Community structure, picoplankton grazing and zooplankton control of heterotrophic nanoflagellates in a eutrophic reservoir during the summer phytoplankton maximum

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ABSTRACT. An intensive 5 wk study was conducted to investigate the role of protists, especially heterotrophic nanoflagellates (HNF), in microbial food webs during the summer phytoplankton bloom in the epilimnion and metalimnion of the eutrophic Řimov reservoir (South Bohemia, Czech Republic). On average, protists consumed ~90% of bacterial production in both layers. The community composition of HNF and the relative importance of different HNF groups as picoplankton consumers were determined. Small HNF (<8 μm), as chrysomonads, bodonids and choanoflagellates, usually accounted for <30% of total HNF biomass but numerically dominated the community in both layers. They consumed most of (~70 to 85%) the bacterioplankton as well as autotrophic picoplankton (APP, exclusively cyanobacteria) production in the reservoir, with the rest consumed by ciliates. Both ciliates and HNF had higher clearance rates on APP than on bacteria and their grazing was likely responsible for a sharp decrease in APP abundance (from 3-4 x 105 to <2 x 103 ml⁻¹) and a very constant size structure of bacterioplankton in which short rods in the size class of 0.4 to 0.8 μm constituted 55 to 80% of the total bacterial biomass in both layers. The proportion of HNF to total picoplankton biomass in the epilimnion indicated that the picoplankton biomass was sufficient high to support HNF growth for most of the study. Uptake of picoplankton by less numerous, but larger, HNF (kathablepharids, Goniomonas sp., and Streptomonas sp.) was negligible, while their biomass, especially in the metalimnion, exceeded that of small HNF and the total biomass of picoplankton. This suggested that food items other than picoplankton were consumed to meet their carbon requirements. Analyzing potential bottom-up and top-down factors controlling HNF numbers and biomass, we did not find a tight relationship between HNF and the concentration of bacteria and chlorophyll. Variability of HNF abundance and biomass in the epilimnion could largely be explained by cladocerans or by pooled abundances of all potential crustacean consumers of HNF. In the metalimnion, the mean cell volume of HNF was positively linked to chlorophyll but negatively to the abundances of Cyclopidae and to the pooled abundances of Ceriodaphnia quadrangula and Diaphanosoma brachyurum.

KEY WORDS: Nanoflagellates · Community structure · Picoplanktivory · Bacterial production · Food webs

INTRODUCTION

Small protists, especially heterotrophic nanoflagellates (HNF) and ciliates, are major consumers of picoplankton and nutrient regenerators; they serve as prey for numerous micro- and metazooplankters in pelagic systems (e.g. Fenchel 1987, Sherr & Sherr 1988, Sanders & Wickham 1993). Principal top-down and bottom-up factors regulating HNF dynamics in freshwater aquatic systems have been intensively studied (e.g. Pace et al. 1990, Berninger et al. 1991, Jürgens 1994, Gasol et al. 1995). However, there have been
very few studies on the trophic role and food-web relationships of this functionally diverse group of protists, labelled ‘HNF’, which contains species with different survival strategies and feeding ecology. Studies with a taxonomic insight at least at the functional level of the major components are required.

There is now increasing evidence from freshwater systems that HNF are numerically dominated by small, primarily bacterivorous forms (Sanders et al. 1989, Bennett et al. 1990, Weisse 1991). The process of ‘bacterivory’ has been largely defined as feeding on heterotrophic bacteria, but it can be considered to include (Sherr & Sherr 1994) the consumption of both heterotrophic and autotrophic picoplankton (APP). The role of APP as a component of pelagic microbial food webs has been recently revised as well (Stockner 1988, Weisse 1993). However, it received little attention as a quantitatively and qualitatively important carbon source for protists, especially for HNF (e.g. Nagata 1988, Caron et al. 1991, Weisse 1993 and references therein). On the other hand, frequent observations of larger forms of HNF within natural communities (Mathes & Arndt 1994), constituting a major proportion of HNF biomass, exceeding even the biomass of picoplankton, indicate that along with bacterivory other feeding modes are also likely of importance for some forms of planktonic HNF (Sanders 1991).

Protists, both HNF and ciliates, are very important consumers of both picoplankton and phytoplankton (Sherr & Sherr 1994), which implies that pelagic protozoa have very different physiologies, survival strategies, and feeding modes (Fenchel 1987, Sanders 1991). However, there are only a few comparative studies on protists combining a wide range of both taxonomic and ecological approaches (Finlay et al. 1988, Sanders et al. 1989, Bennett et al. 1990, Šimek et al. 1995).

As the doubling times of protozoa and other microorganisms in meso- and eutrophic waters are usually on the order of 1 to 4 d, studies with high sampling frequency are required to elucidate the functional interactions within microbial food webs. The objective of this study was to assess the role of HNF communities at the highest achievable taxonomic level in the eutrophic Římov reservoir during the summer phytoplankton peak. Whereas detailed information on the particular role of different ciliate taxa have been given in Šimek et al. (1995), we report here on: (1) community structure and grazing rates of separate groups of HNF on bacteria and picocyanobacteria, and the role of HNF in controlling picoplankton population in the epilimnion and metalimnion of the reservoir; (2) the importance of picoplankton in the diet of various HNF taxa; and (3) factors of bottom-up and top-down control of HNF.

**MATERIALS AND METHODS**

**Study site and sampling scheme.** From 16 August to 23 September 1993, water samples were collected 3 times a week between 08:30 and 09:30 h from the eutrophic Římov reservoir (South Bohemia, Czech Republic). The study site was located at the deepest point of the reservoir about 250 m from the dam. Samples were taken with a 2 l Friedinger sampler from 2 layers: the epilimnion (a mixed sample from 0.5, 1, and 1.5 m) and the metalimnion (a mixed sample from the thermocline and from 0.5 m above and below this depth). A detailed description of the sampling procedure is given in Šimek et al. (1995). From a final volume of 42 l collected from the epilimnion and metalimnion, respectively, 2 l of the mixed sample was used for further phytoplankton and microzooplankton analyses. The remaining 40 l of water was filtered through a 100 μm plankton net to concentrate and remove zoo- plankton >100 μm.

