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Mineralization of organic pollutants by anodic oxidation using reactive electrochemical membrane synthesized from carbothermal reduction of TiO₂

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A B S T R A C T

Reactive Electrochemical Membrane (REM) prepared from carbothermal reduction of TiO₂ is used for the mineralization of biorefractory pollutants during filtration operation. The mixture of Ti₄O₇ and Ti₅O₉ Magnéli phases ensures the high reactivity of the membrane for organic compound oxidation through *OH mediated oxidation and direct electron transfer. In cross-flow filtration mode, convection-enhanced mass transport of pollutants can be achieved from the high membrane permeability (3300 LMH bar⁻¹). Mineralization efficiency of oxalic acid, paracetamol and phenol was assessed as regards to current density, transmembrane pressure and feed concentration. Unprecedented high removal rates of total organic carbon and mineralization current efficiency were achieved after a single passage through the REM, e.g. 47 g m⁻² h⁻¹ 72% and 6.7 g m⁻² h⁻¹ 47% for oxalic acid and paracetamol, respectively, at 15 mA cm⁻². However, two mechanisms have to be considered for optimization of the process. When the TOC flux is too high with respect to the current density, aromatic compounds polymerize in the REM layer where only direct electron transfer occurs. This phenomenon decreases the oxidation efficiency and/or increases REM fouling. Besides, O₂ bubbles sweeping at high permeate flux promotes O₂ gas generation, with adverse effect on oxidation efficiency.

1. Introduction

Anodic oxidation (AO) is an electrochemical advanced oxidation process (EAOP) increasingly recognized as a promising next-generation technology for the treatment of contaminated effluents (Panizza and Cerisola, 2009; Chaplin, 2014; Radjenovic and Sedlak, 2015; Moreira et al., 2017). The process is based on both direct electron transfer (DET) from organic compounds (R) to the anode surface (eq. (1)) and generation of hydroxyl radicals (*OH) from water oxidation on the surface of electrodes (M) with high O₂ overvoltage (eq. (2)) (Panizza and Cerisola, 2009). *OH is a very strong oxidant (E° = 2.8 V vs SHE) (Latimer, 1952) allowing the degradation of a large range of bio-refractory organic compounds. Mineralization of various micropollutants and complex effluents has been achieved using AO (Ozcan et al., 2008; Balci et al., 2009; Oturan et al., 2015; Martínez-Huitle et al., 2015; Brillas and Martínez-Huitle, 2015; Antonin et al., 2015; Trellu et al., 2016). However, several scientific challenges still need to be overcome in order to promote the application of AO for water treatment.

\[ \text{R} \rightarrow (\text{R}^*)^+ + e^- \]  
\[ \text{M} + \text{H}_2\text{O} \rightarrow \text{M}(*\text{OH}) + \text{H}^+ + e^- \]  

*OH are produced in a heterogeneous way in the electrochemical cell, therefore, new approaches are necessary in order to break the high cost bottleneck related to the limited mass transfer of organic pollutants from the bulk to the electrode surface (Yang et al., 2009; Chaplin, 2014; Radjenovic and Sedlak, 2015). Electro-oxidation is controlled by mass transport phenomenon when the concentration of organic compounds is too low by comparison to the electron transfer rate (current density). In the potential region of water discharge, secondary reactions such as oxygen evolution...
strongly decreases the current efficiency (Panizza et al., 2001; Panizza and Cerisola, 2009). Unfortunately, most of anode materials used in AO can only be produced under the form of conventional plate electrodes (e.g. boron-doped diamond), for which mass transport is limited by surface diffusivity in the external film (Brillas and Martinez-Huitle, 2011). This technological constraint also limits the potential for scaling up. Implementation of flow-through porous electrodes is the most promising way to overcome mass transfer limitations (Zaky and Chaplin, 2013; Ganiyu et al., 2015; Radjenovic and Sedlak, 2015; Ronen et al., 2016).

