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Rotating disk electrodes to assess river biofilm thickness and elasticity

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ABSTRACT

Keywords: Epilithon Periphyton Biofilm architecture Biofilm deformation Voltammetry Electrochemistry The present study examined the relevance of an electrochemical method based on a rotating disk electrode (RDE) to assess river biofilm thickness and elasticity. An in situ colonisation experiment in the River Garonne (France) in August 2009 sought to obtain natural river biofilms exhibiting differentiated architecture. A constricted pipe providing two contrasted flow conditions (about 0.1 and 0.45 m s $^{-1}$ in inflow and constricted sections respectively) and containing 24 RDE was immersed in the river for 21 days. Biofilm thickness and elasticity were quantified using an electrochemical assay on 7 and 21 days old RDE-grown biofilms $(t_7 \text{ and } t_{21}, \text{ respectively})$. Biofilm thickness was affected by colonisation length and flow conditions and ranged from $36 \pm 15 \,\mu\text{m}$ (mean \pm standard deviation, n = 6) in the fast flow section at t_7 to 340 \pm 140 μ m (n = 3) in the slow flow section at t_{21} . Comparing the electrochemical signal to stereomicroscopic estimates of biofilms thickness indicated that the method consistently allowed (i) to detect early biofilm colonisation in the river and (ii) to measure biofilm thickness of up to a few hundred µm. Biofilm elasticity, i.e. biofilm squeeze by hydrodynamic constraint, was significantly higher in the slow (1300 \pm 480 μ m rpm^{1/2}, n = 8) than in the fast flow sections (790 \pm 350 μ m rpm^{1/2}, n = 11). Diatom and bacterial density, and biofilm-covered RDE surface analyses (i) confirmed that microbial accrual resulted in biofilm formation on the RDE surface, and (ii) indicated that thickness and elasticity represent useful integrative parameters of biofilm architecture that could be measured on natural river assemblages using the proposed electrochemical method.

1. Introduction

River epilithic biofilms are complex microbial consortia of algae, bacteria and other micro- and meso-organisms that develop on solid substrata (Lock, 1993). Embedded in a mucilage matrix of microbially generated biopolymers (EPS: *extracellular polymeric substances*), these aggregates have relatively high mechanical stability and cell density. River biofilm dynamics influences

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various instream processes such as primary production (Wetzel, 1975), river food web (Feminella and Hawkins, 1995), organic matter and nutrient cycling (Paul et al., 1991; Battin et al., 2003a; Teissier et al., 2007), and accumulation of contaminants such as pesticides (Dorigo et al., 2007) and toxic metals (Cheng et al., 2008; Thuy Dong et al., 2008).

Biofilm architecture (e.g. thickness, cohesion) varies with community maturation and resistance to current velocity, both for monospecific biofilms (e.g. Mukherjee et al., 2008) or for complex river biofilms (Peterson, 1996). Architecture partly conditions biofilm functions affecting mass transfer between aggregates and bulk water, influencing for example the relative uptake of substrates differing in bioavailability (Battin et al., 2003b). In spite of its major interest, the in situ characterisation of biofilm architecture remains a challenge since tools are very scarce, inconvenient to use in the field and somewhat semiquantitative. Among architectural parameters, thickness is the most integrative and informative with respect to variation in key parameters including volume, wet weight, and number of species. However river biofilm thickness is rarely measured and studies often intentionally use biomass as an indirect estimation of thickness (Dodds et al., 1999). Several destructive (scanning electron microscopy, cryoembedding) and nondestructive (light microscopy, scanner with an image acquisition system, a laser triangulation sensor, confocal laserscanning microscopy and two-photon excitation microscopy) optical methods are available to measure biofilm thickness (Paramonova et al., 2007). They are ideal tools for biofilm monitoring at the micrometer scale spatial resolution. Investigations on bacterial biofilms are also oriented towards nanoscopic spatial arrangement using a combination of confocal laser-scanning microscopy and atomic force microscopy (Schmid et al., 2008). The main drawback for their application to river biofilm is the incompatibility between their observation scale and the centimetre or metre scale of biofilm development in rivers (e.g. on rock substrates such as pebbles). An optical method (Bakke and Olsson, 1986), periodically applied for river and estuarine biofilms (Sekar et al., 2002; Rao, 2003) determines biofilm thickness as the vertical sample displacement required to move the focal plane of the microscope from the water--biofilm interface to the biofilm-substratum interface. It is limited in that an estimate of the refractive index of the transparent film is required and it can only be applied to biofilm thinner than 100 µm (Paramonova et al., 2007).

Herbert-Guillou et al. (1999) reported an electrochemical method based on the analysis of a tracer oxidation current on a rotating disk electrode (RDE) where biofilm has developed. This electrochemical technique was applied to detect very thin bacterial biofilms developed in sea and tap waters (Herbert-Guillou et al., 2000; Gamby et al., 2008). Beside thickness measurement, Herbert-Guillou et al. (2000) showed that the RDE method could be used to provide complementary information on biofilm functional properties relative to biofilm elasticity.

