Ciliate community structure and interactions within the planktonic food web in two alpine lakes of contrasting transparency

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SUMMARY

1. Climate warming is accelerating the retreat of glaciers and recently, many ‘new’ glacial turbid lakes have been created. In the course of time, the loss of the hydrological connectivity to a glacier causes, however, changes in their water turbidity and turns these ecosystems into clear ones.

2. To understand potential differences in the food-web structure between glacier-fed turbid and clear alpine lakes, we sampled ciliates, phyto-, bacterio- and zooplankton in one clear and one glacial turbid alpine lake, and measured key physicochemical parameters. In particular, we focused on the ciliate community and the potential drivers for their abundance distribution.

3. In both lakes, the zooplankton community was similar and dominated by the copepod Cyclops abyssorum tatricus and rotifers including Polycarthra dolichoptera, Keratella hiemalis, Keratella cochlearis and Notholca squamula. The phytoplankton community structure differed and it was dominated by the planktonic diatom Fragilaria tenera and the cryptophyte alga Plagioselmis nannoplanctica in the glacial turbid lake, while chrysophytes and dinoflagellates were predominant in the clear one.

4. Ciliate abundance and richness were higher in the glacial turbid lake (∼4000–27 800 Ind L⁻¹, up to 29 species) than in the clear lake (∼570–7150 Ind L⁻¹, up to eight species). The dominant species were Balanion planctonicum, Askenasia cf. chlorelligera, Urotricha cf. furcata and Mesodinium cf. acarus. The same species dominated in both lakes, except for Mesodinium cf. acarus and some particle-associated ciliates, which occurred exclusively in the glacial turbid lake. The relative underwater solar irradiance (i.e. percentage of PAR and UVR at depth) significantly explained their abundance distribution pattern, especially in the clear water lake. In the glacial turbid lake, the abundance of the dominating ciliate taxa was mainly explained by the presence of predatory zooplankton.

5. Our results revealed an unexpected high abundance and richness of protists (algae, ciliates) in the glacial turbid lake. This type of lake likely offers more suitable environmental conditions and resource niches for protists than the clear and highly UV transparent lake.

Keywords: climate change, high mountain lakes, protists, turbidity, UV radiation

Introduction

Recently, molecular studies identified alpine lakes as hotspots of eukaryotic protistan diversity (Triadó-Margarit & Casamayor, 2012; Kammerlander et al., 2015; Filker et al., 2016). However, ciliate assemblages are still an underexplored group in those lakes (Wille et al., 1999; Sonntag, Summerer & Sommaruga, 2011), even though within the microbial food web they are ecologically important as major link to higher trophic levels (Sommer et al., 2012; Lischke et al., 2016). Planktonic ciliate assemblages of oligo- and mesotrophic lowland lakes...
characteristically include small algivorous prostomatid and oligotrich ciliates and their community and vertical distribution patterns are related to food availability, predation and abiotic factors (Weisse & Müller, 1998; Sonntag et al., 2006; Van Wichelen et al., 2013; Posch et al., 2015). In the clear alpine Gossernköllesee (Austrian Central Alps), only Balanion planctonicum, Urotricha cf. castalia and the mixotrophic Askenasia chlorelligera are found during the ice-free season, when they present a characteristic species-specific distribution in the water column (Wille et al., 1999; Sonntag et al., 2011). Among the environmental factors known to influence the vertical distribution patterns of zooplankton, ultraviolet radiation (UVR) is important (Williamson et al., 2011). In clear alpine lakes, the harmful solar UV-B radiation (280–320 nm) can easily reach the lake bottom (Laurion et al., 2000). In such lakes, vertical downward migrations during the day are known for phyto- and zooplankton (Tilzer, 1973; Tartarotti et al., 1999) but not for ciliates (Sonntag et al., 2011). For zooplankton, UVR has been identified to be an important driver in transparent lakes (Fischer et al., 2015), whereas high nutrient concentrations near the lake bottom can influence the vertical distribution of phytoplankton (Tilzer, 1973; Saros et al., 2005). Nevertheless, surface avoidance alone might be insufficient in these relatively shallow and highly UVR transparent alpine lakes. Other strategies, including the synthesis or accumulation of photoprotective compounds, or effective DNA repair mechanisms, are prerequisites and are found in a variety of planktonic organisms (Mitchell & Karentz, 1993; Sommaruga, 2001; Tartarotti et al., 2014).

As climate warming is accelerating glacier melting processes resulting in many newborn proglacial and ice-contact lakes (Carrivick & Tweed, 2013; Vaughan et al., 2013), the high loads of inorganic particles (‘glacial flour’) in glacier-fed lakes strongly influence abiotic factors and consequently their biota (Sommaruga, 2015). These ecosystems are light-limited and subsequently, the primary production, the distribution of phyto- and zooplankton and the growth and survival of protists are affected (Hylander et al., 2011; Sommaruga & Kandolf, 2014; Sommaruga, 2015). Recently, next-generation sequencing revealed high diversities of pro- and eukaryotes in a glacial turbid alpine lake (Kammerlander et al., 2015; Peter & Sommaruga, 2016). Although these lake types are in the spotlight of upcoming studies due to the strong progressive retreat of ice sheets in alpine and arctic regions (Sommaruga, 2015; Vorobeyva et al., 2015; Peter & Sommaruga, 2016), still virtually nothing is known about ciliates and other planktonic members in such lakes.

Here, we studied the ciliate community at the species-level of one clear and one glacier-fed alpine lake and the possible drivers for their species distribution against the background of potentially damaging effects of both high UVR transparency and high loads of glacial mineral particles. We expected to find different ciliate communities in respect to the prevailing environmental conditions. We sampled two adjacent alpine lakes located in the Austrian Central Alps, including one glacier-fed turbid (Faselfadsee 3, hereafter Lake FAS 3) and one clear lake (Faselfadsee 4, hereafter Lake FAS 4), for biotic (ciliates, bacterio-, phyto- and zooplankton) and physicochemical parameters on three occasions in two consecutive years.

**Methods**

**Study sites**

The Faselfad (FAS) lake area comprises a group of six adjacent lakes situated at altitudes between 2263 and 2620 m a.s.l. in the western Austrian Central Alps (see Figure S1 in Supporting Information). All six lakes (Lake FAS 1–6) are located below one retreating glacier, the ‘Faselfadferner’. Here, we sampled the glacial turbid Lake FAS 3 and the clear Lake FAS 4 during the ice-free seasons on 12 July 2010 and 5 July 2011, and on 29 August 2011. Lake FAS 3 is located about 200 m below the glacier and is fed by meltwater from the glacier and runoff from the catchment, resulting in high loads of particles in the pelagial (Table 1), while Lake FAS 4 is clear and fed by seepage. Up to our knowledge, both lakes are fishless and cladoceran-free (for more details, see Sommaruga & Kandolf, 2014; Kammerlander et al., 2015; Peter & Sommaruga, 2016).

