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The significance of nitrogen fixation to new production during early summer in the Baltic Sea

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BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Introduction

References

Figures

▶I

Close

Title Page Abstract Conclusions **Tables** 14 Back Full Screen / Esc Printer-friendly Version

EGU

Interactive Discussion

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Abstract

Rates of dinitrogen (N₂) fixation and primary production were measured during two 9 day transect cruises in the Baltic proper in June-July of 1998 and 1999. Assuming that the early phase of the bloom of cyanobacteria lasted a month, total rates of N₂ fixation contributed 15 mmol N m⁻² (1998) and 33 mmol N m⁻² (1999) to new production (sensu Dugdale and Goering, 1967). This constitutes 12-26% more new N than other annual estimates (mid July-mid October) from the same region. The between-station variability observed in both total N₂ fixation and primary productivity greatly emphasize the need for multiple stations and seasonal sampling strategies in biochemical studies of the Baltic Sea. The majority of new N from N₂ fixation was contributed by filamentous cyanobacteria. On average, cyanobacterial cells >20 μ m were able to supply a major part of their N requirements for growth by N₂ fixation in both 1998 (73%) and 1999 (81%). The between-station variability was high however, and ranged from 28-150% of N needed to meet the rate of C incorporation by primary production. Since the molar C:N rate incorporation ratio (C:N_{RATE}) in filamentous cyanobacterial cells was almost twice as high as the molar C:N mass ratio (C:N_{MASS}) in both years, we suggest that the diazotrophs incorporated excess C on a short term basis (for carbohydrate ballasting and buoyancy regulation), released nitrogen or utilized other regenerated sources of N nutrients. Measured rates of total N₂ fixation contributed only a minor fraction of 13% (range 4-24) in 1998 and 18% (range 2-45) in 1999 to the amount of N needed for the community primary production. An average of 9 and 15% of total N₂ fixation was found in cells <5 μ m. Since cells <5 μ m did not show any detectable rates of N₂ fixation, the ¹⁵N-enrichment could be attributed to regenerated incorporation of dissolved organic N (DON) and ammonium generated from larger diazotroph cyanobacteria. Therefore, N excretion from filamentous cyanobacteria may significantly contribute to the pool of regenerated nutrients used by the non-diazotroph community in summer. Higher average concentrations of regenerated N (ammonium) coincided with higher rates of N₂ fixation found during the 1999 transect and a higher level of ¹⁵N-enrichment in cells

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

References

Figures

Close

Title Page Introduction **Abstract** Conclusions **Tables** Back Full Screen / Esc Printer-friendly Version

EGU

Interactive Discussion

<5 μ m. A variable but significant fraction of total N₂ fixation (1–10%) could be attributed to diazotrophy in cells between 5–20 μ m.

1 Introduction

Surface blooms of filamentous diazotrophic cyanobacteria are well documented and a recurring phenomenon in the Baltic Sea in summer (cf. Leppänen et al., 1988). Due to their ability to fix N₂, diazotroph cyanobacteria may contribute significantly to the input of new N to the Baltic Sea ecosystem. The contribution of this source of new N has been subject of a number of investigations (e.g. Rinne et al., 1978; Leppänen et al., 1988; Rahm et al., 2000; Larsson et al., 2001; Wasmund et al., 2001). Our current knowledge may suggest large-scale regional differences in the abundance of the most common filamentous cyanobacteria in the Baltic Sea, Aphanizomenon sp. and Nodularia sp. (Kahru et al., 1994; Wasmund, 1997). During peak abundance in summer, maximum biomass of the filamentous cyanobacteria can be found in the southern and eastern parts of the Baltic proper (Niemistö et al., 1989). Although the filamentous cyanobacteria rarely dominate the autotroph community, conspicuous surface accumulations of the buoyant diazotrophs may congregate in wind and current driven rows or patches in coastal and off shore waters of the Baltic Sea (Niemistö et al., 1989; Walsby et al., 1995). Larsson et al. (2001) clearly demonstrated the great extent of inter-annual variability in timing and total abundance of filamentous cyanobacteria at an off shore station in the Baltic Sea. A similar spatial and temporal variability in measured rates of N₂ fixation during the growth season of diazotrophic cyanobacteria has been found by Ohlendieck et al. (2000) and Wasmund et al. (2001).

The earliest estimates of N_2 fixation in the Baltic Sea were based on the acetylenereduction method. Plankton samples were routinely concentrated in net-tows or by a screen from natural seawater. Due to this pre-treatment of the samples, the enrichment method may have excluded or even physiologically disturbed certain plankton groups and hence, underestimated total rates of N_2 fixation to the Baltic Sea. More

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ← ►I

← ► Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

recent studies however, have applied the sensitive ¹⁵N tracer method using natural water samples to measure N₂ fixation directly (Montoya et al., 1996; Ohlendieck et al., 2000; Wasmund et al., 2001). The ¹⁵N method opened the possibility to investigate the significance of new (nitrate, nitrite and N₂) and regenerated (ammonium) N incorporated into the autotroph community of Baltic Sea phytoplankton (e.g. Sörensson and Sahlsten, 1987; Sahlsten and Sörensson, 1989). A number of studies have also used the ¹⁵N method to monitor the direct transfer of newly fixed N into the planktonic foodweb (Pearl, 1984; Glibert and Bronk, 1994; Ohlendieck et al., 2000).

Historically, decay and cell lysis of cyanobacteria have been viewed as a significant source of regenerated N (Lindahl et al., 1980; Schultz and Kaiser, 1986) and high concentrations of dissolved organic nutrients have been found during and immediately after blooms of Baltic Sea cyanobacteria (Gundersen, 1981; Kuparinen et al., 1984). However, Ohlendieck et al. (2000) found that actively growing diazotroph filamentous cyanobacteria transferred DON to the picoplankton community during the early stages of a summer bloom. A similar observation was made by Gallon et al. (2002), who attributed the discrepancy between the acetylene reduction method (gross N₂ fixation) and the ¹⁵N-tracer method (net N₂ fixation) in filamentous cyanobacteria collected over several diel cycles to be DON excretion by the diazotrophs. Exoenzymes are instrumental in the regenerative process of dissolved organic matter and have been found in association with filamentous cyanobacteria in the Baltic Sea (Stoecker, et al., 2005). Therefore, filamentous cyanobacteria may not only be a source of DON for the nondiazotrophs, but also serve as an active centre of organic nutrient regeneration to the plankton community. Until recently, filamentous cyanobacteria were considered the only diazotroph organisms that contributed significantly to new production by N₂ fixation in the Baltic Sea. Nitrogen fixation by single cell cyanobacteria (Wasmund et al., 2001; Zehr et al., 2001; Montoya et al., 2004) however, may also be of importance and represents another source pathway of new N to the pelagic food-web.

