Interaction among bacterioplankton and macrophytes in shallow lakes with high macrophyte cover

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Received: 17 January 2020 / Accepted: 27 August 2020 / Published online: 5 September 2020
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Abstract
Growth of submerged and emergent macrophytes was studied together with heterotrophic bacterioplankton abundance and production in two Hungarian shallow lakes with dominant macrophyte covers. It was expected that bacterioplankton numbers and activity would have an effect on macrophyte biomass accumulation. Bacterial production and abundance showed a strong seasonal pattern with maximum in the warmest months (July, August). It was found that macrophyte biomass increased with heterotrophic bacterial production and abundance up to 5.6 µg C l⁻¹ h⁻¹ and 5.30*10⁶ cells, respectively, while over that value was negatively associated with macrophyte growth. It was also shown that the relationship between heterotrophic bacteria and macrophytes also varied seasonally, showing a multifaceted relationship. It was demonstrated that macrophytes are not only the most significant carbon and energy source for the bacteria in shallow, macrophyte-dominated lakes, but are also competing organisms that could be supressed by excessive bacterial activity. These findings could help better understand the interaction between macrophytes and bacterioplankton, and assist wetland managers in quantifying what may be a primary cause of reed die-back.

Keywords Bacterioplankton · Macrophytes · Activity · Abundance · Litter · Biomass accumulation

Introduction
Although wetlands and littoral zones cover less than 7% of the Earth’s land surface, they are the most important natural carbon reserves (Reddy and DeLaune 2008) one of the most productive zones globally, as well as sites of intense decomposition, especially of organic carbon (Richey et al. 2002; Duarte and Prairie 2005). These areas are therefore crucial, not only to the freshwater water bodies with which they are associated, but also to connected terrestrial landscapes (Odum 1956; Schindler and Scheuerell 2002).

Macrophytes simultaneously integrate the effects of and changes to several environmental factors (Lacoul and Freedman 2006). Rooted macrophytes, via their roots and rhizomes, are affected by the sediment in a wide variety of ways (Barko et al. 1991; Armstrong et al. 19962006). Some such interactions are universal, while some are specific to the aquatic environment. Plant growth is directly affected by nutrient availability and uptake (Barko and Smart 1986; Lacoul and Freedman 2006; Tóth et al. 2017), and the growth and distribution of macrophytes are also affected by the different chemical and physical characteristics of the sediment (Armstrong and Armstrong 1999; Koch 2001; Tóth 2016). The organic matter content of the sediment influences not only the growth and distribution of individual plants (Barko and Smart 1983; Koch 2001), but also the species composition of the community, and even successional processes within the ecosystem (Carpenter 1981).

The organic carbon of wetlands and littoral zones is an important part of aquatic ecosystem metabolism (Wetzel 1995) and both periphytic and planktonic heterotrophic bacteria assimilate a large portion of it for both energetic and structural purposes (Azam et al. 1983, 1994; Findlay et al. 1986). In small, shallow lakes with large littoral zones, the importance of autochthonous dissolved organic matter originating from macrophytes is high (Jonsson et al. 2001; Cole et al. 2002), moreover higher heterotrophic bacterioplankton...
activities were reported from lakes with high macrophyte cover (Reitner et al. 1999; Jeppesen et al. 2002; Rooney and Kalff 2003). The distribution of dead plant biomass (litter) can therefore directly and indirectly influence microorganisms and, as a result, environmental conditions (Tranvik et al. 2009).