The objectives of the present study were to (i) adapt the RDE method to estimate natural phototrophic biofilm thickness and elasticity and particularly, (ii) improve the biofilm elasticity parameter calculation, (iii) assess the relevance of thickness and biofilm elasticity measurements to differentiate contrasted river phototrophic biofilms and, (iv) prove the suitability of this

method for in situ experiments. As flow rate and biofilm maturation are proved to influence biofilm architecture (Peterson, 1996), we designed an experimental device to produce 7-day and 21-day-old biofilms in situ while varying the flow rate.

2. Materials and methods

2.1. Experimental design

2.1.1. Biofilm production device

An experimental pipe device for biofilm production was designed and scaled to provide two contrasted current velocity conditions within the same pipe, so that all factors affecting biofilm dynamics other than flow could be considered similar. According to the volume continuity equation for an incompressible fluid, through a pipe constriction (from the section #1 of area A_1 to the section #2 of area A_2), (i) the fluid velocity increases and (ii) this increase in velocity (from v_1 to v_2) is set to the decrease in section area as follows: $v_2/v_1 = A_1/A_2$.

The constricted pipe consisted in three main parts: an upstream first cylinder (section #1, slow flow) followed by a converging conical inlet (angle α_1) and a second downstream cylindrical throat (section #2, fast flow) (Fig. 1.). The current velocity v_1 was determined by the local river current velocity and followed river flow variations during the whole experiment. The current velocity v_2 depends on v_1 value and on the ratio between diameters (Φ_2/Φ_1) . Diameter dimensions were chosen (i) to provide a quite easily handling structure, (ii) to ensure relatively homogeneous flow conditions in each section and (iii) to ensure a ratio v_2/v_1 around 4. Inlet and throat diameters were set to 20 and 10 cm respectively. A diverging recovery part (angle α_2) followed by a third cylindrical throat (section #3; diameter $\Phi_3 = \Phi_1$) was added to the structure to ensure a straight exit stream. Convergence and divergence angles were chosen according to values minimising flow detachment and head loss in Venturi pipes: $\alpha_1 = 20^\circ$ and $\alpha_2 = 14^\circ$. Numerous formulas are found to estimate the entrance length (l_e) of cylindrical ducts i.e. the position beyond which flow is fully developed (Anselmet et al., 2009). Application of such formulas to the present flow conditions yields values of l_e/Φ_e between 20 and 30 lead to too long pipe dimensions to be handled in the river. Entry and constricted section lengths were set to 3 and 4 times the diameter, the total length being therefore 186 cm. At RDE locations, viscous shear stress on the cylinder (and incidentally on biofilm) is around 10 times larger in the constricted than in the entry section, ensuring relative homogeneous and contrasted local flows at RDE surfaces.

The constricted pipe was made of 3-mm thick Plexiglas[®] to ensure light diffusion. Pipe sections for which diameter was smaller than Φ_1 were surrounded with another 20-cm diameter Plexiglas[®] pipe to form a single continuous pipe and decrease detachment of the external flow around the pipe. The additional sheath did not affect light penetration: irradiance in both sections of the pipe, as measured using a LI-COR Li100 quantameter at sunlight, exhibited similar values within a 10% range.

2.1.2. Experimental procedure

Twelve RDE were incorporated at each downstream extremity of both sections of the apparatus (Fig. 1). The RDE were labelled



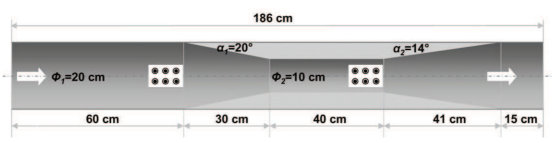


Fig. 1 – Photograph and schematic representation of the experimental pipe device. The position of the RDEs is indicated on the photograph by its labelling. SF: slow flow section; FF: fast flow section; 7: 7 days; 21: 21 days. Arrow shows current direction.

SF (for slow flow) or FF (for fast flow) according to which section they were located. In each section, the surface of 6 RDE per pipe side (right and left) was vertically positioned at the equator line to prevent particle sedimentation during the colonisation process. The RDE were positioned next to each other to ensure homogeneous environmental conditions between replicates. They were maintained in order to arise to the pipe internal surface with nylon cable gland allowing an easy recovery.

The constricted pipe was immersed parallel to the water current at the bottom of the River Garonne at the study site of l'Aouach (01°18′00″E; 43°23′08″N). This site is a typical reach for biofilm development (Lyautey et al., 2005; Boulêtreau et al., 2006). During the low-water period (from July to October), the study river reach is characterised by a shallow (<1.5 m), wide (100 m), and unshaded bed. Water exhibits low turbidity (<30 NTU) and nutrient concentrations of about 10 μ g P L⁻¹ of soluble reactive phosphorus, 1 mg N L^{-1} of both ammonium and nitrates, and 1.5 mg C L⁻¹ of dissolved organic carbon. The constricted pipe was maintained on the river bottom in a zone where the riverbed was flat and homogeneous (boulder rocks), shallow (water depth around 50 cm) and current velocity was slow (around 0.1 m s^{-1}). The experiment was performed on August 2009 during a low-flow period to exploit the most stable current velocities as possible, and to enable biofilm accrual especially in the fast flow section. Data on daily mean flow were supplied by DIREN Midi-Pyrénées (gauging station: Portet-sur-Garonne) and mean current velocity was measured at the pipe entry using an FLO-MATE portable flowmeter (Model 2000, Marsh-McBirney, USA).

