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Highly Efficient Electrochemical Production of Hydrogen Peroxide Using the GDE Technology

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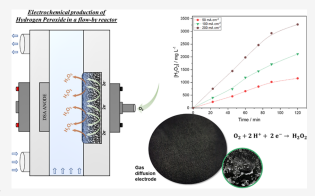


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s **ABSTRACT:** This work examines the role of oxygen supply in the 6 improvement of the hydrogen peroxide (H_2O_2) electrochemical 7 production efficiency and the generation of high H_2O_2 concentrations 8 in electrochemical processes operated in a discontinuous mode. To 9 conduct this study, a highly efficient Printex L6 carbon-based gas 10 diffusion electrode (GDE) as a cathode was employed for the 11 electrogeneration of H_2O_2 in a flow-by reactor and evaluated the 12 effects of lowering the operation temperature (to increase solubility) 13 and increasing the air supply in the system on H_2O_2 electrogeneration. 14 The results obtained in this study show that unlike what is expected in 15 flow-through reactors, the efficiency in the H_2O_2 production is not 16 affected by the solubility of oxygen when GDE is employed in the 17 electrochemical process (using the flow-by reactor); i.e., the efficiency



18 of H_2O_2 production is not significantly dependent on O_2 solubility, temperature, and pressure. The application of the proposed 19 PL6C-based GDE led to the generation of accumulated H_2O_2 of over 3 g L⁻¹ at a high current density. It should be noted, however, 20 that the application of the electrocatalyst at lower current densities resulted in higher energy efficiency in terms of H_2O_2 production. 21 Precisely, a specific production of H_2O_2 as high as 131 g kWh⁻¹ was obtained at 25 mA cm⁻²; the energy efficiency (in terms of H_2O_2 production) values obtained in this study based on the application of the proposed GDE in a flow-by reactor at low current 23 densities were found to be within the range of values recorded for H_2O_2 production techniques that employ flow-through reactors.

1. INTRODUCTION

The past few decades have seen a dramatic increase in the search for new technologies that are capable of producing chemical oxidants at substantial concentrations and in a highly efficient manner. Hydrogen peroxide (H_2O_2) is a highly efficient, eco-friendly chemical oxidant, which has a wide range of applications in different sectors. 1 H_2O_2 has a high reduction potential ($E^0 = 1.77$ vs standard hydrogen electrode, SHE) and produces nontoxic water when applied; as a result, this oxidant is widely applied in several industrial processes, including the synthesis of chemical products, paper bleaching, and wastewater treatment. $^{2-6}$ As part of the efforts to combat the SARS-SoV-2 virus, H_2O_2 was widely employed as a reagent in the formulation of decontamination and disinfection products and for the cleaning of contaminated respiratory masks for reutilization due to its antimicrobial properties. 3

The range of application of H_2O_2 has progressively increased 40 in recent years, and the annual consumption of this oxidant is 41 estimated to increase to 6 million tons in 2027. Most of 42 the current production of H_2O_2 occurs through the 43 anthraquinone process. To meet the increasing demand for 44 H_2O_2 , alternative efficient techniques for the production of the

oxidant are currently being studied, and one of the techniques 45 that have been found to be highly promising is the 46 electrochemical production of H_2O_2 via oxygen reduction 47 reactions (ORR)—see eq 1.2,4,9-11 The ORR technique 48 employs oxygen as the raw material in the electrochemical 49 process. Over the past few years, there has been a huge interest 50 among researchers in the use of ORR via the 2-electron 51 pathway for the electrogeneration of H_2O_2 ; this technique has 52 become extremely popular because it is an energy-intensive 53 multistep process, which has been proven to have the following 54 advantages: high efficiency, good operational safety, and low 55 environmental impact. There have been several reports in 56 the literature regarding the mechanism of operation of the 57 ORR process. As demonstrated in the literature, through the 58 application of carbon-based cathode materials, O_2 is easily 59

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(B) Flow-by Electrochemical Reactor Setup

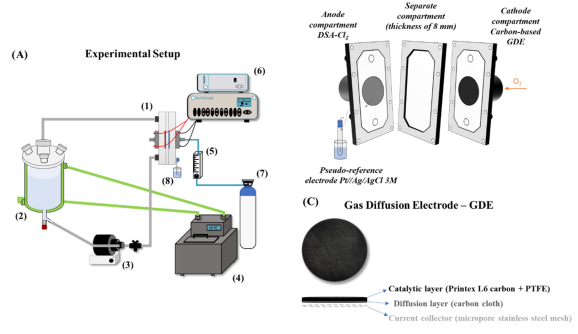


Figure 1. (A) Experimental setup: (1) electrochemical cell, (2) reservoir tank, (3) peristaltic pump, (4) thermostatic bath, (5) gas flowmeter, (6) power supply, (7) O₂ gas cylinder, and (8) pseudo-reference electrode Pt//Ag/AgCl 3M. (B) Flow-by electrochemical reactor setup and (C) gas diffusion electrode composition.

 60 reduced to $\rm H_2O_2$ via the transfer of two electrons at a potential 61 of 0.682 V vs $\rm SHE^{4,11-13}$

$$O_2 + 2H^+ + 2e^- \rightarrow H_2O_2$$
 (1)

63 Some studies reported in the literature have pointed out 64 different ways to improve the production of H_2O_2 through 65 ORR via the 2-electron pathway; some of these ways include 66 the following:

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- (i) Improvement of the catalytic properties of the cathode; this can be done by studying and developing new highly efficient cathode materials based on carbon black (CB) or through the addition/doping of organic or inorganic catalysts into carbonaceous materials; 4,11,14–18
- (ii) Improvement of oxygen supply in the electrochemical cell since the low solubility of oxygen in the cell causes the efficiency of the process to be controlled by diffusion. As reported in the literature, to help tackle this problem, some important progress has been made by
 - (a). performing the electrochemical operation under high pressure and at low temperature so as to improve oxygen solubility. 1,6,19-21
 - (b). optimizing the shape/configuration of carbon-based cathodes by employing gas diffusion electrodes (GDEs) in flow-by electrochemical reactors instead of flow-through electrode reactors. 1,18,19,22
 - (c). the design of more efficient electrochemical reactors with enhanced turbulence. 1,18,19