Recently, carbon nanotube electrochemical filters have been developed for the removal of organic pollutants, however they are not active for *OH production (Tsierkezos and Ritter, 2012; Schnoor and Vecitis, 2013; Gao and Vecitis, 2013; Liu et al., 2015). Besides, research works from Guo et al. (2016) showed that a convection-enhanced rate constant for Fe(CN)₆³⁻ oxidation of 1.4 × 10⁻⁴ s⁻¹ – the highest reported in the literature, close to the kinetic limit – was reached by using porous sub-stoichiometric titanium oxide (TiOₓ) reactive electrochemical membrane (REM) (Guo et al., 2016). The high membrane permeability (3200 L m⁻² h⁻¹ bar⁻¹) resulted in a strong increase of the convective mass transport of pollutants from the bulk solution to electron-transfer sites at the surface and in the porosity of the material. The reactivity of the material has been investigated by using probe molecules (oxalic acid, coumarin, terephthalic acid) showing that both DET and formation of *OH were involved in the oxidation of organic compounds (Zaky and Chaplin, 2013, 2014; Ganiyu et al., 2016). First studies on TiOₓ REM used either commercial Ebonex® membranes (Walsh and Wilks, 2010; Zaky and Chaplin, 2013) or TiO₂ ultrafiltration membranes reduced to TiOₓ under 1 atm H₂ during 50 h at 1050 °C (Guo et al., 2016). The latter showed the most promising results because of a higher permeability and predominance of the TiO₂ phase but high production costs are associated to the synthesis method.

The focus of this study is on testing a novel TiOₓ REM synthesized from carbothermal reduction of TiO₂. The REM is characterized by scanning electron microscopy (SEM), X-ray diffraction (XRD), Hg porosimetry and water permeability. In cross-flow filtration mode, the effect of permeate flux, pollutant concentration and current density on mineralization efficiency of various organic compounds (oxalic acid (OA), paracetamol (PCT), phenol (PHE)) is assessed. A focus is given on phenomena limiting the efficiency of the REM during treatment of aromatic molecules at high concentration or high permeate flux. From the different behaviors observed, electro-oxidation mechanisms and potential of the REM to be applied for water treatment are highlighted.

2. Materials and methods

2.1. Chemicals

All chemicals were of reagent grade and purchased from Sigma Aldrich (Na₂SO₄, PCT, OA) or Alfa Aesar (PHE). Solutions were prepared using ultrapure water (Purrelab®, Elga LabWater, 18.2 MΩ cm).

2.2. REM synthesis and characterization

Porous Magneli phase was synthesized from carbothermal reduction of TiO₂ by using an extrusion die, based on the method developed by Saint-Gobain CREE. The extrusion paste contained 74.1% of anatase form of TiO₂ (Altichem, median particle diameter of 0.37 μm), 2.5% of carbon black (Thermox N990), 18.4% of water and 4.5% of organic binders used as extrusion adjuvant. Organic binders were removed during the debinding step under air atmosphere at 350 °C, 2 h. Finally, sintering and carbothermal reduction of TiO₂ to TiOₓ was performed during 2 h under argon atmosphere at 1300 °C.

REM morphology was analyzed from images obtained by SEM at high resolution (Hitachi S4800) at 30 and 3000 magnification. Hg porosimetry analysis was performed on a AutoPore IV (MicroMetrics) device. The nature of Magneli phases was characterized by powder XRD using X’Pert Pro (Pan Analytical) system with a Cu X-ray tube (40 kV and 20 mA). Scans were collected by Pan Aytlyal software and processed with Full Prof and Origin Pro software.

