INTRODUCTION

About 20 yr ago, the central role of bacteria and flagellates in many marine pelagic systems was recognized. The term 'microbial loop' was introduced (see also Azam et al. 1983) to describe the coupling between eukaryotic phytoplankton, the particulate and dissolved organic carbon they produce, bacteria (heterotrophic, photoheterotrophic, photoautotrophic, or chemoautotrophic), archaea, viruses and their pro-

Microbial dynamics in autotrophic and heterotrophic seawater mesocosms. I. Effect of phytoplankton on the microbial loop

Uta Passow¹,*, Christina L. De La Rocha¹, Carol Arnosti², Hans-Peter Grossart³, Alison E. Murray⁴, Anja Engel¹

¹Alfred-Wegener-Institut für Polar- und Meeresforschung, Am Handelshafen 12, 27570 Bremerhaven, Germany
²Department of Marine Sciences, University of North Carolina, Chapel Hill, North Carolina 27599-3300, USA
³Institute of Freshwater Ecology and Inland Fisheries, Dept. of Limnology of Stratified Lakes, Alte Fischerhütte 2, 16775 Stechlin, Germany
⁴Division of Earth and Ecosystem Science, Desert Research Institute, 2215 Raggio Parkway, Reno, Nevada 89512, USA

ABSTRACT: The effect of phytoplankton on the other compartments of a simple microbial loop consisting of dissolved matter, bacteria and nanoflagellates was investigated in three 1000 l mesocosms. These mesocosms were inoculated with a natural bacterial community and (1) no other additions (No Addition tank), (2) Phaeocystis globosa (Phaeocystis tank), or (3) 2 diatom species (Diatom tank). During the 20 d experiment, autotrophic activity was negligible in the No Addition tank. In contrast a small P. globosa bloom developed in the Phaeocystis tank and a large diatom bloom dominated the Diatom tank. In this paper we describe the experiment, the changes in chlorophyll a and heterotrophic nanoflagellate concentrations, as well as the cycling of nitrogen, phosphorus, and silica. Then we provide a synthesis of the structure and functioning of the microbial loops in these 3 systems using cluster analysis, a statistical pattern recognition tool. The goal was to test the hypothesis that differences in the resident phytoplankton populations would be reflected in (1) the composition and concentration of dissolved organic matter, (2) the composition of the bacterial community, (3) the food web, and (4) the cycling of elements and organic matter. In all 3 mesocosms, nitrate and silicic acid remained abundant. Orthophosphate was preferred by diatoms, whereas Phaeocystis appeared to prefer dissolved organic phosphorus. The hypothesis that phytoplankton composition shapes the structure and functioning of the microbial loop was partially supported: 6 d after inoculation each mesocosm exhibited a distinct organic matter signature. After 10 to 12 d, concentrations of heterotrophic nanoflagellates were high enough to exert significant grazing pressure in all 3 mesocosms. A parallel shift in bacterial community composition was visible in all mesocosms at this time, possibly reflecting grazing pressure. The food-web structure developed divergently in the 3 mesocosms during the second half of the experiment. Differences in biochemical cycling between mesocosms were predominantly driven by the large quantitative differences in autotrophs.

KEY WORDS: Mesocosm · Microbial loop · Structure and function · Food-web structure · Cluster analysis

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tist grazers, as well as mixotrophic protists. The relationship of the different components of the microbial loop to one another is complex, with each compartment both influencing and being influenced by the others. Bacterial activity and community composition, for example, modify and are modified by the quality and quantity of dissolved organic matter (DOM), which, in turn, is a result of DOM release, which is a function of phytoplankton composition, growth and exudation (Pinhassi et al. 1999, Riemann et al. 2000, Castberg et al. 2001, Fandino et al. 2001, Schäfer et al. 2001), grazing (Šimek et al. 1997, 2001, Lebaron et al. 1999), and viral lysis (Fuhrman 1999).

These links between organic carbon, bacteria (used here to encompass Bacteria and Archaea), viruses, and protists mean that the species composition and physiological state of the autotrophic community in a given system may profoundly affect the bacterial and protist communities. Whereas we recognize the importance of phytoplankton composition (e.g. diatoms vs. coccolithophores or nitrogen fixers) for biochemical cycling, potential differences in the structure and functioning of the associated microbial loops are rarely considered. Potentially, shifts in phytoplankton composition could, for example, cause modifications in DOM composition, resulting in adjustments of the bacterial community structure and degradation rates, leading to variations in the biochemical cycling of elements and organic matter and in the export efficiency of organic matter. Highly controlled experiments that would be required to study the effect of phytoplankton on the microbial loop are impossible in the field due to many potentially confounding physical and biological factors: the patch under investigation might, for example, be diluted due to mixing or advection, or the arrival of swimmers may change the food web.

