RESEARCH ARTICLE

High diversity of protistan plankton communities in remote high mountain lakes in the European Alps and the Himalayan mountains

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One sentence summary: Diversity of protistan communities in remote high mountain lakes in the European Alps and the Himalayan mountains suggests impact of glacier retreat on community structure and function.

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ABSTRACT

We analyzed the genetic diversity (V4 region of the 18S rRNA) of planktonic microbial eukaryotes in four high mountain lakes including two remote biogeographic regions (the Himalayan mountains and the European Alps) and distinct habitat types (clear and glacier-fed turbid lakes). The recorded high genetic diversity in these lakes was far beyond of what is described from high mountain lake plankton. In total, we detected representatives from 66 families with the main taxon groups being Alveolata (55.0% OTUs 97%), Stramenopiles (34.0% OTUs 97%), Cryptophyta (4.0% OTUs 97%), Chloroplastida (3.6% OTUs 97%) and Fungi (1.7% OTUs 97%). Centrohelida, Choanomonada, Rhizaria, Katablepharidae and Telonema were represented by <1% OTUs 97%. Himalayan lakes harbored a higher plankton diversity compared to the Alpine lakes (Shannon index). Community structures were significantly different between lake types and biogeographic regions (Fisher exact test, P < 0.01). Network analysis revealed that more families of the Chloroplastida (10 vs 5) and the Stramenopiles (14 vs 8) were found in the Himalayan lakes than in the Alpine lakes and none of the fungal families was shared between them. Biogeographic aspects as well as ecological factors such as water turbidity may structure the microbial eukaryote plankton communities in such remote lakes.

Keywords: diversity; alpine lakes; next-generation sequencing

INTRODUCTION

In the last years, molecular tools have seized biodiversity studies, and next-generation sequencing (NGS) approaches have become popular instruments to estimate protist diversity (Margulies et al., 2005). To date, most studies on microbial diversity including protists and fungi have focused on marine ecosystems (e.g. Alexander et al., 2009; Stoeck et al., 2010; Edgcomb et al., 2011; Bik et al., 2012; Stock et al., 2012). Even though the diversity of freshwater microbial plankton is presumably much higher than in the marine environment...
(Logares et al., 2009; Auguet, Barberán and Casamayor 2010; Barberán and Casamayor 2010; Barberán et al., 2011; Triadó-Margarit and Casamayor 2012), only few freshwater lakes were examined so far by using such molecular tools (e.g. Chen et al., 2008; Lefèvre et al., 2008; Steele et al., 2011; Charvet et al., 2012a; Charvet, Vincent and Lovejoy 2012b; Charvet, Vincent and Lovejoy 2014; Stoeck et al., 2014). Hitherto only very few sequence data are available from protistan plankton in high mountain lakes (e.g. Triadó-Margarit and Casamayor 2012).

Yet, these data emphasized high mountain lakes as diversity hotspots for (hitherto unknown) eukaryotic microbial plankton. Such a scarce knowledge on microbial eukaryotic plankton in high mountain lakes is unsatisfying considering the ecological importance of these organisms in the energy and carbon transfer within aquatic food webs (e.g. Azam et al., 1983; Sherr and Sherr 1988; Weisse and Müller 1998; Sonntag et al., 2006; Zingel et al., 2007).

Likewise, microscopy studies on protist diversity in high mountain lakes are scarce (Félik et al., 1999; Straškrabová et al., 1999; Wille et al., 1999; Sonntag, Summerer and Sommaruga 2011). The difficulties that occur in protist investigation by microscopy are their small sizes, few morphological characters available to identify especially flagellated species and low abundance particularly in high mountain lakes. Though, as protists include manifold groups that have specific demands on their environment concerning their nutrition or temperature, there is a need to identify them as exact as possible to obtain an overall picture of the food web interactions. To overcome such inconveniences, NGS approaches such as pyrosequencing (Margulies et al., 2005) are promising techniques to unravel the hidden diversity of microbes in environmental samples (Sogin et al., 2006; Caron et al., 2012). One major strength of NGS is the depth of sequencing which allows elucidating the local ‘rare biosphere’ (Sogin et al., 2006), a seed bank of low abundant taxa that may play a pivotal role in ecosystem response to environmental changes as well as in ecosystem stability and function(ing) (Pedróš-Alió 2007; Dawson and Hagen 2009; Stoeck and Epstein 2009).

