Nutrient competition experiments with periphyton from the Baltic Sea

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ABSTRACT. Nutrient competition experiments were performed with periphytic microalgae from the Kiel Fjord, western Baltic Sea. Diatoms were dominant competitors at high Si:N ratios (under N-limited conditions) and high Si:P ratios (under P-limited conditions). Under low silicate supply, cyanobacteria became dominant at low N:P ratios and chlorophytes at high ones. Changes in light intensity influenced the outcome of competition at the species level but not at the level of higher taxa. The nitrogen source (ammonium or nitrate) had no impact on the outcome of competition.

KEY WORDS: Periphyton, Microphytobenthos, Competition, Nutrients

INTRODUCTION

Tilman's (1982) competition theory is one of the best tested theories explaining dominance and replacement of species by biotic interactions between populations. Its most prominent prediction is the resource ratio hypothesis, according to which the ratio of the limiting resources determines the taxonomic outcome of competition. So far it has been most intensively applied to phytoplankton communities, both in culture experiments (Sommer 1983, 1986, 1994a, Kilham 1986, Tilman 1977) and in situ (Riegman 1991, Makulla & Sommer 1993, Sommer 1993, Sommer et al. 1993). Similar attempts with benthic microalgae have been rare. An exception is the study of Fairchild et al. (1985) during which clay pots filled with nutrient enriched agar had been incubated for several weeks in an oligotrophic lake in Alaska. Here, I report on laboratory experiments with benthic microalgae (including filamentous forms) from the Kiel Fjord, western Baltic Sea. The main purpose of my experiments was to investigate whether the major, gross taxonomic findings of competition research with phytoplankton would also apply to periphyton. The working hypotheses to be tested were:

1. High Si:N or Si:P ratios select for diatoms.
2. Low N:P ratios at low Si supply select for cyanobacteria.
3. High N:P ratios at low Si supply select for green algae.
Additionally, the influence of the nitrogen source (nitrate vs ammonium) and of light intensity was tested.

The study was not designed to mimic the physical conditions under which competition takes place in situ. Factors such as substratum roughness, exposure to wave action, water current velocity, and presence of grazers certainly influence the outcome of competition and must be considered as this line of research continues to be developed. However, for the purpose of starting experimental competition research with periphyton it was assumed, similarly as for plankton, that there are robust physiological differences between major taxonomic groups which permit the detection of basic patterns in the taxonomic outcome of nutrient competition even under extremely simplified laboratory conditions.

MATERIALS AND METHODS

Periphyton was obtained from the dock in front of the Institute of Marine Research, Kiel, Germany. It was
scraped off from hard surfaces just below the water line and suspended in filtered water from the sampling site. Then the algal suspension was sieved through a 250 μm screen in order to exclude larger filamentous algae and animals. The filtered suspension was equally divided between the different cultures and subjected to the experimental conditions for 6 to 9 wk.

**Agar-plate experiments.** The agar-plate experiments consisted of ca 4 mm thick 2% agar layers covered by a 2 mm water layer in petri dishes. The agar was prepared in filtered in situ water which was enriched by nutrients as shown in Table 1, plus f/2 medium (Guillard & Ryther 1962) which contains a mixture of trace elements. Nutrient concentrations were designed to provide one set of cultures with a balanced N:P ratio (15:1 by atoms), one set with N strongly limiting relative to P (N:P = 1.55:1), and one set with P strongly limiting relative to N (N:P = 145:1). The nitrogen enrichment in the P-limited and in the N:P-balanced media consisted of equimolar amounts of nitrate and ammonium, while there were 2 media in the N-limited set, one with nitrate and the other with ammonium enrichment. Silicate concentrations were 1102.5, 112.5, and 2.5 μM, respectively. The experimental design was fully factorial, each N:P ratio combined with each Si concentration. In addition, the thirteenth medium was not N-enriched at all in order to select for nitrogen-fixing cyanobacteria.

Light was supplied by fluorescent tubes. To balance the low emission of white tubes (Osram 'Biolux') in the red and blue spectral ranges, 1 combined them with purple ones (Osram 'Fluora'). The light intensities at the surfaces of the petri dishes were 100, 35, and 5 μE m⁻² s⁻¹. All 3 light intensities were used for the first series of experiments (begun on 12 April 1995). For the second series (begun on 15 May 1995) the lowest light intensity was omitted. The light-dark periodicity was 14:10 h, and the experimental temperature was 18°C. Each light intensity was combined with each nutrient treatment, each combination was duplicated within both series. This means there were 4 replicates for all treatments with the medium and high light intensities and 2 replicates for all treatments with the low light intensity.

