Fluvial biofilm responses to joint changes in nutrients, temperature, turbidity and water velocity: an *ex situ* experiment

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**Abstract.** The aim of this study was to explore the responses of the epipelic biofilm of a Pampean stream with little impact from human activity to two environmental conditions, with joint modifications in nutrients, temperature, water velocity and turbidity. The experiment was conducted using artificial channels and lasted five weeks. The biological variables measured included chlorophyll-*a* content, bacterial biomass, ash-free dry weight, total carbohydrate concentration, total respiratory activity, and biofilm composition. Results show that the species’ composition of the biofilm was affected, although no other structural or metabolic variables measured were. These results highlight the importance of including structural parameters to measure rapid changes in water quality, even when analyzing the effects of co-occurring variables.

**Keywords:** Epipelic biofilms; Water quality; Artificial channels; Temperature; Nutrients.

**Introduction**

The land use change to produce goods and services represents one of the most important human alterations, affecting the structure and functioning of aquatic ecosystems in general. The loss of habitats, water extraction, pollution, resource overuse and the introduction of non native species have influences negatively in freshwater ecosystems. As a consequence to these pressures, the freshwater ecosystems are among the most vulnerable in the world (Revenga et al., 2005). Particularly in lotic fluvial systems, their sensitivity to human impact is magnified by their lineal and unidirectional nature; almost any activity conducted in the basin has the potential to influence the characteristics downstream up to a great extent (Malmqvist and Rundle, 2002).

In the Pampean plains, for example, are greatly influenced by human activity. The southernmost sector of these plains in
Argentina has over 21 million inhabitants (INDEC, 2010) that conduct an intense agricultural and industrial activity, favouring the input of contaminants to the water bodies. On the other hand, the climatic models for the region also predict higher rainfalls (Hulme and Sheard, 1999), which can increase erosion and generate flooding, increasing the transport of sediments, nutrients, and contaminants into the water (Davies-Colley et al., 1992; Davies-Colley and Smith, 2001). Although the lotic systems from the Pampean plains have naturally high concentrations of nutrients when compared to other lotic systems in the world (Meybeck, 1982; 1987; Giorgi et al., 2005; Feijó and Lombardo, 2007), these modifications in land use, intensified by the effect of changes in climatic patterns, have repercussions on their water quality, altering their physical, chemical and hydrologic properties (Rodrigues Capitulo et al., 2010). For instance, an experimental addition of nutrients in a Pampean stream evidenced responses of the biotic communities different to those observed in water bodies from other biomes (Artigas et al., 2013).

Among the diverse communities that inhabit the streams, biofilms composed of algae, bacteria, fungi and protozoa, all embedded in an extracellular matrix (Lock et al., 1984) represent a pertinent bioindicator of environmental perturbations within the aquatic ecosystem (Bonnineau et al., 2010; Romaní et al., 2016). This community is particularly sensitive to light and nutrients availability, and to the physical characteristics of water flow (Horner et al., 1990; Stevenson, 1996). On the one hand, increments in some factors such as light availability, concentrations of inorganic nutrients, temperature and water velocity, usually have a positive effect on biofilm development, that gets expressed as a proliferation in algal and bacterial biomass and elevated metabolism (e.g. Horner and Welch, 1981; Guasch et al., 1995; Dodds et al., 2002; Olapade and Leff, 2005), along with reductions in the proportion of carbohydrates (e.g. Freeman and Lock, 1995; Sutherland, 2001), and the proliferation of polysaprobic and eutrophic species. On the other hand, increments in other factors such as water velocity or turbidity usually have the opposite effects on the community (e.g. von Schiller et al., 2007; Romani and Sabater, 2000; Davies-Colley and Smith, 2001).

Research concerning the responses of the epipelic biofilms (those that develop on fine sediments) to environmental changes in template areas is scarce (Sierra and Gómez, 2007; 2010; 2013; Gómez et al., 2009; Cochero et al., 2013). In natural streams, however, biofilms are almost always subjected to multiple stressors that influence the overall water quality rather than to the modifications of a single factor (Breitburg et al., 1998; Venter et al., 2006; Halpern et al., 2007, 2008; Crain et al., 2008). Therefore, the aim of this study was to explore the possible responses of epipelic biofilms to two distinct environmental conditions, with joint increments of nutrients, temperature, water velocity and turbidity.

