Differential Responses of Lichen Symbionts to Enhanced Nitrogen and Phosphorus Availability: An Experiment with Cladina stellaris

SARI MAKKONEN1,*, RIIKKA S. K. HURRI1 and MARKO HYVÄRINEN2

1Department of Environmental Sciences, University of Kuopio, POB 1627, FI-70211 Kuopio, Finland and 2Botanical Gardens, Department of Biology, University of Oulu, POB 3000, FI-90014 Oulu, Finland

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Key Results

Background and Aims
Lichens can be both nitrogen- (N) and phosphorous- (P) limited and thus may be susceptible to nutrient enrichment. Nutrient enrichment with N and P may have differing impacts on the lichen structure because of different physiological responses of fungal and algal partners to these nutrients. The hypothesis was tested that the differential responses of lichen symbionts to enhanced availability of N and P is reflected in the lichen thallus structure and the wall-to-wall interface between the algal and fungal cells.

Methods
Lichen cushions of Cladonia stellaris were treated with one P and two N concentrations alone and in combination that yielded total depositions of approx. 300 (moderate) and 1000 (high) mg N m−2 and 100 (high) mg P m−2 over an experiment lasting 14 weeks. The effects of N and P inputs on the relative volumes of fungal and algal cell in the medullary tissue and on the thallus structure were studied using light microscopy. The interface between algal and fungal cell walls was examined using transmission electron microscopy.

Key Results

The influence of excess P on the lichen thallus structure was stronger than that of additional N. Addition of P reduced the N : P ratio in podetia, the proportion of the medullary layer volume occupied by the algal cells, the thallus volume occupied by the internal lumen, and the algal cell-wall area covered by fungal hyphae.

Conclusions
Ecologically realistic changes in the availability of key macronutrients can alter the growth of symbionts. Reduction in the proportion of photobiont cells indicates that the application of P either stimulates fungal hyphal growth in the medullary tissue or impairs the cell division of the algal cells. The results suggest that both the N and P availability and thallus N : P ratio affect the growth rates of lichen symbionts.

Key words: Lichen, N, P, nutrient status, algal cell, fungal cell, thallus structure, Cladina stellaris.

INTRODUCTION

Several field and laboratory studies have indicated that the growth of large foliose and fruticose lichens may be limited by nitrogen (N; Crittenden et al., 1994 and references therein; Kurina and Vitousek, 1999; Palmqvist and Dahlman, 2006). However, little is known about the effects of nutrient deficiency on lichen structure. The results of fertilization experiments (Kauppi, 1980; Roy-Arcand et al., 1989) suggest that the impact of enhanced N on lichen growth may be biased towards the growth of the photobiont rather than the mycobiont.

Increases in photobiont populations and a concomitant disruption of the symbiosis, followed by thallus disintegration in extreme cases, have long been known to be a response to ecologically unrealistic N concentrations (e.g. Smith and Griffiths, 1969). Over-growth of the algal cells has been reported in highly N-enriched Cladina lichens (e.g. Kauppi, 1980). However, the impact of moderate or ecologically realistic changes in the N input on lichen anatomy and especially on the symbiotic interaction between photobiont and mycobiont are still poorly understood.

The growth of lichens, especially N-fixing lichens, may also be limited by P (Crittenden et al., 1994). However, to our knowledge, there are no experimental data available regarding the effect of P-limitation on lichen growth. Hyvärinen and Crittenden (2000) have shown with Cladina portentosa that P recycles from the senescing parts of thalli to the growing apices. They interpreted these findings as a source–sink relationship within the thallus and an adaptation of lichen to P-limited sites. In fungi, P translocation is suggested to be symplastic, based on cytoplasmic streaming through multi-perforate septa (Wetmore, 1973; Hyvärinen and Crittenden, 2000). N has also been shown to resorb from senescent basal tissue and to be recycled within the thallus (Kytoväita, 2000). N has also been shown to resorb from senescent basal tissue and to be recycled within the thallus (Kytoväita, 2000). Both key macronutrients must be actively translocated from the older parts to promote the growth of apices. Thus alterations in the availability of N and P may also result in physiological changes in the lichen thallus.

