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Zooplankton-mediated nutrient limitation patterns in marine phytoplankton: an experimental approach with natural communities

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ABSTRACT: Zooplankton nutrient recycling has been shown to substantially affect nutrient availability for phytoplankton. However, investigations are required to determine whether zooplankton also influence nutrient limitation in marine phytoplankton communities, and whether grazing by different zooplankton groups results in different patterns of phytoplankton nutrient limitation. We performed laboratory experiments under different nutrient supply conditions on a variety of phytoplankton communities with natural densities of copepods and rotifers, and tested phytoplankton nutrient limitation in bioassays for nitrogen, phosphorus, and the combination of the two. After 5 d incubation with zooplankton, we observed a significant increase in phytoplankton biomass in the zooplankton treatments. We relate this largely to nutrient recycling effects, which are amplified through possible trophic cascade effects. In copepod treatments, the highest phytoplankton biomass was reached under Redfield and nitrogen excess nutrient supply conditions, while the highest biomass in rotifer treatments was registered under phosphorus excess conditions. In most cases, nutrient limitation assays revealed a co-limitation of phytoplankton by nitrogen and phosphorus. With increasing nitrogen supply, we observed an increase in phosphorus limitation in the copepod treatments and a decrease in nitrogen limitation in the rotifer treatments. The phytoplankton community was driven into phosphorus limitation under nitrogen excess conditions in copepod treatments. Our results indicate that natural densities of zooplankton are able to promote nitrogen and phosphorus co-limitation in phytoplankton communities.

KEY WORDS: Nutrient limitation · Marine plankton · Zooplankton-driven nutrient recycling · Stoichiometry

INTRODUCTION

Nutrient limitation in phytoplankton communities is commonly observed and may eventually lead to growth constraints (Hessen et al. 2004), as well as stoichiometric changes in phytoplankton biomass (Klausmeier et al. 2004a). While nutrient limitation in freshwater lakes is mainly associated with phosphorus (Schindler 1977, Hecky & Kilham 1988, Elser & Hassett 1994), nutrient limitation in marine environments is not focused on just one nutrient (Hecky et al. 1993). Nitrogen (N), phosphorus (P) and iron can potentially limit the primary productivity of oceans (Boyd et al. 2000, Wu et al. 2000, Sañudo-Wilhelmy et al. 2001, Brzezinski et al. 2011). However, the type of nutrient limitation is dependent on the temporal and spatial distribution of dissolved nutrients (e.g. Tyrrell 1999, Weber & Deutsch 2010), in addition to species-specific requirements of phytoplankton (Quigg et al. 2003, Lagus et al. 2004). Seasonal variations in the nutrient limitation of phytoplankton can be high in coastal areas that are affected by river runoff and
turnover events, which cause seasonally changing ratios of dissolved nutrients, notably N and P (e.g. Fisher et al. 1992, Beman et al. 2005).

In general, the stoichiometric composition of phytoplankton is dependent on the rate of carbon fixation per assimilated nutrient and can be characterised as the atomic carbon to nitrogen to phosphorus (C:N:P) ratio. The average relationship between C:N:P in seston is found to be relatively stable, ranging between 106:16:1 (Redfield 1958, Copin-Montegut & Copin-Montegut 1983) and 166:20:1 (Sterner et al. 2008). Alterations of phytoplankton C:N:P ratios can be caused by changes in various factors, such as growth rates (Klausmeier et al. 2004a), light intensity (e.g. Healey 1985, Sterner et al. 1997, Diehl 2002, Klausmeier et al. 2004b), and dissolved nutrient concentrations (Klausmeier et al. 2004a).

It has further been shown that the availability of dissolved nutrients is also affected by the nutrient recycling of zooplankton (Sterner et al. 1992, Urabe 1999). Since consumer bodies have relatively conservative stoichiometric compositions and specific nutrient requirements (Sterner et al. 1992), the ratio of excreted nutrients is altered through the nutrient demands of the consumer. Hence, the withdrawal of nutrients from zooplankton in ratios required by consumers may consequently affect the nutrient limitation of phytoplankton (Elser et al. 1988, Andersen & Hessen 1991, Urabe 1999, Elser & Hassett 1994). This concept of phytoplankton stoichiometry being affected by a feedback mechanism of zooplankton consumer grazing is termed consumer-driven nutrient recycling (CNR; Sterner 1990, Elser & Urabe 1999).

