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Anodic microbial community analysis of microbial fuel cells based on enriched inoculum from freshwater sediment

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1 Anodic microbial community analysis of microbial fuel cells based on enriched

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Abstract

The characterization of anodic microbial communities is of great importance in the study of microbial fuel cells (MFCs). These kinds of devices mainly require a high abundance of anode respiring bacteria (ARB) in the anode chamber for optimal performance. This study evaluated the effect of different enrichments of environmental freshwater sediment samples used as inocula on microbial community structures in MFCs. Two enrichment media were compared: ferric citrate (FeC) enrichment, with the purpose of increasing the ARB percentage, and general enrichment (Gen). The microbial community dynamics were evaluated by polymerase chain reaction denaturing gradient gel electrophoresis (PCR-DGGE) and real time polymerase chain reaction (qPCR). The enrichment effect was visible on the microbial community composition both during precultures and in anode MFCs. Both enrichment approaches affected microbial communities. Shannon diversity as well as β -Proteobacteria and γ -Proteobacteria percentages decreased during the enrichment steps, especially for FeC (p<0.01). Our data suggest that FeC enrichment excessively reduced the diversity of the anode community, rather than promoting the proliferation of ARB, causing a condition that did not produce advantages in terms of system performance.

Keywords: Freshwater sediment; microbial communities; DGGE; real time qPCR; MFC.

Introduction

A biological approach to the study of microbial electrochemical technologies (METs) can increase knowledge within the microbial electrochemistry field. The need for a better understanding of anodic microbial community composition is of great importance in studying microbial electrochemical cells, including microbial fuel cells (MFCs). MFCs are biocatalyzed systems able to convert chemical energy into electrical energy using anaerobic respiration by electroactive microorganisms known as anode respiring bacteria (ARB) [1, 2]. Typically, they consist of two compartments separated by a proton exchange membrane (PEM) with an external circuit connecting anode and cathode electrodes. In these devices, the anode biofilm acts as a biocatalyst to hydrolyse the substrate and release protons and electrons in MFCs; therefore, the higher the amount of ARB in the anodic biofilm, the higher the electrical energy production [3]. In MFCs, the anode acts as a terminal electron acceptor in the same manner as any other natural acceptor, e.g. oxygen, nitrate or Fe(III) [4, 5].

It is known that electrochemically active microorganisms can interface with the electrode in several ways
[6], although exoelectrogenic mechanisms in microbial species are still an open field of research, and
many different new microbial strains show extracellular electron transfer capacities [7]. ARB can be
found and enriched from many different environmental sources, such as freshwater and marine sediments
salt marshes, anaerobic sludge, industrial effluent and sludge from wastewater treatment plants [1, 5, 8,
9]. The microbial community composition is affected by the inoculum source and the type of enrichment
as well as by the more thoroughly investigated system design and operating parameters [3].
Different approaches have been reported in the literature to optimize the formation of a high performing
anode microbial community [10, 11]. However, most of them are based on the use of already formed
biofilm, or other elements of pre-existing bioelectrochemical systems [12–14]. On the contrary,
enrichment procedures directly acting on the inoculum represent a more efficient approach [5].
Considering the substantial agreement on the pivotal role of ARB in MFC microbial community, it has
been proposed to promote their specific selection by means of iron enrichment methods [5, 15]. To this
aim, it is possible to use ferric citrate (FeC), which selects for microorganisms that are able to reduce
Fe(III); meanwhile, other bacteria without the required ability are eliminated, consequently increasing the
percentage of electrochemically active bacteria [8]. Pierra et al. [5] evaluated the use of an iron
enrichment method to target dissimilatory metal-reducing bacteria. Sathish-Kumar et al. [15] compared
an Fe(III) enrichment method against an electrochemical procedure as well as a combination of the two.
According to literature, enrichment with Fe(III) citrate allowed the selection of ARB to mimic the role of
a MFC carbon electrode working as an electron acceptor [6, 15, 16].
Different opinions are reported in literature about the correlation between microbial diversity and the
performance of MFCs. Torres et al. showed that the lower-diversity type of MFC exhibits higher
performance [17], but some years later, Stratford et al. obtained a strong correlation in a regression model
between the power output of the system and the Shannon index, which was then proposed as a predictor
of good performances [18].
Yamamoto et al. pointed out the relevance of the analysis of both the planktonic and attached
components; in his study, he showed that the correlation between biofilm and planktonic microbes was
important to achieve better performance [19].
In the present study, experiments were designed to find the reason why the performance of Fe(III)-type
MFC was lower than the control MFC.

77	In line with this, we investigated the impact of freshwater sediment enrichments on the structure and
78	composition of microbial community sampled from planktonic component, carbon felt biofilm and
79	graphite rod biofilm.
80	The effect of a Fe(III) enrichment procedure was compared with a general (non-specific) enrichment.
81	Electrochemical performance and biofilm morphology were previously evaluated in Agostino et al. [20].
82	A combination of denaturing gradient gel electrophoresis (DGGE), a community structure technique and
83	real time quantitative polymerase chain reaction (qPCR), a population technique [21] was applied to
84	investigate the changes in and the diversity of the microbial community.
85	This work represents the first attempt to describe the effect of ferric citrate enrichment on a microbial
86	community in a MFC system from a biomolecular point of view.
87	Materials and Methods

A freshwater sediment sample (Bagnère Creek, Valle D'Aosta, Italy) was enriched under anaerobic

Experimental set up

conditions with two different media: a Ferric Citrate (FeC) medium and a General (Gen) medium. Sodium acetate was used as an electron donor and carbon source in both enrichment methods; media composition and operational protocol were as previously reported [20]. Briefly, the composition of FeC medium was the following: Fe(III) citrate 13.70 g/L; NaHCO₃ 2.50 g/L; NH₄Cl 1.50 g/L; NaH₂PO₄ 0.60 g/L; KCl 0.10 g/L; Na acetate 2.50 g/L; Wolfe's Vitamin solution 10 mL/L (ATCC) and Wolfe's trace mineral solution 10 mL/L (ATCC). The composition of the Gen medium was the following: NH₄Cl 1.50 g/L; NaH₂PO₄ 2.45 g/L; Na₂HPO₄ 4.28 g/L; KCl 0.10 g/L; Na acetate 2.50 g/L; Wolfe's Vitamin solution 10 mL/L (ATCC) and Wolfe's trace mineral solution 10 mL/L (ATCC).