Dahlan et al. (2003) measured the concentrations of chlorophyll a, ergosterol and chitin in Hypogymnia physodes and Platismatia glauca subjected to N-rich fertilization for several years. They concluded that the relative investment of N in the lichen thallus was radically changed in favour of the photobiont and suggested that increased photosynthetic capacity may be an adaptation to alleviate the carbon cost of high-N tissues. These indirect measurements of the relative investment of N into the different symbionts show that the physiological changes in the photobiont and mycobiont may be coupled with the general nutrient regime. Whether these changes are also reflected in the lichen thallus structure remains to be shown.

The aim of this study was to investigate the effect of moderately enhanced N or P availability, alone and in...
combination, on the growth of alga and fungus, and on the podetium structure in *Cladina stellaris*. Each cell of the mature photobiont is usually enveloped by hyphae (Nash, 1996). We hypothesized that the proportion of the algal cell-wall area covered by fungal hyphae might vary if the growth rate of lichen partners either increase or decrease. It was expected that the combined N and P treatments would not necessarily bias the growth of symbionts as both the alga and fungus might benefit from the moderate nutrient enhancement.

**MATERIALS AND METHODS**

*Study species*

*Cladina stellaris* (Opiz) Brodo is a mat-forming fruticose lichen that dominates field layers of the oligotrophic and well-drained environments of boreal and arctic zones (Longton, 1988; Ahti and Oksanen, 1990). *Cladina stellaris* grows vegetatively by producing new growth at the top of each podetium, thus lengthening the internode formed in the previous year. The thallus structure of *C. stellaris* and other *Cladina* spp. differs from many other fruticose lichens in that it lacks an outer cortex. The outer layer of *C. stellaris* consists of algal cells and loosely packed hyphal tissue resembling the medulla (Figs 1 and 2). Below the upper layer is the inner cortex in which fungal cells are tightly packed. The core of podetium, which is empty, is called here an internal lumen.

*Experimental design and fertilization treatment*

Intact 20 × 26 cm cushions of *Cladina stellaris* together with 5-cm depth of topsoil were transported from the island of Hailuoto (65°01′N, 24°47′E) in northern Finland to the experimental field of the University of Oulu (approx. 25 km east from the site of collection). Six blocks, each of which contained seven 20 × 26 cm quadrats separated by 30–40 cm buffer zones, were set up randomly in the field roughly 30 m apart from each other, with lichen cushions originating from the same place forming one block. Lichen cushions and the underlying soil were kept in plastic boxes with drainage holes in the bottom and embedded within soil.

The following treatments were applied in factorial combinations to lichen cushions within each block: P (one concentration), N (in two concentrations), and distilled water that represented the ‘control’ level of the nitrogen and phosphorus treatments (subsequently referred to as P: control and high; and N: control, moderate and high). In addition, a dry control (DC) that was not treated with any solution was included in each block in order to test whether the addition of water alone influenced thallus structure.

Nitrogen was applied as NH₄NO₃ in concentrations of 0.16 and 0.48 mM for the ‘moderate’ and ‘high’ N treatments, respectively, and P as Na₂PO₄.H₂O at a concentration of 0.043 mM for the ‘high’ P treatment. Applications were made three times a week, and the pH of solutions ranged from 5.8 to 6.2. The concentrations were selected to yield total depositions of approx. 100 (high) mg P m⁻², and 300 (moderate) and 1000 (high) mg N m⁻² over 14 weeks that the experiment was carried out. The level of N deposition in the ‘high’ treatment was roughly the same magnitude as in industrialized areas in Britain where *Cladina* lichens are still found (see Hyvärinen and Crittenden, 1998b and references therein). In the combined N and P treatments the salts were prepared in the same solution to keep the spraying volume the same for all treatments.
The background deposition of total N during the experiment was 48.5 mg N m\(^{-2}\) and 51.9 mg N m\(^{-2}\) as collected in two rain gauges located 0.7 km south-east and 2.4 km south-west from the study site. The background deposition of P was not measured at those sites since it is generally considered to be negligible. The study site was surrounded by forests and hence protected from potential short-distance deposition of P-rich particulates (e.g. from agriculture), which are generally regarded to be the main form of P deposition (e.g. Newman, 1995).

All the quadrats except DC were sprayed with 100 mL of solution three times a week during the study period from 28 May to 3 September, 1998. Spraying took place only during natural rainfall events or early in the morning when the lichens were naturally moist after the morning dew. Precipitation in the area was 241.2 mm during the exposure period (Helminen et al., 1998), and the spray irrigation increased the amount of water by approx. 31% in the experimental plots. A 2-week period without irrigation preceded the harvest in order to allow the surface deposits to be assimilated or washed away by rainwater.