According to the nutrient demands of herbivorous consumers, recycled food should have different effects on the nutrient budget of an aquatic system. Hypothetically, since copepods have a relatively high N demand (Andersen & Hessen 1991, Gismervik 1997, Hassett et al. 1997), they should excrete less N, and consequently drive the system towards N limitation (Sterner & Hessen 1994, Walve & Larsson 1999). Conversely, consumers with low N:P ratios and/or higher growth rates (Main et al. 1997), such as daphnids, rotifers (Stemberger & Miller 1998) or appendicularians (Sommer 2003), tend to manoeuvre the system towards P limitation due to higher P demands. Such effects have been observed and described in freshwater lakes (cf. Table 1 in Elser & Urabe 1999), but are more difficult to detect in marine environments. The effects of nutrient recycling are presumed to be weaker in marine systems, due to a smaller stoichiometric mismatch between phytoplankton and consumers (Elser & Hassett 1994). Until now, only one study has attempted to investigate the effect of zooplankton on phytoplankton stoichiometry and nutrient limitation in a marine environment (Sommer et al. 2004). Sommer et al. (2004) performed mesocosm experiments in Atlantic waters off Norway to investigate the influence of density-manipulated copepod and appendicularian communities on phytoplankton nutrient limitation using bioassays. In the appendicularian treatments, nutrient limitation of phytoplankton was not detected. The authors attributed the absence of phytoplankton nutrient limitation in these treatments to the low densities of appendicularians. However, as the employed concentrations represented natural densities, the authors concluded that appendicularians would not be able to drive phytoplankton into nutrient limitation in the natural environment. In the copepod treatments, Sommer et al. (2004) found that N limitation was primarily enhanced through a trophic cascade effect. Ciliates were reduced through copepod grazing, releasing nanophytoplankton from ciliate grazing pressure, while at the same time reducing the amount of N-rich ciliate excretions. As a consequence, nanophytoplankton was affected by N limitation due to the reduced amount of ciliate excretions.

In terms of ecological modelling, 2 studies (Touratier et al. 2001, Nugraha et al. 2010) have specifically approached the question of how herbivorous zooplankton may influence N and P distributions in aquatic systems. Both studies are based on the model of Sterner (1990) and extend the understanding of the relationship between the N:P ratio of zooplankton excretions and the quality of the phytoplankton food (the phytoplankton N:P ratio). Touratier et al. (2001) showed that the type of nutrient limitation in a phytoplankton community is dependent on the zooplankton group (in their case copepods and cladocerans), and its stoichiometric requirements. The model of Nugraha et al. (2010) was specifically designed for marine systems, and predicts that global primary productivity would be significantly affected by herbivorous zooplankton. The model also shows that if zooplankton had a higher N:P ratio than phytoplankton, global primary productivity would increase in the presence of herbivorous zooplankton. The resulting low N:P re-supply ratio from zooplankton would favour the growth of P-limited nitrogen-fixing phytoplankton, which in turn would promote the growth of N-limited phytoplankton (Nugraha et al. 2010).

However, the influence of zooplankton on nutrient limitation and growth of phytoplankton in marine systems has never been tested under controlled labo-
ratory conditions. Therefore, we performed a laboratory study to investigate the short-term effects of natural zooplankton densities on nutrient limitation patterns of phytoplankton communities. We investigated whether zooplankton is able to affect the nutrient limitation of phytoplankton on an ecologically reasonable timescale of 5 d. In highly variable environments, such as the Bay of Biscay, longer periods of stable oceanographic conditions for plankton are unlikely to occur (e.g. Koutsikopoulos & LeCann 1996). In addition, we examined whether different zooplankton groups (copepods and rotifers in our case) can cause different nutrient limitation patterns in phytoplankton communities. To obtain a wide range of responses, we investigated the nutrient limitation patterns in different phytoplankton communities arising from a common natural community, and under different nutrient supply conditions.

MATERIALS AND METHODS

Experimental design

The experiment was performed in the laboratory facilities of the European Institute for Marine Studies, Plouzané, France, using 6 mesocosm tanks (350 l) at constant temperature (~18°C), and 17 h light exposure (60 µmol quanta m−2 s−1) with natural spectrum fluorescence tubes.