For this purpose, biofilms from a stream with little human impact were exposed to two environmental conditions, named Low and High treatment, that combined increments in temperature, nutrients (phosphorous and nitrogen), turbidity and water velocity. Changes in the structure and metabolism of the epipelic biofilms were measured for five weeks, predicting growth in both algal and bacterial biomass and total respiration, declines in the total concentration of carbohydrates in the sediment, and changes in the specific composition that favor species tolerant to more eutrophic conditions.

**Materials and methods**

A laboratory experiment in artificial channels was carried out using epipelic biofilm from a site in the “Martin Stream” (34° 54’ 51” S - 58° 04’ 39” W), which is exposed to the impact of low agricultural activity. The water quality of the site (Table 1) was assessed *a priori* by the concentrations of Soluble Reactive Phosphorous (SRP, mg P L⁻¹), Dissolved...
Inorganic Nitrogen (DIN, mg N L⁻¹), Dissolved Oxygen (DO, mg L⁻¹), Biochemical Oxygen Demand (BOD₅, mg L⁻¹), and Chemical Oxygen Demand (COD, mg L⁻¹) (Bartram and Balance, 1996).

Table 1. Dissolved oxygen concentration (DO), soluble reactive phosphorous (SRP), dissolved inorganic nitrogen (DIN), biochemical oxygen demand (BOD₅) and chemical oxygen demand (COD) measured in the “Martin” stream before the experimental stage.

| Parameter         | Value (± Standard Deviation) |
|-------------------|------------------------------|
| Temperature (°C)  | 23.43 (± 0.68)               |
| pH                | 8.77 (± 0.02)                |
| Conductivity (µS cm⁻¹) | 1536 (± 4.12)             |
| DO (mg L⁻¹)       | 7.70 (± 0.25)                |
| SRP (mg P L⁻¹)    | 0.37 (± 0.01)                |
| DIN (mg N L⁻¹)    | 0.22 (± 0.05)                |
| BOD₅ (mg L⁻¹)     | 6 (± 0.11)                   |
| COD (mg L⁻¹)      | 15 (± 0.57)                  |

Nine indoor artificial channels measuring 1 m (length) x 0.15 m (width) x 0.10 m (height) were used, each with an access ramp (40° slope) that ensures a laminar input flow (water depth was 0.10 m). Water exiting the channel flowed through a slit, and fell into a holding reservoir before being pumped back to the access ramp (Figure 1). All artificial channels were exposed to a photoperiod of 14 h light-10 h dark. Light was provided by GE® E-biax Helical lights (6,500°K, IRC82%) with an intensity of 110-115 µE m⁻² s⁻¹ of photosynthetically active radiation.

Figure 1. Experimental design employed (a) and schematics of one of the nine artificial channels used in the experiments (b). The physical-chemical variables in the control channels were kept similar to the values obtained in the field, while the treatment channels were exposed to the combined increments in four variables in two distinct levels.

Out of the nine channels, three were used as controls (C) and their physical-chemical variables were kept similar to the values measured at the stream. Another three channels (HIGH treatment) were exposed to a 4° C increase in temperature, 300% increase in nutrients (SRP and DIN), 50% increase in suspended solids and 20% increase in water velocity. These values were selected as they represent a realistic
characteristic of highly eutrophic sites in Pampean streams (Licursi and Gómez, 2002; Gómez et al., 2008; Sierra and Gómez, 2010), and by considering the increments forecasted in temperature and rainfall for the Pampean Region for the next decades (Hulme and Sheard, 1999) and the temperature-runoff relationship (Labat et al., 2004). The last three channels (LOW treatment) were exposed to intermediate levels of the manipulated variables: 1 °C increase in temperature, 50% increase in nutrients (SRP and DIN), 15% increase in suspended solids and 5% increase water velocity. These values were selected as they represent an intermediate alteration of the experimental variables.