Macrophytes directly affect their environment. Among other things, the availability of litter increases the amount of organisms participating in the decomposition of macrophyte biomass (Findlay et al. 1986; Reitner et al. 1999; Jeppesen et al. 2002; Rooney and Kalff 2003), which consequently influences larger ecosystem processes. While it is believed that each lake has a specific set of bacteria typical to the given lake (Lindström 2000; Yannarell and Triplett 2005; Jones et al. 2009), the data also suggest that these specific or characteristic sets of bacteria come from bacterial groups with global distributions (Lindström et al. 2005; Schauer et al. 2005; Sommaruga and Casamayor 2009). These bacteria, which contribute to litter decomposition in freshwater lakes, play an outstanding role in the regulation and directing the processes within the aquatic ecosystem, and affect the water quality of the whole lake (Newton et al. 2011). The amount of nutrients (N and P) available determines the nature of the process of respiration and the associated terminal electron acceptor compound, and consequently the oxidation-reduction conditions of the environment: high carbon and low N and P conditions promote anaerobic processes, such as methanogenesis, sometimes resulting in extremely low oxidation-reduction potential (Laanbroek 1990; Reddy and DeLaune 2008; Middelburg and Levin 2009). Thus, bacterial activity also significantly affect the macrovegetation of wetlands and littoral zones via the decomposition of C.

Whereas the majority of studies suggest direct bacterial dependence on macrophytes (Findlay et al. 1986; Reitner et al. 1999; Jeppesen et al. 2002; Rooney and Kalff 2003), very little is known about the effects of bacterioplankton on macrophyte growth and biomass accumulation. In our work, we analysed the growth and biomass accumulation of submerged and emerged macrophyte stands in two shallow Hungarian lakes in relation to the amount and activity of bacterioplankton. We hypothesised (i) that bacterioplankton production affects the growth and biomass accumulation of macrophytes, and (ii) that the amount of non-autotrophic bacteria affects the growth of macrophytes and biomass accumulation.

Materials and methods

Site description

Lake Fertő (Neusiedler See) (Supplementary Fig. 1a) is a 315 km² soda lake situated on the border between Hungary and Austria. It is shallow (<1.4 m depth), turbulent, 36 km long, and 12 km wide at the widest part of the lake. While more than 55% of Lake Fertő is covered by macrovegetation, 64 km² (~85%) of the Hungarian part of the lake is covered by emergent macrophytes, predominantly Phragmites australis (Cav.) Trin. ex Steud. (Supplementary Fig. 1a). Beginning from the early 1980s, reed stands in the Hungarian part of Lake Fertő were affected by die-back. Approximately 30% of the Hungarian stands are now at different stages of the related degradation (Márkus et al. 2008; Márkus and Király 2012), while on the Austrian side of the lake, reed stand deterioration is at ~10% (Schmidt and Csaplovics 2010).

Lake Kolon is a ~6 km long, 27.3 km² lake that is almost fully covered with macrovegetation, with occasional openings at the surface, located in the middle of Hungary, in the Danube-Tisza interfluve (Supplementary Fig. 1b). The lake receives water from the discharge of a short local flow system on its eastern side and recharges another short gravity system to the west. The dominant chemical type of the lake is Ca–Mg–bicarbonate (Mádl-Szönyi and Tóth 2009), and its average water level is less than 2 m depth.

Two areas were chosen for sampling in Lake Fertő and two in Lake Kolon (Supplementary Fig. 1). In the southern part of the Lake Fertő, we used a sampling site (47°38’57.41″N, 16°43’44.81″E) within a reed stand and a sampling site in a relatively large (0.03 km²) opening covered with submerged macrophyte (47°39’51.61″N, 16°42’53.33″E). In Lake Kolon a sampling site within the reed stand (46°48’5.67″N, 19°20’9.65″E) and a site in a relatively large (0.05 km²) opening covered with submerged macrophyte (46°46’24.70″N, 19°20’24.79″E) were selected. Biomass and water sampling was performed on a monthly basis in 2015 and 2016.

Plant morphological measurements

Macrophytes from both emergent and submerged macrophyte-dominated sites were sampled (repeatedly harvested from areas near the plots) throughout the vegetation period (April–October) at least once a month. Due to plant high heterogeneity at sites, plants from 4 to 5 random 0.25 m² quadrants were collected, and samples were pooled, and considered a single observation. Submerged macrophytes were sorted into species, oven dried for a week at 65 °C and weighed, while emergent macrophytes were cut at the water level, sorted into species, the green and dry parts were oven dried separately for a week at 65 °C and weighed separately. Biometric measurements were performed only on green Phragmites plants; plant height was determined as the length from the cut surface to the tip of the top leaf or seed head, the diameter in the middle of the most basal internode of each cut reed stem was measured to the nearest 0.1 mm with a vernier calliper, and both green and dry leaves, as
well as the number of nodes of each plant, were counted. Stem, leaves, and seed heads of green plants were weighed separately after drying.

