Fate of increased production in late-summer plankton communities due to nutrient enrichment of the Baltic Proper

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ABSTRACT: The fate of increased production due to nutrient enrichment was studied experimentally in late-summer plankton communities in the central Baltic Sea in 1997 and 1998. In the first year, inorganic nitrogen and phosphorus, as well as labile organic carbon, were added to natural and 0.8 µm pre-filtered seawater in a 2³ factorial design, and the response of the heterotrophic bacterial community was followed. In the second year, 2 experiments were carried out, where the response of the whole microbial community to nutrient and carbon enrichments was studied. In the first of those experiments, the seawater was pre-filtered through a 40 µm filter to exclude filamentous cyanobacteria. In the second experiment, a 40-fold concentrate of >90 µm plankton was added, resulting in a 9-fold concentration of phytoplankton biomass compared to natural levels. All experiments were run for 3 d in a deck incubator equipped with continuous seawater flow. Phytoplankton primary production (PP) was limited by both nitrogen and phosphorus, while bacterial production (BP) was co-limited by inorganic nutrients and labile carbon source. The increased PP resulted in a higher biomass of picocyanobacteria and phototrophic flagellates, while filamentous cyanobacteria showed no positive response. The increased BP did not result in a higher bacterial biomass (BB), indicating grazing control by protozoa. In accordance, higher flagellate and/or ciliate biomasses were observed in the enriched treatments. Picocyanobacteria therefore seemed to be less edible to protozoa than the heterotrophic bacteria. The increase in BP by carbon addition increased biomasses at the higher trophic levels more than the increase in PP, indicating a tighter link in the heterotrophic part of the microbial food web and an important pathway for organic material in regenerated conditions.

KEY WORDS: Nutrient limitation · Microbial food web · Baltic Sea · Cyanobacteria · Picoplankton

INTRODUCTION

Since eutrophication has become an increasing problem in marine coastal and estuarine areas, the question of nutrient limitation of phytoplankton production and biomass has been central. It has been studied by a variety of methods such as nutrient contents of cells, nutrient uptake rates and physiological indicators (e.g. Granéli 1984, 1987, Sakshaug & Olsen 1986, Zevenboom 1986, Hecky & Kilham 1988 and references therein, Graziano et al. 1996, Hameed et al. 1999, Holmboe et al. 1999, Mallin et al. 1999, Seppälä et al. 1999). Bioassays, in which the response of phytoplankton primary production (PP) and biomass accumulation is followed after nutrient enrichment, have been used in several studies (Granéli et al. 1990, Kivi et al. 1993, Downing et al. 1999 and references therein). The overall conclusion of the bioassays carried out in
coastal areas of the Baltic Sea is that the limiting nutrient varies not only areally but also seasonally between phosphorus and nitrogen, while the open sea areas are largely nitrogen limited (Granéli et al. 1988, Kivi et al. 1993, HELCOM 1996, Seppälä et al. 1999). Besides N and P, Fe has also been suggested as limiting the growth of filamentous cyanobacteria in the open areas of the Baltic Sea (Stal et al. 1999).

The Baltic Sea is one of the many estuaries in the world that suffers from severe eutrophication problems (HELCOM 1996). In the open areas of the Baltic Sea, filamentous cyanobacteria regularly form late summer blooms, which are reported to have intensified and increased in frequency and duration (Kahru et al. 1994, Bianchi et al. 2000). These blooms are dominated by Aphanizomenon flos-aquae Ralfs, and Nodularia spumigena Mertens, which are both able to fix atmospheric nitrogen. N. spumigena blooms are frequently toxic (Sivonen et al. 1989).

Filamentous cyanobacterial blooms occur at the time when planktonic production relies largely on nitrogen and phosphorus regenerated within the planktonic food web. There is indication that the blooms are initiated in fronts, i.e. areas where different water masses meet, and where water with a low mineral N:P ratio is introduced to the trophogenic layer from below the thermocline (Kononen et al. 1996 and references therein), thus giving the N-fixing cyanobacteria a competitive advantage in a N-limited late-summer situation.

The role of the microbial food web in Baltic cyanobacterial blooms has been studied little (Heinänen et al. 1995). It is well known that microbes, especially protozoa, play an important role in the regeneration of nutrients in oligotrophic conditions (e.g. Johannes 1965, Caron & Goldman 1990 and references therein). On the other hand, picoplankton, such as Synechococcus-type cyanobacteria and heterotrophic bacteria, are effective competitors of larger phytoplankton for nutrients, and can grow at maximal rates under oligotrophic conditions (Raven 1987, 1998). In many studies, planktonic bacteria have been found to be limited not only by inorganic nutrients, but also by a source of labile carbon (e.g. Kivi et al. 1993, Kuparinen & Heinänen 1993, Zweifel et al. 1993, Rivkin & Anderson 1997, Thingstad et al. 1998, Carlsson & Caron 2001). Most nutrient enrichment bioassays with bacteria have been carried out in cultures free of predators and phytoplankton, where competition over inorganic nutrients between phyto- and bacterioplankton cannot be estimated. Moreover, the nutrient regeneration and release of labile DOC by protists (e.g. Johannes 1965, Taylor et al. 1985) cannot be taken into consideration in a system consisting only of bacteria.