**Lake sampling and in situ measurements**

Water samples were collected with a 5-L Schindler-Patalas sampler from an inflatable boat above the deepest point of the lakes at 0, 1, 2, 4, 6, 8, 10, 12 and either 13 m (Lake FAS 4) or 14 m (Lake FAS 3) depth around midday (within 3 h of solar noon). For ciliates, subsamples were taken (0.2–1.0 L total volume per sample and depth; triplicates) and immediately preserved with 5% Bouin’s solution (final concentration). Further subsamples were fixed for phytoplankton (100 mL total volume per depth) with acidified Lugol’s solution, for bacteria (50 mL total volume per depth; triplicates) with 2% formalin (final concentration) and for zooplankton (5 L per depth; triplicates). Zooplankton was concentrated on a 45-µm mesh, transferred into 100 mL glass flasks and fixed with 4% formalin (final concentration).
Table 1  Geographic location, altitude, lake area, maximum depth \((Z_{\text{max}})\), weather conditions, physicochemical parameters including temperature, \(pH\), conductivity, dissolved organic carbon (DOC), dissolved phosphorus (DP), total phosphorus (TP), dissolved nitrogen (DN), optical properties such as turbidity, diffuse attenuation coefficients at 320 nm \((K_{d\ 320})\) and of photosynthetically active radiation \((K_{d\ PAR})\), depth of 1% of surface irradiance for 320 nm UV \((Z_{\%\ 320})\) and PAR \((Z_{\%\ PAR})\) and fraction of the water column to which 1% of the surface irradiance at 320 nm \((Z_{\%\ 320}\ Z_{\max})\) and of PAR \((Z_{\%\ 320}\ Z_{\max})\) penetrated and chlorophyll \(a\) (chl \(a\)) of Lakes FAS 3 (turbid) and FAS 4 (clear). Data are shown as means, minimum and maximum values. n.d., not determined. For additional parameters see Table S1 in Supporting Information.

| Lake         | FAS 3 (turbid) | FAS 4 (clear) |
|--------------|---------------|---------------|
| Latitude     | 47°N 04' 15"  | 47°N 04' 27"  |
| Longitude    | 10°E 13' 15"  | 10°E 13' 34"  |
| Altitude (m a.s.l.) | 2414          | 2416          |
| Lake area (km\(^2\)) | 0.02          | 0.02          |
| \(Z_{\max}\) (m)  | 17.0          | 15.0          |

| Sampling date | July 2010 | July 2011 | August 2011 | July 2010 | July 2011 | August 2011 |
|---------------|-----------|-----------|-------------|-----------|-----------|-------------|
| Weather       | Cloudy    | Cloudy    | Sunny       | Cloudy    | Cloudy    | Sunny       |
| Temperature (°C) | 5.4 (3.1–7.1) | 5.7 (4.6–7.2) | 6.6 (5.0–8.6) | 7.0 (5.8–9.3) | 7.4 (5.9–8.4) | 10.0 (9.1–10.6) |
| pH            | 7.5 (7.1–7.8) | 7.6 (7.5–7.8) | 8.0 (7.2–8.8) | 7.2 (7.1–7.3) | 7.4 (7.4–7.5) | 7.3 (7.2–7.4) |
| Conductivity (\(\mu S\  cm^{-1}\)) | 39.7 (32.8–53.4) | 37.6 (37.0–39.7) | 43.2 (42.0–45.8) | 51.6 (48.9–54.0) | 50.6 (49.9–51.8) | 49.8 (48.8–51.5) |
| DOC (\(\mu g\  L^{-1}\)) | 226.9 (182.0–282.0) | 165.3 (148.0–206.0) | 266.9 (220.0–318.0) | 245.1 (228.0–311.0) | 299.2 (245.0–364.0) | 305.1 (268.0–374.0) |
| DP (\(\mu g\  L^{-1}\)) | 1.9 (1.5–2.4) | 1.3 (1.0–1.9) | 1.3 (1.3–1.6) | 1.0 (0.8–1.1) | 0.7 (0.4–0.9) | 0.7 (0.6–0.9) |
| TP (\(\mu g\  L^{-1}\)) | 7.3 (5.9–9.3) | 6.9 (5.0–9.1) | 8.8 (7.3–10.6) | 2.3 (1.8–3.4) | 2.2 (1.4–4.9) | 2.5 (1.7–5.7) |
| DN (\(\mu g\  L^{-1}\)) | 168 (143–184) | 168 (162–178) | 145 (112–199) | 210 (202–230) | 178 (167–183) | 172 (170–176) |
| Turbidity (NTU) | 3.2 (1.0–4.5) | 4.4 (3.9–4.6) | 8.6 (5.7–11.8) | 0.2 (0.1–0.3) | 0.2 (0.2–0.3) | 0.2 (0.02–0.3) |
| \(K_{d\ 320}\) \(\mu m\) | n.d. | 1.5 | 3.9 | n.d. | 0.2 | 0.3 |
| \(K_{d\ PAR}\) \(\mu m\) | n.d. | 0.5 | 1.3 | n.d. | 0.1 | 0.1 |
| \(Z_{\%\ 320}\) \(\mu m\) | n.d. | 3.1 | 1.2 | n.d. | 21.3 | 15.9 |
| \(Z_{\%\ PAR}\) \(\mu m\) | n.d. | 8.5 | 3.7 | n.d. | 41.6 | 33.1 |
| \(Z_{\%\ 320}\) \(\mu m\) \(Z_{\max}\) | n.d. | 0.2 | 0.1 | n.d. | 1.4 | 1.1 |
| \(Z_{\%\ PAR}\) \(Z_{\max}\) | n.d. | 0.5 | 0.2 | n.d. | 2.8 | 2.2 |
| Chl \(a\) (\(\mu g\  L^{-1}\)) | 3.5 (0.4–7.8) | 1.3 (0.4–2.2) | 8.0 (0.2–18.7) | 0.8 (0.2–1.7) | 0.6 (0.4–0.7) | 1.9 (1.5–3.1) |

In addition, living ciliates were collected by vertical net hauls (10 \(\mu m\) mesh size) and the water was prescreened through a 250-\(\mu m\) net to remove large predatory zooplankton. These samples were stored in cooling boxes at \textit{in situ} temperature.

Furthermore, subsamples were collected for the analyses of dissolved organic carbon (DOC), turbidity, water chemistry, photoprotective compounds (i.e. mycosporine-like amino acids; MAAs) and chlorophyll \(a\) (chl \(a\)). Temperature was measured with a thermometer attached inside the water sampler. The downwelling irradiance at 305, 320, 340 and 380 nm (UVR) and between 400 and 700 nm (photosynthetically active radiation, PAR) was measured in July and August 2011 with a profiling ultraviolet radiometer (PUV-501B, Biospherical Instruments).

Sample processing

\textit{Ciliate abundance and taxonomy.} Living ciliates were identified using an Olympus BX50 microscope (bright field, differential interference contrast) following the taxonomic keys of Foissner et al. (1991), Foissner, Berger & Kohmann (1994), Foissner et al. (1995), Foissner, Berger & Schaumburg (1999) and Berger (1999, 2006). Preserved samples were filtered onto cellulose nitrate filters (Sartorius, Goettingen; 0.8 \(\mu m\) pore size, with integrated counting grid) and stained following the quantitative protargol staining method (QPS) after Skibbe (1994) and Pfister, Sonntag & Posch (1999). From these permanent preparations, ciliates were identified and quantified under bright field (400–1000× magnification). The taxonomical assignment followed the revision of Adl et al. (2012).

