

1 **ON THE EFFECTS OF FERRICYANIDE AS CATHODIC MEDIATOR ON THE**
2 **PERFORMANCE OF MICROBIAL FUEL CELLS**

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18

19 **ABSTRACT**

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21 This study provides an insight into the long term influence of the use of ferricyanide in
22 the cathode chamber of a MFC on the power generated and the COD removal attained.
23 Two MFCs were operated in semicontinuous mode, using winery wastewater as fuel,
24 activated sludge as the anodic inoculum and concentrations of 0.05M and 0.25M of
25 ferrocyanide added in the cathode chamber as redox mediators. The MFC used had two
26 chambers separated by a proton exchange membrane Sterion®. The results shows that
27 permeability of the membrane to mediators is a factor of the major significance. Under
28 no crossover, the mediator produced a positive effect on the electricity generation and
29 COD removal. However, as the experiments progressed a significant concentration of
30 mediator was detected in the anode chamber and the performance of the MFC gets worse.
31 This work reports results that helps to understand the main processes happening in the
32 MFC.

33

34 **HIGHLIGHTS**

- 35
- 36 • Crossover of ferrocyanide and ferricyanide to the anodic chamber reduces
37 efficiency in the production of electricity
 - 38 • The couple ferro/ferricyanide in the cathode chamber of a MFC can improve the
39 performance in terms of COD removal and energy efficiency
 - 40 • At low mediator concentration, a maximum in the power generation and COD
41 removal can be attained
 - 42 • At high mediator concentration, there was inhibition of biodegradation of winery
43 wastewater
- 44

45 **Keywords:** wastewater treatment; energy recovery; winery wastewater; microbial fuel
46 cell; ferrocyanide

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50 **INTRODUCTION**

51

52 Nowadays, there is a tremendous need to develop cost-effective and less energy intensive
53 technologies for the treatment of wastewater. In this context, novel systems for the
54 simultaneous recovery of energy and treatment of wastewater have gained interested and
55 among them, microbial fuel cells (MFC) seem to be promising and worth of study, in
56 spite of being still far away from practical applications. MFC consist of an anode, where
57 the microorganisms oxidize the organic matter of wastewater to generate electrons and
58 protons, and a cathode where the electrons and protons reduce an electron acceptor . The
59 protons in anode chamber can travel through the membrane or through the liquid, while
60 the electrons pass by an external circuit. Both react with an electron acceptor in cathode
61 chamber resulting in power generation [1, 2].

62 Many previous studies have demonstrated that MFC is greatly influenced by many
63 factors, such as the microorganism, the material of electrodes, the operational parameters
64 [1, 2]. Among these factors, the cathodic electron acceptor has been found to be crucial
65 to MFC performance. Most studies used oxygen as the electron acceptor in MFC, because
66 of its high oxidation potential, availability; economically attractive and environmentally
67 friendly because water is formed as the only end product. However, the performance of
68 MFC is low using oxygen due to the slow kinetics of oxygen reduction (Ter Heijne et al.,
69 2006;).

70 Recently, a number of studies have shown that ferrocyanide/ferricyanide is excellent
71 cathodic electron acceptor, due to its good performance in the MFC. Several researches
72 obtained very high power densities using ferricyanide in the cathodic chamber of MFC
73 such as 4310 mWm^{-2} and 3600 mWm^{-2} [3, 4] while the power generation is typically less
74 than 1000 mWm^{-2} using oxygen dissolved [5]. The rapid oxidation of ferrocyanide by

75 oxygen to form ferricyanide and the reversibility of this reaction, that favor the easier
76 cathodic reduction of the ferricyanide (as compared to the oxygen reduction) are the main
77 characteristic looked for in this mediator [6]. However, toxicity for microorganisms of
78 the couple ferrocyanide/ferricyanide is well-known and it becomes a major challenge in
79 MFC, making necessary a good isolation of the anodic chamber to prevent the inhibition
80 of the biological processes, in particular those carried out by bioelectrogenic
81 microorganisms.

82 Obviously, the best way to prevent the transport of ferrocyanide and ferricyanide to the
83 anodic chamber is a proton exchange membrane, in particular taking into account that
84 both species are anions. This is not a perfect solution because of the permeability of the
85 membrane, which may prevent the perfect isolation of the anodic chamber with respect
86 to the cathode mediator.

