Increased Air Temperature during Simulated Autumn Conditions Does Not Increase Photosynthetic Carbon Gain But Affects the Dissipation of Excess Energy in Seedlings of the Evergreen Conifer Jack Pine

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Temperature and daylength act as environmental signals that determine the length of the growing season in boreal evergreen conifers. Climate change might affect the seasonal development of these trees, as they will experience naturally decreasing daylength during autumn, while at the same time warmer air temperature will maintain photosynthesis and respiration. We characterized the down-regulation of photosynthetic gas exchange and the mechanisms involved in the dissipation of energy in Jack pine (Pinus banksiana) in controlled environments during a simulated summer-autumn transition under natural conditions and conditions with altered air temperature and photoperiod. Using a factorial design, we dissected the effects of daylength and temperature. Control plants were grown at either warm summer conditions with 16-h photoperiod and 22°C or conditions representing a cool autumn with 8 h/7°C. To assess the impact of photoperiod and temperature on photosynthesis and energy dissipation, plants were also grown under either cold summer (16-h photoperiod/7°C) or warm autumn conditions (8-h photoperiod/22°C). Photosynthetic gas exchange was affected by both daylength and temperature. Assimilation and respiration rates under warm autumn conditions were only about one-half of the summer values but were similar to values obtained for cold summer and natural autumn treatments. In contrast, photosynthetic efficiency was largely determined by temperature but not by daylength. Plants of different treatments followed different strategies for dissipating excess energy. Whereas in the warm summer treatment safe dissipation of excess energy was facilitated via zeaxanthin, in all other treatments dissipation of excess energy was facilitated predominantly via increased aggregation of the light-harvesting complex of photosystem II. These differences were accompanied by a lower deepoxidation state and larger amounts of β-carotene in the warm autumn treatment as well as by changes in the abundance of thylakoid membrane proteins compared to the summer condition. We conclude that photoperiod control of dormancy in Jack pine appears to negate any potential for an increased carbon gain associated with higher temperatures during the autumn season.

Temperature and daylength are important drivers of physiological changes in boreal evergreen conifers because they determine the length of the growing season. In late summer and early autumn the decrease of the daylength is a signal that initiates cold hardening, a transition of physiological processes that allow hardy plants to survive severe winter conditions. The cold hardening process includes the cessation of growth and long-term changes in the metabolism of the tree (Weiser, 1970; Bigras et al., 2001). Short days are supposed to induce the cold hardening most effectively, whereas decreasing temperatures affect this process only to some extent (Christersson, 1978).

As evergreen conifers keep their needles during winter, the cold hardening process involves a reorganization of the photosynthetic machinery (Ensminger et al., 2004), as the needles retain a substantial amount of chlorophyll (Chl) and proteins and therefore continue to absorb light. Absorption of light under winter conditions can cause photooxidative damage of the photosynthetic apparatus, created by an imbalance between the photochemical generation of electrons and their diminished utilization due to decreasing sink capacity, i.e. the down-regulation of metabolism and growth in winter (Oquist and Huner, 2003). To maintain the balance between light capture and energy utilization under conditions with altered sink capacity, energy flow and photosynthesis have to be adjusted by a process defined as photostasis (Oquist and Huner, 2003). Plants have developed a wide range of mechanisms to achieve this balance and to avoid photooxidative damage. This includes adjustments of the Chl
and protein concentration, changes in antenna size and organization, stoichiometry and aggregation of components of the photosynthetic apparatus, and a range of alternative energy dissipation pathways (Demmig-Adams et al., 1996; Asada, 1999; Oquist and Huner, 2003; Munekaga et al., 2004; Kanervo et al., 2005; Ensminger et al., 2006). Recently, the carotenoid zeaxanthin in combination with the PSII subunit PsbS has gained interest as both play an important role in the safe thermal dissipation of excess energy via nonphotochemical quenching (NPQ; Li et al., 2000, 2002; Savitch et al., 2002; Niyogi et al., 2005). NPQ actually consists of three different components that are commonly defined by referring to their relaxation properties in darkness following a period of illumination (Muller et al., 2001). A dynamic quenching mode is rapidly inducible and driven by the conversion of violaxanthin into zeaxanthin and protonation of the PsbS protein in response to acidification of the thylakoid lumen. State transition quenching relaxes within tens of minutes and is assumed not to be of great importance for photoprotection in terrestrial plants under high light (Niyogi, 1999; Kanervo et al., 2005). The sustained quenching mode ($q_s$) in turn is induced in connection with a decrease in PsbS levels and a loss of functional PSII (Ensminger et al., 2004; Ebbert et al., 2005). In evergreen conifers, this $q_s$ is also associated with the termination of growth and induction of frost hardiness (Oquist and Huner, 2003). In this $q_s$ state, a reorganization of the photosynthetic apparatus may protect conifer needles by dissipating excess energy as heat independent of a pH gradient (Gilmore and Ball, 2000; Horton and Ruban, 2005; Ensminger et al., 2006). Overwintering evergreen conifers seem to be able to shift between the dynamic and the sustained antenna quenching ($q_A$) modes for dissipating excess energy (Oquist and Huner, 2003). Aside from NPQ of excess energy in the antenna, a zeaxanthin independent way of quenching has been described in the reaction center (RC; Ivanov et al., 2001, 2006; Lee et al., 2001; Sane et al., 2003; Finazzi et al., 2004). In addition to antenna and RC quenching, excess energy can be funneled into a range of alternative electron sinks, most notably photorespiration (Osmond and Grace, 1995; Streb et al., 1998), the water-water cycle (Asada, 1999), and cyclic electron transport (Ivanov et al., 2001; Kanervo et al., 2005). For recent reviews, see Scheibe et al. (2005) and Ensminger et al. (2006).

