Phosphate addition to phosphorus-deficient Baltic Sea plankton communities benefits nitrogen-fixing Cyanobacteria

Emil Vahtera1,2,* , Riitta Autio1,3, Hermanni Kaartokallio1,3, Maria Laamanen1,4

1Department of Biological Oceanography, Finnish Institute of Marine Research, PO Box 2, 00561 Helsinki, Finland
2Present address: Woods Hole Oceanographic Institution, Redfield 332 MS#32, Woods Hole, Massachusetts 02543, USA
3Present address: Finnish Environment Institute, PO Box 140, 00251 Helsinki, Finland
4Present address: Helsinki Commission (Baltic Marine Environment Protection Commission), Katajanokanlaituri 6 B, 00160 Helsinki, Finland

ABSTRACT: The effect of single nutrient additions (ammonia or phosphate) on the biomass and phosphorus dynamics of natural Baltic Sea phytoplankton bloom communities dominated by nitrogen-fixing Cyanobacteria was studied. The aim was to elucidate bloom-time phosphorus deficiency characteristics of the community. In addition, the intention was to study the effect of an increased phosphate supply on phosphate uptake in different size fractions and to test the hypothesis of increased phosphate supply favoring large nitrogen-fixing Cyanobacteria during general phosphate deficiency. Based on specific phosphate uptake affinity and specific alkaline phosphatase activity, the bloom communities showed varying degrees of phosphorus deficiency. The bulk of phosphate uptake was dominated by the 8 to 20 µm size fraction. However, if the community initially showed signs of phosphorus deficiency, phosphate addition increased the relative amount of phosphate taken up by the size fraction dominated by nitrogen-fixing Cyanobacteria. The 2 main bloom-forming filamentous diazotrophic species, Aphanizomenon sp. (considered non-toxic in the Baltic Sea) and Nodularia spumigena (toxic), showed differing levels of phosphorus deficiency. N. spumigena constantly had a higher percentage of the population expressing alkaline phosphatase activity, as indicated by enzyme-labeled fluorescence assays. Approximately one-fifth of added phosphate was estimated to be taken up by the diazotrophic Cyanobacteria during phosphorus deficiency.

KEY WORDS: Nitrogen-fixing Cyanobacteria · Phosphate uptake · Baltic Sea · Aphanizomenon sp. · Nodularia spumigena

INTRODUCTION

During thermally stratified conditions, intrusions of deeper water alter surface layer nutrient ratios and cause species selection in Baltic Sea phytoplankton communities (Kononen et al. 1996, Vahtera et al. 2005). Quasi-stationary salinity fronts that run along the Gulf of Finland axis in the western gulf are frequently observed. Here, frontal upwelling events and horizontal advection of water affect stratification conditions and nutrient concentrations, which again affect the phytoplankton community composition (Kononen et al. 1996, Moisander et al. 1997, Pavelson et al. 1997). The northern coast of the Gulf of Finland is an area of intense coastal upwelling (Myrberg & Andrejev 2003). Coastal upwelling events affect the surface layer in a similar way as frontal upwelling, but with a more pronounced effect on surface layer temperatures. Temperatures may decrease by 10°C within hours, and phosphate and ammonium concentrations can increase up to 4- and 10-fold, respectively (Haapala 1994).

The occurrence of diazotrophic Cyanobacteria (cyanobacteria that possess the capacity to fix dissolved atmospheric nitrogen) is generally promoted by
low nitrogen to phosphorus (N:P) ratios and high P concentrations, with some species seemingly indifferent to P supply during the growth period (Smith 1983, Kangro et al. 2007). The main bloom-forming filamentous diazotrophic species in the Baltic Sea, *Nodularia spumigena* Mertens ex Bornet & Flahault 1886, and *Aphanizomenon* sp. Morren ex Bornet et Flahault 1886, typically bloom during conditions deficient in dissolved nutrients, in areas south of approximately 62° N (see Fig. 1). During these conditions, their growth and N2-fixation are promoted, to varying degrees, by the additions of inorganic P and/or trace elements and dissolved organic matter (Stal et al. 1999, Moisander et al. 2003, Stolte et al. 2006), while the phytoplankton productivity of non-N2-fixing species generally experiences N limitation (Granéli et al. 1990, Tamminen & Andersen 2007).

