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Bacterial community response to changes in a tri-trophic cascade during a whole-lake fish manipulation

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Abstract. Microbial communities play a key role in biogeochemical processes by degrading organic material and recycling nutrients, but can also be important food sources for upper trophic levels. Trophic cascades might modify microbial communities either directly via grazing or indirectly by inducing changes in other biotic or in abiotic factors (e.g., nutrients). We studied the effects of a tri-trophic cascade on microbial communities during a whole-lake manipulation in which European perch (Perca fluviatilis) were added to a naturally fishless lake divided experimentally into two basins. We measured environmental parameters (oxygen, temperature, and nutrients) and zooplankton biomass and studied the changes in the bacterial community using next generation sequencing of 16S rRNA genes and cell counting. Introduction of fish reduced the biomass of zooplankton, mainly Daphnia, which partly altered the bacterial community composition and affected the bacterial cell abundances. However, the microbial community composition was mainly governed by stratification patterns and associated vertical oxygen concentration. Slowly growing green sulfur bacteria (Chlorobium) dominated the anoxic water layers together with bacteria of the candidate division OD1. We conclude that alterations in trophic interactions can affect microbial abundance, but that abiotic factors seem to be more significant controls of microbial community composition in sheltered boreal lakes.

Key words: anaerobic microbes; bacterial community composition; boreal lakes; next generation sequencing; top-down vs. bottom-up control; trophic cascades.

INTRODUCTION

Microbes have a significant role in biogeochemical cycles in ecosystems, but they also constitute a large fraction of ecosystem biomass and can be an important food source for higher trophic levels, especially in aquatic environments (Tranvik 1992). Thus understanding the abiotic and biotic factors that regulate microbial communities is important. Seasonal succession together with changing temperature and pH (Lindström et al. 2005, Yannarell and Tripplett 2005, Crump et al. 2009, Eiler et al. 2012), light and humic substances (Kent et al. 2004, Haukka et al. 2005), oxygen (Shade et al. 2008, Peura et al. 2012a), and the availability of inorganic and organic nutrients (Zeng et al. 2009, Zhang et al. 2015) are important “bottom-up” factors controlling bacterial community composition. In addition, trophic interactions (i.e., “top-down control”) may shape bacterial communities. Previous studies have focused on demonstrating how factors of direct trophic control such as predation (Pernthaler 2005) or resource competition (Peura et al. 2012b, de Vries and Shade 2013) have selective community-level effects on bacterial diversity.

Boreal humic lakes receive high organic carbon inputs, especially of dissolved organic carbon (DOC), from forest and peatland catchments. In these environments, microbes have an important role, not only in controlling greenhouse gas fluxes, but in transferring this incoming allochthonous carbon to higher trophic levels via a “microbial link” (Jones 1992, Jansson et al. 2000). Boreal lakes, and particularly smaller ones, exhibit steep vertical gradients of temperature, oxygen, nutrients, and light, meaning that each water layer provides a unique sub-environment for bacteria. Therefore, the composition and function of microbial communities differ markedly between the oxic and anoxic water layers (Shade et al. 2008, Taipale et al. 2009a, Peura et al. 2012a). For example, anaerobic phototrophs, such as green sulfur bacteria (GSB) of the genus Chlorobium, can dominate the anoxic water layers of these lakes (Arvola et al. 1992, Karhunen et al. 2013), whereas methane-oxidizing bacteria (MOB) thrive at the oxic-anoxic interface (Kankaala et al. 2006, Taipale et al. 2009a). The main microbial groups in the oxic upper water layer (epilimnion) have been assigned to Actinobacteria (Lindström and Leskimen 2002, Haukka et al. 2005, Newton et al. 2011) and Betaproteobacteria (Burkert et al. 2003, Grossart et al. 2008). In addition, the suboxic and anoxic lower water layers (meta- and hypolimnion) of these lakes have been...
shown to harbor previously unknown groups, such as candidate division OD1 (Peura et al. 2012a).

