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Growth of two peat-forming mosses in subarctic mires: species interactions and effects of simulated climate change

Mats Sonesson, Bengt Å. Carlsson, Terry V. Callaghan, Sven Halling, Lars Olof Björn, Monika Bertgren and Ulf Johanson


In patches of co-occurring species in natural plant communities, there is a finely poised balance between species in the ways in which they respond to prevailing moisture and temperature regimes. However, environmental change scenarios, in which temperature, moisture and ultraviolet-B radiation are suggested to increase, may favour one of the species. The imbalance is likely to occur at the levels of interactions between patches of the different species and at the shoot level when neighbouring shoots belong to different species. We increased temperature and UV-B in a two-way factorial experiment and increased water supply independently in two subarctic mire communities dominated by the mosses Sphagnum fuscum and Dicranum elongatum. The effects of simulated increase in UV-B were studied using two separate radiation systems, i.e. a "square wave" system and a "modulated" system.

When precipitation was enhanced, both species showed an increase in growth but this was not sustained beyond 5 mm per day. S. fuscum showed a 50% greater response to enhanced precipitation than did D. elongatum, as would be expected from their habitat preferences. Under ambient temperature, S. fuscum grew 67% faster than D. elongatum and this relative difference in response was maintained after one year under a temperature enhancement. The response by species over the winter period was moderated by their neighbours. S. fuscum growth was enhanced when it grew next to D. elongatum whereas D. elongatum grew better with neighbours of its own species. Increased temperature and UV-B radiation did not affect the interaction between the species.

Although a balance was maintained between the two species over the short duration of the experiment, potential was shown for an imbalance to occur over longer periods and particularly if winter warming and precipitation are greater than those in summer.

During the peak growing season 20% increased UV-B over ambient had a negative effect on S. fuscum under increased temperature but there were no overall seasonal effects on either species, irrespective of method of UV supplementation.

Approximately 3% of the Earth’s land surface is covered with peat and about 75% of this is in boreal and arctic areas (Kivinen and Pakarinen 1980). Peatlands sequester atmospheric carbon and store it in peat. They are estimated to contain 300 Gt carbon which is equivalent to about 40% of the carbon content of the atmo-
sphere (Sjörs 1982). There is little doubt that both the formation and decomposition of peat and the factors controlling these processes are of great importance for the global atmospheric carbon balance, particularly if predicted climatic change occurs and peatlands shift from carbon sinks to source status (Billings et al. 1982, 1983, Oechel et al. 1993, 1995, IPCC 1996, Malmer and Wallén 1996).

Subarctic and arctic peatlands are on permafrost to a large extent (Kalela 1939, Kalliola 1939, Ruuhijärvi 1960, Sonesson 1970a, Botch and Masing 1983). This affects decomposition as well as the formation of plant biomass and hence peat. Where the permafrost surface is deep in the soil profile below a thick active layer, oxidising conditions exist and aerobic decomposition is rapid (Christensen et al. 2000). In contrast, a high permafrost table during the growing season leads in general to wet and anaerobic soil conditions in the unfrozen active layer which in turn prevents decomposition of surface peat. However, in a climate change scenario with an increase in temperature and precipitation (Maxwell 1992, Cattle and Crossley 1995), particularly in the northern areas which have already experienced warming of up to 1.5°C per decade (Chapman and Welsh quoted in Weller 2000), both decomposition and plant growth will be affected. However, plant growth and decomposition will probably not be affected to the same extent because temperature is likely to differ between aboveground plant organs and subsurface peat. An effect of increasing humidity may be increased thermal conductivity of the peat. This can then result in a lowering of the permafrost table and in combination with increased temperature enhance decomposition and flux of CO₂. However, if surface peat layers are oxidised, the remaining layers will be less labile and the decomposition of these is also less temperature sensitive (Christensen et al. 1999).