The microbial cultures were subjected to three sequential enrichments for 21 days of total growth at room temperature (21 ± 2 °C) and with gentle orbital shaking (150 rpm). They were then inoculated into the two-chamber MFCs, with a ratio of 10% v/v of the total anode volume. At each step, 10% (v/v) of the microbial cultures were inoculated in fresh anaerobic media. Before inoculation, the media were purged by high-speed N₂ flow for 15 minutes in order to reach anaerobic conditions. Biofilm formation into the anodic chamber was carried out by applying a low external resistance (47 Ω), resulting in a positive anode potential polarization.

MFCs operated in continuous mode with a hydraulic retention time of 5 days (0.5 mL/h). The analyte consisted of 1 g/L per day of CH₃COONa and 0.31 g/L per day of NH₄Cl dissolved into a phosphate buffer solution (PBS: NaH₂PO₄ 2.45 g/L; Na₂HPO₄ 4.28 g/L; KCl 0.10 g/L) with 10 mL/L of Wolfe's vitamin solution (ATCC) and 10.00 mL/L Wolfe's trace mineral solution (ATCC). The catholyte was comprised of 6.58 g/L K₃[Fe(CN)₆]) dissolved into PBS. Carbon felt (Soft felt SIGRATHERM GFA5, SGL Carbon, Germany) was used as the material for anode and cathode electrodes. A cation exchange membrane (CEM, CMI-7000, Membranes International Inc., USA) was used to separate the two compartments. Electrical contacts to the electrodes were made using graphite rods. The experiments were performed in duplicate, at room temperature conditions (20-22°C), and lasted 90 A data acquisition system (Agilent 34972A) was used to monitor cell voltage continuously across the external resistor and anodic potentials. Electrochemical characterization and bioanode imaging analysis were performed as previously reported [20] to evaluate the performances of the devices. Polarization curves were obtained at the end of the biofilm acclimation phase during changing external circuit resistances. Anode impedance spectra were recorded using a multi-channel VSP potentiostat in a 3-electrode configuration for each polarization condition. Cyclic voltammetry (CV) was performed using the same potentiostat to obtain the putative electron transfer redox centre. Bioanode imaging was acquired by fluorescent microscopy to characterize the biofilm distribution within the electrode after LIVE/DEAD staining. DNA extraction Carbon felt biofilm and graphite rod samples were subjected to a pre-treatment; 1.25 g each of wet anode electrode and graphite rods were washed twice with 4 mL of 0.9% NaCl. Supernatants were centrifuged

Carbon felt biofilm and graphite rod samples were subjected to a pre-treatment; 1.25 g each of wet anod electrode and graphite rods were washed twice with 4 mL of 0.9% NaCl. Supernatants were centrifuged for 20 min at 10000 rpm. Pellets were re-suspended in 0.9% of NaCl solution.

DNA was extracted from each sample using a commercial kit (UltraClean™ Microbial DNA Isolation Kit, MO-BIO Laboratories, Inc., Carlsbad, CA), following manufacturing information. Genomic DNA integrity was checked by electrophoresis gel on a 2% agarose gel and 1X TBE (Tris-Borate-EDTA) buffer after each extraction as previously described [22].

133	The quantification of the extracted DNA was performed by fluorometric quantification using a Qubit™
134	Fluorometer and a Qubit™ dsDNA HS Assay by Invitrogen (Life Technology, Ltd., Paisley, UK)
135	according to manufacturer instructions.
136	PCR-DGGE and Sequencing
137	Primer 357F with GC clamp and 518R were used to amplify the V3 region of the 16S rRNA genes from
138	the bacterial community [23]. The PCR amplification was performed in a 50-µl volume containing 0.2
139	μM of each primer, 0.2 mg/ml of Bovine Serum Albumin (BSA) and 1X of Master Mix for PCR (Bio-
140	Rad).
141	PCR was performed in T100 thermal cycler (Bio-Rad, Italy) as follows: ten cycles of denaturation at 94
142	°C for 30 s, annealing at 55 °C for 30 s and extension at 72 °C for 1 min; twenty-five cycles of
143	denaturation at 92 °C for 30 s, annealing at 52 °C for 30 s and extension at 72 °C for 1 min; followed by a
144	single final extension at 72 °C for 10 min.
145	The PCR products were approximately 190 bp in length. DGGE was carried out using the DCode TM
146	Universal Detection System (Bio-Rad Laboratories, CA, USA) as previously described by Webster et al.
147	[24]. Ten microliters of the PCR product were loaded onto 8% polyacrylamide gels (acrylamide: bis-
148	acrylamide, 37.5:1) with denaturing gradients ranging from 30% to 50% (where 100% denaturant
149	contains 7 mol L-1 urea and 40% formamide) in 1X TAE buffer. The electrophoresis was run at a constant
150	voltage of 200 V at 60 °C for 5 h. After that, the gel was stained with SYBR® Green I nucleic acid gel
151	stain (Sigma-Aldrich), visualized on a UV transilluminator and photographed (Gel Doc XR+ System,
152	Bio-Rad). The computerized images of DGGE profiles were analysed with the Quantity One software,
153	Version 4.6.7 (Bio-Rad Laboratories, CA, USA).
154	DGGE bands recurrent at the site level, or shared among different sites, were excised and rinsed in 50 μL
155	of deionized water. The gel bands were then crushed in 10 μL of sterile mmQ water and stored at -20 $^{\circ}C.$
156	DNA extracts from excised DGGE bands were used as templates and PCR was performed as described
157	above, except for the elimination of BSA and the employment of modified bacterial reverse primers
158	(357F-GC-M13R and 518R-AT-M13F), as previously described [25]. PCR products were sent to the
159	Genechron (Ylichron S.R.L.) laboratory for Sanger sequencing. The sequences were then compared with
160	the NCBI database using nucleotide Basic Local Alignment Search Tool (BLASTn) analysis
161	(http://www.ncbi.gov/).

