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Design of a novel fuel cell-Fenton system: a smart approach to zero energy depollution

Thi Xuan Huong Le, Roseline Esmilare, Martin Drobek, Mikhail Bechelany, Cyril Vallicari, Duy Linh Nguyen, Anne Julbe, Sophie Tingly and Marc Cretin

A model azo dye pollutant, Acid Orange 7 (AO7), was removed efficiently from an aqueous medium by a smart eco-friendly Fuel Cell-Fenton (FC-Fenton) system without any external power supply. In this approach, AO7 was degraded by an electro-Fenton process at a designed cathode (Carbon Felt (CF)/porous Carbon (pC)) supplied by direct clean electrical energy from abiotic glucose oxidation at a CF/gold anode (CF@Au). The highly active cathode was fabricated by an attractive route combining Atomic Layer Deposition (ALD) of ZnO on commercial carbon felts (CFs) followed by subsequent solvothermal conversion of the metal oxide to a metal organic framework (here ZIF-8). The as-prepared composite material was further calcined at high temperature under a controlled atmosphere. A pC-based support with high specific surface area and nitrogen as a dopant was thus obtained, enhancing both conductivity and electrocatalytic properties toward H2O2 production from oxygen reduction. Degradation kinetics of AO7 (0.1 mM initial concentration) at the CF@pC cathode was monitored by UV-vis spectrophotometry and High-Performance Liquid Chromatography (HPLC) to prove the efficiency of the composite material for the degradation of such a bio-refractory model molecule. Benefitting from the H2O2 production rate (9.2 mg L−1 h−1) by the pC layer, AO7 (35.0 mg L−1) was degraded by the electro-Fenton process in acidic medium (pH = 3) with removal efficiency reaching 90% in 10 h. The durability of the system was extended for more than 2 months with an average power output of 170 mW m−2, confirming this abiotic FC-Fenton system as a promising, green, future technology for both environmental and energy-related areas, including membrane-coupled reactor systems.

1. Introduction

The scarcity of pure water worldwide is dramatically affecting the economic development of Third-World countries, but also the industrial growth as a whole. When considering water recycling and reuse, new technologies such as Electrochemical Advanced Oxidation Processes (EAOP), currently deserve strong attention. Indeed, such processes are of high interest since they are very efficient for the degradation of refractory pollutants that cannot be eliminated by conventional techniques. Among EAOP, the electro-Fenton process (EF) allows for in situ generation of highly reactive and non-selective 'OH radicals through oxygen reduction to hydrogen peroxide (eqn (1)) and its further reduction to 'OH through the Fenton reaction (eqn (2)) in the presence of Fe2+, which induces the total mineralization of pollutants.

\[
\begin{align*}
O_2 + 2H^+ + 2e^- & \rightarrow H_2O_2 \quad (1) \\
Fe^{2+} + H_2O_2 + H^+ & \rightarrow Fe^{3+} + 'OH + H_2O \quad (2)
\end{align*}
\]

In the EF process, the efficiency of pollutant degradation strongly depends on the cathode material where 'OH radicals have to be generated. Carbon-based electrodes like vitreous carbon, carbon felt (CF), and carbon sponge are commonly used as cathodes in oxygen-dissolved solutions to produce hydrogen peroxide, crucial species for the effective destruction of Persistent Organic Pollutants (POPs). The enhancement of cathode material performances for the electro-generation of H2O2 has, thus, received increasing interest through the development of composite electrodes prepared with polycrylonitrile, ethanol/hydrazine hydrate, nitrogen-functionalized carbon nanotubes, multi-walled carbon nanotubes/surfactants, polypyrrole/anthraquinonedisulphonate composite films, thermal treatments and graphene. Enhanced performance is mainly due to an increase of both the electronic conductivity of materials and the electrochemical active surface area in the developed microporous nanostructures.