The tanks were filled in late February 2010 with 1 µm filtered sea water. The natural phytoplankton community of the Bay of Brest on the Atlantic coast of Brittany, France, served as initial phytoplankton community. To obtain a range of different phytoplankton communities, a different amount (10, 5, 1, 0.1, 0.01, 0.001 ml) of a Bay of Brest water sample (250 µm pre-filtered) was added to each of the 6 mesocosm tanks. The intentionally introduced higher level of environmental noise across the tanks allows the observation of broad instead of community-specific responses. Phytoplankton communities in the tanks represented different combinations of species originating from a single sample of the Bay of Brest community and were therefore treated as replicates. The tanks served as growth chambers for the phytoplankton communities, as well as incubation basins for the zooplankton experiments and nutrient bioassays. To attain similar phytoplankton biomasses before zooplankton inoculation, 4 wk of growth was allowed. Aeration of the mesocosms was provided by constant air bubbling.

To investigate nutrient limitation patterns of phytoplankton under different nutrient supply ratios, the following nutrient fertilisation ratios were selected (based on natural dissolved P concentrations): Redfield conditions (16:1 N:P ratio with addition of 8 µmol l−1 N and 0.5 µmol l−1 P) for a balanced environment, N excess conditions (48:1 N:P ratio with addition of 24 µmol l−1 N and 0.5 µmol l−1 P) for a P-depleted environment, and P excess conditions (5:1 N:P ratio with addition of 8 µmol l−1 N and 1.5 µmol l−1 P) for a N-depleted environment (Fig. 1). N was added as 1:1 molar solution of KNO3 and (NH4)2SO4 and P was added as KH2PO4 solution. For each nutrient supply condition, 3 bottles (800 ml) per mesocosm tank were filled with tank water, fertilised and incubated in the respective tanks. To ensure reliable handling of the manipulation procedure, samples for the 3 nutrient supply conditions were taken out one after the other.

Two days after the initiation of phytoplankton growth in the fertilised 800 ml bottles, copepods and rotifers were each added to 1 bottle per nutrient condition at similar concentrations (~200 µg zooplankton carbon l−1). The third bottle served as the control
without zooplankton (Fig. 1). For zooplankton treatments, the copepod *Acartia tonsa* and the rotifer *Brachionus plicatilis* were used, since both are well established experimental zooplankton species. *B. plicatilis* lives mainly in coastal waters and salt lakes and was chosen for practical culturing reasons. Rotifers were fed with a mixture of *Rhodomonas* sp. and *Dunaliea* sp. and were washed over a 63 µm gauze and put into filtered sea water before being transferred into the experimental units. Numbers were estimated by averaging 4 counts of 1 ml each. Copepods were hatched from eggs in filtered natural sea water, fed with *Rhodomonas* sp. and cultured until adult stages were reached. They were transferred into the experimental units by individual hand picking after being kept for 1 night in fresh filtered sea water without food algae. After zooplankton addition, the bottles were incubated for 5 d in the respective tanks, and subsequently sub-sampled for stoichiometric analysis and nutrient limitation bioassay (Fig. 1). Altogether, the experiments consisted of 54 experimental units with 9 factor combinations (3 nutrient conditions × 3 zooplankton treatments) per tank (n = 6).

**Sampling procedure**

To ensure the elimination of rotifers and copepods from the samples, the water from all 800 ml bottles was filtered through a 63 µm mesh. For the determination of seston particulate organic carbon (POC), nitrogen (PN), and phosphorus (PP), 50 ml (for POC and PN) and 20 ml (for PP) of the residual water was filtered onto pre-combusted acid-washed glass fibre filters (GF/C, Whatman). POC and PN were measured with a CHN-Elemental Analyzer (CE Instruments), while PP was measured via a molybdate reaction following sulphuric acid digestion. To determine the elemental composition of the zooplankton species, 2 × 10 adult *Acartia tonsa* and 2 × ~1000 *Brachionus plicatilis* were brought onto pre-combusted acid-washed glass fibre filters, and analysed accordingly.

