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**Official URL:** [https://doi.org/10.1016/j.bioelechem.2019.03.002](https://doi.org/10.1016/j.bioelechem.2019.03.002)

**To cite this version:**

Champigneux, Pierre and Renault-Sentenac, Cyril and Bourrier, David and Rossi, Carole and Délia, Marie-Line and Bergel, Alain. *Effect of surface roughness, porosity and roughened micro-pillar structures on the early formation of microbial anodes*. (2019) Bioelectrochemistry, 128. 17-29. ISSN 1567-5394

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Effect of surface roughness, porosity and roughened micro-pillar structures on the early formation of microbial anodes

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Abstract

The early formation of electroactive biofilms was investigated with gold electrodes inoculated with Geobacter sulfurreducens. Biofilms were formed under an applied potential of 0.1 V/SCE, with a single batch of acetate 10 mM, on flat gold electrodes with different random surface roughness. Roughness with arithmetical mean height (Sa) ranging from 0.5 to 6.7 μm decreased the initial latency time, and increased the current density by a factor of 2.7 to 6.7 with respect to nano-rough electrodes (Sa = 4.5 nm). The current density increased linearly with Sa up to 14.0 A·m⁻² for Sa of 6.7 μm. This linear relationship remained valid for porous gold. In this case, the biofilm rapidly formed a uniform layer over the pores, so porosity impacted the current only by modifying the roughness of the upper surface. The current density thus reached 14.8 ± 1.1 A·m⁻² with Sa of 7.6 μm (7 times higher than the nano-rough electrodes). Arrays of 500-μm-high micro-pillars were roughened following the same protocol. In this case, roughening resulted in a modest gain around 1.3-fold. A numerical model showed that the modest enhancement was due to ion transport not being sufficient to mitigate the local acidification of the structure bottom.

Keywords:
Bioanode
Geobacter sulfurreducens
Gold
Microbial fuel cell
Bioelectrochemical system

1. Introduction

Over the past fifteen years, the electroactive properties of microorganisms have motivated a profusion of inspiring work. In the domain of microbial electrochemical technologies [1,2], microbial anodes are the core element of many innovative processes [3], such as microbial fuel cells [4], microbial electrolysis cells for hydrogen production [5], methane synthesis [6] or metal recovery [7], passive systems such as the microbial snorkel for environmental bioremediation [8], various kinds of biosensors [9,10], etc.

Beyond the framework of microbial electrochemical technologies, electroactive biofilms are at work in microbial corrosion [11], and microbial anodes have been envisioned as a possible way to inhibit corrosion [12]. Electroactive species have also been used as markers of surface biofouling [13,14]. The early detection of electroactivity has successfully been implemented for monitoring equipment biofouling in marine and fresh water environments [15]. In this case, the sensors were based on cathodic catalysis but, given the ubiquity of exoelectrogenic species [16,17], there is no doubt that anodic sensors will also be developed in the future to investigate or monitor environmental biofilms.

In this very broad context, the research community is still facing an important gap in basic knowledge when the objective is to control the electrode/bacteria interface. The predominant role of electrode topography has been pointed out in many studies [18–26] and in a recent review article [27], but controversial results have been reported. For instance, surface roughness has sometimes been reported as the main parameter that impacts the current density produced by microbial anodes, being even more important than the anode material itself [28]. In contrast, in the case of stainless steel electrodes, 5 μm roughness has shown no positive effect vs. a smooth surface [22]. In another study, stainless steel electrodes inoculated with Geobacter sulfurreducens showed increased biofilm coverage and current production when the average surface roughness was increased from 2 to 4 μm, while no effect was observed at lower values, from 0.1 to 2 μm [23]. A third study, which used flame oxidation to increase the roughness of stainless steel electrodes from 75 to 151 nm, reported higher current and better biofilm coverage with higher roughness [29].

These examples illustrate the extreme variability of the effect of roughness reported in the literature. This situation is, to a great extent, due to the concomitant variation of roughness and other surface parameters. Variation of roughness is often linked to the variation in surface chemistry, which depends on the technique used to modify the roughness. It is consequently difficult to unravel the impact of the surface topography from that of other surface parameters, and notably those
related to chemical modification of the surface [29]. For these reasons, several research teams have turned to surface structuration as a way to control the surface topography by minimizing the changes of other surface properties [30–34].

We have recently implemented surface structuring of gold electrodes to compare the nano-rough surface (Rq = 4.5 nm), either flat or structured with 500-μm-high micro-pillars [34]. The current density produced was increased from 2 to 8.5 A·m⁻² by the micro-pillars. This current increase was attributed to the larger surface area available for biofilm development, due to the micro-pillars.

Here, the same experimental approach was implemented in order to assess the impact of random roughness at the micrometre level. As previously, gold electrodes were used because gold is a suitable electrode material for forming efficient microbial anodes [35–38] and it allows accurate control of the surface topography, while keeping similar surface chemical properties. In is thus possible to focus on the impact of the electrode topography by avoiding variation of other surface parameters. As previously, electrodes were colonized by Geobacter sulfurreducens, a widely-used model electroactive bacterium [39].

Attention was focused on the early stage of biofilm formation. Early formation is the phase when there are the most opportunities to act on the biofilm properties. It is consequently important to progress in understanding this key phase and to identify the parameters that can impact it, if the objective is to establish basic guidelines for electroactive biofilm design. The early stage of biofilm formation is the phase where levers are most likely to be found, not only to enhance electroactive properties, but also to mitigate them, as in the context of microbial corrosion or biofouling. Mastering the early formation of electroactive biofilms is also essential to decrease the response time and the detection threshold of sensors based on the detection of microbial electroactivity.

The previous study dealt with surface structuring in the form of 500-μm-high micro-pillars [34]. The present study completes the range of surface topographies in three ways:

- random roughness ranging from 0.5 to 6.7 μm was elaborated on flat electrodes by using gold electrodeposition,
- the same gold electrodeposition protocol was pushed to create a thick porous layer with pore sizes from a few to around 50 μm,
- the random roughness was elaborated on the electrode surface that had previously been structured with the 500-μm-high micro-pillars.
- The sum of the two studies gives a comprehensive view of the impact of surface topography over a very wide range.

Finally, it would be useful to emphasize a general comment on the results provided by the present work. It is shown here that random micrometre-sized surface roughness has a great impact on the current produced by microbial anodes. This roughness size can be achieved unintentionally when treating, modifying or coating an electrode surface. Consequently, micrometre-sized roughness should now be considered as a key parameter in the assessment of any electrode surface modification. When comparing the performance of a non-treated, control electrode with a treated, modified or coated electrode, the possible variation of the surface roughness must be assessed first, before concluding on the possible impact of other parameters related to the interface modification. Beyond the specific results described below, this general warning should be one of the major conclusions of the study.