In high mountain lakes organisms are confronted with short growing seasons, low food availability or high incident solar radiation (Sommaruga 2001; Rose et al., 2009). Characteristically, high mountain lakes can be very transparent or turbid dependent on the connection to a glacier. The lower the particle load and the concentration of chromophoric dissolved organic matter, the more transparent are such lakes. Lake transparency is crucial for the biota because it determines the penetration of photosynthetically active radiation and of ultraviolet radiation (UVR). For example, in clear high mountain lakes the potentially harmful wavelengths of UVR can reach down to the lake bottom (Morris et al., 1995; Laurion et al., 2000; Sommaruga and Augustin 2006). As for clear high mountain lakes, several surveys have concentrated on the effects of UVR onto the planktonic community and their key players, but it is largely unknown how sensitive species from turbid alpine lakes are to UVR. However, a recent comparative study with one copepod species living in clear and turbid alpine lakes suggests that photoprotection and DNA repair mechanisms are important in adapting to the shift in water transparency (Tartarotti et al., 2014).

In our study, we sampled four different lakes, including two biogeographic regions (Austrian Central Alps and the Himalayan mountains, Nepal) and two lake types (clear vs glacier-fed turbid lakes) using massively parallel tag sequencing (pyrosequencing) of the hypervariable V4 region of the small subunit ribosomal DNA to determine the plankton diversity. The reasoning for these habitat choices is among others to maximize the extent of diversity that can be identified from different mountain ranges and different lake types. Our data shows a much higher diversity of fungal and protistan plankton as known from previous diversity studies in high mountain lakes, with ca. 60% of our detected sequences showing a high genetic divergence to deposited sequence data.

**MATERIALS AND METHODS**

**Faselfad lakes**

The study site Faselfad (FAS) is located in the western Austrian Central Alps and comprises a group of six adjacent lakes situated between 2263 and 2620 m above sea level (a.s.l.). All six lakes originate from one glacier, the ‘Faselfadferner’, and mainly differ in altitude and water transparency. Out of these, we selected one clear (FAS 4) and one glacier-fed turbid lake (FAS 3) (Fig. 1, Table 1). FAS 3 is located approximately 200 m below the glacier and fed by glacial melt water enriched with high particle loads, so-called ‘glacial flour’ and partially by water from the catchment. The clear lake FAS 4 has lost its connectivity to the glacier and is fed by seepage from its catchment (Sommaruga and Kandolf 2014).

On 29 August 2011, water samples were collected with a 5-L Schindler–Patalas sampler at the deepest point of each lake from an inflatable boat. Mixed water samples (10 L) in a ratio of 1:1.1 were taken from the uppermost meters (0, 1 and 2 m) and 1 m above and below the chlorophyll a (chl a) maximum. This sampling strategy was based on previous samplings along vertical depth gradients indicating that most of the taxa were found around the chl a maximum (Kammerlander et al., unpublished). Prior to sampling, the chl a maximum in the water column was detected with a Backscat I-Fluorometer (Haardt, model 1101.1, excitation 380–540 nm, emission 685 nm). Further, subsamples were taken for the analysis of abiotic parameters, i.e. turbidity, conductivity, pH, total phosphorus (TP), dissolved organic carbon (DOC), nitrate (NO3-N) and ammonium (NH4-N). The chemical parameters were measured in the laboratory of the Institute of Ecology at the University of Innsbruck as described in Sommaruga-Wöggrath et al. (1997). For measuring the turbidity a handheld turbidimeter (WTW Turb 430 T) was used. For details, see Sommaruga and Kandolf (2014). The DOC analyses were measured with a high temperature catalytic oxidation method (Shimadzu TOC-VCPH–total organic carbon analyzer). For details, see Laurion et al. (2000) and Sommaruga and Augustin (2006). For nucleic acid extractions, triplicate water samples were collected in clean plastic carboys and 2–3 L each was drawn onto Durapore membranes (0.65 μm, 47 mm, Millipore) using a peristaltic pump. Filters were frozen immediately in liquid nitrogen. Samples were stored at −20° C until DNA extraction.

**Himalaya lakes**

One glacier-fed turbid (HL 5) and one clear (HL 15) lake were sampled in the Khumbu Valley region (Nepal) close to Mount Everest on 8 and 9 October 2004 (Fig. 1, Table 1). For more details on this study site, see Tartarotti et al. (1998) and Sommaruga and Casamayor (2009).