87 Winery wastewater is a promising feedstock for MFC technology [7, 8] due its rich
88 composition in sugars, vitamins, minerals, and redox-active mediators such as tannic acid,
89 and in a recent study, a power density as high as 465 mW m^{-2} was obtained with a two
90 compartment MFC directly fed with this type of wastes [9]. The aim of this work is to
91 assess how the use of ferrocyanide can help to improve the performance of this type of
92 MFC by enhancing the efficiency of the cathodic reduction in such devices. Therefore, in
93 the present study, potassium ferricyanide is studied as the electron acceptor in the cathode
94 chamber of a two-chamber MFC, in order to evaluate the removal of winery wastewater
95 in the power generation. A Sterion[®] proton exchange membrane has been used to separate
96 the compartments of the cell. This membrane shows low permeability as compared to
97 other commercial membranes[10]. However, as the permeability of membranes is always
98 a factor to be considered, concentration of the mediator couple ferrocyanide/ferricyanide
99 was decided to be monitored in the anodic chamber. Likewise, it was decided in the design

100 of the tests not to modify it during the test, in order to have a clear view of the performance
101 of this type of MFC, pointing out their advantages and disadvantages. On the other hand,
102 taking into account the typical low reproducibility of the MFC devices (clearly justified
103 in terms of the great deal of parameters that influence on their performance) it was decided
104 to run simultaneously (and under the same operation conditions except for the
105 ferro/ferricyanide concentration) two MFC during approximately 40 days, and then to
106 reverse the ferro/ferricyanide concentration in their respective cathode chambers,
107 without changing the electrodes or other components of the electrochemical cell

108

109 MATERIAL AND METHODS

110

111 **MFC configurations and operation.** Two MFC (so-called MFC₁ and MFC₂) were used
112 in this study. The MFCs were made of acrylic tubes (inner diameter 40 mm; length 180
113 mm) and divided by a Sterion[®] membrane into two chambers, with 70 mL (anode) and
114 100 mL (cathode), respectively. The Sterion[®] membrane was pretreated with 3% (v:v)
115 hydrogen peroxide solution, 0.5 mol L⁻¹ sulfuric acid and ultrapure water. Carbon felt
116 (KFA10, SGL Carbon Group[®]) were used as electrode in both chambers without being
117 copped with Platinum. A stainless steel wire and an external resistance of 120 Ω
118 connected the anode and the cathode.

119 The MFCs were operated in parallel in semi-continuous mode and at room temperature.
120 Every day, 50 mL of liquid was removed from anode chamber and replaced by fresh
121 winery wastewater. A peristaltic pump was used to recirculate catholyte from the
122 reservoir (500 mL) through the cathode chamber of the MFC at 1.66 mL s⁻¹. Oxygen was
123 supplied in cathode reservoir using an aquarium aerator and porous stone diffuser in all
124 experiment. When ferricyanide is used in the catholyte of MFC, the ferricyanide is

125 reduced to ferrocyanide. Then, the ferricyanide can be recovered by the oxygen supplied
126 in the reservoir, oxidizing the ferrocyanide formed in the cathode.

127 Firstly, the MFCs were started-up using HCl (pH 3.0) in the cathode compartment for 6
128 days. Then, to evaluate the influence of ferricyanide concentration (in the cathode) on
129 MFC performance, two concentrations were chosen: 0.05 and 0.25 M. The MFC₁ was
130 started up with 0.05 M of ferricyanide in the cathode and was operated for 46 days. Then,
131 the MFC₁ was run using the solution of 0.25M of ferricyanide between the 47th and 80th
132 day of operation. The MFC₂ was operated just in the reverse conditions of MFC₁, firstly
133 with 0.25 M and then with 0.05 M of ferricyanide, in order to evaluate the potential
134 hysteresis of the system.

135

136 **Inoculum and wastewater.** The inoculum was collected from the activated sludge
137 reactor at the municipal Wastewater Treatment Plant of Ciudad Real (Spain) and
138 concentrated by sedimentation. The concentration of total solids and total volatile solids
139 were 15.8 and 11.1 g L⁻¹ respectively. To inoculate the MFCs, a solution of 90% of seed
140 sludge and 10% winery wastewater was added into the anode chamber and kept for 24
141 hours before start the operation. The winery wastewater used in this work was taken from
142 the collecting tank of the Wastewater Treatment Facility of the winery Bodegas Crisve
143 (Socuellamos, Spain), and stored at 4°C before being used. Table 1 shows the
144 composition of this winery wastewater. NaHCO₃ (6000 mg L⁻¹) were used to adjust and
145 to buffer the pH to 6.5. Dibasic sodium phosphate (Na₂HPO₄·2H₂O) and ammonium
146 sulfate ((NH₄)₂SO₄) were added to increase the phosphorous and nitrogen concentrations
147 to 10 mg P-PO₄³⁻ L⁻¹ and 100 mg N-NT L⁻¹, because in a previous study it was determined
148 that the unbalanced COD:N:P ratio is one of the biggest problem in treating this type of
149 wastewater in biological system [9].

150

151

Table 1: Characteristics of winery wastewater used.