**Water sample analyses**

Total suspended matter (TSM) was measured gravimetrically after filtering the water through a previously dried and pre-weighed GF-5 fibreglass filter (nominal pore size = 0.4 µm), drying for 2 h at 105°C, then weighing again and subtracting the filter weight from the total weight. TSM concentration was calculated based on the predetermined volume of the filtered sample. The concentration of coloured dissolved organic matter (CDOM) was measured spectrophotometrically at 440 nm (Hitachi, U-2900) after filtering the water through a cellulose acetate filter (pore size 0.45 µm), and was expressed as Pt (platina) units (mg Pt l⁻¹) (Cuthbert and Del Giorgio 1992). Nitrogen forms (ammonium, nitrate, and urea) (Newell et al. 1967; Elliott and Porter 1971; Mackereth et al. 1978), total phosphorus (TP) (Menzel and Corwin 1965) and total soluble phosphorus (TSP) (Gales et al. 1966) were determined. Chlorophyll-α of the water column was determined from freshly collected samples. 1 L of sampled water was filtered through Whatman GF-5 filter papers, and pigments were analysed following extraction in 60 °C methanol and clarification by centrifugation (10 000 rpm for 10 min) (Iwamura et al. 1970).

**Bacterial production and count measurements**

Bacterial production (BP) was measured using the radiolabelled leucine L-[4,5-³h] incorporation method (Kirchman et al. 1985; Gasol 1999). Mixed leucine solution (radiolabelled leucine: normal leucine = 1:10) was added to fresh water samples in situ at varying concentrations (20 nM, 40 nM, 70 nM, 100 nM, and 200 nM) in triplicate. Incubation was performed at actual lake temperatures measured during sampling. The reaction was stopped with 50 v/v % trichloroacetic acid (TCA). TCA-killed controls were also used to correct for the abiotic uptake of labelled leucine.

Unincorporated radiolabelled leucine was separated by centrifugation. A scintillation cocktail was added to our pellet and, after one day, we measured the disintegration per minute of samples by TRI-CARB 2100TR Liquid Scintillation Analyzer. From this, we calculated the leucine uptake rate (Kirchman et al. 1985) and the bacterial biomass production (Simon and Azam 1989).

Total bacteria were detected by DAPI (4′,6-diamidino-2-phenylindole)-staining (Porter and Feig 1980). Formaldehyde-fixed water samples were collected onto 0.2 µm polycarbonate filters, cells were identified under ultraviolet (350–450 nm) excitation with a colour camera (Olympus DP71) on an Olympus BX51 microscope. Heterotrophic bacterial abundance, i.e. bacterial abundance (BA) was calculated by subtracting the abundance of autotrophic picoplankton (data not shown) from the total number of bacteria. The abundance of total bacteria was determined using YABBA-2011 (Yet Another Bacterial Biovolume Algorithm - http://www.technobiology.ch) software (Zeder et al. 2011), while heterotrophic bacterial abundance was calculated by subtracting autotrophic picoplankton types (data not shown) from the total bacteria number.

**Oxidation-reduction potential and oxygen content measurement**

The oxidation-reduction potential (ORP) of the sediment was determined at each sampling point every sampling date. ORP was measured with a multisensory probe containing 5 platinum coils mounted 10 cm apart on a 50 cm long fibreglass reinforced epoxy tube and attached to a 1-m long aluminium probe (Paleo Terra, Netherland). ORP data were registered with a custom-built voltmeter recorder (Paleo Terra, Netherland) against a saturated Ag/AgCl reference electrode and related to the standard hydrogen electrode.

Water oxygen content was measured in situ in every 25 cm of the water column using HQ30d multimeter with a LDO101 DO Probe (Hach, USA).