In order to drive the electrons required for the oxygen reduction reaction (ORR), an electric generator is generally

\[\text{Institute Européen des Membranes, IEM UMR-5635, Université de Montpellier, ENSCM, CNRS, Place Eugène Bataillon, 34095 Montpellier Cedex 5, France. E-mail: mikhael.bechelany@univ-montp2.fr; marc.cretin@umontpellier.fr\]

\[\text{Van Lang University, 45 Nguyen Khac Nhu, District 1, Ho Chi Minh City, Vietnam}\]
required. To overcome this issue, innovative solutions have been proposed. It was previously shown that microbial fuel cells can be used for wastewater treatment as a sustainable alternative energy source. The anode side contains an electroactive biofilm working as a biocatalyst to produce electrons while the EF process takes place at the cathode. The biocatalyst has been grown on graphite rods, porous graphite felt, granular graphite, CF brush or CF. In spite of their very promising performance for both energy savings and wastewater treatment, these processes still suffer from some restrictions associated to complex experimental conditions, limited bacteria durability and low electrocatalytic electrode properties.

An attractive alternative to microbial fuel cell-driven EF processes is the direct production of electrons through the glucose oxidation process. Glucose fuel cells can be classified into three types according to the nature of the catalyst: (i) enzymatic catalyst, (ii) microbial catalyst and (iii) metal catalyst. The electro-oxidation of non-enzymatic glucose with metallic nanoparticles displays long-term stability and can be performed on electrode materials containing Ni, Pt, Pd-Rh, Pt-Ru, Au or Au–Pt–Pd.

In order to move towards a clean electrical energy system, we propose, in this study, a smart system whose working principle is based on abiotic glucose oxidation to drive the EF process. We focused our work on the development of relevant electrodes with electrocatalytic properties resulting from modifications of commercial CFs. In order for the system to operate efficiently, the electric potential difference between the two electrodes must be large enough and the redox reactions must be sufficiently fast at the electrodes to limit kinetic losses. At the anode, the glucose oxidation reaction was performed on gold particles deposited onto commercial CFs (CF@Au); whereas at the cathode, a novel porous carbon-based material (CF@pC) was prepared with relevant catalytic properties to facilitate H₂O₂ production, to extend the system durability and to ensure an easy and fast procedure, unlike microbial fuel cell-Fenton systems.

According to our best knowledge, such a concept of an abiotic fuel cell-Fenton (FC-Fenton) system has never been described in the literature and its application for efficient pollutant degradation represents an innovative green technology approach for both environmental and energy-related areas.

2. Experimental

2.1. Materials

The carbon felt was purchased from A Johnson Matthey Co., Germany. Diethyl zinc (DEZ), Zn(CH₂CH₃)₂ (95%), 2-methylimidazole (2-mim, 97% purity), chloroauric acid trihydrate (HAuCl₄·3H₂O, 99.99%), Acid Orange II sodium salt (AO7), D(+)-glucose monohydrate, sodium sulfate (anhydrous, 99.0–100.5%), iron(ii) sulfate heptahydrate (99%), methanol (99.8%) and sulfuric acid (98%) were bought from Sigma-Aldrich; potassium hexacyanoferrate (≥99%) and potassium nitrate (≥99%) from Fluka; potassium hydroxide (KOH pellets pure) from Merck and all chemicals were used without further purification.

2.2. The fuel cell-Fenton system

The FC-Fenton system was composed of two cylindrical compartments separated by a Nafion® 117 perfluorinated membrane (3.5 cm x 5 cm), as depicted in Fig. 1. The anode electrode consisted of CF@Au while the cathodic electrode consisted of porous carbon deposited onto commercial CF (CF@pC). The anode compartment contained 90 mL of glucose (0.5 M) in 0.1M KOH (pH 11.0) saturated by N₂ under magnetic stirring; whereas the cathode compartment contained 90 mL aqueous solution of AO7 (0.1 mM) with FeSO₄·7H₂O (0.2 mM) as the catalyst and Na₂SO₄ (50 mM) as a supporting electrolyte adjusted at pH = 3 and saturated by oxygen. The pH of each compartment was maintained by adding either H₂SO₄ (1 M) or KOH (1 M). The decomposition of AO7 was monitored by measuring the dye absorbance at the selected single wavelength of 485 nm, specific for azo bonds, with a Jenway 6300 spectrophotometer (Barloworld Scientific Ltd, Dunmow, UK).