Parameter	Value
pH	4.11
Conductivity(mS cm ⁻²)	2030
COD (mg L ⁻¹)	6850
TOC (mg L ⁻¹)	1030
Total Nitrogen(mg L ⁻¹)	18.3
Total Phosphorous(mg L ⁻¹)	0.95

152

153

154 **Analytical methods.** A GLP22 Crison[®] pH-meter, a GLP 31 Crison[®] conductivity meter
155 and an Oxi538 WTW[®] oxy-meter were used to measure pH, conductivity and dissolved
156 oxygen, respectively. The total suspended (TSS) and volatile suspended (VSS) solids
157 were measured gravimetrically [11]. The COD and concentration of phosphorous were
158 measured using a spectrophotometric method (DR2000, HACH[®]). The total nitrogen was
159 monitored using a Multi N/C 3100 Analytik Jena analyzer. The ferrocyanide
160 concentrations were analyzed by spectrophotometer method (KODAK, 1999).

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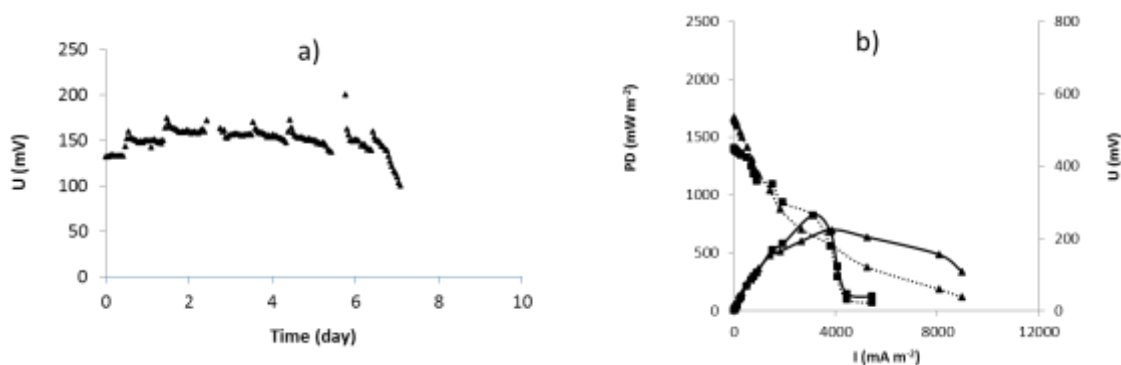
162 **Electrochemical measurements.** A digital multimeter (Keithley[®] 2000) was connected
163 to the system to continuously monitor the value of the cell potential and the data were
164 recorded in a personal computer. The polarization curves from the MFC were obtained
165 by varying the resistance in the circuit and measuring the voltage. Power density (mW m⁻²)
166 and current density (mA m⁻²) were based on the surface area of anode (7.0 cm⁻²). The
167 current (I) was calculated using Ohm's Law ($I = E/R$), and the output power of the cell
168 using $P = I E$, where I (A) is the current, E (V) is the voltage, R (Ω) is the external

169 resistance and P (W) is the power. Coulombic efficiency (CE) was based on total current
170 generation and the maximum current that can be produced from COD oxidation and was
171 calculated according to a procedure described in a previous work [11].

172

173 RESULTS AND DISCUSSION

174 **Electricity production.** Figure 1 shows the changes in the cell potential (resistance of
175 120 ohm) and the polarization curves of a mediatorless double chamber microbial fuel
176 cell operated in the same conditions than the two MFC tested in this work and used as a
177 reference.



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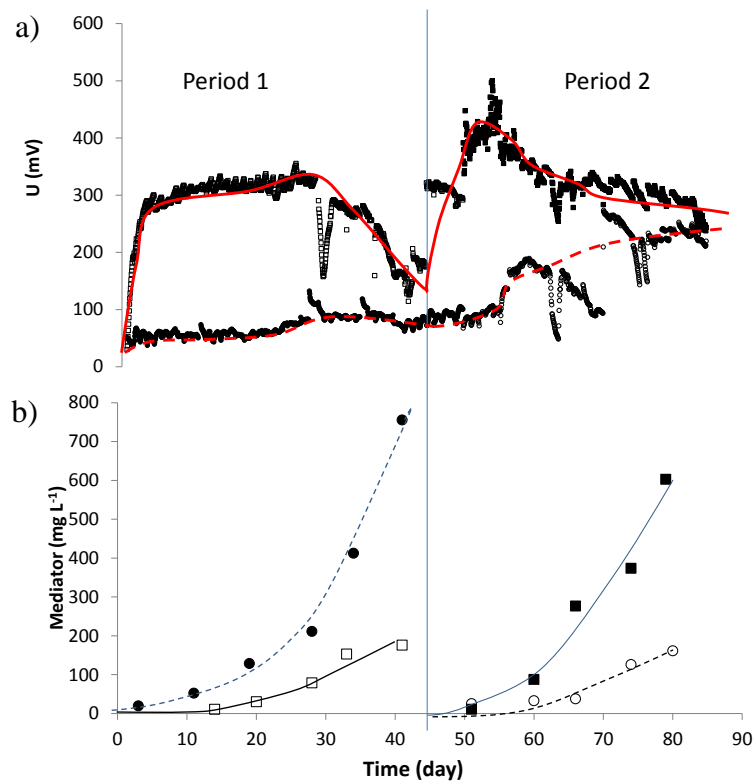
179 **Figure 1.** Electricity produced in MFC_0 treating winery wastewater without ferricyanide mediator in
180 cathode chamber (\blacktriangle). B) polarization curves obtained at days 4th and 6th.