As in other ecosystems, the composition and diversity of microbial communities in small boreal lakes are controlled by abiotic bottom-up factors, such as DOC and nutrient concentrations (Peura et al. 2013), and by biotic top-down factors, i.e., grazing (Lindström 2000, Kent et al. 2004). The relative importance of bottom-up and top-down factors varies with respect to the trophic status of the lake, with higher nutrient concentrations increasing bottom-up effects and nutrient-limitation enhancing top-down control (Gasol et al. 2002). In stratified boreal lakes, the oxic epilimnion becomes nutrient-limited during summer, while the anoxic hypolimnion can have high concentrations of phosphate and ammonium, creating different patterns of top-down and bottom-up effects on microbial communities between these depth zones. However, as microbes are an important food source for consumers at higher trophic levels, they can also be controlled indirectly via trophic cascades, whereby predators, by decreasing the abundance of bacterivores, suppress grazing pressure on microbes (Thakur et al. 2014).

Naturally fishless small humic lakes provide excellent environments to study the potential impacts of cascading trophic interactions from higher trophic levels to bacterial communities, as introducing a new top predator is relatively straightforward. Fishless lakes often have dense populations of large zooplankton, such as cladocera from the genus Daphnia. Due to their ability to filter feed on a broad spectrum of particle sizes, Daphnia can exert a significant grazing pressure on all components of microbial food webs, including pico-, nano-, and microplankton (Jürgens et al. 1997, Muylaert et al. 2002). Daphnia can also migrate vertically in the water column, allowing them to utilize resources below the oxic water layers and thereby enabling the use of microbial biomass as energy subsidies when phytoplankton is scarce (Salonen and Lehtovaara 1992). For example, in a small and highly humic Finnish lake Mekkojärvi, Daphnia longispina are known to feed extensively on methanotrophs and partly also on GSB (Taipale et al. 2009b). Altering the food web structure by adding fish as new top predators to this kind of sheltered boreal lake was shown to reduce the zooplankton biomass and alter the zooplankton community structure (Järvinen and Salonen 1998), and can therefore be expected to modify bacterial communities as well.

We experimentally tested the effect of a tri-trophic cascade on microbial communities by dividing a small humic and naturally fishless lake into two basins with a plastic curtain, and adding fish to one basin. Zooplankton biomass decreased rapidly after fish addition to the fish-present basin, whereas, in the fish-absent basin, the zooplankton biomass increased through the summer toward the autumn (described by Devlin et al. 2015). We studied the changes in environmental parameters and microbial cell numbers, as well as evaluating bacterial community composition using next generation sequencing (NGS) of the 16S rRNA genes. The study covered three open-water seasons and provided a data set with exceptional coverage of temporal and spatial observations of epi-, meta-, and hypolimnetic microbial communities. We expected higher bacterial cell densities after Daphnia removal, taking into account the strong effect of Daphnia on microbes previously observed in this lake (Arvola et al. 1992). A more pronounced effect of Daphnia abundance on the microbial community was expected in the oxic epilimnion where Daphnia mainly feed than in the deeper anoxic hypolimnion, which Daphnia can only visit during short vertical migration forays. In addition, altered trophic cascades were expected to modify microbial community composition via decreased predation pressure, thus favoring fast-growing microbial species.

**Methods**

**Study lake and design**

Mekkojärvi is a small, shallow (mean depth 3 m), and highly humic lake located in southern Finland in the Evo forest area. The lake is steeply stratified and has a shallow (~0.5 m during the open-water period) oxic epilimnion. The lake is ice covered from November to April and develops total and persistent anoxia during the winter. Mekkojärvi is partially meromictic, as the annual spring mixing can remain incomplete. The lake lacks fish due to the persistent winter under-ice anoxia, leading to exceptionally high summer zooplankton abundance, mainly of Daphnia longispina (further characteristics of Mekkojärvi are given in Salonen and Lehtovaara 1992, and Taipale et al. 2008).