In this study, we approached the question of nutrient and labile carbon limitation in the late-summer planktonic community in the oligotrophic central Baltic by nutrient enrichment bioassays. In the experiments, N and/or P limitation of phytoplankton, as well as N, P and C limitation of bacteria, was studied. The focus was on the potential nutrient competition between phytoplankton and bacteria, and food-web effects subsequent to the increased production of phyto- and bacterioplankton.

**MATERIALS AND METHODS**

**Experimental setup.** Cruises for project BASYS (EU/MAST III) to the Gotland Basin, central Baltic Sea, were carried out in late summer 1997 and 1998 (July 29 to August 10), on RV ‘Baltica’. In 1997, heavy late-phase blooms of filamentous cyanobacteria, with dense surface flocs as a consequence of sunny, calm and warm weather conditions, were studied. Before and during the 1998 cruise, the weather was stormy, with heavy southwesterly winds, and the blooms of filamentous cyanobacteria were still under development.

In the first experiment (August 3 1997), the effects of resource and grazing on bacterial production (BP) and biomass were studied. Water for the experiment was taken at Stn A1 (57° 21'N, 19° 41'E, depth 100 m; Fig. 1) from 4 m depth. Additions of carbon (36 µM glucose-C), nitrogen (6 µM NH₄-N) and phosphorus (0.6 µM PO₄-P) were made according to a 2³ factorial
design (control treatment with no additions, and additions of N only, P only, combined N+P, C only, combined C+N, combined C+P and combined C+N+P) to 1 l unfiltered or 0.5 l 0.8 µm filtered seawater in acid-rinsed and seawater-rinsed polycarbonate bottles. All treatments were performed in duplicate. The bottles were placed in a deck incubator with a continuous flow-through of seawater to keep the temperature at the ambient level. The unfiltered treatments were incubated under a neutral density filter that removed 90% of the sunlight. To inhibit autotrophic plankton growth, the filtered treatments were incubated in darkness. Bacterial biomass (BB) and BP were followed for 72 h with a 24 h sampling interval.

In Expts 2 and 3 (July 31 and August 6 1998, respectively), effects of nutrient additions on the whole microbial food web were studied. Water for the experiments was taken from 5 m depth. For Expt 2 (<40 µm experiment) the water sample was taken from Stn J1 (57° 16′ N, 20° 04′ E, depth 240 m; Fig. 1) and prefiltred through 40 µm plankton sieve to remove mesozoo-plankton and large aggregates. For Expt 3 (>90 µm experiment) the water sample from Stn T1 (57° 36′ N, 20° 04′ E, depth 110 m; Fig. 1) was treated similarly, but an inoculum of ca. 40-fold >90 µm concentrate, obtained by net hauls from 8 m depth to the surface, was added to the experimental water. Concentrated inoculum from the net was first thoroughly mixed in the <40 µm filtrate and then distributed into acid-washed 8 l polycarbonate flasks. In both 1998 experiments, nitrogen (5.7 µM NH4-N), phosphorus (0.6 µM PO4-P) and carbon (68 µM sucrose-C) were added on Day 0 according to a 23 experimental design without replication. The phosphate addition was adjusted to reach the below-thermocline concentration. Ammonium was added in a N:P ratio of 9.5, and C:P ratios were adjusted to the Redfield ratio. The experiment was run for 3 d in a deck incubator, with a continuous flow-through of seawater ensuring that the in situ temperature was maintained. Excess light was screened with a neutral filter (Rosco) attached to a polythene sheet, allowing ca. 12% of the irradiance to pass to the experimental flasks. The experiments were started at ca. 09:00 h, and sampling was carried out at 0, 24, 48 and 72 h after the start.

**Measurements.** Inorganic nutrients were measured onboard with a Hitachi U-1100 spectrophotometer using standard procedures (Koroleff & Grasshoff 1983). Chl a was measured onboard with a Shimadzu RFPC-5001 spectrofluorometer, calibrated with pure chl a (Sigma), after filtration of 50 to 100 ml aliquots onto Whatman GF/F filters and extraction in 96% ethanol for 24 h at room temperature.

In the <40 µm experiment, PP was measured in total, <10 µm and <2 µm prefractionated subsamples using the 14C method (Steemann-Nielsen 1952, Lignell 1992). After addition of NaH14CO3 (1 µCi per 20 ml), the samples were incubated for 4 h in a water bath adjusted to in situ temperature and under artificial light (160 to 200 µmol quanta m–2 s–1), which corresponded to, on average, 12% of the surface irradiance measured during the experiments. After incubation, acidified (pH < 2) 4 ml subsamples were left uncapped for 24 h (no bubbling), after which their radioactivity was measured with the LKB Wallac Rackbeta 1215 Liquid Scintillation Counter. The daily PP values were calculated using the irradiation measured onboard (Li-Cor LI-193SA, Spherical Quantum Sensor connected to a LI-1400 Datalogger).