The aim of this study was to assess the significance of N₂ fixation to new production during the early stages of a cyanobacterial bloom. During two successive years,

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◀ ▶I

■ Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

rates of N_2 fixation and primary production were monitored in early summer at 9 stations along a N-S transect covering the major parts of the Baltic proper. By exclusion filtration of natural seawater prior to and after in situ incubations, we identified the significance of single cell diazotrophs and the magnitude of DON excretion by filamentous cyanobacteria incorporated by organisms $<5\,\mu m$. By applying known molar C:N_{RATE} ratios for filamentous cyanobacteria, we estimated the significance of N_2 fixation contributing to new N growth in filamentous cyanobacteria.

2 Materials and methods

CTD profiles and seawater samples were collected at designated stations (Fig. 1) on two transect cruises in the Central Baltic Sea in 1998 (23 June-1 July) and 1999 (21 June–29 June). During a drift station experiment (see Gallon et al., 2001), three daystations were kept during 5 additional days in 1998 (6 July-11 July; drift experiment stations 12-14; Fig. 1). Physical parameters were measured using a Neil-Brown unit equipped with conductivity, temperature and density (CTD) sensors. Only temperature profiles are reported here. Seawater was collected at pre-determined depths using Hydrobios water samplers attached to the CTD unit (1998) or the same water samplers attached to a hydro-wire (1999). Upon retrieval on deck, a small quantity of water (approx. 150 mL) was collected immediately from each water sampler for nutrient analysis (see below). The majority of the water sample from each depth was transferred to a 20 L canister. Each canister was gently homogenized by inverting it a couple of times prior to sub-sampling. Sub-sample aliquots were immediately dispensed into parallel incubation bottles for in situ N₂ fixation and primary production incubations using the ¹⁵N and ¹⁴C tracer methods (see below) and for HPLC pigment analysis (see below). The incubation bottles were deployed at ambient depths on a free-floating array from 07:00-19:00 h (1998 transect) or 06:00-18:00 h (1999 transect). During the 3 drift experiment stations in 1998 (Fig. 1), incubations commenced from 04:00-16:00 h daytime (N₂ fixation and primary production) and 16:00-04:00 h at night (N₂ fixation).

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◆ ▶ I

◆ ▶ Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

At noon each day, samples were also collected for particulate organic carbon (POC), particulate nitrogen (PN) and particulate total phosphorus (PTP) analysis (see below).

2.1 Nutrient analysis

Nutrient samples were collected in the morning and afternoon at each station visited. The nutrient samples were stored at 4°C and analysed daily (typically within 4h of sampling) using an onboard auto-analyser. Dissolved inorganic nitrogen (DIN; nitrate, nitrite, ammonium) and soluble reactive phosphate (SRP) were determined according to the methods described by Grasshoff et al. (1983). A daily average using the morning and afternoon data was calculated for each station visited.

o 2.2 N₂ fixation

Sample aliquots were collected in replicates of 0.5 or 1 L polycarbonate bottles filled to the brim and sealed with a butyl rubber septum cap. Great care was taken to avoid airbubbles inside the sealed incubation bottles. A gas-tight syringe was used to inject 15 Nlabelled dinitrogen (N₂) gas (99 atom % ¹⁵N, Campro Scientific) and excess water was allowed to bleed out through a second injection needle to equalize pressure. Final ¹⁵N₂ enrichments ranged between 5.4-6.1 atom % N depending on the physical parameters of the sample water (Weiss, 1970). Each incubation was terminated by collecting the cells on pre-combusted (450°C, 6h) 25 mm Whatman GF/F filters using low vacuum (<100 mm Hg) filtration. At all stations in both years, one whole seawater sample from each depth was collected on a GF/F filter (total N₂ fixation) and one sample was screened through a 5 μ m membrane filter and cells <5 μ m were collected on a GF/F filter upon terminating the incubation. In another set of samples (see Table 2 for details) cells >20 µm were collected on a Nytex screen and transferred to a GF/F filter using GF/F filtered seawater. A number of samples were also screened prior to deployment and cells <5 µm were incubated separately as a control to test for N₂ fixed by this fraction, during the 1998 transect (all stations) and the 1999 transect (stations 9, 11,

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◀ ▶I

■ Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

13 and 15). During the 1998 transect (stations 6–9) cells <20 μ m were also separated prior to the in situ incubations to test for N₂ fixation by smaller sized cyanobacteria. Incubations with pre-screened cells <5 μ m and <20 μ m were terminated and collected on separate GF/F filters. Each filter sample was wrapped in pre-combusted aluminium foil and stored at –20°C until further analysis. In the laboratory, the samples were dried at 60°C, wrapped in tin cups (Heraeus CHN cups) and packed into pellets. The ¹⁵N enrichments were analysed on a Europa Mass Spectrometer 20–20, with an ANCA-SL preparatory unit (Crewe, England, UK). The mass spectrometer was standardised with bovine serum albumin and the precision of the analytical procedure, as measured by standard deviation of the nitrogen isotope, was typically 0.04%. The rate of N₂ fixation was calculated according to Montoya et al. (1996).