Over the past decades substantial warming has occurred in northern latitudes, especially in the winter (Serreze et al., 2000). It has been suggested that this is likely to increase the length of the growing season and thus the productivity of northern hemisphere forests (White et al., 2000; Saxe et al., 2001). Increased temperatures in autumn allow trees to maintain a higher quantum yield (Methy et al., 1997). However, little data are available regarding the metabolic responses of evergreens of the boreal forests to extended growing seasons as a result of warmer air temperatures in late autumn. In this study, we focus on the interaction of altered autumn growth conditions and photosynthesis in Jack pine (Pinus banksiana Lamb.), an important evergreen conifer forming dominant stands on dry sandy soils in the western boreal zone of Canada, e.g. Northern Saskatchewan. Climate change will alter the phasing of temperature and daylength, the signals that trigger low temperature acclimation and development during autumn in conifer trees. The question is how the seasonal development is affected once trees experience the naturally decreasing daylength regulating growth cessation and limiting sink capacity while at the same time increased air temperature allows to maintain photosynthetic and respiratory capacity. Here, we used a factorial experiment with 16-h versus 8-h photoperiod (representing long day during July/August and short day during October/November, respectively, at e.g. 54°N, 105°W in northern Saskatchewan) and 22°C versus 7°C (summer and autumn, respectively) to separate the effects of daylength and temperature on photosynthesis and respiration to distinguish which processes are regulated by temperature or daylength only or by a combination of these abiotic factors. In particular, we focused on the effects of these treatments on the acclimation of photosynthetic capacity and energy dissipation and the composition of the photosynthetic apparatus.

**RESULTS**

**Gas Exchange**

Needle level gas exchange was performed on samples of seedlings of all four treatments (Fig. 1). At 1,000 μmol photons m⁻² s⁻¹, we observed the highest rate of light-saturated net assimilation ($A_{sat}$) in the summer conditions with 16-h photoperiod and 22°C (LD/HT) treatment. In the natural autumn control with 8-h photoperiod/7°C (SD/LT), $A_{sat}$ was decreased by 19%. Within the other two treatments, $A_{sat}$ was considerably lower, with the warm autumn conditions with 8-h photoperiod/22°C treatment (SD/HT) showing 41% lower values than LD/HT and 28% lower values than SD/LT. This pattern was also observed when net assimilation was measured at growth light conditions (Fig. 1). While assimilation did not show any clear effect of either temperature or photoperiod and only showed a significant difference in the interaction of both factors, we observed a clear response of respiration to either factor plus the interaction of both factors. Within the warm temperature treatments, the respiration rate in LD/HT was more than twice the rate observed in SD/HT (Fig. 1), suggesting an effect of the shorter photoperiod. Low temperature imposed an additional effect on dark respiration, being responsible for a further decrease in cold summer conditions with 16-h photoperiod/7°C (LD/LT) and SD/LT needles. These results are also valid when data are expressed on a fresh weight basis because the ratio of fresh weight to leaf area (grams per meter) only changes minimally between treatments over the duration of the...
Figure 1. The effect of daylength and temperature on Chl fluorescence parameters. Bars indicate light-saturated net CO₂ assimilation, measured at 1,000 µmol photons m⁻² s⁻¹ PPFD (A₅₀), CO₂ assimilation at growth light conditions of 350 µmol photons m⁻² s⁻¹ PPFD (A₁), and Black bars indicate dark respiration measured after 20 min of dark acclimation. All measurements were performed at growth temperature (22°C in LD/HT and SD/HT, 7°C in LD/LT and SD/LT plants). Each bar represents the average of n = 7 ± se biological replicates. ** and * Significant differences due to daylength, temperature, and their interactive effect, respectively. Two symbols, P ≤ 0.01; three symbols, P ≤ 0.001.

