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Multiple electron transfer systems in oxygen reducing biocathodes revealed by different conditions of aeration/agitation

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A B S T R A C T

Oxygen reducing biocathodes were formed at −0.2 V/SCE (+0.04 V/SHE) from compost leachate. Depending on whether aeration was implemented or not, two different redox systems responsible for the electrocatalysis of oxygen reduction were evidenced. System I was observed at low potential (−0.03 V/SHE) on cyclic voltammetries (CVs). It appeared during the early formation of the biocathode (few hours) and resisted the hydrodynamic conditions induced by the aeration. System II was observed at higher potential on CV (+0.46 V/SHE); it required a longer lag time (up to 10 days) and quiescent conditions to produce an electrochemical signal. The hydrodynamic effects produced by the forced aeration led to its extinction. From their different behaviors and examples in the literature, system I was identified as being a membrane-bound cytochrome-related molecule, while system II was identified as a soluble redox mediator excreted by the biofilm. This study highlighted the importance of controlling the local hydrodynamics to design efficient oxygen reducing biocathodes able to operate at high potential.

1. Introduction

Because it is abundantly available in the environment and because its complete reduction only produces water, molecular oxygen has been targeted as the final electron acceptor in fuel cells, including microbial fuel cells (MFCs). This particular kind of fuel cell relies on bacterial biofilms that are developed on anodes and are able to catalyze the oxidation of various kinds of organic and/or inorganic matter, which makes these devices of great combined economic and environmental interest [1].

The vast majority of MFCs presented in the literature rely on abiotic oxygen reducing (OR) cathodes [2,3]. These cathodes often use expensive catalysts, e.g. derivatives of platinum-based chemicals, and are sensitive to poisoning and biofouling in natural media like the sludge or sediments used to feed MFCs. These drawbacks make them difficult to scale up for industrial applications and OR-biocathodes would constitute worthwhile alternatives. Electroactive bacteria coming from many different natural environments have been reported in OR-biocathodes, the most used inocula being aerobic sludge [4–8], seawater or marine sediment [9–14], and, to a lesser extent, soils [15,16]. The majority of studies on OR-biocathodes have implemented multispecies environmental consortia and studies using pure cultures of a single bacterium remain scarce [9, 11,17].

Various set-ups and methodologies have been used for developing OR biocathodes but analytical investigation of electron transfer mechanisms ultimately calls upon cyclic voltammetry. Different values of oxygen reduction reaction (ORR) half-wave potential (read on the reductive scan) have been obtained by various authors using this technique. As ORR potential depends on pH, values reported in works realized at unusually acidic [18] or alkaline pH [19] that would have changed the ORR potential are omitted here. Only values obtained in studies realized at neutral pH are presented in Schematic 1.

The reported potentials can be clustered in two different groups. In a first group, ORR catalysis was observed at rather low potentials, ranging between −0.1 and 0.1 V/SHE. In a second group, signals for ORR were observed at higher potentials, ranging between 0.4 and 0.5 V/SHE. Examples of “high potential” signals were notably found by Ter Heijne et al., who compared the performances of biocathodes developed from aerated sludge using different, but deliberately high, polarization potentials: 0.255, 0.355 and 0.455 V/SHE [8]. They described an identical voltammetric ORR signal centered at 0.49 V/SHE, whatever the polarization potential. The main difference between the biocathodes resided in a faster start-up time with the reactors polarized at lower potential (0.255 and 0.355 V/SHE). Xia et al. conducted a similar experiment to compare the behavior of OR-biocathodes designed at 0.142, 0.302 and 0.442 V/SHE [5]. The cyclic voltammograms showed an ORR signal at 0.46 V/SHE with a mature biocathode. They identified 0.302 V/SHE as the best polarization potential, resulting from a compromise between fast start-up and high current densities, and consequently used a similar potential, 0.311 V/SHE to design efficient OR-biocathodes for an MFC application in a subsequent report. Strycharz-Glaven et al. found a

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without forced aeration in the electrochemical reactor. Encroachments in the reductive scan at neutral pH for various biocathodes described in the literature. Fig. 1. microbial solar cell inoculated with marine sediments [10].