The pelagic phytoplankton community of the Baltic Sea undergoes a typical annual succession for coastal temperate regions. A spring bloom of dinoflagellates and diatoms is initiated by increased solar irradiation after ice break and stratification of the upper water column (Wasmund et al. 1998). The spring bloom in the Gulf of Finland and in coastal regions of the Baltic is typically N limited (Kivi et al. 1993, Kuosa et al. 1997, Tamminen & Andersen 2007), and dissolved inorganic N (DIN) is exhausted from the upper mixed layer by the end of April. Various amounts of dissolved inorganic and organic P are left over after the spring bloom in the upper mixed layer (e.g. Larsson et al. 2001, Naush et al. 2008), constituting an excess of P beyond the requirements of the non N2-fixing phytoplankton that dominate the community during spring and early summer. During summer, the dissolved inorganic N:P ratio has been below the Redfield ratio of 16:1 at least since the late 1970s in the northern Baltic Proper (Suikkanen et al. 2007). The calculated excess P, based on winter DIN and dissolved inorganic P (DIP) concentrations and a consumption of nutrients in the Redfield ratio of 16:1 during spring, has shown an increasing trend at least since the beginning of the 1990s (Laanemets et al. 2006). Late winter phosphate concentrations and subsequent excess P in various forms (biologically available dissolved inorganic and organic P forms), is hypothesized to eventually fuel the growth of a summer community often dominated by diazotrophic *Cyanobacteria* (Larsson et al. 2001, Janssen et al. 2004). Cellular P storage and processes that regenerate inorganic P in the surface layer during thermal stratification have also been noted to be important regulative factors for bloom formation of diazotrophic *Cyanobacteria* (Larsson et al. 2001, Lignell et al. 2003).

Blooms of diazotrophic *Cyanobacteria* have been observed to initiate in frontal regions and upwelled water (Kononen et al. 1996, Vahtera et al. 2005), and the frequency and intensity of pre-bloom upwelling events have been noted to be important factors promoting *Aphanizomenon* sp. blooms (Lips & Lips 2008).

We conducted a field experiment to investigate the nutrient limitation characteristics of diazotrophic cyanobacterial bloom communities during the bloom period in the Gulf of Finland and northern Baltic Proper and the effects of altered nutrient supply stoichiometry on phosphate uptake by a size fractionated community. Series of 48 h incubations with single pulsed nutrient addition treatments of ammonia or phosphate were made. The phytoplankton community biomass (as chlorophyll [chl] a and particulate P), alkaline phosphatase activity, and size fractionated community phosphate uptake were measured. In addition, species-specific P deficiency of the diazotrophic *Cyanobacteria* was studied with an enzyme-labeled fluorescence (ELF) technique.

**MATERIALS AND METHODS**

**Description of study sites and experiments.** Our study was conducted in July 2003 along a northeast to southwest transect from the central Gulf of Finland to the northern Baltic Proper (Fig. 1), encompassing a salinity gradient in the upper 5 m layer (5.4 to 6.6). In 2003, the Gulf of Finland was characterized by a moderate cyanobacterial bloom dominated by *Aphanizomenon* sp. (Lips & Lips 2008). A homogeneous upper mixed layer approximately 5 m deep was observed at all experimental stations, and the surface water temperature ranged between 18 and 20°C.

Three geographically distinct experimental stations were visited. The first station was revisited after 7 d. The experimental stations are hereafter denoted Expt A (14 July, western Gulf of Finland), B (16 July, central Gulf of Finland), C (21 July, Baltic Proper), and D (23 July, northern Baltic Proper).

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**Fig. 1.** Baltic Sea (a) and study area (b) located in the central part of the Baltic Sea. The 4 experimental stations (A to D) are marked with dots in (b).
July, western Gulf of Finland). Sea water for the experimental treatments was acquired using a 30 l sampler. Samples were taken from a depth interval of 2 to 3 m.

The experimental treatments consisted of (1) a control (no addition), (2) ammonium (NH$_4^+$) addition, and (3) phosphate (PO$_4^{3-}$) addition (hereafter denoted as Control, +N, and +P treatments, respectively). N was added as ammonium (8.0 µmol l$^{-1}$) and P as phosphate (0.5 µmol l$^{-1}$). The final nutrient concentrations in the treatments mimicked natural concentrations occurring below the seasonal thermocline during the cruise. Immediately after the nutrient additions, two 14 l aliquots of each treatment were transferred to duplicate experimental units (acid-washed transparent polyethylene containers) that were incubated for 48 h on deck in large water baths at ambient sea water temperature. Samples for determination of particulate P, alkaline phosphatase, chl a, solubilized reactive P (SRP), dissolved inorganic nitrogen (DIN) concentrations and ELF assays were acquired on 3 occasions: 1 or 2, 24 and 48 h from the start of the experiment from the 14 l experimental units. In order to determine orthophosphate turnover times ($T_t$; h), duplicate 250 ml samples were taken on 3 occasions (2, 24 and 46 h) from the 14 l experimental units and incubated on deck for 30 to 120 min in 500 ml acid-washed glass flasks.

To determine size fractionated P incorporation, 500 ml experimental units receiving the same nutrient additions as the 14 l experimental units were incubated in the on deck flow-through pool for 48 h in an acid washed 1 l polycarbonate flasks. These 500 ml experimental units were sampled at 4 occasions at 2, 8, 25 and 46 h from the start of the experiments.