In 2011, 2012, and 2013 Mekkojärvi was divided into two separate basins with a plastic curtain extending from surface to bottom. The curtain was set in place each spring after ice melt and spring mixing. In early July each year, European perch (Perca fluviatilis) were added in numbers representing naturally occurring perch biomass in similar lakes in the Evo forest area (approximately 1 g fresh mass per m² of lake surface area). Adult perch (>12 cm) were added to one basin and juveniles (<3 cm) to the other basin. However, due to a brief pulse of hypoxic conditions (repeated in every study year) in one basin, the established perch population died shortly after introduction, leaving the basin fishless and thus, in practice, creating two contrasting treatments of fish present and fish absent. Further details of the fish treatments are given in Devlin et al. (2015).

**Sample and data collection**

Both basins of the lake were sampled one to two times per month during the ice-free period (see Devlin et al. [2015], for full description of sampling, measurement of environmental factors and nutrients, and of zooplankton identification and calculation). Briefly, on each sampling occasion, temperature and oxygen profiles were measured in situ using a portable field meter (YSI model 58; Yellow Springs Instruments, Yellow Springs, Ohio, USA). Integrated water column samples were collected with a Limnos water sampler (height 60 cm, volume 4.25 L; Limnos, Turku,
Finland) from epilimnion (0–0.5 m), metalimnion (0.5–1.0 m), and hypolimnion (1–3 m), delimited according to thermal and oxygenic stratification. Concentrations of dissolved inorganic P (PO$_4^{3-}$) and N (NH$_4^+$, NO$_2^-$, NO$_3^-$), and of DOC in the lake water were determined following standard laboratory methods (available online). Zooplankton samples were collected with a Limnos water sampler and 6 L of water were passed through a 50-μm mesh net to retain zooplankton. The zooplankton were later identified, counted, and measured under a stereo microscope. Daphnia biomass (w, mg C) was then calculated using length–mass relationships $w = aL^b$ from Rahkola et al. (1998) where the intercept $a$ was estimated as 5.66, $L$ is the Daphnia body length in mm, and the slope $b$ was 1.72. Bacterial cell counts were measured from 2012 samples ($n = 42$, covering seven sampling occasions from the three depth layers and both lake basins) by flow cytometry (FACSCanto II; Becton Dickinson Biosciences, Oxford, UK). Water samples (1.5 mL) were fixed with particle-free 20% w/v paraformaldehyde dissolved in Milli-Q water, 1% final concentration. The fixed samples were stored at $-20{\degree}C$. Before the analyses the samples were thawed and stained with SYBR Green I DNA stain (Sigma-Aldrich) in the 1:5000 final dilution of the initial stock. Yellow-green 0.5-mm long, 0.5-μm diameter beads (Fluoresbrite Microparticles; Polysciences, Warrington, Pennsylvania, USA) were added with known final concentration (10$^5$ beads/μL) to determine absolute cell concentrations (Zubkov and Burkill 2006). Flow cytometry counts were enumerated with Flowing Software (Flowing Software ver 2.5, Turku, Finland).