Water temperature increments were achieved using regulated Atman 70 W water heaters placed in the individual tanks of each artificial channel. The increments in water velocity were achieved by calibrating the Chosen® Champion CX-500 water pumps, also placed in the individual tanks of each channel, at the proper speeds. Increases in turbidity were achieved by adding sterilized suspended solids to each channel from the corresponding stream where the biofilm was collected from. Nutrient increments (SRP and DIN) were achieved by adding dissolved Nitrofoska® fertilizer (of frequent use in the Pampean plain for agricultural purposes, 12% N - 12% P- 17% K) in the appropriate concentration for each channel. For every physical-chemical variable measured, three samples were collected from each channel, and the results obtained for each sample were averaged to be used in all statistical analyses.

For the biofilms to develop, each channel contained Falcon® multiwell polystyrene microplates first filled with sterilized sediment from the sampling site, for a total volume in each well of 3.4 cm³. Biofilm inocules were brought from the sampling site during the summer and added to the microplates, and water from the same site was circulated in each channel for 36 days. Water from all channels was partially renewed with filtered stream water twice a week to prevent metabolite accumulation.

### Physical-chemical variables
Dissolved Oxygen (DO, mg L⁻¹), temperature (°C), conductivity (µS cm⁻¹) and pH were measured using a CONSORT C933 sensor. Turbidity (NTU) was measured using an HORIBA U10 sensor, and water velocity (m s⁻¹) using a Schiltknecht MiniAir20. Nutrient samples were filtered through glass fiber filters (Whatman GF/F, Whatman International); ammonia, nitrites, nitrates and soluble reactive phosphorous were analyzed according to standard methods (APHA, 1998). Total dissolved inorganic nitrogen (DIN, mg N L⁻¹) was calculated as the sum of nitrate, nitrite and ammonia.

### Epipelic biofilm sampling
The epipelic biofilm samples were collected by pipetting the first 10 mm of the superficial layer (3.14 cm³) from wells selected at random in each channel. Samples for all biological analyses consisted of three subreplicates (three wells). All measurements performed in the biofilm samples were normalized to square centimeters of sand surface area, which was calculated as described in Marxsen and Witzel (1991).

### Community analysis
Three samples from each channel were fixed with formaldehyde (4%) and used to identify the community composition. Density of consumers and producers of the microbenthic community (size < 1 mm) were estimated using a Sedgwick–Rafter chamber (APHA, 1998) in an inverted optical microscope (Olympus BX 50) at 400X. The following keys were used for species identification: Bourrely (1966, 1968, 1970), Krammer and Lange-Bertalot (1986, 1988, 1991a, b), Tell and Conforti (1986), Streble and Krauter (1987), and Komárek and Anagnostidis (1999, 2005).

### Chlorophyll-a
Epipelic biofilm samples were filtered through Sartorious GF/C filters.
Samples were sonicated for 2 min in a Cleanson CS-1106 sonicator and filters were stored in the dark and frozen until they were analyzed. Chlorophyll-a (mg cm⁻²) was then extracted with 90% acetone for 12 h. The supernatant was read in a UV-VIS Auto 2602 spectrophotometer, and the concentration was calculated according to Strickland and Parsons (1968).

**Bacterial biomass**

Epipelic biofilm samples were stored in sterile glass vials with formalin 2% v/v. Bacterial density was estimated after sonication (three 2 min cycles) and appropriate dilution (1:100 to 1:400) of the samples. Diluted samples were stained for 10 min with DAPI (4',6-diamidino-2-phenylindole) to a final concentration of 1 µg mL⁻¹ (Porter and Feig, 1980), and filtered through a 0.2 µm black polycarbonate filter (GE Osmonics). Bacteria were then counted using an epifluorescence microscope (Olympus BX-50) under 1,000 x magnification. Twenty fields were counted for a total of 400 to 800 organisms per replicate. Bacterial biovolume was calculated assuming a 0.1 µm³ constant volume per bacterial cell (Romani et al., 2009), and bacterial biomass (µgC cm⁻²) was calculated from bacterial cell biovolume using the conversion factor of 2.2 x 10⁻¹ gC µm⁻³ (Bratbak and Dundas, 1984).

**Ash-free dry weight**

Samples were dried for 48 h at 60 °C, weighed, and re-weighed to determine the ash-free dry weight (AFDW) content (APHA, 1998).