**Statistical analyses**

A two-way ANOVA was used to determine differences in abiotic and biotic factors with site (Lake Kolon and Lake Fertő) and macrophyte type (emergent and submerged) as independent variables, while measured parameters as dependent. Time-, site-, and habitat-specific differences were analysed by 3-way ANOVAs following data normalisation, where time, site (Lake Kolon and Lake Fertő) and macrophyte type (emergent and submerged) were as independent variables, while measured parameters as dependent. All data were graphed and statistically analysed (linear or non-linear regression) using SigmaPlot 12.5. We used generalized additive models (gam in R) to evaluate associations among water temperature, macrophytes and bacteria.

**Results**

In Lake Fertő, within the studied opening dominated by submerged macrophytes *Najas marina* L., *Utricularia vulgaris* L., *Hydrocharis morsus-ranae* L., and *Ranunculus circinatus* L. were observed, although the vast majority of the collected biomass (>99%) was *N. marina*. Submerged plants were of a detected in June, and, by the end of August, biomass peaked at 585 g [dry mass] m⁻² (Fig. 1). The emergent site in Lake Fertő was dominated by *Phragmites* (Supplementary
Fig. 1. Reed biomass accumulation began at the end of April and maximum biomass (1030 ± 350 g [dry mass] m⁻²) was reached at the end of July (Fig. 1). In the process of senescence, biomass gradually decreased to 600 ± 180 g [dry mass] m⁻² by the end of October (Fig. 1).

In Lake Kolon (Supplementary Fig. 1b), within the submerged macrophyte-dominated opening Nymphaea alba L., Nuphar lutea L., Sm, Ranunculus circinatus L., Utricularia vulgaris L., Hydrocharis morsus-ranae L., Stratiotes aloides L., and Ceratophyllum demersum L were found. These species were unevenly distributed; in the sampling area at the centre of the opening, only Nymphaea and Nuphar were dominant. By the end of April, plants were visible at the site, while their biomass peaked at the end of June, with 612 ± 83 g [dry mass] m⁻². By the end of summer, biomass was approximately 300 g [dry mass] m⁻² (Fig. 1).

The emergent macrophyte-dominated area of Lake Kolon (Supplementary Fig. 1b) consisted of Phragmites (> 74% of all biomass), Typha latifolia L., Carex sp., Nymphaea alba, Ceratophyllum demersum, Utricularia vulgaris. At the end of April, 70 ± 18 g [dry mass] m⁻² was collected from the sampling site, and maximum biomass was measured in August (1544 ± 356 g [dry mass] m⁻²) (Fig. 1).

The difference between the macrophyte biomass in Lake Fertő and in Lake Kolon was significant both for submerged (2-way ANOVA; F = 91.6; P = 0.011) and emergent (2-way ANOVA; F = 327.3; P < 0.001; Fig. 1) macrovegetation. The large difference between the emergent sites was a result of the smaller size of the Phragmites plants in Lake Fertő; at the peak of the growing season, the biomass of reed plants in Lake Kolon was 2.5–3 time higher than that of plants in Lake Fertő (42.3 ± 6.0 and 14.8 ± 1.6, respectively) as a result of both the taller and larger diameter plants (Supplementary Fig. 2). Nevertheless, Phragmites plants at both sites had nearly identical numbers and relative biomass of green leaves, yet during senescence, Phragmites plants from Lake Fertő lost ~55% of their leaf biomass, while plants from Lake Kolon lost ~21% (Supplementary Fig. 3). Further analysis showed that, except for nitrate content (Pearson: r = 0.976, P = 0.025), biomass accumulation did not correlate with any edaphic properties of the studied sites (Table 1) and was not related to the limnological properties of the sites (data not shown), probably because for the majority of parameters the lakes were similar with minor differences (Supplementary Tables 1–5).