Samples were collected from a boat and integrated over the water column. Filters for nucleic acid extraction were prepared as described above and transported to Innsbruck in liquid nitrogen.
Pyrosequencing

DNA isolation and construction of pyro-amplicon libraries

DNA was isolated directly from the Durapore membranes using Qiagen’s AllPrep kit according to the manufacturer’s instructions. The samples (filters) were extracted and pooled. From these extracts, the hypervariable V4 region of the 18S rRNA gene was amplified using the eukaryote specific primer pair TAReukV4F and TAReukREV (5′-ACTTTCGTTCTTGATYRA-3′, Stoeck et al., 2010) yielding ca. 500 basepair (bp) fragments. To distinguish the different samples in downstream processes, the V4 forward primer was tagged with specific 10-bp identifiers (MIDs) at the 5′-end. The PCR protocol followed the description of Stoeck et al. (2010). To minimize PCR-bias, we ran three individual reactions per sample. The resulting PCR products were purified (MinElute PCR purification kit, Qiagen, Germany) and pooled prior to sequencing. The V4-DNA amplicon libraries were sequenced on one-half of a PicoTiter Plate with a Roche FLX GS20 sequencer and the Titanium chemistry (FAS lakes: EnGenCore, SC, USA; HL lakes: LGC Genomics, Berlin, Germany). and a minimum length of 300 bp were kept. The remaining sequences were then checked for chimeras and clustered at different threshold levels (90, 95, 96, 97, 98, 99 and 100%) using the OTUpipe script (Edgar et al., 2011) implemented in QIIME. The OTUpipe script was done with the following adjustments: -m usearch -l –reference_chimera_detection -j 1 –s 0.xx –word_length yy. The word length value was calculated according to an equation given in Edgar et al. (2011). For our pyro-amplicons, a value of 64 was used. For taxonomic assignments, one representative sequence (longest) from each OTU was extracted and analyzed with the software package JAguc (Nebel et al., 2011a) and GenBank’s nr nucleotide database release 187 as reference database. JAguc employs BLASTn searches, with algorithm parameters adjusted for short reads (-m 7 -r 5 -q -4 -G 8 -E 6 -b 50). Using a custom Java-based script, the output files from QIIME’s OTUpipe and JAguc were merged. Non-target OTUs (metazoans and embryophytes) were excluded and the resulting file converted into a biom file, which was then used as a basis for statistical and network analyses.

Statistical and network analyses

Rarefaction profiles and Shannon index (alpha-diversity), as well as Chao-Jaccard beta-diversity, were calculated in QIIME. For this purpose, data were normalized and resampled 1000 times to account for uneven sample sizes (Logares et al., 2012). UPGMA (Unweighted Pair Group Method with Arithmetic mean)-clustering was used to construct Chao-Jaccard distance dendrograms.
A table including the number of observed OTUs (only amplicons were considered that were at least 95% similar to database entries) across each sample and their taxonomic assignment (rank: family) was generated and subjected to QIME for calculation of a network data file. Cytoscape (Cline et al., 2007) was used to visualize and analyze shared and exclusive families in samples. Nomenclature follows Adl et al. (2012). We created a network graph with an ‘edge-weighted spring embedded layout’ where every dot represented a taxonomic family, which was colored according to its phylogenetic affiliation. For more details, see http://qiime.org/tutorials/making_cytoscape_networks.html.

Additionally, Fisher’s exact tests (Fisher 1922) were run to test the non-random independency of the datasets among the lakes, i.e. the null hypothesis was that the taxon distribution was equal in all lakes. For this statistical analysis, we used the vegan package of R.

**RESULTS**

**Lake characteristics**

The turbidity in FAS 3 was 16-fold higher than in FAS 4 (Table 1). Though turbidity was not directly measured in HL 5 and HL 15, the optical appearance of both lakes was similar to the according turbid and clear FAS lakes (Sommaruga pers. obs., Fig. 1). The nitrate (NO\(_3\)-N) values were generally higher in the Alpine lakes than in the Himalayan ones (mean NO\(_3\)-N of Alpine lakes vs HL lakes: ~145 vs 31 \(\mu\)g L\(^{-1}\)).

In the clear lakes, conductivity and DOC concentrations were higher than in the turbid lakes and the concentrations of TP and the pH were lower in the clear lakes, with the highest TP concentration (mean TP: 8.8 \(\mu\)g L\(^{-1}\)) observed in FAS 3 (Table 1).

**Overview of V4 amplicon data**

After quality check, 226267 sequences in total with >300 bp length (mainly between 350 and 450 bp) were used for the taxonomic assignments (Table 2; Fig. S1, Supporting Information). Our target organisms were eukaryotic unicellular organisms and fungi checked at least at the family level. Non-target sequences (1.4%), unassigned sequences (0.9%) and singletons/doubletons were removed finally resulting in 219155 sequences, and grouping into 1804 operational taxonomic units (OTUs) called at 97% sequence similarity. Different cluster thresholds (90–100%) were applied (Fig. S2, Supporting Information) and the rarefaction curves (Fig. S3, Supporting Information) showed that we obtained saturated sampling profiles for OTUs called at 97% sequence similarity. Approximately 60% of the target sequences had a best BLAST hit with >95% sequence similarity to a deposited sequence of a described taxon (Fig. S4, Supporting Information), indicating that a relatively large proportion of the data points to an as yet unsequenced novel diversity in high mountain lakes.

