Communication

Benthic Metabolism in Fluvial Sediments with Larvae of Lampetra sp.

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Abstract: Lampreys spend their larval stage within fine sand fluvial sediments, where they burrow and act as filter feeders. Lamprey larvae (ammocoetes) can significantly affect benthic-pelagic coupling and nutrient cycling in rivers, due to high densities. However, their bioturbation, feeding and excretion activities are still poorly explored. These aspects were investigated by means of laboratory incubations of intact sediments added with ammocoetes and of animals alone. Oxygen respiration, nutrient fluxes and excretion rates were determined. Individual ammocoete incubations suggested that biomass-specific oxygen consumption and ammonium, reactive phosphorus and silica excretion were size-dependent, and greater in small compared to large individuals. The comparison of ammocoetes metabolic rates with rates measured in intact sediments revealed that ammocoetes activity decreases significantly when they are burrowed in sediments. Furthermore, results suggest that a major fraction of ammonium excreted by ammocoetes was assimilated by benthic microbes or microalgae to overcome in situ N-limitation. Alternatively, part of the excreted ammonium was oxidized and denitrified within sediments, as nitrate uptake rather increased along with ammocoetes density. Ammocoetes excreted reactive phosphorus and silica but such production was not apparent in bioturbated sediments, likely due to microbial or microalgal uptake or to immobilization in sediments.

Keywords: ammocoetes; sediments; bioturbation; metabolism; fluxes; engineering species

1. Introduction

Sediment bioturbation is an extensively studied subject, both in marine and freshwater systems [1–5]. It consists of a variety of processes affecting sediment physical structure and biogeochemistry and is grouped into two main actions: the reworking of particles and the ventilation of burrows [2]. The reworking of particles occurs during burrowing activities and includes the mixing of old and recent organic matter, the ingestion of sediments and the production of biodeposits. All these processes may determine a priming effect on the mineralization of sedimentary organic matter pools, in particular of their refractory fractions. This results in nutrient recycling and contrasts their burial. Burrows ventilation introduces oxygen (O₂) and sometimes nitrate (NO₃⁻)-rich water within sediments, at depths where these electron acceptors are generally depleted [6,7]. Oxygen and NO₃⁻ diffuse through burrow walls and create oxidized niches, producing multiple biogeochemical consequences, such as the oxidation of chemically reduced compounds (e.g., metals, sulphides or methane), nitrogen (N) removal via coupled nitrification and denitrification and phosphorus (P) precipitation [8,9].
Microbioturbation by small organisms as meiofauna (e.g., organisms with body size < 1 mm) was demonstrated to increase organic matter mineralization and CO₂ fluxes [10] and N removal via denitrification [11]. However, most bioturbation studies have addressed the role of macrofauna (e.g., organisms with body size > 1 mm); benthic macrofaunal communities consist of organisms characterized by different functional groups, each colonizing different sediment horizons, with specific bioturbation mode and feeding strategy [12]. Bioturbation by macrofauna was demonstrated to produce important effects on benthic metabolism, significantly enhancing mineralization rates and benthic-pelagic coupling or facilitating the growth of primary producers [13,14]. Bioturbation is performed also by large vertebrate organisms as fish when they resuspend sediments during feeding, hiding or reproduction [3,4]. Cyprinids, for example, are known to feed on sediment macrofauna as snails and chironomids and determine particles resuspension and nutrient mobilization in the water column [15]. Egg hiders like salmonids or lampreys, during the gravel nest digging, may significantly affect bottom properties [4,16,17]. Having very close sympatric distribution in rivers and streams of the Northern Hemisphere, salmonids and lampreys with their coarse-substrate bioturbation may produce profound ecological effects on their reproduction conditions [18] and on the stream ecosystems [4,16].