**Total carbohydrates**

Epipelic biofilm samples were ground in 5 mL of 1 M H₂SO₄ with a glass rod in glass tubes, covered in aluminum foil and placed in a thermobath at 100 °C for an hour. An aliquot of 1 mL was separated from the supernatant and 1 mL of 5% phenol and 5 mL of concentrated H₂SO₄ were then added. After allowing the tubes to cool down for 30 min the samples were read in a UV-VIS Auto 2602 spectrophotometer at 485 nm (based on Dubois et al., 1956). Total carbohydrate (µg mL⁻¹) values were obtained using a glucose calibration curve.

**Total respiratory activity**

The activity of the electron transport system (ETS) was assayed by measuring the reduction of the electron transport acceptor INT (2-3 tetrazolium chloride) into INT-formazan (iodonitrotetrazolium formazan) (Blenkinsopp and Lock, 1990). Epipelic biofilm samples were incubated for 12 h on a shaker in the dark with 0.02% INT (2-(4-Iodophenyl)-3-(4-nitrophenyl)-5-phenyltetrazolium chloride) at room temperature. To terminate the reaction, 8 mL of methanol at 4 °C were added, and samples were filtered through Sartorius GF/C filters before reading them in a UV-VIS Auto 2602 spectrophotometer at 480 nm (Blenkinsopp and Lock, 1990). Total respiratory activity values (mg formazan cm⁻² h⁻¹) were calculated using an INT-formazan calibration curve (Sigma I7375).

**Data analysis**

Significant differences for the measured biological variables were analyzed using a two-way repeated measures analysis of variance (RM ANOVA) to test for the differences among treatments and dates (Winer, 1971). Homogeneity of variances was first checked using Cochrane’s test, and probabilities within groups were corrected for sphericity using the Greenhouse–Geisser correction. Differences in the physical-chemical variables between treatments were analyzed using one-way ANOVA, and the same analysis was used to monitor that the manipulated variables fell within the planned values. All post-hoc comparisons were made by Student–Newman–Keuls Test (SNK), and generalized eta² (ηG²) was computed as a measure of the effect size (Olejnik and Algina, 2003). This statistic has two major advantages over the traditional eta² and partial-eta²: first, it provides measures of effect size that are comparable across a wide variety of research designs; and
second, these effect-size measures provide indices of effect that are consistent with Cohen’s (1998) guidelines for defining the magnitude of the effect (Olejnik and Algina, 2003). These guidelines state that an effect size ≤ 0.20 is considered small, around 0.50 is considered a medium effect, and ≥ 0.80 is a large effect. η² provides comparability across between-subjects and within-subjects in repeated measures designs (Bakeman, 2005), and is estimated as:

$$ \eta^2 = G^2 = \frac{SS_A}{SS_A + \frac{SS_B}{A} + \frac{SS_P}{A}} $$

where: SS represents a Sums of Squares, A represents a between-subjects factor (Treatment), P represents a within-subjects factor (Time) and s represents the subjects factor.

Also, the overall differences between the controls and treatments in the composition (autotrophs and consumers) were analyzed conducting an Analysis of Similarity (ANOSIM), and a two-way similarity percentage analysis (SIMPER), based on the Bray–Curtis similarity measurement was used to determine the percent contribution of each taxon to the average dissimilarity between groups across all times (Clarke 1993).

Results

Physical-chemical parameters

The values obtained for the physical-chemical variables measured in the sampling site were used as the target values for the control channels in the laboratory. The manipulated variables in the channels (SRP, DIN, temperature, water velocity and turbidity) were significantly increased in both LOW and HIGH treatments when compared to the controls (p < 0.05, Table 2). Results from the analysis of variance show that these increments were in accordance with the planned increments for the experiment.

Table 2. Mean ± standard deviation of the physical-chemical variables measured in the experiment. A posteriori test Student-Neuman-Keuls (SNK) are shown when significant differences were found.