Increased precipitation and temperatures in combination will also improve plant growth which will counteract the effect of greater CO₂ emission from increased decomposition. Hence, to what extent the subarctic-arctic peatlands will be sources or sinks for carbon in the future may largely depend on the rate and magnitude of change in thermal conductivity of the peat and the rate and magnitude of biomass formation.

UV-B radiation associated with stratospheric ozone depletion may also have an effect on plant growth (Searles et al. 2001). However, studies on the effects on mosses are few and with conflicting results. Experimentally added UV-B has been found to either stimulate (Sonesson et al. 1996), depress (Gehrke et al. 1996, Gehrke 1998) or to have no effect (Björn et al. 1999, Gwynn-Jones et al. 1999, Searles et al. 1999) on the performance of bryophytes.

Two moss species are particularly important for ombrotrophic peat formation in subarctic (continental) peatlands, viz. the hummock-forming Sphagnum fuscum Klingsgr. and Dicranum elongatum Schleich. (Kalliola 1939, Ruuhijärvi 1960, Sonesson 1970a). They form structures that may often be large, elongated, surrounded by or adjacent to, wet depressions (Sonesson 1970a, 1980a). Normally D. elongatum and lichens are dominant on the summits and S. fuscum on the slopes. The plants rely nearly exclusively on water and nutrients from precipitation (Malmer and Nihlgård 1980). This makes them particularly strongly exposed to climatic impacts. Stratigraphic studies have shown that there is usually a progressive succession in these sites from moss communities of wet depressions to S. fuscum-dominated slopes to D. elongatum-dominated summit communities, i.e. there is a successive development towards drier stages (Sonesson 1970b). The opposite pattern has also been observed, however, when S. fuscum and even mosses of wet areas succeed D. elongatum. Regressive development may be brought about by changes in the hydrology of the ecosystem. How much of the change that is due to inherent properties of the ecosystem (e.g. species interactions) and how much is due to external environmental factors (such as weather) cannot usually be determined from stratigraphic studies only (Sonesson 1970b). However, by experimentally manipulating the live plant cover, information may be gathered about some of the major mechanisms involved.

This study has two main objectives. Firstly, we use environmental manipulation experiments to study the likely response of communities to increase in precipitation, temperature, and ultraviolet-B radiation. Secondly, we seek to understand how interaction between co-existing species may be affected by such climate change.

Materials and methods
Experimental sites and vegetation
Two mires on permafrost, situated ca 10 km apart, at Abisko (68°21’N; 18°49’E) and Stordalen (68°21’N; 19°04’E) were used for the field studies. The studies at the Stordalen site were completed in 1973, those at Abisko in 1997 and 1998. S. fuscum and D. elongatum are dominant mosses on the higher, ombrotrophic parts in both mires (“hummock sites”, Sonesson 1970a). Plant communities in which these mosses are particularly prominent have been described by Sonesson (1970a, b, 1980a) as variants of an “Empetrum hermaphroditum–Vaccinium microcarpum association”. Lichens dominate in variants of the driest habitats while moisture demanding mosses such as Sphagnum balticum (Sw.) Lange become prominent over the wettest habitats of the association. Extreme (dry or wet) habitats have been avoided in the present study. We
have concentrated on habitats in the moderately wet to moderately dry range. The seasonal median, total and interquartile range of water tables of the habitats in relation to the bryophyte surface are shown in Sonesson (1970b). For an extensive description of other aspects of these mires, i.e. their vegetation and general ecology, see also Sonesson (1970a, 1980a).

Data on temperature and precipitation during the study periods are shown in Table 1. The Stordalen site is located at the same altitude as the Abisko meteorological station, i.e. 390 m above sea level, and the mean temperatures are similar, whereas precipitation is on average about 6% higher at Stordalen (Rydén 1980).