Chl Fluorescence

Chl a fluorescence was measured simultaneously with gas exchange. Maximum photochemical efficiency of PSII in the dark-adapted state (Fᵥ/Fₘ) was highest in the LD/HT control (Table I). The sole effect of a decreased length of the photoperiod resulted in a 10% decrease of Fᵥ/Fₘ in SD/HT plants. Low temperature had a much stronger effect on decreasing Fᵥ/Fₘ in the LD/LT compared to the LD/HT treatment. The effect was additive when combining both factors in the SD/LT treatment (51% decrease) with no interactive effect between daylength and temperature. In both high temperature treatments the quantum yield of PSII did not differ significantly from each other, but low temperature decreased the yield to less than one-half in LD/LT and to about one-quarter in SD/LT compared to the LD/HT treatment (Fig. 2A). qₑ as a measure of the fraction of open PSII RCs revealed large differences that were temperature dependent but did also depend on the interactive effect of daylength and temperature; e.g. values of the SD/HT treatment were 56% higher than in the summer control LD/HT treatment (Fig. 2B). Using nonphotochemical (qₒ) and antenna quenching (qₛ), we calculated the ratio of qₒ/qₛ to indicate the relative component of the qₒ from all nonphotochemical processes (Table I). qₒ/qₛ values were lowest during the typical LD/HT treatment, whereas under SD/HT conditions or under SD/LT conditions, the respective values were 40% and 38% higher than in LD/HT (Fig. 2C), showing solely a photoperiod dependence.

Effects of Photoperiod and Temperature on the Aggregation State of Chl-Protein Complexes

Nondenaturing SDS-PAGE was used to estimate the amount of Chl-binding protein complexes in the thylakoid membrane. The ratio of monomeric light-harvesting complex of PSII (LHCII³) and dimeric LHCII (LHCII²) to oligomeric LHCII (LHCII¹) was determined from gel scans to assess the degree of aggregation of LHCII, with a low ratio indicating a high aggregation state of the LHCIIIs (Fig. 3). The bulk of LHCII in all four treatments was found in form of oligomeric LHCII¹; only one-fifth to one-third depending on the treatment, was found as LHCII² and LHCII³. In LD/HT controls, the ratio of LHCII²¹/LHCII³ was 0.66 ± 0.23 and thus about twice the ratio determined in SD/HT (0.35 ± 0.07), LD/LT (0.33 ± 0.08), and SD/LT (0.29 ± 0.08), which indicates a much higher aggregation state in the latter treatments.

Photosynthetic Pigments

Chl a and Chl b levels differed by less than 10% in the two warm temperature treatments, and Chl b increased significantly in the two cold treatments. Photoperiod had no effect on Chl levels (Fig. 4A). However, because the low temperature increases in Chl b exceeded the slight increases in Chl a, we observed a temperature-dependent decrease in Chl

Table 1. The effect of daylength and temperature on Chl fluorescence parameters

<table>
<thead>
<tr>
<th>Treatment</th>
<th>LD/HT</th>
<th>SD/HT</th>
<th>LD/LT</th>
<th>SD/LT</th>
</tr>
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<tbody>
<tr>
<td>Fᵥ/Fₘ</td>
<td>0.80 ± 0.01</td>
<td>0.72 ± 0.02</td>
<td>0.51 ± 0.05</td>
<td>0.39 ± 0.02</td>
</tr>
<tr>
<td>qₒ</td>
<td>0.36 ± 0.03</td>
<td>0.52 ± 0.02</td>
<td>0.35 ± 0.05</td>
<td>0.38 ± 0.03</td>
</tr>
<tr>
<td>qₛ</td>
<td>0.93 ± 0.01</td>
<td>0.96 ± 0.01</td>
<td>0.74 ± 0.06</td>
<td>0.71 ± 0.05</td>
</tr>
</tbody>
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| Significance | Interactive Effect |
|--------------|-------------------|-------------------|-------------------|-------------------|
| Daylength    | Temp              |                   |                  |                  |
| n.s.         | n.s.              |                  |                  |                  |
| **           | **                |                  |                  |                  |
| ***          | *                 |                  |                  |                  |
| n.s.         |                   |                  |                  |                  |