The next step in the experimental analysis was to test the phytoplankton for potential nutrient limitation in each of the 54 experimental units in a bioassay experiment. For these bioassays, 4 × 20 ml of the filtered water were filled into small sub-sample bottles (Fig. 1). In 3 of the sub-samples, the nutrients N, P, and a combination containing N and P (NP) were added as nutrient pulses (24 µmol l⁻¹ N, 1.5 µmol l⁻¹ P). One sub-sample served as a control without nutrient addition. In all bioassays, chlorophyll fluorescence was measured with an AquaPen-C AP-C 100 (Photon Systems Instruments) fluorometer at 455 nm excitation wavelength. Integrated raw fluorescence served as proxy for the chlorophyll a content. Measurements were performed immediately after the addition of the nutrient pulse (0 h) and subsequently at 2, 24, 48 and 72 h after the nutrient pulse (±30 min).

**Calculations**

To identify cases of nutrient limitation in each bioassay, the slopes of linear regressions of chlorophyll a versus time (h) were calculated and tested for significant differences between sub-samples with nutrient addition (N, P, NP) and the control (5% level 1-sided *t*-test). The presence of significant differences were then classified either as no limitation (no differences from control), P limitation (P or P + NP different from control), N limitation (N or N + NP different from control), or co-limitation (NP or N + P + NP different from control). Following Tamminen & Andersen (2007), we used bootstrapping in our nutrient limitation classification by the random selection of slopes within factor combinations. Analyses of variance (ANOVARAs) were performed on the integrated raw fluorescence data (as proxy for chlorophyll a content) to highlight significant differences (p < 0.05) among tanks and experimental units.

**RESULTS**

Similar phytoplankton biomasses were observed in the tanks after 4 wk of growth and remained in the same order of magnitude among nutrient treatments throughout the experimental duration (n = 18; chlorophyll a values among tanks over time, Kruskal-Wallis ANOVA p = 0.344; starting chlorophyll a values among the 3 nutrient approaches over all tanks, Kruskal-Wallis ANOVA p = 0.505). The phytoplankton communities in the tanks represented typical phytoplankton communities that can be found in the Bay of Brest (e.g. Del Amo et al. 1997) and were dominated by diatoms (*Nitzschia* sp., *Chaetoceros* sp.) and dinoflagellates (*Gymnodinium* sp.).

All 54 experimental units (800 ml bottles) were successfully analysed, and the respective type of nutrient limitation in phytoplankton communities could be determined. All nutrient limitation assays were analysed; 72 measurements (2 h measurement under N excess conditions) of the total 1080 measurements could not be performed due to technical problems.
Zooplankton effects on algal biomass

We observed significant differences between zooplankton treatments averaged over all nutrient conditions (2-way ANOVA p < 0.001), expressed as enhanced phytoplankton biomass in the copepod and rotifer treatments after 5 d of incubation compared to the control treatment without zooplankton (Fig. 2a). The effect of the zooplankton was dependent on the applied nutrient conditions (2-way ANOVA zoo × nutrient interaction p < 0.001). In the control as well as in the rotifer treatments, the highest phytoplankton biomass was found under P excess conditions. In the copepod treatments, the Redfield nutrient conditions promoted the highest phytoplankton biomass.

Nutrient limitation assay results

In the bioassays, the highest growth rates (indicated by the slopes in the nutrient limitation assays) were reached under P excess conditions, and the lowest growth rates were reached under Redfield conditions (Fig. 3; same pattern for each zooplankton treatment, not shown separately). A significant increase in total phytoplankton abundance related to the addition of nutrients was determined in over 75% of all cases. The biomass of phytoplankton communities increased significantly in 76% of the control treatments, in 80% of the copepod treatments, and in 87% of the applied nutrient conditions (2-way ANOVA zoo × nutrient interaction p < 0.001). In the control as well as in the rotifer treatments, the highest phytoplankton biomass was found under P excess conditions. In the copepod treatments, the Redfield nutrient conditions promoted the highest phytoplankton biomass.
The largest increase in phytoplankton biomass occurred in the zooplankton treatments as a result of combined NP addition, indicating a prevalent NP co-limitation. Under Redfield conditions, nutrient addition did not increase the phytoplankton biomass in most cases (no limitation in 66% of cases in control, 38% in copepod and 61% in rotifer treatments; Table 2). Under N excess conditions, most cases in the control treatment were also classified as not being nutrient limited, closely followed by those classified as P limited. Under P excess conditions, most cases in the control treatment were also classified as not being nutrient limited, closely followed by those classified as NP co-limited (38%) and P limited (36%). In the copepod treatments, phytoplankton was classified as being NP co-limited under P excess (46%) and P limited under N excess conditions (37%). In the rotifer treatments, phytoplankton was classified as NP co-limited under P excess and N excess conditions (45% and 47%, respectively; Table 2).