**Plankton diversity and partitioning of diversity**

The main groups detected in all lakes belonged to the alveolates (55.0% of total OTUs\(_{97}\%\)), the stramenopiles (34.0% of total OTUs\(_{97}\%\)), the cryptophytes (4.0% of total OTUs\(_{97}\%\)), the chloroplastids (3.6% of total OTUs\(_{97}\%\)) and the fungi (1.7% of total OTUs\(_{97}\%\)). The contribution of the phyla Centrohelida, Choanomonada, Rhizaria, Katablepharidae and Telsonema was <1% of the total OTUs\(_{97}\%). The number of total sequences
followed the same ranking except for the fungi, which represented <1% of the total sequences (Fig. 2).

In the turbid FAS 3, OTUs were assigned to 34 different taxonomic families, and in FAS 4, we detected 28 different families (Fig. 3, Table S1, Supporting Information). Interestingly, in the HL lakes, the clear HL 15 harbored a larger number of families (n = 39) as HL 5 (n = 35). Accordingly, Shannon diversity in both HL lakes was higher than in the FAS lakes, and the clear HL 15 was more diverse than HL 5 (Fig. 4).

Differences in community composition between the glacier-fed and the clear lakes in both geographic regions were confirmed by significant Fishers exact tests (P < 0.01). In addition, communities between the HL and the Alpine lakes were significantly different (Fishers exact tests, P < 0.01). Partitioning of diversity (Chao–Jaccard) shows that the two FAS lakes were more similar to each other regarding their protistan plankton communities than to either of the two HL lakes. Furthermore, the Chao–Jaccard distance among the FAS lakes was smaller than the distance among the HL lakes. This applies to both, analyses conducted with OTUs97% obtained from all four lakes (Fig. 5) as well as to taxonomic families observed in the four samples (Fig. S5, Supporting Information).

In comparison to the FAS lakes, the network analysis shows that the HL lakes had notably more families of the Chloroplastida (10 vs 5) and the Stramenopiles (14 vs. 8), whereas alveolate (18 vs 16) and fungal families (5 in both cases) were in the same order of magnitude (Fig. 3). None of the fungal families was shared between the two geographic regions. The HL lakes additionally harbored one cercozoan, one telonemid and one choanomond family that were not detected in the FAS lakes. On the other hand, one acanthoid family (Centrohelida) was exclusively detected in the FAS lakes.

The FAS lakes shared 24 families (Fig. 3), most of them assigned to the Alveolata (12 families) and Stramenopiles (6 families). Interestingly, fungi (five families) were exclusively detected in FAS 3. In total, ten families were unique to FAS 3 and only four to the clear one.

In total, 52 different families were detected in the two HL lakes, 22 of which were shared. The only choanomond family was exclusively found in the turbid HL 5. Additionally, HL 5 harbored only three unique alveolate families, whereas in the clear HL 15 seven unique alveolate families were found. An overview about the families detected in each lake is given in Table S1 (Supporting Information).

**DISCUSSION**

The few available studies focusing on microbial eukaryote plankton diversity in high mountain lakes have exposed only the tip of the iceberg of an as yet undiscovered microbial diversity in this type of extreme freshwater ecosystem (Félix et al., 1999; Stráskrábová et al., 1999; Wille et al., 1999; Sonntag, Summerer and Sommaruga 2011; Triadó-Margarit and Casamayor 2012).

In our study, we found 3252 molecular OTUs generated after 454 data processing with 97% cluster threshold target sequences = eukaryotic unicellular organisms and fungi, checked at least at the family level. Non-target sequences = multicellular organisms such as higher plants (Embryophyta) and Metazoa, prokaryotes and unassigned sequences.

| Total reads | After QIIME quality check | Sequences | OTUs | Non-target sequences | Target sequences without singletons/doubletons |
|-------------|---------------------------|-----------|------|----------------------|-----------------------------------------------|
| 350 803     | 226 267                   | 322       | 3241 | 219 155              | 1804                                          |
| 221 011     | 9073                      | 305 803   | 322  | 2015                 | 118                                           |