181

182 As it can be observed, the system rapidly met a steady state cell voltage of 150 mV and
183 the maximum power density was around 750 mW m^{-2} . These values were going to be used
184 in this work as reference for comparison purposes because the MFC_0 was operated under
185 the same conditions of the two MFC with mediators. At this point, Figure 2 shows the
186 changes in the voltage (resistance of 120 ohm) and the concentration of
187 ferrocyanide/ferricyanide measured at the anodic chambers during the four tests carried
188 out. During the first period (which lasted 46 days) the cathodic chamber of MFC_1
189 contained 0.05 M of mediator (ferrocyanide/ferricyanide) while the cathodic chamber of

190 the MFC₂ five times more (0.25M). From day 46th on, the concentrations of mediators in
191 both cells were changed in order to evaluate the differences and the robustness of the
192 microbiological processes.

193 It can be clearly observed that low concentration of mediator in the cathode have a very
194 positive effect on the production of electricity. Thus, when a MFC was operated with acid
195 chloride solution (pH of 3.0) in the cathode, without ferrocyanide/ferricyanide redox
196 couple, the average voltage was around 150 mV. As a consequence of using 0.05 M of
197 ferrocyanide/ferricyanide redox mediator in cathode, the average voltage increased to 275
198 mV as it shows the first period of operation of the MFC₁ and to the 188 mV reached by
199 MFC₂ during its second operation period. According to previous works [5, 12, 13], this
200 was an expected behavior because of the lower mass transfer limitations of the transport
201 of reduction agents to the cathode surface and the improved electrochemical reduction of
202 the mediator. A very important aspect to be pointed out is that in both cases, the
203 concentration of mediator in the anode is not negligible and it increases with time up to
204 values of around 150 mg dm⁻³ at the end of the 45 days-long period. This high
205 concentration could be the reason for the decrease in the production of electricity of MFC₁
206 during the last ten days of the first testing period.



207

208 **Figure 2:** a) Electricity produced during the experimental period of the MFC treating with winery
 209 wastewater in different ferricyanide concentrations in cathode: 0.05 in MFC₁ (□), 0.25 in MFC₂ (●), 0.25
 210 in MFC₁ (■) and 0.05 in MFC₂ (○). b) Temporal variation of ferricyanide in anolyte when 0.05 M of
 211 ferricyanide was used in MFC₁ (□) and MFC₂ (○) and when 0.25 M of ferricyanide in MFC₁ (■) and
 212 MFC₂ (●).