BP was measured using the tritiated thymidine incorporation method (TTI) (Fuhrman & Azam 1982, Smith & Azam 1992). In 1997, 2 x 1 ml samples from each experimental unit were incubated with 10 nM of tritiated thymidine (final concentration). The incubation was stopped after 60 min by adding ice-cold 50% trichloroacetic acid (TCA). Non-incorporated thymidine was washed away with ice-cold 5% TCA, and centrifugation. In 1998, 3 replicates were used, the thymidine addition was 20 nM, and the 30 min incubation was stopped by adding 10 µl of 37% formalin solution to the samples, which were then extracted with 10% TCA in ice-cold conditions, and washed 3 times with ice-cold 5% TCA using centrifugation. The controls were treated in the same way as the samples after the bacteria had been prekilled by the addition of 50% TCA (1997) or 37% formalin (1998). Incorporated thymidine was counted with a Beckman LS 18101 or an LKB Wallac Rackbeta 1215 liquid scintillation counter. A conversion factor of 1 x 1018 cells per mole of incorporated thymidine was used to calculate cell production (Heinänen & Kuparinen 1992).

For bacterial enumeration, 1 to 3 ml of formaldehyde-fixed subsample was filtered onto a 0.2 µm black polycarbonate filter (Poretics) and stained with acridine orange (Hobbie et al. 1977). Nanoflagellates and picoalgae were fixed with unbuffered glutaraldehyde (final concentration 1%), stained with protamine (Haas 1982, Kuosa 1988) and filtered onto black Poretics filters (pore size 0.2 µm).

The cells were counted using an epifluorescence microscope, with a 100x oil immersion objective under blue (bacteria, nanoflagellates, and picoplanktonic eukaryotes) or green (picocyanobacteria) excitation. A total of 100 picocyanobacterial and bacterial cells were measured by eye using an ocular graticule at the beginning and end of the experiment. Their biovolume was calculated using a formula of a globe (cocci) or a cylinder (elongated cells). In 1997, an image analyser was used (Blackburn et al. 1998) for quantification of bacteria. Heterotrophic nanoflagellates were classified into 6 size
categories, based on their cell length, using an ocular micrometer. In each size category, the biovolume was estimated using the formula of an ellipsoid with a flattened, round bottom. Autotrophic nanoflagellates were distinguished by their cell shape and red chlorophyll or orange phycocerythrin (cryptomonads) fluorescence. Their biovolume was calculated from cell dimensions using species-specific formulas (Tikkanen 1986). Ciliates and net-phytoplankton were counted with a Leitz Diavert inverted microscope from Lugol-preserved samples using the Utermöhl method (Utermöhl 1958).

Calculations. The biovolumes were converted to carbon using the following coefficients: 0.35 pg C µm–3 for bacteria (Bjørnsen 1986), 0.22 pg C µm–3 for heterotrophic nanoflagellates (Børsheim & Bratbak 1987) and picocyanobacteria (Li 1986), and 0.11 pg C µm–3 for autotrophic nanoflagellates and picoeukaryotes (Edler 1979). Ciliates and net-phytoplankton were converted to biomass using their measured volumes and 0.11% carbon content (Edler 1979) in the calculations.

To evaluate the fate and effects of the increased bacterial and pico- and nanoalgal production on the microbial food web, a carbon budget was constructed from the <40 µm experiment. For the calculations, the control, NP, and the CNP units were chosen, because major PP and BP responses to the inorganic nutrient and labile carbon additions were observed in those treatments. PP and BP values from the 14C and TTI measurements were integrated over the 3 d period using the measured irradiance curve in the PP calculations. Biomass change was calculated as the difference between the initial and final biomasses in the experiment, and loss was calculated as the difference between the production-based biomass estimate and the microscopically observed biomass.

In the 1997 experiment the response in heterotrophic bacterial abundance and production to nutrient addition or predator removal was tested using ANOVA (SYSTAT). To fulfil the assumptions of homogeneity of variance, the data on BP was ln-transformed. To identify which treatments differed from the control, a 1-way ANOVA and Tukey’s test was used.

RESULTS

Effect of nutrient enrichment on the heterotrophic bacterial community

The first experiment (1997) was started in a situation with high surface temperature and low inorganic nutrient concentrations (Table 1). The phytoplankton community was dominated by cyanobacteria and nanoflagellates, and a filamentous cyanobacterial bloom was floating in the upper 5 m of the water column. The filamentous cyanobacteria formed aggregates with associated pennate diatoms, bacteria and protists. Heterotrophic bacteria, flagellates and ciliates showed maximum abundances in the top 10 m (3–5 × 10⁶, 1–6 × 10³ and 0.2–11 × 10¹ cells m⁻¹, respectively).