**Bacterial community composition**

Water for DNA extraction was collected from epilimnion (0–0.5 m), metalimnion (0.5–1.0 m), and hypolimnion (1.0–3.0 m) and pooled samples from each stratum were stored in 100-mL plastic containers and frozen at $-20{\degree}C$. The DNA samples were then freeze dried (Alpha 1–4 LD plus, Christ) and extracted using a PowerSoil DNA extraction kit (MoBio Laboratories) according to the manufacturer’s instructions. Altogether, 96 DNA samples were included, covering 16 different sampling time points during the three-year experiment ($n = 4$ in 2011, $n = 7$ in 2012, and $n = 5$ in 2013) from all three depth layers and from both lake basins. After extraction, DNA concentrations of the samples were measured with a Qubit 2.0 Fluorometer (Invitrogen) using the Qubit dsDNA HS Assay kit (Life Technologies). Amplification of bacterial 16S rRNA genes ($E. coli$ positions 27 to 338) was conducted using general bacterial primers 27F (5’-AGAGTTTGATCMTGGCTCAG-3’) and 338R (5’-GTCTGGCTCCGTAGGAGT-3’). Primer 27F carried an Ion Torrent adaptor A at the 5’ end and an 11–12 bp barcode followed by a KEY tag, and primer 338R carried a P1 adaptor at the 5’ end. PCR amplification for Ion Torrent sequencing was done in 20-μL volumes containing 1 μL of template DNA, 0.5 μmol/L of each primer, 1× Phusion HotStartII buffer (Finnzymes), 0.02 U Phusion HotStartII polymerase, and 0.3 mmol/L dNTPs. Cycling conditions involved an initial denaturing step at 98°C for 2 min, followed by 25 cycles of 98°C for 10 s, 52°C for 10 s, and 72°C for 30 s, and a final elongation step of 72°C for 10 min. The PCR amplicons were purified twice using the Agencourt AMPure XP purification system (Beckman Coulter) and quantified with a Qubit 2.0 Fluorometer. Equal concentrations of bar-coded amplicons were pooled for the emulsion PCR, which was performed using an IonOneTouch2400template kit (Life Technologies). Templated beads were purified using the OneTouch ES instrument (Life Technologies) according to the manufacturer’s protocol. NGS of the pooled library was conducted starting from the A adaptor using the Ion Torrent Personal Genome Machine (PGM; Life Technologies) with Ion Sequencing 400 kit and Ion 314 Chip. Sequence quality control was done using Mothur (Schloss et al. 2009), whereby sequences shorter than 200 bp, low-quality sequences with mismatches in bar code and primer sequences or with homopolymers longer than eight nucleotides, as well as barcodes, primers, and chimeras were removed. A final total of 258 572 reads was obtained. The Ion Torrent sequences have been deposited in the NCBI Short read Archive under accession number SRP061976.

To assign reads to operational taxonomic units (OTUs) a 97% sequence similarity cut-off was used in Mothur (available online). OTUs containing at least 50 reads were included in further analysis (covering 79% of all sequences). A representative sequence from each of the 597 OTUs obtained was classified using the Least Common Ancestor tool of the SINA ALIGNER v1.2.11 (Pruesse et al. 2012). Classification was based on the SILVA taxonomy for the 10 nearest neighbors retrieved from the curated SILVA SSU Ref database Release 106 (Pruesse et al. 2007) with minimal identity of 0.80. From the 597 OTUs, only seven remained as unclassified at the phylum level.

**Statistical analyses**

For studying the bacterial $\alpha$-diversity, the data set was resampled with the command “subsample” in Mothur to gain equal numbers of sequences in each sample (2019 sequences). The $\alpha$-diversity was estimated with the inverse Simpson index as a measure of diversity and with the Chao index as a measure of richness. All statistical analyses were conducted using R version 3.1.1 (R Core Team 2014). Disparities in $\alpha$-diversity (Chao or inverse Simpson index) between treatments and water layers within lake sides were studied with nested two-way ANOVA with post hoc tests. Differences in the relative abundances of key bacterial taxa (Chlorobium, Burkholderiales, OD1, and Rhodospirillales) in the hypolimnion were examined with two-way ANOVA with post hoc tests, or with two-way adjusted ranks transformation test (ART; Leys and Schumann 2010). In the