### Experimental design

#### Stordalen

Natural growth of the two species was measured in 16 plots (each approximately 10 × 10 cm) selected over a large sub-site (about 150 × 300 m) in each of three habitats for each species, viz. 2 (species) × 3 (habitats) × 16 (plots) = 96 plots in total (Sonesson and Johansson 1974). The statistical sampling design is described in Sonesson et al. (1980). The habitats were selected for being “moderately dry”, “intermediate”, and “moderately wet” according to community characteristics. The measurements were performed in 1973 between June 5 and September 15.

During part of the 1973 season the effect of increased precipitation on growth was simulated in an experiment on a neighboring sub-site, using forty plots selected to represent moderately dry habitats of each species. The experiment started on July 1 and was finished on September 15. Distilled and deionized water was mist-sprayed onto plots twice a day on 20 rain-free days during the first four weeks after which no further irrigation was carried out. The experimental design consisted of 2 species × 4 levels of daily water supply (control, 1, 5, 10 mm) × 5 replicate plots. Half of the daily water doses were applied in the mornings and half in the afternoons to minimize leaching effects. No irrigation was made during three rainy days during the first experimental period (July 1–15) and during five rainy days during the second period (July 16–27).

#### Abisko

Twenty-four years later, a two-way factorial experiment simulating increased temperature and ultraviolet-B radiation was established over a 30 × 75 m sub-site to study the effects on growth of the same two species.

Thirty plots were subjectively selected in moderately wet and intermediate habitats where *D. elongatum* and particularly *S. fuscum* were abundant. Each plot measured approximately 10 × 10 cm. The plots were distributed in a non-systematic way over the whole sub-site. Twenty plots were randomly selected for the experiment simulating future elevated UV-B levels using a modulated supplementation system (Bowles 1997, based on Caldwell et al. 1983) and ten plots were selected for a comparative experiment applying a “square wave technique” for UV-B supplementation (Johanson et al. 1995, Bertgren 1997, Gehrke 1998).

Temperature was increased by placing on the plots hexagonal open, UV-B-transparent plexiglas chambers of an ITEX-type (ITEX = International Tundra Experiment, Marion et al. 1997) that increase the average air temperature by about 2.0°C over ambient (Marion et al. 1997). The chambers were placed on small wooden blocks a few cm above the moss surface to improve ventilation (van Oijen et al. 1999) and to reduce possible effects of extreme warming events.

The UV dose was administered from small (15 cm) UV-B fluorescent tubes (Philips TL 12/4W) covered...
with 0.13 mm cellulose diacetate (Courtaulds, Derby, U.K.) to exclude UV-C radiation (Bertgren 1997). These were suspended under square frames on thin aluminium legs, 20 cm above the moss surface. The UV-B tubes were pre-burned for 48 h to get a stable output which was checked with a UV-B sensor. The UV-B dose over ambient corresponded to 20% ozone depletion under clear sky at this latitude. Control plots were equipped with UV-B frames under which 0.1 mm Mylar film/ filters were placed to exclude all UV radiation below 318 nm. All filters were changed once a week to avoid the effects of ageing.

The modulated system measured ambient UV-B radiation continuously and added the difference between ambient radiation and that corresponding to a 20% depletion below 318 nm. All filters were changed once a week to avoid the effects of ageing.

The modulated system measured ambient UV-B radiation continuously and added the difference between ambient radiation and that corresponding to a 20% ozone depletion under clear sky at this latitude. Control plots were equipped with similar frames and fluorescent tubes but with the total extra daily UV-B dose in a square wave centred on noon (Johanson et al. 1995, Bertgren 1997, Gehrke 1998). For the treatments and replicate numbers of the temperature and UV-B experiment, see Table 2.

All experiments were performed during two consecutive seasons. In 1997 work started on July 10 and was finished on September 5, in 1998 work started on June 12 and was finished on September 12. The equipment was removed during winter. The studies of the 1998 season comprised nearly a complete growing season in contrast to those of 1997. Thus we present results mainly from 1998.