Mesocosms (~200 to 15 000 l) allow for the controlled manipulation of a water body large enough to adequately host the different compartments of the microbial food web. While mesocosms are generally too small to support a food web that includes large zooplankton, they sustain microbial food webs adequately for days to weeks and have been used successfully to investigate microbial dynamics after perturbations of the environment. Some such studies have focused on the development or decline of phytoplankton blooms (e.g. Allredge et al. 1995, Engel et al. 2002) or on the interactions between different microbial loop organisms (Castberg et al. 2001, Larsen et al. 2001, Brussaard et al. 2005). Other studies have concentrated on changes in the composition of organic matter (Meon & Kirchman 2001) or on the bacterial community. Enrichment of natural seawater with nutrients often permits a growth spurt of bacteria and a subsequent increase in protists, allowing investigations of the effects of a growth and a grazing phase on the bacterial community (Lebaron et al. 1999, 2001, Schäfer et al. 2001, Troussellier et al. 2005). Other investigations have examined possible direct effects of nutrient addition (organic or inorganic) on microbial communities (Pinhassi et al. 1999, Øvreås et al. 2003), or on the competition between bacterial and phytoplankton communities (Havskum et al. 2003). Studies focusing on the post-bloom phase of phytoplankton, when colonization of decaying diatoms is accompanied by shifts in bacterial community composition and increases in abundance and hydrolytic activity of bacteria, have also been conducted (Riemann et al. 2000). Most of these investigations have studied specific aspects or processes of the microbial loop rather than trying to understand and characterize the structure and functioning of the microbial loop in its entirety. Furthermore, the effect of phytoplankton composition on the different components of the microbial loop has not previously been addressed.

We therefore set up a series of mesocosms to investigate the influence of the phytoplankton community on organic matter composition, the activity and composition of the bacterial community, the food-web structure, and associated biochemical cycling. The phytoplankton chosen, Phaeocystis globosa and the diatoms Chaetoceros decipiens and Thalassiosira pseudonana, are all bloom formers, potentially able to build up a large autotrophic biomass, but they belong to different functional groups. Diatoms require silicic acid for growth and play a significant role in transporting carbon to the deep ocean. Diatoms generate transparent exopolymer particles (TEP) and form large aggregates which sink (Passow 2002). Phaeocystis flagellates, in contrast, are small and mobile, but may form large carbon-rich colonies. Phaeocystis spp. generate abundant TEP and are known for their generally high production of mucilage, which often accumulates as foam at the water surface. Sedimentation events are possible after Phaeocystis spp. blooms, but depend on the presence of other particles in the water (Wassman et al. 1995).

The main hypothesis of this experiment was that differences in the dominating phytoplankton population would be reflected in the composition and concentration of DOM, and in the microbial community and the cycling of elements. The 3 mesocosms initially differed in their autotrophic population only. The biology and biochemistry of the evolving microbial loop community was monitored over a 20 d period. In the present paper, which gives an overview of the experiment, we present data on the distributions of chlorophyll a (chl a) and heterotrophic nanoflagellates (HNAN) and on nitrogen, phosphorus, and silica cycling. Additionally, analyses of the similarities and differences in the
development of the microbial loops in the 3 mesocosms based on all data collected (Grossart et al. 2007, this issue, Murray et al. 2007, this issue) are provided. Murray et al. (2007) relate temporal changes in the bacterial community composition to variations in organic matter composition and utilization by comparing the composition of the bacterial community structure to changes in enzymatic hydrolysis rates and organic matter composition. Grossart et al. (2007) present changes in the composition of organic matter and assembles biogeochemical budgets.

MATERIALS AND METHODS

Experimental set-up. Three rectangular, 1000 l polypropylene tanks (1.65 m depth) were filled with pre-filtered water collected from 2 m depth in the North Sea, near Helgoland, Germany. Mesocosms and filtration systems were cleaned thoroughly before use to avoid contamination with dissolved organic carbon (DOC). Mesocosms were scrubbed and wiped with alcohol, the 1.2 µm cartridge was rinsed and autoclaved, and the ultrafiltration system was acid rinsed (phosphoric acid), followed by a rinse with sodium hydrochloride. Because high wind conditions at the time of collection promoted the inflow of water from the Elbe River, the collected water was high in nitrate (≥56 µM) and relatively low in salinity (30‰). Fresh water influence was also visible in the bacterial composition (Murray et al. 2007). The water was stored in the dark for 48 h prior to the start of the experiment so that coarse particulates could settle. Half of the water was then filtered through a 1.2 µm cartridge filter. To diminish the initial microbial population and the concentrations of colloidal organic carbon, the remaining half of the water was ultra-filtered to remove all material larger than 50 kDa (Millipore, Pellicon). The filled tanks were mixed overnight by an airlift system before the experiment was started.

To the No Addition tank, nothing further was added. To the Phaeocystis tank, 1.5 l of a culture of solitary P. globosa was added to produce a cell concentration of 700 cells l−1 at Day 0. The Diatom tank was supplemented with 15 µmol l−1 of silicic acid as sodium metasilicate and inoculated with 3 l each of axenic cultures of Chaetoceros decipiens (CCMP173) and Thalassiosira pseudonana (CCMP 1013), producing an initial cell concentration of approximately 300 and 3000 cells l−1, respectively. Assuming a growth rate of 0.5 and 0.25 d−1 for the diatoms and Phaeocystis spp., respectively, these initial cell concentrations would have resulted in blooms after 8 d equivalent to particulate organic nitrogen (PON) concentrations of 10 µM. The first samples were taken 17 h after addition of the inoculi to ensure that the added phytoplankton was well mixed within the mesocosms.