\textit{Phytoplankton abundance, taxonomy and chlorophyll \(a\).} For qualitative and quantitative phytoplankton analysis, samples were settled in Utermöhl chambers and observed with an inverted microscope (Reichert-Jung, Vienna) at four different magnifications (48×, 128×, 320× and 504×). For more details see Rott (1981).

For the analysis of chl \(a\), 0.9–2.5 L were pre-filtered through a 100-\(\mu m\) net to remove zooplankton,
concentrated on pre-combusted glass-fibre filters (Whatman, GF/F) and stored at −20 °C. Pigments were extracted in the dark with 90% alkaline acetone (v:v) at 4 °C for 24 h. After sonication (1 min at 35 W; Sonoplus, HD2070, Bandelin, Berlin) on ice, the extracts were cleared by filtration through glass-fibre filters (Whatman, GF/F) and the absorbance was measured before and after acidification in a double-beam spectrophotometer (Hitachi U-2001, Inula, Vienna). The chl a concentration was calculated according to the formula of Lorenzen (1967) correcting for phaeopigments.

**Bacterial abundance.** Samples were filtered onto black polycarbonate filters (Merck Millipore GBTP, Darmstadt; 0.22 μm pore size), stained with DAPI (4', 6'-diamidino-2-phenylindole; Molecular Probes, Eugene) and semi-automatically counted (Zeiss AxioImager; software IMI 1.5 and Zeiss Axiovision; Zeder 2005–2010; http://www.technobiology.ch). On average, ~20 images per filter were selected for counting (at least 500 cells) using a 63× objective.

**Zooplankton abundance and taxonomy.** Zooplankton was identified following the taxonomic keys of Einsle (1993), Voigt & Koste (1978a,b). The different life stages (i.e. nauplii, copepodid CI–CIV and copepodid CIV–adult females and males) of *Cyclops abyssorum tatricus* as well as the number of rotifers were counted with an inverted microscope (Leitz, Labovert) after sedimentation in Utermöhl chambers at a magnification of 100×. For biomass estimation, the *C. abyssorum tatricus* body length was measured at a magnification of 40× or 100×, and the body length and width of the rotifers were measured at 320×. Biomass was then calculated according to Bottrell et al. (1976) (rotifers) and Praptokardiyi (1979) (*C. abyssorum tatricus*).

**Physicochemical parameters**

For dissolved organic carbon (DOC), water was stored in pre-combusted (2 h at 500 °C) glass bottles with glass stoppers in the dark at 4 °C and subsequently filtered through pre-combusted (2 h at 450 °C) glass-fibre filters (Whatman GF/F, Maidstone; rinsed with 20 mL of Milli-Q water and 10 mL of lake water). DOC was measured with a high-temperature catalytic oxidation method (Shimadzu TOC-V CPH, Vienna; equipped with a Shimadzu platinized-quartz catalyst for high sensitivity analysis).

Analyses of pH, conductivity, ion concentrations and nutrients (Tables 1 and S1) were determined using standard protocols (Sommaruga-Wögrath et al., 1997).

Turbidity was measured using a portable turbidimeter (WTW Turb 430 T, Weilheim), which mainly detects the scattering of ‘white light’ by small particles such as glacial flour (Sommaruga & Kandolf, 2014).

**MAAs analysis**

For the analysis of MAAs in the seston, 0.25–0.5 L lake water were prefilted through a 100-μm mesh to remove zooplankton, filtered onto pre-combusted glass-fibre filters (Whatman GF/F) and frozen at −80 °C. The extraction, analysis and calculation followed the protocol of Tartarotti & Sommaruga (2002).

**Statistical analyses**

The diversity index for the ciliate community in each lake was calculated according to Shannon–Wiener (Mühlenberg, 1993). We used Canoco 5.04 (ter Braak & Šmilauer, 2012) in order to visualise the variation among the lakes at the three sampling events. We used principal component analyses (PCA) on independent data sets: (i) based only on the environmental data (Table 1 and S1, Fig. 1) and (ii) based solely on the ciliate community (abundance data, Fig. 2). We then used redundancy analyses (RDA) with forward selection of statistically significant parameters to analyse the variation in the ciliate species pattern in relation to the environmental parameters. The first parameter included within the forward selection process typically has the highest significant impact on the ciliate abundance distribution. Any parameter covariant with an already selected parameter will then not be available for selection within the next forward selection step (see ter Braak & Šmilauer, 2012, pages 339 ff). In our analyses, several of the explanatory variables were auto-correlated, especially PAR and UVR (all measured wavelengths, Figure S2). Although the stepwise forward selection process allows only for one of these variables to be selected, the effect of any co-varying parameters on the ciliate community is a combined impact and cannot be interpreted independently. In case that an alternative explanatory parameter is an equally plausible candidate for selection as major parameter, we state this in Table 2 together with the percentage this parameter would explain if chosen as the first parameter.

Redundancy analyses were chosen because of the short gradients observed in the ciliate data. The ciliate abundance data were log (y + 1) transformed, centred for analyses and the samples were standardised (Chord distance; Legendre & Gallagher, 2001). The arch effect in
the species data remained with and without Chord or Hellinger distance transformation. Explanatory variables were centred and standardised. The RDA was then calculated for samples with PAR and UVR measurements (i.e. July and August 2011; Figure S3) with the following data: (i) for all ciliates in each lake and (ii) for the dominant ciliate groups (*B. planctonicum*, *Askenasia cf. chlorelligera*, *Urotricha* spp., *Mesodinium cf. acarus*) within each lake. The following explanatory variables were included in the analyses: depth, turbidity, physicochemical parameters (Tables 1 and S1), the relative underwater solar irradiance (i.e. % intensity of UVR at 320 nm and of PAR at the respective depths), chl *a*-specific MAAs, abundance of bacteria, zooplankton (>1% of total abundance) including rotifers and copepods (nauplii, CI–CIII stages, CIV–adult), abundance and biovolume of phytoplankton (>5% of total biovolume) and chl *a*. Statistical significance of the explanatory parameters was assessed with 999 unrestricted Monte Carlo permutations and corrected for false discovery rate Benjamini–Hochberg.
Table 2 Major drivers for the total abundance of ciliates and the dominant species (>1% of total mean abundance) assessed by RDA with forward selection of significant parameters ($P_{adj.} < 0.05$). The percentage of the amount explaining the variation in the composition of the ciliate community is indicated in brackets. Note that Mesodinium cf. acarus is present only in the glacial turbid Lake FAS 3 and that the samples only for which the depth profiles of ultraviolet radiation (UVR) and photosynthetically active radiation (PAR) measurements were available (i.e. July and August 2011) were included in the analyses. See also Figure S2 with a PCA of the abiotic parameters included in these analyses.

|                             | Clear lake – FAS 4 | Glacier-fed turbid lake – FAS 3 |
|-----------------------------|--------------------|----------------------------------|
| All ciliates                | Underwater solar irradiance (65% UV-B**)$^1$ | Underwater solar irradiance (41% PAR*) |
| Balanion planctonicum       | Underwater solar irradiance (67% UV-B**)$^1$ | No significant parameter |
| Askenasia cf. chlorelligera | Underwater solar irradiance (78% UV-B**$^1$ | Potential predators (46% rotifers*)$^3$ |
| Urotricha cf. furcata       | Potential predators (62% rotifers*)$^3$ | No significant parameter |
| Mesodinium cf. acarus       | –                  | Potential predators (78% zooplankton*) |

$P$ levels with Benjamini–Hochberg correction for false discovery rates during multiple testing: **<0.01, *<0.05.
Alternative factors: $^1$Zooplankton abundance (58%, **). $^2$Zooplankton abundance (63%, **). $^3$Phytoplankton abundance (46%, **).

