k_{abs(ASP)} \) by competition kinetic method according to the pseudo first-order kinetics between ASP and BA during the electro-Fenton process. Experimental conditions: \([\text{ASP}]_0 = 0.1 \text{ mM}, ([\text{BA}]_0 = 0.1 \text{ mM}, [\text{Fe}^{2+}] = 0.2 \text{ mM, } I = 50 \text{ mA, pH} 3.0, [\text{Na}_2\text{SO}_4] = 50 \text{ mM}\)
3.3.4 The effect of catalyst concentration and applied current on ASP mineralization

Fig. 3-4 shows the influence of Fe\textsuperscript{2+} concentration on the mineralization of ASP solution in terms of TOC abatement under the same conditions of Fig. 3-1.

For both Pt/carbon-felt and BDD/carbon-felt cell, the TOC removal efficiency decreased as Fe\textsuperscript{2+} concentration increased from 0.1 to 0.5 mM. With the increasing Fe\textsuperscript{2+} concentration, the percentage of scavenged hydroxyl radicals from Fe\textsuperscript{2+} ions increased, so the side reactions between Fe\textsuperscript{2+} ions and •OH (Eq. (3-2)) became more significant (Özcan et al. 2008). This reaction becomes particularly important at longer treatment time because of lower concentration of organic matter in the solution. Furthermore, as can be seen from Fig. 3-4, the TOC removal rate was fast during the first 120 min for all the Fe\textsuperscript{2+} concentration in both cells. After 120 min, the TOC removal values gradually decreased and reached a steady state. This is due to the quick decomposition of ASP and its aromatic derivatives which are more easily oxidizable than the ring opening products such as aliphatic acids obtained at longer treatment times (Hammami et al. 2008). The slower mineralization rate observed at longer electrolysis times for both Pt and BDD anodes can also be attributed to the formation of stable ferro- and/or ferri-complexes with carboxylic acids (Oturan et al. 2000; Brillas et al. 2003; Guivarch et al. 2003; Hammami et al. 2008; Haidar et al. 2013).

Even so, it is evident that the use of BDD anode lead to the acceleration of ASP mineralization (Fig. 3-4). For example, at the end of the 360 min electrolysis, 84.5% and 98.0% of the initial TOC value was removed, in the case of Fe\textsuperscript{2+} concentration of 0.1 mM, for Pt and BDD anode, respectively. Moreover, BDD/carbon-felt cell achieved > 95% mineralization in only 240 min at 0.1 mM Fe\textsuperscript{2+}, while 84.5% mineralization reached at 360 min for Pt/carbon-felt cell. It can be explained that BDD anode contributed to the degradation of all the byproducts, even the most refractory ones (Dirany et al. 2012) due to the high reactivity of BDD(•OH) largely formed at this anode.
The effect of applied current on the mineralization of ASP solution in terms of TOC abatement under the same conditions of Fig. 3-2 was investigated and the results were shown in Fig. 3-5. With the raising applied current from 100 to 200 mA, the TOC removal efficiency was found to increase from 72.2% to 81.7% in Pt/carbon-felt cell. The TOC removal efficiency increased from 93.1% to 97.5% in BDD/carbon-felt cell at
the end of 360 min electrolysis under the same conditions. Higher applied current lead to the higher accumulation of hydroxyl radicals in the bulk, and then a greater mineralization efficiency can be achieved because of the simultaneous degradation of ASP and its byproducts (Dirany et al. 2012). Further increase in the applied current to 500 mA, the TOC removal efficiency declined to 71.6% and 93.4% for Pt and BDD anode, respectively. In the treatment of ASP, the application of current higher than 200 mA would increase the extent of parasitic reactions (Eq. (3-5)-(3-8)), leading to the decrease of the process efficiency (Mousset et al. 2014).

**Fig. 3-5** Effect of applied current on mineralization of 0.2 mM ASP with Pt (a) and BDD (b) anode versus carbon-felt cathode. Experimental conditions: \([\text{Fe}^{2+}] = 0.2 \text{ mM}, \ pH_0 \ 3.0, \ [\text{Na}_2\text{SO}_4] = 50 \text{ mM}\)
3.3.5 Identification and evolution of carboxylic acids

It is well known that the cleavage of the benzenic ring of aromatics can lead to the release of short-chain carboxylic acids in electro-Fenton process (Sirés et al. 2007a; Özcan et al. 2009b; Isarain-Chávez et al. 2010; Dirany et al. 2012; El-Ghenemy et al. 2013). The concentration of carboxylic acids were monitored by ion-exclusion HPLC when 0.2 mM ASP solutions were electrolyzed under the conditions of Figs. 3-4 and 3-5. In this study, oxalic, oxamic and maleic acid at retention time ($t_R$) of 8.90, 14.26 and 11.33 were observed, respectively, in both Pt/carbon-felt and BDD/carbon-felt cell. Among these carboxylic acids, the concentration of maleic acid was detected in trace and only exist in the first 10 min in both cells, so changes of maleic acid concentration were not shown in Fig. 3-6.

As shown in Fig. 3-6, all the carboxylic acids were generated at the beginning of the electrolysis and followed the accumulation-destruction cycles. A similar behavior has been observed for the treatment of other organic products (Özcan et al. 2009b; Dirany et al. 2012). Oxalic and oxamic acids were remained even at the end of the electrolysis indicating their lower reactivity with hydroxyl radicals and need much more time to destruction (Dirany et al. 2012). It also depicted by comparing Fig. 3-6a and Fig. 3-6b that all the acids were nearly disappeared at the end of electrolysis (360 min) in BDD/carbon-felt cell, in agreement with the mineralization results shown in Fig. 3-4b and Fig. 3-5b. However, oxalic and oxamic acid were still existed in Pt/carbon-felt cell in 360 min reaction, which was corresponding to the residual TOC remaining at the end of mineralization treatments (Fig. 3-4a and Fig. 3-5a).
Fig. 3-6 Evolution of short-chain carboxylic acids formed during electro-Fenton processes with Pt (a) and BDD (b) anodes versus carbon-felt cathode. Experimental conditions: [ASP]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, I = 200 mA, pH₀ = 3.0, [Na₂SO₄] = 50 mM

3.3.6 Identification and evolution of inorganic ions

During the mineralization of ASP, the N atoms were released as inorganic ions upon the cleavage of the ASP molecules. The qualitative and quantitative monitoring of NO₃⁻ and NH₄⁺ was performed by the ionic chromatography analysis. The evolution of the ions concentration identified during the electro-Fenton process was shown in Fig.
The final concentration of NO$_3^-$ was much higher with BDD anode, which agrees with the faster mineralization mentioned in Fig. 3-4. NO$_3^-$ and NH$_4^+$ were formed from the beginning of the electrolysis. In Pt/carbon-felt cell, the release of ammonium ions was much larger than that of nitrate ions in 240 min reaction, but it decreased to 0.016 mM after 360 min reaction. The release of NO$_3^-$ in Pt/carbon-felt cell increased from 0 to 0.011 mM. Moreover, Fig. 6a indicated that 0.15 mM oxamic acid was existed in Pt/carbon-felt cell at 360 min reaction. At the end of electrolysis, 43.2% of initial nitrogen of ASP as NO$_3^-$ (6.35% of total), NH$_4^+$ (9.25% of total) and oxamic acid (84.4% of total) was quantified. This indicated some other refractory nitrogenated compounds are formed in the Pt/carbon-felt cell that account for the remaining TOC at 360 min. In BDD/carbon-felt cell, the concentration of NH$_4^+$ increased to 0.07 mM in the first 60 min, and then it decreased to 0.004 mM at 360 min. NO$_3^-$ increased persistently from the beginning to the end of the reaction and the concentration reached 0.116 mM at 360 min. Since TOC was nearly completely removed in BDD/carbon-felt cell at 360 min reaction, the non equilibrating of the nitrogen mass balance could be attributed to the transformation of nitrogen to other nitrogen species, such as NO$_2^-$, gas N$_2$ and NH$_3$ (Hammami et al. 2008).
Fig. 3-7 Evolution of the concentration of the inorganic ions released during electro-Fenton processes with Pt (a) and BDD (b) anode versus carbon-felt cathode. Experimental conditions:

\[ [\text{ASP}]_0 = 0.2 \text{ mM}, \ [\text{Fe}^{2+}] = 0.2 \text{ mM}, \ I = 200 \text{ mA}, \ \text{pH}_0 = 3.0, \ [\text{Na}_2\text{SO}_4] = 50 \text{ mM} \]

### 3.3.7 The evolution of solution toxicity with reaction time

In order to monitor the potential toxicity of ASP and its byproducts, 0.2 mM ASP was electrolyzed in the presence of 0.2 mM Fe\(^{2+}\) at 200 mA using Microtox method for both Pt and BDD anode. The percentage of bacteria luminescence inhibition versus the electrolysis time was recorded after 5 and 15 min exposure times of *V. fischeri* luminescent bacteria to the ASP solutions. In Fig. 3-8, only the curves obtained after a 15 min exposure time were presented, because the curves recorded after a 5 min exposure time were very similar.

For Pt/carbon-felt cell, the curves of inhibition percentage were characterized by a strong increase of the toxicity at 10 min. This indicated that the formed intermediates of ASP at the beginning of the electrolysis, which might mainly contain the cyclic compounds, were more toxic than ASP. The percentage of inhibition decreased sharply after 10 min, which showed the decrease of toxic intermediate products. The secondary peak which appeared at 120 min was much lower than the primary peak, showing the toxicity of secondary intermediates were much lower than the former byproducts, even a little lower than the untreated ASP solution. After 120 min, the inhibition percentage
decreased slowly to the end of the electrolysis.

The curves obtained for BDD anode showed an analogous behavior at the beginning of the electrolysis as Pt anode. The toxicity increased significantly and attained the maximum luminescence inhibition peak at 15 min. The secondary peak appeared at 40 min and showed the secondary intermediates still more toxic than the original ASP solution. After 40 min treatment, the percentage of inhibition decreased and became stable from 120 min. This might due to the destruction of ASP and its cyclic byproducts.

**Fig. 3-8** Evolution of the inhibition of marine bacteria, *Vibrio fischeri* luminescence (Microtox method) during electro-Fenton processes. Experimental conditions: [ASP]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, I = 200 mA, pH₀ 3.0, [Na₂SO₄] = 50 mM

### 3.4 Conclusions

Aqueous solution of 0.2 mM ASP was degraded effectively by electro-Fenton process using a carbon-felt cathode and a Pt or BDD anode. Compete removal of ASP was attained by both anodes, due to the formation of •OH in the bulk from Fenton reaction and at the anode surface from water oxidation. For both anodes, ASP could be completely removed in less than 30 min and the ASP concentration decay followed pseudo-first-order kinetics. Absolute rate constant of hydroxylation reaction of ASP was determined as $(5.23 \pm 0.02) \times 10^{9}$ M⁻¹ s⁻¹. The use of BDD instead of Pt anode yields a faster mineralization rate because of the higher oxidation power of BDD(•OH)
comparing with Pt(•OH). Short-chain aliphatic carboxylic acids such as oxalic, oxamic and maleic acid were identified as aliphatic intermediates by ion-exclusion chromatography. The bacteria luminescence inhibition showed the toxicity of ASP solution increased at the beginning of electrolysis, and then it declined until lower than the untreated ASP solution at the end of the reaction.

References


Chapter 4 Treatment of saccharin (SAC) in aqueous solution by

electro-Fenton process
4.1 Introduction

In this part, the effect of different anode materials and different supporting electrolyte on the removal of sweetener saccharine (SAC) was investigated by electro-Fenton process. The absolute rate constant of the reaction between SAC and \textsuperscript{\textprime}OH was determined by the competition kinetic method. The effect of the Fe\textsuperscript{2+} concentration and applied current on the degradation and mineralization of SAC was examined. The aliphatic short-chain carboxylic acids released during electro-Fenton process were monitored. The variation of toxicity of SAC solution and its intermediates was determined by Microtox method.

4.2 Materials and methods

4.2.1 Chemicals

Saccharin (1,1-dioxo-1,2-benzothiazol-3-one (or o-benzoic sulfimide), \(\text{C}_7\text{H}_5\text{CNO}_3\text{S}\)) was purchased from Sigma-Alorich. Analytical grade anhydrous sodium sulfate and ferrous sulfate heptahydrated were obtained from Sigma-Aldrich and Acros Organics, respectively. Regent grade benzoic acid was purchased from Problabo (France) used as the competition substrate in several kinetic experiments. Analytical grade carboxylic acids and other chemicals used for chromatographic analysis were purchased from Acros, Merck, Sigma, Riedel-de Haën and Fluka. Ultrapure water used for the preparation of the working solutions and HPLC eluting solutions was obtained from a Millipore Milli-Q (simplicity 185) system with resistivity > 18 M\(\Omega\) cm at room temperature.

4.2.2 Electrochemical apparatus and procedures

Electrolyses were performed at constant current and room temperature using a Hameg HM8040-3 triple power supply (Germany) in an open, cylindrical undivided glass cell of 6 cm diameter and 250 mL capacity containing 220 mL SAC solution.
Electro-Fenton oxidation was conducted using three anodes: a cylindrical Pt mesh (4.5 cm height, i.d. = 3.1 cm, Platecxis, France), a 25 cm\(^2\) thin-film BDD (CONDIAS GmbH, Germany) and a commercial DSA (mixed metal oxide Ti/RuO\(_2\)-IrO\(_2\), Baoji Xinyu GuangJiDian Limited Liability Company, China). A 87.5 cm\(^2\) piece of carbon felt (17.5 \(\times\) 5 cm, Carbon-Lorraine, France) was used as cathode.

In all cases, the anode was centered in the electrolytic cell and was surrounded by the cathode that covered the inner wall of the cell. H\(_2\)O\(_2\) was produced from reduction of O\(_2\) dissolved in the solution. Continuous saturation of O\(_2\) at atmospheric pressure was ensured by bubbling compressed air passing through a frit at about 0.5 L/min, starting 5 min before the beginning of the electrolysis. Prior to the electrolysis, a catalytic quantity of ferrous ion and 50 mM Na\(_2\)SO\(_4\) (supporting electrolyte) were added to the SAC solutions with constant stirring by using a magnetic stirrer (IKA, Germany).

The pH of initial solutions (pH\(_0\)) was set at 3.0 by the addition of 1 M sulfuric acid because this value was reported as the optimal pH value for the electro-Fenton processes (Özcan \textit{et al.} 2008a). The pH of SAC solutions was measured with a CyberScan pH 1500 pH-meter (Eutech Instrument, USA).

### 4.2.3 Analytical methods and procedures

The concentration of residual SAC was monitored by high performance liquid chromatography (HPLC), which consist of a Merck Lachrom liquid chromatograph, equipped with a L-2130 pump and fitted with a Purospher RP-18, 5 \(\mu\)m, 25 \(\times\) 4.6 mm (i.d.) column at 40 °C, and coupled with a L-2400 UV detector at maximum absorption wavelength of 218 nm for SAC. The analyses were performed using a phosphoric acid (pH 3.0)/methanol (80:20, \(v/v\)) as mobile phase in isocratic mode at a flow rate of 1.0 mL/min. The injection volume was 20 \(\mu\)L.

The short-chain carboxylic acids were identified and quantified by ion-exclusion HPLC using a L-7100 pump, a Merck Lachrom liquid chromatograph equipped with a Supelcogel H column (250 \(\times\) 4.6 mm, 9 \(\mu\)m) and a L-7455 photodiode array detector at the wavelength of 220 nm. The mobile phase was 0.1% H\(_2\)SO\(_4\) solution and flow rate
was fixed to 0.2 mL/min. Calibration curves were achieved using standard solutions of related carboxylic acids. The identification of the carboxylic acids was performed by the retention time \((t_R)\) comparison and standard addition methods using standard substances.

The TOC of the samples withdrawn from the treated solution at different electrolysis times was determined by Shimatzu TOC-VCSH analyser consisting of a non dispersive infra-red absorption detector (NDIR) according to the 680 °C combustion catalytic oxidation method. Platinum based catalyst was used to facilitate the combustion at 650 °C. The carrier gas was oxygen with a flow rate of 150 mL/min. The injection volume was 50 μL.