Lampreys are acknowledged as burrowing ecosystem engineers, not only at the adult spawner stage [16], but also during the whole larval stage [19,20], which is very unique within vertebrates. Lampreys have a 3–8 year larval stage in between the free embryonic and metamorphosis phases during which they are called ammocoetes and spend most of their time burrowed in the soft sediments of streams and rivers [21]. Ammocoetes burrow by inserting their head into the sediments with vigorous contractions of the tail [22]. Elective substrates are made of fine sands or coarse organic matter, allowing burrow construction, water exchange and protection against predators [23,24]. Ammocoetes feed on particulate matter suspended in the water column. They are thus able to circulate the water through their mouth and perform filter-feeding activity. This is a very specific behavior of a vertebrate: ammocoetes burrow as any polychaete and feed on suspended matter as mollusks. While in their burrow, larvae exploit water flow through their branchial chamber for the provision of food, respiratory needs and excretion of metabolic wastes [22]. Due to the length of their larval stage, good-quality burrowing substrates are important for the vitality of lamprey populations. The ammocoetes abundance generally displays positive correlation with soft sediment, organic matter, low flow and low depth, which characterize streams marginal habitats [25–27]. Lampreys larvae are able to affect via their bioturbation, the sediment physical structure, oxidation status and microbial communities of these marginal habitats [22,28]. Ammocoetes may also modify nutrient dynamics at the sediment–water interface during their burrowing phase, but the biogeochemical effects, extent and significance are still poorly studied [19,20]. The understanding of the effects produced by the ammocoetes during their burrowing phase is consequently of interest both in management-conservation and in benthic biogeochemistry-river functioning perspectives.

In this study we measured benthic processes in sediments added with lamprey larvae as a first assessment of their contribution to benthic respiration and nutrient cycling. We focused on the dissolved inorganic forms of nitrogen (NH₄⁺ and NO₃⁻), silica (SiO₂) and phosphorus (PO₄³⁻) as these chemical species represent the main nutrients that support the growth of phytoplankton and microphytobenthos. Any variation in their biogeochemical cycle, either from external pressures or resulting from the activity of the macrobiota, may produce deep impacts in the functioning of aquatic ecosystems. Ammocoetes respiration (O₂ consumption) and excretion rates (dissolved nitrogen, phosphorus and silica release) were also quantified to analyze the contribution of lamprey larvae metabolism to benthic processes. This study falls within investigations addressing the functional role of organisms that enhance nutrient recycling and benthic-pelagic coupling and might have an important role in sustaining primary and secondary production, and in general river functioning [29]. Such an investigation is urgent as lamprey larvae abundance is declining in many areas,
and the net loss of ecosystem functions associated to the disappearance of these organisms is unknown.

Deep alterations of riverine ecosystems, including water abstraction, discharge regulation, hydropeaking, dredging, impoundment, embankment and eutrophication have decreased larval habitats, and impacted lamprey populations [25,30,31]. The ecological niche of lampreys, and of ammocoetes in particular, is still poorly explored and more information on the interactions between larvae and river benthic and pelagic compartments is needed to understand the biogeochemical role of these organisms and to improve the recovery of lamprey populations.

2. Materials and Methods

2.1. Sampling Procedure and Microcosms Set-Up

Ammocoetes of Lampetra sp. were collected in July 2015 from the River Šventoji located in the region of western Lithuania (56°03′ N, 21°11′ E) (Figure 1). At the sampling site, sediments were colonized by a variety of submerged, floating-leaved and emergent macrophytes; but large patches devoid of vegetation were also found. The width of the wet section at the study site was 5–6 m, the maximum depth was 1 m and the average water velocity of the whole water column, measured at several transects with a current meter (Flow Probe FP101, Global Water Instrumentation, College Station, TX, USA) was 0.1 ± 0.1 m s⁻¹. The river water was yellowish, likely due to elevated concentrations of dissolved organic matter, whereas dissolved inorganic nutrient concentrations were low, with silicon (Si) in excess to N and P. Ammonium and NO₃⁻ concentrations averaged 3.1 ± 0.9 and 6.4 ± 1.7 µM, respectively. Soluble reactive phosphorus and SiO₂ averaged 1.2 ± 0.1 and 76.5 ± 3.4 µM, respectively. Water temperature and electrical conductivity were measured by an YSI multiple probe (Pro 1030, YSI, Yellow Springs, OH, USA).