### Phytoplankton stoichiometry

In our experiments, mean C:N ratios of the phytoplankton biomass (control treatment without zooplankton) ranged around 4.3 (average ratio of column 1 in Table 3). These C:N ratios were clearly below the mean seston stoichiometry in marine and freshwater environments described by Sterner et al. (2008; ratio 8.3) and Redfield (1958; ratio 6.63). These results indicate N excess conditions as prevalent under all initial nutrient conditions (Redfield conditions as well as N excess and P excess fertilisation). Phytoplankton C:N ratios in the copepod treatments (average 5.6; Table 3) were slightly higher (statistically not significant) than in the control treatments and the biomass C:N ratio of the copepods (4.5; Table 4). This discrepancy between the C:N ratios may indicate that copepods draw an high amount of N from the system into their bodies, resulting in higher phytoplankton C:N ratios. In the rotifer treatments, the mismatch between the phytoplankton C:N ratio (average 5.6) and the rotifer C:N

<table>
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<tr>
<th>N:P supply condition</th>
<th>Control</th>
<th>Copepods</th>
<th>Rotifers</th>
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<td></td>
<td>C:N</td>
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<td>5:1</td>
<td>4.1</td>
<td>90.8</td>
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<td>(±1.0)</td>
<td>(±32.3)</td>
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<tr>
<td>16:1</td>
<td>3.6</td>
<td>94.6</td>
<td>31.2</td>
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<td>48:1</td>
<td>5.2</td>
<td>93.4</td>
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<td>(±2.2)</td>
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an = 5
Alternatively, an effect of compensatory growth could be related to bacterioplankton. If rotifers focus on bacteria as food, phytoplankton could increase due to reduced grazing pressure, nutrient recycling of P-rich bacterial food and less competition for nutrients. Similar to the rotifer treatments, the copepod treatments showed an increase in phytoplankton biomass compared to the control treatments. Copepod grazing on bacterioplankton and ciliates, which were as equally low in abundance as in the rotifer treatments, may have had a minor compensatory effect on the total phytoplankton abundance in our experiments. A more probable link in trophic cascades could involve heterotrophic dinoflagellates, which have been found to be high quality food for copepods (Chen & Liu 2011). Since dinoflagellates were abundant in our phytoplankton communities, their selective removal by copepods could release diatoms from grazing pressure. However, since different effects in different nutrient supply conditions were observed in our experiments (Fig. 2), mechanisms other than trophic cascades must have played a role in stimulating phytoplankton growth.

Another process that may have promoted phytoplankton growth in the zooplankton treatments is the supply of trace metals through zooplankton excretions. Although the input of trace metals in coastal areas might be high, iron was found to limit phytoplankton growth even in nutrient-rich upwelling regions (Fe up to 3 nmol l\(^{-1}\), Hutchins & Bruland 1998). It is known that zooplankton release trace metals (Fowler 1977, Masuzawa et al. 1988) that stimulate phytoplankton growth (Coale et al. 1996, Boyd et al. 2000). For example, Sarthou et al. (2008) showed that copepod grazing has a significant influence on iron recycling, and hence its residence time in the water, and in turn, on phytoplankton productivity. However, it is doubtful that concentrations up to 100 nmol l\(^{-1}\) total dissolved iron, as in the Bay of Brest (Laes et al. 2005), are still limiting primary productivity. Nevertheless, trace metals also appear to influence the uptake and assimilation of macronutrients in phytoplankton (Wu et al. 2000, Wang & Dei 2001, Wang 2002). Therefore, the release of trace metals other than iron in the zooplankton treatments through nutrient recycling could have supported phytoplankton biomass growth, by promoting the assimilation of the required macronutrients N and P. As phytoplankton biomass almost doubled under some nutrient conditions in the zooplankton treatments, an increased nutrient use efficiency is possible since under zooplankton grazing only a fraction of digested nutrients is excreted and made available for the build-up of new phytoplankton biomass.