Freguia et al. working with Pseudomonas putrefaciens [20] and by Cournet et al. when testing the ORR activity of Cothurnus, a Gram negative one [21]. ORR signals were observed at −0.05 and −0.03 V/SHE, respectively, with these bacteria [21,22]. Marine biofilms formed with [9,11] or without [23] prior polarization produced low potential signals and voltammetric tests performed on pure strain bacteria isolated from such biofilms also presented low potential signals [9,24]. Blanchet et al. observed an ORR signal at +0.03 V/SHE during the cathodic phase of reversible bioelectrodes designed from garden compost [25]. A similar signal, centered on +0.04 V/SHE was observed during the straightforward formation of OR-biocathodes [15].

In conclusion, the data, presented in Fig. 1, revealed that:

- The low and high potential signals were observed with marine [9,10,24] or soil inocula [15,25]. Both types of signal were also observed when using the same electrode material, for example carbon cloth [15,25] or carbon paper [6,20]. In consequence, low or high potential signals are correlated neither with the inoculum source nor with the electrode material.

- Signals at low potential were recorded mainly with pure strain cultures [9,20,22,21], when no polarization was applied prior to voltammetric experiments [20,22,21], or, at the most, a polarization at low potential (minimum −0.04 V/SHE in Desmond-Le Quemener et al. [15]).

- Low potential signals appeared quickly after the experiment began (a few hours up to 2 days for Desmond-Le Quemener et al., whereas high potential signals needed a higher polarization potential (minimum 0.142 V/SHE, in [6]) and more time to develop (up to 15 days in Ter Heijne’s experiments [8]).

The importance of the polarization potential was particularly highlighted by Desmond-Le Quemener et al. Using two different polarization potentials, they experienced the coexistence of both low and high potential signals when the biocathode was formed at 0.34 V/SHE, but only the low potential signal for biocathodes formed at −0.16 V/SHE.

The poor solubility of oxygen can be an important rate limiting factor of OR-cathodes operating in aqueous media. Controlling the aeration conditions is thus a crucial element for a better understanding of OR-cathode behavior. Accordingly, different conditions of aeration were tested in this work in order to better understand the relative developments of low and high potential signals for oxygen reduction by microbial biofilms. OR-biocathodes were designed using compost leachate as the medium, under constant polarization. In a first experimental run, OR-biocathodes were either in reactors exposed to permanent forced aeration using a vigorous air flux or had no forced aeration. In the second experimental run, biocathodes initially designed under quiescent conditions were later exposed to different aeration conditions, using air or nitrogen fluxes. The electrochemical behavior of the different biocathodes was characterized by chronoamperometry (CA) and cyclic voltammetry (CV). The structures of the biofilms developed on the electrodes were imaged by epifluorescence microscopy.

2. Experimental

2.1. Experimental medium: garden compost leachate

The leachate used as medium and source of electroactive microorganisms was produced from commercially available garden compost (Eco-Terre) using the method described elsewhere [26]. The leachate was supplemented with 20 mM KCl in order to insure a minimum conductivity (4 mS·cm⁻¹). Its initial pH was 7.8.