The device was immersed for 21 days, and six RDE per section were sampled after 7 (t_7) and 21 (t_{21}) days of colonisation. Replicate RDE were named as follows: SF or FF when

collected in the slow flow or fast flow section followed by 7 or 21 according to the sampling time, and followed by the replicate number; RDE SF7#3 stands for one of the RDE sampled in the slow flow section after 7 days of colonisation. Sampled RDE were kept in river water at 4 °C in the dark during transport to the laboratory and measurements were performed within 5 h. At t_7 the 12 sampled RDE were replaced by stainless-steel cylinders of similar diameter.

2.2. Biofilm architecture measurements

2.2.1. Electrochemical measurement theory

The method consists of measuring the steady-state diffusion current on the RDE interface at a fixed potential and at a fixed rotation speed Ω without biofilm (t₀) and after biofilm development (t_7 and t_{21}). To impose this constant potential, a 3electrode-system immersed in an electrochemical cell filled with a tracer solution and connected to a potentiostat was used: (i) RDE, the working metallic electrode on which biofilm develops; (ii) the reference electrode that controls the potential of the working electrode and (iii) the counter electrode that closes the electrical circuit and the overall current goes through. Diffusion current results in the oxidation of a reduced species at the RDE-electrolyte interface. Without biofilm, diffusion current depends directly on the diffusion boundary layer thickness at the RDE-electrolyte interface. With RDE rotating at a constant rotation speed around its axis, the diffusion boundary layer thickness is maintained constant. Biofilm is considered as an inert porous layer with respect to mass transport since it contains more than 95% of water (Characklis, 1990). The biofilm is also considered as a layer of stagnant water on the RDE surface, and the slow convection existing inside the biofilm is neglected. The diffusion coefficient in biofilm was shown to be the same as the diffusion coefficient in water (L'Hostis et al., 1996), this property is extended for the thicker river biofilms under investigation in the present study. This layer adds to the hydrodynamic boundary layer one, inducing a decrease in diffusion current intensity.

2.2.2. Electrochemical measurement setting

The RDE was made of a 5-mm diameter platinum cylinder (electrical conductor) coated with a Teflon[®] cylinder (electrical insulator). The reference electrode was a saturated calomel electrode (SCE) (REF421, Radiometer Analytical, France). The counter electrode was a cylindrical grid of platinum immersed into the electrolyte solution that surrounded the working electrode. A 0.01M potassium ferrocyanide [Fe(CN)₆]²⁻ and ferricyanide [Fe(CN)₆]³⁻ solution was used as tracer in 1M KCl. Ferrocyanide oxidation current intensity was measured at 0 V/ SCE at which no water electrolysis and no oxygen reduction occur. Measurements were performed at 20 °C.

In the laboratory, the RDE was mounted on a motor axis plugged using mercury contacts and was rotated by a DC motor system. The motor speed was controlled with a servo system and measured using a tachometer. Prior to diffusion current measurements, the equilibrium potential of the ferrocynanide/ ferricyanide couple at the same concentration was measured between -0.240 and -0.236 V/SCE in accordance with the reference potential (-0.237 V/SCE). Diffusion current was then measured at the potential 0 V/SCE for each RDE rotation speed between 100 and 1200 rpm by steps of 100 rpm. Rotation speed was limited to 1200 rpm to prevent biofilm erosion. Before t_0 measurements, every RDE were polished using sandpaper (grade 1200) and cleaned with distilled water. After t_7 and t_{21} measurements, each RDE was individually conditioned into river water until further analyses.

Biofilm thickness δ (µm) was calculated from diffusion current intensity measurements with (i(t))and without biofilm (i(0)) for each RDE rotation speed (Ω in rpm) as follows:

$$\delta = nFDC^*S[i(t)^{-1} - i(0)^{-1}] \times 10,000$$
(1)

with *n* is the number of electrons, F the Faraday constant (96485 C mol⁻¹ or s A mol⁻¹), D the diffusion coefficient in both water and biofilm set to 6.8×10^{-6} cm² s⁻¹ at 20 °C according to Deslouis et al. (1980), C* the electroactive species concentration in the bulk solution (0.00001 mol cm⁻³), and S the active RDE area (0.196 cm²).

2.2.3. Image acquisition and analysis

For RDE biofilm cover estimations, stereomicroscopy (Olympus SZX10, 24× magnification) images of the bare RDE (t_0) and the wet colonised RDE (t_7 or t_{21}) surfaces were captured using an Olympus U-TV0.63XC camera (Olympus Corporation, Tokyo, Japan) as TIFF files (1600 by 1200 pixels) and imported in Photoshop CS3 (Adobe Photoshop v 10.0.1). No staining was performed. The image of the bare RDE surface was used as control. Binary images were generated by affecting the white color to the bare pixels and the black color to the colonised pixels. RDE biofilm cover (surface %) was determined on the platinum surface as the ratio of the surface area of black pixels to the total

surface area (sum of white and black pixels) with Image J 1.37v (Wayne Rasband, National Institutes of Health, USA).