2.3 Primary production

Sample water was collected in acid cleaned (10% hydrochloric acid) 250 mL Erlenmeyer polycarbonate incubation bottles. Three replicate incubation bottles, a time zero and a dark bottle were collected for each depth. The primary production method was adapted to the trace-metal clean procedures outlined by Fitzwater et al. (1982). Under low light conditions ^{14}C -labelled sodium bicarbonate was added to each flask at the start of the assay (final specific activity = 1.6 μg C nCi $^{-1}$) and the time zero was immediately filtered and processed. At the end of the incubation and under low light conditions, each bottle was immediately filtered onto a 25 mm Whatman GF/F filter. Thereby particles >20 μm were collected separately at all stations in both years. The size fraction >20 μm was collected on a Nytex screen and transferred to a 25 mm Whatman GF/C filter using GF/F filtered seawater. Each filter was placed in a 20 ml scintillation vial and added 250 μl HCl (0.5%) until further processing after each cruise. In the laboratory all sample vials were uncapped and the residual ^{14}C -bicarbonate was left to evaporate off the filters in a fume hood. Ultima Gold XR scintillation cocktail was added to each filter sample and the activity was counted on a Wallac 1409 Liquid

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



Scintillation Counter using the external channel ratio method.

2.4 HPLC pigment analysis

Sample water aliquots of 2 L were filtered onto Whatman GF/F filters. Cells >20 µm were collected separately at stations 1-7 in 1998 and on all stations (except stations 9-11) in 1999. The size fraction >20 μ m was collected on a Nytex screen and transferred to a 25 mm Whatman GF/C filter using GF/F filtered seawater. The filter samples were wrapped in aluminium foil and stored in liquid nitrogen until further analysis. Each pigment sample was placed in 2.5 ml (90%) acetone and, to ensure complete extraction, each sample was added a mixture of 2 and 4 mm glass beads and shaken for 5 min in a cooled Vibrogen cell mill (Buehler, Germany). Subsequently, the extracts were centrifuged (5000 rpm) for 10 min at -10°C and the supernatant analysed according to the method of Derenbach (1969). The entire processing and analysis was executed under low light conditions to prevent photo-oxidation of the pigments. Pigment concentrations were determined by reverse-phase high-performance liquid chromatography (RP-HPLC) according to the method of Barlow et al. (1997) using a diode array spectrophotometer (Waters) and commercially available pigment standards (Sigma Chemical Company, USA and International Agency for ¹⁴C Determination, Denmark). The pigment composition from each sample was used to calculate the relative proportion of phytoplankton classes to total chlorophyll a (Chl a) using the CHEMTAX program (Mackey et al., 1996). Only the fraction of total Chl a associated with filamentous cyanobacteria is shown in this study. The ratio between the class specific pigment of filamentous cyanobacteria (echinenone) and Chl a was established from natural samples of the filamentous diazotrophs collected specifically for this purpose (Meyerhöfer, 1994).

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



2.5 POC and PN analysis

Sample aliquots of $0.5 \, \text{L}$ from each depth were filtered onto pre-combusted ($5 \, \text{h}$, $500^{\circ} \, \text{C}$) 25 mm Whatman GF/F filters, that were wrapped in pre-combusted aluminium foil and stored at $-20^{\circ} \, \text{C}$. Particles $> 20 \, \mu \text{m}$ were collected separately at all stations in both years. A sample aliquot of 1 L was collected on a $20 \, \mu \text{m}$ Nytex screen and the retained particles transferred to a 25 mm Whatman GF/C filter. Prior to analysis, the filters were dried at $65^{\circ} \, \text{C}$, fumigated by concentrated HCl, dried again and packed in tin capsules. The POC and PN content was analysed using a Control Equipment Corporation 240-XA Elemental Analyzer (1998) or a Carlo-Erba Strumentazione Elemental Analyser (1999).

2.6 PTP analysis

Sample aliquots of $0.5\,L$ were filtered onto HCl washed $25\,\text{mm}$ Whatman GF/F filters. Particles $>\!20\,\mu\text{m}$ were collected separately at all stations in both years. A sample aliquot of $1\,L$ was collected on a $20\,\mu\text{m}$ Nytex screen and the retained particles transferred to a $25\,\text{mm}$ Whatman GF/C filter. Analysis of cellular P by the ashing-hydrolysis method was performed as discussed in Solorzano and Sharp (1980). The filters were soaked in phosphate-free, deionised water (particle free) and were dried prior to use. The filter-collected samples were rinsed with $0.17\,\text{M}$ Na $_2\text{SO}_4$ and stored in acid cleaned containers at $-20\,^{\circ}\text{C}$. Prior to analysis, the samples were thawed and $0.017\,\text{M}$ MgSO $_4$ was added before being left overnight at $95\,^{\circ}\text{C}$. The samples were then combusted at $500\,^{\circ}\text{C}$ for $3\,\text{h}$ and the remaining P was extracted in 2 ml of $0.2\,\text{M}$ HCl at $80\,^{\circ}\text{C}$ for a minimum of $1\,\text{h}$. The acid extraction solution was centrifuged ($30\,\text{min}$, $5000\,\text{rpm}$), and the supernatant analysed as for SRP according to Grasshof et al. (1983).

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



Results

Average surface water temperature between the two years was approximately 2°C higher in 1999 than in 1998 (Fig. 2). The thermocline was more pronounced in 1999 and during the drift experiment stations in 1998 (Fig. 2). DIN $(0.1-0.2 \,\mu\text{M})$ was slightly higher in 1999 than during the 1998 transect with average concentrations of ammonium twice as high in 1999 (Fig. 3A, B). The average concentration of SRP in surface waters (0–14 m) was uniformly low during the 1999 transect (0.08 μ M) (Fig. 3C, D) and slightly higher in 1998 (average of $0.2 \mu M$ at the southern stations and $0.05 \mu M$ at the northern stations). Due to these differences between the years, the median (geometric mean) of the molar N:P ratio (1-3) was almost twice as high during the 1999 transect. Cursory microscopy of live samples during the two cruise transects showed that Aphanizomenon sp. was the predominant filamentous cyanobacteria in surface waters at the northern stations (data not shown), whereas *Nodularia* sp. appeared more frequent at the southern stations.