**Bacterial abundance and biomass.** Subsamples were fixed with formaldehyde (2% final conc.), stained with DAPI (final conc. 0.2% wt/vol) and enumerated by epifluorescence microscopy (Olympus BH2). Between 400 and 600 bacteria were sized by an interactive image analysis system (Lucia, Laboratory Imaging, Prague, Czech Republic). For further details see Psenner (1993). Bacterial biomass was calculated according to the allometric relationship between cell volume and carbon content given by Norland (1993). As the variation of bacterial width was often below the minimal size differences reliably detectable by image analysis and, moreover, had little variability, we chose the variation in cell length to subdivide bacterioplankton into 7 size classes at 0.4 μm intervals (for the treatment used see Pernthaler et al. 1996a).

**Bacterial production.** Bacterial production was measured via thymidine incorporation (BP-thy) with the method modified from Riemann & Sondergaard (1986); for details see Šimek et al. (1995). To determine empirical conversion factors (ECF) between thymidine incorporation rate and bacterial cell production rate, replicate 750 ml subsamples of water filtered through 1 μm filters (Poretics) were incubated for 24 h in the dark at in situ temperature. To avoid a possible breakdown of algal cells during the filtration procedure, only a gentle vacuum was applied. The bacterial cell production rate (BP-num) was calculated from the slope of the increase of ln bacterial abundance over time (0, 12, 24 h). Only samples in which no HNF were observed during and after the incubation were taken into account. The average ECF (2.25 ± 1.38 and 2.74 ± 2.27 × 10⁻¹⁸ cells mol⁻¹, n = 16, for the epilimnion and metalimnion, respectively) were used for calculations.
Picocyanobacteria abundance. Among autotrophic picoplankton (APP), only picocyanobacteria dominated, especially *Synechococcus*-like species (mean cell volume ca 0.5 μm³) and occasionally *Microcystis incerta* of similar size. The latter species was partly floc forming; samples were therefore sonicated for 1 min (Ultrasonic Homogenizer 471, Cole Palmer Instruments Co., Chicago, USA, set at 1) before counting APP, using the autofluorescence of phycobiliproteins for cell detection. The epilimnion and metalimnion APP were sized only twice during the course of the study using a fine scale eyepiece micrometer. A conversion factor of 200 fg carbon μm⁻³ was assumed (Weisse 1993).

Protistan grazing and abundance. Two kinds of prey were used to measure protistan grazing upon picoplankton: fluorescently labelled bacterioplankton (FLB) and fluorescently labelled *Synechococcus*-like cyanobacteria (FLC) prepared according to the protocol of Sherr & Sherr (1993). For details of the concentration procedures used for reservoir bacterioplankton and a *Synechococcus*-like species (mean cell volume ± SD, 0.48 ± 0.23 μm³) see Šimek & Straskrabařová (1992) and Šimek et al. (1995), respectively.

We determined protistan grazing impacts on bacterioplankton and APP in short-time FLB or FLC direct uptake experiments according to Šimek et al. (1995). Flagellate and ciliate uptake rates were determined in the same treatment, but separately for the 2 different tracers used, FLB and FLC. Tracer amounts of FLB accounted for 5 to 15% of natural bacterial abundances. The FLC addition constituted generally 10 to 25% of total APP abundance; however, at the beginning of September, when APP abundances had declined to ~2 × 10⁴ ml⁻¹, this was up to 40%. For the rest of the experimental period APP abundances were too low for the tracer technique. For protistan enumeration and tracer ingestion determinations, 30 ml subsamples were taken 0, 3, 6, 10, 20, and 30 min after tracer addition and preserved according to Sherr & Sherr (1993, for details see Šimek et al. 1995).

We determined flagellate and ciliate grazing rates in subsamples from time series from 10 to 30 and 3 to 10 min, respectively. Samples from time zero were also inspected to avoid a potential bias due to attachment of non-ingested tracers on protistan surfaces. Within 24 h after fixation, 5 ml (flagellates) or 20 to 30 ml (ciliates) subsamples were stained with DAPI, filtered through 1 μm black Poretics filters, and inspected via epifluorescence microscopy. Only flagellate individuals without plastids (checked for chl a fluorescence) were included in the counts of HNF. To estimate total protistan grazing rate (TPGR), we multiplied average uptake rates of HNF and ciliates by their total in situ abundances. For the 3 most abundant, relatively easily distinguishable groups of HNF, i.e. small *Spumella*-like chrysomonads, bodonids and choanoflagellates, we determined tentatively their group-specific grazing rates on bacteria. A comparison of clearance rates on FLB and FLC was used to calculate relative selectivity by HNF for or against one of these 2 prey types.

Biomass and community structure of HNF. Geometrical approximations (prolate spheroids) were used to determine HNF mean cell volumes based on measurements of cell length and width (by an ocular micrometer) of at least 50 individuals per sample. For biomass estimations we assumed a conversion factor of 220 fg C μm⁻³ (Børshøj & Bratbak 1987). A rough evaluation of HNF community structure was attempted 4 times combining the following approaches: (1) DAPI-stained samples in epifluorescence microscopy, (2) live sample observation, (3) protargol staining (Skibbe 1994, for details see Šimek et al. 1995), and (4) scanning electron microscopy according to a slightly modified method of Marchant (1972). Criteria for grouping HNF into different taxonomic categories were: cell size and shape; size, position, and form of nuclei; number, position, and length of flagella; lack/expression of internal/external cell organelles such as flagella, collar, kinetoplasts, vacuoles, etc. Since protargol staining and live sample observation were always applied in parallel with fluorescence microscopy, we were able to group most of the HNF into taxonomic units at the level of an order or family and, occasionally, even genus. Moreover, some of the dominant HNF species in the reservoir have been isolated and identified, so we used these reference cultures in both protargol-stained and fluorescence microscopy to confirm our determinations. Using the above approaches, however, usually between 15 to 20% of the inspected HNF could not be identified. Generally, we based our identifications on the publication of Patterson & Larsen (1991) and references therein.

Disappearance of labelled cells. To evaluate the time-integrated grazing impact of all bacterivores <100 μm (i.e. mostly protists and small rotifers), 750 ml of 100 μm prescreened subsamples was incubated for 24 h in the dark at in situ temperature with a FLB addition constituting 3 to 6% of bacterial abundance. Subsamples (20 ml) for quantifying labelled (FLB) and natural bacterial cells were taken at times 0, 12, and 24 h. To calculate the rate of cell disappearance (FLB-disap) we used a linear model which also takes into account changes in natural bacterial abundances (Salat & Marrašé 1994).