Water temperature in the indoor mesocosms was always between 11.0 and 11.3°C, and each mesocosm received 250 µE m−2 s−1 light at the water surface under a 16:8 h light:dark cycle, mirroring the contemporaneous light cycle at the sampling site. The mesocosms were continuously mixed using an airlift system. The airlift system slowly moved water from the bottom of the mesocosm to the top, through a tube into which air was injected into the lower half of the tube (Egge & Aksnes 1992). Mixing by this airlift system did not promote aggregation of small particles as can be seen from the size distributions of TEP and Coomassie-stained particles (CSP) (U. Passow unpubl. data). Conductivity/temperature/depth (CTD) profiles indicated no stratification in the tanks, and pH remained between 8.0 and 8.2 in all mesocosms at all times.

The experiment was conducted for 20 d, and samples were taken every 2 d (except initially, when the interval was 3 d) from about 20 cm below the surface. Data were graphed in real time, but are referred to in whole days for ease of reading, e.g. Day 0.7 is called Day 1. Samples were processed immediately after collection.

Analysis of chlorophyll and abundance of heterotrophic flagellates. Chl a samples were collected on pre-combusted GF/F filters (Whatman), extracted in 90% acetone, and measured using a fluorometer (Turner TD 10). HNAN were collected on black filters (0.8 µm polycarbonate, Poretics, filtering 3 to 20 ml filter−1), fixed with glutaraldehyde, stained immediately with DAPI, and stored frozen for a few weeks until triplicate filters were counted via fluorescence microscopy (Kemp et al. 1993) at a magnification of 400× to 1000×. Autofluorescence was used to distinguish heterotrophic from autotrophic flagellates.

Analysis of inorganic nutrients, dissolved organic phosphorus (DOP), particulate organic phosphorus (POP), particulate organic nitrogen (PON), and biogenic silica (BSi). Samples for nitrate, nitrite, and phosphorus were filtered (GF/F, Whatman) and kept frozen until determination on a nutrient autoanalyzer (Technicon ASM II). Concentrations of soluble reactive phosphorus and silicic acid were measured manually using a spectrophotometer. Autoanalyzer and manual determinations of orthophosphate (PO43−) agreed well, and averages of both were used. DOP was calculated from TP (total phosphorus) and PO43−. TP was determined colorimetrically after persulfate oxidation (Koroleff 1983). Silicic acid concentrations were determined via molybdate blue spectrophotometry (Strickland & Parsons 1968) using a reverse-order reagent blank.

Samples for PON were collected on pre-combusted GF/F filters and then measured using an elemental
RESULTS AND DISCUSSION

Autotrophic and heterotrophic biomass and activity

The 3 mesocosms differed appreciably with respect to the build up of autotrophic standing stocks (Fig. 1a) and primary production (Grossart et al. 2007). In the No Addition tank, net growth of phytoplankton was negligible, whereas a small *Phaeocystis globosa* bloom (chl a concentrations 2 µg l−1) developed on Days 12 to 14, followed by a secondary rise at the end of the experiment (<2 µg chl a l−1) in the *Phaeocystis* tank. Overall, the No Addition tank represented a net heterotrophic system, whereas the autotrophic and heterotrophic activities were generally balanced in the *Phaeocystis* tank (Grossart et al. 2007). In the Diatom tank, a net autotrophic system developed, with chl a concentrations increasing exponentially to 55 µg l−1 at the end of the experiment (Day 20).

HNAN had mostly been removed by the pre-filtration step, but a small inoculum remained, growing to significant concentrations by Days 8 to 10 in all 3 mesocosms (Fig. 1b). The presence of phytoplankton had, however, an effect on flagellate abundance, with higher peak concentrations (16 × 10^3 cells ml−1) in both algal tanks compared to the No Addition tank (<4 × 10^3 cells ml−1). HNAN introduced grazing pressure on bacteria during the second half of the experiment in all 3
Table 1. List of parameters measured during the experiment, the method used, and the paper in which the data are discussed in detail. Parameters are categorized as for the clustering analysis