**DISCUSSION**

In our experiments, we observed, on average, positive effects of zooplankton presence on total phytoplankton abundances compared to the control treatments (Fig. 2). In general, higher phytoplankton biomass may result from nutrient recycling or trophic cascade effects of selective grazing and compensatory growth. Compensatory growth has been observed in freshwater lakes with low zooplankton densities, where grazing caused a significant increase in the biomass of grazing resistant algae (e.g. Sommer et al. 2001). Since we were not analysing the grazing efficiency on different phytoplankton species, we cannot verify this hypothesis. A trophic cascade effect was also considered to be responsible for higher phytoplankton abundances in a mesocosm study with North Atlantic phytoplankton (Sommer et al. 2004). Since copepods fed on ciliates and released nanophytoplankton from microzooplankton grazing pressure, total phytoplankton increased in biomass. This mechanism cannot be excluded for our experiments, since large ciliates (>100 µm with abundances <1 ml\(^{-1}\)) were occasionally present, but may not apply for the rotifer treatments. Although rotifers may feed on small ciliates (<60 µm, Gilbert & Jack 1993), the upper natural limit of their food size is determined by body size, and therefore would not include ciliates larger than 100 µm. In addition, the abundance of rotifers in our experiments greatly exceeded that of ciliates (by a factor of 150). The release of phytoplankton from the relatively low grazing pressure of ciliates through rotifers would thus not suggest a considerable effect on the phytoplankton biomass. Therefore, we exclude a trophic cascade effect involving ciliates in the rotifer treatments. Alternatively, an effect of compensatory growth could
The most prominent result of our experiments was that highest total phytoplankton abundances were observed in zooplankton treatments and differed among nutrient supply conditions. This can only be explained if consumer-driven nutrient recycling comes into consideration as a driving factor for the different total phytoplankton abundances, since effects of trophic cascades or micronutrient release should be comparable under all nutrient conditions. The results indicate better phytoplankton fertilisation by nutrient recycling under N-enriched conditions in the copepod treatments, and under P-enriched conditions in the rotifer treatments (Fig. 2). Differences become apparent when comparing N excess with P excess conditions: phytoplankton biomasses in the copepod treatments significantly exceeded those in the rotifer treatments under N excess (Fig. 2), and vice versa under P excess. These results can only be explained by differences in consumer-driven nutrient recycling, with respect to growth rate differences of the 2 zooplankton groups. Since copepods have a lower growth rate compared to rotifers (Allan 1976), they should require less P and excrete more P. Thus, under N-rich conditions, P scarcity could be compensated by the P-rich excretions of copepods, consequently stimulating phytoplankton growth and biomass accumulation. P excretion of a copepod-dominated zooplankton community has been estimated to account for 15 to 20% of the primary production requirements in the north-western Mediterranean Sea, with seasonal peaks up to 90% (Gaudy et al. 2003). Increased P release by herbivorous zooplankton was also suggested as a reason for increased primary productivity by model results published by Nugraha et al. (2010). Interestingly, nutrient limitation assays show P limitation of phytoplankton under N excess conditions (Table 2), which is not evident by the stoichiometry of 103:16:1 (Table 3) and may be related to higher growth rates under N excess than under Redfield conditions (Fig. 3). In contrast to the copepod treatments, phytoplankton in the rotifer treatments reached the highest biomass and growth under P excess conditions at 90% (Gaudy et al. 2003). Rotifers have higher growth rates than copepods (Allan 1976) and an accordingly higher P demand. Therefore, rotifer excretions contain low P concentrations (high N:P ratio) and thus do not stimulate phytoplankton growth in systems with P scarcity (N excess) but do it under N scarcity. As observed in our experiments, the relatively higher N concentrations in rotifer excretions seem to have stimulated phytoplankton growth and biomass accumulation under P excess conditions (Fig. 2d).

Under P excess conditions, rotifer biomass may also have influenced nutrient recycling. Rothhaupt (1995) found that Brachionus rubens reached highest biomass under non P-limited food conditions, while growth rates declined with P-limited food. In our experiments, there is no indication of severe nutrient limitation from phytoplankton N:P stoichiometry (Table 3). However, since we did not monitor the rotifer biomass after the start of the experiments, it is not possible to estimate the excess of P supply through higher rotifer biomass or the amount of P withdrawal through higher growth rates. Nevertheless, we observed that P availability did play a significant role in our experiments, since the responses of phytoplankton in the nutrient limitation assays were highest under P excess conditions (Fig. 3; same pattern for each zooplankton treatment, not shown).