In the unfiltered treatments, the number of bacteria decreased with time (Fig. 2), indicating that grazing exceeded production. Concomitantly, BP increased in all treatments. The highest response, 3-fold increase in the BP over the control, was observed in the treatment with both nutrients and carbon added (Fig. 2). These results indicate that the BB was limited by predation, while the BP was substrate limited. In the 0.8 µm-prescreened treatments, both biomass and production of bacteria increased 48 h. The positive response of the bacterial number to removal of predators larger than 0.8 µm confirms the result from the first part of the experiment. A significant positive effect of nutrient treatment (CNP) on the BP (ANOVA, p < 0.05), and a significant positive response of BB to predator exclusion (ANOVA, p < 0.05), were found after 48 h of incubation. Of the single nutrient additions, carbon gave the strongest response in BP (1-way ANOVA, p < 0.05 and Tukey’s test). Additions of CP and CNP also gave significantly increased production rates (Tukey’s test, p < 0.05). In conclusion, this experiment showed that increased BP due to nutrient enrichment was directly channelled to higher trophic levels. This led us to continue with studies on the entire microbial community.

Effect of nutrient enrichment on the whole microbial community

Initially, the <40 µm experiment represented a typical late summer situation in the Baltic Sea.
Inorganic nutrients were almost depleted from the upper water column, and total phytoplankton biomass was low (Table 1). Unicellular picocyanobacteria constituted 50% of the total phytoplankton biomass. Of the picoplankton, less than 1% were the eukaryotic *Micromonas pusilla*. Photosynthetic nanoflagellates consisted mainly of Pedinellids, Cryptophytes, Pyramimonads and Chrysophytes. Although surface blooms of filamentous cyanobacteria had not yet developed, their share of the initial phytoplankton biomass was ca. 20% (mainly *Nodularia spumigena* and *Aphanizomenon* sp.). The abundance of heterotrophs was typically high for late summer; their share was 60% of the community biomass. Organisms constituted 57% of the total particulate organic carbon (POC), while the rest was detritus.

The >90 µm experiment was started 1 wk after the <40 µm experiment, with a water sample from Stn T1 (Fig. 1). Because of the addition of the >90 µm concentrate, the total phytoplankton biomass was ca. 9-fold larger than in the <40 µm experiment, which was mainly due to the 13-fold concentration of filamentous cyanobacteria and 8-fold concentration of dinoflagellates. Small filaments of *Pseudanabaena acicularis* were observed, but otherwise the algal communities were similar at the beginning of the >90 µm and <40 µm experiments. The abundances of heterotrophs were at the same level as in the <40 µm experiment, with the contribution to the microscopically observed biomass being 18%. A total of 32% of the POC was in autotrophic organisms, leaving 50% of the POC in the detritus and mesozooplankton.

**Nutrient dynamics**

At the beginning of Expts 2 (<40 µm experiment) and 3 (>90 µm), the concentrations of inorganic ammonium and phosphate were 0.25 to 0.35 µM N and 0.02 to 0.05 µM P in the units without nutrient enrichment, and 6.1 µM N and 0.7 µM P in the units with the nutrient enrichment. In both experiments, added nutrients were most rapidly used, within 2 d, in the CNP treatments (Fig. 3). In the NP treatment, nutrient depletion was never complete (1 and 20% of NH₄-N and 24 and 15% of PO₄-P was left at the end of the <40 and >90 µm experiments, respectively). In the single
nutrient treatments, the percentage of ammonium and phosphate that was left at the end of the experiments was even higher (on average 60% for NH<sub>4</sub>-N and 46% for PO<sub>4</sub>-P). Stimulation of the nutrient depletion by the other added nutrient could be calculated for the first experimental day (0 to 1 d). The highest stimulation of both ammonium and phosphate depletion rates were observed by the addition of a bacterial carbon source combined with the other inorganic nutrient (either N or P). The C addition alone did not speed up the depletion rate of either N or P. This indicates a situation where phytoplankton was co-limited by both N and P, and bacteria were limited by C, N and P.

**Primary production**

In the <40 µm experiment, PP decreased or showed no response during the experiment in all treatments except NP and CNP (Fig. 4). The total PP increase due to combined NP addition was almost entirely due to the 2 to 10 µm algae, with the picoalgal contribution to the increase significantly smaller (ca. 1/3). However, picoalgae also benefited from the NP addition, by increasing their production 5-fold compared to the control. The increase in the PP was less apparent in the CNP treatment where the bacteria satisfied their carbon demand from the added labile carbon, and competed for inorganic nutrients. PP decreased in the <2 and <10 µm size classes, and started to level off in the ‘total’ <40 µm fraction during the third experimental day, when both ammonium and phosphate were exhausted in the experimental bottles (Fig. 4). In the >90 µm experiment, the PP rate decreased from the initial 17.5 µg C l<sup>-1</sup> h<sup>-1</sup> in all treatments except NP and CNP, in which the rate increased to 30.2 and 20.9 µg C l<sup>-1</sup> h<sup>-1</sup>, respectively (Fig. 4). The increase in PP was much less than in the <40 µm experiment, where most of the increase was due to <10 µm phytoplankton. The >20 µm size fraction (filamentous cyanobacteria) contributed, on average, 43% to the PP.