Solar irradiation intensity in the on deck flow-through pools was reduced to levels equaling irradiation intensities at approximately 2 to 5 m depth (~100 µmol q m$^{-2}$ s$^{-1}$) by covering the pools with fine black mesh.

**Analytical methods.** For phytoplankton identification and quantification, 250 ml samples were preserved with acid Lugol’s solution and processed using the Utermöhl (1958) technique. Cyanobacterial filaments were counted as 100 µm segments and other phytoplankton as single cells or colonies. Biovolumes were estimated according to HELCOM guidelines (Olenina et al. 2006) with the PhytoWin software.

For analysis of chl a concentrations, duplicate 200 ml samples were filtered on GF/F (Whatman) glass fiber filters, which were extracted at room temperature for 24 h in 10 ml of 96% ethanol. Chl a concentrations were measured using a Perkin-Elmer LS-2B fluorometer, after filtration through GF/F filters for the removal of interfering particles (HELCOM 1988). The fluorometer was calibrated with pure chl a (Sigma Chemical).

SRP and DIN concentrations were determined from single 10 ml water samples according to the guidelines for the second stage of the Baltic Monitoring Programme (Grashoff et al. 1983, HELCOM 1983). The analysis detection limits were 0.05, 0.1, 0.06, and 0.25 µmol l$^{-1}$ for SRP, NO$_3$, NO$_2$, and NH$_4$, respectively. Particulate nutrients were analyzed from single 250 ml samples filtered on pre-combusted and acid washed GF/F (Whatman) filters according to the same protocol as for SRP after boiling the filter in 50 ml of persulfate solution.

Samples for determination of the orthophosphate turnover times ($T_t$) were allowed to acclimate in the water baths used for incubation for at least 30 min before addition of carrier free $^{33}$PO$_4^{3-}$ (20 000 to 50 000 DPM) to each bottle. Using a 12-hole Micropore filtration manifold, $T_t$ was measured by filtration of 10 ml sub-samples onto 25 mm, 0.2 µm pore size polycarbonate membrane filters (Whatman) at 3 time points. The filtration intervals ranged from 10 to 120 min, depending on expected turnover times based on SRP concentrations. The samples were filtered under low suction (<20 kPa) to reduce error caused by breakage of fragile cells) until completely dry and then transferred to liquid scintillation vials with immediate addition of scintillation liquid (Instagel). The incorporated $^{33}$PO$_4^{3-}$ was assayed with liquid scintillation techniques (Wallac RackBeta), and all results were corrected for decay time of $^{33}$P. Formalin killed triplicate samples (500 µl formalin 10 ml$^{-1}$ sample), treated similarly to live samples, were used to estimate the amount of abiotic adsorption. The use of formalin may cause damage to cell walls and thus leakage of P from cells. $T_t$ was calculated according to Thingstad et al. (1993), where a consumed fraction of added radioactive label $r(t)$ is assumed to follow the theoretical expression $r(t) = (1 – e^{-t/T_t})$. The incubation time following the addition of the isotope is denoted by $t$ (h). $T_t$ can then be determined with 1 time point measurement by solving the equation for $T_t = t/–ln(1−r(t))$. $T_t$ was estimated 3 times during each of the 4 experiments, at the beginning (2 h after nutrient additions) and after 24 and 46 h, except for Expt D, when only 1 estimate was made after 2 h.