**Growth measurements**

Straight stainless wires (Ø 0.6 mm) were inserted into the bryophyte carpet (extending from ca 10 cm underneath to 3 cm above the bryophyte surface) to serve as reference points for growth increment measurement (Sonesson and Johansson 1974, Sonesson 1980b). Points shaded by vascular plants were avoided. Leaves of Rubus chamaemorus L. in particular, expanding over the points were cut and removed. Every wire was marked with paint in a way that it could be individually recognized. Readings of the length increments were made using a thin (2 mm) glass tube (0.23 g) with a fused millimetre scale and equipped with a narrow flat collar at the lower end (Sonesson and Johansson 1974, Sonesson 1980b). The maximum error in reading was estimated to be ± 0.2 mm (Sonesson and Johansson 1974). All measurements were made by the same person. The wire method was a variant of a method proposed and pioneered by Clymo (1970). Ten wires were inserted in each plot at Stordalen. The mean value of the ten readings was the replicate value of the plot.

Growth interactions between D. elongatum and S. fuscum and the effects of increased UV-B and temperature were studied at Abisko in each plot by inserting wires so that they touched individuals of the same species or adjacent individuals of both species. In this way growth of individuals surrounded by individuals of the same species was measured, i.e. S. fuscum in S. fuscum (S/S) and D. elongatum in D. elongatum (D/D), and of individuals immediately adjacent to/surrounded by the other species, i.e. S. fuscum at/in D. elongatum (S/D) and D. elongatum at/in S. fuscum (D/S). Four points for each combination of individuals in each plot were equipped with wires, i.e. 4 × 4 wires per plot. For the evaluation of interactions, mean values of each set of four wires were the replicate values of each plot. For the evaluation of the effects of temperature and UV-B, mean values of the eight wire measurements of each species (S/S + S/D and D/D + D/S, respectively) were the replicate values of each plot.

Growth of the same individual S. fuscum capitula was easy to follow throughout the seasons while the glass tube usually hit more than one apex of the tiny D. elongatum individuals. This was probably the reason why bending of the D. elongatum apices caused by the weight of the tube was only rarely observed.

Possible movements of the wires due to thawing and freezing during early winter and early spring were controlled using three sets of ten wires which were inserted in three different parts of the Abisko site. They were inserted in moss carpets dominated by either of the two mosses studied or by a combination of both. The distance between these wires was approximately 10 cm. The experiment was established on September 9, 1997 when the levels of the wire tops were carefully determined using a precision levelling instrument. This procedure was repeated on June 9, 1998 (after thaw). There was nothing in the measurements that could be explained by instabilities of the reference wires due to frost heaving. The variance ratios (Zar 1984) between the autumn and spring readings were for Set 1 = 1.93, \( P > 0.20 \); Set 2 = 1.25, \( P > 0.20 \); Set 3 = 1.23, \( P > 0.20 \).

**Table 2. Treatments and replicate numbers of the temperature and UV-B supplementation experiment.**

<table>
<thead>
<tr>
<th>Modulated system</th>
<th>Number of plots</th>
</tr>
</thead>
<tbody>
<tr>
<td>UV-B</td>
<td>5</td>
</tr>
<tr>
<td>UV-B + temperature</td>
<td>5</td>
</tr>
<tr>
<td>Temperature</td>
<td>5</td>
</tr>
<tr>
<td>Control</td>
<td>5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Square wave system</th>
</tr>
</thead>
<tbody>
<tr>
<td>UV-B</td>
</tr>
<tr>
<td>Control</td>
</tr>
</tbody>
</table>

**Data analysis**

Natural growth and the precipitation experiments were analysed using three-way ANOVAs with Species, Habitat/Irrigation level and Time of season as independent
factors. For the two-way factorial experiment at Abisko, means of length increments over the summer season were analysed with a repeated-measures ANOVA with Time of season as a within-subject factor and Species, Neighbour identity, Temperature, and UV-B flux as between-subject factors. A separate two-way ANOVA with Species and Neighbour identity as factors was run for the winter season. All analyses were carried out using SPSS for Windows 10.0 (SPSS Inc., Chicago, IL).