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1 http://www.sfs.fi/
2 http://www.mothur.org/
result, Basin refers to the different lake sides that are separated with the plastic curtain and Treatment refers to fish addition (meaning before and after fish addition). For inspecting bacterial community composition, functions from the Vegan package (Oksanen et al. 2013) were used. Nonmetric multidimensional scaling (NMDS, conducted with metaMDS function) plots calculated based on Morisita-Horn distance matrix (Horn 1966) were used to visualize dynamics in community structure ($\beta$-diversity) of bacteria (OTUs represented by at least 50 reads) for total data as well as separately for the three water layers. Before NMDS, Wisconsin and square-root transformations were applied to OTU abundance data. Environmental data (oxygen concentration, temperature, DOC concentration, inorganic nitrogen concentrations ($\text{NO}_2^- + \text{NO}_3^- + \text{NH}_4^+$), phosphate concentration ($\text{PO}_4^{3-}$) as well as $\text{Daphnia}$ density and season were fitted to the ordination using the function “envfit.” Differences in bacterial community between water layers as well as changes in the bacterial community within a water layer, between lake basins, and treatments (separately and their interactions) were tested with permutational multivariate analysis of variance (PERMANOVA) using the function “adonis” in R. To normalize the data, log-transformation was used for the inverse Simpson index and cube-root transformation for bacterial cell counts.

**Results**

Mekkojärvi developed steep stratification every year from May to September, resulting in a shallow oxic epilimnion (maximum depth 0.5 m) and anoxic meta- and hypolimnion (Appendix S1: Table S1). Average nutrient concentrations ($\text{NH}_4^+$, $\text{NO}_2^- + \text{NO}_3^-$ and $\text{PO}_4^{3-}$) were 10-fold higher in the hypolimnion than in the epilimnion (Table S1), but the concentrations did not differ between the two experimental lake basins in the epilimnion or in the hypolimnion (Mann-Whitney $U$, $P > 0.05$). However, in the metalimnion (0.5–1.0 m) $\text{NH}_4^+$ and $\text{PO}_4^{3-}$ concentrations were higher in the fish-absent basin than in the fish-present basin (Mann-Whitney $U$, $P = 0.004$ and $P = 0.004$, respectively; Table S1).

Prior to fish addition, oxygen concentrations were slightly higher in the epilimnion and metalimnion (Mann-Whitney $U$, $P = 0.009$ and $P = 0.03$, respectively) in the fish-present basin, whereas after fish addition no differences were observed (Mann–Whitney $U$, $P > 0.05$). The bacterial cell counts from 2012 indicated that the fish addition and water depth were the most important factors affecting bacterial density (Fig. 1). Before fish addition, bacterial cell densities increased from epi- to hypolimnion in both basins. After fish addition, cell densities were higher in the fish-present than in the fish-absent basin in all depth layers. When comparing the situation before fish addition to that after fish addition within basins, cell-densities decreased in all depth layers in the fish-absent basin, whereas in the fish-present basin, they remained almost at the same level (Fig. 1). The observed difference in the cell densities was smaller in the epilimnion (28% difference) than in the metalimnion or hypolimnion (68% difference in both). However, due to the low number of replicates ($n = 2$) no statistical analysis could be done with reliable results.

Number of observed OTUs at 97% similarity level varied between 244 and 910 within samples, and was on average 509, 566, and 603 in epili-, meta-, and hypolimnion samples, respectively. The bacterial community composition visualized with NMDS clearly differed between the different depth layers (Fig. 2), with epilimnion and

![Fig. 1. Bacterial cell counts (mean ± SE) from 2012 from different depth layers and lake basins before fish addition (three sampling times), after fish addition (two sampling times), and during the transition phase just after fish addition (two sampling times).](image-url)
The microbial community composition was affected by the treatment (i.e., fish presence/absence) and study year in all depth layers, while lake basin was found to shape community composition in the epilimnion and the hypolimnion (Table 1). Both Chao richness index and inverse Simpson index remained similar throughout the season regardless of fish addition (Table 2). Bacterial community richness (Chao) in the fish-present basin increased from the epilimnion to the hypolimnion ($P < 0.001$), but no difference in richness was seen between these layers in the fish-absent basin ($P > 0.05$, Appendix S1: Table 2). The inverse Simpson index was lower in the fish-absent basin than in the fish-present basin ($P = 0.024$). Because the epilimnetic community compositions were found to be distinct between the two lake basins ($P = 0.003$), we evaluated separately which environmental factors had most influence on the communities. NO$_3^-$+NO$_2^-$ concentrations correlated with the epilimnetic community composition in both basins, oxygen correlated with the community in the fish-present basin and DOC in the fish-absent basin (Fig. 4). The metalimnetic and hypolimnetic communities were controlled by the same environmental parameters and thus assessed together without basin distinction. Highest correlations were observed between oxygen concentration in the metalimnion, and temperature in the hypolimnion, and season in both depth layers (Fig. 4).