Results

Physicochemical parameters

Although the lakes slightly differ in temperature (Lake FAS 3 <6 °C; Lake FAS 4 >7 °C) in both years, they showed a weak but at least temporally stable temperature stratification with a gradual decrease in temperature near the lake bottom (Fig. 3a–f; Table 1).

The PCA of the environmental data of the FAS lakes distinctly separates the samples from the clear and the glacial turbid lake (Fig. 1a,b). The PCA axis one reflects sulphate and magnesium versus turbidity, phosphorus (total and dissolved phosphorus), potassium and pH, with sulphate and magnesium being higher in the clear and the other parameters being higher in the glacial turbid lake. We additionally calculated a PCA taking into account UVR and PAR (available only for 2011, Figure S2). UVR and PAR are scoring negatively on PCA axis one, together with sulphate and magnesium, and opposing the trends of turbidity. Thus, the first PCA axis also reflects a gradient of the intensity of downwelling irradiance. At the same time, the single sampling events (except for the July samples in the glacial turbid Lake FAS 3) are also clearly separated along the second PCA axis which is not only reflecting weather conditions (sunny versus cloudy), but also nitrate and dissolved reactive silica versus dissolved organic carbon and calcium. Still, the difference in ordination space was larger between the two lakes than among the sampling occasions.

Ciliates

In both lakes, the dominant ciliates were the heterotrophic prostatomatsids B. planctonicum (Lake FAS 3 versus Lake FAS 4: 71% versus 60% on average, considering all sampling dates) and Urotricha cf. furcata (1% versus 37%), haptorids such as Askenasia cf. chlorelligera (4% versus 3%) and Mesodinium cf. acarus (22%; present only in Lake FAS 3) (Table S2, Figs 3a–f & 4). In the glacial turbid lake, most ciliates belonged to the Prostomatea, Phylopharyngea, Colpodea, Oligohymenophorea, Haptoria, Hypotrichia and Oligotrichia. In the clear lake, mainly Prostomatea, Haptoria and single individuals of Hypotrichia were identified (Table S2). The Shannon diversity index ($H_s$) was similar in both lakes, however, the $H_s$ of the glacial turbid Lake FAS 3 (0.75) was slightly higher, though not significant, when compared with Lake FAS 4 (0.69). By contrast, species richness and the mean abundance were higher in Lake FAS 3 than in the clear Lake FAS 4 (Table S2). The PCA analysis showed clearly that the ciliate communities were different in both lakes. Based on the abundance data, the lakes were separated along the first PCA axis reflecting the presence of Urotricha cf. furcata versus Mesodinium cf. acarus and Enchelys cf. simplex (Fig. 2a,b). The latter two species were found only in the glacial turbid lake. The second PCA axis stands for the presence of B. planctonicum. In contrast to the abiotic parameters, the ciliate composition was similar within each lake at the three sampling events. For both lakes, the relative underwater irradiance (i.e. % of UV-B and PAR at depth) appeared to be a major explanatory variable for the overall ciliate abundance pattern (Table 2). This was particularly the case in the clear lake, where the pattern of the abundant species was significantly explained by the UV-B irradiance at depth, such as for B. planctonicum and Askenasia cf. chlorelligera. However, the abundance of potential predators (e.g. rotifers, copepods) significantly explained the abundance distribution of Askenasia cf. chlorelligera and Mesodinium cf. acarus (present only in the glacial turbid lake FAS 3) in the glacial turbid Lake, but also of...
Urotricha cf. furcata and B. planctonicum in the clear lake (Table 2).

The vertical distribution pattern and the number of species were similar in both July profiles in Lake FAS 3 (Fig. 3a,b; Table S2). Balanion planctonicum dominated between 8 and 10 m depth, Mesodinium cf. acarus at 14 m and Urotricha cf. furcata between 0 and 8 m. In contrast, we observed less species in lower abundance in the August 2011 profiles (Fig. 3c). Concurrently, B. planctonicum and Askenasia cf. chlorelligera both occurred in the upper water layers, Enchelys cf. simplex between 8 and 10 m depth, while Mesodinium cf. acarus was evenly distributed along the water column. The vertical ciliate distribution within Lake FAS 4 was similar at all sampling dates with the dominant species found between 8 and 13 m depth. The total mean abundance was similar in July 2010 and 2011, but lower in August 2011 (Fig. 3d–f; Table S2).

Zooplankton

In both lakes, we found one copepod species, Cyclops abyssorum taticus, and the rotifers Polyartha dolichoptera, Keratella hiemalis, Keratella cochlearis and Notholca squamula were dominant (Table S3; Fig. 5a–f). The zooplankton community in Lake FAS 3 consisted mainly of copepods (67% of the relative abundance and 99% of the relative biomass of zooplankton) in July 2010, while rotifers made up 82–93% of the relative zooplankton abundance, but only 8–16% of the zooplankton biomass in July and August 2011. The relative abundance of the different copepod life stages was dominated by copepodid CIV–adult life stages (48% of the relative C. abyssorum taticus abundance and 80% of the relative biomass respectively) in July 2010, while nauplii dominated (69% of the relative abundance and 21% of the relative biomass of C. abyssorum taticus) in July 2011. The copepodid CI–CIII life stages contributed with ~15% to the relative abundance of C. abyssorum taticus (14–25% of the relative biomass) in all three sampling events. The rotifer community was composed of P. dolichoptera (22–69% of the relative abundance of rotifers and 24–73% of the relative biomass respectively), K. hiemalis (5–40% and 4–53%), Kelikkottia longispina (1–7% and 2%) and K. cochlearis (1–4% and <1%). Synchaeta spp. (1–9% and up 17%) and Filinia longiseta (~2% and 1%) were only abundant in July 2010, N. squamula in July and August 2011 (26–40% and 9–15%) and Ascomorpha sp. (1% and <1%) in August 2011. The highest rotifer abundance was found between 0 and 4 m in July 2010 and between 6 and 12 m in 2011, whereas the copepods were mainly found at greater depths (8–14 m) (Fig. 5a,b). In August 2011, the zooplankton was more homogeneously distributed over the whole water column (Fig. 5c).