88 So far, when it comes to the development and application of 89 techniques for the production of H_2O_2 through ORR, most of 90 the effort has been devoted toward the development of new 91 cathodic materials (which use lab-scale cells) or specific 92 applications of H_2O_2 (such as cells for wastewater treatment). $^{19,22-24}$ No substantial effort has been devoted toward 93 investigating and developing new efficient techniques that are 94 capable of producing H_2O_2 at high concentrations. 14,16 Thus, 95 the present work aims to develop and optimize the operational 96 parameters of a new carbon-based gas diffusion electrode 97 (GDE) applied in a flow-by electrochemical reactor with a 98 view to obtaining high H_2O_2 concentrations and high 99 production efficiency. The choice of the electrochemical 100 reactor operating mode to be flow-by is since the use of 101 GDE in flow-by reactors has advantages in relation to the flow- 102 through reactors, as it minimizes the formation of bubbles on 103 the electrode surface, which increases the ohmic drop and also 104 reduces the possibility of the electrolyte salt precipitation 105 inside the GDE, blocking its channel structure and deactivating 106 it over time. 25

2. EXPERIMENTAL SECTION

- **2.1. Chemicals.** The following reagents were used to 108 perform the experiments: sodium sulfate (PanReac Appli- 109 Chem), sulfuric acid (Scharlab), 60% w/w poly- 110 (tetrafluoroethylene) dispersion (PTFE—Uniflon), and 111 titanium(IV) oxysulfate solution (Sigma-Aldrich). The aque- 112 ous solutions were prepared using ultrapure water (Milli-Q 113 system with resistivity > 18 M Ω cm). Printex L6 carbon 114 (PL6C) was purchased from Evonik Ltd. (Brazil).
- **2.2. Electrochemical Reactor Setup.** As can be seen in 116 Figure 1, the experimental system was set up using a flow-by 117 f1 electrochemical reactor with Printex L6 carbon/PTFE 118 deposited on carbon cloth employed as the gas diffusion 119 cathode and a dimensionally stable anode-chlor alkali (DSA- 120 Cl₂) used as the anode. The interelectrode gap was 8.0 mm, 121 and both electrodes occupied a geometric area of 20.0 cm². 122 The electrolyte solution, 0.1 mol L⁻¹ Na₂SO₄, pH 2.5, was fed 123 to the electrochemical cell from the reservoir tank (with a 124 capacity of 1.0 L) through a peristaltic pump operating at a 125

126 flow rate of 50.0 L h⁻¹ (under an electrolyte flow of 50 L h⁻¹, 127 the flow regime in the reactor is laminar, with a Reynolds 128 number of $\sim\!600$ and an internal rate (ν^0) of $\sim\!0.190$ m s⁻¹). 18 129 A thermostatic bath coupled to the reservoir tank was used to 130 control the temperature.

The flow-by electrochemical reactor used in this work operates under atmospheric pressure because the reactor is not hermetically closed due to the continuous entry of gas into the cathode compartment. O_2 gas was continuously injected into the cathode compartment, and this was monitored with the aid of a gas flowmeter. An Autolab PGSTAT302N potentiostat coupled with a BOOSTER 10A was used as a power supply. A 138 Pt//Ag/AgCl 3.0 M was employed as a pseudo-reference electrode and was coupled to the electrochemical cell, as the described by Beati et al. The platinum wire and the Ag/AgCl reference electrode were added to an external chamber containing the same electrolyte as the electrolyte from the electrochemical reactor (0.1 mol L^{-1} of Na_2SO_4).

2.3. Preparation of Gas Diffusion Electrode. The 145 carbon black (CB) material used for the conduct of the 146 experiments was Printex L6 carbon (PL6C)—acquired from 147 Degussa Brazil. Before manufacturing the GDE, the catalytic 148 material—Printex L6 carbon is heat-treated at 120 °C for 24 h 149 to remove water residues and organic interferences. 15 After 150 that, the carbon black material was mixed with 20 or 40% (w/ 151 w) of PTFE dispersion in 400 mL ultrapure water for 2 h until 152 the mixture was completely homogenized. The catalytic mass 153 was then filtered to remove excess water. Ten grams of the wet 154 catalytic mass was deposited and spread over the carbon cloth 155 (geometric area of 126 cm²). The electrode was dried at 120 156 °C for 15 min and was subsequently treated through the 157 application of a pressure of 5 tons and a temperature of 290 °C 158 for 2 h. The electrode was then cut into a circular shape of 20 159 cm².

2.4. Electrochemical Study. The following experimental parameters were investigated in this study: (i) temperature (25, 15, and 5 °C); (ii) O_2 flow applied to the cathode (10, 25, 50, 100, 200, and 300 mL min⁻¹ or 0.5, 1.25, 2.5, 5, 10, and 15 cm min⁻¹, respectively); (iii) PTFE (%) loading in the cathode composition (20 and 40%); and current density.

166 For the experimental tests (i) and (ii), electrolysis was 167 performed by applying a constant current density of 50 mA 168 cm $^{-2}$ for a period of 60 and 120 min, respectively. The PTFE 169 (%) loading was evaluated by electrolysis with different 170 currents applied (25, 50, 75, 100, 125, 150, 175, and 200 171 mA cm $^{-2}$).

2.5. Quantification of H_2O_2 . Hydrogen peroxide was quantified (in mg L^{-1}) using titanium(IV) oxysulfate solution as an indicator reagent, and the quantification analysis was performed by UV–vis spectroscopy (at $\lambda=408$ nm) using an Agilent 300 Cary series UV–vis spectrophotometer. The method adopted for the quantification of H_2O_2 in this study was based on the technique proposed in previous studies reported in the literature. 1,6,19

2.6. Service Lifetime Test. The lifetime of the electrode 181 was evaluated by applying a current density of 200 mA cm⁻² 182 using Arbin Instruments (model FBTS—20 V). Cyclic 183 voltammetry analysis was performed before and after the 184 electrochemical durability tests in a potential window of 0.0 to 185—0.8 V and at a scan rate of 50 mV s⁻¹ using Autolab 186 PGSTAT302N equipment and Ag/AgCl 3M as a reference 187 electrode.