**Bacterial production**

BP was stimulated by the CNP addition in the <40 µm experiment (Fig. 5). The thymidine incorporation rate increased 5-fold (max. 5.2 µg C l<sup>-1</sup> h<sup>-1</sup>) during the 3 experimental days. The BP in the <40 µm experiment showed a pattern very similar to that in the predator-free treatment in 1997. The combined NP addition without sucrose, but also the single and combined nitrogen and carbon additions, slightly enhanced the thymidine incorporation at the beginning (Fig. 5). Apparently, shortage of carbon or nutrients started to limit the bacterial growth after the first experimental day, and the only significant response was observed with the combined addition of nitrogen, phosphorus and carbon. In the >90 µm experiment, BP responded to the nutrient and carbon (CNP) enrichment immediately during the first experimental day. The final level of BP, however, was lower than in the <40 µm experiment. The other treatments gave practically no response to the nutrient manipulations, except in the CP treatment at the end of the experiment. Integrated thymidine incorporation rates in the CNP treatment of the <40 µm experiment was 25% higher than that of the >90 µm experiment.
Biomass development

Chl-a concentration followed roughly the same pattern as PP. The highest (4-fold) increase in the <40 µm experiment was observed in the NP treatment (Fig. 6), and a ca. 2.5-fold increase was observed in the CNP treatment. In the treatments with the single N or combined NC addition, chl-a increased slightly, but otherwise remained at the initial level. The highest biomass increase in the <40 µm experiment was observed in the small nanoflagellate (2 to 10 µm) size fraction, whereas chl-a in the picoalgal fraction decreased slightly from the initial level (Fig. 6). In the >90 µm experiment, grazing by the concentrated zooplankton apparently reduced any increase in chl-a concentration, because the maximum chl-a increase in the NP treatment was only ca. 60% compared to the control. In the >90 µm experiment, in which mesozooplankton was concentrated in the experimental bottles, the chl-a increase was mainly due to the increase in the <2 µm size-fraction (Fig. 6).

Picoalgal abundances were high, from 4.5 × 10^5 to 1 × 10^6 cells ml^{-1}, with the highest abundances in both experiments found in treatments enriched with both N and P (Fig. 7), although with large irregular variations. This was partly due to small picocyanobacterial colonies that were included in the counts. Practically all picoalgae were cyanobacteria of *Synechococcus*-type. The cell volume of picoplanktonic cyanobacteria increased in all treatments from the initial 0.51 µm³ to 0.86–0.80 µm³. The largest cells were found in the CN treatment of both experiments.

Bacterial abundances varied in all experimental treatments (Fig. 7), but with no apparent trend except for a slight increase in the >90 µm experiment (Fig. 7). Bacterial cell volume increased from the initial 0.033 µm³ to 0.044–0.053 µm³, with the largest increase in the N treatment.

Abundances of photosynthetic nanoflagellates increased from the initial 2.4 and 1.2 × 10^3 cells ml^{-1} (<40 and >90 µm experiments, respectively; Fig. 7). Cryptomonads were clearly favoured by the addition of nitrogen, especially when combined with phospho-
rus in both experiments, while the pedinellids reached highest cell numbers in the CNP treatments. Other treatments did not show such significant responses.

Heterotrophic nanoflagellate abundances increased 3-fold from the initial value towards the end of the >90 µm experiment (Fig. 7), but only on average 1.4-fold in the <40 µm experiment. This indicates that grazing control of the trophic cascade from mesozooplankton (in the >90 µm experiment) via ciliates prevailed in the community. In the CNP and NP additions, where picoplankton production was higher than in the other experimental treatments, the increase of heterotrophic nanoflagellates was most pronounced, indicating that they were at the same time limited by food availability.

Ciliate numbers decreased from the initial 5.1 and 5.6 x 10^4 cells l⁻¹ in both experiments (Fig. 8), but less so in the <40 µm experiment. The difference between the <40 and >90 µm experiments implies that larger zooplankton controlled the ciliate abundance, while at the same time ciliate growth was limited by food availability, as seen by the higher abundances in the NP and CNP treatments, where the production of their prey—nanoflagellates and picoplankton—was elevated.

### Carbon budget <40 µm

#### Initial community

In the initial plankton community, the share of heterotrophs was 59% of the total biomass (Fig. 9). BB (94 µgC l⁻¹) was ca. 2 times higher than that of the picoalgae (57 µgC l⁻¹), which, in turn, made up 60% of total phytoplankton biomass. Detritus comprised an approximately equal amount of carbon as picoplankton. The ‘detritus’ pool did not include mesozooplankton because it was excluded from the experimental units by the 40 µm sieve. The ratio between the biomasses of heterotrophs and autotrophs (H:A) was 1.73.

To compare the control, NP and CNP treatments during the experiment, BP and PP values were integrated over the 3 d period.