**Nutrient limitation patterns**

The results of nutrient limitation bioassays indicate NP co-limitation of phytoplankton in the zooplankton treatments for all conditions other than Redfield (Tables 1 & 2). Averaged over all nutrient conditions, the control treatments showed no clear dominance of any limitation type; all frequencies ranged between 20% and 30% (Table 1). This indicates that zooplankton in general increased the phytoplankton NP co-limitation and that the macronutrients N and P had a synergistic effect on the phytoplankton communities. Nutrient limitation patterns for marine phytoplankton are known to be heterogeneous and may change over time (e.g. Karl et al. 2001b), and phytoplankton appears to be less affected by nutrient limitation in marine than in freshwater environments (Hecky & Kilham 1988). In open oceans, phytoplankton growth is predominantly P limited (Karl 1999, Wu et al. 2000, Hannides et al. 2009, Weber & Deutsch 2010), whereas in coastal areas, it is mainly N limited (Ryther & Dunstan 1971, Beman et al. 2005) or subject to seasonally changing nutrient limitation (Delmas et al. 1983, Fisher et al. 1992, Del Amo et al. 1997). However, metadata analyses have revealed combined NP co-limitation as the most abundant nutrient limitation type in both terrestrial and aquatic environments (Elser et al. 1990, 2007, Harpole et al. 2011). These studies imply that combined NP addition has the highest synergistic effects on phytoplankton growth, which coincides with our findings and seems especially promoted by zooplankton.

Besides the dominance of NP co-limitation in our study, the data showed trends of increasing P limita-
tion in the copepod treatments and decreasing N limitation in the rotifer treatments with increasing N supply ratio (Table 2). Single nutrient limitation was only found in 1 case (P limitation in copepod treatments under N excess; Table 2). The nutrient limitation patterns of phytoplankton influenced by zooplankton nutrient recycling are dependent on zooplankton densities and incubation time. Compared to other mesocosm experiments (Sommer et al. 2001, 2004), the zooplankton densities employed in our experiment fall within the lower range. Higher zooplankton densities would certainly increase the nutrient recycling effect, and force phytoplankton into single nutrient limitation more quickly.

Concerning the incubation time, the increased phytoplankton biomass in the zooplankton treatments indicates a significant influence of zooplankton during the 5 d incubation. In coastal pelagic environments such as the Bay of Biscay, high temporal and spatial small-scale variability of plankton is commonly observed (e.g. Koutsikopoulos & LeCann 1996). Our applied incubation time of 5 d without disturbance is already an optimistic value for the investigation of ecologically reasonable short-term effects of zooplankton on phytoplankton growth.

Another reason for the few cases of single nutrient limitation of phytoplankton in the presence of zooplankton may be related to the similar phytoplankton stoichiometry under all applied nutrient conditions. Phytoplankton biomass N:P ratios showed a conservative relationship compared to the initial N:P supply conditions (Table 3). Despite the range of fertilisation from 5:1 to 48:1 dissolved N:P, phytoplankton biomass N:P ratios did not reflect these nutrient conditions. The stoichiometry of our phytoplankton communities differed from the classical Redfield ratio, and were more similar to the findings of Sterner et al. (2008; our Table 3), who observed an average N:P ratio of 20:1 in particulate matter from a variety of aquatic environments. In terms of nutrient limitation, these stoichiometric ratios do not indicate severe single nutrient limitation, fitting the overall NP co-limitation pattern of our bioassays, and only tend towards weak P limitation of the phytoplankton community. Studies investigating seston C:N:P ratios have indicated that nutrient limitation seems less pronounced, and biomass carbon to nutrient stoichiometry less variable in marine than in freshwater seston (Hecky et al. 1993, Sterner et al. 2008). The biomass N:P ratios of marine phytoplankton cultures can range from <3:1 to >45:1 under extremely nutrient limiting conditions only, while natural marine particulate matter N:P ratios seem to be far less variable (Geider & La Roche 2002). According to Geider & La Roche (2002) and Guildford & Hecky (2000), critical ratios of dissolved N and P for nutrient limitation in phytoplankton growth are found below 10 and above 22 N:P. Model predictions indicate that, depending on ecological conditions, variations in phytoplankton biomass N:P ratios in a range of 8 to 45 may be interpreted as optimal growth conditions (Klausmeier et al. 2004b). Our results seem to support the predicted modelled ranges, and show that marine phytoplankton seem to be mainly NP co-limited in ranges between 5 and 48 N:P.