2.7. Scanning Electron Microscopy (SEM) Images. The 188 Printex L6 carbon-based GDE was morphologically charac- 189 terized by scanning electron microscopy (SEM) using the 190 HRSEM-Gemini-500 equipment. The images were taken with 191 40× magnification.

3. RESULTS AND DISCUSSION

3.1. Effect of Temperature. One of the most efficient 193 electrochemical experimental setups employed in the produc- 194 tion of H₂O₂ via ORR through the 2-electron pathway involves 195 the coupling of electrodes in a flow-through reactor operating 196 at high pressure. The production of hydrogen peroxide is 197 limited by the availability of oxygen on the cathode surface, 198 and the solubility of this gas can be increased significantly 199 when the system is operated under high pressure and at low 200 temperature, as has been previously demonstrated. In 201 addition, the low temperature applied in the process can 202 help decrease the rate of H₂O₂ decomposition. By performing 203 the electrolysis under these optimal conditions using a flow-by 204 reactor in a discontinuously operated bench-scale plant, it was 205 possible to obtain a maximum accumulated H₂O₂ concen- 206 tration of approximately 400 mg L^{-1} at 0.9 Ah L^{-1} with a 207 temperature of 11.5 °C and a pressure of 2 bar and a specific 208 production of H₂O₂ of about 110 g kWh⁻¹. The production of 209 higher H₂O₂ concentration was not feasible in this 210 discontinuous process because, under these conditions, there 211 is an equilibrium between the rates of H₂O₂ production and 212 decomposition, and from this point onwards, the process 213 becomes unproductive. Thus, the only way to obtain higher 214 H₂O₂ production efficiency is to change the operation mode 215 from discontinuous to continuous mode, where the hydrogen 216 peroxide removed is protected against self-decomposition.

It should be noted that the physical mechanisms associated 218 with the reduction of oxygen in GDE are very different from 219 those that occur in electrodes made up of flow-through cells; 220 furthermore, it is interesting to evaluate whether the decrease 221 in temperature also exerts an influence over the physical 222 mechanisms related to oxygen reduction in GDE. Thus, to 223 evaluate the effects of temperature and 02 solubility in flow-by 224 reactors using GDE, electrolysis experiments were carried out 225 at 5, 15, and 25 °C, with a maximum 02 solubility of $^{14.8}$ – 226 11.2, 10.4, and 8.69 mg 14 , respectively. As can be seen in 227 Figure 2A, unlike what is observed under the application of 228 129 gas solubility do not play an influential role in the efficiency of 230 GDE when applied in a flow-by electrochemical cell.

The electrolysis experiments carried out at 5, 15, and 25 $^{\circ}$ C, 232 under atmospheric pressure and O_2 flow at 50 mL min⁻¹, 233 yielded very similar concentrations of H_2O_2 , with a mean value 234 of 543 mg L^{-1} and a standard deviation of 5.6 mg L^{-1} at 1 Ah 235 L^{-1} . The average concentrations of H_2O_2 electrogenerated 236 under the three temperature levels amounted a specific 237 production of H_2O_2 of 93.5 g kWh⁻¹—this is slightly lower 238 than the values obtained for flat electrodes equipped in flow- 239 through cells—the values reported for these electrodes ranged 240 between 101 and 135 g kWh⁻¹. 1,19

Furthermore, as can be seen in eqs 2 and 3, the 242 decomposition of H_2O_2 in the bulk solution or on the anode 243 surface did not cause a decline in H_2O_2 concentration; this 244 behavior was observed by Monteiro et. al in a flow-through 245 cell. The main advantage of the flow-through electrochemical 246 reactor is that the solution flow passes through the anode and 247 cathode, which improves the oxidation or reduction rate, as 248

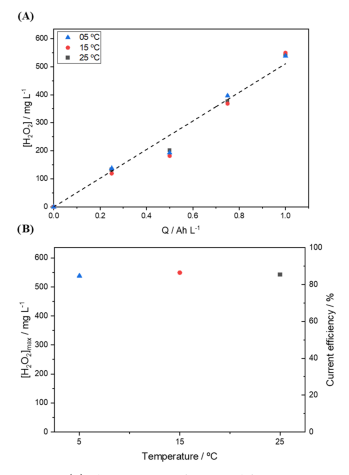


Figure 2. (A) Electrogeneration of H_2O_2 at different temperature levels (5.0, 15.0, and 25.0 °C); (B) maximum concentration of H_2O_2 produced in 60 min and current efficiency relative to the applied temperature for Printex L6 carbon employed at a current density of 50 mA cm⁻² using 0.1 mol L^{-1} Na₂SO₄, at pH 2.5, as a supporting electrolyte. O₂ flow rate employed: 2.5 cm min⁻¹.

249 well as the efficiency of the electrochemical process, because it 250 improves the convection of material transfer in the electrode 251 surface. According to Lu and Zhang, 27 flow-through reactors 252 have two advantages over the flow-by electrochemical reactors, 253 which are (i) high mass transfer efficiency and (ii) electron-254 transfer efficiency. In the flow-by reactor, the mass transfer is 255 highly limited by the fact that the electrodes are in parallel with 256 the solution flow. Thus, the rate of decomposition of H₂O₂ is 257 more pronounced in flow-through cells than in flow-by electrochemical cells; as such, the application of flow-through 259 cells does not allow one to obtain higher electrogenerated 260 H₂O₂ concentrations. It is worth emphasizing that the GDE 261 works by increasing the efficiency of O2 mass transfer at the 262 cathode, which improves this limitation of mass transfer in the 263 flow-by electrochemical cell; however, the DSA-Cl₂ anode, 264 which is parallel to the GDE, is limited to the mass transfer

$$H_2O_2 \to H_2O + \frac{1}{2}O_2$$
 (2)

$$_{266}$$
 $H_2O_2 \rightarrow O_2 + 2H^+ + e^-$ (3)

 267 It is noteworthy that the ORR process involving the 268 production of H_2O_2 mostly occurs in the triple phase of the 269 GDE since the O_2 that is solubilized in the electrolyte does not 270 practically interact with the electrode surface, and as such, it

does not take part in the process. This phenomenon, observed 271 in flow-through cells and which is characterized by the 272 occurrence of higher H_2O_2 production efficiency at low 273 temperature, would occur if H_2O_2 production was higher at 274 5 °C, once the solubility of O_2 is higher at this temperature 275 than at other temperature levels.