### 4.2.4 Toxicity measurements

The toxicity of SAC and its intermediates generated in electro-Fenton processes was investigated on samples collected from solutions at different electrolysis times. Experiments were performed with the bio-luminescence marine bacteria *V. fischeri* (Hach Lange France SAS) by Microtox method according to the international standard process (OIN 11348-3). Two values of the inhibition of the luminescence (%) were measured after 5 min and 15 min of exposure to samples at 15 °C using a Microtox model 500 system.

### 4.3 Results and discussion

#### 4.3.1 Effect of supporting electrolyte on the degradation of SAC

In order to investigate the effect of the supporting electrolyte on the degradation kinetics of SAC aqueous solutions, experiments were performed in acidic medium (pH 3.0) containing different supporting electrolytes as 50 mM Na₂SO₄, 100 mM NaNO₃ and 100 mM NaCl with three anodes (Fig. 4-1). The removal of SAC followed the first-order kinetic model and the apparent rate constant values \((k_{app})\) were determined.
accordingly and given in table 4-1.

Table 4-1 Apparent rate constants ($k_{app}$) obtained in electro-Fenton processes for SAC degradation, assuming pseudo-first order kinetic model under different operating conditions.

<table>
<thead>
<tr>
<th>Electrode</th>
<th>$[\text{Fe}^{2+}]$/mM</th>
<th>$I$/mA</th>
<th>supporting electrolyte</th>
<th>$k_{app}$ (min$^{-1}$)</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>DSA</td>
<td>0.2</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.181±0.004</td>
<td>0.997</td>
</tr>
<tr>
<td>DSA</td>
<td>0.2</td>
<td>200</td>
<td>NaNO$_3$ (100 mM)</td>
<td>0.170±0.030</td>
<td>0.998</td>
</tr>
<tr>
<td>DSA</td>
<td>0.2</td>
<td>200</td>
<td>NaCl (100 mM)</td>
<td>0.019±0.001</td>
<td>0.994</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.211±0.008</td>
<td>0.990</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>200</td>
<td>NaNO$_3$ (100 mM)</td>
<td>0.125±0.007</td>
<td>0.977</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>200</td>
<td>NaCl (100 mM)</td>
<td>0.038±0.001</td>
<td>0.999</td>
</tr>
<tr>
<td>BDD</td>
<td>0.2</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.194±0.008</td>
<td>0.988</td>
</tr>
<tr>
<td>BDD</td>
<td>0.2</td>
<td>200</td>
<td>NaNO$_3$ (100 mM)</td>
<td>0.143±0.002</td>
<td>0.998</td>
</tr>
<tr>
<td>BDD</td>
<td>0.2</td>
<td>200</td>
<td>NaCl (100 mM)</td>
<td>0.023±0.001</td>
<td>0.999</td>
</tr>
<tr>
<td>Pt</td>
<td>0.05</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.047±0.004</td>
<td>0.945</td>
</tr>
<tr>
<td>Pt</td>
<td>0.1</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.108±0.003</td>
<td>0.995</td>
</tr>
<tr>
<td>Pt</td>
<td>0.3</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.147±0.012</td>
<td>0.953</td>
</tr>
<tr>
<td>Pt</td>
<td>0.5</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.063±0.004</td>
<td>0.972</td>
</tr>
<tr>
<td>BDD</td>
<td>0.05</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.077±0.002</td>
<td>0.994</td>
</tr>
<tr>
<td>BDD</td>
<td>0.1</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.106±0.003</td>
<td>0.995</td>
</tr>
<tr>
<td>BDD</td>
<td>0.3</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.117±0.001</td>
<td>0.999</td>
</tr>
<tr>
<td>BDD</td>
<td>0.5</td>
<td>200</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.095±0.002</td>
<td>0.996</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>50</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.070±0.001</td>
<td>0.999</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>100</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.176±0.005</td>
<td>0.994</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>300</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.256±0.005</td>
<td>0.998</td>
</tr>
<tr>
<td>Pt</td>
<td>0.2</td>
<td>500</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.259±0.005</td>
<td>0.999</td>
</tr>
<tr>
<td>BDD</td>
<td>0.2</td>
<td>50</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.089±0.001</td>
<td>0.998</td>
</tr>
<tr>
<td>BDD</td>
<td>0.2</td>
<td>100</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.137±0.005</td>
<td>0.991</td>
</tr>
<tr>
<td>BDD</td>
<td>0.2</td>
<td>300</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.187±0.006</td>
<td>0.993</td>
</tr>
<tr>
<td>BDD</td>
<td>0.2</td>
<td>500</td>
<td>Na$_2$SO$_4$ (50 mM)</td>
<td>0.156±0.002</td>
<td>0.998</td>
</tr>
</tbody>
</table>

For all the anodes, the complete removal of SAC almost finished in a 30 min electrolysis period in the presence of Na$_2$SO$_4$ and NaNO$_3$. However, it was removed
only about 50% in 30 min in the case of NaCl. The main reason of this low degradation is related to the formation of active chlorine generated by the oxidation of chloride ions at the surface of DSA, Pt and BDD anode when NaCl used as supporting electrolyte (Eq. (4-1)) (Özcan et al. 2008b; Loaiza-Ambuludi et al. 2013; De Moura et al. 2014). Therefore, part of electrical energy provided is lost in this reaction.

\[ 2\text{Cl}^- \rightarrow \text{Cl}_2 + e^- \quad (4-1) \]

The electrogenerated active chlorine can acts as oxidation mediator in the bulk of the solution, which can accelerate the removal rate of organic pollutants through the production of HClO (Eq. (4-2)) according to some researches (Özcan et al. 2008b; Loaiza-Ambuludi et al. 2013).

\[ \text{Cl}_2 + \text{H}_2\text{O} \rightarrow \text{HClO} + \text{Cl}^- + \text{H}^+ \quad (4-2) \]

However, in electro-Fenton process, the electrogenerated chlorine can also react with Fe\(^{2+}\) (Eq. (4-3)) or decompose hydrogen peroxide (Eq. (4-4)) (Loaiza-Ambuludi et al. 2013), and then reducing the production rate of strong oxidant \(^{\cdot}\text{OH}\) by Fenton’s reaction.

\[ 2\text{Fe}^{2+} + \text{Cl}_2 \rightarrow 2\text{Fe}^{3+} + 2\text{Cl}^- \quad (4-3) \]

\[ 2\text{H}_2\text{O}_2 + \text{Cl}_2 \rightarrow 2\text{HCl} + \text{H}_2\text{O} + \text{O}_2 \quad (4-4) \]

Moreover, Cl\(^-\) could consume \(^{\cdot}\text{OH}\) and lead to the formation of ClOH\(^{\cdot}\) through Eq. (4-5) (Nidheesh et al. 2014). Oxidative capacity of ClOH\(^{\cdot}\) towards organic compounds is much lower than that of \(^{\cdot}\text{OH}\) formed during the EF process.

\[ \text{Cl}^- + ^{\cdot}\text{OH} \rightarrow \text{ClOH}^{\cdot} \quad (4-5) \]

Therefore, the removal efficiency of SAC was much lower when NaCl used as background electrolyte.

It can also be seen from Fig. 4-1 and Table 4-1 that the removal rate of SAC for Na\(_2\)SO\(_4\) was a little higher than that obtained for NaNO\(_3\) in three electrolysis cells. For example, when DSA was used as anode and at 200 mA applied current, the acceleration for Na\(_2\)SO\(_4\) was 0.181 min\(^{-1}\), while it was 0.170 min\(^{-1}\) for NaNO\(_3\). Therefore, Na\(_2\)SO\(_4\) was chosen to be the supporting electrolyte in the following experiments.
4.3.2 Effect of anode materials on the removal and mineralization of SAC

The removal of SAC by electro-Fenton process using DSA, Pt and BDD as anodes was investigated by keeping the same cathode as carbon-felt. Experiments were performed at Fe$^{2+}$ concentration 0.2 mM, Na$_2$SO$_4$ concentration 50 mM, applied current 200 mA and pH$_0$ 3.0. 0.2 mM SAC could be completely removed in a 30 min reaction for all the anode materials (Fig. 4-2a). The $k_{app}$ values for SAC removal using DSA, Pt and BDD anodes were very similar (0.181, 0.194 and 0.211 min$^{-1}$, respectively). However, when it comes to mineralization, BDD anode showed its great superiority. In a 360 min reaction, the TOC removal efficiencies for SAC were 55.8%, 76.1% and 96.2% for DSA, Pt and BDD anodes, respectively (Fig. 4-2b). On the one hand, the BDD(’OH) radicals can effectively mineralize short chain carboxylic acids generated in the electro-Fenton process, which are relatively recalcitrant to mineralization (Oturan et al. 2013). On the other hand, Fe(III)–carboxylic acid complexes formed in electro-Fenton process were difficultly oxidizable with ’OH produced in the medium from Fenton’s reaction and at the DSA and Pt anode surface from Eq. (4-6) (Boye et al. 2002; Sirés et
These complexes are completely destroyed using a BDD anode due to the great amount of \( \cdot \text{OH} \) generated on its surface (Sirés et al. 2007) because of its large O\(_2\) evolution overpotential. Since poor TOC removal efficiency was obtained using DSA anode, only Pt and BDD anodes were applied in the following experiments.

\[
\text{M (H}_2\text{O)} \rightarrow \text{M (OH)} + \text{H}^+ + e^- \tag{4-6}
\]

**Fig. 4-2** Comparison of the performance of different anode in the degradation (a) and mineralization (b) of SAC ([SAC]\(_0\) = 0.2 mM, [Fe\(^{2+}\)] = 0.2 mM, [Na\(_2\)SO\(_4\)] = 50 mM, \(I = 200\) mA, \(pH_0\) = 3.0)**
4.3.3 Effect of Fe\(^{2+}\) concentration on the removal of SAC

The effect of Fe\(^{2+}\) (catalyst) concentration on the removal of SAC was examined by using the Fe\(^{2+}\) concentration of 0.05, 0.1, 0.2, 0.3 and 0.5 mM. The time-course of SAC concentration decay was determined by HPLC, where SAC displayed a well-defined peak at the retention time \((t_R)\) 14.2 min under the operating conditions of Fig. 4-3.

As can be observed in Fig.4-3 and Table 4-1, the removal rate for SAC was enhanced by increasing the Fe\(^{2+}\) concentration from 0.05 to 0.2 mM for both Pt and BDD anode. Moreover, the kinetics of SAC removal increases more rapidly (1.8 times) in the case of Pt/carbon-felt cell when the Fe\(^{2+}\) increased from 0.05 to 0.2 mM, compared to BDD/carbon-felt cell. Since the applied current was kept constant at 200 mA, it can be assumed that the production rate of H\(_2\)O\(_2\) via the oxygen reduction would be identical for all the Fe\(^{2+}\) concentrations under this study (Oturan et al. 2010). Then increasing the Fe\(^{2+}\) concentration could lead to the increase of hydroxyl radicals generated via Fenton’s reaction. These hydroxyl radicals reacted with SAC immediately, resulting in the increase of SAC degradation. On the contrary, further increasing the Fe\(^{2+}\) concentration to 0.5 mM, the removal rate declined. The observed decrease in SAC degradation rate upon increasing the Fe\(^{2+}\) concentration might due to the role of Fe\(^{2+}\) as scavenger of hydroxyl radicals (Eq. (4-7)) which takes place with a large rate constant 
\((k = 3.20 \times 10^8 \text{ M}^{-1} \text{ s}^{-1})\) (Oturan et al. 2010). Therefore, the optimal Fe\(^{2+}\) concentration for the removal of SAC was 0.2 mM.

\[
\text{Fe}^{2+} + \cdot \text{OH} \rightarrow \text{Fe}^{3+} + \text{OH}^- \quad (4-7)
\]
Effect of Fe$^{2+}$ (as catalyst) concentration on the removal of SAC with Pt (a) and BDD (b) anode ([SAC]$_0$ = 0.2 mM, [Na$_2$SO$_4$] = 50 mM, $I$ = 200 mA, pH$_0$ 3.0)

**4.3.4 Effect of applied current on the removal of SAC**

The applied current is another important parameter for the effectiveness of the electro-Fenton process. In order to investigate the effect of current intensity on the
oxidative degradation of SAC, several experiments were performed by varying applied current in the range of 50–500 mA at the optimal Fe^{2+} concentration of 0.2 mM as assessed above. The results were shown in Fig. 4-4 and Table 4-1. When Pt was used as anode, the removal rate of SAC increased as the applied current increasing. The electrolysis time for complete disappearance of SAC has been changed from >> 30 min for 50 mA to 30, 25, 20 and 15 min for 100, 200, 300 and 500 mA current intensity, respectively (Fig. 4-4a). The \( k_{\text{app}} \) was increased from 0.07 to 0.26 min^{-1} when applied current increased from 50 to 500 mA. The oxidation of SAC was accelerated by increasing the applied current because of progressively large production of \(^{\cdot}\)OH (Loaiza-Ambuludi et al. 2013).

As in the case of BDD anode, the removal rate of SAC increased significantly from 0.09 to 0.19 min^{-1} when applied current increased from 50 to 200 mA. However, it decreased slightly to 0.16 min^{-1} as applied current increased to 500 mA. At higher applied current than 200 mA, the 4e^- reduction of \( O_2 \) leading to the formation of \( H_2O \) (Eq. (4-8)) would compete with the formation of \( H_2O_2 \) (Eq. (4-9)) (Özcan et al. 2008a).

\[
\begin{align*}
O_2 + 4H^+ + 4e^- & \rightarrow 2H_2O \quad (4-8) \\
O_2 + 2H^+ + 2e^- & \rightarrow H_2O_2 \quad (4-9)
\end{align*}
\]

In addition, the progressive enhancement of other parasitic reactions at higher current, such as hydrogen evolution (Eq. (4-10)) at the cathode and oxygen evolution (Eq. (4-11)) at the anode (Yahya et al. 2014), also contributed to the lower SAC removal rate. It could be concluded that the comparatively better performance of BDD was more pronounced at lower current intensities.

\[
\begin{align*}
2H_2O + 4e^- & \rightarrow H_2 + 2OH^- \quad (4-10) \\
2H_2O & \rightarrow O_2 + 4H^+ + 4e^- \quad (4-11)
\end{align*}
\]
Fig. 4-4 Effect of applied current on the destruction kinetics of SAC with Pt (a) and BDD (b) anode

([SAC]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, [Na₂SO₄] = 50 mM, pH₀ 3.0)

4.3.5 Determination of the rate constant of reaction between SAC and 'OH

The absolute rate constant for the second order kinetics of the reaction between SAC and 'OH (kₘₐₓ,SAC) was determined by using the competitive kinetics method
Benzoic acid (BA) was employed as the standard competitor because the absolute rate constant of the reaction between BA and \( \cdot \text{OH} (k_{\text{abs,BA}}) \) is known as \( 4.30 \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \) (Özcan et al. 2008a; Oturan et al. 2010). Experiments of competitive kinetics were carried out in the presence of equal concentrations of SAC and BA (0.1 mM). The hydroxylation rate constant of SAC can be calculated according to Eq. (4-12) and determined from Fig. 4-5 to be \( k_{\text{abs,SAC}} = (1.85 \pm 0.01) \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \).

\[
\ln \left( \frac{[\text{SAC}]_0}{[\text{SAC}]_t} \right) = \left( \frac{k_{\text{abs,SAC}}}{k_{\text{abs,BA}}} \right) \ln \left( \frac{[\text{BA}]_0}{[\text{BA}]_t} \right)
\]

(4-12)

Interestingly, this rate constant value is the same as that reported by Toth et al. (Toth et al. 2012).