Total Chl a as a function of depth did not show any pronounced inter-annual differences (Fig. 4A). Integrated rates of total N₂ fixation were a fraction of 0.71 lower at night than in daytime at the drift station (Table 1). The daytime rates of N₂ fixation measured on all transect stations in both years, were factor corrected for the night-time activity to yield daily rates (Fig. 4B). Since primary production was measured from dawn to dusk, no factor correction was used for this parameter. With the exception of the drift experiment stations, the rates of N₂ fixation and primary production were twice as high in surface waters in 1999 compared to 1998 (Fig. 4B, C). Very low or no rates of N₂ fixation and primary production were detected below 14 m depth in both transects. Integrated median rates of total N₂ fixation (0-14 m) were 0.5 mmol N m⁻²d⁻¹ in 1998 (range 0.1–0.8) and 1.0 mmol N $m^{-2}d^{-1}$ in 1999 (range 0.3–2.7). Integrated median rates of total primary production were 32.3 mmol C m⁻²d⁻¹ in 1998 (range 20.4–40.9) and 57.9 mmol C m⁻²d⁻¹ in 1999 (range 25.9–85.2).

Rates of ^{15}N enrichment in cells $<5 \,\mu\mathrm{m}$ (separated after incubation) were plotted as

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page Introduction **Abstract** Conclusions References **Tables Figures** Back Close Full Screen / Esc Printer-friendly Version Interactive Discussion

▶I

a linear function of total N_2 fixation for both transects in 1998 and 1999 (Fig. 5) and the average fraction of ^{15}N enrichment in cells $<5\,\mu\text{m}$ (the slope) was 0.09 in 1998 and 0.15 in 1999. Cells $<5\,\mu\text{m}$ (separated prior to incubation) showed no detectable uptake. Cells $<20\,\mu\text{m}$ (separated prior to incubation) at stations 6–9 in 1998 showed low rates of N_2 fixation (median fraction 0.05 of total N_2 fixation) but the ratios were variable (range 0.01–0.1) between stations (Table 2).

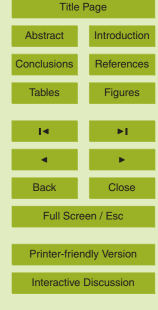
Diazotroph specific Chl a was plotted as a linear function of Chl a>20 µm (Fig. 6). The fraction of N_2 fixation in cells >20 μ m ranged between 0.92–0.88 of the total rate in surface waters in 1998, and was 0.48 of the total at 14 m depth (Table 3). The fraction of N_2 fixation in cells >20 μ m was 0.81–0.85 in surface waters in 1999 (Table 3). The average fractions of N₂ fixation in cells >20 µm were used to calculate the integrated rate of N_2 fixation in cells >20 μ m at each station of both transects. In 1999, the depths without data were substituted with the results from the 1998 transect (Table 3). The fraction of primary production in cells >20 µm varied widely between stations in each year (Table 4) but the median value in 1998 (0.17) was similar to the median calculated in 1999 (0.16). The median fractions of integrated ChI a and PTP in particles $>20 \,\mu m$ were similar in both years and approximately one fifth of the total (Table 4). The fractions of integrated POC and PN in particles $>20 \mu m$ (relative to total amount) were slightly higher and approximately one third of the total (Table 4). Both the molar elemental ratios of C:N in particles $>20 \,\mu\text{m}$ (C:N_{MASS}>20 μm) and total integrated $C:N_{MASS}$ were lower in 1999 than in 1998 (Table 4). The median of total $C:P_{MASS}$ was higher in 1999 and the integrated C:P in particles $>20 \,\mu\text{m}$ (C:P_{MASS}>20 μ m) varied little between the two years, but was higher than total C:P_{MASS} (Table 4). The median of the total C:N incorporation ratio was more than 5 times higher than the molar C:N incorporation in cells >20 μ m (C:N_{BATE} >20 μ m) and both size fractions were similar between the years (Table 4). The calculated C:N_{BATE} in cells >20 μ m was plotted as a function of daily integrated PAR (Fig. 7).

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



4 Discussion

4.1 Early blooms of diazotrophic cyanobacteria

We found a striking difference in surface water hydrography between the two summer transects investigated in 1998 and 1999. Surface waters during the 1999 transect was warmer and had a variable, but more pronounced thermocline than in 1998. This difference appeared to be reflected in the overall plankton community production. Given the great variability between stations in each year (Fig. 3) there was a general trend towards higher rates of both N₂ fixation and primary production in surface waters in 1999. This was also evident in the median rates of total N₂ fixation and primary production that were a factor of 2 higher during the 1999 transect. The hydrography at the drift stations visited 5 days after the 1998 transect (st. 12-14) were characterized by a stronger thermocline and slightly warmer surface waters than observed earlier in the cruise that year. The higher rates of N₂ fixation and primary production at the drift station (Fig. 3B, C) may suggest that, once in place, the plankton community was able to respond quickly to changes in hydrography and the strengthening of the thermocline. Integrated estimates by Wasmund et al. (2001) for the central Baltic Sea in 1997–1998 showed that 75 mmol N m⁻² was incorporated in surface waters in summer (mid July-mid August). Their estimate for late summer and fall amounted to 50 mmol N m⁻² (mid August-mid October), but Wasmund et al. (2001) did not include the early summer phase of the bloom of filamentous cyanobacteria. Assuming that the early onset of the bloom lasted about a month, our study may suggest that a significant amount of N₂ fixation contributed to new production in 1998 (15 mmol N m⁻²) and 1999 (33 mmol N m⁻²). By including the early bloom, 12-26% more N can be added to the annual estimate given by Wasmund et al. (2001), which brings the annual budget in the range of 140–158 mmol N m⁻² yr⁻¹. The between-station variability in both primary production and N₂ fixation that appeared in this study greatly emphasize the need for implementing spatial as well as temporal sampling strategies in ecological studies of

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

the Baltic Sea.