2.2. Electrochemical set-up and tests

The electrochemical reactors were designed from Schott bottles, filled with 650 mL garden compost leachate, which left around 40 mL gas headspace. The reactors were closed by four-hole caps. Three holes were used to connect the electrodes and the fourth was devoted to the gas supplier. The working electrode was a 6 cm² piece of carbon cloth (Paxitech, Grenoble, France) connected to a platinum wire. The counter electrode was a 10 cm² platinum grid (Héraeus, France) connected to a platinum wire. Potential was controlled and expressed versus a saturated calomel reference electrode (SCE, E₀ = 0.24 V/SHE, Biologic, France), which was regularly checked versus another SCE throughout the experiment. The electrode saturation was maintained by addition of KCl crystals to the electrode compartment to avoid any drift of the potential. Gas supply (air or nitrogen) was achieved with a glass tube ending in a porous fitted glass tip. Gas fluxes were adjusted to produce gentle bubbling. Air supply brought the oxygen concentration up from 5.4 ± 0.3 to 6.5 ± 0.1 mg L⁻¹. Ten minutes of N₂ flux reduced the oxygen concentration to below 0.5 mg L⁻¹. Oxygen concentration was measured using an optical dissolved oxygen meter (Multi 3410, WTW GmbH, Germany). Temperature in the reactors was maintained at 40 °C using a thermostatic bath. All experiments...
were performed at least in duplicate. For clarity, only one set of data per experiment is presented below.

Electrochemical experiments were carried out using a multichannel potentiostat and the EC-Lab software (Biologic, France). OR-biocathodes were designed under polarization (chronoamperometry, CA) at $-0.2 \text{ V/SCE}$ ($+0.04 \text{ V/SHE}$), a potential high enough to develop both low- and high-potential signals [15]. Punctually, biocathode polarization was stopped for 30 min to allow the biocathodes to reach a stable open circuit potential (OCP) before cyclic voltammetry (CV) was performed. CVs were recorded from OCP up to 0.4 V/SCE and then down to $-0.7 \text{ V/SCE}$, at a scan rate of $1 \text{ mV s}^{-1}$. Each CV recording consisted of three successive scans, which were generally identical. For clarity, only the second scans are presented here. CVs were recorded at $t = 0$, prior to biofilm establishment, and at the end of the experiments with new, uncolonized electrodes, which served as abiotic controls.

2.3. Epifluorescence imaging

The electrodes were extracted from the reactors and washed carefully with sterile physiological water to remove all soluble and solid materials except the attached biofilms. Bioelectrodes were stained with $0.03\%$ acidine orange (A6014, Sigma) for 10 min, rinsed with sterile physiological water and then left to dry at room temperature. Biofilms were imaged with a Carl Zeiss Axio Imager-M2 microscope equipped for epifluorescence with an HXP 200 C light source and the Zeiss 09 filter (excitor HB450e 490, reflector FT 10, barrier filter LP520). Images were acquired with a digital camera (Zeiss AxioCam MRm) every $0.5 \mu\text{m}$ along the Z-axis and the set of images was processed with the Zen® software. The biovolume was calculated by the automated image analysis of the Zen® software on an acquisition stack that was representative of the colonization observed at various locations of the electrode surface.

3. Results and discussion

3.1. Effect of permanent air flux

3.1.1. Electrochemistry

Chronoamperometries at $-0.2 \text{ V/SCE}$ ($+0.04 \text{ V/SHE}$) were performed in parallel in four reactors. Reactors R1 and R2 were run with permanent forced aeration, while reactors R3 and R4 remained under quiescent conditions up to day 20, when forced air flux was introduced (Fig. 1).

Similar mean current densities of $-0.029 \pm 0.006$ and $-0.030 \pm 0.010 \text{ A m}^{-2}$ were measured initially between day 0 and day 5 whether the reactors were exposed to forced aeration or not. In the presence of forced aeration, the reduction current density remained barely higher, around $-0.038 \pm 0.008 \text{ A m}^{-2}$, up to the end of experiment. In contrast, under quiescent conditions, the reduction current started to increase strongly after 6.5 days of polarization. A stable maximal current density of $-0.31 \pm 0.02 \text{ A m}^{-2}$ was reached between day 11 and day 19. Finally, when air was forced into the electrochemical reactor on day 20, the reduction current rose back to its initial value of around $-0.035 \text{ A m}^{-2}$ within 10 h. Two stable but different current densities were thus experienced here, depending on the aeration of the electrochemical reactor.