Zooplankton. Zooplankton >100 μm were concentrated from 40 l (see above), preserved in 4% formaldehyde (final conc.) and quantified by direct microscopic counting of several subsamples (McCauley 1984). Rotifers were enumerated separately from un-
RESULTS

Characteristics of bacterial and protistan abundances and cell volumes

During the study period, water temperatures in the epilimnion and metalimnion varied between 15.3 and 22.2°C, and 13.8 and 20.2°C, respectively. The ranges of bacterial abundances (Fig. 1), mean cell volumes, and the corresponding mean carbon content per cell were similar in the epilimnion (2.47 to 4.40 x 10^6 cells ml^-1, 0.039 to 0.066 μm^3, 11 to 15 fg C cell^-1) and the metalimnion (2.03 to 4.40 x 10^6 cells ml^-1, 0.034 to 0.045 μm^3, 10 to 13 fg C cell^-1); for details see Simek et al. (1995, 1996). Cells between 0.4 and 0.8 μm length accounted for 55 to 80% of the total bacterial biomass throughout the study (Fig. 2). Cells <0.4 μm and those between 0.8 and 1.2 μm length constituted together another ~10 to 28% of total bacterial biomass, while the cells >1.2 μm length were generally of minor importance.

HNF abundances ranged similarly in both layers, between 1.4 and 4.5 x 10^3 ml^-1, except for the last week of August with their remarkably higher numbers in the epilimnion (Fig. 1). While the mean cell volume of preserved HNF in the epilimnion was between 19 and 39 μm^3, it sharply increased (up to 170 μm^3) in the screened water as follows: samples (1 l) were preserved with acid Lugol's solution (1% final conc.) and, after sedimentation, postfixed with Bouin's fixative (5% final conc.) added into the 10-fold concentrated samples. The concentrated subsamples (5 ml) were filtered onto nitrocellulose membrane filters (Sypor 1, 4 μm pore size, 10 mm diam.; Barvy Laky, Czech Republic) and stained with carbolerythrosine (Sorokin & Overbeck 1972). The whole filter area was inspected.
metalimnion during the second half of the study in parallel with a rising phytoplankton biomass (data not shown). Numbers of ciliates ranged from 6 to 55 cells ml⁻¹ and from 13 to 70 cells ml⁻¹ in the epilimnion and metalimnion, respectively. For a detailed description of phytoplankton and ciliate dynamics and species compositions see Šimek et al. (1995).

**Comparison of bacterioplankton production and consumption**

To compare bacterioplankton production and consumption by protists, we used 2 different approaches. The first one was based on a short-term incubation of samples (thymidine uptake assay, BP-thy, vs total protozoan grazing rate, TPCR, measured via direct FLB uptake) whereas the second one followed changes in bacterial cell number (BP-num) versus FLB disappearance rate (FLB-disap), integrated over 24 h incubation.

Bacterioplankton production and grazing rates on bacteria (Fig. 3) showed different time courses (independent of the approach we used), but a very similar range of rates. On average, 91% of bacterial production in the epilimnion and 92% in the metalimnion was consumed by protozoa (Table 1). HNF accounted, on average, for 82 and 79% of the total protozoan grazing in the epilimnion and metalimnion, respectively. The role of ciliates (Fig. 3) has already been reported in detail by Šimek et al. (1995).

The estimates based on 24 h incubations showed slightly higher rates of bacterial consumption by bacterivores <100 μm compared to bacterial cell production (Table 1). Both short-term and 24 h incubations yielded very similar means both for bacterioplankton production and its consumption (cf. Fig. 3, Table 1). On the other hand, there was very little coincidence in the time courses of the corresponding parameters as evidenced by weak (TPGR vs FLB-disap in the epilimnion and in the pooled data) or insignificant correlations (BP-thy vs BP-num in both layers and in the pooled data; Table 2). BP-num was significantly positively linked to bacterial numbers for all but the epilimnion data set, and to FLB-disap for all but the metalimnion data set. TPCR was significantly and positively correlated with abundances of bacteria and HNF in all 3

<table>
<thead>
<tr>
<th></th>
<th>Epilimnion (10⁶ bacteria l⁻¹ h⁻¹)</th>
<th>Metalimnion (10⁶ bacteria l⁻¹ h⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacterial production based on thymidine incorporation rate</td>
<td>64.0 ± 26.7</td>
<td>58.0 ± 23.9</td>
</tr>
<tr>
<td>Total protozoan grazing rate on bacteria</td>
<td>58.1 ± 20.7</td>
<td>53.3 ± 21.3</td>
</tr>
<tr>
<td>Bacterial production based on changes in cell numbers</td>
<td>56.4 ± 29.4</td>
<td>51.7 ± 43.8</td>
</tr>
<tr>
<td>Disappearance rate of fluorescently labelled bacteria</td>
<td>65.7 ± 38.4</td>
<td>54.3 ± 26.3</td>
</tr>
</tbody>
</table>
Table 2. Spearman rank correlations of HNF and bacterial abundances and parameters of bacterial production and consumption, August-September 1993 (n = 16 for the epi- and metalimnion, n = 32 for the pooled sample). Epi: epilimnion; Meta: metalimnion; All: pooled data from both layers; HNF: total abundance of heterotrophic nanoflagellates; Bac: bacterial abundance; BP-thy: bacterial production based on thymidine incorporation rate; BP-num: bacterial production based on changes in cell numbers; TPGR: total protozoan grazing rate on bacteria; FLB-disap: FLB disappearance rate; *p < 0.05; **p < 0.01; ***p < 0.001.