For thickness estimation, stereomicroscopy (Leica MZ 12.5, $16 \times$ magnification) images of a side view of each colonised RDE standing in water were captured using a Leica DFC320 camera (Leica Microsystems DI Cambridge). Several focal planes corresponding to various cross sections ((x, z)-planes in a (x, y, z) coordinate system) were visible on the picture thanks to the setting of an appropriate depth of field. The projected image of the various focal planes was converted to binary image after biofilm pixels selection. The maximal biofilm height (maximal z-coordinate of the (y, z)-plane) on each abscissa of the image (x-axis) was measured automatically in pixels using Image J. Conversion from pixel to μ m was performed using a line scale standard. This gives the mean maximal biofilm thickness (mean z_{max}) of the whole colonised RDE surface ((x, y)-plane).

2.2.4. Cell numeration

After electrochemical measurements, material on the RDE surface was removed with a sterile scalpel and placed into 1 mL of filter-sterilized (0.2 μm pore-size filter) river water and preserved for storage at 4 $^\circ C$ with the addition of 100 μL of neutralized formaldehyde to the biofilm suspension. Biofilm suspension was sonicated in an ultrasonic bath (Elmasonic S900H, Elma, South Orange, NJ) at 37 kHz (15 min) and vortexed (15 min) according to Buesing and Gessner (2002). For bacterial counts, 500 µL aliquot of the appropriate cell suspension dilution was stained with 200 μ L DAPI (0.01 mg mL⁻¹) and collected by filtration on 0.2 µm pore-size black polycarbonate filters (Nuclepore, Whatman, Maidstone, UK) according to Garabetian et al. (1999). Counts were carried out on an Olympus BH2 RLFA microscope at 1250× magnification and results were expressed as cell number per cm². Diatom density in biofilm suspension was estimated directly (t_7) or after 5-fold dilution (t_{21}) using a Nageotte counting chamber, by counting the total number of diatoms in 30 fields (1.25 mL each, 0.5 mm depth), using light microscopy at 250 \times magnification (Olympus BH2 RLFA).

2.2.5. Statistical analyses

Electrochemical parameters (biofilm thickness and elasticity) were deduced by statistical adjustment using Origin 8.1 SR1 (v8.1.13 88, OriginLab Corporation, Northampton, USA). Agreement between simulated and measured thickness was evaluated by X^2 and R^2 application. The non-parametric Mann–Whitney U-test procedure was used to test for flow effects on biofilm thickness, biofilm elasticity, RDE biofilm cover, bacterial and diatom cell numbers. Correlation between biofilm architecture parameters was explored by using the Pearson r coefficient. All values are given as average \pm standard deviation (SD). Statistical analyses were performed with SPSS 15.0 software for Windows, and were considered significant at $p \leq 0.05$.

3. Results

3.1. Determination of biofilm thickness and elasticity

The reciprocal steady-state current intensity (mA^{-1}) was plotted against the reciprocal square root of the RDE rotation speed (rpm^{-0.5}) in the Koutecky-Levich coordinates in the

Fig. 2. For each EDT, before (t₀) or after biofilm colonisation $(t_7 \text{ or } t_{21})$, the current increased with the RDE rotation speed according to the Levich law (Levich, 1962). For a given rotation speed, the current decreased with biofilm formation (t_7 vs. t_0 and t₂₁ vs. t₀). This decrease in the current intensity measured between to and t7 or t21 was significant and allowed thickness determination using equation (1) for 22 RDE over 24. Connecting issues were at the origin of the defects on 2 RDE (SF7#1 and FF21#16). For 22 RDE (and even the most colonised ones), minimal recorded current intensities (i.e. intensity measured at the minimal rotation speed of 100 rpm) were higher than several tens of μA suggesting that the measurement was relevant (see Appendix). The slope is higher for 21- than for 7-day-old biofilms, and for slow than for fast flow grown biofilms (Fig. 2). Biofilm thickness measured at each RDE rotation speed (Ω) was represented on Fig. 3. The relationship between thickness and rotation speed can be analysed by considering the following law:

$$\delta = \frac{1}{(\delta_0)^{-1} + K \Omega^{0.5}}$$
(2)

 δ_0 (µm) is biofilm thickness at zero RDE rotation speed and, in other words, the theoretical biofilm thickness without any particular hydrodynamic constraint. The coefficient K (µm⁻¹ rpm^{-1/2}) relates the dependence of thickness with RDE rotation speed and was used to parametise biofilm elasticity as 1/K(µm rpm^{1/2}).

Parameter values are resumed in the Table 1. The derivative of δ vs. Ω may tend towards infinity when the rotation speed tends towards zero. This can result in a loss of accuracy on δ_0 yielding to unrealistic too large δ_0 for SF21#6, SF21#10 and SF21#12 parameter fits (as indicated using the infinity sign in Table 1). Such unrealistic values led us to exclude the corresponding RDE results. The poor agreement between measured and simulated thicknesses at high rotation speed for these RDE is likely to suggest that the law is not applicable under high rotation speeds for thick biofilms. Nevertheless weak X² values confirmed good fit quality for 19 out of 22 RDE; the calculated δ_0 values are reliable and ranged from 16 μ m after 7 days of colonisation to 500 µm after 21 days of colonisation. Electrochemically measured biofilm thicknesses were significantly correlated with stereomicroscopic estimates (Table 2). Electrochemical biofilm thickness estimates were 1.8-fold lower than stereomicroscopic estimates, ranging from 70 to 540 µm (Fig. 4).