The CV recorded with the biocathodes formed under quiescent conditions showed a reduction wave around $+0.22 \text{ V/SCE}$ ($+0.46 \text{ V/SHE}$) (noted II on Fig. 2). The CV recorded with the same biocathodes after exposing them to 24 h forced aeration showed a very different reduction wave, around $-0.27 \text{ V/SCE}$ ($-0.03 \text{ V/SHE}$) (reduction wave noted I on Fig. 2). The biocathodes formed under permanent forced aeration displayed similar CV with the reduction wave at $-0.28 \text{ V/SCE}$ ($-0.04 \text{ V/SHE}$) (wave I). The abiotic control CV carried out before the development of the biofilm was different from voltammograms with wave I and wave II. It evidenced their biotic origin.

The shape of wave II – a reduction peak followed by a plateau – was previously observed by researchers working with enzymatic OR-biocathodes under quiescent conditions [14,27]. It was the signature of a mass-transport limited reaction. Similar maximal current densities were observed with waves I and II, around $-0.44 \text{ A m}^{-2}$, indicating that both waves were limited by oxygen mass transport. The two different kinds of waves, I and II, corresponded to two different redox systems responsible for OR-catalysis and for the two different stable current densities evidenced under polarization at $+0.04 \text{ V/SHE}$.

3.1.2. Biofilm morphology

The forced aeration of the solution bulk with continuous bubbling of fine gas bubbles induced a shear stress on the electrode surface that had significant impact on their colonization and the morphology of biofilms (Fig. 3).

Under continuous air flux (R1 and R2), electroactive biofilms were diffuse, in the form of dispersed colonies. The cell density was sparse and areas of the carbon fibers making up the carbon cloth electrode were still visible even after 3 full weeks of biofilm development. In contrast, electroactive biofilms obtained without bubbling (R3 and R4) were much thicker; the carbon fibers were no longer visible in the three-dimensional biofilm images. The determination of biomass quantity by image analysis of the z-stack acquisition reported a ratio of 40 between the volume of biofilm (bio-volume) calculated on electrodes R3 and R4 and that calculated on electrodes R1 and R2. In conclusion, the shear stress induced by the forced aeration had a considerable negative impact on the colonization of the carbon cloth electrodes.

3.2. Effects of air and $N_2$ fluxes

Four OR-biocathodes were developed in independent electrochemical reactors under quiescent conditions. After 12 days of polarization, when the chronoamperometric response was maximal, two reactors were exposed to air flux for 24 h and the other two to nitrogen flux for 3 h (Fig. 4). Both gas fluxes led to a fast decrease in the reduction current.

Stationary electrochemical responses of the OR-biocathodes before gas bubbling were mostly identical, each displaying $-0.166 \pm 0.009 \text{ A m}^{-2}$ between days 10 and 12. The current was mainly due to redox system II, which was the only system detected on CV performed at this time (red CVs, Fig. 5).

When nitrogen gas was forced into the medium (Fig. 4, gray line), reductive current density immediately fell to zero and neither system I nor system II was detected by CV as long as the bubbling lasted, because oxygen was completely chased out of the medium (Fig. 5A, gray line)}
This confirmed that both systems catalyzed oxygen reduction. When the nitrogen flux stopped, oxygen reduction resumed immediately (Fig. 4, gray line), producing a current density of $-0.025 \text{ A m}^{-2}$ that corresponded to the redox activity of system I only (Fig. 5A, blue line CV). The presence of system II was not detected on CV immediately after $N_2$ flux was stopped. The chronoamperometry showed that it started to produce current again only later, after day 15.

When air was flushed through the medium (Fig. 4, black line), OR current density at $+0.04 \text{ V/SHE}$ was immediately enhanced, to $-0.35 \text{ A m}^{-2}$ for a few minutes, due to oxygen concentration increase and the higher convection. This enhancement was transient and the reduction current density decreased thereafter as the forced aeration continued. It stabilized at around $-0.05 \text{ A m}^{-2}$ after the air flux stopped. CV recorded when quiescent conditions resumed revealed that the reduction current was due to the activity of redox system I alone (Fig. 5B, blue curve). Oxygen reducing activity of system II progressively resumed on the CV recorded at the end of the experiment (green line in Fig. 5B).