3.2. Effect of O₂ Flow Rate. As previously stated, the main 277 limitation of the electrochemical production of H₂O₂ is the ²⁷⁸ solubility of O2. High-pressured devices or accessories that 279 promote the drag of bubbles such as the venturi mixer 280 (improving the gas-liquid contact surface area) have been 281 shown to enhance the efficiency of the electrochemical process 282 by effectively supplying the O₂ needed as a raw material.^{6,19} 283 GDE is a suitable alternative device that helps to minimize the 284 mass transfer constraint in the reactor, but the flow rate of O2 285 gas that passes through the cell may influence the performance 286 of the electrochemical system in very different ways. 18,22 In 287 view of that, one needs to evaluate the O2 gas input so that 288 there is no shortage or excess of the reagent, as this will impact 289 the efficiency of H_2O_2 production. In the present study, the O_2 290 gas injection flow into the cathode compartment was evaluated 291 by varying the O₂ reagent input from 0.5 to 15 cm min⁻¹. The 292 experimental conditions were kept at 15 °C. As can be seen in 293 Figure 3, one can clearly observe that an increase in the gas 294 f3

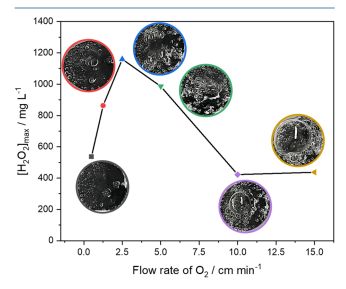


Figure 3. Maximum concentration of H_2O_2 produced in 120 min relative to the O_2 flow rate. The experiments were performed at a current density of 50 mA cm⁻² using 0.1 mol L^{-1} Na₂SO₄, at pH 2.5, as a supporting electrolyte.

flow resulted in an increase in the maximum concentration of 295 H_2O_2 accumulated in the electrochemical device up to 1,159 \pm 296 $13.6~{\rm mg~L^{-1}}$ at 2.5 cm ${\rm min^{-1}}.$ With regard to the O_2 flow 297 between 0.5 and 1.25 cm ${\rm min^{-1}},$ the amount of O_2 recorded 298 was lower; in other words, the oxygen did not interact with all 299 of the ORR active sites available on the GDE and because of 300 that the efficiency of H_2O_2 production and the accumulated 301 concentration of H_2O_2 produced were found to be lower in the 302 discontinuous process (536.4 and 863.1 mg L^{-1} for 0.5 and 303 1.25 cm ${\rm min^{-1}},$ respectively).

Looking at the images in the inset of Figure 3, one will 305 observe the formation of large air pockets at conditions above 306 5 cm min⁻¹; this is attributed to the excess gas that entered the 307 cathode compartment, which increased in diameter as the 308

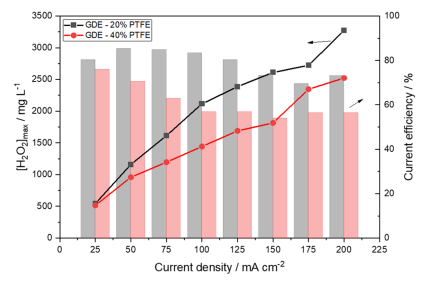


Figure 4. Maximum concentration of H_2O_2 obtained in 90 min of electrolysis based on the application of different current densities for each GDE. The supporting electrolyte employed: 0.1 mol L^{-1} Na₂SO₄, at pH 2.5.

309 injected O_2 flow increased. At 15 cm min⁻¹, the air pockets 310 covered a large part of the electrode surface, reducing the 311 three-phase contact area of the GDE; this resulted in a 312 decrease in the amount of H_2O_2 accumulated by 510 mg L^{-1} . 313 This negative effect represented a 2.7-fold decrease in H_2O_2 314 production efficiency.

The nondependence of the GDE on O_2 solubility and on its 316 3D-multichannel structure allows the electrode to provide an 317 unlimited supply of oxygen at the electrode/electrolyte 318 interface. The O_2 that passes through the GDE can directly 319 interact with the ORR active sites present in the Printex L6 320 carbon, and this promotes the generation of high concen-321 trations of H_2O_2 in the system. In addition, the versatility of 322 operation and easy installation of the GDE allow it to be 323 operated in bench cells and in full-scale electrochemical 324 reactors with volumes ranging from milliliters to hundreds of 325 liters.

A point worth mentioning is that the O2 gas humidity does 327 not play an important role in the ORR when using the GDE as cathode. The catalytic layer of the GDE is almost totally 329 hydrated due to the partial permeation of the electrolyte; in 330 this way, when the O2 gas flow is passing through the channels of the GDE, the gas tends to increase its degree of humidity.²⁸ Thus, regardless of the gas being prehumidified (relative humidity-RH of 50%) or dry (RH < 10%), a selfhumidification occurs as the gas passes through multichannels of the GDE, as well as a decrease in proton resistance decreases due to water enrichment along the channel.²⁸ Also, it is worth emphasizing that the oxygen reduction reaction must occur in an aqueous environment so that O_2 is reduced to H_2O_2 . Thus, is essential that the catalytic layer of GDE is partially wet, and this parameter can be controlled by the PTFE content. Xia et al. used gas diffusion electrodes based on carbon