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a/Chl b (Fig. 4B), which also showed an interactive effect. Total carotenoids per total Chl was constant among all treatments, except for SD/HT needles, which revealed a significant increase of about 15% compared to the other three treatments (Fig. 4C). In general, the share of neoxanthin and lutein made up approximately one-half of the carotenoid pool and did not vary much between treatments (Fig. 5). In contrast, the fractions of the remaining carotenoids were more variable, indicated, for example, by an increase in β-carotene and an almost complete absence of zeaxanthin in the SD/HT treatment. In turn, this treatment showed the highest amount of violaxanthin but the least amount of antheraxanthin. As a result, the deep-oxidation state was considerably lower in SD/HT, reaching only about one-quarter of the values obtained in the other treatments (Fig. 4D). This effect has been observed in two independent experiments carried out in two separate years.

Changes in Proteins of Photosynthesis and Carbohydrate Metabolism

The acclimation in response to daylength and temperature (Fig. 6) shows the highest levels of RC proteins of PSII and PSI (PsbA and PsaA/B, respectively) under LD/HT conditions compared to decreased levels in cold temperature treatments with lowest values in the LD/LT treatment. However, although the overall amount of RC proteins was variable among the treatments, the ratio of PsaA/B over PsbA remained constant (Fig. 6). PsbS levels decreased by 19% in response to short photoperiod and by 15% in response to low temperature as compared to the summer control. The additive effect of decreased photoperiod and low temperature resulted in PsbS levels in the SD/LT treatment to be 34% lower than in LD/HT, indicating that its regulation is independent with no interactive effect of temperature and daylength (Fig. 6).

There was also a clear response of the LHC proteins Lhca1, Lhca4, and Lhcb5 to daylength (Fig. 6). Protein levels in LD/HT and LD/LT clearly exceeded the levels in SD/HT and SD/LT, while there was no effect of temperature. By contrast, Lhcb1 was significantly affected by temperature, as indicated by the accumulation

Figure 2. A, The effect of daylength and temperature on quantum yield of PSII under steady-state condition at 1,000 μmol photons m⁻² s⁻¹ PPFD. B, Estimated qP, qN as an estimate of the amount of qO. All measurements were performed at growth temperature (22°C for LD/HT and SD/HT and 7°C for LD/LT and SD/LT). Each bar represents the average of n = 8 ± SE biological replicates. ⋆, ⋄, and ⋆, Significant differences due to daylength, temperature, and their interactive effect, respectively. Two symbols, P ≤ 0.01; three symbols, P ≤ 0.001.

Figure 3. The effect of daylength and temperature on the degree of aggregation of LHCII. Bars indicate the ratio of LHCII⁺ and LHCII⁻ to LHCII as determined from nondenaturing SDS-PAGE. Each bar represents the average of n = 4 ± SE biological replicates.
of this protein, with LD/LT being 35% and SD/LT being 26% higher than LD/HT and SD/HT levels (Fig. 6). Lhca2 and Lhcb2 exhibited minimal differences between LD/HT and the three other treatments (Fig. 6). The accumulation of the large subunit of Rubisco (RbcL) followed the same pattern observed for the RC proteins of PSII and PSI. Cytosolic Fru-1,6-bisphosphatase (cFBPase) and UDP-Glc pyrophosphorylase (UGPase), both cytosolic enzymes that are required to convert triose phosphate exported from the chloroplast into Suc, were clearly affected by daylength and UGPase also by temperature. Short-day conditions alone induced the

**Figure 4.** The effect of daylength and temperature on the composition of photosynthetic pigments in needles of Jack pine. A, Total Chl per fresh weight; Chl b was affected by temperature. B, Chl a to Chl b ratio. C, Total carotenoids per total Chl. D, Deepoxidation status of the xanthophyll cycle pigments, calculated as \((0.5A + Z)/(V + A + Z)\). Each bar represents the average of \(n = 8 \pm \text{SE}\) biological replicates. *, Significant differences due to daylength, temperature, and their interactive effect, respectively. One symbol, \(P \leq 0.05\); two symbols, \(P \leq 0.01\); three symbols, \(P \leq 0.001\).