2.3. Cross-flow filtration and electro-oxidation setup

The REM (inner and outer diameter of 6 and 10 mm, active length of 9 cm) was used as anode in inside-outside cross-flow filtration mode. A 3 mm diameter stainless steel rod used as cathode was placed at the center of the REM. A recirculating pump and a pressurized reservoir tank ensured constant cross-flow rate (0.88 m s⁻¹) and transmembrane pressure (TMP, 40–1000 mbar). Experiments were performed with OA (0.75–36 mM; TOC = 18–870 mg L⁻¹), PCT (0.1–2.3 mM; TOC = 9.6–220 mg L⁻¹) or PHE (0.25–2 mM; TOC = 18–140 mg L⁻¹) as model organic compounds. Initial pH of the solution was not adjusted and the evolution of the pH in the permeate and retentate was monitored. Sodium sulfate (50 mM) was added as supporting electrolyte. Only the retentate was 100% recirculated, while the permeate was continuously sampled for analysis. Experiments were performed in galvanostatic mode, using a power supply from ELC (AL924A). Current density (j) was in the range 6–30 mA cm⁻². Potential difference in the electrochemical cell was in the range 4.0–6.5 V, according to current intensity. All experiments were performed with a constant volume of permeate collected (350 mL). A schematic view of the setup is provided in Fig. 1.

2.4. Analytical methods

Concentration of PCT was analyzed by HPLC (Agilent 1200) with a C18 column and a UV detector (λ = 254 nm). The mobile phase was a mixture of water (with 0.1% formic acid) and acetonitrile (with 0.1% formic acid). HPLC was operated using a constant flow of 1.2 mL min⁻¹ and gradient elution (0–3.5 min: 25% acetonitrile; 6.5 min: 80% acetonitrile; 7.5–10.5 min: 25% acetonitrile). The total organic carbon (TOC) was measured with a Shimadzu TOC-L analyzer using the 680 °C combustion catalytic oxidation method. The relative standard deviation on TOC measurements is 2%.

2.5. Kinetic analysis and figures of merit

Different performance indicators were used for the determination of the oxidation efficiency. The TOC flux, percentage of removal (PR), removal rate (RR) and mineralization current efficiency (MCE, percentage of current directed towards the mineralization of the substrate passing through the REM) of the substrate i (molecule or TOC) were calculated as follows (Brillas et al., 2009; Garcia-Segura and Brillas, 2011; Zaky and Chaplin, 2013):

\[ \text{TOC flux} = \frac{C_{f,i} - C_{p,i}}{C_{f,i}} \times J \]  

\[ \text{PR}_i = \frac{C_{f,i} - C_{p,i}}{C_{f,i}} \times 100 \]  

\[ \text{RR}_i = \frac{C_{f,i} - C_{p,i}}{C_{f,i}} \times J \]
MCE = \frac{n F J (TOC_f - TOC_p)}{4.32 \times 10^7 m j} \times 100

where $C_{f,i}$ and $C_{p,i}$ are the concentration of the feed and permeate (g L$^{-1}$), $J$ is the permeate flux (L h$^{-1}$ m$^{-2}$), $n$ is the number of electrons consumed per molecule of substrate mineralized, $F$ is the Faraday constant, $4.32 \times 10^7$ is a conversion factor (3600 s h$^{-1}$ x 12,000 mg mol$^{-1}$), $m$ is the number of carbon atoms of the substrate studied, $j$ is the current density (A m$^{-2}$) and TOC$_f$ and TOC$_p$ are TOC of the feed and permeate (mg L$^{-1}$), respectively.

As explained in the results and discussion section, oxidation of the substrate in the feed stream was negligible.

One experiment was performed in quadruplicate in order to assess the reproducibility of experimental data.