2. Materials and methods

2.1. Preparation of gold electrode with different surface roughness values

Nano-rough gold (NG) surfaces 1 cm² in area were formed on Si (100), P-type silicon wafers. The electrodes were formed by evaporation of a 250 nm thick gold layer followed by electrolytic deposition of gold to achieve a 1.75 μm thick gold layer. The average roughness, measured by AFM, was 4.5 nm for the NG surfaces.

Random micro-rough gold (MG) was formed by electrolytic growth of an additional gold layer on the NG surfaces. The NG surfaces were immersed in a bath of chloroauric acid (H[AuCl₄], 2 mM) and sulfuric acid (H₂SO₄, 96%, 2 mM). Four different current densities were applied: 0.35, 0.69, 1.39 and 2.08 A·m⁻², which led to 4 different surface topographies named MG-1 to MG-4. The MG surfaces were then mounted on polychlorobiphenyl (PCB) supports. The gold surface was connected to the contact pad of the PCB using a KnS 4526 micro-welding machine and electrical contacts were insulated with biocompatible silicon (Locite 4562).

Porous gold (PG) was formed following the same protocol at a high rate using 5 A·m⁻² for 10 min. The porous structure obtained was around 100 μm thick, measured with a TENCOR P16 mechanical profilometer. Three similar electrodes were made in this way.

Micro-pillar arrays with micro-rough surface (μP-MG): surface micro-structuring was performed on Si (100) P-type silicon wafers with an epoxy resin (SU-8 3050, MicroChem Corp.) patterned by photolithography to form the pillar array. The array consisted of square pillars 100 μm wide and 500 μm high, spaced 200 μm apart. The structured electrodes had 1156 pillars on a 1 cm² surface. A first metallization was performed by sputtering Cu/Cr and then smooth electrolytic deposition of gold to create the usual NG gold layer with 4.5 nm roughness. Then the surface of the micro-pillars was roughened by electrolytic deposition of gold, performed for 10 min at three different current densities: two electrodes were roughened by electrodeposition at 0.8 A·m⁻² (μP-MG-1), one at 1.5 A·m⁻² (μP-MG-2) and another at 2.1 A·m⁻² (μP-MG-3).

2.2. Electrochemical set-up

Colonization of the gold surfaces was carried out in tightly sealed 500 mL reactors, filled with 300 mL solution and continuously sparged with N₂:CO₂ (80:20). The gas bubbling was carefully arranged to avoid any disturbance on the working electrode. A water-bath kept the temperature steady at 30 °C. The experiments were performed with 3-electrode set-ups in single compartment reactors. The auxiliary electrode was a platinum grid (Heraeus SAS, Germany) and a saturated calomel electrode was used as the reference (SCE, potential +0.24 V/SHE). The gold surfaces were mounted and welded onto a PCB support and connected to the potentiostat. The bioanodes were formed under constant polarization at 0.1 V/SCE (VSP potentiostat, Bio-Logic SA, France) and the current densities were recorded throughout the experiment every 10 min. Cyclic voltammetries at 1 mV·s⁻¹ were performed before inoculation and when the current was close to reaching a maximum.

The electroactive biofilms were formed under constant polarization at 0.1 V/SCE in 4 different runs. Each run corresponded to four or five reactors run in parallel and inoculated with the same pre-culture of Geobacter sulfurreducens:

- run #1: 4 reactors with the MG-1 to MG-4 micro-rough surfaces and 1 reactor with the nano-rough (NG) surface used as a control, 
- run #2: identical to run #1, 
- run #3: 3 reactors with the PG surface and 2 reactors with the NG surface used as a control, 
- run #4: 4 reactors with micro-rough micro-pillar arrays, μP-MG-1 to μP-MG-3.

The current densities were expressed with respect to the 1 cm² projected surface area of the gold surfaces. The chronoamperograms (current density vs. time) were characterized by three parameters: the maximum current density reached (Jmax A·m⁻²), the starting time (tstart days), defined as the time required for the current to reach 10% of Jmax, and the current density increase rate (r, A·m⁻²·d⁻¹), which was calculated as the average value of the slope of the current density vs. time over 1 day around the inflection point of the curve. Maximum current densities were recorded by checking that the
maximum value was reached during at least 18 records during the chronoamperogramm, which represented a duration of at least 3 h (record every 10 min.). According to the 10 min frequency current recording, the experimental error of the starting time values was ±0.01 d.

2.3. Inocula and media

*Geobacter sulfurreducens* (ATCC 51573) purchased from DSMZ was grown in specific growth medium that contained, per litre: 0.1 g KCl, 1.5 g NH₄Cl, 2.5 g NaHCO₃, 0.6 g NaH₂PO₄, and 0.82 g CH₃COONa. The medium was sterilized by autoclaving at 121 °C for 20 min. After sterilization, the medium was completed with 8 g/L sodium fumarate filtered to 0.22 µm, 10 mL/L Wolfe’s vitamin solution (ATCC MD-VS) and 10 mL/L modified Wolfe’s minerals (ATCC MD-TMS). A pre-culture of *G. sulfurreducens* was prepared for 3 days in a fresh deoxygenated culture medium (2% v/v, 30 °C) to reach a final absorbance of around 0.4 at 620 nm. Reactors were filled with growth medium, vitamin solution and modified Wolfe’s minerals and constantly sparged with N₂/CO₂ (80:20). After 20 min of deoxygenation under open circuit potential, the electrodes were characterized by an initial cyclic voltammetry. Then, reactors were inoculated with the pre-culture. The inoculation volume was adjusted to obtain the cell density in each reactor that corresponded to an inoculation ratio of 10% v/v with an inoculum at 0.4 absorbance. After the inoculation, acetate 10 mM was added to the medium as the sole electron donor; no fumarate was present.

2.4. Microscopic imaging and electrode characterization

The clean electrode surfaces were observed on a Hitachi S-4800 scanning electron microscope (SEM) in low vacuum mode at an accelerating voltage of 15 kV and an intensity of 15 µA. Various magnifications and locations were used to characterize the surfaces. The surface roughness was characterized by the spatial arithmetical mean height (Sₐ) measured with a non-contact 3D optical profiler (S-Neox, Sensofar). The values were averaged over 20% of each surface.

The electrochemically active surface area (Aₑ) was measured following the technique described by Burke and Nugent [40]. The electrodes were first soaked in an H₂O₂/H₂SO₄ (1:3) solution to reduce the hydrophobicity and thus avoid gas retention on the surface during the measurement. Cyclic voltammetry was performed between 0 and 1.8 V/SCE in H₂SO₄ 0.5 M, following two successive phases: firstly, 25 cycles were performed at 500 mV·s⁻¹ in order to clean the surface, then 25 cycles were performed at 50 mV·s⁻¹ for the measurement. A reduction peak appeared between 0.8 and 0.9 V/SCE, which was assumed to be due to the formation of a monolayer of gold oxide. The charge related to the peak (Q, µC) was calculated by integration and was linked to the electrode surface area (Aₑ, cm²) with the equation:

\[ Aₑ = \frac{Q}{CF} \]  

where CF = 386 µC·cm⁻² is the conversion factor given by Burke and Nugent for gold [40].