Relative incorporation of $^{33}$P in 3 different size fractions (0.2–8, 8–20, and >20 µm) was measured by filtration onto membrane filters of different pore sizes (0.2 µm, 8 µm) after incubation and subsequent subtraction of relevant fractions. The >20 µm size fraction was obtained by filtration of samples through a 20 µm mesh, after which the sample was filtered onto a 0.2 µm pore size membrane filter. To obtain the >20 µm fraction, the value of the sample that was first filtered through the 20 µm mesh was subtracted from the value of the total >0.2 µm fraction. Pre-killed sample results were not subtracted from the size fractionation samples to account for abiotic adsorption of $^{33}$PO$_4^{3-}$ onto Cyanobacteria aggregate associated sur-
concentration of 125 µmol l–1. The assays were run for
(MUF-P) was used as the substrate at a final saturating
washed test tubes. 4-methylumbelliferyl-phosphate
samples were incubated at 20°C in the dark in acid-
as described by Kononen et al. (1993). Triplicate 4 ml
sured according to Petterson (1980) with modifications
for phosphate uptake,
thingstad & Rassoulzadegan (1999) for 2 size
fractions (0.2–20 µm and >20 µm). The specific affinity
for phosphate uptake, α, is estimated by dividing the
fraction of radioactive label incorporated in a size frac-
tion by the product of the turnover time (Tt) and esti-
ated P biomass (Bt), α = I / TtB.
Total alkaline phosphatase activity (APA) was mea-
sured according to Petterson (1980) with modifications
as described by Kononen et al. (1993). Triplicate 4 ml
samples were incubated at 20°C in the dark in acid-
washed test tubes. 4-methylumbelliferyl-phosphate
(MUF-P) was used as the substrate at a final saturating
centration of 125 µmol l–1. The assays were run for
60 min with measurements every 15 min, and the
results were checked for linearity. Fluorescence read-
ings were made with a Jasco 750 spectrofluorometer at
an excitation wavelength of 365 nm and an emission
wavelength of 460 nm. Fluorescence units were cali-
brated with MUF standard solutions over the range
from 0.01 to 1 µmol l–1, and the fluorescence inten-
sity increase in the samples was used to calculate the
potential APA given as µmol MUF-P hydrolyzed h–1.
APA was normalized to particulate P to obtain biomass
specific APA (SAPA) and account for bacterial and zoo-
 plankton biomass.
Species SAPA of diazotrophic Cyanobacteria was
studied using the ELF technique (González-Gil et al.
1998). The ELF 97 endogenous phosphatase detection
kit (Invitrogen E6600) was used on samples concen-
trated on a 20 µm nylon mesh that was folded into a
funnel. The sample was transferred from the mesh with a pipette into 1.5 ml centrifuge tubes, by first
washing the mesh with the filtrate to flush all of the
sample to the bottom of the funnel. The samples were
centrifuged at 1000 × g for 5 min, after which the supernatant was removed without disturbing the pel-
et. The pellet was incubated in the dark for 30 min in
500 µl of 70% ethanol, after which the centrifugation
procedure was repeated and the supernatant removed.
A 70 µl mix of the ELF-P substrate and buffer solution
(1:20) was added on the pellet, and the sample was
incubated in the dark for another 30 min. The incuba-
tion was stopped by repeatedly washing the sample
with filtered (0.2 µm) and autoclaved seawater. Sam-
ps were stored in 200 to 300 µl of filtered seawater in
the dark at 5°C. Samples were counted within 1 mo of
the preparation using epifluorescence microscopy
(Leitz Aristoplan) using a UV excitation filter set with a
long pass emission filter (excitation/emission: 340 to
380/425 nm). The total amount of filaments and the
amount of filaments showing ELF alcohol (ELFA, the
product of ELF-P hydrolysis) fluorescence were
counted. In total, at least 100 filaments were counted.

**Statistical analyses.** Mixed model repeated mea-
sures analyses (PROC MIXED) of the Statistical Analy-
sis System (SAS V8.02, SAS Institute) were used to test
for mean differences in particulate P concentration,
SAPA, chl a concentration, size fractionated phosphate
uptake, and relative amount of filaments showing
ELFA fluorescence. We tested for the main effect of
treatment and time and for the interaction of these
terms on the measured parameters separately for each
experiment for all variables except phosphate uptake
and ELFA. Differences in phosphate uptake were
tested for the main effects of treatment for each size
fraction separately. Differences in relative amounts of
filaments showing ELFA fluorescence were tested for
the main effects of species and treatment and the inter-
action of species and treatment. An autoregressive
covariance structure was used, and the experimental
unit was used as the within-subject factor and the
treatment as the between-subject factor. Logarithm
transformations were applied where necessary to
ensure a normal distribution of model residuals. When
the criteria for parametric tests could not be met by
transformations, we used a Kruskal-Wallis non-para-
metric test to test for differences between treatments
separately for each time point.

**RESULTS**

**Phytoplankton community composition and chl a
biomass**

The phytoplankton community standing stock bio-
mass at the beginning of the experiments was domi-
nated by diazotrophic Cyanobacteria in all cases. In
Expt A, 64% of total wet weight biomass consisted of
Aphanizomenon sp. (635 mg m–3), whereas the contribu-
tion of Nodularia spumigena (4.6 mg m–3) was very
modest (<1%). The biomass of haptophytes of the
genus Chrysochromulina constituted 9% (96 mg m–3)
of the total wet weight biomass. In Expt B, Aphani-
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zomenon* sp. (866 mg m$^{-3}$) contributed 62% and *N. spumigena* (110 mg m$^{-3}$) 8% to total wet weight biomass. *Chrysochromulina* spp. biomass (165 mg m$^{-3}$) comprised 11% of the total wet weight biomass. At the beginning of Expt C, in contrast to the previous experiments, *N. spumigena* biomass (929 mg m$^{-3}$) represented 55% of total wet weight biomass and *Aphani-zomenon* sp. represented 14% (238 mg m$^{-3}$) of total wet weight biomass. *Pyramimonas* spp. (Prasinophyceae; 238 mg m$^{-3}$) and the diatom *Nitzschia paleacea* (140 mg m$^{-3}$) comprised 14 and 8% of total biomass, respectively. Expt D, which was the same sampling station as Expt A, showed a decrease in *Aph- anizomenon* sp. (217 mg m$^{-3}$) biomass from the previous sampling to 45% and a slight increase of *N. spumigena* (35 mg m$^{-3}$) biomass to 7% of total wet weight biomass. *Chrysochromulina* spp. (57 mg m$^{-3}$) comprised 11% of the total wet weight biomass.