3. Results and discussion

3.1. REM characterization

The REM was characterized in order to anticipate its suitability for the removal of organic pollutants from water. SEM images presented in Fig. 2a and b show the uniform and interconnected structure of REM pores. Pore structure was further characterized by Hg porosimetry (Fig. 2c and d). Differential and cumulative intrusion pore volume indicate a mono-modal pore size distribution ranging between 0.8 and 1.9 $\mu$m with median pore diameter of 1.4 $\mu$m. Hg porosimetry analysis also determined a porous volume of 41% and specific surface area of 0.40 m$^2$ g$^{-1}$. These characteristics correspond to microfiltration membranes allowing the separation of particles and bacteria, while dissolved species pass through the membrane. The porous structure of the REM is tailored for reaching high permeate flux. Water permeability was determined as 3300 L m$^{-2}$ h$^{-1}$ bar$^{-1}$. This is 67-fold higher than commercial Ebonex® electrodes (Vector Corrosion Technologies, Inc.) consisting primarily of Ti$_4$O$_7$, which present a bimodal pore size distribution. While micro-sized pores range between 1 and 6 $\mu$m, nano-sized pores (<10 nm) strongly increase pressure drop across Ebonex® membranes (Zaky and Chaplin, 2013). Similar flux (3208 L m$^{-2}$ h$^{-1}$ bar$^{-1}$) was obtained by Guo et al. (2016) with TiO$_2$ membranes reduced to Ti$_4$O$_7$ using a thermal method at 1050 ºC under 1 atm H$_2$ during 50 h (Guo et al., 2016). The median pore size diameter of these membranes was 2.99 $\mu$m, however, the 70 $\mu$m thick active layer with smaller pores could not be characterized by Hg porosimetry.
Porosimetry. Detailed study on limiting mechanisms for electro-oxidation showed three different behaviors according to permeate flux through the membrane (Zaky and Chaplin, 2013; Guo et al., 2016). Mass transport rate is the key parameter since electro-oxidation reactions occur at the membrane/pore surface. At low permeate flux and convection rate, diffusion can limit organic compound oxidation, similarly to conventional plane configurations (parallel plate electrodes). Convection-enhanced mass transport of pollutants is observed at higher permeate fluxes and the diffusion boundary layer is eliminated (Guo and Vecitis, 2012; Zaky and Chaplin, 2013). By continuously increasing the permeate flux, an ultimate limitation can finally arise from the electron transfer rate. This kinetic limitation depends on the nature of the substrate to oxidize, intrinsic kinetic reactivity of the REM and current supplied. From the comparison to values reported by Guo et al. (2016) with a similar configuration, it is anticipated that the high permeability of the REM used during this study avoid diffusion limitations during electro-oxidation of organic compounds, even at the lowest flux used (TMP = 40 mbar; J = 120 L h⁻¹ m⁻²).

The electrochemical performance of REM also tightly depends on the nature of the Magnéli phase. Comparison of XRD data of the REM and TiO₂ standards shows the presence of a mixture of Ti₄O₇ and Ti₃O₅ phases (Fig. 2e, f and 2g). None studies focused on the electrochemical performance of Ti₃O₅ for organic compound oxidation but the most conductive Magnéli phase is known to be Ti₄O₇ and recent studies demonstrated that this phase is able to produce *OH from water oxidation (Guo et al., 2016; Ganiyu et al., 2016, 2017). Overall, physical characteristics of REM synthesized from carbothermal reduction of TiO₂ show that this material is tailored for water treatment applications, particularly for oxidation of organic pollutants. Moreover, this innovative synthesis method has the great advantage to reduce thermal treatment times and to avoid H₂ consumption.

### 3.2. Electrochemical oxidation of organic compounds

The efficiency of the REM for mineralization of organic molecules was assessed using various model compounds involving different electro-oxidation mechanisms. Hydroxyl radicals generated at the surface of a BDD anode are able to participate to the oxidation of OA (Martinez-Hüitle et al., 2004; García-Segura and Brillas, 2011). However, based on the very low absolute rate constant for the reaction of *OH with OA (∆ = 1.4 × 10⁶M⁻¹ s⁻¹ for oxidation of PCT by *OH (Weiss et al., 2007; Ferro et al., 2010) – Guo et al. (2016) considered that OA could be used as a direct oxidation probe. As regards to PHE and PCT, both DET and *OH mediated oxidation can participate to the mineralization (rate constant of 1.4 × 10⁴mol⁻¹ s⁻¹ for oxidation of PCT by *OH (Land and Ebert, 1967); rate constant of 9.8 × 10⁸mol⁻¹ s⁻¹ for oxidation of PHE by *OH (Bisby and Tabassum, 1988)). Besides, poisoning effect or anode fouling from electro-polymerization is often observed during electro-oxidation of PHE at the surface of anode materials with low O₂ overpotential, i.e. when oxidation mainly occurs by DET (Belhadj Tahar and Savall, 2009a, 2009b; Panizza and Cerisola, 2009).