162	Real time qPCR
163	DNA was also used for qPCR absolute quantification assays. Specific primers targeting different bacterial
164	phyla and classes were selected from the scientific literature based on sequence analysis (Table 1).
165	Moreover, we selected primers to detect typically electroactive microorganisms such as <i>Geobacteriaceae</i>
166	and Pseudomonas spp.
167	The qPCR reactions targeting specific regions of 16S rRNA were performed with SYBR® Green
168	chemistry in 20 μ L total volume of SsoAdvanced TM Universal SYBR® Green Supermix (Bio-Rad, Italy)
169	$1X,2~\mu L$ of $1{:}10~DNA$ as template and $250~nM$ of each primer. Total bacteria were quantified with
170	TaqMan® chemistry in 20 μL total volume of iQ TM Multiplex Powermix (Bio-Rad, Italy) 1X, 2 μL of
171	1:10 DNA as template, 250 nM of each primer and 100 nM of probe.
172	Each reaction was performed in triplicate with a CFX96 Touch TM Real-Time PCR Detection System
173	(Bio-Rad, Italy). The bacterial concentration from each sample was calculated by comparing the threshold
174	cycle values obtained from the standard curves using the CFX Manager™ software. Standard curves for
175	absolute quantifications were constructed using 10-fold serial dilutions of specific standard genomics
176	(Table 1); the number of bacteria was expressed in terms of the number of gene copies, which is
177	comparable between different samples.
178	The different strains used were obtained from ATCC (<i>Alcaligenes faecalis</i> ATCC® 8750D-5™,
179	Bacteroides fragilis ATCC® 25285D-5™, Clostridium difficile ATCC® 9689D-5™, Desulfovibrio
180	vulgaris ATCC® 29579D-5™, Geobacter metallireducens ATCC® 53774D-5™, Pseudomonas
181	aeruginosa ATCC® 15442 TM).
182	Negative controls containing all of the elements of the reaction mixture except template DNA were
183	performed in every analysis and no product was ever detected. The amplification efficiency of the qPCR
184	for all primer pairs was determined using the linear regression slope of a dilution series.
185	Reaction protocols are reported in Table 2. Melt curve analysis was performed at the end of each
186	amplification reaction, with the exception of total bacteria, by slowly heating the qPCR products from 65
187	$^{\circ}$ C to 95 $^{\circ}$ C, in increments of 0.5 $^{\circ}$ C for 5 seconds with simultaneous measurement of the SYBR Green
188	signal intensity. Melting-point-determination analysis allowed the confirmation of the specificity of the
189	amplification products. All qPCRs were considered valid if they had linear standard curves with an $R^2\!>\!$
190	0.980 and an efficiency between 90 and 105% (Bio-Rad, Real-Time PCR Applications Guide).

Data analysis and statistics

The DGGE profiles were compared using cluster analysis (BioNumerics software, version 7.6, Applied

Maths, Ghent, Belgium) using band-based similarity coefficients (i.e. Jaccard coefficients) for the

194 construction of similarity matrices and the UPGMA algorithm was used to obtain the dendrograms [33–

37].

The Shannon index was calculated using BioNumerics software version 7.6 (Applied Maths, Ghent,

197 Belgium). Absolute and relative quantifications were calculated using the qPCR data.

Student's t-tests and one-way ANOVAs with a Tukey's post hoc analysis were performed to compare two

or more groups of independent samples.

For the Student's t-test, variance homogeneity was first assessed using the Levene's test; thus, an equal

variance for Tukey's test was assumed for multiple post hoc comparisons. The differences between means

were considered significant at p < 0.05. Statistical analysis was performed using SPSS software (version

203 24.0 for Windows).

Results and Discussion

Freshwater sediment samples were anaerobically enriched with two different media, with and without

Fe(III)citrate in order to evaluate the effect of enrichment on the microbial community that developed in

the anode chamber of the MFCs. Each third sequential enrichment was then inoculated into the MFCs and

acclimatized.

Enrichment effect on precultures

The effect of both kinds of enrichment (i.e. FeC or Gen) on the freshwater sediment was already detectable after the first preculture step. DGGE analysis showed all enrichment steps differed considerably from the freshwater sediment sample (Jaccard similarity = 28.0%). The similarity between the Gen enrichment precultures was higher than between the FeC ones (Jaccard similarity = 76.9% and 52.9%, respectively) (Figure 1a). The presence of an anaerobic environment and FeC in the medium affected the diversity of the microbial community; the Shannon diversity index decreased during the enrichment steps, especially in the third FeC preculture (ANOVA, Tukey's post hoc: p < 0.001) (Figure 1b). This can be interpreted as a marker of the effect of the specific enrichment on the microbial