Chl $a$ concentrations varied between the experiments, with Expts B and C having on average the highest total chl $a$ concentrations (Fig. 2). Treatment effects in Expt A were non-significant. However, the concentrations changed significantly during the experiment, and the model interaction term was significant (Table 1) indicating differing slopes, i.e. differing responses through time, for the treatments. The Control treatment showed a declining chl $a$ concentration during the 48 h incubation; this is in contrast to the increasing trend observed during the +P and +N treatments (Fig. 2). All treatments showed an initial decline in chl $a$ biomass after 24 h of incubation. Expt B showed clear and significant treatment effects and a significant interaction term (Table 1). Chl $a$ concentrations declined in the Control and +N treatments in a similar manner, whereas the +P treatment showed slightly increasing concentrations. Expt C showed a

![Fig. 2. Average chlorophyll a (chl a) concentrations (µg l$^{-1}$) in the experimental units in Expts A to D during the 48 h incubations. Error bars denote SD, n = 2 (duplicate treatments). +P and +N indicate the phosphate- and ammonium-addition treatments, respectively.](image-url)
slight increase in the +P treatments and an initial increase and subsequent slight decrease in chl \(a\) concentrations in the +N treatment. The Control treatment showed stable chl \(a\) concentrations. Treatment effects were non-significant, but the differences in the response of treatments through time were mirrored by a significant interaction term (Table 1), similar to Expt A. Expt D showed no significant treatment effects. However, significant differences of chl \(a\) concentrations between days were observed (Table 1).

### Particulate and dissolved nutrient concentrations

Particulate P concentrations showed generally clear treatment effects (Fig. 3), albeit with occasional variability. In Expt A, no significant trends were discernible (Table 1), and SRP concentrations at the beginning and the end of the experiment indicated no net uptake of phosphate, except for the +N treatment (Table 2). DIN concentrations indicated net uptake of N in the Control and +P treatments.

Expt B showed significant treatment effects for particulate P concentrations, and a significant interaction term (Table 1). Particulate P had overall increasing concentrations in the +P treatment and more stable concentrations in the Control unit with a slight increase at the end of the experiment. The +N treatment showed an initial increase with a subsequent decrease by the end of the experiment (Fig. 3). The results are consistent with the SRP concentrations that show net uptake in the +P and Control treatments but no decrease in the +N treatment (Table 2). In the +P treatment, average (±SD) particulate P concentrations increased from 0.43 ± 0.03 to 0.94 ± 0.02 \(\mu\text{mol l}^{-1}\), nearly equaling the phosphate addition of 0.50 \(\mu\text{mol l}^{-1}\). DIN concentrations indicated a net uptake of N only in the Control treatment, although large variation was observed.

Expt C showed a similar pattern as Expt B in the development of particulate P (Fig. 3). We were unable to normalize the variable residuals, thus a non-parametric Kruskal-Wallis test was used for statistical analysis. The Control and +N treatments showed rather stable particulate P concentrations, and the +P treatment displayed a marked average (± SD) increase from 0.57 ± 0.01 to 0.99 ± 0.03 \(\mu\text{mol l}^{-1}\), being close to the added concentration of 0.5 \(\mu\text{mol l}^{-1}\). By the end of the experiment, the treatment effects were significant \((\chi^2 = 7.8, \text{df} = 2, p > \chi^2 0.02)\). SRP concentrations in the +P treatment showed net uptake of phosphate, whereas the SRP concentration did not change in the +N treatment. The Control and +N treatments had SRP concentrations at and below the detection limit (0.05 \(\mu\text{mol l}^{-1}\)) of the method used (Table 2). DIN concentrations showed a net uptake of N in the +N treatment. The +P treatment also showed net uptake, despite low concentrations of DIN. The Control treatments had low and variable concentrations.

Expt D showed a significant treatment effect for particulate P (Table 1). Concentrations were fairly stable during the experiment in all treatments. In the +P treatment, particulate P concentrations were approximately 0.15 to 0.20 \(\mu\text{mol l}^{-1}\) higher than in the Control and +N treatments, respectively. SRP concentrations indicated net uptake of phosphate in the +P treatment. The +N and Control treatments had similar concentrations at the beginning and end of the experiment due to very low concentrations and high variability (Table 2). The +N treatment showed a decrease in DIN, indicating a net uptake of N. DIN concentrations decreased in the Control and +P treatments as well, despite low concentrations.
Table 2. Average (±SD) soluble reactive phosphorus (SRP) and dissolved inorganic nitrogen (DIN) concentrations (µmol l⁻¹) and the average DIN:SRP ratio at the start and end of the experiments. Detection limits (DL) were SRP: 0.05 µmol l⁻¹, NH₄: 0.25 µmol l⁻¹, NO₂: 0.06 µmol l⁻¹, NO₃: 0.1 µmol l⁻¹. NA: measurement not available. +P and +N indicate the phosphate- and ammonium-addition treatments, respectively.