Four phylogenetic groups dominated the microbial community throughout the water column in all the study years (Appendix S2: Figs S1–S3). The most abundant OTU (OTU001) overall in the water column was assigned to the green sulfur bacteria (GSB) group, genus *Chlorobium*. This OTU contributed on average 8%, 36%, and 41% of all sequences in the epi-, meta-, and hypolimnion, respectively (Fig. 3). Altogether, from the 597 OTUs that contained at least 50 reads, 64 OTUs were classified as *Chlorobium,*
which means that from the hypolimnetic and metalimnetic communities almost half of all bacterial sequences were assigned to these anaerobic phototrophs (Fig. 4). In the epilimnion, the order Burkholderiales, belonging to a class Betaproteobacteria, was the most abundant bacterial group comprising, on average, 24% of epilimnetic bacterial sequences, being the closest match for 67 OTUs (including the second most abundant OTU, OTU002). Members of group candidate division OD1 were also abundant and found in high densities from the hypolimnion and metalimnetic community, the second most abundant OTU, OTU002). Members of group comprising, on average, 24% of epilimnetic bacterial sequences, being the closest match for 67 OTUs (including the second most abundant OTU, OTU002). Members of group candidate division OD1 were also abundant and found in high densities from the hypolimnion and metalimnion (Fig. 4), where OTU003 was the second- and third-most abundant OTU: 12% of all sequences in the hypolimnion and 9% of those in the metalimnion were assigned to this division. A biogeochemically important group, MOB, was present in high numbers in all water layers, being most abundant in the metalimnion where they comprised, on average, 9% of the microbial community. All the sequences of methanotrophs belonged to Gammaproteobacteria group Methylococcales, which formed the dominant group in the sequences of Gammaproteobacteria. Of the key microbial groups of the hypolimnion (Chlorobium, Burkholderiales, OD1, and Rhodospirillales), Burkholderiales and Rhodospirillales abundance stayed at the same relative abundance throughout the whole season in both basins, whereas the proportion of Chlorobium decreased, and OD1 increased toward autumn (Appendix S3: Fig. S4).

### Discussion

Consistent with our expectations, an altered trophic cascade was seen to manifest at the microbial level in the lake experiment; the number of bacterial cells was higher in all depth layers in the basin where Daphnia were removed by the fish, indicating that fish addition decreased Daphnia grazing on bacteria. Daphnia are also capable of migrating vertically in the water column despite strong oxygen stratification (Arvola et al. 1992), explaining why we found higher bacterial cell numbers throughout the water column in the fish-present basin. We found distinct microbial community structures and dominant OTUs in each water layer and in each study year. Fish addition was found to alter microbial community composition in the epilimnion and hypolimnion, where in addition the abiotic environmental factors exerted important controls on the community structures. Although differences in community composition were found between years, the same core phylogenetic groups dominated the microbial communities in each study year.