![Figure 1](image-url)

Figure 1. The figure depicts the location of the sampling site, indicated with the asterisk (a), a detail of the river course with floating-leaved vegetation (b), Lampetra sp. larvae retrieved from sieved sediment (c), a burrowed ammocoete (d), an intact sediment core (e), a sediment cores incubation tank with stirring units and temperature control system (f) and a bottle with glass beads used for ammocoetes individual incubation (g).

Lampreys are of conservation concern in Europe and are listed on Annex II of the European Union Habitats Directive (92/43/EEC) and collection of any Lampetra sp. ammocoete is prohibited. Sampling of lamprey larvae for our study was carried out with a special fishing permit, in the framework of a fish communities sampling program. Ammocoetes
were collected from shallow marginal areas without vegetation (depth 0.2–0.3 m, low to undetectable water velocity) from muddy sand oxidized sediments. To this purpose, surface sediment layers (0–10 cm depth) were collected with squared plastic frames exploring areas of 300 cm$^2$ and sieved with 1 mm sieve in order to retrieve larvae. Alive lamprey larvae were transferred into buckets filled with river water. In addition, 32 intact sediment cores (i.d. 8 cm, length 30 cm) were collected by hand from the adjacent undisturbed area in order to have nearly 10 cm of sediments and 20 cm of clear water phase. Three smaller intact cores (i.d. 3.5 cm, length 25 cm) were collected for general characterization of sediment and ~200 L of in situ river water was collected for ammocoetes bioturbated sediment preincubation and incubation procedures.

2.2. Measurement of Ammocoetes Respiration and Excretion Rates

Single ammocoetes collected from the river were selected according to their body size in order to cover a wide range of lengths (from 4 to 10 cm, $n = 25$). In the laboratory, they were gently transferred to 100 mL BOD (biological oxygen demand) glass vials containing sterilized glass beads to reproduce a sedimentary environment and 0.22 µm-filtered river water. Vials with ammocoetes were incubated in the dark at in situ temperature (18 °C) (Figure 1g). At the beginning and at the end of the incubation ~60 mL of water was collected with plastic syringes from the incubation tank ($t_0$) and from the bottles ($t_f$). Initial and final water samples were dispensed and treated for different purposes. An aliquot of 30 mL was transferred to 12-mL Exetainer (Labco Scientific, Lampeter, UK) flushing the volume twice for dissolved O$_2$ analysis and immediately preserved with 50 µL of 4% HgCl$_2$. Another aliquot of 30 mL was filtered (GF/F filters) and transferred to scintillation vials for inorganic nutrient analyses (NH$_4^+$, NO$_3^-$, PO$_4^{3-}$ and SiO$_2$). Dissolved inorganic nutrients were measured within 12 h (see Section 2.4). Incubations lasted from 0.5 to 1.5 h in order to keep final O$_2$ concentrations within 30% of the initial value. At the end of the incubations, test animals were euthanized overdosing anesthetic 2-phenoxethynol, their length was measured and the dry weight (DW) of each ammocoete was measured after drying at 60 °C to a constant weight. Biomass specific ammocoetes respiration and excretion rates ($\mu$mol g$_{DW}^{-1}$h$^{-1}$) were calculated according to Equation (1):

$$\text{Flux} = \frac{[(C_f - C_i) \times V]}{t \times B},$$

where $C_f$ and $C_i$ are the final and initial concentrations of the target solute (µM), $V$ is the volume of the BOD glass vials corrected for the volume of glass beads (L), $t$ is the incubation time (h) and $B$ is the ammocoetes biomass (g$_{DW}$).