In Lake FAS 4, the rotifers made up 92% of the relative zooplankton abundance (10% of the relative zooplankton biomass) in July 2010, while the copepods contributed with 46–50% to the zooplankton abundance.
Fig. 5 (a–f) Vertical distribution of zooplankton (rotifers and the copepod *Cyclops abyssorum tatricus*) in the glacial turbid Lake FAS 3 (a–c) and in the clear Lake FAS 4 (d–f) during the ice-free periods of 2010 and 2011. Error bars indicate standard deviation and note the different scale bars between sampling sites.

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and up 98% of the zooplankton biomass in 2011 (Table S3; Fig. 5d–f). Copepodid CI–adults were the most abundant life stages in July 2010 (50% of the relative abundance and 84% of the relative biomass of C. abyssorum tatricus), however, nauplii dominated in July 2011 (73% of the relative abundance of C. abyssorum tatricus and 28% of the relative biomass respectively) and the copepodid stages CI–CIII in August 2011 (61% of the relative abundance and biomass of C. abyssorum tatricus). Polyarthra dolichoptera dominated the rotifer community (94–97% of the relative abundance and up 99% of the relative biomass of rotifers) in this clear lake. Depending on the sampling date, K. hiemalis, K. cochlearis, K. longispina and N. squamula contributed between 2% and 6% to the relative abundance and biomass of rotifers. The copepods were mainly distributed close to the bottom of the lake at all sampling occasions, also the highest rotifer abundance was found near the lake bottom in July 2010 (Fig. 5d), but in 2011 most of the rotifers were distributed between 6 and 10 m (Fig. 5e,f).

**Phytoplankton**

In both lakes, the phytoplankton community differed considerably in respect to quantities and species composition with higher values in the glacial turbid Lake FAS 3. In this lake, the phytoplankton community was dominated by diatoms (Bacillariophyceae, 10–76% of total biovolume), cryptophytes (22–53% of total biovolume) and chlorophytes (0.1–30% of total biovolumes; Fig. 6a–c; Table S4; Figure S4a–c). The highest quantities of phytoplankton were found in August 2011, when diatoms dominated and reached a peak biovolume of >4 mm³ L⁻¹ (Table S4). In July 2010, the cryptophycean flagellate Plagioselmis nanoplanctica was dominant (53% of total biovolume) in the lower part of the water column, along with the needle-shaped Koliella sp. (Chlorophyta) known from snow habitats which contributed up to 30%. The relative dominance of the planktonic diatom Fragilaria tenera (Bacillariophyceae) was lowest in July 2010 and highest in August 2011 (10–76% of total biovolume).

In all samples of Lake FAS 4, mainly flagellated chrysophytes (29–44% of total biovolume), dinoflagellates (28–34% of total biovolume) and the diatom F. tenera (5–20% of total biovolume) were common (Fig. 6d–f; Table S4; Fig. S4d–f). The cryptophyte Plagioselmis nanoplanctica contributed with only ~5% to the total biovolume, whereas the coccal green alga Coenococcus sp. reached up to 19% in August 2011.

Overall, the phytoplankton biovolume and abundance depth distribution was closely related to chl a values in both lakes and at all sampling occasions (Fig. 6 and Figure S4). In both lakes, we observed a strong inverse depth distribution of chl a and phytoplankton with highest values in the lower water column, especially in July 2010 and 2011 (Fig. 6 and Figure S4). Only in the glacial turbid Lake FAS 3, we observed higher values of both chl a and phytoplankton biovolume in the upper water column in August 2011 (Figs 6c & S4c).

**Bacteria**

In Lake FAS 3, the bacterial abundance ranged from 0.5 to 1.1 × 10⁶ cells mL⁻¹ (Fig. 6a–c). In July 2010 and 2011, the bacterial abundance peaked near the lake bottom (10–14 m; Fig. 6a,b), while the highest abundance was found in the upper 0–4 m in August 2011 (Fig. 6c). In Lake FAS 4, the vertical distribution of bacteria showed a clear pattern with the lowest abundance found in the upper water layers and the highest in the deeper zone between 8 and 13 m with 3.8–4.7 × 10⁵ cells mL⁻¹ (Fig. 6d–f).

**Discussion**

Recent morphological and molecular studies have shown that glacial systems harbour a high microbial diversity including particle-associated ciliate groups among others (Anesio & Laybourn-Parry, 2012; Mieczan et al., 2013; Lazzaro, Risse-Bühl & Brankatschk, 2015; Lutz et al., 2015). In our studied lakes, the species richness is significantly higher in the glacial turbid Lake FAS 3 than in the clear Lake FAS 4. Mainly particle-associated ciliates contributed to the species richness in the still glacially influenced turbid Lake FAS 3. Such turbid lakes underlie glacial meltwater fluctuations, several mixing events and episodically high loads of glacial particles (Sommaruga, 2015). During glacial melting periods, glacier streams and rain events ‘wash out’ the glacier forefield and may thus introduce diverse microorganisms into a lake. The latter process also influences clear lakes, though particle-associated protists likely do not find optimal conditions for attachment in the pelagic region. This may explain why particle-associated bacterivorous and detritus-feeding ciliate species were only present in the glacial turbid Lake FAS 3 (Table S2). Omnivory and life strategies, including cyst formation, probably enable these species to survive under disadvantageous conditions (Foissner et al., 1991, 1994; Berger, 1999, 2006; Foissner, 2008). We therefore

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Fig. 6 (a–f) Vertical distribution of the phytoplankton (biovolume of Bacillario-, Chloro-, Chryso-, Crypto- and Dinophyta), chlorophyll $a$ (chl $a$) and bacterial abundance in the glacial turbid Lake FAS 3 (a–c) and the clear Lake FAS 4 (d–f) during the ice-free periods of 2010 and 2011. Note the different scale bars between sampling sites.
argue that glacial turbid lakes offer suitable environmental conditions and resource niches for certain specialised protists that probably are excluded when this type of lake turns into a clear one.

In both lakes, the total ciliate abundance distribution pattern was significantly explained by the relative underwater solar irradiance (Table 2). In the clear Lake (FAS 4), the major factor explaining the distribution patterns of the dominant species was also the relative underwater solar irradiance, but this was not the case in the glacial turbid Lake (FAS 3) where zooplankton predation was more important (Table 2). When excluding the relative underwater solar irradiance from the redundancy analyses, phytoplankton abundance was the main explanatory variable for the total abundance of ciliates (31%, $P < 0.01$) in the glacial turbid Lake FAS 3. This makes sense since the dominant species found in this lake are known to be algivorous. Suitable food algae for most ciliates are unicellular chryso- and cryptophytes in a size range of 2–15 μm (Sonntag et al., 2006) such as Plagioselmis nanoplanctica that was highly abundant in the glacial turbid, but not in the clear lake (Fig. 6 and Figure S4, Table S4). In the glacial turbid Lake FAS 3, the abundance (and biomass) of phytoplankton, as well as the chl a concentration, were higher than in Lake FAS 4, especially in August 2011 (Fig. 6 and Figure S4, Table S4). Such high chl a peaks are not unusual in glacially influenced alpine lakes (Tolotti & Cantonati, 2000). Glacial meltwater can be a source of nutrients, especially phosphorus adsorbed to silt particles. Although only a small fraction of the phosphorus seems to be bio-available, this input may trigger periods of production in glacier-fed turbid lakes (Hodson, 2006; Mindl et al., 2007; Hodson et al., 2008; Sommaruga, 2015). Certain phytoplankton groups such as cryptophytes are well known to be adapted to low-light conditions because they possess a unique light-harvesting pigment complex in their photosynthetic apparatus (Gantt, Edwards & Provasoli, 1971; Spear-Bernstein & Miller, 1989; Grossman et al., 1993; Gervais, 1997) and also, Teubner et al. (2001) observed higher chl a concentrations in a smaller size fraction of algae (<10 μm) at low-light intensities. Interestingly, the cold- and low-light-adapted green alga Koliella sp. (found only in the glacial turbid Lake FAS 3; Table S4) is equipped with a highly efficient photosystem that quickly reacts to light fluctuations (La Rocca et al., 2015). This effective adaptation could be advantageous in light-limited lakes such as the glacial turbid Lake FAS 3.