3.2. In situ experimental settings

The RDE supporting device was designed to be immersed into the river ensuring both *in situ* environmental variability (algal and bacterial inoculum, light, temperature, nutrient, etc.) and two contrasted flow conditions. Flow velocity level in the pipe was controlled by natural temporal hydraulic changes in the river. Other than days 5-6-7 when the daily mean flow peaked at 99 m³, the river experienced a period of quite stable and low

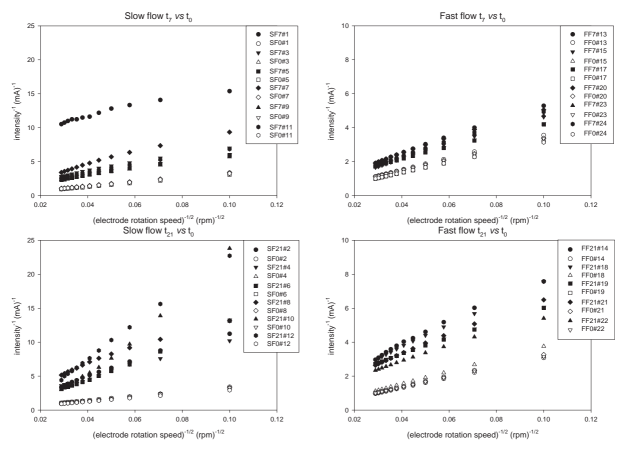


Fig. 2 – Inverse current intensity evolution with the electrode rotation speed measured on electrodes after different colonisation times (0 day, t_0 : closed symbols; 7 days, t_7 and 21 days, t_{21} : open symbols) in two flow sections (slow flow, SF and fast flow, FF) with the ferro-/ferricyanide tracer. Each symbol corresponds to one RDE.

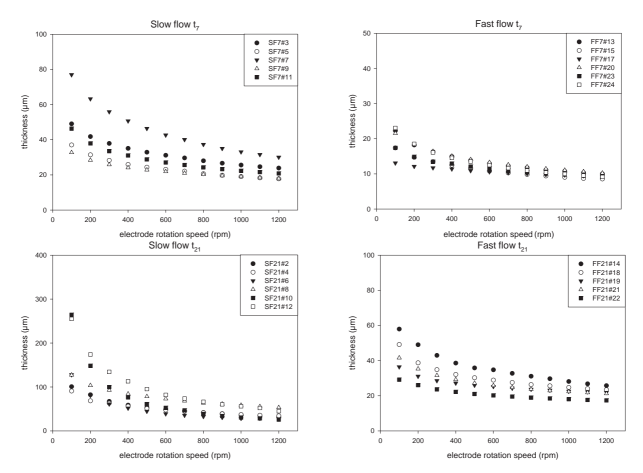


Fig. 3 - Thickness evolution with the electrode rotation speed measured on electrodes after two colonisation times (7 days, closed symbols and 21 days, open symbols) in two flow conditions (slow flow, SF and fast flow, FF) with the ferrocyanide tracer. Each symbol corresponds to one RDE.

flow (64 \pm 10 m³ s⁻¹) during the experiment, favouring biofilm development (data not shown). While measurement on day 7 highlighted the above mentioned 3-day period of hydraulic disturbance, other discrete measurements on days 0 and 21 in the slow flow section (i.e. inlet of the pipe) showed quite similar flow velocity values around 0.11 m s⁻¹ that correspond to a theoretical Reynolds number of 23,000 (Table 3). According to the device dimensions, flow velocity and Reynolds number in the fast flow section can be calculated from the former data to be around 0.46 m s⁻¹ and 46,000, respectively.

3.3. Biofilm features

Diatom accrual contributed to biofilm formation on the RDE. Diatom density increased during colonisation with 27×10^3 and 102×10^3 individuals per cm² in the slow flow section and with 8×10^3 and 33×10^3 individuals per cm² in the fast flow section on average at t_7 and t_{21} respectively (Fig. 5a). Consistently bacterial densities increased during colonisation reaching 32×10^6 and 27×10^6 cells per cm² on average at t_{21} in the slow and fast flow sections, respectively (Fig. 5b.). Comparing the two sections, diatoms densities were significantly different, whereas bacterial densities were not. As expected, RDE biofilm cover significantly increased between t_7 and t_{21} from 36 to 59% on average in the slow flow section and from 54 to 85% on average in the fast flow section (Fig. 5c.). Stereomicroscopic thickness significantly increased between t_7 and t_{21} and significantly decreased from the slow to the fast flow section (Fig. 5d).

Biofilm thickness significantly increased with time, means ranging from 100 to 340 μ m in slow flow and from 36 to 72 μ m in fast flow (Fig. 5e). Biofilm thickness was significantly affected by flow conditions at both sampling times.

Significant (or quasi significant) changes in biofilm elasticity values (1/K) occurred between t_7 and t_{21} and between flow conditions (Fig. 5f.). Mean (1/K) values were significantly higher in the slow (1300 μ m rpm^{1/2}) than in the fast flow section (790 μ m rpm^{1/2}) (Mann–Whitney U-test, p = 0.032).