Besides dissolved nutrient availability, factors that affect the stoichiometric N:P composition of marine phytoplankton include growth rates (Klausmeier et al. 2004a) and genetic disposition (Quigg et al. 2003). Highly taxon-specific biochemical requirements are reflected in the N:P ratios of phytoplankton biomass (Rhee & Gotham 1980, Hecky & Kilham 1988, Geider & La Roche 2002, Quigg et al. 2003, Lagus et al. 2004), and influence the dissolved nutrient distribution in the oceans (Weber & Deutsch 2010). Experiments have shown that the phytoplankton N:P ratio decreases with increasing growth rate (Goldman et al. 1979). Models suggest that under high growth rates, phytoplankton assimilate nutrients in the ratio they require, while changes in the phytoplankton N:P ratio may only occur after a threshold in dissolved N and P availability is exceeded (Klausmeier et al. 2004a).

Another possible way in which stoichiometric ratios and nutrient limitation patterns of phytoplankton might be influenced concerns bacterioplankton. By competing for dissolved nutrients and changing the relative availability of nutrients, bacteria can promote nutrient limitation of autotrophs (Cherif & Loreau 2009). Under certain circumstances, the influence of bacteria can cause changes in phytoplankton stoichiometry. Danger et al. (2007) have shown that in high N:P growth media (>95:1), bacteria contribute to high C:N ratios in phytoplankton biomass and promote P limitation in phytoplankton growth. Since our plankton communities also contained the natural bacterial populations, their potential influences on the observed limitation patterns cannot be excluded. Since we neither applied very high N:P conditions nor used high zooplankton densities, the obtained results should include only the natural and not an artificially promoted influence of bacteria on the phytoplankton community. The more or less homogeneous stoichiometric ratios of the phytoplankton communities over all applied nutrient conditions (with or without zooplankton) may suggest that a process smoothened potential stoichiometric differ-
ences between the different N:P conditions (Table 3). This process may be related to bacterioplankton nutrient competition and nutrient recycling.

We conclude that zooplankton exerts most influence on the nutrient limitation patterns in our experiment by nutrient recycling, leading to NP co-limitation in phytoplankton communities. Trophic cascade effects including bacteria and dinoflagellates could have amplified the observed differences in total phytoplankton biomass between treatments. Nutrient recycling by bacterioplankton and the genetic disposition of phytoplankton may be responsible for the homogeneous phytoplankton stoichiometry over all applied nutrient conditions.

Implications

Our findings suggest a strong impact of herbivorous zooplankton on phytoplankton productivity and biomass even at low zooplankton densities. In relatively nutrient-rich environments, such as our mesocosm experiments or in coastal areas (Elser et al. 1990, 2007), the nutrient recycling of zooplankton seems to lead to nutrient co-limitation patterns. However, this might not be valid in extreme nutrient-poor marine regions, such as subtropical gyres, which occupy approximately 40% of the earth’s surface (Polovina et al. 2008). In such ecosystems, the near-surface concentrations of below 100 nmol for both inorganic N and P in the euphotic zones (Karl 1999, Karl et al. 2001a) indicate that phytoplankton growth is severely nutrient limited (e.g. Wu et al. 2000, Karl et al. 2001b, Moutin et al. 2005), with zooplankton having a measurable influence on nutrient availability (e.g. Corner & Davies 1971, LeBorgne 1982, Steinberg et al. 2002, Hannides et al. 2009). Therefore, in nutrient-poor marine regions, zooplankton might drive phytoplankton communities into single nutrient limitation, rather than nutrient co-limitation. For example, Hannides et al. (2009) showed that the active export of P from the euphotic zone by excretions of migrant zooplankton has the same magnitude as the passive export of P through the settlement of particles in the North Pacific Gyre. The same effect of P removal appears to hold for fish: their high demands for P can influence nutrient dynamics even in nutrient-rich regions (Hjerne & Hansson 2002) and may influence nutrient limitation patterns for phytoplankton through storage of P, mostly in fish bones, and low P recycling (Czamanski et al. 2011). Thus, active P removal from other trophic levels would certainly aggravate P scarcity and limitation of phytoplankton communities, and influence primary production. To begin with, further experimental investigations should include the effects of zooplankton on nutrient limitation patterns in nutrient-poor environments to obtain more insights into the influence of zooplankton nutrient recycling on phytoplankton communities.

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