community. In fact, Shannon diversity indexes provide information about richness (the number of present species) and evenness (how abundances are distributed across species), and was proved to be positively correlated with power output [18]. Sequencing analysis revealed that, in our samples, the majority of bacteria belonged to the Proteobacteria and Firmicutes phyla (Table 3). Microorganisms belonging to β-Proteobacteria, γ- Proteobacteria, ε-Proteobacteria, δ-Proteobacteria and Bacilli classes dominated our community, which was in line with previous literature [8, 38]. Many uncultured bacteria such as Comamonas spp., Dysgonomonas spp., Acrobacter spp., Alcaligenes spp. and Citrobacter spp. were also detected, probably because of the environmental origin of the inoculum [39]. Coherently with the sequencing, quantitative analysis of the main microbial components revealed that the inoculum was mainly comprised of β -Proteobacteria (36.1%) and γ -Proteobacteria (41.0%) (Figure 2). Their percentages decreased during enrichment steps, especially in the third FeC enrichment steps (β-Proteobacteria, < 1% and γ-Proteobacteria, 3.6%), confirming that the more selective FeC enrichment method has a major effect on the equilibrium of the microbial community when compared to the Gen enrichment method [20]. β-Proteobacteria subclasses consist of several groups of aerobic or facultative bacteria, which are often highly versatile in their degradation capacities [40]. Decrease in their relative quantities during the enrichment steps was possibly due to an anaerobic environment during precultures. The microbial community at the third Gen enrichment step resulted in a population more dominated by Proteobacteria (beta, gamma and delta classes) and Bacteroidetes phyla as compared to the first FeC step (22% and 3% vs 4% and 1%, respectively). Only the Firmicutes phylum had higher percentage at the third FeC enrichment step than at the Gen steps (1% vs 0.3%, respectively) (Figure 2). Electrochemically active microorganisms, such as Geobacteriaceae spp. and Pseudomonas spp., decreased more in FeC enrichments than in general ones (Figure 3). FeC enrichment steps negatively affected Geobacteriaceae spp. (ANOVA, Tukey's post hoc: p < 0.001). On the contrary, they did not differ between the inoculum and third Gen enrichment step (ANOVA, Tukey's post hoc; p > 0.05). Pseudomonas spp. decreased at a statistically significant rate throughout the steps of both Gen and FeC enrichment (ANOVA, Tukey's post hoc: p < 0.01), especially in FeC enrichment. The lower diversity and presence of these microorganisms, both in absolute and relative quantification, in the third FeC enrichment step as compared to the Gen steps explains the performance of the devices

inoculated by Gen enriched preculture: Gen-MFCs exhibited higher current and power density than FeC-MFC ones ($74 \pm 4 \text{ mA/m}^2 \text{ vs } 50 \pm 3 \text{ mA/m}^2$; $79 \pm 12 \text{ mW/m}^2 \text{ vs } 38 \pm 2 \text{ mW/m}^2$, respectively) and shorter start-up time (5 days vs 10 days, respectively) (data shown in [20]). Although previous studies showed that the FeC enrichment improved MFCs' performance [5, 15], this could be dependent on its concentration. Of note, in a recent work by Liu et al., [41] optimal community development was obtained at a Fe(III) concentration much lower than the one used in the present research.

Enrichment effect on MFC communities

As observed for the preculture steps, FeC enrichment also affected the microbial community developed in MFCs anodes. Biological analyses performed on the MFCs anodic compartments revealed that the kind of enrichment is the main source of diversity (similarity 48.7%) (Figure 4a). Even though qPCR showed no differences between Gen-MFCs and FeC-MFCs for all types of strains researched (t-test, p > 0.05), Shannon diversity was higher in the Gen-MFCs when compared with those of FeC-MFCs (t-test, p < 0.05) (Figure 4b). As observed during the early steps of the test, and until the end of the start-up time, the Shannon diversity, which was strongly associated with power [18], could explain the better performance of Gen-MFCs. Three-electrode EIS analysis suggested a more efficient electron transfer mechanism in the Gen-MFCs' bioanodes as opposed to the FeC-MFCs' bioanodes. By this impedance analysis, it is possible to recognize two features: a high-frequency process, which is related to the electron charge transfer (activation resistance) and a low frequency process, accounting for the anodic biofilm masstransfer limitation (diffusion resistance), mainly dependent upon the diffusion of the organic substrate in the biofilm. Gen-MFCs bioanode, with its higher Shannon diversity and more dense and active mixed consortia, is associated to a higher consumption rate of substrate, that decreases diffusion resistance, and hence accelerates electrons-transfer mechanisms, with respect to FeC-MFCs, where the diffusion time constant is about 4 times lower than Gen-MFCs. This resistance strongly depends on the applied external resistances, which are higher at open circuit voltage conditions and lower at the maximum power point. Moreover, the presence of a higher percentage of dead/inactive bacteria covering the interface between the bulk solution and the anodic electrode of FeC-MFCs, which was detected by Fluorescence Microscopy (Agostino et al., [20]) contributed to an increase in the resistance related to the interfacial process, i.e. double layer capacitance.