| Variable       | C          | LOW       | HIGH      | SNK                    |
|----------------|------------|-----------|-----------|------------------------|
| pH             | 8.7 (± 0.1)| 8.6 (± 0.1)| 8.5 (± 0.2)| C = LOW < HIGH         |
| Conductivity   | 633.6 (+ 256.7) | 723.5 (+ 266.8) | 1,075.4 (+ 292.4) | C = LOW < HIGH         |
| DO (mg L⁻¹)    | 7.7 (+ 0.02)| 7.7 (+ 0.02)| 7.6 (+ 0.02)| C = LOW < HIGH         |
| Temperature    | 24.7 (+ 1.3) | 25.8 (+ 1.2) | 28.9 (+ 1.4) | C < LOW < HIGH         |
| Turbidity      | 32.3 (+ 7.5) | 36.9 (+ 8.1) | 48.3 (+ 10.8) | C = LOW < HIGH         |
| Water Velocity | 0.35 (+ 0.01) | 0.37 (+ 0.01) | 0.43 (+ 0.01) | C < LOW < HIGH         |
| SRP (mg L⁻¹-P) | 0.20 (+ 0.12)| 0.327 (+ 0.19)| 0.891 (+ 0.53) | C < LOW < HIGH         |
| DIN (mg L⁻¹-N) | 0.43 (+ 0.40)| 0.75 (+ 0.65) | 1.39 (+ 1.30) | C < LOW < HIGH         |

C = Control, LOW = Low treatment, HIGH = High treatment.

Bacterial biomass

The bacterial biomass remained unvaried throughout the first three weeks in all channels, and increased in the fourth week indistinctly from the treatments (Figure 2a). The biomass in the controls had a mean of 16.64 (+25.96) µgC cm⁻², of 24.61 (+44.08) µgC cm⁻² in the LOW channels and of 30.09 (+69.45) µgC cm⁻² in the HIGH channels; the variation in the parameter throughout the experiment was high (ηG² = 0.64) (Table 3).

Algal biomass

The chlorophyll-α concentration in the channels remained similar throughout the experiment in all channels (Figure 2b). The mean values in the controls (0.41 ± 0.23 mg cm⁻²) were not significantly different from the values in the LOW
treatment (0.42 ± 0.21 mg cm⁻²) or the HIGH treatment (0.33 ± 0.14 mg cm⁻²).

**Ash-free dry weight**

The mean ash-free dry weight values were of 0.03 (±0.01) mg cm⁻² in the controls, of 0.05 (±0.08) mg cm⁻² in the LOW channels, and of 0.07 (±0.11) mg cm⁻² in the HIGH channels. The variation of this parameter throughout the experiment (Figure 2c) was similar to the one measured in the bacterial biomass, which suggests that the bacterial community is responsible for most of the organic matter content in the samples. However, there were no significant differences between the treatments and the controls (Table 3).

**Total respiratory activity**

The total respiration values, as measured by the electron transport system, showed a mean value of 20.41 (±19.51) µg formazan cm⁻² h⁻¹ in the controls, of 16.67 (±10.61) µg formazan cm⁻² h⁻¹ in the LOW channels and of 12.57 (±8.01) µg formazan cm⁻² h⁻¹ in the HIGH channels (Figure 2d). The response in this variable was dependant on the sampling time (significant Treatment*Date interaction in Table 3), being significantly higher in the controls in the first two weeks (one-way ANOVA, p < 0.05).

**Total carbohydrates**

The concentration of total carbohydrates had a mean value of 1329 (±1662) µgC cm⁻² in the controls, of 1526 (±1799) µgC cm⁻² in the LOW channels and of 1535 (±1797) µgC cm⁻² in the HIGH channels. Although the total carbohydrate concentration was not significantly different between each treatment (Table 3), attributed to the large variability of the parameter throughout the experiment (η² = 0.79), their concentration increased by the final two weeks in all channels (Figure 2e).

**Table 3.** Repeated measures ANOVA summary results for the biological variables examined in the experiment, considering two factors: Treatment (Control, Low, High) and Date (1 through 6). Measures of the biological effect are also shown (η²), and significant differences are highlighted in bold (p < 0.05).

| Source of variation               | Treatment | Date | Treatment x Date |
|-----------------------------------|-----------|------|------------------|
|                                   | p         | η²   | p               | η²   | p          | η²   |
| **Bacterial biomass**             | 0.90      | 0.01 | **0.01**         | 0.64 | 0.92       | 0.03 |
| Chlorophyll-a                     | 0.29      | 0.09 | 0.08             | 0.30 | 0.25       | 0.30 |
| Ash-free dry weight               | 0.27      | 0.10 | **0.04**         | 0.46 | 0.25       | 0.32 |
| Total respiratory activity        | 0.30      | 0.18 | **0.03**         | 0.30 | **0.01**   | 0.69 |
| Carbohydrates                     | 0.90      | 0.01 | **0.01**         | 0.79 | 0.94       | 0.02 |

