Electrochemical jet-cell for the in-situ generation of hydrogen peroxide

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Abstract

In this work, a proof of concept of a novel reactor design for the in-situ electrochemical production of H_2O_2 from oxygen reduction reaction is presented for the first time. The innovative design incorporates a venturi-based jet aerator to supply atmospheric oxygen without additional energy consumption to a 3D flow-through modified carbon felt (CF) cathode. Preliminary experiments confirmed that electro-generation of hydrogen peroxide is possible in a system as the one proposed. Comparison with a flow-by cell with a gas diffusion cathode under similar conditions revealed that current efficiency towards hydrogen peroxide accumulation is even higher (72 vs 65 % at 1 h) than in the case of the conventional system. Jet aerator stands as a promising oxygen supply thanks to its excellent performance and both low investment cost and energy consumption. Considering all the above, the electrochemical jet cell stands as a rather promising design for the efficient hydrogen peroxide electro-generation.

Keywords: hydrogen peroxide, gas diffusion electrode, jet electrochemical cell, carbon felt
1. Introduction

Hydrogen peroxide is a widely used chemical in industries such as textile, pulp and paper bleaching, electronics, metallurgic or environmental applications as it is the case of water treatment, among others. Hydrogen peroxide is mostly manufactured on an industrial scale by means of the Riedl-Pfleiderer process (anthrahydroquinone autoxidation) since its implementation in 1940 by IG FarbenIndustrie [1]. Despite the fact that the yield is high and the process is commercially profitable, it does exhibit various disadvantages in environmental terms on account of its high energy consumption and large amounts of waste generated [2]. In addition, the product has to be transported over long distances from the production point to the end user. An alternative option for the generation of hydrogen peroxide in aqueous medium is the in-situ production from dissolved oxygen reduction reaction (ORR), according to Equation 1:

\[ \text{O}_2 + 2e^- + 2\text{H}^+ \rightarrow \text{H}_2\text{O}_2 \] (1)

This way of production presents various advantages, e.g., ease of automation, mild reaction conditions, its applicability on a small scale and it being fully electricity-powered. Moreover, the electrochemical method avoids all the issues related to transport, storage and handling of this hazardous chemical [3].

The production rate and current efficiency is often limited by the low solubility of oxygen in water which is approximately 40 mg dm\(^{-3}\) (oxygen atmosphere) or 8 mg dm\(^{-3}\) (air atmosphere) at 1 atm, 25 °C and deionized water, depending basically on its partial pressure [4]. This problem has been addressed by substituting conventional electrodes for gas diffusion electrodes (GDEs) in which air or oxygen is directly supplied to the cathode without the need for it to be dissolved in the electrolyte [5]. However, \(\text{O}_2/\text{air}\) utilization efficiencies are extremely small, in the order of \(< 0.1\%\) [6], resulting in a waste of energy and oversizing of compressors in the event of industrial application.

Because of this, an appropriate reactor design is of utmost importance. In the recent years, different authors have proposed novel electrochemical reactors based on flow-through electrodes because of the favorable mass transport rate [7-9]. In the particular case of electrochemical \(\text{H}_2\text{O}_2\) reactors, it is important to ensure a continuous and sufficient supply of oxygen to the electrode, i.e., minimizing mass transport limitations to the minimum energy consumption and maximum simplicity.
In this work, an innovative approach to electrochemical cell designing is presented for the first time. The key feature is a jet aerator to supply oxygen to the system eliminating the need for an external compressor. Also, a 3D flow-through modified carbon felt cathode to achieve high space-time yield is used. With this novel approach to ORR reactor design, higher \( \text{H}_2\text{O}_2 \) generation rate and lower energy consumption is intended compared to conventional GDEs systems.

2. Materials and methods

2.1 Electrolyte solution

All the experiments were performed in a Milli-Q water solution with a concentration of 50 mM Na\(_2\)SO\(_4\) as widely used in literature [10-12], supplied by Panreac and a total electrolyte volume of 1 dm\(^3\).

2.2 Experimental set-ups

Two different experimental set-ups were used: the novel jet electrochemical cell (henceforth referred to as \textit{jet cell}) and a conventional flow-by cell with a gas diffusion cathode (henceforth referred to as \textit{GDE cell}). A conceptual picture of both cells is shown in Figure 1.