**Figure 5.** The effect of daylength and temperature on the carotenoid composition in needles of Jack pine. Each carotenoid component is normalized to the total amount of carotenoids. The relative size of each pie reflects the amount of carotenoids based on a per total Chl basis. \(βC\), \(β\)-carotene; \(V\), violaxanthin; \(A\), antheraxanthin; \(Z\), zeaxanthin; \(L\), lutein; \(N\), neoxanthin. Values represent the average of \(n = 8 \pm \text{SE}\) biological replicates.
accumulation of cFBPase by 27% and exposure to low temperature by 13%. A similar pattern was observed in the expression of UGPase (Fig. 6).

DISCUSSION

Short Photoperiod and Low Temperature Promote Down-Regulation of Photosynthetic Capacity

It is remarkable that net CO₂ assimilation in the SD/HT treatment is considerably lower than in the LD/HT and comparable to that of the two low temperature treatments (Fig. 1). Photosynthesis therefore appears to be not only temperature dependent but is also strongly influenced by the length of the photoperiod. In conifers, shortened daylength acts as a signal for the induction of terminal budset and the cessation of growth (Repo et al., 2001). In combination with low temperature, this is an early autumn prerequisite to induce freezing resistance and therefore an important mechanism to prepare for the harsh winter conditions in northern forest environments (Bigras et al., 2001). This actual effect of daylength on photosynthesis probably reflects feedback regulation through an overall decrease in metabolic activity and growth. Such down-regulation of photosynthesis due to a reduced sink capacity is associated with the cessation of growth (Savitch et al., 2000). A decreased rate of CO₂ uptake is therefore likely due to the cessation of growth in pine after cold acclimation. Carbohydrate partitioning undergoes a tremendous shift as growth ceases and metabolic activity decreases under short photoperiod and low temperature. Increased expression of cFBPase, one of the key regulatory enzymes controlling the conversion of triose phosphate exported from the chloroplast into Suc (Strand et al., 2000), is associated with short photoperiod (Fig. 6) and indicates increased capacity for Suc synthesis. There is a dual role for Suc during autumn; it can be exported to the roots to be used for starch synthesis for winter storage and it may serve as a cryoprotectant to increase frost resistance in needle tissue. This clearly shows that the regulation of an important pathway for the cold hardening process strictly requires proper phasing of low temperature and daylength. However, levels of UGPase, acting downstream of cFBPase in the Suc synthesis pathway (Kleczkowski et al., 2004), did not show this exclusive increase due to short daylength but increased also in response to temperature (Fig. 6).

Down-regulation of metabolic processes in response to low temperature and short photoperiod was also reflected in decreased rates of dark respiration in SD/HT, LD/LT, and SD/LT compared to LD/HT. The response of plants to a shorter daylength was a more than 50% decrease in the rate of respiration. However, as a result of warmer temperatures, the respiration rate in the SD/HT treatment remained higher than in the SD/LT treatment.

Acclimation of Photosynthetic Energy Conversion and Composition of the Photosynthetic Apparatus to Growth Conditions