**N : P ratio**

Biomass N : P ratio was calculated as a quotient of [N] and [P] at each application. The concentrations of N and P in the apices of *C. stellaris* were determined as described by Hyvärinen et al. (2003).

**Microscopy**

Microscopic examination was conducted on five *C. stellaris* thalli chosen randomly from each treatment in six blocks. The fourth internode from the top of each podetium was fixed in 2.5% glutaraldehyde in phosphate buffer (pH 7), and post-fixed in 1% buffered OsO\(_4\) (Holopainen, 1982). Samples were dehydrated in ethanol and embedded in Ladd's LX-112 resin. Semi-thin sections were cut with an Ultratome III (LKB-Producer AB, Sweden) and stained with toluidine blue. A longitudinal section through the medulla, cortex and internal lumen within the thallus was photographed (TMAX ASA 100) with a NIKON Microphot-FXA light microscope (Nikon Corporation, Japan) at \(x \times 4\) magnification.

The proportions of the lumen, inner cortex (i.e. a central cylinder of *Cladina*; Nash, 1996) and medulla within the secondary thallus were analysed using point frequency analysis as described in Tarhanen et al. (1997). A 10 \(\times\) 15 grid with a random point within each grid cell was placed on the photograph and numbers of points hitting the target were counted. The longitudinal section size of the internode was estimated by multiplying the area of the grid cell by the number of counted points. The relative volumes of medulla, cortex and internal lumen within each thallus internode were derived from the surface area measurements on longitudinal sections. The volume density of algal cells (%) was estimated by calculating the proportion of the medullary layer volume occupied by the algal cells. The medullary layer volume consisted of the algal and fungal cells, and the interhyphal air spaces.

Ultrathin sections of the lichens were cut with an Ultracut E ultramicrotome (Reichert-Jung AG, Austria) and stained with uranyl acetate and lead citrate. The samples were examined with a JEM 12000 EX electron microscope (JEOL Ltd, Japan), and images of contacts between algal and fungal symbionts were acquired with a digital camera. The length of interface between algal and fungal cells was measured on the cell-wall surface (i.e. perimeter) of cross-sectional algal cell using NIH-Image software for Windows (Scion Corporation, Maryland, USA). The wall-to-wall interface between symbionts was expressed as a percentage of the algal cell perimeter (\(L_{\text{fungal wall}} : L_{\text{algal cell perimeter}}\)). In addition, size and maximum and minimum diameters of the algal cells were analysed. The shape of cells was determined by comparing the ratio of maximum and minimum diameters. The mean number of algal cells examined per treatment was 369-6 (range, 286–485).

**Statistical analysis**

The impact of N and P application on the thallus structure in *C. stellaris* was tested by factorial ANOVA with randomized blocks using the statistical package SPSS (SPSS Inc, Chicago, IL, USA). The model (N + P + block) + (N \(\times\) P) was fitted to the data. Whenever the influence of ‘block’ was statistically insignificant the term was dropped from the final model.

**RESULTS**

In control lichens, the average thallus N : P ratio was 10 : 1 (± 1-0, s.d.) and increased with increasing N supply \((F_{2,25} = 4.58, P < 0.05; \text{Fig. 3})\). The addition of P reduced this ratio by 15–20% in all the N treatments \((F_{1,25} = 10.66, P < 0.01)\).

The blocks in the experimental design did not have any effect on the structural characteristics of the thallus that were measured, and the term was therefore dropped from the models given in Tables 1, 2 and 3.

The main effect of P was observed as a significant change in the proportions of thallus volume occupied by the lumen and medulla (Table 1). The high P treatment increased the proportion of thallus volume occupied by the medullary tissue and reduced the volume density of the internal lumen (Fig. 4). In contrast to the P treatment, addition of N, particularly in the moderate N treatment, reduced the volume density of the medullary tissue and increased that of the lumen (Table 1 and Fig. 4). The lumen volume density was correlated negatively with [P] and positively with the thallus N : P ratio (Table 4). Both [N] and [P] (% of thallus dry mass) were positively correlated with the thallus volume occupied by the inner cortex (Table 4). However, the main effect of N and P treatments on the volume density of inner cortex was not statistically significant (Fig. 4 and Table 1). The total longitudinal section area of internode was positively correlated with the internal lumen volume density (%; \(r = 0.351, P < 0.05, n = 42\)). The total longitudinal section area of the thallus internode was not significantly affected by the different nutrient treatments applied (data not shown).
The proportion of medullary layer volume occupied by the algal cells (% of medulla) was significantly affected by P (Table 2). The high P treatment reduced the volume density of the algal cells (Fig. 5) which varied from 5 to 29% in the low P treatment, and from 5 to 11% in the high P treatment. Addition of P also significantly reduced the wall-to-wall interface between algal and fungal cells (Table 3 and Fig. 6). The volume density of the algal cells and wall-to-wall contact between symbionts were not significantly affected by the N supply (Table 2), nor were there significant changes in the size and shape of algal cells in any of the fertilization treatments (data not shown).