Figure 1. Conceptual picture of the electrochemical reactors. a) Jet cell b) GDE cell.
In both electrochemical cells, a modified carbon felt (CF) electrode was used as the cathode. The electrode was modified following a similar procedure to the one described by Yu et al. [11] which mainly consists in immersing a carbon felt piece into an ink in an ultrasonic bath. The composition of the ink was slightly modified with respect to Yu’s and follows the receipt: 10 mg/mL-H\(_2\)O of carbon black (Vulcan ® XC72R), 50 mg/mL-H\(_2\)O of polytetrafluoroethylene (60% Teflon ® emulsion solution from ElectroChem, Inc.) and 20 mL/L-\(n\)-butanol. On its part, a titanium anode with active coating based on iridium oxides (Tianode, India) was used as counter electrode. Apart from the cell, each bench-scale installation is properly configured with auxiliary equipment (reservoir tanks, power supply, pump, rotameter, etc.)

2.3 \(\text{H}_2\text{O}_2\) measurement

The hydrogen peroxide concentration was measured by the potassium titanium (IV) oxalate method [13] according to standard DIN 38 409, part 15, DEV-18. The titanium solution was supplied by Fluka and the absorbance was determined at \(\lambda = 407\) nm by means of an Agilent 300 Cary series UV-Vis spectrophotometer.

3. Theory

The design of the novel electrochemical jet cell incorporates a key feature with the objective of enhancing efficiency of hydrogen peroxide electro-generation: a venturi-based jet aerator as oxygen supply to a 3D flow-through modified carbon felt cathode with high space-time yield.

Regarding the former, a jet aerator is used to create a suction that entrains air into the water flow. This form of aeration does not need extra equipment, such as a compressor, representing a huge advantage with respect to conventional GDE systems because of the savings in investment and operating costs. Because of the high velocity in the throat of the jet aerator, the bubbles are broken into smaller pieces as a consequence of the high shear rates [14]. A fraction of the air flow sucked in the system is dissolved, saturating water in oxygen, while the rest of it remains undissolved in the form of air bubbles. This way, the water is supersaturated, i.e., it contains a concentration of oxygen higher than the one of the equilibrium described by Henry’s law for a short period of time.

For hydrogen peroxide electro-generation, the oxygen supplied to the system must also be in contact with electrons and protons at some point, according to Equation 1. The
physical place where those three reagents coincide can be referred to as triple-contact point (TCP). An adequate cathode must provide a large surface area with many TCPs where reactants are supplied in sufficient quantities. Considering this, a graphite felt was selected as the cathode material since it possesses a large three-dimensional surface [10, 11]. Nevertheless, the H$_2$O$_2$ generation rate was reported to be low in bare graphite felt, requiring changes to increase its productivity. Yu et al. reported a simple method to modify this material, adding a carbonaceous powder (carbon black) and polytetrafluoroethylene (PTFE) as a hydrophobic agent [11]. By doing this, the surface area is increased [7] (and so do TCPs) and hydrophobicity facilitates gas diffusion into the inner parts of the porous structure favoring the oxygen supply to active sites. The last property is truly important since hydrogen peroxide electro-generation kinetic from ORR is normally limited by the low concentration of oxygen in water [4] and, thus, being O$_2$-mass transfer controlled [15]. To minimize this limitation, the water flow is forced to pass through the porous structure of the cathode, as shown in Figure 1.

These special hydrodynamic conditions are used to achieve a high space-time yield in small concentrations of reactants by creating favorable mass transport conditions [7-9]. In fact, at similar linear empty tube flow velocities, the mass transport coefficient is expected to be 5 to 6 times higher than for a plate electrode with parallel flow [16]. All the aforementioned characteristics configure a promising reactor design for fast and efficient hydrogen peroxide electro-generation.

4. Results and discussions

A concept-proof experiment was conducted in the jet cell to demonstrate whether the electro-generation of hydrogen peroxide is possible or not. This experiment was carried out in 1 dm$^3$ 0.05 M Na$_2$SO$_4$ medium at a current density of 50 mA per cm$^3$ of electrode material. All the experiments were performed in triplicate to ensure reproducibility being the value in Figure 2 the average of the three experiments. It shows a logarithmic trend line, which is the typical dynamic response in non-divided electrochemical cells for the electro-generation of H$_2$O$_2$ [5, 17]. High concentrations and current efficiencies of 450 (75 %), 725 (61 %) and 960 (51 %) mg H$_2$O$_2$ dm$^3$ were obtained after 1, 2 and 3 hours of electrolysis, respectively. Therefore, the proof of concept is positive and it is demonstrated that hydrogen peroxide can be generated in a flow-through modified carbon felt cathode by using a jet aerator as oxygen supply.
Figure 2. Concentration curves for hydrogen peroxide generation in the jet cell ■ Jet cell □ GDE cell. Current density 50 mA cm\(^{-3}\). Electrolytic medium: 0.05 M Na\(_2\)SO\(_4\). Inter-electrode gap: 18 mm