In both FAS lakes, the ciliate community was dominated by the common euplanktonic heterotrophic ciliates B. planctonicum and Urotichia spp. (Table S2; Weisse & Müller, 1998; Wille et al., 1999; Sonntag et al., 2006, 2011). These species are known to be omnivores and to have an optimal growth strategy (i.e. K-strategy in B. planctonicum and r-strategy in Urotichia spp.; Weisse et al., 2001). This probably explains their success in alpine lakes, even if survival strategies during unfavourable periods such as cyst formation are unknown. Balanion planctonicum was highly abundant particularly in the glacial turbid Lake FAS 3 and we speculate that a higher phytoplanктon abundance is responsible for this. Furthermore, this species is known to be an efficient and selective feeder on small flagellates (Weisse & Müller, 1998). The different depth preferences of both prostomatid species (Fig. 3a–f) suggest an interspecific interaction probably competition for food, as found in other alpine lakes (Sonntag et al., 2011).

As mentioned above, in the clear lake, the abundance pattern of B. planctonicum and the mixotrophic Askenasia cf. chlorelligera were significantly explained by the relative underwater solar UVR (Table 2). Balanion planctonicum is sensitive to surface UVR (Sonntag et al., 2011), which may explain its preference for deeper water layers (Fig. 3d–f). Mixotrophic ciliates such as Askenasia cf. chlorelligera can accumulate sunscreen compounds known as mycosporine-like amino acids (MAAs) that are crucial for survival under high levels of UVR (Sonntag, Summerer & Sommaruga, 2007; Summerer, Sonntag & Sommaruga, 2008). Although we were not able to measure MAAs in this species, MAA concentrations in seston were significantly higher in FAS 4 than in FAS 3 reflecting the different degree of UV stress for planktonic organisms in these systems (Tartarotti et al., 2014). In our study, the abundance of ciliate species bearing algal symbionts and described or known as mixotrophic was higher in the glacial turbid than in the clear lake (Table S2). The reduced UV stress and the capability to switch from auto- to heterotrophy may be advantageous in glacial turbid, but at the same time light-limited habitats (Hinder et al., 1999; Sommaruga, 2015).

The abundance of predatory zooplankton significantly explained the abundance pattern of Mesodinium cf. acarus and Askenasia cf. chlorelligera in the glacial turbid lake, and of Urotichia cf. furcata and B. planctonicum in the clear lake (Table 2). The copepod C. abyssorum tatricus and rotifers such as P. dolichoptera and Keratella spp. are commonly found in Alpine lakes (Table S3; Tartarotti et al., 1999; Obertegger, Flaim & Sommaruga, 2008; Tiberi et al., 2013). Beside algae, ciliates are also in their food spectrum (Arndt, 1993; Wickham, 1995; Weisse &
1962  B. Kammerlander et al.

Frahm, 2002). However, ciliates are equipped with several defensive mechanisms/strategies, such as changing their morphology by sensing chemical cues (kairomones), repel an aggressor with extrusomes or performing escape jumps (Weisse & Sonntag, 2016 and references therein). For example, prostatomad and haptorial ciliates have characteristic swimming behaviour including spinning and jumping plus a motionless moment allowing them not only to prey efficiently, but also to avoid zooplankton predation (Foissner et al., 1999 and references therein; Jakobsen, Everett & Strom, 2006). Predatory zooplankton react on moving prey (e.g. Bruno, Andersen & Klørboe, 2012) and typical swimming patterns as mentioned above could be seen as an effective escape strategy (Wickham, 1995; Weisse & Frahm, 2002; Weisse & Sonntag, 2016).

In conclusion, our study unveiled an unexpected highly diverse planktonic protistan community in a glacier-fed turbid lake. Adaptive life strategies and specific traits seem to be a prerequisite for them to thrive in these extreme habitats. Therefore, it is important to include species-specific traits, as well as food and predator–prey interactions in future studies.

Acknowledgments

We thank J. Franzoi, G. Larsen and S. Morales-Gomez for chemical analyses and bacterial analyses, as well as H. W. Breiner, G. Kandolf, P. Kirschner, D. Pöll, K. Schwab and T. Stoeck for help during field work, C. Grubbauer and U. Scheffel for help during lab work and H. Berger for identifying the hypotrich ciliates. We thank two anonymous reviewers for critical comments. The study was financed by the Austrian Science Fund (FWF P21013-B03 and I2238-B25 to RS and TWF UNI-0404/140 to BT), the University of Innsbruck (to BT) and by a Doctoral Fellowship of the Austrian Academy of Sciences (OEAW, DOC-fForte 22883 to BK) and of the University of Innsbruck (2012/3/BIO9 to BK).

References

Adl S.M., Simpson A.G.B., Lane C.E., Lukes J., Bass D., Bowser S.S. et al. (2012) The revised classification of eukaryotes. Journal of Eukaryotic Microbiology, 59, 429–493.

Anesio A.M. & Laybourn-Parry J. (2012) Glaciers and ice sheets as a biome. Trends in Ecology and Evolution, 27, 219–225.

Arndt H. (1993) Rotifers as predators on components of the microbial web (bacteria, heterotrophic flagellates, ciliates) – a review. Hydrobiologia, 255/256, 231–246.

Berger H. (1999) Monograph of the Oxytrichidae (Ciliophora, Hypotrichia). Monographiae Biologicae, 78, 1–1080.

Berger H. (2006) Monograph of the Urostyloidae (Ciliophora, Hypotrichia). Monographiae Biologicae, 85, 1–1304.

Bottrell H.H., Duncan A., Gliwicz Z.M., Grygierek E., Herzig A., Hillbricht-Ilksowska A. et al. (1976) A review of some problems in zooplankton production studies. Norwegian Journal of Zoology, 24, 419–456.

Bowser S.S. (2012) The revised classification of the Ciliophora, Hypotricha). Monographiae Biologicae, 255/256, 1–1080.

Bottrell H.H., Duncan A., Gliwicz Z.M., Grygierek E., Herzig A., Hillbricht-Ilksowska A. et al. (1976) A review of some problems in zooplankton production studies. Norwegian Journal of Zoology, 24, 419–456.