The technique was firstly used as proposed by Burke and Nugent with the NG electrodes. Eq. (1) gave a surface area of 2.73 cm². This value was clearly overestimated as the projected surface area was only 1 cm². It was extremely unlikely that the surface area would be multiplied by a factor of 2.73 with a surface roughness as low as 4.5 nm. It was speculated that the electrodeposition technique used here to prepare the electrodes led to the formation of more than one molecular layer of oxide, while the Burke and Nugent procedure assumes that plain gold is covered by only a monolayer. Consequently, we chose to take the nano-rough electrode as the reference, by taking its electrochemically active surface area to be equal to its projected surface area of 1 cm². Accordingly, Eq. (1) was recalibrated by using a conversion factor of 1053 µC·cm⁻².

At the end of the experiments, the bioanodes were stained with Syto 9 (Invitrogen, Molecular Probes) at 2.5 µM for 10 min, then carefully washed with medium and dried at ambient temperature. The electrodes were imaged with a Carl Zeiss Axiovalger M2 microscope equipped for epifluorescence with an HBO 50 W ac mercury light source and the Zeiss 09 filter (excitor HI450–490, reflector FT 10, Barrier filter LP520). Images were acquired with a monochrome digital camera (Evolution VF). Several different spots were imaged in order to have an appropriate representation of the general pattern of the colonization.

Epifluorescence images were analysed by greyscale interpretation to calculate the ratio of the electrode surface area covered by bacteria. The grey intensity threshold between the areas covered by the bacteria and the non-covered areas was set manually. Grey levels greater than the threshold value were considered to correspond to colonized areas, while lower grey levels were considered to show clean areas. Six images, of dimensions 866 by 645 µm, were treated for each electrode to determine the average coverage ratio, the edges of the electrodes being excluded to avoid the impact of side effects.

The biofilm structures were observed by scanning electron microscopy (SEM) with a Leo 435 VP-Carl Zeiss SMT. A 4% glutaraldehyde phosphate buffer (400 mM, pH 7.4) was used to fix the biofilms. The surfaces were then rinsed in phosphate buffer containing saccharose (400 mM). The dehydration process consisted of immersion in successive concentrations of acetone (50%, 70%, 100% solutions), followed by acetone and hexamethyldisilazane (HMDS) solution (50:50), and finally 100% hexamethyldisilazane. Complete evaporation of the last solution was achieved to complete the process. The samples were sputtered with a thin (10 nm) layer of gold (Sancoat Six SEM sputter coater) to guarantee their electrical conductivity.

3. Results and discussion

3.1. Electrode topography of micro-rough (MG) and porous (PG) flat surfaces

SEM imaging of the MG surfaces formed at 0.35 and 0.69 A·m⁻² (MG-1 and MG-2) showed randomly distributed deposits that covered around 50 and 70% of the surface, respectively (Fig. 1). The higher electrodeposition current densities of 1.39 and 2.08 A·m⁻² led to round craters about 10 µm in diameter (MG-3 and MG-4). During the electrodeposition phase, gas bubbles evolved from the electrode surface. At the lower current densities (MG-1 and MG-2), electrodeposition was slow and the bubbles detached from the surface without impacting the deposition. For MG-3 and MG-4, gold deposition was faster and tended to entrap the bubbles in the deposited structure. This phenomenon created a crater pattern with characteristic size of around 10 µm.

Higher magnification showed that gold formed flr-shaped deposits of nano-dendrites on the MG-1 and MG-2 electrodes. These nano-dendrites tended to disappear on the MG-3 and MG-4, probably due to the visible densification of deposition.

All these MG surfaces showed roughness patterns added on to the NG basal surface, which remained visible at the bottom. In contrast, the PG surfaces formed at 5 A·m⁻² showed a porous structure uniformly deposited over the whole surface area, so the basal NG surface was no longer visible (Fig. 2A). Deep pores were observed, with pore sizes ranging from a few microns to around 50 µm (Fig. 2B). The largest pores were interconnected, creating a highly porous network. The MG surfaces could be considered as 2-dimensional rough surfaces, while the PG surfaces presented a 3-dimensional porous layer with well-formed pores, which fully masked the basal surface.

For the MG surfaces, the surface area measured electrochemically (Aₑ) increased with the electrodeposition current density but the increase was lower at the highest current densities, when deposition became denser (Table 1). This trend was consistent with SEM images,
revealing a deposition pattern of isolated clusters of nano-dendrites at low current density, which tended to disappear with the densification of the deposition at higher current densities.

The arithmetical mean height ($S_a$) of the MG surfaces increased linearly from 0.5 to 6.7 $\mu$m with the electrodeposition current density, while this linearity was clearly broken for the PG surfaces (Fig. 3). This confirmed the visual difference observed by SEM between MG and PG surfaces. Higher current density made the gold layer develop a 3-dimensional porous structure but did not considerably increase the surface roughness.

![SEM imaging of the micro-rough gold (MG) surfaces produced by electrodeposition with increasing current densities from MG-1 to MG4: A) MG-1, $S_a = 0.5 \mu$m; B) MG-2, $S_a = 1.7 \mu$m; C) MG-3, $S_a = 4.6 \mu$m; D) MG-4, $S_a = 6.7 \mu$m.](image1)

![SEM imaging of the porous gold (PG) electrodes.](image2)
3.2. Electrode topography of micro-roughened micro-pillar arrays (\(\mu\text{P-MG}\))

The micro-pillar arrays with smooth surfaces implemented in a previous work [34] were used again here and, in addition, the walls were roughened by gold electrodeposition at 3 different current densities (Table 1).

The rough surface structure was not uniform along the micro-pillars. The top parts of the pillars were very notably roughened, whereas the bases presented low gold deposition. It was not possible to determine roughness parameters on such structures and the surface characterization consequently consisted of an extended visual description based on SEM imaging (Table 2). At the bottom of \(\mu\text{P-MG-1}\), the surface was slightly modified, with roughness evaluated visually at under 1 \(\mu\text{m}\), whereas on \(\mu\text{P-MG-2}\) and \(\mu\text{P-MG-3}\), the roughness was greater, reaching a few micrometres, but did not present any three-dimensional organization. Going up along the pillars, the roughness increased to reach a maximum on the top.

The deposit was thickest on the top edges of the pillars and three different deposition patterns could be distinguished (Fig. 4). On the \(\mu\text{P-MG-1}\) surfaces, obtained with the lower electrodeposition current densities, the deposit presented a thickness of under 10 \(\mu\text{m}\) (Fig. 4A and B). The initiation of a porous structure could be guessed but remained low and mainly composed of 10-\(\mu\text{m}\) wide craters. In contrast, the \(\mu\text{P-MG-2}\) surface, which was roughened at higher current density, showed a 20- to 30-\(\mu\text{m}\) thick deposit consisting of a loose porous structure with pore size around 10 \(\mu\text{m}\) (Fig. 4C). This micro-porous surface extended 50 to 100 \(\mu\text{m}\) down the pillars, decreasing gradually. The \(\mu\text{P-MG-3}\) surface, treated at the highest current density, displayed surface morphology similar to that of \(\mu\text{P-MG-2}\) but with a tighter porous structure (Fig. 4D). The total deposit height was similar but the pore size and the void degree were clearly smaller.