Decreased sink capacity requires acclimation of the energy partitioning process to balance the flow of...
energy between energy captured by the light reactions and the energy utilized metabolically. This may be achieved by changing properties of the thylakoid membrane-bound LHC proteins, thereby altering the efficiency of capture, conversion, and dissipation of light energy. Several of the LHC proteins showed a response to photoperiod. Lhca1, Lhca4, and Lhcb5 reached the lowest levels in the SD/HT treatment (Fig. 6). This coincided with a low deepoxidation state of the xanthophyll cycle pigments and a lack of zeaxanthin (Figs. 4 and 5), pigments that are preferentially bound in bulk by these proteins (Morosinotto et al., 2002). The down-regulation of photosynthesis in autumn was reflected in partial losses of PSII and PSI RCs as well as the RbcL content, which clearly reflects adjustment of photosynthesis to the reduced need for photosynthates under cold temperatures (see also Zarter et al., 2006). However, in the SD/HT treatment, the decrease of PsbA was balanced by an increase in the fraction of open RCs of PSII, resulting in a similar yield as in LD/HT (Figs. 2, A and B, and 6). While in LD/HT the photosynthetic apparatus remains fully functional and associated with a considerable sink for photosynthates, in low temperature treatments a reduced sink capacity was clearly accompanied by a decrease in the photochemical efficiency. A decrease in photochemical efficiency was not apparent in SD/HT plants, reflecting an energetic imbalance in these plants. Apparently this imbalance was not reflected by qN (Fig. 2B), suggesting alternative sinks for electrons that could include photorespiration, water-water cycle, and cyclic electron transport. Increased photorespiration has previously been shown to be an effective photoprotective strategy in high mountain plants under low temperature (Streb et al., 1998). Ivanov et al. (2001) suggested that cyclic electron transport around PSI is necessary to dissipate excess energy and to retain functionality of the photosynthetic apparatus in winter-stressed pine needles. Our observation of a significant increase in β-carotene in SD/HT (Fig. 5) points toward an increased formation of singlet-excited oxygen ($^1\text{O}_2^*$), which can efficiently be quenched by β-carotene (Cantrell et al., 2003; Krieger-Liszkay, 2005; Telfer, 2005). $^1\text{O}_2^*$ is generated from singlet-excited Chl via the Chl triplet state, if the energy of excited Chl cannot be used for photochemistry or dissipated as heat (Adams et al., 2004). There are at least two possible sites of an increased production of $^1\text{O}_2^*$ (Krieger-Liszkay, 2005). It could either originate from the antenna, where singlet Chl cannot be quenched due to low concentrations of zeaxanthin and hence passes its energy down to produce $^1\text{O}_2^*$, or in the RC of PSI via charge recombination. Here, β-carotene is the principal quencher of excess energy because triplet Chl cannot be quenched directly. In this case, charge recombination would contribute to filling the gap between electrons generated, as measured by the yield and electrons used by CO$_2$ fixation in the SD/HT treatment.

Dissipation of Excess Energy Depends on the Aggregation State of LHCII

Not only did the yield of photochemistry differ between treatments but we also observed different strategies to dissipate energy that is in excess to be used for photochemistry. It has been suggested that high $q_O$ in relation to $q_N$ is an indicator for high $q_O$ (Bukhov et al., 2001; Sane et al., 2003). We found $q_O/q_N$
values in the LD/HT treatment considerably lower than in the other treatments (Fig. 2C). Previous work also showed that a higher amount of aggregated LHs is associated with improved photoprotection in overwintering evergreens and other plants (Horton et al., 1991; Ruban et al., 1993; Ottander et al., 1995; Gilmore and Ball, 2000; Krol et al., 2002). Reorganization of pigment-protein complexes, which is indicated by an increased aggregation state of LHII, was observed in all but the LD/HT control treatment (Fig. 3) and typically coincides with higher qE in these treatments. Short photoperiod, as well as low temperature, seem to provide a separate signal that invokes elements of the cold acclimation pathway, resulting in a structural reorganization of the photosynthetic apparatus. Through a constitutive and nonregulated quenching component facilitated by aggregated antenna complexes, SD/HT plants that have been grown under the same temperature regime as the LD/HT plants do not require to accumulate large quantities of zeaxanthin and hence show a much lower deepoxidation of the xanthophyll cycle pigments. Based on these observations, we conclude that zeaxanthin is not the main component required to facilitate the quenching of excess energy in the SD/HT treatment.

Our observations support the LHII conformation model for NPQ proposed by Horton et al. (2005). In this model, the amount of quenching is regulated within minutes by the aggregation state of LHII in combination with binding of either violaxanthin or zeaxanthin, with the highest quenching efficiency resulting from an aggregated configuration binding zeaxanthin. A similar mechanism might be responsible for the acclimation to seasonal changes in photoperiod and temperature (Fig. 7), with the zeaxanthin-binding aggregated state representing the qL component of NPQ (Fig. 7C).