<table>
<thead>
<tr>
<th>Category/Parameter</th>
<th>Method</th>
<th>Source</th>
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<tbody>
<tr>
<td><strong>% monomer composition</strong></td>
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<tr>
<td>Dissolved free neutral carbohydrate (DFCHO) composition</td>
<td>HPLC (Mopper et al. 1992)</td>
<td>Grossart et al. (2007)</td>
</tr>
<tr>
<td>Dissolved combined neutral carbohydrate (DCCHO) composition</td>
<td>HPLC after acid hydrolysis (Mopper et al. 1992)</td>
<td>Grossart et al. (2007)</td>
</tr>
<tr>
<td>Dissolved free amino acid (DFAA) composition</td>
<td>HPLC after ortho-phthaldialdehyde derivatization (Lindroth &amp; Mopper 1979)</td>
<td>Grossart et al. (2007)</td>
</tr>
<tr>
<td>Dissolved combined amino acid (DCAA) composition</td>
<td>HPLC after ortho-phthaldialdehyde derivatization and acid hydrolysis</td>
<td>Grossart et al. (2007)</td>
</tr>
<tr>
<td>Particulate combined amino acid (PCAA) composition</td>
<td>HPLC after ortho-phthaldialdehyde derivatization and acid hydrolysis</td>
<td>Grossart et al. (2007)</td>
</tr>
<tr>
<td>Particulate combined neutral carbohydrate (PCCHO) composition</td>
<td>HPLC after acid hydrolysis</td>
<td>Grossart et al. (2007)</td>
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<tr>
<td><strong>Phylotypes</strong></td>
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<td><strong>Food web</strong></td>
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<tr>
<td>Chlorophyll a</td>
<td>Fluorometer</td>
<td>Present paper</td>
</tr>
<tr>
<td>Virus-like particles (VLP)</td>
<td>SYPER Gold counts (Chen et al. 2001)</td>
<td>U. Passow et al. (unpubl. data)</td>
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<tr>
<td>Bacterial abundance</td>
<td>DAPI counts (Porter &amp; Feig 1980)</td>
<td>Murray et al. (2007, this issue)</td>
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<tr>
<td>Heterotrophic nanoflagellates (HNAN) in 3 size classes</td>
<td>DAPI/autofluorescence counts (Kemp et al. 1993)</td>
<td>Present paper, U. Passow et al. (unpubl. data)</td>
</tr>
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<td>Autotrophic nanoflagellates (PNAN) in 3 size classes</td>
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<td>U. Passow et al. (unpubl. data)</td>
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<tr>
<td>Transparent exopolymer particles (TEP)</td>
<td>Colorimetric and microscopical determination (Passow &amp; Alldredge 1995)</td>
<td>U. Passow et al. (unpubl. data)</td>
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<tr>
<td>Coomassie-stained particles (CSP)</td>
<td>Microscopical determination (Long &amp; Azam 1996)</td>
<td>U. Passow et al. (unpubl. data)</td>
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<tr>
<td><strong>Biochemical</strong></td>
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<td>PO₄³⁻, Si(OH)₄, dissolved organic phosphorus (DOP)</td>
<td>Manually (Strickland &amp; Parsons 1968)</td>
<td>Present paper</td>
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<tr>
<td>NO₃, PO₄³⁻</td>
<td>Autoanalyzer</td>
<td>Present paper</td>
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<tr>
<td>Particulate organic phosphorus (POP)</td>
<td>Manually (Koroleff 1983)</td>
<td>Present paper</td>
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<td>CHN analyzer</td>
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<td>Particulate organic carbon (POC)</td>
<td>CHN analyzer</td>
<td>Present paper</td>
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<tr>
<td>Dissolved organic carbon (DOC)</td>
<td>High-temperature combustion</td>
<td>Grossart et al. (2007)</td>
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<tr>
<td>Dissolved monosaccharides (MCHO)</td>
<td>TPTZ-method (Myklestad et al. 1997)</td>
<td>Grossart et al. (2007)</td>
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<tr>
<td>Total dissolved carbohydrates (TCHO)</td>
<td>TPTZ-method after acid hydrolysis (Myklestad et al. 1997)</td>
<td>Grossart et al. (2007)</td>
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<td>Sum of dissolved free neutral carbohydrates (ΣDFCHO)</td>
<td>HPLC (Mopper et al. 1992)</td>
<td>Grossart et al. (2007)</td>
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<td>Sum of dissolved combined neutral carbohydrates (ΣDCCHO)</td>
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<tr>
<td>Sum of particulate combined amino acids (ΣPCAA)</td>
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mesocosms (U. Passow et al. unpubl. data). Grazing was estimated using the method of Gasol (1994), which suggested that HNAN concentrations during the second half of the experiment were regulated by their food (U. Passow et al. unpubl. data). Phytoplankton composition or quantity appeared to have little effect on average bacterial concentration, but height and timing of peak concentrations (Murray et al. 2007), as well as the food-web interactions differed between mesocosms (U. Passow et al. unpubl. data).

Cycles of silica, nitrogen, and phosphorus

Silica cycle

In the No Addition and Phaeocystis tanks, concentrations of silicic acid did not decrease from their initial values (Fig. 2a). In the Diatom tank, to which silicic acid had been added, concentrations barely changed until Day 10 and then decreased rapidly to about 10 µM at the end of the study (Fig. 2a). Simultaneously, concentrations of BSi increased over time in the Diatom tank, mirroring the drop in silicic acid concentrations (Fig. 2b) and clearly reflecting the growth of a diatom bloom. By about Day 14 of the experiment, the sum of suspended BSi and dissolved silicic acid was no longer equal to the concentration of silicic acid at the beginning of the experiment (Fig. 2b). This apparent loss of silicon from the tank coincided with the visible settling of diatoms to the bottom of the tank in aggregates too large to remain in suspension under the influence of the airlift system. Loss of particulates, presumably due to settling, was also observed in POC (Grossart et al. 2007).