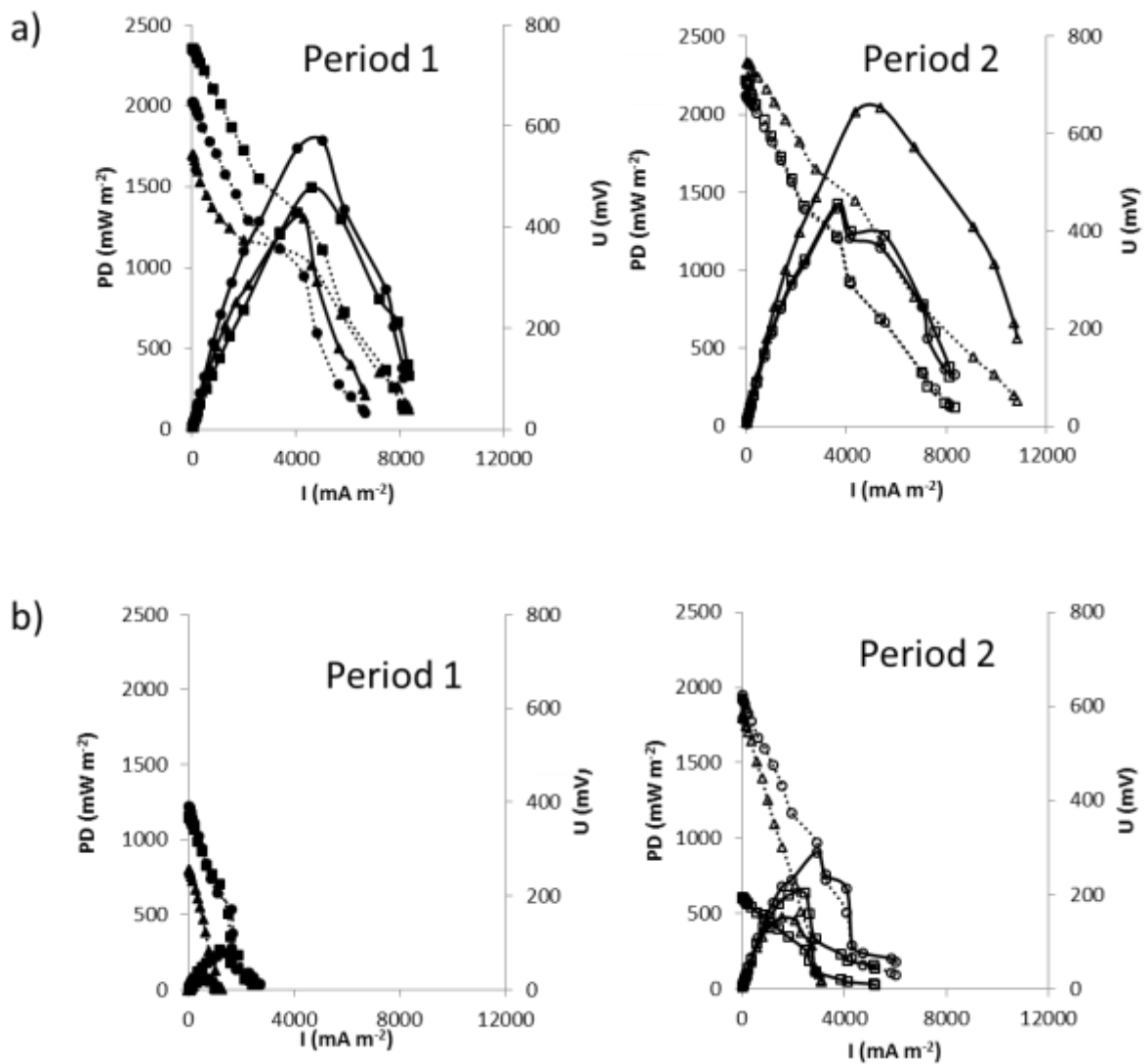
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214 Regarding the use of a higher concentration of mediator in the cathode chamber, the first
 215 period of MFC₂ showed a significantly lower value of the electricity produced, even if
 216 comparing this production with the production of the mediatorless MFC. In fact, the
 217 exponential growth phase was not observed, at least as clearly as in the mediatorless cell
 218 and in MFC₁. In this case, it can be observed that concentration of the mediator in the
 219 anodic chamber was significantly higher and a possible explanation for these worse
 220 results is that this concentration may have been affecting bioelectrogenic microorganism
 221 growth during the very sensible first acclimation period. Thus, at the end of the first
 222 testing period, the final concentration of the mediator in MFC₂ was 5 times higher than

223 in MFC₁. On the contrary, the increase of the mediator concentration in the cathode
224 chamber of MFC₁ by 5 times up to 0.25 M during second period of operation showed a
225 significant improvement in the electricity production reaching a considerable voltage
226 peak of 500 mV. However, after reaching this value it was observed a continuous drop
227 down to almost 280 mV at the end of the 35 days length of the second period. In turn, the
228 concentration of mediator in the anode chamber increases with almost the same rate that
229 in the MFC₂ operated with the same concentration of mediator, suggesting that these high
230 concentrations of mediator in the anode chamber may have been inhibiting the
231 performance of the cell [14]. Even though the permeability of membrane with
232 ferricyanide was expected to be too low, the diffusion coefficient of the ferricyanide ion
233 across the PEM membrane is $2 \cdot 10^{-8} \text{ cm}^2 \text{ s}^{-1}$ [15, 16]. As can be observe in high
234 concentration of ferricyanide in the cathode, significant amount of the ion can diffuse
235 across the membrane decreasing the MFC performance in terms of electricity generation,
236 because the microbial activity is reduced and membrane was clogged making more
237 difficult the transport of protons through the cathode chamber [17, 18].