Concentrations of AO7 during the experiments were measured in parallel by an HPLC system (Waters 717 autosampler and Waters 616 Pump) with a photodiode array detector (Waters 2996 photodiode array detector) fitted with a reverse phase (RP) Thermo Scientific C18 column (L = 250 mm, I.D. = 4.6 mm, and 5 μm particles).

The electrochemical performance of the FC-Fenton system was evaluated by connecting the abiotic anode and the electro-Fenton cathode under a discharge resistor of 5 ohms. The evolution of the potential of each electrode was measured independently by introducing a reference electrode (Ag/AgCl) in each compartment. The delivered current density and the output power density were derived from the resistance and the cell voltage (U), with P (mW m⁻²) = U (mV) × j (mA m⁻²).

2.2.1. Fabrication of the CF@Au anode. The anode was firstly fabricated by electro-deposition of a gold layer onto the surface of a commercial carbon felt (raw CF, Johnson Matthey Co., Germany) (4.0 cm length x 1.0 cm width x 1.0 cm...
thickness). Electrodeposition was performed using cyclic voltammetry (CV) by running 70 scans from −0.9 to 0 V versus SCE at a scan rate of 10 mV s⁻¹ in a N₂ saturated solution containing 0.05 mg mL⁻¹ chloroauric acid. The process was recorded on a µ3AUT70466 Autolab system (Eco Chemie BV, Netherlands) at a scan rate of 10 mV s⁻¹ using a three-electrode system with the CFs as the working electrode, a saturated calomel electrode (SCE) as the reference electrode and platinum foil as the counter electrode. In order to form the gold particles by dewetting, as described in our previous work, an additional thermal treatment was required. It was performed in a tubular furnace (Vecstar Ltd) under flowing nitrogen (200 mL min⁻¹) with a heating rate of 5 °C min⁻¹ up to a temperature of 1000 °C.

2.2.2. Fabrication of the CF@pC cathode. Cathodes have been prepared by an original strategy combining 3 steps: (i) atomic layer deposition (ALD) of a metal oxide (MO), (ii) solvothermal conversion of MO to a metal organic framework (MOF) and finally (iii) MOF carbonization, forming pC with a high specific surface area.

(i) A home-made ALD setup was used for depositing ZnO thin layers onto commercial CF (4.0 cm length × 1.0 cm width × 1.0 cm thickness) at 100 °C using sequential exposures of DEZ and deionized water separated by a purge with dry argon (Ar flow rate of 100 sccm). The deposition protocol was as follows: (a) 0.5 s pulse of DEZ, 30 s exposure and 50 s purge with dry Ar; (b) 2 s pulse of H₂O, 40 s of exposure and 60 s purge with dry Ar. The selected pulse, exposure and purge times were chosen both to ensure completion of the ALD surface reactions and to prevent any mixing of the reactive species. The characteristics of the ZnO ALD deposits (thickness, crystalline phase and grain/crystallite sizes) were found to be insensitive to both support morphology and composition. The growth rate per cycle of the zinc oxide layers was controlled by measuring the films thickness deposited on Si-wafer companion substrates placed in the ALD reaction chamber.

(ii) The ZnO-coated CFs were submitted to a solvothermal treatment in a closed pressure vessel (Teflon-lined stainless-steel 45 mL autoclave) containing a solution of 2-mim in methanol (10 wt% 2-mim). The autoclave was heated at 100 °C for 5 h in a conventional oven. After this solvothermal treatment, the resulting material was washed five times with methanol and dried for 2 h at 70 °C.