The electrochemically active surface area \(A_e\) (Table 2) was consistent with the SEM images. The loose porous structure of the \(\mu\text{P-MG-2}\) electrode resulted in the highest active surface area, of 98 \(\text{cm}^2\). The active surface area was slightly lower for \(\mu\text{P-MG-3}\) (89 \(\text{cm}^2\)), which is consistent with the tight porosity observed by SEM. The two \(\mu\text{P-MG-1}\) electrodes gave a fairly reproducible \(A_e\) of 48 and 53 \(\text{cm}^2\), respectively. In comparison, the same micro-pillar networks, when implemented with a nano-rough surface of 4.5 nm in a previous work [34], led to considerably lower \(A_e\) of 3.26 \(\text{cm}^2\). The roughening technique used here was efficient in increasing the electrochemically active surface area, ensuring multiplicative factors of 15 to 30.

### Table 1

<table>
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<th>Anode</th>
<th>(J_{\text{depositing}}) (A-m(^{-2}))</th>
<th>(S_a) ((\mu\text{m}))</th>
<th>Roughness morphology</th>
<th>(A_e) ((\text{cm}^2))</th>
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<td>None</td>
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</tr>
<tr>
<td>MG-2</td>
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<td>1.7</td>
<td>Fir-shaped nano-dendrites</td>
<td>18.2</td>
</tr>
<tr>
<td>MG-3</td>
<td>1.39</td>
<td>4.6</td>
<td>Micro-craters</td>
<td>36.8</td>
</tr>
<tr>
<td>MG-4</td>
<td>2.08</td>
<td>6.7</td>
<td>Micro-craters</td>
<td>40.7</td>
</tr>
<tr>
<td>PG</td>
<td>5.0</td>
<td>7.6</td>
<td>Porous 3D layer</td>
<td>--</td>
</tr>
</tbody>
</table>

3.3. Effect of micro-roughness on electroactivity establishment and biofilm growth

The four nano-rough (NG) electrodes, used as controls, showed reproducible values (Table 3) of starting time (2.6 ± 0.3 d) and \(J_{\text{max}}\) (2.1 ± 0.5 A·m\(^{-2}\) on 3 values). The \(J_{\text{max}}\) values were consistent with the average value of 2.5 ± 2.0 A·m\(^{-2}\) obtained in a previous study performed with 7 similar NG electrodes [34]. Here the experimental deviation was the most pronounced on the rate of increase of current density \(r_i\), which ranged from 0.42 to 1.8 A·m\(^{-2}\)·d\(^{-1}\). The large experimental deviation already observed with NG electrodes has been explained by the poor reproducibility of the surface area colonization ratio. Here, epifluorescence imaging performed after 15 days of polarization, when the current had fallen to zero, showed a similar situation. NG electrode surfaces mainly displayed isolated cells and isolated micro-colonies of different sizes, as described previously [34]. The biofilms appeared at an early stage of formation, with micro-colonies, which started to form larger patches at a few places. The NG electrodes consequently showed scattered values of biofilm coverage ratios, ranging from 26 to 57%.

The starting times of the eight MG electrodes (Table 3) were clearly shorter (1.6 ± 0.1 d) than those of the NG electrodes. The presence of micro-roughness reduced the starting time by around 1 day with respect to that recorded with the NG electrodes. As observed on the clean surfaces (Fig. 1), increased surface roughness created asperities of the size of Geobacter cells or larger, thus offering a larger contact area for bacteria to adhere to. In contrast, the lack of anchoring points on the NG surface was detrimental to bacterial adhesion and delayed the current production.

The MG electrodes showed current increase rates between 8 and 12 times higher than the NG electrodes, with \(r_i\) values of 3.8 to 7.6 A·m\(^{-2}\)·d\(^{-1}\) obtained between days 1.5 and 3 (Fig. 5A). The current increase rates rose with the micro-roughness. The roughness and, probably, the presence of cavities tens of microns wide, accelerated biofilm formation.

The maximum current densities \(J_{\text{max}}\) recorded with the MG electrodes, ranging from 5.6 to 14.0 A·m\(^{-2}\), were significantly higher than those of the NG electrodes (2.1 ± 0.5 A·m\(^{-2}\) on average). Actually, after day 3.5, the current continued to increase slowly in run #1, while it tended to stabilize in run #2, which was disturbed by an accidental ingress of air into the gas supply network. The \(J_{\text{max}}\) values from run #2 were consequently underestimated (Table 3). In spite of this technical issue, all the records showed that the MG electrodes produced considerably higher current than the NG electrodes, with a multiplication factor of 2.7 to 6.7. The presence of micro-roughness clearly enhanced the current production.

The \(J_{\text{max}}\) values increased linearly with \(S_a\) parameter values from 0.5 to 6.7 \(\mu\text{m}\), with a slope of 0.80 A·m\(^{-2}\)·\(\mu\text{m}^{-1}\) (Fig. 6). Only the \(J_{\text{max}}\) values obtained from run #1 were used for this correlation (Table 3). Nevertheless, even though the values from run #2 were underestimated, they followed the same linear relationship with respect to \(S_a\), with a slope of 0.82 A·m\(^{-2}\)·\(\mu\text{m}^{-1}\) (Fig. S1 in Supplementary data). This linear relationship was not valid for the NG electrodes, which showed that the presence of micro-roughness caused an actual jump in the current produced.
Cyclic voltammograms were performed after the electrodes reached \( J_{\text{max}} \), on day 6. For comparison the theoretical curve, corresponding to the Nernst–Michaelis–Menten kinetics \([41,42]\), was added:

\[
J = J_{\text{max}} \frac{1}{1 + \exp\left(\frac{-F}{RT} (E-E_k)\right)}
\]

where \( F \) is the Faraday constant (96,485 C per mole e\(^{-}\)), \( R \) is the gas constant (8.3145 J mol\(^{-1}\) K\(^{-1}\)), \( T \) is the temperature (303 K), \( E \) is the applied potential (V) and \( E_k \) is the value of the potential extracted from the experimental curves at which \( J = J_{\text{max}}/2 \).

Fig. 5B shows that all the electrodes were efficient at oxidizing acetate from low potentials, with kinetics not far from the Nernst kinetics. The rate of electron transfer from the bacterial cell to the gold surface was consequently not far below the rate necessary to ensure the Nernst equilibrium at the electrode surface. The normalized presentation of all the CV curves in the form of \( J/J_{\text{max}} \) used in Fig. 5B points out that the equilibrium at the electrode surface. The normalized presentation of all the curves closer to the theoretical Nernst curve. Increasing roughness may thus lead to slightly less efficient electron transfer kinetics.