Canoco Reference Manual and User’s Guide: Software for Ordination (version 5.0). Microcomputer power, Itaca. Available at: http://www.canoco.com.

Carriwick J.L. & Tweed F.S. (2013) Proglacial lakes: character, behaviour and geological importance. Quaternary Science Reviews, 78, 34–52.

Einsle U. (1993) Crustacea: Copepoda: Calanoida und Cyclopoida. Gustav Fischer Verlag, Stuttgart, Jena, New York.

Filker S., Sommaruga R., Vila L. & Stoeck T. (2016) Protistan plankton communities of high-mountain lakes from three continents exhibit strong biogeographic patterns. Molecular Ecology, 25, 2286–2301.

Fischer J.M., Olsen M.H., Theodore N., Williamson C.E., Rose K.C. & Huang J. (2015) Diel vertical migration of copepods in mountain lakes: the changing role of ultraviolet radiation across a transparency gradient. Limnology and Oceanography, 60, 252–262.

Foissner W. (2008) Protist diversity and distribution: some basic considerations. Biodiversity and Conservation, 17, 235–242.

Foissner W., Berger H., Blatterer H. & Kohmann F. (1995) Taxonomische und ökologische Revision der Ciliaten des Saprobiensystems. Band IV: Gymnostomatea, Loxodes, Suctoria. Informationsberichte des Bayer. Landesamtes für Wasserwirtschaft, München.

Foissner W., Berger H. & Kohmann F. (1994) Taxonomische und ökologische Revision der Ciliaten des Saprobiensystems. Band III: Hymenostomatea, Prostomatida, Nassulida. Informationsberichte des Bayer. Landesamtes für Wasserwirtschaft, München.

Foissner W., Berger H. & Schaumburg J. (1999) Identification and ecology of limnetic plankton ciliates. Informationsberichte des Bayer. Landesamtes für Wasserwirtschaft, München.

Foissner W., Blatterer H., Berger H. & Kohmann F. (1991) Taxonomische und ökologische Revision der Ciliaten des Saprobiensystems. Band II: Chytridophora, Oligotrichida, Hypotrichia, Colpoda. Informationsberichte des Bayer. Landesamtes für Wasserwirtschaft, München.

Gant E., Edwards M.R. & Provasoli L. (1971) Chloroplast biliproteins within intrathylakoidal spaces. Journal of Cell Biology, 48, 280–290.

Gervais F. (1997) Light-dependent growth, dark survival, and glucose uptake by cryptophytes isolated from a freshwater chemocline. Journal of Phycolgy, 33, 18–25.
Grossman A.R., Schaefer M.R., Chiang G.G. & Collier J.L. (1993) The phycobilisome, a light-harvesting complex responsive to environmental conditions. *Microbiological Reviews*, 57, 725–749.

Hinder B., Baur I., Hanselmann K. & Schanz F. (1999) Microbial food web in an oligotrophic high mountain lake (Jörli Lake III, Switzerland). *Journal of Limnology*, 58, 162–168.

Hodson A.J. (2006) Phosphorus in glacial meltwaters. In: Hodson A.J. (Ed. P.G. Knight), pp. 81–82. Blackwell Publishing, Malden.

Hodson A.J., Anesio A.M., Tranter M., Fountain A., Osborn M., Priscu J. et al. (2008) Glacial ecosystems. *Ecological Monographs*, 78, 41–67.

Hylander S., Jepson T., Lebret K., Von Einem J., Fagerberg T., Balseiro E. et al. (2011) Climate-induced input of turbid glacial meltwater affects vertical distribution and community composition of phyto- and zooplankton. *Journal of Plankton Research*, 33, 1239–1248.

Jakobsen H.H., Everett L.M. & Strom S.L. (2006) Hydrochemical signaling between the ciliate *Mesodinium pulex* and motile protist prey. *Aquatic Microbial Ecology*, 44, 197–206.

Kammerlander B., Breiner H.-W., Filker S., Sommaruga R., Sonntag B. & Stoeck T. (2015) High diversity of protistan plankton communities in remote high mountain lakes in the European Alps and the Himalaya mountains. *FEMS Microbiology Ecology*, 91, fiv010, doi: 10.1093/femsec/fiv010.

La Rocca N., Scuto K., Meneghesso A., Moro I., Rascio N. & Morosinotto T. (2015) Photosynthesis in extreme environments: responses to different light regimes in the Antarctic alga *Kollya antarctica*. *Physiologia Plantarum*, 153, 654–667.

Laurion I., Ventura M., Catalan J., Psenner R. & Sommaruga R. (2000) Attenuation of ultraviolet radiation in mountain lakes: factors controlling the among- and within-lake variability. *Limnology and Oceanography*, 45, 1274–1288.

Lazzaro A., Risse-Buhl U. & Brankatschk R. (2015) Molecular and morphological snapshot characterisation of the protist communities in contrasting alpine glacier forefields. *Acta Protozoologica*, 54, 143–156.

Legendre P. & Gallagher E. (2001) Ecologically meaningful transformations for ordination of species data. *Oecologia*, 129, 271–280.

Lischke B., Weithoff G., Wickham S.A., Attermeyer K., Grossart H.-P., Scharnweber K. et al. (2016) Large biomass of small feeders: ciliates may dominate herbivory in eutrophic lakes. *Journal of Plankton Research*, 38, 2–15.

Lorenzen C.J. (1967) Determination of chlorophyll and phaeo-pigments: spectrophotometric equations. *Limnology and Oceanography*, 12, 343–346.

Lutz S., Anesio A.M., Edwards A. & Benning L.G. (2015) Microbial diversity on Icelandic glaciers and icecaps. *Frontiers in Microbiology*, 6, 307. doi: 10.3389/fmicb.2015.00307.

Mieczan T., Görniak D., Świątecki A., Zdanowski M. & Tarlowa-Kukury M. (2013) The distribution of ciliates on Ecology Glacier (King George Island, Antarctica): relationships between species assemblages and environmental parameters. *Polar Biology*, 36, 249–258.

Mindl B., Anesio A.M., Meier K., Hodson A.H., Laybourn-Parry J., Sommaruga R. et al. (2007) Factors influencing bacterial dynamics along a transect from supraglacial runoff to proglacial lakes of a high Arctic glacier. *FEMS Microbiology Ecology*, 59, 307–317.

Mitchell D.L. & Karentz D. (1993) The induction and repair of DNA photodamage in the environment. In: *Environmental UV Photobiology* (Eds A.R. Young, L.O. Björn, J. Moan & W. Nultsch), pp. 345–377. Springer, New York.

Mühlenberg M. (1993) *Freilandökologie*, 3rd edn. Quelle und Mayer Verlag, Heidelberg, Wiesbaden.

Obertegger U., Flaim G. & Sommaruga R. (2008) Multifactorial nature of rotifer water layer preferences in an oligotrophic lake. *Journal of Plankton Research*, 30, 633–643.

Peter H. & Sommaruga R. (2016) Shifts in diversity and function of lake bacterial communities upon glacier retreat. *The ISME Journal*, 10, 1545–1554. doi: 10.1038/ismej.2015.2.