Nitrogen cycle

Nitrate concentrations in the source water were very high (57.6 µM), and some nitrate was added inadvertently to the tanks with algae during the inoculation with phytoplankton. In the No Addition and Phaeocystis-
tis tanks, no significant drop in nitrate concentrations was observed, with values fluctuating around 55 ± 3 and 54 ± 3 µM, respectively, during the whole experiment (Fig. 2c). In the Diatom tank, nitrate concentrations decreased during the first 14 d of the development of the diatom bloom, but remained at ≥24 µM thereafter. Nitrite concentrations remained stable around 0.6 µM in the No Addition tank, but fluctuated more, between 0.3 and 0.7 µM in both algal mesocosms. Ammonia concentrations were not monitored, but were 2.5 µM in the source water. Overall growth limitation due to inorganic nitrogen sources was unlikely for all mesocosms.

PON fluctuated around low (17 µg l⁻¹) values in the No Addition tank (Fig. 3a), reflecting the presence of a small heterotrophic grazer community. In the Phaeocystis tank, PON accumulated to levels 3 times that of the No Addition tank, with concentrations peaking on Day 16 and remaining high thereafter (Fig. 3b). Peak PON lagged 2 d behind peak chl a concentrations of the small Phaeocystis bloom observed from Days 12 to 14. In the Diatom tank, concentrations of PON increased exponentially and, by the end of the experiment, were 25 times greater than in the No Addition tank (Fig. 3c), reflecting the exponential build up of a large diatom bloom. As nitrogen concentrations were so high that it was assumed that nitrogen was not limiting production, we focused our energies on the phosphorus cycle, and DON was not determined. The median DON concentration measured on 9 dates in February and April 2002 near the routine sampling station was 11.8 µM (range 3.1 to 75.7 µM, n = 9, written communication from M. Schütt, Institute of Biogeochemistry and Marine Chemistry, Hamburg, Germany; confirmed by data from the MUDAB database of the Bundesamt für Seeschifffahrt und Hydrographie), implying that bioavailable DON can be, but is not necessarily, present in large amounts. At this time of the year nitrogen fixation may safely be neglected. The active participation of ammonia and DON in the cycling of nitrogen during our study is suggested by the fact that nitrate plus nitrite plus PON concentrations fluctuated by about 6% over time in the No Addition and Phaeocystis tanks. In the No Addition tank, no trend was observed in any of the measured nitrogen pools, implying that net DON concentrations did not change significantly during the experiment. An estimate based on budget calculations suggests that the build up of about 2 µM PON (27 µg l⁻¹) in the Phaeocystis tank was supported to 33% by nitrate, with the remaining fraction coming from DON or ammonia. An overall large decrease in the sum of nitrate, nitrite, and PON by 22 µM (308 µg l⁻¹) in the Diatom tank (the increase of 10 µM PON was offset by a decrease of 32 µM nitrate) must have been the result of sedimentation and a net increase in DON in the Diatom tank. The PON loss due to sedimentation may be roughly estimated from the silica or the phosphorus budget and suggests that less than half of the lost nitrogen can be explained by sedimentation, while the remaining >10 µM augmented the ammonia or DON pool. Increases in the DON concentration are common during diatom blooms, especially if those are growing under high nitrogen conditions (e.g. Søndergaard et al. 2000). High concentrations of bioavailable nitrogen in all 3 mesocosms are also confirmed by the fact that PON closely tracked POC (Grossart et al. 2007), with a molar C:N ratio never above 6.5 (data not shown).

Phosphorus cycle

Concentrations of PO₄³⁻ did not decrease significantly from initial values in the No Addition tank, whereas DOP decreased linearly over the whole study period from just above 1 µM initially to undetectable concentrations on Day 20 (Fig. 4a). The heterotrophic community of the No Addition tank evidently preferred DOP over PO₄³⁻, and no PO₄³⁻ was used even when DOP concentrations were low (0.2 µM). POP remained near the detection limit in the No Addition tank (Fig. 3a), and the particulate N:P ratio (molar) was low at ≤13 until Day 10, but jumped up to an average of 183 during the next days, suggesting growth under
low phosphorus conditions after Day 10, although PO$_4^{3-}$ was available.