Furthermore, Fig. 2B also shows that the NG electrode seemed to provide slightly more efficient electron transfer rates, displaying CV curves closer to the theoretical Nernst kinetics. The NG electrodes offered a large, smooth surface area on which the cell could settle completely, which probably promoted electron transfer. The NG electrodes thus provided conditions tending to have opposite effects: on the one hand, their smoothness was detrimental to cell anchoring and biofilm formation but, on the other, the large contact surface they offered to the cells might benefit the electron transfer rate. On smooth surfaces, cell adhesion is difficult and fragile but electron transfer may be promoted. Nevertheless, considering the small difference in the curves, to be fully confirmed, this speculative comment would need to be supported by a systematic study of the electrode kinetics.

The MG electrode images after 15 days of polarization showed greater colonization then the NG controls, with biofilm coverage ratios between 80 and 100% on all electrodes. The average values were \( 92 \pm 7\% \), \( 87 \pm 4\% \), \( 94 \pm 3\% \) and \( 91 \pm 7\% \) on electrodes MG-1 to 4, respectively (Fig. 7).

Two different biofilm structures were observed. For the lowest roughness (MG-1 and MG-2), the biofilm coverage was uniform. The biofilm presented a high coverage ratio and great uniformity, while it did not look dense or thick. The biofilms on the MG-3 and MG-4 electrodes appeared denser and less uniform, showing a thickening in some areas and growing structures up to about ten micrometres high, with some mushroom-like growths. On these electrodes, craters of about 10 \( \mu \)m were superimposed on the micrometre-sized roughness (Fig. 1). This larger scale pattern may have been responsible for the less uniform biofilm structure. The superimposed crater pattern provided a good environment for bacterial adhesion. As reported in many studies, the presence of cavities favours biofilm formation because it creates areas where shear forces are low and reduces the strength needed for primary attachment of cells to the surface \([43–45]\). Furthermore, bacterial communities communicate via quorum sensing systems to develop specific biofilm architectures. Biofilm shaping is a metabolic response that is triggered by the high concentration of signalling molecules. Here, the 10-\( \mu \)m-sized craters may have enhanced the local accumulation of quorum sensing molecules, which favoured biofilm growth and maturation. The presence of mushroom structures may thus have been the result of the more rapid biofilm maturation promoted inside the craters.

**Table 2**

<table>
<thead>
<tr>
<th>( J_{\text{depositing}} ) (A·m(^{-2}))</th>
<th>Bottom roughness (( \mu )m)</th>
<th>Aspect of the surface of the top of pillars</th>
<th>( A_e )(cm(^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \mu P - MG - 1 )</td>
<td>0.8</td>
<td>( \approx ) 1</td>
<td>5–10 ( \mu )m cavities</td>
</tr>
<tr>
<td>( \mu P - MG - 1 ) (duplicate)</td>
<td>0.8</td>
<td>( \approx ) 1</td>
<td>5–10 ( \mu )m cavities</td>
</tr>
<tr>
<td>( \mu P - MG - 2 )</td>
<td>1.5</td>
<td>3–4</td>
<td>Loose porous structure</td>
</tr>
<tr>
<td>( \mu P - MG - 3 )</td>
<td>2.1</td>
<td>3–4</td>
<td>30 ( \mu )m thick with 10 ( \mu )m pores</td>
</tr>
</tbody>
</table>

Fig. 4. SEM imaging of the top of the \( \mu P - MG \) surfaces: (A and B) \( \mu P - MG - 1 \) duplicates; (C) \( \mu P - MG - 2 \); (D) \( \mu P - MG - 3 \).
Table 3
Electrochemical parameters measured on the chronoamperograms as described in Section 2.2: maximum current density ($J_{\text{max}}$, A·m$^{-2}$), the starting time ($t_{\text{start}}$, days), defined as the time required for the current to reach 10% of $J_{\text{max}}$, and the current density increase rate ($r_i$, A·m$^{-2}$·d$^{-1}$), calculated as the average value of the slope of the current density vs. time over 1 day around the inflection point of the curve.

<table>
<thead>
<tr>
<th>Surface</th>
<th>$t_{\text{start}}$ (d)</th>
<th>$r_i$ (A·m$^{-2}$·d$^{-1}$)</th>
<th>$J_{\text{max}}$ (A·m$^{-2}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NG</td>
<td>2.4</td>
<td>0.52</td>
<td>1.7</td>
</tr>
<tr>
<td></td>
<td>2.8</td>
<td>0.42</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td>2.4</td>
<td>1.8</td>
<td>2.1</td>
</tr>
<tr>
<td></td>
<td>2.9</td>
<td>1.4</td>
<td>2.6</td>
</tr>
<tr>
<td>MG-1</td>
<td>1.5</td>
<td>3.8</td>
<td>9.0</td>
</tr>
<tr>
<td></td>
<td>1.7</td>
<td>3.9</td>
<td>6.9</td>
</tr>
<tr>
<td>MG-2</td>
<td>1.6</td>
<td>4.6</td>
<td>10.3</td>
</tr>
<tr>
<td></td>
<td>1.6</td>
<td>4.6</td>
<td>7.0</td>
</tr>
<tr>
<td>MG-3</td>
<td>1.5</td>
<td>6.2</td>
<td>13.2</td>
</tr>
<tr>
<td></td>
<td>1.6</td>
<td>5.0</td>
<td>8.6</td>
</tr>
<tr>
<td>MG-4</td>
<td>1.5</td>
<td>7.6</td>
<td>14.0</td>
</tr>
<tr>
<td></td>
<td>1.8</td>
<td>7.0</td>
<td>11.3</td>
</tr>
<tr>
<td>PG</td>
<td>2.7</td>
<td>7.6</td>
<td>15.9</td>
</tr>
<tr>
<td></td>
<td>2.7</td>
<td>7.9</td>
<td>14.7</td>
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<tr>
<td></td>
<td>2.8</td>
<td>7.3</td>
<td>13.8</td>
</tr>
<tr>
<td>μ-P-MG-1</td>
<td>0.9</td>
<td>1.4</td>
<td>5.9</td>
</tr>
<tr>
<td>μ-P-MG-1</td>
<td>0.9</td>
<td>1.4</td>
<td>5.8</td>
</tr>
<tr>
<td>μ-P-MG-2</td>
<td>1.0</td>
<td>2.1</td>
<td>7.8</td>
</tr>
<tr>
<td>μ-P-MG-3</td>
<td>0.9</td>
<td>1.9</td>
<td>7.0</td>
</tr>
</tbody>
</table>

* The values of $J_{\text{max}}$ from run #2 (see Section 2.2) were underestimated because of air ingress into the gas network supply.