**DISCUSSION**

The present study revealed that moderately enhanced availability of the macronutrients N and P affects the structure of podetia in *C. stellaris*. Enhancements of N and P, when given independently, caused changes in the thallus volumes occupied by the internal lumen and the medulla, in both high P and moderate N treatments. Moreover, it appears that the influence of P fertilization on the lichen thallus structure was stronger than that for N fertilization. Given the fact that both N and P treatments moderately increased the total concentrations of those elements in the thalli (Hyvärinen et al., 2003), it can be speculated that thallus anatomy may change due to the nutrient content of the lichen rather than nutrient availability.

The N uptake efficiency varies depending on the N economy of the lichen as well as the mode of uptake of N into the thallus. In general, when N is received in ionic form in low concentrations, either from natural rainfall (Hyvärinen and Crittenden, 1998b) or applied solutions (Crittenden, 1996), the estimated uptake efficiency of non-N-fixing lichens has ranged between 90–100%. The uptake efficiency of NO$_3^-$ and NH$_4^+$ may be considerably

**TABLE 1. Factorial ANOVA for the volumes (%) of medulla, inner cortex and internal lumen in the thallus of *C. stellaris***

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>d.f.</th>
<th>Sum of squares</th>
<th>Mean squares</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medulla</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N treatment</td>
<td>2</td>
<td>0.02</td>
<td>0.01</td>
<td>2.26</td>
<td>0.118</td>
</tr>
<tr>
<td>P treatment</td>
<td>1</td>
<td>0.04</td>
<td>0.04</td>
<td>8.24</td>
<td>0.007</td>
</tr>
<tr>
<td>N x P</td>
<td>2</td>
<td>0.02</td>
<td>0.00</td>
<td>1.75</td>
<td>0.189</td>
</tr>
<tr>
<td>Residuals</td>
<td>36</td>
<td>0.16</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inner cortex</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P treatment</td>
<td>1</td>
<td>0.00</td>
<td>0.00</td>
<td>0.79</td>
<td>0.380</td>
</tr>
<tr>
<td>N x P</td>
<td>2</td>
<td>0.00</td>
<td>0.00</td>
<td>0.25</td>
<td>0.783</td>
</tr>
<tr>
<td>Residuals</td>
<td>36</td>
<td>0.07</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Internal lumen</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P treatment</td>
<td>1</td>
<td>0.06</td>
<td>0.06</td>
<td>18.57</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>N x P</td>
<td>2</td>
<td>0.02</td>
<td>0.02</td>
<td>3.13</td>
<td>0.056</td>
</tr>
<tr>
<td>Residuals</td>
<td>36</td>
<td>0.11</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**TABLE 2. Factorial ANOVA for the algae volume (%) in the medulla layer of *C. stellaris***

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>d.f.</th>
<th>Sum of squares</th>
<th>Mean squares</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>N treatment</td>
<td>2</td>
<td>0.00</td>
<td>0.00</td>
<td>0.14</td>
<td>0.892</td>
</tr>
<tr>
<td>P treatment</td>
<td>1</td>
<td>0.09</td>
<td>0.00</td>
<td>25.96</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>N x P</td>
<td>2</td>
<td>0.00</td>
<td>0.00</td>
<td>0.12</td>
<td>0.883</td>
</tr>
<tr>
<td>Residuals</td>
<td>36</td>
<td>0.12</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**TABLE 3. Factorial ANOVA for the wall-to-wall interface between algal and fungal cells (%) in the thallus of *C. stellaris***