For comparative purposes, the hydrogen peroxide electro-generation was also performed in a conventional flow-by cell using the same modified carbon felt electrode as a gas diffusion electrode. To compare on equal terms, current density, anode material, electrolytic medium, inter-electrode gap and ratio electrical charge/electrolyte volume was kept constant in both cells. Considering Figure 2, it can be observed that the jet cell accumulates a higher concentration of hydrogen peroxide than in the case of GDE cell. The accumulation rate is the difference between the generation rate at the cathode (Eq. 1) and the destruction rates, mainly as a consequence of a cathodic reduction (Eq. 2) and an anodic oxidation (Eq. 3) [3, 18]. Destruction reactions leads to a severe decrease in current efficiency since the applied electric charge not only is not used in producing hydrogen peroxide molecules but, even worse, it decomposes the already existing ones. The current efficiency can also be diminished by other parasitic reactions that do not involve hydrogen peroxide destruction such as hydrogen evolution (Eq. 4):

\[
H_2O_2 + 2H^+ + 2e^- \rightarrow 2H_2O \quad (2)
\]

\[
H_2O_2 - 2e^- \rightarrow O_2 + 2H^+ \quad (3)
\]
At the beginning of the experiment, the destruction rates were low because the hydrogen peroxide concentration was also low and the destruction may suffer from mass transfer limitations. It is precisely at this time when the generation and accumulation rates are quite similar and both reactors can be properly compared. According to Figure 2, the tendency lines are overlapped in the first instants of the experiments indicating that the generation is quite similar and the jet cell is as efficient as the conventional GDE cell for the electro-generation of hydrogen peroxide.

The role of the jet aerator was further investigated in a non-stop experiment under three different aeration conditions. Before starting, nitrogen was bubbled in the reservoir tank for 10 minutes to strip all the oxygen and the valve controlling air flow into the system through jet aerator was closed. Once the O\textsubscript{2} concentration was set to zero, the experiment began and the electrolysis was carried out for 30 minutes. At minute 30, nitrogen bubbling was substituted by air sparging and the electrolysis continued for another 30 minutes. Finally, air sparging was stopped and during the last 30 minutes of electrolysis, the oxygen was supplied by using the jet aerator. The concentration-time curves obtained in this experiment are shown in Figure 3.
Figure 3. Hydrogen peroxide generation under different aeration conditions at different current densities. ■ 50 mA cm\(^{-3}\) □ 75 mA cm\(^{-3}\). Electrolytic medium: 1 dm\(^3\) 0.05 M Na\(_2\)SO\(_4\).

The electrolysis under nitrogen bubbling led, as expected, to a very low concentration of hydrogen peroxide since no oxygen is available for electro-reduction in the cathode. For the next 30 minutes, hydrogen peroxide concentration was slightly higher but remains very low, thus, obtaining poor current efficiencies, indicating that the dissolved oxygen is not sufficient for an efficient hydrogen peroxide generation. It is not until the jet aerator was used when hydrogen peroxide concentration takes off, especially in the case of the higher current density. The cell potential fluctuates slightly (± 0.2 V) when bubbles (non-conductive) are flowing through the cathode, without resulting in an operational problem. This tremendous improvement in efficiency is associated with the formation of small to medium air bubbles as a consequence of the turbulence generated in the throat of the venture aerator [19] which may reach TCPs along the entire cathode surface by diffusing through the pores thanks to the hydrophobicity of the PTFE added to modify the carbon felt [20].
5. Conclusion

Hydrogen peroxide electro-generation in a new electrochemical reactor design based on a jet aerator and a flow-through modified carbon felt (CF) cathode is demonstrated. The comparison with a conventional GDE cell confirmed that this new approach is highly efficient as demonstrated by its high current efficiency of 75, 61 and 51% at 1, 2 and 3 h, respectively. The jet aerator proved to be an excellent oxygen supply with a minimum investment and energy costs. The electrochemical jet cell design is truly promising and could mean a giant leap forward in the applicability of electro-generated hydrogen peroxide as a green oxidant to environmental remediation in general, and water treatment in particular.

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References