4.2 Primary production and N₂ fixation in filamentous cyanobacteria

We found that cells collected in the $>20 \,\mu m$ fraction were predominantly filamentous cyanobacteria (Fig. 6) and the contribution of N₂ fixation to new production in open waters of the Baltic Sea was almost entirely found in cells >20 μ m (Table 3). Gallon et al. (2002) found that filamentous cyanobacteria incorporate C and N at ratios much higher than the Redfield relationship of 6.6 in surface waters. By applying these C:N incorporation ratios to our specific depths (C:N_{BATE} [0-7 m]=17.6; [7-14 m]=5.1; Gallon et al., 2002) we calculated how much new N from N₂ fixation did support the measured rates of primary production. Integrated rates of N₂ fixation in filamentous cyanobacteria contributed an average of 73% in the 1998 transect (range 28-119%) and 81% in 1999 (range 46–150%) towards the measured rates of primary production in cells >20 μ m. This may suggest that filamentous cyanobacteria were able to support their photosynthetic growth requirement with new N at some of the stations, while at other stations they were not able to compliment their growth by diazotroph N at the days visited. The average integrated C:N_{RATE} ratios in cells >20 µm in this study (Table 4) were almost two times higher than the Redfield relationship of 6.6 in both years but showed high variability in both years (C:N_{RATE} ranged between 7-28). Gallon et al. (2002) found a similar C:N_{BATE} ratio to ours and noted a temporal uncoupling between C and N incorporation rates in the filamentous diazotrophs investigated over 3 diel cycles. In high light conditions, filamentous cyanobacteria do incorporate excess amounts of C as glucose, lipids and polyhydroxybutyrate storage products (Stal and Walsby, 1998). Loss of buoyancy results from excessive C-incorporation and carbohydrate ballasting (Ibelings et al., 1991) limiting the exposure of filamentous cyanobacteria to UV irradiance in the top layer of the water column. In our study integrated C:N_{RATE} in cells >20 μ m was frequently higher on days with high surface irradiance (Fig. 7) showing that measured rates of primary production and N₂ fixation were at times out of phase. These rates were not reflected however, in C:N_{MASS} of all particles combined or >20 μ m (Table 3).

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◀ ▶I

■ Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

Indeed, the C:N_{MASS} ratios that were similar to the Redfield relationship, may suggest that excess C incorporated was stored on a very short term basis (<24 h). Also a low cellular P content causes loss of buoyancy in filamentous cyanobacteria (Konopka et al., 1985; Klemer et al., 1995) and particles >20 μ m had a low P content in our study (Table 4). Low P is commonly found in filamentous cyanobacteria in the Baltic Sea in summer (Larsson et al., 2001; Nausch et al., 2004).

Other studies in the Baltic Sea have found that diazotrophs can meet (Sörensen and Sahlsten, 1987) or even exceed (Stal and Walsby, 2000) their growth requirements for new N by N₂ fixation. The biogeochemical estimates by Larsson et al. (2001), based on measurements of total nitrogen, also suggested that Baltic Sea cyanobacteria were able to assimilate more new N through N₂ fixation, than what was needed to support their growth. The Gallon et al. (2002) study noted a discrepancy in the rates of N₂ fixation determined by the acetylene reduction method (gross N₂ fixation) and the ¹⁵Ntracer method (net N₂ fixation). They concluded that a significant amount of newly fixed N may have been released by the filamentous cyanobacteria (Gallon et al., 2002) and this was not being accounted for with the ¹⁵N tracer method used in this study. We found evidence of exudation of newly fixed N (see below) and this may be a reason for the discrepancy between our study and the gross estimates of N₂ fixation mentioned above (Stal and Walsby, 2000; Sörensen and Sahlsten, 1987; Larsson et al., 2001). A crude comparison between these gross estimates from literature and our results based on the ¹⁵N method (73% and 81%, see above), indicate that filamentous cyanobacteria may excrete as much as one third of newly fixed N during the course of a day. Stal et al. (2003) however, suggested that up to 80% of recently fixed N can at times be released over shorter periods of time in surface waters of the Baltic Sea. High C:N_{BATE} ratios in filamentous cyanobacteria may also be caused by uptake of other sources of N (Sörensson and Sahlsten, 1987). Mulholland et al. (2004) found rapid uptake of previously released NH₄ and DON in their cultures of Trichodesmium. Evans et al. (2000) found that nitrogenase activity in filamentous cyanobacteria decreased over the course of a day in the Baltic Sea and we suggest that this can be attributed to

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



simultaneous uptake of regenerated N. Therefore, the exceedingly high C:N_{RATE} ratios found in this study may be as a consequence of both excess C incorporation as well as release of newly fixed N and regenerated N incorporation.

4.3 Diazotroph release of DON and single cell N₂ fixation

Cells $<5 \,\mu m$ incorporated 9% in 1998 and 15% in 1999 of total N₂ fixation and the same size fraction showed no detectable rates of N2 fixation in this study. Therefore, the observed ^{15}N enrichment in cells $<5 \,\mu m$ may have resulted from recent N_2 fixation excreted as DON or NH₄ and incorporated by non-diazotroph phytoplankton. Cyanobacteria are known to exudate nitrogenous material as DON (Flynn and Gallon, 1990; Glibert and Bronk, 1994) and NH₄ (Mulholland and Capone, 2001) at times of excess N₂ fixation relative to cellular synthesis (Gallon et al., 2002). Our study confirms the findings of Ohlendieck et al. (2000), who suggested that newly fixed N was transferred from filamentous cyanobacteria to smaller plankton. In their study, an increase in abundance of Synechococcus sp. was observed between cruises and over the course of a month, and this co-occurred with a developing bloom of filamentous cyanobacteria in the central Baltic Sea in 1995. Ohlendieck et al. (2000) suggested that the observed increase in biomass of the non-diazotroph cyanobacteria may have been fuelled by newly excreted DON from the larger filamentous diazotrophs. Pico-cyanobacteria are strongly N-limited in the Baltic Sea and may benefit from diazotroph cyanobacteria as a nutrient source, since they show similar temporal and spatial distribution (Stal et al., 2003). The release of N by filamentous cyanobacteria represents an important and significant way of making new N available for the pelagic food-web at the early stage of cyanobacterial blooms in the Baltic Sea.