2.3. Measurements of Benthic Fluxes in Lamprey Larvae Bioturbated Sediments

In the laboratory, all cores were maintained at the same temperature of the natural environment (18 °C) submerged with the top open in temperature-controlled incubation tanks containing aerated river water (Figure 1f). Then, variable numbers of ammocoetes were added to the cores (from 0 to a maximum of 4 individuals per core) in order to study the effect of their biomass on benthic processes. Once added, all lampreys were rapidly digging in surface sediments, creating with their body U-shaped burrows, with head and tail at the sediment–water interface and the body entirely hidden within the sediments. Thereafter sediments with lampreys were pre-incubated overnight in the tanks. The day after, dissolved O$_2$ and inorganic nutrient fluxes were measured in the dark according to a start-end standard incubation procedure [32]. Before starting the incubation, the water in the tanks was renewed with fresh in situ water. At the beginning of the measurements, water samples (ca. 100 mL, $n = 5$, $t_0$) were collected from each tank with plastic syringes and all cores were closed with rubber stoppers. An incubation period of 4–5 h was set in order to keep the final O$_2$ concentration within 20–30% of the initial value. At the end of the incubation, a water sample (ca. 100 mL, $t_f$) was collected from each core. All samples were
processed as described in Section 2.2. Net fluxes of all measured solutes (µmol m\(^{-2}\)h\(^{-1}\)) across the sediment–water interface were calculated according to Equation (2):

\[
\text{Flux} = \frac{[(C_f - C_i) \times V]}{t \times A},
\]

where \(C_f\) and \(C_i\) are the final and initial concentrations of the target solute (µM), \(V\) is the volume of the water phase in the core (L), \(t\) is the incubation time (h) and \(A\) is the sediment surface in the core (m\(^2\)).

2.4. Sediment Characterization and Analytical Methods for Water Samples

Three small sediment cores were extruded and sliced for chlorophyll \(a\), organic matter and grain size analysis. Sediment chlorophyll \(a\) was measured spectrophotometrically in the upper 0.5 cm sediment layer according to [33], after 24 h extraction in 90% acetone. Organic carbon (C\(_{\text{org}}\)) was analyzed in dried sediments from the upper 0–2 cm layer with a Shimadzu total organic carbon (TOC) analyzer (Tokyo, Japan) equipped with a compartment for solid analysis. An aliquot of wet sediment was employed for sediment grain size analysis with a laser particle size analyzer (Analysette 22 MicroTec plus, Fritsch GmbH, Idar-Oberstein, Germany).

Dissolved O\(_2\) concentration was measured with a microelectrode (OX-50, Unisense A/S, Aarhus, Denmark). Dissolved inorganic nutrients (NO\(_3^-\), PO\(_4^{3-}\) and SiO\(_2\)) were measured with a continuous flow analyzer (San++, Skalar Analytical B.V., Breda, The Netherlands) using standard colorimetric methods [34]. Ammonium was analyzed manually by means of the salicylate-hypochlorite method, using nitroprussiate as a catalyst [35].

2.5. Statistical Analysis

Data on respiration and nutrient excretion were plotted against ammocoetes dry biomass and logarithmic regression curves were fitted where appropriate. Data on benthic respiration and nutrient fluxes were plotted against ammocoetes dry weight and analyzed via linear regression. The coefficient of determination and statistical significance are provided for the regression models. Significance level for all tests was \(p < 0.05\). All statistical analyses were performed via Sigma Plot 12.0 (Systat Software Inc., San Jose, CA, USA).