<table>
<thead>
<tr>
<th></th>
<th>HNF</th>
<th>Bac</th>
<th>BP-thy</th>
<th>BP-num</th>
<th>TPGR</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All</td>
<td>0.312</td>
<td>0.290</td>
<td>0.265</td>
<td>0.614***</td>
</tr>
<tr>
<td></td>
<td>Epi</td>
<td>0.146</td>
<td>0.109</td>
<td>0.247</td>
<td>0.721**</td>
</tr>
<tr>
<td></td>
<td>Meta</td>
<td>0.344</td>
<td>-0.087</td>
<td>0.559*</td>
<td>-0.087*</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>0.344</td>
<td>0.297</td>
<td>0.379</td>
<td>0.572**</td>
</tr>
<tr>
<td></td>
<td>Epi</td>
<td>0.161</td>
<td>0.775**</td>
<td>0.398</td>
<td>0.514**</td>
</tr>
<tr>
<td></td>
<td>Meta</td>
<td>0.364</td>
<td>0.287</td>
<td>0.392</td>
<td>0.506*</td>
</tr>
</tbody>
</table>

Data sets and with FLB-disap in all but the metalimnion data set. FLB-disap was positively related to bacterial abundance and BP-thy for all but the epilimnion data set, however, no significant relation was found with abundance of HNF (Table 2, Fig. 3).

We also did not find any significant correlation between ciliate abundance or biomass versus the parameters of bacterial production and protistan bacterivory listed in Table 2.

Community structure and grazing of heterotrophic nanoflagellates on picoplankton

HNF appeared to be a very diverse group of organisms belonging to different taxa (Table 3) with distinct efficiencies as picoplanktivores (see below). We made a rough evaluation of the HNF community structure on a group or genus level on 4 separate occasions (Fig. 4). On average, 3 groups of HNF of the smallest cell size (mostly 3 to 8 μm length), chrysomonads, bodonids, and choanoflagellates, accounted for about 70% (~49, 11, and 10%, respectively) of the total HNF abundance in both the epi- and metalimnion. Owing to a high cell-specific grazing rate on picoplankton (Table 3), these HNF groups were clearly the most important picoplanktivores in the reservoir. In terms of biomass, however, they accounted only for 10 to 25% of the total HNF (except for >30% on 16 August in the metalimnion).

Other HNF taxa of larger cell volume, i.e. Kathablepharis sp., Goniomonas sp., Cercomonas sp., and Streptomonas sp. (Table 3), showed cell-specific grazing rates on bacteria which were lower by more than 1 order of magnitude. We did not present these grazing rates due to their very high statistical uncertainty, as

Table 3. Cell-specific grazing rates on bacteria for the 3 most abundant bacterivorous flagellate groups, mean cell volumes and community structure of heterotrophic nanoflagellates on group or genus level (based on evaluation of 4 sampling dates, for details see Fig. 4). Epi: epilimnion; Meta: metalimnion.

<table>
<thead>
<tr>
<th>Flagellate group</th>
<th>Grazing rate (bact flag⁻¹ h⁻¹) Mean ± SD</th>
<th>n</th>
<th>Cell volume (μm²) Mean ± SD</th>
<th>n</th>
<th>% of total number of HNF Epi</th>
<th>Meta</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spumella-like chrysomonads</td>
<td>21 ± 11</td>
<td>102</td>
<td>16 ± 6</td>
<td>102</td>
<td>50.2</td>
<td>48.7</td>
</tr>
<tr>
<td>Bodonids</td>
<td>30 ± 17</td>
<td>50</td>
<td>60 ± 33</td>
<td>50</td>
<td>10.7</td>
<td>11.3</td>
</tr>
<tr>
<td>Choanoflagellates</td>
<td>53 ± 19</td>
<td>50</td>
<td>43 ± 15</td>
<td>50</td>
<td>11.0</td>
<td>9.5</td>
</tr>
<tr>
<td>Kathablepharis sp.</td>
<td>97 ± 33</td>
<td>48</td>
<td>97 ± 33</td>
<td>48</td>
<td>5.4</td>
<td>7.0</td>
</tr>
<tr>
<td>Cercomonas sp.</td>
<td>80 ± 27</td>
<td>50</td>
<td>80 ± 27</td>
<td>50</td>
<td>5.1</td>
<td>0</td>
</tr>
<tr>
<td>Goniomonas sp.</td>
<td>146 ± 44</td>
<td>44</td>
<td>996 ± 439</td>
<td>55</td>
<td>0.7</td>
<td>4.5</td>
</tr>
<tr>
<td>Streptomonas sp.</td>
<td>78 ± 63</td>
<td>70</td>
<td>78 ± 63</td>
<td>70</td>
<td>11.5</td>
<td>15.9</td>
</tr>
</tbody>
</table>
mostly zero uptake rates on FLB were observed among 44 to 50 individuals inspected per taxon which, moreover, were not present in all samples. Actually, Kathablepharis sp. was slightly more abundant in the metalimnion, but Goniomonas sp. in the epilimnion. Cercomonas sp., however, was found exclusively in the epilimnion whereas the largest (996 \( \mu m^3 \)) heart-shaped Streptomonas sp. was remarkably more abundant in the metalimnion. The occurrence of these HNF taxa strongly influenced total HNF biomass (Fig. 4). For instance on 30 August and 13 September, Streptomonas sp. in the metalimnion accounted for only 5 and 13% of total HNF numbers, but for 45 and 70% of total HNF biomass, respectively. Similarly, Goniomonas sp. accounted for 13% of total HNF numbers, but for ~35% of the total HNF biomass in the epilimnion on 22 September.

From microscopic observations, chrysomonads, bondiniids and choanoflagellates appeared to be the major APP consumers in the reservoir. Flagellates contributed equally to the total protistan grazing on APP in both layers, on average 83%, the remaining 17% was consumed by ciliates (Fig. 5). APP mean cell volumes were around 0.48 \( \mu m^3 \) in both layers (Simek et al. 1996), but their numbers dropped sharply from ~3 to 4 \( \times 10^5 \) ml\(^{-1}\) at the beginning to <2 \( \times 10^5 \) ml\(^{-1}\) by the end of the study (Fig. 5), likely due to protozoan grazing. In both layers, the amount of APP grazed per day was roughly equal (at the beginning) to or even severalfold higher (in the middle part of the study) than the APP standing stock. In parallel with the drop of APP numbers, there was a decrease in the cell-specific grazing rate of HNF on APP in both layers from ~6 to 1 cell HNF\(^{-1}\) h\(^{-1}\) to or even severalfold higher (in the middle part of the study) than the APP standing stock. In parallel with the drop of APP numbers, there was a decrease in the cell-specific grazing rate of HNF on APP in both layers from ~6 to 1 cell HNF\(^{-1}\) h\(^{-1}\) (Fig. 5). The cell-specific grazing rates of HNF on bacteria ranged from 9 to 32 and from 8 to 41 cells HNF\(^{-1}\) h\(^{-1}\) in the epilimnion and metalimnion, respectively (Fig. 6).