It was necessary to test the extent to which the nutrient supply ratios could be distorted relative to the concentration ratios due to differential release of the different ions from the agar. For that purpose 300 ml agar blocks containing the first 5 media were put into 200 ml sterile filtered seawater without nutrient enrichment. The water was replaced by fresh water at logarithmically increasing intervals (1, 2, 4, 8, 16 d). At each replacement the nutrient concentrations in the water were measured according to standard oceanographic methods (Strickland & Parsons 1968). Release rates were calculated by dividing the concentration increase in the water by the interval length.

Two times per week, 5 randomly selected samples of 4 mm² were taken from the plates, fixed by Lugol's iodine and placed into hot (>80°C) seawater to dissolve the agar. The mixed sample was treated like a plankton sample and counted under an inverted microscope (Utermöhl 1958). Cell volumes were obtained from linear measurements of at least 20 individuals per species and approximation to the nearest standard geometric figure. The competitive success of a species was expressed by its contribution to total biomass ('relative biomass', \( p_r = B/B_{total} \)) at the end of the experiments.

**Tissue-culture-plate experiments.** The tissue culture plates consisted of transparent polystyrene. Each plate comprised 12 chambers. Each chamber was filled initially with 1 ml of the microalgal suspension mixed with 2 ml of one of the 12 experimental media (Table 2). There were 4 levels of silicate (2.5, 5, 20, 80 μM) combined with 3 N:P ratios (P-limited: 45:1, balanced: 15:1, N-limited: 5:1). Nitrogen was supplied as NH₄NO₃. Three parallel plates were started on 16 August 1995. The light-dark periodicity was 14:10 h, and the light intensity was 100 μE m⁻² s⁻¹. Once per week 2 ml of the supernatant medium was replaced by fresh medium.

The algal film on the bottom of the chambers was inspected directly under an inverted microscope twice per week. This gave a biased view of the algal commu-

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**Table 1. Final nutrient concentration (enrichment plus original seawater) in the medium of the agar-plate experiments. Values in μM**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Silicate</th>
<th>Phosphate</th>
<th>Nitrate</th>
<th>Ammonium</th>
<th>N:P</th>
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<tbody>
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<td>1</td>
<td>1102.5</td>
<td>36.14</td>
<td>271.4</td>
<td>270.7</td>
<td>15:1</td>
</tr>
<tr>
<td>2</td>
<td>1102.5</td>
<td>3.74</td>
<td>274.1</td>
<td>270.7</td>
<td>145:1</td>
</tr>
<tr>
<td>3</td>
<td>1102.5</td>
<td>36.14</td>
<td>1.4</td>
<td>54.7</td>
<td>1.55:1</td>
</tr>
<tr>
<td>4</td>
<td>1102.5</td>
<td>36.14</td>
<td>55.4</td>
<td>0.7</td>
<td>1.52:1</td>
</tr>
<tr>
<td>5</td>
<td>112.5</td>
<td>36.14</td>
<td>271.4</td>
<td>270.7</td>
<td>15:1</td>
</tr>
<tr>
<td>6</td>
<td>112.5</td>
<td>3.74</td>
<td>271.4</td>
<td>270.7</td>
<td>145:1</td>
</tr>
<tr>
<td>7</td>
<td>112.5</td>
<td>36.14</td>
<td>1.4</td>
<td>54.7</td>
<td>1.55:1</td>
</tr>
<tr>
<td>8</td>
<td>112.5</td>
<td>36.14</td>
<td>55.4</td>
<td>0.7</td>
<td>1.55:1</td>
</tr>
<tr>
<td>9</td>
<td>2.5</td>
<td>36.14</td>
<td>271.4</td>
<td>270.7</td>
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<td>2.5</td>
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<td>274.1</td>
<td>270.7</td>
<td>145:1</td>
</tr>
<tr>
<td>11</td>
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<td>1.4</td>
<td>54.7</td>
<td>1.55:1</td>
</tr>
<tr>
<td>12</td>
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<td>55.4</td>
<td>0.7</td>
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<tr>
<td>13</td>
<td>2.5</td>
<td>36.14</td>
<td>1.4</td>
<td>0.7</td>
<td>0.06:1</td>
</tr>
</tbody>
</table>
Table 2. Nutrient concentrations (µM) in the media used for the tissue-culture-plate experiments (note that N was supplied as NH₄NO₃, which implies 2 atoms of N per molecule).