In the *Phaeocystis* tank, PO$_4^{3-}$ concentrations dropped from 0.9 to 0.7 μM between Days 1 and 4 and then remained constant until Day 12 (no significant decrease between Days 1 and 12), but were drawn down to 0.5 μM within 2 d and fluctuated around that value for the remainder of the study (Fig. 4b). The initial decrease in DOP was almost twice as fast as that in the No Addition tank, with values reaching 0.2 μM already by Day 10 and remaining low for the remainder of the study (Fig. 4b). The higher initial rate of DOP utilization in the *Phaeocystis* tank compared to the No Addition tank and the initial lack of a PO$_4^{3-}$ draw down during the growth phase of the *P. globosa* bloom (peak: Days 12 to 14) suggest that *P. globosa* preferred DOP over PO$_4^{3-}$. *P. globosa* is known to grow on organic phosphorus when PO$_4^{3-}$ concentrations are ≤0.4 μM (Veldhuis & Admiraal 1987, van Boekel 1991). Phosphorus-limited growth by *P. globosa* was suggested by POP values (Fig. 3b). A significant increase in POP (0.1 μM) was not observed until Day 18 in the *Phaeocystis* tank, 4 to 6 d after the *P. globosa* bloom peak. The particulate molar N:P ratio, which was 17 on Day 4, increased appreciably to above the Redfield ratio thereafter, with values of 115 when the *P. globosa* bloom peaked. Low phosphorus availability during growth of *P. globosa* may have been responsible for the scarcity of colonies in our experiment (U. Passow et al. unpubl. data). The effect of phosphorus availability on the formation of colonies of *Phaeocystis* spp. is controversial in the literature (Veldhuis & Admiraal 1987, van Boekel 1991, Riegman et al. 1992). In our experiment, the lack of substantial colony formation allowed grazing of *P. globosa* flagellates by large HNAN to become a significant factor for bloom termination (U. Passow et al. unpubl. data). The growth of a population of small autotrophic flagellates after the *P. globosa* bloom was supported by PO$_4^{3-}$. The particulate N:P ratio decreased to 30 between Days 18 and 20, also providing evidence that phosphorus was available, even if not bioavailable for *P. globosa*. Clearly future studies of *Phaeocystis* spp. need to include detailed measurements on the cycling of phosphorus to clarify some of these issues.

In the Diatom tank, PO$_4^{3-}$ concentrations fell linearly between Days 1 and 8 (Fig. 4c) and more steeply during the next days to <0.2 μM on Day 14, reflecting the growth of the diatoms. DOP dropped more steeply than PO$_4^{3-}$ during the first 8 d, then fluctuated around an average low value of 0.3 μM for the remainder of the experiment. A higher rate of initial DOP utilization in the Diatom tank compared to the No Addition tank is consistent with more vigorous bacterial growth during this initial phase (U. Passow et al. unpubl. data), which was also associated with a higher rate of DOC utilization (Grossart et al. 2007). Concentrations of POP increased exponentially in the Diatom tank, and were 25 times greater than in the No Addition tank by the end of the experiment (Fig. 3c). The particulate N:P ratios after Day 4 were always above the Redfield ratio thereafter, with values of 55, implying growth conditions under low phosphorus with rapid phosphorus turn-over during growth of the diatom bloom.

**Synthesis: comparison between the 3 mesocosms**

Differences in the start up conditions

The 3 mesocosms were intended to initially differ only with regard to their phytoplankton population and their initial silicic acid concentration, as the origi-
nal water as well as environmental conditions were the same in each of the mesocosms. Silicic acid concentrations were always >10 µM and are assumed not to have affected heterotrophic dynamics. However, *Phaeocystis globosa* could not be acquired axenically, and bacteria were added into this mesocosm along with the phytoplankton inoculum. DGGE profiles of bacteria initially present in the *Phaeocystis* tank were quite different (only ~20% similar) from those in the other 2 tanks; however, by Day 4, they were much more similar (~70%) (Murray et al. 2007), suggesting that most of the introduced bacteria vanished rapidly. In the Diatom tank, some of the introduced diatoms lysed due to the lower salinity of the mesocosm water, introducing additional DOM of a different composition than that already present (Grossart et al. 2007). This material supported a larger growth spurt of the bacterial population during the first 4 d compared with the development in the other 2 mesocosms (Murray et al. 2007). However, after Day 4, the DOM composition and the bacterial concentration of the Diatom tank returned to conditions similar to those in the other 2 mesocosms (Grossart et al. 2007, Murray et al. 2007), suggesting that the additional DOM was removed within this time span. Bacterial activity can utilize freshly released DOM within a few days (Grossart et al. 2006). Differences between the mesocosms after Days 4 to 6 are consequently interpreted to be the direct or indirect result of the growth and activities of the differing phytoplankton populations and not the transient side effects of the inoculations themselves.

Overall comparison of the mesocosms based on all parameters measured

Comparison of averages neglects temporal trends. The 1-dimensional clustering approach used here allowed a separation of temporal trends within meso-
cosms and treatment-based differences between mesocosms (Fig. 5). This analysis allowed us to test the main hypothesis that microbial communities dominated by different phytoplankton communities would differ appreciably (1) in the composition of DOM sustained in the surrounding water, (2) in the composition of the bacterial community, (3) in the food-web structure and functioning, and (4) in the biochemical cycling. The results of the 4 clustering analyses reflect these 4 parts of the hypothesis.