Preliminary studies without current supply showed the absence of adsorption of these organic compounds in membrane pores and other parts of the pilot. It was also observed that oxidation mainly resulted from convection through the REM. Results from an electro-oxidation experiment at 30 mA cm⁻² and TMP = 40 mbar during 90 min showed that >98% degradation and mineralization of 0.18 mM (TOC = 18 mg L⁻¹) of PCT was achieved in the permeate along the experiment, while PCT and TOC concentration in the feed stream only decreased by 9% and <5% (after 90 min), respectively. Molecules oxidized at the outer surface of the REM are not accumulated in the retentate because back diffusion is low compared to forward convection (Gao and Vecitis, 2012). Thus, reaction in the feed stream was neglected, compared to the reaction through the REM. A quasi-permanent steady-state of both feed and permeate flux/concentration was reached few minutes after anodic polarization of the REM. Permeate was sampled until the same volume of solution was filtered (350 mL) for all experiments. Therefore, each data point reported in Figs. 3.4 and 7b refers to a single experiment. The efficiency was calculated by comparing the concentration in the collected permeate and in the feed stream (considered as constant).

One experiment (TMP = 100 mbar; [PCT] = 0.33 mM; TOC = 32 mg L⁻¹; J = 15 mA cm⁻²) was performed in quadruplicate in order to assess the reproducibility of experimental data. Low standard deviations were obtained for TOC flux (11.9 ± 0.4 g m⁻² h⁻¹), percentage of TOC removal (60 ± 3%), TOC removal rate (7.1 ± 0.3 g m⁻² h⁻¹) and MCE (49 ± 1%).

#### 3.2.1. Oxidation of oxalic acid

Fig. 3 shows results obtained for the two-electrode oxidation of OA to CO₂ and H₂O at 15 mA cm⁻². For example, 75% of TOC removal from a solution containing 354 mgC L⁻¹ was achieved after a single passage through the membrane with a permeate flux of 139 L h⁻¹ m⁻² (TOC flux = 49 g h⁻¹ m⁻²). Results from increasing concentrations of oxalic acid (TMP = constant = 40 mbar) are compared with results from increasing TMP (C₁ constant = 18 mg C L⁻¹). An ultimate current limitation was drawn according to the maximum amount of electron that can be used for oxidation of OA (black solid lines in Fig. 3). This depends on the amount of electron supplied to the system (current density) and on the amount of OA available for oxidation (TOC flux).

Both OA concentration and permeate flux influences the TOC flux through the REM. When experiments were performed at the same TMP (40 mbar) and increasing OA concentrations (C₁ = 18–800 mg C L⁻¹), TOC removal rate and MCE reached a plateau around 47 g m⁻² h⁻¹ and 72%, respectively, indicating that a kinetic limitation was reached. At relatively low OA flux (<30 g m⁻² h⁻¹), almost full mineralization of OA was achieved. However, at the highest OA flux, the kinetic limitation observed for OA oxidation could not reach the ultimate current limitation (MCE = 100%). In fact, a portion of electrons is also consumed in oxygen evolution reactions i.e. the generation of O₂ and *OH that do not react or react slowly with OA. Maximum TOC removal rate of OA was 4.9-fold higher than the maximum reported in the literature using REM at a constant anodic potential of 2.94 V (47 vs 9.6 g m⁻² h⁻¹), most probably because of the higher current density used in this study leading to an increase of the kinetic limit (Guo et al., 2016). Different behavior was observed when experiments were performed at constant OA concentration (C₁ = 18 mg C L⁻¹) and increasing TMP (i.e. increasing permeate flux). TOC removal rate and MCE of OA followed a bell curve with maximum of 14 g m⁻¹ h⁻¹ and 21% achieved for a TOC flux of 29 g m⁻² h⁻¹ (J = 1650 L m⁻² h⁻¹). Similarly to the behavior observed with increasing OA concentration, OA oxidation also reached a kinetic limit when increasing the permeate flux. The difference actually lies in the increase of oxygen evolution reactions. High permeate fluxes increased O₂ gas bubbles sweeping from the REM surface, thus resulting in the promotion of the competitive reaction of oxygen evolution. Guo et al. (2016) reported the same behavior in a cross-flow filtration configuration using TiO₂ REM from thermal reduction of TiO₂ under H₂. Lower TOC removal rate (9.6 g m⁻² h⁻¹) but higher MCE (84%) was achieved for an optimal value of OA flux of 13.5 g m⁻² h⁻¹ (J = 561 L m⁻² h⁻¹). Lower TOC removal rate and optimal value of permeate flux is ascribed to the lower current density (lower electron transfer rate). However, higher maximum MCE was reached because O₂ gas bubbles generation and sweeping
was reduced because of the lower current density and lower optimal value of permeate flux, respectively.