Results

Natural growth

Growth was clearly related to type of habitat for both species \( (P < 0.001, \text{Fig. 1, Table 3}) \). The lowest increment was measured for dry habitats and the highest for wet habitats. This was particularly evident during the early and peak summer periods (Habitat × Time of season interaction, \( P < 0.001, \text{Table 3} \)). The increment of \( D. \) elongatum in wet habitats was much higher than that in dry habitats, fivefold during the early period, tenfold during the peak season. Overall, \( S. \) fuscum had a 67% larger increase in length than \( D. \) elongatum \( (P < 0.001) \) although the relative differences between the habitats were smaller in comparison with those of \( D. \) elongatum. The highest increments were measured during the peak season when growth in the wet habitats was ten times higher and nearly four times higher than in the dry habitats for \( D. \) elongatum and \( S. \) fuscum, respectively.

Simulated climate change

Increased precipitation had a significant effect on growth which generally increased \( (P < 0.01, \text{Fig. 2, Table 4}) \). This was particularly evident for plots irrigated at 1 mm per day, whereas the higher amounts resulted in no enhanced growth in comparison with the control \( (P > 0.05) \). During the early season, a 1 mm increase in precipitation nearly trebled the growth of \( D. \) elongatum. During the late season when irrigation had ceased, mean length increase was still about two thirds greater than that of the controls. Overall \( S. \) fuscum showed a 50% larger response than \( D. \) elongatum to enhanced precipitation \( (P < 0.001) \). An increase in precipitation by 1 mm (during the first two periods) tended to increase growth of \( S. \) fuscum by half over the controls during the early season and by approximately one third during the peak and late season. The insignificant effects of precipitation at a rate higher than 1 mm per day were similar to those for \( D. \) elongatum.

Increased temperature had a strong positive effect on the growth of both mosses \( (P < 0.001, \text{Fig. 3, Table 5}) \). Over the whole summer season of 1998 (June 12–September 12), \( S. \) fuscum increased 23% more than \( D. \) elongatum \( (P = 0.064, \text{Fig. 3, Table 5}) \). Significant interactions involving “Time of season”, “Species”, and “Temperature” indicated, however, that as the season progressed, the difference in growth between the two species increased, especially in plots with increased temperature (Fig. 4).

Table 3. ANOVA results for the natural growth investigation at Stordalen 1973.

<table>
<thead>
<tr>
<th>Source</th>
<th>Df</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>1</td>
<td>0.0116</td>
<td>16.4</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>Habitat</td>
<td>2</td>
<td>0.0157</td>
<td>22.3</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>Time</td>
<td>2</td>
<td>0.0130</td>
<td>18.5</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>Species × Habitat</td>
<td>2</td>
<td>0.0005</td>
<td>0.78</td>
<td>0.46</td>
</tr>
<tr>
<td>Species × Time</td>
<td>4</td>
<td>0.0018</td>
<td>2.51</td>
<td>0.084</td>
</tr>
<tr>
<td>Habitat × Time</td>
<td>4</td>
<td>0.0051</td>
<td>7.23</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>Species × Habitat × Time</td>
<td>4</td>
<td>0.0001</td>
<td>0.16</td>
<td>0.96</td>
</tr>
<tr>
<td>Error</td>
<td>255</td>
<td>0.0007</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*** \( P < 0.001 \).
Fig. 2. Mean daily length increment (mm ± 1 SE) of Dicranum elongatum and Sphagnum fuscum in moderately dry habitats at Stordalen during three summer periods (1973). Three intensities of increased precipitation over ambient were simulated. Irrigation was made during the first two periods only. (*Note that the mean figure and SE for the third period at 5 mm for D. elongatum were negligible, 0.0002 and ± 0.0001, respectively). N = 5. Increased precipitation had a significant effect on growth for both species (P < 0.001). S. fuscum had an overall 50% larger response than D. elongatum (P < 0.001, Table 4).