#### Control treatment

The picoplankton biomass, especially the autotrophic picocyanobacteria increased during the experiment (Fig. 9: control). Nano- and microphytoplankton biomass remained at the initial level, while filamentous cyanobacteria doubled their bio-

![Fig. 7. Abundances of picoalgae, bacteria, photosynthetic nanoflagellates (Pnan), and heterotrophic nanoflagellates (Hnan) in the experiments. Left panel: <40 µm experiment, right panel: >90 µm experiment](image_url)

![Fig. 8. Number of ciliates at the beginning (initial) and at the end of the experiments](image_url)
mass in all treatments. The contribution of heterotrophs decreased slightly to 43% of the biomass. Integrated PP during the experiment was 96 µg C L⁻¹, to which the <2 µm size fraction contributed 38% (36 µg C L⁻¹). Integrated BP (52 µg C L⁻¹) was higher than the picoalgal production and 54% of total PP. Practically all of the <2 µm PP was incorporated into the picoalgal biomass, while half of the total PP and 75% of BP were lost.

NP treatment

The addition of nitrogen and phosphorus led to a 4.5-fold increase in PP of nano- and microphytoplankton fractions (200 µg C L⁻¹) compared to the control, and more than an 8-fold increase in the picophytoplankton PP (105 µg C L⁻¹; Fig. 9: NP), while BP doubled (91 µg C L⁻¹). The contribution of the <2 µm size fraction to the total PP was slightly lower than in the control (34%), whereas BP, compared to PP, was only 30%. The major fraction (67%) of the <2 µm PP was incorporated into the picocyanobacterial biomass, whereas 65 and 74% of nano- and microphytoplankton and BP, respectively, were lost (Fig. 9: NP). Compared to the control treatment, the biomass increase in the NP treatment was 2-fold in the picoplankton fraction (‘Synechococcus’ and bacteria), and 9-fold in the nano- and microphytoplankton fraction, while the increase of the biomass of filamentous cyanobacteria was negligible (Fig. 6: NP). The heterotrophic nanoflagellate and ciliate biomass decreased slightly, and the detritus pool to ca. 67% from the initial.

CNP treatment

In the unit with the labile carbon source in addition to the inorganic ammonium and phosphate, BP (167 µg C L⁻¹) increased nearly 4- and 2-fold compared to the control and NP unit, respectively, while the PP of both the pico- (89 µg C L⁻¹) and nano- and microphytoplankton (159 µg C L⁻¹) fractions were 20 to 25% lower than in the NP treatment (Fig. 9). The biomasses of pico- and nanoalgae increased as much as in the NP treatment. Microalgae and filamentous cyanobacteria increased their biomass 35% less than in the NP treatment. As a consequence of the labile carbon addition, BB increased 38% more than in the NP treatment. Most of the >2 µm PP and BP was lost (Fig. 9), but only 10% of the <2 µm PP. In contrast to the control and the NP treatments, heterotrophic nanoflagellate and microheterotroph (ciliate) biomasses increased slightly in the CNP treatment, indicating that the increased BP was directly channelled to the protists. The amount of detritus was higher in the CNP treatment than in the others (Fig. 6: CNP), and heterotroph biomass made up 41% of the total biomass.

Fig. 9. Carbon budget of the <40 µm experiment. Left panel: biomasses of the organisms at the beginning (initial biomass) and at the end of the experiment in the control, NP and CNP units. Detritus: measured POC — microscopically derived biomass; Apico: picoplanktonic cyanobacteria and eukaryotes; Fcyano: filamentous and colonial cyanobacteria; Pnan: photosynthetic nanoflagellates; Pmicro: photosynthetic microalgae; bacteria: heterotrophic bacteria; Hnan: non-photosynthetic nanoflagellates; Hmicro: ciliates. Right panel: integrated production and loss rates of phytoplankton and bacteria. <2 µm PP: primary production in the <2 µm size fraction; <2 µm loss: primary production in the <2 µm size fraction – biomass increase of picoautotrophs; >2 µm PP: primary production in the >2 µm size class; >2 µm loss: primary production in the <2 µm size fraction – biomass increase of cyanobacteria and bacteria; Hnan: non-photosynthetic nanoflagellates; Hmicro: ciliates. Percentages show the share fraction of loss of the production: (loss/production) × 100.
In spite of the elevated production rates, the same percentage (25%) of BP appeared in the biomass of bacteria, whereas roughly 70 to 100% of the <2 µm PP was incorporated into the picoautotrophic biomass. For the >2 µm phytoplankton, the corresponding loss of PP was calculated to be ca. 50 to 69%.

The H:A ratio decreased from the initial 1.73 to 0.92 in the control unit, 0.67 in the NP and 0.85 in the CNP treatment.

Carbon turnover rate (production:biomass, P:B) of bacteria increased from the initial 0.15 d⁻¹ in the NP and CNP treatments, while both P:B and assimilation efficiency (production:chlₐ) of phytoplankton were lower at the end of the experiment compared to the initial community (Table 2).

### DISCUSSION

**Phytoplankton nutrient limitation**

On the 1997 cruise dense cyanobacterial blooms in their late phase were floating in the mixed surface layer. The initial situation of the experiments in 1998 represented an earlier phase of the bloom: filamentous cyanobacterial biomass was fairly high, but because of the prevailing winds they were distributed in the water column and no surface accumulations were seen. Water temperature, ca. 15°C, was still too low to allow intensive growth of *Aphanizomenon flos-aquae* and *Nodularia spumigena* (Kononen 1992), which were the dominating genera in the filamentous cyanobacterial community.