238 Figure 3 shows polarization curves recorded at different times during all experimental
239 periods to determine the maximum power output and the main electrochemical features
240 of the MFC. It is worth to mention that the polarization curves had the same behavior
241 than the cell voltage. Thus, when the MFC was run without mediator in cathode chamber
242 (Figure 2), the MFC showed a maximum of power density of 828 mW m^{-2} (265 mV).
243 Using a 0.05 M ferricyanide concentration in the cathode chamber resulted in a substantial
244 increase in power density, with values of 1783 and 906 mW m^{-2} for MFC₁ and MFC₂,
245 respectively). By increasing the mediator concentration up to 0.25M, the maximum power
246 densities obtained were 1420 and 279 mW m^{-2} in MFC₁ and MFC₂ respectively, values

247 that were under those observed when 0.05M mediator was used and that are in agreement
248 with the discussion made before for the cell potential.



249

250

251 **Figure 3.** A) Polarization curves obtained in the MFC₁ fed with winery wastewater: period 1) 0.05 M of

252 mediator; period 2) 0.25 M of mediator. B) Polarization curves obtained in the MFC₂ fed with winery

253 wastewater: period 1) 0.25 M of mediator; period 2) 0.05 M of mediator

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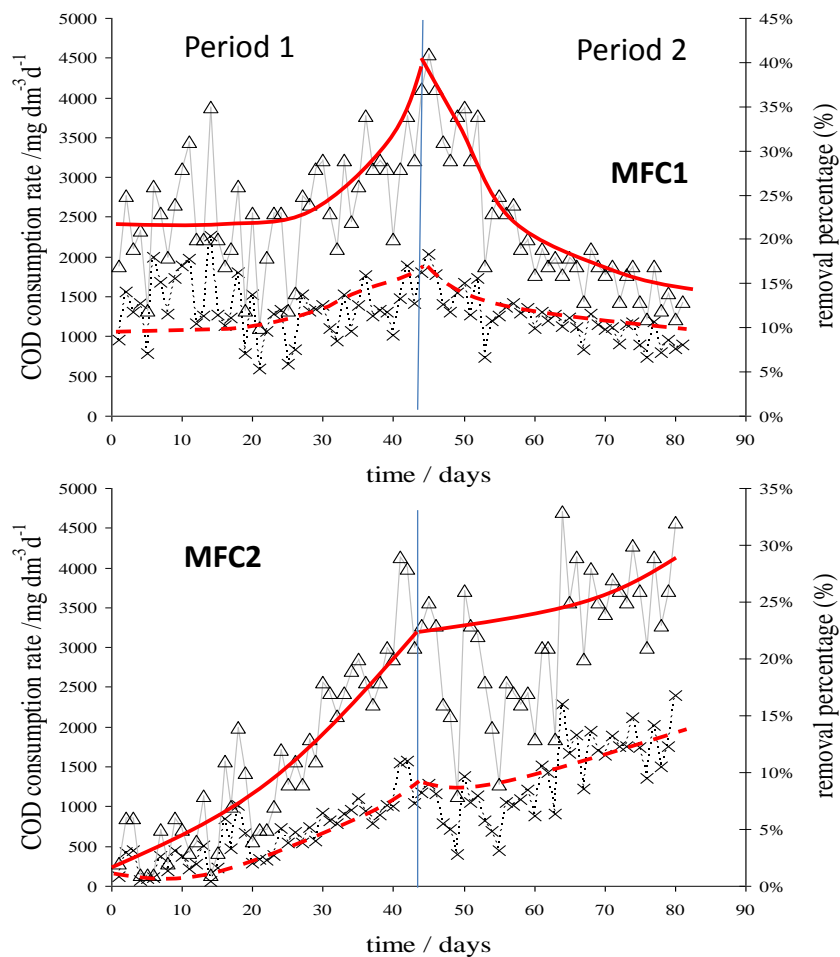
255 **COD removal.** Electricity production in MFC is carried out by bioelectrogenic

256 microorganisms, and according to the results shown, the effect of the use of a mediator

257 in the cathodic chamber of a MFC can be initially good, but its transport to the anodic
258 chamber may turn down the production of electricity.

259 The COD consumption is a very interesting parameter because it does not only inform
260 about the activity of bioelectrogenic microorganisms but it gives information about the
261 performance of the complete community of microorganisms contained in the microbial
262 fuel cell. Figure 4 shows the changes in this parameter during the performance of the two
263 MFC in the two periods studied.

264



265

266 **Figure 4:** Changes in the anodic chamber the COD consuming rate (x), the COD removal efficiency (Δ)
267 and trend (-) during the performance of the two MFC operated with mediator in the cathode chamber. The
268 MFC₁ was operated with 0.05 M of mediator in period 1 and with 0.25 M of mediator in period 2. The
269 MFC₂ was operated with 0.25 M of mediator in period 1 and with 0.05 M of mediator in period 2.