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Figure 2. Distribution of the protistan and fungal communities among the lakes (expressed in number of sequences in% per lake; Faselfad lakes, FAS; Himalaya lakes, HL). Note that most of the sequences were assigned to the Alveolata (45.4% of total sequences), Stramenopiles (45.2%), Cryptophyta (6.8%) and Chloroplastida (1.5%). All other groups were <1% of the total sequences.

fragments (Stoeck et al., 2014). For further reasons, discussed in detail previously, we refer to Caron et al. (2009), Nebel et al. (2011b) and Stoeck et al. (2014). Therefore, to compare our molecular data with previous studies, we here discuss our taxonomically assigned OTUs preferably on a higher taxonomic level, namely the family rank, which is a relatively solid and reliable assignment rank for short V4 tags in microbial eukaryotes (Stoeck et al., 2010; Dunthorn et al., 2012).

The major taxon groups detected in the four lakes corroborate with those found in the gene-based study of Triadó-Margarit and Casamayor (2012) for lakes located in the Central Pyrenees, Spain. The authors found that most of the sequences belonged to the stramenopiles, alveolates and cryptophytes, but also to sequences of ophistokonts (fungi), chloroplastids, rhizaria, and a few katablepharids, euglenozoans, and Telonema. Although different molecular techniques have been applied in the latter survey, our study supports their taxonomic pattern (except for euglenozoans) because we also found that most of the sequences belong to the stramenopiles, alveolates and cryptophytes (Fig. 1).

In the HL lakes (Figs 2 and 3), we detected that >90% of the target sequences belong to the alveolates (>80% of which were Dinophyceae) and stramenopiles (>70% chrysophytes, i.e. Chrysophyceae and Synurophyceae). In the Alpine lakes, Chrysophyceae and Dinophyceae are key algal groups (Tołłoti et al., 2009) belonging to the most abundant taxa (e.g. Rott 1988). They can be indicators for environmental changes such as acidification or nutrient availability (Tołłoti et al., 2003). Therefore, a regular (molecular-based) time-series survey of these taxa may be important to assess environmental changes in high mountain lakes, which are extremely sensitive ecosystems (e.g. Catalan, Curtis and Kernan 2009; Modenutti et al., 2013; Rogora et al., 2013). Our study set the baseline and benchmark for such monitoring processes. Dinophyceae and Chrysophyceae seem well adapted to this extreme cold and nutrient-limited habitat type. This is coincident with their wide distribution even in the high arctic (e.g. Charvet et al., 2012a; Charvet, Vincent and Lovejoy 2012b) and in glacial ice (García-Descalzo et al., 2013). Survival strategies of Dinophyceae and Chrysophyceae are, for example, mixotrophy enabling them to adapt to low food and light supply by combining their heterotrophic and autotrophic nutrition modes (Stoecker 1999; Holen and Boraas 2009; Stoecker et al., 2009; Charvet, Vincent and Lovejoy 2012b; McMinn and Martin 2013). Also recently, experiments have shown that mixotrophic chrysophycean species such as Dinobryon divergens, are less sensitive to the effect of glacial flour (Sommaruga and Kandolf, 2014). Furthermore, specialized life stages such as resting stages or cysts guarantee the survival under harsh environmental conditions. For example, nutrient shortage or low temperature are well known to induce cyst formation in many protists including dinoflagellates and chrysophytes, but also in ciliates (Kristiansen 1996; Foissner 2006, 2008; Mertens et al., 2012). For instance, ~24% of the freshwater dinoflagellates produce cysts (Mertens et al., 2012) and siliceous cysts are also a substantial part of the chrysophyte life cycle (e.g. Sandgren 1991).

The Shannon diversity in the HL lakes exceeded that of the FAS lakes in the Austrian Alps (Fig. 4). For this high diversity in the HL lakes, several reasons could be taken into account, all of which, however, require in-depth investigations. One possibility for this higher diversity is input from atmospheric transport or from the catchment through precipitation in the Himalayan mountain region and/or nutrient availability. In general, atmospheric transport of microorganisms by air and dust is widespread among bacteria, fungal spores, protist cysts and pollen (e.g. Foissner 2006; Kellogg and Griffin 2006). Nepal and especially the Khumbu Valley region are strongly influenced by monsoon rainfalls. Particularly, from August to September, over 98% of the total annual precipitation falls in the Khumbu Valley (Lami et al., 2010) so that probably more taxa are ‘washed
Figure 3. Network of the protistan and fungal communities in the glacier-fed turbid FAS 3 and the clear FAS 4 lake in the Austrian Alps and the glacier-fed turbid HL 5 and clear HL 15 lake in the Himalayan mountains using Cytoscape (version 2.8.3). Each dot represents a taxonomic family, which was colored according to its phylogenetic affiliation; Faselfad lakes, FAS; Himalaya lakes, HL.