Electrochemical thickness measurements were significantly correlated with RDE biofilm cover, diatom and bacterial densities (Table 2). In addition, significant correlation was also observed between biofilm elasticity and other parameters except bacterial density.

4. Discussion

Ecologists agree to consider thickness increase as the driving force of biofilm structural and functional properties (Sabater Table 1 – Results of parameter fits (minimisation Chisquare): parameter values (average ± square deviation) and fit quality (χ^2 /degree of freedom; R²) for each RDE.

	$\delta_0(\mu m)$	K (μm ⁻¹ rpm ^{-1/2})	$\frac{\chi_2}{dof}$	R ²			
Slow flow t	7						
SF7#3	87 ± 2	0.00084 ± 0.00001	0.13	0.9978			
SF7#5	65 ± 1	0.00116 ± 0.00001	0.04	0.9987			
SF7#7	193 ± 13	0.00076 ± 0.00002	1.52	0.9926			
SF7#9	49 ± 1	0.00103 ± 0.00001	0.03	0.9986			
SF7#11	90 ± 1	0.00106 ± 0.00001	0.04	0.9993			
Fast flow t ₇							
FF7#13	44 ± 1	0.00222 ± 0.00002	0.01	0.9995			
FF7#15	30 ± 1	0.00241 ± 0.00003	0.02	0.9978			
FF7#17	16 ± 0	0.00124 ± 0.00002	0.00	0.9970			
FF7#20	39 ± 0	0.00203 ± 0.00001	0.01	0.9995			
FF7#23	27 ± 0	0.00205 ± 0.00003	0.02	0.9973			
FF7#24	58 ± 1	0.00260 ± 0.00002	0.02	0.9991			
Slow flow t ₂₁							
SF21#2	501 ± 108	0.00077 ± 0.00003	5.81	0.9869			
SF21#4	252 ± 18	0.00071 ± 0.00002	1.70	0.9939			
SF21#6	+∞ ^a	0.00090 ± 0.00004	58	0.9328			
SF21#8	277 ± 13	0.00042 ± 0.00001	1.43	0.9971			
SF21#10	+∞ ^a	0.00053 ± 0.00018	1222	0.7460			
SF21#12	$+\infty^{a}$	0.00044 ± 0.00002	265	0.9304			
Fast flow t_2	1						
FF21#14	114 ± 3	0.00084 ± 0.00001	0.26	0.9973			
FF21#18	86 ± 4	0.00094 ± 0.00003	0.78	0.9861			
FF21#19	48 ± 2	0.00076 ± 0.00003	0.40	0.9780			
FF21#21	69 ± 1	0.00097 ± 0.00001	0.09	0.9976			
FF21#22	40 ± 1	0.00099 ± 0.00002	0.07	0.9949			
a $+\infty$ Indicates an unrealistic too large thickness value.							

and Admiraal, 2005), but, studies on river biofilms suffer from a lack of available tools to characterise biofilm architecture. The present study intended to assess the ability of an electrochemical method based on rotating disk electrode to measure and evaluate two features of biofilm architecture: thickness and elasticity.

Previously, the electrochemical method measured only very thin bacterial biofilms, between 0.9 and 3.5-µm thick in tap water (Gamby et al., 2008), and up to 10-µm thick in seawater (Herbert-Guillou et al., 1999). The use of 1 M KCl in the electrochemical assay could be expected to cause thickness underestimation due to EPS constriction (Frank and Belfort, 1997).

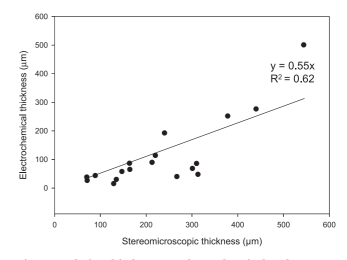


Fig. 4 – Relationship between electrochemical and stereomicroscopic measurements of biofilm thickness.

However, in their previous experiments, electrochemical estimates of biofilm thickness were validated by means of confocal laser-scanning microscopy (L'Hostis, 1996). In the present study, stereomicroscopy was used since the whole colonised RDE surface can be examined, and microbial counts can then further be done on fresh material since it does not require any previous processing such as staining, cryoembedding or cryosectioning. Stereomicroscopic measurements cannot provide absolute thickness values, but gave the upper limit of biofilm thickness range for each RDE. Nevertheless, the agreement between electrochemical measurements and stereomicroscopic estimates of biofilm thickness, 2-fold higher than the electrochemical one, confirmed the relevance of the electrochemical approach to usefully measure thicknesses ranging from a few μm to several hundreds of $\mu m.$ The electrochemical method is suitable for studying biofilms containing not only prokaryotic but also eukaryotic microorganisms such as microphytobenthic algae, and particularly diatoms. Stacking of diatom cells, typically several $10 \,\mu m$ in size, would give a biofilm cluster of hundreds of μm in thicknesses. Our measurements are thus consistent with the expected thicknesses for such biofilms.