Using the data from both layers we plotted HNF clearance rates on bacteria against clearance rates on APP (Fig. 7). The clearance rates were calculated as grazing rates divided by prey abundance and all calculations of grazing rates were based on total numbers of HNF.

The position of all points above the 1:1 line indicated higher clearance rates of HNF on APP which were, in terms of cell volume, about 1 order of magnitude larger than the reservoir bacterioplankton (for details see

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**Fig. 4.** Community structure of heterotrophic nanoflagellates (HNF) on 4 sampling dates, based on cell abundance and total biovolume.
Fig. 6. Total picoplankton biomass (bacteria and APP) and total biomass and cell-specific grazing rate of heterotrophic nanoflagellates (HNF) on bacteria in the (a) epilimnion and (b) metalimnion, August-September 1993.

Fig. 7. Comparison of clearance rates (CR) of heterotrophic nanoflagellates calculated from ingestion rates of bacterioplankton and APP. All points from the epilimnion and metalimnion with simultaneous uptake measurements of both picoplankton food items pooled (16 August to 8 September, cf. Fig. 5). Dashed lines: 1:1 and 10:1 ratios; solid line: the best linear fit with the model $y = k \times x$.

The calculated correlation forced through zero ($y = k \times x$) indicated 3.5-fold higher clearance rates on APP than on bacteria. The bulk of organic carbon in picoplankton (38 to 84 and 27 to 93 pg l$^{-1}$ in the epilimnion and metalimnion, respectively) was dominated by bacteria (Fig. 6). APP in both layers constituted between 16 and 41% of the total picoplankton organic carbon during the first half of the study, but its proportion dropped to negligible levels during the second half of the study. As the numerical majority of HNF were picoplanktivorous, we compared the ratio of organic carbon in picoplankton biomass to HNF biomass (Fig. 6). In the epilimnion, this ratio (avg. 2.7 ± 1.3, range 1.6 to 5.7) was lowest (<2) in the middle part of the study period when the species with low uptake rates on picoplankton, such as Kathablepharis sp. and Goniomonas sp., contributed significantly to the total HNF biomass (Fig. 4). In the metalimnion, the picoplankton:HNF biomass ratio (avg. 1.3 ± 1.4, range 0.5 to 5.3) was >2 only at the beginning of the study, decreasing then to values of ~0.5 (13 September) in parallel with an increasing biomass of Streptomonas sp. (cf. Figs. 4 & 6). For the last 2 sampling dates the ratio was around 1.

Relations of HNF to zooplankton groups and phytoplankton biomass

Abundances and fluctuations of 6 zooplankton groups showed some notable differences between the epilimnion and metalimnion (Fig. 8). Daphnia (>95% D. galeata), nauplii, and Cyclopidae were more abundant in the epilimnion, the former 2 groups especially during the second half of the study. At the beginning of the study, Eudiaptomus gracilis (generally <20 ind. l$^{-1}$) and rotifer abundances (mostly 30 to 600 ind. l$^{-1}$, dominated by Keratella cochlearis, Brachionus sp., and Polyarthra sp.) peaked in the epilimnion and metalimnion, respectively. The sum of Ceriodaphnia quadrangula and Diaphanosoma brachyurum showed similar abundances in both layers with increased numbers during the second half of the study. Chl $a$ concentrations ranged from 7 to 112 µg l$^{-1}$ and from 10 to 60 µg l$^{-1}$ in the epilimnion and metalimnion, respectively; for more details see Šimek et al. (1995).

Potential top-down and bottom-up control of HNF abundance and biomass was analyzed using non-parametric Spearman rank correlations (Statgraphics) of HNF parameters with zooplankton and chl $a$ concentrations. As the epilimnion and metalimnion significantly differed in some parameters, we analyzed data from both layers separately and pooled data as well (Table 4). Here, we list only those correlations with plankton groups that were significant in at least 1 layer or in the
pooled data, i.e. all respective correlations with ciliates, nauplii, rotifers and *Eudiaptomus gracilis* were insignificant. Among zooplankton groups tested, cladocerans (especially the *Ceriodaphnia quadrangula* and *Diaphanosoma brachyurum* group) showed the strongest negative influence on numbers and biomass of HNF, at least in the epilimnion. This influence was even stronger when a group of the total herbivorous crustacean zooplankton was used (*Daphnia galeata, C. quadrangula, D. brachyurum, E. gracilis* and nauplii together), i.e. all those crustaceans potentially feeding on particles covering the HNF size range. Changes in HNF abundance and biomass were significantly negatively correlated with changes in the abundance of the total herbivorous crustacean zooplankton in the epilimnion. The corresponding correlations for the metalimnion were insignificant, likely due to mostly lower numbers of these zooplankters within the layer (Fig. 8).

The mean cell volume of HNF decreased with the abundance of the group of *Ceriodaphnia quadrangula* and *Diaphanosoma brachyurum*, and Cyclopidae in the metalimnion, and, moreover, with the numbers of *Daphnia galeata* in the pooled data set. These relationships indicated a top-down control of these zooplankton groups on large HNF dominated by *Streptomonas* sp. in the metalimnion. On the other hand, positive and highly significant correlations were found between chl *a* concentrations and the biomass and the mean cell volume of HNF in the metalimnion, indicating a tight linkage between the concomitant development of high numbers of *Streptomonas* sp. (Fig. 4) and a high phytoplankton biomass dominated by chlorophytes (especially *Staurastrum pingue; Simek et al. 1995*).