<table>
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<tr>
<th></th>
<th>Expt A</th>
<th>Expt B</th>
<th>Expt C</th>
<th>Expt D</th>
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<tr>
<td><strong>SRP</strong></td>
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<tr>
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<td>0.07 ± 0</td>
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<tr>
<td><strong>DIN</strong></td>
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<tr>
<td>Control</td>
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<td>165 ± 26</td>
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<td>106 ± 52</td>
<td>110 ± 10</td>
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</table>

Fig. 3. Average particulate phosphorus concentrations (µmol l⁻¹) in the experimental units in Expts A to D during the 48 h incubations. Error bars denote SD, n = 2 (duplicate treatments). Measurements for Expt A at Time 1 h are not available. +P and +N indicate the phosphate- and ammonium-addition treatments, respectively.
**SAPA and APA**

SAPA and APA were on average lowest in Expt A and highest in Expt C (Fig. 4, data for APA not shown). SAPA ranged from 0.02 to 1.18 nmol MUF h$^{-1}$ nmol P$^{-1}$ during all experiments and was lowest in the +P treatments. SAPA and APA decreased in all +P treatments, indicating an alleviation of P deficiency and a repression of enzyme synthesis by the phosphatase-producing organisms. In contrast, SAPA and APA increased in the Control and +N treatments during the time of the experiment.

Expt A showed no significant treatment effects, variation in the data was relatively large, and the overall level of SAPA was low (Table 1, Fig. 4). The treatments showed significant effects, and the model interaction term was also significant (Table 1), depicting opposite responses between the +P treatment and the Control/+N treatments during the experiment. Expt C showed a similar pattern in SAPA development to Expt B, with a significant treatment effect. The level of SAPA was nevertheless higher, indicating a higher hydrolytic enzyme activity in relation to particulate P concentrations. This manifested as a significant interaction term in the model (Table 1). Temporal and treatment related patterns of APA were similar to the patterns of SAPA (data not shown).

**P turnover times ($T_t$)**

Phosphate addition to the experimental units increased $T_t$ markedly in all experiments (Fig. 5). In Expt A, $T_t$ remained high (>$17 \text{ h}$) throughout the experiment in the +P treatment, indicating an inability of the community to assimilate the added phosphate (0.5 µmol l$^{-1}$). The +N treatment showed a decreasing $T_t$, and it was lower compared to the
Control treatment. In Expt B, $T_T$ was initially an order of magnitude lower in the Control and +N treatments compared to the +P treatment. The Control and +N treatments showed decreasing $T_T$. The +P treatment showed a rapidly decreasing $T_T$ after 24 h to levels comparable to the other treatments. Expt C showed constantly very fast $T_T$ in the Control and +N treatments. The +P treatment showed a steady $T_T$, approximately an order of magnitude faster than for the Control and +N treatments. Expt D showed similar initial patterns in $T_T$ to the other experiments. Unfortunately, subsequent measurements are not available for Expt D.

**Size fractionated phosphate incorporation and specific affinity for phosphate uptake**

The size fractionated relative incorporation of phosphate showed a dominance of incorporation in the 8–20 µm size fraction in all experiments and co-dominance with the 0.2–8 µm fraction in Expts C and D, particularly in the Control and +N treatments (Fig. 6). Phosphate addition slightly increased the relative phosphate incorporation in the >20 µm fraction in Expt B and significantly so in Expts C and D (Table 3). During Expts B, C, and D, which were considered P deficient (see below), the apparent uptake of added phosphate was evident, as approximately 20% of the added phosphate was taken up and accumulated in the size fraction >20 µm. The >20 µm size fraction contained on average 87% of *Aphanizomenon* sp. filaments and 89% of *Nodularia spumigena* filaments of the total counted filaments. Most of the *Anabaena* spp. filaments (87%) passed the 20 µm mesh used in size fractionation, and only 13% remained in the >20 µm size fraction. *Anabaena* spp. were nevertheless scarce in all experiments. Of the other large species, *Dinophysis acuminata* and *Oblea rotunda* were almost completely retained by the 20 µm mesh. Large species other than
the filamentous *Cyanobacteria* were present only as minor fractions of the total biomass in the samples.

In Expts C and D, the 0.2–8 µm fraction in the +P treatments showed a significant decrease in uptake when uptake by the >20 µm size fraction increased (Table 3). The 8–20 µm fraction in +P treatments showed only little difference from the other treatments in Expts B and C. However, for Expts A and D, the increase in uptake by this size fraction with phosphate addition was significant (Fig. 6, Table 3).
The percentage of abiotic adsorption of total uptake ranged from 0.4 to 21.4%, with an average of 2.5%. The highest proportion of abiotic adsorption was encountered in the +P treatment of Expt A.