DGGE analysis showed higher similarity between the anode suspension and the carbon felt biofilm, as shown by the dendrogram in Figure 4a. Jaccard similarity of 92.3% and 84.6% was found for suspension and carbon felt biofilm of Fe-MFCs and Gen-MFCs, respectively. This is, to some extent, an unanticipated result, since higher similarity between the carbon felt biofilm and the graphite rod biofilm might be expected. In fact, graphite rod and anode carbon felt are constantly in contact. This unexpected result could be due to the carbon felt properties. Indeed, it is a porous material, and during the experiment, it was soaked in the anode medium. Thus, at the moment of the analysis, it also contained suspension, which could have led to high similarity between bacteria communities of anode biofilm and planktonic component. Real time qPCR analysis at the end of the MFCs' operation suggested that the community was dominated by β-Proteobacteria both in planktonic samples and the attached component (i.e. carbon felt biofilm and graphite rod biofilm) of all MFCs (Gen-MFCs: 26.13% and 24.42%; FeC-MFCs: 44.45% and 37.06%, respectively) (Figure 5). B-Proteobacteria in the planktonic component was statistically significantly higher in the FeC-MFCs than in Gen-MFCs (ANOVA: Tukey's post hoc, p < 0.05). These data confirmed the outcome of the sequencing analysis (Table 4).β-Proteobacteria was found to be the most abundant class within the Proteobacteria phylum in numerous previous studies using two-chamber MFCs, with different inocula and a variety of substrates, like synthetic wastewater or a liquid fraction of pig slurry (see for example [42, 43]). Interestingly, δ -Proteobacteria percentage increased from the inoculum (2.17%) to the end of the experiment (Gen-MFCs: 22.36% and 22.78%; FeC-MFCs: 22.85% and 15.87% in planktonic and attached components, respectively). Moreover, their percentage was doubled on the carbon felt biofilms compared to the graphite rod biofilms of Gen-MFCs (24.19% vs 12.99%). On the other hand, the reverse condition was found for FeC-MFCs (15.23%vs 31.35%). As in Chae et al. [40], δ -Proteobacteria were the second most frequently detected bacteria class in MFCs. The δ-Proteobacteria class, including Geobacteraceae spp., was found to be statistically significantly higher in the attached component of Gen-MFCs than in the FeC ones (ANOVA: Tukey's post hoc, p < 0.05). Differently from δ -Proteobacteria, the percentage of γ -Proteobacteria, which are largely facultative anaerobes [7], decreased from inoculum (40.96%) to end of the experiment (Gen-MFCs: 17.98% and 14.28%; FeC-MFCs: 17.28% and 9.90% in planktonic and attached component, respectively). The γ -Proteobacteria class was found in statistically significantly higher amounts in the attached component of

Gen-MFCs (ANOVA: Tukey's post hoc, p < 0.05). Bacteroidetes were found to be statistically significantly higher in all components of Gen-MFCs (ANOVA: Tukey's post hoc, p < 0.05), although their percentages were quite similar in both Gen and FeC-MFCs, ranging between 1.79 and 2.53%. Bacteria in the Firmicutes phylum, which contains both obligate anaerobes (such as *Clostridia* spp.) and facultative anaerobes (such as *Bacilli* spp.) [7] was a smaller component in the MFCs, as they already were in the inoculum. Their percentage was always less than 0.1%. Firmicutes quantification indicated their higher presence in the attached component instead of in the suspension of both types of MFCs (ANOVA: Tukey's post hoc, p < 0.05); their presence was also higher in the FeC-MFCs attached components than in Gen ones (ANOVA: Tukey's post hoc, p < 0.05). Along with Proteobacteria, Firmicutes is often among the predominant phyla composing the anode biofilm, regardless of configuration, inoculum or substrate [38, 42, 44, 45]. However, qPCR detected it in low percentages in each step of our analysis, for either FeC or Gen enrichment. This result could depend on the specific freshwater inoculum used for our work. As was pointed out by Hu et al. [46], Firmicutes seem to be more characteristic of planktonic samples, rather than sediments, as it was the case of the present study. Real time data confirmed the high similarity between suspension and carbon felt biofilms for both Gen-MFCs and Fe-MFCs; the Gen planktonic component was statistically similar to Gen biofilm for all kinds of strains with the exception of Geobacteriaceae and Pseudomonas spp., which were higher in the biofilm (ANOVA: Tukey's post hoc, p < 0.001) [7, 47], as well as higher in their relative percentage. Similar behaviour is shown by FeC-MFCs; the planktonic component that differs from carbon felt biofilm only for the large number of total bacteria and Geobacteriaceae spp. in the biofilm (ANOVA: Tukey's post hoc, p < 0.001). In comparing the attached components in the MFCs, we can confirm that, from CV analysis (data shown in [20]), Geobacteriaceae spp. had the main role in the electron transfer in Gen-MFCs (midpoint potential equal to -0.4 V vs Ag/AgCl) and *Pseudomonas* spp. had the main role in FeC-MFCs (midpoint potential equal to -0.215 V). Real time quantification showed Geobacteriaceae spp. higher in Gen-MFCs than in FeC-MFCs and Pseudomonas spp. higher in FeC-MFCs than in Gen MFCs, even if there was not a statistically significant difference (ANOVA: Tukey's post hoc, p > 0.05). The lack of significant variation within the data of that comparison could be explained by the presence of a higher number of dead microorganisms in the FeC anode biofilms according to quantification. Indeed, qPCR quantifies both living and dead microorganisms; therefore, we had to take in to account the morphological

characterization of the anode microbial biofilm by fluorescence microscopy; the ratio between living and dead microorganisms was higher in Gen-enriched bioanodes than in FeC-enriched ones (2.9 ± 0.5 and 1.4 ± 0.4 , respectively) [20].

Conclusion

Combined use of DGGE and qPCR biological approaches to study the microbial community growing in anode chambers allowed us to characterize 86% of the freshwater sample and between 64% - 87% of the anode community. This is a higher percentage than found from previous studies [22, 48] and is in line with the results from other biological approaches in MFC investigation [49–51]. The qPCR confirmed sequencing analysis performed on the bands cut from the DGGE gels. This approach proved to be useful to have a quite full and detailed understanding of the dynamic evolution of the anodic microbial communities, without high costs in terms of budget and time [21, 52–54].