3. Results and Discussion

3.1. Sampling Site Features

Large portions of the Šventoji River marginal areas were covered by fine sand deposits (Md = 0.195 ± 0.012 mm), in which the silt fraction (<63 µm) contributed a small percentage of the total (16.6 ± 3.9% on average). The studied sediments are typical for most of Lithuanian lowland rivers marginal habitats occupied by lamprey larvae. In a prior study on habitat selection, lamprey larvae were demonstrated to be habitat specialists that prefer fine-sand sediments, and to a lesser extent organic-rich bottom environments [24,36]. Such selection is likely driven by much lower microbial activity and O\(_2\) consumption in organic-poor sediments. Lotic ecosystems generally do not display O\(_2\) shortage due to water flow, however shallow areas with low current velocity are sites of fine sediments accumulation where O\(_2\) demand and supply can be decoupled, resulting in local undersaturation. Another important aspect related to habitat selection by lamprey larvae is the build-up of anaerobic metabolism end-products (e.g., NH\(_4^+\), chemically reduced metals, sulphides) in sediments receiving elevated inputs of labile organic matter. These end-products may accumulate in pore water, and be toxic to ammocoetes. The sediments of Šventoji River marginal areas, where ammocoetes were collected, appeared light brown and quite oxidized along the upper 10–20 cm horizon, without vertical redox discontinuities (Figure 1e). Sediment oxidation may derive by low heterotrophic microbial activity at the study site. The latter is due to low organic matter content in sediments (C\(_{\text{org}}\) = 0.6 ± 0.2% of the sediment dry weight) and to the presence of coarse fragments of leaves from terrestrial
origin, that are generally refractory to mineralization due to their high cellulosic content. Such oxidation can also be a direct consequence of elevated bioturbation by lamprey larvae, favoring the oxidation of reduced chemical forms [19]. Experimental studies on ammocoetes behavior in reconstructed sediments demonstrated a high mobility of larvae within the upper 4–5 cm sediment horizon during the day and during the night, resulting in elevated bioturbation-induced oxidation and increased sediment redox [19,37]. Such experiments revealed that ammocoetes move inside the substrate without leaving it and that surface activity was observed rarely.

When ammocoetes dig within sediments they stabilize the fine sand particles of burrow walls by producing mucus, as demonstrated for other macrofauna groups [38]. Fine sediments are a specific requirement for ammocoetes as larger particles are more difficult to penetrate by larvae or to solidify with mucus and would result in burrow collapse. This may explain the large densities of ammocoetes at the study site, whereas elsewhere lampreys are declining in numbers due to specific habitat loss (e.g., increase of sedimentation and organic content) [31]. At the study site, water transparency and shallowness favored the growth of microphytobenthos and surface sediments displayed high content of chlorophyll \( a \), averaging 162.4 ± 55.7 mg Chl \( a \) m\(^{-2} \) (\( n = 3 \)). Benthic algae represent high quality organic matter at the sediment surface, and may constitute a food source for ammocoetes [39]. Nutrient concentrations in the Šventoji River water were low, and typical of the dry season and of lowland streams within forested catchments.

### 3.2. Size, Weight and Metabolic Activity of the Ammocoetes Retrieved in the Šventoji River

In the Šventoji River, lamprey larvae represent Lamproptera genus and could belong to both river (\( L. \) fluviatilis) and brook lamprey (\( L. \) planeri); however, the area is mostly used for river lamprey spawning and rearing habitat. It is very difficult to distinguish the two species during their larval stage due to rather similar morphology [31]; moreover, there are no clear differences between brook and river lamprey ammocoetes sediment preference, burrowing and feeding behavior. Sampled river-marginal area hosted a Lamproptera sp. population including all size classes (from 0+ sub-yearlings to 5 year old larvae). The length of larvae retrieved from sediments (\( n = 114 \)) varied between 11 and 105 mm (average 47 ± 16 mm), corresponding to a dry weight between 0.002 and 0.305 g\(_{\text{DW}}\) ind\(^{-1}\) (average 0.046 ± 0.053 g\(_{\text{DW}}\) ind\(^{-1}\)). However, the distribution of frequencies of body length (not shown) suggests that individuals with length between 40 and 60 mm represented nearly 50% of the retrieved animals, and animals with a size within that range were used in the sediment core incubations.