(iii) The as-modified CFs were further carbonized at 1000 °C for 10 h under nitrogen in order to obtain a pC layer deposited onto the commercial CFs. The resulting CF@pC cathode was expected to offer a high specific surface area, enhanced conductivity and enhanced electrocatalytic properties toward H₂O₂ production from oxygen reduction.

2.3. Material characterization

Chemical and structural characterizations of the prepared materials were performed by scanning electron microscopy (SEM, Hitachi S-4800), X-ray diffraction (XRD) (PANalytical Xpert-PRO diffractometer equipped with a X'celerator detector using Ni-filtered Cu-radiation), EDX analysis (Silicon Drift Detector (SDD), X-MaxN, Oxford Instrument) coupled to a Zeiss EVO HD15 SEM analyzer. X-ray photoelectron spectroscopy (XPS) measurements were conducted using an ESCALAB 250 Thermal Electron with AlKα (1486.6 eV). Binding energies were calibrated using the contained carbon (C1s = 284.8 eV).

2.4. Electrocatalytic activity measurements

Electrochemical analysis was performed with a µ3AUT70466 Autolab system (Eco Chemie BV, Netherlands) with a conventional three-electrode assembly composed of a SCE reference, a Pt foil counter electrode and the developed electrodes as working electrode. CV was performed in 1.0 M KNO₃ containing the redox probe Fe(CN)₆³⁻/⁴⁻ 10 mm. The electroactive surface area of the cathodes was evaluated according to the Randles-Sevcik relation.

To evaluate the catalytic activity of the CF@pC cathode towards H₂O₂ electro-generation, linear sweep voltammetry (LSV) measurements were performed in 45 mL Na₂SO₄ (50 mM) solution at pH = 3.0. Before the measurements, the solution was saturated by oxygen for 10 min and the potential was scanned from the open circuit potential to −1.2 V versus Ag/AgCl at 5 mV s⁻¹ in the three-electrode system. The electro-generation of H₂O₂ was carried out at the potential of −0.21 V vs. Ag/AgCl by chronopotentiometry, and the resulting H₂O₂ concentrations were determined by spectrophotometry using potassium titanium(IV) oxalate as a colored indicator. The absorbance of the yellow per-titanic acid complex between H₂O₂ and potassium titanium oxalate in acidic solution was measured by a Jenway 6300 spectrophotometer (Barioworld Scientific Ltd, Dunmow, UK) at λ = 400 nm through a 1 cm polystyrene cuvette. The calibration curve was obtained from standard a H₂O₂ solution that was titrated against potassium permanganate solution (0.1 N).

To evaluate the catalytic activity of the CF@Au anode towards glucose electro-oxidation, CV measurements were performed at 50 mV s⁻¹ in 0.1 M KOH solution containing 10 mM glucose. The glucose solution was stirred for 24 h before testing. The glucose oxidation potential was fixed from the LSV in 0.1 M KOH solution containing 0.5 M glucose.

3. Results and discussion

To develop an efficient FC-Fenton system, the innovation proposed in this work involves relevant modifications of commercial CFs according to two different strategies: (i) preparation of a cathode from the modification of CFs with a pC layer and (ii) preparation of an anode by electrodeposition of gold particles onto CFs. The key parameters of both steps were examined through a description of the deposits (location, quantity and physico-chemical characteristics) and investigation of the electro-catalytic activity of the as-prepared materials assembled in a FC-Fenton cell. The electrodes were assembled in a FC-Fenton cell whose characteristics were described and discussed in relation with the electrode structure and morphology, and with the degradation kinetics of the pollutant AO₇ in an aqueous solution.
3.1. Characterization of both the CF@Au anode and CF@pC cathode