3.2.2. Oxidation of paracetamol

PCT is an interesting model organic pollutant because both DET and \(^\text{OH}\) mediated oxidation is required in order to achieve efficient mineralization of the molecule. On one hand, PCT presents a high rate constant for oxidation by \(^\text{OH}\) (9.8 × 10^8 \text{M}^{-1} \text{s}^{-1}) and the well-known degradation by-product 1,4-benzoquinone is strongly refractory to DET (Brillas et al., 2005). On the other hand, OA and some other short-chain carboxylic acids generated as organic intermediates have lower oxidation rate by \(^\text{OH}\) and faster mineralization can be achieved by DET (Weiss et al., 2007).

The efficiency of PCT degradation and mineralization by the REM is shown in Fig. 4. Very high degradation rate of PCT was obtained, e.g. >99.9% degradation of PCT was reported until TOC flux reach 7, 14 and 20 g m\(^{-2}\) h\(^{-1}\) at 6, 15 and 30 mA cm\(^{-2}\), respectively. However, higher efficiency was observed at high concentration/low TMP compared to experiments conducted at low concentration/high TMP. There is actually a competition between the reaction of free \(^\text{OH}\) with each other and with organic compounds (Kapaïka et al., 2008). The dimerization reaction of \(^\text{OH}\) decreases mineralization efficiency through the consumption of \(^\text{OH}\) in non-organic events (scavenging reactions). Particularly, \(^\text{OH}\) dimerization has been observed to form hydrogen peroxide, and subsequently O\(_2\) (Kapaïka et al., 2009). Therefore, the increase of O\(_2\) gas bubble sweeping at high permeate flux might lead to a shift of the equilibrium of these reactions and could promote the scavenging reactions of \(^\text{OH}\) dimerization.

Using PCT as model compound, the ultimate current limitation for TOC removal (by assuming MCE = 100%) is 5.7, 14.2 and 28.3 g m\(^{-2}\) h\(^{-1}\) at 6, 15 and 30 mA cm\(^{-2}\), respectively. Compared to experiments with OA at the same current density (ultimate current limitation for TOC removal was 65.8 g m\(^{-2}\) h\(^{-1}\), at 15 mA cm\(^{-2}\)), a lower amount of TOC can be removed because more electrons are required to mineralize 1 mole of carbon (4.6 moles of e\(^{-}\) per mole of C for PCT and 1 mole of e\(^{-}\) per mole of C for OA).