**Fig. 4-5** Determination of the absolute constant of reaction between SAC and \( \cdot \text{OH} \) using competition kinetics method by taking BA as standard competitors ([SAC]_0 = 0.1 mM, ([BA]_0 = 0.1 mM, [Fe^{2+}] = 0.2 mM, I = 50 mA, pH_0 3.0, [Na_2SO_4] = 50 mM)

**4.3.6 The effect of Fe^{2+} concentration and applied current on SAC mineralization**

The mineralization ability of SAC by electro-Fenton process was assessed from the TOC abatement over electrolysis time. The influence of the Fe^{2+} concentration in the Pt/carbon-felt and BDD/carbon-felt cells under the same condition of Fig. 4-3 was shown in Fig. 4-6.
Figure 4-6 indicated the optimal Fe\(^{2+}\) concentration for both Pt/carbon-felt and BDD/carbon-felt cells was 0.2 mM. When the Fe\(^{2+}\) concentration was higher than 0.2 mM, the percentage of scavenged hydroxyl radicals from Fe\(^{2+}\) ions increased, thus the side reactions between Fe\(^{2+}\) ions and \(^{\cdot}\)OH (Eq. (4-7)) became more significant (Özcan \textit{et al.} 2008a).

It can be seen by comparing Fig. 4-6a and Fig.4-6b that the mineralization of SAC was accelerated by the employ of BDD anode. At optimal conditions, BDD/carbon-felt cell achieved 93.3% mineralization in only 180 min, while this rate was about 70% for or Pt/carbon-felt cell at same time. A mineralization rate of 76.1% was reached by this cell at 360 min. BDD anode has the much higher O\(_2\) evolution overpotential (1.27 V) than Pt (0.27 V) (Oturan \textit{et al.} 2011). So the oxidative action of BDD(\(^{\cdot}\)OH) is much more efficient than Pt(\(^{\cdot}\)OH) (Martínez-Huittle and Brillas 2009). Moreover, the loosely bound BDD(\(^{\cdot}\)OH) formed at the anode surface (Eq. (4-6)) can readily react with organic pollutant due to the low adsorption ability of \(^{\cdot}\)OH on BDD, in contrast to the chemisorbed radicals Pt(\(^{\cdot}\)OH) which relatively strongly attached to the surface and less reactive (Oturan \textit{et al.} 2011; Dirany \textit{et al.} 2012).

![Graph showing TOC removal efficiency for different Fe\(^{2+}\) concentrations over time](image)
Fig. 4-6 Effect of Fe$^{2+}$ concentration on mineralization of SAC by electro-Fenton process with Pt (a) and BDD (b) anode ([SAC]$_0$ = 0.2 mM, [Na$_2$SO$_4$] = 50 mM, $I$ = 200 mA, pH$_0$ 3.0)

The effect of applied current on the mineralization of 0.2 mM SAC solution was investigated under the same condition as Fig. 4-4 and the results were shown in Fig. 4-7. When applied current raised from 50 to 200 mA, the TOC removal efficiency was increased from 64.2% to 76.1% in Pt/carbon-felt cell and from 88.9% to 96.2% in BDD/carbon-felt cell. However, the TOC removal efficiency decreased slightly for both anodes when current intensity further increased to 500 mA. As applied current increased, the higher electrogenerated H$_2$O$_2$ concentration was obtained and larger amounts of OH$^-$ was yielded from Fenton’s reaction, and then a greater mineralization efficiency can be achieved due to the simultaneous degradation of SAC and its byproducts (Dirany et al. 2012). But a current intensity higher than 200 mA would increase the extent of parasitic reactions (Eqs. (4-8), (4-10) and (4-11)) and decrease the mineralization efficiency of SAC (Mousset et al. 2014).
Fig. 4-7 Effect of applied current on mineralization of SAC by electro-Fenton process with Pt (a) and BDD (b) anode ([SAC]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, [Na₂SO₄] = 50 mM, pH₀ 3.0)

4.3.7 Identification and evolution of short-chain carboxylic acids

Generation of short chain carboxylic acids is expected from the oxidative breaking of aryl moieties of organic/cyclic oxidation intermediates. Experiments were performed
when initial SAC concentration was 0.2 mM, Fe$^{2+}$ concentration was 0.2 mM, Na$_2$SO$_4$ concentration was 50 mM, applied current was 200 mA and pH$_0$ was 3.0.

Oxalic, formic, and maleic acid at retention time ($t_R$) of 8.90, 16.03 and 11.33, respectively, were observed during electro-Fenton process. Since the concentration of maleic acid was detected in trace level, the changes of maleic acid concentration were not shown in Fig. 4-8. In Pt/carbon-felt cells, formic acid presented only in the first 20 min and then it undergo to the mineralization (Fig. 4-8a) (Özcan et al. 2008a). Oxalic acid showed the largest accumulation and the higher persistence due to its lower reactivity with $^\cdot$OH, which suggested that it can be generated from different ways as ultimate reaction intermediate (Dirany et al. 2010). In addition, oxalic acid was still existed with a concentration of 0.04 mM after a 360 min reaction. This relatively weak degradation of oxalic acid can be attributed to its high stability in the presence of ferric ions and its weaker reactivity toward $^\cdot$OH radicals ($k_2 = 2.8 \times 10^7$ M$^{-1}$ s$^{-1}$) (Oturan et al. 2008). This was also corresponding to the residual TOC remaining at the end of mineralization treatments in Pt/carbon-felt cells.

On the other hand, oxalic acid showed high persistence, but it disappeared at the end of the reaction in BDD/carbon-felt cell (Fig. 4-8b). This phenomenon was in agreement with the faster TOC removal in BDD/carbon-felt cells as described in section 4.3.6.
Evolution of short-chain carboxylic acids formed during electro-Fenton processes using Pt (a) and BDD (b) anodes ([SAC]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, [Na₂SO₄] = 50 mM, I = 200 mA, pH₀ = 3.0).

**Fig. 4-8** Evolution of short-chain carboxylic acids formed during electro-Fenton processes using Pt (a) and BDD (b) anodes ([SAC]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, [Na₂SO₄] = 50 mM, I = 200 mA, pH₀ = 3.0).

4.3.8 Evolution of SAC solution toxicity during electro-Fenton process

In order to determine the potential toxicity of SAC and its intermediates,
mineralization of a 0.2 mM SAC solution was examined at 200 mA constant current. The evolution of the luminescence inhibition as function of the electrolysis time for exposition time of 5 and 15 min are tested. Only the curves obtained at both Pt and BDD anodes after a 15-min exposure time were presented in Fig. 4-9, because the curves recorded after a 5-min exposure time were very similar. For both Pt and BDD anode, the toxicity increased significantly and reached the maximum luminescence inhibition peak at 30 min, indicating the formation of significantly more toxic by-products at the beginning of the treatment. The inhibition ratio decreased after 30 min, in relation to the destruction of aromatics and the increase of the less toxic by-products. These less toxic byproducts mainly included the carboxylic acids, which present a very low toxicity toward *V. fischeri* bacteria (Dirany *et al.* 2012).

![Graph showing inhibition of luminescence of *V. fischeri* bacteria during electro-Fenton processes with Pt and BDD anode (|SAC|₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, I = 200 mA, pH₀ = 3.0, [Na₂SO₄] = 50 mM)](image)

**Fig. 4-9** Inhibition of luminescence of *V. fischeri* bacteria during electro-Fenton processes with Pt and BDD anode (|SAC|₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, I = 200 mA, pH₀ = 3.0, [Na₂SO₄] = 50 mM)

### 4.4 Conclusions

It was demonstrated that electro-Fenton process with a Pt or BDD anode and a carbon felt cathode was a very effective method for the degradation of SAC in water.
The effect of Fe\textsuperscript{2+} concentration and applied current on the removal and mineralization of SAC was investigated. For both anodes, SAC could be completely degrade in less than 30 min. The oxidative degradation of SAC followed pseudo-first-order kinetics. Absolute rate constant of hydroxylation reaction of SAC was determined as (1.85 ± 0.01) × 10\textsuperscript{9} M\textsuperscript{-1} s\textsuperscript{-1} using competition kinetics method. The optimal conditions for SAC removal were SAC concentration 0.2 mM, Fe\textsuperscript{2+} concentration 0.2 mM, Na\textsubscript{2}SO\textsubscript{4} concentration 50 mM, applied current 200 mA and initial pH 3.0. The use of BDD anode yields a faster mineralization rate because of the higher oxidation power of BDD comparing with Pt. Short-chain aliphatic carboxylic acids such as oxalic, formic and maleic acids are identified as aliphatic by-products. The bacteria luminescence inhibition showed the toxicity of SAC solution increased at the first 30 min of electrolysis, and then it decreased after a 60 min reaction for both Pt and BDD anodes.

References
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Chapter 5 Electrochemical mineralization of sucralose in aqueous medium at ambient temperature by electro-Fenton process
5.1 Introduction

In this part, the effect of the Fe^{2+} concentration and applied current on the mineralization of sucralse (SUC) was examined. The effect of applied current on the mineralization efficiency was assessed. The aliphatic short-chain carboxylic acids and inorganic ions released during electro-Fenton process were monitored by ion-exclusion chromatography and ion chromatography (IC). The variation of toxicity of SUC solution and its intermediates was determined by Microtox method.

5.2 Materials and methods

5.2.1 Chemicals

Sucralose (1,6-dichloro-1,6-dideoxy-β-D-fructofuranosyl-4-chloro-4-deoxy-α-D-galactopyranoside, C_{12}H_{19}Cl_{3}O_{8}) was purchased from Sigma-Aldrich. The chemical structure and main characteristics of SUC were presented in Table 2-1. Analytical grade anhydrous sodium sulfate (supporting electrolyte) and ferrous sulfate heptahydrated (catalyst source) were obtained from Sigma-Aldrich and Acros Organics, respectively. Analytical grade carboxylic acids and other chemicals used for chromatographic analysis were purchased from Acros, Merck, Sigma, Riedel-de Haën and Fluka. Ultrapure water used for the preparation of the working solutions and HPLC eluting solutions was obtained from a Millipore Milli-Q (simplicity 185) system with resistivity > 18 MΩ cm at room temperature.

5.2.2 Electrochemical apparatus and procedures

Bulk experiments were conducted in an open, undivided and cylindrical glass cell of 250 mL capacity containing 220 mL SUC solution. Either a cylindrical Pt mesh (4.5 cm height, i.d. = 3.1 cm, Platecxis, France) or a 24 cm² thin-film BDD electrode (CONDIAS GmbH, Germany) was used as anode, and a 87.5 cm² piece of carbon felt (17.5 × 5 cm, Carbon-Lorraine, France) was used as cathode. In all electrolys, the
anode was centered in the cell, surrounded by the carbon felt which placed on the inner wall of the cell covering the totality of the internal perimeter. \( \text{H}_2\text{O}_2 \) was generated from the reduction of \( \text{O}_2 \) dissolved in the solution. The continuous saturation of oxygen at atmospheric pressure was assured by bubbling compressed air through a frit at about 0.5 L/min, starting 5 min before electrolysis. All the electrolytic trials were conducted under constant current conditions by a Hameg HM8040-3 triple power supply (Germany).

The degradation experiments were performed using 0.2 mM SUC solution with constant stirring by using a magnetic stirrer. 50 mM Na\(_2\)SO\(_4\) was added into the SUC solution as supporting electrolyte. A catalytic quantity of ferrous ion was added into the solution before the beginning of electrolysis. The initial pH (pH\(_0\)) of SUC solutions was measured with a CyberScan pH 1500 pH-meter (Eutech Instrument, USA) and set at 3.0 (±0.1), adjusting by the addition of 1 M sulfuric acid.

### 5.2.3 Analytical methods and procedures

The TOC of the samples withdrawn from treated solutions at different electrolysis times were determined by Shimatzu TOC-V\(_{\text{CSH}}\) analyser consisting of a non-dispersive infra-red absorption detector (NDIR) according to the 680 °C combustion catalytic oxidation method. Platinum was applied as catalyst to facilitate the combustion at 650 °C. The carrier gas was oxygen with a flow rate of 150 mL/min. The injection volume was 50 μL.

The short-chain carboxylic acids were identified and quantified by ion-exclusion HPLC using a L-7100 pump, a Merck Lachrom liquid chromatograph equipped with a Supelcogel H column (250 × 4.6 mm, 9 μm) and a L-7455 photodiode array detector at the wavelength of 220 nm. The mobile phase was 0.1% H\(_2\)SO\(_4\) solution and flow rate was fixed to 0.2 mL/min. Calibration curves were achieved using standard solutions of related carboxylic acids. The identification of the carboxylic acids was performed by the retention time (\( t_R \)) comparison and standard addition methods using standard substances.

Inorganic ions (Cl\(^-\)) released in the electro-Fenton process were monitored by ion
chromatography with a Dionex ICS-1000 Basic Ion Chromatography (IC) System equipped with an IonPac AS4A-SC (anion exchange) 250 × 4 mm column and fitted to a DS6 conductivity detector containing a cell heated at 35 °C under control through a Chromeleon SE software. The mobile phase was a mixture of 3.6 mM Na₂CO₃ and 3.4 mM NaHCO₃ solution with a flow rate of 2.0 mL/min. The volume of injections was 25 μL.

5.2.4 Toxicity measurements

The toxicity of SUC and its intermediates generated during the electrolyses were investigated via samples collected from treated solutions at different times. Experiments were performed with the bio-luminescence marine bacteria *V. fischeri* (Hach Lange France SAS) by Microtox method according to the international standard process (OIN 11348-3). Two values of the inhibition of the luminescence (%) were calculated after 5 and 15 min of exposure to samples at 15 °C.

5.3 Results and discussion

5.3.1 Effect of Fe²⁺ concentration on the mineralization of sucralose

Mineralization efficiency in both Pt/carbon-felt and BDD/carbon-felt cells was assessed by solution TOC (as equivalent of dissolved organic carbon) measurements during electrolysis of SUC solutions. The effect of Fe²⁺ concentration on the mineralization of SUC solution was monitored by using the Fe²⁺ concentration of 0.05, 0.1, 0.2, 0.3 and 0.5 mM. The SUC concentration was fixed at 0.2 mM, applied current was 200 mA and initial pH was 3.0. As can be observed in Fig. 5-1, the mineralization for SUC was enhanced by increasing the Fe²⁺ concentration from 0.05 to 0.2 mM for both Pt and BDD anode. Since the applied current was kept constant at 200 mA, it can be assumed that the production rate of H₂O₂ via the oxygen reduction would be identical for all the Fe²⁺ concentrations under this study (Oturan *et al.* 2010). On the basis of the same H₂O₂ production rate, a higher Fe²⁺ concentration could lead to the increase of
hydroxyl radicals generated in bulk of solution via Fenton's reaction (Eq. (5-1)). These hydroxyl radicals would react with SUC immediately, resulting in the increase of SUC mineralization.

\[ \text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{OH}^- + '\text{OH} \]  

(5-1)

However, further increasing the Fe\(^{2+}\) concentration to 0.5 mM, the mineralization efficiency of SUC declined evidently in both Pt/carbon-felt and BDD/carbon-felt cells. The negative influence of higher Fe\(^{2+}\) concentration might due to the role of Fe\(^{2+}\) as scavenger of hydroxyl radicals (Eq. (5-2)) which takes place with a large rate constant \((k = 3.20 \times 10^8 \text{ M}^{-1} \text{s}^{-1})\) (Sirés et al. 2007b; Oturan et al. 2010). Therefore, the value of 0.2 mM was chosen as the optimal Fe\(^{2+}\) concentration under these conditions and used in the following experiments.

\[ \text{Fe}^{2+} + '\text{OH} \rightarrow \text{Fe}^{3+} + \text{OH}^- \]  

(5-2)

By comparing Fig. 5-1a and Fig. 5-1b, it can be seen that the use of BDD anode lead to the acceleration of SUC mineralization. At optimal condition (Fe\(^{2+}\) concentration 0.2 mM), 96.1% of the initial TOC value was removed in BDD/carbon-felt cell after only 180 min reaction, while it takes 240 min for Pt/carbon-felt cell to achieve 94.2% of mineralization efficiency. This fact can be explained by following reasons: Firstly, a much higher quantities of BDD('OH) than Pt('OH) is provided when electrolysis is operated at the current within the water discharge region (Eq. (5-3) and (5-4)) (Brillas et al. 2005; Panizza and Cerisola 2005; Brillas et al. 2009).

\[ \text{BDD} + \text{H}_2\text{O} \rightarrow \text{BDD('OH)} + \text{H}^+ + e^- \]  

(5-3)

\[ \text{Pt} + \text{H}_2\text{O} \rightarrow \text{Pt('OH)} + \text{H}^+ + e^- \]  

(5-4)

Secondly, since BDD anode presents the highest O\(_2\)-overvoltage (1.27 V), which is much higher than Pt anode (0.27 V) (Martínez-Huitle and Brillas 2009; Oturan et al. 2011), the oxidative action of BDD('OH) is rather effective than Pt('OH) (Oturan et al. 2011). Thirdly, due to the low adsorption ability of 'OH on BDD, the physisorbed and labile BDD('OH) formed at the anode surface (Eq. (5-3)) can readily react with organic pollutant, in contrast to the chemisorbed radicals typically formed at Pt surface (Eq. (5-4)) which limited the oxidation ability of Pt('OH) (Oturan et al. 2011; Dirany et al.
Finally, BDD anode contributed to the degradation of all the byproducts, even the most refractory ones (Dirany et al. 2012).