The 4 dendrograms that depict the degree of similarity between samples based on the 4 categories show different patterns. The dendrogram based on the percent monomer composition (Fig. 5a) shows large relative Euclidian distances (none under 20%), implying relatively large differences between samples. All initial mesocosm samples cluster together, mirroring the fact that initially the composition of DOM was similar in all mesocosms and affected by pre-filtration (Grossart et al. 2007). The lysing of diatoms is reflected in the percent monomer composition of D4 (Diatom tank Day 4), which falls outside the initial cluster. After Days 4 to 6, each mesocosm clusters in its own group, indicating that the measurable fraction of the organic matter differed appreciably between mesocosms after Day 6. This suggests that phytoplankton composition did affect, either directly or indirectly, the composition of DOM appreciably. Although monomer composition of DOM changed considerably over time, temporal changes were smaller than differences between mesocosms. The very large shift (60%) in organic matter composition of the diatom bloom on Day 20 may reflect the transition into senescence of the bloom.

The dendrogram based on DGGE phylotypes shows that both temporal and treatment-based differences in bacterial community structure were large, >25% and up to 90% between any 2 samples (Fig. 5b). Nevertheless, the data grouped into 2 large clusters, 1 representing the first phase (up to Day 8 or 10) and 1 representing the second phase (after Day 8 or 10), and 2 ‘outliers’, P1 (Phaeocystis tank Day 1) and D18 (Diatom tank Day 18). The outlier position of P1 was caused by the addition of non-axenic inoculum, whereas that of D18 cannot be explained easily with the data available. The temporal division between the 2 large clusters suggests that the phytoplankton community composition had a relatively small effect on bacterial community composition compared to the temporal shift occurring in parallel in all 3 mesocosms. The most prominent difference between the 2 phases was the appearance of HNAN exerting grazing pressure on bacteria. We, thus, hypothesize that grazing was responsible for this temporal shift in bacterial composition. DOP concentration in all 3 mesocosms remained fairly constant during this second phase. Possibly grazing introduced some steady state with respect to DOP recycling. Although differences in DGGE phylotypes between mesocosms were small compared to these temporal changes, bacterial community structure also reflected some differences between the mesocosms, as indicated by the clustering of samples from the No Addition tanks within the cluster of Phase 2. The presence of phytoplankton per se appeared to have had a larger effect on bacterial community composition than the type of phytoplankton, as bacterial community composition in the 2 tanks with algae were not separated, also confirming that flagellate abundance (e.g. grazing) rather than phytoplankton might have been responsible for the observed differences in bacterial composition. These results gained from the clustering approach used here are in accordance with the more detailed analysis of the bacterial community composition in the companion paper by Murray et al. (2007).

A different picture emerges when looking at the dendrogram based on food-web structure, where all samples from the No Addition tank, the first 10 d of the Phaeocystis tank, and many Diatom tank samples from the early phase were grouped together in a cluster (Fig. 5c), with a relatively high degree of similarity (<20% difference). This suggests that, during the first phase of the experiment, development in standing stocks and activities were similar in all mesocosms, and dominated by the start up conditions, e.g. inoculation and pre-filtration. The initial lysis event in the Diatom tank was probably responsible for the slightly different grouping of samples from Days 4 and 8. The cluster analysis furthermore implies that food-web structure in the No Addition tank changed little over time. Food webs in both algal mesocosms developed differently (different clusters) during the second phase, after Day 10, and went through large temporal changes, as represented by the large Euclidian distance between consecutive samples. These temporal changes reflect large differences in organism and particle concentrations, as well as in primary production, community respiration, and hydrolysis rates of leucine, whereas the extracellular hydrolysis rates of 6 complex polysaccharides were remarkably similar in all 3 mesocosms (Murray et al. 2007). On average, the enzymatic hydrolysis of 6 different complex polysaccharides appeared unaffected by autotrophic production, whereas average leucine hydrolysis differed significantly between mesocosms, with higher values in mesocosms with higher primary production (Murray et al. 2007).

The results of the clustering analysis of the biochemical parameters (Fig. 5d) suggest that all samples were very similar, with a relative Euclidian distance of <20%, except for the samples from the last days of the experiment (Days 16 to 20) from the Diatom tank. Although comparably similar, samples grouped into 4
clusters, reflecting different levels of biomass. The first cluster contained all samples with very low biomass, e.g. Diatom tank samples on or before Day 4, Phaeocystis tank samples on or before Day 10, and both initial and late No Addition tank samples. Biomass in the No Addition tank fluctuated during the experiment. The second cluster contained samples of slightly elevated biomass, including samples from the late Phaeocystis and intermediate samples from the Diatom tank. The third cluster includes some intermediate to late samples from the No Addition tank, emphasizing the difference between the No Addition and the 2 tanks with algae. Samples from the last days of the experiment from the Diatom tank underwent large temporal changes as the bloom grew, reflected in the high degree of dissimilarity between the individual samples of this fourth cluster. The biomass build up, which is a quantitative rather than qualitative process, dominated the clustering of the biochemical parameters more than any other single process.