In the CF@Au material, the as-deposited gold particles play the catalyst role for the direct electro-oxidation of glucose in the FC-Fenton system. The gold particles, with an average size of 300 nm calculated by the ImageJ software, are uniformly distributed onto the CFs, as evidenced on the SEM images at different magnifications (Fig. 2d compared to raw CFs in 2a) and EDX mapping results (Fig. S1†). The XRD analysis of the as-prepared electrode (Fig. 2e) features both a broad diffraction peak around $2\theta = 23^\circ$, corresponding to a hexagonal graphite structure (002), and a less intense peak at $43^\circ$. Crystalline gold particles on the surface of CF were proved through the emergence of sharp peaks at $38.2^\circ$ (111), $44.2^\circ$ (200) and $64.6^\circ$ (220), which are specific for the cubic phase of gold.\textsuperscript{31,32} Gold particles were thus successfully obtained by reduction of the chloroauric anions by electrodeposition of gold layers onto the CF surface subsequently transformed into particles by thermal treatment.

The catalytic activity of the CF@Au electrode was tested by running CVs in 0.1 M KOH solution in either the presence or absence of glucose (10 mM). In the absence of glucose (Fig. 2f), the CV of the CF@Au electrode shows two peaks relative to gold surface-oxides formation and reduction, respectively, in alkaline media.\textsuperscript{33,34} In the presence of glucose, raw CFs are inactive for glucose electro-oxidation in the investigated potential range; whereas the CF@Au electrode shows the typical voltammetric behavior characterized by three electrochemical processes: (1) formation of an adsorbed glucose layer on the gold electrode surface through dehydrogenation of the anomeric carbon in the glucose molecules,\textsuperscript{35,36} (2) oxidation of the as-adsorbed glucose forming the gluconolactone species (appearance of a large left shoulder peak around 0.3 V) and (3) re-adsorption and oxidation of another glucose.\textsuperscript{37}

The active cathode was fabricated by combining ALD of metal oxide (ZnO) on CFs followed by its solvothermal conversion to ZIF-8, and a subsequent calcination under a controlled atmosphere to forming pC-based deposits. The presence of ZnO is highlighted by the diffraction peaks at $2\theta = 31.7^\circ$, $34.4^\circ$ and $36.2^\circ$, corresponding to the diffraction lines (100), (002) and (101) of the wurtzite hexagonal ZnO phase. After 250 ALD cycles, the thickness of the ZnO layer was $\sim 50$ nm and the mean size of crystallites in the ZnO layer was 18 nm, as calculated using the Debye–Scherrer equation reported elsewhere.\textsuperscript{38} The solvothermal conversion of the as-modified CFs resulted in a ZIF-8-based composite material with ZIF-8 crystals (0.1–1 $\mu$m in size) homogeneously covering the CFs (Fig. 3a). Both ZIF-8 and ZnO were found to co-exist on the CFs, as confirmed in Fig. 3e by the characteristic XRD peaks of both ZIF-8 ($2\theta = 7.3^\circ$ (011), 10.4$^\circ$).
formula consisting mainly of micropores (pore volume area that is 9.3 times higher (calculated by Randles the CF@pC electrode exhibits an estimated electroactive surface solution at the CF@pC electrodes (Fig. 3e). Compared to raw CF, and thus, the CF@pC material was explored for the in situ electrochemical production of H2O2.

The presence of carbonaceous structures and the absence of ZnO in the resulting CF@pC material confirm that a metal-free carbon material was successfully synthesized. Besides, the elemental analysis of CF@pC by XPS indicated this as well as the presence of nitrogen (1.73%) on the porous carbon that could be attributed to N-containing functional groups like pyridinic-N (398.2 eV), pyrrolic-N (399.8 eV), quaternary-N (403 eV) and N-oxide (392 eV) (Fig. 3h), as reported elsewhere.39,40 These N-containing groups improve the hydrophilic characteristics that promote both electron transport and mass transfer, and thus, electrochemical performance toward ORR during the EF process.41,42

According to N2 physisorption measurements, it is apparent that the carbon layer on the CFs generates a porous structure consisting mainly of micropores (pore volume = 0.082 cm$^3$ g$^{-1}$) increasing the specific surface area of the CF material from 0.0915 m$^2$ g$^{-1}$ to 64 m$^2$ g$^{-1}$. As previously mentioned,41,44 the microstructure of such a cathode material can facilitate the diffusion and transformation of oxygen to H$_2$O$_2$ on the surface, and thus, the CF@pC material was explored for the in situ electrochemical production of H$_2$O$_2$.