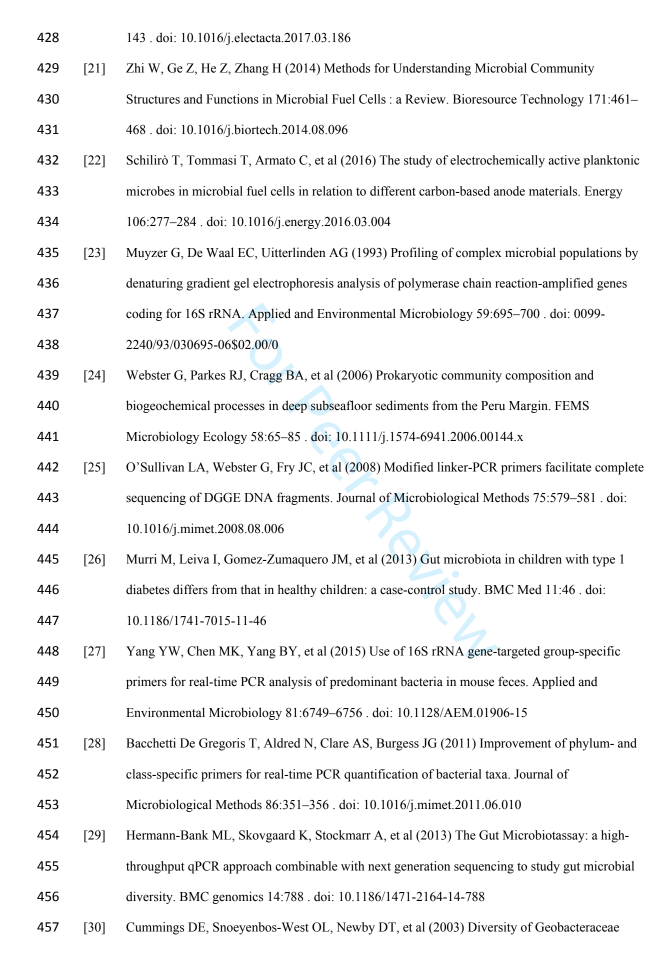
Pre-enrichment steps with FeC strongly affected the equilibrium of the microbial community. However, rather than facilitating the growing of ARB, which were thought to be the most responsible for current production, the procedure just generally reduced the population diversity. On the contrary, the MFCs exposed to Gen enrichment showed a more heterogeneous community and had a better performance than the FeC ones. Our findings suggest that the use of a highly selective method of enrichment seems to be detrimental to the formation of an anode microbial community adequate for operating inside MFCs.

Declaration of interest

The authors declare no conflict of interest.

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547 Tables

Table 1. Targets, primers, amplicon size, annealing temperature and genomic standards used in real time qPCR. Ta= annealing temperature, bp= base pair.

Target	Primers	Amplicon size (bp); Ta	Standard	Reference
Bacteroidetes	F CATGTGGTTTAATTCGATGAT	126	Bacteroides fragilis	[26]
	R AGCTGACGACAACCATGCAG	Ta: 60°C		
β- Proteobacteria	Beta979F AACGCGAAAAACCTTACCTACC	174	Alcaligenes faecalis	[27]
	Beta1130R TGCCCTTTCGTAGCAACTAGTG	Ta: 50°C		
γ- Proteobacteria	1080yF TCGTCAGCTCGTGTYGTGA	170	Shewanella oneidensis	[28]
	Y1202R CGTAAGGGCCATGATG	Ta: 52°C		
δ- Proteobacteria	F: GGTGTAGGAGTGAARTCCGT	534	Geobacter metallireducens	[29]
	R: TACGTGTGTAGCCCTRGRC	Ta: 55°C		
Firmicutes	F: ATGTGGTTTAATTCGAAGCA	126	Clostridium difficile	[26]
	R: AGCTGACGACAACCATGCAC	Ta: 60 °C		
Geobacteraceae spp.	Geo564F AAGCGTTGTTCGGAWTTAT	277	Geobacter metallireducens	[30]
	Geo84OR GGCACTGCAGGGGTCAATA	Ta: 55°C		
Pseudomonas spp.	gacA1 GBATCGGMGGYCTBGARGC	425	Pseudomonas aeruginosa	[31]
	gacA2 MGYCARYTCVACRTCRCTGSTGAT	Ta: 61°C		
Total Bacteria	16S RNA F AGAGTTTGATCMTGGCTCAG	About 600	Desulfovibrio vulgaris	[32]
	16S RNA R TTACCGCGGCKGCTGGCAC	Ta: 55°C		
	Probe CCAKACTCCTACGGGAGGCAGCAG			

Table 2. Thermal protocol of qPCR for all strains under study. Each reaction is 40 cycles long. In the last
 row are the Efficiency (Eff) and R² obtained from each standard curve.

Target	Total bacteria	β- Proteoba cteria	γ- Proteoba cteria	δ- Proteoba cteria	Bacteroid etes	Firmicute s	Geobacteria ceae spp.	Pseudomo nas spp.
Initial denaturation	95°C for 3 min	95°C for 2 min	95°C for 3 min	95°C for 3 min	95°C for 2.5 min	95°C for 2.5 min	95°C for 3 min	95°C for 2.5 min
Denaturation	95°C for 30 s	95°C for 10 s	95°C for 30 s	95°C for 30 s	95°C for 10 s	95°C for 10 s	95°C for 30 s	95°C for 30 s
Annealing	55°C for 30 s	50°C for 30 s	52°C for 30 s	55°C for 30 s	60°C for 20 s	60°C for 20 s	55°C for 30 s	61°C for 30 s
Extension	72°C for 30 s		72°C for 30 s	72°C for 36 s	72°C for 15 s	72°C for 15 s	72°C for 30 s	72°C for 36 s
	_	Melt curve	Melt curve	Melt curve	Melt curve	Melt curve	Melt curve	Melt curve
Standard curve parameters	Eff= 90.8 R ² =0.992	Eff= 99.9 R ² =0.999	Eff= 95.3 R ² =0.991	Eff= 95.4 R ² =0.99	Eff= 97.1 R ² =0.999	Eff= 100.7 R ² =0.999	Eff= 104.0 R ² =0.998	Eff= 99.7 R ² =0.999

Table 3. Sequencing analysis of the band excised from DGGE lanes of inoculum and first and third preculture enrichment steps. Accession number and percentage of similarity are also reported.