**Does Increased Autumn Air Temperature Increase Photosynthetic Gain?**

Our results indicate an experimentally extended growing season does not necessarily result in increased CO2 uptake and carbon gain in an evergreen conifer. On the contrary, short-day photoperiod and warm temperatures might even have the opposite effect due to increased rates of respiration and decreased maximum capacity for carbon uptake. Based on these experiments using seedlings in climate controlled chambers, we cannot predict how short photoperiod and warm autumn temperature might affect entire boreal forests in the future. In addition, in our treatments we used a relatively large temperature difference in the SD/LT (7°C/5°C) versus the SD/HT (22°C/18°C, warm autumn) treatment compared to the predicted increase of mean annual land air surface temperature that is in the range of 4°C to 6°C by the end of 2100 (IPCC, 2001). Nonetheless, our results suggest that increased autumn air temperature has the potential to interrupt the regulation of the seasonal development in conifer trees. One can now speculate that there is a temperature within the range of predicted climate change at which field-grown trees cannot behave optimally and thus cannot exploit the growing season optimally. The temperature increase apparently alters the phasing of the two critical environmental stimuli, thereby not only decreasing the sink capacity of the trees but even turning it into a potential source for the respiratory release of CO2. Thus, photosynthetic down-regulation due to periodic control of growth cessation during the autumn appears to offset any potential carbon gain resulting from a prolonged growing season in the autumn (Saxe et al., 2001). In response to experimentally increased autumn temperature, Jack pine seedlings exhibit an energy-quenching mechanism that does not follow the well-described PsbS- and zeaxanthin-dependent dissipation pathways. Thus, further investigations are essential to determine whether our findings from controlled environment experiments are consistent with climatic changes under natural environmental conditions on an ecosystem scale.

**MATERIALS AND METHODS**

**Plant Material and Growth Conditions**

Rooted Jack pine (Pinus banksiana Lamb.) seedlings were obtained from a local nursery (Somerville Seedlings) and planted in a mixture of ProMix (Premier Horticulture) and low nutrient mineral sand (1:2, v/v). The plants were kept outside underneath a light shelter for 1 year. In the second year, plants were transferred to controlled experimental summer conditions at the end of July, 2005 (Conviron growth chambers). Following 10 d at experimental summer conditions (see below) to allow for acclimation to chamber conditions, plants were exposed for 4 weeks to either 22°C/18°C (day/night) with a photoperiod of 16 h (LD/HT), 22°C/18°C with an 8-h photoperiod (SD/HT), 7°C/5°C with a 16-h photoperiod (LD/LT), or 7°C/5°C with an 8-h photoperiod (SD/LT). All treatments were provided with a photosynthetic photon flux density (PPFD) of 350 μmol photons m⁻² s⁻¹.

**Photosynthetic Gas Exchange**

CO2 exchange rates were measured on detached current year needles using a LiCor 6202 infrared gas analyzer connected to a modified LD2/3 cuvette (Hansatech). The needles were collected right before the measurement in the early afternoon, after seedlings had been exposed to growth light of 350 μmol m⁻² s⁻¹ for at least 4 h. The CO2 concentration was maintained at 375 ppm in air with 21% O2. Dark respiration was measured in these needles after 20 min of dark adaptation. Subsequently, plants were exposed to a PPFD of 350 μmol m⁻² s⁻¹ for 7 min to obtain measurements of steady-state photosynthesis, followed by a shift to a saturating PPFD of 1,000 μmol m⁻² s⁻¹ for another 7 min. Steady-state photosynthesis was usually attained within 3 to 6 min, depending on actinic light intensity and temperature. Gas exchange rates are averages over a measuring period of 30 s. All gas exchange measurements were performed at growth temperature.

**Chl Fluorescence**

Chl a fluorescence was measured with a PAM 2100 Chl fluorometer (Heinz Walz). The fiber optic of the PAM 2100 was connected to the LD2/3 Hansatech cuvette via a custom-made port to allow simultaneous fluorescence and gas exchange measurements after seedlings had been exposed to growth light of 350 μmol m⁻² s⁻¹ for at least 4 h (see above). Initial (minimum) PSII fluorescence in the dark-adapted state (F0) and Fm were determined after 20 min of dark adaptation in the cuvette. Fv’/Fm’ and transient fluorescence (Ft) were obtained concomitantly with the gas exchange measurements after
steady-state photosynthesis was achieved (Sveshnikov et al., 2006). Optimum quantum efficiency of PSI was calculated as $F_o/F_m = (F_m - F_o)/F_m$ and the quantum yield of PSI in the light as $F_o/F_m' = (F_m' - F_o)/F_m'$ (Genty et al., 1989). The fraction of PSI RCs in an open state was estimated as $q_o = (F_m' - F_o)/(F_m' - F_o)$ (Schreiber and Bilger, 1987). Nonphotochemical and antenna quenching were calculated as $q_o = 1 - (F_m' - F_o)/(F_m' - F_o)$ and $q_o = 1 - F_o/F_m'$ according to Rees et al. (1990).