**Fig. 5-1** Effect of Fe$^{2+}$ concentration on mineralization of SAC during electro-Fenton process using Pt (a) and BDD (b) anode ($[\text{SUC}]_0 = 0.2 \text{ mM}$, $[\text{Na}_2\text{SO}_4] = 50 \text{ mM}$, $I = 200 \text{ mA}$, pH$_0$ 3.0)
5.3.2 Effect of current intensity on the mineralization of SUC aqueous solutions

In order to investigate the effect of current intensity on the mineralization of SUC, several experiments were carried out by varying applied current in the range of 100–500 mA at the optimal Fe$^{2+}$ concentration of 0.2 mM as assessed above. The results were shown in Fig. 5-2. In both Pt/carbon-felt and BDD/carbon-felt cells, the mineralization efficiency of SUC solution increased significantly as the applied current increased from 100 to 200 mA. Taking Pt/carbon-felt cell as an example, when current intensity was 100 mA, the TOC removal efficiency was 69.7% in a 360 min treatment. Increasing the current to 200 mA resulted in 94.1% of mineralization efficiency in a 240 min reaction. The mineralization of SUC aqueous solution was accelerated by increasing the applied current because both Fe$^{2+}$ regeneration and H$_2$O$_2$ production would be promoted, and then large amount of 'OH would be produced (Boye et al. 2002; Lin et al. 2014). When the applied current increased from 200 to 500 mA, the mineralization efficiency of sucralose in both cells changed slightly. On the one hand, a higher current leads to the increase of parasitic reactions, such as hydrogen evolution (Eq.(5-5)) and oxygen evolution (Eq. (5-6)) reactions (Dirany et al. 2010; Dirany et al. 2012).

\[
\begin{align*}
2\text{H}_2\text{O} + 2e^- & \rightarrow \text{H}_2 + 2\text{OH}^- \quad (5-5) \\
2\text{H}_2\text{O} \rightarrow \text{O}_2 + 4\text{H}^+ + 4e^- \quad (5-6)
\end{align*}
\]

On the other hand, when the current density was further increased, excessive hydroxyl radicals would be consumed via the following side reactions (Eqs. (5-7) and (5-8)) (Wu et al. 2012; Lin et al. 2014). As a result, 200 mA was selected as the optimal current value and used in the remaining experiments.

\[
\begin{align*}
\text{H}_2\text{O}_2 + \cdot\text{OH} & \rightarrow \text{HO}_2^- + \text{H}_2\text{O} \quad (5-7) \\
\text{HO}_2^- + \cdot\text{OH} & \rightarrow \text{H}_2\text{O} + \text{O}_2 \quad (5-8)
\end{align*}
\]

It also be observed from Fig. 5-2 that SUC solutions could be almost totally mineralized after 240 min of electrolysis at optimal conditions (Fe$^{2+}$ concentration 0.2 mM and applied current 200 mA) in both Pt/carbon-felt and BDD/carbon-felt cells, indicating that electro-Fenton process with the use of Pt or BDD anode and a piece of carbon-felt cathode was quite effective for the mineralization of SUC in aqueous
medium.

![Graph](image)

**Fig. 5-2** Effect of applied current on mineralization of SAC by electro-Fenton process with Pt (a) and BDD (b) anode ([SUC]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, [Na₂SO₄] = 50 mM, pH₀ 3.0)

### 5.3.3 Mineralization current efficiency (MCE)

The mineralization capacity of the process can be expressed by MCE which can be

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determined from the following equation (Eq. (5-9)) (Skoumal et al. 2008; Brillas et al. 2009; Oturan et al. 2011):

\[
MCE = \frac{nFV_s \Delta (\text{TOC})_{\text{exp}}}{4.32 \times 10^7 m t}
\]  

(5-9)

where \(n\) is the number of electrons consumed per SUC molecule according to Eq. (5-10) (= 48), \(F\) is the Faraday constant (= 96,487 C/mol), \(V_s\) is the solution volume (L), \(\Delta (\text{TOC})_{\text{exp}}\) is the experimental TOC decay (mg/L), \(4.32 \times 10^7\) is the conversion factor to homogenize units (= 3600 s/h \times 12,000 mg of C/mol), \(m\) is the number of carbon atoms of sucralose (= 12), \(I\) is the applied current (A) and \(t\) is the given treatment time (h).

\[
\text{C}_{12}\text{H}_{19}\text{Cl}_3\text{O}_8 + 16\text{H}_2\text{O} \rightarrow 12\text{CO}_2 + 51\text{H}^+ + 3\text{Cl}^- + 48\text{e}^-
\]  

(5-10)

By comparing Fig. 5-3(a) and Fig. 5-3(b), it can be seen that the MCE in Pt/carbon-felt cell was lower at all the chosen time than that in BDD/carbon-felt cell. This was attributed to the low oxidation power of Pt anode (Oturan et al. 2011).

Figure 5-3 indicated that the MCE values decreased with rising applied current from 100 to 500 mA in both Pt/carbon-felt and BDD/carbon-felt cells. For example, the MCE values after 60 min of electrolysis in BDD/carbon-felt cell for 100, 200, 300 and 500 mA were 31.49%, 20.17%, 14.40% and 8.90%, respectively. This was related to the involvement of some parasitic reactions and particularly the hydrogen gas evolution reaction (Eq. (5-5)), which competed with the formation of \(\text{H}_2\text{O}_2\) (Eq.(5-11)) (Özcan et al. 2008).

\[
\text{O}_2 + 2\text{H}^+ + 2\text{e}^- \rightarrow \text{H}_2\text{O}_2
\]  

(5-11)

Figure 5-3 also showed that the MCE values decreased continuously from the beginning to the end of the electrolysis for every trial in both cells. This was due to the gradual formation of intermediates such as carboxylic acids that are more difficult to destroy by BDD(’OH) and also due to mass transport limitations because of the presence of small concentration of organic matter (Sirés et al. 2007a; Özcan et al. 2008).
Fig. 5-3 Dependence of mineralization current efficiency calculated from Eq. (5-10) on the electrolysis time for the experiments reported in: (a) Fig. 5-2a and (b) Fig. 5-2b

5.3.4 Identification and evolution of short-chain carboxylic acids

To identify the generated carboxylic acids released in the electro-Fenton process,
0.2 mM SUC solution aqueous solution was treated when Fe$^{2+}$ concentration was 0.2 mM, Na$_2$SO$_4$ concentration was 50 mM, applied current was 200 mA and pH$_0$ was 3.0.

The concentration of carboxylic acids was monitored by ion-exclusion HPLC and the results were shown in Fig. 5-4. In both Pt/carbon-felt and BDD/carbon-felt cells, oxalic, pyruvic, formic and glycolic acid at retention time ($t_R$) 8.90, 11.57, 16.03 and 14.53, respectively, were observed. As shown in Fig. 5-4, formic acid was observed as soon as the electrolysis started in both cells. Its concentration reaches the maximum value of 0.090 mM after a 20 min reaction in Pt/carbon-felt cell and 0.096 mM after only 10 min in BDD/carbon-felt cell. Then it decreased gradually to be completely mineralized after 180 min and 60 min in Pt/carbon-felt and BDD/carbon-felt cells, respectively. Pyruvic acid was also generated at the beginning of the electrolysis and followed the accumulation-destruction cycles in two cells. It disappeared after 120 min electrolysis in both cells. The accumulation of glycolic acid was a little slower and appeared after 7 min of electrolysis. Oxalic acid showed the largest accumulation and the higher persistence in both cells due to its high stability in the presence of ferric ions and its weaker reactivity toward •OH radicals ($k_2 = 2.8 \times 10^7$ M$^{-1}$ s$^{-1}$) (Oturan et al. 2008). This suggested that oxalic acid would be generated from different ways as ultimate reaction intermediates.

Moreover, all the carboxylic acids were vanished after a 360 min electrolysis in Pt/carbon-felt cell and a 240 min electrolysis in BDD/carbon-felt cell, which was corresponding to the residual TOC remaining at the end of mineralization treatments (Fig. 5-2a and Fig. 5-2b).
Fig. 5-4 Evolution of short-chain carboxylic acids formed during electro-Fenton processes with Pt (a) and BDD (b) anodes ([SUC]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, [Na₂SO₄] = 50 mM, I = 200 mA, pH₀ = 3.0).

5.3.5 Identification and evolution of chloride ions

Upon bond cleavage of the sucralose molecules, the Cl atoms are released into the
solution as inorganic ions and the concentration of Cl\(^-\) was shown in Figure 5-5. When Pt was used as anode, a progressive accumulation of Cl\(^-\) was observed, until reaching the expected maximum value of 0.6 mM. In contrast, the release of Cl\(^-\) was increased significantly and reached a maximum concentration of 0.544 mM after 30 min, whereupon it decreased until a final value of 0.063 mM at 360 min. This phenomenon could be explained by the oxidation of Cl\(^-\) to Cl\(_2\) and/or ClO\(^-\)/HClO, ClO\(_3^-\), and ClO\(_4^-\) by BDD(•OH) and/or •OH in the bulk (Randazzo et al. 2011; Dirany et al. 2012).

\[\text{Cl}\] (mM) \hspace{1cm} \text{Time (min)}
\begin{tabular}{c|c|c|c|c|c|c}
0 & 50 & 100 & 150 & 200 & 250 & 300 & 350 \\
0.0 & 0.1 & 0.2 & 0.3 & 0.4 & 0.5 & 0.6 & \\
\end{tabular}

\[\text{BDD} \quad \text{Pt}\]

**Fig. 5-5** Evolution of chloride ions released during electro-Fenton processes with Pt (a) and BDD (b) anode ([sucralose]\(_0\) = 0.2 mM, [Fe\(^{2+}\)] = 0.2 mM, [Na\(_2\)SO\(_4\)] = 50 mM, \(I = 200\) mA, pH\(_0\) 3.0)

### 5.3.6 The evolution of toxicity with reaction time

In order to monitored the potential toxicity of sucralose and its intermediates, 0.2 mM sucralose was electrolyzed in the existence of 0.2 mM Fe\(^{2+}\) at 200 mA using Microtox method for both Pt/carbon-felt and BDD/carbon-felt cells. The percentage of bacteria luminescence inhibition versus the electrolysis time after 15 min exposure times of *V. fischeri* luminescent bacteria to the sucralose solutions was shown in Fig. 5-6.

When Pt was used as anode, there were several peaks due to the degradation primary intermediates and formation to secondary/tertiary intermediates that were more
toxic than the previous sucralose solution in the first 60 min electrolysis (Feng et al. 2014). The other peak appeared at 120 min, indicating the generation of other toxic byproducts. From 120 to 240 min, the inhibition percentage decreased sharply. The toxicity was much lower than the untreated sucralose solution at 240 min, in relation to the increase of the less toxic byproducts, such as carboxylic acids, which present a very low toxicity toward *V. fischeri* bacteria (Dirany et al. 2012). Consequently, the toxicity kept stable from 240 min to the end of the electrolysis.

The curves obtained for BDD anode showed a similar behavior at the beginning of the electrolysis as Pt anode. Several peaks were obtained, indicating the formation of some intermediates that were more toxic than the previous sucralose solution in a 60 min reaction. The secondary peak appeared at 40 min and showed the secondary intermediates still more toxic than the original sucralose solution. The secondary peak which appeared at 240 min was much lower than the primary peak, showing the toxicity of secondary intermediates were much lower than the primary byproducts. After 240 min, the inhibition percentage decreased significantly and was much lower than the untreated sucralose solution at the end of the electrolysis, showing the disappearance of toxic intermediate products.

![Graph](image)

**Fig. 5-6** Evolution of the inhibition of marine bacteria, *Vibrio fischeri* luminescence (Microtox method) during electro-Fenton processes with Pt (a) and BDD (b) anode ([sucralose]₀ = 0.2 mM, [Fe²⁺] = 0.2 mM, [Na₂SO₄] = 50 mM, I = 200 mA, pH₀ 3.0)
5.4 Conclusions

Electro-Fenton process using a carbon-felt cathode and a Pt or BDD anode was successfully applied to the mineralization of sucralose in aqueous solutions in this study. The optimal conditions for the mineralization of 0.2 mM sucralose solution were Fe$^{2+}$ concentration 0.2 mM, Na$_2$SO$_4$ concentration 50 mM, applied current 200 mA and initial pH 3.0. At optimal conditions, sucralose could be totally mineralized in 240 min for BDD anode and 360 min for Pt anode. Short-chain aliphatic carboxylic acids such as oxalic, pyruvic, formic and glycolic are identified as aliphatic byproducts during the electro-Fenton process. Cl atoms were found in the structure of sucralose and were converted to Cl$^-$. The bacteria luminescence inhibition showed the toxicity of sucralose solution increased at the beginning of electrolysis, and then it declined until much lower than the original sucralose solution at the end of the reaction.
References


Oturan, M.A., Pimentel, M., Oturan, N. and Sirés, I. (2008) Reaction sequence for the


Chapter 6 Decolorization of Orange II in water by 

electro/\alpha-FeOOH/PDS process
6.1 Introduction

In this part, peroxydisulfate (PDS), which has the similar structure of O-O bond contained in H₂O₂ (Hou et al. 2012), was activated by goethite to generate sulfate radicals and remove Orange II from water in a electrochemical (EC) reactor (electro/α-FeOOH/peroxydisulfate process).

The traditional one-factor-at-a-time approach has been widely used to study the effects of various factors in most experiments. Experimental factors are varied one at a time, with the remaining factors being held constantly (Zhang et al. 2011). This method fails to investigate the interactive influences of different variables. When a combination of several independent variables and their interactions affect decolorization efficiency, response surface methodology (RSM) is effective for optimizing the operating parameters in multivariable systems (Gong et al. 2010; Zhang et al. 2011). As one of the RSM designs, Box-Behnken design (BBD) is known as a modified central composite experimental design (Ay et al. 2009; Zhang et al. 2011). BBD is an independent, rotatable quadratic design with no embedded factorial or fractional factorial points where the variable combinations are at the midpoints of the edges of the variable space and at the center. In addition, BBD requires fewer runs than other RSM designs with the same number of variables. For example, only 15 runs are needed for a three-parameter experimental design. As a result, BBD was applied in this study to investigate the effect of electric current, α-FeOOH concentration and PDS concentration on the results of the pseudo-first-order decolorization rate constant of Orange II (a kind of widely used zao dyes) in electro/α-FeOOH/peroxydisulfate process.

6.2 Materials and methods

6.2.1 Chemicals

Orange II (4-(2-hydroxynaphthylazo) benzenesulfonic acid sodium salt) was purchased from Sinopharm Chemical Reagent Co., Ltd. (China) and used without further purification. The Chemical structure and main characteristics of Orange II were
presented in Table 6-1. α-FeOOH was obtained from Sigma-Aldrich. Stock solution was prepared by dissolving a selected amount of Orange II in deionized water and the initial concentration (C₀) was fixed at 50 mg/L. 50 mM Sodium sulfate was added as a supporting electrolyte.