3.4. Effect of porosity on electroactivity establishment and biofilm growth

The porous gold electrodes (PG) showed reproducible behaviour (Fig. S2A in Supplementary Data) for the phase of current establishment. Cyclic voltammetries performed at day 5, when the bioanodes were close to reaching maximum current production, confirmed the great reproducibility of the three bioanodes (Fig. S2B in Supplementary Data) and also their excellent kinetics, close to the theoretical Nernst-Michaelis-Menten kinetics.

The PG electrodes showed $r_i$ and $J_{\text{max}}$ values comparable to those of the MG-4 electrodes (Table 3). The rates of current density increase were identical to those of the MG-4 electrodes, the maximum current densities were only slightly improved and the start-up times were longer for the porous electrodes. The thick porous layer was consequently not markedly more efficient than micro-roughness. Starting times were longer and maximum current densities were only slightly improved.

The spatial arithmetic mean height ($S_a$) was 6.7 μm for the MG-4 electrodes and 7.6 μm for the porous electrodes (Table 1). The linear relationship between $J_{\text{max}}$ and $S_a$ observed for the MG electrodes was fully validated with the PG electrodes (Fig. 6). Actually, the 100-μm thick porous structure behaved like a 2-dimensional micro-rough surface. This conclusion is consistent with pore clogging, which was observed on most of the electrode surface area.

The reactors were stopped at day 18 to image the biofilms. Epifluorescence microscopy showed a dense, uniform biofilm, with a thickness of around 20–30 μm. In some areas, the biofilm was punctured above the pores (Fig. 8A) but, in most spots, it completely covered the pores (Fig. 8B). This resulted in an almost uniform biofilm with a bumpy appearance as seen on the 3-dimensional reconstruction (Fig. 8C).

SEM imaging of the third PG electrode confirmed that the biofilm covered the majority of the surface area with only some rare spots where unclogged pores appeared. SEM imaging also showed that the walls of the open pores were often lined with a continuous biofilm going as deep as could be observed (Fig. 9A). During the drying process, in a few spots, a layer around 50 μm thick of the porous deposited gold cracked and pulled away. This allowed observations of pores in depth, below the top surface. Hence, it was observed that the pores clogged by the 20-μm thick biofilm over their opening had been deeply colonized, similarly to the unclogged pores (Fig. 9B). A biofilm around 5 to 10 cells thick was observed on most of the pore walls. This internal colonization of the pores could be observed as deep as the porous structure was cracked, i.e., around 50 μm. Finally, when zooming in at the cell scale, we observed gold rods of tens of nanometres tightly interdigitated with Geobacter cells (Fig. 9C), which suggests that the small scale roughness was fully suitable for entrapping cells. The gold deposit made here appeared highly suitable for bacterial colonization, leading to tight interdigitation between gold structures and cells.

3.5. Effect of roughened micro-pillars on electroactivity establishment and biofilm growth

One way to avoid clogging of a porous structure by a biofilm covering the top was to design larger pores or a structure with more space...
between protuberances. To this end, we re-used the micro-pillar arrays implemented in a previous work [34], with a pillar spacing of 200 μm.

The four electrodes (μPG-MG) exhibited similar chronoamperograms (Fig. S3 in Supplementary Data). The two μPG-MG-1 electrodes gave identical chronoamperogram characteristics (Table 3). The μPG-MG-2 electrode gave slightly higher r_i and J_{max} values, which suggested that the thicker and looser porous structure observed on the top of the μPG-MG-2 surface was the most suitable for the development of electroactive biofilm. Nevertheless, close J_{max} values were obtained, with only around 30% variations (Table 3), while their electrochemically active surface area varied by a factor of two (Table 2). The current produced by the bioanodes was thus slightly impacted by the electrochemically active surface area.

Cyclic voltammograms recorded at day 7, when the electrodes were not far from producing the maximum current density, confirmed the fair reproducibility observed on the electrochemical characteristics (Fig. S3B in Supplementary Data).

At the end of the experiments, electrode colonization was characterized by epifluorescence microscopy (Fig. 10). For all the bioanodes, the bottom of the electrode structure displayed faint fluorescence, indicating a thin biofilm. In contrast, the top of the pillars looked well colonized by a thicker biofilm. On the μP-MG-1 electrodes, the colonization on top of the pillars was uniform with a fully covering biofilm (Fig. 10A). On μP-MG-2 and -3, the biofilm did not clog the pores and had penetrated the porous structure, at least for a few micrometres (it was not possible to see deeper into the porous structures). The microscopic observation revealed a clear difference in biofilm formation: well-formed biofilms on the top and poor colonization at the bottom.

The previous study [34], which implemented the same micro-pillar arrays but with a nano-rough surface (R_a = 4.5 nm), led to current densities of 5.9 and 6.3 A·m^{-2} related to the projected surface area. Here, roughening the pillar surface led to a very modest gain, of 1.3-fold at most.

In comparison, here, in flat configuration, the MG surfaces obtained by roughening the same NG surface increased the J_{max} by a factor of 2.7 to 6.7 (Table 3). Roughening greatly impacted the current produced by the flat bioanodes, but it lost much of its interest in the case of the micro-pillar arrays.

The biofilm structure described above suggests an explanation for this difference between flat surfaces and micro-pillar arrays.
The biofilms were well-developed on the top of the micro-pillars, while the bottom was poorly colonized. It can consequently be assumed that the current was mainly produced by the fully developed biofilm on top of the micro-pillars (Fig. 10). Only the top part of the micro-pillar array was efficient.

An elementary quantification supports this hypothesis. On flat micro-rough surfaces, the highest current densities recorded were of the order of 14 A·m$^{-2}$ (Table 3). The highest current density recorded on roughened micro-pillars was roughly of the order of 8 A·m$^{-2}$. Assuming that the surface of the roughened micro-pillars produces the same current density as the roughened flat surface, 0.57 cm$^2$ of micro-pillar surface area would be sufficient to obtain 8 A·m$^{-2}$. A micro-pillar array of 1 cm$^2$ projected surface area corresponds to a geometric surface area of 3.18 cm$^2$ (500 μm height, 100 μm wide, 200 μm spacing). This means that around 18% of the surface area of the micro-pillar array working at 14 A·m$^{-2}$ would be sufficient to produce 8 A·m$^{-2}$. This surface ratio corresponds to the top of the micro-pillars and around 100 μm of their upper sides. Only the upper part of the micro-pillar array was working.