By monitoring total N in surface waters of the Baltic Sea in summer, Larsson et al. (2001) estimated the annual new production by N_2 fixation and found a sizeable amount of N that could not be attributed to their estimates of filamentous cyanobacteria biomass (determined from cell counts). Larsson et al. (2001) concluded that the N that could not be accounted for may have been generated by DON excretion from 1293

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◆ ▶I

◆ ▶ Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

filamentous cyanobacteria, or generated by other diazotrophs that were not investigated in their study. N_2 fixation by single cell cyanobacteria represents another source pathway of newly fixed N to the pelagic food-web in early summer. In our study we found N_2 fixation in the pre-screened cells <20 μ m (Table 2). Since the group specific pigment echinenone was not found in size fractions <20 μ m, we assume that this size fraction did not contain any significant amounts of filamentous cyanobacteria. Wasmund et al. (2001) also detected N_2 fixation in the smaller size fractions of Baltic Sea phytoplankton. They estimated that cells <10 μ m contributed 43% of total N_2 fixation in the upper 20 m of the water column (Wasmund et al., 2001) and suggested that this was caused by non-heterocystous, coccoid cyanobacteria. Recent studies have also demonstrated that significant amounts of active single cell diazotrophs are found in temperate regions of the northern Pacific (Zehr et al., 2001; Montoya et al., 2004). Since we did not detect any significant rates of N_2 fixation in cells <5 μ m in our daylight incubations, we conclude that cells between 5–20 μ m in the Baltic Sea in early summer may at times use N_2 fixation to support their nutrient requirements.

4.4 The importance of N₂ fixation to total primary productivity

Assuming the same C:N_{RATE} assimilation ratios as above (Gallon et al., 2002), integrated rates of total N₂ fixation contributed 13% in 1998 (range 4–24%) and 18% in 1999 (range 2–45%) to total primary production. In recent years, a wide range in C:N_{RATE} ratios has emerged from Baltic Sea studies. By using the 15 N-tracer method, Wasmund et al. (2001) found a C:N_{RATE} of 6.8 at the height of the bloom of filamentous cyanobacteria in August 1997. During other stages of the bloom however, molar C:N_{RATE} approached 20 to >100 and in August the year after, only 17% of the average N demand could be met by N₂ fixation (Wasmund et al., 2001). Studies taking into account gross N₂ fixation on the other hand, e.g. acetylene reduction by Stal and Walsby (2000) and total N biogeochemistry by Larsson et al. (2001), have generally measured a higher contribution (>50%) to new production. This discrepancy can to a large extent be attributed to N excretion (see Gallon et al., 2002) that is not being accounted for

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



in N₂ fixation estimates using the ¹⁵N-tracer method. Sörensson and Sahlsten (1987) however, measured N₂ fixation by the acetylene reduction method and found that N₂ fixation contributed only 16% of the total N utilized by the phytoplankton community at station BY31 (stations 8 in 1998 and 9 in 1999 in this study). The low contribution to new production by N₂ fixation in the Sörensson and Sahlsten (1987) study may suggest that other nutrient sources (such as regenerated N) were utilized by the phytoplankton community (including the diazotroph cyanobacteria). The Sörensson and Sahlsten (1987) study took place in late summer during the last stages of a cyanobacterial bloom. At that time we can assume that there was a closely coupled food-web dominated by regenerated production with ammonium as one of the major sources of N. In contrast, our study took place at the beginning of the cyanobacterial growth season with a plankton community still attuned towards new production and relatively low levels of ammonium. We found that 9 and 15% of the total rate of N2 fixation (dominated by filamentous cyanobacteria), was incorporated by cells $<5 \,\mu m$ in early summer (Fig. 5). This is almost a factor of ten less than what Sahlsten and Sörensson (1989) found during a declining cyanobacterial bloom in the same area (71–93% of total N). In our study we found relatively low concentrations of ammonium (median conc. $0.05 \mu M$) in 1998 and the median concentration of ammonium in 1999 was almost twice as high (approx. $0.1 \mu M$). The higher median values of ammonium in surface waters during the 1999 transect (Fig. 3) also coincided with higher rates of N₂ fixation during that transect and a higher level of 15 N-enrichment in cells <5 μ m (Fig. 5). The diel increase in ammonium concentrations reported by Stal et al. (2003) for the 1999 transect may be as a result of enzymatic regeneration of newly released DON or release of NH₄ and hence, the larger filamentous diazotrophs may have contributed new N to the pool of regenerated nutrients. Surface waters of the Baltic Sea in summer are characterized by a low N:P nutrient ratio and we can assume that regenerated nutrients are readily utilized by the non-diazotroph components of the phytoplankton community. Therefore, we concur with Larsson et al. (2001) in that N₂ fixation may sustain a considerable fraction of net community production in the Baltic Sea in summer. Rolff (2000) also demonstrated

BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page Introduction **Abstract** Conclusions References **Tables Figures** Back Close Full Screen / Esc Printer-friendly Version Interactive Discussion

that the effect of N_2 fixation (trophic $\delta^{15}N$ enrichment) was found in all size fractions of phytoplankton in the Baltic Sea in late summer. Hence, N_2 fixation in summer may contribute significantly to new as well as regenerated production in surface waters of the Baltic Sea.

5 Summary

This study shows that N_2 fixation by filamentous cyanobacteria may provide a significant input to new production in the surface waters of the Baltic Sea in early summer. Disproportionate amounts of C are at times incorporated relative to N in filamentous cyanobacteria. Since the C: N_{MASS} is fairly constant and similar to the Redfield relationship, we assume that excess C is not stored in cells over longer periods of time and possibly used within the course of a day. Filamentous cyanobacteria and non-diazotroph phytoplankton may also utilize significant amounts of regenerated N. Even though the phytoplankton community is dominated by non-diazotroph organisms in the Baltic Sea, a small fraction of single cell diazotrophs may at times contribute significantly towards new production. Total rates of N_2 fixation, measured by the N_2 tracer method, contribute only a minor portion of new N to the phytoplankton community in summer. DON release from filamentous diazotroph cyanobacteria however, may contribute significantly towards the utilization of regenerated N by the non-diazotroph community of phytoplankton.