*Increased UV-B* had no statistically significant overall effect regardless of experimental system. However, there was a significant Time × UV-B interaction (P = 0.03, Table 5), which was mainly caused by S. fuscum responding negatively to increased UV-B under increased temperature at the peak of the growing season (Fig. 4, Table 5).

Species interactions

Neighbour identity had no effect on the length increment of either species during the summer season, neither in itself nor in interactions involving temperature or UV-B (Table 5). During the “winter period”, however, S. fuscum performed much better when growing with D. elongatum as a neighbour than in monoculture (Fig. 5). This was manifest as a significant Species × Neighbour interaction (P = 0.01; Table 6).

Discussion

Humidity and temperature effects

As humidity and temperature increased so did moss growth both in the specific habitats where the species grow and in the experimental manipulations. Overall there was 50% larger growth in S. fuscum than in D. elongatum due to enhanced precipitation. The experiment indicated that ambient precipitation is sub-optimal for growth of both species. The optimum value is unknown but seems to be closer to an extra 1 mm per

Table 4. ANOVA results for the irrigation experiment at Stordalen 1973.

<table>
<thead>
<tr>
<th>Source</th>
<th>Df</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>1</td>
<td>0.0072</td>
<td>22.9</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>Irrigation</td>
<td>3</td>
<td>0.0013</td>
<td>4.14</td>
<td>0.008**</td>
</tr>
<tr>
<td>Time</td>
<td>2</td>
<td>0.0106</td>
<td>33.7</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>Species × Irrigation</td>
<td>3</td>
<td>0.0001</td>
<td>0.39</td>
<td>0.76</td>
</tr>
<tr>
<td>Species × Time</td>
<td>2</td>
<td>0.0022</td>
<td>7.08</td>
<td>0.001**</td>
</tr>
<tr>
<td>Irrigation × Time</td>
<td>6</td>
<td>0.0002</td>
<td>0.49</td>
<td>0.81</td>
</tr>
<tr>
<td>Species × Irrigation × Time</td>
<td>6</td>
<td>0.0001</td>
<td>0.36</td>
<td>0.90</td>
</tr>
<tr>
<td>Error</td>
<td>96</td>
<td>0.0003</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**P<0.01, ***P<0.001.
day than to higher quantities. Some of the insignificant or negative effects of the higher precipitation may be an artefact of the method of application. Leaching of nutrients from the plants and ensuing reduced growth is difficult to avoid using distilled water even when precautions are taken to minimize such effects. Natural precipitation water contains nutrients of importance for growth, particularly in ombrotrophic sites (Malmer and Nihlgård 1980). Hence an increase of natural precipitation exceeding 1 mm per day together with its content of nutrients may enhance growth in contrast to what was indicated by the experiment. Our simulated 1 mm daily addition, however, corresponded to 55%–135% increase over the present normal summer values, which was much more than the predicted annual 20–25% increase for the middle of this century, particularly during winter (Maxwell 1992).

Increased temperature too has a large effect on growth even when increases are only approximately 2°C over ambient, corresponding to the lower range of predicted values for the middle of the 21st century (Maxwell 1992). As length and biomass are positively correlated (Sonesson and Johansson 1974), a combination of enhanced temperature by 2°C and precipitation by 20–25% may substantially increase the sequestering of carbon in habitats typical of the species. The superior growth of \textit{S. fuscum} should result in a dominance over \textit{D. elongatum} in moderately wet to moderately dry habitats. Also the natural growth today under ambient conditions indicates superior growth of \textit{S. fuscum}.