Table 6-1 Chemical structure and main characteristics of Orange II

<table>
<thead>
<tr>
<th>Name</th>
<th>Orange II</th>
<th><img src="image" alt="chemical_structure" /></th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical structure</td>
<td></td>
<td><img src="image" alt="chemical_structure" /></td>
</tr>
<tr>
<td>Molecular formula</td>
<td>C₁₆H₁₁N₂NaO₄S</td>
<td></td>
</tr>
<tr>
<td>CAS number</td>
<td>633-96-5</td>
<td></td>
</tr>
<tr>
<td>Molecular weight</td>
<td>350.32</td>
<td></td>
</tr>
<tr>
<td>λ_{max}(nm)</td>
<td>485</td>
<td></td>
</tr>
<tr>
<td>Water solubility (g/L)</td>
<td>116</td>
<td></td>
</tr>
</tbody>
</table>

6.2.2 Electrochemical apparatus and procedures

Batch experiments were conducted in an undivided electrolytic reactor (glass beaker) containing 400 mL of Orange II solution. Electrolyses were performed at constant current controlled by a direct current (DC) power supply (Model WYK-305) from Yangzhou Jintong Source, Co., Ltd. (China). One 5 × 11.9 cm plate anode (Ti/RuO₂-IrO₂) and one plate cathode (stainless steel) of the same dimensions were arranged parallel to each other at a distance of 4.0 cm. Prior to the electrolysis, the initial pH (pH₀) of Orange II solutions were measured with a Mettler-Toledo FE20 pH-meter (Mettler-Toledo Instruments Co., Ltd., Shanghai) and adjusted by sulfuric acid and sodium hydroxide. A magnetic stirrer (model 78-1, Hangzhou Instrument Motors Factory, China) was applied to achieve the transport of electroactive matter toward the electrode and ensure homogeneity throughout the reaction. The DC power supply was initiated when PDS solutions and α-FeOOH were added to the electrolytic cell. Samples were taken from electrolytic reactor at each predetermined time interval.
These samples were filtered through 0.22 μm membranes (Millipore Co.) before analysis and then mixed with the same volume of methanol to quench the reaction before analysis.

6.2.3 Analytical methods and procedures

The absorbance of Orange II was measured at maximum absorption wavelength of 485 nm using a Rayleigh UV-9100 spectrophotometer (Rayleigh Co., China). The UV-visible spectrum from 200 to 800 nm was achieved by a spectrophotometer (Shimadzu, UV-1700). The concentration of residual PDS was measured by an iodometric titration method (Wahba et al. 1959). Total organic carbon (TOC) analyses were carried out with non dispersive infra red absorption detector (NDIR) using an Analytikjena multi N/C 3100 analyzer. The carrier gas was oxygen with a flow rate of 150 mL/min. The structure of α-FeOOH was characterized by X-ray diffraction (XRD). XRD patterns are collected on a X’Pert Pro Xray diffractometer (PANalytical B.V., Holland), using a Cu Kα radiation (λ = 1.5406 Å). The X-ray photoelectron spectroscopy (XPS) was operated on a ESCALAB 250Xi spectrometer (Thermo fisher, USA) which equipped with AlKα X-ray source. The XPS data analysis was performed using the XPSPeak4.1 program with a symmetric Gaussian–Lorentzian sum function and Shirley background subtraction.

6.2.4 Box-Behnken design (BBD)

Box-Behnken statistical experiment design and response surface methodology were employed to investigate the effects of the three independent variables on the response functions. The independent variables were electric current (X₁), α-FeOOH dosage (X₂) and PDS concentration (X₃). The low, center and high levels for each variable are designed as -1, 0 and +1, respectively as illustrated in Table 6-2. The experimental levels for each variable were determined based results from preliminary experiments of operating parameters. The dependent variable or objective function was the pseudo-first-order decolorization rate constant (Y).
Table 6-2 Experimental range and levels of the independent variables

<table>
<thead>
<tr>
<th>Variables</th>
<th>Symbol</th>
<th>-1</th>
<th>0</th>
<th>+1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electric current (A)</td>
<td>X₁</td>
<td>0.2</td>
<td>0.5</td>
<td>0.8</td>
</tr>
<tr>
<td>α-FeOOH dosage (g/L)</td>
<td>X₂</td>
<td>0.2</td>
<td>0.5</td>
<td>0.8</td>
</tr>
<tr>
<td>PDS concentration (g/L)</td>
<td>X₃</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

Each independent variable was coded as $X_i$ and determined by the following equation:

$$ X_j = \frac{x_j - x_0}{\Delta x_j} $$  \hspace{1cm} (6-1)

where $X_i$ is dimensionless code value of the $i^{\text{th}}$ independent variable, $x_i$ is the uncoded value of the $i^{\text{th}}$ independent variable, $x_0$ is the uncoded $i^{\text{th}}$ independent variable at the center point and $\Delta x_i$ is the step change value between low level (-1) and high level (+1).

As can be seen in Table 6-3, the total number of experiments in this study was 15 based on 3 levels and a 3 factor experimental design, with three replicates at the center of the design for estimation of a pure error sum of squares. Experimental data from the BBD could be analyzed and fitted to a second-order polynomial model using Design Expert 8.0 software:

$$ Y = \beta_0 + \sum \beta_i X_i + \sum \beta_{ii} X_i^2 + \sum \beta_{ij} X_i X_j $$  \hspace{1cm} (6-2)

where $Y$ is the response and $X_i$ and $X_j$ are the independent variables, square effects and interaction effects; $\beta_i$, $\beta_{ij}$ and $\beta_{ii}$ are the linear coefficients, interaction coefficients and squared coefficients; $\beta_0$ is the intercept parameter.

Table 6-3 Design matrix in coded units and the experiments

<table>
<thead>
<tr>
<th>Stand no.</th>
<th>$X_1$</th>
<th>$X_2$</th>
<th>$X_3$</th>
<th>$k$ (min⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Observed</td>
</tr>
<tr>
<td>1</td>
<td>-1</td>
<td>-1</td>
<td>0</td>
<td>0.0079</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>-1</td>
<td>0</td>
<td>0.021</td>
</tr>
<tr>
<td>3</td>
<td>-1</td>
<td>1</td>
<td>0</td>
<td>0.011</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0.022</td>
</tr>
<tr>
<td>5</td>
<td>-1</td>
<td>0</td>
<td>-1</td>
<td>0.011</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>0</td>
<td>-1</td>
<td>0.017</td>
</tr>
<tr>
<td>7</td>
<td>-1</td>
<td>0</td>
<td>1</td>
<td>0.011</td>
</tr>
<tr>
<td>8</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0.021</td>
</tr>
</tbody>
</table>
6.3 Results and discussion

6.3.1 Decolorization of Orange II under different systems

In order to investigate the decolorization efficiency of Orange II under different oxidation systems, 50 mg/L Orange II solution was treated for 120 min by PDS alone, α-FeOOH alone, α-FeOOH/PDS process, EC alone, EC/α-FeOOH process, EC/PDS process and EC/α-FeOOH/PDS process. The results were shown in Fig. 6-1. It can be seen from Fig. 6-1a that little decolorization occurred when Orange II was treated by PDS alone, because PDS is stable at ambient temperature and its oxidation power is limited ($E^0 = 2.01$ V) (Zhao et al. 2010; Wu et al. 2012b), which could hardly degrade Orange II. What's more, the remaining percentage of PDS after 120 min reaction in the PDS alone process was 98.2% (Fig. 6-1b), indicating almost no PDS decomposed in this case. Negligible color removed was also observed when α-FeOOH alone was applied, indicating the effect of adsorption of Orange II decolorization was not obvious under the condition investigated. Orange II was hardly removed in α-FeOOH/PDS process and the remaining percentage of PDS was 95.7%, because PDS can not be activated by Fe(III) on the surface of α-FeOOH. When treated by EC alone, 51.8% of 50 mg/L Orange II was decolorized because a certain amount of hydroxyl radicals ($\text{M}^\cdot\text{OH}$) was formed as intermediate of water discharge on DSA anode (M) (Eq. (9)) (Özcan et al. 2009).

$$\text{M(H}_2\text{O}) \rightarrow \text{M}^\cdot\text{OH} + \text{H}^+ + \text{e}^-$$

Adding 0.5 g/L α-FeOOH to the electrolytic reactor (EC/α-FeOOH process) could
not improve the decolorization efficiency of Orange II. The decolorization efficiency was 57.2% after 120 min reaction in EC/α-FeOOH process. The decolorization efficiency improved significantly to 81.4% in EC/PDS process, which might be attributed to sulfate radicals produced via Eq. (6-4). 50.0% of PDS was decomposed in this case (Fig. 6-1b).

\[ \text{S}_2\text{O}_8^{2-} + e^- \rightarrow \text{SO}_4^{2-} + \text{SO}_4^{2-} \]  \hspace{1cm} (6-4)

The decolorization efficiency of Orange II was 90.9% and the remaining percentage of PDS was 38.7% in the process of EC/α-FeOOH/PDS, indicating Fe(II) can be generated by the cathodic reduction of Fe(III) (Eq. (6-5)), and then more sulfate radicals can formed by Fe(II) activating PDS reaction (Eq. (6-6)).

\[ \equiv\text{Fe(III)} + e^- \rightarrow \equiv\text{Fe(II)} \]  \hspace{1cm} (6-5)

\[ \equiv\text{Fe(II)} + \text{S}_2\text{O}_8^{2-} \rightarrow \equiv\text{Fe(III)} + \text{SO}_4^{2-} + \text{SO}_4^{2-} \]  \hspace{1cm} (6-6)
Fig. 6-1 Decolorization of Orange II (a) and remaining percentage of PDS (b) after 120 min of reaction under different systems \( (C_0 = 50 \text{ mg/L}, [\alpha-\text{FeOOH}] = 0.5 \text{ g/L}, [\text{PDS}] = 2 \text{ g/L}, I = 0.5 \text{ A}, \) \( \text{pH}_0 = 3.0, [\text{Na}_2\text{SO}_4] = 50 \text{ mM}) \)

In order to investigate the mechanism of Orange II decolorization in EC/\( \alpha \)-FeOOH/PDS process, the catalyst \( \alpha \)-FeOOH was characterized by XRD and XPS. The XRD analysis was applied to define the structure of fresh \( \alpha \)-FeOOH and the obtained patterns were shown on Fig. 6-2. The peaks at the angle of 20 21.223°, 33.241°, 34.700°, 36.649°, 39.984°, 41.186°, 50.613°, 53.237° and 59.023° are specific of \( \alpha \)-FeOOH (Jaiswal et al. 2013).
Fig. 6-2 XRD pattern of $\alpha$-FeOOH (Reference X-ray lines belonging to $\alpha$-FeOOH (JCPDS card n° 29-0713) are given at the bottom of the figure)

XPS was used to applied to observe the surface properties of $\alpha$-FeOOH before and after reaction. The results were showed in Fig. 6-3. The domain peak for O 1s was at 531.6 eV. The O 1s peak of $\alpha$-FeOOH is shifted significantly higher (0.4 eV) than that of $\alpha$-Fe$_2$O$_3$ (531.2 eV) (Guo et al. 2010), indicating the proton on the hydroxyl oxygen has some interaction with the O$_2^-$ oxygen (McIntyre and Zetaruk 1977). Fig. 6-3a showed the O 1s peak of $\alpha$-FeOOH did not change obviously after the reaction. So the properties of oxygen on the surface of $\alpha$-FeOOH did not change during the EC/$\alpha$-FeOOH/PDS process. The peaks of Fe 2p spectrum at 711, 719, and 725 eV represented the binding energies of Fe 2p$_{3/2}$, shake-upsatellite Fe 2p$_{3/2}$, and Fe 2p$_{1/2}$, respectively (Nie et al. 2009). As can be seen from Fig. 6-3b, there was no Fe (II) on the surface of $\alpha$-FeOOH before the reaction. However, The ratio of Fe(II)/Fe(III) on the surface of $\alpha$-FeOOH adsorbent increased to 0.80 after 120 min reaction. This implied more Fe(II) was generated on the surface of $\alpha$-FeOOH particles during EC/$\alpha$-FeOOH/PDS process (Eq. (6-5)).
Based on the results and the previous studies, (Liang et al. 2007; Özcan et al. 2009; Lin et al. 2013), the mechanism of EC/α-FeOOH/PDS process was shown in Fig. 6-4. Fe(II) was generated by the reduction and transformation of Fe(III) on the surface of α-FeOOH due to the direct electron transfer (Eq. (6-5)), and then Fe(II) activated PDS
to generate sulfate radicals (Eq. (6-6)). Meanwhile, hydroxyl radicals were generated both by water discharge on DSA anode (Eq. (6-3)) and the reaction between sulfate radicals and \( \text{H}_2\text{O} \) or \( \text{OH}^- \) (Eqs. (6-7) and (6-8)) (Liang et al. 2007; Saien et al. 2011). Orange II was degraded by both sulfate radicals and hydroxyl radicals.

\[
\text{SO}_4^{2-} + \text{H}_2\text{O} \rightarrow \text{H}^+ + \text{SO}_4^{2-} + \cdot\text{OH} \quad (6-7)
\]

\[
\text{SO}_4^{2-} + \text{OH}^- \rightarrow \text{SO}_4^{2-} + \cdot\text{OH} \quad (6-8)
\]

**Fig. 6-4** Proposed mechanism of EC/\(\alpha\)-FeOOH/PDS process

### 6.3.2 Effect of initial pH

The effect of pH on degradation of Orange II was investigated when initial Orange II concentration was 50 mg/L, \(\alpha\)-FeOOH dosage was 0.5 g/L, PDS concentration was 2 g/L, \(\text{Na}_2\text{SO}_4\) solution concentration was 50 mM and the electrical current was 0.5 A. The initial pH values investigated were 3, 7 and 10. As it indicated in Fig. 6-5, no significant effect of pH on the decolorization efficiency was observed. The final efficiency was 90.9%, 92.1% and 89.1% at initial pH 3.0, 7.0 and 10.0, respectively. It is reported that sulfate radicals could react with \(\text{H}_2\text{O}\) or \(\text{OH}^-\) to produce hydroxyl radicals (Eqs. (6-7) and (6-8)) (Liang et al. 2007; Saien et al. 2011). Either \(\text{H}^+\) could be generated or \(\text{OH}^-\) could be consumed, which led to the reduction of solution pH in EC/FeOOH/PDS process. Moreover, the reaction of hydroxyl radical and sulfate radical
(Eq. (6-9)) as well as the decomposition of PDS in water (Eq. (6-10)) would generate HSO$_4^–$. The dissociation of HSO$_4^–$ (Eq. (6-11)) would further release H$^+$ and decrease the solution pH (Kusic et al. 2011; Wu et al. 2012b).

\[
\begin{align*}
\text{SO}_4^{2–} + \cdot \text{OH} & \rightarrow \text{HSO}_4^{–} + \frac{1}{2}\text{O}_2 \quad (6-9) \\
\text{S}_2\text{O}_8^{2–} + \text{H}_2\text{O} & \rightarrow 2\text{HSO}_4^{–} + \frac{1}{2}\text{O}_2 \quad (6-10) \\
\text{HSO}_4^{–} & \rightarrow \text{SO}_4^{2–} + \text{H}^+ \quad (6-11)
\end{align*}
\]

Therefore, the final pH was almost the same value (about 2.5) and the decolorization of Orange II tended to be operated under similar conditions as the reaction proceeded. This indicated that the EC/α-FeOOH/PDS process could be successfully applied at a wide range of initial pH. Since pH of most actual textile effluent was neutral, pH 7.0 was chosen as the optimal initial value used in the following experiments.

**Fig. 6-5** The effect of initial pH value on the degradation of Orange II ($C_0 = 50$ mg/L, [α-FeOOH] = 0.5 g/L, [PDS] = 2 g/L, $I = 0.5$ A, [Na$_2$SO$_4$] = 50 mM)
6.3.3 Response surface analysis

6.3.3.1 Main variables

The full factorial BBD with three factors in three levels as well as results of Orange II decolorization rate constant for each run are listed in Table 6-3. Based on Table 6-3, the main effects plot and the interaction plots for Orange II decolorization rate constant at 120 min were developed. The main effects plot illustrated that the effects of electrical current, α-FeOOH dosage and PDS concentration on Orange II decolorization rate constant were positive (Fig. 6-6). This means that the greater removal efficiency could be achieved at high level (+1) of each factor than that at low level (–1) of the factor. Among the variables, the influence of electrical current was the greatest, and the second one was the PDS concentration. The influence of α-FeOOH dosage was softer than the other two variables at 120 min.