out’ from the atmosphere or the catchment area. For example, typical Chlamydomonaceae (Chloroplastida) such as Chlamydomonas nivalis (‘Red Snow’) or Chloromonas nivalis live in snow and ice water and may be easily washed into a lake during snow and ice melting processes or rainfall (Hoham and Duval 2001; Remias et al., 2010). Interestingly, Chlamydomonaceae (C. raudensis) are also found in permanently ice-covered polar lakes (Pocock et al., 2004; Bielewicz et al., 2011). In addition, Zhang et al. (2007) reported a higher diversity of cultivable bacteria in glacial ice in the Himalayan mountain region especially during the monsoon period. These authors showed evidence that this was associated with long transport of continental dust and marine air masses.

The diversity (Shannon diversity, Fig. 4) was higher in lakes with a lower nitrogen concentration (HL lakes; Table 1). Not surprisingly the nitrogen deposition and concentration is higher in the European Alps (Table 1) than in the Himalayan mountain region because of increased anthropogenic input (e.g. Rogora et al., 2008). Consequently, nitrogen deposition can affect not only production and biomass of phytoplankton, but also their taxonomic composition as reviewed in Slemmons, Saros and Simon (2013). For example, certain taxa of the diatom family Fragilariaphyceae such as Asterionella formosa and Fragilaria crotonensis occurs in high abundances by nitrogen enrichment (e.g. Saros et al., 2005). In our study, the most abundant diatoms belong to this family and were predominantly found in the FAS lakes (Suppl. Table 1) with higher nitrogen concentrations.

Therefore, the higher availability of reactive nitrogen in Alpine lakes could have been important in structuring the phytoplankton community in the FAS lakes.

In the HL lakes and in the FAS lakes, protistan community structures were significantly different between the glacier-fed turbid lakes and the clear ones (Figs 3 and 5). Less pronounced differences between the two FAS lakes compared to the differences between the two HL lakes may be attributed to the close vicinity of the FAS lakes, located in the same catchment area.

Figure 4. Shannon diversity (taxonomic rank: family) between the glacier-fed turbid FAS 3 and the clear FAS 4 lake in the Austrian Alps and the glacier-fed turbid HL 5 and clear HL 15 lake in the Himalayan mountains. Only amplicons are considered that were at least 95% similar to database entries; Faselfad lakes, FAS; Himalaya lakes, HL.

Figure 5. UPGMA clustering of Chao-Jaccard beta-diversity based on OTUs called at 97% sequence similarity (for details, see the section ‘materials and methods’). Lakes from Alps in Austria (FAS) are more similar to each other than to either of the two HL lakes regarding protistan (incl. fungi) community composition. Furthermore, similarity between the glacier-fed turbid and the clear lake in the Alps (FAS 3 and FAS 4, respectively) was notably higher than similarity between communities in the glacier-fed turbid and clear lakes in the Himalayan mountains (HL 5 and HL 15, respectively). The same pattern was observed when the analysis was conducted with taxonomic families detected in the four lakes (Fig. S5, Supporting Information).
Some years ago, the two FAS lakes were connected to the same glacier. This makes it reasonable to assume that they had a similar seed community of fungi and protists before FAS 4 lost connectivity to the glacier, giving rise to the evolution of a new plankton community. In contrast, the two HL lakes are located in different catchment areas and thus, these lakes may receive different input to maintain and support the corresponding plankton community structures.

Possible explanations for differences in plankton community structures in turbid and clear lakes are for example, the high loads of suspended particles (‘glacial flour’) from retreating glacier leading for example to a higher mortality of heterotrophic flagellates (Sommaruga and Kandolf 2014). Whereas, in clear alpine lakes, the potentially harmful UV-B can reach the lake bottom (Sommaruga and Psenner 1997; Laurion et al., 2000; Sommaruga and Augustin 2006) and protists in such transparent habitats developed different strategies to protect themselves from high levels of incident solar radiation. Avoidance of high levels of solar radiation during solar noon or the synthesis or accumulation of photoprotective compounds and/or the presence of effective DNA repair mechanisms can be found among various planktonic organisms (e.g. Tilzer 1973; Sommaruga and Psenner 1997; Zagarase, Feldman and Williamson 1997; Alonso et al., 2004; Sonntag, Summerer and Sommaruga 2007, 2011; Tartarotti et al., 2014).