Xia et al. used gas diffusion electrodes based on carbon anotubes with 40% PTFE loading to produce H_2O_2 in an undivided electrochemical cell with a volume of 0.16 L. The attained authors evaluated the effect of the O_2 flow injected directly into the GDE—this was one of the parameters investigated in their study. Based on their results, the increase in O_2 flow promoted an increase in O_2 mass transfer within the GDE, and this in turn led to an increase in H_2O_2 production. However, when the O_2 flow rate reached 280 mL min⁻¹, there was a

decline in the accumulated H_2O_2 concentration (compared to 350 the O_2 flow rate of 210 mL min⁻¹). The excess of O_2 flow led 351 to the formation of bubbles that covered the surface of the 352 electrode, and this led to a decrease in the production of H_2O_2 . 353 A similar outcome was noted in our present study. Xia et al. 354 obtained the best H_2O_2 production efficiency at a flow rate of 355 210 mL min⁻¹, where the accumulated H_2O_2 concentration 356 was 1291 mg L^{-1} (with a current efficiency of 88.5% in 60 min 357 of electrolysis). Remarkably, under the operating conditions 358 employed by Xia et al., the amount of O_2 consumed was 359 twice as high as the amount of O_2 consumed in the present 360 work even though our proposed system operated for an 361 additional 1 h (1159 \pm 13.6 mg L^{-1} in 120 min).

Another study that deserves being mentioned is that of Lima 363 et al. where the authors employed a Printex L6 carbon-based 364 GDE (similar to the electrocatalyst employed in our present 365 study) to evaluate H₂O₂ production in an electrochemical cell. 366 For comparison purposes, Lima et al. employed different 367 amounts of carbon (8 g) and PTFE loading (40%) in their 368 analysis (in the present study, 0.67 g carbon loading and 20% 369 PTFE loading were employed).²⁴ With the GDE exhibiting a 370 relatively larger thickness due to the higher amount of carbon 371 in its composition, the authors had to apply a pressure of 0.2 372 bar of O₂ gas in the cathode compartment for the electrode to 373 work in the best condition.²⁴ Under these conditions, Lima et 374 al. reported having obtained an accumulated H₂O₂ concen- 375 tration of \sim 750 mg L^{-1} after 120 min of electrolysis in an $_{376}$ electrochemical cell. Interestingly, despite consuming a higher 377 amount of reagent (O2), the amount of H2O2 concentration 378 obtained in their study²⁴ corresponds to only 65% of H₂O₂ 379 concentration obtained from the application of the Printex L6- 380 based GDE proposed in our present study. This shows that 381 high amounts of carbon or high PTFE loadings are not 382 required in the composition of the GDE since the ORR 383 process, involving H₂O₂ production, occurs slightly below the 384 electrode surface (on the catalytic layer of GDE), and thus the 385 use of thinner electrodes can lead to satisfactory results.

3.3. Effect of PTFE (%) Loading and Current Density. 387 The percentage content of PTFE employed in the GDE exerts 388 an influential role on the hydrophobicity of the electrode; in 389 other words, increasing the PTFE content in the GDE 390

391 composition makes the electrode more hydrophobic and this 392 inhibits the permeability of the aqueous solution through the 393 electrode. With regard to the electrode proposed in the present 394 study, the application of more than 40% PTFE in the GDE was 395 found to render the device excessively hydrophobic, and this 396 made the system behave like a conventional/flat electrode. On 397 the other hand, the application of lower contents of PTFE 398 resulted in higher permeability of the solution in the GDE 399 (higher degree of wettability). With low contents of PTFE, the 400 preliminary assays indicated that at values below 20%, the 401 solution completely permeates the electrode according to the 402 use of the GDE; this fact was observed by an electrolyte 403 soaking in the cathode compartment when a GDE containing 404 10% PTFE was used. Thus, it was not possible to generate 405 H₂O₂ electrosynthesis data for GDE containing values below 406 20%.

It should be noted, however, that there is a minimum PTFE loading value that will prevent the solution from soaking through the electrode. Thus, it is essentially important to find an ideal PTFE loading that helps to prevent high hydrothin phobicity or high wettability of the electrode. The results obtained from the thorough analysis conducted in the present study helped obtain some useful insights in this regard. Based on our findings, the ideal PTFE loading should be between 20 and 40%; this is because below 20% PTFE loading, flooding occurs on the electrode, while there is high resistance to solution permeability when one applies a PTFE loading above 418 40% (see Figure 4).

At horough investigation was carried out to evaluate $\rm H_2O_2$ generation at different current densities using the GDE with 421 PTFE loadings of 20 and 40% (see Figure 4 for the results 422 obtained). One will observe that the accumulated $\rm H_2O_2$ 423 concentrations (obtained in 120 min of electrolysis) for the 424 20% PTFE–GDE were between 1.1- and 1.4-fold higher than 425 those obtained for the 40% PTFE–GDE at all of the current 426 densities evaluated. The 20% PTFE–GDE contains the 427 equivalent of 80% of Printex L6 carbon by mass; this is 20% 428 more than the amount of PL6C in the 40% PTFE–GDE.

The difference in the carbon content between the two data electrodes (20% PTFE–GDE and 40% PTFE–GDE) resulted that in an increase of almost 46% in current efficiency for the production of $\rm H_2O_2$ at a current density of 100 mA cm⁻²; this data effect can be attributed to a higher amount of ORR active sites available for $\rm O_2$ adsorption on the carbon surface. It is also worth emphasizing that lower contents of PTFE contain a data higher amount of carbonaceous matrix (since the catalytic mass of GDE is composed of Printex L6 carbon and PTFE). The carbon matrix is responsible for promoting the electrosum characteristic production of $\rm H_2O_2$, and thus, the greater the content data of carbonaceous material, the greater the amounts of ORR data active site present in the GDE.

In previous works reported in the literature, 4 it shows that 443 Printex L6 carbon contains only carboxyl-type oxygenated 444 functional groups (COOH) in its chemical composition 445 (18.6% content—data referring to an analysis by X-ray 446 photoelectron spectroscopy, XPS). As reported in the 447 literature, 2,4 the carboxyl group is the oxygenated functional 448 group that has the greatest influence on H_2O_2 electrosynthesis, 449 followed by the carbonyl functional group (C=O). The 450 oxygenated functional group on the surface of the carbona-451 ceous material is responsible for the displacement of electrons 452 from its adjacent carbon, making it an excellent active site for 453 the adsorption of the O_2 molecule, and for tending to the

formation of the OOH* intermediate, which is the only 454 intermediate that favors the formation of H_2O_2 (the * 455 symbolizes that the species is adsorbed at the active site).