Band	Closest relative	Phylum	Class	accession n°	Similarity %
1	Uncultured γ-proteobacterium	Proteobacteria	γ-proteobacteria	HE856452	95
2	Acinetobacter sp.	Proteobacteria	γ-proteobacteria	JN082732	98
3	Uncultured β-proteobacterium	Proteobacteria	β-proteobacteria	KC602997	86
4	Alcaligenes sp.	Proteobacteria	β-proteobacteria	KX345927	93
5	Acinetobacter sp.	Proteobacteria	γ-proteobacteria	KP943116	100
6	Bacterium enrichment culture		nd	FJ842606	98
7	Arcobacter sp.	Proteobacteria	ε-proteobacteria	KP182157	99
8	Uncultured Trichococcus sp.	Firmicutes	Bacilli	KR911832	99
9	Uncultured bacterium		nd	LN651052	96
10	Uncultured bacterium		nd	KU362745	99
11	Uncultured Clostridium sp.	Firmicutes	Clostridia	KU764678	99
12	Trichococcus sp.	Firmicutes	Bacilli	KT424954	93
13	Bacillus sp.	Firmicutes	Bacilli	KJ743291	100
14	Uncultured Agrobacterium sp.	Proteobacteria	α-proteobacteria	JN625545	100
15	Uncultured Comamonas sp.	Proteobacteria	β-proteobacteria	KX010337	100
16	Rhizobium sp.	Proteobacteria	α-proteobacteria	KC252885	99
17	Uncultured Geobacter sp.	Proteobacteria	δ-proteobacteria	LC001501	99
18	Uncultured Clostridiales sp.	Firmicutes	Clostridia	KJ185096	100

Table 4. Sequencing analysis of the band excised from DGGE lanes of planktonic, carbon felt biofilm and graphite rod components at the end of the test. Accession number and percentage of similarity are reported.

Б		D	- CI		Similarity
Band	Closest relative	Phylum	Class	accession n°	%
1	Comamonas sp.	Proteobacteria	β-proteobacteria	KX225279	98
2	Pseudomonas sp.	Proteobacteria	γ-proteobacteria	AY954288	98
3	Uncultured Alcaligenes sp.	Proteobacteria	β-proteobacteria	LC001185	99
4	Uncultured β-proteobacteria	Proteobacteria	β-proteobacteria	CU920026	91
5	Uncultured Comamonas sp.	Proteobacteria	β-proteobacteria	AB793337	100
6	Comamonas sp.	Proteobacteria	β-proteobacteria	KX225279	100
7	Rhizobium sp.	Proteobacteria	α-proteobacteria	JN688942	99
8	Uncultured Arcobacter sp.	Proteobacteria	ε-proteobacteria	JX944559	99
9	Uncultured Alcaligenes sp.	Proteobacteria	β-proteobacteria	LC001185	99
10	Uncultured Alcaligenes sp.	Proteobacteria	β-proteobacteria	LC001185	100
11	Arcobacter sp.	Proteobacteria	ε-proteobacteria	FJ968638	99
12	Uncultured bacterium		nd	LN651026	83
13	Pseudomonas sp.	Proteobacteria	γ-proteobacteria	LN885540	99
14	Arcobacter sp.	Proteobacteria	ε-proteobacteria	FJ968638	99
15	Comamonas sp.	Proteobacteria	β-proteobacteria	KX225279	100
16	Uncultured Geobacter sp.	Proteobacteria	δ-proteobacteria	JX944527	98
17	Acinetobacter sp.	Proteobacteria	γ-proteobacteria	KP943121	99
18	Bacterium		nd	AJ630288	100
19	Comamonas sp.	Proteobacteria	β-proteobacteria	KC622039	96
20	Uncultured Achromobacter sp.	Proteobacteria	β-proteobacteria	LC070644	98
21	Uncultured Microvirgula sp.	Proteobacteria	β-proteobacteria	LC070638	99
22	Comamonas sp.	Proteobacteria	β-proteobacteria	KX225279	100
23	Uncultured γ-proteobacterium	Proteobacteria	γ-proteobacteria	AJ871053	89
24	Comamonas sp.	Proteobacteria	β-proteobacteria	KM083034	100
25	Comamonas sp.	Proteobacteria	β-proteobacteria	KX225279	99
26	Rhizobium sp.	Proteobacteria	α-proteobacteria	JN688942	99

588 589	Figure legends
590	Figure 1. a) DGGE profile and cluster analysis of bacterial community profile of freshwater sediment (I)
591	and the first and third precultures (Gen = green; FeC = red). The trees were generated using Jaccard
592	similarity. b) Shannon Index in the inoculum, Gen and FeC first (Pre1) and third (Pre3) precultures (green
593	and red, respectively).
594	Figure 2. Graphical representation of relative abundance of real time qPCR products in the inoculum (top)
595	and during the first and third preculture steps of General and FeC enrichment (bottom left and bottom
596	right, respectively). 1° and 3° refer to the first and third step of precultures, respectively.
597	Figure 3. Histograms of <i>Geobacteraceae</i> spp. and <i>Pseudomonas</i> spp. real time quantification in the
598	inoculum and during the first (Pre1) and third (Pre3) enrichment steps for the general (Gen) and ferric
599	citrate (FeC) precultures. Geobacteraceae spp.: Inoculum vs FeC enrichment steps $p < 0.001$ (ANOVA,
600	Tukey's post hoc). Inoculum vs third Gen enrichment step $p > 0.05$ (ANOVA, Tukey's post hoc).
601	$\it Pseudomonas \ spp. \ inoculum \ vs \ both \ third \ enrichment \ steps \ p < 0.001 \ (ANOVA, \ Tukey's \ post \ hoc).$
602	Figure 4. a) Cluster analysis of bacterial community profile of anodic plankton (P), biofilm (B) and rod
603	(R) of the Gen-MFC (green) and FeC-MFC (orange). The trees were generated using Jaccard similarity.
604	b) Shannon Index at the end of test in the three different component of MFC anode (P = plankton; B =
605	biofilm; R = rod). Gen-MFC in green and FeC-MFC in red.