Worldwide, the retreat of glaciers is a given fact (e.g. Vaughan et al., 2013) resulting in the emergence of numerous newborn proglacial lakes and loss in connectivity with glaciers. The latter process again is associated with major changes in lake physical-chemical conditions in high mountain lakes. Selective mechanisms/behaviors as mentioned above are most likely major evolutionary forces governing shifts in plankton community structures when glacier-fed turbid lakes turn into clear ones after loss of glacier connectivity. In a first step, some turbid-lake taxa are probably eliminated from the original seed community, when turbid lakes shift to clear ones. This is because these taxa have low adaptive capabilities to survive in clear UVR-flooded waters. In a second step, succession of new taxa preferring clear-lake conditions complete the community shifts. To address this issue in detail future studies are needed focusing on the factors that affect community structures and ecosystem function(ing). Nevertheless, our study gives a first insight into microbial communities of glacier-fed and clear lakes revealing a high diversity and an important reservoir of largely unseen protistan diversity.

SUPPLEMENTARY DATA

Supplementary data is available at FEMSEC online.

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REFERENCES

Adl SM, AGB Simpson, Lane CE, et al. The revised classification of eukaryotes. J Eukaryot Microbiol 2012;59:429–514.

Alexander E, Stock A, Breiner HW, et al. Microbial eukaryotes in the hypersaline anoxic L’Atalante deep-sea basin. Environ Microbiol 2009;11:360–81.

Alonso C, Rocco V, Barriga JP, et al. Surface avoidance by freshwater zooplankton: field evidence on the role of ultraviolet radiation. Limnol Oceanogr 2004;49:225–32.

Auguet JC, Barberán A, Casamayor EO. Global ecological patterns in uncultured Archaea. ISME J 2010;4:182–90.

Azam FT, Fenchel T, Field JG, et al. The ecological role of water column microbes in the sea. Mar Ecol Prog Ser 1983;10:257–63.

Barberán A, Casamayor EO. Global phylogenetic community structure and beta-diversity patterns in surface bacterio-plankton metacommunities. Aquat Microb Ecol 2010;59:1–10.

Barberán A, Fernández-Guerra A, Auguet JC, et al. Phylogenetic ecology of widespread uncultured clades of the Kingdom Eurarchaeota. Mol Ecol 2011;20:1988–96.

Bielewicz S, Bell E, Kong W, et al. Protist diversity in a permanently ice-covered Antarctic Lake during the polar night transition. ISME J 2011;5:1559–64.

Bik HM, Sung W, De Ley P, et al. Metagenomic community analysis of microbial eukaryotes illuminates biogeographic patterns in deep-sea and shallow water sediments. Mol Ecol 2012;21:1048–59.

Bragg L, Stone G, Imelfort M, et al. Fast, accurate error-correction of amplicon pyrosequences using Acacia. Nat Methods 2012;9:425–6.

Caporaso JG, Kuczynski J, Stombaugh J, et al. QIIME allows analysis of high-throughput community sequencing data. Nat Methods 2010;7:335–6.

Caron DA, Countway PD, Jones AC, et al. Marine protistan diversity. Ann Rev Mar Sci 2012;4:467–93.

Caron DA, Countway PD, Savai P, et al. Defining DNA-based operational taxonomic units for microbial-eukaryote ecology. Appl Environ Microb 2009;75:5797–808.

Catalan J, Curtis CJ, Kernan M. Remote European mountain lake ecosystems: regionalisation and ecological status. Freshwater Biol 2009;54:2419–32.

Charvet S, Vincent WF, Comeau A, et al. Pyrosequencing analysis of the protist communities in a High Arctic meromictic lake: DNA preservation and change. Front Microbiol 2012a; 3:422.
Sogin ML, Morrison HG, Huber JA, et al. Microbial diversity in the deep sea and the underexplored 'rare biosphere'. P Natl Acad Sci USA 2006;103:12115–20.

Sommaruga R. The role of solar UV radiation in the ecology of alpine lakes. Photochem Photobiol 2001;62:35–42.

Sommaruga R, Augustin G. Seasonality in UV transparency of an alpine lake is associated to changes in phytoplankton biomass. Aquat Sci 2006;68:129–41.

Sommaruga R, Casamayor EO. Bacterial ‘cosmopolitanism’ and importance of local environmental factors for community composition on remote high-altitude lakes. Freshwater Biol 2009;55:994–1005.

Sommaruga R, Kandolf G. Negative consequences of glacial turbidity for the survival of freshwater planktonic heterotrophic flagellates. Sci Rep 2014;4:1–5.

Sommaruga R, Psenner R. Ultraviolet radiation in a high mountain lake of the Austrian Alps: air and underwater measurements. Photochem Photobiol 1997;65:957–63.