| Table 4 Spearman rank correlations of HNF parameters with selected zooplankton parameters and phytoplankton biomass (chl *a*), August-September 1993. For chl *a*, *n* = 16 for the epilimnion and metalimnion, *n* = 32 for the pooled sample. For relations with zooplankton groups, *n* = 11 for the epilimnion and metalimnion, *n* = 22 for the pooled sample. Epi: epilimnion; Meta: metalimnion; All: pooled data from both layers. *p* < 0.05, **p** < 0.01, ***p** < 0.001 |
| --- | --- | --- | --- | --- |
| **Number of HNF** | **Daphnia galeata** | **Cer + Dia** | **Total cladocerans** | **Total herbivorous crustaceans** | **Cyclopidae** | **Chl *a*** |
| All | 0.054 | -0.541* | -0.527* | -0.262 | 0.162 | -0.282 |
| Epi | -0.501 | -0.706* | -0.623* | -0.724* | -0.200 | -0.259 |
| Meta | 0.482 | -0.337 | -0.237 | 0.054 | 0.373 | -0.365 |
| **Biomass of HNF** | | | | | | |
| All | -0.537* | 0.131 | -0.290 | -0.525* | -0.468* | 0.148 |
| Epi | -0.509 | -0.509 | -0.518 | -0.627* | -0.164 | 0.937 |
| Meta | -0.091 | 0.333 | -0.342 | -0.255 | -0.309 | 0.746*** |
| **Mean cell volume of HNF** | | | | | | |
| All | -0.572** | -0.479* | 0.057 | -0.356 | -0.519* | 0.143 |
| Epi | 0.109 | 0.355 | 0.300 | 0.136 | 0.181 | 0.377 |
| Meta | -0.364 | -0.679* | -0.027 | -0.155 | -0.673** | 0.825*** |

Fig. 8 Changes in abundance of zooplankton species or groups in the epilimnion and metalimnion, August-September 1993: *Daphnia (>95% D. galeata), sum of Ceriodaphnia quadrangula and Diaphanosoma brachyurum, Cyclopidae, Eudiaptomus gracilis, nauplii, and rotifers*
DISCUSSION

Comparison of bacterial production and its consumption

One might have expected significant correlations between parameters of bacterial production, BP-thy and BP-num (Table 2), however, all relations between these 2 parameters were insignificant, likely due to the different treatment and the length of sample incubation. One possible source for the imbalance between parameters of bacterial production and bacterivory could be oscillations of these parameters over a diurnal cycle (Wikner et al. 1990, Christoffersen 1994), which might especially affect parameters derived from the short-term incubations, such as BP-thy and TPG. A more detailed discussion of potential biases of the latter parameters has already been reported (Šimek et al. 1995). In contrast, 24 h incubation of samples under constant conditions in the dark (BP-num and FLB-disap) may reduce the influence of a diurnal cycle. Moreover, prefiltration through 1 μm pore-size filters (BP-num) might stimulate bacteria, giving faster and more effective growth (Smits & Riemann 1988), but it might also decrease bacterial growth rate as a consequence of a lower dissolved organic carbon availability caused by the absence of algae.

As the major part of bacterial production was consumed by protists, one could expect a very tight relationship between these parameters. Some typical seasonal studies with a sampling frequency of 1 wk or longer (e.g. Sanders et al. 1989, Chrzanowski & Šimek 1993), or the comparison of different ecosystems (Sanders et al. 1992), documented this tight predator-prey linkage by highly significant correlations between protistan bacterivory and bacterioplankton growth rate. Surprisingly, we did not find such highly significant correlations (Table 2) in this study with a sampling frequency of 3 times a week. This might be explained by very frequent oscillations of both parameters (Fig. 3), or a time shift between them in the order of several hours which was not detectable with our experimental design.

Very similar mean values of TPG and FLB-disap and significant correlations (Tables 1 & 2) indicated that protozoa were the major source of bacterial mortality in the different incubations. Some rotifers, which may have been present along with protozoa in the samples for estimations of FLB disappearance rate (water screened through a 100 μm mesh), could have a small influence on the total rate of bacterivory. We did not find, however, any significant correlations between numbers and species composition of rotifers on the one hand, and bacterial parameters and HNF (as the major bacterivores) on the other.

During population peaks, cladocerans become the main consumers of bacterial production in lakes (e.g. Šimek et al. 1990, Vaqué & Pace 1992, Jürgens 1994). However, they seemed to be insignificant bacterivores during the summer season in the reservoir for several reasons: (1) we estimated that ~90% of bacterial production was consumed by protists (Fig. 3), (2) cladocerans were generally not abundant enough to exert a significant grazing impact on bacterioplankton (Fig. 8), and (3) no significant correlation was found between cladoceran abundance and parameters of bacterial numbers, biomass and production (data not shown).

Community composition of HNF and their role as picoplankton consumers

We observed almost no uptake of fluorescent tracers by pigmented flagellates, and HNF were the main bacterivores in the reservoir, as is common in many freshwater lakes (e.g. Berninger et al. 1991, Sanders et al. 1992, Chrzanowski & Šimek 1993). However, our data showed a different role of picoplankton in the diet of several taxonomic groups of HNF.

The smallest HNF, chrysomonads, bodonids, and choanoflagellates, were the most abundant (especially 2.5 to 6 μm large Spumella-like chrysomonads; Fig. 4, Table 3) as in the case of other temperate lakes (Sanders et al. 1989, Bennett et al. 1990, Springmann 1993, Mathes & Arndt 1994). They also showed the highest taxon-specific uptake rates on bacteria (Table 3), but as we did not measure the corresponding values on APP, only a mean uptake rate of APP per HNF is presented (2.9 cells HNF -1 h -1). Taxon-specific cell volumes of HNF and cell volumes of their ingested picoplankton prey were transformed to carbon units using the following conversion factors (in fg C μm -3): HNF, 220 (Börsheim & Bratbak 1987); bacteria, 287 (according to Norland 1993, corresponding to the mean cell volume of the reservoir bacteria of 0.047 μm3); picocyanobacteria, 200 (Weisse 1993). Assuming a gross growth efficiency of 40% for bacterivorous HNF (Fenchel 1982) and using our estimates of ingestion, the doubling times of chrysomonads, bodonids, and choanoflagellates would be 18, 38, and 24 h, respectively. Edelmann (1993) reported a very similar average doubling time of HNF <5 μm (37 h, >50% chrysomonads) for the period April–October 1993 in the Röm reservoir, estimated from changes in HNF abundance in the absence of their predators. Also, Jürgens (1992) found that small chrysomonads are well adapted to exploit and grow on in situ bacterial concentrations. Since monodispersed FLB and FLC were used, the grazing data for bodonids might be partly underestimated, as these forms are usually asso-
associated with surfaces and specialized in feeding on attached bacteria (Caron 1987). Choanoflagellates were mostly observed as sessile forms on cells of *Staurastrum pingue* and *Fragilariella crotonensis*, constituting the major portion of phytoplankton biomass in the reservoir (Simek et al. 1995).