When experiments were performed at the same TMP (40 mbar) and increasing PCT concentrations (\(C_1 = 18-220 \text{mg C L}^{-1}\)), TOC removal rate and MCE did not reach a plateau; bell curves were observed and maximum MCE were reached for TOC flux around 5 (MCE = 49%), 8 (MCE = 47%) and 16 g m\(^{-2}\) h\(^{-1}\) (MCE = 43%) at 6, 15 and 30 mA cm\(^{-2}\), respectively. Much higher MCE was obtained compared to a previous study using boron-doped diamond plate electrode (Brillas et al., 2005). The highest maximum TOC removal rate was obtained at high current density (12 g m\(^{-2}\) h\(^{-1}\) at 30 mA cm\(^{-2}\)) because of the increase of the electron transfer rate. However, the different bell curves observed for the MCE at different current density clearly overlapped, meaning that the current density has to be adapted to the concentration of PCT in order to achieve maximum MCE. A dark-orange color was also observed in the permeate for experiments performed with operating conditions corresponding to the downward phase of the bell curve. The decrease of oxidation efficiency at high PCT concentration can be explained by taking into consideration the different oxidation mechanisms occurring in the REM according to the depth into the porous electrode. Jing et al. (2016) provided an approximation of the potential distribution in an ultrafiltration TiO\(_2\) REM, based on the analytical solution developed by Lasia (2008). The resistance of the electrode material and the concentration polarization of electroactive species were neglected. The main parameters influencing the calculation were pore radius, specific solution resistance, exchange current density and overpotential. The simulation showed that the potential dramatically drops in a cylindrical pore and reaches a value below the one required for \(^\text{OH}\) production at the
Results from increasing concentrations of PCT (TMP = constant = 40 mbar) are compared with results from increasing transmembrane pressure (TMP) (C = constant = 18 mgCL⁻¹). Experiments have been performed at 6, 15 and 30 mA cm⁻², [Na₂SO₄] = 50 mM. Solid and dotted lines represent piece-wise interpolation and are included for easier reading of the figure.

Higher permeate flux due to the increase of the electron transfer rate. However, the upward phase of bell curves observed during theses experiments shows that the efficiency of the electro-oxidation process for the removal of low concentration of organic compounds can be strongly improved by increasing the permeate flux to an optimal value, thanks to the increase of the convective mass transport of organic pollutants towards the anode surface.

A particular attention was also given to the evolution of the permeate flux during experiments performed at 30 mA cm⁻² (Fig. 5). Using only the supporting electrolyte, permeate flux was observed to increase during the first 3–5 min of filtration due to the electro-osmotic flow. The electric field results in enhanced bulk liquid motion due to the net surplus of ions in the diffuse part of the electrical double layer formed at the surface of charged pores (Bowen and Clark, 1984; Huisman et al., 1998).

This effect decreases at lower current density. Then, the permeate flux dropped around 65% of the initial water flux, followed by a period of stabilization (or slight decrease) between 10 and 90 min. The release of O₂ bubbles in the bulk from oxygen evolution at the anode reduces the liquid bulk volume active for filtration. In filtration mode, O₂ gas bubbles are continuously swept from the anode surface and a steady state is reached between the amount of O₂ produced and flushed out from the REM. When current was not anymore supplied, the permeate flux was restored to its initial value in 20 min, once all O₂ gas bubbles were flushed out. Very similar behavior was observed using low concentration of PCT. The permeate flux was restored to its initial value and none permeability loss was observed even after several successive experiments. At high concentration of PCT, the
The evolution of the permeate flux was slower and stabilized around 80% of the initial flux, indicating a lower production rate of O₂ gas bubbles. In fact, the increase of the concentration of PCT favors the reaction of \( ^* \text{OH} \) with organic compounds and hinders the reaction of \( ^* \text{OH} \) dimerization as well as subsequent reactions leading to the formation of O₂ (Fig. 6). Besides, PCT may also increase the over-potential of oxygen evolution, which inhibited the reaction of oxygen evolution. Further electrochemical characterizations would be required in order to clarify the explanation of this phenomenon. The evolution of the permeate flux was also observed to depend on the current density (data not shown). The higher the current density, the higher the drop of the permeate flux because of the increase of oxygen evolution reaction and generation of a greater amount of O₂ bubbles.

3.2.3. Phenol electro-oxidation

Different results were observed during electro-oxidation of PHE. Permeate flux continuously decreased during the whole experiment and could not be restored to its initial value when current was not anymore supplied. Permeability loss of the REM after 90 min of electro-oxidation is reported in Table 1, according to PHE concentration and current density. REM fouling observed during the treatment of PHE with a high ratio between the PHE concentration and the current density can be explained by taking into account similar electro-oxidation mechanisms than those reported in this study for PCT. When the concentration of PHE is too high with respect to the current supplied, \( ^* \text{OH} \) produced in the top layer of the REM are not sufficient to oxidize all PHE molecules. Therefore PHE and degradation by-products can react in the deeper reactive layer.