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Silicate</th>
<th>Phosphate</th>
<th>Nitrogen</th>
<th>N:P ratio</th>
</tr>
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<tr>
<td>1</td>
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<td>2</td>
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<tr>
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<td>20</td>
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<tr>
<td>5</td>
<td>80</td>
<td>2</td>
<td>45</td>
<td>45:1</td>
</tr>
<tr>
<td>6</td>
<td>20</td>
<td>2</td>
<td>45</td>
<td>45:1</td>
</tr>
<tr>
<td>7</td>
<td>5</td>
<td>2</td>
<td>45</td>
<td>45:1</td>
</tr>
<tr>
<td>8</td>
<td>2.5</td>
<td>2</td>
<td>45</td>
<td>45:1</td>
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<tr>
<td>9</td>
<td>80</td>
<td>6</td>
<td>15</td>
<td>5:1</td>
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<td>10</td>
<td>20</td>
<td>6</td>
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<td>11</td>
<td>5</td>
<td>6</td>
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<td>5:1</td>
</tr>
<tr>
<td>12</td>
<td>2.5</td>
<td>6</td>
<td>15</td>
<td>5:1</td>
</tr>
</tbody>
</table>

nity because filamentous algae and their epiphytes extended vertically into the medium. Therefore, the entire content of the chambers was scraped out by a narrow scoop and treated like a plankton sample at the termination of the experiments.

RESULTS

Agar-plate experiments

The release rates (R, in µM d⁻¹) of nutrients from the agar blocks declined rapidly with time. Their change in time could be fitted to the model \( R = at^b \) by regression analysis, where \( t \) was defined as the midpoint of the incubation intervals (in d). The exponents of the regression \( (b) \) were around -0.5 (Table 3). Differences between the exponents of the individual regressions were insignificant. Therefore, the ratio of nutrient release rates could be calculated from the parameter \( a \) of the regressions. The ratios of the release rates were similar to the concentration ratios in the medium (Table 4). Therefore, the latter were used for further analysis of data.

Species composition stabilized after 3 to 4 wk. A regression analysis of the relative biomass of each species against time showed that the slope was no longer significant. This was taken as an indication that competitive equilibrium had been attained. The species composition at the end of the experiments (Figs. 1 to 3) showed good replication. Only in 2 treatments (high light, low Si, N:P balanced; and high light, low Si, P-limited) was there a difference in one of the coexisting species under similar nutrient/light conditions but different starting times of the experiments. However, the species replacing each other (Cladophora pygmaea Reinke and Rhizoclonium riparium Harvey) belonged to the same family (Cladophoraceae). In most other cases, the differences between the \( p_i \) values of individual species were <0.2 between replicates.

Distinctive competitive optima were found for a number of species:

Cyanobacteria: Microcoleus chthonoplastes Thuret, a known nitrogen-fixer (Paerl 1988), was the dominant competitor at high and medium light intensities, low Si levels and low N:P ratios. At low light intensities under these nutrient conditions the cyanobacteria consisted of Lyngbya gracilis Rabenhorst.

Chlorophyta: Cladophora pygmaea or Rhizoclonium riparium was dominant at high light intensities, low Si levels and high N:P ratios. At low light intensities under these nutrient conditions the cyanobacteria consisted of Lyngbya gracilis Rabenhorst.