BP at all sites showed a typical seasonal pattern, with low values in late autumn, at winter and the beginning of spring, and highest production between July and September (Fig. 2). BP of the submerged macrophyte dominated site in Lake Fertő (seasonal average 4.9 ± 0.6 µg C l⁻¹ h⁻¹) was significantly higher than that of Lake Kolon (2.1 ± 0.6 µg C l⁻¹ h⁻¹; 2-way ANOVA F = 11.0, P = 0.013) (Fig. 2a). The emergent-dominated sites have high BP values with seasonal averages of 4.9 ± 0.7 and 3.6 ± 0.5 µg C l⁻¹ h⁻¹ at Lakes Fertő and Kolon, respectively, though this difference was not significant (Fig. 2a; 2-way ANOVA F = 2.3, P = 0.169). BA in Lake Fertő and Lake Kolon were very similar, with maximal counts of 5.3*10⁶ and 7.0*10⁶ cells ml⁻¹ respectively (2-way ANOVA, P > 0.46; Fig. 2b).

At low BP and low BA the plant biomass was also low. Plant biomass peaked at intermediate heterotrophic bacterial production (Fig. 3) and heterotrophic bacterial abundance (Fig. 4). At heterotrophic bacterial production 5.6 µg C l⁻¹ h⁻¹ and heterotrophic bacterial abundance values of 5.3*10⁶ cells the macrophyte biomass peaked (~ 1.5 kg m⁻²) (Figs. 3 and 4). At time of high BP (> 8 µg C l⁻¹ h⁻¹) and high BA (> 6.0*10⁶ cells) had low macrophyte biomass (Figs. 3 and 4). GAM revealed that BP was not predicted by either macrophyte biomass or temperature. Bacterial abundance was significantly related to macrophyte biomass.

Table 1 Pearson product moment correlation (coefficient r and significance P) between seasonal maximal biomass and seasonal nutrient average at the study site in Lakes Fertő and Kolon

|        | NH₄-N | NO₃-N | TSN  | TN   | SRP  | TP   |
|--------|-------|-------|------|------|------|------|
| r      | -0.23 | 0.98  | 0.73 | 0.94 | 0.67 | 0.89 |
| P      | 0.775 | 0.025 | 0.268| 0.064| 0.334| 0.108|
Interaction among bacterioplankton and macrophytes in shallow lakes with high macrophyte biomass 

Interaction among bacterioplankton and macrophytes in shallow lakes with high macrophyte biomass and temperature was shown to be important (p < 0.05) but not temperature (Supplementary Table 6). Additional analysis showed some temporal and thermal relations between macrophytes biomass and bacterioplankton activity/abundance (Supplementary Figs. 4 and 5). Biomass accumulation rates also showed a relation to BP and BA (Fig. 5).

The original source of carbon for bacteria is the coloured dissolved organic matter coming from macrophytes (Fig. 6 and Supplementary Figs. 4 and 5). Our data show that both the BP and the BA in warm (above annual average, Supplementary Table 1) waters correlated with CDOM content of the water (Pearson Product Moment Correlation r = 0.63, P = 0.001 and r = 0.536, P = 0.010, respectively; Fig. 6). The bacterial activity changed the chemistry and the O2 content of the sediment and the water column. The oxidation reduction potential (ORP) of the upper layer of the sediment in the second half of the summer correlated with both BP (Pearson Product Moment Correlation r = −0.70, P = 0.012) and BA (Pearson Product Moment Correlation r = −0.66, P = 0.021; Fig. 7). The anoxic ORP of the sediment was apparent throughout the season and at different depths (Supplementary Fig. 7), driving the oxygen content of the water column into the unsaturated zone (Supplementary Fig. 8).

Discussion

Littoral zones are important and very productive parts of aquatic ecosystems (Costanza et al. 1997; Ziegler et al. 2015). Due to the characteristics of the littoral zone (inundation, lack of grazing, etc.), the majority of the macrovegetation biomass will eventually become litter, which could be substantial (Duarte and Kalf 1986; Rooney and Kalf 2000; Windham 2001; Karunaratne et al. 2003; Tóth 2016). The average annual production of submerged and emergent patches at Lake Fertő and Lake Kolon was ~0.5 and 1.4 kg m⁻² respectively. While no difference in the biomass accumulation of submerged macrophytes in Lakes Fertő and Kolon was shown, the above-ground production of Phragmites in Lake Kolon was higher than that in Lake Fertő (1.5 and 1.2 kg m⁻² y⁻¹, respectively). Besides the difference in accumulated biomass, the seasonal dynamics of litter production in the Phragmites stands was different too; in Lake Fertő it started earlier (July) than in Lake Kolon (September). Moreover, in Lake Fertő this litter production was higher (470 g m⁻²) than in Lake Kolon (346 g m⁻²), which results in a substantial carbon accumulation and a possible explanation for higher BP rates in Lake Fertő.