The specific affinity for phosphate uptake (α) was lowest during Expt A (Table 4), in all treatments and both studied size fractions. Based on a definition of P deficiency (α > 10 l µmol-P⁻¹ h⁻¹) by Tanaka et al. (2006), the 0.2–20 µm size fraction was P deficient during Expts B, C, and D, and ammonium addition drove these communities into further P deficiency, depicted by increasing specific affinity for phosphate uptake. P addition markedly lowered the P deficiency of both size fractions, as shown by the decrease in the specific affinity for phosphate uptake by an order of magnitude lower than the control and ammonium addition treatments.

**DISCUSSION**

Our aim was to elucidate the effect of altered nutrient supply stoichiometry on phosphate uptake in natural cyanobacterial bloom communities during the generally nutrient depleted bloom period. In our experiments, phosphate addition always increased the particulate P concentrations and Tf, and decreased SAPA and the specific affinity for phosphate uptake. Phosphate addition also decreased the relative amount of ELFA fluorescent filaments of *Aphanizomenon* sp. and *Nodularia spumigena* to some degree. Effects of phosphate addition on chl a concentrations were subtle. Ammonium addition treatments responded similarly to the control treatments, with some exceptions. During Expt A, Tf decreased notably with ammonium addition, indicating an increased P turnover with the ammonium addition. The Tf of the control treatments was notably different from Expt A and the rest of the experiments. Tf was initially more than an order of magnitude higher in Expt A than in the other experiments, showing a faster turnover of the P pool and potentially indicating a more pronounced P shortage in Expts B, C, and D.

Tf on its own is to be considered a relative index of P availability and not an estimate of phosphate uptake. Tanaka et al. (2006) suggested that the specific phosphate uptake affinity and/or the specific APA can be used as measures of P deficiency. The specific affinity for phosphate uptake can be seen as an analogue for the clearance rate by phagotrophic organisms: the lower the affinity, the poorer the organism is at competing for a specific resource. The concept of specific affinity is valid when the substrate that is under study is not limiting growth, i.e. it is assumed that uptake of the substrate is intimately coupled with growth. If the substrate is not limiting growth, the value of the specific affinity will be an underestimate of the ‘true’ value because the substrate concentration is beyond the range of linear uptake of the substrate (Thingstad & Rassoulzadegan 1999). This scenario was probably the case during Expt A, and therefore the values for the specific affinity are lower limits of the true values.

Compared to the control and N-addition treatments, P addition lowered the average affinity for phosphate uptake by an order of magnitude in most cases. A distinct difference between control treatments was that during Expt A, the specific affinity for phosphate uptake was an order of magnitude lower than for Expts B to D. These
results show that the osmotroph community in Expt A was more P sufficient than the communities in the subsequent experiments. According to Tanaka et al. (2006), values of specific affinity for phosphate uptake >10 l µmol-P–1 h–1 indicate P deficiency of the community. However, the estimates of the specific phosphate affinity are subject to difficulties in measuring the P biomass of the osmotroph community. In the present study, we used the particulate P concentrations, subjecting the results to error due to the presence of detritus, and thus, a potential underestimation of true values.

The SAPA results further support the observation of a more P sufficient community during Expt A. SAPA was very low in Expt A and initially higher in all other experiments. Phosphate addition decreased SAPA in Expts B to D, in accordance with the specific affinity estimates. APA was determined using 4 ml samples,
which might cause high variation in the data due to
presence of large cyanobacterial aggregates unevenly
sampled by such a small volume. However, the data
did not show markedly high standard deviation, except
during Expt A, when APA was generally low.

N addition stimulated the turnover of the P pool dur-
ing Expt A. SRP concentrations declined to levels
below the detection limit by the end of Expt A, even
though no marked increase in either chl a nor P bio-
mass was observed due to initially very low SRP con-
centrations. The specific affinity for phosphate uptake
showed no increase in the 0.2–20 µm size fraction,
whereas ammonium addition seemed to increase the
affinity of the larger size fraction. Compared to the
other experiments, the affinities were nevertheless
very low. The relatively large P biomass concentration
estimates during Expt A, however, may have caused
underestimation of the specific affinities.

Dependent upon the initial conditions of the commu-
nity (i.e. P sufficiency or deficiency) phosphate addi-
tions increased the relative uptake of phosphate the
greatest in the >20 µm size fraction.

During Expt A, a proportionally larger fraction of
added phosphate (up to 20%) accumulated in particu-
late form through abiotic adsorption, compared to the
average value of 2.5% during the experiments. Abiotic
adsorption of phosphate onto Cyanobacteria-aggre-
gate surfaces might be a significant P storage method
in addition to intracellular stores. Abiotic adsorption of
phosphate has been noted to occur in filamentous
Cyanobacteria and is known to contribute to growth
(Sañudo-Wilhelmy et al. 2004, Fu et al. 2005).