Sommaruga-Wögrath S, Koinig K, Schmidt R, et al. Temperature effects on the acidity of remote alpine lakes. Nature 1997;387:64–7.

Sonntag B, Posch T, Klammer S, et al. Phagotrophic ciliates and flagellates in an oligotrophic deep alpine lake: contrasting variability with seasons and depths. Aquat Microb Ecol 2006;43:193–207.

Sonntag B, Summerer M, Sommaruga R. Sources of mycosporine-like amino acids in planktonic Chlorellabearing ciliates (Ciliophora). Freshwater Biol 2007;52:1476–85.

Sonntag B, Summerer M, Sommaruga R. Factors involved in the distribution pattern of ciliates in the water column of a transparent alpine lake. J Plankton Res 2011;33:541–6.

Steele JA, Countway PD, Xia L, et al. Marine bacterial, archaean and protistan association networks reveal ecological linkages. ISME J 2011;5:1414–25.

Stock A, Breiner HW, Pachiadaki M, et al. Microbial eukaryote life in the new hypersaline deep-sea basin Thetis. Extremophiles 2012;16:21–34.

Stoeck T, Bass D, Nebel M, et al. Multiple marker parallel tag environmental DNA sequencing reveals a highly complex eukaryotic community in marine anoxic water. Mol Ecol 2010;19:21–31.

Stoeck T, Breiner HW, Filker S, et al. A morpho-genetic diversity survey on ciliate plankton from a mountain lake pinpoints the necessity of protist barcoding in microbial ecology. Environ Microbiol 2014;16:430–40.

Stoeck T, Epstein S. Protists and the rare biosphere. Crystal Ball. Environ Microbiol Rep 2009;1:20–2.

Stoecker DK. Mixotrophy among Dinoflagellates. J Eukaryot Microbiol 1999;46:397–401.

Stoecker DK, Johnson MD, de Vargas C, et al. Acquired phototrophy in aquatic protists. Aquat Microb Ecol 2009;57:279–310.

Straškrabová V, Callieri C, Carrillo P, et al. Investigations on pelagic food webs in mountain lakes—aims and methods. J Limnl 1999;58:77–87.

Tartari GA, Panzani P, Adreani L, et al. Lake Cadastre of Khumbu Himal Region: geographical-geological-limnological data base. Mem Ist Ital Idrobiol 1998;57:151–235.

Tartarotti B, Saul N, Chakrabarti S, et al. UV-induced DNA damage in Cyclops abyssorum taticus populations from clear and turbid alpine lakes. J Plankton Res 2014;36:557–66.

Tilzer MM. Diurnal periodicity in the phytoplankton assemblage of a high mountain lake. Limnol Oceanogr 1973;18:15–30.

Tolotti M, Forström L, Morabito G, et al. Biogeographical characterization of phytoplankton assemblages in high altitude, and high latitude European lakes. Adv Limnl 2009;62:55–75.

Tolotti M, Thies HJ, Cantonati M, et al. Flagellate algae (Chrysophyceae, Dinophyceae, Cryptophyceae) in 48 high mountain lakes of the Northern and Southern slope of the Eastern Alps: biodiversity, taxa distribution and their driving variables. Hydrobiologia 2003;502:331–48.

Triadó-Margarit X, Casamayor EO. Genetic diversity of planktonic eukaryotes in high mountain lakes (Central Pyrenees, Spain). Environ Microbiol 2012;14:2445–56.

Vaughan DG, Comiso JC, Allison I. Observations: Cryosphere. In: Stocker TF, Qin D, Plattner GK, et al. (eds). Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change. Cambridge, NY, USA: Cambridge University Press, 2013, 335–44.

Weisse T, Müller H. Planktonic protozoa and the microbial food web in Lake Constance. Arch Hydrobiol Spec Iss Adv Limnl 1998;53:223–54.

Wille A, Sonntag B, Sattler B, et al. Abundance, biomass and size structure of the microbial assemblage in the high mountain lake Gossenköllesee (Tyrol, Austria) during the ice-free period. J Limnl 1999;58:117–26.

Zagarese HE, Feldman M, Williamson CE. UV-B-induced damage and photoreactivation in three species of Boecella (Copepoda, Calanoida). J Plankton Res 1997;19:357–67.

Zhang S, Hon S, Ma X, et al. Culturable bacteria in Himalayan glacial ice in response to atmospheric circulation. Biogeosciences 2007;4:1–9.

Zingel P, Agasild H, Noges T, et al. Ciliates are the dominant grazers on pico- and nanoplanクトon in a shallow, naturally highly eutrophic lake. Microb Ecol 2007;53:134–42.