Pfister G., Sonntag B. & Posch T. (1999) Comparison of a direct live count and an improved quantitative protargol stain (QPS) in determining abundance and cell volumes of pelagic freshwater protozoa. *Aquatic Microbial Ecology*, 18, 95–103.

Posch T., Eugster B., Pomati F., Pernthaler J., Pittsch G. & Eckert E.M. (2015) Network of interactions between ciliates and phytoplankton during spring. *Frontiers in Microbiology*, 6, 1289. doi: 10.3389/fmicb.2015.01289.

Praptokardiyo K. (1979) *Populationsdynamik und Produktion von Cyclops abyssorum tauricus* (Kozinski 1927) im Gossenköllesee (2413 m ü.N.N., Kühtai, Tirol). PhD Thesis, University of Innsbruck, Innsbruck.

Rott E. (1981) Some results from phytoplankton counting intercalibrations. *Schweizerische Zeitschrift für Hydrologie*, 43, 34–61.

Saros J.E., Interlandi S.J., Doyle S., Michel T.J. & Williamson C.E. (2005) Are the deep chlorophyll maxima in alpine lakes primarily induced by nutrient availability, not UV avoidance? *Arctic, Antarctic, and Alpine Research*, 37, 557–563.

Skibbe O. (1994) An improved quantitative protargol stain for ciliates and other planktonic protists. *Archiv für Hydrobiologie*, 130, 339–347.

Sommaruga R. (2001) The role of solar UV radiation in the ecology of alpine lakes. *Journal of Photochemistry and Photobiology B: Biology*, 62, 35–42.
Sommaruga R. (2015) When glaciers and ice sheets melt: consequences for planktonic organisms. *Journal of Plankton Research*, 37, 509–518.

Sommaruga R. & Kandolf G. (2014) Negative consequences of glacial turbidity for the survival of freshwater planktonic heterotrophic flagellates. *Scientific Reports*, 4, 1–5.

Sommaruga-Wögrath S., Koenig K.A., Schmidt R., Sommaruga R., Tessadri R. & Psenner R. (1997) Temperature effects on the acidity of remote alpine lakes. *Nature*, 387, 64–67.

Sommer U., Adrian R., De Senerpont Domis L., Elser J.J., Tartarotti B. & Sommaruga R. (2002) The effect of different phagotrophic ciliates and flagellates in an oligotrophic deep alpine lake: contrasting variability with seasons and depths. *Aquatic Microbial Ecology*, 43, 193–207.

Sonntag B., Summerer M. & Sommaruga R. (2007) Sources of mycosporine-like amino acids in planktonic *Chlorella*-bearing ciliates (Ciliophora). *Freshwater Biology*, 52, 1476–1485.

Sonntag B., Summerer M. & Sommaruga R. (2011) Factors involved in the distribution pattern of ciliates in the water column of a transparent alpine lake. *Journal of Plankton Research*, 33, 541–546.

Spear-Bernstein L. & Miller K.R. (1989) Unique location of the phycobiliprotein light-harvesting pigment in the Cryptophyceae. *Journal of Phycology*, 25, 412–419.

Summerer M., Sonntag B. & Sommaruga R. (2008) Ciliate-symbiont specificity of freshwater endosymbiotic *Chlorella* (Trebouxiiophyceae, Chlorophyta). *Journal of Phycology*, 44, 77–84.

Tartarotti B., Cabrera S., Psenner R. & Sommaruga R. (1999) Survivorship of *Cyclops abyssorum taticus* (Cyclopoida, Copepoda) and *Boeckella gracilipes* (Calanoida, Copepoda) under ambient levels of solar UV radiation in two high mountain lakes. *Journal of Plankton Research*, 21, 549–560.

Tartarotti B., Saul N., Chakrabarti S., Trattner F., Steinberg C.E.W. & Sommaruga R. (2014) UV-induced DNA damage in *Cyclops abyssorum taticus* populations from clear and turbid alpine lakes. *Journal of Plankton Research*, 36, 557–566.

Tartarotti B. & Sommaruga R. (2002) The effect of different methanol concentrations and temperatures on the extraction of mycosporine-like amino acids (MAAs) in algae and zooplankton. *Archiv für Hydrobiologie*, 154, 691–703.

Teubner K., Sarobe Egiguren A., Vadrucci M.R. & Dokulil M. (2001) ¹⁴C photosynthesis and pigment pattern of phytoplankton as size related adaptation strategies in alpine lakes. *Aquatic Sciences*, 63, 310–325.
Wickham S.A. (1995) Trophic relations between cyclopoid copepods and ciliated protists: complex interactions link the microbial and classic food webs. *Limnology and Oceanography, 40*, 1173–1181.

Wille A., Sonntag B., Sattler B. & Psenner R. (1999) Abundance, biomass and size structure of the microbial assemblage in the high mountain lake Gosenköllesee (Tyrol, Austria) during the ice-free period. *Journal of Limnology, 58*, 117–126.

Williamson C.E., Fischer J.M., Bollens S.M., Overholt E.P. & Breckenridge J.K. (2011) Toward a more comprehensive theory of zooplankton diel vertical migration: integrating ultraviolet radiation and water transparency into the biotic paradigm. *Limnology and Oceanography, 56*, 1603–1623.

**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Table S1.** Additional physicochemical parameters of Lakes FAS 3 (turbid) and FAS 4 (clear).

**Table S2.** Species list of the ciliate community from the Lakes FAS 3 (turbid) and FAS 4 (clear).

**Table S3.** Species list of the zooplankton community from the Lakes FAS 3 (turbid) and FAS 4 (clear).

**Table S4.** Species list of the phytoplankton community from the Lakes FAS 3 (turbid) and FAS 4 (clear).

**Figure S1.** The study site Faselfad (FAS): The glacier-fed turbid Lake FAS 3 and the clear Lake FAS 4 are located at ~ 2416 m a.s.l. Photo: B. Kammerlander.

**Figure S2.** PCA of the environmental parameters in the glacial turbid Lake FAS 3 and the clear Lake FAS 4 during the sampling dates when changes in the underwater solar irradiance were measured (i.e. in July and August 2011). Together with the biological explanatory parameters (e.g. bacterio-, phyto- and zooplankton), these data were used in the redundancy analyses to explain the ciliate distribution. Note the distinct correlation of PAR and UVR. The impact of covariant environmental parameters on the ciliates is a combined effect and cannot be separated within the redundancy analyses (RDA). The singular, independent effect of each of these strongly auto-correlated variables on the distribution of the ciliates is marginal.

**Figure S3.** Depth profiles of ultraviolet radiation (UVR) at 305, 320, 340, 380 nm, and of photosynthetically active radiation (PAR) in the glacial turbid Lake FAS 3 (a, b) and in the clear Lake FAS 4 (c, d).

**Figure S4.** Vertical distribution of the phytoplankton community (abundance data of Bacillario-, Chloro-, Chryso-, Crypto- and Dinophyta), chlorophyll a (chl a) and bacterial abundance in the glacial turbid Lake FAS 3 (a–c) and the clear Lake FAS 4 (d–f) during the ice-free periods of 2010 and 2011. Note the different scale bars between sampling sites.

*(Manuscript accepted 5 August 2016)*