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BGD

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◆ ▶I

◆ ▶ Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

References

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BGD

3, 1279–1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page Introduction Abstract Conclusions References **Tables Figures** 14 **▶**I Back Close Full Screen / Esc Printer-friendly Version Interactive Discussion

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20

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BGD

3, 1279–1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◀ ▶I

■ Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

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3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◆ ▶I

◆ ▶ Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

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3, 1279–1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



Table 1. Rates of N_2 fixation measured in daylight (04:00–16:00 h) and at night (16:00–04:00 h) in whole seawater from the Baltic Sea in 1998. The rates from each station were integrated (0–14 m) and the N_2 fixation at night was calculated as a fraction of the daytime.

Station	Day nmol N m ⁻² h ⁻¹	Night nmol N m ⁻² h ⁻¹	Night:Day
12	30.34	21.65	0.71
13	30.45	21.54	0.71

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page			
Abstract	Introduction		
Conclusions	References		
Tables	Figures		
I∢	►I		
	•		
Back	Close		
Full Screen / Esc			
Printer-friendly Version			
Interactive Discussion			

Table 2. Integrated rates of N_2 fixation (0–14 m) in whole seawater and cells <20 μ m in the Baltic Sea in 1998. Cells <20 μ m were separated prior to the in situ daytime incubations and the rates of N_2 fixation in cells <20 μ m were also calculated as a fraction of the total rate.

Station	Total N ₂ fixation μ mol N m ⁻² h ⁻¹	N_2 fixation <20 μ m μ mol N m ⁻² h ⁻¹	Fraction <20 μm ratio
6	27.15	0.18	0.01
7	22.51	2.06	0.09
8 ^(*)	18.13	1.79	0.10
9	38.21	3.89	0.10

^{(*)=0-9} m integration (sample at 14 m depth missing)

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page			
Abstract	Introduction		
Conclusions	References		
Tables	Figures		
I∢	►I		
4	•		
Back	Close		
Full Screen / Esc			
Printer-friendly Version			
Interactive Discussion			

Table 3. Rates of N_2 fixation measured in cells larger and smaller than $20 \,\mu\text{m}$ (separated after incubation) in the Baltic Sea in 1998 (stations 6–9) and 1999 (stations 7–15). The rates from each station were averaged per depth and the rate of N_2 fixation in cells >20 μ m was calculated as a fraction of the total rate.

Depth m	Total N ₂ fixation nmol N L ⁻¹ h ⁻¹ (range)	N_2 fixation >20 μ m nmol N L ⁻¹ h ⁻¹ (range)	Fraction >20 μm ratio (range)
1998:			
0	3.08 (2.06-5.18)	2.86 (1.74-4.54)	0.92 (0.84-1.00)
4	2.68 (2.14-4.36)	2.38 (1.70-3.80)	0.89 (0.82-0.97)
9	1.16 (0.54–1.54)	1.00 (0.48–1.28)	0.88 (0.81-1.00)
14	0.22 (0.12–0.40)	0.12 (0.04–0.26)	0.48 (0.29–0.65)
1999:			
0	5.42 (1.22-13.52)	4.28 (1.02-10.92)	0.81 (0.69-0.99)
4	5.38 (1.44-11.96)	4.50 (1.22–10.12)	0.85 (0.70-1.00)

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page			
Abstract	Introduction		
Conclusions	References		
Tables	Figures		
I₹	►I		
■	•		
Back	Back Close		
Full Screen / Esc			
Printer-friendly Version			
Interactive Discussion			

Table 4. Integrated mass and primary production in cells >20 μ m calculated as the fraction of the total, during two transect cruises (1998, 1999) in the Baltic Sea. The molar ratios of integrated mass and elemental incorporation in two size fractions (total mass and particles >20 μ m) were also calculated.

	1998	1999
	median (range)	median (range)
Fraction >20 μm (0–14 m):		
Primary production	0.17 (0.13-0.25)	0.16 (0.06-0.27)
Chl a	0.16 (0.07-0.27)	0.24 (0.12-0.46)
POC	0.37 (0.31-0.43)	0.27 (0.18-0.58)
PN	0.34 (0.31-0.42)	0.29 (0.18-0.68)
PTP	0.18 (0.12–0.31)	0.20 (0.10-0.36)
Integrated molar ratios (0–14 m):		
$C:N_{MASS}>20 \mu m$	10.1 (7.8–12.0)	6.8 (5.8–7.8)
C:N _{MASS} total	9.2 (8.6–10.2)	7.4 (7.1–7.9)
$C:P_{MASS}>20 \mu m$	204.7 (124.5-285.3)	195.9 (127.3-355.1)
C:P _{MASS} total	98.4 (84.2-113.5)	145.3 (123.8-166.5
$C:N_{BATF}>20 \mu m$	12.1 (8.2–27.7)	11.6 (6.5–17.6)
C:N _{RATE} total	65.0 (36.9–185.0)	60.0 (22.3–249.6)

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page		
Abstract		Introdu
Conclusions		Refere
Tables		Figu
I∢		•
•		•
Back		Clo
Full Scr	ee	n / Esc
Printer-frie	nc	lly Vers

EGU

Interactive Discussion

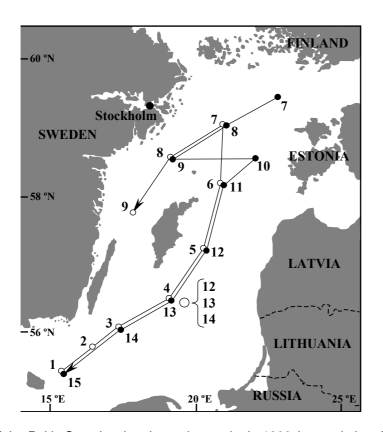
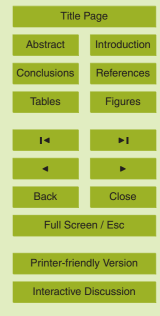


Fig. 1. Map of the Baltic Sea showing the cruise tracks in 1998 (open circles, S-N) and 1999 (closed circles, N-S). Location of the drift experiment stations in 1998 (larger open circle) 12–14 are also shown.