Mosses are strongly dependent on environmental conditions due to their poikilohydric nature (Sveinbjörnsson and Oechel 1992, Tenhunen et al. 1992, Green and Lange 1994). Hydration has a direct effect on performance and temperature has an indirect effect through its control of hydration. Thus, the interaction between hydration and temperature may be particularly important for moss performance. Growth in \textit{Hylocomium splendens} was found to be highest in humid and mild temperature sites (Callaghan et al. 1997) and there was an increase in biomass and rates of ramification during years with wetter climate than normal (Ökland 1997). Increased hydration and temperature together resulted in increased length in \textit{Polytrichum commune} (Potter et al. 1995), whereas irrigation alone stimulate length growth of \textit{Dicranum majus} (Hanslin et al. 2001).

During winter (i.e. late autumn–spring), \textit{S. fuscum} grew better with \textit{D. elongatum} as neighbour than in monoculture, whereas the opposite was true for \textit{D. elongatum} itself (Fig. 5). An explanation for the negative length values could be a dehydration of peat during winter and differences in mechanical strength between the two species. The strength of \textit{S. fuscum} shoots appears to be much lower than that of \textit{D. elongatum} and dehydration might cause the shoots to collapse. Differences in mechanical strength of the stems of \textit{Hylocomium splendens} have been shown to be related to erect and prostrate growth forms (Ross et al. 1998). The decrease may also be an effect of surface tension due to a lowering water level of the peat (Clymo, pers. comm.). During winter, \textit{S. fuscum} individuals growing adjacent to \textit{D. elongatum} will get support resulting in improved growth. After the thaw of seasonal frost above the permafrost table, hydration of the \textit{S. fuscum} mats will be restored and the shoots will return to their former shape.

**UV effects**

The negative UV-B effect on \textit{S. fuscum} during the peak season might have been an interaction with a drought

<table>
<thead>
<tr>
<th>Source</th>
<th>df</th>
<th>MS</th>
<th>(F)</th>
<th>(P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time of season</td>
<td>2, 208</td>
<td>10.8</td>
<td>23.7</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>Species</td>
<td>1, 104</td>
<td>8.39</td>
<td>3.49</td>
<td>0.064+</td>
</tr>
<tr>
<td>Neighbour identity</td>
<td>1, 104</td>
<td>0.17</td>
<td>0.07</td>
<td>0.79</td>
</tr>
<tr>
<td>Temperature</td>
<td>1, 104</td>
<td>70.5</td>
<td>29.3</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td>UV-B</td>
<td>1, 104</td>
<td>0.12</td>
<td>0.05</td>
<td>0.82</td>
</tr>
<tr>
<td>Time (\times) Species</td>
<td>2, 208</td>
<td>2.37</td>
<td>5.20</td>
<td>0.006**</td>
</tr>
<tr>
<td>Time (\times) Neighbour</td>
<td>2, 208</td>
<td>0.99</td>
<td>2.18</td>
<td>0.12</td>
</tr>
<tr>
<td>Time (\times) Temp.</td>
<td>2, 208</td>
<td>2.85</td>
<td>2.64</td>
<td>0.020**</td>
</tr>
<tr>
<td>Time (\times) UV-B</td>
<td>2, 208</td>
<td>1.62</td>
<td>1.55</td>
<td>0.039*</td>
</tr>
<tr>
<td>Species (\times) Neighbour</td>
<td>1, 104</td>
<td>0.08</td>
<td>0.03</td>
<td>0.86</td>
</tr>
<tr>
<td>Species (\times) Temp.</td>
<td>1, 104</td>
<td>0.18</td>
<td>0.08</td>
<td>0.78</td>
</tr>
<tr>
<td>Species (\times) UV-B</td>
<td>1, 104</td>
<td>2.62</td>
<td>1.09</td>
<td>0.30</td>
</tr>
<tr>
<td>Neighbour (\times) Temp.</td>
<td>1, 104</td>
<td>1.22</td>
<td>0.51</td>
<td>0.48</td>
</tr>
<tr>
<td>Neighbour (\times) UV-B</td>
<td>1, 104</td>
<td>2.75</td>
<td>1.15</td>
<td>0.29</td>
</tr>
<tr>
<td>Temp. (\times) UV-B</td>
<td>1, 104</td>
<td>1.24</td>
<td>0.52</td>
<td>0.47</td>
</tr>
<tr>
<td>Time (\times) Species (\times) Temp.</td>
<td>2, 208</td>
<td>1.82</td>
<td>3.99</td>
<td>0.020*</td>
</tr>
<tr>
<td>Time (\times) Species (\times) UV-B</td>
<td>2, 208</td>
<td>1.06</td>
<td>1.31</td>
<td>0.10+</td>
</tr>
<tr>
<td>Time (\times) Temp. (\times) UV-B</td>
<td>2, 208</td>
<td>1.36</td>
<td>2.99</td>
<td>0.053+</td>
</tr>
</tbody>
</table>