Isolation of Thylakoid Membranes and Separation of Chl-Protein Complexes

Thylakoids of fresh needles were isolated at 4°C according to Krol et al. (2002) by grinding needles in 50 mM Tricine, pH 7.8, containing 0.4 M sorbitol, 10 mM NaCl, 5 mM MgCl₂, and 20% (w/v) polyethylene glycol 4000. The ground samples were filtered through Miracloth, followed by three washing steps in double-distilled water, 1 mM EDTA, and washing buffer, containing 50 mM Tricine, pH 7.8, 10 mM NaCl, and 5 mM MgCl₂ by centrifugation at 4°C at 5000g for 10 min. The pellets were solubilized in 300 mM Tricine, pH 8.8, containing 13% (v/v) glycerol, 0.1% (w/v) SDS, and 0.45% (w/v) dodecylmaltoside:Chl ratio of 20:1 (w/w). The Chl-protein complexes were separated in the dark at 4°C in nondenaturating 1.5 M Tris-HCl polyacrylamide gels with a Tris/Gly running buffer that contained 0.2% (w/v) Deriphat 160. The gels were scanned at 671 nm and the relative amount of Chl in each complex was determined as ratio of the peak area to the total area.

Photosynthetic Pigments

Needle samples for the pigment extraction were taken around noon after seedlings had been exposed for 4 h to growth light of 350 μmol m⁻² s⁻¹. Needles were ground to a fine powder in liquid nitrogen and extracted for 2 h in the dark on ice in 100% acetone buffered with NaHCO₃. Extracts were separated by HPLC with a Spherisorb ODS-1 analytical column (S.P.E.), containing 4% (w/v) SDS, 15% (w/v) Suc, 20 mM dithiothreitol, and Complete, EDTA-free, proteinase inhibitor cocktail (Roche Diagnostics). Membranes were solubilized in 300 mM Tricine, pH 7.8, 10 mM NaCl, and 5 mM MgCl₂ by centrifugation at 4°C at 5000g for 10 min. The pellets were solubilized in 300 mM Tricine, pH 8.8, containing 13% (v/v) glycerol, 0.1% (w/v) SDS, and 0.45% (w/v) dodecylmaltoside:Chl ratio of 20:1 (w/w). The Chl-protein complexes were separated in the dark at 4°C in nondenaturating 1.5 M Tris-HCl polyacrylamide gels with a Tris/Gly running buffer that contained 0.2% (w/v) Deriphat 160. The gels were scanned at 671 nm and the relative amount of Chl in each complex was determined as ratio of the peak area to the total area.

Protein Extraction, SDS-PAGE, and Immunoblotting

For protein extraction, needles were ground to a fine powder in liquid nitrogen. Forty milligrams of sample were extracted in 800 μL of ice-cold extraction buffer for 15 min on ice followed by 15 min of extraction at room temperature. The extraction buffer consisted of 60 mM Tris-HCl, pH 6.8, 10 mM NaCl, 5 mM MgCl₂, and 20% (w/v) polyethylene glycol 4000. The ground samples were filtered through Miracloth, followed by three washing steps in double-distilled water, 1 mM EDTA, and washing buffer, containing 50 mM Tricine, pH 7.8, 10 mM NaCl, and 5 mM MgCl₂ by centrifugation at 4°C at 5000g for 10 min. The pellets were solubilized in 300 mM Tricine, pH 8.8, containing 13% (v/v) glycerol, 0.1% (w/v) SDS, and 0.45% (w/v) dodecylmaltoside:Chl ratio of 20:1 (w/w). The Chl-protein complexes were separated in the dark at 4°C in nondenaturating 1.5 M Tris-HCl polyacrylamide gels with a Tris/Gly running buffer that contained 0.2% (w/v) Deriphat 160. The gels were scanned at 671 nm and the relative amount of Chl in each complex was determined as ratio of the peak area to the total area.

**ACKNOWLEDGMENT**

The authors are grateful to Dr. P.E. Jensen (The Royal Veterinary and Agricultural University, Copenhagen) for providing the PSI antibody.

Received October 30, 2006; accepted January 12, 2007, published January 26, 2007.

**LITERATURE CITED**


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