This hypothesis is reinforced by the theoretical approach developed previously [34], which showed strong mass transfer limitation of the buffer species HCO$_3^-$ inside a pillar array. Acetate oxidation causes acidification in the vicinity of the bioanode surface:

$$\text{CH}_3\text{COO}^- + 2\text{H}_2\text{O} \rightarrow 2\text{CO}_2 + 7\text{H}^+ + 8\text{e}^- \quad (3)$$

which has been shown to be heavily detrimental to electroactive biofilms [46–48]. According to the equations developed previously, the modified Damköhler number ($\text{Da}_{\text{mod}}$) relative to HCO$_3^-$ is:

$$\text{Da}_{\text{mod}} = \frac{J_{\text{loc}} \delta}{nFDC_0} \quad (4)$$

where $J_{\text{loc}}$ (A·m$^{-2}$) is the local current density, i.e. the current density expressed with respect to the geometric surface area of the pillars, $\delta$ (500 10$^{-6}$ m) is the micro-pillar height, $n$ is the number of electrons produced per molecule of HCO$_3^-$ consumed, and $F$ (96 485C·mol$^{-1}$) is the Faraday constant, $D$ (1.18 10$^{-9}$ m$^2$s$^{-1}$ [49]) is the diffusion coefficient of HCO$_3^-$, and $C_0$ (mol·m$^{-3}$) is the concentration at the top of the micro-pillar array.

The value of $n$ was taken as 8/7 because, for 8 electrons produced, 7 mol of HCO$_3^-$ are consumed to neutralize the 7 protons produced (Reaction 3). The concentration of HCO$_3^-$ was 29.6 mM in the bulk of the solution and $C_0$ was calculated by assuming a diffusion layer of
100 μm above the top of the micro-pillar array (Fig. 9), which is a common value for an electrochemical process in quiescent solution. $D_{\text{admod}}$ expresses the rate of consumption by the electrochemical reaction with respect to the maximum diffusion rate along the micro-pillar. It must be corrected by a geometric ratio ($\zeta$):

$$\zeta = \frac{I_{loc}\delta}{S}$$  

(5)

where $I_{loc}$ (m) is the electrochemically active length in the section of the elementary unit and $S$ (m$^2$) is the section of the elementary unit that is free for diffusion (Scheme 1). $\zeta$ expresses the overall electrochemically active area in the elementary unit with respect to the section available for diffusion.

It was assumed that, locally, the surface of the roughened micro-pillars produced the same current density as the roughened flat surface, so $I_{loc} = 14$ A·m$^{-2}$. In this condition, $\zeta D_{\text{admod}}$ is equal to 5.7, showing that the rate of consumption by the electrochemical reaction is considerably higher than the maximum diffusion rate of the buffering species. With this theoretical approach, the $\text{HCO}_3^-$ concentration and the pH profiles along the micro-pillars can be calculated (Fig. 11, continuous lines). The model predicts that $\text{HCO}_3^-$ is consumed very early in the upper part of the micro-pillar structure, so the pH sinks to very low values. The order of magnitude of the penetration depth of $\text{HCO}_3^-$ is around 60 μm. This value is consistent with the elementary calculation made above, which indicated that 100 μm of pillar height was sufficient to ensure the current density of 8 A·m$^{-2}$. Actually, the penetration depth of the $\text{HCO}_3^-$ should be enhanced by the migration flux, which has been neglected in the reaction-diffusion approach used to establish the Damköhler number.

For comparison, the calculation made in the previous work [34] for nano-rough micro-pillars that produced 6.1 A·m$^{-2}$ with $I_{loc}$ of 1.87 A·m$^{-2}$ is also plotted in Fig. 11 (dashed lines). In this case, the diffusion of $\text{HCO}_3^-$ along the micro-pillar also showed significant limitation, but it reached the bottom of the structure. This confirmed that the fall of $\text{HCO}_3^-$ concentration in the upper part of the micro-pillar structure was due to the increase of the local reaction rate ($I_{loc}$) because of the roughened structure.

In conclusion, the three approaches:
- observation of the very different levels of biofilm development at the top and bottom of the micro-pillar structure,
- comparison with the current density produced by flat micro-rough surfaces,
- theoretical modelling of $\text{HCO}_3^-$ transport,

converge to show that roughening the micro-pillar surface resulted in high efficiency of the upper part, which precluded the exploitation of lower area of the micro-pillars. The high local current density achieved on the upper part of the micro-pillars consumed the whole flux of the buffering species, which could consequently not mitigate the acidification of the lower part of the array. This resulted in severe acidification of the lower part of the micro-structure (Scheme 2).

![Scheme 1. Elementary unit of a micro-pillar array. A) top view; B) the side view. S is the section free for diffusion, $\delta_{dl}$ in the diffusion layer above the micro-pillar array; on the side view the concentration profile of a species consumed by the electrochemical reaction is schematized.](image1)

![Scheme 2. Roughened micro-pillar array. The efficient upper part (red) of the micro-pillars causes a severe acidification of the lower part (yellow).](image2)

4. Discussion and further research directions

A correlation was sought by expressing the maximum current density with respect to the electrochemically active surface area ($A_e$). In this form, the current density produced by the micro-rough electrodes (MG) was lower than that produced by the nano-rough electrodes (NG), and the current density decreased as $S_a$ increased (Table 4). When the roughness increased, the increased electrochemically active surface area was less and less accessible for microbial settlement. Furthermore, the current density related to $A_e$ was even smaller for the roughened micro-pillar arrays than for the MG electrodes. This raises the question of the most appropriate way to measure the active surface area available for biofilm settlement. Here, expressing the current density with respect to $A_e$ gave considerably higher values for the NG electrodes while, in practice, the NG electrodes produced smaller currents. The electrochemical technique used here to measure...
Ae overestimated the surface area that could actually be reached by the microbial cells. The electrochemical measurements were based on the oxidation and reduction of the oxide layers, which involves only a few monomolecular layers on the gold surface. This measurement scale, at the Angstrom level, was much too small to assess the surface available for microbial cells, which have characteristic sizes of at least 1 μm. The electrochemically active surface area is a useful parameter for assessing the physical effect of the roughening technique, but it cannot be used to calculate the local current density. This surface area is overestimated with respect to the surface area that is really available for cell settlement. In the case of the micro-pillar arrays, the mass transfer limitation is superimposed and causes the current density to drop further. The electrodeposition technique used here succeeded in producing gold deposits favouring tight bacterial colonization and ensuring efficient electron transfer with electroactive cells. The proximity of the voltammogram curves with the theoretical Nernst-Michaelis-Menten kinetics was a proof of the electron transfer efficiency that occurred between these gold surfaces and the microbial cells.

Roughness at the micrometre scale greatly increased the current density with respect to the nano-rough surface (Ra = 4.5 nm), by a factor of 2.7 to 6.7. Actually, nano-rough electrodes were not fully appropriate for biofilm formation. Difficult bacterial adhesion resulted in low and poorly reproducible biofilm coverage even after more than 12 days of polarization. Micro-roughening of the surface, from 0.5 to 6.7 μm, favoured bacterial adhesion and led to full biofilm formation and higher current production. The complete, uniform biofilm coverage promoted by the micro-roughness is undoubtedly a major reason for the great current enhancement.