### 3.3. Discussion

The two redox systems evidenced here corresponded well with the two signals previously experienced by [15]. Their respective characteristics fitted those of low- and high-potential signals shown in the literature (see introduction). Redox system I, observed around $-0.27 \text{ V/SCE}$ ($-0.03 \text{ V/SHE}$), demonstrated similar behavior to low-potential signals: fast development and a relative independence from the potential at which the biocathode was formed [15]. A contrario, redox system II, observed around $+0.22 \text{ V/SCE}$ ($+0.46 \text{ V/SHE}$), needed more polarization time to establish itself (up to 10 days) and to show an electrochemical response.

Neither of the two systems displayed an electrochemical response during nitrogen flux, which confirmed that they were both related to oxygen reduction (Figs. 4 and 5A). However, they behaved differently under air flux. The air flux never altered the signal produced by redox system I, which showed OR-activity under permanent air supply (Figs. 1 and 2). This system demonstrated an enduring OR-activity when air was supplied in reactors previously conditioned in quiescent conditions (Figs. 4 and 5B). The OR-signal due to system II was observed only under quiescent conditions and, after a brief enhancement, it vanished when air was supplied in the reactors (Figs. 1, 2, 4 and 5B). Forced aeration brought more oxygen into the reactor and enhanced signal II for a brief moment but it ultimately led to its extinction. Due to its oxidative properties, oxygen at high concentrations could be toxic for microbial cells. However, its concentration only changed from $5.4 \pm 0.3$ to $6.5 \pm 0.1 \text{ mg L}^{-1}$ during air flux, and this moderate increase is unlikely to have been responsible for the loss of signal II through oxygen toxicity.

Forced aeration induced shear forces on the electrode surface, which impacted the electrode colonization (Fig. 3). Shear forces are known to strongly influence the settlement, the structure and the morphology of biofilms [28,29]. Other studies on the formation of biofilms from natural consortia or pure microbial strains under various hydrodynamic conditions have shown that the thickness, the amount of fixed biomass, the nature and amount of exopolymeric substances produced and the distribution of bacterial colonies vary widely depending on the shear mode applied [30,31]. It is generally agreed that the greater the physical...
force applied to the biofilm is the more resistant, compact and thin the biofilm becomes [32,33]. Under high shear stress, biofilm detachment could occur via erosion, resulting in thinner biofilms that are rather smooth and rich in microbial cells in the colonized areas. Here, such a phenomenon of biofilm erosion should impact redox systems I and II similarly. If erosion was responsible for the disappearance of system II observed under forced aeration, it would similarly cause the destruction of system I. Consequently, biofilm erosion cannot explain the behavior of the biocathodes under forced aeration.

The most likely explanation of the difference between the two redox system behaviors is that they constituted two different mechanisms for external electron transfer (EET) from the electrode to oxygen. Different types of EET mechanisms from cathodes to various final electron acceptors were reported in the literature reviewed by Rosenbaum et al. [34].

The CV characteristics of system I were very close to those previously reported by Blanchet et al., who worked with the same inoculum (compost leachate) and polarization potential (+ 0.04 V/SHE) [25]. Actually, system I characteristics were similar to many low-potential signals reported in other works performed with other media and/or inocula [24, 22,21]. The voltammograms presented in these works showed characteristics (potential, current density, fast development even without polarization) that may link them to redox system I observed here. Parot et al. demonstrated a close similarity between CVs recorded with marine bacterial strains and those of an iron porphyrin adsorbed on the electrode surface [24]. Iron porphyrins are redox molecular substructures present in numerous proteins such as cytochromes. Cytochromes have been proven to achieve direct external electron transfer between bacteria and electrodes [35]. They are generally anchored in cellular membranes and are consequently not removed from the biofilm by hydrodynamic effects. Therefore, all these observations led us to postulate that redox system I may be related to membrane-bound cytochromes.