The benefit of the pC structure on the electroactive surface area is also obvious from the increase in the electrochemical redox signal of the redox probe [Fe(CN)$_6$]$_3$$^3^-$/[Fe(CN)$_6$]$_4$$^4^-$. In solution at the CF@pC electrodes (Fig. 3e). Compared to raw CF, the CF@pC electrode exhibits an estimated electroactive surface area that is 9.3 times higher (calculated by Randles-Sevcik formula$^{29,30}$). This result may be attributed to an enhanced specific surface area, a high degree of graphitization and the presence of abundant graphitic nitrogen atoms.44 Fig. 4a shows the LSV of H$_2$O$_2$ production by running the potential of the cathode to generate H$_2$O$_2$ at a low potential in an electro-Fenton process for pollutants degradation.

### 3.2. FC-Fenton system efficiency

The FC-Fenton system was operated by connecting both the abiotic CF@Au anode and the CF@pC cathode in a two-chambers cell separated by a Nafion membrane. Fig. 4 presents the operation feature of the FC-Fenton system where direct clean electrical energy from abiotic glucose oxidation at the CF@Au electrode and ZnO in the resulting CF@pC material a

production at a constant potential of $-0.21$ V vs. Ag/AgCl. After 60 min, a stable concentration of H$_2$O$_2$ (9.2 mg L$^{-1}$) was obtained at pH = 3 whereas no H$_2$O$_2$ generation was detected at the raw CF electrodes, revealing the outstanding performance of the newly designed CF@pC cathode.

Besides, as observed in Fig. 4a, glucose oxidation at CF@Au anode starts at $-0.46$ V vs. Ag/AgCl. For efficient operation, it is thus obvious that the raw CF cathode cannot be combined with the CF@Au anode to build a FC-Fenton system due to the low electric potential difference between the two electrodes. However, the CF@pC electrode has great promise as an efficient cathode to generate H$_2$O$_2$ at a low potential in an electro-Fenton process for pollutants degradation.
the hydroxyl radicals (·OH) were formed. Fig. 4b clearly demonstrates that an average power output of 170 mW m⁻² was continuously produced for at least two months. The CF@Au electrode exhibits stable electroactivity toward glucose oxidation at around -220 mV due to the presence of the gold nanoparticles. In the cathodic compartment, regular potential oscillation takes place due to both proton consumption in ORR (eqn (1)) and exchange through the Nafion membrane, leading to pH variations affecting the cathodic potential values. As a result, pH adjustment plays a key role in maintaining a stable performance of the FC-Fenton system. The characterization of both the CF@Au anode and the CF@pC cathode before/after two months use in the FC-Fenton system has been performed by SEM (Fig. S4 and S8†), EDX mapping (Fig. S1 and S7†), XPS analysis (Fig. S2, S5 and S9†), EDX analysis (Fig. S3, S6 and S10†) and elemental composition measurements (Tables S1-S4†) in order to prove the durability of both fabricated electrodes. As observed from Fig. S4,† the gold particles were still steadily deposited on the CF@Au anode after two months use in the FC-Fenton system. In the case of the CF@pC cathode, some additional impurities (F, Si, S) at very low quantities (Table S4†) occurred after two months of system operation. This contamination comes from the dye solution continuously supplied into the cathodic compartment. However, no negative impact on the FC-Fenton efficiency was observed. These results clearly confirm the robustness of both cathode and anode electrodes, attesting to their long-term stability in the proposed fuel cell system.