Lamprey larvae alone incubations revealed large differences, generally by one order of magnitude, of the biomass-normalized metabolic and excretion rates (Figure 2). Smaller individuals displayed higher metabolic and excretion activities than larger individuals, as reported in analogous studies on invertebrate macrofauna (e.g., the filter-feeding bivalves Dreissena polymorpha or Sinanodonta woodiana) [32,40]. Oxygen respiration rates varied between 20 and 200 \( \mu \text{mol} \) \( \text{O}_2 \) g\(_{\text{DW}}\)\(^{-1}\)h\(^{-1}\) (Figure 2a). Ammocoetes displayed lower metabolic activity and lower excretion rates when compared to teleosts of similar size, likely due to the environment where they spend most of their time (i.e., the sediment, which is a hostile environment for \( \text{O}_2 \) exigent organisms) [41]. However, ammocoetes respiration rates aligned with ranges reported for other bioturbators (e.g., 10–260 \( \mu \text{mol} \) \( \text{O}_2 \) g\(_{\text{DW}}\)\(^{-1}\)h\(^{-1}\) for different bivalves and 90–140 \( \mu \text{mol} \) \( \text{O}_2 \) g\(_{\text{DW}}\)\(^{-1}\)h\(^{-1}\) for burrowing chironomid larvae) [40,42]. Rates of \( \text{NH}_4^+ \) excretion (~2 to ~13 \( \mu \text{mol} \) \( \text{NH}_4^+ \) g\(_{\text{DW}}\)\(^{-1}\)h\(^{-1}\)) were one order of magnitude lower than the absolute values of \( \text{O}_2 \) consumption but displayed a similar trend with much higher excretion measured in smaller organisms (Figure 2b). Similar results are reported also in [32,40]. Ammocoetes mostly released \( \text{NH}_4^+ \) to the water but different authors report that up to 15–20% of excreted \( \text{N} \) is in the form of urea, not quantified in the present study [43]. During larvae alone incubations, net accumulation of \( \text{NO}_3^- \) in the water was also measured (Figure 2c). However, ammocoetes-related \( \text{NO}_3^- \) fluxes were generally lower than 2 \( \mu \text{mol} \) \( \text{NO}_3^- \) g\(_{\text{DW}}\)\(^{-1}\)h\(^{-1}\), and did not exhibit a clear
size-dependent trend. Such NO$_3^-$ could be produced via nitrification by bacteria colonizing the skin of ammocoetes, oxidizing part of the excreted NH$_4^+$, as reported for other burrowing organisms [44]. The absolute value of the molar ratio between O$_2$ and NH$_4^+$ fluxes measured in incubations of individual ammocoetes averaged 17.1 ± 4.1, whereas the O$_2$ to NH$_4^+$+NO$_3^-$ molar ratio averaged 13.8 ± 2.9. Such ratios, that do not consider urea excretion, can be considered as proxies of the C:N ratio of the ammocoetes food particles and suggest high quality, N-rich organic matter as phytoplankton, microphytobenthos or labile organic detritus as food source [39]. Excreted phosphorus is generally associated with feces, but high fluxes of PO$_4^{3-}$ from ammocoetes were also measured (0.4 to 2.4 µmol PO$_4^{3-}$/gDW$^{-1}$h$^{-1}$) (Figure 2d). To our knowledge, this is the first time that PO$_4^{3-}$ excretion was reported for ammocoetes, as most studies considered only N compounds. The molar ratio between NH$_4^+$ and PO$_4^{3-}$ release was surprisingly low and averaged 8.2 ± 3.7. This may suggest that part of the ammocoetes excreted N, rather than P, is associated to biodeposits. Ammocoetes alone incubations revealed also variable release of SiO$_2$, from <1 to 25 µmol SiO$_2$/gDW$^{-1}$h$^{-1}$ (Figure 2e). Such release falls within the range reported for filter-feeding bivalves [32,40]. In some of the incubated ammocoetes, the NH$_4^+$ to SiO$_2$ excretion ratio was close to unity. Silica excretion supports studies reporting diatoms as a food source of ammocoetes [39]. Reactive P and SiO$_2$ excretion displayed a tendency towards higher rates in smaller organisms, as shown for O$_2$ and NH$_4^+$, but with a larger degree of variability.