The second parameter measurable by electrochemistry is biofilm elasticity. Initially Herbert-Guillou et al. (2000) found direct variation of bacterial biofilm thickness with electrode

Parameter	δ_0	1/K	Bacterial density	Diatom density	RDE biofilm cover	Stereomicroscopic thickness
δ ₀	1.000	0.615**	0.480*	0.764***	0.680***	0.833***
1/K		1.000	0.428	0.696***	0.700***	0.781***
Bacterial density			1.000	0.533**	0.561**	0.646***
Diatom density				1.000	0.714***	0.755***
RDE biofilm cover					1.000	0.822***
Stereomicroscopic thickness						1.000

Table 3 – Theoretical hydraulic characteristics in the slow and fast flow sections at t_0 (first day), t_7 (7 colonisation days) and t_{21} (21 colonisation days) estimated from measurements at the inlet of the pipe and pipe dimensions.

	Parameter	t _o	t ₇	t ₂₁
Slow flow	υ (m s ⁻¹)	0.11	0.30	0.12
	Re	22,000	60,000	24,000
Fast flow	ν (m s ⁻¹)	0.44	1.20	0.48
	Re	44,000	120,000	48,000

speed rotation, depending on biofilm development conditions. Therefore, they calculated biofilm deformation as the difference between electrochemical thickness at 100 rpm and thickness at a given rotation speed, and represented this latter as a function of electrode rotation speed. This simple relationship was not observed in the present study, probably because the studied biofilms contained algae and inorganic particles. Adapted from Foret (2006) that demonstrated the dependence of electrochemical thickness with $kQ^{-0.5}$ in water circuit biofilms, an original parameterisation of biofilm elasticity resulting from the assessment of an empirical

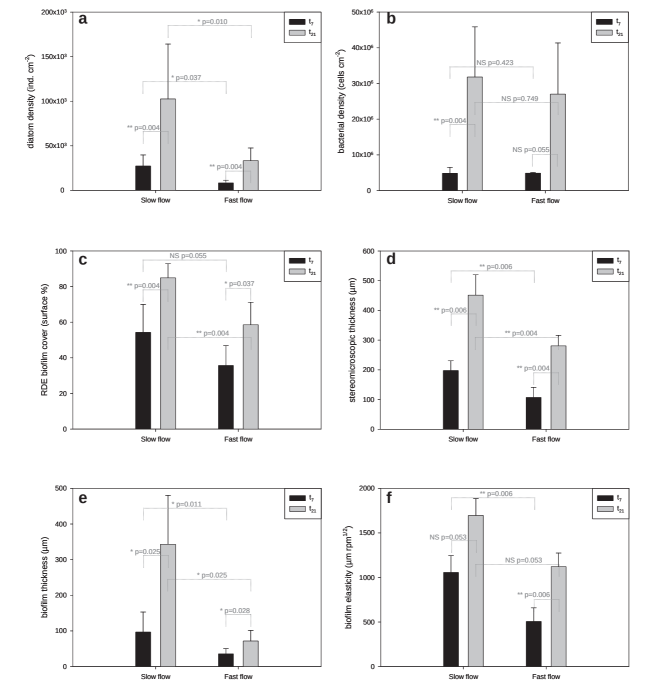


Fig. 5 – Effects of flow conditions (slow flow vs. fast flow) and colonisation time (t₇, black vertical bar vs. t₂₁, grey vertical bar) on diatom density (a), bacterial density (b), biofilm (electrochemical) thickness (c), elasticity (d), biofilm cover (e), and stereomicroscopic thickness (f).

relationship between biofilm thickness and RDE rotation speed $\Omega^{-0.5}$ was proposed here. Resulting elasticity values, displaying a wide range of magnitude from about 400 to 2400 μ m rpm^{1/2}, express the magnitude of biofilm thickness variation due to increasing rotation speed and quantify the extent to which biofilm can be reduced by hydrodynamics constraint. The values cannot be compared to existing data, however.

The in situ experiment was designed to compare core biological parameters to electrochemical parameters on natural river biofilms. As time is one of the main drivers of biofilm structuring, biofilms were sampled at two stages of biofilm accrual pattern, colonisation and maturation. Successional changes driven by changes in benthic microalgal species strategies result in temporal changes in biofilm structure (McCormick and Stevenson, 1991; Biggs et al., 1998; Wellnitz and Brader, 2003). Successional processes were also reported for river biofilm bacterial communities (Jackson et al., 2001; Lyautey et al., 2005; Lear et al., 2008). In the studied section of the River Garonne, biofilm bacterial richness proved to increase from 0 to 7 days, and decrease from 7 to 21 days (Lyautey et al., 2005), justifying the selected sampling times. The biofilm support material is known to influence biofilm community composition (Cattaneo and Amireault, 1992) and biofilms colonising RDE platinum may have exhibited distinctive taxonomic assemblages as compared to biofilms colonising river pebbles. An in-depth comparison of biofilm structure, biomass and composition between platinum and natural substrata is still to be performed, since no data on assemblage composition was recorded in the present study. Abundances of bacteria and diatoms were monitored, showing evidence of a microbial accrual on immersed RDE surfaces. Recovered densities were comparable to those previously observed in the River Garonne biofilms for diatoms, namely $10^5 - 10^7$ individuals per cm² (Eulin, 1997) and bacteria, about $10^7 - 10^8$ cell per cm² (Lyautey et al., 2010). Temporal evolution of microbial densities of RDE biofilms fitted with measured thickness enhancement. Interestingly, RDE biofilm cover increased with microbial densities and thickness suggesting that phototrophic river biofilms extend both horizontally and vertically in accordance with the typical model of biofilm development from isolated column forming clusters to connected mushrooms (Costerton et al., 1987). The proposed electrochemical assay was recommended to detect and survey fouling of man-made devices in marine and drinking waters (Herbert-Guillou et al., 1999; Gamby et al., 2008). It could also be used to evaluate the early dynamics of river biofilm e.g. the kinetics in the very early stage of colonisation in time course experiments or the patchiness of early accrual zones in microscale experiments.