Our splitting of HNF into taxonomic groups supplemented by their uptake rates on picoplankton showed that only a portion, i.e. chrysomonads, bodonids, and choanoflagellates, are true bacterivores. For larger HNF (see Fig. 4) we observed negligible uptake rates on picoplankton which would not allow them to meet their carbon requirements. Cercomonas spp. were mostly considered as bacterivorous (Patterson & Zölle 1991) which was not clearly confirmed in our study. *Goniomonas* sp., tentatively identified as *G. truncata*, was likely able to grow without particulate food (Sanders 1991), which supported both our field data (this study) and observations from laboratory stock cultures (Simek unpubl. data), in which almost no uptake of both FLB and FLC was recorded. Kathablepharids were rarely found ingesting picoplankton, but we frequently observed them with ingested autofluorescing cells of blue-greens or algae in a size range from 1.5 to 4 μm. However, FLB could be substantially different from natural bacterioplankton regarding cell surface properties, motility, etc. Thus, we could not completely exclude the possibility of selection for or against such tracers by respective taxa of HNF (Landry et al. 1991, Sherr & Sherr 1993).

There are no literature data about a planktonic mass occurrence of *Streptomonas* sp. This heart-shaped flagellate (12 to 20 μm) with a rigid cell and 2 equal flagella inserting into an anterior depression (Patterson & Zölle 1991) only rarely ingested picoplankton or algae in the size range of 1 to 4 μm, thus its prevailing feeding mode remains unclear. Its abundance (>300 ind. ml⁻¹) and biomass (4 times higher than that of the total picoplankton!) peaked in the metalimnion on 13 September, concomitantly with a heavy bloom of *Staurastrum pingue* and large blocks of *Fragilariella crotonensis* (cf. Simek et al. 1995). It indicated that the community contains specific food niches, where the flagellates profited from feeding within the surrounding or at the surfaces of the large algae. The specific conditions of the algal bloom also favored the development of populations of detritophagous ciliate species, such as *Coleps* sp., *Crythocyclops mucicola* and *Cinetochilum margaritaceum* (Simek et al. 1995). Thus, detritivory could also be of some importance for *Streptomonas* sp. An important part of flagellate — although not of HNF — biomass was composed by a mixotrophic dinoflagellate, *Ceratium hirundinella*. We did not observe phagotrophy so we included this species into the phytoplankton biovolume (Simek et al. 1995).

The contributions of different groups to total numbers and biomass of HNF (Fig. 4) in the Rimov reservoir were very similar to those reported for temperate lakes, such as Lake Constance (Springmann 1993), Mondsee (Salbrechter & Arndt 1994), Piburger See (Pernthaler et al. 1996b), and lakes in Northern Germany (Mathes & Arndt 1994). Small picoplanktivorous HNF numerically dominated throughout the study (Fig. 4). However, in terms of biomass, especially in the metalimnion, usually HNF ≥10 μm became more important (see also Mathes & Arndt 1994) with predatory forms dominating, and also very likely with an important role for osmotrophs (*Goniomonas* sp.) or detritivores (*Streptomonas* sp.).

While the biomass range in picoplankton was very similar in the epi- and metalimnion (Fig. 6), there were significant differences in the picoplankton:HNF carbon ratio. Assuming a 40% gross growth efficiency of HNF, total picoplankton in the epilimnion would be quantitatively a carbon source sufficient for HNF growth, except for a short period in the middle of the study. In the metalimnion, the picoplankton:HNF carbon ratio indicated that picoplankton was not even quantitatively sufficient to support the HNF growth, except for a short period at the beginning of the study. Moreover, considering the dominance of very small bacterioplankton cells (Fig. 2), this ratio does not reflect the availability of different sizes of picoplankton as food for HNF (cf. Simek & Chrzanowski 1992, Shimeta 1993).

Generally, our APP abundances fit well in the range of values reported by Weisse (1993) and references therein) for lakes and reservoirs of similar trophy. Numerous planktonic organisms of different taxonomic position have been identified as APP feeders (see review of Stockner & Antia 1986), but HNF and ciliates are considered as primary consumers of APP both in marine and fresh waters (e.g. Nagata 1988, Kuosa 1991, Weisse 1993, Simek et al. 1995). We paid specific attention to the role of protozoa in controlling APP abundance, the sharp decrease of which could be explained by protozoan grazing. We did not measure APP growth rate, but assuming 1 doubling daily (mean value reported for freshwaters; Weisse 1993), our data indicate that APP would have been subjected to significant grazing pressure by protozoa.

This study and our recent reports (Simek et al. 1995, 1996) are among the first presenting species- or taxon-specific grazing rates of dominant freshwater planktonic protists on APP (see also Sherr et al. 1991). Among the reservoir HNF, accounting for 83% of the total protistan grazing on APP, small *Spumella*-like chrysomonads, bodonids and choanoflagellates contributed 40 to 60, 5 to 15, and 5 to 15%, respectively, while grazing by other groups of HNF was negligible.
However, as APP disappeared by the middle of the experiment (Fig. 5) we could not collect a sufficiently large set of data as presented for taxon-specific grazing rates on bacteria (cf. Table 3). Among ciliates, small oligotrichs, dominated by \textit{Halteria grandinella} and \textit{Strombidium hexakinetum}, were the major APP consumers (Šimek et al. 1995). Also in the oligo-mesotrophic lake Piburger See, choanoflagellates and chrysomonads among HNF, and an oligotrichous ciliate, \textit{Pelagohalteria viridis}, were found to be the major picoplanktivores (Pernthaler et al. 1996b, Šimek et al. 1996).