![Graphs showing relationships between ammocoetes biomass and various nutrient fluxes](image)

**Figure 2.** Biomass-normalized rates of oxygen respiration (a), nutrient excretion (b,d,e) by ammocoetes and NO$_3^-$ production (c) plotted as a function of their individual biomass. All rates were determined via incubations of ammocoetes of variable size (n = 25) in 100 mL glass bottles provided with an artificial substrate (glass beads).

### 3.3. Benthic Respiration and Nutrient Fluxes in Sediments Bioturbated by Ammocoetes

Lamprey density in incubated cores varied between 0 and 600 ind m$^{-2}$ and the average larvae biomass added to sediments was 0.028 ± 0.008 g$_{DW}$m$^{-2}$. Through their respiration and burrowing activity, lampreys enhanced benthic O$_2$ and NO$_3^-$ consumption and NH$_4^+$ excretion (Figure 3a–c), whereas PO$_4^{3-}$ and SiO$_2$ fluxes tended to increase, but slopes were not significantly different from zero (Figure 3d,e).
Figure 3. Total oxygen uptake (a) and net nutrient fluxes (b–e) measured in sediments with different biomass of ammocoetes \( (n = 32) \). Graphs report also the linear regressions between process rates and ammocoetes biomass. Slopes were significant for total oxygen uptake (TOU) \( (p < 0.0001) \), \( \text{NH}_4^+ \) \( (p = 0.002) \), \( \text{NO}_3^- \) \( (p < 0.0001) \), \( \text{PO}_4^{3-} \) \( (p = 0.340) \) and \( \text{SiO}_2 \) \( (p = 0.100) \) fluxes.

Similar stimulation of total \( O_2 \) uptake in sediments with ammocoetes was demonstrated in macrofauna bioturbation studies \([6,8,14,40]\). Burrowing organisms increase benthic \( O_2 \) consumption due to their metabolic activity and to the subsurface burrow structures, that expand the surface through which dissolved \( O_2 \) can diffuse. Ammocoetes do not construct deep burrows and it is likely that most of the increase of benthic \( O_2 \) demand is due to the ammocoetes respiration. What is known is that ammocoetes frequently change their position within sediments, so they dig frequently, but it is not clear whether they perform a ventilation of their burrows. From visual observations this is unlikely, as they feed directly from the water, taking advantage of continuous water renewal in lotic ecosystems. At 18 °C the respiration of bare sediments (without ammocoetes) was 1253 \( \mu \text{mol} \text{ m}^{-2} \text{ h}^{-1} \), which is expected for sandy sediments with low organic matter content as at the study site and as reported in similar substrates \([8]\). Such respiration increased by \( \sim 35 \mu \text{mol} \text{ O}_2 \text{ m}^{-2} \text{ h}^{-1} \) per gram dry weight of ammocoetes (Figure 3a). Such value falls within the range of biomass-normalized ammocoetes respiration rates, and suggests that most of the increase of benthic \( O_2 \) consumption in sediments with ammocoetes was due to the larvae respiratory needs. In a similar experiment, the addition of the oligochaete \textit{Sparganophilus tamesis} to muddy and sandy sediment determined an increase of 26 and 17 \( \mu \text{mol} \text{ O}_2 \text{ m}^{-2} \text{ g}_{\text{DW}}^{-1} \text{ h}^{-1} \) whereas large individuals of \textit{S. woodiana} stimulated \( O_2 \) demand by less than 10 \( \mu \text{mol} \text{ O}_2 \text{ m}^{-2} \text{ g}_{\text{DW}}^{-1} \text{ h}^{-1} \) \([8,40]\). Single individual incubations revealed large \( \text{NH}_4^+ \) excretion, which was substantially reduced when ammocoetes were in sediments, likely due to benthic assimilation (by bacteria or microphytobenthos) or nitrification in the aerobic sediment volume. Increasing ammocoetes biomass resulted in more negative \( \text{NO}_3^- \) fluxes, supporting what is generally found with burrowing organisms via isotope-based techniques: an increase of denitrification rates \([6]\). Similarly, high rates of \( P \) and \( \text{Si} \) excretion did not result in increased fluxes in lamprey larvae bioturbated sediment incubations, likely due to coupled retention (e.g., via dark assimilation or co-precipitation) of these nutrients in the sandy sediments of the Šventoji River. Overall, the combination of ammocoetes alone and bioturbated sediment incubations suggest that lamprey larvae retain high quality particles via ingestion and recycle large quantities of nutrients that do not accumulate in
the water column but are likely reused within sediment by microbial community or on the sediment surface by microphytobenthos.