Another known EET pathway relies on the secretion of soluble mediators used for shuttling electrons. A large variety of bacterial extracellular redox mediators have been identified in both bioanodes and biocathodes. Among them are phenazines such as piocyanine [36], flavin derivatives such as riboflavin [37], or quinones such as pirroloquinoline quinone and 2-amino-3-dicarboxy-1,4-naphtoquinone [20,38]. Such molecules are usually locally concentrated inside the biofilm, which ensures proximity between bacteria and electrodes and facilitates the electrochemical transfer. The soluble mediators are subjected to mass transport; they can be dispersed in the bulk in presence of increased hydrodynamic forces and thus lose the possibility to play their electron mediator function. As redox system II behavior fitted this description in all respects, it probably corresponded to a diffusible extracellular redox mediator that was poorly linked to the biofilm and was easily released to the bulk by hydrodynamic conditions.

The nature of the soluble redox mediator postulated here for system II cannot be enlarged to all similar high-potential systems reported in the literature, because they have shown different sensitivities to hydrodynamics. The high-potential system observed by Desmond-Le Quemener et al. using the same inoculum (compost leachate) was not sensitive to hydrodynamics [15]. This difference may have come from the higher polarization potential they used to form their biocathode (+ 0.34 V instead of + 0.04 V/SHE), which may have resulted in selecting other electron exchange pathways between electrode and biofilm. Xia et al. used a continuously-fed electrochemical cell with forced aeration to form the biocathode, but prevented problems caused by the hydrodynamic conditions by interposing a plastic plate in the cathodic compartment [6,5].

Ter Heijne et al. used an electrochemical cell in continuous mode [8]. This operating mode may imply a certain degree of shear stress on the cathode surface, but it did not impact the electrochemical behavior of their system. Moreover, increasing the flow rate from 0 to 21 L h$^{-1}$ led to increased current densities for OR from 40 to 270 mA m$^{-2}$. Strycharz-Graven et al. identified the presence of immobilized cofactors inside the biofilms, which achieved a direct, non-mediated EET [10].

Due to its higher redox potential, system II is of greater interest than system I if the objective is to design efficient OR-biocathodes for MFC applications. The results presented here, and the past works referred to, recall that high potential oxygen reductive systems are observed when rather high polarization potentials are applied to form the OR-biocathodes. This highlights the benefit of designing OR-biocathodes under polarization. However, as reflected by the discussion above, the nature of redox system II may hinder the development of practical applications. Its sensitivity to turbulence may restrict its application to systems in which quiescent conditions are ensured. Continuous flow MFCs, for example, may be suitable provided that high retention time and low flow rate are used to mitigate the possible release of the redox mediator from the biofilm. Precautions as implemented by Xia et al., i.e. a plastic plate interposed between the aeration source and the biocathode, would certainly be of interest [6,5]. The use of a small-volume cathodic compartment in order to limit the dilution effect of the mediator in the solution bulk would also merit attention.

4. Conclusions

Oxygen-reduction biocathodes formed from compost leachate under constant polarization at + 0.04 V/SHE showed two different
redox systems linked to oxygen reduction. A low-potential system (I) was detected in all bioanodes, whatever the experimental conditions, and ensured a basal oxygen reduction activity. A high-potential system (II) developed only under quiescent conditions and vanished when the solution was stirred. In the light of the literature, system I was probably a cytochrome derived compound, firmly attached in the biofilm system. System II behavior was characteristic of a soluble redox mediator, the chemical nature of which remains to be determined. Its solubility made it sensitive to turbulence inside the reactor. This work shows that implementing efficient oxygen-reducing bioanodes in MFCs requires careful control of the local hydrodynamics in order to favor the development of the high-potential redox system.

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Appendix A. Supplementary data

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