Fig. 6-6 Main effects plot for Orange II decolorization rate constant: Applied current (A); α-FeOOH dosage (B); PDS concentration (C)

6.3.3.2 Interaction between variables

Fig. 6-7 illustrated interaction plots showing the existence or not of interaction
among the factors. One factor was fixed at its high (+1) or low (−1) level while the other was investigated (Domínguez et al. 2010). As can be seen in Fig. 6-4, the curves of interaction between α-FeOOH dosage and PDS concentration are across, indicating the interaction between two variables is significant. The curves of interaction between electrical current and PDS concentration are tending to cross. It may assume that the interaction between two variables is significant. The interaction effect between electrical current and α-FeOOH dosage are less significant than that between α-FeOOH dosage and PDS concentration. This was also confirmed by the high probability value ((Prob > F) > 0.1) through analysis of variance (ANOVA).

![Interaction plots for Orange II decolorization rate constant: Applied current (A); α-FeOOH dosage (B); PDS concentration (C)](image)

**Fig. 6-7** Interaction plots for Orange II decolorization rate constant: Applied current (A); α-FeOOH dosage (B); PDS concentration (C)

### 6.3.3.3 Fitting model and analysis of variance

In order to justify the significance and adequacy of the model, the ANOVA analysis is required. The value of (Prob > F) was used as a tool to check the significant of each model terms. The smaller the value of (Prob > F) means the more significant of the corresponding model term. Generally, the value of (Prob > F) over 0.1 implies that the model term is insignificant (Zhang et al. 2010). Based on the results shown in Table 6-4, RSM model of second order polynomial equations was determined as follows,
\[ Y = 0.021 + 4.85 \times 10^{-3} X_1 + 5.38 \times 10^{-4} X_2 + 1.86 \times 10^{-3} X_3 - 5.50 \times 10^{-4} X_1X_2 + 7.50 \times 10^{-4} X_1X_3 + 2.38 \times 10^{-3} X_2X_3 - 3.93 \times 10^{-3} X_1^2 - 1.55 \times 10^{-3} X_2^2 - 1.65 \times 10^{-3} X_3^2 \]

(6-12)

The coefficients of determination \( R^2 \) gave the proportion of the total variation in the response variable explained. The \( R^2 \) obtained in this work (0.9507) ensured a satisfactory adjustment of the quadratic model to the experimental data. The coefficient of variation (CV) is a measure expressing standard deviation as a percentage of the mean. It illustrates the extent to which the data were dispersed. The lower values of CV give better reproducibility (Ahmad et al. 2005; Zinatizadeh et al. 2006). In this work, the CV value (10.6%) was within the acceptable range (0.5–13.5%) (Zinatizadeh et al. 2006; Wu et al. 2012a). In addition, adequate precision was used to assess the signal to noise ratio. The value of adequate precision greater than 4 is considered to give accurate prediction (Körbahtli 2007). Therefore, the adequate precision value of 9.126 indicated adequate signals for the models to be used to navigate the design space.

Table 6-4 ANOVA for response surface quadratic model

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>Sum of squares</th>
<th>Degree of freedom</th>
<th>Mean square</th>
<th>F-value</th>
<th>p-value</th>
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<td>Model</td>
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<td>9</td>
<td>( 3.48 \times 10^{-4} )</td>
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<td>0.0089</td>
</tr>
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<td>( X_1 )</td>
<td>( 1.88 \times 10^{-4} )</td>
<td>1</td>
<td>( 1.88 \times 10^{-4} )</td>
<td>57.98</td>
<td>0.0006</td>
</tr>
<tr>
<td>( X_3 )</td>
<td>( 2.78 \times 10^{-5} )</td>
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<td>( 2.78 \times 10^{-5} )</td>
<td>8.55</td>
<td>0.0329</td>
</tr>
<tr>
<td>( X_2X_3 )</td>
<td>( 2.26 \times 10^{-5} )</td>
<td>1</td>
<td>( 2.26 \times 10^{-5} )</td>
<td>6.95</td>
<td>0.0462</td>
</tr>
<tr>
<td>( X_1^2 )</td>
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<td>( 3.25 \times 10^{-6} )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lack of fit</td>
<td>( 1.52 \times 10^{-5} )</td>
<td>3</td>
<td>( 5.08 \times 10^{-6} )</td>
<td>10.30</td>
<td>0.0898</td>
</tr>
<tr>
<td>Pure error</td>
<td>( 9.87 \times 10^{-7} )</td>
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<td>( 4.93 \times 10^{-7} )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cor. Total</td>
<td>( 3.29 \times 10^{-4} )</td>
<td>14</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

6.3.3.4 Adequacy check of the model

The examination of residuals was employed to investigate the model adequacy. It
would give poor or misleading results if the model was not adequate fit (Wu et al. 2012a). The normal probability and studentized residuals plots illustrate whether the studentized residuals follow a normal distribution. Normal plot presented at Fig. 6-8 is normally distributed and resemble a straight line, indicating that there is no apparent problem with normality and no need for transformation of response (Khataee et al. 2010). Fig. 6-9 shows the studentized residual versus predicted value, and the residuals appear to be a random scatter. It suggests that the equality of variance does not seem to be violated (Grčić et al. 2009). Furthermore, the similarity between predicted and actual values of the response which was illustrated in Fig. 6-10 indicated that there was no significant violation of the models and the models were satisfactory and accurate.

![Normal Plot of Residuals](image)

**Fig. 6-8** The internally studentized residuals and normal % probability plot for Orange II decolorization rate constant
6.3.3.5 Response surface analysis

The 3D response surfaces plots were developed based on the RSM equation and showed in Fig. 6-11. There is an optimum current value for Orange II decolorization rate. In other word, increasing electrical current to certain extent could increase Orange II decolorization rate, while further increasing electrical current would decrease the
The decolorization rate. On the one hand, increase of applied current could promote the conversion of Fe(III) to Fe(II) on the surface of the catalyst (Eq. (6-5)) which resulted in more sulfate radicals generated on α-FeOOH surface. On the other hand, higher electrical current improves the generation of sulfate radicals and hydroxyl radicals via electron transfer reaction (Eq. (6-4)) and water discharge on DSA anode (M) (Eq. (6-3)) (Lin et al. 2014; Nidheesh and Gandhimathi 2014), respectively. However, further increasing the electrical current might lead to the side reactions and particularly the hydrogen gas evolution (Eq. (6-12)) which lead to the decline of the decolorization rate (Wu et al. 2012a).

\[
2\text{H}_2\text{O} + 2\text{e}^- \rightarrow \text{H}_2 + 2\text{OH}^- \quad (6-12)
\]

Fig. 6-11 also showed that the decolorization rate of Orange II increased slightly with the increase of α-FeOOH dosage and PDS concentration, because higher concentration of catalyst and PDS favored the generation of sulfate radicals.
Fig. 6-11 Response surface showing removal efficiency as a function of two independent variables:
(a) electrical current (A) and α-FeOOH dosage (B), (b) electrical current (A) and PDS concentration (C), (c) α-FeOOH dosage (B) and and PDS concentration (C)

130
6.3.4 Stability of \( \alpha \)-FeOOH

The recycle experiments were performed to investigate the stability of \( \alpha \)-FeOOH. The solid was easily removed from the electrochemical reactor after each repetitive oxidation process, then washed by deionised water, dried in the vacuum oven and stored at ambient temperature. The initial concentration of Orange II solution was fixed at 50 mg/L, PDS concentration was 2 g/L, \( \alpha \)-FeOOH dosage was 0.8 g/L, \( \text{Na}_2\text{SO}_4 \) solution concentration was 50 mM, initial pH was 7.0 and the applied current was 0.5 A. As shown in Fig. 6-12, the decolorization efficiencies of Orange II during three reaction cycles ranged from 86.5% to 92.1%, indicating \( \alpha \)-FeOOH is stable and can be reused.

![Graph showing the recycling study of \( \alpha \)-FeOOH during Orange II degradation](image)

**Fig. 6-12** Recycling study of \( \alpha \)-FeOOH during Orange II degradation (\( C_0 = 50 \) mg/L, [\( \alpha \)-FeOOH] = 0.5 g/L, [PDS] = 2 g/L, \( I_0 = 0.5 \) A, \( \text{pH}_0 = 7 \), [\( \text{Na}_2\text{SO}_4 \)] = 50 mM)

6.3.5 Changes in the UV-visible spectrum and mineralization efficiency of Orange II

To clarify the changes in molecular and structural characteristics of Orange II as a result of oxidation in the electro/\( \alpha \)-FeOOH/PDS process, representative UV–visible spectra changes in the dye solution as a function of reaction time were depicted and the
corresponding spectra are indicated in Fig. 6-13. As can be seen from these spectra, there was one main band located at 485 nm in the visible region, which was originated from the azo-band chromophore (−N=N−) (Wu et al. 2012b). Meanwhile, the absorption at 310 nm and 228 nm in the ultraviolet region were corresponded to naphthalene ring and benzene ring structures in the dye molecule (Zhang et al. 2009). As shown in Fig. 6-13(a), the visible band disappeared as the reaction proceeded, due to the fragmentation of the azo links by the reaction. The decay of the absorbance at 310 nm was considered as evidence of aromatic fragment degradation in the dye molecule and its intermediates. However, the absorbance at 228 nm increased, indicating that the azo structure was destroyed and some aromatic fragment such as benzene rings were formed.

In order to investigate the mineralization efficiency of Orange II electro/α-FeOOH/peroxydisulfate process, the changes of TOC were observed with initial Orange II concentration 50 mg/L, α-FeOOH dosage 0.5 g/L, PDS concentration 2 g/L, Na2SO4 solution concentration 50 mM, initial pH 7.0 and the electrical current 0.5A. As can be seen in Fig. 6-13(b), the TOC removal efficiency was only 12.8% after 120 min compared with 92.1% of Orange II degradation efficiency. Interestingly, by extending the reaction time to 180 min the TOC removal efficiency slightly increased to 16.7%, indicating most Orange II were converted to more simple intermediate products in electro/α-FeOOH/peroxydisulfate process.
Fig. 6-13 UV-visible spectra changes with reaction time in EC/α-FeOOH/PDS process (a) and the changes of TOC removal efficiency with reaction time (b) (C₀ = 50 mg/L, [α-FeOOH] = 0.5 g/L, I = 0.5 A, pH 7, [Na₂SO₄] = 50 mM)

6.4 Conclusion

Aqueous solutions of Orange II were effectively degraded by a novel
electro/α-FeOOH/peroxydisulfate process. Orange II decolorization efficiency first increased with the increase of electrical current, α-FeOOH dosage and PDS concentration, then it decreased when electric current, α-FeOOH dosage and PDS concentration were further increased. Initial pH value has insignificant influence on the decolorization of Orange II. ANOVA indicated both the interaction effect and the quadratic effect were significant on Orange II decolorization rate. The models were significant and could fit the experimental data well.
References


Saien, J., Soleymani, A.R. and Sun, J.H. (2011) Parametric optimization of individual and hybridized AOPs of Fe$^{2+}$/H$_2$O$_2$ and UV/S$_2$O$_8^{2−}$ for rapid dye destruction in aqueous media. Desalination 279(1–3), 298-305.


Application of response surface methodology to the removal of the antibiotic tetracycline by electrochemical process using carbon-felt cathode and DSA (Ti/RuO$_2$–IrO$_2$) anode. Chemosphere 87(6), 614-620.


Chapter 7 Decolorization of Orange II in water by

electro/Fe₃O₄/PDS process
7.1 Introduction

In this part, Orange II was degraded by electrochemical process combined with Fe\textsubscript{3}O\textsubscript{4} activated PDS process (EC/Fe\textsubscript{3}O\textsubscript{4}/PDS process). The effect of various reaction parameters such as initial pH, current density, PDS concentration and Fe\textsubscript{3}O\textsubscript{4} dosage on the removal of Orange II was investigated. The X-ray photoelectron spectroscopy (XPS) was applied to investigate the surface properties of Fe\textsubscript{3}O\textsubscript{4} before and after reaction. Gas chromatography–mass spectrometry (GC–MS) was applied to identify the intermediates and the degradation pathway of Orange II is proposed accordingly. The mineralization of Orange II in terms of TOC removal was also investigated. The change of acute toxicity during the treatment was investigated by activated sludge inhibition test.

7.2 Materials and methods

7.2.1 Materials

Orange II (4-(2-hydroxynaphthylazo) benzenesulfonic acid sodium salt) with reagent purity grade was obtained from Sinopharm Chemical Reagent Co., Ltd. (China) and used without further purification. Fe\textsubscript{3}O\textsubscript{4} used in this study was from Prolabo Co. (Paris, France). Sodium peroxydisulfate (Na\textsubscript{2}S\textsubscript{2}O\textsubscript{8}, 98%) and anhydrous sodium sulfate (Na\textsubscript{2}SO\textsubscript{4}, 99%) were purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China) and of analytical grade.

7.2.2 Experimental

The experimental set-up used for this study is illustrated in Fig. 7-1. Bulk electrolyses were carried out in undivided glass beaker with a capacity of 250 mL. A plate mixed metal oxide (DSA, Ti/RuO\textsubscript{2}–IrO\textsubscript{2}) anode (5.0 cm × 11.9 cm) is served as the anode and a plate of stainless steel with the same dimensions is used as cathode. The gap between the electrodes was 4.0 cm. The reactor was immersed in a water bath to
keep the temperature constant at 25 °C. A Model RYI-3010 direct current (DC) power supply from Shenzhen Yizhan Source, Co., Ltd. (China) was employed to provide the constant current.

Before each run, a fresh stock solution of Orange II was prepared in deionised water and the initial concentration \( (C_0) \) was kept at 25 mg/L. Sodium sulfate (50 mM) was added as a background electrolyte. The initial pH \( (\text{pH}_0) \) of Orange II solutions was measured with a Mettler-Toledo FE20 pH-meter (Mettler-Toledo Instruments Co., Ltd., Shanghai) and adjusted by sulfuric acid or sodium hydroxide. Before the beginning of electrolysis, a certain amount of PDS and Fe\(_3\)O\(_4\) were added into 200 mL of Orange II solution with constant stirring by using a mechanical stirrer (RW 20, IKA, Germany).

![Fig. 7-1 Schematic of the experimental setup employed in the present study](image)

7.2.3 Analysis

The absorbance of Orange II was measured at \( \lambda_{\text{max}} = 485 \) nm using a Rayleigh UV-9100 spectrophotometer (Rayleigh Co., China). Decolorization efficiency was calculated according to Eq. (7-1),

\[
\text{Decolorization efficiency (\%)} = \frac{A_0 - A}{A_0} \times 100
\]  

(7-1)

where the \( A_0 \) and \( A \) were the absorbance of the sample at time 0 and \( t \), respectively. The concentration of residual PDS was measured by an iodometric titration method (Wahba et al. 1959). The quantification of the dissolved iron ions concentration in the solution...
was systematically evaluated by Atomic Absorption Spectroscopy (AAS, Perkin Elmer A Analyst 800). XPS was operated on a KRATOS XSAM800 spectrometer equipped with MgKα X-ray source. GC–MS (VARIAN 450–GC/320–MS) was employed to determine the intermediates. Before analysis, Orange II solution was extracted with dichloromethane for ten times, and then the extracts were concentrated by rotary evaporator at 40 °C to about 1 mL (Zhong et al. 2011a; Zhong et al. 2011b). A VF–5 MS capillary column (30 m length × 0.25 mm ID × 0.25 μm film thickness) was employed for GC separation. The GC equipments were operated in a temperature programmed mode with an initial temperature of 35 °C held for 2 min, then ramped to 280 °C with a 10 °C/min rate. Helium was used as the carrier gas. Electronimpact (EI) mass spectra were scanned from 0 to 550 m/z.