The AO7 degradation was followed by HPLC analysis during electrolysis. As shown in Fig. 5b, the concentration of AO7 decreases rapidly at the early stage of electrolysis with approximately 70% conversion within 3 hours. Subsequently, degradation decelerates because the decomposition of AO7 by the EF process leads to the formation of various aromatic compounds and short-chain carboxylic acids that deactivate the ·OH radicals. Hence, the AO7 decomposition increases by only -20% in the seven following hours, resulting in about 90% elimination of AO7 after 10 h of the electro-catalytic process. On the other hand, the absorption peaks at 254 and 310 nm (spectrophotometry), attributed to aromatic amines and naphthalene in the initial mother solution, are also weakened as a result of the increased treatment time (elimination of the aromatic ring structure by ·OH – Fig. 5c). In order to evaluate the durability of the studied FC-Fenton system, the decay kinetics of AO7 was monitored over two months. The cathodic compartment was weekly replenished with fresh AO7 (0.1 mM) solution with Na₂SO₄ and 0.20 mM Fe²⁺ at pH = 3; while in the anodic compartment, the abiotic anode and the solution were kept unchanged. The breaking of the azo bond in the AO7 structure (Fig. 5a) by ·OH was identified by measuring the absorbance of the solution at λ = 485 nm. As a result of the high consistency of both gold nanoparticles and the pC layer at the anode and cathode electrodes, respectively, the cell current output remained stable during at least two months and led to a steady degradation of AO7 from water (Fig. 5d). From Fig. 4a, it could be clearly noticed that the ORR occurs at less negative potentials after modification of CF (−0.21 V vs. Ag/AgCl on CF@pC instead of −0.67 V on bare CF). This observation is of an utmost importance when applying such a cathode in the proposed FC-Fenton system. In fact, the glucose oxidation at the anode takes place at −0.46 V vs. Ag/AgCl, which would make it impossible to employ a combination of the CF@Au anode and raw CF cathode in the experimental setup. These results prove the promising efficiency of the proposed FC-Fenton cell for the zero-energy depollution systems of acidic solutions and could be further extended to membrane-coupled reactor systems. This study is currently in progress and will be the subject of our next communication.

Compared with other studies from the literature,14,15,37 the present FC-Fenton system shows remarkable advantages such as: (i) virtually zero energy costs (no need of any external power generator), (ii) simple and fast operating cell at ambient temperature with no need of an anaerobic medium (contrary to bacterial electrodes and biofilms), (iii) accelerated ORR leading to H₂O₂ production at higher potentials, (iv) low negative potentials leading to more efficient pollutants removal and finally, (v) long-term stability enabling multiple degradation cycles without decreasing the catalytic activity over time.

4. Conclusions

The present work demonstrates the potential of a smart eco-friendly approach for the removal of persistent organic pollutants (POPs) from water without using any external power input. The proposed green technology relies upon a FC-Fenton system composed of an originally designed porous CFs cathode and CF@Au anode placed in a cell with two-chambers separated by a protonic exchange membrane (Nafion). The original synthesis of the cathode electrode, based on ALD of ZnO and its subsequent solvothermal conversion to ZIF-8 and pyrolysis, produced a porous carbon material with beneficial electrocatalytic
properties for the diffusion and the reduction of oxygen into \( \text{H}_2\text{O}_2 \). The presence of gold particles on the CFs anode allowed the direct electro-oxidation of glucose to supply green electrons for the ORR. The catalytic properties of both anode and cathode induce a stable output current density of 360.3 ± 51.5 mA m\(^{-2}\) at 400 ± 50 mV that could be maintained for long period of time. As a consequence, 90% of the initial concentration of the azo dye pollutant AO7, identified by HPLC analysis, was eliminated upon extended EF degradation for 10 h; and the cell power output of 170 mW m\(^{-2}\) was stable at least for two months. Hence, this first proof of concept of an abiotic FC-Fenton system demonstrates high efficiency towards pollutant degradation with a huge potential in both energy-related areas and environmental protection.

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