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



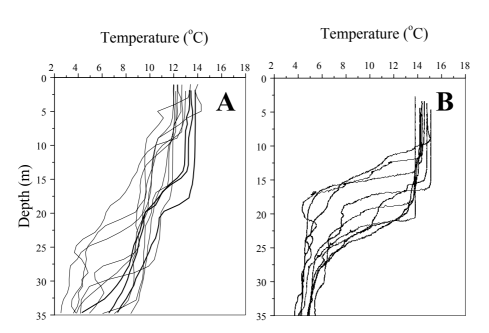


Fig. 2. Seawater temperature measured as a function of depth at the transect stations in 1998 **(A)** and 1999 **(B)**. Temperature profiles at the drift stations 12–14 (thick line) are also shown.

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



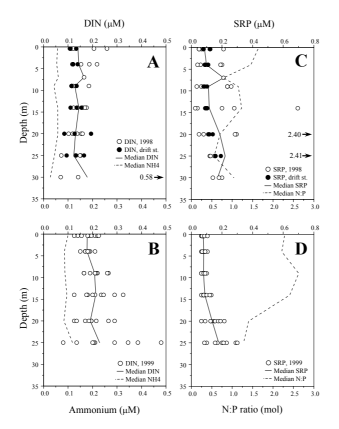


Fig. 3. DIN concentrations (open circles) as a function of depth in 1998 **(A)** and 1999 **(B)**. The geometric average of DIN (whole line) and ammonium (dashed line) is shown for each cruise transect each year. SRP concentrations (open circles) as a function of depth in 1998 **(C)** and 1999 **(D)**. The geometric average of SRP (whole line) and the molar N:P ratio (dashed line) is shown for each cruise in each year. Nutrient sample were also collected at the drift station casts 12–14 (whole circles). Stations 1, 2, 6 and 7 in 1998 did not have all nutrients analysed and were not included in the graph.

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



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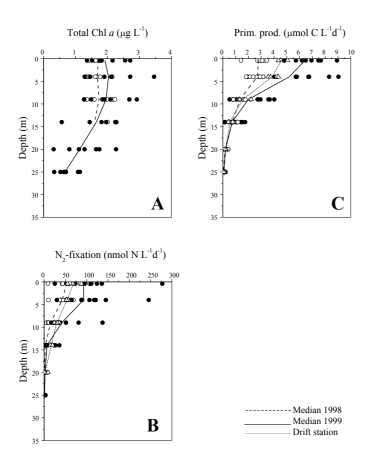


Fig. 4. Total Chl a (**A**), rates of N_2 fixation (**B**) and primary production (**C**) plotted as a function of depth in 1998 (open circles) and 1999 (closed circles). Where available, the geometric average at each depth was calculated for the 1998 transect cruise (dashed line) and drift station (dotted line) and the cruise transect in 1999 (whole line).

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page



Printer-friendly Version

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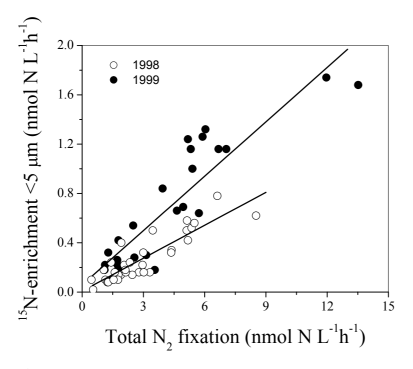
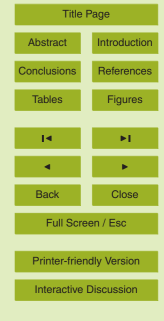


Fig. 5. Rates of ¹⁵N enrichment in cells <5 μ m (screened after incubation) as a function of total N₂ fixation measured from water samples at the transect stations 1–9 and at the drift experiment stations 12–14 in 1998 (open circles) and from the transect stations 7–15 in 1999 (closed circles). Solid lines shows the linear regression in 1998; y=0.089(±0.009)·x+0.008(±0.030), n=33, R=0.881, P<0.0001; and in 1999; y=0.147(±0.014)·x+0.058(±0.075), n=26, R=0.904, P<0.0001. Rates of ¹⁵N enrichment in cells <5 μ m at stations 1–2 in 1998 were below detection and not included in this figure.

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



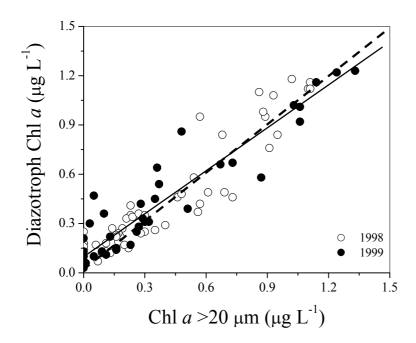


Fig. 6. Diazotroph Chl *a* plotted as a function of Chl $a>20\,\mu\mathrm{m}$ in samples collected from the cruise transects in 1998 (open circles) and 1999 (closed circles). Dashed line is the 1:1 relationship and the solid line shows the linear regression of both cruises combined; $y=0.872(\pm0.036)\cdot x+0.098(\pm0.019)$, n=90, R=0.933, P<0.0001.

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.



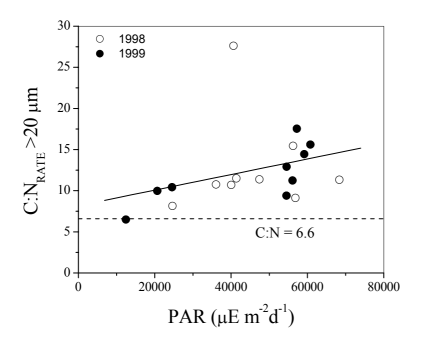


Fig. 7. Integrated C:N_{RATE}>20 μ m as a function of daily integrated PAR in 1998 (open circles) and 1999 (closed circles). The dashed line is the Redfield relationship and the solid line is the linear regression; y=0.000095(±0.000069)·x+8.18(±3.31), n=18, R=0.322, P=0.192.

3, 1279-1311, 2006

Significance of nitrogen fixation in early summer

U. Ohlendieck et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

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