The clustering analysis suggests that parts of our hypothesis were accurate in our mesocosm system, whereas other parts can be rejected. Phytoplankton composition affected DOM composition after only 6 d, and food-web structure in the 3 mesocosms diverged after 10 d. In terms of biochemical cycling, the Diatom tank digressed from the other 2 mesocosms after about 14 d, reflecting differences in biomass production. The effect of phytoplankton on bacterial community composition was unexpectedly small, with the modest, distinct development in the No Addition tank after Day 12. However, temporal changes in bacterial community composition were marked.

OVERALL SUMMARY AND CONCLUSIONS

The overarching goal of this mesocosm study was to characterize the effect of phytoplankton on the associated microbial food webs. The 3 mesocosms developed divergently in terms of autotrophic build up of organic carbon: a net heterotrophic system prevailed in 1 mesocosm, an overall balanced system with a small bloom of Phaeocystis globosa existed in the second, and a large diatom bloom formed in the third mesocosm. Considering these large quantitative and qualitative differences in autotrophs between the 3 mesocosms the differences in the structure and function of the associated microbial loops were small. Initial inspection of the very large data set led to the impression that organic matter composition and the heterotrophic grazer communities of the different mesocosms appeared not to differ significantly from each other. Using statistical pattern recognition tools, it was, however, possible to tease the differences apart and identify interesting connections between phytoplankton and the other components of the microbial loop.

The cycling of phosphorus differed appreciably as a function of phytoplankton, with DOP utilized preferentially by Phaeocystis globosa and PO$_4^{3-}$ utilized by diatoms.

After about 6 d, the organic matter composition (to the degree it was measurable, 20 to 70%) developed differently as a function of phytoplankton composition, and each mesocosm displayed its own distinct organic matter signature. However, overall DOC concentrations and dynamics were similar in all mesocosms, and were hiding the compositional differences (Grossart et al. 2007).

Turnover of freshly released material, especially amino acids appeared larger in the presence of algae compared to in the heterotrophic mesocosm (Grossart et al. 2007), leading to higher bacterial leucine uptake rates (Murray et al. 2007).

Peak concentrations of flagellates were higher in the algal mesocosms than in the No Addition tank, as food availability was higher in the algal mesocosms (U. Passow et al. unpubl. data). The quantitative difference between the autotrophic biomass in both tanks with algae was, however, not reflected in flagellate numbers. In our experiments, the presence or absence of autotrophs, which introduced organic matter into the system rather than phytoplankton composition or quantity, appeared relevant in determining flagellate numbers (U. Passow et al. unpubl. data).

After about 10 d, food-web structure developed somewhat divergently in all 3 mesocosms, with differences visible in the feeding relationships. Higher flagellate numbers compared to the No Addition tank were, for example, supported by elevated concentrations of bacteria in the Phaeocystis tank and by higher TEP concentrations in the Diatom tank (U. Passow et al. unpubl. data). High concentrations of HNAN >6 µm were only supported in the Phaeocystis tank (U. Passow et al. unpubl. data). Rapid temporal change in all mesocosms overlaid these developments.

Bacterial community composition was minimally affected by phytoplankton composition, and temporal variation was as large as differences between mesocosms, obscuring possible differences caused by different autotrophic communities. Two phases were visible in the bacterial community composition of all mesocosms. Possibly high grazing pressure during the second phase was responsible for this shift.

Although differences were observed in the hydrolysis of individual polysaccharides, which could be related to community composition, the temporal changes in patterns were similar among mesocosms, suggesting that polysaccharide turn-over was largely independent of phytoplankton species, but a function of bacterial community composition.
The microbial loop systems investigated in this study were simple compared to natural ones, where a more diverse population of protists is to be expected. Prefiltration had eliminated fragile and larger protists. Nevertheless, this study allows some insights into the effect of phytoplankton on the structure and function of the associated microbial loop systems, which could not have been investigated in situ. Our results suggest that large quantitative differences in the phytoplankton concentration (chl a) result in relatively small differences in the heterotrophic and non-living compartments. A more complex food web, as is common in nature, is likely to level out potential quantitative differences in the non-autotrophic biomass even more. Qualitative differences in phytoplankton composition were reflected in DOM composition, but not echoed in bacterial community composition. These results suggest that differences in bacterial community composition and specific polysaccharide hydrolysis rates between different natural systems might be small even if DOM composition is likely to vary. This could mean that the organic substances commonly generated in marine systems may be utilized well to the growth conditions in our mesocosms. This is supported by indications that hydrolysis rates were related to bacterial community composition. These possibilities need to be addressed in future studies. In our experimental bacterial community composition appeared to be largely a function of the grazer population, implying that top-down rather than bottom-up control determined DOM composition. A more complicated food web, where the bacterial predators are themselves under more severe grazing pressure, may result in a shift towards bottom-up control of bacterial community composition.

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Lebaron P, Servais P, Trousselier M, Courties C and 7 others

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