Another main driver of biofilm structuring is flow. The RDE supporting device was imagined on the pattern of one Venturi pipe immersed into the river ensuring both *in situ* environmental variability (algal and bacterial inoculum, light, temperature, nutrient, etc.) and two contrasted flow conditions. As intended, generated current velocities, 0.11 and 0.46 m s⁻¹, were in the velocity range that favours such biofilm development (Horner and Welch, 1981). Despite disturbed hydraulic conditions for a 3-day period, stable and low daily mean flows occurred during most of the experiment especially during the whole maturation period. During stable and low-flow periods, typical Reynolds numbers (23,000 and

46,000) discriminated between optimal (Re near 22,000) and suboptimal biofilm growth conditions (Re > 40,000; Godillot et al., 2001). Consistently, higher diatom densities and biofilm thicknesses were found in the optimal flow section as compared to the other section. To our knowledge, only one study has quantified the effect of hydrodynamics on the thickness of stream microbial biofilms (Battin et al., 2003b): thicknesses deduced from confocal laser-scanning microscopy images of cryosections of biofilm were significantly higher for biofilms cultivated on ceramic coupons in the slow flow condition (0.065 m $s^{-1};\, \mbox{Re}=$ 1869) than in the fast flow condition (0.23 m s⁻¹; Re = 7559). The relationship between biofilm thickness and Reynolds number in the former and in the present study were consistent with Godillot et al. (2001) showing a maximum biofilm biomass for Re about 22,000. As for biofilm elasticity in the present study, biofilms produced in the slow flow section exhibited higher elasticity values than biofilms produced in the fast flow section. Most of the microorganisms that formed river biofilm biovolume are fitted with cellular structures maintaining cellular shape (e.g. bacterial cell walls, and diatom siliceous frustules). Biofilm elasticity most probably resulted rather from intercellular space reduction than from cell size constriction. Indeed, biofilm elasticity as defined in the present study might thus refer to voids (pores and channels) within biofilm and/or the looseness of cell adhesion in biofilm. Biofilm elasticity could thus fit with the sinuosity index of Battin et al. (2003b). The multiplication of pores or voids within biofilm contributes to enlarge biofilm surface area within biofilm and therefore facilitates biofilm - water interactions and advective solute transport (De Beer et al., 1996). Such mechanical property is well studied in biofilm models used to design and evaluate performance of biofilm reactors (e.g. Picioreanu et al., 1998). Biofilm elasticity as defined in the present study could be considered as an integrative parameter of biofilm-water interaction ability, in analogy with biofilm surface enlargement in studies of bacterial biofilms of industrial environments. For example, the reduction of biofilm-water interactions forming a barrier for advective solute transport could be an adaptative response of biofilm submitted to chemical stress. Indeed communities exposed to cadmium were primarily dominated by short stalked and ad-pressed diatom species whereas control communities were dominated by filamentous diatom species (Feurtet-Mazel et al., 2003). River biofilm architecture was also affected by chronic copper exposure through the growth of the chain-forming diatom Melosira varians changing from long filaments to short tufts (Barranguet et al., 2002). Such a qualitative observation might be quantified by measuring biofilm elasticity using the proposed electrochemical method. Further studies, addressing the relationship between biofilm architecture and the proposed measure of elasticity, might then allow to test whether biofilm physiognomic properties would reflect biofilm fitness at the community scale.

5. Conclusion

The present study showed the suitability of an electrochemical method based on rotating disk electrode to assess river biofilm thickness up to 500 μ m thick. Results extended the application domain of the method previously developed for tap water and seawater biofilms to complex biofilms mainly constructed by algae. The method reliably detected very thin biofilms, as well as measuring biofilm thickness of several hundred-µm. By analysing thickness evolution vs. electrode rotation speed, the electrochemical method can be used to calculate biofilm elasticity as an estimate of the extent to which biofilm is reduced by hydrodynamic constraint. This trait of biofilm architecture would relate to biofilm-water interactions. Very few studies have been conducted on the physical properties of river biofilms, due in part to technical difficulties associated with such complex biofilms. The electrochemical method developed here combined rotating disk electrodes which can be immersed directly in the river, and an electrochemical assay requiring only a few minutes. This nondestructive method is compatible with further analyses on the same sample e.g. bacterial or algal counts, pigment, or DNA extraction and analysis. Expanding the toolbox of biofilm characterisation techniques, the rotating disk electrode electrochemical method can be used to provide novel information on river biofilm architecture.

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Appendix. Supplementary data

Supplementary data related to this article can be found online at doi:10.1016/j.watres.2010.10.016.

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