The TOC of the samples were determined by an Analytik Jena Multi N/C 3100 analyzer with non dispersive infra red absorption detector (NDIR). The carrier gas was oxygen with a flow rate of 158 mL/min. The acute toxicity of intermediates generated during the treatment was assessed by means of activated sludge inhibition test (Olmez-Hanci et al. 2009). The activated sludge used as the heterotrophic biomass was obtained from a municipal wastewater treatment located in Wuhan. The mixed liquor suspended solids (MLSS) concentration used in the activated sludge inhibition experiments was adjusted to 1800 mg/L. To conduct the activated sludge inhibition test, 80 mL activated sludge was aerated 5 min and mixed with 20 mL treated wastewater at room temperature. The dissolved oxygen (DO) concentration of the mixed solution was measured by HQ30d (HACH) oxygen meter at selected time intervals. Oxygen uptake rate (OUR) was measured through the slope of the DO concentration versus time (Wu et al. 2012a):

\[
\text{OUR} = \frac{\Delta \text{DO}}{\Delta t}
\]

(7-2)

Specific oxygen uptake rate (SOUR) was determined by dividing OUR to the MLSS concentration:

\[
\text{SOUR} = \frac{\text{OUR}}{\text{MLSS}}
\]

(7-3)

where the MLSS were measured by filtration and evaporation the solution at 105 °C for 2 h.
7.3 Results and discussion

7.3.1 Decolorization of Orange II under different systems

To evaluate the decolorization efficiency of Orange II under different systems, the following blank experiments were conducted: Fe$_3$O$_4$ alone, PDS alone, Fe$_3$O$_4$/PDS process, EC alone, EC/Fe$_3$O$_4$ process, EC/PDS process and EC/Fe$_3$O$_4$/PDS process. The results are shown in Fig. 7-2a. It can be obviously seen that no appreciable Orange II decolorization was observed when Fe$_3$O$_4$ was added into the Orange II solution alone, indicating the effect of adsorption on Orange II decolorization was not obvious under the condition investigated. Little decolorization also occurred when PDS alone was applied. PDS is stable at ambient temperature and its oxidation power was limited ($E^0 = 2.01$ V) (Zhao et al. 2010b; Wu et al. 2012b), which could hardly degrade Orange II. The remaining percentage of PDS after 60 min reaction in the PDS alone process was 98.0% (Fig. 7-2b), indicating almost no PDS decomposed in this case. The concentration of Orange II was hardly decreased in Fe$_3$O$_4$/PDS process and the remaining percentage of PDS was 90%. This may due to the lack of Fe(II) on the surface of Fe$_3$O$_4$ particles. When treated by EC alone, 61.1% of 25 mg/L Orange II was decolorized due to a certain amount of hydroxyl radicals (M(•OH)) formed as intermediate of water discharge on DSA anode (M) (Eq. (7-4)) (Özcan et al. 2009).

\[ \text{M(H}_2\text{O)} \rightarrow \text{M(•OH) + H}^+ + e^- \] (7-4)

When 0.5 g/L Fe$_3$O$_4$ was employed to the electrolytic reactor, the decolorization efficiency of Orange II increased to 82.8%. It can be explained that in EC/Fe$_3$O$_4$ process, Fe$_3$O$_4$ was used as particle electrodes and could increase mass transfer coefficient and reduce energy consumption (Lin et al 2013). The decolorization efficiency slightly improved to 85.1% in EC/PDS process, which might be attributed to sulfate radicals produced via Eq. (7-5). 54.4% of PDS was decomposed in this case (Fig. 7-2b).

\[ \text{S}_2\text{O}_8^{2-} + e^- \rightarrow \text{SO}_4^- + \text{SO}_4^{2-} \] (7-5)

The decolorization efficiency of Orange II was 93.0% and the remaining percentage of PDS was 33.7% in the process of EC/Fe$_3$O$_4$/PDS, indicating Fe(II) can be
generated by the cathodic reduction of Fe(III) (Eq. (7-6)), thus more sulfate radicals can
formed by Fe(II) activating PDS reaction.

\[ \equiv \text{Fe(III)} + e^- \rightarrow \equiv \text{Fe(II)} \]  
(7-6)

![Graph showing decolorization of Orange II and remaining percentage of PDS over time.](image)

**Fig. 7-2** Decolorization of Orange II (a) and remaining percentage of PDS (b) after 60 min of
reaction under different systems ($C_0 = 25$ mg/L, [PDS] = 20 mM, [Fe$_3$O$_4$] = 0.5 g/L, $j = 8.4$ mA/cm$^2$, pH$_0$ 3.0, [Na$_2$SO$_4$] = 50 mM)
To verify the enhancement of the conversion of Fe(III) into Fe(II) in the process of EC/Fe$_3$O$_4$/PDS, the X-ray photoelectron spectroscopy was applied to characterize Fe$_3$O$_4$ before and after reaction. The domain peak for O 1s at 530.0 eV is characteristic of oxygen in a metal oxide (Zhao et al. 2012). The peak of Fe 2p spectrum at 710.6 and 724.4 eV represent Fe 2p3/2 and 2p1/2, respectively, indicated the existence of Fe$_3$O$_4$ (Huang et al. 2012). As can be seen from Fig. 7-3(a), no significantly change was observed about O 1s XPS spectra before and after reaction. Before the reaction, the Fe2p$_{3/2}$ spectrum fitted to four peaks located at binding energies (BE): 710.2, 711.3, 712.3 and 714.1 eV (Guo et al. 2010; Zhao et al. 2012). The peak at 710.2 eV could be assigned to the formation of ferrous oxide, where other peaks would be considered as the Fe(III) oxide (Fig. 7-3(b)). The ratio of Fe(II)/Fe(III) on the surface of Fe$_3$O$_4$ adsorbent increased from 0.75 to 1.09 after 60 min reaction. This implied more Fe(II) was generated on the surface of Fe$_3$O$_4$ particles in the electrochemical reactor (Eq. (7-6)).
Based on the results and the literature (Liang et al. 2007; Özcan et al. 2009; Yan et al. 2011; Lin et al. 2013), the mechanism of EC/Fe₃O₄/PDS process was indicated in Fig. 7-4. Fe(II) was generated by the reduction and transformation of Fe(III) on the surface of Fe₃O₄ due to the direct electron transfer (Eq. (7-6)), and then Fe(II) activated PDS to release sulfate radicals via Eq. (7-7)). Meanwhile, hydroxyl radicals were generated both by water discharge on DSA anode (Eq. (7-4)) and the reaction between sulfate radicals and H₂O or OH⁻ (Eqs. (7-8) and (7-9)). Orange II was degraded by both sulfate radicals and hydroxyl radicals.

$$\text{≡Fe(II) + S}_2\text{O}_8^{2-} \rightarrow \equiv\text{Fe(III) + SO}_4^{2-} + \text{SO}_4^{2-}$$ \hspace{1cm} (7-7)

$$\text{SO}_4^{2-} + \text{H}_2\text{O} \rightarrow \text{H}^+ + \text{SO}_4^{2-} + \cdot\text{OH}$$ \hspace{1cm} (7-8)

$$\text{SO}_4^{2-} + \text{OH}^- \rightarrow \text{SO}_4^{2-} + \cdot\text{OH}$$ \hspace{1cm} (7-9)
Fig. 7-4 Proposed mechanism of EC/Fe₃O₄/PDS process

The degradation kinetics of different systems was investigated and the results were summarized in Table 7-1. It can be seen that the decolorization of 25 mg/L Orange II by EC, EC/Fe₃O₄, EC/PDS and EC/Fe₃O₄/PDS process followed apparent first order kinetics and the apparent rate constants ($k_{app}$) were 0.015, 0.028, 0.030 and 0.043 min⁻¹, respectively, at pH 3.0.

<table>
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<th>Run</th>
<th>pH₀</th>
<th>Current density (mA/cm²)</th>
<th>[PDS] (mM)</th>
<th>[Fe₃O₄] (g/L)</th>
<th>$k_{app}$ (min⁻¹)</th>
<th>$R^2$</th>
<th>Residual [PDS] (mM)</th>
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<tr>
<td>1</td>
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<td>0.998</td>
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<td>0.043</td>
<td>0.992</td>
<td>6.74</td>
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</table>
7.3.2 Effect of initial pH

The effect of initial pH was observed when Orange II concentration was 25 mg/L, PDS concentration was 20 mM, Fe$_3$O$_4$ dosage was 0.5 g/L, Na$_2$SO$_4$ solution concentration was 50 mM and the electrical density was 8.4 mA/cm$^2$. As can be seen from Fig. 7-5, the decolorization efficiencies of Orange II were 93.0%, 91.2% and 92.1% at initial pH 3.0, 6.0 and 9.0, respectively. When initial pH ranged from 3.0 to 9.0, the rate constants varied slightly from 0.037 to 0.043 min$^{-1}$ (Table 7-1). Sulfate radicals generated in EC/Fe$_3$O$_4$/PDS process could react with H$_2$O or OH$^-$ to produce hydroxyl radicals (Liang et al. 2007; Saien et al. 2011). Either H$^+$ could be generated or OH$^-$ could be consumed (Eqs. (7-8) and (7-9)), which resulted in the reduction of solution pH. Moreover, the dissociation of HSO$_4^-$ (Eq. (7-10)), which was formed through Eq. (7-11), would further release H$^+$ and decrease the solution pH (Kusic et al. 2011).

\[
\begin{align*}
\text{HSO}_4^- & \rightarrow \text{SO}_4^{2-} + \text{H}^+ \quad \text{(7-10)} \\
\text{SO}_4^{2-} + \cdot \text{OH} & \rightarrow \text{HSO}_4^- + 1/2\text{O}_2 \quad \text{(7-11)}
\end{align*}
\]

Therefore, the pH of Orange II solution after 60 min at different initial pH was almost the same (about 2.3). As a result, the decolorization efficiency of Orange II and the decomposition of PDS (Table 7-1) were similar at different initial pH values. This

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<table>
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<tr>
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<tr>
<td>7</td>
<td>6.0</td>
<td>3.36</td>
<td>20</td>
<td>0.5</td>
<td>0.027</td>
<td>0.988</td>
</tr>
<tr>
<td>8</td>
<td>6.0</td>
<td>16.8</td>
<td>20</td>
<td>0.5</td>
<td>0.071</td>
<td>0.989</td>
</tr>
<tr>
<td>9</td>
<td>6.0</td>
<td>25.2</td>
<td>20</td>
<td>0.5</td>
<td>0.117</td>
<td>0.973</td>
</tr>
<tr>
<td>10</td>
<td>6.0</td>
<td>8.40</td>
<td>5</td>
<td>0.5</td>
<td>0.030</td>
<td>0.998</td>
</tr>
<tr>
<td>11</td>
<td>6.0</td>
<td>8.40</td>
<td>10</td>
<td>0.5</td>
<td>0.053</td>
<td>0.994</td>
</tr>
<tr>
<td>12</td>
<td>6.0</td>
<td>8.40</td>
<td>10</td>
<td>0.2</td>
<td>0.029</td>
<td>0.991</td>
</tr>
<tr>
<td>13</td>
<td>6.0</td>
<td>8.40</td>
<td>10</td>
<td>0.8</td>
<td>0.057</td>
<td>0.969</td>
</tr>
<tr>
<td>14</td>
<td>6.0</td>
<td>8.40</td>
<td>10</td>
<td>1.2</td>
<td>0.049</td>
<td>0.976</td>
</tr>
</tbody>
</table>
proved that the EC/Fe$_3$O$_4$/PDS process could be successfully applied at a wide range of initial pH from 3.0 to 9.0. Since pH 6.0 was the natural pH of 25 mg/L Orange II solution, it was selected in the following experiments.

![Graph showing the effect of initial pH on the oxidation efficiency of Orange II.](image)

**Fig. 7-5** The effect of initial pH on the oxidation efficiency of Orange II ($C_0 = 25$ mg/L, [PDS] = 20 mM, [Fe$_3$O$_4$] = 0.5 g/L, $j = 8.4$ mA/cm$^2$, [Na$_2$SO$_4$] = 50 mM)

### 7.3.3 Effect of current density

Fig. 7-6 illustrated the effect of current density with initial Orange II concentration of 25 mg/L, PDS concentration 20 mM, Fe$_3$O$_4$ dosage 0.5 g/L, Na$_2$SO$_4$ solution concentration 50 mM and the initial pH 6.0. The decolorization efficiency and apparent rate constant were 80.3% and 0.027 min$^{-1}$ (Table 7-1) in a 60 min reaction when current density was 3.4 mA/cm$^2$. When current density increased to 25.2 mA/cm$^2$, the decolorization efficiency was nearly 100% in 30 min reaction and the rate constant increased to 0.117 min$^{-1}$ (Table 7-1). On the one hand, when current density increased, more hydroxyl radicals would be formed on anode surface (Eq. (7-4)) and more sulfate radicals would be generated via electron transfer reaction (Eq. (7-5)). On the other hand, the increase of current density favored the cathodic reduction of Fe(III) (Eq. (7-6)), and more PDS would be activated to generate sulfate radicals via Eq. (7-7). Meanwhile, Table 7-1 demonstrated that the increase of current density from 3.34 to 25.2 mA/cm$^2$
did cause the increase of PDS decomposition. This also suggested more sulfate radicals would be generated when current density increased.

Fig. 7-6 The effect of current density on the oxidation efficiency of Orange II ($C_0 = 25$ mg/L, [PDS] = 20 mM, $[\text{Fe}_3\text{O}_4] = 0.5$ g/L, pH$_0$ 6.0, $[\text{Na}_2\text{SO}_4] = 50$ mM)

7.3.4 Effect of PDS concentration

As the source of sulfate radicals, PDS plays a significant role in EC/$\text{Fe}_3\text{O}_4$/PDS process. The effect of PDS concentration was investigated by varying the concentration of PDS from 5 to 20 mM when Orange II concentration was 25 mg/L, $\text{Fe}_3\text{O}_4$ dosage was 0.5 g/L, $\text{Na}_2\text{SO}_4$ solution concentration was 50 mM, initial pH was 6.0 and the electrical density was 8.4 mA/cm$^2$. As can be seen from Fig. 7-7, when PDS concentration increased from 5 to 10 mM, the Orange II decolorization efficiency and apparent rate constants over a 60 min reaction increased from 84.1% to 95.8% and the rate constants increased from 0.030 to 0.053 min$^{-1}$ (Table 7-1). Meanwhile, the amount of PDS decomposed also increased, indicating more reactive radicals would be generated to degrade Orange II at higher PDS concentration. When PDS concentration increased to 20 mM, the Orange II decolorization efficiency and rate constants after 60 min declined slightly to 91.2% and 0.039 min$^{-1}$ (Table 7-1). This was due to the fact that the side reaction between $S_2\text{O}_8^{2-}$ and $\text{SO}_4^{2-}$ became more significant (Eq. (7-12))
when PDS concentration was excessive (Hori et al. 2005; Wu et al. 2012b).

\[
S_{2}O_{8}^{2-} + SO_{4}^{-} \rightarrow S_{2}O_{8}^{2-} + SO_{4}^{2-}
\]  

(7-12)

Therefore, 10 mM was selected as the optimal initial PDS concentration and used in the following experiments.

![Fig. 7-7 The effect of PDS concentration on the oxidation efficiency of Orange II (C_0 = 25 mg/L, [Fe_3O_4] = 0.5 g/L, pH_0 6.0, j = 8.4 mA/cm^2, [Na_2SO_4] = 50 mM)](image)

**7.3.5 Effect of Fe_3O_4 dosage**

The comparison of Orange II decolorization efficiency under different Fe_3O_4 dosage was investigated with initial Orange II concentration 25 mg/L, PDS concentration 10 mM, Na_2SO_4 solution concentration 50 mM, the initial pH 6.0 and the electrical density 8.4 mA/cm^2. The results were shown in Fig. 7-8. When Fe_3O_4 dosage varied from 0.2 to 0.8 g/L, the decolorization efficiency of Orange II within 60 min reaction increased from 83.9% to 97.6% and apparent rate constants increased from 0.029 to 0.057 min^{-1} (Table 7-1). At higher Fe_3O_4 dosage, more Fe(II) could be generated through Eq. (7-5), and then a considerable amount of sulfate radicals would be produced via Eq. (7-6). These sulfate radicals reacted immediately with Orange II, leading to a fast increase of the Orange II decolorization. When Fe_3O_4 dosage further increased to 1.2 g/L, the Orange II decolorization efficiency declined slightly to 96.0%
and apparent rate constants decreased to 0.049 min\(^{-1}\) (Table 7-1). When Fe\(_3\)O\(_4\) particles were added in excess, an excess of sulfate radicals would be generated, resulting in considerable disappearance of sulfate radicals without the degradation of Orange II due to the combination between sulfate radicals themselves (Eq. (7-13)). Moreover, the interaction between excess Fe(II) on the surface of Fe\(_3\)O\(_4\) particles and sulfate radicals might directly quench sulfate radicals via Eq. (7-14) (Yan et al. 2011).

\[
\text{SO}_4^{\cdot-} + \text{SO}_4^{\cdot-} \rightarrow 2\text{SO}_4^{2-} \quad (7-13)
\]

\[
\equiv\text{Fe(II)} + \text{SO}_4^{\cdot-} \rightarrow \equiv\text{Fe(III)} + \text{SO}_4^{2-} \quad (7-14)
\]

As a result, the decomposition of PDS increased with an increase in Fe\(_3\)O\(_4\) dosage of up to 0.8 g/L (Table 7-1), and the highest decolorization efficiency of Orange II was obtained at Fe\(_3\)O\(_4\) dosage of 0.8 g/L. This value was chosen as the optimal Fe\(_3\)O\(_4\) dosage and applied in the following experiments.