Our data indicate that the macrophyte-derived organic matter (CDOM) affects planktonic heterotrophic bacterial production and bacterial abundance throughout the whole study period, although this relation becomes more significant at warmer water temperatures. In return, this bacterial activity and abundance has influence on the biomass
and biomass accumulation rates of macrophytes. At activities higher than 5.6 µg C l⁻¹ h⁻¹ and bacterial abundance higher than 5.3*10⁶ cells the biomass of macrophytes could be affected and decrease by up to 50–60%. It is also shown that bacterial activity in spring could negatively impact macrophyte growth rates.

In our study we showed that increase in the number of heterotrophic bacteria and increase in their activity mainly due to increased litter decomposition resulted in unfavourable conditions, mostly by decreasing the ORP to anoxic conditions as previously described (Tóth 2016). Although some morphological, physiological and physical (Armstrong and Armstrong 1990; Amsberry et al. 2000) factors help macrophytes to survive these adverse conditions, extremely low ORP (oxygen depletion) eventually could not be tolerated by majority of macrophytes.

We were unable to relate the observed differences in macrophyte biomass to any measured limnological or chemical factor, and found that the bacteriological data do correlate with biomass accumulation, though not in a linear way. Similar results (i.e., plant biomass was positively related to bacterial activity), have also been observed in other cases (López et al. 1995; Isaksen and Finster 1996; Boschker et al. 2000). The increased activity and more abundant microbiota, change the oxygenation of area, implying accompanying biogeochemical changes, such as increased volatile fatty acid and sulphide pools (Findlay et al. 1986; Reitner et al. 1999; Jeppesen et al. 2002; Rooney and Kalff 2003), which might directly or indirectly affect plant growth.

Besides the absolute amount, the composition of the litter could affect bacterial activity and quantity. *Phragmites* litter, due its slow rate of decay, could cover the sediment surface for a long time, while submerged macrophyte litter will decompose significantly faster. N and P contents of the decomposing litter have a significant effect on bacterial activity (Güsewell and Gessner 2009), thus submerged macrophytes due to their higher N and P contents decompose...
significantly faster, while the decay rate of emergent macrophytes is lower, due to their extremely limited N and P (Enríquez et al. 1993; Windham and Lathrop 1999; Güsewell and Gessner 2009). Moreover, since the stems and leaves of Phragmites have different decay rates (Windham 2001; Balogh et al. 2006), the composition of the litter could also affect activity of the microbiota. In our study, in Lake Fertő the foliar litter production was higher (~230 g m⁻²) than in Lake Kolon (66 g m⁻²), which results in a higher, easy to access carbon pool and could be a possible explanation for higher heterotrophic bacterial production rates in Lake Fertő.

In conclusion, our data showed that bacterial activity could affect macrophyte growth, both directly and indirectly through changing the oxygenation of the sediment and water. Over specific bacterial activity and abundance macrophyte biomass accumulation is showing their multifaceted relationship: macrophytes are not only the most significant carbon source for the bacteria, but are also a competing organism.

Acknowledgements This research was supported by grants National Research, Development and Innovation Office—NKFIH, K-116666 and National Research, Development and Innovation Office—NKFIH, KH-129505 of the Hungarian National Research, Development and Innovation Office. The authors are grateful to Stephanie C. J. Palmer for her help with the English of the text.

Funding Open access funding provided by ELKH Centre for Ecological Research.

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Fig. 7 Relation between bacterial production (a), heterotrophic bacterial abundance (b) and the oxidation reduction potential of the upper 10 cm of the sediment at the studied sites near the population climax (July–August–September). Pearson product moment correlations: a—r= −0.70, P= 0.012; b—r = − 0.66, P= 0.021
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