Phaeophyta: The brown algae were only represented by Ectocarpus siliculosus Lyng. which contributed substantially to algal biomass at medium light intensities, low and medium Si levels and low N:P ratios.
Fig 1. Final species composition of the periphyton biomass in the agar-plate experiments at 100 μE m⁻² s⁻¹. The first 2 bars in each block refer to the experiments started on 12 April; the third and fourth bar refer to the experiments started on 15 May. Achn: Achnanthes brevipes; Amph: Amphipleura rutjlan; Clad: Cladophora pygmaea; Melo: Melosira nummuloides; Micr: Microcoleus chthonoplastes; Rhiz: Rhizoclonium riparium; Stau: Stauroneis constricta

Diatoms: Melosira nummuloides Ag. was the only centric species of importance. It had its competitive optimum at medium light intensities, high Si levels and a balanced N:P ratio. Achnanthes brevipes Ag. was dominant at low light intensities, high Si levels and high N:P ratios. Stauroneis constricta Cleve was a dominant species at all light intensities, high Si levels and low N:P ratios. Amphipleura rutjlan Cleve was most important at medium light intensities, medium Si levels and high N:P ratios. Synedra (= Fragilaria) tabulata Kütz. also preferred medium light intensities and high N:P ratios, but high Si levels. An unidentified, small (<8 μm length) Navicula sp. had its optimum at low light intensities, medium Si levels and low N:P ratios.

Fig 2. Final species composition of the periphyton biomass in the agar-plate experiments at 35 μE m⁻² s⁻¹. The first 2 bars in each block refer to the experiments started on 12 April; the third and fourth bar refer to the experiments started on 15 May. Achn: Achnanthes brevipes; Amph: Amphipleura rutjlan; Ecto: Ectocarpus siliculosus; Melo: Melosira nummuloides; Micr: Microcoleus chthonoplastes; Stau: Stauroneis constricta; Syne: Synedra tabulata; Ulot: Ulothrix impexa

The working hypotheses stated in the introduction were tested by a multiple regression analysis with stepwise variable selection. The dependent variable was the arcsine square-root transformed relative biomass (arcsin√p) of higher taxa (diatoms, chlorophyta, cyanobacteria). The arcsine square-root transformation was necessary to obtain a normal distribution of the residuals. The independent variables were the log₁₀ transformed values of the light intensity (I) and of the stoichiometric nutrient ratios, N:P, NO₃:NH₄ and Si(N or P). The latter ratio needs some explanation: nutrient ratios are only meaningful predictors if both nutrients are limiting (Tilman 1982). In order to make Si:N and Si:P ratios numerically comparable, the molar
Sornmer. Penphyton nutrient competition experiments

AGAR, LOW LIGHT

high Si medium Si low Si

Cyanobacteria: arcsin√p
(Si:N or P) = 0.89 - 0.15 log₁₀
r² = 0.69; p < 0.0001

Chlorophyta: arcsin√p
(Si:N or P) = 0.14 - 0.21 log₁₀
r² = 0.66; p < 0.0001

The conclusions are that the proportion of diatoms is positively related to Si:N or Si:P and to N:P ratios. Cyanobacteria are negatively influenced by high Si:N or Si:P ratios and by high N:P ratios. Chlorophyta are negatively influenced by high Si:P or Si:N ratios but positively influenced by high N:P ratios.

Tissue-culture plate experiments

The tissue-culture experiments were conducted for 9 wk, although the species composition of the bottom flora had already stabilized after 4 wk. Again, there was a satisfactory replication between parallel treatments. At the level of higher taxa, the same trends showed up as in the agar-plate experiments (Fig. 4). Under P limitation (N:P = 45:1) diatoms dominated at high Si:P ratios and were gradually replaced by green algae at declining ratios. Under a balanced supply of N and P (15:1) diatoms were displaced by green algae and cyanobacteria as Si:P or Si:N ratios declined. Under N-limitation (N:P = 5:1) diatoms dominated at high Si:N ratios and cyanobacteria at low ones. A multiple regression analysis performed in the same way as for the agar-plate experiments confirmed these trends:

Diatoms: arcsin√p
(Si:N or P) = 0.50 + 0.69 log₁₀
r² = 0.94; p < 0.0001

Cyanobacteria: arcsin√p
(Si:N or P) = 1.47 - 0.34 log₁₀
r² = 0.91; p < 0.0001

Chlorophyta: arcsin√p
(Si:N or P) = -0.30 - 0.32 log₁₀
r² = 0.81; p < 0.0001

At the species level there were differences to the agar-plate experiments:

Calothrix aeruginosa Thuret was the only cyanobacterium under N-limiting and N:P-balanced conditions. This species fixes molecular nitrogen. Microcoleus chthonoplastes formed small populations under P limitation and low Si:P ratios. The green algae were represented only by Rhizoclonium riparium.