Figure 5. Graphical representation of relative abundance of real time qPCR products in the planktonic (P)

and attached (B+R) components at the end of the test in the general (Gen) and ferric citrate (FeC) MFCs.

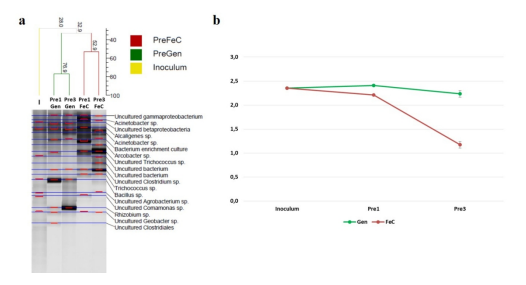


Figure 1. a) DGGE profile and cluster analysis of bacterial community profile of freshwater sediment (I) and the first and third precultures (Gen = green; FeC = red). The trees were generated using Jaccard similarity. b) Shannon Index in the inoculum, Gen and FeC first (Pre1) and third (Pre3) precultures (green and red, respectively).

114x60mm (300 x 300 DPI)

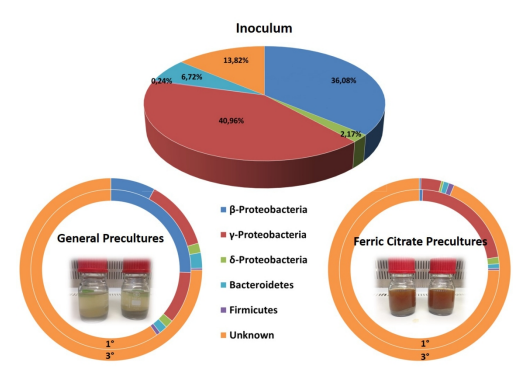


Figure 2. Graphical representation of relative abundance of real time qPCR products in the inoculum (top) and during the first and third preculture steps of General and FeC enrichment (bottom left and bottom right, respectively). 1° and 3° refer to the first and third step of precultures, respectively.

121x86mm (300 x 300 DPI)

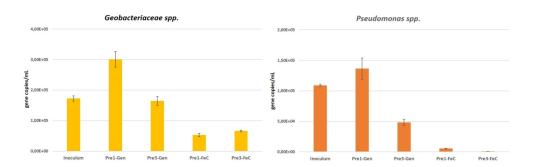


Figure 3. Histograms of Geobacteraceae spp. and Pseudomonas spp. real time quantification in the inoculum and during the first (Pre1) and third (Pre3) enrichment steps for the general (Gen) and ferric citrate (FeC) precultures. Geobacteraceae spp.: Inoculum vs FeC enrichment steps p < 0.001 (ANOVA, Tukey's post hoc). Inoculum vs third Gen enrichment steps p > 0.05 (ANOVA, Tukey's post hoc). Pseudomonas spp. inoculum vs both third enrichment steps p < 0.001 (ANOVA, Tukey's post hoc).

169x52mm (300 x 300 DPI)

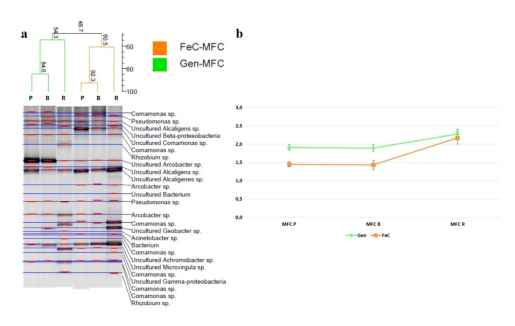


Figure 4. a) Cluster analysis of bacterial community profile of anodic plankton (P), biofilm (B) and rod (R) of the Gen-MFC (green) and FeC-MFC (orange). The trees were generated using Jaccard similarity. b) Shannon Index at the end of test in the three different component of MFC anode (P = plankton; B = biofilm; R = rod). Gen-MFC in green and FeC-MFC in red.

120x71mm (300 x 300 DPI)

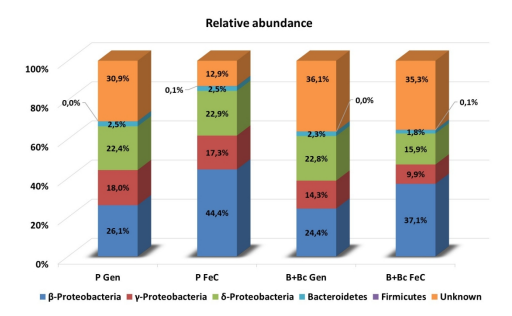
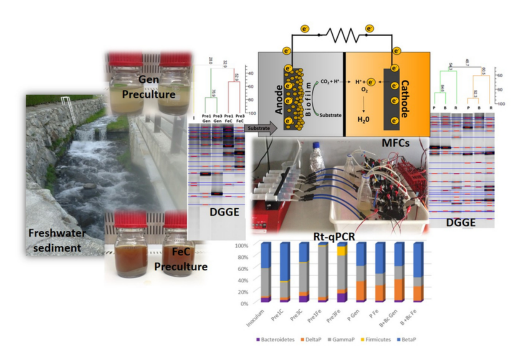


Figure 5. Graphical representation of relative abundance of real time qPCR products in the planktonic (P) and attached (B+R) components at the end of the test in the general (Gen) and ferric citrate (FeC) MFCs.

135x81mm (300 x 300 DPI)



TOC/Graphical Abstract

67x44mm (300 x 300 DPI)