Looking at Figure 4, one will observe that an increase in the 457 current density resulted in an increase in the accumulated 458 $\rm H_2O_2$ concentrations obtained for both GDEs, but there was 459 no ideal current density to work with. The accumulated $\rm H_2O_2$ 460 concentrations for 20% PTFE–GDE were 1614.3 \pm 19.1, 461 2610.7 \pm 40.6, and 3271.0 \pm 47.3 mg L $^{-1}$ at 75, 150, and 200 462 mA cm $^{-2}$; for 40% PTFE–GDE were 1196.3 \pm 14.3, 1443.1 \pm 463 23.6, and 2523.9 \pm 38.5 mg L $^{-1}$ at 75, 150, and 200 mA cm $^{-2}$, 464 respectively.

Thus, for a better understanding of the electrochemical 466 production of H₂O₂, one needs to observe Figure 5, which 467 fs

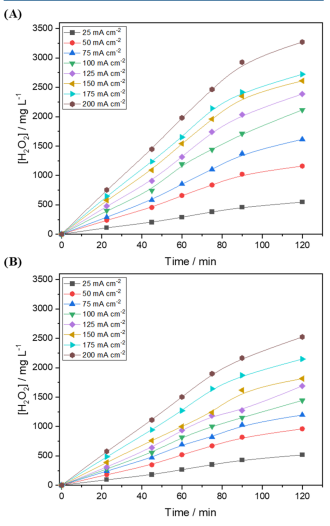


Figure 5. Amount of H_2O_2 electrogenerated (in mg L^{-1}) for (A) Printex L6 carbon with 20% PTFE loading and (B) Printex L6 carbon with 40% PTFE loading at different current densities using 0.1 mol L^{-1} Na₂SO₄, at pH 2.5, as a supporting electrolyte. O₂ flow rate: 2.5 cm min⁻¹; electrolyte temperature: 15 °C.

shows the relationship between the concentrations of H_2O_2 468 generated as a function of time for both GDEs. It can be noted 469 that the application of current densities higher than 150 mA 470 cm⁻² led to a decrease in H_2O_2 concentration after 90 min of 471 electrolysis due to the decomposition of H_2O_2 within the 472 solution and on the anode surface (as discussed in eqs 2 and 473 3). The consumption of H_2O_2 by these parallel reactions 474

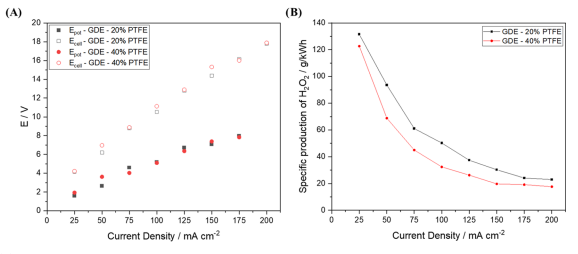


Figure 6. (A) Average cathode potential and cell potential values obtained for the electrodes investigated in 120 min of electrolysis at different current densities. (B) The specific production of H_2O_2 in g kWh⁻¹ vs applied current density.

475 causes a decline in the current efficiency; at very high current 476 densities, the current efficiency may also decrease by favoring 477 the ORR via 4-electron transfer on the cathode surface. The 478 20% PTFE–GDE exhibited a maximum current efficiency of 479 85.3% at a current density of 50 mA cm⁻². From this value 480 onwards (85.3%), the current efficiency only decreased until it 481 reached 73% at 200 mA cm⁻².

The same behavior (decline in current efficiency) was observed for the 40% PTFE—GDE; however, this electrode that recorded a maximum current efficiency of 75.9% at a current density of 25 mA cm $^{-2}$. Thus, one can conclude that for an operation aimed at obtaining a higher current efficiency, one needs to employ low current densities. However, when the aim that is to obtain high $\rm H_2O_2$ concentration in a short period of time, one will need to employ a high current density.

As can be seen in the first 60 min of the graph in Figure 5, 491 the 20% PTFE-GDE recorded kinetic constant values that 492 were 24, 47, 51, and 34% higher than those of the 40% PTFE-493 GDE at current densities of 50, 100, 150, and 200 mA cm⁻², 494 respectively. Both electrodes exhibited an apparent pseudo-495 order kinetic constant of zero (i.e., the generation of H₂O₂ is 496 independent of the concentration of O₂ and H⁺). The 20% 497 PTFE-GDE showed H₂O₂ production rate values of 10.7, 498 15.5, 19.9, and 28.5 mg L^{-1} min⁻¹ at current densities of 50, 499 100, 150, and 200 mA cm⁻², respectively. Valim et al.³⁰ 500 reported a H₂O₂ production rate of 5.9 mg L⁻¹ min⁻¹ when 501 operated at -1.0 V vs Ag/AgCl, whereas Carneiro et al. 15 502 reported a slightly higher rate of 7.6 mg L⁻¹ min⁻¹ at the same conditions. In both works, the GDE was modified with metallic oxides, whose modification is to improve the selectivity and 505 catalytic activity of the carbonaceous material. Moreira et al.³¹ 506 reported surprising values of 38 mg L⁻¹ min⁻¹ when operated at 100 mA cm⁻² using a Sudan-Red-modified Printex L6 carbon-based GDE. 508

Finally, an interesting element to consider in our analysis of the efficiency of the electrochemical process is the cell potential ($E_{\rm cell}$) and the difference of potential between the cathode and pseudo-reference electrode Pt//Ag/AgCl ($E_{\rm cat-ref}$) values; this is because these potentials may indicate changes in the reactor setup or even in the electrode fabrication method. With that in mind, an analysis was conducted to evaluate whether the amount of carbon in the composition of the GDE can affect the cathode potential and cell potential values since

it affects the conductivity of the electrode. Interestingly, both 518 the 20% PTFE–GDE and 40% PTFE–GDE recorded very 519 similar $E_{\rm cat-ref}$ and $E_{\rm cell}$ values, as seen in Figure 6A; this shows 520 f6 that both electrodes exhibit similar electrochemical behavior, 521 despite the difference in carbon content. 522

As expected, an increase in the current density resulted in an 523 increase in the potential values; also, higher $E_{\rm cell}$ values were 524 recorded at higher current densities. This outcome suggests 525 that, under these working conditions, the anode DSA-Cl₂ plays 526 an essential key role in the process, especially in the parallel 527 reactions that lead to the decomposition of H_2O_2 , as can be 528 observed in eq 3 or in the formation of predator species for 529 $H_2O_2^{-21}$ such as ozone, which plays a role in the decomposition 530 of H_2O_2 once it is formed and disappears immediately 531 afterward. It is worth noting that the use of more active 532 anodes, such as BDD anode, can increase the intensity of 533 parallel reactions and negatively influence the production of 534 H_2O_2 .