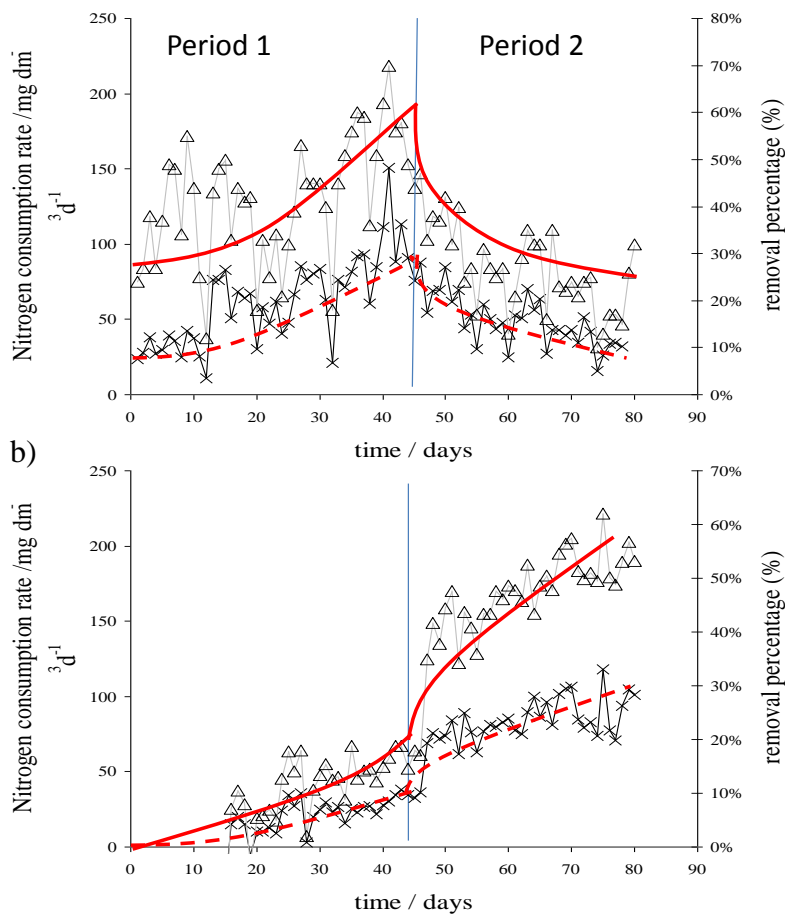
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271 As it can be observed the COD consuming rate in MFC₁ when 0.05 M of mediator
272 solution was used increased continuously until reaching a value around 1400 mg L⁻¹ d⁻¹.
273 During this period, the maximum COD removal efficiency occurred in 45th day with 41%
274 of the influent COD biodegraded. By increasing the mediator concentration in the
275 cathodic chamber of this cell up to 0.25 M, the COD consuming rate in MFC₁ decreased
276 to 1000 mg L⁻¹ d⁻¹ and the COD removal efficiency was reduced to 15%. This result
277 clearly indicates that the increase in the ferricyanide concentration in the anode chamber
278 influenced negatively the microbial activity, not only in bioelectrogenic microorganisms
279 but in the complete COD degrading community.

280 Regarding MFC₂, when it was operated with 0.25M ferricyanide, the COD consuming
281 rate and the COD efficiency removal increased during the experimental period up to
282 values of 1000 mg L⁻¹ d⁻¹ and 20%, respectively. Comparing with MFC₁, these lower
283 values was a consequence of the ferricyanide passed through PEM affecting the
284 performance of the microbial community but also indicating that microorganisms could
285 get over in the presence of important concentrations of mediator in the anode chamber.
286 However, when the mediator concentration fell by 5 times (during the second operation
287 period), it can be observed that the COD consuming rate almost doubled and the COD
288 removal efficiency was increased up to 30%. These results clearly shows that the decrease
289 in the mediator concentration in anode compartment enhanced the organic matter removal
290 by microorganisms. In relation with the literature, these values of COD removal
291 efficiency were lower than those observed by [7] and similar to the reported by [19], who
292 reported yields of about 67 % and 27 %, respectively, for the same type of wastewater.

293

294 Figure 5 shows the time course of nitrogen concentration consuming rates and the
295 resulting nitrogen removal efficiency during the complete period studied.



297

298 **Figure 5:** Changes in the anodic chamber the nitrogen consuming rate (x), the nitrogen removal efficiency
 299 (Δ) and trend (-) during the performance of the two MFC operated with mediator in the cathode chamber.