At the beginning of the <40 µm experiment in 1998, phytoplankton PP and biomass were low. PP in all size fractions was strongly co-limited by both nitrogen and phosphorus (cf. Fig. 4). Whereas the single N-addition increased the PP slightly, phosphorus was also needed to have a major effect on production. The same pattern was found in the response of chlₐ. Granéli et al. (1990) found overall N-limitation in the Baltic Proper, except during late-summer filamentous cyanobacterial blooms, when they found the system P-limited. Temporal variability in the nutrient limitation has been found to occur not only in coastal areas (e.g. Paasche & Erga 1988, Kivi et al. 1993) but also in the open Baltic Sea (Seppälä et al. unpubl.).

The nutrient additions especially affected the nano- and picophytoplankton production, whereas the filamentous cyanobacteria were less affected. This was obvious also in the >90 µm experiment, in which PP was already initially at a higher level as a result of the addition of the >90 µm concentrate, and the responses were less pronounced. The increases in nano- and picoalgal abundance and biomass were apparent (cf. Fig. 7), as opposed to the less significant changes in filamentous cyanobacteria. In other experiments, carried out in this area (Seppälä et al. unpubl.) in the northern Baltic coastal zone (Kuuppo-Leinikki et al. 1994), it was found that nutrient enrichments in late summer are primarily channelled to nano- or microalgae, which, under a prolonged period of nutrient enrichment, may form monospecific blooms in mesocosms (e.g. Olli et al. 1996). It is often considered that picoplanktonic cyanobacteria do not experience nutrient limitation (Raven 1987, Wehr 1989) due to their large surface:volume ratio and they are capable of growing at the maximum growth rate in oligotrophic environments. However, in this experiment both their production and biomass showed clear responses to the combined additions of nitrogen and phosphorus, indicating that their growth was nutrient-limited. A decrease in the turnover rate and assimilation efficiency of <2 µm phytoplankton (cf. Table 2) indicated, however, that much of the increase in picoalgal production was connected to the increase in their biomass.

Lifting the water sample to the deck incubator obviously gave the filamentous cyanobacteria favourable conditions for growth. Their biomass also doubled from the initial level in the control treatment (cf. Fig. 9), and the addition of inorganic N and P did not further enhance their growth in the experiments. For this the filamentous cyanobacteria most probably used intracellular P reserves and N fixation (Kromkamp 1987, Larsson et al. 2001). Deficiency of iron has been found to limit the growth and photosynthesis of oceanic nitrogen-fixing cyanobacteria (e.g. Behrenfeld et al. 1996), and may perhaps occur also in the Baltic Sea (Stal et al. 1999).

### Limitation of BP

BP was limited by both inorganic nutrients and a labile carbon source on both cruises, as indicated by the increased thymidine incorporation rate only in the
CNP treatment. Bacterial cell volume increased the most in the N treatment. Many recent studies, carried out with bacterivore-free seawater or freshwater cultures, suggest that phosphate limits BP and biomass either alone, or in combination with labile DOC (Kuparinen & Heinänen 1993, Thingstad et al. 1998, Carlsson & Caron 2001). Rivkin & Anderson (1997) reported from their study in the oligotrophic Sargasso Sea that labile carbon alone, or with either inorganic nitrogen or phosphorus, limited bacterial growth.

While most of the bacterial limitation bioassays have been carried out with predator-free seawater cultures, we included the whole microbial food web in our experiments, and in the >90 µm experiment we even concentrated phytoplankton and mesozooplankton communities. Even if nutrients were regenerated by protists and zooplankton during our study in 1998 (Caron & Goldman 1990), and especially in the >90 µm experiment, the CNP limitation of bacteria was similar to that in 1997 with predator-free seawater cultures. This implies that regeneration processes could not provide inorganic nutrients or labile carbon rapidly enough to ensure the maximal growth of bacteria. The 5- and 3-fold turnover rates in the CNP treatment compared to the initial and control and the NP treatment, respectively (cf. Table 2) also indicate that bacteria were depleted by nitrogen, phosphorus, and carbon.

In the >90 µm experiment, bacterial thymidine incorporation increased little compared to the <40 µm experiment (>2-fold difference). This may have been a methodological artefact, since we did not measure the conversion factor for TTI, but used the same conversion factor (1 × 10^{18} cells per mole of incorporated thymidine; Heinänen & Kuparinen 1992) in both experiments. On the selective grazing on large, elongated and dividing bacteria (Gonzales et al. 1990, Kuuppo-Leinikki 1990, Šimek & Chrzanowski 1992) may have suppressed the BP in the >90 µm experiment.

**Competition between phytoplankton and bacteria**

When bacteria were provided with a labile carbon substrate, the depletion rate of ammonium and phosphate increased. PP and the increase of phytoplankton biomass (chl a) were suppressed by 20 to 25% in the CNP treatment, when compared to the NP treatment in both experiments, even though the same amount of ammonium and phosphate was added. The inorganic nutrients were exhausted earlier from the CNP rather than the NP treatment (cf. Fig. 3). This could be due to the increased depletion rates of both NH_4-N and PO_4-P in the CNP treatment compared to the NP treatment through increased BP. This deficiency led to decreased PP in all size fractions on the last day of both experiments (cf. Fig. 4).