Among the diatoms, Amphipleura rutilans Cleve dominated under P limitation and high Si:P ratios, whereas Entomoneis paludosä Reimer dominated under N limitation and high Si:N ratios. Under a bal-
TISSUE PLATES. HIGH LIGHT

<table>
<thead>
<tr>
<th>PO₄</th>
<th>NH₄NO₃</th>
<th>Si</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 µM</td>
<td>45 µM</td>
<td>80 µM</td>
</tr>
<tr>
<td>6 µM</td>
<td>15 µM</td>
<td>2 µM</td>
</tr>
</tbody>
</table>

Fig. 4. Final species composition of the periphyton in the tissue-culture-plate experiments. Amph: Amphipleura rutilans; Aora: Amphora flebilis; Calo: Calothrix aeruginosa; Ento: Entomoneis paludosa; Nicl: Nitzschia closterium; Micr: Microcoleus chthonoplastes; Rhiz: Rhizoclonium riparium

The major findings of my experiments were:

- Light has a strong impact on the selection of competitively dominant species, but it does not determine at which Si:P or Si:N ratios diatoms become dominant. The same has been found for marine phytoplankton (Sommer 1994a).
- The nitrogen source (ammonium vs nitrate) has no important impact on the outcome of nitrogen competition, although in theory changes in the ratio of substitutable resources could be important (Tilman 1982). Similar results have been found for marine (Sommer 1995, 1996) and for freshwater phytoplankton (Makulla 1995).
- Under P-limited conditions diatoms become increasingly dominant with increasing Si:P ratios. Under N-limited conditions the same happens with increasing Si:N ratios. So far, the increase of diatoms with increasing Si:P ratios has been reported from experiments with freshwater phytoplankton (Sommer 1983, Kilham 1986, Tilman et al. 1986) and the increase of diatoms with increasing Si:N ratios from experiments with marine phytoplankton (Sommer 1986, 1994a, 1995, 1996).
- Under low Si supply cyanobacteria dominate at low N:P ratios and chlorophytes at high ones. This pattern has been reported for freshwater phytoplankton (Tilman et al. 1986) but not for marine phytoplankton where chlorophytes are usually quite unimportant.

DISCUSSION

It was not my intention to mimic natural conditions in these experiments. Instead, I intended to discover robust patterns in the competitive response of periphyton communities to various nutrient ratios. This philosophy had been successful in phytoplankton ecology and there was no a priori reason why it should not work in periphyton. Conditions in in vitro competition experiments differ considerably from natural conditions. Some of the differences are an inherent necessity in experimentation because the effects of competition as the mechanism under study have to be isolated from confounding influences such as grazing and physical disturbances. Other differences concern the surface properties of growth substrata, restrictions of 3-dimensional growth, and the impact of water movements which transport both organisms and nutrients.

The potential impact of some of the differences in physical conditions can be seen by comparing the agar-plate and the tissue-culture-plate experiments. Agar is known to be a highly selective medium, thus restricting the number of species from which competitive winners can be selected. In addition to the surface properties of agar, the agar plates enforced a 2-dimensional growth mode with tight contact to the surface of the agar. In the tissue-culture plates, filamentous algae could grow in an erect mode and serve as a substratum for the growth of epiphytic unicells. This difference in the 'architecture' (sensu Wetzel 1983) of the periphyton community probably accounted for much of the difference at the species level between both types of cultures. Despite the species differences there was good agreement in the competitive performance of higher taxa. This agreement even extends to the competitive performance of higher taxa in phytoplankton communities. Therefore, it seems plausible that the physiological requirements and abilities which are decisive for the outcome of nutrient competition show a sufficient degree of uniformity within higher taxa to make comparisons between laboratory experiments and natural sites possible.

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- Under low Si supply cyanobacteria dominate at low N:P ratios and chlorophytes at high ones. This pattern has been reported for freshwater phytoplankton (Tilman et al. 1986) but not for marine phytoplankton where chlorophytes are usually quite unimportant.
The dominance of diatoms decreases with decreasing N:P ratios, but this effect is not strong enough to prevent diatom dominance at high Si:N ratios. This pattern has not previously been reported.

LITERATURE CITED


Manuscript first received: December 22, 1995
Revised version accepted: June 6, 1996