The current density increased linearly with S4. It was observed that the 10-μm micro-craters (MG-3 and MG-4) accelerated biofilm growth locally but did not increase the current density beyond the value due to the increase in S4. It can be concluded that S4 is a fully appropriate parameter for checking the effect of surface topography on the current density provided by electroactive biofilms. Nevertheless, the decoupling observed between the status of biofilm development and the current density shows that biofilm electroactivity is a complex process that does not correspond to the simple addition of similar contributions from individual microbial cells.

The porous layers produced current densities of up to 15.9 A·m⁻², i.e. 14.8 ± 1.1 A·m⁻² on average for 3 electrodes. Nevertheless, the porous structure was far from being fully exploited because of the formation of a covering biofilm on the upper surface that clogged the porosity. As a major message, it should be kept in mind that pores several tens of micrometres in diameter were quickly clogged by the biofilm.

A similar result has already been suggested in the literature by Kipf et al., who observed that Geobacter sulfurreducens unexpectedly gave similar performance on activated carbon and graphite felt [50]. The thick biofilm formed was suggested to be an element of the explanation. Similarly, microbial anodes formed on 2-dimensional carbon cloth have been shown to perform as efficiently as those formed on porous felt, because the biofilm formed on the upper side of the carbon felt clogged the porosity of the felt electrode [51]. In this case, the operating conditions were very different, microbial anodes were formed in wastewater, fed with food waste and inoculated with biological sludge. The present study confirmed that, even in well-controlled pure culture conditions, electroactive biofilms can clog pores several tens of micrometres in diameter quite fast during their formation phase.

With respect to the basic nano-rough electrode, current density was multiplied by a factor greater than 7 when the porous layer was used. The porous structure greatly improved the current density. This improvement was directly linked to the increase in surface roughness. Porosity acted as an increase of surface roughness, i.e. through the 2-dimensional effect it induced on the upper surface, rather than through a 3-dimensional effect. In this context, Sa remains the appropriate parameter to assess the effect of electrode topography on the current density produced. Moreover, the linear relationship observed with the micro-rough electrodes remained valid with the porous ones. This is a major conclusion that should be kept in mind to possibly reinterpret the impact of porosity on microbial anode performance. Actually, some current increases that have been roughly attributed to the effect of 3-dimensional porosity in the literature so far may have been due to the impact of the porosity on the upper surface roughness.

As illustrated in the introduction, the results reported in the literature on the impact of the surface micro-roughness on electroactive biofilms are diverse. The results obtained here are in line with the reports that have observed an impact of the roughness at the size of a few micrometres [23,28,52,53]. These studies didn’t propose correlations between roughness and the produced current density. Here, taking care to modify the surface roughness, without affecting other surface parameters, revealed a significant and increasing impact of surface roughness over a wide range from 0.5 to around 8 μm, including the effect of porosity. This range of action is larger than reported in a study carried out with stainless steel, which observed an effect of roughness only above 2 μm [23].

Protuberant micro-structures have been shown to improve the performance of microbial anodes, mainly by increasing the surface area available for biofilm growth. Gold cross-shaped micro-pillars 40 μm wide and 8 μm high have been shown to increase the current produced by S. cerevisiae 4.9 fold because they expanded the surface area to volume ratio by a factor of 4.5 [54]. The presence of 20 μm high and 20 μm diameter cylindrical micro-pillar arrays improved the current produced by S. marcescens 1.5 times [55]. In our previous study, 100 μm wide and 500 μm high micro-pillar patterning led to current densities being improved by factors of up to 8, related to higher surface area [34].

Here, it was decided to use the same micro-pillar arrays as previously investigated, with the objective of improving the performance by roughening the pillar surfaces. The effect of roughening was not as impressive as expected. Current density values showed a modest 1.3-fold gain with respect to the non-roughened micro-pillar arrays. The theoretical approach indicated that current enhancement was effective but the local high current density produced at the top of the micro-pillars exacerbated the mass transfer limitation. Mass transfer of the buffering species was no longer able to mitigate the acidification at the bottom of the micro-pillar array. This issue points out, once again, the crucial importance of interfacial acidification [46–48] and the importance of ion transport to mitigate it [56].

The upper part of the roughened micro-pillars was too efficient and precluded functioning of the bottom part. The theoretical approach indicated that, with the roughened surface, micro-pillars of around 60- to 100-μm height would be sufficient. Higher pillars could be used if the space between them was increased in order to favour mass transport. An optimum has to be found, and the simple mass transfer calculation used here provides some first basic guidelines for optimizing
the topography of bioanodes. Furthermore, it may be favourable to decrease the micro-pillar width to a few tens of micrometres, or less if technologically possible, with the objective to induce ultra-philic electroactive strains, but they remain rare and not really efficient [58]. Increasing the concentration of the buffer species by acclimating electroactive species to high buffer concentrations may be another promising way to progress on bioanode performance.

In conclusion, the present results indicate that creating surface roughness of a few micrometres is a fully appropriate way to accelerate the formation of bioanodes and to increase the current they produce in their formation phase. Progress might still be made by micro-structuring the surface. In this context, the optimal topography is directly dependent on the capacity of the strain to resist acidification. A simple theoretical tool is described here to start optimizing the micro-design of bioanode surfaces.

5. Conclusion

Current density was increased by a factor of up to 6.7 by surface micro-roughness with respect to nano-rough surfaces. For $S_b$ ranging from 0.5 to 7.6 $\mu$m, current density increased linearly with $S_b$ and this relationship remained valid for porous structures. Porosity was shown to impact the bioanode by modifying the 2-dimensional surface topography here rather than through 3-dimensional colonization. Combining roughness and micro-pillars resulted in lower current density than flat roughness. This result was explained by a limitation on ion transport inside the micro-pillar arrays and a simple theoretical tool was provided to progress towards optimal surface topography.

It must be kept in mind that the conclusions drawn here concern the early formation of Geobacter sulfurreducens biofilms. The main objective was to impact the starting phase in order to accelerate the biofilm formation and the current production. These results should now be confronted with different situations: mature biofilms, biofilms of other strains and multispecies biofilms [59]. Unravelling the impact of the surface topography on the formation and performance of microbial anodes will be a long and complex task. Here, we hope to have provided the research community with a basic brick that will help to build up a sound foundation for future works.

Declarations of interest

None.

Acknowledgements

This work received the support of the French state, managed by the Agence Nationale de la Recherche (ANR), within the framework of the project Koropokkuru (ANR-14-CEDS-0004).

The authors would like to acknowledge Luc Etcheverry, Frédéric DaCosta, Jean-Pierre Escavit and Vincent Loisel (Laboratoire de Génie Chimique) for their help in designing and setting up the reactors. Cécile Pouzet (Plateforme Imagerie-FR AIB) and Benjamin Erable (Laboratoire de Génie Chimique) are gratefully acknowledged for their implication in the fluorescence imaging acquisition, and Vincent Baylac (CIRIMAT-NNC OVM RTS) for his help in the topographical analysis.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.bioelechem.2019.03.002.

References