It is not certain whether this can, as such, be interpreted as bacterial superiority over phytoplankton in the competition of inorganic nutrients (Currie & Kalff 1984). It is clear, however, that bacteria were co-limited by labile carbon and inorganic nutrients, and when provided with C, they took up the inorganic NH_4 and PO_4 from the water, leaving less for the phytoplankton.

**Food-web effects of increased production**

Trophic cascade control in the food web could be observed in the experiments. Even if the removal of resource-limitation by nutrient and labile carbon additions increased the production of bacteria (>3-fold in CNP treatment over the control), a comparable increase in their biomass could not be observed. The grazer control of bacteria was further verified by the 1997 experiment, in which BB increased in the <0.8 µm prefilted units, while a decrease was observed in the unfiltered units.

Heterotrophic nanoflagellates and ciliates are known to be the main grazers of bacteria and picocyanobacteria in aquatic food webs (e.g. Fenchel 1982, Sherr & Sherr 1987, Kuuppo-Leinikki 1990, Caron et al. 1991, Kuosa 1991, Šimek et al. 1997). Unlike bacteria, the growth of picoplanktonic cyanobacteria seemed not to be grazer-controlled in these experiments. It is possible that the <2 µm PP measurements were slightly underestimated. There is evidence that part of picocyanobacterial cells may be retained on the 2 µm filter during the prefiltration, especially when they form colonies during the late-summer phase (Pick & Agbeti 1991, Uronen & Kuuppo unpubl.). The share of the <2 µm size fraction of the whole (<40 µm) PP was 52 to 60% in the control, NP and CNP treatments, which is reasonably high compared to the values published (e.g. Agawin et al. 2000), indicating that our PP measurements are not far from realistic.

In our experiments, heterotrophic nanoflagellates also seemed to be grazer-controlled. This was obvious from the difference in their growth in experiments without mesozooplankton (<40 µm prescreened experiment), in which ciliates could feed on the flagellates, and with mesozooplankton (>90 µm experiment), in which mesozooplankton eliminated the ciliates (cf. Figs. 7 & 8). The differences were smaller, however, than between the nutrient treatments. Ciliates on the next trophic level showed grazer control as well, by decreasing more, in numbers, in the experiment with
mesozooplankton (>90 µm) than without mesozooplankton (<40 µm). On the other hand, when the production rates of their prey increased in the NP and CNP treatments, ciliate abundances also increased significantly over that in the other treatments (cf. Fig. 8). This agrees with the results from mesocosm experiments in Baltic coastal areas, where Kuuppo-Leinikki et al. (1994) found that the food-web effects of increased production are visible on the ciliate and mesozooplankton levels of the food web, whereas heterotrophic nanoflagellates were mainly controlled by grazing from above.

The increased BP in the CNP treatment also resulted in a slight biomass increase in heterotrophic nanoflagellate and ciliate biomasses. On the other hand, the increased PP of pico- and nanophytoplankton did not result in comparable biomass changes of protists. This indicates that the heterotrophic carbon pathway, bacteria-flagellate-ciliate or bacteria-ciliate, was more effective than the photosynthetic pathway in the microbial food web.

Duarte et al. (2000) calculated from their mesocosm experiment that nutrient enrichment shifts the biomass ratio of the planktonic food web towards dominance by autotrophs. The same happened in our <40 µm experiment. The initial H:A ratio (1.73) decreased in all treatments, but most (0.67) in the NP treatment. This implies a lower protozoan consumer biomass in relation to that of autotrophs, and a more direct channeling of PP to the mesozooplankton or into the dissolved pool. In this study, the percentage of total PP and BP not appearing in the biomasses of the organisms, i.e. the losses, was rather high (cf. Fig. 9). Moreover, the increase of the detritus pools accounted for 0, 25 and 37% of the losses in the control, NP and CNP treatments, respectively, thereby leaving the remaining 63 to 100% as losses into the dissolved pool or by respiration.

**CONCLUSION**

In our experiments in the late-summer cyanobacterial bloom, we could not detect nutrient limitation in the growth of filamentous cyanobacteria. Nano- and microphytoplankton and picocyanobacteria were limited by both nitrogen and phosphorus, and BP was co-limited by both inorganic nutrients and the labile carbon source. The additions of NH₄-N and PO₄-P were channelled mainly to flagellated nanoplankton. The increased BP was not observed as increased biomass, indicating grazing control by protists, whereas a large fraction of the measured <2 µm PP was accumulated into the picoautotrophic biomass. The increase in production at the bottom of the microbial food web led only to weak effects in the biomasses, because of tight grazing control in the food web at all trophic levels. The increase in BP by carbon addition increased biomasses at the higher trophic levels more than an increase in PP, indicating a tighter link on the heterotrophic part of the microbial food web.

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