Fig. 5. Evolution of the permeate flux during the treatment of paracetamol (PCT) at 17.7 mgC L⁻¹ or 141 mgC L⁻¹, phenol (PHE) at 141 mgC L⁻¹ and in the absence of organic compound. Current density is set at 30 mA cm⁻² during 90 min, then, none current was supplied during 50 min [Na₂SO₄] = 50 mM. Solid lines represent piece-wise interpolation and are included for easier reading of the figure.

Fig. 6. Recapitulative scheme of the main electro-oxidation mechanisms of organic compounds in the reactive electrochemical membrane.
3.2.4. Evolution of the pH
The evolution of the pH was monitored both in the permeate and in the retentate (Fig. 7a). During an experiment at 30 mA cm⁻² and [PCT] = 0.18 mM, it was observed a rapid increase of the pH in the retentate (11.1 at \( t = 10 \) min; 11.8 at \( t = 30 \) min), followed by a period of slow increase (12.2 at \( t = 90 \) min). In the permeate, pH rapidly reached a stable value around 1.8. When the feed water is passing through the membrane, \( H^+ \) generated from water oxidation at the anode (eq. (2)) are accumulated in the permeate due to the convective mass transport. On the contrary, \( OH^- \) generated at the cathode (stainless steel rod) are accumulated in the retentate. Stabilized values of pH observed in the permeate according to operating conditions are plotted in Fig. 7b. pH was mainly affected by current density and permeate flux. During experiments with PCT as model organic compounds, pH in the permeate ranged between 1.8 and 3.2 according to operating conditions. Lowest pH was observed at high current density (high amount of \( H^+ \) generated) and low permeate flux (low dilution of generated \( H^+ \)). Further investigations would be required in order to emphasize the influence of this pH gradient on electro-oxidation mechanisms and efficiency.

4. Conclusion
Unprecedented high efficiency of mineralization of organic compounds is presented in this study by using an innovative REM synthesized from carbothermal reduction of TiO₂. The homogeneous pore structure (median pore size = 1.4 μm) and suitable Magnéli phase synthesized (mixture of Ti₄O₇ and Ti₅O₉) resulted in high permeability (3300 L h⁻¹ m⁻²), high convection-enhanced mass transfer and excellent reactivity of the REM. From the high TOC removal rates (6.7 g m⁻² h⁻¹ and 4.7 g m⁻² h⁻¹ for PCT and OA, respectively, at 15 mA cm⁻²) and mineralization current efficiency (47% and 72% for PCT and OA, respectively, at 15 mA cm⁻²) achieved, the great promise of this new water treatment technology has been emphasized. Two limitations have been also highlighted. A decrease of the process efficiency has been observed at high TOC flux due to the effect of the decrease of the anodic potential according to the REM depth. Hydroxyl radicals can be only generated in the top layer of the REM. Therefore, adverse effects arises from the presence of refractory organic compounds and from electro-polymerization in the deeper layer where only DET occurs. In the case of PHE, interaction of electro-generated polymers with the...
pore surface strongly reduced the membrane permeability (fouling). Besides, at high permeate flux, a decrease of the electro-oxidation efficiency was observed due to enhanced sweeping of O2 bubbles from the anode surface, which promotes oxygen evolution reactions. Whatever the organic compound studied, current density appeared as the crucial parameter for limiting these adverse effects. Fortunately, the advantage of electro-oxidation is the possibility to easily tune current intensity in order to adapt the process to effluent concentration and/or permeate flux. While the efficiency of the REM was observed to be constant during this study, the assessment of the long-term lifetime of these electrodes is the focus of ongoing research in order to confirm the suitability of this process for water treatment. Then, the use of REM for the treatment of real effluents will be investigated since particles and colloids in real water matrix may affect the efficiency of the process.

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