300 The MFC₁ (a) was operated with 0.05 M of mediator in period 1 and with 0.25 M of mediator in period 2.

301 The MFC₂ (b) was operated with 0.25 M of mediator in period 1 and with 0.05 M of mediator in period 2

302

303 As it can be seen, the changes were consistent with those observed in the COD removal.

304 When 0.05 M of ferrocyanide/ ferricyanide mediator solution was used in MFC₁ and

305 MFC₂, it can be seen an increase in the consumption of nitrogen during the performance

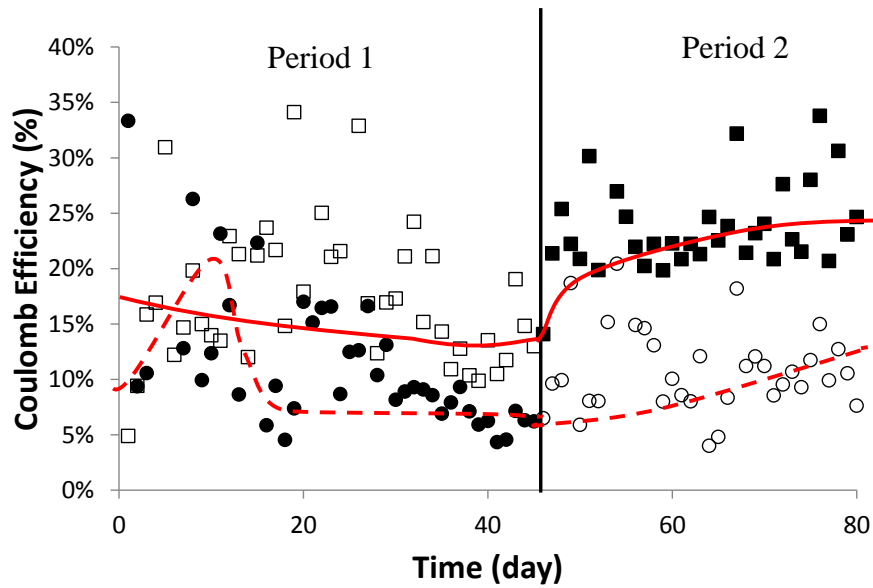
306 of both cells and the nitrogen consuming rate was around 100 mg NT L⁻¹ d⁻¹ at the end of

307 the tests. On the contrary, when MFC₁ was operated with 0.25M

308 ferrocyanide/ferricyanide mediator in the cathode chamber, the NT consuming rates were

309 slightly smaller than in other operational conditions and they settled down in a value of
310 around $50 \text{ mg NT L}^{-1} \text{ d}^{-1}$ indicating the negative effect of the transport of mediator to the
311 anode chamber. In the case of MFC_2 (that initially was operated with this mediator
312 concentration), the increase in the nitrogen consumption was much slower although there
313 was a significant increase, pointing out the capacity of adaptation of microorganisms to
314 this mediator.

315 It is worth pointing out that although significant changes in the COD consuming rate were
316 observed for the different concentrations of mediators applied in cathode chamber, the
317 activity of electrogenic microorganisms in terms of coulombic (or electric) efficiency was
318 kept almost constant as it can be clearly seen in Figure 6. When the MFC_1 was run using
319 0.05 of mediator, the Coulomb Efficiency was 16%. Because of the lower COD removal
320 and the similar production of electricity the Coulomb Efficiency increased up to 34% in
321 MFC_1 when it was operated with 0.25 M of ferrocyanide. Regarding the MFC_2 , operated
322 during the first period with 0.25 M of mediator, the Electric Efficiency increased in the
323 first moment as a consequence of low COD removal. Then the Electric Efficiency
324 decreased to around 7%. In changing the concentration of the mediator in this cell
325 dropping down to 0.05 M an increase was observed up to value around 12%. Values
326 obtained were within the typical range and suggested that other microorganisms such as
327 methanogens and sulphatogens were competing for the COD and the transport of
328 mediator affected significantly not only to bioelectrogenic microorganisms but to the
329 whole community of microorganisms contained in the anodic chamber. The important
330 crossover of the mediator through the PEM membrane advice against the use of this
331 technology for the improvement of the performance of MFC.



332

333 **Figure 6:** Electrical Efficiencies during the life test of the MFC fed with winery wastewater in different
 334 concentration of mediator in cathode chamber. The MFC₁ was operated with 0.05 M of mediator in period
 335 1 (□) and with 0.25 M of mediator in period 2 (■). The MFC₂ was operated with 0.25 M of mediator in
 336 period 1 (●) and with 0.05 M of mediator in period 2 (○).

337

338

339 **Conclusions**

340

341 There is an important influence of ferricyanide concentration on the performance of dual
 342 chamber MFC operated in semi-continuous mode, both in terms of energy recovery and
 343 COD removal. Under no crossover of mediator to the anodic chamber, performance of
 344 the MFC is improved. However, permeability through PEM lead to significant increases
 345 in the concentration of mediator in the anodic chamber, which affects both the electricity
 346 production and the COD consumption rate. Taking into account the low robustness of the
 347 membrane technology, it can be advisable not to use this type of mediator in MFC for the
 348 treatment of winery wastewater.

349

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