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>d.f.</th>
<th>Sum of squares</th>
<th>Mean squares</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>N treatment</td>
<td>2</td>
<td>0.00</td>
<td>0.00</td>
<td>0.01</td>
<td>0.991</td>
</tr>
<tr>
<td>P treatment</td>
<td>1</td>
<td>0.02</td>
<td>0.02</td>
<td>9.48</td>
<td>0.004</td>
</tr>
<tr>
<td>N x P</td>
<td>2</td>
<td>0.00</td>
<td>0.00</td>
<td>0.21</td>
<td>0.812</td>
</tr>
<tr>
<td>Residuals</td>
<td>36</td>
<td>0.07</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
lower in N-fixing lichens. For example, Dahlman et al. (2002) estimated that only 2–27% of N supplied as aqueous NH$_4^+$ or NO$_3^-$ solution to Peltigera aphthosa and Nephroma arcticum could be taken up by the lichen thalli.

There are few estimates of P uptake efficiency in lichens. Farrar (1976) demonstrated that Hypogymnia physodes can rapidly absorb P from bathing solutions containing PO$_4^{3-}$ in concentrations typical of rainfall (Newman, 1995). In addition, Hyvärinen and Crittenden (1998a) measured PO$_4^{3-}$ uptake efficiency of Cladina portentosa in a spray chamber (lysimeter). In their experiment, the lichen mat was able to scavenge approx. 80% of P supplied at a concentration of 30 µg P L$^{-1}$ and at a rate of approx. 100 mL h$^{-1}$. In our study the P concentration of the solution applied was considerably higher (0.043 mM, i.e. 1.33 mg P L$^{-1}$) and therefore it is difficult to estimate how much of the P would have been immediately absorbed. Nevertheless, the increase in P concentration was statistically significant (Hyvärinen et al., 2003), confirming that a great proportion of the P was scavenged by C. stellaris.

One problem commonly faced with studies involving fertilization of lichens is the adverse effects of ammonium ions. Toxicity of ammonium ions to ATP formation in chloroplasts and mitochondria is well known in the higher

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Fig. 4. Box-plot showing the volumes (% of thallus volume) of medulla, inner cortex and internal lumen in C. stellaris after N and P treatments. The line in the box indicates the median value of the data values. The ends of the vertical lines indicate the minimum and maximum data values. Asterisk = extremes; circle = outliers.
plants (e.g. Salisbury and Ross, 1992). Because ion uptake occurs over the entire thallus surface of lichens, it may be assumed that they could be more susceptible to harmful effects of ions than vascular plants. However, such toxic effects of NH₄⁺ usually occur only at concentrations beyond the levels used in our study. For example, Gaio-Oliveira et al. (2004) studied the effect of ammonium ion addition on Xanthoria parietina. They concluded that ammonium ions resulted in toxic effects on the mycobiont and photobiont at different threshold concentrations (0.34 and 0.69 m NH₄⁺, respectively), the latter being more tolerant to ammonium ions. In terms of N, these concentrations are approx. 350–700 times higher than the highest value used in the present study (including background deposition; see Methods).

It should be noted that the relative growth rate of lichens in terms of dry mass was not significantly promoted by addition of N and P (Hyväriinen et al., 2003). Given a weak positive relationship between the total internode longitudinal sectional area and the thallus volume occupied by the lumen, the lumen volume density gain may indicate an increase in radial growth and lichen area (i.e. cover) expansion. Data on lichens with green algal symbionts such as C. stellaris are lacking, but it has been recently shown with nitrogen-fixing lichens (i.e. Nephroma arcticum and Pertigera aphthosa) that lichen expansion is limited by thallus N status (Sundberg et al., 2001; Dahlman et al., 2002), while lichen weight gain was mainly affected by irradiance and carbon assimilation, and not by N supply (Sundberg et al., 2001).

The low volume density of the algal cells (% of medulla) together with the increased thallus volume occupied by medullary tissue after the high P treatment might indicate an increase in fungal hyphal growth in relation to the number of algal cells. The reduced wall-to-wall interface between the algal and fungal cells also seems to point to enhanced fungal growth. Studies on ectomycorrhizal associations have shown that additional P application to the hyphal compartment stimulates external fungal hyphal growth, simultaneously improving the nutrient status of host plant (Jentschke et al., 2001). In this study, P enrichment may have resulted in nutrient imbalances in the lichen that led to enhanced hyphal production in the medullary layer. The reduced proportion of thallus volume occupied by algal cells might also be explained either by the lack of growth stimulation or impaired algal cell division due to excess P. Data for lichens are lacking, but Martínez-Abaigar et al. (2002) found in a laboratory study that high tissue P concentration in liverwort (Jungermannia exsertifolia) was capable of reducing net photosynthesis and increasing the proportion of carotenoids to chlorophylls, indicating P toxicity in the bryophyte.