No three- or higher-order interactions other than those reported significant at or below \(P = 0.10\), \(P < 0.10\), \(P < 0.05\), \(P < 0.01\), \(P < 0.001\).
Fig. 4. Length increments for three periods at Abisko during the summer season 1998 (June 12–September 12) for Sphagnum fuscum and Dicranum elongatum growing under different levels of temperature and UV-B. Error bars are 1 SE. N = 5–10. During the peak season increased UV-B had a negative effect on S. fuscum under increased temperature (P = 0.03). There were no overall seasonal UV effects on either species (Table 5).

Gehrke's was about 55%, however (estimated from the figures in Gehrke 1998). This slight (possible) difference in variability cannot explain the disagreeing results. Different weather conditions during the experimental periods might have been a cause. The precipitation figures were approximately similar, but the mean air temperatures were 0.1°C–2°C lower than normal in 1994 and in 1995 but 1°C–2°C warmer in 1998. It is known that the enzyme system for DNA repair works better at higher temperatures (Takeuchi et al. 1996, Pakker et al. 2000, Paulsson and Björn 2001). The higher temperatures in 1998 might have reduced the sensitivity of the mosses to UV-B in contrast to the effects in 1994–95.

The insignificant or lack of effect of enhanced UV-B in our experiment is similar to what was reported for other mosses in the Arctic (Björn et al. 1999, Gwynn-Jones et al. 1999) and for the Antarctic population of Sphagnum magellanicum (Searles et al. 1999).

**Long-term effects**

If temperatures and precipitation increase, D. elongatum is likely to be hampered by shading from the overtopping S. fuscum and may eventually become extinct in wet to moderately dry habitats. However, this

**Table 6. ANOVA results for the “winter period” at Abisko, i.e. September 5, 1997–June 12, 1998.**

<table>
<thead>
<tr>
<th>Source</th>
<th>df</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>1</td>
<td>0.27</td>
<td>0.54</td>
<td>0.46</td>
</tr>
<tr>
<td>Neighbour identity</td>
<td>1</td>
<td>0.47</td>
<td>0.94</td>
<td>0.33</td>
</tr>
<tr>
<td>Species × Neighbour</td>
<td>1</td>
<td>3.36</td>
<td>6.72</td>
<td>0.010**</td>
</tr>
<tr>
<td>Error</td>
<td>76</td>
<td>0.50</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**P<0.01.**
may not apply to the “extreme dry habitats”, i.e. on the
summits of the hummocks, where S. fuscum does not
grow. The only competitors to D. elongatum on these
tops seem to be lichens. D. elongatum might survive in
these habitats even in the next 50–100 year climate
However, temperature and climate are changing con-
currently with other environmental factors such as ni-
trogen deposition. Over the last 20 years, many of the
ombrotrophic parts of the Stordalen mire have become
more eutrophic and formerly abundant S. fuscum has
become very restricted in abundance (Svensson et al.
1999). Although this study has identified the relative
importance of three environmental change factors for
two moss species, responses to other factors in com-
bination need to be studied before implications for bog
growth and atmospheric carbon sequestration can be
made.

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