4. Conclusions

Taken together, results from this experimental study support previous findings that lamprey larvae are ecosystem engineers capable via their own metabolism and burrowing activities to affect sediments physical, chemical and biological features. The impact of an ecosystem engineer on the environment is a function of three main biological aspects: its behavior, body size, and population density, all mediated by the abiotic conditions [4]. Following this concept, Lampestra ammocoetes seem to be influential bioturbators, because: (a) they are vigorous burrowers, actively bioturbating and displacing sediments; (b) lamprey community consists of several year cohorts, with the biggest individuals up to 7–10 cm; and (c) the density in a healthy system could reach high numbers, as in Šventoji River case, and as tested in our experimental setup. Larval densities have been determined by a number of researchers and found to be very variable, both from place to place and from year to year in the same place [31]. In some Lithuanian rivers, proposed as NATURA 2000 sites for lampreys, Lampestra sp. density ranged from a few ind. m$^{-2}$ in large rivers to density up to 46 ind m$^{-2}$ in Šventoji River [45].

We demonstrated the important role of lamprey larvae as regulators of nutrient cycling in riverine systems, coupling pelagic production with the benthic compartment and taking an active role in nutrient spiraling. This role is comparable to that of filter-feeding mollusks, that in pristine lotic ecosystems, via filtration and excretion, were promoting the recycling of otherwise limiting nutrients. Besides excreting all the key nutrients and enriching sediments with biodeposits, lamprey larvae actively bioturbated sediments, improving their chemical quality (e.g., increasing O$_2$ and NO$_3^-$ penetration and pore water redox potential), augmenting the communities of aerobic bacteria and mixing refractory and labile particles, producing a stimulatory effect on organic matter microbial mineralization. As such, ammocoetes act as burrowing macrofauna, even if they do not assume a static feeding position in the substrate. Contrarily to burrowing fauna, ammocoetes do not consolidate and ventilate their burrows creating biogeochemical hotspots, rather they seem to produce a much more diffuse effect on surface sediments. Incubation of single ammocoetes revealed elevated excretion of inorganic N, Si and P, but such fluxes were attenuated when ammocoetes were buried. Frequent movements of ammocoetes in sediments result in particles reworking and in multiple injections of oxic or nitrate-rich water in sediments, potentially affecting pore water chemistry and microbial community composition. Such movements might also favor the retention of excreted nutrients in microbial biomass within nutrient-poor sandy sediments. Ammocoetes might therefore increase the nutritional quality of sediments, with cascade implications for different trophic levels, likely sustaining the heterotrophic microbial communities, benthic primary producers, meio- and macrofauna [5,46]. Even more, ammocoetes tend to drift downstream with their aging [21,31], displacing the biogeochemical services they provide from upstream spawning sections to downstream reaches with low or no local ammocoetes recruitment.

Further investigations, combining molecular and biogeochemical techniques may reveal hidden and interactive associations between ammocoetes and microbial community, which are important for the functioning of lotic ecosystems. Such investigations should also expand the sediment typologies of marginal areas where ammocoetes burrow to allow generalization of the presented results.

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