In accordance with our expectations, the effect of an altered trophic cascade was seen in the epilimnion, where microbial communities differed between fish-present and fish-absent basins. In addition, abiotic factors, such as nutrient and oxygen concentrations, were found to control the microbial community compositions. DOC concentration has been repeatedly linked to lake microbial communities in which Betaproteobacteria and Bacteroidetes, both abundant in our samples, have been shown to favor high DOC concentrations (Burkert et al. 2003, Eiler et al. 2003, Hutalle-Schmelzer et al. 2010). Nitrate concentration was connected to epilimnetic communities, and has also been shown to control the sediment community composition in freshwaters (Zhang et al. 2015) where it can further affect biogeochemical processes and greenhouse gas \( \text{N}_2\text{O} \) production (Piña-Ochoa and Álvarez-Conbelas 2006, Saarenheimo et al. 2009). In addition, the difference in microbial community composition between lake basins can be explained by different selection forces. In the fish-present basin, the top-down selection force (Daphnia) was removed enabling within-microbial-community interactions, such as competition. In the fish-absent basin, Daphnia were presumed to exert high unselective grazing pressure. However, we did not find any clear interactions between Daphnia density and the main key microbial groups, possibly because the dominant OTUs were insensitive to Daphnia grazing, masking changes in minor OTUs. In contrast to our expectations, removal of Daphnia also modified the hypolimnetic microbial community composition, which probably reflects the absence of vertically migrating Daphnia (Arvola et al. 1992, Taipale et al. 2009a). We suggest that the hypolimnetic microbial community in the fish-absent basin was modified by both Daphnia grazing and abiotic factors, which might have

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**Table 1.** Results of permutational multivariate analysis of variance (PERMANOVA) comparing epilimnetic, metalimnetic, and hypolimnetic bacterial communities between treatments and basins, within each layer.

<table>
<thead>
<tr>
<th>Factor</th>
<th>Epilimnion</th>
<th>Metalimnion</th>
<th>Hypolimnion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>df</td>
<td>P</td>
<td>F</td>
</tr>
<tr>
<td>Year</td>
<td>1.19</td>
<td>5.049 &lt;0.05</td>
<td>2.560 &lt;0.05</td>
</tr>
<tr>
<td>Treatment</td>
<td>2.19</td>
<td>2.836 &lt;0.05</td>
<td>1.902 &lt;0.05</td>
</tr>
<tr>
<td>Basin</td>
<td>1.19</td>
<td>4.757 &lt;0.05</td>
<td>1.416 ns</td>
</tr>
<tr>
<td>Year × treatment</td>
<td>2.19</td>
<td>2.062 &lt;0.05</td>
<td>1.150 ns</td>
</tr>
<tr>
<td>Year × basin</td>
<td>1.19</td>
<td>1.094 ns</td>
<td>0.967 ns</td>
</tr>
<tr>
<td>Treatment × basin</td>
<td>2.19</td>
<td>1.351 ns</td>
<td>1.119 ns</td>
</tr>
<tr>
<td>Year × treatment × basin</td>
<td>2.19</td>
<td>0.862 ns</td>
<td>1.024 ns</td>
</tr>
</tbody>
</table>

*Note: Boldface type highlights significant differences; ns, not significant.*
favored fast growing bacterial taxa. Furthermore, we suggest that only seasonal factors (e.g., temperature and oxygen) controlled the community in the fish-present basin, leading to dominance of slow-growing bacterial groups. Similarly, the metalimnetic community was affected by seasonal factors in both basins.

Our results corroborate previous findings that anoxic waters of steeply stratified lakes harbor unique microbial communities (Shade et al. 2008, Peura et al. 2012), which are not usually classified as belonging to the common freshwater tribes (Newton et al. 2011). We found that the anoxic water layers (metalimnion and hypolimnion) were overwhelmingly dominated by green sulfur bacteria (Chlorobium), which are obligate anaerobic photoautotrophs and thus an important but largely unacknowledged part of the carbon cycle (Van Gemerden and Mas 1995, Karhunen et al. 2013). In addition, the bacterial group belonging to candidate division OD1 was found to be abundant. The role of OD1 in lake ecosystems is currently unknown. We found OD1 continuously throughout the open water seasons from both metalimnion and hypolimnion in relative abundances similar to those earlier reported from Mekkojärvi and other nearby lakes (Peura et al. 2012). We could also see that the proportion of OD1 increased in both lake basins toward autumn in the hypolimnion. The relative proportion of OD1 might even have been underestimated due to the primer choice (Peura et al. 2012). In contrast, a few epilimnetic samples might have contained some OTUs from the metalimnion, as strictly anaerobic GSB and presumably anoxic OD1 were recorded from those epilimnetic samples; this probably means that the bottom of our 60-cm Limnos sampler may have sometimes penetrated below the extremely narrow oxic epilimnion in the lake.