Based on the cell potential values, one can estimate the 536 specific production of H_2O_2 in g kWh⁻¹; this is an important 537 and more realistic parameter that can help evaluate the 538 applicability of the electrodes in real systems. The term specific 539 production of H_2O_2 represents how much hydrogen peroxide 540 is produced per power generated per time, which can be 541 expressed in grams per kW per hour (g kWh⁻¹). The specific 542 production of H_2O_2 was calculated using eq 4, where $C_{H_2O_2}$ is 543 the concentration of H_2O_2 (in mg L⁻¹), V is the volume (in L), 544 E is the cell potential (in V), I is the current (in A), and t is the 545 time (in h)

specific production of
$$H_2O_2\left(\frac{g}{kWh}\right) = \frac{C_{H_2O_2} \cdot V}{E \cdot I \cdot t}$$
 (4) ₅₄₇

Figure 6B shows the results obtained from the analysis of the 548 specific production of H_2O_2 as a function of current density. 549 One will observe that an increase in current density (j), which 550 consequently results in an increase in the cell potential value, 551 causes a decrease in the specific production of H_2O_2 ; in other 552 words, smaller grams of H_2O_2 are produced per kWh at high 553 current densities (the application of j > 100 mA cm⁻² results in 554 the specific production of H_2O_2 values less than 37 g kWh⁻¹). 555 On the other hand, at lower current density; i.e., 25 mA cm⁻², 556 an extremely high specific production of H_2O_2 values were 557

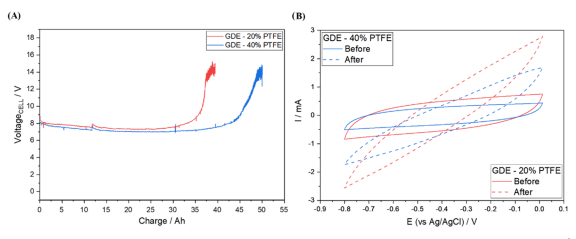


Figure 7. (A) Durability test for 20% PTFE–GDE (red line) and 40% PTFE–GDE (blue line) at a current density of 200 mA cm $^{-2}$; (B) cyclic voltammetry analysis performed on the cathode potential (in the potential range of 0.0 to -0.8 V and at a scan rate of 50 mV s $^{-1}$) before and after the durability test using O₂-saturated 0.1 mol L $^{-1}$ Na₂SO₄ (pH 2.5 adjusted with H₂SO₄) as an electrolyte solution.

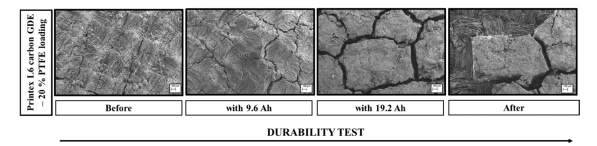


Figure 8. SEM images (with 40× magnification) obtained for 20% PTFE-GDE before and after the durability test.

 558 recorded—the 20% PTFE–GDE and 40% PTFE–GDE 559 recorded the specific production of $^{12}O_2$ of 131.5 and 122.6 560 g kWh⁻¹, respectively.

A comparison of the specific production of H₂O₂ values 562 obtained for the proposed 20% PTFE-GDE and 40% PTFE-GDE with the values obtained for other gas diffusion electrodes reported in the literature pointed to the superior 565 performance of the electrodes proposed in the present study. 566 For illustration purposes, Lima et al. 24 employed 40% PTFE-567 GDE at a current density of 50 mA cm⁻², where they obtained 568 a maximum H₂O₂ concentration of 750 mg L⁻¹, with a specific 569 production of H_2O_2 of ~23.5 g kWh⁻¹; this is roughly 2.9- and 570 4-fold smaller compared to the values obtained for the 40% 571 PTFE-GDE and 20% PTFE-GDE proposed in our present 572 work. Moreira et al.³¹ also employed 20% PTFE-GDE, where 573 they obtained the specific production of H₂O₂ of ~9.5 g 574 kWh⁻¹ at 100 mA cm⁻²; this is roughly 5 times lower than the 575 value obtained in this work. Barros et al. 16 employed an 576 unmodified GDE for the electrochemical generation of H₂O₂ 577 in a potentiostatic mode at -1.1 V (the cell current value 578 should be approximately 150 mA cm⁻²), where they obtained a 579 specific production of H₂O₂ of approximately 28 g kWh⁻¹ and 580 H₂O₂ accumulated a concentration of 6,400 mg L⁻¹; a 581 comparison of the conditions applied in their work with the 582 GDE developed in our present work showed that our proposed GDE exhibited a slightly higher H₂O₂ production efficiency of 584 30 g kWh⁻¹.

The efficiency in the production of H_2O_2 of the GDE seed developed in this work is because it is composed of a diffusion layer based on a carbon cloth and a catalytic layer based on seed PTFE. Some GDE reported in the

literature (e.g., the work by Lima et al., 24 Moreira et al., 31 and 589 Barros et al. 16) employ a single layer that acts as both a 590 diffusion and catalytic layer. In those case, up to the point at 591 which the solution permeates the GDE, it is called the catalytic 592 layer, while after this point, where the solution does not 593 permeate, it is called the diffusion layer. The use of carbon 594 cloth as the diffusion layer facilitates the diffusion of O2 gas to 595 the catalytic layer, and therefore, it was possible to achieve a 596 higher production of H2 O₂.