![Fig. 7-8](image)

**Fig. 7-8** The effect of Fe\(_3\)O\(_4\) dosage on the oxidation efficiency of Orange II \((C_0 = 25 \text{ mg/L, [PDS]} = 10 \text{ mM, pH}_0 = 6.0, j = 8.4 \text{ mA/cm}^2, [\text{Na}_2\text{SO}_4] = 50 \text{ mM})\)

### 7.3.6 Stability of Fe\(_3\)O\(_4\)

The recycle experiments were performed to investigate the stability of Fe\(_3\)O\(_4\). The solid was easily removed from the electrochemical reactor after each repetitive oxidation process, then washed by deionised water, dried in the vacuum oven and stored
at ambient temperature. The initial concentration of Orange II solution was fixed at 25 mg/L, PDS concentration was 10 mM, Fe$_3$O$_4$ dosage was 0.8 g/L, Na$_2$SO$_4$ solution concentration was 50 mM, initial pH was 6.0 and the electrical density was 8.4 mA/cm$^2$. As shown in Fig. 7-9a, the decolorization efficiencies of Orange II during three reaction cycles ranged from 98.0% to 95.3%. The remaining percentages of PDS in each recycle experiment were similar (Fig. 7-9b). The metal leaching level was not obviously changed after three recycle time, with a Fe content in the solution at the end of the test almost constant at 13.05-32.72 mg/L. This indicated Fe$_3$O$_4$ is stable and can be reused.

**Fig. 7-9** Recycling study of the degradation of Orange II (a) and remaining percentage of PDS (b)
(C₀ = 25 mg/L, [PDS] = 10 mM, [Fe₃O₄] = 0.8 g/L, pH₀ 6.0, j = 8.4 mA/cm², [Na₂SO₄] = 50 mM)

7.3.7 The degradation pathway of Orange II

GC–MS was employed to identify the intermediate products formed during EC/Fe₃O₄/PDS process. The main intermediate products detected were illustrated in Table 7-2 and a plausible degradation pathway was proposed (Fig. 7-10) based on the results and previous studies (Özcan et al. 2009; Zhao et al. 2010a; Zhong et al. 2011b). The cleavage of –N=N– of Orange II caused by electron transfer of sulfate radicals (Yuan et al. 2011) leads to the formation of sodium sulfanilamide and 1-amino-2-naphthol. Sodium sulfanilamide is thermally stable and highly soluble in water, whereas 1-amino-2-naphthol is oxygen sensitive and may be decomposed under aerobic conditions (Zhao et al. 2010a; Zhong et al. 2011b). Therefore, neither of these two intermediates was detected during GC–MS analysis. Afterwards, sodium sulfanilamide was subsequently oxidized to 1,4-benzoquinone (A). 1-amino-2-naphthol was firstly oxidized to 1,2-naphthalenedione (B). 1,2-naphthalenedione (B) was subsequently oxidized to the quinoid structures, resulting in the formation of 1(3H)-isobenzofuranone (C), phthalic anhydride (D), methyl hydrogen phthalate (E) and 2-(hydroxymethyl) benzoic acid (F) (Özcan et al. 2009, Zhao et al. 2010a)[1,50]. Further oxidation resulted in aromatic ring cleavage and generation of carboxylic acids and finally converted to CO₂ and H₂O.
<table>
<thead>
<tr>
<th>Compound</th>
<th>Retention time (min)</th>
<th>Chemical name</th>
<th>Chemical structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>6.03</td>
<td>1,4-benzoquinone</td>
<td><img src="image" alt="1,4-benzoquinone" /></td>
</tr>
<tr>
<td>B</td>
<td>16.02</td>
<td>1,2-naphthalenedione</td>
<td><img src="image" alt="1,2-naphthalenedione" /></td>
</tr>
<tr>
<td>C</td>
<td>12.75</td>
<td>1(3H)-isobenzofuranone</td>
<td><img src="image" alt="1(3H)-isobenzofuranone" /></td>
</tr>
<tr>
<td>D</td>
<td>12.24</td>
<td>phthalic anhydride</td>
<td><img src="image" alt="phthalic anhydride" /></td>
</tr>
<tr>
<td>E</td>
<td>12.26</td>
<td>methyl hydrogen phthalate</td>
<td><img src="image" alt="methyl hydrogen phthalate" /></td>
</tr>
<tr>
<td>F</td>
<td>12.83</td>
<td>2-(hydroxymethyl) benzoic acid</td>
<td><img src="image" alt="2-(hydroxymethyl) benzoic acid" /></td>
</tr>
</tbody>
</table>
7.3.8 The changes of TOC and toxicity with reaction time

In order to clarify the mineralization efficiency of Orange II in EC/Fe$_3$O$_4$/PDS
process, the changes of TOC were investigated with initial Orange II concentration 25 mg/L, PDS concentration 10 mM, Fe$_3$O$_4$ dosage 0.8 g/L, Na$_2$SO$_4$ solution concentration 50 mM, initial pH value 6.0 and the current density 16.8 mA/cm$^2$. As can be seen in Fig. 7-11, Orange II was almost completely removed after a 60 min reaction, but the TOC removal efficiency was only 19.6%. Further prolong of reaction time to 90 min, the TOC removal efficiency achieved 30.0%. This indicated part of Orange II was converted to smaller intermediates. The variation of acute toxicity in EC/Fe$_3$O$_4$/PDS process was monitored by oxygen uptake (Olmez-Hanci et al. 2009; Olmez-Hanci et al. 2010). As indicated in Fig.7-11, the SOUR value decreased at the beginning of the reaction, indicating more toxic products were generated. It has been illustrated in Section 7.3.7 that the main intermediates formed at the early stage of the treatment are 1,4-benzoquinone (A) and 1,2-naphthalenedione (B). Therefore, the higher toxicity observed at the beginning of EC/Fe$_3$O$_4$/PDS process can be attributed to the formation of these intermediates, because the toxicity of quinones is well-known (Hammami et al. 2008). After 30 to 90 min reaction, the value of SOUR increased gradually. This means further treatment could convert these intermediates to less toxic products and mineralization led to the detoxification of treated solution.

![Fig. 7-11 The changes of TOC, Orange II concentration and SOUR with reaction time (C$_0$ = 25 mg/L, [PDS] = 10 mM, [Fe$_3$O$_4$] = 0.8 g/L, j = 16.8 mA/cm$^2$, pH$_0$ 6.0, [Na$_2$SO$_4$] = 50 mM)](image-url)
7.3.9 Electrical energy consumption estimation

In order to evaluate the energy consumption of EC/Fe$_3$O$_4$/PDS process, an electrical energy consumption per order of magnitude (EE/O) was employed. EE/O is defined as the number of kilowatt hours of electrical energy required for bringing about the degradation of a pollutant by one order of magnitude (90%) in 1 m$^3$ of contaminated water (Daneshvar et al. 2007; Sadik 2007a; Sadik 2007b; Behnajady et al. 2009).

$$EE/O \ (\text{kWh m}^{-3} \text{order}^{-1}) = \frac{U_{\text{cell}}It}{V \log(A_0/A)} \quad (7-15)$$

where $U_{\text{cell}}$ is the average cell voltage (V), $I$ is the current (A), $t$ is the electrolysis time (h) and $V$ is the volume (L).

The electrical energy consumption for Orange II degradation under different reaction systems were summarized in Table 7-3. It can be seen from Table 7-3 that the electrocatalysis processes were more energy saving than the photocatalysis processes. For example, EE/O value was 273.9 (kWh m$^{-3}$ order$^{-1}$) for UV/TiO$_2$ process, while EC/Fe$_3$O$_4$ process only needed 19.65 (kWh m$^{-3}$ order$^{-1}$) electrical energy consumption. What’s more, the presence of oxidant (H$_2$O$_2$ or PDS) could improve electrocatalysis and photocatalysis, and the associated electrical energy consumption decreased. Table 4 indicated that the EE/O value of EC/Fe$_3$O$_4$/PDS process was less than one-half of EC/Fe$_3$O$_4$ process.

<table>
<thead>
<tr>
<th>Method</th>
<th>Operating parameters</th>
<th>EE/O (kWh m$^{-3}$ order$^{-1}$)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>UV/TiO$_2$</td>
<td>[Orange II]$_0$ = 17.5 mg/L, [TiO$_2$] = 0.5 g/L, pH$_0$ = 6.1</td>
<td>273.9</td>
<td>Sadik 2007b</td>
</tr>
<tr>
<td></td>
<td>43 W UV lamp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>EC/Fe$_3$O$_4$</td>
<td>[Orange II]$_0$ = 25 mg/L, [Fe$_3$O$_4$] = 0.5 g/L, $j = 8.$ mA/cm$^2$, pH$_0$ = 3.0, [Na$_2$SO$_4$] = 50 mM</td>
<td>19.65</td>
<td>Lin et al. 2013</td>
</tr>
<tr>
<td></td>
<td>3.0, [Na$_2$SO$_4$] = 50 mM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>UV/TiO$_2$/H$_2$O$_2$</td>
<td>[Orange II]$_0$ = 17.5 mg/L, [H$_2$O$_2$] = 71 mM, [TiO$_2$] = 0.:</td>
<td>40.45</td>
<td>Sadik 2007b</td>
</tr>
<tr>
<td></td>
<td>37 g/L, pH$_0$ = 6.1, 43 W UV lamp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>UV/TiO$_2$+ZnO/PDS</td>
<td>[Orange II]$_0$ = 17.5 mg/L, [PDS] = 8.5 mM, [TiO$_2$+ZnO]</td>
<td>18.08</td>
<td>Sadik</td>
</tr>
</tbody>
</table>

Table 7-3 Comparison of electrical consumption under the different reaction system
Aqueous solutions of Orange II have been degraded effectively in the EC/Fe₃O₄/PDS process, in which Fe₃O₄-activated PDS process is enhanced by the electrochemical process. The effect of initial pH, current density, PDS concentration and Fe₃O₄ dosage, on the Orange II decolorization was investigated. The results indicated that Orange II can be totally decolorized in a 60 min reaction when initial Orange II concentration was 25 mg/L, PDS concentration was 10 mM, Fe₃O₄ dosage was 0.8 g/L, current density was 8.4 mA/cm² and initial pH was 6.0. Recycle experiments showed Fe₃O₄ particles were stable and can be reused. XPS spectrum indicated Fe(II) was generated on the surface of Fe₃O₄ particles after reaction. The main intermediates were separated and identified by GC–MS technique and a plausible degradation pathway of Orange II was proposed.
References


by a novel electro/Fe$^{3+}$/peroxydisulfate process. Separation and Purification Technology 117, 18-23.


Chapter 8 General conclusion and future perspectives
8.1 General conclusion

In this paper, electro-Fenton and electro-Fenton-like processes were used to degrade artificial sweeteners and azo dyes. The results obtained during this thesis work concern the removal efficiency, the oxidation mechanism, degradation pathway and toxicity evolution of target pollutants.

(1) Electro-Fenton process was an effective method for the degradation of ASP in water. The oxidative degradation rate and mineralization efficiency were affected mainly by the catalyst (Fe$^{2+}$) concentration and applied current. The absolute rate constant of hydroxylation reaction of ASP was found to be $(5.23 \pm 0.02) \times 10^9$ M$^{-1}$ s$^{-1}$. Short-chain aliphatic acids such as oxalic, oxamic and maleic acid were identified as aliphatic intermediates in the electro-Fenton process. The bacteria luminescence inhibition showed the toxicity of ASP solution decreased, during the treatment, after reached a maximum during the first period of the oxidation reaction.

(2) Artificial sweetener SAC could be degraded effectively by electro-Fenton process with a DSA, Pt or BDD anode. However, the using of BDD anode could accelerate the mineralization of SAC. The optimal conditions for SAC removal were SAC concentration 0.2 mM, Fe$^{2+}$ concentration 0.2 mM, Na$_2$SO$_4$ concentration 50 mM, applied current 200 mA and initial pH 3.0. Oxalic, formic, and maleic acids were observed as aliphatic by-products of SAC during electro-Fenton process. The bacteria luminescence inhibition showed the toxicity of SAC solution increased at the beginning of electrolysis, and then it declined until the end of the reaction.

(3) Artificial sweetener SUC could be completely mineralized in a 360 min reaction by electro-Fenton process with a Pt or BDD anode. The mineralization rate was affected mainly by the Fe$^{2+}$ concentration and applied current. The mineralization current efficiency (MCE) decreased with rising applied current from 100 to 500 mA with both Pt and BDD anodes. Oxalic, pyruvic, formic and glycolic acids were detected during the oxidation of SUC.

(4) Orange II was effectively decolorized by EC/α-FeOOH/PDS process. The initial pH of Orange II solution had little effect on the decolorization of Orange II. RSM
based on Box-Behnken statistical experiment design was applied to analyze the experimental variables. The response surface methodology models were derived based on the results of the pseudo-first-order decolorization rate kinetics and the response surface plots were developed accordingly. The results indicated that the applied current showed a positive effect on the decolorization rate constant of Orange II. The interaction of α-FeOOH dosage and PDS concentration were significant. The ANOVA results confirmed that the proposed models were accurate and reliable for the analysis of the variables of EC/α-FeOOH/PDS process. The catalyst α-FeOOH showed good structural stability and could be reused.

(5) Aqueous solutions of Orange II have been degraded effectively in the EC/Fe$_3$O$_4$/PDS process. The decolorization rate was affected by the initial pH of Orange II solution, current density, PDS concentration and Fe$_3$O$_4$ dosage. Orange II can be totally decolorized in a 60 min reaction time when initial Orange II concentration was 25 mg/L, PDS concentration was 10 mM, Fe$_3$O$_4$ dosage was 0.8 g/L, current density was 8.4 mA/cm$^2$ and initial pH was 6.0. Recycle experiments showed that Fe$_3$O$_4$ particles were stable and can be reused. XPS spectrum indicated that Fe(II) was generated on the surface of Fe$_3$O$_4$ particles after reaction. The main reaction intermediates of Orange II were separated and identified by GC–MS technique and a plausible degradation pathway was proposed.

8.2 Future perspectives

Further studies can be developed as follows:

(1) Finding out the suitable method to detect the intermediate products of artificial sweeteners and then obtain the degradation pathway;

(2) In the sulfate radical-based electro-Fenton-like process, persulfate is remained in the solution after the reaction. Moreover, persulfate can be decomposed to SO$_4^{2-}$ and H$^+$, resulting the decline of the pH value. So it is important to separate persulfate and SO$_4^{2-}$ from the treated solution.

(3) Both sulfate radicals and hydroxyl radicals are contributed to the degradation
of organic contaminants in the sulfate radical-based electro-Fenton-like process. When attacking organic compounds, hydroxyl radicals are more likely to do through hydrogen abstraction or addition reactions, while sulfate radicals participate in electron transfer reaction. Therefore, further investigation can focus on the different by-products generated under the attack of these two radicals.

(4) All the solutions contained organic pollutants are synthetic wastewaters which were prepared in the lab. In the future, we can use electro-Fenton and sulfate radical-based electro-Fenton-like processes to degrade target pollutants in actual wastewater.
Publications