### TABLE 4. Correlation matrix of thallus morphological characteristics, nitrogen and phosphorus concentrations, and N : P ratio in C. stellaris

<table>
<thead>
<tr>
<th></th>
<th>[N] (% of thallus d. wt)</th>
<th>[P] (% of thallus d. wt)</th>
<th>N : P ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>32</td>
<td>33</td>
<td>31</td>
</tr>
<tr>
<td>[P] (% of thallus d. wt)</td>
<td>0.640***</td>
<td>0.789***</td>
<td></td>
</tr>
<tr>
<td>N : P ratio</td>
<td>-0.094</td>
<td>-0.036</td>
<td>-0.021</td>
</tr>
<tr>
<td>Medulla (% of thallus internode)</td>
<td>0.024</td>
<td>0.305*</td>
<td>-0.348*</td>
</tr>
<tr>
<td>Fungal volume (% of medulla)</td>
<td>0.066</td>
<td>-0.173</td>
<td>0.180</td>
</tr>
<tr>
<td>Algal volume (% of medulla)</td>
<td>0.376*</td>
<td>0.369*</td>
<td>-0.261</td>
</tr>
<tr>
<td>Inner cortex (% of thallus internode)</td>
<td>-0.154</td>
<td>-0.409*</td>
<td>0.360*</td>
</tr>
<tr>
<td>Lumen (% of thallus internode)</td>
<td>-0.229</td>
<td>-0.170</td>
<td>0.102</td>
</tr>
<tr>
<td>Wall-to-wall interface between symbionts (%)</td>
<td>-0.229</td>
<td>-0.170</td>
<td>0.102</td>
</tr>
</tbody>
</table>

*P < 0.10, *P < 0.05, **P < 0.01, ***P < 0.001.

**FIG. 5. Box-plot of relative algal cell volume (% of medullary layer volume) after N and P treatments. The line in the box indicates the median value of the data values. The ends of the vertical lines indicate the minimum and maximum data values.**
It has been shown previously that P addition increases the concentration of usnic acid in *C. stellaris* but that the increase was less evident in moderate N + P treatments (Hyvärinen *et al.*, 2003). The enhanced usnic acid level might be associated with increased fungal hyphal growth in the medulla because usnic acid mainly occurs on the surface of fungal hyphae. The P-induced elevation in the usnic acid concentration may also explain the low medullary layer volume occupied by the algal cells. Bacˇkor *et al.* (1998) demonstrated that usnic acid inhibited the growth of the photobiont *Trebouxia irregularis* isolated from *Cladina mitis*. By increasing hyphal growth and secondary metabolite production, the fungal partner probably inhibits the cell division of the algal partner.

Knowledge is still limited about the habitat factors or environmental conditions that affect the symbiotic interaction between algae and fungi in relation to lichen growth. Sun and Friedmann (2005) found a positive relationship between alga-to-fungus ratio and habitat summer temperature in *Cladina rangiferina*. They suggested that regulation of the ratio of producer (alga) to consumer (fungus) directly contributes to adaptation to a wide range of thermal regimes and to the distribution of lichens. The differential responses of fungal and algal growth to N and P fertilization observed in the present study suggest that the tissue nutrient content, and particularly the nutrient balance, affect resource allocation in the lichen thallus. In aquatic ecosystems (Sridhar and Bärlocher, 2000; Fong *et al.*, 2004), as well as in isolated lichen symbions (Crittenden *et al.*, 1994), the growth of free-living algal and fungal cells have been shown to be N- and P-limited. In the lichen symbiosis, the fungal partner primarily determines thallus form and the structural characters of tissue layers, but the algal partner may also influence it (Nash, 1996). The balance between N and P supply, as reflected in N : P ratios in plant biomass, has been used mainly to assess whether N or P is more limiting for biomass production. Variation in [N] is often more important in determining the N : P ratios of bryophytes or lichens (Güsewell, 2004). In this short-term experiment, addition of P clearly reduced the N : P ratios. The negative effect of P on the algal partner suggests that algal growth was N-limited even though the N concentration of the thalli increased.

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**LITERATURE CITED**