Flow-by electrochemical cells are characterized by higher cell 598 voltage compared to pressurized nondivided microfluidic 599 electrochemical cells. The main advantage of the pressurized 600 nondivided microfluidic cells lies in the short separation 601 distance between the cathode and the anode; it is this short 602 distance between the cathode and the anode that allows these 603 cells to have lower cell voltage and ohmic resistance compared 604 to flow-by cells. It is worth noting that the lower the cell 605 voltage, the less amount of specific production of H2O2. In 606 previous studies conducted by Moratalla¹⁹ and Monteiro, ¹ the 607 authors obtained the specific production of H₂O₂ values that 608 ranged between 101 and 135 g kWh⁻¹ at a current density of 5 609 mA cm⁻²; the values they obtained are slightly higher than 610 those obtained in our present study at 25 mA cm⁻². It should 611 be noted, however, that the aforementioned studies [1, 20] 612 employed different electrode technologies in the electro- 613 chemical process, which was more dependent on temperature 614 and pressure.

3.4. Durability Test for GDE. The electrochemical $_{616}$ resistance of the 20% PTFE-GDE and 40% PTFE-GDE $_{617}$ was evaluated by applying a current density of 200 mA cm $^{-2}$ $_{618}$ and the operation time needed for the $E_{\rm cell}$ to increase $_{619}$ f7

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620 exponentially. As can be seen in Figure 7A, the electrode 621 containing 20% of PTFE loading maintained the E_{cell} constant 622 for 36 Ah (this corresponds to 7.5 days), after which the 623 voltage increased significantly. The electrode with 40% PTFE 624 loading exhibited a longer lifetime, reaching an uninterrupted 625 life span of 48 Ah (10 days). After the aforementioned lifetime, 626 both GDEs exhibited a more resistive current profile, as can be 627 seen in the cyclic voltammograms obtained before and after 628 the durability test (see Figure 7B); this behavior can be 629 attributed to the fact that the electrodes lost a significant part 630 of the catalytic film (the catalytic mass containing Printex L6 631 carbon and PTFE) deposited under the carbon cloth. In 632 addition, the GDEs were found to have been soaked by the electrolyte.

Figure 8 shows the SEM images related to the removal of the 635 catalytic film from the carbon cloth in the 20% PTFE-GDE. 636 Before the durability test, the catalytic mass was deposited 637 homogeneously and uniformly over the carbon cloth substrate (Figure 8, before). During the durability test, the catalytic film (Printex L6 carbon + PTFE) started to exhibit some cracks 640 (like cracked soil). An increase was observed in the thickness 641 of the cracks over time until parts of the catalytic film were 642 removed from the carbon cloth substrate (Figure 8, after).

4. CONCLUSIONS

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643 The present work reported the development and application of 644 a new optimized carbon-based gas diffusion electrode 645 supported on carbon cloth employed as a cathode in a flow-646 by electrochemical reactor for the electrochemical production 647 of H₂O₂. The following conclusions can be drawn from the 648 results obtained in this study:

- The use of the proposed PTFE-GDE (as cathode) in a flow-by reactor helped eliminate the dependence of the electrochemical process on temperature and O₂ solubility in terms of the electrochemical production of H_2O_2 ; the application of the proposed system allowed O₂ to pass through the 3D-multichannel structure of the GDE until it reached the triple phase where the ORR active sites were located, giving rise to the production of H₂O₂. The application of the following temperature levels, 5, 15, and 25 °C, resulted in the average H₂O₂ production of 8.51 mg L⁻¹ min⁻¹ and a specific production of H_2O_2 of 93.5 g kWh⁻¹.
- An analysis of the mass transfer of O₂ injected into the cathode compartment showed that the ideal O2 flow rate was 2.5 cm min⁻¹; the application of this O₂ flow rate in the electrochemical system led to the production of $1,159 \pm 13.6 \text{ mg L}^{-1} \text{ of H}_2\text{O}_2 \text{ in } 120 \text{ min of electrolysis.}$ The application of an O₂ flow rate below 2.5 cm min⁻¹ caused a significant decrease in mass transfer, and this undermined the interaction of O2 with the active sites of the ORR; in these conditions, large air pockets were formed, which covered a great part of the electrode surface, reducing the H₂O₂ production efficiency.
- The hydrophobic characteristic of the GDE is determined by the PTFE loading. The findings of this study showed that the application of GDE composed of PTFE loading below 20% led to the soaking of the electrode, while the incorporation of PTFE loading above 40% into the GDE elevated the resistance of the electrode to partial permeability, impeding the smooth operation of the triple phase. The 20% PTFE-GDE

- sample produced a high accumulated H₂O₂ concen- 680 tration, which was 1.4 times higher than the amount 681 obtained for the 40% PTFE-GDE.
- The application of low current densities favored the 683 current efficiency and H2O2 production efficiency; the 684 application of the current density of 50 mA cm⁻² 685 resulted in current efficiency and specific production of 686 H₂O₂ of 80.3% and 131 g kWh⁻¹, respectively. It should 687 be noted, however, that the application of a current 688 density of 200 mA cm⁻² in 120 min of electrolysis led to 689 the production of an accumulated H2O2 concentration 690 of nearly 3,270 \pm 47.3 mg L⁻¹ but with a low current 691 efficiency of 64.5% and a specific production of H2O2 of 692 23 g kWh⁻¹. This is attributed to the increase in parallel 693 reaction rates associated with H₂O₂ decomposition as a 694 result of an increase in applied current density.
- The durability/lifetime test conducted showed that the 696 40% PTFE-GDE recorded a lifetime of 48 Ah (which 697 corresponds to 10 days of uninterrupted use) at 200 mA 698 cm⁻². The lifetime of the 40% PTFE-GDE was found 699 to be 1.3 times higher than that of the 20% PTFE- 700 GDE; in essence, this result shows that an increase of the 701 PTFE loading in the GDE resulted in an increase in the 702 electrode lifetime. Over time, the catalytic film showed 703 surface cracks, which increased in thickness until the film 704 was removed from the substrate.

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Notes

The authors declare no competing financial interest.

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