**Autotrophs**

|                  | Treatment | Date | Treatment x Date |
|------------------|-----------|------|------------------|
| Diatoms          | 0.71      | 0.14 | 0.24             | 0.14 | 0.54       | 0.13 |
| Chlorophytes     | 0.86      | 0.02 | 0.63             | 0.04 | 0.46       | 0.17 |
| Euglenophytes    | 0.06      | 0.11 | 0.49             | 0.10 | 0.33       | 0.29 |
| Cyanophytes      | 0.52      | 0.05 | 0.09             | 0.30 | 0.66       | 0.12 |

**Consumers**

|                  | Treatment | Date | Treatment x Date |
|------------------|-----------|------|------------------|
| Nematods         | 0.22      | 0.16 | 0.33             | 0.12 | 0.34       | 0.23 |
| Cladocerans      | 0.62      | 0.02 | 0.26             | 0.18 | 0.29       | 0.29 |
| Rotifers         | 0.55      | 0.04 | 0.43             | 0.10 | 0.35       | 0.25 |
| Protozoans       | 0.17      | 0.09 | 0.67             | 0.06 | 0.48       | 0.21 |

**Community composition**

The most represented autotrophs throughout the experiment in all channels were diatoms (Figure 3), mainly *Surirella linearis* Smith, *Denticula elegans* Kützing, *Navicula erifuga* Lange-Bertalot, *Nitzschia frustulum* (Kützing) Grunow var. *frustulum*, and *Placoneis clementis* (Grunow) Cox.

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In abundance, the diatoms were followed by cyanophytes, mainly \textit{Oscillatoria tenuis} Agardh \textit{ex} Gomont, and chlorophytes, mainly \textit{Coelastrum microporum} Nägeli \textit{in} A. Braun, \textit{Scenedesmus opoliensis} Richter and \textit{Pediastrum duplex} Meyen. Euglenophytes, although scarce, were represented by \textit{Euglena acus} (Müller) Ehrenberg and species of the genus \textit{Phacus} Ehrenberg. The abundance of the different algal groups did not exhibit significant differences between the controls and the treatments (Table 3). Among the consumers, the nematods were the most abundant in all

**Figure 2.** Variations in the biological variables measured throughout the experiment: (a) Bacterial biomass; (b) Chlorophyll-\(a\) concentration; (c) AFDW; (d) Total respiration; (e) Total carbohydrates. Bars indicate standard deviation.
Figure 3. Total densities of autotrophs (left panels) and consumers (right panels) in the controls, LOW and HIGH treatments, represented as mean values (point), standard error (boxes) and standard deviation (bars).
channels, followed by cladocerans and protozoans (ciliates and testate amoebeae), while rotiferans were the least abundant (Figure 3). However, no significant differences were found in these groups due to the treatments (Table 3).

The ANOSIM results showed significant differences between the controls and the LOW treatment (p = 0.04) and between the controls and the HIGH treatment (p = 0.03) when considering the species composition. The average dissimilarity between the controls and the LOW treatment was slightly lower (71.9%) than the dissimilarity between the controls and the HIGH treatment (73.2%), and the taxa that contributed more to the dissimilarity between the groups belonged to the autotrophs, particularly diatoms and cyanobacteria (Table 4).

### Table 4. Percentage of dissimilitude between the controls and both treatments, and percentage of contribution of the species that contributed most to the dissimilarity between treatments (Control vs. Low treatment and Control vs. High treatment) as expressed by the SIMPER results.

| Species                        | Control vs. LOW (% dissimilitude) | Control vs. HIGH (% dissimilitude) |
|--------------------------------|-----------------------------------|-----------------------------------|
| Oscillatoria tenuis Agardh ex Gomont | 12                                | 17                                |
| Nitzschia palea (Kützing) Smith | 7                                 | 8                                 |
| Ulnaria ulna (Nitzsch.) Compère | 8                                 | 12                                |
| Nitzschia linearis Smith | 6                                 | 13                                |
| Diadesmis converfacea Kützing | 6                                 | 13                                |
| Sellaphora seminulum (Grunow) Mann | 2                                | 8                                 |
| Placoneis placenta (Ehrenberg) Heinzerling | 4                                | 8                                 |
| Nitzschia amphibia Grunow | 2                                 | 3                                 |
| Calloneis bacillum (Grunow) Cleve | 2                                | 3                                 |
| Planothidium lanceolatum Lange-Bertalot | 2                                | 3                                 |