APP have been frequently observed in food vacuoles of protozoa in lakes (e.g. Nagata 1988, Sherr et al. 1991, Christoffersen 1994, Šimek et al. 1995), but their digestibility, nutritional value, and uptake selectivity are largely unknown. In laboratory experiments, Caron et al. (1991) investigated ingestion of \textit{Paraphysomonas} sp. and 2 species of ciliates on 3 strains of \textit{Synechococcus} sp. and 2 strains of heterotrophic bacteria. Only 1 ciliate selected against \textit{Synechococcus} when offered a mixed diet of cyanobacteria and bacteria. When cyanobacteria were the sole food source, the growth efficiencies of the protozoa were substantially lower than in the presence of bacteria, likely due to an incomplete digestion of \textit{Synechococcus}. In contrast, Müller (1966) reported that \textit{Paraphysomonas} sp. selected for 2 out of 3 \textit{Synechococcus} species over bacteria in laboratory experiments. From field experiments on picoplankton consumption by protozoa in coastal waters, Caron et al. (1991) concluded that protists usually do not select for or against natural \textit{Synechococcus} cells. These inconsistent conclusions suggest that species-specific properties of predator and prey might strongly influence digestibility and the selection among \textit{Synechococcus} strains.

In food vacuoles of ciliates we observed APP in various degrees of digestion. Moreover, a tight predator-prey linkage supported significant correlations between the abundance of ciliates, \textit{Halteria grandinella} (Šimek et al. 1995, this study) and \textit{Pelagohalteria viridis} (lake Piburger See, Šimek et al. 1996) and abundance of APP. Also the proportion of small picoplanktivorous forms in the total HNF numbers and biomass was higher during the first half of the study, when APP accounted for a more significant part of the total picoplankton biomass (cf. Figs. 4 & 6). From the ingestion rates of chrysomonads, bodonids, and choanoflagellates we calculated that picocyanobacteria constituted 49, 30, and 39% of the total picoplanktonic carbon ingested, respectively. Although we had to use only the mean grazing rate for all HNF on APP, instead of taxon-specific grazing rates as used for grazing on bacteria, the above data strongly support the idea that APP are important in the diet of HNF.

The data from 2 very different systems, the eutrophic Rimov reservoir and the oligo-mesotrophic lake Piburger See, revealed much higher clearance rates of protists on APP compared to bacteria, for HNF they were on average ~3 times (Pernthaler et al. 1996b, this study), and for ciliates 2 to 4 times higher (Šimek et al. 1995, 1996). It has been reported that both HNF and ciliates appear to be size-selective consumers of various picoplankton prey in freshwaters (Šimek & Chrzanowski 1992, Jürgens & Güde 1994, Šimek et al. 1994, 1995). The protists not only cropped the major proportion of bacterial production in the reservoir (Fig. 3), they also seemed to be responsible for keeping a very stable bacterial size structure dominated by cells within a length range of 0.4 to 0.8 μm. On the basis of a model, Shimeta (1993) suggested that the size around 0.4 μm acts as the refuge at which bacterial particles have the lowest encounter rate resulting in the minimal probability of being grazed by suspension feeders. This also fits well the concept of grazing-resistant bacteria of Jürgens & Güde (1994). With respect to the significant positive correlation we found between TPGR and bacterial abundance, the very stable size structure could result from protistan grazing upon larger and fast growing bacterial cells (see also Pernthaler et al. 1996a). Thus, although the majority of bacterial cells were extremely small and other size classes had little effect on the total biomass (Fig. 2), the reservoir bacterioplankton grew with an average doubling time of 2 d. As doubling times of bacterivorous HNF (Edelmann 1993) and ciliates (Macek et al. 1996, Šimek et al. 1996) also ranged between 1 and 3 d, the data indicate a very tight predator-prey linkage within the reservoir microbial food webs.

**Bottom-up and top-down control of HNF**

Gasol et al. (1995) showed, based on an extensive data set from 16 Quebec lakes, that HNF resource factors (total phosphorus, chl. bacterial numbers) best explained the abundances of HNF among lakes differing in trophic. Within lakes, however, HNF abundance was not a direct function of resource concentration. The analysis of seasonal patterns among lakes further showed that the bottom-up variables were more important in spring and top-down variables in summer. Our study was timed from mid- to late summer, when heavy algal blooms, typically dominated by large inedible phytoplankters, occur (Komáreková 1994), resulting, together with strong grazing pressure exerted by fish, in moderate to low abundances of metazooplankton in the reservoir (Štraškrabová et al. 1993, Brandl 1994). We did not find a tight linkage of HNF abundance to the resource parameters by statisti-
tor-prey linkage as neither total abundance of rotifers nor abundances of presumably nano- or microphagous species of rotifers, i.e. Keratella cochlearis, Polyarthra sp., and Brachionus sp. (Sanders & Wickham 1993), were correlated with HNF parameters (data not shown). Basically the same holds also for ciliates, which might partly result from the prevailing sizes of reservoir HNF and ciliates. Although oligotrichs are very efficient consumers of small HNF under laboratory conditions (Jürgens et al. 1997), and HNF are generally considered as potentially important food source for ciliates (e.g. Weisse 1991), our data did not support the tight food linkage between these protists.

Conclusions

A complex food web structure developed during the period of the summer phytoplankton peak in the eutrophic Římov reservoir (Figs. 4, 5 & 6). Bottom-up (picoplankton abundance, phytoplankton composition, and chl) and top-down factors (especially cladocerans and the sum of all herbivorous crustaceans; Table 4) varied in their impact on abundance, biomass and taxonomic structure of the reservoir HNF. To determine the main carbon fluxes within microbial food webs, the dominant protists must be identified at least at a functional level. Among protists, small HNF (chrysomonads, bodonids and choanoflagellates) consumed the major part (ca 80 to 85%) of both bacterial and APP production in the reservoir, while the rest was grazed by ciliates (Figs. 3 & 5 in this study, and Šimek et al. 1995). In terms of available organic carbon and uptake selectivity, APP represents a food source of high importance, but further studies are necessary to assess their digestibility and nutritional adequacy for planktonic protozoa on a species-specific basis.

Both the sharp decrease in APP abundance and the typical size structure of bacterioplankton cells dominated by short rods were apparently caused by heavy protozoan grazing. Besides picoplanktivory, recognized as the most important feeding mode of numerically dominating small HNF, other feeding modes such as algivory, detritivory, and osmotrophy were likely of high importance for larger HNF dominating in terms of biomass (see also Sanders 1991, Mathes & Arndt 1994), especially in the metalimnion of the reservoir.

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