The relative amount of filaments showing ELFA fluo-
rescence was always lower for Aphanizomenon sp.
Peccarily, the majority of Nodularia spumigena fila-
ments nevertheless showed ELFA fluorescence in Expt A
with very low SAPA and APA. This might be an artifact
caused by a relatively small N. spumigena population
present during Expt A, or evidence for an adaptation
to chronically low phosphate concentrations and slow
response to changes in environmental conditions
(Degerholm et al. 2006). High relative amounts of ELFA
fluorescent filaments might also be due to the utilization
of the original ELF protocol developed by González-Gil
et al. (1998). The original protocol includes an ethanol
incubation, which might increase the permeability of
the cell membrane. Therefore, the method might also
detect intracellular phosphatases (Nedoma et al. 2003).
Nedoma et al. (2003) also pointed out the importance
of variable time lag in ELFA precipitate formation. Due
to this time lag, labeling kinetic studies should be done
to determine the suitable incubation time for each envi-
ronmental sample. The use of a 30 min incubation time
in the present study might therefore have underesti-
mated the amount of ELFA fluorescence in our samples.

Furthermore, samples were stored in filtered and
autoclaved sea water of which the absolute phosphate
concentration was not known. Storage of ELF samples
in sea water with low concentrations of phosphate
might not stop the formation of ELFA. This would
cause an overestimation of ELFA fluorescent filaments
(Nedoma et al. 2003). However, since all samples were
prepared in the same manner, the errors induced by the
above methodological caveats are similar in all sam-
ple, making the samples comparable to each other.

The results from this short-term experimental study
are in general accordance with previous field research.
Pulsed phosphate supplies introduced to the surface
layer by frontal or coastal upwelling events have been
noted to promote blooms of Aphanizomenon sp.
(Kononen et al. 1996, Vahtera et al. 2005), and experi-
mental studies have shown a stimulating effect of
phosphate addition on N₂-fixation (Moisander et al.
2003, 2007). Recently, Nausch et al. (2009) showed that
phosphate enrichment of the surface layer by
upwelling markedly lowered the C:P ratio of filamentous
Cyanobacteria and that uptake of phosphate was
intensified most in P-deplete cells occurring at frontal
regions. We used non-diluted surface water samples
that do not reflect the conditions in upwelling regions;
however, they might be more representative of frontal
regions where ‘old’ surface waters with P-starved cells
are mixed with the newly upwelled phosphate-rich
intermediate waters, reflecting a situation similar to
what Nausch et al. (2009) observed.

We experimentally showed that a portion of the
added phosphate is directly assimilated into the
cyanobacterial size fraction, alleviating P deficiency. In
our experiments, no apparent effects on growth (chl a
concentrations) could be seen, presumably because
the diazotrophic filamentous Cyanobacteria generally
have slow growth rates (Lehtimäki et al. 1997, Stolte et
al. 2002); phosphate uptake has been noted to be
decoupled from growth in phytoplankton (Lean & Pick
1981). Since ammonium addition did not markedly
increase chl a concentrations in any of the experiments
in relation to the control treatments, the biomass of
non-N-fixing constituents of the plankton community
were deemed P or N and P limited. Without a com-
bined N and P addition, it is impossible to say whether
P or N and P together were the limiting factors. How-
ever, during Expt A, ammonium addition markedly
decreased phosphate turnover times. This indicates
that in our experiments, N limitation was subordinated
to P limitation. Only when P was sufficient, in this case
accumulated in the organisms as observed in Expt A,
did ammonium addition evoke an effect in phosphate
turnover times, indicating P as the dominant limiting
factor. The differing responses between experiments
are likely due to differing nutritional histories of the
CONCLUSIONS

It has been stated that the nutrient storage capacity of the bloom-forming and N-fixing Cyanobacteria would be the main strategy for their P acquisition and that winter phosphate concentrations and cellular P storage, to a large extent, govern the magnitude of the blooms (Larsson et al. 2001, Janssen et al. 2004). Based on our results, we propose that elevated phosphate concentrations cause a shift of phosphate uptake towards larger plankton size fractions during cyanobacterial blooms, when non-N-fixing species are mainly N or N and P co-deficient. In our experiments, most of the phytoplankton biomass in the >20 µm size fraction, and >50% of the entire phytoplankton community wet weight biomass, consisted of N-fixing Cyanobacteria. Elevated phosphate concentrations, caused by a single addition pulse during generally P-deficient conditions, alleviated the P deficiency of the phytoplankton community by lowering the specific affinity for phosphate uptake, the P biomass SAPA, and the species-specific expression of APA. This was especially evident for Aphanizomenon sp., and these results are in accordance with previous findings (Degerholm et al. 2006). Ammonium addition did not cause an increase in growth, measured as chl a biomass, but P addition caused an increase in phosphate uptake, with a relatively greater effect in the >20 µm size fraction dominated by the N-fixing Cyanobacteria.

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