### Discussion

The results from this study show that only the specific composition of the autotrophic assemblage changed in the analyzed time period, as indicated by increase in the dissimilitude between the control and treatments. Although the structural parameters of biofilms are usually last to be affected by changes in the environment, the species’ composition within the biofilm was significantly impacted by the treatments, reaching dissimilitude values of 71.9% due to the low impact treatment, and 73.2% due to the more intense treatment. Density variations in species such as Oscillatoria tenuis, Nitzschia palea, Ulnaria ulna, Nitzschia linearis, Diadesmis converfacea, Sellaphora seminulum and Placoneis placenta, which proliferate in eutrophic conditions (Palmer 1969; Van Dam et al., 1994; Licursi et al., 2016), contributed the most to the variability between the controls and both treatments.

These changes led to a modification in the autotrophic assemblage, with rises in species most tolerant to a worse habitat and water quality condition, in only a few weeks of the experiment. Similar changes were recorded in a field nutrient addition in a Pampean stream (Artigas et al., 2013; Cochero et al., 2013), where the basal phosphorous levels were tripled, and rapid occurring changes were observed in the algal assemblage, particularly in the proportion of diatoms as a consequence of the input of nutrients to the system (Licursi et al., 2016), while the bacterial biomass and its metabolic activity took longer periods of time to be significantly affected (Cochero et al., 2013).

Although it has been reported that the co-occurrence of the increments in temperature and nutrients lead to metabolic and structural responses in epilithic
biofilms in short exposition periods (60 days) (Villanueva et al., 2011), and similar responses are caused by nutrients and water current velocity (Horner and Welch, 1981), the changes in the total respiratory activity in this experiment, with an epipelic biofilm exposed to a greater number of stressors, were not evident immediately. The dependence of bacteria for autochthonous organic matter in Pampean streams (Cochero et al., 2013) suggests that this heterotrophic community needs longer exposition times to show significant changes in the biofilm. On the other hand, the effect of nutrients alone can be less predictable in ecosystems with higher basal nutrient levels, such as Pampean streams. Although changes are slow, it is possible that the chronic input of nutrients will affect the functioning and the services that these water bodies provide (Artigas et al., 2013). Our results show that the epipelic biofilms subjected to the co-occurrence of several constant stress factors reveal that the specific composition of the autotroph assemblage responds sensitively at short term exposures, in comparison with the other variables analyzed.

Pampean streams are subject to a growing demographic pressure, and the expansion of cultivated land along with the effects of global changes might have significant effects in temperature and rainfall patterns that influence these lotic systems. Therefore, the combination of studies with various temporal and spatial scales can provide with a more precise comprehension of the factors that determine the development and dynamics of these epipellic communities. Studies that include the effects of multiple stressors in freshwater communities, such as those carried out by Ormerod et al. (2010), Proia (2012), Piggott et al. (2012), Lange et al. (2014), Piggott et al. (2015), require a more intricate and complex experimental design than the one employed in this article, to be able to identify the individual contribution of each environmental variable. However, the results shown in this article increase the current knowledge of the overall effects of co-occurring variables on epipelic biofilms of these nutrient rich streams, and highlights the sensitivity of the specific composition to measure rapid changes in water quality.

Conclusion

In summary, this study shows that the responses of nutrient-rich biofilms to the effects of co-occurring physical-chemical changes are not easily predictable, since they do not necessarily coincide with the results obtained in the large variety of studies that modified the same physical-chemical variables in an individual manner. This could be the result of the inherent resistance of nutrient-rich biofilms to stressors, or could be a consequence of an antagonistic interaction between the manipulated variables.

However the case, more research that includes the effects of multiple stressors is needed to understand how the biofilms respond in natural settings, where the variations of a single variable due to either environmental or human-induced factors are unusual. Also, these results highlight the sensitivity of community-based bioindicators, such as species composition, to environmental changes at short term experiments.

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Conflict of interest statement

Authors declare that they have no conflict of interests.

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