Spring and autumn mixing events have been shown to modify the microbial communities of small humic lakes by destroying the conditions necessary for strictly anaerobic groups and supporting the growth of minor groups (Taipale et al. 2011, Garcia et al. 2013). Although communities were found to differ between years, the same dominant phylogenetic groups developed repeatedly in each study year in the two lake basins after the possible spring mixing. The community compositions of the lower water layers were clearly affected by the seasonal succession, as reported previously (Shade et al. 2007, Crump et al. 2009, Eiler et al. 2012). Thus it seems that abiotic factors are more significant in controlling the microbial community than biotic factors (e.g., Daphnia) in this kind of sheltered boreal lake. One reason for this could be that Daphnia removal opened a feeding niche for other
bacterivores, such as rotifers and ciliates (Järvinen and Salonen 1998, Jürgens et al. 1999, Zöllner et al. 2003), which could then have exerted similar grazing pressure and top-down control towards the microbial communities as *Daphnia* had before fish addition. In particular, top-down grazing pressure by ciliate and flagellate protists on bacteria has been shown to modify both the size fractionation and structure of microbial communities (Pernthaler 2005), where especially Alphaproteobacteria, Actinobacteria, and Flavobacterium have been shown to develop grazing-resistant forms and become the most abundant bacterial groups (Jürgens et al. 1999, Pernthaler 2005). We do not have quantitative information about the abundance of rotifers or ciliates during the experiment. However, while counting crustacean zooplankton, rotifers were observed only in low numbers throughout the ice free season and within each treatment basin, and the altered trophic cascade was not seen to induce any noticeable change in the abundance of rotifers. Regarding ciliates, the minor decrease in bacterial cell densities in the epilimnion but not in the hypolimnion in the fish-absent basin in 2012 could reflect a greater presence of ciliates in the epilimnion, but their densities must have been low because there was still an observable difference between lake basins. Hence, although the removal of *Daphnia* by the introduced fish may have been partly compensated by greater development of other bacterivorous zooplankton groups, the responses seen in the microbial community composition and abundance suggest that any such compensation remained limited.

In conclusion, the microbial community composition was affected partly by the top-down tri-trophic cascade, but more strongly by natural abiotic factors, which were linked to the season and other abiotic characteristics of the lake. The steep stratification patterns with low pH, low oxygen concentrations, and restricted light intensities in lower water layers created an extreme environment in the deeper layers where only certain specialized groups were able to grow. However, there was an effect of tri-trophic cascades on bacterial abundance, as cell numbers were higher in the fish-present than in the fish-absent basin. This finding is further supported by previous results where quantitative abundance of MOBs was assessed throughout the entire three-year experiment, and similar responses in the bacterial cell numbers were observed (Devlin et al. 2015). The effect of the altered trophic cascade could even be further connected to lake biogeochemical processes as reduced methane emissions were measured from the fish-present basin (Devlin et al. 2015). It seems that in the future, when climate change and anthropogenic activities are expected to modify characteristics in aquatic ecosystems (reviewed in Jeppesen et al. 2009), microbial abundance and community composition may be modified not only directly by changing nutrient inputs or oxygen concentrations, but also by altered trophic cascades. Furthermore, changes in microbial community composition might lead to changes in biogeochemical cycles and in greenhouse gas emissions (Thakur et al. 2014, Devlin et al. 2015).

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Supporting Information

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